Enhanced production of andrographolide –effect of gamma irradiation on *Andrographis paniculata*

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**ABSTRACT**

One of the most widely applied empirical approaches to optimize the synthesis and accumulation of secondary plant products is the use of physical mutagen like γ –ray. The effects of gamma irradiation on secondary metabolite production in callus cultures derived from four doses of γ-irradiated seedlings of *A.paniculata* were studied. The result of this study indicated that γ-irradiation positively influenced andrographolide production in *A.paniculata* from the callus cultures. The significant impact of this study was that the callus cultures produced from non-irradiated seedlings showed the absence of andrographolide. The effect of four doses of γ-irradiation on the *invitro* production of andrographolide from the leaf explants of *A.paniculata* was studied. An enhanced rate of andrographolide was found in all the doses of samples using HPLC analysis after gamma irradiation. Tissue culture increased the efficiency of mutagenic treatments, which lead to new possibilities in plant improvement and the effect of it is a vast field for research investigations. The presence of the important secondary metabolite andrographolide is not yet seen reported from the callus cultures of A*. paniculata.*.So far andrographolide is extracted from intact plants only. The *in vitro* production of this compound is not yet reported. This study provides the feasibility to produce the important secondary metabolite *in vitro.*

**INTRODUCTION**

Deforestation, globalwarming and pollution pose a threat to the welfare of plants in many parts of the world. The acute shortage of raw materials has also meant poor quality of the life saving drugs and necessities their replenishment through concerted efforts. Now -a -days natural plant extracts are manufactured on a large scale by several multinational companies for the preparation of many formulations.

*Andrographis paniculata* Nees. (Acanthaceae) known, as *Kalmegh* in Ayurveda is an annual herb. The genus *Andrographis* consists of twenty species of small herbs of which *Andrographis paniculata* is the most popular. The herb is reported to possess an astringent, anodyne, tonic and alexipharmic properties and is useful in dysentery, cholera, diabetes, consumption, influenza, bronchitis, swellings, itches and piles. *A. paniculata* is found in the Indian pharmacopoeia and is prominent in at least twenty-six Ayurvedic formulas(Anon, 1985)(1). In Traditional Chinese medicine, *A. paniculata* is an important “cold property herb”. Most of the biological actions of *A. paniculata* has been described due to the presence of Andrographolide, a diterpene lactone present in these plants.) Chakravartni and Chakravarthi (1952) observed that the leaves of Kalmegh contain the maximum amount of andrographolide content (2.5%) from the leaves and the stem posses the lesser amount (2).Secondary metabolites isolation from callus will help to satisfy the needs to some extent.

Owing to the high economical and pharmacological importance of secondary metabolites a large number of industries are deeply interested in the production of these compounds. The plant tissue culture technique has widened the scope and opened new vistas for the production of secondary metabolites.

The land area is shrinking due to fast increasing population growth. Significant numbers of young plants are lost due to unscrupulous cutting and destruction. So the need of the hour is a quantum jump in to non-conventional methods of tissue culture. Plant tissue culture based on the principle of totipotency, is a boon to studies of the biosynthesis of secondary metabolites and provides an efficient means of producing economically important plant products.

MATERIALS AND METHODS

The seedlings were subjected to four doses of gamma irradiation at Rubber Board, Puthuppally, and Kottayam,Kerala.India. The doses were 0.5kR, 1kR, 1.5kR and 2kR. The time taken for each irradiation was calculated using the formula 276.74750Kilorad/Hour. Based on this calculation 0.5kR irradiation was done by giving gamma irradiation for 7 seconds, 1kR by 13 seconds, 17 seconds for 1.5 kR and 26 seconds for 2kR.

The seedlings after irradiation were allowed to grow in the green house. After ten days the third leaf from the apical portion was selected for *in vitro* inoculation..For callus induction, each treatment consisted of 10 explants in test tubes, with three replications and observations were recorded in Table1.

**Table1: Hormones used with Full Ms for Callus initiation**

|  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Medium strength | Auxins | | | | Cytokinins | | Gibberrellin | Additive |
| Full MS | 0.25 |  | 0.05 |  |  |  |  |  |
| 0.10 |  |  |  |  |  |  |  |
| 0.05 |  | 2.50 |  |  |  |  |  |
|  |  | 2.50 | 0.05 | 0.10 |  |  |  |
|  |  | 0.50 | 2.50 |  |  |  |  |
|  |  | 0.05 | 0.05 |  |  |  |  |
|  | 0.10 |  | 0.10 |  |  |  |  |
|  |  |  |  | 0.10 | 0.05 | 1.00 |  |
|  |  |  |  | 0.10 |  | 0.10 |  |
|  |  | 0.05 |  |  | 0.05 |  |  |
|  |  |  |  |  | 0.05 | 0.05 |  |
|  |  |  |  |  | 0.10 | 0.10 | 2.00 |

**Standardization of the medium for callus proliferation**

The proliferated calli, after fifteen days, were sub-cultured into callus proliferating medium containing basal MS medium supplemented with various concentrations of cytokinins (BA/KIN) or in combination with auxins (NAA, 2,4D).The medium was also enriched with vitamins, phenylalanine and the additive polyethyleneglycolin various concentrations of auxins and cytokinins. For callus proliferation, each treatment consisted of ten experimental bottles, with three replications and the fresh weight of the callus after forty and seventy days of inoculation were taken. Relative growth rate (RGR) of calluses was estimated in GRAPH 4.5,4.6.

