- 1 Full title
- 2 Transcriptome of the coralline alga *Calliarthron tuberculosum* (Corallinales, Rhodophyta) reveals
- 3 convergent evolution of a partial lignin biosynthesis pathway
- 4
- 5 Short title
- 6 *Calliarthron* transcriptomics reveals monolignol biosynthesis pathway
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18

20 Abstract

21

21	
22	The discovery of lignins in the coralline red alga Calliarthron tuberculosum raised new questions about
23	the deep evolution of lignin biosynthesis. Here we present the transcriptome of C. tuberculosum
24	supported with newly generated genomic data to identify gene candidates from the monolignol
25	biosynthetic pathway using a combination of sequence similarity-based methods. We identified
26	candidates in the monolignol biosynthesis pathway for the genes 4CL, CCR, CAD, CCoAOMT, and CSE
27	but did not identify candidates for PAL, CYP450 (F5H, C3H, C4H), HCT, and COMT. In gene tree
28	analysis, we present evidence that these gene candidates evolved independently from their land plant
29	counterparts, suggesting convergent evolution of a complex multistep lignin biosynthetic pathway in this
30	red algal lineage. Additionally, we provide tools to extract metabolic pathways and genes from the newly
31	generated transcriptomic and genomic datasets. Using these methods, we extracted genes related to
32	sucrose metabolism and calcification. Ultimately, this transcriptome will provide a foundation for further
33	genetic and experimental studies of calcifying red algae.
34	
35	Keywords: Red algae, lignification, calcification, transcriptome, gene identification, phenylpropanoid
36	pathway, monolignol
37	
38	Introduction
39	
40	Coralline red algae (Corallinales, Sporolithales, Hapalidiales) are a diverse lineage of calcified seaweeds
41	that play important ecological roles in nearshore ecosystems worldwide: they stabilize coral reefs by
42	creating a calcium carbonate matrix [1–3], induce settlement of invertebrate taxa [4–6], and contribute to

43 the storage of blue carbon through the creation of biogenic calcium carbonates [7,8]. In recent years, there

- 44 has been increased global attention paid to coralline algae. Taxonomists are clarifying their vastly
- 45 underestimated species diversity [9–12]; ecologists and physiologists are documenting interspecific

variation in coralline growth and calcification, particularly in response to climate stress, which may
ultimately impact marine communities [13–17]; evolutionary biologists are examining patterns in
coralline trait evolution [18–20] and using >100 million-year-old coralline fossils to strengthen modern
phylogenies [21,22].

50

51 The discovery of ligning within cell walls of the coralline species *Calliarthron cheilosporioides* 52 (Corallinales, Rhodophyta) dramatically changed our perspective on the evolution of lignin biosynthesis 53 [23]. Lignins are complex aromatic polymers predominantly found in the secondary cell walls of plant 54 support tissues [24,25] and were long considered to have evolved when land plants emerged from the 55 oceans, enabling upright growth in air [26]. Among the principal chemical components of wood, lignins 56 in plant secondary cell walls help reinforce tissue mechanical properties, permit hydraulic transport, and 57 increase pathogen resistance [27,28]. In the articulated coralline C. cheilosporioides, lignins were found 58 predominantly within decalcified flexible joints, called genicula [23], that have remarkable biomechanical 59 properties, permitting this articulated coralline species to thrive along wave-battered coastlines [29,30]. 60

61 Because lignin biosynthesis is physiologically complex and involves several enzymes in the monolignol 62 pathway [31–33], Martone et al. [23] proposed that much of the lignin biosynthetic pathway may have 63 predated land plants altogether, evolving in a common ancestor of red and green algae more than one 64 billion years ago. Alternatively, some (or all) of the monolignol biosynthetic pathway may have evolved 65 independently in the embryophyte and rhodophyte lineages. For example, one important enzyme involved 66 in S-lignin production (F5H) evolved independently in lycopods and embryophytes [34,35]. Moreover, 67 candidate genes related to monolignol biosynthesis have since been found in diverse algal lineages such 68 as diatoms, dinoflagellates, haptophytes, cryptophytes, and green and red algae [36], raising questions 69 about how the monolignol pathway may have evolved across such evolutionarily divergent lineages. Until 70 now, questions about monolignol evolution have largely gone unanswered as transcriptomic and genomic 71 data have mostly been limited to non-coralline red algae (e.g. [37–40] but see [41]).

73	Here we present a transcriptome of the articulated coralline Calliarthron tuberculosum (a sister species of
74	C. cheilosporioides) to investigate the evolutionary history of monolignol biosynthesis. Additionally,
75	though a complete mitochondrial genome [42] and a draft nuclear genome [43] of C. tuberculosum were
76	previously published, herein we generated a revised nuclear genome assembly using new short-read
77	sequence data to aid validation of transcriptomic reads. Based on comparative analysis of genome and
78	transcriptome data, we identify gene candidates for a putative monolignol biosynthetic pathway in C.
79	tuberculosum and investigate evolutionary relationships of these enzymes with those from other
80	taxonomic groups, including their land plant counterparts. We also provide a list of annotated genes in the
81	C. tuberculosum transcriptome and a simplified method for extracting genes from metabolic pathways.
82	We illustrate the utility of this dataset by extracting gene candidates involved in sucrose metabolism and
83	calcification. This transcriptomic dataset provides a foundation for future studies of coralline algal
84	ecology, physiology, and evolution.
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86	Results
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 86 87 88 89 90 91 92 93 94 95 96 	Results The C. tuberculosum transcriptome is complete and supported by genomic data Two transcriptomic datasets were generated from Calliarthron thalli: one from whole tissue (calcified intergenicula plus uncalcified genicula; sample I+G/PTM1 in the deposited data) and a second from intergenicular (i.e., calcified) tissue only (sample I/PTM2). Transcriptome sequencing based on RNA-Seq produced 38.8 total Gb of sequence data (17.3 Gb for sample I+G; 21.5 Gb for sample I). Reads were assembled <i>de novo</i> using Trinity. The whole tissue dataset had 172,700,376 total reads and the intergenicular tissue dataset had 215,491,160 total reads with an overall average coverage of 677-fold. A third reference transcriptome combining data from both tissues was assembled independently. All three

98	transcriptome data were considered complete based on the recovery of core eukaryotic genes (e.g. 94.5%
99	of CEGMA and 87.8% of BUSCO genes based on TBLASTN; Fig S1A). Genomic sequences were also
100	assembled for C. tuberculosum (Table S1), but these remain highly fragmented and were used only as
101	additional support to the transcriptome data in subsequent searches below. More than half (18840; 56.6%)
102	of the 33301 transcripts in the reference transcriptome were supported by the genome data (BLASTN, $E \le$
103	10-5).
104	
105	The incomplete monolignol biosynthetic pathway in Calliarthron tuberculosum
106	
107	The combined C. tuberculosum transcriptomic dataset was searched for genes encoding enzymes from the
108	monolignol biosynthetic pathway. The transcriptomic dataset was translated into all six reading frames
109	and queried with a combination of homology-based approaches, including HMMER searches and KEGG
110	based annotations. Closest homologs from <i>Arabidopsis thaliana</i> were also verified (BLASTN, $E \le 10^{-30}$).
111	We identified gene candidates of 4CL, CCR, CAD, CSE, and CCoAOMT, but not HCT, COMT, PAL,
112	TAL, or PTAL (Fig 1). PAL/TAL/PTAL was considered absent as only fragmented (and no full length)
113	sequences were identified. Evidence for the presence of homologous p450 enzymes (C3H, C3H, and
114	F5H) was weak; as a result, their status was classified as ambiguous (Fig 1). All sequences identified had
115	genomic support (BLASTN, $E \le 10^{-5}$) except for those identified for PAL/TAL/PTAL.
116	
117	Fig 1. The presence of <i>C. tuberculosum</i> sequence candidates in the monolignol pathway.
118	Red indicates presence of a putative homolog in C. tuberculosum; blue indicates no significant hits; green
119	indicates ambiguous presence. Note how the PTAL/PAL/TAL sequences obtained from the HMMER
120	search were indicated as absent as all sequences found were too short, 1/4-1/3 in length relative to those
121	in land plants. All sequences identified have genomic support except for PTAL/PAL/TAL.

123	Candidate sequences from <i>C. tuberculosum</i> (bolded as contig_gene_isoform in Figs 2, 3, and 4) were
124	characterized by comparing key residues with their land plant homologs in multiple sequence alignments.
125	The evolutionary relationships between the identified C. tuberculosum sequences, closely related
126	sequences in additional taxa, and sequences from the broader protein family of their land plant homologs
127	were analyzed in gene trees. Below we describe in detail results for the main biosynthetic enzymes 4CL,
128	CCR, and CAD (Figs 2, 3, and 4). Descriptions of the other biosynthetic enzymes CCoAMT, CSE, and
129	the cytochrome P450 sequences C3H, C4H, F5H are found in Appendix S1 and Figs S2-S4.
130	
131	Fig 2. 4CL candidates from <i>C. tuberculosum</i> in relation to plants and other taxa
132	(A) Partial alignment of C. tuberculosum candidates (bolded) and embryophyte 4CL sequences. Residues
133	involved in hydroxycinnamate binding are indicated with black triangles [61,62]. Phenylalanine substrate
134	binding pocket is indicated with Box I and Box II.
135	(B) Maximum likelihood acyl-activating enzyme (AAE) gene tree showing relationships between
136	Calliarthron sequences (magenta dots) and other taxa (Embryophyta – dark green, Chlorophyta – light
137	green, Rhodophyta – red, Animalia and Opisthokonta – purple, Bacteria and Cyanobacteria – blue,
138	Oomycota, Mycetozoa and Fungi – yellow, Ochrophyta – brown). Functionally demonstrated plant 4CLs
139	are labelled (+). Additional functional groups are labelled [44,45]. Ultrafast bootstrap values > 95 are
140	marked by *. Model = WAG+F+G4. Sites with \leq 80% occupancy were removed. Accession numbers can
141	be found in Appendix S1.
142	
143	Fig 3. CCR candidates from C. tuberculosum in relation to plants and other taxa

144 (A) Partial alignment *C. tuberculosum* candidates (bolded) and land plant CCR sequences. Catalytic

residues are labelled with NWYCY [64] and additional residues are indicated above with a black box.

146 NADPH binding pocket residues are indicated with black triangles [65] and the GXXGXX[A/G] motif is

147 underlined [66]. Hydroxycinnamonyl binding pocket residues are indicated with a gray triangle [65].

