

Genome-wide identification of the *SWEET* gene family mediating the cold stress response in *Prunus mume*

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The *SWEET* (Sugars Will Eventually be Exported Transporter) gene family encodes a family of sugar transporters that plays essential roles in plant growth, reproduction, and biotic and abiotic stresses. *Prunus mume* is a considerable ornamental wood plant with high edible and medicinal values; however, low temperature severely limits its geographical distribution. To investigate whether this gene family mediates *P. mume*'s response to cold stress, we identified its 17 *SWEET* genes from *P. mume* and divided them members into four groups. Sixteen of these genes were anchored on six chromosomes, and one gene was anchored on the scaffold with four pairs of segmental gene duplications and two pairs of tandem gene duplications. *Cis*-acting regulatory element analysis indicated that the *PmSWEET* genes are potentially involved in the *P. mume* developmental procedure, such as circadian control, abscisic acid-response and light-response, and responses to numerous stresses, such as low-temperature and drought. We performed low-temperature treatment in the cold-tolerant cultivar 'Songchun' and cold-sensitive cultivar 'Zaolve' and found that the expression of four of 17 *PmSWEETs* was either upregulated or downregulated with prolonged treatment times, which indicates that these family members may potentially play a role in cold stress responses in *P. mume*. Our study provides a basis for further investigation of the role of *SWEET* proteins in the development of *P. mume* and its responses to cold stress.

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13

14 Abstract

15 The SWEET (Sugars Will Eventually be Exported Transporter) gene family encodes a family of
16 sugar transporters that plays essential roles in plant growth, reproduction, and biotic and abiotic
17 stresses. *Prunus mume* is a considerable ornamental wood plant with high edible and medicinal
18 values; however, low temperature severely limits its geographical distribution. To investigate
19 whether this gene family mediates *P. mume*'s response to cold stress, we identified its 17 *SWEET*
20 genes from *P. mume* and divided them members into four groups. Sixteen of these genes were
21 anchored on six chromosomes, and one gene was anchored on the scaffold with four pairs of
22 segmental gene duplications and two pairs of tandem gene duplications. *Cis*-acting regulatory
23 element analysis indicated that the *PmSWEET* genes are potentially involved in the *P. mume*
24 developmental procedure, such as circadian control, abscisic acid-response and light-response,
25 and responses to numerous stresses, such as low-temperature and drought. We performed low-
26 temperature treatment in the cold-tolerant cultivar 'Songchun' and cold-sensitive cultivar
27 'Zaolve' and found that the expression of four of 17 *PmSWEETs* was either upregulated or
28 downregulated with prolonged treatment times, which indicates that these family members may
29 potentially play a role in cold stress responses in *P. mume*. Our study provides a basis for further
30 investigation of the role of *SWEET* proteins in the development of *P. mume* and its responses to
31 cold stress.

32 **Keywords:** *Prunus mume*, SWEET gene family, expression pattern, cold response.

33

34 1. Introduction

35 Sucrose is the main carbohydrate in most plants; it is synthesized in the leaves during
36 photosynthesis and then transported by phloem sap to storage organs, such as roots, stems,
37 flowers, seeds and fruits (Rennie and Turgeon, 2009; Lemoine et al., 2013). Sucrose provides
38 energy and carbon sources for plants and acts as an important signal and resistance molecule that
39 participates in the normal growth of higher plants (Chen et al., 2015). However, these sugars
40 must be assisted by appropriate sugar transporters as it cannot be transported independently to
41 the storage organs (Ainsworth and Bush, 2011). At present, three transporter families have been
42 identified as essential sugar transporters: monosaccharide transporters (MSTs), sucrose
43 transporters (SUTs), and Sugar Will Eventually be Exported transporters (SWEETs) (Chen et al.,
44 2010; Chen et al., 2015; Eom et al., 2015). Of these three families, *SWEETs* were the final gene
45 family to be uncovered and were first identified by Chen et al. in *Arabidopsis* (Chen et al., 2010).
46 *SWEET* proteins act as sugar transporters that mediate the inflow or outflow of phloem
47 parenchyma sugar into the phloem apoplast (Slewiniski, 2011; Braun, 2012; Chen, 2014). Unlike
48 the SUT and MST families, which require energy to transport sugar across the plasma membrane

49 (Maynard and Lucas, 1982; Lemoine, 2000), SWEET proteins promote the diffusion of sugar
50 across concentration gradients at the cellular membrane or vacuolar membrane, regardless of the
51 proton gradient or pH of the cellular environment (Chen et al., 2012; Chen et al., 2015).

52 SWEET proteins are characterized by conserved MtN3_saliva (MtN3_slv) transmembrane
53 (TM) domains (Chen et al., 2012), also known as PQ-loop repeats (Eom et al., 2015; Feng and
54 Frommer, 2015). SWEETs in eukaryotes commonly consist of seven transmembrane helices
55 (TMHs), which contain a pair of 3-TMH repeats detached by an added helix (Xuan et al., 2013),
56 and this structure has been described as the “3-1-3” TM SWEET structure (Chen et al., 2010). In
57 contrast to the structure of eukaryote SWEET proteins, prokaryote SWEET proteins, known as
58 SemiSWEETs, are composed of only three TMHs (Xuan et al., 2013). In eukaryotes, proteins
59 that contain 6 or 7 TMHs are prevalent, but SemiSWEETs with 3 or 4 TMHs have also been
60 detected in plant genomes. In a study of SWEET genes from 25 plant genomes, 140 of the 411
61 SWEET sugar transporters identified were semiSWEET; with all of the identified semiSWEETs
62 either lacking the first or second 3-TM domain or exist only in partial form (Patil et al., 2015).
63 This data therefore demonstrates that the presence of semiSWEETs in higher plant genomes is
64 not unusual, and further, that SWEETs may in actual fact have formed by direct fusion from
65 SemiSWEETs (Jia et al., 2017). In addition, a novel extraSWEET protein consisting of 14 and
66 15 TMHs has been reported from *Vitis vinifera* (Patil et al., 2015) and *Oryza punctata* (Jia et al.,
67 2017); it is speculated that this extraSWEET may have formed from the duplication of a 7 TMH
68 SWEET gene in these two species. Recent research on 3, 249 SWEET proteins also identified a
69 superSWEET with > 18 TMHs in oomycetes, which carry 5–8 repeats of a semiSWEET (Jia et
70 al., 2017). According to phylogenetic analysis, the SWEET genes in *Arabidopsis* can be divided
71 into four clades: Clade I (SWEET1–3) and Clade II (SWEET4–8) mainly transport glucose,
72 while Clade I also transports hexose (Chen et al., 2010; Lin et al., 2014). Clade III members
73 (SWEET9–15) mainly transports sucrose (Chen et al., 2012; Eom et al., 2015), and Clade IV
74 members (SWEET16–17), which are located on the tonoplast membrane, mainly transports
75 fructose (Eom et al., 2015). The phylogenetic relationships of the SWEET genes described
76 hereafter are all based on results from *Arabidopsis*.

77 Advances in whole-genome sequencing have enabled genome-wide identification of
78 SWEET genes in numerous species. These include important crops, fruits and vegetables, such
79 as rice (*Oryza sativa*) (Yuan and Wang, 2013), sorghum (*Sorghum bicolor*) (Mizuno et al.,
80 2016), soybean (*Glycine max*) (Patil et al., 2015), apple (*Malus domestica*) (Wei et al., 2014),
81 grape (*Vitis vinifera*) (Chong et al., 2014), banana (*Musa acuminata*) (Miao et al., 2017), tomato
82 (*Solanum lycopersicum*) (Feng et al., 2015), rapeseed (*Brassica napus*) (Jian et al., 2016), potato
83 (*Solanum tuberosum*) (Li et al., 2020) and valencia sweet orange (*Citrus sinensis*) (Yao et al.,
84 2021). Additionally, many SWEET genes have been confirmed to play diverse and complex
85 roles in physiological processes, such as nectar secretion (Ge et al., 2000; Lin et al., 2014),

86 pollen development (Sun et al., 2013), senescence (Quirino et al., 1999), and seed filling (Sosso
87 et al., 2015). Moreover, SWEET genes are also involved in biotic and abiotic stress responses
88 (Yuan and Wang, 2013), including the reaction of plants to stress at low temperatures. For
89 example, overexpression of *AtSWEET16* and *AtSWEET17* increases cold tolerance (Chardon et
90 al., 2013; Klemens et al., 2013; Guo et al., 2014); overexpression of *AtSWEET4* increases plant
91 size and frost resistance (Chong et al., 2014; Liu et al., 2016); and *AtSWEET11* and *AtSWEET12*
92 are involved in responses to stress caused by cold or dehydration (Le Hir et al., 2015; Durand et
93 al., 2016). *AtSWEET15* is also known as SAG29 (where SAG stands for senescence-associated
94 gene); however, its transcription level gradually increases at low temperature, high salinity, and
95 drought during natural leaf senescence (Quirino et al., 1999). Cold stress significantly inhibits the
96 expression of *CsSWEET2*, *CsSWEET3*, and *CsSWEET16* in *Camellia sinensis*, while the
97 expression of *CsSWEET1* and *CsSWEET17* increases sharply (Yue et al., 2015). A functional
98 study of *CsSWEET16* in *C. sinensis* revealed that it is located in the vacuolar membrane and
99 regulates cold resistance in transgenic *Arabidopsis* plants (Wang et al., 2018). The transcriptional
100 activity of many *SISWEET* genes increases under low-temperature stress in tomato (Feng et al.,
101 2015). Studies have shown that expression of the *MaSWEET* gene in banana is upregulated in
102 response to low temperature, salt, and osmotic stress (Miao et al., 2017). Using genome-wide
103 analysis of the *BoSWEET* gene in *Brassica oleracea* var. *capitata*, five possible candidate genes
104 were found to promote sugar transport and thereby enhance chilling tolerance in cabbage (Zhang
105 et al., 2019).

