A histone H3K4me1-specific binding protein is required for siRNA accumulation and DNA methylation at a subset of loci targeted by RNA-directed DNA methylation

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In plants, RNA-directed DNA methylation (RdDM) is a well-known de novo DNA methylation pathway that involves two plant-specific RNA polymerases, Pol IV and Pol V. In this study, we discovered and characterized an RdDM factor, RDM15. Through DNA methylome and genome-wide siRNA analyses, we show that RDM15 is required for RdDM-dependent DNA methylation and siRNA accumulation at a subset of RdDM target loci. We show that RDM15 contributes to Pol V-dependent downstream siRNA accumulation and interacts with NRPE3B, a subunit specific to Pol V. We also show that the C-terminal tudor domain of RDM15 specifically recognizes the histone 3 lysine 4 monomethylation (H3K4me1) mark. Structure analysis of RDM15 in complex with the H3K4me1 peptide showed that the RDM15 tudor domain specifically recognizes the monomethyllysine through an aromatic cage and a specific hydrogen bonding network; this chemical feature-based recognition mechanism differs from all previously reported monomethyllysine recognition mechanisms. RDM15 and H3K4me1 have similar genome-wide distribution patterns at RDM15-dependent RdDM target loci, establishing a link between H3K4me1 and RDM15-mediated RdDM in vivo. In summary, we have identified and characterized a histone H3K4me1-specific binding protein as an RdDM component, and structural analysis of RDM15 revealed a chemical feature-based lower methyllysine recognition mechanism.

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DNA methylation mainly refers to the addition of a methyl group to the fifth position of cytosine, resulting in 5′-methylcytosine (5-mC). Such methylation is involved in gene imprinting, fruit ripening, maintenance of genome integrity, and other important biological functions. In mammals, DNA methylation mainly occurs in the CG context, but can be found in all three sequence contexts, including CG, CHG, and CHH (H = T, A, or C) in plants. In plants, DNA methylation is involved in the biogenesis that involves not only Pol IV but also several other enzymes, such as CHROMOMETHYLASE 3 (CMT3) and CMT2, depending on the chromatin context. In mammals, DNA methylation mainly refers to an addition of a methyl group to the fifth position of cytosine, resulting in 5′-methylcytosine (5-mC). Such methylation is involved in gene imprinting, fruit ripening, maintenance of genome integrity, and other important biological functions. In mammals, DNA methylation mainly occurs in the CG context, but can be found in all three sequence contexts, including CG, CHG, and CHH (H = T, A, or C) in plants. In plants, DNA methylation is involved in the biogenesis that involves not only Pol IV but also several other enzymes, such as CHROMOMETHYLASE 3 (CMT3) and CMT2, depending on the chromatin context.

The RdDM pathway is responsible for DNA methylation and transcriptional silencing of transposons and other repetitive elements and is critical for pathogen defense, stress response, and many other processes in plants. This pathway comprises two main steps that depend on the plant-specific RNA polymerases Pol IV and Pol V. The first step is Pol IV-dependent siRNA biogenesis that involves not only Pol IV but also several other RdDM factors, such as RDR2 (RNA-DEPENDENT RNA POLYMERASE 2) and DCL3 (DICER-LIKE 3). The second step is siRNA-guided DNA methylation, in which Pol V-transcribed long-noncoding RNA serves as a scaffold RNA, and DRM2 is recruited to catalyze DNA methylation. This recruitment of DRM2 involves base-pairing between AGO4 (ARGONAUTE 4) and AGO6-bound siRNAs and the scaffold RNA and the participation of RdDM factors such as KTF1 (KOW DOMAIN-CONTAINING TRANSCRIPTION FACTOR 1), DRD1, DMS3, and RDM1 (RNA-DIRECTED DNA METHYLATION 1).

Different epigenetic marks, such as DNA methylation and histone modifications, can interact to regulate the chromatin state and DNA methylation level in the genome. Histone modifications are established by histone modification enzymes known as writer proteins and can be recognized by reader proteins to regulate downstream effector molecules. To date, with few exceptions, all of the reported recognition mechanisms of histone mark readers are conserved between plants and animals.

In RdDM, the recruitment of Pol IV requires SHH1 (SAWADEE HOMEODOMAIN HOMOLOG 1), which possesses a SAWADEE domain that can bind to both unmethylated H3K4 and methylated H3K9. At RdDM target loci, active histone marks, such as H3K4me2 and H3K4me3, function to prevent RdDM. On the other hand, the recruitment of Pol V requires the participation of SUVH2 and SUVH9. Although these two proteins lack histone methyltransferase activities, they both have an SRA domain that can bind to methylated DNA, indicating that they contribute to the interplay between Pol V and chromatin by binding to pre-existing methylated DNA. The crosstalk between histone modification and RdDM has been extensively studied, but the existing model cannot explain all of the targeting mechanisms of DNA methylation. Additional studies are required to increase our understanding of the interaction between epigenetic modification and RdDM components.

