Chloroplastic acyl carrier protein synthase I and chloroplastic 20 kDa chaperonin proteins are involved in wheat (*Triticum aestivum*) in response to moisture stress

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*ABSTRACT*

In this study, two bread wheat (*Triticum aestivum* L.) cultivars, Pishgam (drought-tolerant) and Shahryar (drought-sensitive), were grown in the greenhouse under control and moisture stress conditions. Based on phenological and morpho-physiological results, Pishgam was confirmed as a moisture stress tolerant cultivar. In the following, at the start of heading time, its treated and untreated flag leaves were sampled for two-dimensional electrophoresis (2-DE) based on proteomics approach. Among approximately 263 protein spots appearing in two-dimensional gels, 23 and 10 protein spots were up- and down-regulated, respectively. Among these differentially expressed proteins, 11 proteins with more differences were identified by MALDI TOF/TOF MS which allocated to six functional protein groups involved in photosynthesis or respiration, carbohydrate metabolism, energy metabolism, chaperon, lipid metabolism and unknown function. We report this for the first time that chloroplastic acyl carrier protein synthase I and chloroplastic 20 kDa chaperonin proteins were significantly changed in wheat in response to moisture stress.

**Introduction**

Abiotic stresses can change physiological and biochemical traits (Jamshidi Goharrizi, Baghizadeh, et al. 2020; Jamshidi Goharrizi, Moosavi, et al. 2020), genes expression template (Jamshidi Goharrizi et al. 2018) as well as proteome pattern of plants (Nazari et al. 2018). One crucial issue in plant production is access to water, which affects the plant growth cycle. Low moisture leads to water stress, which is most often observed in areas with low rainfall and low irrigation (Wang et al. 2005). Wheat (*Triticum aestivum* L.), as one of the most important crops in the world, accounts for about 20% of the calories consumed by humans (Brenchley et al. 2012); meanwhile, due to the phenomenon of global warming and limitation of available water resources, its performance is decreasing annually. In recent years, numerous studies have focused on interactions between bread wheat and drought stress, which their results have demonstrated that recognizing genetic diversity in various traits can be effective in tolerance to stress (Hameed et al. 2011). Moisture stress leads to adverse effects on the quantity and quality of wheat yield, in addition, it creates complex responses at the cellular and physiological levels in the plant (Nazari et al. 2019). Obtaining and identifying high yielding plants with identifying and improving molecular mechanisms under water stress conditions could be the best strategy to deal with drought stress (Kamal et al. 2010). Proteins are essential biomolecules in organisms that affect all of cell functions, and their expression measurements can provide a broad overview of molecular events and specific physiological conditions (Ngara and Ndimba 2014). Proteomics is known as an effective technique for identifying potential proteins which are presented in tissues, cells or subcellular compartments, in different conditions (Ghatak et al. 2017). Proteomics is a powerful technique to directly assess of proteins influenced by a particular environmental stimuli, to identification of biochemical pathways and the complex response of plants to environmental stress. So far, several proteomics studies have been performed on crop species of wheat under drought stress (Caruso et al. 2009; Bazargani et al. 2011; Ge et al. 2012; Budak et al. 2013; Kamal et al. 2013; Cheng et al. 2015, 2016; Fotovat et al. 2017; Li et al. 2018).

In the present study, we first assessed the effect of imposing moisture stress in two bread wheat cultivars; Pishgam (drought-tolerant) and Shahryar (drought-sensitive). After determining the tolerant cultivar to moisture stress, we used a proteomics approach to identify responsive protein classes involved in tolerance to moisture stress in tolerant bread wheat (Pishgam cultivar). This study provides more information about the flag leaf protein profiles of tolerant bread wheat in response to moisture stress and identification of molecular differences and tolerance mechanisms.

