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DOMAINS REARRANGED METHYLTRANSFERASE3 controls DNA methylation and regulates RNA polymerase V transcript abundance in *Arabidopsis*

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Contributed by Steven E. Jacobsen, December 11, 2014 (sent for review November 7, 2014; reviewed by James C. Carrington and Or Gozani)

DNA methylation is a mechanism of epigenetic gene regulation and genome defense conserved in many eukaryotic organisms. In *Arabidopsis*, the DNA methyltransferase DOMAINS REARRANGED METHYLASE 2 (DRM2) controls RNA-directed DNA methylation in a pathway that also involves the plant-specific RNA Polymerase V (Pol V). Additionally, the *Arabidopsis* genome encodes an evolutionarily conserved but catalytically inactive DNA methyltransferase, DRM3. Here, we show that DRM3 has moderate effects on global DNA methylation and small RNA abundance and that DRM3 physically interacts with Pol V. In *Arabidopsis* *drm3* mutants, we observe a lower level of Pol V-dependent noncoding RNA transcripts even though Pol V chromatin occupancy is increased at many sites in the genome. These findings suggest that DRM3 acts to promote Pol V transcriptional elongation or assist in the stabilization of Pol V transcripts. This work sheds further light on the mechanism by which long noncoding RNAs facilitate RNA-directed DNA methylation.

DNA methylation | epigenetic regulation | RNA polymerase | non-coding RNA | gene silencing

In eukaryotes, DNA methylation plays significant roles in gene silencing and controls many important biological processes, including genome imprinting (1), X chromosome inactivation (2), genome stability, and the silencing of transposons, retroviruses, and other harmful DNA elements (3–5). In *Arabidopsis*, DNA methylation occurs in CG, CHG, and CHH (where H = A, T, or C) sequence contexts and is controlled by at least four DNA methyltransferases: METHYLTRANSFERASE 1 (MET1), CHROMOMETHYLASE2 (CMT2), CHROMOMETHYLASE3 (CMT3), and DOMAINS REARRANGED METHYLASE 2 (DRM2). MET1 and DRM2 are the plant homologs of mammalian methyltransferases Dnmt1 and Dnmt3, respectively, whereas CMT2 and CMT3 are plant-specific methyltransferases. MET1, like Dnmt1, is important for maintenance of CG methylation during DNA replication (6–8), and DRM2, CMT3, and CMT2 control the maintenance of non-CG methylation (9–13). The establishment of DNA methylation is controlled by DRM2 (9) via a process termed RNA-directed DNA methylation (RdDM) (14–16).

In the current RdDM model, the DNA-DEPENDENT RNA POLYMERASE IV (Pol IV), RNA-DEPENDENT RNA POLYMERASE 2, and DICER-LIKE 3 function together to produce small interfering RNAs (siRNAs) that are bound by ARGONAUTE4 (AGO4). The AGO4/siRNA complex is thought to base pair with noncoding RNA transcripts produced by a DNA-DEPENDENT RNA POLYMERASE V (Pol V). Production of Pol V transcripts, as well as genome-wide Pol V chromatin occupancy, requires a complex termed DDR consisting of the putative chromatin-remodeling factor DEFECTIVE IN RNA-DIRECTED DNA METHYLATION 1 (DRD1), structural

maintenance of chromosome domain protein DEFECTIVE IN MERISTEM SILENCING 3 (DMS3), and RNA-DIRECTED DNA METHYLATION 1 (17–20). The SUPPRESSOR OF VARIATION 3–9 HOMOLOG 2 (SUVH2) and -9 (SUVH9) proteins are required for genome-wide Pol V chromatin association by binding to DNA methylation (21, 22). The co-occurrence of Pol IV-dependent siRNAs, Pol V-dependent noncoding RNA transcripts, and AGO4-DRM2 interaction (23) is thought to ultimately guide DRM2 to specific genomic sequences to cause DNA methylation.

The *Arabidopsis* genome encodes another DNA methyltransferase-like gene, *DOMAINS REARRANGED METHYLTRANSFERASE 3* (*DRM3*), that is homologous to *DRM2*. Similar to *DRM2*, *DRM3* has N-terminal ubiquitin-associated (UBA) domains and a C-terminal methyltransferase domain (24). Sequence alignments of the *DRM3* methyltransferase domains from many plant species revealed the absence of highly

Significance

DNA methylation, a chemical mark on chromatin, while not affecting DNA's primary sequence, plays important roles in silencing "bad DNA" that would become deleterious to cells if abnormally expressed. This DNA methylation-mediated silencing system against bad DNA is tightly regulated to prevent the misplacement of methylation on "good DNA." In *Arabidopsis thaliana*, DOMAINS REARRANGED METHYLTRANSFERASE2 (DRM2) controls RNA-directed DNA methylation in a pathway that also involves the plant-specific RNA Polymerase V (Pol V). The *Arabidopsis* genome also encodes an evolutionarily conserved but catalytically inactive methyltransferase, DRM3. Here, we investigate the molecular mechanism of DRM3 action on DNA methylation and its dynamic regulation of Pol V transcription. Together, this study sheds further light on the mechanism of RNA-directed DNA methylation.

