

Comprehensive genomic characterisation of the NAC transcription factor family and its response to drought stress in *Eucommia ulmoides*

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The NAC transcription factor family enhances plant adaptation to environmental challenges by participating in signalling pathways triggered by abiotic stressors and hormonal cues. We identified 69 NAC genes in the *Eucommia ulmoides* genome and renamed them according to their chromosomal distribution. These EuNAC proteins were clustered into 13 sub-families and distributed on 16 chromosomes and 2 scaffolds. The gene structures suggested that the number of exons varied from 2 to 8 among these *EuNACs*, with a multitude of them containing three exons. Duplicated events resulted in a large gene family; 12 and 4 pairs of *EuNACs* were the result of segmental and tandem duplicates, respectively. The drought-stress response pattern of 12 putative *EuNACs* was observed under drought treatment, revealing that these *EuNACs* could play crucial roles in mitigating the effects of drought stress responses and serve as promising candidate genes for genetic engineering aimed at enhancing the drought stress tolerance of *E. ulmoides*. This study provides insight into the evolution, diversity, and characterisation of NAC genes in *E. ulmoides* and will be helpful for future characterisation of putative *EuNACs* associated with water deficit.

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17

18 Abstract

19 The NAC transcription factor family enhances plant adaptation to environmental challenges by
20 participating in signalling pathways triggered by abiotic stressors and hormonal cues. We
21 identified 69 *NAC* genes in the *Eucommia ulmoides* genome and renamed them according to their
22 chromosomal distribution. These *EuNAC* proteins were clustered into 13 sub-families and
23 distributed on 16 chromosomes and 2 scaffolds. The gene structures suggested that the number of
24 exons varied from 2 to 8 among these *EuNACs*, with a multitude of them containing three exons.
25 Duplicated events resulted in a large gene family; 12 and 4 pairs of *EuNACs* were the result of
26 segmental and tandem duplicates, respectively. The drought-stress response pattern of 12 putative
27 *EuNACs* was observed under drought treatment, revealing that these *EuNACs* could play crucial
28 roles in mitigating the effects of drought stress responses and serve as promising candidate genes
29 for genetic engineering aimed at enhancing the drought stress tolerance of *E. ulmoides*. This study
30 provides insight into the evolution, diversity, and characterisation of *NAC* genes in *E. ulmoides*
31 and will be helpful for future characterisation of putative *EuNACs* associated with water deficit.

32 **Keywords:** *Eucommia ulmoides*, NAC family, drought-responsive, gene expression, phylogenetic
33 analysis

34

35 Introduction

36 *Eucommia ulmoides* Oliver is a highly valued tertiary relict perennial tree species native to
37 China (Deng et al. 2022). It is widely used in industry because it not only produces wood but is
38 also a valuable raw biomaterial for extracting the active ingredients of Chinese medicine and *trans*-
39 rubber (Wei et al. 2021; Zhu & Sun 2018). Compared with *cis*-rubber, *trans*-rubber has unique

40 characteristics, such as high hardness, resistance to acid and alkali corrosion, good insulation, and
41 a low thermal expansion/contraction coefficient (Enoki et al. 2003; Kent & Swinney 1966; Rose
42 & Steinbuechel 2005). To improve the viscosity, resilience, elasticity, weather resistance, and
43 tensile strength of *trans*-rubber, it can be made into *cis*-rubber by vulcanisation (Yan 1995). The
44 vulnerability of *Hevea brasiliensis* to pests and diseases, as well as its narrow habitat, has led to
45 significant challenges for the rubber industry (Tang et al. 2016). *Eucommia ulmoides* has wide
46 adaptability, few pests and diseases, and its leaves, bark, and pericarp are rich in *trans*-rubber.
47 Therefore, *E. ulmoides* is considered an ideal alternative or complementary tree species to *H.*
48 *brasiliensis* (Wuyun et al. 2018). During the growth of *E. ulmoides*, some environmental factors,
49 such as drought and low-temperature stress, can prevent its full genetic potential, resulting in
50 reduced yield and even plant death (Zuo et al. 2022). The identification and utilisation of resistance
51 genes is the basis for breeding resistant varieties. Transcription factors (TFs), which activate or
52 inhibit their expression by specifically binding to *cis*-acting elements on the promoters of target
53 genes, play an important role in many biological processes (Yuan et al. 2020). As a plant-specific
54 supergene family, *NAC* has been demonstrated to play a key role in plant growth and development
55 and in the response to abiotic stress (Du et al. 2022b; Hussain et al. 2017). Notably, *NAC* is very
56 important in plant adaptation to land (Xu et al. 2014). Therefore, *NAC* family genes have been
57 widely studied in many species. However, the identification and analysis of *NAC* genes in *E.*
58 *ulmoides* have not been emphasised.

59 The *NAC* [no apical meristem (NAM), *Arabidopsis* transcription activator (ATAF1/ATAF2),
60 and cup-shaped cotyledon (CUC2)] family is one of the largest gene families in plants. The N-
61 terminus region has a highly conserved NAM domain, and the C-terminus consists of variable
62 transcriptional regulatory regions, the latter of which have been implicated in specific biological
63 functions (Shao et al. 2015). Increasing evidence suggests that *NAC* TFs have multiple functions
64 in plant-responses to biotic and abiotic stress. *SINAC1* is involved in the process of fruit softening
65 and fruit pigmentation based on the phytohormone pathway (Ma et al. 2014). *NAC13* has important
66 significance in popular responses to salt stress (Zhang et al. 2019). In wheat, overexpression of
67 *TaNAC1-D1* and *TaNAC071-A* improves resistance to *Fusarium* head blight (Perochon et al. 2019)
68 and drought (Mao et al. 2022), respectively; *TaNAC30* negatively regulates stripe rust (Wang et
69 al. 2018). Furthermore, *TaNAC29* improves salt tolerance by strengthening the antioxidant system
70 (Xu et al. 2015). *GhirNAC2* regulates ABA biosynthesis and stomatal closure by regulating
71 *GhNCED3a/3c* expression, thus playing an active role in cotton drought resistance (Shang et al.
72 2020). Although *NAC* TFs are related to various developmental processes and stress responses in
73 plants, the specific functions of most *NAC* genes remain obscure, especially in *E. ulmoides*.

74 The chromosome-level genome of *E. ulmoides* was recently sequenced (Li et al. 2020); this
75 provides the opportunity to systematically study the *NAC* gene family and to explore the potential
76 functional involved in *E. ulmoides* biotic and abiotic responses. In the present study, we performed
77 genome-wide identification and characterisation of *NAC* proteins based on the genome of *E.*
78 *ulmoides*. In addition, we surveyed their expression under drought stress by transcriptome

79 sequencing. This study will lay the foundation for further studies of the molecular mechanisms of
80 NAC TFs in *E. ulmoides* response to drought stress.

81 **Materials & Methods**

82 **Identification of EuNAC proteins from the *E. ulmoides* genome**

83 The complete genome data of *E. ulmoides* were obtained from the Gene Warehouse
84 (<https://ngdc.cncb.ac.cn/gwh/Assembly/25206/show>). The NAC protein sequences of *Arabidopsis*
85 were obtained from *Arabidopsis* Information Resources (TAIR,
86 <https://www.arabidopsis.org/index.jsp>), and the protein sequences of *poplar* and rice were both
87 derived from the Ensembl Plants website (<http://plants.ensembl.org/index.html>). The Hidden
88 Markov model (HMM) files of the NAC domain (PF01849) and the NAM domain (PF02365) were
89 obtained from the Pfam database (<https://pfam.sanger.ac.uk>), which were used for identification
90 analysis. HMMER 3.3.2 (<http://hmmer.org/>) was then employed to scan the NAC proteins from
91 the *E. ulmoides* genome with the default parameters. The candidate *EuNACs* were further validated
92 by the NCBI Conserved Domain Search Service (CD Search) (<https://www.ncbi.nlm.nih.gov>),
93 SMART (<http://smart.embl-heidelberg.de>), and Pfam database. Proteins without NAC and NAM
94 domains and duplicates were manually deleted. The molecular weight (MW) and isoelectric point
95 (pI) of each protein were analysed using the ExpASY pI/Mw tool (<https://www.expasy.org>).

