The SvFUL2 transcription factor is required for inflorescence determinacy and timely flowering in Setaria viridis

Jiani Yang,1 Edoardo Bertolini,1 Max Braud,1 Jesus Preciado,2,‡ Adriana Chepote,1 Hui Jiang,1 and Andrea L. Eveland1,*

1 Donald Danforth Plant Science Center, Saint Louis, Missouri, 63132, USA
2 National Science Foundation Research Experiences in Plant Science at the Danforth Center, Saint Louis, Missouri, 63132, USA

*Author for communication: aeveland@danforthcenter.org
†Senior author.
‡Present address: Plant Molecular and Cellular Biology Program, University of Florida, Gainesville, Florida, 32611, USA.

J.Y. and A.L.E. designed and advised the research. J.Y., M.B., J.P., A.C., and H.J. performed genetics and morphological experiments. E.B. performed GRN analyses. M.B. performed gene editing. J.Y. and E.B. performed data analyses. J.Y., E.B., and A.L.E. wrote the paper. All authors have read and approved the manuscript for submission.

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 Author (https://academic.oup.com/plphys/pages/general-instructions) is: Andrea L. Eveland (aeveland@danforthcenter.org).

Abstract

Inflorescence architecture in cereal crops directly impacts yield potential through regulation of seed number and harvesting ability. Extensive architectural diversity found in inflorescences of grass species is due to spatial and temporal activity and determinacy of meristems, which control the number and arrangement of branches and flowers, and underlie plasticity. Timing of the floral transition is also intimately associated with inflorescence development and architecture, yet little is known about the intersecting pathways and how they are rewired during development. Here, we show that a single mutation in a gene encoding an AP1/FUL-like MADS-box transcription factor significantly delays flowering time and disrupts multiple levels of meristem determinacy in panicles of the C4 model panicoid grass, Setaria viridis. Previous reports of AP1/FUL-like genes in cereals have revealed extensive functional redundancy, and in panicoid grasses, no associated inflorescence phenotypes have been described. In S. viridis, perturbation of SvFUL2, both through chemical mutagenesis and gene editing, converted a normally determinate inflorescence habit to an indeterminate one, and also repressed determinacy in axillary branch and floral meristems. Our analysis of gene networks connected to disruption of SvFUL2 identified regulatory hubs at the intersection of floral transition and inflorescence determinacy, providing insights into the optimization of cereal crop architecture.

Introduction

Inflorescence structure determines fruit, seed, and pollen production, which are critical for reproductive success of plants and global food security. During the shift from vegetative to reproductive growth, the indeterminate shoot apical meristem (SAM), which patterns the vegetative organs, transitions to an inflorescence meristem (IM). Like the SAM, the IM continues indeterminate growth but instead, leaf growth is suppressed and axillary meristems (AMs) grow out into reproductive organs on its flanks. In
The regulatory modules that control inflorescence growth habit are somewhat conserved between eudicots and grasses. In maize and rice, TFL1-like genes delay flowering time and prolong the indeterminate status of the developing inflorescence (Nakagawa et al., 2002; Danilevskaya et al., 2010; Kaneko-Suzuki et al., 2018). In rice, AP1/FUL-like genes have overlapping roles in flowering time (Kobayashi et al., 2012). Over-expression of OsMADS14, OsMADS15, or OsMADS18 all result in early flowering phenotypes (Jeon et al., 2000; Fornara et al., 2004; Lu et al., 2012), and in the case of OsMADS15, reduced panicle size and branch number (Lu et al., 2012). In winter wheat and barley varieties, expression of VERNALIZATION 1 (VRN1), an AP1/FUL-like gene, has been well-characterized as an early signal in promoting timely vegetative-to-reproductive transition in response to vernalization (Yan et al., 2003; Preston and Kellogg, 2008; Li et al., 2019). Expression of FUL2 and FUL3 genes in wheat are also induced by vernalization to promote flowering (Chen and Dubcovsky, 2012; Li et al., 2019). A recent study revealed that AP1/FUL-like genes in wheat and the genetic interactions among them contribute to maintenance of IM and SM determinacy, as well as flowering time (Li et al., 2019). Loss-of-function in both VRN1 and FUL2 genes converted the normally determinate IM of the wheat spike to an indeterminate habit, and also enhanced indeterminacy in primary AMs. Introduction of a single functional copy of either VRN1 or FUL2 reverted the vrn-null; ful1-null mutant IM back to a determinate habit (Li et al., 2019).

While evidence across the plant kingdom supports conserved roles for AP1/FUL-like genes in floral transition and inflorescence architecture, to date, there have been no inflorescence phenotypes described for loss-of-function AP1/FUL-like genes in the subfamily Panicoideae, which includes agronomically important crops such as maize, sorghum, and sugarcane (Sacccharum officinarum). This is likely due to functional redundancy (Litt and Irish, 2003; Preston and Kellogg, 2007). In this study, we show that a single loss-of-function mutation in an AP1/FUL-like gene in model panicoide grass, Setaria viridis (green foxtail), is sufficient to confer both strong flowering time and inflorescence determinacy phenotypes despite its overlapping expression pattern with three closely related paralogs. S. viridis is a weedy, C₄ species that has demonstrated promise as a model system for elucidating molecular mechanisms in panicoide crops (Li and Brutnell, 2011; Huang et al., 2017; Yang et al., 2018). It also represents a key evolutionary node between domesticated and undomesticated grasses. Like wheat, S. viridis produces a determinate inflorescence that terminates in a spikelet, but AMs undergo multiple orders of branching (Doust and Kellogg, 2002; Zhu et al., 2018). We isolated the SvFUL2 mutant in a genetic screen, which displayed a “barrel”-like panicle morphology due to enhanced indeterminacy in AMs. The determinate IM was also converted to an indeterminate habit resembling a maize ear. Further investigation of SvFUL2 loss-of-function at the molecular level using genomics approaches revealed regulatory modules that link floral transition and inflorescence determinacy pathways through interactions among MADS-box transcription factors (TFs)
and several other developmental regulators. This mutant and the analyses presented here, provide insights into the complex interface of flowering time and inflorescence development, and potential targets for fine-tuning inflorescence ideotypes in cereal crops.

Results
Characterization of the *barrel 1* mutant in *S. viridis*

In a forward genetics screen of approximately 3,000 N-methylurea (NMU) mutagenized M2 families of *S. viridis* (Huang et al., 2017; Yang et al., 2018), we isolated the *barrel 1* (*brl1*) mutant, named for its abnormal, barrel-shaped panicle (*brl1-ref* allele). Compared with mature panicles of the wild-type mutagenized reference line (A10.1), mutant panicles were shorter and thicker and appeared more branchy (Figure 1A and B; Table 1). Mutant plants were shorter in stature and produced tillers with more leaves (Figure 1A; Table 1). In addition to morphological defects, flowering time was obviously delayed in *brl1* mutants. To test the effect of different photoperiods (Ppds) on floral transition, we examined flowering time of the mutant compared to control plants under both short day (SD; 12-h light/12-h dark) and long day (LD; 16-h light/8-h dark) conditions. Under SD conditions, which typically promote flowering in *S. viridis* (Doust et al., 2017), *brl1* mutant panicles emerged ~6 d later (average 29.20 d after sowing [DAS]) than those of wild-type (average 22.94 DAS; Figure 1C). Under LD conditions, flowering time in *brl1* mutant (avg. 31.21 DAS) and A10.1 wild-type plants (average at 28.44 DAS) was delayed compared to under SDs, but *brl1* mutants still flowered significantly later than wild-type (average 3 d; Figure 1D; Table 1).

