1	Motif analysis in co-expression networks reveals regulatory elements
2	in plants:
3	The peach as a model
4	
5	Running title: In silico prediction of peach regulatory motifs
6 7	Najla Ksouri <sup>1</sup> , Jaime A. Castro-Mondragón <sup>2,3</sup> , Francesc Montardit-Tardà <sup>1</sup> , Jacques van Helden <sup>2</sup> , Bruno Contreras-Moreira <sup>4,5,6*</sup> and Yolanda Gogorcena <sup>1*</sup>
8 9 10	<sup>1</sup> Laboratory of Genomics, Genetics and Breeding of Fruits and Grapevine, Estación Experimental de Aula Dei-Consejo Superior de Investigaciones Científicas, 50059 Zaragoza, Spain.
11 12	<sup>2</sup> Aix-Marseille Univ, INSERM UMR_S 1090, Theory and Approaches of Genome Complexity (TAGC), F-13288 Marseille, France.
13 14	<sup>3</sup> Current address: Centre for Molecular Medicine Norway (NCMM), Nordic EMBL Partnership, University of Oslo, 0318 Oslo, Norway.
15 16 17	<sup>4</sup> Laboratory of Computational and Structural Biology, Department of Genetics and Plant Production, Estación Experimental de Aula Dei–Consejo Superior de Investigaciones Científicas, 50059 Zaragoza, Spain.
18	<sup>5</sup> Fundación ARAID, Zaragoza, Spain
19 20	<sup>6</sup> Current address: European Molecular Biology Laboratory, European Bioinformatics Institute, Wellcome Genome Campus, Hinxton, Cambridge CB10 1SD, UK
21	*Senior authors
22	One sentence summary
23 24 25	Motifs prediction depends on the promoter size. A proximal promoter region defined as an interval of -500 bp to +200 bp seems to be the adequate stretch to predict <i>de novo</i> regulatory motifs in peach
26	Footnotes:
27	Authors' contribution
28	NK, BC-M and YG devised the study objectives, designed the experiment, discussed
29	data and wrote the manuscript. NK performed the bioinformatics analysis, FM-T
30	contributed to delimit the proximal promote region. JA-CM aided to prepare the figures

31 and provided critical feedback. JvH contributed in the critical discussion of results. YG

and BC-M contributed the analysis tools and YG conceived the experiment and
 supervised the activities. All authors read and approve the manuscript.

### 34 Fundings

This work was partly funded by the Spanish Ministry of Economy and Competitiveness grants AGL2014-52063R, AGL2017-83358-R (MCIU/AEI/FEDER/UE); and the Government of Aragón with grants A44 and A09\_17R, which were co-financed with FEDER funds. N. Ksouri was hired by project AGL2014-52063R and now funded by a PhD fellowship awarded by the Government of Aragón.

#### 40 Abstract

Identification of functional regulatory elements encoded in plant genomes is a 41 42 fundamental need to understand gene regulation. While much attention has been given to model species as Arabidopsis thaliana, little is known about regulatory motifs in 43 44 other plant genera. Here, we describe an accurate bottom-up approach using the online workbench RSAT:: Plants for a versatile ab-initio motif discovery taking Prunus persica 45 46 as a model. These predictions rely on the construction of a co-expression network to generate modules with similar expression trends and assess the effect of increasing 47 48 upstream region length on the sensitivity of motif discovery. Applying two discovery 49 algorithms, 18 out of 45 modules were found to be enriched in motifs typical of well-50 known transcription factor families (bHLH, bZip, BZR, CAMTA, DOF, E2FE, AP2-ERF, Myb-like, NAC, TCP, WRKY) and a novel motif. Our results indicate that small 51 number of input sequences and short promoter length are preferential to minimize the 52 amount of uninformative signals in peach. The spatial distribution of TF binding sites 53 revealed an unbalanced distribution where motifs tend to lie around the transcriptional 54 start site region. The reliability of this approach was also benchmarked in Arabidopsis 55 thaliana, where it recovered the expected motifs from promoters of genes containing 56 57 ChIPseq peaks. Overall, this paper presents a glimpse of the peach regulatory components at genome scale and provides a general protocol that can be applied to 58 many other species. Additionally, a RSAT Docker container was released to facilitate 59 similar analyses on other species or to reproduce our results. 60

Keywords: Motif prediction, cis-regulatory elements, *Prunus persica*, Transcription
Factor binding motifs

2

# 63 **1. Introduction**

Peach [Prunus persica (L.) Batsch], a member of Prunus genus, is one of the best 64 genetically characterized species within the Rosaceae family. With a small size diploid 65 66 genome (2n = 2x = 16; 230 Mbp), and relatively short generation time (2-3 years), peach has become a model species for fruit genetic studies (Abbott et al., 2002). Obtaining 67 elite genotypes with broad environmental adaptations and good fruit quality are the 68 fundamental targets of all Prunus breeding programs, since they directly affect the 69 70 economical relevance of this crop (Gogorcena et al., 2020). Indeed, previous works have reported strong affinity between environmental cues and the fruit quality and 71 72 aroma (Wong et al., 2016; Tanou et al., 2017). To stand the environmental stimuli and 73 ensure edible fruit development, a complex re-arrangement of the gene expression 74 network is required.

75 The modulation of gene expression is a complex process occurring at various levels from which the transcriptional regulation is the core control code (Petrillo et al., 2014). 76 The transcription machinery is regulated by an interplay between DNA-binding proteins 77 78 called transcription factors (TFs) and cis-regulatory elements (CREs). TFs bind short 79 sequences known as TF binding sites (TFBS) or motifs located at CREs (e.g., 80 promoters, enhancers, silencers). TFs may act as either activators or repressors of gene expression, leading to dynamic changes of the cellular pathways. For peach, annotation 81 82 of TFs is available in the plant transcription factor database (plantTFDB) (Tian et al., 83 2019).

As of February 2020, plantTFDB v5.0 stores 2780 peach TFs classified into 58 families 84 (http://planttfdb.cbi.pku.edu.cn/). While much is known about TF families, TF-binding 85 motifs remain elusive. Deciphering the cis-regulatory network has become a 86 prerequisite toward scoping out the foundations of transcriptional regulation in P. 87 persica. The computational exploration of these DNA motifs has been greatly 88 stimulated by the availability of genomic data and the release of whole genome 89 90 sequence assemblies (Verde et al., 2013; Verde et al., 2017). In this context, a variety of plant motif finders has emerged. Notwithstanding their value, they are hampered by 91 certain limitations such as, a restricted range of species, Promzea for maize (Liseron-92 93 Monfils et al., 2013), and AthaMap for Arabidopsis (Steffens et al., 2005), and limited 94 analysis capabilities around experimentally defined motifs as PlantCare, (Rombauts et

3

al., 1999) or PlantPAN, (Chang et al., 2008). Thereby, to circumvent these pitfalls, we
have adopted a plant-customized tool for *de novo* motifs discovery, RSAT::Plants
(http://rsat.eead.csic.es/plants/). RSAT has both a friendly user interface and commandline tools for versatile analyses in a wide collection of plants (Nguyen et al., 2018).
Since the analysis of proximal promoter regions is easier in small genomes with short
intergenic regions, most of cis-regulatory motif predictions so far have been conducted
in *Arabidopsis thaliana* (Ma et al., 2012; Korkuc et al., 2014; Cherenkov et al., 2018).

102 In *P. persica* there are only two examples of regulatory motif discovery, in particular on a set of 350 dehydrin promoter sequences (Zolotarov and Strömvik, 2015) and 30 103 104 heat responsive genes (Gismondi et al., 2020). In contrast to these case studies, we 105 propose a structured bottom-up framework to identify statistically over-represented 106 motifs on a genome scale. Our probabilistic approach relies on the hypothesis that genes 107 within co-expressed modules are likely co-regulated by the same TFs. This approach 108 has been successfully tested in other species, for example in Arabidopsis thaliana 109 (Koschmann et al., 2012; Ma et al., 2013) and maize (Yu et al., 2015). According to 110 Bianchi et al., 2015, an arbitrary defined segment of 1500 bp upstream of the transcription start site (TSS) can be considered as the proximal promoter in peach. 111 However, recent studies about the genomic delimitation of proximal promoters in 112 113 Prunus persica effectively reduced this region to a window of approximately 500 nt (Montardit-Tardà, 2018). 114

115 The proposed approach relies on three fundaments, i) an accurate definition of co-116 expressed gene modules, ii) an assessment of the effect of upstream region length 117 regarding the effectiveness of motif discovery and, finally iii) disclosing the effect of 118 splitting the analysis around the TSS site in discovering potential cis-elements. All 119 together, we demonstrate the utility of our strategy in analyzing genome wide data to provide insights on gene regulation dynamics across tissues and specific conditions. To 120 the best of our knowledge, no work has been reported on cis-elements present in P. 121 122 *persica* on genome wide level, hence the originality of our survey. Additionally, the 123 predicted motifs can be browsed (https://eead-csicfrom this study at compbio.github.io/coexpression motif discovery/peach/), where we provide readers 124 125 with direct links to the results, source code and a Docker container to reproduce the 126 analysis on any other plant species.

#### 127 **2. Results**

# 128 2.1 Identification of differentially expressed transcripts and construction of 129 weighted co-expression network

130 After quality assessment and pseudo-alignment, an expression matrix was generated 131 from eight peach published transcriptomes, including treated and control samples with their corresponding biological replicates. Differential analysis yielded 11,335 altered 132 133 transcripts using Q-value < 0.01 and  $|\beta| > 1$  thresholds. The number of differentially expressed transcripts (DETs) identified in each RNA-seq experiment is listed in Table 134 1. Detailed information about quality control, pseudo-alignment and differential 135 expression analyses is shown in Table  $S_1$ . An overview of our workflow is provided in 136 Figure 1. 137

The WGCNA R-package was adopted to construct an unsigned co-expression network 138 for 11,335 stress-related transcripts. All samples and DETs were considered in the 139 140 network construction, as neither outliers nor transcripts with missing values, were 141 detected (Figure S<sub>1</sub>, A). Using a dynamic tree cut algorithm, 45 co-expression modules 142 were retained with size ranging from 29 to 1795 transcripts per module (Figure  $S_1$ . C). 143 The 45 distinct modules (labeled with different colors) are shown in a dendrogram in which major tree branches constitute modules and leaves correspond to DETs (Figure 144 145 **S**<sub>1</sub>. **B**).

# 146

147

# 2.2.1 Effect of proximal promoter length on prediction accuracy

2.2 Transcription factor binding site (TFBS) prediction

As a first step towards extracting regulatory signatures, upstream region boundaries 148 149 were defined from -1500 bp to +200 bp relative to TSS (Up 1). Six out of 45 modules 150 were found to display positive signals and higher significance when compared to the 151 random clusters. Upstream regions of modules (M9, M10, M11, M18, M21 and M41) matched known core DNA-binding elements corresponding to Myb-like, BZR, 152 153 CAMTA, bZip, E2FE, and TF families. Modules with their corresponding regulatory elements are represented in Figure 2 and further information is provided in Table  $S_2$ . 154 Motifs resulting from both oligo and dyad analysis correspond to signatures with strong 155 156 confidence estimation. Besides, eight poly (AT)-rich signals were discarded from M1, M2, M3, M4 and M6 due to their low complexity. Curiously, these (AT) patterns were 157 158 also detected in the random clusters and their occurrence seemed to be associated with

the size of the module (**Table S**<sub>3</sub>). For instance, M1 is the largest module with 1795 sequences and (AT)-repetitive signals were detected in 40 out of the corresponding 50 random clusters.