96 **Phylogenetic analysis of EuNAC proteins**

97 The *Arabidopsis* NAC protein sequences were downloaded from the TAIR database
98 (<http://www.Arabidopsis.org>). Full-length protein sequence multiple alignments were performed
99 using the ClustalW programme (Larkin et al. 2007). MEGA 6.0 software (Hall. 2013) was
100 employed to construct an unrooted phylogenetic tree of *E. ulmoides* and *Arabidopsis* NAC proteins
101 using the neighbour-joining (NJ) method with 1000 bootstrap iterations. All *EuNAC* proteins were
102 classified according to the NAC protein classification criteria in *Arabidopsis* (Ooka et al. 2003a).

103 **Conserved motif and gene structure analysis of *EuNAC* genes**

104 The Gene Structure Display Server (GSDS; <http://gsds.cbi.pku.edu.cn/>) programme was used
105 to explore the exon/intron structure pattern of the *EuNAC* genes by comparing their predicted
106 coding sequence with the corresponding full-length gDNA sequence. Multiple Expectation
107 Maximization for Motif Elicitation (MEME) (<http://meme-suite.Org/>) programmes were
108 employed to identify the conserved domains for candidate *EuNAC* proteins with default
109 parameters. The conserved motifs and exon/intron structure were visualised using Tbttools (Chen
110 et al. 2020).

111 **Genome distribution, selective pressure, and synteny analysis of *EuNAC* genes**

112 The location of each *EuNAC* gene on the chromosome was determined based on the *E.*
113 *ulmoides* genome annotation file and visualised using Tbttools software (Chen et al. 2020).
114 MCScan X software (Wang et al. 2012) with default parameters was employed to analyse
115 duplication events, and the intra-species and inter-species collinearity relationships. Circos
116 software (Krzywinski et al. 2009) and Tbttools were used for visualisation. Tbttools were also used
117 to calculate the nonsynonymous (Ka) and synonymous (Ks) rates of *EuNAC* homologous genes.
118 The selection pressure acting on the gene pairs was calculated based on the Ka/Ks ratio, and the

119 dates of each duplication event were further deduced with the formula $T = Ks/2\lambda$, the mean
120 synonymous substitution rate (λ) was assumed to be 6.5×10^{-9} (Liu et al. 2021; Lynch & Conery
121 2000a).

122 Promoter region analysis of *EuNAC* genes

123 The 2-kb promoter sequences upstream of the *EuNAC* genes start codon (ATG) were
124 extracted, and the cis-acting elements and their potential related functions were predicted with the
125 PlantCARE online server (<http://bioinformatics.psb.ugent.be/webtools/plantcare/html/>). The 20
126 cis-acting elements with the highest frequency were visualised using Tbttools (Chen et al. 2020).

127 Expression analysis of *EuNAC* genes under drought stress

128 Two-year-old ‘Qinzhong 1’ grafted potted plants with consistent growth were placed in the
129 Agricultural College of Shihezi University and well managed in the natural environment. Three
130 months later, they were subjected to drought stress treatment. For drought treatment, the soil was
131 saturated with water, and watering was terminated. Leaves were collected at 0, 15, 30, and 45 d
132 and labelled CK, D15, D30, and D45, respectively. Each sample was pooled from six individual
133 plants, and three biological replicates were set for each treatment. Samples were quickly cleaned
134 with distilled water, immediately frozen in liquid nitrogen, and stored at -80°C for future use.

135 The above samples were submitted to Beijing Novogen Bioinformatics Technology Co., Ltd.
136 (Beijing, China) for cDNA library construction and transcriptome sequencing. The data were
137 uploaded to the National Center for Biotechnology Information (NCBI) Sequence Read Archive,
138 with accession number PRJNA961078. Gene expression levels were normalised with FPKM
139 (fragment per kilobase per million mapped reads) and visualised with TBtools.

140

141 Results

142 Identification of *EuNAC* proteins from the *E. ulmoides* genome

143 We identified 69 *NAC* genes in the *E. ulmoides* genome, named *EuNAC1* TO *EuNAC69*,
144 according to their order on the chromosomes (Table S1). The *EuNAC* proteins varied significantly
145 in length and molecular weight. The length of proteins encoded by *EuNAC* genes ranged from 86
146 (*EuNAC50*) to 617 (*EuNAC64*) amino acids (aa), and the molecular weight varied from 9.81
147 (*EuNAC50*) to 70.46 kDa (*EuNAC64*); the isoelectric points (pIs) ranged from 4.51 (*EuNAC59*) to
148 10.01 (*EuNAC2*). This study also analysed other basic information about 69 *NAC* genes, including
149 homologous genes in *Arabidopsis*, open reading frame (ORF) length, location coordinates,
150 chromosomal positions, and exon numbers (Table S1).

151 Phylogenetic analysis and classification of *EuNAC* proteins

152 To investigate the evolutionary relationships among *EuNAC* family genes, MEGA6.0
153 software was employed to construct a neighbour-joining phylogenetic tree based on full-length
154 protein sequences of *NAC* in *E. ulmoides* and *Arabidopsis* (Figure 1). According to the
155 classification system for *Arabidopsis*, *NACs* from *E. ulmoides* were divided into 13 distinct
156 subfamilies, namely *NAC2*, *ONAC022*, *AtNAC3*, *NAP*, *ANAC063*, *ANAC011*, *ONAC003*, *NAM*,
157 *NAC1*, *OSNAC7*, *OSNAC8*, *TIP*, and *ANAC011*, using a phylogenetic tree; however, no *EuNAC*
158 members were identified in the *ATAF* subfamily. Of these 13 subfamilies, *OSNAC7* had 11 *EuNAC*

159 members, which was the most abundant, followed by *NAM*, which had eight members. *ANAC001*
160 was the least frequent with only two members. Phylogenetic analysis revealed that EuNAC
161 proteins had evolved in some diversity, similar to a report in *Arabidopsis* (Ooka et al. 2003a).

162 **Conserved motif and gene structure analysis of *EuNAC* genes**

163 To gain insight into the functional regions of EuNACs, the conserved motifs for each EuNAC
164 protein were analysed using the MEME programme. A total of 10 conserved motifs were identified
165 and named motifs 1–10 (Table S2). These conserved motifs had large variations in length, with a
166 distribution range of 11–50 amino acid residues. As shown in Figure 2A, motif 3 had the highest
167 frequency in the *EuNAC* family, and it existed in almost all members except *EuNAC2*, *19*, and *33*.
168 In addition, motifs 1, 2, 4, 5, and 6 were very abundant in the *EuNAC* family, but none of the
169 *ONAC003* subfamily members had these motifs. Most of the conserved motifs were distributed in
170 the N-terminus of the NAC proteins, indicating that the N-terminal region plays an important role
171 in *NAC* gene function. In addition, similar motif compositions existed among different members
172 of the same subfamily, indicating that members of the same subfamily had similar functions.

173 To investigate the structural features of *EuNACs*, we analysed the intron/exon distribution
174 patterns of each *EuNAC* gene. The exon distribution within the *EuNAC* genes varied from 2 to 8
175 (Table S1, Figure 2B). Forty-five (65.2%) genes had three exons, 10 (14.5%) genes had six exons,
176 and *EuNAC45* had the largest number of exons, with 8 exons (Figure 2B, Table S1). Forty-nine
177 (71.0%) *EuNAC* genes possessed less than three exons, indicating a low structural diversity among
178 *EuNAC* genes.

179 **Genome distribution, selective pressure, and synteny analysis of *EuNAC* genes**

180 To investigate the distribution of *EuNAC* genes on the chromosomes of *E. ulmoides*, TBtools
181 software was employed to map the chromosome locations for all *EuNACs* identified in this study
182 (Figure 3). The 69 *EuNACs* were unevenly scattered on 16 chromosomes and two scaffolds, and
183 the length of each chromosome showed no correlation with the number of genes contained.
184 Chromosomes 8 and 12 had the most *EuNACs*, both with seven genes. Only one *EuNAC* was
185 distributed on chromosomes 3 and 16, and no *EuNACs* were distributed on chromosome 11.
186 Notably, most *EuNACs* were distributed near the ends of the chromosome.

187 Furthermore, the duplication events of *EuNAC* gene family members were examined using
188 MCScanX software. A total of 12 pairs of segmental duplications were identified in the *EuNAC*
189 family, which were distributed across 15 chromosomes, except for chromosomes 3 and 11, and
190 four pairs of tandem replications (*EuNAC7/8*, *EuNAC30/31*, *EuNAC48/49*, and *EuNAC53/54*)
191 were identified, which were distributed on chromosomes 2, 7, 12, and 13, respectively (Figure 4,
192 Table S3). The results showed that segmental duplication events might be the crucial driving force
193 in *EuNAC* gene family expansion. To evaluate the selection pressure of *EuNACs*, the *Ka*, *Ks*, and
194 *Ka/Ks* for duplicated gene pairs were calculated. In general, a *Ka/Ks* > 1 indicates positive
195 selection, *Ka/Ks* = 1 indicates neutral selection, while *Ka/Ks* < 1 indicates purifying selection
196 (Vahdati & Lotfi 2013). The *Ka/Ks* ratios of 16 replicated *EuNAC* gene pairs were all less than 1,
197 indicating that the evolution of the *EuNAC* gene family was subjected to purification selection
198 (Table S4).