Previous studies in *S. viridis* showed that flowering time impacted both plant architecture and biomass (Doust, 2017). Under both LD and SD conditions, plant height and panicle length of *brl1* mutants were significantly shorter than wild-type plants at maturity (Table 1). Under SD conditions, above-ground dry weight was increased in mutants compared to wild-type, largely due to biomass of vegetative tissue (leaves and stems; Table 1). In LDs, above-ground dry weight of *brl1* mutants was comparable to wild-type,
however, we still observed a significant increase in dry weight of vegetative tissue (leaves and stems) in the mutant (Table 1). Seed shape and size were also different with the mutant seeds being longer and narrower than those of wild-type (Figure 1E; Table 1).

Examination of the inflorescence morphology revealed that brl1 mutants displayed various levels of indeterminacy. At the tip of the panicle, the IM appeared indeterminate in mutants, and newly formed branch meristems (BMs) were still visible at maturity (Figure 1F). At the base of the mutant panicle, rudimentary primary branches were observed, which were not found in wild-type (Figure 1G). Primary branches were longer in brl1 mutants and the panicle rachis was clearly thicker (Figure 1H; Supplemental Figure S1A). Bristles, which are modified branches paired with spikelets in Setaria sp, did not elongate to the length of wild-type bristles, and so were largely found buried under spikelets (Figure 1H). In Setaria spp., spikelets form upper and lower FMs. During differentiation, the upper floret (uf) develops whorls of floral organs, that is, lemma and palea, lodicules, stamens, and pistil, within a pair of subtending glumes, and the lower floret is aborted. Development of spikelets and flowers was also affected in brl1 mutants, but phenotypes showed low penetrance with various severities of indeterminacy. For example, ~17% of brl1 mutant panicles produced additional flowers, bristles, and/or spikelets within spikelets compared to the typical one flower per spikelet in wild-type (Figure 1H–K). The lemma and palea of mutant flowers were more elongated in the mutant and were more rigid, which is likely contributing to the elongated seed shape (Supplemental Figure S1, B and C).

brl1 mutants show loss of determinacy in various stages of inflorescence development

We used scanning electron microscopy (SEM, Figure 2) to compare the developmental progression of inflorescence primordia from the brl1 mutant with that of wild-type S. viridis. By 11 DAS, the vegetative SAM of wild-type plants had finished transitioning to the reproductive IM, as the first primary BMs were initiated on its flanks (Figure 2A). In the brl1 mutant, the vegetative-to-reproductive transition was delayed to 15 DAS (Figure 2B), consistent with its late-flowering phenotype (Figure 1C). After the transition, wild-type inflorescences initiated primary branches in a spiral pattern (Figure 2C), and then secondary and tertiary axillary branches sequentially in a distichous pattern, as previously described (Figure 2E; Doust and Kellogg, 2002; Yang et al., 2018). The brl1 IM was elongated compared to that of wild-type (Figure 2B and Table 1; Supplemental Figure S1, D and E), and this appeared to enable capacity for increased initiation of primary and higher-order branches (Figure 2, D and F), consistent with the mature panicle phenotype. By 17 DAS, the wild-type IM had become determinate and terminated as a spikelet (Figure 2G). BMs then began to differentiate from the tip of the inflorescence primordium into either an SM or a sterile bristle, and this continued basipetally (Figure 2, G and I). Conversely, the IM of the brl1 mutant remained indeterminate and continued to produce primary BMs at 21 DAS (Figure 2H), where SMs and bristles began to differentiate toward the top of the inflorescence primordium (Figure 2). By the end of the developmental series analyzed by SEM, the brl1 IM remained indeterminate, which is consistent with its mature phenotype in Figure 1F.

While differentiation of SMs and bristles appeared normal in the mutant (Figure 2, I and J), the onset was delayed compared to wild-type and after additional rounds of higher-order branching (Figure 2F). SMs developed similarly in brl1 mutants and wild-type, initiating glumes and upper and lower FMs; the uf typically develops into a perfect flower with lemma, palea, anther, and carpel and the lower

| Table 1 | Phenotypic measurements of brl1 mutant plants compared to wild-type (A10.1) controls |
|-----------------|-------------------------------|--------------|--------------|
| Photoperiod     | Trait                          | A10.1        | brl1         |
| LD              | Days to panicle emergence (DAS)| 28.44 ± 1.06 | 31.21 ± 1.82** |
| 16-h Light/8-h Dark | Weight of leaves and stems (g)  | 3.83 ± 0.98  | 4.73 ± 0.65** |
|                 | Weight of panicles with seeds (g) | 3.06 ± 1.14  | 2.01 ± 0.62** |
|                 | Plant height (cm)              | 61.72 ± 11.23 | 35.81 ± 3.75** |
|                 | Panicle length (cm)            | 6.25 ± 0.61  | 5.03 ± 0.71** |
| SD              | Days to panicle emergence (DAS)| 22.94 ± 0.97 | 29.20 ± 3.00** |
| 12-h Light/12-h Dark | Weight of leaves and stems (g)  | 0.89 ± 0.25  | 2.00 ± 0.38** |
|                 | Weight of panicles with seeds (g) | 1.45 ± 0.51  | 1.18 ± 0.53** |
|                 | Plant height (cm)              | 30.95 ± 4.75 | 16.74 ± 2.00** |
|                 | Tiller No.                     | 11.20 ± 3.93 | 16.60 ± 3.68** |
|                 | Leaf No.                       | 38.00 ± 12.94 | 126.25 ± 29.97** |
|                 | Panicle length (cm)            | 3.53 ± 0.41  | 3.16 ± 0.50** |
|                 | Primary branch No.             | 36.55 ± 3.80 | 47.95 ± 9.56** |
|                 | Weight of 100 seeds (mg)       | 159.10       | 149.40       |
|                 | Seed length (mm)               | 2.08 ± 0.06  | 2.39 ± 0.15** |
|                 | Seed width (mm)                | 1.28 ± 0.05  | 1.21 ± 0.07** |
|                 | IM* length (μm)                | 202.92 ± 23.55 | 310.80 ± 34.51** |
|                 | IM* width (μm)                 | 65.50 ± 7.60 | 89.99 ± 10.45** |

All data except for IM length and width (n ≥ 9) are based on mean values of n ≥ 20 individuals (±sd).

*Length and width of IM were measured at the end of the reproductive transition and before layout primary branches (Supplemental Figure S1, D and E).

**Indicates a significant difference in brl1 mutant compared to A10.1 (wild-type) determined by Student’s t test (P < 0.01).
floret aborts (Figure 2K; Doust and Kellogg, 2002; Yang et al., 2018). In some cases, we observed aberrant meristematic outgrowths in brl1 FMs (Figure 2L) which may explain our observations of additional spikelets and bristles within some spikelets (Figure 1, I and J). Our SEM analysis showed that a determinacy program was delayed in the IM, BMs, and SMs of brl1 mutants.

