Extensive \textit{de novo} activity stabilizes epigenetic inheritance of CG methylation in \textit{Arabidopsis transposons}

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Abstract

Cytosine methylation within CG dinucleotides (mCG) can be epigenetically inherited over many generations. Such inheritance is thought to be mediated by a semiconservative mechanism that produces binary present/absent methylation patterns. However, we show here that in *Arabidopsis thaliana* hiddm1 mutants, intermediate heterochromatic mCG is stably inherited across many generations and is quantitatively associated with transposon expression. We develop a mathematical model that estimates the rates of semiconservative maintenance failure and *de novo* methylation at each transposon, demonstrating that mCG can be stably inherited at any level via a dynamic balance of these activities. We find that DRM2 – the core methyltransferase of the RNA-directed DNA methylation pathway – catalyzes most of the heterochromatic *de novo* mCG, with *de novo* rates orders of magnitude higher than previously thought, whereas chromomethylases make smaller contributions. Our results demonstrate that stable epigenetic inheritance of mCG in plant heterochromatin is enabled by extensive *de novo* methylation.
Introduction

Cytosine methylation provides a mechanism to heritably alter the genome without permanent modification of the DNA sequence (M. Kim & Costello, 2017). DNA methylation represses transposable element (TE) transcription and transposition in plants and vertebrates (H. Zhang et al., 2018). Methylation also regulates endogenous genes: methylation close to the transcriptional start site generally causes gene silencing (Nuñez et al., 2021), whereas methylation of other genic regions can promote or counteract expression (Williams & Gehring, 2020). Genetic defects in the methylation machinery lead to human disease such as cancer (Baylin & Jones, 2016). Disruption of methylation patterns during clonal propagation of plants causes developmental defects that hamper agriculture, and methylation patterns within genes account for a substantial fraction of phenotypic variation in natural Arabidopsis thaliana populations (Lloyd & Lister, 2021; Ong-Abdullah et al., 2015; Shahzad et al., 2021). Faithful propagation of DNA methylation is clearly essential, yet our understanding of the underlying processes is far from complete.

Across eukaryotes, cytosine methylation is most prevalent within CG dinucleotides (Schmitz et al., 2019). The core model for the epigenetic inheritance of CG methylation (mCG) – proposed over forty years ago (Holliday & Pugh, 1975; Riggs, 1975) – was inspired by the symmetry of the CG site, and posits that the methylation status of the old strand is used to reproduce the pattern on the strand synthesized during DNA replication. This elegant, semiconservative model has strong experimental support (Catania et al., 2020; Gowher & Jeltsch, 2018; Jeltsch, 2006), but contains a potential flaw because it lacks a mechanism to recover DNA methylation following maintenance failure (Haerter et al., 2014).

Two classes of models have been proposed to address the above issue. The first, developed most explicitly with data from the fungus Cryptococcus neoformans, proposes that very high
fidelity semiconservative maintenance (estimated failure rate of $9.3 \times 10^{-5}$ per CG site per cell cycle) combined with rare, random and potentially non-enzymatic *de novo* methylation and natural selection can produce stable epigenetic inheritance of mCG over million-year timescales (Catania et al., 2020). The second, developed with mammalian data, deemphasizes the semiconservative mechanism, and instead proposes a combination of inefficient maintenance (*e.g.* failure rate of 8.0% per site per cell cycle; Wang et al., 2020) and high rates of untemplated *de novo* methylation (*e.g.* *de novo* rate of 4.7% per site per cell cycle; Wang et al., 2020) that is specifically targeted to methylated regions by an unknown mechanism (Haggerty et al., 2021; Lökvist et al., 2016; Wang et al., 2020). The maintenance methyltransferases in the two systems are different (*Dnmt5* in *C. neoformans* and *Dnmt1* in mammals; Dumesic et al., 2020; Huff & Zilberman, 2014), which might account for the distinct mechanisms. The overall lack of long-term, transgenerational epigenetic inheritance of mammalian mCG (Kazachenka et al., 2018; Seisenberger et al., 2013; Tucci et al., 2019) may also be compatible with higher rates of maintenance failure.

The existing plant data (mostly from *Arabidopsis*) appear to be more compatible with the former mechanism. Plants exhibit stable, transgenerational mCG inheritance (Becker et al., 2011; Quadrana & Colot, 2016; Schmitz et al., 2011), and although plants use MET1 (*a Dnmt1 family enzyme*; Tirot et al., 2021), the global maintenance failure rate of $6.3 \times 10^{-4}$ per generation (Van Der Graaf et al., 2015) or about $1.9 \times 10^{-5}$ per cell cycle (assuming 34 cell cycles per generation, Watson et al., 2016) reported for *Arabidopsis* is lower than that reported for *C. neoformans* ($9.3 \times 10^{-5}$, Catania et al., 2020). Even the higher maintenance failure rate of $4.4 \times 10^{-5}$ per cell cycle ($1.5 \times 10^{-3}$ per generation) reported for *Arabidopsis* genes (Van Der Graaf et al., 2015) is two times lower than the *C. neoformans* rate. Epigenetic inheritance of plant mCG is thought to depend only on MET1 (Cokus et al., 2008; Lister et al., 2008; H. Zhang et al., 2018), which, like *C. neoformans*
Dnmt5, does not have known de novo activity (H. Zhang et al., 2018). The reported rates of de novo mCG are also low in Arabidopsis (global rate of $2.6 \times 10^{-4}$ per generation or $7.6 \times 10^{-6}$ per cell cycle; Van Der Graaf et al., 2015), and the source of the de novo activity is unknown.

Plant and animal genomes also contain cytosine methylation outside CG dinucleotides (Du et al., 2015). In plants, the CMT3 methyltransferase family catalyzes methylation of CNG trinucleotides, conventionally described as CHG (where H is any non-G base) to avoid overlapping CG sites (Law & Jacobsen, 2010). The related CMT2 family can methylate cytosines outside CG and CNG contexts (Zemach et al. 2013; Stroud et al. 2014), a pattern of specificity referred to as CHH. CMT2 and CMT3 rely on a positive feedback loop with dimethylation of lysine 9 of histone H3 (H3K9me2) (Du et al., 2012, 2015; Rajakumara et al., 2011) – a hallmark of heterochromatin (Sequeira-Mendes et al., 2014) – and therefore preferentially methylate heterochromatic TEs (Stroud et al., 2014; Zemach et al., 2013; Y. Zhang et al., 2018). Plants also possess an RNA-directed DNA methylation (RdDM) pathway, in which 24 nucleotide RNA molecules guide DRM methyltransferases (homologs of animal Dnmt3) to initiate DNA methylation in all sequence contexts, and to maintain CHH methylation at relatively euchromatic TEs (Matzke & Mosher, 2014).

Regardless of the mechanism, plants do not perfectly copy non-CG methylation each cell division, which leads to a probabilistic distribution of methylation states at CHG and CHH sites that contrasts with the more binary mCG patterns (Cokus et al., 2008; Lister et al., 2008). Although the RdDM pathway can establish DNA methylation in every context, including CG (Chan et al., 2004; Cuerda-Gil & Slotkin, 2016; Pélissier et al., 1999; Teixeira et al., 2009), several studies have concluded that mCG maintenance is independent of the non-CG methylation pathways (Lister et al., 2008; Stroud et al., 2014; To et al., 2022). The de novo mCG rates reported for TEs (where the
non-CG pathways operate) are also lower than those for genes (Van Der Graaf et al., 2015), ranging down to 6.8×10⁻⁸ per cell cycle (2.3×10⁻⁶ per generation; Hazarika et al., 2022). Thus, inheritance of mCG within TEs is thought to be essentially semiconservative.

Previously, we described widespread intermediate mCG within heterochromatic TEs of Arabidopsis h1ddm1 mutants lacking the nucleosome remodeler DDM1 and linker histone H1 (Zemach et al. 2013; Lyons et al. 2017). The reduced mCG maintenance efficiency in h1ddm1 mutants combined with the reported low de novo rates in TEs (Hazarika et al., 2022; Van Der Graaf et al., 2015) imply that these intermediate mCG patterns should be unstable and eventually degrade to very low levels, but this has not been evaluated. Here we analyze h1ddm1 DNA methylation over ten generations of inbreeding. We find that, contrary to the above expectation, intermediate mCG patterns are stably inherited. Using a mathematical model, we find a predicted de novo component of mCG inheritance orders of magnitude stronger than implied by the previously reported transgenerational rates of mCG gain in TEs. This prediction is confirmed by analysis of h1ddm1 compound mutants lacking either CMT2, CMT3, or the key RdDM methyltransferase DRM2. Our results contradict the established view that mCG epigenetic inheritance within TEs is essentially semiconservative. Instead, our data indicate that epigenetic inheritance of heterochromatic mCG is stabilized by extensive de novo methylation, with RdDM contributing most of the de novo activity and CMT2/3 making smaller contributions.

