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Expression profile analysis of 9 heat shock protein genes throughout the life cycle and under abiotic stress in rice

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Plant heat shock proteins (Hsps) facilitate protein folding or assembly under diverse developmental and adverse environmental conditions. Nine OsHsps were identified in our previous study from a proteomic analysis of rice cv. IRAT 109 leaf samples at the seedling stage under drought stress. To obtain additional information on the 9 *OsHsp* genes, this study focused on an expression profile analysis at different development stages throughout the life cycle of rice, and under different abiotic stresses and phytohormone treatments. The 9 genes exhibited distinctive expression patterns in different organs or development stages. Five of the genes (*OsHsp72.90*, *OsHsp72.57*, *OsHsp71.18*, *OsHsp24.15* and *OsHsp18.03*) showed high expression in the endosperm, indicating that *OsHsp* genes may play important roles in rice seed development. All 9 OsHsps were up-regulated under heat, polyethylene glycol, and abscisic acid treatment, whereas salt stress caused up-regulation of 6 genes (*OsHsp93.04*, *OsHsp71.10*, *OsHsp71.18*, *OsHsp72.57*, *OsHsp24.15* and *OsHsp18.03*) and cold stress resulted in down-regulation of *OsHsp93.04* and *OsHsp72.57*. These diverse expression profiles imply potential functional diversity of the *Hsp* gene family in rice.

heat shock protein, drought, heat, real-time PCR, rice

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Plants are frequently challenged by persistently changing stress factors such as drought, salinity, cold, and high temperatures. These abiotic stresses usually cause protein dysfunction. Maintenance of proteins in their functional conformations and prevention of the aggregation of non-native proteins are particularly important for cell survival under stress. Heat-shock proteins (Hsps) function as molecular chaperones that assist protein folding or assembly and prevent irreversible protein aggregation to maintain cellular homeostasis under both optimal and adverse developmental conditions [1].

Based on their approximate molecular weights, 5 major families of Hsps are recognized: Hsp100, Hsp90, Hsp70, Hsp60, and the small Hsp (sHsp) families. The Hsp100 ranges in molecular weight from 100 to 104 kD, which is a

major heat-regulated protein family in diverse organisms that interacts transiently with non-native protein species and prevents the aggregation and promotion of correct folding of proteins. Hsp101 was assumed to play a pivotal role in heat tolerance in *Arabidopsis thaliana*. Transgenic *Arabidopsis* plants expressing lower amounts of Hsp101 showed a severely diminished capacity to acquire heat tolerance after mild conditioning pretreatments [2].

The Hsps in the molecular weight range of 82 to 90 kD are classified as the Hsp90 family. These proteins have been isolated and cloned from many plant species. A rice (*Oryza sativa*) Hsp90 plays an important role in salt stress [3]. In *Arabidopsis*, AtHsp90s are essential for tolerance to biotic and abiotic stresses [4,5]. AtHsp90.3 was shown to confer heat tolerance in *Saccharomyces cerevisae*, but resulted in heat sensitivity in transgenic *Arabidopsis* plants [6]. PgHsp90 from *Pennisetum glaucum* was indicated to play

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an adaptive or protective role to counter stress-induced protein damage [7].

The Hsp70 family represents one of the most highly conserved classes of heat shock proteins and has molecular weights in the range of 68 to 75 kD. At least 18 genes encode Hsp70 proteins in *Arabidopsis* [8]. Cytosolic Hsp70 plays important roles under abiotic and biotic stresses [9]. AtHsp70s in the plastid stroma are important for thermotolerance of germinating seeds, which indicates that plastid physiology is important for seeds to endure heat stress [10]. In addition, several researchers showed that the Hsp70 family in other plant species is involved in plant development and tolerance to a variety of stresses [1,11–14].

The proteins of the Hsp60 family vary in size from 58 to 65 kD. It is generally agreed these proteins are important for assistance of plastid proteins such as Rubisco, and bind different proteins after their transcription and before folding to prevent aggregation of the proteins [15].