The brl1 locus encodes SvFUL2, a MIKC-type MADS-box TF

F2 populations were generated from a cross between the brl1 mutant and the parental line, A10.1. Wild-type and barrel-like panicle phenotypes segregated with the expected Mendelian 3:1 ratio (139:48; $P [\chi^2, 1 \text{ d.f.}] = 0.83$), which indicated that brl1 is a single locus recessive allele. To map the brl1 locus, Bulk Segregant Analysis (BSA) was performed (Michelmore et al., 1991; Schneeberger, 2014) with a pool of DNA from 30 brl1 mutant individuals from the segregating F2 population that was sequenced to approximately 92x coverage (244.2M reads). Reads were aligned to the A10.1 reference genome (phytozome.jgi.doe.gov; version 2.1; Mamidi et al., 2020) and single nucleotide polymorphisms (SNPs) were called using GATK. Three high-confidence, non-synonymous SNPs were identified and supported by high observed allele frequency (Supplemental Figure S2; Supplemental Data Set S1). One candidate SNP disrupted the start codon of Sevir.2G006400, a MIKC-type MADS-box gene, and was supported by whole-genome sequencing of the brl1 mutant (Figure 3, A and B). Sevir.2G006400 had previously been annotated as SvFUL2 based on phylogenetics and evolutionary developmental analyses (Preston et al., 2009; Zhu et al., 2018). The other two non-synonymous changes which generated missense mutations in Sevir.2G174301 and Sevir.7G048300 on chromosomes 2 and 7, respectively, were not identified by whole-genome sequencing of the brl1 mutant, indicating that they might be randomly fixed mutations by selfing (Supplemental Data Set S1). Unlike our previous experience with mapping by BSA in this population (Yang et al., 2018), we did not resolve a clear peak in the genomic region surrounding SvFUL2, likely due to its position at the end of chromosome 2 where recombination rates are relatively higher and break linkage.

We designed a dCAPs marker specific for this SNP and genotyped over 200 segregating F2 individuals. Our genotyping results showed that this SNP co-segregated with the barrel panicle phenotype at 100% (Supplemental Figure S3). Transcript levels of SvFUL2 were strongly reduced in the mutant inflorescence primordia compared to wild-type early in development based on RT-PCR (Supplemental Figure S4).

SvFUL2 encodes the ortholog of OsMADS15 in rice, an AP1/FUL-like MADS-box gene in the MIKC-type...
subfamily. Consistent with previous phylogenetic studies (Wu et al., 2017; Li et al., 2019), our phylogenetic analysis of AP1/FUL-like MADS-box genes from *S. viridis* as well as Arabidopsis, rice, wheat, maize, and sorghum showed that *SvFul2* was located in the FUL2 subclade along with three copies of wheat *Ful2*, rice *OsMADS15*, and maize *zap1* and *mads3* (Figure 3A; Supplemental Data Set S2). *SvFul2* is more closely related to *SvFul1* (Sevir.9G087300) in the FUL1 subclade, which includes wheat VRN1s, rice *OsMADS15*, maize *zmm15*, and *zmmads4*. *SvFul3* (Sevir.2G393300) and *SvFul4* (Sevir.3G374401) are located in the FUL3 and FUL4 subclades, respectively. By examining a previously generated transcriptomics resource across six sequential stages of early *S. viridis* inflorescence development (Zhu et al., 2018), we found that *SvFul1*, *SvFul2*, and *SvFul3* shared similar spatiotemporal expression patterns, increasing during branching and then decreasing during floral development with a small drop during spikelet specification (Figure 3C). *SvFul1* was expressed highest at 10 and 12 DAS and *SvFul2* expressed more at later stages, which indicate the two may have different functions. Comparatively, *SvFul4* was expressed at lower levels throughout inflorescence development, its expression gradually decreasing after the reproductive transition.

**Figure 3** Phylogeny of AP1/FUL-like MADS-box genes in grasses and Arabidopsis and their expression profiles during *S. viridis* inflorescence development. A, Phylogenetic analysis of AP1/FUL-like MADS-box genes from *S. viridis*, Arabidopsis, and several grass species based on protein coding sequence. *SvFul2* is highlighted in red. *Os* = *Oryza sativa*, *Ta* = *Triticum aestivum*, *Zm* = *Zea mays*, *Sb* = *Sorghum bicolor*, *Sv* = *Setaria viridis*. B, Exon–intron structure of the *SvFul2* gene consists of seven exons (solid rectangles) and six introns (horizontal line). The 5′- and 3′-untranslated regions are shown as gray rectangles. Gray triangle indicates the location of the SNP that disrupts the start codon within the *SvFul2* gene. C, Expression profiles (RNA-seq) of four *S. viridis* AP1/FUL-like MADS-box genes across six stages of early inflorescence development based on the transcriptomics resource described in (Zhu et al., 2018). Error bars indicate standard errors of three to four biological replicates.

**Gene editing of SvFul2 validates the mutant phenotype in S. viridis**

To validate that Sevir.2G006400 (*SvFul2*) is responsible for the observed phenotypes of the br1 mutant, we used genome editing. A clustered regularly interspaced short palindromic repeats (CRISPR)/Cas9 construct was designed containing two guide (g)RNAs that specifically targeted the first exon and the first intron of *SvFul2*, respectively, in the highly transformable *S. viridis* accession, ME034 (Figure 4A; Acharya et al., 2017; Van Eck, 2018). In the T1 generation, individual plants carrying a homozygous 540-bp deletion in the first exon of *SvFul2* were selected (Figure 4A; Supplemental Figure S5). We called this genotype *SvFul2_KO*. These were moved forward to generation T2 where they were then outcrossed to ME034 and then selfed to select Cas9-free *SvFul2_KO* plants for phenotyping. *SvFul2_KO* plants displayed phenotypes consistent with
observed in brl1 mutants, panicles of SvFul2_KOs took on an indeterminate growth habit (Figure 4D). Flowering time was also delayed in the SvFul2_KOs (avg. 20.76 DAS) compared to the ME034 wild-type siblings (avg. 17.87 DAS) by ~3 d (Figure 4G; Supplemental Table S1). The ME034 accession flowers earlier than A10.1, consistent with the shift in flowering time shown here. SEM analysis of SvFul2_KO during early inflorescence development showed floral transition consistent with the ME034 panicle emergence data, and increased branching capacity and IM indeterminacy as observed in brl1-ref (Supplemental Figure S6).

We further tested the allelic relationship between brl1-ref and SvFul2_KO by a genetic cross. brl1-ref and SvFul2_KO fail to complement and are allelic (Figure 5). Taken together, our analyses support SvFul2, Sevir. 2G006400, as the locus responsible for the brl1 mutant phenotypes in S. viridis.

Loss of SvFul2 function alters expression of flowering and meristem determinacy pathways

To determine the molecular mechanisms underlying the complex phenotypes of the Svful2 mutant, we used RNA-seq to profile gene expression in mutant inflorescence primordia across three key developmental transitions and compared them to equivalent stages in wild-type controls: right before (Stage 1) and after (Stage 2) the floral transition, and during the initiation of spikelet specification (Stage 3; Supplemental Figure S7; Supplemental Data Set S3). Here, we expect to capture transcriptional changes related to both differences in flowering time and meristem determinacy. For each stage, we profiled four biological replicates, each consisting of pooled, hand-dissected inflorescence primordia. Differential expression was determined using DESeq2 (1.22.2). Our analysis found 382, 2,584, and 2,035 differentially expressed genes (DEGs) at Stages 1–3, respectively (Supplemental Data Set S4).

