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#### **RESEARCH PAPER**

# Translatome and metabolome effects triggered by gibberellins during rosette growth in *Arabidopsis*

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#### **Abstract**

Although gibberellins (GAs) are well known for their growth control function, little is known about their effects on primary metabolism. Here the modulation of gene expression and metabolic adjustment in response to changes in plant (*Arabidopsis thaliana*) growth imposed on varying the gibberellin regime were evaluated. Polysomal mRNA populations were profiled following treatment of plants with paclobutrazol (PAC), an inhibitor of GA biosynthesis, and gibberellic acid (GA<sub>3</sub>) to monitor translational regulation of mRNAs globally. Gibberellin levels did not affect levels of carbohydrates in plants treated with PAC and/or GA<sub>3</sub>. However, the tricarboxylic acid cycle intermediates malate and fumarate, two alternative carbon storage molecules, accumulated upon PAC treatment. Moreover, an increase in nitrate and in the levels of the amino acids was observed in plants grown under a low GA regime. Only minor changes in amino acid levels were detected in plants treated with GA<sub>3</sub> alone, or PAC plus GA<sub>3</sub>. Comparison of the molecular changes at the transcript and metabolite levels demonstrated that a low GA level mainly affects growth by uncoupling growth from carbon availability. These observations, together with the translatome changes, reveal an interaction between energy metabolism and GA-mediated control of growth to coordinate cell wall extension, secondary metabolism, and lipid metabolism.

**Key words:** Gibberellin, growth, paclobutrazol, primary metabolism, translatome.

#### Introduction

Gibberellins (GAs) are tetracyclic diterpenoid plant growth hormones involved in the regulation of diverse physiological processes including seed germination, stem elongation, leaf expansion, root growth, and the development of reproductive organs (Olszewski et al., 2002; Hedden, 2003; Ueguchi-Tanaka et al., 2007; Schwechheimer and Willige, 2009). Mutants deficient in GA synthesis or signalling have germination deficiencies, remain dwarfed, and typically exhibit delayed flowering phenotypes. The metabolic pathways of GA biosynthesis and degradation, as well as elements of GA signalling pathways, have been reported (reviewed in Hedden and Phillips, 2000; Olszewski et al., 2002; Yamaguchi, 2008). Terpene cyclases convert geranylgeranyl diphosphate to

ent-kaurene which is then further channelled into ent-kaurenoic acid and GA<sub>12</sub> by ent-kaurene oxidase (KO) and ent-kaurenoic acid oxidase (KAO), respectively (Sun and Kamiya, 1997; Helliwell et al., 2001; Yamaguchi, 2008). Restricting KO activity in mutants limits GA biosynthesis in Arabidopsis (Helliwell et al., 1998). Similarly, inhibiting KO by the chemical paclobutrazol (PAC) results in a decrease in GA content concomitant with a reduction in growth (Rademacher, 2000). The inhibitory effect of PAC can be reversed by GA application, making it a valuable tool for gaining insights into the effects of GA on plant growth and metabolism (Lenton et al., 1987; Yim et al., 1997; Rademacher, 2000; Toh et al., 2008; Filardo et al.,

2009; Lu *et al.*, 2010). In the final step of the GA biosynthetic pathway,  $GA_{12}$  is converted to  $GA_4$ , a bioactive form, through oxidations by GA20-oxidase (GA20ox) and GA3-oxidase (GA3ox) (Hedden and Phillips, 2000; Olszewski *et al.*, 2002). Inactivation of bioactive GA occurs through  $2\beta$ -hydroxylation by GA2-oxidase (GA2ox). GA homeostasis is controlled by induction of GA2ox genes as well as negative feedback regulation of GA20ox and GA3ox expression by elevated levels of GA (Phillips *et al.*, 1995; Thomas *et al.*, 1999; Elliott *et al.*, 2001).

Despite many reports in the literature on the roles of GAs, as yet little information is available concerning their effects on the coordination of primary metabolism and growth. Among the studies carried out to date, the overexpression of GA2ox (involved in GA catabolism) or GA20ox (involved in GA biosynthesis) induced stunted or enhanced growth, respectively, in transgenic tobacco (Nicotiana tabacum) plants (Biemelt et al., 2004). Silencing GA2ox in tobacco triggered growth and the formation of xylem fibre cells (Dayan et al., 2010). Similarly, several 'Green Revolution' genes responsible for dwarfing traits in crops interfere with the action of GAs (Hedden *et al.*, 2003). In the study of Biemelt et al. (2004), vegetative growth was positively correlated with the rate of photosynthesis. In contrast, despite slow growth of GA-deficient mutants A70 and W335 of tomato (Solanum lycopersicum L.), the rate of photosynthesis per unit leaf area was unchanged relative to that of the wild type (Nagel and Lambers, 2002). There is also conflicting evidence with respect to the effect of chemical compounds that inhibit GA biosynthesis. For example, application of the KO inhibitor PAC reduces growth in rice, but it did not affect the rate of photosynthesis (Yim et al., 1997). Uniconazole is another plant growth retardant which primarily acts by inhibiting KO (Rademacher, 2000). Plants treated with uniconazole displayed growth depression, while net photosynthesis was stimulated (Thetford et al., 1995). Thus, it remains unclear how exactly GA is involved in the regulation of growth and metabolism.

Expression profiling studies have revealed many genes regulated by GA (Yamaguchi, 2008); however, in only a few cases has the physiological affect of GA on primary metabolism and its underlying genes been investigated. One example is OsPDK1 from rice, which encodes pyruvate dehydrogenase kinase, a negative regulator of the mitochondrial pyruvate dehydrogenase complex (mtPDC) that plays an important role in intermediary metabolism. The expression level of OsPDK1 increased after gibberellic acid (GA<sub>3</sub>) treatment, resulting in lowered pyruvate dehydrogenase activity. Blocking PDK1 expression in transgenic rice by RNA interference reduced vegetative growth compared with control plants, indicating that GA affects mtPDC activity by regulating the expression of OsPDK1 which then affects growth and biomass accumulation (Jan et al., 2006). This observation suggests that GA can challenge primary metabolism at the entry point of the tricarboxylic acid cycle.

The analysis of the GA signalling pathways led to the discovery of a class of negative regulators of growth called DELLA proteins, which in *Arabidopsis* are encoded by the

genes GAI (GA Insensitive), RGA (Repressor of GA 1-3), and RGL 1-3 (RGA-like 1-3). Arabidopsis plants lacking DELLA proteins show greater root elongation and biomass accumulation under salt stress than the wild type (Achard et al., 2006), and transgenic Populus trees overexpressing Arabidopsis DELLA-domainless versions of the DELLA proteins GAI and RGL1 had profound consequences for plant morphology and cellular metabolism, suggesting increased respiration in roots and reduced carbon flux through the lignin biosynthetic pathway in shoots as well as a shift towards defence compounds associated mainly with the phenylpropanoid pathway (Busov et al., 2006). Although these studies have advanced our understanding of the effect of GAs in specific developmental phases, they provide limited information concerning the general role of GAs in the regulation of plant metabolism and growth.

Here detailed kinematic analysis of Arabidopsis plants treated with PAC and/or GA3 is combined with metabolite profiling to provide an evaluation of GA-coordinated primary plant metabolism and growth. The expression of genes regulated by GA<sub>3</sub> in polysome-trapped RNA pools was also evaluated (as a measure of translational activity). For polysome isolation, a recently established procedure which allows efficient immunopurification of mRNAs in ribosome complexes by the use of a FLAG-tagged ribosomal protein L18 (RPL18) in Arabidopsis was used (Zanetti et al., 2005; Mustroph et al., 2009a, b). The metabolic and transcriptional/translational consequences of the altered GA levels were analysed, and the data collected are discussed in the context of current models linking plant energy metabolism to GA biosynthesis and growth. The analysis indicates that the effect of GAs on plant growth is an integral component of a large regulatory response of primary metabolism. The rise in GA level is a signal that integrates carbon metabolism and growth. However, under GA deprivation conditions, the relationships linking carbon availability and growth are modified, thus uncoupling growth from carbon availability.

#### Materials and methods

Plant material and growth conditions

Arabidopsis thaliana (L.) Heynh. Col-0 wild-type and 35S:HF-RPL18 seeds (provided by J. Bailey-Serres, University of California, Riverside, CA, USA) were sown on standard greenhouse soil (Stender AG; Schermbeck, Germany) in plastic pots with 100 ml capacity. The trays containing the pots were placed under a 16/8 h day/night or an 8/16 h day/night cycle (22/16 °C) with 60/75% relative humidity and 180  $\mu$ mol m $^{-2}$  s $^{-1}$  light intensity.