**Secondary metabolite analysis**

The medium used for callus proliferation was taken as medium for callus andrographolide accumulation (Table2). The calli kept in the proliferating medium were grown for 70 days and were taken out and allowed to dry in the hot air oven at 330C. Before drying the fresh weight of the calli was taken. The dry weights of the calli were also taken after proper drying and they were stored at 40C, after proper labeling. The andrographolide content was estimated by HPLC analysis

**RESULTS AND DISCUSSION**

EFFECT OF THE GAMMA IRRADIATION ON ANDROGRAPHOLIDE PRODUCTION ( L-PHENYL ALANINE IN COMBINATION WITH KINETIN AND NAA)

0.5kR

Stock calluses were transferred into MS medium fortified with 50mg|l phenyl alanine, 1mg|l kinetin and 2mg|l NAA and the impact of auxins and cytokinins together with L-phenyl alanine on andrographolide synthesis was analysed after a period of seventy days. The fresh weight observed was 10930mg. The presence of alkaloid was 0.0049%W|W which was greater in the non-irradiated plants .The precursor L-phenyl alanine and gamma irradiation promoted the alkaloid production when the seedlings of *A.paniculata* were irradiated with 0.5kR of γ ray

1kR

Stock calluses transferred to full MS medium favoured an increase in the andrographolide production. The positive influence of precursor, L-phenyl alanine was exhibited by the callus cultures with an increase in andrographolide production, 0.0039%w|w. In non-irradiated calli the andrographolide content was (0.0). In the MS medium supplemented with 1mg|lkinetin, 2mg|l NAA and 50mg|l L-phenyl alanine the andrographolide showed a remarkable increase with gamma irradiation at the dose of 1kR to the seedlings (TABLE 3)

1.5kR

1.5kR of γ-irradiation together with L-phenyl alanine induced andrographolide de production significantly and a progressive increase in the andrographolide production was noticed (0.0042%w|w). 1.5kR gamma irradiation favoured callus proliferation (fresh weight: 11040mg) and andrographolide production. Even though the precursor was supplemented into MS medium there was no significant impact on andrographolide production in control explants where the content was negligible (0.0). Thus γ- irradiation favoured andrographolide production and maximum accumulation of andrographolide using precursor was reported in this dose.

2kR

The positive influence of gamma irradiation and L-phenyl alanine was significant in callus proliferation and andrographolide production. The andrographolide production enhanced up to 0.0072%w|w. In the control plant calluses, the presence of andrographolide was not reported. Maximum callus growth and proliferation recorded was 19242mg of fresh weight. It was found that by increasing the dose of γ – irradiation the accumulation of andrographolide was also increased.(GRAPH 4.7)

Gamma irradiation had a significant impact on andrographolide production in *A.paniculata*. In non–irradiated seedlings, the presence of andrographolide is not reported from calluses produced from nutrient media supplemented with various growth regulators.

In the present study, the 2kR irradiated seedlings exhibited a large increase in andrographolide production in the callus cultures when the medium was fortified with 50mg|l phenylalanine, 1mg|l kinetin and 2mg|l NAA. The andrographolide produced was 0.0072%w|w.The gamma irradiation up to 2kR increased the andrographolide production from the calluses.

Maximum andrographolide production was found in the callus produced from seedlings irradiated with 2kR of γ-irradiation proliferated in a medium supplemented with the precursor-phenylalanine. In *A.paniculata* the andrographolide content in callus produced from nonirradiated explants raised in a medium containing phenylalanine was negligible (0.0). But a significant increase in andrographolide accumulation (0.0072%w\w) was noticed in the callus produced from explants after 2kR of γ-irradiation. This observation was in contrast to the observations made by Butcher and Conolly (1971)(3). They reported that callus and suspension cultures of *A.paniculata* had been shown to produce three new sesquiterpene lactones, which had been named paniculides, A, B, C and not the andrographolide (diterpenes) produced by the intact plant. They also reported that the callus derived from leaves, stems, hypocotyls, roots and embryos accumulated paniculides, but no andrographolides were detected. They attributed this abnormal terpenoid metabolism is due to the lack of organized tissues in these cell cultures. This result was in agreement with the reports of Zhang-Hua et al (1999)(4). They studied the influence of gamma ray on Shikonin formation on cultured cells of *Onasma paniculatum*. The optimum gamma radiation dose was 1500 R, which gave a 144.6% increase in Shikonin content compared to the control (unirradiated) callus gamma irradiation together with the presence of higher level of PEG. Andrographolide production did not show gamma irradiation dose dependence in all the samples studied.