148	(B) CCR maximum likelihood gene tree showing relationships between <i>C. tuberculosum</i> (magenta dots)
149	and other taxa (Embryophyta – dark green, Chlorophyta – light green, Rhodophyta – red, Animalia and
150	Opisthokonta – purple, Bacteria and Cyanobacteria – blue, Oomycota, Mycetozoa and Fungi – yellow,
151	Ochrophyta – brown). Functionally demonstrated plant CCRs are labelled (+). Additional functional
152	groups are labelled. Ultrafast bootstrap values >95 are marked by *. Model = LG+G4. Sites with $\leq 80\%$
153	occupancy were removed. Accession numbers can be found in Appendix S1.
154	
155	Fig 4. CAD candidates from C. tuberculosum in relation to plants and other taxa
156	(A) Partial alignment of <i>C. tuberculosum</i> CAD sequence candidates (bolded) with land plant CAD
157	sequences. Zn ⁺² ion coordinating and proton shuttling residues are indicated with the black triangle,
158	NADPH or NADH interacting residues are boxed. Hydrostatic interaction forming residues are indicated
159	with a black box. Putative substrate-binding residues are indicated with grey boxes. [67-69]
160	(B) CAD maximum likelihood gene tree showing relationships between <i>C. tuberculosum</i> (magenta dots)
161	and other taxa (Embryophyta – dark green, Chlorophyta – light green, Rhodophyta – red, Animalia and
162	Opisthokonta – purple, Bacteria and Cyanobacteria – blue, Oomycota, Mycetozoa and Fungi – yellow,
163	Ochrophyta - brown). Alcohol dehydrogenase (ADH) sequences from yeast, and aldehyde reductase
164	(YAHK and AHR) sequences from <i>E. coli</i> were used as the ADH family is closely related to that of CAD
165	[70,71]. Functionally demonstrated plant CADs are labelled (+). Additional functional groups are
166	labelled. Ultrafast bootstrap values >95 are marked by *. Model = LG+G4. Sites with \leq 80% occupancy
167	were removed. Accession numbers can be found in Appendix S1.
168	

169 *Identification of 4CL candidates*

170

171 4CL is an acyl-CoA synthase in the monolignol pathway and a member of the acyl-activating enzyme

172 (AAE) superfamily. 4CL converts p-coumaric acid, caffeic acid, and ferulic acid into their respective

173 hydroxycinnamoyl-CoA thioesters. We identified 11 candidate 4CL-coding transcripts: two based on

174	KEGG analysis and nine additional sequences based on HMMER searches (Fig 2A). A query of these
175	sequences against the A. thaliana proteome returned related proteins within the acyl-activating enzyme
176	superfamily but not the A. thaliana 4CL (Table S2). Moderate sequence conservation exists in substrate
177	binding and hydroxycinnamate binding residues between 4CL candidates in C. tuberculosum (bolded)
178	and 4CLs in land plants (identity similarity [IS] > 70% Fig 2A).
179	
180	In the 4CL gene tree analysis, most C. tuberculosum sequences grouped with sequences from other
181	Rhodophytes (Fig 2B). In addition, C. tuberculosum sequences grouped within several functional clades
182	including malonate CoA ligase (ultrafast bootstrap support [BS] = 100%), succinylbenzoate CoA ligase
183	(BS = 87%), oxylate CoA ligase $(BS = 100%)$, acetyl CoA synthase $(BS = 100%)$, and the long chain
184	fatty acid CoA ligase (BS = 89%) (magenta dots, Fig 2B) [44,45]. In contrast, embryophyte 4CL
185	sequences form a clade separated from candidate 4CL sequences in C. tuberculosum (BS = 99% Fig 2B)
186	by the luciferase containing outgroup. Thus, 4CL candidates in C. tuberculosum did not show any clear
187	homology to functionally demonstrated 4CL sequences from embryophytes.
188	
189	Identification of CCR candidates
190	
191	CCR is the first committed enzyme in the monolignol pathway, reducing cinnamoyl-CoA esters to
192	cinnamaldehydes. We identified three sequences as candidate CCR-coding transcripts: one based on
193	KEGG analysis and two additional sequences based on HMMER searches (Fig 3A). A query of these
194	sequences against the A. thaliana proteome returned sequences within the CCR family (CCR7, CCR4,
195	CCR-Like6) (Table S2). Substrate-binding residues (NWYCY) and the hydroxycinnamonyl-binding
196	pocket showed low sequence conservation (IS <80%). In contrast, the core catalytic residues (S, T, and
197	K) and NADPH-binding residues appear to be conserved (IS >90%) between the candidate sequences in
198	C. tuberculosum and CCRs in land plants (Fig 3A).

200 In the CCR gene tree analysis, C. tuberculosum sequences varied in their relatedness to other taxa with 201 some sequences closer to Rhodophytes and others more closely related to Oomycota/Mycetozoa/Fungi 202 (Fig 3B). Additionally, CCR candidates in C. tuberculosum were mapped with epimerase dehydratase 203 type sequences that included the A. thaliana CCR family (Fig 3B). Sequences from C. tuberculosum 204 grouped with epimerase dehydratase type sequences of non-embryophyte origin. In contrast, embryophyte 205 CCR, class 2 CCR, and CCR-like form an independent clade (BS >97%). The embryophyte CCR clade 206 and the non-embryophyte epimerase dehydratase clade (containing sequences from C. tuberculosum) 207 were more closely related than the embryophyte dihydroflavonol-4-reductase protein (DFR) group within 208 the overall epimerase dehydratase family. 209 210 Identification of CAD candidates 211 212 CAD, the final step in the monolignol pathway, is an alcohol dehydrogenase converting various 213 hydroxycinnamaldehydes to their respective hydroxycinnamyl alcohols. SAD, proposed to catalyze this 214 same reaction for sinapyl monolignols [46], is added into our analysis despite debate over their function. 215 We identified five sequences as candidate CAD-encoding transcripts: two based on KEGG analysis and 216 three additional sequences based on HMMER searches (Fig 4A). A query of these sequences against the 217 A. thaliana proteome returned CAD2 and other alcohol dehydrogenases (Table S2). NADPH-binding 218 motifs show moderate conservation (IS >80%) (Fig 4A). One C. tuberculosum sequence showed high 219 conservation with land plant counterparts, suggesting a promising CAD candidate (+ in Figs 3A and 3B). 220 221 In the CAD gene tree analysis, all *C. tuberculosum* sequences grouped with sequences from other 222 Rhodophytes (Fig 4B). CAD candidates in C. tuberculosum were mapped with their embryophyte CAD 223 counterparts and closely related alcohol dehydrogenases. Sequences from C. tuberculosum grouped 224 together with oxidoreductases (BS = 100%), sorbitol dehydrogenases (BS = 100%), general alcohol 225 dehydrogenases (BS = 100%), and an algal CAD clade (BS = 100%). Sequences in this algal CAD clade

226	were based on previous sequence similarity-based annotation and have not been functionally
227	demonstrated. In contrast, the land plant CAD and SAD sequences form their own clades (BS 100%; Fig
228	4B) that are separated from the <i>C. tuberculosum</i> candidates by the functionally distinct alcohol
229	dehydrogenases, such as yeast alcohol dehydrogenase 7 (ADH7) and E. coli aldehyde reductase (YAHK).
230	
231	Identification of additional metabolic pathways in Calliarthron tuberculosum
232	
233	To enable broad and rapid identification of C. tuberculosum genes involved in specific metabolic
234	processes, we present two general tools for gene identification within the C. tuberculosum transcriptome
235	dataset using KEGG based annotations. This involves extracting whole metabolic pathways or individual
236	genes (see Appendix S1; Fig S5). We included annotations for all metabolic genes recovered in the C.
237	tuberculosum transcriptome (Table S3). We identified 36 putative C. tuberculosum genes present in the
238	starch and sucrose metabolism pathway (Fig S5; Table S4). In addition, we individually searched for
239	genes potentially involved in calcification [41,47,48] and identified 13 sequence candidates related to
240	calcium transport, six related to inorganic carbon transport, five related to pH homeostasis, 19 putative
241	carbonic anhydrases, and 12 putative HSP90 genes (Table S5).
242	
243	Discussion
244	
245	Evidence for convergent evolution of monolignol biosynthesis
246	
247	Using sequence similarity methods with genes from the monolignol pathway in land plants, we identified
248	candidates for five genes related to monolignol biosynthesis (4CL, CCR, CAD, CCoAOMT, and CSE)
249	from the newly generated C. tuberculosum transcriptomic dataset. These gene candidates are supported
250	by genomic evidence, retain major motifs from their respective gene family, and return their A. thaliana

251 counterpart in reciprocal BLAST analyses, suggesting that these enzymes may function similarly in 252 monolignol biosynthesis in C. tuberculosum.

253

254 Despite supporting evidence from sequence similarity analyses, functional predictions for candidate 255 sequences in the monolignol pathway within C. tuberculosum are obscured by the gene tree analysis. If 256 the monolignol pathway in embryophytes and C. tuberculosum evolved in a common ancestor and was 257 retained through conserved evolution, we would expect their sequences to form functional clades 258 uninterrupted by functionally divergent protein sequences. However, with the exception of the 259 CCoAOMT candidate, our gene tree analyses consistently showed that monolignol biosynthetic genes in 260 land plants are not sister to those in C. tuberculosum. C. tuberculosum sequences were found within each 261 respective overall protein family, but consistently grouped with land plant genes of non-monolignol 262 forming function. If these C. tuberculosum sequences are functionally homologous to the monolignol 263 biosynthesis counterpart in land plants, then they likely arose independently in *C. tuberculosum*. 264 Convergent evolution in protein function, with phylogenetic patterns of protein sequences with similar 265 functions intersected by sequences with dissimilar functions, is not uncommon in cell wall synthesizing 266 enzymes [49]. Biosynthetic enzymes in C. tuberculosum could have evolved similar substrate specificity 267 after the divergence of red algae and land plants or, alternatively, may reflect genes that were individually 268 acquired. Previous evidence suggests that the core monolignol biosynthesis genes (4CL, CCR, and CAD) 269 in C. tuberculosum may have been acquired through horizontal gene transfer from a bacterial source [36]. 270 Thus, over evolutionary time genes in C. tuberculosum may have developed enough synchronicity in gene 271 expression and protein regulation to produce an ad hoc monolignol biosynthetic pathway.