**Materials and methods**

**Plant materials and moisture stress treatment**

The materials used in the greenhouse pot experiment included two bread wheat cultivars (*Triticum aestivum* L.): drought-sensitive Shahryar and drought-tolerant Pishgam developed in Seed and Plant Improvement Institute (S.P.I.I), Karaj, Iran. Pishgam cultivar was bred and introduced as a tolerant cultivar for arid and semi-arid regions of Iran by S.P.I.I. (Table 1).

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Before the start of the proteomics experiment, tolerance to moisture stress in Pishgam was re-approved using our pheno-logical and morpho-physiological data analysis. So that the preliminary experiments was conducted in a research-green-house at Bu-Ali Sina University (located in Hamedan province, west of Iran) in 2014–2015 growing season. The seeds were surface sterilized by washing with 70% ethanol, followed by immersion in 5% sodium hypochlorite for 30 min and were finally washed three times with distilled water. Then, wheat seeds were germinated and vernalized in the dark condition for 21 days at 4°C before transplanted into pots. Next, five germinated seedlings of similar growth were transferred to black plastic pots, with 40 cm of diameter and 80 cm of height, containing 15 kg soil comprised of 50% agronomy-field soil (silty-loam), 25% sand and 25% manure. Plants were grown under two controlled and imposed moisture stress conditions with 10–12 h photoperiod and 25 ± 3°C temperature in three replications. In the following, soil moisture was maintained at 95% and 45% soil pod capacity (S.P.C) in controlled and imposed moisture stress conditions, respectively. At the first-three weeks of normal growth, the irradiation of plants was applied daily with tap water while adding the necessary volume to bring soil to field capacity (determined by weighing pots). After the first-three weeks, the applying moisture stress treatment (45% S.P.C) was started when the seedling had approximately 4–6 leaves and followed until the harvest time. At heading time, the expanded flag leaves of treated and untreated plants were harvested, quickly wrapped in aluminum foil pouch, immediately frozen in liquid nitrogen and stored at −80 °C for protein extraction.

**Measurement of the morpho-physiological traits**

In both non-stress and stress conditions, 31 traits related to pheno-logy, morpho-physiology, root-characters, and grain yield were measured at heading and harvest times on five plants in each pot. These traits were included days to heading (DTH), days to anthesis (DTA), days to maturity (DTM), grain filling period (GFP), chlorophyll content (SPAD), plant height (PH), peduncle length (PEL), tiller number per plant (TN), leaves number per plant (LN), fertile spike number per plant (FSNPP), spikelet number per spike (SNPS), seed number per main spike (SNPMS), seed number per plant (SNPP), main spike weight (MSW), seed weight per main spike (SWPMS), peduncle weight (PEW), main stem weight (MSTW), 1000-grain weight (TGW), economical yield per plant (EYPP), biological yield per plant (BYPY), plant harvest index (PHI), leaf area index (LAI), relative water content (RWC), water use (WU), excised leaf water retention (ELWR), water use efficiency (WUE), main root length (MRL), root volume (RV), root dry weight (RDW), root area (RA), root to shoot dry weight ratio (RDW/SDW). At heading time, leaf physiological traits including relative water content (RWC), excised leaf water retention (ELWR) and chlorophyll content (SPAD index) were measured on the second leaves per plant. So that, RWC and ELWR were measured according to Mguis et al. (2013), and SPAD index was recorded using a chlorophyll meter (SPAD-502; Konica Minolta Sensing, Inc. Osaka, Japan). The total water use (WU) was calculated as the amount of water use during the plant growth.

At physiological maturity, all of plants were cut off from the soil surface, the different parts of the plant were separated, and the characters related to root and grain yield were measured. Water use efficiency (WUE) determined by the ratio between the economical yield per plant (EYPP) and the total water use (WU).

**Data analysis for phenological and morpho-physiological traits**

Before analyzing data, the normal distribution of the total data and homogeneity of variance were verified using the Kolmogorov–Smirnov test. A t-test was used to test the statistical differences between the means of wheat materials in non-stress and stress conditions using SPSS software.

