Induced genetic variations in *Cuminum cyminum* through supplemental UV-B radiation

**Abstract**

**Aim**: The present study was conducted to investigate the genotoxic effect of UV-B radiation on morphological, biochemical and cytological parameters of *Cuminum cyminum*.

**Methodology**: The germinated roots of cumin were exposed to UV-B radiation at different intervals viz., 20 min, 40 min and 60 min, respectively, and some treated seeds were sown on pots for morphological and biochemical observation.

**Results**: Chromosomal studies divulged that UV-B radiation has substantial impact on Active Mitotic Index (AMI %). On increasing UV-B radiation dose, the chromosomal aberration rate elevated, thus AMI % moderately decreased and Total Abnormality percentage (TAB %) gradually increased. Different types of chromosomal abnormalities were ascertained, among which scattering was more prominent. The morphological observation showed that survival percentage and plant height decreased at elevated dose of UV-B. Biochemical results indicated that chlorophyll a, chlorophyll b and carotenoid contents decreased as compared to control, but proline content showed significant increment at higher doses of UV-B.

**Interpretation**: The observation elucidates that UV-B causes chromosomal aberrations during cell division and acts as a potent genotoxic agent for roots. Thus, it can be concluded from the above experiment that UV-B rays promote plant growth at low doses but at the higher doses, it subsequently hinders plant growth by damaging important constituents of plant cell.

**Key words**: Active mitotic index, Chromosomal abnormality, *Cuminum cyminum*, Total abnormality percentage, UV-B rays

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Introduction

Earth incorporates both ionizing and non-ionizing radiations. All living and non living organisms are inevitably exposed to these types of radiation. Ultra-violet rays are part of electromagnetic spectrum. Ultra-violet radiation is enhanced due to dissolution of protective ozone layer. Among three types of UV rays (UV-A, UV-B, UV-C), UV-B light (280 - 315 nm) is a natural component of sunlight, and due to its short wavelength, it has the highest energy of sunlight spectrum at the Earth’s surface (Jansen et al., 1998). Exposure of UV-B is increasing due to depletion of ozone layer linked to chlorofluorocarbons and breaking of \( \text{O}_2 \) structure by chlorine (Rowland, 2006). In plants, wide inter- and intraspecific differences have been reported in response to UV-B irradiation with respect to growth, production of dry matter and biochemical changes (Milchunas et al., 2004). Different species have different responses to the level of UV-B irradiation (Frohmeyer et al., 2003).

Strong absorption of UV-B photons by biologically important macromolecules (i.e., proteins and nucleic acids has a large effect on plant and animals metabolism (Caldwell et al., 2007; Heisier et al., 2003). The effects of frequent UV exposure on plants are reduced growth, photosynthetic activity and flowering (Teramura et al., 1991; Santos et al., 2004; Jansen and Bornman, 2012). Growth parameters like plant height and leaf area are significantly reduced in wheat, rice, maize, rye, soybean, sunflower, cucumber and spinach under high UV-B radiation (Chatterji, 2015; Mishra et al., 2014). Ultraviolet-C and B influences DNA through the production of cyclobutane pyrimidine dimers leading to mutations (McLennan, 1987).

Information on the action of UV-B radiation as well as of endogenous or exogenous antioxidant on the meristematic cells is very limited (Perennes et al., 1999; Potters et al., 2000). Cumin is an important spice crop of India, belonging to family Apiaceae. Cumin plant showed wide range of pharmacological activities including antimicrobial, antioxidant, anticancer, hypolipidemic, cardiovascular, central nervous, respiratory, immunological, anti-inflammatory, analgesic antipyretic and many other pharmacological effects (Al-Snafi, 2016). Reason behind preference of meristematic root tips for study is that treatments can be carried out in dark to avoid photoreactivation and due to low photolyase activity; root tips are prone to UV-B radiation. In view of the above, the present study was carried out to determine the exogenous effects of UV-B rays on root meristems of Cuminum cyminum.

Materials and Methods

**Plant Material:** Healthy Cumin seeds of variety GC 4 (Gujarat Cumin 4) were procured from Seed Spices Research Station S.D. Agriculture University, Jagudan Gujarat.

