



Genome-wide identification and analysis of the evolution and expression pattern of the *HVA22* gene family in three wild species of tomatoes

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ABSTRACT

Wild tomato germplasm is a valuable resource for improving biotic and abiotic stresses in tomato breeding. The *HVA22* is widely present in eukaryotes and involved in growth and development as well as stress response, such as cold, salt, drought, and biotic stress. In the present study, we identified 45 *HVA22* genes in three wild species of tomatoes. The phylogenetic relationships, gene localization to chromosomes, gene structure, gene collinearity, protein interactions, and *cis*-acting element prediction of all 45 *HVA22* genes (14 in *Solanum pennellii*, 15 in *S. pimpinellifolium*, and 16 in *S. lycopersicoides*) were analyzed. The phylogenetic analysis showed that the all *HVA22* proteins from the family Solanaceae were divided into three branches. The identified 45 *HVA22* genes were grouped into four subfamilies, which displayed similar number of exons and expanded in a fragmentary replication manner. The distribution of *HVA22* genes on the chromosomes of the three wild tomato species was also highly similar. RNA-seq and qRT-PCR revealed that *HVA22* genes were expressed in different tissues and induced by drought, salt, and phytohormone treatments. These results might be useful for explaining the evolution, expression patterns, and functional divergence of *HVA22* genes in *Lycopersicon*.

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INTRODUCTION

The *HVA22* gene was first isolated from the dextrin layer of barley (*Hordeum vulgare*) in 1993 (Shen, Uknes & Ho, 1993). Homologs to the *HVA22* gene have been identified in eukaryotes, such as yeast, cereals, *Arabidopsis thaliana*, nematodes, mice, and humans. Approximately 367 *HVA22* homologs have been found in eukaryotes, which present a conserved TB2/DP1 (deleted-in-polyposis) domain (PF03134) (Guo & David Ho, 2008; Sharon & Suvarna, 2017; Gomes Ferreira et al., 2019). Notably, no homologs have been described in prokaryotes to date, suggesting that *HVA22* is likely involved in eukaryote-specific functions.

In plants, the expression patterns of *HVA22* during development are well characterized. The accumulated transcripts of *HVA22* homologs in leaves are highly induced by abscisic acid (ABA), drought, cold, and salt stresses in both barley and *A. thaliana* (Shen, Uknes & Ho, 1993; Shen et al., 2001; Chen et al., 2002). The heterologous expression of the barley *HVA22* protein improves the salt tolerance and the survival rate of *E. coli* under cold stress by delaying its proliferation (Lu, 2013). Additionally, a recent study on barley discovered that the amount of the transcription product of the *HVA22* gene was significantly higher in barley *hvabi5d* mutant strains than in wild-type plants under drought conditions (Collin et al., 2020). Subsequently, the *HVA22* gene was also shown to be induced by drought and salt stresses in different plants. For example, the accumulation of transcript of *HVA22* homologs in common wheat roots under drought stress was reported (Grzesiak et al., 2019). The expression of the *HVA22* gene was significantly higher in drought-tolerant bermudagrass subjected to external simulated drought (Liu et al., 2014). The *HVA22*-like protein encoded by BG598159 was found to play an important role in salt and drought stress in potato by interacting with the StPDI1 protein and participating in the sucrose transport pathway (Eggert et al., 2016). The expression of the tomato *HVA22* gene was also significantly induced by salt and drought stresses (Wai et al., 2022). Also, the heterologous expression of the *Citrus clementina* *CcHVA22d* gene in tobacco enhanced dehydration tolerance and significantly reduced the H₂O₂ content in a short-term dehydration environment (Gomes Ferreira et al., 2019).

Studies have shown that the *HVA22* gene also plays an important role in endoplasmic reticulum-related pathways. The Yop1p gene, a homolog of *HVA22* in yeast, appears to be involved in the translocation of substances from the endoplasmic reticulum to the Golgi apparatus during cellular activity (De Antoni et al., 2002). Previous studies on the yeast Yop1p protein revealed that the Yop1p/DP1 protein interacted with the Rtn4/NogoA protein, thereby co-regulating the interactions between other proteins *in vivo* as well as endoplasmic reticulum function (Hu et al., 2008). The *HVA22* gene in the dextrin layer of barley seeds also has a similar function as yeast Yop1p. The accumulation of the *HVA22* gene in the dextrin layer after induction by abscisic acid (ABA) inhibits vesicle transport in cells, thereby delaying the incorporation of storage protein vesicles, which is a process thought to play a role in regulating seed germination and seedling growth (Guo & David Ho, 2008). More recent studies in rice have shown that the rice *HVA22* family gene *OsHLP1* promotes disease resistance mechanisms in plants by maintaining endoplasmic reticulum homeostasis (Meng et al., 2022). To date, studies on the involvement of *HVA22* homologs in the cellular vesicle transport pathway are scarce.

The supply of food and vegetable production have become major issues with the continuous rise in the world population. This is compounded by the potential impact of an increasingly changing climate on crop productivity. Extreme temperatures, drought, and soil salinization are the main adverse environments often encountered by plants (Gong et al., 2020). Tomatoes are a favorite vegetable worldwide. However, adverse environments such as salt, drought, and cold severely affect tomato growth and development (Chaudhary et al., 2019a; Chaudhary et al., 2019b). Wild tomatoes belonging to the genus *Lycopersicon* have higher tolerance to salt, drought, and cold than cultivated tomatoes (Szymański et al., 2020).

Thus, wild tomatoes were an important genetic resource for our study on tomato response to adversity. It has been demonstrated that *HVA22* genes are significantly upregulated in rice (Zhao *et al.*, 2021), *A. thaliana* (Chen *et al.*, 2002), barley (Shen *et al.*, 2001), and tomatoes (Wai *et al.*, 2022) in response to salt and drought stresses. However, systematic studies on *HVA22* family genes in wild tomatoes have not been reported. In this study, we used the bioinformatics methods to comprehensively identify *HVA22* family genes in three species of wild tomatoes (*S. pimpinellifolium*, *S. pennellii*, and *S. lycopersicoides*). This study might provide a theoretical reference for elucidating *HVA22* family gene members and mining tomato genes for resistance to abiotic stresses.

MATERIALS & METHODS

Identification of *HVA22* family genes in the family Solanaceae

The protein sequence of the *Arabidopsis thaliana* *HVA22* family gene was downloaded from the Ensembl database (<http://plants.ensembl.org/index.html>) (Yates *et al.*, 2022). Protein sequence files for three species of wild tomatoes (*S. pimpinellifolium*, *S. pennellii*, and *S. lycopersicoides*), tobacco (*Nicotiana benthamiana*), pepper (*Capsicum annuum*), eggplant (*Solanum melongena*), and potato (*Solanum tuberosum*), as well as genome files from the Solanaceae genome database (<https://Solgenomics.net/>), were downloaded (Fernandez-Pozo *et al.*, 2015). The hidden Markov model of the structural domain of the *HVA22*-like protein TB2/DP was obtained from the Pfam (PF03134) (<http://pfam.xfam.org/>) and PANTHER (PTHR12300) (<http://www.pantherdb.org/>) database (Mistry *et al.*, 2021; Thomas *et al.*, 2022). The screened *HVA22* protein sequences were validated using the online protein structural domain prediction tool HMM search (<http://hammer.org/>) (Finn, Clements & Eddy, 2011), and genes that did not contain the TB2/DP structural domain were removed. The physicochemical properties of the screened tomato *HVA22* family of proteins were predicted on the ExPASy website (<https://www.expasy.org/protparam/>) (Artimo *et al.*, 2012). The subcellular localization prediction of *HVA22* family genes in three species of wild tomatoes was performed on the WoLF PSORT online tool (<https://wolfpsort.hgc.jp/>) (Horton *et al.*, 2007).

Construction of conserved motifs, *cis-acting* elements, and phylogenetic tree of *HVA22* gene in three species of wild tomatoes

The *HVA22* family gene motifs (Grundy *et al.*, 1997) in the three species of wild tomatoes were searched using the MEME online tool (<https://meme-suite.org/meme/tools/meme>). The number of search base sequences was set to 20, and the minimum and maximum widths were set to 6 and 50, respectively. The results from the MEME search were used to map the conserved modal motifs and gene structures using TBtools. Multiple sequence comparisons were performed using MEGA 11 software, and a phylogenetic tree was constructed using the maximum likelihood method (ML) (Tamura, Stecher & Kumar, 2021). The constructed phylogenetic trees were embellished using the online tools ITOOL (<https://itol.embl.de/>) (Letunic & Bork, 2021). The 2,000-bp promoter sequence upstream of the *HVA22* family gene in the three species of wild tomatoes was extracted, and the *cis-acting* element of the *HVA22* family gene was predicted using the PlantCARE database

(<http://bioinformatics.psb.ugent.be/webtools/plantcare/html/>) (Rombauts *et al.*, 1999) and visualized using TBtools (Chen *et al.*, 2020).