272

273 Alternatively, the phylogenetic evidence might suggest that gene candidates in *C. tuberculosum* do not

274 function in monolignol biosynthesis and instead have a function similar to their sister sequences within

- 275 their distinct phylogenetic groupings. For example, considering only clustering patterns in the
- 276 phylogenetic data, perhaps C. tuberculosum contig 141618 functions as a CoA ligase that acts on

277	malonate and not coumarate (4CL enzyme) (Fig 2B). However, the tandem use of stricter curated
278	sequences in our predictive HMM models and more flexible HMM models with previously annotated
279	data, such as KEGG annotations, improves our confidence in finding potential gene candidates.
280	Biochemical or functional assays will ultimately be needed to verify the function of candidate gene
281	sequences.
282	
283	The monolignol biosynthesis pathway and missing steps in Calliarthron tuberculosum
284	
285	Several key steps in the monolignol biosynthetic pathway were not recovered in the C. tuberculosum
286	transcriptome, including PAL, TAL, PTAL, HCT, COMT, C3H, C4H, or F5H. Although we cannot
287	dismiss that these observations may be due to fragmented sequences in the assembled genome and
288	transcriptome data, we present several other possibilities.
289	
290	The ammonia-lyase PAL, TAL, or PTAL creates the first substrates in the monolignol biosynthetic
291	pathway [50-52]. Although no full-length homologs were identified in the C. tuberculosum
292	transcriptome, short sequence candidates identified may represent a fragmented gene. However, these
293	short sequences lacked genomic support, indicating they may be contaminants of non-Calliarthron origin.
294	For this reason, PAL, TAL, and PTAL are currently indicated as absent (Fig 1). If these are indeed from
295	C. tuberculosum, RACE amplification could help determine if the short ammonia-lyase we identified has
296	a longer transcript. C. tuberculosum likely has an ammonia-lyase acting on phenylalanine or tyrosine
297	since PAL and TAL are also key enzymes in producing flavanoids and coumarins, which have been
298	previously detected in both fleshy and coralline red algae [53]. Further validation will be required to
299	elucidate their presence.
300	
301	C3H, C4H, or F5H are p450 monooxygenases responsible for converting substrates across the monolignol

302 pathway eventually resulting in H to S to G type monolignols, respectively (Fig 1). P450 sequence

303 candidates have been identified, but their substrate-specific identity as C3H, C4H, or F5H homologs is 304 unclear. The cytochrome P450 sequence candidates from the C. tuberculosum transcriptome form two 305 divergent groups. One group is likely involved in carotenoid biosynthesis, positioned within the CYP97 306 clade, while the other group forms their own clade of unknown function (Fig S2B). The identified 307 candidates from C. tuberculosum may have multi-substrate specificities, acting on various substrates, 308 including monolignol intermediate products. Some substrate promiscuity has previously been observed 309 within members of the cytochrome P450 enzyme family [54,55]. Alternatively, each of the identified 310 P450 clades in C. tuberculosum could contain a new class of cytochrome P450 capable of functioning in 311 H-, G-, or S- unit monolignol biosynthesis. This proposed convergent evolution of a distinct and 312 independently-evolved cytochrome P450 involved in monolignol production has previously been 313 documented in the clubmoss Selaginella moellendorffii (F5H) [34,35]. In any case, the presence of unique 314 P450s represents an interesting avenue of exploration to elucidate substrate specificity and functionality 315 in the monolignol pathway in C. tuberculosum. 316

317 HCT is one alternative route shifting monolignol synthesis from H- to G- to S- types using a temporary 318 shikimate decoration (Fig 1) [56]. Its absence could suggest that C. tuberculosum does not utilize an HCT 319 enzyme or create G lignin using this route. Another alternative route in G- and S- type monolignol 320 synthesis utilizes a CSE enzyme that acts on caffeoyl shikimate, an HCT downstream product (Fig 1). 321 The absence of an HCT is at odds with the CSE enzyme identified in this study (Fig 1), suggesting that 322 the CSE candidate identified may not be utilized in the monolignol biosynthetic pathway for C. 323 tuberculosum. Though this absence could be due to fragmentation in the transcriptome, more data are 324 required for further validation.

325

326 COMT is necessary for S type monolignol production in angiosperms [57–59]. The absence of this
 327 enzyme raises questions about how *C. tuberculosum* can produce sinapyl alcohol, a precursor component
 328 for S monolignols. Some evidence exists for a bifunctional enzyme in pine that can function as both

329	COMT and CCoAOMT (named AEOMT) in heterologous systems [60]. However, only moderate-to-low
330	sequence similarity is shared among CCoAOMT, COMT, and the bifunctional AEOMT. Perhaps a
331	similar protein with broad substrate specificity is present in C. tuberculosum but has yet to be identified
332	based on sequence similarity.
333	
334	Conclusion
335	
336	In summary, we have identified several gene candidates in the C. tuberculosum transcriptome that
337	represent central components in the monolignol biosynthetic pathway, helping to explain the surprising
338	presence of lignins in this coralline red alga. Despite the complexity of monolignol biosynthesis, and
339	contrary to the predictions outlined in Martone et al. [23], our gene trees do not demonstrate a deeply
340	conserved evolution of monolignol biosynthesis, but instead suggest that each of the enzymes identified
341	in C. tuberculosum likely evolved independently from those found in land plants. Interestingly, there
342	remain several key enzymes in the monolignol pathway whose sequences have not been identified,
343	including those related to pathway entry and to shifting the types of monolignols produced that would
344	form H-, G-, and S-lignins within the cell wall. Further biochemical evidence and validation of sequence
345	expression will be necessary to provide functional support for both the genes identified and to elucidate
346	potential alternative routes in the monolignol biosynthetic pathway in C. tuberculosum. By providing
347	methods to easily identify additional gene candidates from the C. tuberculosum transcriptome, we aim to
348	facilitate future research on this fascinating organism.
349	
350	Methods
351	
352	Data and code availability
353	

354	All sequencing data generated from this study are available at European Nucleotide Archive
355	(transcriptome data: accession PRJEB39919; genome data: accession PRJEB39919). Genome supported
356	transcripts, transcriptome assemblies, annotations, and an example of metabolic pathway extraction are
357	available on Github (<u>https://github.com/martonelab/geneAnnotCalliarthronTranscriptome/</u>).
358	
359	Experimental model and subject details
360	
361	Specimen collection and sequencing
362	
363	Two male, haploid specimens of Calliarthron tuberculosum were collected October 6, 2013, from
364	Bluestone Point (48.81952, -125.1640), Bamfield, British Columbia, Canada and verified as haploid male
365	specimens by microscopy. A portion of each collected sample was pressed and deposited into the UBC
366	herbarium with voucher codes A89970 and A89985. Voucher codes can be queried at
367	https://herbweb.botany.ubc.ca/herbarium/search.php?Database=algae for more information.
368	Calcified intergenicula and non-calcified genicula from each individual were divided into two portions for
369	data collection: either whole tissue (Sample I+G/PTM1 in the dataset) or calcified tissue only (Sample
370	I/PTM2 in the dataset). Total RNA was extracted using the Spectrum Plant Total RNA kit (Cat #
371	STRN50, Sigma-Aldrich) and sequenced on the Illumina HiSeq 2000 platform (paired-end 2x100bp,
372	insert size ~220bp).
373	
374	Abbreviation of enzyme names
375	
376	CAD, (hydroxy)cinnamyl alcohol dehydrogenase; SAD, sinapyl alcohol dehydrogenase; CCoAOMT,
377	caffeoyl-CoA O-methyl transferase; CCR, (hydroxy)cinnamoyl-CoA reductase; C3'H, p-coumaroyl
378	shikimate 3'-hydroxylase; C4H, cinnamate 4 hydroxylase; 4CL, 4-hydroxycinnamoyl-CoA ligase;
379	COMT, caffeic acid O-methyltransferase; F5H, ferulic acid/coniferaldehyde/coniferyl alcohol 5-

380 hydroxylase; HCT, hydroxycinnamoyl-CoA:shikimate hydroxycinnamoyl transferase; PAL,

- 381 phenylalanine ammonia-lyase
- 382
- **383** *Transcriptome assembly and annotation*
- 384

385 Illumina sequence reads were assembled using Trinity with the *de novo* mode at default setting [72]. 386 independently for each anatomical sample (I+G/PTM1; I/PTM2 in the ENA database). A reference 387 transcriptome was also assembled *de novo* using Trinity by independently combining the sequence reads 388 generated from both samples. The assembled transcripts were annotated using Blast2GO [73]. Briefly, 389 each transcript was searched against the NCBI RefSeq protein database (BLASTX, $E \le 10^{-5}$), and its 390 putative function was inferred based on the top protein hit and Gene Ontology (GO) terms. These proteins 391 were then mapped onto the corresponding metabolic pathways in the Kyoto Encyclopaedia of Gene and 392 Genomes (KEGG) database [74]. Identification of genes present in KEGG annotated pathways were 393 extracted using the pathview package [75].

394

395 Filtering contaminant sequences in genome assembled data

396

397 To identify putative contaminant sequences in the genome assembly, each genome scaffold was searched 398 (BLASTN) against a database of archaeal, bacterial and viral genome sequences retrieved from the NCBI 399 RefSeq database. Sequences with a significant hit ($E \le 10^{-5}$, covering > 50% of the query length) were 400 considered putative contaminants and removed from the genome assembly. To identify broad differences 401 in sequence characteristics, genomic scaffolds with and without transcriptomic support were compared 402 for G+C content and transcript length (Fig S1B). Scaffolds with no transcript support and low recovery of 403 eukaryotic genes (< 6% BUSCO or CEGMA recovery) were also identified as likely putative 404 contaminants and removed from the genome assembly.

