

# Global Analysis of DELLA Direct Targets in Early Gibberellin Signaling in *Arabidopsis* <sup>W</sup>

Rodolfo Zentella,<sup>a,1</sup> Zhong-Lin Zhang,<sup>a,1</sup> Mehea Park,<sup>a,2</sup> Stephen G. Thomas,<sup>a,3</sup> Akira Endo,<sup>b</sup> Kohji Murase,<sup>a</sup> Christine M. Fleet,<sup>a,4</sup> Yusuke Jikumaru,<sup>b</sup> Eiji Nambara,<sup>b</sup> Yuji Kamiya,<sup>b</sup> and Tai-ping Sun<sup>a,5</sup>

<sup>a</sup>Department of Biology, Duke University, Durham, North Carolina 27708

<sup>b</sup>RIKEN Plant Science Center, Tsurumi-ku, Yokohama, 230-0045, Japan

**Bioactive gibberellins (GAs) are phytohormones that regulate growth and development throughout the life cycle of plants. DELLA proteins are conserved growth repressors that modulate all aspects of GA responses. These GA-signaling repressors are nuclear localized and likely function as transcriptional regulators. Recent studies demonstrated that GA, upon binding to its receptor, derepresses its signaling pathway by binding directly to DELLA proteins and targeting them for rapid degradation via the ubiquitin-proteasome pathway. Therefore, elucidating the signaling events immediately downstream of DELLA is key to our understanding of how GA controls plant development. Two sets of microarray studies followed by quantitative RT-PCR analysis allowed us to identify 14 early GA-responsive genes that are also early DELLA-responsive in *Arabidopsis thaliana* seedlings. Chromatin immunoprecipitation provided evidence for in vivo association of DELLA with promoters of eight of these putative DELLA target genes. Expression of all 14 genes was downregulated by GA and upregulated by DELLA. Our study reveals that DELLA proteins play two important roles in GA signaling: (1) they help establish GA homeostasis by direct feedback regulation on the expression of GA biosynthetic and GA receptor genes, and (2) they promote the expression of downstream negative components that are putative transcription factors/regulators or ubiquitin E2/E3 enzymes. In addition, one of the putative DELLA targets, *XERICO*, promotes accumulation of abscisic acid (ABA) that antagonizes GA effects. Therefore, DELLA may restrict GA-promoted processes by modulating both GA and ABA pathways.**

## INTRODUCTION

Bioactive gibberellins (GAs) control a wide range of processes during plant development, including seed germination, leaf expansion, stem and root elongation, flowering time, and flower and fruit development (Davies, 2004; Fleet and Sun, 2005; Swain and Singh, 2005). Genetic and molecular studies have identified the GA receptors and several positive and negative components in the GA signaling cascade (Sun and Gubler, 2004; Hartweck and Olszewski, 2006). Among them, three major players are the GA receptors, the DELLA repressor proteins, and the F-box proteins that control the stability of DELLA proteins. Elegant work by Ueguchi-Tanaka et al. (2005) demonstrated that GA-INSENSITIVE DWARF1 (GID1) is a soluble GA receptor in rice (*Oryza sativa*). Subsequently, the GID1 homologs (GID1a, GID1b, and GID1c) in *Arabidopsis thaliana* were identified (Nakajima

et al., 2006). Null mutations in the single *GID1* gene in rice or in all three genes in *Arabidopsis* lead to an extremely dwarf and GA-insensitive plant (Ueguchi-Tanaka et al., 2005; Griffiths et al., 2006; Willige et al., 2007).

The DELLA proteins are conserved repressors of GA signaling that act immediately downstream of the GA receptor to modulate all aspects of GA-induced growth and development in plants (Thomas and Sun, 2004; Griffiths et al., 2006; Nakajima et al., 2006). Recent studies further suggest that DELLA proteins may also restrict plant growth by integrating signals from other hormone pathways and environmental cues (Achard et al., 2003, 2006; Fu and Harberd, 2003). There are five members of the *DELLA* gene family in *Arabidopsis*: *REPRESSOR OF ga1-3 (RGA)*, *GA-INSENSITIVE (GAI)*, *RGA-LIKE1 (RGL1)*, *RGL2*, and *RGL3*. Characterization of mutant combinations of null alleles in each *DELLA* gene demonstrates the overlapping and distinct functions of these genes in plant development. GA-induced vegetative growth and floral initiation are repressed by *RGA* and *GAI* (Dill and Sun, 2001; King et al., 2001). GA-promoted seed germination is mainly regulated by *RGL2*, although the remaining *DELLA* genes also play a minor role (Lee et al., 2002; Wen and Chang, 2002; Tyler et al., 2004; Cao et al., 2005; Tyler, 2006). In addition, *RGA*, *RGL1*, and *RGL2* are involved in flower and fruit development (Cheng et al., 2004; Tyler et al., 2004; Yu et al., 2004). Structurally, DELLAs are a subgroup of proteins that belong to the GRAS (for *GAI*, *RGA*, *SCARECROW*) family of transcriptional regulators and share a conserved C-terminal GRAS domain (Pysh et al., 1999; Bolle, 2004). DELLA proteins are named after a conserved motif at their N termini, absent in other

<sup>1</sup> These authors contributed equally to this work.

<sup>2</sup> Current address: Department of Molecular Genetics and Cell Biology, University of Chicago, Chicago, IL 60637.

<sup>3</sup> Current address: Rothamsted Research, Harpenden, Hertfordshire, AL5 2JQ UK.

<sup>4</sup> Current address: Department of Biology, Emory and Henry College, Emory, VA 24327.

<sup>5</sup> Address correspondence to tps@duke.edu.

The author responsible for distribution of materials integral to the findings presented in this article in accordance with the policy described in the Instructions for Authors (www.plantcell.org) is: Tai-ping Sun (tps@duke.edu).

<sup>W</sup> Online version contains Web-only data.

www.plantcell.org/cgi/doi/10.1105/tpc.107.054999

GRAS members (Silverstone et al., 1998; Peng et al., 1999; Pysh et al., 1999). All DELLA proteins also contain a polymeric Ser and Thr region that may include sites of phosphorylation or glycosylation, Leu heptad repeats that may mediate protein–protein interactions, and putative nuclear localization signals. DELLA-green fluorescent protein (GFP) fusion proteins are nuclear localized when expressed in transgenic plants (Silverstone et al., 2001; Gubler et al., 2002; Itoh et al., 2002). Although DELLAs do not have a clearly identified DNA binding domain, they may act as coactivators or repressors by interacting with other transcription factors. In support of this idea, two other GRAS proteins, SHORT-ROOT (SHR) and SCARECROW (SCR) have been shown recently to be associated with the promoter sequences of their target genes in vivo (Levesque et al., 2006; Cui et al., 2007).

Recent studies demonstrated that DELLA proteins are rapidly degraded in response to GA treatment and their N-terminal DELLA domain plays a vital regulatory role in DELLA protein stability (Sun and Gubler, 2004). Deletion or specific amino acid substitutions within the conserved DELLA motif (e.g. *gai-1* and *rga-Δ17* in *Arabidopsis*) stabilize mutant DELLA proteins and confer GA-insensitive dwarf phenotypes (Peng et al., 1997; Dill et al., 2001; Gubler et al., 2002; Itoh et al., 2002).

The F-box proteins SLEEPY1 (SLY1) in *Arabidopsis* and GID2 in rice are part of the SCF<sup>SLY1</sup> and SCF<sup>GID2</sup> E3 ubiquitin ligase complexes, respectively (McGinnis et al., 2003; Sasaki et al., 2003). Both *sly1* and *gid2* mutants display GA-unresponsive dwarf phenotypes and accumulate extremely high levels of DELLA proteins. Yeast two-hybrid and pull-down assays showed that both SLY1 and GID2 interact directly with DELLA proteins, indicating that SLY1 and GID2 recruit DELLA proteins for degradation by the 26S proteasome (Sasaki et al., 2003; Dill et al., 2004; Fu et al., 2004; Gomi et al., 2004). With the recent discovery of the GA receptor, the mechanism involved in GA-induced DELLA proteolysis has been elucidated further. It appears that GA promotes interaction of its receptor GID1 with the DELLA proteins via their DELLA domains (Ueguchi-Tanaka et al., 2005; Griffiths et al., 2006; Nakajima et al., 2006; Willige et al., 2007). This in turn may cause a conformational change in the DELLA protein that facilitates the F-box protein recognition (Griffiths et al., 2006), resulting in rapid degradation of DELLA through the ubiquitin-proteasome pathway.

Mutant studies also identified *SPINDLY* (SPY) as another GA signaling repressor in *Arabidopsis* (Jacobsen and Olszewski, 1993). SPY and its homologs in other species share high sequence similarity to the O-linked N-acetylglucosamine transferases (OGTs) in mammals (Thornton et al., 1999; Roos and Hanover, 2000). The animal OGTs modify target proteins by glycosylation of Ser/Thr residues, which either interfere or compete with kinases for phosphorylation sites (Wells et al., 2001). The recombinant SPY protein exhibits OGT activity in vitro (Thornton et al., 1999), and like animal OGTs, SPY is localized to the cytoplasm and nucleus (Swain et al., 2002). Target proteins of SPY have not been identified, but potential candidates include the DELLA proteins (Olszewski et al., 2002; Shimada et al., 2006; Silverstone et al., 2007).

Further downstream in the GA signaling pathway, the transcription factor GAMYB induces transcription of  $\alpha$ -amylase genes in the barley (*Hordeum vulgare*) aleurone (Gubler et al., 1995, 1999). GAMYB acts downstream of DELLA, although it is unlikely a direct target of DELLA because of a 1-h lag time between GA-

dependent DELLA protein degradation and GAMYB mRNA induction (Gubler et al., 2002). Mutant and transgenic studies in barley, rice, and *Arabidopsis* indicate that GAMYB also modulates GA-regulated floral development (Murray et al., 2003; Kaneko et al., 2004; Millar and Gubler, 2005). Moreover, one of the *Arabidopsis* GAMYBs (MYB33) is likely to play a role in GA-mediated floral induction by activating expression of *LEAFY* (Blazquez and Weigel, 2000; Gocal et al., 2001; Millar and Gubler, 2005). In *Arabidopsis*, GA also induces trichome initiation by activating *GLABROUS1*, another MYB gene (Perazza et al., 1998).

Although the earliest events of GA signaling, from GA perception to DELLA degradation, are now better understood, the gene regulatory network directly downstream of DELLA is unclear. Several microarray experiments have examined the effects of GA and DELLA proteins on gene expression in germinating seeds, seedlings, and flowers in *Arabidopsis* (Ogawa et al., 2003; Cao et al., 2006; Nemhauser et al., 2006). These experiments have taken advantage of mutants with defects in the GA biosynthetic pathway and/or in DELLA genes. The null mutant *ga1-3* is severely GA deficient because *GA1* encodes *ent-copalyl* diphosphate synthase, which catalyzes the first committed step in GA biosynthesis (Sun and Kamiya, 1994). This mutant is an extreme dwarf and has delayed flowering and male sterility (Koornneef and van der Veen, 1980; Wilson et al., 1992). These defects can be completely rescued by exogenous application of GA. The work by Ogawa et al. (2003) identified GA-regulated genes during germination by analyzing *ga1-3* seeds after GA treatment for 3 to 12 h. GA-induced genes include those that are involved in cell wall metabolism (for cell elongation) and cell division. Genes that function in other plant hormone pathways were also modulated by GA (Ogawa et al., 2003). Cao et al. (2006) monitored differential gene expression in imbibed seeds and developing flowers of the wild type, *ga1-3*, and a quintuple null mutant *ga1 rga gai rgl1 rgl2*. Their study uncovered a large number of GA-regulated genes, ~50% of which were also DELLA dependent (Cao et al., 2006). However, this analysis could not distinguish early DELLA targets from those that are located further downstream in the GA response pathway. Nemhauser et al. (2006) compared the initial responses of *Arabidopsis* seedlings to GA and six additional plant hormones by analyzing publicly available microarray data from the AtGenExpress consortium ([http://www.arabidopsis.org/servlets/TairObject?type=expression\\_set&id=1007966175](http://www.arabidopsis.org/servlets/TairObject?type=expression_set&id=1007966175)). Interestingly, within a 3-h treatment period, genes involved in each hormone signaling pathway are largely specific, although each hormone appears to alter expression of other hormone metabolism genes (Nemhauser et al., 2006).

Because DELLA proteins play a central role in modulating GA responses in plants, elucidating the molecular events immediately downstream of DELLA should shed light on how GA controls plant development. In this study, we identified early GA-responsive genes and specifically those directly controlled by DELLA proteins in shoots of *Arabidopsis* seedlings by microarray analysis. Our results indicate that DELLA proteins participate in two aspects of the GA signaling network: they help establish GA homeostasis by feedback regulating the expression of GA biosynthetic genes and GA receptors, and they promote the expression of downstream regulatory proteins that are putative negative components in GA signaling. In addition, DELLA may mediate interaction between GA and ABA pathways by upregulating

expression of a putative E3 ligase gene, *XERICO*, which in turn promotes ABA accumulation.

## RESULTS

DELLA proteins are conserved GA signaling repressors that act immediately downstream of the GA receptors (GID1) and play a pivotal role in modulating all aspects of GA responses (Thomas and Sun, 2004; Ueguchi-Tanaka et al., 2005). These growth repressors may function as transcriptional regulators, although their direct targets have not been identified. To uncover early GA-responsive genes and those controlled by DELLA, we carried out two sets of microarray experiments. The first set aimed to identify early GA-regulated genes by analyzing the global gene expression profile in the GA-deficient mutant *ga1-3* in the presence or absence of GA treatment; the second set aimed to uncover DELLA-regulated genes by induced expression of a dominant DELLA mutant protein (*rga-Δ17*) using a glucocorticoid (dexamethasone [DEX])-inducible system. All microarray data were generated using shoots of 8-d-old seedlings. The next two sections describe our initial experiments for determining the optimal GA treatment time point and for generating the reagents for the DEX-inducible *rga-Δ17* system.

### Early GA Responses Were Observed within Minutes of GA Treatment

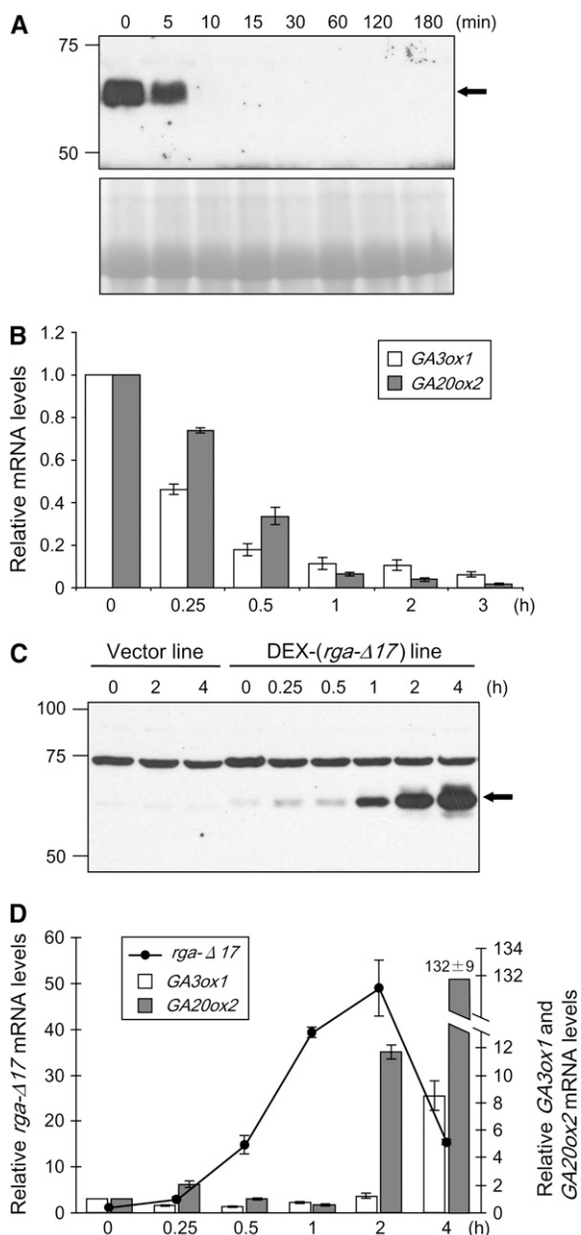
In the wild-type background, responses to exogenous GA may be attenuated because of preexisting levels of endogenous GA. To maximize changes in gene expression between water versus GA treatment, the severely GA-deficient mutant *ga1-3* was used. To determine the appropriate time point(s) of GA treatment for the microarray experiments, we first examined the rate of RGA protein disappearance and changes in transcript levels of two known GA-downregulated genes (*GA3ox1* and *GA20ox2*) (Chiang et al., 1995; Phillips et al., 1995). In the *ga1-3* mutant, RGA was rapidly degraded upon addition of 2 μM GA<sub>4</sub> (Figure 1A). After 5 min of GA treatment, the amount of RGA was reduced to ~50%. At 10 min, ~90% of the RGA protein disappeared, and by 30 min it was no longer detectable (Figure 1A; see Supplemental Figure 1A online). *GA3ox* and *GA20ox* enzymes catalyze the last and penultimate steps for the synthesis of bioactive GAs in *Arabidopsis*, respectively (Hedden and Phillips, 2000). Transcript levels of *GA3ox1* and *GA20ox2* were shown to be under feedback regulation by GA treatment and by activity in the GA response pathway (Hedden and Phillips, 2000; Sun and Gubler, 2004). In the *ga1-3* mutant, the amounts of *GA3ox1* and *GA20ox2* transcripts rapidly declined upon GA application, and changes were noticeable as early as 15 min after hormone treatment (Figure 1B). Our results showed that GA-induced RGA degradation precedes GA downregulation of *GA3ox1* and *GA20ox2* mRNA levels. Based on these results, we decided to perform microarray analysis using the *ga1-3* samples that were treated with 2 μM GA<sub>4</sub> or water for 1 h.

