H3K9 methylation is usually associated with DNA methylation, and together they symbolize transcriptionally silenced heterochromatin. A number of proteins involved in epigenetic processes have been characterized. However, how the stability of these proteins is regulated at the post-translational level is largely unknown. Here, we show that an Arabidopsis JmjC domain protein, JMJ24, possesses ubiquitin E3 ligase activity. JMJ24 directly targets a DNA methyltransferase, CHROMOMETHYLASE 3 (CMT3), for proteasomal degradation to initiate destabilization of the heterochromatic state of endogenous silenced loci. Our results uncover an additional connection between two conserved epigenetic modifications: histone modification and DNA methylation.

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Methylation of DNA and of H3K9 (H3K9me) are two conserved epigenetic modifications generally associated with gene silencing in eukaryotes. DNA methylation, which is established by DNA methyltransferase 3 (DNMT3) and maintained by the maintenance methyltransferase DNMT1, occurs almost exclusively in the symmetric CG context in mammals. In contrast, in plants, cytosines in all classes of sequence context (that is, CG, CHG, and CHH, where H = A, C, or T) can be methylated (Law and Jacobsen 2010). After DNA methylation has been set up in H3K9me levels to enforce the transcriptional silencing in both animals and plants (Johnson et al. 2007; Bernavichute et al. 2008; Du et al. 2015). In Arabidopsis, the CHG methylation is highly correlated with H3K9me2 (Bernavichute et al. 2008; Du et al. 2012) owing to the establishment of a self-reinforcing loop between these two epigenetic marks. CMT3 is recruited by H3K9me2, which is deposited by KRYPTONITE (KYP) to methylate CHG; in turn, methylated CHG DNA recruits KYP to maintain H3K9me2 levels (Johnson et al. 2007; Law and Jacobsen 2010; Du et al. 2012). Histone methylation, including H3K9me, can be reversed by a JmjC domain-containing family of proteins [Klose et al. 2006]. Genes encoding 21 JmjC proteins have been identified in Arabidopsis, and several proteins have been characterized as active histone demethylases [Lu et al. 2008; Luo et al. 2014].

Transposable elements and repetitive sequences, which are major targets of DNA methylation and H3K9me, are maintained as transcriptionally silenced heterochromatin. Paradoxically, transcription from the silenced loci is required to initiate heterochromatin formation, and the regulation of this process is largely unclear [Grewal and Elgin 2007; Matzke and Mosher 2014; Holoch and Moazed 2015]. In fission yeast, a JmjC protein, Epe1, functions specifically to promote the transcription of silenced loci to dynamically maintain heterochromatin [Ayoub et al. 2003; Zofall and Grewal 2006; Trewick et al. 2007]. Recently, we reported that JMJ24 functionally resembles Epe1 and plays a conserved role in promoting basal-level transcription of silenced loci to reinforce the RNA-based silencing in Arabidopsis [Deng et al. 2015]. However, the molecular mechanism of Epe1 and JMJ24 to counteract H3K9me is still unknown, since both proteins harbor an atypical JmjC domain whose demethylase activity is unclear [Klose et al. 2006; Zofall and Grewal 2006; Baba et al. 2011; Deng et al. 2015]. Epe1 was assumed to be a protein hydroxylase that affects the stability of a heterochromatin protein, such as Swi6 or Clr4 methyltransferase, thereby regulating the extent and stability of heterochromatin domains [Trewick et al. 2007]. Arabidopsis JMJ24 harbors a RING motif, and we found that this protein has E3 ubiquitin ligase activity. We also demonstrated that JMJ24 ubiquitinated CMT3 in vitro and destabilized it in vivo. JMJ24 decondenses heterochromatin probably through degradation of CMT3.