**Protein extraction**

The flag leaves of tolerant cultivar (Pishgam) were selected for this experiment based on statistical analysis from previous stage. Leaf samples were ground in liquid nitrogen and acetone/trichloroacetic acid (TCA) precipitated according to Damerval et al. (1986) with some modifications. The protein concentrations were measured according to the Bradford assay using BSA as standard.

**Two-dimensional electrophoresis (2-DE)**

2-DE was carried out according to Görg et al. (1988). At the first dimension separation, 320 µL of rehydration solution containing 120 µg of protein was taken up into an immobilized pH gradient (IPG) strip (17 cm, pH 4–7 linear, BioRad) during rehydration over night, then proteins were separated by subjecting the IPG gel strips to electrophoresis for 100 kV-h in a PROTEAN IEF system. At the second dimension separation, the IPG strips were equilibrated in DTT-containing equilibration solution at room temperature for 30 min, sealed at the top of a 12.5% SDS-PAGE gel. Proteins were separated based on molecular weight by using a PROTEAN II Xi Cell two electrophoresis unit (BioRad, USA). Protein spots in analytical gel were visualized by silver staining according to Blum et al.’s protocol (1987).

**Image analysis**

Six silver-stained 2-D gels provided from three replication of each non-treated and treated plants, were scanned using a GS800 calibrated densitometer (Bio-Rad). The scanned images were processed and statistically evaluated with Melanie 7 software (Genebio, Geneva, Switzerland). The spot volumes were normalized as a percentage of the total volume, quantified, and subjected to t-test. Finally, protein spots with significant differences ($p \leq .01$) were considered as regulated proteins for further analysis.

### Table 1. Some characteristics of the plant materials used in the present research.

| Cultivar name | Release year | Breeding phases | Type of wheat | Main breeding characteristics |
|---------------|--------------|-----------------|---------------|------------------------------|
| Pishgam       | 2008         | Crossing Winter | Drought tolerant, cold tolerant, brown and yellow rust tolerant |
| Shahryar      | 2002         | Crossing Winter | Drought susceptible |

Source: Seed and Plant Improvement Institute (S.P.I.I), Karaj, Iran.
Protein spots were carefully excised from the preparative CBB stained gels and subjected to in-gel trypsin digestion according to previous study from our group (Nazari et al. 2018). Afterward, peptide mixtures were analyzed using MALDI-TOF/TOF MS. MALDI TOF TOF (Framingham, MA, USA). Data acquisition and data processing were respectively done using a TOF TOF Series Explorer and Data Explorer (both from AB Sciex). Rehydration of the gels after storage in 15% (v/v) acetonitrile, 20% (v/v) methanol, 1% (v/v) trifluoroacetic acid and mixed with the peptide sample at 1:1 ratio (v/v).

**In-gel digestion and protein identification**

**Results**

**Phenological and morpho-physiological responses of wheat to moisture stress**

Overall, imposed moisture stress significantly (p < 0.05) decreased the whole phenological and morpho-physiological traits, except excised leaf water retention (ELWR) and water use efficiency (WUE) that increased in both wheat cultivars (Table 2). Some of the traits related to root-characters including root dry weight (RDW) and root to shoot dry weight ratio (RDW/SDW) only increased in Pishgam cultivar (Table 2). As expected, mean values of the most measured traits in Pishgam (drought-tolerant cultivar) was significantly (p < 0.05) higher than Shahryar (drought-sensitive) under different moisture conditions (Table 2). High mean values of measured traits in Pishgam, especially economical yield per plant (EYPP), yield-related trait and water use efficiency (WUE) indicate that its productivity and moisture stress-tolerance was significantly greater than Shahryar cultivar under different moisture conditions. As tolerance to moisture stress in Pishgam has previously been reported by the S.P.I.I, according to our results, tolerance to moisture stress was again confirmed in this cultivar. Therefore, its flag leaves were used in the proteomic experiment to evaluate the changes on flag leaf proteome in response to imposed moisture stress.

