

# The Arabidopsis *SLEEPY1* Gene Encodes a Putative F-Box Subunit of an SCF E3 Ubiquitin Ligase<sup>W</sup>

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**The Arabidopsis *SLEEPY1* (SLEEPY1) gene positively regulates gibberellin (GA) signaling. Positional cloning of *SLEEPY1* revealed that it encodes a putative F-box protein. This result suggests that *SLEEPY1* is the F-box subunit of an SCF E3 ubiquitin ligase that regulates GA responses. The DELLA domain protein RGA (repressor of *ga1-3*) is a repressor of GA response that appears to undergo GA-stimulated protein degradation. RGA is a potential substrate of *SLEEPY1*, because *sly1* mutations cause a significant increase in RGA protein accumulation even after GA treatment. This result suggests SCF<sup>SLEEPY1</sup>-targeted degradation of RGA through the 26S proteasome pathway. Further support for this model is provided by the observation that an *rga* null allele partially suppresses the *sly1-10* mutant phenotype. The predicted *SLEEPY1* amino acid sequence is highly conserved among plants, indicating a key role in GA response.**

## INTRODUCTION

Bioactive gibberellins (GAs) are tetracyclic diterpenoid phytohormones required for germination, stem elongation, induction of flowering, and fertility (Richards et al., 2001). Severe mutants defective in GA biosynthesis (e.g., the deletion of the *GA1* gene in *ga1-3*) display failure to germinate, dwarfism, delayed flowering, and reduced fertility. All of these phenotypes are rescued by GA treatment. By contrast, mutants impaired in GA signaling display similar phenotypes but are not rescued by GA.

Elements of the GA response pathway in Arabidopsis have been defined through the genetic analysis of mutants (Richards et al., 2001; Olszewski et al., 2002). Positive regulators of GA signaling include *SLEEPY1* (*SLEEPY1*) and *PKL* (*PICKLE*). *PKL* encodes a putative chromatin-remodeling factor, CHD3 (Ogas et al., 1999). Recessive mutations in this gene result in adult plants in which the primary root meristem retains embryonic characteristics. This phenotype is enhanced by GA biosynthetic inhibitors.

Negative regulators of GA signaling include *SHI*, *SPY*, *RGA*, *GAI*, *RGL1*, and *RGL2*. Overexpression of the *SHI* (*SHORT INTERNODES*) gene results in a GA-insensitive semidwarf phenotype. The predicted *SHI* gene product is homologous with the RING-finger domain that mediates protein–protein interaction in ubiquitylation and transcription (Fridborg et al., 1999, 2001). Loss of *SPY* (*SPINDLY*) function results in a GA-overdose phenotype that includes increased internode length, parthenocarpy, and

increased resistance to the GA biosynthetic inhibitor paclobutrazol during vegetative growth and germination (Jacobsen and Olszewski, 1993). *SPY* encodes an O-GlcNAc transferase (OGT) (Jacobsen et al., 1996; Swain et al., 2002). OGTs may regulate the target protein function by competing with protein kinases for modification of phosphorylation sites. *RGA* (*REPRESSOR OF GA1-3*) and *GAI* (*GA-INSENSITIVE*) encode members of the DELLA (VHID) domain subfamily of the GRAS family of putative transcription regulators (Richards et al., 2001; Olszewski et al., 2002; Peng and Harberd, 2002). Loss of *RGA* or *GAI* function results in decreased sensitivity to the GA biosynthetic inhibitor paclobutrazol during vegetative growth. Conversely, mutations in the DELLA domain of *GAI* and *RGA* result in a gain-of-function (semidominant) semidwarf phenotype (Peng et al., 1997; Dill et al., 2001). *RGA* and *GAI* share 83% amino acid identity and act as GA-repressible repressors of stem elongation in Arabidopsis (Dill and Sun, 2001; King et al., 2001). Recent evidence shows that RGA protein accumulation decreases in response to GA treatment. This finding suggests that RGA is subject to GA-induced proteolysis (Silverstone et al., 2001). Other members of the DELLA gene family in Arabidopsis include *RGL1*, *RGL2*, and *RGL3* (*RGA-LIKE*) (Sanchez-Fernandez et al., 1998). *RGL2* is a negative regulator of germination whose transcript levels are increased transiently during dormant seed imbibition (Lee et al., 2002). *RGL1* appears to be a negative regulator of more diverse GA responses, including germination, stem elongation, leaf expansion, flowering, and flower development (Wen and Chang, 2002).

The DELLA family of GA response genes is a highly conserved gene family of considerable agronomic importance (Peng et al., 1999). Introduction of DELLA domain semidwarf mutations into crop plants resulted in the 16 to 31% increase in yield referred to as the Green Revolution (Allan, 1986; Peng et al., 1999). These genes are negative regulators of GA response.

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Loss of function leads to increased GA signaling, whereas gain of function results in reduced GA signaling and dwarfism. Although there are five members of the DELLA family in Arabidopsis, there is only a single DELLA gene in rice (*SLR1* [*SLENDER-RICE*]) and in barley (*SLN1* [*SLENDER*]). Like RGA, both *SLR1* and *SLN1* apparently are subject to GA-regulated proteolysis (Chandler et al., 2002; Gubler et al., 2002; Itoh et al., 2002). Recently, proteolysis of *SLN1* was shown to depend on the 26S proteasome, indicating a role for ubiquitin in GA signal transduction (Fu et al., 2002).

The *sly1* mutants were isolated as recessive GA-insensitive dwarf mutants in two independent screens based on their increased seed dormancy, a property expected in a GA response mutant. The first screen recovered the ethyl methanesulfonate-induced *sly1-2* allele that suppressed the ability of *abi-1-1* (*abscisic acid-insensitive*) to germinate on 3  $\mu$ M abscisic acid. The second screen identified *sly1-10* based on brassinosteroid-dependent germination (Steber et al., 1998; Steber and McCourt, 2001). Loss of *SLY1* function results in all of the phenotypes expected of a GA response mutant, including increased seed dormancy, growth as a dark green dwarf, delayed flowering, and reduced fertility.