Results and Discussion

JMJ24 is an E3 ubiquitin ligase

In addition to the JmjC domain, JMJ24 harbors a RING motif and a coiled-coil (CC) domain [Fig. 1A]. As RING
finger proteins are known to have ubiquitin E3 ligase activity (Joazeiro et al. 1999; Lorick et al. 1999; Xie et al. 2002), we examined whether JMJ24 has this activity in vitro. Figure 1B shows that a purified MBP (maltose-binding protein)-JMJ24 fusion protein was able to self-ubiquitinate, whereas JMJ25/IBM1, a close relative of JMJ24, was inactive (Supplemental Fig. S1). The E3 ligase activity of the JMJ24 was abolished when the RING domain was deleted (Fig. 1C). We further constructed two mutant proteins, Cys243 → Ser, Cys263 → Ser(C243S, C263S) and His244 → Ala, Cys263 → Ser(H244A, C263S), in which the RING motif was disrupted. In vitro ubiquitination assays showed that neither mutant protein possessed self-ubiquitination activity (Fig. 1D), suggesting that an intact RING domain is necessary for E3 activity of JMJ24. This is consistent with results obtained from other RING E3 ligases (Xie et al. 2002; Seo et al. 2003). Dimerization is known to activate RING E3 ligase activity, and the CC domain can function to mediate protein dimerization (Lupas 1996; Xie et al. 2002; Seo et al. 2003). To examine the role of the CC domain in JMJ24 dimerization, we expressed MBP and glutathione-S-transferase (GST) fused to wild-type JMJ24 [MBP-JMJ24 and GST-JMJ24] and CC domain-deleted JMJ24 [MBP-JMJ24ΔCC and GST-JMJ24ΔCC] in Escherichia coli. Figure 1E shows that JMJ24 can form dimers through the CC domain in vitro. We further confirmed that JMJ24 formed dimers in vivo by coimmunoprecipitation assay using Flag- or Myc-tagged proteins from plant extracts (Fig. 1F). Collectively, these results indicated that JMJ24 possesses properties typical of the RING E3 ligase.

**JMJ24 ubiquitinites CMT3 in vitro**

The E3 ligase activity of JMJ24 prompted us to hypothesize that JMJ24 may target components involved in heterochromatin formation for degradation. To examine this possibility, we performed a screen and found that JMJ24 interacted with CMT3 in yeast cells (Supplemental Fig. S2; Fig. 2A). Using purified proteins, we confirmed by
pull-down assays the direct association of JMJ24 with CMT3, but no interaction was seen between JMJ25/IBM1 and CMT3 (Fig. 2B), although the latter protein pair was reported to function in the same genetic pathway (Saze et al. 2008). We further examined whether CMT3 was a target of JMJ24 E3 ligase by in vitro ubiquitination assays. Indeed, CMT3 protein was ubiquitinylated by wild-type JMJ24 but not the JMJ24(H244A, C263S) mutant in the presence of E1 and E2 (Fig. 2C). The E3 activity of JMJ24 was inhibited by the RING motif mutant JMJ24(H244A, C263S) (Fig. 2D), consistent with a previous report that the RING mutant of the E3 ligase can function as a dominant-negative mutant (Xie et al. 2002; Seo et al. 2003).

**JMJ24 destabilizes CMT3 in vivo**

Next generated double-transgenic plants harboring 35S:Myc-CMT3 and a β-estradiol-inducible (Zuo et al. 2000) XVE:Flag-JMJ24 to determine whether JMJ24 would associate with CMT3 in vivo. Myc-CMT3, but not Myc-SGS3 (Deng et al. 2015), could be detected in the anti-Flag immunoprecipitates when Flag-JMJ24 was induced, but Myc-CMT3 was not detected in the absence of an inducer (Fig. 3A), indicating a specific association between JMJ24 and CMT3 in vivo. In Arabidopsis, Myc-CMT3 levels were reduced when Flag-JMJ24 was transiently over-expressed (Fig. 3B), with no change in CMT3 mRNA (Supplemental Fig. S3A), indicating that JMJ24 destabilized CMT3 in vivo. In contrast, CMT3 levels were increased upon induction of the dominant-negative mutant of JMJ24, JMJ24(H244A, C263S) (Fig. 3C; Supplemental Fig. S3B). Our results suggest that JMJ24(H244A, C263S) sequestered wild-type JMJ24 and stabilized CMT3. To further confirm that JMJ24 regulates CMT3 stability, we expressed 35S:Myc-CMT3 in wild-type Arabidopsis and the jmj24-null mutant. CMT3 protein was hardly detected in wild-type without the 26S proteasome inhibitor MG132. However, CMT3 was readily detected in the jmj24 mutant even without MG132 treatment with similar mRNA expression levels (Fig. 3D). A protein decay experiment showed that the half-life of CMT3 was increased in the jmj24 mutant compared with the Col-0 wild type (Fig. 3E, Supplemental Fig. S4). Taken together, these results indicate that JMJ24 destabilizes CMT3 in plants.

