Genomewide identification and analysis of heat-shock proteins 70/110 to reveal their potential functions in Chinese soft-shelled turtle *Pelodiscus sinensis*

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Abstract
Heat-shock proteins 70/110 (Hsp70/110) are vital molecular chaperones and stress proteins whose expression and production are generally induced by extreme temperatures or external stresses. The Hsp70/110 family is largely conserved in diverse animals. Although many reports have studied and elaborated on the characteristics of Hsp70/110 in various species, the systematic identification and analysis of Hsp70/110 are still poor in turtles. In this study, a genomewide search was performed, and 18 candidate *PsHSP70/110* family genes were identified in Chinese soft-shelled turtle, *Pelodiscus sinensis*. These *PsHSP70/110* proteins contained the conserved "heat shock protein 70" domain. Phylogenetic analysis of *PsHSP70/110* and their homologs revealed evolutionary conservation of Hsp70/110 across different species. Tissue-specific expression analysis showed that these *PsHSP70/110* genes were differentially expressed in different tissues of *P. sinensis*. Furthermore, to examine the putative biological functions of *PsHSP70/110*, the dynamic expression of *PsHSP70/110* genes was analyzed in the testis of *P. sinensis* during seasonal spermatogenesis following germ cell apoptosis. Notably, genes such as *PsHSPA1B-L*, *PsHSPA2*, and *PsHSPA8* were significantly upregulated in *P. sinensis* testes along with a seasonal decrease in apoptosis. Protein interaction prediction revealed that *PsHSPA1B-L*, *PsHSPA2*, and *PsHSPA8* may interact with each other and participate in the MAPK signaling pathway. Moreover, immunohistochemical analysis showed that *PsHSPA1B-L*, *PsHSPA2*, and *PsHSPA8* protein expression was associated with seasonal temperature variation. The expression profiling and interaction relationships of the *PsHSPA1B-L*, *PsHSPA2*, and *PsHSPA8* proteins implied their potential roles in inhibiting the apoptosis of germ cells in *P. sinensis*. These results provide insights into *PsHSP70/110* functions and will serve as a rich resource for further investigation of HSP70/110 family genes in *P. sinensis* and other turtles.

Keywords
Chinese soft-shelled turtle *Pelodiscus sinensis*, expression profiling, heat-shock proteins 70/110, protein interaction, regulation of apoptosis
Heat-shock proteins (Hsps), which were first reported in Drosophila, are ubiquitously found in bacteria, plants, and animals (Arya, Mallik, & Lakhotia, 2007). When organisms are exposed to extreme temperatures or external stresses such as disease, toxins, and hypoxia, Hsps can be synthesized as stress proteins and accumulate to respond to various environmental insults (Gupta, Sharma, Mishra, Mishra, & Chowdhuri, 2010; Srivastava, 2002). Based on the approximate molecular weight and amino acid sequence homologies of Hsps, they are distinctly classified into five major families: Hsp110, Hsp90, Hsp70, Hsp60, and small Hsps (sHsps) (Lindquist & Craig, 2003). Among these proteins, Hsp70 is one of the most heat inducible and evolutionally conserved in terms of structure and function (Murphy, 2013). Hsp70 proteins generally form two groups: stress-inducible Hsp70 proteins and constitutively expressed heat-shock cognate 70 (Hsc70) proteins, according to expression profiling (Kiang & Tsokos, 1998). Two major conserved domains, a nucleotide-binding domain (NBD) and a substrate-binding domain (SBD), are characteristic of Hsp70 family proteins (Bertelsen, Chang, Gestwicki, & Zuiderweg, 2009). The intrinsic activities and allosteric coupling of NBD and SBD are associated with the functions of Hsp70 (Bertelsen et al., 2009).

Specifically, Hsp110, which exhibits a longer C-terminal extension, shares the same domain organization and exhibits highly similar crystal structures to Hsp70, which reveals the close relationships of the Hsp110 and Hsp70 protein families (Dragovic, Broadley, Shomura, Bracher, & Hartl, 2006).

| Gene name     | NCBI accession          | Conserved domains                                                                 |
|---------------|-------------------------|-----------------------------------------------------------------------------------|
| PsHSPA1A-L    | XM_014572517.1, XP_014428003.1 | cl17037, NBD_sugar-kinase_HSP70_actin                                             |
| PsHSPA1B-L    | XM_006134687.2, XP_006134749.1 | cl17037, NBD_sugar-kinase_HSP70_actin; PT00009, heat-shock 70-kDa protein         |
| PsHSPA2       | NM_001287561.1, NP_001274490.1 | cd10233, HSPA1-2, 6-8-like_NBD; PT00009, heat-shock 70-kDa protein                |
| PsHSPA4-X1    | XM_006128531.2, XP_006128593.1 | cl17037, NBD_sugar-kinase_HSP70_actin                                             |
| PsHSPA4-X2    | XM_006128532.2, XP_006128594.1 | cd11737, HSPA4_NBD; pfam00012, Hsp70 protein                                      |
| PsHSPA4L      | XM_014577892.1, XP_014433378.1 | cl17037, NBD_sugar-kinase_HSP70_actin; pfam00012, Hsp70 protein                  |
| PsHSPA5       | NM_001286892.1, NP_001273821.1 | cd10241, HSPA5-like_NBD; pfam00012, HSP70                                        |
| PsHSPA8*      | NM_001286908.1, NP_001273837.1 | cd10233, HSPA1-2, 6-8-like_NBD; PT00009, heat-shock 70-kDa protein; cl00788, MttA_Hcf106 |
| PsHSPA9       | XM_014569866.1, XP_014425352.1 | cd11733, HSPA9-like_NBD; PRK00290, molecular chaperone DnaK                       |
| PsHSPA12A     | XM_006120999.2, XP_006121061.1 | cl17037, NBD_sugar-kinase_HSP70_actin                                             |
| PsHSPA12A-L1  | XM_006127821.2, XP_006127883.1 | cd10229, HSPA12_like_NBD                                                          |
| PsHSPA12A-L2  | XM_014573716.1, XP_014429202.1 | cl17037, NBD_sugar-kinase_HSP70_actin                                             |
| PsHSPA12A-L3  | XM_014572222.1, XP_014427708.1 | cd10229, HSPA12_like_NBD                                                          |
| PsHSPA12A-L4  | XM_006127819.2, XP_006127881.1 | cd10229, HSPA12_like_NBD                                                          |
| PsHSPA12B-L   | XM_014570751.1, XP_014426237.1 | cl17037, NBD_sugar-kinase_HSP70_actin                                             |
| PsHSPA13      | XM_006136928.2, XP_006136990.1 | cd10237, HSPA13-like_NBD; PRK13930, rod shape-determining protein MreB            |
| PsHSPA14      | XM_014570505.1, XP_014425991.1 | cl17037, NBD_sugar-kinase_HSP70_actin                                             |
| PsHSPH1       | XM_006125320.1, XP_006125382.1 | cl17037, NBD_sugar-kinase_HSP70_actin                                             |

