# Spatiotemporal control of axillary meristem formation by interacting transcriptional regulators

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**KEYWORDS:** axillary meristem, branching, stem cell, transcription **SUMMARY STATEMENT** 

Shoot branching utilizes interacting transcriptional regulators to fine-tune the spatiotemporal expression of *STM*, and thus precisely regulate axillary meristem initiation.

#### **ABSTRACT**

Branching is a common feature of plant development. In seed plants, axillary meristems (AMs) initiate in leaf axils to enable lateral shoot branching. AM initiation requires a high level of expression of the meristem marker SHOOT MERISTEMLESS (STM) in the leaf axil. Here, we show that modules of interacting transcriptional regulators control STM expression and AM initiation. Two redundant AP2-type transcription factors, DORNRÖSCHEN (DRN) and DORNRÖSCHEN-LIKE (DRNL), control AM initiation by regulating STM expression. DRN and DRNL directly up-regulate STM expression in leaf axil meristematic cells, as does another transcription factor REVOLUTA (REV). The activation of STM expression by DRN/DRNL depends on REV, and vice versa. DRN/DRNL and REV have overlapping expression patterns and protein interactions in the leaf axil, which are required for the up-regulation of STM expression. Furthermore, LITTLE ZIPPER3, another REV-interacting protein, is expressed in the leaf axil and interferes with the DRN/DRNL-REV interaction to negatively modulate *STM* expression. Our results support a model in which interacting transcriptional regulators fine-tune the expression of STM to precisely regulate AM initiation. Thus, shoot branching recruits the same conserved protein complexes used in embryogenesis and leaf polarity patterning.

#### INTRODUCTION

In contrast to animals, plants are sessile organisms with an enormous developmental plasticity to adapt to the changing environment. To this end, plants innovated a branching growth habit (Coudert et al., 2015). In seed plants, shoot branches arise from axillary meristems (AMs, also termed lateral meristems) in, or near, the leaf axils (Schmitz and Theres, 2005; Wang and Jiao, 2018). Axillary bud development comprises two stages: initiation in the leaf axil and subsequent outgrowth or dormancy. The final pattern of branches to a large extent determines the architecture of the shoot system.

Recent study has shown that AMs initiate from a group of leaf axil cells that constitute a meristematic cell lineage (Burian et al., 2016; Shi et al., 2016).

These meristematic cells continuously express the meristem marker *SHOOT MERISTEMLESS* (*STM*). Whereas a low level of *STM* expression maintains meristematic competence, high levels of expression lead to AM initiation (Greb et al., 2003; Long and Barton, 2000; Shi et al., 2016). The maintenance of a low level of *STM* expression requires a low auxin concentration and response (Wang et al., 2014a; Wang et al., 2014b). Before the formation of axillary buds, REVOLUTA (REV) up-regulates *STM* expression to promote AM initiation (Shi et al., 2016). Subsequently, cytokinin activates *WUSCHEL* expression *de novo* to establish the AM (Wang et al., 2017). In addition, genetic studies have identified several transcription factor-encoding genes that regulate AM initiation in *Arabidopsis*, including *LATERAL SUPPRESSOR* (*LAS*),

REGULATOR OF AXILLARY MERISTEMS (RAX), CUP-SHAPED

COTYLEDON (CUC) and REGULATOR OF AXILLARY MERISTEM

FORMATION (ROX) (Greb et al., 2003; Hibara et al., 2006; Müller et al., 2006;

Raman et al., 2008; Yang et al., 2012). Genetic and molecular studies have revealed direct and indirect interactions among these genes to form a regulatory network (Raman et al., 2008; Tian et al., 2014).

REV belongs to the class III homeodomain-leucine zipper (HD-ZIPIII) family of transcription factors and plays pleiotropic roles in embryo, meristem, leaf, and vascular development, including AM initiation (Emery et al., 2003; Huang et al., 2014; Otsuga et al., 2001; Prigge et al., 2005; Talbert et al., 1995; Zhong and Ye, 1999). During leaf polarity patterning, the activity of REV and related HD-ZIPIII proteins is inhibited by microRNAs 165/166 (Emery et al., 2003; Mallory et al., 2004), and by interacting LITTLE ZIPPER (ZPR)-type microProteins (Kim et al., 2008; Wenkel et al., 2007).

The AP2 family transcription factors *DORNRÖSCHEN* (*DRN*), also named *ENHANCER OF SHOOT REGENERATION 1* (*ESR1*), and the related *DORNRÖSCHEN-LIKE* (*DRNL/ESR2*) also function in embryonic meristem and lateral organ development (Banno et al., 2001; Capua and Eshed, 2017; Chandler et al., 2007; Cole et al., 2009; Ikeda et al., 2006; Kirch et al., 2003; Nag et al., 2007). During embryogenesis, both DRN and DRNL heterodimerise with HD-ZIPIII proteins (Chandler et al., 2007). We have recently shown that AM initiation is compromised in the *drn-1* mutant (Tian et al., 2014),

highlighting a novel function of DRN.

In this study, we show that DRN and DRNL redundantly promote AM initiation during the vegetative phase and show that DRN/DRNL and REV coordinately up-regulate *STM* transcription in mature leaf axils. In early leaf development, ZPR3 is strongly expressed and may destabilize the DRN/DRNL—REV interaction in leaf axils, resulting in a low level of *STM* expression. These findings emphasize the dynamic interaction of transcriptional regulators as a core feature of developmental control. In addition, we show that the same DRN/DRNL—REV and ZRP3—REV interactions are shared by AM initiation, embryo development, and leaf patterning, although with different downstream targets.

#### RESULTS

## DRN and DRNL redundantly control axillary bud formation

We have recently shown via a genome-wide study that *DRN* is required for AM initiation (Tian et al., 2014). This current study aimed to understand in more detail how *DRN* regulates the formation of AMs in rosette leaf axils in the vegetative shoots. Firstly, we tested whether the *DRN* paralogue *DRNL* is also required for this process. In wild-type *Arabidopsis* plants, the first visible evidence of AM formation by scanning electron microscopy was a cluster of

small and proliferating cells at the adaxial leaf base (Fig. 1A). However, these proliferating cell clusters were absent in most early rosette leaf axils in *dm-1*, *dml-1*, *dml-2*, and *dm-1 dml-1* mutants (Fig. 1B–E). The absence of expression of the corresponding genes was confirmed (Fig. S1). In wild-type plants grown in short days, axillary buds develop from each rosette leaf, except from the two cotyledons and some of the first-formed true leaves (Fig. 1F, G). In contrast, *dm* and *dml* mutants flower earlier and show a strong reduction in axillary bud formation, especially in rosette leaves formed in the early and mid-phase of vegetative development (Fig. 1F, G). The *dm-1 dml-1* double mutant shows more serious defects in axillary bud formation than either *dm* or *dml* single mutant (Fig. 1F, G), suggesting that *DRN* and *DRNL* have important redundant functions in AM initiation.

#### DRN/DRNL and REV genetically co-regulate AM initiation

During embryogenesis, REV can dimerize with DRN/DRNL (Chandler et al., 2007). To test whether DRN/DRNL–REV heterodimers are also recruited during AM initiation, we sought genetic evidence by constructing *drn-1 rev-6*, *drnl-2 rev-6*, and *drn-1 drnl-1 rev-6* double and triple mutants. Although short-day-grown *rev-6* loss-of-function plants showed a strong reduction in axillary bud formation (Otsuga et al., 2001; Talbert et al., 1995), axillary buds occasionally formed in the axils of cauline leaves and of rosette leaves formed during late vegetative development. The *drn-1 rev-6* and *drnl-2 rev-6* mutants showed further reductions in axillary bud formation. Additionally, the defect in

axillary bud formation in the drn-1 drnl-1 rev-6 mutant was almost completely penetrant (Fig. 1G). This increased phenotypic penetrance in double and triple mutant plants compared to rev single mutants suggests that DRN/DRNL and REV have combinatorially important functions that converge to form axillary meristems, but that they might also provide individual contributions. It has been proposed that AM initiation during vegetative and reproductive stages requires different sets of genes (Hempel and Feldman, 1994; Huang et al., 2012; Yang and Jiao, 2016). Compared to early rosette leaves, the AM defect becomes less severe in axils of later-initiated rosette leaves and cauline leaves in drn, drnl, and drn drnl mutants. Expression of pREV::REV-GR-HA could not complement the drn-1 drnl-1 AM defect from early rosette leaf axils, but resulted in slightly more AMs in late rosette leaves and cauline leaves (Fig. S2A). Consistently, p35S::DRN-GR could not complement the rev-6 AM defect from early rosette leaf axils (Fig. S2B). Thus, DRN and DRNL preferentially affect the AM initiation pathway during early vegetative stages, more than during later stages and reproduction, which affects cauline leaves and later-initiated rosette leaves.

