Characterization of histone deacetylases and their roles in response to abiotic and PAMPs stresses in *Sorghum bicolor*

Qiaoli Du¹†, Yuanpeng Fang¹†, Junmei Jiang²†, Meiqing Chen¹, Xiaodong Fu¹, Zaifu Yang¹, Liting Luo¹, Qijiao Wu¹, Qian Yang¹, Lujie Wang¹, Zhiguang Qu¹, Xiangyang Li² and Xin Xie¹*  

Abstract  

**Background:** Histone deacetylases (HDACs) play an important role in the regulation of gene expression, which is indispensable in plant growth, development, and responses to environmental stresses. In *Arabidopsis* and rice, the molecular functions of HDACs have been well-described. However, systematic analysis of the HDAC gene family and gene expression in response to biotic and abiotic stresses has not been reported for sorghum.  

**Results:** We conducted a systematic analysis of the sorghum HDAC gene family and identified 19 SbHDACs mainly distributed on eight chromosomes. Phylogenetic tree analysis of SbHDACs showed that the gene family was divided into three subfamilies: RPD3/HDA1, SIR2, and HD2. Tissue-specific expression results showed that SbHDACs displayed different expression patterns in different tissues, indicating that these genes may perform different functions in growth and development. The expression pattern of SbHDACs under different stresses (high and low temperature, drought, osmotic and salt) and pathogen-associated molecular model (PAMPs) elf18, chitin, and flg22) indicated that SbHDAC genes may participate in adversity responses and biological stress defenses. Overexpression of SbHDA1, SbHDA3, SbHDT2 and SbSRT2 in *Escherichia coli* promoted the growth of recombinant cells under abiotic stress. Interestingly, we also showed that the sorghum acetylation level was enhanced when plants were under cold, heat, drought, osmotic and salt stresses. The findings will help us to understand the HDAC gene family in sorghum, and illuminate the molecular mechanism of the responses to abiotic and biotic stresses.  

**Conclusion:** We have identified and classified 19 HDAC genes in sorghum. Our data provides insights into the evolution of the HDAC gene family and further support the hypothesis that these genes are important for the plant responses to abiotic and biotic stresses.  

**Keywords:** Genome analysis, Histone deacetylases, Expression profile, Abiotic stress, Pathogen-associated molecular model (PAMPs), *Sorghum bicolor*

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**Background**  
Epigenetic regulation is an important regulatory mechanism that helps plants adapt to environmental stresses. Moreover, epigenetic regulation of gene expression proceeds through DNA or histone modification without changing the DNA sequence, which is fast, reversible, and heritable [1]. Chromatin remodeling, an important type of epigenetic regulation, is one of the key regulators of gene expression in higher plants. It affects various
cellular processes by regulating gene expression in different growth and development stages [2]. Reports have shown that different environmental stresses can cause different types of modifications in histones [3]. For example, histone post-translational modifications mainly include histone acetylation, ubiquitination, phosphorylation, ADP-ribosylation, and methylation [4]. Among these, histone acetylation has attracted the most attention, and histone acetylation modification plays a vital role in the regulation of eukaryotic transcription activity. Histone acetylation mainly occurs on lysine residues at histone tails. It is a reversible dynamic equilibrium process, mainly controlled by histone acetyltransferases (HATs) that catalyze histone acetylation and histone deacetylases (HDACs) that control the co-regulation of histone deacetylation. HATs transfer the acetyl group of the acetyl-CoA to lysine residues at the end of histones to eliminate the positive charge and force the chromatin structure into a more elongated state, which is beneficial for transcription factor binding and is related to transcriptional activation of genes. Meanwhile, HDACs remove the acetyl groups from the ends of histones, leaving chromatin in a tighter, more condensed state, which is not conducive to the binding of transcription factors or transcriptional regulators to DNA, and is associated with transcriptional suppression/silencing [5]. Thus, histone modification plays an important role in the regulation of gene expression; HATs promote gene expression, and HDACs inhibit gene expression.

Histone acetylases (HATs) are divided into five families based on sequence characteristics: GNAT, MYST, p300/CREB binding protein (CBP) coactivator, TAFII250, and HATs. HDACs in plants are divided into three subfamilies: RPD3/HDA1, SIR2, and HD2 [6], the first two of which are homologous to the yeast RPD3/HDA1 and SIR2 families, and RPD3/HDA1 is the largest subfamily of the HDAC family. These HDACs are mainly distributed in the nucleus or cytoplasm, or they shuttle between the nucleus and cytoplasm [7]. Members of the SIR2 family are conserved from prokaryotes to eukaryotes, and use Nicotinamide adenine dinucleotide (NAD+) as a coenzyme to regulate the activity of HDACs [8]. The HD2 family is unique to plants and has not been found in yeasts or animals [9, 10]. In Arabidopsis, HD2 family members have a conserved amino acid region (EFWG motif) at the N-terminus, and both HDT1 and HDT3 contain a C2H2 zinc finger domain that may mediate DNA-protein or protein-protein interactions [11]. All members of this family contain a typical histone deacetylase domain, the enzymatic activity of which requires the presence of Zn^{2+}. A large number of studies have shown that in the human body, HDACs are closely related to the occurrence and development of cancer [12]. In 1988, HDACs were also found in plants [13]. Histone acetylation and deacetylation play an important role in the growth and development of plants, including root development [14], flower development [15], gametophyte development [16], and cell proliferation during organ growth [17]. They also participate in plant responses to changes in the external environment, such as light signals [18], salt stress and abscisic acid (ABA) signaling [19], cold stress [20], heat stress [21] and other hormone signaling pathways [22].