### An Inducible *rga-Δ17* Protein Accumulated to High Levels and Caused Altered Expression of GA-Regulated Genes

Among the five DELLA proteins in *Arabidopsis*, RGA and GAI are the major regulators of GA-mediated vegetative growth (Dill and

Sun, 2001; King et al., 2001). To identify direct targets of DELLA proteins in vegetative phase, we generated transgenic *Arabidopsis* carrying an inducible *rga-Δ17* transgene using the DEX-mediated transcriptional induction system (Aoyama and Chua, 1997). The *rga-Δ17* protein, which contains a DELLA motif deletion, is resistant to GA-induced degradation and therefore confers an extreme dwarf phenotype when expressed using the *RGA* promoter (Dill et al., 2001). The use of the DEX-inducible system allowed us to generate transgenic plants that grew normally in the absence of DEX treatment. The *rga-Δ17*-inducible construct contains a chimeric transcription factor (GVG), which is constitutively expressed under the cauliflower mosaic virus 35S promoter (*CaMV35S*) and six copies of the *GAL4* upstream activating sequence fused to the *rga-Δ17* coding sequence. GVG consists of the DNA binding domain of the yeast transcription factor *GAL4*, the transactivating domain of the herpes viral protein VP16, and the hormone binding domain of the rat glucocorticoid receptor. In plant cells, GVG exists as a cytoplasmic complex with the 90-kD heat shock protein (HSP90). Application of DEX results in dissociation of the GVG-HSP90 complex and rapid movement of GVG into the nucleus, where it activates transcription of the 6xUAS-driven transgene (*rga-Δ17* in our experiment).

We identified multiple homozygous transgenic lines that contain the vector (control) or the DEX-inducible *rga-Δ17* construct [DEX-(*rga-Δ17*)] in the T3 generation. For the DEX-(*rga-Δ17*) lines, we then screened for those that had low basal levels of transgene expression and demonstrated rapid and high induction levels of the *rga-Δ17* transcript and protein following DEX treatment. We also analyzed GVG transcript levels in the vector control and the DEX-(*rga-Δ17*) lines and identified those with similar levels of GVG expression to minimize artifacts due to differential GVG levels (Kang et al., 1999). To determine the appropriate time points for microarray experiments, a DEX induction time course was carried out. Eight-day-old seedlings, pretreated for 16 h with 2 μM GA<sub>4</sub> to saturate the GA responses in the plants, were then treated with 10 μM DEX plus 2 μM GA<sub>4</sub> or only GA. Application of DEX caused rapid accumulation of *rga-Δ17* protein in the DEX-(*rga-Δ17*) line but not in a vector control line (Figure 1C). By 1 h of DEX treatment, the *rga-Δ17* protein was readily detectable and it continued to accumulate for up to 4 h, reaching ~16 times the level of RGA protein present in *ga1-3* (see Supplemental Figure 1B online). Although a small amount of *rga-Δ17* protein was detected in the DEX-(*rga-Δ17*) line before DEX treatment (time 0 h; Figure 1C), this transgenic line did not show any growth defects without DEX induction (data not shown). Transcript levels of *GA3ox1* and *GA20ox2* were also determined. Accumulation of *GA3ox1* transcript was only observed after 4 h of DEX induction. However, for *GA20ox2*, a >11-fold increase was observed after 2 h. By 4 h, the levels of *GA20ox2* mRNA were >100-fold higher than those in the uninduced sample. It is worth noting that induction of both GA biosynthetic genes was preceded by accumulation of the *rga-Δ17* transcript and protein (Figures 1C and 1D). In addition, the transcript levels of *GA3ox1* and *GA20ox2* were not induced in the vector control line (see Supplemental Figure 2A online). Based on these results, the 2- and 4-h time points were selected for microarray experiments.



**Figure 1.** Characterization of *Arabidopsis* Lines Used for Microarray Experiments.

**(A)** GA treatment triggers rapid degradation of endogenous RGA. Eight-day-old *ga1-3* seedlings were treated with 2  $\mu\text{M}$   $\text{GA}_4$  for 0 to 3 h. Proteins extracted from shoots were analyzed by immunoblotting using affinity-purified anti-RGA antibody. Ponceau staining was used to confirm equal loading. Arrow indicates the position of RGA. The amounts of RGA remaining after 5- to 30-min treatments were estimated by serial dilutions as shown in Supplemental Figure 1A online.

**(B)** GA application causes a decrease in mRNA levels of GA biosynthetic genes. Seedlings were grown and treated as in **(A)** except that total RNA was isolated at the indicated time points. Transcript levels were determined by qRT-PCR, and the level at time 0 h was set to 1.0. Bars indicate the average transcript level  $\pm$  SE of four replications from two independent experiments.

**(C)** DEX treatment strongly induces the accumulation of *rga-Δ17* in a

## Identification of Early GA-Responsive and DELLA Target Genes by Microarray Analysis

Using the conditions defined in the previous two sections, we produced two sets of data using Affymetrix ATH1 GeneChips: the first one allowed us to identify early GA-responsive genes, whereas the second one aimed to uncover early DELLA-responsive genes. Our assumption was that among the DELLA targets that function in the GA response pathway, DELLA-induced genes would be downregulated by GA treatment, and DELLA-repressed genes would be upregulated by GA.

## Early GA-Responsive Gene Data Set

For the first data set, we used 8-d-old shoots of *ga1-3* seedlings treated for 1 h with either water or 2  $\mu\text{M}$   $\text{GA}_4$ . Four biological replicas were performed. The CEL files that contained the raw hybridization signal values of each microarray were analyzed using GeneSpring 7.2. The data were normalized using the GC-RMA (robust multiarray analysis with correction for GC content of the oligonucleotide) algorithm. The statistical analysis consisted of a one-way analysis of variance (ANOVA), using a cutoff value of  $P \leq 0.01$  because a more stringent filtering procedure (e.g., false discovery rate [FDR]  $< 0.05$ ; Benjamini and Hochberg, 1995) did not recover any GA-responsive genes. The resulting list included 81 genes (see Supplemental Table 1 online). When a 2 fold change (FC) in gene expression was used to filter our gene list, only 20 genes were recovered, all of them downregulated. When the threshold was lowered to 1.5 FC, 42 genes were obtained, 33 of them being downregulated and nine upregulated by GA. Our final list of 67 GA-responsive genes contained all genes that had at least 1.2 FC with respect to the water-treated control (Table 1). Among these, 45 were downregulated by GA and 22 induced. As expected, the two GA biosynthetic genes, *GA3ox1* and *GA20ox2*, were present in this list. Recently, several GA-related data sets using the ATH1 arrays have become available. One of them, generated by the AtGenExpress consortium ([http://www.arabidopsis.org/servlets/TairObject?type=expression\\_set&id=1007966175](http://www.arabidopsis.org/servlets/TairObject?type=expression_set&id=1007966175)), examined the GA response of Columbia (Col-0; wild type) and *ga1-5* mutant seedlings growing in liquid culture for 7 d and treated with 1  $\mu\text{M}$   $\text{GA}_3$  for 0.5, 1, and 3 h. Another data set contains gene expression profiles of Landsberg *erecta* (*Ler*), *ga1-3*, and the *ga1-3 rga gai rgl1 rgl2* quintuple mutant of

DEX-inducible *rga-Δ17* line. Eight-day-old seedlings were pretreated for 16 h with 2  $\mu\text{M}$   $\text{GA}_4$  followed by treatment for 0 to 4 h with either 2  $\mu\text{M}$   $\text{GA}_4$  only or in combination with 10  $\mu\text{M}$  DEX. Shoot proteins were extracted and analyzed by immunoblotting using crude anti-RGA antibody. The 75-kD nonspecific band serves as evidence for equal loading. Arrow indicates the position of *rga-Δ17*.

**(D)** DEX treatment caused rapid accumulation of *rga-Δ17* transcript, which precedes induction of *GA3ox1* and *GA20ox2*. Seedlings were pretreated and treated as in **(C)**. Total RNA from shoots was collected at the indicated time points. Transcript levels were determined by qRT-PCR, and the level at time 0 h was set to 1.0. Bars are the means  $\pm$  SE of three replicates. In **(B)** and **(D)**, the housekeeping gene *GAPC*, whose expression is not responsive to GA (Dill et al., 2004), was used to normalize different samples.

**Table 1.** GA-Responsive Genes and Overlaps with Other Microarray Data Sets

AGI Locus	Description	<i>ga1-3</i>						Cao et al. (2006)				Nemhauser et al. (2006) <sup>a</sup>					
		DEX-( <i>rgaΔ17</i> )			AtGenExpress**			(Flowers)		(Seeds)		0.5 h	1 h	3 h			
		1 h	2 h	4 h	0.5 h	1 h	3 h	DELLA-Dep. <sup>b</sup>	DELLA-Ind. <sup>b</sup>	DELLA-Dep. <sup>b</sup>	DELLA Ind. <sup>b</sup>						
1	At2g45900	Expressed protein (Exp-PT1)*	-13.5	31.5	31.8	-1.3	-4.3	-15.6	+								
2	At1g15550	GA3ox1*	-10.9	1.0	2.4	-1.5	-4.8	-6.8	+								
3	At5g51810	GA20ox2*	-10.6	4.3	45.4	-1.4	-5.3	-10.7	+								
4	At1g50420	SCL3*	-6.8	4.0	6.4	-1.4	-2.8	-3.4	+								
5	At4g19700	RING-E3 HCa type (RING)*	-5.5	1.7	1.9	-2.8	-4.0	-4.3	+								
6	At3g63010	GID1b*	-5.1	2.9	4.3	-1.5	-2.6	-2.5	+								
7	At3g05120	GID1a*	-4.1	1.3	1.7		-1.8	-1.9	+								
8	At4g36410	UBC17*	-4.1	12.8	17.0				+								
9	At2g04240	XERICO*	-3.7	1.9	1.7	-1.5	-2.4	-2.4	+								
10	At1g54120	Expressed protein	-3.4				-1.4	-1.4	+								
11	At1g56650	MYB75	-3.1								+						
12	At5g19340	Expressed protein	-2.9				-1.3	-1.5									
13	At4g23060	IQD22	-2.8			-1.3	-1.9	-2.0									
14	At1g17830	Expressed protein	-2.3						+								
15	At5g18840	Sugar transporter, putative similar to ERD6	-2.2						+								
16	At5g03670	Expressed protein	-2.2														
17	At1g29270	Expressed protein	-2.2								+						
18	At3g30180	BR6ox2	-2.2														
19	At1g68570	H <sup>+</sup> -dependent oligopeptide transport family protein	-2.1														
20	At5g67480	BT4 (BTB/TAZ domain protein 4)*	-2.0	1.7	1.4	-1.2	-1.6	-1.9									
21	At5g47550	Cys protease inhibitor, putative/cystatin	-2.0														
22	At1g21250	WAK1 (Wall-Associated Kinase1)	-1.9						+								
23	At5g05180	Expressed protein	-1.9														
24	At4g27730	Oligopeptide transporter family protein	-1.8														
25	At4g36220	FAH1 (Ferulate-5-hydroxylase1)	-1.8						+								
26	At3g54320	WRI1 (WRINKLED1)	-1.7								+						
27	At2g41180	SigA binding protein-related	-1.7						+								
28	At2g02080	Zinc finger (C2H2 type) family protein	-1.7														
29	At1g76990	ACT domain-containing protein	-1.7				-1.3	-1.3									
30	At1g80870	Protein kinase family protein	-1.6														
31	At3g61460	BRH1 (BR-responsive RING-H2)	-1.6														
32	At4g27300	S-locus protein kinase, putative	-1.6						+								
33	At1g69160	Expressed protein	-1.5														
34	At3g47160	Expressed protein	-1.5														
35	At3g19850	Phototropic-responsive NPH3 family protein	-1.5														
36	At3g02910	Expressed protein	-1.5				-1.5	-1.6	+								
37	At3g52870	CaM binding family protein (CaM-BP)*	-1.5	-1.8	-1.6												
38	At3g11280	MYB-like protein*	-1.4	2.4	2.6				+								
39	At4g38580	Copper chaperone-related	-1.4														
40	At3g12670	CTP-synthase/UTP-ammonia ligase, putative	-1.4														
41	At4g31590	Glycosyl transferase family 2 protein	-1.4														
42	At2g34340	Expressed protein (Exp-PT2)*	-1.4	1.3	3.7												
43	At2g31730	bHLH154*.c	-1.4	2.0	2.0												
44	At4g39630	Expressed protein	-1.3														
45	At2g33310	IAA13	-1.3														
46	At4g28220	NADH dehydrogenase-related	1.2														
47	At5g23290	c-MYC binding protein, putative/prefoldin	1.2														
48	At2g21185	Expressed protein	1.3														
49	At1g55190	Prenylated rab acceptor (PRA1) family protein	1.3														
50	At5g04420	No apical meristem (NAM) family protein	1.3														
51	At5g05250	Expressed protein	1.3														
52	At5g22310	Expressed protein	1.3														

(Continued)

**Table 1.** (continued).

AGI Locus	Description	<i>ga1-3</i>	DEX-( <i>rga-Δ17</i> )						Cao et al. (2006)				Nemhauser et al. (2006) <sup>a</sup>				
			DEX-( <i>rga-Δ17</i> )		AtGenExpress**		(Flowers)		(Seeds)								
			1 h	2 h	4 h	0.5 h	1 h	3 h	DELLA-Dep. <sup>b</sup>	DELLA-Ind. <sup>b</sup>	DELLA-Dep. <sup>b</sup>	DELLA Ind. <sup>b</sup>	0.5 h	1 h	3 h		
53	At5g12980	RCD1-like cell differentiation protein, putative	1.4														
54	At1g14440	Zinc finger homeobox family protein	1.4														
55	At3g01470	HB-1/HAT5	1.4														
56	At5g60970	TCP family transcription factor, putative	1.4														
57	At5g16590	LRR transmembrane protein kinase, putative	1.4														
58	At3g07010	Pectate lyase family protein	1.5														
59	At2g38090	MYB family transcription factor	1.5														
60	At5g03555	Permease, nucleotide, allantoin family protein	1.6														
61	At3g60520	Expressed protein	1.6														
62	At4g30850	Expressed protein	1.6														
63	At5g08130	bHLH046 <sup>c</sup>	1.6														
64	At2g19310	Expressed protein	1.6														
65	At1g54050	17.4-kD class III heat shock protein	1.7														
66	At2g41940	Zinc finger (C2H2 type) family protein	1.7														
67	AT3g50750	BZR1-like BR-signaling positive regulator-related	1.9														

Single asterisks indicate genes that overlap between the *ga1-3* and DEX-(*rga-Δ17*) microarray data sets, which are also listed in Table 2. Double asterisks indicate raw data from the AtGenExpress consortium analyzed in this work. The + indicates that the gene was identified by Cao et al. (2006) as a GA-responsive gene (either DELLA-dep. or DELLA-ind.). AGI, Arabidopsis Genome Initiative.

<sup>a</sup> AtGenExpress data analyzed by Nemhauser et al. (2006).

<sup>b</sup> DELLA-dep., DELLA-dependent genes; DELLA-ind., DELLA-independent or partially dependent genes.

<sup>c</sup> Names according to Bailey et al. (2003).

germinating seeds and developing flowers (Cao et al., 2006). To compare our data with those generated by AtGenExpress, we downloaded the Affymetrix *CEL* files and analyzed them using the same procedure and cutoffs as our own data set ( $FC \geq 1.2$  and  $P \leq 0.01$ ). The wild-type and *ga1-5* data sets were analyzed together, and a list of 173 genes was obtained (see Supplemental Table 2 online). When we compared this list with our *ga1-3* list of 67 genes, only 14 genes overlapped (~21%), all of them GA repressed (Table 1). Similarly, 28 to 32% of our *ga1-3* list overlapped with GA-responsive genes in seeds and flowers, identified by Cao et al. (2006). The differences among these gene lists could be caused by genetic backgrounds (extreme GA-deficient *ga1-3* versus the wild type and *ga1-5* leaky mutant, *Ler* versus *Col*), growth conditions (agar plates versus liquid culture), tissues (shoots versus whole seedlings, seeds or flowers), or GA treatments ( $2 \mu\text{M GA}_4$  versus  $1 \mu\text{M GA}_3$ ). In addition, after 1- to 3-h GA treatments, the changes in transcript levels for most GA-responsive genes were very subtle (with  $FC \leq 2$ ). Therefore expression of these genes would be extremely sensitive to experimental conditions. In fact, none of the GA-responsive genes in our *ga1-3* list or in the AtGenExpress data set could pass a more stringent filtering method (Nemhauser et al., 2006; this work).

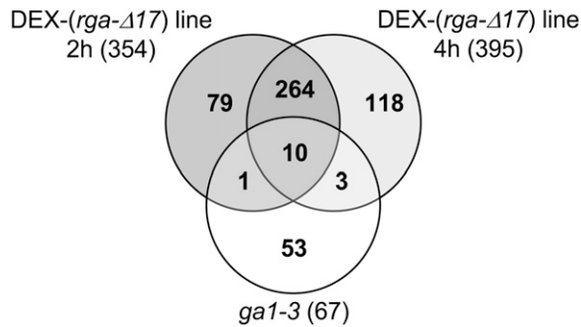
### Early DELLA-Responsive Gene Data Set

A second set of microarray data was generated using 8-d-old shoots of the DEX-(*rga-Δ17*) and vector control transgenic lines

that were pretreated for 16 h with  $2 \mu\text{M GA}_4$  and then exposed to  $2 \mu\text{M GA}_4 \pm 10 \mu\text{M DEX}$  for 2 and 4 h. Three biological replicates were performed for the DEX-(*rga-Δ17*) line and two replicates for the vector control line at 4 h.

