

# Transposable elements contribute to the establishment of the glycine shuttle in Brassicaceae species

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## Abstract

$C_3$ - $C_4$  intermediate photosynthesis has evolved multiple times convergently and independently in the Brassicaceae, although the family lacks *bona fide*  $C_4$  species. Evolution of  $C_3$ - $C_4$  intermediate photosynthesis requires a reconfiguration of gene-regulatory networks, modified spatial expression patterns of multiple genes and ultrastructural adjustments. Mechanisms underpinning the reconfiguration of these networks are mostly unknown. In this study, we use a pan-genomic association approach to identify genomic features that might confer differential gene expression towards  $C_3$ - $C_4$  intermediate traits. We found a strong correlation between transposon insertions in *cis*-regulatory regions and the  $C_3$ - $C_4$  intermediacy trait. Our study revealed 222 orthogroups where presence of a TE within a gene correlates with  $C_3$ - $C_4$  intermediate photosynthesis. In this set, genes involved in the photorespiratory glycine shuttle are enriched. The P-protein of the glycine decarboxylase enzyme complex is known to play a crucial role in the establishment of the photorespiratory glycine shuttle. We found independent transposon insertions in the promoter sequences of the gene encoding the P-protein and hypothesize that these insertions lead to a spatial shift in gene expression. Our findings hint at a pivotal role of TEs in the evolution of  $C_3$ - $C_4$  intermediacy, especially in mediating differential gene expression.

## Introduction

$C_4$  photosynthesis convergently and independently evolved more than 60 times in flowering land plants (Sage et al. 2012).  $C_4$  photosynthesis functions as a biochemical carbon concentrating mechanism that reduces the rate of photorespiration and thereby increases photosynthetic efficiency. Species that perform  $C_4$  photosynthesis are mainly found in warm and dry areas where leaf internal  $CO_2$  concentrations tend to be low and light conditions are frequently saturating, both promoting the oxygenation ratio of Rubisco (Betti et al. 2016, Sage et al. 2012). Although the trait has evolved convergently across multiple plant lineages, the complexity of anatomical, biochemical, and developmental adaptations makes understanding and modifying  $C_4$  photosynthesis a difficult undertaking.

Plants that exhibit  $C_3$ - $C_4$  intermediate phenotypes are promising research subjects to study the early adaptations towards  $C_4$  photosynthesis (Bellasio & Farquhar 2019, Kennedy & Laetsch 1974, Lundgren 2020, Schlüter & Weber 2016).  $C_3$ - $C_4$  intermediate species exhibit specialized anatomical traits that are also observed in  $C_4$  species. They differ from  $C_4$  species in terms of metabolism and do not show a fully integrated  $C_4$  cycle. The  $C_3$ - $C_4$  intermediate trait is characterized by, e.g., a lowered carbon compensation point (CCP), a reduced number of mesophyll cells (MC), chloroplast-rich bundle-sheath cells (BSC) and in some cases an increased vein density (Christin et al. 2011, Dengler et al. 1994, Schlüter et al. 2017).

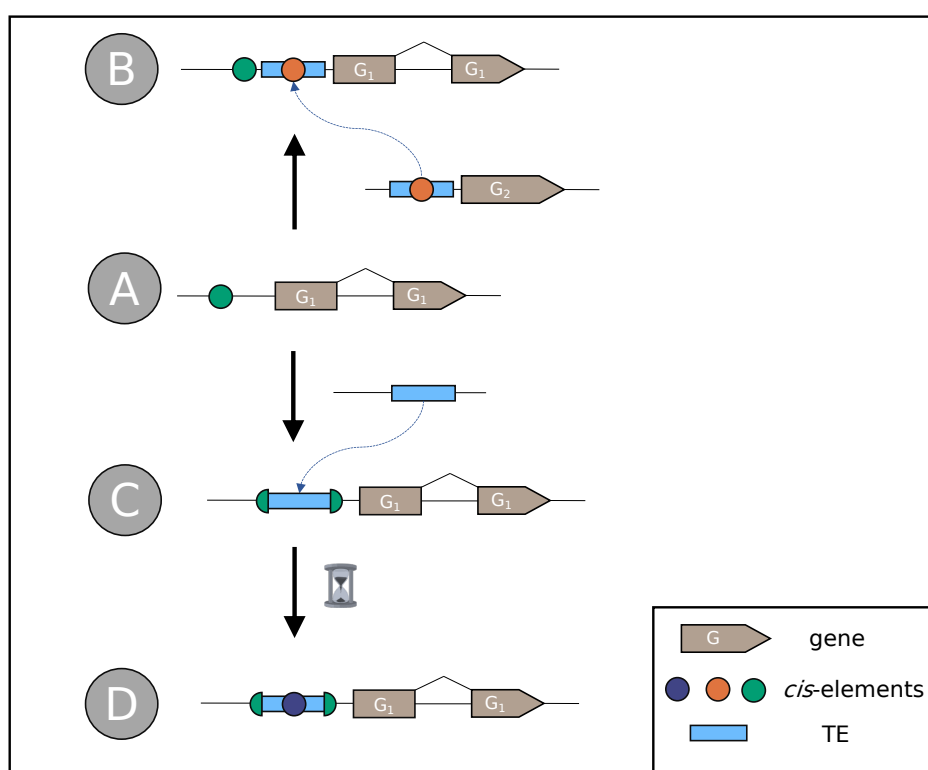
One trait that is commonly shared between multiple C<sub>3</sub>-C<sub>4</sub> intermediate species from independent origins is the photorespiratory glycine shuttle, sometimes referred to as C<sub>2</sub> photosynthesis (reviewed in Schlüter & Weber (2016)). It relies on the BSC-specific decarboxylation of photorespiratory glycine, leading to an elevated CO<sub>2</sub> concentration around Rubisco. The increased partial pressure of CO<sub>2</sub> around the site of its fixation leads to a larger proportion of the Rubisco carboxylation reaction, which suppresses photorespiration and decreases the CCP (Kennedy & Laetsch 1974, Monson & Edwards 1984, Schlüter et al. 2017).

It was shown that the BSC-specific decarboxylation of glycine is caused by the differential activity of the glycine decarboxylase complex (GDC). In C<sub>3</sub>-C<sub>4</sub> intermediate species from the genera *Moricandia*, *Flaveria* and *Panicum*, the P-protein of the GDC was reported to be only present in BSC mitochondria, but not in MC mitochondria (reviewed in Schulze et al. (2016)). This is a notable example of convergent evolution, since these species belong to the distant families Brassicaceae, Asteraceae and Poaceae, respectively. Loss of the GDC P-protein from the MC restricts glycine decarboxylation to the BSC in C<sub>3</sub>-C<sub>4</sub> intermediate species (Morgan et al. 1993, Rawsthorne et al. 1988, Schulze et al. 2016). In C<sub>3</sub> *Flaveria*, the gene encoding the GDC P-protein (GLDP) is present in two differentially regulated copies, *GLDPA* and *GLDPB*. In C<sub>3</sub>-C<sub>4</sub> intermediate *Flaveria* species, the ubiquitously expressed *GLDPB* is downregulated compared to C<sub>3</sub> *Flaveria* species, whereas the BSC-specific *GLDPA* is active (Schulze et al. 2013). In C<sub>3</sub>-C<sub>4</sub> intermediate *Moricandia*, the differential expression of *GLDP1* is thought to be mediated by the loss of one gene copy and differential regulation of the other: whereas *GLDP2* is absent in the Brassicaceae clade containing the C<sub>3</sub>-C<sub>4</sub> intermediate species, *GLDP1* was reported to be differentially expressed by loss of a potential *cis*-element called M-Box. The M-Box confers a basal mesophyll- and bundle sheath expression in the *Arabidopsis thaliana* *GLDP1* promoter (Adwy et al. 2015, 2019) and was not found in *GLDP1* promoter orthologs of C<sub>3</sub>-C<sub>4</sub> intermediate *Moricandia* species (Schlüter et al. 2017). A second *cis*-element, the V-Box, was shown to confer expression to the BSC and is present in all so far analyzed Brassicaceae *GLDP1* promoter sequences (Adwy et al. 2015, 2019). Spatially and temporally differential gene expression is crucial for the evolution of C<sub>3</sub>-C<sub>4</sub> intermediate photosynthesis (Hibberd & Covshoff 2010, Reeves et al. 2017). The underlying genomic variation can be mediated by single-nucleotide polymorphisms (SNPs) or structural variation such as insertions or deletions.

