Glyceraldehyde-3-phosphate dehydrogenase subunits A and B are 1

- essential to maintain photosynthetic efficiency 2
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21 **Short title:** Evaluation of GAPDH subunits A & B in Arabidopsis

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ABSTRACT

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In plants, glyceraldehyde-3-phosphate dehydrogenase (GAPDH; EC 1.2.1.12) reversibly converts 1,3-bisphosphoglycerate to glyceraldehyde-3-phosphate coupled with the reduction of NADPH to NADP⁺. The GAPDH enzyme that functions in the Calvin Benson Cycle is assembled either from four glyceraldehyde-3-phosphate dehydrogenase A subunits (GAPA) proteins forming a homotetramer (A₄) or from two GAPA and two glyceraldehyde-3phosphate dehydrogenase B subunit (GAPB) proteins forming a heterotetramer (A_2B_2) . The relative importance of these two forms of GAPDH in determining the rate of photosynthesis is unknown. To address this question, we measured the photosynthetic rates of Arabidopsis (Arabidopsis thaliana) plants containing reduced amounts of the GAPDH A and B subunits individually and jointly, using T-DNA insertion lines of GAPA and GAPB and transgenic GAPA and GAPB plants with reduced levels of these proteins. Here we show that decreasing the levels of either the A or B subunits decreased the maximum efficiency of CO₂ fixation, plant growth, and final biomass. Finally, these data showed that the reduction in GAPA protein to 9% wild-type levels resulted in a 73% decrease in carbon assimilation rates. In contrast, eliminating GAPB protein resulted in a 40% reduction in assimilation rates. This work demonstrates that the GAPA homotetramer can compensate for the loss of GAPB, whereas GAPB alone cannot compensate fully for the loss of the GAPA subunit.

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Keywords: Glyceraldehyde-3-phosphate dehydrogenase; GAPDH; Photosynthesis; biomass

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INTRODUCTION

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order to feed the growing world population against the backdrop of climate change (IPCC, 2014; Pereira, 2017; IPCC, 2019; NASA, 2020). The photosynthetic capacity of a crop over the season determines the rate of growth and hence yield potential. A number of reports have now been published demonstrating that under glasshouse and field conditions improvements in photosynthesis, including the Calvin-Benson Cycle (CBC) can improve the productivity and yield of the plant (Driever et al., 2017; Kubis and Bar-Even, 2019; Simkin, 2019; Simkin et al., 2019; Burgess et al., 2022; De Souza et al., 2022; Raines et al., 2022). In the CBC, glyceraldehyde-3-phosphate dehydrogenase (GAPDH) catalyses the conversion of 1,3bisphosphoglycerate to the triose phosphate, glyceraldehyde 3-phosphate (GAP) (Cséke and Buchanan, 1986). Previous work has shown that the antisense suppression of the GAPDH gene had no effect on the rate of CO₂ assimilation until GAPDH activity had decreased to 30-40% of WT levels (Price et al., 1995; Ruuska et al., 2000). However, more recently a study showed that overexpression of GAPDH in rice (Oryza sativa) resulted in increased photosynthetic CO₂ assimilation under elevated [CO₂] conditions (Suzuki et al., 2021), raising the possibility that GAPDH could be a target for future manipulations to improve photosynthesis. The CBC GAPDH is highly regulated and in plants is comprised of two distinct subunits, the glyceraldehyde-3-phosphate dehydrogenase A subunits (GAPA) and the glyceraldehyde-3-phosphate dehydrogenase B subunits (GAPB) that function as either as a homotetramer A₄ or heterotetramer A₂B₂ (Cerff, 1979; Iadarola et al., 1983; Howard et al., 2011). The primary structures of these two subunits show considerable similarity and are produced from separate nuclear genes (GapA1, GapA2 and GapB) (Cerff, 1995). The GAPA subunits share 92.6% identity and the major difference in the primary sequence between the

In recent years there has been a focus to develop strategies to increase crop yields in

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GAPA and GAPB subunits is a C-terminal extension (CTE) on the GAPB, with substantial similarity to the C-terminus of Chloroplast protein of 12 kDa (CP12) (Baalmann et al., 1996; Pohlmeyer et al., 1996), This CTE contains cysteine residues which have been shown to confer thioredoxin-mediated redox regulatory capacity onto the GAPDH A₂B₂ complex (Baalmann et al., 1996; Scheibe et al., 1996; Sparla et al., 2002; Marri et al., 2005; Fermani et al., 2007).

In the chloroplast of vascular plants, the predominant active form of GAPDH is believed to be the A₂B₂, however, leaves of many species contain other, less abundant forms of GAPDH including the A₄ form and the 2(A₂B₂) and 4(A₂B₂) multimers involved in deactivation of the enzyme (Wolosiuk and Buchanan, 1976; Scagliarini et al., 1998; Sparla et al., 2005; Fermani et al., 2007). The homotetramer A₄ form of GAPDH has been termed 'non-regulatory' (GAPDH_N), firstly because of the absence of the CTE identified in GAPB and secondly, it fails to aggregate into larger oligomers and the A2B2 regulatory form (GAPDH_R) (Scagliarini et al., 1998). In the absence of the CTE, GAPDH_N is thought to be regulated by the formation of the CP12/GAPDH/PRK (Phosphoribulokinase) complex (Trost et al., 2006; Lopez-Calcagno et al., 2014). Although the regulation of the two tetramers of GAPDH are different, results of Scagliarini et al (1998) showed that the kinetic properties of GAPDH_N are similar to GAPDH_R. The activity data for the GAPDH A₄ and the A₂B₂ showed that both of these isoforms have similar kinetic parameters, with a V_{max} (NADPH) of 130 and 114 μmol min⁻¹ mg⁻¹ respectively and a Km (BPGA) of 2.0 and 2.3 μM respectively. Based on these data it was proposed that that the B-subunits are mostly responsible for regulation of the enzyme (Sparla et al., 2005) and the A-subunits for catalytic activity (Scagliarini et al., 1998). Over and above the multimers of the A₂B₂ complex a further level of redox regulation of GAPDH activity occurs through the formation of a high molecular weight complex which

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98 includes the CBC enzyme PRK and the small regulatory protein CP12 (Howard et al., 2008; 99 Carmo-Silva et al., 2011; Lopez-Calcagno et al., 2014; Lopez-Calcagno et al., 2017).

