Arabidopsis *GLABROUS7* **Gene Requires Downstream Sequences for Function**

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The Arabidopsis *GLABROUSl (GL1)* **gene is a myb gene homolog required for the initiation of trichome development. In situ hybridiration revealed that the highest levels of** *GL1* **transcripts were present in developing trichomes. In contrast, previous work had shown that putative promoter sequences from the 5' noncoding region of the** *GL1* **gene directed the expression of a B-glucuronidase** *(GUS)* **reporter gene only in stipules. Deletion analysis of the 3' noncoding region of** *GL1* **has identified an enhancer that is essential for** *GL1* **function. Sequences fmm the region containing the enhancer, in conjunction with** *GL1* **upstream sequences, direct the expression of a** *GUS* **reporter gene in leaf primordia and developing trichomes in addition to stipules, indicating that the downstream enhancer is required for the normal expression pattern of** *GL1.*

INTRODUCTION

A fundamental process during development is the commitment of cells to different developmental fates, a process known as determination (Slack, 1983; McDaniel, 1985). Determination can occur at several levels: cells may become committed to form a particular organ, tissue, or cell type. The differentiation of trichomes (hairs) on the epidermis of Arabidopsis shoots provides an excellent opportunity to study the commitment of cells to a specific pathway leading to a single, differentiated cell type (reviewed in Marks et al., 1991).

Arabidopsis trichomes are unicellular structures 200 to 500 pm in height, with a thickened cell wall and a papillate surface. They are found on the surfaces of leaves, sepals, petioles, and stems. Trichomes are not essential for viability, and mutations affecting trichome development have been isolated in 10 different genes (Koornneef et al., 1982; Haughn and Somerville, 1988; Marks and Esch, 1993). Multiple mutant alleles have been identified for most of these genes, indicating that most of the genes specifically involved in trichome development have been identified. Recessive mutations in two genes, *GLABROUS7 (GL7)* and *TRANSPARENT TESTA, GLABRA (TTG),* block the initiation of most trichomes. All known mutant alleles at the *TTG* locus are pleiotrophic; mutant plants lack seed coat mucilage and seed coat color as well as trichomes. All alleles of *GL7,* including a null allele in which the entire gene is deleted **(g/7-7;** Oppenheimer et al., 1991), affect only trichome development.

The *GL7* gene has been cloned by T-DNA tagging (Herman and Marks, 1989; Marks and Feldmann, 1989). DNAsequence analysis revealed that the N-terminal domain of the *GL7* gene product contained sequence similarity to the *myb* class of transcriptional regulators (Oppenheimer et al., 1991). Plant *myb* genes are usually present as gene families, although the functions of few plant *myb* genes have been determined to date (reviewed in Larkin et al., 1993).

The *GL7* transcript is very low in abundance and has not been detected on RNA gel blots, although cDNAs have been isolated using polymerase chain reaction. When 1.4 kb of *5'* noncoding sequences containing the putative GL7 promoter were fused to the P-glucuronidase *(GUS)* reporter gene and introduced into Arabidopsis plants, **GUS** activity in the vegetative meristem was detected only in paired stipulelike structures at the bases of the leaves. This result was unexpected, because the trichomes develop from cells on the protoderm of the leaf and not from the stipules.

The existence of additional regulatory sequences for *GL7* was suggested by the nature of the *9/1-43* mutant allele, the T-DNA-induced insertion allele that was used to clone the *GL7* locus (Herman and Marks, 1989; Marks and Feldmann, 1989). The T-DNA insertion in the *9/1-43* allele is located 581 bp downstream of the *GL7* polyadenylation sites (Oppenheimer et al., 1991). This allele exhibits only a partia1 **loss** of *GL7* function; no trichomes are found on the stems of mutant plants, but normal trichomes are present on leaves and sepals. The phenotype of this mutant indicated that some *GL7* regulatory sequences might be located downstream of the coding region. In this report, we demonstrate that *GL7* transcripts are present in leaf primordia and developing trichomes. We have also identified an enhancer-like element located downstream of the

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GL1 transcribed region that is required for all *GL1* functions. Sequences from the region containing the enhancer extend the range of expression of the GUS reporter gene directed by *GL1* 5'sequences to include expression in leaf primordia and developing trichomes.