Paclobutrazol and gibberellic acid treatment

Fourteen days after sowing, plants growing singly in pots (100 ml) were watered with 10 ml of PAC solution (0.17 mM). For GA treatment, each plant was sprayed every second day with 1 ml of 50  $\mu$ M GA<sub>3</sub> containing 0.1% (v/v) Tween-80. *Arabidopsis* plants used in the assays were placed in trays together in a random arrangement with 35 pots per tray.

#### Kinematic analysis

Rosette area, number of rosette leaves, and rosette fresh weight (FW) and dry weight (DW) were measured at 2 d intervals from 14 d to 40 d after sowing. Rosette growth was described by the sigmoidal function:

$$f(t) = A/(1 + \exp\{-[(t - t_0)/b]\})$$
 (1)

where A is the difference between maximal and minimal growth values,  $t_0$  is the time when growth is maximal, and b is the steepness of maximum growth.

To calculate the rosette growth rate, the differential of the sigmoidal function (1) was determined:

$$d/dt \ f(t) = A \exp(-\{(t-t_0)/b\})/\left[b(1 + \exp\{-(t-t_0)/b\})^2\right]$$
(2)

Final rosette growth was calculated as the upper asymptote (A) of the sigmoidal curve. The moment of occurrence of the limiting point in the upper threshold sector of the curve indicated the duration of rosette growth, when a rosette reached 95% of its final growth. The peak of the first derivative of the curve corresponding to the highest growth rate occurred at the zero value of the second derivative.

The relative growth rate (RGR; mg g<sup>-1</sup> d<sup>-1</sup>) was calculated using the classical approach (Hunt, 1982):

$$RGR = In(M_2) - In(M_1)/t_2 - t_1$$
 (3)

where M1 and M2 are the plant mass at times  $t_1$  and  $t_2$ , respectively.

#### Metabolite analysis

Whole rosettes were harvested successively at 15, 20, 25, or 30 d after sowing (1, 6, 11, or 16 d after onset of PAC and/or GA<sub>3</sub> treatment), in the middle of the photoperiod. For metabolite analysis in entire plants (shoot and root), plants were harvested 27 d after sowing, when they were in the exponential growth phase. Harvests of 30 plants (six independent samples containing five whole rosettes or five root systems per samples) were performed per treatment and immediately frozen in liquid nitrogen. Chlorophyll, sucrose, starch, total protein, total amino acid, and nitrate contents were determined as described by Cross et al. (2006). Malate and fumarate contents were determined as described by Nunes-Nesi et al. (2007). Metabolite extraction for gas chromatography-mass spectrometry (GC-MS) was performed on the same samples as used for RNA extraction and metabolite determination by spectrophotometric methods. Derivatization and GC-MS analysis were performed as described previously in Lisec et al. (2006). NAD, NADH, NADP, and NAPH were determined following the protocol of Gibon and Larher (1997).

#### Measurements of photosynthetic parameters

Gas exchange measurements were performed with an open-flow gas exchange rates Li-Cor 6400 (Li-Cor Inc., Lincoln, NE, USA) with a portable photosynthesis system to fit a whole-plant cuvette. Light was supplied from a series of light-emitting diodes (LEDs) located above the cuvette, providing an irradiance of 300 µmol m<sup>-2</sup> s<sup>-1</sup>. The reference CO<sub>2</sub> concentration was set at 400 μmol CO<sub>2</sub> mol<sup>-1</sup> air. Dark respiration was measured on whole rosettes using the same protocol on plants kept in the dark. All measurements were performed at 25 °C, and the vapour pressure deficit was maintained at 2.0±0.2 KPa, whilst the amount of blue light was set to 10% of photon flux density to optimize stomatal aperture. Fluorescence emission measurements to estimate the actual flux of photons driving photosystem II were performed using a leaf chamber fluorometer (Model 6400-40, Li-Cor). Plants 20 d after sowing were used for determination of the photosynthetic parameters.

Total and polysomal RNA isolation

Polysomes from the 35S::HF-RPL18 line (27 d after sowing, as for metabolite analyses) were immunoprecipitated using powdered tissue and extracted in polysome extraction buffer, and clarified crude extract was incubated with ANTI-FLAG-Affinity GEL (Sigma-Aldrich) as described by Mustroph et al. (2009a). RNAs of the crude extract and immunoprecipitated eluate were extracted using an RNeasy Mini Kit (Qiagen, Hilden, Germany).

#### cDNA synthesis and real-time PCR analysis

cDNA was synthesized from 2 µg of polysomal and non-polysomal RNAs using Superscript III reverse transcriptase (Invitrogen, Darmstadt, Germany). Absence of genomic DNA contamination and RNA integrity were analysed as described (Piques et al., 2009). Real-time PCRs were performed in a 384-well microtitre plate with an ABI PRISM 7900 HT sequence detection system (Applied Biosystems Applera, Darmstadt, Germany), using Power SYBR Green PCR Master Mix. PCRs and data analysis were performed as described (Caldana et al., 2007; Balazadeh et al., 2008). PCR primers were designed using QuantPrime (Arvidsson et al., 2008). Three biological replicates were processed for each experimental condition.

#### ATH1 expression profiling and data analysis

Immunopurified RNA samples from GA<sub>3</sub>- and PAC-treated plants as well as non-treated control plants (two biological replicates for each treatment) were subjected to transcriptome profiling using Affymetrix ATH1 microarrays (Atlas Biolabs; http://www. atlas-biolabs.com/). For quality check and normalization, the raw intensity values were processed with Robin software (Lohse et al., 2010) using default settings. For background correction, the robust multiarray average normalization method (Irizarry et al., 2003) was performed across all arrays. A factorial design (PAC treatmentcontrol and GA<sub>3</sub> treatment-control) was applied for data analysis. Statistical analysis of differential gene expression for treatment versus control samples was carried out using the linear model-based approach (Smyth, 2004). The obtained P-values were corrected for multiple testing using the nestedF procedure, applying a significance threshold of 0.05 in combination with the Benjamini and Hochberg (1995) false discovery rate control. Expression data were submitted to the NCBI Gene Expression Omnibus (GEO) repository (http:// www.ncbi.nlm.nih.gov/geo/) under accession number GSE29699.

#### Statistical analyses

All experiments were designed in a completely randomized distribution. Analysis of variance (P < 0.05) was carried out to determine effects of treatments. Differences among means in figures and tables were examined by the Tukey or t-test. All statistical analyses were carried out using SPSS 8.0 for Windows statistical software (SPSS, Inc., Chicago, IL, USA).

#### **Results**

Inhibition of gibberellin biosynthesis reduces rosette expansion rate but not duration

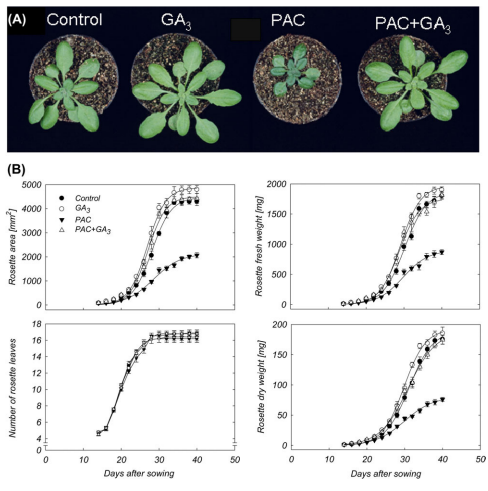
Despite the fact that many mutants with altered GA biosynthesis or signalling are known (Yamaguchi, 2008), the manipulation of GA content by PAC treatment was chosen in an attempt to monitor the translational regulation of individual mRNAs in transgenic Arabidopsis expressing a FLAG-tagged form of ribosomal protein L18. A benefit of this approach is that polysomes are purified from crude cell extracts without contamination by other mRNA

ribonucleoprotein complexes (Zanetti et al., 2005); furthermore, establishing transgenic GA mutants expressing the FLAG-tagged L18 was not needed. A clear decrease of rosette area was observed for 35S::HF-RPL18 Arabidopsis plants grown under a low GA regime, namely when plants were treated with PAC and grown side by side with controls (Fig. 1A). On the other hand, application of GA<sub>3</sub> alone caused a slight increase in the growth of the rosette as compared with control. Kinematic analysis of rosette growth revealed that FW, DW, and area of plants treated with PAC and/or GA<sub>3</sub> follow a sigmoidal function (Fig. 1B), revealing three growth phases: an early phase during which the absolute rate of rosette growth increased, a mid phase encompassing the period of the maximum absolute rate of rosette growth, and a late phase during which the absolute rate of rosette growth decreased. PAC treatment reduced the maximal rates of rosette expansion and FW and DW accumulation by  $\sim$ 60%, and rosette area and FW and DW by >50% at the end of the experiment (Table 1). However, the time at which maximal biomass accumulation occurred (on the basis of FW and DW, as well as rosette expansion) and the total duration of rosette growth were not affected by the PAC treatment (Table 1). Moreover,

