SOMATIC EMBRYOGENESIS AND CALLUS PROLIFERATION IN *Picrorhiza kurroa* ROYLE ex. BENTH

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**KEYWORDS**

*Picrorhiza kurroa*

*In vitro*

Endangered

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Somatic embryos

**ABSTRACT**

*Picrorhiza kurroa* Royle ex. Benth is an endangered and important medicinal plant of alpine Himalayas. Developing an efficient protocol for its mass multiplication is essential to meet the requirements of pharmaceutical industries and also in conservation of this plant under its natural habitat. Present study was undertaken to develop a protocol for *in vitro* mass multiplication of *P. kurroa*. Result of study revealed that highest frequency of shoot regeneration was achieved on Murashige and Skoog’s basal medium supplemented with 1.0 mg/l BAP, 0.5 mg/l Kn and 1.0 mg/l GA<sub>3</sub>, while the best rooting was observed in MS medium supplemented with 2.5 mg/l IBA. MS medium supplemented with 3.0 mg/l 2,4-D resulted in highest frequency of embryogenic callus. Callus inoculated on MS media supplemented with BAP and IAA resulted in both shoot and root formation while the callus on MS media supplemented with NAA and IBA resulted only root formation. The somatic embryos were established from callus on MS medium supplemented 2.5 mg/l 2,4-D after four weeks. MS medium containing 1.0 mg/l BAP and 1.0 mg/l GA<sub>3</sub> resulted into shoots from well developed somatic embryos. This protocol will provide a system for the germplasm conservation in *P. kurroa* by multiplication and regeneration of true to type plants.
1 Introduction

The increasing demand of medicinal plants for the development of several drugs and chemotherapeutics has threatened their natural habitats and now they are in brink of extinction (Mugula et al., 2012; Bodeker et al., 2013). These plants are demanding to be conserved for their potential utility. One of such plant is Picrorhiza kurroa Royle ex. Benth belonging to family Scrophulariaceae which is endemic to Himalayan region (Patial et al., 2012). The plant grows from Kashmir to Sikkim at an altitude of 3,000-5,000 meters. The rhizomes of the plant are the principle source of iridoid glycosides out of which the major is Picrosides and kutkosides (Patil et al., 2013). These iridoids have numerous therapeutic uses like antihepatotoxic, choleretic, hypolipidemic, antiinflammatory, antispasmodic (Tiwari et al., 2012; Sultan et al., 2016). The dried extracts from the plant has a cure for liver disorders, fever, jaundice and allergies (Baruah et al., 1998). The herbal preparation of the plant plays a major role in anti diabetic and anticancerous activity (Pradhan 2011; Kumar & Ramesh 2014; Mallick et al., 2015).

The over exploitation of the rhizomes of P. kurroa for the extraction of bioactive constituents along with the poor cultivation and small population size has depleted this species from its natural habitat (Verma et al., 2012). As a result, the plant is now listed as an endangered plant species in red data book by International Union for Conservation of Nature (IUCN). The conventional method of propagation using rhizomes and seeds restricts multiplication as seed germination is very poor in P. kurroa, so conservation using in vitro approaches is essential (Rawat et al., 2013).

It is necessary to develop a protocol for mass multiplication of this endangered high value medicinal plant so as to assure its conservation and the pharmaceutical demand. The present study was undertaken with the objective to improve the protocol for high frequency shoot multiplication and callus mediated somatic embryogenesis in P. kurroa under in vitro conditions.

2 Materials and Methods

2.1 Plant material and surface sterilization

The explants were excised from young and healthy growing parts of P. kurroa collected from high altitudes of Kishwar area of Jammu and Kashmir. The plant parts were washed under running tap water so as to remove all the soil completely. Different explants like auxiliary buds and internodal segments were excised from plant parts and were then collected in a beaker containing 2-3 drops of Tween 20 (Himedia) in distilled water. After 10 min the explants were washed thrice with distilled water.

