Genome-wide identification and abiotic stress-responsive pattern of heat shock transcription factor family in *Triticum aestivum* L.

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**Abstract**

**Background:** Enhancement of crop productivity under various abiotic stresses is a major objective of agronomic research. Wheat (*Triticum aestivum* L.) as one of the world’s staple crops is highly sensitive to heat stress, which can adversely affect both yield and quality. Plant heat shock factors (Hsfs) play a crucial role in abiotic and biotic stress response and conferring stress tolerance. Thus, multifunctional Hsfs may be potentially targets in generating novel strains that have the ability to survive environments that feature a combination of stresses.

**Result:** In this study, using the released genome sequence of wheat and the novel Hsf protein HMM (Hidden Markov Model) model constructed with the Hsf protein sequence of model monocot (*Oryza sativa*) and dicot (*Arabidopsis thaliana*) plants, genome-wide TaHsfs identification was performed. Eighty-two non-redundant and full-length TaHsfs were randomly located on 21 chromosomes. The structural characteristics and phylogenetic analysis with *Arabidopsis thaliana*, *Oryza sativa* and *Zea mays* were used to classify these genes into three major classes and further into 13 subclasses. A novel subclass, TaHsfC3 was found which had not been documented in wheat or other plants, and did not show any orthologous genes in *A. thaliana*, *O. sativa*, or *Z. mays* Hsf families. The observation of a high proportion of homeologous TaHsf gene groups suggests that the allopolyploid process, which occurred after the fusion of genomes, contributed to the expansion of the TaHsf family. Furthermore, TaHsfs expression profiling by RNA-seq revealed that the TaHsfs could be responsive not only to abiotic stresses but also to phytohormones. Additionally, the TaHsf family genes exhibited class-, subclass- and organ-specific expression patterns in response to various treatments.

**Conclusions:** A comprehensive analysis of Hsf genes was performed in wheat, which is useful for better understanding one of the most complex Hsf gene families. Variations in the expression patterns under different abiotic stress and phytohormone treatments provide clues for further analysis of the TaHsfs functions and corresponding signal transduction pathways in wheat.

**Keywords:** Genome-wide, Heat shock factor, *Triticum aestivum*, Expression profile, Abiotic stress
Background
Wheat (*Triticum aestivum* L.) is a temperate cereal crop that often encounters heat stress during the reproductive stage in warm-climate wheat production regions. Heat stress has a substantial adverse impact on carbon assimilation and starch synthesis, resulting in the reduction of grain yield and quality. Wheat is one of the world's staple crops, and the most crucial target of wheat breeding is high and stable yield and quality. Wheat is a cool season crop having an optimal daytime growing temperature during its reproductive development of 15°C, and for every degree Celsius above this optimum temperature a reduction of 3–4% in the yield has been observed [1]. Additionally, it is reported that the average global temperature is increasing at a rate of 0.18°C every decade [2], and starch accumulation in wheat grains decreases by >30% at temperatures between 30°C and 40°C [3]. Therefore, the likely impact of heat stress on wheat and the genetic improvement of heat tolerance and its underlying mechanisms have been extensively investigated in recent years.

As sessile organisms, plants could not escape from harmful environments by changing sites, and are exposed to multiple abiotic and biotic stresses frequently [4]. Therefore, a complex stress regulation and response network was developed at biochemical, physiological and molecular levels for stresses adaptation [5, 6]. Many genes which exert a crucial part in this complex stress regulation and response network or confer stress tolerance are mainly regulated by transcription factors [7]. Transcription factors that perform a crucial function in stress signal perception and transduction processes could induce the expression of stress-responsive genes by recognizing and interacting with cis-acting elements in their promoter region, thereby the stress tolerance of plants is enhanced by activated stress signal cascade and whole downstream functional genes of this network [8]. Therefore, transcription factors are considered as potent candidates for developing the next-generation transgenic crops with strong stress tolerance.

Among plant transcription factors, Hsfs have recently attracted particular interest because as terminal components of signal transduction chains, plant Hsfs can regulate the expression of genes involved in various abiotic stress responses [9]. Most types of abiotic stresses disrupt the metabolic balance of cells, resulting in an increase in the production of reactive oxygen species (ROS) [10], and the general concept of HS (heat shock) signaling activation is mainly centered on the disruption of cytosolic protein homeostasis and depletion of the pool of free chaperones [11]. Hsfs and their products protect cells from extreme proteotoxic damage via the expression of molecular chaperones such as heat shock proteins (Hsps). Furthermore, it has been proven that Hsfs participate in some stress-related phytohormone signaling pathways such as abscisic acid (ABA) and salicylic acid (SA) [12, 13].

The plant Hsf gene was firstly cloned from tomato in 1990 [14]. The structure analyses revealed that a modular Hsf contains five conserved domains, including two indispensable domains DNA-binding domain (DBD) and oligomerization domain (OD) comprising central helix-turn-helix (HTH) motif and hydrophobic heptad repeats (HR-A and HR-B) respectively in N-terminal, beside three typical domains nuclear localization signal (NLS), nuclear export signal (NES) and activator peptide motif (AHA) are presented in C-terminal of Hsfs [11, 14]. Based on the number of amino acid residues inserted between HR-A and HR-B, plants Hsf can be grouped into three major classes, namely: A, B, and C. There are 21 and 7 amino acid residues insertion between the HR-A and HR-B region of HsfAs and HsfCs respectively. Compared with HsfAs and HsfCs, HsfBs have a shorter HR-A/B without any amino acid residue insertion [9, 15]. Furthermore, the AHA activation domains are identified in HsfAs uniquely which are absent in HsfBs and HsfCs [16]. HsfBs are characterized by the tetrapeptide-LFV-, which is located within the C-terminal and is predicted to be a repressor motif based on its observed activity in other plant transcription factors [17, 18]. Based on phylogenetic comparisons with the Hsf family of model species such as *Arabidopsis* and rice, plant Hsf family members could further be divided into several subclasses. The specific function of plant Hsf subclasses in model plants also been reported in previous work. *AtHsfA1s* have been proved to be the master regulators of heat stress response in *Arabidopsis*, which could induce the expression of diverse transcription regulators, including other Hsf subclasses (A2, A3, A7, B1 and B2) [19, 20]. *AtHsfA2* not only confers heat and osmotic stress tolerance, but also plays a significant role in the growth and development of plants [21–23]. The tomato (*Lycopersicon esculentum*) HsfA4s are potent activators of heat stress gene expression, whereas HsfA5s act as specific repressors of HsfA4s activity [24]. The function of OsHsfB2b is considered as a negative regulator in response to drought and salt stresses in rice [25]. *OsHsfC1b* is induced by salt, mannitol and ABA, but not by H₂O₂, and play an important role in salt and osmotic stress response [26]. The above results reveal that plants have evolved both subclass-specific and multiple functions in some members of the Hsf family.

The identification of plant Hsf family genes is usually based on the characteristics of the two conserved domains, DBD and HR-A/B, which is the core of the Hsf HMM model, as a query in searching the proteome [27]. Unlike the simple and small Hsf family in animals and yeast, plants have relatively complex and large Hsf gene families. Bread wheat (*T. aestivum*) has one of the most...
complex genomes known to science, with an overall size of more than 17 billion bases [28]. Additionally, the tetraploid emmer wheat (T. turgidum; BBAA) and hexaploid bread wheat (BBAADD) originated within the past few hundred thousand years and ten thousand years ago [29, 30], respectively. It can be predicted that wheat that underwent two rounds of whole genome duplications in recent years must have one of the most complex and largest Hsf family in plants. The recently available T. aestivum genome TGACv1 has allowed the identification of TaHsfs at the genome-wide level [31].

