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4	How to Build a Fruit: Transcriptomics of a Novel Fruit Type in the
5	Brassiceae
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19 20	Abstract: Comparative gene expression studies are invaluable for predicting how existing genetic

21 pathways may be modified or redeployed to produce novel and variable phenotypes. Fruits are

22 ecologically important organs because of their impact on plant fitness and seed dispersal,

23 modifications in which results in morphological variation across species. A novel fruit type in 24 the Brassicaceae known as heteroarthrocarpy enables distinct dispersal methods in a single fruit 25 through segmentation via a lateral joint and variable dehiscence at maturity. Given the close 26 relationship to Arabidopsis, species that exhibit heteroarthrocarpy are powerful models to 27 elucidate how differences in gene expression of a fruit patterning pathway may result in novel 28 fruit types. Transcriptomes of distal, joint, and proximal regions from *Erucaria erucarioides* and 29 Cakile lanceolata were analyzed to elucidate within and between species differences in whole 30 transcriptome, gene ontology, and fruit patterning expression profiles. Whole transcriptome 31 expression profiles vary between fruit regions in patterns that are consistent with fruit anatomy. 32 These transcriptomic variances do not correlate with changes in gene ontology, as they remain 33 generally stable within and between both species. Upstream regulators in the fruit patterning 34 pathway, FILAMENTOUS FLOWER and YABBY3, are expressed in the distal and proximal 35 regions of *E. erucarioides*, but not in the joint, implicating alterations in the pathway in 36 heteroarthrocarpic fruits. Downstream gene, INDEHISCENT, is significantly upregulated in the 37 abscissing joint region of C. lanceolata, which suggests repurposing of valve margin genes for 38 novel joint disarticulation in an otherwise indehiscent fruit. In summary, these data are consistent 39 with modifications in fruit patterning genes producing heteroarthrocarpic fruits through different 40 components of the pathway relative to other indehiscent, non-heteroarthrocarpic, species within 41 the family. Our understanding of fruit development in Arabidopsis is now extended to atypical 42 siliques within the Brassicaceae, facilitating future studies on seed shattering in important 43 Brassicaceous crops and pernicious weeds.

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46 Introduction

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48 Studying gene expression patterns across plant structures and species can elucidate how their 49 modification may produce morphological variation (1,2). Fruits are diverse and ecologically 50 relevant plant structures to investigate because their morphological variation determines how 51 their seeds are dispersed (3,4). There are multitudinous fruit morphologies in nature, and they are 52 often categorized as fleshy or dry. Fleshy fruits are distributed primarily by animals, as the seeds 53 are discarded before or after consuming. Dry fruits however, may be dispersed by animals, wind, 54 or water. Dry fruits are further classified by whether they are dehiscent, releasing seeds into the 55 environment, or indehiscent, releasing seeds in a protected fruit wall propagule. Thus, variation 56 in fruit morphology is directly tied to differences in dispersal capabilities. 57 58 Arabidopsis thaliana (Brassicaceae) is the premier model for dry dehiscent fruits. Arabidopsis 59 fruits have been characterized from gynoecium formation to seed release, and many genes 60 responsible for fruit development are described, as are their interactions (5-7). This knowledge 61 forms a basis of comparison in the investigation of complex trait morphologies that diverge from 62 Arabidopsis, especially amongst close relatives e.g., the loss of dehiscence in many species

63 across the Brassicaceae (1).

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Brassicaceae fruits vary markedly in shape, structure, and size (1,8). Their variation in
dehiscence is a focal point for research because it fundamentally changes fruit structure,
subsequently affecting dispersal and diversification (9). A prerequisite for exploring how
differences in fruit morphology are achieved across the Brassicaceae is familiarity with both the
fruit structure and underlying genetic pathways in *Arabidopsis* (10.11). *Arabidopsis* fruits.

70 hereafter referred to as typical siliques, are composed of five basic elements: valve, replum, 71 seeds, septum, and valve margins. The valve, synonymous with ovary wall in Arabidopsis, is the 72 outermost tissue of the fruit that protects the developing seeds and is separated from the replum 73 at maturity to release seeds. The replum is the persistent placental tissue to which the seeds are 74 attached. The septum, which connects to the replum, divides the fruit into two locules or 75 chambers. The valve and replum are separated by the valve margin, which consists of a 76 lignification and separation layer. Thus, proper fruit formation relies on the establishment of 77 medial (replum) and lateral (valves and valve margin) components (12). As the fruit dries, 78 tension is created via the lignified layer, which facilitates the separation of the valves from the 79 replum at the separation layer (13). This general morphology is stable across most dehiscent 80 members of Brassicaceae (1).

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82 The causal factors for dehiscence have been well characterized in Arabidopsis (14–17), with 83 proper formation and positioning of the valve margin being a key to this process. The valve 84 margin pathway is essential for spatial regulation and development of valve, replum, and valve 85 margin tissues (11,18–23). Briefly, FRUITFULL (FUL) and REPLUMLESS (RPL), as well as 86 other upstream regulators, restrict the expression of the valve margin genes to two cell layers 87 between the valve and replum, respectively. The valve margin genes, SHATTERPROOF 1/2 88 (SHP1/2), INDEHISCENT (IND), SPATULA (SPT), and ALCATRAZ (ALC), are responsible for 89 the formation of the valve margin, specifically of the separation and lignification layers that 90 control dehiscence (Fig 1). Upstream regulators of FUL and RPL, e.g., APETALA2 (AP2), 91 FILAMENTOUS FLOWER (FIL), YABBY3 (YAB3), and JAGGED (JAG) are also key to precise 92 positioning of the valve margin because they tightly regulate downstream processes. In sum,

93 replum and valve genes function in an antagonistic manner to ensure proper formation of these94 regions of the fruit(12).

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96 Figure 1. Diagram of simplified valve margin pathway for fruit dehiscence in *Arabidopsis*

97 *thaliana;* valve margin. R, replum. Sl, separation layer. ll, lignification layer. Valve margin = sl

98 + II. Modified from data available in (10-11,14) and figure 2 (36).

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100 Most of the *Arabidopsis* valve margin genes are pleiotropic and many of them share

101 indehiscence as a phenotype of mutation. For example, a mutation in any of the following genes

102 results in indehiscent fruits in *Arabidopsis: SHP1/2, SPT, ALC* and *IND* (24–27). Overexpression

103 of FUL or NO TRANSMITTING TRACT (NTT) also results in indehiscent fruits (28,29); FUL

104 overexpression completely suppresses *SHP1/2*, resulting in reduced lignification in the en*b* layer

and reduced valve margin formation; overexpression of NTT phenocopies the *ful* mutation

106 resulting in valve margin specific genes being expressed throughout valve. In summary, a

107 modification of many components in this pathway results in a loss of dehiscence. Because

108 indehiscence is observed in at least 20 different lineages across the family, it is likely that this

109 phenotype evolved via multiple modifications to this pathway (30). As such, there is no singular

alteration to the fruit patterning pathway implicated in this shift for all tribes.

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To date, little is known about the genetic basis of indehiscence in the Brassicaceae, although it is currently being bridged by studies in taxa with varying indehiscent morphologies. Recently, a study demonstrated a deviation in expression of eight key genes between pod shatter sensitive species and shatter resistant species of *Brassica* and *Sinapis* (2). In *Lepidium*, there has been an evolutionary shift from dehiscence to indehiscence, e.g., valve margin genes that are conserved

between the dehiscent *L. campestre* and *Arabidopsis* have been lost in the indehiscent *L. apellianum* (31,32). Upregulation in upstream regulator *AP2* has been suggested as a factor in
this indehiscence (32).

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121 A notable morphological adaptation is the evolution of a complex fruit type known as 122 heteroarthrocarpy, which is only found in some members of the tribe Brassiceae (30,33,34). This 123 modified silique is defined by the presence of a variably abscising central joint, an indehiscent 124 distal region, and a variably dehiscent proximal region (Fig 2). As such, this novel morphology 125 offers an opportunity to investigate fruit variation beyond shifts from dehiscent to indehiscent. 126 Anatomically, heteroarthrocarpic fruits appear most like *Arabidopsis* siliques in their proximal 127 regions, varying by a lack of a valve margin cell layer in indehiscent variants (35–37). There are 128 three described variations of heteroarthrocarpy: a non-abscising joint with a dehiscent proximal 129 region, an abscising joint with an indehiscent proximal region or an abscising joint with a 130 dehiscent proximal region (36). These subtypes have evolved multiple times, perhaps as a bet 131 hedging strategy in response to selective pressure from hostile desert environments (9,37). 132 Heteroarthrocarpic subtypes may be developmental enablers that have facilitated changes in fruit 133 morphology across the tribe, which would explain heteroarthrocarpy's evolutionary lability (36). 134 Regardless of lability, all types are linked by the mechanism in which seeds from the same fruit 135 are released by different means. In other words, the joint is the novel and unifying feature of 136 heteroarthrocarpy (36).