Members of the HDAC gene family have been identified in Arabidopsis [23], rice [24], corn [9], tomato [25], cotton [26], tea [27], and other plants. Eighteen HDAC members have been identified in the Arabidopsis genome, all of which belong to the above three HDAC families. Increasing evidence shows that in the response of Arabidopsis to biological and abiotic stresses, HDACs play a vital role in the process of epigenetic regulation. For example, AtHDA6 and AtHDA19 function in the abscisic acid (ABA) signaling pathway, and can also be induced by jasmonic acid (JA) and ethylene, while HDA6 interacts with EIN3 and HDA proteins in ethylene and JA signaling [28]. AtHDA19 inhibits WRKY38 and WRKY62 transcription factors and regulates the expression of Pathogenesis Related 1 (PR1) by participating in the plant defense response mediated by salicylic acid (SA) [29]. Overexpression of AtHD2C in Arabidopsis leads to increased tolerance to salt and drought stress [30]. HD2 family members also play a cardinal role in plant growth and development. For example, silencing the HD2A gene in Arabidopsis causes seed development to cease [31], while overexpression of HD2A alters leaf and flower morphology [11], defects, delayed flowering, and suspension of seed development. Expression of HD2A, HD2B, HD2C, and HD2D is inhibited by ABA and sodium chloride [19]. In rice, HD17705 participates in the regulation of seed germination in response to abiotic stress [32]. Research on HDAC genes in rice showed that overexpression of HDT701 in transgenic rice leads to increased susceptibility to rice blast (Magnaporthe oryzae) and bacterial blight (Xanthomonas oryzae pv oryzae) (Xoo), but silencing of the HDT701 gene increases resistance to these two pathogens, and stimulates the production of reactive oxygen species (ROS) under the action of the PAMP elicitors flg22 and chitin. The level of histone H4 acetylation has a clear regulatory effect; it negatively regulates the transcription of defense-related genes, indicating that HDT701 regulates the level of histone H4 acetylation of pattern recognition receptors and defense-related genes [33]. SIR2 proteins are also important in the response to pathogen
infection. For example, the SIR2 family protein AtSRT2 in *Arabidopsis* plays a negative regulatory role in the basic defense of plants against the pathogen PstDC3000, and expression of the AtSRT2 gene is inhibited under pathogen infection [34]. In rice, the OsSRT1 gene inhibits the expression of starch metabolism–related genes in seeds [35]. Reports showed that different HDAC family members have been found to mediate different aspects of plant growth and development. They also play a role in biological and abiotic stress responses, but the characteristics of HDAC genes in sorghum have not yet been reported.

In the present study, we identified 19 HDAC-encoding genes in sorghum. Using bioinformatics, we systematically analyzed phylogenetic relationships, conserved domains, and motifs of the sorghum HDAC gene family, and we performed real-time fluorescence quantitative PCR (qRT-PCR) to assess the responses of the 19 genes in different tissues and under different stresses. Changes in acetylation in sorghum seedlings were assessed under adverse stress conditions (low and high temperature, NaCl, PEG 6000). In addition, we also performed prokaryotic expression analysis of SbHDA1, SbHDA3, SbHDT3, and SbSRT2 genes, and carried out spot experiments under heat, salt, osmotic and drought stresses, to verify whether expression of these genes in *E. coli* could enhance stress tolerance. The findings lay a molecular foundation for further exploring the functions of sorghum HDAC genes in sorghum resistance breeding.

**Results**

**Genome-wide identification of HDAC genes in Sorghum bicolor**

We performed a comprehensive identification of HDAC genes in sorghum genome, and removed the redundant HDAC genes based on conserved domains. Totally, 19 HDAC protein sequences containing conserved histone deacetylase domains were obtained. Bioinformatics analysis showed that genes encoded proteins ranging from 269 (SbHDT1) to 703 (SbHDA2) amino acid residues. The coding sequences of the sorghum SbHDAC gene family range from 834 bp (SbHDT2) to 1905 bp (SbHDA1), the molecular weight of the corresponding proteins ranges from 29.54679 (SbHDT1) to 45.30 (SbHDA2) kDa, and the pI of most SbHDACs proteins is < 7.0, making them acidic at neutral pH (Table 1), except for SbHDT2 (pI = 7.92), SbSRT1 (pI = 8.88), and SbSRT2 (pI = 9.17). The 19 HDAC genes are distributed on nine chromosomes; chromosome 3 contains the largest number of HDAC genes (4), followed by chromosomes 9 and 10 (3), chromosomes 2, 4, and 6 each contain 2, chromosomes 5, 7, and 8 each contain 1, and chromosome 1 does not contain any HDAC genes (Fig. S1).

### Table 1  Identification of SbHDAC gene family members

| Gene Name | Accession Number | Chr | Range | CDS (bp) | Protein (aa) | Molecular weight (kDa) | PI | Subcellular localization |
|-----------|-----------------|-----|-------|----------|--------------|------------------------|----|------------------------|
| SbHDA1    | LOC110432512    | 2   | 4,103,729~4,115,614 | 1905 | 634 | 68.29599 | 5.68 | Nucleus |
| SbHDA2    | LOC8060653      | 2   | 72,626,015~72,634,000 | 2112 | 703 | 76.72727 | 5.78 | Chloroplast. Nucleus. |
| SbHDA3    | LOC8079300      | 3   | 52,802,297~52,806,221 | 1404 | 467 | 51.90450 | 5.78 | Nucleus |
| SbHDA4    | LOC8085432      | 3   | 66,687,911~66,692,549 | 1203 | 430 | 46.69788 | 6.13 | Nucleus |
| SbHDA5    | LOC8082952      | 4   | 7,759,256~7,763,600 | 1551 | 516 | 57.75734 | 5.29 | Nucleus |
| SbHDA6    | LOC8085534      | 4   | 8,149,916~8,155,266 | 1059 | 352 | 38.76319 | 5.5 | Chloroplast. Nucleus. |
| SbHDA7    | LOC8056459      | 5   | 10,412,638~10,414,364 | 1077 | 358 | 38.86902 | 9.08 | Nucleus |
| SbHDA8    | LOC8073075      | 6   | 42,785,502~42,802,511 | 1344 | 447 | 50.82971 | 6.55 | Cytoplasm. Nucleus. |
| SbHDA9    | LOC8065827      | 9   | 50,989,573~50,993,878 | 1173 | 390 | 42.25157 | 5.49 | Nucleus |
| SbHDA10   | LOC8085527      | 9   | 50,989,573~50,993,878 | 1173 | 390 | 42.25157 | 5.49 | Nucleus |
| SbHDA11   | LOC8079300      | 10  | 50,823,203~50,827,657 | 1053 | 350 | 38.61221 | 6.01 | Chloroplast. Nucleus. |
| SbHDA12   | LOC8065038      | 10  | 51,422,040~51,430,431 | 1557 | 518 | 57.76654 | 5.48 | Cytoplasm. Nucleus. |
| SbHDT1    | LOC8078709      | 3   | 70,646,814~70,650,720 | 810 | 269 | 29.54679 | 5.56 | Nucleus |
| SbHDT2    | LOC8078710      | 3   | 70,659,296~70,662,320 | 834 | 277 | 29.70739 | 5.02 | Nucleus |
| SbHDT3    | LOC8068982      | 9   | 55,768,231~55,771,439 | 900 | 299 | 32.30415 | 4.64 | Nucleus |
| SbHDT4    | LOC8065675      | 9   | 59,367,712~59,370,487 | 924 | 307 | 33.37051 | 4.80 | Nucleus |
| SbHDT5    | LOC8065099      | 10  | 58,010,970~58,014,442 | 1461 | 486 | 52.29158 | 7.92 | Nucleus |
| SbSRT1    | LOC110436344    | 6   | 48,919,950~48,935,447 | 1431 | 476 | 53.23976 | 8.88 | Chloroplast |
| SbSRT2    | LOC8076304      | 8   | 6,041,680~6,047,145 | 1455 | 484 | 53.66489 | 9.17 | Chloroplast |