199 To further understand the phylogenetic mechanisms of the *EuNAC* gene family, we
200 constructed a comparative homologous map of *NAC* genes in *E. ulmoides*, *Arabidopsis*, and rice.
201 In total, 31 *EuNACs* had a collinear relationship with 27 *AtNACs* and 10 *OsNACs*. Thirty-one and
202 eleven pairs of *NAC* homologous gene pairs were formed between *E. ulmoides* and *Arabidopsis*
203 and between *E. ulmoides* and rice, respectively (Figure 5, Table S5). The results indicate that the
204 *NAC* genes underwent significant evolution and replication after differentiation in
205 monocotyledonous and dicotyledonous plants.

206 **Promoter region analysis of *EuNAC* genes**

207 To investigate the potential functions of *EuNACs*, we employed PlantCARE to predict the
208 cis-acting elements within the 2.0 kb sequence upstream of the initiation codon (ATG) of *EuNACs*
209 (Table S6). As expected, both TATA and CAAT boxes with good characteristics were found in
210 the results; we also found several other CIS regulators (Table S7 and Figure S1). They were mined
211 in the promoter region of *EuNACs*. As shown in Figure 6, we divided the homeopathic elements
212 into four categories according to their functions. The first category was phytohormone-responsive
213 elements, such as CGTCA-motif, TGACG, TCA-element, and ABRE, wherein ABREs have been
214 associated with ABA responses and TCA-element have been associated with salicylic acid
215 responsiveness. The second category was elements of cis-regulation related to the response to
216 external or environmental pressure. This category includes low-temperature response elements
217 (LTR), which respond to external abiotic stress, abundant cis-regulatory elements (AREs), which
218 are required for anaerobic induction, and MYB binding site (MBS) elements. Notably, 29 of the
219 69 *EuNAC* promoters contained MBS elements that were involved in drought induction as MYB
220 binding sites and could be predicted based on their responses to drought stress treatments. The
221 third category included light-responsive elements, such as G-box, Box-4, and GT1-motif, in which
222 at least one photo-responsive element was detected in almost every promoter region of *EuNACs*.
223 The last category was cis-regulatory elements related to growth and development. CAT-box and
224 O2-site were mainly detected, which also indicated that most *EuNACs* may be involved in *E.*
225 *ulmoides* meristem expression, zein metabolism regulation, and cell cycle regulation. Finally,
226 based on the above results, *EuNACs* may be involved in stress response, light, hormones, and
227 growth pathways.

228 **Expression analysis of *EuNAC* genes under drought stress**

229 To further investigate the potential function of *EuNACs* in response to drought stress,
230 comparative transcriptomics of *E. ulmoides* under drought stress were used to analyse the
231 expression patterns of *EuNACs*. Of the 69 *EuNAC* genes, 20 showed high expression levels with
232 $\text{FPKM} \geq 20$, including *EuNAC1*, 2, 12, 15, 20, and 25. Forty-three *EuNACs* showed low expression
233 levels, with $\text{FPKM} \leq 20$, including *EuNAC10*, 11, 13, 14, 16, 17, and 18. In addition, *EuNAC19*,
234 47, 52, 54, 55, and 58 were not expressed in *E. ulmoides* leaves (Table S8). Differential gene
235 expression (DEG) analysis showed that 12 *EuNAC* genes were significantly differentially
236 expressed, of which *EuNAC2*, 3, 11, and 14 were upregulated after drought stress treatment, and
237 *EuNAC1*, 36, 37, 64, and 66 were downregulated. The expression levels of *EuNAC8* and 13 first
238 decreased and then increased with prolonged treatment time; in contrast, the expression level of

239 *EuNAC61* first increased and then decreased (Figure 7, Table S8). The variable expression patterns
240 of *EuNACs* may indicate their differential roles in the drought stress response of *E. ulmoides*. In
241 particular, differentially expressed *EuNACs* may play a crucial role in *E. ulmoides*' response to
242 drought stress.

243

244 Discussion

245 The NAC transcription factor family is one of the largest gene families in plants. These factors
246 are involved in regulating hormone signalling pathways, biotic and abiotic stress responses, and
247 plant growth and development (Yuan et al. 2020; Zhang et al. 2019). Several plant genomes have
248 defined the *NAC* gene family. The evolutionary connection and duplication patterns of the *NAC*
249 gene family in *E. ulmoides* can be better understood thanks to the genome of this organism.
250 Previous studies have shown that genes with close evolutionary relationships often share similar
251 functions (Lynch & Conery 2000b). Therefore, by studying the evolutionary relationships across
252 gene families, we can learn more about and perhaps even anticipate how genes operate (Balazadeh
253 et al. 2011; Zhang et al. 2019).

254 According to our study, the genome of *E. ulmoides* included 69 *NAC* genes, which is fewer
255 than that of *A. thaliana* (117 *NAC* genes) (Ooka et al. 2003b), but similar to *K. obovate* (79 *NAC*
256 genes) (Du et al. 2022a). Our results indicate that the majority of *EuNACs* did not experience
257 environmental selection-induced elimination, but rather demonstrated a high level of conservation
258 throughout evolution, underlining the necessity for more research from an evolutionary standpoint.
259 All 69 *NAC* proteins were divided into 13 subgroups based on their sequence homology and
260 classification relative to *Arabidopsis* (Ooka et al. 2003b). *NACs* in *Arabidopsis* exhibit a high
261 degree of similarity among members of the same class or *NAC* subgroup. Four *EuNACs* in the
262 *ANAC011* subgroup are orthologous to *Arabidopsis* genes, including *AtNAC071* and *AtNAC096*,
263 which are in charge of tissue reunification, dehydration, and other processes.

264 Our research indicates that the *NAM* subgroup has 8 *EuNACs* that are orthologous to
265 *AtNAC054* and *AtNAC059* in *Arabidopsis*, which are known to be crucial for organ development,
266 programmed cell death, secondary wall building, and biotic and abiotic stress responses (Kim et
267 al. 2007). The 7 *EuNACs* in subgroup *NAP* are orthologous to *AtNAC018*, *AtNAC025*, and
268 *AtNAC56* and are essential for leaf senescence (Guo & Gan 2006). Three genes of the *EuNAC* gene
269 family's subgroup *TIP* are orthologs of the *Arabidopsis* genes *AtNAC060* and *AtNAC091*. These
270 orthologous genes have been demonstrated to be crucial in the stress response and abscisic acid
271 (ABA) signalling (Donze et al. 2014; Jeong et al. 2008; Li et al. 2014). Four *EuNACs* in the
272 *ANAC011* subgroup are orthologous to *Arabidopsis* genes, including *AtNAC071* and *AtNAC096*,
273 which are in charge of tissue reunification, dehydration, and osmotic stress (Asahina et al. 2011;
274 Yang et al. 2020). Five orthologous *EuNACs* to *AtNAC016*, which are known to be involved in
275 chlorophyll degradation, are found in the *NAC2* subgroup. This indicates that this subgroup of
276 *EuNACs* may also control how chlorophyll degrades in plants (Sakuraba et al. 2015). Similar to
277 *AtNAC003* and *AtNAC068*, the *ANAC001* subgroup also has two *EuNACs* that are orthologous to

278 them. These genes control salt and osmotic stress tolerance, in addition to DNA damage responses
279 (Xu et al. 2013; Yoshiyama et al. 2014).

280 Except for individuals from the *ONAC003* and *ANAC063* subgroups in this investigation, the
281 N-terminus of the EuNAC protein had motifs 1, 2, 3, 4, 5, and 6. *ONAC7* comprised motifs 7 and
282 10, indicating that NAC transcription factors had a highly conserved N-terminus and a very varied
283 C-terminus. The range of *EuNAC* introns was 2 to 8, which is comparable to the number observed
284 in many plants (Du et al. 2022a; Liu et al. 2019). In general, the deletion or insertion of introns
285 can have diverse effects on gene function. Sometimes, intron deletions can lead to the loss of gene
286 function if they result in a frameshift mutation or the removal of critical regulatory sequences. Our
287 analysis supported the findings of Jeffares' research, which demonstrated that genes susceptible to
288 abrupt changes in stress expression levels have much less intron density (Jeffares et al. 2008);
289 *EuNACs* with fewer introns merit higher consideration if the study objective is to concentrate on
290 genes that react instantly to environmental stress.