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140	Figure 2. Mature and young heteroarthrocarpic fruits. (A), Mature Erucaria erucarioides fruit in
141	lateral view before dehiscence (left), and medial view after dehiscence (right). (B), Young E.
142	erucarioides fruit in medial view. C, Cakile lanceolata fruit in lateral view before dehiscence
143	(left), and medial view after joint abscission (right). (D), Young C. lanceolata fruit in medial
144	view; Modified from figure 1 (36). White arrows indicate joint region; blue arrows indicate
145	replum. Scale bars = 5mm
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148	A comparison of expression patterns between heteroarthrocarpic subtypes is potentially
149	informative for formulating hypotheses about its evolutionary origins. Erucaria erucarioides and
150	Cakile lanceolata, hereafter referred to as Erucaria and Cakile, are two well-studied
151	representatives for heteroarthrocarpy because of their close relation and divergent subtypes (Fig
152	2) (9,10,36,37). In previous studies it was hypothesized that the formation of heteroarthrocarpy
153	is the result of repositioning of the valve margin, such that the valve is only present in the
154	proximal region of the fruit, unlike in Arabidopsis where it is found in the entire ovary (36). In
155	other words, the joint is the distal portion of the valve margin. This hypothesis was partially
156	supported by comparative gene expression data of some, but not all, genes in the valve margin
157	pathway using a candidate gene approach (10). However, that study did not definitively
158	determine how the pathway has been repositioned because it did not investigate upstream genes.
159	Candidate gene approaches will, by design, overlook non-targeted genes, and a lack of in situ
160	hybridization does not necessarily indicate a lack of expression. Further, the basis of the joint
161	remains unknown.
162	No study to date has investigated transcriptional variation of heteroarthrocarpic fruits sectioned

163 transversely into distal, joint and proximal regions. This approach is complementary to prior

178	Materials and Methods
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172	in heteroarthrocarpy.
171	expression of fruit patterning transcripts will be consistent with repositioning of the valve margin
170	expect gene expression to be consistent with anatomical features within fruits, and that
169	patterns, unique or shared, between and within, two variant heteroarthrocarpic species. We
168	future research regarding the evolution of the joint. Herein, the objective is to uncover transcript
167	expression patterns between and within Erucaria and Cakile, and will set the groundwork for
166	players involved in the formation of heteroarthrocarpy. They will clarify unique and shared gene
165	Expression profiles from these regions will elucidate broad patterns and potentially identify key
164	research because it quantifies expression of all transcripts in discrete regions of a whole system.

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180 Plant material

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182 Seeds from *Erucaria erucarioides* (Coss. and Durieu) Müll.Berol and *Cakile lanceolata* (Willd.)

183 O.E.Schulz were obtained from the late César Gómez-Campo's and KEW royal botanical

184 garden's seed collections, respectively. Vouchers for *Cakile* and *Erucaria* have been deposited in

185 the Vascular Plant Herbarium at the University of Alberta, and the Harvard University Herbaria,

respectively. Seeds were germinated in 1% agar and transferred to clay pots containing a 2:1 soil

187 (Sungro sunshine mix #4, Agawam, MA, USA) to perlite mixture. Plants were grown under a

188	16/8-hour light/dark schedule at 24°C with scheduled watering in the University of Alberta,
189	Department of Biological Sciences, growth chambers.
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191	Distal, joint, and proximal regions from 10mm fruits (~10 days post fertilization) were collected
192	and flash frozen in liquid nitrogen prior to storage at -80°C. Distal and proximal regions were
193	classified as all tissue ~1mm above or below the joint, and the joint is remaining tissue between
194	distal and proximal regions (Fig 2). The 10mm fruit size is roughly equivalent to Arabidopsis
195	stage 17A fruits (7), which go through elongation and cell expansion before maturity. This size
196	was chosen to capture late stage valve margin gene expression because the valve margin is easily
197	distinguished at this stage, and an increase in lignification is observed in key layers, e.g., enb.
198	(36).
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202 203	RNA isolation and cDNA library preparation
204	RNA was extracted from frozen tissue using manual grinding and a Qiagen RNeasy micro kit
205	(Hilden, Germany) with the following amendments to protocol: RNA was incubated in nuclease
206	free water for five minutes prior to elution, and this eluate was spun through the same extraction
207	column to maximize RNA yield. RNA concentration was verified using a Nanodrop ND-1000

- spectrophotometer (Software version 3.1.2), and quality was confirmed using the Agilent 2100
- bioanalyzer (Software version B.02.09.SI720). All cDNA samples were set at the same
- 210 concentration of the most dilute RNA extraction. Samples were processed using the Illumina
- 211 TruSeq stranded mRNA LT sample prep kit RS-122-2101 (California, U.S.), and the procedure

was followed as described in the low sample protocol. The mRNA from each sample was
isolated and purified using AMPure XP magnetic beads (Agencourt; Beverly, Massachusetts)
before primary and secondary strand cDNA synthesis. Unique Illumina adapters were ligated,
and each sample was PCR amplified before validation. Samples were normalized, pooled, and
sequenced by the center for applied genetics (TCAG) facilities of the Toronto Sick Kids hospital,
Ontario, Canada.

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De novo transcript assembly, differential expression, and annotation 220

221 Raw reads were trimmed and quality checked using Trim Galore! (Version 0.4.1) (38) and 222 FastQC (Version 0.11.3) (39) then assembled using Trinity (Version 2.2.0) (40). Corset (Version 223 1.0.6) (41) was used to estimate contig abundance by grouping contigs into representative gene 224 clusters as the first step of the differential expression analysis. Contigs are defined as continuous 225 overlapping paired-end reads. Next, edgeR (Version 3.6.2) (42,43) was used to perform pairwise 226 differential expression analysis of Trinity gene, Trinity contig, and Corset clusters between 227 proximal, joint, and distal regions of the same species. Genes, contigs, and clusters were 228 classified as significantly differentially expressed if log2(fold-change)> 2 and the False 229 Discovery Rate (FDR)-corrected p-value (α) < 0.05. The analyze diff expr.pl script, provided 230 with Trinity, was used to generate z-score heatmaps of all significantly differentially expressed 231 contig clustered transcripts (α) < 0.05. A z-score is used to indicate how many standard 232 deviations a value is above the mean. The transcriptomes were annotated using the Basic Local 233 Alignment Search Tool (BLAST) (44) algorithm on a local copy of both the National Center for 234 Biotechnology Information (NCBI) non-redundant protein (nr) database and The Arabidopsis 235 Information Resource (TAIR) database (45). BLASTx (E-value<10⁻¹⁰) was used to identify highly

236	similar sequences, and transcripts with the highest bit-score from the TAIR database were used
237	as representative transcripts for heatmap generation. Whole transcriptome and fruit patterning
238	heatmaps were generated using ggplot2 (46) and ggplot in R, respectively (Version 3.4.2) (47).
239	
240 241	Orthologous Clustering
242	Orthofinder (Version 1.1.8) (48) was used to match orthologous transcripts from unfiltered
243	Erucaria and Cakile transcriptomes. Orthogroups containing transcripts from both species as
244	well as top BLAST matches for fruit patterning genes of interest were used to generate heatmaps
245	For Venn diagram generation, high-throughput sequencing (HTS) (49) filtered transcripts, sorted
246	by regions, were translated to longest open reading frame (ORF) protein fasta files using
247	TransDecoder (Version 5.0.0) (50). These files were uploaded for comparison using the

248 Orthovenn webserver (51). HTS filtering was used to reduce file size due to the web server

249 upload limit, and to reduce the number of insubstantial transcripts.