RESULTS

Localization of *GL1* **Transcripts**

The localization of *GL1* transcripts by in situ hybridization is shown in Figure 1. Tissue sections were hybridized with antisense or sense strand digoxigenin-labeled RNA probes derived from the 3' end of the *GL1* coding region downstream of the conserved *myb* domain. This region of the *GL1* gene has no significant DNA sequence similarity to any of the other Arabidopsis *myb* genes that we have sequenced, and the probe sequences are present in the Arabidopsis genome as a single copy, as determined by DNA blotting (D.G. Oppenheimer, unpublished data).

The strongest hybridization signal was detected in developing trichomes of the leaf and stem (Figures 1A and 1B, filled arrows). Weak hybridization was also detected in other cells of the leaf, particularly in the adaxial epidermis (Figure 1C, white arrow). Occasional strongly hybridizing, unexpanded individual cells in the epidermis were observed (Figure 1C, filled arrow). These cells may represent trichome precursor cells.

No *GL1* transcripts have been detected in mature stipules (Figures 1A and 1D, open arrows). However, low levels of *GL1* transcripts have been observed in developing stipules, young floral primordia, developing stamens, and developing carpels (D. G. Oppenheimer, unpublished data). No hybridization was observed with the sense strand probe (Figure 1D).

Identification of an Enhancer-Like Element Downstream of *GL1* **That Is Required for** *GL1* **Function**

Because the stipule-specific expression pattern of the *GUS* reporter gene in plants containing a fusion of 1.4 kb of *GL1* upstream sequences to the *GUS* coding region (pGL1p/GUS; Oppenheimer et al., 1991) does not accurately reflect the expression pattern of the *GL1* transcript, additional regulatory sequences must be present at the *GL1* locus. The structure of the *gl1-43* allele suggested that sequences required for proper *GL1* function are located downstream of the gene. To test for the presence of such sequences, two sets of nested deletions were constructed in the GL7 3' noncoding region, as shown in Figure 2. The starting point for both sets of deletions was the 4.5-kb Xhol-Bglll fragment previously shown to be sufficient to complement the *g!1-1* deletion mutant (Oppenheimer et al., 1991). Nucleotide coordinates are given relative to the first nucleotide of the start codon, which is 1402 bp downstream of the Xhol site. The production of trichomes on the leaves of regenerating shoots provided a rapid assay

Figure 1. *GL1* Transcript Distribution in Vegetative Meristems.

(A) Longitudinal section through a wild-type vegetative apex hybridized with a GL1-specific antisense probe.

(B) Cross-section through wild-type leaf primordia hybridized with a GL1-specific antisense probe.

(C) Higher magnification view of a portion of the leaf primordia shown in (B).

(D) Longitudinal section through a wild-type vegetative meristem hybridized with a *GL1* sense strand control probe. Filled arrow indicates a possible trichome precursor cell.

Hybridization signals are visible as a blue-purple depositions of pigment. Filled arrows show developing trichomes; open arrows, mature stipules; and white arrow, hybridization in adaxial epidermis.

Sequences retained in each construct are represented by a black bar below the schematic of the *GL1* locus. All of the pGL1-3′∆ constructs contain **the** entire GL7 transcribed region as well as 1.4 kb of upstream sequences. In the pGGE constructs, various portions of the *GL7* 3' noncoding region have been inserted downstream of a chimeric reporter gene in which the GUS coding sequence has been placed under the control of GL1 5' sequences. In pGL1-3'A1 through pGL1-3'A9, all GL1 sequences are in the same relative orientations as in the wild-type gene. For pGL1-**3'AIO** through pGL1-3'A15 and for the pGGE constructs, the relative orientation of the *GL7* 3' noncoding segment present in each construct is indicated by an arrowhead. See Methods for details of plasmid constructions. Complementation results for **the** pGL13'A constructs are based on the ability of the construct to induce the production of trichomes on leaves of regenerating transgenic plantlets **in** culture **(see** Table **1** for data). The restriction sites indicated on the GL7 schematic are X, Xhol; **RV,** EcoRV; B, BamHi. A hatched box indicates the region between the end points of pGL1-3'AI and pGL1-3'A9, which must contain at least a portion of the essential downstream sequences. The positions of the **three** *GL1* exons (black boxes) and the *gl1-43* T-DNA insertion site are also indicated. Functional complementaton of the *gl1-1* mutant phenotype, based on the data in Table 1, is indicated by a +, while failure to complement *g/7-7* is indicated by a -.

for complementation of the $q11-1$ -conferred phenotype (Herman and Marks, 1989). The deletion constructs were used to transform g /1-1 plants and assayed for the ability to complement the mutant phenotype. The plasmid pBINXB, which contains the entire GL7 Xhol-Bglll restriction fragment, was included in each transformation experiment as a positive control.

