UV-B Induced Changes to the Physiological and Phytochemical Parameters of Phyllanthus amarus Schum

N. Shanthi1, S. Murugesan1, S.M. Janetta Nithia2, M. Kotteswari1, S. Shyamala Gowri1,2

1PG and Research Department of Botany, Unit of Plant stress physiology, Pachaiyappa’s College, Chennai-600 030, Tamil Nadu, India
2Department of Botany, Sri Meenakshi Government Arts College for Women, Madurai, India

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Abstract

UV-B is a growing concern due to the rise in UV-B levels on the surface of the earth as a result of the loss of stratospheric ozone. Increased levels of UV-B radiation can in fact negatively alter plant physiological processes, growth and productivity. However, when researching the effects of UV-B on medicinal plants like Phyllanthus amarus and in the tropical area under field conditions, there are some curious phenomena have been discovered. Enhanced UV-B radiation has greatly improved the growth of P. amarus. The outcome of the photosynthetic pigment showed increased UV-B enhanced synthesis of Chlorophyll or the accumulation of Chlorophyll pigments in the treated plants compared to non-UV-B treated plants. The concentration of UV-B absorbing pigments also increased due to enhanced UV-B radiation in P. amarus. The synthesis of secondary metabolites such as flavonoid and phenol content was increased under UV-B treatment as compared to control. The UV-B radiation enhances the grade of the medicinal plant by improving the medicinally active compounds. This enhanced impact of UV-B could be important to observe when studying the phytotherapeutic function of P. amarus in health aspects of human life.

Keywords: UV-B, Chlorophyll, Flavonoids, Phenol, Saponin, Alkaloids, Phyllanthus amarus.

INTRODUCTION

The depletion of the ozone layer caused by CFCs and other ozone-depleting compounds increases the amount of UV-B radiation (280-320 nm) on the emission on the earth's surface. The UV-B radiation is one of the most critical abiotic stresses in the ecosystem and it’s influence on all physiological and phytochemical parameters. On the other hand, the UV-B effect of plants depends on plant species, the position where plants grow and the interactive action another environmental issue [1].

The result of UV-B on the tropical organism is an explicit link, since ozone depletion and corresponding improvement in UV-B are mostly in the low latitude region. UV-B decreased plant height, leaf area and plant dry weight increased auxiliary branching and curling of the leaf [2]. Metwally ([3] revealed that when an exposure of UV-B in a number of weeks, leaf area and plant dry weight of rice was considerably was significantly diminish. Several of the previous studies have agreed to react to UV-B irradiation with relevancy growth, production of dry matter and physiological and biochemical changes depending on the species [4, 5].

Many plant species are unaffected by UV-B irradiation and many others tend to be stimulated by their development and other biological processes [6, 7].

Measurements of various alternative key parameters like chlorophyll and antioxidant content and levels of UV-B engrossing compounds have additionally established to be helpful indicators of UV-B tolerance or sensitivity, since these parameters show the foremost speedy response to UV-B stress [8]. UV-B affects the reduction of photosynthetic activity mainly associated with the degradation of PS II proteins. It may induce photobleaching and photodegradation of photosynthetic pigments [9]. Chlorophylls and carotenoids could also be adversely suffering from relative great amounts of UV-B, with carotenoids typically being less affected than chlorophylls [10].

In constant time, plants respond to environmental stress by synthesizing a variety of secondary metabolites for defense purpose. Ultraviolet (UV) radiation influences the accumulation of endogenous plant secondary metabolites [11, 12]. Several studies have investigated the UV-mediated...
accumulation of flavonoids and associated phenolics, alkaloids, terpenes, lignans, steroids, curcumines, saponins, phytosterol and glucosides. However, there is an increasing awareness that a far larger variety of metabolites accumulate in ultraviolet light exposed plants. Especially tropical plants have developed a variety of mechanisms to protect against UV-B induced damage. The impact of increased UV-B radiation on healthy plants in tropical regime has gained considerable attention in today's world.

