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Comparative transcriptomics in ferns reveals key innovations and divergent evolution of the secondary cell walls

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1 **Title:** Comparative transcriptomics in ferns reveals key innovations and divergent
2 evolution of the secondary cell walls

3

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35

36 **Abstract**

37 Despite ferns being crucial to understanding plant evolution, their large and complex
38 genomes has kept their genetic landscape largely uncharted, with only a handful of
39 genomes sequenced and sparse transcriptomic data. Addressing this gap, we generated
40 extensive RNA-sequencing data for multiple organs across 22 representative species
41 over the fern phylogeny, assembling high-quality transcriptomes. These data facilitated
42 the construction of a time-calibrated fern phylogeny covering all major clades, revealing
43 numerous whole-genome duplications and highlighting the uniqueness of fern genetics,
44 with half of the uncovered gene families being fern-specific. Our investigation into fern
45 cell walls through biochemical and immunological analyses identified occurrences of the
46 lignin syringyl unit and its independent evolution in ferns. Moreover, the discovery of an
47 unusual sugar in fern cell walls hints at a divergent evolutionary path in cell wall
48 biochemistry, potentially driven by gene duplication and sub-functionalization. We provide
49 an online database preloaded with genomic and transcriptomic data for ferns and other
50 land plants, which we used to identify an independent evolution of lignocellulosic gene
51 modules in ferns. Our data provide a framework for the unique evolutionary path that ferns
52 have navigated since they split from the last common ancestor of euphyllophytes more
53 than 360 million years ago.

54

55 **Introduction**

56 Since they diverged from a shared ancestor with seed plants more than 360 million years
57 ago, ferns have played a significant role in life on Earth ¹. They occupy various niches in
58 different ecosystems, acting as pioneer species, key ecological players, invasive entities,
59 and contributors to agriculture. They are the second most diverse group of vascular plants
60 after angiosperms, with over 10,500 existing species ²⁻⁶. Ferns exhibit great
61 morphological and physiological diversity, and have evolved equally diverse strategies to

62 cope with environmental challenges ⁷, such as adaptations to low-light environments ⁸.
63 The secondary metabolites produced by ferns and the genes responsible for their
64 biosynthesis are of great interest for environmental clean-up efforts, agriculture, and the
65 discovery of new pharmaceuticals ^{9–11}.

66 Despite fern's ecological importance, the phylogenetic relationship of major clades
67 in Monilophyta (ferns) remains elusive ¹². Based on a Maximum Likelihood (ML) tree of a
68 concatenated matrix of 146 low-copy nuclear genes, Qi and coauthors ¹³ inferred
69 Marattiales to be sister to Polypodiidae (i.e. the leptosporangiate ferns) as proposed in
70 Pteridophyte Phylogeny Group (PPG) I (2016). Conversely, Shen and coauthors ¹²
71 inferred Marattiales to be sister to Ophioglossidae (consisting of Psilotales and
72 Ophioglossales), based on a coalescent-based tree of two low-copy nuclear gene sets of
73 69 transcriptomes. Nitta and coauthors ¹⁴ inferred Gleicheniales as a monophyletic clade
74 based on a ML tree of a concatenated matrix of 79 plastome loci as opposed to the
75 paraphyletic inference of Shen and coauthors ¹² and Qi and coauthors ¹³. It is also unclear
76 whether horsetails are a sister group to the last common ancestor of all the remaining
77 ferns or other fern clades, such as Marattiales ^{5,12}.

78 Given ferns' critical evolutionary position as the sister group to seed plants,
79 investigating their genomes, coding sequences, and gene families offers unparalleled
80 insights into the evolution of plants ¹⁵, especially the key aspects of vasculature and cell
81 walls. The evolution of vasculature and secondary cell walls precipitated a 10-fold
82 increase in plant species numbers (<http://www.theplantlist.org/>) and shaped the Earth's
83 geo- and biosphere ¹⁶. Ferns thus harbour key information for the evolution of vascular
84 plant form and function ¹⁷.

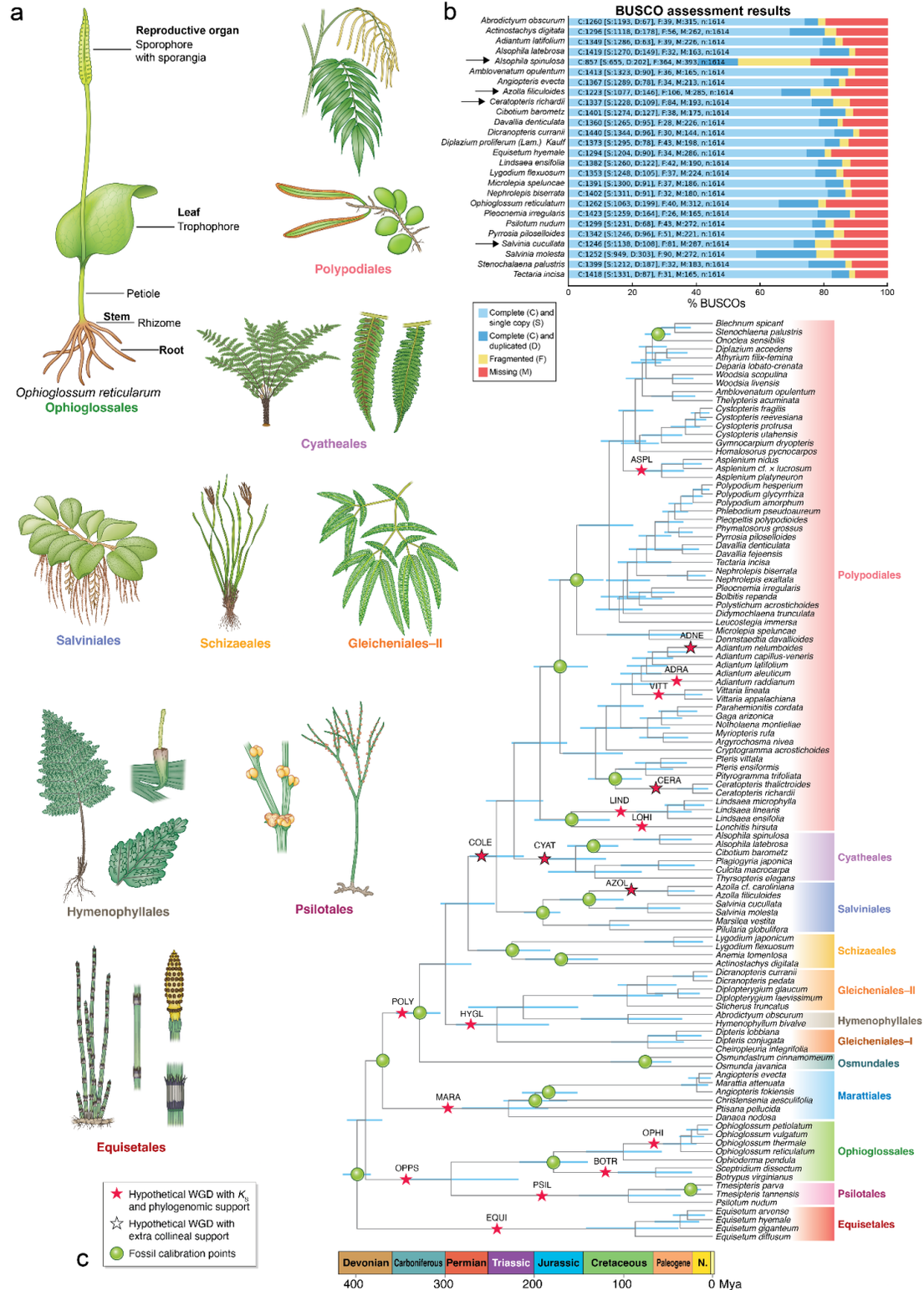
85 However, ferns are infamous for their exceptionally large genomes (on average
86 12.3 billion base pairs), with one of the largest genome of any living organism - 160 billion
87 base pairs - found in ferns¹⁸. They also have exceptionally high numbers of chromosomes
88 (averaging at 40.5, with a peak at 720)¹⁹, which are believed to result from multiple
89 instances of whole-genome duplication ^{20–22} and a relatively slow genome downsizing
90 process ²³. Among plants, ferns show the highest rate of polyploidy-driven speciation ²⁴,
91 a direct relationship between genome size, chromosome number and the age of long
92 terminal repeat-retrotransposon (LTR-RT) insertions ^{25–27}, and a high rate of whole

93 genome duplications (WGDs) among several fern lineages^{23,28}. However, the genetic
94 and genomic evidence for widespread whole-genome duplication in ferns remains largely
95 unexplored^{29–31}.

96 Thus, comparative studies that investigate the evolution of ferns have been
97 hampered by their large, complex genomes, limiting our understanding of fern genome
98 evolution and the genetic underpinnings of the evolution of vasculature and cell walls. To
99 date, only few fern genomes and transcriptomes are available^{20,32–35}, and no studies that
100 conducted a comprehensive comparison of gene inventories, transcriptional programs
101 and biochemical properties of their cell walls have been reported.

102 To address this, we generated 405 RNA-sequencing samples to generate coding
103 sequences and gene expression atlases for 22 fern species, capturing major
104 representatives of ten fern orders. We investigated ancient polyploidy, the distribution of
105 fern-specific gene families, how gene age correlates with organ-specific expression, and
106 predicted the functions of fern-specific genes. To better understand how fern cell walls
107 have evolved, we performed a comprehensive histological and biochemical analysis of
108 fern tissues and propose a novel biosynthetic pathway of lignocellulose. We further
109 detected a wide-spread occurrence of the unusual hemicellulose mixed-linkage glucan,
110 and show that it evolved independently in ferns, and propose candidate
111 glucosyltransferases responsible for its synthesis. We also detected a novel type of a
112 methylated sugar, a 2-O-Methyl-D-glucopyranose. We also show that ferns likely
113 independently evolved secondary cell walls through several duplication events in the
114 cellulose synthase family. Finally, we make our fern genomic and transcriptomic data
115 easily accessible with the CoNekt database (<https://conekt.plant.tools/>).

116 Our data, findings, and tools shed light on the evolution of cell walls, lignin,
117 specialised metabolism, and organ development in ferns and other land plants. We
118 envision that similar large, comparative studies will elucidate the evolution of plants and
119 other organisms.



120

121 **Figure 1. Sampling, transcriptome assembly, and species tree of major**
 122 **representatives of ferns. a) *Ophioglossum reticulatum* with sampled organs labelled,**
 123 **together with samples representative of the other fern ordersb) Completeness of**

124 transcriptome assembly measured by Benchmarking Universal Single-Copy Orthologs
125 (BUSCO). *Alsophila spinulosa*, *Ceratopteris richardii*, *Azolla filiculoides* and *Salvinia*
126 *cucullata* have available genomes, while the remaining values are for the transcriptome
127 assemblies reported here. C) The evolutionary timescale of Monilophyta based on the
128 inferred consensus fern cladogram. The species tree shows the inferred consensus
129 phylogenetic topology with branch lengths representing absolute divergence time
130 estimated by Bayesian molecular dating analysis. The horizontal coordinates of each
131 internal node denotes the posterior mean divergence time while the bars represent the
132 95% Highest Posterior Density. Hypothetical WGDs are indicated at corresponding
133 phylogenetic nodes as red stars with four-letter identifiers. Black outline around a red star
134 indicates events with additional collinear support. Fossil calibrations are indicated at
135 corresponding phylogenetic nodes with green circles. The clade strips indicating
136 affiliations at order level are shown as vertical bars with distinct colours. The geological
137 timeline refers to the International Commission on Stratigraphy (ICS) v2023/09.
138

139 **Results**

140 **Construction of fern coding sequences by transcriptome assembly**

141 To capture the diversity of ferns, we selected 22 candidate species representing ten
142 orders, which were photographed and dissected on site (Figure S1), with organs
143 categorised with localities and vouchers attached (Table S1). We collected 405 RNA-seq
144 samples (Table S2), capturing 25 specific organs at different developmental stages,
145 categorised into four major organs - leaves, roots, stems, and reproductive organs - for
146 simplified comparison (Figure 1a)(Table S1). Our transcriptome assembly pipeline
147 combined TRINITY and k-mer SOAPdenovo-Trans assemblies concatenated with
148 EvidentialGene (Figure S2)^{36,37}. We removed any potential non-fern mRNA contaminants
149 and any sequences with aberrant GC content due to assembly artefacts, low transcripts
150 per million (TPM) values, or sequence similarity higher to non-fern species than to ferns
151 (see methods, Figure S2). The assembly yielded 30,000–100,000 coding sequences
152 (CDSs) per species with high Benchmarking Universal Single-Copy Orthologs (BUSCO)
153 scores (Figure 1b, Table S3) that rivalled the scores of the four sequenced genomes
154 (Figure 1b, black arrows).
155

156 **Reconstruction of the evolutionary timeline of ferns**

157 Given the standing phylogenetic discordance, we reconstructed a phylogenetic tree of
158 108 fern species (22 from this study, 7 sequenced genomes, and 79 from the 1000 Plant
159 Transcriptomes Initiative (1KP) and other studies)^{13,20,32–35,38–40}, covering the whole

160 backbone of Monilophyta (Table S4, Supplemental Methods 1). Four datasets, each with
161 a different outgroup (horsetails, seed plants, lycopods or bryophytes), were first used in
162 nucleotide with three different methods including ASTRAL-Pro2 ⁴¹, concatenation-based
163 method and STAG ⁴²(Supplemental Methods 1) on the 107 ferns dataset and then
164 reanalyzed in both nucleotide and peptide on the 108 ferns (adding the latest *Marsilea*
165 *vestita* genome ³⁹) using the favored method ASTRAL-Pro2. We recovered a well-
166 supported fern backbone phylogeny in which Marattiales was inferred to be sister to
167 Polypodiidae (i.e. leptosporangiate ferns) and Gleicheniales was inferred as a
168 paraphyletic clade (Supplemental Methods 1, Figure SM5-6). A closer phylogenetic
169 relationship between Marattiales and Polypodiidae than between Marattiales and
170 Ophioglossidae was supported in all datasets with Local Posterior Probability (LPP) as
171 1.00, except the one with bryophytes as outgroup based on peptide alignment.
172 Phylogenies derived from this dataset favored Marattiales and Ophioglossidae as sister
173 groups, with LPP as 0.82 (Supplemental Methods 1, Figure SM6). The monophyly of
174 Gleicheniales was not supported, whereas the Gleicheniales-II (Gleicheniaceae) was
175 closer to Hymenophyllales than Gleicheniales-I (Dipteridaceae) in all datasets, except
176 the one with bryophytes as outgroup based on nucleotide alignment, whose conflicting
177 branching pattern was only supported by LPP as 0.46 (Supplemental Methods 1, Figure
178 SM5).

179 In addition, these analyses provided strong support for a scenario wherein
180 horsetails are a sister group to the last common ancestor of all the remaining ferns in the
181 phylogeny inferred from every combinational setting of non-horsetails outgroups and
182 methods (Supplemental Methods 1). We selected phylogeny derived from the nucleotide
183 dataset with horsetails as outgroup using ASTRAL-Pro2 as the consensus tree and used
184 in all our subsequent analyses (Figure 1c). This selection was based on the general
185 consistency across datasets with varied outgroups, and on the ASTRAL-Pro2 derived
186 phylogeny with STAG method (Supplemental Methods 1).

187 To estimate the absolute divergence time of the 108 ferns, we used Bayesian
188 molecular dating under the independent rate and LG general amino acid substitution

189 model⁴³ with 18 soft fossil constraints (indicated in Figure 1c as green dots, Table S5).
190 The 95% Highest Posterior Density (HPD) and posterior mean of the stem ages of major
191 fern clades was summarised (Supplemental Methods 1, Table S6). The resulting high-
192 confidence fossil-calibrated tree thus resolved a long-standing discussion on fern
193 phylogenetics.

194

195 **Identification of 18 separate whole genome duplication events in ferns**

196 It has been proposed that ferns have a large number of chromosomes due to repeated
197 rounds of whole-genome duplication (WGD)⁴⁴. Here, we used K_S -age distributions and
198 phylogenomic methods to unveil remnants of ancient WGDs⁴⁵. In total, we found support
199 for 18 hypothetical WGDs within the backbone of ferns, of which five with attained
200 collinear support (red stars, Figure 1c, see Supplemental Methods 1). Seven of the WGDs
201 were found within Polypodiales, two in Ophioglossales, and one in each of the lineages
202 of Equisetales, Psilotales, Marattiales, Salviniiales and Cyatheaales, together with four
203 shared by more than one order. WGDs were previously identified in other studies with
204 different phylogenetic locations^{28,46}. We reevaluated possible scenarios thereof and
205 proposed WGDs with both K_S and phylogenomic support after correcting rate variation
206 and taking into account the uncertainty in gene tree and gene tree-species tree
207 reconciliation (Supplemental Methods 1). Next, we tested the species richness and
208 genome size as a function of the number of ancient WGD events shared by different major
209 clades (excluding Polypodiales due to numerous nested WGD events therein), and
210 observed a significant positive correlation between the number of ancient WGD events
211 and the number of species in a lineage (Figure S3, adjusted $R^2 = 0.41$, p-value = 0.02).
212 Conversely, we did not observe any significant correlation between the number of ancient
213 WGD events and genome size (Figure S3).

214

215 **Diversity and conservation of fern gene functions and expression patterns**

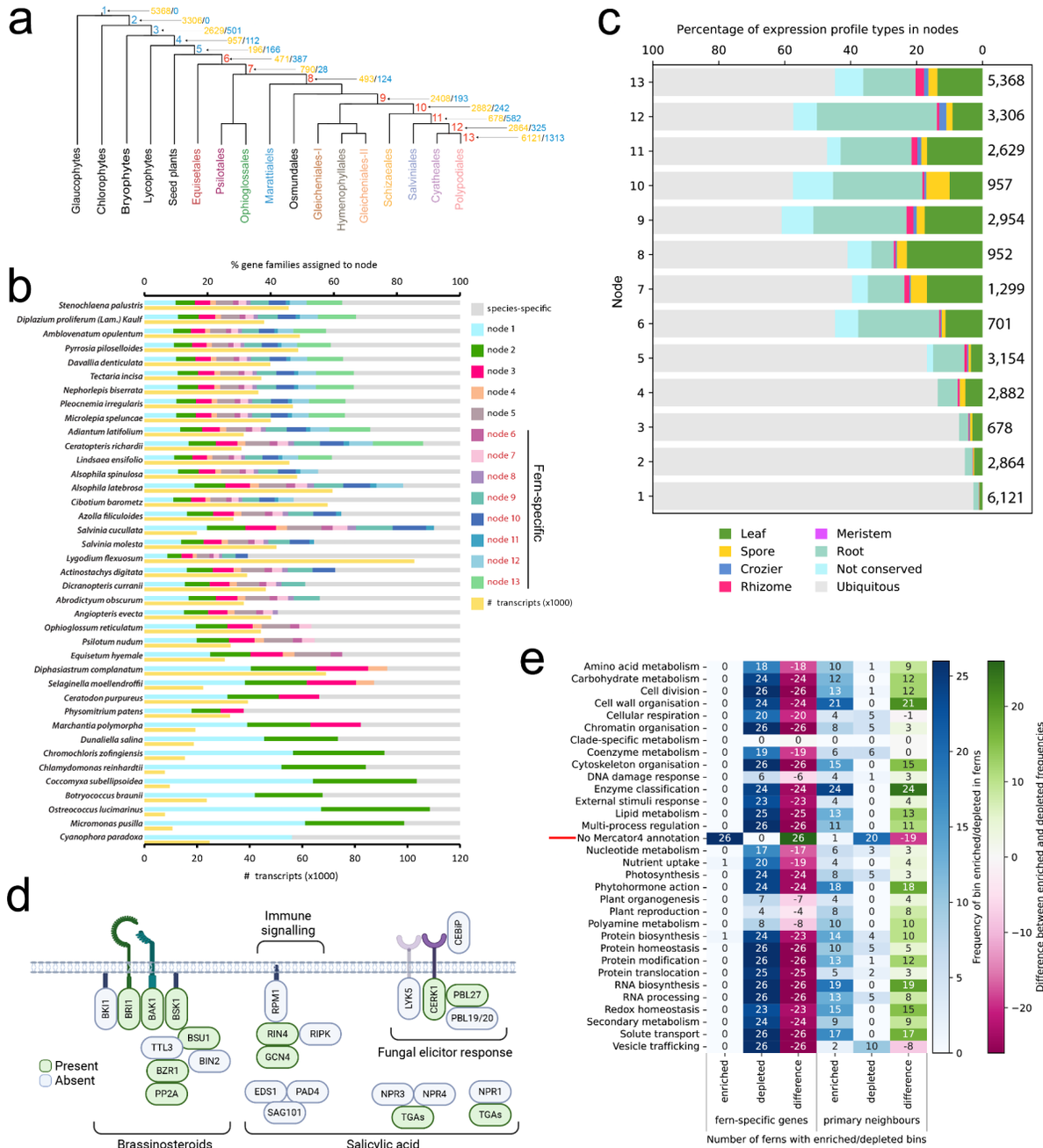
216 To investigate predicted gene functions within ferns, we conducted a phylostratigraphic
217 analysis encompassing one glaucophyte, seven chlorophytes, three bryophytes, two
218 lycophytes, 26 ferns (22 fern transcriptomes and four genomes), two gymnosperms, and
219 six angiosperms (see methods). The 47 species are used to study the gain and losses of

220 gene families across Archaeplastida, by assigning the orthogroups to nodes ranging from
221 node 1 (the earliest ancestor of Archaeplastida) to node 13 (the ancestor of
222 Polypodiales)(Figure 2a). The nodes are based on the fern phylogeny tree shown in
223 Figure 1c and known relationships between Archaeplastida ⁴⁷.

224 Orthogroup gains, a measure of new gene family acquisition, were highest in the
225 early stages of algae evolution (nodes 1 and 2 gained 5368 and 3306 orthogroups,
226 respectively) and when plants colonized the land (node 3, 2629 gained
227 orthogroups)(Figure 2a). However, a substantial number of orthogroup gains and losses
228 were also observed within the different fern lineages (e.g., 2408 gained and 193 lost
229 orthogroups in node 9, Figure 2a).

230 The analysis further revealed that ~50% of fern gene families are fern-specific
231 (Figure 2b nodes 6–13 in red, Table S7). For example, >50% of gene families in
232 *Stenochlaena palustris* belong to nodes 6–13 (fern-specific nodes 6–13 in red, Figure 2a,
233 Table S7), suggesting that the fern lineage has evolved genes with novel, unexplored
234 functions. We analysed sequences of 25 orthogroups comprising at least ten fern species.
235 Representatives of 17 orthogroups show no significant sequence or structural similarity
236 to non-fern species (Figure S4), with eight of them being disordered (few to no secondary
237 structures). These results show that >50% of gene families in ferns represent novel,
238 uncharacterized proteins, indicating that studying ferns will likely provide new insights into
239 plant biology and evolution.

240 To understand how gene age correlates with gene expression specificity, we
241 identified organ-specific genes with specificity measure (SPM) analysis (distributions of
242 SPM values are shown in Figure S5, expression profiles of organ-specific genes are
243 shown in Figure S6, Table S8)⁴⁷. Genes belonging to older nodes (nodes 1–4) were less
244 organ-specific (<20%) than younger nodes (nodes 5–13, 30–60%, Figure 2c). Younger
245 genes (nodes 5–13) had specialised functions in a particular organ, with roots having the
246 highest number of specifically expressed genes (Figure 2c). Furthermore, species-
247 specific genes also tended to show a less ubiquitous, more organ-specific expression
248 (Figure S7), indicating an overall negative association between gene age and organ
249 specificity, which is in line with similar observation in land plants ⁴⁷.



