

Regulation of *WUSCHEL* Transcription in the Stem Cell Niche of the *Arabidopsis* Shoot Meristem ^W

Isabel Bäurle¹ and Thomas Laux²

Institute of Biology III, Freiburg University, D-79104 Freiburg, Germany

Pluripotent stem cells are localized in specialized microenvironments, called stem cell niches, where signals from surrounding cells maintain their undifferentiated status. In the *Arabidopsis thaliana* shoot meristem, the homeobox gene *WUSCHEL* (*WUS*) is expressed in the organizing center underneath the stem cells and integrates regulatory information from several pathways to define the boundaries of the stem cell niche. To investigate how these boundaries are precisely maintained within the proliferating cellular context of the shoot meristem, we analyzed the transcriptional control of the *WUS* gene. Our results show that the *WUS* promoter contains distinct regulatory regions that control tissue specificity and levels of transcription in a combinatorial manner. However, a 57-bp regulatory region is all that is required to control the boundaries of *WUS* transcription in the shoot meristem stem cell niche, and this activity can be further assigned to two adjacent short sequence motifs within this region. Our results indicate that the diverse regulatory pathways that control the stem cells in the shoot meristem converge at these two short sequence elements of the *WUS* promoter, suggesting that the integration of regulatory signals takes place at the level of a central transactivating complex.

INTRODUCTION

Plants produce most of their organs postembryonically from stem cells at the shoot and root apices. Similar to animal stem cells, plant stem cells are located in niches where neighboring cells provide signals to maintain them in an undifferentiated state (Spradling et al., 2001; Weigel and Jürgens, 2002; Laux, 2003). Cells that leave the stem cell niche initiate differentiation and give rise to lateral organs such as leaves and flowers. In general, the decision between stem cell fate and differentiation can be regulated in two ways: either in a lineage mechanism in which each stem cell divides asymmetrically to give rise to one stem cell and one cell prone to undergo differentiation, or in a population-based mechanism by which the outcome of an individual division cannot be predicted; rather, the stem cell population as a whole is kept constant by external cues (Spradling et al., 2001). The shoot meristem stem cell niche operates in a population mode and thus requires precise spatial regulation of stem cell-inducing signals to maintain the correct position and number of stem cells (Bäurle and Laux, 2003). Because all cells in the shoot meristem, including the signaling niche cells, continuously divide, a long-standing question of plant development is how the position and the boundaries of the stem cell niche are stably maintained, or, as Newman (1965) put it several decades ago, how the pattern (of

the shoot apex) can be maintained while the matter (the constituent cells) constantly changes.

The regulation of transcriptional domains of regulatory genes plays a pivotal role for many developmental processes; thus, the analysis of transcriptional control is crucial to gain insight into the mechanisms that govern spatial and temporal patterning in development (Watanabe and Okada, 2003). In *Arabidopsis thaliana* shoot and floral meristems, transcriptional regulation of the *WUSCHEL* (*WUS*) homeobox gene controls the stem cell pool. *WUS* is expressed in a small group of cells underneath the stem cells termed the organizing center and is required to keep the stem cells in an undifferentiated state, indicating that the organizing center cells act as signaling cells of the shoot meristem stem cell niche (Laux et al., 1996; Mayer et al., 1998). Ectopic expression of *WUS* inhibits differentiation and can result in the formation of ectopic stem cells or even somatic embryos, indicating the necessity to locally restrict *WUS* activity (Schoof et al., 2000; Brand et al., 2002; Zuo et al., 2002; Gallois et al., 2004). Recent findings indicate that the regulation of *WUS* transcription is a central checkpoint in stem cell control, integrating information from several regulatory pathways. First, the size of the stem cell population is controlled through the size of the *WUS* expression domain. This is achieved by a dynamic feedback loop, with *WUS* indirectly activating the expression of the signaling peptide *CLAVATA3* (*CLV3*) in the stem cells and *CLV3* repressing *WUS* transcription through the *CLV1* receptor kinase signaling pathway (Brand et al., 2000; Schoof et al., 2000; Rojo et al., 2002; Lenhard and Laux, 2003). Second, temporal control of stem cell activity in the determinate floral meristem is achieved by the repression of *WUS* transcription through *AGAMOUS* (*AG*) activity (Lenhard et al., 2001; Lohmann et al., 2001). Furthermore, based on changes of its expression domain in shoot meristem mutants, several other regulatory pathways have been implicated in the control of *WUS* gene expression

¹ Current address: John Innes Centre, Colney Lane, Norwich NR4 7UH, UK.

² To whom correspondence should be addressed. E-mail laux@biologie.uni-freiburg.de; fax 49-761-203-2745.

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: Thomas Laux (laux@biologie.uni-freiburg.de).