The array data were filtered through several rounds of analyses. The *CEL* files from the Affymetrix output were analyzed with GeneSpring 7.2 as described above. To identify DELLA-responsive genes after DEX treatment, a two-way ANOVA statistical analysis was performed to compare gene expression in the DEX-(*rga-Δ17*) line  $\pm$  DEX (for 2 or 4 h). A list of 666 DELLA-responsive genes was generated when filtered by  $FDR \leq 0.01$  and  $FC \geq 1.5$ . An  $FDR \leq 0.01$  indicates 1% or less false positives. This gene list was then filtered by subtracting genes whose expression was affected by DEX treatment in the vector control line at 4 h (see Supplemental Table 3 online) to exclude those genes mainly affected by the *GVG* transgene. Because the vector control line accumulated an approximately three fold higher *GVG* transcript than the DEX-(*rga-Δ17*) line (see Supplemental Figure 2B online), genes with a  $P \leq 0.01$  and a  $FC \leq 3$  in the vector line were eliminated from the DELLA-responsive gene list. In addition, the *RGA* gene was removed from this list. As a result, 475 genes were listed as putative RGA-regulated, 336 being upregulated and 139 downregulated (see Supplemental Table 4 online).

To identify putative DELLA target genes that function in the GA response pathway, we compared the two microarray data sets and found only 14 overlapping genes (Figure 2, Table 2). All overlapping genes, except for At3g52870, were induced by *rga-Δ17*



**Figure 2.** Overlap between Microarray Data Sets of Early GA- and DELLA-Responsive Genes.

A total of 475 genes found to respond to *rga-Δ17* at 2 and/or 4 h after DEX induction ( $FDR \leq 0.01$ ;  $FC \geq 1.5$ ) are shown in the upper circles. The 67 GA-responsive genes in *ga1-3* after 1 h GA treatment ( $P \leq 0.01$ ;  $FC \geq 1.2$ ) are shown in the lower circle. Fourteen overlapping genes (listed in Table 2) were identified to be present in both GA- and DELLA-responsive gene lists.

and repressed by GA based on the microarray data (Table 2), suggesting that the direct downstream targets of RGA are mainly repressors of GA signaling. Among these 14 overlapping genes, nine of them were also found to be affected by the DELLA protein mutations in flowers and three in seeds (Cao et al., 2006) (Table 1).

In addition to these 14 genes, we found four more genes, *bHLH137*, *LBD40*, *WRKY27*, and *IQD22*, that exhibited upregulation in the DEX-(*rga-Δ17*) data set and GA downregulation in the *ga1-3* experiment. However, the P values in the *ga1-3* microarray data set for the first three genes were above 0.01. In the case of *IQD22*, its FC value in the vector control line was  $>3$ . Further quantitative RT-PCR (qRT-PCR) analysis corroborated their GA and RGA responses in *ga1-3* and in a transgenic line that expresses the *rga-Δ17* gene under the control of the endogenous RGA promoter,  $P_{RGA}:(rga-Δ17)$  (Dill et al., 2001), respectively (Table 2, Figure 3). These results indicated that these four additional genes are indeed early GA and RGA responsive. The transcript levels of gene *At3g52870* did not exhibit GA or RGA responses by qRT-PCR (Table 2); therefore, it was removed from the putative DELLA target gene list.

Among the early DELLA-induced genes, four of them encode either GA biosynthetic enzymes (*GA20ox2* and *GA3ox1*) or GA receptors (*GID1a* and *GID1b*). Transcript levels of these genes are known to be reduced by GA treatment and by the loss-of-function *DELLA* mutations (Hedden and Phillips, 2000; Griffiths et al., 2006). Our microarray data suggested that DELLA proteins may directly regulate expression of these genes.

The physiological roles of the rest of the early DELLA-induced genes have not been reported previously. Three genes encode putative ubiquitin E2 conjugating enzyme (UBC17) or RING-type

**Table 2.** Putative DELLA Targets

AGI Locus <sup>a</sup>	Description <sup>a</sup>	Microarray Data		qRT-PCR Data <sup>b</sup>			ChIP-qPCR <sup>c</sup>			
		DEX- <i>(rga-Δ17)</i> 2 h DEX	4 h DEX	<i>ga1-3</i> 1 h GA	<i>ga1-3</i> 1 h GA	<i>ga1-3</i> 3 h GA	<i>PRGA:(rga-Δ17)</i> <sup>d</sup>	Fold Enrichment	P Value <sup>e</sup>	
1	<b>At5g51810</b>	<b>GA20ox2</b>	4.3	45.4	-10.6	-4.4 ± 0.3	-23.5 ± 3.5	6.5 ± 0.0	1.0 ± 0.1	0.521
2	<b>At1g15550</b>	<b>GA3ox1</b>	1.0	2.4	-10.9	-7.1 ± 0.6	-18.0 ± 3.6	6.3 ± 0.2	1.4 ± 0.3	0.287
3	<b>At3g05120</b>	<b>AtGID1a</b>	1.3	1.7	-4.2	-2.1 ± 0.4	-2.3 ± 0.1	2.0 ± 0.2	2.1 ± 0.4	0.043
4	<b>At3g63010</b>	<b>AtGID1b</b>	2.9	4.3	-5.1	-2.9 ± 0.2	-2.5 ± 0.6	3.1 ± 0.3	2.1 ± 0.2	0.019
5	<b>At3g11280</b>	<b>MYB</b>	2.4	2.6	-1.4	-1.3 ± 0.0	-2.1 ± 0.2	2.1 ± 0.0	2.7 ± 0.1	0.002
6	<b>At5g50915</b>	<b>bHLH137<sup>f,g</sup></b>	4.7	10.0	-1.6	-1.4 ± 0.1	-6.7 ± 1.8	3.1 ± 0.2	1.8 ± 0.4	0.087
7	<b>At2g31730</b>	<b>bHLH154<sup>g</sup></b>	2.0	2.0	-1.4	-1.3 ± 0.0	-2.2 ± 0.2	1.8 ± 0.2	2.1 ± 0.7	0.228
8	<b>At5g52830</b>	<b>WRKY27<sup>f</sup></b>	4.6	5.0	-1.6	-1.6 ± 0.1	-1.4 ± 0.1	2.0 ± 0.1	1.3 ± 0.1	0.058
9	<b>At1g50420</b>	<b>SCL3</b>	4.0	6.4	-6.8	-7.7 ± 0.4	-8.2 ± 1.4	2.8 ± 0.4	3.5 ± 0.2	<0.0001
10	<b>At1g67100</b>	<b>LBD40<sup>f</sup></b>	3.1	6.9	-2.2	-2.2 ± 0.7	-9.3 ± 0.9	3.9 ± 0.7	2.0 ± 0.1	0.011
11	<b>At4g23060</b>	<b>IQD22<sup>f</sup></b>	3.9	2.6	-2.8	-3.0 ± 0.2	-2.3 ± 0.4	2.6 ± 0.2	2.3 ± 0.8	0.152
12	At5g67480	BT4 <sup>h</sup>	1.7	1.4	-2.0	-2.2 ± 0.2	-2.2 ± 0.3	1.3 ± 0.4	ND	
13	<b>At2g45900</b>	<b>Exp-PT1</b>	31.5	31.8	-13.5	-7.9 ± 1.3	-10.8 ± 0.4	5.9 ± 0.8	1.9 ± 0.6	0.243
14	At2g34340	Exp-PT2 <sup>h</sup>	1.3	3.7	-1.4	-1.7 ± 0.2	-2.2 ± 0.2	1.2 ± 0.1	ND	
15	At4g36410	UBC17 <sup>h</sup>	12.8	17.0	-4.1	-1.8 ± 0.1	-4.1 ± 0.7	-1.4 ± 0.0	ND	
16	<b>At2g04240</b>	<b>XERICO</b>	1.9	1.7	-3.7	-2.4 ± 0.3	-3.0 ± 0.5	1.9 ± 0.1	1.3 ± 0.1	0.080
17	<b>At4g19700</b>	<b>RING</b>	1.7	1.9	-5.5	-2.4 ± 0.4	-2.8 ± 0.9	2.8 ± 0.1	1.4 ± 0.3	0.262
18	At3g52870	CaM-BP <sup>i</sup>	-1.8	-1.6	-1.5	-1.1 ± 0.0	1.0 ± 0.3	1.2 ± 0.0	ND	

<sup>a</sup> The genes in boldface are responsive to both GA and DELLA, determined by microarray and qRT-PCR analyses.

<sup>b</sup> Average values of three repeats ± SE.

<sup>c</sup> Means of three independent ChIP-qPCR ± SE. ND, not determined.

<sup>d</sup> FC between the *Ler/PRGA:(rga-Δ17)* transgenic line and wild-type *Ler*.

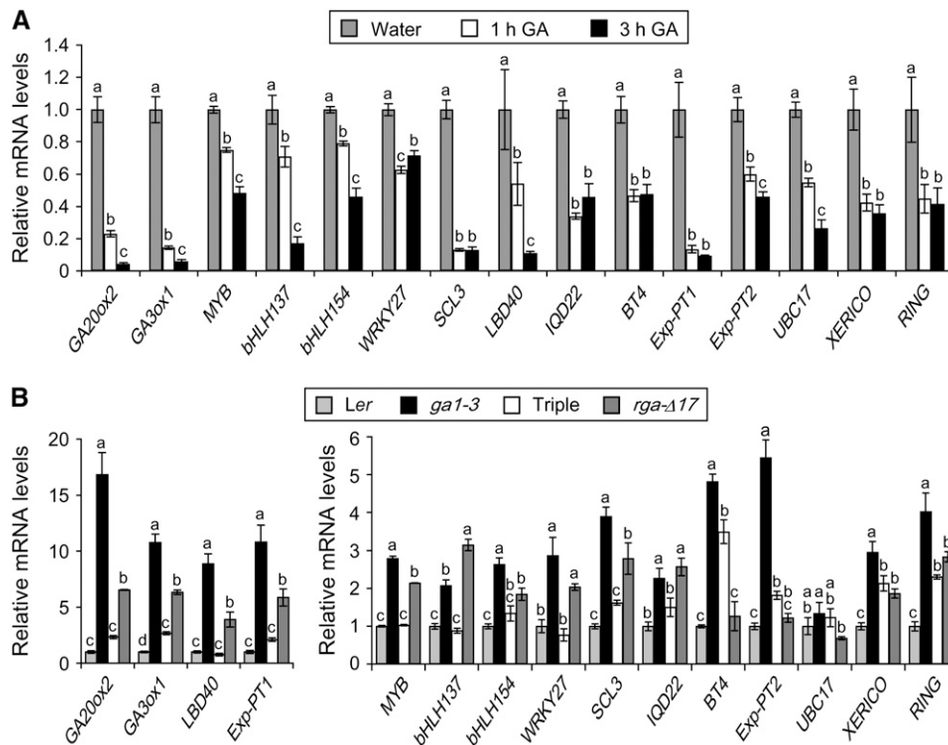
<sup>e</sup> A *t* test analysis was performed using the statistical package SAS 9.1.3.

<sup>f</sup> Genes that do not overlap between the *ga1-3* and the DEX-(*rga-Δ17*) microarray data sets but that were rescued after further testing by qRT-PCR (see text).

<sup>g</sup> Name according to Bailey et al. (2003).

<sup>h</sup> GA-responsive genes with weak response to DELLA, measured by qRT-PCR.

<sup>i</sup> This gene is considered a false positive because its expression, when measured by qRT-PCR, was not responsive to GA or DELLA.



**Figure 3.** Regulation of Transcript Levels of Putative RGA Target Genes by GA and DELLA.

**(A)** GA treatment downregulates mRNA levels of putative RGA targets in *ga1-3*. The means of three replicates of quantitative RT-PCR  $\pm$  SE are shown. Relative mRNA levels of individual genes after GA treatment were calculated in comparison to the water-treated control at each time point. Similar results were obtained when qRT-PCR was performed using two additional sets of biological replicates.

**(B)** Relative transcript levels of putative RGA target genes in the wild type, *ga1-3*, the triple homozygous mutant *rga-24 gai-t6 ga1-3*, and the transgenic line carrying *PRGA:(rga-Δ17)* (all lines are in the *Ler* background). The means of three replicates of qRT-PCR  $\pm$  SE are shown. The expression level in *Ler* was arbitrarily set to 1.0. Similar results were obtained when qRT-PCR was performed using a second set of samples.

In **(A)** and **(B)**, the housekeeping gene *GAPC*, whose expression is not responsive to GA (Dill et al., 2004), was used to normalize different samples. One-way ANOVA was performed with least significant difference multiple comparison tests at an  $\alpha$  level of 0.05 using SPSS version 11.5.0. When two samples show different letters (a to d) above the bars, the difference between them is significant ( $P < 0.05$ ).

ubiquitin E3 ligases (XERICO and At4g19700), which may ubiquitinate and modify activity of downstream components (Conaway et al., 2002) or regulate their stability via the proteasome pathway (Kraft et al., 2005; Stone et al., 2005). At4g19700 will be referred to as RING in the rest of this article. Several DELLA-responsive genes are predicted to encode nuclear transcription factors (*MYB*, *bHLH137*, *bHLH154*, and *WRKY27*) or transcriptional regulators (*SCL3*, *LBD40*, *IQD22*, and *BT4*). *MYB*, *bHLH*, and *WRKY* proteins belong to large families of transcription factors that regulate a myriad of processes during plant growth and development (Kranz et al., 1998; Eulgem et al., 2000; Toledo-Ortiz et al., 2003). *SCL3*, like the DELLA proteins, belongs to the GRAS family of putative transcriptional regulators (Pysh et al., 1999). *LBD* is a member of the ASYMMETRIC LEAVES2/LATERAL ORGAN BOUNDARIES domain (AS2/LBD) family (Iwakawa et al., 2002; Shuai et al., 2002). *IQD22* belongs to the IQD (IQ domain) family of calmodulin (CaM) binding proteins (Abel et al., 2005), which contain putative nuclear localization signals and may mediate  $Ca^{2+}$  signaling to regulate gene expression in the nucleus (Levy et al., 2005). *BT4* is a member of another family of nuclear  $Ca^{2+}$ /CaM binding proteins, which contain a BTB/POZ

domain at their N terminus and a zinc finger TAZ domain at the C terminus (Du and Poovaiah, 2004) and may function in transcriptional regulation (Kanai et al., 2000). At2g45900 and At2g34340, which will be referred to as Exp-PT1 and Exp-PT2, respectively, were annotated as expressed proteins and both are predicted to be localized to the nucleus (Nakai and Kanehisa, 1992; Heazlewood et al., 2005, 2007; Nair and Rost, 2005).

To corroborate our putative DELLA target gene list, we analyzed their transcript levels in GA time-course experiments and in *DELLA* mutant seedlings by qRT-PCR.

#### Putative RGA Targets Were GA and DELLA Responsive by qRT-PCR Analysis

We have previously shown, by qRT-PCR analysis, that the transcript levels of *GA3ox1*, *GID1a*, and *GID1b* are reduced by GA treatment and by the loss-of-function *DELLA* mutations (Griffiths et al., 2006). Similar results were also obtained in a microarray study that compared gene expression profiles in the wild type, *ga1-3*, and the *rga gai rgl1 rgl2 ga1* quintuple mutant (Cao et al., 2006). These observations are consistent with our microarray

data showing that these three genes may be direct targets of DELLA proteins. To verify our microarray results, we examined the transcript levels of the remaining 14 putative DELLA targets by qRT-PCR analysis. *GA3ox1* was also included in the analysis as a control. Consistent with our *ga1-3* microarray data, the transcript levels of all of these genes were downregulated by GA treatment (Figure 3A).

To confirm that these genes are DELLA responsive, we compared their transcript levels in *Ler*, *ga1-3*, the *rga-24 gai-t6 ga1-3* triple null mutant, and the  $P_{RGA}:(rga-\Delta 17)$  transgenic line by qRT-PCR. As expected, almost all (except *UBC17*) displayed elevated mRNA levels in *ga1-3* compared with *Ler* (Figure 3B, Table 2). Consistent with the microarray data indicating that they are DELLA-induced genes, transcript levels of these genes, except *BT4*, *Exp-PT2*, and *UBC17*, were increased by *rga-17* (comparing *Ler* versus *rga-17*). Moreover, their expression in the triple null mutant was lower than in *ga1-3* but similar to that in *Ler* (Figure 3B). These results support the idea that these genes are immediate targets of both RGA and GAI. Interestingly, the mRNA levels of *XERICO* and *RING* in the triple mutant remained higher than in *Ler*, suggesting that additional DELLA proteins may be also involved in controlling GA signaling through these two genes.

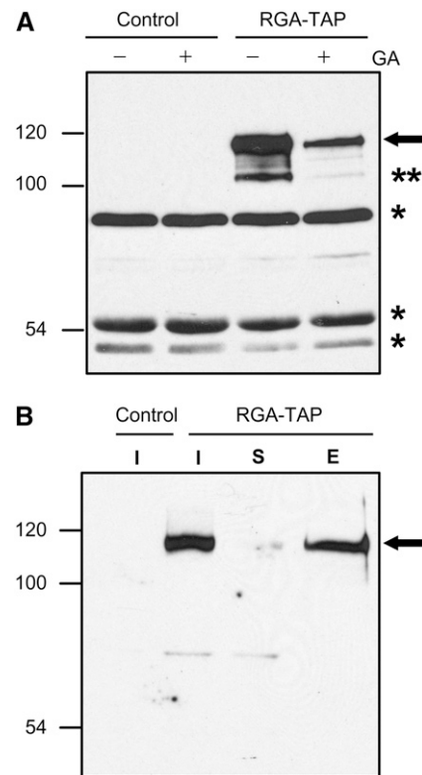
Our qRT-PCR analysis confirmed that 14 of the 17 RGA putative targets listed in Table 2 are both GA and DELLA responsive. *BT4*, *Exp-PT2*, and *UBC17* were clearly GA responsive, but their expression was not significantly affected in the  $P_{RGA}:(rga-\Delta 17)$  transgenic line or in the *rga gai ga1* background. Therefore, these genes were classified as only GA responsive but not DELLA responsive. In the rest of this article, we focus on the remaining 14 genes.