Interaction network and expression analysis of *HVA22* family homologous genes

MCSanX was used to analyze the *HVA22* gene in four species of *Lycopersicon* (*S. lycopersicum*, *S. lycopersicoides*, *S. pennellii*, and *S. pimpinellifolium*) and Solanaceae [tobacco (*N. benthamiana*), pepper (*C. annuum*), potato (*S. tuberosum*) and eggplant (*S. melongena*)] interspecies as well as the intraspecific collinearity in four species of *Lycopersicon*. The substitution rate of paralogous homologous genes was calculated using Ka/Ks_Calculator 2.0 (Wang *et al.*, 2010). The direct homologous genes between species and the paralogous homologous gene collinearity within species were visualized using TBtools. The STRING online website was used to predict protein–protein interaction relationships (Szklarczyk *et al.*, 2019), interactions, after which the relationship data given by the predictions were visualized using Cytoscape 3.9.1. The expression matrices for different tissues and developmental stages of *S. pimpinellifolium* were downloaded from the Tomato Function Genomics database (<http://ted.bti.cornell.edu/>), which included expression data for root, stem, leaves, young flower buds, anthesis flowers, 10 days post anthesis, 20 days post anthesis, 30 days post anthesis, and ruptured fruit (Fei *et al.*, 2010). From these, *HVA22* family genes were selected and the expression profiles were heat-mapped using TBtools.

Total RNA extraction and reverse transcription

The extraction of plant leaf RNA was accomplished using a Tiangen plant polyphenol polysaccharide total RNA extraction kit (Beijing, China). The cDNA synthesis of extracted total RNA was performed using a 5 × All-In-one RTMasterMix (AccuRT Genomic DNA Removal Kit; G492, ABM, Vancouver, Canada) reverse transcription kit. qRT-PCR-specific primers were designed using the NCBI online primer tool (https://www.ncbi.nlm.nih.gov/tools/primer-blast/index.cgi?LINK_LOC=BlastHome), and the designed qRT-PCR primers were sent to Biotech Biologicals (Shanghai, China) for synthesis (Table S1). Quantitative PCR (qPCR) analysis was subsequently performed on a LightCycler machine using ChamQ Universal SYBR qPCR Master Mix (Q711, Vazyme, Nanjing, China) with *Slactin* as the internal reference gene. Three replicates of each treatment were performed. The relative expression was calculated by the $2^{-\Delta\Delta C_t}$ Method (Livak & Schmittgen, 2001).

Plant material and growing conditions

The plant material used in this study was a wild tomato variety (*S. pimpinellifolium*, LA1589). For stress treatment, the seeds were sown in nutrient soil and vermiculite ($v/v = 2:1$) in a growth room at 24 ± 2 °C under 16-h light /8-h dark cycle. The seedlings were treated with Hoagland every week until use. Then, the plants were treated with different stresses, including 100 mM abscisic acid (ABA) or methyl jasmonate (MeJA), 200 mM NaCl, and 15% PEG6000. The leaves were collected and stored in liquid nitrogen quickly

for RNA extraction at different time points (0, 2, 6, 12, and 24 h). Three independent biological replicates were included for each sample in the experiment.

RESULTS

Identification of physicochemical properties and prediction of subcellular localization of *HVA22* family proteins from the three species of wild tomatoes

In the present study, we used the Simple HMM Search function of the TBtools tool to search the *HVA22* gene in *S. lycopersicoides*, *S. pennellii*, and *S. pimpinellifolium*. The retrieved genes were then subjected to structural domain validation using the HMM search (<http://hmmer.org/>) (Finn, Clements & Eddy, 2011) and InterPro (<https://www.ebi.ac.uk/interpro/result/InterProScan/#table>) online tool, and those without TB2/DP1 structural domains were discarded. The 45 *HVA22* family genes were finally determined in three species of wild tomatoes (15 in *S. pimpinellifolium*, 16 in *S. lycopersicoides*, and 14 in *S. pennellii*) and named as *HVA22a–HVA22p* according to their positions on chromosomes. Subsequently, we characterized the physicochemical properties and predicted the subcellular localization of the *HVA22* gene family member proteins from these three species of wild tomatoes. All *HVA22* family proteins in three species of wild tomatoes (15 in *S. pimpinellifolium*, 14 in *S. pennellii*, and 16 in *S. lycopersicoides*) had amino acid lengths between 88 and 603 and protein molecular masses between 9989.65 to 68493.8 Da, with an isoelectric point of 5.53–10.09 and aliphatic index of 51.68–123.66. In terms of hydrophilicity, most of the proteins exhibited hydrophilic proteins (GRAVY <0) and a few exhibited hydrophobic proteins (GRAVY >0). The subcellular localization predictions showed that *HVA22*-like was localized to multiple organelles in three species of wild tomatoes; most *HVA22* family genes were localized to the endoplasmic reticulum, chloroplasts, cytoplasm, and nucleus; only the *SpiHVA22k* gene was localized to the vesicle (Table 1).

Phylogenetic tree analysis of *HVA22* family genes

We combined 45 *HVA22* genes from three species of wild tomatoes (16 in *S. lycopersicoides*, 14 in *S. pennellii*, and 15 in *S. pimpinellifolium*) with *HVA22* genes from *A. thaliana* and cultivated tomatoes (*S. lycopersicum*) to construct a complete phylogenetic tree (Table S3). The phylogenetic tree showed that *HVA22* genes in the cultivated tomatoes, three species of wild tomatoes, and *A. thaliana* were divided into two clades: I and II (Fig. 1A). The subgroups I and II were included in clade I, and the subgroups III and IV were included in clade II. The *HVA22* family genes in the three species of wild tomatoes in groups I, II, III, and IV clustered on a subgroup with the *A. thaliana* and cultivated *HVA22* genes. To enable a comprehensive understanding of the quantitative distribution of *HVA22* family gene members in tobacco (*N. benthamiana*), potato (*S. tuberosum*), eggplant (*S. melongena*), pepper (*C. annuum*), and three kinds of wild tomatoes (*S. pimpinellifolium*, *S. pennellii*, and *S. lycopersicoides*). We identified 120 *HVA22* family genes in selected Solanaceae species, constructed a phylogenetic tree, and designated these genes based on their chromosomal locations (Table S4). All *HVA22* genes were classified into three

Table 1 Physicochemical properties of three species from wild tomato HVA22 protein.

Species	Gene id	Gene name	Length	MW(Da)	pI	Aliphatic index	GRAVY	Subcellular localization
<i>S.pimpinellifolium</i>	Spim05g006320.1.1	<i>SpiHVA22h</i>	171	20248.41	7.67	98.71	-0.101	E.R
	Spim03g025310.1.1	<i>SpiHVA22c</i>	182	20623.89	6.65	102.86	0.066	Chlo
	Spim06g022990.1.1	<i>SpiHVA22i</i>	133	15554.63	9.49	115.11	0.249	Extr
	Spim06g026920.1.1	<i>SpiHVA22j</i>	135	15665.57	8.8	104.74	0.21	Chlo
	Spim11g010400.1.1	<i>SpiHVA22o</i>	167	19963.73	9.32	113.23	0.34	Chlo
	Spim03g033180.1.1	<i>SpiHVA22d</i>	176	20507.66	6.41	96.93	0.005	Extr
	Spim04g011050.1.1	<i>SpiHVA22e</i>	105	12446.7	8.66	104.86	0.13	Cyto
	Spim10g007900.1.1	<i>SpiHVA22l</i>	156	18030.22	9.26	95.64	0.026	Chlo
	Spim04g032680.1.1	<i>SpiHVA22g</i>	323	35618.35	8.81	71.05	-0.346	Chlo
	Spim01g007550.1.1	<i>SpiHVA22a</i>	180	21270.52	5.95	92.11	-0.116	Cyto
	Spim09g019680.1.1	<i>SpiHVA22k</i>	241	28172.67	8.93	84.94	-0.026	Vacu
	Spim10g015050.1.1	<i>SpiHVA22m</i>	603	68493.8	8.68	81.51	-0.408	Nucl
	Spim04g027770.1.1	<i>SpiHVA22f</i>	494	56507.39	9.34	89.82	-0.226	Nucl
	Spim01g044830.1.1	<i>SpiHVA22b</i>	255	29350.56	8.68	108.94	0.308	Nucl
	Spim10g025730.1.1	<i>SpiHVA22n</i>	150	17597.28	9.1	80.73	-0.339	Cyto
	Sopen05g003250.1	<i>SpHVA22g</i>	171	20234.38	7.67	98.13	-0.102	E.R.
	Sopen11g005670.1	<i>SpHVA22m</i>	142	16848.74	8.58	95.42	0.063	Cyto
	Sopen03g028180.1	<i>SpHVA22b</i>	136	15845.88	9.37	115.44	0.21	Extr
	Sopen04g006560.1	<i>SpHVA22e</i>	131	15267.96	5.53	106.41	0.186	Extr
	Sopen03g030110.1	<i>SpHVA22c</i>	180	20364.57	6.65	104	0.076	Nucl
Sopen06g029050.1	<i>SpHVA22h</i>	135	15652.57	8.8	104.74	0.231	Chlo	
<i>S.pennellii</i>	Sopen10g022980.1	<i>SpHVA22k</i>	187	21851.19	7.02	83.9	-0.07	E.R.
	Sopen03g035280.1	<i>SpHVA22d</i>	176	20507.66	6.41	96.93	0.005	Extr
	Sopen10g003720.1	<i>SpHVA22i</i>	156	18020.24	9.24	93.14	0.013	Chlo
	Sopen12g031160.1	<i>SpHVA22n</i>	302	33844.67	9.5	68.15	-0.447	Chlo
	Sopen04g034960.1	<i>SpHVA22f</i>	304	33623.99	8.95	68.12	-0.379	Chlo
	Sopen10g032550.1	<i>SpHVA22l</i>	188	22108.6	8.63	86.7	-0.227	Extr
	Sopen10g018090.1	<i>SpHVA22j</i>	559	63615.22	8.69	81.82	-0.444	Nucl
	Sopen01g003320.1	<i>SpHVA22a</i>	155	18113.25	10.09	51.68	-0.977	Nucl
	Solyd05g052440.1	<i>SlydHVA22h</i>	171	20176.3	7.67	97.6	-0.113	E.R.
	Solyd11g055150.1	<i>SlydHVA22n</i>	123	14697.04	7.9	91.14	-0.112	Cyto
	Solyd11g055100.1	<i>SlydHVA22m</i>	145	17473.82	8.63	123.66	0.675	Chlo
	Solyd03g072250.1	<i>SlydHVA22d</i>	190	21617.78	6.16	90.37	-0.049	Cyto
	Solyd06g073510.1	<i>SlydHVA22i</i>	135	15689.63	8.8	104.74	0.205	Chlo
	Solyd03g077950.1	<i>SlydHVA22e</i>	176	20507.66	6.41	96.93	0.005	Extr
Solyd12g070530.1	<i>SlydHVA22o</i>	229	26296.16	9.05	65.15	-0.345	Chlo	
Solyd12g070590.1	<i>SlydHVA22p</i>	302	33831.62	9.28	68.15	-0.456	Chlo	
Solyd04g078410.1	<i>SlydHVA22g</i>	313	34770.38	8.87	68.02	-0.35	Chlo	
Solyd01g052910.1	<i>SlydHVA22a</i>	182	21641.02	5.91	95.88	-0.036	Nucl	
Solyd10g061800.1	<i>SlydHVA22l</i>	241	27758.68	7.05	78.92	-0.22	Chlo	