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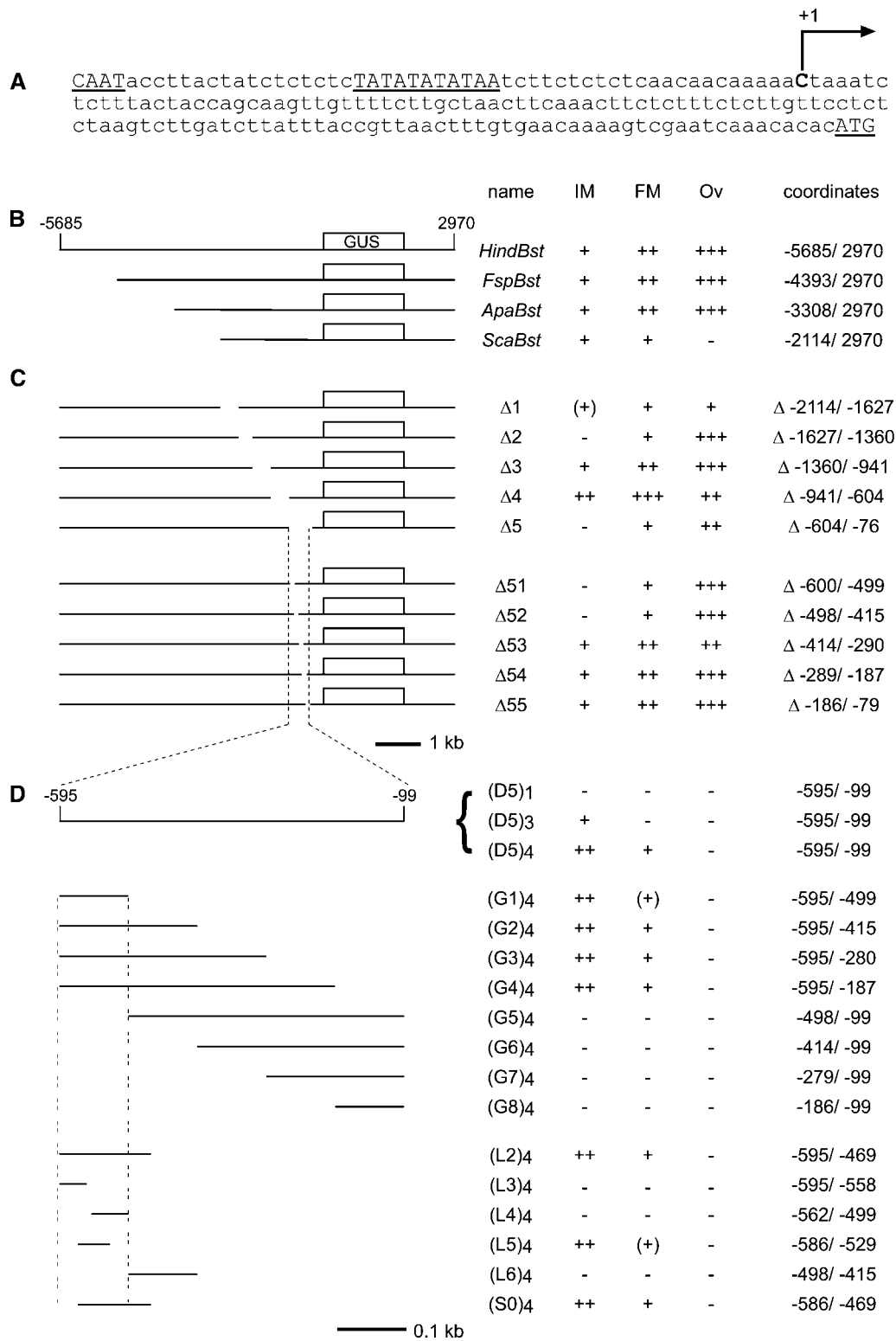


Figure 1. Diagram of the Reporter Constructs Used in the *WUS* Promoter Analysis.

(A) The putative *WUS* transcription start site (+1) was determined by RACE PCR. Putative CAAT and TATA boxes and the start codon are underlined.
(B) to (D) For each construct, a scheme is shown at left. At right, the name, the relative staining intensities in the inflorescence meristem (IM), the floral

(Laufs et al., 1998; Kaya et al., 2001; Stuurman et al., 2002; Bertrand et al., 2003; Ueda et al., 2004; Zhao et al., 2004; Carles et al., 2005; Wu et al., 2005). In addition to the inhibitory CLV3 signal, a positive signal originating from the stem cells has been postulated that would activate *WUS* transcription and anchor the organizing center to the shoot tip (Schoof et al., 2000). However, no direct regulator of *WUS* transcription has been identified to date.

Therefore, understanding how the boundaries of the *WUS* transcriptional domain are regulated is central to gaining insight into how the position and size of the stem cell niche are maintained at the tip of the shoot meristem. Here, we have identified two short sequence motifs within the *WUS* promoter that act as central integrating elements in stem cell control.

RESULTS

Regulatory Elements That Control *WUS* Transcription in the Stem Cell Niche

To delimit the control region of the *WUS* gene, we analyzed the expression pattern of β -glucuronidase (*GUS*) driven by *WUS* promoter fragments. The putative transcription start site of the *WUS* gene was determined by two independent rapid amplification of cDNA ends (RACE) experiments as being 126 nucleotides upstream of the ATG start codon (Figure 1A). As a starting point for promoter analysis, we chose an 8.7-kb *WUS* genomic fragment, *HindBst* (–5685/2970; referring to the putative transcriptional start site), in which the *WUS* coding region was replaced with the coding region of the *GUS* gene (Figure 1B). This reporter comprised 5.7-kb upstream and 1.3-kb downstream sequences and mimicked the described mRNA expression pattern of *WUS* in meristems (Mayer et al., 1998), showing strong *GUS* expression in young floral meristems and weaker *GUS* expression in the organizing center of the vegetative and inflorescence meristems (Figure 2A). The construct also recapitulated ovule-specific mRNA expression (Figure 2A), in which *WUS* is expressed in the apical nucellus during early developmental stages and is required for ovule patterning (Gross-Hardt et al., 2002; Sieber et al., 2004). In addition, *GUS* staining in stamens was detected in a pattern similar to the mRNA expression (Wellmer et al., 2004) but was very weak and was not analyzed further. We focused our analysis on inflorescence and floral meristems but obtained corresponding results for the vegetative meristem of the seedling where analyzed (Figures 2 and 3E).

