



Genome-wide identification and comparative expression analysis of *LEA* genes in watermelon and melon genomes

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Abstract Late embryogenesis abundant (LEA) proteins are large and diverse group of polypeptides which were first identified during seed dehydration and then in vegetative plant tissues during different stress responses. Now, gene family members of LEA proteins have been detected in various organisms. However, there is no report for this protein family in watermelon and melon until this study. A total of 73 *LEA* genes from watermelon (*CILEA*) and 61 *LEA* genes from melon (*CmLEA*) were identified in this comprehensive study. They were classified into four and three distinct clusters in watermelon and melon, respectively. There was a correlation between gene structure and motif composition among each LEA groups. Segmental duplication played an important role for *LEA* gene expansion in watermelon. Maximum gene ontology of *LEA* genes was observed with poplar *LEA* genes. For evaluation of tissue specific expression patterns of *CILEA* and *CmLEA* genes, publicly available RNA-seq data were analyzed. The expression analysis of selected *LEA* genes in root and leaf tissues of drought-stressed watermelon and melon were examined using qRT-PCR. Among them, *CILEA-12-*

17-46 genes were quickly induced after drought application. Therefore, they might be considered as early response genes for water limitation conditions in watermelon. In addition, *CmLEA-42-43* genes were found to be up-regulated in both tissues of melon under drought stress. Our results can open up new frontiers about understanding of functions of these important family members under normal developmental stages and stress conditions by bioinformatics and transcriptomic approaches.

Keywords *LEA* gene family · *Citrullus lanatus* · *Cucumis melo* · Genome-wide analysis · Drought stress

Introduction

Late embryogenesis abundant (LEA) proteins were firstly identified in developing cotton seeds (Dure and Galau 1981) and then, abundance of this protein family was discovered in different organisms such as *Brassica napus*, *Arabidopsis thaliana*, *Caenorhabditis elegans*, *Dictyostelium discoideum*, etc. (Gal et al. 2004; Eichinger et al. 2005; Kikawada et al. 2006; Dalal et al. 2009; Denekamp et al. 2010). Especially most of LEA proteins are called as hydrophilins because of their high Arg/Lys, Glu, Ala, Thr and Gly amino acids contents (Battaglia et al. 2008). Analysis of LEA proteins revealed that this protein group can be divided into seven distinct classes named as LEA 1–5, dehydrin and SMP (Seed Maturation Protein) according to their different motif contents. However, their classification is still controversial and there are differences between authors (Dure et al. 1989; Hunault and Jaspard 2010). In this study, Pfam nomenclature was used for LEA protein classification.

Plant LEA proteins have many important functions in normal plant growth and in preserving cells from the

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detrimental effect of different stress conditions. For example, these proteins have been considered to prevent the inactivation of malate dehydrogenase, lactate dehydrogenase, catalase or citrate synthase enzymes under desiccation or cold conditions according to *in vitro* studies (Hara et al. 2001; Goyal et al. 2005; Reyes et al. 2008). Besides, membrane preservation with sugars and sequestration of ions were other roles of LEA proteins (Wolkers et al. 2001; Grelet et al. 2005; Liu et al. 2010). In addition, premature seed dehydration and maturation at the distal end of siliques were reported in *LEA* gene (*ATEM6*) knocked out Arabidopsis plants, which demonstrates the role of *LEA* genes in normal seed development and plant growth (Manfre et al. 2006).

This group of proteins were considered to be related with abiotic stress tolerance. As evidence, over expression of PM2 protein from soybean in *Escherichia coli* provided living of bacteria in high salt or extreme temperature conditions (Liu and Zheng 2005; Liu et al. 2010). Besides, Arabidopsis plants lack of one or two of three LEA 4 proteins were sensitive to water limitation conditions (Olvera-Carrillo et al. 2010). In addition, a LEA 3 like protein of *Artemia franciscana* revealed the increased survival capacity of *Drosophila melanogaster* cells under water deficiency conditions (Marunde et al. 2013). *E. coli* cells expressing OsLEA protein presented an increase resistance to cold, salinity, osmotic, heat and UV radiation conditions (He et al. 2012). Furthermore, some LEA proteins have a membrane stabilization function by association of anionic phospholipid vesicles under drought or cold conditions (Kosová et al. 2007; Tolleter et al. 2010).

There are a few studies subjected on LEA protein family by genome wide search. Up to date, studies, which make identification and characterization of LEA proteins in Arabidopsis, rice, soybean, tomato, potato, maize, poplar, sweet orange, cucumber, etc. were performed. In these studies, chromosomal location and gene duplications, cis-elements, phylogenetic relationships and conserved domains of LEA proteins were determined. Also, expression patterns of this gene family under abiotic stress conditions were evaluated using publically open RNA seq data and quantitative PCR results (Wang et al. 2007; Hundertmark and Hinch 2008; Li et al. 2011; Lan et al. 2013; Cao and Li 2015; Charfeddine et al. 2015; Pedrosa et al. 2015; Altunoglu et al. 2016; Li and Cao 2016).

Melon (*Cucumis melo*) and watermelon (*Citrullus lanatus*) are important crops and belong to Cucurbitaceae family, which includes cucumber (*Cucumis sativus* L.), squash (*Cucurbita* spp.), pumpkin (*Cucurbita* L. spp.) and gourd (*Lagenaria* Ser. spp., *Luffa* Mill. spp., and *Cucurbita* spp.). Production of watermelon is about 104 million Mg

and it is in the top five of the most eaten fresh fruits around the world according to Food and Agriculture Organization (FAO) (<http://faostat.fao.org>). In addition, Turkey is the third largest producer of watermelon after China and Iran (Kumar and Wehner 2011). Besides, melon production is about 26 million tons worldwide according to FAO reports (<http://faostat.fao.org>), which has an economic value in some European and East Asian countries (Garcia-Mas et al. 2012).

Cucurbitaceae family is a useful model system for studying vascular biology and sex determination (Liu et al. 2004). In addition, whole genome sequences of melon and watermelon were reported in 2012 and 2013, respectively (Garcia-Mas et al. 2012; Guo et al. 2013). To our knowledge, LEA proteins of melon and watermelon have not been characterized yet. Therefore, we have performed genome wide analysis for these important family members and have identified 61 and 73 *LEA* gene members in melon and watermelon genomes, respectively. Then, bioinformatics analysis and molecular characterization of these *LEA* genes were performed. Besides, expression patterns were evaluated using RNA seq data and quantitative real time PCR analysis (qRT-PCR).

Materials and methods

Identification of *LEA* genes in melon and watermelon genomes

Three different applications were used for identification of *LEA* genes in *Cucumis melo* and *Citrullus lanatus*. *LEA* protein sequences of 14 plants (*Arabidopsis thaliana*, *Gossypium hirsutum*, *Oryza sativa*, *Glycine max*, *Sorghum bicolor*, *Triticum aestivum*, *Triticum durum*, *Pisum sativum*, *Zea mays*, *Hordeum vulgare*, *Brassica napus*, *Medicago truncatula*, *Nicotiana tabacum*, *Vitis vinifera*) were obtained by using LEAP database which is a data bank including *LEA* proteins from different organisms (<http://forge.info.univ-angers.fr/~gh/Leadb/index.php?action=0&mode=0>) (Hunault and Jaspard 2010).

These sequences were used in specification of cucumber and homologous peptides by making BLASTP search in open accessed PHYTOZOME v9.1 database which provides access to the sequenced and annotated green plant genomes (Goodstein et al. 2012). In addition to this, the database was scanned using *LEA* keywords. Further, Hidden Markov Model (HMM) profiles of *LEA* proteins in the Pfam database (<http://pfam.sanger.ac.uk/>) were compared with *Cucumis sativus* in the Phytozome database. Similar scanning of *Cucumis sativus*, *Cucumis melo*

and *Citrullus lanatus* genomes EST sequences were conducted with TBLASTN in NCBI database for possible LEA proteins. By using decrease redundancy tool (web.expasy.org/decrease_redundancy), related sequences were added to the study and non-related sequences were removed from the study. Preserved areas of every sequence added to the study were also controlled by using SMART database which provides the identification and annotation of genetically mobile domains and the analysis of domain architectures (<http://smart.emblheidelberg.de/>) (Letunic et al. 2012) and Pfam database which is a large collection of protein families (<http://pfam.sanger.ac.uk/>).

Identification of chromosomal location and estimation of genomic distribution of *LEA* genes

Chromosomal locations of *LEA* genes were identified using Cucurbit Genomics database (<http://www.icugi.org/cgi-bin/ICuGI/tool/blast.cgi>) which allows search in genomes of Cucurbitaceae family members for watermelon.

Tandem and segmental duplications were identified with Plant Genome Duplication Database (PGDD) (Tang et al. 2008) which is a public database and provides to identify and catalog plant genes in terms of intragenome or cross-genome relationships. Briefly, top five matches $\leq 1e-05$ were taken as potential anchors after BLASTP search against all potential peptide sequences of melon and watermelon *LEA* proteins. Collinear blocks were searched by MCSScan and $\leq 1e-10$ alignments were considered as meaningful matches (Tang et al. 2008; Du et al. 2013).