The small heat shock proteins (sHsps) include proteins with molecular weights in the range of 15–42 kD [16]. The sHsps can bind to partially folded or denatured proteins, which prevents irreversible unfolding or incorrect protein aggregation, or binds to unfolded proteins and allows further refolding by Hsp70/Hsp100 complexes [15]. Cloning and function analysis of sHsps have been performed in a variety of plant species. In rice, higher expression levels of sHsp17.7 were recorded in sHsp17.7 transgenic plants. A drought tolerance experiment showed overproduction of sHsp17.7 could increase drought tolerance in transgenic rice seedlings [17].

Rice is the main staple food for a large segment of the world's population and the productivity of rice is restricted by a broad range of environmental conditions [18]. In particular, drought and heat threaten food security. Thus, discovery and identification of stress-related genes is a very important topic. Proteomic analysis of the upland rice IRAT 109 at the seedling stage under drought stress was undertaken in our previous study, and 9 OsHsps involved in drought stress were identified [19]. To obtain additional information on the 9 *OsHsp* genes, the present study fo-

cused on an expression profile analysis of the *OsHsp* genes at different development stages throughout the life cycle of rice and under different abiotic stress and phytohormone treatments.

1 Materials and methods

1.1 Identification of the 9 OsHsp genes

Nine OsHsps were identified from the proteomic analysis in our previous study [19]. Molecular weights and theoretical isoelectric points (pI) were calculated with the EXPASY tool (http://www.expasy.org/tools/protparam.html). The characteristics of the genes are listed in Table 1. *Cis*-elements in the 1-kb upstream region of the 9 *OsHsp* genes were identified by analyzing the sequences in the Plant *cis*-acting Regulatory DNA Elements database (PLACE, http://www.dna. affrc.go.jp/PLACE/signalscan.html).

1.2 Genome-wide expression analysis of OsHsps in 2 rice cultivars

Expression profile data for OsHsps in 27 tissues of Minghui 63 and Zhenshan 97 were extracted from the CREP database [20]. Twenty-seven tissues (Table 2) from throughout the rice life cycle were selected. Expression values of each gene were logarithmized and cluster analyses were performed in *R* with euclidean distances and complete linkage hierarchical clustering.

Expression data was analyzed with Student's t-test to identify differentially expressed genes. The genes that were up- or down-regulated by more than 2-fold and with P < 0.05 were considered to be differentially expressed.

1.3 Expression analysis of the 9 OsHsp genes under abiotic stresses

Seeds of rice cv. IRAT109 were imbibed for 3 d in water, sown on moist paper and germinated in the dark at 25°C. Germinated seeds were planted in a plastic container with

Gene name	Molecular weight (Dalton)	TIGR locus	Isoelectric point	Family
OsHsp18.03	18030.82	LOC_Os01g08860.1	5.61	sHsp
OsHsp24.15	24153.29	LOC_Os02g52150.1	7.96	sHsp
OsHsp71.10	71100.37	LOC_Os03g16860.1	5.1	Hsp70
OsHsp71.18	71184.71	LOC_Os11g47760.1	5.1	Hsp70
OsHsp72.57	72565.29	LOC_Os03g02260.1	5.71	Hsp70
OsHsp72.90	72897.5	LOC_Os02g53420.1	5.49	Hsp70
OsHsp85.88	85878.03	LOC_Os08g38086.1	4.98	Hsp90
OsHsp93.04	93044.79	LOC_Os06g50300.1	4.89	Hsp90
OsHsp108.99	108985.34	LOC_Os03g31300.1	6.25	Hsp100