The explants were then surface sterilized with 70% ethanol, mercuric chloride (HgCl₂), sodium hypochlorite and 1ppm Potassium permanganate (KMnO₄) in laminar air flow. A single or combinations of two or more sterilants were used for the surface sterilization of the explants.

2.2 Culture medium and conditions

MS (Murashige & Skoog 1962) basal medium (Himedia) was used for the shoot establishment. The medium was supplemented with 3% (w/v) sucrose and 0.8% (w/v) agar. The pH of the medium was adjusted to 5.8 before autoclaving. MS medium with varying concentrations of growth hormones was prepared for shoot induction, multiplication and callus proliferation. Cultures were maintained at 25 ±2°C in 16/8 h light/dark cycle in tissue culture room. Subculturing was carried after 4–6 weeks interval. Care was taken to prevent any further contamination of subcultures from the microbes.

2.3 Shoot proliferation and rooting

Two weeks after the initial establishment of aseptic cultures, the shoot cultures thus obtained were further transferred for multiplication on to MS medium containing different combination and concentrations of BAP (Benzyl Amino Purine), Kn (Kinetin) and GA3 (Gibberellic Acid). Ten different MS medium combinations with growth regulators were tried for shoot multiplication (Table 1). For root induction, after 6 weeks the well developed in vitro grown shoots were separated singly and implanted on MS medium supplemented with different concentrations of IBA (Indole Butyric Acid) (Table 2).

2.4 Callus induction and organogenesis

Fully expanded young leaves of in vitro grown plants of P. kurroa were cut into small pieces using sterilized blade and then inoculated on MS medium containing 2,4 D (2, 4-Dichlorophenoxy Acetic Acid) at different concentrations (1.0mg/l to 4.0mg/l) for callus induction and proliferation. The proliferated callus after six weeks was transferred to regeneration media containing BAP, IAA and NAA (Naphthalene Acetic Acid).

2.5 Somatic embryogenesis and plant regeneration

After eight weeks of culturing, the small pieces of friable and embryogenic callus were transferred to MS medium containing 2,4-D (2.0, 2.5, 3.0 mg/l) for induction of somatic embryos. After 4 weeks of subculturing, greenish yellow embryos were observed. These pro embryoids were subcultured on same medium for maturation for next two weeks. Mature somatic embryos were transferred to regeneration MS medium containing BAP (0.5, 1.0, 1.5, 2.0 mg/l) and GA3 (1.0mg/l).
Table 1 Different concentrations and combinations of growth hormones used for shoot and root multiplication in *P. kurroa*.

| Growth regulators                  | Concentration of growth regulators (mg/l) |
|-----------------------------------|------------------------------------------|
| MS + BAP                          | 0.5, 1.0                                  |
| MS + Kinetin                      | 0.5, 1.0                                  |

**Shoot multiplication medium**

| Treatment | Concentration |
|-----------|---------------|
| I. MS + BAP + Kinetin             | 1.0 mg/l BAP + 0.5 mg/l Kn               |
| II. MS + BAP + Kinetin            | 0.5 mg/l BAP + 1.0 mg/l Kn               |
| III. MS + BAP + Kinetin + GA3     | 1.0 mg/l BAP + 1.0 mg/l Kn + 1.0 mg/l GA3|

**Root multiplication medium**

| Treatment                  | Concentration |
|----------------------------|---------------|
| MS + IBA                  | 0.5, 1.0, 1.5, 2.0, 2.5, 3.0 |

3 Results and Discussion

3.1 Surface Sterilization

The explants used in the present study were taken from the *P. kurroa* plant parts, collected from field. These were surface sterilized with sodium hypochlorite used at the concentration of 5.0% which resulted in 46.6% of uncontaminated explants. Best sterilization treatment was 70% ethanol for 30 sec followed by HgCl₂ (0.1%) for 2 min and then KMnO₄ (1ppm) for 3 min resulting in 80.0% of uncontaminated explants. Similar results were also reported by Sen et al. (2014) where they observed that more than 3 min of surface sterilization with mercuric chloride is lethal to all of his tested varieties of *A. aspera*. Thus the herbaceous explants are always treated with low concentration of sterilants so that their soft tissues are not affected or damaged.