Structural variation can be caused by the activity of mobile genetic elements, also called transposable elements (TEs) or transposons. TEs can be divided into two classes (Wicker et al. 2007) based on their transposition mechanisms: Class I transposons proliferate via a “copy-and-paste” mechanism involving an RNA intermediate, whereas Class II transposons transpose directly via a “cut-and-paste” mechanism. It has been frequently proposed that TEs can contribute greatly to genome evolution and the evolution of novel genetic and phenotypic features (Buchmann et al. 2012, Feschotte 2008, Qiu & Köhler 2020, Wicker et al. 2007). Decades ago, Britten and Davidson put forward that co-option of mobile sequences containing gene regulatory elements can connect genes to the same gene regulatory networks (Britten & Davidson 1971). The co-option of TEs for regulatory purposes is called “exaptation” (Brosius & Gould 1992). With today’s vast amount of available genomic data, a deeper understanding of the role of transposable elements in genetic regulation allows linking genomic mechanisms with the evolution of complex traits.

TEs can rewire gene regulatory networks using different modes of action and influence the interplay of regulatory proteins (*trans*-elements) and the DNA sequences they are binding to (*cis*-elements). One mode is the exaptation of a *cis*-regulatory element (CRE) from other genes (Fig 1). If the CRE inside a TE is copied from one gene and retained by the other gene, both genes are afterwards controlled by a mutual CRE, thus connected in a gene regulatory network (Fig 1, B). In contrast to this, TE integration into a CRE can suppress its function, either by interrupting the CRE sequence or altering the chromatin state of the respective CRE locus (Fig. 1, C) (Feschotte 2008). Another possibility is the *de novo* generation of new CRE by point mutations in degrading TEs (Fig. 1, D). New CREs, e.g. a 10-mer promoter element, can arise by random point mutations in between 700,000-4.8 million years (Behrens & Vingron 2010).

Several examples for the role of TEs in rewiring gene regulatory networks in plants have been reported: in rice, the *mPing* DNA transposon was found preferentially in the 5' region and was associated with the upregulation of stress response genes (Naito et al. 2009). In Brassicaceae, the evolution of heat-tolerance was linked to the activity of *Copia* retrotransposons containing heat-shock factor binding elements (Pietzenuk et al. 2016). Furthermore, TEs were also connected to endosperm development, e.g., the distribution of the PHEREs1 MADS-box transcription factor binding motifs by *Helitron* transposons in *Arabidopsis thaliana*



**Figure 1. Schematic illustration of gene regulation rewiring by TE exaptation.** A: The hypothetical gene G1 is controlled by a CRE (green dot). B: Gene G2 is regulated by a different CRE (orange dot) located within a TE (blue box). Upon transposition of the TE to the upstream region of G1, G1 might co-opt the function of the orange CRE, thus connecting G1 and G2 to the same gene regulatory network. C: TE transposition can also lead to destruction or suppression of the CRE. D: During TE decay, new CREs (blue dot) might occur through accumulation of point mutations.

(Batista et al. 2019). The *Youren* miniature inverted-repeat TE (*MITE*) was shown to be transcribed in rice endosperm, putatively mediated by a NUCLEAR FACTOR Y binding motif in the vicinity of the 5' terminal inverted repeat (TIR) of *Youren* (Nagata et al. 2022).

Investigating potential roles for TEs in the evolution of C<sub>4</sub> photosynthesis, Cao et al. (2016) analyzed 40 C<sub>4</sub> gene orthologs between rice and maize for the presence of BSC-specific promoter motifs. Of over 1,000 promoter motifs that were differentially distributed between C<sub>3</sub> and C<sub>4</sub> orthologs, more than 60 % were associated with TEs and potentially co-opted by TE integration. These motifs may originate from non-photosynthetic genes and transposed to C<sub>4</sub> genes, which connected gene regulatory networks. The authors showed that TEs play a significant role in the evolution of C<sub>4</sub> photosynthesis in maize. The study of Cao et al. (2016) focused on evolutionary distant grasses, which makes it difficult to draw conclusions about the early evolutionary events towards C<sub>4</sub> photosynthesis.

Here, we focus on TE-mediated exaptation of CREs in C<sub>3</sub>-C<sub>4</sub> intermediate Brassicaceae species that mediate decisive steps in the evolution of the photorespiratory glycine shuttle and could play a crucial role in C<sub>4</sub> evolution. We analyzed independent evolutionary events in closely related species from one plant family, the Brassicaceae. The Brassicaceae family contains multiple important and well-studied model plant species such as *A. thaliana*, *Arabis alpina* as well as relevant crop and vegetable plants such as *Brassica oleracea* and *Diplotaxis tenuifolia* (arugula). Current studies (Guerreiro et al., in preparation) described at least five independent origins of C<sub>3</sub>-C<sub>4</sub> intermediate photosynthesis in this family.

To test whether TE insertions might be involved in establishing the C<sub>3</sub>-C<sub>4</sub> intermediate photosynthetic trait, we performed a pan-genomic association study to analyze the TE landscape of 15 Brassicaceae species. We

tested for correlations between TE positions and the presence of C<sub>3</sub>-C<sub>4</sub> intermediate traits. Specifically, we  
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searched for correlations between the presence or absence of upstream co-occurring TEs with the CCP. In  
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this unbiased approach, we aimed at finding genes that retained upstream TEs selectively only in C<sub>3</sub>-C<sub>4</sub>  
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intermediate plants. Based on the results of our analysis, we examined the promoter orthologs of relevant  
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photorespiratory genes in closer detail to reveal the influence of TE insertions towards the establishment of  
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C<sub>3</sub>-C<sub>4</sub> photosynthesis traits. Here, we present evidence that the insertion of TEs in *cis*-regulatory regions are  
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associated with the establishment of C<sub>3</sub>-C<sub>4</sub> photosynthesis in the Brassicaceae.  
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## Materials and Methods

### Genomes and carbon compensation points

The genomes for *Brassica gravinae* (Bg), *Brassica tournefortii* (Bt), *Carrichtera annua* (Ca), *Diplotaxis erucoides* (De), *Diplotaxis tenuifolia* (Dt), *Diplotaxis viminea* (Dv), *Hirschfeldia incana* (accessions HIR1 and HIR3), *Moricandia nitens* (Mn) and *Moricandia suffruticosa* (Ms) were obtained from Guerreiro *et al.* (in preparation). The genome of *Arabis alpina* (Aa) was obtained from Jiao *et al.* (2017). The genome of *Arabidopsis thaliana* (At) was obtained from Lamesch *et al.* (2012). The genome sequences of *Moricandia arvensis* (Ma) and *Moricandia moricandoides* (Mm) were obtained from Lin *et al.* (2021). The genome assembly for *Brassica oleracea* (Bo) was obtained from Parkin *et al.* (2014). The genome for *Gynandropsis gynandra* (Gg) was obtained from Hoang *et al.* (2022). A full list of species names and accession number and sources can be found in Supplemental Table 1. Gas exchange data was obtained from Schlüter *et al.* (in preparation). Data for the phylogenetic tree was obtained from Guerreiro *et al.* (in preparation).