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conserved key amino acids known to be critical for catalytic activity. However, DRM3 is required for full levels of DRM2-mediated DNA methylation (24, 25). Similarly, mammalian genomes encode a catalytically inactive methyltransferase termed Dnmt3L that also lacks catalytically critical residues but is required for de novo DNA methylation in vivo (26–29). Dnmt3L forms a hetero-tetrameric complex with Dnmt3a, and the interaction of Dnmt3a with Dnmt3L stimulates its activity (27–29). Despite this similarity, DRM3 and Dnmt3L differ significantly in their N-terminal domains. Dnmt3L has an N-terminal ADD domain that specifically recognizes histone 3 unmethylated at lysine 4 and targets Dnmt3 to chromatin (30, 31). In contrast, DRM3 has N-terminal UBA domains of unknown function (23, 24). Despite the importance of DRM3 for DRM2-mediated DNA methylation, it is unclear whether DRM3 acts similarly to mammalian Dnmt3L by interacting with DRM2 and stimulating DRM2 activity. It is also unknown whether DRM3 is a general or a locus-specific factor required for DNA methylation. Furthermore, the relationship between DRM3 and other components in the RdDM pathway remains elusive.

Here, through a combination of genetic, genomic, and biochemical approaches, we describe aspects of the molecular mechanism of DRM3 action in *Arabidopsis*. We show that DRM3 acts as a general factor controlling global DNA methylation and small RNA abundances and interacts physically with Pol V. To further understand the molecular basis of the DRM3–Pol V interaction, we determined Pol V genome-wide chromatin occupancy in *drm3* and found surprisingly that DRM3 target sites have either gain or loss of Pol V in *drm3*. Similar to *drm2*, the sites that lost Pol V occupancy in *drm3* had relatively low methylation levels, siRNAs, and cytosine contents, suggesting that DNA methylation is important for retaining Pol V at these sites. This methylation-dependent Pol V retention is consistent with our recent observation in *met1* (CG methyltransferase) that loss of DNA methylation results in loss of Pol V chromatin association (21). We also observed that many additional sites gain Pol V occupancy in *drm3*. However, despite gaining Pol V occupancy, we observed a reduction in the abundance of Pol V-dependent transcripts in *drm3*, suggesting that DRM3 may stabilize Pol V transcripts and/or mediate Pol V elongation. Together, our results indicate that DRM3 controls DNA methylation through its functional interaction with Pol V and by regulating Pol V transcript levels.

Results and Discussion

DRM3 Is a General Factor That Has Moderate Effects on DNA Methylation at RdDM Targets. A previous large-scale DNA methylome study of 86 *Arabidopsis* gene-silencing mutants revealed that genome-wide DNA methylation is partially reduced in *drm3* mutant (25). To test whether DRM3 acts as a general or specific factor for genome-wide DNA methylation, we performed an extensive analysis of the bisulfite-sequencing (BS-seq) data in *drm3*, as well as that of *drm2* and *nrpe1* mutants (the catalytic subunit of Pol V) for comparison. We first defined differentially methylated regions (DMRs) in each of these mutants for the CHH context because the RdDM pathway primarily impacts CHH methylation (25). We observed that the majority of *drm3* DMRs (~98%) overlap with DMRs of *nrpe1* (Fig. 1A), suggesting that DRM3 acts mainly at RdDM target sites. Consistently, we noted a strong overlap in the DMRs defined in *drm2* and *nrpe1* mutants although there were a greater number of DMRs in *nrpe1* than in *drm2*, which is likely due to the higher sequencing depth of the *nrpe1* library relative to *drm2* (*Materials and Methods*). Additionally we observed many fewer DMRs in *drm3* than in *drm2* (Fig. 1A). It is possible that DRM3 specifically acts at certain RdDM targets. Alternatively, DRM3 might have a generally weak methylation defect at most RdDM targets, in which case *drm3*-specific DMRs might be mainly due to

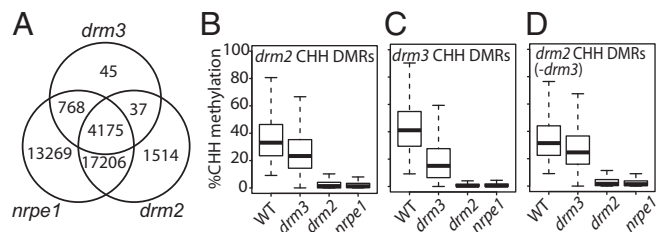


Fig. 1. DRM3 has moderate effects on global DNA methylation at RNA-directed DNA methylation targets. (A) Overlap of differentially methylated regions (DMRs) identified in *drm3*, *drm2*, and *nrpe1* mutants. CHH methylation levels over (B) *drm2* hypo-DMRs, (C) *drm3* hypo-DMRs, and (D) *drm2* hypo-DMRs excluding *drm3* hypo-DMRs in WT, *drm2*, *nrpe1*, and *drm3*. The CHH methylation in *drm3* is significantly reduced compared with WT ($P < 2.2e-16$; Wilcoxon signed rank test). CHH methylation levels in *drm2* and *nrpe1* are both significantly reduced compared with *drm3* ($P < 2.2e-16$; Wilcoxon signed rank test).

