Comparative transcriptomics identifies candidate genes involved in the evolutionary transition from dehiscent to indehiscent fruits in *Lepidium* (Brassicaceae)

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**Abstract**

**Background:** Fruits are the seed-bearing structures of flowering plants and are highly diverse in terms of morphology, texture and maturation. Dehiscent fruits split open upon maturation to discharge their seeds while indehiscent fruits are dispersed as a whole. Indehiscent fruits evolved from dehiscent fruits several times independently in the crucifer family (Brassicaceae). The fruits of *Lepidium appelianum*, for example, are indehiscent while the fruits of the closely related *L. campestre* are dehiscent. Here, we investigate the molecular and genetic mechanisms underlying the evolutionary transition from dehiscent to indehiscent fruits using these two *Lepidium* species as model system.

**Results:** We have sequenced the transcriptomes and small RNAs of floral buds, flowers and fruits of *L. appelianum* and *L. campestre* and analyzed differentially expressed genes (DEGs) and differently differentially expressed genes (DDEGs). DEGs are genes that show significantly different transcript levels in the same structures (buds, flowers and fruits) in different species, or in different structures in the same species. DDEGs are genes for which the change in expression level between two structures is significantly different in one species than in the other. Comparing the two species, the highest number of DEGs was found in flowers, followed by fruits and floral buds while the highest number of DDEGs was found in fruits versus flowers followed by flowers versus floral buds. Several gene ontology terms related to cell wall synthesis and degradation were overrepresented in different sets of DEGs highlighting the importance of these processes for fruit opening. Furthermore, the fruit valve identity genes *FRUITFULL* and *YABBY3* were among the DEGs identified. Finally, the microRNA miR166 as well as the TCP transcription factors *BRANCHED1* (BRC1) and *TCP FAMILY TRANSCRIPTION FACTOR 4* (TCP4) were found to be DDEGs.

**Conclusions:** Our study reveals differences in gene expression between dehiscent and indehiscent fruits and uncovers miR166, BRC1 and TCP4 as candidate genes for the evolutionary transition from dehiscent to indehiscent fruits in *Lepidium*.

**Keywords:** Transcriptome, Fruit development, Dehiscence, Differentially expressed genes, Lepidium appelianum, Lepidium campestre

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Background

Flowering plants (angiosperms) form fruits to protect and disperse their seeds. Fruits come in many different types with different morphologies and different properties such as dry or fleshy, and dehiscent or indehiscent [1]. There is a tremendous variation in fruit types both across and within different plant lineages [2]. However, the evolutionary mechanisms that enabled such dramatic shifts to occur, often in a relatively short period of time, remain largely unknown.

The crucifer family (Brassicaceae) includes a number of economically important plants such as cabbage, broccoli, mustard, radish, and turnips. The model plant Arabidopsis thaliana is also a member of this family [3]. Typical fruits of Brassicaceae species are dehiscent, i.e. that the fruits open upon maturation to release the seeds. Dehiscent fruits also likely represents the ancestral fruit type of Brassicaceae [4]. However, indehiscent fruits, i.e. fruits that only release the seed upon decomposition of the fruit, are found in many tribes distributed across the Brassicaceae phylogeny [5]. The scattered distribution of indehiscent fruits indicates that this property evolved independently several times. This situation is mirrored in the genus Lepidium belonging to Brassicaceae: Species of this genus typically produce two-seeded dehiscent fruits, but the genus also includes species with indehiscent fruits [6].

Brassicaceae fruits are composed of two fruit valves that are connected to the replum and enclose the developing seeds. Dehiscent fruits, such as those of A. thaliana and Lepidium campestre (also known as field pepperwort or field cress), form a well-defined dehiscence zone (DZ) at the valve margin [7]. The DZ consists of the lignified layer, a stripe of lignified cells, and a separation layer, a region of small thin-walled cells [8, 9]. During fruit ripening, the whole fruit dries and shrinks. Only the lignified structures stay rigid. Thereby a spring-like tension is created within the fruit. At the same time, the middle lamellae of the separation layer cells degenerate to form a pre-determined breaking zone at which the pressure tears the valves apart from the replum. Consequently, the fruit bursts open to release the seeds [9–11]. In contrast, the indehiscent fruits of the closely related Lepidium appelianum do not form a DZ. Instead, a continuous ring of lignified cells surrounds the seeds such that the fruit cannot open [7].

Much of the gene regulatory network underlying the proper formation of the fruit valves, replum and DZ has been elucidated in A. thaliana (reviewed in Ballester and Ferrándiz [12]). Establishment of the DZ requires expression of the two redundant MADS-box genes, SHATTERPROOF1 (SHP1) and SHATTERPROOF2 (SHP2). The SHP1 and SHP2 proteins act as transcription factors and activate the basic helix-loop-helix protein-encoding genes INDEHISCENT (IND), ALCATRAZ (ALC) and SPATULA (SPT), and also autonomously contribute to DZ development [8, 13–15].

For correct fruit patterning, it is crucial that the expression of the SHP genes is restricted to the DZ. Three genes encoding transcription factors contribute to this process: The MADS box gene FRUITFULL (FUL) which is expressed in the fruit valves [16, 17], the BELL1-like homeobox gene REPLIMLESS (RPL) [18], also known as PENNYWISE [19], BELLRINGER [20], VAAMANA [21], and BLH9 [22] which is expressed in the replum, and the floral homeotic gene APETALA2 (AP2) which negatively regulates RPL [23].

Transcription factors controlling the expression of these regulators have also been determined. High levels of the C2H2 zinc finger proteins JAGGED (JAG) and the two closely related YABBY1 group proteins FILAMENTOUS FLOWER (FIL) and YABBY3 (YAB3) activate the expression of FUL [24]. In contrast, lower levels of JAG/FIL/YAB3 expression promote expression of SHP genes. The expression of RPL is activated by the KNOTTED1-like homeobox protein BREVIPEDICELLUS (BP) [25] whose gene is in turn activated by the C2H2 zinc finger protein NO TRANSMITTING TRACT (NTT) [26]. AP2 is negatively regulated by the microRNA miR172 [27].

Additionally, other factors which influence the size and the position of the DZ have been identified. The WUSCHEL-RELATED HOMEOBOX gene 13 (WOX13) controls replum width and negatively regulates JAG/FIL/YAB3 [28]. The auxin-response factors ARF6 and ARF8, which are regulated by miR167 [29], activate miR172 together with FUL [27]. The MYB protein ASYMMETRIC LEAVES 1 (AS1), likely in collaboration with the leucine zipper protein ASYMMETRIC LEAVES 2 (AS2), negatively regulates BP [25].

In general, proteins encoded by genes expressed in the replum often negatively regulate genes expressed in the valves and vice versa. Apart from the already mentioned interactions, this includes negative regulation of the replum gene BP by the valve proteins encoded by JAG/FIL/YAB3, and negative regulation of JAG/FIL/YAB3 by the replum protein RPL [30].

In a previous study, we have shown that orthologues of the valve margin genes are expressed in a similar way in L. campestris (dehiscent fruits) as in A. thaliana fruits but that expression of the respective orthologues is abolished in the corresponding tissues of indehiscent Lepidium appelianum fruits [7]. As parallel mutations in different genes are unlikely, we concluded that the changes in gene expression patterns are probably caused by changes in upstream regulators such as FUL, RPL or AP2.
To conduct a more unbiased approach to identify the genetic changes that lead from dehiscent to indehiscent fruits than the analysis of candidate genes, we have sequenced the transcriptomes of floral buds, flowers and fruits of both, *L. campestre* and *L. appelianum* in the present study. We have identified differentially expressed genes (DEGs) and differently differentially expressed genes (DDEGs) where the latter refers to genes for which the change in expression level between two structures is significantly different in one species than in the other. More DEGs were identified in flowers than in fruits and floral buds and a higher number of DDEGs was found in fruits versus flowers than in flowers versus floral buds. Cell wall synthesis and degradation are important processes for fruit opening as revealed by gene ontology (GO) analysis. The fruit valve identity genes *FRUITFULL* and *YABBY3* were identified as DEGs such that the possible cause for the evolutionary transition from dehiscent to indehiscent fruits in *Lepidium* may even be an upstream factor of these genes. Possible candidates are *BRANCHED1* (*BRC1*), an ortholog of which may determine whether dehiscent or indehiscent fruits develop on the dimorphic plant *Aethionema arabicum*, and *TCP FAMILY TRANSCRIPTION FACTOR 4* (*TCP4*) which may regulate *YABBY3*. These two genes were found to be DDEGs. Our study elucidates differences in gene expression patterns between dehiscent and indehiscent fruits and reveals *BRC1* and *TCP4* as possible causes for the evolutionary transition from dehiscent to indehiscent fruits in *Lepidium*.

