Key changes in gene expression identified for different stages of C₄ evolution in *Alloteropsis semialata*

Running title: Evolution of C4 transcriptomes in Alloteropsis

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Comparative transcriptomics in a phylogenetic context show that the initial emergence of C₄ photosynthesis in *Alloteropsis semialata* coincides with few changes in gene expression within mature leaves, with secondary adaptation occurring in geographically isolated populations.

Abstract

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 C_4 photosynthesis is a complex trait that boosts productivity in tropical conditions. Compared to C_3 species, the C_4 state seems to require numerous novelties, but species comparisons can be confounded by long divergence times. Here, we exploit the photosynthetic diversity that exists within a single species, the grass *Alloteropsis semialata*, to detect changes in gene expression associated with different photosynthetic phenotypes. Phylogenetically-informed comparative transcriptomics show that intermediates with a weak C_4 cycle are separated from the C_3 phenotype by increases in the expression of 58 genes (0.22% of genes expressed in the leaves), including those encoding just three core C_4

55 enzymes: ASP-AT, PCK, and PEPC. The subsequent transition to full C₄ physiology was accompanied by increases in another 15 genes (0.06%), including only the core C₄ enzyme PPDK. These changes likely created a rudimentary C₄ physiology, and isolated populations subsequently improved this emerging C₄ physiology, resulting in a patchwork of expression for some C₄-accessory genes. Our work shows how C₄ assembly in *A. semialata* happened in incremental steps, each requiring few

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alterations over the previous one. These create short bridges across adaptive landscapes that likely facilitated the recurrent origins of C₄ photosynthesis through a gradual process of evolution.

Keywords: adaptation, C4 photosynthesis, complex trait, intermediates, phylogenetics, transcriptomics

65 Introduction

The origins of traits composed of multiple anatomical and/or biochemical components have always intrigued evolutionary biologists (Darwin, 1859; Meléndez-Hevia *et al.*, 1996; Lenski *et al.*, 2003). If such traits gain their function only through the co-ordinated action of multiple components, their evolution via natural selection must cross a valley in the adaptive landscape. Despite this obstacle,

- complex traits have evolved repeatedly in diverse groups of organisms. This apparent paradox is solved for most traits by the existence of intermediate stages, which act as evolutionary enablers, creating bridges over the valleys of the adaptive landscape (Jacob, 1977; Dawkins, 1986; Weinreich *et al.*, 2006; Blount *et al.*, 2012; Vopalensky *et al.*, 2012; Werner *et al.*, 2014). The accessibility of new traits likely depends on the length and complexity of such bridges, which are generally unknown.
- 75 Quantifying the evolutionary gap between phenotypic states is therefore crucial to contextualise the likelihood of a novel trait evolving.

An excellent system to study the evolutionary trajectories of an adaptive trait is C₄ photosynthesis. This metabolic pathway increases CO₂ concentration at the active site of assimilation via the Calvin-Benson cycle (Hatch, 1987; Sage, 2004; Christin & Osborne, 2014). This avoids the energetically costly process of photorespiration, effectively increasing photosynthetic efficiency in warm and arid conditions (Sage *et al.*, 2012, 2018). This CO₂-concentrating mechanism relies on a set of specific leaf anatomical properties and the co-ordinated action of up to ten enzymes carrying the C₄ reactions (hereafter 'core C₄ enzymes') and numerous associated proteins (Table S1; Hatch, 1987; Bräutigam *et al.*, 2011; Sage *et al.*, 2012; Külahoglu *et al.*, 2014; Lundgren *et al.*, 2014; Yin and Struik 2017). Despite its apparent complexity, C₄ photosynthesis is a textbook example of convergent evolution, having independently evolved more than 60 times within flowering plants (Sage *et al.*, 2011). The origins of C₄ photosynthesis were likely facilitated by the presence of anatomical enablers

- in some groups (Christin *et al.*, 2013b; Sage *et al.*, 2013), but the processes leading to a functioning C₄ biochemical pathway within these anatomical structures are less well understood. All C₄ enzymes
- 90 studied so far exist in C₃ plants, but are involved in different pathways (Aubry *et al.*, 2011). There is a bias in the recruitment of genes into the C₄ system, with genes ancestrally abundant in the leaves of C₃ plants preferentially co-opted for C₄ (Christin *et al.*, 2013a; John *et al.* 2014; Emms *et al.*, 2016; Moreno-Villena *et al.*, 2018). Changes to their expression patterns and/or kinetic properties of the encoded enzyme then followed (Bläsing *et al.*, 2000; Hibberd & Covshoff, 2010; Huang *et al.*, 2017;
- 95 Moreno-Villena *et al.*, 2018), with cell-specific expression realized in some cases through the recruitment of pre-existing regulatory mechanisms (Brown *et al.*, 2011; Kajala *et al.*, 2012; Cao *et al.*,

2016; Reyna-Llorens & Hibberd, 2017; Borba et al., 2018; Reyna-Llorens et al., 2018).

The evolutionary transition between C_3 and C_4 phenotypes involves intermediate stages that only have some of the anatomical and biochemical modifications typical of C_4 plants (Monson *et al.*,

- 100 1989; Sage *et al.*, 2012, 2018). In particular, some C₃+C₄ plants perform a weak C₄ cycle that is responsible for only part of their carbon assimilation (these correspond to 'type II C₃-C₄ intermediates'; Ku *et al.*, 1983; Monson *et al.*, 1986; Schlüter & Weber, 2016). This weak C₄ cycle might have emerged through the upregulation of C₄-related enzymes to balance nitrogen among cellular compartments in the multiple lineages of plants that use a photorespiratory pump (Sage *et al.*, 2011,
- 105 2012; Mallmann *et al.*, 2014; Bräutigam & Gowik, 2016). Metabolic models suggest that any increase in flux of CO_2 fixed through the C_4 cycle in intermediate plants directly translates into biomass gain, leading to gradual increases in C_4 gene expression (Heckmann *et al.*, 2013; Mallmann *et al.*, 2014). The current model of C_4 evolution therefore assumes gradual, yet abundant changes in plant transcriptomes and genomes during the transition from C_3 ancestors to physiologically C_4 descendants. Indeed,
- 110 comparisons of C₃ and C₄ species have typically identified thousands of differentially expressed genes encoding C₄ enzymes, regulators, and accessory metabolite transporters (Bräutigam *et al.*, 2011, 2014; Gowik *et al.*, 2011; Külahoglu *et al.*, 2014; Li *et al.*, 2015; Lauterbach *et al.*, 2017). These large numbers might partially result from the comparison of species typically separated by millions of years of divergence (Christin *et al.*, 2011), which leaves ample time for the accumulation of secondary
- changes linked to the C₄ trait beyond the minimal requirements, as well as variation in other unrelated traits (Heyduk *et al.* In press). Even within a single species where photosynthetic transitions can be induced, the number of differentially expressed genes identified in transcriptome comparisons can be extremely high (Chen *et al.*, 2014). Previous efforts have however typically targeted very few individuals per C₄ lineage, such that the initial bout of co-option that generated a C₄ cycle cannot be distinguished from subsequent adaptation via natural selection and diversification caused by genetic drift (Christin & Osborne, 2014; Reeves *et al.*, 2018; Heyduk *et al.* In press).