significance cutoff effects we impose in the DMR calling procedure. Toward this end, we determined the percent methylation in each cytosine context (CG, CHG, CHH) across *drm2* DMRs and noted a moderate loss of DNA methylation in *drm3* (Fig. 1B and Fig. S14). We also quantified DNA methylation patterns at *drm3* CHH DMRs and showed that *drm3* mutants had a stronger effect at *drm3* CHH DMRs than those of all *drm2* DMRs (Fig. 1C). However, even at *drm3* CHH DMRs, the effects of the *drm2* and *nrpe1* mutations were much stronger than those of the *drm3* mutation (Fig. 1C). Furthermore, at *drm2* CHH DMRs that were not called as *drm3* DMRs, we still noted a small but significant effect ($P < 2.2e-16$; Wilcoxon signed rank test) of the *drm3* mutation on DNA methylation (Fig. 1D). Together, these analyses suggest that DRM3 is a weak RdDM factor acting at most sites, rather than a locus-specific methylation factor.

DRM3 Acts Downstream of the Production of siRNAs. RdDM has two main phases, siRNA production and methylation targeting. To place DRM3 in the RdDM pathway, we analyzed the accumulation of siRNAs in the *drm3* mutants. We first assessed the abundance of siRNAs derived from the 5S *rDNA*, *AtSN1*, *siRNA 02*, and *siRNA 1003* loci in *drm3* by Northern blotting. The 5S *rDNA*, *AtSN1*, and *siRNA 1003* were shown previously to be dependent on Pol IV and Pol V (32) whereas *siRNA 02* was shown to be dependent only on Pol IV (32), but unaffected in downstream effector mutants such as *nrpe1*, *ago4*, and *drm2*. Consistent with a previous report (24), we observed partially reduced accumulation of siRNAs at 5S *rDNA*, *AtSN1*, and *siRNA 1003*, but not at *siRNA 02* (Fig. 2A), suggesting that DRM3 likely acts in the downstream portion of the RdDM pathway. To further place DRM3 within the RdDM pathway, we generated siRNA libraries and performed high-throughput sequencing from WT, *nrpe1*, *drm2*, and *drm3*. Consistent with the partial reduction of DNA methylation levels, we found that the abundance of siRNAs in *drm3* was also slightly but significantly reduced relative to WT at *drm2* DMRs (Fig. 2B and Fig. S1B). Furthermore, using small RNA clusters previously defined as being affected only in the upstream mutants versus clusters affected in downstream mutants (33), we were able to see a significant loss of 24 nucleotide small RNAs at the downstream clusters (Fig. 2C) ($P = 4.611e-10$; Wilcoxon signed rank test) but not at the upstream clusters in *drm3* mutants (Fig. 2D). Together, these data suggest that DRM3 acts in the downstream part of the RdDM pathway.

DRM3 Interacts with Pol V. To gain further insights into the mechanism of DRM3 action, we purified an HA-epitope-tagged DRM3 expressed from a transgene introduced into *Arabidopsis* *drm3* and

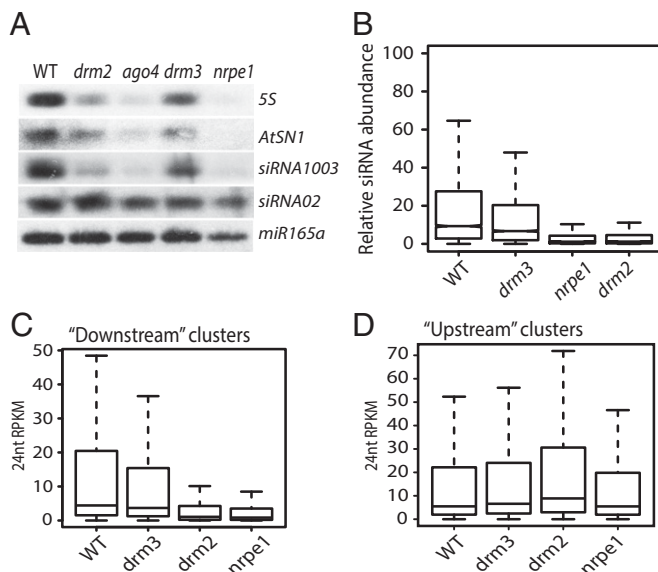


Fig. 2. DRM3 acts downstream of the production of siRNAs. (A) Northern blots show accumulation levels of small RNAs at 5S rDNA, *AtSN1*, and *siRNA1003* loci. The *siRNA02* and *miR165a* serve as two internal controls. (B) Average distribution of 24-nt siRNA reads over *drm2* CHH hypo-DMRs. The siRNA abundance is significantly reduced compared with WT in all mutants ($P < 2.2 \times 10^{-16}$; Wilcoxon signed rank test). Levels of 24-nt siRNAs in WT and mutants over previously defined siRNA clusters affected in the "downstream" (C) and "upstream" (D) RdDM mutants. The 24-nt siRNA levels are only significantly reduced in *drm3* ($P = 4.611 \times 10^{-10}$; Wilcoxon signed rank test), *drm2* ($P < 2.2 \times 10^{-16}$; Wilcoxon signed rank test), and *nrpe1* ($P < 2.2 \times 10^{-16}$; Wilcoxon signed rank test) compared with those of WT in the downstream clusters (C).