Experiments on medicinal plants in tropical ecosystem have received huge attention in nowadays world. As a result of the impact of UV-B on medicinal plant analysis is extremely less compared to agricultural plants. *P. amarus* is one in every of the standard healthful plants belong to Euphorbiaceae family and it’s a little annual herb full-grown on tropical and subtropical region. Because of its wide selection medicinal properties it’s utilized in the Indian Ayurvedic systems from the traditional times. The plant extract primarily used for Jaundice, Anemia, Bronchitis, Leprosy, Asthma etc. We hypothesized that the impact of increased UV-B on growth, photosynthetic pigment synthesis and accumulation of phytochemical compounds would be greater than the UV-B radiation not received by the plant.

**MATERIALS AND METHODS**

**Plant materials**
Certified seeds of *Phyllanthus amarus* obtained from the Farm aid, Chennai was shown in experimental plots in the Pachaiyapppa's College Botanical Garden, Chennai. One set of plants was grown under ambient solar radiation and other under 20% UV-B enhanced solar radiation.

**Plant growth and UV-B treatment**

The seeds were soaked overnight in the running water. Separate soil beds were prepared for control (ambient) and UV-B treatment and seeds were sown in these experimental plots. The plants were watered regularly and care was taken to avoid microbial or pest infection during the experimental period. Plants with the first foliage leaf stage were used for UV-B treatment. UV-B treatment was given to these plants for 4 hrs daily from 10 a.m to 2 p.m. Treatment was continued under ambient solar radiation and 20% UV-B enhanced solar radiation supplemented by a Philips TL40W/12 sunlamp (Gloelampenfabrieken, Holland). The first formed leaves were collected at different time periods and all the physiological and biochemical analyses were carried out.

**Measurement of radiation**

A Li-Cor Li-188B quantum/radiometer (Li-Cor., Inc., USA) with suitable photodetector was used to measure all the visible and photosynthetically active radiation. Radiation below 400 nm was determined by an IL 700 radiometer with a SEE 400 photodiode detector (International Light Inc., USA).

**Determination of growth**

Shoot length was determined soon after the seedlings were uprooted. All the measurements were the mean of twenty randomly selected samples.

**Estimation of pigments**

**A. Chlorophyll**

Pigments were extracted in 80% acetone and the amount of total Chl, Chl *et al.*, Chl *b* and carotenoid was quantified using the formulae of Wellburn and Lichtenthaler [13].

\[
\text{Chlorophyll } a \quad (\text{mg/l}): \quad (12.21 \times A_{666}) - (2.81 \times A_{646})
\]

\[
\text{Chlorophyll } b \quad (\text{mg/l}): \quad (20.13 \times A_{666}) - (5.03 \times A_{663})
\]

\[
\text{Total Chlorophyll} \quad (\text{mg/l}): \quad (7.18 \times A_{663}) + (17.32 \times A_{646})
\]

**B. Carotenoid**

The concentration of total amount of carotenoids was estimated in the 80% acetone extract by measuring the absorbance at 480 nm. Mackinney’s (14) formula was used to correct the Chl interference.

\[
\text{Carotenoids } \quad (\text{mg/l}): \quad 1000 A_{470} - (3.27 \text{ Chl } a) - (104 \text{ Chl } b)
\]

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**C. Flavonoid**

Fresh leaf samples equivalent to 100 mg were cut into small pieces and incubated overnight in 5 ml of 80% acidified methanol (80:20:1 of methanol:water:HCl) at 4°C in the dark. After centrifugation to remove debris, the absorbance at 315 nm was taken and the flavonoids content was expressed as µg/g leaf fresh weight [15].

**D. Anthocyanin**

Anthocyanin were extracted from the leaves by grinding the leaves in 80% acidified methanol (80:20:1 of methanol:water:HCl). After centrifugation, the clear extract was used to estimate the concentration of anthocyanin by measuring the absorbance at 530 and 657 nm, according to Mancinelli et al., [16].

\[
\mu g/g \text{ fresh weight: } (A_{530}) - (0.3 \times A_{657})
\]
Preparation of extracts
The fresh leaves *P. Amarus* (Euphorbiaceae) of control and UV-B irradiated plant leaves were soaked individually at room temperature in methanol for 72 hrs in the ratio of 4:1. The extract was filtered using Whatman filter paper. This was repeated for 2 to 3 times and similar extracts were pooled together and concentrated at 45°C under reduced pressure using a vacuum rotary evaporator. The concentrated crude methanol extract was subjected to the preliminary phytochemical screening. The concentrated crude methanolic extracts were subjected to qualitative phytochemical analysis [17].