In the first set of deletion constructs, sequences were removed progressively from the 3'end of the Xhol-Bglll fragment (Figure 2, pGL1-3'Al-pGL1-3'A5). The proportion of regenerating shoots that produced trichomes following transformation with each of these constructs is given in Table 1. The deletion of sequences to position $+2741$ (pGL1-3' Δ 1) had no effect on the ability to complement *917-7,* in comparison to pBlNXB transformants. In contrast, none of the regenerating shoots transformed with a construct deleted to position +2414 (pGL1- 3'A2) produced trichomes (Figure 2 and Table 1). **AI1** deletions with end points upstream of this position also failed to complement *gl1-1* efficiently (pGL1-3' Δ 3-pGL1-3' Δ 5; Figure 2 and Table 1). As shown in Table 2, the analysis of transgenic plant families confirms the results of the tissue culture assay. Most of the transgenic families containing pGL1-3' Δ 2, pGL1-3' Δ 3,

a Kan', kanamycin resistant; trichome', explants producing trichomes. The trichome phenotype of regenerating shoots transformed with various plasmids was scored 4 to 5 weeks after the explants were transferred to kanamycin-containing media.

b Four of the five trichome⁺ pGL1-3' Δ 4 regenerants produced very few trichomes.

^c All four trichome⁺ pGL1-3'∆5 regenerants produced very few trichomes.

^aThe number of independently segregating inserts was determined by following both the segregation of kanamycin resistance and the production of trichomes. The presence of approximately one to three intact copies of *GLI* 3'A transgene was confirmed by DNA gel blotting. **b** The number of trichomes was counted on one member of the first leaf pair of 10 plants in F_3 families segregating for the insert(s). $c +$, Trichomes produced; $-$, trichomes not produced.

d This family was segregating for two independent kanamycinresistant inserts, **but** only one of the inserts was responsible for the production of trichomes.

or pGL1-3'A5 produced no trichomes. Those families containing $pGL1-3' \Delta 5$ that did produce some trichomes produced many fewer trichomes than did pBlNXB transformants (Table 2). These results demonstrate that sequences downstream of the GL7 coding region are required for trichome initiation. The downstream boundary of these essential sequences must lie between the deletion end points of pGL1-3' Δ 1 and pGL1-3' Δ 2.

A second deletion set was constructed removing interna1 sequences from the GL1 3' noncoding region. A common upstream end point was maintained for these deletions at an EcoRV site located at position +1640, which is 17 bp downstream of the first *GL7* polyadenylation site. Four deletions were constructed that removed 568, **774,** 782, and 949 bp downstream of this EcoRV site (Figure 2, pGL1-3'A6-pGL1-3'A9). **All** four constructs complemented the *911-7* mutation for the production of trichomes on leaves in tissue culture (Table 1). pGL1-3'Ag **also** complemented the *g/14* mutant phenotype in transgenic plants (Table 2). The results from this set of deletions indicate that 3'sequences sufficient for *GL7* function must

reside downstream of position +2589, the end point of the pGL1-3'Ag deletion. At least a portion of the essential 3' sequences must lie between positions +2589 and +2741 (hatched section of the GL7 3' noncoding region in Figure 2).

Two possible explanations must be considered regarding the nature of this essential downstream region. This region may contain a transcriptional enhancer required for the proper leve1 and expression pattern of the *GL7* transcript. Alternatively, the region might contain a coding exon missed by the original analysis of the GL7 gene. The latter hypothesis seemed unlikely because two polyadenylation sites were identified at positions +1628 and +1643 by polymerase chain reaction amplification of the 3'ends of the GL7 transcript; longer transcripts were not detected (Oppenheimer et al., 1991). However, two short open reading frames (ORFs) do exist downstream of the three identified coding exons. The first of these, ORF 4 (Oppenheimer et al., 1991), is completely deleted in the pGL1- $3'$ Δ 9 construct, which acts as a functional GL1 gene. The second downstream ORF (ORF 5) is located between the end points of pGL1-3' Δ 1 and pGL1-3' Δ 2, which is within the region required for GL7 function.