Results

TE CG methylation decays to stable intermediate levels within ten h1ddm1 generations
To understand DNA methylation inheritance in h1ddm1 plants, we crossed a plant homozygous for mutations in both canonical Arabidopsis H1 genes to a heterozygous ddm1 plant from a line in which the ddm1 mutation had never been homozygous. The resulting h1.1/+;h1.2/+;ddm1/+ F1 was allowed to self-fertilize to generate an F2 founder in which all three mutant alleles were homozygous (and the ddm1 allele was homozygous for the first time). As DDM1 predominantly functions in heterochromatin (Zemach et al., 2013), we focused our analysis on heterochromatic TEs (hTEs) that lose much of their DNA methylation in all sequence contexts when this remodeler is inactivated (Fig. 1A-C and Fig. S1A-B). Consistent with the positive association between DNA methylation loss and H3K9me2 in ddm1 plants (Zemach et al., 2013), F2 h1ddm1 mCG varies substantially between hTEs and shows a strong negative correlation with H3K9me2 in the parental h1 genotype (r = -0.60, Fig. 1D). Unlike ddm1 where most mCG is lost by the F2 (Fig. S1C), average and median h1ddm1 mCG across hTEs decreases substantially from the F2 to the F3 generation (median change 11%), less so from the F3 to the F4 generation (~5%), and even less from the F4 to the F5 (Fig. 1A-C). Overall mCG remains stable at intermediate levels after the F5 generation, up to generation F11 (ten generations of inbreeding; Fig. 1A-C, Fig. S1D). Across all ten h1ddm1 generations analyzed, intermediate mCG spans vast swaths of heterochromatin (>12Mbp, ~10% of the genome), as can be appreciated from genome browser views (Fig. 1B). CG methylation patterns are well-correlated across generations (Fig. 1E), indicating that the heterogeneous mCG landscape of h1dddm1 hTEs is accurately reproduced.

CHG methylation (mCHG) behaves similarly to mCG over h1ddm1 generations, dropping substantially initially, then leveling off in the subsequent generations (Fig. S1A,E), whereas h1ddm1 CHH methylation (mCHH) decreases overall, though not monotonically (Fig. S1B,F). In both cases relative losses are much smaller than for mCG. Early generation mCHG and mCHH
patterns are poorly correlated with mCG, but the correlation improves over time (Fig. 1E), suggesting that methylation in all contexts evolves in concert across h1ddm1 generations. Although correlations are strongest between methylation patterns of the late generations, early generation mCHG and mCHH patterns are better correlated with late generation mCG than with early generation mCG (Fig. 1E). Therefore, to some extent, early generation non-CG methylation can predict late generation mCG. This observation suggests that the processes shaping mCG epigenetic inheritance in TEs are related to non-CG methylation.

**Intermediate mCG is caused by methylation fluctuations at individual CG sites**

A simple explanation for the observed intermediate mCG across h1ddm1 generations would be methylation heterogeneity across CG sites, with fully methylated and fully unmethylated sites producing an intermediate pattern when averaged in bins. In this scenario, the gradual decrease and stabilization of mCG in h1ddm1 would be caused by an increasing number of CG sites permanently switching to an unmethylated state, until only sites with stable mCG maintenance remain methylated around the F5 generation. For example, because DDM1 is more important for mCG maintenance in nucleosomes than in the connecting linker DNA (Lyons & Zilberman, 2017), the observed intermediate mCG patterns could be caused by stable mCG maintenance in linkers and unstable maintenance in nucleosomes. This should cause mCG in TEs with poorly positioned nucleosomes to stabilize at lower levels, because shifting nucleosome positions across cell cycles should prevent efficient maintenance at any CG site. However, this is not the case, as mCG behaves similarly across h1ddm1 generations in TEs with various levels of nucleosome positioning (Fig. S1G).
To test the broader possibility of bimodal CG site methylation heterogeneity, we plotted per-base methylation in wild type (WT), h1, ddm1 and h1ddm1 plants for all heterochromatic CG sites methylated above 5% in WT. The near-binary nature of mCG is evident in WT versus ddm1 plants, and even more so in h1 versus ddm1 mutants (Fig. 1F), because h1 enhances heterochromatic mCG (Fig. 1A-B; Lyons & Zilberman, 2017; Zemach et al., 2013). In contrast, F2 h1ddm1 hTE mCG is much more uniformly distributed, covering with substantial representation all methylation values from 0 to 1 (Fig. 1F). This pattern largely persists over the generations (Fig. S1H) and is correlated across generations (Fig. 1G and S1I). Therefore, CG sites that are typically consistently methylated across a WT population of cells exist in a mixture of methylation states in h1ddm1 cells.

Another possibility to generate intermediate mCG is methylation heterogeneity across entire loci (TEs, for example), with fully methylated and fully unmethylated loci producing an intermediate pattern when averaged. This scenario entails dynamic switching of locus methylation, with the fraction of loci that lose mCG increasing over the first few generations before stabilizing. To examine this possibility, we analyzed methylation of sequencing reads that correspond to individual DNA molecules, where locus-level variation should produce reads with either low or high methylation. When considering all heterochromatic reads with >3 CG dinucleotides in early, middle, and late generations, the observed distribution of per-read mCG is relatively uniform in h1ddm1 plants (Fig. 1H), similar to the per-CG h1ddm1 mCG (Fig. 1F), indicating that many individual DNA molecules have a mixture of methylated and unmethylated CG sites. This becomes apparent when mCG patterns of individual reads are visualized, with h1ddm1 reads exhibiting a mixture of mCG levels and patterns (Fig. 1I). These results demonstrate that the intermediate h1ddm1 mCG patterns do not arise from averaging of fully methylated and
unmethylated CG sites, or from entire loci dynamically switching their methylation states during development. Instead, our results indicate that individual CG sites regularly switch their methylation states, implying a substantial flux of de novo mCG in h1ddm1.

Although WT CG sites tend to have higher methylation levels than in h1ddm1, individual CG sites nonetheless show methylation variability between DNA molecules in WT (Fig. 1I). This is reflected in a broader distribution of per-read methylation levels in WT compared to h1 (Fig. 1H) and in a similarly broader distribution of per-CG methylation (Fig. 1F-G). This suggests that mCG epigenetic inheritance in WT heterochromatin also involves substantial de novo methylation.

**Intermediate CG methylation is heritable in h1ddm1 plants**

A possible explanation for the apparently stable inheritance of intermediate heterochromatic mCG in h1ddm1 plants is that we are assaying the wrong tissue. We measure DNA methylation in leaves, which do not contribute to the subsequent generation. Cells that mediate inheritance, such as reproductive cells, show more robust maintenance of mCG than leaves and other somatic tissues (Hsieh et al., 2016; Park et al., 2016). Therefore, the h1ddm1 genotype might primarily destabilize somatic, but not reproductive methylation maintenance. In this scenario, the intermediate mCG patterns we observe in leaves may be produced in each generation through methylation decay during somatic development from an initially high level expected to be compatible with primarily semiconservative inheritance. To test this hypothesis, we quantified sperm DNA methylation in the F4 and F5 generations of h1ddm1 plants and WT, h1 (only the F5 generation), and ddm1 siblings. WT plants have increased mCG in sperm compared to leaf (Fig. 2A-B), in agreement with previous work (Hsieh et al., 2016), whereas mCG is similarly low in ddm1 sperm and leaf (Fig. 2A,C). Sperm mCG is unchanged in h1 compared to WT (Fig. 2A-C), consistent with the low H1
levels in male reproductive cells (He et al., 2019). Additionally, we find a large reduction of mCHH (but not mCHG) in sperm compared to leaf in all genotypes (Fig. S2A-B), indicating that mCHH reprogramming in sperm (Calarco et al., 2012) does not require DDM1.

Importantly, h1ddm1 plants possess nearly the same average mCG in sperm and leaf in the F4 and the F5 generations (Fig. 2A-B), a phenomenon readily apparent in the genome browser (Fig. 2D). Furthermore, there is a high correlation of per-site mCG between sperm and leaf of the same generation (Fig. 2E, top) as well as sperm across generations (Fig. 2E, bottom). Similar to h1ddm1 leaves (Fig. 1F), h1ddm1 sperm per-site mCG shows a continuous distribution of methylation frequencies (Fig. 2F). Per-read sperm methylation patterns in h1ddm1 (Fig. 2G-H) are also similar to those of leaves (Fig. 1H-I), indicating a mixture of mCG levels and patterns in h1ddm1 sperm cells.

Because sperm cells directly initiate the next generation, our results demonstrate that patterns of intermediate methylation are indeed inherited. The extensive mCG heterogeneity between DNA molecules (Fig. 2G-H), which represents heterogeneity between the haploid sperm, indicates that widespread de novo methylation of CG sites mediates stable inheritance of heterochromatic mCG in h1ddm1 plants. Notably, per-read mCG heterogeneity is also apparent in WT sperm (Fig. 2G-H), suggesting an important role for de novo methylation in maintaining WT heterochromatic mCG.

**TSS methylation is quantitively associated with TE expression in ddm1 and h1ddm1 plants**

Because loss of DNA methylation is accompanied by widespread TE activation in ddm1 mutants (Miura et al., 2001; Tsukahara et al., 2009), we examined whether the increased levels of DNA methylation in h1ddm1 (compared to ddm1) are associated with reduced TE expression. After
assembling transcripts based on the Araport11 assembly (Pertea et al., 2015), which includes ddm1-specific TE transcripts (Panda & Slotkin, 2020), we quantified expression in WT, h1, ddm1 and h1ddm1 plants. Of the 1375 transcripts upregulated in ddm1, 1075 overlap heterochromatic TEs. 796 of these are bona fide expressed heterochromatic loci, based on their H3K9 methylation profiles and expression level (H3K9me1 or -me2 > 0.25 in WT, >2.5 TPM in ddm1; Fig. S3A).

Expression of these heterochromatic transcripts is generally reduced in h1ddm1 compared to ddm1 (Fig. 3A-B), with 303 of these loci significantly downregulated (Fig. S3B, qval<0.05). In contrast, expression of upregulated euchromatic ddm1 transcripts is not substantially altered in h1ddm1 (Fig. S3C).