The A₄ homotetramer has been shown in spinach (Spinacia oleracea) chloroplast preparations to constitute 15–20% of the total GAPDH activity (Scagliarini et al., 1998). Howard et al. (2011) examined stromal extracts from dark-adapted leaves of species from Leguminosae (pea (Pisum sativum 'Onwards'), Medicago (Medicago truncatula 'Jemalong'), broad bean (Vicia faba 'The Sutton'), French bean (Phaseolus vulgaris 'Vilbel')), Solanaceae (potato (Solanum tuberosum 'Desiree'), tomato (Solanum lycopersicon 'Gardener's Delight'), and tobacco (Nicotiana tabacum 'Samson')), Amaranthaceae (spinach), and the Brassicaceae (Arabidopsis). This study revealed that the relative amounts of the A₂B₂ and the A₄ complexes vary among species. Whereas all species were found to accumulate the A₂B₂ heterotetramer, in contrast, in some plant species the A₄ tetramer was not detected (Howard et al., 2011). This raises the question of the role of the A₄ form for the activity of GAPDH in the CB cycle. To date the relative importance of the A₄ versus the A₂B₂ form of plastid GAPDH, in determining the rate of CO₂ assimilation, has not been elucidated. In this manuscript, to explore this question we have used insertion mutants in both the GapB and GapA-1 genes together with transgenic lines where the relative amounts of the GAPA and GAPB proteins have been decreased individually.

RESULTS

Identification and analysis of Arabidopsis lines with reductions in GAPDH A and B transcript and protein levels

We identified T-DNA insertion mutants for *gapa-1* (SAIL_164_D01) and *gapb* (SAIL_308_A06) from The Arabidopsis Information Resource (TAIR) database (http://www.arabidopsis.org/). All T-DNA insertion sites were confirmed using PCR analysis of genomic DNA followed by sequencing of the T-DNA/gene junctions. The positions of the

123 T-DNA inserts are presented in Fig. 1. Homozygous plants (identified by PCR) were used to 124 assess the effect of each T-DNA insertion on the expression of the GAPDH transcripts. RT-125 qPCR analysis confirmed that the transcript abundance encoding the GAPA subunit in the 126 gapa-1 mutant was reduced by approx. 45%, the remaining transcript being produced by the 127 gapa-2 gene (Fig. 2A). The transcript for the GAPB subunit in the gapb mutant was not 128 detected in the T-DNA insertion line, evidencing the gapb mutant as a true knockout (KO) 129 In order to obtain additional independent mutant lines, GAPA and GAPB 130 expression levels were downregulated using antisense constructs (Supplemental Fig. S1, Fig. 131 **2A** and **B**). Additionally, two GAPA co-suppressed lines were identified from plants 132 transformed with a GAPA over-expression construct using the Arabidopsis sequence in the 133 sense orientation (Supplemental Fig. S1C). In GapA co-suppressed transformants, we 134 identified two lines with 5% (cA1) and 11% (cA2) of total GapA (GapA-1 and GapA-2) 135 transcript levels and antisense lines with 35% (aA1) and 11% (aA2) of GapA transcript (Fig. 136 **2A**). In anti-sense *GapB* transformants, we identified three lines with 11% (aB1), 15% (aB2) 137 and 41% (aB2) levels of the GapB transcript (Fig. 2B). Western blot analysis of these mutant 138 lines was used to determine changes in GAPDH protein levels, which showed a reduction in 139 bands at 37.6 kDa representing GAPA and 47.7 kDa (Fig. 2A and B). In the GAPA1 140 insertion line and the GAPA1 antisense lines the level of the GAPA protein was reduced to 141 51-43% of the CN plants and in the GAPA co-supressed lines only 9% of the GAPA protein 142 was detected (Table 1). In the GAPB insertion line no band was detected indicating the 143 absence of the B subunit in this mutant (Fig.2A). The level of protein in the GAPB antisense 144 lines was between 15 and 40% of the CN plants (Fig.2B and Table 1).

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Chlorophyll fluorescence imaging reveals that PSII efficiency is maintained in plants showing a significant reduction in GAPA protein levels.

In order to explore the impact of a decrease in the subunits of the GAPDH enzyme on photosynthetic capacity, the quantum efficiency of PSII (F_q '/ F_m '), chlorophyll a fluorescence was analysed (Baker, 2008; Murchie and Lawson, 2013). No significant decrease in F_q '/ F_m ' was observed in the gapa-I insertion line 164 which had a 45% reduction in GapA transcript levels and 46% reduction in GAPA protein or in the gapb insertion line 308 with no detectable level of the GAPB protein (**Table 1**). The GapA antisense lines, containing 46% of WT protein levels, maintained equivalent PSII photosynthetic efficiency to controls. GapB antisense lines also showed no significant differences in Fq'/Fm' consistent with the observed result in the gapb insertion line 308, suggesting that in the absence of GAPB, the presence of the GAPDH A subunit is sufficient to maintain the photosynthetic capacity of these plants.

