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1 **Title: Transcriptome profiling reveals the crucial biological pathways involved in**
2 **cold response in Moso bamboo (*Phyllostachys edulis*)**

3
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19 **Running title:** Transcriptomic profiling of cold response in Moso bamboo

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41

42 **Abstract**

43 Most bamboo species including Moso bamboo (*Phyllostachys edulis*) are tropical or
44 subtropical plants that greatly contribute to human wellbeing. Low temperature is one
45 of the main environmental factors restricting bamboo growth and geographic
46 distribution. Our knowledge of the molecular changes during bamboo adaption to cold
47 stress remains limited. Here, we provided a general overview of the cold-responsive
48 transcriptional profiles in Moso bamboo by systematically analyzing its
49 transcriptomic response under cold stress. Our results showed that low temperature
50 induced strong morphological and biochemical alternations in Moso bamboo. To
51 examine the global gene expression changes in response to cold, 12 libraries
52 (non-treated, cold-treated 0.5 h, 1 h and 24 h at -2°C) were sequenced using an
53 Illumina sequencing platform. Only a few differentially expressed genes (DEGs) at
54 early stage while a large number of DEGs at late stage were identified in this study,
55 suggesting that the majority of cold response genes in bamboo are late-responsive
56 genes. A total of 222 transcription factors from 24 different families were
57 differentially expressed during 24h cold treatment, and the expressions of several
58 well-known C-repeat/dehydration responsive element-binding factor (CBF) negative
59 regulators were significantly up-regulated in response to cold, indicating the existence
60 of special cold response networks. Our data also revealed that the expression of genes
61 related to cell wall and the biosynthesis of fatty acids were altered in response to cold
62 stress, indicating their potential roles in the acquisition of bamboo cold tolerance. In
63 summary, our studies showed that both plant-kingdom conserved and species-specific
64 cold response pathways exist in Moso bamboo, which lays the foundation for
65 studying the regulatory mechanisms underlying bamboo cold stress response and
66 provides useful gene resources for the construction of cold-tolerant bamboo through
67 genetic engineering in the future.

68

69

70 **Introduction**

71 Bamboo is one of the most important non-timber forest products, covering over 30
72 million hectares (ha) worldwide and accounting for 68.8 billion US dollars in
73 international trade in 2018 (King 2019). Moso bamboo (*Phyllostachys edulis*), is one
74 of the most economically important bamboo species, serves as a promising
75 bio-resource for renewable forestry products, and accounts for over two-thirds of total
76 bamboo growing area (4.43 million ha) in China (Peng et al. 2013). The distribution
77 of bamboo in nature is greatly influenced by agro-climatic zones, human interventions,
78 and climatic factors (Lucina Yeasmin et al. 2017). Most bamboo species, including
79 Moso bamboo, are commonly located in tropical or subtropical climatic regions, and
80 temperature is one of the major environmental factors that control Moso bamboo
81 growth and geographic distributions (Gu et al. 2010; Numata et al. 1957; Wenwei
82 1991; Xu and Qin 2003; Yeasmin et al. 2015). Over the past two decades, significant
83 progress has been made in understanding the plant responses to chilling (0–15 °C) and
84 freezing (<0 °C) in *Arabidopsis thaliana* (Lee et al. 2005), rice (Zhang et al. 2014),
85 cotton (Kargiotidou et al. 2010), soybean (Calzadilla et al. 2016) and tomato (Weiss
86 and Egea-Cortines 2009). In contrast, our understanding of the mechanisms that
87 underlie bamboo cold stress response is surprisingly limited.

88 In the currently accepted model, cold stress first acts on the signal perception and
89 transduction pathways, which induces transcriptional control, and consequently
90 activates a variety of cold-regulated (COR) proteins (Guo et al. 2018; Zhu 2016).
91 Molecular, physiological and metabolic studies demonstrated that low temperature
92 leads to changes in membrane fluidity, initiating the cellular cold response through
93 calcium (Ca^{2+}) signaling pathways (Sangwan et al. 2001; Zhang et al. 2014). Ca^{2+} is
94 recognized by calcium-binding proteins such as calmodulin (CaM), CaM-like proteins
95 (CML), Ca^{2+} dependent protein kinases (CDPKs), and calcineurin B-like proteins
96 (CBLs) (Kudla et al. 2018). These calcium-binding proteins also known as Ca^{2+}
97 sensors are induced early (3 h of exposure to 0 °C) during cold stress in *Arabidopsis*
98 and rice (Abbasi et al. 2004; Lee et al. 2005). Mitogen-activated protein kinase

99 (MAPK) cascades, which are activated by various stress signal messengers, also play
100 a role in cold response. A typical MAPK cascade is composed of three protein kinases:
101 MAP kinase kinase kinase (MAPKKK or MEKK), MAP kinase kinase (MAPKK,
102 MKK, or MEK), and MAP kinase (MAPK or MPK). Among them, MKK2 induces
103 the expression of *COR* genes to enhance freezing tolerance in Arabidopsis (Teige et al.
104 2004). MPK3/6, on the other end, negatively regulates freezing tolerance via
105 phosphorylation and destabilization of the inducer of *CBF* expression 1 (ICE1), which
106 is a basic-helix-loop-helix (bHLH) transcription factor and acts as the master
107 regulator of cold response in Arabidopsis (Li et al. 2017; Zhao et al. 2017). The
108 integration of these signals is mediated through the coordination of transcriptional
109 activators and repressors, many of which have been well characterized (Chinnusamy
110 et al. 2007). The typical transcriptional regulation pathways of cold stress are C-repeat
111 binding factor (CBF)-dependent and CBF-independent pathways. The key
112 components of the CBF-dependent pathway are ICE1-CBF-COR, and they play a
113 predominant role in cold tolerance (Guo et al. 2018; Zhou et al. 2011). ICE1 mediates
114 the CBF-dependent pathway by positively regulating the expression of *CBFs* (Guo et
115 al. 2018). Overexpression of *ICE1* leads to an increased expression of *CBFs* and
116 improves cold tolerance in transgenic Arabidopsis (Chinnusamy et al. 2003). A large
117 number of genes directly or indirectly participate in cold regulation through regulating
118 ICE1 at the level of transcription, translation and post-translation (Agarwal et al. 2006;
119 Maruyama et al. 2014). CBFs bind to the promoter of *COR* genes to activate their
120 expression and confer increased freezing tolerance in plants (Gilmour et al. 2004).
121 CBF homologs have been characterized in many plant species such as rice (Dubouzet
122 et al. 2003), maize (Qin et al. 2004), barley (Morran et al. 2011) and soybean
123 (Kidokoro et al. 2015). Transgenic plants overexpressing *CBFs* show enhanced cold
124 tolerance compared to wild type (Ito et al. 2006; Kasuga et al. 2004). The expressions
125 of *CBFs* are also negatively regulated by a number of transcription factors such as
126 *MYB15* and *ZAT12* (Agarwal et al. 2006; Maruyama et al. 2009).

127 Plant adaption to cold stress involves changes at cellular and molecular levels,
128 which are governed by plant hormones (Lado et al. 2016). Abscisic acid (ABA) is the

129 key plant hormone that is involved in plant responses to abiotic stresses (Gusta and
130 Wisniewski 2013). In many species, cold stress is accompanied by the increased
131 expression level of the 9-cis-epoxycarotenoid dioxygenase (*NCED*) gene, which
132 encodes one of the key enzymes for ABA biosynthesis and leads to the induction of
133 endogenous ABA (Mantyla et al. 1995). The ABA signal is perceived through the
134 ABA receptor complex, which is composed of PYRABACTIN RESISTANCE 1
135 (PYR1), PYR1-like protein (PYL) and regulatory components of the ABA receptor
136 (RCAR) family of START proteins, and induces broad gene expressions in response
137 to abiotic stresses (Lee and Luan 2012). The current consensus is that both
138 ABA-dependent and ABA-independent pathways are involved in the plant responses
139 to cold stress (Lado et al. 2016). The ethylene pathway seems to play a negative role
140 in regulating freezing tolerance partly by inhibiting the functions of *CBF* or *DREB* in
141 *Arabidopsis* (Kazan 2015; Shi et al. 2012). Recent reports showed that the ethylene
142 pathway also plays a positive role in cold stress in *Arabidopsis*, tomato, rice and
143 tobacco (Catala and Salinas 2015; Tian et al. 2011; Zhang et al. 2009) . Therefore, the
144 role of ethylene in cold tolerance appears to be species dependent. A key plant
145 response to cold is growth repression, through which plants might re-allocate
146 resources from growth to processes that help to increase cold tolerance (Eremina et al.
147 2016). Gibberellins (GAs) are well-known growth promoting hormones, and both GA
148 metabolism and signaling are targeted by cold stress (Achard et al. 2008; Seo et al.
149 2009). Cold induces the expression of *GA 2-oxidases* (*GA2OX*) gene, which encodes a
150 key enzyme for the inactivation of bioactive GAs (Xu et al. 1999). Studies from
151 *Arabidopsis* revealed that *CBF3* promotes the accumulation of DELLA proteins,
152 which are key negative regulators in GA signaling pathway, and lead to retarded plant
153 growth in response to cold (Zhou et al. 2017). Auxin not only plays vital roles in plant
154 growth and development, but also mediates the cold response (Rahman 2013). It was
155 reported that cold stress affects the auxin response pathway primarily through the
156 repression of the auxin transport pathway instead of a signaling pathway, and this
157 effect is linked to the inhibition of intracellular trafficking of a subset of auxin efflux
158 and influx carriers in *Arabidopsis* (Shibasaki et al. 2009). However, no study reported

159 the role of plant hormone in bamboo responses to cold stress.

160 Cold stress often leads to multiple physiological changes, such as cell membrane
161 damage associated with ion leakage (Whitlow et al. 1992), changes in MDA content
162 (Kong et al. 2016) and proline content (Hayat et al. 2012). Cold stress also stimulates
163 the accumulation of some anti-stress enzymes, such as SOD (Abid et al. 2016; Reddy
164 et al. 2004), POD (Miller et al. 2010) and APX (Caverzan et al. 2012). Evaluating
165 these physiological responses to cold stress in Moso bamboo would effectively
166 determine the effects of cold stress, as well as broaden our understanding of the cold
167 adaptation process in this important species.

168 The draft genome sequence of Moso bamboo was released in 2013 (Peng et al.
169 2013), and an advanced version has recently been mapped at the chromosomal level
170 (Zhao et al. 2018). The draft genomes of four other bamboo species, *Olyra latifolia*,
171 *Raddia guianensis*, *Guadua angustifolia* and *Bonia amplexicaulis* have been
172 published very recently (Guo et al. 2019), providing an excellent opportunity for
173 cold-related studies of this economically and ecologically important grass to be
174 undertaken. The Moso bamboo genome contains 24 *DREB* transcription factors, and
175 *PeDREB1* is strongly induced by cold treatment (Liu et al. 2012; Wu et al. 2015). A
176 recent study revealed that the MYB transcription factor *PheMYB4-1* regulates the cold
177 response in Moso bamboo. Transgenic *Arabidopsis* plants overexpressing *PheMYB4-1*
178 display increased cold and freezing tolerance, and *PheMYB4-1* may
179 induce *CBF* expression and activate the downstream *COR* genes (Hou et al. 2018). In
180 addition, 13 *TIFY* family transcription factors show up-regulation in response to cold
181 stress (Huang et al. 2016). All these data suggest that the transcriptional regulation is
182 crucial for Moso bamboo's tolerance to low temperature; thus, it is very important to
183 identify the cold-regulated transcription factors in Moso bamboo.