Based on Principal Component Analysis (PCA), we observed fewer differences in the mutant transcriptome at Stage 1, suggesting that the main influence of SvFul2 on inflorescence development begins once the SAM has initiated transition to the IM (Figure 6A). We also observed dynamic shifts in DEGs among the three stages; only 33 DE genes were shared across all three stages, and 149, 451, and 68 were shared between Stages 1 and 2, Stages 2 and 3, and Stages 1 and 3, respectively (Figure 6B). This suggests that SvFul2 potentially modulates different target genes in various spatiotemporal contexts. Indeed, indirect effects due to differences in cell populations, especially at Stage 3 after phenotypes of Svful2 and wild-type inflorescences have diverged, likely underlie a portion of stage-specific DEGs.

As expected, the SvFul2 gene itself was significantly downregulated in mutant inflorescences at all three stages (Figure 6C). The other three S. viridis AP1/FUL-like genes were significantly upregulated in the mutant, suggesting that those of the brl1 mutant (Figure 4, B–G). Compared with ME034 normal plants, SvFul2_KOs were shorter and branchy with more leaves (Figure 4B; Supplemental Table S1), and panicles displayed increased densities of longer primary branches (Figure 4, C and F; Supplemental Table S1).
the four AP1/FUL-like genes may provide some level of functional compensation during inflorescence development. Two B-class genes, SvMads16 (AP3) and SvMads4 (PISTILLATA), which in grasses are typically expressed at low levels prior to floral organ development (Whipple et al., 2004), were upregulated in Svful2 Stage 2 inflorescences (Figure 6C). In addition, two E-class genes were differentially expressed in mutant inflorescences: SvMads34 was upregulated at Stages 1 and 2, while SvMads5 was downregulated at Stage 2 (Figure 6C). In rice, OsMADS34 coordinates with AP1/FUL-like genes, and physically interacts with some of them, to specify IM identity (Kobayashi et al., 2012). In general, E-class genes play partially redundant roles in specifying floral organ identities via protein–protein interactions with other MADS box proteins (Pelaz et al., 2000; Honma and Goto, 2001; Theissen and Saedler, 2001; Ditta et al., 2004).

Since the transition to reproductive growth is delayed in Svful2 mutants, we expected to see changes in genes and pathways associated with flowering time (Figure 6D). Functional categories related to flowering time were overrepresented among DEGs post-transition at Stage 2, including “vegetative to reproductive phase transition of meristem” (GO:0010228; P adj = 3.75e-07) and “vernalization response” (GO:0010048; P adj = 3.68e-06; Figure 6E; Supplemental Data Set S6). Upregulated DEGs included homologs of AP2-like genes in maize known to suppress indeterminate growth in the SM, including indeterminate spikelet1 (Sevir.9G034800) and sister of indeterminate spikelet1 (SvSid1, Sevir.2G093800; Supplemental Figure S8; Chuck et al., 1998, 2007, 2008). The homolog of rice MOTHER OF FT AND TFL1 (Sevir.4G169200, Svmf1), which represses SM identity, was also upregulated in Svful2, and the ortholog of maize ramosa2 (SvRd2; Sevir.5G116100), which functions to promote meristem determinacy (Bortiri et al., 2006), was downregulated (Supplemental Figure S8).

As a consequence of increased meristem indeterminacy, Svful2 mutant inflorescences branch more. We also found that genes associated with “anatomical structure formation involved in morphogenesis” (GO:0048646) were overrepresented among DEGs at stages 1 (P(adj) = 0.0027) and 2 (P(adj) = 3.65e-06; Figure 6E; Supplemental Data Set S6), consistent with enhanced expression of genes involved in organogenesis. Among this functional class were putative orthologs of known genes that specify abaxial cell fate, for example, Sevir.1G255800 (SvYab1S) and milkweed pod1 (Sevir.6G158800; Candela et al., 2008), and adaxial cell fate,
Figure 6 Transcriptional changes in Svful2 mutants across three stages of inflorescence development. A, PCA showed that biological replicates were well-correlated with each other and that PC1 (explaining 56% of the variance) was associated with developmental stage. Loss-of-function in Svful2 resulted in fewer transcriptional changes prior to the floral transition with larger changes between genotypes appearing later in development. B, DEGs showed dynamic transcriptional changes across three stages of inflorescence development in Svful2 mutants. Among DEGs were several encoding MADS-box TFs (C) and known regulators of flowering time (D). TPM values were Log2 transformed to generate heatmaps. Yellow and black asterisks indicate up and downregulated DEGs (FDR < 0.05), respectively. E, Subsets of GO terms that were overrepresented among DEGs at each of the three developmental stages. *P* adj < 0.05.
for example, *Rice outermost cell-specific gene5 (Sevir.SG077800)* and *PHABULOSA (Sevir.9G157300)*; *McConnell et al., 2001; Juarez et al., 2004; Zou et al., 2011*), which were upregulated in the mutant (Supplemental Figure S8). Alternatively, the putative ortholog of rice *DWARF3, SvD3 (Sevir.4G068300)*, which functions in suppression of branching through the strigolactone signaling pathway (*Zhou et al., 2013*), was downregulated (Supplemental Figure S8). The major transcriptome changes observed in loss-of-function mutants at Stage 2 reflect a core function for *SvFUL2* in modulating the reproductive transition at the molecular level, but also how it links delayed flowering to suppression of meristem determinacy programs.

**Transcriptional rewiring by perturbation of *SvFUL2* reveals sub-networks connecting reproductive transition and determinacy pathways**

To further investigate how *SvFUL2* connects within a larger gene network to regulate flowering time and meristem determinacy pathways, we used a computational strategy based on weighted gene co-expression network analysis (WGCNA) and a random forest classifier to construct a gene regulatory network (GRN) representing normal inflorescence development in *S. viridis* (A10.1). Here, we integrated RNA-seq data from a previous study that captured precise stages of A10.1 inflorescence primordia spanning the IM transition to the development of floral organs (*Zhu et al., 2018*) with the staged wild-type data collected in this study. Using the WGCNA algorithm (*Langfelder and Horvath, 2008*), we clustered 26,758 genes into 27 co-expression modules (Figure 7A; Supplemental Figure S9 and Supplemental Data Set S7). Module eigengenes (MEs; expression pattern that best fits an individual module) were evaluated for their significant associations with four key developmental events represented in the network: the vegetative-to-reproductive transition (8 and 10 DAS), branching (11, 12, and 14 DAS), meristem determinacy (15–17 DAS), and flower development (18 DAS; Figure 7A). Within each module, we tested for enrichment of genes that were differentially expressed in the *SvFUL2* mutant, and found several that showed enrichment during key developmental events (Figure 7A). Among these, MEmagenta showed a strong positive correlation with the floral transition and a negative correlation with meristem determinacy (Figure 7, A and B). MEmagenta showed enrichment for DEGs in Stages 1 and 2 (Figure 7A). Network analysis of this module revealed *SvMads51 (Sevir.5G438700)*, which is orthologous to *mads69* from maize, was predicted as a hub node in control of many genes, including the regulator of meristem determinacy *SvSid1* (*Liang et al., 2019; Supplemental Figure S10*). In maize, *mads69* has been associated with both floral transition and meristem determinacy. It is located in the maize QTL Vgt3 associated with flowering time and latitudinal variation, and it was recently validated as the transcriptional regulator of FT genes. *ZmMADS69* was also associated with tassel branch number and tassel branch length (*Liang et al., 2019*). Alternatively, MEbrow showed a positive correlation with meristem determinacy, but was negatively correlated with the floral transition (Figure 7, A and B). *SvFUL2* and *SvFUL3* were both co-expressed in the brown module, along with 428 DEGs largely at Stages 2 and 3.