In this study, the transcriptomes of mature leaves are compared among plant populations using a phylogenetic approach. The work aims to quantify the phenotypic differences in gene expression between the C_3 phenotype and plants using a weak C_4 cycle (C_3+C_4 state), independently from those responsible for the transition to the full C_4 type, and finally from those involved in the adaptation of an existing C_4 phenotype. The time elapsed between transitions, and therefore the number of changes unrelated to C_4 emergence, is reduced by focusing on a single species containing a diversity of photosynthetic types, the grass *Alloteropsis semialata*. Congeners of *A. semialata* are C_4 , but previous comparative transcriptomics and leaf anatomy have shown that C_4 biochemistry emerged multiple

- times in the genus, from a common ancestor with some C₄-like characters (Fig. 1; Dunning *et al.*, 2017). Capitalizing on the physiological diversity existing within *A. semialata*, leaf transcriptomes from multiple individuals originating from diverse populations of each photosynthetic type in this species are analysed, together with closely related C₃ and C₄ species, to detect the changes in gene expression linked to (i) the phenotypic difference between C₃ plants and C₃+C₄ intermediates, (ii) the
- 135 shift to fixing carbon exclusively via the C_4 pathway in solely C_4 plants, and (iii) the adaptation of the C_4 cycle after its evolution in geographically isolated C_4 populations. This deconstruction of the genetic origins of a complex biochemical pathway sheds new light on the number of genetic changes needed to move to another part of the adaptive landscape during different stages of a stepwise physiological transition.

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Material and Methods

Species sampling and growth conditions

Three biological replicates from ten separate populations/species were used for differential gene expression analyses. Seven of these were geographically distinct *Alloteropsis semialata* populations
including: two C₃ populations from South Africa (RSA6) and Zimbabwe (ZIM1502) that represent extremes of the C₃ geographic range (Fig. 1B; Lundgren *et al.*, 2015), two geographically distant C₃+C₄ populations from Tanzania (TAN1602) and Zambia (ZAM1503) that are hypothesised to operate a weak C₄ cycle (Lundgren *et al.*, 2016), and three C₄ populations from Cameroon (CMR1601), Tanzania (TAN4) and the Philippines (PHI1601) that sample the two C₄ genetic subgroups (Olofsson *et al.*, 2016; Fig. S1). The C₄ populations of *A. semialata* have decreased CO₂-compensation points,

- increased carboxylation efficiencies, and shifts in carbon isotopes compared with the C₃ populations that confirm their photosynthetic type (Lundgren *et al.*, 2016). The C₄ leaves are characterized by increased vein density, PEPC protein abundance, and transcript abundance of genes encoding some C₄ enzymes compared with the C₃ types (Lundgren *et al.*, 2016, 2019; Dunning *et al.*, 2017). The C₃+C₄
- A. semialata also show elevated leaf levels of PEPC protein and genes for some C₄ enzymes and increased concentration of chloroplasts in bundle sheaths in comparison with the C₃ populations, but no increase in vein density (Lundgren *et al.*, 2016; Dunning *et al.*, 2017). However, while slightly shifted compared to their C₃ conspecifics, their carbon isotope ratios are not in the C₄ range, which is common in plants performing a weak C₄ cycle, responsible for only part of their CO₂ uptake (i.e. 'type II
- intermediates'; Monson *et al.*, 1988; von Caemmerer, 1992; Sage *et al.*, 2012; Lundgren *et al.*, 2016).
 This results in a reduced CO₂-compensation point and oxygen inhibition (Lundgren *et al.*, 2016), as

observed in other species acquiring part of their carbon via a weak C₄ cycle (Ku et al., 1991). In addition to the seven A. semialata populations, we included one population of each of the C₄ congeners A. angusta (AANG1 from Uganda) and A. cimicina (from Madagascar) to enable comparison of

convergent C₄-related changes in gene expression (Fig. S1). Finally, an *Entolasia marginata* population 165 from Australia was included as a C₃ outgroup. Three distinct genotypes for eight of the ten populations described above were retrieved from a recent dataset (Dunning *et al.* In press) or sequenced here. For the two other populations, sufficient biological replicates were not available. For A. angusta, we sequenced three clones of a single wild collected plant that were established more than one year before the study, while for *E. marginata* we sequenced two different genotypes and a clone of these 170 genotypes, similarly established before the study (See Table S2 for detailed sample collection information).

To evaluate the diversity of gene expression across the diversity of photosynthetic types and the genetic diversity within each photosynthetic type, we supplemented the above data with a single biological replicate from a further 15 geographic distinct populations (12 from previously published 175 data; Dunning *et al.*, 2017, In press; Fig. 1A). The three newly sequenced individuals are two $C_4 A$. semialata from Sri Lanka (SRI1702, lat: 6.81 long: 80.92) and Zambia (ZAM1726, lat: -14.21 long: 28.60), and a C₃ individual from Zimbabwe (ZIM1503, lat: -18.78 long: 32.74). In total, we had 45 RNA-Seq libraries from 25 populations/species, with three biological replicates sampled from 10 populations and a single biological replicate sampled from the remaining 15 populations (Fig. 1A). 180

All plants were collected from the field as seeds or live cuttings, and subsequently grown under controlled conditions at the University of Sheffield as previously described (Dunning et al., 2017). In brief, plants were potted in John Innes No. 2 compost (John Innes Manufacturers Association, Reading, England) and maintained under wet, nutrient-rich conditions in controlled environment chambers (Conviron BDR16; Manitoba, Canada) set to 60% relative humidity, 500 μ mol m⁻² s⁻¹ light intensity, 185

14h photoperiod, and day/night temperatures of 25/20°C. After a minimum of 30 days in these growth conditions, young fully expanded leaves were sampled for transcriptome analyses.

RNA extraction, sequencing, and transcriptome assembly

RNA extraction, library preparation and sequencing were performed as previously described (Dunning 190 et al., 2017). In brief, total RNA was extracted from the distal half of fully expanded fresh leaves, sampled in the middle of the light period, using the RNeasy Plant Mini Kit (Qiagen, Hilden, Germany) with an on-column DNA digestion step (RNase-Free Dnase Set; Qiagen, Hilden, Germany). Total RNA was used to generate 34 indexed RNA-seq libraries using the TruSeq RNA Library Preparation Kit v2

(Illumina, San Diego, CA). Each library was subsequently sequenced on 1/24 of a single Illumina HiSeq 2500 flow-cell (with other samples from the same or unrelated projects), which ran for 108 cycles in rapid mode at the Sheffield Diagnostic Genetics Service.