*Also known as PsHsc70.
By functioning as a molecular chaperone in the folding, denaturation, degradation, and inhibition of proteins and controlling regulatory proteins, Hsp70 plays essential roles in heat adaptation and protection against stresses in diverse species (Murphy, 2013). Extensive evidence has suggested that Hsp70 not only exhibits ATP-dependent chaperoning function but is also a negative apoptosis-inducing factor (AIF) in response to a wide range of stimuli (Goloudina, Demidov, & Garrido, 2012; Jiang et al., 2011; Sabirzhanov, Stoica, Hanscom, Piao, & Faden, 2012). In rodent models, overexpression of Hsp70 provides a survival advantage to tumor cells because Hsp70 can interact with multiple components of the apoptotic machinery (Jäättelä, 1995). Conversely, it has been reported that Hsp70 knockdown leads to decreased cell proliferation and facilitates the induction of apoptosis in multiple cancer cell models (Kotoglou et al., 2009; Zhang et al., 2013). Indeed, Hsp70 can block apoptosis by binding to apoptosis protease-activating factor 1 (Apaf1), thereby preventing the recruitment of procaspase-9 to the apoptosome (Beere et al., 2000). Similarly, Hsp70 regulates the important apoptotic mediator Bax and prevents Bax from translocating to mitochondria, which is necessary for the disruption of the mitochondrial membrane (Stankiewicz, Lachapelle, Foo, Radicioni, & Mosser, 2005). The regulatory roles of Hsp70 in apoptosis are also due to the effects of Hsp70 on stress-induced kinases, including SAPK/JNK, p38, and apoptosis signal-regulating kinase (Park et al., 2002; Park, Lee, Huh, Seo, & Choi, 2001). Additionally, Hsp70 can inhibit caspase-independent apoptosis by directly interacting with AIF and cathepsins (Jesper et al., 2004; Ravagnan et al., 2001). Despite considerable research advances, the antiapoptotic mechanism of Hsp70 is still controversial, especially in nonmodel animals. In recent studies, many genes encoding Hsp70 have been identified and characterized from nonmodel animals such as amphibians, insects, crustaceans, mollusks, and fishes (Luan et al., 2010; Simoncelli, Morosi, Rosa, Pascolini, & Fagotti, 2010; Song et al., 2016; Wang et al., 2019; Wang, Wu, Jian, & Lu, 2009), enriching knowledge of the phylogenetic relationships and biological functions of Hsp70. However, few studies have focused on the genomewide identification and functional analysis of the Hsp70 gene family in turtles.

Chinese soft-shelled turtle (Pelodiscus sinensis), a reptile, presents important economic value and is widely distributed in Asian countries such as China, Japan, and Korea. P. sinensis is an ectothermic aquaculture species with a specific evolutionary role linking ectothermic amniotic animals (fishes and amphibians) and endothermic amniotic animals (birds and mammals) (Zimmerman, Vogel, & Bowden, 2010) and can thus be used as a potential animal model to study the evolution of critical genes or species (Liu, Chu, et al., 2016). The body temperature of P. sinensis is dependent on the ambient temperature, similar to other ectotherms, resulting in typical hibernation patterns in midwinter (Chen et al., 2015). Previous studies have revealed distinct seasonal apoptosis in the testis of P. sinensis on the basis of morphological and molecular evidence (Liu et al., 2017). Furthermore, it is well known that Hsp70/110 presents the obligatory function of responding to adverse external stimuli, especially heat shock, and exhibits survival-promoting effects and suppression of apoptosis (Gao

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**FIGURE 1** Characterization of the identified PsHSP70/110 genes in Pelodiscus sinensis. (a) Phylogenetic analysis of PsHSP70/110 proteins using bootstrap values with 1,000 replications. (b) A diagram of conserved domains in PsHSP70/110 proteins.
et al., 2014). However, the relationship between apoptosis and stress-related Hsp70/110 is poorly understood in P. sinensis. Fortunately, the public genomic sequences and RNA-seq data of P. sinensis (Liu et al., 2017; Liu, Yang, et al., 2016; Wang et al., 2013) can provide rich resources for the identification, phylogenetic analysis and functional exploration of PsHsp70/110 genes in P. sinensis.

The primary goals of this study were to systematically identify candidate PsHSP70/110 family genes based on the whole-genome sequence of P. sinensis and to analyze their classification, conserved structures, and phylogenetic relationships. Furthermore, we focused on the mRNA and protein expression of PsHSP70/110 genes to investigate their putative roles in germ cell apoptosis following seasonal temperature variation. These results will provide useful information for the exploration of PsHSP70/110 functions in P. sinensis and facilitate further investigation of Hsp70/110 family genes in turtles.

2 | MATERIALS AND METHODS

2.1 | Animals

In this study, all sample procedures and animal care were conducted according to the guidelines of the Animal Research Institute Committee (Northwest A&F University, Shaanxi, China). The protocol was approved by the Science and Technology Agency of Shaanxi Province under permit NO. SYXK (SN) 2018-0003. All efforts were made to minimize animal suffering. Healthy adult Chinese soft-shelled turtles (3–4 years old with an average weight of 1.09 ± 0.15 kg) were captured from Yangcheng Lake in Suzhou (31°N, 120°E), Jiangsu province, China. Suzhou has a typical temperate climate with four distinct seasons, including spring (March–May), summer (June–August), autumn (September–November), and winter (December–February). Prior to the experiments, the turtles were acclimated in a tank with a recirculating water system for one week. To analyze the tissue-specific expression of PsHSP70/110 mRNA in P. sinensis, five male and five female turtles that were breeding at 25 °C were anesthetized by intraperitoneal administration of sodium pentobarbital (20 mg/kg) and killed by cervical dislocation. Tissue samples (including muscle, heart, blood, brain, kidney, liver, lung, spleen, intestine, testis, and oviduct) were collected immediately. For dynamic expression analysis of PsHSP70/110 genes in the testis, male turtles collected in April (spring, average temperature 15.7°C), July (summer, average temperature 28.6°C), and October (autumn, average temperature 18.4°C) in 2017 were used to obtain testis samples with five replicates. The testis sample from one side of each turtle was fixed for immunohistochemistry (IHC) analysis. The other side of the testis was placed in liquid nitrogen immediately and kept at −80°C until RNA extraction.