**UV-B treatment:** Seeds were surface sterilized in sodium hypochlorite for 10 min followed by washing with distilled water. Presoaked cumin seeds were allowed to germinate at 25°C in an incubator. Fully germinated cumin seeds of uniform size root tips were selected. Petriplates containing germinated seeds were placed inside UV chamber. One petriplate out of three was removed from UV chamber after 20 min. and subsequently other two were removed after 40, 60 min after which petriplates were left undisturbed for one hour for recovery. After treatment some germinated seeds were sown in pots and some were used for cytological study.

**Cytological analysis:** Irradiated roots were fixed in Carnoy’s fixative along with control sets. After 24 hrs, roots were preserved in 90% alcohol. UV-B irradiated roots were hydrolyzed in 1N HCl and then washed under running water to remove additional chemical and dried on blotting paper. Roots were stained using 2% aceticarmine, slides were prepared by squash technique and the photographs were taken under PCTV Vision Photography Software. Experiment was performed in three replicates along with control. Active Mitotic Index was calculated according to Edgar (1961) and Balog (1982). The morphological parameters were recorded, the survival percentage was taken after 21 days from sowing and plant height was taken after 45 days in centimeters.

**Biochemical analysis:** Photosynthetic pigments were analyzed by preparing leaf extract using 80% acetone and optical density of supernatant was taken at three different wavelengths (470 nm, 646 nm and 663 nm) following the procedure of Lichtenthaler and Wellburn (1983). Chlorophyll \( a \), Chlorophyll \( b \) and Carotenoids contents were computed there upon. The proline content was quantified according to Bates et al. (1973).

**Statistical analyses:** Data was analyzed using SPSS 16.0 software. One way analysis of variance (ANOVA) and Duncan’s Multiple Range Test (\( p \leq 0.05 \)) was performed and the graph was plotted by using Sigma plot 10.0 software.

**Results and Discussion**

Cytological studies of UV-B treated cumin seeds produced toxic effect on chromosomes. With respective increase in UV-B exposure on root meristems, AMI percentage decreased, while TAB % increased. Thus, AMI and TAB percentage showed inverse relationship with each other in UV-B treated sets. Highest AMI % was recorded in control sets 11.49± 0.18 which was reduced to 7.75 ± 0.44 at UV 60 min dose while the lowest TAB% was observed at UV 20 min dose i.e. 1.85± 0.04, which show elevation up to 6.93 ± 0.20 in UV-60 min sets (Table 1). Reduced AMI% may be due to inhibition of DNA synthesis at S phase that probably occurred due to decrease in ATP level and due to the pressure from the functioning of the energy production centre (Sudhakar et al., 2001). In-depth cytogenetic study of chromosomal morphology in case of control revealed that chromosome complement was \( 2n=14 \) as seen at equatorial plate in metaphase and 14:14 poleward migration was recorded at anaphase (Fig.1A, B). Various chromosomal anomalies viz.,

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scattering, stickiness, precocious movement of chromosome, orientation, laggard etc were detected at both metaphasic and anaphasic stages in treated sets (Fig. 1). The most dominant chromosome abnormality observed was scattering which may be due to the loss of microtubules of spindles fibers, unequal separation whereas orientation of chromosomes may be attributed to disturbance in spindle formation. (Fig.1C, J). Fig. 2a and 2b indicates graphical representation of metaphasic and anaphasic abnormalities which progressively increased along with exposure of UV-B radiation. Regular mitotic division was disturbed at higher doses of UV-B radiation resulting in C-metaphase and loop formation at metaphase. According to Levan (1938), C metaphase is caused due to inactivation of spindle fibres, followed by a random scattering of chromosomes over the cell. (Fig.1D). Looping of chromosome is mainly due to abnormal function of spindle which fails to attach with kinetochore resulting in improper chromosomal division. (Fig.1E). Unorientation (Fig.1F, K) may occur due to disruption of spindle structure or function as per UV-B radiation, leading to imbalance of spindle fibre on both the sides of centromere traction power or chromosome acentric fracture cannot cause normal movement of chromosome (Han et al., 2007).

