Characterization of ASR gene and its role in drought tolerance in chickpea (Cicer arietinum L.)

Abstract

Chickpea has a profound nutritional and economic value in vegetarian society. Continuous decline in chickpea productivity is attributed to insufficient genetic variability and different environmental stresses. Chickpea like several other legumes is highly susceptible to terminal drought stress. Multiple genes control drought tolerance and ASR gene plays a key role in regulating different plant stresses. The present study describes the molecular characterization and functional role of Abscisic acid and stress ripening (ASR) gene from chickpea (Cicer arietinum) and the gene sequence identified was submitted to NCBI Genbank (MK937569). Molecular analysis using MUSCLE software proved that the ASR nucleotide sequences in different legumes show variations at various positions though ASR genes are conserved in chickpea with only few variations. Sequence similarity of ASR gene to chickpea putative ABA/WDS induced protein mRNA clearly indicated its potential involvement in drought tolerance. Physiological screening and qRT-PCR results demonstrated increased ASR gene expression under drought stress possibly enabled genotypes to perform better under stress. Conserved domain search, protein structure analysis, prediction and validation, network analysis using Phyre2, Swiss-PDB viewer, ProSA and STRING analysis established the role of hypothetical ASR protein NP_001351739.1 in mediating drought responses. NP_001351739.1 might have enhanced the ASR gene activity as a transcription factor regulating drought stress tolerance in chickpea. This study could be useful in identification of new ASR genes that play a major role in drought tolerance and also develop functional markers for chickpea improvement.

Introduction

Chickpea (Cicer arietinum L.), one of the earliest food legume crop with a diploid chromosome number of 16 is cultivated in the tropics all over the world [1] and belongs to the family Fabaceae [2]. India being the largest producer of chickpea produces 68% of the total world
production and about 9.21Mha area is under chickpea cultivation producing 8.88Mt [3]. Chickpea, characterized by different desi and kabuli cultivars has a profound nutritional and economic value [4]. A considerable decrease in chickpea productivity has been observed in the last thirty years due to change from lower to higher temperature regions of cultivation in South-East Asia and East Africa [5]. Presently, the world average productivity is about 995 kg/ha which is very low [3] and has stagnated in recent years due to vulnerability of chickpea crop to various abiotic (drought, terminal heat, high salt, cold stress), and biotic (Ascochyta blight, Fusarium wilt, Helicoverpa) stresses [6]. Average losses upto 60% have been reported due to abiotic stresses globally in chickpea [7]. Drought drastically affects the plant growth processes and reduces plant yield [8]. Development of chickpea varieties tolerant to drought has been very slow due to its narrow base and limited genomic resources [9] necessitate improving its genetic potential [10]. Plants combat these stresses through a series of physiological mechanisms controlled by several stress related genes which in-turn is regulated by specific transcription factors [11–13].

Drought is a genetically complex trait [14]. Among various transcription factors, abscissic acid is involved in signaling drought stress in general and their levels have a direct effect on different parts of the plant or plant as a whole [15]. Drought tolerance may be governed by some pathways that are ABA dependent [16] or pathways that are independent of ABA [17]. Particularly, transcription factors of the Asr (abscisic acid, stress, ripening) family of genes are expressed only in plants that interact with ABRE elements as a regulatory mechanism of ABA dependent pathways under stress [18, 19]. The major domain is of Pfam family which is an ABA/WDS domain. This group of genes is a part of regulating complex wherein they play a major role. Particularly in processes involving metabolism of sugars like in fruit ripening, maturation of pollen, senescence and differential responses to various abiotic stresses like drought, salinity, reduced light intensity, cold [20–25]. The protein products of this gene act like chaperons and help in preventing thawing and freezing type denaturation [26]. ASR genes are expressed in different organs viz., potato tubers [27], and fruits of tomato, apricot and pomelo [28], pollen of lily [29], leaves and roots of tomato, rice, maize and pine [30,31]. All ASR genes known till date have a DNA binding activity at the N-terminus that is sequence specific and dependent on Zn$^{2+}$ with a nucleus localization signal at the C terminus. Sub-cellular fractionation studies also proved that ASR protein occur in the nucleus and cytoplasm. The potential role of ASR1 gene in drought tolerance in common bean was studied and strong selection pressures, lower gene diversity was found in the accessions [32]. Transgenic Arabidopsis with over-expressed ASR gene showed an increase in tolerance to drought and salt and decrease in sensitivity on exposure to exogenous ABA [33]. ASR3 gene was identified as a putative candidate gene for association mapping for tolerance to drought in rice [34]. So far, the characterization studies on ASR genes in chickpea are limited.

Huge genome sequence information available online in public domains (http://www.ncbi.nlm.nih.gov) have served as excellent sources for identification of important genes for insect resistance, quality traits, resistance to different abiotic stresses viz., drought, salinity, heat. Such computational studies are valuable in areas of comparative genomics and have enabled us to identify and characterize chickpea ASR genes. In the present study, single ASR homologue was identified in chickpea. Analysis of conserved domains, phylogenetic relationships, three dimensional structure prediction and validation, and functional partners within the query sequence identified a hypothetical protein NP_001351739.1, potentially involved in drought tolerance. The drought responses and relative expression levels of chickpea ASR genes under different treatments were also studied. Our study gives an idea about the role of ASR genes in drought tolerance in chickpea and also indicates similarities with the already characterized proteins which may be possibly used in improvement of chickpea and related pulse crops.
Results

Relative water content, chlorophyll, protein content and membrane stability index in control and water stressed plants

Ten genotypes were used to examine drought responses at different time points viz., control (0day), 6day and 12day as defined for chickpea [35]. Susceptible checks (ICCV2, Pusa 1003 and Pusa 362) were also included in analysis for comparative study. Variations in relative water content (RWC,%), chlorophyll index (CI, Spad Units), protein content (μg/ml) and membrane stability index (MSI, mS/cm) were measured to test their responses for improved drought tolerance and better understand their resistance mechanisms, presented in Table 1. There was a significant decrease in all the parameters measured from day 0–12 of drought stress in all the genotypes (Table 2). Control plants maintained a higher RWC (%), CI, protein content and MSI in comparison to drought stressed plants (Fig 1A–1D). Reduction in RWC (%) ranged from 8% to 28.3% in all the genotypes at 12th day after imposing drought stress (Fig 1A). Maximum decrease in RWC (%) was 28.3% in ICCV2; however, BGD72 (8.73%), ICCV10 (9.29%) and ICCV3311 (10.81%) maintained the RWC (%) under drought stress (Table 2 and Fig 1A). Drought stress affected the CI significantly, and the decrease was noteworthy in ICCV2 (20.31%) (Fig 1B). On the contrary, resistant genotypes BGD72 (6.77%), ICCV10 (7.5%) and ICCV3311 (7.57%), retained higher CI at 12th day of drought stress treatment (Table 2 and Fig 1C). Our results also signified a substantial decrease in soluble protein content (leaf) in all the chickpea genotypes under drought stress (Fig 1B). Though, BGD72 showed minimal decrease in protein content (7.89%) over the control, this decrease was prominent in ICCV2 (24.3%) (Table 2 and Fig 1B). Decline in Membrane stability index (MSI) was prominent in all the genotypes on imposition of drought stress (Fig 1D). BGD72 (7.69%), ICCV10 (9.09%) and ICCV3311 (12%) showed minor decrease in MSI under stressed conditions in comparison to control conditions. Conversely, the susceptible genotypes viz., ICCV2 (29.62%), Pusa362 (29.15%) and Pusa1003 (28.57%) displayed a much higher reduction in MSI at 12th day of drought stress treatment (Table 2 and Fig 1D). Significant changes observed from control to stressed samples demonstrate stress at morpho-physiological and biochemical levels in all the genotypes.

