

Identification and expression analysis of xyloglucan endotransglucosylase/hydrolase (XTH) family genes in grapevine (*Vitis vinifera* L.)

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Xyloglucan endotransglucosylases/hydrolases (XTH) are key enzymes in cell wall reformulation. They have the dual functions of catalyzing xyloglucan endotransglucosylase (XET) and xyloglucan endonuclease (XEH) activity and play a crucial role in the responses against abiotic stresses, such as drought, salinity, and freezing. However, a comprehensive analysis of the *XTH* family and its functions in grapevine (*Vitis vinifera* L.) has not yet been completed. In this study, thirty-four *XTHs* were identified in the whole grapevine genome and then named according to their distribution on chromosomes. Based on a phylogenetic analysis including *Arabidopsis XTHs*, the *VvXTHs* were classified into 3 groups. *Cis*-element analysis indicated that these family members are related to most abiotic stresses. We further selected 14 *VvXTHs* from different groups and then examined their transcription levels under drought and salt stress. The results indicated that the transcription levels of selected *VvXTHs* in the leaves and roots presented the largest changes, suggesting that *VvXTHs* are likely to take part in the responses to drought and salt stress in grapevines. These results provide useful evidence for the further investigation of *VvXTHs* function in response to abiotic stresses in grapevine .

1 **Identification and expression analysis of the**
2 **xyloglucan endotransglucosylase/hydrolase (XTH)**
3 **family in grapevine (*Vitis vinifera* L.)**

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20 **Abstract**

21 Xyloglucan endotransglucosylases/hydrolases (XTH) are key enzymes in cell wall reformulation.
22 They have the dual functions of catalyzing xyloglucan endotransglucosylase (XET) and
23 xyloglucan endonuclease (XEH) activity and play a crucial role in the responses against abiotic
24 stresses, such as drought, salinity, and freezing. However, a comprehensive analysis of the *XTH*
25 family and its functions in grapevine (*Vitis vinifera* L.) has not yet been completed. In this study,
26 thirty-four *XTHs* were identified in the whole grapevine genome and then named according to their
27 distribution on chromosomes. Based on a phylogenetic analysis including *Arabidopsis XTHs*, the
28 *VvXTHs* were classified into 3 groups. *Cis*-element analysis indicated that these family members
29 are related to most abiotic stresses. We further selected 14 *VvXTHs* from different groups and then
30 examined their transcription levels under drought and salt stress. The results indicated that the
31 transcription levels of selected *VvXTHs* in the leaves and roots presented the largest changes,
32 suggesting that *VvXTHs* are likely to take part in the responses to drought and salt stress in
33 grapevines. These results provide useful evidence for the further investigation of *VvXTHs* function
34 in response to abiotic stresses in grapevine.

35

36 **Subjects:** Agricultural Science, Molecular Biology, Plant Science

37

38 **Keywords:** *VvXTH*, *Vitis vinifera*, phylogenetic analysis, expression pattern, abiotic stress

39 Introduction

40 As one of the most economically fruit crops, Grapevine (*Vitis vinifera* L.) is cultivated worldwide
41 (*Feng et al., 2000*). However, the growth of grapes in natural environment is inevitably impacted
42 by a series of abiotic stresses, including salinity, drought, and extreme temperatures, which damage
43 the cell walls of the plants, disrupt the biofilm system, and ultimately affect the quality and yield
44 of the fruit (*Araujo et al., 2016; Liu et al., 2019; Ning et al., 2017*). Xyloglucan
45 endotransglucosylase/hydrolase (XTH) can carry out cell wall structural modification and
46 rearrangement by severing and repolymerizing cellulose mono-xyloglucan cross-linked structures
47 (*Campbell et al., 2010*). It belongs to the glycoside hydrolase 16 (GH16) family, which is a
48 subfamily of glycoside hydrolases containing diverse enzymes with different specific targets, such
49 as keratan sulfate, β -1,3-glucans, mixed linkage β -1,3(4)-glucans, xyloglucans, j-carrageenan, and
50 agarose (*Mark et al., 2009; Stratilova et al., 2020*). All XTH proteins present the typical structure
51 of XTH enzymes: the (D/N)E(I/L/A/V/F)(D/T)(F/I)E(F/I/L)LG motif, which includes the catalytic
52 active-site residues ExDxE (*Matsui et al., 2005; Liu et al., 2007; Miedes et al., 2009; Singh et al.,*
53 *2011*). XTH proteins may present one or two enzyme activities: xyloglucan endonuclease (XEH)
54 activity and/or xyloglucan endotransglucosylase (XET) activity. The former specifically
55 hydrolyzes xyloglucan β -1,4 glycosidic bonds, cleaving the xyloglucan chain, thereby shortening
56 the xyloglucan chain, and the latter can transfer xyloglucan fragments between xyloglucan chains,
57 elongating xyloglucan chains (*Han et al., 2016*).

58 Thus far, the XTH family has been identified and analyzed in species such as *Arabidopsis*
59 *thaliana* (33), *Hordeum vulgare* (24), *Glycine max* (61), and *Nicotiana tabacum* (54). (*Nomchit et*
60 *al., 2010; Li et al., 2018; Meng et al., 2018; Cheng et al., 2021; Strohmeier et al., 2004; Tiika et*
61 *al., 2021*). The XTH family was initially classified into three groups, named groups I, II, and III,

62 in *Arabidopsis* (Campbell et al., 1999). However, a subsequent study in *Oryza sativa* found that
63 there was no clear distinction between groups I and II; therefore, rice *XTHs* were divided into only
64 2 groups: group I/II and group III (Eklof et al., 2010). The *XTH* members in group III could be
65 further divided into two subgroups (IIIA and IIIB) according to their three-dimensional structures
66 (Baumann et al., 2007; Fu et al., 2019). Moreover, a small outlier group was identified close to
67 the root of the tree and was named the ancestral group. The *XTHs* of group I/II and group IIIB
68 showed primarily or only XET activity, while the *XTHs* in group IIIA mainly displayed XEH
69 activity (Eklof and Brumer 2010; Nomchit et al., 2010; Opazo et al., 2017). Further studies
70 revealed that each type of enzyme activity was determined by several structural characteristics.
71 For example, in the protein structure of *TmNXG1*, loop 2 is the key structure affecting hydrolysis
72 and transglycosylase activity (Mark et al., 2009). *PttXET16-34* contained an important N-glycan
73 structure, which is found in all group I/II members but absent in almost all IIIA groups (such as
74 *TmNXG1*). Interestingly, the N-glycosylation site shifts from the C-terminus to the other side of
75 the active-site cleft in group IIIB (Mark et al., 2009; Eklof et al., 2010).

76 Increasing evidence has revealed that *XTHs* are instrumental for coping with abiotic stresses
77 through cell remodeling and enhanced cell wall biogenesis in plants (Eklof and Brumer, 2010;
78 Tiika et al., 2021). For instance, the constitutive expression of *CaXTH3* has been verified to
79 enhance resistance to salinity and drought stress in tomato plants (Choi et al., 2011). *AtXTH11*,
80 *AtXTH29*, and *AtXTH33* were observed to be upregulated through different secretory pathways in
81 *Arabidopsis* seedlings treated with heat stress and drought stress (Caroli et al., 2021). A recent
82 study revealed that the overexpression of persimmon *DkXTH1* promotes tolerance to salt and
83 drought stress by improving photosynthesis and reducing lipid peroxidation (Han et al., 2017).
84 Additionally, transgenic tobacco with estradiol-inducible expression of *SIXTH10* shows stronger

85 growth under salinization and hypothermia conditions (*Norbert et al., 2020*), and *GmXTH*
86 expression levels have been reported to be significantly associated with flooding stress (*Li et al.,*
87 *2018*). Transgenic soybeans overexpressing *AtXTH31* also exhibit increased tolerance to flooding
88 stress (*Li et al., 2018*). Moreover, an *AtXTH19* mutant was demonstrated to show lower freezing
89 tolerance during cold and subzero acclimation than the wild type, which is likely related to
90 differences in the cell wall composition and structure (*Daisuke et al., 2020*).

91 Taken together, the above studies highlight the essential functions of *XTHs* in resisting abiotic
92 stress. In fact, the identification of novel genes involved in abiotic stress resistance and their
93 application in genetic breeding is now considered an effective approach for the improvement of
94 stress resistance in grapes. The existence of a high-quality de novo-assembled grape genome has
95 made it possible to identify gene families in this species. In this study, we isolated and identified
96 the *XTH* family members from grapevine and performed a complete bioinformatics analysis of the
97 *XTH* family. Interestingly, we identified some putative members with potential functions under
98 abiotic stresses, especially salt and drought stress. These findings allow in-depth research on the
99 potential functions of the selected *VvXTHs* in grapevine.