PAC did not affect the total number of rosette leaves (Fig. 1B). Thus, the effects of a low GA regime on the rate of expansion and on the rate of FW and DW accumulation determine to what extent the final rosette area and biomass accumulation is affected by the GA deficit. Importantly, growth inhibition induced by PAC was completely reversed by application of GA<sub>3</sub> (Fig. 1A, B; Table 1), suggesting that the application of PAC had a specific effect on GA biosynthesis. Similar results were obtained both in the wild type (Col-0; see Supplementary Fig. S1 available at JXB online) and the transgenic 35S::HF-RPL18 line, indicating that overexpression of the FLAG-tagged RPL18 protein does not influence the response to GA<sub>3</sub>.

Translatome profiling of paclobutrazol- and gibberellic acid-treated plants

Here, the immunopurification of mRNA–ribosome complexes (Mustroph *et al.*, 2009*a*) was used to discover genes underlying the contrasting growth behaviour of GA<sub>3</sub>- and PAC-treated plants. Global profiling of mRNAs in ribosome complexes was done by using Affymetrix ATH1 microarrays. To this end, ribosome-associated mRNA was



**Fig. 1.** Phenotypic changes of 35S::HF-RPL18 Arabidopsis plants caused by treatment with PAC and/or GA<sub>3</sub>. (A) Shoots of 27-day-old plants. (B) Time course of rosette growth of plants treated with PAC and/or GA<sub>3</sub>. Rosette growth was described by the sigmoidal function  $y = A/(1+\exp\{-[(x-x_0)/b]\})$ . Values are presented as means  $\pm$ SE of 10 individual determinations.

Table 1. Effect of GA regime on components of growth dynamics [final rosette area, final rosette fresh weight (FW) and dry weight (DW); maximal rosette expansion rate, maximal rate of FW and DW accumulation; time to maximal rosette expansion, time to maximal FW and DW accumulation; rosette expansion duration, duration of FW and DW accumulation] of plants treated with PAC and/or GA<sub>3</sub> Comparisons were made in each column by Tukey test at the 5% level. Values are presented as means ±SE of 10 individual plants. Identical letters indicate values that are not statistically different.

Treatment	Final rosette area (mm²)	Maximum rosette expansion rate (mm <sup>2</sup> d <sup>-1</sup> )	Time to maximum rosette expansion (d)	Rosette expansion duration (d)
Control	4473±174 a	433±19 a	28±2 a	36±2 a
$GA_3$	4871±168 a	474±15 a	27±1 a	34±2 a
PAC	2139±104 b	147±8 b	28±1 a	37±1 a
PAC+GA <sub>3</sub>	4501±208 a	454±23 a	27±2 a	34±3 a

	Final rosette FW (mg)	Maximum rate of FW weight accumulation (mg d <sup>-1</sup> )	Time to maximum FW accumulation (d)	FW accumulation duration (d)
Control	1869±70 a	163±10 a	30±1 a	39±1 a
GA <sub>3</sub>	1969±48 a	178±5 a	29±2 a	37±2 a
PAC	900±39 b	67±2 b	29±2 a	39±1 a
PAC+GA <sub>3</sub>	1758±66 a	160±7 a	29±1 a	37±1 a

	Final rosette DW (mg)	Maximum rate of DW accumulation (mg d <sup>-1</sup> )	Time to maximum DW accumulation (d)	DW accumulation duration (d)
Control	190±8 a	15±1 a	31±1 a	41±1 a
GA <sub>3</sub>	193±10 a	17±1 a	30±2 a	38±2 a
PAC	81±3 b	5±1 b	30±2 a	41±2 a
PAC+GA <sub>3</sub>	189±7 a	13±1 a	31±2 a	42±1 a

isolated from plants treated for 13 d with either GA<sub>3</sub> or PAC at a final age of 27 d, corresponding to the time point of the maximal leaf expansion rate (Table 1). Analysis with the Robin software (Lohse et al., 2010) revealed that, compared with controls, 98 genes were differentially expressed by at least 2-fold after PAC treatment (39 up, 59 down), and 140 genes were differentially expressed upon GA<sub>3</sub> treatment (138 up, two down) (Supplementary Table S1 at JXB online; see Supplementary Fig. S2). The translatome profiles revealed major changes in the expression of genes associated with the cell wall, secondary metabolism, hormone signalling, and transcription factors (Table 2).

To confirm the array analysis, 18 genes representative of GA biosynthesis and signalling were selected in order to analyse their expression by quantitative reverse transcription-PCR (qRT-PCR). The expression of GA-related genes measured in plants treated with PAC and/or GA<sub>3</sub> relative to control plants (log<sub>2</sub> ratios) is shown in Fig. 2. Relative expression values of all measurements are provided in Supplementary Table S2 at JXB online.

The expression of the GA biosynthesis genes KO, KAO1, and KAO2 is regulated in a GA-negative feedback manner (Fig. 2A, B). Overall relative expression of KO, KAO1, and KAO2 is lower at the translatome level as compared with the total mRNA level. At the translatome level, only KO showed a significant down-regulation upon GA or PAC plus GA treatment, whereas KAO1 and KAO2 expression was not affected. In addition, GA20ox1, GA20ox2, and

GA20ox3 showed a similar response at the total RNA and translatome level, whereby they are down-regulated by GA<sub>3</sub>, but up-regulated by PAC, suggesting that their mRNA abundance directly influences polysome loading. For GA3ox1 and GA3ox2, the expression was downregulated by GA<sub>3</sub> and by PAC, although for GA3ox2 this was only observed at the total RNA level. Expression of genes for GA-inactivating enzymes (GA2ox1, GA2ox2, GA2ox3, GA2ox4, GA2ox7, and GA2ox8) was up-regulated in plants treated with GA<sub>3</sub> alone or PAC plus GA<sub>3</sub>. Moreover, GA-inactivating genes were down-regulated in PAC-treated plants (Fig. 2A, B). Only GA2ox7 and GA2ox8 did not show any response in the polysome fraction. The feedback and feedforward mechanisms also operate at the level of GA perception, with GA<sub>3</sub> alone or PAC plus GA<sub>3</sub> negatively regulating expression of GID1B, whereas PAC treatment up-regulated GID1B (Fig. 2A) both at the total mRNA and the translated mRNA level. In contrast, RGL1, RGL2, and RGL3, encoding DELLA proteins, showed upregulation upon GA<sub>3</sub> alone or PAC plus GA<sub>3</sub> treatment in both the total mRNA and polysome-associated fraction, and were down-regulated by PAC application.

Regulation of metabolism and growth processes in response to gibberellin

To characterize the effect of GA on primary metabolism, the levels of key primary metabolites were next monitored

**Table 2.** List of genes affected by  $GA_3$  or PAC treatment Expression changes are given as the  $log_2$  ratio of immunopurified polysomal mRNA from plants treated with PAC or  $GA_3$ .