(continued on next page)

Table 1 (continued)

Species	Gene id	Gene name	Length	MW(Da)	pI	Aliphatic index	GRAVY	Subcellular localization
	Solyd10g059860.1	<i>SlydHVA22k</i>	547	62120.36	8.28	79.85	−0.452	Nucl
	Solyd10g052770.1	<i>SlydHVA22j</i>	119	14080.39	9.14	85.21	−0.242	Chlo
	Solyd01g086740.1	<i>SlydHVA22b</i>	306	34201.23	8.87	110.56	0.206	Nucl
	Solyd03g070260.1	<i>SlydHVA23c</i>	88	9989.65	9.64	84.2	−0.172	Cyto

groups (Fig. 2B). Large differences were found in amino acid length and structural domains of HVA22 family genes. HVA22 family proteins in group III had the most pronounced differences in length and structural domains. However, the amino acid lengths were essentially similar in each cluster in group III. The HVA22 family proteins in group III also possessed the Zf-met, RVT-3, and LRR_8 structural domains besides the TB2/DP structural domain unique to HVA22 proteins. In addition, the Solanaceous HVA22 family proteins with Zf-met (*SpiHVA22m*, *SpHVA22j*, *SlydHVA22k*, *StHVA22v*, *CaHVA22i*, *SmHVA22o*, *NbHVA22c*, *NbHVA22u*, *SpiHVA22b*, *SlydHVA22b*, *StHVA22h*, and *NbHVA22n*) and RVT-3 (*SpiHVA22f*, *SlydHVA22f*, *StHVA22g*, *SmHVA22f*, *CaHVA22g*, and *NbHVA22g*) structural domains clustered in group III. On the contrary, *NbHVA22r* containing the LRR_8 structural domain was independently classified into a distinct cluster, which might be related to the fact that only *NbHVA22r* contained the LRR_8 structural domain among the numerous HVA22 family proteins in species belonging to Solanaceae. Moreover, the distribution position of the TB2/DP structural domain in the amino acid sequence of the HVA22 family in Solanaceae was at the N-terminal, except for *SlydHVA22m*, *NbHVA22a*, *CaHVA22l*, and *SmHVA22d*, which were at the C-terminal.

Conserved motif and gene structure analyses of the HVA22 gene family

The conserved motifs of the HVA22 family genes of three wild tomato species were predicted using the online tool MEME to understand the specific distribution of conserved motifs in HVA22 genes in the three species of wild tomatoes, and a total of 20 conserved motifs were identified (Fig. 2). Motifs 1, 2, and 3 formed the TB2/DP structural domain, which was distributed in HVA22 family genes in the three species of wild tomatoes. Additionally, the Zf-met structural domain consisting of Motifs 9, 10, and 14 was also distributed in some wild tomato HVA22 family of proteins. The HVA22 protein possessing the Zf-met structural domain was mainly concentrated on a small cluster in group I. The analysis indicated that the amino acid motif composition of the HVA22 family in the same group was approximately similar. HVA22 genes containing a Zf-met structural domain exhibited more motifs, with the exception of *SpiHVA22f*, *SpiHVA22b*, and *SlydHVA22b*.

The structural differences in exon–intron arrangement are an important source of gene family variation and plant diversity. Different structures lead to differences in gene expression and function (Xu et al., 2012). Our results showed that the HVA22 family genes in the three species of wild tomatoes were divided into three major groups by the phylogenetic tree, with a large degree of similarity in the exon–intron arrangement in most of the same clusters (Fig. 2). However, large differences existed in the arrangements in

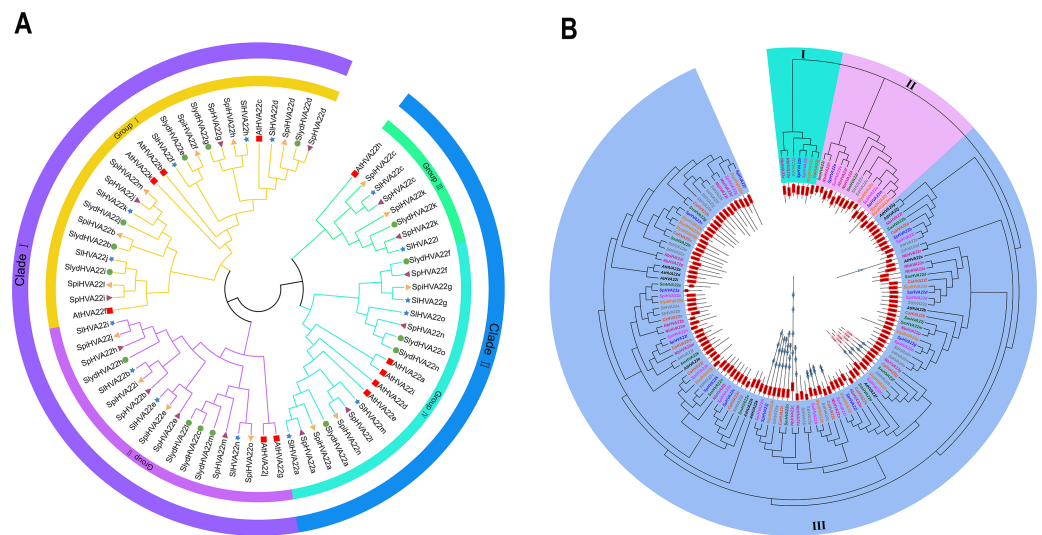


Figure 1 Phylogenetic tree analysis of the *HVA22* gene. (A) Phylogenetic trees were constructed for 60 *HVA22* genes from *Lycopersicon* using the ML method with 1,000 bootstrap repetitions. A species abbreviation was provided prior to each *HVA22* protein name: Sl, *Solanum lycopersicum*; Sp, *Solanum pennellii*; At, *Arabidopsis thaliana*; Spi, *Solanum pimpinellifolium*; and Slyd, *Solanum lycopersicoides*. (B) Phylogenetic tree of the *HVA22* family in Solanaceae. The phylogenetic tree was constructed using the ML method with 1,000 bootstrap repetitions. The different coloured *HVA22* genes were derived from different Solanaceae species, and the conserved structural domains of the corresponding *HVA22* genes are shown inside the evolutionary tree, with the TB2/DP1 structural domain in red, the Zf-met structural domain in blue, and the RVT-3 structural domain in pink.

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some of the clusters. In groups I and II, the number of exons was mostly 5, and the number of exons in a few individual *HVA22* family genes was 2 (*SpiHVA22e*), 3 (*SlydHVA22c*, and *SlydHVA22j*), 4 (*SlydHVA22n*, *SlydHVA22m*, and *SpiHVA22i*), and 8 (*SlydHVA22b*, *SlydHVA22k*, *SpiHVA22m*, and *SpHVA22j*). The number of group III exons was highly variable, ranging from 2 to 9. Despite the large variation in the number of exons in group III, the *HVA22* genes in each subgroup in group III exhibited similar gene structures.