### Gene annotation

Consistent structural gene annotations were generated for each species with *Helixer* (Stiehler *et al.* 2020) using the hybrid convolutional and bidirectional long-short term memory model, *HybridModel*, specifically the trained instance of `land_plant_v0.3_m_0100`. This was followed by post-processing the raw predictions into final primary gene models with *Helixer Post* (Bolger 2022, personal communication). Default parameters were used except as specified below. During data preparation, the subsequence length was increased to 106920 bp. During inference, overlapping was activated using a core-length of 53460 bp and an offset of 13365 bp. Phase prediction was used. During post processing, a window size of 100 bp, an edge threshold of 0.1, a peak threshold of 0.8, and a minimum coding length of 60 bp were used. The exact code-state can be found at the following commits: *Helixer bb840b4*, *Geenuff 1f6cffb* and *Helixer Post 08c6215*, while the process is compatible with all releases beginning with v0.3.

### Annotation of transposable elements

TEs were *de novo* annotated using *EDTA* 1.9.9 (Ou *et al.* 2019) using the `-anno 1` and `-sensitive 1` flags. For the calculation of genomic composition (Fig. 2, Fig. 3), intact and fragmented TEs were used. To reduce the influence of false-positive hits, the pan-genomic gene-TE association study was performed for intact TEs only. The LTR insertion time was calculated using

$$t_{insertion} = \frac{1 - LTRidentity}{2 \cdot \mu}$$

assuming a neutral mutation rate of  $\mu = 1.4 \cdot 10^{-8}$  substitutions per site per year (Cai *et al.* 2018).

### Analysis of differential transposable element insertion

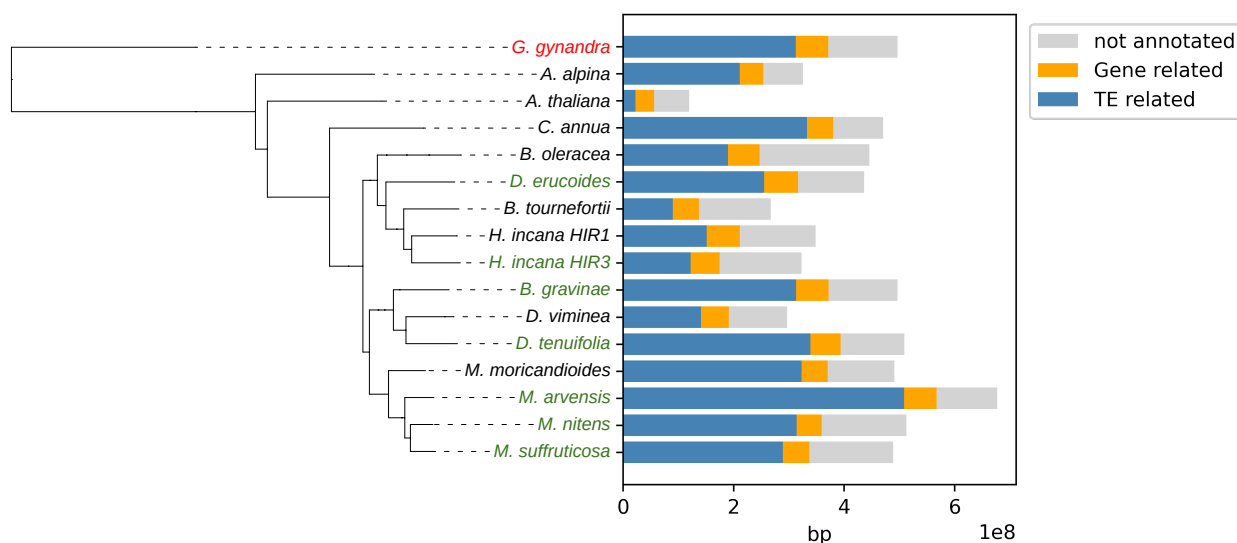
All downstream analyses were performed using *Python* 3.6 including *pandas* 1.2.4, *numpy* 1.20.1, *matplotlib* 3.4.1, *scikit-learn* 0.24.1, *scipy* 1.6.2 and *statsmodels* 0.12.2. All scripts are available on GitHub. The annotation files for genes and intact TEs were compared for each species. TEs were considered co-occurring

with genes if their position matched one of the five cases described in Fig. 5. Genes with TEs within 3000 bp upstream were filtered and collapsed into orthogroups (OGs) using *OrthoFinder* 2.5.4 (Emms & Kelly 2015). For each OG, the presence or absence of an upstream TE was correlated to the CCP using one-way ANOVA. Each OG was functionally annotated using *Mercator4* 4.0 (Schwacke et al. 2019). A randomly chosen representative protein sequence from each OG was uploaded to the *Mercator4* web tool for functional annotation. Enrichment of *Mercator* bins for genes with correlating upstream TEs was calculated using Fisher's exact test. The identities of TEs in the *GLDP1* promoter were validated using the *CENSOR* webtool (Kohany et al. 2006).

## Results

To screen genomic features of potential relevance to the evolution of the C<sub>3</sub>-C<sub>4</sub> photosynthesis trait, we conducted a pan-genomic association study of eight C<sub>3</sub> Brassicaceae species, seven C<sub>3</sub>-C<sub>4</sub> intermediate Brassicaceae species from five independent origins, and one C<sub>4</sub> Cleomaceae as an outgroup species. The five independent origins of C<sub>3</sub>-C<sub>4</sub> intermediate photosynthesis can be found in the *Moricandia arvensis*, -*nitens*, and -*suffruticosa* monophylum, as well as in *Diplotaxis erucoides*, *Diplotaxis tenuifolia*, *Brassica gravinae* and *Hirschfeldia incana* HIR3 (Fig. 2).

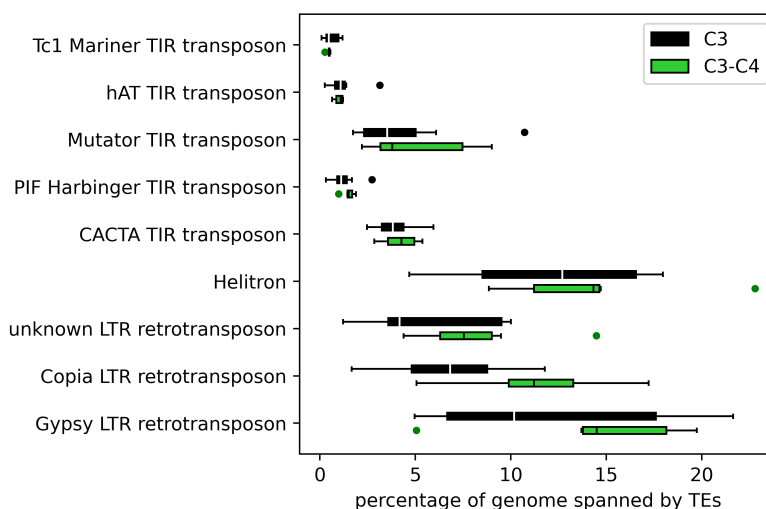
The species panel exhibits genome sizes ranging from 120 Mbp in *Arabidopsis thaliana* to 677 Mbp in *Moricandia arvensis*. We found no significant difference in genome size between species exhibiting either the C<sub>3</sub> or C<sub>3</sub>-C<sub>4</sub> intermediate photosynthesis phenotype (Fig 2; one-way ANOVA p>0.05). We next *de novo* annotated TEs using the *EDTA* pipeline (Ou et al. 2019). Overall, the annotated fragmented and intact transposons made up between 18 % of the genome in *Arabidopsis thaliana* and 75 % in *Moricandia arvensis*. We observed differences in genome size and TE content also in closely related species, e.g. between *M. arvensis* and *M. moricandoides* or between *B. gravinae* and *D. viminea*. Furthermore, we observed that differences in genome size are mainly due to the different TE content.



