Somatic DNA demethylation generates tissue-specific methylation states and impacts flowering time

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Abstract

Cytosine methylation is a reversible epigenetic modification of DNA. In plants, removal of cytosine methylation is accomplished by the four members of the DEMETER (DME) family of 5-methylcytosine DNA glycosylases, named DME, DEMETER-LIKE2 (DML2), DML3, and REPRESSOR OF SILENCING1 (ROS1) in Arabidopsis thaliana. Demethylation by DME is critical for seed development, preventing experiments to determine the function of the entire gene family in somatic tissues by mutant analysis. Here, we bypassed the reproductive defects of dme mutants to create somatic quadruple homozygous mutants of the entire DME family. dme; ros1; dml2; and dml3 (drdd) leaves exhibit hypermethylated regions compared with wild-type leaves and rdd triple mutants, indicating functional redundancy among all four demethylases. Targets of demethylation include regions co-targeted by RNA-directed DNA methylation and, surprisingly, CG gene body methylation, indicating dynamic methylation at these less-understood sites. Additionally, many tissue-specific methylation differences are absent in drdd, suggesting a role for active demethylation in generating divergent epigenetic states across wild-type tissues. Furthermore, drrd plants display an early flowering phenotype, which involves 5′-hypermethylation and transcriptional down-regulation of FLOWERING LOCUS C. Active DNA demethylation is therefore required for proper methylation across somatic tissues and defines the epigenetic landscape of intergenic and coding regions.

Introduction

Phenotypic stability depends on epigenetic stability. Cytosine DNA methylation is both stably inherited and dynamic. In many plant species, DNA methylation is concentrated in transposable elements and other repetitive sequences, transcriptionally silencing these genomic regions. DNA methylation also accumulates in gene coding regions, where it does not induce transcriptional silencing (Niederhuth et al., 2016; Bewick and Schmitz, 2017). Methylation patterns can be highly conserved over evolutionary time but also dynamic in specific developmental contexts, such as during reproduction (Gehring, 2019). These patterns are a result of methylation activity and demethylation activity, the coordination of which is required to maintain transgenerational epigenetic and phenotypic stability (Williams and Gehring, 2017). Methylation is established by de novo DNA methyltransferases, which are guided to their targets by small RNAs in a process termed RNA-directed DNA methylation (RdDM) (Matzke and Mosher, 2014). Symmetric DNA methylation is maintained by
DNA demethylation is initiated by the activity of a family of bifunctional HhH-GPD DNA glycosylases/lyases (Agius et al., 2006; Gehring et al., 2006; Morales-Ruiz et al., 2006; Penterman et al., 2007; Ortega-Galisteo et al., 2008). In Arabidopsis thaliana, there are four gene family members: DEMETER (DME), REPRESSOR OF SILENCING1 (ROS1), DEMETER-LIKE2 (DML2), and DML3 (Choi et al., 2002; Gong et al., 2002; Penterman et al., 2007; Ortega-Galisteo et al., 2008). These enzymes remove methylated cytosines through base excision repair (Roldán-Arjona et al., 2019). DME and ROS1 genes are present in plants from algae onward (Pei et al., 2019). The distribution of DML2 and DML3 is more limited, with DML3 present in monocots and dicots and DML2 present only in a subset of dicots (Pei et al., 2019).

Molecular and morphological phenotypes have been described for plants with mutations in one or more of the 5-mC DNA glycosylases in Arabidopsis and rice (Oryza sativa). Arabidopsis plants heterozygous for dme mutations have a visibly striking phenotype: 50% seed abortion (Choi et al., 2002). Seeds that inherit a mutant dme allele from the mother abort after several days of development and thus homozygous dme mutants have only rarely been recovered. In some Arabidopsis accessions, dme is also not fully transmitted through the male parent (Schoft et al., 2011). DME is expressed in the polar nuclei and the central cell (the female gamete that is the progenitor of the endosperm) before fertilization and is required to establish gene imprinting in the endosperm after fertilization (Choi et al., 2002; Gehring et al., 2006). In the central cell, DME demethylates specific loci (Park et al., 2016) and this hypomethylated state is transmitted to the endosperm after fertilization, such that the maternally inherited endosperm genome is hypomethylated compared with the paternally inherited endosperm genome (Gehring et al., 2006, 2009; Hsieh et al., 2009; Ibarra et al., 2012). DME-dependent endosperm hypomethylated sites are enriched for fragments of transposable elements that reside near genes (Gehring et al., 2009). DME is also active in the pollen vegetative cell and similar targets are hypomethylated in a DME-dependent manner in both the vegetative cell and endosperm (Calarco et al., 2012; Ibarra et al., 2012). Like Arabidopsis, rice central cells and vegetative cells are hypomethylated (Kim et al., 2019; Park et al., 2016). Mutations in a rice DME homolog (termed ROS1a) have a similar phenotype as Arabidopsis dme mutants—maternal null alleles disrupt endosperm development and mutant maternal or paternal alleles are only rarely transmitted to progeny (Ono et al., 2012). ROS1a is also responsible for DNA demethylation in the vegetative cell (Kim et al., 2019) and likely also in the central cell. Although Arabidopsis DME is mostly highly expressed in reproductive tissues, expression is also detected in vegetative tissues (Mathieu et al., 2007; Park et al., 2017; Schumann et al., 2019). The function of DME outside of reproductive tissues is less well understood.

Arabidopsis plants with mutations in any of the other three DNA glycosylases lack visibly dramatic phenotypes under standard growth conditions and all null mutants are viable singly and in combination (Gong et al., 2002; Penterman et al., 2007; Ortega-Galisteo et al., 2008). However, some phenotypes have been observed. Stomatal precursor-cell density is increased in ros1 leaves due to hypermethylation and transcriptional silencing of a negative regulator of precursor cells (Yamamuro et al., 2014). Additionally, ros1 and rdd triple mutants exhibit impaired tracheary element differentiation, resulting in a high frequency of protoxylem discontinuities (Lin et al., 2020). Plants with mutations in ros1 and rdd exhibit enhanced susceptibility to bacterial and fungal pathogens (Yu et al., 2013; Le et al., 2014; López Sánchez et al., 2016), which is associated with decreased expression of biotic stress genes and, in a handful of examined genes, promoter hypermethylation at TE sequences (Le et al., 2014; Halter et al., 2021). Genome-wide profiling of DNA methylation in ros1 plants and rdd plants indicates DNA hypermethylation at hundreds or thousands of intergenic regions and transposable elements that are also targeted by RdDM and are enriched near genes (Penterman et al., 2007; Lister et al., 2008; Tang et al., 2016). Most of the DNA methylation changes in rdd are not obviously associated with changes in gene transcription (Penterman et al., 2007; Lister et al., 2008). Thus, it is thought that one function of the DNA demethylation in vegetative tissues is to “clean up” after a robust DNA methylation system to keep genes free from methylation.