TEs that are similarly expressed between ddm1 and h1ddm1 possess a striking, localized depletion of DNA methylation in all contexts near the transcriptional start site (TSS) in h1ddm1, so that their methylation levels approach those of ddm1 plants at the TSS (Fig. 3C-D). In both CG and CHG contexts, average methylation downstream of the TSS increases to an intermediate level within 1 kb (Fig. 3C-D), illustrating that, as in genes (X. Zhang et al., 2006; Zilberman et al., 2007), DNA methylation in the TE body is compatible with high levels of expression. In contrast, TEs with reduced expression in h1ddm1 compared to ddm1 show little or no methylation depletion near the TSS and overall non-CG methylation levels similar to WT (Fig. 3E-F), indicating that the increased levels of CG and non-CG methylation in h1ddm1 plants (compared to ddm1) attenuate TE expression. Considering ddm1 and h1ddm1 together, \( \log_2 \) of the expression level shows a strong negative correlation with mCG around the TSS of downregulated TEs (+/-500 bp; TEs longer than 2 kb; \( r = -0.60 \), Fig. 3G), suggesting that DNA methylation around the TSS quantitatively reduces expression. Taken together, our results support the hypothesis that intermediate mCG gives rise to intermediate TE expression at the cell population level.
TSS methylation correlates with TE expression in ddm1 x WT F1 plants

Some TEs continue to be highly expressed following ddm1 outcrossing to WT despite DDM1 restoration, whereas others are silenced (Teixeira et al., 2009). A recent study reported that TE silencing in F1 progeny of ddm1 and WT is not correlated with DNA methylation but is instead linked to deposition of the H2A.W histone variant (Osakabe et al., 2021). Analysis of the published TE expression categories indeed shows no methylation difference between silenced (inactive) and expressed (neutral) TEs in the F1 plants (Fig. 3H, left), but these categories also show little difference in TE expression (Fig. 3I and S3D). We therefore calculated expression values from the published raw RNA-seq data to identify 235 TEs that are at least 4-fold downregulated in the F1 plants and compared them to 225 TEs not downregulated in the F1 (Fig. 3H-I and S3E). Consistent with published results (Teixeira et al., 2009), we find a strong recovery of DNA methylation in all contexts around the TSS of re-silenced TEs, but not at expressed TEs (compare "Osakabe et al." to "reanalysis", Fig. 3H), as would be expected if TE expression is reduced due to increased DNA methylation.

Considering that loss of DDM1 disperses heterochromatic foci (Soppe et al., 2002), greatly reduces DNA methylation in all contexts (Vongs et al., 1993; Zemach et al., 2013), and strongly activates TE expression (Kato et al., 2004; Miura et al., 2001; Tsukahara et al., 2009), whereas loss of H2A.W does not substantially disperse heterochromatin, alter DNA methylation, or activate TE expression (Bourguet et al., 2021), reduced H2A.W occupancy in heterochromatin is not likely to be a primary cause of the ddm1 phenotype. Instead, our data indicate that TE activation in ddm1 mutants is caused by loss of DNA methylation, and reduced TE expression in h1ddm1 and ddm1 x WT plants is a consequence of increased methylation. As H2A.W occupancy within TE bodies
is anticorrelated with TE expression (Fig. 3J), transcription-coupled H2A.W eviction plausibly explains at least some of the observed H2A.W loss from *ddm1* heterochromatin (Osakabe et al., 2021).

**Morphological phenotypes are ameliorated in *h1ddm1* compared to *ddm1***

In addition to extensive disruption of heterochromatin, loss of DDM1 function causes CHG hypermethylation of genes and severe morphological phenotypes (Ito et al., 2015; Kakutani et al., 1995, 1996; Zemach et al., 2013). As heterochromatic methylation and TE silencing are partially restored in *h1ddm1* plants, we evaluated whether other *ddm1* defects are also ameliorated. We find that the widespread *ddm1* genic mCHG increase is largely abolished in *h1ddm1* (Fig. 4A-B). The *ddm1* morphological phenotypes are also markedly improved in *h1ddm1* plants, which have more rosette leaves with more total area, are not delayed in bolting, and grow taller and produce more siliques (fruit) than *ddm1* (Fig. 4C-D). We have propagated *h1ddm1* plants until generation F14, at which point they remain fertile and generally healthy (Fig. S4A). Thus, in addition to stabilized intermediate DNA methylation of heterochromatin, *h1ddm1* plants have a stable, relatively normal morphology in comparison to *ddm1*.

**Genic mCG declines across *ddm1* and *h1ddm1* generations***

Although DDM1 is primarily important for TE methylation, some genes, especially those with low levels of expression, lose mCG in *ddm1* plants (Fig. S4B) (Ito et al., 2015; Lyons & Zilberman, 2017). Unlike *ddm1* hTEs, in which mCG declines precipitously in the F2 generation and shows little change thereafter (Fig. S1C), these genes show a gradual transgenerational mCG decline (Fig. 4E). As in TEs, mCG in *ddm1*-affected genes is substantially higher in *h1ddm1* than in *ddm1* (Fig.
4E). Compared to TEs (Fig. 1C), genic mCG declines more slowly and stabilizes more gradually across h1ddm1 generations (Fig. 4E). The slower decline of genic mCG in ddm1 and h1ddm1 compared to TEs suggests that mCG maintenance is more robust in genes in the absence of DDM1. This is consistent with most genes retaining roughly WT mCG in ddm1 (Lyons & Zilberman, 2017), so that some loci (hTEs) have strongly compromised mCG maintenance, others (lowly expressed genes) have modestly reduced mCG maintenance, and others (most genes) have effectively normal mCG maintenance.

**Quantification of h1ddm1 mCG de novo and maintenance failure components**

The pattern of transgenerational mCG loss and stabilization in h1ddm1 resembles an exponential decay (Fig. 1C) that should be sensitive to the rates of both maintenance failure and de novo methylation. To characterize h1ddm1 mCG dynamics more precisely, we developed a mathematical model that describes the CG methylation reaction using three processes expressed as linear difference equations: de novo methylation per cell cycle (delta, \( \delta \)), maintenance failure per cell cycle (epsilon, \( \epsilon \)), and DNA replication, assuming 34 cell cycles per generation based on published estimates for long day conditions (Fig. 5A and Methods, Watson et al., 2016). Over many cell cycles, any given CG site can be affected multiple times by these processes, dynamically changing back and forth between methylated and unmethylated. Overall, each TE has an initial WT mCG level, \( M_0 \), and a lower steady-state mCG level, termed \( M^* \), that is approached after multiple generations \( n \) of inbreeding, such that \( M_n \rightarrow M^* \) for large \( n \). The two unknowns \( \delta \) and \( \epsilon \) can be derived on a per-TE basis by fitting the time-series methylation data to an exponential decay, with a timescale \( [(\delta+\epsilon)/2]^ {-1} \), that tends to \( M^*=\delta/(\delta+\epsilon) \) (Methods). Longer TEs (>2 kb) proved to be more amenable to modeling due to lower levels of noise. Excluding TEs with an
average $WT$ mCG $\leq 0.65$, the mCG trajectory of the vast majority of TEs $>2$ kb ($1617/1883$, 86%) could be reliably fit to an exponential decay (Methods). The fit to the generational averages is shown (Fig. 5B). Although many short TEs were excluded, the remaining TEs comprise ~80% of total heterochromatic sequence length in the *Arabidopsis* genome.

By individually modeling the above TEs, we find that *h1ddm1* de novo and maintenance failure rates are within comparable ranges and show an intriguing linear correlation of 0.51 ($\delta_{\text{median}}=0.022$, $\delta_{\text{mean}}=0.021$, sd=0.010; $\varepsilon_{\text{median}}=0.033$, $\varepsilon_{\text{mean}}=0.035$, sd=0.015) (Fig. 5C). We were unable to calculate $\delta$ and $\varepsilon$ for *ddm1* due to the very rapid loss of mCG in this genotype (Fig. 1A and S1C). However, we find that TEs that most readily regain mCG in *ddm1 x WT* F1 plants (Osakabe et al., 2021) have the highest $\delta$ values in *h1ddm1* (Fig. S5A, r=0.48), suggesting that the *h1ddm1* $\delta$ values are relevant for other genotypes.

By altering the values of $\delta$ and $\varepsilon$, essentially any steady-state methylation level can be achieved, as shown in the breadth of observed M* at individual TEs, ranging from near 0 to greater than 60% (Fig. 5D). Further, different $\delta$ and $\varepsilon$ combinations can achieve the same M*. To help visualize this, we overlaid lines on a plot of $\varepsilon$ vs. $\delta$ that correspond to given steady-state mCG states (M*) (green lines, Fig. 5C). These lines emphasize that a low $\varepsilon$ (efficient maintenance) is usually required for the $WT$ mCG level, and that even a small increase of the absolute maintenance failure rate can have a substantial impact on the steady-state methylation level. Thus, although the average *h1ddm1* $\varepsilon$ is only about 3%, this compounds over many cell cycles to produce a substantially lower steady-state mCG compared to $WT$ (Fig. 1A-C).

To appreciate the implications of this model, consider a TE comprised of 100 CG sites, beginning with 90% methylation, and where $\delta$ and $\varepsilon$ are equal to their *h1ddm1* mean values. De novo activity affects unmethylated CG sites (uCG), and therefore plays only a small role in the
initial cell cycles following the creation of the mutant state (generation 0, only 10 uCG, Fig. 5E). However, as time progresses, unmethylated sites increase in prevalence and the absolute contribution of \textit{de novo} methylation increases, whereas maintenance failure becomes relatively less important as the number of mCG sites decreases. A steady-state is approached around cell cycle 150 (corresponding to just over four generations), when the numbers of gains and losses are balanced, both converging to a constant equal to (on average) 0.64 CG sites gaining/losing per cell cycle (dashed line in Fig. 5E).