To test for the presence of a transcriptional enhancer and to further localize the sequences required for *GL7* function, Several additional constructs were generated containing alterations of the 3' noncoding region (Figure 2). A required exon would not function in the inverse orientation, whereas an enhancer would be expected to function after inversion. In the plasmid pGL1-3' Δ 10, all of the sequences downstream of position +1739 were inverted, and no sequences were deleted (Figure 2). In addition, sequences from three regions, +1795 to +2705 (pGL1-3' Δ 11 and pGL1-3' Δ 12), +2423 to +2914 (pGLl-YA13 and pGL1-3'A14), and +2589 to +2914 (pGL1- $3'$ Δ 15) were inserted downstream of the EcoRV site at +1640 in both orientations (Figure 2). The pGL1-3'A10, pGL1-3'A11, pGL1-3' Δ 12, pGL1-3' Δ 13, and pGL1-3' Δ 14 constructs complemented the *911-7* mutation (Figure 2 and Tables 1 and 2). Thus, the sequences between +2423 and +2914 downstream of the GL7 coding region, in either orientation, are both necessary and sufficient to function as the essential downstream element. However, because the pGL1-3' Δ 15 construct does not complement **g/7-7,** the sequences between +2589 and +2914 are not sufficient for function, at least when present in an inverted orientation.

Sequences from the **3'** Noncoding Region Extend the Expression Pattern of the GL7 Promoter-GUS Fusion Construct

Several portions of the GL7 3' noncoding region were inserted downstream of the GUS reporter gene in the plasmid pGL1p/ GUS to examine whether these sequences would alter the stipule-specific expression pattern. The GUS histochemical staining patterns observed in plants that contain these pGGE (GL1-GUS Enhancer) reporter gene constructs are listed in Table 3. A variety of sequences from the 3' end of the GL1 gene resulted in the expression of GUS activity in leaf primordiaand developing trichomes in addition to expression in stipules, as shown in Figure 3. These sequences are also able to direct the expression of GUS activity in leaf primordia and developing trichomes regardless of orientation. Aberrant staining patterns were exhibited by *two* of 17 families, presumably due to the influence of sequences near the insertion site (Topping et al., 1991). GUS activity was initially expressed throughout the leaf primordium (Figure 3A). In transgenic plants from families with somewhat lower overall expression levels, the highest levels of GUS activity were observed in developing trichomes (Figure 36, open arrow). In older leaves of plants from families exhibiting high levels of expression, GUS activity persisted longest in the trichomes (Figure 3C). GUS activity was also detected in the stem and stem trichomes of bolting plants (Figure 3D). The staining visible within the developing flower buds in Figure 3D is due to weak staining in developing stamens and carpels (data not shown). As expected, in plants transformed with the original pGL1p/GUS plasmid, GUS activity was only detected in stipules (Figure 3E, filled arrow).

Transgenic plants containing the constructs pGGE2 and pGGE3 exhibit histochemical staining for GUS activity in leaf primordia and developing trichomes in addition to expression

^a-, No variation among transformants, except for minor variations in staining intensity.

Figure 3. Addition of *GL1* Downstream Sequences to pGL1p/GUS Increases the Range of the *GUS* Expression Pattern.

- (A) Young seedling apex of a pGGES plant stained with X-gluc.
- (B) Leaf primordium of a pGGE1 plant stained with X-gluc.
- (C) Older leaf of a pGGES plant stained with X-gluc.
- (0) Stem and inflorescence of a bolting pGGES plant stained with X-gluc.
- (E) Seedling of a pGL1p/GUS plant stained with X-gluc.