**JMJ24 regulates CHG methylation and H3K9me2 through CMT3**

Cytosines in all classes of sequence context (that is, CG, CHG, and CHH, where H = A, C, or T) can be methylated in plants. CMT3 is the main CHG methyltransferase in Arabidopsis, and CHG methylation is strongly depleted in cmt3 mutants (Lindroth et al. 2001; Law and Jacobsen 2010; Stroud et al. 2013). To determine whether JMJ24 regulates CHG methylation on a genome-wide basis, we performed Southern blot analysis with methylation-sensitive enzymes. HpaII and MspI are isoschizomers that recognize 5′-CCGG-3′ sequences with differential sensitivity to methylation. HpaII is inhibited by methylation of either cytosine of the recognition site, whereas MspI is sensitive only to methylation of the outer cytosine and thus detects CHG methylation. Using a highly repetitive Athila retrotransponson long terminal repeat (LTR) as a probe (Lindroth et al. 2001), we found that the Athila LTR showed slight but reproducibly greater resistance to MspI digestion in jmj24 mutants (Fig. 4A). In contrast, the digestion was increased in cmt3 mutants (Fig. 4A), consistent with previous reports (Lindroth et al. 2001; Jackson et al. 2002). These results indicate that the CHG methylation increased in jmj24 but decreased in cmt3. The increased CHG methylation in jmj24 was dependent on CMT3, since the MspI digestion pattern of jmj24;cmt3 double mutants was identical to cmt3 single mutants (Fig. 4A). No change of HpaII digestion was detected in either jmj24 or cmt3 single mutants or double mutants (Fig. 4A), indicating that neither JMJ24 nor CMT3 had any effect on CG methylation within the Athila LTR. The same results were also detected for two additional loci: 5s-rDNA repeats (Fig. 4B; Supplemental Fig. S5A; He et al. 2009) and centromere satellites (Supplemental Fig. S5B; Vongs et al. 1993). Sequencing of bisulfite-treated DNAs confirmed that the CHG methylation on a representative Athila LTR region (Lindroth et al. 2001) was increased by ~20% in jmj24 compared with wild-type but was abolished in either the cmt3 single or jmj24;cmt3 double mutant [Supplemental Fig. S5C]. The direct repeats present on the FWA promoter were methylated predominantly at CG sites, leading to the silencing of FWA in wild type (Kinoshita et al. 2007). We found the CG and CHH methylation levels remained unchanged, whereas CHG methylation

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**Figure 3.** JMJ24 destabilizes CMT3 in vivo. **(A)** JMJ24 associates with CMT3 in vivo. Double-transgenic seedlings carrying 35S:Myc-CMT3 and an estradiol-inducible XVE:Flag-JMJ24 were treated with (+) or without (−) β-estradiol followed by immunoprecipitation with anti-Flag antibody. Double-transgenic plants carrying Myc-SGS3 and Flag-JMJ24 were used as a control. **(B,C)** CMT3 protein levels in response to transient induction (+β-estradiol) of wild-type JMJ24 (B) or JMJ24(H244A, C263S) (C) in double-transgenic plants. **(D)** CMT3 was stabilized by the 26S proteasome inhibitor MG132. Myc-CMT3 expressed in wild-type or the jmj24 mutant was analyzed in the presence or absence of MG132 with an antibody against c-Myc. The bottom panel shows CMT3 transcript levels. **(E)** CMT3 protein decay rate in wild-type (top panel) or the jmj24 mutant (bottom panel). Samples were treated with cycloheximide (CHX) and collected at the indicated time points for immunoblot analysis using an antibody against c-Myc. Tubulin (TUB) was used a loading control in **B–E.**
was elevated by ~50% in jmj24 compared with wild type [Fig. 4C]. The CHG methylation was recovered to wild-type levels in JMJ24/jmj24 in which a wild-type functional JMJ24 was introduced to complement the jmj24 mutant. However, no recovery of the CHG methylation was seen in JMJ24/H244A, C263S) was used for complementation [Fig. 4C, Supplemental Fig. S5D,F]. Similar results were also obtained for QQS [Fig. 4D, Supplemental Fig. S5E], a de novo protein-coding gene located by both CG and non-CG methylation [Silveira et al. 2013].