 Table 2
 Tissues selected from different developmental stages throughout

 the life cycle of rice

Sample No.	Developmental stage	Tissues
1	3 d after sowing	embryo and radicle
2	15 d after subculture	calli
3	screening stage	calli
4	5 d after regeneration	calli
5	72 h after imbibition	seed
6	trefoil stage	seedling
7	seedling with 2 tillers	shoot
8	seedling with 2 tillers	root
9	5 d before heading	stem
10	5 d before heading	flag leaf
11	heading stage	stem
12	heading stage	panicle
13	14 d after heading	flag leaf
14	young panicle at stage 3 (secondary branch primordium differentiation stage)	leaf
15	young panicle at stage 3	sheath
16	young panicle at stage 3	panicle
17	young panicle at stage 4 (pistil/stamen primordium differentiation stage)	panicle
18	young panicle at stage 5 (pollen-mother cell formation stage)	panicle
19	4–5 cm young panicle	leaf
20	4–5 cm young panicle	sheath
21	4–5 cm young panicle	panicle
22	1 d before flowering	hull
23	1 d before flowering	stamen
24	3 d after pollination	spikelet
25	7 d after pollination	endosperm
26	14 d after pollination	endosperm
27	21 d after pollination	endosperm

Hoagland nutrient solution, and were incubated in a growth chamber under a 12 h light/12 h dark cycle with 80% RH. The trefoil-stage seedlings were subjected to the following five treatments. To simulate drought stress, salt stress, and for abscisic acid (ABA) treatment, seedlings were irrigated with Hoagland liquid culture containing 20% polyethyleneglycol 6000 (PEG), 150 mmol/L NaCl, and 100 µmol/L ABA, respectively. For cold and heat shock treatments, seedlings were exposed to 4 and 42°C. Samples were collected after treatment exposure for 0, 3, 6, 12, and 24 h and after recovery for 24 h. Ten shoots were combined as one biological replicate and each treatment consisted of three repeats.

Samples were ground in liquid nitrogen using a mortar and pestle. Total RNA was extracted using TRIzol reagent (Invitrogen) according to the manufacturer's instructions. First-strand cDNA was synthesized using the SuperScript First-strand Synthesis Kit (Invitrogen, USA). Real-time PCR was performed in an optical 96-well plate, including 12.5 μ L 2× SYBR Green Master Mix Reagent (TaKaRa), 1 μ L cDNA samples, and 0.2 mmol/L of each gene-specific primer, in a final volume of 25 μ L, using the following thermal cycling procedure: 50°C for 2 min, 95°C for 10 min; 40 or

45 cycles of 95°C for 30 s; 60°C for 30 s; and 72°C for 1 min. Disassociation curve analysis was performed as follows: 95°C for 15 s; 60°C for 20 s; and 95°C for 15 min. The *actin1* gene was used as an endogenous control to normalize variance in the quality of RNA and the amount of cDNA used. Primers were designed according to the sequences within the boundaries defined by the cDNA ends (Table S1) and synthesized by Invitrogen.

2 Results

2.1 Identification of the 9 OsHsp genes

In the proteomic analysis, 13 spots representing 9 OsHsp proteins were up-regulated under drought stress. These OsHsps were designated as OsHsp18.03, OsHsp24.15, OsHsp71.10, OsHsp71.18, OsHsp72.57, OsHsp72.90, OsHsp85.88, OsHsp93.04 and OsHsp108.99, which were divided into 4 classes according to the molecular weight. Two genes (*OsHsp18.03* and *OsHsp24.15*) belong to the sHsp family, 4 OsHsps (*OsHsp71.10*, *OsHsp71.18*, *OsHsp72.57* and *OsHsp72.90*) are part of the OsHsp70 family, 2 genes (*OsHsp85.88* and *OsHsp93.04*) belong to the OsHsp90 family, and *OsHsp108.99* belongs to the OsHsp100 family.

2.2 *Cis*-elements in the upstream region of the 9 *OsHsp* genes

A search of the PLACE database with the 1-kb upstream region of the 9 *OsHsp* genes as queries found at least 19 previously identified putative stress-responsive or organ-specific *cis*-elements in the promoter regions of the *OsHsp* genes (Table S2). These *cis*-elements include the heat shock-responsive CCAATBOX [21], the ABA-responsive ABRE-LATERD1, the dehydration-responsive ACGTATERD1 and MYBCORE [22,23], the light-regulated GT1CONSENSUS and IBOXCORE [24], and the defense-responsive WBO-XATNPR1 [25]. The dehydration-responsive ACGTATE-RD1 was the most frequent stress-related *cis*-element, with 10 and 14 found in *OsHsp72.9* and *OsHsp93.4*, respectively.