Progressive truncations from the 5' end revealed that the region upstream of position –3308 was dispensable for promoter activity (Figures 1B and 2A, *ApaBst*). By contrast, the region between –3308 and –2114 was essential for expression in ovules and for high-level expression in floral meristems (Figures 1B and 2A, *ScaBst*). In plants carrying the *ScaBst* (–2114/2970) reporter construct, the spatial *GUS* expression pattern in inflorescence and floral meristems was unchanged, indicating that the regulatory sequences controlling the boundaries of *WUS* expression in the stem cell niche are present (Figures 2A, 3I, and 3J). Corresponding *WUS* genomic versions of these promoter truncations were able to complement the inflorescence and floral meristem defects observed in *wus* mutants (Figure 2A, Table 1). In accordance with the loss of *GUS* activity in ovules, seed set in *wus* mutants complemented with the *ScaBst* genomic construct was strongly reduced.

To further characterize the *WUS* regulatory regions, we analyzed the effects of internal deletions of 267 to 528 bp in length within the 2.1-kb upstream region (Figure 1C). We performed the experiments in the context of the complete *HindBst* reporter to minimize positional effects attributable to different integration sites of the transgenes. Deletion $\Delta 1$ (Δ –2114/–1627) resulted in qualitatively unaltered but generally weaker *GUS* expression (Figure 2B), indicating that the deleted sequence harbors general transcriptional enhancer element(s). Deletion $\Delta 3$ (Δ –1360/–941) did not alter the *GUS* expression pattern (Figure 1C), and deletion $\Delta 4$ (Δ –941/–604) gave stronger *GUS* expression in the inflorescence and floral meristems (Figure 2B). However, replacement of the latter region by an unrelated DNA fragment of the same length produced normal *GUS* expression levels (data not shown), suggesting that the increased expression strength in $\Delta 4$ was caused by spacing effects rather than by the excision of a negative regulatory element. Again, the corresponding *WUS* genomic deletion constructs ($\Delta 1$, $\Delta 3$, and $\Delta 4$) complemented the inflorescence and floral meristem defects of the *wus* mutant, and seed set was restored to wild-type levels (Figure 2B, Table 1; data not shown).

Deletions $\Delta 2$ (Δ –1627/–1360) and $\Delta 5$ (Δ –604/–76) completely abolished *GUS* expression in the inflorescence meristem and reduced expression in floral meristems (Figure 2B), indicating the presence of essential regulatory elements in these regions. However, the genomic fragment carrying the $\Delta 2$ deletion complemented the inflorescence and flower phenotype conferred by the *wus* mutant (Figure 2B, Table 1). This finding suggests that the $\Delta 2$ deletion reduced transcription below the detection limit of the *GUS* assay but still allowed for sufficient transcription of *WUS* to rescue stem cell maintenance in the

Figure 1. (continued).

meristem (FM), and the ovule (Ov) [–, none; (+), very faint; +, weak; ++, moderate; +++, strong], and the exact coordinates of the *WUS* promoter fragments or deletions (Δ) are given. The *WUS* coding region was replaced by the *GUS* coding sequence (box).

(B) Diagram of the truncation constructs analyzed.

(C) Diagram of the internal deletion constructs analyzed.

(D) Diagram of the constructs used during the functional definition of *cis*-regulatory sequences. To generate the reporter constructs, the monomers were multimerized as indicated and fused to –60 *CaMV*:*GUS*. D (deletion), G (gain of function), L (little deletion), and S (linker scanning) denote series of constructs.