In the current research, we found a chromatin-binding protein, RDM15, that is involved in RdDM. Through whole-genome bisulfite sequencing and siRNA sequencing of rdm15 mutants and the wild type (WT), we demonstrate that RDM15 is required for RdDM-dependent DNA methylation and RdDM-dependent siRNA accumulation at a subset of RdDM target regions. In addition, protein structure analysis revealed that RDM15 can specifically recognize H3K4me1 through its C-terminal Tudor domain. The RDM15 Tudor domain uses an aromatic cage and a hydrogen-bonding network to achieve the monomethyllysine-specific recognition of H3K4me1; this chemical feature-based monomethyl lysine state-specific readout mechanism has not been discovered previously. Our results also show RDM15 protein enrichment at RdDM-dependent RdDM target regions and physical interaction between RDM15 and a PolV subunit. In summary, our study has identified an RdDM component, RDM15, and has elucidated the molecular mechanism underlying RDM15 function in the RdDM pathway.

Results
RDM15 is required for transcriptional silencing at RdDM target loci. In this study, we used a T-DNA mutagenized population in the Arabidopsis ros1-1 mutant background (C24 ecotype) to identify genetic factors involved in DNA methylation and transcriptional gene silencing. This screen was based on a stress-responsive RD29A promoter-driven LUCIFERASE transgene (pRD29A-LUC) in the ros1-1 mutant background. Dysfunction of ROS1 silences the pRD29A-LUC transgene and endogenous RD29A gene through heavy methylation at the RD29A promoter. A CaMV 35S promoter-driven kanamycin-resistance gene NPTII (p35S-NPTII) that is linked to the pRD29A-LUC transgene is also silenced in the ros1-1 mutant. By screening for ros1 suppressor mutants, we have discovered several components of the RdDM pathway, such as RDM1, RDM4, and KFT1. In previous studies, all RdDM mutants isolated through this screen can release the silencing of pRD29A-LUC but not of p35S-NPTII, because silencing of p35S-NPTII is not fully dependent on the RdDM pathway. In the current study, we isolated a ros1-1 suppressor mutant, rdm15-1, that can partially release transcriptional silencing of pRD29A-LUC in the ros1-1 background but cannot release the silencing of p35S-NPTII (Fig. 1a and Supplementary Fig. 1a). As shown in Fig. 1a, the luminescence after cold treatment was partially recovered in the rdm15-1/ros1-1 double mutant compared to ros1-1. Consistently, the transcript level of LUC was higher in the rdm15-1/ ros1-1 double mutant than in ros1-1 (Fig. 1b). The silencing of the endogenous RD29A gene in ros1-1 was also released by the rdm15 mutation (Fig. 1b). However, the silencing of p35S-NPTII in ros1-1 was not released by the rdm15-1 mutation (Supplementary Fig. 1a). In addition to silencing the transgene, RdDM is also required for the silencing of endogenous genomic loci. AtGP1 and AtMEA-ISR are known RdDM targets that are silenced in WT and ros1-1 mutant plants, and this silencing can be released in ago4, a known RdDM mutant, and also in rdm15 mutants (Supplementary Fig. 1b). These results suggest that RDM15 is involved in RdDM-dependent gene silencing.

To clone the RDM15 gene, we used TAIL-PCR and found a T-DNA insertion in the 7th exon of AT4G31880 (Fig. 1c). A complementation test showed that a transgene containing the native promoter-driven genomic sequence of AT4G31880 can rescue the pRD29A-LUC silencing phenotype of the rdm15-1/ ros1-1 double mutant (Supplementary Fig. 1c). To confirm the silencing function of RDM15, we ordered two T-DNA insertion alleles of RDM15 (Col-0 background): Salk_013481 and Salk_024055 (hereafter referred to as rdm15-2 and rdm15-3). Both of these mutants contain T-DNA insertions in the 7th exon of RDM15 (Fig. 1c and Supplementary Fig. 1d). The silencing of AtGP1 and AtMEA-ISR, two known RdDM targets, is released by both rdm15-2 and rdm15-3 mutations (Fig. 1d), supporting the notion that RDM15 is required for the silencing of RdDM target loci.

RdDM controls the expression of the DNA demethylase ROSI via methylation of the MEMS sequence located in the promoter region of ROSI. ROSI expression is dramatically decreased in...
many RdDM mutants, including *nrpd1a*, *nrpd1b*, and *ago4*24,25. We found that *rdm15* mutants have lower *ROS1* transcript levels (Fig. 1e and Supplementary Fig. 1e), although the levels are not as low as in *nrpd* or *ago4* mutants. These data suggest that *RDM15* functions in RdDM-dependent transcriptional gene silencing (TGS).

*RDM15* is required for RdDM-dependent DNA methylation. To characterize the effect of *rdm15* on DNA methylation, we generated single-base resolution maps of DNA methylation for Col-0, *rdm15-2*, and *rdm15-3*, with two biological replicates for each mutant. Principal components analysis showed very good consistency between the replicates (Supplementary Fig. 2a). Using the R package methylKit26, which considers variations among replicates of two mutant alleles, we identified 1390 differentially methylated regions (DMRs) in *rdm15* mutants compared with the WT; most of the DMRs (1354 out of 1390) were hypomethylated, suggesting that *RDM15* is required for DNA methylation at more than 1300 endogenous genomic regions (Supplementary Data 1).

We compared the DNA methylation patterns of *rdm15* mutants with those of known RdDM mutants. NRPD1 and NRPE1 are the largest subunits exclusive to Pol IV and Pol V, respectively. Using the same methylKit method and compared to the WT, we identified 4293 hypo DMRs in *nrpd1-3* (Col-0 ecotype) and 4629 hypo DMRs in *nrpe1-11* (Col-0 ecotype) (Supplementary Data 1).