184 In this work, Moso bamboo was used as it is the most common type of bamboo in
185 tropical and subtropical areas. The morphological and physiological changes were
186 recorded after cold treatment; RNA-seq was used to analyze the dynamic changes in
187 transcription that occur at different time points during cold treatment. Two objectives
188 were addressed in this study, namely the identification of candidate genes

189 participating in cold regulation pathways, and the analysis of expression profiles of
190 key genes involved in cold regulation in Moso bamboo. Overall, our study revealed a
191 broad overview of the Moso bamboo cold-responsive transcriptome, and uncovered
192 cold signal perception and the responsive pathway in Moso bamboo. To the best of
193 our knowledge, this is the first systematic study of the transcriptome profiling of
194 Moso bamboo under cold stress. Our study revealed cold-regulated candidate genes
195 that may potentially be used for generating plants with enhanced cold tolerance.

196

197 **Material and Methods**

198 **Plant material and growth conditions**

199 The seeds of Moso bamboo (*Phyllostachys edulis*) and Ma bamboo (*Dendrocalamus*
200 *latiflorus* Munro) used in this study were collected from Guangxi Zhuang
201 Autonomous Region (Guangxi, China). Bamboo seeds were thoroughly washed with
202 sterile water and soaked in sterile water for 16 h, and germinated in soil at 22 °C
203 under long-day conditions (16 h of cool white fluorescent light, photon flux of 70 μ
204 $\text{mol m}^{-2} \text{s}^{-1}$). For cold treatments, 3-weeks old seedlings at the three-leaf stage were
205 subjected to -2 °C in a freezing chamber (LGX-400B-LED) for 24 h or 72 h, and then
206 allowed to recover at 25°C for 5 days. Seedlings of the control group were grown at
207 25 °C continuously. Surface structural changes of the abaxial side of the bamboo
208 leaves were imaged using a HITACHI TM3030 PLUS Tabletop Scanning Electron
209 Microscope (SEM) (Hitachi, Japan). To calculate the survival rate, around 30
210 seedlings were treated with cold for the indicated time, and then allowed to recover
211 for 5 days. The numbers of seedlings alive and dead were calculated and the data were
212 statistically analyzed. All experiments were repeated independently at least 3 times.

213

214 **Measurement of electrolyte leakage, relative malondialdehyde (MDA) content** 215 **and superoxide dismutase (SOD), peroxidase (POD), and ascorbic acid** 216 **peroxidase (APX) activities**

217 Measurement of electrolyte leakage was performed as described previously with

218 some modifications (Duan et al., 2017). Briefly, leaves were detached from the
219 cold-treated plants and immersed in 50 mL tubes containing 30 mL water, and then
220 the conductivities were measured immediately (S0) with an electrical conductivity
221 meter (type starter 300C, OHAUS, America). The samples were collected after
222 shaking at 120 rpm for 15 min in a vacuum condition, and the conductivities (S1)
223 were determined. Subsequently, the samples were boiled in a water bath with agitation
224 at 120 rpm for 15 min, and the conductivities were measured again after cooling to
225 25 °C (S2). The relative electrolytic leakage (%) was calculated as
226 $(S1-S0)/(S2-S0)\times 100$.

227 All the antioxidant enzymes were measured based on the protocol reported
228 previously with some modifications (Ara et al. 2013). For each sample, five bamboo
229 whole seedlings were pooled together for analysis. For MDA content measurement,
230 whole seedlings (around 0.1 g) were homogenized and mixed with 1 mL MDA
231 reaction buffer consisting of 0.5 % (v/v) thiobarbituric acid and 20 % (v/v)
232 trichloroacetic acid. The mixture was incubated in a water bath at 100 °C for 30 min,
233 and then the reaction was stopped in an ice bath. The mixture was then centrifuged at
234 10,000 g for 10 min, and the absorbance of the supernatant was measured at 450 nm,
235 532nm, and 600 nm respectively. The MDA content was calculated based on the
236 protocol of the MDA content Assay Kit (Solarbio, China). For the measurement of the
237 activities of SOD, POD, and APX, Moso bamboo seedlings (0.1 g) were homogenized
238 thoroughly in 50 mmol potassium phosphate buffer (pH 7.8) containing 1 %
239 polyvinylpyrrolidone. The homogenate was centrifuged at 13,000 g for 20 min at 4 °C.
240 The activities of those enzymes were measured using a SOD activity Assay Kit
241 (Solarbio, China), POD activity Assay Kit (Solarbio, China) and APX activity Assay
242 Kit (Solarbio, China), respectively according to the manufacturers' instructions.

243

244 **Determination of proline (Pro) content**

245 Proline content in Moso bamboo seedling was measured by sulfosalicylic acid-acid
246 ninhydrin method using Pro content Assay Kit (Solarbio, China) (Abraham et al.
247 2010). Briefly, around 0.1 g of tissues were boiled in 1 mL of 3% sulphosalicylic acid

248 at 95 °C for 15 min. The homogenate was centrifuged at 10,000 g for 10 min. About
249 0.5 mL of supernatant was transferred to a new tube containing 0.5 mL of acetic acid
250 and 0.5 mL of acidified ninhydrin reagent. After 30 min of incubation at 95 °C,
251 samples were kept at room temperature for 30 min and 1 mL of toluene was added to
252 the samples, which were then shaken at 150 rpm to extract red products. The
253 absorbance of the toluene layer was determined at 520 nm using a spectrophotometer.
254 The Pro content was calculated following the manufacturer's instructions (Solarbio,
255 China).

256

257 **Library preparation and transcriptome sequencing**

258 A total amount of 1 µg RNA from each sample was used for sample preparations.
259 Sequencing libraries were generated using a NEBNext® Ultra™ RNA Library Prep
260 Kit for Illumina® (NEB, USA) and following the manufacturer's instructions. Index
261 codes were added to attribute sequences to each sample. Briefly, mRNA was purified
262 from total RNA using poly-T oligo-attached magnetic beads. Fragmentation was
263 carried out using divalent cations under elevated temperatures in NEB Next First
264 Strand Synthesis Reaction Buffer (5X). First strand cDNA was synthesized using the
265 random hexamer primer and M-MuLV Reverse Transcriptase. Second strand cDNA
266 synthesis was subsequently performed using DNA Polymerase I and RNase H.
267 Remaining overhangs were converted into blunt ends via exonuclease/polymerase
268 activities. After adenylation of 3' ends of DNA fragments, NEB Next Adaptor with
269 hairpin loop structure were ligated to prepare for hybridization. To select cDNA
270 fragments of preferentially 240 bp in length, the library fragments were purified using
271 the AMPure XP system (Beckman Coulter, Beverly, USA). 3 µL USER Enzyme
272 (NEB, USA) was incubated with the size-selected, adaptor-ligated cDNA at 37°C for
273 15 min followed by 5 min at 95°C before the PCR was started. PCR was performed
274 with Phusion High-Fidelity DNA polymerase, Universal PCR primers, and Index (X)
275 Primer. Ultimately, PCR products were purified (AMPure XP system) and library
276 quality was assessed on the Agilent Bioanalyzer 2100 system. The clustering
277 generation of the index-coded samples was performed on a cBot Cluster Generation

278 System using TruSeq PE Cluster Kit v4-cBot-HS (Illumina) according to the
279 manufacturer's instructions. After cluster generation, the library preparations were
280 sequenced on an Illumina platform and paired-end reads were generated.

281

282 **Transcriptome sequencing**

283 Raw data (raw reads) of fastq format were initially processed through in-house perl
284 scripts. In this step, clean data were obtained from the raw data by removing reads
285 containing adapters, poly-N regions and low quality reads from raw data. These clean
286 reads were then mapped to the *P.edulis* genome sequence as a reference. Only reads
287 with a perfect match or one mismatch were further analyzed and annotated based on
288 the reference genome. Hisat2 tools were used in mapping with the reference genome
289 (Pertea et al. 2016). At the same time, Q20, Q30, GC-content and sequence
290 duplication levels of the clean data were calculated. All the downstream analyses
291 were based on clean data with high quality. The Fragments per kilobase of transcript
292 per million fragments mapped (FPKM) of each gene was calculated based on the
293 length of the gene and the read counts mapped to the gene.

294 Differential expression analysis of two time points were performed using
295 the DESeq2 package version 1.22.2 (Love et al. 2014). The resulting *p* values were
296 adjusted using the Benjamini and Hochberg's approach for controlling the false
297 discovery rate. Genes with an adjusted *p*-value ≤ 0.05 and a fold change (FC) ≥ 1.5
298 found by DESeq2 were assigned as differentially expressed.

299

300 **Gene functional annotation**

301 Gene function was annotated based on the following databases: Swiss-Prot
302 (Boeckmann et al. 2003), EuKaryotic Orthologous Groups (KOG) (Tatusov et al.
303 2000), and Kyoto Encyclopedia of Genes and Genomes (KEGG) (Minoru et al.
304 2008). GO enrichment analysis of the DEGs was implemented by the Goseq R
305 packages based on Wallenius non-central hyper-geometric distribution (Young et al.

2010), which can adjust for gene length bias in DEGs. DIAMOND software (version 0.9.22, <https://github.com/bbuchfink/diamond>) was used to align the DEGs to the proteins in KEGG, which is a compendium of databases covering both annotated genomes and protein interaction networks for all sequenced organisms. KEGG pathway is part of KEGG database, and is a compilation of manually verified pathway maps to categorize gene functions with the emphasis on biochemical pathways (Minoru et al. 2008). The output of plant-specific KEGG pathways were populated with the KEGG Orthology (KO) assignments in this study.

314

315 **RNA extraction and qPCR**

316 Total RNA was extracted using the Plant RNA Kit (OMEGA) and reverse
317 transcription using the PrimeScript™ RT Reagent Kit with gDNA Eraser (TaKaRa,
318 Japan) according to the manufacturer's instructions. The quantitative real-time PCR
319 (qPCR) was performed using the TB Green PCR Master Mix Kit (TaKaRa, Japan).
320 The relative expression levels were calculated as described (Huang et al. 2010), and
321 the specific primers for selected 14 DEGs including *PeCML* (PH01000133G0880),
322 *CBL-interacting protein kinase* (*PeCIPK1*, PH01000445G0310), *cold-responsive*
323 *protein kinase* (*PeCRPK1*, PH01000300G0810), *PeMKK4* (PH01003465G0120),
324 *PeMPK3* (PH01000033G1790), *PeICE1* (PH01001045G0070), *PeMYB15*
325 (PH01001287G0090), *PeZAT12* (PH01001038G0420), the *phytochrome-interacting*
326 *factor 3* (*PePIF3*, PH01000595G0290), a MYB family transcription factor
327 *REVEILLE1* (*PeREVI*, PH01000160G0940), *PeWRKY40* (PH01001777G0070),
328 *PeCBF3* (PH01000842G0220), *PeCBF4* (PH01001480G0400), and *PeCOR47*
329 (PH01000447G0290), are listed in Supplemental Table 26. The expression of the
330 housekeeping gene *PeUBQ* (PH01000093G1330) in Moso bamboo was used as
331 internal control as reported previously (Fan et al. 2013).