As a major cereal crop, wheat is widely cultivated around the world, following maize in grain, and provides carbohydrates and proteins for approximately 40% of the world’s population [32]. Wheat is hypersensitive to heat stress, particularly at the early grain filling and reproductive stages, significantly limiting wheat production [33]. This study aimed to elucidate the abiotic stress-responsive pattern of TaHsfs and identify candidates for genetic improvement of abiotic tolerance in this species.

Results
Identification of Hsf genes in T. aestivum
The constructed HMM for Hsf was based on the protein sequence of A. thaliana and Oryza sativa, which was queried in BLASTP searches for possible homologous TaHsfs in the T. aestivum proteome. A total of 154 candidate Hsf protein sequences were identified in this process. Subsequently, the putative wheat Hsf protein sequences were surveyed to further determine whether these included a DBD and HR-A/B domain using SMART software (http://smart.embl-heidelberg.de/). Consequently, 74 of the candidate Hsf protein sequences were excluded from further analysis based on the absence of DBD or HR-A/B domains and overlapping genes. Eighty nonredundant wheat Hsfs with the DBD and HR-A/B domains were identified and characterized. Additionally, two members of subclass TaHsfA2 (TaHsfA2–11 and TaHsfA2–18) were obtained by homologous cloning (Fig. 1, Additional file 1). The deduced protein sizes ranged from 209 amino acids (TaHsfB2–2) to 701 amino acids (TaHsfB2–7) (Table 1). Based on the number of amino acid residues inserted into the HR-A/B domain, 82 TaHsfs were classified into three major classes; class A contained the highest number of TaHsf members (40), classes B and C consisted of 16 and 26 TaHsf members, respectively (Fig. 1B).

Except for three TaHsfs (TaHsfA2–11, TaHsfC3–9 and TaHsfC3–11) located on the unanchored scaffolds, 79 nonredundant wheat Hsfs were mapped to 21 wheat chromosomes (Fig. 2, Additional file 2). TaHsfs were distributed among 21 wheat chromosomes, but the number of TaHsf genes on each chromosome extensively differed. The highest number of TaHsf genes (N = 8) was observed on chromosomes 3B and 5A, whereas the lowest number was detected on chromosomes 6A, 6B and 6D, each of them including only one TaHsf gene. Chromosomes 5 and 3 showed the highest density of TaHsf genes, with 19 and 18 members, respectively. Chromosomes 3A and 3B have previously been reported to likely harbor key genes conferring heat tolerance in wheat [34].

Phylogenetic analysis and classification of TaHsf members
To investigate the evolutionary features and characteristics of the TaHsf genes, an unrooted phylogenetic tree was constructed. Phylogenetic analysis was performed based on the Hsfs amino acid sequences of the N-terminal domains of Hsfs, including the DNA-binding domain, the HR-A/B domain and the linker between two domains from T. aestivum, A. thaliana, O. sativa and Z. mays. The three major classes, namely, TaHsfA (green), TaHsfB (yellow) and TaHsfC (blue), could also be clearly distinguished by phylogenetic analysis (Fig. 3). The TaHsf family was further divided into 13 subclasses based on their bootstrap values and phylogenetic relationship with the orthologous genes from O. sativa and A. thaliana (Additional file 3). Class A was further subdivided into seven subclasses, namely, TaHsfA1, TaHsfA2, TaHsfA3, TaHsfA4, TaHsfA5, TaHsfA6 and TaHsfA9. Both classes B and C consisted of three subclasses, which were named as TaHsfB1, TaHsfB2 and TaHsfB4 and TaHsfC1, TaHsfC2 and TaHsfC3, respectively. Although classes A, B and C in both eudicots (Arabidopsis) and monocots were conserved, whereas subclasses HsfA2, A6, A9 and B1 were divided into different clade between monocots and dicots. The monocots and dicots uniquely consist of subclasses B4, C2 and A7, A8, B3 respectively. The monocots T. aestivum, O. sativa and Z. mays followed the same subclassification and were very closely clustered. Subclass A2 had the highest number of Hsfs (N = 18) in the wheat Hsf family, whereas the lowest number of Hsfs (N = 3) was observed in subclasses A1, A5, A6, A9 and B1. A clade of TaHsfC without orthologs detected in other plants was designated as TaHsfC3. All homeologous TaHsf gene groups with a copy on each of the A, B and D homeologous chromosome were closely clustered. The wheat Hsf genes also exhibited a closer phylogenetic relationship with the monocot rice than with the Z. mays and the dicot Arabidopsis.

Gene structure and conserved domains of Hsfs in T. aestivum
To explore the structural diversity of the TaHsf members, the intron-exon organization of each TaHsf gene was analyzed by comparing the cDNA sequences with the corresponding genomic DNA sequence (Fig. 4, Additional file 4). Analysis of the intron-exon boundaries of all TaHsfs indicated a highly conserved organization, particularly within the homeologous TaHsf gene groups.
and same subclass TaHsf members. This observation further validated the precision of the classification. However, the number of exons and introns differed among the TaHsfs, 62 of the 82 Hsf genes exclusively contained two exons, whereas 10 Hsf genes contained three exons. We identified nine intronless genes, and 10 TaHsfs are consisted of a single exon, all of which belonged to subclass TaHsfC3. A previous study showed that most of Hsf DBDs underwent an insertion of a conserved intron, separating the DBD into two parts [9]. 53 of the 82 TaHsfs contained a single intron. Two TaHsfA2 members (TaHsfA2–13, TaHsfA2–15) contained four introns, which is the highest number in the TaHsf family genes. Additionally, the lengths of the TaHsf introns were highly variable, which ranged from 80 bp (TaHsfA4–2) to 5836 bp (TaHsfC1–6).

Conserved motif analysis was conducted using MEME, and 15 motifs were identified in TaHsf family members (Fig. 5, Additional file 5). To further determine the structural characteristics of the TaHsf family members, the SMART online tool was employed to predict the conserved domains. The DBD is the most conserved domain, which is composed of Motifs 1 and 2 in most TaHsfs (76 of 82). Six TaHsfs had shorter DBD domains. TaHsfC1–1, TaHsfC2–3 and TaHsfC3–12 only contained Motif 1; TaHsfA6–1, TaHsfC3–11 and TaHsfC3–13 only contained Motif 2. TaHsfA2–18, TaHsfC1–1 and TaHsfC2–3 had partial α1-helices; TaHsfA6–1 and TaHsfC3–11 did not contain β4-sheets; TaHsfC3–12 and TaHsfC3–13 lacked α1, β3 and β3, β4, respectively (Fig. 1A). The HR-A/B regions are indispensable domains that are characterized by the predicted coiled-coil structure, which had two typical motifs (3 and 4) in the TaHsf family. Motif 3 is the predominant motif, corresponding to the HR-A/B regions in 76 members of the TaHsf family. The DBD and HR-A/B
Table 1 Details of various TaHsfs, including chromosome locations, transcript ID, protein sequence length (number of amino acids, AA), molecular weight (MW), nuclear export signal (NES), nuclear localization signal (NLS), activator motifs (AHA) and endoplasmic reticulum (ER) membrane retention signals