We found that the Pol IV, Pol V, and *RDM15* targets have comparable genomic compositions. Among the hypo DMRs, 82% in *nrpd1-3*, 81% in *nrpe1-11*, and 79% in *rdm15* are in TE regions (Fig. 2a). Similar to known RdDM targets, *rdm15* DMRs show DNA hypomethylation in all three sequence contexts (mCG, mCHG, and mCHH) compared to WT (Fig. 2b), although the methylation level in *rdm15* mutants is not as low as in *nrpd1-3* (Fig. 2b). We further analyzed the overlap between RDM15 targets and known RdDM targets. As shown in Fig. 2c,
94% (1269/1354) and 95% (1281/1354) of rdm15 hypo DMRs overlap with the nrpd1 and nrpe1 hypo DMRs, respectively. DNA methylation profiles of several representative rdm15 hypo DMRs are shown in Fig. 2d and Supplementary Fig. 2b. These results revealed that RDM15 is required for methylation at a subset of RdDM target loci, suggesting that RDM15 may be an RdDM component.

RDM15 regulates RdDM-dependent siRNA accumulation. We generated genome-wide siRNA profiles for WT (Col−0), rdm15-2, and rdm15-3 with two biological replicates for each genotype (Supplementary Fig. 3a). In Arabidopsis, RdDM has two main steps, i.e., biogenesis of 24-nt siRNAs and guidance of DRM2 to RdDM target by the siRNAs. Genomic regions showing RdDM-dependent DNA methylation always have siRNA enrichment. To determine whether RDM15 is involved in siRNA biogenesis, we compared rdm15 mutants with Col−0, and identified 2808 RDM15-dependent siRNA clusters (hereafter referred to as RDM15 siRNAs) (Supplementary Data 2) that showed decreased 24-nt siRNA accumulation in rdm15 mutants relative to Col−0. This result suggested that RDM15 is required for 24nt siRNA accumulation in vivo (Fig. 3a).

RdDM-dependent siRNAs can be classified into upstream siRNAs (siRNAs dependent on Pol IV only) and downstream siRNAs (siRNAs dependent on both Pol IV and Pol V)\(^\text{16}\). The upstream siRNAs are affected only in mutants defective in upstream RdDM components, such as nrpd1, whereas the downstream siRNAs are affected not only in these mutants but also in mutants defective in downstream RdDM components, such as nrpe1 and drm2 (Fig. 3b). To position RDM15 in the RdDM pathway, we examined the enrichment of RDM15-dependent siRNAs and found that RDM15 siRNA levels are lower in nrpd1-4, nrpe1-12, and
drm2-2 compared to the WT (Fig. 3b). In addition, the Pol IV-only siRNAs are not significantly affected in rdm15 mutants. In contrast, Pol V-dependent siRNAs are significantly reduced in rdm15 mutants (Fig. 3a). These results showed that RDM15 is mainly required for the accumulation of downstream siRNAs in the RdDM pathway, suggesting that RDM15 functions in a downstream step of RdDM.

To further understand the relationship between the siRNAs and DNA methylation in rdm15 mutants, we monitored changes in siRNA enrichment at hypo DMRs in rdm15 mutants. The siRNA levels were clearly lower at hypo-DMRs in rdm15 mutants than in the WT (Fig. 3c). Consistent with this result, the DNA methylation level, including mCG, mCHG, and mCHH, was decreased in regions of RDM15-dependent siRNA clusters (Fig. 3d).
and Supplementary Fig. 3b). The siRNA levels at several representative hypo-DMRs are shown in Fig. 3e and Supplementary Fig. 3c. These results suggested that the change of DNA methylation is associated with the change of siRNA levels at RDM15-dependent RdDM targets.

RDM15 physically interacts with NRPE3B, a subunit exclusive to Pol V. To further investigate how RDM15 affects RdDM, we identified RDM15-interacting proteins by performing immunoprecipitation (IP) followed by liquid chromatography–tandem mass spectrometry (LC–MS/MS) with RDM15-3xMYC/rdm15-3 and RDM15-3xFLAG/rdm15-3 plants (Supplementary Fig. 4a). WT plants that do not express RDM15-3xMYC or RDM15-3xFLAG were used as controls. We found that NRPE3B, a subunit exclusive to Pol V, was co-immunoprecipitated with anti-MYC antibodies and anti-FLAG antibodies in the RDM15-3xMYC and RDM15-3xFLAG transgenic plants, respectively, but not in the control plants (Fig. 4a and Supplementary Data 3). We validated the interaction between RDM15 and NRPE3B using a split luciferase complementation assay and a BiFC assay in tobacco leaves (Figs. 4b, c). In addition, we examined Pol V-dependent transcripts in rdm15 mutants. The transcripts of IGN25 and IGN27 were identified as Pol-V dependent transcripts in previous studies. We found that the transcript levels of IGN25 and IGN27 were decreased in rdm15-2 and rdm15-3 mutants compared to WT (Supplementary Fig. 4b), suggesting that RDM15 is required for Pol V-dependent transcription. These data indicated that the RdDM function of RDM15 may involve its interaction with Pol V.