3.2 Shoot multiplication

The explants were established in MS basal medium after two weeks of inoculation (Figure 1a). The established shoots were transferred to shoot multiplication medium after four weeks. The best shoot multiplication was observed in MS medium supplemented with 1.0 mg/l BAP, 0.5 mg/l Kn and GA₃ (1.0 mg/l). After six weeks of subculturing in shoot multiplication medium the plants regenerated into multiple shoots (Figure 1b). After six weeks the number of shoots per explant was counted and also their length was recorded.

The maximum percentage of shoot growth observed after six weeks was 85.63% with an average number of 8.7± 0.18 shoots per explant. The average shoot length was 5.92 ± 0.29 cm. However, when the concentration of kinetin was increased to 1.0 mg/l, the percentage shoot growth decreased to 80.21% and the average number of shoots was 7.8± 0.15. (Table 3). Venkatachalam et al. (2012) reported a high frequency (87.3%) of shoot proliferation (6.2 shoots/explant) in MS medium supplemented with 2.0 mg/l BAP in combination with 0.5 mg/l kinetin in *Gloriosa Superba* while Patial et al. (2012) observed maximum number of shoots on MS medium supplemented with kinetin in *P. kurroa*.

In the present study, it was also observed that the shoots obtained on MS medium supplemented with BAP only, were healthy with long and thick leaves but the number of shoots was less whereas the number of shoots was more in MS media containing kinetin but the shoots were thin (Figure 2). Reduction in the number of shoots regenerated per explant with BAP as only growth regulator was also reported in *A. squamosa* (Nagori & Purohit, 2004).

Table 2 Effect of different sterilants on the surface sterilization of explants.

| S. No. | Treatments             | Time duration | Percent survival of the explants |
|--------|------------------------|---------------|---------------------------------|
| 1      | 70% ethanol            | 30-60 sec.    | Nil                             |
| 2      | KMnO₄ (1ppm)           | 3 min         | Nil                             |
| 3      | 70% ethanol + HgCl₂ (0.05%) | 30 sec. + 2 min | 13.3                           |
| 4      | 70% ethanol + HgCl₂ (0.1%) | 30 sec. + 2 min | 36.6                           |
| 5      | 70% ethanol + HgCl₂ (0.2%) | 30 sec. + 4 min | 30.0                           |
| 6      | 70% ethanol + HgCl₂ (0.5%) | 30 sec. + 2 min | 15.9                           |
| 7      | Sodium hypochlorite    | 5 min         | 46.6                            |
| 8      | Sodium hypochlorite    | 5 min         | 20.0                            |
| 9      | 70% ethanol + HgCl₂ (0.2%) + KMnO₄ (1ppm) | 30 sec. + 2 min + 3 min | 66.6                           |
| 10     | 70% ethanol + HgCl₂ (0.1%) + KMnO₄ (1ppm) | 30 sec. + 2 min + 3 min | 80.0                           |
Figure 1 (A) Proliferation of leaves from auxiliary buds after two weeks MS medium  (B) Shoot multiplication in MS medium with BAP, Kn and GA$_3$.

Figure 2 Effect of BAP and Kinetin in *P. kurroa* after 4 weeks of subculturing.