Quantitative real-time PCR (qRT-PCR) analysis

To analyze the expression pattern of ASR gene under drought stress in selected genotypes of chickpea, real time quantitative PCR was performed. The Beta Actin gene was used as the reference gene. Samplings were done in triplicates at each time point (0day, 6th day and 12th day after drought stress treatment). The mean fold change in the ASR gene, normalized to Beta Actin gene at different time points was calculated by Ct (cycle threshold) values (S1 Table of S1 File). Normalization with Actin gene produced more consistent and similar results in drought

| Source of Variation | df | RWC | CI | Protein | MSI | F crit |
|---------------------|----|-----|----|---------|-----|-------|
| Treatment           | 2  | 376.428** | 99.351** | 40.72* | 386.881** | 3.354 |
| Error               | 27 | 111.155 | 15.891 | 8.771 | 114.133 |       |
| F value             |    | 3.386 | 6.251 | 4.642 | 3.389 |       |

** Significance at p≤0.05

https://doi.org/10.1371/journal.pone.0234550.1001
Table 2. Mean values of relative water content (RWC, %), leaf protein content, chlorophyll index (CI, Spad units) and membrane stability index (MSI, mS/cm) of the ten chickpea genotypes under drought stress conditions.

| Genotype   | Physiological parameters |
|------------|--------------------------|
|            | RWC (%)                  | Protein (leaf) (μg/ml) | CI (Spad units) | MSI (mS/cm) |
|            | C 12D                    | C 12D                  | C 12D          | C 12D      |
| ICCV97309  | 73.80^a                  | 28.10^a                | 52.30^a        | 74.00^a    |
|            | 64.00^a                  | 23.40^a                | 50.36^a        | 64.20^a    |
| ICCV3311   | 74.00^a                  | 28.93^a                | 54.50^b        | 75.00^b    |
|            | 66.00^b                  | 25.53^b                | 50.36^b        | 66.00^b    |
| ICCV10316  | 66.00^b                  | 25.69^b                | 53.10^a        | 67.00^b    |
|            | 50.00^c                  | 21.36^c                | 44.02^c        | 50.00^c    |
| ICCV9307   | 69.00^d                  | 26.85^d                | 52.95^a        | 69.00^d    |
|            | 54.00^e                  | 22.96^e                | 47.11^d        | 60.00^d    |
| BGD 72     | 76.70^f                  | 30.40^f                | 55.96^d        | 78.00^e    |
|            | 70.00^g                  | 28.00^g                | 52.16^e        | 72.00^e    |
| ICCV10     | 74.97^e                  | 30.99^e                | 55.65^d        | 77.00^f    |
|            | 68.00^f                  | 27.00^f                | 51.46^d        | 70.00^f    |
| ICCV5313   | 69.00^f                  | 26.05^f                | 51.34^e        | 68.00^g    |
|            | 51.00^g                  | 21.83^e                | 45.13^f        | 52.00^f    |
| Pusa 1003  | 55.00^h                  | 25.48^h                | 47.00^d        | 56.00^h    |
|            | 40.50^i                  | 20.53^i                | 40.46^i        | 40.00^i    |
| Pusa 362   | 57.00^i                  | 24.78^i                | 50.06^g        | 43.66^g    |
|            | 42.00^j                  | 20.95^j                | 43.93^g        | 41.80^j    |
| ICCV2      | 51.00^k                  | 23.29^k                | 47.60^l        | 59.00^l    |
|            | 38.00^l                  | 17.62^k                | 37.93^l        | 41.80^l    |
| Mean       | 66.64                    | 27.06                  | 52.07          | 67.70      |
| CV (%)     | 84.51                    | 6.22                   | 9.53           | 21.55      |

Means followed by different letters within a column are significantly different from each other according to Tukey's Studentized Range (HSD) test at p ≤ 0.05; C: control conditions; 12D: 12th day after imposition of drought stress; CV: coefficient of variation.

https://doi.org/10.1371/journal.pone.0234550.t002

Fig 1. Drought responses of selected chickpea genotypes measured after different periods of stress (0, 6 and 12 days after drought stress treatment). Changes in relative water content (leaf) (A), Protein content (leaf) (B), Chlorophyll index (CI), MSI (D) of selected genotypes of chickpea. Samplings were done at 0day (control), 6th day and 12th day after imposition of drought stress. All the measurements were recorded in three replications and mean values were plotted against the selected chickpea genotypes.

https://doi.org/10.1371/journal.pone.0234550.g001
susceptible (ICCV2, Pusa1003 and Pusa362) and tolerant (BGD72, ICCV10 and ICCV3311) genotypes that clearly illustrated differential expression of transcripts of ASR gene (Fig 2A). In general, with the induction of drought stress, the expression of ASR gene was up-regulated in all the chickpea genotypes over controls and reached maximum at 6th day, followed by a slight decrease at 12th day. Expression patterns revealed high and significant expression of Abscissic acid and stress ripening gene (ASR) in the drought tolerant genotypes (BGD72, ICCV10 and ICCV3311) in comparison to the susceptible genotypes (ICCV2, Pusa1003 and Pusa362) indicating its sensitivity to drought stress treatment (Fig 2B).

Highest relative expression value was observed for BGD72 (2.54 fold higher than the control) at 6th day after imposition of stress. The increase in ASR gene expression was pronounced in ICCV10 (up to 1.86-fold change, 6th day) and ICCV3311 (1.67-fold change, 6th day) followed by a small decrease in fold change to 1.49 by 12th day of drought stress. Decreased relative expression level was evident in ICCV2 (0.69-fold change) at 12th day after stress imposition. No significant differences in the relative expression level of ASR were observed from control and treated samples in Pusa1003 and Pusa362, though the induction was higher in ICCV97309, ICCV9307 and ICCV10316.

Amplification of ASR genes from chickpea

The ASR gene homologues (Fig 3) were isolated from Medicago gene sequence available at NCBI EST database (DbEST-http://www.Ncbi.nm.nih.gov/dbEST/) [36]. Sequences of the ASR genes ranged from 680bp to 700bp nucleotides. Conserved region was observed in all the chickpea genotypes with very minor variations (Fig 4). BLASTn results confirmed 99.32% similarity with chickpea putative ABA/WDS induced protein (LOC101493413), mRNA with E value of 0.00.

Molecular analysis of chickpea ASR homologue with other legume plants

Comparison of the chickpea ASR homologue with other legume plants available at NCBI database revealed conserved nucleotides at various positions (Fig 5A–5B). ‘A’ at position number

---

Fig 2. Differential expression of chickpea ASR genes under drought stress conditions. (A) Heat-map showing differential expression of chickpea ASR genes during drought stressed periods (0day, 6th day and 12th day) in selected genotypes of chickpea. The scale at the top represents log fold change, maximum value is displayed as dark red and minimum value is displayed as light green. (B) Real-time PCR analysis to validate the differential expression of chickpea ASR genes during drought conditions. The Beta Actin gene was used as a reference. Expression was measured after 0, 6 and 12 days after imposition of stress. In control, expression was recorded on day 0 of stress. The mean fold change in ASR gene expression at each time point was calculated using the 2^-ΔΔCT method where ΔΔCT = (C_{Target} - C_{Actin})_{Time x} - (C_{Target} - C_{Actin})_{Time 0}. Data are means ± SD of triplicate samples.

https://doi.org/10.1371/journal.pone.0234550.g002
Neighbour joining analysis

Multiple sequence alignment of ASR gene was done using MUSCLE software and a phylogenetic tree was constructed by Neighbour joining method with 1000 replications in bootstrap test using Treedyne software. The *Cicer arietinum* gene encoding putative ABA/WDS domain containing protein was grouped with *Medicago truncatula* glycine-rich cell protein encoding gene with bootstrap value 21 and *Phaseolus vulgaris* Asr genes, *Arachis hypogea* glycine rich TATA-binding protein encoding genes and *Cajanus cajan* POU domain class 4 transcription factor-1 gene with 100 bootstrap value. The *Glycine soja*, *Glycine max* Asr genes were grouped closer to *Vigna unguiculata* and *Vicia faba* sequences (Fig 6).

Target protein sequence analysis

Blastx results showed 99.44% similarity with putative ABA/WDS induced protein (*Cicer arietinum*) with accession ID NP_001351739.1 and E value 7e-22, the sequence of which was downloaded from NCBI database (Fig 7). The protein was reported from *Cicer arietinum* with 257 amino acids and has been found to encode a family of plant proteins induced by water deficit stress or abscisic acid (ABA) stress and ripening.
Structural analysis of the protein NP_001351739.1

Conserved domains NCBI CDD tool identified pfam02496 ABA/WDS induced protein and one superfamily ABA_WDS with an E-value of 9.03e-22 (Fig 8). Expasy Protparam tool assessed different characteristics of the predicted protein. The molecular weight of the

Fig 4. Jalview of multiple sequence alignment of ASR gene homologues of seven chickpea genotypes. Conserved region was observed in all the chickpea genotypes with very minor variations (www.ebi.ac.uk).

https://doi.org/10.1371/journal.pone.0234550.g004
predicted protein was found to be 27.1 KDa and isoelectric point 5.05. This domain comprised mainly of glycine (G) and asparagine (D) with 22.6% and 9.3% respectively. The atomic
composition of ABA-WDS domain with formula $C_{1151}H_{1661}N_{343}O_{428}S_{1}$ and 3584 atoms showed presence of 25 positively charged residues (Arg + Lys) and 46 negatively charged residues (Asp + Glu). The instability index of 27.62 indicated stability of the ABA-WDS domain induced protein.

Alignment and neighbour joining analysis

Alignment of conserved ABA/WDS domain induced protein and phylogenetic tree was constructed using Neighbour joining analysis (Fig 9). The phylogram divided the ASR proteins into three major clusters. Multiple sequence alignment showed *Cicer arietinum* ABA/WDS induced protein belonging to cluster I grouped with *Brachypodium distachyon* abscissic stress ripening protein 3 with bootstrap value 71. Closely related ABA/WDS induced proteins from other crop plants in the cluster II included *Triticum uratu* abscissic stress ripening protein with bootstrap value 98, putative bundle sheath specific protein_1_Os01g0963600 and homeotic protein female sterile protein *Oryza sativa Japonica* group with bootstrap value 93, hypothetical protein_Os1_15903_ group with bootstrap value 80, putative fruit ripening protein_Os01g0959100_protein and unnamed protein_Triticum aestivum with bootstrap value of 95.

