Exogenous Polyamine Treatment Prevents UV-A Induced Cellular Damage and Photoinhibition of Photosystem II in Lichen Physcia semipinnata (J. F. Gmel.) Moberg

Dilek Ünal¹, İnci Tüney Kızılkaya²*

¹University of Bilecik, Faculty of Science and Literature, Department of Molecular Biology and Genetics, Bilecik, Turkey
²Ege University, Faculty of Science, Department of Biology, Bornova, Izmir

Received: 28 March 2018
Accepted: 06 November 2018
DOI: 10.18466/cbayarfbe.410546

Abstract

In this study the protection ability of polyamines on Photosystem II (PSII), the proline content of the cells and necrosis formation induced by Ultraviolet-A irradiation were studied. Although 24 h Ultraviolet-A exposure decreased the photosynthetic quantum yield (Fv/Fm) ratio in Physcia semipinnata (J. F. Gmel.) Moberg, thalli that treated with 1 mM polyamine, especially spermidine and spermine, were slightly influenced by the UV-A exposure. It was also found that spd and spm treated samples had lower proline content than putrescine (put)-treated samples. Moreover, it was found that the samples that were treated with spd and spm had lower percentage of DNA damage than put-treated samples and non-treated groups. In this study, Ultraviolet-A induced DNA damage detected by acridine orange/ethidium bromide staining assay. The DNA damage in thalli of P. semipinnata was detected in situ in phycobiont layer after 24 h of Ultraviolet-A exposure, in contrast, polyamine-treated samples had a lower rate of necrotic cells.

Keywords: Polyamine, photosystem II, proline, Ultraviolet-A, lichen.

1. Introduction

After the discovery of the severe depletion of the stratospheric ozone, the studies increased about the potential effects of enhanced UV-A (315-400 nm) radiation on all living organisms. The detrimental effects of UV are varied and well known now. Some of these effects are reduction in photosynthesis, increase of growth rate and cell differentiation also damages in the protein contents of PSII and chlorophyll a synthesis, increase in lipid peroxidation, chlorophyll photobleaching, phycobiliprotein degradation, and direct damage on DNA. DNA lesions such as strand breaks are one of the most dangerous damage since they disrupt the linear structure of the DNA and prevent replication and gene expression. In addition, UV radiation generates oxygen and hydroxyl radicals that react with DNA to form monomeric photoproducts such as cytosine and thymine photohydrates and causes strand breaks and DNA-protein crosslinks. The most important toxic and mutagenic photoproducts produced as a result of UV-induced DNA damage is the cyclobutane dimmers. In response to such damages, the cell adapts itself by possessing several strategies such as photoreactivation, nucleotide excision repair and recombination repair in order to remove the photoproducts. A failure in repairing the damage could lead mutations or general deterioration of cell function.

UV radiation affects photosynthetic systems by two ways. It may cause a direct damage on key components such as D1 protein of PS II, so that the photosynthetic pigments are break down or by inhibiting the activation of RUBISCO [1].

Most organisms developed some strategies to avoid these UV-damages. Plants have efficient damage repair mechanisms, such as, rapid repair of the photosynthetic machinery and a variety of other UV-stress responses including UV stress proteins. Often, microalgae produce myosporin-like amino acids (MAAs), an effective UV sunscreen compound that absorbs harmful UV rays to prevent UV-induced damage [2]. Besides MAAs, other pigments such as carotenoids, scytonemines and usnic acids have important photoprotective role against UV damages. Especially carotenoids, scavenge toxic oxygen species formed in the chloroplast. Scytonemins are specific to cyanobacteria and cyanobacteria associated lichens while usnic acid is a lichen specific compound. Similar to MAAs, the synthesis of scytonemins and usnic acid are triggered by UV radiation since the main role of this compounds is UV protection.