We also integrated the co-expression network with information derived from regulatory interactions among TFs and their putative targets based on the GENIE3 algorithm (*Huynh-Thu et al., 2010*). This complementary approach helped us to infer the directionality and connectivity of important hub genes within the gene network, yielding a directed graph of regulatory interactions. Based on the assumption that expression of a given gene is a function of the expression of the other genes in the network, we applied a regression trees method using as inputs TFs expressed in our dataset (*n = 1,295*) selected based on PlantTFDB (*Jin et al., 2017*), their expression trajectories in the network and those of other genes, to identify potential TF targets (*Huynh-Thu et al., 2010*). Regulatory genes and their predicted targets were restricted based on information from differential expression analysis between wild-type and the *SvFUL2* mutant. We used the resulting regulatory framework to explore functional relationships between *SvFUL2*, its predicted direct targets, and predicted upstream regulators, particularly in the context of connecting floral transition and meristem determinacy. For example, magenta and brown modules were connected through a bHLH TF (Sevir.2G248500) that was predicted to directly target both *SvMADS51* and *SvFUL2* (*Liang et al., 2019*), placing it as a potential regulator at the intersect of flowering time and meristem determinacy (Supplemental Figure S11).

Our GRN also predicted that *SvFUL2* controls several co-expressed TFs previously implicated in developmental processes, and localized to modules that positively associated with branching/meristem determinacy and negatively associated with floral transition (Figure 7C). Among these were *SvRA2*, and an INDETERMINATE DOMAIN TF, several members of which have been involved in both the floral transition and determinacy, including the founding member from maize, *indeterminate 1 (id1; Colasanti et al., 1998; Kozaki et al., 2004)*. Our analyses point to a possible feedback loop mechanism between *SvFUL2* and *SvRA2*, where *SvFUL2* is also a predicted direct target of *SvRA2*. We also observed putative feedback regulation between *SvFUL2* and TFs encoded by putative orthologs of maize *knotted 1 (kn1; TALE TF, Sevir.9G107600)* and *fasciated ear 4 (fea4; bZIP TF, Sevir.4G119100)*, which promote meristem maintenance and differentiation, respectively (*Bolduc et al., 2012; Pautler et al., 2015*). Extensive feedback among these developmental TFs could represent endogenous mechanisms for fine-tuning development during the floral transition and patterning of the inflorescence.

**Discussion**

According to the classic ABCDE model of floral development in Arabidopsis, A-class AP1/FUL-like MADS-box genes
Network analysis reveals transcriptional rewiring in the $Sufu2$ mutants that links reproductive transition and determinacy pathways. A, (left) Heatmap represents the WGCNA ME association with key events during early inflorescence development (vegetative-to-reproductive transition, branching, meristem determinacy, and flower development) in $S. viridis$. Network modules are represented and named with different colors based on the WGCNA default module annotation. Number of genes co-expressed in each module is indicated to the left. Student asymptotic $P$ for the ME association are indicated: *** $P < 0.001$; ** $P < 0.01$, and * $P < 0.05$. (right) Heatmap represents enrichment of $Sufu2$ DEGs at each of the...
have essential functions in modulating the floral transition and floral organ development (Irish and Sussex, 1990; Ferrándiz et al., 2000; Kaufmann et al., 2010). The roles of A-class genes have been the focus of extensive evolution and development studies (Litt and Irish, 2003; Preston and Kellogg, 2007); however, relatively little is known about their functions in grasses. Given the complex branching patterns that arise post-floral transition and prior to flower development in grasses, it is expected that there would be some variation in function. In general, grasses show subtle variations on the traditional ABCDE model, however, the underlying mechanisms are generally conserved (Ambrose et al., 2000; Whipple et al., 2007). Functional redundancy among A-class genes is widespread in grasses, and only recently have simultaneous perturbations in multiple paralogs revealed informative mutant phenotypes, for example, in rice and wheat (Wu et al., 2017; Li et al., 2019). So far there have been no inflorescence phenotypes reported for AP1/FUL-like genes in any panicoid species, which include major cereal and energy crops. Therefore, we know little about their specific functions in regulating important agronomic traits such as flowering time and inflorescence determinacy.

In this study using S. viridis as a model, we characterized a loss-of-function mutant in an AP1/FUL-like gene, SvFUL2, that displayed strong developmental phenotypes, which was unexpected for a single mutant allele. Our morphological and molecular analyses of the Svful2 mutant provide insights into the roles of AP1/FUL-like genes in connecting flowering time and inflorescence determinacy in panicoid grasses, as well as predictions on conserved and novel regulatory interactions underlying the complex phenotypes.

SvFUL2 is necessary for proper timing of flowering and determinacy programs

Phylogenetic studies have reconstructed the evolutionary history of AP1/FUL-like genes in angiosperms (Litt and Irish, 2003; Preston and Kellogg, 2006; Soltis et al., 2007; Wu et al., 2017). The monocot AP1/FUL-like clade members evolved independently after the split of monocots and eudicots (Litt and Irish, 2003; Preston and Kellogg, 2007). In the Poaceae clade, four copies of AP1/FUL-like genes are derived from three duplication events in the AP1/FUL lineage. The first likely occurred during early monocot evolution, giving rise to the FUL3 clade. The second occurred near the base of the Poaceae, which generated the FUL1 and FUL2 clades (Preston and Kellogg, 2006). The last duplication produced the FUL3 and FUL4 clades, and FUL4 was lost in some grass species during evolution (Wu et al., 2017). Such duplication events can lead to functional redundancy and subsequent diversification. In grasses, AP1/FUL-like genes are expressed much earlier than in eudicots (Preston and Kellogg, 2007), and their transcripts have been detected in IM, BMs, and SMs in addition to floral organs. In several grass species studied, FUL1/VRN1/OsMADS14 and FUL2/OsMADS15 have redundant and/or overlapping spatiotemporal expression patterns in these three meristem types, yet show different patterns within the spikelet. This suggests that in certain grasses, these two genes play redundant roles during the floral transition and in SM identity, but diversified roles in floral organ identity.

Among the grasses, AP1/FUL-like genes have been most studied at the functional level in rice and wheat (Wu et al., 2017; Li et al., 2019), where clear functions in flowering time have been demonstrated. The role of SvFUL2 in controlling flowering time is consistent with the significant accumulation of SvFUL2 transcripts (over 100-fold change in expression in the IM) during the transition from SAM to IM (Figure 6C). SvFUL1 and SvFUL3 were also induced during this time (Figure 6C), but fold changes were not as large as for SvFUL2, similar to what has been shown in rice (Kobayashi et al., 2012). Based on our results and previous studies, the accumulation of AP1/FUL-like transcripts in the IM upon the induction of FT is likely required for promoting the reproductive transition, and this is a conserved mechanism in grasses.