| Gene name | Chromosome | Transcript ID | No. of AA | MW (kDa) | NES | NLS | AHA Motif | ER Membrane Retention Signals |
|-----------|------------|---------------|-----------|----------|-----|-----|-----------|--------------------------------|
| TaHsfA1–1 | 4A         | AA058750      | 521       | 57.36    | (126)RRKP(252)KKRR | (464)DSEFWEQFLCA |
| TaHsfA1–2 | 5B         | AA128550      | 529       | 58.19    | (129)RRKP(256)KKRR | (472)DSEFWEQFLCA |
| TaHsfA1–3 | 5D         | AA140370      | 522       | 57.46    | (124)RRKP(249)KKRR | (465)DSEFWEQFLCA |
| TaHsfA2–1 | 5A         | AA1215960     | 346       | 38.99    | (157)LKRDRQLLM | (125)KRRKP(223)RKELEDAISNKRRRRID | (313)DDFWEDLLHE | (2)SHRM(342)AEKL |
| TaHsfA2–2 | 5B         | AA12081150    | 353       | 39.73    | (164)KRRDRQLLM | (132)KRRKP(230)RKELEDAISNKRRRRID | (320)DDFWEDLLHE | (2)SHRM(349)AQKM |
| TaHsfA2–3 | 5D         | AA142920      | 348       | 38.91    | (159)KRRDRQLLM | (127)KRRKP(225)RKELEDAISNKRRRRID | (315)DDFWEDLLHE | (344)AEKL |
| TaHsfA2–4 | 2A         | AA0357000     | 413       | 45.51    | (130)RRRKP(229)RKELHDAISKKRRRRID | (370)DNFWELLNK |
| TaHsfA2–5 | 2B         | AA0492290     | 405       | 44.93    | (131)KRRKP(239)KKRRRR | (362)DNFWELLNK |
| TaHsfA2–6 | 2D         | AA0573260     | 412       | 45.44    | (130)KRRKP(229)RKELHDAISKKRRRRID | (369)DNFWELLNK |
| TaHsfA2–7 | 4A         | AA1018260     | 341       | 39.61    | (133)KRRKP(241)PTKR KRRR | (311)DDFWELLSE | (2)DRVL |
| TaHsfA2–8 | 4B         | AA1062420     | 341       | 39.6     | (133)KRRKP(241)PTKR KRRR | (311)DDFWELLSE | (2)DRVL |
| TaHsfA2–9 | 4D         | AA126990      | 341       | 39.5     | (133)KRRKP(241)PTKR KRRR | (311)DDFWELLSE |
| TaHsfA2–10| 5A         | AA1256510     | 372       | 41.14    | (320)LLSLGLE | (153)KRRRP(259)RRKELAEALLSKKRGRP | (314)ESFWKELSL |
| TaHsfA2–11| U          | 373           | 41.44    | (322)LLSLGLE | (153)KRRRP(261)RRKELA DALLSKKRGRP | (314)ESFWKELSL |
| TaHsfA2–12| 5D         | AA1405180     | 377       | 41.6     | (326)LLSLGLE | (153)KRRRP(261)RRKELA DALLSKKRGRP | (320)ESFWKELSL |
| TaHsfA2–13| 1A         | AA0048030     | 364       | 41.03    | (166)IDRLKRDKNL II | (433)DTFFQSSCSG |
| TaHsfA2–14| 1B         | AA0114190     | 364       | 41.03    | (166)IDRLKRDKNL II | (435)KRRKP(234)RKELEDAISKKRRRRPI | (318)NDWAEFLFD |
| TaHsfA2–15| 1D         | AA0199110     | 370       | 42.05    | (172)IDRLKRDKNL II | (441)KRRKP(240)RKELEDAISKKRRRRPI | (324)NDWAEFLFD |
| TaHsfA2–16| 3A         | AA0636800     | 397       | 44.24    | (129)KRRRP(237)KKRRR | (354)DDFWELLMSR |
| TaHsfA2–17| 3B         | AA0745290     | 433       | 48.01    | (128)KRRRP(236)KKRRR | (353)DDFWELLMSR |
| TaHsfA2–18| 3D         | 407           | 46.23    | (369)LDVYKLDD | (81)KRRRP(189)KKRRR(353)KKRRR | (306)DDFWELLMSR |
| TaHsfA3–1 | 2A         | AA0373860     | 467       | 51.63    | (438)FDALDGDGLHL | (402)DTFFQSSCSG | (463)GNMK |
| TaHsfA3–2 | 2A         | AA0367680     | 502       | 55.45    | (437)DTFFQSSCSG | (498)GNMK |
| TaHsfA3–3 | 2B         | AA0455720     | 475       | 52.09    | (29)LEPKLEM | (483)DTFFQSSCSG |
| TaHsfA3–4 | 2D         | AA0576700     | 499       | 54.79    | (25)LLLEPKLEM(47)EALDDGDGLHL | (484)DTFFQSSCSG | (495)GNMK |
| TaHsfA4–1 | 1A         | AA0026630     | 443       | 49.57    | (263)MELALVSM | (112)KRRKP(220)KKRRR | (386)DLDFWFRFLTD | (439)AQK |
| TaHsfA4–2 | 1B         | AA0108940     | 445       | 49.88    | (263)MELALVSM | (110)KRRKP(219)KKRRR | (388)DLDFWFRFLTD | (441)AQK |
| TaHsfA4–3 | 1D         | AA0193800     | 442       | 49.71    | (266)MELALVSM | (110)KRRKP(218)KKRRR | (385)DLDFWFRFLTD | (438)AQK |
| TaHsfA4–4 | 3A         | AA0661360     | 432       | 48.38    | (133)KCDNASLKL | (100)KRRKP(198)KKRRR | (370)DGFWQQLFLE | (428)AEK |
| TaHsfA4–5 | 3B         | AA0700580     | 441       | 49.47    | (133)KCDNASLKL | (100)KRRKP(198)KKRRR | (379)DGFWQQLFLE | (437)AEK |
| TaHsfA4–6 | 3D         | AA0847210     | 433       | 48.46    | (133)KCDNASLNL | (100)KRRKP(198)KKRRR | (370)DGFWQQLFLE | (429)AEK |
| TaHsfA5–1 | 6A         | AA1567660     | 458       | 49.88    | (340)LI TL | (109)KRRKP(212)HKKRRR | (414)DNFWEQFLTE | (454)EQLK |
Table 1  Details of various TaHsfs, including chromosome locations, transcript ID, protein sequence length (number of amino acids, AA), molecular weight (MW), nuclear export signal (NES), nuclear localization signal (NLS), activator motifs (AHA) and endoplasmic reticulum (ER) membrane retention signals (Continued)