Furthermore, when we restricted the motif discovery to the region with [-500 bp, +200
bp] boundaries (Up 2), fifteen modules were found to discern statistically significant
motifs. These were then grouped into 10 TF families as illustrated in Figure 2 (TCP,
bHLH, BZR, bZip, NAC, WRKY, AP2-ERF, Myb-like, CAMTA and E2FE).

An in-depth look at the major changes occurring when trimming the upstream segments 166 167 to 500 bp resulted in interesting observations, summarized as follows. Spurious (AT) rich events considered as low quality predictions were limited to M2 and were replaced 168 by relevant regulatory elements in M1, M3, M4 and M6 (Table S<sub>2</sub>). Significant signals 169 buried in the long upstream region (Up 1) were inferred in modules M24, M28 and M43 170 171 (Figure 2, Table  $S_2$ ). Besides, shortening the upstream promoter region size to 500 bp enhances the statistical relevance of the predicted motifs, compared to the negative 172 173 controls, regardless of the algorithm applied.

Overall, these findings suggest that shortening the upstream region increases the signalto-noise ratio to detect biologically relevant motifs and, at the same time, reduces the occurrence of low complexity AT-rich motifs. In **Figure 3**, we illustrate a clear showcase of this observation. Indeed, with both oligo and dyad analysis, the statistical significance of motif E2FE found in Module M41 (black bars) has noticeably increased compared to those identified in random clusters (gray bars). Hence, more significant motif discovery was accomplished in the window of [-500 bp, +200 bp].

181

182

# 2.2.2 Effect of splitting the promoter region around the TSS on motif prediction

183 Next, due to the difference in nucleotide composition in coding and non-coding regions, 184 we subdivided the proximal promoter region in two segments around the TSS, with 185 each interval examined separately: upstream, from -500 bp to 0 bp (Up 3), and 186 downstream, from 0 to +200 bp (Up 4). Doing so, motifs of two additional TF families 187 were discovered, BCP in module M1, DOF in modules M7, M9 and M21. In contrast to 188 BCP sites lying downstream the TSS (Up 4), DOF sites were found across both

6

intervals (see Figure 2, Table  $S_2$ ). Intriguingly, an uncharacterized motif was overrepresented in upstream 4 of module M25 requiring further research.

191 In conclusion, a total of 77 TF binding motifs were revealed from the different assessed 192 promoter regions (Table  $S_2$ ). Modules with candidate predicted motifs might be 193 classified in two types depending on their potentially matching TF. Indeed, across the 194 four examined upstream tracts, we recognize those with motifs bound by a single TF family, considered as single TF-driven modules (e.g., M6, M11, M18, M28 and M41). 195 Conversely, modules having multiple TFBS for several distinct TFs suggest a possible 196 combinatorial regulation under particular circumstances. However, more evidence is 197 needed to address this issue. On the other hand, we observed that the majority of cis-198 199 regulatory elements yielded in this study were mainly detected in the upstream region 200 Up 2, defined from -500 bp to + 200 bp (Figure 2, Table  $S_2$ ).

# 201 2.3 Gene Ontology enrichment

202 A Gene Ontology analysis was conducted to annotate the potential function of the 203 gene modules. Thirteen modules were significantly enriched with biological processes 204 (Figure 4). Six GO terminologies were particularly intriguing and will be briefly 205 described. In modules M1 and M18, transcripts were over-represented respectively in leaf and root tissues under drought experiment which is in line with the 206 207 "photosynthesis" and "response to water" enrichment. Similarly, module M2 was 208 enriched for "response to stimulus" with high TPM values in fruit tissue at different ripening stage. Transcripts within M5 were mostly abundant in fruit tissue under cold 209 stress, in line with the "cold acclimation" enrichment. Not surprisingly, "response to 210 stress" was over-represented in fruit in module M10 as we are dealing with stress 211 212 conditions. Finally, hormonal levels are known to imbalance under stress explaining the enrichment of "response to hormone stimulus" in M21. Overall, we consider that the 213 214 GO enrichment results (Figure 4.A) are in harmony with the expression profiles of 215 transcripts in Figure 4.B.

#### 216 2.4 TFs annotation and prediction of their TFBS using footprintDB

The predicted modules were examined for genes encoding TFs. In total 39 annotated TFs were shortlisted in **Figure 5.** Myb and Myb-like TFs were exclusively expressed in modules M1 and M2. They were particularly over-represented in fruit and leaf tissues in agreement with their transcript profiling illustrated in **Figure 4.B**. We hypothesize that

Myb factors may act as regulators of drought stress and ripening in peach. In the same 221 222 vein, bHLH genes identified in M3 were notably abundant in stigma tissue, which is in 223 accordance with Figure 4.B. NAC and E2FE transcription factors were respectively 224 annotated in M4 and M41, and their coding genes were repressed among experiments in all tissues. The WRKY TFs assigned to module M6 were abundant under hyper 225 226 hydricity fitting with **Figure 4.B** and suggesting a regulatory function of the WRKY in such a condition. Module M7 was associated with genes encoding three TFs with 227 228 different expression profiles (DOF, bHLH and ERF). Calmodulin binding proteins 229 identified in M11 and bZip annotated in M18 and M21 were highly abundant among all 230 experiments indicating that they may be involved in multiple biological processes.

231 Subsequently, we verified whether the disclosed motifs in each module are the actual binding sites of the aforementioned TFs (Figure 5). TFs were individually examined for 232 233 their potential DNA-site using footprintDB and results were compared to those derived 234 from RSAT. Consensus sequences predicted from genes coding TFs showed high 235 similarity to consensus sequences predicted from modules (Table 2). As for instance, 236 the binding motif "tTTGGCGGGAAA" identified in module M41 is almost identical to E2FE-predicted site "TTTTGGCGGGAAAA" from the same module. This suggests 237 that E2FE may modulate gene expression in M41 and "tTTGGCGGGAAA" motif 238 239 could be the *bona fide* binding site of this transcription factor.

#### 240 **2.5 Motif scanning**

To identify the position of transcription factor binding sites (TFBS) in the promoter 241 region of *P. persica* genes, position-specific scoring matrices (PSSMs) of all candidate 242 motifs (77) were in silico scanned to the long (Up 1) upstream stretch [-1500, +200 bp]. 243 We observed a clear positional bias of the TFBS close to the TSS, more precisely within 244 the interval [-500 bp, +200 bp], then it progressively declines towards the 5' limit 245 (Figure 6). For motifs detected respectively in Up 1 (yellow color), Up 2 (green) and 246 Up 3 (blue), sites were notably concentrated upstream the TSS showing a bell-shaped 247 248 distribution from -500 bp to +0 bp with a maximum of density around -250 bp. Conversely, the positional distribution of motifs predicted along the upstream 4 was 249 250 biased toward downstream the TSS with the flatter peak reaching its limit at the TSS (Up 4, purple). Detailed scanning results can be accessed at https://eead-csic-251 252 compbio.github.io/coexpression motif discovery/peach. On the other side, (AT)

repetitive elements were also scanned to check their relevance, e.g., whether they correspond to the TATA box. The underlying data included in **Figure S**<sub>2</sub>, showed that TFBSs of these motifs were remarkably distant to the TSS and were distributed across the whole proximal region.

#### 257 **2.6** Validation of the protocol for *de novo* cis-element discovery

258 To demonstrate the performance of the motif finding approach, we evaluated the effect 259 of variable proximal promoter lengths on uncovering true DNA-binding sites in Arabidopsis thaliana. Experimentally proven motifs from a selection of A. thaliana 260 261 transcription factors belonging to different families were successfully recovered by at 262 least one algorithm. As summarized in Figure 7, JASPAR and *de novo* identified motifs displayed high consensus similarity. Moreover, in order to refine the comparison, we 263 264 annotated the newly reported motifs JASPAR to ensure that they correspond to the TF 265 family in question. As expected, *de novo* motifs shared the same annotation as the 266 reference JASPAR motifs, which underlines the predictive performance of the proposed 267 methodology.

#### 268 **3. Discussion**

In the present study, transcriptional profiling of eight independent data sets was 269 270 conducted to decipher the intricate process of gene regulation in peach and to reveal 271 meaningful biological signatures. DETs were grouped into 45 co-expression modules 272 undergoing similar changes in their expression patterns. Unlike conventional clustering 273 methods (such as k-means and hierarchical clustering), which are based on geometric 274 distances, WGCNA is a graph-based approach relying on network topology as inferred from the correlation among expression values (Li et al., 2018). In our hands, the 275 276 WGCNA algorithm robustly and accurately defined modules within a complex multi-277 condition dataset.

Discerning regulatory signals from blocks of co-expressed genes is a common presumption used to identify functional genomic elements. It has been successfully applied and approved in various plants species like *Arabidopsis thaliana* (Koschmann et al., 2012; Ma et al., 2013), *Zea mays* (Yu et al., 2015) and *Hordeum vulgare L.* (Cantalapiedra et al., 2017). However, little is known about its applicability to woody species. To our knowledge, this article is the first in which this hypothesis has been tested in *Prunus persica* genome wide.

For each predicted module, two-motif discovery algorithms (oligo and dyad analysis) 285 286 were ran to discover significant motifs in the upstream promoter region. As suggested by Bianchi and colleagues, we initially defined the upstream promoter size as an 287 interval of [-1500 bp to +200 bp] relative to the TSS (Bianchi et al., 2015). Discovered 288 motifs with significant poly-(AT) sites were discarded due to their low complexity and 289 scarcity of information concerning their specific-regulatory function. We reasoned that 290 291 low complexity sequences might be linked to repetitive stretches of DNA, extensively 292 present in plant genomes (Yu et al., 2015). Interestingly, when tuning the promoter 293 upstream length to a tract of [-500 bp, +200 bp] relative to the TSS, these low 294 complexity motifs were limited to module M2. It would seem that long upstream 295 promoter regions unbalance the signal-to-noise ratio exacerbating the identification of such AT motifs. Along the same lines, we observe a dependence of (AT)-rich sites on 296 297 the dataset size. Indeed, AT-low-complexity motifs were only detected in the first six 298 modules, which contain from 560 to 1795 upstream sequences. In light of these 299 considerations, we believe that in our study case, they may result in part due to the 300 properties of DNA sequences (both upstream region length and dataset size) rather than 301 the performance of the chosen algorithms. In Table S<sub>3</sub>, the results revealed that AT-rich occurrence in random cluster increases in parallel with the module size. 302

To check whether the AT-rich patterns overlap the TATA boxes, a position scanning experiment was conducted. It is well documented in plants that a TATA box region lays between -30 and +35 bp with respect to the TSS (Zhu Qun et al., 1995; Smale, 2001) However, the scanning results portrayed that peaks were located far from this interval, confirming that they are distinct signals (**Figure S**<sub>2</sub>).