AGI	Name	Log₂		Description	
		GA <sub>3</sub> -control	PAC-control		
Cell wall					
AT1G03870	FLA9	0.38	<b>-1.81</b>	Fasciclin-like arabinogalactan-protein 9	
AT1G10550	XTH33	1.02	-0.31	Xyloglucan:xyloglucosyl transferase	
AT1G11545	XTH8	0.04	<b>-1.55</b>	Xyloglucan endotransglucosylase/hydrolase 8	
AT1G35230	AGP5	1.07	-0.07	Arabinogalactan-protein (AGP5).	
AT1G69530	EXP1	-0.70	<b>-1.27</b>	α-Expansin gene family	
AT2G20870		1.14	<b>-2.08</b>	Cell wall protein precursor	
AT2G40610	EXP8	-0.52	-2.82	α-Expansin gene family	
AT2G43570	CHI	1.29	2.46	Chitinase, putative	
AT3G07010	J	0.07	-1.53	Pectin lyase-like protein	
AT3G10720		1.12	-1.19	Invertase/pectin methylesterase inhibitor	
AT3G29030	EXP5	0.04	-1 <b>.33</b>	α-Expansin gene family	
AT3G44990	XTH41	- <b>1.28</b>	0.49	Xyloglucan endo-transglycosylase	
AT3G45970	EXPL1	2.33	- <b>1.20</b>	Expansin-like	
AT4G02330	PMEPRCB	2.26	-0.12	Pectinesterase activity	
AT4G12730	FLA2	0.50	- <b>1.56</b>	Fasciclin-like arabinogalactan-protein 2	
AT4G25810	XTH23	1.09	0.56	Xyloglucan endotransglycosylase-related protein	
AT4G30270	XTH24	1.08	-0.48	Xyloglucan transferase in sequence	
AT5G49360	BXL1	0.58	−0.46 − <b>1.16</b>	$\beta$ -D-Xylosidase/ $\alpha$ -L-arabinofuranosidase	
AT5G57550	XTH25	1.27	0.15	Xyloglucan endotransglycosylase-related protein	
		3.46	- <b>1.99</b>	, ,	
AT5G57560	XTH22	3.40	- 1.99	Cell wall-modifying enzyme	
Primary metabolisi	m				
AT1G61800	GPT2	0.24	2.03	Glucose6-phosphate/phosphate transporter 2	
AT2G43820	SGT1	-0.02	1.16	UDP-glucose:salicylic acid glucosyltransferase	
AT3G10720		1.12	-1.17	Plant invertase inhibitor	
AT3G47380		-0.40	<b>-1.26</b>	Plant invertase inhibitor	
AT4G18010	5PTASE2	1.12	0.01	Inositol polyphosphate 5-phosphatase	
AT4G23600	CORI3	0.19	1.23	Encodes cystine lyase	
AT4G33150	LKR/SDH	0.24	1.11	Lysine-ketoglutarate reductase, lysine catabolism	
AT5G27420	CNI1 / ATL31	1.50	-0.31	Ubiquitin ligase that functions in the carbon/nitrogen response	
Secondary metabo	olism				
AT1G02205	CER1	0.52	<b>-1.74</b>	Aldehyde decarbonylase involved in wax biosynthesis	
AT1G03495		-0.06	3.02	Acyl-transferase family protein	
AT1G54040	TASTY	0.49	<b>-1.23</b>	Epithiospecifier protein, interacts with WRKY53	
AT3G29590	At5MAT	-0.12	1.41	Anthocyanidin 5-O-glucoside-6"-O-malonyltransferase	
AT3G55970	JRG21	-0.19	1.30	Oxoglutarate/iron-dependent oxygenase	
AT4G14090		-0.32	2.54	Anthocyanidin 5-O-glucosyltransferase	
AT4G22870		0.02	2.95	2-Oxoglutarate and Fe(II)-dependent oxygenase	
AT4G30470		1.21	-0.39	Lignin biosynthesis	
AT4G34135		1.26	0.73	Flavonol 7-O-glucosyltransferase	
AT4G37410	CYP81F4	-0.31	2.23	Indole glucosinate metabolism	
AT5G13930	TT4 / CHS	-0.01	1.84	Chalcone synthase	
AT5G17050		-0.02	1.08	Anthocyanidin 3-O-glucosyltransferase	
AT5G17220	TT19	-0.28	2.75	Glutathione transferase	
AT5G42800	TT3	0.17	2.87	Dihydroflavonol reductase	
AT5G54060	UF3GT	-0.04	2.15	UDP-glucose:flavonoid 3-O-glucosyltransferase	
Lipid metabolism					
AT1G06080	ADS1	0.65	<b>-1.67</b>	Δ9-Acyl-lipid desaturase	
AT1G06350	7.001	0.28	-1.86	Fatty acid desaturase family protein	
AT2G38180		-0.23	-1.00 1.12	SGNH hydrolase-type esterase	
AT2G38530	LTP2	-0.23 -0.18	2.13	Involved in lipid transfer between membranes	
				•	
AT3G02040	SRG3	1.06	0.24	Senescence-related gene 3	

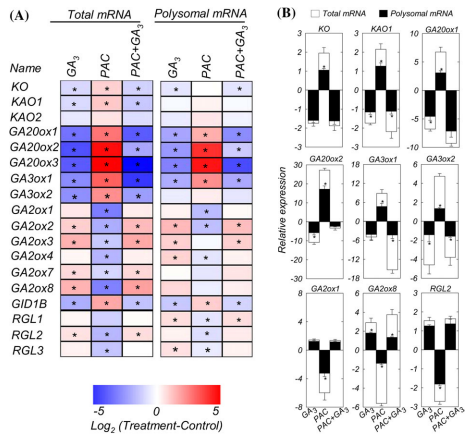
Table 2. Continued

AGI	Name	Log <sub>2</sub>		Description	
		GA <sub>3</sub> -control	PAC-control		
AT3G16370		0.10	-1.16	GDSL-like lipase/acylhydrolase	
AT3G56060		0.51	<b>-2.56</b>	Glucose-methanol-choline oxidoreductase	
AT4G18970		0.15	<b>-2.09</b>	GDSL-like lipase/acylhydrolase	
AT4G28780		0.21	<b>-1.67</b>	GDSL-like lipase/acylhydrolase	
AT4G26790		0.09	-1.41	GDSL-like lipase/acylhydrolase	
AT4G38690		0.48	<b>-1,21</b>	PLC-like phosphodiesterase	
AT4G39670		1.52	0.40	Glycolipid transfer protein (GLTP) family protein	
AT5G24210		1.11	-0.53	α/α-Hydrolases superfamily protein	
AT5G48490		0.33	<b>-1.46</b>	Bifunctional inhibitor/lipid-transfer protein	
Hormone biosyntl	hesis and signalling				
AT1G15550	GA3OX1	-0.70	2.32	Gibberellic acid biosynthetic pathway	
AT1G29440		-0.38	<b>-1.30</b>	SAUR-like auxin-responsive protein family	
AT1G29450		-0.37	<b>-1.48</b>	SAUR-like auxin-responsive protein family	
AT1G29500		-0.33	<b>-2.17</b>	SAUR-like auxin-responsive protein family	
AT1G29510	SAUR68	-0.36	<b>-1.49</b>	SAUR auxin-responsive protein family	
AT1G30040	GA2OX2	1.07	-0.60	Gibberellin 2-oxidase	
AT1G72520	LOX4	1.66	0.31	PLAT/LH2 domain-containing lipoxygenase	
AT2G21220		1.00	-0.49	SAUR-like auxin-responsive protein family	
AT2G30020	AP2C1	1.41	-0.17	PP2C-superfamily; ABA signalling	
AT2G34600	JAZ7	2.18	0.28	Jasmonic-acid responsive	
AT3G03840		-0.31	<b>-1.83</b>	SAUR-like auxin-responsive protein family	
AT3G25780	AOC3	1.23	0.42	Allene oxide cyclase; jasmonic acid biosynthesis	
AT3G48520	CYP94B3	1.26	0.44	Involved in catabolism of jasmonoyl-L-isoleucine	
AT3G53250		0.32	<b>-1.48</b>	SAUR-like auxin-responsive protein family	
AT3G57530	CPK37	1.40	0.02	ABA signalling	
AT4G11280	ACS6	2.23	-0.52	1-Aminocyclopropane-1-carboxylate (ACC) synthase	
AT4G25420	GA20OX1	-0.65	1.48	Gibberellin 20-oxidase	
AT4G34760		0.23	<b>-1.67</b>	SAUR-like auxin-responsive protein family	
AT4G38850	SAUR15	-0.68	<b>-1.52</b>	SAUR auxin-responsive protein family	
AT4G38860		-0.61	<b>-1.70</b>	SAUR-like auxin-responsive protein family	
AT5G07200	GA20OX3	-0.13	1.45	Gibberellin 20-oxidase	
AT5G51810	GA20OX2	-0.15	1.39	Gibberellin 20-oxidase	
AT5G45340	CYP707A3	2.44	-0.09	ABA 8'-hydroxylase activity; ABA catabolism	
AT5G56300	GAMT2	1.09	-0.15	Gibberellic acid methyltransferase 2	
Transcription fact					
AT1G21910	DREB26	2.33	-0.32	DREB subfamily A-5 of ERF/AP2 TF family	
AT1G33760		3.03	-0.06	DREB subfamily A-4 of ERF/AP2 TF family	
AT1G50420	SCL3	-0.43	1.24	Antagonist of DELLA proteins	
AT1G52830	IAA6	1.23	-1.05	IAA/AUX protein	
AT1G53160	SPL4	-0.71	1.38	Regulation of vegetative phase change	
AT1G56650	PAP1	-0.33	2.19	MYB TF involved in anthocyanin metabolism	
AT1G66350	RGL1	0.30	<b>-1.46</b>	Gibberellin signalling	
AT1G73805	SARD1	1.09	-0.41	Salicylic acid biosynthesis and signalling	
AT1G77640	MELOTIC	1.95	-0.72	DREB subfamily A-5 of ERF/AP2 TF family	
AT1G80840	WRKY40	3.27	0.12	Plant defence	
AT2G17040	ANAC036	1.73	-0.16	Leaf and stem morphogenesis	
AT2G33810	SPL3	0.35	<b>-1.46</b>	Regulation of vegetative phase change	
AT2G38470	WRKY33	1.83	0.07	Camalexin biosynthesis; defence	
AT2G40140	CZF1	1.24	-0.11	Stress responsive CCCH-type zinc finger	
AT2G46400	WRKY46	1.92	-0.47	WRKY family Group III	
AT3G49530	ANAC062	1.33	0.02	NAC domain protein involved in plant defence	
AT3G55980	SZF1	2.64	-0.50	Stress responsive CCCH-type zinc finger	
AT3G58120	bZIP61	0.84	<b>-2.75</b>	Encodes a member of the BZIP family	
AT4G17490	ERF6	1.39	-0.47	ERF subfamily B-3 of ERF/AP2 TF family	