Filled arrows in (A), (B), (D), and (E) indicate stipules. The open arrow in (B) indicates a developing trichome.

in stipules. However, these constructs do not include the region identified above as essential for GL7 function (Figure 2 and Table 1). In an attempt to resolve this inconsistency, we performed quantitative assays for GUS activity on transgenic seedlings containing various pGGE constructs, as shown in Table 4. The levels of GUS activity detected were quite low, with the highest levels only five- to eightfold above background levels observed in untransformed plants. However, these higher levels of GUS activity were detected only in pGGE4 and pGGE5 transformants, which contain downstream sequences that are required for GL1 function. In contrast, all pGL1p/GUS, pGGE2, and pGGE3 transformed families examined exhibited lower levels of GUS activity (Table 4).

DISCUSSION

Our previous work demonstrated that sequences 5'to the *GL7* gene direct the expression of a GUS reporter gene only in the stipules of developing leaf primordia. However, in this work, we show that *GL7* transcripts are most abundant in leaf primordia and developing trichomes. In addition, we have shown that sequences more than 900 bp downstream of the 3'end of the GL7 transcript contain an enhancer-like element essential for trichome initiation. Sequences from the same 490-bp region downstream of *GL1*, in conjunction with *GL1* 5' sequences, direct the expression of a GUS reporter construct in stipules, leaf primordia, and developing trichomes. The pattern of GUS activity observed in plants containing the pGGE constructs is very similar to the distribution of GL7 transcripts detected by in situ hybridization (compare Figures 1A and 1B with Figure 38). Thus, the same downstream region that is required for GL7 function also acts as an enhancer-like element that confers the proper *GL7* expression pattern on a GUS reporter gene construct. These results strongly suggest that the expression of GL7 in leaf primordia and developing trichomes is essential for GL7 function.

We have not examined the nature of the interaction between the 5' and 3' noncoding sequences of GL7 in promoting

a Data are presented as the mean f SE of the GUS activities (pmol 4-methylumbelliferone/min/mg protein) in tissue extracts from equivalently staged seedlings from independent transformants.

expression of the reporter gene in leaf primordia and developing trichomes. Thus, we cannot yet distinguish whether the 3'sequences contain a true tissue-specific enhancer or a constitutive enhancer acting through the *GL7* 5' sequences. Experiments to discriminate between these possibilities are in progress.

We have not been able to detect high levels of *GL7* transcripts in stipules, despite the high levels of GUS activity detected in the stipules of plants containing pGL1p/GUS or the pGGE constructs. It seems unlikely that the histochemical localization of GUS activity in stipules is artifactual. The tissue was prefixed in 0.1% formaldehyde to prevent the induction of enzyme activity during the assay, and $K_3Fe(CN_6)$ was included in the assay buffer to promote the in situ precipitation of the indigo blue precipitate (De Block and Debrouwer, 1992). One possible explanation for our failure to detect GL7 transcripts in older stipules may be that RNA in stipules is **less** accessible to hybridization. Arabidopsis stipules appear to be glandular and rather different from the surrounding tissues in their histological properties. **We** have been unable to detect GL7 transcripts in the stipules of plants containing a *GL7* coding region under the control of the cauliflower mosaic virus (CaMV) 35s promoter, whereas high levels of the *GL7* transcript were detected in all surrounding tissues (D.G. Oppenheimer, unpublished observations). In contrast, the 35s promoter drives GUS expression at high levels in both stipules and the surrounding tissues.

Another minor difference between the pattern of GL7 transcript accumulation and the pattern of GUS activity in the pGGE constructs is that GUS activity is detected in all cell layers of the developing leaf primordium, whereas the strongest in situ hybridization signals are detected in the protoderm. However, in older leaves, in which the GUS histochemical staining of nontrichome cells has become weaker, preferential staining of the epidermis has been observed (J.C. Larkin, unpublished data). It seems likely that differences in the relative stabilities of the GL7 transcript and the GUS protein (andlor transcript) contribute to the minor differences that we have observed between the in situ hybridizations and the GUS histochemical staining patterns. However, we cannot rule out the possibility that additional regulatory sequences are located within the transcribed region of *GL7.*

