PRODUCTION OF LIMONOIDS THROUGH CALLUS AND CELL SUSPENSION CULTURES OF CHINABERRY (MELIA AZEDARACH L.)

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Abstract

To study the in vitro production of limonoid contents through callus and cell suspension cultures of chinaberry (Melia azedarach L.) in different explants were inoculated in Murashige and Skoog (MS) medium for the callus induction with various plant growth regulators (PGRs) separately as well as in combinations. The highest callus induction (73.3%) was observed in nodular stem sections and further callus was subcultured for multiplication and finally transferred to cell suspension medium. The optimized parameters for the production of total limonoids were adjusted and UV-visible spectrophotometer was used to quantify total limonoids at 577 nm. Production of total limonoids through callus cell suspension cultures on the optimized medium was highest (141.7 µg/ml) followed by other sources like KH2PO4 (0.1 g/l) extra supplementation with optimized medium produced (138 µg/ml). Plant callus cell suspension cultures through optimized medium may be considered as a good source for the production of bio-products and its purified form could be used as a medicinal sources.

Introduction

The Chinaberry (Melia azedarach L.) belonging to Meliaceae is widely distributed throughout the tropics, subtropics and temperate zones. It is a fleshy-fruited small to medium-size tree native to northwestern India. Chinaberry plant has been introduced to many subtropical countries is also cultivated in different areas of Pakistan up to 1700-meter height which can be used as ornamental shade-tree and also as insect repellent (Hammad et al. 2000 and Nasir et al. 1972). In last 20 years, this family identified as one of the most important sources of bioproducts of genera Melia and Azadirachta, were found very effective (Schmuttnerer and Wilps1995).

Plants growing in natural environment defend themselves against biotic constraints to maintain their survival, and that they synthesize different secondary metabolites having important physiological and ecological functions was reported by Guerriero et al. (2018), Jassim and Naji (2003) and Wallace (2004). M. azedarach is a multipurpose tree because it is used in multidirectional and widespread purposes in medicine, therapeutics as well as for other economic implications (Shekhawat et al. 2014). Secondary metabolites are the main constituents of plants involved in the protection against herbivores, bacteria, viruses, fungi and other competing with plants. Some plants made use of bioproducts as serve to attract pollinators and seed dispersers (Wink 2003, Yang et al. 2018). Plant secondary metabolites are biologically active, non-nutrients and having good role in the maintenance of human life are used as food, drink (Cherif 2012). A large number of these plant-based chemicals are used as important sources for varieties of pharmaceuticals, agrochemicals, food additives, flavors, pesticides, resins, dyes and oils (Balandrin and Kloeke1988, Parr 1988).

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Chinaberry is rich source of secondary compounds including tetrnorotriterpenoids (limonoids) which are important phytochemicals due to their wide variety of properties like antioxidant, antimalarial, antibacterial, antifungal and insecticidal effects against disease-causing vectors (Coria et al. 2008, Khanavi et al. 2008, Mazumder and Rahman 2008, Thirumalai et al. 2009, Abdel-Aty and Abdel-Aty 2015). In some countries, the chinaberry plant is used for the medicinal purpose of treating humans against different diseases such as nausea, vomiting, skin diseases, anthelmintic, diuretic, vermifuge, typhoid fever, and pain in the pelvic region and also for the treatment of asthma (Hammad et al. 2001). Fruit extracts of chinaberry have multiple effects against insects like growth retardation, antifeedant, moulting disorders and changes of behavior (Schmidt et al. 1998, Hammad et al. 2001, Wandscheer et al. 2004).

The amount of plant based medicinal and insecticidal compounds in different parts of chinaberry are low. This might be due to effect of geographical distribution of plants as well as seasonal variations affecting the production of these limonoids from targeted plants. Similarly, that these plants may be susceptible to environmental influences like light, heat, etc. was reported by Sundaram et al. (1995), Szeto and Wan (1996) and Jarvis (1998). The plant products (secondary metabolites) are limited due to above mentioned problems for production. As awareness towards safer, environment friendly plant products are increasing, demand of chinaberry based limonoids is also increasing continuously. Limonoids have complex chemical structures, which are difficult to ways in the laboratory. To overcome these problems, there is urgent need to develop alternative route for limonoids production. The use of biotechnological techniques like plant cell and tissue culture for production of these metabolites is getting importance. That in vitro callus is an alternative source for the production of secondary metabolites was reported by Samar et al. (2013). The advantage of in vitro technique is the continuous production of these metabolites, which is not affected by any seasonal variation or other environmental influences and diseases. Thus the plant cell and tissue culture technique can be used for the enhanced production of these metabolites (Ushiyama and Hibino 1997, Ferri et al. 2011).