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251 Gene ontology

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253 Transcriptome fasta files from *Erucaria* and *Cakile* were imported to BLAST2GO (Version 2.8)

254 (52). Annotation files were exported and filtered to generate gene ontology (GO) terms for each

region and species. These GO terms were used to produce graphs containing transcriptome hits

256 for chosen terms. Terms were chosen based on searches for lignin, abscission, dehiscence,

257 specific hormone keywords, and top hits. For comparison between transcriptomes, the log2 of

258 selected GO term counts were divided over the log2 of all GO term counts (log2(n)/log2(N)).

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261 **Results**

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263 **De novo Assembly of** *Erucaria* and *Cakile* **Transcriptome Data** 264

265 RNA-seq libraries were constructed from 9 total replicates of triplicate distal, proximal, and joint

- regions. RNA samples from segmented fruits of two distinct plants were combined before
- sequencing to achieve optimal yield for library preparation. Sequencing from both libraries
- averaged 27.41 and 29.41 million paired-reads for *Erucaria* and *Cakile*, respectively. After
- quality trimming read counts were reduced to 27.36 million and 28.36 million high quality reads,
- 270 respectively. Inter-quartile ranges per base were minimally 33 for *Erucaria* for the first 5 base
- pairs, and minimally 32 in the 90th percentile; *Cakile's* inter-quartile ranges were minimally 33
- 272 for the first 5 base pairs, and minimally 29 in the 90^{th} percentile.
- 273

The transcriptome from *Erucaria* had an average contig length of 942.83, and *Cakile's* had an average length of 877.15. The total transcript count for *Erucaria* and Cakile was 227,530 and 314,194 reads, respectively (Table 1). Corset cluster counts averaged 365,257 (*Erucaria*) and 436,177 (*Cakile*). Notably, the first replicate for *Cakile* had a read count of 269,732, which is minimally 130,000 fewer than replicate 2 and 3. This inconsistency may have caused some issues in downstream analyses, but overall, both transcriptomes were of adequate quality and read-depth.

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284 Table 1. Statistics for de novo Trinity assembly of *Erucaria erucarioides* and *Cakile lanceolata*

285 pairwise reads.

All (Longest Isoform)	Erucaria	Cakile
N50	1544 (1017)	1464 (835)
Median Contig Length	578 (374)	517 (330)
Average Contig length	942.83 (656.94)	877.15 (577.55)
Total Assembled bases	214,521,562 (92,098,767)	275,595,508 (108,815,069)
Total Trinity Genes	140194	184945
Total Trinity Transcripts	227530	314194
GC%	41.89	42.05

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289 Annotation of Assembled Transcripts

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Both transcriptomes were compared to the nr and TAIR peptide database using a BLASTx

algorithm, and all downstream analyses used the TAIR10 annotation for facilitated comparison

to Arabidopsis. A total of 254,592 (*Cakile*) and 213,757 (*Erucaria*) transcripts with e-values≤

294 10⁻⁵ were matched to the TAIR10 database with multiple transcripts matches per gene. The GO

analysis averaged 8,644 and 8,941 terms for *Erucaria* and *Cakile*, respectively. The top 15 GO

terms consisted of 11 cellular component, three molecular function, and one biological process.

- 297 Nucleus, plasma membrane, and protein binding were the top three terms, all of which are
- biological processes (Fig S1).

300	The majority of selected orthogroups were similar between and within species (lignin,
301	abscission, and dehiscence processes, and hormone response) (Fig 3). Exceptions include: cell
302	wall modification related to abscission, general abscission, and catabolic lignification. Cakile has
303	a greater ratio of cell wall modification processes and a lower ratio of general abscission
304	processes relative to Erucaria. Erucaria has a higher ratio of catabolic lignification processes in
305	the joint region despite having similar ratios relative to Cakile in the distal and proximal regions
306	(Fig 3). Overall, the GO analysis results are consistent between and within species.
307	
308	Figure 3. Graph of select Gene Ontology (GO) terms for <i>Erucaria erucarioides</i> and <i>Cakile</i>
309	lanceolata. Sample(n) and total(N) raw counts log2 transformed for interspecies comparison. GO
310	terms chosen based on search terms: lignin, abscission, dehiscence, and response to hormone.
311	
312	Additional results from OrthoVenn showed minimal difference in orthologous clustering within
313	species, but some differences between species (Fig 4). There are a greater number of shared
314	clusters between the proximal and distal regions in Erucaria (2548) than Cakile (2306) despite
315	<i>Cakile</i> having substantially more overall clusters than <i>Erucaria</i> (50,003 vs 32,757). Additionally,
316	there are fewer clusters unique to the joint for Cakile (21) than Erucaria (112). In sum, there are
317	fewer orthologous clusters in common within regions of Cakile fruits than within regions of
318	Erucaria fruits.
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323	Figure 4. Venn diagrams of three-way and pairwise High Throughput Sequencing (HTS) filtered
324	transcripts for Erucaria erucarioides and Cakile lanceolata transcriptomes. (A), Three-way
325	Venn diagrams of Erucaria and Cakile orthologous clusters for distal, joint, and proximal
326	regions. (B), Pairwise Venn diagrams of Erucaria and Cakile orthologue-clustered transcripts
327	(Erucaria region vs Cakile region).
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330	Cakile shares a greater number of representative transcripts from the valve margin pathway with
331	Erucaria (12) than Erucaria does with Cakile (7), i.e., more representative BLAST transcripts
332	from Cakile are orthologous with transcripts from the Erucaria transcriptome than vice versa. Of
333	the representative valve margin gene transcripts, only two are orthologous between both species,
334	ASYMETRIC LEAVES 2 (AS2) and SHP2. (Fig 5 and 6). Representative transcripts are those with
335	the highest bit-score after a BLAST search against the TAIR database.
336	
337	Figure 5. Heatmap of edgeR contig clustered transcripts from <i>Erucaria erucarioides</i> expressed
338	in log2 TPM with TMM normalization. Representative transcripts based off largest bitscore hit
339	against TAIR database. Bolding indicates shared orthogroup with Cakile lanceolata. TPM,
340	Transcripts Per Million; TMM, Trimmed Mean of M-values. FULa,b,c,d are copies of FUL that
341	are present in some species across the Brassicaceae (72). Asterisks indicate significant
342	differential expression between proximal and joint region (FDR-corrected α =0.01)
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346	Figure 6. Heatmap of edgeR contig clustered transcripts from <i>Cakile lanceolata</i> expressed in
347	log2 TPM with TMM normalization. Representative transcripts based off largest bitscore hit
348	against TAIR database. Bolding indicates shared orthogroup with Cakile lanceolata. TPM,
349	Transcripts Per Million; TMM, Trimmed Mean of M-values. FULa,b,c,d are copies of FUL that
350	are present in some species across the Brassicaceae (72). Asterisks indicate significant differential
351	expression between distal and joint region (FDR-corrected α =0.05)
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Identification of Differentially Expressed Transcripts in 10mm fruit 356

357 For whole transcriptome comparison, two heatmaps of significant pairwise differentially 358 expressed transcripts ($\alpha = 0.01$) were generated (Figs 7 and 8). Contig clustering was chosen for 359 this analysis because it is a more conservative estimation of significant differential expression at 360 the transcript level, i.e., there are a greater number of transcripts being compared with more 361 stringent FDR correction relative to corset clustering. Values were then converted to z-score to 362 facilitate interspecies comparison, and for visual clarity. The joint and proximal regions of 363 Erucaria are most alike in expression and are both dissimilar to the distal region (Fig 7). All 364 three regions in *Cakile* have different expression patterns, and the distal region has a relatively 365 large inter-replicate variance (Fig 8). There are 15,345 (*Erucaria*) and 74 (*Cakile*) significantly 366 differentially expressed (SDE) transcripts in each transcriptome. There were no SDE Cakile 367 transcripts with FDR-adjusted p-values < 0.01. The low number of SDE genes between *Cakile* 368 regions indicates a lack of regional distinction in terms of transcript expression. These data 369 demonstrate a large difference in significant differential expression between the distal region

370 relative to the joint and proximal region in *Erucaria*, and little significant variation between all371 three *Cakile* regions.

372

373 Figure 7. Heatmap of all significant edgeR contig clustered transcripts in *Erucaria erucarioides*,

374 expressed as z-scores (FDR-corrected α =0.01). Row and column dendrograms indicate clustering

of transcripts (n=15,345) and biological replicates (n=3 per region), respectively.

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377 Figure 8. Heatmap of all significant edgeR contig clustered transcripts in *Cakile lanceolata*,

378 expressed as z-scores (FDR-corrected α =0.01). Row and column dendrograms indicate clustering

379 of transcripts (n=74) and biological replicates (n=3 per region), respectively.