Structural prediction and validation

The three-dimensional model of the hypothetical protein was constructed using Phyre2 database (Fig 10A). The results revealed that the protein comprised of only alpha helixes and had no β-sheets. Alpha helixes accounted for 35% of the total protein. The Psi-Phi plot showed that 72.2% amino acid with 184 residues were present in the most favored regions, 14.5% amino acids with 37 residues in the allowed regions and 13.3% amino acids with 34 residues in the

Fig 7. Target protein sequence downloaded from NCBI (http://www.ncbi.nlm.nih.gov). Blastx revealed 99.44% similarity with putative ABA/WDS induced protein (*Cicer arietinum*) with accession ID NP_001351739.1.

https://doi.org/10.1371/journal.pone.0234550.g007

Fig 8. Conserved domains in chickpea ASR gene homologue. NCBI tool for conserved domain search (CDD) identified ABA-WDS domain-containing protein and ABA_WDS superfamily induced by water deficit stress.

https://doi.org/10.1371/journal.pone.0234550.g008
disallowed regions (Fig 10B). ProSA-web revealed that the z-score of the protein was -3.59 and within the range of scores found for proteins of similar size (Fig 11). The energy plot indicating the energies as a function of amino acid position was also plotted (Fig 12A). The parts of the predicted model that contribute to the overall bad score were also indicated (Fig 12B).

**STRING analysis**

Protein-protein interactions of the hypothetical protein NP_001351739.1 were analyzed using STRING database and the network obtained is shown in Fig 13. The network analysis revealed our target protein interacts with ten different proteins for carrying out its functions (Fig 14).

**Discussion**

Pulses have a great potential to improve human health as a rich source of protein, soil health through nitrogen fixation and helps in attaining food and nutritional security. In recent years pulses are consistently coming under the centre stage of research focus because of its importance. The up swinging prices of pulses and the nutritional importance have forced the policy makers to pay attention towards pulses particularly in the vegetarian population and increasing vegetarian community worldwide. Year 2016 has been declared by the UNO as the "International year of Pulses". There has been a reduction in the pulse availability per person per day in the last 50 years from 70 grams/capita/day to 34 grams/capita/day (http://www.faostat3.fao.
The major challenges impeding the pulse production and productivity are narrow base in
the cultivated varieties, genotype and environment interaction, multiple biotic and abiotic
stresses, difficulty in screening and precisely identifying the target traits [37].

Chickpea with nutritionally essential components and lesser anti-nutritional factors has the
best composition among legumes [38]. Chickpeas are the major source of protein in vegetarian
society [4]. Chickpea is also called as gram or Bengal gram in common language. It is self-polli-
nated crop with out-crossing rate less than 1%. Two chickpeas desi (dark coloured seed coat)
and kabuli (white coloured seed coat) are known having varied gene pool [39]. Chickpea pro-
ductivity remained stagnated and low since many years partly because of numerous environ-
mental stresses and insufficient genetic variability in various traits due to the domestication
process [40]. Drought is one of the major abiotic stresses affecting the crop productivity all
over the world and chickpea like several other legumes is highly susceptible to terminal
drought stress. Reduction in yield parameters has been linked to the adverse effects of drought
stress on growth processes of plants viz., cell growth, biomass, leaf area index and plant yield.

Plants survive the environmental stresses and overcome the harmful effects of drought stress
with the help of numerous mechanisms. Phenomics assisted with genomic approaches appear
to be a dependable solution to decipher these mechanisms and identify solutions for combat-
ing drought at a fast pace and improving yield [41]. The present study describes comprehen-
sively the isolation and characterization of ASR gene. The studies on this gene and its
molecular function in drought tolerance are very limited in chickpea.

In the present study, seven chickpea genotypes were selected to examine drought responses
at 0day, 6th day and 12th day after stress treatment and variations in their physiological param-
eters were assessed for speedy characterization of their drought tolerance (Table 1). All the
chickpea genotypes showed a significant decline in RWC(%), CI, protein content and MSI
under stressed conditions in comparison to control conditions (Table 2 and Fig 1A–1D).

RWC was considered as the best measure for water status of a plant in mid 80s as it indicates
the balance between water absorbed and consumed through transpiration. Under stressed
conditions, reduction in RWC has been established at various stages of growth in chickpea.
Higher retention of water in tolerant genotypes under stress has been confirmed by many workers [42, 43]. Under drought stress, significant reduction in RWC (%) was observed in all the genotypes at different time points in comparison to controls (Table 2 and Fig 1A). Per cent decrease in RWC was found to be highest in ICCV2 (28.3%) followed by Pusa1003 (26.36%). BGD72 (8.73%), ICCV10 (9.29%) and ICCV3311 (10.81%) maintained a considerably higher RWC (%) in both control and stressed conditions enabling them to perform better in terms of physiological processes under stress. In contrast, ICCV2 (28.3%), Pusa1003 (26.36%) and Pusa362 (26.31%) showed maximum decrease in RWC signifying their vulnerability to drought stress. MSI indicates the cell membrane damage by measuring electrical conductivity of cell leachates under drought. Membrane injuries and leakage of electrolytes from the membrane triggers programmed cell death in plants and also assist in remobilization for seed development [44]. Reduction in membrane stability index (MSI) at different stages under moisture stress has also been confirmed in earlier studies [42]. The per cent decrease in MSI in the chickpea genotypes under stressed conditions ranged from 7–30% (Table 2 and Fig 1D). Many workers have also reported severe influence of drought stress on membrane thermo-stability, canopy temperature depression and yield traits mainly filled pods and seeds per plant [45]. Increased temperature under
drought stress damages the cell walls and increases leakage of electrolytes [44, 46]. Relatively lesser decrease in MSI was observed in BGD72 (7.69%), ICCV10 (9.09%) and ICCV3311 (12%) with higher RWC (%) signifying their tolerance to drought. RWC and MSI have been suggested for screening the germplasm for drought tolerance [2]. Drought stress also hinders photosynthetic machinery of plants by bringing about changes in the chlorophyll content [47]. Modification in total chlorophyll, phenolics and proteins of the plants may govern their drought tolerance [48, 49]. Significant reduction was observed in chlorophyll content of all the selected chickpea genotypes under stressed conditions. The percent decrease in chlorophyll index under stress ranged from 6–21%. The tolerant genotypes BGD72 (52.16), ICCV10 (51.46) and ICCV3311 (50.36) maintained high SPAD values in contrast to Pusa362 (43.66), Pusa1003 (40.46) and ICCV2 (37.93) that showed significant decrease under stressed conditions (Table 2 and Fig 1C). Diversity studies among 43 lentil genotypes confirmed significant variations for twelve different phenotypic traits and positive correlation between stable lines and SPAD index establishing use of chlorophyll index as a standard measure for tolerance to drought [50]. Considerable reduction of soluble proteins has been seen in chickpea varieties viz., Bivaniej and ILC482 and Pirouz under stressed conditions [31]. Remarkable changes in quantity and quality of soluble proteins under stress have been detected in chickpea [52]. The percentage reduction in soluble protein content (leaf) ranged from 7–25% in the selected genotypes of chickpea. Maximum soluble protein was found in BGD 72 (28 mg/ml) followed by ICCV10 (27mg/ml) and ICCV3311 (25.53mg/ml) having higher SPAD values whereas minimum protein was found in ICCV2 (17.63 mg/ml) followed by Pusa 362 and Pusa 1003 (20.96 mg/ml and 20.53 mg/ml, respectively) with lower SPAD values (Table 2 and Fig 1B). Progressive increase in water stress significantly decreased net photosynthesis rate and protein content in moth bean genotypes [53].