250

251 **Figure 2. Gene functions in ferns.** a) Division tree of Archaeplastida. Leaves represent
 252 orders, while node numbers correspond to the phylostrata. The orange and blue numbers
 253 indicate the number of gains and losses of orthogroups, respectively. b) Stacked bar plot
 254 showing the percentage of gene families belonging to a phylostrata (node). Species-
 255 specific gene families comprised genes from only one species and were not assigned to
 256 nodes. c) Percentage of gene families identified as organ-specific for each node.
 257 Numbers on the right side of the plot indicate the number of orthogroups per node. d)
 258 Examples of signalling components present (green shapes) or absent (grey shapes) in
 259 ferns. e) Clustered heatmap showing enrichment and depletion of biological processes of

260 fern-specific genes and primary co-expression neighbours in 26 (22 from this study and
261 four sequenced genomes) fern species analysed. The left column indicates the biological
262 processes defined by Mapman bins. The scores in the 'enriched' and 'depleted' column
263 indicate in how many of the 26 species fern-specific genes are significantly (BH adj.p <
264 0.05), connected or disconnected to genes belonging to a specific bin, respectively. A
265 higher value in difference (# enriched - # depleted) indicates overall enrichment, while
266 lower values indicate overall depletion for respective bins in the 26 ferns.

267

268 Finally, we investigated whether organ-specific genes are conserved across ferns
269 and other land plants. While many organs express significantly similar sets of genes
270 across ferns and other land plants, we observed a clear difference between fern and seed
271 plant transcriptomes (Figure S8). Not surprisingly, the organ-specific gene sets of seed-
272 containing plants show higher mutual similarity to those of other seed plants (Figure S8,
273 blue box), while those of ferns show the highest similarity to those of other ferns (Figure
274 S8, green box).

275

276 **Ferns lack several genes essential for hormonal signalling, defence and**
277 **development in angiosperms, indicating their unique developmental and**
278 **environmental strategies**

279 Terrestrialization and the evolution of seeds and flowers required the evolution of many
280 biological functions⁴⁸, which is readily visible when comparing gene inventories of algae,
281 land, seed and flowering plants (Figure S9, gene function completeness indicated by
282 darker cells). We used MapMan sequence-based annotations and compared the gene
283 function repertoire of ferns and model angiosperms and observed the absence of several
284 components in ferns (missing functional categories indicated with red text in Table S9,
285 Supplemental Methods 2).

286

287 Hormone signalling

288 Missing components include abscisic acid regulation, perception, and transport, auxin
289 methylation-based degradation, brassinosteroid signalling (Figure 2d)⁴⁹ and degradation,
290 cytokinin degradation and transport, and degradation of gibberellins and jasmonic acid
291 and their transport genes.

292 For example, several components of the salicylic acid (*SAG101*, *EDS1*, *PAD4*,
293 Figure 2d)⁵⁰ and strigolactone signalling pathways were missing (Table S9), as is the
294 degradation component of the former hormone. To test this further, we investigated the
295 presence of canonical NPR domains (NPR1-like C superfamily, BTB/POZ NPR plant
296 domain or BTB/POZ superfamily, and an ANKYRIN domain) in our fern transcriptomes.
297 Of the 26 ferns we studied, 22 had at least one canonical NPR (Supplemental Methods
298 2, Figure SM7). This is consistent with previous evidence that the duplication of NPR1/3/4
299 happened sometime during angiosperm diversification, long after the split between
300 flowering plants and ferns⁵¹. The *SAG101/PAD4/EDS1* module, on the other hand,
301 appears to be a more recent invention of flowering plants, as it is mainly absent in non-
302 seed plants (Supplemental Methods 2, Figure SM8).

303 Further, we investigated perception and downstream signalling of jasmonic acid
304 (JA), focusing on the JA receptor COI1 and the JAZ transcriptional repressors. Both COI1
305 and JAZ candidates are encoded in fern transcriptomes. While *Arabidopsis thaliana* and
306 *Marchantia polymorpha* both have only one copy of COI1, ferns show several gene
307 duplication events, some of which are species-specific, and some of which appear more
308 ancient (Supplemental Methods 2, Figure SM9). The current evolutionary model of JA
309 perception is that the COI1 ligand switched from *dn-cis-OPDA* to JA-Ile in the ancestor of
310 vascular plants⁵². The radiation of COI1 in ferns, however, suggests functional
311 divergence in jasmonate perception, possibly complicating its evolutionary history. This
312 highlights ferns as a key lineage for further functional investigation to understand the
313 evolution of plant immunity.

314

315 Secondary metabolism

316 Phytochemical studies on ferns have revealed that they contain a wide range of
317 secondary metabolites, many of which are function herbivore defense and show bioactive
318 properties⁹. For secondary metabolite pathways associated with biotic interactions, we
319 observed that multiple genes known to act in the flavonoid biosynthesis pathway were
320 missing in all fern species analysed (Table S9), agreeing with previous datasets on the
321 evolution of red pigmentation in land plants^{53,54}. Yet, ferns are able to synthesise
322 flavonoids⁵⁵.

326 cortical sclerenchyma (Cs), endodermis (En), epidermis (Ep), metaxylem (MX), pericycle
327 (Pe) and phloem (Ph). Red arrows without labels indicate stained cell walls. b)
328 Percentage of lignin (1st column), H, G, S (2nd-4th) and H+G+S (5th) thioacidolysis lignin
329 units. c) The H, G, and S lignin unit biosynthesis pathway for angiosperms (red text) and
330 lycophytes (blue text). Intermediate metabolites are indicated by black text, while the grey
331 box contains enzymes involved in lignin polymerization. d) BLAST scores (x-axis) of
332 AtC3H, AtC4H, SmoF5H and AtF5H against the translated transcriptomes found in the
333 CoNekT database. Each point represents a protein. e) Comparative co-expression
334 network analysis of *PAL* genes from grape and fern *Dicranopteris curranii*. Nodes
335 represent genes, solid edges connect co-expressed genes, while dashed edges connect
336 orthologs. Coloured shapes represent different orthogroups, while the gene names are
337 based on the best BLAST hits to *Arabidopsis thaliana*. f) Phylogenetic tree of land
338 CYP450s of *P. irregularis* (genes starting with Pir), *S. palustris* (Spa), *A. obscurum* (Aob),
339 *Selaginella moellendorffii* (Smo), *Marchantia polymorpha* (Mp). The Arabidopsis (AT)
340 lignin-related C4H, C3'H and F5H and flavonoid-related F3H are included.

341

342

343 External stimuli response

344 Plants have evolved elaborate signalling and response pathways to cope with the
345 changing environment. For several of these pathways, we observed that orthologs of
346 phototropin-mediated receptors, all CO₂ sensing and signalling components, and many
347 gravity-sensing proteins were absent in our fern transcriptomes (Table S9). Genes known
348 to be essential for sensing and responding to temperature in flowering plants were
349 present in ferns, but acquired thermotolerance factors were missing. Other missing
350 proteins include those involved in several pathogenesis-related processes, such as
351 pattern- and effector-triggered immunity (16 out of 36 factors)(Figure 2d), WRKY33-
352 dependent immunity, pathogen polygalacturonase inhibitors, and basic chitinases. While
353 some components of symbiosis pathways are present in our fern transcriptomes (Table
354 S9), many factors are absent, such as mycorrhizal response genes and transporters.

355

356 Transcript control and modification

357 Several components controlling mRNA and protein levels are also absent, such as more
358 than half of the subgroups of MYBs and most REMs. Organellar RNA processing is
359 lacking plastidial and mitochondrial CFM-type splicing factors, a majority of mitochondrial
360 RBA splicing components (>20), C-to-U RNA editing (>50 factors), and mRNA

361 stabilisation and deadenylation factors. For protein homeostasis, we found only class C-
362 I and C-II small HSP holdase chaperones, while the ten other classes were absent (C-III
363 to ER), together with E3 ubiquitin ligases from groups IV and V.

364

365 Reproduction and organ development

366 Not surprisingly, our transcriptomes indicate that ferns differ from flowering plants in their
367 gene inventories related to reproduction. Ferns lack genes associated with anther
368 dehiscence (*PCS1*, *NST1/2*, *MYB26*), pollen aperture formation (*INP1/2*), pollen tube
369 growth (except *GEX3*), embryo axis formation (except *ATML1*), endosperm formation
370 (exception *GLAUCE*) and seed formation and dormancy (Table S9). On the other hand,
371 ferns contain nearly all male gametogenesis (e.g., *DUO1/3*, *DAZ*, *APD*) and exine
372 (*ROCK/TEX2*, *DEX1*, *NEF1*) formation factors, stamen (*TPD1*, *EMS1*, *JAG*) and tapetum
373 (*DYT1*, *TDF1*) regulators and most factors important for female gametophytes (*AMP1*,
374 *CYP78a*, *RKD*, *MAA3*) but lack genes essential for central cell formation (Table S9).
375 Surprisingly, while ferns are seedless, they contain most genes important for seed
376 maturation and globulins.

377 Interestingly, most flower formation photoperiodic and autonomous promotion
378 pathway genes are present in ferns and bryophytes. However, as expected, most genes
379 important for floral transition are missing (*FRIGIDA*, *FRL1/2*, *FES1*), except *FRI-C* effector
380 complex genes, floral meristem identity (*LMI2*, *AP1/3*, *PISTILLATA*, *SEPALLA*), and
381 morphogenesis (*BLR*, *ETT*). This suggests that the flower formation pathways have other
382 roles in ferns, possibly linked to photoperiodic response, developmental timing,
383 sporulation control, or other process.

384 For organ development, ferns lack several key genes essential for leaf adaxial and
385 abaxial polarity, guard cell formation, and stomatal density (Table S9). Their root
386 developmental programs are likely also different from flowering plants, as they lack the
387 entire MYB-bHLH-WD40 transcriptional regulatory module, and genes controlling
388 columella apical meristem (*WOX5*, *FEZ*, *SMB*, *BRN1/2*), and endodermis meristem
389 regulation and signalling (*SHR*, *SCR*, *KOIN*, *IRK*). More than half of Casparian strip
390 factors are missing, and nearly all vascular system formation factors (only 2 out of 14
391 transcription factors present).

392 Taken together, the analyses shown in Figure 2a-e provide further support for the
393 presence of unique growth, development, and survival strategies in ferns, and suggests
394 that additional research on them is worthwhile.

395

396 **Co-expression-based prediction of gene functions in the fern lineage**

397 The presence of >50% of fern-specific orthogroups (Figure 2b) indicated that ferns might
398 have evolved as-yet unknown gene functions on a massive scale. To investigate whether
399 the fern-specific genes can be annotated by sequence-similarity approaches, performed
400 an enrichment analysis of their biological functions. We observed enrichment for MapMan
401 bin containing uncharacterized genes ('No Mercator4 annotation', red line, Figure 2e),
402 and a depletion of bins related to known biological processes. This indicates that
403 sequence similarity approaches cannot infer the functions of most fern-specific genes.

404 Gene co-expression networks can reveal the functions of non-annotated genes
405 based on the guilt-by-association principle⁵⁶, where genes with similar expression profiles
406 tend to be involved in the same biological process. To predict the functions of fern-specific
407 genes without relying on sequence similarity to genes with known functions, we calculated
408 the functional enrichment of their direct neighbours in the co-expression networks.
409 Interestingly, fern-specific genes are significantly co-expressed (> 17 fern species) with
410 biological processes such as 'Cell wall organisation', 'Enzyme classification',
411 'Phytohormone action' and 'RNA biosynthesis' and moderately co-expressed (> 10 fern
412 species) with 'Cell division', 'Cytoskeleton organisation', 'Lipid metabolism', 'Multi-
413 process regulation', 'Protein biosynthesis', 'Protein modification', and others (Figure S10,
414 Figure 2e). This indicates that these genes are involved in most biological processes in
415 ferns, especially cell wall, development and new metabolic pathways.

416

417 **S lignin has evolved independently in the fern lineage**

418 Lignin is a complex phenolic polymer that forms essential structural materials in the
419 support tissues of vascular plants. Importantly, this polymer also confers hydrophobicity
420 to xylem vessels, allowing water transport from roots to leaves and enabling plants to
421 grow out on land. Lignin is primarily composed of three monolignols: *p*-coumaryl, coniferyl
422 and sinapyl alcohols, which are named *p*-hydroxyphenyl (H), guaiacyl (G) and syringyl

423 (S) units when incorporated into the polymer⁵⁷. Several studies, including one focused
424 on ferns⁵⁸, suggest a complex evolutionary history that may include independent
425 evolutionary paths for lignin synthesis, particularly the S units, among different plant
426 lineages⁵⁹. However, testing this was difficult without additional genomic information. Our
427 fern transcriptome datasets provided an opportunity to explore the evolution of lignin
428 across the entire fern family.

429 We first characterised the presence and sites of deposition of the different lignin
430 units in nine ferns from three orders: Equisetales, Cyatheaales and Polypodiales, using a
431 simple staining procedure. We stained cross-sections of stems and petioles with
432 Phloroglucinol-HCl, which reacts with coniferaldehyde residues of lignin to generate a red
433 condensation product^{60,61}. All nine ferns showed the presence of lignin. However staining
434 was not or poorly discernable in vessels of *Equisetum hyemale* and *Adiantum latifolium*
435 (Figure 3a, Figure S11). In many species, lignin was mostly found in the subcortical
436 sclerenchyma or outer layers of the metaxylem tissues, as expected (Figure S11).
437 Interestingly, Mäule staining on stem cross-sections indicated the presence of S-units in
438 *Pleocnemia irregularis* and *Stenochlaena palustris*⁶² (Figure 3a, Figure S11), indicating
439 that this subunit predominantly found in angiosperms⁶³, is also widespread in ferns.

440 Given these results, we further determined lignin content and structure within all
441 22 fern species, using the CASA method⁶⁴ and thioacidolysis followed by Gas
442 Chromatography-Mass Spectrometry (GC-MS)^{65,66}, respectively (Figure 3b). Overall, we
443 observed a large variability in total lignin content and the different units among the ferns.
444 Not surprisingly, *Equisetum hyemale* showed the lowest CASA lignin content (1 %), and
445 *Stenochlaena palustris* the highest (19 %)(Figure 3b). Most ferns contain lignin composed
446 of G units (130 - 13000 $\mu\text{mol/g}$ of CASA lignin), while H units are less abundant and, in
447 some cases, not detectable (0 - 57 $\mu\text{mol/g}$). Interestingly, we observed substantial
448 differences in the lignin content of multiple organs when analyzed. For example, H units
449 were detectable in petioles but not in rhizomes of *D. denticulata* and *A. opulentum* (Figure
450 3b, Table S11). We observed S units in high quantities in *P. irregularis* and *S. palustris*
451 (>4000 $\mu\text{mol/g}$), medium quantities in *A. obscurum* (86.9 $\mu\text{mol/g}$ CASA) and minute but
452 detectable quantities in *S. molesta*, *Cibotium barometz*, *Lindsaea ensifolia*, *Nephrolepis*
453 *biserrata* (<5.0 $\mu\text{mol/g}$ CASA).

454 Next, we set out to identify the biosynthetic pathways of lignin, focusing on S units.
455 To analyze the pathways and make the fern gene expression data easily accessible, we
456 uploaded the expression data for the 22 ferns to our CoNeKT database
457 (<https://conekt.sbs.ntu.edu.sg/>)⁶⁷, upgrading the database to comprise 39 species,
458 including angiosperms, lycophytes, bryophytes and algae. S unit synthesis evolved
459 independently in the lycophyte *Selaginella* and angiosperms⁶⁸, illustrated in Figure 3c.
460 Unlike angiosperms, which require *p*-coumarate 3-hydroxylase (C3'H) and ferulate 5-
461 hydroxylase (F5H) to make S units (Figure 3c, black arrows), *Selaginella* utilises a
462 multifunctional F5H that skips several steps of the pathway to make caffealdehyde and
463 caffeyl alcohol, which can then be utilised to make G and S lignin (Figure 3c, blue text)⁶⁹.
464 Blasting *AtC3H* and *AtC4H* (Cinnamate 4-hydroxylase) against all species proteomes in
465 the CoNekT database (<https://conekt.sbs.ntu.edu.sg/blast/>), showed identity scores of
466 >60% for ferns (Figure 3d, orange points), indicating that ferns likely contain C3'H and
467 C4H enzymes (Table S12). However, *AtF5H* and *SmF5H* showed only low sequence
468 identity to fern proteomes (~40%), which according previous studies indicates an absence
469 of known F5H enzymes in ferns⁷⁰.

470 To identify candidate fern F5H enzymes, we took advantage of the observation
471 that lignin biosynthetic genes tend to be tightly coexpressed and that these relationships
472 are conserved even across large evolutionary distances⁷¹. Indeed, comparing the co-
473 expression networks of *PAL* genes from fern *Dicranopteris* and angiosperm *Vitis vinifera*
474 (grape, Vitaceae) revealed many of the expected enzymes and a CYP98A3-like gene that
475 could likely represent *C3H* (Figure 3e, query gene *Dcu_g01768*, co-expression networks
476 of the lignin genes are in Supplemental Data 1). Furthermore, most of the fern lignin
477 biosynthetic genes are co-expressed with at least two other relevant enzymes (Figure
478 S12a), and the co-expression networks can suggest the unknown components (Figure
479 3f), including transcription factors and CYP450 enzymes (Figure 3f, Figure S12). To
480 suggest the identity of fern F5H enzymes, we first performed phylogenetic analysis of all
481 CYP450s of S lignin-producing ferns Pir, Spa, Aob, angiosperm *Arabidopsis*, lycophyte
482 *Selaginella* and included the outgroup bryophyte *Marchantia* that does not produce S
483 units (Figure 3g). We then indicated which CYPs are co-expressed with at least one lignin
484 enzyme. As expected, *C3H* genes are co-expressed with the other lignin biosynthetic

485 enzymes (co-expressed genes indicated with grey boxes, Figure 3g). However, we
486 observed several clades in the tree that likely emerged independently in ferns and
487 contained groups of co-expressed CYP450 enzymes (Figure 3g, indicated by red, green,
488 and blue arrows). These enzymes comprise prime candidates for the discovery of F5H
489 enzymes in ferns.

490

491 **Members of the Polypodiales contain a non-canonical cell wall sugar**

492 To further understand the evolution of fern cell walls, we carried out a monosaccharide
493 composition analysis using Gas Chromatography on the 22 ferns that were part of our
494 transcriptomic study (Table S13). The most abundant sugars were glucose (a building
495 block of cellulose, mixed-linkage glucans, xyloglucans), mannose (mannans) and xylose
496 (xylans)(Figure 4a). Less abundant sugars were rhamnose (pectic rhamnogalacturonan
497 I, arabinogalactan-protein), fucose (rhamnogalacturonan II, xyloglucan, arabinogalactan-
498 protein), arabinose (hemicellulose arabinoxylan, rhamnogalacturonan I and II,
499 arabinogalactan-protein) and galactose (rhamnogalacturonan I, hemicellulose
500 galactomannans, arabinogalactan-protein). The proportions of various sugars changed
501 among species and among different organs of the same species, which is in line with
502 previous observations⁷². For example, the *T. incisa* rhizome exhibited a higher proportion
503 of glucose than the petiole of the same species, and higher than rhizomes in of *Davaillia*
504 *denticulata* and *Ambloventanum opulentum* (Figure 4a). In addition to these sugars, we
505 also observed trace amounts of methylated rhamnose (3O-MeRhap), a known sugar
506 found in arabinogalactan proteins in ferns⁷³, in all species except for *Psilotum nudum*
507 and *Dicranopteris curranii* (Figure S13),

508 Interestingly, we detected an unknown peak from samples derived from three
509 species, *T. incisa*, *A. opulentum* and *D. proliferum* (Figure 4a, red bars). Because this
510 peak was not observed with our common standards during Gas Chromatography analysis
511 (data not shown), it likely represented a novel sugar. Since initial GC-MS analyses
512 suggested the sugar to be a methylated hexose (data not shown), we synthesised a panel
513 of methylated sugars (Figure S14). Out of the six methylated sugars, only 2-O-Methyl-D-
514 glucopyranose (2O-Me-Glcp) showed identical retention time and mass spectrum to the
515 unknown sugar (Figure 4b, Figure S15), indicating that these three species of ferns

520 **Figure 4. Polysaccharide analysis.** a) Percentage of total neutral sugars estimated by
521 GC-MS. The sugars are rhamnose, fucose, arabinose, xylose, mannose, galactose and
522 glucose. b) GC spectra of the unknown peak from c) The number of orthogroups involved
523 in cell wall biosynthesis in land plants. Columns represent species, while rows correspond
524 to a given gene family. The rows are further divided into different polysaccharide classes,
525 separated by horizontal lines. Red and blue numbers indicate that a given species
526 contains significantly more/less genes than others (adjusted p-value < 0.05). Darker
527 colors of boxes indicate more gene copies in each row. The red arrows indicate rows
528 which are particularly depleted in ferns. Ferns are indicated with bold names and thick
529 black lines. d) Relative epitope abundance for five fern species quantified by
530 Comprehensive Microarray Polymer Profiling (CoMMP). Rows indicate the different
531 species and organs, while columns represent the obtained signal from the different
532 antibodies. The colours of the cells correspond to the signal strength. e) Schematic
533 drawing of the major cell wall polysaccharides. Each colour-coded shape represents a
534 sugar or amino acid. The antibodies binding to a respective epitope are coloured with
535 blue letters, while the black bold letters indicate the biosynthetic enzymes.
536

537 **Ferns contain most but not all cell wall polysaccharides of angiosperms**

538 We next performed a large-scale comparative analysis of Carbohydrate-Active enZymes
539 (CAZymes) in land plants (Figure 4c)⁷⁵. The CAZyme database contains genes involved
540 in cell wall biosynthesis, allowing us to compare similarities and differences of ferns and
541 other land plants. To do this, we calculated with species contain significantly (adjusted p-
542 value <0.05) more (red numbers) or less (blue numbers) than the other species. Ferns
543 contain fewer xyloglucan-related gene families involved in remodelling (*BGAL10*)⁷⁶,
544 fucosylation (*FUT1*) and no genes involved in O-acetylation (*AXY4*)⁷⁷ (Figure 4c, red
545 arrows)⁷⁸. Although the number of *FUT1* homologs are fewer in ferns, the evolutionary
546 history of GT37 sequences was shown to be more complex when it comes to substrate
547 specificity⁷⁹. For xylans, ferns showed a near absence of genes involved in methylation
548 of glucuronic acid in glucuronoxylan (*IRX15*, *GXMT1-3*)⁷⁴ and xylan acetylation (*ESK1*)⁸⁰.
549 Homogalacturonan pectins showed fewer fern genes involved in xylogalacturonan
550 synthesis (*XGD1*)⁸¹ and galacturonan acetylation (*PMR5*)⁸². Finally, rhamnogalacturonan
551 I pectins showed fewer fern genes involved in remodelling (*TBG4*)⁸³. A similar analysis of
552 hydroxyproline-rich glycoproteins did not reveal significant absences of these proteins in
553 ferns (Figure S16). Overall, ferns tend to contain a lower number of genes per family and
554 do not have the *DUF579* (sugar methylesterification)⁸⁴ and *DUF231* (sugar acetylation)⁸⁵

555 gene families (Figure 4c).