The RDM15 Tudor domain specifically recognizes H3K4me1. The results above show that RDM15 functions in the regulation of siRNA accumulation and DNA methylation at a subset of RdDM target loci. To further investigate how RDM15 functions, we analyzed its protein sequence and found two domains with known functions: an N-terminal ARM repeat, which is known to mediate protein–protein interactions, and a C-terminal Tudor domain, which functions as a histone mark reader. The Tudor domain has been found in many proteins related to epigenetic regulation and acts as a histone mark reader module that recognizes methylated lysine or arginine marks. In addition, we found histone H3 in the affinity purification of RDM15 in vivo (Fig. 4a). In plants, histone mark readers sometimes have plant-specific binding targets and functions. This prompted us to
that no in vivo H4K20 methylation could be detected by mass RDM15. In addition, researchers previously reported that the indicating that H4K20me1 is not the optimal binding partner for further con

modi

while the H4K20me1 yielded a 15-fold lower binding af

strongest binding (1.47

f)

Tudor domain fold with

crystallization, only the H3A1 to H3Q5 segment of the peptide

barrel structure that resembles the structure of other Tudor
cation (Fig. 6c). Mutations of residues of either the aromatic cage or of those involved in hydrogen bonding interactions with the peptide resulted in a significant decrease in the binding affinity (Fig. 6d, e), revealing that these residues are critical for the recognition of H3K4me1 by RDM15.

We expressed the mutated RDM15 (W616A/Y623A/D643A/ D645A) incapable of binding to H3K4me1 in rdm15-3 mutant plants, generating mutated RDM15-3×MYC/rdm15-3 plants. We found that the H3K4me1 was co-immunoprecipitated by wild-type RDM15 but not by mutated RDM15 (Supplementary Fig. 5e), suggesting that the mutations in the Tudor domain interrupted the binding of H3K4me1 in vivo. We examined the methylation level using methylation-sensitive PCR at several RDM15-dependent RdDM loci and found that the wild type but not the mutated RDM15 rescued the methylation phenotype in rdm15-3 (Supplementary Fig. 5d), suggesting that when RDM15 is mutated to lose the binding of H3K4me1, the RdDM function of RDM15 is also impaired.

In the structure of the RDM15-H3K4me1 complex, the recognition of the H3 tail by the RDM15 Tudor domain highlights the specific recognition of H3R2 and H3K4me1, while other surrounding residues and the intervening H3T3 do not

The structure of the RDM15 Tudor domain in complex with H3K4me1 peptide. To further investigate the molecular mechanism underlying the specific recognition of the H3K4me1 mark by RDM15, we carried out structural studies. The crystal structure of the RDM15 Tudor domain in complex with the H3 (1–15) K4me1 peptide was determined using the SAD method and was refined to 1.7 Å resolution (Supplementary Table 2 and Fig. 5c). Although a 15-residue H3K4me1 peptide was used in the H3K4me0 NDB no detectable binding. The RDM15 Tudor domain bound much more strongly to the RdDM function (Fig. 5c). In general, both H3R2 and H3K4me1 are specifically recognized by a hydrogen-bonding network and electrostatic interactions involving their charged side chains, as well as by an aromatic cage that captures the methyl group of the mono- methylation states. NDB no detectable binding.

chains of H3R2 and H3K4me1 form a pincer-like conformation, thereby anchoring on two adjacent negatively charged surface pockets of the Tudor domain and highlighting a significant structural and chemical complementarity (Fig. 5d). In detail, the H3R2 inserts its side chain into a negatively charged pocket and forms hydrogen-bonding interactions with Glu647 and Gln654 of RDM15, as well as a salt bridge interaction with Glu647 (Fig. 6a). For H3K4me1, the interactions can be divided into two parts. Three aromatic residues, Trp616, Tyr623, and Tyr641, form an aromatic cage to accommodate the methyl group of the mono-methyllysine from one side, which is similar to the other canonical methyllysine readers29,32 (Fig. 6b). On the other side, the two free protons of the monomethylammonium group are fully coordinated with Asp643 and Asp645 by hydrogen bonding and salt bridge interactions (Fig. 6b), representing a mono-methyllysine recognition mechanism (Supplementary Fig. 6 and see detailed analysis in the Discussion). The H3T3 positioned between H3R2 and H3K4me1 stretches its side chain against the protein and is not involved in the interaction with the RDM15 Tudor domain (Fig. 5c). In general, both H3R2 and H3K4me1 are specifically recognized by a hydrogen-bonding network and electrostatic interactions involving their charged side chains, as well as by an aromatic cage that captures the methylation modification (Fig. 6c). Mutations of residues of either the aromatic cage or of those involved in hydrogen bonding interactions with the peptide resulted in a significant decrease in the binding affinity (Fig. 6d, e), revealing that these residues are critical for the recognition of H3K4me1 by RDM15.
contribute to the recognition. The RDM15 Tudor domain therefore recognizes an RXKme1 (X here stands for any residue) motif on histone tails. Among all four types of Arabidopsis histone proteins, there are two additional sites that contain the same motif and that might be recognized by the RDM15 Tudor domain: H2A R3T K5 and H4 R3G K5. A previous mass spectrometry study showed that the lysine residues of both of these two sites can be acetylated, but no methylation modification was identified in vivo. It follows that although the RDM15 Tudor domain only recognizes two residues on the histone tail, the specific RXKme1 pattern only occurs in the H3K4 region, which ensures that recognition only occurs at H3K4me1, thereby explaining the sequence specificity.