Sequence alignment, phylogenetic analysis and identification of preserved motifs

Amino acid sequences were loaded to MEGA6 (Tamura et al. 2013) program which enables to conduct statistical analysis of molecular evolution. Then, multiple sequence alignments were done by using ClustalW. Aligned file was used for constructing of phylogenetic tree by neighbor joining method with bootstrap analysis for 1000 iterations (Saitou and Nei 1987). Jones–Taylor–Thornton (JTT) substitution model was applied to phylogeny reconstruction and rates among sites were gamma distributed (G). Protein sequence motifs were identified by MEME motif search tool (<http://meme.nbcr.net/meme3/meme.html>) (Bailey and Elkan 1994) which allows to discover novel motifs in collections of unaligned nucleotide or protein sequences. Specified MEME motifs were scanned in InterPro database with InterProScan which enables functional analysis of proteins by classifying them into families and predicting domains and important sites (Quevillon et al. 2005).

Gene ontology analysis (GO)

Functional analysis of *LEA* proteins was performed by using Blast2GO (<http://www.blast2go.com>) program (Conesa and Götz 2008). Amino acid sequences of *LEA* proteins were loaded to Blast2GO program and three steps accomplished for functional analysis with this program (i) matching was conducted with the loaded sequences in the program (BLASTp) (ii) mapping associated with BLAST results was completed (Mapping) (iii) dump file related to sequences was created (Annotation). Briefly, with the program, determination of biological functions, cellular content and molecular functions which are the three categories of the GO categorization were provided.

Comparative physical mapping of watermelon *LEA* genes with other organisms

For the purpose of revealing orthologous relations between watermelon and other species including Arabidopsis, soybean, poplar, potato, grape and maize, peptide sequences were scanned in response to amino acid sequences of watermelon *LEA* proteins using BlastP program. Those having e -value $\leq 1e-5$ and are at least 80% identical were considered as meaningful. Orthologous relations of *LEA* genes between watermelon and Arabidopsis, soybean, poplar, potato, grape and maize were viewed with MapChart program (Voorrips 2002).

Calculation of homologous and nonhomologous change rates

Beside the amino acid sequences of *LEA* genes coding duplicated protein, orthologous gene pairs between melon and watermelon and Arabidopsis, rice and soybean were also aligned by utilizing the ClustalW based multiple sequence alignment tools. CODEML (<http://www.bork.embl.de/pal2nal/>) program (Suyama et al. 2006) was used for calculating homologous (Ks) and nonhomologous (Ka) change rates through amino acid sequences of *LEA* proteins and their original complementary DNA sequences. Duplication and separation time of every *LEA* gene was calculated by using homologous mutation change of (million years ago, MYA) λ changes in response to every homologous area and year ($T = Ks/2\lambda$ ($\lambda = 6.5 \times 10^{-9}$)) (Lynch and Conery 2000; Yang et al. 2008).

In silico identification of miRNAs targeting *LEA* genes

Identification of miRNA controlled gene targets is important for understanding the miRNA functions. Plant miRNA database (<http://bioinformatics.cau.edu.cn/PMRD/>) was

used for identification of miRNAs targeting melon and watermelon *LEA* genes. In order to obtain previously known plant miRNAs, miRBase v20.0 (<http://www.mirbase.org/>) program was utilized. All known plant miRNAs and melon, watermelon *LEA* gene transcripts were aligned together. Then, all melon and watermelon miRNAs were defined using internet based psRNA Target Server (<http://plantgrn.noble.org/psRNATarget/>). Using parameters described earlier by Zhang (2005), all known plant miRNAs and their potential target or targets were evaluated. These miRNA targets determined with computer method were analyzed by utilizing BLASTX scans in order to identify default gene homologous for the confirmation.

Estimation of predicated structures of CILEA and CmLEA proteins

Melon and watermelon *LEA* genes were scanned in Protein Data Bank (PDB) which gives information about the 3D structures of proteins. In order to determine the best 3D structure of similar sequence, known three dimensional structures were used for BLASTP analysis (Berman et al. 2000). For the obtained information from Phyre2 program (Protein Homology/Analog/Recognition Engine; <http://www.sbg.bio.ic.ac.uk/phyre2>), protein structure was estimated with homology modeling (Kelley et al. 2015).

Expression profiling of melon and watermelon *LEA* genes by using transcriptome data

For RNA-Seq analysis, all the Illumina HiSeq readings were obtained from an open data bank archive (SRA, Sequence Read Archive). Entry numbers used in readings are as follows for melon; SRR411102, SRR411100, SRR411106, SRR411104, SRR1033647, SRR1033646, SRR2082958, SRR2082965, SRR2082865, SRR2082935, SRR2082943, SRR2082953, SRR2082831, SRR2082832, SRR2082790, SRR2082791, SRR2082796, SRR2082813 and for watermelon; SRR1724899, SRR1724900, SRR1724901, SRR1724902, SRR1724903, SRR1724943, WM-UR-1/SRR1001435, WM-UR-2/SRR1001436, WM-IM-1/SRR1001437, WM-IM-2/SRR1001438, WM-PM-1/SRR1001439, WM-PM-2/SRR1001440, WM-MA-1/SRR1001441, WM-MA-2/SRR1001442, SRR494474, SRR518988, SRR518988, SRR494479, SRR518992, SRR518993.

All readings were downloaded as raw sequence data in “sra” format and converted into “fastq” format. After removing of the readings at low quality [Phred quality (*Q*) score <20], all the clear readings were subjected to FastQC analysis in order to control their reading qualities in terms of sequence quality for every base, quality score for every base, nucleotide content for every base and sequence duplication level.

Plant samples, growing conditions and stress application

Melon and watermelon seeds were supplied from Monsanto Gıda ve Tarım Tic. Ltd. Şti (Antalya). Seed shells were removed and seeds were washed with distilled water three times. Afterwards, seeds were taken into plastic pots and grown inside the plant grow box in culture medium containing Hoagland solution (Hoagland and Arnon 1950) for 14 days at $400 \mu\text{mol m}^{-2} \text{s}^{-1}$ light intensity at $24 \pm 2 \text{ }^\circ\text{C}$ and 16 h of light and 8 h of dark photoperiod. For drought stress, 10% polyethylene glycol 6000 (PEG-6000) was added to Hoagland solution. Considering morphological features of grown plants and taking parameters used by Baloglu et al. (2014) in the cucumber plant into account, at the zero, third, twelfth and twenty fourth hours of the stress application, samples were taken from stressed plants and control plants for RNA isolation. Stress applied plants and control plants were grown in the plant grow box at the same conditions. The zero hour was used as a control. Root and leaf samples of the grown plants were collected separately in order to use in the tissue specific expression analysis. Tissue samples taken by making biological sampling at three at a time were used in tissue specific gene expression analysis by being frozen with liquid nitrogen.

RNA isolation and quantitative real time PCR (qRT-PCR) analysis

Total plant RNA was isolated using Invitrogen Trizol reagent (Life Technologies Corporation, NY, USA). The quality and quantity of isolated RNA was checked using agarose gel electrophoresis and MultiscanGO nano-spectrophotometer (ThermoFisher Scientific, USA). Single and double stranded DNA fragments present in RNA samples were removed by DNase I enzyme (Thermo, Lithuania). cDNA synthesis was done using RevertAid First Strand cDNA Synthesis Kit according to manufacturers' instructions (ThermoFisher Scientific, USA). For the qRT-PCR analysis, *LEA* genes in which it was seen an increase in gene expression were detected and gene specific primers were designed for those *LEA* genes which were showing increased expression using Primer 5 software program and a list of used primers is given in Supplementary Table 11. *18S rRNA* gene was used as internal control gene in the study. Primer sequences of the used *18S rRNA* gene is as follows (GenBank ID: X51542.1): 5-GTGACGGGTGACGGAGAATT-3 and 5-GACACTAATGCGCCCGGTAT-3 (Baloglu et al. 2014). Sampling was performed for three times for each step and each repetition, triple qRT-PCR test was done using SYBR Green master mix in Light Cycler 480 Real-Time PCR System (Roche Applied

Science, Germany) according to manufacturers' recommendations.

Taking *18SrRNA* gene expression as reference, relative gene expressions were calculated. Proportioning gene expression levels to *18SrRNA* gene, control and stress time periods were compared with each other. Calculating values of cycle threshold number (CT) for every uprising curve, calibration curves were formed. Δ CT and $\Delta\Delta$ CT values were calculated as Δ CT = CT sample – CT reference and $\Delta\Delta$ CT = Δ CT stress given sample – Δ CT control (in 0 h) and the difference at the expression level was determined as $2^{-\Delta\Delta Ct}$ (Livak and Schmittgen 2001).

Statistical analysis

Statistical calculations and comparisons were performed by using Minitab 17 package program. Student *T* test was used in the analysis of the differences between the stress given samples and control samples. If *p* value is <0.01; it is assumed that the expression difference in the relevant *LEA* gene is meaningful.