E-value ≤ 1 × 10^{-5} was used to identify the HDAC gene family members in sorghum as the minimum threshold.
Phylogenetic analysis and sequence alignment of SbHDACs

Using the amino acid sequences of HDACs in *Arabidopsis*, rice, and tomato, together with HDACs in sorghum, MEGA7.0 was employed to construct a phylogenetic tree of the HDAC gene family (Fig. 1). The HDAC sequences of sorghum are clustered into three subfamilies: RPD3/HDA1, SIR2, and HD2. Among them, RPD3/HDA1 has the most member (12). Based on sequence similarity, this subfamily can be divided into RPD3/HDA1-Class1, RPD3/HDA1-Class2, and RPD3/HDA1-Class3, each group contains 6, 2 and 4 sorghum SbHDAC members, respectively. All members of the RPD3/HDA1 subfamily share similarity with the HDAC domain of sorghum; only two SIR2 members containing the SIR2 domain are found in sorghum. In addition, sorghum HD2 contains five members with a C2H2 zinc finger domain that recognize and bind DNA, indicating that they may bind DNA or mediate protein-protein interactions. DNAMAN7.0 software was used to compare the amino acid sequences of members of the sorghum SbHDAC family. The results showed that the overall sequence identity of the 19 SbHDAC proteins was only 16.59% (Fig. S2).

Collinearity analysis of the SbHDACs gene family

Collinearity analysis of the sorghum HDAC gene family identified only one pair of collinearity gene (*SbHDA5/SbHDA11*) pair in sorghum genome, indicating a replication event may have occurred in the recent past. Besides, this event did not cause significant amplification of HDACs, or significant gene loss after genome duplication event (Fig. 2).

Gene structure and conserved motifs of SbHDACs proteins

In order to further explore the structural characteristics of the sorghum HDAC gene family, we analyzed intron-exon numbers and conserved motifs of all members. The results showed that the number of introns and exons in *SbHDAC* genes was 0-15 and 1-18, respectively (Fig. S3). And there are almost no differences between members of the HD2 subfamily. The more closely related genes in the phylogenetic tree

![Phylogenetic relationship of HDAC gene family among sorghum, tomato, rice, and Arabidopsis](image)

*Fig. 1* Phylogenetic relationship of HDAC gene family among sorghum, tomato, rice, and *Arabidopsis*. Multiple sequences alignment and phylogenetic tree were constructed by MEGA7.0 and the bootstrap test was performed with 1000 iterations. The five groups are indicated with camber lines.
appeared to have similar structural components, hence it can be concluded that genes in the same subgroup may perform similar functions.

In order to further analyze the diversity of sorghum SbHDACs, 30 conserved motifs were predicted using MEME online software. In general, proteins clustered in the same subfamily share similar motifs, indicating that family protein members in the same subfamily may have similar functions. As can be seen in Fig. S4, all 12 members of the HDA contain motifs 1 and 5; the five members of the HD2 subfamily all contain motif 7, motif 9, motif 16, and motif 19; the two members of the SIR2 subfamily SbSRT1 and SbSRT2 both contained motif 29. In addition, motif 12 was only present in SbHDA4, SbHDA5, SbHDA8, SbHDA9, and SbHDA12. Thus, it can be inferred that these genes may have some special functions, but the conserved motifs of the entire HDAC family were quite different, which may be due to the different functions of family members.

**Putative Cis-acting regulatory elements in the promoter region of SbHDACs genes**

In order to obtain the cis-acting regulatory elements of the SbHDAC gene family, we analyzed the sequence of the putative promoter region of each SbHDAC gene, and identified 30 putative cis-acting regulatory elements. The results showed that sorghum mainly had the core cis-acting elements TATA and CAAT, as well as some elements related to stress, development, and plant hormone responses such as MYB binding site (MBS), MYC elements (involved in drought tolerance), LTR elements and DRE (involved in low temperature responses), ABRE elements (related to the ABA signaling pathway), P-box elements (related to gibberellin responses), and DRE1 (related to salt stress). The promoter region of each SbHDAC gene differed in the type and number of regulatory
cis-acting elements (Fig. 3, Table S2). Although RPD3/HDA1, HD2, and SIR2 share most of the cis-regulatory elements in their promoters, some cis-elements are missing in certain subfamilies. For example, LTR cis-acting elements are only present in the promoter regions of SbHDA2, SbHDA3, SbHDA7, and SbHDA11. MBS was found in the promoter regions of SbHDA2, SbHDA3, SbHDA5, SbHDA9, SbHDA11, SbHDA12, SbHDT3, and SbSRT2. The drought response-related element MYB is present in all HDACs. This indicates that different members of the SbHDAC gene family may be involved in different abiotic stresses.

Expression pattern of SbHDACs genes in different tissues
We explored the expression patterns of SbHDAC family genes in different tissues including roots, stems, leaves, buds, and seeds. It can be seen from Fig. 4 that these 19 SbHDAC genes were successfully detected in five tissues. Nine genes (SbHDA1, SbHDA1, SbHDA2, SbHDA3, SbHDA4, SbHDA5, SbHDA6, SbHDA12, SbHDT1, and SbHDT2) were expressed at high levels in leaves, and relatively low levels in roots, stems, leaves, and seeds. SbHDT3 was expressed in roots, while SbHDA8, SbHDA10, and SbHDA11 were highly expressed in stems. SbHDA7 was most highly expressed in buds. Unlike the other 18 SbHDAC genes, SbSRT2 is expressed in roots and stems, but expression levels in leaves and buds were relatively low and comparable. These results showed that the 19 SbHDAC genes were constitutively expressed in five tissues of sorghum, and most were expressed in root, stem and leaf tissues.