Yield of methanol extracts
The concentrated crude methanol extracts were weighed independently and stored in 4°C in the refrigerator until further usage.

Qualitative phytochemical analysis
The different qualitative chemical tests were performed for establishing the profile of the given extract for its chemical composition. The crude drug was redissolved in methanol and subjected to various phytochemical analyses. The standard protocols were used for qualitative analysis of samples to check for the presence of alkaloids, flavonoids, phenols, saponins, tannins and terpenoids.

STATISTICAL ANALYSIS
The data on cell viability were analyzed by using the one way ANOVA followed by the Dennett’s multiple comparison tests with equal sample size by using SPSS 17.0. The difference was considered significant when *p*<0.005. All the values were expressed as mean ± standard deviation (S.D).

RESULTS AND DISCUSSION
Effects of increased UV-B radiation were studied on one of our native medicinal plants, *P. amarus*. The improvement in solar UV-B was 20% above ambient radiation. The analysis of growth parameters, biochemical constituents and synthesis of secondary metabolites, once exposure to increased solar UV-B, was disbursed at associate interval of five days.

All the parameters were analyzed once ten days from the onset of UV-B treatment.

Changes in the growth characteristics
The effect of enhanced UV-B radiation accelerates the shoot length in throughout the growth period (Fig 1). Where as in non UV-B treated plant showed the normal growth level.

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Fig-1: Typical morphological change in *P. amarus* grown under ambient and enhanced UV-B radiation

The growth was promoted from 10th day onward. The difference between ambient and UV-B treated plants was found to be 30% (Fig 2). Elevated UV-B radiation directly affects plants through growth and development, biomass production, leaf characteristics and flowering time [18,19]. Sullivan and Teramura [20] stated that Gymnosperms tree species are tolerant of UV-B, whereas, some conifers show an increase in growth in response to UV-B radiation species. Many plant species are unaffected by UV-B radiation and a number of others are apparently aroused in their growth, however most species are vulnerable and damage, results such as rice and maize [21, 22]. Current studies under semi natural field conditions have found that UV-B radiation is not detrimental to growth and physiology [23]. Furthermore, UV-B radiation impacts are species specific and depend upon interaction with alternative environmental factors (1).

Fig-2: Changes in the shoot length in *P. amarus* grown under ambient and enhanced UV-B radiation. The values represent an average of 20 plants, Mean ± SE, n = 20
Changes in photosynthetic pigment

Changes in chlorophyll pigments are evidences of high UV radiation tolerance of analyzing plants. Chlorophyll-a and b are the main photosynthetic pigments in all higher plants. Whereas, carotenoids are the main UV-protecting pigments. Chlorophyll content levels have been determined to provide evidence of UV-B effects on chlorophyll metabolism. Spectral analysis showed that the total chlorophyll content of control plant was significantly less when compared to UV-B treated plants (Fig 3). The total chlorophyll content was promoted speedily in the 10th Day to the 25th day. UV-B stress exaggerated a production of chlorophyll, significantly 20th day. On the 25th day the chlorophyll synthesis was declining due to senescence. However, the percentage of promotion increased with an increase in plant growth. The content of chlorophyll was step by step exaggerated throughout the complete development period. An increase in chlorophyll following enhanced UV-B was observed in leaves of Bromus catharticus [24] and in wheat seedlings [25]. Chlorophyll synthesis, though addicted to the light absorbed by the phytochrome, could most likely be a result of the co-action between UV-B and phytochrome photo sensors [26]. The response of chlorophyll levels to exaggerated UV-B radiation depends on status and on the biological phase of plants [27]. Likewise, Li et al., [28] found that UV-B radiation increased the chlorophyll content synthesis during their growth of P. persicae. As a result of increased UV-B radiation, miRNA/genes rates have been elevated compared to untreated UV-B plants.