PCR amplicons were then isolated from seven chickpea cultivars and sequenced by gene specific markers [36] using an ABI3500xL genetic analyser (Applied Biosystems, USA). These
gene sequences were subjected to sequence alignment using MUSCLE software. The chickpea ASR homologue was compared with other legumes and the conserved nucleotides were identified. Jalview results proved that all the ASR nucleotide sequences in comparison with other legumes show evidence of variations at various positions in the sequence region and the

Fig 13. Protein-protein interactions of the target protein NP_001351739.1. STRING network analysis was done to study the interactions of our target protein (indicated in red) with other proteins (http://string-db.org/) for its functioning.

https://doi.org/10.1371/journal.pone.0234550.g013

...gene sequences were subjected to sequence alignment using MUSCLE software. The chickpea ASR homologue was compared with other legumes and the conserved nucleotides were identified. Jalview results proved that all the ASR nucleotide sequences in comparison with other legumes show evidence of variations at various positions in the sequence region and the

Fig 14. Predicted functional partners of the target protein NP_001351739.1. Functional partners required for its functioning of the target protein were predicted using STRING database signifying its role in drought tolerance.

https://doi.org/10.1371/journal.pone.0234550.g014
conserved sequence region comprising possibly the ABA/WDS domain (Fig 5A and 5b). The phylogenetic tree showing relatedness with other legumes was generated using Treedyne software by neighbor joining method. *Cicer arietinum* gene encoding putative ABA/WDS domain containing protein mRNA showed high similarity with *Phaseolus vulgaris* Asr genes, *Arachis hypogea* glycine rich TATA-binding protein encoding genes and *Cajanus cajan* POU domain class 4 transcription factor-1 gene with maximum bootstrap value 100. *Glycine soja*, *Glycine max* Asr genes were grouped in a different cluster closer to *Vigna unguiculata* and *Vicia faba* sequences (Fig 6) with high bootstrap values. Similar findings confirmed that *Cicer arietinum* gene family was closely related to *Medicago trunculata* while soybean, pigeonpea and common bean sharing a common ancestor grouped in a separate cluster [54].

The changes in the expression of chickpea ASR gene relative to the β-actin gene at different points (control, 6th day and 12th day after drought stress treatment) were studied using qRT-PCR. Results confirmed that drought stress significantly increased the ASR gene expression that possibly elicited increased responsiveness towards drought tolerance in chickpea. Expression patterns revealed high and significant expression of ASR gene in all the genotypes under drought stress. Expression of ASR gene was prominent in BGD72, ICCV10 and ICCV3311 at 6th day of stress compared with controls with a slight decrease at 12th day of stress, however, there was no major change in its expression in ICCV2, Pusa1003 and Pusa362 under stressed conditions. It has been proposed ASR1 gene expression increases under stress in a variety of species [55]. The results of qRT-PCR analysis of OsASR1 and OsASR3 expression revealed that drought stress mainly regulates the expression of all ASR genes in rice and their over-expression in transgenic plants improved their drought and cold stress tolerance. Transgenic *Brachypodium distachyon* L. plants over-expressing BdASR4 gene retained more water and displayed higher tolerance to drought and antioxidant activities in comparison to the wild plants [56]. Physiological studies also proved the involvement of ASR1 gene in stress tolerance in transgenic tobacco, tomato, maize and rice [57].

The ASR gene sequences were confirmed at protein level using BlastX and Expasy server (http://expasy.org/cgi-bin/protparam). BlastX results showed 100 per cent similarity with chickpea hypothetical ABA/WDS induced protein (NP_001351739.1) with highest score of 486 and lowest Evalue of 0.00. The conserved domain search tool in NCBI identified a pfam02496 ABA/WDS induced protein and an ABA/WDS superfamily with a low E-value of 9.03e-22. Expasy tool revealed that the ABA/WDS domain of the 27.1KDa predictive protein (pl 5.05) was primarily composed of 25 positively charged residues and 46 negatively charged residues. The instability index value of 27.62 also confirmed that the predicted protein is stable. All ASR proteins largely have Glu, Ala, His, Lys and Gly residues and a continuous ABA/WDS domain [58].

Tomato ASR1 protein with DNA binding activity and pl 7.3 is a chromatin bound protein that interacts directly with DNA or indirectly by interacting with other proteins. ASR1 proteins may be nuclear or may be dispersed in the cytoplasm. In general, 40-60KDa molecules diffuse passively through nuclear pores depending on their concentration gradient, whereas other molecules are transported actively [59] through the nuclear localization signals. These NLS (rich in basic amino acids) are recognized by certain docking molecules viz., receptor molecules at the nuclear pore [60]. Sub-cellular localization studies proved ASR1 being smaller in size can pass through the nuclear pore with ease and contains a signal sequence KKDARKKEKKKKLR near the C-terminus [26]. Maize ASRs also have a Zn-binding domain at the N-terminal and a nuclear targeting signal with two abscissic acid/water deficit stress domains (ABA/WDS) at the C-terminal. Hybrid assays and sub-cellular fractionation studies confirmed the role of ZmASR proteins as transcription factors or molecular chaperons in different plant species [61].
Multiple sequence alignment of the chickpea conserved ABA/WDS domain containing protein (NP_001351739.1) with those of model plants available in NCBI was also done using Treebyne software and a phlogenetic tree was generated using neighbour joining method. The *Cicer arietinum* ABA/WDS induced protein grouped with *Brachypodium distachyon* ASR protein with bootstrap value 71. The *Cicer arietinum* ASR protein was found to be closely related to monocotyledonous Gramineae species, *Zea mays* and *Oryza sativa*, *Sorghum bicolor* in comparison to other plant species including cottonwood, banana species. Similar phylogenetic relationships were found while comparing the apple ASR genes with those of other plant species [58].

Three-dimensional structure of the hypothetical protein NP_001351739.1 was predicted using homology modeling in Phyre2 database (http://www.sbg.bio.ic.ac.uk/~phyre2/html/page.cgi?id=index). The protein comprised of alpha helixes accounting for 35% of the total protein (Fig 10A). ASR proteins adapt two different conformations viz., alpha helical or a polyproline type II under different environmental stresses. Polyethylene glycol (PEG) and glycerol stabilize the α-helical conformation in general while lower temperature, lower pH and increased NaCl stabilize the PII conformation. This structural plasticity is critical for plant stress resistance, facilitating their response to drought and interaction with target proteins [62]. The predicted model was then evaluated through Ramachandran plot using Swiss-Pdb Viewer v4.1.0 program by plotting the Psi-Phi angles of the amino acid residues against each other. The Psi-Phi pairs had 72.2% residues in the most preferred regions, 14.5% residues in allowed regions and 13.3% residues in outlier regions as shown in Fig 10B.

Protein Structure Analysis (ProSA) tool identifies the regions that contribute to an overall bad score in the predicted model and has long been used for their refinement and validation. The energy plots and the Z-scores indicate the problems in the predicted structure of the target protein. The z-score of NP_001351739.1 was found to be -3.59 indicating the high quality of the predicted protein model as shown in Fig 11. Energy plot of NP_001351739.1 is shown in Fig 12A. The positive values indicate the erroneous parts of the model in general. Residual error plots indicating reliability of the local model were also plotted and visualized by color gradients. Blue colored regions indicate more reliable regions and red regions indicate the probable unreliable regions. Models that slide towards blue regions from light red color are considered to be of high quality (Fig 12B).

Software packages available online offer great opportunities for analyzing biological systems. AraNet, GeneMania, and STRING being user-friendly [63–65] have been used for studying the protein interactions, regulatory networks, gene associations and their biological pathways [66–68]. STRING database was used to predict the direct and indirect interactions of NP_001351739.1 (http://string-db.org/). The hypothetical protein was found to interact with ten different proteins (Fig 13) for its functioning viz., XP_004497781.1 late embryogenesis abundant protein 1-like (90 amino acid), XP_004508082.1 embryonic protein DC 8-like (424 amino acid), XP_004500781.1 Dehydrin ERD 14 like (225 amino acid), XP_004485799.1 uncharacterized protein ECU09_1610-like (225 amino acid), XP_004509086.1 uncharacterized protein LOC101513321 (89 amino acid), XP_004504366.1 uncharacterized protein LOC101493439; upstream in-frame stop codon (89 amino acid), XP_004509025.1 uncharacterized protein LOC101490213, protein LE25-like (116 amino acid), XP_004509025.1 late embryogenesis abundant protein 2; upstream in-frame stop codon (155 amino acid), CapLEA-1 late embryogenesis abundant protein 1-like (177 amino acid) and XP_004506729.1 protein SLE2; upstream in-frame stop codon (98 amino acid). The predicted functional partners of the hypothetical protein NP_001351739 confirm its role in drought tolerance in chickpea (Fig 14).

Integrated high-throughput approaches employing molecular networks with phenomics together may provide assumptions and address precise biological queries [69].
Drought QTLs have been identified using different approaches viz., QTL mapping [70], sequence similarity based candidate gene allele diversity analysis [36] and genome wide association study (GWAS) [71] by some workers. Ten genes were found to express under abiotic stresses [36]. Functional validation was also done for these genes using the already reported genes from model plants. Linkage analysis and association mapping approach was also used to identify and validate the chickpea genes and QTLs for moisture stress tolerance based on sequence similarity approach [72]. Candidate genes identified may be used to develop cultivars with desired tolerance to drought and ensure greater genetic gains and also enhance the probability of breeding widely adapted high yielding hybrids in chickpea. ASR is the most widely reported drought stress responsive gene [73]. The ASR gene family has evolved from Spermatophyta. ASR gene is regulated by water deficit, salinity stress and hormone Abscisic acid (ABA), low temperature and intensity of light [74]. ASR genes were first recognized in tomato [75] and consequently, in different plant species *Zea mays* (nine), *Sorghum bicolor* (seven), *Oryza sativa* (six), *Brachypodium distachyon* (six), *Pinus taeda* (four), *Phaseolus vulgaris* (two) and *Vitis vinifera* (one) [76,77,18]. Transgenic studies confirmed ASR genes could be involved in ABA signalling pathways enabling the plants to respond to external stresses [78,79] and transgenic plants with over-expressed ASR gene were found to be more tolerance to water and salt stress [26]. During late embryogenesis accumulation of tomato ASR1 was observed. Electrophoretic assays and direct visualization also confirmed formation of homodimers in DNA by ASR1 in response to water stress [23, 80]. In contrast, activity of tomato ASR2 promoter was enhanced in response to ABA in papaya and tobacco, while reduction was observed in tomato and potato [81]. Transgenic *Arabidopsis* lines over-expressing maize ASR genes exhibited better growth performance and higher survival rates as compared to wild type under drought conditions. These lines had lower malondialdehyde content and higher ABA and proline content improving their drought tolerance. The results thus, proved Zm ASR3 enhance drought tolerance via an ABA dependent pathway [82]. Their precise role in conferring improved tolerance to drought and salt has also been established in tomato, rice and lily [33, 34, 26]. Consequently, reports on involvement of ASR gene in legumes in drought responses are insufficient.

