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Genome-wide identification of HSF family in peach and functional analysis of *PpHSF5* involvement in root and aerial organ development

Bin Tan^{*}, Liu Yan^{*}, Huannan Li, Xiaodong Lian, Jun Cheng, Wei Wang, Xianbo Zheng, Xiaobei Wang, Jidong Li, Xia Ye, Langlang Zhang, Zhiqian Li and Jiancan Feng

College of Horticulture, Henan Agricultural University, Zhengzhou, China Henan Key Laboratory of Fruit and Cucurbit Biology, Zhengzhou, China These authors contributed equally to this work.

ABSTRACT

Background. Heat shock factors (HSFs) play important roles during normal plant growth and development and when plants respond to diverse stressors. Although most studies have focused on the involvement of HSFs in the response to abiotic stresses, especially in model plants, there is little research on their participation in plant growth and development or on the HSF (PpHSF) gene family in peach (*Prunus persica*).

Methods. DBD (PF00447), the HSF characteristic domain, was used to search the peach genome and identify *PpHSFs*. Phylogenetic, multiple alignment and motif analyses were conducted using MEGA 6.0, ClustalW and MEME, respectively. The function of *PpHSF5* was confirmed by overexpression of *PpHSF5* into Arabidopsis.

Results. Eighteen *PpHSF* genes were identified within the peach genome. The *PpHSF* genes were nonuniformly distributed on the peach chromosomes. Seventeen of the PpHSFs (94.4%) contained one or two introns, except PpHSF18, which contained three introns. The in silico-translated PpHSFs were classified into three classes (PpHSFA, PpHSFB and PpHSFC) based on multiple alignment, motif analysis and phylogenetic comparison with HSFs from Arabidopsis thaliana and Oryza sativa. Dispersed gene duplication (DSD at 67%) mainly contributed to HSF gene family expansion in peach. Promoter analysis showed that the most common cis-elements were the MYB (abiotic stress response), ABRE (ABA-responsive) and MYC (dehydration-responsive) elements. Transcript profiling of 18 PpHSFs showed that the expression trend of PpHSF5 was consistent with shoot length changes in the cultivar 'Zhongyoutao 14'. Further analysis of the PpHSF5 was conducted in 5-year-old peach trees, Nicotiana benthamiana and Arabidopsis thaliana, respectively. Tissue-specific expression analysis showed that *PpHSF5* was expressed predominantly in young vegetative organs (leaf and apex). Subcellular localization revealed that PpHSF5 was located in the nucleus in N. benthamiana cells. Two transgenic Arabidopsis lines were obtained that overexpressed *PpHSF5*. The root length and the number of lateral roots in the transgenic seedlings were significantly less than in WT seedlings and after cultivation for three weeks. The transgenic rosettes were smaller than those of the WT at 2–3 weeks. The two transgenic lines exhibited a dwarf phenotype three weeks after transplanting, although there was no significant difference in the number of internodes. Moreover, the PpHSF5-OE lines

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Corresponding author Jiancan Feng, jcfeng@henau.edu.cn

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exhibited enhanced thermotolerance. These results indicated that PpHSF5 might be act as a suppresser of growth and development of root and aerial organs.

Subjects Agricultural Science, Biotechnology, Genomics, Molecular Biology, Plant Science **Keywords** Heat shock factors family, Root development, Peach (*Prunus persica*), Functional identification, PpHSF5, Aerial organ

INTRODUCTION

Plant growth and development are affected by a range of abiotic stress, including cold, heat, salinity and drought stress (*Guo et al.*, 2016). Heat shock factors (HSFs) act with heat shock proteins (HSPs) as key transcriptional activators during responses to abiotic stress (*Hu*, *Hu* & *Han*, 2009). Recent studies indicated that HSFs act as key components of signal transduction in response to different abiotic stresses in plants (*Guo et al.*, 2016; *Scharf et al.*, 2012).

HSFs in plant genomes can be identified by a conserved DNA-binding domain (DBD). The DBD domain is located in the N-terminal of all HSFs and specifically binds to heat stress (HS) motifs in the promoters of target genes (*Wang et al., 2018*). The adjacent HR-A/B region is linked to the DBD by a connector of variable length (15–80 amino acid residues) that contains a bipartite heptad pattern of hydrophobic amino acid residues, which constitutes a coiled-coil domain for protein interaction. According to the number of amino acid residues inserted into the HR-A/B region, HSFs are divided into three main groups, each with subgroups, namely HSFA (A1-A9), HSFB (B1-B5) and HSFC (C1-C2) (*Koskull-Doring, Scharf & Nover, 2007; Yang et al., 2014*). HSFA members contain an acidic motif (AHA activation domain) at their C-terminus and act as transcriptional activators. The members of HSFB act as transcriptional repressors.

In a wide range of plants, a number of HSFs have been shown to be involved in resistance to heat (*Guo et al., 2016*) and other abiotic or biotic stresses (*Yu et al., 2019*). Of the 21 HSF family members in Arabidopsis, a number act as pioneer regulators of the response to heat shock. HSFA1a, HSFA1b, HSFA1d, HSFA1e and HSFA2 play active regulatory roles in the response to HS in plants (*Busch, Wunderlich & Schoffl, 2005*; *Nishizawa et al., 2006*). In Arabidopsis, the assembly of the HSFA1/A2 super-activated complex regulates heat stress genes (*Chan-Schaminet et al., 2009*). *HSFB1* and *HSFB2b* participate in disease resistance regulation of Arabidopsis and expression of *Pdf1.2* (*Kumar et al., 2009*). *OsHSFB4b* and *OsHSFA2c* participate in the regulation of the heat shock response by regulating the expression of HSP100 (*Singh et al., 2012*). *OsHSFC1b* is related to the regulation of salt stress and plant development (*Schmidt et al., 2012*).