100

101 **Materials & Methods**

102 **Identification and biochemical analysis of *XTHs* in grapevine**

103 The sequence annotations of the whole genome and the gene GFF3 file were downloaded by using
104 CRIBI v2.1 (<https://urgi.versailles.inra.fr/Species/Vitis/Annotations>) (*Canaguier et al., 2017*).
105 We also downloaded hidden Markov models (PF00722 and PF06958) of the *XTH* domain from
106 the Pfam database (<http://pfam.xfam.org>) and obtained the candidate gene sequence numbers of
107 the grapevine *XTH* family with HMMer software (*Potter et al., 2018*). To avoid duplication and

108 the inclusion of sequences without XTH family domain characteristics, sequences without the
109 XTH domain and sequences showing alternative splicing were removed. The EMBL-EBI online
110 tool (<http://pfam.xfam.org/search/sequence>) (*Gaia et al., 2021*) was used to further analyze
111 secondary structure domains, and the sequences without typical XTH domains were removed.

112 The relative molecular weight (MW), hydrophilicity (GRAVY), and isoelectric point (pI) of
113 these VvXTHs were predicted and analyzed using ExPASy (<https://www.expasy.org/>) (*Duvaud et*
114 *al., 2021*). Single peptide (SP) prediction was performed on the SignalP v4.1 server
115 (<http://www.cbs.dtu.dk/services/SignalP/>).

116 Gene structures were analyzed with Gene Structure Display Server software
117 (<http://gsds.cbi.pku.edu.cn/>) (*Hu et al., 2015*). Conserved motifs in VvXTHs were statistically
118 identified with the online Multiple EM for Motif Elicitation (MEME) software ([https://meme-](https://meme-suite.org/meme/tools/meme)
119 [suite.org/meme/tools/meme](https://meme-suite.org/meme/tools/meme)) (*Bailey et al., 2009*), and TBtools was then used for the clustering
120 and visualization of VvXTHs in grapevine. Multiple protein sequence alignments were performed
121 with ClustalX software and the Esript 3.0 online program
122 (<https://esript.ibcp.fr/ESPrpt/ESPrpt/>) (*Larkin et al., 2007*).

123 **Phylogenetic analysis of VvXTHs in grapevine**

124 To investigate the phylogenetic relationship of VvXTHs, the 34 VvXTH protein sequences from
125 grapevine and the 33 AtXTH protein sequences from *Arabidopsis* were used for multiple sequence
126 alignment by using the Clustal W program within MEGA 11.0 software (*Sudhir et al., 2018*). The
127 phylogenetic tree was built using the neighbor-joining (NJ) method with 1000 bootstrap
128 replications and the p-distance model and was then validated by the maximum likelihood method.
129 To better visualize the phylogenetic tree, the final tree diagram file (*.nwk) was uploaded from

130 MEGA to Figtree and EVOLVIEW online software (<http://www.evolgenius.info/evolview/>)
131 (*Balakrishnan et al., 2019*).

132 The Grape Genome Browser (12X)
133 (<http://www.genoscope.cns.fr/externe/GenomeBrowser/Vitis/>) provided chromosomal location
134 data for all *VvXTHs*. We used TBtools to identify and illustrate the distribution of genes on
135 chromosomes. MCScanX with the default parameters was applied to identify gene duplication
136 events. The CIRCOS program (<https://github.com/CJChen/TBtools>) was used to analyze syntenic
137 relationships among *VvXTHs*. *VvXTHs* falling within the identified collinear blocks were regarded
138 as segmental events, and any two genes separated by a distance of less than 100 kb whose similarity
139 exceeded 75% were considered tandem duplications. To visualize the synteny relationships of
140 orthologous *XTHs* derived from grapes and *Arabidopsis*, Dual Synteny Plotter software
141 (<https://github.com/CJ-Chen/TBtools>) was applied to construct a syntenic analysis map (*Xie et al.,*
142 *2018*). The *Arabidopsis* sequences were obtained from The *Arabidopsis* Information Resource
143 (TAIR) database (<https://www.arabidopsis.org/>) (*Han et al., 2013*). TBtools software was used to
144 calculate the nonsynonymous (Ka) and synonymous (Ks) substitution rates and Ka/Ks ratio of
145 each gene pair. Divergence times were calculated as follows: $T = Ks/2\lambda$ ($\lambda = 6.5 \times 10^{-9}$ for
146 grapevine) (*Li et al., 2019*).

147 **Cis-Element analysis of XTHs in grapevine**

148 The sequences within 1500 base pairs (bp) upstream of the starting codon of the *VvXTHs* were
149 obtained from Ensembl Plants (<http://plants.ensembl.org/index.html>) as the promoter regions (*Dan*
150 *et al., 2017*). The *cis*-elements were predicted with PlantCARE Web Tools
151 (<http://bioinformatics.psb.ugent.be/webtools/plantcare/html/>) (*Magali et al., 2002*) and New

152 PLACE Web Tools (<https://www.dna.affrc.go.jp/PLACE/?action=newplace>). TBtools was used
153 to draw heatmaps and build clustering trees.

154 **Gene expression analysis of *XTHs* in different grapevine organs and tissues**

155 To understand the spatial and temporal expression patterns of *VvXTHs* during development, a high-
156 throughput microarray data, from a gene expression atlas generated from different organs/tissues
157 at different developmental stages (*Marianna et al., 2012*), was employed for further analysis.
158 According to the gene ID, the expression profiles of *VvXTHs* was extracted from the GSE36128
159 data set, and we then normalized the average expression value of each gene in 54 samples
160 (including green and woody tissues and organs at different developmental stages as well as
161 specialized tissues such as pollen and senescent leaves). TBtools was used to draw heatmaps and
162 build clustering trees.

163 To verify the reliability of the results obtained from the GSE36128 data set, the organ-specific
164 expression patterns were examined with quantitative real-time RCR (qRT-PCR) using the five
165 different organs (tendrils, root, stem, leaf and flower) from 5-year-old trees of grapevine “Crimson”
166 growing at the experiment station of Shandong Agricultural University (Tai'an, Shandong, China).

167 **RNA extraction and expression analysis of *VvXTHs***

168 The tissue culture seedlings of *Vitis vinifera* cv “Crimson” seedless were grown on 1/2 Murashige
169 and Skoog (MS) solid medium with 0.2 mM indole-3-butyric acid (IBA) under a 16-h-light/8-h
170 dark cycle at $24 \pm 1^\circ\text{C}$ for six weeks. Six-week-old seedlings, which transcription level changes
171 more pronounced, were transferred to liquid medium containing 200 mM NaCl or 200 mM
172 mannitol for salt and drought stress treatments, respectively. The treated seedlings were extracted
173 and separated into leaves and roots for 0, 3, 6, 9, 12, and 24 h upon treatment, immediately frozen

174 in liquid nitrogen and stored at -80°C for RNA extraction. For each sample, three biological
175 replicates were collected.

176 Total RNA was extracted from the samples treated with NaCl and mannitol using a HiPure
177 HP Plant RNA Mini Kit (Magen, Guangzhou, China) based on the supplier's instructions.
178 Subsequently, first-strand cDNA was synthesized from total RNA with the PrimeScript™ RT
179 reagent kit with gDNA Eraser (Vazyme Biotech Co., Nanjing, China). qRT-PCR was performed
180 using a SYBR® PrimeScript™ RT-PCR Kit (TaKaRa, Dalian, China) according to the supplier's
181 instructions with a CFX96™ Real-Time PCR Detection System. Gene expression levels were
182 normalized against the average expression of the internal reference gene, and the baseline and Ct
183 (threshold cycles) value were automatically determined by the CFX Manager software program.
184 The relative expression levels of *VvXTHs* were calculated using the $2^{-\Delta\Delta\text{Ct}}$ comparative Ct method.
185 The internal reference gene used in this study was *Vvβ-actin7* (XM_034827164), which has been
186 proved to be a most stable gene for normalization by comparison with other reference gene (*Vvβ-*
187 *actin101*: XM_002265440) (Fig. S1). All experiments were performed with three biological
188 replicates, and all the primers used in this study are listed in Table S1. To visualizing the relative
189 difference, the expression level of 0 h treatments for salt and drought stresses and tendrils for plant
190 tissue specificity was set as 1, respectively. Then, TBtools was used to draw a heatmap for
191 visualization.

192 **Results**

193 **Identification and analysis of *VvXTHs* in grapevine**

194 Forty-two sequences were identified by searching for two domains (Pfam: PF00722 and PF06955)
195 with the HMMer program. We deleted six alternative splicing sequences and two sequences
196 without typical XTH domains. As a result, we finally identified 34 *VvXTHs*. Described by previous

197 studies (Cao *et al.* 2016; Fu *et al.*, 2019; Li *et al.*, 2018; Wan *et al.*, 2014), we named these genes
198 according to their chromosomal locations and named them *VvXTH1-VvXTH34*.