2.2 | Identification and conserved domains of Hsp70/110 family genes in P. sinensis

The public P. sinensis genome sequences (Wang et al., 2013) were used to identify potential Hsp70/110 genes via genomewide searches. A BLASTp homology search against the NCBI database was also performed by using the reported Hsp70/110 sequences from human and mouse as a query. To confirm the candidate PsHSP70/110 genes, the obtained protein sequences were further subjected to analysis of conserved domains against public databases including NCBI Conserved Domain Database (http://www.ncbi.nlm.nih.gov/cdd), Pfam (http://pfam.xfam.org), and InterProScan (http://www.ebi.ac.uk/Tools/pfa/interproscan5). The nucleotide and amino acid sequences of PsHSP70/110 genes were retrieved and used for further analysis. These candidate PsHSP70/110 family genes were named according to the homologous gene names, annotations, and gene descriptions in the NCBI database. The physical and chemical characteristics of PsHSP70/110 genes were analyzed by using ExPASy (http://web.expasy.org/protparam). The molecular weights, theoretical pI, instability index, aliphatic index, and grand average of hydropathicity (GRAVY) were calculated.

2.3 | Phylogenetic analysis and alignment of Hsp70/110 genes in different species

To detect the phylogenetic relationship of PsHSP70/110 genes, the protein sequences of Hsp70/110 family genes from humans (Homo sapiens), house mouse (Mus musculus), platypus (Ornithorhynchus anatinus), turkey (Meleagris gallopavo), chicken (Gallus gallus), American alligator (Alligator mississippiensis), green anole (Anolis carolinensis), western painted turtle (Chrysemys picta bellii), three-toed box turtle (Terrapene mexicana triunguis), green sea turtle (Chelonia mydas), African clawed frog (Xenopus laevis), and zebrafish (Danio rerio) were downloaded from the NCBI database. The phylogenetic relationships among different species were analyzed using the TimeTree server (http://www.timetree.org). The full-length sequences of Hsp70/110 proteins from different species were used for sequence alignment and phylogenetic analysis. Sequence alignment was performed by using MAFFT software. The phylogenetic tree was constructed by using PhyML 3.0 software (Guindon et al., 2010) based on the maximum-likelihood (ML) method and bootstrap values with 1,000 replications.

2.4 | Quantitative real-time PCR analysis

Quantitative real-time PCR (qRT-PCR) analysis was performed according to previous reports (Liu et al., 2017; Liu, Yang, et al., 2016). Total RNA was extracted using TRIzol reagent (Life Technologies). cDNA was synthesized using the SuperScript First-Strand Synthesis System (Invitrogen). Primer sequences for gene expression analysis were designed using Beacon Designer software (Premier Biosoft International). The relative expression levels of genes were normalized to β-actin and analyzed using the $2^{-ΔΔCT}$ method (Livak & Schmittgen, 2001).

2.5 | Transcriptomic expression analysis of PsHSP70/110 genes in P. sinensis

The dynamic expression patterns of PsHSP70/110 genes in the testis of P. sinensis in different months were detected by using published
RNA-seq data from *P. sinensis* testis to calculate gene expression levels. Transcriptomic analysis of *P. sinensis* testes in April (spermatogenically quiescent phase, MT-1), July (intermediate spermatogenesis, MT-2), and October (late spermatogenesis, MT-3) was performed in our previous study (Liu et al., 2017). RNA-seq data from the three libraries are available in the NCBI Sequence Read Archive (SRA, http://www.ncbi.nlm.nih.gov/Traces/sra) under accession numbers SRX2351846 (MT-1), SRX2352135 (MT-2), and SRX2352136 (MT-3). The expression patterns of *PsHSP70/110* genes in different months were calculated by the fragments per kb per million reads (FPKM) method (Mortazavi, Williams, McCue, Schaeffer, & Wold, 2008). The FPKM value associated with an abundance of zero was set to 0.01 for gene expression analysis. Heat maps of gene expression were generated by using Cluster software (de Hoon, Imoto, Nolan, & Miyano, 2004) and Java Treeview software (Saldanha, 2004).

### 2.7 Immunohistochemistry

Immunohistochemistry was performed as previously described (Liu et al., 2017). Briefly, sections of *P. sinensis* testis were dewaxed in xylene, dehydrated in an ethanol series, and blocked with 3% H$_2$O$_2$ in distilled water. The sections were incubated with anti-HSPA1L/ HSPA2 (ab154374, Abcam Inc., Cambridge, MA, USA; 1:100), anti-HSPA1B-L (ab154409, Abcam Inc., Cambridge, MA, USA; 1:100), and anti-Hsc70/HSPA8 (ab19136, Abcam Inc.; 1:150) antibodies separately overnight at 4°C. Negative controls were reacted with PBS instead of the specific antibodies. A biotinylated secondary antibody and Vector ABC reagent (Vector Laboratories) were subsequently added according to the manufacturer's instructions. Then, the sections were stained using the FAST DAB Peroxidase Substrate (Sigma) and counterstained with hematoxylin for 10 s. The slides were dehydrated and analyzed under a light microscope.

### 2.8 Prediction of 3D protein structures

The 3D structures of *PsHSP70/110* proteins were predicted for further analysis of protein functions. The 3D protein structures were generated by a homology modeling method using the known homologous structure as a template. The Phyre2 server (Kelley, Mezulis, Yates, Wass, & Sternberg, 2015) was used for homology modeling, secondary structure prediction, and domain analysis. The comparison of 3D protein structures was performed by using PyMOL Viewer software. The prediction of binding sites in proteins was performed using similar structures on the 3DLigandSite server (Wass, Kelley, & Sternberg, 2010).
2.9 | Data analysis

The data were expressed as the means ± SEM. One-way ANOVA was performed using SPSS 16.0 software to assess the differences in gene expression levels. The level of significance was set at a p-value < 0.05.