Results and discussion

Determination and characterization of *LEA* gene family members in melon and watermelon

LEA protein sequences of 14 plant from LEAP database were used for determination of melon and watermelon *LEA* genes. After detailed searches in Phytozome v9.1, Pfam and NCBI etc. databases, Melonomics database for melon and Cucurbit Genomics Database for watermelon were employed to find *LEA* sequences belong to these plants. Consequently, 73 *LEA* genes in watermelon (named as *CILEA-1* to *CILEA-73*) and 61 *LEA* genes in melon (*CmLEA-1* to *CmLEA-61*), mostly which were carrying conserved *LEA* domain, were determined. Protein lengths of identified *LEA* proteins changed between 64 and 464 amino acids in melon and between 64 and 1162 amino acids in watermelon. Also, molecular weights of these proteins varied between 6779.4 and 129,808.3 Da in watermelon. The smallest *LEA* protein was of 51.242 Da, whereas the biggest *LEA* protein was of 6795.4 Da in melon. Moreover, according to physicochemical analysis, 68 and 75% of *LEA* proteins were of a basic character in watermelon and melon, respectively. Most of the *LEA2* and *LEA1* group proteins were basic whereas most of the *LEA4*, *LEA5*, *SMP* and dehydrin group proteins were of an acidic character in watermelon and melon. *LEA3* protein group composed of acidic and basic proteins in both plants. Our results were consistent with the results of Filiz et al.

(2013) who found that *LEA4*, *LEA5* and *LEA6* group proteins were acidic and most of *LEA* proteins were basic (72.2%) in purple false brome. More information about results of used parameters for *CmLEA* and *CILEA* proteins is available in Supplementary Table 1 and Supplementary Table 2.

Similar studies on genome wide identification of *LEA* family members in other plants revealed, 79 *LEA* genes in cucumber (Altunoglu et al. 2016), 53 *LEA* genes in poplar (Lan et al. 2013), 51 *LEA* genes in Arabidopsis (Hundertmark and Hinch 2008), 36 *LEA* genes in soybean (Li et al. 2011), 34 *LEA* genes in rice (Wang et al. 2007), 32 *LEA* genes in maize (Li and Cao 2016), 29 *LEA* genes in potato (Charfeddine et al. 2015), 27 *LEA* genes in tomato (Cao and Li 2015) and 72 genes in sweet orange (Pedrosa et al. 2015). Accordingly, watermelon has the second highest number of *LEA* protein genes after cucumber enabling a lot more different types of searches in different databases and to include more plants for inquiry.

Chromosomal distribution and duplication of *CmLEA* and *CILEA* genes

CILEA genes were mapped on 11 watermelon chromosomes. Chromosome 1 and 10 had the most *LEA* genes with the number of 11 whereas chromosome 1 and 8 had a few *LEA* genes with the number of 3. *CILEA* genes locating on chromosome 1, 5 and 9 were sited at the upper end of chromosomes but *CILEA* genes locating on chromosome 2, 7, 10 and 11 were sited at the lower end of these chromosomes. Only *CILEA-73* gene couldn't be mapped on any chromosome (Fig. 1). Chromosomal distributions of *CmLEA* genes could not be determined because locations of these genes were still in scaffold level.

Tandem and segmental duplications events for *CILEA* and *CmLEA* genes were evaluated because duplication of genes is one of the reason for occurrence of orthologous genes (Mehan et al. 2004). Twenty of *CmLEA* genes (32%) showed duplication events, however, we couldn't elucidate the duplication type as a result of absence of exact chromosomal locations of *CmLEA* genes (Supplementary Table 4). Segmental duplications between *LEA* genes were determined in watermelon with the percentage of 31%. Seven *CILEA* genes showed (9.5%) tandem duplication events. These tandem duplications located on chromosome 1 (*CILEA-6-9*) and chromosome 10 (*CILEA-64-65-66-67-68*) (Supplementary Table 3). According to tandem duplication analysis, frequencies were 70 and 30%, in cucumber and poplar, respectively. Our findings suggest that tandem duplication is not prevalent in watermelon as potato and rice. Besides, segmental duplication rates were 51% in

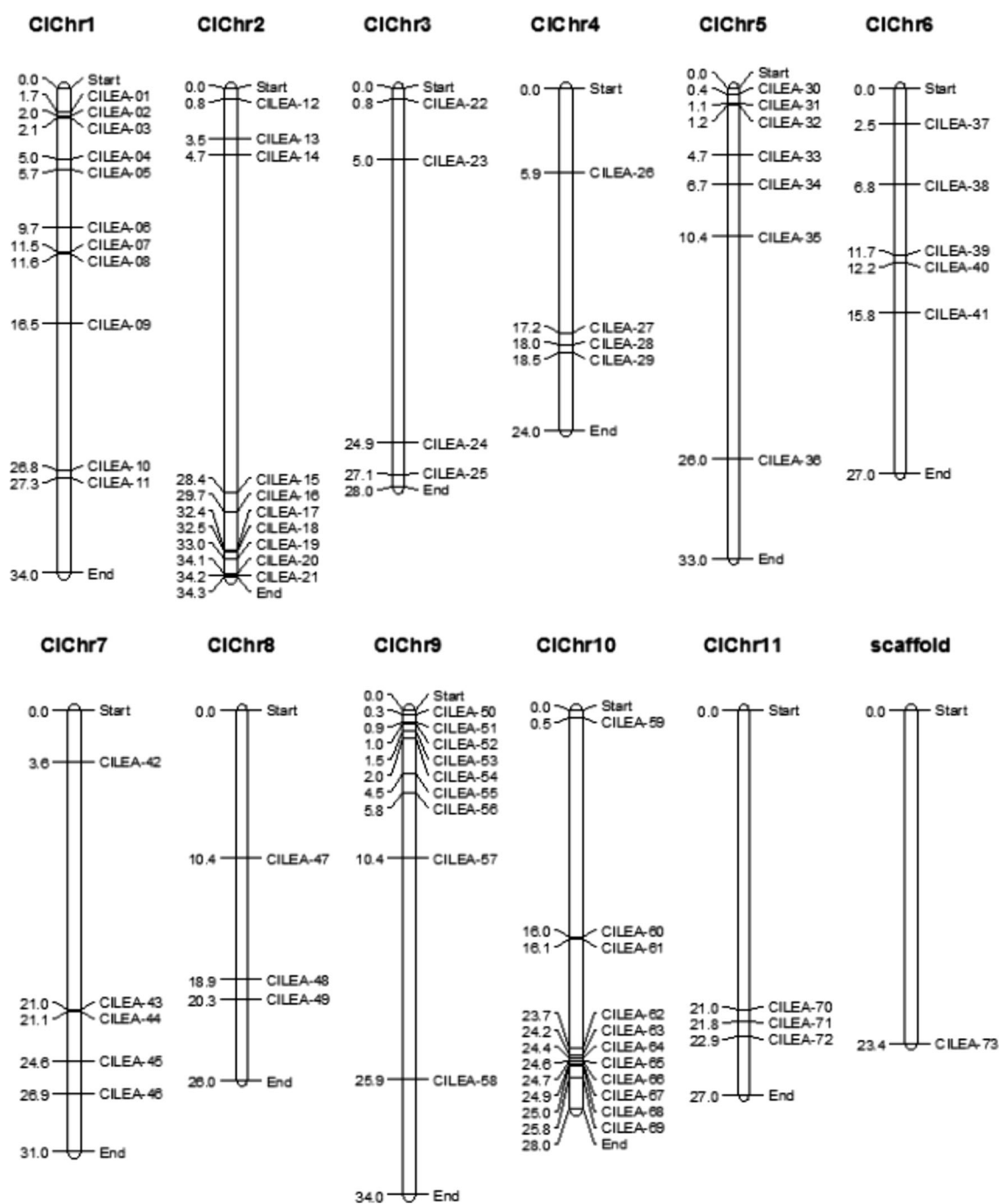


Fig. 1 Chromosomal distribution of 73 *CILEA* genes. Physical locations of watermelon *LEA* genes were presented on 11 watermelon chromosomes numbered from C1Chr1 to C1Chr11

tomato, 43% in Arabidopsis, 25% in maize, 20% in potato, 17% in Chinese plum and 15% in rice (Wang et al. 2007; Hundertmark and Hincha 2008; Lan et al. 2013; Cao and Li 2015; Charfeddine et al. 2015; Altunoglu et al. 2016; Li and Cao 2016; Du et al. 2013). Our results indicate that segmental duplication events may be one of the main reason for *LEA* gene family expansion in watermelon.

Phylogenetic distribution and identification of conserved motifs

Phylogenetic analysis was implemented by Neighbour-Joining method with bootstrap analysis for 1000 repetitions for 61 melon and 73 watermelon *LEA* proteins to analyze evolutionary relationships between this groups of proteins. Jones-Taylor-Thornton (JTT) substitution model was used

in MEGA v.6 software (Molecular Evolutionary Genetics Analysis) (Tamura et al. 2013) and rates among sites were gamma distributed (G). Four distinct clusters (Cluster I–IV) including 9, 23, 20 and 21 proteins were determined for watermelon LEA proteins, respectively (Fig. 2b). LEA 2 group proteins were dominant in watermelon according to Pfam nomenclature. LEA1 group proteins were located in Cluster I (CILEA-26), Cluster II (CILEA-62) and Cluster III (CILEA-59). Three members of LEA3 group proteins (CILEA-55-60-61) were in Cluster III whereas one member of this group (CILEA-52) was located in Cluster III. In addition, only one member of LEA4 group protein (CILEA-69) which located in Cluster IV was determined in watermelon. LEA5 (CILEA-28-29), LEA6 (CILEA-11-70) and SMP (CILEA-25-33-38-56) group of proteins accumulated only in the Cluster III. Besides, dehydrin group of proteins were distributed in Cluster II (CILEA-72), Cluster III (CILEA-14-43-44) and Cluster IV (CILEA-48) and LEA2 group of proteins were in all clusters mainly in Cluster IV.