556 To directly compare the polysaccharide inventories of angiosperms and ferns, we
557 performed a Comprehensive Microarray Polymer Profiling (CoMPP)⁸⁶, where we probed
558 102 cell wall extracts from nine ferns from six fern orders, including 36 organs at different
559 developmental stages, with 48 antibodies targeting different cell wall epitopes. The
560 epitopes recognised by the antibodies are given in Table S14.

561 Our analysis revealed the relative abundance of cell wall polysaccharides (Table
562 S15), which showed that different organs from the same species tend to have similar cell
563 wall composition (Figure S17a) and the polysaccharide profiles within species tend to be
564 more correlated than across species (Figure S17b).

565 We found that Eusporangiate ferns were generally richer in easily extracted
566 polymers than leptosporangiate ferns, with *Adiantum* as a notable outlier (Figure 4d).
567 Surprisingly, while mixed-linkage β -glucan (MLG) was only reported so far in *Equisetum*
568 ^{87,88}, we observed a clear MLG signal outside of Equisetidae in *O. pendulum*, *Angiopteris*
569 *sp1*, *A. capillus-veneris* and *S. palustris* (Figure 4d). For pectins, we observed a high
570 abundance of homogalacturonan at different grades of methyl esterification (antibodies
571 CCRCM38, LM18, LM19, LM20, JIM5, JIM7), but low signal from RG-I backbone (INRA-
572 RU1, INRA-RU2), galactosylated RG-I (LM16) and arabinan (LM13). For hemicelluloses,
573 we observed a strong signal for xyloglucan (CCRC-M87), both fucosylated (CCRC-M102,
574 CCRC-M1, CCRC-M39) and non-fucosylated (CCRC-M50), suggesting that xyloglucan
575 might be a quantitatively important hemicellulose, which contrasts with a previous study
576 reporting mainly mannan-rich cell walls ⁸⁹. We also observed signals from xylan (CCRC-
577 M154, LM11, CCRC-M159, LM10, LM23) and galactomannan (CCRC-M175) and various
578 mannan-containing polysaccharides (LM22, LM21), with galactomannan showing signal
579 only in *E. palustre* (CCRC-M170, -M167). The weakest signals were observed for
580 epitopes in arabinogalactan-proteins (AGPs) and extensins, as only a few antibodies
581 gave moderate signals (JIM8, MAC207, JIM13). Other antibodies showed weak signals
582 (AGPs: JIM16, LM2, LM14, extensins: JIM20, JIM11). Finally, feruloylated
583 polysaccharides that crosslink with arabinan and galactan residues of cell wall pectin via
584 ester bonds ⁹⁰ showed no signal (Figure 4d). Taken together, these results indicate that
585 fern and angiosperms share most of the polysaccharides and their biosynthetic enzymes,

586 but ferns might lack certain sugar modifications and AGP structures found in flowering
587 plants.

588

589 **Evolution of the cellulose synthase superfamily in Archaeplastida**

590 In addition to lignin, cellulose is one of the major load-bearing polymers. Angiosperms
591 contain primary and secondary cell walls enriched in cellulose, which are biosynthesised
592 by CESA1,3,6 and CESA4,7,8 in *Arabidopsis thaliana*⁹¹. The CESA complexes are
593 arranged in hexameric complexes called rosettes in angiosperms^{92,93}, or as linear
594 terminal complexes in bacteria^{94,95}. *Selaginella* is the latest diverging plant known to
595 possess both CesA hexameric complexes and CesAs of the type that forms linear
596 complexes in bacteria⁹⁶. The plant kingdom has also evolved cellulose synthase-like
597 (CSL) genes to produce other polysaccharides⁹⁷, such as mannans (CSLA)⁹⁸, glucan
598 chain of xyloglucan (CSLC)⁹⁹, cellulose in tip growing cells (CSLD)^{100,101} and mixed-
599 linkage glucans (CSLF)¹⁰². However, the evolution of the CESA superfamily is not well
600 understood in ferns.

601 A phylogenetic analysis of the *CESA* superfamily built from algal and land plant
602 protein sequences showed that ferns contain both linear (*CESA* linear) and hexameric
603 (*CESA1/3/10* and *CESA6*) *CESA* genes (Figure 5a, Figure S18). The *CSLA*, *CSLC* and
604 *CSLD* families were found in all land plants, including ferns (Figure 5a). The *CSLB* and
605 *CSLG* families are only found in seed plants¹⁰³, but ferns contain one clade of genes that
606 is likely ancestral to the two families (Figure 5a, green arrow). We also observed that the
607 angiosperm secondary cell wall enzymes *CESA4,7,8* and the fern *CESAs* do not form a
608 monophyletic group (Figure 5a, black arrows), indicating that ferns either lack *CESA4, 7,*
609 *8* or have evolved versions that no longer form clear clades with them. Conversely, ferns
610 form two distinctive groups with *Arabidopsis CESA6* and *CESA1,3,10* (Figure 5a),
611 suggesting that the ancestor of ferns and seed plants contained two *CESAs* that gave
612 rise to *CESA6-like* and *CESA1-like* clades. The phylogenetic tree revealed four
613 independent duplication events of the *CESAs* within ferns (Figure 5a, light blue arrows),
614 suggesting that ferns have likely evolved cell walls with properties distinctive from
615 flowering plants.

616

617 **Fern-specific evolution of secondary cellulose synthases**

618 To better understand the function of the duplicated CESAs ferns, we first analyzed the
619 gene tree of two CESA clades (Figure 5a, red and blue arrow). Both clades contain genes
620 from *Dicranopteris curranii* (red clade: *Dcu_g31359*, blue clade: *Dcu_g12277*, black
621 arrows), suggesting a duplication in the ancestor of Gleicheniales-II (Figure 5b). To
622 suggest the function of the red and blue clades, we first examined the expression profiles
623 of two representative genes (*Ceric.13G049300.1*) and (*Ceric.09G024100.1*) from
624 *Ceratopteris richardii* (Figure 5b, blue and red solid arrows, respectively). While
625 *Ceric.13G049300.1* showed the highest expression in vegetative fronds,
626 *Ceric.09G024100.1*'s expression was highest in leaf and shoot tips, suggesting different
627 biological processes for the two genes (Figure 5c).

628 We next compared the co-expression network of *Ceric.13G049300.1* to other ferns
629 using the CoNekT's Expression Context Conservation (ECC) panel
630 (<https://conekt.sbs.ntu.edu.sg/sequence/view/2353618>). The fern gene with the most
631 similar expression network was *Stenochlaena palustris Spa_g26805*, which happens to
632 be found in the same clade (Figure 5b, blue solid and double arrow). The networks of
633 *Ceric.13G049300.1* and *Spa_g26805* contain orthologs involved in lignocellulose
634 production, such as CESAs, 4CLs, OMT1s, CYP98A3 (lignin-related C3'H), CS11¹⁰⁴ and
635 laccases¹⁰⁵(Figure 5d). Thus, the genes from the blue clade are likely involved in
636 secondary well wall biosynthesis, indicating that ferns independently evolved this module.

637 Conversely, *Ceric.09G024100.1* and its most similar co-expression ortholog was
638 *Pyrrhosia piloselloides Ppi_g22229* (Figure 5b, green solid and double arrow) were co-
639 expressed with genes unrelated to lignocellulose production (e.g., genes similar to
640 monoterpenol-associated CYP76C2¹⁰⁶)(Figure 5d). This suggests that the second CESA
641 clade might be involved in another unknown biological process. Taken together, this
642 indicated that ferns have independently evolved a secondary cell wall module, and further
643 duplicated the CEASs to perform yet unknown functions.

654 represent genes, while solid and dashed edges connect co-expressed and orthologous
655 genes. Coloured shapes indicate the different orthogroups. e) Order tree summarising
656 the duplication events of cell wall-related genes. The tree is based on the gene tree of
657 CESAs and lignin-related genes. Coloured shapes represent the different gene classes.

658

659 **The evolution of lignocellulose-biosynthesizing genes in land plants**

660 To better understand how the genes involved in lignin and cellulose synthesis have
661 evolved in land plants, we mapped the timing of gene duplications onto the land plant
662 species tree (Figure 5e). Because both *CESA6-like* and linear *CESA* clades both contain
663 bryophytes (Figure 5a), we propose that the ancestor of land plants contained a *CESA6-*
664 *like* and a *CESA* of linear-type. The ancestor of ferns and seed plants evolved *CESA1/3-*
665 *like* genes, that further expanded in seed plants into *CESA1* and 3 and secondary
666 *CESAs4,7,8* (Figure 5a). Within ferns, we observed duplications of *CESA1/3-like* in the
667 ancestor of Gleicheniales-II (Figure 5a, duplication 3), *CESA6-like* in Hymenophyllales
668 (duplication 4), and two duplications of the fern-specific CESAs (duplications 1,2) in
669 Gleicheniales-II. We also observed a complete gene set of lignin biosynthetic genes in
670 early-diverging land plants, and evidence of duplication of *4CL* in the ancestor of ferns
671 and seed plants and the ancestor of Hymenophyllales (Figure S19).

672 Taken together, the ferns show a prolific duplication of *CESA* genes deeply within
673 the fern lineage (Figure 5e), further suggesting that ferns have evolved cell walls with yet
674 unknown features.

675

676 **Discussion**

677 Despite ferns' critical evolutionary position as the sister group, no large-scale studies that
678 investigated their phylogeny, biological pathways and cell walls had been performed. To
679 remedy this, we generated gene expression atlases for 22 ferns and covered ten out of
680 12 fern orders (Figure 1), allowing us to generate a high-quality species tree that resolved
681 the long-standing relationship between Gleicheniales and Hymenophyllales^{12,107,108}. The
682 tree is supported by outgroups comprising lycophytes, horsetails or seed plants, but not
683 bryophytes. We speculate that the greater phylogenetic distance between bryophytes and
684 the other lineages, combined with reduced single-copy gene dataset obtained from
685 OrthoFinder (Supplemental Methods 1), and degenerated phylogenetic signals in amino

686 acid sequences contributed to the discordance between bryophyte-based and the other
687 outgroups.

688 The species tree of ferns allowed us to estimate the time of speciation and whole
689 genome duplication events. The WGD analysis revealed that WGD events likely
690 contributed to species diversity (Figure S3), but we observed no correlation between the
691 number of WGDs and genome size. This suggests that alternative evolutionary sources
692 contribute to the exceptional genome size of ferns, and that recent transposon activities
693 and ploidy variation might play a bigger role.

694 The stem age of early diverging ferns Equisetidae (consisting of Equisetales), with
695 95% HPD time estimates are in line with the estimate from ¹⁰⁹ and the oldest unequivocal
696 euphyllophyte fossils ¹. Polypodiidae was originated in the time range between Lower
697 Carboniferous ($323.2 \pm 0.4 - 358.9 \pm 0.4$ mya) and Middle Devonian ($382.7 \pm 1.6 - 393.3$
698 ± 1.2 mya), with 95% HPD time estimate as 345.09 - 389.61 mya and posterior mean
699 369.01 mya, which might have first survived the Hangenberg and Kellwasser extinction
700 events before its substantial diversification. The early diverging leptosporangiate fern
701 order Osmundales originated amid Upper to Lower Carboniferous ($298.9 \pm 0.15 - 358.9$
702 ± 0.4 mya), consistent with ¹³. The aquatic Salviniales, the only extant ferns with
703 heterospory, was originated between Upper Triassic ($201.4 \pm 0.2 - 237$ mya) and Lower
704 Permian ($273.01 \pm 0.14 - 298.9 \pm 0.15$ mya), with 95% HPD time estimate as 211.15 -
705 273.19 mya and posterior mean 241.70 mya, which might correlate with the P-T event
706 after which a vast majority of aquatic environment became empty and the innovative
707 microspores might facilitate their spread and survival. The two most species-diverse
708 suborders, Polypodiineae (i.e., eupolypods I) and Aspleniineae (i.e., eupolypods II) of the
709 Polypodiales were originated amid the Cretaceous (66.0 - 145.0 mya), a relatively warm
710 and ice-free period, with 95% HPD time estimate as 89.75 -144.92 mya and posterior
711 mean 115.93 mya, coincident with the burst of angiosperms in the mid-Cretaceous as
712 highlighted by Darwin ¹¹⁰ and the decline of gymnosperms ¹¹¹.

713 Our gene inventory analysis shows massive gains of genes in the fern lineage
714 (Figure 2a), resulting in ~50% being fern-specific (Figure 2b). Expression analysis
715 revealed that the fern-specific genes tend to be organ-specific, suggesting their role in
716 fern-specific adaptations. Conversely, older genes are ubiquitously expressed (Figure

717 2c), which aligns with our previous observation that these genes tend to have basal,
718 essential functions (e.g., photosynthesis, protein synthesis, DNA duplication)⁴⁷. Many of
719 the genes involved in angiosperms' hormonal and developmental pathways were missing
720 (Table S9), showing that ferns have organised these pathways differently.

721 Signalling and biosynthetic pathways may significantly vary within land plants, and
722 these pathways tend to expand to support increased anatomical and lifestyle complexity
723 ¹¹². Thus, the arguably simpler fern hormonal pathway genes might suggest that these
724 pathways can function in ferns without their angiosperm counterparts. Alternatively, ferns
725 might have evolved equally complex but alternative signalling pathway components that
726 show no sequence similarity to known angiosperm genes. This idea is exemplified by our
727 analysis of flavonoid biosynthesis genes. The lack of these enzymes in ferns—but the
728 presence of flavonoids—indicates that the 'canonical' flavonoid pathway is an angiosperm-
729 specific invention and suggests that ferns have either convergently evolved other
730 enzymes with similar functions or use a different pathway to synthesise these
731 compounds. As fern-specific genes are co-expressed with genes involved in
732 development, reproduction and various signalling pathways (Figure 2e), ferns likely have
733 independently expanded these pathways.

734 The observed high amounts of lignin S units in *P. irregularis* and *S. palustris*
735 (Figure 3ab), and the absence of angiosperm- or lycophyte-specific *F5H* enzymes
736 suggest that the S unit has independently evolved at least four times in the plant lineage:
737 angiosperms, lycophytes, gymnosperms and now ferns ¹¹³. The re-emergence of S lignin
738 in distantly related plant lineages implies that it may have an essential role in plants'
739 environmental adaptation, such as improved mechanical properties or herbivore
740 resistance ¹¹³. While lycophytes have evolved a C3'H-independent pathway by inventing
741 a dual meta-hydroxylase *SmF5H* (Figure 3c, blue pathway)⁶⁹, we observed the presence
742 of *C3'H* genes in ferns (Figure 3d), suggesting that ferns have independently evolved a
743 *F5H* enzyme, and likely follow the biosynthetic route of angiosperms. By combining
744 phylogenetic and co-expression analysis, we propose that the red clade shown in Figure
745 3h, which contains the highest density of *CYP450s* co-expressed with the lignin
746 biosynthetic genes, comprises the fern *F5H* enzymes. The biosynthetic activity of these
747 genes could be tested by in vitro studies, as done for Selaginella *SmF5H* ⁶⁹.

748 Our comparative analysis of cell wall-related genes indicated that ferns and
749 angiosperms contain similar gene sets but that ferns have smaller acetyl and methyl-
750 transferase gene families (Figure 4c). Cell wall composition varies considerably between
751 fern species, corroborating findings in earlier glycan array surveys of ferns ¹¹⁴.
752 Surprisingly, we observed a clear mixed-linkage β -glucan signal from *O. pendulum*,
753 *Angiopteris sp1*, *A. capillus-veneris* and *S. palustris* (Figure 4d), demonstrating that this
754 unusual polymer is found outside of fern Equisetum ^{87,88}. While the AGP epitopes showed
755 weak signals, ferns contain AGPs with special features, such as 3-O-methylrhamnose,
756 that are not known in angiosperms ^{73,115}.

757 Surprisingly, we observed a wide-spread occurrence of mixed-linkage glucans
758 outside of Equisetidae (Figure 4d), and we propose two candidate enzyme families that
759 could produce this hemicellulose. First, bryophytes, ferns and Selaginella both contain
760 MLGs ⁹⁶ and linear CesAs (Figure 5a). In the moss *Physcomitrium*, linear CesAs produce
761 arabinoglucan ¹¹⁶, and the authors point out that these CESAs are related to an
762 ascomycete MLG synthase and thus represent an early system for MLG-synthesis. A
763 second candidate could be the fern CSLB/G-related clade, as the functions of these
764 genes are currently unknown in ferns and angiosperms. The biosynthetic activity of these
765 enzymes could be tested in vivo, as done for the barley MLG synthase ¹¹⁷.

766 Separate sets of CesAs for primary and secondary cell wall synthesis are a shared
767 feature of spermatophytes ⁹¹. Whereas tracheids probably evolved once uniting all
768 tracheophytes ¹¹⁸, vessels have evolved in angiosperms and independently in
769 *Selaginella*, the *Gnetales*, *Equisetum* and other ferns ¹¹⁹. Surprisingly, *Selaginella* does
770 not have two sets of CesAs, and we did not observe CesA related to angiosperm
771 secondary cell walls in ferns (Figure 5a), suggesting that their vascular elements result
772 from convergent evolution ⁹⁶. Convergent evolution is supported by a fern-specific CesA
773 clade containing genes involved in lignocellulose biosynthesis (Figure 5). We observed
774 several duplications of the cell wall-related genes within ferns (Figure 5e), which aligns
775 with similar observations in angiosperms and bryophytes ^{120,121}. Combined with the
776 presence of a non-canonical sugar 2-O-Methyl-D-glucopyranose observed in
777 Polypodiales (Figure 4a), our data suggest that cell walls underwent independent
778 innovations within ferns. While it is unclear whether 2-O-Methyl-D-glucopyranose is

779 biosynthesized by ferns or bacterial or fungal organisms found in the environment, their
780 significant presence in the fern cell walls indicates that they might have a role in fern
781 biology.

782 We anticipate that the availability of the comprehensive coding sequence and
783 transcriptomic data from ferns - and their availability as a user-friendly CoNekT database
784 (<https://conekt.sbs.ntu.edu.sg/>)-will be mined to lead to vital insights into the evolution of
785 plant genes and gene families. Implementing fern data into the existing comparative
786 genomic framework will enhance our understanding of the plant tree of life.

787

788 **Methods**

789 **Sampling of ferns**

790 22 ferns from 22 families were sampled across Singapore (Table S1). Fern organs were
791 sampled as three biological replicates, where organs were selected to capture the highest
792 variance in the developmental stages and morphological characteristics (Figure S1).
793 Samples were placed into 15 ml falcon tubes and kept in liquid nitrogen and subsequently
794 at -80°C to prevent degradation of RNA.

795

796 **RNA isolation and sequencing**

797 After collection, each sample was ground in liquid nitrogen to a fine powder. RNA was
798 extracted from 100 mg of plant material using Spectrum™ Total Plant RNA Kit (Sigma)
799 Protocol A following the manufacturer's instructions. Quality control of all extracted RNA
800 was carried out by Novogene (Singapore). Each sample was evaluated for its quantity,
801 integrity and purity using agarose electrophoresis and Nanodrop. Library construction
802 was performed by Novogene, and mRNA was enriched from total RNA with oligo(-dT)
803 magnetic beads. The library was then quantified with Qubit and real-time PCR and
804 sequenced using Illumina NovaSeq 6000, with paired-end sequencing of 150 base pairs
805 (bp) per read and a sequencing depth of approximately 60-70 million reads.

806

807 **Transcriptome assembly**

808 Low-quality RNA-seq reads were removed, and the remaining reads were trimmed with
809 Fastp (v0.23.2)¹²². Reads were assembled via a curated transcriptome assembly pipeline

810 (Figure S2). Reads were assembled in three biological replicates for each organ, with all
811 organs concatenated and filtered. Each organ consisting of three reads was assembled
812 using SOAPdenovo-Trans (v1.03)³⁷ with 10 single *K-mer* (21-39) and Trinity (v2.8.5)³⁶
813 with 25 single *K-mer*. All reads were concatenated into a Trinity-SOAPdenovo assembly
814 and filtered through the Evidential Gene Pipeline
815 (<http://arthropods.eugenescience.org/EvidentialGene/>) using trformat.pl and tr2aacds.pl for
816 removal of any redundant coding sequences. The filtered transcriptome was evaluated
817 using BUSCO (v5.4.3)¹²³ and embryophyta as the dataset. All organ assemblies within
818 the sampled fern species were concatenated and filtered again through the
819 EvidentialGene Pipeline, with the output transcriptome being filtered via transcripts per
820 million (TPM) reads using kallisto (v0.50.1)¹²⁴ against each RNA-seq. TPM scores were
821 averaged per organ, and coding sequences with scores < 1 were removed from the
822 assembly. Assembly was then filtered using GC% content, with redundancy coding
823 sequences with less than 40% and more than 60% removed. All transcriptomes were
824 blasted against the NCBI database, and sequences with a percentage identity of more
825 than 70% and an e-value less than e^{-10} were removed from the final assembly. The quality
826 of assembly was determined by BUSCO using the embryophyta dataset (Table S2).

827

828 **Construction of K_S -based age distributions**

829 K_S -age distributions for all paralogous genes (paranome) of genomes and transcriptomes
830 were constructed by ksrates (v1.1.1)¹²⁵. In brief, the ksrates pipeline entails firstly
831 translating the coding nucleotide sequences into peptide sequences assuming standard
832 genetic code, filtering out sequences whose sequence length is not divisible by 3,
833 containing invalid codons or in-frame stop codon, after which an all-versus-all BLASTP
834 was implemented with *E*-value set as 1×10^{-10} in BLASTP (v2.11.0+)¹²⁶ and the resultant
835 subject-query hit table was fed into MCL (v14-137)¹²⁷ with clustering inflation factor set
836 as 3.0 to delineate paralogous gene families while filtering out gene families whose size
837 is larger than 200, secondly calling the aligner MUSCLE (v3.8.1551)¹²⁸ under default
838 parameter to obtain a multiple sequence alignment (MSA) at the protein level for each
839 paralogous gene family while filtering out sequence pairs whose gap-stripped alignment
840 length was shorter than 100, which was then back-translated into a codon alignment and

841 subsequently fed into the CODEML function within PAML (v4.9j)¹²⁹ to acquire the
842 maximum likelihood estimate (MLE) of K_S values under non-pairwise mode using the
843 default control file defined by wgd (v1.1.1)¹³⁰ and then calling FastTree (v2.1.11)¹³¹ upon
844 the peptide MSA under default parameter to attain a midpoint-rooted phylogenetic tree of
845 each paralogous gene family for retrieving the weight of each paralogous gene pair with
846 or without outliers, and eventually building the K_S -age distribution with de-redundancy
847 achieved by node-weighted method after excluding outliers. The collinear gene pairs
848 (anchor pairs) were identified by i-ADHoRe (v3.0.01)¹³² under the default control file
849 defined by wgd, and the weight values for anchor pairs whose corresponding K_S values
850 were between 0.05 and 20 were recalculated and reassigned while the weight of
851 remaining pairs was set as zero. For orthologous K_S -age distributions, the process of
852 MCL clustering was superseded as reciprocal best hits (RBH) searching to identify
853 orthologous gene pairs while the weighting process was revoked on that only one-versus-
854 one orthologues were inferred. CD-HIT (v4.8.1)¹³³ was applied for the de-redundancy of
855 transcriptome assemblies with the clustering threshold set as 0.99 before K_S -age
856 analysis.

