High temperature induced biochemical changes in pearl millet genotypes at seedling stage

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Abstract

Thirteen genotypes of Pearl millet (Pennisetum glaucum (L.) R. Br.) selected on the basis of heat tolerance were studied to see changes at biochemical and physiological levels. The plants were initially raised under normal condition in small plastic pots and 20 days old seedling were exposed at (40 °C for 4 hrs, 44 °C for 4 hrs and 46 °C for 2 hrs) in BOD incubator at Mandor to create the heat stress condition. After two days treatment data were recorded for MSI, chlorophyll content, antioxidative enzymes and MDA content. MSI, chlorophyll content and MDA decreased significantly due to heat stress while SOD, CAT and HB 17-6 performed better having antioxidative enzymes. The better performance of these genotypes may be due to high RWC, MSI, chlorophyll, SOD, CAT and low MDA content under high temperature stress.

Keywords: MSI, MDA, SOD, Pearl millet and Biochemical

Introduction

Pearl millet [Pennisetum glaucum (L.) R. Br.] is the staple food of majority of the poor and small land holders, as well as feed and fodder for livestock in rainfed regions of the country. In the semiarid tropics, inadequate seedling establishment due to heat stress can reduce

features, RNA species and proteins, and alters the effectiveness of enzymatic. Plants turn on enzymatic and non-enzymatic ROS scavenging systems to protect this ROS production. The main ROS scavenging enzymes are superoxide dismutase (SOD), catalase (CAT), peroxidase (POX), ascorbate peroxidase (APX), and glutathione reductase (GR), and the non-enzymatic systems include ascorbic acid (ASC) and glutathione (GSH) [5]. High-minded levels of these antioxidants are crucial for imparting thermo tolerance in plants [6]. Present investigation was conducted to study High temperature induced biochemical changes in Pearl millet genotypes at seedling stage.
Materials and Methods
The experiment was conducted at Mandor, Jodhpur to identify the heat tolerance mechanism in pearl millet at seedling stage. The pearl millet entries HB 17-6, HR 17-6, JMSB20143, JMSB 20101, J-2594, CZI 2010/11, CZI 2007/9, 411B, PPMI 1053, PPMI 1087, PPMI 1213, BIB 240 and BIB 238 used under this study. The plants were initially raised under normal condition in small plastic pots and 20 days old seedling were exposed at (40°C for 4 hrs, 44°C for 4 hrs and 46°C for 2 hrs) in BOD incubator to create the heat stress condition. All the measurements were taken after 2 days of treatment. Second fresh leaves were collected for the analysis.

Chlorophyll extraction
100 mg of finely cut fresh leaves were taken and grind with 10 ml of 80% acetone. It was then centrifuged at 5000 –10000 rpm for 5 mins. The supernatant was transferred. The absorbance of the solution was red at 645 nm and 663 nm.

Estimation of Chlorophyll content
The concentrations of chlorophyll a, chlorophyll b and total chlorophyll were calculated using the following equation:

\[
\text{Chlorophyll a: } 12.7(A_{663}) - 2.69(A_{645}) \\
\text{Chlorophyll b: } 22.9(A_{655}) - 4.68(A_{663}) \\
\text{Total Chlorophyll: } 20.2(A_{665}) + 8.02(A_{663})
\]

Membrane stability index
The procedure described by Premchandra et al. [8] modified and by Sairam [9] was used for calculating membrane stability index. Leaf samples (0.1 g) were placed in distilled water (10 ml). One set was kept at 40°C for 30 minutes and its conductivity of electrolytic leakage (C1) was recorded using conductivity meter. The second set was kept in boiling water bath (100°C) for 10 minutes and its conductivity (C2) was recorded after cooling at room temperature.