Although genetic approaches have proved fruitful in understanding the function of 5-mC DNA glycosylases, the inability to generate homozygous mutant dme plants has stymied full understanding of this gene family. RNAi was used to knockdown DME expression in vegetative tissues of the rdd triple mutant (Schumann et al., 2019). These plants are even more susceptible to a fungal pathogen and additional hypermethylation was shown at a few loci. These data suggest that vegetative tissue of rdd plants retain some demethylation activity from DME. Recently, Kim et al. (2021) isolated homozygous mutant dme-2 plants in the Landsberg erecta (Ler) background, which display root and shoot defects related to meristem alterations. To identify the full extent of demethylation activity, we created plants where dme was complemented only in the central cell, allowing us to examine methylation and transcription on a genome-wide scale in vegetative tissues null for all four 5-methylcytosine DNA glycosylases.

### Results

**Isolating quadruple demethylase mutants**

In order to more fully understand the role of the DME family of 5-methylcytosine DNA glycosylases in Arabidopsis, we sought to isolate homozygous mutations in all four homologs in this family: **DME**, **ROS1**, **DML2**, and **DML3**. To rescue
the \textit{dme} seed abortion phenotype, we created a transgene in which the genomic coding sequence of \textit{DME} was expressed under the central cell-specific promoter of \textit{AGL61} (Figure 1, A; Steffen et al., 2008). The transgene was transformed into the previously isolated \textit{rdd} triple mutant in the Columbia-0 (Col-0) background (Penterman et al., 2007). Transgenic \textit{rdd} plants were pollinated by heterozygous \textit{dme} mutants to create heterozygous quadruple mutants (Figure 1, B). These heterozygotes were self-fertilized and seed abortion rates were quantified to test the ability of the \textit{pAGL61:DME} transgene to complement the \textit{dme} seed abortion phenotype. Whereas non-transgenic plants harboring the \textit{dme} mutation exhibit 50\% seed abortion, transgenic lines expressing \textit{pAGL61:DME} exhibited minimal seed abortion, indistinguishable from \textit{rdd} plants. Expression of \textit{DME} in the central cell before fertilization is therefore sufficient to rescue the post-fertilization seed abortion phenotype.

After self-fertilizing the heterozygous quadruple mutant, the following four genotypes were isolated over two subsequent generations: homozygous quadruple mutants (hereafter termed \textit{drdd}), homozygous \textit{rdd} mutants, homozygous \textit{dme} mutants, and homozygous wild-type segregants (hereafter termed WT, and to serve as genetically closely related wild-type controls for subsequent experiments). All four of these genotypes were determined to be homozygous for a single transgene insertion and displayed seed abortion rates of \(<2\%,\) similar to non-transgenic \textit{rdd} (Figure 1, C).

To determine whether the \textit{AGL61} promoter was active outside of the central cell, we compared the expression of \textit{DME} in leaves of wild-type Col-0 plants lacking the transgene to leaves of WT, \textit{dme}, and \textit{drdd} plants that were transgenic for \textit{pAGL61:DME} (Figure 1, D). There was no significant difference in \textit{DME} expression between Col-0 and WT, suggesting that \textit{pAGL61:DME} is not significantly active in leaves. \textit{DME} expression was 6.3- and 10-fold lower in \textit{dme} and \textit{drdd}, respectively (Figure 1, D), suggesting the \textit{AGL61} promoter can direct a low level of expression in leaf tissues. Thus, although perhaps not representing a complete loss-of-function of \textit{DME}, this material is suitable to examine the consequences of substantially reduced \textit{DME} activity in vegetative tissues.

**DME is required for full demethylation in somatic tissue**

To fully understand the function of \textit{DME} outside of the central cell and the combined activity of all members of the \textit{DME} family, paired DNA and RNA samples were collected from rosette leaf tissue of WT, \textit{dme}, \textit{rdd}, and \textit{drdd} plants. Individual leaves from four separate biological replicate plants were sliced along the midvein, and DNA and RNA were extracted from each half, so that DNA methylation and gene expression could be analyzed from the same leaf.

Whole-genome bisulfite sequencing (BS-seq) was performed on DNA samples (Supplemental Dataset S1). In order to improve read depth for subsequent analyses, bisulfite-sequencing reads were combined for pairs of replicates to create two high-depth replicates for each genotype. Previously, the \textit{rdd} triple mutant was constructed by back-crossing \textit{ros1} and \textit{dml2} T-DNA insertions originally from a Wassilewskija (Ws) background into a Col-0 background (Penterman et al., 2007). We therefore first analyzed the zygosity of Ws SNPs throughout the sequenced WT, \textit{dme}, \textit{rdd}, and \textit{drdd} methylomes to determine the presence of chromosomal regions originating from Ws. Consistent with previous reports (Penterman et al., 2007), we observed regions of Ws homoyzosity surrounding the \textit{ros1} and \textit{dml2} T-DNA insertions on chromosomes 2 and 3, respectively, in both

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**Figure 1** Generation of homozygous \textit{dme} and quadruple \textit{drdd} mutants. A, Schematic showing the construct designed to express \textit{DME} specifically in central cells using the \textit{AGL61} promoter. B, Schematic showing the segregation of mutant genotypes. Transgenic \textit{rdd} plants were pollinated by heterozygous \((+/-)\) \textit{dme} to generate heterozygous \textit{drdd} \textit{F}\textsubscript{1} progeny. WT, \textit{dme}, \textit{rdd}, and \textit{drdd} homozygotes were isolated from two subsequent generations. C, Proportion of aborted seeds in non-transgenic WT, \textit{dme}, \textit{rdd}, and \textit{drdd} plants homozygous for the \textit{pAGL61:DME} transgene. In total, 250–300 seeds were evaluated for each genotype. D, qRT-PCR measurement of \textit{DME} expression in WT Col-0 plants and WT, \textit{dme}, and \textit{drdd} plants expressing \textit{pAGL61:DME}. **\textit{P} \leq 0.001** (versus WT).
**rrd** and **drdd** (Supplemental Figure S1). These regions were discarded in all methylation analyses, as strain-specific methylation differences between Col-0 and Ws could contribute to false positives in identifying differentially methylated regions (DMRs).

To assess the extent of methylation differences in **dme**, **rrd**, and **drdd**, differentially methylated dinucleotides (CG context) or individual cytosines (CH context) were identified between WT and each mutant genotype. Methylation differences were restricted to a minimum percentage difference of 35% for CG, 20% for CHG, and 15% for CHH. Differentially methylated CGs/Cs (DMCs) were then aggregated into windows, which were assigned a differential methylation score based on the quantity and proportion of DMCs (Williams and Gehring, 2017). This score was then used to identify DMRs (Figure 2, A). Using this approach, very few DMRs were identified between biological replicates of the same genotype (33 between WT replicates and 5 between **drdd** replicates) and similar numbers of differences were identified between WT and mutant genotypes, regardless of which pairwise comparison of biological replicates was performed (Supplemental Figure S2). In total, 8–13% of DMRs identified with one biological replicate were not differentially methylated in other replicate comparisons, suggesting little difference between biological replicates. To account for these differences between replicates and identify a final list of DMRs between genotypes, only CGs/Cs that were differentially methylated in both mutant replicates were included (see methods). As this approach required DMCs to have adequate read depth in both replicates in order to be counted, we estimate that our analysis has few false positives, but as a consequence likely offers an underestimate of the true number of methylation differences among the genotypes.