Different regimes of $\delta$ and $\varepsilon$ manifest as different steady-state levels and/or mCG trajectories (with faster or slower decays) (Fig. 5F-G). Efficient maintenance and low \textit{de novo} each cause a slower decay to the mCG steady-state, so that TEs with excellent maintenance and low \textit{de novo} fail to reach steady-state even after 10 generations (Fig. 5F-G, #3, n=146 TEs), whereas those with highest \textit{de novo} but poor maintenance converge to steady-state rapidly (Fig. 5F-G, #2, n=149 TEs). At the most mCG-depleted TEs, \textit{de novo} rates are low, with larger maintenance failure, and average mCG stabilizes at $\sim$10% (Fig. 5F-G, #4, n=17 TEs). At the other end of the spectrum, TEs with high $\delta$ and low $\varepsilon$ lose $\sim$20% of their WT mCG (Fig. 5F-G, #1, n=13 TEs). However, because $\delta$ and $\varepsilon$ are positively correlated (Fig. 5C), TEs with poor maintenance tend to have better \textit{de novo}, and vice versa, so that the above extremes (Fig. 5F-G, #1 and #4) are unusual. Although we could not fit genic methylation dynamics well enough to confidently quantify $\delta$ and $\varepsilon$, the more gradual decline of mCG in genes (Fig. 4E) than in TEs (Fig. 1C) is consistent with lower $\delta$ and $\varepsilon$ (less \textit{de novo} activity and better maintenance) in genes.

\textbf{Contributions of DRM and CMT methyltransferases to CG methylation in h1ddm1}
Our model indicates that the initial mCG decline is dominated by maintenance failure, whereas steady-state mCG has a strong dependence on de novo methylation (Fig. 5E-F). The correlation of late generation (steady-state) mCG with mCHG and mCHH (Fig. 1E) therefore suggests that the RdDM and/or CMT pathways that mediate mCHG/CHH contribute (directly or indirectly) to de novo mCG activity. Consistent with this, δ shows strong linear correlations with mCHG and mCHH (\( r_{\text{mCHG}}=0.66 \), \( r_{\text{mCHH}}=0.59 \)) (Fig. S5B-C). To elucidate the enzymatic origins of de novo mCG, we generated three h1ddm1 mutant lines, each lacking one of the three principal Arabidopsis non-CG methyltransferases: DRM2, CMT2 and CMT3. We inbred each line through the F7 generation and analyzed leaf methylomes as we did for h1ddm1. The h1ddm1drm2 and h1ddm1cmt2 lines were created in the same way as the initial h1ddm1 line: a plant homozygous for all mutations except ddm1 was allowed to self-fertilize to create the founder F2 in which the ddm1 mutation was homozygous for the first time. We were unable to generate h1ddm1cmt3 plants this way, and instead created an F2 h1ddm1(cmt3/+ ) plant (first generation of ddm1 homozygosity), which was allowed to self-fertilize to create F3-equivalent h1ddm1cmt3 plants (first generation of cmt3 homozygosity).

Average hTE mCG is greatly reduced in h1ddm1drm2 compared to h1ddm1, which becomes obvious after the F2 generation (Fig. 6A-B). Consistently, the overall morphology of h1ddm1drm2 plants resembles ddm1 more than h1ddm1 (Fig. S6A). In contrast, neither h1ddm1cmt2 nor h1ddm1cmt3 lines exhibit an overall mCG decrease compared to h1ddm1 (Fig. 6A-B), despite corresponding strong losses of mCHH and mCHG, respectively (Fig. S6B-C), and these lines morphologically resemble h1ddm1 (Fig. S6A).

Although the loss of CMT3 does not substantially reduce average heterochromatic mCG, we identified 86 TEs with mCG in h1ddm1cmt3 mutants comparable to h1ddm1drm2 (>50% mCG
decrease; p < 0.01, Fisher’s exact test, Fig. 6C-E and S6D). Because non-CG methylation promotes RdDM (Choi et al., 2021; Johnson et al., 2014; Liu et al., 2014), we suspected the loss of CMT3 might reduce RdDM activity at these loci. Indeed, mCHH (the hallmark of RdDM) is nearly eliminated at these TEs in F7 h1ddm1cmt3 plants, as it is in F7 h1ddm1drm2 (Fig. 6F-G, mCHG shown in Fig. S6E), indicating that RdDM activity depends on CMT3 at a subset of heterochromatin. These CMT3-dependent TEs also have reduced mCHH in F7 h1ddm1cmt2 plants (Fig. 6F-G), as well as reduced mCG (Fig. 6D-E), so that steady-state mCG is affected by all three of the tested methyltransferases. Notably, non-CG methylation at these TEs does not collapse immediately in the compound methyltransferase mutants, but can be similar to h1ddm1 in early generations, and tends to decrease in tandem with mCG (Fig. 6F and S6F-H). This is consistent with our observation that CG and non-CG methylation changes are correlated across h1ddm1 generations (Fig. 1E), and with the known dependence of non-CG methylation on mCG at some TEs (Choi et al., 2020; Choi et al., 2021; Y. Zhang et al., 2018). The CG and non-CG methylation decreases may therefore be mutually reinforcing. Together, these data demonstrate that DRM2 is a key contributor to intermediate mCG in h1ddm1 heterochromatin, but also that epigenetic inheritance of mCG involves integration of the RdDM, CMT and MET1 pathways, at least at some loci.

**DRM2 mediates most of the de novo mCG in heterochromatin**

Application of our discrete methylation model to the methyltransferase mutant data allows us to calculate the relative contributions of each methyltransferase to de novo and maintenance mCG (Fig. 7A-C). Surprisingly, we find that ε decreases for all three compound mutants compared to h1ddm1 (Fig. 7D), indicating maintenance becomes more efficient when these methyltransferases
are removed. This is likely related to the observation that F2 h1ddm1 mCG shows a strong negative correlation with H3K9me2 (Fig. 1D). As the initial mCG decline is dependent in large part on ε (Fig. 5E-F), ε should also correlate with H3K9me2, which is indeed the case (Fig. 7E). Because non-CG methylation and H3K9me2 vary in tandem (Stroud et al., 2014), our data suggest that the drm2, cmt2 and cmt3 mutations improve mCG maintenance by rendering TEs less heterochromatic (less non-CG methylation and H3K9me2) and therefore less dependent on DDM1.

The cmt compound mutants exhibit modest decreases in δ that are comparable to the decrease in ε. In h1ddm1cmt2, δ decreases by 33% (from 0.021 to 0.014) and ε decreases by 26% (from 0.035 to 0.026), and in h1ddm1cmt3, δ decreases by 43% (to 0.012) and ε decreases by 34% (to 0.023; Fig. 7D,F-G). The changes between δ and ε are correlated (Fig. S7A-C), resulting in a form of dynamic compensation that, in the case of cmt mutants, keeps steady-state mCG (which depends on the ratio of δ and ε) nearly the same as in h1ddm1 (Fig. 7H-J). However, in h1ddm1drm2, δ falls 75% (to 0.0053; Fig. 7F-G) but maintenance does not improve at a similar level (ε decreases by 23% to 0.027; Fig. 7D,G). The much greater relative decrease in δ causes greatly reduced steady-state mCG compared to h1ddm1 (Fig. 7H,K) and accordingly, TEs with both high δ and low ε are very uncommon in h1ddm1drm2 as compared to low δ and high ε (Fig. 7L-M, black and orange points, respectively). Our results indicate that although CMT2, CMT3 and DRM2 all participate in the epigenetic dynamics of heterochromatic mCG, DRM2 contributes the bulk of the de novo mCG activity.

De novo rate estimates are consistent between h1ddm1 and met1

Because we calculate TE de novo mCG rates in the h1ddm1 background, it is important to ascertain how relevant these rates are for WT, especially because loss of histone H1 substantially perturbs
RdDM activity (Choi et al., 2021; Papareddy et al., 2020). To address this issue, we calculated *de novo* rates in *met1* and *h1met1* mutants using published data (Choi et al., 2020). These mutants are assumed to lack maintenance activity (ε=1), and δ values can therefore be straightforwardly inferred based on measured steady-state mCG (δ=mCG, Methods). The *met1* and *h1met1* δ values are correlated with *h1ddm1* δ values (Fig. S7B-C, \( r=0.28 \) for *met1* and \( r=0.34 \) for *h1met1*), supporting the conclusion that *h1ddm1* δ values are relevant for other genotypes. Some TEs retain RdDM and CMT activity in *met1* and *h1met1* mutants (MET1-independent TEs), whereas others lose methylation in all sequence contexts (MET1-dependent TEs; Choi et al., 2020; Choi et al., 2021; Y. Zhang et al., 2018). As expected, *h1ddm1* δ values show similar distributions at these TE categories (Fig. 7N). However, the *met1* and *h1met1* δ values are much higher in MET1-independent TEs, where they are much more comparable to those for *h1ddm1* (Fig. 7N), as would be expected if *de novo* activity is mediated by the non-CG methylation pathways.

Bisulfite sequencing has an intrinsic error rate caused by incomplete chemical conversion of unmethylated cytosine, as well as PCR and sequencing errors (Krueger & Franke, 2012). This rate can be estimated based on the measured methylation frequency in the unmethylated chloroplast genome, and is 0.17% for both *met1* and *h1met1* (dashed lines in Fig. 7N). Because per-TE error rates will be distributed around 0.17%, *met1* and *h1met1* δ estimates that are close to this value should be treated as upper bounds, with real δ values potentially much smaller. Therefore, many MET1-dependent TEs may experience very low *de novo* rates in *met1* and *h1met1* plants (Fig. 7N, left panel). In contrast, MET1-independent TEs generally have δ values well above the error rate (Fig. 7N, right panel), with the average *met1* and *h1met1* δ values (0.013 for both) in MET1-independent TEs only 1.7-fold lower than that for *h1ddm1* in MET1-independent TEs.
Because RdDM activity at MET1-independent TEs is similar between met1 and WT (Choi et al., 2021), WT heterochromatic de novo rates should be in a comparable range.