**Material and methods**

**Experimental material, soil selection, drought stress treatment**

Seven promising genotypes of chickpea (*Cicer arietinum*) viz., ICCV97309, ICCV3311, ICCV10316, ICCV9307, BGD72, ICCV10, and ICCV5313 were selected from Pulse Research Laboratory, Division of Genetics, Indian Agricultural Research Institute (IARI), Pusa, New Delhi. The amalgamated soil (peat to vermiculite, 1:1) with pH 7.6 and conductivity 0.4 ds/m was taken from the IARI field and each genotype was sown in 13cm diameter plastic pots in three replications under glasshouse conditions at the National Phytotron Facility, Indian Agricultural Research Institute, New Delhi (28°08’N 77°12’) with a photoperiod of 12h in a completely randomized design (CRD) design in the year 2019–20. The temperature was maintained at 24°C in the day and 18°C in the night. These pots were irrigated with 200 ml water on daily basis. Drought stress was imposed on 12 day old plants by withholding water for 6 days and 12days respectively [35]. Control plants were watered regularly for the same duration. Leaf tissues of the control and drought stressed plants were collected at different time points viz., 0d (control), 6d and 12d and fixed in liquid nitrogen and stored at -80°C for RNA isolation. These genotypes were identified on their relative basis of tolerance to drought [83].
**Determination of relative water content**

Relative water content (RWC) of selected chickpea genotypes was measured at different time points (0 day, 6th day and 12th day) as per the standard method [84]. Young leaf tissues of chickpea were collected and their fresh weights were recorded. The leaf tissues were then incubated in petriplates containing distilled water for 4 hours for calculating their turgid weights. Oven drying of the leaves was done for 72 hours at 60˚C and then the plant dry weights were recorded. RWC was calculated using the following formula:

\[
RWC (\%) = \left( \frac{FW - DW}{TW - DW} \right) \times 100
\]

Where, FW - Fresh weight; DW—Dry weight; TW—Turgid weight

**Estimation of chlorophyll index and protein content**

Konica Minolta SPAD 502 Plus chlorophyll meter was used for measuring the chlorophyll index of the selected chickpea genotypes at three different time points (0 day, 6th day and 12th day). For estimating protein content (leaf) at different time points, crushing of leaves was done in 50mM phosphate buffer with pH 7.8. The protein content was estimated by colorimetric method [85]. Absorbance of the samples was recorded on Beckman DU® 640 spectrophotometer at 595 nm [86]. The Bovine Serum Albumin (Sigma, USA) was used as a standard. The protein content was expressed in μg ml⁻¹.

**Determination of membrane stability index**

400 mg fresh leaf sample was taken and added to test tubes containing 10ml of distilled water. The test tubes were kept in a water bath maintained at 45˚C for 30 minutes and conductivity (C1) was noted using a portable conductivity meter. These test tubes were again placed in water bath maintained at 100˚C for 10 minutes and then conductivity was noted again (C2) [87]. The MSI was calculated using formula:

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

**Statistical analysis**

The data for all physiological parameters were subjected to standard method of statistical analysis such as analysis of variance (ANOVA) using XLSTAT software. The mean values and coefficient of variation (CV) were calculated for each parameter. The standard errors of the mean were presented in the figures as error bars. The mean comparisons were performed using Tukey’s Studentized Range (HSD) test. The Tukey’s Studentized Range (HSD) test at p = 0.05 was employed to test the differences among the treatment means for the measured parameters at 0 (control), 6th day and 12th day after imposing drought stress.

**RNA extraction and quantitative real-time PCR (qRT-PCR) analysis**

RNA isolation was done by using NucleoZOL (Takara Bio). Genomic DNA and other contaminants were removed by precipitation. One phase RNA extraction was followed by conversion to first strand cDNA using the Accuscript high fidelity cDNA synthesis kit (Agilent). Brilliant III ultra fast SYBR Green was used to measure the relative changes in the expression of chickpea ASR genes under water stressed conditions. The cDNA from leaf tissues were used as template. The Beta Actin gene was used as the reference gene. The qRT-PCR was conducted on a CFX 96 Real Time PCR (Biorad) in a reaction volume of 25μL, that comprised of 2μL chickpea
samples cDNA, 0.5 μL each ASR specific forward and reverse primer, 12.5 μL Brilliant III ultra fast SYBR Green QPCR master mix (Agilent), and 9.5 μL nuclease-free molecular biology grade water. The qRT-PCR reaction cycle included 95˚C for 3 min followed by 40 cycles at 95˚C for 5 s, 60˚C for 12 s. The relative expression levels of chickpea ASR genes under different treatments were then calculated using the $2^{-\Delta\Delta CT}$ method [88].

DNA extraction and PCR amplification

The genomic DNA of seven chickpea genotypes was extracted from young leaves using the CTAB method [39]. Purified DNA was used for PCR amplification using ASR gene specific forward 5’-GGGAACATATCTTTTCAACCA-3’ and reverse 5’-CTGCAGCACCTAACCACCA-3’ primer custom synthesized by G-Biosciences, USA [83]. PCR was carried out in the Chickpea Molecular Breeding Laboratory, Division of Genetics, ICAR-IARI using a G-STORM thermal cycler (Labtech, France). The PCR master mix comprised of 20ng of the template DNA, 1.6 μl of 10X Tris borate-ethylenediamine-tetra acetic acid (TBE), 1μl of 10mM dNTP mix (Genei, Bangalore), 1μl each of 5μM forward and reverse primer and 0.3μl of 3U Taq Polymerase (Genei, Bangalore). The PCR cycling reaction consisted of three steps, starting with initial denaturation (90˚C for 3min) followed by 38 cycles of denaturation (94˚C for 20sec), annealing (55˚C for 50sec) and elongation (72˚C for 50sec) followed by final elongation (72˚C for 7mins) [89]. The PCR amplicons were visualized on a 3% 1X TBE buffer using 100bp DNA ladder (Thermo Scientific, USA). The gel image was documented using UV light gel documentation system (UVITECH Imaging System, UK).

Nucleotide sequence analysis

PCR amplicons of the seven chickpea genotypes were purified using BigDye terminator v3.1 kit (Applied Biosystems, USA) and were sequenced using an ABI3500XL genetic analyser (Applied Biosystems, USA). Sequencing data was analyzed in Sequencing Analysis v 5.4. Raw sequences were assembled using the forward and reverse sequences of each genotype in KB v 1.4.1.8(KB base caller) tool. The assembled chickpea ASR gene sequence was submitted to NCBI GenBank (MK937569) using the web-based submission tool Sequin from the NCBI home page (https://www.ncbi.nlm.nih.gov/Sequin/). A comparison BLAST tools (blastn) were used determine the sequence identities of the assembled sequence (http://www.ncbi.nlm.nih.gov). Sequence alignments were performed using MUSCLE software and phylogram of Cicer ASR gene with ASR gene sequences of different legumes available at NCBI database was constructed by neighbor joining method.

Computational protein analysis

The ASR sequences were also verified at the protein level using blastx and Expasy server (http://expasy.org/cgi-bin/protparam) and their conserved domains were identified using the NCBI tool for conserved domain search (CDD). Common characteristics of the predicted protein including molecular weight, isoelectric point (pI), amino acid composition, aliphatic and instability index were assessed using protparam tool. The amino acid sequences for the ABA/WDS conserved domain from different crop plants were retrieved from NCBI CDD tool. Sequence alignments were performed and a circular phylogenetic was constructed using Tree-dyne software (https://www.phylogeny.fr/). The three dimensional structure was predicted by homology modeling using the Phyre2 database and validated with Swiss-PDB Viewer (SPDBV). The target protein model was further refined by the Program structure Analysis (ProSA) program that predicts the structure of the target protein using the molecule viewer Jmol in order to find out the regions that contribute to errors in the protein model. Network
analysis was done using the STRING database (http://string-db.org/) to find out functional and physical interactions of the predicted protein.

