Original article

Chromatographic quantification of polyphenol in relation to potential antioxidant activity and isolation of DNA from in vitro cultivated Decalepis nervosa Wight & Arn. leaf explant

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Abstract

The objective of the present study was to perform in vitro micropropagation as well as the growth of callus from the leaf explant of Decalepis nervosa Wight & Arn. (DN) (Family: Apocynaceae), a climber woody medicinal plant. Addition of silicon (Si) as sodium and potassium silicate in the MS media helped in fast micropropagation, callus development and organogenesis of the explant which was modified technique in plant tissue culture of the said plant. Silicon ion also helped in avoiding browning of callus. Therefore, an alternate in vitro tissue culture method was established using sodium silicate and potassium silicate using varying concentrations of 0.5, 1.0, 2.0, 2.5, 3.0 and 3.5 mg/l to overcome the challenges that compared with the normal MS media. Half strength MS medium supplemented with BAP at 0.2 mg/l and NAA at 2.5 mg/l gave callus growth in 21 days and resulted in more accumulation of gallic acid. Thereafter, direct rooting and shoots of meristem occurred with IBA and Si as sodium silicate (IBA at 2 mg/l and Na₂SiO₃ at 2.5 mg/l), supplemented in half-strength MS media along with coconut water, within 18 days. Further, gallic acid in the methanol callus extract was estimated by the HPLC method and analyzed for antioxidant activity, using DPPH assay method. Finally, leaf DNA extraction was carried out (starting of genotype analysis) by CTAB (cetyl trimethylammonium bromide) method for purification of DNA and nanodrop method for quantification of the same. Finally, the results concluded that potassium silicate enhanced the production of phenolic content when analyzed by HPLC method, proved antioxidant activity as well as improved root formation in a very short time. Thereafter, single DNA pure band was identified and quantified which is essential for future research on new drug discovery.

Key words: Antioxidant, callus, chromatography, DNA, in vitro culture, silicates

1. Introduction

In biological research, plant tissue culture is an interesting, important and also challengeable experiment for the noble drug discovery. The technique has a high impact on increased plant secondary metabolites, easy isolation of constituents and genetic manipulation as per desired requirements. The plant tissue culture method not only helps in the enhancement of phytoconstituents but also helps in mass production of economically important plant species that may be any known cultivated plants or may be endangered plants. There are many forms of tissue culture techniques, in that callus culture and micropropagation are the well-known methods of interest for mass production (Dave and Purohit, 2002; Debnath et al., 2006).

Of late, Decalepis nervosa (DN) Wight & Arn. Venter is among the species of Decalepis which became endangered plant species with medicinal liana of Apocynaceae. Though, the plant is distributed in the Western Ghats regions of Tamil Nadu, Kerala, and Karnataka due to the over-exploitation and habitat loss, the plant is in high demand to the economic Indian market (Jonta, 2009; Udayan et al., 2013). The plant is climbing shrub, leaves are the simple opposite, with elliptic or elliptic-oblong and acute to acuminate shape. The roots of this plant are odorous, curvy and hard. Other species of Decalepis roots are aromatic odor and traditionally used in the preparation of pickles, flavoring principle (Murthi and Seshadri 1941) and several Ayurvedic drug formulations (Anonymous, 2003), appetizer and blood purifier (Jacob, 1937). It was documented that tribal people use more than 9500 plant species for their various requirements among that for the medicinal purpose, they use more than 7500 species and remaining for food, fiber, and cordage, fodder, gum, resin and dyes (Anonymous, 1962; Anonymous, 2000). Hence, the demand for the plant is enormous and this leads to abundant procurement of the plant parts which created the existence of the same species. The overexploitation by destructive harvesting, habitat loss, limited pollinator, delayed seed germination, etc., resulted, the said plant become endangered in the natural habitat but still some species of the plant exists in the wild forest, especially in the Western Ghats regions of the South Indian zones. Hence, conservation of such a selected plant is essential for economic benefits. In the present study, a micropropagation protocol was...
developed for the said plant for mass propagation with high biomass yield. Many works of literatures revealed various micropropagation techniques for Decalepis hamiltonii and other Decalepis species (Bais et al., 2000; Giridhar et al., 2005), but no such literature is available on DN leaf in vitro culture. Thereafter, phenolic acid such as gallic acid is an important plant secondary metabolites which is present in the leaf of DN plant, shows many pharmacological activities such as an antioxidant, antimutagenic, lipid-lowering, antiatherosclerotic, anti-liver injury, anti-tumor, and anticarcinogenic, etc. (Jiang and Yang, 2010; Zhang et al., 2010).