Polyamines (PAs) are the other group of molecules play an important role in regulating the sensitivity and the tolerance of organisms under stress conditions. PAs are low-molecular weight polycationic compounds at
physiological pH. Their binding ability to negatively charged molecules, such as nucleic acids, proteins, and membrane phospholipids [3] protects these vital cell components against stress-induced damage. Diamine put, triamine spd and tetramine spm are the main PAs found in all living cells. Kramer et al. [4] demonstrated the important role polyamines in the plant protection mechanisms during UV-B radiation exposure. Kotzabasis et al. [5] also mentioned the similar findings of their study by reporting the association of spm, spd and put with light harvesting complex (LCH) and the PSII.

Lichen-like organisms exist about 600 million years when the UV-B radiation had reached to earth’s surface, and they must already then have developed efficient mechanisms for protection and repair. Studies on the UV radiation effects and the protection mechanisms against UV-induced stresses are increasing, our knowledge about avoidance mechanisms of lichens is still restricted. Furthermore, there is scarce information on other mitigation mechanisms and the role of external polyamines on cell damage of lichens induced by UV-A radiation.

First aim of the present study is to determine the time- and irradiance- relationships of UV-A damage on thalli of P. semipinnata. Second aim is to investigate if there is a protective feature of the polyamines on cell damage, regulation effect on the rate of photosynthetic quantum yield and proline accumulation.

2. Materials and Methods

2.1 Lichen Material

P. semipinnata samples were collected from the tree branches in Karagöl (Izmir-Turkey) (38°33’N 27°13’E, 840 m) in September. After collection, samples were transferred to the laboratory, cleaned from contaminants and dust by washing three times with distilled water. All the experiments, including UV-treatment conducted under light with intensity of 120 μmol m⁻² s⁻¹.

2.2 UV-A radiation treatment

Experiment consists of two parts. In the first experiment thalli were exposed to UV-A (Black Ray, 352 nm, 50 Hz, 0.60 Amps, model XX-20BLB), intensity J/cm², in petri dishes for 1 h, 2 h, 3 h, 5 h and 24 h without any treatments. UV-A lamps FSX24T12-UVB-HO, Philips) fixed 15 cm above the samples. Temperature was stabilized at 24°C and checked constantly to avoid heat effects. One thallus was not exposed to UV-A (as negative control).

2.3 Polyamine Treatment

After determining the most harmful UV-exposure duration the second part of the experiment carried out. In this experiment P. semipinnata thalli were incubated with 1 mM of polyamines, put, spd and spm for 30 minutes. After incubation, thalli were exposed to UV-A radiation in Petri dishes for 24 h.

2.4 Chlorophyll a fluorescence

Chlorophyll a fluorescence was measured with the plant efficiency analyzer (Handy PEA, Hansatech). Before measurement, Lichen thalli were dark-adapted for 15 min before measurements. The Fv/Fm (maximum quantum yield efficiency of PSII) parameters were calculated by the fluorescence induction curves of 5 seconds durations recorded at an irradiance of 1800 μmol m⁻²s⁻¹ from light emitting diodes. The Fv/Fm parameter was used as a stress indicator by determining the photosynthesis rate. Each treatment was repeated ten times.

2.5 Proline Analysis

Proline content was measured by modified method of Bates et al [6]. 0.5 mg tissue of lichen homogenized in 3% (v/v) sulfosalicylic acid and homogenate keep in laboratory condition for 24 h. The homogenates filtered through six layers of cheeseclothes. Extracts were treated with acid-ninhydrin at 90°C for an hour in water bath. After incubation the tubes transferred to ice boxes to end the reaction. After termination the solution extracted in toluene. Absorbance recorded at 520 nm. in spectrophotometer (Pharo 300, Merck). Absorbance values were calculated by using the standard curve.

2.6 Tissue section preparation

The thallus was chopped and fixed in %4 paraformaldehyde for overnight. The fixed samples were dehydrated in 5 different dilutions (50%, 70%, 80%, 95%, and 100%) of ethanol for 15 min. The dehydrated samples were dipped into ethanol-xylene (1:1) for 10 min and in 100% xylene for 10 min, respectively. The samples were embedded in paraffin, and sectioned in 5 μm coarseness via Leica RM 2145 microtome.