Conclusion

Present study reveals that increased expression of ASR gene under drought stress possibly enabled the tolerant chickpea genotypes to perform better under stressed conditions. The results show a close relationship between qRT-PCR data and physiological characterization of the genotypes under drought stress conditions which displayed higher RWC(%), MSI, CI and protein content in BGD72, ICCV10 and ICCV3311 in comparison to the susceptible genotypes ICCV2, Pusa1003 and Pusa362. Modifications have occurred at various nucleotides in gene sequence of ASR genes during evolution. The NCBI CDD tool, Expasy Protparam, Phyre2, Swiss PDB viewer and ProSA tool revealed important features viz., primary structure, secondary structure, z-scores of hypothetical protein NP_001351739. NP_001351739 with Arginine, lysine, glutamic acid, Asparagine and ABA/WDS conserved domain that might comprise the nuclear localization signals (NLS) and pass through the nuclear pores inducing different plant stress inducible genes. The predicted functional partners identified by STRING network analysis also proved that NP_001351739 interacts with various LEA proteins, dehydrins for its functioning and is likely to play an important role in drought tolerance in chickpea. This hypothetical ASR protein might have enhanced the ASR gene activity as a transcription factor mediating drought responses in chickpea. This study could be useful in identification of new ASR genes that play a major role in drought tolerance and also develop functional markers for chickpea improvement.

Supporting information

S1 File.
(PDF)

S1 Raw image. PCR Amplification of seven chickpea genotypes using ASR gene specific marker. PCR amplification of seven chickpea genotypes viz., ICCV97309, ICCV3311, ICCV10316, ICCV9307, BGD72, ICCV10, and ICCV5313 was done using ASR gene specific primer and revealed a single amplicon ranging from 680-700bp; Marker-100 bp Banglore Genei DNA ladder.
(PDF)

Acknowledgments

The authors would like to thank the Division of Genetics for providing all the facilities to carry out genotyping and software analysis.

Author Contributions

Conceptualization: C. Bharadwaj, P. K. Jain.
Data curation: Rajesh Kumar Singh.
Formal analysis: Supriya Sachdeva.
Funding acquisition: C. Bharadwaj.
Investigation: Supriya Sachdeva.
Methodology: Supriya Sachdeva, B. S. Patil.
Project administration: C. Bharadwaj.

Resources: C. Bharadwaj, B. S. Patil.

Software: Supriya Sachdeva, Rajesh Kumar Singh, P. K. Jain, Manish Roorkiwal, Rajeev Varshney.

Supervision: C. Bharadwaj.

Validation: Rajesh Kumar Singh.

Visualization: C. Bharadwaj, Rajeev Varshney.

Writing – original draft: Supriya Sachdeva, C. Bharadwaj.

Writing – review & editing: C. Bharadwaj, P. K. Jain, Manish Roorkiwal, Rajeev Varshney.

References

1. Gaur PM, Jukanti AK, Varshney RK. Impact of genomic technologies on chickpea breeding strategies. Agronomy. 2012; 2(3):199–221.

2. Sachdeva S, Bharadwaj C, Sharma V, Patil BS, Soren KR, Roorkiwal M, et al. Molecular and phenotypic diversity among chickpea (Cicer arietinum) genotypes as a function of drought tolerance. Crop and Pasture Science. 2018; 69(2):142–53.

3. Kumar N, Bharadwaj C, Satyavathi CT, Pal M, Kumar T, Singhal T, et al. Morpho Physiological Characterization and Grouping (SAHN) of Chickpea Genotypes for Salinity Tolerance. Vegetos. 2017; 30:3.

4. Varshney RK, Song C, Saxena RK, Azam S, Yu S, Sharpe AG, et al. Draft genome sequence of chickpea (Cicer arietinum) provides a resource for trait improvement. Nature biotechnology. 2013; 31(3):240. https://doi.org/10.1038/nbt.2491 PMID: 23354103

5. Krishnamurthy L, Kashiwagi J, Upadhyaya HD, Gowda CL, Gaur PM, Singh S, et al. Partitioning coefficient-A trait that contributes to drought tolerance in chickpea. Field Crops Research. 2013; 149:354–65.

6. Varshney RK, Thudi M, Nayak SN, Gaur PM, Kashiwagi J, Krishnamurthy L, et al. Genetic dissection of drought tolerance in chickpea (Cicer arietinum L.). Theoretical and Applied Genetics. 2014a; 127(2):445–62. https://doi.org/10.1007/s00122-013-2230-6 PMID: 24326458

7. Supriya S, Bharadwaj C, Vinay S, Neeraj K, Kv B, Patil BS, et al. Morpho-physiological grouping of chickpea (Cicer arietinum L.) genotypes on the basis of their response to drought stress. International Journal of Tropical Agriculture. 2017; 35(1):5–13.

8. Da Silva AF, Sediyama T, Matsu E, Silva FD, Cruz CD, Borém A, et al. Phenotypic and molecular diversity among soybean cultivars as a function of growing season. Agronomy Science and Biotechnology. 2015; 1:52–61.

9. Jaganathan D, Thudi M, Kale S, Azam S, Roorkiwal M, Gaur PM, et al. Genotyping-by-sequencing based intra-specific genetic map refines a “QTL-hotspot” region for drought tolerance in chickpea. Molecular Genetics and Genomics. 2015; 290(2):559–71. https://doi.org/10.1007/s00438-014-0932-3 PMID: 25344290

10. Sudupak M, Akkaya MS, Kence A. Analysis of genetic relationships among perennial and annual Cicer species growing in Turkey using RAPD markers. Theoretical and Applied Genetics. 2002; 105(8):1220–8. https://doi.org/10.1007/s00122-002-1060-8 PMID: 12582902

11. Shimray PW, Bajaj D, Srivastava R, Daware A, Upadhyaya HD, et al. Identifying transcription factor genes associated with yield traits in chickpea. Plant molecular biology reporter. 2017; 35(5):562–74.

12. Kujur AL, Bajaj DE, Saxena MS, Tripathi SH, Upadhyaya HD, Gowda CL, et al. Functionally relevant microsatellite markers from chickpea transcription factor genes for efficient genotyping applications and trait association mapping. DNA research. 2013; 20(4):355–74. https://doi.org/10.1093/dnares/dst015 PMID: 23633531

13. Century K, Reuber TL, Ratcliffe OJ. Regulating the regulators: the future prospects for transcription-factor-based agricultural biotechnology products. Plant physiology. 2008; 147(1):20–9. https://doi.org/10.1104/pp.108.117887 PMID: 18443109

14. Cortés AJ, Chavarro MC, Madriñán S, This D, Blair MW. Molecular ecology and selection in the drought-related Asr gene polymorphisms in wild and cultivated common bean (Phaseolus vulgaris L.). BMC genetics. 2012; 13(1):58.
15. Bartels D, Sunkar R. Drought and salt tolerance in plants. Critical reviews in plant sciences. 2005; 24(1):23–58.

16. Saumoneau A, Laloï M, Lallemand M, Rabot A, Atanassova R. Dissection of the transcriptional regulation of grape ASR and response to glucose and abscisic acid. Journal of experimental botany. 2011; 63(3):1495–510. https://doi.org/10.1093/jxb/err391 PMID: 22140241

17. Taiz L, Zeiger E. Stress physiology. Plant physiology. 2006;4.

18. Frankel N, Carrari F, Hasson E, Iusem ND. Evolutionary history of the Asr gene family. Gene. 2006; 378:74–83. https://doi.org/10.1016/j.gene.2006.05.010 PMID: 16822623

19. Taiz L, Zeiger E. Stress physiology. Plant physiology. 2006;4.

20. Frankel N, Carrari F, Hasson E, Iusem ND. Evolutionary history of the Asr gene family. Gene. 2006; 378:74–83. https://doi.org/10.1016/j.gene.2006.05.010 PMID: 16822623

21. Carrari F, Fernie AR, Iusem ND. Heard it through the grapevine? ABA and sugar cross-talk: the ASR story. Trends in plant science. 2004; 9(2):57–9. https://doi.org/10.1016/j.tplants.2003.12.004 PMID: 15106586

22. Jeanneau M, Gerentes D, Fouelliasar X, Zivy M, Vidal J, Toppan A, et al. Improvement of drought tolerance in maize: towards the functional validation of the Zm-Asr1 gene and increase of water use efficiency by over-expressing C4–PEPC. Biochimie. 2002; 84(11):1127–35. https://doi.org/10.1016/s0300-9084(02)00024-x PMID: 12595141

23. Masklin L, Gudesblat GE, Moreno JE, Carrari FO, Frankel N, Sambade A, et al. Differential expression of the members of the Asr gene family in tomato (Lycopersicon esculentum). Molecular and General Genetics MGG. 1998; 258(1–2):1–8. https://doi.org/10.1007/s004380050700 PMID: 9613566

24. De Vienne D, Leonardi A, Damerval C, Zivy M. Genetics of proteome variation for QTL characterization: application to drought-stress responses in maize. Journal of Experimental Botany. 1999; 50(332):303–9.