2.3 Expression analysis of OsHsp genes in different tissues

The corresponding probes for all of the 9 *OsHsp* genes were included on the Crep chip data. The average signal values of the 9 *OsHsp* genes are listed in Table S3.

Hierarchical cluster analysis based on the signal values showed that the *OsHsp* genes had very diverse expression patterns (Figure 1). The 9 *OsHsp* genes were expressed in at least one developmental stage or tissue. Hierarchical clustering, which was based on the mean log-transformed intensity value of the genes, enabled the expression profiles to be divided into two major groups. Group I contained 4 genes (*OsHsp93.04*, *OsHsp71.10*, *OsHsp85.88* and *OsHsp108.99*).

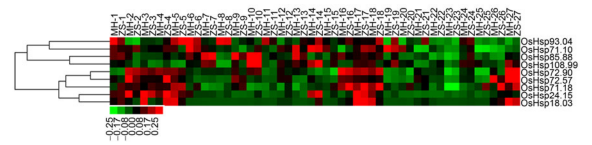


Figure 1 Hierarchical clustering display of *OsHsp* family genes in rice based on the average signal values in 27 tissues. Numbers refer to: 1, embryo and radicle after germination; 2, calli, 15 d after subculture; 3, calli, screening stage; 4, calli, 5 d after regeneration; 5, seed, 72 h after imbibition; 6, root and leaf at three-leaf stage; 7, shoot, seedling with 2 tillers; 8, root, seedling with 2 tillers; 9, stem, 5 d before heading; 10, flag leaf, 5 d before heading; 11, stem, heading stage; 12, panicle, heading stage; 13, flag leaf, 14 d after heading; 14, leaf, young panicle at stage 3; 15, sheath, young panicle at stage 3; 16, young panicle at stage 3; 17, young panicle at stage 4; 18, young panicle at stage 5; 19, leaf, 4–5 cm young panicle; 20, sheath, 4–5 cm young panicle; 21, panicle, 4–5 cm young panicle; 22, hull, 1 d before flowering; 23, stamen, 1 d before flowering; 24, spikelet, 3 d after pollination; 25, endosperm, 7 d after pollination; 26, endosperm, 14 d after pollination; 27, endosperm, 21 d after pollination. Each number comprises three parallel lines that indicate Minghui 63 (MH) and Zhenshan 97 (ZS), respectively. The colour scale (representing average signal values) is shown at the bottom.

Transcripts of both *OsHsp93.04* and *OsHsp71.10* were more frequent in the embryo, leaf and panicle, and *OsHsp71.10* was also highly expressed in the root. High expression levels of *OsHsp85.88* and *OsHsp108.99* were detected in the embryo, leaf, stem, panicle and mature endosperm (21 d after pollination).

The expression patterns of the genes in Group II differed from those of Group I. All 5 genes were highly expressed in calli, the panicle and endosperm. Up-regulation of OsHsp72.90 was detected 21 d after pollination in endosperm, and transcripts of OsHsp72.59 were up-regulated in the endosperm at 14 and 21 d after pollination. Expression levels of OsHsp71.18 were lower in Minghui 63 than those in Zhenshan 97. In addition, gradual up-regulation of OsHsp71.18 was detected in the endosperm from 7 d after pollination to 21 d after pollination. OsHsp24.15 was predominantly expressed in calli, the embryo, leaf (young panicle at secondary branch primordium differentiation stage), young panicle at stage 4 (pistil/stamen primordium differentiation stage) and stage 5 (pollen mother cell formation stage), and moderately expressed in the endosperm. OsHsp18.03 was expressed preferentially in calli, the young panicle at stage 4 (pistil/stamen primordium differentiation stage), and was gradually up-regulated in the endosperm from 7 d after pollination to 21 d after pollination.

Overall, the *OsHsp* genes in Group I showed a higher expression level in vegetative tissues, and fewer transcripts were detected in calli and the endosperm compared to those of OsHsps in Group II. The expression patterns were almost identical between Minghui 63 and Zhenshan 97, except for the expression levels.