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382 We compared expression profiles of 21 genes important for valve margin formation and 383 positioning in Arabidopsis (2,11,16,53–64) (Fig 5 and 6). Contig clustered transcripts were also 384 chosen for this analysis based on matches against the TAIR database. Most fruit patterning genes 385 for both species have no significant differences in expression across all regions, except for FIL 386 and YAB3 which were significantly upregulated in the distal region relative to the joint in 387 *Erucaria*, and *IND* which is significantly upregulated in the joint relative to both the distal and 388 proximal regions in *Cakile*. Upstream regulators *FIL* and *YAB3* are not expressed in late stage 389 *Cakile* fruits, despite global expression in *Erucaria* fruits. Downstream regulator *IND* is 390 expressed in the whole fruit in *Erucaria*, but only in the joint region of *Cakile* (Fig 5 and 6). 391

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393 **Discussion**

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Gene ontology of heteroarthrocarpic fruits 396

397 Overall, GO terms within fruits and between species are similar (Fig 3 and S1), as expected,

398 because all sections and replicates are from developing fruit with shared components (e.g., ovary

399 wall, septum). Because dehiscence is susceptible to misexpression and loss of function mutations

400 in the valve margin pathway (21,24–28), broad changes in gene ontology are unnecessary to

401 explain heteroarthrocarpy. Additionally, GO analyses of top terms do not usually vary between

402 closely related species (65,66). However, similarities in gene ontology do not imply similarity

403 between all expressed transcripts, so variation of just a few transcripts may still be the driving

404 factor behind heteroarthrocarpy.

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406 Global transcript expression of heteroarthrocarpic fruits are 407 consistent with anatomy

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409 Transcript expression patterns are consistent with anatomical variances within and between 410 fruits. The distal region of *Erucaria* has opposing transcript expression relative to both its joint 411 and proximal regions (Fig 7), i.e., when transcripts are upregulated distally in *Erucaria* they are 412 downregulated proximally. This pattern is consistent with heteroarthrocarpic fruit anatomy, as 413 distal regions contain no valve or valve margin, and proximal regions have both (36). In contrast, 414 all regions of *Cakile* have variable transcript expression, with the clearest distinction between the 415 proximal and joint regions, i.e., when genes are upregulated proximally they will be 416 downregulated in the joint (Fig 8). As with *Erucaria*, expression profiles in *Cakile* vary in a

417 manner consistent with anatomy. Superficially, one would expect the *Cakile* silique to have 418 similar expression between all regions because the entire fruit is indehiscent, however the distal 419 region of *Cakile* is more like the distal region of *Erucaria* than to its own proximal region (36). 420 Further, its abscising joint is anatomically reminiscent to a valve margin (36). Abscission zones 421 are also found between septum and seeds, and they too share similar anatomy and expression to 422 typical silique valve margins (67). Heteroarthrocarpic distal regions are unlike indehiscent non-423 heteroarthrocarpic siliques such as *L. appelianum*, because heteroarthrocarpic distal regions have 424 no remnant valve margin in contrast to indehiscence observed in *Lepidium* and the proximal 425 region of *Cakile* (32,36). Thus, we expect different expression patterns within heteroarthrocarpic 426 fruits, as well as between heteroarthrocarpic and non-heteroarthrocarpic fruits. In summary, there 427 is a clear difference between distal and proximal expression profiles for both *Erucaria* and 428 *Cakile*, which is consistent with a repositioning of the valve margin, i.e., the distal region is quite 429 distinct from the proximal region due to the lack of valve margin, or its remnant, in the distal 430 region. This consistency is further explored by analysis of fruit patterning transcript expression 431 involved in valve margin formation. 432 433 434 435 436 Fruit patterning genes 437 438 439 Despite the substantial differences in anatomy, most valve margin genes reveal similar

440 expression patterns across fruits in both *Erucaria* and *Cakile* (Fig 5 and 6). These differences are

441 initially surprising because previous studies showed variation in expression patterns across fruits

442 with in-situ hybridization (10). *EeFUL1*, one of two *FUL* homologs found in *Erucaria*, was 443 previously shown to only be expressed in the proximal region in earlier stages of carpel 444 development (10), but all FUL transcripts are expressed across all regions in this study of later 445 stage development (Fig 5). This discrepancy may be due to dynamic gene expression at different 446 stages or because our methodology cannot distinguish within region differences (e.g., genes 447 expressed in valve but not replum), so differences within regions cannot be distinguished. In 448 contrast to *EeFUL1*, our data are consistent with a previous publication which demonstrated that 449 other fruit patterning genes have broader expression domains than found in Arabidopsis (10). 450 *EeALC* and *EeIND* and *ClALC* were expressed in the septum of *Erucaria* and *Cakile*. 451 respectively, which is found throughout all regions sampled in this study. The replum is also 452 found throughout all sampled regions of *Erucaria* and *Cakile*, so expression patterns of 453 pleiotropic genes, e.g., AP2, show broader expression patterns than expected in valve and valve 454 margin alone. 455

456 It is a compelling finding that upstream regulators FIL/YAB3 and JAG have variable expression 457 across *Erucaria* (Fig 5). These three genes positively regulate expression of *FUL* and valve 458 margin genes in Arabidopsis such that their cooperative function has been designated together as 459 JAG/FIL activity (12). Our data suggest a decoupling of this cooperation in heteroarthrocarpic 460 fruits because these three genes do not exhibit the same expression patterns across Erucaria 461 fruits (Fig 5 and 6). That is, no expression of JAG was detected in any region of either species at 462 this stage. FIL and YAB3 showed differently expression patterns across fruits of Erucaria, but 463 neither were detected in *Cakile*. It is important to note that the double mutant of *fillyab3* in 464 Arabidopsis have fruits that are remarkably reminiscent of heteroarthrocarpy: they lack valve 465 margin in the distal region of fruit while maintaining ovary wall identity (11). In contrast to

466 heteroarthrocarpy, these mutants have ectopic valve margin in the proximal region of their fruits 467 (11). As these genes exhibit different patterns across *Cakile* and *Erucaria* and are expressed in 468 both proximal and distal regions of *Erucaria*, heteroarthrocarpy cannot be explained by a simple 469 lack of expression of these key regulators. Further, FIL/YAB are absent in the joint region of 470 *Erucaria* (Fig 5), which is confounding since the joint contains small portions of both proximal 471 and distal regions, an unavoidable consequence of segmentation during tissue collection. 472 Nonetheless, deviation in expression patterns of these upstream regulators between Arabidopsis 473 and heteroarthrocarpic fruits implicates variation in their expression profiles in the origin of 474 heteroarthrocarpy. 475 476 When exploring heteroarthrocarpy, we need to consider fruit patterning beyond the basal-apical 477 differences that distinguish distal, joint, and proximal regions. That is, the lateral (valve and 478 valve margin) and medial (replum) patterning is maintained in heteroarthrocarpic fruits whereas 479 the apical-basal is not. In other words, not only is replum tissue present in distal, joint, and 480 proximal regions of heteroarthrocarpic fruits, it is appropriately sized. FIL/YAB3 and JAG 481 function antagonistically with replum promoting gene, WUSCHEL RELATED HOMEOBOX 13 482 (WOX13), which positively regulates RPL in turn. This interaction is necessary for proper 483 medial-lateral formation of Arabidopsis fruits. Further, ASYMMETRIC LEAVES1 (ASI) and AS2 484 collaborate with JAG/FIL function as promoters of lateral factors (12). The loss of both AS1/2 485 and JAG/FIL in Arabidopsis results in dramatic medial-lateral differences and substantially 486 enlarged repla, which is interestingly more pronounced in the basal portion of the fruit (12,68). 487 As AS1/2 and AS1 are expressed throughout Cakile and Erucaria regions, respectively, this 488 pattern suggests that AS1 alone is sufficient for proper replum (aka medial-lateral) formation in 489 heteroarthrocarpic fruits. In other words, the collaboration between JAG/FIL function and AS1/2

490 is not maintained in heteroarthrocarpic fruits. Further, in at least *Cakile JAG/FIL* activity is non491 detectable in the entire fruit, at least at later stages of development. Thus, it appears that some
492 redundancy in lateral-medial patterning of Arabidopsis fruits has been lost in heteroarthrocarpic
493 fruits, while simultaneously gaining apical-basal differences.