We identified 276 DMRs between WT and **dme** mutants (Figure 2, B). This suggests that DME does not have a large number of target loci in leaves that are independent of the other three demethylases. We observed 2,091 regions with increased methylation in **rrd** compared with WT. The quadruple mutant **drdd** exhibited even greater methylation differences compared with WT, with 1,265 loci more than **rrd**, including almost twice the number of regions with differential non-CG methylation. This implies that DME does function in somatic tissue, but redundantly with **ROS1**, **DML2**, and **DML3**. The set of loci hypermethylated in **drdd** compared with WT (2,601 regions after merging sequence contexts) therefore represents a set of genomic targets for the entire DME family, which we hereafter refer to as “**DRDD target loci**” (Supplemental Dataset S2).

To better understand the extent of DME activity at DRDD target loci, we quantified methylation levels at these regions in all four genotypes studied. Single mutant **dme** replicates had methylation levels similar to WT, suggesting that DME functions redundantly with the other demethylases in removing methylation at these targets (Figure 2, C). Whereas **rrd** triple mutants showed hypermethylation in all three contexts at DRDD target loci, this hypermethylation was much more pronounced in **drdd** (Figure 2, C). At the single-cytosine level, 72% of CGs within DRDD target loci were more highly methylated in **drdd** compared with **rrd**, as well as 53% of non-CG cytosines. These data suggest that in **rrd** plants, DME still actively removes methylation from the majority of DRDD target loci. To further verify the 2,601 DRDD target loci identified, we performed a single replicate of enzymatic methyl-sequencing (EM-seq; Feng et al., 2020) on WT and **drdd** leaf tissues, which confirmed the increased methylation of all 2,601 loci in **drdd** (Figure 2, D).

We next sought to better understand the genomic context of DRDD target loci, as well as their characteristic DNA methylation patterns. To do this, we identified the chromatin state of each DRDD target using a published dataset of chromatin states that incorporates information from 16 characteristics and histone modifications (Sequeira-Mendes et al., 2014). The overall set of DRDD targets was dispersed over a distribution of chromatin states highly similar to the entire Arabidopsis genome (Figure 2, F). DRDD target loci could be classified in either of two groups: “CG-only” loci, which lack non-CG methylation in both WT and **drdd**, and exhibit increased CG methylation in **drdd**, and “multi-context” loci, which exhibit hypermethylation in CG and either CHG and/or CHH contexts in **drdd** (Figure 2, E). While equally common (Supplemental Dataset S2), these two types of target loci occupied distinct chromatin states. CG-only DMRs were more commonly associated with gene bodies and the immediately adjacent 5’- and 3’-regions, whereas multi-context loci were more common in distal 5’-regions and other intergenic DNA associated with the repressive histone mark H3K27me3 (Figure 2, F).

Intergenic DNA enriched for H3K27me3 is typically adjacent to heterochromatin and colocalizes with many RdDM targets in euchromatin as well as the borders of heavily methylated TEs (Sequeira-Mendes et al., 2014). These differences in chromatin state distribution likely reflect the well-characterized differences between the distributions of gene body methylation (gbM), which is typically only on CG-dinucleotides, and the activities of the RdDM pathway, which is active at repetitive DNA and intergenic sequences within euchromatin and the borders of TEs (Stroud et al., 2014; Bewick and Schmitz, 2017). Gene bodies have not been previously reported as targets for active DNA demethylation. We found that two-thirds of CG-only DMRs hypermethylated in **drdd** were also more methylated in **rrd** than WT, but failed to meet cutoffs for differential methylation (Supplemental Figure S3). We note that these CG-only DMRs are unlikely to represent the trivial differences in CG methylation that can spontaneously arise between generations, termed spontaneous epimutations (Johannes and Schmitz, 2019); our approach for identifying DMRs was designed to avoid identifying large numbers of epimutations, as evidenced by the small number of significant CG methylation differences identified between WT and **dme** (<100 CG DMRs) and between WT replicates (33 CG DMRs).
which shared the same grand-parental heritage as WT and *drdd*.

To further understand the role of active demethylation in targeting methylation within gene bodies, we sought to examine *drdd* methylation patterns within gene body methylated (gbM) genes (Zilberman et al., 2007; Bewick et al., 2016; Picard and Gehring, 2017). Across 4,234 Arabidopsis gbM genes identified by Bewick et al. (2016), we observed no differences in methylation between WT and *drdd* (Figure 3). We therefore sought to identify the genes directly overlapping CG-only hyperMRs, which we termed DRDD target genes. We identified 732 DRDD target genes, which minimally overlapped previously described gbM genes (175 out of 4,234). DRDD target genes display distinct characteristics in WT. Most notably, CGs were intermediately methylated, suggesting they are methylated in a subset of cells, and consequently display a lower average methylation level than typical gbM genes (Figure 3, B). Active demethylation by DRDD at these target genes appears to be biased toward the 5'- or 3'-ends of the coding sequence (Figure 3, B). We therefore propose that the gene bodies targeted by DRDD represent a new class of gbM genes that appear to be specifically targeted in a
subset of cell and tissue types, raising the possibility that gbM may play a functional role in some contexts.

**Demethylation by DRDD establishes tissue-specific methylation states**

DRDD target loci exhibit low but often non-zero levels of CG methylation in wild-type leaves and are by definition hypermethylated in *drdd*. This suggests that both methylation and demethylation processes occur at these sites in wild-type rosette leaves (Figures 2, C, 3, A). The low level of CG methylation at DRDD target loci in WT leaves is in contrast to most CG sites in the genome, which are either not methylated or fully methylated (Figure 3, A). We hypothesized that the balance between methylation and demethylation occurring at DRDD target loci could be altered in different tissues or over developmental time. To test this, we generated whole-genome bisulfite sequencing data for four tissues from individual wild-type Col-0 plants: rosette leaves (of the identical developmental stage as our prior leaf samples), cauline leaves, closed flower buds, and mature green embryos. Cauline leaves and flower buds were collected simultaneously. Two replicates were sequenced for each tissue type, with each set of tissues collected from one individual plant.

We identified hundreds of regions differentially methylated at CG dinucleotides among tissues and thousands of regions differentially methylated in CHG or CHH contexts (Supplemental Figure S4). Embryos exhibited 2,612 regions with increased CHH methylation when compared with rosette leaves and 2,189 regions when compared with cauline leaves, suggesting increased activity of the RdDM or CMT2 pathways at this developmental stage, consistent with previous reports documenting embryo hypermethylatation (Bouyer et al., 2017; Kawakatsu et al., 2017; Narsai et al., 2017). In addition, we observed increased CHG methylation at 11,588 regions in embryos or flower buds compared with both rosette and cauline leaves. CHH and CHG methylation sites typically exhibit partial methylation, with only a fraction of sites methylated in whole-tissue bisulfite sequencing data, suggesting cellular heterogeneity. It is therefore possible that differences in the cell type composition of different tissues could cause large differences in the levels of non-CG methylation observed in whole tissues. Rosette and cauline leaves exhibited highly similar methylation profiles in our dataset, suggesting that cell-type composition is predictor of methylation patterns.