The raw RNA-Seq data were cleaned using the Agalma pipeline v.0.5.0 to remove low quality reads (Q<30), and sequences corresponding to ribosomal RNA or containing adaptor contamination

- 200 (Dunn *et al.*, 2013). *De novo* transcriptomes were assembled using Trinity (version trinityrnaseq_r20140413p1; Grabherr *et al.*, 2011). All raw data and transcriptome assemblies have been submitted to the NCBI repository (Bioproject PRJNA401220). Coding sequences (CDS) longer than 500 bp were predicted for each population using OrfPredictor (Min *et al.*, 2005), which uses homology to a user supplied reference protein database or *ab initio* predictions if no suitable match is
- 205 found. The protein database used comprised the complete coding sequences of eight model species: *Arabidopsis thaliana, Brachypodium distachyon, Glycine max, Oryza sativa, Populus trichocarpa, Setaria italica, Sorghum bicolor* and *Zea mays.*

Phylogenetic reconstruction using core-orthologs

- 210 Single-copy orthologs were extracted from the newly and previously published transcriptome assemblies (Dunning *et al.*, 2017) to infer phylogenetic relationships among individuals. Homologous sequences to 581 single-copy plant core-orthologs previously determined in the Inparanoid ortholog database (Sonnhammer & Ostlund, 2014) were identified. A Hidden Markov Model based search tool (HaMSTR v.13.2.3; Ebersberger *et al.*, 2009) was used to screen the CDS of the transcriptomes.
- 215 Sequences of the single copy plant core-orthologs were subsequently aligned using a previously described stringent alignment and filtering pipeline (Dunning *et al.*, 2017). In brief, the CDS were translation aligned and filtered using T-COFFEE v. 11.00.8cbe486 (Notredame *et al.*, 2000) before trimming with gblocks v.0.91 (Castresana, 2000). Sequences shorter than 100 bp after trimming, and ortholog alignments with a mean nucleotide identity <95% were discarded, retaining 504 markers. A</p>
- 220 maximum likelihood tree was inferred using IQ-TREE v.1.6.3 (Nguyen *et al.*, 2014), which determined the most appropriate nucleotide substitution model prior to inferring a phylogeny with 1,000 ultrafast bootstrap replicates.

Differential expression analyses

For differential expression analysis, we used the 45,144 cDNA sequences from the *A. semialata* reference genome (Dunning *et al.*, In press; accession number QPGU00000000) as a reference.
 Cleaned reads were mapped to the reference using Bowtie2 v.2.3.4.1 (Langmead & Salzberg, 2012)

recording all alignments. Counts for each transcript were then calculated using eXpress v.1.5.1 (Roberts & Pachter, 2013) with default parameters, and are reported in reads per kilobase of transcript

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per million mapped reads (rpkm). A multivariate analysis was used to assess similarities and differences in overall transcriptome expression profiles between samples. Clustering of expression profiles based on the biological coefficient of variation (BCV) were identified with multidimensional-scaling (MDS) in edgeR v3.4.2 (Robinson *et al.*, 2010).

- Differential expression analysis in edgeR was restricted to the ten populations with three 235 biological replicates. For each pair of populations, differentially expressed genes were identified as those with an associated false discovery rate (FDR) below 0.05. The overlap between pairwise comparisons was used to identify changes associated with specific branches of the phylogenetic tree inferred from core orthologs. Changes were assigned to a branch if significant results were detected for all pairwise tests involving one member of the descending clade and one population outside the clade,
- and the direction of expression change was consistent. This summary of pairwise tests was done separately for each C₃+C₄/C₄ clade (*A. cimicina*, *A. angusta*, and *A. semialata*) with all C₃ populations so that convergent gene expression shifts could be detected. Overall, by grouping the differential expression results based on the phylogenetic clades, we are able to identify changes in gene expression that coincide with specific physiological transitions, as well as those that precede or follow these
 transitions.

Results

Transcriptome sequencing

Over 190 million 108-bp paired-end reads were used in this study, including more than 167 million for
the ten populations sampled in triplicate (Table S3). For these 30 samples used in differential
expression analyses, the data comprised 36.13 Gb, with a mean of 1.20 Gb per library (SD=0.54 Gb;
Table S3). Over 95% of reads were retained after cleaning, and a *de novo* transcriptome was assembled
for each of the populations using all available reads.

255 Phylogenetic relationships based on concatenated ortholog alignments

A phylogenetic tree was inferred from a concatenated alignment of 504 'core-orthologs' extracted from the predicted coding sequences from 25 transcriptome assemblies (12 assembled here), for a total of 573,762 bp after cleaning. Each population was represented by at least 126,048 bp (mean=468,507 bp; SD 94,782 bp). The concatenated alignment had 21.1% gaps and 6.3% of sites were parsimony

- informative. The phylogeny was inferred using the GTR+F+R4 substitution model, which was the best fit model according to the BIC. The phylogenetic relationships were congruent with previous genome-wide nuclear trees (Olofsson *et al.*, 2016; Dunning *et al.* In press), and confirmed that all the sampled C₄ populations of *A. semialata* form a monophyletic group, which is sister to the C₃+C₄ populations (Fig. 1). These two are in turn sister to the C₃ populations, so that previously inferred nuclear clades I
 (C₃), II (C₃+C₄), III and IV (both C₄) are retrieved, with the polyploid populations (RSA3 and RSA4)
- branching in between and the Cameroonian population at their base (Olofsson *et al.*, 2016; Fig. 1). *A. angusta* and *A. cimicina* branched successively outside of *A. semialata* (Fig. 1), again mirroring previous results (Lundgren *et al.*, 2015; Olofsson *et al.*, 2016; Dunning *et al.* In press).

270 Transcriptome-wide patterns

A mean of 57.4% (SD=12.05%) of cleaned reads from the 45 RNA-Seq libraries mapped back to the 45,144 cDNA sequences extracted from the reference *A. semialata* genome (only *A. semialata* samples n=34, mean=64.1%, SD=4.3%). In total, 59.8% (n=26,975) of gene sequences had expression levels of >1 read per million of mapped reads in at least three samples and were retained for differential

- expression analysis. Based on their expression profiles, samples group strongly by species (Fig. 2A).
 When focusing on *A. semialata*, the main phylogenetic groups are recovered, which match the photosynthetic types (Fig. 1 and 2B). There is no apparent effect of the source study, with previous and new transcriptomes of the same species grouping together (Fig. 2). Differential expression analysis was performed for each pair of the ten populations that had three biological replicates. The 45 pairwise tests
 performed returned an average of 4,880 (SD=2,125) significantly (FDR<0.05) differentially expressed genes (Fig. 3; Table S4). The number of differentially expressed genes is highest between the most
- distantly-related populations and lowest among close relatives (Fig. 3). Complete expression results are available in Tables S4 and S5.
- 285 Differences between the C_3 and C_3+C_4 states of A. semialata

As expected, the long divergence time between the C₃ outgroup (*Entolasia marginata*) and *A*. *semialata* results in a large number of significant expression changes (branch A in Fig. 4). A total of 825 genes are downregulated along this branch (3.1% of those expressed in leaves), including two genes encoding phosphoenolpyruvate carboxylase (PEPC; *ppc-1P2* and *ppc-2P1*;