Here, we report map-based cloning of the *SLY1* gene, a putative F-box subunit of an SCF E3 ubiquitin ligase. A BLAST (Basic Local Alignment Search Tool) search revealed *SLY1* homologs in many plant species, suggesting that its role as a positive regulator of GA response also is conserved. In addition, mutations in *SLY1* resulted in high-level RGA protein accumulation even in the presence of GA. This result indicates that the *SLY1* gene is needed for the GA-stimulated proteolysis of RGA. The dwarf phenotype of *sly1* plants is suppressed by the *rga-24* null mutation. Thus, accumulation of high levels of RGA is required for expression of the dwarf phenotype of *sly1* mutants. These results suggest that an SCF<sup>*SLY1*</sup> complex mediates the GA-induced degradation of RGA.

## RESULTS

### Effect of the *sly1-10* Mutant on *GA3ox1* Expression

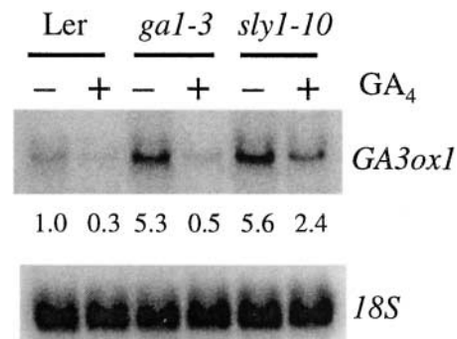
GA biosynthesis is subject to feedback regulation. Decreased GA biosynthesis or response results in increased mRNA accumulation for most of the late GA biosynthetic genes that encode GA-20 oxidase (*GA20ox*) and GA-3 oxidase (*GA3ox*), whereas the application of exogenous GA results in decreased expression of the same GA biosynthetic genes (reviewed by Hedden and Phillips, 2000). Because mutations in *SLY1* result in reduced GA response, we would expect increased expression of these GA biosynthetic genes in the *sly1-10* mutant. To test this hypothesis, we compared the transcript accumulation of one of the GA-3 oxidase genes in Arabidopsis, *GA3ox1*, in the wild type, *ga1-3*, and *sly1-10* (Figure 1). As shown previously, in the absence of GA, *GA3ox1* transcript levels were fivefold higher in the severe GA biosynthetic mutant *ga1-3* than in the wild type (Chiang et al., 1995; Cowling et al., 1998; Yamaguchi et al., 1998). Similar to *ga1-3*, the *sly1-10* mutant accumulated fivefold greater *GA3ox1* mRNA levels compared with the wild type in the absence of exogenous GA. Application

of GA<sub>4</sub> resulted in a 10-fold decrease in *GA3ox1* expression in *ga1-3* but only a 2-fold decrease in *sly1-10*. This result indicates that the *sly1-10* mutant retains some residual sensitivity to GA<sub>4</sub>.

### Map-Based Cloning of *SLY1*

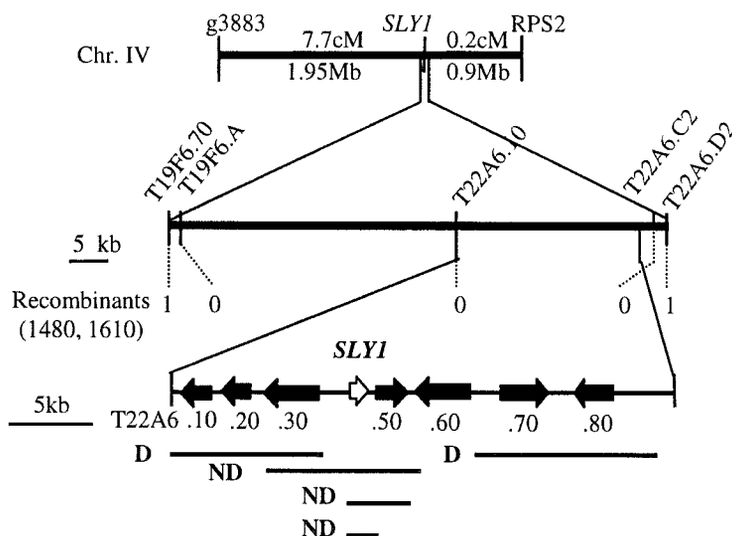
The *SLY1* gene was cloned using a map-based approach to further elucidate its function in GA signal transduction. The *sly1-2/sly1-2* mutant in the Landsberg *erecta* (*Ler*) ecotype was crossed to wild-type ecotype Columbia to generate an F<sub>2</sub> mapping population of *sly1-2/sly1-2* plants segregating for *Ler*- and Columbia-specific physical markers. *sly1-2/sly1-2* F<sub>2</sub> seeds were germinated by cutting the seed coats. As a result of the poor germination and poor fertility of *sly1-2*, all mapping was performed relative to PCR-based physical markers that differ between the two ecotypes (Konieczny and Ausubel, 1993; Bell and Ecker, 1994). *SLY1* was 7.7 centimorgan (cM) from g3883 and 0.2 cM from *RPS2* (Figure 2). *RPS2* was used as a point from which to begin a chromosome walk toward g3883. A population of 805 *sly1-2/sly1-2* F<sub>2</sub> plants (1610 chromosomes) was scored for the *RPS2* cleaved amplified polymorphic sequence marker (Bent et al., 1994). Four F<sub>2</sub> plants heterozygous for the *RPS2* marker were recovered and used as a basis for a chromosome walk of 0.9 Mb. The recombination rate across this region was very low, with a ratio of 4500 kb/cM compared with the Arabidopsis average of 200 kb/cM (Schmidt et al., 1995). *SLY1* was localized to a 70-kb region between markers T19F6.7 and T22A6.D2.