332

333 **Results**

334 **Effects of cold on Moso bamboo**

335 To assess the effects of cold on Moso bamboo, 3-week-old bamboo seedlings were
336 treated under cold conditions as we described in the material and methods section.
337 Our results showed that the initial wilting and curling of leaves appeared at 24 h and
338 the freezing injury symptoms became more severe at 72 h (**Figure 1a, upper panel**).
339 After cold treatment, seedlings were transferred to 25 °C and allowed to recover for 5
340 days. Bamboo seedlings recovered after 24 h cold treatment had reduced leaf
341 expansion and leaf wilting phenotypes, while seedlings recovered from 72 h- cold
342 treatment almost entirely lacking in the chlorophyll and the tissues began
343 wilting-to-death (**Figure 1a, lower panel**). The mortality ratio after 5 days' recovery
344 was 39 % for 24 h- cold treated plants and 69 % for 72 h- cold treated plants
345 respectively (**Supplemental Figure 1**). Results from scanning electron microscopy
346 (SEM) clearly demonstrated the collapsed and shrunken trichomes on the abaxial side
347 of the bamboo leaves after 24 h freezing treatment, which were even more obvious
348 after 72 h (**Figure 1b**).

349 To evaluate the cold-induced phenotypes, several abiotic stress related to the
350 biochemical parameters were measured. Electrolyte leakage reflects the degree of
351 membrane dysfunction caused by stress, and the increased conductivity is indicative
352 of more severe membrane damage (Whitlow et al. 1992). Our results showed the
353 relative electrolyte leakage of bamboo was increased dramatically with the
354 progression of cold treatment (**Figure 1c**). The MDA content exhibited a significant
355 increase after 24 h- and 72 h- cold treatment (**Figure 1c**). SOD, POD and APX work
356 as crucial enzymatic antioxidants to detoxify ROS (Abid et al. 2016; Reddy et al.
357 2004), and those antioxidant enzyme activities increased significantly after 24 h- and
358 72 h- cold treatment (**Figure 1c**). The proline content significantly increased at 24 h
359 and was maintained at a high level at 72 h (**Figure 1c**). Results from the
360 morphological physiological observations and biochemical assays showed that cold
361 stress at -2 °C was detrimental to Moso bamboo, and also indicated that the broad
362 change in gene expression happen within 24 h.

363

364 **Characterization of the cold-treated Moso bamboo transcriptome**

365 To provide a comprehensive profile of the transcriptome of Moso bamboo in
366 response to cold, we performed RNA-Seq analysis. To optimize the conditions for this
367 experiment, bamboo seedlings treated at -2°C were harvested at different time
368 points (0 h, 0.5 h, 1 h, 3 h, 6 h, 12 h, and 24 h) to analyze the expression patterns of
369 several cold responsive marker genes such as *PeCRPK1*, *PeCML*, *PeCBF3*, *PeMPK3*,
370 *PeZAT12*, *PePIF3*, *PeMKK4*, *PeICE1*, and *PeCOR47* (Kidokoro et al. 2017; Pareek et
371 al. 2017; Shi et al. 2015). Our results showed that in most cases, these genes were
372 responsive at 0.5 h or 1 h, and had the most significant change at 24 h (**Supplemental**
373 **Figure 2**). Therefore, samples from cold treatment under -2°C at 0 h, 0.5 h, 1h and 24
374 h were used for RNA sequencing. A total of 12 samples, including three biological
375 duplicates at each of the four time points were performed.

376 Illumina platform generated 269,435,030 raw reads. After filtering, 266,080,018
377 clean reads containing a total of 79.27 Gb clean nucleotides with 91.68% Q30 bases
378 (base quality > 30) were obtained through stringent quality assessment and data
379 filtering. The quality of the sequencing data is summarized in **Supplemental Table 1**.
380 The clean reads were mapped to the *P.edulis* genome using the HISAT2 tool. The
381 average mapping ratio ranged from 85.21% to 89.35%. Based on the read alignments,
382 StringTie was applied to transcript assembly (Pertea et al. 2015). After optimal gene
383 structure prediction and alternative splicing analysis, a total of 47,092 genes were
384 identified, with 15,105 (32.1%) new genes.

385 To validate and annotate the assembled transcriptome library, we searched against
386 the Non-redundant (Nr) peptide database, Swiss-Prot protein database, KOG and
387 KEGG, using BLASTx with a cutoff E-value of 10^{-5} . The results indicated that over
388 65% of the transcripts had significant similarity to at least one target from these
389 databases (**Supplemental Table 2**).

390

391 **Global change of the cold-responding transcripts in Moso bamboo**

392 The DEGs were determined as cold-responsive genes if the fold change in
393 expression levels was at least 1.5 fold change and the adjusted p -value ≤ 0.05 at any
394 time point compared to control using DESeq2. The DEGs identified at 0.5 h and 1 h

395 were defined as early responsive genes, and those that changed exclusively at 24 h
396 were regarded as the late responsive genes.

397 In total, 2,463 DEGs which cover 5.2% of all Moso bamboo genes were identified
398 under cold treatment, of which, 1,177 (47.8%) were up-regulated (**Figure 2a**,
399 **Supplemental Table 3**) and 1,286 (52.2%) were down-regulated (**Figure 2a**,
400 **Supplemental Table 4**). Our results demonstrated that 73 and 59 genes were
401 up-regulated at 0.5 h or 1 h respectively while 1,137 genes were up-regulated at 24 h
402 (**Supplemental Table 5, 6 and 7**). Among all the cold up-regulated genes, only 26
403 genes had increased expression levels at all the time points (**Supplemental Table 8**).
404 In the down-regulated category, only 16 genes were down-regulated at 0.5 h and 20
405 genes at 1 h, while 1,263 genes were down-regulated at 24 h (**Supplemental Table 9**,
406 **10 and 11**), with only 2 genes showing decreased expression at all time points
407 (**Supplemental Table 12**). The expressions levels of 1,072 up-regulated and 1,253
408 down-regulated genes were completely changed at 24 h. The comparison of the
409 number of early responding genes (132 up-regulated and 36 down-regulated genes)
410 suggested that the observed gene up-regulation may play a key role in the early
411 response to cold stress.

412 To visualize the expression patterns of these DEGs at early and late stages, a
413 heatmap was constructed on the basis of the fragments per kilobase of transcript per
414 million (FPKM) values (**Figure 2b**). DEGs with similar expression patterns were
415 grouped, and the heatmap results showed that most DEGs changed their expression
416 profile significantly at 24 h. Our data suggested that the majority of the cold-regulated
417 genes are late-response genes under our treatment conditions. This observation in
418 bamboo is similar to the previous reports in *Arabidopsis* which showed that most
419 induced or repressed genes appeared at 24 h cold treatment (Lee et al. 2005).

420

421 To further verify the RNA-seq data, we performed qPCR analysis for 12 selected
422 DEGs that are known to be related to cold stress, including *PeCML*, *PeCIPK1*,
423 *PeCRPK1*, *PeMKK4*, *PeMPK3*, *PeICE1*, *PeMYB15*, *PeZAT12*, *PePIF3*, *PeREV1*,
424 *PeWRKY40* and *PeCBF3*. The transcripts of those 12 genes showed similar

425 expression patterns to the results from RNA-seq (**Figure 2c**). These results support
426 the validity of the Moso bamboo cold-regulated transcriptome from the *in silico*
427 analysis.

428

429 **Functional annotation and classification of cold-regulated genes**

430 To characterize the functional classifications of the cold-regulated genes, 77.03 %
431 of up-regulated transcripts and 67.86 % of down-regulated transcripts were matched
432 to the Gene Ontology (GO) database, resulting in three categories being identified:
433 biological processes, cellular components, and molecular functions (**Supplemental**
434 **Figure 3a and 3b**). In each of the three main GO classifications, “Cellular process”,
435 “Cell” and “Binding” exhibited the highest match numbers among up-regulated and
436 down-regulated genes (**Supplemental Table 13 and 14**). We noticed a higher
437 percentage of genes from the “Rhythmic process” and “Positive regulation of
438 biological process” existed in the up-regulated genes compared with down-regulated
439 genes (**Supplemental Figure 3, Supplemental Table 13**). “Cell killing” and
440 “Detoxification” genes were only present among the up-regulated genes. On the other
441 hand, the “Growth” and “Cellular component organization or biogenesis” groups were
442 more abundant amongst down-regulated genes (**Supplemental Figure 3,**
443 **Supplemental Table 13 and 14**). This data suggested that cold stress might provoke
444 the expressions of genes involved in cell killing and detoxification processes, which is
445 consistent with the phenotypes observed and shown in **Figure 1**.

446 For further functional prediction and categorization, all cold-regulated genes were
447 subject to phylogenetic classification using the KOG database. The up-regulated and
448 down-regulated genes were assorted in 22 and 24 KOG categories, respectively
449 (**Supplemental Figure 4**). “Signal transduction mechanisms”, “General function
450 prediction only”, and “Posttranslational modification” genes were the most
451 represented categories (**Supplemental Table 15 and 16**). “Secondary metabolites
452 biosynthesis”, “Energy production and conversion”, and “Carbohydrate transport and
453 metabolism” were over-represented amongst the up-regulated genes compared to
454 down-regulated genes (**Supplemental Table 15**), which implied that genes involved

455 in carbohydrate and secondary metabolism pathways might contribute to cold
456 resistance processes. “Chromatin structure and dynamics”, “Replication,
457 recombination and repair” and “Cell cycle control, cell division, chromosome
458 partitioning” were more abundant in down-regulated genes compared to up-regulated
459 genes (**Supplemental Table 16**). In addition, “Nucleotide transport and metabolism”
460 and “Extracellular structures” were only identified among down-regulated genes
461 (**Supplemental Figure 4b, Supplemental Table 16**). These findings suggested that in
462 bamboo chromatin remodeling was crucial in regulating cold response, and also
463 hinted that the genes involved in the repair process and cell division were heavily
464 inhibited in response to cold stress.