106 *Prunus mume* is a traditional flower native to southwest China and the middle and lower
107 reaches of the Yangtze River. In the northern China, low temperatures severely limit the growth
108 and distribution of this species. Although SWEET sugar transporters have been associated with
109 responses to cold in other species, little is known about the role of *PmSWEETs* in cold responses
110 in *P. mume*. This study aims to conduct a genome-wide analysis of the SWEET gene family in *P.*
111 *mume*, with a specific focus on SWEET gene transcriptional responses to cold stress, providing a
112 starting point to study the detailed role of *PmSWEETs*.

113 2. Materials and Methods

114 2.1 Plant Genomic Resources

115 To explore the phylogeny of the SWEET genes in *P. mume* and other species, we downloaded
116 SWEET proteins from two model plants (*Arabidopsis thaliana* and *Oryza sativa*, representing
117 dicotyledons and monocotyledons, respectively) and eight other Rosaceae species. The protein
118 sequences of 17 *AtSWEETs* and 21 *OsSWEETs* were downloaded from the TAIR 10 database
119 (<http://www.arabidopsis.org/>) and TIGR (<http://rice.plantbiology.msu.edu/>), respectively. The *P.*
120 *mume* genome sequence and annotation files were obtained from the *P. mume* genome project
121 (<http://prunusmumegenome.bjfu.edu.cn/>); the genomes of eight other Rosaceae species, *Malus*

122 *domestica* (Daccord et al., 2017), *Prunus avium* (Shirasawa et al., 2017), *Prunus persica* (Verde
123 et al., 2013), *Prunus yedoensis* (Baek et al., 2018), *Pyrus communis* (Linsmith et al., 2019), *Rosa*
124 *chinensis* (Raymond et al., 2018), *Prunus salicina* (Liu et al., 2020), and *Prunus armeniaca* (Jiang
125 et al., 2019), were downloaded from the Genome Database for Rosaceae ([https://www.
126 rosaceae.org/](https://www.rosaceae.org/)).

127 **2.2 Identification of SWEET Genes in *P. mume* and Other Species**

128 The hidden Markov model (HMM) profiles of the MtN3_slv domain for the SWEET gene
129 family (PF03083) were downloaded from the Pfam database (<http://pfam.xfam.org/>) and used as
130 queries to search for SWEET proteins in the proteomes of *P. mume* and other species with
131 HMMER software (version 3.1b2, <http://hmmer.org/>) (Finn et al., 2015). To ensure confidence,
132 the E-value cutoff was set at 10^{-5} . Then, all putative SWEET proteins were screened to confirm
133 the presence of the MtN3_slv domain by SMART (<http://smart.embl-heidelberg.de/>), the Pfam
134 database (<http://pfam.xfam.org/>) and NCBI-CDD (<https://www.ncbi.nlm.nih.gov/cdd>), and
135 sequences with MtN3_slv domain were retained.

136 The SWEET genes were named based on their location information in the genome. In
137 addition, the number of amino acids, molecular weight (MW) and isoelectric point (pi) were
138 calculated using the online ExpASy program ([https://web.expasy.org/cgi-
139 bin/protparam/protparam](https://web.expasy.org/cgi-bin/protparam/protparam)). The distributions of TM helices were predicted by TMHMM Server
140 v. 2.0 (<http://www.cbs.dtu.dk/services/TMHMM/>).

141 **2.3 Phylogenetic and Conserved Domain Analysis**

142 To examine the phylogeny between SWEET genes in *P. mume* and other species, alignment
143 of full-length SWEET protein sequences from three species (*P. mume*, *A. thaliana*, and *O.*
144 *sativa*) and eight Rosaceae species was performed by using MAFFT software with the FFT-NS-1
145 strategy (Kato and Standley, 2013). Subsequently, maximum likelihood (ML) phylogenetic
146 trees were constructed using FastTree (version 2.1.11) (Price et al., 2010) with default
147 parameters. Then, iTOL v4.0 (<https://itol.embl.de/itol.cgi>) (Letunic and Bork, 2019) and AI CS6
148 software were used to annotate and embellish the phylogenetic tree.

149 **2.4 Conserved Motif and Gene Structure Analysis**

150 The conserved motifs of *PmSWEETs* were predicted by MEME Suite Version 5.3.3
151 (<https://meme-suite.org/meme/tools/meme>) (Bailey et al., 2009), where the maximum number of
152 motifs for the conserved domains was set to 10, motif width was set to 6-50, and the residuals
153 were designated as the default parameters. Gene structure data was extracted from the *P. mume*
154 genome gff file, visualized using TBtools software (Chen et al., 2020), and then edited in AI CS6
155 software.

156 **2.5 Chromosome Location, Duplication and Synteny Analysis**

157 The location and chromosome length information of *PmSWEETs* was obtained from the gff
158 file downloaded from the *P. mume* genome project (<http://prunusmumegenome.bjfu.edu.cn/>). A
159 chromosomal location figure was drawn using the online tool MG2C
160 (http://mg2c.iask.in/mg2c_v2.0/). Gene tandem and segment replication events were analyzed
161 using the Multiple Collinearity Scan Toolkit (MCScanX) and Circos in TBtools, respectively,
162 with the default parameters. The synteny of the *PmSWEETs* across *A. thaliana*, *P. armeniaca*,
163 and *P. salicina* was mapped using MCScanX in TBtools. The Ks and Ka values for duplicated
164 gene pairs were calculated based on the coding sequence alignments using the Ka/Ks calculator
165 in TBtools. According to two ordinary rates (λ) of 1.5×10^{-8} and 6.1×10^{-9} substitutions per site
166 per year (Lynch and Conery, 2000; Blanc and Wolfe, 2004), the formula $t = Ks/2\lambda \times 10^{-6}$ Mya
167 was used to calculate the divergence time.

168 **2.6 Cis-Acting Element Analysis of *PmSWEET* Gene Promoter Regions**

169 The upstream sequences (2.0 kb) of the *PmSWEETs* were retrieved from the genomic
170 sequence data in TBtools and then submitted to the PlantCARE database
171 (<http://bioinformatics.psb.ugent.be/webtools/plantcare/html/>) (Lescot et al., 2002) for *cis*-acting
172 element analysis. We finally selected 12 elements, including those induced by hormones, such as
173 methyl jasmonate (MeJA)-responsive, abscisic acid (ABA)-responsive, and stress-responsive
174 elements; the stress-responsive factors included those involved in defense and stress, low
175 temperature, and light. By combining these data with phylogenetic tree information (nwk file),
176 the map was constructed by TBtools and edited by AI CS6 software.

177 **2.7 *PmSWEET* Genes Expression Analysis**

178 To investigate the function of *PmSWEETs* involved in tissue development and cold
179 tolerance, we used root, stem, leaf, bud and fruit data from RNA sequencing (Zhang et al., 2012)
180 to analyze the *PmSWEET* expression patterns in different tissues and then used flower bud
181 dormancy data from RNA sequencing of *P. mume* ('Zaolve') (Zhang et al., 2018) to analyze
182 *PmSWEET* responses to low temperature from November to February. Furthermore, we explored
183 the expression of SWEET gene family members in the stem of *P. mume* ('Songchun') in
184 geographically distinct locations, including Beijing (BJ, N39°54', E116°28'), Chifeng (CF,
185 N42°17', E118°58') and Gongzhuling (ZGL, N43°42', E124°47') and for three different periods
186 of the year, including cold acclimation (October, autumn), the final period of endo-dormancy
187 (January, winter), and deacclimation (March, spring) (Jiang, 2020). TBtools (Chen et al., 2020)
188 was used to create the heatmap.

189 **2.8 qRT-PCR Analysis of *PmSWEET* Genes**

190 To examine the response of *PmSWEET* to low temperature, the annual branches of the cold-
191 sensitive cultivar ‘Zaolve’ and the cold-tolerant cultivar ‘Songchun’ were collected. Before
192 chilling treatment, the shoots were incubated overnight at 22 °C and then transferred to 4 °C for
193 0, 1, 4, 6, 12, 24, 48, and 72 h under long-day conditions (16-h light/8-h dark). The stems were
194 collected immediately stored in liquid nitrogen until their longterm storage at -80 °C in readiness
195 for RNA extraction. Each treatment had three biological replicates.

196 Total RNA of each sample was extracted using the RNAPrep Pure Plant Plus Kit (Tiangen,
197 Beijing, China). Complementary cDNA was synthesized using ReverTra Ace® qPCR RT Master
198 Mix with gDNA Remover (Toyobo, Osaka, Japan). The specific primers were designed by
199 Primer 3 (<https://bioinfo.ut.ee/primer3-0.4.0/>) based on the cDNA sequences (Table S1). The
200 expression levels of *PmSWEETs* at low temperature were analyzed using quantitative real-time
201 polymerase chain reaction (qRT-PCR) with a PikoReal real-time PCR system (Thermo Fisher
202 Scientific, CA, USA) with SYBR® Green Premix *Pro Taq* HS qPCR kit (Accurate biology,
203 China). The reactions were performed in a 10 µL volume, including 5.0 µL SYBR®Green
204 Premix *Pro Taq* HS qPCR master mix, 0.5 µL each of forward and reverse primers, 1.0 µL of
205 cDNA and 3.0 µL of ddH₂O. The reactions were performed according to the following
206 procedure: 95 °C for 30 s, followed by 40 cycles of 95 °C for 5 s and 60 °C for 30 s. Via the use
207 of the phosphatase 2A gene of *P. mume* as the reference gene, the relative expression was
208 calculated by using the formula $2^{-\Delta\Delta C_t}$ method (Livak and Schmittgen, 2001). Each real-time
209 qRT-PCR was conducted in three biological replicates. The statistical analyses of ‘Zaolve’ and
210 ‘Songchun’ were independent carried out using SPSS22.0, the one-way ANOVA analysis of
211 variance was calculated by least significant difference (LSD) and Student-Newman-Keuls test
212 with significant difference at level $p = 0.05$. GraphPad Prism6 software was used to draw the
213 diagram.