494

495 Valve margin pathway recruitment and abscission in the Cakile 496 joint. 497

498 The fruit of *Cakile* is distinct in that the joint abscises (disarticulates) at maturity. The joint, 499 which represents the distal portion of the valve margin, thus represents a novel abscission zone in 500 *Cakile*, completely separating the distal portion of the fruit. This is an unusual feature of certain 501 heteroarthrocarpic subtypes, as there is no equivalent abscission zone in Arabidopsis. Our data 502 strongly implicate the recruitment of downstream valve margin genes as responsible for joint 503 abscission, although how that zone is positioned remains elusive. *IND* is significantly 504 upregulated in joint region (Fig 6) and is primarily responsible for formation of separation and 505 lignification layers in typical siliques (24,26), a juxtaposition of cell types also observed in the 506 abscising joint region. Its presence in the joint may be due to a co-option of downstream valve 507 margin pathway genes to facilitate formation of the joint abscission zone. Similar co-option is 508 observed in seed abscission zones, although these zones typically involve SEEDSTICK (STK) in 509 lieu of SHP, and the functionally similar transcription factor HEC3 in lieu of IND (67). SHP1/2 510 and ALC expression are both consistent with this co-option, as they are expressed in all three 511 regions (Fig 6). Additionally, SPT expression is consistent with expression of IND, as expected 512 from its downstream role in valve margin formation. (Fig 6) (14). Further, both representative 513 transcripts are among the 21 unique orthologous clusters in the joint of *Cakile* (Fig 4). This

514 pattern is consistent with in situ hybridization data that showed SHP2 expressed in septum and 515 ovules of Cakile, and in ovules of Erucaria (10). Thus, the likely function of SHP1/2 and ALC in 516 the joint region would be to promote expression of IND (SHP1/2), and the formation of the 517 separation layer (ALC). What is unusual about joint abscission is that for the joint to separate, the 518 distal and proximal regions of the replum must also separate. This expression pattern then 519 implies that the mechanism used to physically separate valve from replum may also be in play 520 for replum in the joint region. Taken together with anatomical studies, our data strongly suggests 521 that there is a repurposing of the valve margin pathway in an otherwise indehiscent *Cakile* fruit, 522 and that this pathway may be capable of initializing disarticulation in multiple tissue types. 523

524 Conclusion

525

526 Transcriptomic expression from late stage *Erucaria* and *Cakile* fruits is consistent with some 527 conservation and some deviation of the valve margin pathway, specifically in upstream 528 regulation, e.g., YAB/FIL/JAG. Thus, different upstream regulators are implicated in the loss of 529 dehiscence in Brassiceae relative to *Lepidium*, where AP2 is likely responsible (32). Loss of 530 expression of YAB/FIL/JAG in Arabidopsis results in differing apical and basal phenotypes. 531 which may help to explain the apical/basal differences in heteroarthrocarpic fruits (11). Further, 532 heteroarthrocarpic fruits likely recruit the same mechanism used in valve and seed abscission for 533 joint abscission (Fig 6). Functional tests are necessary to confirm whether redeployment of 534 FIL/YAB3, IND, and possibly SPT have key roles in the origin of heteroarthrocarpy as well as 535 joint abscission.

537	There have been multiple whole genome duplications in the Brassicales, which has resulted in
538	many polyploids within the Brassicaceae family (69–71). We considered the possibility of
539	transcriptional differences between gene copies in distal, joint, and proximal regions that were
540	undetected because we were unable to determine copy number in our transcriptome. For
541	example, there are four copies of FUL in the Brassiceae (72), but each potential FUL copy had
542	multiple hits from the same transcripts in both transcriptomes, so there is no definitive answer
543	about copy number and expression (Fig 5 and 6). That is, we could not confirm or refute
544	subfunctionalization of some fruit patterning genes as having a role in the origin of
545	heteroarthrocarpy. An analysis of multiple transcripts for every fruit patterning gene showed
546	generally similar expression for each, but further analyses are needed to determine if
547	neo/subfunctionalization plays a role in heteroarthrocarpy.
548	
549	Understanding the nature of heteroarthrocarpy, and how it relates to fruit development in
550	Arabidopsis, will facilitate future studies on seed shattering in important Brassicaceous crops,
551	and pernicious heteroarthrocarpic weeds. Further, these studies inform on the origin of important
552	variation in seed packaging and dispersal capabilities.
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563

564 Author Contributions Statements

- 565 SC and JH contributed concept and project design. SC and KM contributed to plant care, RNA
- 566 extraction and cDNA library preparation. KM designed scripts and was lead in initial
- 567 bioinformatic analyses; SC completed later analyses using scripts produced by KM. SC wrote
- the first manuscript draft; SC and JH wrote subsequent manuscript drafts. All authors contributed
- to revision and proofreading of the final submitted version.

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571 Conflict of interest statement