## Expression patterns of *DRN* and *DRNL* in leaf axils

Because AMs initiate from organ boundary cells located at the adaxial side of leaf axils, we investigated whether *DRN* and *DRNL* are expressed in the leaf axils. We initially used *pDRN::GUS* and *pDRNL::GUS* reporter lines that recapitulate RNA *in situ* hybridization patterns (Kirch et al., 2003; Nag et al.,

and young primordia for both *pDRN::GUS* and *pDRNL::GUS* (Fig. S3). We subsequently analyzed the expression patterns of the *pDRN::DRN-GFP* and *pDRNL::DRNL-CFP* reporters (Chandler et al., 2011; Cole et al., 2009) in young leaf primordia, which were comparable to those of RNA *in situ* hybridization (Fig. 2A, B). We observed broad *pDRN::DRN-GFP* signals in leaf primordia, including boundary cells (Fig. 2A, Fig. S4C). In addition, DRN was very strongly expressed in AMs (Fig. S4C). A similar pattern of *pDRNL::DRNL-CFP* expression was detected in primordia, boundary cells and AMs (Fig. 2B, Fig. S4D). Similar to *DRN* and *DRNL*, *REV* was also expressed in leaf axils (Fig. 2C) and this expression was relatively low in leaves after P<sub>8</sub> (Shi et al., 2016). In contrast to the expression of *DRN* and *DRNL*, that of *REV* was more adaxial-specific (Otsuga et al., 2001).

We next analysed *DRN* and *DRNL* expression in older leaf axils. Imaging of *pDRN::DRN-GFP* in leaves showed that GFP signal was absent from the adaxial side of the leaf axil between P<sub>8</sub> and P<sub>9</sub> stages (Fig. 2D, Fig. S4A). Starting from P<sub>10</sub>, the GFP signal became restricted to the center of the leaf axils (Fig. 2E, F, Fig. S4B), the site of presumptive AM initiation. We also observed similar dynamic expression of *pDRNL::DRNL-CFP* in leaf axils (Fig. 2G–I). The expression of *REV* was similarly restricted to the center of leaf axils in P<sub>9</sub> and older leaves (Shi et al., 2016). The enrichment of *DRN*, *DRNL* and *REV* expression in leaf axils is consistent with the up-regulation of *STM* 

expression in  $P_{10}$  and older leaves, which is critical for AM initiation (Shi et al., 2016).

## DRN and DRNL regulate STM expression

AM initiation requires cells that continuously express STM. A low level of STM expression maintains meristematic competence, whereas high STM expression leads to AM initiation (Greb et al., 2003; Long and Barton, 2000; Shi et al., 2016). In wild-type plants, the expression of STM persists in young leaf axils (Fig. 3A). We found that STM expression was maintained in drn-1, drnl-2, and drn-1 drnl-1 mutants, but was much lower than in wild-type plants (Fig. 3B–D, Fig. S5). In wild-type, the number of STM-expressing cells and the level of STM expression increased from the P<sub>11</sub> stage, just prior to the stage at which AMs become morphologically visible (Fig. 3E) (Shi et al., 2016). However, STM was not upregulated in the mutants during leaf maturation (Fig. 3F–H, Fig. S5). The level of STM expression in drn and drnl mutants was similar to that in the rev-6 mutant (Shi et al., 2016) (Fig. 3I, J). Reverse transcription quantitative PCR (RT-qPCR) analysis also confirmed the up-regulation of STM by DRN and DRNL. The level of STM expression in shoot apices that are enriched with leaf axils by leaf removal, was significantly reduced in *drn-1*, *drnl-2*, and *drn-1 drnl-1* plants compared to in wild-type plants (Fig. 3K). This low level of STM expression in leaf axils is insufficient for AM initiation (Shi et al., 2016), and thus explains the AM initiation defects in drn and drnl mutants.

Previous studies have shown that ectopic STM activity can induce meristems from undifferentiated (and presumably meristematic) cells, but not differentiated cells (Brand et al., 2002; Shi et al., 2016). To test whether leaf axil cells in drn and drnl mutants remain meristematic cells that are competent to respond to STM activity, we introduced p35S::STM-GR into drn-1, drnl-2 and drn-1 drnl-1 plants. In these plants, dexamethasone (Dex) can induce the nuclear translocation of a STM-glucocorticoid-receptor (GR) fusion protein to activate STM function. We found that axillary buds could be induced from young and mature leaf axils in these mutants by Dex application, although at a slightly lower frequency in the drn-1 drnl-1 double mutant (Fig. 4). REV also up-regulates STM expression to promote AM initiation. Induced STM expression can similarly complement the rev mutant phenotype in bud formation (Shi et al., 2016). Genetically, STM overexpression also complemented branch suppression in the drn-1 drnl-1 rev-6 triple mutant (Fig. S6). Thus, leaf axil cells in drn-1, drnl-2, as well as drn-1 drnl-1 that express STM at a low level, remain meristematic, as observed in rev-6 (Shi et al., 2016).

## DRN and DRNL directly activate *STM* expression

To test whether DRN can directly activate *STM* expression, we generated Dex-inducible *p35S::DRN-GR* lines and measured the effect of DRN activation on the expression of *STM* by RT-qPCR. DRN activation resulted in a rapid increase in *STM* mRNA levels within 2 h of treatment in the presence or

absence of the protein synthesis inhibitor cycloheximide (CHX) (Fig. 5B).

These results suggest that induction of *STM* does not require *de novo* protein synthesis and that *STM* is probably a direct target of DRN.

We next performed chromatin immunoprecipitation (ChIP) assays using leaf axil-enriched shoot tissues to examine whether DRN and DRNL directly bind to the *STM* promoter *in vivo*. We designed primers upstream of the start codon that covered the ~2.6 kb region or spanned the start codon (Fig. 5A). Using antibodies against GFP, we found that DRN-GFP strongly associated with a region containing the start codon and a GCCGCC motif (GCC box), the conserved binding site for DRN and related AP2-type transcription factors (Banno et al., 2006), in vegetative-stage shoot apex tissue that was enriched for axils by leaf removal (Fig. 5C). Similarly, we found that DRNL-CFP could associate with the same region (Fig. 5D), which we term fragment 1. It should be noted that fragment 1 also contains multiple ATGAT motifs, which are the conserved binding site for REV and we recently demonstrated that REV also binds to fragment 1 (Shi et al., 2016).

A transient transfection assay in protoplasts further confirmed that DRN and DRNL could bind to *STM* genomic regions, especially fragment 1, and up-regulate *STM* expression (Fig. 5E). Although both DRN and DRNL activated a 3.0 kb *STM* promoter-driven *Luciferase* reporter gene (*pSTM::Luc*), the activation of *pSTM*2::*Luc*, in which fragment 1 was replaced with a CaMV 35S minimal promoter (contains the TATA box), was substantially reduced.

To further confirm the importance of fragment 1 for STM expression, we constructed a GUS reporter (pSTM::GUS) containing 6.3-kb of the STM promoter upstream of the start codon, which also included fragment 1. We also constructed a pSTM\(\textit{2}::GUS\) reporter, in which fragment 1 was replaced by a CaMV 35S minimal promoter. From multiple independent transgenic lines (>5), we repeatedly found that *pSTM::GUS* expression recapitulated RNA *in situ* patterns, with strong expression in the shoot apex and leaf axils (Fig. 5F, Fig. S7A-D). By contrast, pSTM\(\Delta::GUS\) plants showed barely detectable GUS signals in these tissues (Fig. 5G, Fig. S7E-H), which might be partly due to the inhibitory *cis*-element K-box and RB-box remaining in the promoter (Aguilar-Martinez et al., 2015; Uchida et al., 2007). To directly analyze the contribution of fragment 1 to STM expression, we also generated a fragment 1::GUS reporter, in which GUS expression remained detectable but was less enriched in the leaf axil compared to that using the 6.3-kb promoter (Fig. 5H, Fig. S7I–L). We therefore conclude that fragment 1, which is bound by DRN, DRNL and REV, is critical for *STM* expression.

#### DRN and DRNL interact with REV to activate *STM* expression

DRN, DRNL and REV can directly activate *STM* expression by binding to the same promoter region, and these transcription factors significantly overlap in expression. Similar to *rev* mutants, *drn* and *drnl* mutants have AM initiation defects. During embryogenesis, DRN/DRNL can physically interact with REV (Chandler et al., 2007). During AM initiation, *REV* is also expressed at

progressively higher levels in the leaf axil (Shi et al., 2016), in a similar way to *DRN* and *DRNL*. We therefore speculate that the DRN/DRNL–REV interaction occurs during AM initiation, and is critical for the activation of *STM* expression.

To examine the interaction between DRN and REV during *in vivo* development, we performed co-immunoprecipitation (Co-IP) using *pDRN::DRN-GFP p35S::REV-MYC* plants. Using leaf axil-enriched shoot tissues, we detected interaction between DRN-GFP and REV-MYC when immunoprecipitated with an anti-GFP antibody and probed with an anti-MYC antibody. Furthermore, using the same amount of DRN-GFP, we found that the band indicating the amount of (35S-driven) REV-MYC was about three-fold stronger in old leaf axil-enriched tissues than in young leaf axil-enriched tissues (Fig. 6A).

To test whether the detected DRN/DRNL–REV interaction is required for the activation of *STM* expression, we introduced *pREV::REV-GR-HA* into the *drn-1 drnl-1* background and measured the effect of REV activation on *STM* expression, which was substantially reduced compared to that in wild-type siblings (Fig. 6B). Consistently, the effect of DRN activation on *STM* expression was also compromised in the *rev-6* background in comparison to that in wild-type siblings (Fig. 6C).

A transient transfection assay in protoplasts also confirmed that the DRN/DRNL–REV interaction can up-regulate *STM* expression (Fig. 6D).

Although transformation with only DRN, DRNL or REV resulted in the activation of *STM* expression (Fig. 5E, Fig. 6D), co-transformation of DRN and REV or DRNL and REV resulted in a ~2.5-fold stronger activation of *STM* expression (Fig. 6D).