CHG methylation colocalizes with H3K9me2, and these two epigenetic marks are interdependent on one another in Arabidopsis [Jackson et al. 2002; Law and Jacobsen 2010; Du et al. 2012]. Consistent with this, H3K9me2 levels were also increased in promoter regions

of FWA [Fig. 4E], QQS [Fig. 4F], and SDC [Fig. 4I] in jmj24 mutants in accordance with the elevated CHG methylation in the mutants [Fig. 4C,D; Deng et al. 2015]. Furthermore, a correlation between the CHG methylation/H3K9me2 and transcripts levels was also observed in these mutants for QQS [Fig. 4H] and SDC [Fig. 4I], two loci regulated by non-CG methylation. No significant changes for FWA transcript levels were detected for all mutants analyzed here [Supplemental Fig. S5G]; this is not surprising, as FWA is controlled by CG methylation [Lindoeth et al. 2001; Kinoshita et al. 2007]. Taken together, these results suggest that the RING motif plays an indispensable function for JMJ24, and JMJ24 regulates CHG methylation and H3K9me2 probably through degradation of CMT3.

Post-translational modification plays a critical role in regulating protein activity and stability. The latent H3K9me demethylase activity of a human homolog of Epe1/JMJ24, PHF2, which also harbors an atypical jmjC domain, was stimulated by phosphorylation [Baba et al. 2011]. Furthermore, ubiquitin-mediated degradation of DNA-associated proteins adds another regulatory layer to transcriptional regulation, which may be necessary to fine-tune expression patterns in response to environmental signals or developmental cues. Indeed, transcriptional regulators and chromatin-modifying enzymes tend to be unstable [Schwanhausser et al. 2011]. The putative H3K9me demethylase Epe1 was rapidly degraded by 26S proteasomes in order to define the proper heterochromatin boundary in yeast [Braun et al. 2011]. While it is not clear whether JMJ24 may reveal its demethylase activity after phosphorylation, our results uncover a novel mechanism of the JmjC protein to counteract H3K9me in plants. The RING finger was specifically acquired in the plant clade of JmjC histone demethylase, and this is consistent with the fact that CMT3 is also plant-specific. Our data here present a good example for both conserved (convergent evolution) and diversification mechanisms during evolution.

Materials and methods

DNA construction

The coding sequences of JMJ24, JMJ25/IBM1, and CMT3 were PCR-amplified from an Arabidopsis cDNA library, digested, and ligased into entry vector pENTR3C [Invitrogen]. Deletion variants of JMJ24 were generated by overlapping PCR, and point mutation mutants were generated with a QuickChange site-directed mutagenesis kit [Stratagene]. Entry vectors were recombined into pgEX-DC and pMAL-DC for E. coli expression or pBA-DC [SS5 promoter] and per-DC [XVE-inducible promoter] for plant expression [Zhang et al. 2005]. The JMJ24am complementation vector was generated with a QuickChange site-directed mutagenesis kit [Stratagene] from a previous wild-type complementation construct [Deng et al. 2015]. Primers are listed in Supplemental Table S1.

Plant material and growth conditions

All Arabidopsis used here were in the Col-0 background. The jmj24 and cmt3 mutants have been reported: jmj24-1 [GK-085H03] [Deng et al. 2015] and cmt3-11 [SALK_148381] [Chan et al. 2006]. The jmj24/cmt3 double mutant was generated by genetic crossing with single mutants. Transgenic plants were generated by the floral dip transformation method [Clough and Bent 1998]. Seeds were sterilized and grown on half-strength Murashige-Skoog (MS) medium for 2 wk before being transferred to soil. Plants were grown in a growth chamber at 22°C under 16-h light/8-h dark long-day photoperiod cycles.
Yeast two-hybrid assays

The Matchmaker GAL4-based two-hybrid system (Clontech) was used to perform yeast two-hybrid assays. The full-length JM24 coding sequence was inserted into the pG5TR9-GW vector (Clontech) to generate a binding domain (BD) #JM24 fusion (BD-JM24) construct. Coding sequences of HSK9 methyltransferases [SUVRs and SUVRs and DNMTrs (Supplemental Fig. S2)] were introduced into the pGAD424-GW vector (Clontech) to generate activation domain [AD] fusion constructs. Interactions between AD and BD fusions were assayed according to the manufacturer’s instruction.