406 *Genome annotation guided by transcriptome evidence*

407

407	
408	Repetitive elements in the genome assembly were identified and masked using RepeatMasker version
409	open-4.0.6 [76]. To maximize recovery of transcript support for genome scaffolds, the transcriptomes
410	(I+G/PTM1; I/PTM2 in the dataset) were mapped against the masked genome scaffolds using PASA
411	v2.0.2 [77], and full-length coding sequences (CDSs) were predicted with TransDecoder v5.0.1 [72].
412	These CDSs represent the primary set of putative genes and were used as extrinsic hints to guide <i>ab initio</i>
413	gene prediction using AUGUSTUS v3.2.1 [78] from the genome scaffolds.
414	
415	HMM based gene candidate search
416	
417	Monolignol biosynthesis gene candidates were identified from the C. tuberculosum transcriptomic dataset
418	using Hidden Markov Model (HMM) based searches [79]. Transcriptomic sequence contigs were
419	translated into all six reading frames using EMBOSS Transeq [80]. This amino acid database was used
420	for subsequent sequence searches. HMM profiles used to search for homologs in the transcriptome were
421	produced by aligning amino acid sequences of a given protein or protein family using MUSCLE [81] with
422	no manual adjustment. The profiles were searched against the translated C. tuberculosum dataset in
423	HMMER searches [79] to look for putative sequence homologs. Sequences more than 100 amino acids
424	long were retained for subsequent analysis. These sequences were then searched against the Arabidopsis
425	(GenBank taxid:3701) proteome using NCBI's BLAST [82] to verify their closest homolog match
426	(BLASTP, $E \le 10^{-30}$).
427	
428	Domain and motif comparison
429	
430	The monolignol biosynthetic genes and their overall gene families contain sequence domains that

431 influence protein shape and function. To compare these key domains, multiple sequence alignments

432	(MSA) of candidate amino acid sequences from C. tuberculosum with their land plant counterpart protein
433	were produced. Sequences were aligned using MUSCLE under default settings [81]. Key domains and
434	motifs were chosen based on available literature and highlighted in the MSA as indicated in each figure
435	legend. In each MSA, an asterisk (*) represents full conservation; and a period (.) represents sites with
436	conservation >50%. Accession numbers can be found in Appendix S1.
437	
438	Gene tree analysis
439	
440	Gene trees were reconstructed for the candidate sequences of C. tuberculosum identified. For each gene
441	tree analysis, sequence candidates from C. tuberculosum, the functionally demonstrated enzyme sequence
442	from land plants, enzyme sequences from the overall protein family from land plants, and the top 20
443	sequences identified by NCBI BLAST using C. tuberculosum candidates as a query against the total
444	database using default settings (BLASTP, $E \le 10^{-20}$) were compiled. Land plant sequences identified to
445	represent the functional gene and overall gene family were curated by a literature search. For each set of
446	sequences, a multiple sequence alignment was performed using MUSCLE with default setting [81]. Sites
447	with <80% coverage were removed using trimAl [83]. IQTree was used to search for the evolutionary
448	model alignment under a BIC criterion [84,85]. A maximum likelihood tree was reconstructed using
449	IQTree [86], with node support calculated based on 1000 ultrafast bootstrap pseudoreplicates in IQTree
450	[86]. A clade is considered strongly supported when bootstrap value $\ge 95\%$. FigTree was used to edit
451	branch width and colors [87]. Accession numbers can be found in Appendix S1.
452	
453	Generation of genome data as additional support for transcriptome data
454	
455	Genome data of <i>C. tuberculosum</i> were generated using Illumina IIx platform (paired-end 2×150bp reads,
456	insert size ~350 bp). An overview of the summary statistics for the genome assembly can be found in

457 Table S1. Adapter sequences were removed using Trimmomatic v0.33 [88] (LEADING:25

458 TRAILING:25 HEADCROP:10 SLIDINGWINDOW:4:20 MINLEN:50). The generated filtered 459 sequence reads and the previously published genome data (GenBank accession #: SRP005182) generated 460 using the 454 pyrosequencing platform [43] were used in a *de novo* genome assembly using SPAdes [89]. 461 The 454 reads were treated as unpaired, single-end reads in the assembly process. This *de novo* assembly 462 was further scaffolded with the transcriptome data using the L RNA Scaffolder [90]. Putative 463 contaminant sequences were removed based on shared similarity against known genome sequences from 464 bacterial, archaeal, and viral sources in NCBI RefSeq (BLASTN, $E \le 10^{-5}$), and subsequently based on 465 discrepancy in G+C content of the assembled scaffolds, and the recovery of core eukaryotic genes 466 (CEGMA and BUSCO). Because the genome assembly is fragmented, genome scaffolds on which no 467 transcripts were mapped were filtered out, yielding the final genome assembly (21,672 scaffolds, total 468 bases 64.15 Mbp). These genome scaffolds were used as additional support for the transcriptome data. 469 For the reference transcriptome (combined I+G/PTM1 ; I/PTM2), putative coding sequences were 470 predicted based on alignment of the assembled transcripts against the genome scaffolds using PASA [77] 471 and TransDecoder [72], from which the coded protein sequences were predicted. 472 473 Completeness of transcriptome and genome data 474 475 The completeness of the genome and transcriptome data was assessed by the recovery of core conserved 476 eukaryote genes with the Core Eukaryotic Genes Mapping Approach (CEGMA) [91] and Benchmarking 477 Universal Single-Copy Orthologs (BUSCO) [92] datasets. CEGMA and BUSCO datasets (eukaryote 478 odb9 and Viridiplantae odb10) were independently used as query to search against the predicted proteins 479 from the reference transcriptome (combined IG and IO) using BLASTP ($E \le 10^{-5}$) and against the same 480 transcriptome using TBLASTN ($E \le 10^{-5}$). The core CEGMA and BUSCO proteins were also queried 481 against the 21,672 genome scaffolds using TBLASTN ($E \le 10^{-5}$). 482

484 Key Resources Table

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Biological Samples		
Calliarthron tuberculosum	This paper	A89970 and A89985
(sample vouchers)		at
		https://herbweb.bota
		ny.ubc.ca/herbarium/
		search.php?Database
		<u>=algae</u>
Critical Commercial Assays	_	
HiSeq 2000 (Transcript reads)	Illumina	
IIx platform (Genomic reads)	Illumina	
Spectrum Plant Total RNA kit	Sigma-Aldrich	STRN50
Deposited Data		
Raw sequencing reads for transcriptomic and genomic	This paper	PRJEB39919
data		
Genome supported transcripts, transcriptome	This paper	https://github.com/m
assemblies, annotations		artonelab/geneAnnot
		<u>CalliarthronTranscri</u>
		ptome/
Additional Calliarthron Genomic Reads	[43]	SRP005182
Pyropia genomic data	[39]	MXAK00000000
Arabidopsis Proteome		taxid:3701

Software and Algorithms		
TransDecoder v5.0.1	[72]	https://github.com/T
		ransDecoder/TransD
		ecoder/wiki
Trinity	[72]	https://github.com/tri
		nityrnaseq/trinityrna
		<u>seq/wiki</u>
Blast2GO	[73]	https://www.blast2g
		<u>o.com/</u>
Kyoto Encyclopedia of Genes and Genomes (KEGG)	[74]	https://www.genome
		.jp/kegg/
Pathview R Package	[75]	https://www.biocond
		uctor.org/packages/r
		elease/bioc/html/pat
		hview.html
HMMER	[79]	http://hmmer.org/
EMBOSS Transeq	[80]	http://emboss.source
		forge.net/apps/releas
		e/6.6/emboss/apps/tr
		anseq.html
MUSCLE v3.5	[81]	http://www.drive5.c
		om/muscle/muscle.h
		tml
IQtree	[85,86]	http://www.iqtree.or
		<u>g/</u>

TrimAl	[83]	http://trimal.cgenomi
		<u>cs.org/</u>
FigTree	[87]	http://tree.bio.ed.ac.u
		k/software/figtree/
Trimmomatic v0.33	[88]	http://www.usadella
		b.org/cms/?page=tri
		mmomatic
SPAdes	[89]	https://cab.spbu.ru/s
		oftware/spades/
L_RNA_Scaffolder	[90]	https://github.com/C
		<u>AFS-</u>
		bioinformatics/L_R
		NA_scaffolder
PASA v2.0.2	[77]	https://github.com/P
		ASApipeline/PASA
		pipeline
CEGMA	[91]	http://korflab.ucdavi
		s.edu/datasets/cegma
		<u> </u>
BUSCO	[92]	https://busco.ezlab.o
		<u>rg/</u>
RepeatMasker version open-4.0.6	[76]	http://www.repeatma
		sker.org/
AUGUSTUS v3.2.1	[78]	https://github.com/n
		extgenusfs/augustus

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487

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500

501 Author Contributions

502

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504 J.X., K.H., M.A.L., C.X.C.; Visualization, J.X., E.J.; Writing - Original Draft, J.X. and P.T.M.; Review

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506

507 Declaration of Interests

508

509 The authors declare no competing interests.

510

511 References

513	1.	Adey WH. The algal ridges and coral reefs of St. Croix: their structure and Holocene
514		development. Atoll Research Bulletin. 1975; 1-67. doi:https://doi.org/10.5479/si.00775630.187.1
515	2.	Borowitzka MA. Algal calcification. Oceanography and Marine Biology Annual Review. 1977;
516		189–223.
517	3.	Goreau TF. Calcium carbonate deposition by coralline algae and corals in relation to their roles as
518		reef-builders. Annals of the New York Academy of Sciences. 1963;109: 127-167.
519	4.	Harrington L, Fabricius K, De'ath G, Negri A. Recognition and selection of settlement substrata
520		determine post-settlement survival in corals. Ecology. 2004;85: 3428-3437. doi:10.1890/04-0298
521	5.	O'Leary JK, Barry JP, Gabrielson PW, Rogers-Bennett L, Potts DC, Palumbi SR, et al. Calcifying
522		algae maintain settlement cues to larval abalone following algal exposure to extreme ocean
523		acidification. Scientific reports. 2017;7: 5710-5774. doi:10.1038/s41598-017-05502-x
524	6.	Swanson RL, de Nys R, Huggett MJ, Green JK, Steinberg PD. In situ quantification of a natural
525		settlement cue and recruitment of the Australian sea urchin Holopneustes purpurascens. Marine
526		ecology Progress series (Halstenbek). 2006;314: 1-14. doi:10.3354/meps314001
527	7.	Fisher K, Martone PT. Field study of growth and calcification rates of three species of articulated
528		coralline algae in British Columbia, Canada. Biological Bulletin. 2014;226: 121-130.
529		doi:10.1086/BBLv226n2p121
530	8.	van der Heijden LH, Kamenos NA. Reviews and syntheses: Calculating the global contribution of
531		coralline algae to total carbon burial. Biogeosciences. 2015;12: 6429-6441. doi:10.5194/bg-12-
532		6429-2015
533	9.	Gabrielson PW, Hughey JR, Diaz-Pulido G. Genomics reveals abundant speciation in the coral
534		reef building alga Porolithon onkodes (Corallinales, Rhodophyta). Journal of phycology. 2018;54:
535		429–434. doi:10.1111/jpy.12761
536	10.	Hind KR, Miller KA, Young M, Jensen C, Gabrielson PW, Martone PT. Resolving cryptic species
537		of Bossiella (Corallinales, Rhodophyta) using contemporary and historical DNA. American