199 The analysis of the physical and biochemical data of the 34 *VvXTHs*, including their amino
200 acids (AAs), MWs, SPs, pIs, total average hydrophilicity (GRAVY) and subcellular localization,
201 revealed that they contained 251~369 AAs. The MW ranged from 28.5 to 41.7 kDa, while the pI
202 ranged from 4.61 to 9.45. All XTHs exhibited hydrophilicity. Subcellular location prediction
203 results showed that most of the genes are localized in the plasma membrane (29), while a few were
204 localized extracellularly (5), including *VvXTH10* and *VvXTH12* in group IIIA and *VvXTH2*,
205 *VvXTH32*, and *VvXTH33*. The majority of the proteins (80%) contained signal peptides, which
206 were approximately 25-AA long (Table 1).

207 **Phylogenetic analysis and classification of *VvXTHs***

208 To investigate the evolutionary relationships and functional associations of *VvXTHs* with
209 *AtXTHs*, we built a phylogenetic tree utilizing the protein sequences of *XTHs* from *Vitis vinifera*
210 and *Arabidopsis* (Fig. 1). The *VvXTHs* were grouped according to the previous grouping method
211 applied for *AtXTHs* and the evolutionary relationship between grapes and *Arabidopsis*. The results
212 of the phylogenetic analysis indicated that the 34 *VvXTHs* could be divided into three groups,
213 including twenty-seven *VvXTHs* in group I/II, two in group IIIA, and five in group IIIB. In
214 addition, one XTH protein (*VvXTH11*) was classified into the original ancestral group. Group I/II
215 contained most of the members, and substantial similarity could be observed between some
216 members of the group. The termini of the phylogenetic tree branch showed a total of twenty-two
217 sister pairs, eight of which were orthologous pairs between *Arabidopsis* and grapevine, and six
218 were grape homolog gene pairs. This analysis revealed that the number of *VvXTHs* was slightly
219 expanded in comparison to the number of *XTHs* in *Arabidopsis*.

220 Thirty-four *VvXTHs* were unevenly distributed on 13 chromosomes. In particular, Chr.11
221 contained the largest number of *VvXTHs* (15), whereas other chromosomes contained considerably
222 fewer genes. For example, a total of 4, 3, and 2 genes were located on Chr.5, Chr.10 and Chr.1,
223 respectively. In addition, Chr.2, Chr.3, Chr.6, Chr.7, Chr.8, Chr.12, Chr.15, Chr.16, and Chr.17
224 each contained only 1 gene (Fig. 2A). Therefore, it can be inferred that there should be no
225 observable association between the number of *XTHs* and the length of chromosomes. Furthermore,
226 the genes located on Chr.11 and Chr.5 were closely clustered together. According to the
227 chromosome location and genome annotation information, a total of 81 tandem duplicate gene
228 pairs were obtained (Fig. 2A). To determine the relationships among *VvXTH* members, we
229 performed a collinearity analysis and found no *VvXTH* within the identified collinear blocks, which
230 indicated that segmental duplication is not involved in *VvXTH* expansion (Fig. 2B). These results
231 prove that the expansion of *VvXTHs*, especially group I/II gene members, was driven by tandem
232 duplication. We also traced the duplication time of *VvXTHs* by analyzing their Ka, Ks and Ka/Ks
233 ratio. The Ka/Ks ratios of all *VvXTHs* were less than 1, ranging from 0.07 to 0.28. The duplication
234 times of all *VvXTHs* were also calculated. The duplication times ranged from 2.91-78.39 Mya
235 (million years ago) (Table S2). To further evaluate the evolution and development of the *VvXTH*
236 family, we constructed a comparison diagram of grapes and *Arabidopsis*. Eight *VvXTHs* were
237 shown to be synonymous with *XTHs* of *Arabidopsis*. Among these genes, *VvXTH10* is collinear
238 with *AtXTH31* and *AtXTH32* in *Arabidopsis*, while *VvXTH1* is collinear with *AtXTH27* and
239 *AtXTH28* in *Arabidopsis* (Fig. 2C). It is speculated that there may be functional redundancy among
240 these genes, which implies that they have important roles in evolutionary progress.

241 **Gene structural and multiple sequence alignment analysis of *VvXTHs***

242 Gene structural analysis revealed that the closely related genes within this subfamily are
243 characterized by a similar structure, which can be further verified by the results of phylogenetic
244 analysis (Fig. 3A). With the exception of *VvXTH32*, which lacks any introns, all other *VvXTH*
245 members contain 2~4 different introns. In particular, group I/II contains a large number of
246 members, and most members have 2 introns. The gene sister pairs, including *VvXTH23/26*,
247 *VvXTH25/28*, *VvXTH14/15*, and *VvXTH8/9* at the terminal branch of the evolutionary tree, have
248 highly similar exon/intron structures. In addition, compared with the adjacent gene *VvXTH27*,
249 *VvXTH24* has lost an exon and exhibits different intron and exon lengths. The members of group
250 IIIA all have 3 introns and 4 exons and present high structural similarity. The members of group
251 IIIB have developed different numbers and lengths of intron/exon structures during the long
252 evolutionary process (Fig. 3C). In general, most *VvXTHs* present the same intron/exon structure
253 pattern and remain conserved during evolution, which is consistent with the results obtained in
254 other plants.

255 Based on the results of MEME motif analysis, motif 3 and motif 4 are highly conserved in all
256 *VvXTHs*. Motif 3 is a characteristic domain that catalyzes enzymatic contact reactions, which
257 denoted as (D/N)E(I/L/A/V/F)(D/T)(F/I)E(F/I/L)LG (Fig. 3B and 3D). Among these, the first
258 glutamate residue (E) indicates the catalytic nucleophile that initiates the enzymatic reaction, and
259 the second E residue functions represents a base to activate the entrant substrate. In addition, all
260 members except for *VvXTH30* contain motif 1; moreover, members of the same group share a
261 similar motif composition (Fig. 3B). For instance, motif 2 only exists in group I/II, and motif 8
262 only exists in group IIIA and IIIB. Genes in the same clade, especially those that are closely related,
263 such as 1) *VvXTH23*, *VvXTH26*, *VvXTH25*, and *VvXTH28*; 2) *VvXTH6*, *VvXTH7*, *VvXTH8*, and
264 *VvXTH9*; and 3) *VvXTH10* and *VvXTH12*, can share much more similar motif structures (Fig. 3B).

265 In addition, motif 7, motif 9, and motif 10 only exist in group I/II, and most of the group members
266 contain the motifs mentioned above (Fig. 3B). The members of group IIIA contain 5 motifs with
267 the same distribution. Group IIIB members contain 4-6 motifs; while they all share 4 identical
268 motifs, only VvXTH4 and VvXTH32 have motif 6, and only VvXTH33 does not have motif 5.
269 The results of multiple sequence alignment also confirmed that the conserved domain active site
270 is present in all VvXTHs. Moreover, with the exception of VvXTH2 (IIIB), VvXTH10 (IIIA), and
271 VvXTH12 (IIIA), potential N-glycosylation residues are located near the active site in the 31 other
272 VvXTHs (Fig. 3E). Conserved domain predictions suggested that members of the same subfamily
273 may have similar structures and may be involved in similar functions. Thus, we need to focus on
274 distinctive members that may present surprising functions that remain to be discovered.

275 **Organ-specific expression pattern analysis of VvXTHs**

276 Through the expression profile (GSE36128) analysis of the GEO data set, we obtained the specific
277 expression patterns of *VvXTH* members in different organs and developmental periods of
278 grapevine to predict the functions of *VvXTHs* in growth and development (Fig. 4). According to
279 the results of cluster analysis, the *VvXTH* families were classified into 4 groups (A-D): group A
280 contains seven genes with high expression levels in berry peels, skins, shafts, and tendrils; group
281 B includes four members with high expression levels only in stems and tendrils but low expression
282 in other organs; group C includes eight genes with very low expression levels in all organs; and
283 group D is the largest (15 members) subfamily and shows the highest expression in berries, shafts,
284 and tendrils. In addition, the *VvXTHs* had higher expression levels in the pulp, peel, and stem
285 during the V (veraison), MR (mid-ripening), and R (ripening) periods, indicating that *VvXTHs* may
286 be related to fruit ripening. In short, the four groups of *VvXTHs* present specific expression profiles

287 depending on the organ and developmental stage. This interesting phenomenon may be due to the
288 specific functions of these specific genes in related tissues.

289 To verify the reliability of the organ-specific expression profiles, qRT-PCR analysis was
290 conducted on five different tissues (tendrils, root, stem, leaf and flower) of grapevine “Crimson”
291 for *VvXTHs*, then the qRT-PCR results were compared with the data obtained from GSE36128
292 data set of the cultivar “Corvina” (*Marianna et al., 2012*) with the same tissues at the
293 corresponding developmental stages. It was found that the expression patterns of *VvXTHs* were
294 generally consistent with the data obtained from the GSE36128 data set (Fig. S2), which suggests
295 that temporal and spatial expression of *VvXTHs* is generally similar in different cultivars, even
296 grown in different conditions.