Characterization of chromatin targets of RDM15 binding. We performed chromatin immunoprecipitation using native promoter-driven RDM15-3xFLAG/rdm15-3 followed by high-throughput sequencing (ChIP-seq). We observed higher RDM15 protein enrichment in RDM15-dependent siRNA regions than in Pol IV-only siRNA regions (Fig. 7a), which is consistent with our above finding that RDM15 is required for the accumulation of RDM15 siRNAs but does not affect the accumulation of Pol IV-only siRNAs (Fig. 3). When we ranked RDM15 siRNA regions by the difference in siRNA accumulation between rdm15 and the WT, we found that the change in siRNA accumulation (rdm15 vs. the WT) was positively correlated with RDM15 protein enrichment; the change in siRNA accumulation in nrdp1 vs. the WT

Fig. 6 Details of the specific interaction between the RDM15 Tudor domain and the H3K4me1 peptide. a The side chain of H3R2 is specifically recognized by Glu647 and Gln654 of the Tudor domain through a salt bridge and hydrogen-bonding interactions. The hydrogen bonds are highlighted with dashed red lines. b The methyl group of H3K4me1 is specifically accommodated within an aromatic cage formed by Trp616, Tyr623, and Tyr641. The two monomethylammonium protons of monomethyllysine form hydrogen bonding and salt bridge interactions with two negatively charged Asp643 and Asp645 residues. c A schematic representation of the intermolecular interactions between RDM15 and the H3K4me1 peptide. d, e The ITC binding curves between the H3K4me1 peptide and the RDM15 Tudor domain mutations showing that the disruption of the aromatic cage (d) or the residues involved in hydrogen bonding interactions (e) dramatically decreases the binding affinity.
served as the negative control (Fig. 7b and Supplementary Fig. 7a). These results suggest that the influence of RDM15 on siRNA accumulation is correlated with its protein enrichment. In addition, the distribution pattern of H3K4me1 but not of H3K4me2 and H3K4me3 was similar to that of RDM15 (Fig. 7c and Supplementary Fig. 7b), which is consistent with our finding of high-affinity binding of RDM15 to H3K4me1. The enrichment of RDM15 and H3K4me1 is shown for several representative RDM15 target regions in Fig. 7d and Supplementary Fig. 7c.

It has been reported that H3K4me1 is enriched in the gene body and flanking regions. Here, we showed an association of H3K4me1 with RDM15-dependent RdDM targets. Pol IV-only siRNA regions, which were not affected in rdm15, served as the control. In the boxplots, the center is the median of the data; upper bound of the box is an upper hinge, 75% quantile of the data; lower bound of the box is a lower hinge, 25% quantile of the data; minima is the smallest data point greater than or equal to lower hinge – 1.5 x interquartile range (IQR); maxima is the largest data point less than or equal to upper hinge + 1.5 x IQR.

**Discussion**

RdDM is an important DNA methylation pathway in plants and has been extensively studied during the past two decades. Several forward and reverse genetics screens, including the rosl suppressor screen in this study, were used to identify RdDM components. Through this rosl suppressor screen, several RdDM mutants, such as rdm1, rdm4, and ktf1, were identified. In the current study, an RdDM mutant, rdm15, was isolated and characterized. The methylome and siRNA analyses revealed that RDM15 is required for RdDM-dependent siRNA accumulation and DNA methylation at a subset of RdDM targets. In addition, the siRNA analysis and the interaction of RDM15 with Pol V suggested that RDM15 functions in the downstream steps of the RdDM pathway.

Several histone modifications, such as H3K9me2, have been reported to be associated with RdDM targets. Here, we showed an association of H3K4me1 with RDM15-dependent RdDM.
targets. The structural analysis demonstrated that RDM15 can specifically recognize and bind to H3K4me1 (Fig. 5). In addition, ChIP-seq results showed that the genomic distribution of RDM15 protein enrichment is similar to that of H3K4me1 but not H3K4me2/3 (Fig. 7 and Supplementary Fig. 7). H3K4me1 is usually thought to be associated with actively transcribed genomic regions. Pol V-transcribed regions are presumably more active than untranscribed regions. We speculate that RDM15 may help recruit Pol V to a subset of RdDM targets by recognizing the H3K4me1 mark at these genomic regions. In addition, we found that RDM15 is also associated with H3K4me1 in genic regions, which are mostly not RdDM target regions. This result indicated that RDM15 may have an RdDM-independent function in these regions.