**Figure 2. Phylogeny and genomic composition of 15 selected Brassicaceae species and the Cleomaceae outgroup.** C<sub>3</sub>-C<sub>4</sub> intermediate species are highlighted in green, the C<sub>4</sub> outgroup *G. gynandra* is highlighted in red. TE-related nucleotides are defined as spanning intact and fragmented transposons.

Class I type retrotransposons represented the majority of identified TEs in C<sub>3</sub> and C<sub>3</sub>-C<sub>4</sub> species (Fig. 3). Within each analyzed genome, between 60 % and 68 % of all annotated TEs were Class I retrotransposons for C<sub>3</sub> and C<sub>3</sub>-C<sub>4</sub> species, respectively. Among all analyzed species, the proportion of TE classes in the

genomes varied greatly (Fig. 3; Supplemental Table S2). In the genomes analyzed here, the TE Class II was dominated by TEs from the *Helitron* group, making up between five and 20 percent of the genome (Fig. 3). The percentage of the genome made up of TEs from the different classes varied between the photosynthesis types, with a significantly higher amount of TEs in C<sub>3</sub>-C<sub>4</sub> genomes (two-way ANOVA, p = 0.013). 156  
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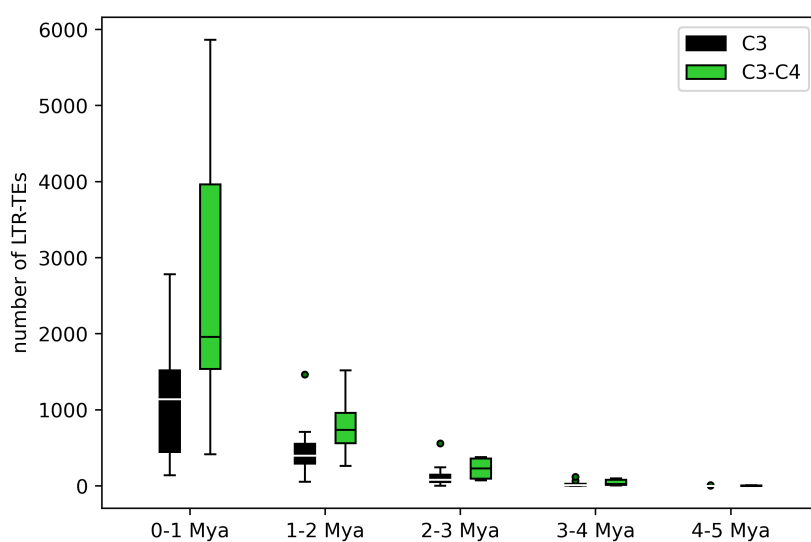


**Figure 3.** Boxplot indicating the percentage of the genome spanned by nine classes of intact and fragmented TEs in eight C<sub>3</sub> and six C<sub>3</sub>-C<sub>4</sub> intermediate species. The y-axis shows the TE classes, the x-axis indicates the fraction of the genome made up by the respective TE class. Black boxes depict C<sub>3</sub> species, green boxes depict C<sub>3</sub>-C<sub>4</sub> intermediate species. The boxes start and end at the lower/upper quartiles of the data, the whiskers extend to the upper and lower range of the data. Outliers are indicated by circles. The median is indicated by a white or black bar for C<sub>3</sub> or C<sub>3</sub>-C<sub>4</sub> intermediate species, respectively.

To analyze recent increases of TE activity and their potential roles in the evolution of C<sub>3</sub>-C<sub>4</sub> intermediate photosynthesis, we determined the LTR insertion times (Fig. 4; Supplemental Table S3). *LTR retriever*, which is the LTR annotation tool of the *EDTA* pipeline, detected LTR transposons down to a threshold for repeat identity of 91 %. Assuming a neutral mutation rate of  $\mu = 1.4 \times 10^{-8}$  substitutions per site per year (Cai et al. 2018), LTR insertion times could thus be dated back to a maximum of 4 million years. Both C<sub>3</sub> and C<sub>3</sub>-C<sub>4</sub> intermediate species showed mainly the same pattern of LTR bursts. There was an increased frequency for LTR-TEs younger than two million years. The increase was more pronounced for C<sub>3</sub>-C<sub>4</sub> intermediate species, which was largely due to the high number of young LTR-TEs in *M. arvensis*. Statistical analysis revealed a significant correlation between the age distribution of LTR-transposons and the photosynthesis phenotype (2-way ANOVA, p = 0.033). 160  
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We next analyzed the differential co-occurrence of TEs with genes. We reduced our species set to five C<sub>3</sub> species and six C<sub>3</sub>-C<sub>4</sub> intermediate species based on genome quality and to analyze a comparable number of C<sub>3</sub> and C<sub>3</sub>-C<sub>4</sub> species (At, Bo, Bg, De, Dt, Dv, HIR1, HIR3, Ma, Mm, Mn). Co-occurring TEs were defined as follows (Fig. 5): (I) the TE starts or ends in a 3,000 bp window upstream of the gene (upstream), (II) the TE starts or ends in a 3,000 bp window downstream of the gene (downstream), (III) the TE spans a whole gene (spanning), (IV) the TE starts in a gene (start), or (V) the TE ends in a gene (end).

Genes that were spanned by TEs (III) were mostly open reading frames related to the transposase genes of the respective TEs. Genes with overlapping TEs (IV and V) might have broken coding sequences and may result from imprecise annotations. Across the selected eleven species, 61,092 TEs were identified to be co-occurring with a gene in at least one species, whereas 22,053 co-occurring TEs belonged to C<sub>3</sub> and 39,039 co-occurring TEs belonged to C<sub>3</sub>-C<sub>4</sub> species. In both C<sub>3</sub> and C<sub>3</sub>-C<sub>4</sub> intermediate species, around 70 % of the TEs co-occurring with genes were located up- or downstream of the gene (Fig. 5). Analyzing

