Root initiation in cuttings and \textit{in vitro} raised shoots of \textit{Pinus roxburghii}

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\textbf{ABSTRACT}

Rooting of \textit{in vitro} produced buds and shoots is often the limiting step during micropropagation. Therefore, a better understanding of the various stages before and during root formation is needed. Reviewing the work done so far on pines the present investigation was carried out to study the factors that affect \textit{in vitro} rooting. Different parameters that influence rooting viz. donor age, phytohormones, and substrate were investigated. Shoots took from the field; it was found that the juvenility of the explant and position on the mother plant greatly affects the \textit{in vitro} responses. \textit{In vitro} raised shoots and hypocotyl cuttings of \textit{Pinus roxburghii} showed the best response over the other explants on $\times \frac{1}{2}$ DCR medium. Among the various auxins used in the present investigation, N-acetyl-aspartate (NAA) at lower concentrations found best for root initiation. Agar at 0.6% concentrations resulted in more healthy roots. Further elongation was achieved on $\times \frac{1}{2}$ DCR medium supplemented with lower concentrations of NAA. The present investigation was an attempt to establish an operative micropropagation protocol by improving the rooting of “hard to root” \textit{P. roxburghii}. \textit{In vitro} rooting studies on \textit{P. roxburghii} will be vital for enhanced multiplication and genetic improvement of this economically important forest tree species.

\textbf{KEY WORDS:} Auxins, hypocotyl, \textit{Pinus roxburghii}, rooting

\textbf{INTRODUCTION}

Two difficult stages of conifer tissue culture are rooting and acclimatization. The efficiency of \textit{in vitro} adventitious rooting is highly variable [1] and is the key problem in conifer plantlet regeneration. Juvenile plant material exhibit greater potential for adventitious rooting [2]. The difference in the rooting response by the difference in age groups in \textit{Pinus pinaster} was reported by Dumas and Monteuuis (1955) [3]. The size of shoot (more than 5 mm) [4] and the removal of callus from the base of shoots also found to show the improved effect on rooting [5].

Several aspects of rooting phytohormones viz. type, concentration, mode of application, and duration of response needs to be considered. For most of pine species, N-acetyl-aspartate (NAA) is reported as most potent auxin compared to indole-3-butyric acid (IBA) [4,6-9]. Low concentration of NAA generally found most effective for root induction in pines [10,11]. Combinations of NAA, indole-3-acetic acid (IAA), or IBA are more effective [6,9]. Sometime results in the formation of callus [11]. Root induction is also affected by the shoot quality [12], donor age, and genetic origin [13,14].

The shoot induction medium, type of cytokinins, and their concentrations also reported having profound effect on rooting [4,11] and sometime attributed as “carry-over-effect” of cytokinins at higher concentration [15]. Similarly, elongation medium also influences the rooting as reported for \textit{Pinus radiate} by Horgan and Aitken-Christie (1981) [16].

\textit{In vitro} rooting is usually done using the agar-solidified medium as the substrate. Advantages include uniform distribution of auxins and nutrients and good contact between shoots and substrate that results in more synchronous rooting. However, the quality of root produced on agar is not always acceptable [2]. Agar hinders gas exchange as well as the production of root hairs and development of the vascular system. Permeable substrates like peat:perlite or peat:vermiculite found more recommended [17]. Peat:vermiculite (1:1) moistened with $\times \frac{1}{2}$ medium was reported better by Pulido \textit{et al.} (1994) [11] in \textit{Pinus canariensis}. Similar findings were
reported by Horgan and Holland (1989) [5] in P. radiata, he suggested that peat:perlite:vermiculite (1:1:1) is better over agar. 50% rooting in soil under mist chamber in Pinus taeda was reported by Mehra-Palta et al. (1978) [4].

MATERIAL AND METHODS

Rooting of In Vitro Raised Shoots

In vitro raised shoots, hypocotyl cuttings from juvenile seedlings, shoots from 1 + year old plants, and hedge shoots (7 + year old) collected from different positions on mother plant were used as explants. All the explants were subjected to in vitro rooting. Detailed studies of rooting behavior in vitro because of different parameters: Shoot quality, donor age, growth regulators, and substrate were studied.