Many structural and functional studies have identified the general principles explaining the specific recognition of histone lysine methylation marks29,30,32. However, only a few studies have investigated the specific recognition of lower methyllysine, especially the monomethyllysine. L3MBTL1, a previously reported low-methylated lysine-specific binding protein, can specifically recognize mono- and dimethyllysine via the narrow and deep surface cavity of the protein (Supplementary Fig. 6a)40. In the recognition mode of L3MBTL1, almost the entire side chain of the methyllysine inserts into a deep cavity on the L3MBTL1 protein surface (Supplementary Fig. 6a). The size of the cavity therefore limits the size of the substrate. For L3MBTL1, the methyllysine must be in the lower methylation states of di- or monomethyllysine, representing a physical selection mechanism. The molecular mechanism for the specific recognition of H3K4me1 by the CW domain of Arabidopsis SDG8 was recently determined41. Like L3MBTL1, the SDG8 CW domain has a narrow cavity that accommodates the small side chain of H3K4me1 (Supplementary Fig. 6b), i.e., the recognition of H3K4me1 by SDG8 also involves physical selection41. In contrast, H3K4me1 is anchored within a surface groove of RDM15, with only the tip of the methyl group inserting into the canonical aromatic cage (Supplementary Fig. 6c). The specificity for the monomethylammonium protons by the two negatively charged residues, which increases the binding affinity for monomethyllysine. In other words, the specificity of H3K4me1 for RDM15 involves a chemical selection mechanism. The increase of the methylation state of the methyllysine from H3K4me1 to H3K4me2 or H3K4me3 would disrupt or even fully eliminate the hydrogen bonding interaction observed between RDM15 and the H3K4me1 side chain, resulting in a decreasing binding affinity as revealed by our ITC data (Fig. 5b). This selection of a lower methylation state based on chemical features clearly differs from selection based on physical shape previously reported for L3MBTL1 and SDG8, and therefore represents a hydrogen bonding interaction-based but not a shaped-based mechanism for specific monomethyllysine recognition.

The Tudor domain can function in multiple modes, including single Tudor, double Tudor, tandem Tudor, or triple-linked Tudors like Spindlin124. A single Tudor domain has been reported to be able to recognize H3K36me3, H4K20me2, and H3K4me42,28,42. The Tudor domains of PHF1 and PHF19 have similar structures and recognition mechanisms and can specifically recognize the H3K36me3 mark; the structures of their complexes with H3K36me3 peptide have been reported42-45, which enables us to perform a structure-based comparison. The RDM15 Tudor and PHF1 Tudor have a generally similar folding topology, with their aromatic cages occupying almost identical positions (Supplementary Fig. 6d)45. It is interesting that the H3K4me1 and H3K36me3 peptides occupy different positions, although their methionylammonium groups occupy the same position (Supplementary Fig. 6d), suggesting that the conserved aromatic cage of the Tudor domain serves as a docking site for methyllysine, with the diversified surrounding residues contributing to the histone mark specificity. PHF1 uses a four-residue aromatic cage to accommodate H3K36me3 (Supplementary Fig. 6d)45. Although Tyr47 and Phe65 of PHF1 adopt an almost identical conformation as Tyr623 and Tyr641 of RDM15, the Trp41 of PHF1 has a distinct torsion angle relative to Trp616 of RDM15, resulting in a broader pocket in PHF1, which is suitable for accommodating trimethyllysine; RDM15, in contrast, has a narrow pocket that only accommodates monomethyllysine. We also compared the structure of the RDM15-H3K4me1 complex with those of the JMJD2A double Tudor-H3K4me3 complex (PDB code: 2GFA) and SFG29 tandem Tudor-H3K4me3 complex (PDB code: 3MEA)46,47. Although it is difficult to superimpose the RDM15 Tudor to the corresponding Tudor domains of JMJD2A and SFG29 that bind to methylated histone peptides, the RDM15 Tudor can be well superimposed to the remaining Tudor domains of JMJD2A and SFG29 without peptide binding (Supplementary Fig. 6e), suggesting that single Tudor and tandem Tudor provide different platforms to recognize histone marks with diversified modes and histone binding surface.

Our genetic and epigenomic analyses identified RDM15 as a factor required for siRNA accumulation and DNA methylation at a subset of genomic regions targeted by RdDM. Our biochemical and structural results demonstrated that RDM15 is a reader of H3K4me1, and revealed a hydrogen bonding interaction-based mechanism of recognition of lower methyllysine marks. Future work will determine whether RDM15 may affect the recruitment of Pol V to RdDM target regions and whether the mechanism of recognition of lower methyllysine mark by RDM15 may be applicable to other histone mark readers.

Methods

Plant materials and growth conditions. Arabidopsis C24 ecotype plant materials were used in this study (WT, ros1-1, ros1-1 rdm15-1, ros1-1, nrpd1-1) carried a homozygous T-DNA insertion that contained the p35S-NPTII and P35DA-LUC transgenes. T-DNA mutagenized ros1-1 populations were used for ros1-1 suppressor screening according to luciferase signals37. Plant materials of Columbia-0 (Col) ecotype included rdm15-2 (Salk_033481), rdm15-3 (Salk_024055), nrpd1-3 (SALK_128428C), and nrpe1-11 (SALK_029919C). Plants were grown in growth rooms at 22 °C with a 16–8 h light-dark cycle.

Whole-genome bisulfite sequencing and data analyses. Genomic DNA was extracted from 2-week-old seedlings using the Plant DNeasy mini kit (Qiagen) and was sent to the Core Facility for Genomics at the Shanghai Center for Plant Stress Biology (PCSB) for whole-genome sequencing (WGBS).

For WGBS data analysis, the raw data were trimmed using Trimmomatic48 with parameters “LEADING:20 TRAILING:20 SLIDINGWINDOW:4:15 MINLEN:50”. Only clean paired-end reads were mapped to the TAIR10 genome using tool SAMtools50 with parameters “–LEADING:0 –TRAILING:0 –SLIDINGWINDOW:4:15 –MINLEN:50”.