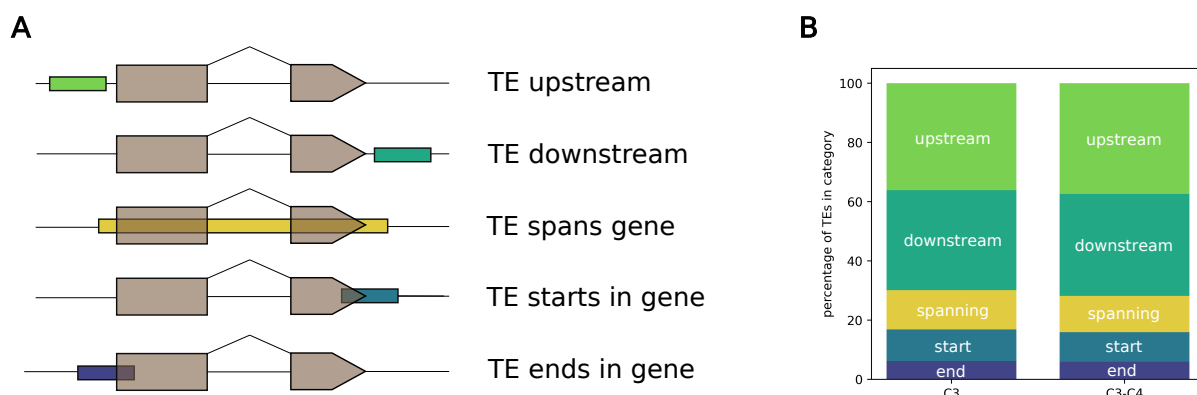


**Figure 4. Boxplot of LTR-TE insertion times for eight C<sub>3</sub> and six C<sub>3</sub>-C<sub>4</sub> intermediate species.** The x-axis shows the insertion time in bins of 1 million years before today (Mya). The y-axis depicts the number of identified LTR-TEs calculated to be inserted within this timeframe. The boxes start and end at the lower/upper quartiles of the data, the whiskers extend to the upper and lower range of the data. Outliers are indicated by circles. The median is indicated by a white or black bar for C<sub>3</sub> or C<sub>3</sub>-C<sub>4</sub> intermediate species, respectively. Calculation was performed using the LTR similarity of each LTR-TE and a neutral mutation rate of 1.4\*10<sup>-8</sup> substitutions per site per year. Black boxes represent C<sub>3</sub> species, green boxes represent C<sub>3</sub>-C<sub>4</sub> species.

potentially exapted CREs, we further analyzed TEs in the up to 3000 bp 5' region of the gene. A total of 22,560 genes with co-occurring upstream TEs were collapsed into 12,087 OGs using *OrthoFinder* (Emms and Kelly, 2015). By this, we filtered out false positive gene models from the *Helixer* annotation and allowed for between-species comparisons. For each of these OGs, one-way ANOVA was employed, correlating the presence or absence of a co-occurring upstream TE with the CCP of the respective species. We identified 222 OGs where the co-occurrence of one of the orthologs with an upstream TE correlated with the CCP ( $p \leq 0.05$ ; Tab. 1, Supplemental Table S4). Among the ten OGs with the lowest p-value were genes involved in photorespiration such as GLYCOLATE OXIDASE or the genes encoding the T- and P-subprotein of the glycine decarboxylase complex (Fig. 6A). Strikingly, in the ten OGs with the lowest p-values, the C<sub>3</sub>-C<sub>4</sub> intermediate OGs exhibited upstream TEs, whereas the C<sub>3</sub> OGs lacked upstream OGs. Thus, in the OGs with the lowest p-values, we observed a “gain” in upstream TEs (Fig. 6A).

In the list of OGs with a correlation between the CCP and the presence of upstream TEs, three photorespiratory genes occurred (*GOX*, *GLDP*, *GLDT*). To quantify putative enrichment of certain gene ontologies, each OG was functionally annotated with a *Mercator* bin. Statistical enrichment analysis using Fisher's exact test revealed that the *Mercator* bin “Photosynthesis.Photorespiration” ( $p = 0.000904$ ) was enriched in the set of genes that co-occur with upstream transposons (Tab. 2). The occurrence of this *Mercator* bin was increased twenty-fold over the background, which is higher than for any other analyzed *Mercator* bin (Tab. 2; Supplemental Table S5).

From the enriched photorespiratory genes, we selected *GLDP*, the gene encoding the P-protein of the GDC, for a deeper analysis. Only one *GLDP* gene copy is present in species from the Brassicaceae tribe that contains all known C<sub>3</sub>-C<sub>4</sub> intermediate species of the Brassicaceae (Schlüter et al. 2017). In contrast, the other two photorespiratory genes with correlating upstream TEs (Tab. 1; Fig. 6A) are found in higher copy numbers, which complicates a detailed genetic analysis. Furthermore, it is known that the differential expression of



**Figure 5.** **A:** Different contexts of TEs co-occurring with genes. **B:** Bar charts indicating the fractions of TE co-occurring with genes within five contexts: starting or ending in a gene (start / end), spanning the gene (over) or residing within a 3000 bp window upstream or downstream the gene.

*GLDP* contributes to the establishment of the photorespiratory glycine shuttle (Monson & Edwards 1984, Rawsthorne et al. 1988, Schulze et al. 2013) and several studies about the underlying regulatory genetics have been conducted (Adwy et al. 2015, 2019, Dickinson et al. 2020, Schulze et al. 2016).

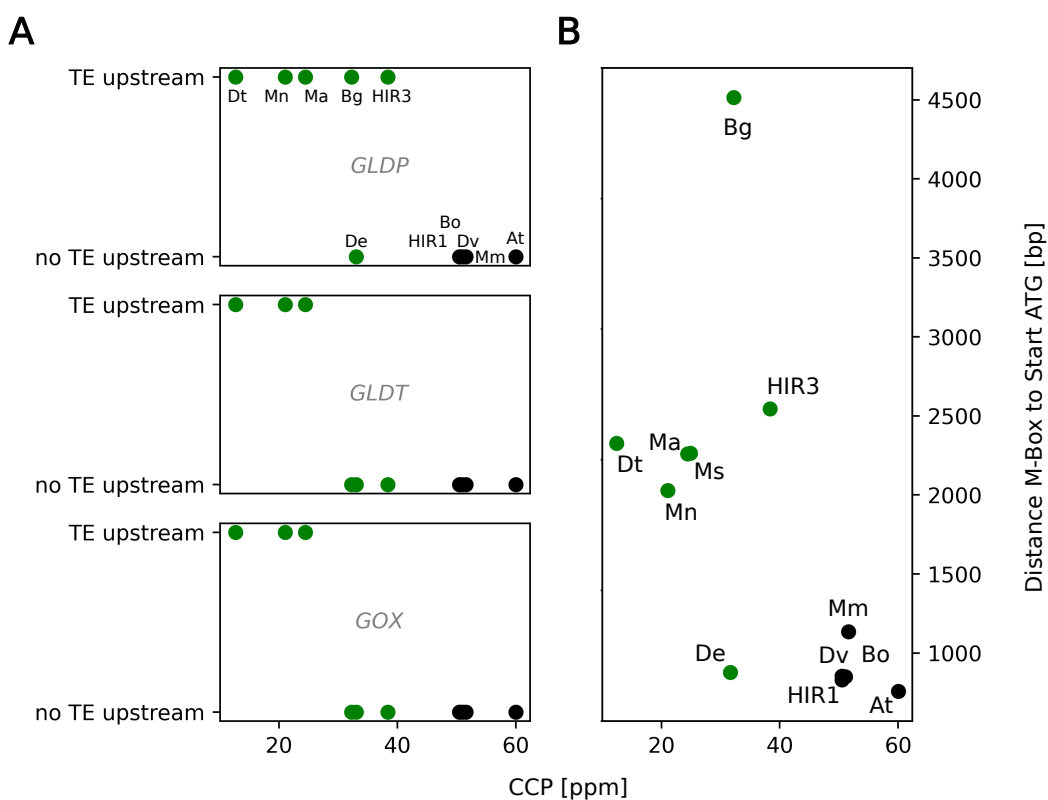
We found three independent TE insertions in the promoter of C<sub>3</sub>-C<sub>4</sub> intermediate *GLDP1* orthologs. In *Diplotaxis tenuifolia* a Mutator TE starts at 1970 bp upstream of the *GLDP1* start codon. In *Hirschfeldia incana* HIR3 a TE of the *Helitron* class is located around 2240 bp upstream. In orthologs from the monophyletic clade *Moricandia arvensis*, *Moricandia nitens* and *Moricandia suffruticosa* a MITE DNA transposon was detected, starting 1950 bp upstream of the *GLDP1* start codon. We calculated the minimum timespan since the MITE insertion by pairwise multiple sequence alignments of the MITE in the three *Moricandia* *GLDP1* promoters using the neutral mutation rate formula that was also employed for the calculation of LTR ages. We found the *GLDP1* promoter MITE to be at least 6.5 million years old.