To evaluate the relationship between tissue-specific methylation differences and DRDD, we calculated the methylation levels of DRDD target loci in each tissue of our dataset, as well as a set of randomly selected 200 bp windows that had the characteristics necessary to be identified as a hypermethylated region (hyperMR) (see Methods). DRDD target loci exhibited increased methylation in flower buds and embryos compared with the leaf samples, most obviously at regions with increased CG methylation in *drdd* (Figure 4, A and B and Supplemental Figure S5). To control for genomewide differences in the activity of the RdDM pathway between different tissue types (Bouyer et al., 2017), we also examined the methylation level of DRDD targets in all four tissues of an *rdr2* mutant. RDR2 is involved in biogenesis of
the small RNAs that direct DNA methylation. We found that many DRDD targets exhibited tissue-specific methylation differences in the absence of RdDM activity (Figure 4, A and Supplemental Figure S5). We also calculated the standard deviation in DNA methylation among tissues at DRDD target loci and control regions in both WT and rdr2. The standard deviation of methylation levels between tissues was much higher at DRDD target loci than at randomly selected windows (Figure 4, C), suggesting that DRDD targets overlap with loci that exhibit developmentally dynamic epigenetic states, in many cases independent of RdDM activity. Lastly, >50% of regions identified to be CG hypomethylated in either rosette or cauline leaves compared with buds or embryos overlapped directly with DRDD target loci (Supplemental Figure S4). Thus, hundreds of loci exhibited high methylation levels in WT buds and embryos but reduced methylation in cauline and rosette leaves, as well as high methylation levels in drdd rosette leaves (Figure 4, B

Figure 4 Demethylation by DRDD establishes tissue-specific methylation differences. A, CG methylation levels of four different tissues of WT and rdr2 mutants at 2,601 DRDD target loci. **P < 0.0001 (t-test, compared with rosette leaves). The black dots represent the median value and vertical lines denote the interquartile range. B, Heatmap showing methylation level of DRDD target loci across four different WT tissues. Methylation levels of rosette leaves from WT and drdd + pAGL61:DME are shown for comparison. C, Standard deviation of DNA methylation levels across tissues at DRDD target loci compared with random windows. D, Genome browser snapshots depicting typical examples of multicontext and CG-only DRDD targets in different WT tissues.
We observed this DRDD-dependent demethylation in WT leaves at both CG-only and multi-context DRDD targets (Figure 4, D). The activity of DRDD at multi-context targets likely opposes the RdDM pathway, as these targets exhibited reduced variability between tissues in rdr2 mutants, whereas CG-only targets did not (Supplemental Figure S5).

Demethylation by DRDD is associated with target gene regulation

DNA methylation has variable impacts on gene expression. While many genes are not affected by changes to DNA methylation at proximal sequences, a subset of genes are strongly impacted by DNA methylation, often by transcriptional silencing and/or heterochromatin formation (Lister et al., 2008), although occasionally by promoting increased expression (Williams et al., 2015; Pignatta et al., 2018). To identify which genes might be impacted by the activity of DRDD demethylases, we performed RNA-seq on WT, dme, rdd, and drdd tissue isolated from the same leaf samples used for methylation profiling (Supplemental Dataset S1). WT and dme transcriptomes were almost identical (Figure 5, A), consistent with the minimal differences observed in methylation patterns between the two genotypes. We identified 168 downregulated and 174 upregulated genes in rdd...
compared with WT, and 170 downregulated and 252 upregulated genes in *drdd* compared with WT. Overall, the *drdd* transcriptome encompassed a larger dynamic range of expression changes relative to WT, suggesting that the most extreme expression changes were exacerbated in *drdd* compared with *rdd* (Figure 5, A, Supplemental Figure S6, and Supplemental Dataset S3).

To examine whether these expression changes were linked to DNA methylation differences between *drdd* and WT, we calculated the distance between up and down-regulated genes and the closest DMR. Regions of increased DNA methylation (hyperMRs) were frequently adjacent to down-regulated genes, with 40% (45 out of 113 genes, *P* < 0.0001 (Ws regions excluded)) of down-regulated genes residing within 2 kb of a hyperMR (Figure 5, B). This association is consistent with previous studies of *ros1* and *rdd* mutants, which suggested that a small subset of methylation gains in these mutants caused transcriptional silencing of proximal genes (Lister et al., 2008; Yamamuro et al., 2014; Halter et al., 2021). We also observed proximity between hyperMRs and some up-regulated genes, with 17% of up-regulated genes residing <2 kb of a hyperMR (34 out of 205 genes, *P* = 0.03). This association is marginally enriched compared with the number of genes expected by chance, as determined by calculating the distance between hyperMRs and control groups of randomly selected 200 bp windows (Figure 5, B).

The proximity between a subset of hyperMRs and down-regulated genes was detectable for both multi-context (30 genes) and CG-only (19 genes) hyperMRs (Figure 5). An association between down-regulated genes and multi-context hyperMRs was expected, as methylation in multiple sequence contexts is a hallmark of the RdDM pathway, which typically functions in transcriptional silencing. The association of CG-only DMRs with decreased expression in *drdd* was more surprising, as CG gbM has mostly been associated with minimal impacts on gene regulation in plants (Bewick and Schmitz, 2017; Picard and Gehring, 2017). For example, the locus encoding the defense and cell death regulator *SPEECHLESS* (Figure S7). Conversely, we also observed that increased *SPEECHLESS* expression in *drdd* was associated with increased CG gbM (Figure 5, D–E).

To further understand the link between expression and gbM, we identified 13 genes that were differentially expressed between WT and *drdd* and directly overlap CG-only hyperMRs. We then analyzed their tissue-specific methylation and expression levels (as defined by Klepikova et al., 2016), to see if developmentally modulated methylation patterns could be associated with gene regulation. In all down-regulated genes, there was a strong inverse correlation between expression and methylation level across tissues (Supplemental Figure S8). Additionally, four out of six upregulated genes exhibited strong positive correlations between expression and methylation level, suggesting that CG gbM can be associated with up-regulation. DRDD may actively remove methylation from these coding regions to facilitate higher or lower expression, or the altered transcriptional output of these genes may induce a different epigenetic state in a DRDD-dependent manner (Figure 5, D and E).

To better understand if the gene expression differences between WT and *drdd* mutants are associated with any particular biological functions, we performed GO term analysis (Supplemental Figure S6). Genes with decreased expression in *drdd* were significantly associated with one GO term, immune response, which was also associated with genes that increased in expression (Supplemental Figure S6). Several defense-response genes down-regulated in *drdd* were previously identified as targets of ROS1 and are in transposon-rich rapidly evolving regions of chromosomes, which tend to be frequently targeted by the RdDM pathway (Le et al., 2014; Halter et al., 2021). We observed greater increases in the expression of epidermal development genes in *drdd* than in *rdd* (Supplemental Figure S6). GO terms for regulation of hormone levels, response to osmotic stress, and response to cold were also significantly enriched among upregulated genes (Supplemental Figure S6).