Three main clusters (Cluster I–III) were observed among melon LEA proteins, which comprised of 11, 5 and 45 proteins, respectively (Fig. 2a). LEA2 group was the dominant group between CmLEA proteins. SMP (CmLEA-23-32-33), LEA3 (CmLEA-34-38) and LEA4 (CmLEA-12) group proteins accumulated in Cluster I. LEA5 group proteins (CmLEA-58-59) were in Cluster II. Dehydrin group of proteins were located in Cluster II (CmLEA-36-46) and Cluster III (CmLEA-60). Cluster III contained only

LEA2 group of proteins. LEA6 group proteins weren't seen in melon.

Especially, SMP, LEA3 and LEA5 group proteins were distributed in a cluster whereas the dehydrin and LEA2 group proteins were allocated to different clusters in melon and watermelon. This kind of expansions may explain the different motif contents of LEA2 group proteins. According to phylogeny based analysis for LEA proteins in other plants, the highest number of LEA protein clusters were LEA4 group in Arabidopsis (Hundertmark and Hinch 2008), dehydrin group in tomato (Cao and Li 2015), LEA2 and LEA3 groups in maize (Li and Cao 2016), dehydrin and LEA2 groups in Chinese plum (Du et al. 2013), LEA2 group in cucumber (Altunoglu et al. 2016) and LEA4 group in poplar (Lan et al. 2013). LEA2 group was the mainly distributed group in watermelon, melon and cucumber which were in the Cucurbitaceae family. LEA2 group proteins may more required than other group of LEA proteins for the specific functions in this important plant family.

Moreover, 17 pairs of segmentally and 5 pairs of tandemly duplicated watermelon LEA genes were classified in the same cluster in phylogenetic tree. As an example, CILEA-16-27 and CILEA-24-20 proteins which were segmentally duplicated and located in the fourth cluster and all of them belonged to LEA2 group proteins. Tandemly duplicated CILEA-64-65 and CILEA-65-68 proteins were in the second cluster. These motif and sequence conservations can be attributed to the reliability of phylogenetic

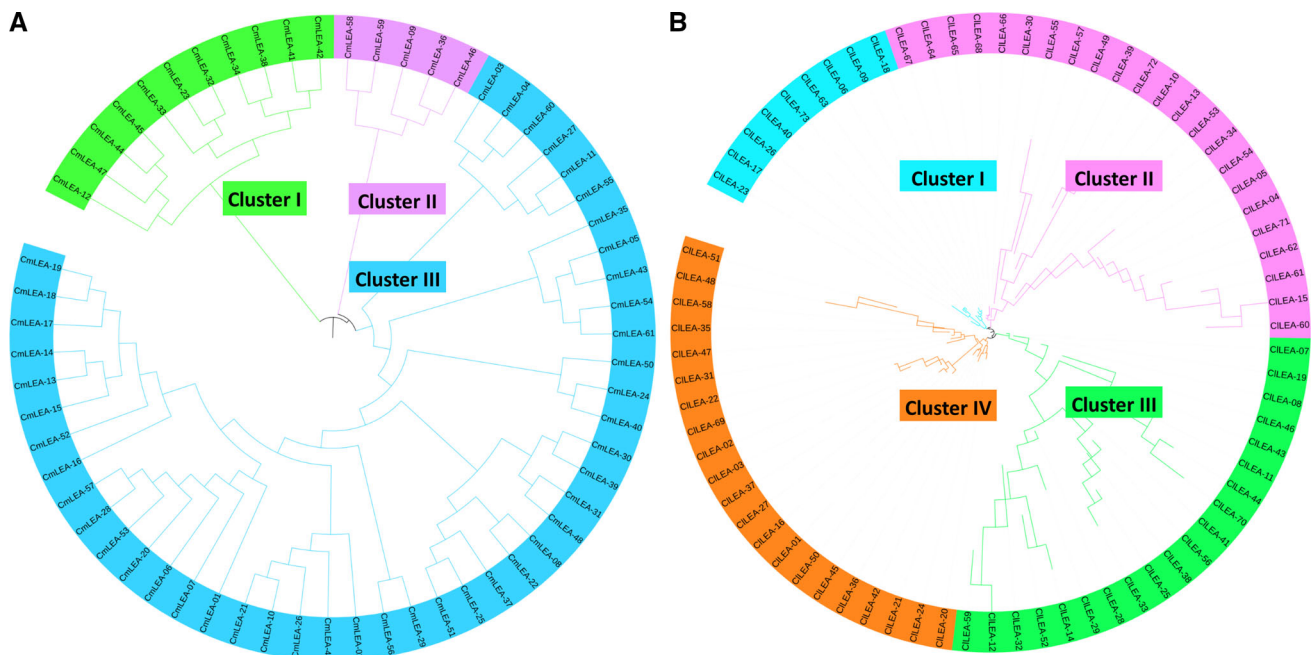


Fig. 2 Phylogenetic comparison of *CILEA* and *CmLEA* genes. **a** Phylogenetic tree of melon *LEA* genes with three clusters. **b** Phylogenetic tree of watermelon *LEA* genes with four clusters

tree and may be related to various of biological functions of this group of proteins.

Another phylogenetic tree which was containing all LEA proteins from cucumber, watermelon and melon was constructed (Altunoglu et al. 2016) (Supplementary Fig. 4). Six different clusters were observed among these LEA proteins from different origins. LEA2 group of proteins was the most dominant group in cluster 6. Other LEA groups (LEA1-6, SMP and dehydrin) dispersed to other clusters. Same LEA protein groups were determined in different clusters. This may be arisen from different motif contents of same LEA groups from different plants.

In addition, amino acid motif compositions of melon and watermelon LEA proteins were evaluated by MEME software. According to this analysis, 10 different motifs were determined for 67 CILEA proteins in watermelon (Table 1; Supplementary Fig. 3b). LEA1 (CILEA-26-59-62) and LEA4 (CILEA-69) group proteins had only motif 6. All LEA3 (CILEA-52-55-60-61) and LEA5 (CILEA-28-29) group proteins included only motif 10. CILEA-25-33-

38-56 proteins all of which were in the SMP protein group had motif 5 whereas all of dehydrin group proteins had no motifs with the exception of CILEA-48 which includes motif 6. LEA6 group proteins had no motifs like dehydrin group. LEA2 group proteins especially included motif 1, 2, 3 and 4 also motif 7 and 8 could be found in some members of LEA2 group.

Ten different motifs were found for 52 CmLEA proteins in melon using MEME software (Table 2; Supplementary Fig. 3a). Dehydrin and SMP group proteins had no motifs whereas LEA1, LEA4 group proteins and CmLEA-41-44-45-47 proteins had only motif 10. LEA2 group proteins included motif 1, 2, 3, 4 and 8 and some of LEA2 group proteins also had motif 5 and 7. Especially LEA1 and LEA4 protein groups had only one common motif, when LEA protein motif compositions of cucumber, watermelon and melon were compared (Altunoglu et al. 2016). This suggests that this group of LEA proteins may have similar function and role in these plants. In addition, dehydrin group proteins had no motifs in these plants in contrast to

Table 1 Amino acid sequence composition of 10 different motifs observed in CILEA proteins

Motif No.	Sites	E-value	Amino acid sequence composition of motif	Width (aa)
Motif 1	41	7.9e-115	NPNKRIGIYYD	11
Motif 2	32	3.9e-144	FLVGLAVLILWLVFRPHKQF	21
Motif 3	32	1.1e-096	LPPFYQGHKNTTVWSPFLSG	20
Motif 4	24	1.5e-090	GRVRKVGWKTWRVRIIVNC	21
Motif 5	4	1.5e-040	KVEQSDAAAIQAAEVRATGQNVIIIPGGIAATAQSAATFNTRMMRDEDKI	50
Motif 6	6	5.1e-042	KMSYHAGEAKGQAEKASNMMDKASDAAQEAKEQMGGQAEKAKAQGAA	50
Motif 7	4	5.5e-029	GSDSTGVPTKLLTLNCTLRITYHNPATFFGIIVSSSPIQLMYSQIQIASG	50
Motif 8	16	6.2e-027	RCHWRCCCCCCCW	15
Motif 9	7	3.3e-031	GTMNISSYARIPGRVRLHIFKHHVVSTMSCSMTIDISNHSIQDQWC	47
Motif 10	6	1.1e-029	MGHEEYQEMTEKGGWMPDPMGTGYRPEHNGDEIDEAELRTK	41

Table 2 Amino acid sequence composition of 10 different motifs observed in CmLEA proteins

Motif No.	Sites	E-value	Amino acid sequence composition of motif	Width (aa)
Motif 1	27	4.5e-154	RNPKNRIGIYYDTIEVYAMYK	21
Motif 2	41	3.3e-158	WLVFRPHKQFHVQDVQLTQF	21
Motif 3	32	1.1e-124	QRLCTGWLPPFYQGHKNTTVWSPHFEQQ	29
Motif 4	26	1.2e-118	ELNVEFDGRVRKMGWVKMGRYKPKVNCE	29
Motif 5	13	7.0e-031	CWRCCCCFCL	11
Motif 6	7	7.2e-029	VSVRNPNMASFNYSNSTMQIFYHGMVIGEAQTPGGRVEARCTQYMNVT	48
Motif 7	4	1.1e-023	DFTGVATDLMSLNSTVKLIFRNPATFFGVIVSSTFIDLHYS	41
Motif 8	7	1.8e-020	GQMSISSYARIPGKVKLLHVFKHHVVSKMCSMNIDISSHSIQDQWC	47
Motif 9	4	7.2e-019	DNSQKMSYHAGEAKGQTQEKVSNMMDTASEKAQEAKEYAQETGQQMMEKA	50
Motif 10	5	4.6e-025	EALBEEIAHQAGEKPFVEQSDAEAYQEMERRATGSNMIIPGGIRAEQSAET	50