308 By limiting the promoter length to a window of -500 bp, new regulatory motifs were recovered. Additionally, splitting the proximal promoter region into two intervals 309 310 around the TSS enabled the discovery of further hidden candidate TF motifs. Such 311 observations may strengthen our hypothesis that shorter upstream regions improve the 312 sensitivity motif discovery (from 11 motif sequences identified within Up 1 to 58 313 sequences identified in Up 3 and Up 4 assessed separately). Defining the upstream 314 promoter length has been a controversial issue (Kristiansson et al., 2009). If the interval is too short or too long, the motif of interest may not be captured. Therefore, we reason 315 that an analysis on regions of variable length would yield a more comprehensive picture 316 of the complex regulatory code. 317

The spatial distribution of the occurrences of the 77 inferred motifs along the promoter 318 319 region is crucial to understand gene regulation in Prunus persica. Our findings revealed 320 that TFBSs are not uniformly dispersed across the promoter but they exhibit a strikingly 321 mixture of 2 density profiles: while the majority showed bell-shaped distribution at the interval of [-500 bp, 0 bp], others were diverged downstream the TSS [0 bp, +200 bp] 322 323 (Figure 6). These findings are similar to those described in A. thaliana, with nearly two thirds of the examined TFBSs within the region from -400 bp to +200 bp (Yu et al., 324 325 2016). TFBSs of bHLH, BZR, TCP and WRKY are particularly concentrated from -500 326 bp to 0 bp. This denotes a positional binding preference within this proximal region, 327 which is in agreement with (Yu et al., 2016) reporting that their positional preference is 328 between -100 bp to -50 bp. On the other hand, bZip, CAMTA, E2FE and Myb-like exhibited a dual binding distribution with central peaks upstream and downstream the 329 330 TSS. A possible explanation of this is that some TFs may display different binding preferences depending on their TF-specific structure, biological functions or 331 332 combinatory with other TFs. The degree to which the arrangement of motif sites is 333 associated to their function needs to be further investigated especially that data about 334 TFBS distribution in plants is only limited to Arabidopsis thaliana (Zou et al., 2011; Yu et al., 2016). According to our findings, we may consider that the boundary from -335 500 bp to 0 bp is an adequate region to look for the majority of TFBSs lying in the 336 proximal promoter region in peach. However, we should keep in mind that proximal 337 TFBSs could also occur downstream the TSS. Thus, we suggest defining the peach 338 proximal promoter length as a tract of [-500 bp to +200 bp], analyzing separately the 339 two regions around the TSS for a better motif coverage. In fact, according to Montardit-340 341 Tardà (2018), differences in the nucleotide composition were found upstream and 342 downstream the TSS. At this point, we should mention that gene regulation involves a 343 complex interplay between the proximal (promoter) and distal regulatory regions located thousands of base pairs away from the TSS (e.g., enhancers) (Li et al., 2019). 344 345 Our workflow sheds light mainly on sequence signatures extracted from the proximal promoter. Thus, it might not be adequate to study distal genomic elements. 346

Furthermore, rather than barely returning a list of significant motifs, our methodology
assigned them to different modules to help shape a clear overview of the peach
regulation code. Overall, we were able to distinguish 18 modules harboring 77 motifs
from 11 TF families: bHLH, bZip, BZR, CAMTA, DOF, E2FE, AP2-ERF, Myb-like,

NAC, TCP and WRKY. While some modules, such as M6, M11, M28 and M41, seem to 351 352 be driven by a single TF (WRKY, CAMTA and E2FE, respectively), motifs from 353 different families were annotated in the rest. This can be explained by the fact that some 354 promoter sequences may encompass multiple TFBSs of perhaps interacting TFs. Indeed, TFs have been reported to frequently operate in combination (Guo et al., 2018; Kumar 355 356 et al., 2018). Combinatorial regulation is required to confer specific responses in a particular tissue and under a particular stress. Thus, the hypothesis of cooperative 357 358 interactions between diverse motifs in peach is worthy to be further investigated.

From the inferred list of motifs (**Figure 2**), we found similar binding sequence potentially perceived by different class of transcription factors. For example, motifs "tGaCACGTGtc" and "GaCACGTGkCGg" in module M5 are distinct but can be aligned despite different nucleotide frequencies in some positions. We presume that TFs from related families may have similar DNA recognition sequences, as reported for instance by Franco-Zorrilla et al., 2014 for Myb and AP2 TFs.

The biological significance of modules with significant identified signals was 365 366 determined by Gene Ontology analysis and TF annotation. The enriched modules reflected many biological functions involved in abiotic stress responses such as cold 367 368 acclimation, response to stress, response to water and response to hormone (Figure 4). 369 In this context, modules M1 enriched for "photosynthesis" contained candidate Myb 370 and Myb-related factors. These findings are in line with (Baldoni et al., 2015) reporting 371 that Myb TF family is known to regulate drought tolerance and the stomatal movements in plants. bHLH binding sites were mainly disclosed in modules M3, M5 and M7 372 (Figure 5). Associated TFs among those were abundant under various stress conditions 373 proposing a multi-functional role of bHLH. According to Bianchi et al., (2015), bHLH 374 factors play a central role in flavonoid biosynthesis and cold acclimation in peach. 375 Similarly, bZip TFs were found in both M18 and M21 and their transcripts were mainly 376 377 over-represented in all experiments. Our results are supported by previous studies 378 reporting that bZip were induced by various environmental cues. Indeed it was revealed 379 that they play a pivotal role in responses to cold stress in peach and enhance water use 380 efficiency in almond-peach rootstocks (Hu et al., 2018). WRKYs putative motifs were 381 restricted to M6 and were exclusively activated in leaf tissue under hyper-hydricity 382 (HH) stress. It is well known that HH leads to morphological abnormalities, such as 383 brittle leaves (Carrillo Bermejo et al., 2017). We speculate that WRKY factors may be

implicated in morphological damages produced by HH. Module M11 was found to be a 384 385 potential CAMTA-driven module (Figure 2), where two genes coding CAMTA were annotated (Figure 5). A previous study in A. thaliana demonstrated that cold stress 386 increases the level of calcium sensed by CAMTA (Doherty et al., 2009). This 387 perturbation of calcium levels leads to modification of the CAMTA activity that in turn 388 triggers the induction of cold response genes of the CBF family. For this reason, 389 CAMTA motifs are of great interest. From the perspective of peach breeding, these 390 391 findings may be of great interest, as genes within modules are potential targets for 392 further experimental validation.

393 Finally, a major drawback of motif discovery approaches is their limited performance. To tackle this issue we designed a control experiment in which genomic sites detected 394 395 by ChIP-seq for 10 A. thaliana TFs were analyzed. Comparing the de novo predicted motifs to the corresponding curated motifs in JASPAR we observed a high similarity in 396 397 terms of Ncor scores (Figure 7 and Table S<sub>4</sub>). When searching for *in-vivo* validated motifs, we would ideally expect to get identical predicted motifs. Nonetheless, while 398 399 most consensus sequences had high Ncor values > 0.8, others had lower values. As well, we observed that the choice of upstream region length affects the performance. In some 400 401 cases, particularly Up 1 and Up 3, the expected motif was not even found. Unlike the results found in peach, examining 4 upstream tracts only returned motifs from the same 402 403 query families probably as a consequence of the JASPAR TFBSs profiles being curated. Taken together, we believe that the proposed workflow is robust enough to be extended 404 405 to other species in order to identify reliable regulatory motifs.

406 **4.** Conclusion

407 DNA motif discovery is a primary step for studying gene regulation, however the in silico prediction of regulatory motifs in not straightforward. In contrast to previous 408 surveys that usually assume a fixed promoter length right at the start; this work reports 409 410 regulatory elements while testing different upstream sequence intervals. It is among the first efforts providing a comprehensive collection of Prunus persica motifs without a 411 412 prior knowledge. By coupling gene expression networks and module analysis, we were able to extract interpretable information from a large set of noisy data and to reveal 413 primary candidate TF-target binding sites responding to specific conditions. These 414 results offer a more complete view of the proximal regulatory signatures in P. persica 415

and we believe that it may contribute to address the knowledge gap about thetranscriptional regulatory code in non-model species.

# 418 **5.** Materials and methods

# 419 **5.1 Input data and processing**

Eight peach RNA-sequencing datasets were downloaded from the European Nucleotide 420 421 Archive (https://www.ebi.ac.uk/ena) and were used as raw reads for this project. This 422 comprehensive dataset includes data of various peach cultivars, from various tissues 423 (root, leaf, stigma and fruit), different stress conditions and developmental stages. A 424 detailed list of the project IDs and metadata is provided respectively in Table 1 and 425 **Table S\_{1}.** The obtained reads were quality-processed and trimmed using FASTQC v.0.11.5 and Trimmomatic v.0.36 (Bolger et al., 2014), to discard adaptors and low-426 quality sequences with mean Phred score (Q < 30) and window size of 4:15. The first 427 nucleotides were then head-cropped to ensure a per-position A, C, G, T frequency near 428 429 to 0.25. Following the trimming, only sequences longer than 36 bp were retained for 430 further analysis. The complete workflow is shown in Figure 1 (see step 1).

431 The high quality reads from each RNA-seq project were quantified separately using the 432 pseudo-aligner kallisto v.0.43.1 for fast and accurate transcripts count and abundance 433 (Bray et al., 2016). Kallisto was run in two steps: i) a transcriptome index was built from all cDNA transcripts of Prunus persica v2, from Ensembl Plants release 39 (Verde 434 et al., 2017; Howe et al., 2020). ii) Each sample was pseudo-aligned against the index. 435 436 Transcript level abundance was estimated and normalized to transcripts per million 437 (TPM) using 100 bootstraps (-b 100) to ascertain the technical variation. For single-end 438 read mode, average fragment length and standard deviation were additionally required and were set to (-1 200) and (-s 50), respectively. 439

# 440 5.2 Transcript-level profiling

441 Differential expression analysis was conducted with Sleuth R package v.0.29.0 442 (Pimentel et al., 2017) for each RNA data set separately. The Wald test (WT) was 443 applied to output abundance files in order to retain the significant expressed transcripts 444 from each experiment. Samples and their biological replicates from each experiment 445 were compared with their corresponding control. To reduce the false positives, only 446 transcripts passing an FDR cutoff *Q*-value < 0.01 and beta statistic (approximation of 447 the Log<sub>2</sub> Fold Change between two tested conditions)  $|\beta| > 1$  were retained. Significant

transcripts obtained from each RNA-seq project were merged into a single list with anassigned mean TPM value for each replicate.