**DRDD enzymes remove methylation from the FLOWERING LOCUS C (FLC) locus and are required to delay flowering**

During long-day (16 h) growth conditions, we observed that *drdd* mutants were early flowering compared with WT, *dme*, and *rdd* plants, transitioning to flowering after establishing eight-nine true leaves (Figure 6, A, Supplemental Figure S9, and Supplemental Dataset S4). The early flowering phenotype observed in *drdd* is unlikely to be caused by the presence of Ws regions, as these regions are also present in *rdd* (Supplemental Figure S1). Under short day (8 h) growth conditions, we observed an early flowering phenotype both in *drdd* and *rdd*, which has not been noted in previous studies, although the effect was greater for *drdd* (Figure 6, B, Supplemental Figure S9, and Supplemental Dataset S4). We therefore hypothesized that active DNA demethylation in Arabidopsis might regulate flowering time, perhaps by altering the epigenetic state at genes important for regulating the transition to flowering. In our RNA-seq dataset, the flowering time regulator *FLC* had significantly reduced expression in comparisons of WT versus *drdd* and *rdd* versus *dme*. In fact, *FLC* transcripts were not detected in *rdd* mutants (Figure 6, C). We then sought to verify the expression levels of *FLC* using reverse transcription-quantitative PCR (RT-qPCR). *FLC* expression was clearly reduced in both *rdd* and *drdd* mutant seedlings compared with WT, with a 3-fold and 5-fold lower relative transcript abundance, respectively (Figure 5, D). Complex epigenetic regulation of the *FLC* locus by histone modification pathways has been well-described (Wu et al., 2020), but whether it is connected to DNA methylation in cis remained unknown.

To assess the DNA methylation state of the *FLC* locus, we analyzed the DNA methylation profiles of WT, *dme*, *rdd*, and
plants in our methylation profiling data. We observed minimal DNA methylation in the transcribed region or 3' region of FLC, but saw evidence of DNA methylation in all three sequence contexts in a region 800–2,139 bp 5' of the transcriptional start site of the FLC coding region. To better define the methylation patterns of this region with greater depth, we performed enzymatic-methyl sequencing (EM-seq) on WT and drdd leaves. The EM-seq data clearly showed large increases in methylation in all sequence contexts across this region (Figure 6, E). We further verified this methylation change by performing bisulfite PCR on three separate fragments within this region (Supplemental Figures S10, S11). Across these bisulfite PCRs, we observed no methylation in WT or dme plants, but increased methylation in drdd, and to a lesser extent rdd, consistent with the observed flowering time phenotypes (Figure 5, B and E). Thus, DRDD enzymes target the region 5' of FLC for demethylation. Hypermethylation of this region is correlated with reduced FLC expression, as well as earlier flowering time.

As this region is highly methylated in all three sequence contexts in drdd, we hypothesize that it is targeted by RdDM. Indeed, 23–24 nucleotide small RNAs homologous to this precise region, which could participate in RdDM, are present within multiple WT small RNA datasets (Lee et al., 2012; Jeong et al., 2013; Erdmann et al., 2017). Additionally, we also identified a pair of 100 bp tandem repeats with high sequence similarity within the DRDD target region. Tandem repeats have been shown to recruit the RdDM pathway upstream of other genes, including the flowering time regulator FWA (Chan et al., 2006; Henderson and Jacobsen, 2008). Thus, FLC is an exemplar of a locus that lacks DNA methylation in WT leaves due to active DNA demethylation, rather than due to lack of methylation targeting.

To further investigate the link between DNA methylation 5' of FLC and flowering time, we examined methylation patterns of the FLC 5' region across 653 ecotypes for which methylation coverage was available (Kawakatsu et al., 2016). We identified a short sequence upstream of the tandem

**Figure 6** DRDD enzymes delay flowering and remove DNA methylation proximal to FLOWERING LOCUS C. A, Flowering phenotype of long-day grown WT, dme, rdd, and drdd plants 24 days after germination. B, Flowering time of WT, dme, rdd, and drdd plants grown in long (16 h) and short (8 h) day conditions. Flowering time is measured by the number of true leaves present at the initial time of bolting. Error bars represent standard deviation between biological replicates. C, Differentially expressed genes in drdd compared with WT, with FLC highlighted among strongly down-regulated genes. D, Relative FLC expression in 7-day-old seedlings measured by quantitative real-time PCR. E, Genome browser snapshot of EM-seq methylation data and Col-0 embryo small RNA-seq data (Erdmann et al., 2017) at the intergenic region 5' of FLC.
repeats targeted by RdDM/DRDD that exhibited highly variable methylation levels across ecotypes, which we term the ecotype variable region (region Y, Figure 7, D). Conversely, we found that the tandem repeats were seldom methylated, exhibiting methylation levels comparable to $drdd$ in only 14 ecotypes (region X, Figure 7, D). If DNA methylation of the FLC tandem repeats reduces flowering time as we hypothesize, then we would predict these 14 ecotypes to exhibit reduced flowering time compared with closely related ecotypes. For 11 out of 14 these ecotypes (for which both DNA methylation and flowering time data were available), we identified the most closely related ecotype that lacked methylation at the FLC tandem repeats, using the 1001 genomes project SNP identity tool (tools.1001genomes.org/strain_id/). In all cases, the genomic SNP identity of related ecotypes was >99%. This provided a dataset of genetically similar ecotype pairs with large differences in FLC methylation levels (Figure 7, A). While the 11 highly methylated ecotypes exhibited variable flowering times—likely reflecting a wide variation in geography, reproductive strategy, and developmental phenotypes—overall they exhibited a significantly reduced flowering time compared with closely related and lowly methylated strains (Figure 7, B and C). This data are therefore consistent with the hypothesis that high levels of methylation directed to the tandem repeats upstream of FLC is associated with a reduced flowering time, and may point to a role for DNA (de)methylation in modulating the flowering time of plants in the wild.

**Discussion**

In this study, we sought to elucidate the role of the DME family of 5-mC DNA glycosylases in somatic tissues of A. thaliana. To this end, we generated somatic homozygous mutants of $dme$ as well as quadruple $drdd$ mutants. $dme$ mutants exhibited similar phenotypes, DNA methylation profiles, and transcriptional profiles to WT plants, suggesting that DME does not perform a distinct individual function within leaf tissue, with the caveat that our mutants are likely not a true $dme$ null due to possible low-level expression of the pAGL61:DME transgene outside of the central cell. Nevertheless, when combined with mutations to the other three members of the DME-family to generate $drdd$ mutants, a number of previously unknown targets and functions for active demethylation in somatic tissues were brought to light.
Active DNA demethylation targets a subset of gene bodies

Compared with WT, drdd mutants exhibited DNA methylation gains at 2,601 loci. We estimate the true number of DRDD targets to be even higher, as we employed stringent cutoffs for both read depth and the effect size of methylation changes. Approximately half of the DRDD targets gained methylation in all three sequence contexts, likely due to DNA methylation deposited by the RdDM pathway. These targets are similar in manner to previously described RDD targets—loci proximal to TEs or intergenic sequences that are targeted by RdDM, and exhibit an increase or "spreading" of DNA methylation in the absence of demethylase activity (Penterman et al., 2007; Lister et al., 2008; Tang et al., 2016).

