Molecular characterization and expression studies of *Eucalyptus globulus* stress-responsive gene *DHN-10*

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**Abstract**

The *Eucalyptus globulus* is one of the important forest tree species which reveals enhanced tolerance to multiple abiotic stresses. The enhanced tolerance of *E. globulus* against low temperature is linked to the presence of enhanced accumulation of lysine-rich dehydrin proteins. The accumulation of dehydrin protein of Mr 10 kDa in response to multiple abiotic stresses prevents cells from dehydration. It is encoded by a novel dehydrin-10 (*DHN-10*) gene. In the present study, we have used the *DHN-10* gene of *E. globulus* of Pakistani origin, which was cloned in a bacterial expression vector pET30(a) and sequenced. We have found the 315 bp long coding sequence of this gene which has been enrolled as *Eucalyptus GZJ-2018 DHN-10* gene under GenBank accession number MG948256.1. The *in silico* studies have identified several differences of this gene from the earlier enrolled *DHN-10* genes: the studied gene possessed higher amphipathic character because of the presence of five additional electrically charged amino acids (two positively charged histidine, one negatively charged glutamate, and two negatively charged aspartate residues) and one extra lysine residue. The studied *DHN-10* gene has been successfully expressed in the BL21-DE3 expression strain of *Escherichia coli* and 10 kDa protein has been detected on the nitrocellulose membrane. Our study is the first report of the sub-cloning of the *DHN-10* gene and its expression outside the *Eucalyptus* cell.

**Keywords**: *Escherichia coli* expression strain, *Eucalyptus* dehydrin proteins, *in silico* analysis, sub-cloning.

**Introduction**

Drought, salinity, and frost result in dehydration of plant cells. The dehydration signals in plants trigger the synthesis of dehydration-induced cellular proteins (dehydrins) which were first observed in maize and barley in 1989 as reported in Yao et al. (2005) and Ali et al. (2014, 2016). Dehydrins belong to a multi-protein family called late embryogenesis abundant (LEA) proteins group-2 (Puhakainen et al. 2004, Yang et al. 2012). The dehydrins are located in the cytoplasm, nucleus, mitochondria, and chloroplast (Xu et al. 2005). Further, dehydrins have been found in the endoplasmic reticulum, plasma membrane, and tonoplasts (Close et al. 1993, Close 1996). The extensive accumulation of dehydrins has been observed in the plant embryos during later developmental stages, just like other LEA proteins (Hanin et al. 2011). The dehydrins accumulate extensively in all vegetative tissues when plants are subjected to environmental stresses that may cause the dehydration of cells like osmotic stress, drought, salinity, and heat (Hanin et al. 2011).

The dehydrins are hydrophilic and thermostable...
proteins with high content of electrically charged amino acids. Another characteristic of all dehydrins is that they possess a lysine-rich conserved sequence of 15 amino acids which is designated as the “K segment” (Saavedra et al. 2006, Graether and Boddington 2014). When the cellular hydration becomes low, the K-segments of dehydrin proteins adopt α-helical structural conformation. The alterations in structural conformation of dehydrin proteins result in the altered function of these proteins. The phenomenon of structural alterations is known as “moonlighting” and it is the main characteristic of intrinsically disordered/unstructured proteins (IDPs/IUPs). When IDPs/IUPs perceive signals of changes of hydration (availability/deficiency) in a microenvironment, they undergo conformational changes leading to functional changes. The α-helices of dehydrin proteins (designated as amphipathic α-helices) have been found important for developing interactions with dehydrated surfaces of biomembranes and other cellular proteins. When the dehydrin molecule is in α-helical conformation then its numerous K-segments can form bundles. This feature enhances the interaction of dehydrin with biomembranes and other cellular proteins which enhances the stress tolerance of cells (Hanin et al. 2011). Dehydrins can also be exploited for the management of excessive metal ions and reactive oxygen species (ROS) in the cells under stressful conditions. Because some of the dehydrin proteins have been discovered to contain a relatively higher content of histidine/arginine or other reactive amino acids on the surface, they have also been reported for possessing properties to bind with metal ions and scavenge the ROS or reactive oxygen species in the cells under stress conditions (Hanin et al. 2011).