Hence, it is necessary to enhance the content of gallic acid in the DN leaf through in vitro culture. Application of silicon in plant tissue culture helps in early root formation, micropropagation, and accumulation of plant secondary metabolites. Many research works are proved earlier that the use of silicon compounds especially sodium, potassium, and calcium silicate improves organogenesis, embryogenesis, growth traits, morphological, and physiological characteristics of leaves and prevents oxidative phenolic browning in various plants (Sivanesan and Park, 2014). There are no reports on the in vitro culture of DN plant species. A polyphenolic compound such as gallic acid is present in the selected plant, and also silicate compounds help in accumulation of secondary plant constituents. Hence, an innovative protocol was designed and the present research was carried out with silicon-containing compounds with a combination of coconut water and other plant growth hormones for micropropagation and growth of the plant from stem under the aseptic condition as well as callus induction of the leaf disk. Further, estimation of a polyphenolic compounds such as gallic acid with its antioxidant study vis-a-vis DNA isolation, purification and quantification from plant callus were carried out.

2. Materials and Methods

2.1 Procurement of DN plant

 Matured DN plants were procured from Western Ghats region of Tamil Nadu in month of April, 2018 by Dr. S. Gokul, PDF scholar of CIMP, Bangalore and maintained in the department (Pharmacognosy and Phytochemistry, Krupanidhi College of Pharmacy, Bangalore, Herbarium number: KCP/KD-DN/398/2018) medicinal plant garden after authenticated by Dr. Rajasekharan P. E, Principal Scientist, Department of Plant Biotechnology, Indian Institute of Horticultural Research, Bangalore. Chemicals and plant hormones used in this present study were procured from Himedia Laboratories, Mumbai, India.

2.2 Preparation of culture medium

Murashige and Skoog (MS) basal medium (Murashige and Skoog, 1962) was used throughout the study. The basal medium was supplemented with various plant hormones with varying concentrations such as 1-naphthaleneacetic acid (NAA), indole-3-butyric acid (IBA) and 6-benzylaminopurine (BAP) (0.1 to 3 mg/l), 1 to 3% (w/v) sucrose, media adjuvants such as coconut water (3 to 8%, v/v) and silicate compounds, viz., sodium and potassium silicate (0.5 to 3.5 mg/l). The pH of the medium was adjusted to 5.6 to 5.8. The medium was prepared with added 0.8 to 1.2% (w/v) agar in 1 Lt volume and then melted in a gas burner. The whole medium separately was distributed in various culture vessels, tubes and finally then sterilized in an autoclave at 15 Psi and 121°C, for 15 min.

2.3 Surface sterilization of explants

Fresh young leaves of DN were selected, thoroughly washed in running tap water and 0.3% (v/v) sodium hypochlorite with a few drops of "Teepol" for 5 min. Then rinsed with double distilled water followed by a wash in running tap water. Then leaves were transferred to sterile laminar airflow cabinet and surface sterilized in 0.05% (w/v) mercuric chloride (HgCl2) solution for 5 min and rinsed thoroughly in sterile distilled water.

2.4 Callus establishment and growth

Surface sterilized tender leaves were cut into optimum size (0.5 to 1.0 cm). Few leaf discs were aseptically placed into sterile culture vessels and tubes under aseptic area and incubated under the BOD incubator at 25 ± 2°C. The night environment was maintained properly throughout the callus growth. Proliferated callus was sub-cultured in 10 to 15 days interval for further establishment. After establishment, 0.5 g of callus was transferred to fresh MS medium (50 ml) supplemented with BAP and NAA (0.1 to 3 mg/l), coconut milk (3 to 8%, v/v).