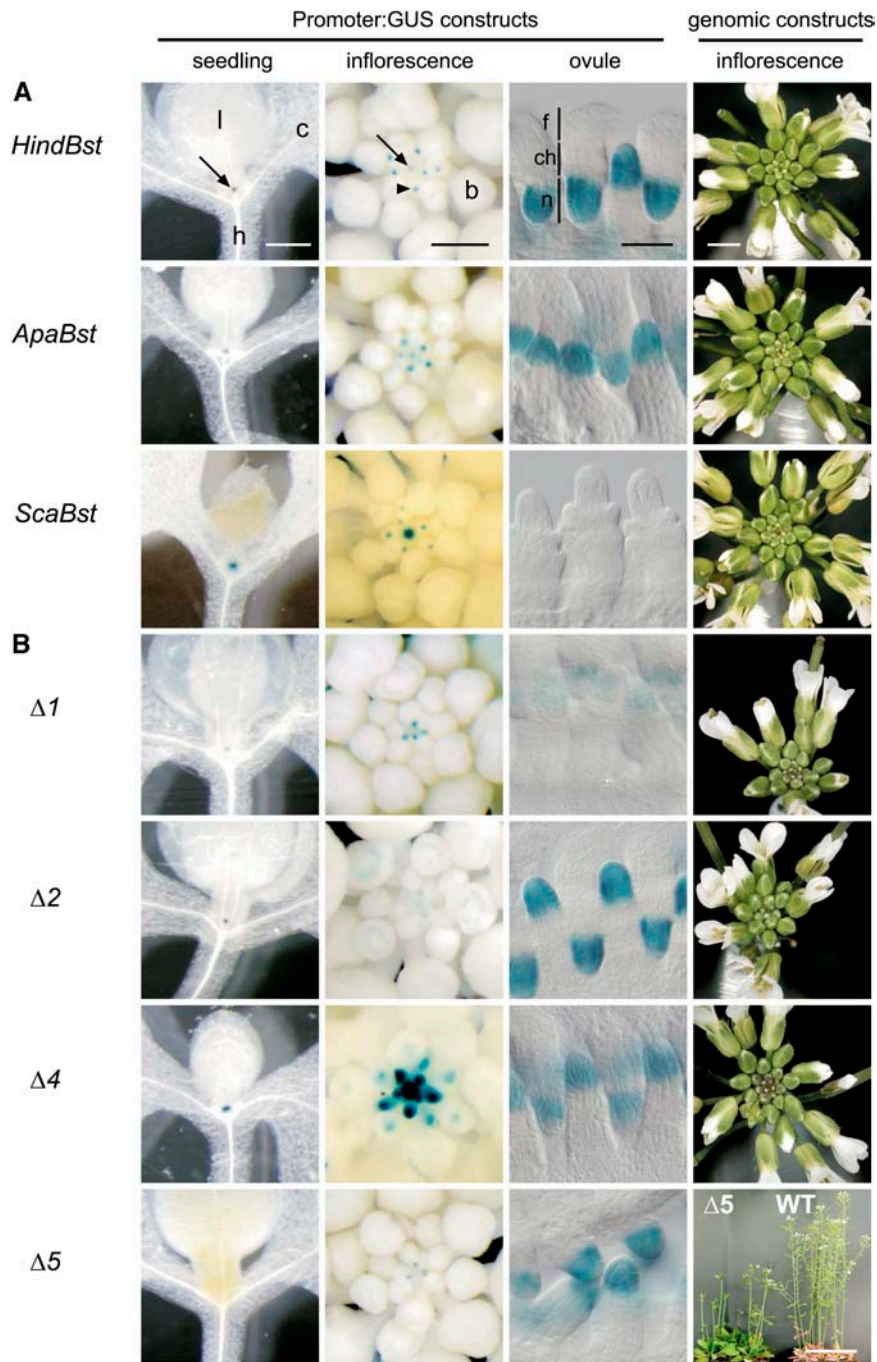


Figure 2. Expression Patterns of *WUS:GUS* Constructs and Complementation of the Inflorescence Phenotype with Corresponding Genomic Fragments.

Each row shows the expression pattern of the indicated *WUS* promoter fragment in (from left to right) seedlings, inflorescences, and ovules and the complementation of a homozygous *wus-1* mutant plant with the corresponding genomic fragment (right column). Seedlings and inflorescences were stained with GUS with 2 mM Fe-cyanide for 1 d except *ScaBst* (2 mM, 2 d) and $\Delta 4$ (5 mM, 1 d). Ovules were stained with GUS with 5 mM Fe-cyanide for 1 d.

(A) Truncation constructs.

(B) Deletion constructs.

b, floral bud; c, cotyledon; ch, chalaza; f, funiculus; h, hypocotyl; l, leaf; n, nucellus. Arrows indicate the shoot meristem, and the arrowhead indicates the floral meristem. Bars = 0.5 mm (seedling, inflorescence), 30 μ m (ovule), 2 mm (genomic constructs, except $\Delta 5$), and 5 cm (genomic construct $\Delta 5$ and the wild type).

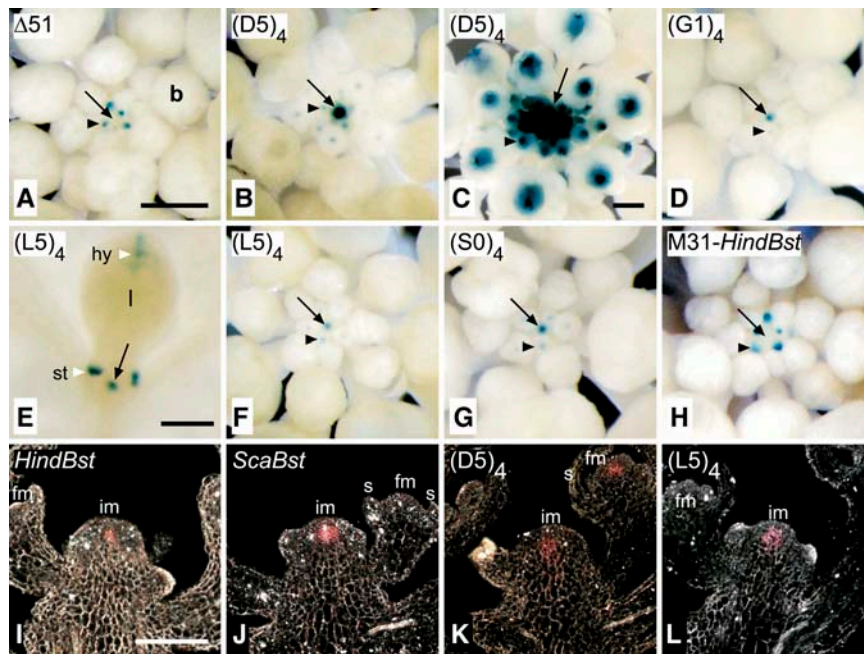


Figure 3. Expression Patterns of *WUS:GUS* Reporter Constructs in Inflorescences.

Inflorescences (**[A]** to **[D]** and **[F]** to **[L]**) or seedling (**[E]**) were stained with either 2 mM Fe-cyanide (**[A]**, **[B]**, and **[D]** to **[H]**) or 5 mM Fe-cyanide (**[C]** and **[I]** to **[L]**) in the staining buffer for 1 d (**[A]** and **[H]** to **[J]**) or 3 d (**[B]** to **[G]**, **[K]**, and **[L]**).

(A) to **(H)** Whole-mount views with bright-field optics. GUS activity is visualized by blue color.