***HVA22* gene promoter analysis in the three species of wild tomatoes**

We performed a *cis-acting* element analysis of the 2,000-bp promoter sequence upstream of the *HVA22* gene in three species of wild tomatoes (*S. lycopersicoides*, *S. pennellii*, and *S. pimpinellifolium*) (Fig. 3). The analysis showed that the *cis-acting* elements in the *HVA22* gene were divided into four categories: light-responsive *cis-acting* elements, phytohormone-responsive *cis-acting* elements, biotic/abiotic stress *cis-acting* elements, and growth and development *cis-acting* elements. *Cis-acting* elements involved in light response, phytohormone response, and development were distributed in *HVA22* family genes in all three species of wild tomatoes, while *cis-acting* elements involved in plant growth and abiotic stress were only present in the promoters of some *HVA22* family genes. In the present study, five hormone response elements were identified to be involved in the transcriptional initiation of the *HVA22* gene: abscisic acid response element (ABRE), salicylic acid response element (TCA-element and SARE), gibberellin response element (TATC-box, GARE-motif



Figure 2 Genetic structure analysis. Phylogenetic relationships, Structure and conserved motifs of *HVA22* genes in three types of wild tomatoes. The different coloured *HVA22* genes were derived from different wild tomato species. Cyan boxes indicate exons, green boxes indicate UTR and black lines indicate introns. The numbers 1–20 and the different colored boxes indicate motifs.

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and P-box), auxin-responsive element (TGA-element, AuxRR-core, and TGA-box), and methyl jasmonate response element (TGACG-motif and CGTCA-motif). Four response plant biotic/abiotic stress elements were found, namely, *cis*-acting elements involved in defense and stress response (TC-rich repeats, and WUN-motif), *cis*-acting elements involved in low-temperature response (LTR), and MYB-binding sites involved in drought-inducing elements (MBS). Six species (CAT-box, AACA_motif, GCN4-motif, circadian, RY-element, and MSA-like) were involved in plant growth and developmental response elements. The largest number of *cis*-acting element types were involved in light response, with eight light response elements identified in the *HVA22* family of genes; among these, except for *SpHVA22j*, the promoter sequences of the remaining *HVA22* family genes were distributed with *cis*-acting elements associated with light response. Among the promoters of the *HVA22* family genes in the three species of wild tomatoes, apart from *SlydHVA22b*, *SlydHVA22c*, *SlydHVA22e*, *SlydHVA22f*, *SlydHVA22g*, *SlydHVA22j*, *SlydHVA22n*, *SlydHVA22o*, *SlydHVA22p*, *SpHVA22b*, *SpHVA22b*, *SpHVA22f*, *SpHVA22h*, *SpHVA22i*, *SpHVA22n*, *SpiHVA22a*, *SpiHVA22g*, and *SpiHVA22l*, all other *HVA22* family genes contained *cis*-acting elements in response to low temperature or drought. Four *cis*-acting elements involved in biotic/abiotic stress were not as widely distributed in the promoters of *HVA22* family genes in the three species of wild tomatoes as were light-responsive elements and phytohormone-responsive elements. This also suggested that some *HVA22* family genes were involved in responding to abiotic stresses.

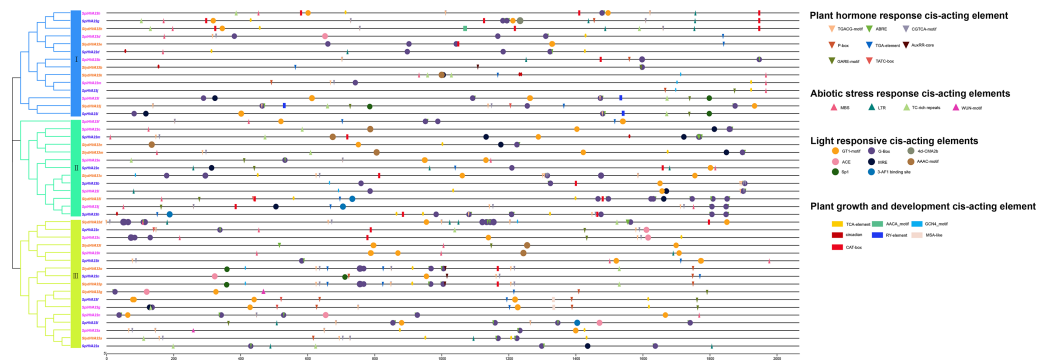


Figure 3 Distribution of CREs of *HVA22* genes in three species from wild tomato. Different colored squares indicate different branches of *HVA22* family genes in the phylogenetic tree. Different CREs were indicated by different shapes, inverted triangles indicate hormone response elements, circles indicate light response elements, boxes indicate growth and development related elements, triangles indicate stress response related elements, and different elements were indicated by different colors.

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Gene localization to chromosomes and collinearity analysis of *HVA22* in wild tomatoes

The *HVA22* family genes in three species of wild tomatoes were mainly distributed on chromosomes Chr 01, Chr 03, Chr 04, Chr 05, Chr 06, Chr 09, Chr 10, Chr 11, and Chr 12 (Fig. 4A). Among these, the chromosomes distribution of the *HVA22* family genes in *S. pennellii* and *S. lycopersicoides* was consistent; they were all distributed on chromosomes Chr 01, Chr 03, Chr 04, Chr 05, Chr 06, Chr 10, Chr 11, and Chr 12. The chromosomal distribution of *HVA22* family genes in *S. pimpinellifolium* differed from that of *S. pennellii* and *S. lycopersicoides*. The *HVA22* family gene in *S. pimpinellifolium* was distributed on chromosome Chr 09; however, no *HVA22* gene family members were found on chromosome Chr 12. We performed intraspecific MCScanX analysis on three species of wild tomatoes to gain a clear understanding of the linear relationships between *HVA22* family genes within species. The results showed five pairs of paralogous homologous genes within the *HVA22* family in *S. pimpinellifolium*: *SpiHVA22o/SpiHVA22e*, *SpiHVA22n/SpiHVA22a*, *SpiHVA22j/SpiHVA22i*, *SpiHVA22h/SpiHVA22d*, and *SpiHVA22f/SpiHVA22b*. Four pairs of paralogous homologous genes existed in *S. lycopersicoides* and *S. pennellii* (*SlydHVA22b/SlydHVA22k*, *SlydHVA22c/SlydHVA22i*, *SlydHVA22e/SlydHVA22h*, and *SlydHVA22g/SlydHVA22p* in *S. lycopersicoides*; *SpHVA22a/SpHVA22l*, *SpHVA22b/SpHVA22n*, *SpHVA22d/SpHVA22g*, and *SpHVA22e/SpHVA22m* in *S. pennellii*) (Fig. 4B). We performed a Ka/Ks analysis of the identified paralogous homologs to enable a comprehensive understanding of the *HVA22* family genes in three species of tomatoes. The final results showed that all 13 pairs of paralogous homologous genes in the *HVA22* families had Ka/Ks greater than 1 (Table 2). This also suggested that the paralogous homologous gene pairs were subjected to stronger environmental stresses, and the gene evolution and protein function were stabilized.

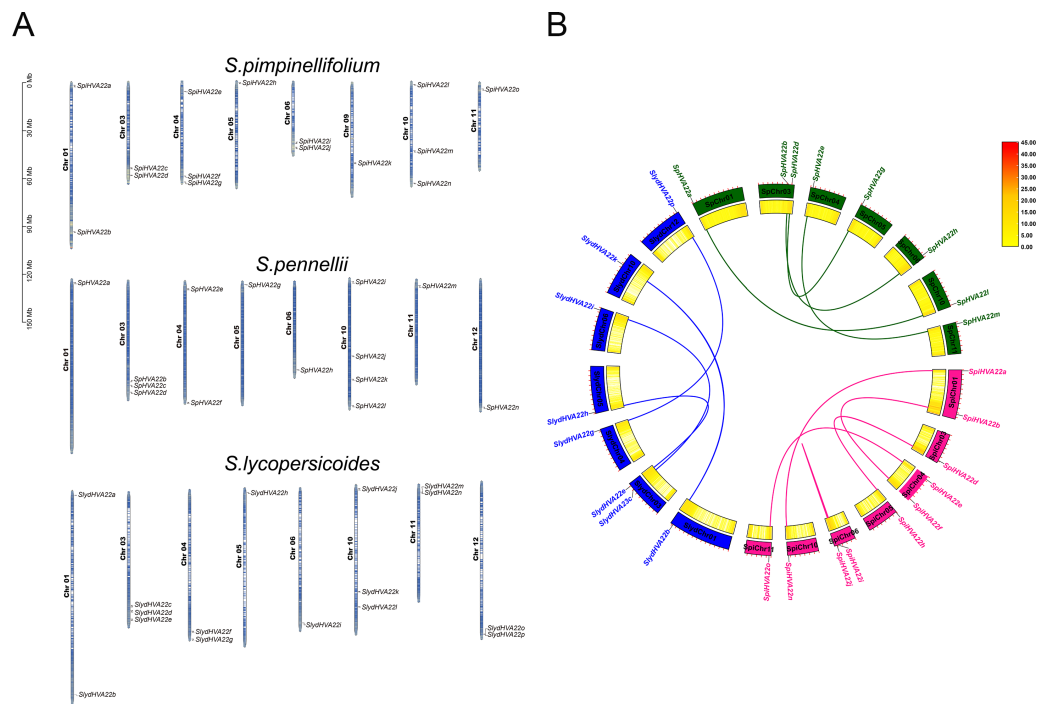


Figure 4 Gene localization to chromosomes and collinearity analysis of *HVA22* within species. (A) Chromosome localization of three species from wild tomato *HVA22* genes. (B) Collinearity analysis of *HVA22* within species. Pink, green, and blue lines, which indicate the collinearity of the *HVA22* gene between *Solanum pimpinellifolium* and *Solanum pennellii*, and *Solanum lycopersicoides* and *Solanum lycopersicoides*, respectively.