Several HSFs are stress-inducible transcriptional factors that participate in the growth and development of root and aerial organs in plant. Overexpression of *AtHsfB4* in Arabidopsis induces specific effects on root development, resulting in shortened roots (*Begum, Reuter & Schoffl, 2013*). The over-expression of *BhHsf1* conferred growth retardation of aerial organs, producing a dwarf phenotype, although the primary roots were not obviously different from those of wild type (*Zhu et al., 2009*). Transgenic Arabidopsis

plants with strong expression of *AtHsfA3* and *AtHsfA2* showed a severely dwarfed phenotype and increased tolerance to heat (*Ogawa, Yamaguchi & Nishiuchi, 2007*; Yoshida et al., 2008). The thermotolerant phenotype was also observed in the cotyledons, rosette leaves, inflorescence stems and seeds of transgenic Arabidopsis plants expressing *OsHsfA2e* (*Yokotani et al., 2007*).

The HSF family have been analyzed genome-wide in several plants, such as rice (Oryza sativa), Arabidopsis (Arabidopsis thaliana), cotton (Gossypium hirsutum), soybean (Glycine max), wheat (Triticum aestivum), pepper (Capsicum annuum), poplar (Populus trichocarpa), Brassica napus, grape (Vitis vinifera) and Tartary buckwheat (Fagopyrum tataricum) (Nover et al., 2001; Chauhan et al., 2011; Wang et al., 2014; Li et al., 2014; Guo et al., 2015; Xue et al., 2014; Zhang et al., 2016; Zhu et al., 2017; Liu et al., 2018; Liu et al., 2019). Peach (Prunus persica L.) is an important economical crop and a popular fruit with consumers. However, there are limited studies on peach HSFs. To remedy this, we identified the HSF family in the peach genome and conducted bioinformatics analysis of the 18 identified *PpHSFs*. Based on the latest transcriptome data (*Lian et al., 2020*), the expression patterns of the PpHSF genes were analyzed during development of the cultivar 'Zhongyoutao 14'. 'Zhongyoutao 14' (derived from 'SD9238'), is a temperature-sensitive peach that exhibits a shorter internode length and a smaller canopy when grown below 30 °C (Lu et al., 2016). PpHSF5 was further analyzed and found to function in the development of the root and aerial organs. Furthermore, the thermotolerant phenotype was analyzed in newly obtained transgenic Arabidopsis plants expressing *PpHSF5*. The genome-wide analysis of *PpHSF* gene family offers a basis for further investigation into the function and evolutionary history of peach HSFs and provides candidate genes for peach molecular breeding.

MATERIALS & METHODS

Plant materials

Established peach trees (*Prunus persica*) cultivar 'Zhongyoutao 14' ('Maotao' as rootstock) have been grown for 5 years at the Experimental Station of the Horticulture College, Henan Agricultural University (Zhengzhou, China). Samples from the apex, young and mature leaves, self-pollinated embryos, and fruit were collected, frozen in liquid nitrogen and stored at -80 °C. Leaves from *Nicotiana benthamiana* were used for subcellular location of PpHSF5. *Arabidopsis thaliana* (L.) Heynh Columbia 0 (Col-0) was used for transformation with *PpHSF5*.

Identification and chromosomal location of HSF genes in peach

The hidden Markov model (HMM) of the DBD domain (PF00447), characteristic of HSFs, was downloaded from the Pfam website (*Finn et al., 2000*) and used to identify HSF genes in peach. The peach genome files (v2.1) were downloaded from JGI database (https://phytozome.jgi.doe.gov/pz/portal.html) (*Verde et al., 2013*), HSF protein sequences were obtained in peach genome by BLASTP and hmmsearch function, and then the DBD domain were further identified by Pfam analysis. The peach HSF gene and protein sequences were named

according to physical location on the chromosomes. Positional information was retrieved from peach genome annotations obtained from Phytozome v12.1, and chromosome locations of the *PpHSFs* were drawn using the Circos software (*Krzywinski et al., 2009*). The isoelectric points and other physical properties were approximated from ExPASy (http://web.expasy.org/compute_pi). Gene structures were predicted using the Gene Structure Display Server 2.0 (http://gsds.cbi.pku.edu.cn/).

Phylogenetic and motif analysis of PpHSFs

The amino acid sequences of 21 AtHSFs (*Arabidopsis thaliana*), 25 OsHSFs (*Oryza sativa*) and 18 PpHSFs (*Prunus persica*) were gathered from Phytozome v12.1 using ClustalW with system default settings. The phylogenetic trees were formulated by the maximum likelihood method (ML) with Jones-Taylor-Thornton (JTT) model in MEGA 6.0 (http: //www.megasoftware.net/download_form).. Conserved motifs of HSF proteins in peach were identified using the MEME tool (http://meme.nbcr.net/meme/cgi-bin/meme.cgi) with default parameters in normal operation mode. The subcellular localization was predicted with Plant-mPLoc (http://www.csbio.sjtu.edu.cn/bioinf/plant-multi/#).