The shift from a determinate to indeterminate fate in the IM of Svful2 mutants (Figures 1F and 2H), which was also observed in the wheat vrn1-null; ful2-null double mutant, is reminiscent of the tfl1 mutant phenotype in Arabidopsis (Shannon and Meeks-Wagner, 1991). Previous studies that examined spatiotemporal expression of FUL1/VRN1/OsMADS14 and FUL2/OsMADS15 in phylogenetically disparate grasses, showed that they are most abundantly expressed in the tip of the IM (Preston and Kellogg, 2007). In both Svful2 and wheat vrn1-null; ful2-null mutants, significant increases in the expression of TFL1 homologs were detected (Figure 6D; Li et al., 2019). These results suggest that the mechanism for controlling IM determinacy in grasses involves an antagonism between AP1/FUL-like genes and TFL1-like genes, as in eudicots. IM determinacy appears to be very sensitive to the activity of AP1/FUL-like genes. In wheat, complete loss of both VRN1 and FUL2 function leads to an indeterminate IM, while a single functional copy of VRN1 or FUL2 in a heterozygous state was able to recover a determinate IM. It has been proposed that indeterminate growth in the IM was derived from a determinate habit in evolution, which involved the modification and/or loss of an early common TFL1 mechanism (Bradley et al., 1997). This hypothesis could explain this apparent sensitivity.

Figure 7 (continued)

three developmental stages profiled in the mutant among the MEs. Gold and black asterisks indicate up- and downregulated genes, respectively, in the Svful2 mutant compared to wild-type. B. Expression trajectories of MEmagenta and MEBrown across S. viridis inflorescence development. Y-axis represents the average TPM of co-expressed genes in each module. C. Subnetwork of predicted direct targets of SvFUL2 and its direct upstream regulators based on our GRN. Genes are represented as circles with edges linking specific regulators to their targets. Differential fold change of expression in the Svful2 mutant background is represented by the colored scale. Darker colors represent higher [fold change].
The strong phenotypes we observed in *S. viridis* by a single knockout of an AP1/FUL-like gene indicate its central role in controlling multiple developmental processes. Interestingly, the co-expression of closely related paralogs, *SvFUL1* and *SvFUL3*, with *SvFUL2* does not seem to provide much functional compensation, but we did see both genes upregulated upon *SvFUL2* perturbation (Figure 6C). *SvFUL2* was expressed at high levels (highest among other AP1/FUL-like genes) at most of the developmental stages we examined (Figure 3B). The localized expression of *SvFUL2* mRNAs was previously analyzed by in situ hybridization in a study by (Preston and Kellogg, 2007) to evaluate developmental expression patterns of AP1/FUL-like genes in inflorescence primordia across grasses. They showed that *SvFUL2* transcripts accumulated largely in the IM during early inflorescence development, consistent with its role in floral transition and determinacy. In addition, transcripts were detected in BMs and SMs, also consistent with indeterminacy in these meristem types in the loss-of-function *SvFUL2* mutant. Expression patterns of *SvFUL1* and *SvFUL2* overlapped in these meristems but interestingly, diverged in developing FMs (Preston and Kellogg, 2007). *SvFUL2* showed distinct expression in marking floral organ primordia of the first two whorls, however, in our analyses of the *SvFUL2* mutant, development of these organs appeared unaffected. This suggests that other factors potentially compensate for loss of *SvFUL2* in proper floral organ development in *S. viridis*.

The functional redundancy of AP1/FUL-like genes in grasses provides an opportunity for diversification of function, and a toolkit for fine-tuning development of desired traits.

**SvFUL2 as an integrator of flowering time and inflorescence determinacy**

Connections between flowering time signals and meristem determinacy pathways in the inflorescence have been highlighted in various grass species. A strong flowering signal can impose meristem determinacy when perceived by the developing inflorescence (Dixon et al., 2018). For example, wheat *Ppd-1*, which functions in a *Ppd*-dependent floral induction pathway, suppresses paired spikelet formation through modulation of FLOWERING LOCUS T (Boden et al., 2015). The paired spikelet phenotype is associated with enhanced indeterminacy. In maize, loss-of-function in *id1*, a key player in the floral transition, leads to complete loss of meristem determinacy; instead of floral organs, plantlets are developed from every spikelet in mutant tassels (Colasanti et al., 1998). Meristem identity genes, for example, AP1/FUL-like genes, have been proposed to function downstream of the flowering signal to promote meristem determinacy and reshape inflorescence architecture (Dixon et al., 2018).

In our study, the important role of *SvFUL2* in coordinating flowering time and meristem determinacy is not only supported by its strong pleiotropic phenotypes, but also reflected in our predictions of regulatory relationships between *SvFUL2* and its upstream modulators and downstream targets. Several MADS-box TFs, most of which are homologs to those implicated in flowering time, were predicted to directly target *SvFUL2* (Figure 7C). We also identified a bHLH TF of unknown function predicted to target both *SvFUL2* and *Svmads51*, which is the syntenic ortholog of the maize flowering time regulator *mads69*, and therefore a potential integrator of floral transition and meristem determinacy (Supplemental Figure S11). Interestingly, several other MADS-box TFs were shown to directly target *SvFUL2* based on predictions in our GRN: *SvMADS37.1* (Sevir. 6G230800), *SvMADS56* (Sevir.9G347400), *SvMADS5* (Sevir.4G060800), *SvMADS34/PAP2* (Sevir.9G087100), and *SvFUL1* (Figure 7C). *SvMads34/Pap2* was also predicted to be a direct target of *SvFUL2*. These predictions are consistent with previous studies showing regulatory interactions among MADS-box TFs during the floral transition and inflorescence development. In rice, *OsMADS37* and *OsMADS56* have been functionally characterized as flowering time regulators. *OsMADS34/PAP2* has been shown to function redundantly with rice AP1/FUL-like genes to promote flowering and SM determinacy (Kobayashi et al., 2010, 2012), and *OsMADS5* is involved in spikelet identity (Wu et al., 2018). Our analysis indicates that *SvFUL2* may function as a core integrator at the interface of these closely linked developmental programs through feedback coordination with several other developmental TFs.

Our network analysis also uncovered potential feedback regulation between *SvFUL2* and *SvRA2* (Figure 7C), which could point to a conserved mechanism by which flowering links to AM determinacy in grasses. *SvRA2* is the ortholog of maize *ra2* and barley *Six-rowed spike4* (*Vrs4*; Bortiri et al., 2006; Koppolu et al., 2013). Both *ra2* and *Vrs4* function in imposing determinacy on spikelet pair meristems and triple SMs in maize and barley, respectively. Although several downstream targets of *ra2* and *Vrs4* have been identified through genetic and/or transcriptomics analyses (Bortiri et al., 2006; Bai et al., 2012; Koppolu et al., 2013; Eveland et al., 2014), upstream regulators have not been described. Unlike other genes in the RAMOSA pathway, *ra2* function is highly conserved across grasses and expresses early during AM initiation, and temporally after the expression of AP1/FUL-like genes (Bortiri et al., 2006; Koppolu et al., 2013; Zhu et al., 2018). In addition, localization studies have shown that *ra2* and *SvFUL2* are expressed in overlapping domains within BMs during early inflorescence development (Bortiri et al., 2006; Preston and Kellogg, 2007; Koppolu et al., 2013). The conserved spatiotemporal expression pattern of *ra2* is consistent with it being downstream of FUL2 to potentially coordinate the flowering signal with regulation of meristem determinacy. Further functional studies are required to determine the genetic and molecular interactions between RA2 and FUL2.