A small decrease of 12% in F_q '/ F_m ' was found in the GapA co-suppressed lines that had the lowest level of GAPA protein (**Table 1**). To investigate further the impact of a combined reduction of both the GAPA and GAPB protein levels, double mutants gapa-1/gapb (164/308) were generated. Homozygous plants with insertions in both gapa-1 (164) and gapb (308) were grown as described above. Interestingly, double mutants of gapa-1/gapb (164/308) showed a significant reduction in F_q '/ F_m ' suggesting that a GAPA protein level of 52% is insufficient to maintain photosynthetic efficiency in the absence of GAPB.

Photosynthetic CO₂ assimilation and electron transport rates are reduced in lines with reduced GAPDH protein level

To assess the impact on photosynthesis of changes in the levels of GAPDH protein, CO_2 assimilation rates were determined as a function of internal CO_2 concentration (C_i) (A/C_i ' curve). Plants were grown in environmentally controlled chambers under short day conditions as described in materials and methods. The gas exchange measurements were made on

mature leaves on plants six weeks after germination. A/C_i curves were determined for the gapa-1 insertion line (164), gapb insertion line (308), co-suppressed line of GapA (cA), GapA antisense line (aA), GapB antisense line (aB) and gapa-1/gapb crossed line (164/308) compared to the controls (CN) plants (Fig. 3). From these A/C_i curves, the maximum rate of CO_2 assimilation (A_{max}) in all mutant lines tested was shown to be significantly lower than for the CN plants. The plants with the lowest levels of the GAPA protein had the greatest decrease in assimilation rate (Fig. 2A), with maximum assimilation rates attained in these plants being approx. 27% of that observed in the CN (Fig. 3A; Table 1). Furthermore, in the gapa-1 mutant (164), an approx. 50% reduction in GAPA protein levels resulted in a 40% reduction in maximum assimilation (Fig. 3A).

Plants with no detectable level of GAPB protein (**Fig 2B**) had a 30% decrease in assimilation rates compared to the 73% reduction observed in a line with 9% GAPA proteins (cA) (**Fig. 3B**). Finally, in line 164/308, representing the double mutant *gapa-1/gapb*, containing no GAPB protein and only 51% of the levels of GAPA protein, the assimilation rates are similar to the single *gapa-1* (164), and *gapb* (308) insertion mutants. This result suggests that the double mutant shows no cumulative impact on assimilation rates under these conditions as long as 51% GAPA protein remains (**Fig. 3C**).

Further analysis of the A/C_i curves using the equations published by von Caemmerer and Farquhar (1981) illustrated that the maximum rate of carboxylation by Rubisco (Vc_{max}) and maximum electron transport rate (J_{max}), were reduced in some lines (Sharkey et al., 2007; Sharkey, 2016) (Table 1). The results for Vc_{max} showed that lines with a reduction in GAPA displayed a significant decrease compared to CN. No significant difference in Vc_{max} was observed in plants with a reduction in GAPB.

Furthermore, the results showed that the lines with reductions in either GAPA or GAPB had a lower rate of photosynthetic electron transport (J_{max}), needed to sustain Ribulose

1,5-bisphosphate (RuBP) regeneration, when compared to control plants (Table 1). As previously noted, the maximum rate of CO₂ assimilation (A_{max}) was significantly lower in all lines compared to CN, however, A_{max} was significantly lower in cA, where GapA transcript and GAPA protein levels were at the lowest levels. No significant differences in A_{max} were observed between the single mutants gapa-1 (164) and gapb (308) compared to the double mutant gapa-1/gapb (164/308).

Growth and vegetative biomass are reduced in both GAPA and GAPB reduced lines

Growth analysis of GapA co-suppressed and insertion lines was carried out on homozygous plants grown in growth chambers at 22 °C under short day length (130 µmol m-2s-1 in an 8 h/16 h light/dark cycle) and relative humidity (RD) 50%. The growth rate of these plants was determined using image analysis of total leaf area over a period of 52 days from planting. Observations of the growth rates of *GapA* co-suppressed lines (cA), CN Columbia (Col-0), and the *gapa-1* and *gapb* insertion lines (164 and 308) showed a statistically significant reduction in all growth parameters (Fig. 4; Supplemental Fig. S2). The co-suppressed and insertion lines were shown to have a statistically significantly slower growth rate when compared to the CN plants at 40 days post planting (Fig. 4A and B). By 52 days post planting, this growth trend continued (Fig. 4B) and the final leaf area was reduced compared to controls (Fig. 4B).

A growth analysis of the *gapa-1* insertion mutant (164), the *GapA* co-suppressed (cA) and the *GapA* antisense lines (aA) showed a significant reduction in dry weight and leaf number (**Fig 5A**) compared to the CN. Significant reductions in the leaf number and final biomass were seen in the *gapb* insertion mutant (308) and *GapB* antisense lines was also observed when compared to CN (**Fig. 5B**). A comparative analysis of the single insertion mutants *gapa-1 and gapb* with the double mutants *gapa-1/gapb* showed that reduction in

both the A and B subunits resulted in a greater decrease in leaf area, biomass and leaf number after 46 days of growth (Fig. 5B), even in the absence of a larger decrease in assimilation rates observed in Fig. 3C (see Table 1).