Table 2. Continued

AGI	Name	Log <sub>2</sub>		Description
		GA <sub>3</sub> -control	PAC-control	
AT4G23810	WRKY53	2.78	-0.13	Regulator of flowering and senescence
AT4G25480	CBF3	<b>-1.16</b>	-0.89	DREB subfamily; cold acclimation
AT4G31800	WRKY18	2.06	0.46	Development-regulated defence response
AT4G34410	RRTF1	2.81	-0.10	ERF subfamily; redox homeostasis
AT5G04340	ZAT6	1.17	-0.15	C2H2 zinc finger; phosphate homeostasis
AT5G22380	ANAC090	2.20	0.06	NAC domain-containing protein
AT5G22570	WRKY38	1.94	-0.23	WRKY family Group III; plant defence
AT5G26920	CBP60g	1.68	0.31	Salicylic acid biosynthesis and signalling
AT5G39860	PRE1	-0.75	<b>-2.86</b>	bHLH136/Paclobutrazol resistance 1
AT5G49520	WRKY48	1.58	0.41	Stress-responsive WRKY member
AT5G62470	MYB96	1.01	-0.37	R2R3 MYB involved in ABA signalling
AT5G67450	AZF1	1.23	-0.11	Stress responsive zinc-finger protein

Significance (global test; P < 0.05) is indicated in bold.



**Fig. 2.** Changes in gene expression in shoots of *Arabidopsis* plants treated with PAC and/or  $GA_3$ , relative to control. (A) Heat map. Different shades of red and blue express the extent of the change according to the colour bar provided ( $log_2$  ratio of control); white indicates no change. Asterisks indicate values determined by the Student's *t*-test to be significantly different from the control (P < 0.05). For absolute values, see Supplementary Table S2 at *JXB* online. Primer sequences are given in Supplementary Table S5. (B) Relative expression of genes selected from the heat map. Data represent means  $\pm$ SE of three independent replicates (each replicate is a pool of five plants). Asterisks indicate a significant difference in gene expression between non-polysomal and polysomal fractions, as determined by the Student's *t*-test (P < 0.05).

across the time course of shoot development. Levels of carbohydrates, organic acids, total amino acids, and nitrate, as well as chlorophyll and total protein were determined in shoots of plants treated with PAC and/or GA<sub>3</sub>. Rosette FW

increased with time in all treatments (Fig. 3A). However, biomass accumulation increased more slowly in PAC-treated plants than in PAC plus GA<sub>3</sub>, GA<sub>3</sub> alone, or control treatments. Ln-transformation of these data revealed that

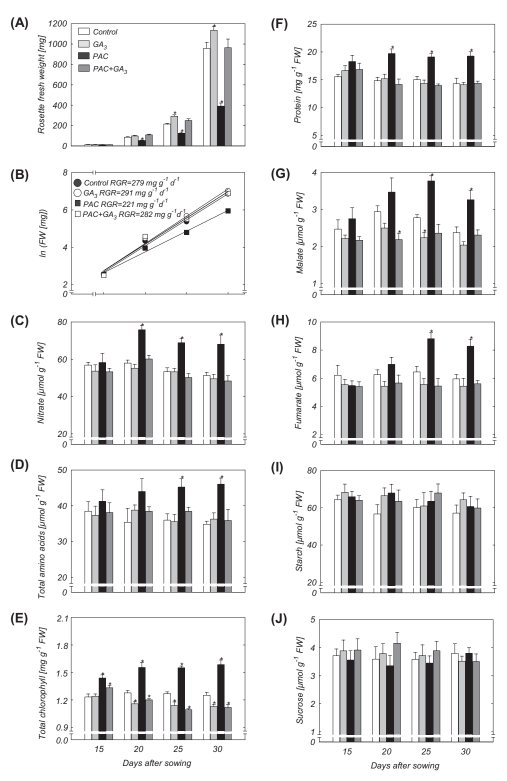


Fig. 3. Developmental changes of biomass and major metabolites in shoots of Arabidopsis plants treated with PAC and/or GA3. (A) Rosette fresh weight. (B) Relative growth rate over vegetative development. (C) Nitrate. (D) Total amino acids. (E) Total chlorophyll. (F) Protein. (G) Malate. (H) Fumarate. (I) Starch. (J) Sucrose. Data are means ±SE of six replicates (each replicate is a pool of five plants). Asterisks indicate values determined by the Student's t-test to be significantly different from the control (P < 0.05).

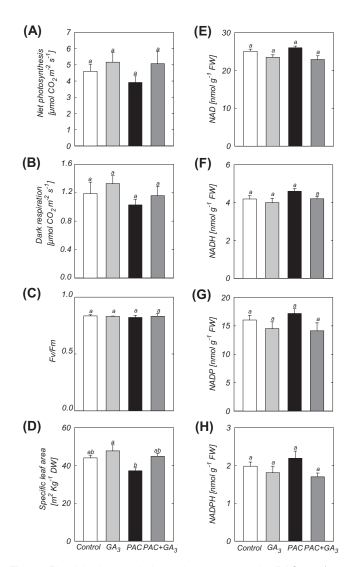
PAC-treated plants showed a reduction of ~20% in RGR across the entire period (15-30 d) in comparison with the other treatments (Fig. 3B). In rosette leaves from plants treated with PAC, nitrate levels increased from an initial value of 57  $\mu$ mol g<sup>-1</sup> FW at day 15 to ~73  $\mu$ mol g<sup>-1</sup> FW at days 20-30 (Fig. 3C). However, nitrate levels remained steady in plants treated with GA<sub>3</sub> alone, PAC plus GA<sub>3</sub>, and in controls during the course of the experiment. In

addition, total amino acids in plants treated with GA<sub>3</sub>, PAC, or PAC plus GA<sub>3</sub> initially remained at the same level as in the controls (Fig. 3D), but changed after PAC treatment at days 25-30. Furthermore, chlorophyll and total protein content were higher in plants treated with PAC than in the other treatments (Fig. 3E, F). The high levels of chlorophyll and protein in plants treated with PAC remained until the end of the experiment. In contrast, total protein contents of plants treated with GA<sub>3</sub>, or PAC plus GA<sub>3</sub>, were similar to those observed in the control plants. Moreover, total chlorophyll decreased in plants under GA<sub>3</sub>, or PAC plus GA<sub>3</sub>, treatment as compared with their respective controls 25–30 d after sowing (Fig. 3E). During rosette development, malate and fumarate levels increased significantly in PAC-treated plants at days 25-30, while they remained stable in plants exposed to GA<sub>3</sub> and PAC plus GA<sub>3</sub> (Fig. 3G, H). Starch and sucrose levels were similar in plants treated with PAC and/or GA<sub>3</sub> as compared with their respective controls 15–30 d after sowing (Fig. 3I, J).