![Fig-3: Changes in the total Chl content of P. amarus grown under ambient and enhanced UV-B radiation. The values represent an average of 3 independent measurements. Mean ± SE, n=3.](image)

Compared to the control plant, the synthesis of carotenoid pigment content in UV-B stressed plants are considerably higher up to the 25th day (Fig 4). Such UV-B evoked pigment synthesis was twofold elevated than control plants. For photosynthetic organisms, the protecting role of carotenoids against high UV-B radiation is the standard and protective role of carotenoids in blue-green algae against UV-A radiation was reported by Monika Ehling-Schulze et al., [29]. UV-B exposure accumulated the pigment synthesis capability of plants [30]. Recent studies have highlighted that ecologically relevant low UV-B levels will trigger the induction of synthetic resin compounds and also the accumulation of carotenoids and GS.

![Fig-4: Changes in the total carotenoid content of P. amarus grown under ambient and enhanced UV-B radiation. The values represent an average of 3 independent measurements. Mean ± SE, n=3.](image)
Changes in non-photosynthetic pigments

The majority of the plants respond to UV-B radiation by producing UV-B absorbing compounds. The present investigation stated that UV-B radiation increased the accumulation of flavonoids in *P. amarus* as compared to the ambient plants. The concentration flavonoids increased from 10th Day onwards (Fig 5). The maximum 20% increase of flavonoids was observed in UV-B treated plant at 25th day as compared to control.

Flavonoids are probable antioxidants and are used as a vital interest recently, due to their impending useful effects on human health in fighting diseases. Several studies used acute increased UV-B levels and targeted on the rise of flavonoids together with anthocyanins, and hydroxycinnamic acids as UV-B protection by serving as active oxygen scavengers [31, 32]. The stimulation of flavonoid biogenesis by UV-B is mainly due to of gene expression [33]. The UV-B stress activates the transcription of genes encoding the primary protein of the flavonoid biogenesis pathway, Chalcone Synthase (CHS) [34], and enzymes that act downstream within the pathway [35]. Analysis accepted deoxyribonucleic acid sequence components within the promoter regions of flavonoid biogenesis genes that are required for UV-B, and transcription produced proteins that act within these components to mediate the reaction. Research has acknowledged DNA sequence elements in the promoter regions of flavonoid biosynthesis genes that are required for induction by UV-B, and transcription factor proteins that interact with these elements to mediate the reaction [35].

The synthesis of anthocyanin content showed a maximum elevation in plants with UV-B stressed an increase in the amount of anthocyanin was determined at any point in the UV-B treated plant, with a limit of 25% at 25th Days (Fig 6). Accumulation of anthocyanin was important due to UV-B interactions and plants. Most of the studies have shown that enhanced anthocyanin was chiefly because of UV-B irradiation impact. Ambasht and Agarwal [36] discovered that over 275% increase of anthocyanin content in maize. Anthocyanin has terribly weak absorption within the ultraviolet radiation-B regions and considered UV screens solely at very high concentration. Escobar-Bravo *et al.* [1] revealed that UV-B evoked accumulation of anthocyanin shield the chemical action equipment from the damaging result of UV-B radiation.
Prefer to anthocyanin, flavonoid concentration was additionally raised in UV-B treated seedlings when eight days of treatment. In general, the flavonoid increase is linearly dependent on the UV-B effect [37]. Del Valle et al., [38] all over that flavonoid concentration will diminish the UV-B penetration and guard the photosynthetic apparatus up to some extent, however, if depends abreast of an intensity level, which can vary in numerous species. Likewise UV-B induction of anthocyanin accumulation was discovered conjointly in Arabidopsis [39], eggplant [40] and grape [41].

Phytochemical analysis of ethanol extracts of leaves of *P. amarus*

A preliminary phytochemical investigation of the ethanol extracts of leaves of *P. amarus* revealed the presence of flavonoids, tannins, phenolic compounds, alkaloids, glycosides, fats and carbohydrates. Phytochemical analysis of leaf extract of *P. amarus* showed the high quantity of secondary metabolites in the UV-B treated plant when compared to the control plant (Table 1).