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DISCUSSION

A reduction in GAPA protein levels inhibits CO₂ assimilation and reduces biomass yield

Previous research showed that a 60-70% reduction in GAPDH activity was needed to affect growth and development in tobacco (N. tabacum 'cv W38') antisense GAPDH lines and that no severe impact on photosynthesis was observed until levels were reduced to less than 35% of wild-type levels (Price et al., 1995). The results presented in this study clearly showed a slow growth phenotype in A. thaliana following reductions in GAPA protein levels by 50%. GapA co-suppressed lines, with more than a 90% reduction in GapA transcript levels and a barely detectable GAPA protein content showed the most statistically significant impact on photosynthetic efficiency (-73%) even with GAPB being present at wild-type levels. The principal form of GAPDH in plant chloroplast has been proposed to be the heterotetrameric A₂B₂. In some plants, including spinach, an A₄ homotetramer has also been detected representing up to 20% of total GAPDH activity (Scagliarini et al., 1998). This A₄ homotetramer was not detected in Arabidopsis by previous studies (Howard et al., 2011), providing evidence that under normal circumstances the A₂B₂ tetramer is the principal active form in Arabidopsis. In this study, plants showing an absence of GAPB protein in the gapb mutant lines-maintained photosynthesis rates at 66% of wild type levels suggesting that under conditions where GAPB is limiting, or absent, that the A₄ form of GAPDH can maintain photosynthesis.

Importantly, the work here also allowed a comparative analysis between plants with different levels of the GAPA and GAPB subunits under the same environmental conditions.

When the *gapa-1* (164) and *gapb* (308) mutant lines were crossed to form the double mutant *gapa-1/gapb* (164/308), the combined effects resulted in a cumulative reduction in biomass (-60%), which was significantly greater than the reductions observed in the *gapa-1* (-35%) or *gapb* (-16%) mutants alone. Interestingly, the assimilation rates for the *gapa-1/gapb* double mutant showed no further reductions compared to *gapa-1*, and *gapb* single mutants. Firstly, suggesting that GAPA, even though reduced in level, is able to maintain the assimilation rate even in the absence of GAPB and secondly, that the decrease in biomass observed in the double mutant may be due to impacts early in development leading to a cumulative effect on growth.

Recent reviews of the literature have shown that over-expressing of some CBC enzymes can lead to increases in photosynthesis and biomass and that a multi-target approach can result in cumulative yield gains in some plants (Simkin, 2019; Simkin et al., 2019; Raines, 2022). The co-overexpression of *GapA* and *GapB* in transgenic rice increased GAPDH activity to more than 3.2-fold of the wild-type levels; under elevated [CO₂], CO₂ assimilation increased by approximately 10% demonstrating that the overproduction of the chloroplast GAPDH proteins is effective at improving photosynthesis at least under elevated [CO₂]. However, under these conditions, no statistically significant differences in biomass were observed compared to wild-type plants, although a small increase in starch accumulation was observed. (Suzuki et al., 2021). In contrast, no statistically significant difference in CO₂ assimilation was observed in ambient [CO₂] (Suzuki et al., 2021). These results suggest that the manipulation of GAPDH activity may have more importance as atmospheric [CO₂] increases due to current climate change models where [CO₂] increases from 416 ppm to 550 by 2050 and 700 ppm by 2100 (Le Quéré et al., 2009; IPCC, 2019; NASA, 2020). Furthermore, given that no increase in growth rate or final biomass was observed at ambient [CO₂], increasing GAPDH may have more value

in a multi-target approach, such as targeting additional CBC enzymes, photorespiratory elements and photosynthetic electron transport in combination with GAPDH in the same plants.

CONCLUSION

Our results have shown that both GAPA and GAPB are essential for normal growth and development in Arabidopsis plants and that the A₂B₂ form of the enzyme is required for maximum photosynthetic efficiency. The phenotypes described in this manuscript provide *in vivo* evidence of the relative importance of the individual subunits of the GAPDH complex on photosynthetic carbon assimilation. In this study we also show that the suppression of GAPA to almost undetectable levels resulted in a 73% decrease in carbon assimilation compared to 34% reduction in photosynthesis in the absence of GAPB providing direct evidence of the importance of GAPA in maintain photosynthetic capacity.

MATERIALS AND METHODS

Identification and analysis of T-DNA GAPDH mutants and production of double

287 mutants

The gapa-1 and gapb mutants in Arabidopsis (Arabidopsis thaliana) were identified in the Arabidopsis Information Resource (TAIR) database (gapa-1: SAIL_164_D01 and gapb: SAIL_308_A06). The mutant insertion sites were identified by PCR and the location of each T-DNA insertion was determined by sequencing the PCR products spanning the junction site (Fig. 1). The GapA-1 was amplified with forward primers GapA1 Fwd (5'gagagcatgtgacataacggg'3) and reverse primer GapA1 Rev (5'accttaagcttggcctcagtc'3) in conjunction with primer Sail_LB3 (5'tagcatctgaatttcataaccaatctcgatacac'3). The GapB was amplified with forward primers GapB Fwd (5'cgacgatgtctcctctcagc'3) and reverse primer GapB Rev (5'gaccgggattcttgagacg'3) in conjunction with primer Sail_LB3. Double mutants

297 gapa-1/gapb (164/308) was obtained by crossing homozygous plants of gapa-1
298 (SAIL_164_D01) and gapb (SAIL_308_A06) and segregating the double homozygous
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Construct Generation

GAPA and GAPB antisense constructs

A partial-length coding sequence of glyceraldehyde-3-phosphate dehydrogenase A subunit (GapA-1: At3g26650) and the glyceraldehyde-3-phosphate dehydrogenase B subunit (GapB: RT-PCR At1g42970) amplified using primers AtGAPAf were by (5'cacctatcgaaggaaccggagtgtt'3) and AtGAPAr (5'tcctgtagatgttggaacaatg'3) and AtGAPBf (5'caccttgatggtaagctcatcaaagtt'3) and AtGAPBr (5'ggtgtaggagtgtggtggttgt'3) respectively. The resulting amplified products were cloned into pENTR/D (Invitrogen, UK) to make pENTR-GAPA1; pENTR-antiGAPA and pENTR-antiGAPB. The cDNA's were introduced into the pGWB2 gateway vector (Nakagawa et al., 2007) AB289765)) by recombination from the pENTR/D vector to make pGWB2-AntiGAPA and pGWB2-AntiGAPB (Supplemental Fig. S1). cDNA are under transcriptional control of the 35s tobacco mosaic virus promoter, which directs constitutive high-level transcription of the transgene, and followed by the nos 3' terminator.