The putative DELLA targets may be coordinately regulated through common *cis*-elements in their promoters. Alternatively, different elements may be present in these genes if DELLA proteins interact with different transcription factors to regulate individual promoters. Promoter analysis of the 14 DELLA-responsive genes using the Web-based promoter analysis tool Athena (<http://www.bioinformatics2.wsu.edu/cgi-bin/Athena/cgi/home.pl>) (O'Connor et al., 2005) did not find any known transcription factor binding site to be significantly enriched. Using the MEME program (Bailey and Elkan, 1994; Bailey and Gribskov, 1998), a consensus sequence [C/T]T[C/T][C/A]TC[T/C][C/T]TCT[C/T][C/T]T[T/C] (named CCT element) with  $P < 7.2 \times 10^{-6}$  was found to be present within 1 kb 5' upstream from the transcription start site of all 14 DELLA-responsive genes. To test whether DELLA directly binds to the promoters of its targets and whether this CCT element plays an important role in DELLA binding, we performed chromatin immunoprecipitation (ChIP) experiments.

### RGA Interacted with Target Promoters in Vivo

ChIP is a powerful technique to detect protein-DNA interactions in vivo (Orlando, 2000) and has been used effectively to verify putative direct target genes of transcription factors in *Arabidopsis* (Wang et al., 2002; Wigge et al., 2005; Levesque et al., 2006). DELLA proteins do not have a bona fide DNA binding domain. However, other members of the GRAS superfamily, SHR and SCR, are capable of interacting with DNA by ChIP-qPCR assays (Levesque et al., 2006; Cui et al., 2007). We therefore employed

this assay to test whether RGA directly interacts with the promoters of the early DELLA-induced genes identified by microarrays. To pull-down RGA protein efficiently from *Arabidopsis*, we generated transgenic lines that express a RGA fusion protein with a TAP (alternative tandem affinity purification) tag. The TAP tag sequence contains nine copies of the Myc epitope, six His residues, the cleavage site of the 3C protease, and two copies of IgG binding sequence (Rubio et al., 2005). This RGA-TAP fusion protein is responsive to GA-induced degradation (Figure 4A) and is functional in planta to rescue the *rga-24* null allele defect (see Supplemental Figure 3 online). Using IgG-coated beads, RGA-TAP can be pulled down efficiently from chromatin preparations from *RGA-TAP* plants (Figure 4B). ChIP, followed by gene-specific real-time qPCR, was performed using chromatin from the control



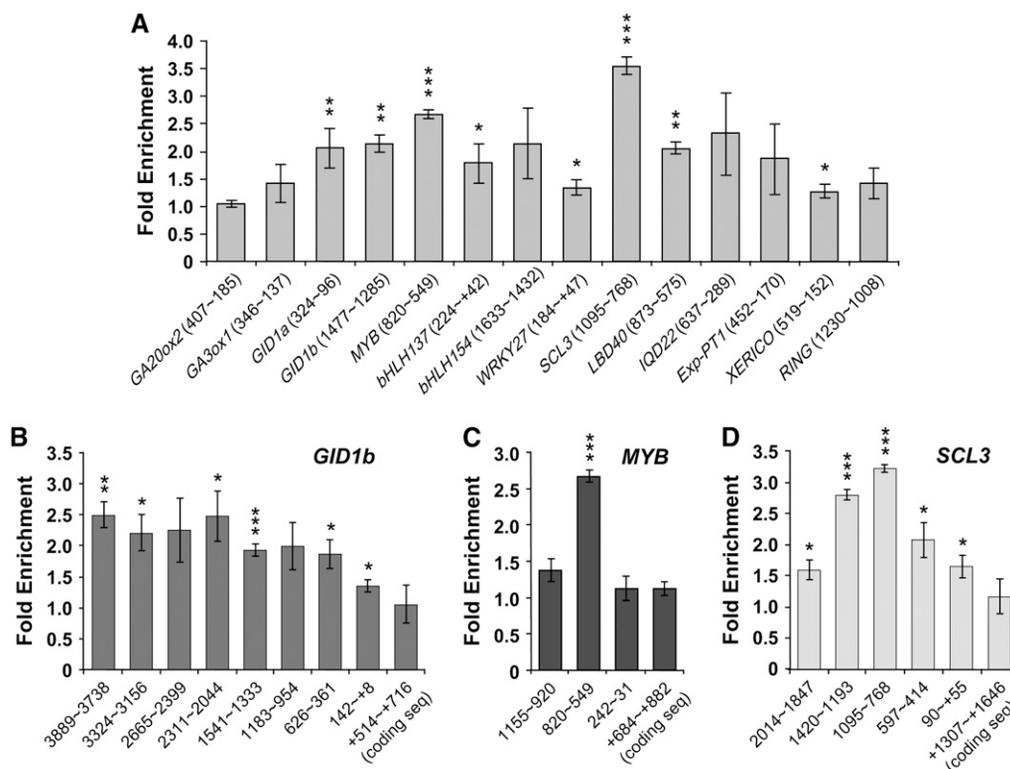
**Figure 4.** GA-Induced Degradation and Efficient Immunoprecipitation of RGA-TAP.

**(A)** The RGA-TAP fusion protein is responsive to GA treatment. Total proteins were extracted from control (*rga-24 ga1-3*) and the *RGA-TAP* line (also in the *rga-24 ga1-3* background) that were treated with water (–) or 2  $\mu$ M GA<sub>4</sub> (+) for 1 h.

**(B)** Immunoprecipitation of RGA-TAP from nuclear extracts with IgG-Sepharose beads. Isolated chromatin from control and *RGA-TAP* lines that were treated with 1% formaldehyde was incubated with IgG beads to pull down RGA-TAP. I, input (total protein extract before immunoprecipitation); S, supernatant after immunoprecipitation; E, eluate. Protein blots were probed with crude anti-RGA antibody. Arrows indicate the position of the RGA-TAP protein. Bands marked with \*\* and \* are truncated RGA-TAP fusion protein and nonspecific cross-reacting proteins, respectively.

*rga-24 ga1-3* and the *rga-24 ga1-3* RGA-TAP transgenic line. For each putative RGA target promoter, qPCR primers were designed to amplify ~250-bp sequences within the 1-kb 5' upstream promoter region. Because the transcription start site has not been determined for all genes, we will refer to the position as bases upstream of the ATG. Whenever possible, the primer sets were designed to amplify promoter sequences containing at least one CCT element. The *18S rRNA* gene was used to normalize the qPCR results in each ChIP sample. A 1.3- to 3.5-fold consistent enrichment was observed for promoter sequences of *GID1a*, *GID1b*, *MYB*, *bHLH137*, *WRKY27*, *SCL3*, *LBD40*, and *XERICO* in the RGA-TAP samples (Figure 5A, Table 2), supporting that these genes are RGA direct targets. The subtle enrichment observed so far may be because the PCR primers do not amplify the optimal RGA binding region of the promoter of the putative target genes. Alternatively, the RGA-DNA interaction may not be direct, but through other transcription factor(s), the cross-linking of RGA-TAP to DNA would be less efficient. The first possibility was tested by performing ChIP-qPCR using additional primer sets spanning different regions within the 5' upstream sequences of

*GA3ox1* (4 kb), *GA20ox2* (3 kb), *GID1b* (4 kb), *MYB* (1.2 kb), and *SCL3* (2 kb). A primer set that amplified the coding region of each gene was also included in the qPCR analysis as an additional control. We observed up to a 2.5-, 2.7-, and 3.2-fold enrichment for *GID1b*, *MYB*, and *SCL3*, respectively (Figures 5B to 5D). However, no significant enrichment was obtained for *GA3ox1* or *GA20ox2* (see Supplemental Figure 4 online). The consistent but moderate enrichment of promoters of eight DELLA-responsive genes in these ChIP experiments suggest that RGA may be associated with its target promoters via interaction with additional DNA binding proteins. The CCT element alone seemed to be insufficient for RGA interaction because several promoter regions containing this element were not significantly enriched (e.g. *Exp-PT1* [452~170], *GA3ox1* [428~257], and *GA20ox2* [201~18]) (Figure 5; see Supplemental Figure 4 online). In addition, some promoter regions without the CCT element were enriched by ChIP with RGA-TAP (e.g. *SCL3* [1095~768] and *GID1b* [1477~1285]). These observations suggest that the CCT element may not be required for RGA binding, consistent with the hypothesis that other transcription factors are involved and



**Figure 5.** RGA-TAP Binds the Promoters of Its Putative Direct Targets in Vivo.

(A) Chromatin preparations of the control line (*rga-24 ga1-3*) or the *rga-24 ga1-3* RGA-TAP line were subjected to ChIP followed by qPCR. Fold enrichment of each promoter region in the RGA-TAP line was calculated by comparing to the control line. The numbers adjacent to the gene names indicate base pairs upstream of the ATG of each gene. A + indicates base pairs downstream of the ATG. The values for fold enrichment for most genes are the average  $\pm$  SE of at least two qPCR reactions from three independent ChIP experiments. The values for *MYB*, *bHLH154*, *LBD40*, and *RING* are the average of three qPCR reactions from one ChIP experiment.

(B) to (D) The promoters of *GID1b*, *MYB*, and *SCL3* were scanned to identify sequences with maximal interaction with RGA-TAP. As an additional negative control, a coding region in each gene was also analyzed by qPCR. The numbers below each bar indicate the region amplified by qPCR as in (A).

In (A) to (D), *t* tests were performed using the statistical package SAS 9.1.3. \*\*\*,  $P < 0.01$ ; \*\*,  $P < 0.05$ ; \*,  $P < 0.1$ .



Besides *IAA13* and *WRI1*, GA treatment appears to also affect several additional genes that are involved in other hormone pathways. *XERICO*, a GA-repressed gene, plays a role in regulating ABA accumulation (Ko et al., 2006). In addition, the *BR6ox2* (*Brassinosteroid-6-oxidase2*) gene, which was downregulated by GA, encodes a cytochrome p450 enzyme (CYP85A2) that catalyzes the last step in brassinosteroid (BR) biosynthesis (Shimada et al., 2003; Nomura et al., 2005).

### Characterization of the *xerico* Mutant and Interaction between GA and ABA Pathways

*XERICO* is one of the putative DELLA target genes, and its transcript levels were induced by DELLA and repressed by GA (Table 2, Figure 3). *XERICO* was named for the drought tolerance phenotype of overexpression of this gene using the *CaMV35S* promoter in *Arabidopsis* (Ko et al., 2006). This drought tolerance phenotype is accompanied by an elevated level of ABA in the plant and hypersensitivity to salt and ABA during seedling growth. These observations suggested that *XERICO* plays a role in ABA metabolism, presumably by upregulating ABA biosynthesis or by downregulating ABA catabolism. Our finding that *XERICO* is a putative DELLA target suggests that DELLA proteins may induce ABA accumulation by upregulating *XERICO*. However, in the previous study (Ko et al., 2006), no loss-of-function *xerico* mutant was included. To verify the physiological role of *XERICO*, we obtained a T-DNA insertion *xerico* mutant (SALK\_075188), in which the T-DNA is inserted into the first exon that is 560-bp upstream from the start codon (Figure 6A). In the homozygous *xerico* mutant, no full-length WT *XERICO* transcript was detected by RT-PCR using primers (P3 and P1r) flanking the T-DNA insertion site (Figures 6A and 6B). Because the T-DNA is inserted into the 5' untranslated region of this gene, we also tested whether any truncated transcripts downstream from the insertion site were present in this mutant. Two pairs of primers (P1f+P1r and P2f+P2r) downstream of the T-DNA insertion site were used for real-time qRT-PCR. In the *xerico* mutant, the gene-specific transcript is still detectable but at ~10-fold lower level than in the wild-type Col-0 plants. Therefore, this *xerico* mutant is a leaky allele rather than a null allele.

Because overexpression of *XERICO* leads to high ABA contents, the loss-of-function *xerico* allele might have lower amounts of endogenous ABA and be more resistant than the wild type to ABA treatment during seedling establishment. Indeed, we found that the *xerico* mutant was more resistant to exogenous ABA treatment (at 0.75 to 1.25  $\mu$ M; Figure 6C). In addition, the seeds of this *xerico* mutant contained lower amounts of endogenous ABA than the wild type (Figure 6D). Although we only characterized one loss-of-function *xerico* allele, its phenotype is opposite to the *XERICO* overexpression line, suggesting that the ABA-deficient phenotype is caused by the *xerico* mutation and supporting the idea that *XERICO* plays a role in ABA metabolism.

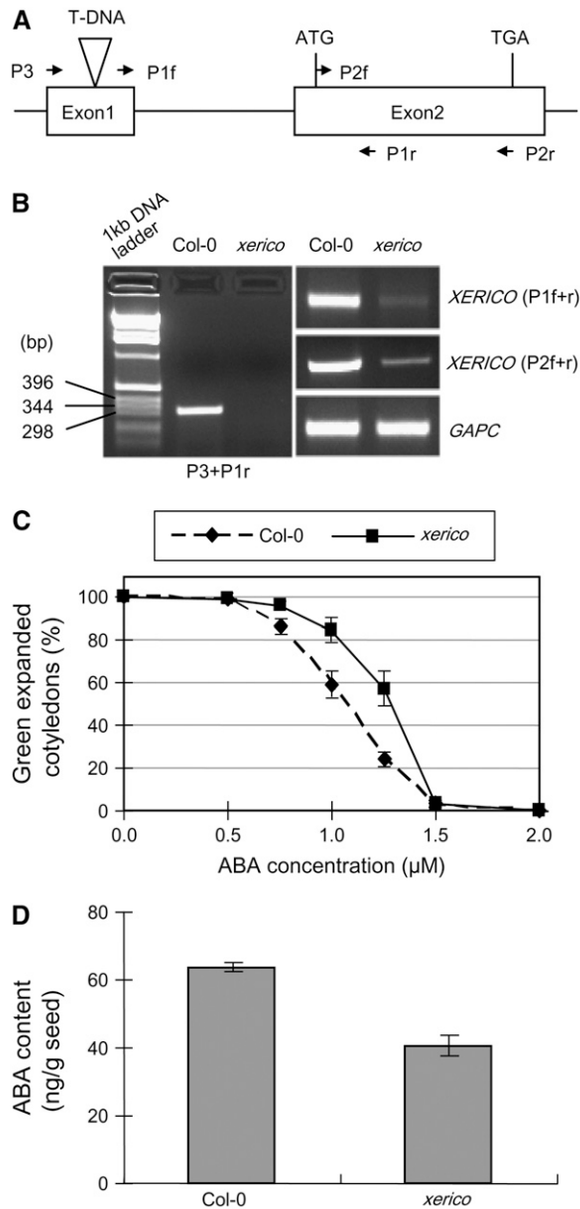
GA and ABA play antagonistic roles, with GA promoting and ABA inhibiting seed germination, seedling growth, and flower initiation (Koornneef et al., 1991; Gazzarrini and McCourt, 2003; Razem et al., 2006). Our data suggest that one function of the DELLA proteins is to upregulate ABA accumulation by inducing *XERICO* expression.

### ABA Inhibits GA Signaling Downstream of DELLA Proteins

Previous studies in cereal aleurone suggest that ABA inhibits GA responses by acting downstream of DELLA proteins (Gómez-Cadenas et al., 2001; Gubler et al., 2002; Zentella et al., 2002). The *sln1a* mutant of barley, which contains a recessive mutation in the *DELLA* gene *SLN1*, exhibits constitutive GA responses. In the aleurone of wild-type embryoless half seeds, expression of  $\alpha$ -amylase genes requires the addition of GA, but in *sln1a*  $\alpha$ -amylase, expression is constitutive. ABA treatment can effectively block  $\alpha$ -amylase production in *sln1a* (Chandler, 1988; Lanahan and Ho, 1988), but it does not protect against GA-induced SLN1 degradation (Gubler et al., 2002). These observations indicate that ABA inhibits GA responses downstream of DELLA proteins. By contrast, by analyzing a GFP-RGA protein in transgenic *Arabidopsis* (*Ler/PRGA:GFP-RGA*), it was reported recently that 2-h 20  $\mu$ M ABA pretreatment inhibited the GA (10  $\mu$ M GA<sub>3</sub>)-induced degradation of this fusion protein (Achard et al., 2006). This discrepancy could be due to the differences in species and/or tissues, or endogenous versus fusion protein. To investigate the effect of ABA on the GA signaling pathway and specifically on DELLA protein stability, we monitored the levels of endogenous RGA in response to exogenous ABA and/or GA in the *ga1-3* mutant. This mutant background was chosen to avoid the indirect effect of ABA on DELLA stability because ABA is known to have an inhibitory effect on GA biosynthesis (Seo et al., 2006; Oh et al., 2007).

Seedlings of *ga1-3* were pretreated for 2 h with water or 20  $\mu$ M ABA, followed by 1 h treatment with water, GA<sub>3</sub> (10  $\mu$ M), ABA (20  $\mu$ M), or a combination of ABA and GA (as in Achard et al., 2006). Figure 7 shows that the endogenous RGA levels did not change when seedlings were treated with either water or ABA. By contrast, GA treatment, in the presence or absence of ABA, led to RGA degradation, similar to what was observed in barley aleurone by Gubler et al. (2002). We also found that RGA responded more rapidly to GA than the GFP-RGA fusion protein (Figure 7; see Supplemental Figure 5A online). Therefore, lower concentrations of GA were also tested in the *ga1-3* mutant. However, ABA did not prevent RGA degradation even in the presence of 0.1  $\mu$ M GA<sub>3</sub> (Figure 7). To ensure that our ABA treatment was effective, we analyzed the transcript levels of *RD29A*, an ABA-responsive gene (Yamaguchi-Shinozaki and Shinozaki, 1994), and found that the amount of *RD29A* mRNA was >200-fold higher in the ABA-treated sample than in the water control (see Supplemental Figure 5C online).