There were 20 predicted genes in the 70-kb region containing *SLY1* (Arabidopsis Genome Initiative, 2000). Transformation of *sly1-2* and *sly1-10* plants with BAC subclones revealed that an 11.7-kb subclone, T22A6.2G10, rescued the dwarf (data not shown) and germination (Figure 3A) phenotypes of *sly1* mutants. This subclone contains three predicted open reading



**Figure 1.** Effect of *sly1* Mutations on *GA3ox1* Transcript.

An RNA gel blot, containing 10  $\mu$ g of total RNA isolated from 8-day-old *Ler* (wild-type), *ga1-3*, and *sly1-10* seedlings treated with (+) or without (-) 1  $\mu$ M GA<sub>4</sub> for 2 h, was hybridized with a labeled *AtGA3ox1* antisense RNA probe. The blot was reprobbed with a labeled 18S oligonucleotide probe. Numerals below the blot indicate the relative levels of *AtGA3ox1* mRNA after standardization using the 18S RNA as a loading control. The level of *AtGA3ox1* mRNA in the water-treated wild type was arbitrarily set to 1.0.



**Figure 2.** Map-Based Cloning of *SLY1*.

Fine mapping delineated a 70-kb region containing *SLY1* between markers T19F6.70 and T22A6.D2. Single recombination events were identified at T22A6.D2 (in 1610 chromosomes) and at T19F6.70 (in 1480 chromosomes). Overlapping BAC clones T22A6 and T19F6 cover this region (Arabidopsis Genome Initiative, 2000). Transformation with subclones of these BACs identified an 11.7-kb complementing subclone, T22A6.2G10. Transformation with T22A6.40 (from  $-1347$  to  $+666$  relative to the ATG) rescued *sly1*. D, dwarf; ND, nondwarf.

frames (ORFs): T22A6.30 (At4g24200), T22A6.40 (At4g24210), and T22A6.50 (At4g24220). Transformation with the T22A6.40 ORF alone rescued the *sly1-10* mutant phenotypes (Figure 3B). DNA sequence analysis of *sly1-2* and *sly1-10* revealed that these alleles contain mutations within this 453-bp ORF (Figure 4A). *sly1-2* has a 2-bp deletion (Cys-337 and Thr-338) that causes a frameshift that eliminates the last 40 amino acids. *sly1-10* contains a 23-bp deletion (amino acids 433 to 456) followed by an  $\sim 8$ -kb insertion. This causes the loss of only the last 8 amino acids and the addition of 46 random amino acids. Thus, T22A6.40 (At4g24210) is the *SLY1* gene. *SLY1* contains no introns and encodes a small predicted protein of 151 amino acids containing a putative F-box domain.

### Expression of *SLY1*

Sequencing of several full-length *SLY1* cDNAs identified 5' untranslated regions (UTRs) of 87 and 105 nucleotides (Figure 4A) (<http://signal.salk.edu/>). The major ORF in this transcript is the *SLY1* gene. The long 380-nucleotide 3' UTR shows a high probability of secondary structure (99.6 kcal/mol) according to Vienna RNA secondary structure prediction (<http://www.tbi.univie.ac.at/cgi-bin/RNAfold.cgi>). The Arabidopsis F-box gene *TIR1* mRNA also contains a long structured 3' UTR. Structured 3' UTRs have been implicated in the control of mRNA stability and localization (Decker and Parker, 1995). Reverse transcriptase-mediated (RT) PCR analysis of wild-type *Ler* plants showed that the *SLY1* transcript is present in all tissues examined (Figure 4B), including rosette leaves, green siliques, flowers, stems, cauline leaves, and seedlings. The *SLY1* mRNA was not detected in the *sly1-10* mutant, as a result of an 8-kb insertion within the PCR product. However, a band of 9 kb was de-

tected in *sly1-10* but not in wild-type *Ler* by RNA gel blot analysis, indicating that the gene is still expressed (data not shown).

### Analysis of the *SLY1* Gene Structure

A BLAST search revealed that the predicted *SLY1* protein sequence is homologous with the cyclin F-box family of proteins (Figure 5A) (Altschul et al., 1990; Zhang et al., 1998; Gagne et al., 2002). The F-box protein is one of the four main subunits of the SCF (SKP1, Cullin, F-box) complex, one type of E3 ubiquitin ligase (Conaway et al., 2002). The F-box subunit directs the interaction of the complex with a specific target for ubiquitylation. Thus, *SLY1* may transduce the GA signal by regulating the stability of GA response proteins through ubiquitin-mediated proteolysis.

SCF-mediated proteolysis regulates many aspects of plant development, including senescence, flower development, circadian rhythm, light receptor signaling, and hormone signaling (Hellmann and Estelle, 2002). The SCF<sup>TIR1</sup> and SCF<sup>COI1</sup> complexes are needed for auxin and jasmonate signal transduction, respectively (Gray et al., 2001; Xu et al., 2002). SCF<sup>TIR1</sup> controls AXR2/IAA7 stability and is the only SCF in Arabidopsis for which a target has been identified to date (Gray et al., 2001). The predicted TIR1 and COI1 F-box proteins share 34% amino acid identity with each other and contain an F-box at the N terminus and a Leu-rich repeat domain at the C terminus. The predicted *SLY1* protein is not homologous with these proteins outside of the F-box motif. F-box proteins often contain a C-terminal protein-protein interaction domain, such as Leu-rich repeats, WD40 repeats, or Kelch repeats, for interaction with its target protein (del Pozo and Estelle, 2000). Although *SLY1* lacks such a conserved protein-protein interaction domain, its



**Figure 3.** Complementation of *sly1* Mutants.

**(A)** GA dose response in germination. Transformation with the T22A6.2G10 subclone rescues the GA-insensitive germination phenotype of *sly1-2*. Percentage germination of dormant *sly1-2* seeds (open triangles), the GA biosynthesis mutant *ga1-3* (open squares), and *sly1-2* transformed with T22A6.2G10 (closed triangles) is shown. Wild-type *Ler* germination was identical to that of T22A6.2G10-transformed *sly1-2*. Error bars represent standard errors for triplicate samples of 50 to 100 seeds.

C terminus clearly is important for function. Both *sly1-2* and *sly1-10* alleles leave the F-box intact but alter the C terminus (Figure 4A). Other examples of F-box proteins lacking a conserved C-terminal protein-protein interaction domain include the mammalian gene *Fbx8* and the Arabidopsis gene *SON1* (Cenciarelli et al., 1999; Ilyin et al., 2000; Kim and Delaney, 2002).