Several lines of evidence suggest that the 3' enhancer contains functionally redundant sequences. First, the pGL1-3' Δ 1 and pGL1-3'Ag deletion constructs define a 152-bp region between positions +2589 and +2741 that must contain at least a portion of the essential sequences (hatched section of the GL1 3' noncoding region in Figure 2). However, pGL1-3' Δ 15, which contains downstream sequences from +2589 to +2913, fails to complement g/1-1. pGL1-3' Δ 9 has the same upstream end point as $pGL1-3' \Delta 15$, but contains additional downstream sequences, whereas pGL1-3' Δ 13 and pGL1-3' Δ 14 have the same downstream end point as $pGL1-3^{\prime}\Delta15$ but contain additional upstream sequences. All three of these constructs complement **g/7-7,** indicating that the addition of sequences either upstream of $+2589$ or downstream of $+2913$ to the sequences present in pGL1-3 $^{\prime}$ Δ 15 is sufficient to restore function. Additional evidence for functional redundancy comes from the pGGE results (Table 3).

The *GL7* 3' noncoding sequences present in pGGE2 and pGGE3 do not qverlap with the 3'sequences present in pGGE4 and pGGE5, yet all of these constructs direct *GUS* expression in leaf primordia and developing trichomes. The downstream region contained in the pGGE2 and pGGE3 constructs is not required for or sufficient for *GL7* function. Consistent with these observations, pGGE4 and pGGE5 transformants have higher levels of GUS activity than pGGE2 and pGGE3 transformants in quantitative GUS assays (Table 4). Thus, it is possible that a combination of several equivalent copies of a leaf primordia/trichome-specific enhancer is required to raise the level of *GL7* expression to some functional threshold. Similar results have been obtained with other plant transcriptional regulatory sequences (Leyva et al., 1992).

We have not detected any stem-specific regulatory elements that could be responsible for the phenotype of the **9/3-43** T-DNA insertion allele during our analysis of the *GL7* 3' regulatory region. As described above, plants homozygous for this allele have normal leaf trichomes but lack trichomes on stems. All of the deletion constructs generated here affect both leaf and stem trichomes (Table 2). It seems likely that the phenotype of the *9/1-43* mutant allele is the result of the nove1 sequences inserted downstream of the gene, rather than the result of the disruption of existing *GL7* regulatory sequences that are specific to stem trichomes. Alternatively, subtle interactions between subdomains of the 3'regulatory region may have been missed by this analysis. However, it should be noted that pGL1- 3'A9 removes sequences from 584 bp upstream of the *9/1-43* T-DNA insertion site to 365 bp downstream of the insertion site and has no effect on trichome development. Thus, it seems unlikely that a stem-specific regulatory sequence is located in this region.

Many examples of enhancers located downstream of the transcription initiation site are known in animal systems (Choi and Engel, 1986; Fischer and Maniatis, 1986). A particularly striking example is provided by the mouse erythropoietin gene (Pugh et al., 1991). An enhancer located 3'of the polyadenylation site of this gene directs oxygen-regulated transcription even in constructs removing the normal transcription start site. In plants, several instances of enhancers or enhancer-like sequences that can function 3'of the transcription start site have been identified. The CaMV enhancer is able to function when placed downstream of a reporter gene (Nagy et al., 1987). Similarly, an enhancer from the **5'** noncoding region of a soybean Pconglycinin gene can confer seed-specific expression on the CaMV 35s promoter when located between the reporter gene coding region and the polyadenylation site but not when located downstream of the polyadenylation site (Chen et al., 1988). Both of these enhancers function more effectively when located 5'of the gene. Dean et al. (1989) have shown that sequences located 3' of the coding region of petunia ribulose bisphosphate carboxylase small subunit genes influence the rates of transcription of the genes. Sequences located within or near the coding region and required for the full range of expression have been described for the pea ferredoxin I gene (Elliot et al., 1989), a parsley 4-coumarate:coenzyme A ligase gene (Douglas et al., 1991), and the oilseed rape AX92 gene (Dietrich et al., 1992), although none of these sequences has been shown to function as a transcriptional enhancer.