**Discussion**

Despite the similarities between the reported rates of mCG change in *C. neoformans* (Catania et al., 2020) and *Arabidopsis* (Becker et al., 2011; Hazarika et al., 2022; Schmitz et al., 2011; Van Der Graaf et al., 2015), our data indicate that epigenetic inheritance of mCG in *Arabidopsis* TEs involves a strong de novo component comparable to mammalian systems (Haggerty et al., 2021; Lövkvist et al., 2016; Wang et al., 2020). The de novo activity is primarily mediated by RdDM, with smaller contributions from CMT2/3. Both of these pathways have self-reinforcing features that focus their activities on TEs (Choi et al., 2021; Du et al., 2014; Johnson et al., 2014; Liu et al., 2014; Rajakumara et al., 2011), thereby solving the problem of targeting de novo mCG to regions with existing methylation. We derive our de novo rates primarily from *h1ddm1* data, but they are in good agreement with the rates in *met1* mutants (Fig. 7N), and should therefore be a good estimate for WT de novo rates.

The de novo mCG rates we find in heterochromatin are about 1% per site per cell cycle ($\delta_{\text{mean}}=0.022$). These rates are at least three orders of magnitude higher than the overall de novo rate of $3.2 \times 10^{-4}$ per generation reported for TEs (Van Der Graaf et al., 2015) and the range of $2.3 \times 10^{-6}$ to $1.7 \times 10^{-4}$ recently reported within different TE-associated chromatin states (CSs; Hazarika et al., 2022). This large difference is likely accounted for by two related factors. First, CS rates were calculated using the AlphaBeta model (Shahryary et al., 2020) that produces a good fit for CSs associated with genes,
explaining up to 88.6% of the data variance in CS5 (Hazarika et al., 2022). However, the model performs less well with the TE-associated states CS30-36. AlphaBeta provides reasonable fits for CS30 (67.6% of variance explained) and CS31 (57% of variance explained), and these states have relatively high reported de novo rates (3.5×10^{-5} for CS30 and 1.7×10^{-4} for CS31). AlphaBeta does not handle other TE CSs well, ranging from 28.3% of variance explained for CS32 to effectively 0% of variance explained for CS36 (variance explained values are from Supplementary Table 2 of Hazarika et al.). This indicates that the AlphaBeta model is not reliable for these regions of the genome.

More generally, AlphaBeta and other published rate estimates rely on the assumption that the rates are low enough that all or nearly all changes are captured by measurements that are separated by one or several generations. This should be a good assumption for genes, but it is not for TEs. Per-generation de novo rates of 30% mean that the half-life of an unmethylated site is short (30% rate implies a half-life of about two generations), leading to underestimates of methylation failure rates, because CG sites that lose methylation are rapidly remethylated. This likely at least partially explains why the reported per-generation maintenance failure rate for genes (1.5×10^{-3}) is 100-fold higher than the rate for TEs (1.2×10^{-5}; Van Der Graaf et al., 2015). Calculation of de novo rates may be confounded by an even more serious issue. Because de novo rate calculations rely on ancestrally unmethylated CG sites, and the half-life of unmethylated sites is inversely related to the de novo rate, CG sites with unusually low de novo rates will dominate the calculation, leading to potentially very large rate underestimates. This might explain the 1000-fold difference between the reported per-generation de novo rate in TEs (3.2×10^{-4}; Van Der Graaf et al., 2015) and our estimate of around 3×10^{-1}. 


Our results indicate that the low levels of mCG divergence observed at plant TEs (Hazarika et al., 2022; Van Der Graaf et al., 2015) are not caused by especially low rates of methylation loss and gain. Instead, this phenomenon is caused by a very high de novo rate that is much greater than the maintenance failure rate, so that maintenance errors are rapidly corrected – an idea consistent with the earlier proposal that TE mCG is stabilized by an epigenetic process that favors methylation gain over methylation loss (Van Der Graaf et al., 2015). Our data indicate that stable epigenetic inheritance of TE mCG is ensured by a combination of DDM1-supported MET1 maintenance activity with the strong de novo activity of RdDM (with smaller contributions from the CMT pathway). Using the reported per-generation maintenance failure rate for genes (1.5×10⁻³) as a baseline, the de novo activity at TEs is strong enough to stabilize mCG in h1ddm1 despite a >100-fold increase in the maintenance failure rate, to ~4×10⁻¹ per generation (equivalent to 1 - 0.985³⁴, Fig. 5, per-cell cycle loss rate=ε/2, Methods). Importantly, there is no compelling reason to assume that maintenance efficiency is the same or similar between genes and TEs in WT plants. Maintenance efficiency at TEs could be higher than at genes, but – even with the support of DDM1 – may also be considerably lower, especially at heterochromatic TEs. One of the main conclusions of our study is that the high de novo rates in TEs enable stable epigenetic inheritance of mCG within a wide range of maintenance rates.

Involvement of the RdDM and CMT pathways implies that de novo mCG rates likely fluctuate during development, as substantial differences in the activities of these pathways have been described across cell types and tissues (Erdmann & Picard, 2020), including during Arabidopsis reproductive and embryonic development (Bouyer et al., 2017; Calarco et al., 2012; Ibarra et al., 2012; Kawakatsu et al., 2017; Long et al., 2021; Zhou et al., 2022). Embryonic development is characterized by a gradual rise of mCHH in TEs that peaks at the onset of
germination and is mediated by RdDM and CMT2 (Bouyer et al., 2017; Kawakatsu et al., 2017). The functional significance of this process has been unclear, and one of its consequences may be to reinforce mCG inheritance by restoring CG sites within TEs to a methylated state.

Our results also bear on the functionality of DDM1 and of plant heterochromatin more broadly. The remarkable alleviation of the *ddm1* phenotype (TE activation, decreased fecundity, increased gene body mCHG, etc.; Fig. 3A-F and 4A-D) that emerges in *h1ddm1* concomitantly with the restoration of heterochromatic DNA methylation illustrates the central importance of chromatin homeostasis in the regulation of organismal viability, and the crucial role of DNA methylation in heterochromatin homeostasis and function. This is consistent with the reports that the mere presence of sufficiently large segments of demethylated heterochromatin in genetically WT plants is sufficient for phenotypic disruption (Johannes et al., 2009; Kooke et al., 2015). Our data do not support the proposal that the *ddm1* phenotype, including TE activation, is primarily caused by loss of the H2A.W histone variant from heterochromatin (Osakabe et al., 2021) – a conclusion that is also inconsistent with the published phenotypes of *ddm1* and *h2a.w* loss-of-function mutants (Bourguet et al., 2021; Jeddeloh et al., 1999; Kakutani et al., 1995, 1996; Tsukahara et al., 2009; Zemach et al., 2013). Instead, our results indicate that *ddm1* phenotypes are primarily caused by loss of heterochromatic DNA methylation, and the ameliorative effects of H1 removal in the *ddm1* background stem directly from increased heterochromatic DNA methylation, especially steady-state mCG.
Materials and Methods

Biological materials and growth conditions

The *h1ddm1* line was previously described (Lyons & Zilberman, 2017). DDM1 is a potent regulator of methylation and its removal initiates the decay of mCG in *h1ddm1*. Therefore, to generate *h1ddm1* we self-fertilized *h1/+;ddm1/+* and isolated *h1;ddm1*. To generate compound *h1ddm1* methyltransferase mutants, we crossed *h1/h1;ddm1/+* (abbreviated *h1;ddm1/+*) to homozygous null *drm2-2* (SALK_150863C), *cmt2-3* (SALK_012874) and *cmt3-11* (SALK_148381C) mutants. We then isolated *h1;drm2;ddm1/+* and *h1;cmt2;ddm1/+* and self-fertilized these to generate each respective F2 *drm2* and *cmt2* compound mutant. For *cmt3* we were unable to obtain *h1;cmt3;ddm1/+* from the initial cross. We did find *h1;ddm1;cmt3/+* frequently however, and therefore used this to generate *h1ddm1cmt3*, but as a result were unable to generate an equivalent F2 for this line. *Arabidopsis* seeds were sown on soil, stratified for 4 days at 4 deg. C and transferred to controlled environment chambers where they were grown in 16h light / 8hr dark at 20 deg. C until tissue harvest.

Leaf DNA isolation and bisulfite library prep

DNA was isolated by pulverizing ~0.5 g of flash frozen rosette leaves of 4 week post-germination plants. ~100 mg of resulting powder was used to extract genomic DNA (gDNA) with the DNeasy plant mini kit (Qiagen cat. no. 69104) per manufacturer’s instructions. gDNA was subsequently sonicated with the Bioruptor Pico (Diagenode) to ~250 bp median fragment length using 10 cycles of 30 seconds on and off. Agencourt Ampure beads (referred to as “beads” henceforth, cat. no. A63881) were then used at 2X volume to purify the sheared DNA. Following ligation of
methylated Truseq sequencing adapters (Illumina) to sheared DNA, bisulfite conversion of DNA was carried out according to manufacturer’s protocol (Qiagen Epitect Kit, cat. no. 59104) except without using carrier RNA. DNA was purified twice with 1.2X beads and converted a second time to ensure complete bisulfite conversion of unmethylated cytosine. Libraries were constructed using NEBnext kits (NEB cat. no. E7645) or Nugen/Tecan Ovation Ultralow (cat. no. 0344NB-08) following the manufacturer’s instructions. NEB next indexing primers (cat. no. E7335S) were used for generating multiplexed libraries during PCR amplification.