214 3. Results

215 3.1 Identification of Members of the *Prunus mume* SWEET Gene Family

216 A total of 17 nonredundant *PmSWEETs* were detected in the *P. mume* genome (sequence
217 information is shown in Supplement File S1), and 175 SWEETs were detected in the eight other
218 species of Rosaceae, including 16 *SWEET* genes in *P. armeniaca*, 19 in *P. avium*, 19 in *P.*
219 *persica*, 19 in *P. salicina*, 16 in *P. yedoensis*, 21 in *P. communis*, 29 in *M. domestica*, and 36 in
220 *R. chinensis* with rigorous filtering. All the newly identified SWEET genes were named
221 according to their chromosome location (Table 1 and Table S2). We determined that candidates
222 with at least one MtN3_slv domain were “genuine” *SWEETs*, all *SWEETs* contained MtN3_slv
223 domains (domain architecture of *PmSWEETs* is shown in Supplement File S2). The number of
224 amino acids, molecular weight (MW), and isoelectric point (pI) were calculated on the basis of
225 the protein sequences. As exhibited in Table 1, the predicted *PmSWEET* proteins ranged from

226 105 (*PmSWEET14*) to 580 (*PmSWEET8*) amino acids in length, with relative molecular weights
227 ranging from 15.96 kDa (*PmSWEET11*) to 63.43 kDa (*PmSWEET8*), and theoretical pIs ranging
228 from 8.30 (*PmSWEET4*) to 9.76 (*PmSWEET3*). The MW and pI of family member *PmSWEET14*
229 could not be determined using this approach however due to the presence of four consecutive
230 undefined amino acids (Table 1). Through prediction and analysis of TMHs of the 17 identified
231 *PmSWEETs*, we found that these *PmSWEET* proteins were predicted to have 2–7 TMHs, and
232 seven members of the *P. mume* SWEET gene family possess 7 TMHs, rarely, there may be only
233 three or two TMHs. Detailed location information of the TMHs is shown in Table S3 and Figure
234 S1.

235 3.2 Phylogenetic Analysis and Classification of *SWEET* Genes

236 To better understand the evolution of homologous *SWEET* genes, we used the ML method
237 to create a phylogenetic tree of all *SWEET* sequences from *A. thaliana* (model dicots), *O. sativa*
238 (model monocots), and *P. mume*. According to previously reported *AtSWEETs* and *OsSWEETs*
239 (Chen et al., 2010; Yuan and Wang, 2013), the 17 identified *PmSWEETs* were divided into four
240 clades (i.e., Clade I, Clade II, Clade III, and Clade IV) (Figure S2). To investigate the
241 evolutionary relationships between *PmSWEETs* and the *SWEETs* of other species, an ML
242 phylogenetic tree of *SWEETs* from 11 species, including 8 other Rosaceae species, was
243 constructed. All members of the *SWEET* gene family in the 11 species were divided into four
244 clades (Figure 1). The largest clade was Clade III, which comprised five *OsSWEET* genes, seven
245 *AtSWEET* genes, and 68 Rosaceae *SWEET* genes; the specific number of genes is shown in Table
246 S4. The smallest clade was Clade IV, which consisted of only two *A. thaliana* *SWEET* genes, one
247 *O. sativa* gene, and 18 Rosaceae *SWEET* genes (Table S4), indicating that *SWEETs* were
248 distributed unevenly among the clades. The numbers of genes in Clade I, II and III varied
249 greatly, suggesting that the *SWEET* gene family expanded, especially in Clades I, II and III,
250 during Rosaceae evolution. The *SWEETs* of Rosaceae were distributed uniformly across each
251 small clade, whereas *SWEETs* from *O. sativa* tended to cluster together. The *PmSWEETs*,
252 *PpSWEETs*, and *PavSWEETs* were clustered together and had similar distributions in the
253 phylogenetic tree.

254 3.3 Conserved Motif and Gene Structure Analysis

255 To explore the sequence features of *PmSWEET* proteins, MEME software and TBtools were
256 used to predict and draw conserved domains. As a consequence, ten distinct motifs were detected
257 in *SWEET* proteins (Figure 2B), and a schematic diagram of *PmSWEET* protein motifs is shown
258 in Figure S3. The number of *PmSWEETs* motifs was distinctive, ranging from 1 to 7. Of them,
259 12 *PmSWEETs* contained more than four motifs, 4 *PmSWEETs* harbored four motifs, and
260 *PmSWEET14* contained only one motifs. Motifs 1, 2, 3, 4 and 6 were highly conserved and
261 present in 15 *PmSWEET*, 13 *PmSWEET*, 16 *PmSWEET*, 11 *PmSWEET* and 12 *PmSWEET*

262 proteins, respectively; while motifs 7, 8 and 10 were relatively unique and existed in only 4
263 PmSWEET, 2 PmSWEET and 2 PmSWEET proteins, respectively. Intriguingly, aside from
264 some unusual proteins, *SWEET* members of the same clade had similar conserved motifs,
265 suggesting that they might have similar functions.

266 To elucidate the structural characteristics of the *PmSWEETs*, the exon-intron structure was
267 further analyzed. As shown in Figure 2C, *PmSWEETs* in Clade II (except *PmSWEET10*)
268 contained four introns. *PmSWEET1*, *PmSWEET9*, and *PmSWEET15* in Clade III had five introns,
269 *PmSWEET8* contained the largest number of introns (12 introns), while *PmSWEET14* contained
270 only one intron. All *PmSWEETs* in Clade IV had five introns. The number of introns in Clade I
271 varied from just two to ten, *PmSWEET17* had two introns, *PmSWEET4* contained five introns,
272 *PmSWEET11* and *PmSWEET12* contained three introns, *PmSWEET3* had ten introns. These
273 results indicated that aside from some unusual proteins, genes clustered together generally
274 exhibited similar gene structures.

275 **3.4 Chromosomal Distribution and Tandem Duplication of *PmSWEET* gene family** 276 **members**

277 According to gene location information, all 17 *PmSWEETs* were mapped, showing that 16
278 *PmSWEETs* were located on chromosomes, and one *PmSWEET* gene was located on scaffold54
279 (Figure 3). The *PmSWEETs* on chromosomes 6 and 7 were clustered in the center of each
280 chromosomes, and all contained four *PmSWEETs*. Two genes each were distributed on
281 chromosomes 2, 3, 4 and 5. *PmSWEET11* and *PmSWEET12* and then *PmSWEET14* and
282 *PmSWEET15* were clustered into two tandem duplication events on chromosomes 6 and 7,
283 respectively. Based on the above results, some *PmSWEETs* gene family members were
284 putatively generated by gene tandem duplication.

285 **3.5 Segmental Duplication and Synteny of the *PmSWEET* Gene Family**

286 Synteny analysis of *PmSWEETs* was performed using the Circos program of TBtools, four
287 segmental duplication events, including *PmSWEET1/PmSWEET14*, *PmSWEET5/PmSWEET8*,
288 *PmSWEET6/PmSWEET9* and *PmSWEET6/PmSWEET16* were detected, and further, each gene
289 pair was located on a different chromosome, as shown with red lines in Figure 4. This finding
290 strongly suggests that some *PmSWEETs* were probability generated by gene segmental
291 duplication. In addition, the selection pressure and divergence time of the duplication events
292 were estimated by the Ka (nonsynonymous) and Ks (synonymous) substitution ratio. In the
293 evolutionary process, the Ka/Ks ratio > 1 indicates positive selection (adaptive evolution), a ratio
294 = 1 indicates neutral evolution (drift), and a ratio < 1 indicates negative selection (conservation).
295 Only one pair of segmentally duplicated *PmSWEETs* (*PmSWEET6/9*) had a Ka/Ks ratio of 0.45,
296 which was significant, and indicated a synonymous change that has been selected during plant

297 genome evolution. The differentiation period of the *PmSWEET6/9* gene pair was 55.34~136.07
298 Mya.

299 To further examine the specific retention of *PmSWEETs*, their collinearity relationship with
300 *AtSWEETs*, *PaSWEETs*, and *PsSWEETs* were detected using the MCScanX procedure of
301 TBtools. A total of 16 homologous gene pairs were detected in *P. mume* and *A. thaliana*.
302 Similarly, 16 pairs of homologous genes between *P. mume* and *P. armeniaca* and 20 between *P.*
303 *mume* and *P. salicina* were detected (Figure 5, Table S5). The collinear complexity of *P. mume*
304 with *P. salicina* was much higher than that with *P. armeniaca* and *A. thaliana*. These results
305 suggested that *P. mume* was relatively distantly related to *A. thaliana* and *P. armeniaca*, but is
306 more closely related to *P. salicina*.

