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In vitro multiplication of *Oroxylum indicum* (L.) through apical and axillary buds and determination of genetic fidelity using RAPD markers

4

6 ABSTRACT

7 Present study involves to develop a protocol for large scale multiplication of Oroxylum indicum L. 8 plants through apical and axillary buds using in vitro techniques. The apical and axillary buds were 9 cultured on Murashige and Skoog (MS) medium supplemented with different combinations of 6-10 benzylaminopurine (BAP) and kinetin. The maximum number of shoots were observed on MS 11 medium supplemented with 8.88 µM of BAP. Rooting was observed in half-strength MS medium 12 supplemented with 8.88 µM of BAP and 7.38 µM of Indole-3-butaric acid. The well-developed plants 13 were hardened in peat and perlite (1:1) and acclimatized in green house and 98% survival rate was 14 observed and then shifted to field for cultivation. Random amplified polymorphic DNA (RAPD) 15 markers were used to evaluate the genetic fidelity of the micropropagated plants. Ten different 16 decamer primers were used to amplify genomic DNA from *in vitro* field grown plants and the mother 17 plant to assess the genetic stability. The RAPD profile analysis revealed that micropropagated plants 18 has no somaclonal variations with the mother plants. These results suggest that the micropropagation 19 protocol developed by this study for rapid in vitro multiplication is appropriate and applicable for clonal 20 mass propagation of Oroxylum indicum which helps in silviculture developments.

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22 Key words: Genetic fidelity, In vitro, Micropropagation, Oroxylum indicum L, RAPD marker

23 **1. INTRODUCTION**

Oroxylum indicum is one of the important medicinal plant, belonging to the family Bignoniaceae. It is found in India, Sri Lanka, South - East Asia, Philippines and Indonesia up to an altitude of 1200 m [1]. This plant has been used as a single drug or as a component of certain polyherbal drug preparations in ayurvedic system of medicine. It is an active ingredient of well known ayurvedic formulations like Chyavanprash and Dashmularista [2].

29 O. indicum are germinating naturally by seeds, however seed dispersal is transpire through 30 anemochory and many seeds may reach unfavourable habitats resulting in poor seed germination. In 31 contrast owing the demand for its medicinal uses over exploitation and unsustainable harvesting are 32 the important factors to threatening its survival. Hence, there is an emerging need of skill to conserve 33 this taxon. At present, the plant tissue culture is a potentially useful technique for ex situ multiplication 34 and restoration of affected taxa. Keeping this in view, an attempt has been made to multiply O. 35 indicum through in vitro technique. Further genetic fidelity test would support our protocol for true to 36 type clones, since some times in vitro raised plantlets might exhibit somaclonal variation by numerous 37 factors such as prolonged culture period or genetic mutation [3, 4]. The morphological, biochemical 38 and molecular analysis have been developed to assure the genetic fidelity of *in vitro* derived clones. 39 However, most of these methods have their own limitations [5]. Polymerase chain reaction (PCR)-40 based techniques like random amplified polymorphic DNA (RAPD) are is very useful in establishing 41 the genetic stability of in vitro regenerated plantlets in many crop species [6, 7]. This method is 42 considered to be sensitive enough to detect the variations or genetic relationship among individuals 43 between and within species.

Quite a number of researchers have been published on *in vitro* multiplication of *O. indicum* however, these studies have not attempted fidelity test on *in vitro* raised hardened plants [8, 9, 10]. Nevertheless RAPD marker studies on *O. indicum* were reported by differentiating the genetic diversity of this plant from different geographic location [11]. Henceforth, the present investigation emphasizes the development of plants through *in vitro* technique and analysis of the genetic fidelity through molecular markers.

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53 2. MATERIALS AND METHODS

54 **2.1 Plant material and surface sterilization**

Apical and axillary buds of *O. indicum* were collected from near Alsoor lake, Bangalore, India and used as a source material. The collected specimens were identified using *The Flora of the presidency of* Madras [11]. These freshly collected apical and axillary buds were washed with running tap water containing soap oil for 30 min to remove the dust particles. The explants were then washed 59 twice with Tween 20 for 2 to 3 min and rinsed five to six times with distilled water. For surface 60 sterilization, the explants were immersed in 5.0% Bavistin (Carbendazim) for 1h to eliminate fungal 61 contamination followed by rinsing with autoclaved doubled-distilled water. Further, the explants were 62 sterilized with 70% alcohol for 2 to 3 min and 0.1% mercuric chloride for 2 to 3 min and rinsed three to 63 four times with sterile distilled water under aseptic conditions in laminar air flow chamber.