Furthermore, our results show that interaction between DRN/DRNL and REV is critical for their binding to the *STM* promoter region. Using the same experimental set-up, we found that both DRN-GFP and DRNL-CFP showed reduced association with fragment 1 of the *STM* promoter region in the *rev-6* background than in wild-type plants (Fig. 6E, F, compare with Fig. 5C, D). Similarly, REV-GR-HA showed weaker binding to the *STM* promoter in the *drn-1 drnl-1* background than in wild-type siblings (compare Fig. 6G, H). These results indicate that the DRN/DRNL–REV interaction is recruited during AM initiation, and this interaction cooperatively promotes binding of the proteins to the *STM* genomic region. Taken together, our data demonstrate that DRN/DRNL and REV function as a complex to promote the expression of downstream target genes during AM initiation from early rosette leaf axils.

#### ZPR3 destabilizes the DRN/DRNL-REV interaction

During leaf polarity patterning, REV can interact with ZPR proteins and this interaction inhibits REV function (Kim et al., 2008; Wenkel et al., 2007). We speculated that ZPR proteins might also participate in the regulation of AM initiation. To this end, we first analyzed the tissue-specific expression pattern of

ZPR3 and ZPR4 and that of the related genes ZPR1 and ZPR2 by RT-qPCR in various organs. ZPR genes were more highly expressed in boundary-enriched tissues such as the shoot apex and flowers than in leaves (Fig. S8A, B). ZPR expression was analysed at different stages of primordium development, and was higher in young leaves and decreased in mature leaves where AMs will initiate (Fig. S8C). Using a pZPR3::GUS reporter (Wenkel et al., 2007), we detected GUS activity in the adaxial domain of leaf primordia, including the leaf axils (Fig. 7A, Fig. S8D–H). Notably, GUS activity in the leaf axils was substantially reduced in P10 and older leaves. The reduction in ZPR3 leaf axil expression during leaf maturation is consistent with the up-regulation of STM expression beginning in P11, prior to the morphological appearance of axillary buds (Greb et al., 2003; Shi et al., 2016). Overexpression of ZPR3 led to fewer axillary buds in the axils of cauline and rosette leaves in p35S::ZPR3 transgenic lines (Fig. S8I–K).

We next tested whether the ZPR3 protein inhibits the DRN/DRNL–REV interaction to inhibit downstream gene expression, using the yeast three-hybrid (Y3H) assay. In yeast, the AP2 domain of DRN or DRNL fused to the activation domain of GAL4 (AD-DRNAP2 or AD-DRNLAP2) can interact with the PAS domain of REV fused to the binding domain of GAL4 (BD-REVPAS) (Fig. 7B), as previously reported (Chandler et al., 2007). Similar results were obtained using full-length DRN/DRNL and REV (Fig. S9). In addition, AD-ZPR3 interacted with BD-REV in yeast, as shown previously (Kim et al., 2008; Wenkel et al., 2007).

The interaction between AD-DRN or AD-DRNL and BD-REV was reduced in the presence of ZPR3 (Fig. 7B, C). Similarly, the introduction of DRN or DRNL in the Y3H assay interfered with the REV–ZPR3 interaction (Fig. S9). To confirm that DRN/DRNL competes with ZPR3 to interact with REV, we also introduced DRN or DRNL without the REV-interacting AP2 domain (Chandler et al., 2007). The truncated DRNΔAP2 and DRNLΔAP2 versions were no longer able to disrupt the ZPR3–REV interaction (Fig. S9).

Using transient transfection assays in protoplasts, we not only confirmed the competition between DRN/DRNL-REV and ZPR3-REV interactions, but also showed that ZPR3 inhibits the DRN/DRNL-REV activation of STM expression. When ZPR3 was co-transformed into protoplasts with REV, the activation of STM was substantially reduced, leading to significantly reduced Luc expression (Fig. 7D). However, co-transformation of DRN/DRNL, ZPR3 and REV fully restored Luc expression to a higher level but still much less than that following REV-DRN or REV-DRNL co-transformation (Fig. 7D, and compare with Fig. 6D). To analyze the effect of DRN-REV interaction on DNA binding activity, we performed an electrophoretic mobility shift assay (EMSA) using biotin-labeled fragment 1 of the STM promoter. The addition of both DRN and REV resulted in a lower mobility than the addition of either single protein, suggesting supershift protein–DNA binding by heterodimerisation. ZPR3 decreased the intensity of the supershift band of DRN and REV (Fig. 7E), suggesting that ZPR3 interferes with the interaction between the DRN–REV

complex and the *STM* promoter fragment, probably through competition with DRN in binding to REV. The reduced *ZPR3* expression in leaf axils of P<sub>10</sub> and older leaves (Fig. 7A, Fig. S8D–H) correlates with, and can explain, the enhanced DRN–REV interaction in old leaf axil-enriched tissues (Fig. 6A). The strong DRN–REV interaction in mature leaves then explains the observed up-regulation of *STM* expression beginning in P<sub>11</sub> (Shi et al., 2016), just prior to the morphological appearance of axillary buds. In fact, overexpression of *ZPR3* leads to AM initiation defects (Fig. S8I–K), which is consistent with a previous report (Kim et al., 2008) and further confirms that reduced *ZPR3* expression in mature leaf axils is critical for the up-regulation of *STM* expression and subsequent AM initiation.

#### DISCUSSION

Shoot branching is fundamental to the radiation of plants, and is a key determinant of plant architecture (Coudert et al., 2015). In seed plants, shoot branching results from the lateral initiation of AMs and subsequently buds, which can become dormant until they perceive permissive environmental or internal cues to allow bud outgrowth and thereby, fine-tune development.

Although the outgrowth of axillary buds has been well studied, their initiation remains less well understood. Recent studies have shown that specification is an early event in which a population of *STM*-expressing meristematic cells is

precisely regulated to initiate AMs (Greb et al., 2003; Long and Barton, 2000; Shi et al., 2016). Although a low level of *STM* expression maintains meristematic competence, a high level of expression leads to meristem initiation, suggesting a threshold model (Shi et al., 2016). The fine tuning of *STM* expression is therefore critical for AM initiation.

In this study, we identified DRN and DRNL as redundant regulators of AM initiation (Fig. 1). Robust axillary bud formation in wild-type plants relies on DRN and DRNL functions. These two related AP2-family transcription factors have highly similar expression patterns in leaf axils (Fig. 2). Furthermore, DRN and DRNL directly promote STM expression by binding to its promoter (Figs. 3–5), which not only explains the roles of DRN/DRNL in AM initiation, but also their overexpression functions. The ectopic activation of STM can at least partially explain the observed ectopic shoot regeneration and the enlarged and disorganized SAM phenotypes following the constitutive overexpression of DRN or DRNL or in the *drn-D* mutant (Banno et al., 2001; Ikeda et al., 2006; Kirch et al., 2003). The transcription of STM is also regulated by additional unrelated cis-elements in its promoter (Aguilar-Martinez et al., 2015; Uchida et al., 2007). These *cis*-elements restrict *STM* expression, and are expected to function together with the cis-elements identified in this study to facilitate spatiotemporal STM expression. We and others have also shown that DRN and DRNL directly activate the expression of CUC genes (Ikeda et al., 2006; Matsuo et al., 2009; Tian et al., 2014), which might indirectly regulate STM

expression and affect AM initiation (Hibara et al., 2006; Raman et al., 2008). How these genes interact temporally and spatially during AM development, especially at reproductive stages, requires further study.

More importantly, our results demonstrate that a novel combination of interacting transcriptional regulators form a regulatory circuit that controls AM initiation during leaf maturation. We have provided in planta evidence that DRN/DRNL and REV form a protein complex in the leaf axils that binds to the STM promoter to fully up-regulate its expression (Fig. 6). In addition, ZPR3 competes with DRN/DRNL for access to REV and thereby titrates the DRN/DRNL-REV interaction (Fig. 7). Thus, the balance between ZPR3 and DRN/DRNL appears to be important for DRN/DRNL–REV complex formation and STM expression. Indeed, temporal changes in DRN, DRNL and ZPR3 expression can be monitored in the leaf axil (Figs. 2, 7). Early in leaf primordium development, ZPR3 is expressed in the leaf axil, where it might inhibit the formation of functional DRN/DRNL-REV complexes, resulting in a low level of STM expression and the absence of axillary bud formation. As the leaf matures, decreasing ZPR3 levels (Fig. 7A) would allow the formation of DRN/DRNL-REV complexes in more mature leaf axils (Fig. 6) to up-regulate STM expression and to promote axillary bud formation (Fig. 8). In an opposing manner, DRN and DRNL transcription is upregulated in the axils of leaves as they mature (Fig. 2). Hence, the competing interaction between DRN/DRNL and ZPR3 for REV, as well as their dynamic expression, provides a very

plausible scenario to explain the observed axillary bud formation process (Greb et al., 2003; Long and Barton, 2000). Further confirmation of the ZPR3–DRN/DRNL interaction in developmental processes requires further experiments, such as manipulating the temporal and spatial expression patterns of these proteins. However, our findings underscore the importance of protein–protein interactions as a recurring feature in transcriptional regulatory networks (Brady et al., 2011).