572 There are no conflicts of interest to report.

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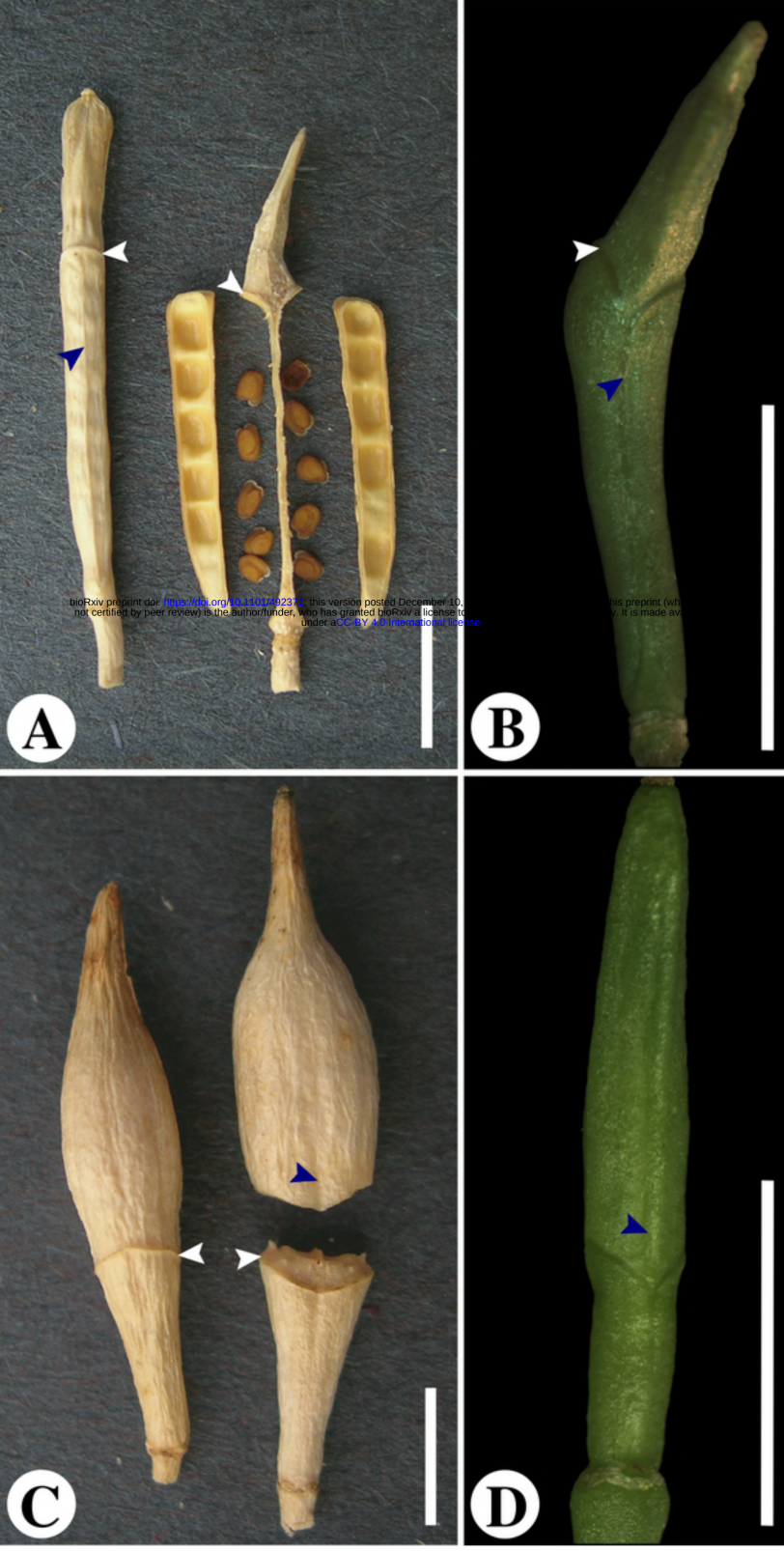
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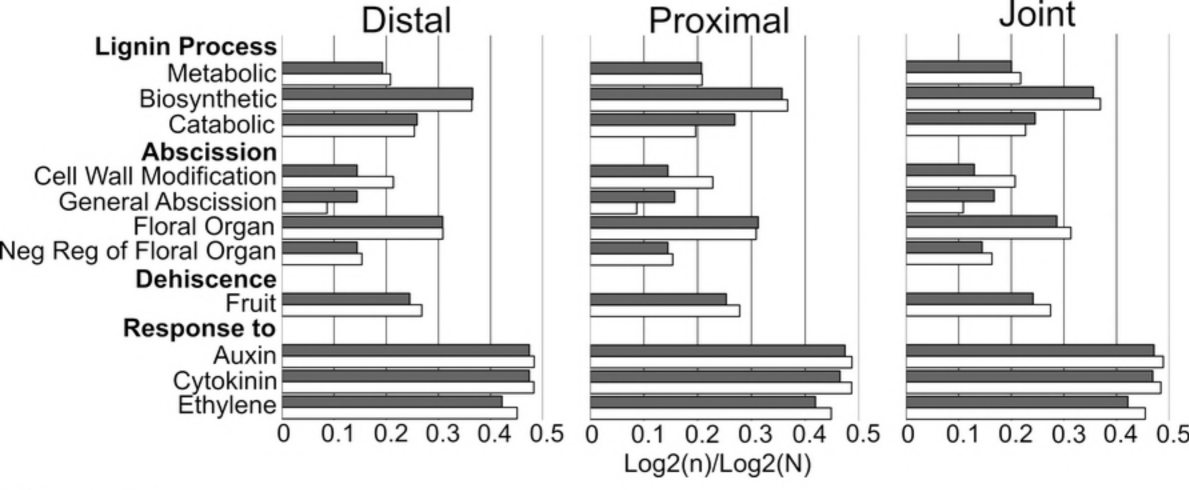
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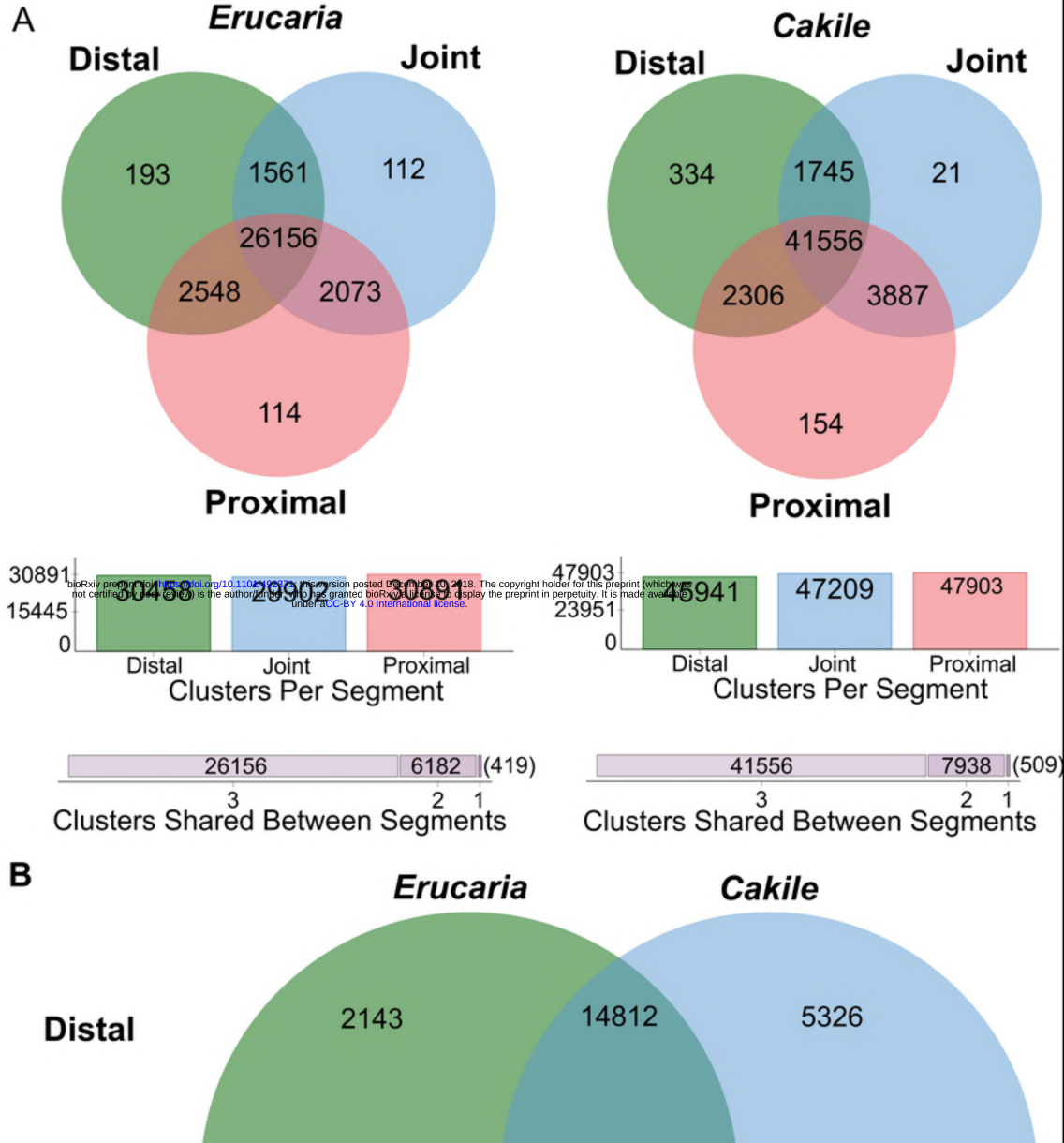
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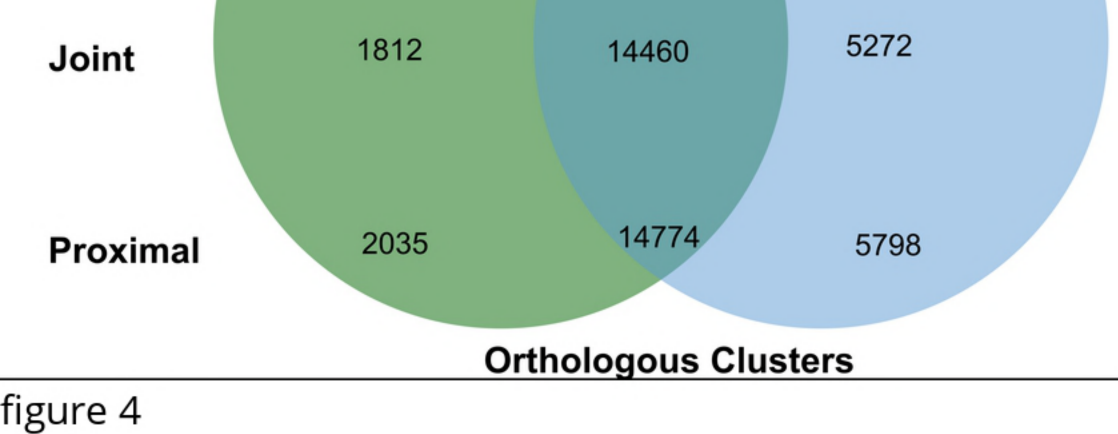
- Figure S1. Graph of top Gene Ontology (GO) terms for *Erucaria erucarioides* and *Cakile*
- *lanceolata*. Sample(n) and total(N) raw counts were log2 transformed for interspecies
- comparison.