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Table 2 The Ka/Ks ratios and date of duplication for duplicate *HVA22* genes in three wild tomatoes.

Species	Duplicated gene pairs	Ka	Ks	Ka/Ks	Selective pressure	Type
<i>S. lycopersicoides</i>	<i>SlydHVA2b/SlydHVA22k</i>	0.80688	1.83965	0.43861	Purify selection	Segmental
	<i>SlydHVA2c/SlydHVA22i</i>	0.43342	1.57164	0.27577	Purify selection	Segmental
	<i>SlydHVA22e/SlydHVA22h</i>	0.22811	3.48793	0.0654	Purify selection	Segmental
	<i>SlydHVA22g/SlydHVA22p</i>	0.18797	0.6627	0.28364	Purify selection	Segmental
<i>S. pimpinellifolium</i>	<i>SpiHVA22a/SpiHVA22n</i>	0.34191	2.79239	0.12244	Purify selection	Segmental
	<i>SpiHVA22b/SpiHVA22f</i>	1.13033	3.13699	0.36032	Purify selection	Segmental
	<i>SpiHVA22d/SpiHVA22h</i>	0.24031	3.49311	0.0688	Purify selection	Segmental
	<i>SpiHVA22e/SpiHVA22o</i>	0.20661	0.78885	0.26192	Purify selection	Segmental
	<i>SpiHVA22i/SpiHVA22j</i>	0.17878	0.9293	0.19239	Purify selection	Segmental
<i>S. pennellii</i>	<i>SpHVA22a/SpHVA22l</i>	0.46107	3.2625	0.14132	Purify selection	Segmental
	<i>SpHVA22b/SpHVA22h</i>	0.18824	1.15914	0.16239	Purify selection	Segmental
	<i>SpHVA22d/SpHVA22g</i>	0.22977	3.50697	0.06552	Purify selection	Segmental
	<i>SpHVA22e/SpHVA22m</i>	0.18263	0.65296	0.27969	Purify selection	Segmental

Evolution and collinearity analysis of the *HVA22* gene family in the three species of wild tomatoes

We performed an interspecific collinearity analysis of *HVA22* family genes in eight Solanaceae species, tobacco, pepper, potato, eggplant, and four species of *Lycopersicon* (*S. lycopersicum*, *S. lycopersicoides*, *S. pennellii* and *S. pimpinellifolium*), based on their divergence times (Wu & Tanksley, 2010), to explore the homology of *HVA22* family genes in Solanaceae plants. Our results showed a significant increase in *HVA22* family homologous genes and a significant acceleration in the rate of evolution from pepper onward. The distribution of *HVA22* homologs was similar on the chromosomes of the remaining Solanaceae members except for tobacco. The *HVA22* genes on chromosomes Chr 03, Chr 04, Chr06, and Chr10 in Solanaceae (excluding tobacco) had high homology. A large similarity and homology were seen in the chromosomal distribution of *HVA22* genes in four species of *Lycopersicon* (Fig. 5A). To gain further insight into the homology of *HVA22* family genes in Solanaceae, we performed a collinearity analysis of *HVA22* family genes in Solanaceae plants one by one; the analysis was also performed in the four species of *Lycopersicon* by the same method. The final results showed that most of the *HVA22* family genes in the three species wild tomatoes were orthologous to each other and the cultivated tomato *HVA22* family genes. Among these, the cultivated tomato *HVA22* genes were found to have corresponding homologs in all three species of wild tomatoes, except *SIHVA22o*, which was not identified as a direct homolog in *S. piminellifolium* (Fig. 5B). The distribution of the *HVA22* family genes on the chromosomes of the four tomato species was also highly similar. The *HVA22* genes on chromosome Chr03 were found to be highly homologous to each other in the co-linearity analysis of Solanaceae (tomato, pepper, potato, and eggplant). However, the *SIHVA22a* and *SIHVA22e* genes located on chromosomes SlChr01 and SlChr04 were found to be orthologous only in eggplant (*SmHVA22a* and *SmHVA22l*), while no orthologous genes were found in pepper and potato. The orthologous genes of *SIHVA22e*, *SIHVA22h*, and *SIHVA22n* located on chromosomes SlChr04, SlChr05, and SlChr11 were present in potato (*StHVA22v*, *StHVA22o*, and *StHVA22v*) and eggplant (*SmHVA22l*, *SmHVA22c*, and *SmHVA22l*); their orthologous genes were not detected in pepper (Fig. 5C).

Protein–protein network analysis of *HVA22* family genes in tomato

We constructed a protein–protein network expression profile of tomato *HVA22* family genes using the STRING database to investigate the interactions between *HVA22*-like proteins and other proteins. Our results showed interactions between *HVA22* family member proteins in tomatoes. Some *HVA22* family proteins (*SIHVA22o*, *SIHVA22g*, *SIHVA22m*, and *SIHVA22a*) interacted with ubiquitin-conjugating enzyme (ubiquitin-conjugating enzyme 13 E2), RNA-binding protein (RNA-binding protein 3.1), and eukaryotic translation initiation factor (ETA). Among these, *SIHVA22j* and *SIHVA22d* interacted with the vesicle-sorting protein (vacuolar protein sorting protein 25). The remaining *HVA22* family proteins (*SIHVA22f*, *SIHVA22k*, *SIHVA22e*, *SIHVA22n*, *SIHVA22b*, *SIHVA22i*, *SIHVA22h*, *SIHVA22l*, and *SIHVA22c*) interacted with proteins

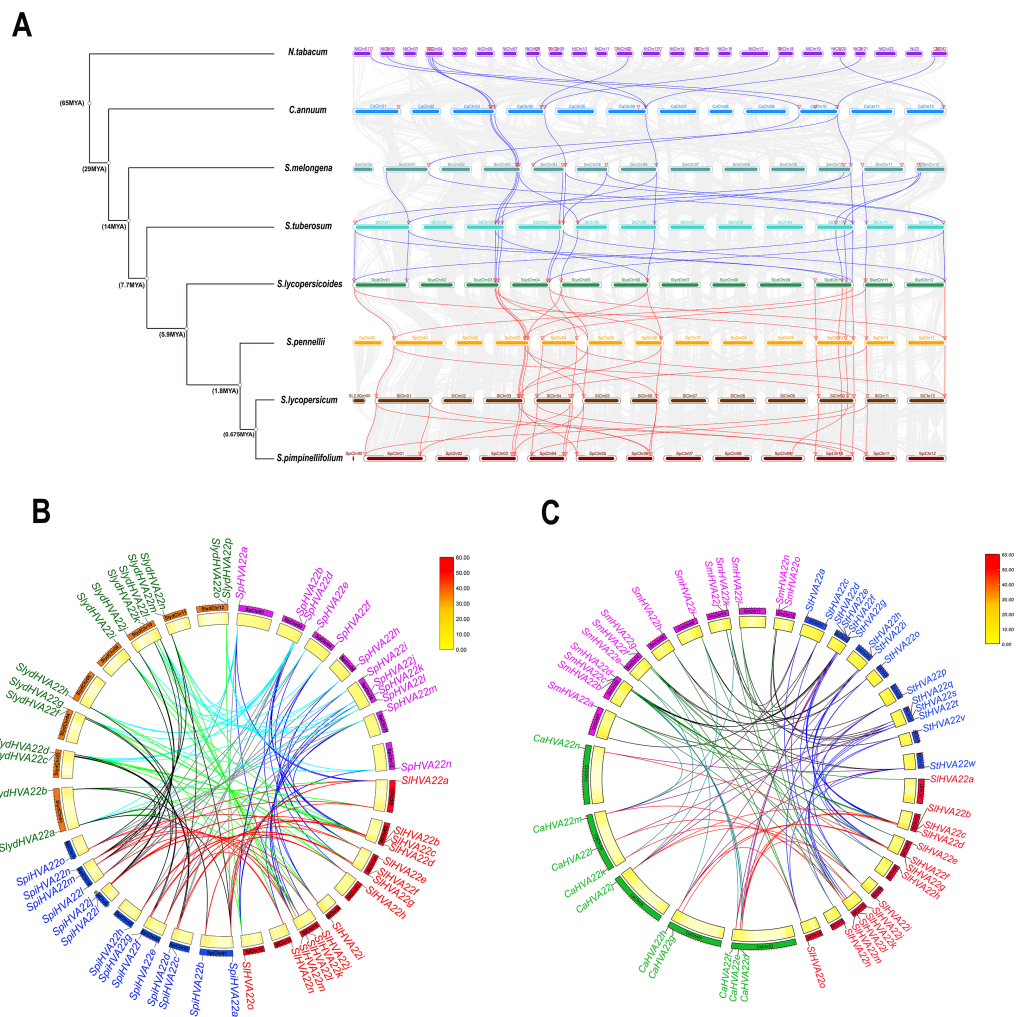


Figure 5 Homologous genes and evolutionary analysis of the HVA22 family. (A) Co-lineage map for Solanaceae species, with species genomes arranged in evolutionary order and colored lines representing HVA22 genes with direct homologous relationships within each species. (B) Collinearity of HVA22 genes within *Solanum pimpinellifolium*, *Solanum lycopersicoides*, *Solanum pennellii*, and *Solanum lycopersicum*, with the outer circle showing the chromosomes of each species, the inner circle showing the gene density, the two ends of the lines representing the direct homologous HVA22 genes, and the different colors indicate comparisons between different Lycopersicon. (C) Co-lineage map of HVA22 genes within Solanaceae (*C. annuum*, *S. melongena*, *S. tuberosum*, and *S. lycopersicum*), with the outer circle showing the chromosomes of each Solanaceae, the inner circle showing gene density, the ends of the lines representing direct homologous HVA22 genes, and the different colors indicate comparisons between different species.