under the control of DRM3 endogenous promoter (Fig. S24). The complementation of the *drm3* mutant by transgenic DRM3 was confirmed by methylation-sensitive enzyme digestion and Southern blot analysis at the *MEA-ISR* locus (Fig. 3A). After affinity purification, DRM3-associated proteins were identified by multi-dimensional protein identification technology mass spectrometry. Table S1 shows a partial list of copurifying proteins. We did not detect any DRM2 peptides in the mass spectrometry data (Table S1), nor were we able to detect DRM2 in coimmunoprecipitation (co-IP) assays with DRM3 (Fig. S2C). This failure detection of DRM3-DRM2 interaction suggests that DRM3 acts differently than its mammalian homolog Dnmt3L and does not interact with DRM2. Instead, we identified seven Pol V subunits with significant sequence coverage, including previously reported Pol V-specific subunits NRPE1, NRPE3B, and NRPE7, as well as Pol IV/Pol V-shared subunits NRPE2, NRPE3A, NRPE11, and NRPE9A (Table S1). To further confirm these interactions, we performed immunoprecipitation and mass spectrometric analysis of the Pol V complex purified from an *Arabidopsis* transgenic line expressing the FLAG-tagged Pol V largest subunit (NRPE1-FLAG). This analysis indeed detected DRM3 peptides in addition to all known Pol V subunits (Table S2). Consistent with previous published purifications (19, 34–36), we also identified other Pol V interactors, including AGO4, DMS3, DRD1, and KTF1/SPT5L. To further validate the mass spectrometry data, we performed co-IP experiments from F1 plants expressing both HA-tagged DRM3 and FLAG-tagged NRPE1. As shown in Fig. 3B, when we pulled down NRPE1 with anti-FLAG beads, we could detect DRM3 with an anti-HA antibody. This interaction was further confirmed by a reciprocal co-IP (Fig. S2B). We were unable to detect an interaction between DRM3 and AGO4, one of the other Pol V interactors, suggesting that DRM3 may form distinct complexes with Pol V, or that the weak association of both AGO4

and DRM3 with Pol V does not allow for detection of a very weak but indirect association of AGO4 and DRM3. In summary, our data suggest that DRM3 physically associates with the Pol V complex.

DRM3 Mediates Pol V Chromatin Association at Specific Target Sites. Given the association of DRM3 with the Pol V complex, we tested whether Pol V chromatin association might be affected in *drm3* mutants in a manner similar to mutants in the DDR complex components (20). Thus, we performed chromatin immunoprecipitation followed by sequencing (ChIP-seq) of Pol V using an endogenous antibody against the NRPE1 subunit of Pol V in WT and *drm3* plants. Additionally, ChIP-seq was performed on *nrpe1* mutants as a negative control and *drm2* mutants as a control for downstream RdDM mutants to enable detection of any specific effects of the *drm3* mutation. Interestingly, using the endogenous Pol V antibody, we were able to identify >60% more high confidence Pol V binding sites ($n = 4,317$) from the resulting sequencing data than we previously observed using an epitope-tagged version of Pol V (20) ($n = 2,656$). The sequencing also revealed that neither *drm2* nor *drm3* mutants displayed large-scale loss of Pol V chromatin occupancy relative to WT, compared with the *nrpe1* mutants (Fig. 4A). This observation is largely consistent with earlier studies showing that loss of DRM2 does not affect Pol V targeting (18). However, upon close inspection of the sequencing data, we found that certain subsets of Pol V targets experienced either loss or gain of Pol V in *drm2* or *drm3* mutants (Fig. 4B and Fig. S3A and B). We also observed that the Pol V signal was increased at a larger number of sites in *drm3* than those that were decreased (Fig. 4C). In *drm2*, the situation was reversed, with more sites losing than gaining Pol V (Fig. 4C). Interestingly, sites that gained Pol V in *drm3* overlapped with sites that gained in *drm2* and vice versa (Fig. 4C). We repeated ChIP-seq analysis of Pol V in *drm3* mutants using the previously described FLAG-tagged NRPE1 transgene two more times and confirmed a similar alteration in patterns of Pol V chromatin occupancy that we observed with the endogenous Pol V antibody, and that *drm3* mutants do not alter the overall levels of Pol V (Fig. S3C and D). These data suggest that DRM2 and DRM3 dynamically regulate Pol V chromatin association and that *drm3* mutants increase Pol V occupancy at many sites.

Pol V Occupancy Is Correlated with Methylation Levels and siRNA Abundance. Pol V occupancy in the genome requires the methyl DNA binding proteins SUVH2 and SUVH9 and is almost completely lost in the strong DNA methylation mutant *met1* (21). We therefore considered the possibility that alterations of DNA methylation patterns in *drm2* or *drm3* might mediate changes in Pol V chromatin association. To test this possibility, we analyzed the methylation levels in *drm2* and *drm3* mutants at Pol V sites

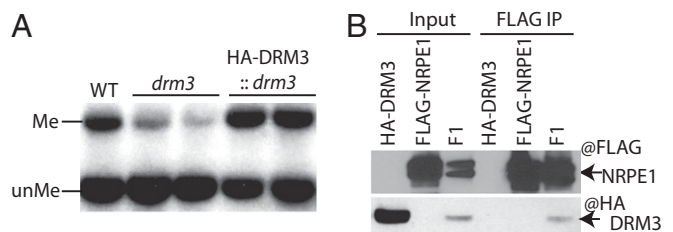


Fig. 3. DRM3 is associated with Pol V in vivo. (A) Complementation test of *drm3* mutant with pDRM3::HA-DRM3 at the *MEA-ISR* locus by restriction digestion with methylation-sensitive *MspI* and Southern blot. Me, methylated DNA; unMe, unmethylated DNA. (B) Coimmunoprecipitation assays confirming DRM3-NRPE1 interaction. Input lanes confirm the expression of the epitope-tagged proteins in the parental lines and F1. F1 represents the first generation from a cross between the two parental lines.