Gene duplication and Cis -element analysis of PpHSFs

Gene duplication was analyzed using MCScanX (*Wang et al., 2012*). Genomic DNA sequences (2000 bps upstream of the start codons) for each *PpHSF* were obtained from the peach genome and skimmed in the PlantCARE database (http://bioinformatics.psb.ugent. be/webtools/plantcare/html/) for *cis*-acting elements analysis of the promoter in *PpHSFs*.

Gene expression analysis of PpHSFs

The FPKM (fragments per kilobase of exon per million fragments mapped) values of the 18 *PpHSFs* (Table S1-1) were obtained from our previous study of gene expression in shoots at four critical growth stages, namely initial period (IP), initial elongation period (IEP), rapid growth period (RGP) and stable growth period (SGP) of temperature-sensitive peach cultivar 'Zhongyoutao 14' (*Lian et al., 2020*). The average maximum temperature of previous week (AMTPW) began to be higher than 30 °C in the first day of RGP (*Lian et al., 2020*). The heat map was generated by TBtools (*Chen et al., 2020*).

Quantitative real-time PCR analysis of PpHSF5

Total RNA of different tissues from 'Zhongyoutao 14' peach and leaves from T_2 transgenic Arabidopsis lines was isolated using the Spin Column Plant Total RNA Purification Kit (ShengGong, Shanghai, China). The cDNA was synthesized using FastQuant RT Kit (Tiangen Biotech, Beijing, China). qRT-PCR was implemented using an ABI PRISM 7500 FAST Sequence Detection System (Applied Biosystems, Madrid, CA, USA) with SYBR Select Master Mix (Applied Biosystems, USA). Primers of *PpHSF5* were designed using Primer Premier 5.0. *PpGAPDH* (Prupe.1G234000) and *AtUBC* (AT5G25760) were used as constitutive controls for either tissue-specific expression in peach or expression analysis in transgenic Arabidopsis, respectively. Primers are shown in Table S1-2. The reaction mixture was as follows: 1 μ L cDNA template (200 ng/ μ L), 1 μ L of each primer (10 μ M), 10 μ L SYBR Premix and 7 μ L ddH₂O. Melting curve analysis was performed after the end of 40 cycles to insure proper amplification of the target. During the melting process, fluorescence readings were continuously collected from 60–90 °C at a heating rate of 0.5 °C s⁻¹. All analysis was repeated three times using biologically replicates. The relative expression levels of *PpHSF5* were calculated as $2^{-\Delta\Delta CT}$ method (*Schmittgen & Livak, 2008*). The relative expression levels of *PpHSF5* was calculated in SPSS using ANOVA at significance levels of *P* < 0.05.

Subcellular localization of PpHSF5

PpHSF5 without the termination codon was amplified by PCR using cDNA from 'Zhongyoutao 14' as the template (Primer details in Tables S1–S2). This coding region was cloned into the pSAK277-GFP vector to construct PpHSF5::GFP fusion proteins that were driven by the CaMV 35S promoter. The p35SPpHSF5::GFP and p35SGFP (control) vectors were transformed into *Agrobacterium tumefaciens* strain GV3101, which were then injected into leaves of *N. benthamiana* according to previously published protocols (*Sparkes et al., 2006*). The leaves were observed 48–72 h after injection using laser scanning confocal microscopy (Zeiss LSM700).

Construction of expression vectors for plant transformation

The CDS of *PpHSF5* was PCR-amplified and cloned into the pSAK277 vector using the restriction enzymes *Xho* I and *Xba* I (Primer details in Table S1–S2). The *p35S::PpHSF5* vector was transformed into *Agrobacterium tumefaciens* strain GV3101. The floral dip method was used to transform *Arabidopsis thaliana* (Col-0) (*Chung, Chen & Pan, 2000*).

Phenotype of overexpression PpHSF5 in Arabidopsis

The seeds from T_2 transgenic Arabidopsis lines were sterilized by 6.25% NaClO for 5 min, and then washed in sterilized ddH₂O. The seeds were cultured at 4 °C for 2 d and then transferred onto MS solid medium under 16/8 h light/dark cycle for one week on square plates. Three biological replicates (with three seedlings of each lines per square plate) were used for observation of root phenotype. The roots of different transgenic lines with three plants per line were measured by a LA2400 Scanner at three weeks to determine the growth status. The leaves were cut from the transgenic seedlings for gene expression analysis. Other seedlings, germinated on agar and grown for one week, were transferred into the soil and grown under normal conditions. The length and width of rosettes (four leaves per plants, five repetition) and number of rosettes (five plants per line) in different transgenic lines and WT were measured and photographed at two weeks and at three weeks after transplanting, respectively. Moreover, the morphology of transgenic lines and WT, including the height of plants (eight plants per line) and the number of branches and blooms (five plants per line) were recorded, three weeks after transplanting.