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from their own family; interactions between them and with other proteins were not observed (Fig. 6).

Expression of tomato HVA22 gene in different tissues

We used published RNA-seq data to map the gene expression heat map of the HVA22 gene in different tissues of *S. pimpinellifolium* to better understand the role of the HVA22 gene in the growth and development of wild tomatoes. The results showed that most of

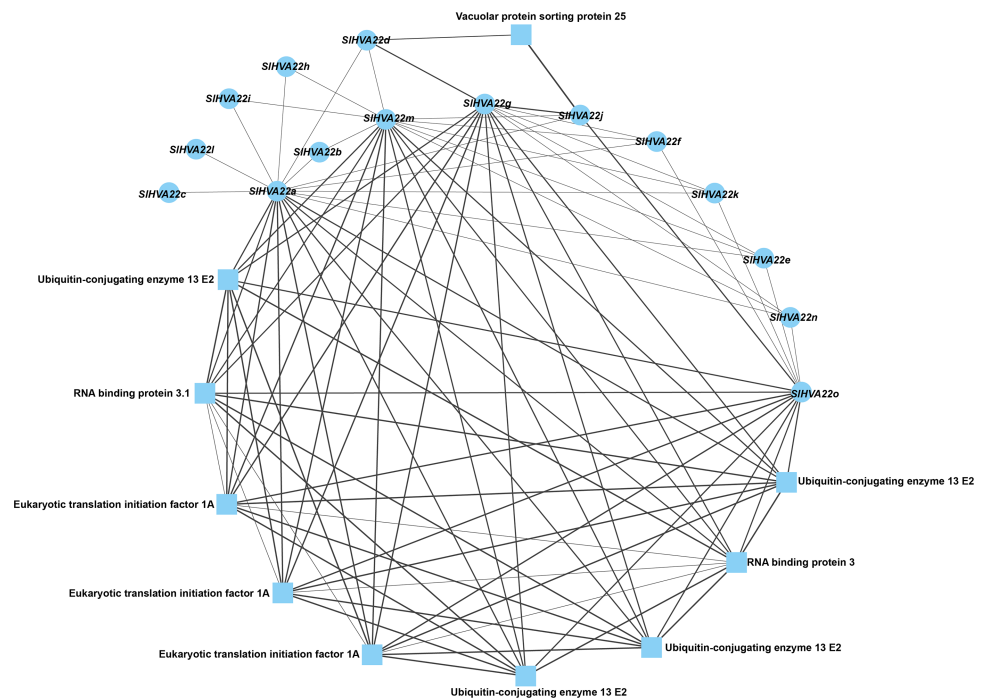


Figure 6 Interaction network of the HVA22 family with other proteins. Each node represents a protein, each connecting line represents the presence of an interaction, and the thickness of the line represents the value of the composite score, with circles representing HVA22 proteins and squares representing other proteins.

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the *HVA22* family members in wild tomatoes were expressed at a lower level in leaves and fruits. In particular, *SpiHVA22c*, *SpiHVA22k*, *SpiHVA22a*, and *SpiHVA22g* family member genes were highly expressed in ripening fruits. In addition, among the wild tomato *HVA22* family genes, *SpiHVA22d* was slightly more expressed in leaves compared with other *HVA22* family members. The roots showed the highest expression of the remaining *HVA22* family members, with the exception of *SpiHVA22g*. The expression of *HVA22* family genes in group I was generally low in flower buds. However, the *HVA22* family member genes in group II showed higher expression levels in the flower buds. In addition, the *HVA22* member genes in group II also showed higher expression levels in the stems. Overall, the expression of tomato *HVA22* family genes was concentrated in roots, flowers, and developing fruits (Fig. 7). The high expression of some *HVA22* family genes in roots also confirmed the possibility that *HVA22* family genes were involved in abiotic stress processes in tomatoes. This result provided important clues for our study of the function of the *HVA22* family gene in tomatoes. We treated *S. pimpinellifolium* seedlings with NaCl, PEG6000, and hormones (ABA and MeJA) to be more confident about how well the wild tomato *HVA22* family genes could respond to salt, drought, and hormones. The expression of *SpiHVA22c*, *SpiHVA22d*, *SpiHVA22g*, *SpiHVA22k*, and *SpiHVA22j* genes in the leaves of the plants obtained from the treatments was analyzed using qRT-PCR. Our results showed that the expression of *SpiHVA22c*, *SpiHVA22d*, *SpiHVA22g*, and *SpiHVA22k* was

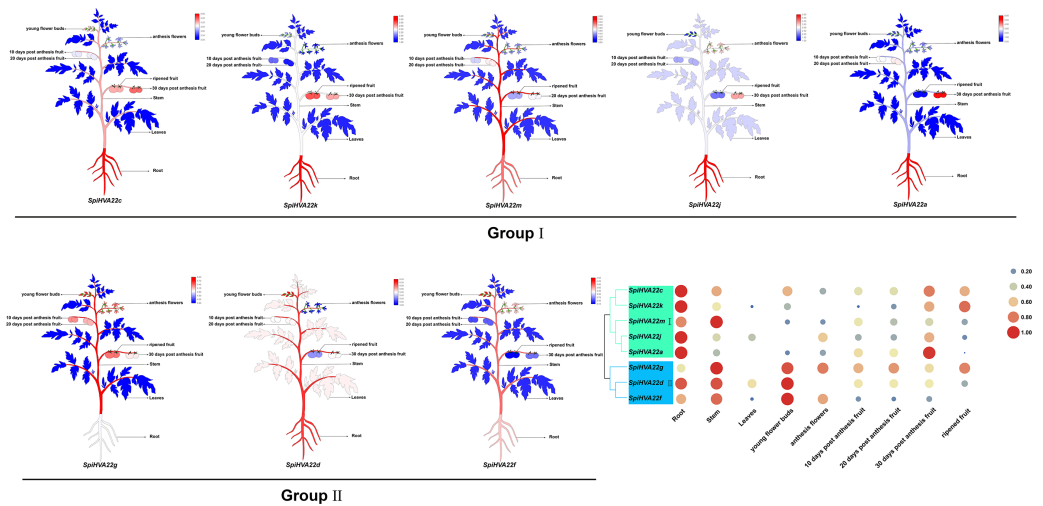


Figure 7 Heat map of tissue-specific expression of HVA22 genes in *Solanum pimpinellifolium*. Root, stem, leaves, young flower buds, anthesis flowers, 10 days post anthesis fruit, 20 days post anthesis fruit, 30 days post anthesis fruit, ripened fruit.

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significantly higher under NaCl and PEG6000 treatment than in the control group. The expression of the *SpiHVA22j* gene was the highest after 2-h treatment and showed lower expression than that in the control group as the treatment time was extended. At the same time, the expression of *SpiHVA22k*, *SpiHVA22g*, and *SpiHVA22c* genes under ABA and MeJA treatment was similar to that under PEG6000 and NaCl treatment. Intriguingly, the *SpiHVA22g* and *SpiHVA22j* gene showed negative regulation under MeJA treatment and positive regulation under ABA treatment (Fig. 8). Apart from this, no significant change was found in the expression of the *SpiHVA22d* gene under ABA treatment, indicating that the *SpiHVA22d* gene was not regulated by the hormone ABA.

DISCUSSION

With advances in gene sequencing technology, next-generation sequencing has improved the accuracy of the genome, thereby avoiding annotation errors in individual gene sequences by whole-gene sequencing. This has also facilitated genetic improvement and directed breeding in tomatoes (Rothan, Diouf & Causse, 2019). The HVA22 gene is commonly found in eukaryotes (Lu, 2013). It is expressed in plants in different tissues such as seeds, stems, and roots and is induced under several environmental stress conditions (e.g., cold, salt, and drought) mediated by ABA (Shen et al., 2001). The HVA22 gene family has been reported in *A. thaliana* (Chen et al., 2002), *C. clementina* (Gomes Ferreira et al., 2019), and cultivated tomatoes (Wai et al., 2022), but has not been reported in wild tomatoes (*S. lycopersicoides*, *S. pennellii*, and *S. pimpinellifolium*). The results of the phylogenetic tree showed that the three wild tomato HVA22 family genes were divided into two clades, which were further divided into four groups (Fig. 1A). This result was also consistent with the classification of tomato HVA22 family genes (Wai et al., 2022). Our results showed that the three wild tomato HVA22 family genes in each group were well assigned to the known HVA22 groups

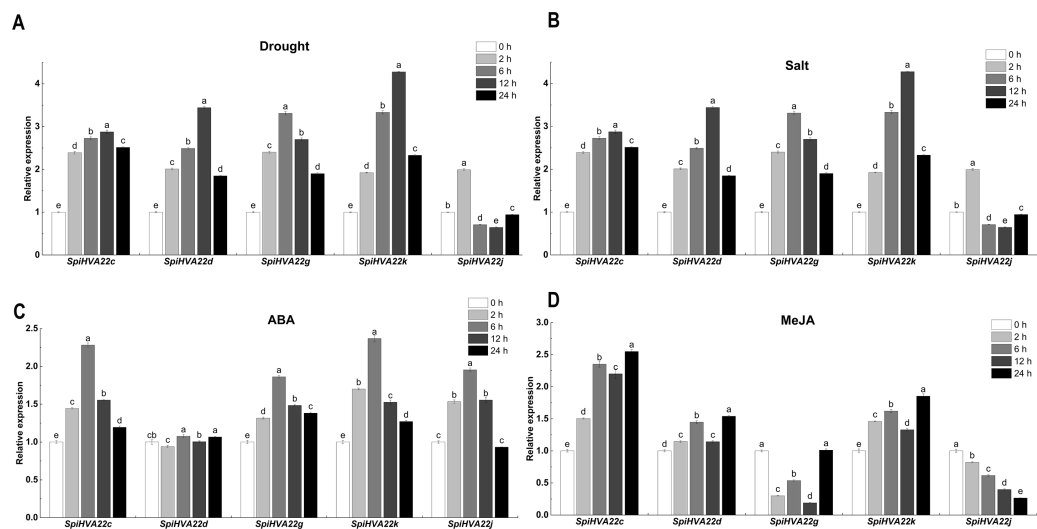


Figure 8 QRT-PCR validation of the *Solanum pimpinellifolium* HVA22 genes under abiotic stress and hormone induction. The standard deviations are shown with error bars.