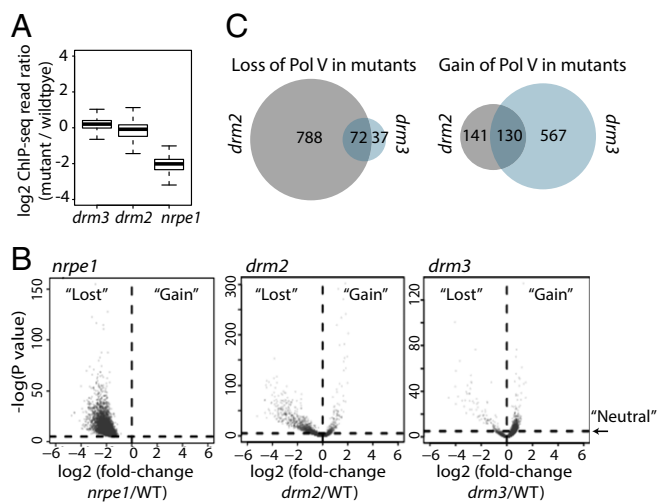


Fig. 4. DRM2 and DRM3 differentially affect Pol V occupancy. (A) Distribution of log₂ ratios of Pol V enrichment in mutant vs. WT over previously defined Pol V sites. The *drm2* shows a significantly greater loss of Pol V compared with *drm3* ($P < 2.2 \times 10^{-16}$; Wilcoxon signed rank test), and *nrpe1* shows a greater reduction than either *drm2* or *drm3* ($P < 2.2 \times 10^{-16}$; Wilcoxon signed rank test). (B) Scatter plots of the relationship between Pol V ChIP-seq fold-change in *nrpe1*, *drm2*, and *drm3* mutants relative to WT and the significance of that change (P value; Fisher's exact test) at defined Pol V binding sites ($n = 4,317$). P value cutoff ($P < 1 \times 10^{-5}$) used to define Pol V sites with either a neutral change or a significant gain or loss of Pol V for each mutant is represented by a horizontal dotted line. (C) Venn diagrams representing the overlaps of Pol V sites (total = 4,317) that lose or gain Pol V in *drm2* and *drm3* mutants.

that were classified as experiencing either loss, gain, or no change in Pol V occupancy. We found that sites that lost Pol V in *drm2* had relatively low levels of methylation in all contexts in *drm2* (yellow boxes in Fig. 5A). These sites were also the sites showing the greatest loss of DNA methylation in *drm2* (yellow boxes in Fig. S4A). In contrast, Pol V sites that experienced little alteration (red boxes) or slight gains of Pol V occupancy (blue boxes) showed high levels of methylation in all sequence contexts (CG, CHG, and CHH) (Fig. 5A and Fig. S4A), suggesting that DNA methylation is a major factor driving retention of Pol V at chromatin consistent with our previous report (21). Similarly, our siRNA analyses revealed that the sites that did not lose Pol V occupancy in *drm2* generally retained higher residual 24-nucleotide siRNA levels (Fig. S4B). Additionally, sites that lost Pol V occupancy in *drm2* showed a lower cytosine content than sites that did not lose Pol V (Fig. S4C). Given this relationship between *drm2* methylation, siRNAs, cytosine content, and Pol V occupancy, we hypothesized that the loss of Pol V chromatin association in *drm3* mutants may be similarly due to an indirect effect of *drm3*-dependent alterations of DNA methylation and siRNAs. We therefore performed a similar analysis on *drm3* mutants and found that altered levels of Pol V in *drm3* mutants followed similar trends as the sites altered in *drm2* mutants. Sites that lost Pol V in *drm3* showed the lowest levels of methylation in all contexts in *drm3* whereas sites that retained or gained Pol V tended to show higher levels of methylation (Fig. 5B). Moreover, sites showing loss of Pol V in *drm3* showed lower starting levels of methylation and cytosines and showed a greater loss of methylation and siRNAs in *drm3* than those sites that retained or gained Pol V (Fig. S4D–F). Together, these results indicate that, in the absence of DRM2 or DRM3, Pol V targeting is largely dictated by sequence composition as well as the remaining DNA methylation and siRNAs.