This study has revealed that the DRN/DRNL–REV module, involved in embryogenesis (Chandler et al., 2007), and the ZPR–REV module, involved in leaf polarity patterning (Kim et al., 2008; Wenkel et al., 2007), are recruited and combinatorially regulate AM initiation. Although AM initiation, embryogenesis, and leaf polarity patterning are clearly distinct developmental processes, they appear to involve conserved regulatory modules. Modifications of ancestral regulatory modules to create novel expression domains may lead to new combinations of regulatory modules and thus to regulatory neofunctionalization (Rosin and Kramer, 2009).

#### MATERIALS AND METHODS

## Plant materials and generation of transgenic plants

Arabidopsis thaliana ecotypes Col-0 and Ler were used as wild-type controls.

The drn-1, rev-6, pREV::REV-GR-HA, pDRN::GUS, pDRNL::GUS,

pZPR3::GUS, p35S::ZPR3, pDRN::DRN-GFP, pDRN::erGFP and

pDRNL::DRNL-CFP lines are in the Col-0 background (Chandler et al., 2011; Cole et al., 2009; Kirch et al., 2003; Nag et al., 2007; Otsuga et al., 2001; Wenkel et al., 2007), and the drnl-1, drnl-2, pREV::REV-Venus, and pSTM::STM-Venus lines are in the Ler background (Chandler et al., 2007; Heisler et al., 2005; Nag et al., 2007). Both drn-1 and drnl-1 are insertion mutants. The position of the insertion in the drn-1 allele is after nucleotide +327 (relative to the ATG) of the DRN gene and for drnl-1 it is after nucleotide +777. The drnl-2 allele has a base substitution from C to T at position +278, resulting in an A to V substitution at amino acid 93. The rev-6 allele has an premature stop codon after R346. Plants were grown in the greenhouse on soil at 22°C under short-day conditions (8 h light/16 h dark) for 28 to 30 d and were induced to flower under long-day conditions (16 h light/8 h dark) for 30 d unless otherwise specified.

The *p35S::DRN-GR* construct was created by inserting the *DRN* coding sequence amplified from cDNA in-frame upstream of the GR-coding sequence in the *pGREEN0229-35S::GR* vector; the T3 generation of homozygous plants of *p35S::DRN-GR* and *pREV::REV-GR-HA* were used for genotyping; *pSTM::GUS* was constructed using an amplified 6.3-kb fragment upstream of the *STM* coding region. To construct *pSTM*Δ::GUS, region 1 (Fig. 5A) was removed, to result in a 6.0-kb fragment. All constructs were transformed into Col-0 plants using the *Agrobacterium*-mediated floral dip method. Multiple transgenic lines (>20 for each construct) were obtained, and lines with

representative phenotypes or expression patterns were used for analysis.

#### Hormone treatment and RT-PCR

For Dex treatment, a 10-mM stock solution of Dex (Sigma-Aldrich) in ethanol was diluted with distilled water to a final concentration of 10 µM. Water with only ethanol was added to leaf axils as the mock control. For expression analyses, plants were grown for 21 d under short-day conditions and meristematic and boundary tissue was enriched for by the manual dissection of leaves from the shoot apex. Total RNA was extracted using the AxyPrep Multisource RNA Miniprep kit (Corning). First-strand cDNA synthesis was performed with 2 µg total RNA using TransScript One-Step gDNA Removal and cDNA synthesis SuperMix (TransGen) and 22-mer oligo dT primers according to the manufacturer's instructions. The RT-PCR analysis was performed in a 20-µL reaction volume using Taq DNA polymerase (TianGen) and gene-specific primers (Table S1). Quantitative RT-PCR (RT-qPCR) was performed on a Bio-Rad CFX96 real-time PCR detection system with a KAPA SYBR FAST qPCR kit (KAPA Biosystems). Relative RT-qPCR expression was normalized to that of TUB6 (At5g12250), which has been shown to be a superior reference gene for RT-qPCR analysis and shows constant expression in various treatments (Kaufmann et al., 2010; Han et al., 2014; Tian et al., 2014). The relative expression level of the positive control was transformed to a value of 1 and was double-normalized by the expression of the reference gene and by the ratio of the positive control. Data for qPCR are  $-\Delta\Delta$ Ct  $\pm$  SD of

three biological replicates, each performed in triplicate. Gene-specific primers (Table S1) were used to amplify and detect each gene.

Tissue preparation, confocal analysis and scanning electron microscopy Seedlings were grown in MS medium in short-day conditions (8 h light at 22°C and 16 h dark at 18°C) for 15 d after seed stratification. Leaves between P<sub>5</sub> and P<sub>11</sub> were then detached from seedlings, laid flat on MS medium and imaged. For sectioning, seedlings were grown in soil in short-day conditions for 21 d. Shoot apices were collected, the leaves were removed and the apices were immediately placed in 2.5% paraformaldehyde (PFA; Sigma-Aldrich) at pH 7.0 at 4°C, and were then vacuum-infiltrated for 30 min and stored overnight at 4°C. Fixed tissue samples were washed with 10% sucrose and 1% PFA at pH 7.0 for 20 min, with 20% sucrose and 1% PFA at pH 7.0 for 20 min, and 30% sucrose and 1% PFA at pH 7.0 for 30 min. Samples were then embedded in 5 to 7% LM agarose (Promega) liquid gel at 30°C and placed at 4°C for 15 min to solidify. Sections of 40 to 70 µm were made using a Leica VT1000S vibratome. For high-resolution images, samples were stained with 50 μg/mL propidium iodide (PI, Sigma-Aldrich). Images were taken with a Nikon A1 confocal and Leica SP5 microscope. Excitation and detection wavelengths for GFP, chlorophyll and Venus were as

Excitation and detection wavelengths for GFP, chlorophyll and Venus were as previously described (Wang et al., 2014). To detect the signal of FM4-64 and propidium iodide staining, a 514-nm laser line was used for excitation and a 561-nm long-pass filter was used for detection. Maximum projection was used

in the Nikon A1 software or LAS AF Lite software.

Scanning electron microscopy was performed using a Hitachi S-3000N variable pressure scanning electron microscope after standard tissue preparation (Jin et al., 2011).

## Chromatin immunoprecipitation and gel-shift assay

ChIP experiments were performed according to published protocols (Kaufmann et al., 2010). Shoots without leaves of ~28-d short-day-grown pREV::REV-GR-HA (induced with Dex for 4 h), pDRN::DRN-GFP and pDRNL::DRNL-CFP plants were harvested and fixed with 1% formaldehyde under vacuum for 10 min. Nuclei were isolated and lysed, and chromatin was sheared to a mean size of 1,000 bp by sonication. The sonicated chromatin served as an input or as a positive control. Immunoprecipitations were performed using an antibody against the glucocorticoid receptor (GR) (PA1 516; Affinity Bioreagents), or against GFP (11814460001, Roche). The precipitated DNA was isolated, purified, and used as a template for PCR. qPCR was performed as described above (see Table S1 for primers). The data are presented as the degree of enrichment of *STM* promoter fragments. The amount of precipitated DNA used in each assay was determined empirically, so that an equal amount of ACTIN2 (At3g18780) was amplified. Three independent biological replicates were performed.

The nucleotide sequences of the double-stranded oligonucleotides for EMSA were *STM* P1. The oligonucleotides were annealed and then labeled

with the Biotin 3´ End DNA Labeling Kit (Pierce). Standard reaction mixtures (20 mL) for EMSA contained 2 mg purified proteins, 2 mL biotin-labeled annealed oligonucleotides, 2 mL binding buffer (100 mM Tris, 500 mM KCl, 10 mM DTT, pH 7.5), 1 mL 50% glycerol, 1 mL 1% NP-40, 1 mL 1 M KCl, 1 mL 100 mM MgCl<sub>2</sub>, 1 mL 200 mM EDTA, 1 mL 1 mg mL<sup>-1</sup> poly (dl-dC) and 8 mL ultrapure water. The reactions were incubated at room temperature (25°C) for 20 min and loaded onto a 10% native polyacrylamide gel containing 45 mM Tris, 45 mM boric acid, 1 mM EDTA, pH 8.3. The gel was sandwiched and transferred to an N+ nylon membrane (Millipore) in 0.56TBE buffer at 380 mA in a 4°C refrigerator for 60 min. The detection of biotin-labeled DNA by chemiluminescence followed the manual of the LightShift Chemiluminescent EMSA Kit (PIERCE).

#### Protoplast transient expression assay

To produce the effector constructs, full-length *REV*, *DRN* and *DRNL* open reading frames were amplified from *Arabidopsis* cDNA and inserted into the pBI221 vector under control of the CaMV 35S promoter. To generate the *pSTM::Luc* reporter gene, the *STM* promoter was amplified from *Arabidopsis* genomic DNA and PCR fragments were inserted into the corresponding sites of the YY96 vector (Zhang et al., 2012) to produce *pSTM::Luc* and *pSTM*Δ::*Luc* (see Table S1 for primers). The YY96 vector contained a CaMV 35S minimal promoter before the *Luc* gene. The PEG-mediated transfection of *Arabidopsis* protoplasts was performed as described previously (Han et al.,

2014; Zhang et al., 2012). The reporter construct, effector plasmid, and a p35S::GUS construct (internal control) were co-transformed into protoplasts. After transformation, the protoplasts were incubated at 23°C for 12–15 h. The protoplasts were pelleted and resuspended in 100  $\mu$ L of 1 × CCLR buffer (Promega). For GUS enzymatic assays, 5  $\mu$ L of extract was incubated with 50  $\mu$ L 4-methylumbelliferyl- $\beta$ -d-glucuronide assay buffer (50 mM sodium phosphate pH 7.0, 1 mM  $\beta$ -d-glucuronide, 10 mM EDTA, 10 mM  $\beta$ -mercaptoethanol, 0.1% sarkosyl, 0.1% Triton X-100) at 37°C for 15 min, and the reaction was stopped by adding 945  $\mu$ L 0.2 M Na<sub>2</sub>CO<sub>3</sub>. For luciferase activity assays, 5  $\mu$ L of the extract was mixed with 50  $\mu$ L luciferase assay substrate (Promega), and the activity was detected using a Modulus Luminometer/Fluometer (Promega) and a luminescence kit. The reporter gene expression levels were expressed as relative LUC/GUS ratios. Three independent biological experiments were each performed in triplicate.

