Physio-biochemical and Molecular Responses of Mung Bean (Vigna radiata L. Wilczek) to Salt Stress

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A B S T R A C T

Soil salinity kindles a cascade of events in the normal functioning of the plant. Plant experience changes in the morphological, physiological and biochemical aspects in response to salt stress. In the current study, several experiments pertaining to the changes due to salt stress has been studied in Mung bean. Morphologically, a significant decrease in fresh weight of roots (1.89 folds) was observed. About 3.27 and 3 fold decrease in the Relative Water Content and Chlorophyll levels, respectively were observed in 300mM NaCl treated and control mung bean plants. Proline level increased to 2.69 folds at 24 hrs in stress imposed plants compared to control. Molecular level changes due to salt stress was analyzed by Semi-RT-qPCR analysis. Total RNA was isolated from the control and salt-stressed samples and cDNA synthesized. Expression analysis of Late embryogenesis abundant (LEA) protein coding genes, VrLEA13 and VrLEA26 revealed amplicons of around 190 bp and 200 bp, respectively. This study creates an insight into the mechanism with which mung bean responds to salt stress by highlighting the significant variation in morpho, physio-biochemical and molecular responses.

Keywords: Mung bean, Salt stress, Biochemical analysis, Molecular analysis, LEA proteins

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Introduction

Pulses are rich sources of protein and hence are rightly pronounced as “Poor man's meat” and “rich man's vegetables”. Vigna radiata L. (Green gram, Mung bean) is an important food legume crops native to India belonging to the family Fabaceae. It is a self-pollinated and a diploid crop (2n=2x=22) with a genome size of 494-579Mbp. The short life span and nitrogen-fixing ability in the soil makes mung bean a valuable crop in most cropping systems (Somta and Srinives, 2007). Salinity stress is one of the most common abiotic stresses in the arid and semi-arid regions (Dutta et al., 2018). Salinity acts as a major environmental constraint that hampers the growth, development, and yield of the plant by altering the morphological, physiological, and biochemical attributes (Taji et al., 2002, Morant-Monceau et al., 2004 and Kandil et al., 2012). The first and foremost effects of salinity stress in pulses are reduction in root length, shoot length, germination percentage,
fresh and dry weight of shoot and root, and chlorophyll content (Abdul-Baki and Anderson, 1973; Magwanga et al., 2018). Saline stress also induces oxidative stress resulting in the generation of Reactive Oxygen Species (ROS), which shows its mischievous effect on the intracellular structure and the cell membrane (Mudgal et al., 2010). Mung bean is known to be a salt-sensitive legume crop (Quddus et al., 2012). Under hydroponic salt stress condition, mung bean’s root length, shoot length, fresh weight, dry weight, and chlorophyll content were diminished when compared to control (El-Kafafi, 2015), whereas the proline and peroxidase content was increased (Ghosh et al., 2015). In Spinach, the total soluble sugar (TSS) content increased as a result of an increase in the salt concentration in the shooting medium (muchate et al., 2019). The present research focused on morphological, physiological, and biochemical adaptation of mung bean in high saline stress conditions. There are several proteins and transcription factors that aid the plants in overcoming abiotic stresses. LEA belongs to a class of proteins that helps the plant in battling stresses like temperature, drought, and salinity. Considering the above antagonistic impact of salinity stress on mung bean, it is important to develop a salt lenient assortment of mung bean which can be used by the farmers for cultivating under high saline conditions.

Materials and Methods

Experimental materials

Seeds of Mung bean variety CO (GG)-8 were collected from the Department of Pulses, Tamil Nadu Agricultural University, Coimbatore. This variety is a popular short duration ruling variety with moderate resistance to Mung Bean Yellow Mosaic Virus.

Imposition of salt stress and Sample collection

The seeds were sown in non-drainage plastic pots (22.8cm X 17.7cm) containing a sterilized mix of red and clay soil (1:1). After the emergence around two-leaf stage, the seedlings were irrigated with different NaCl concentrations viz. 0mM, 100mM, 150mM, 200mM, 250mM, and 300mM NaCl. The pH and Electrical Conductivity (EC) were checked. The samples were collected in different time intervals viz., 0th hrs (Plant with no exposure of stress), 3rd hr, 6th hr, 9th hr, 12th hr, and 24th hr, after which various physiological, biochemical, and molecular analyses were carried out.

Morphological parameter analysis

Followed by the imposition of salt stress, the mung bean seedlings were randomly selected from each treatment and washed thoroughly with distilled water. The length of the root, shoot, and leaf was measured in cm, while the fresh and dry weights of the root and shoot were measured in grams.