In contrast to the reduction of RGR observed in PAC-treated plants (Fig. 3B), neither the rate of net photosynthesis nor the rate of dark respiration was significantly affected by the GA regime (Fig. 4A, B). The photochemical efficiency [maximum variable fluorescence/maximum yield fluorescence ( $F_v/F_m$ )] was also not affected by PAC and/or GA<sub>3</sub> treatment (Fig. 4C), and there was only a small decrease of specific leaf area (SLA; 15%) for whole plants treated with PAC compared with control (Fig. 4D). In agreement with the  $F_v/F_m$ , photosynthesis, and dark respiration results, no significant difference was observed in the pyridine nucleotide [NAD(P)H] levels between treatments (Fig. 4E–H). The DW/FW ratio was similar in controls and plants treated with GA<sub>3</sub>, PAC, or PAC plus GA<sub>3</sub> (0.091, 0.086, 0.093, and 0.088, respectively).

Comparison of the response of the gibberellin regime to a long and short photoperiod in shoot and root

As changes in the photoperiod will alter the amount of carbon fixed each day and the carbon/nitrogen balance in plants (Stitt et al., 2010), the levels of nitrate, total amino acids, sucrose, and protein were investigated in roots and shoots of plants treated with PAC and/or GA<sub>3</sub> to determine whether the inhibition of growth depends on the length of the photoperiod. GA<sub>3</sub> supply increased the shoot-to-root ratio by 1.39- and 1.46-fold, in plants grown under a long or short photoperiod, respectively (Fig. 5B). The increase in the shoot-to-root ratio resulted from a stimulation of shoot growth and a slight inhibition of root growth (Fig. 5A). On the other hand, both root and shoot growth were strongly inhibited in plants growing in the environment that limited GA biosynthesis (PAC treatment), but the shoot-to-root ratio was similar to that observed in control plants under a long or short photoperiod (Fig. 5A, B). GA<sub>3</sub> completely rescued shoot growth of PAC-treated plants, and root growth was also recovered by GA<sub>3</sub>. In roots of low GA plants, the levels of nitrate,



**Fig. 4.** Physiological analysis of plants treated with PAC and/or  $GA_3$ . (A) Rate of photosynthesis. (B) Dark respiration. (C)  $F_v/F_m$ . (D) Specific leaf area. Values are presented as means  $\pm SE$  of 10 individual determinations. (E–H) NAD(P)H levels. Bars labelled with the same letter are not statistically different at the 5% level (Tukey test). Data are means  $\pm SE$  of six replicates (each replicate is a pool of five plants).

total amino acids, protein, and sucrose were similar to those of high GA and control plants under both long- and short-day conditions (Fig. 5C–F). In contrast, nitrate, total amino acids, and protein content were increased under GA deficiency in the *Arabidopsis* shoots. Sucrose content (Fig. 5F) and starch levels (not shown) in shoots of low GA plants were similar to those of high GA and control plants in long- and short-day photoperiods. Thus the level of GA only affects the metabolic composition of the shoot, but not the root.

Changes in metabolite profiles in Arabidopsis leaves in response to gibberellin

In order to verify the effect of GA on other major pathways of primary metabolism, an established GC-MS protocol

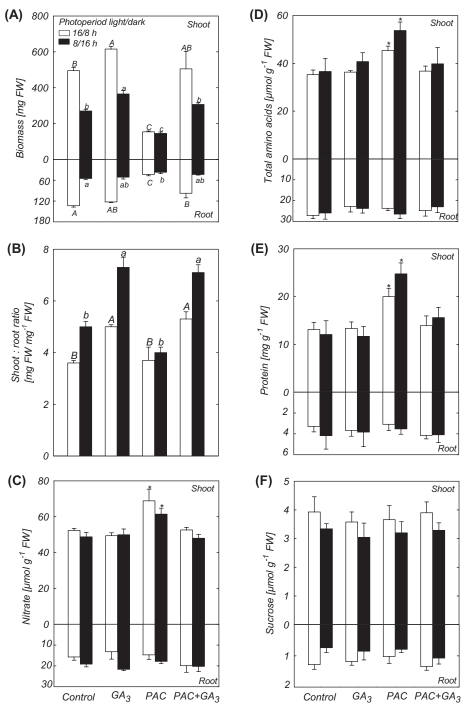
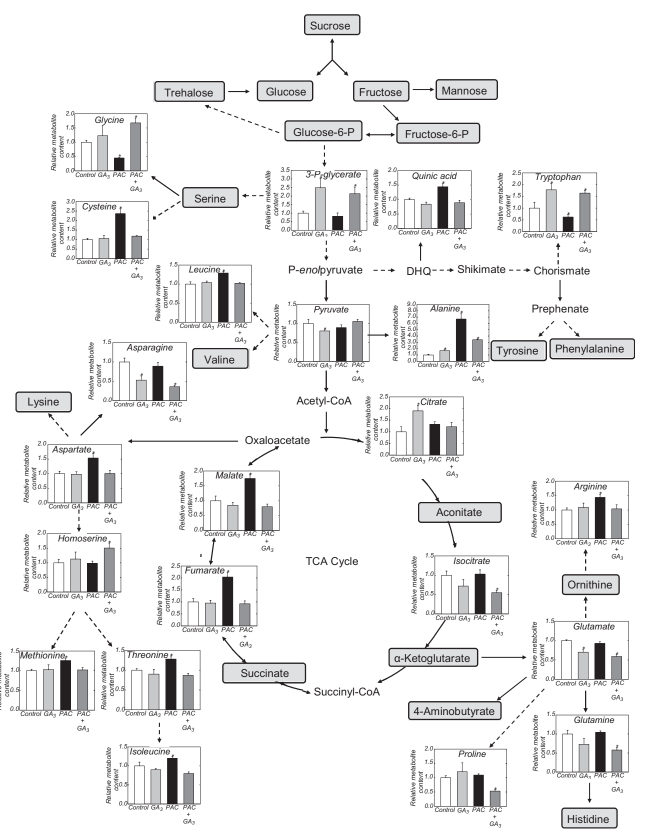


Fig. 5. Comparison of the growth response and metabolite levels in shoots and roots of plants treated with PAC and/or GA<sub>3</sub> in long and short photoperiods. (A) Biomass. (B) Shoot-to-root ratio. (C) Nitrate. (D) Total amino acids. (E) Protein. (F) Sucrose. Bars followed by the same upper case letter (long day) or followed by the same lower case letter (short day) do not differ significantly at the 5% level (Tukey test). Asterisks indicate values determined by the Student's t-test to be significantly different from control (P < 0.05). Values are presented as means of six replicates (each replicate is a pool of five plants) ±SE.

was used for metabolite profiling (Fernie et al., 2004). The analysis revealed that the GA regime did not lead to significant changes in the levels of sugars such as glucose, fructose, mannose, and sucrose. Moreover, there were no significant differences in hexose-phosphates in plants treated with PAC and/or GA<sub>3</sub>. Analysis of amino acid levels revealed an increase in levels of cysteine, leucine, alanine,

aspartate, methionine, threonine, isoleucine, and arginine in plants growing under a low GA regime (PAC treatment), and a decrease in glycine and tryptophan content (Fig. 6; see Supplementary Table S3 at JXB online). On the other hand, there are minor changes in amino acid levels in plants treated with GA<sub>3</sub> alone, or PAC plus GA<sub>3</sub>. Serine content was unaltered by PAC and/or GA treatment, while the



**Fig. 6.** Changes in metabolite profiles in shoots of plants treated with PAC and/or  $GA_3$ . Metabolites without a significant difference between treatments are indicated by a grey square. Metabolites outside grey squares indicate that they were not measured. Continuous arrows indicate a one-step reaction, and broken arrows indicate a series of biochemical reactions. Values are presented as means of six replicates (each replicate is a pool of five plants)  $\pm SE$ . Asterisks indicate values determined by the Student's *t*-test to be significantly different from control (P < 0.05). DHQ, 3-dehydroquinate. A complete list of all metabolites measured by GC-MS can be found in Supplementary Table S3 at *JXB* online.

precursor, 3-P-glycerate, was higher in plants treated with GA<sub>3</sub> alone or PAC plus GA<sub>3</sub>. Glycine and cysteine are linked closely to serine formation. The reduced GA availability caused an increase (>2-fold) in cysteine. Furthermore, glycine was decreased by PAC treatment, while PAC plus GA<sub>3</sub> increased the glycine level, and GA3 alone did not change the glycine content. PAC-treated plants showed an increase in the level of quinic acid, derivatives of which are involved in the synthesis of aromatic amino acids and secondary metabolites (Tzin and Galili, 2010). Among the aromatic amino acids tryptophan showed a clearly elevated level in plants treated with GA<sub>3</sub> alone or PAC plus GA<sub>3</sub>, while the tryptophan content was consistently reduced in plants under GA deficiency (Fig. 6). On the other hand, tyrosine and phenylalanine content remained steady across all treatments.