25. Schneider A, Salamini F, Gebhardt C. Expression patterns and promoter activity of the cold-regulated gene ci21A of potato. Plant physiology. 1997; 113(2):335–45. https://doi.org/10.1104/pp.113.2.335 PMID: 9046587

26. Kalifa Y, Perlson E, Gilad A, Konrad Z, Scolnik PA, Bar-Zvi D. Over-expression of the water and salt stress-regulated Asr1 gene confers an increased salt tolerance. Plant, Cell & Environment. 2004; 27(12):1459–68.

27. Silhavy D, Hutfvångner G, Barta E, Bánfalvi Z. Isolation and characterization of a water-stress-inducible cDNA clone from Solanum chacoense. Plant molecular biology. 1995; 27(3):587–95. https://doi.org/10.1007/BF00019324 PMID: 7894021

28. Iusem ND, Bartholomew DM, Hiltz WD, Scolnik PA. Tomato (Lycopersicon esculentum) transcript induced by water deficit and ripening. Plant Physiology. 1993; 102(4):1353. https://doi.org/10.1104/pp.102.4.1353 PMID: 8278555

29. Wang CS, Liau YE, Huang JC, Wu TD, Su CC, et al. (1998) Characterization of a desiccation-related protein in lily pollen during development and stress. Plant Cell Physiol 39: 1307–1314. https://doi.org/10.1093/oxfordjournals.pcp.a029335 PMID: 10050314

30. Riccardi F, Gazeau P, de Vienne D, Zivy M. Protein changes in response to progressive water deficit in maize: quantitative variation and polypeptide identification. Plant Physiology. 1998; 117(4):1253–63. https://doi.org/10.1104/pp.117.4.1253 PMID: 9701581

31. Chang S, Puryear JD, Dias MD, Funkhouser EA, Newton RJ, Cairney J. Gene expression under water deficit in loblolly pine (Pinus taeda): isolation and characterization of cDNA clones. Physiologia Plantarum. 1996; 97(1):139–48.

32. Cortés AJ, Monseerrate FA, Ramírez-Villegas J, Madriñán S, Blair MW. Drought tolerance in wild plant populations: the case of common beans (Phaseolus vulgaris L.). PLoS One. 2013; 8(5):e62898. https://doi.org/10.1371/journal.pone.0062898 PMID: 23658783

33. Yang CY, Chen YC, Jauh GY, Yang WS. A lily ASR protein involves abscisic acid signaling and confers drought and salt resistance in Arabidopsis. Plant physiology. 2005; 139(2):836–46. https://doi.org/10.1104/pp.105.065458 PMID: 16169963

34. Philippe R, Courtot B, McNally KL, Mourset P, El-Maliki R, Le Paslier MC, et al. Structure, allelic diversity and selection of Asr genes, candidate for drought tolerance, in Oryza sativa L. and wild relatives. Theoretical and applied genetics. 2010; 121(4):769–87. https://doi.org/10.1007/s00122-010-1348-z PMID: 20454772
35. Jain D, Chattopadhayay D. Analysis of gene expression in response to water deficit of chickpea (Cicer arietinum L.). BMC plant biology. 2010; 10(1):24.

36. Roorkiwal M, Nayak SN, Thudi M, Upadhyaya HD, Brunel D, Mounet P, et al. Allele diversity for abiotic stress responsive candidate genes in chickpea reference set using gene based SNP markers. Frontiers in plant science. 2014; 5:248. https://doi.org/10.3389/fpls.2014.00248 PMID: 24926299

37. Bharadwaj C, Srivastava R, Chauhan SK, Satyavathi CT, Kumar J, Faruqui A, et al. Molecular diversity and phylogeny in geographical collection of chickpea (Cicer sp.) accessions. Journal of genetics. 2011:1–7. PMID: 21677383

38. Millan T, Clarke HJ, Siddique KH, Buhairalla HK, Gaur PM, Kumar J, et al. Chickpea molecular breeding: new tools and concepts. Euphytica. 2006; 147(1–2):81–103.

39. Tapan K, Bharadwaj C, Satyavathi CT, Jain PK. A high throughput, improved rapid and reliable genomic DNA extraction protocol from chickpea (Cicer arietinum L.). Vegetos. 2013; 26(2):185–90.

40. Choudhary P, Khanna SM, Jain PK, Bharadwaj C, Kumar J, Lakhera PC, et al. Molecular characterization of primary gene pool of chickpea based on ISSR markers. Biochemical genetics. 2013; 51(3–4):306–22. https://doi.org/10.1007/s10528-012-9564-7 PMID: 23329257

41. Varshney RK, Mir RR, Bhatia S, Thudi M, Hu Y, Azam S, et al. Integrated physical, genetic and genome map of chickpea (Cicer arietinum L.). Functional & integrative genomics. 2014b; 14(1):59–73.

42. Rahbarian R, Khvare-Nejad R, Ganjelai A, Bagheri A, Najafi F. Drought stress effects on photosynthesis, chlorophyll fluorescence and water relations in tolerant and susceptible chickpea (Cicer arietinum L.) genotypes. Acta Biologica Cracoviensia Series Botanica. 2011; 53(1):47–56.

43. Schonfeld MA, Johnson RC, Carver BF, Mornhinweg DW. Water relations in winter wheat as drought resistance indicators. Crop Science. 1988; 28(3):526–31.

44. Pouresmael M, Khavari-Nejad RA, Mozafari J, Najafi F, Moradi F. Efficiency of screening criteria for drought tolerance in chickpea. Archives of Agronomy and Soil Science. 2013; 59(12):1675–93.

45. Leport L, Turner NC, French RJ, Barr MD, Duda R, Davies SL, et al. Physiological responses of chickpea genotypes to terminal drought in a Mediterranean-type environment. European Journal of Agronomy. 1999; 11(3–4):279–91.

46. Bhattacharjee S. An inductive pulse of hydrogen peroxide pretreatment restores redox-homeostasis and oxidative membrane damage under extremes of temperature in two rice cultivars. Plant growth regulation. 2012; 66(3):395–410.

47. Iturbe-Ormaetxe I, Escuredo PR, Arrese-Igor C, Becana M. Oxidative damage in pea plants exposed to water deficit or paraquat. Plant physiology. 1998; 116(1):173–81.

48. Ouvrard O, Cellier F, Ferrare K, Tousch D, Lamaze T, Dupuis JM, et al. Identification and expression of water stress-and abscisic acid-regulated genes in a drought-tolerant sunflower genotype. Plant molecular biology. 1996; 31(4):819–29. https://doi.org/10.1007/BF00019469 PMID: 8864412

49. Jiang Y, Huang B. Protein alterations in tall fescue in response to drought stress and abscisic acid. Crop Science. 2002; 42(1):202–7. https://doi.org/10.2135/cropsci2002.2020 PMID: 11756275

50. Kumar J, Basu PS, Srivastava E, Chaturvedi SK, Naradaj N, Kumar S. Phenotyping of traits imparting drought tolerance in lentil. Crop and Pasture Science. 2012; 63(6):547–54.

51. Mafakheri A. Effect of drought stress and subsequent recovery on protein, carbohydrate contents, catalase and peroxidase activities in three chickpea (Cicer arietinum) cultivars. Australian Journal of Crop Science. 2011; 5(10):1255–60.

52. Ahire RK, Kale AA, Munjal SV, Jamdagni BM. Induced water stress influencing proline accumulation, protein profiles and DNA polymorphism in chickpea cultivars. Indian Journal of Plant Physiology. 2005; 10(3):218.

53. Batra NG, Sharma V, Kumari N. Drought-induced changes in chlorophyll fluorescence, photosynthetic pigments, and thylakoid membrane proteins of Vigna radiata. Journal of Plant Interactions. 2014; 9 (1):712–21.

54. Zheng F, Wu H, Zhang R, Li S, He W, Wong FL, et al. Molecular phylogeny and dynamic evolution of disease resistance genes in the legume family. BMC genomics. 2016; 17(1):402.