The in situ hybridization results, in conjunction with the expression pattern of the reporter gene constructs, indicate that *GL7* is initially expressed throughout the leaf primordium, with higher transcript levels in the protoderm. However, much higher transcript levels are seen in developing trichomes. In fact, high levels of *GL7* transcripts are detected in some protodermal cells that have not shown any sign of cell expansion (Figure 1C, solid arrow). Thus, it appears that a rapid increase in the *GL7* transcript level is an early event in the commitment of a protodermal cell to the trichome development pathway. In some respects, this observation is reminiscent of the pattern of scufe expression during sensory bristle formation in Drosophila (reviewed in Ghysen et al., 1993). Expression of scufe is initially detected in afield of cells but ultimately becomes restricted to a single "sensory mother cell" expressing high levels of scute transcripts and protein. This comparison with the well-studied bristle development system in Drosophila brings into focus an important unanswered question about the trichome developmental pathway: How are the trichome precursor cells selected? The continued analysis of the control of *GL7* expression, in conjunction with the study of other components of the trichome developmental pathway, should provide a model for understanding the commitment of plant cells to specific developmental fates.

METHODS

In Situ Hybridizatlon

The template for the in situ hybridization probe was derived from pGLlexo4.9. This exonuclease Ill-generated deletion (Oppenheimer et al., 1991) completely removes the myb DNA binding region of *GLABROUS7 (GL7*) and leaves the single-copy, carboxy-terminal **do**main of the protein-coding region. The pGL1exo4.9 clone was digested with SstII and religated to remove the majority of the GL1 3' noncoding region. This clone was called pGLIC3. The template for the production of digoxygenen-UTP-labeled RNA probes by runoff transcription was BssHII-digested pGL1C3. T3 RNA polymerase was used to generate antisense strand GL7 probes, and T7 RNA polymerase was used to generate sense strand *GL7* probes, using components obtained from Boehringer Mannheim. Plant tissue was fixed, dehydrated, and embedded in paraffin essentielly **as** described by Jackson (1991). Sections, 8-pm thick, were cut and mounted on poly-L-lysine-coated slides and processed essentially as described by Drews et al. (1991), except that the BMB blocking reagent from a Genius kit (Boehringer Mannheim) was substituted for Denhardt's solution. After washing the slides, they were processed and developed as recommended in the Genius kit. Slides were allowed to develop 1 to 3 days before photographing.

Construction of 3' Deletions

The starting point for the construction of deletions of the *GL7* downstream region was the plasmid pGLl (Oppenheimer et al., 1991). This plasmid contains the 4.5-kb Xhol-Bglll fragment of *GL7* inserted into the Xhol and BamHl sites of the vector pBluescriptll SK+ (Stratagene). To generate nested deletions from the 3' end, the Xhol-Hhal (pGL1- 3' Δ 1, deletion end point +2741), Xhol-Dralll (pGL1-3' Δ 2, deletion end point +2414), Xhol-Nrul (pGL1-3'A3, deletion end point +2208), Xhol-Sstll (pGL1-3' Δ 4, deletion end point +1740), and Xhol-EcoRV (pGL1- $3'$ Δ 5, deletion end point +1641) restriction fragments from pGL1 were each cloned into Sall, Smal-cut pBIN19 (Bevan, 1984). The Sstll and Hhal ends were rendered blunt with T4 polymerase before ligation, according to the manufacturer's instructions (New England Biolabs, Beverly, MA).

The internal deletions of the *GL1* 3' end were generated by cutting pGLl with EcoRV and Sstl (a vector site), treating the plasmid with calf intestinal phosphatase (Boehringer Mannheim), and inserting various isolated restriction fragments. The fragments used were the Nrul-Sstl (pGL1-3' Δ 6), Draill-Sstl (pGL1-3' Δ 7), EcoRV-Sstl (pGL1-3' Δ 8), and Hincll-Sstl (pGL1-3' Δ 9) fragments. The sequences deleted in these constructs are pGLl-3'AC +1641 to +2208; pGL1-3'A7, +I641 to +2414; $pGL1-3' \Delta 8$, +1641 to +2422; and pGL1-3' $\Delta 9$, +1641 to +2488. In a similar manner, the pGL1-3'A11 through pGL1-3'A15 constructs were generated by ligating the following restriction fragments from the *GL7* downstream region to EcoRV, Xbal-cut pGL1, where the Xbal site had been filled in with T4 DNA polymerase: the 911-bp Dral fragment (pGL1- $3'$ Δ 11 and pGL1-3' Δ 12, +1795 to +2415), the 490-bp EcoRV-Accl fragment (pGL1-3'A13 and pGL1-3'A14, +2423 to +2913), and the 324-bp Hincll-Accl fragment (pGL1-3' Δ 15, +2489 to +2913). The pGL1-3' Δ 10 construct was generated by inverting the Sstlt fragment of pGL1. The pGL1-3'A6 through pGL1-3'A15 constructs were excised from the pBluescriptll SK+ vector with Xhol and Sstl and inserted between the Sall and Sstl sites of pBIN19. The plasmids were transferred to Agrobac*terium* tumefaciens LBA4404 (Hoekema et al., 1983; An et al., 1988) by electroporation.