Eucalyptus is an important forest species with enhanced tolerance against abiotic stresses. The dehydrin-10 protein of Eucalyptus of 10 kDa (DHN-10), has been reported responsible along with other two dehydrin proteins (dehydrin-1 and dehydrin-2) for conferring abiotic stress tolerance to Eucalyptus globulus (Fernandez et al. 2012). Dehydrin-10 has still not been well studied for its manipulation purposes and similar expression outside of the Eucalyptus cells. The natural expression of dehydrin proteins is abscisic acid-dependent as well as independent of its normal functioning in the cells. The aim of the presented study was a detailed molecular characterization of dehydrin protein of Eucalyptus globulus and DHN-10 expression in Escherichia coli strain BL21-DE3.

Materials and methods

Plants and isolation of DHN-10 gene: For nucleic acid extraction, the plant tissues were obtained from Eucalyptus globulus trees of saline and water logged areas of Lahore Division (Punjab). The RNA and DNA were isolated from different tissues like fleshy green twig tissues, woody green twig tissue, hard bark, older hard leaves, young soft leaves and young floral buds of apparently older trees as described by Jahangir et al. (2018). The full-length DHN-10, as well as its coding regions (mRNA through cDNA), were amplified using gene specific forward and reverse primers; and cloned via thymine-adenine (TA) cloning of PCR products (TA cloned; Jahangir et al. 2018).

Sequencing and in silico analyses: The TA cloned full-length and coding region of the Eucalyptus globulus gene were sequenced using commercial sequencing facility of Centre for Applied Molecular Biology (CAMB, Lahore, Pakistan). The detected gene sequences were manually analyzed using Chromas v. 1.45 software (http://technelysium.com.au/wp/chromas/). Verification of analyzed sequences was done at NCBI through BLASTn (http://www.ncbi.nlm.nih.gov/BLAST/) by comparing with reported sequences of Eucalyptus DHN-10. Using ExPaSy translating tool (www.expasy.org), the FASTA format of DNA files was translated into amino acid sequences. Further bioinformatic analysis was done using different tools including multiple sequence alignment assay (MSA), UniProt, and ProtParam (MSA performed on Uniprot.org and physiochemical analysis was done on ExPaSy ProtParam tool).

Sub-cloning of DHN-10 gene in pET30(a): The DHN-10 has not been reported earlier to be manipulated for gene expression studies. The coding region of DHN-10 was cloned into E. coli expression vector, pET30(a), to identify the possibility of expression of DHN-10 outside the Eucalyptus cells. The intron-less, central coding region was amplified from TA cloned full-length gene with primers having restriction sites of EcoR1 and HindIII enzymes. The primers were designed using Primer3 software, named as C-1 primers with the following sequences: forward 5'-GAAGCTTATGGCGGGAATCATCCACAAGATC-3′ and reverse 5'-AAGCTTACGCTATCAGTGGCGCAGT-3′. The optimized PCR formulation contained 100 ng plasmid template (TA cloned full length Eucalyptus DHN-10 gene was used as template during its amplification which had central single exon with two introns on both sides), 0.001 nmol of both forward and reverse primers, 1× PCR master mix (Thermo Fisher Scientific Baltics, Vilnius, Lithuania), and nuclease-free water to a final volume of 30 mm³. The optimized cycling conditions were initial denaturation at 95 °C for 5 min, then 95 °C for 45 s, 54 °C for 45 s, 72 °C for 1 min, and final extension at 72 °C for 7 min in Kyurate thermal cycler (Queensland, Australia) super cycler (SC300G) with 30 repeats. The amplified coding sequence of DHN-10 with added restriction sites for EcoR1 and HindIII restriction enzymes on upstream and downstream edges, respectively, was TA cloned into TA cloned plasmid template following the instructions of InsTA PCR cloning kit. The TA cloned construct (named C-1 construct) was transformed into competent E. coli strain Top10F following the previously optimized protocols (Jahangir et al. 2018). Successful ligation of the amplicon was assessed by restriction digestion and PCR amplification of purified plasmid from true transformants. The tested pTZ57R/T plasmid having C-1 construct and empty pET30(a) were double digested with EcoR1 and HindIII enzyme pair. The released insert from construct and linearized pET30(a)
having cut ends were purified with *Quick Gel Elusion* kit (Invitrogen, Carlsbad, USA). The purified gene insert of C-1 with cut ends was ligated into the purified, linear, empty pET30(a) vector using *Thermo Scientific* ligase enzyme and ligation buffer (*InstaClone* PCR cloning kit). The 30 mm³ ligation reaction mixture contained 10 mm³ of purified digestion product (purified insert) and 5 mm³ of digested empty pET30(a) vector, 1 x final concentration of ligation buffer, 8 mm³ of nucleases free water, and 1 mm³ of T4 DNA ligase. It was incubated at 8 - 10 ºC overnight. Then, 10 mm³ of the ligation product of pET30(a) and C-1 (named C-1/pET construct) was transformed into competent *E. coli* Top10F’ following the methodology of Jahanpir et al. (2018). In these experiments, kanamycin (50 µg cm⁻³) was used for the selection of pET30(a) along with 12.5 µg cm⁻³ of tetracyclin for selection of Top10F’. The purified plasmid from true transformants contained 8 mm³ of purified construct plasmid, 10 mm³ of 0.5 M EDTA, and 35 mm³ of nucleases free water (3 h incubation at 37 ºC was provided for complete digestion). The tested plasmid was transformed into competent cells of *E. coli* strain BL21-DE3 following the *E.coli* transformation method (Jahanpir et al. 2018). The obtained transformants were again tested through PCR amplification with gene-specific primers and double restriction digestion with EcoRI and HindIII enzymes.