Expression of SbHDACs in response to phytohormone
In order to explore the hormone responses of SbHDAC gene family members, qRT-PCR was used to analyze the expression patterns of sorghum treated with ABA. The results showed that SbHDA1, SbHDA2, SbHDA3, SbHDA4, SbHDA8, SbHDA9, SbHDA10, SbHDA11, SbHDA12, and SbHDT4 were significantly induced by ABA, with maximum gene expression levels mainly concentrated at 6 and 12 h for SbHDA5, SbHDA6, SbHDA7, SbHDT1, SbHDT2, SbHDT3, and SbHDT5 (Fig. 5). Compared with the control group, SbSRT2 was significantly inhibited under ABA treatment, indicating that ABA could activate and inhibit the expression of different SbHDACs genes.
Expression of SbHDACs genes under abiotic stresses

Figure 6 shows the expression patterns of SbHDACs under low temperature (4°C), high temperature (40°C), PEG6000 (drought stress), D-mannitol (osmotic stress), and NaCl stresses (salt stress). In response to low temperature, SbHDA3, SbHDA4, SbHDA6, SbHDA7, SbHDT2, SbHDT3, SbHDT4, and SbHDT5 were significantly inhibited; meanwhile, SbHDA12 and SbSRT1 reached maximum expression levels after 12 h of continuous treatment, and the relative expression level of SbSRT1 was about 12-fold higher than that of the internal control gene, indicating that these two genes respond to low temperature stress. Most of the SbHDAC genes in sorghum were significantly inhibited under high temperature, except for SbHDA1, SbHDA2, SbHDA6, SbHDT1, SbHDT2, and SbHDT4. After NaCl treatment, expression of the 19 genes was significantly suppressed, except for SbHDA3. Under PEG 6000, D-mannitol drought and osmotic stresses, most genes were suppressed. These results showed that sorghum HDAC genes acted differently in response to adversity stresses.

Expression of SbHDACs genes in response to pathogen-associated molecular patterns (PAMPs)

PAMP elicitors chitin, flg22, and elf18 were used to simulate biological stress, and the expression of SbHDACs was measured to evaluate the effects under PAMPs stress (Fig. 7). It was found that after chitin treatment, expression of SbHDA8 and SbHDA9 genes was increased significantly, and reached a maximum at 3h. Expression of SbHDA12, SbHDT1, SbHDT2, SbHDT3, and SbHDT4 initially increases, then decreased, and reached to the peak at 6h. SbHDA2, SbHDA5, SbSRT1, and SbSRT2 reached maximum expression levels after 12h of continuous treatment, indicating that they were relatively slow in responding to chitin treatment. By contrast, after chitin treatment, expression of SbHDA1, SbHDA3,
and SbHDA4 was suppressed. Meanwhile, SbHDA3, SbHDA4, SbHDA7, SbHDA9, SbHDA10, SbHDA11, SbSRT1, and SbSRT2 were significantly induced by elf18. Under flg22 treatment, the expression patterns of most HDACs revealed an initial increase followed by a decrease in expression level. The above results indicated that chitin, flg22, and elf18 could activate and inhibit the expression of SbHDACs, which implied that SbHDACs were involved in different plant innate immune response processes.

Functional verification of SbHDACs: prokaryotic expression and spot assays of E. coli expressing SbHDA1, SbHDA3, SbHDT3, and SbSRT2 under abiotic stresses

To confirm the function of SbHDACs, SbHDA1, SbHDA3, SbHDT3 and SbSRT2 were amplified and inserted into pET28a for prokaryotic expression (Fig. S5). The coomassie brilliant blue staining results showed that SbHDA1, SbHDA3, SbHDT3, and SbSRT2 proteins were solubly expressed in E. coli when induced at 16 °C, 25 °C or 30 °C (Fig. 8), and western blot results

(See figure on next page.)

Fig. 5 Expression pattern of SbHDAC family genes in response to abscisic acid. Different lowercase letters indicate a significant difference determined by the Duncan's new multiple range test (P-value < 0.05); The reference gene is SbEIF4a; The y-axis value represents the relative expression and abscisic acid (ABA, 200 μM)

Fig. 6 Expression pattern of SbHDAC family genes in response to low temperature, high temperature, drought, osmotic and salty stresses. Different lowercase letters indicate a significant difference determined by the Duncan's new multiple range test (P-value < 0.05); The reference gene is SbEIF4a; The y-axis value represents the relative expression. Low temperature: 4 °C; High temperature: 40 °C; Drought and osmotic stress: 20% PEG6000 and 300 mM D-mannitol; Salty stress: 250 mM NaCl
Fig. 6 (See legend on previous page.)
Fig. 7 Expression pattern of ShHDAC family genes in response to PAMPs. Different lowercase letters indicate a significant difference determined by the Duncan's new multiple range test (P-value < 0.05); The reference gene is SbEIF4a; The y-axis value represents the relative expression. Flagellin (flg22, 100 nM), translation elongation factor (elf18, 100 nM) and chitin (chitin, 8 nM)
showed that the expressed protein is HDAC proteins (Fig. S6). Subsequently, the growth of recombinant bacteria (with pET-28a-SbHDA1, SbHDA3, SbHDT3 or SbSRT2) and control (pET-28a) were further examined under different stresses (PEG 6000, NaCl and D-mannitol) on the LB medium (Fig. 9). The spot growth results showed that after 12h of culture, bacteria harboring SbHDA1, SbHDA3, SbHDT3 or SbSRT2 gene showed higher growth rate than with empty vector under drought (20% PEG 6000) and salt stress (NaCl) (Fig. 9E-G). Meanwhile, all the recombinant bacteria showed the same growth rate under normal condition (Fig. 9A-D). However, under D-mannitol stress, the growth rate of recombinant bacteria was almost insignificant compared with control (Fig. 9F). All the results demonstrated that SbHDA1, SbHDA3, SbHDT3 and SbSRT2 may contribute the tolerance of E. coli under PEG 6000 and NaCl stress.
**Sorghum histone acetylation level are upregulated under stresses treatment**

Additionally, Pan-acetyl lysine antibody was used to examine the acetylation level of sorghum in response to different stresses (cold, heat, osmotic and salt stresses). As shown in Fig. 10, multiple lysine acetylated protein bands were detected in all samples. However, compared with the control group, stronger acetylation level was observed following the stress treatments (Fig. 10B), while all the proteins were loaded in the same amount (Fig. 10A).