#### 450 **5.3 Construction of co-expressed network**

451 Based on the assumption that co-expressed genes may share the same biological signature, weighted gene co-expression network analysis (WGCNA v.1.61) was 452 performed to extract clusters of densely interconnected genes named modules 453 454 (Langfelder and Horvath, 2008). Samples were firstly clustered to remove outliers and 455 transcripts with missing entries. A similarity matrix was constructed by performing 456 pairwise Pearson correlation across all targets. Then an adjacency matrix was built raising the similarity matrix to a soft power ( $\beta$ ). Here  $\beta$  was set to 7 reaching thus 83% 457 of the scale free topology fitting index  $(R^2)$ . To minimize the effect of noise, matrix 458 adjacency was transformed to Topological Overlap Measure (TOM) and its 459 corresponding dissimilarity matrix (1-TOM) was generated. Finally, modules were 460 defined using the cutreeDynamic function with a minimum module size of 20 targets. 461 462 Compared to standard hierarchical clustering, this approach solves the issue of setting 463 the final number of clusters and arranges the genes based on their topological overlap to eliminate spurious associations resulting from the correlation matrix. 464

#### 465 5.4 *De novo* cis regulatory sequences discovery using RSAT::Plants

466 Gene modules resulting from network analysis were subjected to an *ab-initio* motif 467 discovery pipeline using the RSAT::Plants standalone (Figure 1, step 2). For each module, the analysis initiates by generating as negative control 50 random clusters of 468 469 the same size for each module as described previously (Contreras-Moreira et al., 2016). 470 Sequences with four different boundaries around the TSS were retrieved from the genes 471 in the co-expressed modules, random clusters and *Prunus persica* genome v2. The upstream sequences were defined as intervals of i) -1.5 kb to +200 bp ii) -500 bp to 472 +200 bp and iii) two segments around the TSS: -500 bp to 0 and 0 bp to +200 bp. Note 473 474 that the 0 to +200 interval corresponds to the 3' UTR region, which is already downstream. RSAT *peak-motifs* was run under the differential analysis mode, where 475 476 module's upstream sequences served as the test set and all upstream sequences from 477 peach genome were considered as the control set to estimate the background model (a 478 background model was created for each upstream stretch) (Thomas-Chollier et al., 479 2012). Two discovery algorithms were used: i) oligo-analysis, which is based on the 480 over-representation of k-mers in upstream regions, and ii) dyad-analysis, which looks 481 for over-represented spaced pairs of oligonucleotides (Defrance et al., 2008). For each 482 run, up to five motifs were returned per algorithm and were retained to compare their 483 statistical significance with the 50 random clusters considered as negative control.

- 484 Candidate motifs were chosen based on their significance (log E-value) compared to
- 485 negative control and were subsequently annotated by comparison to the footprintDB
- 486 collection of plant curated motifs (http://floresta.eead.csic.es/footprintdb) (Sebastian and
- 487 Contreras-Moreira, 2014) using the *compare-matrix* tool in RSAT (Nguyen et al., 2018)
- 488 requiring a normalized correlation score Ncor  $\geq 0.4$ .
- Finally, selected motifs were scanned along the stretch [-1500 bp, +200 bp] to predict their corresponding binding site positions, using as background model a Markov chain of order 1 (m=1) and a cutoff *P*-value  $\leq 1e^{-4}$ . To ensure the clarity and reproducibility of this strategy, a repository including the source code, links to the results and a tutorial explaining how to reproduce a similar analysis on any species is available at https://eead-csic-compbio.github.io/coexpression\_motif\_discovery/peach.

# 495 **5.5 Transcription factor prediction and Gene Ontology analysis**

Hereafter, the analysis was restricted to modules with significant detected signals. 496 Firstly, genes coding peach TFs were predicted and classified using the iTAK database 497 498 (http://itak.feilab.net/cgi-bin/itak/index.cgi, last accessed January 2020). Protein 499 sequences of TFs were subsequently submitted to footprintDB to predict their interacting DNA-binding site. To functionally interpret the co-expressed modules, Gene 500 Ontology (GO) enrichment was conducted on PlantRegMap / PlantTFDB portal v5.0 501 502 (http://planttfdb.gao-lab.org/, last accessed January, 2020) (Tian et al., 2019). P-value of 0.01 was set to retain the significant GO terms. 503

# 5.6 Validation of the pipeline by detecting *a priori* known motifs in *Arabidopsis thaliana*

To assess the impact of upstream region lengths on the identification of relevant motifs, we used sets of experimentally validated binding sites of 10 *Arabidopsis thaliana* TF families. Sequences of the proven sites were downloaded from JASPAR database (Fornes et al., 2020) and were locally aligned with BLASTN against the *A. thaliana* TAIR10.42 genome from Ensembl Plants to obtain the closest neighbor genes. The following parameters were used: E-value  $\leq 1e^{-5}$ , max\_target\_seqs =1, max\_hsps=1

512 query-coverage of 80% and percentage of identity 98%. Upstream sequences of 513 neighbor genes were obtained with *retrieve-seq* from RSAT::Plants. Similarity between 514 references (JASPAR) and newly discovered motifs was computed with Ncor score (see 515 above).

# 516 Supplemental Data

517 **Supplemental Table S1. A.** Detailed information about the RNA-seq data used for 518 differential analysis

**Supplemental Table S1. B.** Number of survived and dropped reads after quality
processing and pseudo-aligned reads using kallisto program

521 **Supplemental Table S2.** List of candidate regulatory sites discovered within four 522 upstream tracts of different lengths. Motifs are represented as IUPAC consensus 523 sequences. TF match: Transcription factor family of the best match in footprintDB. 524 Ncor: normalized Pearson correlation varying between 0 and 1. Ncor  $\ge$  0.4 indicates 525 high confidence annotations. Gray color indicates that no significant motifs were found.

**Supplemental Table S3.** List of low complexity motifs considered as false positive predictions within a boundary from -1500 bp to +200 bp upstream region length. For each algorithm, sequences are presented both as IUPAC consensus sequences using the degeneracy code and as sequence logos. Last column indicates the occurrence number of AT-rich motifs within the 50 random clusters used as negative control. Ps: W letter refers to (A or T) nucleotide and S refers to (C or G). Number of sites corresponds to the occurrence number of a single motif.

Supplemental Table S4. Similarity of JASPAR motifs (considered as queries) and de *novo* predicted dyad motifs in *Arabidopsis thaliana*. Numbers tagged with asterisks
indicate number of peaks recovered by BLASTN (see Methods). The Ncor scores
correspond to JASPAR databases.

**Supplemental Figure S1.** Co-expression network analysis. (A): Sample clustering to detect outliers. Sample with the same node color are derived from the same RNAexperiment. (B): Topological overlap measure plot. The different shades of color signify the strength of the connections between the genes (from white not significantly correlated to red signifying highly significantly correlated). Modules identified are colored along both column and row and are boxed. (C): Distribution of the module size.

17

544 Supplemental Figure S2. Positional distribution of AT-rich repetitive motifs along
545 upstream 1: [-1500 bp, +200 bp].

#### 546 Acknowledgements

- 547 We thank Eric OLO NDELA for his support and help in creating the HTML report and
- 548 providing useful feedback. We also thank Claudio Antonio Meneses Araya, Dayan
- 549 Sanhueza and Tomás Carrasco for giving us access to the RNA-seq data.

#### 550 Tables

- 551 **Table 1.** Summary of RNA-seq data used as input and the number of differentially
- 552 expressed transcripts (DETs) identified in each RNA-seq experiment.

Project ID	References	Experiments	Tissues	Conditions	DETs
PRJNA271307	(Li et al., 2015)	Ripening stage	Fruit	6	2601
PRJNA288567	(Sanhueza et al., 2015)	Cold storage	Fruit	6	6447
PRJNA248711	(Bakir et al., 2016)	Hyper hydricity	Leaf	2	15
PRJEB12334	(Ksouri et al., 2016)	Drought	Root/Leaf	4	350
PRJNA252780	(Jiao et al., 2017)	Low $T^{\circ}$	Stigma	2	406
PRJNA323761	Unpublished	Drought	Root	2	1118
PRJNA328435	Unpublished	Cold storage	Fruit	2	2963
PRJNA397885	Unpublished	Chilling injury	Fruit	4	2429

553

554	Table 2.	Similarity	comparison	between	RSAT	and	footprintDB	DNA-binding	motif

predictions. The best predictions in footprintDB were selected in *Arabidopsis thaliana*.

556	The TFs grouped in	this table are the same	labeled with a star i	n <b>Figure 5</b>

Modules	lodules RSAT		Gene IDs	FootprintDB	STAMP
	Consensus			Consensus	E-value
M41	tTTGGCGGGAAA	E2FE	Prupe.5G180000	TTTTGGCGGGAAAA	3e <sup>-138</sup>
M21	GaCACGTGkC	bZip	Prupe.1G455300	ACGTGgc	3e <sup>-20</sup>
M18	tGCCACGTGGC	bZip	Prupe.1G419700	TGACGTGGC	1e <sup>-16</sup>
M18	tGCCACGTGGC	bZip	Prupe.1G434500	CACGTGGC	1e <sup>-127</sup>
M18	tGCCACGTGGC	bZip	Prupe.2G182800	TGCCACGT	8e <sup>-125</sup>
<b>M7</b>	tGCCGACa	AP2-ERF	Prupe.3G157100	TGCCGCC	1e <sup>-49</sup>
<b>M7</b>	tGCCGACa	AP2-ERF	Prupe.7G222700	CCGACA	4e <sup>-47</sup>
<b>M7</b>	CACGTGkCGG	bHLH	Prupe.6G303500	CCACGTGr	2e <sup>-84</sup>
<b>M7</b>	aAAAGTc	DOF	Prupe.6G092600	AAAG	2e <sup>-34</sup>
<b>M6</b>	GaAAAGTCaaa	WRKY	Prupe.4G075400	AAAGTCAA	4e <sup>-63</sup>
<b>M6</b>	GaAAAGTCaaa	WRKY	Prupe.5G106700	aAAAGTCAA	2e <sup>-59</sup>
<b>M4</b>	ttAAGCAAata	NAC	Prupe.1G106100	AAGcAAc	7e <sup>-10</sup>
M4	ttAAGCAAata	NAC	Prupe.7G102000	AAGCAA	9e <sup>-35</sup>
M3	CGaCACGTGtCGGtt	bHLH	Prupe.1G252600	CACGTGA	8e <sup>-15</sup>
M3	CGaCACGTGtCGGtt	bHLH	Prupe.2G190100	CACGTGC	3e <sup>-77</sup>
M3	CGaCACGTGtCGGtt	bHLH	Prupe.6G159200	gCACGTG	5e <sup>-20</sup>
M3	CGaCACGTGtCGGtt	bHLH	Prupe.3G064500	CACGTG	9e <sup>-10</sup>

#### 557

# 558 Figure Legends

**Figure 1.** Bottom-up framework for *de novo* motif discovery. <u>Step1</u>: differential expression analysis for transcript detection and extraction of co-expressed modules. <u>Step2</u>: *de novo* motif detection using the peak-motifs tool from RSAT::Plants. Numbers correspond to the different tested upstream tracts, with TSSs anchored on position 0 bp, while letters represent tools within peak-motifs. Green and orange boxes label software and RSAT tools, respectively.