| Gene name | Chromosome | Transcript ID | No. of AA | MW (kDa) | NES | NLS | AHA Motif | ER Membrane Retention Signals |
|-----------|------------|---------------|-----------|----------|-----|-----|-----------|--------------------------------|
| TaHsfA5-2 | 6B         | AA1650270     | 455       | 49.93    | (340) | LTL | (109)RKR(212)HKKRR | (414)DNFWEQFLTE (451)EQLK |
| TaHsfA5-3 | 6D         | AA1745290     | 458       | 49.87    | (340) | LTL | (109)RKR(212)HKKRR | (414)DNFWEQFLTE (454)EQLK |
| TaHsfA6-1 | 7A         | AA1799040     | 310       | 33.78    | (202) | KKK | (247)DMIWYELLEE | (299)DMYWYELLEE |
| TaHsfA6-2 | 7B         | AA2004680     | 351       | 37.98    | (136) | RRR | (250)KKKRR | (299)DMYWYELLEE |
| TaHsfA6-3 | 7D         | AA1009080     | 42.86     | (165) | LKRDKSLLMQQL | (115)KRR | (242)KKKRR | (337)MLHWFEDGE |
| TaHsfA9-2 | 4B         | AA1071190     | 42.88     | (172) | LMKQLVLDDL | (124)KRKKRP(232)RR | (247)DMYWYELLEE |
| TaHsfA9-3 | 4D         | AA142510      | 42.93     | (172) | LMKQLVLDDL | (124)KRKKRP(232)RR | (247)DMYWYELLEE |
| TaHsfB1-1 | 5A         | AA1196580     | 32.15     | (209) | LDVRQLDLRLLM | (114)RRR | (124)KRKKRP(232)RR | (247)DMYWYELLEE |
| TaHsfB1-2 | 5B         | AA1307960     | 32.29     | (211) | LDVRQLDLRLLM | (117)RRR | (247)DMYWYELLEE |
| TaHsfB1-3 | 5D         | AA1418380     | 32.06     | (211) | LDVRQLDLRLLM | (117)RRR | (247)DMYWYELLEE |
| TaHsfB2-1 | 2A         | AA0301090     | 31.99     | (262) | MRTERSDLNWL | (89)RKGKRLGAQQRKGS(165)RR | (247)DMYWYELLEE |
| TaHsfB2-2 | 2D         | AA2166870     | 22.73     | (209) | LDVRQLDLRLLM | (114)RRR | (247)DMYWYELLEE |
| TaHsfB2-3 | 7A         | AA1189330     | 41.11     | (362) | LALENPDLSL | (103)RKGEKQLLCEIHRRKTS (6)ERCG |
| TaHsfB2-4 | 7B         | AA1293070     | 41.46     | (362) | LALENPDLSL | (103)RKGEKQLLCEIHRRKTS (6)ERCG |
| TaHsfB2-5 | 7D         | AA1418740     | 41.4      | (362) | LALENPDLSL | (103)RKGEKQLLCEIHRRKTS (6)ERCG |
| TaHsfB2-6 | 5A         | AA0058470     | 40.4      | (269) | LELDMV | (115)RKGAKHLAEIHRKSS(299)KKKRR |
| TaHsfB2-7 | 5B         | AA2122890     | 35.26     | (269) | LELDMV | (116)RKGAKHLAEIHRKSS(299)KKKRR |
| TaHsfB2-8 | 5D         | AA2122890     | 35.32     | (269) | LELDMV | (116)RKGAKHLAEIHRKSS(299)KKKRR |
| TaHsfB4-1 | 3A         | AA1189330     | 41.36     | (362) | LALENPDLSL | (103)RKGEKQLLCEIHRRKTS (6)ERCG |
| TaHsfB4-2 | 3B         | AA1293070     | 41.46     | (362) | LALENPDLSL | (103)RKGEKQLLCEIHRRKTS (6)ERCG |
| TaHsfB4-3 | 3D         | AA1418740     | 41.4      | (362) | LALENPDLSL | (103)RKGEKQLLCEIHRRKTS (6)ERCG |
| TaHsfB4-4 | 3A         | AA0058470     | 40.4      | (269) | LELDMV | (115)RKGAKHLAEIHRKSS(299)KKKRR |
| TaHsfC1-1 | 3A         | AA0614700     | 31.16     | (175) | QQAAEKKLQRMHL | (105)PRVRRK |
| TaHsfC1-2 | 3B         | AA0722380     | 35.73     | (103) | GLI | (109)PRVRRK |
| TaHsfC1-3 | 3D         | AA0860000     | 35.39     | (103) | GLI | (109)PRVRRK |
| TaHsfC1-4 | 3A         | AA0670400     | 31.16     | (175) | QQAAEKKLQRMHL | (105)PRVRRK |
| TaHsfC1-5 | 3B         | AA0722680     | 33.89     | (109) | PRVRRK |
| TaHsfC1-6 | 3D         | AA0849120     | 25.63     | (109) | PRVRRK |
| TaHsfC1-7 | 3A         | AA0614700     | 26.06     | (103) | PRVRRK |
| TaHsfC1-8 | 3B         | AA0780810     | 24.69     | (103) | PRVRRK |
| Gene name | Chromosome | Transcript ID | No. of AA | MW (kDa) | NES | NLS | AHA Motif | ER Membrane Retention Signals |
|-----------|------------|---------------|-----------|----------|-----|-----|-----------|--------------------------------|
| TaHsfC1–9 | 3D         | AA0873000     | 241       | 26.41    |     |     | (103)PLIVRKK(203)PDVRKK |                               |
| TaHsfC2–1 | 5A         | AA2086000     | 268       | 30       | (197)VKRLRLQL | (118)KRPK(162)RRRK |                               |
| TaHsfC2–2 | 7A         | AA1794130     | 266       | 28.23    | (168)LAFLLTVW | (103)RRGTKAVGGGGKKRKDA(162)RKPK |                               |
| TaHsfC2–3 | 7B         | AA1874420     | 244       | 26.12    | (146)LAFLLTI | (140)RKPK |                               |
| TaHsfC2–4 | 7D         | AA2011440     | 265       | 28.18    | (167)LAFLLTI | (101)RRGTKAAAGGGGGKRKDA(161)RKPK |                               |
| TaHsfC3–1 | 4D         | AA1147440     | 276       | 30.27    | (169)LMLAFLL | (107)PRVRRRR(206)KRRRLLDGEQVSKKKMR |                               |
| TaHsfC3–2 | 5A         | AA1225450     | 274       | 30.36    | (167)LMLAFLL | (163)RRPK(204)KRPRLLLDGEQVQMGKK |                               |
| TaHsfC3–3 | 4B         | AA1084070     | 257       | 28.73    | (169)LMLAFLL | (105)PRVRRR(199)PARARPR |                               |
| TaHsfC3–4 | 3B         | AA0791830     | 274       | 29.98    | (168)LMLAFLL | (103)PRVRRR(203)KRPRLLLDGEQVGKK |                               |
| TaHsfC3–5 | 3B         | AA0777840     | 230       | 25.57    | (170)LMLAFLL | (108)PRVRRRR(202)KRPR |                               |
| TaHsfC3–6 | 4B         | AA1034080     | 257       | 28.5     | (207)LLLNGEVQM | (104)PRVRRR(203)KKPR |                               |
| TaHsfC3–7 | 4B         | AA1067170     | 268       | 30       |     |     | (158)RRPK |                               |
| TaHsfC3–8 | 4B         | AA1045990     | 273       | 29.46    | (174)LKV | (165)RRPK(205)KRPR |                               |
| TaHsfC3–9 | U          | AA1138930     | 276       | 29.86    | (177)LMLAFLL | (171)RRPK(211)KRPR |                               |
| TaHsfC3–10| 5A         | AA1241720     | 273       | 30.18    |     |     | (173)RRPK |                               |
| TaHsfC3–11| U          | AA1138960     | 248       | 26.43    |     |     | (141)RRPK(173)KRPR |                               |
| TaHsfC3–12| 3B         | AA0805650     | 237       | 26.13    | (133)LMLAFLL | (127)RRPK(168)KRPR |                               |
| TaHsfC3–13| 4B         | AA1040480     | 248       | 26.15    |     |     | (145)RRPK |                               |
domains were detected in all members of the TaHsf family. Additionally, Motif 7 corresponded to the NLS and Motif 6 represented the AHA. Overall, the structure of the TaHsf proteins was conserved among the TaHsf family members.