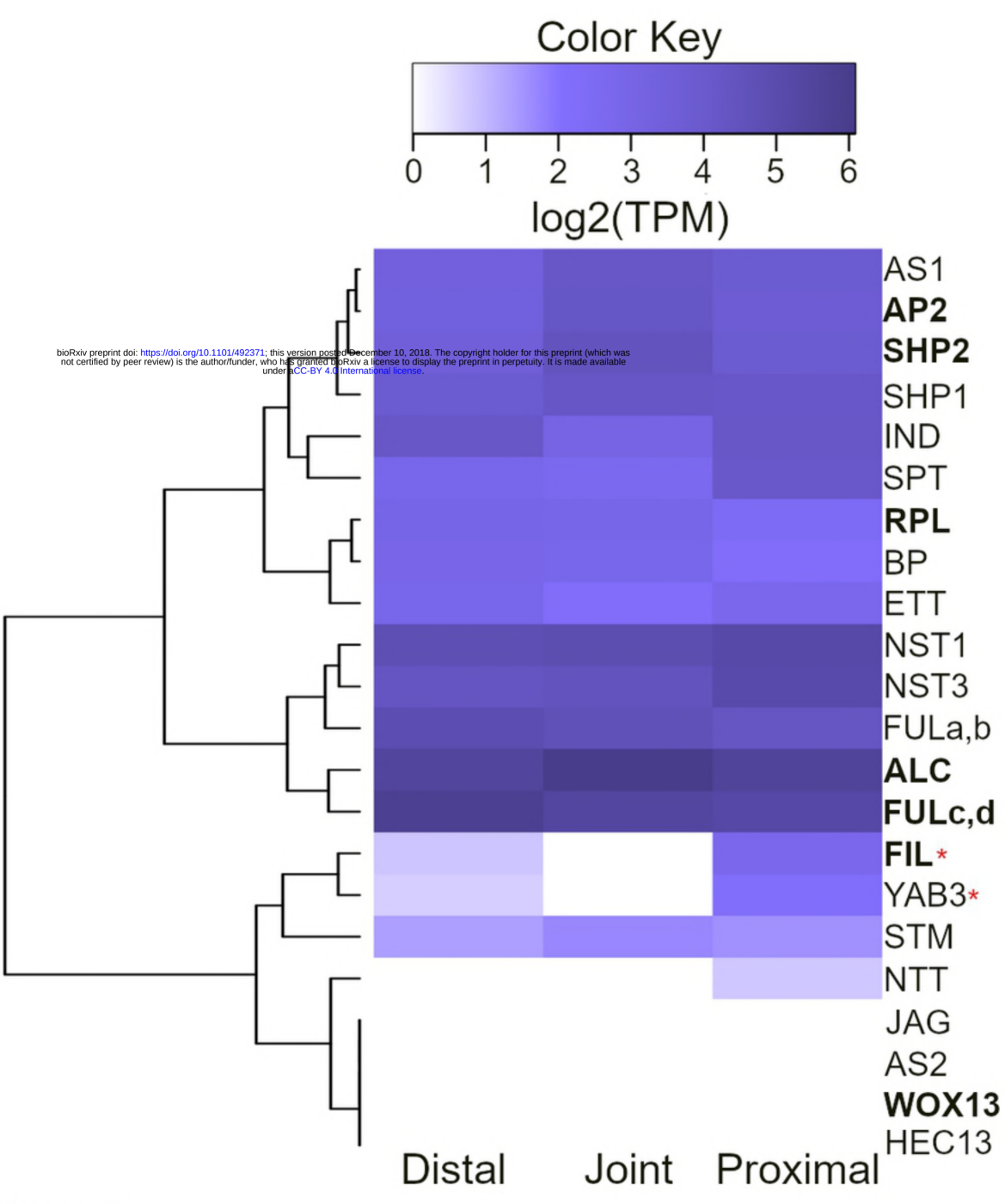


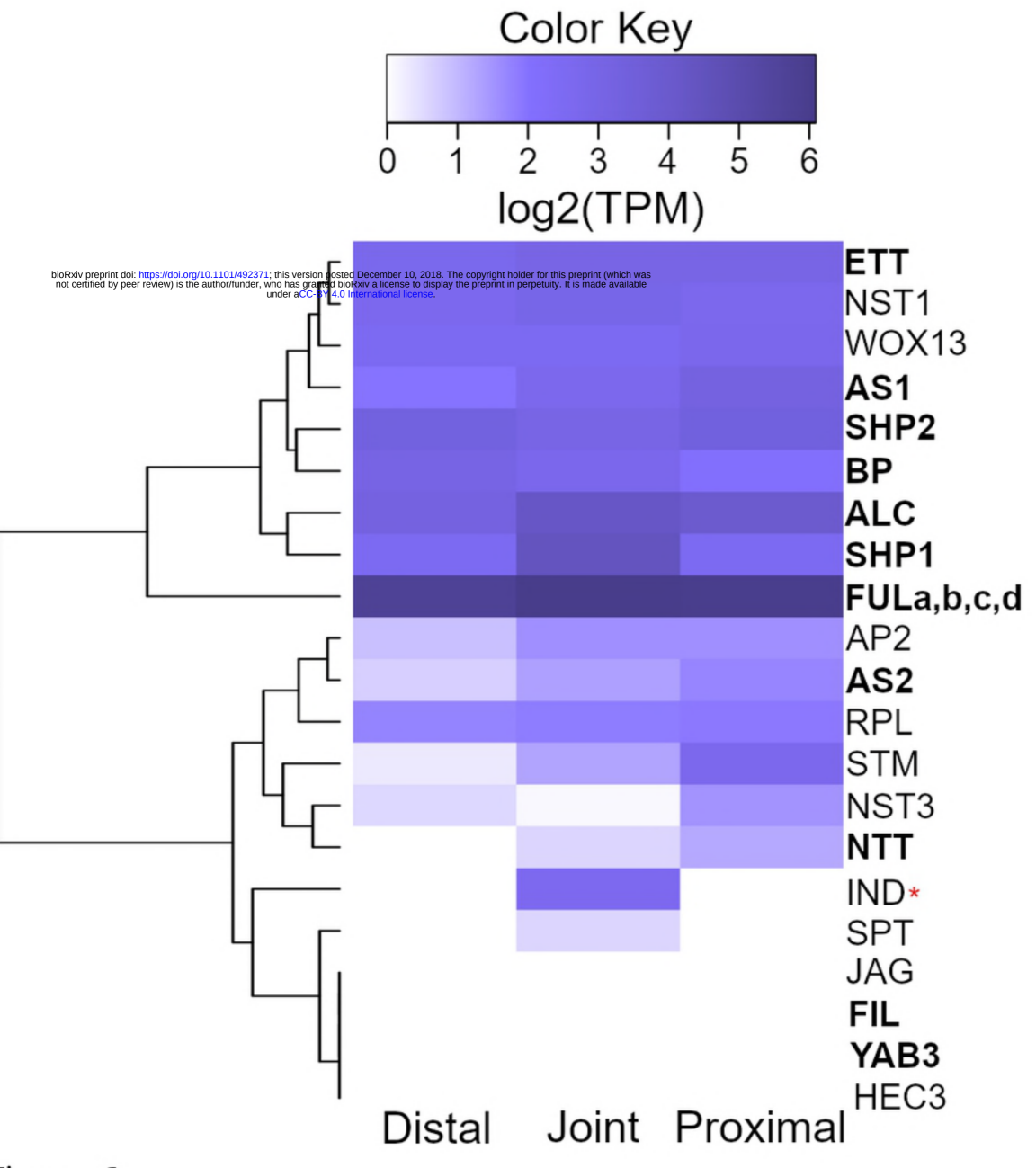


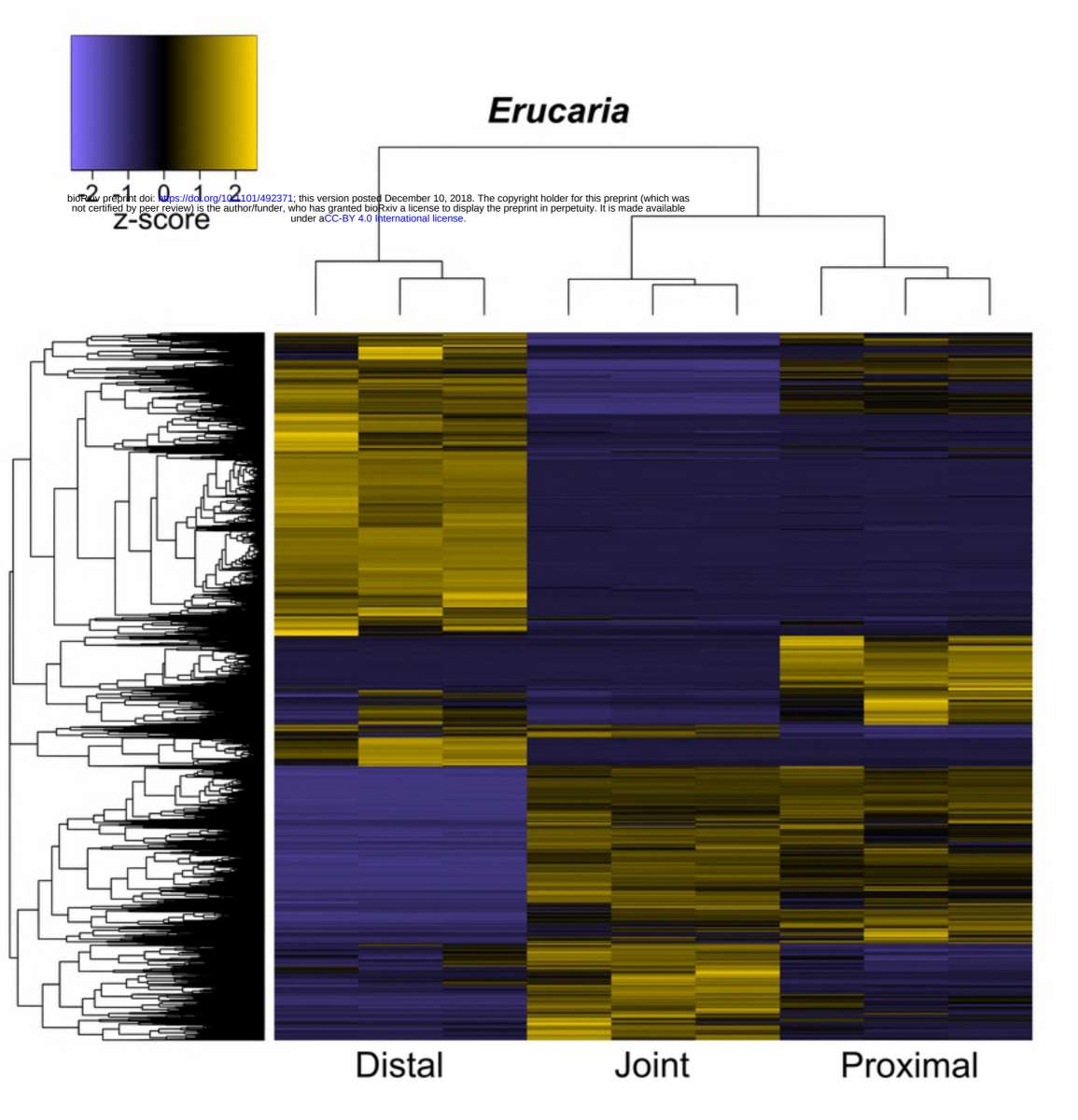