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2.2 Culture media and growth condition

65 The Murashige and Skoog's (MS) basal medium and supplemented with different 66 concentrations of growth regulators like 6-benzylaminopurine (BAP), kinetin (Kn) and with 3% sucrose 67 (w/v) as a carbon source were prepared [12]. The pH of the media was adjusted to 5.6 to 5.8 by using 68 0.1N HCI/NaOH, before addition of 0.8% agar (w/v) and then autoclaved at 121°C for 20 min. The 69 autoclaved medium was stored at room temperature for 24h to check microbial contaminations. The 70 cultures were maintained in culture room at a temperature 25 ± 2°C under built-in white fluorescent light at a photon flux density of 30-50 μ Em⁻²s⁻¹ under a photoperiodic regime of 16 h of light and 8 h 71 72 of dark cycles.

73 2.3 Regeneration of shoots and roots

74 For shoot regeneration, apical and axillary buds (Fig. 1) were cultured on MS medium fortified 75 with different concentrations of cytokinins like BAP (1.11 - 11.11 µM) and Kn (1.16 - 11.62 µM) 76 individually. Percentage of regenerated shoots and the number of shoots per explants were recorded 77 at the end of regeneration period. The shoots were subculture on half strength MS medium fortified 78 with BAP 8.88 μ M and different concentrations of auxins such as α -naphthalene acetic acid (NAA), 79 indole-3-acetic acid (IAA) and indole-3-butyric acid (IBA) with 2% sucrose(w/v) in all the combinations 80 for root development. The mean number of roots and its length were examined periodically up to 4 weeks of culture. 81

82 **2.4 Hardening and acclimatization**

Hardening and acclimatization is the most important steps in the micropropagation process, where it provides plants to tolerate and surviving in the external environmental conditions. The rooted plants were taken out from the culture bottles and washed thoroughly to remove the traces of agar medium and immersed in Bavistin (5%) solution for 5 min. to protect from fungal infection. Then plantlets were transplanted in to pot containing sterilized peat and perlite (1:1) and were kept in green house to maintain high humidity and watered at 2-day intervals. Survival rate of the hardened plants
were observed before transferring to the field.

90 **2.5 Genetic fidelity testing of** *in vitro* raised plants and mother plant

91 Fresh leaf samples of *o.indicum* (obtained from *In vitro*) were collected from The Himalaya Drug 92 Company campus and mother plants from near Alsoor Lake, Bangalore. The samples were washed 93 with running tap water in addition of soap oil for 10 min and rinsed with double distilled water to 94 remove surface contaminants. These cleaned leaves of both samples (100 mg each) were frozen 95 separately in liquid nitrogen and ground in a mortar and pestle to become a fine powder. Total DNA 96 was extracted from the leaves as per the manufacturer's instruction manual of Gen EluteTM Plant 97 Genomic Kit (Sigma Aldrich). The DNA concentration and purity were analysed by 0.8% agarose gel 98 electrophoresis with ethidium bromide staining and spectrometric method.

99 Polymerase chain reaction (PCR) amplification was carried out in 25 µL volume using 10 100 different decamer primers. The reaction mixture consisted of 1U of Tag DNA polymerase (Bangalore 101 Genei), 0.25 µL each of dNTP (10 mM), 5 pmole of decanucleotide primer (Sigma Aldrich), 102 1×polymerase buffer (2.5 µL), 0.25 mM MgCl₂ and 50 ng of DNA sample. The amplification was 103 performed using a thermal cycler (PTC-100TM, MJ Research. Inc., USA). The program consisted of 104 initial denaturation at 93°C for 4 min followed by 40 cycles of denaturation at 93°C for 1 min, 105 annealing at 37°C for 1 min, extension at 72°C for 2 min, and at a final extension cycle of 8 min at 106 72°C. The amplified products were checked in a 2% agarose gel stained with ethidium bromide and 107 documented by a gel documentation system (Pharmacia Biotech).