307 **3.6 Prediction Analysis of Cis-Acting Elements within *PmSWEETs* gene promoters**

308 To further investigate the possible regulatory mechanism of *PmSWEETs* in the process of
309 growth or in plant defence mechanisms, in particular the response to abiotic stress, such as low
310 temperature, we submitted the 2.0 kb upstream sequence from the translation start site of each
311 *PmSWEET* gene to the PlantCARE database to search for the presence of specific *cis*-elements.
312 The *PmSWEET* promoters comprised several conserved regulatory elements that respond to plant
313 hormones and environmental stress, and twelve of these were analyzed further (Figure 6, Table
314 S6). Elements related to light response, anaerobic induction, and ABA response were widespread
315 in the promoter areas of 17, 17 and 16 members of the *P. mume* SWEET gene family,
316 respectively. According to the regulatory elements in their promoters, 14, 12, 11, 10, and 9 *P.*
317 *mume* SWEET gene family members were sensitive to drought inducibility, MeJA, gibberellin,
318 low temperatures and auxin, respectively. By combining these findings with the results of
319 phylogenetic analysis, it was found that gene members of the same clade had similar *cis*-
320 elements. These results indicated that *PmSWEET* genes were involved in the regulatory
321 mechanisms of various stress responses.

322 **3.7 Expression Pattern Analysis of *PmSWEETs***

323 To investigate the role of *PmSWEETs* in development and response to low temperature,
324 the expression patterns of family members in the roots, stems, leaves, buds, fruits and flower
325 buds of different stages of dormancy, were examined based on the RNA-seq dataset (Jiang,
326 2020), and their RPKM values are shown in Tables S7 and S8. As illustrated in Figure 7A, 14 of
327 the *PmSWEET* genes were expressed in at least one tissue, whereas RNA-seq failed to detect the
328 expression of three family members (*PmSWEET5*, *PmSWEET 10* and *PmSWEET 11*). Among
329 them, five *PmSWEETs* presented relatively higher expression levels in fruits (*PmSWEET1*,
330 *PmSWEET6*, *PmSWEET9*, *PmSWEET12* and *PmSWEET17*) and buds (*PmSWEET3*,
331 *PmSWEET13*, *PmSWEET14*, *PmSWEET15* and *PmSWEET16*). Two *PmSWEETs* showed higher

332 expression levels in roots (*PmSWEET4* and *PmSWEET7*) and stems (*PmSWEET2* and
333 *PmSWEET8*). Additionally, several genes (*PmSWEET2*, *PmSWEET3*, *PmSWEET4*, *PmSWEET7*,
334 *PmSWEET8*, *PmSWEET12* and *PmSWEET13*) were expressed in leaves, but their expression
335 levels were low.

336 Most *PmSWEETs* were expressed during the bud dormancy period (except *PmSWEET5* and
337 *PmSWEET16*) as well as being expressed at specific stages of development (Figure 7B). Ten
338 *PmSWEET* genes exhibited specifically higher expressions in the Natural flush (NF) stage
339 (February), *PmSWEET9* was preferentially expressed in the Endo-dormancy I (EDI) stage
340 (November), *PmSWEET10* and *PmSWEET12* showed the highest level of expression in the
341 Endo-dormancy II (EDII) stage (December); and *PmSWEET1*, *PmSWEET3*, *PmSWEET6*,
342 *PmSWEET12* and *PmSWEET13* showed upregulated expression in the Endo-dormancy III
343 (EDIII) stage (January). Among these upregulated genes, eight *PmSWEETs* (*PmSWEET6*,
344 *PmSWEET7*, *PmSWEET10*, *PmSWEET11*, *PmSWEET13*, *PmSWEET14*, *PmSWEET15* and
345 *PmSWEET17*) (Table S6) contained low temperature response elements within their analyzed
346 promoter regions.

347 To further investigate the expression patterns of *PmSWEETs* under cold exposure, we
348 analyzed the stems of the cold-tolerant cultivar *P. mume* ‘Songchun’ at three geographical
349 locations, and their FPKM values are displayed in Table S9. The expression of six *PmSWEET*
350 genes (*PmSWEET5*, *PmSWEET6*, *PmSWEET11*, *PmSWEET14*, *PmSWEET16* and *PmSWEET17*)
351 was not detected. Among the other 11 *PmSWEET* genes, seven *PmSWEETs* (*PmSWEET1*,
352 *PmSWEET2*, *PmSWEET3*, *PmSWEET4*, *PmSWEET7*, *PmSWEET8* and *PmSWEET9*) showed
353 higher expression in spring (3.2~5.3 °C). *PmSWEET13* expression was upregulated in autumn
354 (6.1~7.9 °C) and winter in Beijing (-5.4 °C) and Chifeng (-11.4 °C), but downregulated in
355 spring; the expression levels of *PmSWEET10*, *PmSWEET12* and *PmSWEET15* increased
356 significantly in winter in Beijing (-5.4 °C) (Figure 8A). Among these genes with upregulated
357 expression, four *PmSWEETs* (*PmSWEET7*, *PmSWEET10*, *PmSWEET13* and *PmSWEET15*)
358 (Table S6) contained low-temperature response elements within their analyzed promoter regions.
359 To compare the expression patterns of *PmSWEETs* during different times of the year, another
360 heatmap was generated (Figure 8B). As shown in Figure 8B, *PmSWEETs* expression in the
361 material sourced from the locations, Chifeng and Gongzhuling showed similar expression
362 patterns at the same time of the year, while *PmSWEETs* expressed for the material sourced from
363 the Beijing location showed higher expression in winter (Figure 8B). This may be related to the
364 latitude of the three places, Gongzhuling has the highest latitude, followed by Chifeng and
365 Beijing. There is little difference between the temperature in autumn and spring in these three
366 places, but there is a big difference in winter. In winter, the temperature in Beijing (-5.4 °C) is
367 higher than that in the other two places (Gongzhuling is -22.8 °C, Chifeng is -11.4 °C), which
368 may be the temperature that induces some *PmSWEET* gene expression.

369 3.8 Expression Patterns of *P. mume* SWEETs under Cold Treatment

370 To investigate the role of *PmSWEETs* in response to cold stress, the expression patterns
371 under imposed hypothermia (4 °C) (0, 1, 4, 6, 12, 24, 48 and 72 h) were examined by qRT–PCR
372 using the cold-sensitive cultivar ‘Zaolve’ and the cold-tolerant cultivar ‘Songchun’. We
373 performed a qRT–PCR assay on the 17 identified *P. mume* SWEETs, but the expression of only
374 11 *PmSWEETs* was detectable by this approach, while the remaining 6 *PmSWEETs*
375 (*PmSWEET5*, *PmSWEET6*, *PmSWEET9*, *PmSWEET11*, *PmSWEET15* and *PmSWEET16*) were
376 not detected, consistent with the transcriptome data (Figures 7, 8). As displayed in Figure 9, the
377 changes in expression levels of the 11 *SWEET* genes in the two cultivars differed during the
378 imposed cold stress treatment period. In two varieties, three genes (*PmSWEET2*, *PmSWEET7*
379 and *PmSWEET8*) could be induced to downregulated in both ‘Songchun’ and ‘Zaolve’. In
380 addition, the expression of *PmSWEET13* could be induced to upregulated in both ‘Songchun’
381 and ‘Zaolve’, which rose approximately 11-fold after 6 h of cold treatment in ‘Songchun’, while
382 rose approximately 9-fold after 1 h, and then increased nearly 80-fold after 72 h of cold
383 treatment in ‘Zaolve’. One gene (*PmSWEET3*) changed only slightly in both ‘Songchun’ and
384 ‘Zaolve’. Six genes (*PmSWEET1*, *PmSWEET4*, *PmSWEET10*, *PmSWEET12*, *PmSWEET14*, and
385 *PmSWEET17*) exhibited different expression patterns in the two cultivars. Among those,
386 *PmSWEET1* and *PmSWEET12* were upregulated initially, then downregulated with increasing
387 treatment duration in ‘Songchun’, while in ‘Zaolve’, there was no obvious change in early stage,
388 but rapidly upgraded at 48 h and 72 h, respectively. *PmSWEET4* and *PmSWEET10* were
389 dramatically downregulated with increased cold stress duration in ‘Songchun’, while they were
390 upregulated within 6 h and then decreased with extended treatment in ‘Zaolve’. *PmSWEET14*
391 was no obvious change in early stage, but rapidly upregulated at 72 h in ‘Songchun’, while it was
392 rapidly upregulated at 24 h in ‘Zaolve’, and then downregulated with increasing treatment
393 duration. *PmSWEET17* was upregulated firstly, then downregulated with increasing treatment
394 duration in ‘Songchun’, while it was highly expressed only at 4 h in ‘Zaolve’.