(A) $\Delta 51$.

(B) $(D5)_4$.

(C) $(D5)_4$ *clv1-4*.

(D) $(G1)_4$.

(E) $(L5)_4$. Staining in hydathodes (hy) and stipules (st) is attributable to background activity of the included minimal promoter.

(F) $(L5)_4$.

(G) $(S0)_4$.

(H) *M31-HindBst* (–566/–564 mutated; see Figure 4).

(I) to **(L)** Eight-micrometer sections viewed with dark-field optics. GUS activity is visualized by pink color.

(I) *HindBst*.

(J) *ScaBst*.

(K) $(D5)_4$.

(L) $(L5)_4$.

b, floral bud; fm, floral meristem; im, inflorescence meristem; s, sepal. Arrows indicate the shoot meristem, and arrowheads indicate the floral meristem. Bars = 0.5 mm (**[A]** and **[E]**), 2 mm (**[C]**), and 60 μ m (**[I]**). Magnification of **(B)**, **(D)**, and **(F)** to **(H)** is as in **(A)**; magnification of **(J)** to **(L)** is as in **(I)**.

shoot meristem and to a reduced extent in the floral meristem. Because ovule-specific expression was not affected in $\Delta 2$ (Figures 1C and 2B), the region between –1627 and –1360 presumably contains a meristem-specific enhancer (Figure 4).

By contrast, the genomic fragment carrying the $\Delta 5$ deletion did not rescue the phenotype conferred by the *wus* mutant (Figure 2B, Table 1), suggesting the presence of essential control elements in the region between –604 and –76. Therefore, we further analyzed this sequence by introducing 100-bp deletions within this region in the context of the *HindBst* GUS reporter (Figure 1C). Deletions $\Delta 51$ (Δ –600/–499) and $\Delta 52$ (Δ –499/–415) both abolished GUS expression in the inflorescence meristem and reduced it in floral meristems (Figures 1C and 3A), very similar to what was observed in deletion $\Delta 5$. By contrast,

deletions $\Delta 53$ to $\Delta 55$ (covering –414 to –79) did not affect GUS expression patterns (Figure 1C).

Collectively, these results suggest that the regulation of *WUS* transcription is mediated through several distinct *cis*-regulatory regions. However, the sequences between –600 and –415 are the only ones that are absolutely necessary for *WUS* expression in the inflorescence meristem stem cell niche; therefore, we focused our further analysis on this region.

A 57-bp Element Controls *WUS* Expression Boundaries in Shoot and Floral Meristem Stem Cell Niches

We next asked whether the identified sequences required for the correct expression of *WUS* in the shoot stem cell niche are also

Table 1. Complementation of *wus-1* Mutant Flowers with Promoter Truncations and Internal Deletion Constructs

Construct	Whorl				<i>n</i>
	Sepals	Petals	Stamens	Carpels	
<i>HindBst</i>	4.0 ± 0	4.0 ± 0	5.3 ± 0.4	2.0 ± 0	20
<i>FspBst</i>	4.0 ± 0	4.0 ± 0	5.6 ± 0.5	2.0 ± 0	20
<i>ApaBst</i>	4.0 ± 0	4.0 ± 0	5.8 ± 0.4	2.0 ± 0	20
<i>ScaBst</i>	4.0 ± 0	4.0 ± 0	5.9 ± 0.3	2.0 ± 0	20
Δ1	4.0 ± 0	4.0 ± 0	5.7 ± 0.5	2.0 ± 0	20
Δ2	4.0 ± 0	4.0 ± 0	4.5 ± 1.2	1.3 ± 0.8	20
Δ3	4.0 ± 0	4.0 ± 0	5.9 ± 0.4	2.0 ± 0	20
Δ4	4.0 ± 0	4.0 ± 0	5.9 ± 0.2	2.0 ± 0	18
Δ5	4.5 ± 1.0	4.4 ± 0.8	5.1 ± 1.3	0.1 ± 0.3	20
<i>wus-1</i> ^a	3.7 ± 0.7	3.6 ± 0.9	0.9 ± 0.5	0.0 ± 0	53
<i>Ler</i>	4.0 ± 0	4.0 ± 0	6.0 ± 0.3	2.0 ± 0	20

The average floral organ number and the SD in *n* flowers of the line that showed the best complementation are indicated.

^aData for *wus-1* are taken from Laux et al. (1996).

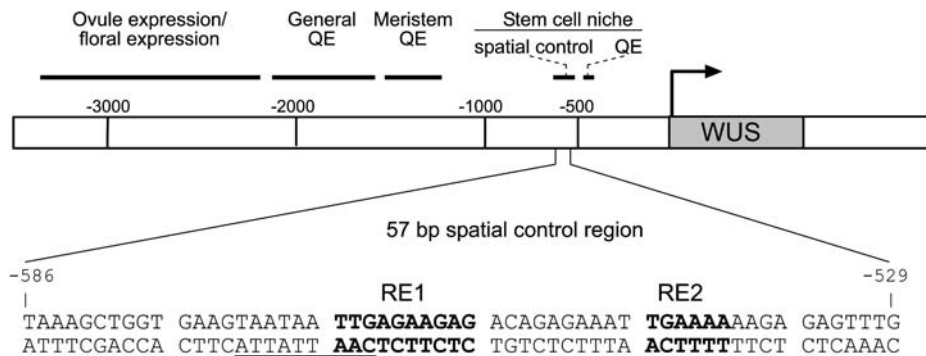
sufficient. For this purpose, we fused tandem repeats of the region deleted in Δ5 (−604/−76), designated D5, to the minimal promoter of the *Cauliflower mosaic virus* (*CaMV*) 35S gene (−60 *CaMV*) followed by the *GUS* coding sequence, and tested whether these synthetic promoters were able to drive *GUS* expression in transgenic *Arabidopsis* plants. We did not detect any *GUS* expression in plants carrying the reporter with only a single copy of D5 [(D5)₁:*GUS*; Figure 1D]. By contrast, three tandem copies of D5 provided moderate levels of *GUS* expression in the organizing center of the inflorescence meristem [(D5)₃:*GUS*; Figure 1D], and four copies conferred strong *GUS* staining in the inflorescence meristem and weaker staining in young floral meristems [(D5)₄:*GUS*; Figures 1D, 3B, and 3K]. Importantly, the spatial expression pattern provided by these multimers precisely recapitulated the one observed with the *ScaBst* reporter (cf. Figures 3J and 3K) and in *WUS* RNA in situ hybridizations (Mayer et al., 1998; Schoof et al., 2000), indicating that the D5 promoter fragment contains all of the necessary sequences for correct transcriptional control in the stem cell