Depending on the balance between nuclear import and export, the intracellular distribution of Hsfs dynamically changes between the nucleus and cytoplasm [35]. The NES, NLS and ER membrane retention signals at the C-terminal of various Hsfs are required for regulating Hsfs subcellular localization. The NESs in 51 TaHsfs were predicted using NetNES, which included 25 TaHsfAs, 10 TaHsfBs and 16 TaHsfCs. The NLS domain was found in 50 members of TaHsf family, including 32 TaHsfAs, 11 TaHsfBs and 7 TaHsfCs; six and one bipartite NLS were found in classes B and C, respectively. Moreover, ER membrane retention signals were detected in 20 members of the TaHsf family. Additionally, the AHA domains, which were identified by sequence comparison, were detected in all TaHsfAs in the center of the C-terminal activation domains. However, these domains were not identified in TaHsfBs and TaHsfCs (Table 1). The tetrapeptide-LFGV domain, which is characteristic of HsfBs, was present in all TaHsfBs, except for TaHsfB2–2,
in the C-terminal of the protein, and is predicted as the repressor motif of transcription [18].

**Transcription profiles of the Hsf genes in *T. aestivum***

Different abiotic stresses and phytohormone signal networks interact and share some common elements that form potential “nodes” for crosstalk [8]. The multifunctional plant Hsf genes are considered as nodes or cross-points that connect several pathways and simultaneously participate in various abiotic and phytohormone signaling pathways. Therefore, the results of transcriptome sequencing analysis under five different treatments (H2O2, heat stress, abscisic acid, salicylic acid, polyethylene glycol) and control in leaf and root tissues are shown in Fig. 6. Transcripts per million (TPM) values were used to measure the transcription level of the TaHsfs (Additional file 6).

The transcription patterns of the 80 TaHsf genes revealed that TaHsfs are responsive to all the phytohormone and stress treatments to different degrees, and almost all TaHsf genes are expressed in the two tissues under different treatments, except for TaHsfA3–1, TaHsfC1–(4–6), TaHsfC2–1 and TaHsfC3–6. The transcription patterns of the TaHsf genes under different treatments and in different tissues significantly differed. However, TaHsfs from the same class, subclass, and most homoeologous TaHsf genes exhibited some degree of similarity in expression patterns.

In *T. aestivum* leaf tissues, the expression of TaHsf family was hardly detectable under normal conditions. TaHsfAs were the most inducible TaHsf genes that were strongly upregulated by both H2O2 and HS treatments. TaHsfC members were mainly induced by ABA treatment. TaHsfB members were induced by H2O2, HS and ABA treatments. The highest expression level in the TaHsf family were deteted in TaHsfA6s under H2O2 and heat treatment in leaf tissues, whereas the expression of TaHsfA6s could hardly be detected in the control and other treatments. Additionally, seven members of subclass A2 (TaHsfA2–1, TaHsfA2–10, TaHsfA2–11, TaHsfA2–12, TaHsfA2–13, TaHsfA2–14 and TaHsfA2–15) were also strongly induced by H2O2 and heat treatments. The members of TaHsfB were responsive to H2O2, HS and ABA treatments. Subclass TaHsfB1 and TaHsfB2–(1–5).
were more sensitive to H$_2$O$_2$ and heat treatments than ABA treatment. However, TaHsfB2-(6–8) were more sensitive to ABA treatment. The members of the TaHsfC subclass were only sensitive to ABA treatment. The members in subclass TaHsfA1 were more sensitive to SA and PEG (polyethylene glycol) treatments than the other subclasses. We could detect very weak expression of TaHsfA3s and TaHsfA4-(1–3), but could hardly detect any expression of TaHsfB4s and TaHsfC1-(1–6) in the leaf tissues under all treatments and the control. Most of the TaHsf members were upregulated after treatment, except for TaHsfC1–7, TaHsfC2–2, TaHsfC2–3 and TaHsfC2–4, which were downregulated after H$_2$O$_2$, HS, SA and PEG treatments.

In T. aestivum root tissues, the class C TaHsf family genes were more sensitive to ABA treatment, among them, TaHsfC3–4 showed the highest expression levels.
Both upregulated and downregulated response modes were observed in the TaHsfBs under ABA treatment, TaHsfB1s were up-regulated, whereas TaHsfB2-(6–8) were downregulated. Most of TaHsfAs were insensitive to ABA treatment, except TaHsfA3s and TaHsfA4–1, TaHsfA4–2 which were only induced by ABA treatment in the root tissues. Members of TaHsfA were only sensitive to H$_2$O$_2$, which were upregulated after treatment. TaHsfA2–10, TaHsfA2–12, TaHsfA2–13, TaHsfA2–14 and TaHsfA2–15 exhibited relatively higher basic transcription levels in the root tissues under control conditions, and were only upregulated under H$_2$O$_2$ treatment, but were suppressed under the other treatments. SA treatment could only upregulate subclass TaHsfA1, as well as inhibit the expression of TaHsf5, which showed high basic transcription level. Most of the upregulation and
high transcription level under PEG treatment were detected in TaHsfC3s, whereas the inhibited expression under PEG treatment was observed in TaHsfC2–2, TaHsfC2–3 and TaHsfC2–4. The TaHsfA6 subclass exhibited H2O2 treatment-specific expression and was hardly detected in the other treatments and control. TaHsfC1–(1–3), TaHsfC1–(7–9), TaHsfC3–10 and TaHsfC3–13 exhibited ABA treatment-specific expression in the root tissues, among them, TaHsfC1–(1–3), TaHsfC3–7, TaHsfC3–12 and TaHsfC3–13 exhibited root-specific expression and
were hardly detected in the leaves. In general, TaHsf in the root tissues had a higher expression diversity than the leaves.

The expression pattern confirmation of selected TaHsf by qRT-PCR

Nine TaHsf genes (TaHsfA1–3, TaHsfA2–1, TaHsfA2–7, TaHsfA2–10, TaHsfA2–12, TaHsfA2–13, TaHsfA2–17, TaHsfB2–6, TaHsfC3–4) from three major classes, were selected for examination of their expression pattern under control and treatments using qRT-PCR, which carried out using wheat seedling leaf treated with H2O2, HS, ABA, PEG, SA and control. The qRT-PCR result of selected TaHsf expression pattern (Fig. 7) showed a high level of consistency with the results of RNA-seq analysis.

Both RNA-seq and qRT-PCR results showed that the TaHsfA1–3 and TaHsfA2–17 were insensitive to all treatments. The other TaHsfA members (TaHsfA2–1, TaHsfA2–7, TaHsfA2–10, TaHsfA2–12, TaHsfA2–13, TaHsfA2–17, TaHsfB2–6, TaHsfC3–4) could only be significantly induced by H2O2 and HS treatments, which could also be observed in RNA-seq analysis. The qRT-PCR results of TaHsfB2–6 revealed that it could be upregulated under H2O2, HS and ABA treatments and insensitive to SA and PEG, and it is more sensitive to ABA than to H2O2 and HS treatments, both of which were consist with RNA-seq results. qRT-PCR and RNA-seq results showed that TaHsfC3–4 could only be intensely induced by ABA treatment.

![Fig. 7 Relative expression level of selected TaHsf analyzed by qRT-PCR under H2O2, heat stress, ABA, SA, PEG treatments. Each bar value represents the Mean ± SD of triplicate experiments](attachment:image.png)
Discussion

Wheat has a large number of Hsf family genes

The common bread wheat, *T. aestivum*, has one of the most complex genomes known to science. The allohexaploid bread wheat genome consists of three closely related subgenomes (A, B and D) and an overall size of at least 17 billion bases [28]. Based on genetic similarity, the 21 pairs of wheat homeologous chromosomes are divided into seven homeologous groups, each containing one pair of chromosomes from the A, B and D subgenomes. The homeologous chromosome groups have similar sets of genes (syntenic genes) and homologous DNA sequences; therefore, the *T. aestivum* genome contains several groups of homeologous genes that have three highly identical sequences in each subgenome [36].