**Table 2.** Mean comparison of two wheat cultivars subjected to non-stress and moisture stress conditions.

| Characters              | Abbreviation | Non-stress          | Moisture stress       |
|-------------------------|--------------|---------------------|-----------------------|
| Days to heading         | DTH          | Pishgam: 112.5 ± 3.5| Shahryar: 99.66 ± 2.66| Pishgam: 101 ± 2.52 | Shahryar: 91 ± 2.01 |
| Days to anthesis        | DTA          | 123 ± 3.22          | 106 ± 2.51            | 116 ± 2.35             | 103 ± 1.88 |
| Days to maturity        | DTM          | 172 ± 3.72          | 163 ± 2.53            | 162 ± 2.66             | 153 ± 2.23 |
| Grain filling period    | GFF          | 49 ± 1.23           | 57 ± 1.33             | 48 ± 1.16              | 50 ± 1.28 |
| Chlorophyll content (%) | SPAD         | 41.56 ± 1.53        | 40.89 ± 1.15          | 53.33 ± 1.74           | 46.43 ± 1.73 |
| Plant height (cm)       | PH           | 81.55 ± 2.56        | 70.94 ± 2.74          | 67.61 ± 1.78           | 59.33 ± 1.68 |
| Peduncle length (cm)    | PEL          | 25.12 ± 1.05        | 24.29 ± 1.18          | 24.50 ± 1.36           | 21.33 ± 1.57 |
| Tiller number per plant | TN           | 5 ± 0.65            | 4.22 ± 0.54           | 4 ± 0.38               | 3.33 ± 0.22 |
| Leaves number per plant | LN           | 18 ± 1.12           | 25.50 ± 1.50          | 17 ± 1.75              | 13.66 ± 1.58 |
| Fertile spikes number per plant | FSNPP     | 3.11 ± 0.14         | 2 ± 0.17              | 1.83 ± 0.44            | 1.67 ± 0.26 |
| Spicle number per spike | SPNPS        | 20.94 ± 2.47        | 18 ± 1.83             | 13.61 ± 1.53           | 10.25 ± 1.13 |
| Seed number per main spike | SNPMS    | 33.44 ± 1.61        | 23.83 ± 1.05          | 25.72 ± 1.44           | 15.89 ± 1.37 |
| Seed number per plant   | SNPP         | 63.45 ± 1.64        | 41.84 ± 1.16          | 35.44 ± 1.22           | 26 ± 1.28 |
| Main spike weight (g)   | MSW          | 2.41 ± 0.16         | 1.75 ± 0.08           | 2 ± 0.19               | 1.53 ± 0.11 |
| Seed weight per main spike (g) | SFWMS      | 1.83 ± 0.03         | 1.23 ± 0.02           | 1.23 ± 0.06            | 0.66 ± 0.03 |
| Peduncle weight (g)     | PEW          | 0.66 ± 0.02         | 0.62 ± 0.01           | 0.58 ± 0.03            | 0.33 ± 0.07 |
| Main stem weight (g)    | MSTW         | 2.49 ± 0.34         | 2.23 ± 0.25           | 1.68 ± 0.25            | 1.13 ± 0.24 |
| 1000-grain weight (g)   | TGW          | 50.57 ± 1.79        | 39.02 ± 1.02          | 48.02 ± 1.67           | 34.76 ± 1.11 |
| Economical yield per plant (g) | EYPP       | 2.94 ± 0.02         | 1.54 ± 0.02           | 1.74 ± 0.03            | 1.44 ± 0.02 |
| Biological yield per plant (g) | BYPP     | 19.55 ± 1.27        | 15.60 ± 1.34          | 9.57 ± 1.13            | 8.72 ± 1.19 |
| Plant harvest index (%) | PHI          | 23.08 ± 1.20        | 8.76 ± 0.89           | 18.50 ± 1.40           | 4.23 ± 0.73 |
| Leaf area index (cm²)   | LAI          | 18.37 ± 1.31        | 15.87 ± 1.38          | 15.87 ± 1.44           | 12.15 ± 1.15 |
| Relative water content (%) | RWC        | 98.38 ± 2.64        | 84.10 ± 1.70          | 83.24 ± 1.51           | 82.67 ± 1.66 |
| Excised leaf water retention (%) | ELWR     | 179.06 ± 3.67       | 97.83 ± 2.23          | 241.63 ± 3.46          | 263.23 ± 4.33 |
| Water use (l)           | WU           | 14210 ± 14.42       | 15410 ± 15.65         | 7772 ± 8.37            | 8868 ± 9.33 |
| Water use efficiency (g/l) | WUE         | 0.08 ± 0.01         | 0.04 ± 0.01           | 0.09 ± 0.02            | 0.05 ± 0.01 |
| Main root length (cm)   | MRL          | 44.55 ± 2.18        | 45.44 ± 1.55          | 34.33 ± 1.15           | 26.75 ± 1.18 |
| Root volume (cm³)       | RV           | 44.66 ± 0.50        | 17.33 ± 0.33          | 13.08 ± 0.68           | 3.67 ± 0.24 |
| Root dry weight (g)     | RDW          | 43.33 ± 0.02        | 5.92 ± 0.05           | 4.95 ± 0.01            | 0.92 ± 0.01 |
| Root area (cm²)         | RA           | 88.31 ± 1.16        | 92.78 ± 1.19          | 77.30 ± 1.19           | 27.93 ± 0.69 |
| Root to shoot dry weight ratio | RDW/SDW   | 0.27 ± 0.02         | 0.42 ± 0.02           | 0.59 ± 0.06            | 0.08 ± 0.01 |