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Small RNA data analysis. Total RNA was extracted from 2-week-old seedlings using the TRIzol (Invitrogen) method. RNA samples were separated on a PAGE gel, and the 18– to 30-nt fraction of the gel was cut for small RNA purification. Linear preparation and sequencing were performed using Illumina reagents according to the manufacturer’s instructions at the Genomics Core Facility of PCSB. Small RNA data were analyzed according to ref. 57 with minor modifications. In
brief, the adapter was removed using the "fastx_clipper" command from FASTX-toolkit. After adapter sequences were trimmed, clean reads with sizes ranging from 18-31 nt were mapped to the Arabidopsis genome (TAIR10) using Bowtie2 with parameters "-v 0 -k 10". Reads that overlapped with annotated rRNAs, snRNA, or snoRNA sequences were excluded. Read counts were normalized to Reads Per Ten Million (RPTM) based on the total abundance of genome-matched small RNA reads. The "hits-normalized-abundance" (HNA) values were calculated by dividing the normalized read count in RPTM by each small RNA hit, where a hit is defined as the number of loci at which a given sequence perfectly matches the genome. The HNA values of small RNAs of all sizes within individual non-overlapping 200-bp windows throughout the whole genome were compared between the mutant and the WT. To focus on regions that show significant small RNA accumulation in the WT plants, HNA values in the mutant and the WT samples were summed, and a cutoff of 25 was applied. Subsequently, only those 200-bp regions that showed ≥2-fold HNA reduction in the mutant were identified as small RNA-depleted regions. In Fig. 3a, small RNA data for Col-0, rdr15-2, and rdr15-3 were sequenced for the project. Small RNA data for nrpi were downloaded from the NCBI GEO database with accession number GSE8380252. In Fig. 3b, all of the small RNA data were downloaded from the NCBI GEO database with accession number GSE4356818.

Affinity purification and mass spectrometry. The RDM15 genomic sequence was fused in-frame to the 3xFLAG tag and 3xMYC tag, and the fused sequences were inserted into the pCAMBIA1305 backbone and were transformed into Arabidopsis plants. A 5′-p-galacturonase fragment was harvested from transgenic plants and ground in liquid nitrogen. The proteins were extracted in 25 ml of lysis buffer [50 mM Tris (pH 7.5), 150 mM NaCl, 5 mM MgCl2, 10% glycerol, 0.1% Nonidet P-40, 0.5 mM DTT, 1 mM PMSF, and 250μl of plant cell protease inhibitor (Sigma)]. After the proteins were extracted, 2 μl anti-Flag M2 (Sigma; F3165) and 2 μl anti-Myc (Sigma; M5546) were added and incubated at 4 °C for 2-3 h. The resins were then washed with the lysis buffer at least five times. The protein samples were sent to the Core Facility for Proteomics at the Shanghai Center for Plant Stress Biology (PSC) for affinity purification and mass spectrometry.

Histone peptide array. Histone peptide array assay was performed using the MODified Histone Peptide Array kit (Active Motif) according to the manufacturer's instructions. In brief, the slide was first blocked for 2 h at room temperature, then washed three times with TBST buffer [10 mM Tris·HCl (pH 7.5), 30 mM NaCl, and 0.05% Tween-20], the slide was incubated overnight with 10 mg of GST-fused purified RDM15 protein at 4 °C in 10 ml of binding buffer [50 mM Hapes (pH 8.0), 150 mM NaCl, 2 mM DTT, and 0.05% Nonidet P-40]. After it was washed three times with TBST buffer, the slide was incubated with anti-GST primary antibody at 1/3000 dilution and secondary antibody at 1/5000 dilution in TBST buffer for 1 h at room temperature, respectively. After the washing, the slide was visualized using Lumi-light ECL substrate (Roche). The intensity of the blotted slide was analyzed according to the instructions of the analysis software of the MODified Histone Peptide Array. The quality of the sequencing data was checked with FastQC. The paired-end reads were mapped to the TAIR10 genome of Arabidopsis thaliana (TAIR10) with bowtie2 with parameters--very-sensitive--no-udem--no-mixed--no-discordant -k 5. To remove potential PCR duplicates, a markup from SAMtools was used. After mapping, only uniquely mapped reads were retained for downstream analysis. The fragment number of interested regions was counted by featureCounts with parameters "-p -O". The RDM15 enrichment (H3K4me1) signal was calculated as log2[(1 + n_CHIP)/N_CHIP] - log2[(1 + n_Input)/N_Input], where n_CHIP and n_Input represent the number of mapped CHIP and Input fragments in the interested regions, and N_CHIP and N_Input are the numbers of all mapped unique fragments. The public data for H3K4me2 and H3K4me3 were downloaded from the NCBI GEO database with accession number GSE113076.

BIFC. Agrobacterium tumefaciens strain GV3101 carrying constructs expressing RDM15-GFP, NRPE3B-βFP, and RDM15-βFP, or NRPE3B-βFP were infiltrated into N. benthamiana leaves. After 2 d, the infiltrated areas were examined with a Leica TCS-SP8 microscope. This experiment was performed once. Real-time RT-PCR analysis. For real-time RT-PCR analysis, total RNA was extracted and contaminating DNA was removed with RNase-free DNase (RNeasy mini kit; Qiagen). mRNA (1 μg) was used for the first-strand cDNA synthesis with Takara RT-PCR Systems following the manufacturer's instructions. The cDNA reaction mixture was then diluted five times, and 1 μl was used as a template in a 15-μl PCR reaction with SYBR Green mix (Takara). Two or three independent biological replicates were used for analysis. Primer sequence information is listed in Supplementary Table 3.