465 To identify candidate metabolic pathways regulated by cold stress, the DEGs were
466 examined using the KEGG pathway analysis tool, which is a compilation of manually
467 verified pathway maps to categorize gene functions with the emphasis on biochemical
468 pathways (Minoru et al. 2008). A total of 65.23% up-regulated genes and 58.03%
469 down-regulated genes were mapped to the KEGG database. For both the up- and
470 down- regulated genes, the clusters for “Metabolism” and “Organismal systems” were
471 significantly enriched (**Supplemental Figure 5**). The cold-regulated genes were
472 further classified into five categories (**Figure 3**). Interestingly, within the
473 “Metabolism” category, “Global and overview maps”, “Carbohydrate metabolism”
474 and “Amino acid metabolism” genes were most highly represented amongst the
475 up-regulated genes (**Figure 3a**). Several genes encoding enzymes associated with
476 cysteine and methionine metabolism, including S-adenosylmethionine synthetase,
477 cystathionine gamma-synthase and adenosylmethionine decarboxylase were
478 specifically induced by cold stress (**Supplemental Table 17**). Moreover, a few genes
479 involved in flavonoid biosynthesis, including phenylalanine ammonia-lyase and
480 chalcone synthase were significantly increased, suggesting that Moso bamboo may
481 use flavonoids as antioxidants to prevent from ROS damage (**Supplemental Table**
482 **17**). Pathways related to “Environmental adaptation” were highly represented in
483 up-regulated genes (**Figure 3a**), with calcium- binding proteins being especially
484 notable (**Supplemental Table 17**). These findings indicated a role of calcium- binding

485 proteins as the main signaling components in cold signal transduction. Furthermore,
486 “Membrane transport” composed of ABC transporters were mainly represented in
487 up-regulated genes (**Figure 3a; Supplemental Table 17**). Several ABC transporters
488 were rapidly induced in Moso bamboo after cold exposure (**Supplemental Table 17**).
489 On the other hand, “Nucleotide metabolism”, “Replication and repair”, “Glycan
490 biosynthesis and metabolism” and “Lipid metabolism” were down-regulated more
491 during cold stress (**Figure 3b**). Many genes related to cell wall modification were
492 found in the category of “Glycan biosynthesis and metabolism” (**Supplemental Table**
493 **18**). Meanwhile, the genes involved in ‘Biosynthesis of unsaturated fatty acids’ and
494 ‘Fatty acid elongation’ were significantly inhibited by freezing treatment
495 (**Supplemental Table 18**), indicating a decrease in fatty acid content in the plasma
496 membrane.

497

498 **Expression profiles of cold-regulated genes**

499 The heatmap demonstrated that dynamic transcriptional changes in response to the
500 cold stress (**Figure 2b**). The identified DEGs were grouped into 12 clusters based on
501 the SOM cluster analysis using the *k*-means method (**Figure 4, Supplemental Table**
502 **19**). The KOG functional category was applied for each cluster to predict the
503 distribution of different functions among the three time periods. The most abundant
504 cluster was Cluster 10 with 573 DEGs induced immediately at 0.5 h and remained
505 up-regulated at 24h. The second most abundant cluster was Cluster 8, which was
506 comprised of 373 genes with significantly increased expression at 24 h. The third
507 most abundant group was Cluster 5, containing 292 genes with decreased expression
508 at 0.5 h and 24 h. The fourth most abundant group was Cluster 3 with 263 genes
509 showing decreased expression at 24 h (**Supplemental Table 19**).

510 Cluster 6 with 164 genes was rapidly induced at 0.5 h but not at later time points,
511 while cluster 1, 4 and 10 showed another peak at 24 h (**Figure 4**). Interestingly,
512 functional categories of “Transcription”, “Lipid transport and metabolism” and
513 “Inorganic ion transport and metabolism” were over-represented in Clusters 1, 6 and
514 10 (**Supplemental Table 20**). This indicated that cells received the cold signal and

515 instantly transmitted through the ion and lipid transport through transcription network
516 within 0.5-1 h cold treatment. The genes with functions of “Amino acid transport and
517 metabolism”, “Carbohydrate transport and metabolism”, and “Energy production and
518 conversion” were identified as up-regulated throughout the 1 - 24 h stage (**Figure 4,**
519 **Supplemental Table 20**), reflecting the downstream metabolic processes activated by
520 signal transduction and transcription in response to cold. Clusters 2, 3, 5, 7 and 11
521 contained 1234 down-regulated genes at 24 h, while cluster 2, 5 and 7 responded to
522 the cold stress rapidly in the early stage (0-0.5 h) by decreasing their expression
523 (**Figure 4**). Functional categories of “Nucleotide transport metabolism”, “Chromatin
524 structure and dynamics”, “RNA processing and modification” were more enriched in
525 those clusters (**Figure 4, Supplemental Table 20**), as shown by those processes
526 appearing to be rapidly negatively regulated by cold stress. In summary, our results
527 indicated that genes responded to cold stress in a hierarchical manner in bamboo.

528

529 **Transcription factors responding to cold stress**

530 Transcription factors play important roles in mediating cold stress related gene
531 expressions (Lee et al. 2005; Zhang et al. 2014). In this study, we identified 222
532 transcription factors from 24 different families, which were differentially expressed
533 throughout the 24 h cold stress (**Figure 5a; Supplemental Table 21**). A total of 111
534 up-regulated transcription factors were identified from 19 different families/groups
535 (**Figure 5b, Supplemental Table 22**). The most up-regulated transcription factors
536 constituted key families that are cold-sensitive, such as APETALA2, ethylene
537 response factors (AP2/ERF), WRKY transcription factors (WRKY), NAC
538 domain-containing proteins (NAC), and basic leucine zipper transcription factors
539 (bZIP) (**Figure 5b, Supplemental Table 22**). The number of down-regulated
540 transcription factors was comparable to that of up-regulated transcription factors,
541 which consisted of 111 genes from 22 families, which were mainly from MYB,
542 homeodomain-leucine zipper transcription factor (HD-ZIP), and B3
543 domain-containing transcription factor (or B3) families (**Figure 5b, Supplemental**
544 **Table 23**).

545 The expression changes of cold responsive transcription factors were illustrated by
546 heatmap analysis, and those with similar expression patterns were categorized
547 (**Figure 5c**). Some transcription factors were induced immediately after the plants
548 were exposed to cold stress, while others were down/up-regulated subsequently,
549 suggesting that a transcriptional cascade triggered by cold stress might be present in
550 Moso bamboo. The analysis highlighted the expression changes of several
551 well-known cold-regulated transcription factors during 24 h cold treatment. For
552 example, *PeCBF3* and *PeWRKY33* responded rapidly to cold treatment at 0.5 h, and
553 their expression levels also increased again at 24 h. The expression level of *PeREVI*
554 and *PeMYB15* increased at 24 h (**Supplemental Table 21**). In addition, the expression
555 of the *PeWRKY40* and *PeZAT12* increased immediately at 0.5 h and maintained a
556 positive slope until 24 h cold treatment (**Supplemental Table 21**), which is consistent
557 with a previous finding that these proteins serve as the markers for early cold response
558 in Arabidopsis (Lee et al. 2005).

559

560 **Expression patterns of selected DEGs in two bamboo species with different cold** 561 **tolerant abilities**

562 Since no Moso bamboo genetic transformation method was available at the time,
563 we could not verify the function of DEGs from our RNA seq results through gene
564 modification method. Alternatively, the gene function might be deduced by comparing
565 the expression patterns in different bamboo populations with different cold-tolerance
566 abilities. Ma bamboo (*Dendrocalamus latiflorus* Munro) is a more cold-sensitive
567 bamboo species compared to Moso bamboo (Liu et al. 2006). To examine the
568 expression patterns of the cold stress-induced genes from RNA seq data, Ma bamboo
569 and Moso bamboo were exposed to low temperature over 24 h and the gene
570 expression patterns were examined by qPCR. Six representative DEGs including two
571 putative positive regulators (*PeCBF3* and *PeCBF4*) and four putative negative
572 regulators (*PeMYB15*, *PePIF3*, *PeZAT12*, and *PeCRPK1*) in cold signaling pathways
573 were selected for this experiment. Our results demonstrated that all genes tested were
574 cold responsive (**Supplemental Figure 6**). More importantly, compared with the less

575 cold-tolerant Ma bamboo, Moso bamboo has higher expression levels of putative
 576 positive regulators and lower expressions of negative regulators in cold signaling
 577 pathway (**Supplemental Figure 6**). These data reflected the effectiveness of our
 578 RNA-Seq results, and support functional significance of DEGs in bamboo cold
 579 signaling pathways.

580

581 **Discussion**

582 **Calcium signaling pathway and MAPK cascades were activated in the early** 583 **stage of the bamboo cold response**

584 We noticed that various Ca^{2+} sensor genes including *PeCaM/CP1*, *PeCML*,
 585 *PeCDPKs* (*PeCDPK19* and *PeCDPK5*), *PeCRPK1*, and *PeCIPK1* were significantly
 586 induced in response to cold stimulus (**Supplemental Table 17**). Our data also
 587 indicated that genes encoding Ca^{2+} binding proteins responded to cold treatment
 588 within 30 min (**Figure 2c**), which was consistent with previous findings that Ca^{2+}
 589 binding proteins rapidly transduced external signals (Kudla et al. 2018). Moreover,
 590 our data demonstrated that the *MAPK* cascades, such as *PeMKK9*
 591 (*PH01003362G0140*), *PeMKK4*, *PeMPK20* (*PH01000298G0100*), and *PeMPK3*
 592 were activated in the early stage of cold treatment (**Figure 2c and Supplemental**
 593 **Table 17**). These data supported previous findings that the activation of the *MAPK*
 594 cascade could be triggered by cold stress, probably due to the over-accumulation of
 595 ROS and MDA content (Zhang et al. 2014).

596 Our data supported the hypothesis that Ca^{2+} and *MAPK* signal transduction
 597 pathways were activated at an early stage when bamboo was challenged with cold
 598 stress to induce downstream gene expression and protect the plant cells. The data are
 599 consistent with previous results from Arabidopsis and rice (Lee et al. 2005; Zhang et
 600 al. 2014). Moreover, we also noticed that several important negative regulators in
 601 Arabidopsis or rice might play opposite roles in bamboo during cold response. For
 602 example, it was reported that Arabidopsis *CRPK1* functions as a negative regulator
 603 through the *CBF* pathway (Liu et al. 2017), and another key regulator, *MPK3*,

604 negatively regulates *ICE1* expression through post-translational modification (Li et al.
605 2017). Interestingly, the expressions of both orthologs in Moso bamboo were
606 enhanced in response to cold treatment, indicating the presence of the bamboo
607 species-specific control mechanisms in response to cold stress.