297 **Transcriptional profiles of *VvXTHs* under abiotic stress**

298 The PlantCARE database and New PLACE database were utilized to identify *cis*-elements in the
299 DNA sequences 1.5 kb upstream of the *VvXTH* start codons. The results showed that all 34 *VvXTHs*
300 contained a variety of abiotic and biotic stress response elements, phytohormone response
301 elements, and growth and development-related response elements (Fig. 5A). Similarly, in the New
302 PLACE database, all 34 *VvXTHs* were predicted to contain phytohormone response elements and
303 elements involved in the responses to a variety of abiotic stresses, including cold and heat, ABA,
304 dehydration and salinity (osmotic) stress (Fig. 5B). As shown in Fig. S3, two drought stress
305 response elements (MYB and MYC) exist in almost all members, and 80% of the gene members
306 contain defense and stress response elements (STREs), which indicates that the *VvXTH* family
307 probably shows important functions when plants are subjected to abiotic or biotic stress. Abscisic
308 acid response elements (ABREs) and salicylic acid response elements (TCA elements) are
309 abundantly present in *VvXTH* family members, which indicates that the *VvXTH* family may also

310 be involved in hormone regulation during plant growth and the response to stress. The presence of
311 a meristem development control element (CAT-box) in most *VvXTH* members suggests that the
312 *VvXTH* family may have significant effects on the regulation of plant growth and development. In
313 particular, the 34 *VvXTHs* were predicted to contain a large number of light response elements in
314 the New PLACE database, and light-responsive elements (GATABOX and Box-4) were present
315 in most *VvXTH* members, suggesting that the *VvXTH* family plays important roles in
316 photosynthesis and photomorphogenesis.

317 Promoter analysis demonstrated the widespread presence of *cis*-elements associated with
318 abiotic stress in the promoter regions of *VvXTHs*, revealing the possible induction of *VvXTH*
319 expression by abiotic stress. To further investigate the potential roles of *VvXTHs* in response to
320 abiotic stress, especially drought stress and salt stress, we selected 14 *VvXTH* members harboring
321 abiotic response *cis*-elements for further study. Six-week-old grape seedlings were exposed to 200
322 mM NaCl or 200 mM mannitol, and the expression of 14 *VvXTHs* was examined in separated
323 leaves and roots. In roots, the expression levels of 11 genes were upregulated under salt stress,
324 among which 4 members were significantly upregulated. The expression of *VvXTH5*, *VvXTH20*,
325 and *VvXTH34* was increased by more than two-fold, and that of *VvXTH4* was increased by more
326 than four-fold. Interestingly, the expression of *VvXTH4* peaked after 9 h of treatment, presenting
327 an obviously different pattern from the other genes. This shows that these genes may respond to
328 salt stress in different ways. Under drought stress, most of the genes whose expression was
329 upregulated reached a peak after 3 h of treatment, and some genes were upregulated by more than
330 four-fold (*VvXTH3* and *VvXTH20*). *VvXTH10* expression reached a peak after 12 h of treatment,
331 indicating that this gene may be expressed at a later time. The number of upregulated *VvXTHs* in
332 leaves relative to roots decreased after stress treatment, but these genes were more highly

333 upregulated. Among these genes, *VvXTH3*, *VvXTH10*, and *VvXTH31* were upregulated by
334 approximately ten-fold. The above genes might play particularly crucial roles in the leaf response
335 to salinity stress. Taken together, our findings indicate that the expression of *VvXTHs* could be
336 altered by salt and drought stress, suggesting that *VvXTHs* may participate in reactions to abiotic
337 damage, especially under salt and drought stress.

338

339 Discussion

340 The *XTH* family consists of modification enzymes that can rebuild cell walls by modulating the
341 construction and composition of xyloglucan cross-links (Campbell *et al.*, 2010). According to
342 previous studies, various members of this family have been identified in *Arabidopsis thaliana*,
343 *Oryza sativa*, *Medicago truncatula*, *Nicotiana tabacum*, *Solanum lycopersicum*, and *Ananas*
344 *comosus*, and these proteins have been verified to play critical roles in development, biotic stress,
345 and abiotic stress (Meng *et al.*, 2018; Li *et al.*, 2019; Yokoyama, 2004; Kurasawa *et al.*, 2009;
346 Xuan *et al.*, 2016). The release of the most recent grape genome database made it possible to
347 identify the grape *XTH* family (Ariga *et al.*, 2007). In this study, thirty-four *VvXTHs* were
348 systematically identified and characterized using bioinformatics approaches. The results showed
349 that the number of identified *VvXTHs* (34) (Fig. 1) was slightly greater than the numbers found in
350 *Arabidopsis thaliana* (33) and *Oryza sativa* (29) (Yokoyama, 2004; Kurasawa *et al.*, 2009), which
351 may be related to pedigree-specific gains and losses as well as gene duplication events. Gene
352 duplication is a primary driver of the expansion of gene families, and tandem duplications and
353 segmental duplications are considered the primary duplication modes (Zhu *et al.*, 2014). Previous
354 studies of the *XTH* family have also reported gene tandem duplications or segmental duplications
355 in barley, soybean, and tobacco (Fu *et al.*, 2019; Li *et al.*, 2018; Meng *et al.*, 2018).

356 Interestingly, we observed that the thirty-four identified *VvXTHs* were located on 13
357 chromosomes, and Chr.11 and Chr.5 contained gene clusters (Fig. 2). Based on the definition of
358 gene tandem duplication, *VvXTH17-VvXTH30* and *VvXTH6-VvXTH9* represent gene tandem
359 duplication events. According to the analysis of the Ka/Ks ratio (Table S2), all genes showed ratios
360 of less than 1, which indicates that they are under intense purifying selection (*Hurst, 2002*). Hence,
361 the role of gene tandem duplication in *VvXTH* family expansion, particularly in increasing the
362 number of *VvXTH* members and their functional diversification, is irreplaceable.

363 According to gene function and Clustal analyses, similar to other plants, the thirty-four
364 *VvXTHs* are divided into groups I/II, IIIA, and IIIB and an ancestral group (Fig. 1). According to
365 previous studies, due to the unclear distinction between groups I and II, these subgroups were
366 combined into one group (group I/II) (*Campbell et al., 1999*). The XTHs in group IIIA mainly
367 display XEH activity, while those of group IIIB showed obvious XET activity, suggesting a
368 functional distinction between groups IIIA and IIIB (*Eklof and Brumer 2010; Nomchit et al., 2010;*
369 *Opazo et al., 2017*). Serines or threonines located near the catalytic center of XET are typical
370 residues for N-glycosylation, and the results of multiple sequence alignment showed that the
371 members of groups I/II and IIIB (except for *VvXTH2*) contain N-glycosylated residues, while
372 those of group IIIA do not (*Mark et al., 2009*). Therefore, we speculated that *VvXTH10* and
373 *VvXTH12* proteins in group IIIA might possess XEH activity and that *VvXTH4*, *VvXTH32* and
374 *VvXTH33* proteins in group IIIB might show XET activity in grape, which is in agreement with
375 previous research findings (Fig. 1) (*Mark et al., 2009; Miedes and Lorences, 2009*).

376 The analysis of gene structure is of great significance to further clarify the origins, evolution,
377 and genetic relationships of species. *XTH* family members show a relatively wide variety of
378 structures. Specifically, most members of the grape *XTH* family contain 3 or 4 introns (Fig. 2C),

379 while others have fewer intron, which may be related to gene splicing (*Mount et al., 2012*). It is
380 precisely because of the existence of multiple introns that gene splicing becomes more
381 complicated, and the number of different *XTH* expression products increases. According to a
382 comparison of the AA sequences of *Arabidopsis thaliana*, *Populus tomentosa*, *Hordeum vulgare*,
383 *Brassica rapa*, and *Brassica oleracea*, even when the difference in protein size is obvious, the
384 active-site domain is still conserved in the reported XTH proteins (*An et al., 2017*). In this study,
385 all 34 VvXTHs were found to contain motif 3 (*Opazo et al., 2017*), which suggests that XTH
386 proteins may play similar roles in the plant kingdom. It has been reported that the active site
387 mediates catalytic activity, which can catalyze hydrolase activity and carry out cell wall structural
388 modification and rearrangement by cutting and repolymerizing cellulose single chains (*Li et al.,*
389 *2018; Behar et al., 2018*). The cross-linked xyloglucan structure has critical functions in
390 maturation and resistance to abiotic stress (*Bulone et al., 2019*).