The huge *T. aestivum* genome and its seven homeologous chromosome groups provide a resource for generating a large TaHsf family. There are at least 56 TaHsfs in the *T. aestivum* genome. Because of the unavailability of the *T. aestivum* genome sequence at that time, the 56 TaHsfs were identified using a BLAST search of a limited database, which was collected from individual EST databases. Thus, the 56 TaHsfs that are identified and show high sequence redundancy, and some sequences were of partial length [37]. The present study employed an advanced method in identifying TaHsfs at the genome-wide level. A new HMM model of plant Hsf protein sequences was constructed, and 80 nonredundant, full-length TaHsfs were identified.

Studies on plant Hsf families have reported more than 20 species to date. The number of Hsf family genes widely differs among plants. There are 21 Hsfs-encoding genes in Arabidopsis, 24 in tomato [11], 25 in pepper [38], 25 in maize [27], 29 in Chinese white pear [39], 64 in *Brassica napus* [40]. Scharf et al. earlier suggested that the expansion of Hsfs in angiosperms is presumably the result of gene duplications and whole-genome duplications (WGDs) at different time points during evolution. Lineage-specific WGDs within the angiosperms are presumably the cause of the observed variations in the number of Hsfs among plant species [11]. The allohexaploid bread wheat genome was generated by the fusion of the *T. urartu* (subgenome A), *Aegilops speltoides* (subgenome B) and *A. tauschii* (subgenome D) genomes several hundred thousand years ago [41]. Approximately 60.1–61.3% of genes in the A, B and D subgenomes have orthologs in all the related diploid genomes [42]. Due to two rounds of allopolyploidy, the TaHsfs have tripled. Although the Hsf genes were always under strong purifying selection pressure [43], the allopolyploid process recently occurred, thus most of the TaHsfs were not eliminated by evolution. The present study found that 21 TaHsf homeologous gene groups consisted of 63 TaHsfs, which included three copies of genes that are located on each homeologous chromosome (A, B and D) and show high nucleotide sequence identity. These results indicate that the high number of TaHsfs in allohexaploid bread wheat was probably generated by allopolyploidy, whereas only 19 of the 82 TaHsf genes did not consist of an integrated homeologous group in the wheat genome, which was probably due to the incomplete genome sequence or gene loss during fractionation from ploidy [44]. The allotetraploid peanut [45] and cotton [46] also have relatively large numbers of Hsf genes in their genome. The number of Hsf genes in *Brassica napus* (genome AACC), which was formed by recent interspecific hybridization, like wheat, has increased to 64 [40]. Therefore, the allopolyploid process, which resulted from the fusion of genomes, apparently contributed to the expansion of the plant Hsf family, which recently underwent allopolyploidy. In addition, allopolyploidy is proved to increase abiotic stress tolerance in plants [47, 48], and the larger number of members of the Hsf family may contribute to the higher abiotic stress tolerance.

Classification and phylogenetic analysis of the TaHsf proteins

The present study showed that all 82 TaHsfs contained the highly conserved domains DBD and HR-A/B, which are essential for its transcriptional functions. All TaHsf genes could be divided into classes A, B and C based on the number of amino acid residues inserted between HR-A and HR-B.

For further classify TaHsfs into subclasses, an unrooted Neighbor-joining (NJ) tree was constructed using previously characterized Hsf and TaHsf families (Fig. 3). The TaHsfs were thus named based on the corresponding subclass name of the orthologous gene in model plants. Additionally, gene orthology analysis was also used as a preliminary method to investigate the function of TaHsfs [49]. The result showed that the class A were the predominant class. Whose number of genes and subclasses were the largest in the Hsf family genes in both monocots and dicots. Distinct differences in subclass species were also observed between monocots and dicots. Monocots tend to have a more complex class C species, whereas dicots have more complex class A and B, and dicots had more complex Hsf subclass species than monocots. The AtHsf family exclusively consisted of AtHsfA7s (AT3G51910, AT3G63350), AtHsfA8 (AT1G67970) and AtHsfB3 (AT2G41690), whereas monocots only included HsfC2. These findings were discordant to the results of Scharf et al., which was caused by the use of a different reference for Hsf subclass nomenclature [11]. The name and sequence of AtHsfs, OsHsfs and ZmHsfs were downloaded from PlantTFBD which do not include subclasses A7 and A8 in monocots. Therefore, we acquired the present results of Hsfs classification, which did not possess
subclasses A7 and A8 in monocots. Our results are consistent with those of the classification and analysis of heat shock transcription factor family in maize which Hsfs classification were based on the data from PlantTFBD too [27].

The present study compared the Hsf family composition of three monocots, and all of them consisted of the same subclass species and proportion distribution, except for the novel subclass, TaHsfC3. Additionally, all of the same subclass genes from monocots were closely clustered. These results indicated that the subclass species and proportion distribution were relatively conserved in monocots, and each Hsf subclass may be indispensable for plants. We also found that the subclass HsfA2, HsfA6, HsfA9, HsfB1, HsfC1 genes from monocots and dicots were grouped into different clades, but the other subclass Hsf genes belonged to the same clades, which indicated that these same subclass Hsf genes from dicots and monocots were not closely related and thus may exist with relatively greater functional differences. A new clade with 13 TaHsfC members, hereby named TaHsfC3, did not have any orthologous genes in the model plant Hsf family. These new subclass genes might have different functions or roles in the stress signaling pathway in wheat.

Conserved structure analysis of TaHsfs
Similar to the Hsf families in other plants, the structure of the TaHsf proteins are well conserved. The DBD domain of plant Hsfs, which is characterized as a key domain, is encoded by two regions that are divided by an evolutionarily conserved intron, which was inserted immediately adjacent to the HTH DNA-binding motif [11]. This intron was detected in most DBD domains of TaHsfs, whereas no intron was found in TaHsfC3–1, TaHsfC3–2, TaHsfC3–3, TaHsfC3–4, TaHsfC3–6, TaHsfC3–7, TaHsfC3–8, TaHsfC3–9, TaHsfC3–10 and TaHsfC3–12 from the novel subclass TaHsfC3, as shown by their gene structure (Fig. 4). Previous study also shown that BnaHsf64 and BnaHsf64 without any introns inserted into the DBD domain [40]. Sequence alignment and MEME analysis identified six TaHsfs with an incomplete DBD domain (Figs. 1 and 5), whereas these genes were proven to be responsive to stress treatments (Fig. 6). These findings indicated that these TaHsfs with incomplete DBD domains can participate in plant stress responses.

Diverse transcription patterns of TaHsf family genes under different treatments
Plant Hsf genes not only could respond to abiotic stresses, but also to phytohormones. Genome-wide expression profiling of plant Hsf families under various abiotic stresses and phytohormones has been extensively studied in different tissues [43, 50, 51]. It was essential to investigate the expression profile of the TaHsf family under different stresses and phytohormones, before further studying a specific TaHsf gene. Our work revealed that the TaHsf family genes could respond to three stresses and two phytohormones, which suggests that TaHsfs might not only improve the heat tolerance of plants, but could also play a crucial role in increasing tolerance to various abiotic stresses as well as in enhancing signaling pathways. Organ-specific expression of TaHsfs was also observed between leaves and roots. Most of the TaHsf homoeologous groups had a similar expression pattern or mode under different treatments, for they shared a high level of sequence identities. The similar expression pattern was also found in the research on TaHsfC2a homoeologous group [32]. But the diverse expression level or pattern also been observed, such as TaHsfA2–1 is more sensitive to H2O2 than TaHsfA2–2 and TaHsfA2–3 in both leaf and root tissue; TaHsfA5–1, TaHsfA5–2 could be responsive to treatments in both leaf and root tissue, but TaHsfA5–3 showed no detectable expression under control and treatments. The duplicated Hsf genes were under strong purifying selection pressure [43], and the wheat genome often undergoes extensive genomic rearrangement, which may cause TaHsf functional differentiation or transcriptional silencing [53, 54]. The diverse expression pattern of homoeologous genes are also suggested to facilitate abiotic acclimation of wheat [55].