395 4. Discussion

396 SWEET genes form a family of sugar transporters that play a role in the transportation of
397 sugars, mainly sucrose, glucose and fructose (Chen et al., 2010; Chen et al., 2012; Feng and
398 Frommer, 2015; Guo et al, 2014; Klemens et al., 2013; Le Hir et al., 2015), and participate in
399 diverse physiological and biological processes in the growth and development of many plants
400 and their responses to biotic and abiotic factors (Lemoine et al., 2013; Li et al., 2017; Li et al.,
401 2018; Zhao et al., 2018). Previous studies have shown that *SWEETs* participate in cold stress
402 responses in several plants (Chardon et al., 2013; Klemens et al., 2013; Guo et al., 2014; Chong
403 et al., 2014; Liu et al., 2016; Le Hir et al., 2015; Yue et al., 2015; Wang et al., 2018; Feng et al.,
404 2015; Miao et al., 2017; Zhang et al., 2019). However, little is known about the potential roles of

405 *P. mume* SWEET genes involved in cold stress. *P. mume* has a high ornamental value, and it can
406 blossom at lower temperatures; but different varieties have different cold resistance, making it a
407 very good material for studying the mechanisms of how *P. mume* SWEET genes function in cold
408 responses. Understanding the link between SWEET genes of *P. mume* and cold-resistance could
409 provide insights into cold-resistance molecular breeding in the future. In this research, we
410 detected a total of 17 *PmSWEETs* in *P. mume*, as many as in *Arabidopsis*, and similar to the
411 numbers in other species of *Prunus*, showing that SWEET genes are still relatively conserved in
412 *Prunus*. The length of *PmSWEET* proteins ranges from 105 aa to 580 aa, and this range provides
413 diversity in the number of TMHs (2–7). *PmSWEETs*, except for *PmSWEET14*, have a theoretical
414 pI larger than 8.0. As an important parameter of proteins, pI is determined by the relative
415 contents of amino acid residues at different pH values, which affects the stability, activity and
416 function of proteins (Gasteiger, 2005). The pI of *PmSWEET14* was not detected, which may be
417 due to its short amino acid sequence.

418 By predicting TMH domains, we found that the number of TMHs in *PmSWEET* genes
419 ranged from 2 to 7, certain *P. mume* SWEETs with only two, three, four, five or six TMHs
420 (Table 1). Fewer than seven TMHs in the eukaryotic SWEET family were also found in other
421 plants, such as wheat (Gao et al., 2018; Gautam et al., 2019), walnut (Jiang et al., 2020),
422 *Kentucky bluegrass* (Zhang et al., 2020) and soybean (Patil et al., 2015). To further validate the
423 accuracy of the SWEET protein, we submitted the protein sequence to the NCBI-CDD and
424 SMART online tools to predict its conserved domains, and it was found that each assessed
425 family member contained the MtN3_slv domain, and therefore, belonged to the SWEET family.
426 The results means that duplication and fusion or genetic loss might take place in the *P. mume*
427 genome. Similar to the case in other plants (Chen et al., 2010; Yuan and Wang, 2013; Patil et al.,
428 2015), *PmSWEETs* can be classified into four clades, and the number of 11 species SWEET
429 genes members in Clade III was larger than that in other clades (Figure 1), suggesting that Clade
430 III may have expanded during evolution. Conserved motif analysis indicates that some special
431 motifs only reside in some certain *PmSWEET* gene members. For instance, motif 8 was uniquely
432 present in *PmSWEET11* and *PmSWEET17*; and motif 10 was uniquely present in *PmSWEET3*
433 and *PmSWEET15*. These results are consistent with those of other plants, such as *Arabidopsis*
434 (Chen et al., 2010), rice (Yuan and Wang, 2013), banana (Miao et al., 2017) and wheat (Gautam
435 et al., 2019). Studies have disclosed that gene structural diversity and conserved protein motif
436 divergence performed key roles in the evolution of the SWEET gene family (Xu et al., 2012),
437 some *PmSWEETs* harbored unique conserved motifs, implying it may be responsible for the
438 functional diversity of SWEET in *P. mume*.

439 Gene duplication, including tandem and segmental duplication events, is the origin of gene
440 family expansion and genomic evolution in plants (Cannon et al., 2004; Ganko et al., 2007). In
441 this study, two pairs of *PmSWEETs* were detected as tandem duplications, and four pairs of

442 *PmSWEETs* were segmental duplications. This outcome was consistent with those of other
443 studies on *SWEET* duplication, including segmental and tandem duplications (Feng et al., 2015;
444 Miao et al., 2017; Gao et al., 2018; Jiang et al., 2020).

445 The *cis*-elements in the promoter play an essential role in gene regulation. All *PmSWEETs*
446 contain at least one light-responsive and anaerobically induced *cis*-element, suggesting that the
447 two elements have an essential role in *PmSWEET* regulation. Moreover, 10 *PmSWEETs*
448 contained one or more low-temperature responsive *cis*-elements (Table S6), indicating that these
449 *PmSWEETs* may play important roles in the response to cold stress. However, whether and how
450 these *cis*-elements work in *P. mume* requires further research.

451 Studies have shown that under low-temperature stress, the soluble sugar content in plants
452 increases, and sugar transporters maintain the balance of osmotic potential through the balance
453 and distribution of sugar, thus improving the cold tolerance of plants (Yamada et al., 2010).
454 Numerous studies have also verified that *SWEETs* are involved in maintaining sugar
455 homeostasis in plant organs and promoting plant adaptation to low temperatures (Seo et al.,
456 2011; Chardon et al., 2013; Klemens et al. 2013; Chandran, 2015; Le Hir et al. 2015; Miao et al.,
457 2017; Wang et al., 2018; Zhang et al., 2019; Zhang et al., 2020). Transcriptome analysis showed
458 that *PmSWEETs* were differentially expressed in different tissues and during dormancy release
459 and cold acclimation. *PmSWEET5* expression was not detected in any tissue/organ that we used,
460 indicating that its expression may be variety -specific or time-specific. Some *PmSWEETs* had
461 specific expression patterns in different organs (Figure 7A). For example, expression of
462 *PmSWEET10* was detected only in ‘Zaolve’ buds at dormancy (stage EDII) and ‘Songchun’
463 stems in winter in Beijing; *PmSWEET16* expression was detected only in *P. mume* buds, which
464 indicates that the genes are expressed only in specific tissues or varieties, such organ-specific
465 expression patterns was also observed in wheat (Gao et al., 2018; Gautam et al., 2019), walnut
466 (Jiang et al., 2020), tea (Wang et al., 2018) and cabbage (Zhang et al., 2019). *AtSWEET5*, the
467 homologue of *PmSWEET10* and *PmSWEET16*, plays a key role in seed germination, and
468 expressed at different stages of pollen development (Engel et al., 2005). The results from
469 expression studies of different organs indicate a role for *PmSWEET10* and *PmSWEET16* in
470 pollen development, suggesting they might have a similar role as *AtSWEET5*. *PmSWEET1*,
471 *PmSWEET6*, *PmSWEET9*, *PmSWEET12* and *PmSWEET17* were strongly expressed in fruit,
472 indicating that these genes may regulate sugar allocation during fruit ripening. Such specific high
473 expression of *SWEETs* in fruits has also been found in pineapple (Guo et al., 2018), sweet orange
474 (Zheng et al., 2014) and apple (Zhen et al, 2018), it can be inferred that *SWEET* protein plays an
475 important role in fruit development and ripening.. *PmSWEET4* (Clade I) and *PmSWEET7* (Clade
476 IV) were strongly expressed in roots, this results had similar expression patterns to previous
477 studies, that *SWEETs* in Clade IV were highly expressed in the root cortex and encoded proteins
478 such as specific fructose uniporters in the root vacuole membrane (Guo et al., 2014).

479 The present results also show that most of the *PmSWEET* genes are expressed more strongly
480 at different endo-dormancy stages in flower bud and fruit tissues than in other tissues and that
481 these genes are differentially expressed during flower development (Figure 7A, 7B). Together,
482 these results suggest that the *P. mume* SWEET family is intimately associated with reproductive
483 development and that different genes are specifically involved during different developmental
484 stages. In rice, *Arabidopsis* and soybean, the expression of SWEET genes is also higher in
485 reproductive tissues than in other tissues (Yuan et al., 2014; Patil et al., 2015). *PmSWEETs* also
486 have different expression levels during dormancy release in flower buds (from November to
487 February). Thus, we speculate that these *PmSWEETs* may participate in the cold reaction at low
488 temperatures to protect the flower bud. In addition, some *PmSWEETs* were expressed more at
489 colder temperatures in the spring (3.2~5.3 °C) and at approximately -5 °C in the winter (Figure
490 8A), indicating that these two temperatures may trigger their cold stress response and increase
491 *PmSWEET* expression to reduce stress injury.

492 The qRT-PCR analysis suggested that six of 17 *PmSWEET* genes (*PmSWEET5*,
493 *PmSWEET6*, *PmSWEET9*, *PmSWEET11*, *PmSWEET15*, and 16) were not expressed in the stem,
494 which was consistent with the transcriptome data. *PmSWEETs* were activated by low
495 temperature (4 °C) and increased or decreased in expression with the extension of treatment time
496 (Figure 9). The expression levels of five *PmSWEETs* (*PmSWEET2*, *PmSWEET4*, *PmSWEET7*,
497 *PmSWEET8*, and *PmSWEET10*) in ‘Songchun’ and three *PmSWEETs* (*PmSWEET2*, *PmSWEET7*
498 and *PmSWEET8*) in ‘Zaolve’ decreased with increasing treatment times (Figure 9), which
499 suggested that these genes might be negatively regulated by low temperatures and result in
500 increased cold sensitivity. The expression levels of two *PmSWEETs* (*PmSWEET13* and
501 *PmSWEET14*) in ‘Songchun’ and three *PmSWEETs* (*PmSWEET1*, *PmSWEET12*, and
502 *PmSWEET13*) in ‘Zaolve’ increased with prolonged treatment (Figure 9), which suggested that
503 these genes might be positively regulated by cold stress responses and increase cold sensitivity.
504 The discrepancy in expression patterns between *PmSWEET1*, *PmSWEET4*, *PmSWEET10*,
505 *PmSWEET12*, *PmSWEET14* and *PmSWEET17* is potentially due to genetic differences between
506 ‘Songchun’ and ‘Zaolve’.