2.4 Expression profiles of the 9 OsHsps under phytohormone treatments at trefoil seedling stage

The responses of the 9 *OsHsp* genes to gibberellin3 (GA3), Naphthaleneacetic Acid (NAA) and Kinetin (KT) were also based on the CREP chip data. Five genes were responsive to all three phytohormones. Three OsHsp genes (*OsHsp71.10*,

OsHsp71.18 and OsHsp93.04) were down- regulated under the GA3, NAA and KT treatments, whereas two genes (OsHsp85.88 and OsHsp108.99) were up-regulated in response to these hormones. The hierarchical cluster display of the 9 genes in the hormonal treatments in Minghui 63 and Zhenshan 97 is shown in Figure 2, and the average signal values are presented in Table S4.

Because the 9 *OsHsp* genes were identified by proteomic analysis of rice IRAT 109 leaf samples at the seedling stage under drought stress, IRAT109 seedlings were treated with ABA at the three-leaf stage. Interestingly, all 9 genes were significantly up-regulated after ABA treatment for 12 h. However, the expression levels decreased back to the levels of non-stressed samples after the stress was released for 24 h (Figure 3).

2.5 Expression patterns of the 9 OsHsps under abiotic stresses

Numerous abiotic stresses are known to trigger alteration in

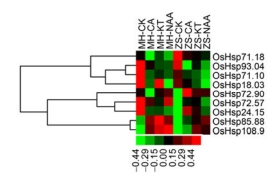


Figure 2 Expression profile of *OsHsp* family genes under different hormonal stress conditions (NAA, GA3 and KT) in Minghui 63 and Zhenshan 97. The colour scale (representing average signal values) is shown at the bottom. MH-CK (Minghui 63, control), MH-GA3 (Minghui 63, GA3), MH-KT (Minghui 63, KT), MH-NAA (Minghui 63, NAA), ZH-CK (Zhenshan 97, control), ZH-GA3 (Zhenshan 97, GA3) ZH-KT (Zhenshan 97, KT) and ZH-NAA (Zhenshan 97, NAA).

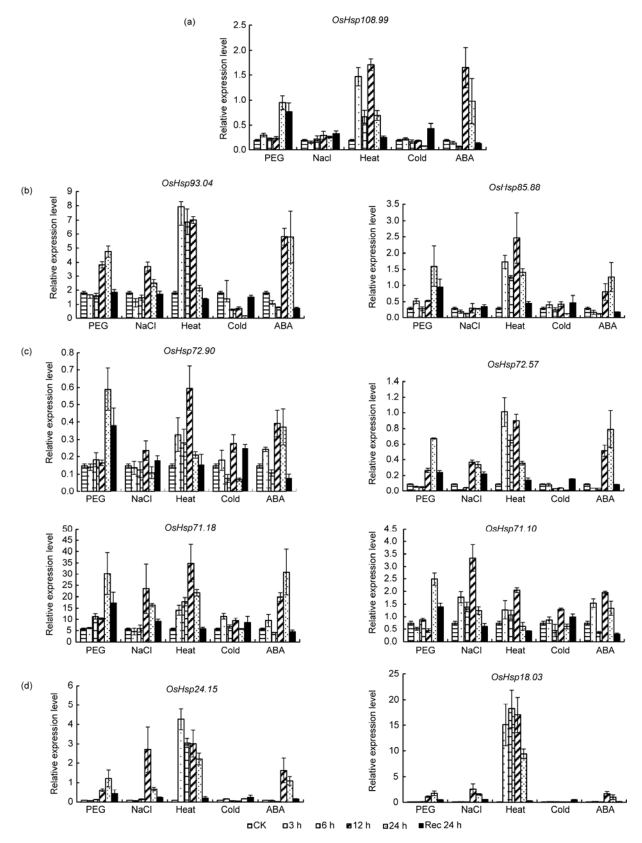


Figure 3 Relative quantification by real-time PCR analysis of the expression levels of *OsHsp* genes in rice IRAT109 in response to abiotic stress treatments. Rice *Actin1* was used as the internal control. The bars are means of the relative fold change of three biological and two technical replicates obtained by real-time PCR. The standard errors of the biological replicates are shown as error bars. Relative expression levels are shown for (a) *OsHsp108.99*, which belongs to the *OsHsp100* family; (b) the two *OsHsp* genes in the *OsHsp90* family; (c) the four *OsHsp* genes in the *OsHsp70* family; and (d) the two *OsHsp* genes in the *sHsp* family.