2.7 Fluorescence microscopy

Thallus sections were stained with acridine orange (AO)-ethidium bromide (EB) dye solution (0.01% (w/v) acridine orange and 0.01% (w/v) ethidium bromide dissolve in 10 mM PBS buffer, pH 7.0) for 10 min. Olympus BX-51 light microscope equipped with blue fluorescence filters used for microscopy and photographed with an Olympus C-5050 digital camera connected to microscope. Fluorescence activities of the cells were counted in 25 different areas of each section.

2.8 Statistical analysis

SPSS software (SPSS for Windows Version 11.0) were used for for statistical analyses. One-way analysis of variance (ANOVA) and Tukey’s pairwise comparisons were performed.
3. Results and Discussion

3.1 Long-term UV-A exposure disrupted photosystem II activity

To determine how UV-A exposure affects the PS II activity, we exposed the thalli to UV-A for different durations ranging from 1 hour to 24 h and then measured permanent reduction in maximal PSII efficiency (Fv/Fm) to determine the rate of photoinhibition. The Fv/Fm values were significantly lower (Table 1) for 5 h and 24 h UV-exposure, indicating the photodestructive effect of UV-A on PS II after prolonged exposure.

Table 1. The results of the photosynthetic quantum yield (Fv/Fm) of *P. semipinnata* thallus exposed to UV-A.

<table>
<thead>
<tr>
<th>n</th>
<th>Control X±SD</th>
<th>1 h X±SD</th>
<th>2 h X±SD</th>
<th>3 h X±SD</th>
<th>5 h X±SD</th>
<th>24 h X±SD</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fv/Fm</td>
<td>0.721±0.4</td>
<td>0.702±0.008</td>
<td>0.708±0.017</td>
<td>0.684±0.05</td>
<td>0.42±0.03</td>
</tr>
<tr>
<td>ANOVA</td>
<td></td>
<td>35.88</td>
<td>F probability</td>
<td>0.0024</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

n=number of replicates, x=mean values, SD=standard deviations.

In the next step, we decided to determine if polyamine treatment would protect the cells from photoinhibition by UV-A exposure. We treated the cells with 1 mM spd, spm, put or H2O and then exposed to UV-A for 24 h. Then, we measured the permanent reduction in maximal PSII efficiency (Fv/Fm). ANOVA and Tukey post hoc test indicated that, the Fv/Fm values were significantly lower in non-treated sample compared to polyamine-treated samples (Table 2), supporting a potential protective effect of polyamines.

Table 2. Photosynthetic quantum yield (Fv/Fm) values of *P. semipinnata* exposed to UV-A light (treated by polyamines or H2O) for 24 h and ANOVA results.

<table>
<thead>
<tr>
<th>Groups</th>
<th>n</th>
<th>Fv/Fm (X±SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>10</td>
<td>0.721±0.05</td>
</tr>
<tr>
<td>H2O</td>
<td>10</td>
<td>0.25±0.06</td>
</tr>
<tr>
<td>1 mM spd</td>
<td>10</td>
<td>0.664±0.08</td>
</tr>
<tr>
<td>1 mM spm</td>
<td>10</td>
<td>0.614±0.12</td>
</tr>
<tr>
<td>1 mM put</td>
<td>10</td>
<td>0.573±0.03</td>
</tr>
<tr>
<td>ANOVA</td>
<td></td>
<td>191.34</td>
</tr>
<tr>
<td>F probability</td>
<td>0.0001</td>
<td></td>
</tr>
</tbody>
</table>

n=number of replicates, x=mean values, SD=standard deviations.

3.2 Proline content did not differ significantly in response to UV-A exposure

The data showed that proline content was nearly the same in spd and spm-treated samples. It was also found that put-treated sample had higher proline content compared all groups (Figure 1).

3.3 Acridine orange/Ethidium bromide

AO is a nucleic acid-specific ultraviolet fluorochrome which stains DNA in bright yellowish-green on a typically green cytoplasmic background. And helps to detect the healthy chromosome structure. As seen in the Figure 2, strong AO signals were detected in the unstressed or lower stressed mycobiont and phycobiont cells of *P. semipinnata* implied that there was a very low DNA damage because AO tends to combine with relatively intact DNA molecules and produced a very strong signal.