108 2.6 Data analysis

All the cultures were examined periodically. Each experiment was repeated 3 times, and subculturing was carried out at 4-week intervals. The mean number of shoots per culture and mean number of roots per shoot and their lengths were recorded. The data were analysed statistically by one-way ANOVA with Tukey's multiple comparison post hoc tests. The minimum level of significance was fixed at p<0.05 with two tailed p value. Statistical analysis was performed using GraphPad Prism version 4.03 for windows (GraphPad Software, San Diego, CA, USA).

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116 **3. RESULTS**

117 3.1 Shoot regeneration

118 Apical and axillary buds did not showed any response for shoot initiation on MS basal medium. 119 Different types and concentrations of growth hormones are used for shoot formation as shown in 120 Table 1. Shoot initiation was observed in all the six BAP combinations, however highest initiation 121 response was observed on MS medium fortified with BAP (8.88 µM) after 4 days of culture. After 30 122 days shoots were ready for first subculture, the shoots attain length (3 - 4 cm) and formed clusters. 123 The maximum number of multiple shoots (9.5 ± 0.10 shoots/explants) and shoot length (6.9 ± 0.44 124 cm) were observed after second subculture (60 days) on the same medium (Fig. 2). Further, 125 increasing the concentration of BAP showed decline in multiplication rate, in addition MS media with 126 Kn failed to induce shoot proliferation in all tested combinations (Table 1).

127 3.2 Rooting

In vitro rooting of multiple shoots was achieved on half strength MS medium enriched with 128 129 different concentrations of auxins (IBA, NAA, IAA) with cytokinin (BAP 8.88 µM) and 2% sucrose. The 130 significant induction of roots (7.5 \pm 0.14) per shoot with an average root length (3.9 \pm 0.20 cm) was 131 found in culture contain 8.88 µM of BAP with 7.38 µM of IBA within 20 to 25 days of subculture (Fig. 132 3). Whereas, number of root formation were declined with respect to its length in lower concentration 133 of IBA. Subsequently, NAA in combination of BAP shoed moderate development of roots when 134 compare to earlier concentrations. However IAA with BAP combinations were failed to produce roots 135 in tested media. As an inference, IBA is referred to as the sole auxin to induce rooting in in vitro 136 condition (Table 2).

137 3.4 Hardening of plantlets

The rooted plantlets were transferred to pots containing sterilized peat and perlite (1:1) and kept in the greenhouse for hardening (Fig. 4). After four weeks of incubation 98% survival rate was noticed in the greenhouse. Later the plants were transferred to a normal environment (Fig. 5).

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3.5 RAPD and genetic fidelity analysis

The evaluation of the genetic fidelity of tissue culture derived plants is an essential for detection of true-to-type clones. In the present study, homogeneity of genetic materials of *in vitro* derived plants was compared to mother plant with the help of RAPD technique. To our knowledge, no report was available till date on the comparative genetic stability of *in vitro* hardened plants and mother plant of *O. indicum* by using RAPD markers. Based on RAPD analysis, no somaclonal variations was observed in tested plants (Figs. 6-8). In the present study, total number of bands observed were 73

out of which 31 bands were monomorphic in nature (16.6 % - 50%). The number of amplification
products obtained by these primers are summarised in Table 3. The size of the monomorphic DNA
bands produced by the tested primers ranged from 250 bp to 2000 bp.

151 4. DISCUSSION

In the present study, standardized the protocol for multiple shoot formation from apical and axillary bud of *O. indicum*. Six different concentrations of growth regulators (BAP and Kn) were tested for shoot proliferation. The maximum number of shoots with significant shoot length was observed in MS medium fortified with 8.88 µM of BAP after second subculture. Out of two cytokinins BAP was identified as significant growth hormone for shoot initiation and multiple shoot formation. Earlier reports have shown that MS with BAP media stimulate the regeneration of plants and gives more shoot regeneration than Kn [13, 14]. The present finding is also agreement with earlier study [15].

The significant development of roots were observed in half strength MS medium containing 8.88 μm of BAP and 7.38 μM of IBA with 2% sucrose. In the present investigation, IBA is used as the sole exogenous auxin to produce rooting in *in vitro* regenerated shoots. The similar kind of success of IBA in promoting efficient root induction have been reported for *Clitoria ternatea* L., *Murraya koenigii* L. Spreng, and *Ocimum basilicum* L. [16, 17, 18].