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GAPA-1 sense constructs

Destination vector pGWPTS1 was generated as described in Simkin et al., (2017). The full-length coding sequencer of *GapA-1* was amplified using primers AtFwd (5'caccatggetteggttactttetetgtee'3) and AtRev (5'ttgatgaaatcacttecagttgttgg'3). The resulting amplified product was cloned into pENTR/D (Invitrogen, UK) to make pENTR-AtGAPA-1 and the sequence was verified and found to be identical. The full-length cDNA was

introduced into destination vector pGWPTS1 by recombination from the pENTR/D vector to make pGWPTS1-AtGAPA-1 (PTS1-GAPA-1) (Supplemental **Fig. S1**). The transgene was under the control of the rbcS2B (1150bp; At5g38420) promoter. In this instance the expression of the cDNA was under transcriptional control of the Rubisco small subunit 2B (rbcS2B) promoter (At5g38420), which directs high-level photosynthetic tissue specific transcription of the transgene and followed by the *nos* 3' terminator.

Generation of transgenic plants

The recombinant plasmid pGWB2-AntiGAPA, pGWB2-AntiGAPB and pGWPTS1-GAPA-1, were introduced into wild type Arabidopsis by floral dipping (Clough and Bent, 1998) using *Agrobacterium tumefaciens* GV3101. Positive transformants were regenerated on MS medium containing kanamycin (50mg L⁻¹). Kanamycin resistant primary transformants (T1 generation) with established root systems were transferred to soil and allowed to self-fertilize. Full details of pGWB2-AntiGAPA, pGWB2-AntiGAPB and PTS1-GAPA-1, construct assembly can be seen in the Supplemental **Fig. S1.**

Plant Growth Conditions

For experimental study, T3 progeny seeds from selected lines were germinated on soil in controlled environment chambers at an irradiance of 130 µmol photons m⁻² s⁻¹, 22°C, relative humidity of 60%, in an 8h/16h square-wave photoperiod. Plants were sown randomly, and trays rotated daily. Leaf areas were calculated using standard photography and ImageJ software (imagej.nih.gov/ij). Wild type plants and null segregants (azygous) used in this study were evaluated independently. Once it was determined that no substantial differences were observed between these two groups, wild type plants and null segregants were combined (null segregants from the transgenic lines verified by PCR for non-integration

of the transgene) and used as a combined "control" group (CN) (Supplemental **Fig. S3**). Four leaf discs (0.6 cm diameter) from two individual leaves, were taken and immediately plunged into liquid nitrogen, and stored at -80°C for determination of transcript levels by RT-qPCR and protein content by western blot.

cDNA generation and RT-qPCR

Total RNA was extracted from Arabidopsis leaf using the NucleoSpin® RNA Plant Kit (Macherey-Nagel, Fisher Scientific, UK). cDNA was synthesized using 1 μg total RNA in 20 μl using the oligo-dT primer according to the protocol in the RevertAid Reverse Transcriptase kit (Fermentas, Life Sciences, UK). cDNA was diluted 1 in 4 to a final concentration of 12.5ng μL⁻¹. For semi quantitative RT-PCR, 2 μL of RT reaction mixture (100 ng of RNA) in a total volume of 25 μL was used with DreamTaq DNA Polymerase (Thermo Fisher Scientific, UK) according to manufacturer's recommendations. For RT-qPCR, the SensiFAST SYBR No-ROX Kit was used according to manufacturer's recommendations (Bioline Reagents Ltd., London, UK). GAPA-1 (At3g26650) and GAPA-2 (At1g12900) transcript were amplified using primers GAPA-F (5'atggttatgggagatgatatgg'3) and GAPA-R (5'ttattggcaacaatgtcagcc'3) and GAPB-F (5'ttcaggtgctctgatgtctctacc'3) and GAPB-R (5' tagccactaggtgagccaaatccacc'3) respectively.

Protein Extraction and Western Blotting

Total protein was extracted in extraction buffer (50 mM 4-(2-Hydroxyethyl)piperazine-1-ethanesulfonic acid (HEPES) pH 8.2, 5 mM MgCl2, 1 mM Ethylenediaminetetraacetic Acid Tetrasodium Salt (EDTA), Glycerol 10% v/v, Triton X-100 0.1% v/v, 2 mM Benzamidine, 2 mM Aminocaproic acid, 0.5 mM Phenylmethanesulfonyl fluoride (PMSF) and 10 mM DTT). Any insoluble material was removed by centrifugation at

14000 g for 10 min (4°C) and protein quantification was determined as previously described (Harrison et al., 1998; Simkin et al., 2017). Samples were loaded on an equal protein basis, separated using 12% (w/v) SDS-PAGE, transferred to polyvinylidene difluoride membrane, and probed using antibodies raised against GAPDH (Pohlmeyer et al., 1996). Proteins were detected using horseradish peroxidase conjugated to the secondary antibody and ECL chemiluminescence detection reagent (Amersham, Buckinghamshire, UK).

Chlorophyll fluorescence imaging screening in seedlings

Measurements were performed on 2-week-old Arabidopsis seedlings that had been grown in a controlled environment chamber at 130 µmol mol⁻²s⁻¹ PPFD and ambient CO₂.Chlorophyll fluorescence parameters were obtained using a chlorophyll fluorescence (CF) imaging system (Technologica, Colchester, UK (Barbagallo et al., 2003; von Caemmerer et al., 2004)). The operating efficiency of photosystem two (PSII) photochemistry, F_q '/ F_m ', was calculated from measurements of steady state fluorescence in the light (F') and maximum fluorescence in the light (F_m ') since F_q '/ F_m ' = (F_m '-F')/ F_m '. Images of F' were taken when fluorescence was stable at 130 µmol m⁻² s⁻¹ PPFD, whilst images of maximum fluorescence were obtained after a saturating at 600 ms pulse of 6200 µmol m⁻² s⁻¹ PPFD (Oxborough and Baker, 2000; Baker et al., 2001; Lawson et al., 2008; Simkin et al., 2017).