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of *A. thaliana* and cultivated tomatoes. This further demonstrated the highly conserved nature of the HVA22 family of genes in *Lycopersicon* crops. We constructed a phylogenetic tree using HVA22 family genes from seven species of Solanaceae to characterize the HVA22 family genes more comprehensively. The results showed that the HVA22 family genes in seven species of Solanaceae were divided into three major branches (Fig. 1B). Although the number of HVA22 family members varies significantly among Solanaceae, they all carry the TB2/DP1 structural domain specific to HVA22-like. In addition, some of the HVA22 proteins have other structural domains besides TB2/DP1, and HVA22 proteins with other structural domains are clustered together in a phylogenetic tree. This also illustrates the reliability of the phylogenetic structure and the high homology of the HVA22 gene in Solanaceae. This homology may result from the TB2/DP1 structural domain specific to the HVA22 protein. In addition, the analysis of the physicochemical properties of HVA22 family proteins in the three species of wild tomatoes found that the average number of amino acids was 231 (*S. pimpinellifolium*), 208 (*S. pennellii*), and 217 (*S. lycopersicoides*) (Table 1). These were close to the 230 amino acids in tomato HVA22 (Wai et al., 2022), but showed a large difference from the 130 amino acids of barley (Shen et al., 2001) and 151 amino acids of *C. clementina* (Gomes Ferreira et al., 2019), which might be related to the large evolutionary distance between different species. Interestingly, we found that the chromosomal distribution positions of the HVA22 family genes in the three species of wild tomatoes showed striking similarities (Fig. 4A). At the same time, these HVA22 proteins with similar distributions on the chromosome showed nearly identical physicochemical properties (e.g., theoretical pI, molecular weight, aliphatic index, and grand average of hydropathicity) (Table 1). This also fully corroborated the conserved nature of the HVA22 family genes in tomatoes.

The prediction of subcellular localization of HVA22 helps us understand the function of this gene. Our findings regarding the subcellular localization of *HVA22* genes in the three species of wild tomatoes revealed that a majority of the *HVA22* genes were predicted to be localized in the endoplasmic reticulum, chloroplast, cytoplasm, and nucleus (Table 1). Evidence showed that after heterologous expression, tomato SIHVA22a, SIHVA22f, and SIHVA22n proteins in rice protoplasts were observed to be localized to the endoplasmic reticulum (Wai et al., 2022). In addition, the rice *HVA22* family gene *OsHLP1* promoted the mechanism of disease resistance by compromising endoplasmic reticulum homeostasis when plants were infected by pathogens (Meng et al., 2022). In yeast, the *HVA22* gene homolog Yop1p played an important role in transporting material between the endoplasmic reticulum and the Golgi apparatus (De Antoni et al., 2002). These findings consistently showed that HVA22 played an important role as an endoplasmic reticulum localization protein in the life activities of eukaryotic cells. The HVA22 protein in the three species of wild tomatoes also seemed to exhibit similar characteristics.

The diversity of exon–intron structures is one of the key factors in the evolution of gene families and underpins the structure of phylogenetic trees (Shiu & Bleecker, 2003; Wang et al., 2014). The structural analysis of the *HVA22* family of genes in the three species of wild tomatoes showed that the number of exons was similar in groups I and II; however, a large difference was found in the number of exons in group III (Fig. 2). This also implied that the *HVA22* gene in the three species of wild tomatoes could be transcriptionally diversified through processes such as selective splicing to regulate a more complex and broad range of functions. A total of 20 motifs were identified in the structural analysis of the *HVA22* family genes in wild tomatoes, among which Motifs 1, 2, and 3 are important motifs that made up the TB2/DP1 structural domain of the *HVA22* gene. A membrane protein TB2-like1 exists in animals and belongs to the same TB2/DP1/*HVA22* family of proteins. This protein may play an important role in the cell membrane transport of retinal ganglion cells (Sato et al., 2005). The Yop1p gene in yeast (*Saccharomyces cerevisiae*) is homologous to the barley *HVA22* gene, and the proteins it encodes have the TB2/DP2 structural domain (Brands & Ho, 2002). Previous studies showed that yeast Yop1p was an integral membrane protein with a hydrophilic region at its N-terminal end. This region bound specifically to the yeast Yip1p protein, and Yop1p interacted with Yip1p to mediate the biological process of intracellular membrane transport (Calero, Whittaker & Collins, 2001). Also, this membrane protein played a critical physiological role in barley, yeast, and humans (Sato et al., 2005; Voeltz et al., 2006). In a study of the TB2/DP1 structural domain of the *HVA22* protein of barley dextrin, the TM1, TM2, and TM3 segments of the *HVA22* gene were separately deleted using truncating mutations, so that the separately deleted *HVA22* genes would all carry their green fluorescent protein (GFP) tags, which were transferred to barley dextrin cells for cellular sublocalization observation. It was found that the sublocalization of the deletion of TM2 presented a results significant differences compared with *HVA22::GFP* (Guo & David Ho, 2008). It further indicated that *HVA22* had the properties of a membrane protein. It also showed that the barley *HVA22* protein had a transmembrane region in the TB2/DP1 structural domain, and the presence of this transmembrane region provided a theoretical basis for the localization of the *HVA22*

protein to the organelle membrane. This result was similar to our predicted subcellular localization of the *HVA22* gene in the three species of wild tomatoes. This also suggested that the *HVA22* gene with the TB2/DP1 structural domain might play an important role in intracellular vesicle transport (Brands & Ho, 2002). However, no relevant studies have been reported on the involvement of the *HVA22* gene in intracellular vesicle transport in tomatoes. Moreover, the Zf-met structural domain was found in the amino acid sequence of *HVA22* in the three species of wild tomatoes. The Zf-met domain is another zinc-finger domain containing the CxxC(12)Hx(6)H motif, which is associated with RNA binding (Yadav, Fernández-Baca & Cannon, 2020). This is similar to the structure identified in the tomato *HVA22* protein. This further indicates that *HVA22* family members are conserved in *Lycopersicon*.

Cis-acting elements are noncoding DNA sequences present in the promoter region of a gene. The distribution of different types of *cis*-acting elements in the promoter region may determine gene regulation and functional roles (Hernandez-Garcia & Finer, 2014). In this study, we characterized a 2,000-bp promoter sequence upstream of the *HVA22* family genes in the three species of wild tomatoes (Fig. 3). The *cis*-acting elements of *HVA22* family genes in the three species of wild tomatoes were divided into four main categories: *cis*-acting elements involved in light response, *cis*-acting elements involved in phytohormone response, *cis*-acting elements involved in biotic/abiotic stress, and *cis*-acting elements involved in growth and development. Common *cis*-acting elements associated with light response were ACE, MRE, G-box, GT1-Mofit, Sp1, 4cl-CMA2b, 3-AF1-binding site, and AAAC-Motif. These light-responsive *cis*-acting elements played an important regulatory role in stress response and growth and development of plants (Kaur et al., 2017). The *cis*-acting elements responding to phytohormones in the promoter of the *HVA22* gene family of three species of wild tomatoes were more widely distributed, including gibberellin, ABA, methyl jasmonate, ethylene, and salicylic acid. This further suggested that the *HVA22* gene might be involved in the life activities of tomatoes through a hormone-regulated network, which was consistent with the findings in tomatoes and *C. clementina* (Gomes Ferreira et al., 2019; Wai et al., 2022). In addition, *cis*-acting elements involved in multiple stresses were predicted in the *HVA22* promoter in the three species of wild tomatoes, such as MBS (drought-inducible), LTR (low temperature responsive), and TC-rich repeats (defense and stress responsive). The *HVA22* gene response to low temperature and drought stress has been reported in *A. thaliana* (Chen et al., 2002), barley (Shen et al., 2001), and rice (Zhao et al., 2021). Six *cis*-acting elements associated with growth and development were identified in the promoters of *HVA22* family genes in the three species of wild tomatoes, including RY-element (involved in seed-specific regulatory element), MSA-like (involved in cell cycle regulatory element), GAT-box (involved in meristematic tissue expression element), circadian (involved in circadian control regulatory element), AACAA_motif (involved in the endosperm-specific negative expression), and GCN4_motif (involved in endosperm expression element) (Zhang et al., 2013). The *HVA22* gene promoter contained *cis*-acting elements associated with plant development, particularly involved in seed-specific regulatory elements, which might be relevant to the function of the *HVA22* gene in seed maturation and dormancy in barley

(Guo & David Ho, 2008). However, further experimental evidence is needed by cloning the upstream promoter of the *HVA22* gene in the three species of wild tomatoes to obtain the corresponding experimental evidence and provide direction for the next study of the *HVA22* gene function.