857

858 **Correction of differences of synonymous substitution rates**

859 Synonymous substitution rates were corrected in ksrates. The principle leans on a
860 number of trios of species, including a focal species, a sister species and an outgroup
861 species. The disparate synonymous substitution rate between focal species and sister
862 species since the divergence is represented indeed by the branch-specific contribution of
863 accumulated synonymous substitutions per synonymous site in respective branches. The
864 mode of orthologous K_S -age distributions inferred from the kernel density estimate (KDE)
865 using Gaussian kernels within the python package scipy was designated as the proxy of
866 the peak K_S value of each orthologous K_S -age distribution. 200 iterations of bootstrap with
867 replacements were implemented for each orthologous K_S -age distribution, and the mean
868 along with standard deviations (STD) of mode across the replicates was determined as
869 the final peak K_S value representing divergence distance and its associated STD. The
870 original accumulated synonymous substitutions per synonymous site of focal species-
871 sister species pair consisting of the branch-specific contribution of both species since

872 diversification was transformed into two times the branch-specific contribution of focal
873 species with the prop of outgroup species to resemble the timescale of focal species. The
874 mean of rescaled peak K_S values of focal species-sister species pair against various
875 outgroup species was taken as consensus-adjusted peak K_S value. The maximum
876 number of trios was set as 20. The species tree inferred by ASTRAL-Pro2 using seed
877 plants as outgroup species were adopted in ksrates.

878

879 **Construction of orthologous families and single-copy gene trees**

880 Orthofinder (v2.5.4)¹³⁴ was performed upon the protein sequences of 107 ferns and
881 outgroup species with an inflation factor set as 3 to delineate the orthologous families. No
882 single-copy gene families were identified by Orthofinder, probably because of the
883 universal and unique gene duplication and loss scenario across species and gene
884 isoforms¹³⁵. To recover reliable and adequate single-copy gene families, we constructed
885 mostly single-copy gene families⁵¹, wherein most species were in single-copy while the
886 remaining species had no more than four copies which were assumed to be transcript
887 variants of the same gene, by retaining the longest copy, if applied, of each species. We
888 referred to the mostly single-copy gene families as single-copy gene families thereafter.
889 In total, 140, 112, 107 and 57 single-copy gene families were constructed from dataset
890 107 ferns, 107 ferns plus seed plants, 107 ferns plus seed plants and lycophytes, 107
891 ferns plus seed plants, lycophytes and bryophytes, respectively. MAFFT (v7.475)¹³⁶ was
892 performed to obtain a peptide multiple sequence alignment (MSA) for each single-copy
893 gene family with the parameter “-auto”. Trimal (v1.4.1)¹³⁷ was then performed to trim the
894 MSA and back-translate it into a codon-level nucleotide MSA with parameter “-
895 automated1”. IQ-TREE (v1.6.12)¹³⁸ was implemented on each codon-level nucleotide
896 MSA wherein ModelFinder¹³⁹ was called to find the best-fit codon substitution model in
897 terms of Bayesian Information Criterion (BIC) upon which a maximum likelihood (ML)
898 gene tree was inferred and assigned with bootstrap support values from 1000 ultrafast¹⁴⁰
899 bootstrap replicates with parameter “-bnni” to further optimize each bootstrap iteration
900 through a hill-climbing nearest neighbor interchange (NNI) search based directly on the
901 corresponding bootstrap alignment to avoid severe model violations. The same process
902 was further applied to the 108 ferns dataset including *Marsilea vestita*, in which a total of

903 136, 108, 103 and 55 single-copy gene families were reconstructed from datasets varied
904 in outgroups with both nucleotide and peptide molecules.

905

906 **Species tree inference**

907 Three methods, ASTRAL-Pro2 ⁴¹, STAG ⁴², and a concatenated-based method were
908 implemented to infer the species tree. The acquired individual ML single-copy gene trees
909 and gene name-species name map files were imported into ASTRAL-Pro2 and STAG
910 under default parameters to estimate a consensus species tree with support values for
911 each bipartition denoting local posterior probabilities (localPP) and the proportion of
912 individual estimates of the species tree that contain that bipartition, respectively. For
913 the concatenated-based method, the individual codon-level nucleotide MSA of single-
914 copy gene families were concatenated and then fed into IQ-TREE to infer a ML super-
915 gene tree as above. *Dicranopteris curranii*, *Dicranopteris pedata*, *Diplopterygium*
916 *laevissimum*, *Diplopterygium glaucum* and *Sticherus truncatus* in the Gleicheniaceae
917 were named as Gleicheniales-II clade while *Cheiropleuria integrifolia*, *Dipteris conjugata*
918 and *Dipteris lobbiana* in the Dipteridaceae were named as Gleicheniales-I clade p. In
919 total, (140), (112), (107) and (57) single-copy gene families were constructed from the
920 (107 ferns dataset), (107 ferns plus seed plants), (107 ferns plus seed plants and
921 Lycopods), (107 ferns plus seed plants, Lycopods and Bryophytes), respectively. The 108
922 ferns dataset including *Marsilea vestita*, were with 136, 108, 103 and 55 single-copy gene
923 families, respectively.

924

925 **Estimation of absolute divergence time**

926 Mcmctree (v4.9j)¹²⁹ was implemented upon the concatenated peptide MSA of single-copy
927 gene families of 108 ferns dataset with Equisetales as outgroup to infer the absolute
928 divergence time for each bipartition. The independent rates model, which assumes a log-
929 normal distribution of evolutionary rates across branches, was selected and 18 fossil
930 calibrations of soft constraint from ¹⁴¹ were adopted to refine the divergence time of
931 internal nodes, as summarised in Table S2. Fossils calibrating clades within
932 Gleicheniaceae or Hymenophyllaceae were avoided for their indefinite phylogenetic

933 location. LG amino acid substitution matrix was selected and a gamma model of rate
934 variation was assumed with alpha as 0.5 and 5 categories in discrete gamma. Parameters
935 controlling the birth-death process were set as 1, 1, 0.1 to generate uniform age priors on
936 nodes that didn't have a fossil calibration. Gamma priors for the transition/transversion
937 rate ratio and shape parameters for variable rates among sites were set as 6 2 and 1 1.
938 A Dirichlet-gamma prior was set upon the mean rate across loci and the variance in
939 logarithm as 2 20 1 and 1 10 1. The first 2000 iterations were discarded as burn-in and
940 then 20,000,000 iterations were performed with sampling per 1000 iterations. The
941 effective sample size (ESS) of all parameters was larger than 200, suggesting adequate
942 sampling and convergence.

943

944 **Phylogenomic analysis of gene tree - species tree reconciliation**

945 To estimate the retention rate and interrogate hypothetical WGDs over competing
946 scenarios in different clades, we implemented 4 categories of statistical gene tree -
947 species tree reconciliation analysis using Whale (v.2.0.3)¹⁴², as shown in Figure SM9-12.
948 Firstly, orthogroups of each category of species were inferred by Orthofinder (v2.5.4)¹³⁴
949 with an inflation factor as 3. Gene families were filtered to assure at least one gene from
950 each descendant present at the root and to avoid large gene family size(which contains
951 noise and causes computational downshift) via "orthofilter.py"
952 (<https://github.com/arzwa/Whale.jl>)1000 gene families were randomly selected as
953 subsequent inputs. PRANK (v.150803)¹⁴³ was utilised to obtain a MSA for each gene
954 family and MrBayes (v.3.2.6)¹⁴⁴ was then applied to infer the posterior distributions of
955 gene trees under the LG + GAMMA model, with iterations set as 110,000 and sample
956 frequency as 10 to get in total 11,000 posterior samples. ALEobserve¹⁴⁵ was
957 subsequently performed on the tree samples to construct the conditional clade distribution
958 with a burn-in of 1000. Two gene family evolution models, the relaxed branch-specific
959 model and the critical branch-specific model, were applied as in a previous study⁴⁵ to
960 estimate the retention rates of hypothetical WGDs for each category, as shown in Figure
961 SM9-12. Hypothetical WGDs with retention rates higher than 0.05 were regarded as
962 supported WGDs, considering the incompleteness of transcriptome assemblies and the
963 stochasticity of sampled gene families.

964

965 **Absolute dating of WGDs**

966 Phylogenetic dating of AZOL and CERA WGD proceeded as follows. Firstly, an
967 orthogroup comprising orthologues from 8 other species and anchor pair which was
968 assumed to be retained from the corresponding WGD was constructed per anchor pair
969 by searching the reciprocal best hits (RBH) between the anchor pair and the
970 transcriptomes or genomes of other species by Diamond (v2.0.5.143)¹⁴⁶ under default
971 parameters (Figure SM14). K_s range 0.36 - 2.00 was confined for the age of anchor pairs
972 to be adopted in terms of densest aggregation of duplicates and avoidance towards
973 saturation for AZOL WGD and K_s range 0.41 - 2.0 was bounded for CERA WGD.
974 Secondly, the peptide sequences of each individual orthogroup were aligned by MAFFT
975 (v7.475)¹³⁶ under default parameters and then concatenated as a single peptide MSA.
976 The numbers of concatenated orthogroups were 45 and 14 for AZOL and CERA WGD,
977 respectively. The adopted fossil calibrations followed Table S2 at corresponding
978 phylogenetic locations while the boundaries of root for AZOL WGD were set as minimum
979 bound 168 mya based on the minimum bound of fossil calibration “Stem Lygodiaceae”
980 and safe maximum bound 345 mya as the fossil calibration “Stem Osmundaceae”, as
981 shown in (Figure SM14). Mcmctree (v4.9j)¹²⁹ was implemented for the Bayesian
982 molecular dating for each WGD with the parameters same as above. The ESS of all
983 parameters was larger than 200, indicating adequate sampling and convergence. The
984 posterior distribution of time estimate for the node joining the anchor pair was retrieved
985 and the 95% HPD, posterior mean, median and mode were adopted to characterise the
986 age of WGD, as shown in (Figure SM13).

987

988 **Identifying organ-specific genes**

989 Organ-specific genes were isolated from each transcriptome via specificity measure
990 (SPM) values⁴⁷. For each gene, we calculated the average TPM values in each organ.
991 Following that, the SPM value of a gene in an organ was calculated by dividing the
992 average TPM in the organ by the sum of the average TPM values of all organs. The SPM
993 value ranges from 0 (gene not expressed in organ) to 1 (gene fully organ-specific). To
994 identify organ-specific genes for each organ, we first identified an SPM value threshold

995 above which the top 5-11% of SPM values were found (Figure S5). These top values
996 varied across the species sampled, depending on the number of organ-specific genes
997 identified. If the SPM value of a gene in an organ was equal or greater than the threshold
998 value, the gene was identified as organ-specific within said organ. Organ-specific genes
999 were then plotted in a heat map to show their distributions (Figure S6).

1000

1001 **Functional annotation of genes**

1002 Assembled sequences of 22 fern species, including four ferns with genomes available
1003 online (*A. spinulosa*, *A. filiculoides*, *C. richardii* and *S. cucullata*) were annotated using
1004 the online tool Mercator4 v.2.0¹⁴⁷. We visualised the Mercator4 annotation using a
1005 heatmap, showing the distribution of Mapman Bins across sampled fern species (Figure
1006 S9).

1007

1008 **Assignment of orthogroups to phylostrata**

1009 Using the coding sequences of the transcriptomes, we constructed orthologous gene
1010 groups (Orthogroups) with Orthofinder (v.2.5.4)¹³⁴. Respective outputs for orthogroups
1011 were used for further analysis. By utilising the theoretical evolutionary line produced by
1012 the phylogenomic analysis of gene trees, phylostratic nodes were assigned to
1013 orthogroups based on plant lineages. This analysis spanned a total of 47 species across
1014 the plant kingdom, and assigned nodes ranging from node 1 (most ancient, ancestor of
1015 Archaeplastida) to node 13 (ancestor of Polypodiales). Nodes were assigned based on
1016 the fern species tree (Figure 1c), as well as known phylogenetic analyses of early plants
1017¹⁴⁸. The nodes are: node 1 (ancestor of Archaeplastida), node 2 (ancestor of green
1018 plants), node 3 (ancestor of land plants), node 4 (ancestor of vascular plants), node 5
1019 (ancestor of ferns and seed plants), and node 6-13 (various fern orders). Specifically,
1020 Equisetales is designated as node 6, Psilotales and Ophioglossales as node 7,
1021 Marattiales as node 8, Hymenophyllales and Gleicheniales-II as node 9, Schizaeales as
1022 node 10, Salviniales as node 11, Cyatheaales as node 12, and Polypodiales as node 13.
1023 Species-specific gene families were characterised by gene families consisting of only one
1024 species, and hence, not assigned to nodes. In cases where nodes encompass multiple
1025 species, such as node 4, orthogroups containing only one node assignment (e.g., those

1026 with genes solely from Ophioglossales and Psilotales) were not designated to specific
1027 nodes.

1028

1029 **Orthogroup gain loss analysis**

1030 Gain and loss of orthogroups were determined by the presence of an oldest clade
1031 member in a particular node. Potential contamination by non-fern sequences due to the
1032 nature of transcriptome assembly was filtered out at this stage by checking for the
1033 presence of at least half of the expected clades in each node. For basal nodes (nodes 1
1034 to 4), the clades used were 'Glaucophytes', 'Chlorophytes', 'Bryophytes', 'Lycophytes',
1035 'Tracheophyta' and 'Spermatophyta'). Nodes were defined as lost based on the clade that
1036 they last appeared in.

1037

1038 **Identification of orthogroup expression profiles**

1039 Analysis of the expression profiles at phylostrata level was performed as in ⁴⁷, by
1040 classifying orthogroups into 'organ-specific', 'ubiquitous' or 'not conserved'. Organ-
1041 specific orthogroups are orthogroups containing organ-specific genes and were
1042 subclassified according to the organ (leaf-, meristem-, crozier-, root meristem-, male-,
1043 spore-, rhizome-, root-specific). Orthogroups that are expressed in different organs for
1044 each species - that is, that do not show an 'organ-specific' expression profile in different
1045 species - were labelled as ubiquitous. Orthogroups that had different organ-specific
1046 expression profiles in different species (orthogroups containing root-specific genes for
1047 *Alsophila latebrosa* and leaf-specific genes for *Equisetum hyemale*) were labelled as not
1048 conserved. Only orthogroups that fulfilled the following criteria were identified as organ-
1049 specific: (1) Contained at least two species with transcriptome data within each
1050 orthogroup. (2) >50% of the genes within the orthogroup supported the expression profile
1051 and (3) ≥50% of the species present in the node supported the expression profile.

1052

1053 **Structural analysis of fern-specific orthogroups**

1054 25 orthogroups containing at least 10 fern species with protein sequence representatives
1055 from sequenced genomes were used to check for sequence similarity by NCBI BLASTp
1056 restricted to Viridiplantae (E-value < 1e-10, Query cover > 50%), prediction of structure
1057 (alphafold 3 -server)¹⁴⁹, structure similarity search using DALI (all PDB, Z score > 8, lali >
1058 0.5 of residues in the protein)¹⁵⁰ and foldseek¹⁵¹. The cif outputs from alphafold 3 were
1059 converted to pdb format for input to DALI and foldseek using UCSF ChimeraX version:
1060 1.8. The sequenced representatives were selected based on the highest similarity to the
1061 consensus sequence, which was derived from the multiple sequence alignment
1062 generated using Seaview v5.0.5 (-align -align_algo 1 -output_format clustal -o) using the
1063 muscle algorithm¹⁵².

1064

1065 **Constructing co-expression networks and addition of the ferns to the CoNekT** 1066 **database**

1067 Co-expression networks were calculated using the CoNekT framework⁶⁷, and were also
1068 used to update the existing database, available at (<https://conekt.sbs.ntu.edu.sg/>).

1069

1070 **CASA lignin quantification**

1071 Methods used for solvent extraction and determination of lignin content by CASA lignin
1072 method closely followed the protocol outlined in⁶⁴. Species organs that were sampled
1073 were ground, with solvent extraction in 80% ethanol for woody samples. For non-woody
1074 samples, extraction with water using sonication was done first to remove proteins and
1075 other water-soluble components. A cysteine stock solution (0.1 g/mL) in 72% sulfuric acid
1076 (SA) was prepared by dissolving 10 g L-cysteine in 100 ml SA. 5-10 mg of the solvent
1077 extract was placed in a glass vial, where 1.0 mL of prepared stock solution was added,
1078 sealed with a Telfon-lined screw cap and stirred at 24 °C (room temperature) via a
1079 magnetic stir bar (400 rpm) for 60 mins until the biomass was completely dissolved. The
1080 dissolving temperature was decreased to 24 °C to identify a milder condition, allowing
1081 convenient operation and minimising interference from carbohydrates. The solution was
1082 diluted with deionized water to a volume of 50 or 100mL in a volumetric flask, depending
1083 on the lignin content and biomass weight used. Absorbance of the diluted solution was

1084 measured at 283 nm (A_{283}) in a 1 cm quartz cell using a UV spectrophotometer against a
1085 blank solution (1 mL stock solution diluted to corresponding volume).

1086

1087 **Thioacidolysis**

1088 This method is adapted from ⁶⁶. Briefly, 10 mg of alcohol-insoluble cell wall residues were
1089 incubated in 3 mL of dioxane with ethanethiol (10%), BF₃ etherate (2.5%) containing
1090 0.1% of heneicosane C₂₁ diluted in CH₂Cl₂ at 100 °C during 4 hr. Three ml of NaHCO₃
1091 (0.2 M) were added after cooling and mixed prior to the addition of 0.1 mL of HCl (6 M).
1092 The tubes were vortexed after addition of 3 mL of dichloromethane and the lower organic
1093 phase collected in a new tube before concentration under nitrogen atmosphere to
1094 approximately 0.5 ml. Then, 10 µL of the mixture was trimethylsilylated (TMS) with 100
1095 µL of N,O- bis(trimethylsilyl) trifluoroacetamide and 10 µL of ACS- grade pyridine. The
1096 trimethylsilylated samples were injected (1 µL) onto an Agilent 5973 Gas
1097 Chromatography–Mass Spectrometry system. Specific ion chromatograms reconstructed
1098 at m/z 239, 269 and 299 were used to quantify H, G and S lignin monomers respectively
1099 and compared to the internal standard at m/z 57, 71, 85.

1100

1101 **Neutral sugar content analysis**

1102 This method is adapted from ⁶⁶. Neutral monosaccharide sugar content was determined
1103 by gas chromatography after acid hydrolysis and conversion of monomers into alditol
1104 acetates as described in Hoebler et al., 1989, and Blakeney et al., 1983. Gas
1105 chromatography was performed on a DB 225 capillary column (J&W Scientific, Folsom,
1106 CA, USA; temperature 205 °C, carrier gas H₂). Calibration was made with standard sugar
1107 solution and inositol as internal standard.

1108

1109 **Synthesis of methylated sugars (Figure S14)**

1110 Please see supplemental methods. The structures of compounds were ascertained by
1111 NMR spectroscopy and were in agreement with reported data.

1112

1113 **Constructing phylogenetic trees for genes controlling primary and secondary cell** 1114 **wall formation**

1115 Genes controlling primary and secondary cell wall formation were identified from a
1116 previous study of *A. thaliana*¹²⁰. Using the known genes as reference, we utilised
1117 OrthoFinder (v.2.5.4)¹³⁴ with 49 species within Table S9. Genes that were grouped in the
1118 same orthogroups with the reference genes were aligned via MUSCLE (v.5.1)¹²⁸ and
1119 analysed via IQTree (v.1.6.12)¹³⁸ to construct trees with bootstrap values of 100 (for gene
1120 families with more than 600 genes) and 1000 (for those less than 600 genes). Trees were
1121 then visualised using MEGAX (v.1.0)¹⁵³, with bootstrap values less than 80% being
1122 condensed. Ancestor-specific duplication events were inferred using trees generated
1123 using previous phylostratigraphic analysis, where branches containing common earliest
1124 ancestors were deemed as such.

1125

1126 **Visualisation of genes controlling primary and secondary cell wall formation**

1127 Annotations of fern genes were based on the phylogenetic trees in the methods
1128 mentioned above, with genes annotated where reference genes from *A. thaliana* were
1129 found. Coexpression coefficients of each gene within 26 fern species were calculated
1130 using Pearson's correlation coefficient (PCC) and transformed into Highest Reciprocal
1131 Rank (HRR)¹⁵⁴ and genes of interest (GOIs) were isolated. Coexpressed GOIs were
1132 visualised using Cytoscape (v 3.10.1) (<https://cytoscape.org/>).

1133

1134 **Analysis of CAZymes and HRGPs**

1135 Protein files from coding sequences of 39 plant species were submitted to the dbCAN2
1136 pipeline¹⁵⁵, which annotates CAZymes using three tools (HMMer against the CAZyme
1137 domain database; DIAMOND for BLASTP against the CAZyme database; dbCAN-sub for
1138 HMMER detection of putative CAZy substrates). A majority vote was used and all
1139 annotations were supported by two or more tools, which were used to filter for relevant
1140 CAZy families. CAZy families, as well as respective functionally described enzymes, were
1141 aligned using MAFFT¹³⁶ (preferably L-INS-i; if the sequence dataset was too big the
1142 automatic mode was used). Sequence alignments were submitted to FastTree in default
1143 mode¹³¹, and homologs of functionally described enzymes were filtered using iTOL¹⁵⁶.
1144 For DUF families (DUF579, DUF231) and selected other enzymes (CGR2-3, BS1,
1145 DARX1, P4H, QUA2 and QUA3), BLASTP was used with the described family members

1146 against the protein files as a database. E-value of e^{-7} was used, with the rest followed as
1147 described above.

1148 Annotation of hydroxyproline-rich glycoproteins (HRGPs) was performed by using
1149 the workflow described in ⁷³. The protein sequences were filtered first for the presence of
1150 N-terminal signal peptides and then classified into 24 classes based on the presence of
1151 distinct amino acid motifs and biases (as outlined in ¹⁵⁷).

1152

1153 **Comprehensive Microarray Polymer Profiling (CoMPP)**

1154 The CoMPP analysis was performed according to the method reported by ⁸⁶. Each
1155 sample was weighed out in triplicate of 10 mg AIR. The samples were sequentially treated
1156 with 300 μ L 50 mM trans-1,2-diaminocyclohexane-N,N,N',N'-tetraacetic acid (CDTA) pH
1157 7.5, followed by extraction with 300 μ L 4 M NaOH containing 0.1% (v/v) NaBH₄. Each
1158 extraction step was carried out for 2 h in a TissueLyser II (Qiagen AB, Sollentuna,
1159 Sweden) at 6 s⁻¹ at room temperature. After each extraction, samples were centrifuged
1160 for 10 minutes at 4000 rpm, and the supernatant was collected. The samples were added
1161 to a 384 well plate and four dilution points were prepared for each sample, then two
1162 technical replicates printed on nitrocellulose using an ArrayJet Marathon printer (ArrayJet,
1163 Roslin, UK).