538 journal of botany. 2015;102: 1912–1930. doi:10.3732/ajb.1500308

- 539 11. Hind KR, Gabrielson PW, Lindstrom SC, Martone PT. Misleading morphologies and the
- 540 importance of sequencing type specimens for resolving coralline taxonomy (Corallinales,
- 541 Rhodophyta): Pachyarthron cretaceum is Corallina officinalis. Journal of Phycology. 2014;50:
- 542 760–764. doi:10.1111/jpy.12205
- 543 12. Twist BA, Neill KF, Bilewitch J, Jeong SY, Sutherland JE, Nelson WA. High diversity of
- 544 coralline algae in New Zealand revealed: Knowledge gaps and implications for future research.
- 545 PloS one. 2019;14: e0225645. doi:10.1371/journal.pone.0225645
- 546 13. Bergstrom E, Ordoñez A, Ho M, Hurd C, Fry B, Diaz-Pulido G. Inorganic carbon uptake
- 547 strategies in coralline algae: Plasticity across evolutionary lineages under ocean acidification and
- 548 warming. Marine environmental research. 2020;161: 105–107.
- 549 doi:10.1016/j.marenvres.2020.105107
- 550 14. Cornwall CE, Comeau S, McCulloch MT. Coralline algae elevate pH at the site of calcification
- under ocean acidification. Global change biology. 2017;23: 4245–4256. doi:10.1111/gcb.13673
- 552 15. Guenther R. The effect of temperature and pH on the growth and biomechanics of coralline algae.
 553 University of British Columbia. 2016.
- McCoy SJ, Ragazzola F. Skeletal trade-offs in coralline algae in response to ocean acidification.
 Nature climate change. 2014;4: 719–723. doi:10.1038/nclimate2273
- 556 17. Noisette F, Egilsdottir H, Davoult D, Martin S. Physiological responses of three temperate
- 557 coralline algae from contrasting habitats to near-future ocean acidification. Journal of
- experimental marine biology and ecology. 2013;448: 179–187. doi:10.1016/j.jembe.2013.07.006
- 18. Hind KR, Gabrielson PW, Jensen C, Martone PT. Evolutionary reversals in Bossiella
- 560 (Corallinales, Rhodophyta): first report of a coralline genus with both geniculate and
- 561 nongeniculate species. Journal of phycology. 2018;54: 788–798. doi:10.1111/jpy.12788
- 562 19. Janot K, Martone PT. Convergence of joint mechanics in independently evolving, articulated
- 563 coralline algae. Journal of experimental biology. 2016;219: 383–391. doi:10.1242/jeb.131755

56420.Steneck RS. The ecology of coralline algal crusts: convergent patterns and adaptive strategies.

565 Ann Rev Ecol Syst. 1986;17: 273–303.

- 566 21. Aguirre J, Perfectti F, Braga JC. Integrating phylogeny, molecular clocks, and the fossil record in
- the evolution of coralline algae (Corallinales and Sporolithales, Rhodophyta) Author (s): Julio
- 568 Aguirre, Francisco Perfectti and Juan C. Braga Published by: Cambridge University P.
- 569 Paleobiology. 2010;36: 519–533.
- 570 22. Rösler A, Perfectti F, Peña V, Aguirre J, Braga JC, Gabrielson P. Timing of the evolutionary
- 571 history of Corallinaceae (Corallinales, Rhodophyta). Journal of Phycology. 2017;53: 567–576.

572 doi:10.1111/jpy.12520

- 573 23. Martone PT, Estevez JM, Lu F, Ruel K, Denny MW, Somerville C, et al. Discovery of Lignin in
- 574 Seaweed Reveals Convergent Evolution of Cell-Wall Architecture. Current Biology. 2009;19:
- 575 169–175. doi:10.1016/j.cub.2008.12.031
- 576 24. Boerjan W, Ralph J, Baucher M. Lignin Biosynthesis. Annual Review of Plant Biology. 2003;54:
 577 519–546. doi:10.1146/annurev.arplant.54.031902.134938
- 57825.Mottiar Y, Vanholme R, Boerjan W, Ralph J, Mansfield SD. Designer lignins: Harnessing the
- 579 plasticity of lignification. Current Opinion in Biotechnology. 2016;37: 190–200.
- 580 doi:10.1016/j.copbio.2015.10.009
- 581 26. Vanholme R, Demedts B, Morreel K, Ralph J, Boerjan W. Lignin biosynthesis and structure. Plant
 582 Physiology. 2010;153: 895–905. doi:10.1104/pp.110.155119
- 583 27. Lange BM, Lapierre C, Sandermann H. Elicitor-induced spruce stress lignin: Structural similarity
 584 to early developmental lignins. Plant Physiology. 1995;108: 1277–1287.
- 585 doi:10.1104/pp.108.3.1277
- 586 28. Tronchet M, BalaguÉ C, Kroj T, Jouanin L, Roby D. Cinnamyl alcohol dehydrogenases-C and D,
- 587 key enzymes in lignin biosynthesis, play an essential role in disease resistance in Arabidopsis.
- 588 Molecular Plant Pathology. 2010;11: 83–92. doi:10.1111/j.1364-3703.2009.00578.x
- 589 29. Martone PT. Kelp versus coralline: Cellular basis for mechanical strength in the wave-swept

- 590 seaweed Calliarthron (Corallinaceae, Rhodophyta). Journal of Phycology. 2007;43: 882–891.
- **591** doi:10.1111/j.1529-8817.2007.00397.x
- 592 30. Denny MW, King FA. The extraordinary joint material of an articulated coralline alga. II.
- 593 Modeling the structural basis of its mechanical properties. Journal of Experimental Biology.
- **594** 2016;219: 1843–1850. doi:10.1242/jeb.138867
- 595 31. Weng JK, Chapple C. The origin and evolution of lignin biosynthesis. New Phytologist. 2010;187:
- **596** 273–285. doi:10.1111/j.1469-8137.2010.03327.x
- 597 32. Dixon RA, Barros J. Lignin biosynthesis: Old roads revisited and new roads explored. Open
- 598 Biology. 2019;9. doi:10.1098/rsob.190215
- 33. Raes, J., Rohde, A., Christensen, J. H., Van de Peer, Y., Boerjan W. Genome-Wide
- 600 Characterization of the Lignification Toolbox in Arabidopsis. Plant Physiology. 2014;133: 1051–
- 601 1071. doi:10.1104/pp.103.026484.role
- 602 34. Weng JK, Akiyama T, Bonawitz ND, Li X, Ralph J, Chapple C. Convergent evolution of syringyl
- 603 lignin biosynthesis via distinct pathways in the lycophyte Selaginella and flowering plants. Plant
- 604 Cell. 2010;22: 1033–1045. doi:10.1105/tpc.109.073528
- 605 35. Weng J-K, Li X, Stout J, Chapple C. Independent origins of syringyl lignin in vascular plants.
- 606 Proceedings of the National Academy of Sciences. 2008;105: 7887 LP 7892.
- 607 doi:10.1073/pnas.0801696105
- 608 36. Labeeuw L, Martone PT, Boucher Y, Case RJ. Ancient origin of the biosynthesis of lignin
 609 precursors. Biology Direct. 2015;10: 1–21. doi:10.1186/s13062-015-0052-y
- 610 37. Matsuzaki M, Misumi O, Shin-i T, Maruyama S, Takahara M, Miyagishima S, et al. Genome
- 611 sequence of the ultrasmall unicellular red alga Cyanidioschyzon merolae 10D. Nature. 2004;428:
- 612 653–657. doi:10.1038/nature02398
- 613 38. Collén J, Porcel B, Carré W, Ball SG, Chaparro C, Tonon T, et al. Genome structure and
- 614 metabolic features in the red seaweed Chondrus crispus shed light on evolution of the
- 615 Archaeplastida. Proceedings of the National Academy of Sciences of the United States of

616 America. 2013;110: 5247–5252. doi:10.1073/pnas.1221259110

- 617 39. Brawley SH, Blouin NA, Ficko-Blean E, Wheeler GL, Lohr M, Goodson H V., et al. Insights into
- 618 the red algae and eukaryotic evolution from the genome of Porphyra umbilicalis (Bangiophyceae,
- 619 Rhodophyta). Proceedings of the National Academy of Sciences of the United States of America.
- 620 2017;114: E6361–E6370. doi:10.1073/pnas.1703088114
- 40. Lee JM, Yang EC, Graf L, Yang JH, Qiu H, Zelzion U, et al. Analysis of the draft genome of the
- 622 red seaweed gracilariopsis chorda provides insights into genome size evolution in rhodophyta.
- 623 Molecular Biology and Evolution. 2018;35: 1869–1886. doi:10.1093/molbev/msy081
- 624 41. Page TM, McDougall C, Diaz-Pulido G. De novo transcriptome assembly for four species of
- 625 crustose coralline algae and analysis of unique orthologous genes. Scientific Reports. 2019;9.
- 626 doi:10.1038/s41598-019-48283-1
- 42. Bi G, Liu G, Zhao E, Du Q. Complete mitochondrial genome of a red calcified alga Calliarthron
 tuberculosum (Corallinales). Mitochondrial DNA. 2016;27: 2554–2556.
- 629 doi:10.3109/19401736.2015.1038801
- 630 43. Chan CX, Yang EC, Banerjee T, Yoon HS, Martone PT, Estevez JM, et al. Red and green algal
- 631 monophyly and extensive gene sharing found in a rich repertoire of red algal genes. Current

632 Biology. 2011;21: 328–333. doi:10.1016/j.cub.2011.01.037

- 633 44. Shockey JM, Fulda MS, Browse J. Arabidopsis Contains a Large Superfamily of Acyl-Activating
 634 Enzymes . Phylogenetic and Acyl-Coenzyme A Synthetases 1. Plant physiology. 2003;132: 1065–
 635 1076 doi:10.1104/j...102.020552 doi:10.1104/j...1020552 doi:1004/j...1020552 doi:10.1104/j...1020552 doi:10.1104/j...1020552 doi:10.1104/j...1020552 doi:10.1104/j...1020552 doi:1004/j...1020552 doi:1004/j...1020552 doi:1004/j.
- 635 1076. doi:10.1104/pp.103.020552.ularly
- 636 45. Shockey J, Browse J. Genome-level and biochemical diversity of the acyl-activating enzyme
- 637 superfamily in plants. Plant Journal. 2011;66: 143–160. doi:10.1111/j.1365-313X.2011.04512.x
- 638 46. Li L, Cheng XF, Leshkevich J, Umezawa T, Harding SA, Chiang VL. The Last Step of Syringyl
- 639 Monolignol Biosynthesis in Angiosperms Is Regulated by a Novel Gene Encoding Sinapyl
- 640 Alcohol Dehydrogenase. The Plant Cell. 2001;13: 1567–1586. doi:10.1105/tpc.010111
- 641 47. Hofmann LC, Schoenrock K, de Beer D. Arctic Coralline Algae Elevate Surface pH and