**Discussion**

Epigenetics mainly refers to the study of heritable gene expression changes that do not involve changes in DNA sequence. It plays a very important role in the growth and development of plants, and histone modification is closely related to gene expression regulation. HDACs, also called lysine deacetylases, are enzymes that regulate gene expression by removing acetyl groups from core histones (H2A, H2B, H3, and H4) [36]. Studies have shown that HDACs play a key role in plant growth and development [7], including genome stability [3], and responses to various environmental stresses [5]. Members of the HDAC family have been widely reported in a variety of plants. However, research on HDAC genes in sorghum is scarce. Therefore, in this study we conducted a comprehensive genomic analysis of the sorghum *SbHDAC* gene family, and the results provided a strong theoretical basis for future functional studies.

The involvement of HDACs in the responses to environmental cues has not been documented in sorghum. Herein, 19 HDACs were identified in the sorghum genome, and characterized in terms of tissue-specific expression profiles, biotic and abiotic stress response expression patterns, prokaryotic expression, and acetylation levels. The 19 SbHDACs belong to three subfamilies: RPD3/HDA1, SIR2, and HD2. The number of SIR2 subfamilies in sorghum is similar to that in *Arabidopsis*, rice, and tomato, all of which have two SIR2 genes, while soybean contains four SIR2 genes [37]. All members of the RPD3/HDA1 subfamily have a conserved HDAC domain, while members of the SIR2 and HD2 subfamilies have SIR2 and C2H2 Zinc finger domains, similar to those reported in *Arabidopsis* [23], *Oryza sativa* [24], and *Zea mays* [38]. Genes belonging to the same subfamily may share similar structures. All *SbHDACs* contain a variety of conserved motifs that are highly similar to those in
other plants, which strongly indicates that SbHDACs may have similar functions to homologous genes.

Previous reports showed that RPD3 type Class I HDACs are localized exclusively in the nucleus in humans, whereas class II HDACs are shuttled between the cytoplasm and the nucleus [39]. In Arabidopsis, previous studies demonstrated that RPD3 type Class II HDACs HDA5, HDA8, and HDA14 are localized in the cytoplasm, whereas HDA15 is localized exclusively in the nucleus. In addition, AtHDA15 was shown to shuttle from the cytoplasm to the nucleus in response to light [18]. In soybean [37], GmHDA6, GmHDA13, GmHDA14, and GmHDA16 are located in the nucleus and cytoplasm, while in sorghum, SbHDA8, SbHDA9, and SbHDA12 are also located in the nucleus and cytoplasm, and SbHDA2, SbHDA6, and SbHDA11 are also located in the chloroplast and nucleus, suggesting a possible shuttling process between these compartments. The two members of the soybean HD2 subfamily are localized in the nucleus, while in the present study, five members of the sorghum HD2 subfamily were found to be localized in the nucleus. SIR2 proteins are reported to occupy discrete subcellular compartments in plants. For example, in rice, OsSRT1 is found in the nucleus [40], and OsSRT2 and AtSRT2 are found in mitochondria [41], but SISRT2 is localized in both the nucleus and cytoplasm [42]. In the present study, we predicted that SbrSRT1 and SbrSRT2 are localized in the chloroplast. Overall, the subcellular localization patterns of different genes in sorghum indicates that they might be differentially regulated and may have distinct roles.

Increasing evidence indicates that responses of HDACs to the environment stress play a key role in plant growth and development [5]. For example, in Arabidopsis, the ATHDAC protein reportedly participates in responses to environmental stresses such as salt, drought, and temperature. It also participates in seed development, senescence, embryonic development, photomorphogenesis, senescence, and flowering processes [43–45]. In rice, HDACs are involved in flowering [46], root development [47], seed germination and responses to environmental stress [48]. In tomato, SIHDA1, SIHDA3, SIHDA4, and SIHDA5 are involved in the response to different abiotic stresses [49]. The PtHDT902 gene in poplar has a strong influence on the formation of the root system, and salt tolerance of poplar has a negative regulatory role [50]; in barley, the HDA1 gene plays a vital role in development and epigenetic regulation [51]. In sorghum, SbHDAC genes belonging to RPD3/HDA1, HD2, and SIR2 groups displayed higher transcription levels in five different tissues. However, in tea plants, only RPD3/HDA1 and HD2 subfamily genes were expressed at high levels in different tissues, while SIR2 subfamily transcription levels were very low, indicating that HDAC genes are expressed differently in species. Under salt, simulated drought, and temperature stress, the expression of most of genes in sorghum was suppressed, consistent with the results reported in Arabidopsis [30], barley [52], and rice. In sorghum, SbHDA3 was up-regulated, and research on Arabidopsis indicates that overexpression of CsHD2C can enhance the sensitivity to ABA and NaCl stress, but whether HDA3 has the same function in sorghum requires further verification. In addition, under treatment with exogenous ABA, the expression of most SbHDAC genes was up-regulated, similar previous results in tea plants [27]. In our current study, SbHDA1, SbHDA3, and SbHDA8 were down-regulated following NaCl treatment and induced by application of ABA, consistent with previous findings. In addition, the NtHD2s helps tobacco to improve the adaptability against salt stress [53]; In rice [32], through yeast two-hybrid screening analysis, it was found that HDA705 can interact with Hsf B1 family protein (RHSF10), and salt responsive WD40 protein (SRWD) to regulate stress response. The above results indicate that sorghum SbHDACs could respond to different abiotic stresses.

Conclusions

In present study, we identified 19 HDACs genes from Sorghum bicolor, which were divided into three subfamilies: RPD3/HDA1, SIR2 and HD2. The cis-acting elements and real-time PCR results indicate that HDACs genes play important roles in participating in stress resistance. Furthermore, sorghum has undergone significant changes in its acetylation level under adversity treatments. Therefore, our research provides help to understand the HDACs gene of sorghum and lay a solid foundation for the improvement of other crops.