Modeling of Pol V Occupancy. We attempted to develop predictive models of the relationship between DNA methylation, siRNAs,

cytosine content, and Pol V occupancy in the *drm2* and *drm3* mutants. We found that models generated from the *drm2* or *drm3* methylome/siRNA profiles performed better in predicting Pol V occupancy levels than models generated using WT profiles (Fig. 5C and Fig. S5A), perhaps because DNA methylation or siRNAs become limiting for Pol V recruitment in these mutant backgrounds. By individually removing predictor variables, we noted that, of the three sequence contexts, only CG-context methylation strongly affected the predictive power of Pol V

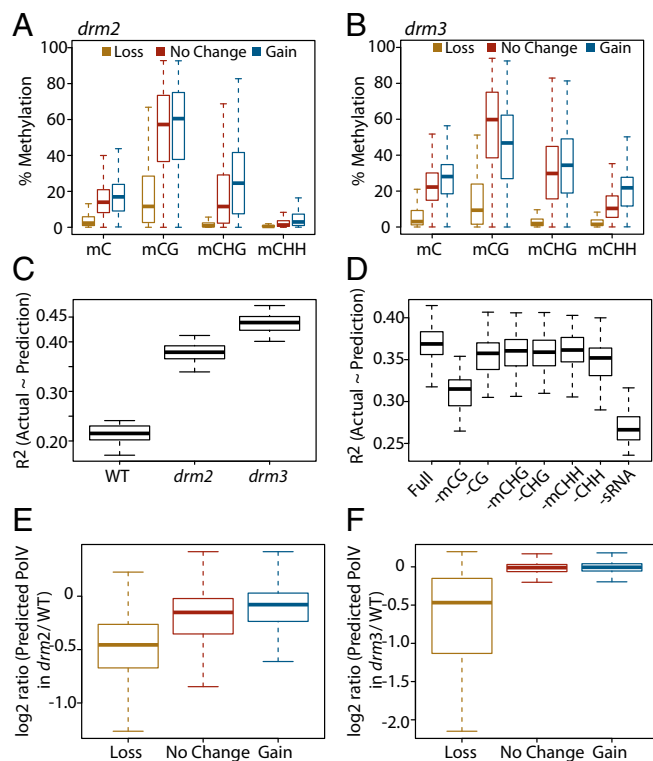


Fig. 5. Pol V association with chromatin in *drm* mutants is correlated to DNA methylation levels. (A) Methylation levels for different cytosine contexts in *drm2* at Pol V sites classified as experiencing a "loss," "gain," or "no change" in Pol V occupancy in *drm2*. (B) Same representation as A, except methylation levels in a *drm3* mutant are shown, and the Pol V sites are classified as experiencing a loss, gain, or no change in Pol V occupancy in *drm3*. The methylation levels of total methylated C (mC) and methylated CG (mCG) are significantly reduced ($P < 2.2 \times 10^{-16}$; Wilcoxon rank sum test) at loss sites compared with gain or no change sites for both *drm2* and *drm3*. (C) Distribution of R^2 values of the fit of predicted Pol V signal to the actual observed Pol V ChIP-seq signal using models built upon methylation, cytosine content, and siRNA data derived from each respective genotype. Models were trained using 3/4 of the defined Pol V peaks and tested against the remaining 1/4 of sites. Training and testing were repeated 25 times for each genotype. The WT R^2 value is significantly less than both the *drm2* and *drm3* values ($P = 5.96 \times 10^{-8}$; Wilcoxon signed rank test). (D) Distribution of R^2 values of predicted versus actual Pol V ChIP-seq signal for models trained on *drm2* data as in C using the full set of parameters ("Full") or subtracting one parameter (–mC, removing methylation data for a given context; –C, removing cytosine abundance data for a given context; –siRNA, removing 24-nt siRNA data). Both the –mCG and –siRNA models perform significantly worse than the next worse model (–CHH, $P = 5.96 \times 10^{-8}$; Wilcoxon signed rank test). (E) Distribution of log₂ ratios between predicted Pol V ChIP-seq signal in *drm2* versus predicted signal in WT at Pol V sites classified as "Loss," "Gain," or "no change" in Pol V enrichment in *drm2*. The model was trained using *drm3* data (without mCHH or siRNAs) at Pol V sites, excluding *drm2* "Loss" or "Gain" sites. (F) Analogous to E, except that sites tested were classified based on Pol V behavior in *drm3* and the model was generated using *drm2* data, excluding the *drm3* "Loss" or "Gain" sites. The loss category is significantly lower than the no change category ($P < 2.2 \times 10^{-16}$; Wilcoxon rank sum test) for both *drm2* and *drm3*.

occupancy modeling (Fig. 5D). These results are consistent with the almost complete loss of Pol V occupancy previously observed in the *met1* mutant (21) and provide additional evidence that CG methylation is critical for retaining Pol V at chromatin. Consistent with the strong role of maintenance methylation in retaining Pol V, we found that, even when we removed both CHH methylation and 24-nucleotide siRNA abundance data (two hallmarks of RdDM), we could predict Pol V occupancy levels reasonably well in *drm2* and *drm3* mutants (Fig. S5B).

Using Pol V occupancy models of *drm3* data (CG and CHG methylation, and cytosine content), we could train new models that successfully predicted the loss of Pol V that was experimentally observed in *drm2* mutants (Fig. 5E). Reciprocally, we could train models using *drm2* data to successfully predict loss of Pol V in *drm3* (Fig. 5F). However, we found that both models failed to predict the observed gain of Pol V in *drm2* and *drm3* mutants (Fig. 5E and F). Together, these results reinforce our hypothesis that the loss of Pol V chromatin occupancy in *drm2* and *drm3* mutants can be explained, and predicted, by alterations of DNA methylation caused by *drm2* and *drm3* mutants.