However, we also identified approximately 1,000 targets unlike those previously described in demethylation mutants, in which DNA methylation present in the CG-only context was removed. These regions are typically located within open chromatin, such as introns, exons, or untranscribed regions and are referred to as gbM. However, the sites we describe here are distinct from typical gbM sites, in which CG methylation at individual sites is close to 100% (Picard and Gehring, 2017). In WT plants, the CG-only regions we identified exhibit intermediate methylation levels, in which CG dinucleotides are methylated at 5–50% (Figure 3). This low to intermediate methylation level implies that the CG sites are methylated in only a fraction of cells in the tissue. Although many of the CG-only target loci exhibit low methylation levels in WT leaves, they are more highly methylated in buds and embryos (Figure 4 and Supplemental Figure S5). Importantly, these loci are also highly methylated in drdd leaves, suggesting their methylation state is established by the active removal of methylation by DRDD enzymes in a subset of leaf cells. The function, if any, of gbM has been enigmatic. Globally, gbM is associated with moderately expressed genes (Zilberman et al., 2007), but some plant species have dispensed with this aspect of methylation patterning altogether (Bewick et al., 2016).

Nevertheless, we observed 11% of down-regulated genes contained a CG-only DRDD target within their gene body, suggesting that methylation at these loci may indeed impact gene expression. These genes also exhibited a strong inverse correlation between CG methylation level and expression level across tissues (Supplemental Figure S8). Thus, we propose that active demethylation may act in a specific subset of cell types, cell cycle, or endoreduplication states in WT tissues, and that the intermediate methylation levels in WT plants are a consequence of heterogeneity in methylation state between cells. The absence of this heterogeneity within drdd mutants could lead to interesting future research on the establishment of methylation patterns within specific cell types.

The discovery of CG-only DRDD targets within gene bodies also raises questions about possible targeting mechanisms for DRDD demethylases. Previous work has proposed that active demethylation by ROS1 may be targeted by specific histone modifications associated with transcriptionally repressed intergenic sequences in Arabidopsis, such as H3K18ac and H3K27me3 (Qian et al., 2014; Tang et al., 2016). Our results are not fully consistent with this view, as we observed DRDD targets in transcriptionally active genes, and across a broad range of chromatin states (Figure 2, F) not dissimilar to the overall distribution of chromatin states in the Arabidopsis genome. It is possible that functional diversification between ROS1 and DME may contribute to such differences in their individual targeting.

Active DNA demethylation targets FLOWERING LOCUS C and is associated with reduced flowering time

FLC is a key regulator of the transition to flowering in Arabidopsis. The transcriptional silencing of FLC under prolonged exposure to cold mediates the vernalization response in some accessions via a Polycomb repressive complex switch (Costa and Dean, 2019). In addition, FLC also regulates the autonomous flowering pathway in early-flowering strains of Arabidopsis, such as Col-0 (Wu et al., 2020). FLC is expressed in the shoots of Col-0 plants in the initial weeks after germination, but expression decreases immediately prior to flowering (Pien et al., 2008; Choi et al., 2009). FLC has been established as a model locus for epigenetic and transcriptional control, and is beginning to be understood with a high degree of complexity (Wu et al., 2020). Despite extensive previous research, connections between FLC regulation and DNA methylation have been limited and indirect. While changes to FLC transcript abundance have been reported in some DNA methyltransferase and DNA methylation-binding mutants (Sheldon et al., 1999; Finnegan et al., 2005; Peng et al., 2006; Yaish et al., 2009), targeting of the FLC locus by either methylation or demethylation pathways has not previously been reported.

FLC is one of the most highly down-regulated genes identified by RNA-seq in drdd compared with WT. This expression change was consistent with the early flowering phenotype of drdd mutants in both long and short day photoperiods. We also observed a difference in methylation patterning proximal to the FLC locus that is correlated with the observed transcript abundance (Figure 6, E). It is possible that DNA methylation at this region could interfere with the binding of key regulators of FLC transcription. The FLC regulator FRIGIDA has been shown to form a super-complex with the H3K4 methyltransferase complex COMPASS-like at FLC (Li et al., 2018). Binding of this complex has been detected at a broad region encompassing 500 bp on either side of the FLC transcriptional start site. While this region does not directly overlap the sequences hypermethylated in rdrd and drdd mutants, or 14 ecotypes that exhibit high methylation levels at this region (Figure 7), it is possible that demethylation of this region is required to establish the chromatin landscape necessary to facilitate binding of this super-complex.

As FLC appears to be targeted by DRDD, yet is completely unmethylated in most WT strains, an interesting question is whether this intergenic region is co-targeted by methylation
and demethylation pathways in WT leaves, resulting in a constant battle between the addition and removal of DNA methylation. Active DNA demethylation at this locus may maintain epigenetic homeostasis (Williams and Gehring, 2020) by preventing the over-reach of methylation-targeting pathways, or methylation of FLC might be important for modulating expression at some point in the life cycle. Small RNAs with homology to the 5′-intergenic region of FLC (Lee et al., 2012; Jeong et al., 2013; Erdmann et al., 2017) could drive the activity of the RdDM pathway, only for methylated cytosines to be subsequently removed by DRDD. Furthermore, FLC is an imprinted gene in the endosperm of some Arabidopsis accessions (Pignatta et al., 2014), a phenomenon that has been observed at other loci with methylated 5′ regulatory intergenic regions. Understanding why dynamic methylation and demethylation pathways appear to be engaged in a tug-of-war at this locus will be an exciting avenue for future research.

In summary, we propose that active demethylation in somatic tissues by DRDD plays an important role in maintaining epigenetic states that can influence transcriptional activity. This activity appears to be important in protecting the chromatin landscape at FLC, a locus at which a number of dynamic epigenetic mechanisms converge. Similar to reproductive development, active demethylation by DRDD may also be developmentally regulated, and may act as a mechanism to establish divergent epigenetic states between cell and tissue types.

Materials and methods

Plant material

Triple homozygous mutant A. thaliana ros-1-3; dml2-1; dml3-1 plants in the Col-0 background (Penterman et al., 2007) were transformed with pAGL61:DME via floral dipping (Clough and Bent, 1998). Twelve single-insertion transformants were selected and pollinated with dme-2 (Col-0 background) (Choi et al., 2002) heterozygote mutant pollen to generate F1 progeny heterozygous for all four DRDD demethylase genes. Two of these T1 lines exhibited a complete rescue of seed abortion. Quadraple heterozygous plants from one of these lines were self-fertilized, and over two subsequent generations of segregation the following genotypes were isolated, each homozygous for the pAGL61:DME transgene: dme, rdd, rdr2, and DRDD WT segregants (to serve as a closely related WT control). The selfed progeny of the initial plant of each genotype was used for all subsequent experiments. To assess seed abortion, siliques were harvested after drying and seeds examined under a dissecting microscope. For tissue methylation profiling, WT plants were segregated from an rdr2-1 ros-1-7 double heterozygote, as described in Williams and Gehring (2017). rdr2 plants were homozygous for the rdr2-1 allele.

pAGL61::DME transgene

The transgene to rescue DME expression in central cells was created by amplifying the promoter of AGAMOUS-LIKE 61 (AGL61), using the primers originally designed by Bemer et al. (2008), (F primer: CAACCGATTGTCAAAATGCC GAAACCCA, R primer: TTTTTGTAGGAGGTTTAG TTGCTTTTCT), the full genomic coding sequence (introns included) of DME (F primer: ATGAATTGAGGG CTGATCCG, R primer: TTAGGTGTTGTCTCTTCAATTT GCT), and cloning both fragments into pENTR-TOPO-D via Gibson assembly (overhang sequences are not included in the primers above). The assembled pAGL61:DME construct was then transferred to the binary vector pMDC99 (Curtis and Grossniklaus, 2003) using LR clonase.