EB mixture with low concentrations of AO identifies normal (AO at high, EB at low fluorescence levels) and late apoptotic/necrotic (AO at low, EB at high fluorescence levels) cells. The results we obtained with AO/EB staining revealed that number of necrotic cells significantly formed by the UV-A treatment and rate of necrotic cells of both photobiont and mycobiont layers significantly vary by the polyamine and UV-A treatments (Table 3). Both photobiont and mycobiont of *P. semipinnata* was stained as brilliant green in control, spd, spm and put-treated samples, while they were stained as brilliant orange in non-polyamine-treated samples (Figure 2). However, thallus section of the put-
treated sample showed that UV-A affected only upper surface part of lichen thallus, especially the mycobiont layer.

One of the responses which help plants to become more tolerant to unfavorable environmental conditions is the accumulation of low molecular-weight osmolytes such as proline and polyamines (PAs). Many studies suggest that biosynthesis of PAs may be a part of the plant response mechanism [3].

Many organisms have some strategies against harmful effects of UV stress. Some researchers believe that polyamines might be one of these strategies. Although the protective role of polyamines under different environmental conditions had been widely discussed, many opinions have been put forward on its accurate mechanism. Previous studies showed that treatment of plants with spd is partially prevented the harmful effects caused by UV-radiation [4, 7]. In another study, UV light has been shown as a put accumulation stimulator in cucumber [4]. Polyamines, especially the thylakoid-associated ones thought to have a constitutive role of photosynthetic apparatus protection. Many researchers pointed out that polyamine accumulation in cucumber leaves is an adaptive mechanism against the UV-radiation stress. Lütz et al. [8] demonstrated the change of polyamine content under UV-light treatment in tobacco. Out of this study, Kotzabasis et al. [5] demonstrated the relation between Light Harvesting Complex (LCH) and PSI and polyamines. In our previous study we showed the protective effects of exogenously added polyamines on Fv/Fm under UV-A stress in lichens [7]. Similarly, in our present work, Fv/Fm did not cause sharply decline in the samples that treated with spd and spm after UV-A radiation exposure for 24h. These results can be interpreted as that the polyamines protect the photosystem II from UV-A radiation (Table 1) [7]. The main question is if polyamines protect whole lichen thalli or only photobiont layer in lichen thalli under UV-A stress. In the present study, we have tried to understand that effect of polyamines on osmoprotectant production such as proline in whole thalli and protective role against cellular damage in both mycobiont and photobiont layer under UV-A radiation.

Proline has been assigned the role of a cytosolute, a storage compound or protective agent for cytoplasmic enzymes and cellular structure [9]. In addition, Alia et al. [10] reported that the UV-induced proline accumulation protects the plants against UV-promoted peroxidative stress. Therefore, the increased proline concentration may act as a hydroxyl radical and singlet oxygen scavenger [11]. However, in the present study, UV-A exposure was not significantly affected on proline accumulation in non-polyamine-treatment groups compared with the control group. Interestingly, polyamine-treated samples showed that proline content were significantly higher than control group, however, especially put significantly induced proline production compare to other polyamine-treated samples (Figure 1). Similarly, Wang et al [12] demonstrated the increase level of proline after spd or spm addition in Nymphaoides peltatum leaves under copper stress. Duan et al [13] also showed that exogenous spd addition to the medium increased proline amount in cucumber roots under salt stress. One suggestion about the relationship between polyamine and proline as a substrate-product complex [14]. Moreover, polyamines and proline synthesis were not only require common precursors which are glutamate, arginine, and ornithine, but also polyamine metabolisms could be related to expression of gene for proline biosynthetic enzymes [15]. Therefore, polyamines induced the accumulation of free proline to protect against oxidative damage [16]. Similarly, in the present study, exogenous polyamine induce proline accumulation, it could be help to protection from UV-A stress.