The level of alanine, a pyruvate-derivated amino acid, was increased >6-fold in PAC-treated plants (Fig. 6). GA<sub>3</sub> decreased the alanine content of PAC-treated plants, while GA<sub>3</sub> alone induced a slight increase in the alanine level. A significant increase in leucine content was only observed in PAC-treated plants, while the valine content remained at the same level in all treatments. Extending this analysis to the precursor of the amino acid family branch, only a marginal decrease of pyruvate was observed in plants treated with GA<sub>3</sub> alone.

Significant changes of aspartate were only observed on PAC treatment (level increased by up to 54%). In plants, aspartate is also precursor of the essential amino acids asparagine, methionine, threonine, and isolecine (Jander and Joshi, 2010). PAC-treated plants showed an increase in the level of methionine, isoleucine, and threonine (Fig. 6). In contrast, methionine, isoleucine, and threonine contents in plants treated with GA<sub>3</sub> alone or PAC plus GA<sub>3</sub> were similar to those observed in control plants. The results also showed a decrease in asparagine content of 47% and 64%, as compared with control, in plants treated with GA<sub>3</sub> alone or PAC plus GA<sub>3</sub>, respectively. On the other hand, asparagine remained at the same level in low GA plants as in the control. Plants treated with PAC and/or GA<sub>3</sub> had no marked effect on 4-ketoglutarate levels (Fig. 6). However, there were reductions in the levels of the amino acids glutamate and glutamine only in plants treated with GA<sub>3</sub> alone or PAC plus GA<sub>3</sub>. In the present experiments, arginine accumulated only in plants grown under a low GA regime. Proline content was unaltered in plants treated with PAC or GA<sub>3</sub> alone, but PAC plus GA<sub>3</sub> led to a decrease of proline content.

### **Discussion**

Gibberellin modifies translation of individual mRNAs

Cell expansion involves the selective loosening and rearrangement of the cell wall to induce turgor-driven growth (Marga et al., 2005). Expansins (Cosgrove, 2000) and xyloglucan endotransglucosylase/endohydrolases (XTHs) (van Sandt et al., 2007) are the two major protein classes

known to drive this process. Treatments with GA<sub>3</sub> or PAC have opposing effects on the expression of genes encoding these proteins, which correlates well with the observed changes in the rosette expansion rate (Table 1). At the level of primary metabolism changes were observed for several genes of cysteine, tryptophan, or lysine metabolism after PAC treatment. GA<sub>3</sub> induced the expression of CNII, which encodes a RING-type ubiquitin ligase (Sato et al., 2009) that associates with 14-3-3 proteins which in turn regulate carbon/nitrogen metabolism by directly binding essential enzymes involved in carbohydrate and nitrogen metabolism (Sato et al., 2011). On the other hand, genes involved in anthocyanin and flavonoid biosynthesis showed a strong up-regulation upon PAC treatment, but no effect of GA<sub>3</sub> application on the expression of these genes was observed. Fatty acid biosynthesis in plants is adjusted to the need for membrane biogenesis during growth or repair (Ohlrogge and Jaworski, 1997). GA deprivation mainly resulted in a decrease in the expression of genes associated with lipid metabolism, while several genes showed an upregulation after GA<sub>3</sub> treatment, correlating with the differences in growth rate. Interestingly, the changes in GA level influence the expression of genes involved in the biosynthesis and response to the plant hormones salicylic acid (SA), jasmonic acid, abscisic acid (ABA), and auxin, suggesting cross-talk between these hormones and GA status, GA<sub>3</sub> treatment results in the up-regulation of CBP60g and SARD1, two transcription regulators of SA biosynthesis and signalling (Zhang et al., 2010; Wang et al., 2011). Furthermore, SA-responsive transcription factors belonging to the WRKY family are highly induced after GA<sub>3</sub> treatment. Taken together, these results indicate that growth is under the direct control of GA levels.

The quantitative profiling of alterations in steady-state and polysomal mRNA populations in response to GA revealed that feedback regulation of GA biosynthesis genes and feedforward regulation of GA catabolism genes also operate at the level of the translatome (Fig. 2). On the other hand, several mRNAs encoding proteins responsible for GA biosynthesis and GA signal transduction induced at the steady-state level were modestly or strongly repressed in the polysome fractions of plants treated with PAC and/or GA<sub>3</sub>. Thus, differential mRNA translation appears to be a crucial mechanism for the control of feedback regulation of GA-related genes and thus biomass accumulation.

Changes in growth and primary metabolism are interlinked with gibberellin level

In an attempt to clarify the effect of GA on primary metabolism and growth, the major carbon metabolites as well as nitrate, chlorophyll, and total protein were determined across plant development. The increase in nitrogen content in shoots of PAC-treated plants was accompanied by an increase in total protein and chlorophyll (Fig. 3). Since leaf expansion was impaired by the low GA regime, the accumulation of protein and chlorophyll in low GA plants may reflect an indirect effect of GA. However, PAC

also increased the levels of protein and chlorophyll in mature leaves (see Supplementary Table S4 at JXB online), suggesting close coordination by the GA regime. The changes in nitrogen levels were also consistent with an increase in malate and fumarate concentration, which acts as a counter-anion for pH regulation during nitrate assimilation (Benzioni et al., 1971; Tschoep et al., 2009). In rosettes of plants grown under a low GA regime, a large amount of fumarate accumulated specifically 25-30 d after sowing. It is also known that fumarate serves as an alternative and flexible sink for photosynthate in Arabidopsis (Chia et al., 2000; Pracharoenwattana et al., 2010). The observation that fumarate is present at high levels in A. thaliana (Chia et al., 2000) in comparison with other plants species (Araújo et al., 2011) provides further evidence that fumarate constitutes a significant fraction of the fixed carbon in this species. In this context, fumarate accumulation in Arabidopsis shoots under low GA regimes may provide an adaptive advantage to allow rapid growth when GA becomes available. Consistent with this hypothesis, GA<sub>3</sub> application to PACtreated plants completely rescued their growth and concomitantly decreased the fumarate content to levels similar to those observed in control plants (see Supplementary Fig. S3 at JXB online).

Despite the fact that biomass is strongly decreased in low GA plants, a relatively small decrease in the RGR was observed. In addition, a reduced capacity for GA biosynthesis also led to a slight decrease in SLA, but did not affect photosynthesis, dark respiration, or the photochemical efficiency of photosystem II  $(F_v/F_m)$  (Fig. 4). These results provide compelling evidence that the reduced biomass was not a consequence of variation in photosynthesis or respiration rates per se. In agreement with these observations, levels of pyrimidine nucleotides [NAD(P)H] were not affected by PAC and/or GA<sub>3</sub> treatment (Fig. 4). Leaf growth is known to be highly dependent on carbon availability (Wiese et al., 2007; Pantin et al., 2011). The low GA plants contained levels of starch and sugars similar to those of GA-treated plants (Fig. 3). However, the maximal rosette expansion rate as well as the maximal FW and DW accumulation rates were strongly affected by the low GA regime. As a consequence, the final leaf area and the final rosette FW and DW were reduced in PAC-treated plants (Table 1). Together, these data indicate that the GA level orchestrates carbon allocation and growth. Interestingly, the duration of rosette expansion and FW and DW accumulation was not affected by the GA regime. This observation suggests that the rosette expansion rate as well as the rate of FW and DW accumulation are more flexible than the duration of leaf elongation and biomass accumulation upon GA deprivation. The observed increase in nitrate, total amino acid, and protein levels in shoots of low GA plants grown under longday conditions (Fig. 3) was also found when low GA plants were grown in an 8 h/16 h (light/dark) photoperiod, indicating a conserved effect of GA deprivation on primary metabolism. The assimilation of inorganic nitrogen into amino acids and the subsequent metabolic conversion of amino acids into protein are energetically expensive processes

(Hachiya *et al.*, 2007). Thus, the accumulation of compounds with high energy storage capacity in shoots of low GA plants can be interpreted as the result of rosette elongation being more reduced than carbon inflow, indicating an uncoupling between carbon availability and shoot elongation. In contrast to GA-deprived plants, GA<sub>3</sub>-treated plants or plants treated with PAC plus GA<sub>3</sub> did not accumulate compounds with high energy storage capacity, showing that GA availability couples primary metabolism and growth.