608

609 **Transcription factors response to cold stress in both plant kingdom-conserved** 610 **and species- specific mechanisms in Moso bamboo**

611 A large number of transcription factors belonging to different transcription factor
612 families have been shown to play a crucial role in regulating the cold response in
613 Arabidopsis (Lee et al. 2005), rice (Zhang et al. 2014), wheat and many other plant
614 species (Calzadilla et al. 2016; Kargiotidou et al. 2010; Wang et al. 2014; Weiss and
615 Egea-Cortines 2009). Here, we identified 222 transcription factors from 24 different
616 gene families responding to cold stress. Among them, MYB, AP2/ERF, WRKY, ZIP
617 families comprise a high proportion of cold-responsive members (**Figure 5a**). We
618 investigated the classical *CBF* regulation pathway including the upstream regulators,
619 such as *PeICE1*, *PeMYB15*, *PeZAT12* and *PePIF3*. Interestingly, *PeICE1*, *PeZAT12*,
620 *PeMYB15* and *PePIF3* were induced rapidly by cold stress at 0.5 h (**Figure 2c**), which
621 demonstrated the effectiveness of our treatments and confirmed the important roles of
622 transcription factors in the early cold response in bamboo. *ICE1* is a key positive
623 regulator of *CBF3* (Chinnusamy et al. 2003), while *ZAT12*, *MYB15* and *PIF3* are all
624 negative regulators of *CBF* genes (Agarwal et al. 2006; Jiang et al. 2017; Novillo et al.
625 2007). The combination of the regulation of *PeICE1*, *PeZAT12*, *PeMYB15* and
626 *PePIF3* could explain the fluctuation of *PeCBF3* expression. Previous studies showed
627 that 8 *WRKYs* display increased expression in the early cold respond response
628 in Arabidopsis (Lee et al. 2005). We found that the enhanced expression of two
629 *PeWRKYs* (*PeWRKY40* and *PeWRKY33*) occurred at an early stage in response to
630 cold stress in Moso bamboo (**Figure 2c and Supplemental Table 21**), which
631 suggested that the *PeWRKYs* might have conserved roles in the cold response in
632 bamboo. We also identified *PeREVI*, whose ortholog in Arabidopsis works as a
633 negative regulator of cold acclimation (Meissner et al. 2013), showing increased

634 expression during the cold treatment. The expression patterns of important regulators
 635 of the *CBF* pathway could explain the cold-sensitive phenotypes of Moso bamboo via
 636 the repression of *PeCBF3*.

637 It is worth noting that except for the *CBF* pathway, other cold stress regulatory
 638 pathways also play a role in plant cold response (Fowler and Thomashow 2002; Kreps
 639 et al. 2002; Monroy et al. 2007; Tian et al. 2013). For example, it was reported that at
 640 least 28% of the cold-responsive genes were not regulated by the *CBF* pathway in
 641 *Arabidopsis* (Fowler and Thomashow 2002). Furthermore, at least one-third of the
 642 cold-inducible genes in wheat were independent of the *CBF* pathway (Monroy et al.
 643 2007). In Moso bamboo, 40 transcription factors were induced immediately at the
 644 transcriptional level upon exposure to cold stress (**Supplemental Table 24**). The early
 645 induced transcription factors, which were closely clustered with known transcription
 646 factors such as *PeWRKY33*, *PeCBF3* and *PeMYB15*, warrant further investigation to
 647 identify a potential *CBF* parallel pathway. For example, the up-regulation of HD-ZIP
 648 transcription factor (*PH01001036G0340*), the NAC transcription factor
 649 (*PH01001177G0140*) and the B3 transcription factor (*PH01000246G0410*) within 0.5
 650 h by cold stress, indicate that those regulators play an important function in cold
 651 stress.

652

653 **The transcriptomic profiles of cell wall related genes changes in response to cold** 654 **stress**

655 Plant cell walls play a structural role in plant abiotic stress defenses (Tenhaken
 656 2015). It has been proposed that increasing the amount of pectin could efficiently
 657 delay plant cell damage by forming hydrated gels (Leucci et al. 2008). Interestingly,
 658 our data revealed that three genes involved in the biosynthesis of pectin, *PeGAE1*
 659 (*PH01000119G0710*) encoding a UDP-D-glucuronate 4-epimerase, *PeRHM1*
 660 (*PH01001109G0280*) encoding a UDP-L-Rhamnose synthase and *PeGATL9*
 661 (*PH01000092G1070*) encoding a galacturonosyl transferase, were highly induced
 662 during the cold treatment (**Supplemental Table 17**). These results suggested that
 663 these genes might function not only in cell wall metabolism but also as candidates for

664 the plant adaption to cold stress in Moso bamboo. On the other hand, we showed
665 several key genes affecting cell wall integrity through xylan modification, such as a
666 1,4-beta-D-xylan synthase (*PeIRX10*, *PH01000002G2800*), two glycosyl transferases
667 (*PeIRX9*, *PH01000428G0570* and *PeIRX9L*, *PH01000256G1170*), and a
668 plant-specific *PeDUF231* (*PeTBL27*, *PH01001319G0100*), were dramatically
669 down-regulated at 24 h (**Supplemental Table 18**). As xylan is the major component
670 of hemicelluloses in the plant cell wall, reduced expression of xylan biosynthesis
671 genes leads to the weakening of the secondary cell wall, resulting in the collapse of
672 xylem vessels (Brown et al. 2007; Lin et al. 2016). Our data indicates cell wall-related
673 genes play important roles in the acquisition of cold tolerance by changing their
674 expression patterns at the transcription level.

675

676 **Lipid metabolism was inhibited under cold stress**

677 It was documented that the most damaging effect of cold stress in plants is plasma
678 membrane damage from dehydration (Steponkus 1993). The cold-treated bamboo
679 displayed obvious dehydration phenotypes, such as wilting (**Figure 1a**) and ruptured
680 trichomes on the leaf surface (**Figure 1b**). Damage of the plasma membrane was
681 demonstrated in terms of increased ion leakage, which implies increased membrane
682 permeability and reduced cell tolerance to low temperature (**Figure 1c**). Furthermore,
683 the increased content of MDA indicated the oxidation of the unsaturated membrane
684 fatty acids (**Figure 1c**). Increased accumulation of unsaturated fatty acids in the
685 plasma membrane improve cold defense by preventing ion leakage (Degenkolbe et al.
686 2012). Based on the KEGG metabolic pathway analysis, the ‘Biosynthesis of
687 unsaturated fatty acids’ and ‘Fatty acid elongation’ clusters were significantly
688 enriched in down-regulated genes (**Supplemental Table 18**). In particular, genes
689 involved in unsaturated fatty acids biosynthesis including an acyl-CoA dehydratase
690 (*PePAS2*, *PH01001117G0220*), a stearyl-ACP desaturase (*PeFAB2*,
691 *PH01001326G0300*), and a NADP-binding protein
692 (*Phyllostachys_edulis_newGene_23189*) were down-regulated at 24 h (**Supplemental**
693 **Table 18**). Several genes involved in cuticle membrane and wax biosynthesis, such as

694 *PeCER3* (*PH01000379G0490*), *PeMYB106* (*PH01005515G0070*), *PeKCS4*
695 (3-ketoacyl-CoA synthase, *PH01000046G1090*), *PeKCS5* (*PH01001011G0290*) and
696 *PeKCS6* (*PH01000101G1030*) were also down-regulated at 24 h (**Supplemental**
697 **Table 18 and 21**). Our data showed that the decreased expression of genes involved
698 in the biosynthesis of fatty acids in response to cold would explain the impaired cold
699 defense and cold-sensitive phenotypes, which are consistent with previous findings
700 (Degenkolbe et al. 2012; Shepherd and Griffiths 2006).

701

702 **Phytohormones play important roles in plant cold-stress response**

703 The crucial roles of plant hormones in the plant cold stress response have been well
704 demonstrated (Shi et al. 2015). Of the DEGs we identified, 71 genes were involved in
705 ABA-, ethylene-, GA- and auxin- related pathways (**Supplemental Table 25**). In cold
706 treated bamboo, expressions of genes involved in ABA biosynthesis, signal reception
707 and downstream signaling pathways were changed. A putative *NCED* gene
708 (*PH01000283G0010*) which encodes a key enzyme in ABA biosynthesis pathway,
709 and a PYR/PYL/RCAR family protein (*PH01002424G0210*), that functions as an
710 ABA sensor, had enhanced expression in the cold-treated bamboo seedlings. In plants,
711 members of the protein phosphatase 2C (PP2C) family may act as positive regulators
712 within ABA-mediated signaling networks activated by diverse environmental stresses
713 or developmental signaling cascades (Xue et al. 2008), with two PP2C family genes
714 (*PH01001115G0280* and *PH01004966G0010*) displaying increased their expression
715 levels after cold treatment in bamboo. A total of 24 putative ABA responsive genes
716 with 17 being up-regulated and 7 being down-regulated were identified in this study
717 (**Supplemental Table 25**), suggesting that ABA related pathways participated in
718 bamboo cold response. The role of ethylene in plant response to cold is different in
719 different species (Kazan 2015). We noticed the down-regulation of a putative *ETO1*
720 (ethylene over-producer) paralog (*PH01000367G0090*) after cold treatment, which
721 acts as a negative regulator of *ACS5* (1-aminocyclopropane-1-carboxylate synthase 5,
722 a key enzyme in ethylene biosynthesis pathway), indicating the enhanced level of
723 ethylene in bamboo after cold treatment. Ethylene signaling pathway also affects plant

724 cold tolerance. In cold treated bamboo, at least 12 ethylene responsive factors (ERF)
725 changed their expression patterns, with 9 being up-regulated and 3 being
726 down-regulated (**Supplemental Table 25**). These results suggest ethylene pathways
727 might be activated during the bamboo cold response. A key response to cold in plants
728 is growth repression, to allow the plant to re-allocate resources from growth to
729 processes responsible for increasing cold tolerance (Eremina et al. 2016). Gibberellins
730 and auxin are well known growth-promoting hormones, and our results indicate their
731 potential roles in the bamboo cold response. In the cold-treated seedlings, one GA
732 2-oxidase gene (*PH01001124G0470*) which deactivates gibberellins was up-regulated
733 (**Supplemental Table 25**), indicating GA homeostasis was required under cold stress
734 in bamboo. The growth hormone auxin essentially regulates all aspects of plant
735 developmental processes under both normal and abiotic stress conditions. The effect
736 of cold stress on auxin is linked to the inhibition of intracellular trafficking of auxin
737 efflux carriers (Rahman 2013). In accordance with these findings, the putative auxin
738 influx carrier *PeLAX2* (*PH01000484G0740*) and efflux carrier *PePIN1*
739 (*PH01000484G0740*) had reduced expression levels in bamboo (**Supplemental Table**
740 **25**). Moreover, a putative AGCVIII kinase (*PH01000023G1420*), which positively
741 activates the PIN-mediated auxin efflux by affecting cell trafficking (Willige and
742 Chory 2015), was also down-regulated (**Supplemental Table 25**). Recently, it was
743 found that abiotic stress-induced growth inhibition involves repression of auxin
744 responsive genes (Shani et al. 2017). Our results align with these findings, since we
745 found that multiple early auxin-responsive genes had altered expression levels in
746 bamboo. For example, a putative transcriptional repressor *AUX/IAA*
747 (*PH01000025G1630*) increased its expression, and 5 auxin-inducible small auxin up
748 RNA genes (*SAUR*) had reduced expression levels (**Supplemental Table 25**),
749 indicating the inhibition of the auxin pathways. Those results suggested that cold
750 stress affected auxin effects mainly through disrupting its transport and signaling
751 pathway in bamboo. In summary, our results indicated that the phytohormone
752 functions in the bamboo cold response, while the detailed precise mechanisms behind
753 this action need to be further investigated.