Methods

Plant materials, growth conditions, and stress treatments

After surface disinfection, the germinated sorghum Bx623 seeds were planted in sterilized soil (Pindstrup, Denmark) and cultivated in a greenhouse at 25/20°C under a 14 h light/10 h dark cycle. When the seedlings grow to the three-leaf stage, abscisic acid (ABA, 200 μM), 20% PEG 6000, mannitol (D-mannitol, 300 mM) and sodium chloride (NaCl, 250 mM) were sprayed to the seedlings separately. For temperature treatments, sorghum seedlings were kept in a constant temperature incubator at 4°C or 40°C, and 25°C was as the control. In response to biological stresses, the seedlings were sprayed with PAMPs such as flg22 (100 nM), elf18 (100 nM) or chitin (8 nM); samples were collected at 0, 3, 6, 9 and 12 h post treatments. Sorghum tissues (roots, stems, leaves, buds and seeds) were sampled under...
normal condition. All the samples, three biological replicates were set up, and three seedlings were processed in each replicate. The samples were quickly frozen with liquid nitrogen and stored in a refrigerator at −80°C for further use in RNA extraction. All methods were performed in accordance with the protocols set up based on the relevant guidelines and regulations.

Identification of HDAC genes in Sorghum bicolor

The whole genome sequence of sorghum was downloaded from NCBI (https://www.ncbi.nlm.nih.gov/) as the local database. The hidden Markov model (HMM) configuration files PF00850 (Hist_deacetyl domain), PF02146 (SIR2 domain) and PF17800 (NPL domain) of the HDAC family were extracted from the Pfam database (http://pfam.sanger.ac.uk). Then, the HMM configuration files were used to search for target sequences with conserved domains in the local sorghum protein database through HMMER 3.0 (http://hmmr.janelia.org/) with the E-value ≤1 × 10⁻⁵. The basic physical and chemical properties of sorghum SbHDAC family proteins are predicted with ExPASy-ProtParam tool (https://web.expasy.org/protscale/). Protein subcellular localization was predicted through Cell-PLoc 2.0 (http://www.cbsbio.sjtu.edu.cn/cgi-bin/PlantPLoc.cgi). Chromosome location and gene structure were separately performed by MG2C (http://mg2c.iask.in/mg2c_v2.0/) and GSDS (http://gds.cbi.pku.edu.cn/). MEME online software (http://meme-suite.org/tools/meme) was used to analyze conservative motif with the motif parameters at 30, and the rest are default.

Phylogenetic tree construction and sequence alignment

The HDAC protein sequences of Arabidopsis, rice and tomato were downloaded from TAIR (https://www.arabidopsis.org/) and NCBI databases, and MEGA 7.0 software was used to construct a phylogenetic tree of HDAC gene family by the neighbor-joining method (Neighbor-joining, NJ; bootstrap = 1000), and DNAMAN 7.0 software was used to align the amino acid sequences.

Collinearity analysis of sorghum HDAC gene family

Sorghum HDAC collinearity was identified by MCScanX verson 1.0, and TBtools (v1.05) [54] was used to drawn the Fig. 2.

Characterization of SbHDAC genes and proteins

Based on CDS sequence, sorghum interspecific phylogenetic tree and genome sequence, TBtools v1.05 was used to predict the number of introns/exons of SbHDACs [54]. MEME online program was used to analyze the conserved motif structure of the SbHDAC protein. To predict cis-acting elements of sorghum SbHDAC genes, the 2000bp upstream sequences of the sorghum transcription start site from the NCBI database were extracted for analyzing through PlantCARE (http://bioinformatics.psb.ugent.be/webtools/plantcare/html/).

Total RNA extraction and qRT-PCR analysis

The total RNA was extracted by TRizol, and the integrity of the extracted total RNA was examined by 1.0% agarose gel electrophoresis. cDNA was synthesized using HiScript® III RT SuperMix (Vazyme Biotech Co. Ltd.). Bio-Rad real-time PCR system (Bio-Rad, Hercules, CA, USA) was used to perform qRT-PCR expression analysis, and sorghum SbELF4a was used as an internal reference gene. The primer pairs used for qRT-PCR were listed in Table S1. After the amplification is completed, the melting curve and amplification curve were checked to evaluate the amplification specificity. All experiments were repeated three times for each sample. Fluorescence quantitative data was analyzed by the 2⁻ΔΔCt [55], and the Duncan’s new multiple range test (based on SPSS software) was employed for significance analysis.

Protein prokaryotic expression in E. coli

The ORF of SbHDA1, SbHDA3, SbHDT3 and SbSRT2 was amplified and inserted into pET28a (+) vector respectively. The obtained recombinant plasmid was transformed into E. coli Rosetta (DE3) competent cells. 1.0 mM isopropyl β-D-thiogalactoside (IPTG) was used to induce protein expression at 16°C, 25°C and 30°C for 24h, 16h and 12h, respectively [56]. The prokaryotic protein was evaluated through 12% SDS-PAGE electrophoresis by coomassie brilliant blue staining. Western blot was used to confirm the size of expressed protein to be HDAC. In order to study the expression of SbHDA1, SbHDA3, SbHDT3 and SbSRT2 in E. coli under different abiotic conditions, a spot assay was conducted in combination with treatment using PEG6000, D-mannitol and sodium chloride. The E. coli cells containing pET28a-SbHDA1, SbHDA3, SbHDT3 and SbSRT2 or pET28a (control) were cultured in LB medium at 37°C until the OD₆₀₀ reaches to 0.6. Then, 1.0 mM IPTG was added and further cultured at 37°C for 12h. Then LB medium (Composition of the medium are tryptone, yeast extract, NaCl and agar) was used to dilute the cultures to 10⁻¹⁰⁻⁴ times. All treatments were cultured at 37°C overnight for photographing.

Protein extraction and immunoblot analysis

The sorghum seedlings were treated by 4°C, 40°C, 300 mM D-mannitol and 250 mM NaCl for 12h, and then the samples including the control were quickly ground in liquid nitrogen. Total protein was extracted by lysis buffer (50 mM Tris-HCl pH7.5, 150 mM NaCl, 0.1% NP-40, 8 M
urea, 1 mM PMSF, 1× cocktail, 50 mM nicotinamide, 3μM Trichostatin A). After adding loading buffer and boiling for 5 min (100°C), the protein was separated by 12% sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE), then transferred to polyvinylidene fluoride membrane (PVDF) membrane. Finally, the acetylated protein was examined with anti-acetyl lysine antibody (1:1000 dilution, PTM Biolabs, Hangzhou, China), the secondary antibody was goat anti-mouse antibody (1:10000 dilution).