Notes: Data are expressed as the mean of all plants in three replications ± standard error. For two cultivars within the same conditions, values followed by different letters are significantly different (p ≤ 0.05).
expressed ($p < .05$) between control and moisture stress-treated plants; 23 spots were upregulated and 10 spots were downregulated by the moisture stress (Figure 1).

Database search and functional classification of moisture stress-responsive proteins

11 spots of 33 spots with more differences (at least two-fold) were selected for identification. These spots were excised from the gels, in-gel digested by trypsin, and analyzed by MALDI-TOF/TOF mass spectrometry. According to Mass data and analyzing them using Mascot program and NCBI non-redundant protein database, ten unique proteins were classified into six functional categories: photosynthesis/respiration (36.36%), carbohydrate metabolism (18.18%), energy metabolism (9.09%), chaperon (9.09%), lipid metabolism (9.09%) and unknown function (18.18%), as listed and shown in Table 3, Supplementary Table 1 and Figure 2.

Discussion

Comparison of the effects of imposed moisture stress on different cultivars

In this study, under different moisture stress conditions, the phenological and morpho-physiological characteristics of wheat cultivars changed significantly ($p < .05$). Pishgam,

![Figure 1. Proteome maps of Pishgam cultivar under the control condition (A), Proteome maps of Pishgam cultivar under the moisture stress condition (B). Responsive protein spots to imposed moisture stress, pH range, molecular weight (MW) and IPG length are shown on the gels.](image-url)