**Expression of Eucalyptus DHN-10 protein in BL21-DE3:** Single colony raised from confirmed C-1/pET/BL21 clone on selection media was incubated overnight at 30 ºC in 3 cm³ LB broth (*Lennox # LAB-173*) having kanamycin (50 µg cm⁻³). The enriched culture was diluted in the same medium (10 cm³) and multiplied at similar conditions until an absorbance A₆₀₀ was 0.7 which was attained in 4 h. The 1 cm³ of enriched culture was saved at -20 ºC to use as an un-induced sample. Isopropyl β-D-1-thiogalactopyranoside (IPTG) was added to the rest of the culture to attain the concentration of 1 mM and incubated again at the same conditions for 3 h. After incubation, the culture was cooled on ice for 20 min and dispensed into 1 cm³ aliquots. The cultured cells were harvested by centrifugation at 5 000 g and 4 ºC for 20 min. The pellets of induced and non-induced C-1/pET/BL21 cultures were resuspended in 100 mm³ of 100 mM phosphate buffer saline (PBS). The 40 mm³ from each suspension was mixed separately with 7 mm³ of 6x protein loading buffer (10 cm³ of 6x protein loading buffer was prepared from 1.2 g sodium dodecyl sulphate (SDS), 1.2 cm³ of 0.5 M Tris-HCl (pH 6.0), 0.01 % (m/v) Bromophenol Blue, 4.7 cm³ of glycerol, 2.1 cm³ of water) and boiled in a water bath for 10 min to denature the proteins. The tubes were then immediately shifted to the ice for 10 min. Both heating and cooling were repeated once again but each for 5 min. Then the tubes were centrifuged at 14 000 g and 4 ºC for 10 min and 20 mm³ of protein samples were loaded on 30 % (m/v) polyacrylamide gel prepared according to standard SDS-PAGE protocol (Laemmli 1970) in *Hoefer Mini*. After resolving the gel at 80 V for 45 min following by 120 V for 2.5 h, it was stained with Coomassie stain for 45 min. After proper staining, the gel was several times destained for clearing of a blue background. Then the gel was analysed for the presence of approximately 10 kDa protein of *Eucalyptus* DHN-10 in a protein mixture of induced and non-induced BL21 cells.

**Protein Dot Blot analysis:** The pellets of 1 cm³ from induced and non-induced C-1/pET/BL21 cultures were directly resuspended in 100 mm³ of protein sample buffer for dot blot analysis (50 cm³ of protein sample buffer was prepared from 24 g of urea, 2.5 g of SDS, 2 cm³ of 1 M Tris-HCl (pH 6.8), 10 mm³ of 0.5 M EDTA, and 0.5 cm³ 2-mercaptoethanol) in 1.5 cm³ centrifuge tubes and incubated in boiling water bath for 10 min. Then the tubes were placed at -20 ºC for 5 min. The tubes were again shifted to boiling water bath for another 10 min. Finally, the tubes were centrifuged at 14 000 g and 4 ºC for 10 min, and 7 mm³ of the supernatant containing crude protein extract was blotted on the *Hybond-C* membrane. The protein samples from induced and non-induced cells were blotted on the membrane separately and in triplicates. The membrane was air-dried in a laminar airflow cabinet for 20 min. Then membrane was blocked in 5 % blocking buffer (100 mM Tris of pH 7.5, 150 mM NaCl, 0.5 % blocking reagent) for 1 h. The membrane was soaked in Anti-His antibody (Roche) diluted to 1 mm³ cm⁻³ in the above-blocking buffer for 35 min at shaking (primary antibody: His-probe H-15, rabbit polyclonal Ig/G 200 µg cm⁻³). Then primary antibody was removed for secondary antibody diluted in the same solution (Anti-Rabbit Ig/G 200 µg cm⁻³). This protein was visualized on *NBT-BCIP* (Roche; 1 tablet per 10 cm² autoclaved double distilled water) and put for colour development in dark for 5 - 10 min.