3 | RESULTS

3.1 | Identification and analysis of Hsp70/110 genes in P. sinensis

In this study, a genomewide search against the P. sinensis genome sequences generated 18 candidate genes belonging to the HSP70/110 family, including 17 Hsp70 genes and one Hsp110 gene (Table 1), and these genes were considered as the PsHSP70/110 family genes of P. sinensis. PsHSP70/110 protein sequences were obtained and used for phylogenetic analysis. According to the genetic distances among the 18 PsHSP70/110 proteins, they were classified into different branches (Figure 1a). Moreover, an analysis of conserved domains showed that all the PsHSP70/110 proteins contained an NBD domain (Table 1; Figure 1b), which was in accord with the typical characteristic of Hsp70/110 proteins. Notably, half of these PsHSP70/110 proteins, such as PsHSPA1A-L, PsHSPA1B-L, PsHSPA12A, and PsHSPA14, exhibited an NBD_sugar-kinase_HSP70_actin (cl17037) domain.

The analysis of amino acid identity showed that members of the same PsHSP70/110 subfamily shared higher sequence similarities and similar domain structures (Figure 1; Table 2). For example, PsHSPA2 and PsHSPA8 shared 89% amino acid identity with each other, and both contained the HSPA1-2_6-8-like_NBD (cd10233) and heat-shock 70-kDa protein (PTZ00009) domains. In addition, PsHSPH1, belonging to the HSP110 proteins, exhibited lower sequence identities with PsHSP70 proteins (less than 40%) with the exceptions of PsHSPA4L, PsHSPA4-X1, and PsHSPA4-X2 (Table 2). Furthermore, analyses of physical and chemical characteristics showed that the molecular weights of most PsHSP70/110 proteins ranged from 50.3 to 77.3 kDa, with the exceptions of PsHSPA4-X1 (94.4 kDa) and PsHSPA12A-L2 (37.2 kDa) (Table S1). The theoretical pI values of most proteins ranged from 5.01 to 5.95, with the exception of PsHSPA2A (7.64). All detailed information on PsHSP70/110 proteins, including their instability index, aliphatic index, and GRAVY, is listed in Table S1.

3.2 | Phylogenetic analysis of Hsp70/110 genes

The homologous Hsp70/110 genes from 12 other species were collected for comparative analysis with the identified PsHSP70/110 genes (Figure 1). The statistics of the Hsp70/110 genes revealed that most species harbored more than 10 Hsp70/110 members, although platypus (O. anatinus), turkey (M. gallopavo), and green sea turtle (C. mydas) exhibited six, eight, and nine members, respectively (Figure 2a). More gene members of the Hsp70/110 family were identified in P. sinensis than in other species. The protein sequences of 18 PsHSP70/110 genes from P. sinensis and 124 homologous genes were used for constructing a phylogenetic tree (Table S2; Figure 2b). The results showed that the 142 Hsp70/110 proteins were grouped into eight distinct clades. The greatest number of Hsp70/110 genes was allocated to the HSPA12 subfamily, which was composed of
HSPA12A, HSPA12B, and their corresponding homologous genes. The lists of PsHSPA4-homologous genes constituted the second largest subfamily. Relatively, few genes belonged to the other subfamilies (Figure 2b). Notably, PsHSPA2 homologs shared the closest phylogenetic relationships with PsHSPA8 homologs, which was consistent with the high similarity between PsHSPA2 and PsHSPA8 in their amino acid sequences (Table 2). HSPA2 and HSPA8 genes were assigned to the same subfamily as HSPA2/8. Compared with the phylogenetic analysis of PsHSP70/110 shown in Figure 1a, the detailed classification shown in Figure 2b was performed to indicate the phylogenetic relationships among PsHSP70/110 genes and the homologous genes in other species.

3.3 | Expression profiling of PsHSP70/110 genes in different tissues of P. sinensis

Tissue-specific expression analysis of PsHSP70/110 genes by qRT-PCR revealed that these PsHSP70/110 genes showed differential expression patterns in different tissues (muscle, heart, blood, brain, kidney, liver, lung, spleen, intestine, testis, and oviduct).
of *P. sinensis* (Figure 3; Table 3). PsHSPA4-X1 and PsHSPA4-X2 exhibited high expression levels in the heart, and PsHSPA4L, PsHSPA12A, PsHSPA12A-L1, and PsHSPA12A-L3 were highly expressed in the blood. Relatively higher expression levels of PsHSPA12B-L and PsHSPA14 were detected in the kidney, and the highest levels of PsHSPA12A-L2 and PsHSPA12A-L4 were found in the lung. Importantly, PsHSPA1A-L, PsHSPA1B-L, PsHSPA2, and PsHSPA8 were highly expressed in the testis, and PsHSPA5, PsHSPA9, PsHSPA13, and PsHSPH1 were highly expressed in the oviduct. The results implied that these PsHSP70/110 genes with specific high expression in reproductive organs (such as the testis and oviduct) may play roles in the reproductive development of *P. sinensis*. In addition, PsHSPA12A-L1, PsHSPA12A-L2, PsHSPA12A-L3, and PsHSPA12A-L4 exhibited no expression in the testis and
oviduct, and PsHSPA1A-L and PsHSPA1B-L were not expressed in the oviduct.

### 3.4 Dynamic expression of PsHSP70/110 genes during spermatogenesis in *P. sinensis*

The dynamic expression analyses of PsHSP70/110 genes revealed that their expression patterns in the testis of *P. sinensis* varied with seasonal temperature changes (different months). A heat map of PsHSP70/110 gene expression based on the FPKM values from RNA-seq data showed that the expression abundances of different PsHSP70/110 genes were extremely diverse in the three different *P. sinensis* libraries (Figure 4; Table 4). The PsHSPA2 gene exhibited the highest expression level, with more than 1,500 FPKM values, whereas the PsHSPA4-X1, PsHSPA12A-L1, PsHSPA12A-L2, PsHSPA12A-L3, and PsHSPA12A-L4 genes were not detected in the three *P. sinensis* testis libraries. Furthermore, qRT-PCR analysis was performed to validate the expression patterns of PsHSP70/110 genes in April, July, and October. Comparative analysis showed that the expression tendency of most PsHSP70/110 genes was coincident between RNA-seq and qRT-PCR analysis (Figure S1). The majority of PsHSP70/110 genes were significantly differentially expressed (*p*-value < 0.05) in July and October compared to April (Figure 5). Moreover, PsHSPA1B-L, PsHSPA2, PsHSPA5, PsHSPA8, PsHSPA9, and PsHSPA13 were significantly upregulated in July and then declined in October, indicating their temperature susceptibility. Additionally, several genes, such as PsHSPA1A-L, PsHSPA4L, PsHSPA12A, PsHSPA14, and PsHSPH1, exhibited the highest expression levels in October.