the transcription of *Hsp* genes. Therefore, we further checked the expression profiles of the 9 *OsHsp* genes under diverse abiotic stress treatments using real-time PCR. Under dehydration stress, transcript levels of the 9 *OsHsp* genes were enhanced. Four genes (*OsHsp93.04*, *OsHsp72.57*, *OsHsp24.15* and *OsHsp18.03*) were up-regulated after 12-h treatment with PEG, and a high level of transcripts of the other five genes was detected after 24-h treatment. After recovery for 24 h, transcript levels of most of the *OsHsp* genes decreased to the level prior to stress treatment.

Significantly increased levels of the 9 *OsHsp* genes were detected after 3 h at 42°C, which indicated these genes are quickly and sharply induced by heat shock stress. This was especially true for the genes *OsHsp24.15* and *OsHsp18.03*, which were up-regulated by more than 48- and 227-fold, respectively.

Salt stress caused up-regulation of 6 genes (*OsHsp93.04*, *OsHsp71.10*, *OsHsp71.18*, *OsHsp72.57*, *OsHsp24.15* and *OsHsp18.03*). However, cold stress caused down-regulation of *OsHsp93.04* and *OsHsp72.57*.

3 Discussion

3.1 OsHsp genes may be involved in seed development

Plant Hsps are expressed during an intriguing array of developmental processes, which indicates they may have important roles in the absence of stresses [26]. The endosperm in monocot seeds plays a very important role in seed development and can be the principal nutrient storage tissue in seeds. The genes in *OsHsp* Group II (*OsHsp72.90*, *OsHsp72.57*, *OsHsp71.18*, *OsHsp24.15* and *OsHsp18.03*) showed high expression in the endosperm, indicating that *OsHsp* genes may play important roles in rice seed development.

In plants, sHsps are generally undetectable in vegetative tissues in the absence of stress, but are expressed during specific stages of plant development, and sHsp expression during seed development is the most extensively characterized example of this non-stress regulation [27]. sHsps accumulated at mid-maturation and were abundant throughout the late maturation period and in the dry seed in *Arabidopsis* [26], pea [28], and sunflower [29].

AtHsp101 is a component of mature seeds and regulates seed development in the absence of stress, in a pattern similar to that seen for Late Embryogenesis Abundant (LEA) proteins and small Hsps; protein accumulates during midmaturation and is stored in the dry seed [29,30].

The mechanisms that regulate Hsp expression during seed maturation remain largely unknown. Using a mutant approach, a genetic framework in which HsfA9 operates as a specialized heat-shock transcription factor (Hsf) for the developmental expression of *Hsp* genes during seed maturation was established, which showed that ABI3 could activate the HsfA9 promoter, whereas HsfA9 in turn was shown to be a potent activator on the promoters of *Hsp* genes [31].

3.2 Function of OsHsps under drought and heat stress

Drought and heat stress are among the abiotic factors that cause the most severe damage to plant crops. In nature, individual stresses, such as drought and high temperature, usually do not occur separately. A combination of drought and heat shock resulted in the closure of stomata, suppression of photosynthesis, enhancement of respiration, and increased leaf temperature. To counter the effects of such stress, plants express a variety of defense genes such as heat shock proteins [32].

The 9 OsHsps analyzed in the present study were identified in a proteomic analysis of rice leaf samples at the seedling stage under drought stress in our previous study [19]. The transcription levels of all 9 *OsHsp* genes were significantly enhanced both under dehydration and heat stress, which showed they may play roles in drought and heat stress.