*SLY1* is highly conserved in the plant kingdom. Many members of the *SLY1* gene family were identified by tBLASTn search of plant ESTs (Figure 5B). Although homologs were detected in many plant species, none were detected outside of the plant kingdom. Amino acid relatedness with the predicted *SLY1* protein ranged from 57% identical/67% similar in the dicot soybean to 42% identical/57% similar in the monocot barley. The rice ortholog of *SLY1* was identified independently by Sasaki et al. (2003) by positional cloning of *GID2* (*GA-INSENSITIVE DWARF*). *GID2* encodes a predicted 212-amino acid protein that is 43% identical and 56% similar to *SLY1*. The regions of highest amino acid identity are the F-box and the 37-amino acid GGF region (Figure 5B). Two variable regions (Figure 5B, VR) are absent in *AtSLY1*. A BLAST search detected a single *SLY1* homolog (154 amino acids, 26% identical) in Arabidopsis, MIF21.6 (At5g48170), on chromosome V (Figure 5B).

#### Accumulation of RGA Is Increased in *sly1* Mutants

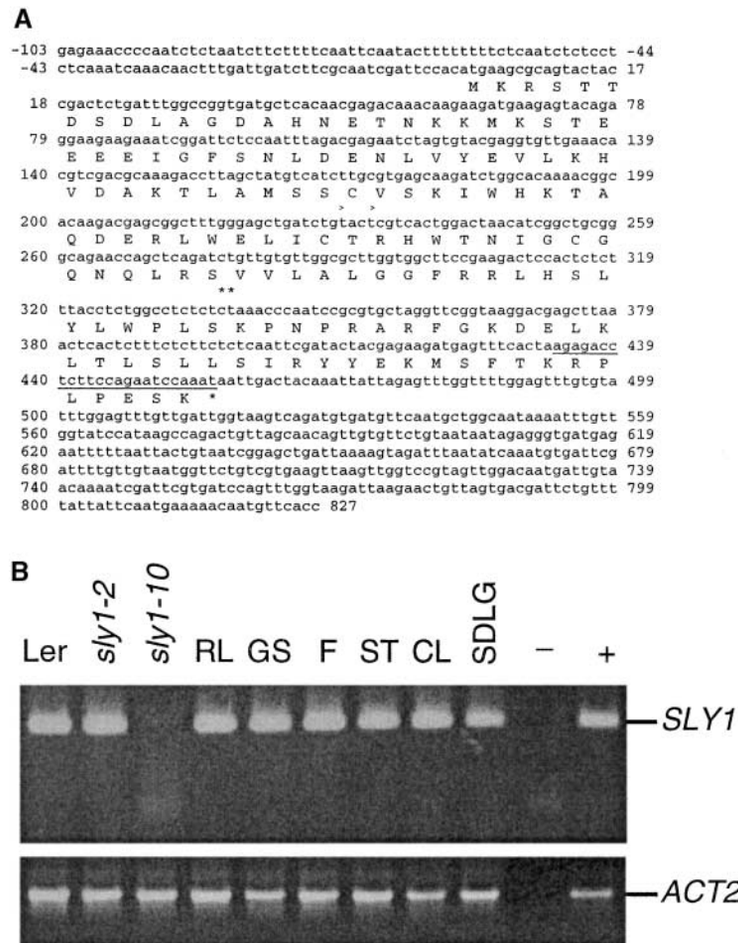
The fact that GA causes reduced accumulation of RGA (Silverstone et al., 2001) prompted the question, Is RGA a target of SCF<sup>*SLY1*</sup> ubiquitin ligase? To explore this hypothesis, we examined the stability of RGA in *sly1-2* and *sly1-10*. RGA accumulates at a much higher level in both *sly* alleles than in the wild type, even after GA treatment (Figure 6A). By contrast, the *RGA* mRNA level was not altered dramatically in the *sly1-10* mutant (Figure 6B). These results show that *SLY1* is directly or indirectly required for the GA-stimulated degradation of RGA. Given that *SLY1* contains an F-box domain, we speculate that RGA may be a target of an SCF<sup>*SLY1*</sup> complex.

#### Analysis of the *sly1-10 rga-24* Double Mutant

If the dwarf phenotype of *sly1* mutants is attributable to the overaccumulation of RGA, we would expect loss-of-function mutations in the *RGA* gene to rescue the *sly1* dwarf phenotype. The *sly1-10* allele was crossed to *rga-24* and used to generate a *sly1-10 rga-24* double mutant. Figure 3C shows a comparison

**(B)** Transformation with T22A6.40 (−1347 to +666) rescues the dwarf phenotype of *sly1-2*. Wild-type *Ler* (left), *sly1-2*+ T22A6.40 (center), and *sly1-2* (right) plants are shown. Bar = 1 cm.

**(C)** Suppression of *sly1-10* by *rga-24*. *sly1-10* (left), the *sly1-10 rga-24* double mutant (center), and wild-type *Ler* (right) plants are shown. *rga-24* partly rescues the dwarf phenotype of *sly1-10*, but not poor fertility. *Ler* and homozygous mutant plants were grown on soil for 60 days under a long-day photoperiod. Bar = 15 mm.



**Figure 4.** *SLY1* mRNA.

**(A)** Sequence of full-length *SLY1* cDNA. The *SLY1* gene contains no introns and predicts a protein of 151 amino acids. The first ATG (+1) in the transcript is the *SLY1* translational start site. *sly1-2* contains a 2-bp deletion (double asterisks). *sly1-10* contains a 23-bp deletion (underlined) and an 8-kb insertion.

**(B)** RT-PCR analysis of *SLY1* mRNA accumulation. An ethidium bromide-stained 2% agarose gel from RT-PCR using 100 ng of total RNA for each sample is shown. *SLY1* (250 bp) plus *ACT2* (471 bp) accumulation are shown for wild-type *Ler*, *sly1-2*, and *sly1-10* whole aerial plants and for wild-type *Ler* rosette leaves (RL), green siliques (GS), flowers (F), stems (ST), cauline leaves (CL), and seedlings (SDLG). For seedlings, tissue was harvested from 4-week-old plants. –, no RNA template; +, genomic DNA.