DRM3 Is Partially Required for Pol V-Dependent Noncoding RNA Transcript Accumulation. Although the loss of Pol V occupancy at certain sites in the *drm3* mutant can be explained by an indirect effect of the loss of DNA methylation, we also observed that *drm3* mutants show an increase of Pol V occupancy at a very large number of sites. The numbers of sites that gain Pol V in *drm3* are much greater than the number of sites that gain Pol V in *drm2* even though *drm3* has a weaker effect on the RdDM than does *drm2*. Furthermore, the sites that gain Pol V in *drm3* have lower levels of DNA methylation than those that gain Pol V in *drm2* (Fig. S4), suggesting that the gains of Pol V in *drm3* may not be simply driven by a redistribution of Pol V to highly methylated sites. The observations that *drm3* shows losses of DNA methylation, and yet shows overall gains of Pol V occupancy, suggest that DRM3 acts at some point downstream of Pol V recruitment. Given the physical interaction of DRM3 with Pol V, we hypothesize that DRM3 might be needed for the efficient production of Pol V transcripts or for their stability. To test this hypothesis, we selected a number of represented Pol V-enriched sites that had either not lost or had gained Pol V occupancy in *drm3* and assayed the Pol V RNA transcript levels in *drm3* using quantitative reverse transcription PCR. We observed

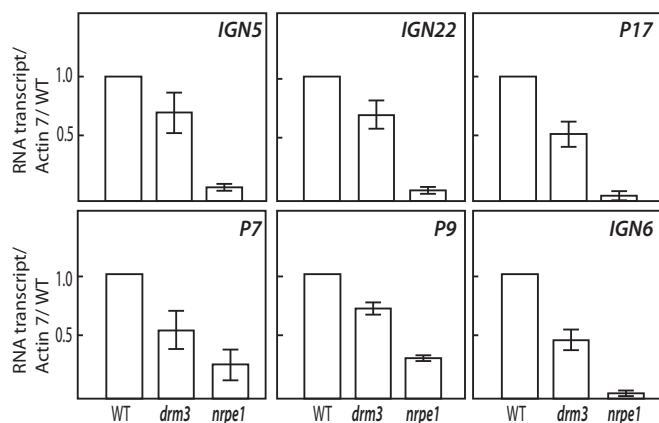


Fig. 6. DRM3 is partially required for Pol V-dependent noncoding RNA transcript accumulations. Quantitative RT-PCR analysis of the abundance of noncoding RNA transcripts from Pol V-enriched loci that either gained or did not lose Pol V occupancy in *nrpe1* and *drm3* mutants. Transcript levels were analyzed in WT, *drm3*, and *nrpe1* plants and normalized to the levels of *ACTIN7*. Error bars represent the SD of more than five replicates.

a partial but reproducible reduction of Pol V transcript at all tested sites (Fig. 6). One possible explanation for these results is that DRM3 may be involved in promoting efficient Pol V transcriptional elongation. This model might also explain the observed increase in Pol V occupancy because a more slowly transcribing Pol V would be expected to spend a longer time on chromatin, and thus give high signal strength in our chromatin immunoprecipitation experiments. A second possibility is that DRM3 might be needed for Pol V RNA transcript stability. Further in vitro Pol V transcription activity assays using a previously developed method (37) will be critical to test these possibilities. Together, these findings suggest a unique role of DRM3 in regulating the output of Pol V transcription activity and help to further elaborate the factors that are needed for proper functioning of the RdDM pathway.

Materials and Methods

Plant Material. All *Arabidopsis* plants used in this study were in the Col-0 ecotype and were grown under long day conditions. The following *Arabidopsis* mutant lines were used: *drm2-2* (SALK_150863), *nrpe1-12* (SALK_033852), *drm3-1* (SALK_136439), and *ago4* (SALK_007523).

Bisulfite Sequencing Data Analysis. The whole-genome BS-seq data from WT plants have been previously described (38), and whole genome datasets for *drm3*, *drm2*, and *nrpe1* have also been described (25). The 50 mer sequencing reads were analyzed. Identical reads were collapsed into single reads, and reads were mapped to the TAIR10 genome using BSMAP (39), allowing up to two mismatches and retaining only uniquely mapping reads. Fractional DNA methylation levels were computed by number of C (#C) / (#C + #T). DMRs were defined as previously described (25) with the following modifications: a false discovery rate (FDR) < 0.001 was required for a DMR to be called, and five cytosines with at least 5x coverage must be present in a candidate DMR. Called DMRs were not merged for downstream analysis.

RNA Analysis. Total RNA was extracted from 0.5 g of flowers using TRIzol (Invitrogen). Small RNAs were purified as previously described (40). Small RNA Northern blot was performed as described previously (41). Small RNA libraries were generated and sequenced following the manufacturer's instructions (Illumina). Adapter sequences were clipped off before mapping. Reads were mapped to the TAIR10 genome using Bowtie (42) by allowing no mismatches and keeping only reads that uniquely map to the genome. For the analyses, the small RNA counts were normalized to the size of each small RNA library by dividing the number of reads to the number of total uniquely mapping reads of 15–30 bp in size. For the comparison of "upstream" and "downstream" clusters the "pol-iv only" and "shh1/drm2/pol-v" clusters from ref. 33 were used, respectively.