| Table 3. Identified moisture responsive proteins using MALDI TOF-TOF in Pishgam cultivar. |
|---|
| **Identity** | **Spot ID** | **Accession number** | **Expression change** | **Coverage %** | **MS Score** | **The pl/MW (kDa)** | **Exp pl/MW (kDa)** |
| Photosynthesis | Ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (plastid) [Melica sublata] | 114 | gi|884999288 | +10.39 | 53 | 220 | 6.13/53.36 |
| | Ribulose bisphosphate carboxylase/oxygenase activase A, partial [Triticum aestivum] | 69 | gi|723047999 | +3.26 | 76 | 171 | 5.35/42.25 | 5.54/46 |
| | Ribulose bisphosphate carboxylase/oxygenase activase A, partial [Triticum aestivum] | 75 | gi|723047999 | +3.17 | 75 | 162 | 5.35/42.25 | 5.86/44 |
| | Ribulose bisphosphate carboxylase small chain PW9, chloroplastic [Aegilops tauschii] | 263 | gi|475591676 | −3.50 | 90 | 142 | 5.85/15.08 | 6.61/18 |
| Carbohydrate metabolism | Fructose-bisphosphate aldolase, chloroplastic-like [Aegilops tauschii] | 93 | gi|1149717357 | +3.92 | 57 | 150 | 6.08/41.83 | 5.40/36 |
| | Phosphoglycerate mutase [Triticum aestivum] | 181 | gi|32400802 | −2.13 | 66 | 158 | 5.3/29.61 | 5.4/30 |
| Energy metabolism | Soluble inorganic pyrophosphatase [Aegilops tauschii] | 152 | gi|1149755635 | −2.82 | 74 | 113 | 5.41/24.49 | 5.43/27 |
| | 20 kDa chaperonin, chloroplastic [Triticum urartu] | 133 | gi|474407512 | −2.83 | 58 | 72 | 6.77/29.80 | 4.99/27 |
| Lipid metabolism | 3-oxoacyl-[acyl-carrier-protein] synthase I, chloroplastic isoform X4 [Glycine max] | 235 | gi|356523620 | +5.52 | 28 | 86 | 8.41/52.55 | 4.39/15 |
| Unknown function | Unnamed protein product [Triticum aestivum] | 167 | gi|669027704 | +5.45 | 56 | 118 | 5.57/27.03 | 5.96/28 |
| | Hypothetical protein GLYMA-13G084400 [Glycine max] | 242 | gi|947069933 | +3.0 | 41 | 75 | 9.10/22.34 | 4.47/12 |

*Number in NCBI SWISS Prot.
$+$ and − indicate protein spots whose abundance increase (+) or decrease (−).
*Percentage of predicted protein sequence covered by matched sequences.
*Statistical probability of true positive identification of the predicted protein calculated by MASCOT.
*TpI/ TMW: Isoelectric point of predicted protein /molecular mass of predicted protein.
*Epl /EMW: Isoelectric point of protein on gel /molecular mass of protein on gel.
Photosynthesis-related proteins

Photosynthesis occurs primarily in the leaves, and some photosynthesis/respiration proteins change by applying moisture stress. In this group, four responsive proteins to moisture stress were identified, including ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco) large subunit (spot 114), chloroplastic ribulose bisphosphate carboxylase small chain (spot 263), and two isoforms of ribulose bisphosphate carboxylase/oxygenase activase A (Rubisco activase A, spots 69 and 75). In this study, rubisco large subunit was strongly upregulated (10.39-fold) and found at the highest levels of protein expression or accumulation by imposed moisture stress. Also, Rubisco small chain (spot 263) was downregulated (3.5-fold) and each of the two rubisco activase isoforms (spots 69 and 75) were upregulated (3.26-fold and 3.17-fold, respectively) by imposed moisture stress. Rubisco is the key enzyme of Calvin cycle (Parry et al. 2003), the only enzyme which can catalyze carbon fixation or oxygenation reaction depending upon the molecular concentration of CO₂ or O₂, and constitutes more than 50% of soluble leaf protein which indicates that this enzyme is important in the plants (Sudhakar et al. 2016). Subunits of rubisco enzyme are susceptible to fragmentation under moisture stress conditions. This phenomenon, possibly leading to isoforms of slightly different molecular weight and isoelectric point, an increase in the number of rubisco large subunit (Salekdeh et al. 2002; Ge et al. 2012; Budak et al. 2013). Some researchers reported upregulation of rubisco large subunits in wheat in response to drought stress (Caruso et al. 2009; Bazargani et al. 2011; Budak et al. 2013; Kamal et al. 2013; Cheng et al. 2015), whereas others found downregulation of rubisco small subunits in wheat (Caruso et al. 2009; Ge et al. 2012) in response to drought stress. Because of rubisco is involved in photosynthetic carbon assimilation and could improve crop yield in C3 plants (Raines 2011), increasing the expression of this enzyme can lead to plant tolerance in moisture stress.