Gas Exchange Measurements

The response of net photosynthesis (A) to intracellular CO_2 (C_i) was measured using a portable gas exchange system (CIRAS-1, PP Systems Ltd, Ayrshire, UK) as previously described (Simkin et al., 2017). Leaves were illuminated with an integral red-blue LED light source (PP systems Ltd, Ayrshire, UK) attached to the gas-exchange system, and light levels

were maintained at saturating photosynthetic photon flux density (PPFD) of 1000 μ mol m⁻² s⁻¹ for the duration of the A/C_i response curve. Measurements of A were made at ambient CO₂ concentration (Ca) at 400 μ mol mol⁻¹, before Ca was decreased to 550, 350, 215, 60 μ mol mol⁻¹ before returning to the initial value and increased to 740, 900, 1140, 1340, 1640 μ mol mol⁻¹. Measurements were recorded after A reached a new steady state (1-2 minutes). Leaf temperature and vapour pressure deficit (VPD) were maintained at 25°C and 1 \pm 0.2 kPa respectively. The maximum rates of Rubisco- (Vc_{max}) and the maximum rate of electron transport for RuBP regeneration (J_{max}) were determined and standardized to a leaf temperature of 25°C based on equations from von Caemmerer (1981), Bernacchi et al. (Bernacchi et al., 2001) and Sharkey (2016). All points below 200 ppm were assigned as rubisco-limited, points above 300 ppm as RuBP-regeneration limited as described (Sharkey, 2016).

Statistical Analysis

- All statistical analyses were done by comparing ANOVA, using Sys-stat (University of Essex, UK). The differences between means were tested using the Post hoc Tukey test (SPSS, Chicago).
- 414 Accession numbers
- Sequence data from this article can be found in the GenBank/EMBL data libraries under accession numbers At3g26650 (NM113576) and At1g42970 (AY039961).

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Author Contributions: A.J.S and M.A generated transgenic plants and performed molecular and biochemical experiments and carried out plant phenotypic and growth analysis. P.E.L.C and L.R.H screened and identified GPADH insertion mutants and assisted in experimental design. M.A performed gas exchange measurement on Arabidopsis. A.J.S and M.A carried out data analysis on their respective contributions. C.A.R conceived and designed the research and C.A.R and A.J.S supervised the research. C.A.R, A.J.S wrote the manuscript with input from all authors.

443 Tables

Table I. The quantum efficiency of Photosystem II (Fq'/Fm'), Maximum electron transport rate (J_{max}), the maximum rate of carboxylation by Rubisco (Vc_{max}) and maximum assimilation (A_{max}) in control and GAPDH lines in relation to reported protein levels.

Plants were grown in short days at 130 μ mol m⁻² s⁻¹ light intensity, 8 h light/16 h dark cycle. Values represent 4-6 plants independent lines (6-8 plants) for group. A_{max} , Vc_{max} and J_{max} derived from A/C_i response curves shown in Figure 3 using the equations published by von Caemmerer and Farquhar (1981) using the spreadsheet provided by Sharkey, (2016). ND = not detected. Statistical differences are shown in boldface (P < 0.05). SE are shown. WT = plants containing Wild Type levels of the transcript and protein subunit. Protein quantities are shown in italics.

Line	Relative % GapA transcript and protein	Relative % GapB transcript and protein	Fq'/Fm' 600 μmol m ⁻² s ⁻¹	J_{max}	Vc _{max}	A _{max}
CN	WT	WT	0.478 +/- 0.014	145.5 +/- 15.75	55.1 +/- 5.2	28.3 +/- 1.15
164	55.2 +/- 15.5 51.6 +/- 1.9	WT	0.479 +/- 0.004	89.0 +/- 11.12	40.2 +/- 5.62	16.2 +/- 2.71
cA	9.4 +/- 4.4 9.1 +/- 3.7	WT	0.420 +/- 0.014	51.9 +/- 4.41	37.6 +/- 3.19	7.67 +/- 0.91
aA	23.3 +/- 11.8 43.6 +/-5.1	WT	0.449 +/- 0.006	119.7 +/- 7.90	40.2 +/- 2.67	23.4 +/- 1.38
308	WT	ND	0.469 +/- 0.003	101.8 +/- 8.91	56.4 +/- 1.12	18.8 +/- 2.27
aВ	WT	22.3 +/- 9.34 26.1 +/- 13.8	0.453 +/- 0.004	110.5 +/- 17.12	52.3 +/- 5.92	18.2 +/- 2.34 ***
164/	55.2 +/- 15.5	ND	0.450 +/- 0.012	101.1 +/- 4.44	38.4 +/- 3.49	20.1 +- 0.54
308	51.6 +/- 1.9					

CN = control. *gapa-1* insertion (164); *gapb* insertion (308); co-supressed GapA (cA); antisense GapA (aA); antisense GapB (aB);

457	Figure 1	Legends
T J/	riguit	Legenus

- 459 Figure 1. Molecular analysis of homozygous GAPDH T-DNA insertion mutants.
- Structure of the two GAPDH genes and the location of T-DNA insertions in the (A) gapa-1
- 461 (At3g26650; SAIL 164 D01) and (B) gapb (At1g42970; SAIL 308 A06) mutants. Protein-
- 462 coding exons are represented by black and intron locations are displayed as inverted white
- 463 triangles above the coding sequence. Location of genomic PCR-screening primers are shown
- by black arrows on each gene model. T-DNA insertion sites are indicated by triangles below
- the sequence and the precise position is given as the number of base pairs from the ATG.
- 466 ATG, translation initiation codon; TGA, translation termination codon. Bolded G (Panel A)
- and C (Panel B) indicate the point of sequence insertion into the promoter region of the
- 468 SAIL 164 D01 and SAIL 308 A06 mutants.