This result was in agreement with the reports of Zhang-Hua et al (1999)(4). They studied the influence of gamma ray on Shikonin formation on cultured cells of *Onasma paniculatum*. The optimum gamma radiation dose was 1500 R, which gave a 144.6% increase in Shikonin content compared to the control (unirradiated). Li et al (2005) studied the effect of γ-irradiation on development, yield and quality of micro tubers *in vitro* in *Solanum tuberosum* L(5).The report of the present study was in agreement with the study on *Solanum tuberosum* done by Li et al, 2005. When plantlets were irradiated with 4Gy a significant increase in the fresh mass and low doses of irradiation increased the starch content of micro tubers and also significantly increased the protein content of micro tubers. Subodh Kumar (1987) had reported that gamma rays induced mutants showed changes in phenolic compounds in *Chrysanthemum.(6)*The positive effect of gamma rays on *Capsicum annum* was observed by the enhanced capsicine content (Bansal, 1969)(7). According to of Kurz and Constabel (1979) the enrichment of the culture medium with additives or amino acids was highly beneficial for the intracellular accumulation of andrographolide(8). Yeoman (1987) was the of the opinion that the rate of cell growth directly regulates secondary metabolism by affecting the kinetic partitioning of precursors between primary and secondary metabolic pathways.(9)

The increase in biomass caused an increase in andrographolide accumulation in 2kR-irradiated explants and this indicates that concentration of andrographolide increased with the increase of cell mass.

Rech S.B et al (1998) reported that in *Rauwolfia sellowii* alkaloids were found in the suspension cultures from early growth stages and the maximum product accumulation was observed with the end of exponential growth phase. They also observed that product formation is growth associated, since the concentrations of alkaloids increased with increase in cell mass.(10)

The accumulation of metabolites in callus produced from γ- irradiated seedlings can be explained on the basis of the experimental studies of Chagvardieff et al (1989) on *Euphorbia characias*. They studied the effect of gamma irradiation in photoautotropic suspension cultures of *E.characias* and observed the arrest of cell divisions and a subsequent over accumulation of sucrose and dry matter when gamma irradiation was applied.(11) The arrest in cell division may cause over accumulation of sugars, which may lead to the synthesis of other metabolites.

In this study, the effect of gamma irradiation on secondary metabolite production may be accelerated by the presence of additives and precursor. Gunckel and Sparrow (1961) reported that gamma rays are known to influence plant growth and development by inducing cytological, genetical, biochemical physiological and morphogenetic changes in cells and tissue.(12) The use of optimal levels of phytohormones for the growth and addition of various chemically defined compounds to the medium as inducers stimulated the production of secondary metabolites.

The studies on accumulation of andrographolide in callus raised from explants after various doses of γ-irradiation proved that the andrographolide accumulation showed a dose dependent variation and maximum andrographolide production was found in the callus produced from seedlings irradiated with 2kR of γ-irradiation.

CONCLUSION

MS medium supplemented with three different concentrations of 2,4-D, kinetin BA, NAA together with additives and precursor were used for callus proliferation and production of secondary metabolite.Among the different media tested, maximum andrographolide accumulation was found in the medium supplemented with phenylalanine. Callus proliferation rate showed somewhat dose dependent increase with increase in γ- irradiation dose in different media tested. The relative growth rate also showed a dose dependent increase from 0.5kR to 2kR of gamma irradiation. The use of additive (PEG 6000) along with growth regulators showed best response in callus produced from seedlings, exposed to 1.5kR of gamma irradiation.Callus produced from 2kR gamma irradiated seedlings showed enhanced growth when the medium was supplied with the precursor along with kinetin and NAA. The use of 20 gms PEG 6000 together with 1mg|l kinetin and 0.5mg/l 2,4 D positively influenced the enhancement of fresh weight and callus proliferation in all doses of gamma irradiation except in 1kR.

Andrographolide accumulation was 0.2089%W\W in the leaves of field grown control plants. When the calli were grown in a medium augmented with plant growth regulators the andrographolide production was 0.0029%W/W. L-phenylalanine acted as a strong precursor for the production of andrographolide in all the four samples of γ-irradiated seedlings.NAA and kinetin also enhanced the rate of andrographolide production in the irradiated seedlings.

Callus developed from seedlings irradiated with 2kR of γ-irradiation proliferated in a medium supplemented with 50mg/lphenylalanine, 1mg/l kinetin and 2mg/l NAA produced maximum andrographolide (0.0072%W/W).

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**4.7 GRAPH SHOWING THE EFFECT OF GAMMA IRRADIATION**

**ON ANDROGRAPHOLIDE PRODUCTION**