Chromosome stickiness leads to inactivation of DNA replication, increased chromosomal contraction and condensation or nucleoproteins probably leading to cell death (Khanna and Sharma, 2013). (Fig.1G, H). Precocious movement of chromosomes observed at metaphase might be formed due to early chiasma terminalisation or univalent chromosome formation at the end of prophase (Kumar and Rai, 2007). Laggard chromosome may depend on the moving speed and process of an individual chromosomes differing from normal ones (Qian, 2004) (Fig.1I). According to Liu et al. (2015), the chromosome bridge may result due to enhanced activity of UV-B radiation, making chromosome breaks, and then the two chromosome sides are respectively healed, producing with double centromere chromosomes i.e., “chromosome bridges” (Fig.1J).

At low doses of UV-B treatment, plants were able to recover from harmful effects of UV-B rays by various means of DNA replication and DNA repair. But in case of higher dose, UV-B radiation breaks down the plant self-protection system and inhibits the action of cell DNA replication transcription and protein synthesis (Liu et al., 2015). DNA damage induced by UV-B radiations might have influenced the expression of number of genes leading to alterations in proteins that control many metabolic processes like plant development, cell cycle (Abdel Haliem et al., 2013). UV-B radiation can make cell DNA base mutate and induce pyrimidine dimer. Once dimers are formed in DNA, double chain of hydrogen bonds are damaged, and the normal DNA replication would not be performed. This could create chromosomal mutations which occur during cell division (Barnes et al., 1990). UV-B treated sets were morphologically analyzed on survival percentage and plant height parameters. The survival percentage and plant height of cumin plants decreased with increased exposure of UV-B radiation. Lowest survival percentage was recorded in UV-60 min i.e., 49.0±2.30% followed by 68.6±2.02% in UV-40 and 86.0±1.15% in UV-20 min whereas highest percentage was 95.0±0.57% in control sets (Fig. 3a). Maximum height was observed in control i.e., 25.59±1.57% followed by 21.85±1.34%, 15.78±0.66%, 9.61±1.03% in UV 20, 40 and 60 min, respectively (Fig. 3b).

UV-B rays causes reduction in survivability at higher dose of UV-B radiation which makes root growth shorter and bender. The growth of plant is reduced in response to UV-B treatment. This reduction is associated with UV-dependent destruction of the growth regulator indole-3-acetic acid (IAA) and formation of growth-inhibiting IAA photoproducts. Reduction in growth could be associated with UV-B induced inhibition in photosynthetic rate and destruction of growth promoting hormone indole acetic acid (IAA) (Kulandaiavelu et al., 1989). It was reported that UV radiation increases ethylene production and that this hormone decreases stem elongation and increases stem thickness (Ros and Tevini, 1995).

Biochemical analysis comprises of photosynthetic pigments quantification and proline estimation. Chlorophyll a, Chlorophyll b and Carotenoid contents were measured in UV-B treated sets. Chl a reduced from 4.24±0.06 (µg ml⁻¹ f. wt.) in control sets to 3.64±0.37 (µg ml⁻¹ f. wt.) in UV 20 min, 3.23±0.04 (µg ml⁻¹ f. wt.) in UV 40 min, 2.23±0.05 (µg ml⁻¹ f. wt.) in UV 60 min. Chl b decreased from 2.80±0.31 (µg ml⁻¹ f. wt.) in control to 2.63±0.27, 2.35±0.10, 1.35±0.13 (µg ml⁻¹ f. wt.) in UV 20, 40, 60 min, respectively. Highest carotenoid content was recorded 1.92±0.14 (µg ml⁻¹ f. wt.) in control while 1.56±0.06, 1.48±0.21, 0.73±0.16 (µg ml⁻¹ f. wt.) in UV-20, 40, 60 min, respectively (Fig.3c). The contents of chlorophyll a, b and total chlorophyll decreases when compared with control values with increasing UV-radiation levels (Salama et al., 2011). Chl a, Chl b and carotenoids are the major light harvesting pigments of photosynthesis and they are involved in synthesis of metabolites that promote the plant growth and development. However, increase in UV radiation damages the chloroplast which causes detrimental effects in various photosynthetic pigments. It has been reported that the concentration of Chl a and Chl b significantly decreased in Barleria obtuse and Vigna unguiculata plants that were exposed to UV radiation (Musil et al., 2002). Thus, chlorophyll content can be studied as one of the markers of cellular stress, and its decreased level can provide the evidence of severity of stress in plants (Ashraf and Harris, 2004).