All three independent TE insertions are located around 100 bp downstream of the M-Box promoter motif. This motif was previously hypothesized to confer mesophyll cell (MC) expression (Adwy et al. 2015) since truncation of the motif from the *AtGLDP1* promoter shifted GUS activity from the whole leaf apex to the veins. Furthermore, the M-Box was reported to be lost in C<sub>3</sub>-C<sub>4</sub> intermediate *Moricandia* species (Adwy et al. 2019). However, upon closer inspection, we found a highly conserved M-Box motif in all analyzed Brassicaceae genomes. Notably, the M-Box was shifted upstream due to the TE insertion in C<sub>3</sub>-C<sub>4</sub> species with the exception of *D. erucoides* (Fig. 6B, Fig. 7, Supplemental Table S6). In *Brassica gravinae*, the EDTA pipeline did not annotate an upstream transposon. However, we found a large insertion of unknown origin in the *B. gravinae* *GLDP1* promoter. This insertion is larger than the three reported TE cases but could be found in a similar position compared to the other *GLDP1* promoter insertions of TE origin (Fig. 7). In the *GLDP1* promoter of C<sub>3</sub>-C<sub>4</sub> intermediate species *D. erucoides* no insertion could be found.

From five analyzed C<sub>3</sub>-C<sub>4</sub> *GLDP1* promoters we found a large insertion behind the conserved M-Box in four cases (monophyletic C<sub>3</sub>-C<sub>4</sub> intermediate *Moricandia* clade, *Diplotaxis tenuifolia*, *Brassica gravinae* and *Hirschfeldia incana* HIR3; Fig. 6B). Out of these four cases, we found evidence for the sequence being a TE in three cases (Fig. 7).

## Discussion

Evolution of new traits such as C<sub>3</sub>-C<sub>4</sub> and C<sub>4</sub> photosynthesis requires the differential regulation of multiple genes. This includes differential expression between MSC and BSC tissue and the installation of light-responsiveness for genes of the core metabolism (reviewed in Hibberd & Covshoff (2010)). In many cases, the evolution of differential gene regulation takes place in promoter sequences, either by introduction or suppression of *cis*-elements.



**Figure 6. A:** Scatter plot for three photorespiratory genes with significant co-associated upstream TEs. The y-axis indicates the presence of an upstream TE (yes/no), the x-axis shows the carbon compensation point. Abbreviations: GLDP/GLDT: P/T-protein of the GLYCINE DECARBOXYLASE COMPLEX; GOX: GLYCOLATE OXIDASE. **B:** Scatter plot for the different architectures of the *GLDP1* promoter. The y-axis indicates the distance between the conserved M-Box sequence and the *GLDP1* start site. Each dot represents a species C<sub>3</sub> species are shown in green, C<sub>3</sub>-C<sub>4</sub> intermediate species are shown in black. Species name abbreviations: At.: *Arabidopsis thaliana*, Bg: *Brassica gravinae*, Bo: *Brassica oleracea*, De: *Diplotaxis erucoides*, Dt: *Diplotaxis tenuifolia*, Dv: *Diplotaxis viminea*, HIR1: *Hirschfeldia incana* HIR1, HIR3: *Hirschfeldia incana* HIR3, Ma: *Moricandia arvensis*, Mm: *Moricandia moricandoides*, Mn: *Moricandia nitens*, Ms: *Moricandia suffruticosa*

A few *cis*-elements for MC specificity were found, such as the MEM1 motif from the *Flaveria trinervia* phosphoenolpyruvate carboxylase gene (Gowik et al. 2017) or the M-Box sequence in Brassicaceae (Adwy et al. 2015, Dickinson et al. 2020).

TEs have the potential to deliver or suppress *cis*-elements upon insertion in a target promoter. Furthermore, they can generate antisense transcription, interrupt or generate heterochromatic regions or serve as raw material for the *de novo* evolution of new *cis*-elements (reviewed in Feschotte (2008)). The role of TEs in the evolution of C<sub>4</sub> photosynthesis has been scarcely investigated. This study presents the *bona fide* first pan-genomic association study correlating the co-occurrence of TEs and genes with the presence of a certain plant phenotype. We analyzed the role of differential TE landscapes in 15 Brassicaceae species. Firstly, we investigated whether genome size and TE content correlate with the presence of the C<sub>3</sub>-C<sub>4</sub> photosynthesis phenotype. Across our species panel a variety of genome sizes is present (Fig. 2), but no correlation of genome size to the presence of the photosynthesis trait was detectable. In our data, different levels of heterozygosity in the sequenced species may confound these findings and genome size estimations have to be handled with

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**Table 1.** Selected subset of ten genes with upstream TEs with the lowest p-values for their association with the CCP.

Gene Name	AGI locus code	p-value
transcription factor (WRKY)	AT5G49520	0.0017
homeobox leucine zipper protein	AT1G27050	0.0017
polyketide cyclase/dehydrase/lipid transport superfamily protein	AT5G06440	0.0017
Em-like protein GEA6	AT3G51810	0.0023
regulatory protein (FLZ) of SnRK1 complex	AT5G49120	0.0023
solute transporter (UmamiT)	AT4G15540	0.0023
Protein GLUTAMINE DUMPER 2	AT4G25760	0.0023
aminomethyltransferase component T-protein of glycine cleavage system	AT1G11860	0.0023
glycolate oxidase	AT3G14420	0.0023
glycine dehydrogenase component P-protein of glycine cleavage system	AT4G33010	0.0023

**Table 2.** Results from two-sided Fisher's exact test for the enrichment of *Mercator* bins within the set of orthologs with significant upstream transposons.