PCR-based DNA methylation analysis. Genomic DNA was extracted using the Plant DNAeasy mini kit (Qiagen) and 100 ng of the genomic DNA was digested with McrBC (NEB) in a 50 μl reaction system for 12 h. Then, a 1 μl digested DNA template was used for the target-specific PCR reaction. PCR products were analyzed by gel electrophoresis. Digestion without GTP was used as the control. Two independent biological replicates were used for analysis. Primer sequence information is listed in the Supplementary Table 3.

Protein expression and purification. The Tudor domain of Arabidopsis thaliana RDM15 (residues 598–662) was cloned into a PET-Sumo vector to fuse a hexahistidine and yeast Sumo tag to the target protein. The recombinant protein was expressed in E. coli strain BL21(DE3) by IPTG induction with a concentration of 0.3 mM. The protein was purified using a HiTrap (GE Healthcare) column. The His-Sumo tag was removed by upl digestion followed by a second step with the HisTrap column. The target protein was further purified using a Superdex G75 column (GE Healthcare). The Se-Met labeled protein was expressed in Strep (Anatrace) containing M9 medium and was purified using the same protocol as used for the WT protein. All of the mutants were generated using a PCR-based mutagenesis method and were expressed and purified using the same protocol as used for the WT protein. All peptides in this research were synthesized by GL Biochem (Shanghai).

Crystallography, data collection, and structure determination. Before crystallization, the purified RDM15 Tudor domain was concentrated to 20 mg/ml and mixed with H3(1-15) K4me1 peptide, which was crystallized in a condition of 0.1 M Tris-Cl, pH 8.5, and 20% PEG 1000. The crystallization was conducted using the sitting-drop vapor diffusion method at 20 °C. The RDM15 Tudor domain in complex with H3K4me1 peptide was crystallized in a condition of 0.1 M Tris-Cl, pH 8.5, and 20% PEG 1000. The crystals were soaked in the reservoir solution supplemented with 15% glycerol and were flash-cooled in liquid nitrogen for data collection. All the diffraction data were collected at the Shanghai Synchrotron Radiation Facility beamline BL17U1 and were processed using the HKL2000 program. The structure of the RDM15 Tudor domain in complex with the H3K4me1 peptide was solved using the SAD method implemented in the Phenix program. Model building was conducted using the Coot program. The geometry of the model was analyzed using the MolProbity program. All of the molecular graphics were generated using the program PyMol (Schrödinger, Inc.). A summary of diffraction data and structure refinement statistics is provided in Supplementary Table 2.

Isothermal titration calorimetry. ITC binding was measured using a Microcal PEAQ-ITC instrument (Malvern). The purified proteins were dialyzed against a buffer containing 100 mM NaCl, 20 mM HEPES, pH 7.0, and 2 mM β-mercaptoethanol and were diluted to a concentration of about 0.8–0.12 mM. The peptide was dissolved in the same buffer with a concentration of about 1.0–1.5 mM. All of the titrations were performed at 20 °C. The data were analyzed using the Origin 7.0 program.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. The data that support this study are available from the corresponding authors upon reasonable request. The sequencing data generated in the course of this study have been deposited in NCBI's Gene Expression Omnibus and are accessible through GEO Series accession number GSE154302. The mass spectrometry data have been deposited in Integrated Proteome Resources with the accession number IPX0002945000. The structure of the Arabidopsis RDM15 Tudor domain in complex with an H3K4me1 peptide has been deposited in the RCSB Protein Data Bank with the accession code 7DE9. Source data are provided with this paper.

Received: 27 October 2020; Accepted: 21 April 2021; Published online: 07 June 2021

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Acknowledgements
We thank the staff at beamline BL17U1 at the Shanghai Synchrotron Radiation Facility for their help with data collection. This work was supported by the National Key R&D Program of China (2018YFD1000200 to Z.L., 2016YFA0503200 to J.D.), the Strategic Priority Research Program of the Chinese Academy of Sciences (XDB27040000 to Z.L., XDB27040101 to J.-K. Z.), Shenzhen Science and Technology Program (JCYJ20200109110403829 and KQTD20190929173906742 to J.D.) and Key Laboratory of Molecular Design for Plant Cell Factory of Guangdong Higher Education Institutes (2019KSY006 to J.D.).

Author contributions
Q.N., L.W., and Hu.Z. performed experiments. C.-G.D. performed histone peptide array. Z.G. and T.B. contributed to the mutation screen. Z.S. and L.C. contributed to the ITC and structure analysis. K.T. did the bioinformatics analysis. C.K. is involved in the discussion of the research. He.Z. performed the sequencing experiments. Q.N., Z.L., J.D., and J.-K.Z. designed the study, interpreted the data, and wrote the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41467-021-23637-4.

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Peer review information Nature Communications thanks Zhanxin Wang, Tatiana Kutateladze, and the other, anonymous, reviewer for their contribution to the peer review of this work. Peer reviewer reports are available.

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