1164 Separate arrays for each probe were first blocked with 5% (w/v) low-fat milk
1165 powder solution in phosphate-buffered saline (MP/PBS), then probed with a set of specific
1166 primary monoclonal antibodies (LM, JIM and MAC 207; Plant Probes, Leeds university,
1167 CCRC; Complex Carbohydrate Research Center, University of Georgia, BS-400;
1168 Biosupplies Australia and INRA-RU donated from Marie Christine Ralet, INRA, France)
1169 (Table S14) for 2 h. After three washes with PBS, the arrays were incubated with 1:5000
1170 solutions of either anti-mouse or anti-rat secondary antibodies (depending on the source
1171 of the primary antibody) conjugated with alkaline phosphatase for another 2 h. Following
1172 three washes with PBS the array had a final wash in Milli-Q water. Arrays were developed
1173 with a 5-bromo-4-chloro-3-indolyl-phosphate (BCIP)/nitro-blue tetrazolium chloride (NBT)
1174 substrate and scanned using a flatbed scanner (CanoScan 9000 Mark II; Canon, Søborg,
1175 Denmark) at 2400 dpi converting the dots to grayscale. The calculated intensity of the
1176 signal was quantified using the microarray analysis software ProScanArray Express

1177 (PerkinElmer, Waltham, Massachusetts, USA). The relative intensity values were
1178 normalised to a scale from 0 to 100 and transformed into a heatmap.

1179

1180 **Enrichment and depletion of biological processes in fern-specific genes and** 1181 **neighbourhood**

1182 Fern-specific genes were defined as genes within orthogroups located in nodes 6-13
1183 (Figure 2a), excluding those orthogroups that contained only genes from one species.
1184 Neighbours of fern-specific genes were defined as genes that are co-expressed with a
1185 fern-specific gene and are not fern-specific genes themselves. The functional annotations
1186 of genes were retrieved (first-level Mapman bins) and subjected to enrichment and
1187 depletion analysis against a background of genes assigned to orthogroups. The analysis
1188 was performed for each fern using a hypergeometric test and adjusted for multiple testing
1189 via Benjamini-Hoechberg correction ($q < 0.05$)¹⁵⁸. The overall trend of enrichment or
1190 depletion of biological processes across fern species was derived by subtracting the
1191 number of depleted Mapman bins across all species from the number of enriched bins.

1192

1193 **Data availability**

1194 The raw sequencing data is available at E-MTAB-13848, while the CDS and protein
1195 sequences are found at <https://doi.org/10.6084/m9.figshare.26347330>. The co-
1196 expression networks are available at <https://conekt.sbs.ntu.edu.sg/species/>.

1197

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1213

1214 **Supplementary Methods**

1215 **Supplementary Methods 1: Inferences of species tree, whole genome duplications,**
1216 **analysis of salicylic acid- and jasmonic acid-mediated signalling in ferns, chemical**
1217 **synthesis of methylated sugars.**

1218

1219

1220 **Supplementary tables:**

1221 **Table S1. 22 species of Tracheophyta from 22 different families.** Samples were
1222 collected from various locations in Singapore, with each species having multiple organs
1223 harvested.

1224 **Table S2. Sequencing statistics.** The columns contain descriptions of the 415 sample
1225 names, including the species, organs, organ types, and data statistics.

1226 **Table S3. Transcriptome assembly statistics for the 22 ferns.** BUSCO value,
1227 MapMan annotation percentage, number of transcripts, GC% content, N50, BUSCO
1228 scores and percentage of genes annotated by MapMan are shown.

1229 **Table S4 Clade, order, family, species and source of data of the 108 ferns used in**
1230 **this study.**

1231 **Table S5. Fossil calibrations of soft constraint adopted in this study.**

1232 **Table S6. The 95% HPD and posterior mean age estimates (mya) for the origin of**
1233 **each major clade**

1234 **Table S7. Phylostratigraphic assignments of orthogroups to nodes.** The table shows
1235 the orthogroups, clades which are present in the orthogroup and the node where the
1236 orthogroup appeared in.

1237 **Table S8. Gene-Organ Specificity.** The table shows the different species (given by
1238 mnemonic), SPM value in a given sample, the number of genes in an organ and the gene
1239 ids specifically expressed in the organ.

1240 **Table S9. Missing/Present Mapman Bins across 39 species, comprising of**
1241 **Glaucophytes, Chlorophytes, Bryophytes, Lycophytes and Ferns.**

1242 **Table S10. The percentage of annotated clades by MapMan bins.** The species
1243 comprising the clades are indicated in column B.

1244 **Table S11. CASA lignin content and thioacidolysis analysis, showing content of H,**
1245 **G and S units.**

1246 **Table S12, BLAST scores of Arabidopsis and Selaginella lignin-related genes**
1247 **against 39 species contained in conekt.sbs.ntu.edu.sg.**

1248 **Table S13. Neutral Sugar Analysis.** The different species and their organs are shown
1249 in rows. The columns indicate the abundance of sugars and the standard deviation.

1250 **Table S14. Antibodies, their immunogens and declared specificities and the**
1251 **references where the antibodies were generated/described.**

1252 **Table S15 CoMPP profiles of the 102 cell wall extracts probed with 48 antibodies.**
1253 The used solvent and antibodies are shown in columns, the species, organs are shown
1254 in rows.

1255

1256 **Supplementary Figures:**

1257 **Figure S1. Pictures of the 22 ferns and their sampled organs.**

1258 **Figure S2. Transcriptome assembly (blue boxes) and subsequent analyses (red**
1259 **boxes).**

1260 **Figure S3.** Genomic properties of ferns in relation to whole genome duplication events.
1261 The plots show the correlation between WGD events (x-axis) and species richness (the
1262 number of species within a lineage), holoploid genome size (total DNA content),
1263 monoploid genome size (DNA content of a single set of chromosomes) and others.

1264 **Figure S4. AlphaFold3-derived structures of the 17 fern-specific proteins.** The colors
1265 indicate confidence scores of the structures.

1266 **Figure S5. Number of genes (y-axis) with a given SPM value (x-axis).** The SPM value
1267 cutoff is indicated by the red line.

1268 **Figure S6. Gene expression profiles of organ-specific genes.** Each gene's
1269 expression has been scaled to range from 0 to 1.

1270 **Figure S7. Expression profiles for species-specific genes.**

1271 **Figure S8. Transcriptome similarity comparison of Archaeplastida.** The heatmap
1272 shows the conservation of organ-specific orthogroups across the species. The Jaccard
1273 index of across species similarities are indicated by red shades, and for within species
1274 similarly with blue shades.

1275 **Figure S9. Gene functions found in Archaeplastida.** Mapman bins (rows) are found in
1276 the different species (columns). The colours indicate the fraction of found bins in a given
1277 species, where 1 indicates that all genes in a given bin are present, while 0 indicates
1278 complete absence.

1279 **Figure S10. Enrichment and depletion of biological processes in neighbours of**
1280 **fern-specific genes.** Cluster map showing significantly enriched and depleted primary
1281 Mapman bins (y-axis) in neighbours of fern-specific genes across ferns. The colour map
1282 indicates the significance of the biological processes, with yellow representing a p-value
1283 of 0.05 and blue representing a p-value of 0.00. P-values above 0.05 are masked.

1284 **Figure S11. Light field, fluorescence, phloroglucinol and Maule staining of the**
1285 **selected ferns.**

1286 **Figure S12. The number (x-axis) of the transcription factors, CYP450s enzyme**
1287 **families and lignin-related enzymes co-expressed with at least two lignin**
1288 **biosynthetic enzymes in the analyzed ferns.**

1289 **Figure S13. GC-MS analysis of 3O-MeRhap.** a) MS profile of 3O-MeRhap. b) *Tectaria*
1290 *incisa* contains 3O-MeRhap (read arrow) and rhamnose. c) *Psilotum nudum* contains
1291 rhamnose but no 3O-MeRhap.

1292 **Figure S14. The protocol of chemical synthesis of 2-O-methyl- and 3-O-methyl- α,β -**
1293 **D-galactopyranose and 2-O-methyl-D-glucopyranose.**

1294 **Figure S15. GC-MS analysis of cell wall sugars.** a) Profile of *Tectaria incisa*, b) *Tectaria*
1295 *incisa* and 2O-MeGlc standard and c) 2O-MeGlc standard. d) GC-MS spectra of the
1296 unknown peak and the methylated sugar standards.

1297 **Figure S16. Gene copy number analysis of hydroxyproline-rich glycoproteins**
1298 **(HRGPs).** Columns represent species, while rows correspond to a given class of HRGP.

1299 Red and blue numbers indicate that a given species contains significantly more/less
1300 genes than others.

1301 **Figure S17. CoMPP analysis of ferns.** a) The clustermap shows fern samples (rows)
1302 and antibodies (columns). The cells indicate the signal scaled from 0 (dark blue) to 1
1303 (bright yellow). b) Pearson Correlation Coefficient (PCC) distribution of CoMPP profiles
1304 within (blue) and across (brown) species.

1305 **Figure S18. Phylogenetic analysis of CESA genes.** The blue circles represent
1306 bootstrap values (value <50 are not indicated by a circle). The leaf colors represent the
1307 different species and orders.

1308 **Figure S19.** Phylogenetic analysis of lignin biosynthetic genes. Any inferred duplication
1309 events are indicated.

1310

1311 **Supplementary Data 1. Co-expression networks of lignin-related genes, cellulose
1312 synthases and cell wall-related transcription factors.** The file should be opened in
1313 cytoscape.

1314

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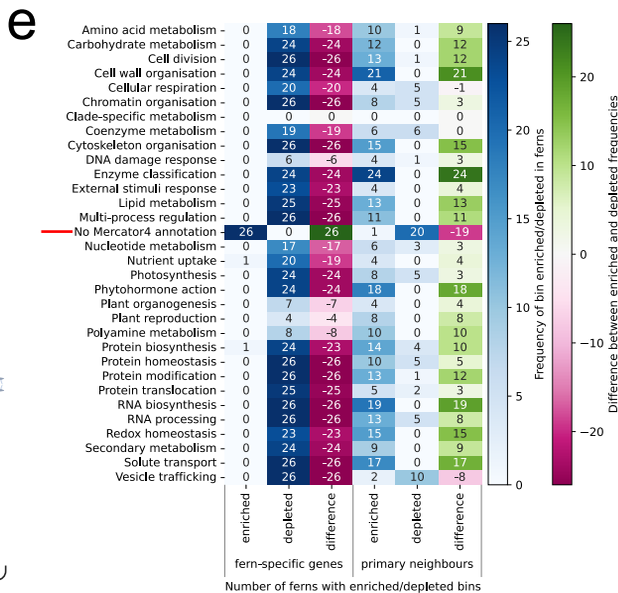
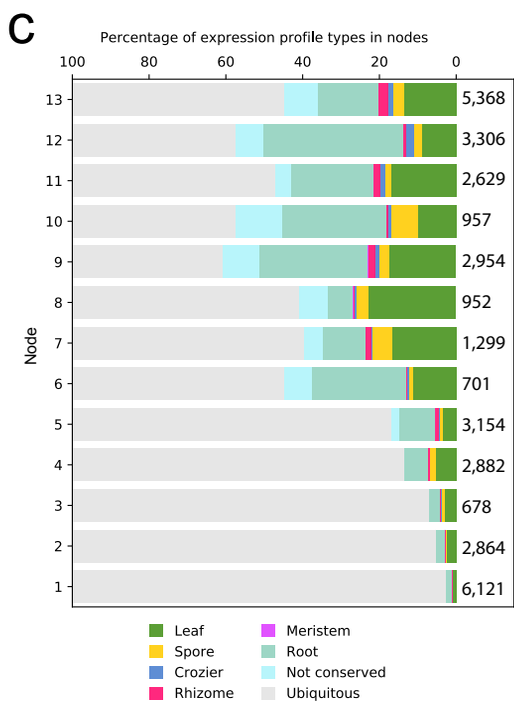
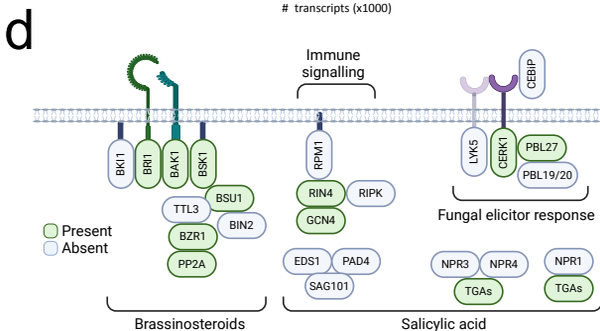
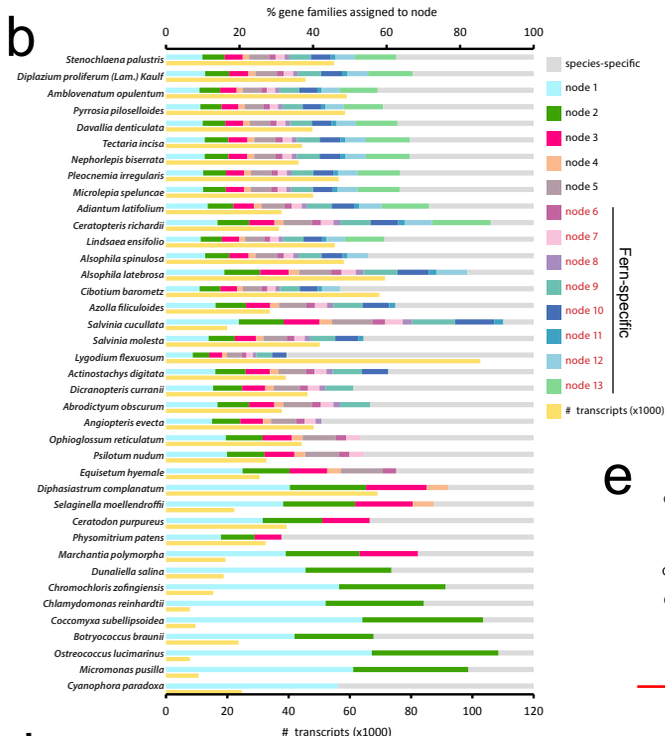
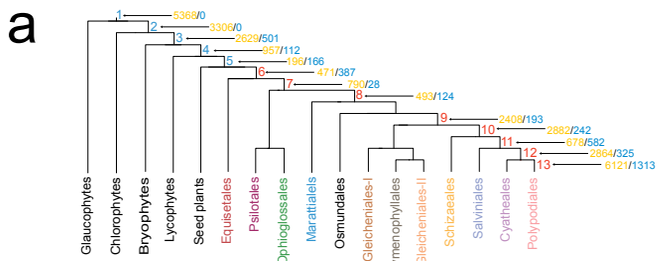
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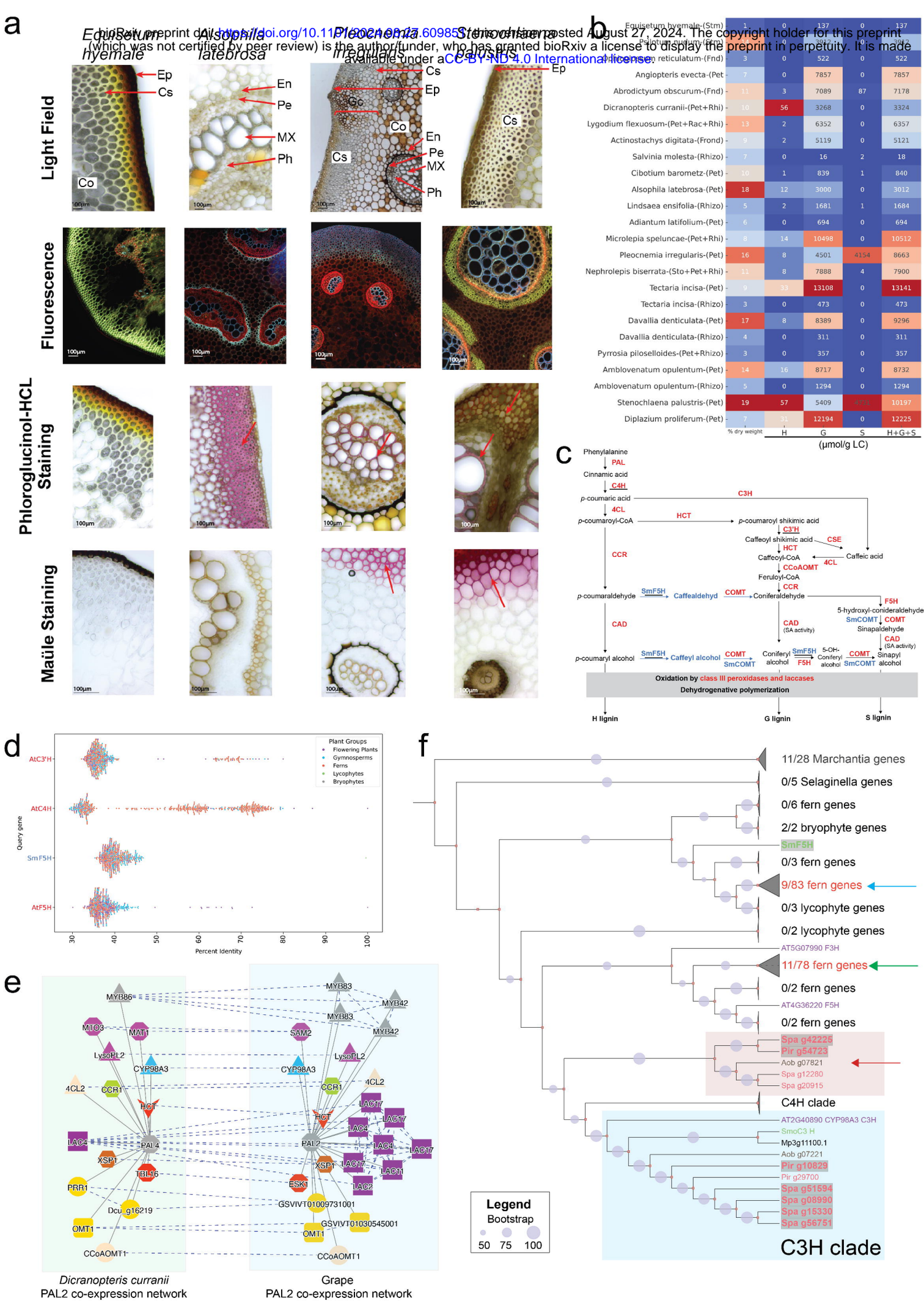
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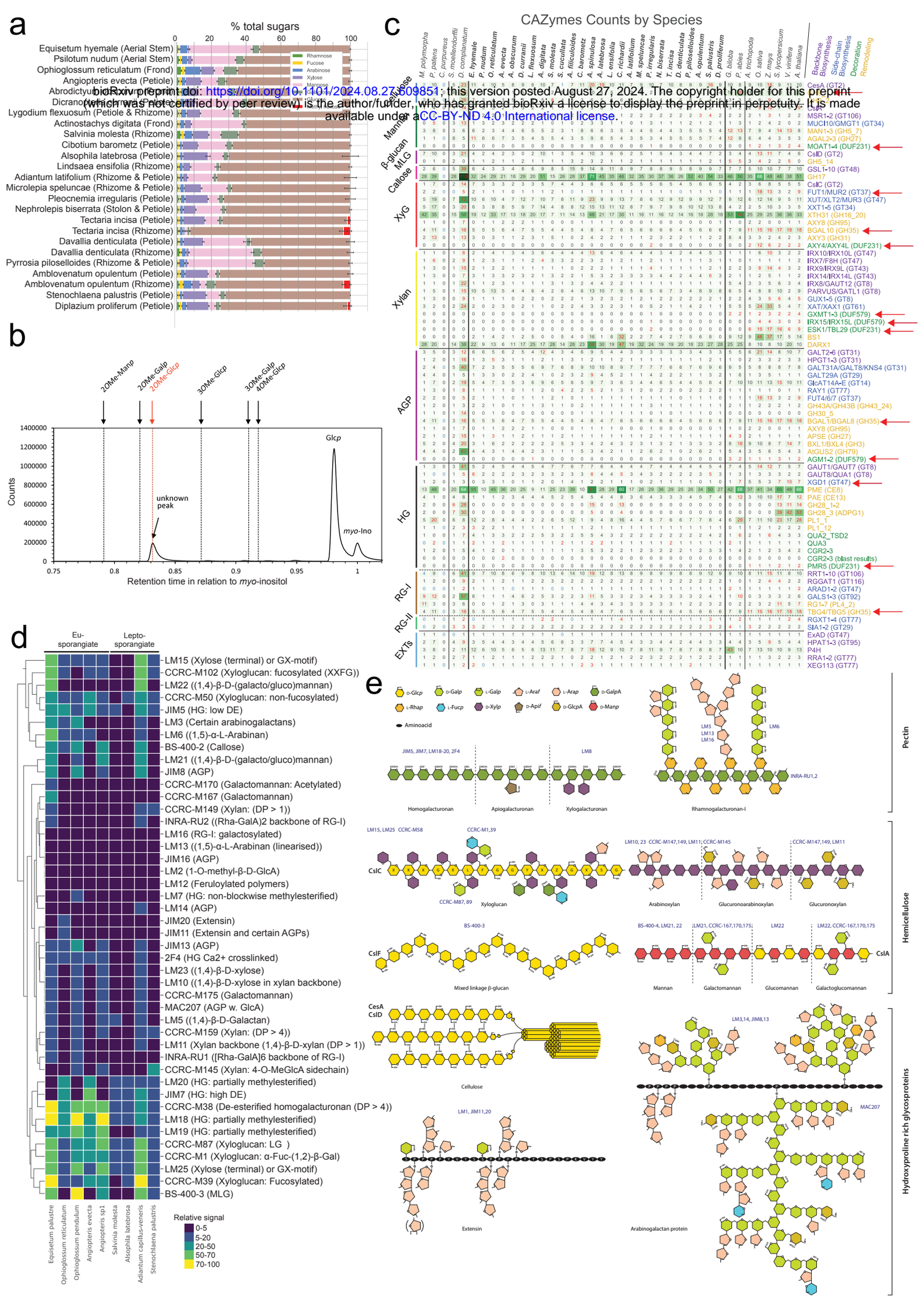
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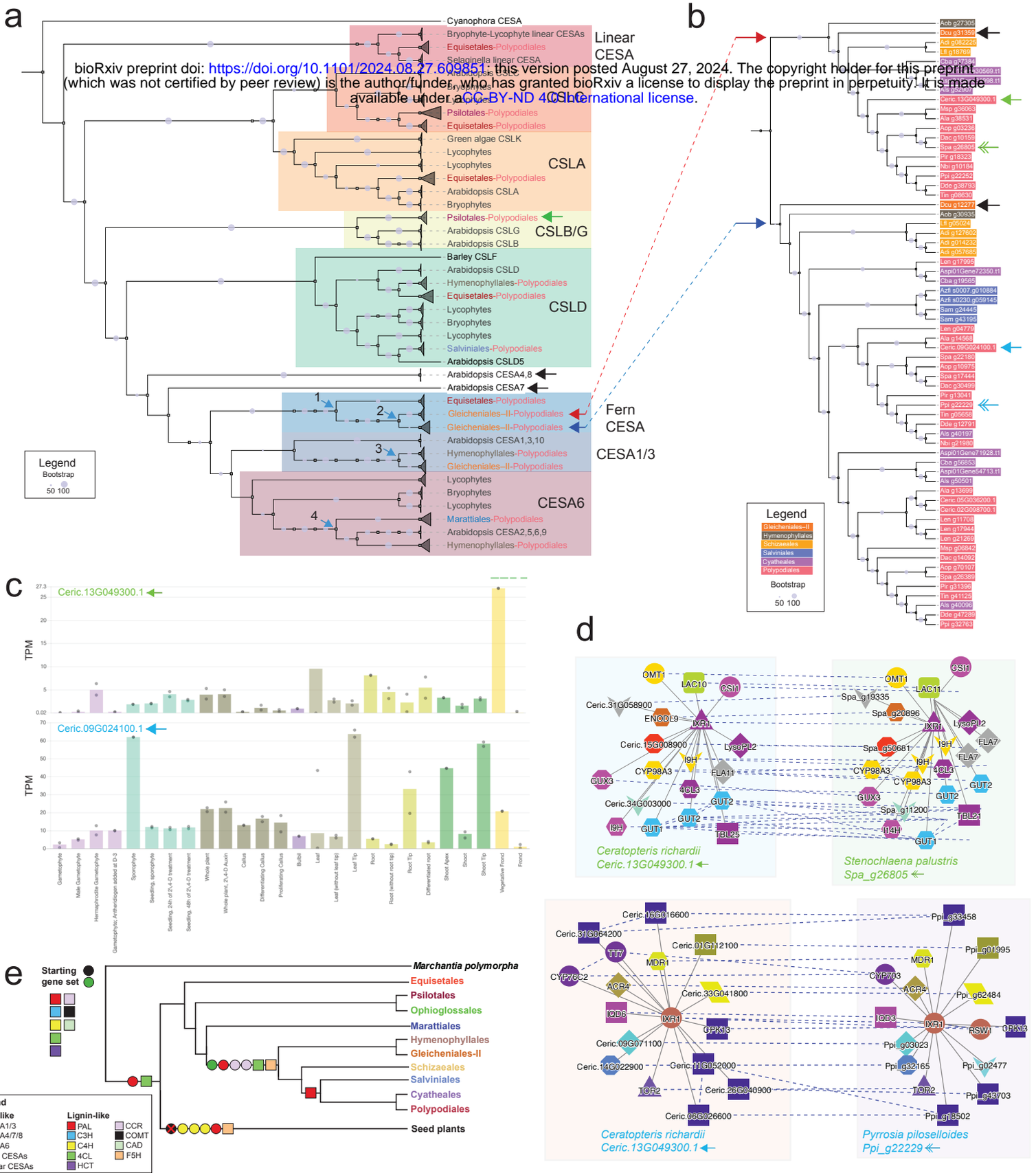
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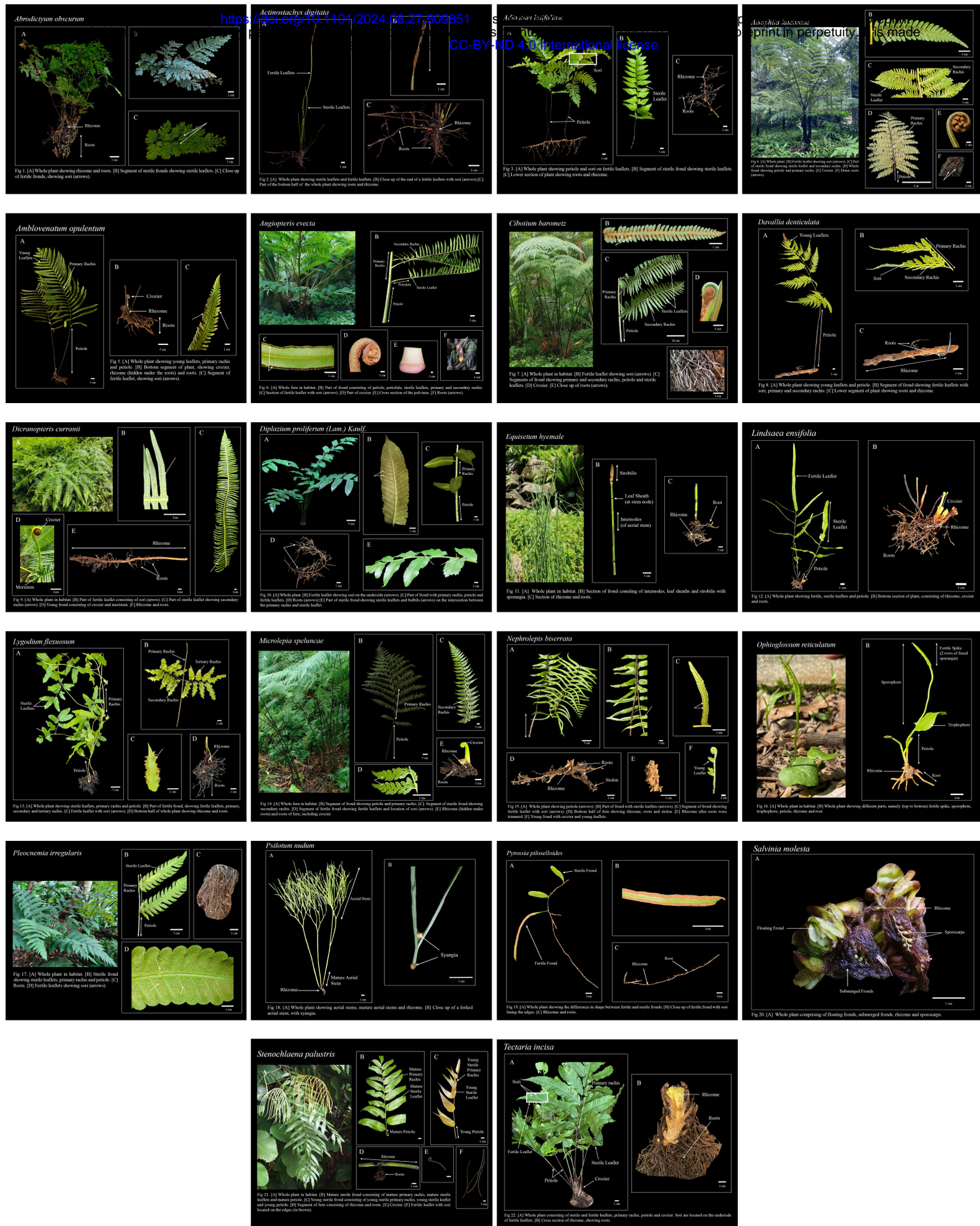