**Results**

**Overview of the RNA-seq analysis and transcriptome assembly**

Sequencing resulted in an average number of reads per library of 56 Mio. for the mRNA and 12 Mio. for the small RNA (Table 1). An initial analysis of the data revealed contamination with sequences from thrips, likely due to infestation of the plants by these insects. Hence, we removed reads matching to the genome of the thrips *Frankliniella occidentalis* [31] as well as uncorrectable and unpaired reads and reads corresponding to organelle sequences. After this filtering step, 42 Mio. reads were retained for further analyses for the mRNA sample. For the small RNA sample many reads seem to be derived from organelle RNA. Hence, after removing uncorrectable reads and those matching to the *Frankliniella occidentalis* genome and organelle sequences, only 1.5 Mio. reads remained on average for the small RNA sample (Table 1).

Assembly using Trinity [32] resulted in a total of 56,413 transcripts for *L. campestre* and 70,380 transcripts for *L. appelianum* after removing putative contaminant sequences but including potential splice variants or chimeric sequences. The assemblies also contained chimeric sequences composed of two different transcripts which were likely a result of mis-assembly [33]. Separation of chimeric sequences increased the number of transcripts to a total of 57,209 for *L. campestre* and 71,332 for *L. appelianum*. We used the Benchmarking Universal Single-Copy Orthologs (BUSCO) tool [34] with the dataset eudicotyledons_odb10 as reference to assess completeness of our transcriptomes. The BUSCO analyses revealed that 94.6% of the expected eudicotylledonous “near-universal single-copy orthologs” are present in our assembly of the *L. campestre* transcriptome while 94.3% of these BUSCOs are present in our *L. appelianum* transcriptome (Fig. 1). It is common that some genes are fragmented in de novo assemblies. Hence, we analyzed the length distribution of our assemblies. For both species there are two peaks (Fig. 2). One peak appears at a length of about 240 nucleotides (log 2 length of about 7.9) and probably represents fragments. The other peak was found at a length of about 1,450 nucleotides (log 2 length of about 10.5) which indicates that there are also a number of full-length transcripts.

To detect conserved miRNAs, we mapped the small RNA reads onto the mature miRNAs of *A. thaliana* as provided by miRBase [35]. We found reads for 64 mature miRNAs belonging to 32 miRNA families in the *L. campestre* small RNA data (Table 2). Using ShortStack [36] and the *L. campestre* genome as available from NCBI, we identified three novel miRNAs. However, no putative target genes could be identified in the transcriptome of *L. campestre* using targetfinder (https://github.com/carringtonlab/TargetFinder). Our *L. appelianum* small RNA data contained reads of 60 mature miRNAs belonging to 30 miRNA families (Table 2). No novel miRNAs could be identified for *L. appelianum* using ShortStack and our transcriptome as reference “genome”.

To assess completeness of our small RNA data, we compared our results to the set of conserved and moderately conserved miRNA families as identified by miRNA sample sequencing of vascular plants [37]. For both species, we identified reads for all 16 miRNA families that were found to have originated before the emergence of eudicots and to be conserved across virtually all corresponding species. Furthermore, we found reads for 6 miRNA families in our *L. campestre* and 7 miRNA families in our *L. appelianum* small RNA data out of 21 miRNA families which were classified as conserved, although missing in a few corresponding species.

**Differential gene expression analysis**

To conduct differential expression analysis, we identified putative ortholog pairs between the transcripts of the two *Lepidium* species. Thereby, we excluded...
ortholog pairs in which the shorter sequence was less half in length than the longer one and we kept only one transcript isoform per gene as described in the methods section. To make sure not to lose genes of interests using this conservative approach, we checked transcripts that were excluded from the ortholog transcriptome and that showed similarity to *A. thaliana* genes annotated to have “DNA-binding transcription factor activity” (GO:0003700) (Supplemental Table 1). Most of these transcripts do not seem to be involved in fruit development. The only transcripts which may have some relation to fruit development are BRANCHED2 (BRC2), MYB26, KANADI 2 (KAN2) and MYB85. BRC2 has similar but weaker effects on branching than BRC1 [38] and may have similar effects on dehiscence as will be discussed for BRC1 later. MYB26 has been shown to have a role in anther dehiscence [39] and hence a role for this TF also in fruit dehiscence is conceivable. KAN2 has been shown to repress ASYMMETRIC LEAVES2 (AS2) in *A. thaliana* [40]. AS2

| Table 1 | Number of reads obtained after sequencing and after correction and pruning steps |
|---------|---------------------------------|
| **Experiment** | **Species** | **Structure** | **Replicate** | **Raw reads** | **Uncorrectable, unpaired reads removed** | **Thrips and organelle sequences removed** |
| mRNA | *L. campestre* | Bud | 1 | 56,364,306 | 47,437,984 | 42,661,682 |
| | &nbsp; | &nbsp; | 2 | 52,626,578 | 44,103,956 | 42,345,608 |
| | &nbsp; | &nbsp; | 3 | 46,984,896 | 38,458,226 | 35,556,348 |
| | *L. appelianum* | Bud | 1 | 61,044,624 | 51,099,508 | 47,872,858 |
| | &nbsp; | &nbsp; | 2 | 57,117,360 | 45,964,748 | 41,954,354 |
| | &nbsp; | &nbsp; | 3 | 56,803,334 | 47,261,624 | 42,970,772 |
| Flower | *L. campestre* | 1 | 53,973,840 | 45,071,412 | 42,388,254 |
| | &nbsp; | &nbsp; | 2 | 54,176,184 | 43,661,258 | 40,639,670 |
| | &nbsp; | &nbsp; | 3 | 47,473,062 | 37,866,352 | 33,766,726 |
| Flower | *L. appelianum* | 1 | 66,540,836 | 59,074,352 | 48,284,508 |
| | &nbsp; | &nbsp; | 2 | 67,087,830 | 57,231,738 | 45,811,592 |
| | &nbsp; | &nbsp; | 3 | 57,259,526 | 48,363,464 | 41,299,370 |
| Flower | *L. campestre* | 1 | 53,973,840 | 45,071,412 | 42,388,254 |
| | &nbsp; | &nbsp; | 2 | 54,176,184 | 43,661,258 | 40,639,670 |
| | &nbsp; | &nbsp; | 3 | 47,473,062 | 37,866,352 | 33,766,726 |
| Flower | *L. appelianum* | 1 | 66,540,836 | 59,074,352 | 48,284,508 |
| | &nbsp; | &nbsp; | 2 | 67,087,830 | 57,231,738 | 45,811,592 |
| | &nbsp; | &nbsp; | 3 | 57,259,526 | 48,363,464 | 41,299,370 |
| smallRNA | *L. campestre* | Bud | 1 | 12,317,448 | 11,640,550 | 1,065,134 |
| | &nbsp; | &nbsp; | 2 | 12,888,040 | 11,434,250 | 1,234,296 |
| | &nbsp; | &nbsp; | 3 | 13,084,802 | 18,556,186 | 4,545,624 |
| | *L. appelianum* | Bud | 1 | 11,107,988 | 10,038,752 | 1,667,408 |
| | &nbsp; | &nbsp; | 2 | 12,171,462 | 10,194,186 | 430,032 |
| | &nbsp; | &nbsp; | 3 | 12,245,803 | 15,075,818 | 1,955,400 |

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itself is part of the fruit development network [25]. MYB85 has a role in lignin biosynthesis in A. thaliana [41]. As lignification is important for fruit dehiscence, MYB85 may also have a role in this process. Future transcriptome analyses with a deeper sampling may help to also include these genes in the analyses. However, in general, we do not seem to miss many factors with potential functions in fruit development.