**Figure 2.** Position Specific Scoring Matrix (PSSM) representation of top scored discovered motifs per modules, along different upstream lengths. The x-axis

corresponds to the four intervals: Up 1: [-1500 bp, +200 bp], Up 2: [-500 bp, -200 bp], 567 568 Up 3: [-500 bp, 0 bp] and Up 4 [0 bp, +200 bp]. The y-axis informs about the motif 569 family revealed per module. Cell colors indicate the statistical significance of the 570 identified motifs while cell sizes represent the normalized correlation (Ncor). Number of sites corresponds to the number of sites used to build the PSSM. When motifs from 571 572 the same family are identified with both algorithms (oligo and dyad-analysis), or in different upstream tracts (Up 1, Up 2, Up 3 and Up 4), only the most significant one is 573 574 represented in the heatmap. Further details are provided in Table S3. An interactive 575 report with source code is accessible at https://eead-csic-576 compbio.github.io/coexpression\_motif\_discovery/peach/

577 Figure 3. Illustrative comparison between predicted motif DEL2 (corresponding to 578 E2FE transcription factor) within two different upstream promoter lengths: -1500 bp to +200 bp (A) and -500 bp to +200 bp (B). The name of the best match among plant 579 580 motifs in footprintDB is labeled in red, next to its Ncor (Normalized correlation) value 581 labeled in blue. The x-axis corresponds to the module of interest (M41) and random 582 clusters ranked ranked by the most significant motifs. The y-axis corresponds to the statistical significance -log10 (P-value). Number of sites corresponds to the occurrence 583 584 number of a single motif. The evidence supporting the putative motifs is Ncor (in blue) 585 and the significance (black bars) when compared to negative controls (gray bars).

586 Figure 4. Functional annotation of relevant gene modules. (A): Gene ontology 587 enrichment. (B): Mean transcript abundance profiling in term of transcripts per million 588 (TPM). The x-axis corresponds the different experimental conditions while the y-axis 589 indicates the number of differentially transcripts per module. Experiment and tissue 590 types are highlighted by different colors (see the color key at the bottom of the figure). 591 Gene profiles along the different conditions are provided at (https://eead-csic compbio.github.io/coexpression\_motif\_discovery/peach). See supplementary Table S<sub>1</sub> 592 for the abbreviations. 593

Figure 5. List of transcription factors within relevant modules. Blue and red squares
indicate transcripts per million while bottom color bars correspond to the tissues types
and different experiments, respectively (See the legend at the right side of the figure).
TFs showing sequence similarity between their footprintDB and RSAT predicted motifs
are labeled with a star.

**Figure 6.** Positional distribution of the detected oligo motifs in promoter genes of *Prunus persica*. Four density distributions were derived from four assessed upstream regions. Up 1: from -1500 bp to 200 bp, Up 2: from -500 bp to +200 bp, Up 3: from -500 bp to 0 bp and Up 4 from 0 bp to + 200 bp. The x-axis corresponds to upstream length in base pairs (bp). The y-axis corresponds to density of captured sites with *P*value <10 e<sup>-4</sup>. Only oligo motifs are presented here, dyads are provided in the report at https://eead-csic-compbio.github.io/coexpression\_motif\_discovery/peach.

- Figure 7. Similarity between JASPAR motifs (considered as queries) and *de novo* predicted oligo motifs found in *Arabidopsis thaliana* along four different upstream
  regions. Numbers tagged with a star indicate number of peaks recovered by BLASTN
  (see Methods). The Ncor scores correspond to JASPAR databases. Only oligo-analysis
  motifs are shown (dyads are available at supplementary Table S<sub>4</sub>). Upstream 1: [-1500
  bp to +200 bp], Upstream 2: [-500 bp to +200 bp], Upstream3: [-500 bp to 0 bp] and
  Upstream4: [0 bp to +200 bp]
- 613 Literature cited
- Abbott AG, Georgi L, Yvergniaux D, Wang Y, Blenda A, Reighard G, Inigo M,
  Sosinski B (2002) Peach: The model genome for Rosaceae. Acta Hortic 575: 145–
  155
- Bakir Y, Eldem V, Zararsiz G, Unver T (2016) Global transcriptome analysis reveals
   differences in gene expression patterns between nonhyperhydric and hyperhydric
   peach leaves. Plant Genome 9: 1–9
- Baldoni E, Genga A, Cominelli E (2015) Plant MYB transcription factors: Their role
  in drought response mechanisms. Int J Mol Sci 16: 15811–15851
- Bianchi VJ, Rubio M, Trainotti L, Verde I, Bonghi C, Martínez-Gómez P (2015)
   Prunus transcription factors: breeding perspectives. Front Plant Sci 6: 1–20
- Bolger AM, Lohse M, Usadel B (2014) Trimmomatic: A flexible trimmer for Illumina
  sequence data. Bioinformatics 30: 2114–2120
- Bray NL, Pimentel H, Melsted P, Pachter L (2016) Near-optimal probabilistic RNAseq quantification. Nat Biotechnol 34: 525–528

Cantalapiedra CP, García-pereira MJ, Gracia MP, Igartua E (2017) Large
differences in gene expression responses to drought and heat stress between elite
Barley cultivar scarlett and a spanish landrace. Front Plant Sci 8: 1–23

- Carrillo Bermejo EA, Alamillo MAH, Samuel David GT, Llanes MAK, Enrique C
   de la S, Manuel RZ, Rodriguez Zapata LC (2017) Transcriptome, genetic
   transformation and micropropagation: Some biotechnology strategies to diminish
   water stress caused by climate change in sugarcane. Plant, Abiotic Stress
   Responses to Clim. Chang. IntechOpen, pp 90–108
- Chang WC, Lee TY, Huang H Da, Huang HY, Pan RL (2008) PlantPAN: Plant
  promoter analysis navigator, for identifying combinatorial cis-regulatory elements
  with distance constraint in plant gene groups. BMC Genomics 9: 1–14
- Cherenkov P, Novikova D, Omelyanchuk N, Levitsky V, Grosse I, Weijers D,
   Mironova V (2018) Diversity of cis-regulatory elements associated with auxin
   response in Arabidopsis thaliana. J Exp Bot 69: 329–339
- 642 Contreras-Moreira B, Castro-Mondragon JA, Rioualen C, Cantalapiedra CP, Van
  643 Helden J (2016) RSAT::Plants: Motif discovery within clusters of upstream
  644 sequences in plant genomes. *In* R Hehl, ed, Plant Synth. Promot. Methods Mol.
  645 Biol. Humana Press, New York, pp 279–295
- 646 Defrance M, Janky R, Sand O, van Helden J (2008) Using RSAT oligo-analysis and
  647 dyad-analysis tools to discover regulatory signals in nucleic sequences. Nat Protoc
  648 3: 1589–1603
- Doherty CJ, Van Buskirk HA, Myers SJ, Thomashow MF (2009) Roles for
   Arabidopsis CAMTA transcription factors in cold-regulated gene expression and
   freezing tolerance. Plant Cell 21: 972–984
- Fornes O, Castro-Mondragon JA, Khan A, van der Lee R, Zhang X, Richmond
  PA, Modi BP, Correard S, Gheorghe M, Baranašić D, et al (2020) JASPAR
  2020: update of the open-access database of transcription factor binding profiles.
  Nucleic Acids Res 48: 87–92
- Franco-Zorrilla JM, López-Vidriero I, Carrasco JL, Godoy M, Vera P, Solano R
   (2014) DNA-binding specificities of plant transcription factors and their potential

to define target genes. PNAS **111**: 2367–2372

- 659 Gismondi M, Daurelio LD, Maiorano C, Monti LL, Lara M V., Drincovich MF,
- Bustamante CA (2020) Generation of fruit postharvest gene datasets and a novel
  motif analysis tool for functional studies: uncovering links between peach fruit
  heat treatment and cold storage responses. Planta 251: 1–18
- Gogorcena Y, Sánchez G, Moreno-vázquez S, Pérez S, Ksouri N (2020) Genomicbased breeding for climate-smart peach varieties. *In* C Kole, ed, Genome Des.
  Clim. fruit Crop. Springer-Nature, pp 291–351
- Guo J, Chen J, Yang J, Yu Y, Yang Y, Wang W (2018) Identification,
  characterization and expression analysis of the VQ motif-containing gene family in
  tea plant (Camellia sinensis). BMC Genomics 19: 1–12
- Howe KL, Contreras-moreira B, Silva N De, Maslen G, Akanni W, Allen J,
  Alvarez-jarreta J, Barba M, Bolser DM, Cambell L, et al (2020) Ensembl
  Genomes 2020 enabling non-vertebrate genomic research. Nucleic Acids Res 1–7
- Hu P, Li G, Zhao X, Zhao F, Li L, Zhou H (2018) Transcriptome profiling by RNASeq reveals differentially expressed genes related to fruit development and ripening
  characteristics in strawberries (Fragaria × ananassa ). Peer J 6: 1–25
- Jiao Y, Shen Z, Yan J (2017) Transcriptome analysis of peach [Prunus persica (L.)
  Batsch] stigma in response to low-temperature stress with digital gene expression
  profiling. J Plant Biochem Biotechnol 26: 141–148
- Korkuc P, Schippers JHM, Walther D (2014) Characterization and identification of
   cis-regulatory elements in Arabidopsis based on single-nucleotide polymorphism
   information. Plant Physiolgy 164: 181–200
- Koschmann J, Machens F, Becker M, Niemeyer J, Schulze J, Bulow L, Stahl DJ,
  Hehl R (2012) Integration of bioinformatics and synthetic promoters leads to the
  discovery of novel elicitor-responsive cis-regulatory sequences in Arabidopsis.
  Plant Physiol 160: 178–191
- Kristiansson E, Thorsen M, Tamás MJ, Nerman O (2009) Evolutionary forces act on
   promoter length: Identification of enriched cis-regulatory elements. Mol Biol Evol

#### **687 26**: 1299–1307

- Ksouri N, Jiménez S, Wells CE, Contreras-Moreira B, Gogorcena Y (2016)
   Transcriptional responses in root and leaf of Prunus persica under drought stress
   using RNA sequencing. Front Plant Sci 7: 1–19
- Kumar N, Dale R, Kemboi D, Zeringue EA, Kato N, Larkin JC (2018) Functional
  analysis of short linear motifs in the plant cyclin-dependent kinase inhibitor
  SIAMESE. Plant Physiol 177: 1569–1579
- Langfelder P, Horvath S (2008) WGCNA: An R package for weighted correlation
   network analysis. BMC Bioinformatics 9: 1–13
- Li E, Liu H, Huang L, Zhang X, Dong X, Song W, Zhao H, Lai J (2019) Long-range
   interactions between proximal and distal regulatory regions in maize. Nat Commun
   10: 1–14
- Li J, Zhou D, Qiu W, Shi Y, Yang JJ, Chen S, Wang Q, Pan H (2018) Application
  of weighted gene co-expression network analysis for data from paired design. Sci
  Rep 8: 1–8
- Li X, Jiang J, Zhang L, Yu Y, Ye Z, Wang X, Zhou J, Chai M, Zhang H, Arús P, et
   al (2015) Identification of volatile and softening-related genes using digital gene
   expression profiles in melting peach. Tree Genet Genomes 11: 1–15
- Liseron-Monfils C, Lewis T, Ashlock D, McNicholas PD, Fauteux F, Strömvik M,
   Raizada MN (2013) Promzea: A pipeline for discovery of co-regulatory motifs in
   maize and other plant species and its application to the anthocyanin and
   phlobaphene biosynthetic pathways and the maize development atlas. BMC Plant
   Biol 13: 1–17
- Ma S, Bachan S, Porto M, Bohnert HJ, Snyder M, Dinesh-Kumar SP (2012)
  Discovery of stress responsive DNA regulatory motifs in Arabidopsis. PLoS One
  712 7: 1–13
- Ma S, Shah S, Bohnert HJ, Snyder M, Dinesh-Kumar SP (2013) Incorporating motif
   analysis into gene co-expression networks reveals novel modular expression
   pattern and new signaling pathways. PLoS Genet 9: 1–20