391 Previous studies have shown that *XTHs* are of vital importance in plant resistance to abiotic
392 stress (*Chen et al., 2019; Dong et al., 2019; Li et al., 2019*). The expression of *CaXTH3* is induced
393 by a variety of abiotic stresses, such as drought, high salt, and low temperature, and the tolerance
394 of *CaXTH3* transgenic tomato plants to salt and drought stress is thereby significantly improved
395 (*Choi et al., 2011*). Additionally, the heterologous expression of *PeXTH* in tobacco improves plant
396 osmotic tolerance by reducing water loss and reducing the speed of stomatal opening (*Han et al.,*
397 *2014*). To study the potential function of *VvXTHs* against abiotic stress, we carried out promoter
398 analysis and tissue expression analysis (Fig. 3 and 4). The results indicated that the upstream
399 promoter regions of almost all members of the grape *XTH* family contain MBS, MYB, MYC, and
400 ARE *cis*-elements for responding to drought stress. Furthermore, 47% of these sequences contain
401 ABREs, to respond to ABA (Fig. 4). Under drought conditions, ABA inhibits root growth and

402 development, represses seed germination, and promotes the shedding of senescent tissues and
403 organs (*Hirayama and Shinozaki, 2007*). Some *VvXTHs* exist in the mature stage, and ABA may
404 promote *XTH* expression and affect organ abscission (Figs. 3 and 6). In addition, a few members
405 of the *VvXTH* family also contain DRE action elements, which means that *XTHs* can potentially
406 respond to salt stress, in addition to drought stress (Fig. 5). It is also interesting that the expression
407 of *VvXTHs* varies in different organs. Genes from the same gene cluster of gene tandem repeat
408 events, such as *VvXTH6* and *VvXTH7* or *VvXTH26*, and *VvXTH30*, show differences in expression
409 in organs at different stages. These results implied that during evolution, closely related genes have
410 undergone subfunctional evolution, functionalization or nonfunctionalization, helping grapes
411 adapt to a variety of growth environments. The expression profiling of *VvXTHs* under different
412 stresses revealed the induction of *VvXTH3*, *VvXTH31*, and *VvXTH10* in roots.

413 Taken together, these findings provide novel information about *VvXTHs* under abiotic stress,
414 especially drought and salt stresses. It can be speculated that the above genes may show increased
415 cell wall-related functions under stress by combining with xyloglucan. Nevertheless, further
416 molecular and genetic identification efforts are needed to verify their functions.

417

418 **Conclusions**

419 In this study, thirty-four *XTHs*, which could be further divided into group I/II, group IIIA, and
420 group IIIB, were successfully isolated and identified in grapevine. It was shown that the *VvXTHs*
421 are unevenly distributed on 13 chromosomes. According to collinearity analysis, tandem
422 duplication of genes may have occurred on Chr.5 and Chr.11. Furthermore, all *VvXTHs* contain
423 conserved *XTH* domains and active sites. Expression analysis showed that some *VvXTHs* can
424 effectively respond to salt and drought stress at the transcriptional level. In this context, the results

425 of the present investigation will lay a foundation for future investigations of the function of
426 *VvXTHs*.

427

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432

433 **Competing Interests**

434 The authors declare that they have no competing interests.

435

436 **Author Contributions**

437 Tian Qiao and Lei Zhang conceived and designed the experiments, performed the experiments,
438 analyzed the data, prepared figures and tables, authored or reviewed drafts of the paper. Yanyan
439 Yu and Yunning Pang performed the experiments, prepared figures and tables, reviewed drafts of
440 the paper. Xinjie Tang, Xiao Wang and Lijian Li analyzed the data, prepared figures and tables,
441 reviewed drafts of the paper. Qinghua Sun and Bo Li conceived and designed the experiments,
442 revised the work critically for important content, reviewed drafts of the paper. All authors read
443 and approved the final manuscript.

444

445 **Data availability**

446 All relevant data and plant materials that support the findings of this study are available from raw
447 data in supplemental files.

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648 Table

649 **Table 1 Molecular characteristics of *VvXTHs* in grapevine.** AA: amino acid; MW: molecular
650 weight; SP: signal peptide; pI: isoelectric point; GRAVY: total average hydrophilicity.

651 Figure Legends**652 Fig. 1 Phylogenetic analysis of XTHs of *Arabidopsis* and grapevine.**

653 The amino acid-based phylogenetic tree was generated using MEGA11.0 software via the
654 neighbor-joining method. Bootstrap test results are indicated in the tree. The different colored
655 branches and arcs represent Group I/II, IIIA, IIIB, and the Ancestral Group, and the blue
656 five-pointed star represents *AtXTH* family members. The red triangle represents *VvXTH* family
657 members.

658 Fig. 2 Systematic analysis of *VvXTHs* in grapevine.

659 (A) Thirty-four *VvXTHs* were mapped on grape chromosomes based on their physical positions.
660 Eighty-one tandemly duplicated gene pairs are indicated by red lines. The scale on the left is in
661 megabases (Mb). (B) Schematic representations of the chromosomal distribution and
662 interchromosomal relationships of *VvXTHs*. Gray lines indicate all synteny blocks in the grape
663 genome. Gene IDs on the chromosomes indicate gene physical positions. (C) Gray lines in the
664 background indicate the collinear blocks identified in grape and *Arabidopsis*, while the different
665 colored lines highlight the syntenic *XTH* gene pairs.

666 Fig. 3 Phylogenetic relationships, structures and conserved motifs of *VvXTHs*.

667 (A) Phylogenetic tree inferred from the protein sequences of *VvXTHs*. Branch colors represent
668 different groups. (B) The motif composition of the *VvXTHs* identified using MEME. The different
669 colored boxes represent different motifs and their positions in each *VvXTH* sequence. Each motif
670 is indicated by a colored box in the legend at the bottom. (C) Gene structure of *VvXTHs*. The boxes

671 represent exons or untranslated regions (UTRs), and lines represent introns. (D) Schematic
672 representation of the conserved domains found in grape. (E) Multiple sequence alignments of the
673 conserved domains of the VvXTHs. The black lines indicate the conserved domains. N-
674 glycosylation residues are indicated with asterisks.

675 **Fig. 4 Expression patterns of *VvXTHs* in different organs and developmental stages.**

676 Rows represent *VvXTH* members, while columns represent different developmental stages and
677 organs. The expression levels of *VvXTHs* are indicated by the intensity of color. The phylogenetic
678 tree on the left side of the heatmap is based on the hierarchical clustering of the expression profiles
679 of *VvXTHs* in 54 samples.

680 **Fig. 5 *Cis*-element analysis in the promoter regions 1500 bp upstream of the start codons of**
681 ***VvXTHs*.**

682 The prediction analysis was performed by using plantCARE (A) and New PLACE (B). The bar
683 graphs represent the total number of *cis*-elements in each gene promoter region. Different colors
684 represent different types of *cis*-elements. Three types of *cis*-elements were predicted in
685 plantCARE. Five types of *cis*-elements were predicted in New PLACE.

686 **Fig. 6 Expression profiles of *VvXTHs* under abiotic stress.**

687 Heatmap showing the relative expression of 14 *VvXTHs*, detected by qRT-PCR, in roots and leaves
688 of 6-week-old “Crimson” grape seedlings after treatment with 200 mM NaCl and 200 mM
689 mannitol for 0, 3, 6, 9, 12, and 24 h (0 h treatment as the control). Experiments were performed in
690 biological triplicates.

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694 **Supplementary materials**

695 **Table S1. Primers used for qRT-PCR in this study.**

696 **Table S2. Ka/Ks analysis and duplication date estimated for duplicating *VvXTHs* paralogs.**

697 **Fig. S1. The relative expression of *VvXTH4* (A) and *VvXTH20* (B) in grapevine roots under**
698 **salt stress normalize using *Vvβ-actin7* and *Vvβ-actin101*, respectively.**

699 **Fig. S2. The comparison between quantitative qRT-PCR data and Microarray data.**

700 **Fig. S3. Heatmap of *cis*-element analysis**

Figure 1

Phylogenetic analysis of XTHs of *Arabidopsis* and grapevine.

The amino acid-based phylogenetic tree was generated using MEGA11.0 software via the neighbor-joining method. Bootstrap test results are indicated in the tree. The different colored branches and arcs represent Group I/II, IIIA, IIIB, and the Ancestral Group, and the blue five-pointed star represents *AtXTH* family members. The red triangle represents *VvXTH* family members.