Finally, we attained two transcriptome datasets, one for L. campestre and one for L. appelianum, each containing 17,755 transcripts and where each transcript in one species has exactly one putative orthologous transcript in the other species. We will refer to these transcriptome datasets as our ortholog-transcriptomes in the following. We reassessed completeness of our ortholog-transcriptomes and found that 89.2% of the BUSCOs remained in our ortholog-assembly for L. campestre while this value was slightly lower at 89.1% for our L. appelianum ortholog-transcriptome.

Reads were mapped independently to the corresponding ortholog-transcriptome and counted using HTSeq-count [42]. A principal component analysis was conducted based on the normalized number of reads mapping to the ortholog-transcriptomes. As expected, the replicates from the same species and structure clustered together (Fig. 3). The species are separable based on first component which explains 54% of the variance while the structures are separable based on second component which explains 30% variance (Fig. 3).

To learn more about the differences in fruit development between the L. campestre and L. appelianum, we analyzed expression in our ortholog-transcriptomes using the programs DESeq2 [43] and edgeR [44]. We used a multi-factor design to not only be able to identify differentially expressed genes (DEGs) between the
Table 2 miRNAs identified in short read data of *L. campestre* and *L. appelianum* by mapping to *A. thaliana* mature miRNAs or using ShortStack with transcriptome or genome data as reference. Highlighted in bold are conserved miRNAs according to Chavez-Montez et al., 2014 [37], bold and italic indicate moderately conserved miRNAs according to Chavez-Montez et al., 2014 [37].

| miRNA family       | *L. campestre* mature miRNA | *L. campestre* ShortStack genome | *L. campestre* ShortStack transcriptome | *L. appelianum* mature miRNA | *L. appelianum* ShortStack transcriptome |
|--------------------|------------------------------|----------------------------------|------------------------------------------|------------------------------|------------------------------------------|
| miR156a-3p/miR156c-3p | x                             | miR156c                          | x                                        | miR156a                      |                                          |
| miR156a-5p/miR156b-5p/miR156c-5p/miR156d-5p/miR156e-5p/miR156f-5p/miR156g/miR156h/miR156i/miR156j | x                             | miR156e, miR156b, miR156c            | x                                        | miR156a, miR156j                      |                                          |
| miR156b-3p         | x                             | miR156b                          |                                          |                              |                                          |
| miR156d-3p         | x                             |                                  |                                          |                              |                                          |
| miR157a-3p/miR157b-3p | x                             |                                  |                                          |                              |                                          |
| miR157a-5p/miR157b-5p/miR157c-5p/miR157d | x                             | miR157c                          |                                          |                              |                                          |
| miR157c-3p         | x                             | miR157c                          |                                          |                              |                                          |
| miR158a-3p/miR158b | x                             |                                  |                                          | miR158a                      |                                          |
| miR159a/miR159b-3p | x                             |                                  |                                          |                              |                                          |
| miR159c            | x                             |                                  |                                          |                              |                                          |
| miR160a-3p         | x                             | miR160a                          |                                          |                              |                                          |
| miR160a-5p/miR160b/miR160c-5p | x                             | miR160a, miR160c                | miR160b                                  |                              |                                          |
| miR160c-3p         | x                             | miR160c                          |                                          |                              |                                          |
| miR161.1           | x                             |                                  |                                          |                              |                                          |
| miR161.2           | x                             |                                  |                                          |                              |                                          |
| miR162a-3p/miR162b-3p | x                             | miR162b                          |                                          | miR162b                      |                                          |
| miR162a-5p/miR162b-5p | x                             | miR162b                          |                                          | miR162b                      |                                          |
| miR164a/miR164b-5p/miR164c-5p | x                             | miR164a                          |                                          |                              |                                          |
| miR164b-3p         | x                             |                                  |                                          |                              |                                          |
| miR164c-3p         | x                             |                                  |                                          |                              |                                          |
| miR165a-3p/miR165b/miR165c-3p/miR166a-3p/miR166b-3p/miR166c/miR166d/miR166e/miR166f/miR166i/miR166j | x                             | miR165b, miR166d, miR166g            | x                                        | miR166a, miR166b, miR166c, miR166f |                                          |
| miR166a-5p         | x                             |                                  |                                          | x                            |                                          |
| miR166b-5p/miR166b-3p | x                             | x                               | miR166a, miR166b                       |                              |                                          |
| miR166e-5p         | x                             | x                               | miR166e                                 |                              |                                          |
| miR167a-3p         | x                             |                                  |                                          |                              |                                          |
| miR167a-5p/miR167b/miR167d | x                             | miR167b                          |                                          |                              |                                          |
| miR167c-3p         | x                             | miR167c                          |                                          |                              |                                          |
| miR168a-3p         | x                             | miR168a                          |                                          |                              |                                          |
| miR168a-5p/miR168b-5p | x                             | miR168a                          |                                          |                              |                                          |
| miR169a-5p/miR169b-5p/miR169c | x                             | miR168a                          |                                          |                              |                                          |
| miR169d/miR169e/miR169f-5p/miR169g-5p | x                             | x                               |                                          |                              |                                          |
| miR169f-3p         | x                             |                                  |                                          |                              |                                          |
| miR170-5p/miR171a-5p | x                             | miR171a                          |                                          | miR170, miR171a              |                                          |
| miR171a-3p         | x                             | miR171a                          |                                          |                              |                                          |
| miR171b-5p/miR171c-5p | x                             | miR171b                          |                                          | miR171b, miR171c             |                                          |
| miR171b-3p/miR171c-3p | x                             | miR171b                          |                                          | miR171b, miR171c             |                                          |
species in the same structure and between structures in the same species, but also to identify genes where the change in expression between the structures is different between the two species. We will refer to the genes identified in the latter analyses as differently differentially expressed genes (DDEGs).

DESeq2 generally identified more DEGs and DDEGs than edgeR, but there is a great overlap of genes identified by both programs (Fig. 4, Supplemental Fig. 1). Only this overlap between the two methods will be considered in the following. More DEGs were observed between the same structure of the different species as compared to different structures of the same species. In L. campestre, there are similar numbers of DEGs between flower and bud as compared to fruit and flower. In L. appelianum, there are more than twice as many DEGs in flowers versus buds as compared to fruits versus flowers (Fig. 4). When looking at DEGs in the same structure of the different species, the highest number of DEGs is observed in flowers, followed by fruits and buds.