642		Carbonate in the Dark. Frontiers in plant science. 2018;9: 1416. doi:10.3389/fpls.2018.01416
643	48.	Nam O, Shiraiwa Y, Jin E. Calcium-related genes associated with intracellular calcification of
644		Emiliania huxleyi (Haptophyta) CCMP 371. ALGAE. 2018;33: 181-189.
645		doi:10.4490/algae.2018.33.4.21
646	49.	Xue J, Purushotham P, Acheson JF, Ho R, Zimmer J, McFarlane C, et al. Functional
647		characterization of a cellulose synthase, CtCESA1, from the marine red alga Calliarthron
648		tuberculosum (Corallinales). Journal of Experimental Botany. 2021; erab414.
649		doi:10.1093/jxb/erab414
650	50.	Kyndt JA, Meyer TE, Cusanovich MA, Van Beeumen JJ. Characterization of a bacterial tyrosine
651		ammonia lyase, a biosynthetic enzyme for the photoactive yellow protein. FEBS letters. 2002;512:
652		240-244. doi:10.1016/S0014-5793(02)02272-X
653	51.	Barros J, Serrani-Yarce JC, Chen F, Baxter D, Venables BJ, Dixon RA. Role of bifunctional
654		ammonia-lyase in grass cell wall biosynthesis. Nature plants. 2016;2: 16050.
655		doi:10.1038/nplants.2016.50
656	52.	Cooke HA, Christianson C V, Bruner SD. Structure and chemistry of 4-methylideneimidazole-5-
657		one containing enzymes. Current opinion in chemical biology. 2009;13: 460-468.
658		doi:10.1016/j.cbpa.2009.06.013
659	53.	Mohy El-Din SM, El-Ahwany AMD. Bioactivity and phytochemical constituents of marine red
660		seaweeds (Jania rubens, Corallina mediterranea and Pterocladia capillacea). Journal of Taibah
661		University for Science. 2016;10: 471-484. doi:https://doi.org/10.1016/j.jtusci.2015.06.004
662	54.	Mallinson SJB, Machovina MM, Silveira RL, Garcia-Borràs M, Gallup N, Johnson CW, et al. A
663		promiscuous cytochrome P450 aromatic O-demethylase for lignin bioconversion. Nature
664		communications. 2018;9: 2412-2487. doi:10.1038/s41467-018-04878-2
665	55.	Guo J, Ma X, Cai Y, Ma Y, Zhan Z, Zhou YJ, et al. Cytochrome P450 promiscuity leads to a
666		bifurcating biosynthetic pathway for tanshinones. The New phytologist. 2016;210: 525-534.
667		doi:10.1111/nph.13790

668	56.	Hoffmann L, Besseau S, Geoffroy P, Ritzenthaler C, Meyer D, Lapierre C, et al. Silencing of
669		Hydroxycinnamoyl-Coenzyme A Shikimate/Quinate Hydroxycinnamoyltransferase Affects
670		Phenylpropanoid Biosynthesis. The Plant cell. 2004;16: 1446–1465. doi:10.1105/tpc.020297
671	57.	Goujon T, Sibout R, Pollet B, Maba B, Nussaume L, Bechtold N, et al. A new Arabidopsis
672		thaliana mutant deficient in the expression of O-methyltransferase impacts lignins and sinapoyl
673		esters. Plant Molecular Biology. 2003;51: 973-989. doi:10.1023/A:1023022825098
674	58.	Lu F, Marita JM, Lapierre C, Jouanin L, Morreel K, Boerjan W, et al. Sequencing around 5-
675		Hydroxyconiferyl Alcohol-Derived Units in Caffeic Acid O -Methyltransferase-Deficient Poplar
676		Lignins. Plant physiology (Bethesda). 2010;153: 569-579. doi:10.1104/pp.110.154278
677	59.	Guo D, Chen F, Inoue K, Blount JW, Dixon RA. Downregulation of Caffeic Acid 3- O -
678		Methyltransferase and Caffeoyl CoA 3- O -Methyltransferase in Transgenic Alfalfa: Impacts on
679		Lignin Structure and Implications for the Biosynthesis of G and S Lignin. The Plant cell. 2001;13:
680		73-88. doi:10.1105/tpc.13.1.73
681	60.	Li L, Popko JL, Zhang X-H, Osakabe K, Tsai C-J, Joshi CP, et al. A Novel Multifunctional O-
682		Methyltransferase Implicated in a Dual Methylation Pathway Associated with Lignin Biosynthesis
683		in Loblolly Pine. Proceedings of the National Academy of Sciences - PNAS. 1997;94: 5461–5466.
684		doi:10.1073/pnas.94.10.5461
685	61.	Hu Y, Gai Y, Yin L, Wang X, Feng C, Feng L, et al. Crystal Structures of a Populus tomentosa 4-
686		Coumarate : CoA Ligase Shed Light on Its Enzymatic Mechanisms. Plant physiology. 2010;22:
687		3093-3104. doi:10.1105/tpc.109.072652
688	62.	Witzel K, Schomburg D, Kombrink E, Schneider K, Ho K, Stuible H. The substrate specificity-
689		determining amino acid code of 4-coumarate : CoA ligase. PNAS. 2003;100: 8601-8606.
690	63.	Stuible H, Kombrink E. Identification of the Substrate Specificity-conferring Amino Acid
691		Residues of 4-Coumarate : Coenzyme A Ligase Allows the Rational Design of Mutant Enzymes
692		with New Catalytic Properties. Journal of Biological Chemistry. 2001;276: 26893-26897.

- 694 64. Jörnvall H, Persson B, Krook M, Atrian S, Gonzàlez-Duarte R, Jeffery J, et al. Short-Chain
- 695 Dehydrogenases/Reductases (SDR). Biochemistry. 1995;34: 6003–6013.
- 696 doi:10.1021/bi00018a001
- 65. Sattler SA, Walker AM, Vermerris W, Sattler SE, Kang C. Structural and Biochemical
- 698 Characterization of Cinnamoyl-CoA Reductases. Plant physiology. 2017;173: 1031–1044.
- 699 doi:10.1104/pp.16.01671
- 700 66. Filling C, Berndt KD, Benach J, Knapp S, Prozorovski T, Nordling E, et al. Critical Residues for
- 701 Structure and Catalysis in Short-chain Dehydrogenases / Reductases. Biological Chemistry.
- 702 2002;277: 25677–25684. doi:10.1074/jbc.M202160200
- 703 67. Bukh C, Nord-Larsen PH, Rasmussen SK. Phylogeny and structure of the cinnamyl alcohol
- dehydrogenase gene family in Brachypodium distachyon. Journal of Experimental Botany.
- 705 2012;63: 6223–6236. doi:10.1093/jxb/ers275
- 706 68. Youn B, Camacho R, Moinuddin SGA, Lee C, Davin LB, Lewis NG, et al. Crystal structures and
- catalytic mechanism of the Arabidopsis cinnamyl alcohol dehydrogenases AtCAD5 and AtCAD4.

708 Organic & Biomolecular Chemistry. 2006;4: 1687–1697. doi:10.1039/B601672C

709 69. Bomati EK, Noel JP. Structural and kinetic basis for substrate selectivity in Populus tremuloides

sinapyl alcohol dehydrogenase. The Plant cell. 2005/04/13. 2005;17: 1598–1611.

- 711 doi:10.1105/tpc.104.029983
- 712 70. Julián-sánchez A, Riveros-rosas H, Piña E. Evolution of Cinnamyl Alcohol Dehydrogenase
- 713 Family Evolution of Cinnamyl Alcohol Dehydrogenase Family. In: Weiner H, Plapp B, Lindahl R,
- 714 Maser E, editors. Enzymology and Molecular Biology of Carbonyl Metabolism. West Lafayette:
- 715 Purdue University Press; 2006. pp. 142–153.
- 716 71. von Borzyskowski LS, Rosenthal RG, Erb TJ. Evolutionary history and biotechnological future of
 717 carboxylases. Journal of Biotechnology. 2013;168: 243–251. doi:10.1016/j.jbiotec.2013.05.007
- 718 72. Haas BJ, Papanicolaou A, Yassour M, Grabherr M, Philip D, Bowden J, et al. De novo transcript
- reference generation and analysis with Trinity. Nature

720 protocols. 2013;8: 1–43. doi:10.1038/nprot.2013.084.De

- 721 73. Conesa A, Götz S, García-Gómez JM, Terol J, Talón M, Robles M. Blast2GO: A universal tool
- for annotation, visualization and analysis in functional genomics research. Bioinformatics.
- 723 2005;21: 3674–3676. doi:10.1093/bioinformatics/bti610
- 724 74. Kanehisa M, Sato Y, Kawashima M, Furumichi M, Tanabe M. KEGG as a reference resource for
- gene and protein annotation. Nucleic acids research. 2016;44: 457–462. doi:10.1093/nar/gkv1070
- 726 75. Luo W, Brouwer C. Pathview: an R/Bioconductor package for pathway-based data integration and
- visualization. Computer applications in the biosciences. 2013;29: 1830–1831.
- doi:10.1093/bioinformatics/btt285
- 729 76. Smit A, Hubley R, Green P. RepeatMasker Open-4.0.
- 730 77. Haas BJ, Delcher AL, Mount SM, Wortman JR, Smith RK, Hannick LI, et al. Improving the
- 731 Arabidopsis genome annotation using maximal transcript alignment assemblies. Nucleic Acids
- 732 Research. 2003;31: 5654–5666. doi:10.1093/nar/gkg770
- 733 78. Stanke M, Schöffmann O, Morgenstern B, Waack S. Gene prediction in eukaryotes with a
- generalized hidden Markov model that uses hints from external sources. BMC Bioinformatics.
- **735** 2006;7: 1–11. doi:10.1186/1471-2105-7-62
- 736 79. Finn RD, Clements J, Eddy SR. HMMER Web Server: Interactive Sequence Similarity Searching.
 737 Nucleic Acids Research. 2011;39: W29–W37. doi:10.1093/nar/gkr367
- Rice P, Longden I, Bleasby A. EMBOSS: The European Molecular Biology Open Software Suite.
 Trends in Genetics. 2000;16: 276–277. doi:10.1016/S0168-9525(00)02024-2
- 740 81. Edgar RC. MUSCLE: a Multiple Sequence Alignment Method With Reduced Time and Space
- 741 Complexity. BMC Bioinformatics. 2004;5. doi:10.1186/1471-2105-5-113
- 742 82. Mahram A, Herbordt MC. Fast and Accurate NCBI BLASTP: Acceleration with Multiphase
- 743 FPGA-based Prefiltering. Proceedings of the 24th ACM International Conference on
- 744 Supercomputing. New York, NY, USA: ACM; 2010. pp. 73–82. doi:10.1145/1810085.1810099
- 745 83. Capella-Gutiérrez S, Silla-Martínez JM, Gabaldón T. trimAl: a tool for automated alignment