291 As a consequence of our findings, which included the identification of 12 segmental
292 duplications and four tandem duplications in 69 *EuNACs*, we concluded that segmental duplication
293 served as the primary catalyst for the growth of the *EuNAC* gene family, in agreement with
294 research on *K. obovata* (Du et al. 2022a). In addition, larger segmental duplications account for
295 most of the *A. thaliana* genome, and at least four large-scale replication events occurred during
296 the formation of angiosperm diversity (100–200 million years ago) (Vision et al. 2000). This might
297 explain why there are more *NAC* members in *Arabidopsis*, while having a smaller genome than *E.*
298 *ulmoides*. Only 10 genes, according to our research, have collinear connections between *O. sativa*
299 and *E. ulmoides*. However, we discovered 27 orthologous pairings in *Arabidopsis*, a
300 dicotyledonous plant. These findings demonstrate a closer evolutionary link between dicotyledons
301 and *EuNACs* than between monocotyledons.

302 Cis-acting elements are specific DNA sequences located in the promoter region of genes that
303 serve as binding sites for transcription factors (Liu et al. 2016; Kaur et al. 2017). In this study,
304 more than half of the 69 *EuNAC* promoters included ABRE homeopathic elements, suggesting that
305 these genes may operate via the control of ABA. MBS elements were found in 29 *EuNACs*,
306 indicating that these genes may be crucial in response to drought stress.

307 Research on gene function can benefit from understanding the patterns of gene expression.
308 According to RNA-seq studies, drought stress drastically altered the expression levels of a few
309 *EuNACs* in the leaves of *E. ulmoides*. Significant differential expression was seen in 12 *EuNACs*.
310 *EuNAC1*, *EuNAC8*, and *EuNAC36* are identical to *ANAC019* (*AT1G52890*), *ANAC055*
311 (*AT3G15500*), and *ANAC072* (*AT4G27410*), which are members of the *AtNAC3* subgroup. Their
312 expression is variably expressed during drought treatment, caused by drought, and stimulated by
313 ABA (Tran et al. 2004). Therefore, we hypothesise that genes *EuNAC1*, *EuNAC8*, and *EuNAC36*
314 belong to the same subgroup and are drought-responsive genes that control *E. ulmoides*' survival
315 ability in drought-stressed environments. Additionally, *ANAC054* (*At3g15170*) and *ANAC059*
316 (*At3g29035*) were identical to *EuNAC3*, *EuNAC11*, and *EuNAC61* and were grouped into the *NAM*
317 subgroup, suggesting that they may be crucial in *E. ulmoides*' response to drought stress.

318

319 Conclusions

320 In summary, 69 *NACs* were identified in *E. ulmoides* in this study. We studied the
321 characteristics of *EuNAC* genes at the genomic level and analysed tissue expression patterns and
322 responses to drought stress. These TFs can be divided into 13 subgroups according to the *NAC*
323 classification method of *Arabidopsis*. Chromosomal localisation and homology analysis showed
324 that segmental duplication was the main driving force for *EuNAC* gene amplification. Genome-
325 wide expression analysis of *NAC* genes in response to drought provides an opportunity to further
326 understand the strong tolerance mechanism of *E. ulmoides* to drought.

327

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331

332 References

- 333 Asahina M, Azuma K, Pitaksaringkarn W, Yamazaki T, Mitsuda N, Ohme-Takagi M, Yamaguchi
334 S, Kamiya Y, Okada K, Nishimura T, Koshihara T, Yokota T, Kamada H, and Satoh S. 2011.
335 Spatially selective hormonal control of RAP2.6L and ANAC071 transcription factors
336 involved in tissue reunion in *Arabidopsis*. *Proceedings of the National Academy of*
337 *Sciences* 108:16128-16132.
- 338 Balazadeh S, Kwasniewski M, Caldana C, Mehrnia M, Zhanor MI, Xue GP, and Mueller-Roeber
339 B. 2011. ORS1, an H₂O₂-responsive NAC transcription factor, controls senescence in
340 *Arabidopsis thaliana*. *Mol Plant* 4:346-360.
- 341 Chen C, Chen H, Zhang Y, Thomas HR, Frank MH, He Y, and Xia R. 2020. TBtools: an integrative
342 toolkit developed for interactive analyses of big biological data. *Molecular plant* 13:1194-
343 1202.
- 344 Deng P, Wang Y, Hu F, Yu H, Liang Y, Zhang H, Wang T, Zhou Y, and Li Z. 2022. Phenotypic
345 Trait Subdivision Provides New Sight Into the Directional Improvement of *Eucommia*
346 *ulmoides* Oliver. *Frontiers in Plant Science* 13.
- 347 Donze T, Qu F, Twigg P, and Morris TJ. 2014. Turnip crinkle virus coat protein inhibits the basal
348 immune response to virus invasion in *Arabidopsis* by binding to the NAC transcription
349 factor TIP. *Virology* 449:207-214.
- 350 Du Z, You S, Yang D, Tao Y, Zhu Y, Sun W, Chen Z, and Li J. 2022a. Comprehensive analysis
351 of the NAC transcription factor gene family in *Kandelia obovata* reveals potential members
352 related to chilling tolerance. *Front Plant Sci* 13:1048822.
- 353 Du Z, You S, Yang D, Tao Y, Zhu Y, Sun W, Chen Z, and Li J. 2022b. Comprehensive analysis
354 of the NAC transcription factor gene family in *Kandelia obovata* reveals potential members
355 related to chilling tolerance. *Frontiers in Plant Science* 13:4735.
- 356 Enoki M, Doi Y, and Iwata T. 2003. Oxidative degradation of cis-and trans-1, 4-polyisoprenes and
357 vulcanized natural rubber with enzyme-mediator systems. *Biomacromolecules* 4:314-320.
- 358 Guo Y, and Gan S. 2006. AtNAP, a NAC family transcription factor, has an important role in leaf
359 senescence. *Plant J* 46:601-612.

- 360 Hall BG. 2013. Building phylogenetic trees from molecular data with MEGA. *Molecular biology*
361 *and evolution* 30:1229-1235.
- 362 Hussain RM, Ali M, Feng X, and Li X. 2017. The essence of NAC gene family to the cultivation
363 of drought-resistant soybean (*Glycine max* L. Merr.) cultivars. *BMC plant biology* 17:1-
364 11.
- 365 Jeffares DC, Penkett CJ, and Bähler J. 2008. Rapidly regulated genes are intron poor. *Trends Genet*
366 24:375-378.
- 367 Jeong R-D, Chandra-Shekara A, Kachroo A, Klessig D, and Kachroo P. 2008. HRT-Mediated
368 Hypersensitive Response and Resistance to Turnip crinkle virus in *Arabidopsis* Does Not
369 Require the Function of *TIP*, the Presumed Guardee Protein. *Molecular plant-microbe*
370 *interactions : MPMI* 21:1316-1324.
- 371 Kaur A, Pati PK, Pati AM, and Nagpal AK. 2017. In-silico analysis of cis-acting regulatory
372 elements of pathogenesis-related proteins of *Arabidopsis thaliana* and *Oryza sativa*. *PLoS*
373 *One* 12:e0184523.
- 374 Kent E, and Swinney F. 1966. Properties and applications of trans-1, 4-polyisoprene. *Industrial &*
375 *Engineering Chemistry Product Research and Development* 5:134-138.
- 376 Kim SG, Kim SY, and Park CM. 2007. A membrane-associated NAC transcription factor regulates
377 salt-responsive flowering via FLOWERING LOCUS T in *Arabidopsis*. *Planta* 226:647-
378 654.
- 379 Krzywinski M, Schein J, Birol I, Connors J, Gascoyne R, Horsman D, Jones SJ, and Marra MA.
380 2009. Circos: an information aesthetic for comparative genomics. *Genome research*
381 19:1639-1645.
- 382 Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, Valentin F,
383 Wallace IM, Wilm A, and Lopez R. 2007. Clustal W and Clustal X version 2.0.
384 *bioinformatics* 23:2947-2948.
- 385 Li P, Zhou H, Shi X, Yu B, Zhou Y, Chen S, Wang Y, Peng Y, Meyer RC, Smeekens S, and Teng
386 S. 2014. The ABI4-Induced *Arabidopsis ANAC060* Transcription Factor Attenuates ABA
387 Signaling and Renders Seedlings Sugar Insensitive when Present in the Nucleus. *PLoS*
388 *Genetics* 10.
- 389 Li Y, Wei H, Yang J, Du K, Li J, Zhang Y, Qiu T, Liu Z, Ren Y, and Song L. 2020. High-quality
390 de novo assembly of the *Eucommia ulmoides* haploid genome provides new insights into
391 evolution and rubber biosynthesis. *Horticulture Research* 7.
- 392 Liu J, Wang X, Chen Y, Liu Y, Wu Y, Ren S, and Li L. 2021. Identification, evolution and
393 expression analysis of WRKY gene family in *Eucommia ulmoides*. *Genomics* 113:3294-
394 3309.
- 395 Liu M, Ma Z, Sun W, Huang L, Wu Q, Tang Z, Bu T, Li C, and Chen H. 2019. Genome-wide
396 analysis of the NAC transcription factor family in Tartary buckwheat (*Fagopyrum*
397 *tataricum*). *BMC Genomics* 20:113.
- 398 Liu Y, Sun J, and Wu Y. 2016. *Arabidopsis ATAF1* enhances the tolerance to salt stress and ABA
399 in transgenic rice. *J Plant Res* 129:955-962. 10.1007/s10265-016-0833-0
- 400 Lynch M, and Conery JS. 2000a. The evolutionary fate and consequences of duplicate genes.
401 *Science* 290:1151-1155.
- 402 Lynch M, and Conery JS. 2000b. The evolutionary fate and consequences of duplicate genes.
403 *Science* 290:1151-1155.