ASEM_AUS1_43423 and ASEM_AUS1_37421; Table S6), which drop to barely detectable levels in all *A. semialata* accessions, and are therefore unlikely to be linked to photosynthetic diversification. A total of 1,500 genes (5.6%) are upregulated in *A. semialata* compared to the C₃ outgroup (branch A in

Fig. 4; Table S6). This includes genes encoding the C₄-related enzymes malate dehydrogenase (NAD-MDH; *nadmdh-2P4*; ASEM_AUS1_14800), adenosine monophosphate kinase (AK; ak-3P3;

- ASEM_AUS1_08191 and ASEM_AUS1_08195), glyceraldehyde 3-phosphate dehydrogenase
 (GAPDH; gapdh-1P2; ASEM_AUS1_06811) and phosphoenolpyruvate carboxylase kinase (PEPC-K; pepck-1P3 and pepck-3P6; ASEM_AUS1_38337 and ASEM_AUS1_12272), although their expression
 levels remain fairly low in all A. semialata regardless of photosynthetic type (mean=42 rpkm; SD=37; Table S5). One gene encoding an enzyme linked to the photorespiratory pathway is also upregulated
- (*hpr-2P3*; ASEM_AUS1_28984), although levels again remain fairly low within *A. semialata* (mean=19 rpkm; SD=13; Table S5). The rest of the numerous genes varying in expression between the whole of *A. semialata* and the outgroup do not have known links to the C₄ pathway. A total of 60 genes (0.22%) are differentially expressed along the branch leading to the C₃ populations of *A. semialata* (branch B in Fig. 4). None of these 60 genes encodes a protein known to function as part of the C₄

305 pathway (Table S6).

Within *A. semialata*, a C₄ cycle, weak or strong, characterizes the monophyletic group of C₃+C₄ and C₄ populations, but not its C₃ sister group. Along the branch leading to C₃+C₄ and C₄ accessions we detect 67 significantly differentially expressed genes (branch E in Fig. 4; Table 1). Of those, 58 (0.22% of all expressed genes) are consistently upregulated in the C₃+C₄ and C₄ populations compared to the C₃ samples, including three genes that encode key C₄ enzymes: aspartate aminotransferase (ASP-AT; *aspat-3P4;* ASEM_AUS1_08268), phosphoenolpyruvate carboxykinase (PCK; *pck-1P1;* ASEM_C4_17510), and PEPC (*ppc-1P3;* ASEM_C4_19029; Table S6). These three genes reach very high levels in the leaves of all C₃+C₄ and C₄ individuals (mean=1,766 rpkm; SD=585; Table S5; Fig. 5), including the C₄ congener *A. angusta* (mean=5,002 rpkm; SD=2,607; Table S5). The other genes

- 315 whose expression changes significantly along the same branch mostly remain at low to moderate levels in all *A. semialata*, but a number of them are also significant in *A. angusta*, and for two of them in *A. cimicina* (Tables 1 and S6). The significant genes include one for Nudix hydrolase, which was previously identified in a comparison of rice and C₄ grasses (Ding *et al.*, 2016). The remaining genes have however not been related to C₄ photosynthesis in previous screens of grasses (Ding *et al.*, 2016;
- Huang *et al.*, 2017). A gene for a callose synthase is downregulated in the C₃+C₄/C₄ group as well as *A*. *angusta* (Table 1), which might be linked to plasmodesmatal widening to facilitate intercellular fluxes, as suggested for other genes linked to callose synthesis (Bräutigam *et al.*, 2011; Huang & Brutnell, 2016). Some of the other differentially expressed genes encode proteins that have been previously suggested as being involved in metabolic/structural differences between photosynthetic types (e.g. acyl
- transferase, pyruvate dehydrogenase; Huang & Brutnell, 2016) or that might be linked to

plasmodesmata (e.g. phosphatidylglycerol/phosphatidylinositol transfer protein), although the functional links with photosynthetic diversification remain to be tested.

Changes during the transition from C_3+C_4 to C_4 in A. semialata

- Within A. semialata, a strong C₄ cycle characterizes a monophyletic group of populations (Fig. 1A), but only 16 genes (0.06% of all expressed genes) were significantly differentially expressed along the branch separating this group from the other populations (branch I in Fig. 4). Of these, 15 were consistently upregulated in the C₄ populations, including one gene encoding the core C₄ enzyme pyruvate orthophosphate dikinase (PPDK; *ppdk-1P2;* ASEM_AUS1_39556), which reaches very high
- 335 levels in all C₄ populations (mean=4,479 rpkm; SD=2,293; Tables 1 and S6; Fig. 5), including the congeners *A. cimicina* (mean=1,766 rpkm; SD=585; Table S5) and *A. angusta* (mean=1,367 rpkm; SD=1,100; Table S5). The other genes upregulated in the C₄ accessions, which include transcription factors and some transporters, reach moderate levels in the C₄ accessions, although some are also significantly upregulated in *A. angusta* (Table 1). Significant changes in the abundance of the genes for
- 340 the phosphatidylglycerol/phosphatidylinositol transfer protein might be linked to modifications of plasmodesmata to facilitate metabolite exchanges (Grison *et al.*, 2015), while aquaporins might be involved in membrane diffusion of CO₂ (Kaldenhoff *et al.*, 2014). However, whether these genes played a direct role in the photosynthetic diversification of *A. semialata* remains speculative.

345 Adaptation of C₄ photosynthesis in independent lineages

- The three C₄ populations included in the differential expression analyses come from geographically distant locations and diverged more than half a million years ago (Lundgren *et al.*, 2015; Olofsson *et al.*, 2016), explaining the large number of differentially expressed genes among them (Fig. 3). Interestingly, this includes enzymes linked to the C₄ cycle with genes encoding PEPC (*ppc-1P3*;
- ASEM_AUS1_12633), NAD-MDH (*nadmdh-1P8;* ASEM_AUS1_25602), PEPC-K (*pepck-1P3;* ASEM_C4_38337), NADP-MDH (nadpmdh-3P4; ASEM_AUS1_33376), and a sodium bile acid symporter (SBAS; *sbas-4P4;* ASEM_AUS1_12098) all upregulated in the C4 plants from the Philippines (PHI1601; Table S6). A comparison of expression levels in the other transcriptomes (including the 15 populations not used for the differential expression) indicates that the gene *sbas-4P4*
- 355 has qualitatively higher expression in all C₄ individuals from clade IV of *A. semialata* (mean=898 rpkm; SD=483), but not in the other C₄ individuals (mean=27 rpkm; SD=19) or the other *A. semialata* populations as a whole (mean=20 rpkm; SD=13; Table S5; Fig. 5). This gene is orthologous to a group of *Arabidopsis* paralogs including BASS6 (At4g22840), which has the ability to transport glycolate,

and appears to be involved in a process decreasing photorespiration (South et al., 2017). The