Seedling vigor index

Seedling vigor index was calculated by using the below formula as reported by Abdul-Baki et al., 1973 expressed in whole number.

\[
SVI = \frac{(Average\ plumule\ length + Average\ Radical\ length)}{Germiation\ percentage}
\]

Relative Water Content (RWC)

The Relative water content estimation was done by Barrs and Weatherley(1962) method. Leaves from control and stress imposed plants were collected under different time interval. Initially, the fresh weight of the leaves was taken followed by the turgid weight by plating the leaves in Petri dishes containing distilled water soaking leaves for 6 hours. The samples
were kept in a hot air oven at 70°C overnight and the dry weight was noted. The RWC was recorded using the formula

$$\text{RWC(\%)} = \frac{F_w - D_w}{T_w - D_w} \times 100$$

Where,

"Fw" is the fresh weight, 
"Dw" is the Dry weight 
"Tw" is the turgid weight and 
"RWC" is expressed in percentage (%)

**Determination of chlorophyll content**

The Chlorophyll content determination was done by the Arnon method (1949). Fresh leaf samples were harvested from control and treated plants and ground using 80% chilled acetone and centrifuged at 3000 rpm for 10 min. The supernatant was made up to a volume of 25 ml with 80% acetone. The color absorbance of the solution was estimated for chlorophyll b at 645 nm for chlorophyll at 663 nm. The total chlorophyll content was measured using the formula,

$$\text{Chlorophyll content} = \frac{(20.2 \times A_{645}) + (8.02 \times A_{663})}{1000 \times W} \times V$$

Where,

"W" is the fresh weight of the material 
"V" is the extraction volume and it is expressed in terms of µg g⁻¹ fresh tissue

**Estimation of Peroxidase content**

The Peroxidase content was estimated by Sumner and Gjessing method (1943). Fresh leaf material (0.5g) harvested from control and NaCl treated samples were homogenized with ice-cold phosphate buffer (pH-6.5) and centrifuged at 10,000 rpm for 20 min. The supernatant was taken and used for enzyme source. Each reaction mixture containing 1 ml of ortho-dianisidine, 0.5 ml of H₂O₂, 1 ml of phosphate, 2.4 ml of distilled water, and the tubes were incubated at 30°C. The reaction was stopped by adding 1 ml of H₂SO₄. The absorbance was recorded at 430 nm by using GENESYS™ 40/50 Vis/UV-Visible spectrophotometer and it was expressed in terms of µM g⁻¹ min⁻¹.

**Estimation of Proline content**

Proline content was estimated according to the method of Bates (1973). Plant materials from control and treated samples were ground in 10 ml of 3% sulpho-salicylic acid and centrifuged at 3000 rpm for 10 min. The supernatant was taken to this 2 ml of acid ninhydrin (1.25 g of ninhydrin in 30 ml of warm glacial acetic acid), 2 ml of glacial acetic acid, 2 ml of 6M orthophosphoric acid were added and the tubes were heated in a water bath (65°C) for one hour. The reaction was terminated by placing the tubes in an ice bath, followed by 4 ml of toluene was added to the test tube and mixed vigorously for 30 sec. The absorbance was recorded at 520 nm against a standard curve prepared using a known concentration of proline. The proline content was calculated by using the following formula and expressed in terms of µg/g.

$$\mu\text{mol proline g}^{-1}\text{ fresh weight} = \frac{(\mu\text{g proline ml}^{-1}\times\text{ml of toluene/115.5})}{(\text{g of sample})}$$

**Estimation of Total soluble sugars**

Total soluble sugar was estimated according to the method Yemm and Willis (1954). 100mg fresh leaf tissue from control and NaCl treated samples were homogenized with 10ml of 80% ethanol followed by centrifugation. The supernatant was collected and the final volume of supernatant was made up to 10 ml with ethanol (80%). 1ml of sample was taken for analysis along with 1ml
of distilled water and 4ml of anthrone reagent was added and kept in the water bath for 1 min at 70°C. For an exothermic reaction, the test tubes were cooled in the air. The absorbance was recorded at 620nm on GENESYS™ 40/50 Vis/UV-Visible spectrophotometer against a standard curve prepared using a known concentration of D-glucose and expressed in terms of mg g⁻¹ fw⁻¹.

Estimation of Ascorbic acid

The Ascorbic acid determination was done by the Omaye method (1979).0.5 g of leaf tissue from control and stress imposed samples were homogenized with 5% tricarboxylic acid and centrifuged at 3500 rpm for 20 min followed by the supernatant and was mixed with 0.2 ml of DTCS reagent (3 g of 2, 4 DNPH, 0.4 g thiourea and 0.05 g of CuSO₄ were dissolved in 9N sulphuric acid and made up to the volume 100ml) incubated at 37°C for 3 hours followed by 1.5 ml of ice-cold sulphuric acid was added and the solution was kept at room temperature for 30 min. The absorbance was recorded at 540nm GENESYS™ 40/50 Vis/UV-Visible spectrophotometer and the results were expressed as µg/mg of protein.