Effect of basal medium

Three basal media viz. Murashige and Skoog’s (MS), DCR, and Woody plant medium (WPM) were used in the present investigation and supplemented with 2.5 μM IAA.

Effect of medium strength

Four strengths of basal medium (DCR) viz. ×¼, ×½, ×1, and ×2 were used for rooting of different types of shoots of Pinus roxburghii.

Effect of auxins

Auxins viz. NAA, IBA, and IAA were supplemented in ×½ DCR medium at 2.5, 5.0, and 7.5 μM concentrations to investigate their effect on rooting.

Effect of agar concentrations

The gelling agent reported to have an important role in adventitious rooting of shoots. To study the effect of agar concentrations on root induction, half strength DCR medium supplemented with 2.5 μM NAA was gelled with 0.6, 0.7, or 0.8% agar.

Effect of pulse treatment

Mode of application of auxin is also known to effect rooting of the shoots. Auxins viz. NAA and IBA were provided at a concentration of 50 μM for 24 h in semisolid medium followed by culturing in semisolid ×½ DCR basal medium gelled with 0.6% agar.

Effect of liquid medium on elongation of root primordia

The shoots with root primordia produced on semisolid auxin supplemented media or in response to pulse treatment were transferred to either liquid (with Filter Paper Bridge) to study their effect on the elongation of root primordia. Liquid and semisolid DCR medium was supplemented with 2.5 μM NAA or IBA singly and in combination (1.25 μM NAA + 1.25 μM IBA).

RESULTS

Rooting of In Vitro Raised Shoots

Effect of basal medium

Basal medium showed differences in rooting from different types of shoots. DCR medium showed a best rooting response in all types of shoot material, especially the in vitro raised shoots, and juvenile seedling shoots showed a best rooting response (Graph 1). MS and WPM medium failed to produce optimum results. Mature shoots of Pinus from 1 + year and 7 + year old plants could not produce appropriate roots in vitro in any of three basal medium.

Effect of medium strength

Among the four strengths of basal medium (DCR) used for rooting of different types of shoots of P. roxburghii, the best rooting was obtained on ×½ strength of DCR medium. At this strength, all types of shoots showed good rooting response except the 7 + year mature shoots of Pinus. Hypocotyl shoots of Pinus seedlings showed the best rooting percentage (88%) followed by in vitro raised shoots (85%) and 1 + year old shoots (65%) (Graph 2).
Effect of auxins

Shoots from the different origin of 2.0-3.0 cm length were used for in vitro rooting. Auxins namely NAA, IBA, and IAA were used at varied concentrations in half strength DCR medium for induction of rooting. Success was obtained only in formation of root primordia at the base of the hypocotyl shoots and in vitro raised shoots when cultured under different experimental combinations of the auxins. It was found that NAA although initiates root primordia but later callus develops which overgrowth the developing root primordia. Root primordia were initiated in 33.31% shoots cultured on 2.5 μM NAA supplemented medium (Graph 3). Increase in NAA concentration led to a decrease in the number of root primordia induced with a simultaneous increase in callus formation. The root primordia produced in all NAA concentrations could not elongate on subculturing to fresh medium, as well as on transfer to the basal medium.

IBA and IAA when used at 2.5-7.5 μM levels produced root primordia in a lesser percentage of shoots. Similar to NAA the root initials formed could not elongate into roots (Graph 3).

Hypocotyl cuttings also responded towards rooting when cultured on DCR basal medium with various concentrations of auxins. NAA (2.5 μM) produced a best rooting response (30%) over other concentrations and auxin types (Graph 4).