We also performed the ABA and GA treatment experiment using *Ler/P<sub>RGA</sub>:GFP-RGA* as described by Achard et al. (2006) and found that GFP-RGA did accumulate to a slightly higher level by ABA in the presence or absence of GA treatment (see Supplemental Figure 5A online). Because ABA may reduce GA biosynthesis and lower bioactive GA levels in the *Ler/PRGA:GFP-RGA* plant, the observed effect of ABA treatment on GFP-RGA stability may be indirect. To test this hypothesis, we measured the relative mRNA levels of GA metabolic genes in this line. Our qRT-PCR data indicated that, indeed, the transcript levels of a GA biosynthetic gene, *GA20ox1*, were significantly reduced by ABA (see Supplemental Figure 5B online). Conversely, mRNA levels of the GA catabolic gene *GA2ox6* were upregulated by ABA.



**Figure 6.** The T-DNA Insertion Site and ABA-Resistant Phenotype of the *xerico* Mutant.

**(A)** The genomic structure of *XERICO* is shown. The position of the T-DNA insertion is indicated by a triangle above the genomic structure. The locations and orientations of primers used to detect wild-type or truncated transcripts are also shown in this diagram.

**(B)** The wild-type or truncated transcripts of the *XERICO* gene in Col-0 (wild type) and the homozygous *xerico* mutant using RNA isolated from the wild type and the homozygous *xerico* mutant using three sets of primers as labeled. Using P3 and P1r, the wild-type transcript was only detected in Col-0 but not in the *xerico* mutant (left panel). However, the mutant still accumulated truncated *xerico* transcripts (~10% of wild-type amounts), which were detected using two pair of primers (P1f + P1r and P2f + P2r) that are downstream of the T-DNA insertion site. The agarose gel image shown contains the qRT-PCR products after 30 cycles of amplification.

**(C)** Seedling establishment of the *xerico* mutant is resistant to ABA.

## DISCUSSION

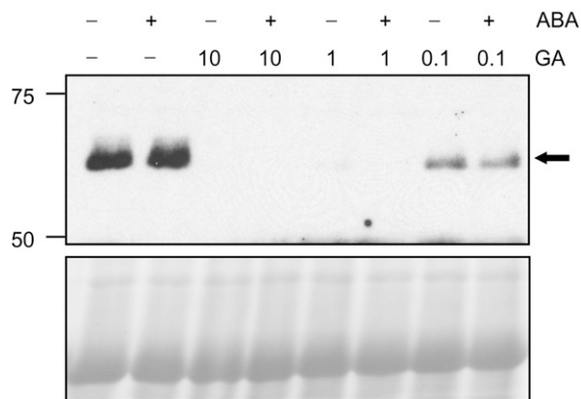
DELLA proteins are repressors that act directly downstream of the GA receptor to modulate all aspects of GA-induced growth and development in plants (Thomas and Sun, 2004; Griffiths et al., 2006; Nakajima et al., 2006). Elucidating the early gene regulatory network downstream of DELLA is crucial to our understanding of how GA controls plant development. By microarray analysis, we identified early GA- and DELLA-responsive genes that are putative DELLA direct targets in shoots of young *Arabidopsis* seedlings. Surprisingly, all of them are GA repressed and DELLA induced. ChIP-qPCR experiments provided evidence for in vivo interaction of a DELLA protein, RGA, with its putative target promoters. Several DELLA targets encode GA biosynthetic enzymes and GA receptors, indicating direct involvement of DELLA in feedback regulation. Most of the remaining DELLA target genes encode transcription factors/regulators, and ubiquitin E2 and E3 enzymes, which may be negative regulators acting downstream of DELLA in the GA response pathway. Our study also revealed a role of DELLA in mediating interaction between GA and ABA pathways.

### Identification of DELLA Target Genes Involved in GA Responses

Because previous mutant analysis indicated that DELLA proteins affect all aspects of GA responses, we originally expected to find early GA-responsive genes to be mostly DELLA responsive. However, only 14 overlapping genes were identified between the GA-responsive and DELLA-responsive data sets in this study. This is likely due to two factors. First, most of the GA-responsive genes have very subtle changes in their expression at the 1-h GA treatment time point and therefore were difficult to detect in our experiments. For example, *bHLH137*, *LBD40*, and *WRKY27* were not among the 14 overlapping genes, although qRT-PCR analysis supported that they are both GA and DELLA responsive (Figure 3). Secondly, the DEX-inducible system has a significant background noise because GVG appears to affect transcription of many genes. We attempted to remove genes whose expression was mainly altered by GVG by filtering the DELLA-induced gene list with those genes that were affected in the vector line. However, the remaining genes on the list are not all DELLA responsive when we compared their expression in the wild type, *ga1-3*, *ga1-3 rga-24 gai-t6*, and *PRGA:(rga-Δ17)* by qRT-PCR (e.g., *BT4*, *Exp-PT2*, and *UBC17*; Figure 3). In addition, *IQD22*, which responded to GA and DELLA (Table 2, Figure 3), was originally excluded from the overlapping gene list because GVG had

Seeds of Col-0 and the homozygous *xerico* mutant were incubated at 22°C for 7 d under continuous light, and seedlings with fully expanded green cotyledons were scored. Data represent means ± SE of three replicates (100 seeds per treatment). Similar results were obtained in three independent experiments.

**(D)** ABA contents are lower in the *xerico* mutant seeds than in Col-0. Data represent means ± SE of three replicates. Similar results were obtained in an independent experiment. A *t* test indicated that the difference between means was highly significant ( $P < 0.01$ )



**Figure 7.** ABA Inhibits GA Signaling Downstream of RGA.

Five-day-old *ga1-3* seedlings were pretreated with water or 20  $\mu$ M ABA for 2 h, followed by treatment with the indicated concentration ( $\mu$ M) of GA<sub>3</sub> for 1 h. Total proteins were extracted from whole seedlings and analyzed by immunoblotting with affinity-purified anti-RGA antibody. Ponceau staining was used to confirm equal loading. Arrow indicates the position of the RGA protein.

a strong effect on its expression (FC = 5.5,  $P = 0.004$ ; see Supplemental Table 3 online). Due to these limitations, we may have missed some of the DELLA targets whose expression is very subtly affected by GA treatment or strongly affected by GVG. In addition, the 475 putative DELLA-responsive genes listed in Supplemental Table 4 online need to be verified by further experimentation. The levels of *rga-Δ17* protein in the DEX-*(rga-Δ17)* transgenic line, after 2 and 4 h of DEX treatment, are ~8 and 16 times higher than RGA in the *ga1-3* mutant, respectively (see Supplemental Figure 1B online). It is possible that some of the DELLA-responsive genes listed in Supplemental Table 4 online are false positives due to overexpression of *rga-Δ17*. However, for our final list of 14 DELLA target genes, we have strong evidence from analyzing their expression in DELLA loss-of-function and gain-of-function mutants that they are indeed responsive to both DELLA and GA. Similarly, the use of a 35S:*RGA-TAP* transgenic line for CHIP-qPCR analysis may raise concerns because overexpression of a transcription factor/regulator may result in nonspecific binding to DNA and/or other proteins. Phenotypic characterization of this line suggests that this is unlikely. The transgenic line 35S:*RGA-TAP* in the double mutant *rga-24 ga1-3* background displayed a dwarf phenotype that is nearly identical to *ga1-3* (see Supplemental Figure 3 online), indicating that this transgene rescued the *rga* defect as effectively as the endogenous RGA.

#### A Direct Role of DELLA in Maintaining GA Homeostasis

GA homeostasis is achieved by a feedback mechanism that appears to coordinate activities in the GA metabolic and response pathways (Hedden and Phillips, 2000; Sun and Gubler, 2004). Under GA-deficient conditions or in mutants with reduced GA signaling (e.g., *gai-1*, *rga-Δ17*, and *sly1*), transcript levels of GA biosynthetic genes, such as *GA20ox* and *GA3ox*, are up-

regulated, whereas expression of the GA catabolic gene *GA2ox* is downregulated. Conversely, GA application or mutations that lead to increased GA signaling, such as *rga* and *gai* null alleles, cause reduced expression of *GA3ox* and *GA20ox* and elevated expression of *GA2ox*. Recently, expression of *GID1* genes in *Arabidopsis* was also found to be under feedback regulation. Transcript levels of all three *GID1* genes are downregulated upon GA treatment (Griffiths et al., 2006). In addition, *GID1* transcript levels were more elevated in *ga1-3* and *P<sub>RGA</sub>:(rga-Δ17)* than in the wild type. In the triple (*ga1 rga gai*) and quintuple (*ga1 rga gai rgl1 rgl2*) null mutants, their transcripts are similar to the wild type (Cao et al., 2006; Griffiths et al., 2006). Although the feedback phenomena have been well documented, the molecular mechanism involved is unclear. Evidence presented in this report supports that *GA3ox1*, *GA20ox2*, *GID1a*, and *GID1b* may be direct DELLA targets. Our microarray and qRT-PCR data showed that these genes are early GA and DELLA responsive. CHIP-qPCR assays further indicate that RGA is associated with the promoters of *GID1a* and *GID1b* in vivo. The lack of significant enrichment of *GA20ox2* and *GA3ox1* promoters could be because DELLA associates with its target sequences via other DNA binding proteins. In this study, *GID1c* was not identified as an early GA- and DELLA-responsive gene, probably because its responses to GA and DELLA are subtler than *GID1a* and *GID1b*. This possibility is supported by our previous qRT-PCR analysis of these genes (Griffiths et al., 2006). Surprisingly, we did not find any *GA2ox* genes in our overlapping gene list or in either of our individual lists for each microarray experiment, suggesting that *GA2ox* may respond at a later time point and is not a direct DELLA target. Our results suggest that DELLA proteins not only are repressors of GA signaling, but they also modulate GA homeostasis by upregulating expression of GA biosynthetic and GA receptor genes. A similar mechanism has been reported recently in the BR pathway, in which a positive regulator of BR signaling (BZR1) represses transcription of several BR biosynthetic genes (He et al., 2005).

#### Putative DELLA Downstream Targets: Transcription Factors/Regulators and Ubiquitin E2/E3 Enzymes

Several putative DELLA target genes are predicted to function in transcriptional regulation or in proteolysis of downstream GA response components. Among the previously studied *Arabidopsis* bHLH genes, *SPATULA* and *PIL5* inhibit seed germination by repressing *GA3ox* transcription and inducing expression of *GA2ox* (Penfield et al., 2005; Oh et al., 2006). Further studies on *PIL5* demonstrated that *RGA* and *GAI* are its direct targets, whereas its regulation of GA metabolic genes is indirect via an unknown mechanism (Oh et al., 2007). MYB and bHLH protein complexes have been shown to function in cellular pathways, such as anthocyanin biosynthesis, trichome formation, and ABA and drought responses (Lloyd et al., 1992; Abe et al., 2003; Zhang et al., 2003). It is conceivable that products of the DELLA-induced genes *bHLH137*, *bHLH154*, and *MYB* may act as repressors of GA signaling, either by themselves or as heterodimers. Characterized *Arabidopsis* WRKY genes function in various plant responses to developmental cues or to pathogens (Eulgem et al., 2000). Two rice WRKYs (Os WRKY51 and 71) when transiently expressed in

aleurone cells, act as transcriptional repressors of GA signaling that interfere with GAMYB and block  $\alpha$ -amylase gene expression (Zhang et al., 2004; Xie et al., 2006). DELLA-induced *WRKY27* may play a similar role in regulating GA signaling in *Arabidopsis*.

*SCL3* belongs to the plant-specific GRAS family of putative transcriptional regulators (Pysh et al., 1999; Bolle, 2004). Two *Arabidopsis* GRAS members, *SCR* and *SHR*, regulate radial root patterning (Di Laurenzio et al., 1996; Nakajima et al., 2001). Recent microarray and ChIP-qPCR analysis indicated that *SHR* binds to promoters of *SCR* and *SCL3* and activates their transcription (Levesque et al., 2006). Together with evidence presented here, GRAS proteins can regulate transcription by interacting with target promoters, either directly or through other transcription factor(s). GRAS proteins also appear to function in transcriptional networks to modulate expression of other GRAS genes.

*LBD40* belongs to the class II subgroup of plant-specific AS2/LBD protein family (Iwakawa et al., 2002; Shuai et al., 2002). All members of this family contain an LBD domain with a conserved Cys-rich motif, which may form a zinc finger. However, only class I LBD proteins contain a coiled-coil protein-protein interaction domain, suggesting that classes I and II may have distinct functions. Two previously characterized class I *LBD* genes (*AS2* and *LBD36*) encode nuclear proteins that control the development of lateral organs (leaf and/or flowers) (Iwakawa et al., 2002; Chalfun-Junior et al., 2005). GA also appears to promote lateral organ differentiation (Sakamoto et al., 2001; Hay et al., 2002). One possible role of *LBD40* and perhaps other class II LBD members is negative regulation of GA-promoted lateral organ development.

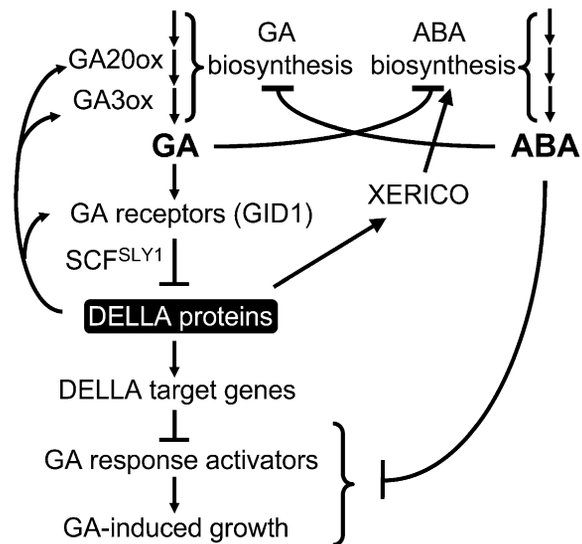
*IQD22* and *BT4* encode nuclear CaM binding proteins, suggesting that  $Ca^{2+}$  may mediate GA-regulated gene expression. *IQD22* belongs to the *IQD* family with a plant-specific IQ domain that contains three CaM binding motifs (Abel et al., 2005). *BT4* has two protein-protein interaction domains: a BTB/POZ domain for dimer or oligomer formation and a zinc finger TAZ domain that is present in transcriptional modulators (Kanai et al., 2000; Du and Poovaiah, 2004). Future studies on *IQD22* and *BT4* will shed light on the regulatory mechanism of these proteins and  $Ca^{2+}$  in GA responses.

*XERICO* is an H2-type RING E3 protein that when overexpressed in *Arabidopsis* causes dramatic ABA accumulation, which leads to increased drought tolerance (Ko et al., 2006). However, transcript levels of ABA biosynthetic genes are not affected by overexpression of *XERICO*, suggesting that *XERICO* may induce ABA accumulation by affecting the activity of an ABA metabolic enzyme(s). *XERICO* contains a putative chloroplast transit peptide (Emanuelsson et al., 1999), although the SubCellular Proteomic Database (<http://www.plantenergy.uwa.edu.au/applications/suba>) predicts that it is equally likely to be cytoplasmic or plastid localized (Heazlewood et al., 2005, 2007). Early portions of the ABA biosynthetic pathway (until the step catalyzed by 9-*cis*-epoxycarotenoid dioxygenase) occur in the plastid, whereas later steps are localized in the cytosol (Nambara and Marion-Poll, 2005). One possible role of *XERICO* would be to mediate inactivation or degradation of a negative regulator of ABA biosynthesis. Our analysis of the loss-of-function *xerico* mutant supports that *XERICO* indeed promotes ABA accumulation. Our data also suggest that DELLA may upregulate ABA accumulation by inducing *XERICO* expression, revealing an interesting regulatory circuitry between GA and ABA pathways.

### Model of the GA Biosynthesis and Signaling Networks and Interaction between GA and ABA Pathways

Our model (Figure 8) reflects the regulatory networks that control GA biosynthesis and GA signaling pathways as well as the interaction between GA and ABA pathways. *GA20ox* and *GA3ox* catalyze the final steps in the synthesis of bioactive GA. Upon binding to the GA receptor *GID1*, the GA-*GID1* complex targets DELLA proteins for ubiquitination and degradation by the 26S proteasome via the *SCF<sup>SLY1</sup>* E3 ligase. Based on our microarray and qRT-PCR data (Table 2), DELLA inhibits GA signaling by activating its downstream target genes (presumably encoding GA signaling repressors). DELLA also feedback induces expression of *GA20ox2*, *GA3ox1*, and *GID1* (Table 2; Griffiths et al., 2006). In addition, DELLA proteins induce *XERICO* expression, which then promotes ABA accumulation (Ko et al., 2006). Inhibition of GA-promoted processes by DELLAs is therefore achieved by modulating both GA and ABA pathways.

Although ABA was shown to stabilize the GFP-RGA fusion protein (Achard et al., 2006; Penfield et al., 2006), the endogenous RGA protein is not protected by ABA treatment in our study. Our results agree with the studies in barley aleurone showing that ABA interferes with the GA signaling pathway by acting downstream of DELLA (Gómez-Cadenas et al., 2001; Gubler et al., 2002). In addition, ABA and GA mutually affect each other's metabolism. For example, the *Arabidopsis* ABA-deficient mutant *aba2* exhibits increased GA biosynthesis (Seo et al., 2006), and the GA-deficient mutant *ga1-3* accumulates higher amounts of ABA (Oh et al., 2007). Also, we observed that ABA treatment may



**Figure 8.** Model of GA Signaling Network and Interaction between GA and ABA Pathways in *Arabidopsis*.

GA and ABA not only mutually inhibit each other's biosynthesis but also promote each other's catabolism (data not shown). Inhibition of GA-promoted processes by DELLA proteins is achieved by modulating both GA and ABA pathways. DELLA also plays a direct role in maintaining GA homeostasis by inducing genes encoding GA biosynthetic enzymes and GA receptors. Our data support the idea that ABA inhibits GA signaling downstream of DELLA, although ABA direct targets need to be elucidated.

affect GA levels by decreasing the expression of GA biosynthetic genes and inducing those involved in GA catabolism (see Supplemental Figure 5B online). Therefore, interactions between GA and ABA pathways occur in both their metabolic and signaling pathways to balance the antagonistic activities of these two hormones in plants.