RNA isolation and Semi-quantitative reverse transcriptase (Semi-qRT-PCR) analysis

Total RNA was extracted with TRIzol reagent, the concentration and quality of the RNA samples were assessed using a nanodrop (Genway Nano, Cole-Parmer, UK) and Agarose gel electrophoresis. To remove DNA contamination from RNA, DNaseI treatment (Promega) was carried out with the isolated RNA. The first-strand cDNA was synthesized using Thermo scientific cDNA synthesis kit. The VrLEA 13, VrLEA 26 genes were amplified by PCR using first-strand cDNA isolated from the seedling of salt-stressed CO (GG) – 8 variety.

Results and Discussion

Effect of NaCl stress on the plant growth parameters

Our experiment showed that there was a significant reduction in the root, shoot, leaf length and leaf area with an increasing NaCl concentration over the control (Fig 1). A reduction in fresh weight and dry weight of roots was observed in the salt-stressed plants when compared with the water control (Fig 2a). In the case of shoot fresh as well as dry weight, there was no evident change observed between the treated and control plants (Fig 2b).

Leaf fresh weight increased only at moderate salinities but decreased significantly at high salt levels like 250 and 300mM NaCl (Fig2c).Total leaf area per plant remained unchanged even up to 100mM NaCl condition. On the other hand, beyond 100 mM NaCl leaf area remained more or less stable (Fig3).The highest value of seedling vigor index (3870) was recorded with the 100mM NaCl treatment. The lowest seedling vigor index was recorded from Control (2825) (Fig 4).

Effect of NaCl stress on physio-biochemical indices

Relative water content

Relative water content is a very sensitive parameter of plant development which is impaired at slightly severe stress. At 24 hours, the plant was subjected to 100mM NaCl stress showed 0.46 fold decreases in the relative water control levels when compared to the control, whereas the plant subjected to 300mM NaCl shows a drastic decrease of 3.27 folds when compared to the control. This indicates that with increasing levels of NaCl there is a decline in the RWC levels (Fig5).
Chlorophyll content

Leaf chlorophyll content decreased progressively with increasing salt levels. For control there were 3 fold decrease in the chlorophyll content in 300mM NaCl treated at seedling stage for 24hrs (Fig6).

Peroxidase activity

A similar trend was also observed after NaCl stress imposition, where the peroxidase activity in the seedling was drastically elevated (2.78 times) at 300 mM NaCl comparative to control (Fig 7).

Proline content

In our experiment with an increase in salinity a significant enhancement in proline activity, 2.69 fold increase in the control seedling (Fig8) was observed.

Total soluble sugar content

Total soluble sugars were gathered in moderately higher quantities at the increased salt level. In our experiment when compared to 0 mM NaCl at 0 hrs and 300mM NaCl at 24 hrs resulted in a 4.6 times increase in the leaf total soluble sugar content over water control (Fig 9).

Ascorbic acid activity

NaCl treatment for 9 hrs (100 and 150mM NaCl) showed significant increases in the ascorbic acid content when compared to 0hrs and 24 hrs (Fig10).
Fig. 3 Impact of salinity stress on leaf area of mung bean seedling

Fig. 4 Average of seedling vigor index as affected by various concentration salinity stress of mung bean seedling
Fig. 5 Effect of NaCl stress on Relative water content of mung bean seedling under different time intervals

Fig. 6 Effect of salinity stress on the total chlorophyll content of mung bean seedling under various time intervals

Fig. 7 Impact of salinity stress on the accumulation of peroxidase content of mung bean seedling under different time intervals
**Fig. 8** Effect of salinity stress on the accumulation of total proline content of mung bean seedling under various time intervals

**Fig. 9** Impact of salinity stress on the accumulation of total soluble sugar content of mung bean seedling under different time intervals

**Fig. 10** Impact of salinity stress on the accumulation of Ascorbic acid content of mung bean seedling under different time intervals
**Fig. 11** Total RNA profile of mung bean from control and stress imposed plants resolved on 1% Agarose gel; L-100 bp ladder; 1 - RNA from control plant; 2 to 6 - RNA from salt-treated plants

**Fig. 12** PCR amplification of mungbean cDNA for *VrLEA* 13 gene resolved on 1% agarose gel; L-100 bp DNA ladder; 1 - Negative control; 2 to 4 - PCR amplified product of cDNA with *VrLEA*13 gene

**Fig. 13** PCR amplification of mungbean cDNA for *VrLEA* 26 gene resolved on 1% Agarose gel; L- 100bp DNA ladder; 1 - Negative control; 2 to 4 - PCR amplified product PCR of cDNA with *VrLEA* 26 gene

Expression analysis of *VrLEA*13 and *VrLEA*26 genes

Distinct rRNA banding patterns were
observed in the total RNA profile of salt control and stressed samples (Fig11). Expression analysis revealed that the amplicon size of the genes VrLEA13 and VrLEA26 were found to be around 190 bp and 200 bp respectively (Fig.12 & Fig 13).