Abbreviations
HDACs: Histone deacetylases; PAMPs: Pathogen-associated molecular model; HATs: Histone acetyltransferases; CBP: CREB binding protein; NAD⁺: Nicotinamide adenine dinucleotide; ABA: Abscisic acid; JA: Jasmonic acid; PR1: Pathogenesis Related 1; SA: Salicylic acid; Xoo: Xanthomonas oryzae pv oryzae; ROS: Reactive oxygen species; E.coli: Escherichia coli; IPTG: Isopropyl β-D-thiogalactoside; SDS-PAGE: Sodium dodecyl sulfate polyacrylamide gel electrophoresis; PVDF: Polyvinylidene fluoride; qRT-PCR: Real-time fluorescence quantitative PCR; ABRE: Abscisic acid responsiveness elements; MYB: v-myb avian myeloblastosis viral oncogene homolog; MYC: Myelocytomatosis protein; DRE: Dehydration-responsive element; ABRE: ABA responsive element.

Supplementary Information
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Additional file 1: Figure S1. Chromosome distribution of SbHDAC genes in sorghum.

Additional file 2: Figure S2. Multiple sequence alignment of the SbHDAC proteins.

Additional file 3: Figure S3. Phylogenetic relationships and gene structures of the SbHDAC family. Exons and introns were shown by filled boxes and single lines, respectively.

Additional file 4: Figure S4. Motif analysis of SbHDAC proteins.

Additional file 5: Figure S5. Amplification and plasmids construction of SbHDAC genes into pET28a. A-D: lane 1, recombinant plasmids. lane 2, double digestion of SbHDAC1-pET28a, SbHDAC3-pET28a, SbHDAC3-pET28a, and SbSRT2-pET28a respectively.

Additional file 6: Figure S6. Detection of SbHDAC1, SbHDAC3, SbHDAC3 and SbSRT2 protein by western blot. M: Protein marker; Lane1: Supernatant of empty vector; Lane 2: Supernatant of SbHDAC1, SbHDAC3, SbHDAC3 and SbSRT2 recombinant cells.

Additional file 7: Table S1. Primer sequences.

Additional file 8: Table S2. cis-acting element analysis of the SbHDAC gene family.

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Authors’ contributions
QY, YF, XL, XX conceived and designed the experiments, ZQ and LW took samples in climate-control green house and performed experiments, MC, XF, JJ performed the experiment, LL, QW, QY, LW and ZY analyzed the data. QD performed the experiment and wrote the paper, XX and XL revised the manuscript. The authors have read and approved the manuscript.

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Availability of data and materials
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Declarations
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Competing interests
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Author details
1 Key Laboratory of Agricultural Microbiological, College of Agriculture, Guizhou University, Guiyang 550025, PR China. 2 State Key Laboratory Breeding Base of Green Pesticide and Agricultural Bioengineering, Key Laboratory of Green Pesticide and Agricultural Bioengineering, Ministry of Education, Guizhou University, Guiyang 550025, PR China.

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References
1. Boyko A, Kovalchuk I. Genome instability and epigenetic modification-hentable responses to environmental stress? Curr Opin Plant Biol. 2011;14(3):260–6.
2. Pfluger J, Wagner D. Histone modifications and dynamic regulation of genome accessibility in plants. Curr Opin Plant Biol. 2007;10(6):645–52.
3. Luo M, Liu X, Singh P, Cui YH, Zimmerli L, Wu KQ. Chromatin modifications and remodeling in plant abiotic stress responses. Biochim Biophys Acta. 2012;1819(2):129–36.
4. Allfrey VG, Faulkner R, Mirsky AE. Acetylation and methylation of histones and their possible role in the regulation of mRNA synthesis. Proc Natl Acad Sci U S A. 1964;51(5):786–94.
5. Liu XC, Yang SG, Zhao ML, Zhao ML, Luo M, Yu CW, et al. Transcriptional repression by histone deacetylases in plants. Mol Plant. 2014;7(5):764–72.
6. Pandey R, Müller A, Napoli CA, Selinger DA, Pikaard CS, Richards EJ, et al. Analysis of histone acetyltransferase and histone deacetylase families of Arabidopsis thaliana suggests functional diversification of chromatin modification among multicellular euakaryotes. Nucleic Acids Res. 2002;30(23):5036–55.
7. Ma XJ, Lv SB, Zhang C, Yang CP. Histone deacetylases and their functions in plants. Plant Cell Rep. 2013;32(4):465–78.
8. Imaj S, Armstrong CM, Kaeberlein M, Guarente L. Transcriptional silencing and longevity protein Sir2 is an NAD-dependent histone deacetylase. Nature. 2000;403(6771):795–800.
9. Lussier A, Borsch G, Loidl A, Haas H, Loidl P. Identification of maize histone deacetylase HD2 as an acidic nucleolar phosphoprotein. Science. 1997;277(5322):88–91.
10. Han Z, Yu H, Zhao Z, Hunter D, Luo X, Duan J, et al. AthD2D gene plays a role in plant growth, development, and response to abiotic stresses in Arabidopsis thaliana. Front Plant Sci. 2016;7:310.
11. Zhou C, Labbe H, Sridha S, Wang L, Tian L, Latoszek-Green M, et al. Expression and function of HD2-type histone deacetylases in Arabidopsis development. Plant J. 2004;38(5):715–24.
12. Li YX, Liu Y, Zhao N, Yang XJ, Li YQ, Zhai FZ, et al. Checkpoint regulator 17X is epigenetically regulated by HDAC3 and mediates resistance to HDAC inhibitors by reprogramming the tumor immune environment in colorectal cancer. Cell Death Dis. 2020;11(9):753.
13. Sendra R, Rodrigo I, Salvador ML, Franco L. Characterization of pea histone deacetylases. Plant Mol Biol. 1988;11(6):857–66.
14. Kornet N, Scheres B. Members of the GCN5 histone acetyltransferase complex regulate PLETHORA-mediated root stem cell niche maintenance and transit amplifying cell proliferation in Arabidopsis. Plant Cell. 2009;21(4):1070–9.

15. Han SK, Song JD, Noh YS, Noh B. Role of plant CBF/p300-like genes in the regulation of flowering time. Plant J. 2007;49(1):103–14.

16. Latrasse D, Benhamed M, Henry Y, Domeninchi S, Kim W, Zhou DX, et al. The MYST histone acetyltransferases are essential for gametophyte development in Arabidopsis. BMC Plant Biol. 2008;8:121.