- 404 Ma N, Feng H, Meng X, Li D, Yang D, Wu C, and Meng Q. 2014. Overexpression of tomato
405 SINAC1 transcription factor alters fruit pigmentation and softening. *BMC plant biology*
406 14:1-14.
- 407 Mao H, Li S, Chen B, Jian C, Mei F, Zhang Y, Li F, Chen N, Li T, and Du L. 2022. Variation in
408 cis-regulation of a NAC transcription factor contributes to drought tolerance in wheat.
409 *Molecular plant* 15:276-292.
- 410 Ooka H, Satoh K, Doi K, Nagata T, Otomo Y, Murakami K, Matsubara K, Osato N, Kawai J, and
411 Carninci P. 2003a. Comprehensive analysis of NAC family genes in *Oryza sativa* and
412 *Arabidopsis thaliana*. *DNA research* 10:239-247.
- 413 Ooka H, Satoh K, Doi K, Nagata T, Otomo Y, Murakami K, Matsubara K, Osato N, Kawai J,
414 Carninci P, Hayashizaki Y, Suzuki K, Kojima K, Takahara Y, Yamamoto K, and Kikuchi
415 S. 2003b. Comprehensive analysis of NAC family genes in *Oryza sativa* and *Arabidopsis*
416 *thaliana*. *DNA Res* 10:239-247.
- 417 Perochon A, Kahla A, Vranić M, Jia J, Malla KB, Craze M, Wallington E, and Doohan FM. 2019.
418 A wheat NAC interacts with an orphan protein and enhances resistance to *Fusarium* head
419 blight disease. *Plant biotechnology journal* 17:1892-1904.
- 420 Rose K, and Steinbuchel A. 2005. Biodegradation of natural rubber and related compounds: recent
421 insights into a hardly understood catabolic capability of microorganisms. *Applied and*
422 *environmental microbiology* 71:2803-2812.
- 423 Sakuraba Y, Piao W, Lim JH, Han SH, Kim YS, An G, and Paek NC. 2015. Rice *ONAC106*
424 Inhibits Leaf Senescence and Increases Salt Tolerance and Tiller Angle. *Plant Cell Physiol*
425 56:2325-2339. 10.1093/pcp/pcv144
- 426 Shang X, Yu Y, Zhu L, Liu H, Chai Q, and Guo W. 2020. A cotton NAC transcription factor
427 GhirNAC2 plays positive roles in drought tolerance via regulating ABA biosynthesis.
428 *Plant Science* 296:110498.
- 429 Shao H, Wang H, and Tang X. 2015. NAC transcription factors in plant multiple abiotic stress
430 responses: progress and prospects. *Frontiers in Plant Science* 6:902.
- 431 Tang C, Yang M, Fang Y, Luo Y, Gao S, Xiao X, An Z, Zhou B, Zhang B, and Tan X. 2016. The
432 rubber tree genome reveals new insights into rubber production and species adaptation.
433 *Nature plants* 2:1-10.
- 434 Tran LS, Nakashima K, Sakuma Y, Simpson SD, Fujita Y, Maruyama K, Fujita M, Seki M,
435 Shinozaki K, and Yamaguchi-Shinozaki K. 2004. Isolation and functional analysis of
436 *Arabidopsis* stress-inducible NAC transcription factors that bind to a drought-responsive
437 cis-element in the early responsive to dehydration stress 1 promoter. *Plant Cell* 16:2481-
438 2498.
- 439 Vahdati K, and Lotfi N. 2013. Abiotic stress tolerance in plants with emphasizing on drought and
440 salinity stresses in walnut. *Abiotic stress—Plant responses and applications in agriculture*
441 10:307-365.
- 442 Vision TJ, Brown DG, and Tanksley SD. 2000. The origins of genomic duplications in
443 *Arabidopsis*. *Science* 290:2114-2117.
- 444 Wang B, Wei J, Song N, Wang N, Zhao J, and Kang Z. 2018. A novel wheat NAC transcription
445 factor, TaNAC30, negatively regulates resistance of wheat to stripe rust. *Journal of*
446 *integrative plant biology* 60:432-443.
- 447 Wang Y, Tang H, Debarry JD, Tan X, Li J, Wang X, Lee T-h, Jin H, Marler B, and Guo H. 2012.
448 MCSanX: a toolkit for detection and evolutionary analysis of gene synteny and
449 collinearity. *Nucleic Acids Research* 40:e49-e49.

- 450 Wei X, Peng P, Peng F, and Dong J. 2021. Natural polymer *Eucommia ulmoides* rubber: A novel
451 material. *Journal of agricultural and food chemistry* 69:3797-3821.
- 452 Wuyun T-n, Wang L, Liu H, Wang X, Zhang L, Bennetzen JL, Li T, Yang L, Liu P, and Du L.
453 2018. The hardy rubber tree genome provides insights into the evolution of polyisoprene
454 biosynthesis. *Molecular plant* 11:429-442.
- 455 Xu B, Ohtani M, Yamaguchi M, Toyooka K, Wakazaki M, Sato M, Kubo M, Nakano Y, Sano R,
456 and Hiwatashi Y. 2014. Contribution of NAC transcription factors to plant adaptation to
457 land. *Science* 343:1505-1508.
- 458 Xu Z, Wang C, Xue F, Zhang H, and Ji W. 2015. Wheat NAC transcription factor TaNAC29 is
459 involved in response to salt stress. *Plant Physiology and Biochemistry* 96:356-363.
- 460 Xu ZY, Kim SY, Hyeon do Y, Kim DH, Dong T, Park Y, Jin JB, Joo SH, Kim SK, Hong JC,
461 Hwang D, and Hwang I. 2013. The Arabidopsis NAC transcription factor ANAC096
462 cooperates with bZIP-type transcription factors in dehydration and osmotic stress
463 responses. *Plant Cell* 25:4708-4724.
- 464 Yan R. 1995. Prospects and research progress on *Eucommia ulmoides* gum. *Progress in Chemistry*
465 7:65.
- 466 Yang JH, Lee KH, Du Q, Yang S, Yuan B, Qi L, and Wang H. 2020. A membrane-associated
467 NAC domain transcription factor XVP interacts with TDIF co-receptor and regulates
468 vascular meristem activity. *New Phytol* 226:59-74.
- 469 Yoshiyama KO, Kimura S, Maki H, Britt AB, and Umeda M. 2014. The role of SOG1, a plant-
470 specific transcriptional regulator, in the DNA damage response. *Plant Signal Behav*
471 9:e28889.
- 472 Yuan C, Li C, Lu X, Zhao X, Yan C, Wang J, Sun Q, and Shan S. 2020. Comprehensive genomic
473 characterization of NAC transcription factor family and their response to salt and drought
474 stress in peanut. *BMC plant biology* 20:1-21.
- 475 Zhang X, Cheng Z, Zhao K, Yao W, Sun X, Jiang T, and Zhou B. 2019. Functional characterization
476 of poplar NAC13 gene in salt tolerance. *Plant Science* 281:1-8.
- 477 Zhu M-Q, and Sun R-C. 2018. *Eucommia ulmoides* Oliver: a potential feedstock for bioactive
478 products. *Journal of agricultural and food chemistry* 66:5433-5438.
- 479 Zuo Y, Li B, Guan S, Jia J, Xu X, Zhang Z, Lu Z, Li X, and Pang X. 2022. EuRBG10 involved in
480 indole alkaloids biosynthesis in *Eucommia ulmoides* induced by drought and salt stresses.
481 *Journal of Plant Physiology* 278:153813.
482

483 **Figures captions**

484 Figure 1. Phylogenetic relationships among *NACs* identified in *E. ulmoides* and *Arabidopsis thaliana*. The
485 unrooted phylogenetic tree was constructed by MEGA 6.0 software using the Neighbor-Joining
486 (NJ) method with 1,000 bootstrap iterations. Each subfamily was distinguished by different
487 colors.