In maize, over-expression of the maize AP1/FUL-like gene, *zmm28*, enhanced grain yield potential through improved photosynthetic capacity and nitrogen utilization (Wu et al., 2019). Direct targets of *ZMM28* revealed through integrated RNA- and ChIP-seq analyses included genes involved in
photosynthesis and carbohydrate metabolism. Homologs of several of these targets were differentially expressed in Sfu12 mutants, including *photosystem I light-harvesting complex gene 6* (Sevir.2G22720), a gene encoding a pyruvate orthophosphate dikinase (Sevir.3G253900), and gene encoding a bZIP TF (Sevir.3G396500). Although Sfu12 encodes a different AP1/FUL-like gene in a different spatiotemporal context, we also observed changes in genes associated with photosynthesis and with starch and sugar metabolism in Stage 3 inflorescences where the mutant was highly indeterminate compared to wild-type. There could be common regulatory interactions between AP1/FUL-like genes associated with photosynthesis, carbon allocation, and sugar signals that link flowering time cues from the leaf to inflorescence architecture. We know little about the mechanisms by which sugar signals interface with development, but clear links, for example with trehalose-6-phosphate, underlie flowering time (Wahl et al., 2013), and meristem determinacy (Satoh-Nagasawa et al., 2006).

The striking phenotype displayed in loss-of-function Sfu12 mutants enables us to more clearly define molecular connections between flowering time and various aspects of IM determinacy. One question that comes to mind is why do the pathways regulated by Sfu12 in *S. viridis* have fewer checks and balances in terms of functional redundancy compared to other grasses? Since *S. viridis* is an undomesticated weed, one hypothesis is that selection against indeterminacy phenotypes in inflorescences of modern cereal crops species masks the ability to recover individual functions of A-class genes at the phenotypic level. Furthermore, perhaps the phenotypes presented in Sfu12 mutants provide plasticity in *S. viridis*’s adaptability to a wide range of environmental conditions. In any case, our analyses of this mutant provide a glimpse into AP1/FUL-like gene function in panicle grasses and predict regulatory interactions linking key yield traits that can be translated to important cereal and energy crops.

**Methods**

**Plant materials and growth conditions**

The *br1l-ref* mutant allele was isolated from an NMU mutagenized M2 population of *S. viridis* (Huang et al., 2017). The mutant allele was backcrossed to the reference mutagenized line (A10.1) and selfed to generate F2 segregating populations. F4 seeds were used for phenotyping, SEM, and RNA-seq experiments. *Setaria viridis* plants for phenotyping were grown under either SD (12-h light/12-h dark) or LD (16-h light/8-h dark) conditions (31°C/22°C [day/night], 50% relative humidity, and light intensity of 400 µmol/m²/s) in a controlled high-light growth chamber at the Danforth Center’s growth facility. *Setaria viridis* plants used for SEM and RNA-seq were grown under the SD conditions.

**SEM analysis**

For SEM analysis, *br1l* mutant and wild-type inflorescence primordia were harvested from young seedlings to examine the developmental defects of mutants. Samples were fixed, hand-dissected, and dehydrated as described (Hodge and Kellogg, 2016). The dehydrated samples were critical point dried using a Tousimis Samdri-780a and imaged by a Hitachi S2600 SEM at Washington University’s Central Institute of the Deaf.

**Histology**

Wild-type and mutant inflorescence primordia were harvested right after the vegetative-to-reproductive transition at 11 and 15 DAS, respectively. The samples were fixed, embedded, and sectioned as described by (Yang et al., 2018). Sections (10 µm) made with a Microm HM 355S microtome (ThermoFisher, Waltham, MA, USA) were deparaffinized, stained with eosin, and imaged with a ZEISS AxioZoom microscope.

**Bulked segregant analysis**

M3 mutant individuals were crossed to the A10.1 reference line and resulting F1 individuals were self-pollinated to generate segregating F2 families. The F2 individuals with mutant and wild-type phenotypes were identified, and the segregation ratio was tested by a χ² test. DNA extracted from 30 *br1l* mutant individuals was pooled to generate a DNA library. The DNA library was made using the NEBNext Ultra DNA Library Prep Kit for Illumina (NEB), size selected for inserts of 500–600 bp, and sequenced with 150-bp paired-end using standard Illumina protocols on Illumina Hi-Seq 4000 platform at Novogene. DNA libraries for whole-genome sequencing were generated from a single *br1l* mutant individual in the M2 generation and sequencing with 100-bp single end on Illumina Hi-Seq 2,500 platform at University of Illinois, Urbana-Champaign W.M. Keck sequencing facility. Read mapping and SNP calling by GATK (3.5-0-version 3.5-0-g36282) were performed as described (Huang et al., 2017).

**Phylogenetic analysis**

The coding sequences of Arabidopsis, *S. viridis*, maize, sorghum, rice, and wheat AP1/FUL-like family genes were obtained from Phytozome (phytozome.jgi.doe.gov; Supplemental Data Set S2) and aligned using ClustalW to build a maximum likelihood tree with bootstrapping (1,000 iterations) in MEGA7 (Kumar et al., 2016).

**CRISPR/Cas9 gene-editing**

The genome sequence of *SfU12* (Sevir.2G006400) was obtained from the *S. viridis* version 2.1 genome (https://phytozome.jgi.doe.gov/). CRISPR-P version 2.0 (Liu et al., 2017) was used to design guide (g)RNAs to minimize off-targets. Two gRNAs targeting *Sfu12* were designed at the first exon and the first intron, 133- and 395-bp downstream of the ATG start codon, respectively. Using a plant genome engineering toolkit (Čermák et al., 2017), gRNAs were combined into a level 0 construct followed by insertion into a plant transformation vector. PCR amplified fragments from pMOD.B_2303 were merged using golden-gate cloning with T7 ligase and SapI/BsmBI restriction enzymes back into the
pMOD_B_2303 backbone to express the two gRNAa from the CmYLCV promoter, each flanked by a tRNA. This construct, along with pMOD_A1110 (a wheat codon-optimized Cas9 driven by the ZmUbi1 promoter) and pMOD_C_0000 modules, were combined in a subsequent golden-gate cloning reaction with T4 ligase and AarI restriction enzyme into the pTRANS_250d plant transformation backbone. The final construct was cloned into Agrobacterium tumefaciens line AGL1 for callus transformation of S. viridis ME034 at the DDPSC Tissue Culture facility. T0 plantlets were genotyped for the presence of the selectable marker, hygromycin phosphotransferase to validate transgenic individuals. In the T1 generation, individual plants with possible mutant phenotypes were selected and the region of the target sites was amplified using PCR and sequenced. A homozygous 540-bp deletion in the first exon of SvFul2 was identified. These T1 mutants were self-pollinated to obtain T2 progeny and outcrossed to ME034 and then selfed to select Cas9-free SvFul2_KO plants. Primer sequences used for vector construction and genotyping are listed in Supplemental Table S2.

RNA-seq library construction, sequencing, and analysis

Poly-A⁺ RNA-seq libraries were generated from pools of hand-dissected inflorescence primordia from wild-type and Svful2 mutant seedlings. Wild-type primordia were sampled at 8, 11, and 17 DAS while, accounting for the mutant’s developmental progression, Svful2 primordia were sampled at 9, 15, and 21 DAS. For each developmental stage, four biological replicates were collected, for a total of 24 data points.