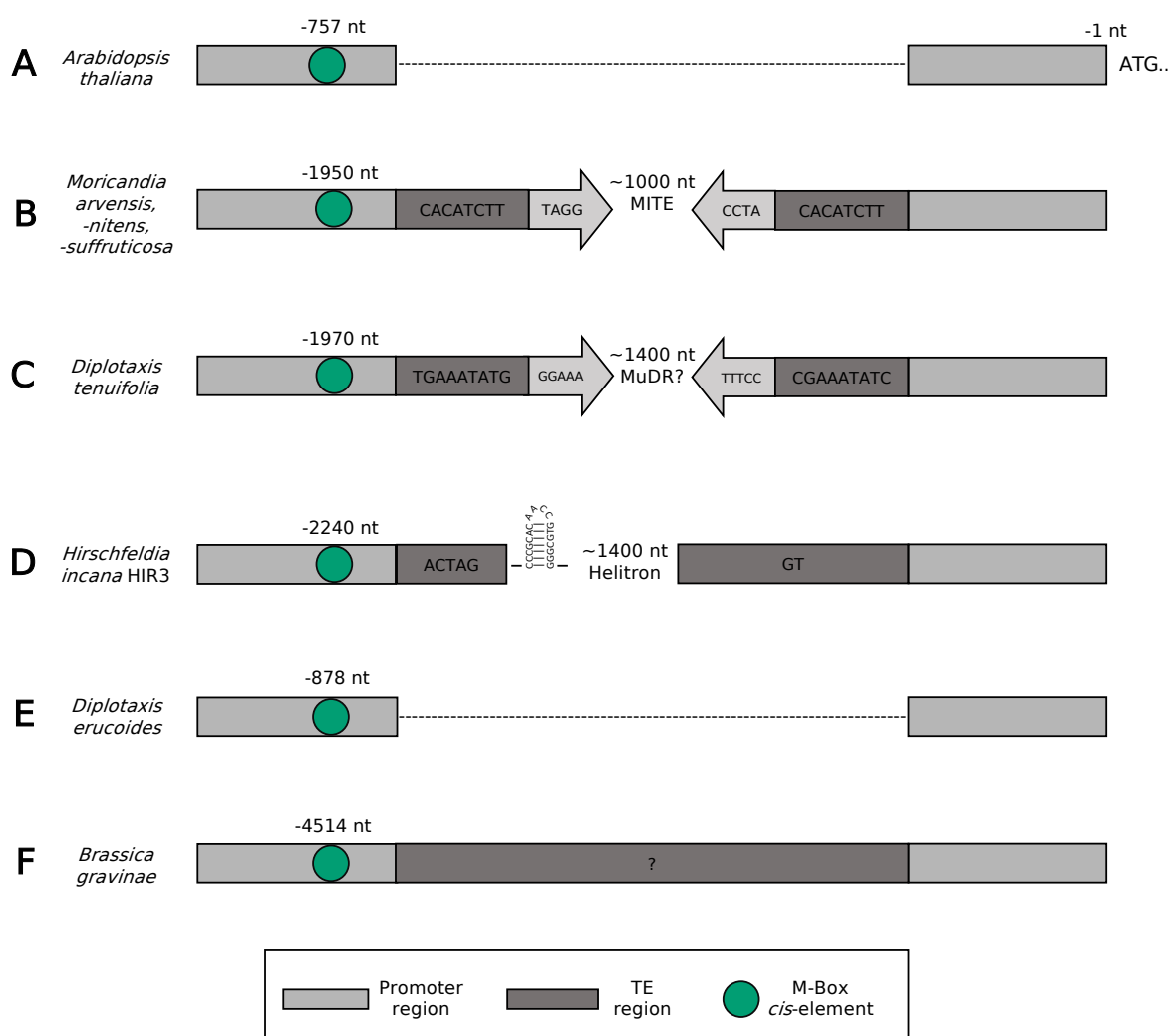
Mercator Bin	OGs with p>0.05	OGs with p<0.05	p-value	Odds ratio
Photosynthesis.Photorespiration	8	3	0.000904	20.3
Not assigned.Not annotated	3132	40	0.004306	0.6
Protein biosynthesis.Ribosome biogenesis	122	7	0.009691	3.1
Multi-process regulation.SnRK1-kinase regulation	7	2	0.011104	15.4
Protein homeostasis.Protein quality control	36	3	0.034403	4.5

care.

Within the Brassicaceae family species exhibiting C<sub>3</sub>-C<sub>4</sub> intermediate traits can only be found in the Brassiceae tribe. Notably, species from this tribe seem to have undergone recent polyploidization events (Walden et al. 2020) and exhibit larger genome sizes than species from neighboring tribes (Lysak et al. 2009). However, genome size alone does not reflect the complexity of a species, as described by the “C-value paradox” (Thomas 1971).

Next, we analyzed the proportion of TEs in the individual genomes. Our estimation of TE proportions is consistent with previously analyzed Brassicaceae genomes (Liu et al. 2020, Mirouze & Vitte 2014) and the *Gynandropsis gynandra* genome (Hoang et al. 2022). We found a significant correlation between the photosynthesis phenotype and the proportion of the genome occupied by TEs in the respective species. Genome size and TE content also vary between closely related species, which may lead to the assumption that these differences are not only due to phylogeny. In all species, we found a recent burst in LTR-TE activity that is consistent with other studies (e.g. Cai et al. (2018)). The recent sharp increase in LTR-TE bursts in C<sub>3</sub>-C<sub>4</sub> species comes mainly from *Moricandia arvensis* and might rather be due to high heterozygosity of LTR-containing genomic regions (Fig. 4). Although we found a significant correlation of LTR content and age and the C<sub>3</sub>-C<sub>4</sub> intermediate phenotype, we cannot ultimately conclude that LTR transposon bursts contributed to the evolution of the C<sub>3</sub>-C<sub>4</sub> intermediacy. Our LTR age analysis is limited to an LTR age of 4 million years. Given the estimated divergence time of 2-11 million years for C<sub>3</sub> and C<sub>3</sub>-C<sub>4</sub> intermediate *Moricandia* species (Arias et al. 2014), our analysis of LTR insertion times will miss the contribution of older LTRs to the evolution of C<sub>3</sub>-C<sub>4</sub> intermediate traits. Based on sequence identity between the C<sub>3</sub>-C<sub>4</sub> intermediate *Moricandia GLDP1* promoters, we estimate the age of the MITE in the *Moricandia GLDP1* promoter to be at least 6.5 million years. This falls well within the proposed divergence time C<sub>3</sub> and C<sub>3</sub>-C<sub>4</sub> intermediate *Moricandia* species of 2-11 million years (Arias, 2014).

The proportion of TEs in the genome can be seen as a proxy for gene duplication events, a process whose role in evolution has been extensively analyzed in the Brassicaceae (Cerbin & Jiang 2018, Oh & Dassanayake 2019, Walden et al. 2020). Evidence for gene duplication events towards C<sub>4</sub> photosynthesis exist, but it is yet unclear which events in the gradual evolutionary pathway along the C<sub>3</sub>-C<sub>4</sub> intermediate stages may require



**Figure 7. Schematic representation of the *GLDP1* promoter region.** "ATG..." depicts the start site of the *GLDP1* gene. Dark grey boxes represent characteristic TE sites such as target site duplications or the *Helitron* insertion sites. Grey arrows depict terminal inverted repeat motifs. The M-Box motif is highlighted as a green circle. In C<sub>3</sub> species such as *Arabidopsis thaliana* no TE is annotated in the promoter sequence, leading to a low spacing between the M-Box and the *GLDP1* start site (A). In the C<sub>3</sub>-C<sub>4</sub> intermediate *Moricandia* species, a MITE TE begins around 1950 bp upstream of the *GLDP1* start codon (B). In *Diplotaxis tenuifolia*, a Mutator TE begins 1970 bp upstream (C). In *Hirschfeldia incana* HIR3 a *Helitron* with a highly conserved hairpin loop structure is inserted around 2240 bp upstream (D). Although being a C<sub>3</sub>-C<sub>4</sub> intermediate species, the *Diplotaxis erucoides* *GLDP1* promoter did not have an insertion behind the M-Box. (E). In *Brassica gravinae* a large insertion of unknown origin could be found behind the M-Box region (F).

sub- or neofunctionalized genes (Emms et al. 2016, Gowik et al. 2011, Huang et al. 2021).

In the descriptive whole-genome view, we observed correlations between TE content and age and the C<sub>3</sub>-C<sub>4</sub> intermediate phenotype. Yet, however, there is an individual TE pattern even in closely related lines (Fig. 2). We therefore conclude that the role of TE activity may have an influence on C<sub>3</sub>-C<sub>4</sub> evolution, but not necessarily via means of general TE activity (TE outbursts or TE purging) but rather via selective TE

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insertions to relevant genes or upstream regions. To analyze this, we employed a pan-genomic *de novo* transposon-gene association study, where we correlated the co-occurrence of TEs with genes to the presence of a C<sub>3</sub>-C<sub>4</sub> intermediate phenotype.