**Stress induced altered morphological indices of mung bean**

The study showed a significant reduction in the root length with an increasing level of NaCl, whereas the shoot and leaf lengths remain unaffected. The same reduction of this root, shoot as well as leaf length was observed by Ghosh et al., 2015. Salt stress caused lower intra-cellular water potential and water shortage around the root zone because of which roots neglected to uptake adequate water and supplements for satisfactory plant development (Kumar et al., 2012). Reduction in biomass under salinity was observed in perennial grass Cynodon dactylon (Hameed et al., 2008). Hence salt stress reduces the fresh weight and dry weight of mungbean seedling was confirmed in the present experiment. It has already been reported that a decrease in leaf area with the induction of salt stress in Cynodon dactylon. In our experiment, leaf area decreased with increasing NaCl concentration. This shows that salt stress highly affects the leaf area. It has already been reported that there was a decrease in seedling vigor index with the induction of salt stress in Brassica napus (Kandil et al., 2012). The present study experiment showed that the vigor index had increased by adding 100mM NaCl when compare to other salt concentration.

**Effect of salt stress on physio-biochemical parameters**

Salt stress causes membrane damage, ion toxicity, decrease the hydrolytic enzyme activity and increase the lipid peroxidation level, and may stimulate the ROS production level. In this study, the variety showed a decrease in leaf water potential with increasing the salt concentration at 300mM. The same reduction in the leaf water potential was observed by Naz and Bano, 2013 at 200mM with the increase in salt stress. It has been reported that the mung bean plants developed under salinity stress demonstrated excess accumulation of leaf Na⁺ and Cl⁻ bringing about increased ROS production and diminished photosynthetic activity and plant development (Panda et al., 2009). Our study showed a significant reduction in the chlorophyll content with increased level of NaCl. The same reduction in the chlorophyll content with the increase in the salt stress was also observed by Roychoudhury et al., 2013.

During salt stress, peroxidase serves as a signal molecule plays an important role in plant defense mechanism. Hydrogen peroxide (H₂O₂) substance is used as a stress marker for oxidative stress and is commonly influenced by abiotic stress. Salt stress increases the peroxidase content in stressed plants of mungbean in the present study. This observation corroborates findings of Dutta et al., 2018.

Under salinity stress, osmolyte like proline maintains cellular equilibrium through diffusion regulation and induces physiological process favorably⁹. The present study experiment showed that the leaf proline content increased consistently with increased salt concentration. Hence, the same accumulation of proline content increased uniformly under salt stress in mung bean was observed by Ghosh et al., ⁹. Total soluble sugars are mainly glucose and fructose was accumulated in relatively higher quantities in high salt level. A strong correlation between sugar accumulation and salt-tolerant has been earlier been reported in
many species (Sumithra et al., 2006). Hence salt stresses significantly increase the total soluble sugar was shown in our experiment. Ascorbic acid is a significant antioxidant agent in photosynthetic and non-photosynthetic tissues which responded straightforwardly with ROS in photosynthetic and non-photosynthetic tissues. The defensive role of ascorbic acid in plant cells from the unfavorable impact of salt stress was depicted by Yang and Guo, 2018.

LEA proteins are a major class of stress associated proteins with multiple roles in abiotic stress tolerance. Highly intrinsically disordered structures of dehydrin group of proteins under stress is reported in Vitisa murensis VamDHN3 (Xu et al., 2020) and several other plant species. Earlier studies from our group have revealed the possible function of Mungbean LEA proteins as a DNA binding protein with its hydrophilic nature similar to other Group 1 LEA proteins (Rajesh and Manickam, 2006). The present study indicated that two of the VrLEA genes expressed during the salt stress could possibly play a protective role during salinity stress.

In conclusion from this present study, it tends to be inferred that the utilization of salt unfavorably influenced the development as well as the metabolism of mungbean seedlings. High concentrations of salt induces stunted growth and loss in chlorophyll content and reduced the elongation of the leaf as well as oxidative damages by altering the antioxidant machinery leading to membrane damage through lipid peroxidation increasing the activity of proline, peroxidase and total soluble sugars in mung bean seedlings under different time intervals. Higher expression of VrLEA 13, VrLEA 26 gene in mung bean plants was observed when imposed with salt stress. Since few studies have simulated the conditions of following salt stress in the plant ecosystem, more studies are needed in the future to understand the effect of salt stress.

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