Rubisco activase A identified in 2 spots (spots 69 and 75), probably the causes of this phenomenon are the existence of protein isoforms, post-translational changes and translation from alternative spliced mRNA (Caruso et al. 2008; Maleki et al. 2014; Liu et al. 2015). The amount of active rubisco in a leaf is an important factor regulating the rate of photosynthetic carbon fixation (Servaites et al. 1984). Rubisco actives can activate the rubisco enzyme by carbamylation, remove tight binding inhibitors from rubisco, thus play a key role in regulating photosynthesis in plants (Keown et al. 2013). It seems upregulation of rubisco activase could lead to more carbon assimilation, more products of photosynthesis, and tolerance to moisture stress in plant. Upregulation of rubisco activase was observed in previous study from our group (Nazari et al. 2018) on tolerant Aegilops wheat in response to moisture stress, also in other studies on tolerant plants in response to various stresses (Budak et al. 2013; De Abreu et al. 2014; Maleki et al. 2014; Boustaní et al. 2017; Yan et al. 2017). On the other hand, downregulation of rubisco activase was reported in sensitive plants in response to various stresses (Sobhanian et al. 2010; Beritognolo et al. 2011; Cheng et al. 2015).

Carbohydrate metabolism

Two different proteins were identified in this group including chloroplastic-like fructose-bisphosphate aldolase (spot 93) and phosphoglycerate mutase (spot 181). As shown in Table 3 and Supplementary Table 1, chloroplastic fructose-bisphosphate aldolase was upregulated (3.92-fold) and phosphoglycerate mutase was downregulated (2.13-fold) in response to moisture stress. Fructose-1,6-bisphosphate aldolase (FBA) or FBPA is a key metabolic enzyme in carbon fixation and sucrose metabolism, catalyzes a reversible reaction that splits the aldol fructose 1,6-bisphosphate (FBP), into dihydroxyacetone phosphate (DHAP) and glyceraldehyde-3-phosphate (GAP) during the pathways of glycolysis or gluconeogenesis (Berg et al. 2010). In this study, the increased expression of FBA enzyme could indicate the maintenance of carbohydrate metabolism and signal transduction in tolerant cultivar (Pishgam) during the period of moisture stress and can be identified as moisture stress-responsive marker protein in chloroplast. In several research studies, upregulation of FBA enzyme was reported from crop plants responses to drought and salt stresses (Kamal et al. 2012; Zadražník et al. 2013; Faghani et al. 2015; Nouri et al. 2015).
Phosphoglycerate mutase (PGM), the other protein in this group, is a key enzymatic activity in glycolysis and catalyses the reversible interconversion of 3-phosphoglycerate to 2-phosphoglycerate (Zhao and Assmann 2011). This enzyme was often affected by various stresses, observed with upregulated (Caruso et al. 2008; Vitámvás et al. 2015; Nazari et al. 2018) and downregulated (Caruso et al. 2009) expression, in wheat and barley plants. Various observations may be due to long-term exposure of the plant to environmental stresses, and the plant adapts to changes without the needing to excess energy.

**Energy metabolism**

Soluble inorganic pyrophosphatase (spot 152) was only identified in the group of proteins related to energy metabolism, which was downregulated (2.82-fold) by imposing moisture stress. Soluble inorganic pyrophosphatase participates in the assimilation of mineral nutrients, especially in sulfur metabolism (Schmidt and Jäger 1992), however, there is little knowledge about the details of this protein function in response to various stresses. It has been reported a decreased abundance of this protein in wheat by drought stress (Bazargani et al. 2011), and only detected in drought-stressed plants (Riccardi et al. 1998).