### Table 2 (continued)

| miRNA family | L. campestre mature miRNA | L. campestre ShortStack genome | L. campestre ShortStack transcriptome | L. appelianum mature miRNA | L. appelianum ShortStack transcriptome |
|--------------|----------------------------|--------------------------------|---------------------------------------|---------------------------|----------------------------------------|
| miR172a/miR172b-3p/miR172c/miR172d-3p/miR172e-3p | x | miR172b (partial) | miR172e | x | miR172b (partial) |
| miR172b-5p/miR172e-5p | x | miR172b (partial) | miR172e | x | miR172b (partial) |
| miR172d-5p | x | miR172b | x | x | miR172b (partial) |
| miR2111b-3p | x | x | x | x | x |
| miR319a/miR319b | x | x | x | x | x |
| miR319c | x | x | x | x | x |
| miR390a-3p | x | x | x | x | x |
| miR390a-5p/miR390b-5p | x | x | x | x | x |
| miR390b-3p | x | x | x | x | x |
| miR393a-3p/miR393b-3p | x | miR393b | x | miR393b | x | miR394b |
| miR393a-5p/miR393b-5p | x | miR393b | x | miR393b | x | miR394b |
| miR394a/miR394b-5p | x | x | x | x | x |
| miR395a/miR395b/miR395c/miR395d/miR395e/miR395f | x | x | x | x | miR395d, miR395f |
| miR396a-3p | x | miR396a | x | x | x |
| miR396a-5p/miR396b-5p | x | miR396a | x | x | x |
| miR396b-3p | x | x | x | x | x |
| miR398a-3p/miR398b-3p/miR398c-3p | x | miR398a | x | miR398b | x | miR398b |
| miR399a/miR399b/miR399c-3p/miR399f | x | miR399a | x | x | x |
| miR399f | x | x | x | x | x |
| miR403-3p | x | x | x | x | x |
| miR403-5p | x | x | x | x | x |
| miR408-3p | x | x | x | x | x |
| miR408-5p | x | x | x | x | x |
| miR472/miR472 | x | miR472 | x | x | x |
| miR8174 | x | x | x | x | x |
| miR8175 | x | x | x | x | x |
| miR824-3p | x | x | x | x | x |
| miR824-5p | x | x | x | x | x |
| miR827 | x | x | x | x | x |
| miR845a | x | x | x | x | x |
| miR845b | x | x | x | x | x |
| miR858a/miR858b | x | x | x | x | x |
| miR863-5p | x | x | x | x | x |
We also analyzed DDEGs in our dataset, i.e. genes which had a significantly different change in expression in flowers versus buds and in fruits versus flowers, respectively, in *L. appelianum* as compared to *L. campestre*. These genes may have a significantly stronger up- or downregulation in *L. appelianum* as compared to *L. campestre* or these genes may be downregulated in one species and upregulated in the other species. We found 70 DDEGs in flowers versus buds and 158 DDEGs in fruits versus flowers when comparing the two species (Fig. 4).

We applied the same methods for the identification of DEGs and DDEGs encoding miRNAs. First, we determined orthologs between the miRNAs based on the *A. thaliana* miRNAs they mapped to. For 56 mature miRNAs belonging to 28 miRNA families reads were found in the small RNA data for both species and these mature miRNAs could thus be used for differential expression analyses (Table 2). We will refer to this dataset as our ortholog-miRNAs. All 16 miRNA families that are conserved across virtually all species according to [37] and 6 out of 21 miRNA families which were classified as moderately conserved belong to our ortholog-miRNAs dataset. Mapping of small RNA reads, counting and differential expression analyses were done as described for the differential expression analysis of the ortholog-transcriptomes.

Only one miRNA was found to be encoded by a DEG or DDEG by both programs DESeq2 and edgeR. The miRNA homologous to miR165a-3p, miR165b, miR166a-3p, miR166b-3p, miR166c, miR166d, miR166e-3p, miR166f and miR166g of *Arabidopsis thaliana* [45] (they all only differ by one nucleotide), which we will refer to as miR165a-3p, was found to be encoded by a DDEG when comparing fruits and flowers. Targets of miR165a-3p are HD-Zip transcription factors like PHABULOSA, REVOLUTA and PHAVOLUTA [46]. However, the expression of these target genes does not change much in our transcriptome analyses (Supplemental Fig. 2). Hence, the role of miR165a-3p for fruit dehiscence remains to be clarified.

**Gene Ontology and transcription factor analyses**

A number of gene ontology (GO) terms [47, 48] of the category molecular function are significantly over- or underrepresented in the DEGs and DDEGs (Table 3). Among them, the terms protein binding (GO:0005515) and RNA binding (GO:0003723) were underrepresented in two datasets of DEGs. Interestingly, several GO terms related to cell wall synthesis and degradation, i.e. pectinesterase activity (GO:0030599), cellulose synthase (UDP-forming) activity (GO:0016760), polygalacturonase activity (GO:0004650) and hydrolase activity, hydrolyzing O-glycosyl compounds (GO:0004553) were overrepresented in different sets of DEGs.
As we were interested in differences in the gene regulatory network involved in fruit dehiscence in the two species, known to be largely composed of transcription factors in Arabidopsis thaliana (reviewed in Ballester and Ferrándiz [12]), we analyzed genes annotated to have “DNA-binding transcription factor activity” (GO:0003700) in more detail. This set includes transcription factors and transcriptional regulators. For simplicity, we will refer to this dataset as genes encoding transcription factors (TFs).

When comparing flowers and buds, 21 and 28 TFs were DEGs in L. campestre and in L. appelianum, respectively. Among them, there are 13 TFs that were DEGs comparing these structures in both species, including four genes with known functions in flower development, AGAMOUS-LIKE 104 (AGL104) [49], SPOROCYTELESS (SPL, also termed NOZZLE) [50], OREARAI (ORE1, also termed ANAC092, ATNAC2, ATNAC6) [51] and ZINC FINGER PROTEIN 2 (ZFP2) [52] (Table 4). Between fruits and flowers, there are 12 TFs in L. campestre and 23 TFs in L. appelianum that
are DEGs. Five of these genes are DEGs in fruits versus flowers in both species (Table 4). TFs with differential expression between structures in both species are probably those TFs with common functions for flower and fruit development.

When comparing the two species, 43 TFs were DEGs in buds, 68 in flowers and 49 in fruits. Among these TFs, 19 were DEGs in all structures (Table 5). Interestingly, four genes involved in flowering time determination, *SQUAMOSA PROMOTER BINDING PROTEIN-LIKE 4* (SPL4) [53], *NUCLEAR FACTOR Y-B2* (NF-YB2) [54], *NUCLEAR FACTOR Y-B10* (NF-YB10) [55] and *FLOWERING LOCUS C* (FLC) [56], as well as the fruit development genes *FRUITFULL* (FUL) [17] and *YABBY3* (YAB3) [24, 30] (Fig. 5) were on this list.

Two TFs were found to be DDEGs when comparing flowers and buds in the two species (Table 6), among them *MASSUGU1 2* (MSG2, also known as *INDOLE-3-ACETIC ACID INDUCIBLE 19*) [58] which has been shown to be involved in stamen filaments development [59]. Comparing fruits and flowers, seven TFs, *PHY-INTERACTING FACTOR 1* (PIF1, also known as *PHYTOCHROME INTERACTING FACTOR 3-LIKE 5*) [60], *MYB DOMAIN PROTEIN 57* (MYB57) [61], *TCP FAMILY TRANSCRIPTION FACTOR 4* (TCP4, also known as *MATERNAL EFFECT EMBRYO ARREST 35*) [62], *BRANCHED 1* (BRC1, also known as *TCP FAMILY TRANSCRIPTION FACTOR 18*) [38], *REVEILLE 6* (RVE6) [63], *TRIPTYCHON* (TRY) [64] and *OBF BINDING PROTEIN 4* (OBP4, also termed *DOF5.4*) [65] are DDEGs in *L. appelianum* as compared to *L. campestre* (Supplemental Fig. 4).

| Dataset           | GO term                                              | Fold enrichment | FDR        |
|-------------------|------------------------------------------------------|-----------------|------------|
| La vs. Lc bud     | protein binding (GO:0005515)                         | 0.74            | 3.43E-02   |
|                   | transferase activity, transferring phosphorus-containing groups (GO:0016772) | 0.44            | 2.84E-02   |
| La vs. Lc flower  | none                                                 |                 |            |
| La vs. Lc fruit   | none                                                 |                 |            |
| Lc flower vs. bud | pectinesterase activity (GO:0030599)                 | 10.69           | 1.48E-02   |
|                   | RNA binding (GO:0003723)                             | 0.15            | 8.99E-03   |
| La flower vs. bud | sodium:proton antiporter activity (GO:0015385)      | 14.17           | 2.35E-04   |
|                   | cellulose synthase (UDP-forming) activity (GO:0016760) | 10.52           | 3.18E-02   |
|                   | polygalacturonase activity (GO:0004650)              | 8.13            | 8.50E-03   |
|                   | iron ion binding (GO:0005506)                        | 3.12            | 3.48E-02   |
|                   | oxidoreductase activity acting on paired donors with incorporation or reduction of molecular oxygen (GO:0,016,705) | 2.89            | 4.07E-02   |
|                   | protein binding (GO:0005515)                         | 0.71            | 3.99E-02   |
|                   | RNA binding (GO:0003723)                             | 0.17            | 1.30E-03   |
| Lc fruit vs. flower| heme binding (GO:0020037)                            | 5.11            | 9.54E-04   |
|                   | hydrolyase activity, hydrolyzing O-glycosyl compounds (GO:0004553) | 4.28            | 1.15E-03   |
| La fruit vs. flower| none                                                 |                 |            |
| flower vs. bud    | acid-amino acid ligase activity (GO:0016881)         | 40.35           | 1.53E-02   |
| fruit vs. flower  | none                                                 |                 |            |