**Sperm DNA isolation and bisulfite library prep**

Open flowers of F4 or F5 plants (which is here denoted as F4 or F5 sperm, respectively) were collected for pollen isolation in Galbraith buffer (45 mM MgCl2, 30 mM sodium citrate, 20 mM MOPS, 1% Triton-X-100, pH7.0) by vortexing at 2000 rpm for 3 min. Flower parts were removed by straining through a 40 micron filter. Pollen grains were obtained by centrifugation at 2600 g for 5 minutes and broken down with glass beads (Sigma). The lysate was then transferred to a 40 micron cell strainer and the flow-through was centrifuged at 800 g for 10 min at 4°C. The pellet was re-suspended in Galbraith buffer and stained with SYBR Green for fluorescence-activated cell sorting (FACS). Vegetative nuclei and sperm nuclei (SN) were separated and collected based on size and fluorescence intensity. DNA was extracted from SN with ChargeSwitch gDNA Micro Tissue Kit (ThermoFisher cat. no. CS11203) following the manufacturer’s instructions. Briefly, SN were lysed in lysis buffer with proteinase K provided by the kit in a 55°C water bath overnight. RNA was digested with RNase A at room temperature for 30 minutes, then DNA was captured with magnetic beads provided, washed, and eluted. After quantification by fluorometry, about 10-20 ng of DNA was used for bisulfite-sequencing library preparation with Ovation Ultralow.
Methyl-Seq DR Multiplex System (Nugen part no. 0336) following the manufacturer’s instructions. Briefly, DNA was sonicated with Bioruptor sonicator, end repaired, ligated to adapters which contain indexes, and bisulfite converted twice with EpiTect Fast DNA Bisulfite Kit (Qiagen 59802) following the manufacturer’s instructions. After PCR amplification, libraries were purified with Agencourt RNAClean XP Beads.

Leaf RNA isolation and library preparation

Leaves (as above) from the F3 generation were flash frozen in liquid nitrogen, pulverized with mortar and pestle on dry ice, and the resulting material was subjected to vortexing in Trizol (Invitrogen, cat. no. 15596-026). Chloroform was then added at one-fifth the total volume and further vortexing was carried out until the solution appeared homogenous. RNA was subsequently pelleted in ice-cold isopropanol. The resuspended RNA was subjected to rRNA removal with Ribo-zero plant kit (Illumina, MRZPL1224) according to the manufacturer’s protocol. 50ng ribo-depleted RNA was used for library preparation with the Scriptseq kit (Epicentre, cat. no. SSV21124) following the manufacturer’s protocol but with the following modifications: the RNA fragmentation step was extended to 10 minutes, and the temperature was increased to 90°C.

Sequencing

For h1dml bisulfite libraries and some control and compound h1dml methyltransferase libraries, we used the Illumina HiSeq 2500 at the QB3 Vincent Coates Genomic Sequencing Lab at UC Berkeley. h1dml compound mutants were mostly sequenced on the NextSeq 500 (Illumina) at the John Innes Centre. Sperm bisulfite libraries were sequenced at the Bauer Core Facility at Harvard University with Illumina HiSeq 2000.
Short read mapping and quantification

Bisulfite libraries were mapped to the genome with BSMAp (Xi & Li, 2009) for all analyses, except single-read analyses, for which we used Bismark (Krueger & Andrews, 2011). BSMAp output was converted to per-base methylation scores with BSMAp’s methratio.py script. RNA was mapped to the genome with Hisat2 (D. Kim et al., 2019), using default settings except "dta" was on. Transcripts were assembled using Stringtie (Pertea et al., 2015) using Araport11 (March 2021 release; available as Araport11_GTF_genes_transposons.Mar202021.gtf at www.aradibopsis.org) as a guide, as we noticed that ddm1 mutant transcripts added in this release (Panda & Slotkin, 2020) were also present in our ddm1 data. For subsequent RNA quantification, we used our merged assembled transcripts to generate indices for use with kallisto pseudomapping (Pimentel et al., 2017) using the following settings for the quant program: --single --fr-stranded -b 100 -l 320 -s 30.

Statistics and visualization of experimental data

Methratio.py output was converted to GFF and further processed and analyzed using commands and workflows outlined in https://github.com/dblyons/modeling_h1ddm1/.

Kallisto output was fed into the R environment for processing and fitting with the Sleuth software package, which was used to perform likelihood ratio tests to derive q-values (i.e. FDR-adjusted p-values) for each TE per genotype comparison, as outlined here: https://rawgit.com/pachterlab/sleuth/master/inst/doc/intro.html.

The ends-analysis.pl script (https://zilbermanlab.net/dzlab-tools-1-5-81-linux-tar/) was used to generate enrichment score matrices of mapped bisulfite data around genomic features of interest. These matrices were imported to R (http://www.R-project.org) for further processing and
visualization using code available at www.github.com/dblyons/R. Both base functions and the ggplot2 library (Wickham 2009) were used for all plots except for heatmaps in Fig. 1, Fig. 2 and Fig. S1, and genome browser images. Heatmaps were generated with pheatmap (https://cran.r-project.org/web/packages/pheatmap/index.html). Built-in R function ‘cor’ was used for calculating Pearson’s correlation. Genome tracks are screenshots of our data were displayed in Integrated Genome Viewer (Robinson et al., 2011). For RNA-seq, we used bedtools genomecov (Quinlan & Hall, 2010) with scaling on to convert Hisat2 bam output to scaled browser tracks with units in reads per million mapped (RPM). For determining MET1-dependency of mCHH, we used the same method as in (Choi et al., 2021) such that: TE mCHH > =5% in WT and > =5% in met1 is MET1-independent, while TEs fitting the following criteria are MET1-dependent: TE mCHH > =5% in WT and < 2% in met1.

Plant leaf area measurements

Flats of the four different genotypes of plants were arranged on a platform to align images. Each plant was segmented manually using ImageJ (Collins, 2007), and then analyzed using ImageJ's mask feature to isolate leaves; the measure feature was used to measure all visible leaf area. Photos were taken using a Canon Rebel T3i.

Reanalysis of data from (Osakabe et al., 2021)

To determine whether DNA methylation changes occur at the promoters of TE genes that are significantly repressed in the ddm1 x Col-0 hybrid, we used the expression values provided in source data for Figure 5 (https://www.nature.com/articles/s41556-021-00658-1#Sec36, Osakabe et al., 2021) to group TE genes as ddm1 active, and if active, downregulated (inactive) or similarly
expressed (neutral) in the ddm1 x Col-0 hybrid F1. We then performed new transcript abundance analysis using kallisto and sleuth (as described above) without using any clustering as was done in (Osakabe et al., 2021), and then generated TE groupings based on the ratio of hybrid/ddm1 TPM as follows (as shown in Fig. S3E): downregulated < 0.25 and q-val < 0.05 (n=253); neutral > 0.3 (n=450). For mapping of methylation at these groups of TEs, we calculated the difference of observed methylation in the hybrid minus the mean of the parental lineages (Col-0 + ddm1) / 2 as 50 bp bins across the relevant loci, and then doubled this number to account for the dilution of the sample by the WT alleles present in the sample. We used the same calculation for estimating mCG gained at the ddm1 allele in Fig. S5A, except for the doubling step. For comparison of ddm1 H2A.W with TE expression, we used H2A.W ChIP-seq processed data from GSE150436 and plotted these against our recalculation of ddm1 TE gene expression, which was remarkably similar to the published results (Osakabe et al., 2021). For the calculation of gained mCG in the hybrid shown in Fig. S5A, we used the difference of observed methylation in the hybrid minus the mean of the parental lineages (Col-0 and ddm1) from (Osakabe et al., 2021).

Transposable element annotations

Heterochromatic TEs used for methylome analysis here are the same as in (Choi et al., 2020). For ddm1 mutant transcriptional analysis shown in Figure 3, we use the assembled transcripts from our analysis outlined above, which closely matches that of the March 2021 Araport11 gene and transposon release. These annotations, as well as genome coordinate files of the modeled TEs, TEs upregulated in ddm1, and the Osakabe et al. reanalysis annotations are available at www.github.com/dblyons/annotations_2022.
Mathematical model for TE methylation dynamics: constructing the recursion relations

We model the transgenerational methylation dynamics of CG sites within TEs. Each CG site is defined to be in one of three states: fully unmethylated, hemi-methylated or fully methylated. Recursion relations are used to express the methylation status of the TE at the end of each cell cycle such that:

\[ U^{(n)} = \text{fraction of sites unmethylated at the end of the } n^{th} \text{ cell cycle} \]

\[ H^{(n)} = \text{fraction of sites hemi-methylated at the end of the } n^{th} \text{ cell cycle} \]

\[ M^{(n)} = \text{fraction of sites methylated at the end of the } n^{th} \text{ cell cycle} \]

where \( U^{(n)} + H^{(n)} + M^{(n)} = 1 \).

As each CG site contains two cytosines, our model relates to the experimentally measured methylation level, \( \langle m_C \rangle \), via

\[ \langle m_C \rangle = \frac{2M^{(n)} + H^{(n)}}{2M^{(n)} + 2H^{(n)} + 2U^{(n)}} = \frac{2M^{(n)} + H^{(n)}}{2(M^{(n)} + H^{(n)} + U^{(n)})}. \]

To derive the recursion relations, we consider three key processes during the cell cycle: a) replication, b) maintenance methylation and c) de novo methylation and assign a probability for the various possible transitions between states. In addition, we assume that there is no active demethylation (selection of appropriate targets in A. thaliana where this criterion holds is discussed in a later section below). In met1 mutants, maintenance is completely compromised, whereas as in the various h1ddml mutants, it is only partially compromised. We therefore consider these two cases separately beginning with the h1ddml mutants.