This study uncovered two roles of DELLA proteins: direct feedback regulation of GA homeostasis and interaction with the ABA pathway. Most of the DELLA immediate targets are regulatory proteins, some of which may function in modulating transcription of downstream GA-responsive genes, and others regulate protein activity or stability of their targets. Future reverse genetic and biochemical studies of these newly identified DELLA targets will help to dissect the downstream regulatory network in GA signaling, which is responsible for the eventual changes in plant growth and development.

## METHODS

### Plasmid Constructions

Primer sequences are listed in Supplemental Table 5 online. DNA sequencing was performed to confirm the absence of sequence errors in plasmid inserts that were generated by PCR amplification. For making the DEX-inducible *rga-Δ17* [DEX-(*rga-Δ17*)] construct (pRG217), the *rga-Δ17* coding region was amplified using the primers 503 and 504 from pRG59 (Dill et al., 2001). The PCR product was purified and digested with *Sall* and *SpeI* and subcloned into the *XhoI-SpeI* sites of the binary vector pTA7001 (Aoyama and Chua, 1997). For RGA-TAP overexpression, the RGA coding sequence was amplified by PCR using primers RGA-224 and NotRGAR-34 and cloned into pCR4Blunt-TOPO (Invitrogen) to generate pCRRGA. The *BamHI-NotI* fragment from pCRRGA was cloned into *BamHI-NotI* sites of pENTR1A (Invitrogen), resulting in pENTRRGA (without stop codon). The RGA fragment of pENTRRGA was inserted into the binary vector pYL436 (Rubio et al., 2005) by Gateway reaction using Gateway LR Clonase II enzyme mix (Invitrogen) to generate pKM30 (*CaMV35S:RGA-TAP*).

### Plant Materials

All mutant and transgenic lines in this study derived from *Arabidopsis thaliana* ecotype *Ler* (wild type), with the exception of the *xerico* mutant that was isolated from the Col-0 ecotype. In the latter case, Col-0 was used as the wild-type control. The homozygous mutants *ga1-3*, *ga1-3 rga-24*, and *ga1-3 rga-24 gai-t6* and the transgenic lines *PRGA:rga-Δ17* and *PRGA:GFP-RGA* were described previously (Dill et al., 2001; Dill and Sun, 2001; Silverstone et al., 2001). All new transgenic lines were generated by *Agrobacterium tumefaciens*-mediated transformation using the floral dip method (Clough and Bent, 1998). For the DEX-inducible system, *Ler* was transformed with pRG217 and pTA7001 to generate the DEX-(*rga-Δ17*) lines and vector control lines, and transformants were selected on Murashige and Skoog (MS) medium containing 35 mg/mL hygromycin B (Sigma-Aldrich). Lines with a 3:1 ratio of resistant:sensitive (in T2) were tested in the T3 generation to identify homozygous transgene plants. The vector control line L7001-465 and DEX-(*rga-Δ17*) line L217-3b11 that accumulated most similar levels of *GVG* mRNA were used for the microarray experiment. To generate the RGA-TAP transgenic lines, *ga1-3 rga-24* was transformed with plasmid pKM30. Lines with a 3:1 (resistant:sensitive) segregation ratio in the T2 generation were selected on MS plates supplemented with 40 mg/L gentamicin sulfate. Line C3-1 fully complemented the *rga-24* mutation. T2 and T3 plants were used for immunoblotting and ChIP experiments.

### Plant Growth Conditions and Sample Preparations

Seeds were surface sterilized and stratified for 4 d at 4°C in water or in 50 μM GA<sub>4</sub> (for seeds containing the *ga1-3* mutation). GA-treated seeds were washed thoroughly before planting. Approximately 500 seeds were arranged in two rows per square plates (100 × 15 mm) and incubated at 22°C under constant light (100 μmol m<sup>-2</sup> s<sup>-1</sup>). For experiments involving line characterizations, microarray analysis, qRT-PCR, and ChIP experiments, seedlings were grown vertically on MS media containing 2% sucrose and 4 g/L of Phytigel (Sigma-Aldrich). A nylon mesh (Nitex, 100 μm pore size; Genesee Scientific) was placed on the surface of the MS media to expedite shoot harvesting. For testing the effects of GA and ABA on RGA and GFP-RGA levels, 5-d-old seedlings were grown as previously described (Achard et al., 2006).

### Immunoblot Analysis

For the *ga1-3* time-course and microarray experiments, 8-d-old seedlings were sprayed with water or 2 μM GA<sub>4</sub> (15 mL per plate) using a fine mister (Preval sprayer; Precision Valve). For time course with DEX-(*rga-Δ17*) line and the vector control, 7-d-old seedlings were sprayed with 2 μM GA<sub>4</sub>. After 16 h, the 8-d-old seedlings were resprayed with 2 μM GA<sub>4</sub> or a combination of 2 μM GA<sub>4</sub> plus 10 μM DEX (plus 0.01% Tween 20) (Sigma-Aldrich). At the indicated time, shoots were harvested, and total proteins were extracted as described (Silverstone et al., 2001) and subjected to immunoblot analysis (50 μg/sample) using affinity-purified or crude anti-RGA antibodies as described previously (Silverstone et al., 2001). To test the GA response of RGA-TAP, 8-d-old seedlings of *ga1-3 rga-24* (control) and the RGA-TAP line were sprayed with water or 2 μM GA<sub>4</sub> and incubated for 1 h. Shoots were collected, and total proteins were extracted and analyzed by immunoblotting using crude anti-RGA antibodies.

For ABA and GA treatment experiments in Figure 7 and Supplemental Figure 5 online, 5-d-old *ga1-3* and *Ler/PRGA:GFP-RGA* seedlings were treated on the plate with water, 0.1, 1, or 10 μM GA<sub>3</sub>, and/or 20 μM ABA. At the indicated time point, whole seedlings were harvested, and total proteins were extracted and analyzed by immunoblotting.

### Microarray Experiments

Total RNA from shoots of 8-d-old *ga1-3* seedlings treated for 1 h with water or 2 μM GA<sub>4</sub> was extracted using the RNeasy plant mini kit (Qiagen) according to the manufacturer's protocol. For the DEX-(*rga-Δ17*) and vector control lines, total RNA was isolated from 8-d-old shoots of seedlings that had been pretreated for 16 h by spraying with 2 μM GA<sub>4</sub> and resprayed with 2 μM GA<sub>4</sub> or 2 μM GA<sub>4</sub> plus 10 μM DEX (plus 0.01% Tween 20) for 2 or 4 h. Biotin-labeled cRNA was produced from 15 μg of total RNA, according to the manufacturer's recommendations (Affymetrix). Twenty micrograms of labeled and fragmented cRNA were hybridized with *Arabidopsis* ATH1 genome arrays. Hybridizations and array scanning were carried out at the Duke University Microarray Core Facility. Four biological replicates per condition were performed for the *ga1-3* experiment. For the DEX-(*rga-Δ17*) line, three biological replicates for each condition and time point were carried out. For the vector control line, two biological replicates per treatment and only at the 4 h time point were performed.

### Microarray Data Acquisition and Statistical Analysis

Raw data were processed with Affymetrix Microarray Suite 5.0, and the resulting CEL files were further analyzed by GeneSpring 7.2 (Agilent Technologies). A GC-RMA normalization was conducted. The data were filtered by expression level using the raw value as a cutoff, where at least one signal value for each probe set had to be ≥50. The remaining genes that are considered to be expressed were then subjected to ANOVA.

For *ga1-3*, a one-way ANOVA with a parametric test and variances not assumed equal was performed.  $P \leq 0.01$  was the cutoff. The gene list was then filtered by  $FC \geq 1.2$  between water versus GA treatment. For the *rga-Δ17* line samples, the same procedure was followed, except that we conducted two-way ANOVA (GA versus GA plus DEX and 2 h versus 4 h) with a parametric test and variances assumed equal. In addition, a multiple testing correction by Benjamini and Hochberg (1995) was applied and an  $FDR \leq 0.01$  was used as cutoff. Then, the gene list was filtered to select for genes with  $FC \geq 1.5$ . Genes whose expression was affected by DEX treatment in the vector control line (one-way ANOVA [ $P \leq 0.01$ ] and  $FC \geq 3$ ) were considered as GVG responsive and were eliminated from the *rga-Δ17*-responsive gene list. The CEL files of the experiment “GA<sub>3</sub> time-course in wild-type and *ga1-5* mutant seedlings,” performed by the AtGenExpress consortium were downloaded directly from The Arabidopsis Information Resource ([http://www.arabidopsis.org/servlets/TairObject?type=hyb\\_descr\\_collection&id=1007966175](http://www.arabidopsis.org/servlets/TairObject?type=hyb_descr_collection&id=1007966175)). The data were processed as described above for the DEX-inducible system experiment with the following modifications: the raw value cutoff was 200 (because of higher average signal values in this data set). This number gave a similar number of present genes (~15,000); no multiple testing corrections were applied to the two-way ANOVA (parameters: water versus 5 μM GA<sub>3</sub> and 0.5 h versus 1 h versus 3 h), and  $P \leq 0.01$  was used instead; finally, a 1.2-fold change in gene expression was used as cutoff. Gene lists for DELLA-dependent and -independent genes in germinating seeds and flowers were obtained from the supplemental data in Cao et al. (2006). Similarly, the gene lists from the analysis of the AtGenExpress data set by Nemhauser et al. (2006) were downloaded directly from their online supplemental data. Gene list comparisons and overlaps were performed using GeneSpring 7.2 and Microsoft Access. The microarray data sets were submitted to the Gene Expression Omnibus and are available under the accession numbers GSE8739 “early gibberellin responses in *Arabidopsis*” and GSE8741 “DELLA protein direct targets in *Arabidopsis*.”

#### Real-Time qRT-PCR Analysis

All real-time qRT-PCR measurements were performed using a Roche LightCycler (Roche Diagnostics). For all RNA samples, total RNA was isolated with the RNeasy plant mini kit (Qiagen) and treated with DNaseI using the DNA-free kit and following the manufacturer's recommendations (Ambion). For time-course transcript measurements, 8-d-old seedlings of *ga1-3*, *rga-Δ17*, and the vector control line were treated with GA and DEX as described in the “Immunoblot Analysis” section. qRT-PCR was performed with 50 ng of DNaseI-treated RNA using the LightCycler RNA Amplification SYBR Green I kit (Roche). We used the housekeeping gene *GAPC* as normalization control because its expression is not affected by GA treatment (Dill et al., 2004). The *GAPC*, *GA20ox2*, *GA3ox1*, and *RGA* primers have been described previously (Dill et al., 2004; Mitchum et al., 2006). The GVG primers and primers for putative DELLA targets are listed in Supplemental Table 5 online. To measure transcript levels of the putative DELLA targets, RNA from shoots of 8-d-old seedlings of *Ler*, *ga1-3*, *rga-24 gai-t6 ga1-3*, and *PRGA:rga-Δ17* were used. For *PRGA:rga-Δ17*, RNA extractions came from both hemizygous and homozygous transgenic seedlings (Dill et al., 2004). To measure transcript levels in response to GA treatments, 8-d-old *ga1-3* seedlings were treated with water (control) or 2 μM GA<sub>3</sub> for 1 or 3 h before harvesting. Total RNA from shoots was isolated and first-strand cDNA synthesized. Real-time qPCR was performed as previously described (Tyler et al., 2004). One-way ANOVA analyses were performed with least significant difference multiple comparison tests at an  $\alpha$  level of 0.05 using SPSS version 11.5.0.

#### ChIP-qPCR Analysis

Chromatin isolation for immunoprecipitations was performed as previously described (Bowler et al., 2004), with the following modifications:

shoots of 9- to 12-d-old seedlings of *ga1-3 rga-24* (control) and *RGA-TAP* were used. The tissue (3 to 5 g) was fixed for 15 min by vacuum infiltration in 1% formaldehyde, washed, and frozen in liquid N<sub>2</sub>. Nuclei were isolated and sonicated to obtain DNA fragments between 0.5 and 1 kb. Sonicated chromatin was aliquoted in fractions of 150 μL in siliconized microfuge tubes and kept frozen at -80°C until use. For *RGA-TAP* immunoprecipitation, IgG-Sepharose 6 Fast Flow beads (GE Healthcare Biosciences) was used. The beads were preincubated with sonicated salmon sperm DNA (20 μg/mL; Ambion) and IgG free BSA (50 μg/mL; Sigma-Aldrich) for 1 h and washed twice with ChIP dilution buffer. A 150-μL aliquot of chromatin of each line (control and *RGA-TAP*) was diluted to 1.5 mL with ChIP dilution buffer, and 150 μL of IgG-Sepharose (50% slurry) was added to each tube. After 1 h of incubation with rotation at 4°C, the beads were washed as previously described (Bowler et al., 2004). Elution and reversed cross-linking of *RGA-TAP*/DNA was done by boiling for 10 min in elution buffer. Eluates were treated with Proteinase K (60 μg/mL; Roche) for 1 h at 45°C, phenol/chloroform extracted, and ethanol precipitated in the presence of 40 μg of glycogen (Roche). The purified DNA was resuspended in 100 μL of TE.

Chromatin enrichment of putative RGA targets was determined by real-time qPCR. DNA primers are listed in Supplemental Table 5 online. The best primer set for each gene within the region of -1250 to +250 bp with respect to the ATG was chosen. All qPCR reactions were carried out using the LightCycler SYBR Green I Fast Start DNA kit (Roche). Reactions were performed with 2 μL of immunoprecipitated DNA, and samples from control and *RGA-TAP* were compared. To normalize the results between samples, the amount of *18S rRNA* gene sequence was quantified using primers described previously (Tyler et al., 2004). Primer efficiencies for each gene set were estimated (between 1.73 and 2.0) and incorporated in the calculations. Additional primer sets for *GA3ox1*, *GA20ox2*, *GID1b*, *MYB*, and *SCL3* were also designed to test enrichment of neighboring regions in their promoters. *t* tests comparing the means of promoter DNA enrichment between control and *RGA-TAP* samples were performed using the statistical package SAS 9.1.3 (SAS Institute).

#### Isolation and qRT-PCR Analysis of the *xerico* T-DNA Insertion Mutant

The *xerico* mutant (Salk\_075188) was identified by searching the SALK Institute Genomic Analysis Laboratory T-DNA express database (Alonso et al., 2003). Homozygous *xerico* mutant plants were identified by PCR using allele-specific primers (see Supplemental Table 5 online). To detect any *XERICO* transcripts produced in this mutant, RNA isolated from 8-d-old Col-0 and the homozygous *xerico* mutant seedlings were analyzed by qRT-PCR using three sets of primers (see Supplemental Table 5 online). All qRT-PCR reactions were performed using 6 mM MgCl<sub>2</sub> with an annealing temperature of 55°C and a total of 45 cycles of amplification.

#### Seedling Establishment Assay of the *xerico* Mutant

For this assay, 1-month-old seeds of Col-0 and the homozygous *xerico* mutant that were harvested at the same time were used. The seeds were surface-sterilized and plated on half-strength MS agar plates (60 × 15 mm) without sucrose in the presence of different concentrations of ABA. The plates were incubated at 4°C for 3 d in the dark and then transferred to 22°C under continuous light. After 7 d, seedlings with fully expanded cotyledons ( $\geq 180^\circ$  angles between the two cotyledons) were counted as established seedlings.

#### Measurement of ABA Levels

Dry seeds (10 mg) were used for ABA measurements. Deuterium-labeled d<sub>6</sub>-ABA (Icon Services) was added to each sample prior to extraction.

Procedures for extraction and purification were described previously (Saika et al., 2007). Resultant extracts were subjected to liquid chromatography (AQUITY UPLC system; Waters) tandem mass spectrometry (MS/MS) (Q-ToF premier; Micromass) analysis to quantify the ABA levels. Extracts were separated by a liquid chromatograph equipped with ACQUITY UPLC BEH C<sub>18</sub> column (2.1 × 50 mm, 1.7 mm; Waters) with a binary solvent system comprising acetonitrile containing water (A) and 0.05% acetic acid (B) at a flow rate of 0.2 mL/min with a linear gradient of solvent B from 3 to 98% in 10 min. The retention time of ABA and d<sub>6</sub>-ABA was 4.18 min. MS/MS conditions were as follows: capillary (kV) = 2.8, source temperature (°C) = 80, desolvation temperature (°C) = 400, cone gas flow (L/h) = 0, desolvation gas flow (L/h) = 500, collision energy = 8.0, MS/MS transition (*m/z*): 263/153 for unlabeled ABA and 269/159 for d<sub>6</sub>-ABA. The amount of ABA was calculated by spectrometer software (MassLynx version 4.1; Micromass).

#### Accession Numbers

Arabidopsis Genome Initiative locus identifiers for the genes mentioned in this article are as follows: *GID1a* (At3g05120), *GID1b* (At3g63010), *RGA* (At2g01570), *GAI* (At1g14920), *GA1* (At4g02780), *GA3ox1* (At1g15550), *GA20ox1* (At4g25420), *GA20ox2* (At5g51810), *XERICO* (At2g04240), *MYB* (At3g11280), *bHLH137* (At5g50915), *bHLH154* (At2g31730), *WRKY27* (At5g52830), *SCL3* (At1g50420), *LBD40* (At1g67100), *IQD22* (At4g23060), *CaM-BP* (At3g52870), *BT4* (At5g67480), *Exp-PT1* (At2g45900), *Exp-PT2* (At2g34340), *UBC17* (At4g36410), *RING* (At4g19700), *GAPC* (At3g04120), *RD29A* (At5g52310), and *GA2ox6* (At1g02400).