Flowering time assay

Plants were sown such that every row of the flats contained one plant of each genotype (WT, dme, rdd, and drdd) with the order iterating by one with each successive row. Flats were grown in a Conviron CMP6050 Control System at 22°C and 50% relative humidity, with 16 h of 120 μmol light from Philips Master TL-D 58W/840 bulbs and 8 h of darkness per day. For short day experiments, the growth cycle was 8 h of light and 16 h of darkness. Starting at 2 weeks of age, all plants were visually inspected three times per week. The number of rosette leaves was recorded once a bolt was visible. Populations were compared using a one-way ANOVA with post hoc Tukey’s test (Supplemental Dataset S4). The long day flowering time assay was repeated three times independently and the short-day assay was repeated twice, using >30 plants of each genotype for each experiment.

Bisulfite sequencing

Four replicate samples of the fifth true leaf of 3-week-old WT, dme, rdd, and drdd individual plants were collected (16 plants total). Leaf samples were split along the midvein, with one half each used for DNA and RNA extractions. DNA was extracted using a CTAB protocol and bisulfite conversion was performed on 200 ng DNA using an Invitrogen MethylCode bisulfite conversion kit. Bisulfite-sequencing libraries were then generated using a QIAGEN Qiaseq Methyl Library Kit, with 11 cycles of amplification and each individual replicate separately indexed. Samples were then sequenced using an Illumina HiSeq 2500 rapid run using 100 × 100 bp paired-end protocol at the Whitehead Institute Genome Technology Core. All 16 samples were multiplexed equally in two separate lanes, to avoid batch effects in sequencing. To improve genomic coverage for downstream analyses, the reads for every two replicates were pooled, creating two high-depth biological replicates for each genotype. To sequence different tissue types, two WT Col-0 replicates and a single rdr2 replicate were sampled throughout each individual plant’s development. The fifth true leaf was collected on day 21; the second cauline leaf and closed flower buds from the primary inflorescence were collected on day 35; and a pool of 30 mature green embryos were collected on day 50. DNA extraction, bisulfite conversion, library preparation, and sequencing were conducted as above.
with the following modifications: (1) embryo samples yielded less than 200 ng of DNA, so reduced quantities were used for each replicate; (2) libraries were amplified with 6–10 cycles; and (3) samples were equally multiplexed across three lanes and sequenced using an Illumina HiSeq2500 standard run with 100 × 100 bp paired-end reads at the Whitehead Institute Genome Technology Core.

DNA methylation analysis for bisulfite-sequencing data

Prior to mapping, adapters were trimmed using Trim Galore (Babraham Bioinformatics), trimming 8 or 10 bp from the 5'-end of reads, and enforcing a 3'-end quality of >25%. Reads were mapped to the Arabidopsis genome using Bismark 0.20.1 (Krueger and Andrews, 2011), allowing for 1 mismatch per read and removing PCR duplicates. Methylation values for each cytosine were calculated using the Bismark methylation extractor function. The efficiency of bisulfite conversion was verified by quantifying the percentage of methylation for reads mapped to the chloroplast. Bisulfite conversion rates were >99.7% across all samples. Before identifying DMRs, SNPs homologous to the Ws-2 genome were identified as described (Picard and Gehring, 2017), and their zygosity was plotted across each chromosome (Supplemental Figure S1). This is because the ros1 and dml2 T-DNA insertions in rdd were originally introgressed from a Ws background. In order to exclude genomic regions (and associated methylation differences) originating from the Ws ecotype, two regions were excluded from all subsequent methylation analyses: chromosome 2 (8,802,496–15,397,296 bp) and chromosome 3 (677,340–5,117,803 bp).

To identify DMRs, the genome was divided into 200 bp windows overlapping by 100 bp. Symmetrical cytosines within CG base pairs were combined to make a single averaged data point, as the two opposite-stranded cytosines within CG base pairs are not statistically independent. A “methylation score” was then calculated for each window based on the density of differentially methylated cytosines (DMCs) within each window (Williams and Gehring, 2017). Each hypermethylated cytosine/CG was assigned a score of +1 and each hypomethylated cytosine/CG was assigned a score of −1, with the added stringency that DMCs had to be present in both biological replicates for each sample. The minimum methylation difference for each cytosine context was as follows: CG—35%, CHG—20%, and CHH 15%. Cytosines with fewer than five reads coverage in each sample being compared were excluded from the analysis. For each window, the number of DMCs were then divided by the total number of cytosines with adequate coverage. A higher score is therefore assigned to windows in which hypermethylated DMCs are proximal to each other, whereas a lower score is assigned to windows in which hypomethylated DMCs are proximal to each other. Windows with a score greater than (not including) +2 were called hyperMRs, windows with a score less than (not including) −2 were called hypoMRs. DRDD targets were identified by combining all windows hypermethylated in drdd compared with WT in any sequence context and merging together using bedtools merge. DRDD targets were split into two categories based on their methylation profiles. CG-only targets were identified based on the absence of non-CG methylation (both CHG and CHH <5%) in drdd, whereas multi-context targets were defined by possessing methylation in both CG and at least one other sequence context (>5% CHG and/or CHH). The presence of DRDD target loci in each chromatin state of the genome was determined by intersecting DRDD targets with the chromatin states identified by Sequeira-Mendes et al. (2014). Chromatin states 4 and 5 were combined to represent “Intergenic polycomb” and chromatin states 8 and 9 were combined to represent “heterochromatin”.

Random control windows were selected from windows with five or more reads coverage for each cytosine and methylation levels low enough in WT to be detected as hypermethylated in drdd (e.g. <65% CG methylation, <80% CHG methylation, and <85% CHH methylation). Random control windows for comparison against multi-context DRDD targets were required to have <65% CG methylation and either <80% CHG methylation or <85% CHH methylation in WT.