Detection of Pol V-dependent transcripts was conducted as described in refs. 19 and 20. Total RNA was extracted from 0.1 g of flowers with TRIzol and treated with DNase I (Invitrogen). First-strand cDNA was synthesized with gene-specific primers (Table S3) using SuperScript III (Invitrogen). Quantification of transcripts was performed as described in ref. 19.

Generation of Epitope-Tagged DRM3 Transgenic Plants. Epitope-tagged DRM3 constructs were generated as described previously (10). The full-length genomic DNA fragment containing a 3.5-kb promoter region was amplified by PCR (Table S3), and PCR products were cloned into the pCR2.1 vector (Invitrogen) with flanking Sal I sites. BamHI and ClaI sites were then introduced at the 5' end of DRM3 by site-directed mutagenesis (200521; Stratagene). A 3xHA (HA = YPYDVPDYA) epitope tag was inserted using BamHI and ClaI sites. This tagged construct was then moved as a SalI fragment into pCAMBIA1300 binary vector and introduced into *Agrobacterium* strain AGLO, followed by transforming into *drm3-1* mutant using the floral dip method (43). The third-generation transgenic plants were used for immunoprecipitation and mass spectrometry analyses.

Affinity Purification and Mass Spectrometry. The immunoprecipitation (IP) was performed as previously described (10). The extracts from ~10 g of flowers expressing HA-DRM3 or FLAG-NRPE1 were incubated with 200 μ L of anti-HA affinity matrix (11815016001; Roche) and M2 Flag agarose beads (A2220; Sigma), respectively, at 4 $^{\circ}$ C for 2–3 h. The bead-bound complex was then washed two times with 40 mL of lysis buffer (LB) and four additional times with 1 mL of LB by mixing at 4 $^{\circ}$ C for 5 min each wash. Bound proteins were released by two times 10-min incubation with 250 μ L of 3xHA peptide (12149; Sigma) for

DRM3 and 3xFLAG peptide (F4799; Sigma) for NRPE1 at room temperature. The eluted protein complexes were precipitated by trichloroacetic acid and subjected to mass spectrometric analyses as previously described (10).

Coimmunoprecipitation. Approximately 1 g of flowers from each parental line, as well as F1 plants expressing complementing, epitope-tagged versions of both proteins, was grounded in liquid nitrogen with 5 mL of lysis buffer (LB), and the lysate was cleared by centrifugation at 10,000 × g in microfuge tubes for 10 min at 4 °C. The supernatants were incubated with 100 μL of either M2 Flag agarose (A2220; Sigma) or anti-HA affinity matrix (11815016001; Roche) for 2 h at 4 °C with rotation. The beads were then washed five times, for 5 min, with 1 mL of LB and resuspended in 100 μL of SDS/PAGE loading buffer. The various proteins were detected by Western blotting using either ANTI-FLAG M2 Monoclonal Antibody-Peroxidase Conjugate (A 8592; Sigma) at a dilution of 1:5,000, anti-HA high affinity (3F10) monoclonal antibody (11867431001; Roche) at a dilution of 1:3,000, anti-Myc 9E10 antibody (AFC-150P; Covance) at a dilution of 1:3,000, and endogenous AGO4 antibody at dilution of 1:1,000. All Westerns were developed using ECL Plus Western Blotting Detection System (RPN2132; GE Healthcare).

Genome-Wide ChIP Sequencing and Library Generation. Two grams of tissue were ground in liquid nitrogen, and ChIP was performed as previously described (20) using endogenous NPPE1 antibody. As further confirmation, we performed ChIP twice from 2 g of plant extracts expressing NRPE1-FLAG either in WT or *drm3* mutant background by FLAG beads. ChIP-enriched DNAs from the three biological replicates were subjected to library preparation and sequencing following the manufacturer's instructions (Illumina). Reads were mapped to the TAIR10 genome using Bowtie (42) by allowing up to one mismatch and keeping only reads that uniquely map to the genome. Reads mapping to identical locations were collapsed into one read. Pol V

sites were called using MACS (44) with a *P* value cutoff of 1e−1, using the WT ChIP-seq library as the treatment and the *nrpe1* library as the control library.

Pol V Modeling. To model Pol V occupancy at Pol V sites, we used methylation data derived from the whole-genome bisulfite libraries and small RNA data from the libraries described above. Cytosine abundance was derived from the TAIR10 build of the *Arabidopsis* genome. Modules were built using the random Forest package (45) using the default parameters. To test the performance of model prediction, the model was trained with ~75% of the identified 4317 Pol V peaks and then tested with the remaining 25% of peaks. This training/testing was iterated 25× for each genotype and subsequent variable subtraction experiments. The reported distribution of *R*² values is based on the correlation between the actual Pol V occupancy and predicted Pol V occupancy for the test sites of each iteration. For prediction of Pol occupancy at mutant loss/gain sites, all of the Pol V sites were used in model training, with the exception of those sites classified as affected in the mutant to be tested (i.e., when using *drm3* data to train the model to predict Pol V occupancy at *drm2* loss/no change/gain sites, those sites that experience loss/gain of Pol V in *drm2* were not included in the *drm3* training data) to avoid overfitting of the model.

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