In vitro regenerated plants do not survive without acclimatization, because it has to adjust many abnormalities in *ex vitro* environment like light, humidity and temperature. However, *in vitro* raised plants need gradual changes in environmental conditions to avoid desiccation, water relations and photoinhibition. In the present study, micropropagated plants were acclimatized [19, 20, 21] in primary hardening chamber and transferred to green house where 98% survival rate were found .

169 The evaluation of the genetic stability of tissue culture derived plants is an essential step to 170 assess true-to-type clones [22]. The present study confirms the homogeneity of micropropagated 171 plantlets compared with that of the mother plant by using RAPD technique. In general, the in vitro 172 raised plants obtained from apical meristems and axillary meristems do not alter any genetic makeup, 173 hence this mode of propagation is considered as safest method to multiply the plantlets [23, 24]. This 174 will further strengthen our RAPD data reveals there is no evidence of somoclonal variations between 175 in vitro raised hardened plants and mother plants of O. indicum. Hence, this is an efficient protocol for 176 the regeneration of O. indicum through apical and axillary buds. However, the present protocol will help the plant breeders and foresters able to generate disease-free, uniform parental clones withgenetic fidelity to generating the population.

179 **5. CONCLUSION**

180 The present investigation is focused on establishment of mass multiplication of O. indicum through 181 micropropagation tool. Researchers who have performed various studies have deduced protocols to 182 obtain plant-lets through high concentration of growth regulators,; whereas, the present study 183 provides a protocol to develop multiple shoots, roots and hardened plants in least concentration of 184 growth regulators with maximum number of shoots and roots followed by well developed plants that 185 are propagated in a normal environment successfully. The assay of genetic fidelity through RAPD 186 markers is a very important aspect for analyzing the somaclonal variation in in vitro propagated 187 plants. Plants developed in this study have not shown any variation in the genetic makeup compared 188 with the mother plant. This kind of investigation is carried out for the first time for O. indicum to 189 ascertain its stability and this protocol could be an important source for forest crop breeders, species 190 conservation and herbal industries

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250 Tables

 Table 1.Effect of different concentrations and combinations of cytokinin on development of multiple shoots in *Oroxylum indicum*

Growth regulator	rs concentration	Mean number of	Mean length of	
in MS me	in MS media (μM)		shoots (cm) \pm SE	
BAP	Kn	-		
1.11	<u>)</u>	8.5 ± 0.08	3.4 ± 0.15	
2.22	-	9.0 ± 0.11^{a}	4.1 ± 0.22	
4.44	-	9.2 ± 0.11^{a}	5.1 ± 0.24^{a}	
6.66	-	9.3 ± 0.11^{ab}	5.8 ± 0.30^{a}	
8.88	-	9.5 ± 0.10^{abcd}	6.9 ± 0.44^{abcd}	
11.11	-	8.6 ± 0.09^{abc}	$5.3\pm0.33^{\rm ac}$	
-	1.16	-		
-	2.32	-		

<i>p</i> value		<i>p</i> <0.001	<i>p</i> <0.001
-	11.62	-	
-	9.30	-	
-	6.97	-	
-	4.65	-	

Means within a column having the same letter did not reach a statistically significant (p<0.001) level

according to one-way ANOVA with Tukey's Multiple Comparison post hoc test.



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Growth regulators	concentration in MS	Mean number of roots	Mean length of roots	
media	a (µM)			
BAP	NAA	-		
8.88	1.35	6.2 ± 0.13	3.0±0.21	
8.88	2.70	6.4 ± 0.14	3.2±0.19	
8.88	5.40	6.8 ± 0.10^{a}	3.3±0.17	
8.88	8.10	$5.8\pm0.14^{\text{cb}}$	2.7±0.13	
8.88	10.80	5.3 ± 0.11^{abc}	2.2±0.20abc	
	IAA			
8.88	1.42	_	<u> </u>	
8.88	2.85		-	
8.88	5.71	$\sqrt{-1}$	-	
8.88	8.21		-	
8.88	11.42	· · ·	-	
	IBA			
8.88	1.23	5.0 ± 0.12	2.0 ± 0.26	
8.88	2.46	5.6 ± 0.15^{a}	2.6 ± 0.12	
8.88	4.92	6.5 ± 0.14^{ab}	3.0 ± 0.18^{a}	
8.88	7.38	7.5 ± 0.14^{abc}	3.9 ± 0.20^{abc}	
8.88	9.84	6.8 ± 0.11^{abc}	2.0 ± 0.26^{ad}	
<i>p</i> value		<i>p</i> <0.001	<i>p</i> <0.05	

Table 2 - Effect of plant growth regulators for development of roots

Means within a column having the same letter did not reach a statistically significant (p<0.05) level according to one-way ANOVA with Tukey's Multiple Comparison post hoc test.