Heat stress treatment

For performing heat stress treatment on the seeds germination and plants grown on the agar medium, seeds of WT and transgenic Arabidopsis lines sown on MS medium at 4 °C for 2 d and in darkness for 2 d (22 °C) were exposed to HS stress at 46 °C for 30 min, and then were transferred into a climate chamber (22 °C, 16 h light/8 h dark cycles). After HS

Table 1 Basic information of <i>PpHSF</i> gene family members.										
Gene name	Gene ID	Length of CDS (bp)	No. of amino acids (aa)	Molecular weight (Da)	Predicted isoelectric point (PI)	Chromosome location	Subcellular localization			
PpHSF1	Prupe.1G021200	1068	355	41166.82	4.9	1	Nucleus			
PpHSF2	Prupe.1G165500	1452	483	54128.99	5.61	1	Nucleus			
PpHSF3	Prupe.1G335100	1227	408	46017.53	4.81	1	Nucleus			
PpHSF4	Prupe.1G410400	1125	374	41952.84	4.95	1	Nucleus			
PpHSF5	Prupe.1G433500	1170	389	43557.76	7.17	1	Nucleus			
PpHSF6	Prupe.2G292100	912	303	33900.98	5.19	2	Nucleus			
PpHSF7	Prupe.3G108700	1314	437	49855.78	5.13	3	Nucleus			
PpHSF8	Prupe.4G046000	591	196	22364.46	8.75	4	Nucleus			
PpHSF9	Prupe.4G068100	1224	407	46118.31	5.23	4	Nucleus			
PpHSF10	Prupe.4G144200	1512	503	56052.61	4.78	4	Nucleus			
PpHSF11	Prupe.5G031100	1551	516	56261.49	4.67	5	Nucleus			
PpHSF12	Prupe.5G093200	996	331	36068.73	4.75	5	Nucleus			
PpHSF13	Prupe.7G056700	735	244	28001.34	5.75	7	Nucleus			
PpHSF14	Prupe.7G117200	1608	535	59567.89	4.98	7	Nucleus			
PpHSF15	Prupe.7G133600	900	299	33339.83	5.07	7	Nucleus			
PpHSF16	Prupe.7G206900	1458	485	53632.01	5.07	7	Nucleus			
PpHSF17	Prupe.7G231100	1002	333	37851.52	5.68	7	Nucleus			
PpHSF18	Prupe.8G234900	1080	359	40936.56	5.58	8	Nucleus			

treatment, the germination of seeds were counted daily and photographed. More than 50 seeds of each line were used in each plate with three replications. Difference in HS stress was confirmed using t-test.

Statistical analysis

Data were analyzed by ANOVA, Tukey HSD^a and Duncan^a's multiple range tests (at P < 0.05) using IBM SPSS Statistics 20 (SPSS, USA).

RESULTS

Genome-wide identification, chromosomal distribution and gene structures of HSF genes in peach

Eighteen *HSF* gene family members were identified from the peach genome and then named *PpHSF1* to *PpHSF18* according to their physical locations (Table 1 and Table S2-1). The *PpHSF* coding sequences ranged from 591 bp (*PpHSF8*) to 1608 bp (*PpHSF14*). In silico-translated PpHSF proteins showed divergent lengths [196 to 535 amino acids (aa)] with different molecular weights (22.36~59.56 kDa) and isoelectric points (4.67 to 8.75) (Table1). All PpHSFs were predicted to be nuclear-localized proteins.

Seven of the 8 peach chromosomes contained at least one *PpHSF*, with the exception being chromosome 6 (Fig. 1 and Table 1). Five *PpHSFs* were located on chromosome 1 (*PpHSF1-5*), and another five (*PpHSF13-17*) on chromosome 7. Chromosomes 2, 3 and 8 carried only one *PpHSF* gene each, while chromosome 5 had two, and chromosome 4 had



Figure 1 Chromosomal location of *HSF* genes in peach (*PpHSFs*). Three syntenic pairs are linked by red lines.

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three. The above results indicated that *PpHSFs* were unevenly distributed across the peach chromosomes.

The structural differences of the *PpHSF* genes were also analyzed. The number of introns ranged from one to three among the *PpHSFs*. The majority of the *PpHSFs* (66.67%) contained one intron, 27.78% contained two introns, and only *PpHSF18* contained three introns (Fig. S1 and Table S2). Interestingly, both *PpHSF18* and *PpHSF12* has predicted introns in the 5'-UTR and 3'-UTR, respectively.

Gene duplication pattern analysis of PpHSFs

To explain the expansion of the *PpHSFs* gene family, the gene duplication patterns of the *PpHSFs* were analyzed and compared across the peach genome (Table S3). There were only two patterns of gene duplications, with 67% of the gene pairs derived from dispersed gene duplication (DSD) and the remaining gene pairs derived from whole-genome duplication (WGD). Three syntenic pairs were identified, and all originated from WGD. The syntenic genes were located on different chromosomes from their partner (Fig. 1).

Classification, phylogenetic and motif analyses of PpHSFs

Among plant species, there are two characteristic amino acid domains in the HSF family, the DBD and adjacent HR-A/B region (*Nover et al., 2001*). The PpHSFs were divided into three classes (PpHSFA, PpHSFB and PpHSFC), according to the number of amino acids

between part A and part B of the HR-A/B domain (Fig. 2A). Multiple sequence alignment analysis of the PpHSF proteins indicated that an insertion of 21 amino acids was found in Class A (11 of the PpHSFs) and a shorter insertion of 14 amino acid in Class C (1 of the PpHSFs) between the HR-A and HR-B regions. Six of the PpHSFs had no aa insertion between the two domains (Class B).