Extension of the gene regulatory network for fruit development

We next investigated how the TFs shown to be DDEGs between fruits and flowers may be involved in the gene regulatory network controlling fruit development (Fig. 5). Therefore, we searched for binding sites of the seven TFs identified by chromatin immunoprecipitation followed by sequencing (ChIP-seq) experiments in the promoters of the genes known to be involved in fruit development. On ChIP-Hub [66], no ChIP-seq data is available for BRC1 and for TRY.

Binding of OBP4 was found in the promoter of all but one of the 18 fruit development genes (Table 7). Binding of RVE6, MYB57, PIF1 and TCP4 was detected in the promoters of 11, 7, 5 and 2 fruit development genes, respectively. PIF1 predominantly binds to the promoters of valve identity genes, with binding to four out of eight valve identity gene promoters and apart from that only binding to one of five valve margin genes. ARF8 is the only fruit development gene for which none of the DDEGs was found to bind to its promoter. To the promoters of YAB3 and FUL, which were found to be
differentially expressed in all structures between L. campestre and L. appelianum, binding of TCP4, RVE6 and OBP4 and of MYB57, RVE6 and OBP4, respectively, was found. It has to be noted, however, that for the ChIP-seq experiments analyzed here, material from young leaves (MYB57, RVE6 and OBP4) or seedlings (PIF1 and TCP4) was used. Hence, as to whether these factors bind to the promoters of fruit development genes in reproductive tissues has still to be determined.

**Discussion**

**Transcriptomes and small RNA datasets of L. campestre and L. appelianum are nearly complete**

We have sequenced the transcriptomes of floral buds, flowers and fruits of L. campestre and L. appelianum. Benchmarking of Universal Single-Copy Orthologs (BUSCO) analysis revealed that the transcriptome assemblies of the two species contain more than 94% of the eudicotyledous "near-universal single-copy orthologs". This number is similar to or more than that for transcriptome assemblies of other Brassicaceae [67–69]. Furthermore, we found members of all 16 miRNA families that were found to have originated before the emergence of eudicots and conserved in eudicots [37]. These findings reveal that our transcriptome and small RNA data includes most of the expected transcripts and miRNAs.

**Differences in gene expression mainly between floral structures**

We identified more DEGs when comparing the same structure between the two species than comparing different structures in the same species (Fig. 4). This indicates that gene regulation has diverged between the two species. This is different to what has been observed other flowering plant species, where the correlation of gene expression is higher in the same structure of different species than in different structures of the same species [70]. However, in this case microarray expression data was analyzed which may select for conserved genes.

The highest number of DEGs was observed in flowers, followed by fruits and buds, and the highest

| Ortholog ID | Ortholog name | Ortholog description (based on TAIR) | Reg. L.c | Reg. L.a |
|-------------|---------------|-------------------------------------|----------|----------|
| AT1G22130.1 | AGL104        | Pollen development and pollen tube growth | -3.0     | -3.6     |
| AT1G61110.1 | anac025, NAC025 | Endosperm cell expansion during germination | -3.6     | -3.2     |
| AT1G69490.1 | ANAC029, ATNAP, NAP | Leaf senescence, drought stress response | 3.4      | 2.8      |
| AT2G47190.1 | ATMYB2, MYB2 | Salt tolerance, Phosphate Starvation Response, Abscisic Acid Signaling, Plant Senescence | 3.7      | 2.6      |
| AT3G04070.1 | anac047, NAC047 | Flood induced leaf movement | 3.3      | 3.8      |
| AT3G23050.1 | ARX2, IAA7 | Auxin response, shoot and root gravitopism | 2.0      | 3.1      |
| AT3G58120.1 | ATBZIP61, BZIP61 | n.a | -3.7 | -2.5 |
| AT4G10240.1 | bbx23 | Temperature-induced hypocotyl elongation together with BBX18, photomorphogenesis activated by PIF1 and PIF3 | -4.8 | -6.3 |
| AT4G27330.1 | NZZ, SPL | Initiation of micro- and megagametogenesis, patterning of the ovule, differentiation of primary sporogenous cells into microsporocytes, regulation of another cell differentiation | -7.9 | -8.7 |
| AT4G28500.1 | ANAC073, NAC073, SND2 | Secondary cell wall development, phloem development | -2.8 | -2.9 |
| AT5G13180.1 | ANAC083, NAC083, VNI2 | Xylem vessel formation, leaf senescence | 2.7 | 3.5 |
| AT5G39610.1 | ANAC092, ATNAC2, ATNAC6, NAC2, NAC6, ORE1 | Leaf senescence, Termination of flower receptivity | 4.0 | 3.5 |
| AT5G7520.1 | ATZFP2, ZFP2 | Abscission of floral organs | 2.1 | 3.4 |
| AT2G01940.3 | ATIDD15, SGR5 | Auxin biosynthesis and transport, aerial organ morphogenesis and gravitropic responses | -3.3 | -4.2 |
| AT2G20180.2 | PIF1, PIL5 | Negative regulation of phytochrome-mediated seed germination | -2.5 | -5.2 |
| AT3G23050.1 | ARX2, IAA7 | Auxin response, shoot and root gravitopism | -2.3 | -3.2 |
| AT5G64530.1 | ANAC104, XND1 | Xylem formation, Regulation of secondary wall synthesis | -3.6 | -5.1 |
| AT5G67300.1 | ATMYB44, ATMYBR1, MYB44, MYBR1 | Abscisic acid signaling, abiotic stress tolerance | -2.3 | -3.9 |

**Table 4** Genes that are differentially expressed in different structures in both species and that are annotated as “DNA-binding transcription factor activity” (GO:0003700). Reg., regulation; L.c., L. campestre; L.a., L. appelianum
number of DDEGs was found in fruits versus flowers as compared to flowers versus floral buds. This indicates that the differences in gene expression between \textit{L. campestre} and \textit{L. appelianum} are most pronounced between flowers and in the transition from flowers to fruits. This is expected as the developmental program leading to fruit dehiscence or indehiscence needs to be initiated before the fruits are formed. Supportingly, in \textit{Aethionema arabicum}, a plant that develops dehiscent and indehiscent fruits on the very same individual, differences between the fruit types start to occur early, two days after anthesis [71].

A number of GO terms related to cell wall synthesis and degradation, e.g. pectinesterase activity, cellulose synthase activity and polygalacturonase activity were overrepresented in different sets of DEGs. It has been recognized that secondary cell wall formation at the valve margins [72] and degeneration of cell walls in the separation layer are essential processes for fruit dehiscence after the DZ is correctly specified [73]. Hence, the overrepresentation of GO terms related to cell wall synthesis and degradation is not surprising.