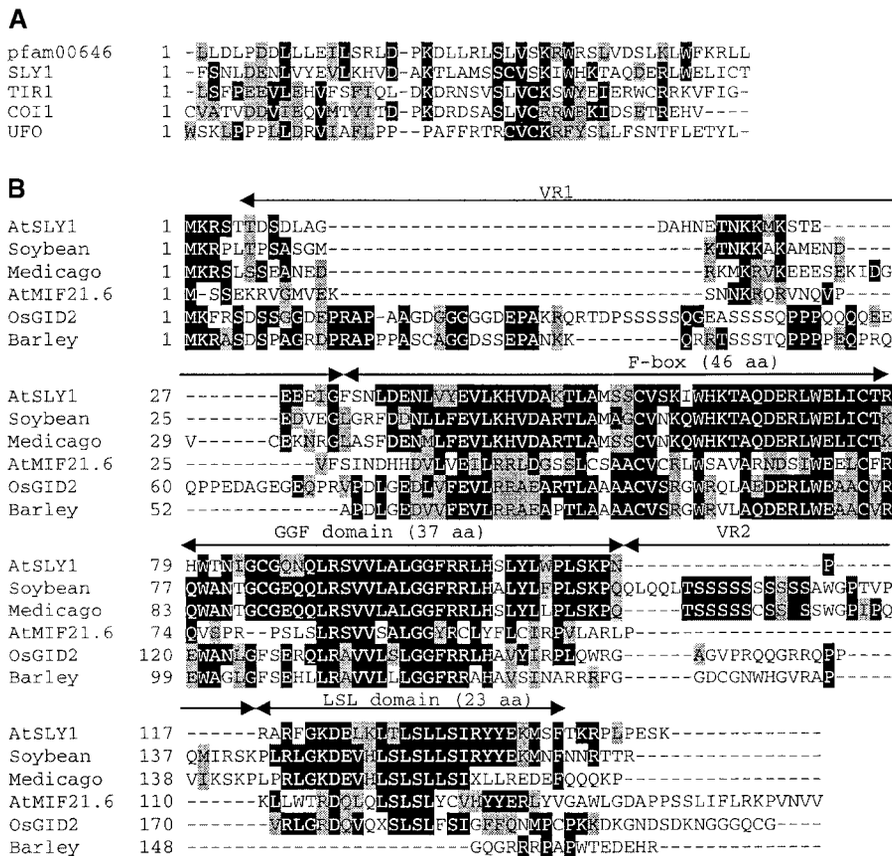
of *sly1-10*, the *sly1-10 rga-24* double mutant, and *Ler* after 60 days of growth on soil. The phenotype of the *rga-24* mutant was almost identical to that of *Ler* (Silverstone et al., 1997). The *rga-24* mutation clearly resulted in a partial rescue of the *sly1-10* dwarf phenotype but did not significantly suppress the germination or fertility defects of *sly1-10*. This result indicates that *rga-24* is epistatic to *sly1-10* for stem elongation and that *RGA* acts downstream of *SLY1* in GA signal transduction.

## DISCUSSION

We report the map-based cloning of the *SLY1* gene of *Arabidopsis*. *SLY1* is a positive regulator of GA response. Recessive mutations in *SLY1* affect the full range of GA phenotypes, including feedback regulation of the *GA3ox1* biosynthetic gene (Figure 1). Thus, the fact that *SLY1* encodes a putative F-box protein sug-

gests that the GA signal is transmitted via an SCF<sup>*SLY1*</sup> E3 ubiquitin ligase.

Ubiquitylation controls target protein activity at multiple levels, including proteolysis and the potentiation of transcriptional activation domains (Conaway et al., 2002). Major members of the SCF complex include homologs of SKP1, cullin, and the RING-finger domain protein Rbx1 (Zheng et al., 2002). The F-box subunit directs the interaction of the complex with a specific target for ubiquitylation. The conserved F-box domain allows the protein to interact with the SKP1 subunit of the SCF. SKP1 tethers the F-box protein to the N terminus of cullin. The RING-finger protein Rbx1 binds the C terminus of cullin and the E2 ubiquitin-conjugating enzyme. The E3 ubiquitin ligase catalyzes the transfer of ubiquitin from E2 to the substrate. The formation of a polyubiquitin chain on the substrate targets it for proteolysis by the 26S proteasome.



**Figure 5.** SLY1 Sequence Alignments.

**(A)** Alignment of the SLY1 F-box with National Center for Biotechnology Information consensus Pfam00646 (an F-box protein family) and other Arabidopsis F-box proteins.

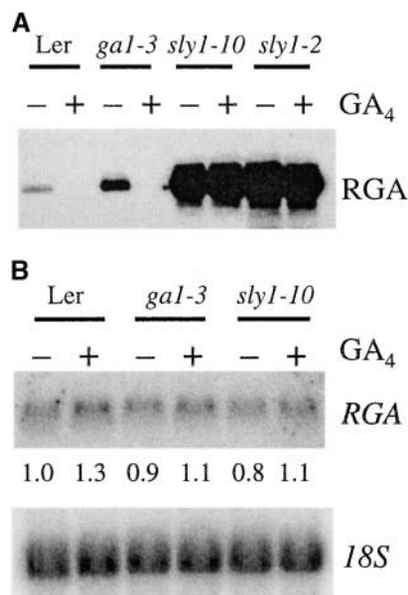
**(B)** Alignment of SLY1 with plant homologs from soybean, *Medicago truncatula*, Arabidopsis MIF21.6, rice OsGID2, and barley. The amino acid sequence predicted from the largest open reading frame in each EST is listed. GGF and LSL refer to conserved residues. aa, amino acids; VR, variable region.