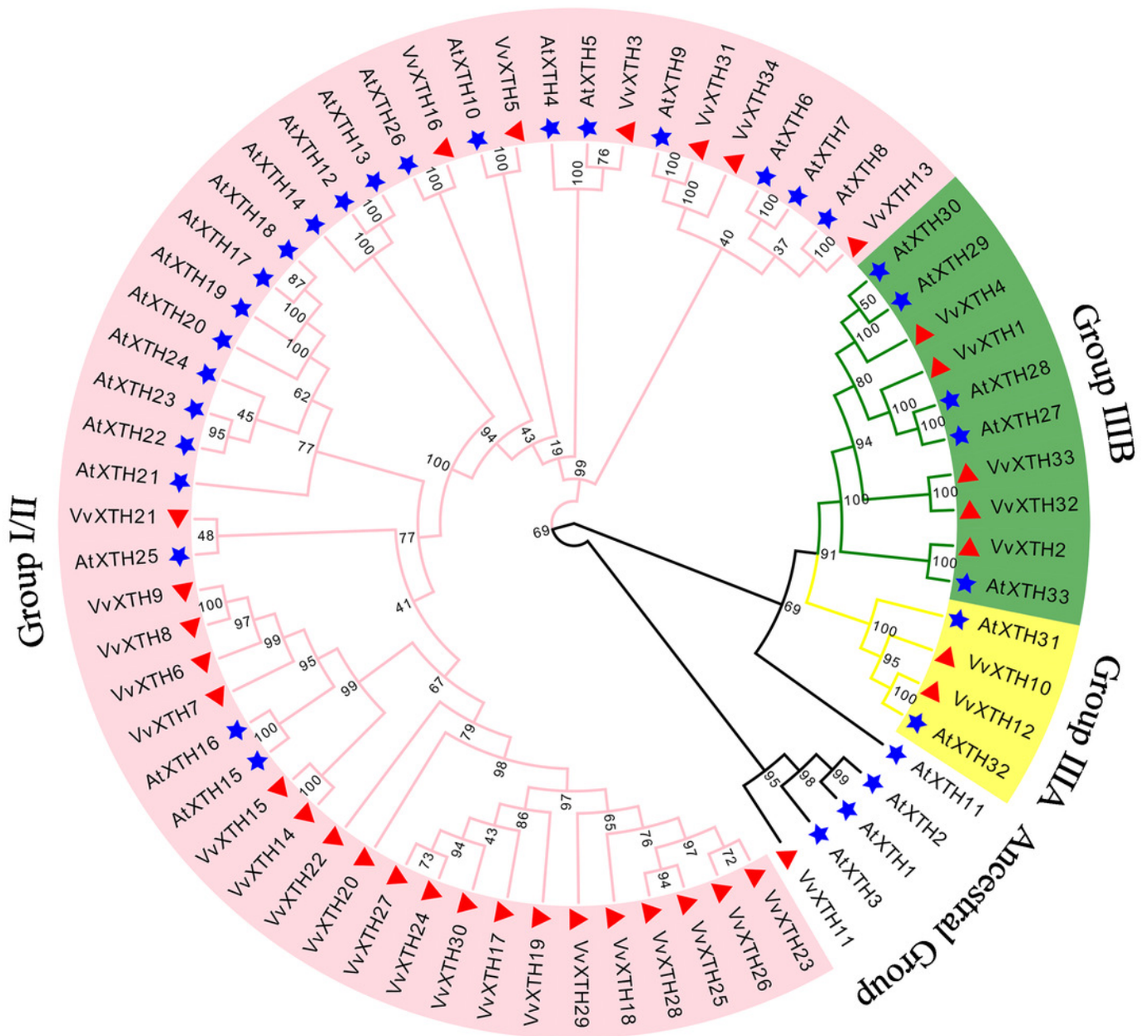


Figure 2

Systematic analysis of *VvXTHs* in grapevine.

(A) Thirty-four *VvXTHs* were mapped on grape chromosomes based on their physical positions. Eighty-one tandemly duplicated gene pairs are indicated by red lines. The scale on the left is in megabases (Mb). (B) Schematic representations of the chromosomal distribution and interchromosomal relationships of *VvXTHs*. Gray lines indicate all synteny blocks in the grape genome. Gene IDs on the chromosomes indicate gene physical positions. (C) Gray lines in the background indicate the collinear blocks identified in grape and *Arabidopsis*, while the different colored lines highlight the syntenic *XTH* gene pairs.

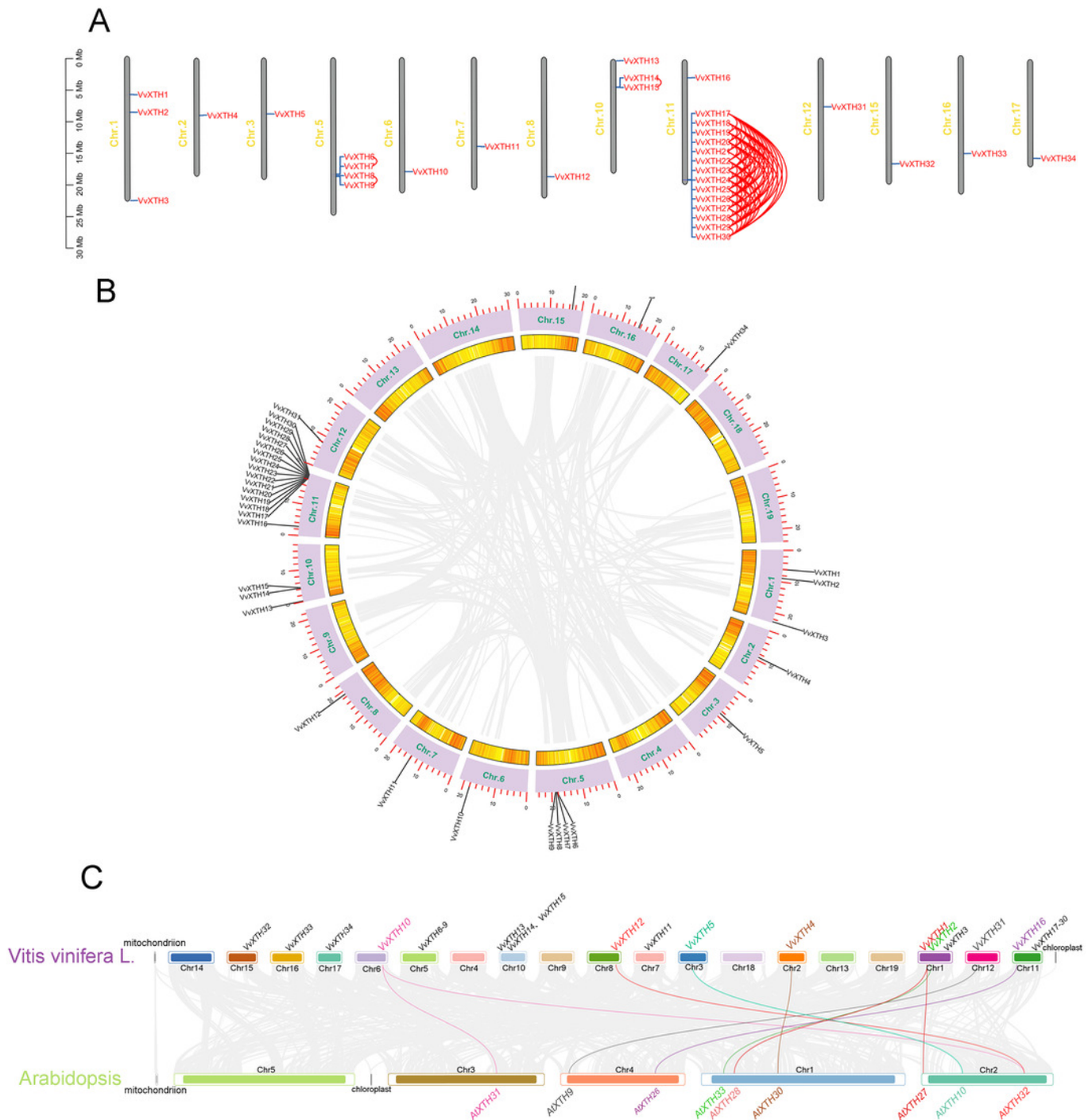


Figure 3

Phylogenetic relationships, structures and conserved motifs of VvXTHs.

(A) Phylogenetic tree inferred from the protein sequences of *VvXTHs*. Branch colors represent different groups. (B) The motif composition of the *VvXTHs* identified using MEME. The different colored boxes represent different motifs and their positions in each *VvXTH* sequence. Each motif is indicated by a colored box in the legend at the bottom. (C) Gene structure of *VvXTHs*. The boxes represent exons or untranslated regions (UTRs), and lines represent introns. (D) Schematic representation of the conserved domains found in grape. (E) Multiple sequence alignments of the conserved domains of the *VvXTHs*. The black lines indicate the conserved domains. N-glycosylation residues are indicated with asterisks.