### 3.5 Interaction network of PsHSP70/110 proteins

A putative interaction network of PsHSP70/110 proteins was proposed, which involved 14 PsHSP70/110 proteins interacting with each other (Figure 6; Table S3). Four proteins (PsHSPA1A-L, PsHSPA12A-L1, PsHSPA12A-L3, and PsHSPA12A-L4) were not found in the network. The protein–protein associations showed that the majority of

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**TABLE 3** The specific primers of PsHSP70/110 genes for qRT-PCR analysis

| Gene name       | Sense primer (5’−3’) | Antisense primer (5’−3’) |
|-----------------|----------------------|--------------------------|
| PsHSPA1A-L      | CTACAAGGGAAAGGAGAA   | CATAGTCTTCAGGCATCT      |
| PsHSPA1B-L      | TTTGGCAAGAAGGAGGAG   | TGGAGTTGAGAAGACAGA      |
| PsHSPA2         | TCTTTGCTTACTACCTCA   | CACTTTGCTTGTCTTTG      |
| PsHSPA4-X1      | GCTCTATATTCTCTCTAA   | GGCTTTCATTCTCATCAG       |
| PsHSPA4-X2      | GCTCTATATTCTCTCTCAA  | GGCTTTCATTCTCATCAG       |
| PsHSPA4L        | TAAGAATGCTTGGAGAAGAT | CTGAACTGCGTCACATA       |
| PsHSPA5         | GTGAGCAGAGAATGCTAT   | ATTCAGACATATTACAT       |
| PsHSPA8         | CTCTGCTTATCAGTAGA    | CGTGTATGGTAGGATTTA      |
| PsHSPA9         | ATACATTCTAGGTTACC    | CTTCATTCTCATAA          |
| PsHSPA12A       | GAACCAATCCACTGAAATA  | CACTAAGATGTTGAATGAT     |
| PsHSPA12A-L1    | ACTACTTCCACCTTCAA    | TCTATTACCTCTGCCTCA     |
| PsHSPA12A-L2    | CTTTGTTTCTGTGCTATT   | AATACATTCTGCTCCTT      |
| PsHSPA12A-L3    | ACAGTACAGAGCAGAAACT  | AAGGCTTCAACACCAGTA     |
| PsHSPA12A-L4    | AAGGAGTATGAGGTAATA   | GGTATAGTGGAGACTA       |
| PsHSPA12B-L     | GGATTACCACCAAGGCTC   | CACCTCTTCCATTCTCAT     |
| PsHSPA13        | ACTCTACACCTTCTCCTC   | TTCAACACCTGCTCAA       |
| PsHSPA14        | AGAGCGATGATGAAAGTTA  | CCTGAGACACATTACACT     |
| PsHSPH1         | GAATGGATGAGTATGCTCTA | ATTTGATGTTCTCTTGA      |
| Psβ-actin       | AGACCAGACAGACTACCTCA | CACCTGACCACAGAAGCAACT  |

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**FIGURE 4** Heat map of PsHSP70/110 gene expression in the testis of *Pelodiscus sinensis* based on RNA-seq analysis
PsHSP70/110 proteins can interact with multiple homologous genes in *P. sinensis*. The blue, purple, and black lines represent the binding, catalysis, and reaction relationships, respectively (Figure 6). The green arrows represent positive interactions with four pairs of PsHSP70/110 proteins. Two proteins, PsHSPA4L and PsHSPA4L, were related to the most proteins, with 11 interacting genes, followed by the PsHSPA4-X1 and PsHSPA4-X2 proteins, with nine interacting genes. Nevertheless, PsHSPA1 and PsHSPA4L were associated with each other only by the

| Gene name       | Gene ID               | FPKM in MT-1 | FPKM in MT-2 | FPKM in MT-3 | Log2 (MT-2/MT-1) | Log2 (MT-3/MT-1) |
|-----------------|-----------------------|--------------|--------------|--------------|------------------|------------------|
| PsHSPA1A-L      | ENSPSIG00000001652    | 6.10         | 2,067.89     | 1,575.81     | 8.40             | 8.02             |
| PsHSPA1B-L      | ENSPSIG00000010036    | 14.86        | 1,655.57     | 1,172.25     | 6.80             | 6.31             |
| PsHSPA2         | ENSPSIG00000001054    | 2,360.52     | 1,815.44     | 2,343.12     | -0.38            | 0.01             |
| PsHSPA4-X2      | ENSPSIG00000013077    | 148.56       | 316.93       | 353.56       | 1.09             | 1.26             |
| PsHSPA4L        | ENSPSIG00000009008    | 456.70       | 846.30       | 943.26       | 0.89             | 1.06             |
| PsHSPA5         | ENSPSIG00000005450    | 470.85       | 285.08       | 474.84       | -0.72            | 0.02             |
| PsHSPA8         | ENSPSIG00000012450    | 415.30       | 638.48       | 446.13       | 0.62             | 0.10             |
| PsHSPA9         | ENSPSIG00000012706    | 266.00       | 169.26       | 169.90       | -0.65            | -0.64            |
| PsHSPA12A       | ENSPSIG00000004375    | 89.51        | 113.45       | 83.67        | 0.34             | -0.09            |
| PsHSPA12B-L     | ENSPSIG00000004539    | 1.35         | 2.19         | 0.01         | 0.70             | -0.70            |
| PsHSPA13        | ENSPSIG00000016198    | 70.71        | 87.39        | 161.55       | 0.31             | 1.20             |
| PsHSPA14        | ENSPSIG00000016234    | 57.48        | 82.66        | 96.88        | 0.52             | 0.76             |
| PsHSPA12A-L     | ENSPSIG00000004375    | 89.51        | 113.45       | 83.67        | 0.34             | -0.09            |
| PsHSPA12B-L     | ENSPSIG00000004539    | 1.35         | 2.19         | 0.01         | 0.70             | -0.70            |
| PsHSPA13        | ENSPSIG00000016198    | 70.71        | 87.39        | 161.55       | 0.31             | 1.20             |
| PsHSPA14        | ENSPSIG00000016234    | 57.48        | 82.66        | 96.88        | 0.52             | 0.76             |
| PsHSPA12A       | ENSPSIG00000004375    | 89.51        | 113.45       | 83.67        | 0.34             | -0.09            |

**TABLE 4** The FPKM values of PsHSP70/110 gene expression in the testis of *Pelodiscus sinensis* determined by RNA-seq analysis.
binding line, and PsHSPA1B-L, PsHSPA2, and PsHSPA8 exhibited the only binding relationships.