#### Supplemental Data

The following materials are available in the online version of this article.

**Supplemental Figure 1.** Quantification of RGA Protein Levels in *ga1-3* after GA Treatment and in the DEX-(*rga-Δ17*) Transgenic Line after DEX Treatment.

**Supplemental Figure 2.** *GA3ox1*, *GA20ox2*, and *GVG* Transcript Levels in the Vector Control and the DEX-(*rga-Δ17*) Transgenic Lines.

**Supplemental Figure 3.** RGA-TAP Is Functional in Planta.

**Supplemental Figure 4.** Lack of Enrichment of the Promoters of GA Biosynthetic Genes in the RGA-TAP Line after ChIP Using IgG Beads.

**Supplemental Figure 5.** Effects of GA and ABA on GFP-RGA and Expression of GA Metabolic Genes.

**Supplemental Table 1.** GA-Responsive Genes in the *ga1-3* Data Set with Significant P Values.

**Supplemental Table 2.** GA-Responsive Genes in the AtGenExpress Data Set.

**Supplemental Table 3.** DEX-Regulated Genes in the Vector Control Data Set.

**Supplemental Table 4.** Putative DELLA-Regulated Genes in the DEX-(*rga-Δ17*) Data Set.

**Supplemental Table 5.** List of Primers and Their Uses.

#### ACKNOWLEDGMENTS

We thank Frédéric Marsolais for his help in determining the conditions for the *ga1-3* microarray experiments, Nam-hai Chua for providing the binary vector pTA7001, Yuri Kanno for support with ABA measurements, and the ABRC for providing the *xerico* T-DNA insertion line pool. We also thank Hong Chang Cui for helpful discussions and advice. This

work was supported by the National Science Foundation (IBN-0235656 and IOS-0641548) to T.-p.S. and by the Japan Society for the Promotion of Science Postdoctoral Fellowships for Research Abroad to K.M.

Received August 11, 2007; revised September 14, 2007; accepted September 20, 2007; published October 12, 2007.

#### REFERENCES

- Abe, H., Urao, T., Ito, T., Seki, M., Shinozaki, K., and Yamaguchi-Shinozaki, K. (2003). Arabidopsis AtMYC2 (bHLH) and AtMYB2 (MYB) function as transcriptional activators in abscisic acid signaling. *Plant Cell* **15**: 63–78.
- Abel, S., Savchenko, T., and Levy, M. (2005). Genome-wide comparative analysis of the IQD gene families in *Arabidopsis thaliana* and *Oryza sativa*. *BMC Evol. Biol.* **5**: 72.
- Achard, P., Cheng, H., De Grauwe, L., Decat, J., Schoutteten, H., Moritz, T., Van Der Straeten, D., Peng, J., and Harberd, N.P. (2006). Integration of plant responses to environmentally activated phytohormonal signals. *Science* **311**: 91–94.
- Achard, P., Vriegen, W.H., van der Straeten, D., and Harberd, N.P. (2003). Ethylene regulates Arabidopsis development via the modulation of DELLA protein growth repressor function. *Plant Cell* **15**: 2816–2825.
- Alonso, J.M., et al. (2003). Genome-wide insertional mutagenesis of *Arabidopsis thaliana*. *Science* **301**: 653–657.
- Aoyama, T., and Chua, N.-H. (1997). A glucocorticoid-mediated transcriptional induction system in transgenic plants. *Plant J.* **11**: 605–612.
- Aoyama, T., Dong, C.H., Wu, Y., Carabelli, M., Sessa, G., Ruberti, I., Morelli, G., and Chua, N.H. (1995). Ectopic expression of the Arabidopsis transcriptional activator Athb-1 alters leaf cell fate in tobacco. *Plant Cell* **7**: 1773–1785.
- Bailey, T.L., and Elkan, C. (1994). Fitting a mixture model by expectation maximization to discover motifs in biopolymers. *Proc. Int. Conf. Intell. Syst. Mol. Biol.* **2**: 28–36.
- Bailey, T.L., and Gribskov, M. (1998). Combining evidence using p-values: Application to sequence homology searches. *Bioinformatics* **14**: 48–54.
- Bailey, P.C., Martin, C., Toledo-Ortiz, G., Quail, P.H., Huq, E., Heim, M.A., Jakoby, M., Werber, M., and Weisshaar, B. (2003). Update on the basic helix-loop-helix transcription factor gene family in *Arabidopsis thaliana*. *Plant Cell* **15**: 2497–2501.
- Benjamini, Y., and Hochberg, Y. (1995). Controlling the false discovery rate - A practical and powerful approach to multiple testing. *J. R. Stat. Soc. B* **57**: 289–300.
- Blazquez, M.A., and Weigel, D. (2000). Integration of floral inductive signals in Arabidopsis. *Nature* **404**: 889–892.
- Bolle, C. (2004). The role of GRAS proteins in plant signal transduction and development. *Planta* **218**: 683–692.
- Bowler, C., Benvenuto, G., Laflamme, P., Molino, D., Probst, A.V., Tariq, M., and Paszkowski, J. (2004). Chromatin techniques for plant cells. *Plant J.* **39**: 776–789.
- Cao, D., Cheng, H., Wu, W., Meng Soo, H., and Peng, J. (2006). Gibberellin mobilizes distinct DELLA-dependent transcriptomes to regulate seed germination and floral development in Arabidopsis. *Plant Physiol.* **142**: 509–525.
- Cao, D., Hussain, A., Cheng, H., and Peng, J. (2005). Loss of function of four DELLA genes leads to light- and gibberellin-independent seed germination in Arabidopsis. *Planta* **223**: 105–113.
- Cernac, A., Andre, C., Hoffmann-Benning, S., and Benning, C. (2006). WR11 is required for seed germination and seedling establishment. *Plant Physiol.* **141**: 745–757.

- Cernac, A., and Benning, C. (2004). WRINKLED1 encodes an AP2/EREB domain protein involved in the control of storage compound biosynthesis in Arabidopsis. *Plant J.* **40**: 575–585.
- Chalfun-Junior, A., Franken, J., Mes, J.J., Marsch-Martinez, N., Pereira, A., and Angenent, G.C. (2005). ASYMMETRIC LEAVES2-LIKE1 gene, a member of the AS2/LOB family, controls proximal-distal patterning in Arabidopsis petals. *Plant Mol. Biol.* **57**: 559–575.
- Chandler, P.M. (1988). Hormonal regulation of gene expression in the “slender” mutant of barley (*Hordeum vulgare* L.). *Planta* **175**: 115–120.
- Cheng, H., Qin, L., Lee, S., Fu, X., Richards, D.E., Cao, D., Luo, D., Harberd, N.P., and Peng, J. (2004). Gibberellin regulates Arabidopsis floral development via suppression of DELLA protein function. *Development* **131**: 1055–1064.
- Chiang, H.-H., Hwang, I., and Goodman, H.M. (1995). Isolation of the Arabidopsis GA4 locus. *Plant Cell* **7**: 195–201.
- Clough, S.J., and Bent, A.F. (1998). Floral dip: A simplified method for *Agrobacterium*-mediated transformation of *Arabidopsis thaliana*. *Plant J.* **16**: 735–743.
- Conaway, R.C., Brower, C.S., and Conaway, J.W. (2002). Emerging roles of ubiquitin in transcription regulation. *Science* **296**: 1254–1258.
- Cui, H.C., Levesque, M.P., Vernoux, T., Jung, J.W., Paquette, A.J., Gallagher, K.L., Wang, J.Y., Bilou, I., Scheres, B., and Benfey, P.N. (2007). An evolutionarily conserved mechanism delimiting SHR movement defines a single layer of endodermis in plants. *Science* **316**: 421–425.
- Davies, P.J. (2004). *Plant Hormones: Biosynthesis, Signal Transduction, Action!* (Dordrecht, The Netherlands: Kluwer Academic Publishers).
- Di Laurenzio, L., Wysocka-Diller, J., Malamy, J.E., Pysh, L., Helariutta, Y., Freshour, G., Hahn, M.G., Feldmann, K.A., and Benfey, P.N. (1996). The SCARECROW gene regulates an asymmetric cell division that is essential for generating the radial organization of the Arabidopsis root. *Cell* **86**: 423–433.
- Dill, A., Jung, H.-S., and Sun, T.-p. (2001). The DELLA motif is essential for gibberellin-induced degradation of RGA. *Proc. Natl. Acad. Sci. USA* **98**: 14162–14167.
- Dill, A., and Sun, T.-p. (2001). Synergistic de-repression of gibberellin signaling by removing RGA and GAI function in *Arabidopsis thaliana*. *Genetics* **159**: 777–785.
- Dill, A., Thomas, S.G., Hu, J., Steber, C.M., and Sun, T.-p. (2004). The Arabidopsis F-box protein SLEEPY1 targets GA signaling repressors for GA-induced degradation. *Plant Cell* **16**: 1392–1405.
- Du, L.Q., and Poovaiah, B.W. (2004). A novel family of Ca<sup>2+</sup>/calmodulin-binding proteins involved in transcriptional regulation: interaction with fsh/Ring3 class transcription activators. *Plant Mol. Biol.* **54**: 549–569.
- Emanuelsson, O., Nielsen, H., and von Heijne, G. (1999). ChloroP, a neural network-based method for predicting chloroplast transit peptides and their cleavage sites. *Protein Sci.* **8**: 978–984.
- Eulgem, T., Rushton, P.J., Robatzek, S., and Somssich, I.E. (2000). The WRKY superfamily of plant transcription factors. *Trends Plant Sci.* **5**: 199–206.
- Fleet, C.M., and Sun, T.P. (2005). A DELLAcate balance: The role of gibberellin in plant morphogenesis. *Curr. Opin. Plant Biol.* **8**: 77–85.
- Fu, X., and Harberd, N.P. (2003). Auxin promotes Arabidopsis root growth by modulating gibberellin response. *Nature* **421**: 740–743.
- Fu, X., Richards, D.E., Fleck, B., Xie, D., Burton, N., and Harberd, N.P. (2004). The Arabidopsis mutant sleepy1<sup>gar2-1</sup> protein promotes plant growth by increasing the affinity of the SCF<sup>SLY1</sup> E3 ubiquitin ligase for DELLA protein substrates. *Plant Cell* **16**: 1406–1418.
- Gazzarrini, S., and McCourt, P. (2003). Cross-talk in plant hormone signalling: What Arabidopsis mutants are telling us. *Ann. Bot. (Lond.)* **91**: 605–612.
- Gocal, G.F.W., Sheldon, C.C., Gubler, F., Moritz, T., Bagnall, D.B., MacMillan, C.P., Li, S.F., Parish, R.W., Dennis, E.S., Weigel, D., and King, R.W. (2001). *GAMYB-like* genes, flowering and gibberellin signaling in Arabidopsis. *Plant Physiol.* **127**: 1682–1693.
- Gómez-Cadenas, A., Zentella, R., Walker-Simmons, M., and Ho, T.H.D. (2001). Gibberellin/abscisic acid antagonism in barley aleurone cells: Site of action of the protein kinase PKABA1 in relation to gibberellin signaling molecules. *Plant Cell* **13**: 667–679.
- Gomi, K., Sasaki, A., Itoh, H., Ueguchi-Tanaka, M., Ashikari, M., Kitano, H., and Matsuoka, M. (2004). GID2, an F-box subunit of the SCF E3 complex, specifically interacts with phosphorylated SLR1 protein and regulates the gibberellin-dependent degradation of SLR1 in rice. *Plant J.* **37**: 626–634.
- Griffiths, J., Murase, K., Rieu, I., Zentella, R., Zhang, Z.L., Powers, S.J., Gong, F., Phillips, A.L., Hedden, P., Sun, T.P., and Thomas, S.G. (2006). Genetic characterization and functional analysis of the GID1 gibberellin receptors in Arabidopsis. *Plant Cell* **18**: 3399–3414.
- Gubler, F., Chandler, P., White, R., Llewellyn, D., and Jacobsen, J. (2002). GA signaling in barley aleurone cells: Control of SLN1 and GAMYB expression. *Plant Physiol.* **129**: 191–200.
- Gubler, F., Kalla, R., Roberts, J., and Jacobsen, J.V. (1995). Gibberellin-regulated expression of a *myb* gene in barley aleurone cells: Evidence for Myb transactivation of a high-pl  $\alpha$ -amylase gene promoter. *Plant Cell* **7**: 1879–1891.
- Gubler, F., Raventos, D., Keys, M., Watts, R., Mundy, J., and Jacobsen, J.V. (1999). Target genes and regulatory domains of the GAMYB transcriptional activator in cereal aleurone. *Plant J.* **17**: 1–9.
- Hartweck, L.M., and Olszewski, N.E. (2006). Rice GIBBERELLIN INSENSITIVE DWARF1 is a gibberellin receptor that illuminates and raises questions about GA signaling. *Plant Cell* **18**: 278–282.
- Hay, A., Kaur, H., Phillips, A., Hedden, P., Hake, S., and Tsiantis, M. (2002). The gibberellin pathway mediates KNOTTED1-type homeobox function in plants with different body plans. *Curr. Biol.* **12**: 1557–1565.
- He, J.X., Gendron, J.M., Sun, Y., Gampala, S.S., Gendron, N., Sun, C.Q., and Wang, Z.Y. (2005). BZR1 is a transcriptional repressor with dual roles in brassinosteroid homeostasis and growth responses. *Science* **307**: 1634–1638.
- Heazlewood, J.L., Tonti-Filippini, J., Verboom, R.E., and Millar, A.H. (2005). Combining experimental and predicted datasets for determination of the subcellular location of proteins in Arabidopsis. *Plant Physiol.* **139**: 598–609.
- Heazlewood, J.L., Verboom, R.E., Tonti-Filippini, J., Small, I., and Millar, A.H. (2007). SUBA: The Arabidopsis subcellular database. *Nucleic Acids Res.* **35**: D213–D218.
- Hedden, P., and Phillips, A.L. (2000). Gibberellin metabolism: New insights revealed by the genes. *Trends Plant Sci.* **5**: 523–530.
- Itoh, H., Ueguchi-Tanaka, M., Sato, Y., Ashikari, M., and Matsuoka, M. (2002). The gibberellin signaling pathway is regulated by the appearance and disappearance of SLENDER RICE1 in nuclei. *Plant Cell* **14**: 57–70.
- Iwakawa, H., Ueno, Y., Semiarti, E., Onouchi, H., Kojima, S., Tsukaya, H., Hasebe, M., Soma, T., Ikezaki, M., Machida, C., and Machida, Y. (2002). The ASYMMETRIC LEAVES2 gene of *Arabidopsis thaliana*, required for formation of a symmetric flat leaf lamina, encodes a member of a novel family of proteins characterized by cysteine repeats and a leucine zipper. *Plant Cell Physiol.* **43**: 467–478.
- Jacobsen, S.E., and Olszewski, N.E. (1993). Mutations at the SPINDLY locus of Arabidopsis alter gibberellin signal transduction. *Plant Cell* **5**: 887–896.
- Kanai, F., Marignani, P.A., Sarbassova, D., Yagi, R., Hall, R.A., Donowitz, M., Hisaminato, A., Fujiwara, T., Ito, Y., Cantley, L.C., and Yaffe, M.B. (2000). TAZ: A novel transcriptional co-activator regulated by interactions with 14-3-3 and PDZ domain proteins. *EMBO J.* **19**: 6778–6791.