In contrast to shoots, the alterations in GA level had no effect on the metabolic status of the root, which suggests that retaining compounds with high energy storage capacity in the leaves is an active process. Because nitrate levels in root remained low in all treatments, it seems unlikely that the rate of nitrogen uptake and unloading to the xylem would be affected by the GA regime in either long- or shortday conditions. Interestingly, limiting GA biosynthesis by PAC treatment led to a strong inhibition of shoot and root growth, but the shoot-to-root ratio resembled that of the control plants in long or short photoperiods. These data reflect an unaltered carbohydrate status of low GA plants as compared with control plants. In agreement with this statement, PAC treatment did not change steady-state sucrose and starch levels. Furthermore, PAC treatment did not lead to substantial changes in the levels of hexosephosphates (Fig. 6). Taken together this indicates that the entry points of carbon into starch synthesis, glycolysis, and cell wall biosynthesis were not affected by the low GA regime. When considered alongside the observed changes in energy metabolism and growth, this strongly implies that the reduction in biomass in low GA plants originates from uncoupling energy metabolism (carbon supply) and growth (demand).

# Gibberellin modulates global changes in primary metabolism

One conspicuous feature of the GA-dependent metabolite profiles was the changes in amino acids and some of their precursors (Fig. 6). The carbon backbone of glycine and cysteine is derived from serine and 3-P-glycerate. 3-P-glycerate is also an intermediate in plastidial phospholipid synthesis and forms the primary substrate for triacylglycerol synthesis (Gibon *et al.*, 2006). It was previously shown that high 3-P-glycerate levels correlate with a high plant growth rate (Meyer *et al.*, 2007), which is in accordance with the observation that its levels are highly induced upon GA<sub>3</sub>, and slightly reduced upon PAC treatment (Fig. 6).

The analysis of aromatic amino acids revealed that GA<sub>3</sub> only affected the tryptophan pool. The increase of the tryptophan pool in plants treated with GA<sub>3</sub> alone or PAC plus GA<sub>3</sub>, as well as the reduction of tryptophan level in plants treated with PAC alone indicates an effect of GA on enzyme activity or substrate availability downstream of this aromatic amino acid. Such an effect could explain certain prominent symptoms of GA availability on growth dynamics such as, for example, the dwarf shoot in low GA plants and accelerated growth following supply of GA<sub>3</sub>. Although

plants have multiple pathways for indole-3-acetic acid (IAA) biosynthesis (Zhao, 2010), the conversion from tryptophan to either indole-3-pyruvic acid (via tryptophan transaminase and indole pyruvate decarboxylase), tryptamine (via tryptophan decarboxylase and amino oxidase), or indole-3-acetaldehyde (via indole aldehyde dehydrogenase) appears to be directly or indirectly disrupted by GA deficiency. Thus, the observed reduction in tryptophan level in low GA plants might contribute to explain the severe decrease in shoot growth in low GA plants since tryptophan deficiency has recently been associated with retardation of aerial organ development by affecting cell expansion (Jing et al., 2009). In light of this fact, GA<sub>3</sub> was able to recover growth of plants treated with PAC and this was accompanied by an increase in tryptophan level. Furthermore, expression of various auxin-up-regulated SAUR (Small Auxin Up RNA) genes was down-regulated in rosettes of low GA plants as compared with control or GA-treated plants (Table 2). In other words, SAUR transcripts are most abundant in tissues that are elongating or programmed to elongate in response to GA, linking growth with the metabolic availability of tryptophan.

Alanine aminotransferase catalyses the interconversion of pyruvate and glutamate to 4-ketoglutarate and alanine (Rocha et al., 2010). Given that PAC reduced plant growth and increased the alanine level, it seems feasible that alanine could contribute to regulate the biosynthesis of pyruvate, facilitating the maintenance of both the carbon/nitrogen balance and the rate of respiration in the plant. In keeping with this observation, GA<sub>3</sub> supplementation rescues shoot and root development and decreases the level of alanine in PAC-treated plants. It is well known that hypoxia alters the carbon/nitrogen balance and alanine accumulates in large amounts via an increase in alanine aminotransferase activity (Miyashita et al., 2007; Rocha et al., 2010). In the present studies, it was found that PAC treatment triggers an enhanced alanine level, and a decreased rosette growth, suggesting a modified relationship between energy metabolism and growth in low GA plants. Glutamine and glutamate serve as nitrogen transport compounds and nitrogen donors in the biosynthesis of several compounds (Masclaux-Daubresse et al., 2006). Nitrogen may be subsequently channelled from glutamine and glutamate to aspartate by aspartate aminotransferase, or to asparagine by asparagine synthase. As the amino acids glutamine and asparagine carry an extra nitrogen atom in the amide group of their side chains they play an important role as nitrogen carriers in cellular metabolism (Urquhart and Joy, 1981). Thus, the reduced levels of asparagine and glutamate observed in plants treated with GA<sub>3</sub> alone and PAC plus GA<sub>3</sub> indicate that both amino acids are being utilized more rapidly when plant growth is stimulated by the hormone. In agreement with this model, plants treated with GA<sub>3</sub> alone and PAC plus GA<sub>3</sub> showed a fast increase in the rate of leaf area growth and biomass accumulation. Together, these data indicate that GA is required for connecting energy metabolism and growth. Aspartate and arginine serve as important nitrogen reserves and intermediates in nitrogen

recycling. These two amino acids accumulated consistently only in low GA plants. Moreover, glutamate, glutamine, and asparagine levels were not altered in low GA plants, showing the maintenance of nitrogen metabolism under GA deprivation although plant growth was decreased. Currently, metabolic profiling studies on GA-deficient mutants in Arabidopsis are scarce. However, metabolic profiling was reported for poplar overexpressing the Arabidopsis proteins GAI and RGL1 with mutated or deleted DELLA domains, respectively (Busov et al., 2006). Overexpression of the modified DELLA proteins caused accumulation of bioactive GA<sub>1</sub> and GA<sub>4</sub>. Most of the metabolic alterations observed in the transgenic poplar leaves indicated a reduced flow of carbon through the lignin biosynthesis pathway and changes in the allocation of secondary phenolic metabolites (Busov et al., 2006). In Arabidopsis, overexpression of GA20ox1 causes an enlargement of leaf size (Gonzalez et al., 2010) similar to that caused by the exogenous application of bioactive GA, but opposite to PAC treatment. As expected, the metabolite profiles of GA20ox1 overexpressors overlap only partly with those obtained here for PAC-treated plants.

In summary, metabolite analyses revealed that a low GA level mainly affects growth by uncoupling it from carbon availability. Given that under a good GA level tight relationships linking carbon availability and growth are observed, strategies to identify genes which orchestrate these relationships will probably present promising ways by which to identify new markers for growth potential.

# Supplementary data

Supplementary data are available at JXB online.

Figure S1. Phenotypic changes of Arabidopsis wild-type plants caused by PAC and/or GA<sub>3</sub> treatment.

Figure S2. MapMan representations of differentially expressed genes.

Figure S3. Effect of treatment with PAC and/or GA<sub>3</sub> on shoot biomass of Arabidopsis plants (27 d after sowing) and fumarate levels.

**Table S1.** Genes differentially expressed in GA-treated versus non-treated control plants and in PAC-treated versus non-treated controls.

Table S2. Expression of GA-related genes in plants treated with PAC and/or GA<sub>3</sub>.

Table S3. Relative metabolite content of rosettes of plants treated with PAC and/or GA<sub>3</sub>.

Table S4. Paclobutrazol increases protein and chlorophyll concentration in fully expanded leaves.

Table S5. Sequence of primers used for qRT-PCR.

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