Photosynthetic organisms exposure to high level of UV radiation causes an oxidative stress resulting in chlorophyll degradation and necrosis. DNA is also one of the most notable targets of UV-radiation. Previous studies showed that UV-A-induced cytotoxicity and oxidative damage depend on radiation intensity and dose distribution [17]. In our previous study we also demonstrated the affected DNA template activity via DNA structural damage [18]. Cellular DNA damage occurs by both the direct and indirect effects of ultraviolet radiation. Indirectly, DNA strand break is induced by the attack of the bases via hydroxyl radical. Previous studies demonstrated the protection effect of polyamines on hydroxyl radical- induced DNA strand breaks [19]. At physiological pH, the spm, spd and put are found as polyammonium ions. Since polycations serve to stabilize active oxygen radicals DNA damage is inhibited via scavenging of these active radicals [19]. This hypothesis is supported by the significantly negative relationship between rate of necrotic cell and the external polyamine application in thalli of P. semipinnata (Table 3, Figure 2).

### Table 3. Necrotic cell rates of P. semipinnata exposed to UV-A light (treated with polyamines or H2O) for 24 h and results of the ANOVA between treatments.

<table>
<thead>
<tr>
<th>Groups</th>
<th>n</th>
<th>necrotic cells% X±SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>control</td>
<td>25</td>
<td>5.03±1.21</td>
</tr>
<tr>
<td>H2O</td>
<td>25</td>
<td>84.2±6.10</td>
</tr>
<tr>
<td>1 mM spd</td>
<td>25</td>
<td>15.3±0.61</td>
</tr>
<tr>
<td>1 mM spm</td>
<td>25</td>
<td>37.0±4.60</td>
</tr>
<tr>
<td>1 mM put</td>
<td>25</td>
<td>48.7±5.70</td>
</tr>
</tbody>
</table>

ANOVA

| F ratio        | 146.29 |
| F probability  | 0.000  |

x=mean values, SD=standard deviations.
Solheim et al. [20] mentioned the remarkable tolerance of lichens to extreme radiation conditions, on the contrary to terrestrial plants. Correlatively, short term (1 h, 2 h, and 3 h) UV-A exposure didn’t changed the Fv/Fm ratio in *P. semipinnata*, in our present study. However, when *P. semipinnata* thalli were exposed to UV-A irradiation for 5 h and 24 h, the results showed the highest damage. It is known that the maximum Fv/Fm rate is a sign of high light and radiation exposure [18]. For this reason, Fv/Fm rate is usually accepted as an indicator of photoinhibition. Under normal conditions, the Fv/Fm value of lichens varies between 0.45-0.65.

In the present study, 0.47 and 0.15 Fv/Fm values indicated the harmful effects of UV-A on photosystem II. These results indicate that the susceptibility or tolerance against UV-A radiation may depend on the exposure time. In the present study, fluorescence staining of *P. semipinnata* thalli demonstrated that UV-A radiation cause necrosis formation (Figure 2).

Relatively strong AO signals were detected in the unstressed or lower stressed mycobiont and phycobiont cells of *P. semipinnata* indicates a low degree of DNA damage. AO tends to bind incorrupt DNA strands with a strong signal. While EtBr with low concentrations of AO identifies normal and late apoptotic/necrotic cells. EtBr staining reflects the density of necrotic cells. Decreased AO staining may be related to destruction of DNA molecules into small fragments. In this study, the AO/EtBr staining process showed that the number of necrotic cells significantly modified by the UV-A treatment (Figure 2). However, the spd and spm-treated samples showed the lower percentage of necrotic cells than the untreated samples, in the mycobiont layer of untreated ones showed an increased level of necrotic cells at higher UV-A exposure.

4. Conclusion
In summary, we have analyzed the changes of photosynthetic efficiency, proline content, and necrosis in lichen thallus during UV-A stress and have demonstrated the protection of polyamines against UV-A induced oxidative damage. As a result, we demonstrated the increase of proline amount after external polyamine treatment (especially the put) in the cell during UV-A exposure. Since proline has an important role as a hydroxyl radical and singlet oxygen scavenger, the increased concentration of proline in *P. semipinnata* cells, might enhanced the tolerance against UV-A stress.

References