- Kaneko, M., Inukai, Y., Ueguchi-Tanaka, M., Itoh, H., Izawa, T., Kobayashi, Y., Hattori, T., Miyao, A., Hirochika, H., Ashikari, M., and Matsuoka, M. (2004). Loss-of-function mutations of the rice GAMYB gene impair alpha-amylase expression in aleurone and flower development. *Plant Cell* **16**: 33–44.
- Kang, H.G., Fang, Y., and Singh, K.B. (1999). A glucocorticoid-inducible transcription system causes severe growth defects in *Arabidopsis* and induces defense-related genes. *Plant J.* **20**: 127–133.
- King, K., Moritz, T., and Harberd, N. (2001). Gibberellins are not required for normal stem growth in *Arabidopsis thaliana* in the absence of GAI and RGA. *Genetics* **159**: 767–776.
- Ko, J.H., Yang, S.H., and Han, K.H. (2006). Upregulation of an *Arabidopsis* RING-H2 gene, *XERICCO*, confers drought tolerance through increased abscisic acid biosynthesis. *Plant J.* **47**: 343–355.
- Koornneef, M., Hanhart, C.J., and van der Veen, J.H. (1991). A genetic and physiological analysis of late flowering mutants in *Arabidopsis thaliana*. *Mol. Gen. Genet.* **229**: 57–66.
- Koornneef, M., and van der Veen, J.H. (1980). Induction and analysis of gibberellin-sensitive mutants in *Arabidopsis thaliana* (L.) Heynh. *Theor. Appl. Gen.* **58**: 257–263.
- Kraft, E., Stone, S.L., Ma, L., Su, N., Gao, Y., Lau, O.S., Deng, X.W., and Callis, J. (2005). Genome analysis and functional characterization of the E2 and RING-type E3 ligase ubiquitination enzymes of *Arabidopsis*. *Plant Physiol.* **139**: 1597–1611.
- Kranz, H.D., et al. (1998). Towards functional characterisation of the members of the R2R3-MYB gene family from *Arabidopsis thaliana*. *Plant J.* **16**: 263–276.
- Lanahan, M.B., and Ho, T.-H.D. (1988). Slender barley: A constitutive gibberellin-response mutant. *Planta* **175**: 107–114.
- Lee, S., Cheng, H., King, K.E., Wang, W., He, Y., Hussain, A., Lo, J., Harberd, N.P., and Peng, J. (2002). Gibberellin regulates *Arabidopsis* seed germination via *RGL2*, a GAI/RGA-like gene whose expression is up-regulated following imbibition. *Genes Dev.* **16**: 646–658.
- Levesque, M.P., Vernoux, T., Busch, W., Cui, H., Wang, J.Y., Blilou, I., Hassan, H., Nakajima, K., Matsumoto, N., Lohmann, J.U., Scheres, B., and Benfey, P.N. (2006). Whole-genome analysis of the SHORT-ROOT developmental pathway in *Arabidopsis*. *PLoS Biol.* **4**: e143.
- Levy, M., Wang, Q., Kaspi, R., Parrella, M.P., and Abel, S. (2005). *Arabidopsis* IQD1, a novel calmodulin-binding nuclear protein, stimulates glucosinolate accumulation and plant defense. *Plant J.* **43**: 79–96.
- Lloyd, A.M., Walbot, V., and Davis, R.W. (1992). *Arabidopsis* and *Nicotiana* anthocyanin production activated by maize regulator-R and regulator-C1. *Science* **258**: 1773–1775.
- McGinnis, K.M., Thomas, S.G., Soule, J.D., Strader, L.C., Zale, J.M., Sun, T.-p., and Steber, C.M. (2003). The *Arabidopsis* *SLEEPY1* gene encodes a putative F-box subunit of an SCF E3 ubiquitin ligase. *Plant Cell* **15**: 1120–1130.
- Millar, A.A., and Gubler, F. (2005). The *Arabidopsis* GAMYB-like genes, MYB33 and MYB65, are microRNA-regulated genes that redundantly facilitate anther development. *Plant Cell* **17**: 705–721.
- Mitchum, M.G., Yamaguchi, S., Hanada, A., Kuwahara, A., Yoshioka, Y., Kato, T., Tabata, S., Kamiya, Y., and Sun, T.P. (2006). Distinct and overlapping roles of two gibberellin 3-oxidases in *Arabidopsis* development. *Plant J.* **45**: 804–818.
- Murray, F., Kalla, R., Jacobsen, J., and Gubler, F. (2003). A role for HvGAMYB in anther development. *Plant J.* **33**: 481–491.
- Nair, R., and Rost, B. (2005). Mimicking cellular sorting improves prediction of subcellular localization. *J. Mol. Biol.* **348**: 85–100.
- Nakai, K., and Kanehisa, M. (1992). A knowledge base for predicting protein localization sites in eukaryotic cells. *Genomics* **14**: 897–911.
- Nakajima, K., Sena, G., Nawy, T., and Benfey, P.N. (2001). Intercellular movement of the putative transcription factor SHR in root patterning. *Nature* **413**: 307–311.
- Nakajima, M., et al. (2006). Identification and characterization of *Arabidopsis* gibberellin receptors. *Plant J.* **46**: 880–889.
- Nambara, E., and Marion-Poll, A. (2005). Abscisic acid biosynthesis and catabolism. *Annu. Rev. Plant Biol.* **56**: 165–185.
- Nemhauser, J.L., Hong, F., and Chory, J. (2006). Different plant hormones regulate similar processes through largely nonoverlapping transcriptional responses. *Cell* **126**: 467–475.
- Nomura, T., Kushiro, T., Yokota, T., Kamiya, Y., Bishop, G.J., and Yamaguchi, S. (2005). The last reaction producing brassinolide is catalyzed by cytochrome P-450s, CYP85A3 in tomato and CYP85A2 in *Arabidopsis*. *J. Biol. Chem.* **280**: 17873–17879.
- O'Connor, T.R., Dyreson, C., and Wyrick, J.J. (2005). Athena: A resource for rapid visualization and systematic analysis of *Arabidopsis* promoter sequences. *Bioinformatics* **21**: 4411–4413.
- Ogawa, M., Hanada, A., Yamauchi, Y., Kuwahara, A., Kamiya, Y., and Yamaguchi, S. (2003). Gibberellin biosynthesis and response during *Arabidopsis* seed germination. *Plant Cell* **15**: 1591–1604.
- Oh, E., Yamaguchi, S., Kamiya, Y., Bae, G., Chung, W.I., and Choi, G. (2006). Light activates the degradation of PIL5 protein to promote seed germination through gibberellin in *Arabidopsis*. *Plant J.* **47**: 124–139.
- Oh, E., Yamaguchi, S., Hu, J., Yusuke, J., Jung, B., Paik, I., Lee, H.-S., Sun, T.-p., Kamiya, Y., and Choi, G. (2007). PIL5, a phytochrome-interacting bHLH protein, regulates gibberellin responsiveness by binding directly to the GAI and RGA promoters in *Arabidopsis* seeds. *Plant Cell* **19**: 1192–1208.
- Olszewski, N., Sun, T.-p., and Gubler, F. (2002). Gibberellin signaling: Biosynthesis, catabolism, and response pathways. *Plant Cell* **14**: S61–S80.
- Orlando, V. (2000). Mapping chromosomal proteins in vivo by formaldehyde-crosslinked-chromatin immunoprecipitation. *Trends Biochem. Sci.* **25**: 99–104.
- Penfield, S., Gilday, A.D., Halliday, K.J., and Graham, I.A. (2006). DELLA-mediated cotyledon expansion breaks coat-imposed seed dormancy. *Curr. Biol.* **16**: 2366–2370.
- Penfield, S., Josse, E.M., Kannangara, R., Gilday, A.D., Halliday, K.J., and Graham, I.A. (2005). Cold and light control seed germination through the bHLH transcription factor SPATULA. *Curr. Biol.* **15**: 1998–2006.
- Peng, J., Carol, P., Richards, D.E., King, K.E., Cowling, R.J., Murphy, G.P., and Harberd, N.P. (1997). The *Arabidopsis* GAI gene defines a signalling pathway that negatively regulates gibberellin responses. *Genes Dev.* **11**: 3194–3205.
- Peng, J., et al. (1999). 'Green revolution' genes encode mutant gibberellin response modulators. *Nature* **400**: 256–261.
- Perazza, D., Vachon, G., and Herzog, M. (1998). Gibberellins promote trichome formation by up-regulating *GLABROUS1* in *Arabidopsis*. *Plant Physiol.* **117**: 375–383.
- Phillips, A.L., Ward, D.A., Uknes, S., Appleford, N.E.J., Lange, T., Huttly, A., Gaskin, P., Graebe, J.E., and Hedden, P. (1995). Isolation and expression of three gibberellin 20-oxidase cDNA clones from *Arabidopsis*. *Plant Physiol.* **108**: 1049–1057.
- Pysh, L.D., Wysocka-Diller, J.W., Camilleri, C., Bouchez, D., and Benfey, P.N. (1999). The GRAS gene family in *Arabidopsis*: Sequence characterization and basic expression analysis of the *SCARECROW-LIKE* genes. *Plant J.* **18**: 111–119.
- Razem, F.A., El-Kereamy, A., Abrams, S.R., and Hill, R.D. (2006). The RNA-binding protein FCA is an abscisic acid receptor. *Nature* **439**: 290–294.
- Roos, M.D., and Hanover, J.A. (2000). Structure of O-linked GlcNAc transferase: Mediator of glycan-dependent signaling. *Biochem. Biophys. Res. Commun.* **271**: 275–280.
- Rubio, V., Shen, Y., Saijo, Y., Liu, Y., Gusmaroli, G., Dinesh-Kumar, S.P., and Deng, X.W. (2005). An alternative tandem affinity purification strategy applied to *Arabidopsis* protein complex isolation. *Plant J.* **41**: 767–778.

- Saika, H., et al.** (2007). Ethylene promotes submergence-induced expression of OsABA8ox1, a gene that encodes ABA 8'-hydroxylase in rice. *Plant Cell Physiol.* **48**: 287–298.
- Sakamoto, T., Kamiya, N., Ueguchi-Tanaka, M., Iwahori, S., and Matsuoka, M.** (2001). KNOX homeodomain protein directly suppresses the expression of a gibberellin biosynthetic gene in the tobacco shoot apical meristem. *Genes Dev.* **15**: 581–590.
- Sasaki, A., Itoh, H., Gomi, K., Ueguchi-Tanaka, M., Ishiyama, K., Kobayashi, M., Jeong, D.-H., An, G., Kitano, J., Ashikari, M., and Matsuoka, M.** (2003). Accumulation of phosphorylated repressor for gibberellin signaling in an F-box mutant. *Science* **299**: 1896–1898.
- Seo, M., et al.** (2006). Regulation of hormone metabolism in Arabidopsis seeds: Phytochrome regulation of abscisic acid metabolism and abscisic acid regulation of gibberellin metabolism. *Plant J.* **48**: 354–366.
- Shimada, A., Ueguchi-Tanaka, M., Sakamoto, T., Fujioka, S., Takatsuto, S., Yoshida, S., Sazuka, T., Ashikari, M., and Matsuoka, M.** (2006). The rice SPINDLY gene functions as a negative regulator of gibberellin signaling by controlling the suppressive function of the DELLA protein, SLR1, and modulating brassinosteroid synthesis. *Plant J.* **48**: 390–402.
- Shimada, Y., Goda, H., Nakamura, A., Takatsuto, S., Fujioka, S., and Yoshida, S.** (2003). Organ-specific expression of brassinosteroid-biosynthetic genes and distribution of endogenous brassinosteroids in Arabidopsis. *Plant Physiol.* **131**: 287–297.
- Shuai, B., Reynaga-Pena, C.G., and Springer, P.S.** (2002). The lateral organ boundaries gene defines a novel, plant-specific gene family. *Plant Physiol.* **129**: 747–761.
- Silverstone, A.L., Ciampaglio, C.N., and Sun, T.-p.** (1998). The Arabidopsis *RGA* gene encodes a transcriptional regulator repressing the gibberellin signal transduction pathway. *Plant Cell* **10**: 155–169.
- Silverstone, A.L., Jung, H.-S., Dill, A., Kawaide, H., Kamiya, Y., and Sun, T.-p.** (2001). Repressing a repressor: Gibberellin-induced rapid reduction of the RGA protein in Arabidopsis. *Plant Cell* **13**: 1555–1566.
- Silverstone, A.L., Tseng, T.-S., Swain, S., Dill, A., Jeong, S.Y., Olszewski, N.E., and Sun, T.-p.** (2007). Functional analysis of SPINDLY in gibberellin signaling in Arabidopsis. *Plant Physiol.* **143**: 987–1000.
- Stone, S.L., Hauksdottir, H., Troy, A., Herschleb, J., Kraft, E., and Callis, J.** (2005). Functional analysis of the RING-type ubiquitin ligase family of Arabidopsis. *Plant Physiol.* **137**: 13–30.
- Sun, T.-p., and Gubler, F.** (2004). Molecular mechanism of gibberellin signaling in plants. *Annu. Rev. Plant Biol.* **55**: 197–223.
- Sun, T.-p., and Kamiya, Y.** (1994). The Arabidopsis *GA1* locus encodes the cyclase *ent*-kaurene synthetase A of gibberellin biosynthesis. *Plant Cell* **6**: 1509–1518.
- Swain, S.M., and Singh, D.P.** (2005). Tall tales from sly dwarves: novel functions of gibberellins in plant development. *Trends Plant Sci.* **10**: 123–129.
- Swain, S.M., Tseng, T.-s., Thornton, T.M., Gopalraj, M., and Olszewski, N.** (2002). SPINDLY is a nuclear-localized repressor of gibberellin signal transduction expressed throughout the plant. *Plant Physiol.* **129**: 605–615.
- Thomas, S.G., and Sun, T.-p.** (2004). Update on gibberellin signaling. A tale of the tall and the short. *Plant Physiol.* **135**: 668–676.
- Thornton, T.M., Swain, S.M., and Olszewski, N.E.** (1999). Gibberellin signal transduction presents...the SPY who O-GlcNAc'd me. *Trends Plant Sci.* **4**: 424–428.
- Toledo-Ortiz, G., Huq, E., and Quail, P.H.** (2003). The Arabidopsis basic/helix-loop-helix transcription factor family. *Plant Cell* **15**: 1749–1770.
- Tyler, L.** (2006). An Analysis of Potential Negative and Positive Components of the Gibberellin Signaling Pathway in *Arabidopsis thaliana*. PhD dissertation (Durham, NC: Duke University).
- Tyler, L., Thomas, S.G., Hu, J., Dill, A., Alonso, J.M., Ecker, J.R., and Sun, T.-p.** (2004). DELLA proteins and gibberellin-regulated seed germination and floral development in Arabidopsis. *Plant Physiol.* **135**: 1008–1019.
- Ueguchi-Tanaka, M., Ashikari, M., Nakajima, M., Itoh, H., Kato, E., Kobayashi, M., Chow, T.Y., Hsing, Y.I., Kitano, H., Yamaguchi, I., and Matsuoka, M.** (2005). *GIBBERELLIN INSENSITIVE DWARF1* encodes a soluble receptor for gibberellin. *Nature* **437**: 693–698.
- Wang, H., Tang, W., Zhu, C., and Perry, S.E.** (2002). A chromatin immunoprecipitation (ChIP) approach to isolate genes regulated by AGL15, a MADS domain protein that preferentially accumulates in embryos. *Plant J.* **32**: 831–843.
- Weijers, D., Benkova, E., Jager, K.E., Schlereth, A., Hamann, T., Kientz, M., Wilmoth, J.C., Reed, J.W., and Jurgens, G.** (2005). Developmental specificity of auxin response by pairs of ARF and Aux/IAA transcriptional regulators. *EMBO J.* **24**: 1874–1885.
- Wells, L., Vosseller, K., and Hart, G.W.** (2001). Glycosylation of nucleocytoplasmic proteins: Signal transduction and O-GlcNAc. *Science* **291**: 2376–2378.
- Wen, C.-K., and Chang, C.** (2002). Arabidopsis *RGL1* encodes a negative regulator of gibberellin responses. *Plant Cell* **14**: 87–100.
- Wigge, P.A., Kim, M.C., Jaeger, K.E., Busch, W., Schmid, M., Lohmann, J.U., and Weigel, D.** (2005). Integration of spatial and temporal information during floral induction in Arabidopsis. *Science* **309**: 1056–1059.
- Willige, B.C., Ghosh, S., Nill, C., Zourelidou, M., Dohmann, E.M.N., Maier, A., and Schwechheimer, C.** (2007). The DELLA domain of GA INSENSITIVE mediates the interaction with the GA INSENSITIVE DWARF1A gibberellin receptor of Arabidopsis. *Plant Cell* **19**: 1209–1220.
- Wilson, R.N., Heckman, J.W., and Somerville, C.R.** (1992). Gibberellin is required for flowering in *Arabidopsis thaliana* under short days. *Plant Physiol.* **100**: 403–408.
- Xie, Z., Zhang, Z.L., Zou, X., Yang, G., Komatsu, S., and Shen, Q.J.** (2006). Interactions of two abscisic-acid induced WRKY genes in repressing gibberellin signaling in aleurone cells. *Plant J.* **46**: 231–242.
- Yamaguchi-Shinozaki, K., and Shinozaki, K.** (1994). A novel cis-acting element in an Arabidopsis gene is involved in responsiveness to drought, low-temperature, or high-salt stress. *Plant Cell* **6**: 251–264.
- Yu, H., Ito, T., Zhao, Y., Peng, J., Kumar, P., and Meyerowitz, E.M.** (2004). Floral homeotic genes are targets of gibberellin signaling in flower development. *Proc. Natl. Acad. Sci. USA* **101**: 7827–7832.
- Zentella, R., Yamauchi, D., and Ho, T.H.D.** (2002). Molecular dissection of the gibberellin/abscisic acid signaling pathways by transiently expressed RNA interference in barley aleurone cells. *Plant Cell* **14**: 2289–2301.
- Zhang, F., Gonzalez, A., Zhao, M.Z., Payne, C.T., and Lloyd, A.** (2003). A network of redundant bHLH proteins functions in all TGT1-dependent pathways of Arabidopsis. *Development* **130**: 4859–4869.
- Zhang, Z.L., Xie, Z., Zou, X., Casaretto, J., Ho, T.H., and Shen, Q.J.** (2004). A rice WRKY gene encodes a transcriptional repressor of the gibberellin signaling pathway in aleurone cells. *Plant Physiol.* **134**: 1500–1513.

**Global Analysis of DELLA Direct Targets in Early Gibberellin Signaling in *Arabidopsis***  
Rodolfo Zentella, Zhong-Lin Zhang, Mehea Park, Stephen G. Thomas, Akira Endo, Kohji Murase,  
Christine M. Fleet, Yusuke Jikumaru, Eiji Nambara, Yuji Kamiya and Tai-ping Sun  
*Plant Cell* 2007;19;3037-3057; originally published online October 12, 2007;  
DOI 10.1105/tpc.107.054999

This information is current as of November 25, 2020

<b>Supplemental Data</b>	<a href="/content/suppl/2007/10/12/tpc.107.054999.DC1.html">/content/suppl/2007/10/12/tpc.107.054999.DC1.html</a>
<b>References</b>	This article cites 125 articles, 64 of which can be accessed free at: <a href="/content/19/10/3037.full.html#ref-list-1">/content/19/10/3037.full.html#ref-list-1</a>
<b>Permissions</b>	<a href="https://www.copyright.com/ccc/openurl.do?sid=pd_hw1532298X&amp;issn=1532298X&amp;WT.mc_id=pd_hw1532298X">https://www.copyright.com/ccc/openurl.do?sid=pd_hw1532298X&amp;issn=1532298X&amp;WT.mc_id=pd_hw1532298X</a>
<b>eTOCs</b>	Sign up for eTOCs at: <a href="http://www.plantcell.org/cgi/alerts/ctmain">http://www.plantcell.org/cgi/alerts/ctmain</a>
<b>CiteTrack Alerts</b>	Sign up for CiteTrack Alerts at: <a href="http://www.plantcell.org/cgi/alerts/ctmain">http://www.plantcell.org/cgi/alerts/ctmain</a>
<b>Subscription Information</b>	Subscription Information for <i>The Plant Cell</i> and <i>Plant Physiology</i> is available at: <a href="http://www.aspb.org/publications/subscriptions.cfm">http://www.aspb.org/publications/subscriptions.cfm</a>