- 360 Arabidopsis paralog previously related to C₄ photosynthesis transports pyruvate (BASS2; Furumoto *et al.*, 2011), but its precise function might differ between the *Alloteropsis* and *Arabidopsis* orthologs. In addition, a gene encoding the photorespiratory enzyme peroxisomal (S)-2-hydroxy-acid oxidase (GLO; *glo*-1P1; ASEM_AUS1_30871) is downregulated in only one of the three C₄ populations (CMR1601; Table S6).
- There is quite large variation in the expression of individual genes encoding some other C₄ enzymes, with some more abundant in the C₄ than C₃+C₄ *A. semialata* populations on average, yet relatively low in other C₄ individuals. These genes include alanine aminotransferase (ALA-AT; alaat-1P5, ASEM_AUS1_25403; C₄ mean=1,105 rpkm; SD=812; C₃+C₄ mean=134 rpkm; SD=59; significantly differentially expressed in 13 of the 15 required pair-wise tests), which has low expression
- in C₄ individuals from Tanzania (TAN4-08; rpkm=135) and Cameroon (CMR1601-07; rpkm=154).
 Similarly, one of the genes encoding the NADP-malic enzyme (*nadpme-1P4*; NADP-ME, ASEM_AUS1_06611; significantly differentially expressed in 7 of the 15 required pair-wise tests) is on average more abundant in the C₄ and C₃+C₄ (mean=300 rpkm; SD=235) than C₃ (mean=75 rpkm; SD=32) *A. semialata* populations, but low within some C₄ individuals (e.g. TAN4-01 rpkm=82; TAN4-01
- 08 rpkm=54; ZAM1503-08 rpkm=50; Fig. 5). This gene is also significantly upregulated in *A. cimicina* and *A. angusta* (Table S5). One of the genes for PEPC kinase (*pepck1P3*) reaches high levels in several C4 accessions of *A. semialata* (Table S5). Similarly, some genes for the small unit of Rubisco reach very low levels in some C4 accessions. For instance, the gene AUS1_20231 is at low levels in most C4 *A. semialata*, yet remains very high in others while the paralog AUS1_26631 reaches extremely low
 levels, specifically in the Asian group of C4 *A. semialata* (Table S5). A third paralog (AUS1_26630) remains high in all accessions, so that the total abundance of genes for Rubisco is not markedly decreased, which is congruent with the high Rubisco protein abundance in the leaf of the C4 *A. semialata* (Ueno & Sentoku, 2006).

The number of genes significantly differentially expressed in the C₄ *A. cimicina* and *A. angusta*lineages is much higher, since only one population represents each of these species (Fig. S3). As
previously reported (Dunning *et al.*, 2017), a high number of genes encoding core C₄ enzymes,
regulatory proteins and transporters are upregulated in *A. cimicina* (Table S7), and to a lesser extent in *A. angusta* (Table S8), while some photorespiration and Rubisco genes are downregulated in both
species. Besides the differentially expressed genes, a number of C₄-related genes are abundant in all
samples independent of their photosynthetic type. This is especially the case of genes encoding βcarbonic anhydrase (βca-2P3; ASEM_AUS1_16750; mean=1,682 rpkm, SD=1,027, min=290) and

malate dehydrogenases: *nadpmdh-1P1* (ASEM_AUS1_23802; mean=443 rpkm, SD=501, min=117), *nadpmdh-3P4* (ASEM_AUS1_33376; mean=447 rpkm, SD=184, min=166), and *nadmdh-3P5* (ASEM_AUS1_22160; mean=157 rpkm, SD=69, min=41). Transcripts for these genes were also

abundant in the leaves of distantly related C₃ grasses, and their upregulation very likely predates the

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Discussion

Sampling the natural diversity to limit false positives

diversification of the group (Moreno-Villena et al., 2018).

- RNA-Seq is routinely used to identify genes differentially expressed between individuals with distinct phenotypes, leading to lists of candidate genes underpinning these differences (e.g. Shen *et al.*, 2014; Dunning *et al.*, 2016; Fracasso *et al.*, 2016). When comparing distinct species, the risk of false positives is very high, as all changes in gene expression unrelated to the studied phenotypic transitions are detected. Here, 77.1% of genes expressed in the leaves are significantly differentially expressed in at least one pairwise comparison between our ten populations (49.8% within *A. semialata*), which all belong to a relatively small group of closely related grasses. A powerful strategy to reduce false positives is to consider multiple independent origins of the trait of interest, and retain only those genes differentially expressed in all lineages (Ding *et al.*, 2016; Rao *et al.*, 2016). Such a filter would however exclude non-convergent changes in gene expression.
- 410 The alternative approach adopted here was to carry out multi-individual comparisons to infer changes along specific branches of the phylogenetic tree. The problem of false positives remains, as changes coinciding with the studied transitions would also be detected. However, working within a species complex decreases the number of false positives, as shorter divergence times are likely to result in fewer unrelated changes in gene expression. Because most changes cluster on terminal branches
 415 (Fig. 4), probably representing neutral changes that do not persist over evolutionary time, the inference of changes on short internal branches is less likely to be affected by drift. Indeed, a comparison of a C₃ *A. semialata* with the C₄ sister species *A. angusta* would identify over 5,000 (18% of genes expressed)
 - in the leaves) differentially expressed genes (Fig. 3). This number drops by approximately 50% when comparing individual C₃ and C₄ populations within *A. semialata*, but still includes all changes that
 occurred before, during, and after the C₃ to C₄ transition. After incorporating multiple populations of each type, only 67 genes (0.25% of genes expressed in the leaves) are identified that differ in expression between the C₃ and C₃+C₄ phenotypes, and 16 (0.06% of genes expressed in the leaves) between the C₃+C₄ and C₄ states. Changes in some of these genes might not be directly linked to the

diversification of photosynthetic types, but several were convergently modified in A. angusta and/or A.

425 *cimicina* (Table 1). These genes represent the best candidates for a role in the emergence and subsequent strengthening of a C₄ cycle in the group.

Emergence and reinforcement of the C_4 cycle in Alloteropsis semialata

The phylogenetic relationships and genus-wide comparisons of transcriptomes and leaf anatomical 430 traits indicate that the last common ancestor of all A. semialata might have possessed a weak C₄ cycle based on the upregulation of some enzymes (Fig. 1; Dunning et al., 2017). A large number of genes are differentially expressed between all A. semialata and the C_3 outgroup, which is not surprising given the evolutionary distance of at least 15 Myr (Christin et al., 2014). However, these include relatively few genes encoding C₄ enzymes (Table S6). We conclude that the transcriptome of the C₃ A. semialata differs from that of other C₃ grasses by relatively few C₄-related genes. The C₃ group might represent a 435 reversal from a C_3+C_4 state to a phenotype with expression levels similar to the C_3 outgroup. In such a scenario, C₄-related changes that happened in the last common ancestor of A. semialata and were reversed in the C_3 group would be assigned to the branch leading to the C_3+C_4 and C_4 groups. Because they focus on the phenotypic gaps in gene expression between the C_3 state and those using a weak or strong C₄ cycle, our transcriptome comparisons are therefore not heavily influenced by potential 440 evolutionary reversals or reticulate evolution.