The MSI was calculated according to the formulae:

\[
\text{MSI} = \frac{(1 - C1/C2) \times 100}{C1/C2}
\]

Determination of Malondialdehyde
Determination of Malondialdehyde (MDA) concentration was determined by the method described by Heath and Packer [10]. Two hundred mg fresh leaf samples were extracted in 5.0 ml of 6% trichloroacetic acid (TCA) solution, centrifuged at 8000 rpm for 10 minute. Two ml of Thio Barbituric Acid (TBA) reagent was added in 1 ml of supernatant, mixed well and incubated for half an hour in a boiling water bath. Later the tubes were cooled to room temperature. The assay mixture was then centrifuged at 5000 rpm for 10 minutes. Supernatant bearing yellow to light orange colour was read on spectrophotometer at two wavelengths viz. 532 nm (major for MDA) and 600 nm (minor for interfering substance) millimolar concentration of MDA was calculated as follows: MDA (mM) = \((O.D.532 – O.D. 600) \times 155\) (extinction coefficient).

Antioxidant enzyme assays
Superoxide dismutase (SOD) assay was performed as per protocol of Dhindha et al. [11]. Leaf sample (0.5 g) was homogenised in 10 cm³ chilled 0.1 M potassium phosphate buffer (pH 7.5) containing 0.5 mM EDTA. The buffer was filtered through cheesecloth, and after centrifugation at 20,000 X g for 20 min, aliquots of the supernatant were used for enzymatic quantifying. The 3.0 cm³ reaction mixture contained 13 mM methionine, 25 mM nitroblue tetrazolium chloride (NBT), 0.1 mM EDTA, 50 mM phosphate buffer pH (7.8), 50 mM sodium bicarbonate and 0.1 cm³ enzyme extract. The reaction was started by adding 2 lμm riboflavin and placing the tubes below 2 X 15.00 W fluorescent lamp for 15 min. It was stopped by switching off the light and covering the tubes with black cloth. Tubes without enzyme develops maximum colour. A non-irradiated complete reaction mixture did not develop colour and served as blank. Absorbance was recorded at 560 nm, and one unit of enzyme activity was taken as that quantity of enzyme, which reduced the absorbance reading to 50% in comparison with the tubes lacking enzymes.

The catalase (CAT) activities were assayed as per the protocol of Chance and Maehly [12]. Samples were prepared by grinding 0.5 g fresh leaves in ice-cold 50 mM potassium phosphate buffer (pH 7.0) containing 0.1 mM ethylene diamine tetra acetie acid (EDTA) and 1% polyvinyl polypyrrolidone (PVP). The homogenate was filtered through four layers of cheese cloth and then centrifuged at 4 C for 20 min at 15,000X g. The supernatant was collected and an appropriate aliquot dilution of the crude extract was used for enzyme assays. CAT activity was measured by following the decomposition of H2O2 at 240 nm (ε = 39.4 mM-1 cm-1) in a reaction mixture containing 50 mM phosphate buffer (pH 7.0) and 15 mM H2O2. Enzyme activity was expressed as lμmol of H2O2 decomposed mg-1 (protein) min-1.