Figure S1. Pictures of the 22 ferns and their sampled organs.

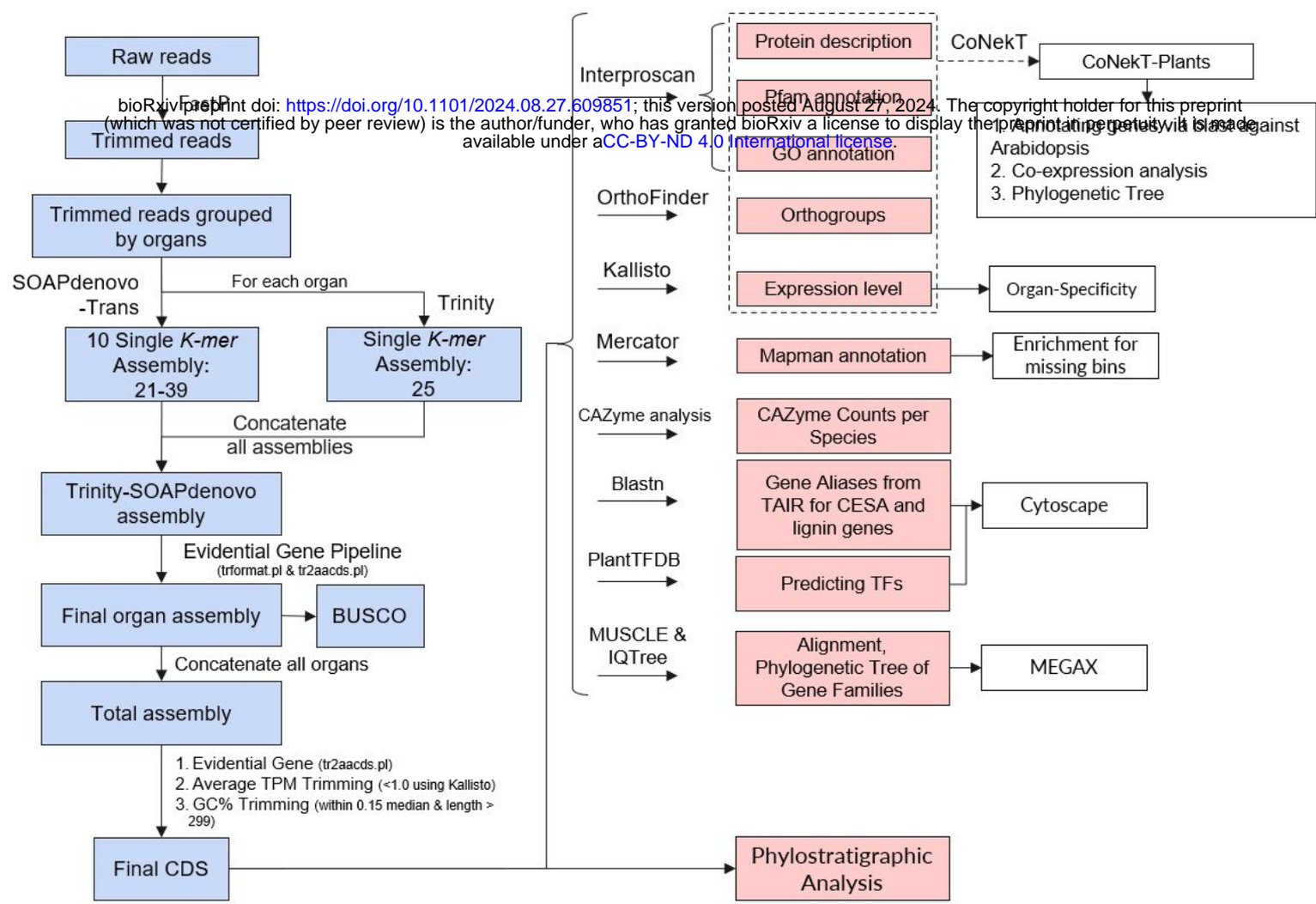


Figure S2. Transcriptome assembly (blue boxes) and subsequent analyses (red boxes).

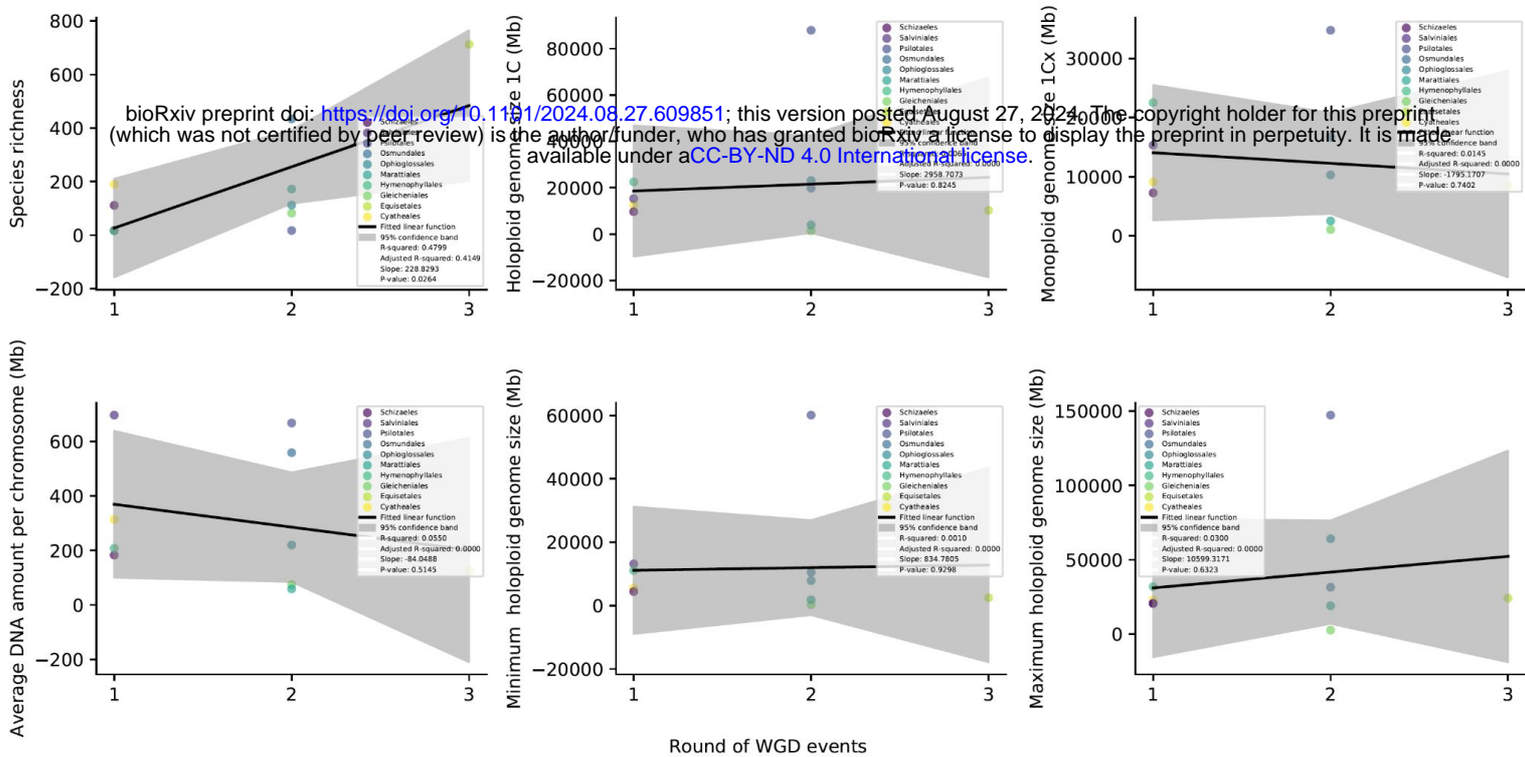


Figure S3. Genomic properties of ferns in relation to whole genome duplication events. The plots show the correlation between WGD events (x-axis) and species richness (the number of species within a lineage), holoploid genome size (total DNA content), monoploid genome size (DNA content of a single set of chromosomes) and others.

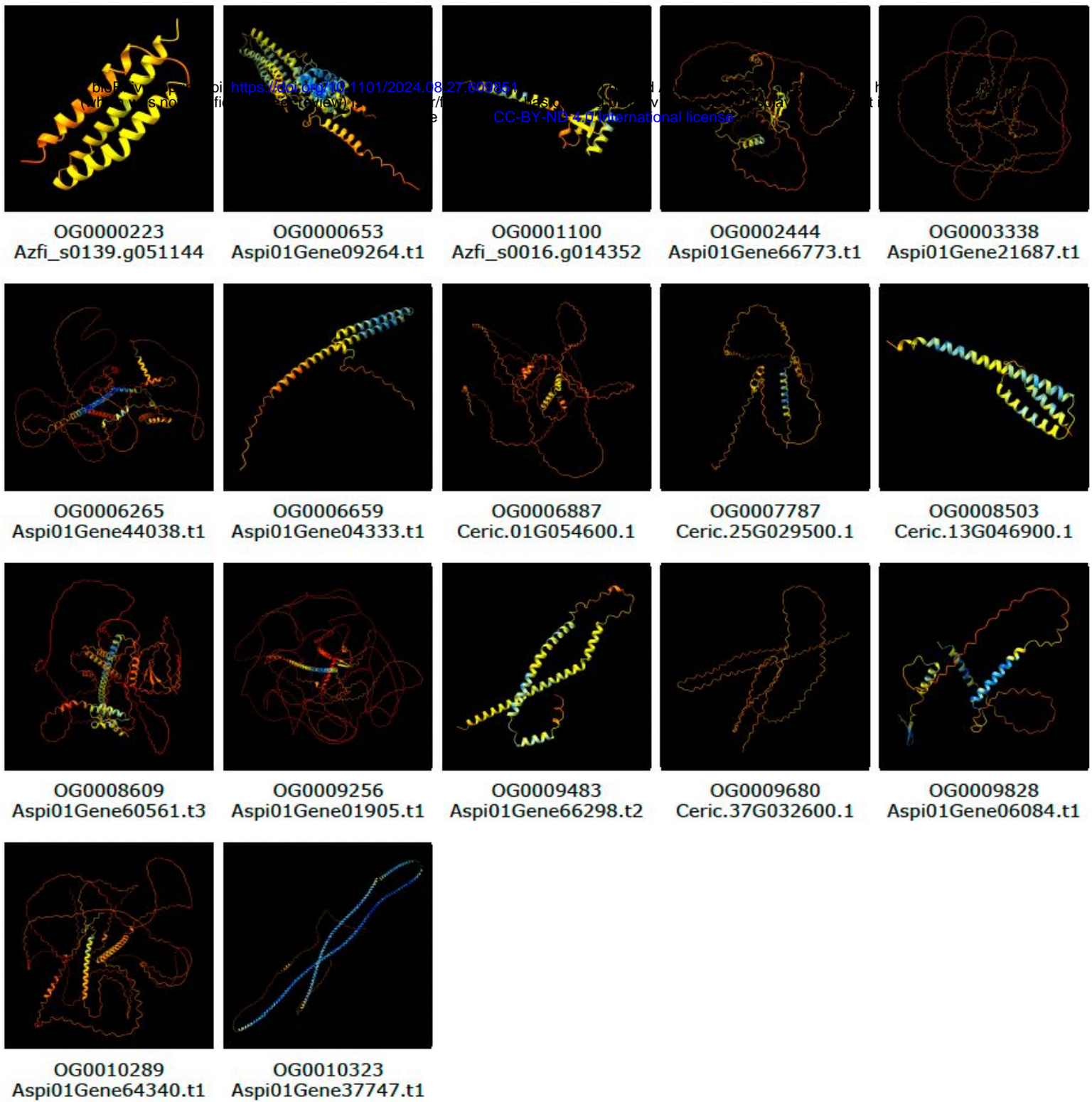


Figure S4. AlphaFold3-derived structures of the 17 fern-specific proteins. The colors indicate confidence scores of the structures.

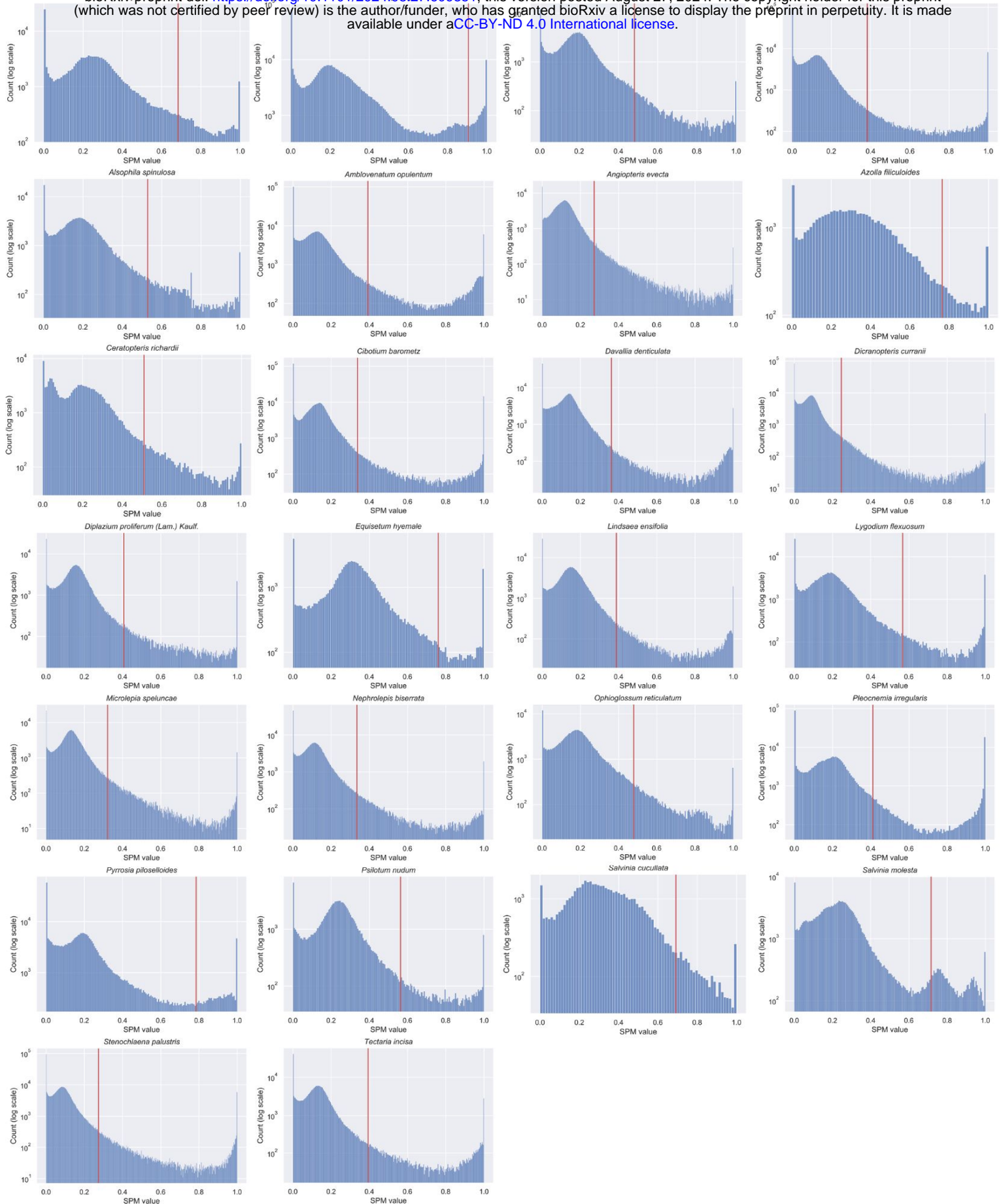


Figure S5. Number of genes (y-axis) with a given SPM value (x-axis). The SPM value cutoff is indicated by the red line.

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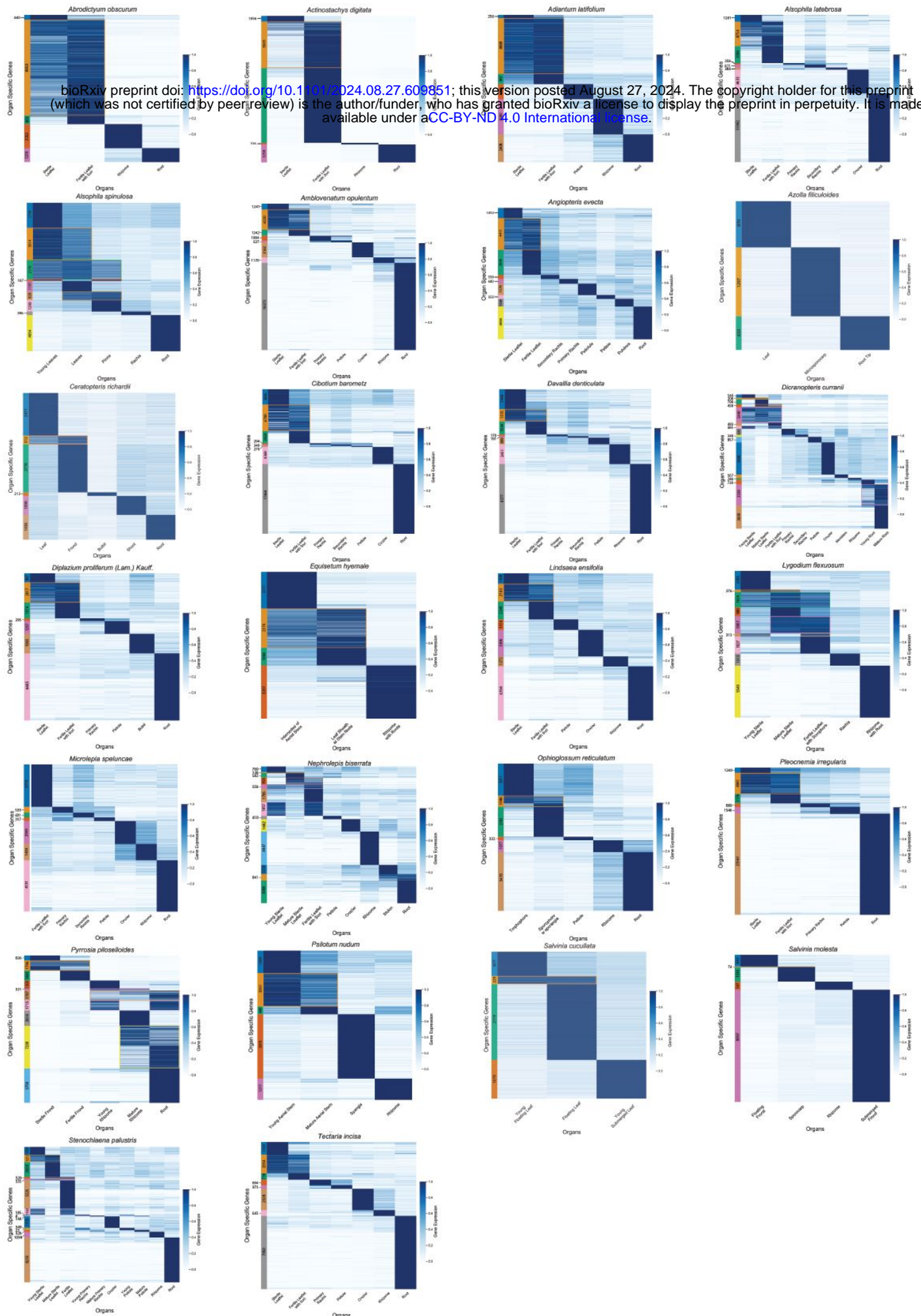


Figure S6 Gene expression profiles of organ-specific genes. Each gene's expression has been scaled to range from 0 to 1.