## Co-IP assay

Arabidopsis plants expressing REV-MYC and DRN-GFP were used in a Co-IP assay. Shoot apices of 1-week-old (younger axils than the tenth leaves) or leaf axils of 3-week-old (mature axils older than the tenth leaves) transgenic plants were ground in IP buffer (20 mM HEPES, pH 7.5, 40 mM KCl, 1 mM EDTA, and 1% Triton X-100), filtered, and centrifuged at 20,000 *g* for 10 min. Supernatants containing equal amounts of DRN-GFP were incubated with anti-GFP coupled to Protein A sepharose beads for 30 min. Beads were

washed four times with wash buffer (20 mM HEPES, pH 7.5, 40 mM KCl, and 0.1% Triton X-100), and bound proteins were eluted with 2 × SDS buffer.

### **Yeast Two-Hybrid and Three-Hybrid assays**

The yeast two-hybrid screens were performed as described (Zhang et al., 2012). To reduce autoactivation, 100 mM 3-AT (3-amino-1,2,4-triazole) was added to the selection medium. We detected no auto-activation of BD-REV in the presence of 3-AT. A yeast three-hybrid assay was performed according to the manufacturer's instructions; the yeast strain AH109 was transformed with pairs of plasmids (pB-REV-DRN/DRNL and pA-ZPR3).

Transformed colonies were selected on synthetic complete medium lacking Leu and Trp. Three independent clones with four respective replicates were used in each experiment. Cell cultures were placed under the indicated light conditions and incubated at 30°C until the OD $_{600}$  was between 0.5 and 0.8 with the conditional expression of the bridge proteins. The relative  $\beta$ -galactosidase activities were calculated as described previously (Ding et al., 2015). At least three independent experiments were performed and the result of one representative experiment is shown.

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## **Competing interests**

The authors declare no competing or financial interests.

#### **Author contributions**

Y.J. and C.Z. conceived and designed the experiments, C.Z. and J.W. performed experiments and analyzed data, S.W., J.W.C. W.W. and X.C. contributed reagents and material, and C.Z., Y.J. and J.W.C. wrote the manuscript.

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## **Data availability**

Sequence data from this article can be found in the *Arabidopsis* Genome initiative database under the following accession numbers: *At1g12980* (*DRN*), *At1g24590* (*DRNL*), *At1g62360* (*STM*), *At2g45450* (*ZPR1*), *At3g60890* (*ZPR2*), *At3g52770* (*ZPR3*), At2g36307 (*ZPR4*) and *At5g60690* (*REV*).

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# **Figures**

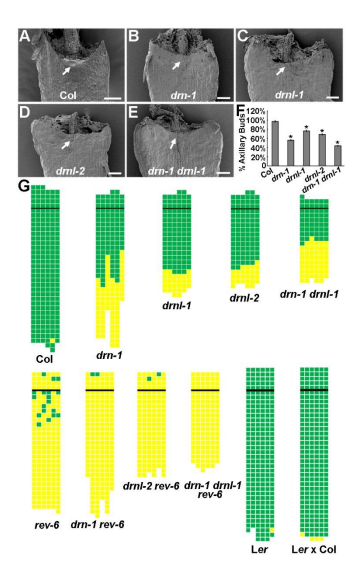
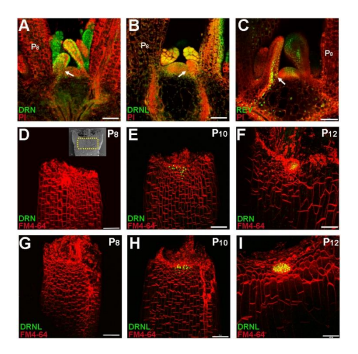


Fig. 1. DRN and DRNL are required for AM initiation.

(A) to (E) Scanning electron micrograph of P<sub>15</sub>-stage rosette leaf axils in Col-0 wild-type with a developing AM (dense cell mass, arrow) (A), and in *drn-1* (B), *drnl-1* (C), *drnl-2* (D), and *drn-1 drnl-1* (E) mutants with bare axils (arrows).

- (F) Graphic representations of axillary bud formation in *drn-1*, *drnl-1*, *drnl-2*, and *drn-1 drnl-1* mutants during vegetative development in comparison with Col-0 wild-type plants. The percentage values indicate the mean proportion of axillary buds formed over the total number of leaves along the shoot axis (plants analyzed >20). Error bars indicate the standard deviation.  $^*P < 0.01$  between wild-type and each mutant.
- (G) Schematic representation of axillary bud formation in leaf axils of Col-0 wild-type plants, drn-1, drnl-1, drnl-2, and drn-1 drnl-1, rev-6, drn-1 rev-6, drn-1 rev-6, and drn-1 drnl-1 rev-6 mutant plants, Ler and mixed Ler and Col-0 ecotypes. The thick black horizontal line represents the border between the youngest rosette leaf and the oldest cauline leaf. Each column represents a single plant, and each square within a column represents an individual leaf axil. The bottom row represents the oldest rosette leaf axils, with progressively younger leaves above. Green indicates the presence of an axillary bud, and yellow indicates the absence of an axillary bud in any particular leaf axil. Bars = 100  $\mu$ m.



**Fig. 2**. *DRN* and *DRNL* are expressed in leaf primordia and accumulate in the leaf axil prior to AM initiation.

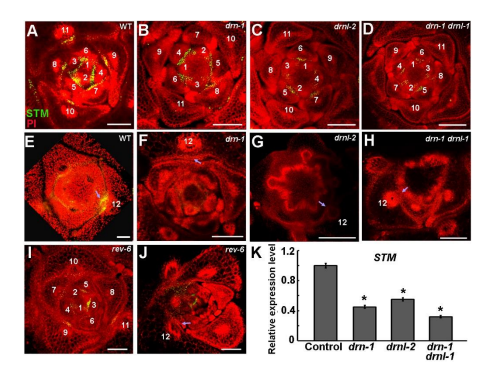
(A) to (C) Expression of *pDRN::DRN-GFP* (A), *pDRNL::DRNL-CFP* (B) and *pREV::REV-Venus* (C) in the vegetative shoot apex and leaf primordia.

Longitudinal sections of 14-d-old plant shoot apices were stained with propidium iodide (PI, red), and fluorescent signals are shown in green. Arrows indicate leaf axils. Note the presence of fluorescent signals in the leaf axils.

(D) to (I) Reconstructed view of the epidermal layer of P<sub>8</sub> stage (D and G), P<sub>10</sub>-stage (E and H), and P<sub>12</sub>-stage (F and I) leaf axils with *pDRN::DRN-GFP*(D) to (F), or *pDRNL::DRNL-CFP* (G) to (I) expression in green and FM4-64 staining in red showing the location of AM progenitor cells. The inset in (D) shows a scanning election micrograph of a rosette leaf axil at a similar stage, and the region within the yellow dotted box roughly corresponds to the imaged

regions shown in (D) to (I). All leaves were removed from 17-d-old plants. Note the enrichment of *DRN-GFP* and *DRNL-CFP* signals in P<sub>10</sub> and P<sub>12</sub> stage leaf axils.

Bars =  $50 \mu m$ .



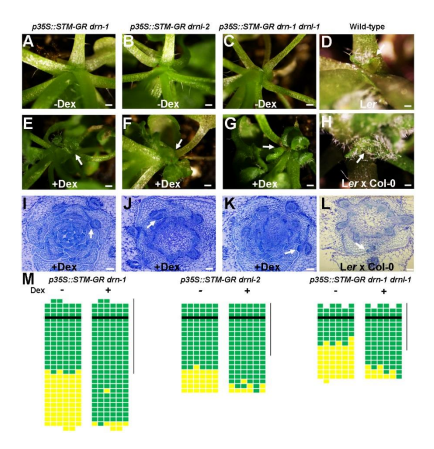
**Fig. 3**. Attenuated *STM* expression in *drn* and *drnl* mutants.

(A) to (D) Patterns of *pSTM::STM-Venus* (green) in transverse sections through the vegetative shoot apex of 28-d-old wild-type Col-0 (A), *drn-1* (B), *drnl-2* (C), *drn-1 drnl-1* (D). Plants stained with propidium iodide (PI, red). *STM* expression level decreased in the mutants compared to wild-type.

(E) to (H) Patterns of *pSTM::STM-Venus* expression in P<sub>12</sub> leaf axils in wild-type Col-0 (E), *drn-1* (F), *drnl-2* (G), *drn-1 drnl-1* (H). Note the lower *STM* expression level in mature leaves in the mutants than in wild-type.