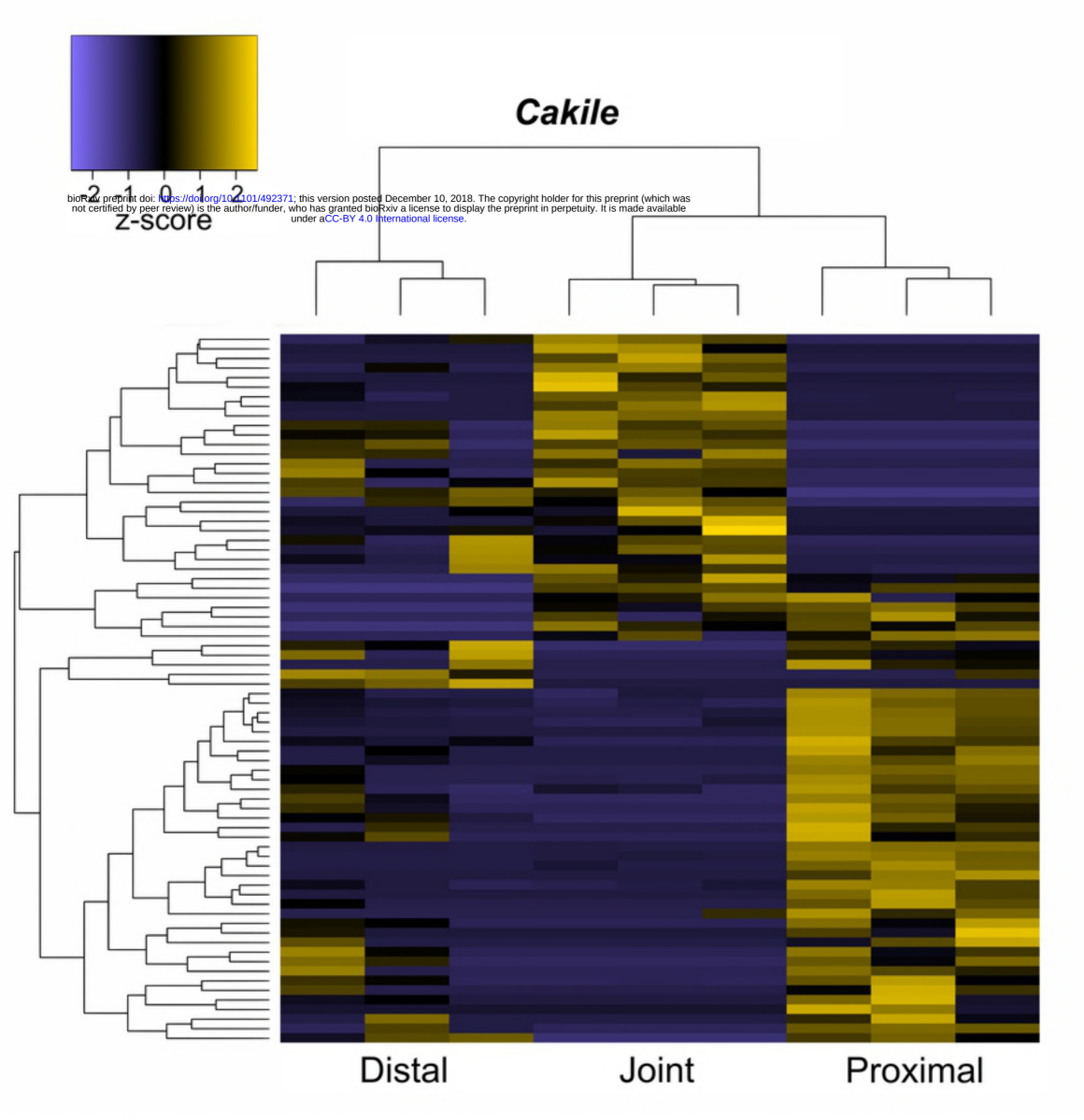






Fig 1 , Hall, Int. J. Plant.Sci 2006

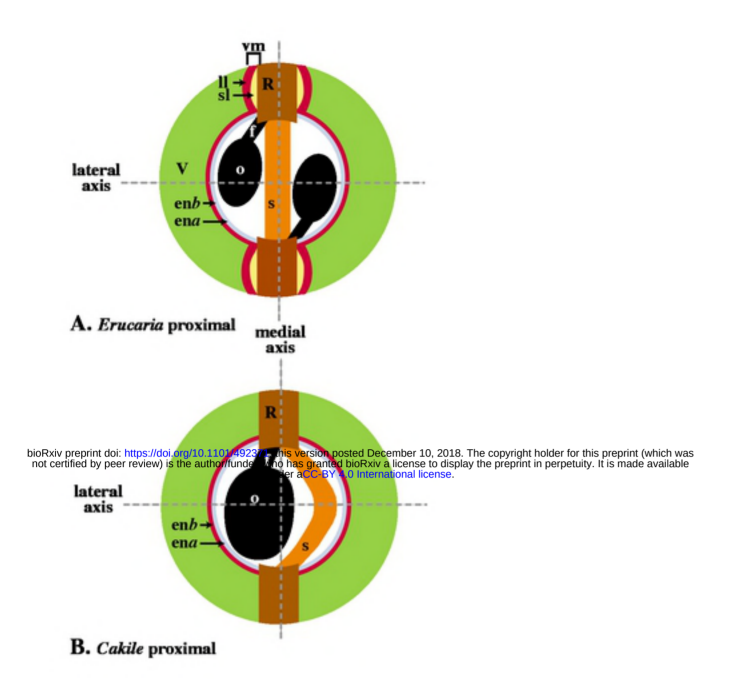


Fig 2 , Hall, Int. J. Plant.Sci 2006