Protein preparation, in vitro pull-down, and ubiquitination assays

The E. coli BL21 strain was used as a host for protein production. Recombinant proteins were purified using amyllose resin [New England Biolabs] or glutathione-Sepharose 4 fast flow [GE Healthcare] according to the manufacturer’s instructions. Interactions between fusion proteins were determined by pull-down as previously described (Jang et al. 2010). In vitro ubiquitination assays were performed as previously described (Xie et al. 2002; Jang et al. 2010). Each reaction mixture [30 µL] contained 100 ng of rabbit E1 [Boston Biochem], 100 ng of human E2 UbeHSH [Boston Biochem], 2 µg of ubiquitin (Sigma-Aldrich), and 500 ng of E3 [MBP-JM24] with or without 500 ng of substrate CMT3. Reactions were incubated for 1 or 2 h at 30°C followed by immunoblot detection.

In vivo coimmunoprecipitation

Two-week-old seedlings of Flag-JM24/Myc-JM24, Flag-JM24/Myc-jm24ΔCC, Flag-JM24/Myc-SC53, or XVE-Flag-JM24/35S:Myc-CMT3 [treated with or without β-estradiol for 16 h] were ground in liquid nitrogen and homogenized in immunoprecipitation buffer (50 mM Tris-HCl at pH 7.5, 100 mM NaCl, 0.2% Triton X-100, 1 mM DTT, 50 µM MG132, protease inhibitor cocktail [Roche]). After centrifugation, supernatants were taken for immunoprecipitation by anti-Flag M2 antibody (Sigma-Aldrich). Antibody-antigen complexes were harvested by protein G agarose (GE Healthcare) and homogenized in nucleoprotein buffer [50 mM Tris-HCl at pH 8.0, 100 mM NaCl, 0.2% Triton X-100, 1 mM DTT]. DNA digestion was performed using the RNase-free DNase set (Qiagen) and cleaned up using an RNeasy minikit (Qiagen). On-column DNA digestion was performed using the RNase-free DNase set (Qiagen) during the purification. cDNA was generated from RNA by using the SuperScript III first strand synthesis system [Invitrogen] according to the manufacturer’s instructions. Quantitative PCR was performed as previously described (Deng et al. 2015). Data are presented as mean ± SD (n = 3).

RNA analysis

Total RNA was extracted using Trizol reagent [Invitrogen] following the manufacturer’s instructions. RNA was treated with a Turbo DNase kit (Ambion) and cleaned up using an RNaseasy minikit (Qiagen). On-column DNA digestion was performed using the RNase-free DNase set (Qiagen) during the purification. cDNA was generated from RNA by using the SuperScript III first strand synthesis system [Invitrogen] according to the manufacturer’s instructions. Quantitative PCR was performed as previously described (Deng et al. 2015). Data are presented as mean ± SD (n = 3).

DNA methylation analysis

Genomic DNAs were extracted from 2-wk-old seedlings using a DNaseasy plant minikit (Qiangen). For Southern blots, 5 µg of genomic DNAs was digested with Hpal or Mspl (New England Biolabs) overnight. The digested DNA was loaded onto a 1.2% agarose gel and transferred to Hybond-N membranes (GE Healthcare). The Althia LTR, SS rDNA repeat, and 180-base-pair [bp] centromere repeat were labeled with α-32PdCTP for Southern hybridization to determine their DNA methylation status. Bisulfite DNA conversion was performed by using 0.5–1 µg of genomic DNAs and an Epitope bisulfite kit (Qiagen) following the manufacturer’s protocol. The purified DNAs were used as a template for PCR amplification, and the product was then cloned into the pGEM-T easy vector (Promega). For each sample, at least 15 individual clones were sequenced. All experiments were repeated at least twice using different biological samples.