Results and Discussion
Pearl millet crop responds to high temperature stress in the form of changes in various physiological, biochemical processes. In the present study, with thirteen genotypes alter in small plastic pots to high temperature stress, physiological index (membrane stability index) have been observe when crop in the heat stress. Besides, key biochemical parameters were also measured (viz., MDA; superoxide dismutase (SOD), catalase (CAT), and chlorophyll. All these parameters helped in assessing tolerant versus susceptible genotypes at biochemical levels to heat stress. The values for stability of cellular membrane in the pearl millet genotypes revealed that there was decline in MSI percent of stressed plant in all genotypes. The MSI values varied from 77.40 to 87.99 percent on fresh weight basis at control 40 °C while under stress varied from 67.17 to 76.47 percent. MSI was high JMSB 20143 followed by J 2594, CZI 2010/11 and BIB 238 at control 40°C, whereas, HB 17-6,HR 17-6 followed by JMSB 20143 and PPMI 1213 high MSI was recorded at 46 °C (Figure-1). The difference in degree of lowering of MSI values was found significant when treatments were compared. Stress induced membrane damage has been biochemically marked by the presence of MDA as one of the thiobarbituric acid reducing substances (TBARS) that accumulates as a consequence of membrane lipid peroxidation. The MDA values in stressed plants were found higher over respective controls in all the genotypes. MDA content varied from 222 to 486 content μmol g-1 f.wt under control, while under stress it varied from 308 to 570 μmol g-1 f.wt. The MDA content was minimum increase in genotype BIB 238 with at par BIB 240 and PP1213 at 46 °C (Figure-2). Reactive oxygen species are known to damage cellular membranes by inducing lipid peroxidation [13]. Membrane stability index may be used as parameter to estimate the cellular injury caused to membrane due to peroxidation of fatty acids of the membrane. In present study, the increased levels of MDA in stress condition indicated the membrane sensitivity/membrane damage due to heat stress. Lower rate of increase of MDA in genotypes...
indicated better membrane strength. In the present study, the MSI reductions were found lowest in genotypes BIB 240 and PPMI 1053 indicating that they are putatively tolerant at 46 °C. Similar result were observed by other researchers [14-18]. Among many quaternary ammonium compounds is synthesized or found abundant mainly in chloroplast where it plays a vital role in adjustment and protection of thylakoid membrane thereby maintaining photosynthetic efficiency. The content of total chlorophyll varied from 0.71 to 1.63 mg g\(^{-1}\) fresh weight under control 40 °C while under 46 °C stress it varied from 0.47 to 1.09 mg g\(^{-1}\) at 46 °C (Figure-3). Plants must be protected from heat-induced oxidative stress so that they can survive under HT. Tolerance to HT stress in crop plants has been associated with an increase in antioxidative capacity [19, 20]. Tolerant plants entail a tendency of protection against the damaging effects of ROS with the synthesis of various enzymatic and nonenzymatic ROS scavenging and detoxification systems [21]. Activities of different antioxidant enzymes are temperature sensitive and activation occurs at different temperature ranges but the activities of these enzymes increase with increasing temperature. Catalase and Superoxide dismutase are the most important enzymes involved in regulation of intracellular level of H\(_2\)O\(_2\). Variability in increasing the activities of these antioxidants across wheat genotypes indicates their differential ability to acquire thermo-tolerance. Catalase activity was maximum in PPMI 1053, PPMI 1213 and HB 17-6 at 46 °C (Figure-4). SOD is usually considered as the first line of defence against oxidative stress. In present study, the activities of catalase and superoxide dismutase enhanced with variable magnitude under heat stress conditions. Among genotypes, maximum percent increase in catalase and superoxide dismutase were recorded in PPMI 1053, PPMI 1087 and HB 17-6 (Figure-5) under high temperature stress conditions. The temperature until which increased activities are maintained varies in the tolerant and susceptible varieties. In the tolerant varieties, they could maintain increased activities at HT in comparison to the susceptible ones [22].

**Fig 1:** Effect of high temperature stress on membrane stability index in pearl millet genotypes. Values of ± SE

**Fig 2:** Effect of high temperature stress on Malondialdehyde (MDA) content in pearl millet genotypes. Values of ± SE
Fig 3: Effect of high temperature stress on Chlorophyll content in pearl millet genotypes. Values of ± SE

Fig 4: Effect of high temperature stress on antioxidant enzyme catalase in pearl millet genotypes. Values of ± SE

Fig 5: Effect of high temperature stress on activity of antioxidant enzyme Superoxide dismutase in pearl millet genotypes. Values of ± SE
Conclusion
High temperature stress is one of the most prevalent types of abiotic stress which directly induces plant desiccation. The plants occurrence strict osmotic stress and production of ROS, which damages the nucleic acid and protein structures. The results of present study reveal that tolerance mechanism for heat stress exists in pearl millet genotypes for a variable extent. On the basis of various parameters analyzed in this study, PPMI 1053, PPMI 1087 and HB 17-6 have been identified as heat tolerant and BIB 238 and BIB 240 as heat susceptible. It is suggested that these heat tolerant genotypes can be used in future breeding programme and information can be used for molecular analysis for study towards the development of a heat tolerant pearl millet hybrids.

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