Replication occurs in a well-defined and relatively narrow time-window of the cell cycle. For the purposes of our model, we define replication to be the start of a new cell cycle, as this is the point that new DNA is synthesized. Instead of modelling the top and bottom strands of DNA specifically,
we make the standard assumption that 50% of hemi-methylated sites become unmethylated upon replication:

\[ P(M \rightarrow H) = 1 \]
\[ P(H \rightarrow H) = \frac{1}{2} \]
\[ P(H \rightarrow U) = \frac{1}{2} \]
\[ P(U \rightarrow U) = 1. \]

After replication, maintenance methylation by MET1 occurs semi-conservatively such that \( H \rightarrow M \). We assume that maintenance occurs both rapidly and very efficiently after replication. Indeed, for all various \( h1ddm1 \) mutants, we find efficient maintenance to hold self-consistently (see below). We therefore re-write \( r_{\text{maint}} \), the contribution from the MET1-maintenance pathway, as \( r_{\text{maint}} = 1 - \epsilon \) where \( 0 \leq \epsilon \ll 1 \) represents the probability of maintenance failure. This gives:

\[ P(M \rightarrow M) = 1 \]
\[ P(H \rightarrow M) = 1 - \epsilon \]
\[ P(H \rightarrow H) = \epsilon \]
\[ P(U \rightarrow U) = 1. \]

Finally, we consider the \textit{de novo} pathways and define \( P(U \rightarrow H) = 2 \ r_{\text{novo}} = \delta \), where the factor of 2 is included as there are two possible cytosine targets in each CG site. In the mutants that we study, \( 0 < \delta \ll 1 \) is also found to hold self-consistently (see below). We neglect the possibility that \textit{de novo} methylation could occur twice at the same CG site (i.e., a \( U \rightarrow M \) transition) within a
single cell cycle, as this process is of order $\delta^2$. We also neglect the possibility that a de novo event could follow a maintenance failure, as this composite process is of order $\epsilon \delta$. The above assumptions are consistent with de novo methylation happening either concurrently with maintenance, or as an extended process throughout the cell cycle.

c) De novo:

$P(M \rightarrow M) = 1$

$P(H \rightarrow H) = 1$

$P(U \rightarrow H) = \delta$

$P(U \rightarrow U) = 1 - \delta$.

Combining the above three processes then provides the recursion relations:

$$M^{(n+1)} = (1 - \epsilon)(M^{(n)} + H^{(n)}/2)$$

$$H^{(n+1)} = \epsilon(M^{(n)} + H^{(n)}/2) + \delta(H^{(n)}/2 + U^{(n)})$$

$$U^{(n+1)} = (1 - \delta)(H^{(n)}/2 + U^{(n)}).$$

Using $U^{(n)} + H^{(n)} + M^{(n)} = 1$ to eliminate $H^{(n)}$ gives:

$$M^{(n+1)} = (1 - \epsilon)(1 + M^{(n)} - U^{(n)})/2$$

$$U^{(n+1)} = (1 - \delta)(1 - M^{(n)} + U^{(n)})/2.$$ 

This pair of equations can be rescaled and added to provide:

$$\frac{M^{(n)}}{(1 - \epsilon)} + \frac{U^{(n)}}{1 - \delta} = 1,$$

which allows the recursion relations to be expressed in terms of a single variable only, where we retain only terms linear in $\epsilon$ and $\delta$:

$$M^{(n+1)} = \frac{\delta}{2} + \left(1 - \frac{\delta + \epsilon}{2}\right)M^{(n)} \quad (1)$$
Next, we consider the met1 mutation. The above derivation is unchanged for replication. We assume complete maintenance failure, so that $\epsilon = 1$. As previously, we neglect processes of order $\delta^2$ within a single cell cycle, and later find that $0 < \delta \ll 1$ also holds self-consistently for met1 mutants. Applying these modifications provides the following recursion relations for $n > 0$:

$$\begin{align*}
M^{(n)} &= 0 \\
H^{(n+1)} &= H^{(n)} / 2 + \delta (H^{(n)} / 2 + U^{(n)}) \\
U^{(n+1)} &= (1 - \delta)(H^{(n)} / 2 + U^{(n)}).
\end{align*}$$

Using $H^{(n)} + U^{(n)} = 1$ then gives:

$$\begin{align*}
H^{(n+1)} &= \frac{1 - \delta}{2} H^{(n)} + \delta \\
U^{(n+1)} &= \frac{1 - \delta}{2} U^{(n)} + \frac{1 - \delta}{2}.
\end{align*}$$

**Mathematical model for TE methylation dynamics: steady state solutions**

At steady state, we define $M^{(n+1)} = M^{(n)} = M^*$, and equivalently for $H$ and $U$. For the various $h1ddm1$ mutants, we then find that to lowest order

$$\begin{align*}
M^* &= \frac{\delta}{\delta + \epsilon} \\
H^* &= 0 \\
U^* &= \frac{\epsilon}{\delta + \epsilon}.
\end{align*}$$

For all mutants considered here, except for met1, the values we extract from the data indeed give $\epsilon, \delta \ll 1$, thereby self-consistently justifying the assumptions made in our derivation. Hence,

$$\langle mC \rangle^* = \frac{H^* + 2M^*}{2(M^* + H^* + U^*)} = M^* = \frac{\delta}{\delta + \epsilon},$$

(2)
Whereas, for \textit{met1} mutants, to lowest order:

\begin{align*}
M^* &= 0 \\
H^* &= 2\delta \\
U^* &= 1 - 2\delta,
\end{align*}

for which,

\[ \langle m_C \rangle^* = \frac{H^* + 2M^*}{2(M^* + H^* + U^*)} = \frac{H^*}{2} = \delta. \quad (3) \]

\textit{Mathematical model for TE methylation dynamics: time dependent solutions}

Using a standard approach to solve difference equations, the recursion relation in Eq. (1) can be solved to express \( M^{(n)} \) as a function \( \delta, \epsilon \) and the initial fraction of methylation, \( M^0 \):

\[ M^{(n)} = \left( M^0 - \frac{\delta}{\delta + \epsilon} \right) e^{-\left| \ln\left(1 - \frac{\delta + \epsilon}{2}\right) \right| n} + \frac{\delta}{\delta + \epsilon}, \]

for \( 0 < \delta \ll 1 \) and \( 0 < \epsilon \ll 1 \), and therefore where we again retain only leading order terms. To allow comparison with the experimentally measured methylation level, the above exponential decay must be expressed as a function of the number of plant generations, \( x \), and the number of cell cycles through the germline per plant generation, such that \( n = n_{cc}x \). The experimentally measured methylation level then becomes:

\[ \langle m_C(x) \rangle = M^{(n)} = \left( M^0 - \frac{\delta}{\delta + \epsilon} \right) e^{-\left(\frac{\delta + \epsilon}{2}\right)n_{cc}x} + \frac{\delta}{\delta + \epsilon}, \quad (4) \]

with the free parameters: \( \delta \), the \textit{de novo} methylation rate for each unmethylated CG site per cell cycle, \( \epsilon \), the maintenance failure rate for each CG site per cell cycle, and \( M^0 \), the initial methylation level before the system was perturbed away from steady state.
Mathematical model for TE methylation dynamics: gain and loss rates per cell cycle

The overall methylation gain-rate per cell cycle (to first order) is:

\[ P(U \rightarrow M) = P(U \rightarrow H)_{\text{de novo}} \times P(H \rightarrow H)_{\text{replication}} \times P(H \rightarrow M)_{\text{maintenance}} \]

\[ = \frac{\delta}{2}. \]

This is referred to in the main text as the ‘de novo rate’.

Similarly, the overall loss-rate per cell cycle (referred to in the main text as the ‘maintenance failure rate’ to emphasize that this is a passive loss of methylation) is:

\[ P(M \rightarrow U) = P(M \rightarrow H)_{\text{replication}} \times P(H \rightarrow H)_{\text{maintenance}} \times P(H \rightarrow H)_{\text{de novo}} \times P(H \rightarrow U)_{\text{replication}} \]

\[ = \frac{\epsilon}{2}. \]

Relationship between the mathematical model and bisulfite sequencing data from leaf tissue

Our model is applied to individual heterochromatic TEs with a length > 2 kb. As the model assumes the absence of active demethylation, we wish to exclude possible ROS1 targets from our analysis. These are known to have depleted methylation levels in WT plants (Lister et al., 2008; Tang et al., 2016; Zhu et al., 2007). Therefore, we select only those TEs with a WT methylation level of \( M_{WT} > 0.65 \) for both replicates. This results in 1883 TEs which we fit using the model.

The transgenerational bisulfite sequencing data of leaf tissues show decreasing methylation levels as a function of time for the following mutants: \( h1ddm1 \), \( h1ddm1cmt2 \), \( h1ddm1cmt3 \) and \( h1ddm1drm2 \). We fit these methylation timeseries to the decaying exponential in Eq. (4) (the numerical details are explained in the following section). Consequently, we need to know the effective timepoints at which the leaf-tissue sequencing data is obtained, as described and illustrated schematically below.
We assume that the heterozygous plant methylation level is at its steady state and define $n = 0$ to occur when the homozygous offspring is formed, coinciding with (we assume) an abrupt potential change in the values of $\delta$ and $\epsilon$. The changed $\delta$ and $\epsilon$ values will now correspond to a different steady-state methylation level. The model predicts that the methylation level of the mutant plant will decay exponentially with time towards this new value. To fit the experimental data to this exponential decay, we first need to know the approximate number of replication events between data points.