507 5. Conclusions

508 In summary, our study is the first to perform genome-wide identification and
509 characterization of SWEETs in *P. mume*, including chromosomal location, duplicated genes,
510 gene structure, phylogenetic relationships and conserved motifs. In addition, the expression
511 profiles of the *PmSWEET* genes in different tissues and geographic locations were also examined
512 based on the RNA-seq data. Furthermore, the expression profiles of these *PmSWEET* genes
513 under cold stress conditions were analyzed by qRT-PCR assay. Our results could provide
514 important information for further research on the biological functions of *PmSWEETs*.

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521 Conflict of Interest

522 The authors declare that the research was conducted in the absence of any commercial or
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524 Author Contributions

525 LS: conceptualization. PL and ML: data curation. ZW: formal analysis and software. LS, QZ and
526 TC: funding acquisition and writing reviews and editing. ZW and JM: methodology. ZW:
527 validation, visualization, and drafting the manuscript. All authors contributed to writing and
528 approved the final manuscript.

529 Data Availability Statement

530 The original contributions presented in the study are included in the article/Supplementary
531 Material, further inquiries can be directed to the corresponding author/s.

532 Supplementary Material

533 Supplemental information for this article can be found online at

534 Supplementary Figure 1 | Schematic representation of *PmSWEET* proteins.

535 Supplementary Figure 2 | Phylogenetic trees of *Arabidopsis thaliana*, *Prunus mume* and Rice

536 Supplementary Figure 3 | Schematic diagram of *PmSWEET* protein motifs

537 Supplementary Table 1 | Primer sequences used for qRT-PCR

538 Supplementary Table 2 | Information for the proteins used in the present study

539 Supplementary Table 3 | TM helix Locus of *PmSWEETs*

540 Supplementary Table 4 | The specific number of genes in the Clades used in the present study

541 Supplementary Table 5 | Duplication events between *P. mume* and *A. thaliana*, *P. armeniaca* and

542 *P. salicina*

543 Supplementary Table 6 | The data of cis-acting element in *PmSWEETs* promoters

544 Supplementary Table 7 | Expression profiles of 17 *PmSWEET* genes in five different tissues

545 (root, stem, leaf, bud and fruit) (RPKM)

546 Supplementary Table 8 | Expression profiles of *PmSWEET* genes during the process of flower

547 bud dormancy release (RPKM)
548 Supplementary Table 9 | Expression profiles of 17 *PmSWEET* genes in different regions and
549 seasons (FPKM)
550 Supplementary Flie 1 | Protein sequences of *P. mume*
551 Supplementary Flie 2 | Domain architecture of *PmSWEETs*
552

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Table 1 (on next page)

Table 1. The *PmSWEET* gene family members in *P. mume*.

Table 1. The *PmSWEET* gene family members in *P. mume*.

Name	Gene ID	Clade	CDS (bp)	No. of amino acids	Molecular weight (kDa)	Theoretical pI	TMHs	No. of MtN3/saliv a domain	Locus
PmSWEET1	Pm007067	III	849	282	31.38	8.34	7	2	Pa2:21184396..21186332
PmSWEET2	Pm008206	IV	759	252	27.74	8.50	7	2	Pa2:31718730..31721555
PmSWEET3	Pm010330	I	1248	415	46.25	9.76	8	2	Pa3:3891190..3895205
PmSWEET4	Pm011260	I	708	235	26.45	8.30	7	2	Pa3:9921623..9924001
PmSWEET5	Pm013198	II	519	172	19.42	8.97	5	1	Pa4:2433448..2434735
PmSWEET6	Pm015728	II	708	235	25.67	9.21	5	2	Pa4:21122646..21124537
PmSWEET7	Pm017566	IV	735	244	26.99	9.14	7	2	Pa5:12327097..12328384
PmSWEET8	Pm018875	III	1743	580	63.43	8.34	6	2	Pa5:20984940..20990591
PmSWEET9	Pm019954	III	828	275	30.68	9.20	7	2	Pa6:436315..437664
PmSWEET10	Pm021931	II	708	235	26.60	8.59	6	2	Pa6:12459796..12461199
PmSWEET11	Pm022695	I	417	138	15.96	9.74	3	1	Pa6:19934418..19935334
PmSWEET12	Pm022696	I	651	216	23.21	8.78	5	2	Pa6:19944525..19945680
PmSWEET13	Pm024167	II	780	259	28.66	9.37	6	2	Pa7:10796671..10798904
PmSWEET14	Pm024554	III	318	105	-	-	2	1	Pa7:13005181..13005663
PmSWEET15	Pm024555	III	891	296	33.14	8.61	7	2	Pa7:13012731..13014646
PmSWEET16	Pm024712	II	639	212	23.95	8.37	5	2	Pa7:13852243..13854234
PmSWEET17	Pm030352	I	510	169	19.26	9.14	4	1	scaffold54:138478..139392

Figure 1

Figure 1. Phylogenetic tree of SWEET sequences from *P. mume* and other plant species.

Clades I, II, III, and IV are indicated by blue, indigo, orange and pale yellow branch lines, respectively. At, *A. thaliana*; Os, *O. sativa*; Pa, *P. armeniaca*; Pav, *P. avium*; Pc, *P. communis*; Pm, *P. mume*; Pp, *P. persica*; Ps, *P. salicina*; Py, *P. yedoensis* var. *nudiflora*; Md, *M. domestica*; Rc, *R. chinensis*.

Figure 2

Figure 2. Phylogenetic relationship, conserved motif and gene structure analysis of *PmSWEET* genes.

A: The ML phylogenetic tree of *PmSWEET* genes. The SWEET genes were grouped into four clades, and blue, purple, red, and green represents Clades I, II, III, and IV, respectively. B: The motif composition of *PmSWEET* proteins. Ten motifs were displayed in different colored rectangles. Motif1: GVVWFLYGLLKKDLFIAIPNGLGFJLGLVQLILYAIYR, Motif2: TKKRSLIVGIJCIVFNIIIMYASPLTIMKLVIKTKSVEYMPFYLSLFLFLN, Motif3: LVITINGFGAVIELIYJAIFIIYAPKKKRKKI, Motif4: APVPTFYRIKKKSTEEFQSVPYVAALLN, Motif5: WYGMPFVHPDN, Motif6: FGILGNIISFLLFL, Motif7: STNWDDDD, Motif8: PMTTLKRIMKKNEFTEQYLSGIPYLMT, Motif9: AMLWLYYGLLKPN, Motif10: NCZGCKDQYQHPQKCKE. Detailed information is shown with logos obtained from the MEME Suite website in Supplementary Figure 3. C: Exon-intron organization of *PmSWEET* genes. Green and black correspond to exons and introns, respectively.

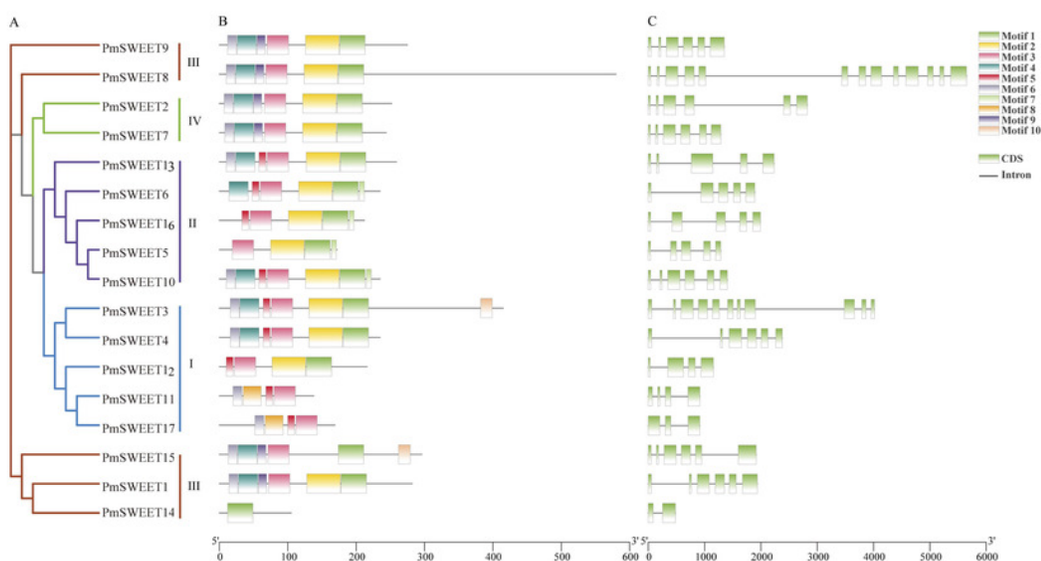


Figure 3

Figure 3. Schematic representations of the chromosomal location of the *PmSWEET* genes.

The chromosome number is indicated on the top of each chromosome and scaffold. Scf54 indicates scaffold54. Green and red gene names indicate tandem duplicated gene pairs.