(I) and (J) STM-Venus expression level in rev-6 in young (I) and mature (J) leaves. (K) RT-qPCR analysis indicates that STM expression is significantly reduced in drn-1, drnl-2, and drn-1 drnl-1 mutant plants. Vegetative shoots with the leaves removed were analysed. Error bars indicate SD. \*P < 0.01 (Student's t-test).

Bars =  $100 \mu m$ .



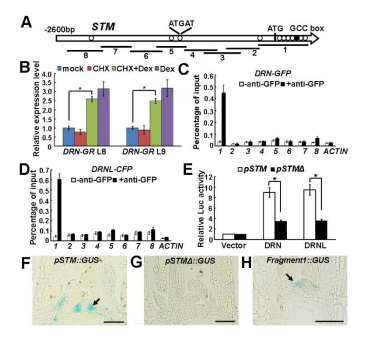
**Fig. 4**. Overexpression of *STM* rescues axillary bud deficiency in *drn* and *drnl* mutants.

- (A) to (C) Close-up of rosette leaf axils in mock-treated *p35S::STM-GR drn-1* (A), *p35S::STM-GR drnl-2* (B), and *p35S::STM-GR drn-1 drnl-1* (C) plants showing the absence of an axillary bud.
- (E) to (G) Close-up of rosette leaf axils in Dex-treated *p35S::STM-GR drn-1* (E), *p35S::STM-GR drnl-2* (F), and *p35S::STM-GR drn-1 drnl-1* (G) plants showing the presence of axillary buds (arrows).
- (D) and (H) Close-up of rosette leaf axils in mock-treated Ler and mixed Ler × Col-0 ecotypes showing the presence of axillary buds (arrows).
- (I) to (L) Transverse sections through vegetative shoot apices of 28-d-old Dex-treated p35S::STM-GR drn-1 (I), p35S::STM-GR drnl-2 (J),

p35S::STM- $GR\ drn$ -1 drnl-1 (K) and mixed Ler  $\times$  Col-0 ecotypes (L) stained with toluidine blue O, showing the presence of axillary buds (arrows) in rosette leaf axils.

(M) Schematic representation of axillary buds in leaf axils with or without Dex induction. Green indicates the presence of an axillary bud, and yellow indicates the absence of an axillary bud. Plants were grown under short-day conditions for 15 d without treatment; leaf axil regions were treated with 10 μM Dex every second day for another 15 d and then transferred to long-day conditions without treatment until axillary buds were counted. The vertical line indicates leaves initiated during Dex treatment.

Bars = 2 cm in (A) to (H), and 100  $\mu$ m in (I) to (L).



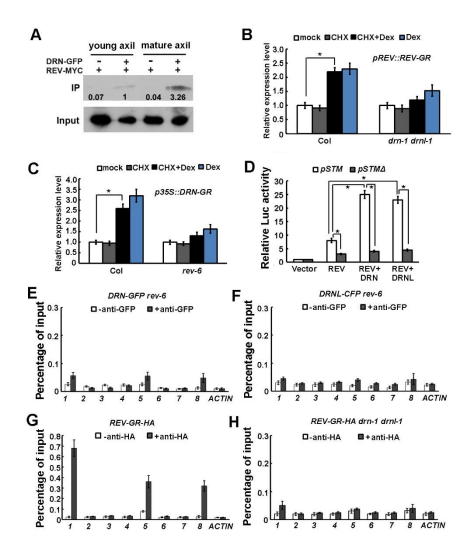
**Fig. 5**. DRN and DRNL regulate *STM* expression via binding to a conserved promoter motif.

- (A) Schematic representation of the *STM* genomic region. The black circle and white circles indicate the GCCGCC motif and the ATGAT motif, respectively; ATG denotes the translation start site. Eight PCR fragments were designed for ChIP analysis.
- (B) RT-qPCR analysis of *STM* expression using *p35S::DRN-GR* vegetative shoots (with the leaves removed) before and after simultaneous Dex and CHX treatment for 2 h. The vertical axis indicates the relative mRNA amount compared with the amount in the mock treatment. Error bars indicate the SD. Two independent transgenic lines were used. \*P < 0.01, (Student's t-test). (C) and (D) ChIP-qPCR analysis indicates binding of DRN-GFP (C), and DRNL-CFP (D) to fragment 1. Error bars indicate the SD.
- (E) Relative *Luc* reporter gene expression in transcriptional activity assays in *Arabidopsis* protoplasts. The 3.0 kb *STM* promoter region (*pSTM*) or the same

region without fragment 1 (as indicated in A,  $pSTM\Delta$ ) was co-transformed with p35S::DRN or p35S::DRNL, and p35S::GUS was the internal control. Data are means  $\pm$  SD for three independent biological experiments, each performed in triplicate. \*P < 0.01, (Student's t-test).

(F) to (H) Pattern of GUS expression driven by *pSTM* (F), *pSTM* (G) and fragment1 (H) in longitudinal sections through a vegetative shoot apex of 30-d-old plants. To compare signals, plants were stained in parallel for 6 h, and sections were placed on the same slides for detection. The GUS signal is barely detectable in leaf axils of *pSTM* (F) but weakly detectable in *fragment1::GUS* (H). Arrows highlight leaf axils. See Fig. S7 for more examples.

Bars =  $100 \mu m$  in (F) to (H).



**Fig. 6**. DRN and DRNL enhance the activation of *STM* expression by REV.

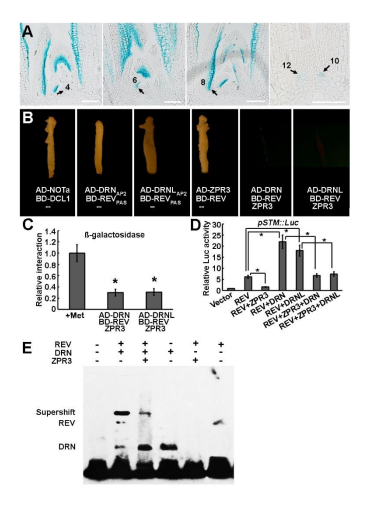
(A) Co-IP assay indicating that DRN and REV interact *in vivo* and the interaction is stronger in mature leaf (older than P<sub>10</sub>) axils than in young leaf (younger than P<sub>10</sub>) axils. The numbers below the blots indicate the relative ratios of the signal intensity between IP and input bands (IP/input). The ratios were normalized to the IP band in young leaf axils of crossed marker lines. An anti-GFP antibody was used for IP and an anti-MYC antibody as a probe. Input shows the amount of DRN-GFP protein used in the IP assay.

(B) RT-qPCR analysis of *STM* expression in *pREV::REV-GR* and

pREV::REV-GR drn-1 drnl-1 vegetative shoots (with the leaves removed)

before and after simultaneous Dex and CHX treatment for 2 h. The vertical axis indicates the relative mRNA amount compared to the amount in the mock treatment. Error bars indicate the SD. Note the reduction in STM activation in  $pREV::REV-GR\ drn-1\ drnl-1.*P < 0.01$ , (Student's t-test).

- (C) RT-qPCR analysis of *STM* expression in *p35S::DRN-GR* and *p35S::DRN-GR rev-6* vegetative shoots (with the leaves removed) before and after simultaneous Dex and CHX treatment for 2 h. The vertical axis indicates the relative mRNA amount compared to the amount in the mock treatment. Error bars indicate the SD. Note the reduction in *STM* activation in *p35S::DRN-GR rev-6*. \*P < 0.01, (Student's t-test).
- (D) Relative *Luc* reporter gene expression in transcriptional activity assays in *Arabidopsis* protoplasts. The *pSTM::Luc* or the *pSTM\Delta::Luc* constructs were co-transformed with *p35S::REV* alone, *p35S::REV* and *p35S::DRN*, or *p35S::REV* and *p35S::DRNL*, and *p35S::GUS* was the internal control. Data are means  $\pm$  SD. Error bars are derived from three independent biological experiments, each performed in triplicate. Note the enhanced activation of *STM* expression by DRN and DRNL. \**P* < 0.01, (Student's *t*-test).
- (E) and (F) ChIP-qPCR analysis demonstrates the reduced binding of DRN-GFP (E), and DRNL-CFP (F) to the *STM* genomic region (as in Figure 5A) in *rev-6* plants. Compare binding with that in Figure 5C and D for Col-0 wild-type plants. Error bars indicate the SD.
- (G) and (H) ChIP-qPCR analysis demonstrates binding of REV-GR-HA to the *STM* genomic region in Col-0 wild-type plants (G), and this is reduced in *drn-1 drnl-1* plants (H). Error bars indicate the SD.



**Fig. 7**. ZPR3 interferes with the DRN/DRNL–REV interaction and inhibits *STM* expression.

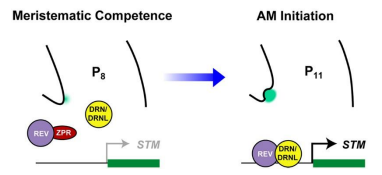
(A) Patterns of *ZPR3*-promoter-driven GUS expression in serial longitudinal 8-μm thick sections through the vegetative shoot apex of a 30-d-old wild-type-like plant. Arrows indicate the leaf axils. Note that the GUS signals are weaker in the leaf axils of P<sub>10</sub> and older leaves. See Fig. S8D–G for additional transverse sections.

Bars =  $100 \mu m$ .