Gene duplication provides the material basis for plant evolution and the generation of new functions (Huang et al., 2022). It occurs in several different modes, such as whole-genome duplication, single-gene duplication, and segmental duplication (De Bodt, Maere & Van de Peer, 2005; Paterson et al., 2010). The chromosomal localization and collinearity analysis of the *HVA22* gene in the three species of wild tomatoes showed spacer regions in the physical location of the *HVA22*-encoding gene on the chromosome in the three species of wild tomatoes (Fig. 4A). Thus, it was tentatively determined that the *HVA22* gene was amplified in the three species of wild tomatoes mainly by large-scale segmental duplication or whole-genome duplication of the gene family. Subsequently, the intraspecific collinearity analysis of the *HVA22* family genes of the three species of wild tomato showed that the *HVA22* genes were amplified in the three species of wild tomatoes in an all-segmental duplication manner, which was consistent with our previous speculation on the distribution of *HVA22* genes on chromosomes (Fig. 4B). At the same time, the same results were obtained in cultivated tomatoes (Wai et al., 2022). The ratio of nonsynonymous to synonymous substitutions reflects, to some extent, the selective pressure of gene evolution. $Ka/Ks > 1$ represents positive selection for accelerated evolution, and $Ka/Ks < 1$ represents the presence of purifying selection for gene duplication (Wang et al., 2010). The *HVA22* paralogous homologous gene pairs in all three wild tomatoes had $Ka/Ks < 1$, which further suggests that these homologous gene pairs underwent more intense environmental selection pressure and exhibited functional homogeneity during evolution. Genome-wide duplication events (WGD) have long been recognized as an important evolutionary force in species formation, adaptation to the environment, and shaping of species diversity (Wood et al., 2009; Soltis & Soltis, 2016). Dicotyledons experienced γ events in gene duplication; however, Solanaceae members experienced another WGD event (T event) about 65 million years ago (Knapp, 2012; Wu, Han & Jiao, 2020). *HVA22* genes were added or lost in Solanaceae driven by genome-wide duplication events. As the Solanaceae diverged during evolution, the *HVA22* family gene members showed a decreasing trend and then an increasing trend in Solanaceae (Fig. 5A). This might be related to the contraction and expansion of family genes in Solanaceae when they experienced the most recent T event. The *HVA22* gene on SlChr 03 in tomatoes is the most conserved member of the Solanaceae family (Fig. 5B). For *HVA22* family genes present only in tomatoes but not in other members of Solanaceae, this might be due to gene deletions caused by genome-wide duplication events (Fig. 5C). The result that *HVA22* family genes were highly conserved in the four species of *Lycopersicon* also indicated the importance of *HVA22* genes in the life activities of tomatoes.

The ubiquitin–proteasome system (UPS) regulates various biological functions in plants, such as hormonal responses (Chen et al., 2018; He et al., 2018), abiotic stress responses (Kim, Jang & Seo, 2016; Shu & Yang, 2017), plant growth and development (Cho et al., 2011; Koops et al., 2011), circadian rhythms (Gil et al., 2017), and plant immune

responses (Lin et al., 2008; Luo et al., 2010). The predicted results of tomato *HVA22* protein interactions indicated that the *SlHVA22* protein was involved in plant signaling and the regulation of plant growth and development (Fig. 6). Among these, the tomato *HVA22* protein was co-expressed with ubiquitin-binding enzymes, suggesting that the tomato *HVA22* gene might be involved in the ubiquitination and regulation of tomato growth and development and abiotic stresses. In addition, the vesicle sorting protein (VPS) (Xiang, Etxeberria & Ende, 2013) plays an important role in the protein sorting pathway as an important protein for the formation of endosomal sorting complex protein (ESCRT) (Gao et al., 2017). Previous studies showed that the growth hormone transporter proteins PIN1, PIN2, and AUX1 were the cargo proteins of ESCRT. To some extent, this suggested that VPS played a role in hormone signaling (Spitzer et al., 2009). The results of the co-expression of *HVA22* protein with VPS suggested that the *HVA22* protein might be a cargo membrane protein of ESCRT and played an important role in intracellular vesicle transport. RNA-binding proteins interacted with RNA through the RNA-binding domain to regulate RNA metabolism and function. Conversely, RNA could bind to RAN-binding proteins and affect their lifespan and function (Hentze et al., 2018). The co-expression of RNA-binding proteins with the tomato *HVA22* protein might indicate an important contribution of RNA-binding proteins in maintaining the function and longevity of the *HVA22* gene. The results of co-expression of the tomato *HVA22* gene with eukaryotic translation initiation factor 1A also further confirmed the conclusion that the *HVA22* gene was present only in eukaryotes. Recently, that the AtGCN2 activation of eukaryotic translation initiation factor 2 phosphorylation was shown to be another key component in response to endoplasmic reticulum stress in *A. thaliana*, and it played an important role in the signaling process of the unfolded protein response (Afrin et al., 2020; Howell, 2021; Liu, Afrin & Pajerowska-Mukhtar, 2019).

The *HVA22* gene was differentially expressed in different tissues of *S. pimpinellifolium*, while the expression of individual *HVA22* genes showed an increasing trend during *S. pimpinellifolium* fruit development (Fig. 7). The root system, as the main tissue directly sensing drought and salt ions, has a high sensitivity to drought and salt ions. The high expression of the *HVA22* gene in roots indicated that the *HVA22* gene played an important role in plant root development and resistance to abiotic stress response. The expression of *HVA22* family genes showed an increasing trend as the fruit grew during fruit development. The same result was shown in the *HVA22* family genes in cultivated tomatoes (Wai et al., 2022). This might be related to the involvement of *HVA22* family genes in tomato fruit growth and development. We treated *S. pimpinellifolium* seedlings with NaCl, PEG6000, and hormones, thereby performing qRT-PCR analysis of the *HVA22* family genes in *S. pimpinellifolium* (Fig. 8). The results showed that *SpiHVA22d*, *SpiHVA22g*, *SpiHVA22k*, and *SpiHVA22c* genes responded positively to salt and drought stresses. This was also similar to the characterization of *HVA22s* described in barley, *A. thaliana*, *C. clementina*, and cultivated tomatoes (Wai et al., 2022). In addition, the responses of *SpiHVA22k*, *SpiHVA22c*, and *SpiHVA22d* genes to ABA and MeJA also suggested that *SpiHVA22k*, *SpiHVA22c*, and *SpiHVA22d* genes played important roles in the ABA and MeJA pathways. Interestingly, the *SpiHVA22j* gene in the *S. pimpinellifolium* *HVA22* family of genes

showed negative regulation under MeJA treatment. This might be related to the fact that the *SpiHVA22j* gene was not involved in the plant jasmonic acid pathway. Previous studies in barley also pointed to the role of the *HVA22* gene as an early ABA-inducible gene (Shen *et al.*, 2001). Our qRT-PCR results further concluded that the *S. pimpinellifolium HVA22* gene might be involved in the plant ABA pathway, which in turn responded to plant regulation of abiotic stresses.

CONCLUSIONS

In the present study, we systematically identified *HVA22* family genes in tomatoes. We used a bioinformatics approach to describe the physicochemical properties, gene structure, *cis*-acting elements, and protein interactions of different *HVA22* genes. The expansion and contraction of *HVA22* family genes during the evolution of Solanaceae species were also discussed. The expression profile data of different tissues of *HVA22* family genes in tomatoes showed that the expression of *HVA22* family genes was mainly concentrated in roots, flowers, and developing fruit. We validated five genes in the tomato *HVA22* family using qRT-PCR. Four of these genes were involved in ABA- and MeJA-mediated regulatory pathways, and played important roles in tomato resistance to abiotic stresses (salt and drought). These results laid the foundation for further investigation of the function of *HVA22* family genes in tomatoes.

ADDITIONAL INFORMATION AND DECLARATIONS

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Competing Interests

The authors declare there are no competing interests.

Author Contributions

- LaiPeng Zhao conceived and designed the experiments, prepared figures and/or tables, authored or reviewed drafts of the article, and approved the final draft.

- Baike Wang performed the experiments, prepared figures and/or tables, and approved the final draft.
- Tao Yang analyzed the data, prepared figures and/or tables, and approved the final draft.
- Huizhuan Yan analyzed the data, prepared figures and/or tables, and approved the final draft.
- Qinghui Yu conceived and designed the experiments, authored or reviewed drafts of the article, and approved the final draft.
- Juan Wang performed the experiments, authored or reviewed drafts of the article, and approved the final draft.

Data Availability

The following information was supplied regarding data availability:

The raw data is available in the [Supplemental Files](#).

Supplemental Information

Supplemental information for this article can be found online at <http://dx.doi.org/10.7717/peerj.14844#supplemental-information>.

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