Percentage of expression profile types in species

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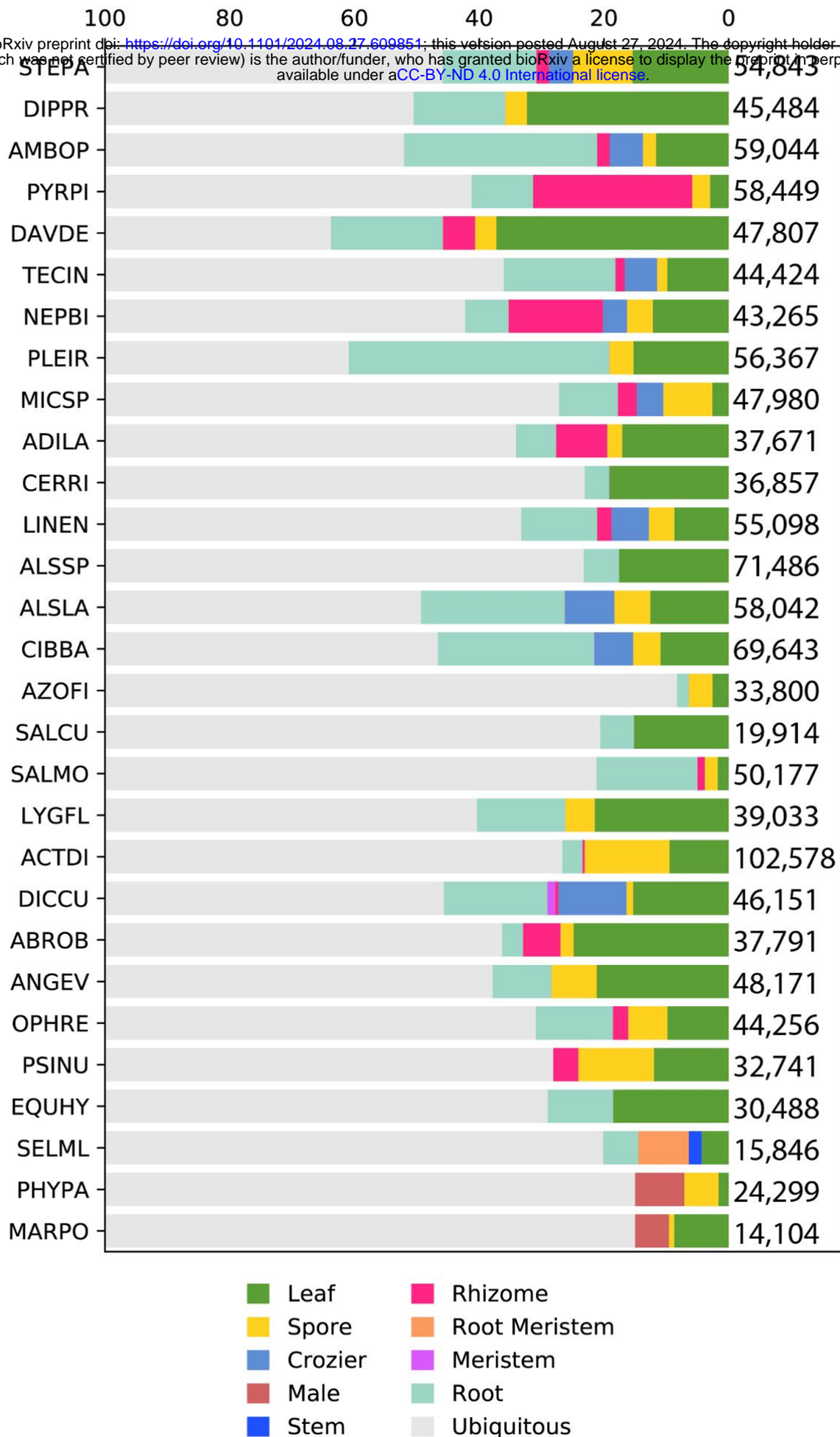


Figure S7. Expression profiles for species-specific genes.

Percentage of Mapman Bins

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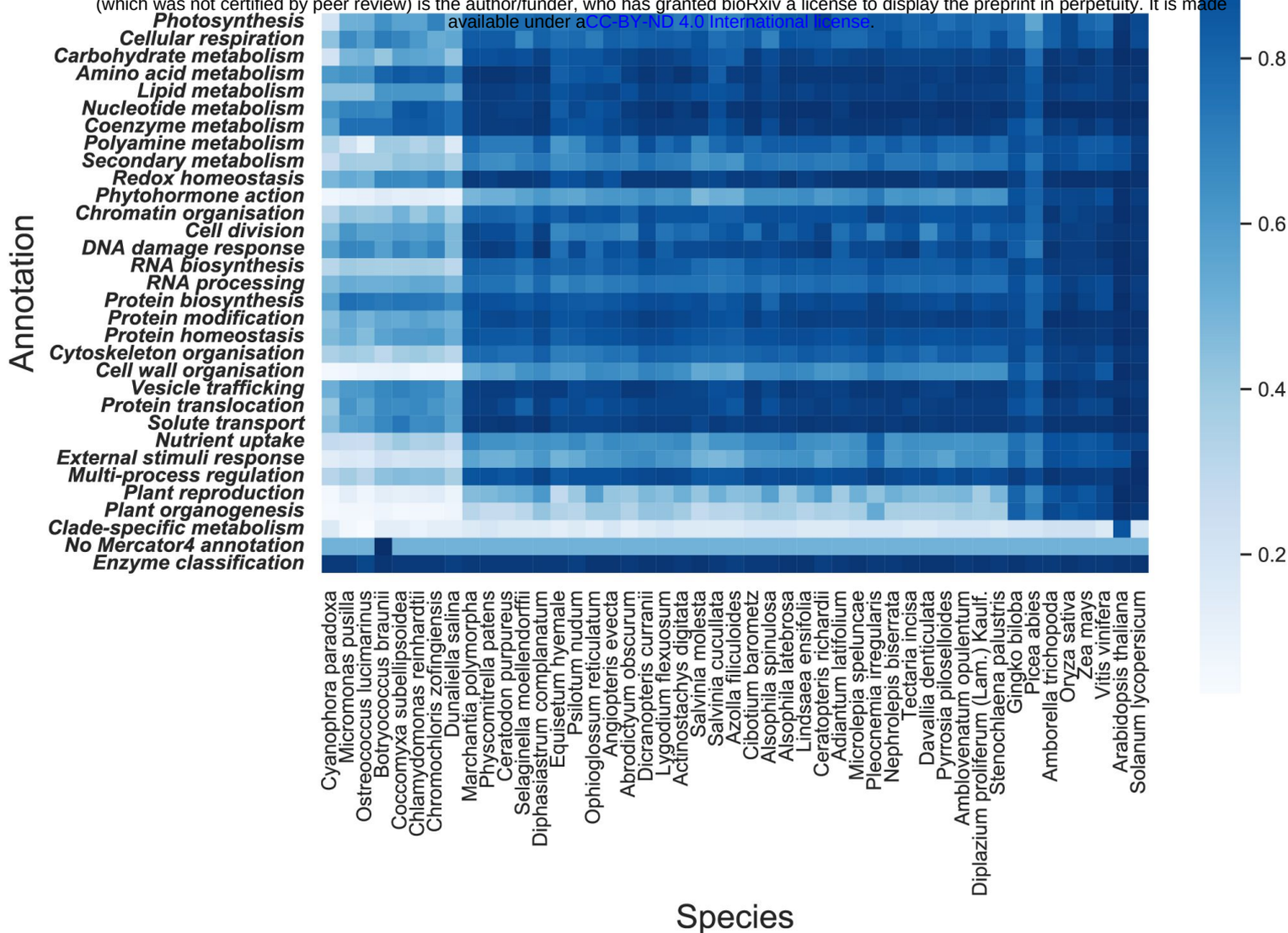


Figure S9. Gene functions found in Archaeplastida. Mapman bins (rows) found in the different species (columns). The colours indicate the fraction of found bins in a given species, where 1 indicates that all genes in a given bin are present, while 0 indicates complete absence.

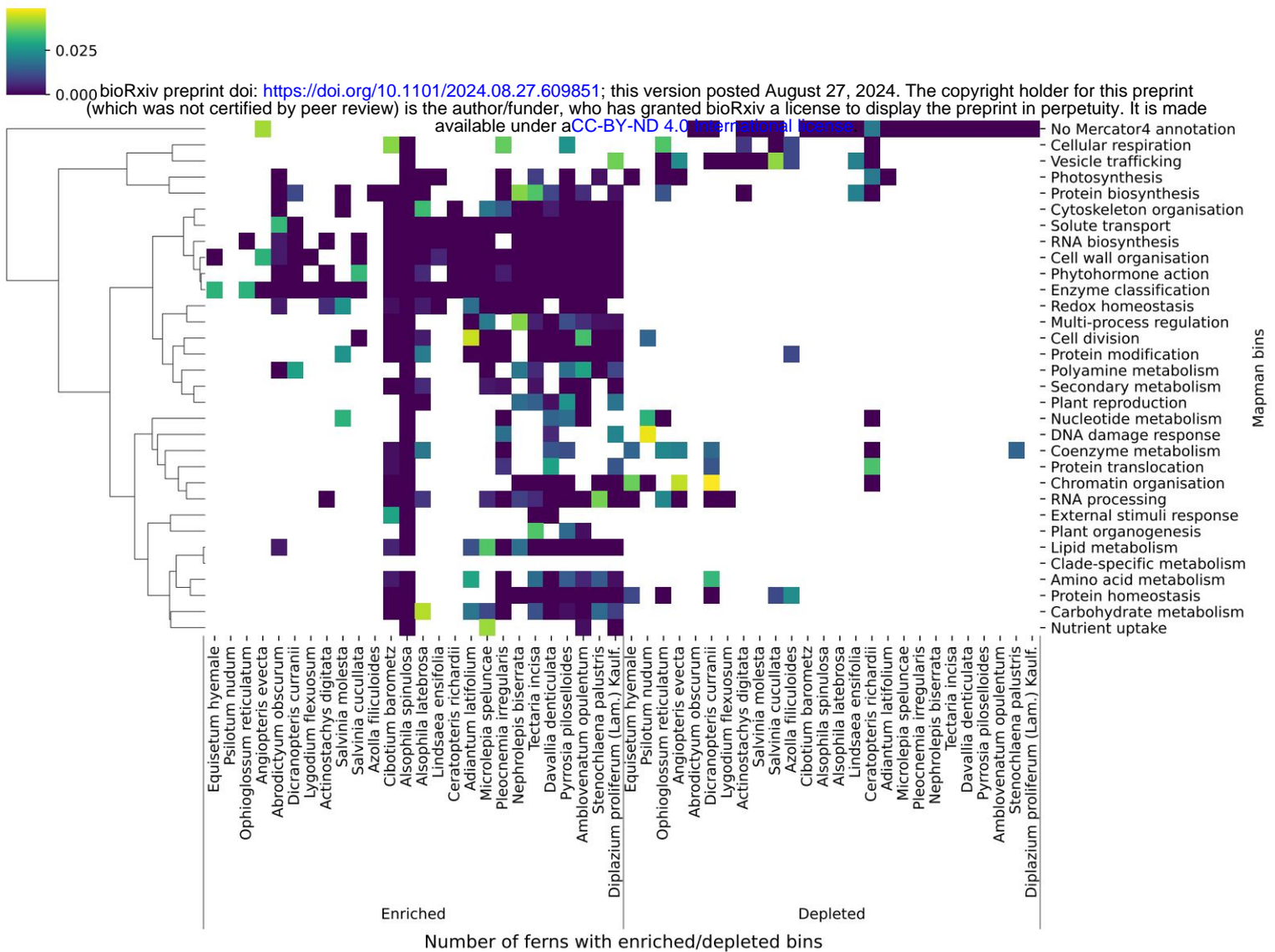
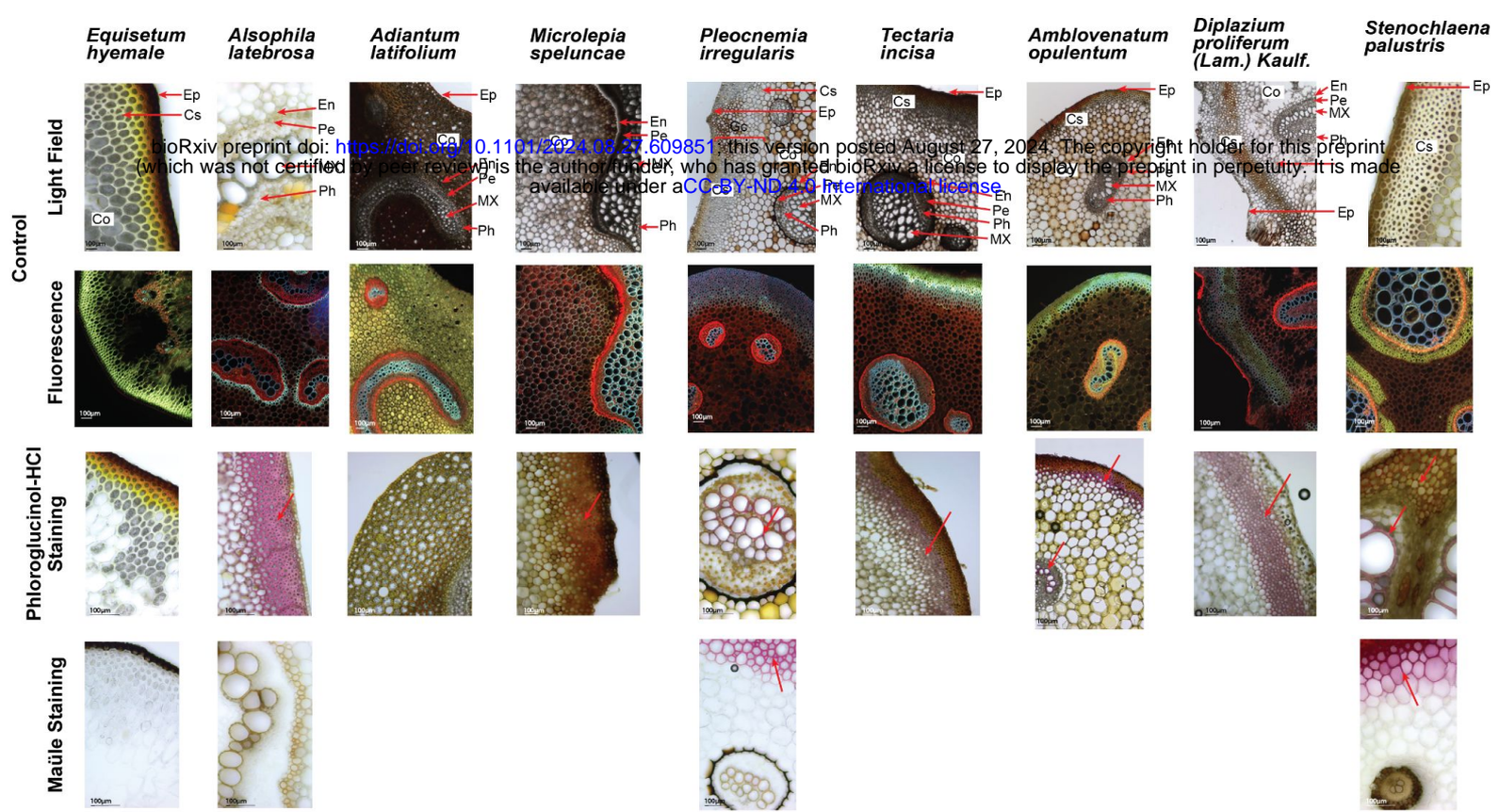


Figure S10. Enrichment and depletion of biological processes in neighbours of fern-specific genes. Clustermap showing significantly enriched and depleted primary Mapman bins (y-axis) in neighbours of fern-specific genes across ferns. The colour map indicates the significance of the biological processes, with yellow representing a p-value of 0.05 and blue representing a p-value of 0.00. P-values above 0.05 are masked.



Ep: Epidermis, Cs: Cortical Sclerenchyma, Co: Cortex, En: Endodermis, Pe: Pericycle, Ph: Phloem, MX: Metaxylem

Figure S11. Light field, fluorescence, phloroglucinol and Maüle staining of the selected ferns.

Bar Chart of Gene Combinations Sorted by Descending Order

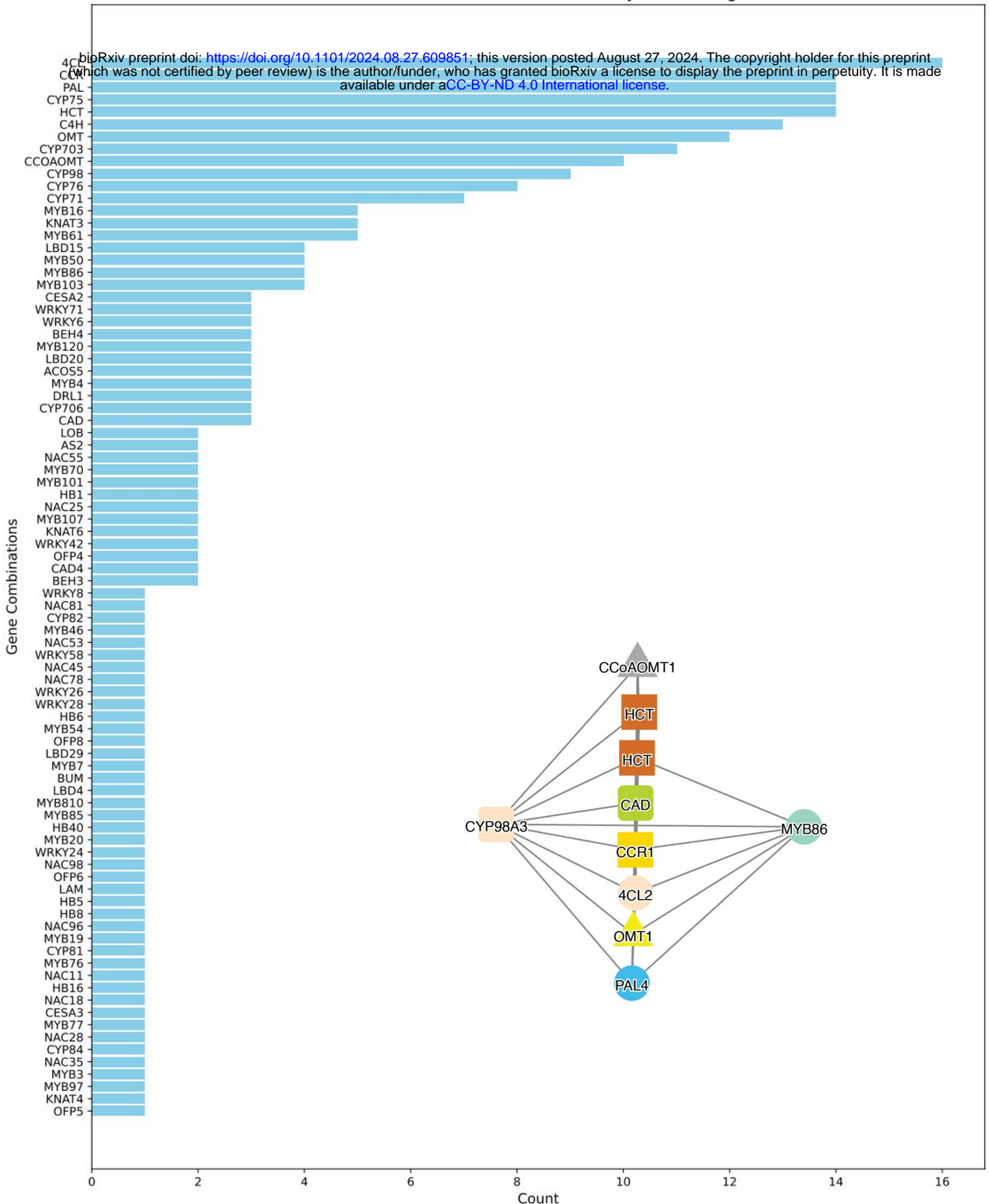
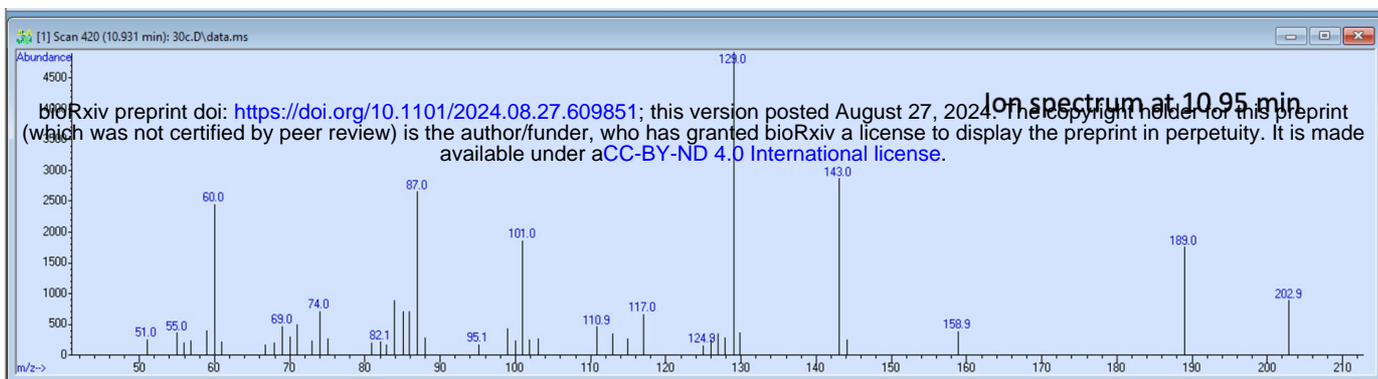
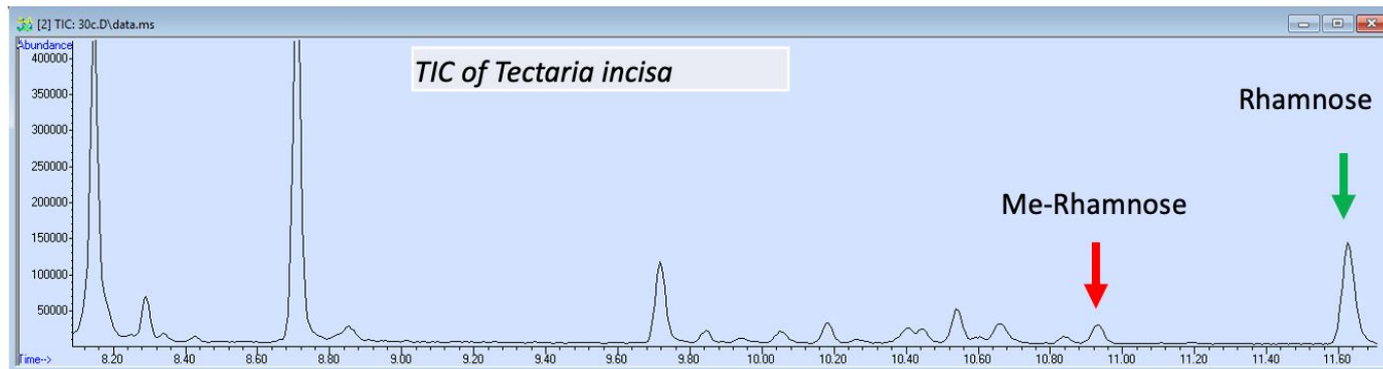