2.5 Micropropagation

Meristem contained nodal segments (1 cm size) of DN plant was selected for micropropagation. Separately, the meristems are inoculated aseptically in test tubes (25 × 150 mm) containing 20 ml of MS medium with the respective sources and concentrations of silicate, then the vessels were maintained in a growth chamber at 25 ± 2°C, an average irradiance of 51 µmol/m²/s and a darkness of 17 h. The number of leaves, shoot length (cm) and number of roots were measured in each case. Furthermore, Silicon concentration in leaves was identified with a scanning electron microscope and determined the content of Si in high yielded micropropagated plant that compared with low yielded micropropagated plant. For the method, the leaves were collected separately and oven-dried at 60°C temperature. Dried leaves samples were coarsely powdered and silicon content was estimated by the molybdenum-blue colorimetric method, described earlier (Gallo and Furlani, 1978).

2.6 Extraction of callus

5 g of fresh callus was reflux with 50 ml methanol for 6 hrs. Evaporated the methanol solvent and the viscous extract was collected for further experiment.

2.7 HPLC analysis

The amount of Gallic acid present in the callus extract was estimated by compared with normal plant hormone content MS medium and MS medium supplemented with silicon compound and plant hormone in HPLC.

**Optimized conditions for chromatography**

| Column | Phenomenex Gemini-NX-5 µm C18, 110 Å, LC Column 250 x 4.6 mm |
| Mobile phase | Methanol: water (60:40) |
| Injection valve | 7725i Rheodyne 20 µl, USA |
| Elution Type | Isocratic |
| Flow Rate | 1 ml/min |
| Detection | UV at 203 nm |
**Standard preparation**

5 mg of standard Gallic acid (purity: 99.5%) was taken in 5 ml volumetric flask and make up the volume to 10 ml with methanol (the concentration of this solution is 1 mg/ml) and was sonicated (at frequency of 30 kHz) for 8 min then the solution was filtered using 0.45 micron Millipore filters.

**Sample preparation**

1 mg of the callus extract was dissolved in 1 ml of methanol (Conc: 1 mg/ml). The solution was a vortex for 5 min. The sample was filtered using 0.45-micron Millipore filters. 20 µl of this sample was injected into the HPLC system.

**Calculation**

The percentage concentration:

\[
\text{Percentage concentration} = \left( \frac{\text{Sample area}}{\text{Standard area}} \right) \times \left( \frac{\text{Standard wt}}{\text{Sample wt}} \right) \times \left( \frac{\text{Sample dilution}}{\text{Standard dilution}} \right) \times \left( \frac{\text{X purity of standard}}{100} \right)
\]

**2.8 Antioxidant study**

**2.8.1 DPPH (2,2-diphenyl-1-picrylhydrazyl hydroxyl) method**

Callus methanol extract was estimated antioxidant activity by the DPPH method. This method is very basic and easy method for estimation of antioxidant nature of extract by spectrophotometric method at 517 nm. The method followed as mentioned in earlier literature (Das et al., 2019) i.e., DPPH solution mixed with extract and kept for 30 minutes in dark for the reaction and finally absorbance measured against blank methanol. Thereafter, percentage inhibition of DPPH was calculated as:

\[
\% \text{inhibition} = \left( \frac{\text{Abs}_{\text{blank}} - \text{Abs}_{\text{sample}}}{\text{Abs}_{\text{blank}}} \right) \times 100
\]