In total, 67 genes are differentially expressed in the group encompassing C₃+C₄ and C₄ phenotypes, and these include only three genes encoding core C₄ enzymes that are upregulated in all C₃+C₄ and C₄ individuals (genes for ASP-AT, PCK and PEPC; Table 1; Table S5). These three enzymes form an aspartate shuttle based on the PCK decarboxylase (Fig. 6), which theoretically cannot sustain a full C₄ pathway on its own without creating an energetic imbalance among cell types (Wang *et al.*, 2014). However, it might create a weak CO₂-concentrating mechanism in C₃+C₄ plants that can function without dramatic energetic consequences due to its coexistence with a C₃ type of photosynthesis. While the functional significance of the other changes detected along the same branch is not always known, several might be linked to the control of plasmodesmata and thereby intracellular exchanges (Table 1). Other small adjustments of the cellular metabolism might remain undetected, but none of the other major C₄ enzymes or transporters are significantly upregulated during the emergence of a weak C₄ cycle (Table 1). The apparently few changes in transcription required to operate a weak

 C_4 cycle in the C_3+C_4 intermediates may be facilitated by C_4 -like anatomical properties and an

455 abundance of genes for some key enzymes in the ancestor, as observed in other C₃ grasses (Christin *et al.*, 2013a, 2013b; Emms *et al.*, 2016; Dunning *et al.*, 2017; Moreno-Villena *et al.*, 2018), and recent

evidence suggests that some anatomical traits themselves might emerge via very few genetic changes (Wang *et al.*, 2017). While it is only responsible for part of the plant's CO_2 uptake, the weak C_4 cycle of C₃+C₄ plants reduces photorespiration (Ku et al., 1991; Lundgren et al., 2016), which confers a selective advantage analogous to that of a complete C₄ cycle in tropical conditions (Sage *et al.*, 2012; Christin & Osborne, 2014; Lundgren & Christin, 2017), and allows the evolution of a stronger C₄ cycle under natural selection for faster biomass accumulation (Heckmann et al., 2013; Mallmann et al., 2014; Bräutigam & Gowik, 2016).

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The transition from a weak to a strong C₄ cycle in A. semialata changes carbon isotope signatures (the method most often used to identify photosynthetic types) from non-C₄ values to values diagnostic of C₄ plants (von Caemmerer, 1992; Lundgren et al., 2015). This shift indicates a strengthened connection between the C_3 and C_4 cycles and a decreased leakiness, so that less atmospheric CO2 is directly fixed by the Calvin-Benson cycle (Monson et al., 1988; von Caemmerer, 1992). Within A. semialata, this might have been mediated by the reduced distance between veins in

- the C₄ A. semialata (Lundgren et al., 2016, 2019; Dunning et al., 2017) and/or biochemical alterations. 470 The upregulation of relatively few genes (0.06%) coincided with the phenotypic transitions, and only one of these encoded an enzyme with a known C₄ function, namely PPDK. This enzyme is responsible for the regeneration of PEP, the substrate of PEPC (Fig. 6). An increased PPDK activity is also observed between species of Flaveria performing a weak and a strong cycle, and it has been suggested
- that this provides PEPC with PEP at higher rates, thereby increasing the efficiency of the C₄ pathway 475 (Monson & Moore, 1989; Sage et al., 2012). Based on the literature and our transcriptome data, the C₄ cycle of A. semialata relies on a minimum of seven enzymes (Fig. 6; Frean et al., 1983; Ueno & Sentoku, 2006). Genes for some of these enzymes (NAD-MDH, and AK) increased in the common ancestor of the whole group, potentially as part of an ancestral weak C₄ cycle (Fig. 1; Dunning et al.,
- 2017). Within A. semialata, further increases in transcript abundance are observed in the C₃+C₄ vs C₃ 480 or C₄ vs C₃+C₄ comparisons (Table 1) for genes encoding PEPC and three other enzymes (i.e. ASP-AT, PCK, and PPDK; Fig. 5). The expression of genes encoding CA and others NAD(P)-MDH in the C_3 ancestor of the group might have been sufficient to sustain a functioning C_4 cycle (Table S5; Moreno-Villena et al., 2018). Genes for the last of these enzymes (NADP-ME) are abundant in some

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C₄ individuals (Table S5; Fig. 5), and might be expressed only in specific conditions, as suggested previously (Frean et al., 1983).

C₄ populations of A. semialata are also characterized by a set of specific anatomical modifications and changes in the cellular localization of some enzymes (Ueno & Sentoku, 2006; Lundgren et al., 2016, 2019; Dunning et al., 2017). Gene expression changes responsible for these

- 490 modifications would not necessarily be captured by our transcriptome analyses of full mature leaves, and the evolution of the C₄ phenotype almost certainly involves more genetic changes than those detected here. While protein abundance is not a direct function of gene expression, the two are correlated (Schwanhäusser *et al.*, 2011; Csárdi *et al.*, 2015; Koussounadis *et al.*, 2015). In the case of *A. semialata*, the three C₄ enzymes with genes differentially expressed in the C₃+C₄/C₄ transcriptomes
- 495 (PEPC, ASP-AT and PCK) are also the ones with large differences in activities between the C₃ and C₄ *A. semialata* in a previous study (Ueno & Sentoku, 2006). Transcriptome comparisons offer a first assessment of the changes underlying adaptive transitions, allowing subsequent investigations of responsible regulatory elements, post-transcriptional processes, changes of the protein kinetics, and verification of gene functions via genetic manipulation (e.g. Wang *et al.*, 2017; Borba *et al.*, 2018).
- Overall, our comparative transcriptomics show that, once the required enablers are present, the transition between C₃ to C₃+C₄ with some C₄ activity, and C₃+C₄ to a rudimentary C₄ metabolism might have required fewer changes in gene expression in *A. semialata* than previously suggested based on other comparisons (Bräutigam *et al.*, 2011, 2014; Gowik *et al.*, 2011; Külahoglu *et al.*, 2014; Li *et al.*, 2015). These changes were spread between the C₃/C₃+C₄ and C₃+C₄/C₄ transitions, supporting a stepwise model of evolution (Mallmann *et al.*, 2014), where evolutionarily stable adaptive peaks can be

reached with few mutations.