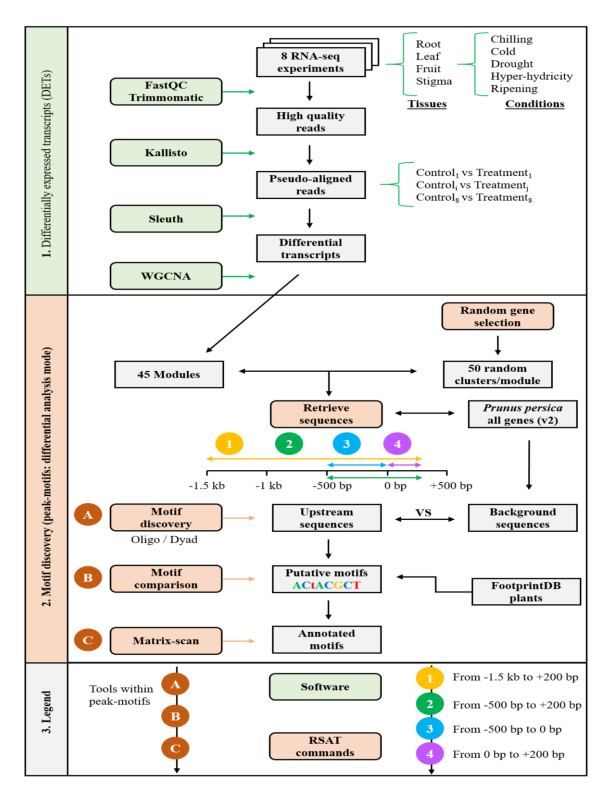
716	Montardit Tardà F (2018) Genomic delimitation of proximal promoter regions: Three						
717	approaches in <i>Prunus persica</i> <u>http://agris.fao.org/agris-</u>						
718	search/search.do?recordID=QC2019600125						
719	Nguyen NTT, Contreras-Moreira B, Castro-Mondragon JA, Santana-Garcia W,						
720	Ossio R, Robles-Espinoza CD, Bahin M, Collombet S, Vincens P, Thieffry D,						
721	et al (2018) RSAT 2018: Regulatory sequence analysis tools 20th anniversary.						
722	Nucleic Acids Res <b>46</b> : 209–214						
723	Petrillo E, Godoy Herz MA, Barta A, Kalyna M, Kornblihtt AR (2014) Let there be						
724	light: Regulation of gene expression in plants. RNA Biol 11: 1215–1220						
725	Pimentel H, Bray NL, Puente S, Melsted P, Pachter L (2017) Differential analysis of						
726	RNA-seq incorporating quantification uncertainty. Nat Methods 1-6						
727	Rombauts S, Déhais P, Van Montagu M, Rouzé P (1999) PlantCARE, a plant cis-						
728	acting regulatory element database. Nucleic Acids Res 27: 295–296						
729	Sanhueza D, Vizoso P, Balic I, Campos-Vargas R, Meneses C (2015) Transcriptomic						
730	analysis of fruit stored under cold conditions using controlled atmosphere in						
731	Prunus persica cv. "Red Pearl." Front Plant Sci 6: 1–12						
732	Scheelbeek PFD, Tuomisto HL, Bird FA, Haines A, Dangour AD (2017) Effect of						
733	environmental change on yield and quality of fruits and vegetables: two systematic						
734	reviews and projections of possible health effects. Lancet Glob Heal 5: 21						
735	Sebastian A, Contreras-Moreira B (2014) FootprintDB: A database of transcription						
736	factors with annotated cis elements and binding interfaces. Bioinformatics 30:						
737	258–265						
738	Smale ST (2001) Core promoters: Active contributors to combinatorial gene regulation.						
739	Genes Dev 15: 2503–2508						
740	Steffens NO, Galuschka C, Schindler M, Bülow L, Hehl R (2005) AthaMap web						
741	tools for database-assisted identification of combinatorial cis-regulatory elements						
742	and the display of highly conserved transcription factor binding sites in						
743	Arabidopsis thaliana. Nucleic Acids Res 33: 397–402						
744	Tanou G, Minas IS, Scossa F, Belghazi M, Xanthopoulou A, Ganopoulos I,						

25

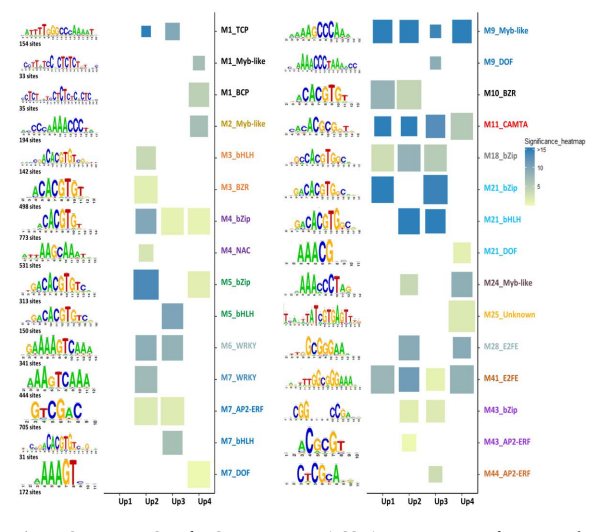
Madesis P, Fernie A, Molassiotis A (2017) Exploring priming responses involved
in peach fruit acclimation to cold stress. Sci Rep 7: 1–14

- Thomas-Chollier M, Darbo E, Herrmann C, Defrance M, Thieffry D, Van Helden
   J (2012) A complete workflow for the analysis of full-size ChIP-seq (and similar)
   data sets using peak-motifs. Nat Protoc 7: 1551–1568
- Tian F, Yang D-C, Meng Y-Q, Jin J, Gao G (2019) PlantRegMap: charting functional
   regulatory maps in plants. Nucleic Acids Res 1: 1–10
- Verde I, Abbott AG, Scalabrin S, Jung S, Shu S, Marroni F, Zhebentyayeva T,
   Dettori MT, Grimwood J, Cattonaro F, et al (2013) The high quality draft
   genome of peach (Prunus persica) identifies unique patterns of genetic diversity,
   domestication and genome evolution. Nat Genet 45: 487–94
- Verde I, Jenkins J, Dondini L, Micali S, Pagliarani G, Vendramin E, Paris R,
  Aramini V, Gazza L, Rossini L, et al (2017) The Peach v2.0 release: highresolution linkage mapping and deep resequencing improve chromosome-scale
  assembly and contiguity. BMC Genomics 18: 1–18
- Wong DCJ, Lopez Gutierrez R, Dimopoulos N, Gambetta GA, Castellarin SD
   (2016) Combined physiological, transcriptome, and cis-regulatory element
   analyses indicate that key aspects of ripening, metabolism, and transcriptional
   program in grapes (Vitis vinifera L.) are differentially modulated accordingly to
   fruit size. BMC Genomics 17: 1–22
- Yu C-P, Chen SC-C, Chang Y-M, Liu W-Y, Lin H-H, Lin J-J, Chen HJ, Lu Y-J,
  Wu Y-H, Lu M-YJ, et al (2015) Transcriptome dynamics of developing maize
  leaves and genomewide prediction of cis elements and their cognate transcription
  factors. Proc Natl Acad Sci 112: 2477–2486
- Yu CP, Lin JJ, Li WH (2016) Positional distribution of transcription factor binding
   sites in Arabidopsis thaliana. Sci Rep 6: 1–7
- Zhu Qun, Dabi T, Lamb C (1995) TATA box and initiator functions in the accurate
   transcription of a plant minimal promoter in vitro. Plant Cell 7: 1681–1689
- 773 Zolotarov Y, Strömvik M (2015) De novo regulatory motif discovery identifies

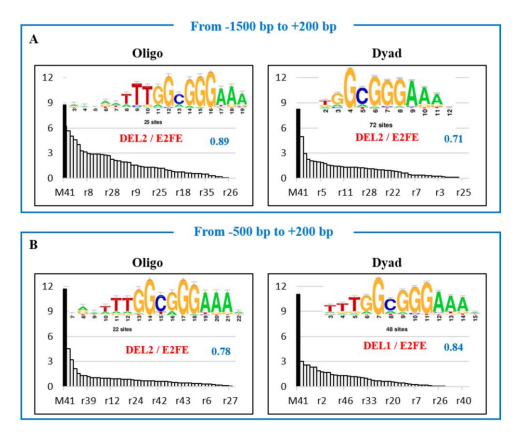
- significant motifs in promoters of five classes of plant dehydrin genes. PLoS One
- 775 **10**: 1–19
- 776 Zou C, Sun K, Mackaluso JD, Seddon AE, Jin R, Thomashow MF, Shiu S-H
- 777 (2011) Cis-regulatory code of stress-responsive transcription in Arabidopsis
- thaliana. PNAS **108**: 14992–14977



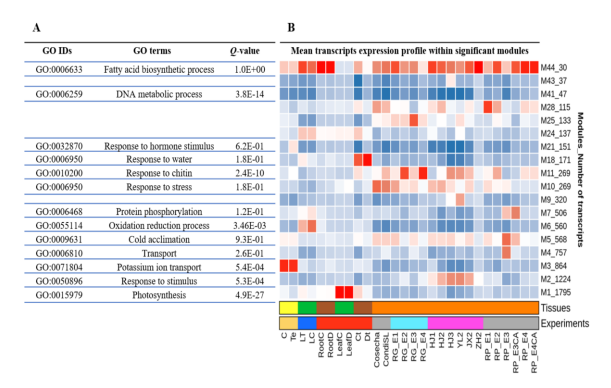
**Figure 1.** Bottom-up framework for *de novo* motif discovery. <u>Step1</u>: differential expression analysis for transcript detection and extraction of co-expressed modules. <u>Step2</u>: *de novo* motif detection using the peak-motifs tool from RSAT::Plants. Numbers correspond to the different tested upstream tracts, with TSSs anchored on position 0 bp, while letters represent tools within peak-motifs. Green and orange boxes label software and RSAT tools, respectively.