In both C<sub>3</sub> and C<sub>3</sub>-C<sub>4</sub> intermediate species, more than 70 % of the analyzed co-occurring TEs were upstream or downstream of the respective co-occurring gene or spanning the gene. This is biologically plausible, since TEs crossing gene borders may disturb gene function and intergenic regions can harbor transposable elements (Buchmann et al. 2012). Therefore we are confident that our association study, relying on *in silico* predicted genes and TEs, is highly robust.

Differential gene regulation mediated by variation in upstream regions was shown to be a driver of C<sub>4</sub> traits in multiple, well documented cases (Adwy et al. 2015, Gowik et al. 2017, Williams et al. 2015, Wiludda et al. 2012). Our analysis revealed 222 OGs with an upstream TE that correlates with the presence of a C<sub>3</sub>-C<sub>4</sub> intermediate phenotype (Fig. 7; p<0.05). Enrichment analysis of *Mercator* bins for this set of genes revealed an enrichment of the codes “Multi-process regulation.sucrose non-fermenting-related kinase (SnRK1) regulation” and “Photosynthesis.Photorespiration”. SnRK1 was shown to act as a central regulator of starvation metabolism that mediates energy homeostasis between organelles (Wurzinger et al. 2018). During nutrient starvation, SnRK1 subcomplexes were found to regulate the differential expression of over 600 target genes (Baena-González et al. 2007). Strikingly, ultrastructural adjustments and re-localization of the GDC P-protein to the BSC were demonstrated as a result of nitrogen starvation in the C<sub>3</sub>-C<sub>4</sub> intermediate species *Chenopodium album* (Oono et al. 2022).

There is a clear bias of TE retention upstream of photorespiratory and SnRK1-regulatory genes in C<sub>3</sub>-C<sub>4</sub> intermediate species, although with a small effect size (3 out of 11 OGs with p<0.05 for “Photosynthesis. Photorespiration”; 2 out of 9 OGs with p<0.05 for “Multi-process regulation.SnRK1 regulation”; see Table 2). We hypothesize that TE retention upstream of these OGs has functional consequences such as differential gene expression, putatively due to the co-option of new- or suppression of existing CREs. Strikingly, the set of genes with significant upstream TEs contains multiple genes involved in photorespiration, such as those encoding GLYCOLATE OXIDASE (GOX) as well as the T- and P- proteins of the glycine decarboxylase complex (GLDT/GLDP). The modification of photorespiration is an important step towards the establishment of the glycine shuttle. The enrichment of TE insertions upstream of photorespiratory genes in C<sub>3</sub>-C<sub>4</sub> intermediates is one hint that TEs play a significant role in the introduction of the glycine shuttle.

*GLDP* is a well-characterized example for differential gene expression at the early stages of C<sub>3</sub>-C<sub>4</sub> evolution across multiple plant lineages (Schlüter & Weber 2016, Schulze et al. 2013). In the Brassiceae tribe, the *GLDP2* copy was lost (Schlüter et al. 2017). Additionally, *GLDP1* was reported to be differentially expressed between C<sub>3</sub> and C<sub>3</sub>-C<sub>4</sub> intermediate *Moricandia* species (Hylton et al. 1988). In *A. thaliana*, GUS activity was restricted to the BSC by truncating the *GLDP1* promoter in the position of the M-Box, a promoter element approx. 800 bp upstream of the *AtGLDP1* gene start site. It was hypothesized that the M-Box confers MC expression, whereas the V-Box, a second promoter element shortly before the At*GLDP1* start site, confers BSC expression (Adwy et al. 2015). Promoter-GUS fusions showed that the *GLDP1* promoter of the C<sub>3</sub> species *M. moricandoides* conferred GUS expression to both MC and BSC, whereas the *GLDP1* promoter of the C<sub>3</sub>-C<sub>4</sub> intermediate species *M. arvensis* restricted GUS expression to the BSC (Adwy et al. 2019).

(Adwy et al. 2019) explain the establishment of the glycine shuttle in *Moricandia* by the loss of the M-Box in C<sub>3</sub>-C<sub>4</sub> intermediate *Moricandia* species. Contrary to this, we found the M-Box sequence in all our analyzed *GLDP1* promoter variants. However, we found the M-Box to be shifted over 1000 bp further upstream by the insertion of three independent TEs in the promoters in three independent evolutionary origins of C<sub>3</sub>-C<sub>4</sub> intermediate photosynthesis. This shift may have led to the M-Box being overlooked in previous studies. Based on the findings by Adwy et al. (2019) we hypothesize that not the loss of the M-Box conferred differential *GLDP1* expression but rather the upstream shift of the element by insertion of a TE. The upstream shift of the M-Box was mediated by three independent TE insertions in lines with independent evolutionary origins of C<sub>3</sub>-C<sub>4</sub> photosynthesis. This hints at a remarkable convergent evolutionary genetic mechanism in C<sub>3</sub>-C<sub>4</sub> evolution. We hypothesize that the loss of *GLDP2* paved the way for subfunctionalization of the *GLDP1* copy in the Brassiceae tribe, the only Brassicaceae tribe containing C<sub>3</sub>-C<sub>4</sub> intermediate species. This subfunctionalization was mediated by the insertion of a TE in the promoter, suppressing the M-Box element and shifting the *GLDP1* expression. It is questionable whether the TE insertion took place before or

after the preconditioning of C<sub>3</sub>-C<sub>4</sub> photosynthesis by anatomical adaptations such as higher vein density and the distinct leaf anatomy. Hypothetically, limited expression of *GLDP1* in the MC may have been deleterious without further adaptations, which could have prevented the TE retention in the promoter. In *D. erucoides* we do not find a transposon in the *GLDP1* promoter region. The spacing of the M-Box to the *GLDP1* start codon is in the range of C<sub>3</sub> plants (Fig. 6B). However, *D. erucoides* shows C<sub>3</sub>-C<sub>4</sub> intermediate phenotypes (Lundgren 2020, Schlüter et al. 2017). We hypothesize that, being an independent evolutionary origin of C<sub>3</sub>-C<sub>4</sub> intermediate photosynthesis, *D. erucoides* either shifted *GLDP1* expression to the BSC by different means or that there must be other additional regulators in the *GLDP1* promoter beyond our transposon-M-Box model. Contrasting the well-studied GDC activity and localization in *Moricandia* species, there is no data on the *D. erucoides* GDC biochemistry and genetics. Therefore, we cannot rule out that the glycine shuttle in *D. erucoides* is mediated by a different GDC regulation compared to the other C<sub>3</sub>-C<sub>4</sub> intermediate species, such as the differential activity of the GDC T-, L- or H- proteins.

Based on a whole-genome view of TE density and gene-TE associations, our study revealed a particular scenario in which independent TE insertions contributed to the convergent evolution of a plant trait. Our study highlights the importance of TEs in the evolution of plant traits like C<sub>3</sub>-C<sub>4</sub> intermediate photosynthesis. Differential *GLDP* expression is one of the most prominent drivers establishing the glycine shuttle. Based on *in silico* data, the proposed genetic mechanism of differential *GLDP1* regulation by a TE-mediated upstream shift of the M-Box must be verified in experimental work. Looking at the genetic mechanisms of gene regulation in C<sub>3</sub>-C<sub>4</sub> intermediate species will pave the way for a better understanding of the C<sub>4</sub> trait and facilitate genetic engineering efforts.

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## Author contributions

A.P.M.W., B.S. and U.S. designed and coordinated the project, S.T. designed and performed all analyses, A.K.D., R.N.F.M.G. and B.S. advised on statistical testing, A.K.D. performed gene annotations using *Helixer*, all authors contributed to writing and accepted the manuscript.

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