488 Figure 2. Motif compositions and DNA structures of *NAC* gene family in *E. ulmoides*. A. The conserved
489 motif distribution of *EuNAC* proteins. Different motifs were distinguished by different colored
490 boxes, and black lines represent non-conserved regions. B. Gene structure of the *EuNAC* gene.
491 Green boxes represent non-coding regions, yellow boxes represent exons, and black lines
492 represent introns.

493 Figure 3. Distribution of 69 *EuNACs* on 16 chromosomes and two scaffolds. Vertical bars represent the
494 chromosomes of *E. ulmoides*. The chromosome number is on the left of each chromosome. The
495 scale on the left represents the length of the chromosome.

496 Figure 4. Schematic representations of the interchromosomal relationships of *EuNAC* genes. The deep red
497 line represents the *EuNAC* gene pairs replicated in tandem, while the remaining colored lines
498 represent the *EuNAC* gene pairs replicated in segments.

499 Figure 5. Synteny analysis of *NAC* genes between *E. ulmoides* and two representative plant species
500 (*Arabidopsis thaliana* and *Oryza sativa*). Green and purple lines represent syntenic *NAC* gene
501 pairs of *E. ulmoides* and *A. thaliana* and *O. sativa*, respectively.

502 Figure 6. The number of each type of cis-acting element in the promoter region of each *EuNAC* gene.

503 Figure 7. Expression levels of 69 *NAC* genes under drought stress in *E. ulmoides* of leaves. The expression
504 level was presented based on the transformed data of \log_2 (FPKM+1) values.

Figure 1

Phylogenetic relationships among NACs identified in *E. ulmoides* and *Arabidopsis thaliana*.

The unrooted phylogenetic tree was constructed by MEGA 6.0 software using the Neighbor-Joining (NJ) method with 1,000 bootstrap iterations. Each subfamily was distinguished by different colors.

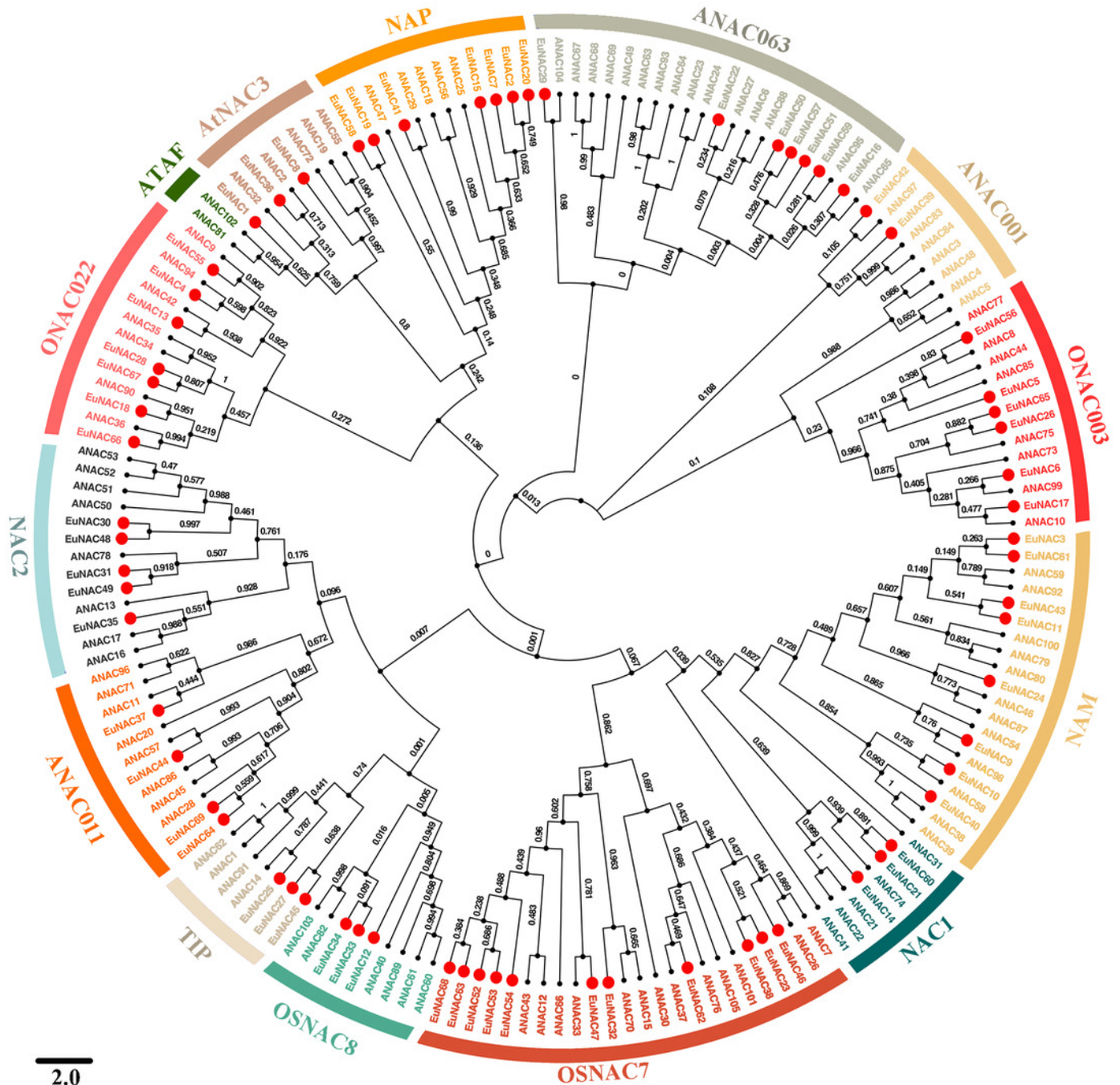


Figure 2

Motif compositions and DNA structures of *NAC* gene family in *E. ulmoides*.

A. The conserved motif distribution of EuNAC proteins. Different motifs were distinguished by different colored boxes, and black lines represent non-conserved regions. B. Gene structure of the *EuNAC* gene. Green boxes represent non-coding regions, yellow boxes represent exons, and black lines represent introns.