In Arabidopsis, there are at least 21 Skp1 homologs (ASK [Arabidopsis Skp1]), 11 cullin homologs, and 2 Rbx homologs (Farras et al., 2001; Lechner et al., 2002; Shen et al., 2002). Recent work has characterized the superfamily of 694 putative F-box protein genes in Arabidopsis (Gagne et al., 2002; Kuroda et al., 2002). According to Gagne et al. (2002), the *SLY1* gene falls into the C2 family of F-box proteins. Representatives of the Arabidopsis F-box family, including C2, were found to interact with Arabidopsis SKP1 homologs using yeast two-hybrid assays, indicating that they are part of functional SCF complexes. A representative of the C2 F-box family was shown to interact with ASK1, ASK2, ASK4, ASK11, and ASK13 (Gagne et al., 2002). SCF complexes in Arabidopsis may be regulated by self-ubiquitylation of the F-box or by RUB modification of the cullin subunit (Conaway et al., 2002; del Pozo et al., 2002). It was shown recently that RUB1 modification of AtCUL1 is required for normal auxin signal transduction. However, the precise role of RUB modification is unknown.

A number of previously published reports support the notion that the GA signal is transmitted by ubiquitylation and proteoly-

sis. Reynolds and Hooley (1992) reported the isolation of a GA-regulated tetraubiquitin cDNA from *Avena fatua*. It is possible that an increased pool of ubiquitin is needed for GA response. Chen et al. (1995) identified a GA-induced cDNA that encodes a ubiquitin-conjugating enzyme in rice. Finally, the negative regulator of GA response, RGA, appears to be subject to GA-stimulated proteolysis (Silverstone et al., 2001). The barley and rice homologs of RGA, SLN1 and SLR1, respectively, also appear to be subject to GA-stimulated proteolysis (Chandler et al., 2002; Gubler et al., 2002; Itoh et al., 2002). This result suggests that this aspect of GA signal transduction is conserved among dicots and monocots. Recently, inhibitor studies suggested that the GA-stimulated proteolysis of barley SLN1 was dependent on the 26S proteasome, suggesting a role for the 26S proteasome in the GA-induced proteolysis of the RGA/SLN1/SLR1 family (Fu et al., 2002).

We propose that GA causes the ubiquitylation of RGA via an SCF<sup>SLY1</sup> E3 ubiquitin ligase. According to this hypothesis, SLY1 acts positively in the GA response because it is the negative regulator of RGA, which is a negative regulator of the GA re-



**Figure 6.** RGA Protein Levels, but Not the *RGA* mRNA, Are Highly Increased in *sly1* Mutants.

**(A)** Eight-day-old seedlings were treated with (+) or without (-) 1  $\mu$ M  $GA_4$  for 2 h, and protein extracts were fractionated by 8% SDS-PAGE. The protein blot was probed with an anti-RGA antibody. Ponceau staining was used to confirm equal loading of the blot (data not shown).

**(B)** A duplicate RNA gel blot (described in Figure 1A) was hybridized with a labeled *RGA* DNA probe. Numerals below the blot indicate the relative levels of *RGA* mRNA after standardization using 18S as a loading control. The level of *RGA* mRNA in the water-treated wild type was arbitrarily set to 1.0.

sponse. Evidence in favor of this model include the following: (1) *sly1* mutants have a GA-insensitive phenotype; (2) the predicted SLY1 protein is homologous with an F-box; (3) RGA accumulates at high levels in *sly1* mutants even in the presence of GA; and (4) the *rga-24* mutation suppresses the dwarf phenotype of *sly1-10*. Figure 6 shows that mutations in *SLY1* result in the high-level accumulation of RGA. Although this evidence does not directly prove an interaction between SLY1 and RGA, it is highly suggestive that an SCF<sup>SLY1</sup> complex may direct the GA-stimulated degradation of RGA. If this hypothesis is correct, SLY1 will be the second F-box protein in plants for which a target has been identified. It is possible that the DELLA motif of RGA is needed for regulation by SCF<sup>SLY1</sup>, because this motif is essential for the GA-induced degradation of RGA (Dill et al., 2001). Like the *sly1-10* mutation, deletion of the DELLA motif stabilizes RGA even in the presence of GA (Dill et al., 2001).

It is tempting to speculate that an SCF<sup>SLY1</sup> complex might target the entire DELLA family, including GAI, RGL1, RGL2, and RGL3, for destruction. However, *RGL2* appears to be regulated in transcription (Lee et al., 2002), and it was shown that RGL1-GFP (green fluorescent protein) and GAI-GFP translational fusion proteins are not subject to GA-regulated proteolysis (Fleck and Harberd, 2002; Wen and Chang, 2002). Although this work has not yet been confirmed by examination of native RGL1 and GAI proteins, it suggests that RGL1 and GAI are not subject to

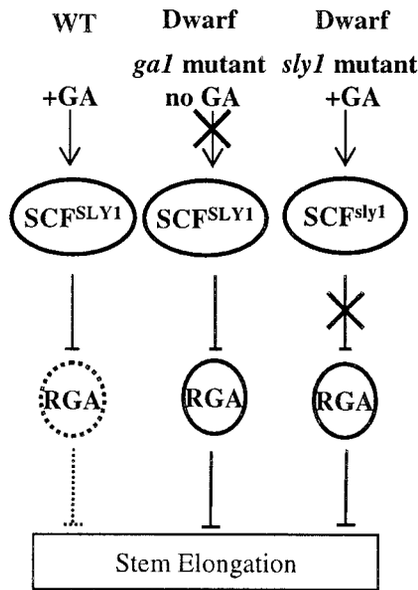
regulation by classic ubiquitin-directed proteolysis. Nevertheless, the *sly1* mutants are defective in the full range of GA responses, whereas *RGA* affects stem elongation, leaf expansion, and flowering time. Thus, the *sly1* phenotypes of increased seed dormancy and reduced fertility likely result from a mechanism other than increased levels of RGA protein. It will be important to determine if *SLY1* regulates other GA response genes, including other members of the DELLA family.

*sly1* mutant phenotypes are not as strong as those of the GA biosynthetic mutant *ga1-3*. The *ga1-3* seeds have an absolute requirement for added GA to germinate. Although *sly1* mutants show increased seed dormancy (5% germinate; Figure 2), they do eventually afterripen and hence germinate (C. Steber, unpublished data). The *ga1-3* mutant has a stronger dwarf phenotype than *sly1* mutants (Steber et al., 1998). Whereas *ga1-3* plants are fully infertile, *sly1* mutants are partially infertile. One possible explanation for this difference may be that the *SLY1* homolog MIF21.6 may be functionally redundant with *SLY1*. This fact also may explain why GA can cause some reduction in *GA3ox1* transcript levels in the *sly1* mutant background. By contrast, GA cannot cause the reduction of RGA protein levels in a *sly1* mutant background. This finding suggests that *SLY1* is involved more directly in the regulation of RGA than in the regulation of *GA3ox1*.