Phylogenetic analysis among the HSF proteins from three plant species, namely 21 AtHSFs (*Arabidopsis thaliana*), 25 OsHSFs (*Oryza sativa*) and 18 PpHSFs (*Prunus persica*), was conducted by constructing a phylogenetic tree. According to the phylogenetic tree, the 64 HSFs derived from the three plant species were divided into three classes and 15 subclasses (Fig. 2B). The peach proteins sorted into the classes of HSFs, within (11 members) in class HSFA, six in HSFB, and one in HSFC. Class A included nine subclasses (A1-A9), the largest number of subclasses. The PpHSFs were grouped into eight of the Class A subclasses, with no PpHSF in Class A7. Class B consisted of 18 total members and was divided into four subclasses (B1-B4). It is noteworthy that PpHSF8 clustered with Class B but as a single branch. Only six members were clustered into Class C, with two subclasses (C1-C2). No PpHSFs clustered with subclass C2.

The conserved motifs in the PpHSF proteins were analyzed using MEME. The results revealed that PpHSFs contained ten conserved motifs (Fig. 2C and Table S4). Motifs 1-3 were found in the N-terminals (the most conserved region) of each PpHSF. Motif 4 was found in Class A and Class B. Motif 5, which was found between the HR-A and HR-B regions, was observed in Class A and Class C. The motif analysis was consistent with the multiple sequence alignment and phylogenetic analyses.

Analysis of the *Cis*-acting regulatory elements in the *PpHSF* gene promoters

The cis-acting elements within the promoters of the 18 *PpHSFs* were analyzed using PlantCARE. Every promoter contained at least two MYB elements (abiotic stress response) (Table 2). All but one promoter contained an ABRE (ABA-responsive element). The next most common elements were MYC elements (dehydration-responsive) (in 88.8% of the promoters), CGTCA- and TGACG- motifs (83.3%), and ARE elements (anaerobic induction) (77.8%). ERE (ethylene-responsive element), MBS (drought inducible), MRE and P-box elements were also present in the promoters of some *PpHSFs*. The TCA-motif was observed in only five *PpHSFs*, namely *PpHSF1*, *PpHSF2*, *PpHSF5*, *PpHSF6* and *PpHSF13*. Previous studies reported several elements, including MYB, ABRE, MYC, play vital roles in stress responses in plants (*He et al.*, 2012; *Li et al.*, 2012). The different *cis-elements* in the promoter regions of these *PpHSFs* implied that the *PpHSFs* may function in plant development and stress responses.

The expression patterns of *PpHSFs* during shoot elongation in 'Zhongyoutao 14'

Based on our previous RNA-seq analysis (*Lian et al., 2020*), the expression patterns of *PpHSFs* were compared in four critical stages of shoot elongation of 'Zhongyoutao 14' grown under elevated temperature in the field (Fig. 3). Most of *PpHSFs* belonging to the A and C classes (except *PpHSF4* and *PpHSF11*) were maintained at lower expression level.



Figure 2 Multiple sequence alignment of the HR-A/B regions (OD), conserved motif and phylogenetic analysis of PpHSFs. (A) Multiple sequence alignment of the HR-A/B regions, from the start of the DNA-binding domain to the end of the HR-A/B region, of the HSF proteins were aligned with MEGA 6. (B) Hylogenetic tree of HSFs from *Prunus persica* (Pp, red star), *Oryza sativa* (Os, blue circle) and *Arabidopsis thaliana* (At, green square) constructed by maximum likelihood method (ML) with Jones-Taylor-Thornton (JTT) model in MEGA 6.0. Both locus ID and subclass numbers are listed. (C) Analysis of conserved motifs in the HSF gene family in peach. Proteins are organized according to the groups in Fig. 2A. Ten motifs were found in the protein sequences as shown in Table S4.

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The *PpHSFs* of B class exhibit diverse expression patterns. The FPKM values of *PpHSF8* and *PpHSF13* remained almost unchanged at the four stages. The transcripts of another three *PpHSFs* (*PpHSF15*, *PpHSF6* and *PpHSF12*) were present at lower levels during the IEP stage and then slightly increased during the RGP and SGP stages. The expression of level of *PpHSF5* showed higher in IEP stage and increased from the RGP to SGP stages. *PpHSF5* might participate in temperature-induced shoot growth of temperature-sensitive peach.

Expression analysis of *PpHSF5* and subcellular localization of PpHSF5

The relative expression of *PpHSF5* was investigated by qRT-PCR in different organs of 'Zhongyoutao 14' (Fig. 4; Table S5-1). The results showed that *PpHSF5* were expressed predominantly in young vegetative organs (leaves and apex), but barely detectable in embryos and mature leaves. This suggested that *PpHSF5* might participated in the growth and development of plants. The 35S::PpHSF5-GFP signal was evident in the cellular nucleus in *N. benthamiana* cells, indicating a nuclear localization (Fig. 5). The result was in concurrence with the prediction from Plant-mPLoc of subcellular localization (Table 1).

Overexpression of *PpHSF5* in arabidopsis results in dwarf phenotypes

To investigate the function of *PpHSF5*, an overexpression vector with *PpHSF5* was transformed into Arabidopsis. The phenotype of two transgenic lines and WT were recorded (Fig. 6). One week after germination on agar medium, the transgenic lines had shorter roots and a smaller number of lateral roots than WT seedlings (Fig. 6A). The average root length in WT was 7.13 cm, in transgenic line L1 was 3.08 cm, and in L2 was 3.50 cm (Figs. 6A and 6B; Table S6-1). Two weeks after transplantation, there was no difference in the number of rosette leaves between the transgenic lines and WT (Fig. 6C and 6D–6A and Table S6-2), although the rosette leaves were significantly shorter and narrower in the transgenic lines (the average length and width; Figs. 6D–6B, 6C and Tables S6-3 and S6-4). The mRNA levels in the PpHSF5-OE lines were obviously higher than WT plants (Fig. 6D and Table S5-2).