The TaHsfs in leaf tissues showed hypersensitivity to H2O2 treatment, particularly in terms of TaHsfAs and TaHsfBs, which exhibited the most significantly upregulated expression. It has been postulated that plant Hsfs act as sensors of ROS levels, resulting in Hsf activation and subsequent expression of other regulatory genes, including other Hsfs [56, 57]. H2O2 activates the Ca2+ signaling pathway at the cell surface, as well as entering the cell through PIP water channels and then activating Ca2+ signaling intracellularly [58]. Ca2+/calmodulin (CaM) could directly regulate the activation of Hsfs by phosphorylation of Hsfs [59]. This indicates that H2O2 may be the direct upstream regulatory component of Hsfs. TaHsfs were most insensitive to SA and PEG treatments in the leaf tissues. TaHsfAs were slightly upregulated, whereas the expression of other TaHsfs could be hardly detected under SA treatment. SA has been found to be involved in both basal and acquired thermotolerance in plants and activates various plant defense responses [60]. Similar results have been observed in tomato, HsfA1 could be induced by SA treatment, but not HsfA2. However, SA treatment could induce the expression of Hsp genes and increase the heat tolerance of plants by enhancing Hsf DNA-binding ability [12]. TaHsfs in leaf tissues were insensitive to PEG treatment, whereas these were relatively sensitive in the root tissues. These results have also been observed in the soybean Hsf
family under PEG treatment [51]. This is consistent with the fact that the leaves are the first to experience heat stress, whereas the roots are the first organs that perceive drought stress. In this study, 6 h of PEG treatment was not enough to lead to active drought stress responses in the leaf tissues. The expression profiles of TaHsfs revealed a specific responsive pattern to different treatments at the class level. Especially in the leaf tissues, TaHsfA1s exhibited sensitivity to H₂O₂ and HS treatment, and TaHsfC3s were sensitive to ABA treatment, whereas TaHsfBs were sensitive to H₂O₂, HS and ABA treatments. TaHsfCs were only inducible upregulated genes in the leaves and roots and were only down-regulated by ABA treatment, whereas TaHsfBs were sensitive to ABA treatment, whereas TaHsfCs were only down-regulated in subclass C3 were only induced by ABA treatment, whereas TaHsfD2s were only induced in subclass D2, which was also observed in other gene family studies [61, 62]. Therefore, we compared the expression patterns of orthologous Hsf genes between wheat and rice in response to different treatments, which revealed that both of these are most sensitive to HS and H₂O₂, as indicated by the upregulation of Hsfs in the seedling leaves. TaHsfs were more sensitive to H₂O₂, whereas OsHsfs were more sensitive to HS treatment. In terms of specific Hsf genes, OsHsfA2a (Os03g0745000), OsHsfA6a (Os06g0565200) and OsHsfB2a (Os04g0568700) were the most inducible upregulated genes in rice, TaHsfA2–10, TaHsfA2–12 as orthologs of OsHsfA2a and TaHsfA6–1, TaHsfA6–2, TaHsfA6–3 as orthologs of OsHsfA6a and TaHsfB2–1, TaHsfB2–2, TaHsfB2–3 as orthologs of OsHsfB2a were also significantly regulated with HS and H₂O₂ treatment. These results suggested that there were certain similarities in the stress response patterns between these two plants, and these Hsf genes may play an essential role in abiotic tolerance in plants. We also observed different expression patterns between rice and wheat orthologous Hsf genes such as TaHsfA2–13, TaHsfA2–14 and TaHsfA2–15, which were significantly upregulated under HS and H₂O₂ treatments, whereas the ortholog OsHsfA2c (Os10g28340) was insensitive to these treatments [50].

ABA is known to play important roles in regulating plant responses to various abiotic stresses, particularly those involving dehydration such as drought, salinity and cold stress [58]. ABA can improve tolerance to various abiotic stresses in plants through the regulation of Hsfs and Hsps. Our result revealed that the ABA-inducible TaHsfs mainly belong to classes B and C regardless of tissue. TaHsfC2–2, TaHsfC2–3 and TaHsfC2–4 were constitutively expressed in the leaves and roots and were only up-regulated by ABA treatment, whereas downregulated by other treatments. TaHsfs in subclass C3 were only induced by ABA treatment in the leaves. Subclasses TaHsfA3, TaHsfA4–1 and TaHsfA4–2 were only induced by ABA in the root tissues, but were hardly detected in the leaves. Huang et al., previously reported that AtHsfA6a (AT5G43840) and AtHsfA6b (AT3G22830) were induced by ABA, but not HS, and these genes are involved in the ABA signal pathway and ABA-mediated thermotolerance and drought tolerance [13]. These findings suggested that TaHsfs might also play a crucial role in the response and acclimation to drought stress in wheat, and ABA-induced TaHsfs seem to hold an independent responsive pathway that differs from heat and oxidative stress. The spatial expression of Hsfs has been investigated in several previous studies. OsHsfA6b (Os01g39020) and OsHsfA9 (Os03g12370) showed seed-specific expression in rice [63]. AtHsfA9 (AT5G54070) is exclusively expressed during the late stages of seed development [64]. GmHsf-02 is uniquely expressed in soybean roots [51]. In this work, we found that the expression pattern of TaHsfs significantly differed between leaves and roots. Several TaHsfs (TaHsfA2–10, TaHsfA2–12, TaHsfB2–6, TaHsfB2–7, TaHsfB2–8, TaHsfC2–2 and TaHsfC3–4) exhibit high basic expression levels in the roots, which in turn could hardly be detected in the leaves. Subclass TaHsfA6 was the most inducible TaHsf genes, which were upregulated by both H₂O₂ and heat treatment in the leaves, whereas it was only responsive to H₂O₂ treatment in the roots. These findings suggest that TaHsfA6s may function specifically in the oxidative stress signaling pathway, but not to osmotic stress. Subclass TaHsfB4 had a high basic expression level in root tissues, which dramatically decreased with all treatments, indicating that TaHsfB4s may be involved in root development.

Conclusions

A new Hsf protein HMM model constructed by the Hsf protein sequence of model monocot (O. sativa) and dicot (A. thaliana) plants was applied to identify novel Hsfs in the proteome of T. aestivum. A total of 82 non-redundant, full-length TaHsfs were identified and localized. Structural characteristics and phylogenetic analysis of T. aestivum, O. sativa, A. thaliana and Z. mays were performed to classify TaHsfs into three major classes and further into 13 subclasses. A novel subclass TaHsfC3 was identified in this study. The allopolyploid process, which occurred after the fusion of genomes, may have contributed to the expansion of the TaHsf genes. Furthermore, the TaHsf expression profiles by RNA-seq revealed that TaHsfs are not only responsive to abiotic stresses but also to specific phytohormones. Additionally, these TaHsfs exhibited class-, subclass- and organ-specific responsive patterns. The new multifunctional TaHsf characterized in this study improves our understanding of the response and acclimation of plants to multifactorial and combinational abiotic stresses, as
well as provides candidate gene resources for further investigations on abiotic stress tolerance in crops.