Chromatin immunoprecipitation (ChIP)

ChIP was conducted as described by Gendrel et al. (2005) with minor modifications. Cross-linked chromatin pellets were resuspended in nucleus lysis buffer and sonicated in a Bioruptor UCD 200 (Diagenode) twice for 10 min at the maximum level. Samples were sonicated for periods of 30 sec with a 30-sec interval in between treatments. HSK9me2 antibody [Millipore, catalog no. 07-411] was used for immunoprecipitation. The purified DNA fragments were analyzed by quantitative PCR as described (Deng et al. 2015). The ChIP signal was first normalized against the input DNA and then to the wild-type value. Data are shown as mean ± SD (n = 3).

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References

Ayoub N, Noma K-i, Isaac S, Kahan T, Grewal SJ, Cohen A. 2003. A novel jmJC domain protein modulates heterochromatization in fission yeast. Mol cell biol 23: 4365–4370.

Baba A, Ohtake F, Okuno Y, Yokota K, Okada M, Imayi N, Ni M, Meyer CA, Igarashi K, Kanno J, et al. 2011. PKA-dependent regulation of the histone lysine demethylase complex PHF2-ARID5B. Nat Cell Biol 13: 668–675.

Bermatavicuiche YV, Zhang X, Cokus S, Pellegrini M, Jacobsen SE. 2008. Genome-wide association of histone H3 lysine nine methylation with CHG DNA methylation in Arabidopsis thaliana. PLoS One 3: e13156.

Braun S, Garcia JF, Rowley M, Rougemaille M, Shankar S, Madhani HD. 2011. The CUL4-DDB1 CDT2 ubiquitin ligase inhibits invasion of a boundary-associated antisilencing factor into heterochromatin. Cell 144: 41–54.

Cao X, Aufsatz W, Zilberman D, Mette MF, Huang MS, Matzke M, Jacobsen SE. 2003. Role of the DRM and CMT3 methyltransferases in RNA-directed DNA methylation. Curr Biol 13: 2212–2217.

Chan SW, Henderson IR, Zhang X, Shah G, Chien JS, Jacobsen SE. 2006. DNA, DRD1, and histone methylation actively target developmentally important non-CG DNA methylation in Arabidopsis. PLoS Genet 2: e83.

Clough SJ, Bent AF. 1998. Floral dip: a simplified method for Agrobacterium-mediated transformation of Arabidopsis thaliana. Plant J 16: 735–743.

Deng S, Xu J, Liu J, Kim S-H, Shi S, Chua N-H. 2015. JM24 binds to RDR2 and is required for the basal level transcription of silenced loci in Arabidopsis. Plant J 83: 770–782.

Du J, Zhong X, Bermatavicuiche YV, Stroud H, Feng S, Caero E, Vashishth AA, Terragini J, Chin HG, Tu A. 2012. Dual binding of chromomethylase domains to HSK9me2-containing nucleosomes directly DNA methylation in plants. Cell 151: 167–180.

Du J, Johnson LM, Jacobsen SE, Patel DJ. 2015. DNA methylation pathways and their crosstalk with histone methylation. Nat Rev Mol Cell Biol 16: 519–532.

Gendrel A-V, Lippman Z, Martienssen R, Colot V. 2005. Profiling histone modification patterns in plants using genomic tiling microarrays. Nat Methods 2: 213–218.
Deng et al.

Grewal SIS, Elgin SCR. 2007. Transcription and RNA interference in the formation of heterochromatin. *Nature* 447: 399–406.

He X-J, Hsu Y-F, Pontes O, Zhu J, Lu J, Bressan RA, Pikaard C, Wang C-S, Zhu J-K. 2009. NRPD4, a protein related to the RPB4 subunit of RNA polymerase II, is a component of RNA polymerases IV and V and is required for RNA-directed DNA methylation. *Genes Dev* 23: 318–330.

Holoch D, Moazed D. 2015. RNA-mediated epigenetic regulation of gene expression. *Nat Rev Genet* 16: 71–84.

Jackson JP, Lindroth AM, Cao X, Jacobsen SE. 2002. Control of CpNpG DNA methylation by the KRYPTONITE histone H3 methyltransferase. *Science* 296: 556–560.