Enzymatic methyl sequencing

Single biological replicates of the fifth true leaf of 3-week-old WT and drdd individuals were collected. DNA was extracted using a CTAB protocol and enzymatic methyl sequencing (EM-Seq) was carried out using 100 ng of input DNA. Input DNA was first sheared in 1 × TE buffer using a Covaris S220 to an average fragment size of 200 bp (175W, 10% duty factor, 200 cycles/burst, 180 s). After shearing, a 2.5 × bead cleanup was performed with a final elution in water. EM-seq was conducted using the NEBNext Enzymatic Methyl-Seq Kit according to the manufacturer’s directions; including the provided spike-in controls (methylated and unmethylated DNA). Six cycles of PCR amplification were performed. Samples were equally multiplexed across three lanes and sequenced using an Illumina HiSeq2500 standard run with 100 bp paired-end reads at the Whitehead Institute Genome Technology Core.

DNA methylation analysis for EM-Seq data

Adapter trimming and read mapping were performed as described above for the bisulfite sequencing data, except that spike-in DNA sequences were included in the reference genome. Methylation values for each cytosine were also calculated as above. The efficiency of enzymatic conversion was determined by quantifying the percent of methylation for reads mapping to the spike-in methylated and unmethylated DNA sequences.

RNA-seq

RNAs were isolated from 3-week-old leaf samples (explained above) using a QIAGEN RNeasy plant mini kit. In total, 400 ng total RNA was used to generate RNA-seq libraries using
a QIAGEN QIAseq Stranded mRNA Select Kit, with 13 cycles of amplification. An additional round of purification using QIAseq beads was performed to remove adapter dimers. RNA-seq was performed on an Illumina HiSeq 2500 using a 50-bp single-end protocol at the Whitehead Institute Genome Technology Core. All samples were multiplexed equally in two separate lanes to avoid batch effects in sequencing.

**Gene expression analysis**

Prior to mapping, adapters were trimmed using Trim Galore (Babraham Bioinformatics), trimming 9 bp from the 5'-end of reads, and enforcing a 3'-end quality of >25%. Reads were mapped to the Arabidopsis genome using STAR, permitting 0.05 mismatches as a fraction of total read length and discarding reads that did not map uniquely (Dobin et al., 2013). Differentially expressed genes were identified by running htseq-count and DESeq2 (Love et al., 2014), ensuring a 0.05 mismatches as a fraction of total read length and discarding reads that did not map uniquely (Dobin et al., 2013). Differentially expressed genes were identified by running htseq-count and DESeq2 (Love et al., 2014), ensuring a minimum of two-fold change in expression and a Benjamini–Hochberg corrected P-value < 0.05. Proximity between differentially expressed genes and DMRs was calculated using BEDTools (closest). As a negative control, the proximity between differentially expressed genes and 10 sets of randomly selected 200 bp windows was calculated for each comparison. For example, proximity to 2,601 DRDD target DMRs was compared with proximity to 2,601 random windows for which hypermethylation could have been detected (e.g. <65% CG, <80% CHG, and <85% CHH methylation in WT). Genes mapping to chromosomal regions with homology to Ws were omitted from the differential expression and DMR proximity analyses shown in Figure 4. GO term analysis was performed using DAVID (Dennis et al., 2003).

**Locus-specific bisulfite-PCR**

DNA was extracted using a CTAB protocol, and bisulfite conversion was performed on 200 ng DNA using an Invitrogen MethylCode bisulfite conversion kit. Bisulfite PCR was performed using a hot-start Taq polymerase [either Ex-Taq HS (Clontech) or DreamTaq HS (Thermo Scientific)] with an annealing temperature of 50°C and an extension temperature of 68°C. PCR products were purified with a MinElute Gel Extraction Kit (QIAGEN) and cloned into the pJET1.2/blunt vector (CloneJET PCR Cloning Kit; Thermo Scientific). Individual colonies (15–25 per locus) were either PCR screened with pJET F and R primers, or plasmids were extracted with a QIAprep Spin Plasmid Miniprep Kit (QIAGEN). PCR screen amplicons or plasmids were sequenced by Sanger sequencing using the pJET1-2F universal primer. Sequences were checked manually for quality, and vector and primer sequences were removed with SnapGene (VS.2.4). Sequences were aligned with Clustal Omega (Sievers et al., 2011) and methylation state was analyzed using CYMATE (Hetzl et al., 2007). Primers for locus-specific bisulfite PCR are in Supplemental Table S1.

**Reverse transcription-quantitative PCR**

RNA was isolated from the 5th leaf of 21-day-old plants using TRIzol Reagent (Invitrogen) according to manufacturer’s instructions. Genomic DNA was removed by treatment with Amplification-grade DNase I (Invitrogen). cDNA was prepared from 500 to 750 ng RNA (standardized within each batch) with Superscript II Reverse Transcriptase (Invitrogen) according to the manufacturer’s instructions, with polyadenylated transcripts selected for through use of an oligo-dT primer. qPCR was performed on a StepONE Plus Real-Time PCR system with Fast SYBR-Green PCR master mix (Applied Biosystems). DME primers were F: CGGGACGGGCT GATTCCTTCAT R: TCCATGGCGAAAAACGTCTATCTC. FLC primer sequences were as previously described (Csorba et al., 2014): FLC_F: AGCCAAAGAGCGAACTCA, and FLC_R: TTTGTCCAGCAGTGACATC. Reactions were normalized to reference gene AT1G58050 (Czechowski et al., 2005). All qPCR reactions were performed with technical triplicates. Cycling conditions were as follows: 95°C for 20 s followed by 40 cycles of 95°C for 3 s and 60°C for 30 s. Relative fold change in expression was determined using the ∆∆Ct method (Livak and Schmittgen, 2001).

**Accession numbers**

All high-throughput sequencing data are available in NCBI GEO under accession GSE191307.

**Supplemental material**

- **Supplemental Figure S1.** Zygosity of Ws SNPs in rdd and drdd mutants.
- **Supplemental Figure S2.** Identification of hyperMRs across individual replicates.
- **Supplemental Figure S3.** Methylation of individual CGs within hyperMRs of rdd and drdd mutants.
- **Supplemental Figure S4.** DNA methylation differences between tissue types.
- **Supplemental Figure S5.** Methylation levels of DRDD targets across tissues in WT and rdr2.
- **Supplemental Figure S6.** Significantly enriched GO terms in drdd versus WT.
- **Supplemental Figure S7.** Bisulfite PCR validation of CG-only DRDD targets.
- **Supplemental Figure S8.** Correlation between expression and CG methylation across tissues.
- **Supplemental Figure S9.** Additional flowering time assay replicates.
- **Supplemental Figure S10.** Bisulfite PCR analysis of FLC in WT and demethylation mutants.
- **Supplemental Figure S11.** Individual cytosine methylation data for FLC.
- **Supplemental Table S1.** Primers for locus-specific bisulfite PCR.
- **Supplemental Dataset S1.** Properties of sequencing libraries generated in this study.
- **Supplemental Dataset S2.** List of DRDD target regions.
**Supplemental Dataset S3.** Differentially expressed genes identified by DESeq2.

**Supplemental Dataset S4.** ANOVA of flowering time data.

**Acknowledgments**

We are grateful to Enrico Calvanese for performing bisulfite PCR validation experiments and Maria José Aranzana for performing a flowering time assay replicate.

**Funding**

Research in this publication was supported by the National Institute of General Medical Sciences of the National Institutes of Health under award R01GM112851 to M.G.

**Conflict of interest statement.** None declared.

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