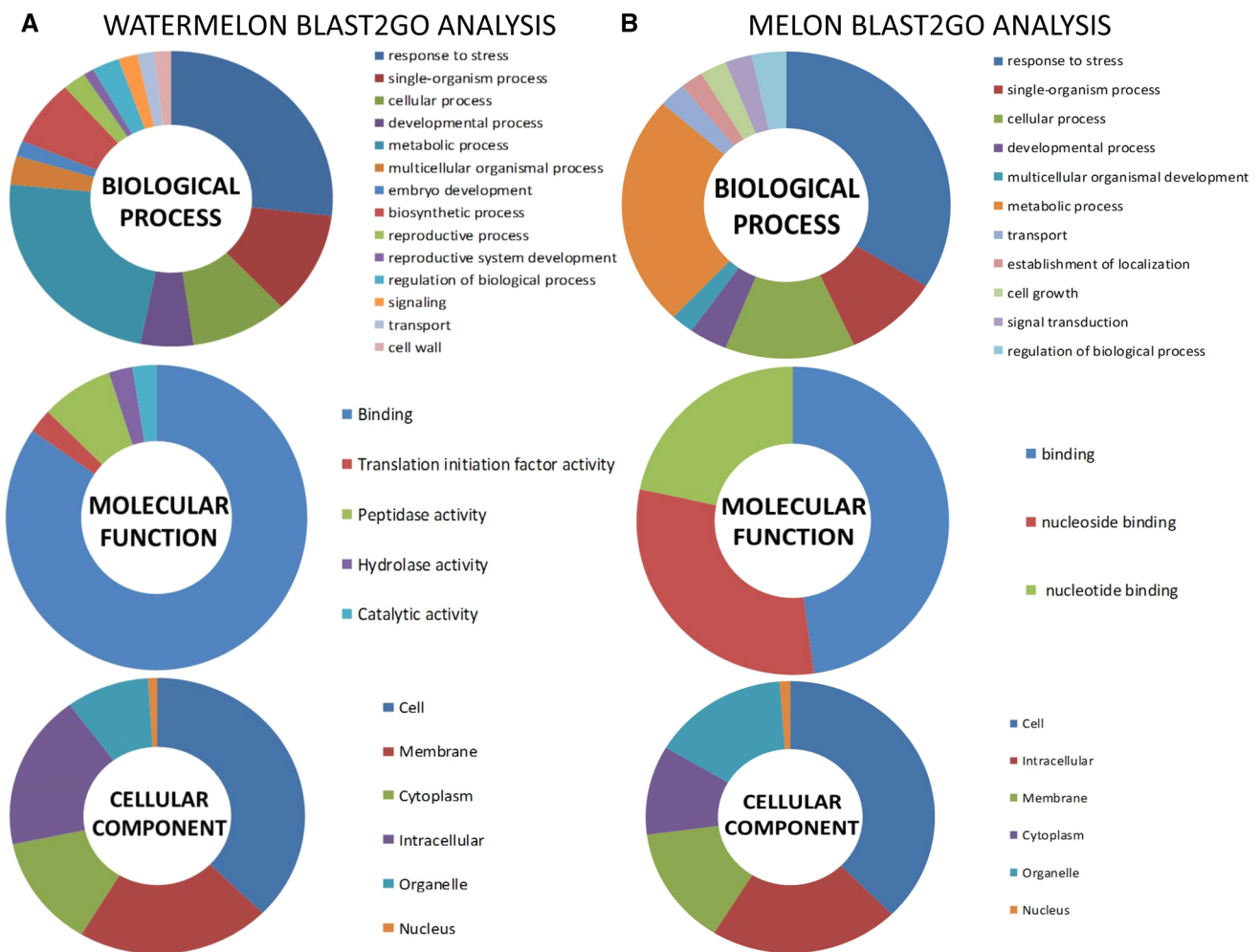


Fig. 3 Gene ontology analysis of **a** *CILEA* and **b** *CmLEA* genes by Blast2Go program with three categories named as biological process, molecular function and cellular component

dehydrin proteins in Chinese plum and potato in which conserved motifs called K (a lysine-rich 15 amino acid sequence) and Y segments were reported before (Charfeddine et al. 2015; Du et al. 2013). This can be attributed to conserved structure of dehydrin group LEA proteins in the Cucurbitacea family.

Gene ontology analysis

Go slim analysis was utilized to determine possible molecular function, cellular localization and biological roles of LEA proteins in melon and watermelon by Blast2GO software (Fig. 3; Supplementary Table 5; Supplementary Table 6). Cellular and membrane localization ratios were higher between *CILEA* and *CmLEA* proteins which may be compatible with their functions on protection of membranes and enzymes to maintain cellular functions under abiotic stress conditions (Goyal et al. 2005; Kosová et al. 2007; Reyes et al. 2008; Tolleter et al. 2010).

In addition, our findings are relevant with the studies which reported that LEA proteins localized commonly in sub-cellular regions such as cytoplasm, mitochondria, nucleus and chloroplast in Arabidopsis, tomato and purple false brome (Hundertmark and Hinch 2008; Filiz et al. 2013; Cao and Li 2015). This may suggest that there is a strong relation between the subcellular localization and the role of the LEA protein in the cell.

Binding to different molecules was the dominant activity for the action of *CILEA* and *CmLEA* proteins as molecular function. Binding of LEA proteins to nucleic acids to protect cellular structures by constructing hydrogen network was reported, which is related to the roles of LEA proteins in stress tolerance (Wolkers et al. 2001; Shih et al. 2004). In addition, membrane stabilization of this protein family members by association with phospholipid vesicles and sugars were demonstrated in model membranes under dehydration conditions (Wolkers et al. 2001; Kosová et al. 2007; Liu et al. 2010; Tolleter et al. 2010).

Binding activity as a molecular function of LEA proteins may be suggested the mode of action of these group of proteins when considering about these reported roles in stress tolerance.

In addition, response to stress is the most observed process when we analyzed biological roles of CILEA and CmLEA proteins and this role was followed by the roles of LEA proteins in metabolic and cellular processes in melon and watermelon. These biological roles were consistent with the reported roles of LEA proteins such as membrane maintenance, enzyme and nucleic acid preservation and oxidant scavenging activity etc. to protect cell structures from the detrimental effects of abiotic stress factors (Hara et al. 2001; Wolkers et al. 2001; Shih et al. 2004; Goyal et al. 2005; Kosová et al. 2007; Reyes et al. 2008) Besides, reports about stress tolerance by overexpression of *LEA* genes in transgenic plants and bacteria can support the defined biological role of *LEA* proteins in our study (Puhakainen et al. 2004; Liu and Zheng 2005; Duan and Cai 2012).

Orthologous *LEA* genes between watermelon and other species

LEA genes which physically mapped on watermelon chromosomes and those of mapped on Arabidopsis, soybean, poplar, potato, grape and maize chromosomes were compared to evaluate orthologous relationships between them (Supplementary Fig. 5). Approximately 41% of *CILEA* genes had orthologous genes with these plants. Maximum number of orthologous genes were with poplar with the ratio of 71%. These results are consistent with our group's previous study which found that cucumber *LEA* genes distributed maximum orthology with *LEA* genes in poplar, which were on the fourth chromosome of poplar (Altunoglu et al. 2016). Detailed analysis revealed that the maximum orthologous relationships were observed between *LEA* genes located on the first chromosome of watermelon and *LEA* genes located on the fourth chromosome of poplar. According to these results, *LEA* genes which belong to Cucurbitaceae family members distributed maximum orthology with the *LEA* genes accumulated on the fourth chromosome of poplar. These results can be useful for the determination of orthologous gene pairs between different organisms and for the evaluation of *LEA* gene family distributions. In addition, these findings suggest that gene duplication and inversion events may have important roles on expansion and organization of this gene family members in watermelon, Arabidopsis, rice, soybean, poplar, potato, grape and maize.

Duplication events and divergence rates of the *CILEA* and *CmLEA* genes

Tandem or segmental gene duplications and transposition events are the reason of the gene family broadening and generation (Cao and Shi 2012). We calculated the approximately dates of duplication events of *CILEA* and *CmLEA* genes to understand broadening ranges of this gene family members. Thus, nonsynonymous (K_a) versus synonymous (K_s) substitutions (K_a/K_s) were estimated for *LEA* genes which showed duplication events in watermelon and melon. In addition, these calculations were repeated for orthologous *LEA* gene pairs between Arabidopsis, rice, soybean, grape, poplar, potato and maize in watermelon (Fig. 4a) and melon (Fig. 4b). The K_a/K_s ratios were between 0.0038 and 0.2503 for segmentally duplicated *LEA* genes and between 0.006 and 0.374 for tandemly duplicated *LEA* genes in watermelon (Supplementary Table 3). Analysis revealed that these segmental and tandem duplication events may occur approximately between 10 and 600 MYA (million years ago) and 3 and 437 MYA, respectively. In addition, the average K_a/K_s ratio was the maximum between watermelon and poplar with the ratio of 0.15. The earliest divergence time of these gene orthologous from other plants was with maize *LEA* genes with the average number of 237 MYA (Fig. 4a; Supplementary Table 7).