Callus cell suspension culture is an efficient means for the production of secondary metabolites with the addition of various plant growth regulators (Zeng et al. 2009). There are also a few reports on the production of azadirachtin related limonoids (Rafiq 2010) and azadirachtin (Prakash et al. 2002, Sujanya et al. 2008) through callus and cell suspension cultures of neem. The work on the production of these bioactive compounds through callus cell and tissue culture is very important, supporting that M. azedarach cell and tissue cultures can be established and may be used as an alternative source for the production of limonoids. The limonoid compounds may be produced in any aseptic place/conditions without any changing season and in at low cost. Attempts were made to develop desired medium with the addition of different concentrations of sucrose.

Materials and Methods

Different explants (Immature flowers, nodular stems, immature embryos, leaves and petioles) of chinaberry plants were collected from Sindh University, Jamshoro, Pakistan. The whole experimental work was carried out at Plant Tissue Culture Lab., Institute of Biotechnology and Genetic Engineering, University of Sindh, Jamshoro, Pakistan. Experiment was carried out following the method reported by Rafiq and Duhot (2010). Initially samples were washed with distilled water thoroughly sterilized with specified concentration of sodium hypochlorite (commercial bleach) for 12 - 17 min, incubated in methanol for one minute, finally washed three times in sterile distilled water. Sterilization was carried out in laminar airflow cabinet.

Various explants cut in to small pieces (about 2 mm were in MS semi-solid medium with 1.0 mg/l 2,4-D, 5.0 mg/l GA3 and 3.0 mg/l NAA. Then cultures were left at 26 ± 2°C and 10/14
dark/light photoperiod for the callus induction. Cultures were maintained under these conditions and then transferred to the suspension liquid media after homogenization in optimized conditions.

Friable callus 0.5 g of fresh callus was taken from semi-solid MS medium, crushed and converted into small pieces and transferred to 5.0 ml MS liquid medium containing 1.0 mg/l of each 2,4-D, 5.0 mg/l GA₃ and 3.0 mg/l of NAA for homogenization. Then suspended homogenized callus cells in liquid media were transferred picking 2.5 ml in 30 ml suspension medium then left on shaking incubator at speed 200 rpm at 26 ± 2°C 18 days of incubation. Media and cultures broth were separated, and methanol was used as a solvent for the extraction of limonoids from cultures broth as well as from biomass.

Different parts of *M. azedarach* were dried at 50°C for 48 hrs, and then 3 g were ground in pestle mortar with autoclaved distilled water and 90% methanol separately. Extracts were centrifuged at 7000 rpm for 10 min, then the supernatant was taken and adjusted the volume up to 30 ml and finally samples were stored at −40°C. The dry callus ground 0.5 g in pestle mortar using autoclaved by D₂O and 90% methanol separately. Sample was centrifuged at 7000 rpm for 10 minutes then separated the supernatant finally adjusted volume up to 5 ml with distilled water and methanol separately. The extraction process was repeated twice. Samples were stored at −40°C for further chemical analysis.

Total limonoid contents were quantified following the method reported by Dai *et al.* (1999), 0.7 ml test sample plus 0.2 ml of methanol solution of vanillin (0.02 mg/ml) were mixed then samples were shaken manually and left at room temperature for 2 min. The 0.3 ml concentrated H₂SO₄ was added in three portions (0.1 ml each), and the mixture was stirred for 10 sec after each addition. With the addition of 0.7 ml methanol, the solution converted into two-layer mixture of a blue-green color. The solution was left at room temperature for 5 min, and then absorbance was noted at 577 nm using UV-visible spectrophotometer.

Per cent callusing response was determined using the method reported of Ding *et al.* (1981). The data were statically calculated by one-way ANOVA test using SAS 16.0 (SAS Inst. Inc., Cary, N.C., and the U.S.A) and Microsoft Excel 2010. The reported results are the average of three replicates and expressed as mean ± Sd.