(B) Y2H and Y3H assay showing the disruption of the DRN/DRNL–REV interaction by ZPR3. Yeast growth on SD-Leu-Trp-His-Ade plates showing that DRN, DRNL and ZPR3 interact with REV, respectively. The interaction of DRN

- or DRNL with REV was weakened after the induction of ZPR3 activity.

  AD-NOTa and BD-DCL1 were used as positive controls.
- (C) Relative  $\beta$ -galactosidase activity of the UAS-driven  $\beta$ -galactosidase reporter measured before and after ZPR3 induction in Y3H. Constructs and additional results are shown in Supplemental Figure S9. The data are mean values of three replicates  $\pm$  SD. \*P < 0.01, (Student's t-test).
- (D) Relative *Luc* reporter gene expression in transcriptional activity assays in *Arabidopsis* protoplasts. The *pSTM::Luc* construct was co-transformed with *p35S::REV* alone, *p35S::REV* + *p35S::ZPR3*, *p35S::REV* + *p35S::DRN*, *p35S::REV* + *p35S::DRNL*, *p35S::REV* + *p35S::DRNL*, or *p35S::REV* + *p35S::ZPR3* + *p35S::DRNL*, and *p35S::GUS* was the internal control. Data are means  $\pm$  SD. Error bars are derived from three independent biological experiments, each performed in triplicate. Note the suppression of REV activation and DRN/DRNL–REV co-activation of *STM* expression by ZPR3. \*P < 0.01, (Student's t-test).
- (E) Supershift in EMSA, indicating that REV and DRN interact and bind to a biotin-labeled *STM* promoter fragment. The addition of ZPR3 decreased the intensity of the supershift band of DRN and REV; 2 μg DRN and ZPR3, and 1 μg REV protein was used for incubation.

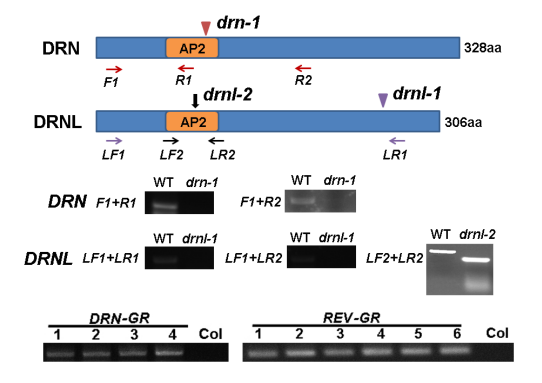


**Fig. 8**. A model summarizing the up-regulation of *STM* expression prior to AM initiation through the licensing of active DRN/DRNL–REV complexes by ZPRs.

## Supplemental Table 1. Primers used in this study.

Primer	Sequence(5'-3')
RT-PCR STM-F	GGCCGCTTATGTCAATTG (95%)
RT-PCR STM-R	AGTAAGCTTCCATGAATTG
ChIP STM-1F	AGGGTTTCCCAATAAATTTA (91%)
ChIP STM-1R	CCATGATGTTGATGTGATG
ChIP STM-2F	GAAGAACACAAAAGAGTTA (93%)
ChIP STM-2R	AAAAGAGTAAGTAACTGTGT
ChIP STM-3F	TACTTTTAAACCAACATC (93%)
ChIP STM-3R	TGCTGAAATATGATTTCTC
ChIP STM-4F	TATTTATGACATTTTTCTTC (99%)
ChIP STM-4R	CATTTGTGTATATGTATATA
ChIP STM-5F	AGGTACAGGTGTCCCATGCT (98%)
ChIP STM-5R	TAATCATAGAGTAAAACATGT
ChIP STM-6F	TAGTGATAACAAAATTAGC (92%)
ChIP STM-6R	GTTTATCCCTGAAAATAGT
ChIP STM-7F	AGTTTCTCAGTTATCTTTTC (96%)
ChIP STM-7R	TTCGAAATATTATTTAGAAG
ChIP STM-8F	AAATATTTATATCTAATAT (96%)
ChIP STM-8R	AATAGATAGAGAGAGATTA
ChIP ACTIN-F	GTCGTACAACCGGTATTGTGC (99%)
ChIP ACTIN-R	CACAAACGAGGGCTGGAACAAG
pSTM-F	<i>AACTGCAG</i> AGACGTAGTAATACTAGA
pSTM-R	<i>GCTCTAGA</i> CTACTTTGTTGGTGGTGTG
pSTM⊿-R	<i>GCTCTAGA</i> GAAACACATTAAAGTACTA
RT-PCR ZPR1-F	GTTCTTCAGAAACATTTTC (98)
RT-PCR ZPR1-R	CTCCACGTAAAGCTTTAG
RT-PCR ZPR3-F	ACTGTTACATAATGAAAG (96%)
RT-PCR ZPR3-R	AGATTGTCCAGAAGCAGA
ChIP ZPR1-F	CTTTAAACGAATAATGCAA (93%)
ChIP ZPR1-R	AGCTCCAAGATGTGTTTAA
DRN F1	GAAACTTCACCGAATCTACCC
DRN R1	GCCGCAGAGTCGTAAGCACAA
DRN R2	GAGCCGTTGAAAGAGTTATT
DRNL F1	CGAAGGTGCCGAGCACAGA
DRNL F2	TACCGCAAAAGCTGCCTC
DRNL R1	TCCACCATTTCCGTTCTGC
DRNL R2	AGCGGCGCAGTCATATGCGCAGTCT
GR F1	GAGGTGATTGAACCCGAGG
GR F2	ACCGTTGCCAGTTCTGAC

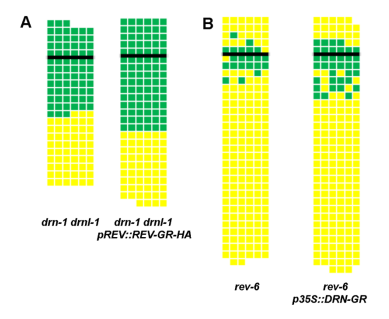
The values in brackets indicate the efficiency of qPCR primers.



**Fig. S1.** Schematic diagrams of DRN and DRNL proteins showing the positions of mutations in the corresponding encoding genes.

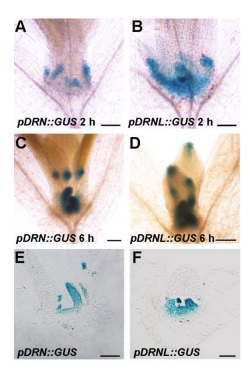
The red arrow at position 107 indicates the insertion site of the *dSpm* element into the AP2 domain-encoding sequence in the *drn-1* allele (Kirch et al., 2003). F1+R1 and F1+R2 could not amply bands from cDNA of *drn-1* mutant, indicating that *drn-1* is a null allele. In *drnl-1*, transposon-flanking DNA was inserted at position 259 near C terminal. LF1+LR1 and LF1+LR2 could not amplify bands from cDNA of *drnl-1*. In *drnl-2* arrow indicates the single nucleotide substitution that results in an A to V substitution at position 93 in the AP2 domain. Homozygosity was confirmed using a dCAPS marker with primers LF2 and LR2, which give a wildtype amplicon of 184 bp that is cleaved in the mutant by *Acc*I into two fragments of 159 and 25 bp. Different

p35S::DRN-GR and pREV::REV-GR-HA transgenic lines (labeled with a number) were genotyped using genomic DNA and primers for GR. Line1 for each transgenic line was used in the experiments, except where otherwise indicated.



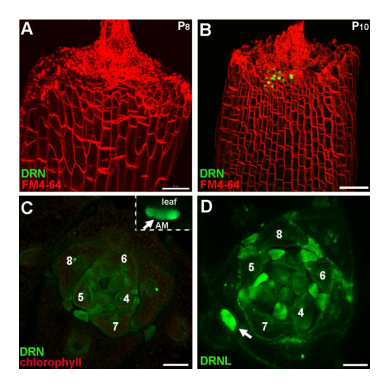
**Fig. S2.** Frequency of axillary buds in *pREV::REV-GR-HA drn-1 drnl-1* and *p35S::DRN-GR rev-6*.

The number of axillary buds was not affected in early rosette leaves but slightly increased in the axil of later rosette leaves and cauline leaves following Dex treatment in *drn-1 drnl-1 pREV::REV-GR-HA* plants (A) and *rev-6 p35S::DRN-GR* plants (B) compared with the controls. The thick black horizontal line represents the border between the youngest rosette leaf and the oldest cauline leaf. Each column represents a single plant, and each square within a column represents an individual leaf axil. The bottom row represents the oldest rosette leaf axils, with progressively younger leaves above. Green indicates the presence of an axillary bud, and yellow indicates the absence of an axillary bud in any particular leaf axil.



**Fig. S3.** *DRN* and *DRNL* expression patterns in shoot apices and young leaf primordia.

- (A) and (C) Expression pattern of *pDRN::GUS* in the vegetative shoot apex and leaf primordia of 10-d-old plants. Seedlings were stained for 2 h in (A) and 6 h in (C).
- (B) and (D) Expression pattern of *pDRNL::GUS* in the vegetative shoot apex and leaf primordia of 10-d-old plants. Seedlings were stained for 2 h in (B) and 6 h in (D).
- (E) and (F) Longitudinal sections for pDRN::GUS (E) and pDRNL::GUS (F), showing expression in the shoot apex. Material was stained for 2 h. Bars = 100  $\mu$ m in (A) to (D), and 50  $\mu$ m in (E) and (F).