Figure 3 Root multiplication in MS medium with IBA after six weeks of sub culturing.
Table 3 Effect of different concentrations of plant growth regulators on in vitro shoot multiplication in *P. kurroa*.

| S.No | Concentration of growth regulators (mg/l) | Percentage Shoot growth | Average no. of shoots per explant | Average shoot length (cms) |
|------|------------------------------------------|--------------------------|-----------------------------------|---------------------------|
| 1    | MS + 0.5 BAP                             | 52.80                    | 2.2± 0.54                        | 2.17 ± 0.58               |
| 2    | MS + 1.0 BAP                             | 66.20                    | 2.7± 0.38                        | 3.61 ± 0.22               |
| 3    | MS + 0.5 Kn                              | 50.80                    | 1.1± 0.32                        | 1.21± 0.37                |
| 4    | MS + 1.0 Kn                              | 59.06                    | 1.9± 0.51                        | 1.92± 0.52                |
| 5    | MS + 1.0 BAP + 0.5 Kn                    | 68.60                    | 6.8± 0.21                        | 4.22± 0.36                |
| 6    | MS + 0.5 BAP + 1.0 Kn                    | 63.41                    | 5.7± 0.25                        | 5.64± 0.80                |
| 7    | MS + 1.0 BAP + 1.0 Kn                    | 72.49                    | 4.2± 0.51                        | 5.21± 0.71                |
| 8    | MS + 1.0 BAP + 0.5 Kn + 1.0 GA3          | 85.63                    | 8.7± 0.18                        | 5.92± 0.29                |
| 9    | MS + 0.5 BAP + 1.0 Kn + 1.0 GA3          | 76.54                    | 8.2± 0.22                        | 5.53± 0.40                |
| 10   | MS + 1.0 BAP + 1.0 Kn + 1.0 GA3          | 80.21                    | 7.8± 0.15                        | 5.22± 0.69                |

*Data is from twenty shoots in three replicates and is represented as mean±SD

3.3 In vitro rooting and hardening

The addition of auxin in the rooting medium results in long length and higher number of roots as compared to hormone free medium. Development of healthy and long rootlets is helpful for the establishment of plantlets in the field. Direct, healthy root initiation and elongation was observed in *P. kurroa* after subculturing of elongated single shoots onto MS basal medium supplemented with different concentrations of IBA (Table 4; Figure 3).

Best rooting medium was MS medium supplemented with 2.5 mg/l IBA resulting in 84.54% of root growth. The average number of roots per explant was 10.35±0.18. The roots were long, healthy with numerous root hairs. As the concentration of IBA was increased to 3.0mg/l, the percent root growth and the average number of roots was decreased to 79.22% and 9.11±0.21 respectively (Table 4). Similar effects of IBA were also observed during in vitro rooting in several other medicinal plant species (Baul et al., 2011; Praveena & Veeresham 2014; Rahdari et al., 2014). The rooted plants were removed from flasks, washed carefully to remove agar and transferred to a potting mixture in plastic pots containing sand: soil (1:1). The plants were initially covered with jars in glass house for two weeks to maintain the humidity. About ninety percent rate of survival was obtained upon hardening.

3.4 Callus induction

Different concentrations of 2,4-D in MS medium resulted in callus formation from the leaf explants (Table 5). The callus was creamy white and friable in MS medium containing 2.0mg/l and 3.0mg/l 2,4-D (Figure 4). It was observed that the high concentration of 2,4-D (4.0mg/l) resulted in browning of explants with no callus formation. Similarly, Zuraida et al., (2015) has also reported that higher concentration of 2,4-D resulted in browning of the callus in *Pelargonium radula*.

Table 4 Effect of IBA concentration on in vitro rooting.

| S. No | MS + Growth regulators (mg/l) | Percentage rooting (%) | Average no. of roots per explant | Types of roots |
|-------|-------------------------------|------------------------|----------------------------------|----------------|
| 1     | MS +0.5 IBA                   | 35.40                  | 1.23±0.27                        | Small thread like roots no root hairs. |
| 2     | MS +1.0 IBA                   | 55.80                  | 4.97±0.62                        | Thin, fragile roots with less root hair. |
| 3     | MS +1.5 IBA                   | 68.56                  | 7.14±0.34                        | Thin, fragile roots with root hairs. |
| 4     | MS +2.0 IBA                   | 74.58                  | 8.44±0.23                        | Thin, fragile roots with few root hairs |
| 5     | MS +2.5 IBA                   | 84.54                  | 10.35±0.18                       | Long, well developed roots with numerous root hairs. |
| 6     | MS +3.0 IBA                   | 79.22                  | 9.11±0.21                        | Long, well developed roots with numerous root hairs. |