where, Abs = Absorbance at UV 517 nm

**2.9 Isolation of DNA from DN callus by CTAB**

The callus tissue is grinded in liquid nitrogen contained a few mg of polyvinylpyrrolidone. Further, 700 µl of CTAB is added in extraction buffer and 10 µl of β-mercaptoethanol for 0.5 gm of tissue. Then keep at 60°C for 30-45 min and then an equal volume of chloroform: isooamylalcohol (24:1 ice cold) is added and mixed well. Thereafter, centrifuged at 8000 rpm for 20 min and then 4 µl of RNase is added into that solution and incubated on a water bath at 65°C for 15 min or 37°C for 1 h in the incubator. Then again the equal volume of chloroform: isooamylalcohol is added and centrifuged at 8000 rpm for 10 min. To the 2/3rd volume of ice-cold isopropanol or double volume of ice-cold ethanol is added to the solution and kept at-70°C. Further, centrifuged at 8000 rpm for 15 min and precipitate is collected. Washed the pellet with 70% alcohol and air-dried, the pellet and resuspended the pellet in 10 mM tris buffer and kept at 4°C for dissolving DNA. Finally, purified DNA was estimated (absorption measured) by nanodrop method by placed 2 µl of water as selected "Blank" after that placed another 2 µl of water to confirm that the measure is 0 and then placed 2 µl of the sample. The ratio of the absorbance at 260: 280 nm and 260: 230 nm (A_{260} / A_{280}, A_{260} / A_{230}) are used to assess the purity of Isolated DNA (Doyle and Doyle, 1987; Healey et al., 2014).

**2.10 Correlation study**

Yield, Si content in the leaf of DN was correlated with the estimated amount of the gallic acid present in the callus methanol extract.

**3. Statistical analysis**

Analysis of variance (ANOVA) and Duncan's multiple range test (Duncan, 1955) were performed. mean ± SD was calculated for the number of roots and length of the shoot. The differences among averages of the recorded parameters for all treatments were tested for significance at 5% level (p<0.05).

**4. Results**

**4.1 Callus initiation and elongation**

Various concentrations of plant hormones with sodium and potassium silicate supplemented MS medium were prepared and leaf disk of DN explant was established. It was resulted that potassium silicate (2.5 mg/l) and coconut water (3 ml/l) containing MS medium supplemented with BAP at 0.2 mg/l and NAA at 2.5 mg/l gave satisfactory initiation of callus in 21 days (Figure 1), whereas in same concentration, sodium silicate and coconut water (2.5 ml/l) containing MS medium supplemented with BAP at 0.3 mg/l and NAA at 2.0 mg/l gave maximum callus growth in 35 days of subcultured callus and thereafter, the growth decreased with increase in concentration. The result showed the highest callus growth with a fresh weight of 1.68 g with sodium silicate at 2.5 mg/l concentration in MS medium whereas the same was reduced (1.23 g) with potassium silicate at 2.5 mg/l concentration (Figure 2).

**4.2 Micropropagation and silicon content in leaf**

Various concentrations of silicate compounds such as sodium and potassium silicates were mixed in MS medium (Full and Half strength) supplemented with IBA and NAA (Auxin hormone for root growth) and coconut water 5% v/v (Cytokinin) were used for direct micropropagation from the DN meristem. It was observed that IBA at 2 mg/l and Na₂SiO₃ at 2.5 mg/l supplemented in half-strength MS media along with coconut water initiated roots and shoot within 18 days and improved the length of roots and shoots after 48 days of subculture. The number of leaves, shoot length (cm) and number of roots was measured and resulted height number of leaves came 16, with a length of shoot 7 cm and 9 numbers of roots with the same condition (Figure 3, Table 1).

The number of roots and length of the shoots is increased up to a certain concentration increase of the sodium and potassium silicate in the MS medium but the same is more significant in sodium silicate containing MS medium (Table 1). Interestingly, the number of leaves is decreased with an increased in concentration (Figure 4) for both the cases (IBA at 2 mg/l and K₂SiO₃ at 2.5 mg/l with coconut water and IBA at 2 mg/l and Na₂SiO₃ at 2.5 mg/l). Thereafter, the accumulation of silicon content in leaves was higher in sodium silicate containing MS medium up to a certain concentration (at 2.5 mg/l) of plant growth than potassium silicate (Figure 5).
4.3 Extraction of callus
Methanol solvent was used for extraction of total phenolic and resulted higher yield with callus formed in sodium silicate contained MS medium (Figure 6).

4.4 HPLC estimation
Quantification of gallic acid present in the extract of various callus was determined by HPLC analysis. Results revealed showed more content of gallic acid (0.68 mg/100 g) in sodium silicate contained callus which was higher than any other extract (Figures 7 and 8). The retention time (Rt) of gallic acid present in the extract of callus showed the same Rt as per standards gallic acid (3.10 min) (Table 2).