754

755 **Hypothetical model occurring in leaves of Moso bamboo upon cold stress**

756 A previous transcriptome study in *Arabidopsis* indicated that 3.9% of all
757 *Arabidopsis* genes were cold stress response genes (Lee et al. 2005). The majority
758 (74%) of the cold-response genes were late-response genes, which only displayed
759 altered expression levels only after 24 h of cold treatment. The significant
760 transcriptomic changes at the late stage were related to primary and secondary
761 metabolism and photosynthesis (Lee et al. 2005). The early-response genes were
762 mainly identified as transcription factors and the genes related to hormone
763 biosynthesis and signaling (Lee et al. 2005). A study of transcriptome reprogramming
764 in cold acclimation of tomato indicated that the early changes in expression are
765 mainly associated with transcription factors. In contrast, the late response that took
766 place after 24 h of cold exposure caused changes in expression of genes involved in
767 metabolism and machinery associated with protein translation (Barrero - Gil et al.
768 2016). Based on the results presented in this study, we propose a model for the cold
769 signal perception and responsive pathways in Moso bamboo (**Figure 6**). According to
770 this model, freezing temperatures are rapidly recognized through calcium signaling
771 pathways, *MAPK* cascades and other pathways such as ABA signaling cascades.
772 These signaling pathways stimulate transcriptional reprogramming including
773 CBF-dependent or CBF-independent pathways that subsequently trigger a complex
774 series of metabolic activities, including antioxidant production, cell wall composition
775 adjustment and lipid metabolism alteration. Notably, several negative regulators of
776 cold tolerance, such as *PeREV1*, *PePIF3*, *PeMYB15* and *PeZAT12* were effectively
777 up-regulated during the early stage of cold stress. This specific expression pattern is
778 speculated to be responsible for the cold-sensitive phenotypes of Moso bamboo. The
779 findings in this study will contribute to the elucidation of the molecular mechanisms
780 underlying the low-temperature response, which could significantly contribute to
781 improving cold tolerance in Moso bamboo.

782

783 **Conclusion**

784 In this study, we demonstrated the physiological and biochemical changes that
785 occur in Moso bamboo in response to cold stress. The genome-wide transcriptome
786 analysis shed light on the DEGs involved in cold regulation. We found that the Ca²⁺
787 signaling pathway and MAPK cascades responded rapidly to cold stress. Additionally,
788 transcription factors involved in the key signaling pathways in response to cold stress
789 were identified in this study. Moreover, our results demonstrated that the expression
790 of genes involved in various metabolism pathways, such as secondary metabolites
791 biosynthesis and lipid metabolism, were altered during cold treatment, revealing the
792 potential role of these genes in cold defense. The results from this study provided
793 information to further elucidate of the possible functions of cold responsive genes in
794 bamboo. In the future, more experimental and bioinformatics work will be needed to
795 reveal the functions of these important candidates in this important species.

796

797 **Availability of data and materials**

798 RNA-Seq in this study had been submitted to GEO under accession number
799 GSE130314.

800

801 **Supplemental Material**

802 Supplemental Figures and Tables are listed

803

804 **Conflict of interests**

805 The authors declare that they have no conflict of interests.

806

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821

822 **Authors' contributions**

823 Q.Z., and Y.L. conceived this project; C.T.L., Y.S.Z., Y.L. and Q.Z. designed
824 experiments and interpreted the results. Y.L., C.W., H.X. and H.G. performed the
825 experiments; Y.W., S.C., H.L., and G.W.L. helped to collect and analyze the data. Y.L.
826 and Q.Z. wrote the manuscript. All authors read and approved the submission of this
827 manuscript.

828

829

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1097

1098 **Figure 1. Effects of freezing stress on Moso bamboo seedlings**

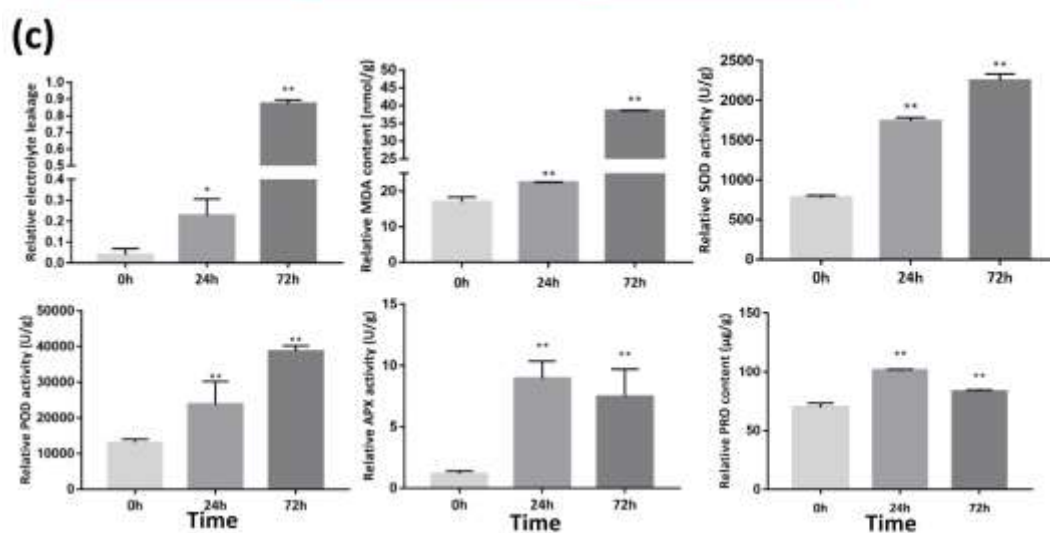
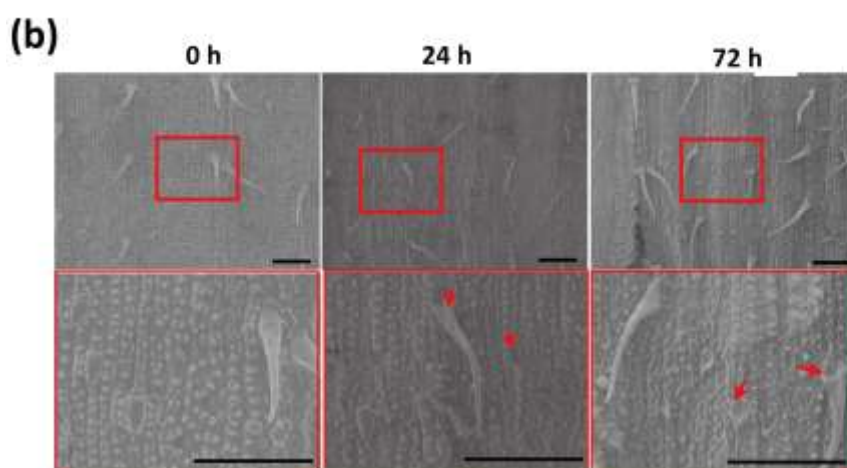
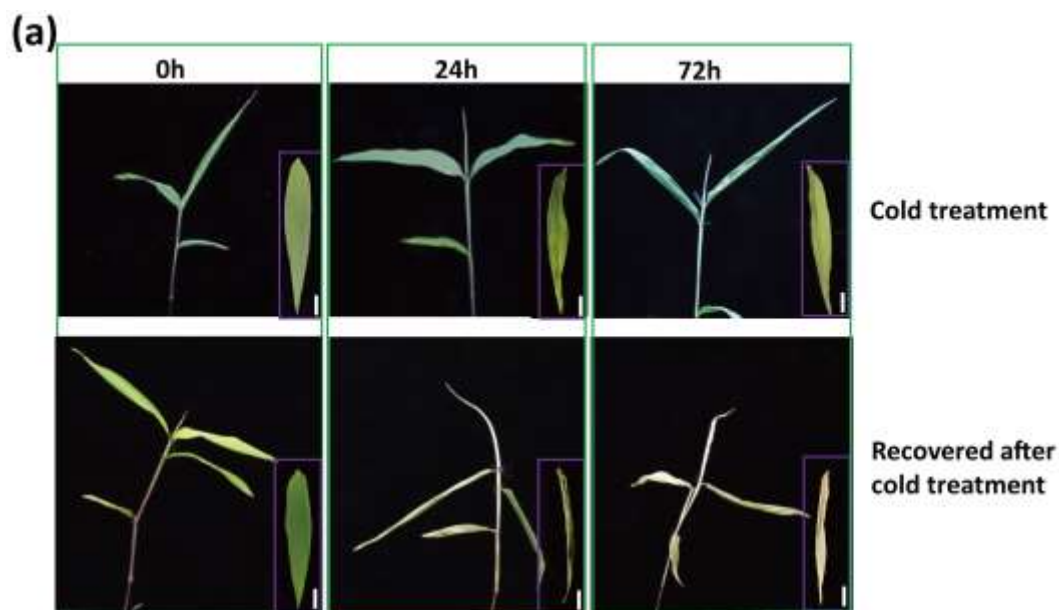
1099 (a) Upper Panel: Morphological changes of bamboo seedlings after cold treatment.
1100 3-week-old Moso bamboo seedlings that were exposed to -2°C for 24 h and 72 h
1101 respectively. Pictures highlighted in boxes show a closer view of the unstressed and
1102 freezing stress-exposed leaves. Bars = 1cm;

1103 Lower Panel: Morphological changes of bamboo seedlings recovered after cold
1104 treatment. 3-week-old Moso bamboo seedlings were exposed to -2°C for 24 h or 72 h,
1105 and then allowed to recover at normal growth temperatures for 5 days. Pictures
1106 highlighted in boxes show a closer view of the unstressed and freezing stress-exposed
1107 leaves. Bars = 1cm;

1108 (b) Scanning electron microscopy images of the lower surfaces of Moso bamboo
1109 leaves showing the collapse of trichomes due to the freezing treatment. The lower
1110 panel shows higher magnification images of the red boxed area in upper panel.
1111 Arrows indicate ruptured trichomes. Bars = 100 μm .

1112 (c) Measurements of physiological and biochemical parameters reflecting damage of
1113 Moso bamboo leaves. Values are means from three replications and error bars
1114 represent the standard deviations. Asterisks indicate significant differences from 24 h
1115 and 72 h to 0 h based on Student's t test data. Statistically significant differences were
1116 indicated by: *, $p < 0.05$, **, $p < 0.01$. All the measurements were performed at least
1117 three times with similar results and representative data from one repetition were
1118 shown.