- trimming in large-scale phylogenetic analyses. Bioinformatics. 2009/06/08. 2009;25: 1972–1973.
- 747 doi:10.1093/bioinformatics/btp348
- 748 84. Luo A, Qiao H, Zhang Y, Shi W, Ho SY, Xu W, et al. Performance of criteria for selecting
- evolutionary models in phylogenetics: a comprehensive study based on simulated datasets. BMC
- 750 evolutionary biology. 2010;10: 242. doi:10.1186/1471-2148-10-242
- 751 85. Nguyen L-T, Schmidt HA, von Haeseler A, Minh BQ. IQ-TREE: A Fast and Effective Stochastic
 752 Algorithm for Estimating Maximum-Likelihood Phylogenies. Molecular Biology and Evolution.
- **753** 2015;32: 268–274.
- 86. Hoang DT, Chernomor O, von Haeseler A, Minh BQ, Vinh LS. UFBoot2: Improving the Ultrafast
 Bootstrap Approximation. Molecular Biology and Evolution. 2018;35: 518–522.
- 756 87. Rambaut A, Drummond A. FigTree v1. 3.1 Institute of Evolutionary Biology. University of
 757 Edinburgh. 2010.
- 88. Bolger AM, Lohse M, Usadel B. Trimmomatic: A flexible trimmer for Illumina sequence data.
 Bioinformatics. 2014;30: 2114–2120. doi:10.1093/bioinformatics/btu170
- 760 89. Bankevich A, Nurk S, Antipov D, Gurevich AA, Dvorkin M, Kulikov AS, et al. SPAdes: A new
- 761 genome assembly algorithm and its applications to single-cell sequencing. Journal of
- 762 Computational Biology. 2012;19: 455–477. doi:10.1089/cmb.2012.0021
- 76390.Xue W, Li JT, Zhu YP, Hou GY, Kong XF, Kuang YY, et al. L_RNA_scaffolder: Scaffolding
- 764 genomes with transcripts. BMC Genomics. 2013;14: 1–14. doi:10.1186/1471-2164-14-604
- 91. Parra G, Bradnam K, Korf I. CEGMA: A pipeline to accurately annotate core genes in eukaryotic
- 766genomes. Bioinformatics. 2007;23: 1061–1067. doi:10.1093/bioinformatics/btm071
- 767 92. Simão FA, Waterhouse RM, Ioannidis P, Kriventseva E V., Zdobnov EM. BUSCO: Assessing
- 768 genome assembly and annotation completeness with single-copy orthologs. Bioinformatics.
- 769 2015;31: 3210–3212. doi:10.1093/bioinformatics/btv351
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772 Supporting Information

773

774 Fig S1. Completeness of the *C. tuberculosum* transcriptome dataset.

- (A) Transcriptome sequences show high recovery of eukaryotic genes in CEGMA/BUSCO analysis.
- Percentage of genomic scaffolds with transcriptome support and transcriptomic scaffolds alone that share
- amino acid sequences with the core eukaryotic gene databases including CEGMA, BUSCO eukaryotic,
- and BUSCO Viridiplantae. Transcriptome encoded amino acid sequences were searched against the
- databases using BLASTP (orange) or TBLASTN (yellow), and genomic scaffolds were searched against
- 780 the databases using TBLASTN (blue)
- **(B)** Transcriptomic support of genomic data analyzed by GC content and transcript length.
- 782 The distribution of GC content (above) against transcript lengths is shown for scaffolds with
- transcriptome support (blue) and scaffolds without transcriptome support (yellow) (right).

Fig S2. C3H, C4H, F5H, P450 candidates from *C. tuberculosum* in relation to plants and other taxa.

- (A) Partial alignment of *C. tuberculosum* P450 candidates with C3H, C4H, and F5H from *A. thaliana*,
- and a novel F5H from *Selaginella moellendorffii*. Heme binding domain residues, secondary structure
- stabilizing K helix residues, PXRX, and the I-helix are indicated [8]. Sites with <80% coverage were
- removed. A strong candidate for beta-carotene synthesis is indicated with a triangle.
- (B) Unrooted CYP450 maximum likelihood gene tree with C. tuberculosum (magenta dots) and
- 790 additional taxa (Embryophyta dark green, Chlorophyta light green, Rhodophyta red, Animalia and
- 791 Opisthokonta purple, Bacteria and Cyanobacteria blue, Oomycota, Mycetozoa and Fungi yellow,
- 792 Ochrophyta brown). Functionally demonstrated plant C3H, C4H, and F5H are labeled (+). Additional
- functional groups are labeled [9]. Ultrafastbootstrap values > 95 are marked by *. Model = VT+F+G4.
- 794

Fig S3. CCoAOMT candidates from *C. tuberculosum* in relation to plants and other taxa.

- (A) Partial alignment of *C. tuberculosum* CCoAOMT sequence candidates with CCoAOMT from land
- 797 plants. Substrate recognition residues (black triangle), divalent metal ion and cofactor binding residues
- 798 (grey triangle), catalytic residues (back square), and the positively charged R220 necessary for substrate
- recognition (grey square) are indicated. Sites with < 70% coverage were removed.
- 800 (B) Unrooted maximum likelihood gene tree of biochemically characterized plant O-methyltransferases
- 801 with *C. tuberculosum* (magenta dots) and additional taxa (Embryophyta dark green, Chlorophyta light
- 802 green, Rhodophyta red, Animalia and Opisthokonta purple, Bacteria and Cyanobacteria blue,
- 803 Oomycota, Mycetozoa and Fungi yellow, Ochrophyta brown). Functionally demonstrated plant

804	CCoAOMT are labeled (+). Additional functional groups are labeled [13]. Ultrafastbootstrap values > 95
805	are marked by $*$. Model = LG + G4. JMT, SAMT, and BAMT are closely related to OMTs.
806	
807	Fig S4. CSE candidates from C. tuberculosum in relation to plants and other taxa.
808	(A) Partial alignment of C. tuberculosum CSE sequence candidates with CSE from land plants. Acyl
809	transferase motifs (HX ₄ D), lipase motifs (GXSXG) and active site residues (triangle) are indicated. Sites
810	with $< 70\%$ coverage were removed.
811	(B) Unrooted maximum likelihood gene tree of C. tuberculosum CSE candidates (magenta dots) and
812	additional taxa (Embryophyta - dark green, Chlorophyta - light green, Rhodophyta - red, Animalia and
813	Opisthokonta – purple, Bacteria and Cyanobacteria – blue, Oomycota, Mycetozoa and Fungi – yellow,
814	Ochrophyta – brown). Functionally demonstrated plant CSE are labeled (+). Additional functional groups
815	are labeled. Ultrafastbootstrap values > 95 are marked by $*$. Model = VT+G4.
816	
817 818 819	Fig. S5. A visual representation of the <i>C. tuberculosum</i> sequences present in the starch and sucrose metabolism pathway from the KEGG based annotation. KEGG based annotation showing the starch and sucrose metabolic pathway with <i>C. tuberculosum</i>
820	annotations highlighted. The gradient map in the top right corner indicates the level of transcription, with
821	white and dark pink coloring representing absence and presence of expression respectively. The annotated
822	map, number "00500", was extracted in the provided R file using the pathview program.
823	
824	Table S1. Summary statistics for the C. tuberculosum genome assembly.
825	Scaffolds are categorized as shared with either red algal (Pyropia yezoensis) genomic scaffolds,
826	eukaryotic sequences, or other bacteria sequences based on sequence similarity.
827	Table S2. Top hits against Arabidopsis thaliana (taxid:3702) using <i>Calliarthron</i> sequences as the
828	search query (BLASTP). Query sequence is indicated by contig number. Result hits are indicated by
829	description (At tax ID 3702) and colored by overall alignment scores with red (>=200), pink (80-200),
830	green (50-80), blue (40-50), and black (<40) that are most to least reliable scores in that order.
831	
832	Table S3. KEGG annotations of Calliarthron tuberculosum reads from the combined transcriptomic
833	dataset. Unique reads are represented by their contig identifier (contig_gene_isoform) and matched with
834	their annotated KEGG based identifier (KO_identifier) and associated protein name.
835	

836 Table S4. Listed representation of the *C. tuberculosum* sequences present in starch and sucrose

837 metabolism pathway from the KEGG based annotation.

838 *C. tuberculosum* sequences were extracted from the KEGG based starch and sucrose metabolism pathway

number "00500". "KEGG Identifier" refers to the specific KEGG code for the gene, "Contig Name"

- 840 refers to the sequence identifier from the *Calliarthron* transcriptome where the values represent the contig
- 841 name_gene number_gene isoform and "Gene Name" refers to the gene acronym, the gene name, and its
- enzyme commission (EC) number. Sequences were extracted in the provided R file using the pathview
- 843 program.
- 844

Table S5. A list of calcification related gene candidates identified from KEGG-based annotations of the *C. tuberculosum* transcriptome.

- 847 Calcification gene candidates were initially selected based on a literature search, and then C.
- 848 *tuberculosum* sequences were identified manually from the KEGG based annotations (annotation file
- available on Github), thus this is not an exhaustive list. The genes are organized by their functional
- 850 classification indicated as "overall function", while "KEGG Identifier" refers to the specific KEGG code
- 851 for the gene, "Contig Name" refers to the sequence identifier from the *Calliathron* transcriptome where
- the values represent the contig name gene number gene isoform and "Gene Name" refers to the gene
- acronym, the gene name, and its enzyme commission (EC) number.
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Figure 1