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- 470 Figure 2. RT-qPCR and Immunoblot analysis of leaf proteins of wild type and
- 471 **experimental GAPDH plants.** (A) Transcript and Protein levels in gapa-1 (At3g26650;
- 472 SAIL 164 D01), GAPA co-suppressed lines (cA), GAPA antisense lines (aA) and control
- 473 (CN). (B) Transcript and Protein levels in gapb (At1g42970; SAIL_308_A06) and GAPB
- antisense lines (aB). Protein (6 µg) extracts from leaf discs taken from two leaves per plant
- and separated on a 12% acrylamide gel, transferred to membranes and probed with antibodies
- 476 to GAPDH which recognises both GAPA and GAPB subunits. Error bars represent SE of 3
- 477 plants per line.

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- Figure 3. Photosynthetic carbon fixation rate determined as a function of increasing
- 480 CO_2 concentrations (A/Ci) at saturating-light levels (1000 μ mol m⁻² s⁻¹). (A) Controls
- 481 (CN) compared to gapa-1 insertion line (164), GAPA co-suppressed (cA) and GAPA
- antisense (aA) lines(. (B) CN compared to gapb insertion line (308) and GAPB antisense
- 483 (aB) line lines and (C) Photosynthetic carbon fixation of CN compared to single insertion
- 484 mutants *gapa-1* (164) and *gapb* (308) and the double mutant *gapa-1/gapb* (164/308).
- Extrapolated data are in Table 1. Error bars represent SE of 6 plants per line).

- Figure 4. Growth analysis of control and experimental lines grown. (A) Plants were
- 488 grown at 130 μmol m⁻² s⁻¹ light intensity in short days (8h/16h days) for 52 days. White bar
- represents a size of 6cm. (B) Plant growth rate evaluated over the first 52 days. Lines co-

suppressing GAPA (cA), Controls (CN), gapa-1 insertion mutant (164) and gapb insertion mutant (308) are represented. Results are representative of 9 to 12 plants per line (CN plants include azygous lines segregated from primary transformants). Significant differences * (p<0.10); **(p<0.05); *** (p<0.01) are indicated. Unless indicated, results are presented as a percentage of CN (CN = 100%). Error bars represent SE.

Figure 5. Growth analysis of control and experimental lines grown in low light. (A) gapa-1 (164) and gapb (308) insertion mutants and GAPA co-suppressed (cA), GAPA antisense (aA) and GAPB (aB) antisense lines were analyzed in parallel. Results are representative of 8 plants per line. (B) gapa-1 (164) and gapb (308) insertion mutants and the double mutant gapa-1/gapb (164/308) crosses were evaluated. Results are representative of 9 to 12 plants per line. Plants were grown at 130 μmol m⁻² s⁻¹ light intensity in short days for 46 days. (CN plants include azygous lines segregated from primary transformants). Data were statistically analysed using 2-way ANOVA. Significant differences * (p<0.10); **(p<0.05); **** (p<0.01) are indicated. Results are presented as a percentage of CN (CN = 100%). Error bars represent SE.