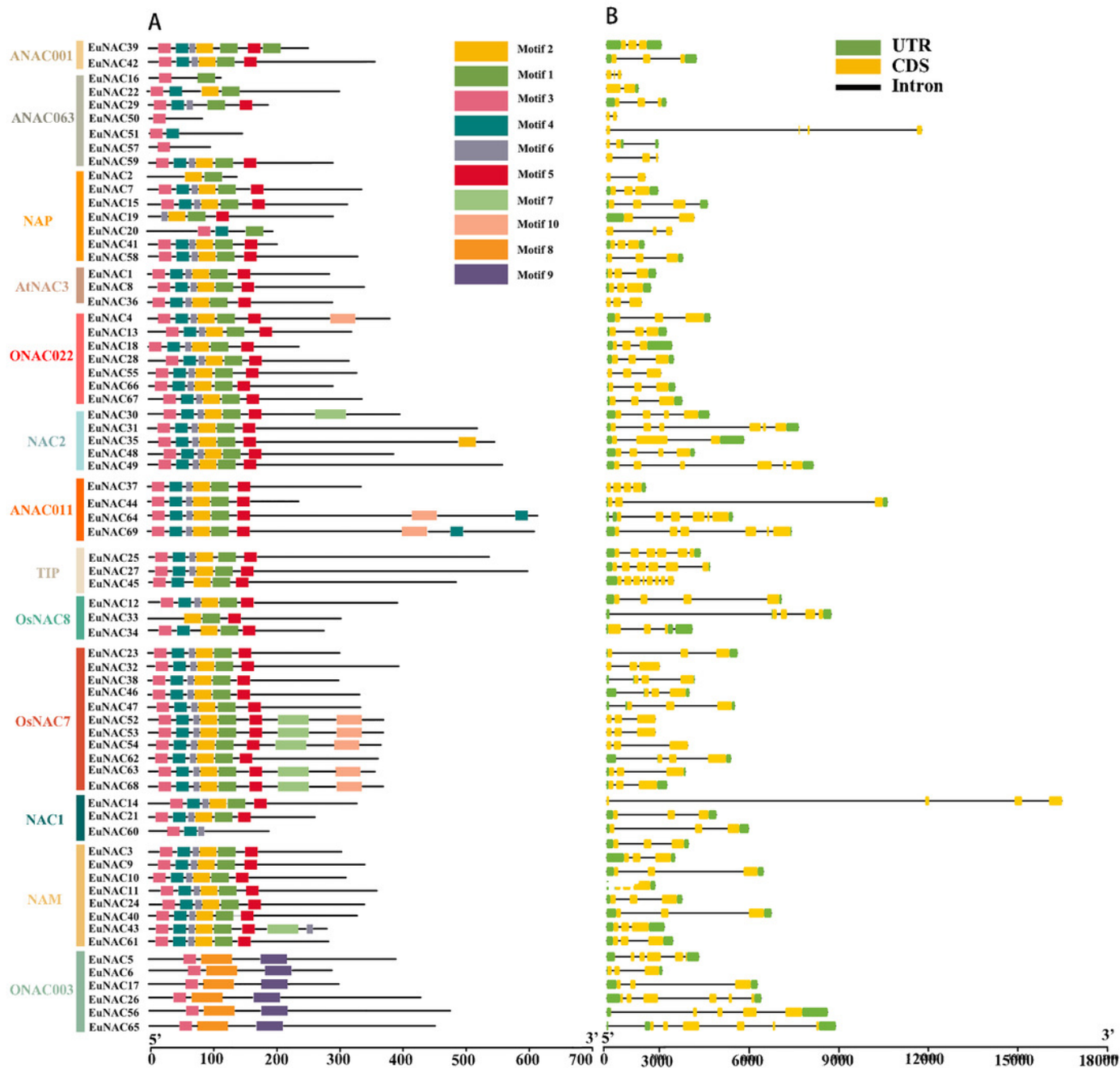


Figure 3

Distribution of 69 *EuNACs* on 16 chromosomes and two scaffolds.

Vertical bars represent the chromosomes of *E. ulmoides*. The chromosome number is on the left of each chromosome. The scale on the left represents the length of the chromosome.

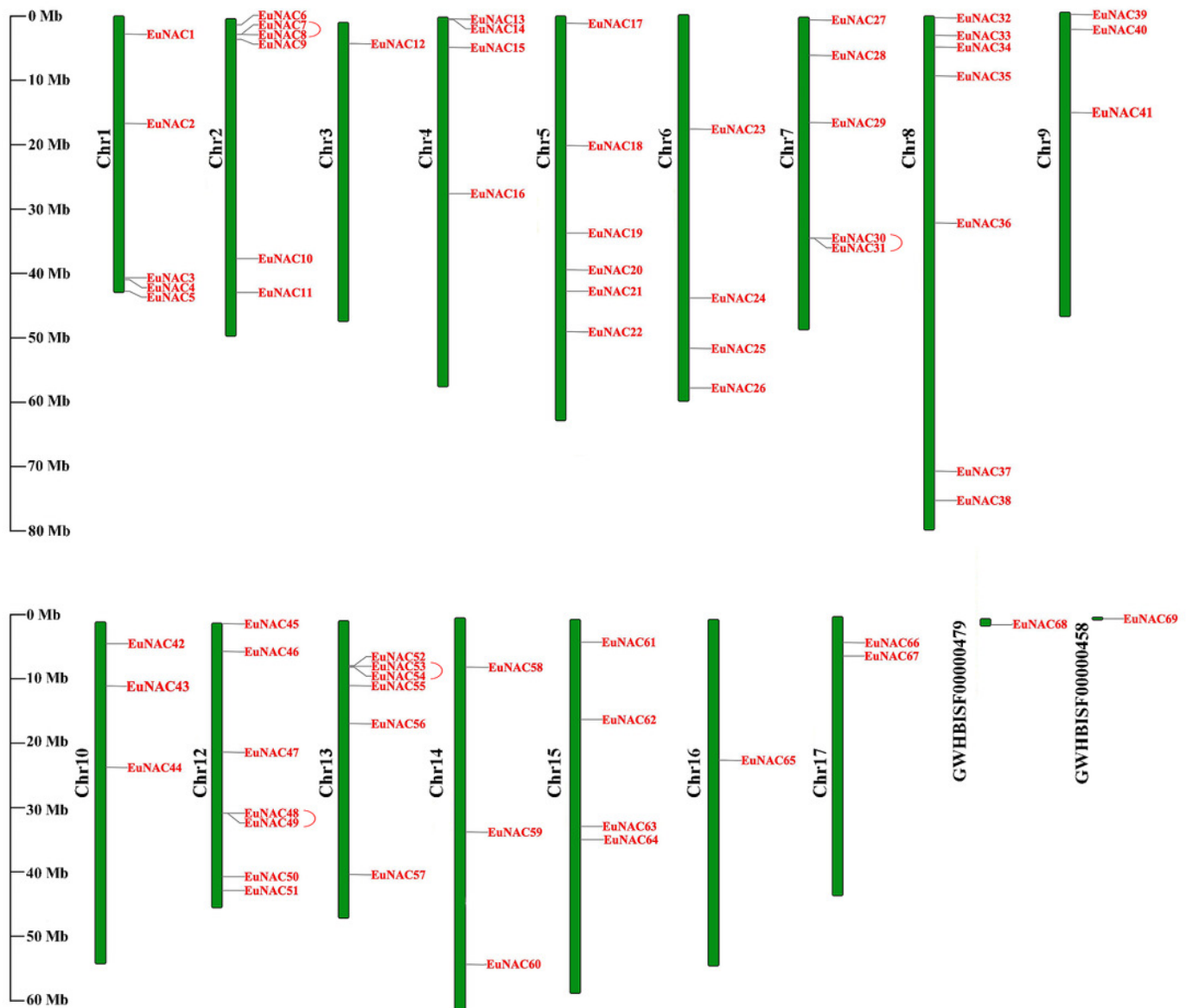


Figure 4

Schematic representations of the interchromosomal relationships of *EuNAC* genes.

The deep red line represents the *EuNAC* gene pairs replicated in tandem, while the remaining colored lines represent the *EuNAC* gene pairs replicated in segments.