*Data is from fifteen plants in three replicates and is represented as mean±SD
After six weeks, MS medium supplemented with 2.5mg/l 2,4-D + 1.0mg/l BAP + 1.0mg/l IAA resulted in the formation of shoots along with the roots from the callus. When the proliferated callus was inoculated on MS medium containing 2.5mg/l 2,4-D + 2.0mg/l NAA + 0.5mg/l IAA, then only root formation was observed (Figure 5a and 5b). The concentration of plant growth regulators needed to induce callus, varies from species to species and also depends on the source of explant (Ngomuo et al., 2014). It has been reported in many plants that 2,4-D is mostly used as auxin for callus induction (Junaid et al., 2007; Silveira et al., 2013) and addition of a low concentration of cytokinins in callus culture medium often enhances callus regeneration (Singh et al., 2009).

3.5 Somatic embryogenesis and shoot regeneration

Somatic embryogenesis has already been reported in many endangered plants for their in vitro conservation (Lee et al., 2011; Kim et al., 2012; Giri et al., 2013; Barberini et al., 2016). In the present study, cytokinins were used as promoters for the induction of somatic embryos. The combination of both 2,4-D along with BAP and GA3 in MS medium significantly affected the number of embryo production. After four weeks, 2,4-D at the concentration of 2.5mg/l resulted in the formation of greenish yellow callus.

The callus when subcultured on the same medium resulted in the formation of many globular somatic embryos. Further subculturing resulted in the formation of globular, torpedo and heart shaped somatic embryos. Mature somatic embryos were transferred to regeneration medium containing BAP and GA3. The mature embryos turned green on MS medium supplemented with 1.0mg/l BAP and 1.0mg/l GA3 after a week and then after two weeks the shoots differentiated with one or two plumules was observed (Figure 6).
Figure 6 Shoot regeneration from somatic embryos in MS medium supplemented with 2,4-D, BAP and GA3

Table 5 Effect of plant growth regulators on callus induction and regeneration after 6 weeks.

| S. No | MS+Growth hormones (mg/l) | Type of Response in P. kurroa | Rate of callus induction |
|-------|---------------------------|------------------------------|--------------------------|
| 1     | MS+1.0 2,4-D              | No callus                    | ---                      |
| 2     | MS+2.0 2,4-D              | Creamy white, friable, small in size | +               |
| 3     | MS+3.0 2,4-D              | Creamy white, friable, large in size | ++              |
| 4     | MS+4.0 2,4-D              | Browning of the explant      | ---                      |
| 5     | MS+2.5 2,4-D+ 1.0 BAP     | Creamy white, friable callus, with shoots | +++             |
| 6     | MS+2.5 2,4-D+ 1.0 BAP + 1.0 IAA | Creamy white, callus, with small shoots and roots | +++          |
| 7     | MS+2.5 2,4-D+2.0 NAA +0.5 IAA | Creamy white callus, with large roots | +++          |

---- Absent, + Less callus, ++ moderate growth, +++ Good growth rate

Conclusions

The in vitro techniques help in conservation and multiplication of medicinally important plants that are overexploited for their bioactive constituents. The present study has developed a protocol for mass multiplication and somatic embryogenesis in P. kurroa which will facilitate its rapid and large scale propagation. This study can facilitate the in vitro conservation of elite genotypes of this medicinally important endangered plant. Further the somatic embryogenesis opens a chance for the improvement of the plant through transgenic approach. Though further studies needs to be done for the improving the germination of somatic embryos and their further establishment in the green house.

Conflicts of interest

Authors would hereby like to declare that there is no conflict of interests that could possibly arise.

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