In the *cis*-element analysis of the 9 *OsHsp* genes, *cis*-elements related to heat shock and dehydration were observed. It is well known that the heat-inducible binding of Hsfs to the heat-shock element (HSE) promoter sequences control the expression of *Hsp* genes during stress and development [33,34]. The analysis carried out in the present study showed that promoters of the 9 *OsHsp* genes harbor a minimum of one module of a putative HSE. The dehydration-responsive elements ACGTATERD1 and MYBCORE existed in the promoter of the 9 *OsHsp* genes. ACGTATERD1 and MYBCORE are important *cis*-elements for ERD1 and MYB proteins, respectively, involved in the regulation of stress-responsive genes [22,23].

In Arabidopsis, the expression level of At-Hsp17.4 was decreased in mutant lines and resulted in a seed desiccation-intolerance phenotype, and At-Hsp17.6A was induced by heat and osmotic stress [27,35]. In rice, tolerance to drought stress was enhanced in transgenic rice plants overexpressing a small heat-shock protein, sHsp17.7, and transgenic seedlings showing higher expression levels of sHsp 17.7 protein could regrow after resumption of watering [17]. In tobacco, transgenic tobacco plants producing elevated levels of the tobacco Hsp70, NtHsp70-1, exhibited tolerance to water stress, as indicated by the higher leaf water content in the transgenic plants compared to that of the wild type [36]. In addition, seven Hsps were induced in response to a combination of drought and heat shock. These included cytosolic Hsp90, Hsp70, and Hsp100, and sHsps (cytosolic, mitochondrial, and chloroplastic). Furthermore, the induction of Hsps was higher in response to drought and heat shock compared with heat shock or drought individually [37]. In maize, Hsp70 and sHsps (sHsp17.2, sHsp17.4 and sHsp26) are responsive to a combination of drought and heat stress [12,38]. Four Hsp genes (Hsp17.8, Hsp26.3, Hsp70 and Hsp101b) in Triticum aestivum plants were monitored via one-step RT-PCR and SDS-PAGE analysis in plants subjected to water deprivation or high temperature individually and in combination. The combination of drought and heat induced Hsp expression more effectively than the individually applied stresses and the induction of Hsps was greater in the drought-tolerant wheat cultivar Katya than in the drought-sensitive cultivar Sadovo [39,40].

3.3 OsHsp genes may be involved in ABA-regulated pathways

ABA regulates many plant responses to environmental stress. The greatest increase in ABA concentration is observed in response to drought stress, and temperature also has an effect on endogenous ABA levels [41].

The 9 *OsHsp* genes identified in the present study were up-regulated under PEG and heat stress as well as in response to ABA treatment. Moreover, ABRELATERD1, a *cis*-acting element involved in the ABA function, was present in all 9 *OsHsp* genes, which indicated they may be associated with the key roles of ABA in plant metabolism.

Hsp genes regulated by ABA have been reported in several plant species, such as maize, tobacco, wheat and rice [12,36,42–44]. Results of studies with an ABA-deficient mutant indicated that endogenous ABA regulated Hsp70 synthesis in the leaves and roots of maize seedlings exposed to drought, heat, and combined drought and heat stress, which implied that ABA may improve plant tolerance to stress by increasing Hsp70 expression. In addition, ABA is involved in the enhancement of enzyme activities and H_2O_2 production in leaves and roots subjected to a number of stress treatments [12].

In summary, analysis of the tissue and phytohormone expression profiles of the 9 *OsHsp* genes has improved the functional dissection of rice *Hsp* genes. Elucidation of the precise role of each *OsHsp* gene, however, requires the use of other experimental approaches including, for example, overexpression and/or RNAi strategies.

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Supporting Information

- Table S1
 Primers used in the real-time PCR analysis
- **Table S2** Number of *cis*-elements in the 1-kb upstream region of *OsHsp* genes
- **Table S3** Average signal values of the nine OsHsp genes expressed in 27 tissues for Minghui 63 and Zhenshan 97
- Table S4 Average signal values of the nine OsHsp genes under the hormonal treatments for Minghui 63 and Zhenshan 97

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