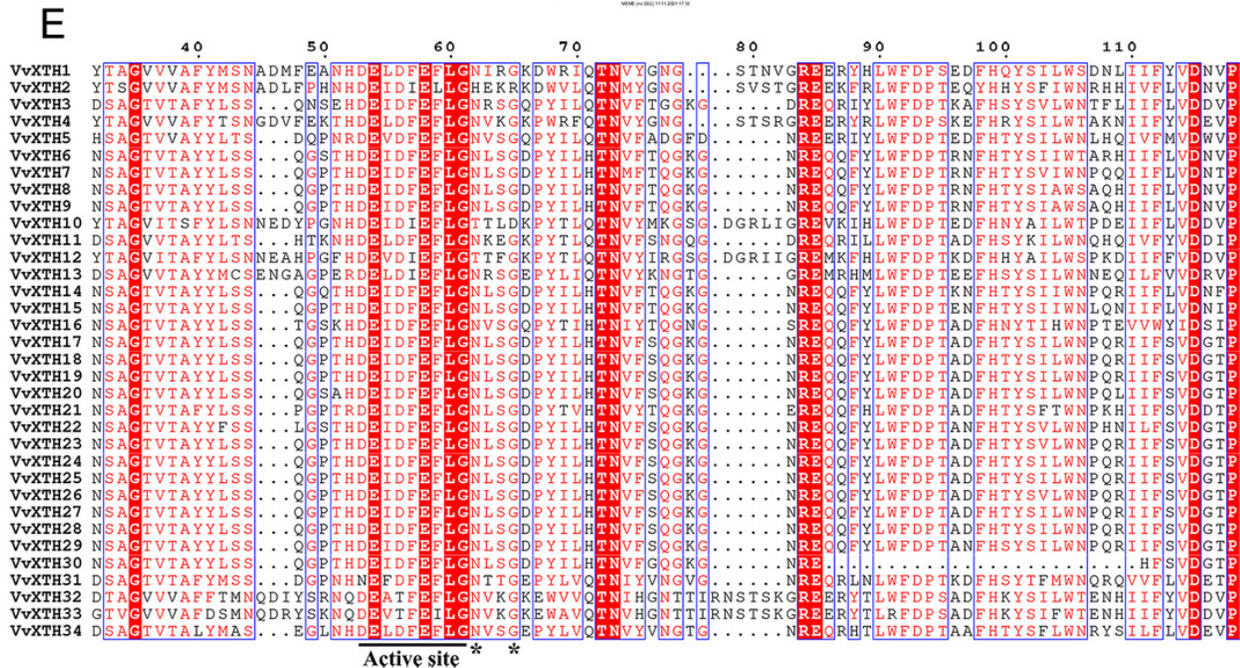
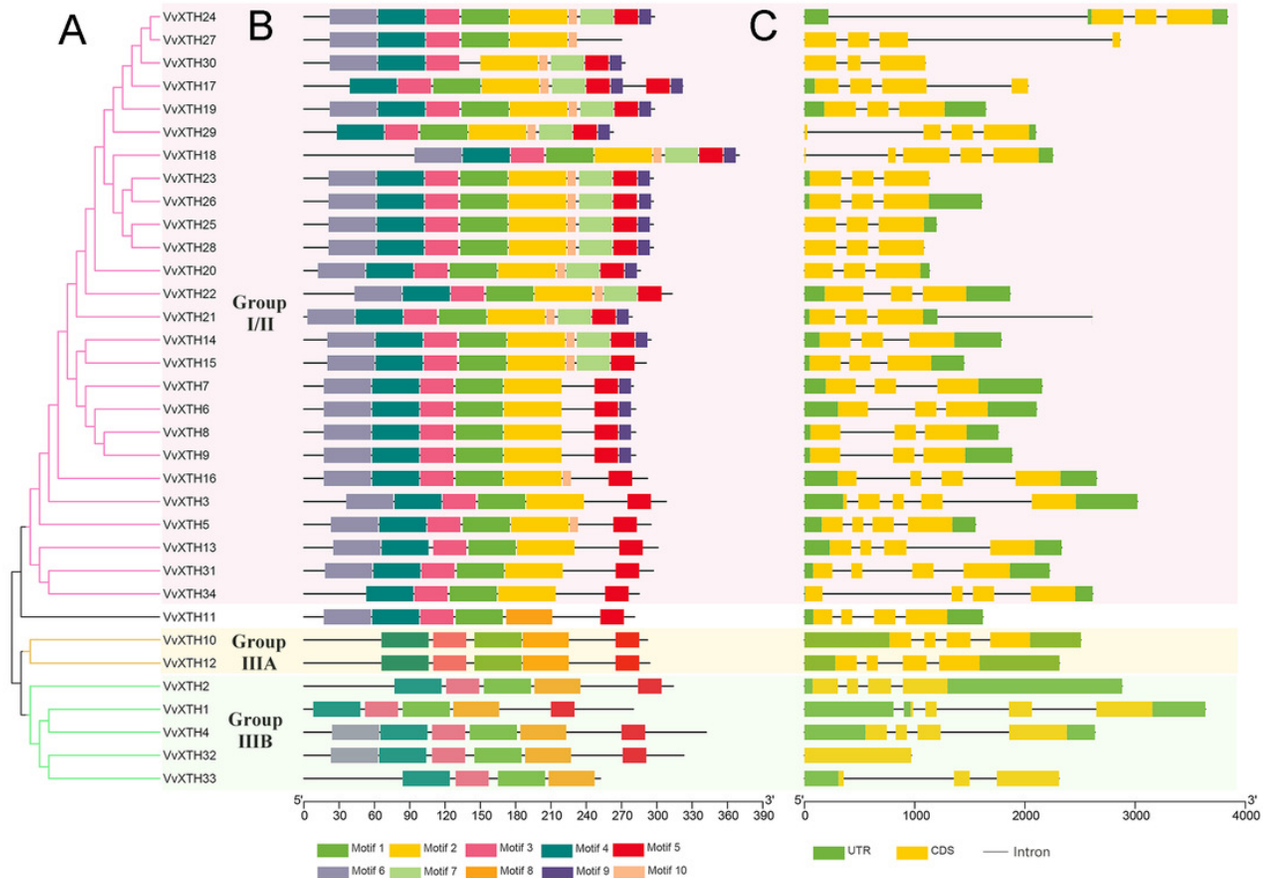


Figure 4

Expression patterns of *VvXTHs* in different organs and developmental stages.

Rows represent *VvXTH* members, while columns represent different developmental stages and organs. The expression levels of *VvXTHs* are indicated by the intensity of color. The phylogenetic tree on the left side of the heatmap is based on the hierarchical clustering of the expression profiles of *VvXTHs* in 54 samples.

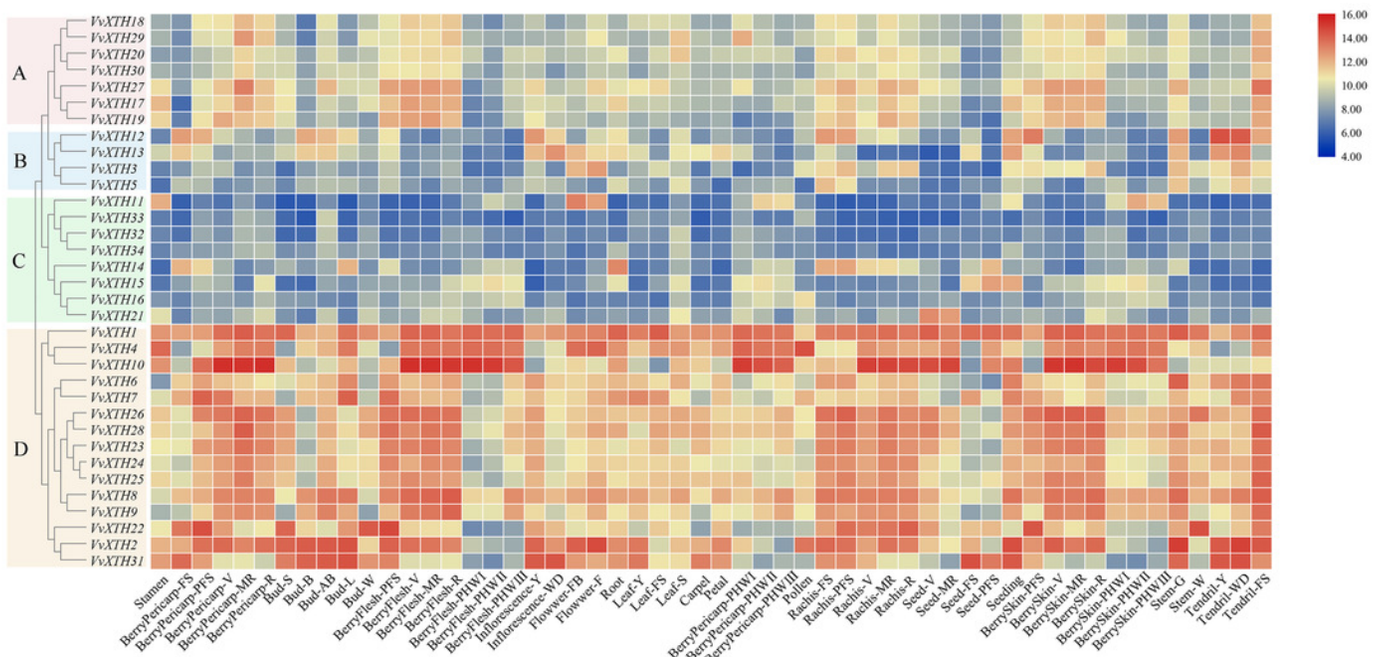


Figure 5

Cis-element analysis in the promoter regions 1500 bp upstream of the start codons of *VvXTHs*.

The prediction analysis was performed by using plantCARE (A) and New PLACE (B). The bar graphs represent the total number of *cis*-elements in each gene promoter region. Different colors represent different types of *cis*-elements. Three types of *cis*-elements were predicted in plantCARE. Five types of *cis*-elements were predicted in New PLACE.