3.6 | Functional enrichment of PsHSP70/110 proteins

The protein interaction associations among PsHSP70/110 proteins were predicted in this study and are described above. Furthermore, functional enrichment analysis of PsHSP70/110 proteins showed that five kinds of INTERPRO protein domains and features and one kind of PFAM protein domain were significantly enriched, with \( p \leq 0.05 \) (Table 5). Protein domain enrichments showed that 14 interacting PsHSP70/110 proteins contained the IPR013126 and PF00012 domains, which indicate structural features of Hsp70 proteins. Moreover, the PsHSP70/110 proteins were assigned to four KEGG pathways, including “Protein processing in endoplasmic reticulum” (ID 4141, six genes), “MAPK signaling pathway” (ID 4010, three genes), “Spliceosome” (ID 3040, three genes), and “Endocytosis” (ID 4144, three genes) pathways (Table 5). Remarkably, three proteins, PsHSPA1B-L (ENSPSIG00000011084), PsHSPA2 (ENSPSIG0000001055), and PsHSPA8 (ENSPSIG00000013985), were involved in the MAPK signaling pathway, which provides valuable information for exploring their functions in *P. sinensis*.

3.7 | Protein expression of PsHSPA1B-L, PsHSPA2, and PsHSPA8 in the testis of *P. sinensis* during spermatogenesis

The protein expression analyses of PsHSPA1B-L, PsHSPA2, and PsHSPA8 by IHC in different months (April, July, and October) showed that the positive reactions for the PsHSPA1B-L, PsHSPA2, and PsHSPA8 proteins in the testis of *P. sinensis* were similar within a given month (Figure 7). Weak immunostaining of the three proteins was observed in April, whereas intense immunostaining was observed in July. In addition, the testis displayed moderate immunoreactivity of the three proteins in October. No staining was detected in the negative control sections. The results of IHC analysis were coincident with the mRNA expression variations of PsHSPA1B-L, PsHSPA2, and PsHSPA8 determined by qRT-PCR analysis in response to seasonal temperature changes.

3.8 | Characterization and protein structure analysis of PsHSPA1B-L, PsHSPA2, and PsHSPA8 proteins

An overview of the analysis between PsHSPA1B-L, PsHSPA2, and PsHSPA8 proteins revealed that they exhibited high amino acid identity (Table 2) and similar conserved protein domains (Figure 1) and that they were enriched in the same KEGG pathway of MAPK signaling (Table 5). Prediction of the 3D protein structures of PsHSPA1B-L, PsHSPA2, and PsHSPA8 showed that the three proteins presented very similar structures and binding sites, especially between PsHSPA2 and PsHSPA8 (Figure 8a; Figure S2; Table 6). More detailed phylogenetic analysis showed that PsHSPA1B-L, PsHSPA2, and PsHSPA8 shared the closest relationships with their corresponding homologous genes from western painted turtle (*Chrysemys picta bellii*), green sea turtle (*C. mydas*), and three-toed box turtle (*Terrapene mexicana triunguis*) (Figure 8b–d). Moreover, the mRNA and protein expression patterns of PsHSPA1B-L, PsHSPA2, and PsHSPA8 genes were similar in the testis of *P. sinensis* (Figure 5; Figure 7), which further validated the putative parallel functions of the three genes in certain biological processes. Furthermore, the PsAPAF1 (ENSPSIG00000011998) protein of *P. sinensis*, a potential interacting protein of PsHSP70/110 (Beere et al., 2000), was selected and subjected to interaction analysis. The predicted protein interaction revealed that PsAPAF1 was associated with the PsHSPA1B-L, PsHSPA2, and PsHSPA8 proteins via both binding and catalysis relationships (Figure S2), which implied...
that PsHSPA1B-L, PsHSPA2, and PsHSPA8 may play similar roles and interact with the PsAPAF1 gene in *P. sinensis*.

## 4 | DISCUSSION

### 4.1 | Overview of PsHSP70/110 genes in *P. sinensis*

The central biological roles of Hsp70/110 in various biological and physiological processes are attributed to their chaperone activity and their structurally and functionally conservative properties in evolution (Lindquist & Craig, 2003). Considerable evidence has demonstrated that the protein structure and conserved domains of Hsp70/110 proteins play roles in modulating multiple cellular processes induced by a wide variety of stimuli (Gupta et al., 2010).

In the present study, protein structure analysis showed that 18 PsHSP70/110 proteins contained the conserved NBD domains that are prominent structural features of Hsp70, consistent with others’ observations (20077; Song et al., 2016). Protein functional enrichments also showed that the majority of PsHSP70/110 proteins were significantly enriched in IPR013126 and PF00012 domains and were described as "Heat shock protein 70." Additionally, the analysis of physical and chemical characteristics showed that the approximate molecular weight of most PsHSP70 family proteins was 70 kDa. These findings revealed the reliability of the identified PsHSP70/110 candidates in *P. sinensis* and supported the following studies.

### 4.2 | Conservation of PsHSP70/110 family genes in evolution

Hsp70 is the most conserved heat-shock stress protein in evolution, and its intrinsic functions and conserved domains are responsible for its conservation across species (Kiang & Tsokos, 1998; Murphy, 2013). In this study, a genomewide search identified 18 PsHSP70/110 family members in *P. sinensis*, which is comparable to the number of Hsp70/110 genes in other species. Phylogenetic analysis and classification of Hsp70 proteins from 13 selected animals indicated that some members of this group, such as HSPA12 genes, were ubiquitous in these species. Specifically, PsHSPA1B-L, PsHSPA2, and PsHSPA8 showed the closest phylogenetic relationships with homologs from *C. picta bellii*, *C. mydas*, and *T. mexicana triunguis*, which is concordant with the evolutionary relationships among these species. In addition, a similar number of Hsp70 family genes were found between *P. sinensis* and other species, suggesting the evolutionary conservation of Hsp70 among different species (Song et al., 2016; Wang et al., 2019). Remarkably, HSPA2, HSPA8, and their homologous genes were assigned to the same subfamily of HSPA2/8 and exhibited the closest phylogenetic relationships, which is consistent with previous reports (Song et al., 2016). In general, comparative analysis can provide a foundation for a better understanding of the evolutionary relationships of PsHSP70/110.