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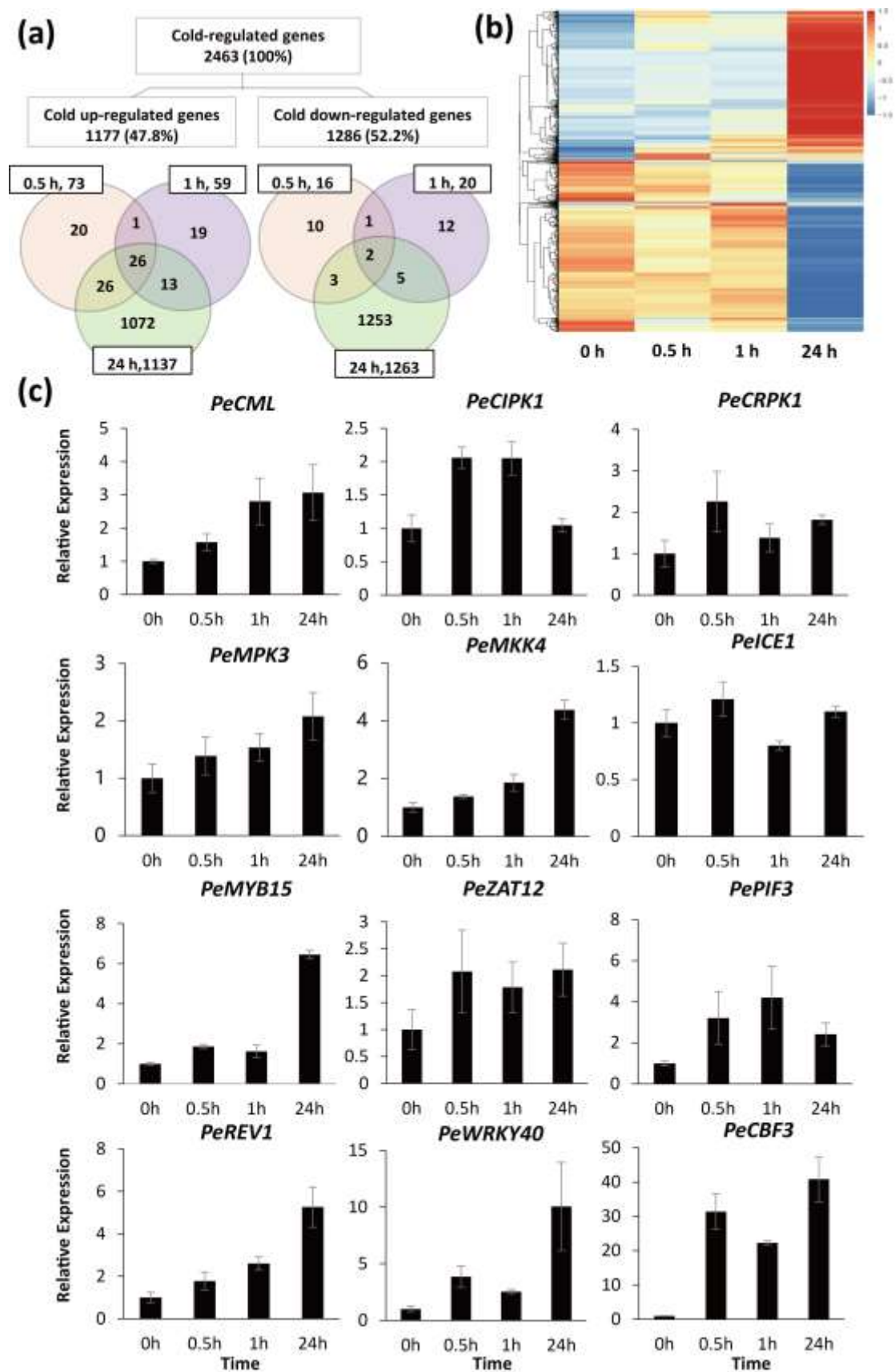
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1124 **Figure 2. Overview of the differentially expressed genes in response to cold stress**
1125 **in Moso bamboo**

1126 (a) Venn diagrams of cold-regulated genes. Figures in rectangles indicate cold
1127 treatment hours (h) and total number of cold-regulated genes at each time point.

1128 (b) Heat map of RNA-Seq transcriptome analysis for 2463 DEGs. Columns and rows
1129 in the heat map represent samples and genes, respectively. Sample names are
1130 displayed below the heat maps. The color bar is the scale for the expression levels of
1131 each gene. (c) Real-time PCR analysis of 12 selected genes in Moso bamboo. Data
1132 represents the average of three independent experiments \pm Standard Error (SE).



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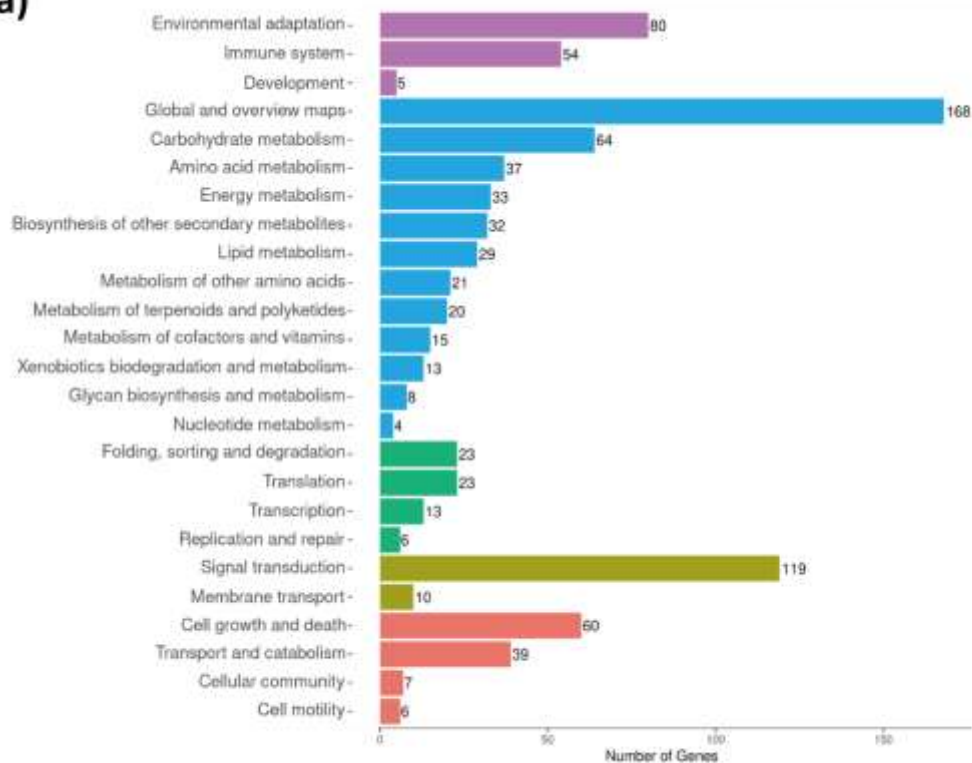
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1136 **Figure 3. Classification of Moso bamboo cold stress responsive genes for each**
1137 **KEGG category**

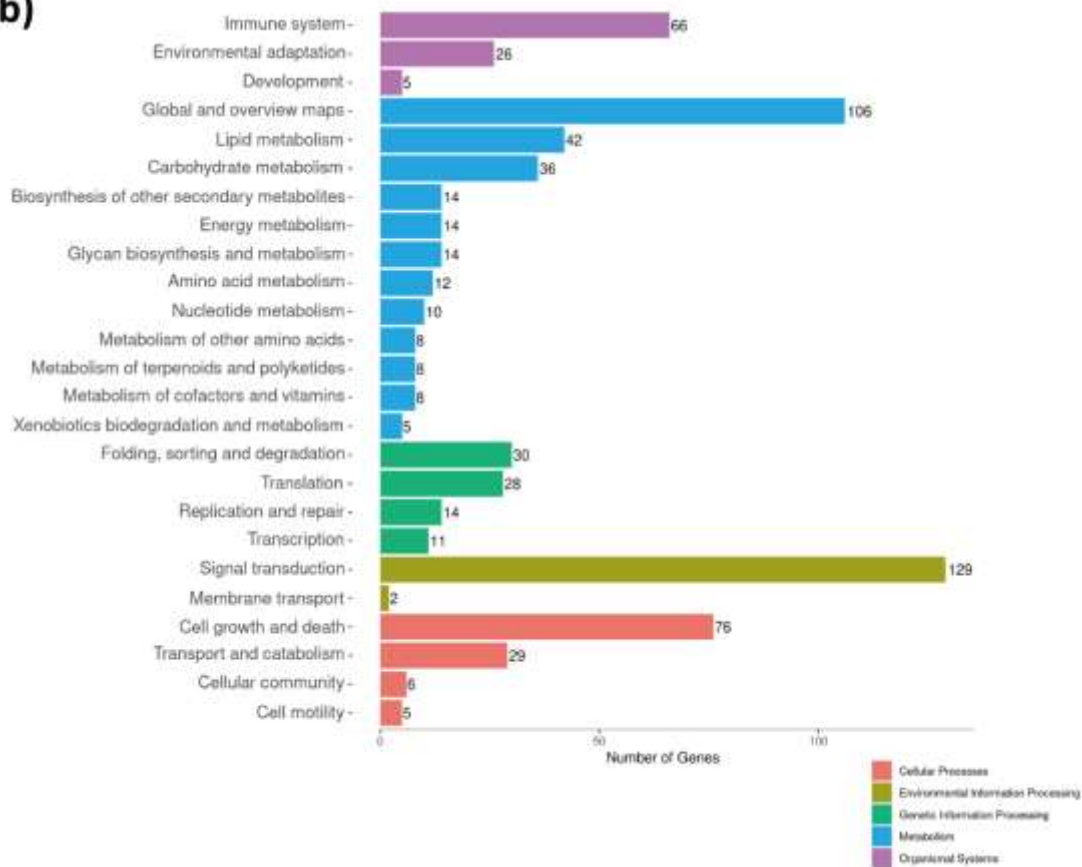
1138 The results are summarized in five categories: Cellular processes, Environment
1139 information processing, Genetic information processing, Metabolism and Organismal
1140 systems. The x-axis indicates the number of genes in a category.

1141 (a) Up-regulated genes; (b) Down-regulated genes.

(a)



(b)