**Results**

The TA cloned full-length DHN-10 gene of *Eucalyptus globulus*, as well as its coding region, were sequenced from *CAMB* commercial sequencing facility. The 315 bp long coding sequence was submitted in *NCBI* data bank as GZJ-2018 dehydrin-10 under accession number GZJ-2018 dehydrin-10 (GenBank: MG948256.1) after confirmation by comparing with reported mRNA sequence of DHN-10 gene of *Eucalyptus globulus* (GenBank: JN052210.1). It produces significant alignment with 100 % query cover and 93 % homology. The submitted sequence was translated on the *Expasy* translate tool and was analyzed on *ProtParam* (Fig. 1 Suppl.). The coding sequence of *Eucalyptus* sp. GZJ-2018 dehydrin-10 gene was 315 bp long whereas the already published coding sequence of the DHN-10 gene (JN052210.1) possesses 297 bp. 114 amino acids are present in GZJ-2018 dehydrin-10 protein with a total molecular mass of 12889.92 Da and theoretical pl of 6.13. This protein contains 29 negatively charged residues (Asp and Glu) and 22 positively charged residues (Arg and Lys).
The submitted GZJ-2018 dehydrin-10 sequence possesses extra integration of two positively charged histidine, one negatively charged glutamate residue, two negatively charged aspartate residues, and one extra lysine residue.
Furthermore, no Trp, Tyr, or Cys amino acids were present in GenBank MG948256.1 (Fig. 1). Hence, the quantification of this protein was not possible through UV spectrophotometry. The *Eucalyptus* sp. GZJ-2018 dehydrin-10 protein showed significant alignment with dehydrin-10 protein of two *Eucalyptus* species, the *E. nitens*, and *E. globulus*; but earlier possesses H, K, E, and S amino acids that are extra inserted than both later ones, the *E. nitens*, and *E. globulus* (Fig. 2 Suppl.).

The coding regions of *Eucalyptus* sp. GZJ-2018 dehydrin-10 gene was amplified from full-length TA cloned *DHN-10* gene with C-1 primer set and *DHN-10* gene with added restriction sites of EcoRI and HindIII on both ends was obtained. The same template was also amplified using a gene-specific primer set for full length (525 bp) amplification as PCR positive control. The gene specific primers that were designed using published sequence of *DHN-10* gene (JN052210.1) amplified the coding sequence of *DHN-10* gene from *Eucalyptus globulus* of Lahore Division with significant differences. The latter (MG948256.1) had 18 base pairs more than the former (JN052210.1). The purified PCR product of the coding region having restriction sites was cloned into pTZ57R/T vector. The resulting C-1/pTZ construct was transformed into *E. coli* Top10F’ strain. Product of double digestion (with EcoRI and HindIII enzymes) of tested and confirmed C-1/pET30(a) plasmid was ligated in empty pET30(a) vector (5 422 bp), digested with the same enzyme pair, and transformed in Top10F’ strain. The recombinant purified plasmid of survived transformants on kanamycin selection medium produced PCR bands of > 315 bp (315 bp coding region plus restriction sites of EcoRI and HindIII restriction enzymes with C-1 primers). A ~315 bp insert released on double digestion of the plasmid with EcoRI and HindIII enzyme pair. The digested pET30(a) vector remained slightly below the upper third band (of 6 000 bp) of the 1 kb DNA ladder. The confirmed plasmid construct (C-1/pET30(a)) was transformed in competent cells of the BL21-DE3 strain of *E. coli* (Fig. 3A). The transformation event was also verified through purification of the C-1/pET30(a) construct (Fig. 3B), PCR (Fig. 4A), and double restriction digestion (Fig. 4B). The protein extract from
induced and non-induced cell pellet of C-1/pET/BL21 clone was observed on SDS-PAGE. A prominent 10 kDa band was visible in induced cell protein whereas no band was observed in a non-induced pellet of the same culture (Fig. 5A). The crude protein extract from the same C-1/pET/BL21 clone was also blotted on nitrocellulose membrane Hybond C for protein dot blot analysis. Secondary antibody-NBT/BCIP substrate color reaction confirmed the production of DHN-10 protein in induced BL21 cells from transformed C-1/pET30(a) plasmid (Fig. 5B). No protein production was observed in the same culture without the induction of a strong T7 promoter. These observations indicated that the DHN-10 gene of Eucalyptus can be successfully expressed outside the parent cell, in an entirely different expression system, if an appropriate promoter is provided.