17. Nellissen H, Fleury D, Bruno L, Robles P, Veylder LD, Traas J, et al. The elongata mutants identify a functional elongator complex in plants with a role in cell proliferation during organ growth. Proc Natl Acad Sci U S A. 2005;102(21):7754–9.

18. Alinsug MV, Chen FF, Luo M, Tai R, Jiang L, Wu K. Subcellular localization of class I HDAs in Arabidopsis thaliana: nucleocytoplasmic shuttling of HDA15 is driven by light. PLoS One. 2012;7(2):e30846.

19. Luo M, Wang YY, Liu XC, Yang SG, Lu Q, Cui YH, et al. HD2C interacts with HDA6 and is involved in ABA and salt stress response in Arabidopsis. J Exp Bot. 2012;63(8):3297–306.

20. Kuang JF, Chen JY, Luo M, Wu KQ, Sun W, Jiang YM, et al. Histone deacetylase interacts with ERF1 and is involved in longin fruit senescence. J Exp Bot. 2012;63(1):441–54.

21. Bharti K, Von Koskul-Doring P, Bharti S, Kumar P, Tintschi-Korbitzer A, Treutler E, et al. Tomato heat stress transcription factor HsB1 represents a novel type of general transcription coactivator with a histone-like motif interacting with the plant CREB binding protein ortholog HAC1. Plant Cell. 2004;16(6):1521–35.

22. Li W, Liu H, Cheng Z, Su YH, Han HN, Zhang Y, et al. DNA methylation and histone modifications regulate de novo shoot regeneration in Arabidopsis by modulating WUSCHEL expression and auxin signaling. PLoS Genet. 2011;7(8):e1002243.

23. Wu K, Malik K, Tian L, Brown D, Mikbi B. Functional analysis of a RPD3 histone deacetylase homologue in Arabidopsis thaliana. Plant Mol Biol. 2000;44(2):167–76.

24. Liu X, Luo M, Zhang W, Zhao JH, Zhang JX, Wu KQ, et al. Histone acetyltransferases in rice (Oryza sativa L.). Phylogenetic analysis, subcellular localization and expression. BMC Plant Biol. 2012;12:145.

25. Aiese Cigliano R, Sanseverino W, Cremona G, Ercolano MR, Conicella C, et al. Histone deacetylases from barley, during seed development and after dormancy, and gene expression analysis under multiple abiotic stresses, DNA methylation and histone modifications. BMC Genomics. 2012;13:286.

26. Yuan LY, Dai HW, Zheng ST, Huang R, Tong HR. Genome-wide identification of rice histone deacetylase HDA705. Plant J. 2012;69(3):448–59.

27. Wu KQ, Tian L, Malik K, Brown D, Miki B. Functional analysis of HD2 as a novel regulator of abscisic acid responses in Arabidopsis by modulating WUSCHEL expression and auxin signaling. J Exp Bot. 2016;67(6):1703–13.

28. Cho LH, Yu JX, Zhang JX, Wu PY, Yang SG, Wu KQ. Identification and characterization of histone deacetylases in tomato (Solanum lycopersicum). Front Plant Sci. 2014;5:760.

29. Lee HG, Seo PJ. MYB696 recruits the HDA15 protein to suppress negative regulators of ABA signalling in Arabidopsis. Nat Commun. 2019;10(1):1713.

30. Yuan LB, Chen X, Chen HH, Wu KQ, Huang SZ. Histone deacetylases HDA6 and HDA9 coordinately regulate valve cell elongation through affecting auxin signalling in Arabidopsis. Biochem Biophys Res Commun. 2019;508(3):695–700.

31. Zheng Y, Ding Y, Sun X, Xie SS, Wang D, Liu XY, et al. Histone deacetylase HD49 negatively regulates salt and drought stress responsiveness in Arabidopsis. J Exp Bot. 2016;67(6):1703–13.

32. Cho LH, Yoon J, Wai AH, An G. Histone deacetylase 701 (HDT701) induces flowering in rice by modulating expression of OsI5D1. Mol Cells. 2018;41(7):665–75.

33. Chung PJ, Kim YS, Jeong JS, Park SH, Nahm BH, Kim JK. The histone deacetylase OsHDAC1 epigenetically regulates the OsNaC2 gene that controls seed root growth in rice. Plant J. 2009;59(5):764–76.

34. Kubo FC, Yasui Y, Ohmori K, Kumamuro T, Tanaka W, Hirano HY. DWARF WITH SLENDER LEAF1 encoding a histone deacetylase plays diverse roles in rice development. Plant Cell Physiol. 2020;61(3):457–69.

35. Guo X, Xie Q, Li B, Su H. Molecular characterization of nine tissue-specific or stress-responsive genes of histone deacetylase in tomato (Solanum lycopersicum). Genet Mol Biol. 2020;43(1):e20180295.

36. Ma XJ, Liang XY, Lv SB, Guan T, Jiang TB, Cheng X. Histone deacetylase gene PHDHT902 modifies adventitious root formation and negatively regulates salt stress tolerance in poplar. Plant Sci. 2020;290:110301.

37. Demetriou K, Kapazoglou A, Bladenopoulos K, Tsafaris AS. Epigenetic chromatin modifiers in barley. B. Characterization and expression analysis of the HD1 family of barley histone deacetylases during development and in response to jasmonic acid. Plant Mol Biol Rep. 2010;28(1):9–21.

38. Demetriou K, Kapazoglou A, Tondelli A, Frasca E, Stanca MA, Bladenopoulos K, et al. Epigenetic chromatin modifiers in barley. I. Cloning, mapping and expression analysis of the plant specific HD2 family of histone deacetylases from barley, during seed development and after hormonal treatment. Physiol Plantarum. 2009;136(3):358–68.

39. Nicolas-Francès V, Grandperret V, Liegard B, Jeandroz S, Vasselon D, Aimé S, et al. Evolutionary diversification of type-2 HDAC structure, function and regulation in Nicotiana tabacum. Plant Sci. 2018;269:66–74.

40. Chen CJ, Chen H, Zhang Y, Thomas HR, Frank MH, He YH. TBtools: an integrative toolkit developed for interactive analyses of big biological data. Mol Plant. 2020;13(8):1194–202.
55. Pfaffl MW. A new mathematical model for relative quantification in real-time RT-PCR. Nucleic Acids Res. 2001;29(9):e45.
56. Guan L, Chen L, Chen Y, Zhang N, Han Y. Expression and activity analysis of fructosyltransferase from aspergillus oryzae. Protein J. 2017;36(6):1–9.

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