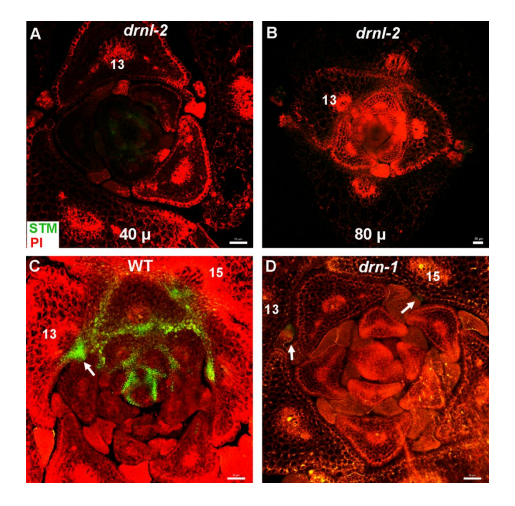


**Fig. S4.** *DRN* is expressed prior to AM initiation in leaf axils.

(A) and (B) Reconstructed view of the epidermal layer of a P<sub>8</sub>-stage (A) and a P<sub>10</sub>-stage (B) leaf axil with *pDRN::DRN-GFP* expression in green and FM4-64 stain in red showing the location of AM progenitor cells. Both leaves were excised from 17-d-old plants. Note the enrichment of *DRN-GFP* signals in the P<sub>10</sub>-stage leaf axil.

(C) and (D) Transverse sections of pDRN::DRN-GFP and pDRNL::DRNL-CFP plants to show the decreased level of DRN and DRNL expression in mature leaves ( $>P_8$ ) and strong expression in the axillary meristem (arrows). The seedling was 21-days-old and the leaf showing an axillary meristem highlighted in the inset in (C) was  $P_{15}$ .

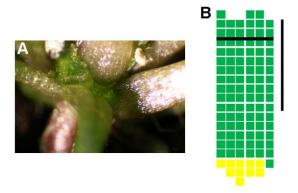
Bars =  $50 \mu m$ .



**Fig. S5.** The expression pattern of *pSTM::STM-Venus* in *drnl-2* and *drn-1* mutants.

- (A) and (B) *STM* was weakly expressed in young and mature leaves. Images were obtained from continuous sections from one *drnl-2* plant.
- (C) and (D) STM-Venus signals were strong in mature leaves of wild-type (arrow represents  $P_{13}$ ), but were severely attenuated in drn-1 mutants (arrows represent  $P_{13}$  and  $P_{15}$ ). Sections for wild-type and drn-1 were at an equivalent position.

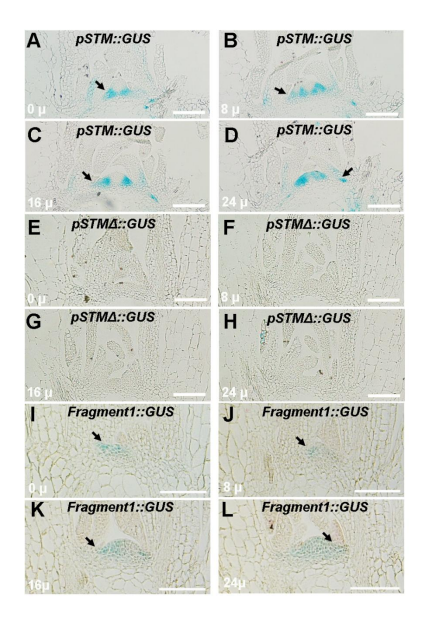
Bars =  $50 \mu m$ .



**Fig. S6.** *STM* overexpression complemented the reduction in axillary meristems in *drn-1 drnl-1 rev-6* mutants.

- (A) Dex induced axillary meristems in p35S::STM-GR drn-1 drnl-1 rev-6 plants.
- (B) The number of axillary buds increased in *p35S::STM-GR drn-1 drnl-1 rev-6* plants after Dex treatment. Plants were grown in short-days for 15 d without treatment; leaf axil regions were treated with 10 μM Dex every second day for another 15 d and then shifted to long-day conditions without treatment until axillary buds were counted. The vertical line indicates leaves initiated during Dex treatment.

The thick black horizontal line represents the border between the youngest rosette leaf and the oldest cauline leaf. Each column represents a single plant, and each square within a column represents an individual leaf axil. The bottom row represents the oldest rosette leaf axils, with progressively younger leaves above. Green indicates the presence of an axillary bud, and yellow indicates the absence of an axillary bud in any particular leaf axil.



**Fig. S7.** The expression patterns of *pSTM::GUS*, *pSTM∆::GUS* and *fragment1::GUS*.

Serial longitudinal 8- $\mu$ m thick sections through vegetative shoot apices of 30-d-old plants showing the pattern of GUS expression driven by pSTM (A–D),  $pSTM\Delta$  (E–H) or fragment1::GUS (I–L). To compare the signals, the plants were stained under the same conditions. Note the reduction in GUS expression in the leaf axils of  $pSTM\Delta::GUS$  plants and the weak GUS expression in fragment1::GUS plants. Arrows highlight leaf axils. Bars = 100  $\mu$ m.

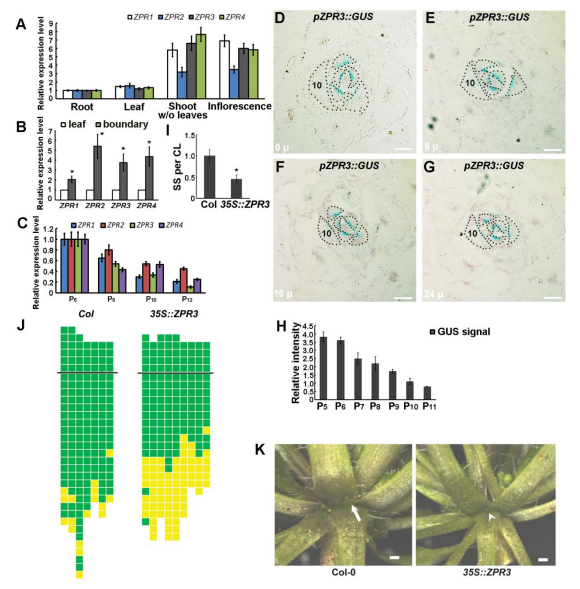
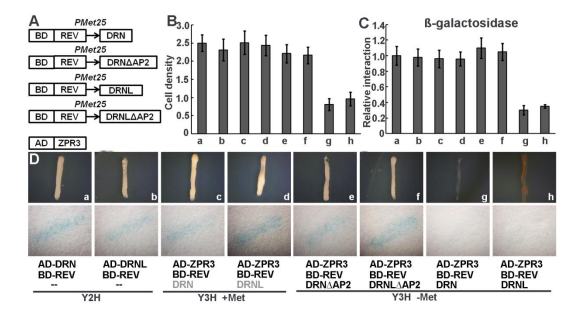


Fig. S8. The expression pattern of ZPR in leaf axils.

- (A) RT-qPCR analysis of *ZPR1*, *ZPR2*, *ZPR3* and *ZPR4* expression levels in roots, leaves, shoots without leaves, and inflorescences. The latter two tissues are enriched with boundary tissues following leaf removal. Error bars indicate the SD.
- (B) RT-qPCR analysis of ZPR expression in leaves and shoot apices without

leaves, which are enriched with boundary tissues. Error bars indicate the SD.  $^*P < 0.01$ , (Student's t-test).

- (C) RT-qPCR analysis of ZPR gene expression at different leaf stages.
- (D) to (G) Pattern of *ZPR3*-promoter-driven GUS expression in serial transverse sections through the vegetative shoot apex of a 30-d-old wild-type-like plant. The 8- $\mu$ m thick sections are ordered from the most apical (D) to the most basal (G); the approximate distance from the summit of the meristem to the midpoint of the section is shown in the bottom left-hand corner of each image. Note that the GUS signals decrease in intensity in the axils of older leaves. Bars = 100  $\mu$ m.
- (H) Measurement of GUS intensity in different leaves of 10, *pZPR3::GUS* plants.
- (I) Frequency of secondary shoots per cauline leaf in wild-type and p35S::ZPR3 transgenic lines. \*P < 0.01, (Student's t-test).
- (J) Schematic of the frequency of axillary buds in *p35S::ZPR3* plants. Green indicates the presence and yellow indicates the absence of an axillary bud in any particular leaf axil.
- (K) Close up of views of an exemplary p35S::ZPR3 transgenic plant showing an axillary meristem defect compared to wild type. Bars = 100  $\mu$ m.



**Fig. S9.** ZPR3 interferes with the DRN/DRNL–REV interaction.

- (A) A schematic representation of the constructs for Y3H vectors. DRN/DRNL and DRN/DRNLΔAP2 were inserted into the BD vector and driven by the *Met25* promoter, which can be suppressed by Met. In a complementary assay, DRN was replaced by ZPR3 in the BD vector.
- (B) to (D) Yeast growth on SD-Leu-Trp-His-Ade plates and staining for  $\beta$ -galactosidase activity. Cell density quantification after incubation for 16 h (B), relative  $\beta$ -galactosidase activity of the transformed yeast (C), and staining for  $\beta$ -galactosidase activity (D) were performed. The addition of Met inhibited the activity of the *Met25* promoter.