Three weeks after transplanting, the soil-grown transgenic lines had fewer rosette leaves and the leaves were shorter and narrower than those in WT plants (Fig. 6F). Moreover, the two transgenic lines (L1 and L2) exhibited a dwarf phenotype (Figs. 6E and 6F). The average height of L1 (16.83 cm) was 40% shorter than that of the WT (26.77 cm). The number of rosette branches was much greater in WT than in transgenic lines, which had just one flowering stalk (Fig. 6F). There was no significant difference in the number of internodes (Figs. 6E and 6F), indicating that the dwarf phenotype of the transgenic lines might be caused by shorter internode length.

Shorter roots were also observed in the transgenic lines for cultivation three weeks after transplanting (Figs. 6G and 6H). Root length and root volume were significantly lower in transgenic lines compared to WT (Figs. 6G and 6H, Tables S6-12, S6-13. The average length of roots in Line 1 was 219.34 cm, which was 54% of the length in WT plants. The root

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Table 2 Cis-elements in the promoters of eighteen PpHSF genes.											
	ABRE	ARE	CGTCA-motif	ERE	MBS	MRE	MYB	MYC	P-box	TGACG-motif	TCA-element
PpHSF1	1	3	2	2	1	1	4	3	2	2	2
PpHSF2	1	2	2	1	-	-	3	1	-	2	1
PpHSF3	1	1	_	2	1	1	4	2	_	1	-
PpHSF4	3	6	3	-	-	-	3	-	1	3	_
PpHSF5	5	_	3	-	2	_	7	_	1	3	3
PpHSF6	4	_	3	_	1	_	7	5	2	3	1
PpHSF7	1	5	_	1	_	_	6	3	_	_	_
PpHSF8	3	3	3	1	1	1	3	8	1	3	_
PpHSF9	5	5	4	-	-	1	13	4	-	4	_
PpHSF10	3	1	5	2	-	-	2	6	-	5	_
PpHSF11	_	1	1	_	2	_	7	8	_	1	_
PpHSF12	6	_	1	1	1	1	13	3	_	1	_
PpHSF13	3	4	1	_	_	_	4	5	_	1	1
PpHSF14	3	4	3	-	3	_	11	3	_	3	_
PpHSF15	8	2	4	1	1	1	6	5	_	4	_

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Figure 3 Heatmap of transcript levels of HSF genes in peach. Transcriptome data were used to measure the expression level of *PpHSFs*. The gene names on the right are organized according to the different subclasses. Samples were harvest from shoots at the IP (initial period), IEP (initial elongation period), RGP (rapid growth period), and SGP (stable growth period), which are four key growth stages during temperature-sensitive peach shoot development. Color scale at the top represents FPKM values. Blue indicates low expression and red indicates high expression. Heatmap was generated using TBtools. Full-size DOI: 10.7717/peerj.10961/fig-3



Figure 4 Relative expression of *PpHSF5* in different tissues of 'Zhongyoutao 14' peach. Established plants were grown under normal conditions. The analyzed tissues include the apex, flower, embryo, young leaf, and mature leaf, which harvested at the same time. The relative expression levels were calculated using the $2^{-\Delta\Delta CT}$ method.

Full-size DOI: 10.7717/peerj.10961/fig-4



Figure 5 Subcellular localization of PpHSF5 in *N. benthamiana* epidermal cells. (A and D) Images of green fluorescence from the GFP protein and the PpHSF5-GFP fusion protein in tobacco cells under the confocal microscope; (B and E) Bright field image of tobacco epidermal cells; (C) Overlay of A and B; (F) Overlay of D and E.

Full-size DOI: 10.7717/peerj.10961/fig-5

volumes in the transgenic lines (Line 1 was 0.19 cm³, Line 2 was 0.36 cm³) was no more than 20% of that in WT (1.95 cm³). Other root indexes output by the root scanner were also less in the PpHSF5-OE lines, including the forks, tips and crossings of roots (Figs. 6G and 6H, Tables S6-14, S6-15, S6-16). Between the two transgenic lines, the higher expression level of *PpHSF5* in L1 resulted in more obvious phenotypes compared to PpHSF5-OE L2 and WT (Figs. 6A, 6C, and 6E and Table S5-2). The above results indicated that *PpHSF5* might participate in plant growth and development and that overexpression of *PpHSF5* results in a dwarf phenotype in transgenic Arabidopsis.

PpHSF5-OE lines exhibit enhanced thermotolerance

The thermotolerance of PpHSF5-OE lines was assayed with that of WT (Fig. 7 and Table S7). As shown in Fig. 7B and 7F, only 8.3% WT seeds germinated, whereas more than 93.3% of the transgenic seeds germinated after HS treatment 1 d. Nearly half of the WT seeds germinated after HS treatment 3 d, whereas 100% of the transgenic seeds were germinated (Fig. 7C and 7F). After HS treatment 5 d and 7 d, 68.4% and 82.6% of WT seeds germinated, respectively (Figs. 7D, 7E and 7F). Compared to WT seedlings, the PpHSF5-OE seedlings exhibited green cotyledons and vigor growth (Figs. 7C, 7D and 7E). These results suggested that the overexpression of *PpHSF5* improves thermotolerance of PpHSF5-OE lines.