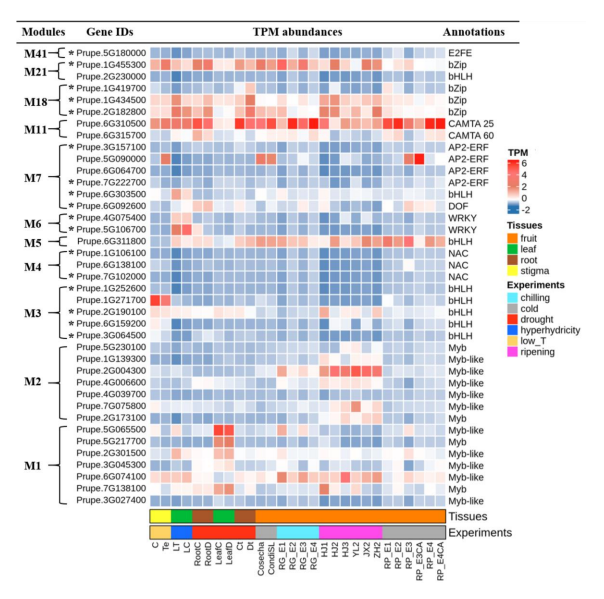
**Figure 2.** Position Specific Scoring Matrix (PSSM) representation of top scored discovered motifs per modules, along different upstream lengths. The x-axis corresponds to the four intervals: Up 1: [-1500 bp, +200 bp], Up 2: [-500 bp, -200 bp], Up 3: [-500 bp, 0 bp] and Up 4 [0 bp, +200 bp]. The y-axis informs about the motif family revealed per module. Cell colors indicate the statistical significance of the identified motifs while cell sizes represent the normalized correlation (Ncor). Number of sites corresponds to the number of sites used to build the PSSM. When motifs from the same family are identified with both algorithms (oligo and dyad-analysis), or in different upstream tracts (Up 1, Up 2, Up 3 and Up 4), only the most significant one is represented in the heatmap. Further details are provided in **Table S3**. An interactive report with source code is accessible at https://eead-csic-compbio.github.io/coexpression\_motif\_discovery/peach/



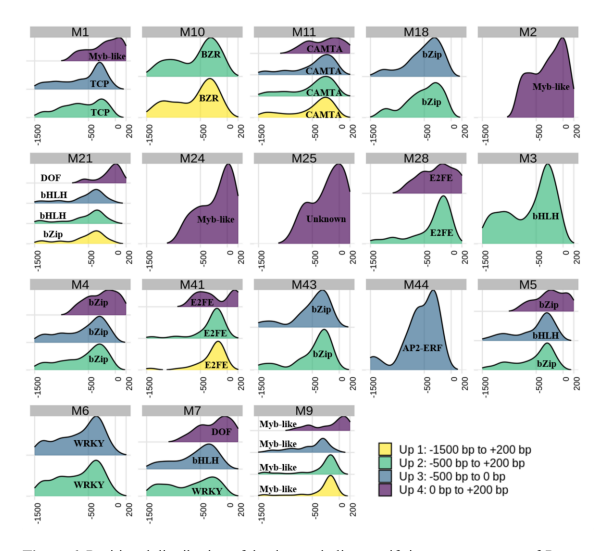
**Figure 3.** Illustrative comparison between predicted motif DEL2 (corresponding to E2FE transcription factor) within two different upstream promoter lengths: -1500 bp to +200 bp ( $\mathbf{A}$ ) and -500 bp to +200 bp ( $\mathbf{B}$ ). The name of the best match among plant motifs in footprintDB is labeled in red, next to its Ncor (Normalized correlation) value labeled in blue. The x-axis corresponds to the module of interest (M41) and random clusters ranked ranked by the most significant motifs. The y-axis corresponds to the statistical significance -log10 (*P*-value). Number of sites corresponds to the occurrence number of a single motif. The evidence supporting the putative motifs is Ncor (in blue) and the significance (black bars) when compared to negative controls (gray bars).



**Figure 4.** Functional annotation of relevant gene modules. **(A):** Gene ontology enrichment. **(B):** Mean transcript abundance profiling in term of transcripts per million (TPM). The x-axis corresponds the different experimental conditions while the y-axis indicates the number of differentially transcripts per module. Experiment and tissue types are highlighted by different colors (see the color key at the bottom of the figure). Gene profiles along the different conditions are provided at (<u>https://eead-csic compbio.github.io/coexpression motif discovery/peach</u>). See supplementary **Table S1** for the abbreviations.



**Figure 5**. List of transcription factors within relevant modules. Blue and red squares indicate transcripts per million while bottom color bars correspond to the tissues types and different experiments, respectively (See the legend at the right side of the figure). TFs showing sequence similarity between their footprintDB and RSAT predicted motifs are labeled with a star.



**Figure 6.** Positional distribution of the detected oligo motifs in promoter genes of *Prunus persica*. Four density distributions were derived from four assessed upstream regions. Up 1: from -1500 bp to 200 bp, Up 2: from -500 bp to +200 bp, Up 3: from -500 bp to 0 bp and Up 4 from 0 bp to + 200 bp. The x-axis corresponds to upstream length in base pairs (bp). The y-axis corresponds to density of captured sites with *P*-value <10 e<sup>-4</sup>. Only oligo motifs are presented here, dyads are provided in the report at <u>https://eead-csiccompbio.github.io/coexpression\_motif\_discovery/peach.</u>

JASPAR TFBS	JASPAR logos	Parameters	Upstream 1	Upstream 2	Upstream 3	Ups tream 4
MA0549.1		Significance		3.44		8.28
BZR	CACGTGG	Logo		<b>CCACGTGG</b>		cCACGTGG
35*		Ncor		0.72		0.73
MA0565.1		Significance				6.52
AP2-B3	TGCATGC	Logo				TGCATGCA
106*		Ncor				0.68
MA0931.1		Significance	32.64	38.66		49.8
bZip	GACACGTG	Logo	tGCCACGTG	tGCCACGTG		tGACGTGGCG
339*		Ncor	0.64	0.49		0.83
MA1197.1		Significance	52.96	66.33	8.32	66.9
CAMTA	aCGCGTg	Logo	ttCACGCG	ttCACGCG	CaCGCGtC	ttCACGCG
351*		Ncor	0.76	0.77	0.89	0.81
MA1224.1		Significance		10.66		20.33
AP2-ERF	GCCGAC	Logo		CGCCGaC		<b>CGCCGaC</b>
470*		Ncor		0.81		0.74
MA1276.1		Significance				1.92
DOF	aaAAAGT	Logo				AAAAGT
223*		Ncor				0.75
MA1289.1		Significance	11.49	17.39		19.18
ТСР	GGGACCAC	Logo	tgGGGaCCACt	gGGGaCCACt		tGGGGACCAC
294*		Ncor	0.84	0.57		0.74
MA1303.1		Significance	14.06	20.46		25.74
WRKY	aaAAGTCAACG	Logo	aaataaaGTCaaCGt	GTtGACtTtt		aAAGTCAACG
303*		Ncor	0.53	0.74		0.8
MA1355.1		Significance	10.38	18.89		26.85
Myb-like	aAACCCTAAtt	Logo	aaCCCtAatta	AaCCCtAac		AACCCTAAttt
231*		Ncor	0.53	0.72		0.65
MA1359.1		Significance	34.74	49.94	6.12	55.41
bHLH	CACGTG	Logo	<b>CCACGTGG</b>	CacGtGcCaCGTG	GaCACGTGtc	cCACGTGGc
258*		Ncor	0.61	0.61	0.61	0.64

**Figure 7.** Similarity between JASPAR motifs (considered as queries) and *de novo* predicted oligo motifs found in *Arabidopsis thaliana* along four different upstream regions. Numbers tagged with a star indicate number of peaks recovered by BLASTN (see Methods). The Ncor scores correspond to JASPAR databases. Only oligo-analysis motifs are shown (dyads are available at supplementary **Table S4**). Upstream 1: [-1500 bp to +200 bp], Upstream 2: [-500 bp to +200 bp], Upstream3: [-500 bp to 0 bp] and Upstream4: [0 bp to +200 bp]

# **Parsed Citations**

Abbott AG, Georgi L, Yvergniaux D, Wang Y, Blenda A, Reighard G, Inigo M, Sosinski B (2002) Peach: The model genome for Rosaceae. Acta Hortic 575: 145–155

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Bakir Y, Eldem V, Zararsiz G, Unver T (2016) Global transcriptome analysis reveals differences in gene expression patterns between nonhyperhydric and hyperhydric peach leaves. Plant Genome 9: 1–9

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Baldoni E, Genga A, Cominelli E (2015) Plant MYB transcription factors: Their role in drought response mechanisms. Int J Mol Sci 16: 15811–15851

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Bianchi VJ, Rubio M, Trainotti L, Verde I, Bonghi C, Martínez-Gómez P (2015) Prunus transcription factors: breeding perspectives. Front Plant Sci 6: 1–20

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Bolger AM, Lohse M, Usadel B (2014) Trimmomatic: A flexible trimmer for Illumina sequence data. Bioinformatics 30: 2114–2120 Pubmed: <u>Author and Title</u>

Google Scholar: Author Only Title Only Author and Title

Bray NL, Pimentel H, Melsted P, Pachter L (2016) Near-optimal probabilistic RNA-seq quantification. Nat Biotechnol 34: 525–528 Pubmed: <u>Author and Title</u>

Google Scholar: Author Only Title Only Author and Title

Cantalapiedra CP, García-pereira MJ, Gracia MP, Igartua E (2017) Large differences in gene expression responses to drought and heat stress between elite Barley cultivar scarlett and a spanish landrace. Front Plant Sci 8: 1–23

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Carrillo Bermejo EA, Alamillo MAH, Samuel David GT, Llanes MAK, Enrique C de la S, Manuel RZ, Rodriguez Zapata LC (2017) Transcriptome, genetic transformation and micropropagation: Some biotechnology strategies to diminish water stress caused by climate change in sugarcane. Plant, Abiotic Stress Responses to Clim. Chang. IntechOpen, pp 90–108

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Chang WC, Lee TY, Huang H Da, Huang HY, Pan RL (2008) PlantPAN: Plant promoter analysis navigator, for identifying combinatorial cis-regulatory elements with distance constraint in plant gene groups. BMC Genomics 9: 1–14

Pubmed: <u>Author and Title</u> Google Scholar: Author Only Title Only Author and Title

Cherenkov P, Novikova D, Omelyanchuk N, Levitsky V, Grosse I, Weijers D, Mironova V (2018) Diversity of cis-regulatory elements associated with auxin response in Arabidopsis thaliana. J Exp Bot 69: 329–339

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Contreras-Moreira B, Castro-Mondragon JA, Rioualen C, Cantalapiedra CP, Van Helden J (2016) RSAT::Plants: Motif discovery within clusters of upstream sequences in plant genomes. In R Hehl, ed, Plant Synth. Promot. Methods Mol. Biol. Humana Press, New York, pp 279–295

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Defrance M, Janky R, Sand O, van Helden J (2008) Using RSAT oligo-analysis and dyad-analysis tools to discover regulatory signals in nucleic sequences. Nat Protoc 3: 1589–1603