The stock solution of the standard (1 mg/ml) was diluted to five different concentrations (5, 10, 20, 40 and 80 µg/ml) and the calibration curve was obtained by plotting peak area versus concentration of the sample. The correlation coefficient of the linearity curve was measured at $R^2 > 0.997$. The percentage of relative standard deviation (RSD) for standard gallic acid was < 1.

4.5 Antioxidant study of callus extracts
All three different callus were estimated antioxidant activity by DPPH method and result revealed sodium silicate contained MS medium gave higher antioxidant value than normal and potassium silicate contained MS medium. IC$_{50}$ was calculated with compared with the standard ascorbic acid and result was depicted in Table 3.
Table 1: Number of roots, length of shoot in various concentrations of hormone in MS medium

| Condition       | Concentration used | Plant hormone | Concentration | Length of shoots (cm) | No. of roots |
|-----------------|--------------------|---------------|---------------|-----------------------|--------------|
| Sodium silicate | 2.5 mg/l           | IBA           | 0.5           | --                    | --           |
|                 |                    |               | 1.0           | --                    | --           |
|                 |                    |               | 1.5           | 03 ± 0.10             | 02 ± 0.01    |
|                 |                    |               | 2.0           | 07 ± 0.11*            | 09 ± 0.13**  |
|                 |                    |               | 2.5           | 07 ± 0.21*            | 07 ± 0.11    |
|                 |                    |               | 3.0           | 07 ± 0.02             | 05 ± 0.32    |
|                 |                    |               | 3.5           | 07 ± 0.13             | 05 ± 0.01    |
| Potassium silicate | 2.5 mg/l       | IBA           | 0.5           | --                    | --           |
|                 |                    |               | 1.0           | --                    | --           |
|                 |                    |               | 1.5           | 02 ± 0.32             | 04 ± 0.03    |
|                 |                    |               | 2.0           | 05 ± 0.12             | 05 ± 0.12    |
|                 |                    |               | 2.5           | 05 ± 0.21             | 04 ± 0.10    |
|                 |                    |               | 3.0           | 05 ± 0.02             | 04 ± 0.13    |
|                 |                    |               | 3.5           | 05 ± 0.11             | 03 ± 0.02    |

Mean ± SD (n = 3); Other concentration of hormones are not shown in this table. **p<0.01; *p<0.05; (Other concentrations of silicate compounds are not showed in table).

Figure 3: Micropropagation of DN meristem in half strength MS medium (IBA at 2 mg/l and K₂SiO₃ at 2.5 mg/l and coconut water 5% v/v).

Figure 4: Number of leaves of DN in various concentration of sodium silicate.

Figure 5: Concentration of silicon in in vitro cultivated leaves of DN by Sod. Silicate supplemented MS medium.
**Figure 6:** Yield and percentage yield in various callus.

**Figure 7:** Standard gallic acid.

**Figure 8:** Gallic acid in sod. silicate contained callus methanol extract.

**Table 2:** Content of gallic acid in various methanol extract of callus

| Type of methanol extract | Gallic acid (mg/100g) | R²   | % RSD | Rt (Minute) |
|-------------------------|-----------------------|------|-------|-------------|
| Normal callus extract   | 0.30                  | 0.996| 0.825 | 3.11        |
| Sod. Silicate contained callus extract | 0.78                  | 0.997| 0.808 | 3.10        |
| Pot. Silicate contained callus extract | 0.57                  | 0.997| 0.812 | 3.12        |

**Table 3:** IC₅₀ values of DPPH scavenging effect in various callus methanol extracts

| Extracts                    | IC₅₀ value (µg/ml) |
|-----------------------------|-------------------|
| Ascorbic acid (standard)    | 7.64 ± 0.32       |
| Normal callus               | 17.23 ± 0.02***   |
| Sod. Silicate contained callus | 11.20 ± 0.16***   |
| Pot. Silicate contained callus | 13.07 ± 0.32***   |

Mean ± SD; (n=3), one way ANOVA study, all data compared with standard ascorbic acid; *** p<0.001= Very high significant.