In agreement with a previous report by 2016, the family of PsHSP70/110 genes examined in our studies was composed of

| TABLE 5 Functional enrichment of PsHSP70/110 genes in Pelodiscus sinensis |
|-----------------------------|-----------------------------|-------------|-----------------------------|
| Category | Pathway ID | Pathway description | Gene count | False discovery rate | Enriched proteins |
| KEGG Pathways | 4141 | Protein processing in endoplasmic reticulum | 6 | 1.77E-07 | PsHSPA1B-L, PsHSPA2, PsHSPA4L, PsHSPA5, PsHSPA8, PsHSPH1 |
| | 4010 | MAPK signaling pathway | 3 | 3.08E-02 | PsHSPA1B-L, PsHSPA2, PsHSPA8 |
| | 3040 | Spliceosome | 3 | 7.94E-03 | PsHSPA1B-L, PsHSPA2, PsHSPA8 |
| | 4144 | Endocytosis | 3 | 2.08E-02 | PsHSPA1B-L, PsHSPA2, PsHSPA8 |
| INTERPRO protein domains and features | IPR013126 | Heat-shock protein 70 family | 14 | 2.16E-35 | PsHSPA1A-L, PsHSPA1B-L, PsHSPA2, PsHSPA4, PsHSPA4L, PsHSPA5, PsHSPA8, PsHSPA9, PsHSPA12A-L1, PsHSPA12A-L3, PsHSPA12A-L4, PsHSPA13, PsHSPA14, PsHSPH1 |
| | IPR018181 | Heat-shock protein 70, conserved site | 11 | 4.54E-32 | PsHSPA1A-L, PsHSPA1B-L, PsHSPA2, PsHSPA4, PsHSPA4L, PsHSPA5, PsHSPA8, PsHSPA9, PsHSPA13, PsHSPA14, PsHSPH1 |
| | IPR029047 | Heat-shock protein 70 kDa, peptide-binding domain | 10 | 1.01E-28 | PsHSPA1A-L, PsHSPA18B-L, PsHSPA2, PsHSPA4, PsHSPA4L, PsHSPA5, PsHSPA8, PsHSPA9, PsHSPA14, PsHSPH1 |
| | IPR029048 | Heat-shock protein 70 kDa, C-terminal domain | 8 | 2.43E-21 | PsHSPA1B-L, PsHSPA2, PsHSPA4, PsHSPA4L, PsHSPA5, PsHSPA8, PsHSPA9, PsHSPH1 |
| | IPR026685 | Heat-shock 70-kDa protein 12A | 5 | 3.62E-06 | PsHSPA12A, PsHSPA12A-L1, PsHSPA12A-L2, PsHSPA12A-L3, PsHSPA12A-L4 |
| PFAM protein domains | PF00012 | Hsp70 protein | 14 | 9.78E-36 | PsHSPA1A-L, PsHSPA1B-L, PsHSPA2, PsHSPA4, PsHSPA4L, PsHSPA5, PsHSPA8, PsHSPA9, PsHSPA12A-L1, PsHSPA12A-L3, PsHSPA12A-L4, PsHSPA13, PsHSPA14, PsHSPH1 |
PsHSP70 and PsHSP110 proteins. In addition to the difference of one domain between the peptide-binding domain and the C-terminal region, Hsp110 exhibits high homology and a similar crystal structure to Hsp70 (Polier, Dragovic, Hartl, & Bracher, 2008), although Hsp110 is a divergent Hsp70 family member. In this study, one PsHSP110 was discovered, which was named PsHSPH1 in *P. sinensis*. On the basis of its structure and sequence, PsHSP110 was included in the subfamily of PsHSP70, and these proteins were studied and discussed together. Sequence alignment showed that PsHSPH1 shared less than 40% identity with most PsHSP70 proteins. Interestingly, PsHSPH1 presented higher sequence identities with PsHSPA4L, PsHSPA4-X1, and PsHSPA4-X2, and these four members were classified into a closer subgroup by phylogenetic analysis, with similar conserved domains. The observations suggested that PsHSP70/110 family genes with close phylogenetic relationships and similar protein structures may present similar potential roles in *P. sinensis*.

### 4.3 Characterization of PsHSP70/110 gene expression and putative roles

The induction and accumulation of Hsp70/110 are tightly associated with a range of environmental and physical stresses (Gupta et al., 2010; Srivastava, 2002). In this study, tissue-specific expression analysis revealed that most PsHSP70/110 genes were constitutively expressed in different tissues of *P. sinensis*, which suggested that PsHSP70/110 genes may be important for organismic homeostasis. Notably, several genes, such as PsHSPA1A-L, PsHSPA1B-L, PsHSPA2, PsHSPA5, PsHSPA8, and PsHSPA9, exhibited specific high expression in the testis and oviduct of *P. sinensis*. Importantly, the high expression of the PsHSPA2 gene in the *P. sinensis* testis was consistent with previous reports that showed a high level of HSPA2 in the human testis (Daugaard et al., 2007; Son et al., 1999; Su et al., 2004), implying a potential special role of PsHSPA2 in the germ cells of *P. sinensis*. Substantial experimental evidence has revealed that Hsp70 is one of the positive necessary factors for tumor cell survival and, on the contrary, that Hsp70 negatively modulates apoptosis (Goloudina et al., 2012; Rérole et al., 2011). In response to stressful conditions and during diverse developmental processes, apoptosis, which is essential for tissue homeostasis, is conspicuous in multicellular organisms (Meier, Finch, & Evan, 2000). In seasonally breeding species, apoptosis is responsible for testicular atrophy during seasonal reproductive regression (Young & Nelson, 2001). Moreover, apoptosis plays a critical role in seasonal spermatogenesis by eliminating defective germ cells or cells carrying DNA mutations (L2017; Russell, Chiarinigarcia, Korsmeyer, & Knudson, 2001). Seasonal spermatogenesis is characteristic of temperate and boreal reptilian species, including *P. sinensis* (Gribbins, 2011; Liu et al., 2017). Our previous studies examining the testis of *P. sinensis* demonstrated dynamic changes in apoptosis during spermatogenesis and showed that apoptosis was closely correlated with seasonal temperature variation: A mass of apoptotic cells was detected in April (spermatogenically quiescent phase), while decreased apoptosis was observed in July and October (intermediate and late spermatogenesis) (Liu et al., 2017). Furthermore, the constitutive or inducible expression of Hsp70 can interfere with stress-induced apoptosis and is implicated in temperature fluctuations (Dang et al., 2015; Murphy, 2013). Studies in the teleost *Prochilodus argenteus*...
suggested that HSP70 may protect germ cells from apoptosis during breeding cycles, and a decrease in HSP70 expression and increase in apoptosis may facilitate testicular remodeling after the reproductive season (Domingos et al., 2013). Additionally, the targeted disruption of Hsp70 in mice leads to developmental arrest and apoptosis of spermatocytes, resulting in infertility (Dix et al., 1996). In this study, both transcriptomic and qRT-PCR analyses showed that the dynamic expression of most PsHS70/110 genes was related to the temperature range. Remarkably, several genes, such as PsHS1B-L, PsHS2, and PsHS8, were significantly upregulated in both July and October compared with April and exhibited an inverse tendency toward apoptosis during seasonal