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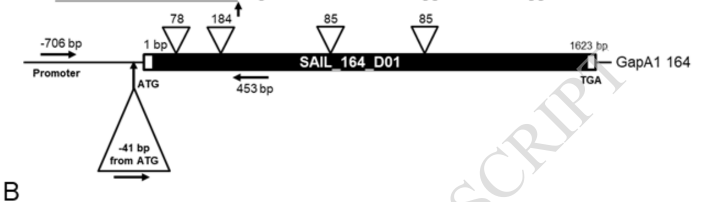
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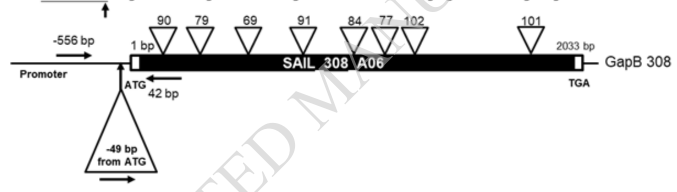
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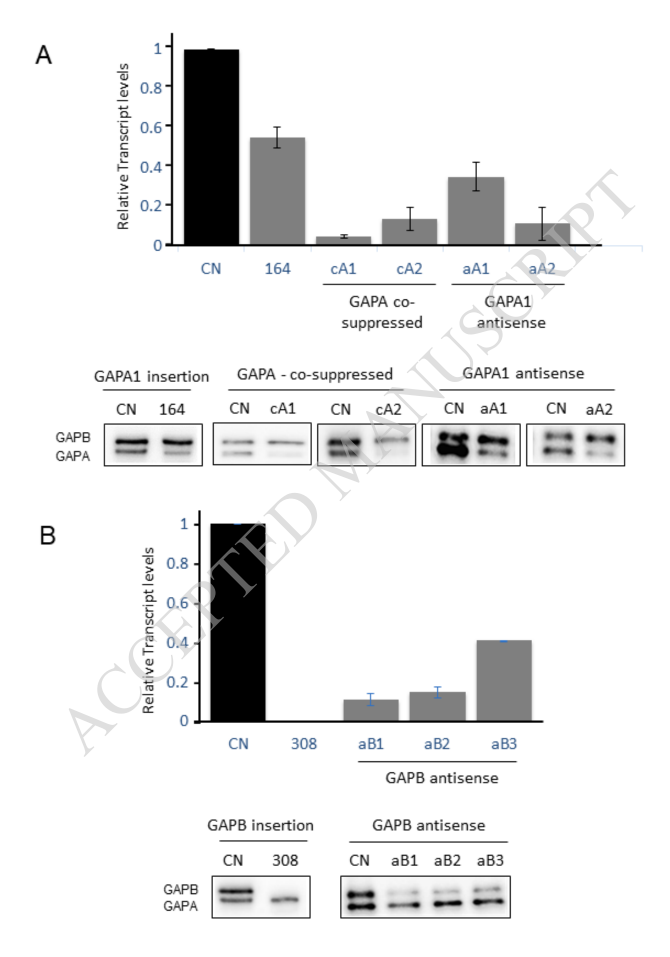
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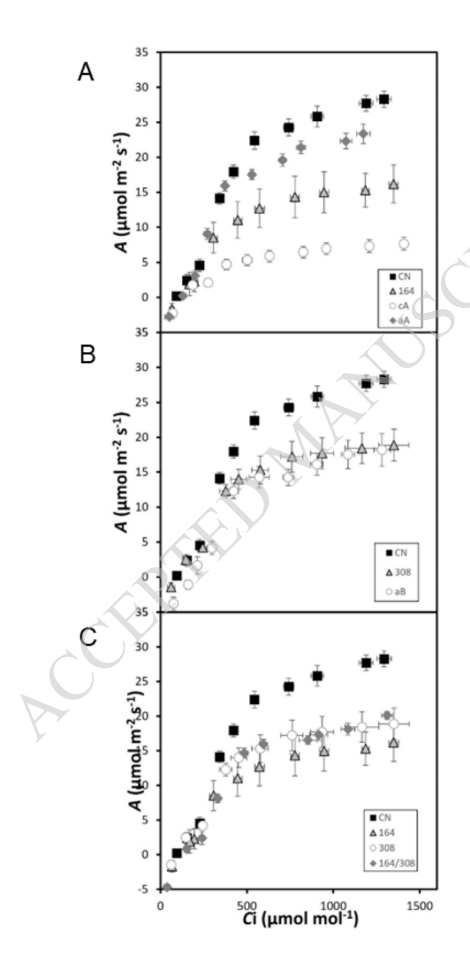
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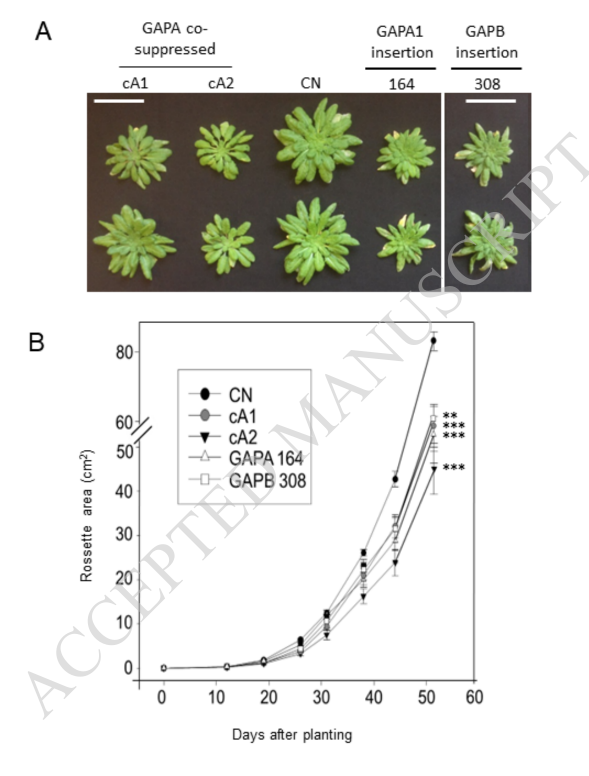


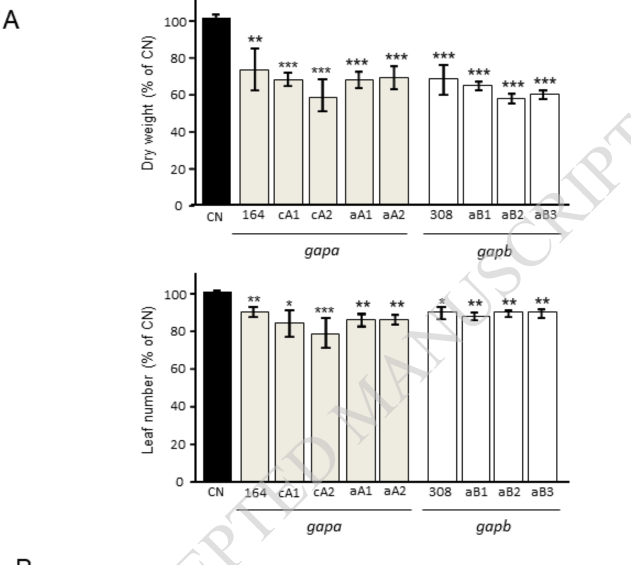
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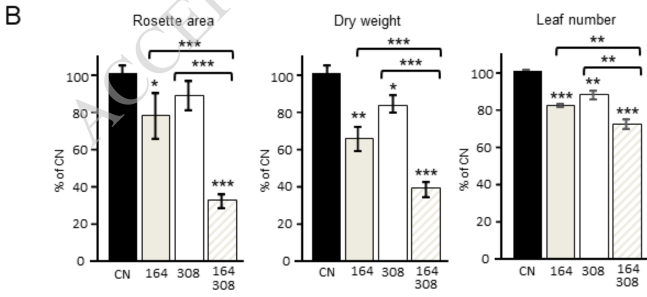












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