**4.6 Isolation of DNA from DN callus**

Isolation of DNA was carried out with the callus of sodium silicate contained MS medium. Result showed single isolated DNA with high purity (Figure 9) and further nano drop method was used for quantification and the result showed in Table 3.

**Table 3:** IC₅₀ values of DPPH scavenging effect in various callus methanol extracts

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|-----------------------------|-------------------|
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| Sod. Silicate contained callus | 11.20 ± 0.16***   |
| Pot. Silicate contained callus | 13.07 ± 0.32***   |

Mean ± SD; (n=3), one way ANOVA study, all data compared with standard ascorbic acid; *** p<0.001= Very high significant.

**4.7 Correlation study**

Based on the previous result, further yield of methanol extract of callus, gallic acid content and silicon content in sodium silicate contained micropropagated leaves was correlated and resulted significant correlation among them (Table 4). The study indicated that yield showed more significant correlation with silicon content in leaves (p<0.05) than content of gallic acid.
Gallic acid content in sod. (Zhao et al., 2001). 2, 4-D and Kinetin combinations were not suitable for cell growth and gallic acid synthesis (Li et al., 2008) because they produced a fast change in the biomass color (from light yellow to light brown) and hence this combination was not applied in this study. Furthermore, it was reported that callus growth was restricted with an increased in phenolic content and the growth occurred in dark conditions (Hoque and Arima, 2002; Castro et al., 2016). A similar result obtained in the present investigation.

Table 4: Correlation study between yield, gallic acid content and Si content in DN plant

| Yield of extract of sod. | Gallic acid content in sod. | Si content in sod. |
|-------------------------|----------------------------|-------------------|
| Silicate contained callus | Silicate contained callus | Silicate contained micropropagated leaf |
| Yield of extract of sod. Silicate contained callus | 1 | 1 |
| Gallic acid content in sod. Silicate contained callus | 0.974* | 0.961* |
| Si content in sod. Silicate contained callus | 0.987** | 1 |

*p<0.05= Significant; **p<0.01 = High significant.

5. Discussion

In the present study, silicate compounds (sodium and potassium silicates) were used for the growth of callus of DN leaf disk. It was reported that silicate compounds increase the morphogenetic potential of plant cells, tissues, and organs in an aseptic culture medium. Not only this, Si also significantly affected the antioxidant enzyme activities in many plants by enhanced the growth and morphogenesis (Liang et al., 2007) of enzymes. The addition of Si to the shoot induction medium significantly increased SOD, POD, APX, CAT activity in regenerated shoot buds as compared with the control (Iyyakkannu and Jeong, 2014). Earlier scientific research reports evident for essential function of Si to the tissue culture medium that enhances callus growth, shoot regeneration and root induction, stimulates somatic embryogenesis, and improves morphological, anatomical and physiological characteristics of plantlets, prolongs the longevity of callus and organs with a potential for plant regeneration (Fadd and Reda, 2014; Iyyakkannu and Jeong, 2014). Based on these concepts, the present study was carried out the first time by applied silicate compounds (sodium silicate and potassium silicate), the growth of callus from the leaf disk of DN explant. Callus initiation resulted with potassium silicate (2.5 mg/l) contained MS medium, supplemented with BAP (at 0.2 mg/l) and NAA at (2.5 mg/l) in 21 days which was also similar with the earlier results (Azra et al., 1997; Dong and Zhan, 2011). Thereafter, sodium silicate helps in the maintenance of callus and its growth and development in MS medium supplemented with BAP at 0.3 mg/l and NAA at 2.0 mg/l in 35 days of subcultured callus. A similar observation was also reported earlier when Si as sodium silicate was added to the modified MS medium, promotes the growth of callus from stem nodal (Sivanesan and Park, 2014). Thereafter, it was reported that the production of gallic acid increased when NAA increased from 0 to 2 mg/l and the same was reduced when concentration was increased at 2.5 mg/l of NAA (Dong and Zhan, 2011). Hence, media composition with sodium silicate and supplemented with 0.3 mg/l of BAP and 2.0 mg/l of NAA was more appropriate for subculturing and maintenance of callus without browning of callus. The similar result was also reported earlier where a combination of NAA 2 mg/l and BAP 0.2 mg/l was suitable for the best cell growth (19.70 g dry weight/l) and jasenosid production (0.3% dry weight) in Saussurea medusa (Zhao et al., 2001). 2, 4-D and Kinetin combinations were not suitable for cell growth and gallic acid synthesis (Li et al., 2008) because they produced a fast change in the biomass color (from light yellow to light brown) and hence this combination was not applied in this study. Furthermore, it was reported that callus growth was restricted with an increased in phenolic content and the growth occurred in
more efficient in extraction of lower molecular weight polyphenols such as gallic acid (Dai and Mumper, 2010; Do et al., 2014). Quantification of gallic acid was performed by HPLC and revealed the same trend where sodium silicate contained MS medium gave higher estimated amount than other extracts and the result was due to the absorption of phenolics by silicate ion in the callus. Furthermore, the same was also revealed in case of accumulation of gallic acid in the leaves. It was reported that plant hormones NAA and BAP were more proper in the presence of coconut water in the accumulation of gallic acid. In this experiment, added coconut water further helped in the accumulation of gallic acid in the callus which showed synergistic action. The result was similar to the earlier research (Darwesh Hadeer et al., 2015) where coconut milk enhanced the accumulation of phenolics in the callus.