Effect of agar concentration

The shoots when cultured on medium gelled with 0.8% agar (Bacteriological grade agar, BDH) produced stunted root primordia with profuse callusing while the shoots cultured in liquid medium were vitrified with no root formation. Thus, attempts were made to check the excessive callusing and induce roots in in vitro raised shoots by culturing them on a gelled medium of softer consistency. To study the effect of agar concentration on rooting, the in vitro raised shoots and hypocotyl cuttings were cultured on medium gelled with 0.8, 0.7, and 0.6% agar (Graph 5 and Graph 6). A limited success was achieved

Graph 3: Effect of different concentrations of auxins on rooting percentage, number of root primordia and callusing percentage in in vitro raised shoots

Graph 4: Effect of different concentrations of auxins on rooting percentage, number of root primordia and callusing percentage in hypocotyl cuttings
in reducing callus formation at basal ends of shoots. In addition, the percentage of root primordia formation was increased slightly. The root primordia induced in 5-7% shoots elongated into roots on further subculturing on basal medium. It was observed that the shoots producing no or moderate callus at shoot base developed root primordia but the shoots. Producing excessive callus at the base produced no root primordia or primordia, which could not elongate to form roots.

Effect of pulse treatment
Attempts were also made to induce rooting by providing a pulse treatment to individual shoots before transferring them for rooting on half strength basal medium. The in vitro raised shoots were cultured on semisolid medium containing high concentration (50 μM) of NAA or IBA individually for 24 h and then transferred to basal medium without auxins. In vitro shoots and hypocotyl cuttings showed the formation of root primordia at the base of 30.11% and 28.11% explants, respectively, (Graph 7 and Graph 8), some of these primordia elongated to a length of 1.0-2.0 cm.
DISCUSSION

Rooting is one of the major steps of micropropagation and affects by a number of factors. Phytohormones, nutrient medium, and physiology of shoots have a profound effect on in vitro rooting. Rhizogenesis requires a lower concentration of nutrient in the medium [2] and hence half strength DCR medium used in the present research was found successful for in vitro rooting response in chirpine. However, different medium with or without modification has been reported to give best rooting response from a variety of explants in Pinus [5,8,18]. Reduction in the medium strength resulted in enhanced rooting response as also reported earlier [11,19].

Three auxins viz. NAA, IBA, and IAA were added alone and in combination to study their effect on root induction. For chirpine, NAA (2.5 μM) was observed as best rooting hormone which coincide with the previous findings [9,18,20]. A lower concentration of NAA showed a best rooting response in present investigations which is also in line with previous findings by Murithii et al. (1993) [10]. However, callus formation was observed as a usual problem in rooting of pines. To overcome callusing, explants were exposed to pulse treatment of hormones. Exogenous application of auxin is generally required for rooting [21]. However, without the pre-treatment of auxin no significant difference in rooting was observed [22,23]. Continuous exposure of NAA (2.5 μM) in medium showed more number of root primordia formation over the pulse treatment of shoots with 50 μM NAA or IBA. Kaul (1987) [24] and Schwarz et al. (1988) [13] also found the same effect of pulse treatment on in vitro rooting.
Agar concentration (0.8%) was reduced to 0.6% as it was not allowing roots to penetrate into the medium and result in their radial growth and callusing at the base of shoots. This change resulted in more number of root formation per micro-shoots. Root primordia produced under the influence of auxins were small and needed elongation, and liquid media was found best for this compared to semisolid medium. The superiority of liquid medium for elongation of root primordia is possibly due to better aeration of the growing roots which is poor in the semisolid medium [2].

**In vitro** developmental stages of plants are having low light level, aseptic conditions and high amount of sugar and nutrient and growth factors and controlled light and humidity in the environment. Such a heterotrophic environment makes these plants more delicate and hard to sustain **in vivo**. Gradual acclimatization of **in vitro** plantlets makes them harder to sustain harsh field condition and provide minimal stress for plant multiplication. **In vitro** rooted plantlets of pine were exposed to further hardening and acclimatization.

**CONCLUSION**

In the present investigation, we attempted **in vitro** rooting of *P. roxburghii* shoots from different aged material as well as from **in vitro** raised shoots. Such an effort of tissue culture of coniferous species can significantly increase forest productivity with the production of selected genotypes. This **in vitro** regeneration protocol will provide an alternative to rooted cuttings for the propagation of conifers.

**REFERENCES**