### Table-1: Comparative analysis of phytochemical compounds in methanol extracts of leaves of *P. amarus*

| Phytochemical test | Control | Treated |
|--------------------|---------|---------|
| **I. Alkaloids**   |         |         |
| 1. Mayer’s Test    | +       | +++     |
| 2. Wagner’s Test   | +       | +++     |
| 3. Hager’s test    | +       | +++     |
| 4. Dragendorff’s test | +     | +++     |
| **II. Flavonoids** |         |         |
| 1. Alkaline reagent test | +   | +++     |
| **III. Fixed oil test** |       |         |
| 1. Spot test       | +       | +       |
| **IV. Carbohydrate** |      |         |
| 1. Fehling’s test  | +       | +++     |
| 2. Benedict’s test | +       | +++     |
| **V. Glycosides**  |         |         |
| 1. Borntrages’s test | +     | +++     |
| **VI. Saponins**   |         |         |
| 1. Foam test       | +       | +++     |
| **VII. Phytosterols** |      |         |
| 1. Libermann-Burhard’s test | +  | +++     |
| **VIII. Phenols**  |         |         |
| 1. Ferric chloride test | + | +++     |
| 2. Gelatin test    | +       | +++     |
| 3. Lead acetate test | +    | +++     |

Note: + = present +++ = Present strongly

The synthesis secondary metabolites like flavonoids and phenol content were increased under UV-B treated condition when compared to control. Such increase was 0.5% higher in UV-B treated plants. At the same time phenol content was more as compared to flavonoid content in UV-B treated plants (Table 2).

### Table-2: Comparative analysis of phenol and flavanoid in ethanol extracts of leaves of *P. amarus*

| S.No | Extract | Phenol content (mg GAE/g DW of Ethanol Extract) | Flavonoid content (mg QE/g DW of Ethanol Extract) |
|------|---------|-----------------------------------------------|-----------------------------------------------|
| 1    | Control | 100.00 ±4.00                                  | 146.02 ± 11.99                                |
| 2    | Treated | 133.33 ± 6.11                                 | 201.59 ± 14.55                                |

Values are mean of triplicates ± Standard deviation.
The plant possesses a high potential for production of valuable phytocomponents under field condition to guard the plant from the damaging effects of UV-B radiation. UV-B exposure additionally led to elevated completely different phytochemical compounds like alkaloids, phenols, phytosterol and flavonoid (Figs. 7 & 8). The improvement was thrice over non stressed plant. The experimental result evidenced that those phytochemical compounds are defence compound. Similar kind of results was projected by Kumari and Agarwal [42] the UV-B radiation having an efficiency to reinforce the amount of terpenoid phenyl propanoid and natural antioxidants.Turtola et al., [43] stated that spruce has high quantity of penolic compound in comparison to Scotch pine as a result of the previous species have higher screening capability against UV-B irradiation. Hagen et al., [44] stated that the results of post harvest UV-B irradiance with UV-B radiation and reportable the augmented price in phenolic content. The phenol will act as shielder after they exposed to UV-B radiation [45]. In other research Behn [46] ascertained that UV-B radiation improves the synthesis of terpenoids in the mint family. Johnson et al., [47] investigated the accumulation of terpenoid, glucoside in basil leaves. The optimum UV-B radiation augmented the production of Glyrrhizic acid, a triterpenoid in Glyrrhiza uralensis [48]. The UV-B light induce alkaloid in C. roseus leaves [49]. Zu et al., [50] reported the augmented content of alkaloids in plant exposed to increased UV-B radiation. Phytochemical compounds are used synthesized by UV-B radiation, which act as UV-B absorbent and so capable protects the plant from the harmful impact of the plant. Increased UV-B radiation raises the elevated level of several individual flavonoids and phenolic acids anywhere they are contained in the Betula pendula response [51].

CONCLUSION

Plants have the ability to adapt the ever-changing climate and improve the growth and physiological activity. Environmental factors affect the chemical composition and induce changes in the amount of secondary metabolites and promote the accumulation of bioactive compounds. It has increased the quality of medicinal plants. Therefore, this changing impact of UV-B may also be important to observe in the discovery of the phytotherapeutic activity of P. amarus, in the health aspects of human life.

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