\textbf{Confirmation of previous expression study}

In a previous study, we have compared expression of the valve margin genes as well as the valve gene \textit{FUL} and the replum gene \textit{RPL} between \textit{L. campestre} and \textit{L. appelianum} by in situ hybridization [7]. We showed that their orthologues from \textit{L. campestre} (dehiscent fruits) are similarly expressed as in \textit{A. thaliana} while expression of the respective orthologues is abolished in valve margins of indehiscent \textit{L. appelianum} fruits. Analysis using qRT-PCR revealed that the valve margin genes \textit{IND} and \textit{SHP1} are expressed at a significantly higher level in flowers and early fruits (the fruit stage for which the transcriptome was sequenced here) of \textit{L. campestre} than in \textit{L. appelianum}. Significantly higher expression was confirmed in the present study for \textit{SHP1} in flowers (Fig. 5). qRT-PCR analysis revealed significantly higher expression of \textit{SHP2} in flowers and of \textit{ALC} in early fruits [7]. Expression was not significantly different in the present transcriptome analysis, but expression was also found to be higher for \textit{SHP2} in flowers and for \textit{ALC} in early fruits. Like qRT-PCR analysis, our transcriptome analysis also found significantly higher expression of \textit{FUL} in flowers of \textit{L. campestre} [7]. Expression was not significantly different in the present transcriptome analysis, but expression was also found to be higher for \textit{FUL} in flowers and for \textit{ALC} in fruits. Like qRT-PCR analysis, our transcriptome analysis also found significantly higher expression of \textit{AP2} in flowers and of \textit{ALC} in early fruits of \textit{L. appelianum} [7]. Expression was not significantly different in the present transcriptome analysis, but expression was also found to be higher for \textit{AP2} in flowers and for \textit{ALC} in fruits. Like qRT-PCR analysis, our transcriptome analysis also found significantly higher expression of \textit{AP2} in flowers and of \textit{ALC} in early fruits of \textit{L. campestre} by both analyses. Again, the difference was significant in qRT-PCR analyses but not in transcriptome analysis. Hence, our transcriptome analysis is in good agreement with the previous qRT-PCR analyses, but in our transcriptome analysis...
Known flower and fruit development genes are differentially expressed
To identify differences in the regulation of flower and fruit development between *L. campestre* and *L. appelianum*, we focused on differentially expressed or differently differentially expressed genes encoding transcription factors (TFs). Among 19 genes encoding TFs which were found to be differentially expressed in all three examined structures, four TFs are involved in flowering time determination. *L. appelianum* and *L. campestre* have different flowering periods according to the Jepson Herbarium (Jepson Flora Project (eds.) 2021, Jepson eFlora, https://ucjeps.berkeley.edu/eflora/, accessed on May 25, 2021), which may be caused by
differences in the expression of the identified flowering time genes.

As mentioned above, the fruit development genes \( SHP1 \) and \( FUL \) were found to be differentially expressed. \( FUL \) is expressed at a significantly lower level in \( L. appelianum \) in all three structures. In \( A. thaliana \), the \( ful \) knockout mutation causes indehiscence \([16, 17]\). \( FUL \) represses the expression of \( SHP1 \) and \( SHP2 \). Hence, lower expression of \( FUL \) in \( L. appelianum \) should lead to higher expression of \( SHP1 \) and \( SHP2 \) in this species. However, only for \( SHP2 \) a non-significantly higher expression has been found in flowers of \( L. appelianum \) (Fig. 5). \( SHP1 \) is unexpectedly expressed at a significantly lower level in \( L. appelianum \) flowers. This may be due to the fact that \( YAB3 \) \([24, 30]\) (Fig. 5) was expressed at a significantly lower level in \( L. appelianum \). \( YAB3 \) does not only activate expression of \( FUL \) but also that of \( SHP1 \) and \( SHP2 \) together with \( JAG \) and \( FIL \) \([24]\). Hence, a lower level of \( YAB3 \) does not only lead to a lower expression of \( FUL \) and to less repression of \( SHP1 \) and \( SHP2 \) by \( FUL \) but also to less activation of \( SHP1 \) and \( SHP2 \) by \( YAB3 \). The contrasting effects of \( YAB3 \) and \( FUL \) on the expression of \( SHP1 \) and \( SHP2 \) may lead to the observed overall similar expression of the \( SHP \) genes in \( L. campestre \) and \( L. appelianum \) but may still lead to differences in dehiscence due to different expression domains as

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**Table 6** DDEGs in different structures annotated as “DNA-binding transcription factor activity” (GO:0003700)

| Ortholog ID | Ortholog name | Ortholog description (based on TAIR) | Reg |
|-------------|---------------|--------------------------------------|-----|
| flower vs. bud | AT3G15540.1 | IAA19, MSG2 | Stamen filaments development | 4.8 |
| fruit vs. flower | AT5G47230.1 | AtMACD1, ERF102, ERF5 | Stress response, leaf growth | 5.3 |
| AT2G20180.2 | FIF1, PILS | Phytochrome-mediated seed germination | 2.7 |
| AT3G01530.1 | ATMYB57, MYB57 | Stamen and nectary development | -4.9 |
| AT3G18550.1 | MEE35, TCP4 | Cotyledon, leaf and petal development, seed oil accumulation | -4.3 |
| AT3G18550.1 | BRC1, TCP18 | Arrests axillary bud development and prevents axillary bud outgrowth. Role in flowering control | -4.3 |
| AT5G52660.2 | RVE6 | Involved in circadian rhythm | -2.6 |
| AT5G32000.1 | TRY | Trichome and root hair patterning, phosphate starvation response | -6.0 |
| AT5G60850.1 | DOF5.4, OBP4 | Cell Cycle Progression and Cell Expansion | -2.3 |

**Table 7** Number of binding sites of TFs found to be DDEGs to the promoters of known fruit development genes

| | PIF1 | MYB57 | TCP4 | RVE6 | OBP4 |
|---------------|------|------|------|------|------|
| Valve | AS1 | 1 | 1 | - | - | 3 |
| | AS2 | - | 1 | - | - | 1 |
| | JAG | 2 | - | - | - | 1 |
| | FIL | 1 | - | - | 3 | 2 |
| | YAB3 | - | - | 1 | 2 | 1 |
| | ARF6 | 1 | - | - | 2 | 1 |
| | ARF8 | - | - | - | - | - |
| | FUL | - | 1 | - | 1 | 1 |
| | AP2 | - | 1 | - | - | 2 |
| Replum | NTT | - | 1 | - | 1 | 3 |
| | BP | - | - | - | 1 | 2 |
| | WOX13 | - | - | - | - | 4 |
| | RPL | - | - | - | 3 | 1 |
| Valve margin | SHP1 | - | - | - | 2 | 5 |
| | SHP2 | - | 1 | - | 1 | 5 |
| | IND | - | - | - | 2 | 3 |
| | ALC | - | 1 | 1 | 2 | 2 |
| | SPT | 1 | - | - | 1 | 2 |
described in [24]. *yab3* single mutants do not have any major defects in dehiscence but *fil yab3* double mutants are largely indehiscent [24]. Hence, decreased expression of *YAB3* in *L. appelianum* as compared to *L. campestre* may have been an important factor for the evolutionary shift from dehiscent to indehiscent fruits in *L. appelianum*. This also shows that there was not only a change in the control of valve margin identity genes but also of the valve identity genes and shifts the causative mutation further upstream in the gene regulatory network of fruit development.

**MiR165 is differently expressed in fruits versus flowers**

Our smallRNA sequencing revealed that the miRNA homologous to miR165a-3p, miR165b, miR166a-3p, miR166b-3p, miR166c, miR166d, miR166e-3p, miR166f and miR166g [45] is encoded by a DDEG when comparing fruits and flowers. Targets of miR165 and miR166 are the mRNAs of HD-Zip transcription factors like PHABULOSA (PHB), REVOLUTA and PHAVOLUTA [46]. Recently, a function of the miR166-PHB module in anther dehiscence has been elucidated [74]. Upregulation of miR166 in the *jba-1D* mutant leads to downregulation of its target gene PHB which results in increased expression of SPOROCYTELESS/NOZZLE (*SPL/NZZ*). *jba-1D* mutants do not develop a dehiscence zone in anthers, i.e. overexpression of miR166 leads to indehiscence of anthers. Expression of miR166 in fruits is much higher in *L. appelianum* (indehiscent fruits) than in *L. campestre* (dehiscent fruits), while the opposite is the case in flowers (Fig. 6). Hence, miR166 may have a role in the development of indehiscent fruits in *L. appelianum* though the details of the regulation remain to be elucidated.