Discussion

The presence of one extra lysine residue in studied DHN-10 (MG948256.1) can be expected for enhanced amphipathic quality which is related to the binding of dehydrin proteins with macromolecules. As dehydrin proteins are molecules with characteristic lysine-rich K-segments, the presence of extra K amino acid residue will aid to all of the specific functions that are linked to or influenced with K-segment. The K-segment is involved in the formation of an amphipathic helix and plays an important role in dehydrin-macromolecules interaction (Close 1996, Close et al. 1993). On the perception of dehydration signals from the microenvironment of cell, K-segments adopt characteristic α-helical conformation (Hanin et al. 2011). It is the response mechanism of the plant cells to abiotic stresses. Koag et al. (2003) have stated that conformational changes in the K segment of dehydrin proteins enhance their role in stabilizing the membranes of a cell during dehydration conditions. The presence of five extra charged amino acid residues like two positively charged histidine, one negatively charged glutamate, and two negatively charged aspartate residues (Fig. 2) in studied DHN-10 protein has enhanced its importance because the dehydrins with higher content of reactive amino acids on the surface possess free metal ion binding and so ROS scavenging properties (Hanin et al. 2011). Free metal ions are also involved, as catalysts, in the synthesis of reactive oxygen species. Therefore, the removal of free metal ions from the medium reduces the synthesis of various ROS (Hara et al. 2001, 2005, Heyen et al. 2002, Hanin et al. 2011). Dehydrins are highly hydrophilic proteins that help other biomolecules to maintain high configurational flexibility and defy the unspecified aggregation and collapse (Mouillon et al. 2006, Koag et al. 2009).

The Eucalyptus DHN-10 gene was not reported earlier to be cloned or expressed in any expression system outside the Eucalyptus species. It is the first report showing the possibility to be sub-cloned in a bacterial expression vector, pET30(a). The induced expression of DHN-10 is also first time reported in E.coli strain BL21-DE3. Cloned DHN-10 gene was found to be successfully translated into a stable protein that was detected in protein dot blot analysis. Although no report could be found upon genetic cloning and expression of Eucalyptus dehydrin genes in the expression vector, however, a significant mass of work has been conducted on sequence study of dehydrin genes. Some other researchers have cloned diverse dehydrin genes from a variety of plant species. For instance, Zeng et al. (2018) have cloned the RNA sequence of the LEA gene family from Gastrodia elata into E. coli to evaluate their function under low-temperature stress. A novel SK3-type dehydrin of the Stipa purpurea plant has been studied and cloned by Yang et al. (2014). Yao et al. (2005) have also reported novel dehydrins in a specific subclass of the LEA protein family from Brassica juncea and Brassica napus. Full-length cDNA of dehydrin-like protein of Pistacia vera has been reported to be cloned by Yakubov et al. (2005).

Conclusions

We successfully cloned and expressed the novel dehydrin-10 gene from the Eucalyptus globulus of Pakistani origin. The in silico analyses of the DHN-10 gene revealed presence of five additional electrically charged amino acids (two positively charged histidine, one negatively charged glutamate, and two negatively charged aspartate residues) and one lysine. The DHN-10 was successfully cloned in a bacterial expression vector (pET30(a)) and its expression was detected in E. coli. The expressed DHN-10 possesses enhanced characteristics of abiotic stress tolerance because of the presence of one additional lysine residue which is directly involved in the amphipathic characteristic of dehydrins. Furthermore, DHN-10 can also be expected to be involved in antioxidants activity because of the presence of five additional charged residues that indicates the possible involvement of DHN-10 (MG948256.1) as an anti-oxidant molecule in the cell which is also a characteristic of a stress response. The present study will be helpful in future practices of cloning and expression of novel dehydrins and dehydrin-like genes.

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