Biosynthesis of carotenoids occur in cellular plastids where they are associated with light harvesting complexes (Kopsell and Kopsell, 2006). Therefore, carotenoid contents may be reduced due to damage caused by UV-B rays on light harvesting complexes. These compounds absorb UV-B before reaching to UV sensitive targets like chloroplast and other organelles, thus acting as solar shield. But when UV-B dose increases over threshold level such substances are unable to screen UV damage, and decrease in chlorophyll pigments indicated the damage of chloroplast caused by UV radiation.
Table 1: Metaphasic and Anaphasic abnormalities induced by UV-B radiation in root meristems of *Cuminum cyminum*

<table>
<thead>
<tr>
<th>Dose</th>
<th>AMI% (Mean±SE)</th>
<th>Metaphasic abnormalities (%)</th>
<th>Anaphasic abnormalities (%)</th>
<th>Oth</th>
<th>TAB%</th>
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<tr>
<td></td>
<td></td>
<td>Sc</td>
<td>St</td>
<td>Un</td>
<td>Pr</td>
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<tr>
<td>Control</td>
<td>11.49±0.18</td>
<td>-</td>
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<tr>
<td>UV-B radiation; 20 min</td>
<td>10.88±0.20</td>
<td>0.41±0.07</td>
<td>0.16±0.08</td>
<td>0.16±0.08</td>
<td>0.17±0.08</td>
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<tr>
<td>UV-B radiation; 40 min</td>
<td>9.12±0.64</td>
<td>0.67±0.17</td>
<td>0.38±0.09</td>
<td>0.17±0.08</td>
<td>0.28±0.02</td>
</tr>
<tr>
<td>UV-B radiation; 60 min</td>
<td>7.75±0.44</td>
<td>1.58±0.03</td>
<td>0.47±0.07</td>
<td>0.49±0.14</td>
<td>0.63±0.16</td>
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Values are mean of replicates ± S.E.; AMI- Active Mitotic Index, TAB- Total Abnormality Percentage; Sc- Scattering, St- Stickiness, Un- Unorientation, Pr- Precocious movement, Cm- C-metaphase, Lg- Laggard formation, Br- Bridge formation, Oth- Others; Means followed by lower case letter are statistically significant at p < 0.05 in Duncan’s Multiple Range Test (DMRT)
Fig. 1: UV-B induced chromosomal aberrations at mitotic stage. A. Normal metaphase (2n=14), B. Normal anaphase (14:14), C. Scattering at metaphase, D. C-metaphase, E. Loop formation, F. Unorientation at metaphase, G. Stickiness at metaphase, H. Stickiness at anaphase, I. 2 Laggards chromosome with one forward chromosome at anaphase, J. Unorientation with Scattering at anaphase, K. Unorientation at anaphase, L. Bridge formation [SCALE BAR = 10 µm].

Fig. 2: UV-B induced metaphasic abnormalities (a) and anaphasic abnormalities (b) in root meristems of Cuminum cyminum.
Proline content of control sets was recorded to be 0.28±0.04 µmol g⁻¹ f. wt., it elevated up to 1.47±0.13c µmol g⁻¹ f. wt. in UV-20 min, 3.20±0.10b µmol g⁻¹ f. wt. in UV-40 min and 3.97±0.07a µmol g⁻¹ f. wt. in UV-60 min treated sets (Fig. 3d).

Proline is a stress marker and play a critical role in protecting plants under stress. On increasing the duration of UV-B exposure proline content in plants is elevated to cope itself from the damages caused by radiation. Proline provides less than 5% of the total pool of free amino acids in plants under stress free condition; whereas the concentration increased up to 80% during stress (Matysik et al., 2002). According to Salama et al. (2011) increasing proline content is referred to as protective mechanism due to the generation of reactive oxygen species by UV radiation. Therefore, it can be concluded from present investigation that when plant exposed to UV –B rays for longer duration, adversely affect the plant growth while lower dose i.e., 20 min shows high AMI% and low TAB% chromosomal aberrations and very slight changes were observed morphologically and biochemically at lower dose as compared to control plants which may be significantly utilized by plant to adapt changing environmental conditions. Further studies can be done at meiotic level for the establishment of some beneficial high yielding traits in cumin through mutation breeding so that this economically and medicinally important crop might be able to withstand hazards of UV-B radiation in the coming years.

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References