**BRC1 and TCP4 as candidate genes for the evolutionary shift from dehiscent to indehiscent fruits**

Our transcriptome analysis also identified seven genes encoding TFs belonging to DDEGs when comparing flowers and fruits (Table 6). PIF1 is a basic helix-loop-helix (bHLH) transcription factor that negatively regulates chlorophyll biosynthesis [60]; it is involved in a variety of biological processes such as the repression of light-induced seed germination and chlorophyll accumulation in light [75]. RVE6 is a MYB protein that controls the pace of the circadian clock together with its close homologs RVE4 and RVE8 [63]. The zinc finger protein OBP4 functions in cell cycle progression and cell expansion [65] and is involved in root development [76, 77]. So far, involvement of these three factors in flower and fruit development has, to the best of our knowledge, not been reported.

Two other MYB genes, *MYB57* and *TRY* have also been found to be DDEGs (Table 6). *MYB57* functions redundantly with MYB21 and MYB24 to regulate stamen development [78]. TRY controls the spacing pattern of trichomes, which are single-celled hairs [64]. Recently it has been found that *TRY* and other MYB genes of the regulatory network for trichome patterning have been modulated to trigger trichome development in fruits [79]. Hence, these two genes are known to function during flower and fruit development but association with fruit dehiscence is not known so far.

More interestingly, the genes encoding for the two TCP transcription factors BRC1 and TCP4 are DDEGs between fruits and flowers when comparing *L. campestre* and *L. appelianum*. Expression of BRC1 correlates with bud inhibition [38, 80] but recently, it has been shown that *BRC1* is neither necessary nor sufficient for bud inhibition [81]. Noticeably, it has been hypothesized that *BRC1* may guide fruit morph determination in the dimorphic Brassicaceae plant *Aethionema arabicum* [71]. *Ae. arabicum* produces two fruit morphs on the same plant, one of which is dehiscent and the other one is indehiscent. qRT-PCR analyses showed that the expression of *BRC1* in *Ae. arabicum* is high in flowers and decreases strongly
in fruits of the indehiscent morph but remains at a low level in flowers and fruits of the dehiscent morph. We observe a very similar pattern in our transcriptome analysis for the indehiscent morph in *L. appelianum* and the dehiscent morph in *L. campestre* (Fig. 7). In Arabidopsis thaliana buds, BRC1 controls a transcription factor cascade that results in abscisic acid (ABA) accumulation [82]. It has been proposed that this cascade also plays a role in the development of indehiscent fruits in *Ae. arabicum* [83]. Thus the effect of BRC1 on fruit indehiscence in *L. appelianum* may be indirect via ABA.

TCP4 has been found to be involved in leaf and flower development as well as in seed oil biosynthesis in *A. thaliana* [62, 84, 85]. Furthermore, TCP4 directly activates the expression of miR167 which targets the TFs ARF6 and ARF8 [86]. This regulation has been hypothesized to be important for flower maturation, but may also be involved in fruit dehiscence as ARF6 and ARF8 are part of the gene regulatory network of fruit development [29] (Fig. 5). Another study found physical interaction of TCP4 and AS2 in yeast-two-hybrid experiments [87]. AS2 has also previously been found to be involved in fruit patterning [25] (Fig. 5). Our analysis of ChiP-seq data on ChiP-Hub [66] additionally revealed that TCP4 binds to the promoter of YAB3 (Table 7), which has been found to be differentially expressed between *L. campestre* and *L. appelianum* in all structures examined. In flowers, TCP4 is expressed at a higher level in *L. appelianum* than in *L. campestre* while the expression pattern is the other way round for YAB3. Hence, it is conceivable that TCP4 represses YAB3 in flowers.

**Conclusions**

Taken together, our study provides insights into the gene regulatory differences in fruit development between *L. campestre* producing dehiscent fruits and *L. appelianum* forming indehiscent fruits. We confirm differences in the expression of the fruit development genes SHP2 and FULL between the two species and reveal the importance of the valve identity gene YAB3 for fruit indehiscence in *L. appelianum*. We uncover the microRNA miR166 and the TCP transcription factors BRC1 and TCP4 as new candidates for causing the evolutionary transition from dehiscent to indehiscent fruits in *L. appelianum*.

**Methods**

**Plant material, RNA extraction and sequencing**

Seeds of *Lepidium appelianum* (KM 1754) were obtained from J Gaskin, USDA, Fremont County, Wyoming, USA and seeds of *L. campestre* (KM 96) were acquired from the Botanical Garden of the University of Zürich. All seeds were subsequently mass propagated in the Botanical Garden of Osnabrück University, Germany. Seeds from these mass propagations were sown on a mixture of seedling substrate (Kammlott, Kammlott GmbH, Erfurt, Germany)/sand/vermiculite (1–3 mm) (8:1:1) which was supplemented with 1 g L⁻¹ each of Osmocote mini (Scotts Miracle-Gro Company, Marysville, OH, USA) and Triabon (http://www.compo-expert.com, COMPO Expert GmbH, Münster, Germany). The seeds were placed for 4 days at 4 °C for stratification and then put in the greenhouse under a light–dark cycle of 16 h light, 8 h dark of artificial light, plus daylight. After 5 weeks in the greenhouse, the plants were vernalized for at least 13 weeks at 4 °C with a light–dark cycle of 8 h light, 16 h dark. After vernalization, the plants were put back in the greenhouse. Plant material was harvested from two batches of independently grown plants 3 to 5 weeks after the end of vernalization. Plant material was harvested between 12 and 4 pm to minimize the effect of circadian rhythm. Late flower buds, flowers and early fruits were harvested and immediately frozen in liquid nitrogen. Three samples were taken for each species and each structure resulting in 18 samples in total. For each sample, about 100 mg plant material was pooled from four individual plants. The material was pulverized in liquid nitrogen using a mortar and pestle. RNA was extracted using Qiazol (Qiagen) according to the manufacturer’s instructions.

![Fig. 7](image-url) Expression data plot of BRC1 in *L. campestre* (Lc) and *L. appelianum* (La). Bars indicate mean normalized count values of reads mapping to BRC1 in the corresponding structure and species. Dark and light grey bars represent the mean values for *L. campestre* (Lc) and for *L. appelianum* (La), respectively. The error bars indicate the standard deviation.
RNA quantity and quality was checked by gel electrophoresis. The samples were sent to the Vienna BioCenter Facility for Next Generation Sequencing where they were quality checked and sequenced on a HiSeqV4. For mRNA sequencing, 125 bp paired-end reads were produced and 50 bp single-end reads were generated for small RNA sequencing.

Preprocessing of RNA-seq data

Raw reads were corrected using Rcorrector [88] with default settings. Uncorrectable reads were excluded using a python script obtained from https://informatics.fas.harvard.edu/best-practices-for-de-novo-transcriptome-assembly-with-trinity.html which was slightly modified for excluding uncorrectable reads from smallRNA libraries. The remaining reads were trimmed with Trim Galore! (https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/) using the following settings: --clip_R1 12, --clip R2 12, --paired, --retain_unpaired, --phred33, --length 36, -q 5, --stringency 5, -e 0.1 for transcriptome reads and the following settings --phred33, --length 18, --max_length 26, -q 5, --stringency 5, -e 0.1, -a adapter for small RNA reads where adapter was replaced by the corresponding adapter sequence identified using FastQC (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/). Thereafter, Poly-A and Poly-T tails were removed from transcriptome reads with PrinSeq [89] using the settings --trim_tail_left 5 and --trim_tail_right 5. Reads that mapped to the genome of *Frankliniella occidentalis* (GenBank: GCF_000697945) or to rRNAs (GenBank: X52320.1), mitochondrial (GenBank: Y08501.2) or chloroplast (GenBank: AP000423.1) sequences from *A. thaliana* as determined using bowtie2 (settings: --very-sensitive-local, --phred33) [90] were excluded from further analyses from both, the transcriptome and the smallRNA libraries.