Additionally, K_a/K_s ratios were 0.136 and divergence dates between duplicated *LEA* genes of melon varied from 2 to 550 MYA. K_a/K_s ratios of melon with Arabidopsis, soybean, rice, poplar, potato, grape and maize were with the average of 0.06, 0.081, 0.02, 0.07, 0.14, 6.65 and 0.03, respectively. The earliest divergence time was with rice with the average of 240 MYA (Fig. 4b; Supplementary Table 8).

According to these results, a vigorous purifying selection effected melon, watermelon and poplar. In addition, divergence times between cucumber and poplar were closer to date with the average of 78.4 MYA (Altunoglu et al. 2016) than other plants as melon and watermelon. The average of K_a/K_s ratio was 0.14 for tandem duplications in cucumber. All K_a/K_s ratios were observed as <1 in tandemly or segmentally duplicated *LEA* genes in Cucurbitaceae family members. These results may again show that *LEA* genes in Cucurbitaceae family members exposed to a large purifying selection. In addition, poplar and Cucurbitaceae family members may be closer genetically than other analyzed plants according to *LEA* gene analysis.

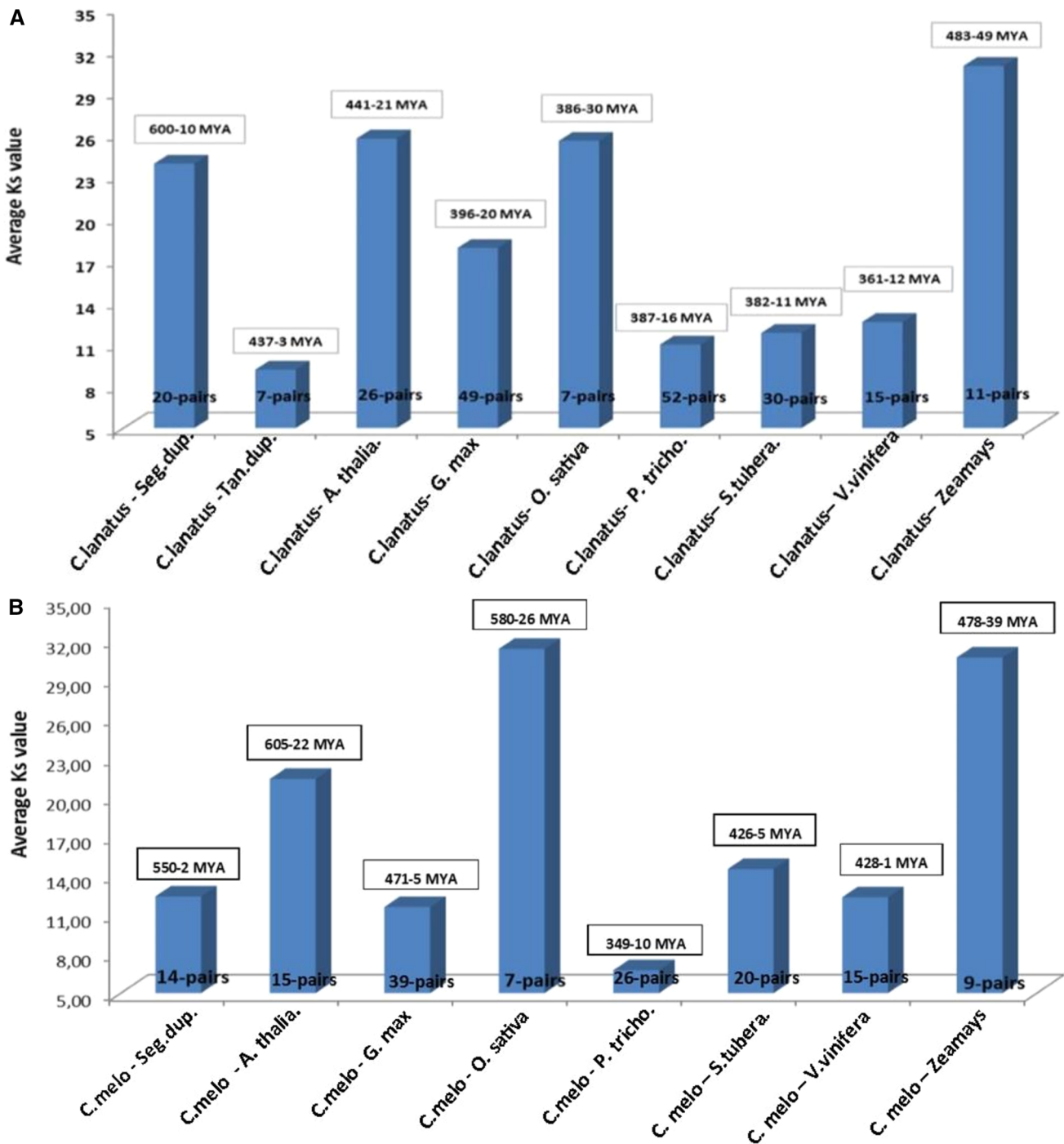


Fig. 4 Estimation of duplication and divergence times of **a** *CILEA* and **b** *CmLEA* genes using duplicated *LEA* genes of melon and watermelon and orthologous *LEA* gene pairs between watermelon–melon and Arabidopsis, soybean, rice, poplar, potato, grape and maize

MicroRNA (miRNA) identification for *CmLEA* and *CILEA* transcripts

MicroRNAs (miRNAs) are small, noncoding regulatory elements in organisms. Hence, they have important roles in gene regulation under stress conditions via destroying

target gene transcripts in plants (Bartel 2004; Ambros and Chen 2007).

miRNAs which targeted *CmLEA* and *CILEA* gene transcripts were identified by psRNA Target Server (<http://plantgrn.noble.org/psRNATarget/>). This server affords two important analyses: (1) reverse complementary matching is

achieved between miRNA and target transcript by scoring scheme of miRU (Dai and Zhao 2011). The threshold of the score is maximum expectation for which cut off threshold was set as 3.0 in the study. (2) Target site accessibility analyses via calculating unpaired energy (UPE) which is necessary to open secondary structure around miRNA's target site on mRNA. Connection of miRNA with target mRNA is going to be easier if needs low UPE.

Analyses revealed that 22 different *CILEA* transcripts (CILEA-06-10-11-13-14-15-17-19-20-33-38-39-42-45-46-52-58-63-64-67-72-73) were targeted by 24 different miRNAs in watermelon (Supplementary Table 9) and 21 different miRNAs targeted 21 different *CmLEA* transcripts (CmLEA-03-04-10-16-18-19-20-26-27-29-30-35-37-40-41-48-49-52-55-56-60) in melon (Supplementary Table 10). Mir854, mir414 and mir2673 were the common miRNAs and the most targeted gene was CILEA-63 in watermelon. CmLEA-20-55 were the most abundant target transcripts and mir854, mir5021 and mir2629 were common miRNAs in melon. Proteins targeted by mir414 in Arabidopsis especially had roles in transcriptional regulation, protein modification, DNA repair or chromatin modification. In particular, mir414 mainly affects regulators of transcription such as bZIP transcription factors, WRKY, MYB, B3, scarecrow, heat shock proteins and TCP. In addition, this important miRNA had roles in posttranscriptional modifications of SNF2 transcriptional regulator, F-Box family proteins, SNF2 transcriptional regulator etc. (Eulgem et al. 2000; Gurley 2000; Jakoby et al. 2002; Suo et al. 2003; Zhang et al. 2003; Flaus et al. 2006; Guo et al. 2007; Guleria and Yadav 2011). This may be indicated that mir414 is very important in the regulation of plant growth. Besides, upregulated expression of another important miRNA family called miRNA854 were reported under water deficiency conditions in rice (Zhou et al. 2010). Mir854 family was present in both plants and animals and this suggests that these miRNAs have a common origin as regulators of basal transcriptional mechanisms (Arteaga-Vázquez et al. 2006). Additionally, a previous study reported that MYB transcription factor which had a key role in regulatory networks controlling development and in responses to stress factors. It was targeted by mir5021 in rosy periwinkle (Pani and Mahapatra 2013). Mir414 and mir854 were found the most abundant miRNAs in cucumber in an earlier study (Altunoglu et al. 2016). This suggested that *LEA* gene family members in Cucurbitaceae family are especially targeted by miRNAs which contributed to the transcriptional regulation. In addition, this may be attributed to the important roles of *LEA* genes like transcription factors in these plants. Therefore, determination of miRNA targets of melon and watermelon *LEA* genes will be useful to elucidate the roles of these genes in melon and watermelon.