Jang I-C, Henriques R, Seo H, Nagatani A, Chua N-H. 2010. Arabidopsis PHYTOCHROME INTERACTING FACTOR proteins promote phytochrome B polyubiquitination by COP1 E3 ligase in the nucleus. *Plant Cell* 22: 2370–2383.

Joazeiro CAP, Wing SS, Huang H-k, Levenson JD, Hunter T, Liu Y-C. 1999. The tyrosine kinase negative regulator c-Cbl as a RING-Type, E2-dependent ubiquitin-protein ligase. *Science* 286: 309–312.

Johnson LM, Bostick M, Zhang X, Kraft E, Henderson I, Callis J, Jacobsen SE. 2007. The SRA methyl-cytosine-binding domain links DNA and histone methylation. *Curr Biol* 17: 379–384.

Kinoshita Y, Saze H, Kinoshita T, Miura A, Soppe WJJ, Koornneef M, Kakutani T. 2007. Control of FWA gene silencing in Arabidopsis thaliana by SINE-related direct repeats. *Plant J* 49: 38–45.

Klose RJ, Kallin EM, Zhang Y. 2006. JmjC-domain-containing proteins and their functions in plants. *Plant Mol Biol Rep* 24: 558–565.

Lupo A. 1996. Coiled coils: new structures and new functions. *Trends Biochem Sci* 21: 375–382.

Matzke MA, Mosher RA. 2014. RNA-directed DNA methylation: an epigenetic pathway of increasing complexity. *Nat Rev Genet* 15: 394–408.

Saze H, Shrairashi A, Miura A, Kakutani T. 2008. Control of genomic DNA methylation by a JmjC domain-containing protein in Arabidopsis thaliana. *Science* 319: 462–465.

Schwanhauss B, Russe D, Li N, Dittmar G, Schuchhardt J, Wolf J, Chen W, Selbach M. 2011. Global quantification of mammalian gene expression control. *Nature* 473: 337–342.

Seo HS, Yang J-Y, Ishikawa M, Bolle C, Ballesteros ML, Chua N-H. 2003. LAF1 ubiquitination by COP1 controls photomorphogenesis and is stimulated by SPA1. *Nature* 423: 995–999.

Silveira AB, Trontin C, Cortijo S, Barau J, Del Bem L, Loudet O, Colot V, Vincentz M. 2013. Extensive natural epigenetic variation at a de novo originated gene. *PLoS Genet* 9: e1003437.

Stroud H, Greenberg MV, Feng S, Bematavitchute YV, Jacobsen SE. 2013. Comprehensive analysis of silencing mutants reveals complex regulation of the Arabidopsis methylome. *Cell* 152: 352–364.

Trewick SC, Mine E, Antonelli R, Urano T, Allshire RC. 2007. The JmjC domain protein Epe1 prevents unregulated assembly and disassembly of heterochromatin. *EMBO J* 26: 4670–4682.

Vongs A, Kakutani T, Martienssen R, Richards E. 1993. Arabidopsis thaliana DNA methylation mutants. *Science* 260: 1926–1928.

Xie Q, Gao H-S, Dullman G, Fang S, Weissman AM, Chua N-H. 2002. SINAT5 promotes ubiquitin-related degradation of NAC1 to attenuate auxin signals. *Nature* 419: 167–170.

Zemach A, Kim MY, Hsieh P-H, Coleman-Derr D, Eshed-Williams L, Thao K, Harmer SL, Zilberman D. 2013. The Arabidopsis nucleosome remodeler DDM1 allows DNA methyltransferases to access H1-containing heterochromatin. *Cell* 153: 193–205.

Zhang X, Garreton V, Chua N-H. 2005. The AIP2 E3 ligase acts as a novel negative regulator of ABA signaling by promoting ABI3 degradation. *Genes Dev* 19: 1532–1543.

Zofall M, Grewal SI. 2006. Swi6/HP1 recruits a JmjC domain protein to facilitate transcription of heterochromatic repeats. *Mol Cell* 22: 681–692.

Zuo J, Niu QW, Chua NH. 2000. An estrogen receptor-based transactivator XVE mediates highly inducible gene expression in transgenic plants. *Plant J* 24: 265–273.