De novo assembly

To simplify de novo assembly, also duplicate reads, i.e. reads with the exact same length and sequence, were removed. The remaining reads were assembled using Trinity [32] with default settings for the two species *L. campestre* and *L. appelianum* separately. To identify remaining contamination in the transcriptome, a BLASTn search was conducted against the nucleotide database of NCBI (nt) using the transcripts in the assembly as query with the option -max_target_seqs 5. Transcripts for which the best BLASTn result came from a non-plant species and had an eValue of $E < 10^{-10}$ were removed from the transcriptomes. The completeness of the assembled transcriptomes was evaluated using the Benchmarking Universal Single-Copy Orthologs tool BUSCO [34] and their accompanying dataset for eudicotyledons with 2121 groups (odb10).

Separation of chimeras in the assemblies

The initial assemblies contained chimeras composed of two different transcripts. As these chimeras often result from misassembly [33], we sought to separate chimeras into their separate transcripts. To recognize chimeras, we first conducted a BLASTn search [91] using the transcripts from the *Lepidium* transcriptomes as query and the cDNA sequences of the representative *A. thaliana* gene model as provided by TAIR10 (TAIR10_cdna_20110103_representative_gene_model_updated.fasta) as database saving the best two subjects (i.e. *A. thaliana* cDNAs) for each query (i.e. for each transcript from the *Lepidium* transcriptomes) (Fig. 8). Using a customized perl-script (Supplementary Data S1) we searched for instances where the two subjects fitted to different regions of the query (i.e. one part of the *Lepidium* transcript has a BLAST hit corresponding to one *A. thaliana* cDNA while another part of the same transcript has a BLAST hit corresponding to another *A. thaliana* cDNA). These instances likely indicate chimeric *Lepidium* transcripts. To identify the best position to split the chimeras, we considered at which position and to what extend the *A. thaliana* cDNAs matched to the *Lepidium* transcripts as shown in Fig. 8. Chimeras were split if the overlap was less than 150 nucleotides either in the middle of the overlap or at the positions corresponding to the corrected end and beginning of the involved transcripts (Fig. 8).

Identification of miRNAs in smallRNA libraries

SmallRNA reads were mapped to mature miRNAs from *Arabidopsis thaliana* as downloaded from miRBase [35] employing bowtie2 with the settings -N 1, -L 18. If a read mapped to a specific miRNA from *A. thaliana* this miRNA was considered to be present in the corresponding *Lepidium* species. Mature miRNAs only differing by one nucleotide were combined to avoid multiple mapping during read-counting. To identify novel miRNAs in *Lepidium*, we used ShortStack [92] with the parameters --foldsize 500, --dicermin 18 and the trinity transcriptome assembly of the corresponding *Lepidium* species as reference “genome”. For *L. campestre*, ShortStack was run a second time, this time using the genome sequence of *L. campestre* as available at the National Centre for Biotechnology Information [93] as reference. The stem-loop sequences classified as “N15” or “Y” by ShortStack were used as query sequences for BLAST searches of pre-miRNAs of *A. thaliana* as
downloaded from miRBase [35] to distinguish known from novel miRNAs. The stem-loop sequences classified as “Y” by ShortStack without similarity to pre-miRNAs of A. thaliana were classified as novel Lepidium miRNAs.

**Determination of orthologs**

For transcriptome data, putative ortholog pairs were determined using a reciprocal best hit approach as follows. BLASTn searches were conducted using the transcriptome assembly with chimeras separated of L. campestre as query and the transcriptome assembly with chimeras separated of L. appelianum as subject and vice versa. For each transcript of L. campestre the best BLAST hits (i.e. all hits having the same eValue, score and alignment length) in L. appelianum were recorded and vice versa. If a transcript Tc from L. campestre had the transcript Ta from L. appelianum among its best BLAST hits and transcript Ta from L. appelianum had transcript Tc from L. campestre in its list of best BLAST hits, these were considered as best reciprocal BLAST hit. Best reciprocal BLAST hits with an alignment length of more than 250 nucleotides and where the length of the shorter sequence was at least 50% of that of the longer sequence were considered as putative ortholog pairs. Additionally, another BLASTn search was conducted using the transcripts in the transcriptomes as query and the Arabidopsis thaliana TAIR10 cDNA dataset as database. The set of putative orthologous transcript pairs was pruned such that only one transcript isoform was kept for each species unless different isoforms fitted to different A. thaliana genes. The isoform with the longest alignment length between the two species was chosen to be kept. This way, for each transcript in the one transcriptome exactly one transcript in the other transcriptome was kept. We refer to this dataset as the ortholog-transcriptome. The transcripts in the ortholog-transcriptome dataset were named using the TAIR10 identifier of the best BLAST result or numbered if no BLAST result was obtained this way.

For the miRNA data, orthologous miRNAs of the two Lepidium species were defined as those miRNAs fitting to the same miRNA from A. thaliana. Comparison of the novel Lepidium miRNAs revealed that none of these was found in both species.

**Read mapping and feature counting**

Preprocessed transcriptome and small RNA reads were mapped against ortholog-transcriptome and mature miRNAs, respectively, using bowtie2 (settings:
Differential gene expression analysis pipeline

Differentially expressed genes were identified using R (https://www.r-project.org/) and the Bioconductor packages edgeR [44] and DESeq2 [43]. Transcript counts were normalized with respect to transcript length. Lowly expressed transcripts with normalized counts and lowly expressed miRNAs with raw counts of less than 19 were discarded. Considering the two species L. campestre and L. appelianum and the structures bud, flower and fruit, the following multi-factor design was used: species + structure + species:structure. A Likelihood Ratio Test (LRT) and a quasi-likelihood F-test were conducted in DESeq2 (command: DESeq(object, test="LRT", reduced=~species + structure)) and EdgeR (command: glmQLFit(object, design)), respectively to identify differentially expressed and differentially expressed genes. Only transcripts and miRNAs having a log-fold change to the base of 2 of more than 2 were considered. For DESeq2 the false discovery rate threshold $\alpha$ was set to 0.001.

For principal component analysis, count data was normalized using regularized logarithm with the option blind=FALSE in DESeq2 and the principal components were plotted using the plotPCA function in R.

GO enrichment analysis

Gene Ontology (GO) enrichment analysis was conducted on the GO website (http://geneontology.org/) using the PANTHER Overrepresentation Test [94]. The TAIR10 identifiers of the transcripts in the ortholog-transcriptome were provided as reference list. The TAIR10 identifiers of the transcripts which were identified as significantly differentially expressed genes by both programs, DESeq2 and EdgeR, were provided as analyzed list. Arabidopsis thaliana was chosen as organism and “GO Molecular function complete” was selected as annotation data set. Enriched GO categories were determined using the Fisher’s Exact Test with False Discovery Rate correction.

GO categories and terms were also determined using the AnnotationDbi in R. Transcripts associated with the term “DNA-binding transcription factor activity” were analyzed further.

Promoter analyses

Binding of transcription factors to the promoters of genes involved in fruit opening was analysed using ChIP-Hub (http://www.chip-hub.org/). ChIP-Hub provides access to data on binding sites determined using chromatin immunoprecipitation followed by sequencing (ChIP-seq). On the ChIP-Hub website, A. thaliana was chosen as species and binding data was visualized on the WashU Epigenome Browser. For each fruit development gene, 1,500 nucleotides upstream of the translation start codon were investigated and each occurrence of binding of one of the transcription factors found to be DDEGs was noted.

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Authors’ contributions
GT conceived the project. LG and GT designed the experiments. KK performed the experiments. LG, NFP and MH analyzed the transcriptome data. LG, GT, MM and SR wrote the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials
The datasets supporting the conclusions of this article are available at NCBI under the BioProject identifier PRJNA769250 https://www.ncbi.nlm.nih.gov/bioproject/769250.

Declarations

Ethics approval and consent to participate
Not applicable.

Consent for publication
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Competing interests
The authors declare that they have no competing interests.

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