niche. Similar to the endogenous *WUS* gene, the (D5)₄:*GUS* reporter displayed strongly enhanced expression in a *clv1-4* mutant background (Figure 3C) (Schoof et al., 2000). In addition, expression of the (D5)₄:*GUS* reporter in wild-type floral meristems was terminated approximately when carpel primordia emerged (data not shown), similar to the time when endogenous *WUS* expression is terminated (Mayer et al., 1998). Together, these findings suggest that the repression of *WUS* transcription mediated by *CLV3* (Brand et al., 2000; Schoof et al., 2000) and *AG* (Lenhard et al., 2001; Lohmann et al., 2001) acts through the D5 control region. However, we did not find an *AG* consensus binding site (Shiraishi et al., 1993) anywhere in the *WUS* promoter, suggesting that *AG* acts indirectly to repress *WUS* expression. It is plausible that multimers of these short fragments are required for detectable levels of transcription because of the lack of transcriptional enhancers present in the natural promoter context, such as those identified by the internal deletions in the region −2114 to −1360.

Notably, the portion of *GUS*-positive plants among all primary transformants was lower with these short synthetic *WUS* promoter constructs than with full-length or nearly full-length *WUS* promoter constructs (see Supplemental Table 1 online). This finding might reflect a stronger influence of the transgene's integration site in the genome on the expression levels of short compared with longer constructs.

We subsequently analyzed a series of reporter genes that carried tetrameric tandem repeats of D5 subfragments (Figure 1D) and identified a 57-bp region named L5 (−586/−529), which provided the correct spatial *WUS* expression pattern in the stem cell niche of the inflorescence meristem (Figure 3L), the vegetative meristem (Figure 3E), and in floral meristems, albeit at a lower level (Figure 3F). Comparison of the expression levels provided by subfragments G1, L2, and L6 (Figures 1D and 3D) revealed that the presence of a 29-bp interval (−498/−469) adjacent to the 57-bp region is required for increased expression but is not sufficient to confer expression on its own (Figures 1D and 4).

Thus, a 57-bp promoter fragment (−586/−529) provides all of the spatial and temporal information necessary for *WUS* transcription in the stem cell niche of shoot and floral meristems.

**Figure 4.** Regulatory Architecture of the *WUS* Promoter.

The approximate positions of regulatory domains are indicated. At bottom, nucleotides within the 57-bp spatial control region essential for promoter activity in the stem cell niche of the inflorescence meristem (RE1 and RE2) are indicated in boldface letters; the predicted HD-ZIP binding site is underlined. QE, quantitative element required for enhanced expression levels.

Two Distinct Sequence Motifs within the 57-bp Regulatory Region Are Essential for WUS Transcription in the Stem Cell Niche

To further define the regulatory sequences present in the 57-bp fragment, we performed linker-scanning mutagenesis using the -586/-469 fragment of the *WUS* promoter, named S0 (for scanning fragment 0), which contained both the 57-bp fragment and the neighboring putative enhancer element (Figure 1D). The S0 fragment was permuted by substituting 10-bp elements with always the same unrelated 10-bp sequence. The resulting mutated promoter fragments were used to create 12 tetrameric *GUS* reporter constructs: (S1)₄:*GUS* to (S12)₄:*GUS* (Figure 5). Plants carrying the unmutated (S0)₄:*GUS* reporter or any 1 of 10 of the mutated reporters [(S1)₄:*GUS*, (S2)₄:*GUS*, (S4)₄:*GUS*, (S6)₄:*GUS* to (S12)₄:*GUS*] displayed *GUS* expression in the inflorescence and floral meristems similar to that of the (D5)₄:*GUS* reporter gene (Figures 3G and 5). By contrast, two mutated constructs, (S3)₄:*GUS* and (S5)₄:*GUS*, in which sequences -566 to -557 and -546 to -537, respectively, had been exchanged, had completely lost promoter activity (Figure 5). By introducing the respective mutations into the *HindBst* reporter gene (*S3-HindBst* and *S5-HindBst*; Figure 5), we confirmed that the two decamers mutated in these constructs are also necessary for *GUS* expression in the inflorescence meristem in the context of the full-length *WUS* promoter.