55. Joo J, Lee YH, Kim YK, Nahm BH, Song SI. Abiotic stress responsive rice ASR1 and ASR3 exhibit different tissue-dependent sugar and hormone-sensitivities. Molecules and cells. 2013; 35(5):421–35. https://doi.org/10.1007/s10059-013-0036-7 PMID: 23620302

56. Yoon JS, Kim JY, Lee MB, Seo YW. Over-expression of the Brachypodium ASR gene, BdASR4, enhances drought tolerance in Brachypodium distachyon. Plant cell reports. 2019; 38(9):1109–25. https://doi.org/10.1007/s00299-019-02429-7 PMID: 31134348

57. Virlouvet L, Jacquemont MP, Gerentes D, Corti H, Bouton S, Gilard F, et al. The ZmASR1 protein influences branched-chain amino acid biosynthesis and maintains kernel yield in maize under water-limited...
58. Huang K, Zhong Y, Li Y, Zheng D, Cheng ZM. Genome-wide identification and expression analysis of the apple ASR gene family in response to *Alternaria alternata* f. sp. mali. Genome. 2016; 59(10):866–78. https://doi.org/10.1104/gen-2016-0043 PMID: 27653246

59. Yoneda Y. Nucleocytoplasmic protein traffic and its significance to cell function. Genes to Cells. 2000; 5(10):777–87. https://doi.org/10.1046/j.1365-2443.2000.00366.x PMID: 11029654

60. Jans DA, Xiao CY, Lam MH. Nuclear targeting signal recognition: a key control point in nuclear transport?. Bioessays. 2000; 22(6):532–44. https://doi.org/10.1002/(SICI)1521-1878(200006)22:6<532::AID-BIES6>3.0.CO;2-O PMID: 10842307

61. Liang Y, Jiang Y, Du M, Li B, Chen L, Chen M, et al. ZmASR3 from the Maize ASR Gene Family Positively Regulates Drought Tolerance in Transgenic *Arabidopsis*. International journal of molecular sciences. 2019; 20(9):2278.

62. Chen Z, Hong X, Zhang H, Wang Y, Li Z, Zhu JK, et al. Disruption of the cellulose synthase gene, AtCesA8/IRX1, enhances drought and osmotic stress tolerance in *Arabidopsis*. The Plant Journal. 2005; 43(2):273–83. https://doi.org/10.1111/j.1365-313X.2005.02452.x PMID: 15998313

63. Warde-Farley D, Donaldson SL, Comes O, Zuberi K, Badrawi R, Chao P, et al. The GeneMANIA prediction server: biological network integration for gene prioritization and predicting gene function. Nucleic acids research. 2010; 38(suppl_2):W214–20.

64. Hwang S, Rhee SY, Marcotte EM, Lee I. Systematic prediction of gene function in *Arabidopsis thaliana* using a probabilistic functional gene network. Nature protocols. 2011; 6(9):1429. https://doi.org/10.1038/nprot.2011.372 PMID: 21886106

65. Szklarczyk D, Morris JH, Cook H, Kuhn M, Wyder S, Simonovic M, et al. The STRING database in 2017: quality-controlled protein–protein association networks, made broadly accessible. Nucleic acids research. 2016; 41(D1):D808–15.

66. Song Q, Wang S, Zhang G, Li Y, Li Z, Guo J, et al. Comparative proteomic analysis of a membrane-enriched fraction from flag leaves reveals responses to chemical hybridization agent SQ-1 in wheat. Frontiers in plant science. 2015; 6:669. https://doi.org/10.3389/fpls.2015.00669 PMID: 26379693

67. Franceschini A, Szklarczyk D, Frankild S, Kuhn M, Simonovic M, Roth A, et al. STRING v9. 1: protein-protein interaction networks, with increased coverage and integration. Nucleic acids research. 2012; 41(D1):D808–15.

68. Maere S, Heymans K, Kuiper M. BiNGO: a Cytoscape plugin to assess overrepresentation of gene ontology categories in biological networks. Bioinformatics. 2005; 21(16):3448–9. https://doi.org/10.1093/bioinformatics/bti551 PMID: 15972284

69. Bassel GW, Gaudinier A, Brady SM, Hennig L, Rhee SY, De Smet I. Systems analysis of plant functional, transcriptional, physical interaction, and metabolic networks. The Plant Cell. 2012; 24(10):3859–75. https://doi.org/10.1105/tpc.112.100776 PMID: 23110892

70. Rehman AU, Malhotra RS, Bett K, Tar’an B, Bueckert R, Warkentin TD. Mapping QTL associated with traits affecting grain yield in chickpea (*Cicer arietinum L.*) under terminal drought stress. Crop Science. 2011; 51(2):450–63.

71. Thudi M, Upadhyaya HD, Rathore A, Gaur PM, Krishnamurthy L, Roorkiwal M, et al. Genetic dissection of drought and heat tolerance in chickpea through genome-wide and candidate gene-based association mapping approaches. Plos one. 2014; 9(5):e96758. https://doi.org/10.1371/journal.pone.0096758 PMID: 24801366

72. Nayak SN. Identification of QTLs and genes for drought tolerance using linkage mapping and association mapping approaches in chickpea (*Cicer arietinum*) (Doctoral dissertation, Osmania University, Hyderabad, India).

73. Hong SH, Kim U, Yang DC, Chung WI. Characterization of an abscisic acid responsive gene homologue from *Cucumis melo*. Journal of experimental botany. 2002; 53(378):2271–2. https://doi.org/10.1093/jxb/erf075 PMID: 12379794

74. González RM, Iusem ND. Twenty years of research on Asr (ABA-stress-ripening) genes and proteins. Planta. 2014; 239(5):941–9. https://doi.org/10.1007/s00425-014-2039-9 PMID: 24531839

75. Iusem ND, Bartholomew DM, Hilt WD, Scolinik PA. Tomato (Lycopersicon esculentum) transcript induced by water deficit and ripening. Plant Physiology. 1993; 102(4):1353. https://doi.org/10.1104/pp.102.4.1353 PMID: 8278555

76. Pérez-Díaz J, Wu TM, Pérez-Díaz R, Ruiz-Lara S, Hong CY, Casaretto JA. Organ-and stress-specific expression of the ASR genes in rice. Plant cell reports. 2014; 33(1):61–73. https://doi.org/10.1007/s00299-013-1512-4 PMID: 24085307
77. Cortés AJ, Chavarro MC, Madriñán S, This D, Blair MW. Molecular ecology and selection in the drought-related Asr gene polymorphisms in wild and cultivated common bean (Phaseolus vulgaris L.). BMC genetics. 2012; 13(1):58.

78. Hu WE, Huang C, Deng X, Zhou S, Chen L, Li Y, et al. TaASR1, a transcription factor gene in wheat, confers drought stress tolerance in transgenic tobacco. Plant, cell & environment. 2013; 36(8):1449–64.

79. Miao H, Wang Y, Liu J, Jia C, Hu W, Sun P, et al. Molecular cloning and expression analysis of the MaASR1 gene in banana and functional characterization under salt stress. Electronic Journal of Biotechnology. 2014; 17(6):287–95.

80. Goldgur Y, Rom S, Ghirlando R, Shkolnik D, Shadrin N, Konrad Z, et al. Desiccation and zinc binding induce transition of tomato abscisic acid stress ripening 1, a water stress- and salt stress-regulated plant-specific protein, from unfolded to folded state. Plant physiology. 2007; 143(2):617–28. https://doi.org/10.1104/pp.106.092965 PMID: 17189335

81. Golan I, Dominguez PG, Konrad Z, Shkolnik-Inbar D, Carrari F, Bar-Zvi D. Tomato abscisic acid stress ripening (ASR) gene family revisited. PLoS One. 2014 13; 9(10):e107117. https://doi.org/10.1371/journal.pone.0107117 PMID: 25310287

82. Liang Y, Jiang Y, Du M, Li B, Chen L, Chen M, et al. ZmASR3 from the Maize ASR Gene Family Positively Regulates Drought Tolerance in Transgenic Arabidopsis. International journal of molecular sciences. 2019; 20(9):2278.

83. Sachdeva SU, Bharadwaj C, Singh S, Roorkiwal M, Sharma VI, Singh AN, et al. Yield plasticity and molecular diversity analysis in chickpea (Cicer arietinum). Indian Journal of Agricultural Sciences (TSI). 2019; 89(5):834–41.

84. Barrs HD, Weatherley PE. A re-examination of the relative turgidity technique for estimating water deficits in leaves. Australian journal of biological sciences. 1962; 15(3):413–28.

85. Prasad AR, Luduena RF, Horowitz PM. Detection of energy transfer between tryptophan residues in the tubulin molecule and bound bis (8-anilinonaphthalene-1-sulfonate), an inhibitor of microtubule assembly that binds to a flexible region on tubulin. Biochemistry. 1986; 25(12):3536–40. https://doi.org/10.1021/bi00360a010 PMID: 3718942

86. Bradford MM. A rapid and sensitive method for the quantitation of microgram quantities of protein utilizing the principle of protein-dye binding. Analytical biochemistry. 1976; 72(1-2):248–54.

87. Blum A, Ebercon A. Cell membrane stability as a measure of drought and heat tolerance in wheat 1. Crop Science. 1981; 21(1):43–7.

88. Livak KJ, Schmittgen TD. Analysis of relative gene expression data using real-time quantitative PCR and the 2− △△CT method. 2001; 25(4):402–8.

89. Shubha Y, Bharadwaj C, Chauhan SK, Rizvi AH, Kumar J, Satyavathi CT. Analysis of genetic diversity in Cicer species using molecular markers. Indian Journal of Genetics and Plant Breeding. 2011; 71 (3):272–5.