Methods

Data collection and identification of Hsf genes

The *T. aestivum* (bread wheat) genome sequence data of TGACv1 were obtained from the plant genome database [31]. The Hsf amino acid sequences and names of *A. thaliana*, *O. sativa* and *Z. mays* were downloaded from PlantTFBD [65]. The HMM model of Hsf was constructed using hmm3r3.0 based on the Hsf amino acid sequences of *A. thaliana* and *O. sativa* [66]. The HMM model of Hsf was used as query to search all possible Hsf protein sequences in the wheat proteome database using BLASTP (E < 0.001). The integrated DBD and HR-A/B domains in the putative wheat Hsfs were examined using SMART [67]. Candidate proteins without the HR-A/B or DBD domains were excluded from further analysis. The NLS and NES domains in the wheat Hsfs were predicted using cNSL Mapper and NetNES 1.1 [68, 69]. The prediction of AHA domains was based on the conserved AHA motif sequence (FWxxF/L, F/I/L) [70]. Protein isoelectric point (pI) and molecular weight (Mw) were calculated using ExPasy [71].

A total of 79 TaHsfs were mapped onto the 21 wheat chromosomes according to the information in the wheat database using MapGene2Chromosome [72]. Homologous gene groups were identified by three high-identity nucleotide sequence [36].

Multiple sequence alignment and phylogenetic analysis

The phylogenetic tree was constructed using the neighbor joining (NJ) method in MEGA (version 5.0) [73]. NJ analysis was conducted with the pairwise deletion option and the Poisson correction. For statistical reliability, bootstrap analysis was performed with 1000 replicates to assess statistical support for each node. For increasing readability of the phylogenetic tree, the N-terminal parts of the proteins containing the DBD, the HR-A/B and the linker between these two regions of TaHsfs were used in the analysis.

Orthologous gene identification and structure analysis

Orthologous gene pairs were identified based on (1) the best hit between *A. thaliana*, *O. sativa* and *T. aestivum*, (2) the position in the phylogenetic tree (bootstrap value > 50), and (3) the identity between orthologous gene pairs (> 90%). Intron-exon organization of the Hsf genes in wheat is illustrated using Gene Structure Display Server program [74] by alignment of the cDNAs with their corresponding genomic DNA sequences. The MEME program [75] was used for identification of conserved motifs, with the following parameters: the optimum motif widths: 6–50 amino acid residues and any number of repetitions: maximum number of motifs:15.

Plant materials and treatments

Cang 6005, the thermo-insensitive cultivated wheat provided by CangZhou Academy of Agriculture and Forestry Sciences, was used in this study. The selected seeds were surface-sterilized and repeatedly rinsed with tap water, then seeded in Hoagland nutrient solution after immersion and imbibition for 12 h, and cultivated in the incubator at 25 °C with a 16 h/8 h photoperiod/dark period. Six two-week-old homogeneous seedlings groups, each of which included 50 seedlings from five biological replicates, were subjected to different treatments, including 10 mM H2O2 for 90 min, heat (37 °C) for 60 min, 0.2 mM ABA for 12 h, 20% PEG6000 for 6 h, 0.8 mM SA for 1.5 h and control. The second leaf was sampled for the experiment. Pooled samples from each group were collected and immediately frozen in liquid nitrogen for RNA extraction.

RNA isolation and RNA-Seq analysis

Total RNA of each treatment was isolated from 50 seedlings from five biological replicates using the Total RNA Extractor (TRizol) kit (B511311, Sangon, China), and RNase-free DNase I was used to remove genomic DNA contamination. RNA integrity was estimated with an Agilent 2100 Bioanalyzer (Agilent Technologies, CA, USA). Approximately 2 μg of RNA from each sample was used as input material for RNA sample preparation. VAHTSTM mRNA-seq V2 Library Prep Kit for Illumina® was used to prepare the sequencing libraries. The HiSeq X Ten sequencers (Illumina, San Diego, CA, USA) were used in paired-end sequencing of the library. FastQC (version 0.11.2) was applied to evaluate the quality of sequenced data. Subsequently, the raw reads were screened using Trimmomatic (version 0.36). HISAT2 (version 2.0) was used to map the clean reads to the reference genome using default parameters. StringTie (version 1.3.3b) was used to calculate the gene expression abundance of the transcripts. TPM was used as transcript measurement, which eliminates the effect of gene sequencing discrepancies and the lengths enabled direct comparisons of gene expression among samples. Differentially expressed genes (DEGs) were identified by DESeq2 (version 1.12.4).

RNA isolation, cDNA synthesis & qRT-PCR analysis

To confirm the expression of representative of TaHsf genes, total RNA was isolated using Redzol (Beijing SBS Genetech Co.,Ltd.). The residual DNA was removed by DNase I (TaKaRa). For reverse transcription, the first-strand cDNA was synthesized using a PrimeScript™ first-strand cDNA synthesis kit (TaKaRa). Quantitative real-time PCR (qRT-PCR) for examination of the *TaHsfs* were
performed with the SYBR Premix ExTaq™ kit (TaKaRa) and an ABI 7500 (Applied Biosystem). Gene-specific and internal reference gene TaRP15 primers were listed in Additional file 7. The qRT-PCR program was set as the following: pre-denaturation at 95 °C for 30 s; denaturation at 95 °C for 5 s; annealing/extension at 60 °C for 34 s, 40 cycles. 2−ΔΔCt method was used to analyze the data. The expression level of TaHsf in leaf was set as 1. Three biological replicates were included for each group of experiments, and three technical replicates were included for each biological sample. The data were represented by mean value ± standard error of three biological replicates, as the previous described [76].

Additional files

Additional file 1: Table S1. Sequences and transcript IDs of novel TaHsf, (XLSX 73 kb)

Additional file 2: Table S2. Physical location of TaHsf. (XLSX 12 kb)

Additional file 3: Table S3. Orthologous gene pairs of Hsf family of *Triticum aestivum*, *Oryza sativa*, *Zea mays* and *Arabidopsis thaliana*. (XLS 25 kb)

Additional file 4: Table S4. Exon-intron structures of TaHsf. (XLSX 18 kb)

Additional file 5: Table S5. Motif sequences identified by MEME tools. (DOC 838 kb)

Additional file 6: Table S6. Gene-specific and internal reference gene TaRP15 primers used for qRT-PCR. (XLSX 10 kb)

Abbreviations

ABA: Abscisic Acid; AHA: Activator motif; DBD: DNA Binding Domain; HMM: Hidden Markov Model; HR: Heptad Repeats; HSF: Heat shock factor; Hsp: Heat shock protein; HTH: helix-turn-helix; NES: Nuclear export signal; OD: Oligomerization Domain; PEG: Polyethylene Glycol; qRT-PCR: quantitative RT-PCR; ROS: Reactive Oxygen Species; SA: Salicylic Acid; TPM: Transcripts Per Million

Acknowledgements

We thank Dr. Amy Flanagan (Noble Research Institute, Oklahoma, USA) for her linguistic assistance during the revision of this manuscript.

Funding

This work was supported by the key project of Natural Science Foundation of Hebei Province (C2016301085), the Science and Technology Innovation Program for Modern Agriculture of Hebei Province (494–0402–JBN–S2XB), the Basic Operating Foundation of Hebei Academy of Agriculture and Forestry Sciences (201810102).

Availability of data and materials

All data generated or analysed during this study are included in this published article (and its supplementary information files).

Authors’ contributions

This study was designed by SND, XLG and GLL. SND performed bioinformatics analysis, generated and analyzed data, and wrote the manuscript. BHL and YYZ performed the experiments. XLG and GLL edited this manuscript and contributed with valuable discussions. All authors read and approved the final manuscript.

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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Received: 20 November 2018 Accepted: 18 March 2019

Published online: 01 April 2019

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