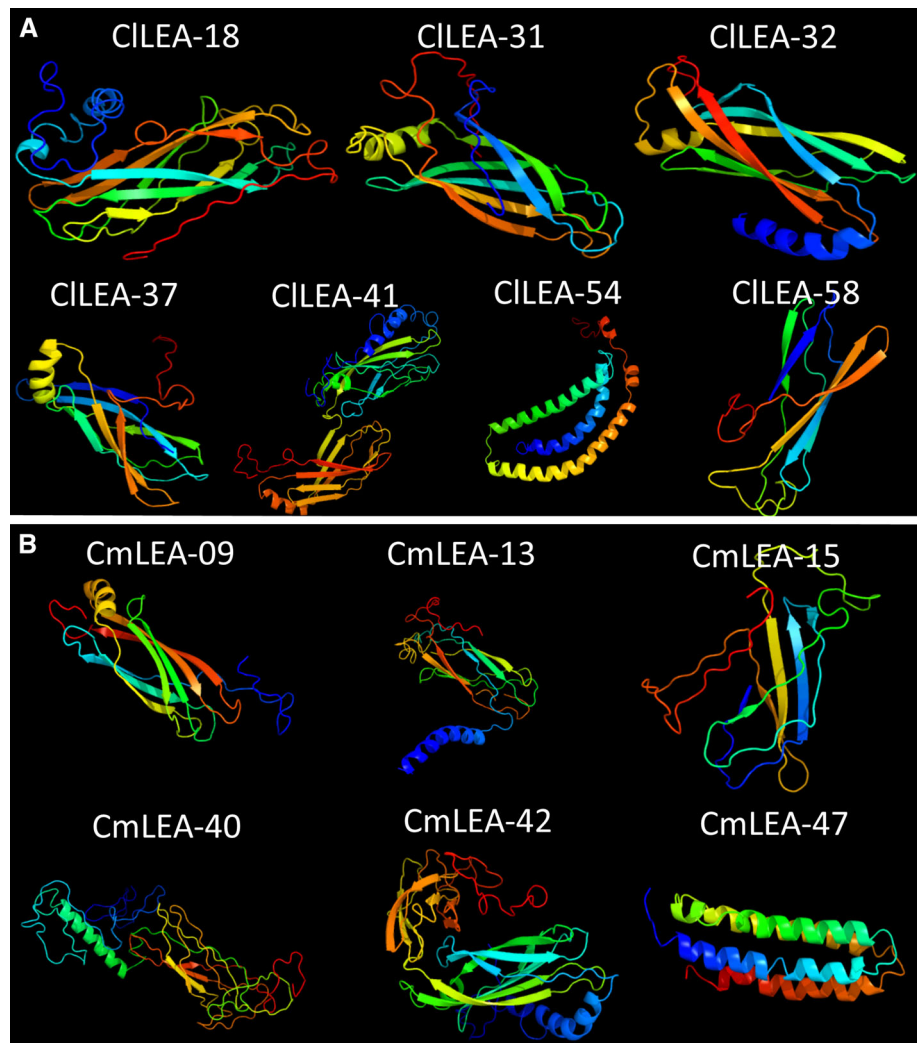
Estimation of predicated 3D-structures of *CILEA* and *CmLEA* proteins

PDB was used by BLASTP search for homology modeling. Seven proteins in watermelon [CILEA-18 (PDB ID:c1yycA), CILEA-31 (PDB ID:d1xo8a), CILEA-32 (PDB ID:d1hx6a1), CILEA-37 (PDB ID:c1yycA), CILEA-41 (PDB ID:d1xo8a), CILEA-54 (PDB ID:c1xq8A), CILEA-58 (PDB ID:c3butA)] (Fig. 5a) and six proteins in melon [CmLEA-09 (PDB ID:d1xo8a), CmLEA-13 (PDB ID:c1yycA), CmLEA-15 (PDB ID:c3butA), CmLEA-40 (PDB ID:d1xo8a), CmLEA-42 (PDB ID:d1xo8a), CmLEA-47 (PDB ID:d1 eq 1a)] (Fig. 5b) which showed high homology were chosen. Detection rates were used in Phyre2 tool with the selection of Hidden Markov Model (HMM) to predict protein structure (Kelley et al. 2015). Percentage of residues modelled were between 73 and 99 for watermelon and between 72 and 99 for melon with the >90% confidence interval. Interestingly, selected proteins from melon and watermelon were in LEA2 group. β sheets were the dominant structures between selected melon and watermelon *LEA* proteins. However, CmLEA-47 and CILEA-54 proteins contained only α helix chains. Predicated structures of cucumber *LEA* proteins distributed structures with predominant β sheets (Altunoglu et al. 2016) and these findings were compatible with the predicated structures of *LEA* proteins in melon and watermelon. This can be attributed to the membership of these plants in the same family. In addition, a previous study suggested that LEA2 group proteins especially had two β sheets and one alpha helix structure, which was consistent with our results. This type of structure resembles the structure of fibronectin Type III residues on the surface of animal cells and this kind of configurations may be important in fluid loss to protect cellular structures from the effects of stress (Li and Cao 2016; Singh et al. 2005). Predicated three-dimensional structures of this proteins analysis may provide to elucidate molecular activity of them to understand their roles in the cell.

Genome-wide expression profiles of *CILEA* and *CmLEA* genes and drought responsive *LEA* genes in watermelon and melon

SRA (Sequence Read Archive) data sets related to melon and watermelon were used to evaluate tissue specific expression patterns of *CmLEA* and *CILEA* genes by RNA Seq approach. Hierarchically clustered heat map was constructed according to transcriptome data from different days of fruit development stages after pollination and phloem and vascular tissues (Supplementary Fig. 1). Expression levels of *CILEA-7-36-44* genes were upregulated in 10 days old white fruit stage while they were

Fig. 5 Predicated three dimensional structures of **a** seven LEA proteins of watermelon and **b** six proteins of melon with 90% confidence level



downregulated in 18, 28 and 34 days old fruit development stages. Besides, expression levels of *CILEA-10-13-16-17-23-29-31-38-43-45-46-50-53-63-70* genes showed a tendency to increase in 10 days old white fruit stage but expression levels of these genes decreased in other stages. Contrast to this, *CILEA-11-34-69* gene expressions showed a decrease in 10 days old white fruit stage while their gene expression levels increased in other fruit stages. In addition, *CILEA-7-31-32-36-40-43-44-47-50-51-52-54-55-73* gene expressions were upregulated in the 10th, 18th and 26th days of fruit development when compared with the expression levels of these genes in the 42nd and 52nd days of fruit development stages. In addition to this, transcriptome data from phloem and vascular tissues of watermelon was investigated. Expressions of *CILEA-7-8-10-16-23-26-31-32-36-37-42-54-59-72* genes were higher in phloem tissues than in vascular tissues.

According to our qRT-PCR results, studied all *LEA* genes responded to drought in leaf tissues (Fig. 6). Especially *CILEA-12-17-46-48* genes expression levels were

upregulated at the 3rd hour of drought stress application. *CILEA-17* gene was highly expressed in leaf tissues in all studied hours of water deficiency. Besides, expression levels of *CILEA-12-41-48* genes were high at the end of 24th hour of stress application in root tissues. In addition, *CILEA-41* was the only gene among the selected genes, whose expression level was upregulated in root and leaf tissues at the end of 24th hour of the stress treatment.

CILEA-17 displayed an increase pattern in phloem and vascular tissues, 10th and 42nd days of fruit development and under drought stress conditions in our study when compared to the qRT-PCR data and the heat map. This may be attributed to its key predicated role in many metabolic processes and in water deficiency conditions. Same situation was observed for *CILEA-12* gene which plays important role in both metabolic processes and stress tolerance. In addition, *CILEA-41* gene had role in fruit development according to transcriptome data and upregulated under drought stress conditions. In addition, 26th *LEA* gene in Arabidopsis (*At2g44060*), which was the orthologous of