Significant antioxidant activity resulted in all the callus extract (p<0.001) which was due to combined effects of silicate compounds, plant hormones and coconut water, supplemented in the MS medium. Polyphenol compounds are itself acts as an antioxidant and, hence the presence of gallic acid was found more in the callus contained silicate compounds and, hence positive synergistic antioxidant activity was reported with the increased content of gallic acid when extract with methanol. The present result was also correlated with the earlier literature (Sunita et al., 2013; Ibrahim et al., 2018; Sekeroglu and Gezici, 2019).

Isolation of DNA from the callus was carried out and resulted in one single band purified DNA and further quantified at two different wavelengths in UV spectrophotometer. The ratio of absorbance at 260 nm and 280 nm was used to assess the purity of DNA. A ratio of ~1.8 is generally accepted as "pure" for DNA. If, the ratio was appreciably lower in either case, it was indicated that the presence of protein, phenol or other contaminants absorb strongly at or near 280 nm. Thereafter, 260/230 ratio was used as a secondary measure of nucleic acid purity. The 260/230 values for "pure" nucleic acid are often higher than the respective 260/280 values. Expected 260/230 values were commonly in the range of 2.0-2.2. If, the ratio was appreciably lower than expected, it was indicated the presence of contaminants which absorb at 230 nm. The study helped to know about the genomic polymorphism of the said plant. In earlier literature also studied the same to identify DNA from the callus extract of carpobrotus species (Diadema et al., 2003). This study will further help the researchers for the beneficial utilization of DNB in various therapeutic efficacies.

Finally, correlation study revealed a significant result when compared silicon and gallic acid content with the yield and silicon content with gallic acid content. A positive significant correlation was observed that established valid scientific evidence for the aim of the present investigation.

6. Conclusion

The results finally concluded that callus and micropropagation of the DN plant successfully carried out through in vitro cultured method, using sodium and potassium silicate compound and further the activity was boosted with the applied coconut water (3% v/v) in MS medium. Callus initiation was observed with potassium silicate contained MS medium, supplemented with BAP and NAA plant hormone whereas the same was subcultured and callus growth was maintained with sodium silicate contained MS medium without browning of callus. Thereafter, shoot length, the number of roots and fresh callus weight, silicon content in leaves were increased with sodium silicate contained MS medium that also associated with high antioxidant activity. Pure DNA was isolated and quantified at various wavelengths and revealed easy manipulation of gene level for better therapeutic efficacies in the near future.

Conflict of interest

The authors declare that there are no conflicts of interest in the course of conducting the research. All the authors had final decision regarding the manuscript and decision to submit the findings for publication.

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