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Doherty CJ, Van Buskirk HA, Myers SJ, Thomashow MF (2009) Roles for Arabidopsis CAMTA transcription factors in cold-regulated gene expression and freezing tolerance. Plant Cell 21: 972–984

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Fornes O, Castro-Mondragon JA, Khan A, van der Lee R, Zhang X, Richmond PA, Modi BP, Correard S, Gheorghe M, Baranašić D, et al (2020) JASPAR 2020: update of the open-access database of transcription factor binding profiles. Nucleic Acids Res 48: 87–92 Pubmed: Author and Title

Google Scholar: <u>Author Only Title Only Author and Title</u>

Franco-Zorrilla JM, López-Vidriero I, Carrasco JL, Godoy M, Vera P, Solano R (2014) DNA-binding specificities of plant transcription factors and their potential to define target genes. PNAS 111: 2367–2372

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Gismondi M, Daurelio LD, Maiorano C, Monti LL, Lara M V., Drincovich MF, Bustamante CA (2020) Generation of fruit postharvest gene datasets and a novel motif analysis tool for functional studies: uncovering links between peach fruit heat treatment and cold storage responses. Planta 251: 1–18

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Gogorcena Y, Sánchez G, Moreno-vázquez S, Pérez S, Ksouri N (2020) Genomic-based breeding for climate-smart peach varieties. In C Kole, ed, Genome Des. Clim. fruit Crop. Springer-Nature, pp 291–351

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Guo J, Chen J, Yang J, Yu Y, Yang Y, Wang W (2018) Identification, characterization and expression analysis of the VQ motif-containing gene family in tea plant (Camellia sinensis). BMC Genomics 19: 1–12

Pubmed: Author and Title Google Scholar: Author Only Title Only Author and Title

Howe KL, Contreras-moreira B, Silva N De, Maslen G, Akanni W, Allen J, Alvarez-jarreta J, Barba M, Bolser DM, Cambell L, et al (2020) Ensembl Genomes 2020 enabling non-vertebrate genomic research. Nucleic Acids Res 1–7

Pubmed: Author and Title Google Scholar: Author Only Title Only Author and Title

Hu P, Li G, Zhao X, Zhao F, Li L, Zhou H (2018) Transcriptome profiling by RNA-Seq reveals differentially expressed genes related to fruit development and ripening characteristics in strawberries (Fragaria × ananassa). Peer J 6: 1–25

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Jiao Y, Shen Z, Yan J (2017) Transcriptome analysis of peach [Prunus persica (L.) Batsch] stigma in response to low-temperature stress with digital gene expression profiling. J Plant Biochem Biotechnol 26: 141–148

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Korkuc P, Schippers JHM, Walther D (2014) Characterization and identification of cis-regulatory elements in Arabidopsis based on single-nucleotide polymorphism information. Plant Physiolgy 164: 181–200

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Koschmann J, Machens F, Becker M, Niemeyer J, Schulze J, Bulow L, Stahl DJ, Hehl R (2012) Integration of bioinformatics and synthetic promoters leads to the discovery of novel elicitor-responsive cis-regulatory sequences in Arabidopsis. Plant Physiol 160: 178–191

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Kristiansson E, Thorsen M, Tamás MJ, Nerman O (2009) Evolutionary forces act on promoter length: Identification of enriched cisregulatory elements. Mol Biol Evol 26: 1299–1307

Pubmed: <u>Author and Title</u> Google Scholar. <u>Author Only Title Only Author and Title</u>

Ksouri N, Jiménez S, Wells CE, Contreras-Moreira B, Gogorcena Y (2016) Transcriptional responses in root and leaf of Prunus persica under drought stress using RNA sequencing. Front Plant Sci 7: 1–19

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Kumar N, Dale R, Kemboi D, Zeringue EA, Kato N, Larkin JC (2018) Functional analysis of short linear motifs in the plant cyclindependent kinase inhibitor SIAMESE. Plant Physiol 177: 1569–1579

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Langfelder P, Horvath S (2008) WGCNA: An R package for weighted correlation network analysis. BMC Bioinformatics 9: 1–13

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Li E, Liu H, Huang L, Zhang X, Dong X, Song W, Zhao H, Lai J (2019) Long-range interactions between proximal and distal regulatory regions in maize. Nat Commun 10: 1–14

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Li J, Zhou D, Qiu W, Shi Y, Yang JJ, Chen S, Wang Q, Pan H (2018) Application of weighted gene co-expression network analysis for data from paired design. Sci Rep 8: 1–8

Pubmed: Author and Title Google Scholar: Author Only Title Only Author and Title

Li X, Jiang J, Zhang L, Yu Y, Ye Z, Wang X, Zhou J, Chai M, Zhang H, Arús P, et al (2015) Identification of volatile and softening-related

genes using digital gene expression profiles in melting peach. Tree Genet Genomes 11: 1-15

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Liseron-Monfils C, Lewis T, Ashlock D, McNicholas PD, Fauteux F, Strömvik M, Raizada MN (2013) Promzea: A pipeline for discovery of co-regulatory motifs in maize and other plant species and its application to the anthocyanin and phlobaphene biosynthetic pathways and the maize development atlas. BMC Plant Biol 13: 1–17

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Ma S, Bachan S, Porto M, Bohnert HJ, Snyder M, Dinesh-Kumar SP (2012) Discovery of stress responsive DNA regulatory motifs in Arabidopsis. PLoS One 7: 1–13

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Ma S, Shah S, Bohnert HJ, Snyder M, Dinesh-Kumar SP (2013) Incorporating motif analysis into gene co-expression networks reveals novel modular expression pattern and new signaling pathways. PLoS Genet 9: 1–20

Pubmed: <u>Author and Title</u> Google Scholar: Author Only Title Only Author and Title

Montardit Tardà F (2018) Genomic delimitation of proximal promoter regions: Three approaches in Prunus persica http://agris.fao.org/agris-search/search.do?recordID=QC2019600125

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Nguyen NTT, Contreras-Moreira B, Castro-Mondragon JA, Santana-Garcia W, Ossio R, Robles-Espinoza CD, Bahin M, Collombet S, Vincens P, Thieffry D, et al (2018) RSAT 2018: Regulatory sequence analysis tools 20th anniversary. Nucleic Acids Res 46: 209–214

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Petrillo E, Godoy Herz MA, Barta A, Kalyna M, Kornblihtt AR (2014) Let there be light: Regulation of gene expression in plants. RNA Biol 11: 1215–1220

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Pimentel H, Bray NL, Puente S, Melsted P, Pachter L (2017) Differential analysis of RNA-seq incorporating quantification uncertainty. Nat Methods 1–6

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Rombauts S, Déhais P, Van Montagu M, Rouzé P (1999) PlantCARE, a plant cis-acting regulatory element database. Nucleic Acids Res 27: 295–296

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Sanhueza D, Vizoso P, Balic I, Campos-Vargas R, Meneses C (2015) Transcriptomic analysis of fruit stored under cold conditions using controlled atmosphere in Prunus persica cv. "Red Pearl." Front Plant Sci 6: 1–12

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Scheelbeek PFD, Tuomisto HL, Bird FA, Haines A, Dangour AD (2017) Effect of environmental change on yield and quality of fruits and vegetables: two systematic reviews and projections of possible health effects. Lancet Glob Heal 5: 21

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only</u> <u>Author and Title</u>

Sebastian A, Contreras-Moreira B (2014) FootprintDB: A database of transcription factors with annotated cis elements and binding interfaces. Bioinformatics 30: 258–265

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Smale ST (2001) Core promoters: Active contributors to combinatorial gene regulation. Genes Dev 15: 2503-2508

Pubmed: Author and Title Google Scholar: Author Only Title Only Author and Title

Steffens NO, Galuschka C, Schindler M, Bülow L, Hehl R (2005) AthaMap web tools for database-assisted identification of combinatorial cis-regulatory elements and the display of highly conserved transcription factor binding sites in Arabidopsis thaliana. Nucleic Acids Res 33: 397–402

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Tanou G, Minas IS, Scossa F, Belghazi M, Xanthopoulou A, Ganopoulos I, Madesis P, Fernie A, Molassiotis A (2017) Exploring priming responses involved in peach fruit acclimation to cold stress. Sci Rep 7: 1–14

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u> Thomas-Chollier M, Darbo E, Herrmann C, Defrance M, Thieffry D, Van Helden J (2012) A complete workflow for the analysis of full-size ChIP-seq (and similar) data sets using peak-motifs. Nat Protoc 7: 1551–1568

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Tian F, Yang D-C, Meng Y-Q, Jin J, Gao G (2019) PlantRegMap: charting functional regulatory maps in plants. Nucleic Acids Res 1: 1–10 Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Verde I, Abbott AG, Scalabrin S, Jung S, Shu S, Marroni F, Zhebentyayeva T, Dettori MT, Grimwood J, Cattonaro F, et al (2013) The high quality draft genome of peach (Prunus persica) identifies unique patterns of genetic diversity, domestication and genome evolution. Nat Genet 45: 487–94

Pubmed: Author and Title Google Scholar: Author Only Title Only Author and Title

Verde I, Jenkins J, Dondini L, Micali S, Pagliarani G, Vendramin E, Paris R, Aramini V, Gazza L, Rossini L, et al (2017) The Peach v2.0 release: high-resolution linkage mapping and deep resequencing improve chromosome-scale assembly and contiguity. BMC Genomics 18: 1–18

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Wong DCJ, Lopez Gutierrez R, Dimopoulos N, Gambetta GA, Castellarin SD (2016) Combined physiological, transcriptome, and cisregulatory element analyses indicate that key aspects of ripening, metabolism, and transcriptional program in grapes (Vitis vinifera L.) are differentially modulated accordingly to fruit size. BMC Genomics 17: 1–22

Pubmed: Author and Title Google Scholar: Author Only Title Only Author and Title

Yu C-P, Chen SC-C, Chang Y-M, Liu W-Y, Lin H-H, Lin J-J, Chen HJ, Lu Y-J, Wu Y-H, Lu M-YJ, et al (2015) Transcriptome dynamics of developing maize leaves and genomewide prediction of cis elements and their cognate transcription factors. Proc Natl Acad Sci 112: 2477–2486

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Yu CP, Lin JJ, Li WH (2016) Positional distribution of transcription factor binding sites in Arabidopsis thaliana. Sci Rep 6: 1–7

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Zhu Qun, Dabi T, Lamb C (1995) TATA box and initiator functions in the accurate transcription of a plant minimal promoter in vitro. Plant Cell 7: 1681–1689

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Zolotarov Y, Strömvik M (2015) De novo regulatory motif discovery identifies significant motifs in promoters of five classes of plant dehydrin genes. PLoS One 10: 1–19

Pubmed: <u>Author and Title</u> Google Scholar: <u>Author Only Title Only Author and Title</u>

Zou C, Sun K, Mackaluso JD, Seddon AE, Jin R, Thomashow MF, Shiu S-H (2011) Cis-regulatory code of stress-responsive transcription in Arabidopsis thaliana. PNAS 108: 14992–14977

Pubmed: <u>Author and Title</u> Google Scholar: Author Only Title Only Author and Title