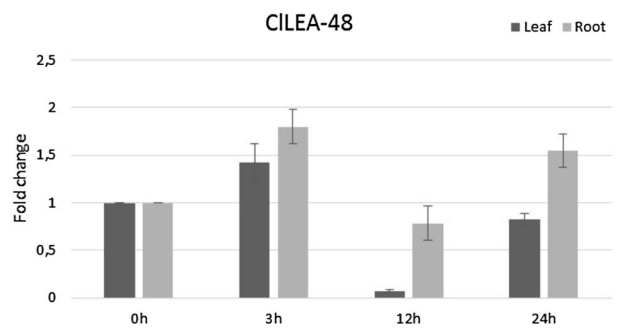
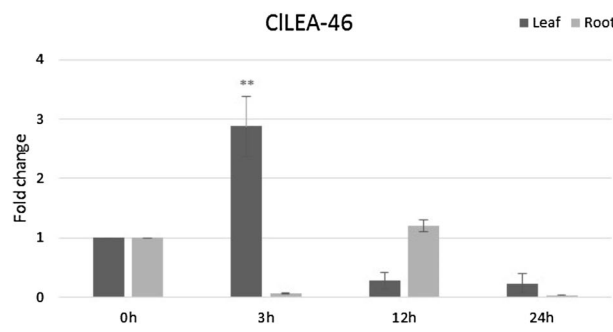
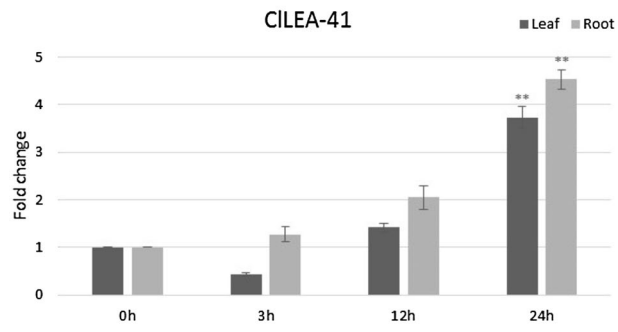
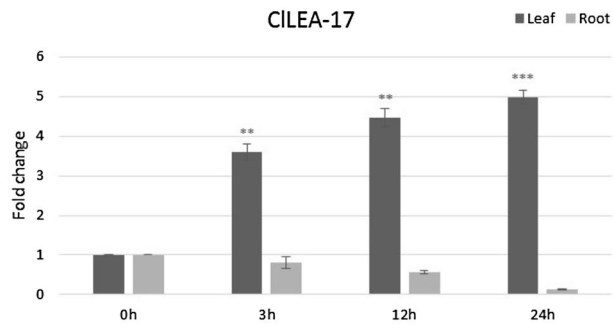
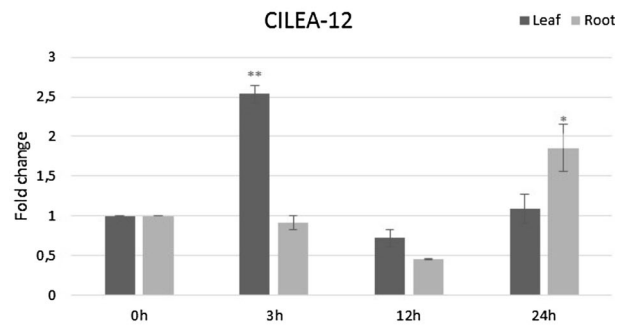
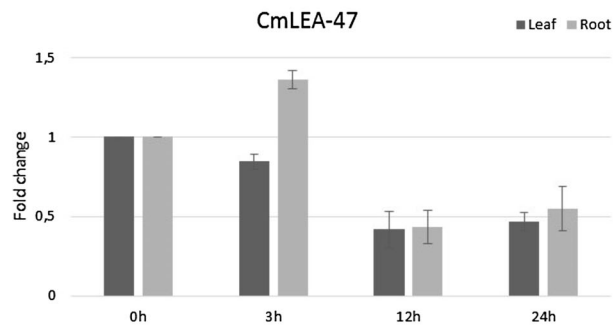
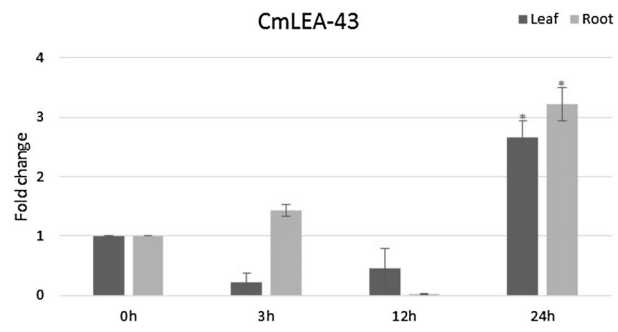
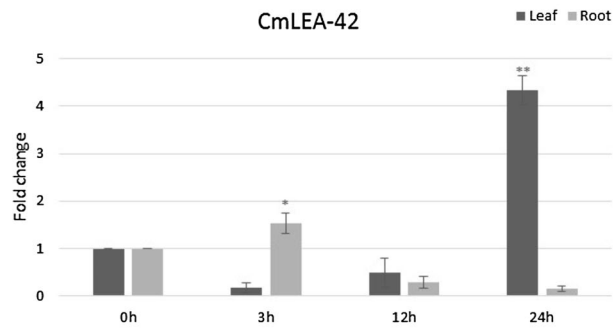
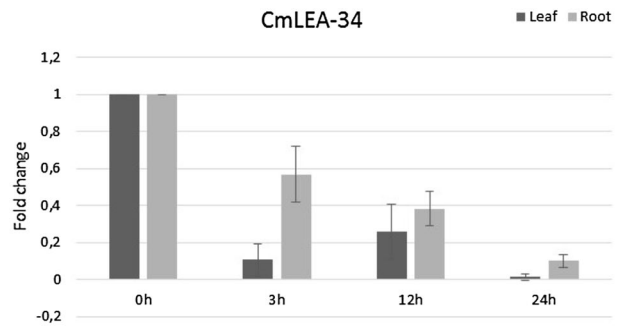
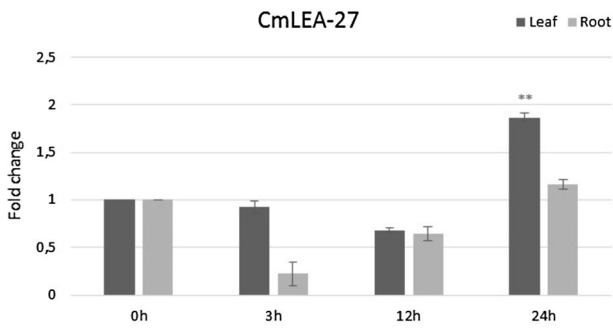


Fig. 6 Expression patterns of selected *CILEA* and *CmLEA* genes under water deficiency conditions in leaf (black) and root (grey) tissues (* means $p < 0.01$; ** means $p < 0.05$; *** means $p < 0.001$)

CILEA-41 gene, had high expression levels in non-seed tissues; also, it was induced by different stresses. Besides, 5th *LEA* gene (At1g20450) and 10th *LEA* gene (At1g76180) in Arabidopsis, which were orthologous of *CILEA-48* gene, were upregulated in different stress conditions. Especially, At1g20450 gene was induced by cold and salt stress conditions (Hundertmark and Hinch 2008). Our results are convenient with these findings, which suggests that *CILEA-41-48* genes and their orthologous in Arabidopsis were stress related genes in these plants. *CILEA-46* gene was related with fruit development while its expression was upregulated in vascular tissues and early in leaf cells under water deficiency. According to these findings, *CILEA-12-17-46* genes can be early response genes in drought stress.

Transcriptome data belonging to different melon varieties, fruit stages and salt stress conditions from SRA database were analyzed to evaluate expression patterns of *LEA* gene family members in melon (Supplementary Fig. 2). Gene expression levels of *CmLEA-1-2-7-8-19-28-31-36-50-51-53-55-60* were upregulated in two melon varieties named as Cantalupo vedrantaïs and Piel de sapo pinonet when compared with expression levels of these proteins in Conomon SC and Piel de sapo-t111 melon varieties. In contrast, *CmLEA-4-5-10-11-12-15-17-23-26-30-32-33-38-40-42-44-45-46-47-48-56-58-61* genes were highly expressed in Conomon SC and Piel de sapo-t111 varieties than in Cantalupo vedrantaïs and Piel de sapo pinonet varieties. Besides, gene expression levels of *CmLEA-2-3-4-6-12-22-23-25-26-27-28-29-31-34-37-38-40-41-47-50-51-53-54-58* decreased while *CmLEA-1-7-9-10-11-18-19-20-21-24-30-35-39-43-52-55-57-59-60* expression levels increased under salt stress conditions. *CmLEA-43* gene was highly expressed in leaf and root tissues under drought stress in our study (Fig. 6). In addition, according to heat map its expression increased in normal vegetative tissues and under salt stress conditions. This gene may be one of the primary response genes to stress; also it may be important in normal cell functions. Expression levels of *CmLEA-42* gene was the highest one in leaf and root tissues between analyzed *LEA* genes under drought stress. In addition, its expression was up regulated in yellow fruit stages when compared with the white fruit stages and a little increase was observed under salt stress based on heat map. 26th *LEA* gene in Arabidopsis (At2g44060), which was orthologous of *CmLEA-42* gene, was upregulated by different stresses (Hundertmark and Hinch 2008). Besides, other gene orthologues of *CmLEA-42* gene in maize [*ZmLEA8* (GRMZ2G053637_T01)] and

ZmLEA10 (GRMZM2G352415_T01)] were upregulated in seed, innermost husk, silks or anthers according to transcriptome data (Li and Cao 2016). These results can support our results and it can be claimed that *CmLEA-42* gene have a function in normal development stages besides its role in stress like function of *CmLEA-43*. However, *CmLEA-34* gene was down regulated under water deficiency and salty conditions and compatible with transcriptome data, so this gene is not seemed to be related with stress.

Present study showed that some melon and watermelon *LEA* proteins have roles in protecting these plants from the effects of abiotic stresses while they have roles in normal cellular functions and fruit development stages which correlated with its abundance in maturing seeds (Galau et al. 1986; Manfre et al. 2006). Previous studies showed that gene expression of a *LEA* gene in citrus belong to *LEA5* group highly increased in fruit than in leaves (Jeon et al. 2006) and also *LEA5* gene expression were reported in cultured cells under freeze, water deficiency and salty conditions (Naot et al. 1995), which are relevant with our findings in melon and watermelon. Besides, expression of some *LEA* genes [*STDHN1*(*YSK2*) and *StDHN25* (*SK3*)] in potato were reported in all tissues (Charfeddine et al. 2015), which may be related with their function in normal plant development stages, and these findings may support our results.

Present study provides a detailed molecular information about *LEA* gene family members in melon and watermelon. *CILEA-17* and *CmLEA-42* genes seems to be important for transgenic interventions to determine their roles. Our results can open up new frontiers about understanding of functions of these important family members under normal and stress conditions by cloning and functional approaches.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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