Next, we replaced trinucleotide and tetranucleotide motifs within the two decamers in the context of the tetrameric (S0)₄:

GUS gain-of-function construct to further restrict the essential *cis*-regulatory sequences (Figure 5). Nucleotides -566/-564 (M31; Figure 3H), -560/-557 (M33), and -546/-544 (M51) were absolutely essential for reporter gene activity in the shoot and floral meristems. Mutating the nucleotides -563/-561 (M32) and -543/-541 (M52) reduced reporter expression in the shoot and floral meristems but did not abolish it, indicating that these nucleotides are important but not essential. By contrast, replacing the nucleotides -540/-537 (M53) did not affect reporter activity. When we introduced the same mutations into the 8.7-kb *HindBst* reporter construct, we obtained analogous results, confirming that the identified nucleotides are also essential for expression in the shoot meristem in the context of the full-length *WUS* promoter (Figure 5). However, these constructs retained expression in the floral meristems, as expected from the presence of a redundant floral meristem-specific control element between -3308 and -2114 (see above).

Together, these results indicate that two distinct sequence motifs that we named RE1 (for regulatory element 1; -566/-557) and RE2 (-546/-541) regulate the boundaries of the *WUS* expression domain in the stem cell niche of shoot and floral meristems (Figure 4).

We noticed that the region mutated in S2 (-576/-567) and S3 (-566/-557) contains a sequence motif (TAATAATTG, -572/-564; Figure 4) similar to the consensus binding site for several HD-ZIP proteins (CAATNATTG) (Johannesson et al., 2001). Because only the S3 but not the S2 mutation affected promoter activity of the tetrameric promoter constructs, we introduced the

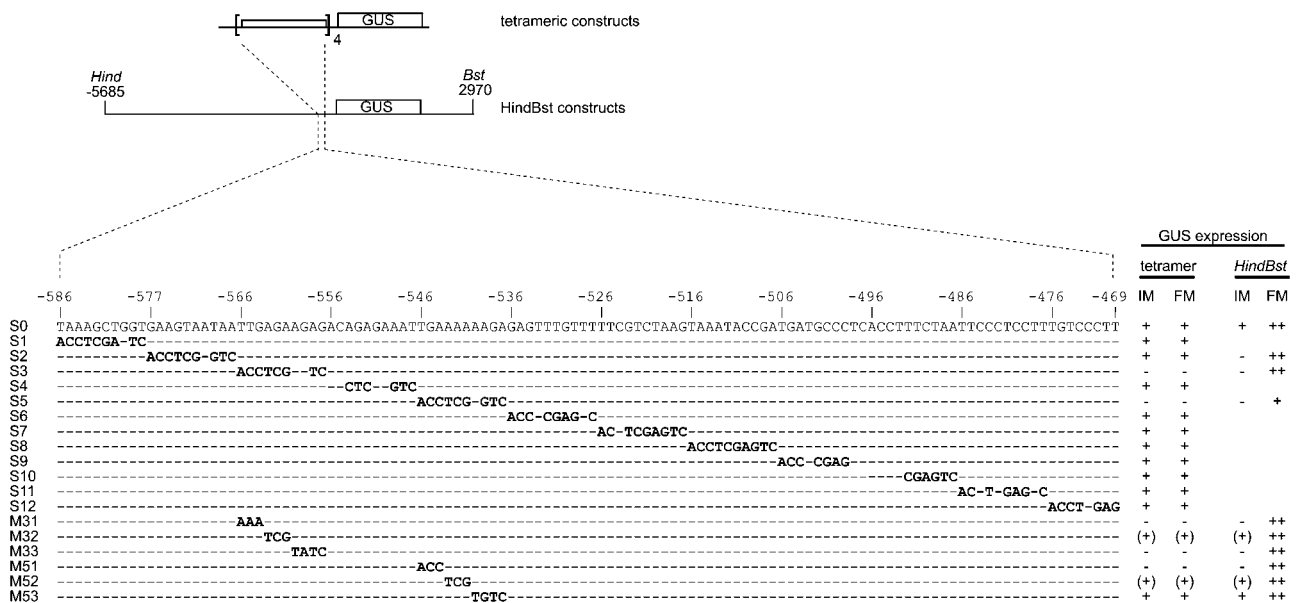


Figure 5. Linker Scanning Analysis.

Schemes of the constructs analyzed. The 118-bp *WUS* promoter fragment S0 (-586/-469) was permuted with the decamer sequence ACCTCGAGTC, generating the mutated fragments S1 to S12. The -566/-557 and -546/-537 regions were also scanned with trinucleotide/tetranucleotide exchanges (M31 to M33 and M51 to M53). For the reporter constructs, each mutated fragment was tetramerized and fused to -60 *CaMV*:*GUS*. Unaltered nucleotides are indicated with dashes. Relative staining intensities in inflorescence meristems (IM) and floral meristems (FM) are indicated at right for tetrameric (tetramer) and full-length *WUS* promoter constructs (*HindBst*). All full-length promoter constructs additionally showed strong staining in ovules unaffected by the indicated mutations.

S2 mutation, which covers the major part of this putative binding site, into the full-length *WUS* promoter. Indeed, no expression in the inflorescence meristem was detected with this *S2-HindBst:GUS* reporter gene (Figure 5), suggesting an essential function of the S2 region in the context of the full-length promoter.

DISCUSSION

The ability to stably maintain multipotent stem cells is crucial for the postembryonic production of new cells in plants and animals. In the plant shoot meristem, the stem cells are specified by *WUS*-dependent signals from underlying organizing center cells, and transcriptional control of the *WUS* gene within the proliferating shoot apex is a key regulatory switch in stem cell regulation. To gain insight into the mechanisms of how the boundaries and the position of the stem cell niche are stably maintained, we identified sequences within the *WUS* promoter that control the spatial and temporal transcription pattern.