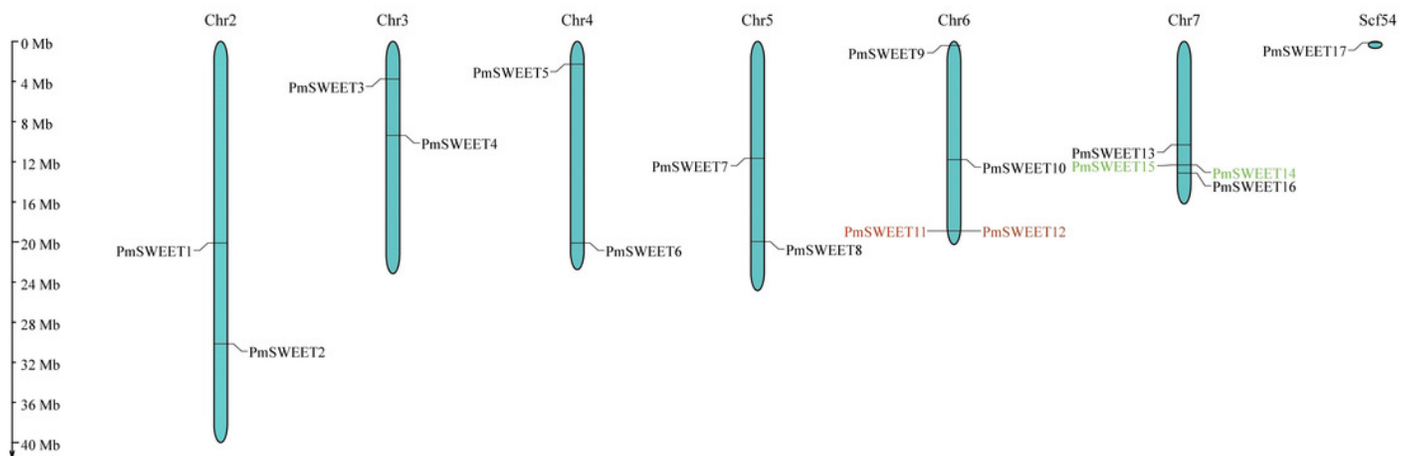


Figure 4

Figure 4. The Circos figure for *PmSWEET* segmental duplication links.

The red lines indicate segmented duplicated gene pairs.

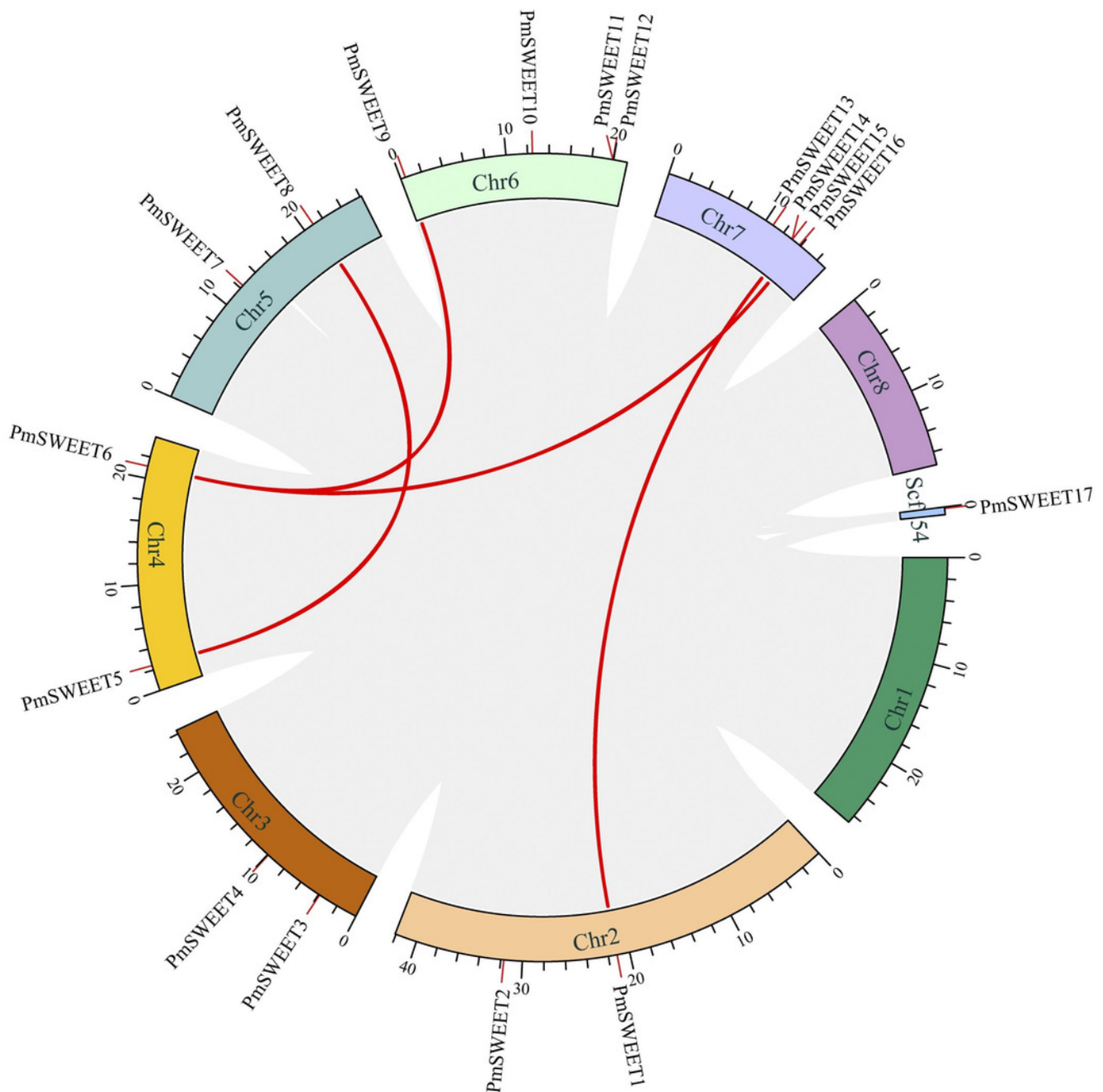


Figure 5

Figure 5. Synteny of SWEET genes in different genome of *P. mume*, *A. thaliana*, *P. armeniaca* and *P. salicina*.

A: Synteny of *PmSWEET* and *AtSWEET* gene pairs. B: Synteny of *PmSWEET* and *PaSWEET* gene pairs. C: Synteny of *PmSWEET* and *PsSWEET* gene pairs.

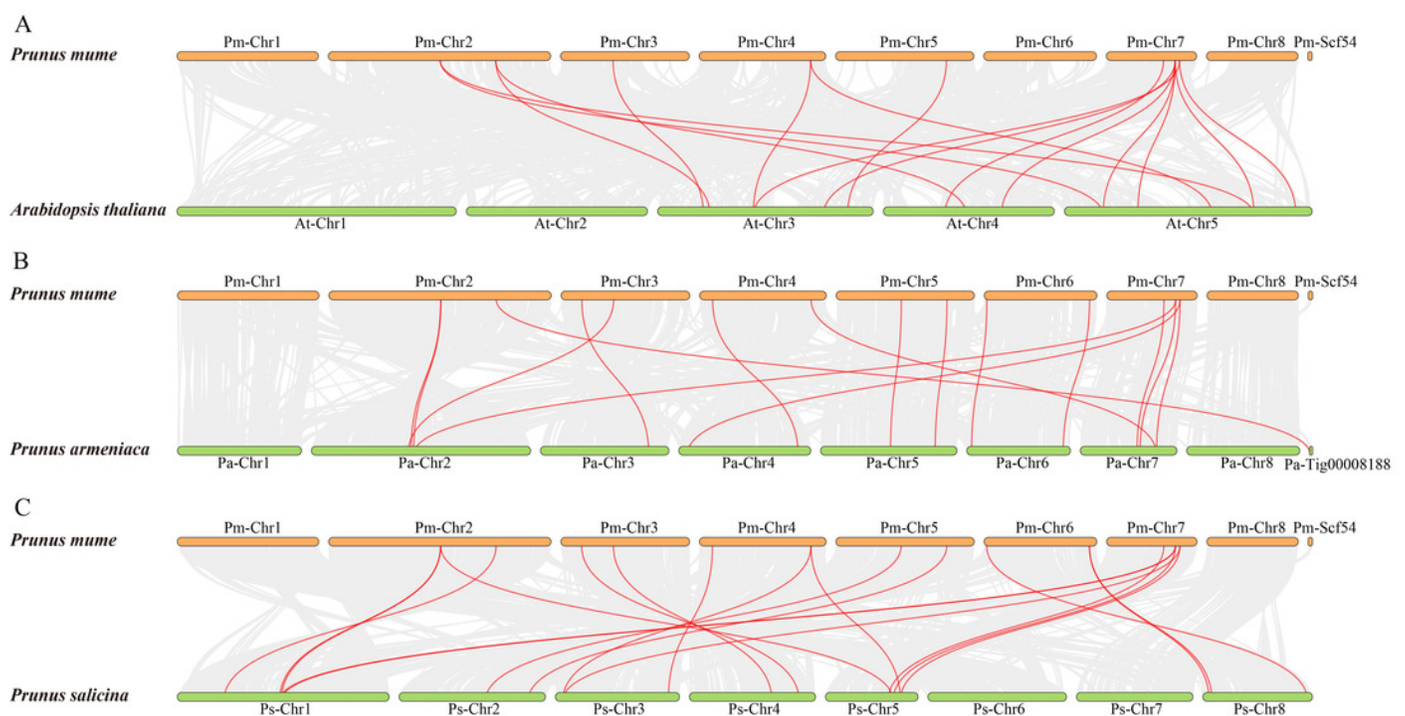


Figure 6

Figure 6. Predicted cis-elements responding to plant growth regulation, hormone response, and stresses response present in the promoter of *PmSWEET* genes

Different colored boxes represent different elements and their positions in each *PmSWEET* promoter. The SWEET genes are classified into four clades, and blue, indigo, purple red, and green represent Clades I, II, III, and IV, respectively.