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Sl.	Sequence (5'-3')	Oligo	No. of	Size of the	No. of	Mono-
No.		primer	bands	amplified	Mono-	morphic
		s name	obtained	bands (bp)	morphic	(%)
					bands	
1.	5'AGAGGCTCAC 3'	HD-1	7	390-2000	3	42.85
2.	5'GCGTTGACTG 3'	HD-2	4	400-600	2	50
3.	5'AGGGCAAACC3'	HD-3	8	250-1300	4	50
4.	5' CCCGTTACAC3'	HD-4	6	600-1300	2	33.3
5.	5'CATACGCCTC 3'	HD-5	12	300-1800	6	50
6.	5'CCGAAGACAC3'	HD-6	6	290-500	2	33.3
7.	5'GCAGTACGAG 3'	HD-7	7	350-1000	3	42.85
8.	5'GCAGTACGTG 3'	HD-8	10	300-1900	5	50
9.	5'CGAGGAGACC3'	HD-9	6	300-1900	1	16.6
10.	5'GTTCCCAGAC3'	HD-10	7	550-1200	3	42.85

Table 3 - The number of amplification products generated of RAPD primers in the analysis ofOroxylum indicum regenerants.





Figure 2

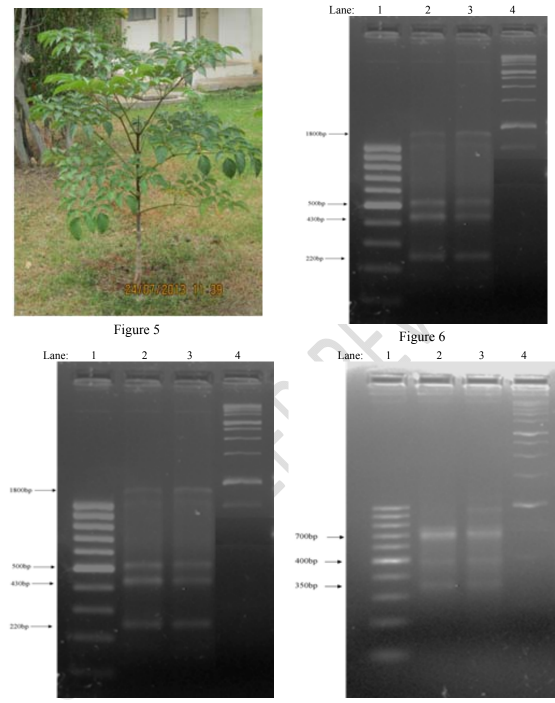






Figure 3

Figure 4







262 Figures legends

- 263 Fig 1 Apical meristems inoculated in medium
- 264 Fig 2 Regeneration of multiple shoots observed in MS medium contain BAP 8.88 µM
- 265 Fig 3 Development of roots in half MS medium with 2% sucrose and IBA (4.92µM)
- 266 Fig 4 Hardened plantlets of Oroxylum indicum in green house
- 267 Fig 5 In vitro raised Oroxylum indicum growing in a normal environment
- Fig 6 A RAPD profile generated by Oligo primer HD 3. Lane 1: Marker (100 bp); Lane 2: Mother plant; Lane 3: *In vitro*-raised hardened plant; Lane 4: Marker (1kb)
- Fig 7 RAPD profile generated by Oligo primer HD 5. Lane 1: Marker (100 bp); Lane 2: Mother
- 271 plant; Lane 3: *In vitro*-raised hardened plant; Lane 4: Marker (1kb)
- 272 Fig 8 RAPD profile generated by Oligo primer HD 7. Lane 1: Marker (100 bp); Lane 2: Mother
- 273 plant; Lane 3: *In vitro*-raised hardened plant; Lane 4: Marker (1kb)