Regulatory Domains of the *WUS* Promoter in Stem Cell Control

Our results show that the *WUS* promoter contains distinct regulatory regions that control tissue specificity and levels of transcription in a combinatorial manner (Figure 4). Among them, a 57-bp region ~550 bp upstream of the putative transcription start provides all information necessary for the correct spatial and temporal transcriptional pattern in the stem cell niches of shoot and floral meristems. Because all other nucleotides within this region were dispensable, two short sequence motifs, RE1 and RE2, mediate this control. In fact, these elements are highly conserved in *WUS* promoter sequences throughout the Brassicaceae family, supporting a central role in stem cell niche transcription (E. Tucker and T. Laux, unpublished results).

Interestingly, the first three nucleotides (TTG) of RE1 overlap with an HD-ZIP consensus binding site-like motif. HD-ZIP proteins can form heteromeric combinations involved in a variety of developmental processes (Johannesson et al., 2001), raising the possibility that HD-ZIP proteins might be involved in the spatial control of *WUS* transcription in the stem cell niche. In accordance with this possibility, a novel HD-ZIP-related protein was isolated in further analysis that specifically binds to this consensus sequence in the *WUS* promoter (I. Bäurle and T. Laux, unpublished results). However, because this consensus-like sequence is only necessary in the context of the full-length promoter but not in multimerized short promoter fragments, we hypothesize that factors binding to it do not mediate the spatial regulation of *WUS* promoter activity but rather enhance transcription levels, the requirement of which can be bypassed by increased copy numbers of RE1 and RE2.

Notably, the regulation of the stem cell niche in shoot and floral meristems, which are homologous systems and share several regulatory mechanisms (Steeves and Sussex, 1989; Schoof et al., 2000), involves not only common but also meristem-type-specific regulatory regions. For example, the 57-bp regulatory region containing RE1 and RE2 is sufficient for the spatial expression of *WUS* in both meristems, but its deletion from the full-length promoter abolishes *WUS* expression only in the shoot

meristem, indicating the presence of redundant *cis* elements that function exclusively in the stem cell niche of the floral meristem. It is conceivable that such differentially used promoter elements might account for differences in growth dynamics, gene expression levels, and temporal control of shoot and floral meristems and have evolved during diversification of their developmental programs.

The expression levels of synthetic tandem repeat promoter constructs containing RE1 and RE2 appear to be highly dependent on the site of integration within the genome, suggesting that their efficacy requires a favorable chromatin state at the integration site. Because this is not the case for constructs with the full-length promoter, some of the identified regulatory regions could act in chromatin organization, such as scaffold attachment (Breyne et al., 1992), nucleosome position and conformation, or recruitment of histone-modifying enzymes (Wagner, 2003) at the *WUS* locus. In fact, the boundaries of the *WUS* expression domain are deregulated in a mutant with compromised histone acetyltransferase activity, although it has yet to be determined whether the *WUS* promoter is directly affected in this case (Bertrand et al., 2003).

Integration of Regulatory Inputs in Stem Cell Control

The spatial and temporal control of the stem cell niche requires the integration of different cues at the level of *WUS* gene expression. One surprising result of this study is that all of these regulatory cues converge at two adjacent small regulatory sites, RE1 and RE2, of the *WUS* promoter. How is this achieved? Two mutually nonexclusive mechanisms can be envisioned. First, different combinations of transcription factors mediating independent regulatory inputs could bind to these motifs alternately or in a combinatorial way. Conversely, different cues could modify the activity of a common central transcription complex. The repression of *WUS* transcription via the stem cell-borne *CLV3* signal could be an example of the latter case. In *clv3* loss-of-function mutants, the *WUS* expression domain is increased from the embryo stage on (Brand et al., 2000; Schoof et al., 2000). However, we did not find a promoter mutation that altered *WUS* expression as expected for a construct lacking *CLV3*-dependent repression or any duplicated sequence motif that would suggest redundancy of such a putative negative regulatory element. Thus, a plausible mechanism for *CLV3* action on *WUS* transcription could involve the phosphorylation of RE1- or RE2-specific transcription factors by the intracellular receptor kinase signaling pathway activated by *CLV3* (Clark, 2001). The knowledge of *WUS*-regulating *cis* elements reported here provides the basis for the search for direct upstream regulators that will eventually allow insight into how transcriptional domains are stably maintained within a changing cellular context of the proliferating shoot apex.

METHODS

Plant Material and Plant Transformation

Arabidopsis thaliana growth conditions and the *wus-1* allele used for the complementation experiments have been described (Laux et al., 1996). The *clv1-4* allele was also described previously (Clark et al., 1993). All

plasmids were introduced into *Agrobacterium tumefaciens* strain GV3101(pMP90) (Koncz and Schell, 1986) by electroporation and transformed into Landsberg *erecta* (*Ler*) wild-type plants by the floral dip method (Clough and Bent, 1998).

GUS Staining

GUS staining was performed as described (Schoof et al., 2000). Material was cleared in 70% ethanol before taking photographs using a Leica MZ12 binocular and a Leica DC300 camera (Leica Microsystems, Wetzlar, Germany). For sections, tissue was dehydrated in an ethanol series up to 50%, postfixed in FAA (50% ethanol:5% formaldehyde:10% acetic acid) for 30 min at room temperature, dehydrated completely, and embedded in Paraplast (Sigma-Aldrich, Taufkirchen, Germany).

Cloning Details

Cloning details are available upon request. For simplicity, fragment names derived from restriction sites are abbreviated as follows: *Hind*III as *Hind*, *Bst*11071 as *Bst*, *Scal* as *Sca*.

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Isabel Bäurle and Thomas Laux

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