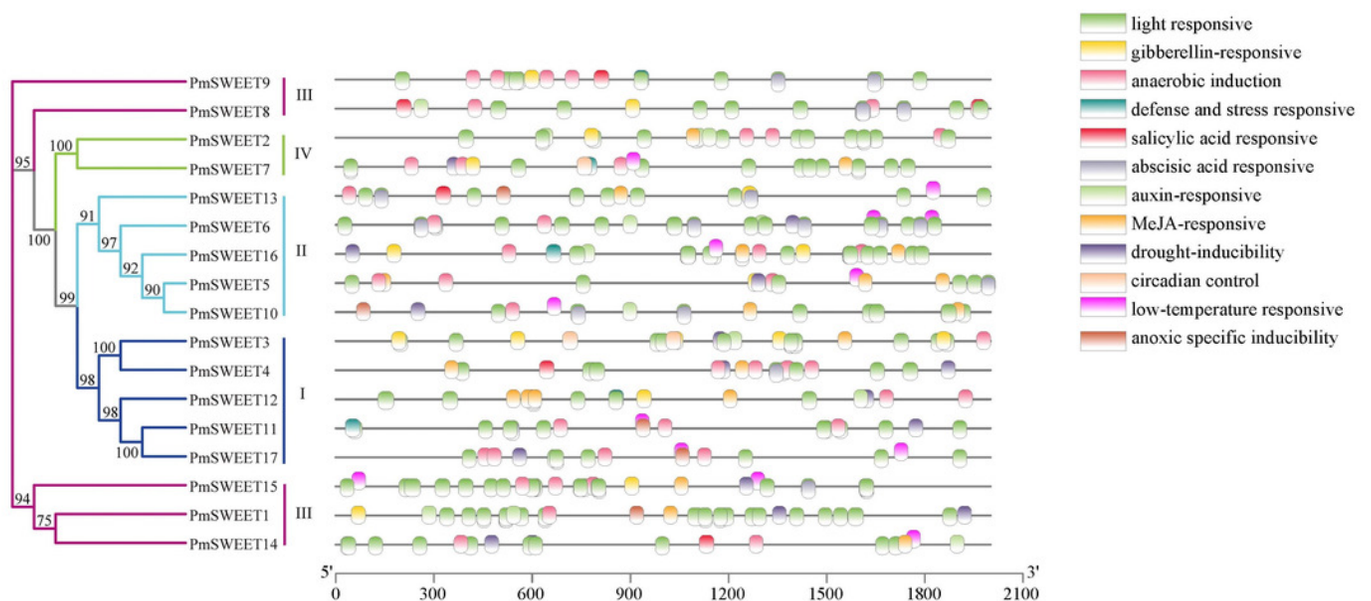


Figure 7

Figure 7. Expression profiles of *PmSWEET* genes in different tissues and different flower buds stage

A: Expression profiles of *PmSWEETs* in different tissues. B: Expression profiles of *PmSWEETs* in the flower bud during dormancy. EDI: Endo-dormancy I, November; EDII: Endo-dormancy II, December; EDIII: Endo-dormancy III, January; NF: Natural flush, February. A 2-based log function conversion is performed on the expression amount, and then normalized by row using min-max method. The color scale on the right of the heat map refers to relative expression level, and the color gradient from blue to red shows an increasing expression level.

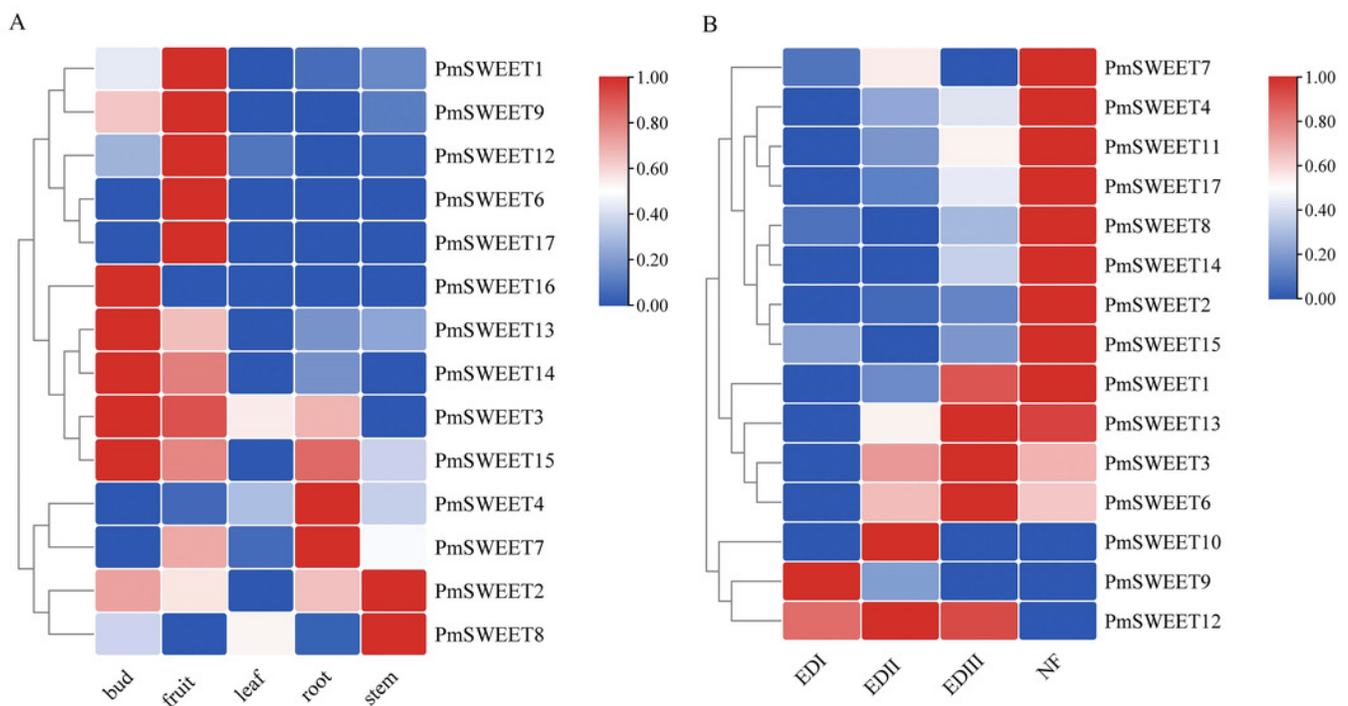


Figure 8

Figure 8. Expression profiles of *PmSWEET*s in stems in different seasons and regions

A: Expression profiles of *PmSWEET*s in stems of ‘Songchun’ in different regions (Beijing, Chifeng and Gongzhuling) and seasons (autumn, winter and spring). B: Comparison of differential expression profiles of stems in Beijing, Chifeng and Gongzhuling during different seasons. A 2-based log function conversion is performed on the expression amount, and then normalized by row using min-max method. The color scale on the right of the heat map refers to relative expression level, and the color gradient from blue to red shows an increasing expression level. Aut, Autumn; Win, Winter; Spr, Spring. BJ, Beijing; CF, Chifeng; GZL, Gongzhuling.

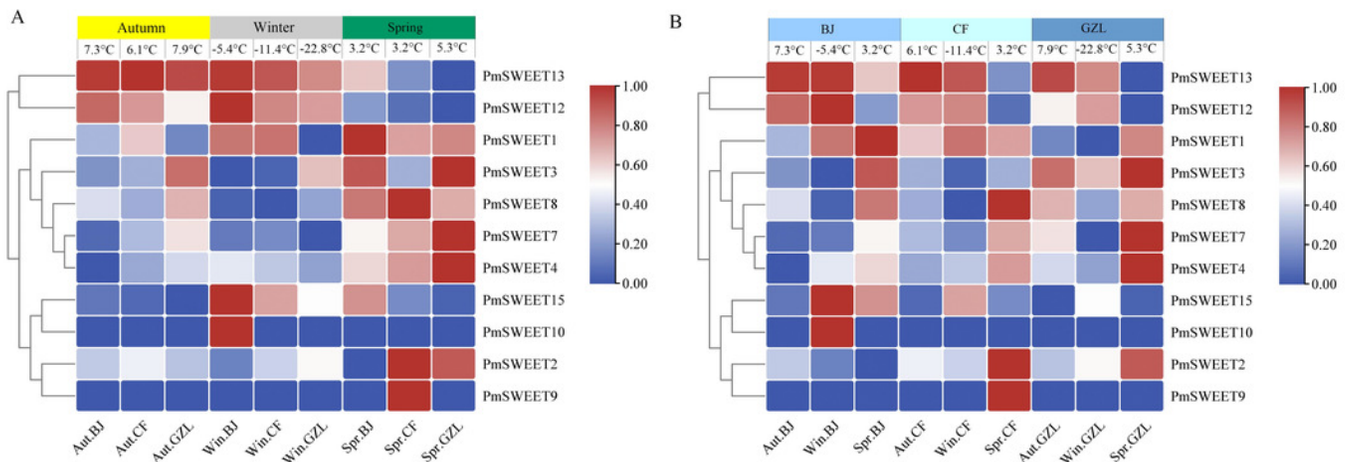


Figure 9

Figure 9. Expression patterns of 11 *PmSWEET* genes under artificial low temperature treatments

The relative quantification method ($2^{-\Delta\Delta C_t}$) was used to evaluate the transcript levels of 11 *PmSWEET* genes. Error bars are standard deviation of three replicates. The statistical analyses of 'Zaolve' and 'Songchun' were independent carried out using SPSS22.0, the one-way ANOVA analysis of variance was calculated by least significant difference (LSD) and Student-Newman-Keuls test, different letters above the bars indicate significant differences ($p = 0.05$). Black letters indicate 'Zaolve', red letters indicate 'Songchun'. GraphPad Prism6 software was used to draw the diagram.