DISCUSSION

Peach contains fewer HSF gene family members among several plant species

HSFs play vital roles in plant growth and defense. Through plant genome sequencing, *HSF* gene family members have been identified in several model organisms and more than 20 plant species (Table S8). Only a single *HSF* was detected in yeast, nematodes and flies (*Nakai, 1999; Nover, 1996*). In this study, 18 *HSF* genes were identified in peach, which is



Figure 6 Phenotypic and expression analysis of transgenic Arabidopsis over-expressing *PpHSF5*. (A) Phenotype of T_2 transgenic plants from two lines over-expressing *PpHSF5* after cultivation for one week. (B) Root length of T_2 transgenic plants over-expressing *PpHSF5*. Three plants were measured in each biological replicate. (C) Phenotype of T_2 transgenic plants from two lines over-expressing *PpHSF5* after cultivation for two weeks. Seeds were transferred to soil after (continued on next page...)

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Figure 6 (... continued)

germination and growth on agar for five days. (D) The morphology and relative expression of T_2 transgenic plants with PpHSF5 and WT after cultivation for two weeks. The number of rosettes, length and width after cultivation for two weeks. Relative expression of *PpHSF5* in transgenic Arabidopsis plants carrying *p35S:PpHSF5*; (E) Phenotype of T_2 transgenic plants over-expressing *PpHSF5* after cultivation in soil for three weeks. (F) The length, width and number of rosette leaves, number of internodes and flower stalks, and the height of plants after cultivation for three weeks. (G) Phenotype of T_2 transgenic plant roots over-expressing *PpHSF5* after cultivation for three weeks. (H) The root length, volume and other indexes were scanned after cultivation for three weeks.



Figure 7 Thermotolerance of the *p35S:: PpHSF5* plants. (A) Five-day-old seedlings of wild type and the *p35S:: PpHSF5* plants were treated at 46 °C for 30 min. Photographs were taken before HS treatment. (B) Photographs were taken after 1d in 22 °C. (C) Photographs were taken after 3d in 22 °C. (D) Photographs were taken after 5d in 22 °C. (E) Photographs were taken after 7d in 22 °C. (F) Comparison of germination rate among wild-type, *p35S::PpHSF5* transgenetic plants after HS treatment. The number of germinated plants was counted daily after HS treatment. For three replication, more than 50 seedlings were used each lines (*t*-test signifificant at *P* < 0.05 and *P* < 0.01, respectively).

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less than in most other plant species, but more than in tea (*Camellia sinensis*), strawberry (*Fragaria vesca*), Chinese plum (*Prunus salicina*) and carnation (*Dianthus caryophyllus*) (*Hu et al.*, 2015; *Liu et al.*, 2016; *Qiao et al.*, 2015; *Li et al.*, 2019).

HSFs in each subgroup are highly similar to each other across a variety of plants. Among these species, Class A contains the largest number of HSFs, followed by Class B, and then Class C. The same phenomenon was also observed in peach, which contained 11 *HSFAs*, six *HSFBs* and one *HSFCs*.

The HSF gene family expanded along with DSD in peach

The number of *HSFs* expanded markedly during plant evolution. The analysis of 51 representative species indicated that the HSF gene family largely expanded along with WGD during plant evolution (*Wang et al., 2018*). In Chinese white pear (*Pyrus bretschneideri*), most *PbHSF* expansions dated back to a recent WGD (*Qiao et al., 2015*). On the other hand, *GmHSFs* in cotton expanded along single gene duplication events (*Wang et al., 2014*). Here, DSD (67%) was the primary type of duplication for the HSF gene family in peach. The same phenomenon was also seen for the E3 ligase gene family in peach (*Tan et al., 2019*). It is probably that peach has not undergone a recent WGD (*Verde et al., 2013*).

HSF gene family was classified into three classes

Plant HSF proteins contain a few conserved characteristic domain (Guo et al., 2016). Generally, HSF families in plant species can be divided into three subfamilies, termed HSFA, HSFB, and HSFC (Liu et al., 2018; Wang et al., 2018). The classification of the PpHSF family was consistent with that in other plant species (tabreftabs7). Multiple sequence alignments revealed that an insertion occurred in the DBD domain near the N-terminus in the PpHSFA and PpHSFC groups. Like in other plants, the PpHSFA and PpHSFC genes contained inserted coding sequence for 21 and 7 aa in the HR-A/B region, respectively, while the HR-A/B region of PpHSFB was compact (Nover, 1996; Scharf et al., 2012). The organization, composition, number of conserved motifs in the HSFs differed among plant species (Wang et al., 2018). In Chinese whit pear, Class A in PbHSFs contained the most conversed motifs, followed by class B and then class C (Qiao et al., 2015). In this study, the number of motifs in the different classes was consistent with those in Chinese white pear. This also showed that members of the same class often have similar sequence structures in peach. For example, motif 5 was present only in PpHSFA and PpHSFC, while all Class B and Class A HSFs contain motif 4. The presence of these motifs may lead to functional group specificity. The similar classifications of HSF families in diverse plants showed that the HSF family was highly conserved during long-term evolution.