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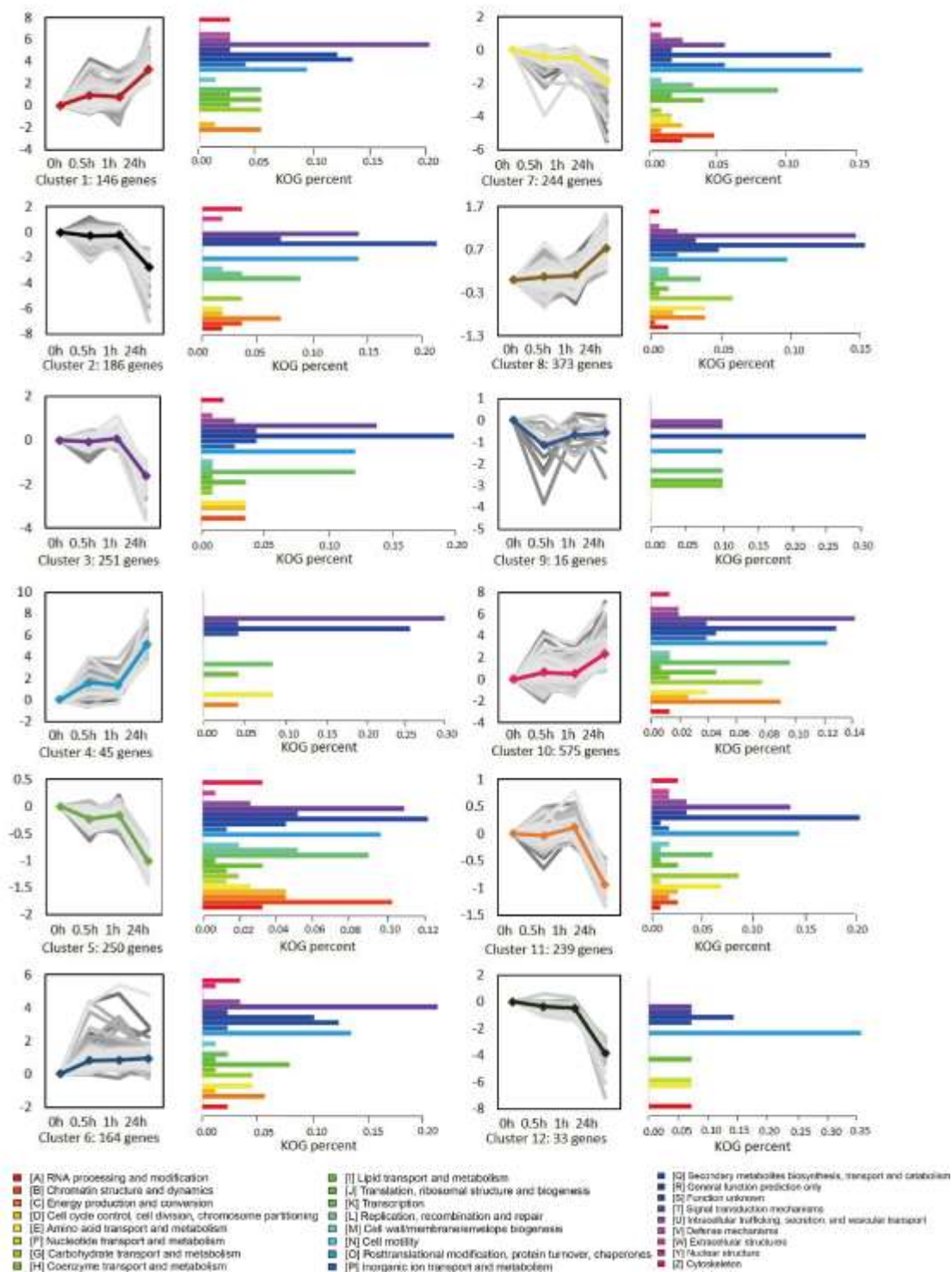
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1145 **Figure 4. Self-Organizing Maps (SOM) cluster analysis of DEGs in 12 different**
1146 **patterns**

1147 Clusters were obtained by the *k*-means method using the gene expression profiles of
1148 the 2,463 DEGs. The y-axis on the left side indicates the absolute value of $\log_2(\text{FC})$.
1149 KOG analysis was applied to each cluster. The x-axis on the right side represents the
1150 percentage of genes in a category.

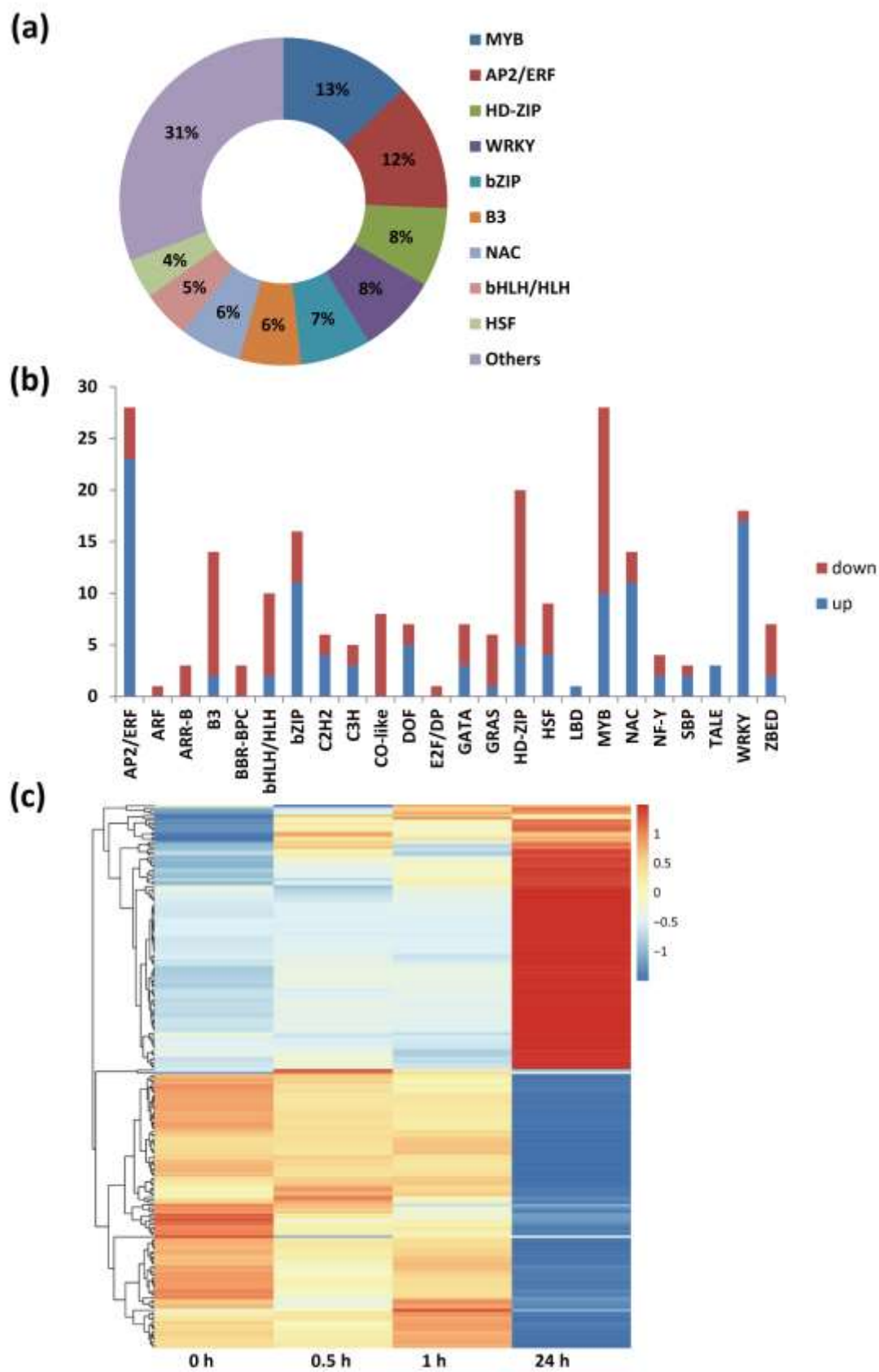
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1159 **Figure 5. Classification of cold regulated transcription factors**

1160 (a) The pie chart presents 222 transcription factors sorted into 24 different families;
1161 (b) The distribution of transcription factors in up- and down- regulated categories; (c)
1162 The heat map representing 222 differentially expressed transcription factors. Columns
1163 and rows in the heat map represent samples and genes, respectively. Sample names
1164 are displayed below the heat maps. The color bar is the scale for the expression levels
1165 of each gene.

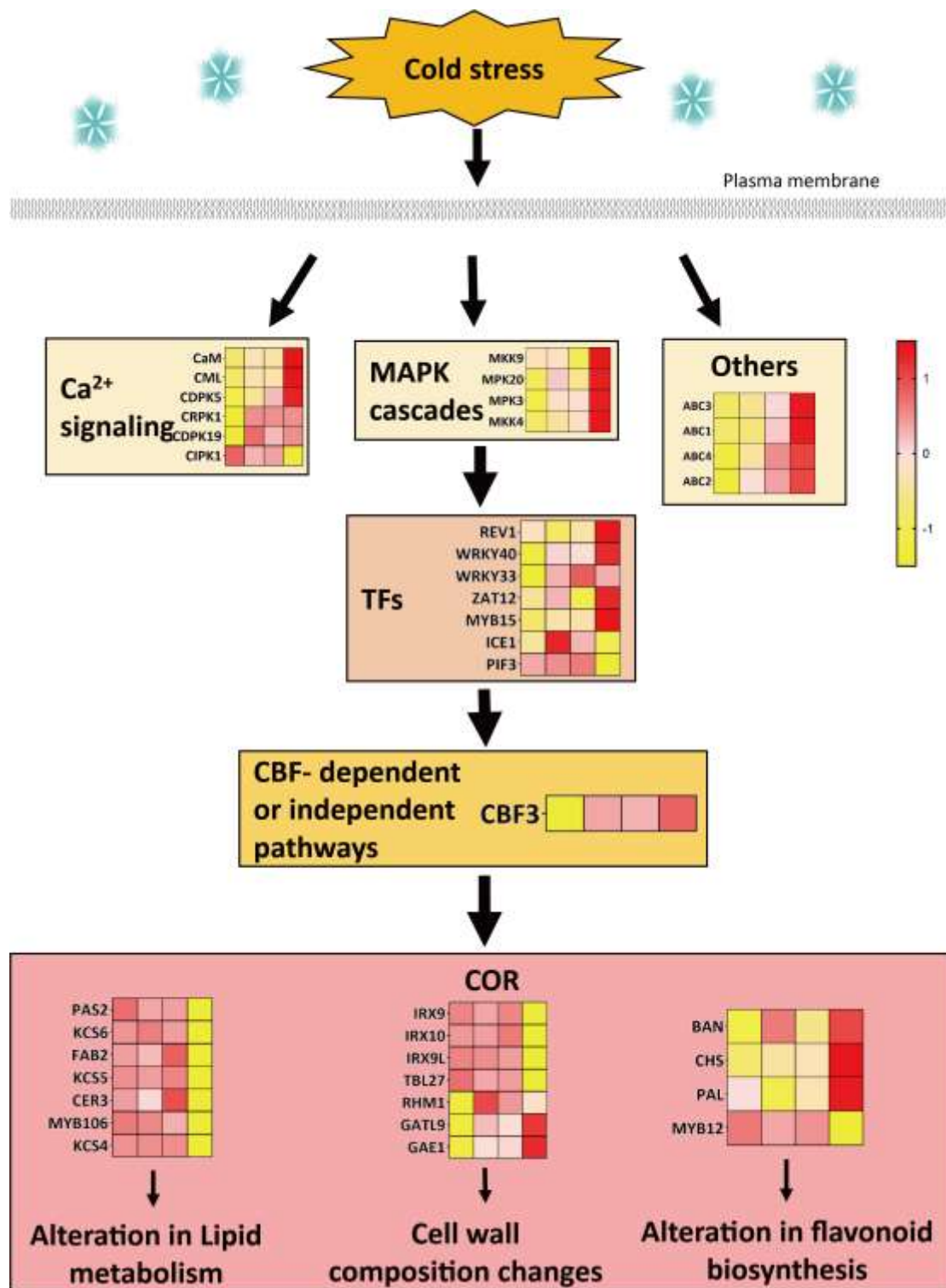


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1169 **Figure 6. A model of the cold response mechanism in Moso bamboo**
1170 Cold stress is rapidly recognized through calcium signaling pathway, MAPK cascades
1171 and other pathways such as ABA signaling cascades. The signaling pathways
1172 stimulate a transcriptional cascade triggered by the CBF-dependent or
1173 CBF-independent pathways. The late responsive genes are associated with a host of
1174 metabolic activities, including antioxidants production, cell wall composition
1175 adjustments and alternations in lipid metabolism. Genes were labeled using individual
1176 heatmaps. The color bar is the scale for the expression levels of each gene on the basis
1177 of FPKM value.



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