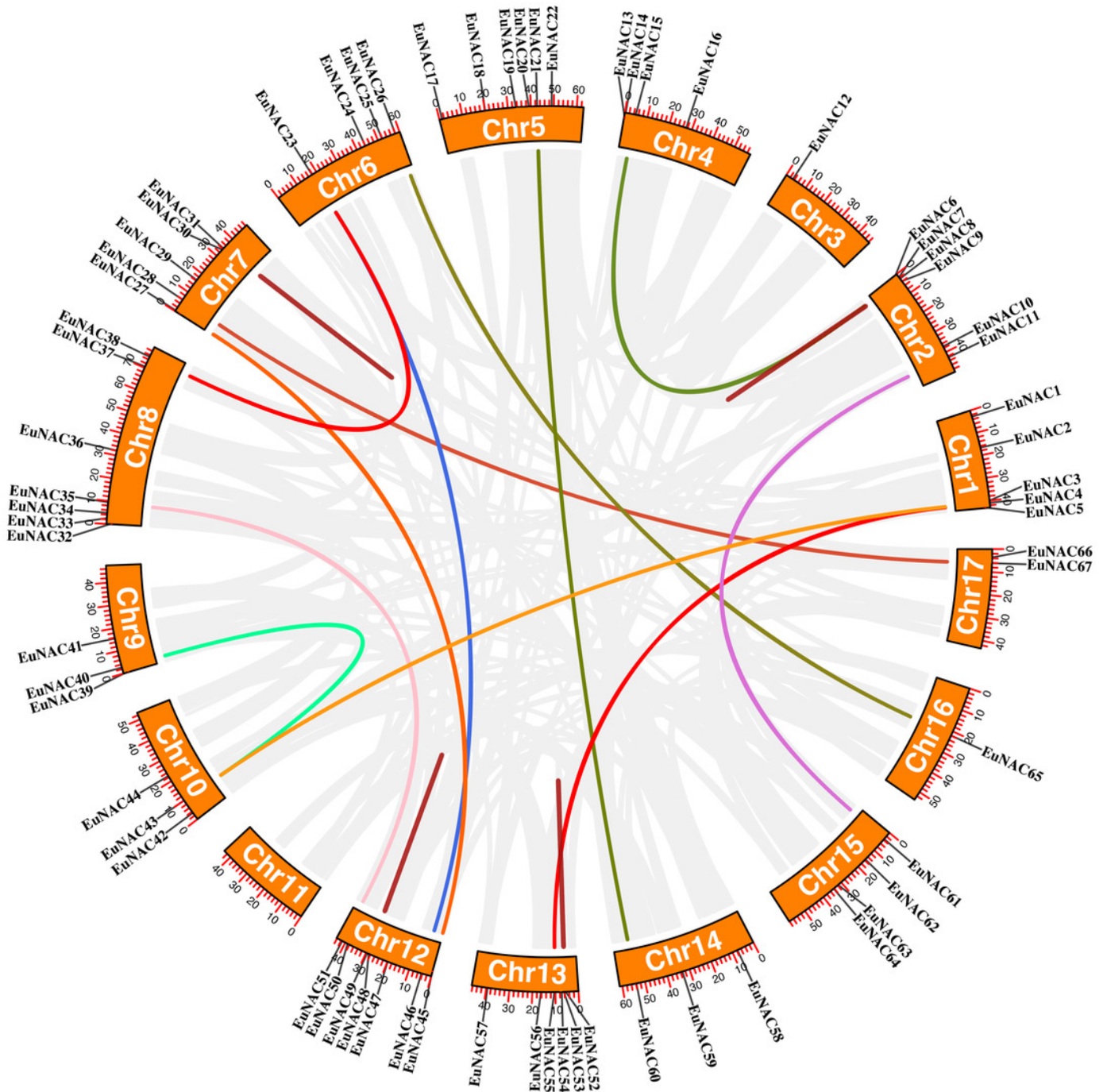


Figure 5

Synteny analysis of *NAC* genes between *E. ulmoides* and two representative plant species (*Arabidopsis thaliana* and *Oryza sativa*)

Green and purple lines represent syntenic *NAC* gene pairs of *E. ulmoides* and *A. thaliana* and *O. sativa*, respectively.

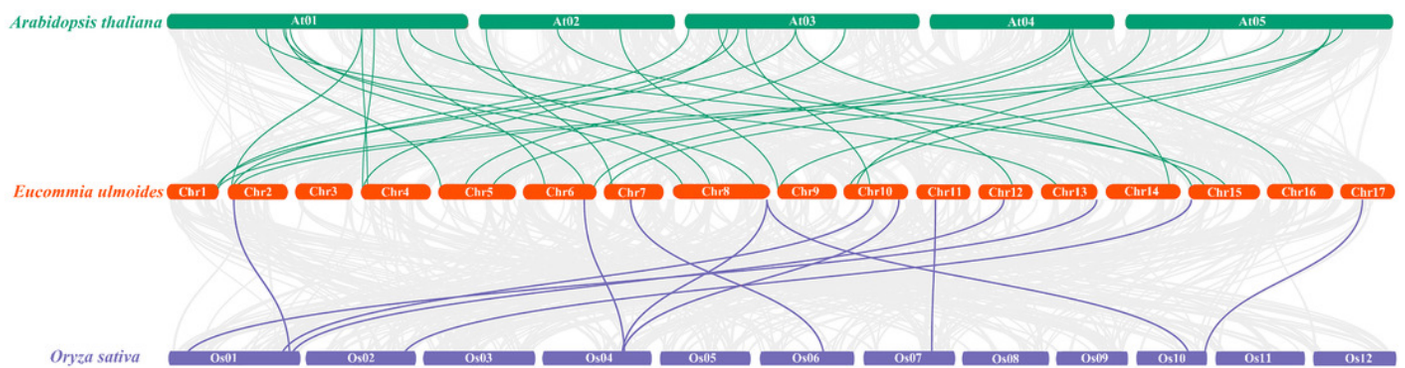


Figure 6

The number of each type of cis-acting element in the promoter region of each *EuNAC* gene.

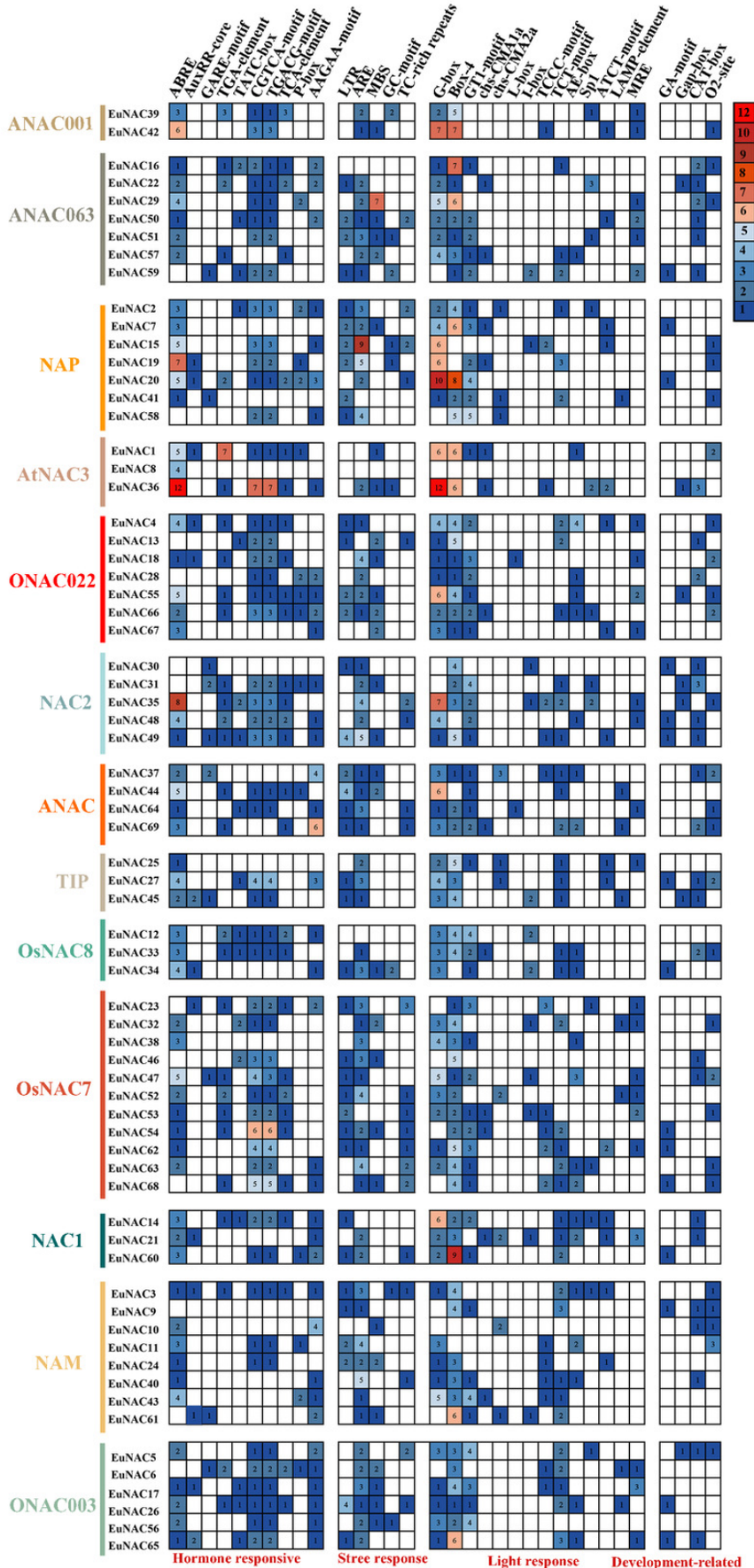


Figure 7

Expression levels of 69 *NAC* genes under drought stress in *E. ulmoides* of leaves

The expression level was presented based on the transformed data of $\log_2(\text{FPKM}+1)$ values.