**Results and Discussion**

Callus induction was made from in different parts of chinaberry with the addition of different PGRs in the medium as shown in Table 1. Variable callus induction effects were observed with the combination of different PGRs like 2, 4-D, BAP, GA₃ and NAA on different explants of *M. azedarach* (flowers, leaves, embryos, stem and petioles). It is seen (Table 1) that the nodules stem section under dark and light condition produced (73.3%) callus in medium with 1.0 mg/l 2,4-D, 5.0 mg/l GA₃ and 3.0 mg/l NAA as compared to other explants and PGRs combinations. Followed by immature flowers produced 66.6% callus induction response with the supplementation of 1mg/l 2,4-D, 1 mg/l BAP and 0.3 mg/l NAA on the 8th week of incubation (Fig. 1).

Total azadirachtin related limonoid contents determined from direct plant source with 90% methanol was used as a solvent for leaf extracts. It was noted that from direct sources highest total limonoid contents were 80.17 ± 8.9 µg/ml determined from leaf followed by 67.01 ± 4.9 µg/ml from flowers. With the use of water as a solvent 45.80 ± 2.5 µg/ml total limonoid contents were noted (Fig. 2). Limonoid contents from dry callus 0.5 g in 5 ml solvent were determined in immature flowers (40.85 ± 5.3 µg/ml) followed by nodular stem sections (26.24 ± 2.3 µg/ml) in methanol extracts. It was observed that limonoid contents were highest in methanol extracts as compared to water extracts (Fig. 3).
Table 1. Effect of different concentrations of plant growth regulators on callus induction.

<table>
<thead>
<tr>
<th>Group</th>
<th>2,4-D</th>
<th>BAP</th>
<th>GA$_3$</th>
<th>NAA</th>
<th>% Callusing response of Chinaberry explants</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Flowers</td>
</tr>
<tr>
<td>A</td>
<td>0.5</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>25.5</td>
</tr>
<tr>
<td>B</td>
<td>1.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>66.2**</td>
</tr>
<tr>
<td>C</td>
<td>1.0</td>
<td>1.0</td>
<td>-</td>
<td>0.3</td>
<td>33.8</td>
</tr>
<tr>
<td>D</td>
<td>1.0</td>
<td>1.5</td>
<td>-</td>
<td>-</td>
<td>NR</td>
</tr>
<tr>
<td>E</td>
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<td>-</td>
<td>5.0</td>
<td>3.0</td>
<td>50</td>
</tr>
<tr>
<td>F</td>
<td>1.0</td>
<td>1.0</td>
<td>-</td>
<td>0.3</td>
<td>30.8</td>
</tr>
<tr>
<td>G</td>
<td>2.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>37.5</td>
</tr>
<tr>
<td>H</td>
<td>2.0</td>
<td>1.0</td>
<td>-</td>
<td>1.0</td>
<td>33</td>
</tr>
<tr>
<td>I</td>
<td>2.0</td>
<td>1.0</td>
<td>-</td>
<td>1.0</td>
<td>NR</td>
</tr>
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</table>

Fig. 1 (A-C) Inoculated flowers, callus and suspension cultures for the production of secondary metabolites.

Fig. 2 (A-B). Total limonoid contents from different intact parts of chinaberry and dry callus.
Callus cell suspension culture showed continuous and stable multiplication of biomass. The time taken for cell growth and limonoids production have been studied by growing callus cells in 250 ml conical flask containing MS liquid medium. As shown in Fig. 3, limonoids production was slow during 1-6 days of incubation and its biomass reached the highest value on 18 days of incubation. Sucrose in cell suspension culture hydrolyzed immediately into glucose and fructose after initiation of culture and disappeared in 48 hrs from callus initiation. In the present work, different percentage of sucrose concentrations supplemented to cells suspension culture of *M. azedarach* are shown in Fig. 3. The highest level of limonoids was noted at 03% sucrose 10/14 dark/light photoperiod followed by 4%. The pH of agar medium and suspension culture was adjusted to 5.8 for the proliferation of callus and production of total limonoids. It was most important to investigate the effective pH value of cell suspension culture because production increased or decreased at pH 5.8 as kept for control. Lowest production of limonoids was determined at pH 5.4 and highest at pH 5.8 (Fig. 3) both in biomass and extracellular media (broth). Temperature also effects on the cells in cultures media. Therefore, its optimization is important. Almost cells belong to plant cultures between the range of 20 and 28°C and production of limonoids increased at temperature ranges (24 - 29°C) in suspension culture. The results mentioned in Fig. 3 showed that highest amount of total limonoids was produced at 28°C. As the temperature increased the reduction effect in limonoids was noted.