Adaptation continued after the emergence of a rudimentary C₄ pathway

- The CO₂-pump generated by the C₄ cycle of *A. semialata* is less efficient than that of other C₄ species
 (Niklaus and Kelly, 2019), as illustrated by the incomplete segregation of enzymes between different cell types (Ueno & Sentoku, 2006) and slightly elevated CO₂-compensation points lying at the upper limit of those observed in C₄ species (Lundgren *et al.*, 2016). Therefore, *A. semialata* may be considered to exhibit an incipient C₄ cycle, which has not been optimised through protracted evolutionary periods, as suggested in the most recent models (Bräutigam & Gowik, 2016). The
- analyses conducted here, which compared all C₄ individuals to the C₃+C₄ or C₃ conspecifics, can detect the changes that happened in the early C₄ members of the group, before the diversification of the C₄ genotypes. However, transcriptome comparisons across C₄ individuals of *A. semialata* show evidence of additional alterations of the leaf biochemistry subsequent to the initial emergence of a C₄ cycle, with the abundance of some C₄-related enzymes varying in abundance across C₄ populations (e.g. NAD-
- 520 MDH) and photorespiratory proteins downregulated in only some of the C₄ populations (Tables S5 and S6). These changes likely represent the adaptation of the C₄ cycle after its initial emergence (Heyduk *et al.* In press; Niklaus and Kelly, 2019), previously illustrated for *A. semialata* by variation in the identity

of genes responsible for an abundance of the key C4 enzyme PEPC across C4 genotypes (Dunning et al., 2017) and leaf anatomy (Lundgren et al. 2019), and recently reported for Gynandropsis gynandra (Reeves et al., 2018).

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The C₄ pathway proposed for A. semialata, based on the upregulation of four core C₄ enzymes in addition to those present in C_3 ancestors (Fig. 6), might serve as an intermediate stage toward more complex and more efficient C_4 cycles. The congeneric $C_4 A$. *cimicina* and A. *angusta* have transcriptomes more typical of other C₄ species, with very high levels of numerous C₄-related enzymes,

- including a number of regulatory proteins and metabolite transporters (Table S5), as would be 530 predicted from other study systems, and an abundance of amino acid transitions adapting the proteins for the new catalytic context (Bräutigam et al., 2011, 2014; Gowik et al., 2011; Mallmann et al., 2014; Christin et al., 2015; Dunning et al., 2017). These two species might have undergone more adaptive changes, due to an earlier C4 origin or faster evolutionary rate. As illustrated by the additional C4-
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related genes upregulated in the C₄ plants from the Philippines, the rudimentary C₄ trait of A. semialata is likely to undergo similar secondary adaptations over evolutionary time.

Conclusions

In this study, the transcriptomes of individuals from the grass Alloteropsis semialata are analysed in a phylogenetic context to show that the changes in gene expression required for a physiological 540 innovation can be spread over time. The relatively few changes required for the initial emergence of a metabolic pathway contrasted with the numerous modifications involved in the adaptation of this new pathway. Indeed, the emergence of a weak C_4 cycle in our study system was accompanied by the upregulation of three enzymes with a known C₄ function and 55 others proteins. The evolution of a stronger C₄ cycle then involved the upregulation of one other C₄ enzyme and 14 other proteins. 545 However, adaptation of C₄ photosynthesis, illustrated here by population-specific expression of C₄specific enzymes, continues when the plants are already in a C₄ state. The evolutionary modifications

- 550 responses of the species. This creates an opportunity for natural selection to act on the standing variation, new mutations and, in some cases, laterally acquired genes, to assemble a trait of increasing complexity, allowing the colonization and gradual dominance in a larger spectrum of ecological conditions.

required to generate a rudimentary C_4 pathway can therefore be modest in species possessing C_4

enablers, but even a suboptimal C₄ pathway is important because it changes the environmental

Data deposition

555 All raw DNA sequencing data (Illumina reads) and transcriptome assemblies generated as part of this study have been deposited with NCBI under Bioproject PRJNA401220.

Supplementary Data

Supplementary data are available at *JXB* online.

Table S1: List of enzymes considered as core C₄ enzymes.
Table S2: Information for populations sampled in triplicates.
Table S3: RNA-Seq data and mapping statistics for ten populations with triplicates.
Table S4: Pairwise differential expression test results for all genes.
Table S5: Leaf abundance, annotation, and summary of significance for all genes.
565 Table S6: Summary of differentially expressed genes referred to in Fig. 1.
Table S7: Summary of differentially expressed genes referred to in Fig. S1A.
Table S8: Summary of differentially expressed genes referred to in Fig. S1B.
Figure S1: Phylogenetic patterns of changes in gene expression in (A) *Alloteropsis angusta*, and (B) *Alloteropsis cimicina*.

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Author contributions

580 LTD, JJMV, AB, CPO, and PAC designed the research; LTD, MRL, JD, PS, CA, FN, JKO, AM, IMA, CJK, LAD, FK, JT, GB, WPQ, CPO, and PAC identified and collected plant material; LTD and JJMV generated and analysed the transcriptome data, with the help of AB and PAC; LTD, JJMV, and PAC wrote the paper with the help of all co-authors.

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Table 1: List of genes differentially expressed in key comparisons within *Alloteropsis semialata* from C_3 to C_3+C_4 , and C_3+C_4 to C_4 that have SwissProt annotations. SwissProt protein description and *Arabidopsis* ortholog information is based on top-hit blast matches. Mean rpkm is derived from the seven *A. semialata* populations used for differential expression analysis (full summary of results can be found in Table S6).