PpHSF5 acts as repressor of organ size in plants

In plants, organ size is primarily controlled by internal developmental signals (Mizukami, 2001; *Dubrovsky et al., 2006; Spradling, Drummond-Barbosa & Kai, 2001; West, Inzé & Beemster, 2004*). Previous research in the model organism *Arabidopsis thaliana* indicates that plant hormones and transcription factors, including HSFs, play crucial roles in growth and development (*Petricka, Winter & Benfey, 2012; Begum, Reuter & Schoffl, 2013*). HSFs as key transcription factors protect plants from various abiotic stresses and then participate in the growth and development (*Guo et al., 2016*). For example, *OsHsfA1a, OsHsfA1b* and *OsHsfA1d* are the main positive regulators of gene expression on heat stress-responsive, and four HSFA proteins play significant roles in gene expression of plant growth and development (*Yoshida et al., 2011*). In poplar (*Populus trichocarpa*), the transcripts of three *PtHsfs* in the B4 subfamily (-B4b, -B4c and -B4d) were maintained at higher levels during the leaf expansion stages (*Liu et al., 2019*). In carnation (*Dianthus caryophyllus*), five *DcaHsfs*, namely *DcaHsf-A1, A2a, A9a, B2a, B3a*, were involved in early flowering

stages (*Li et al.*, 2019). Transgenic Arabidopsis plants overexpressing *AtHSFB4* contained massively enhanced levels of *AtHSFB4* mRNAs and exhibited shorter roots (*Begum, Reuter* & *Schoffl*, 2013). In this study, overexpression of 35S:PpHSF5 in Arabidopsis resulted in not only shorter roots but also in lesser root volume and fewer lateral roots and root forks compared to WT.

The root system of a plant is instrumental to its growth and productivity because it is responsible for the extraction of water and mineral nutrients from the soil and their transport to aboveground parts of the plant (*Hochholdinger & Feix, 1998*). In this study, the *35S:PpHSF5* transgenic lines produced smaller aerial organs compared with WT. For example, the size (length and width) of rosette leaves were smaller than WT two and three weeks after transplanting, while the number of rosette leaves was not affected. The height of the overexpression lines was significantly lower than that in WT, while the number of internodes was not. Overexpression of *OsHsfA2e* in rice caused a dwarf phenotype (*Yokotani et al., 2007*). In plants overexpressing *BhHsf1*, the reduced organ size was mainly attributed to decreased cell proliferation (*Zhu et al., 2009*). The overexpression of *PpHSF5* in peach suggested that the dwarf phenotype of transgenic plants was caused by shorter internodes.

It is still unknown how *PpHSF5* regulates root and aerial organs development. *PpHSF5* is homologous to AtHSFB4 and thus may play similar roles in root development. Confocal laser scanning of roots in AtHSFB4-overexpression transgenic lines showed that ectopic division of the lateral root cap cells (LRC) occurred (Begum, Reuter & Schoffl, 2013). Previous studies indicated that auxin acts in the production of lateral root primordium (LR) (Casimiro et al., 2003; West, Inzé & Beemster, 2004). In the promoter of PpHSF5, there are three *cis-acting* regulatory elements that contain the auxin-inducible TGACG-motif. Two auxin-inducible TGA-box elements in the *GmGH3* promoter were strong binding sites of plant nuclear proteins and improved the auxin inducibility of the GmGH3 promoter (Zhan-Bin Liu et al., 1994). Moreover, the HS assays indicated PpHSF5-OE lines exhibited enhanced thermotolerance compared to WT. Similarity results were observed in transgenic Arabidopsis plants with AtHsfA3 and rice plants with OsHsfA2e (Ogawa, Yamaguchi & Nishiuchi, 2007; Yokotani et al., 2007). Therefore, PpHSF5 might be as a responsive factor for temperature change and involved in auxin signal transduction due to the TGA motifs in its promoter and might serve to negatively regulate root elongation and lateral root development, ultimately affecting the growth of aboveground parts of the plant.

CONCLUSIONS

In this report, 18 *PpHSF* genes were discovered in peach and found to be nonuniformly distributed on the peach chromosomes. The PpHSF family could be classified into three classes (PpHSFA, PpHSFB and PpHSFC) through multiple alignment, motif analysis and phylogenetic comparison. The expansion of the HSF gene family in peach occurred through DSD (67%) and WGD (33%). *PpHSF5* was expressed in diverse tissues and organs of the peach cultivar 'Zhongyoutao 14', with higher levels in young vegetative organs (leaf and apex). Transgenic Arabidopsis lines overexpressing *PpHSF5* showed massively enhanced

levels of *PpHSF5*. Ectopic expression *PpHSF5* repressed the length and number of roots, length and width of rosette leaves, and the height of plants, and enhanced thermtolerance in Arabidopsis after heat stress treatment. Our results further supplied functional and annotation information of the HSF gene family in general and revealed potential roles, outside of the response to heat stress, for *PpHSF5* during plant development.

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Competing Interests

The authors declare there are no competing interests.

Author Contributions

- Bin Tan conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Liu Yan and Huannan Li performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Xiaodong Lian, Jun Cheng and Wei Wang analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Xianbo Zheng and Xiaobei Wang performed the experiments, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Jidong Li, Xia Ye, Langlang Zhang and Zhiqian Li performed the experiments, authored or reviewed drafts of the paper, and approved the final draft.
- Jiancan Feng conceived and designed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.

Data Availability

The following information was supplied regarding data availability: Raw data is available in the Supplemental Files.

Supplemental Information

Supplemental information for this article can be found online at http://dx.doi.org/10.7717/ peerj.10961#supplemental-information.

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