Figure S12. The number (x-axis) of the transcription factors, CYP450s enzyme families and lignin-related enzymes co-expressed with at least two lignin biosynthetic enzymes in the analyzed ferns. Co-expression network of PAL4 (blue circle) from *Dicranopteris curranii*. CYP96A3 and MYB86 are connected to seven and five enzymes involved in lignin biosynthesis.

a



b



c

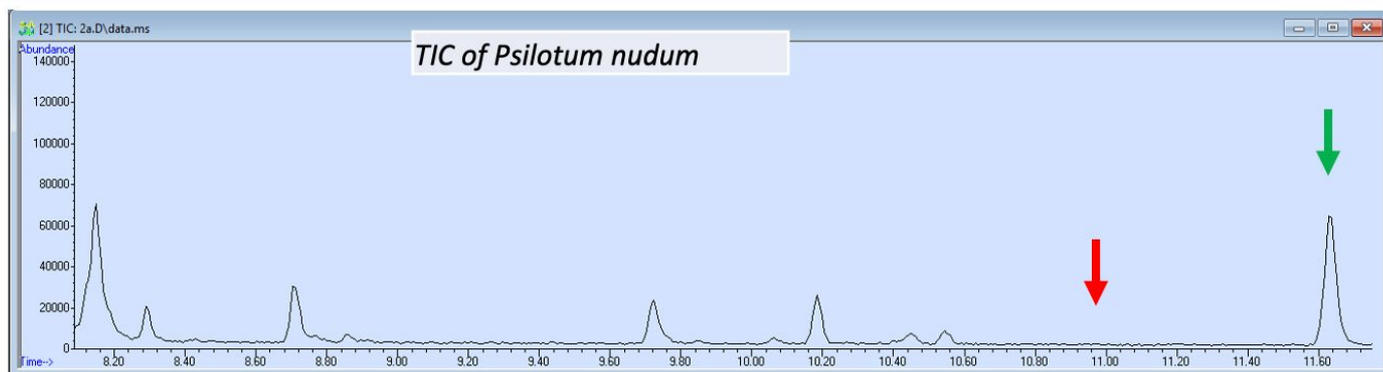


Figure S13. GC-MS analysis of 3O-MeRhap. a) MS profile of 3O-MeRhap. b) *Tectaria incisa* contains 3O-MeRhap (red arrow) and rhamnose. c) *Psilotum nudum* contains rhamnose but no 3O-MeRhap.

Chart 1: The protocol of chemical synthesis of 2-*O*-methyl- and 3-*O*-methyl- α,β -D-galactopyranose **5a** and **5b** and 2-*O*-methyl-D-glucopyranose **10**

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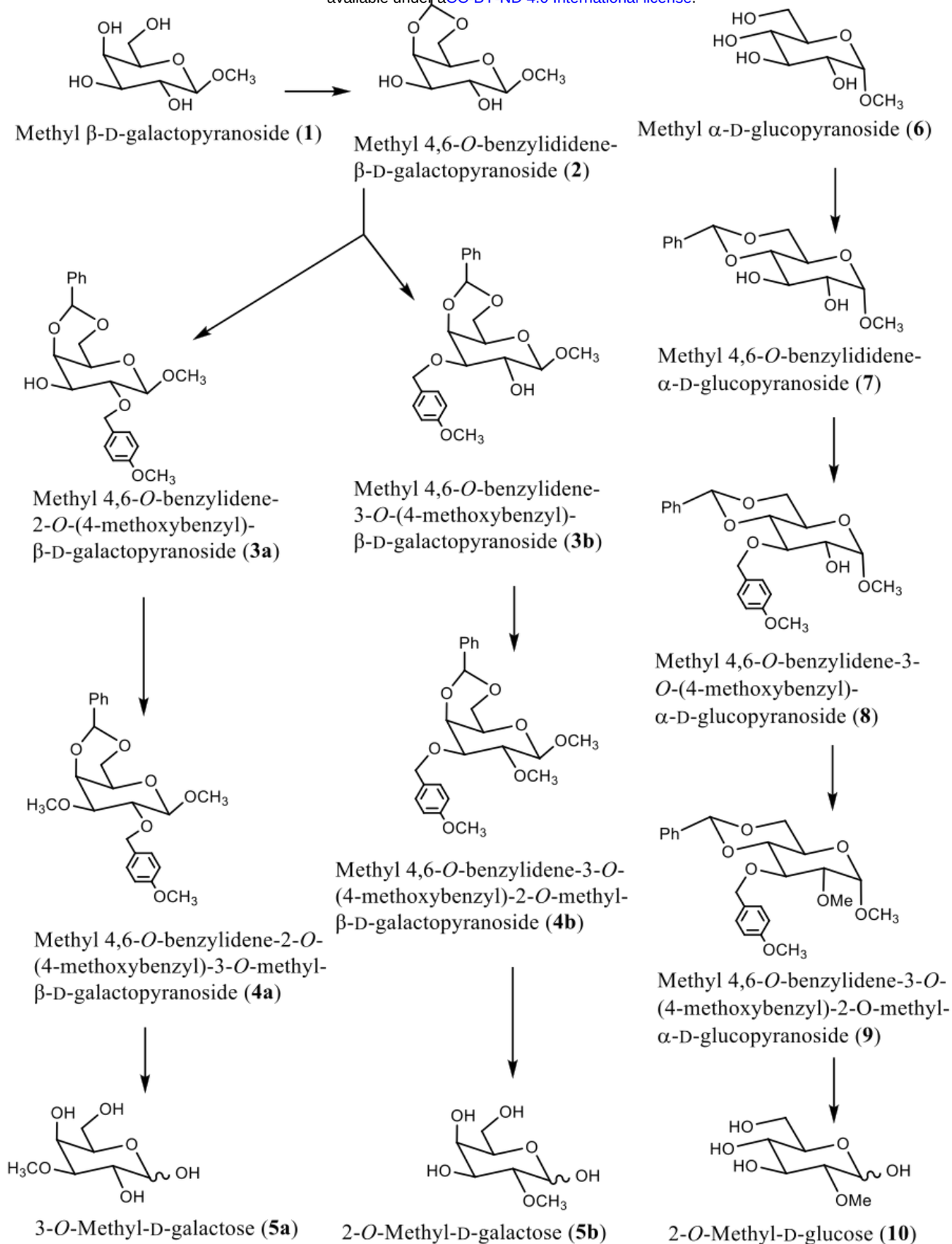


Figure S14. The protocol of chemical synthesis of 2-*O*-methyl- and 3-*O*-methyl- α,β -D-galactopyranose **5a** and **5b** and 2-*O*-methyl-D-glucopyranose **10**.

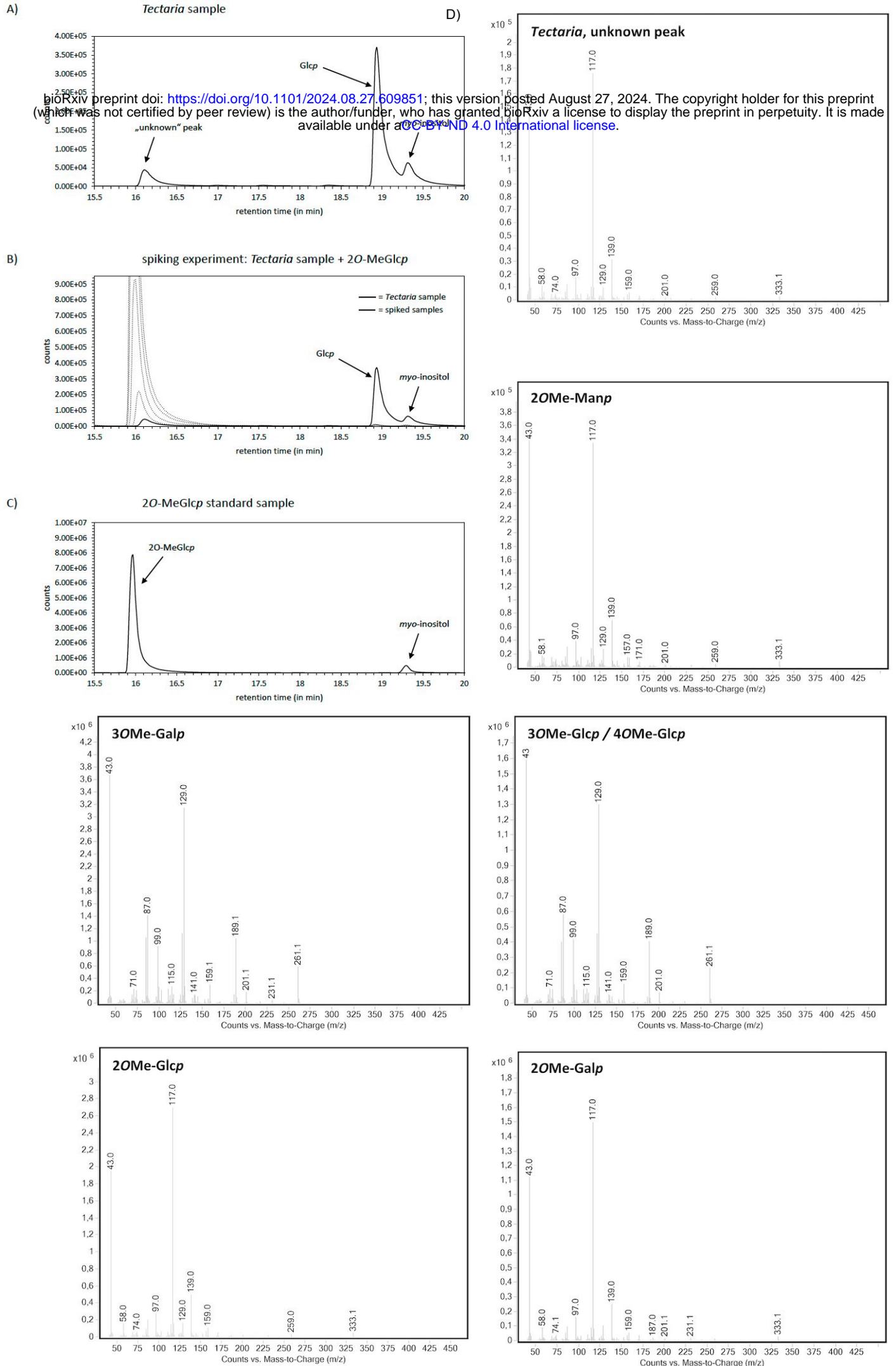


Figure S15. GC-MS analysis of cell wall sugars. a) Profile of *Tectaria incisa*, b) *Tectaria incisa* and 2O-MeGlcP standard and c) 2O-MeGlcP standard. d) GC-MS spectra of the unknown peak and the methylated sugar standards.

HRGP Counts by Species

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1KP Groups	M. polymorpha	P. patens	C. purpureus	S. moellerae	D. completum	E. hyemale	P. nudum	O. reticulatum	A. evecia	A. obscurum	D. curranii	L. flexuosum	A. digitata	S. moellerae	S. cucurbitaria	A. filicoides	C. barobitz	A. spiralis	A. latifolia	L. ensifolia	C. richardii	A. latifolia	M. spallanzanii	P. irregularis	N. bisecta	T. incisa	D. dentata	P. pilosoides	A. opulentum	S. palustris	D. proliferum	G. bilobata	P. abies	A. trichomanes	O. sativa	Z. mays	S. lycopersium	V. vinifera	A. thaliana		
GPI-AGPs	20	16	6	3	2	1	2	7	2	11	6	1	14	4	3	0	22	3	3	9	9	3	10	13	10	0	1	1	8	6	4	5	9	3	12	20	11	0	17		
CL-extensins	1	0	1	6	5	2	0	2	1	4	2	2	2	0	1	0	2	1	3	4	5	0	3	4	1	2	3	1	6	7	3	8	3	4	0	0	5	0	18		
PRPs	0	0	0	0	0	0	0	0	1	0	0	0	1	0	0	0	0	0	0	0	1	0	0	0	0	0	1	0	2	0	0	0	0	0	0	0	0	0	0	0	
non-GPI-AGPs	9	18	28	8	13	9	7	11	5	11	10	9	38	13	4	6	18	15	11	13	29	4	9	8	8	23	15	28	26	16	9	8	3	5	21	25	14	1	7		
Hybrid AGPs	0	1	0	1	0	0	0	0	0	0	0	0	1	0	0	1	1	1	1	0	1	0	0	1	1	1	0	0	1	0	0	1	0	1	2	3	3	0	1		
GPI-extensins	1	1	0	0	2	0	1	1	1	0	1	0	0	0	0	0	1	1	0	0	1	0	0	2	1	1	2	1	0	0	0	0	0	0	0	0	0	0	0	1	
Hybrid extensins	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	1	
Hybrid PRPs	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	
Shared	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	
< 15% motif	1	2	2	1	0	0	1	0	1	2	2	0	6	0	0	0	1	1	0	3	0	0	2	0	0	2	2	3	3	1	0	1	3	1	0	1	2	0	1		
	0	0	0	1	1	1	0	1	2	0	0	0	0	1	0	0	0	0	0	0	0	0	0	0	1	0	1	0	1	0	1	0	0	2	7	2	0	0	1	0	6
	1	1	1	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	1	0	0	0	0	1	0	0	0	0	0	0	0	0	0	0	0	0	0	0	1
	0	0	0	0	0	0	1	1	0	0	0	0	1	0	0	0	0	0	0	1	2	0	0	0	0	1	0	1	2	1	1	0	0	0	0	0	0	0	1	0	0
	0	0	0	0	0	0	0	0	0	0	0	1	0	0	0	0	0	0	0	0	1	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	2
	13	1	6	2	3	10	2	7	9	15	9	3	19	10	2	1	13	0	6	9	12	3	8	13	4	1	5	18	18	11	7	8	3	3	25	16	6	0	7		

Figure S16. Gene copy number analysis of hydroxyproline-rich glycoproteins (HRGPs). Columns represent species, while rows correspond to a given class of HRGP. Red and blue numbers indicate that a given species contains significantly more/less genes than others.

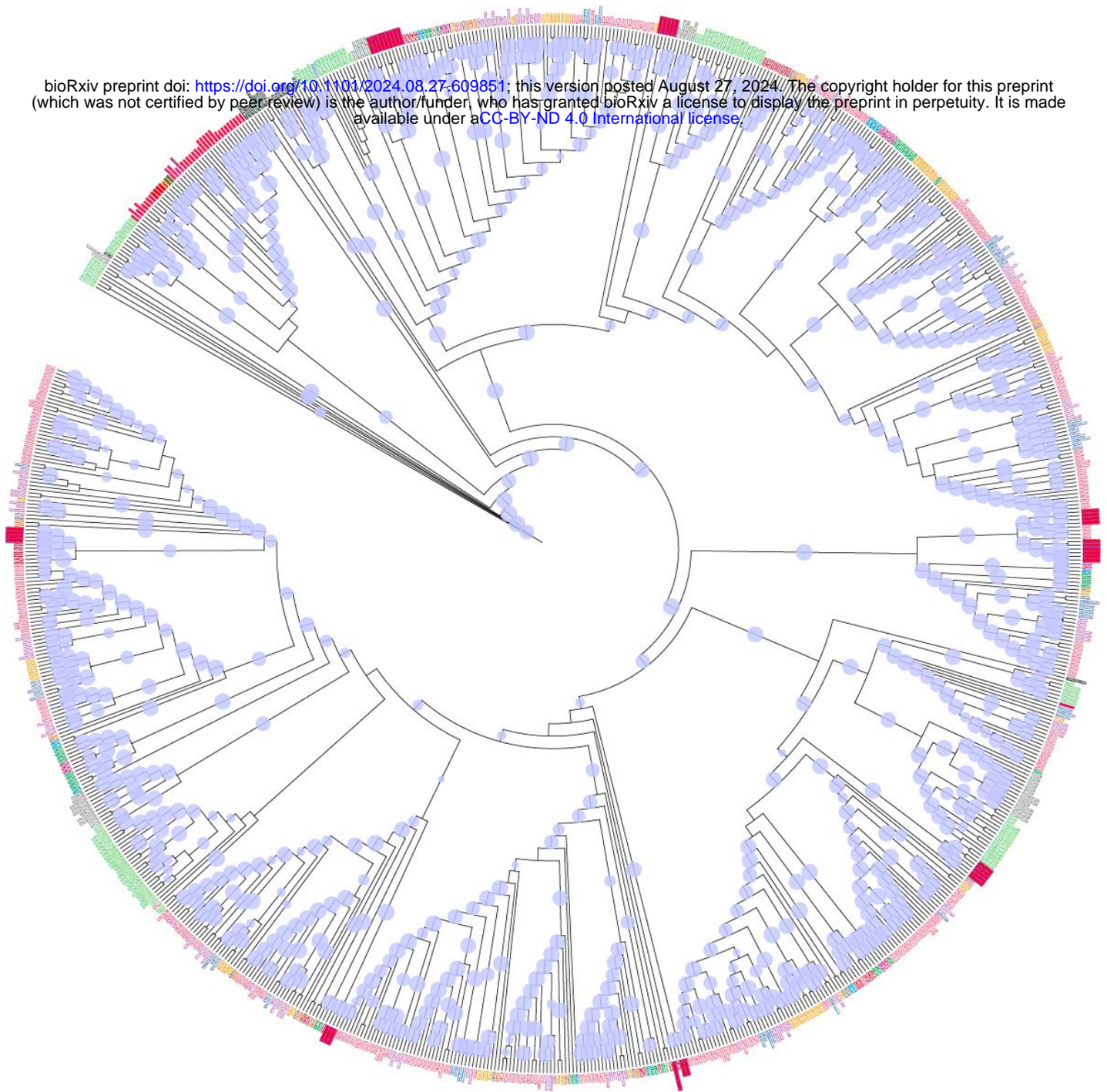


Figure S18. Phylogenetic analysis of CESA genes. The blue circles represent bootstrap values (value <50 are not indicated by a circle). The leaf colors represent the different species and orders.

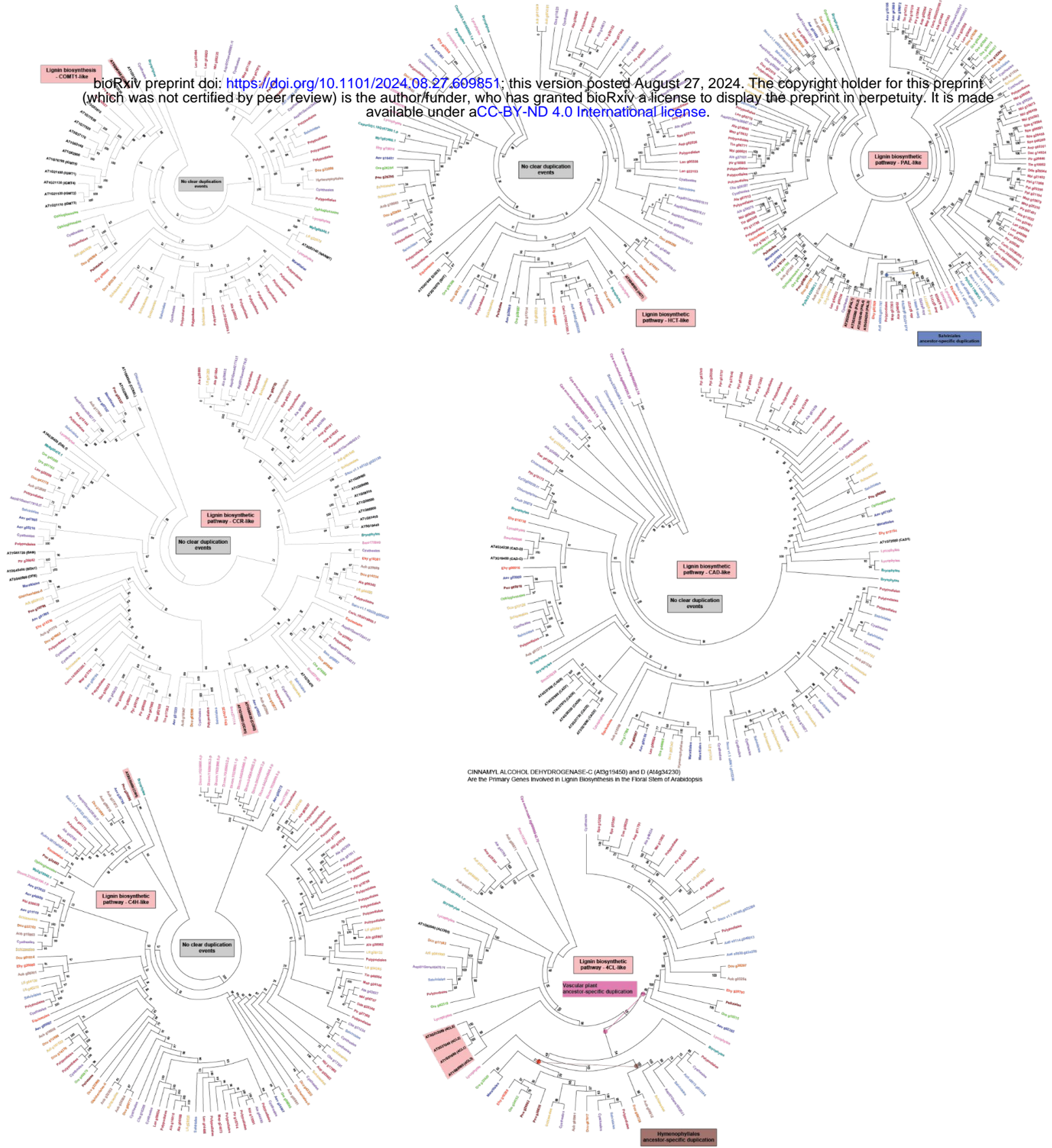


Figure S19. Phylogenetic analysis of lignin biosynthetic genes. Any inferred duplication events are indicated.