RNA was extracted (PicoPure RNA isolation kit; Thermo Fisher Scientific) and subjected to library preparation from 500 ng of total RNA using the NEBNext Ultra Directional RNA Library Prep Kit (Illumina, San Diego, CA, USA), size-selected for 200-bp inserts, and quantified on an Agilent bioanalyzer using a DNA 1,000 chip. RNA-seq libraries were processed using an Illumina HiSeq 4000 platform at Novogene with a 150-bp paired-end sequencing design. On average, for each data point ~20 million cleaned reads were generated. RNA-seq reads were quality checked and processed using the wrapper tool Trim Galore (version 0.4.4_dev) with the parameters “—length 100 —trim-n —illumine.” Cleaned reads were mapped to the S. viridis transcriptome (Sviridis_500_version 2.1; Phytozome version 12.1, phytozome.jgi.doe.gov) using Salmon (0.13.0) with the parameters “—validateMappings —numBootstraps 30,” based on an index generated by primary transcripts (n = 38,209). Gene normalized expression levels (transcript per kilobase million [TPM]; Supplemental Data Set S3) and the count matrix for downstream analyses were determined from Salmon output files and imported in R using the Bioconductor package tximport (Soneson et al., 2015).

Sample variance was computed based on PCA with the function dist and plotPCA on variance stabilizing transformation (vst) scaled data. Analyses of differential expression were performed using the Bioconductor package DESeq2 (version 1.22.2) with default parameters for the Wald test. The Benjamini–Hochberg method for multiple testing correction was used to classify DEGs passing the P-value adjusted cut-off of 0.05.

For GO enrichment analysis, we generated a refined S. viridis GO annotation (Supplemental Data Set S5) using the GOMAP pipeline (https://gomap-singularity.readthedocs.io; Wimalanathan and Lawrence-Dill, 2019) to determine over-representation of GO terms within gene sets with the Bioconductor package topGO. GO testing was performed based on the Fisher’s exact test method.

DE genes enriched in GCN modules were obtained based on the enrichment analysis using the function enricher from the Bioconductor package clusterProfiler (Yu et al., 2012) Benjamini–Hochberg multiple test corrections.

Weighted gene co-expression network analysis

In addition to the samples described above, we included previously described wild-type S. viridis inflorescence primordia samples (Zhu et al., 2018): 23 additional data points from 6 inflorescence stages (10, 12, 14, 15, 16, and 18 DAS). This dataset (GSE118673) was re-processed using the same methods described above and used to build a reference wild-type gene co-expression network spanning S. viridis inflorescence organogenesis, from the transition to reproductive phase to flower development. To reduce samples bias, we first filtered out genes with less than 10 counts (row sum ≤10), then we calculated the Euclidean distance and Pearson’s correlation among samples and removed all replicates with rho coefficient <0.92 or with a Euclidean score <0.8. Based on this, two samples were removed (8 DAS rep 4 and 17 DAS rep 3). Read counts from genes (n = 26,758) and samples (n = 33) passing the above filters were normalized with vst using the function vst from the Bioconductor package DESeq2.

A signed co-expression network was built using the blockwiseModules function from the WGCNA R package (Langfelder and Horvath, 2008) with the parameters: “power = 16, corType = “bicor”, minModuleSize = 30, mergeCutHeight = 0.25, maxBlockSize = 30,000, MaxPoutlier = 0.05, minModuleSize = 20.” The topological overlap matrix was calculated from the blockwiseModules function using the parameter “TOMType = ‘signed.’”

The module-to-developmental stage association was conducted evaluating the significance correlation of the ME and four key developmental stages defined as: (1) vegetative-to-reproductive transition (8 and 10 DAS); (2) branching (11, 12, and 14 DAS); (3) meristem determinacy (15–17 DAS); and (4) flower development (18 DAS). To conduct this analysis, we created a metafile where all samples were classified according to the four key stages. The R function cor and corPvalueStudent were used to test the correlation between ME and the stages. Rho values were used to identify relationship between modules and developmental stages.
Modules with $\rho > 0.8$ were considered strongly correlated to the developmental stages.

To predict targets of *S. viridis* TFs we built a complementary network using a machine learning approach with the Bioconductor package GENIE3 (Huynh-Thu et al., 2010). *Setaria viridis* TFs were downloaded from PlantTFDB (http://planttfdb.gao-lab.org; Jin et al., 2017) and overlapped with the expression matrix used in the WGCNA analysis to identify the expressed TFs in our dataset ($n = 1,265$). These TFs were used as probes to predict regulatory links between the putative targets and their expression trajectories in our dataset. We ran GENIE3 with the parameters “treeMethod = "RF", nTrees = 1,000” and putative target genes were selected with a weight cutoff $\geq 0.005$. Networks were explored and plotted using the R package igraph.

**Data accessibility**

All sequence data, including whole-genome sequence data, raw and processed RNA-seq data, and associated metadata have been deposited in the NCBI Gene Expression Omnibus under accession number GSE156047.

**Accession numbers**

GenBank accession numbers for *S. viridis* AP1/FUL-like genes are as follows: *SvFul1* (XM_034716904; Sevir.9G087300), *SvFul2* (XM_034727736; Sevir.2G006400), *SvFul3* (XM_034726136; Sevir.2G393300), and *SvFul4* (XM_004962888; Sevir.3G74400). A more complete list of accession numbers for AP1/FUL-like genes discussed in this manuscript can be found in Supplemental Data Set S2.

**Supplemental data**

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

**Supplemental Figure S1.** Additional characteristics of the *brl1* mutants (supports Figure 1).

**Supplemental Figure S2.** BSA analysis of the *brl1* locus.

**Supplemental Figure S3.** Genotyping of *brl1* F$_2$ seedlings using a designed dCAPS marker for the SNP located at the start codon of *SvFul2*.

**Supplemental Figure S4.** RT-PCR results showing the reduced *SvFul2* expression level in mutant inflorescence primordia compared with A10.1.

**Supplemental Figure S5.** PCR genotyping of the *SvFul2_KO* CRISPR edited line.

**Supplemental Figure S6.** Morphological analysis of early inflorescence development in *SvFul2_KO* by SEM.

**Supplemental Figure S7.** Three key developmental transitions of *brl1* mutant and wild-type inflorescence development captured by RNA-seq profiling gene expression.

**Supplemental Figure S8.** Dynamic expression differences of genes related to meristem maintenance, plastochnon, abaxial/adaxial cell fate, and meristem determinacy between wild-type and *SvFul2* mutant.

**Supplemental Figure S9.** Dendrogram representing relatedness among genes based on expression across all samples and their respective module assignments (indicated by color classification).

**Supplemental Figure S10.** Gene co-expression subnetwork of the magenta module showing the relationship among key transcriptional regulators and hub genes.

**Supplemental Figure S11.** Gene regulatory sub-network showing predicted targets of *SvMADS51* and its direct upstream regulators.

**Supplemental Table S1.** Phenotypic measurements of *SvFul2-KO* plants

**Supplemental Table S2.** Table of primers used in this study

**Supplemental Data Set S1.** High-confidence SNP calls for the *brl1* mutants.

**Supplemental Data Set S2.** Alignment of coding sequences of *AP1/FUL-like* genes by ClustalW.

**Supplemental Data Set S3.** Transcript abundances (TPM) for all annotated *S. viridis* genes (version 2) in the wild-type compared with *Svful2* mutant inflorescence primordia.

**Supplemental Data Set S4.** DEGs with annotation at three developmental stages determined by DESeq2.

**Supplemental Data Set S5.** GOMAP for *S. viridis*.

**Supplemental Data Set S6.** Overrepresentation of functional classes among DEGs based on GO term enrichment.

**Supplemental Data Set S7.** Weighted gene co-expression gene network generated with the R package WGCNA (available at GEO: GSE156047).

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