**FIGURE 8** Characterization of PsHS1B-L, PsHS2, and PsHS8 proteins. (a) Comparison of the predicted 3D protein structures among PsHS1B-L (green), PsHS2 (blue), and PsHS8 (red), (b–d) Phylogenetic analysis of PsHS1B-L, PsHS2, PsHS8 (in green circles), and their homologs from other species using bootstrap values with 1,000 replications. Ac, Anolis carolinensis; Am, Alligator mississippiensis; Cm, Chelonia mydas; Cp, Chrysemys picta bellii; Dr, Danio rerio; Gg, Gallus gallus; Hs, Homo sapiens; Mg, Meleagris gallopavo; Mm, Mus musculus; Oa, Ornithorhyncus anatinus; Tm, Terrapene mexicana triunguis; Xl, Xenopus laevis
spermatogenesis, which implied that these genes may negatively regulate seasonal apoptosis in *P. sinensis*.

Furthermore, PsHSPA1B-L, PsHSPA2, and PsHSPA8 were implicated in the MAPK signaling pathway in the present study by functional enrichment, indicating that these proteins may function through participating in MAPK signaling. MAPK plays fundamental roles in cellular stress responses and regulates apoptosis through the specific phosphorylation of apoptotic factors, such as p53 (Chang & Karin, 2001; Lee et al., 2006; Taylor, Zheng, Liu, & Thompson, 2013). Apaf1, one of the important components of the p53 signaling pathway, can mediate apoptosis through its association with procaspase-9 (Zou, Li, Liu, & Wang, 1999). Other evidence has strongly suggested that the event of Hsp70 binding to Apaf1 seems to eliminate the oligomerization of Apaf1 and procaspase-9 and then suppresses apoptosis (Beere & Green, 2001; Beere et al., 2000; Saleh, Šrinivasula, Balkir, Robbins, & Alnemri, 2000). As expected, we found protein interactions of PsHSPA1B-L, PsHSPA2, PsHSPA8, and PsAPAF1 with binding relationships. Moreover, IHC analysis showed that the protein expression of PsHSPA1B-L, PsHSPA2, PsHSPA8, and PsAPAF1 with binding relationships. Moreover, IHC analysis showed that the protein expression of PsHSPA1B-L, PsHSPA2, PsHSPA8 exhibited significant upregulation in July and October. More importantly, a distinct decrease in apoptosis in the testis of *P. sinensis* was detected in October (MT-3) (Liu et al., 2017). These observations indicated that the high protein levels of PsHSPA1B-L, PsHSPA2, and PsHSPA8 as well as the interaction with PsAPAF1 may favor the inhibition of apoptosis during spermatogenesis in *P. sinensis*. In addition, in the current study, PsHSPA8, also known as PsHsc70, encoding the predominant cognate member of the PsHSP70 family, was systematically characterized in *P. sinensis* for the first time to the authors' knowledge. Hsc70 is the co-chaperone of the antiapoptotic modulator BAG1, and BAG1 can bind to its ATPase domains to modulate chaperone activities and influence apoptotic responses (Song, Takeda, & Morimoto, 2001; Stuart et al., 1998). Taken together, these findings reveal the potential functions of critical PsHSP70/110 proteins in regulating apoptosis and associated with spermatogenesis in *P. sinensis*.

## 5 | CONCLUSION

In this study, a total of 18 PsHSP70/110 family genes were identified and comprehensively analyzed in *P. sinensis*. All the PsHSP70/110 proteins contained the conserved NBD domain, which represents a typical structural feature of Hsp70. Classification and phylogenetic analysis of PsHSP70/110 and their homologs demonstrated that Hsp70/110 was conserved in evolution across various species. Moreover, the expression profiling of these PsHSP70/110 genes varied considerably in different tissues of *P. sinensis*. Notably, several genes, such as PsHSPA1B-L, PsHSPA2, and PsHSPA8, were significantly differentially expressed in the testis of *P. sinensis* in different months. Functional enrichments showed that PsHSPA1B-L, PsHSPA2, and PsHSPA8, with putative interaction relationships, were involved in the MAPK signaling pathway. Further analysis indicated that the high protein expression of PsHSPA1B-L, PsHSPA2, and PsHSPA8 and their binding to PsAPAF1 may account for the decrease in apoptosis during spermatogenesis in *P. sinensis*, which suggested a putative vital role of critical PsHSP70/110 in regulating apoptosis. The findings of this study will provide insights into the potential functional roles of PsHSP70/110 in *P. sinensis*.

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CONFLICTS OF INTEREST

The authors declare that they have no competing interests.

AUTHOR CONTRIBUTIONS

The authors have made the following declarations about their contributions: TL and HZ conceived and designed the study. TL, YH, and YL analyzed the data and performed the experiments. TL wrote the manuscript. TL and HZ revised the manuscript. All authors read and approved the final manuscript.

DATA ACCESSIBILITY

Transcriptomic data: NCBI SRA: SRX2351846 (MT-1), SRX2352135 (MT-2), and SRX2352136 (MT-3).

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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