Construction of pGLlp/GUS Enhancer Plasmlds

The plasmid pGLlp/GUS (Oppenheimer **et** al., 1991) contains 1.4 kb of *GL1* 5' noncoding sequence fused to the P-glucuronidase *(GUS)* coding region of pBl101 (Jefferson et al., 1987). The pGGE plasmids were constructed by ligating restriction fragments from pGLl into the EcoRl site (filled in with T4 polymerase and deoxynucleotides) that is located downstream of the nopaline synthetase terminator of pGLlp/GUS. The restriction fragments used were the 911-bp Dral fragment (pGGE1), the 782-bp EcoRV fragment (pGGE2 and pGGE3), and the 490-bp EcoRV-Accl fragment (pGGE4 and pGGE5).

Plant Transformation

Arabidopsis thaliana root explants were transformed and regenerated as described previously (Feldmann and Marks, 1986; Valvekens et al., 1988; Oppenheimer et al., 1991). The pGGE plants were regenerated from explants of the Wassileskija genotype **(WS).** Most of the **g/7-7** mutant explants that were transformed with the 3' deletion constructs were of the genotype *9/14, an* in a Columbia genetic background (Oppenheimer et al., 1991). A few experiments were conducted with explants from plants in which the **g/7-7** mutation had been backcrossed three times into the WS background, which yields transformants at a much higher frequency. No differences between the two genotypes in the frequency of explants producing trichomes were observed in control experiments. Data on the proportion of regenerating explants producing trichomes from both backgrounds are pooled in Table 1. In each transformation experiment, pBINXB-transformed explants were

included as positive controls. In **most** experiments, pGL1-3'A5-transformed explants were included as negative controls.

All of the transgenic plant families described in Table 2 were regenerated from the **g/7-7,** an background. The number of independent inserts was determined by following the segregation of both kanamycin resistance and trichome phenotype. The presence of approximately one to three intact copies of the transgene in each family was confirmed by DNA gel blotting. Trichome counts given in Table 2 were determined by counting the number of trichomes on one member of the first leaf pair of T3 seedlings in families segregating for the insert(s), as described in Table 2.

Hlstochemlcal and Quantitative GUS Assays

Plant tissue was prefixed under vacuum in 0.1% formaldehyde, 100 mM NaPO₄, pH 7.0, 0.1% Triton X-100, rinsed in 50 mM NaPO₄, pH 7.0, and immersed in 1 mglmL **5-bromo-4-chloro-3-indoyl** glucuronide (X-gluc), 1 mM $K_3Fe(CN_6)$, 0.1% Triton X-100, 50 mM NaPO₄, pH 7.0, at 37% for 6 to 18 hr (Jefferson et al., 1987; De Block and Debrouwer, 1992). After staining, the tissue was fixed in 3.6% formaldehyde, 5% acetic acid, 50% ethanol for 15 min under vacuum, cleared in 95% ethanol at 65°C for 5 min, and rehydrated through an ethanol series. Tissue samples were stored at 4°C in 10 mM Tris-HCI, 1 mM EDTA before undergoing photomicroscopy. For quantitative GUS assays, pooled seedlings of each genotype were ground to a powder in liquid N_2 and homogenized in extraction buffer (50 mM NaPO₄, pH 7.0, 1 mM EDTA, 0.1% Triton X-100, 0.1% sarkosyl, 10 mM p-mercaptoethanol). After centrifuging for 10 min in a microcentrifuge, the samples were frozen at -70°C. Aliquots of these samples were assayed for GUS activity in extraction buffer containing 1 mM 4-methylumbelliferyl p-o-giucuronide, as described by Jefferson et al. (1987). Fluorescence was measured in a minifluorometer (model No. TK0100; Hofer, San Francisco, CA).

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