815

Arabidopsis Mean rpkm Gene SwissProt protein description ortholog C3+C4 **C**₄ **C**₃ Genes upregulated in C_3+C_4 and C_4 A. semialata (branch E in Fig. 4) ASEM AUS1 17510^a Phosphoenolpyruvate carboxykinase (PCK) AT4G37870 2 1168 3017 ASEM AUS1 08268^a Aspartate aminotransferase (ASP-AT) AT5G11520 158 1843 1196 ASEM_AUS1_19029^a Phosphoenolpyruvate carboxylase (PEPC) 95 828 AT2G42600 1118 ASEM AUS1 30031^a Fruit bromelain AT1G06260 11 260 497 ASEM_AUS1_08709 Iron-sulfur cluster assembly protein 1 AT4G22220 394 473 67 ASEM AUS1 11198 Bifunctional TENA2 protein AT3G16990 10 43 80 ASEM_AUS1_19914 50S ribosomal protein L17 AT5G64650 78 1 58 ASEM_AUS1_02887^a Cysteine proteinase 1 AT2G32230 0 44 54 ASEM_AUS1_16281^a Probable carboxylesterase 15 AT5G06570 1 16 50 ASEM_AUS1_11666 Putative protease Do-like 14 AT5G27660 1 63 39 ASEM_AUS1_18766^a Nudix hydrolase 16 AT3G12600 4 24 38 ASEM_AUS1_21431^a DNA-binding protein MNB1B 94 30 AT4G35570 0 ASEM AUS1 24040^{a,b}Putative phosphatidylglycerol/phosphatidylinositol transfer protein AT3G11780 4 32 24 ASEM_AUS1_08934 Putative F-box protein 0 18 23 AT4G38870 ASEM AUS1 44075 Indole-3-acetaldehvde oxidase AT5G20960 0 28 22 Dihydrolipoyllysine-residue acetyltransferase component 1 of ASEM_AUS1_24692 AT3G52200 0 13 20 pyruvate dehydrogenase complex 0 ASEM_AUS1_38810 UDP-glycosyltransferase AT1G05680 35 17 0 19 ASEM_AUS1_24427 Putative F-box protein AT1G65770 16 ASEM AUS1 43609^a Flavin-containing monooxygenase FMO GS-OX-like 9 0 7 13 AT5G07800 ASEM AUS1 40960 Cysteine-rich receptor-like protein kinase 26 AT4G23240 1 18 13 ASEM AUS1 16960^a Valine--tRNA ligase AT1G14610 0 26 12 ASEM_AUS1_27461^b Aspartic proteinase nepenthesin-2 AT2G03200 0 2 12 ASEM_AUS1_15840 Tyrosine--tRNA ligase AT2G33840 0 4 10 ASEM AUS1 22664 Probable nucleolar protein 5-1 0 19 AT5G27120 8 ASEM_AUS1_39034 Putative protease Do-like 14 0 7 AT5G27660 11 ASEM AUS1 21913 Protein NEN1 AT5G07710 0 5 6 7 ASEM_AUS1_01903 Disease resistance protein RPM AT3G07040 0 2 Genes downregulated in C_3+C_4 and C_4 A. semialata (branch E in Fig. 4) ASEM_AUS1_21734 60S ribosomal protein L23a 0 AT3G55280 206 72 ASEM_AUS1_01414^{a,b}Acyl transferase 4 AT3G62160 150 18 17 ASEM_AUS1_31537 Pumilio homolog 23 49 12 9 AT1G72320 7 7 ASEM_AUS1_00061 40S ribosomal protein SA AT3G04770 42 ASEM AUS1 22162 Tubulin alpha-3 chain AT4G14960 32 6 3 ASEM_AUS1_22449^a Callose synthase 3 AT5G13000 30 2 1 ASEM_AUS1_04268^a 40S ribosomal protein S21 AT5G27700 20 0 0 ASEM AUS1 06562^{a,b}PTI1-like tyrosine-protein kinase 3 AT3G59350 5 1 1 Genes upregulated in C_4 A. semialata (branch I in Fig.4) ASEM_AUS1_39556^{a,b}Pyruvate, phosphate dikinase 1 (PPDK) AT4G15530 60 133 1149 ASEM_AUS1_24184^a Phosphatidylglycerol/phosphatidylinositol transfer protein AT3G11780 0 1 104 2 ASEM_AUS1_29700 Protein SRG1 AT1G17020 1 86 ASEM_AUS1_16577^a Lactoylglutathione lyase 0 0 46 AT1G11840 ASEM AUS1 06220 S-norcoclaurine synthase 1 AT1G17020 1 1 39 ASEM_AUS1_24241 DnaJ homolog subfamily A member 1 AT3G14200 1 1 33

ASEM_AUS1_44200 ^a Aquaporin TIP1-1	AT2G36830	0	0	17
ASEM_AUS1_13652 Transcription factor TGAL4	AT1G08320	0	0	7
ASEM_AUS1_00246 Nicotinamide adenine dinucleotide transporter 2	AT1G25380	0	0	2
Genes downregulated in C_4 A. semialata (branch I in Fig.4)				
ASEM_AUS1_43847 ^{a,b} Short-chain dehydrogenase TIC 32	AT4G23420			0
^a Significant change in the same direction in A. angusta; ^b Significant change in the same direction in A. cimicina				

Figure captions

820 Figure 1: Phylogenetic tree inferred from multiple nuclear markers.

- (A) This phylogeny was inferred under maximum likelihood using transcriptome-wide markers.
 Scale indicates number of nucleotide substitutions per site, and bootstrap support values are indicated near nodes. AANG = A. angusta. For *A. semialata*, population names indicate the country of origin; AUS = Australia, BUR = Burkina Faso, CMR = Cameroon, MAD = Madagascar, PHI =
- Philippines, RSA = South Africa, TAN = Tanzania, SRI = Sri Lanka, TPE = Chinese Taipei, ZAM = Zambia, ZIM = Zimbabwe. Populations sampled with biological replicates and used for differential expression analysis are indicated by the large circles and bold population names. Nuclear clades from Olofsson *et al.* (2016) are indicated. Branch colours indicate the ancestral photosynthetic types, based on the transcriptomes and leaf anatomy detailed investigations of
- Big Dunning *et al.* (2017). The hashed green at the base of *A. semialata* indicates uncertainty between C_3 and C_3+C_4 states. (B) Distribution of *A. semialata photosynthetic* types and sampling locations, with color codes as in panel A. Shadings indicate the approximate ranges of the three photosynthetic types of *A. semialata*, based on Lundgren *et al.* (2016).

Figure 2: Expression profile similarity across all samples.

Expression profiles are clustered in multidimensional scaling (MDS) plots using (A) all samples (B) only *A. semialata* samples. Species and nuclear clades from Olofsson *et al.* (2016) are delimited and population names are as in Fig. 1.

Figure 3: Number of differentially expressed genes among pairs of populations.

The heatmap shows the number of significantly differentially expressed genes detected for each pair of populations. The phylogenetic relationships among populations are indicated on the side, using an ultrametric version of the tree presented in Fig. 1.

Figure 4: Phylogenetic patterns of changes in gene expression.

The maximum-likelihood phylogeny from Fig. 1 is shown unrooted after pruning the populations not used for expression analyses. For each branch, the number of differentially expressed genes is indicated, with numbers next to arrows indicating those that are consistently up- or down-regulated as you move along the tree from the outgroup *Entolasia marginata*. Each population has three

biological replicates, and colours indicate the photosynthetic type (blue = C_3 ; green = C_3+C_4 ; red = C_4). Scale indicates number of nucleotide substitutions per site, with truncated branches highlighted

by two bars. The two greyed out C_4 congeners were excluded from these analyses, and results that involve them can be found in Fig. S3.

Figure 5: Expression levels of exemplar genes across accessions.

Expression levels in reads per kilobase of transcript per million mapped reads (rpkm) are shown for four example genes. Standard deviation for populations with biological replicates is indicated.
Colours indicate the photosynthetic types; blue = C₃; green = C₃+C₄; red = C₄.

Figure 6: Putative C4 pathway in Alloteropsis semialata

- A C₄ cycle is suggested for A. semialata based on the transcript abundance of C₄-related genes, and 860 the literature (Frean et al., 1983; Ueno & Sentoku, 2006). Pathway components are coloured per the differential expression analysis, with those in black being putatively sufficiently abundant in C₃ ancestors, parts of the pathway in green upregulated during the transition to C_3+C_4 , and parts in red upregulated during the transition from C_3+C_4 to C_4 . ALA-AT = alanine aminotransferase, ASP-AT = aspartate aminotransferase, CA = carbonic anhydrase, NADP-MDH = NADP malate 865 dehydrogenase, NAD(P)-ME = NAD(P) malic enzyme, PCK = phosphoenolpyruvatecarboxykinase, PEPC = phosphoenolpyruvate carboxylase, PEPP = phosphoenolpyruvate phosphatase, PPDK = pyruvate orthophosphate dikinase, PCR = photosynthetic carbon reduction (С 870 a 1 v i 875 n _ В e n 880 S 0 n с 885 y с 31 1 e
- 890