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Figure 2



Figure 3



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c124623_g1_i1	26AVVTGASGFIGGHIVKGLLERRYNVVAVVRDVTPSKIGHLILLKQQYADLLSFATVSDLTVKSDALHSA
c118951_g1_i1	67VAVSGITGFIGGHIANDLLSKGYIVHGTLRKNTPERTAHLTSHKNAAKCLELFEADL-VKSGSFDEC
Ta/CCR2	20VCVTGAGGELASWLVKRLLQXGYNVRGTVRNPVDPKNDHLRAFDGAADRLVLLRADL-MEPETLVEA
PtCCR	13VCVTGAGGEIASWLVKLLLERGYTVRGTVRNPEDOKNAHLROLEGAEERLTLVKADL-MDYNSLLNA
LpCCR	18VCVTGAAGYIASWLVKLLLERGYTVKGTVRNPDDPKNAHL&ALDGAVERLILCKADL-LDYDAICAA
HVCCR	21VCVTGAAGYIASWLVKLLLERGYTVKGTVRNPOOPKNAHLKALDGAAERLVLCKADL-LDYDAICAA
TaCCR1	22VCVTGAAGYIASWLVKLLLERGYTVKGTVRNPDDPKNAHLKALDGAAERLVLCKADL-LDYDAICAA
AtCCR1	13VCVTGAGGYIASWIVKILLERGYTVKGTVRNPDDPKNTHLRELEGGKERLILCKADL-ODYEALKAA
StCCR	9VCVTGAGGEIASWLVKLLLEKGYTVRGTVRNPDDPKNGHLKELEGAKERLILLRADL-LDYQSLREA
PtCCR	15ICVTGAGGEIASMMVKLLLDKGYTVRGTARNPADPKNSHLBGLEGAEERLTLCKADL-LDYESLKEA
EgCCR	13VCVTGAGGF1ASWTVKLLLERGYTVRGTVRNPDDPKNGHLRELEGASERLTLYKGDL-MDYGSLEEA
consensus	81
c95473 g1 i1	155 FKGATAVVII-AAPVGVOMRHISALLPSSKVMDAVDAVDAASAAGATVRRVVFLSTEMSVFDPLAPPPTGERRPLGEEDN
c124623 g1 i1	95 MSSSDTVFRVANPMSSAEGSDAFVSAST-GAVRAVLETAKDVGARR/VLTASMASVCGDOASLNPRHVY
c118951 g1 i1	133 LTGCKMAMEVASPYHMDAKDPOKELVNPAV-NGTLNFLRSCKKAGVEKWLTSSVAAIAGEGRCEHTF
TaCCR2	86 FTGCEGIFHAASPVTDOPEKNIEPAI-BOTKYVITAAADMGIKRVVFTSTIGTVYMNENEDPSKPVDDTCN
Pt.CCR	79 INCCOVERVASPVTDOPRENVEPAV-NOTKINVLDACAVAGVBRVVFTSSIGAVYMDPSRDYDALVDENCH
LDCCR	84 AEGCHGVFHTASPVTDDPEONVEPAV-BGTEYVINAADAG-TVREVVFTSSIGAVTMDPNRGPDVVVDESCH
HVCCR	87 VEGCHGVFHTASPVTDOPROMVEPAV-BOTEYVIDAADAG-TVBRVVFTSSIGAVTMDPNRGPDV/VDESCN
TaCCR1	88. VEGCHGVFHTASPVTDOPROMVEPAV-BUTEYVINAAADAG-TVBRVGVTSSIGAVTMDPNRGPDV/VDESCN
At CCR1	79 IDOCDOVENTASPVTDDPEONVEPAV-NGAKEVINAAAEAKVKKVVITSSIGAVTMDPNRDPEAVVDESCH
SLCCR	75 IYOCDOVENTASPVTDOPROMVEPAV-IGTKNVITAAAEAKVGRVVPTSSIGTVYMDPNRAPDKVVDETCN
Pt/CCR	81 TOCCDOVERTASEVTDOPERMVEPAV-NOTKINVI LAARAKVBRVVETSSTGAVYMDPNKOPDVV LDESCN
EgCCR	79 TROCDOWNTASPVTDOPROMVEPAV-LOTKINVTVAAAEAKVBRVVPTSSLGAVTMDPNBGPDVVVDESCN
CODSEDSUS	161
	NUVCY motif
a95473 al il	214 YOVSENDELSEDY LATERTVACHEL REPAIR-PST PETVOLUPTEAMOR/MEMVOTSTANPMETLISCHAGEPPE

c95473_g1_i1	234 YOVSRNDRLSSDYTATINETVAEMKLMSRATR-PSIPFTVCSLVPTFAMGFVMSHVQISTAHPMRFLRSCMAGRFPE
c124623_g1_i1	163 TESDANNECGSS
c118951_g1_i1	200 TEADMNTKSSLRCLPYTISKVEAEKAAMKFIKE-EAPDMKLVVINPIVVIGPNLTKSK-NASVKIFEHIVNGT-FGGILD
Ta/CCR2	156 SDLEYCKKTAWYCIWKTVAEQDALETARQ-RGIELIVVNPVLVLGPLLQPTV-NASTEHVMKYLTGS-AKTYVN
PtCCR	149 SNLDYCKETNWYCT KTVAEKAAMERAKD-KGLDLVVVNPCVVLGPVLQSSI-NSSIIHILKYLTGS-AKTYAN
LPCCR	155 SDLEFCKKTRINNYCH SKAVAEQAAMEAARK-RGIDLVVVNPVLVVGPLLQPTV-NASAAHILKYLDGS-AKKYAN
HVCCR	158 SDLEFCKKTNWYCT KAVAEQAAWEKARA-RGVDLVVVNPVLVVGPLLQPTV-NASAAHILKYLDGS-ARKYAN
Ta/CCR1	159 SDLEFCKKTNWYCT KAVAEQAMEKAAA-RGVDLVVVNPVLVVGPLLQPTV-NASAAHILKYLDGS-AKKYAN
AtCCR1	149 SDLDFCKNTNWYCT BMVAEQAAWETAKE-KGVDLVVLNPVLVLGPPLQPTI-NASLYHVLKYLTGS-AKTYAN
StOCR	145 SDLGFCKNTRWWYCI SKTVAEKTAMDEARE-KGVDLVVINPVLVLGPLLQPTV-NASVLHILKYLTGS-AKTYAN
PtCCR	151 SDLEFCRNTNWYCT BAVAEQAAMDMAKE-RGVDLVVVNPVLVLGPLLQPTV-NASITHILKYLTGS-AKTYAN
EgCCR	149 SDLEFCKSTNWYCT KAVAEKAAWPEAKE-RGVDLVVINPVLVLGPLLQSTI-NASIIHILKYLTGS-AKTYAN
Consensus	241



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c145607_g1_11	158	FGDVTNDPL	LPRTAGAFAEYAWWKATLLA	PVPESVTFEQAAALPVAVGTSMQAFD
c137515_g1_i1	368	YNLCAEMRFAAT	PPTDGCLARRVNHSAAFCY	KLPDRVSFEAAALLEPLSVALEALA
c137498 g1 i1	155	TNLOOK IRSTOGNGVMPDGTSRITADGK	SIYHYMOCSSFSEYTVVADISVV	KVRDDAPLDKVCLLACGI TTGLGAVR:
AtADH1	110	SNMCDL-LRINTEROGMIHDGESRFSINGK	PIYHFLGTSTFSEYTVVHSGQVA	KINPDAPLDKVCIVSCGLSTGLGATLA
c121452_g1_i1	337	ENVCERGYQGLFLGPSSGIWGKSKLEYH	-TMGOCFARVORIEARFAT	11PGNVPSGIVCPLVCGGGTVPEP11
c121452 gl 12	364	ENVCEKGYQGLFLGPSSGIWGKSKLEYH	-THGGCFARVQR IEARFAT	11PG8VPSGIVCPLVCGGGTVPEPIII
AtCADS	111	EQYCPKKIWSYNDVYINGQ	PTQGG-FAKATVVBQKFVV	KI PEGMAVEQAAPLLCAGVTVYSPLSI
PaCAD	91	EQYCSKRIWTYNDVNHDGT	PTQGG-FASSMVVDQMEVV	RIPENLPLEQAAPLLCAGVTVFSPMKI
AtCAD9	114	ENYCPQMSFTYNAIGSDGT	90NYGG-YSENIVVDQRFVL	RFPENLPSDSGAPLLCAGITVYSPMK
PaSAD1	91	EQYCERVVWTYNSIYLDGS	PTPGG-YSSLMVCDQRFIV	KIPENLPPDAAAPLLCAGITVYSPMQ
AtCAD8	110	ENYCPKSIQTYGFPYYDNT	ITY GG-YSDHMVCEEGFVI	RIPONLPLDAAAPLLCAGITVYSPMK
consensus	401			
		NADPH BD	NADPH BD	
c145607_g1_i1	214	MGLKSGQKVFISGDAGGVGVHAIQIAKSVF	GASEVATEASEAR COFVDRAGAD	IIIDYKKDDAGEVLKGWADVVLDCTGI
c137515_g1_i1	424	RAKLSAGDAVFIAGAGPIGLMCALAAR-AA	GAARVVIADVDAAR	LRVATDA-G-
c137498_g1_i1	233	TSNVEKGATVAVNGLGCVGLSCISAAK-ER	GASRIIGVDINPGC	FAKAMEF-G
AtADH1	189	VAKPEROOSVAIPOLGAAGLGAAEGAR-IA	GASRI IGVDFNSKR	FDQAKEF-G
c121452_g1_i1	410	YGFPGAILAVSSVGGIGTAAIKLAR-LR	GL-TVWAISSTPSK	RDGALSA-G-
c121452 g1 12	437	YGFPGAILAVSGVGGIGTAAIKLAR-LR	GL-TVWAISSTPSK	RDGALSA-G-
AtCAD5	175	FGL#QPGLRGGI1GLGGVGHMGVKIAK-AM	GH-HVTVISSSNSK	REEALQDLG
PaCAD	155	FONTEPOKRCGILGLOGVGHMGVKIAK-AF	GL-HVTVISSSDEK	KEEMEVLG
AtCAD9	178	YGMTEAGKHLGVAGLGGIGHVAVKIGK-AF	GL-KVTVISSSSTC	AEEAINHLG-
PaSAD1	155	FONTEPOKSLOVVGLOGIGHMAVKFGK-AF	GL-KVTVISTSPSK	EKEAKEYLG
AtCAD8	174	HGLDKPGMHIGVVGLGGIGHVGVKFAK-AM	GT-KVTVISTSERK	RDEAINRLG
consensus	481	···· ··· ··· [*]* ······ ·	•	•
			NADPH BD	
c145607_g1_i1	453	NTDKTAPTSHARWGAIRLCSKKPSARFCIV	HLGPSSXTILETPSEVAXSINVF	PMAFARPVRGSTLALQSSYSSVAVRR
c137515_g1_i1	512	NARSG-GVVVLN	MGAPVVRVPVLD	A-GCRE8870-A
c137498 g1 11	328	ACHKGWGESCII	WAASGKEISTRP	FQLVTG
AtADH1	284	CVHDGWGVAVLV	WPSKDDAFKTHP	NFLNE
c121452_g1_i1	496	CLKIN-GTFVRV	GIPPSSDMMFEHNFIP	-IF00
c121452 g1 12	523	CLKIN-GTFVRV	SIPPSSOMMFERNFIP	-IFQQ
AtCADS	264	LLKLD-GKLILM	WINNPLQF-LTP	LIMLOR
PaCAD	244	LLETN-GKLVML	WVPEPTHF-VTP	LILGR
AtCAD9	267	LLKVN-GKLIAI	SLPERPLELPMFP	L-VLOR
PaSAD1	244	LLKVN-GKLVLV	MPEKPLSLPPVA	-T80-
AtCAD8	263	LLKHK-GKLVMV	SAPEKPLELPVMP	L-IFER
consensus	721			

Figure 4

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