The phosphate, sulfate and nitrogen sources like KH₂PO₄, NH₄NO₃, KNO₃, NH₄SO₄ and urea (CO(NH₂)₂) at different concentration were added in the callus cell suspension culture media for the of production total amount of limonoid. With the addition of all chemicals, overall production of limonoids almost remains the same, although decreased but not increased as compared to without addition of chemicals (Fig. 4) in the presence of optimized chemicals in MS cell suspension culture medium.
The quality and frequency of callus and cell suspension culture depend on optimization of PGRs and culture environment. Like MS medium 2,4-D (1 mg/l), NAA (3.0 mg/l), GA3 (5.0 mg/l) and 3% sucrose were added in MS medium and inoculated at 26 ± 2°C. In present work, supplement of additional chemicals did not increase the product as compared to without chemicals. Good quality of callus either increased yield or improved the nutritional quality.

Plant bio-products produced through cell and tissue culture technology has been extended by Simões et al. (2012) for commercial use and tissue culture with optimized conditions like time of incubation, pH, temperature. Different additives like PGRs and nitrogen sources may increase the production through callus cell suspension cultures (Abbas et al. 2018). As mentioned in Fig. 3, production of limonoids and biomass were found to be slow and same type of results were reported by Cheng et al. (2006) where decreased of biomass and limonoids production after 22 days of time of incubation was observed. After every 18 days of incubation 10% media with suspension culture was transferred to new liquid media having same PGRs for long time stable production of limonoids. Sucrose is an important as carbon and energy source for all plants (Al-Khayri et al. 1996, Nhut et al. 2000). It was reported that the number of parameters related to secondary metabolites in plant cell cultures such as growth rate and yield are hardly affected by initial sucrose concentrations (Benavides 1997, Wang et al. 1997). Sucrose in cell suspension culture hydrolyzed immediately into glucose and fructose after initiation of culture and disappeared in 48 hrs from callus initiation. The present results are closely supported by Babu et al. (2008), who worked on enhancement of salannin (limonoid having insecticidal activity) productions in cell suspension cultures of Azadirachta indica. Results indicated that changing in sucrose concentration is an effective method for the production of high level of secondary metabolites in cell cultures (Pasqua et al. 2005). Optimized pH is extremely important as uptake of nutrients and PGRs are influenced by pH by regulating the solubility in cell suspension medium (Bhatia and Aswath 2005). Most of the reports showed that the pH of plant cell suspension culture media was maintained at 5.8 (Fig. 3), which justifies present results; it may be possible that the stability of azadirachtin is higher in acidic pH (Jarvis et al. 1998). It has been reported that A. indica cells with the reduction of pH in long term lose viability and died (Potters et al. 2007). Temperature also effects on the cell cultures and optimization is important. In present work, the highest amount of total limonoids were produced at 28°C with Choi et al. (2000), who reported
that cells belong to plant cultured between was in conformity 20 and 28°C and production of limonoids increased at 24 to 29°C by suspension culture in *Taxus chinensis*.

Optimized production of valuable bioactive compounds through callus cell suspension cultures could be useful for humans and used safely as medicines without any side effects as compared to synthetic medicines. This technique can also be applied to the other medicinal plants for multipurpose applications and may be baseline investigations of the production of limonoids through callus cell suspension under controlled conditions. Most of the medicinal plants could be selected and used for the production of secondary metabolites through callus cell suspension cultures.

The production of limonoids from direct source of *M. azedarach* L. is laborious and much expensive than callus cell suspension culture considered to be the most appropriate method for the production of total limonoids. Optimized conditions as discussed here can be used for the production of biologically active compounds from other medicinal plants. This method could be more valuable alternative for the production of bioactive compounds, which could be used for biological control as well as other medicinal purposes.

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