**Chaperone**

In the group of chaperone proteins, the identified protein was chloroplastic 20 kDa chaperonin (spot 133) which was downregulated (2.83-fold) by imposing moisture stress. Chaperones, a group of functional accompanying proteins, are involved in protein folding, assembly, degradation, and protection of nascent proteins during their transport into specific organelles, in both optimal and adverse conditions (Wang et al. 2004; Cheng et al. 2015). Chaperonin 20 kDa (CPN20) is a well-known chloroplast-localized co-chaperonin and help chaperonin CPN60s in protein folding in an ATP-dependent reaction (Horwich et al. 2007). Several studies have shown that CPN20 mediates an antioxidant enzyme (FeSOD) activation in Arabidopsis chloroplasts (Kuo et al. 2013), and negatively regulates abscisic acid signaling in Arabidopsis (Zhang et al. 2013). It seems the reduction of CPN20 was first observed in wheat in response to moisture stress. Also, low-abundance of this protein was reported in maize leaves under drought stress (Zhao et al. 2016).

**Lipid metabolism**

Chloroplastic 3-oxoacyl [acyl-carrier-protein] synthase I (spot 235) was only identified in this group, which was strongly upregulated (5.52-fold) in response to imposed moisture stress. Acyl carrier protein (ACP) is one of the most abundant proteins in the cell which plays an important role in the pathway of fatty acid biosynthesis in the most of organisms. In this pathway, ACP is converted to its active form by acyl carrier protein synthases (AspS) (White et al. 2005). Fatty acids are essential components of cellular membranes, suberin, and cutin waxes which can be structural barriers to the environment. Also, they lead to resistance to environmental stresses through the remodeling of membrane fluidity (Upchurch 2008). In this study, high increased abundance of AspS was observed in wheat in response to moisture stress for the first time which can be shown this protein is probably able to regulate the fluidity of the membrane to maintain the function of the essential proteins in tolerant wheat under moisture stress conditions. In the study of Kamal et al. (2012) on a wheat cultivar, this protein showed an increase in abundance in response to salinity stress.

**Unknown/hypothetical proteins**

In this group, two differentially expressed unknown/hypothetical proteins were detected, including an unnamed protein product (spot 167) with high upregulation (5.45-fold), and hypothetical protein GLYMA-13G084400GLYMA (spot 242) with an upregulation (3-fold). An unknown protein could be defined as a protein whose function has not yet been characterized, and a hypothetical protein could be defined as a protein that is supposed to exist in an organism however its existence has not been shown experimentally (Park et al. 2012). These unknown or hypothetical proteins may contribute to moisture-stress tolerance in wheat cultivar.

**Conclusion**

In this study, we performed a phenological and morpho-physiological analysis with two wheat cultivars subjected to non-stress and moisture stress conditions. After evaluating and determining the moisture stress-tolerant cultivar (Pishgam), its protein changes in flag leaves were identified by 2-DE and MALDI-TOF-TOF MS. Differential expression of 33 moisture stress-responsive proteins revealing the significant effect of moisture stress on the flag leaf proteome of tolerant wheat and the use of that from various signaling pathways and molecular processes in response to moisture stress. Imposed moisture stress significantly increased the abundance of some proteins involved in photosynthesis (rubisco large subunit and rubisco activase A isoform), carbohydrate metabolism (FBA), lipid metabolism (AspSI) and two unknown or hypothetical proteins. In this study, strongly upregulation of AspSI protein in wheat was observed for the first time, in response to moisture stress. Since the highest frequency and level of expression was observed in the upregulated proteins, they can candidate for major roles in tolerance to moisture stress. The abundance of some proteins was significantly reduced by moisture stress which involved in photosynthesis (rubisco small subunit), carbohydrate metabolism (phosphoglycerate mutase), energy metabolism (soluble inorganic pyrophosphatase) and CPN20 chaperonin protein. Furthermore, significant reduction in expression of CPN20 protein in wheat was first observed in this study. The differences in proteome levels may provide an insight into the high tolerance of bread wheat to abiotic stresses. According to the obtained results, proteomics, as a complementary tool, could be useful for identifying candidate genes or proteins to moisture stress-tolerance in bread wheat.

**Disclosure statement**

No potential conflict of interest was reported by the author(s).

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