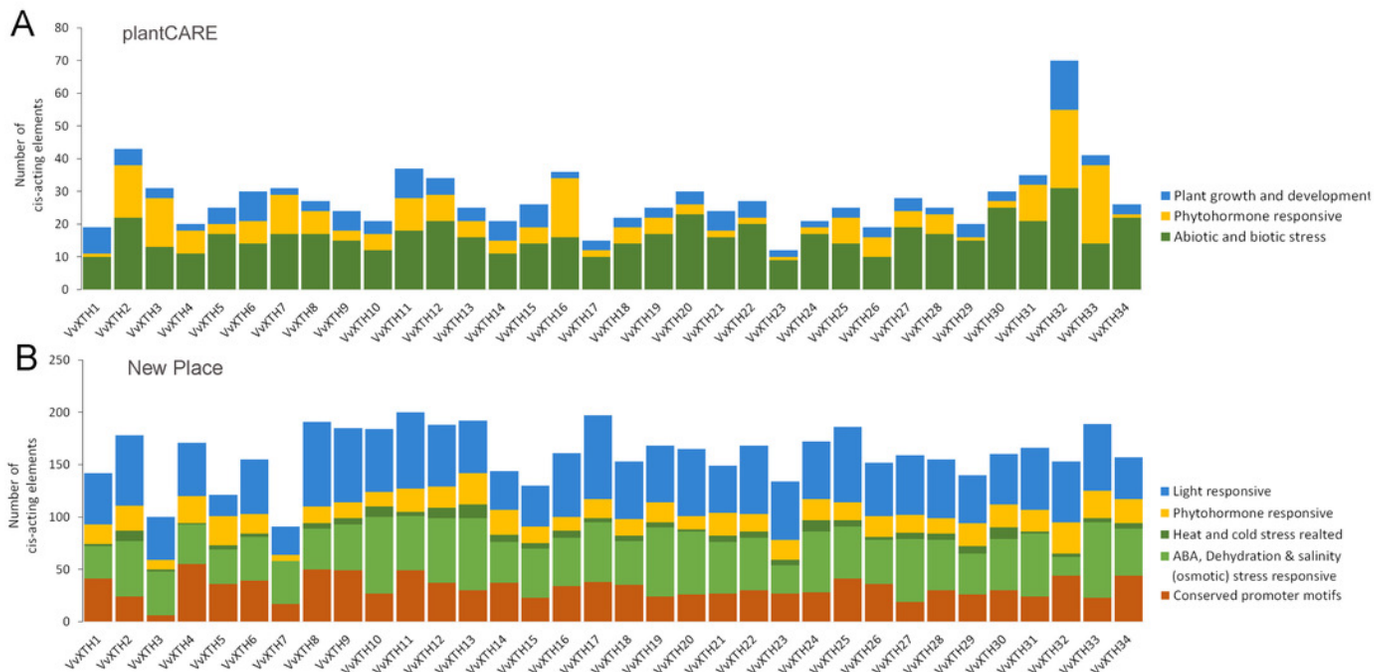


Table 1 (on next page)

Molecular characteristics of *VvXTHs* in grapevine.

AA: amino acid; MW: molecular weight; SP: signal peptide; pI: isoelectric point; GRAVY: total average hydrophilicity.

1 **Table 1 Molecular characteristics of *VvXTHs* in grapevine.**

| Name | Gene Identifier | AA | MW(Da) | SP | pI | GRAVY | Subcellular Localization |
|----------------|--------------------|-----|----------|----|------|--------|--------------------------|
| <i>VvXTH1</i> | VIT_201s0011g06250 | 279 | 32099.88 | – | 6.60 | -0.649 | Plasma membrane |
| <i>VvXTH2</i> | VIT_201s0026g00200 | 313 | 35198.85 | 24 | 6.83 | -0.296 | Extracellular |
| <i>VvXTH3</i> | VIT_201s0150g00460 | 307 | 35270.14 | 35 | 8.65 | -0.366 | Plasma membrane |
| <i>VvXTH4</i> | VIT_202s0012g02220 | 341 | 38867.80 | – | 8.99 | -0.374 | Plasma membrane |
| <i>VvXTH5</i> | VIT_203s0088g00650 | 295 | 34401.83 | 25 | 7.12 | -0.372 | Plasma membrane |
| <i>VvXTH6</i> | VIT_205s0062g00240 | 281 | 32143.11 | 24 | 9.22 | -0.389 | Plasma membrane |
| <i>VvXTH7</i> | VIT_205s0062g00250 | 279 | 32239.31 | 24 | 9.07 | -0.449 | Plasma membrane |
| <i>VvXTH8</i> | VIT_205s0062g00480 | 281 | 32088.01 | 24 | 9.08 | -0.406 | Plasma membrane |
| <i>VvXTH9</i> | VIT_205s0062g00610 | 281 | 32173.18 | 24 | 9.14 | -0.408 | Plasma membrane |
| <i>VvXTH10</i> | VIT_206s0061g00550 | 291 | 32696.72 | 18 | 5.74 | -0.438 | Extracellular |
| <i>VvXTH11</i> | VIT_207s0185g00050 | 280 | 32102.92 | 19 | 7.11 | -0.555 | Plasma membrane |
| <i>VvXTH12</i> | VIT_208s0007g04950 | 293 | 33761.16 | 18 | 9.45 | -0.457 | Extracellular |
| <i>VvXTH13</i> | VIT_210s0116g00520 | 300 | 34816.90 | 27 | 4.61 | -0.584 | Plasma membrane |
| <i>VvXTH14</i> | VIT_210s0003g02440 | 294 | 33673.17 | 27 | 9.44 | -0.375 | Plasma membrane |
| <i>VvXTH15</i> | VIT_210s0003g02480 | 290 | 32860.23 | 27 | 8.18 | -0.278 | Plasma membrane |
| <i>VvXTH16</i> | VIT_211s0016g03480 | 291 | 33246.42 | 17 | 8.24 | -0.338 | Plasma membrane |
| <i>VvXTH17</i> | VIT_211s0052g01180 | 321 | 36502.43 | – | 4.81 | -0.596 | Plasma membrane |
| <i>VvXTH18</i> | VIT_211s0052g01190 | 369 | 41704.56 | – | 6.36 | -0.454 | Plasma membrane |
| <i>VvXTH19</i> | VIT_211s0052g01200 | 297 | 32951.68 | 29 | 5.22 | -0.392 | Plasma membrane |
| <i>VvXTH20</i> | VIT_211s0052g01220 | 285 | 31821.45 | 19 | 5.92 | -0.404 | Plasma membrane |
| <i>VvXTH21</i> | VIT_211s0052g01230 | 278 | 31187.75 | – | 5.13 | -0.405 | Plasma membrane |
| <i>VvXTH22</i> | VIT_211s0052g01250 | 312 | 35059.55 | – | 8.42 | -0.338 | Plasma membrane |
| <i>VvXTH23</i> | VIT_211s0052g01260 | 296 | 32919.61 | 26 | 5.07 | -0.393 | Plasma membrane |
| <i>VvXTH24</i> | VIT_211s0052g01270 | 297 | 33155.89 | 29 | 4.97 | -0.361 | Plasma membrane |
| <i>VvXTH25</i> | VIT_211s0052g01280 | 296 | 32939.66 | 26 | 5.63 | -0.420 | Plasma membrane |
| <i>VvXTH26</i> | VIT_211s0052g01300 | 280 | 31322.79 | 26 | 5.37 | -0.419 | Plasma membrane |
| <i>VvXTH27</i> | VIT_211s0052g01310 | 269 | 30176.71 | 29 | 5.69 | -0.275 | Plasma membrane |
| <i>VvXTH28</i> | VIT_211s0052g01320 | 296 | 33018.79 | 26 | 5.93 | -0.425 | Plasma membrane |
| <i>VvXTH29</i> | VIT_211s0052g01330 | 262 | 29508.94 | 18 | 5.62 | -0.459 | Plasma membrane |
| <i>VvXTH30</i> | VIT_211s0052g01340 | 272 | 29935.10 | 29 | 4.96 | -0.457 | Plasma membrane |
| <i>VvXTH31</i> | VIT_212s0134g00160 | 296 | 33657.80 | 27 | 5.60 | -0.320 | Plasma membrane |
| <i>VvXTH32</i> | VIT_215s0048g02850 | 322 | 37077.07 | 23 | 6.11 | -0.346 | Extracellular |
| <i>VvXTH33</i> | VIT_216s0100g00170 | 251 | 28460.18 | – | 6.65 | -0.232 | Extracellular |
| <i>VvXTH34</i> | VIT_217s0053g00610 | 284 | 32329.43 | 21 | 5.50 | -0.361 | Plasma membrane |

2 AA: amino acid; MW: molecular weight; SP: signal peptide; pI: isoelectric point; GRAVY: total

3 average hydrophilicity.