Although the *sly1* dwarf phenotype is not as strong as that of *ga1-3*, the *sly1* mutants result in a considerably higher (approximately fivefold) level of RGA protein accumulation than *ga1-3* (Figure 6). If plant height were a direct function of RGA protein levels, we would expect *sly1* plants to be smaller than *ga1-3* plants. There are two possible explanations for this finding. One is that the plant may compensate for RGA overabundance in *sly1* mutants by downregulating other DELLA family proteins. Another is that the RGA protein that accumulates in *sly1* mutants may not be fully functional.

How does GA control the activity of SCF<sup>SLY1</sup>? Most SCF ubiquitin ligase-regulated proteins are targeted for ubiquitylation and degradation by phosphorylation (Willems et al., 1999). However, there are examples of ubiquitin ligase-regulated proteins being targeted for ubiquitylation and destruction by Pro hydroxylation or by glycosylation (Huang et al., 2002; Yoshida et al., 2002). Phosphorylation was implicated recently in the regulation of the 26S proteasome-mediated proteolysis of the barley DELLA protein SLN1 (Fu et al., 2002). In addition, Sasaki et al. (2003) report evidence that the RGA homolog SLR1 is targeted for degradation by phosphorylation. Given that proteolysis is a conserved mechanism for regulating the DELLA family of proteins in plants, it is reasonable to speculate that RGA also may be regulated by phosphorylation. However, at present, there is no direct evidence of this phenomenon. In this model, SCF<sup>SLY1</sup> would recognize RGA only when it is phosphorylated by a GA-stimulated kinase. Thus, a GA-regulated kinase and/or phosphatase may play a crucial role in GA signaling.

In summary, we propose a model for the role of SLY1 in GA signal transduction (Figure 7). Wild-type plants (+GA) reach normal height because GA stimulates SCF<sup>SLY1</sup> to target RGA for degradation, thus alleviating the RGA inhibition of stem elongation. In GA biosynthetic mutants (no GA), there is insufficient GA to stimulate the SCF<sup>SLY1</sup> complex to target RGA for



**Figure 7.** Model for the Role of *SLY1* in GA Signaling.

The dwarf phenotype of GA biosynthetic mutants results from the accumulation of active RGA protein. In wild-type (WT) plants (+GA), GA stimulates SCF<sup>SLY1</sup> (directly or indirectly) to target RGA for degradation, resulting in normal height. The *gal1* biosynthetic mutant is a dwarf because there is insufficient GA to stimulate SCF<sup>SLY1</sup> to target RGA for proteolysis. The *sly1* mutant is a dwarf because a lack of functional SCF<sup>SLY1</sup> results in increased accumulation of RGA.

proteolysis. The overabundance of RGA inhibits stem elongation, leading to a dwarf phenotype. Mutations in *SLY1* prevent the degradation of RGA in both the presence and absence of GA, leading to RGA inhibition of stem elongation and a dwarf phenotype.

## METHODS

### Materials and Growth Conditions

*Arabidopsis thaliana* ecotypes Landsberg *erecta* and Columbia and BAC clones used in this study were obtained from the ABRC (<http://www.arabidopsis.org/>). Plants were grown under a regimen of 16-h days at 22°C and 8-h nights at 16°C under halide lights at 100 to 150  $\mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ . *sly1* mutants are less fertile if grown under continuous light. Before plating on 0.5 $\times$  Murashige and Skoog (1962) basal salt mixture (Sigma) plus 0.8% (g/v) agar, seeds were surface-sterilized by incubation in 10% (v/v) bleach and 0.01% SDS for 10 min followed by six washes in sterile water. Seeds were allowed to imbibe for 4 days at 4°C to ensure synchronous germination and then moved to continuous fluorescent light at 50  $\mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$  and 22°C. Seeds with emerging radicals were scored as germinated after 7 days under lights. Stock solutions of GA<sub>4</sub> (Sigma or a gift from Tadao Asami, RIKEN, Saitama, Japan) were made in ethanol. Plant hormones were added to autoclaved medium cooled to 55°C.

For the analysis of RGA protein levels and expression of *RGA* and *GA3ox1* transcripts, Arabidopsis seedlings were grown as described previously (Silverstone et al., 2001). To facilitate germination, *ga1-3* seeds were pretreated with 100  $\mu\text{M}$  GA<sub>4</sub> during the stratification period

and then washed five times with water. Treatments with GA<sub>4</sub> or water were performed by adding 1 mL of solution directly to the surface of the agar. Whole seedlings were harvested and ground directly in liquid nitrogen using a pestle and mortar.

### Map-Based Cloning

Simple sequence length polymorphism and cleaved amplified polymorphic sequence (CAPS) markers were used to localize *SLY1* to chromosome IV (Konieczny and Ausubel, 1993; Bell and Ecker, 1994). CAPS markers used to fine map the *SLY1* gene are listed in the supplemental data online. Clones marked "cereon" were based on the Cereon markers at the TAIR World Wide Web site (<http://www.arabidopsis.org/>). All other markers were derived in this study. The germination and dwarf phenotypes of the *sly1* mutant necessitated using PCR-based mapping markers. Nineteen new CAPS markers were defined by screening 56 2-kb PCR products using 4-bp cutters (see supplemental data online). To identify the *SLY1* gene by complementation, sublibraries were generated from BAC clones T19F6 and T22A6 by partial digestion with *Sau3A*I. Inserts ranging from 5 to 20 kb were inserted into the *Bam*HI cloning site in pBIN19. Subclones representing the region were transformed into *sly1-2* or *sly1-10* (Bechtold et al., 1993; Clough and Bent, 1998). Sequence alignments were determined using tBLASTn to search the TIGR EST and GenBank databases (Altschul et al., 1990; Smith et al., 1996; [www.ncbi.nlm.nih.gov](http://www.ncbi.nlm.nih.gov/), 2002; [www.tigr.org](http://www.tigr.org), 2002).

### The *SLY1* Transcript

The *SLY1* cDNA 3' end was determined by sequencing the 3' ends of full-length cDNAs recovered from an Arabidopsis Columbia uni-Zap XR cDNA library (Stratagene, La Jolla, CA) to identify a polyadenylated sequence. The 3' end was amplified (2  $\mu\text{L}$  of cDNA library, 0.2 mM deoxy-nucleotide triphosphates, 1.5 mM MgCl<sub>2</sub>, 0.5 mM primer, 1 $\times$  reaction buffer, and 5 units of Ex-Taq polymerase [Takara Bio, Shiga, Japan] for 40 cycles of 96°C for 25 s, 60°C for 40 s, and 72°C for 80 s) using *sly1-2f* (5'-AGACGAGCGGCTTTGGGAGC-3') for five cycles before the addition of T7 primer (5'-TAATACGACTCACTATAGGG-3'). This PCR product was used as a template for the reaction using 2-63f (5'-TCTCTCTAAACC-CAATCCG-3') for five cycles before the addition of T7 primer. PCR products were gel purified (Qiaex II; Qiagen, Valencia, CA) and cloned using the TOPO-XL PCR cloning kit (Invitrogen, Carlsbad, CA). Clones were sequenced using fluorescence-based dideoxy terminators and Ampli-Taq polymerase and run on an Applied Biosystems sequencer (model 377; Perkin-Elmer Applied Biosystems, Norwalk, CT).

Because of the presence of the MIF21.6 *SLY1* homolog on chromosome V, it was difficult to specifically detect the *SLY1* transcript by RNA gel blot analysis. Thus, qualitative reverse transcriptase-mediated PCR was used to search for the presence or absence of *SLY1* mRNA in various aerial tissues and in *sly1* mutants. Tissue was harvested from 8-week-old Arabidopsis plants or from seedlings (where indicated). Total RNA was extracted from 0.1 g of plant tissue using the RNeasy Plant Mini Prep Kit (Qiagen) according to the manufacturer's instructions with the addition of an RNase-free DNaseI treatment (Roche Diagnostics, Indianapolis, IN). Reverse transcriptase-mediated PCR was conducted on 100 ng of total RNA using a Roche LightCycler and the LightCycler RNA Amplification Kit SYBR Green I (Roche) according to the manufacturer's instructions at an annealing temperature of 56°C with 5 mM MgCl<sub>2</sub>. A no-reverse transcription control was included for all RNA preparations to confirm the absence of genomic DNA contamination. The primers used for *SLY1* amplification were 5'-TCTCTCTAAACCCCAATCCG-3' and 5'-CCAGCATTGAACATCACATCTGAC-3'. The primers used for *ACT2* amplification were 5'-CTGGATTCTGGTGATGGTGATGTC-3' and 5'-TCT-

TTGCTCATACGGTCAGCG-3' (An et al., 1996). The products of amplification were separated on a 2% agarose gel for 3 h at 60 V (Sambrook et al., 1989).

### Immunoblot Analysis

Total protein was extracted from water- or GA<sub>4</sub>-treated seedlings as described previously (Silverstone et al., 2001). For each sample, 20 μg of total protein was fractionated on an 8% SDS-polyacrylamide gel and examined by immunoblot analysis (Silverstone et al., 2001) using a 2000-fold dilution of an anti-RGA antibody from rat and a 7500-fold dilution of peroxidase-coupled goat anti-rat IgG. Immunoreactive species on the blots were detected using Supersignal Dura Reagent (Pierce). Ponceau staining was performed by incubating the blot with 0.2% (w/v) Ponceau S (Sigma) in 1% (v/v) acetic acid.

### RNA Gel Blot Analysis of GA3ox1 and RGA Transcripts

Total RNA was isolated and GA3ox1 mRNA was detected using an antisense GA3ox1 RNA probe as described previously (Yamaguchi et al., 1998). The RGA transcripts were analyzed using a random primed labeled 2.3-kb RGA DNA probe as described (Silverstone et al., 1998). As a loading control, blots were reprobated with a labeled 18S oligonucleotide probe as described previously (Dill and Sun, 2001). The RNA blots were exposed to a Storage Phosphor Screen (Molecular Dynamics, Sunnyvale, CA) and quantified with a PhosphorImager (model 455Si; Molecular Dynamics) using Imagequant version 5.1 software.

### Generation of the sly1-10 rga-24 Mutant

We isolated the *sly1-10 rga-24* homozygous double mutant by crossing *rga-24* and *sly1-10*. The genotypes of homozygous F2 plants were confirmed using allele-specific PCR primers. Genotyping of the *rga-24* allele was performed as described previously (Dill and Sun, 2001). Primers *sly1-10f* (5'-TCGTCCTGGACTAACATCGGCTG-3') and 2-63r (5'-GCTAACAGTCTGGCTTATGGATAC-3') were used to amplify a 350-bp product in *SLY1* but not *sly1-10*. Primers *sly1-10f* and *sly1-10r2* (5'-GAGCATGCTTGATCCATAGGA-3') were used to amplify a 320-bp product in *sly1-10* but not in *SLY1*. PCR was performed as described (Dill and Sun, 2001).

Upon request, all novel materials described in this article will be made available in a timely manner for noncommercial research purposes.

### Accession Numbers

The GenBank accession number for the *SLY1* gene is NM\_118554. Other accession numbers are as follows: Arabidopsis *TIR1*, AF327430; *SLY1* plant homologs from soybean, BI785351; *Medicago truncatula*, BQ239225; Arabidopsis MIF21.6, AB023039; rice *OsGID2*, AB100246; and barley, BF622212.

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**The Arabidopsis *SLEEPY1* Gene Encodes a Putative F-Box Subunit of an SCF E3 Ubiquitin Ligase**

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<b>Supplemental Data</b>	<a href="/content/suppl/2003/05/01/15.5.1120.DC1.html">/content/suppl/2003/05/01/15.5.1120.DC1.html</a>
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