Following (Watson et al., 2016), we assume a constant number of cell divisions through the cell lineage leading to the generative lineage ("germline"), $n_{cc} = 34$. We also assume a constant number of somatic cell divisions to produce the sequenced leaf tissue: $n_{st}$, this value, however, is unknown. Under these assumptions, for plants of generation F2 and later, there is a uniform number of cell divisions between each generation of $\Delta n = n_{cc}$. The heterozygous plant,
however, is more problematic as it is assumed to stay at its original steady-state methylation level up until the formation of the homozygous zygote. The F2 generation is therefore separated by $\Delta n = n_0 + n_{st}$, where $n_0$ is the number of replications before the leaf tissue branches from the germline, both values of which are unknown.

The data points for early generations are particularly valuable as the methylation level changes most rapidly between these. We treat biological replicates of the same generation as multiple data points with the same time-value. As an initial test of the model, we first fit to the methylation level averaged over all the selected TEs (Fig. 5B). Here all generations from F2 onwards are included. We find excellent fits to both the $h1ddm1$ and the $h1ddm1drm2$ datasets, along with a good fit for $h1ddm1cmt2$. However, in the latter case, the replicates have a greater spread, increasing the fit uncertainty. Furthermore, for $h1ddm1cmt3$, the important F2 generation is not available significantly reducing our confidence in this fit.

To partly overcome this difficulty, we use our other data sets to generate an extra effective datapoint corresponding to the WT plant. As demonstrated by the fit to the average mCG level for $h1ddm1$ F2 to F11 replicates (fit without WT: Fig. 5B solid line), the mCG level clearly decays from a high initial value at $x = 0$ ($\sigma_{fit} = 0.011$). By projecting the WT leaf methylation level onto the $h1ddm1$ fit without WT, we find (coincidentally) that the WT leaf methylation level corresponds to $x = 0$ (to within the fit uncertainty). For each individual TE, therefore, we approximate the initial methylation level (at $x = 0$) by the two WT leaf replicates and confirm that the fit to average methylation levels when including these two extra effective datapoints (fit with WT: Fig. 5B dashed line) is highly consistent to the fit without WT. We then fit to the methylation timeseries for individual TEs to generate distributions of $\delta$ and $\epsilon$ values for each mutant (described in detail below). We confirmed that fits excluding the WT leaf replicates at $x = 0$ provide
comparable distributions, however, manual inspection revealed these to provide less reliable estimates of the decay constant than the fits with the WT leaf replicates included (due to some fits generating anomalously low initial methylation levels and consequently unreliable values for the decay constant). All presented fits, therefore, include the WT replicates at $x = 0$ unless otherwise stated. For $metl$ mutants the methylation decay is too rapid to be accessible in the transgenerational data. Only the steady-state methylation level is available, which corresponds directly to $\delta$ (see Eq. (3)).

**Mathematical model for TE methylation dynamics: numerical fits**

The methylation timeseries, $\langle mC \rangle(x)$, are fit using `curve_fit` from the `optimize` library in `scipy` (Virtanen et al., 2020). First, we performed fits using three free parameters: $M_0$, $M^*$ (both as defined previously), and $B$, the decay constant:

$$\langle mC(x) \rangle = (M_0 - M^*) e^{-Bx} + M^*$$

and the total fit uncertainty, $\sigma_{fit}$, is estimated using the sum of the diagonal of the covariance matrix:

$$\sigma_{fit} = \sigma_{M_0}^2 + \sigma_B^2 + \sigma_{M^*}^2.$$ 

The fit parameters were constrained to $0 < M_0 < 1$, $0 < M^* < 1$ and $B > 0$, while using initial values of: $M_0 = 0.8$, $B = 1$ and $M^* = 0.3$. We do not fit to $\delta$ and $\epsilon$ directly, as the numerical routine is more reliable when fitting to parameters with an order of magnitude close to one. As $B = n_{cc}(\delta + \epsilon)/2$, we reject any fits for which $B > n_{cc}$ as this corresponds to an unphysical value for $\delta$ or $\epsilon$. The majority of TEs produce good fits and a small minority give exceptionally poor fits. To be conservative, we reject all fits with $\sigma_{fit} > 1$. A reliable estimate of the decay constant
requires an appreciable decrease in the methylation level with time. We therefore exclude TEs with \( \langle mCG \rangle_{(F2,F3)} \leq \lambda M^* \) for \( h1ddml, h1ddmlcmt2 \) and \( h1ddmlcmt3 \) or \( \langle mCG \rangle_{(F3,F4)} \leq \lambda M^* \) for \( h1ddmlcmt3 \) using a threshold of \( \lambda = 1.05 \), where \( \langle mCG \rangle_{(F2,F3)} \) is the average methylation level of all F2 and F3 replicates for a particular mutant. Upon manual inspection of the remaining fits, we also excluded nine further TEs. Finally, we selected the subset of TEs with acceptable fits for all four mutants (amounting to 1617 TEs, over 85% of the initial set that we attempted to fit).

The values of \( \delta \) and \( \epsilon \) are then given by:

\[
\delta = \frac{2BM^*}{n_{cc}}
\]

\[
\epsilon = \frac{2B(1 - M^*)}{n_{cc}}.
\]

For a function \( f(x_1,x_2,x_3) \) the uncertainty in \( f \) is given by

\[
\sigma_f^2 = \mathbf{v}^\top \cdot \mathbf{\sigma} \cdot \mathbf{v},
\]

where \( \mathbf{\sigma} \) is the covariance matrix for \( x_1, x_2 \) and \( x_3 \) and \( \mathbf{v} \) is a vector with elements \( v_i = \frac{\partial f}{\partial x_i} \). The uncertainties in the fitted values of \( \delta \) and \( \epsilon \) are therefore:

\[
\sigma_\delta = \frac{2BM^*}{n_{cc}} \sqrt{\left(\frac{\sigma_B}{B}\right)^2 + \left(\frac{\sigma_M^*}{M^*}\right)^2 + \frac{2\sigma_{BM^*}}{BM^*}}
\]

\[
\sigma_\epsilon = \frac{2B(1 - M^*)}{n_{cc}} \sqrt{\left(\frac{\sigma_B}{B}\right)^2 + \left(\frac{\sigma_M^*}{1 - M^*}\right)^2 - \frac{2\sigma_{BM^*}}{B(1 - M^*)}},
\]

where \( \sigma_{BM^*} \) is the covariance of \( B \) and \( M^* \).
Two-sided paired t-tests were performed for all mutant pairings of $\epsilon$-distributions (p-value $< 1 \times 10^{-170}$ for all pairings) and, similarly, for all mutant pairings of $\delta$-distributions p-value $< 1 \times 10^{-200}$ for all pairings). Finally, we note that the presence of the small peak at low $\delta$-values (Fig. 7F) is likely an artifact arising for TEs with a small decay constant. Due to their slow approach to steady-state, the fit algorithm may underestimate the precise value of $M^*$, and hence $\delta$.

*Mathematical model for TE methylation dynamics: limitations of the model*

We have assumed that the *de novo* rate, $\delta$, is independent of the existing methylation level. This may not always be the case. For example, a decrease in RdDM targeting as more methylation is lost is possible, as appears to be the case for a small subset of TEs (Fig. S6F-H). Alternatively, there may be an analogous cooperative *de novo* pathway to that proposed for clustered CG sites as found in CG islands of mammalian genomes (Lövkvist et al., 2016), in which case $\delta$ would represent the average *de novo* rate across the whole TE. Due to the cooperativity, the effective $\delta$ could then reduce across generations as the overall methylation level declines. In the case of small variations of the *de novo* rate over successive generations, the model will still produce a good fit to a decaying exponential with effective rates for both $\delta$ and $\epsilon$. A much larger variation in $\delta$ as a function of time, however, would likely result in a clearly visibly non-exponential decay of the methylation level, which is not observed in our data.

A further issue is that the values of $\delta$ and $\epsilon$ could each be different in the germline versus in somatic tissue (e.g., in the leaf). Our model assumes that these values are the same. Experimental data indicates that F5 mCG levels in the *h1ddm1* leaf and sperm are similar (Fig. 2A), which supports similar values of $\delta$ and $\epsilon$ in the germline and somatic tissue.
The model also assumes that all CG sites within a given TE are equivalent and all exhibit the same dynamical turnover of methylation, parameterized through $\delta$ and $\epsilon$. However, as is apparent from the per-CG site methylation histograms (Fig. 1F, S1H), a minority of CG sites are always highly methylated or always exhibit a near-absence of methylation, configurations that are essentially excluded within our model. Future model developments may need to better incorporate this heterogeneity.

Use of previously published datasets

GSE179796: ChIP-seq data from (Choi et al., 2021).

GSE122394: met1 and h1met1 bs-seq data from (Choi et al., 2020).

GSE96994: h1 bs-seq and nucleosome data from (Lyons & Zilberman, 2017).

GSE150436: H2A.W ChIP, ddm1 x Col-0 bs-seq, and RNA-seq from (Osakabe et al., 2021).

DRS019614: ddm1 G9 (F10) bs-seq data from (Ito et al., 2015)


Author contributions

D.B.L. conceived the research plan, performed experiments and analysis, and wrote the paper

A.B. developed the mathematical model, performed analysis, and wrote the paper

S.H. performed sperm isolation and library preparation

J.Choi contributed to RNA-seq analysis

E.H. contributed to DNA methylation analysis

J. Colicchio measured plant phenotypes
I. A. measured plant phenotypes

X.F. supervised experiments on sperm DNA methylation

M.H. developed the mathematical model, supervised analysis, and wrote the paper

D.Z. conceived the research plan, supervised experiments, and wrote the paper

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Figure legends

Figure 1. Heterochromatic mCG decays to an intermediate equilibrium in h1ddm1.

A) Average fractional CG methylation (mCG) is plotted for all heterochromatic TEs >250 bp (Choi et al., 2020) in the indicated genotypes and h1ddm1 generations.

B) Genome browser view of a 400 kb window of pericentromeric heterochromatin illustrating mCG across h1ddm1 generations (Chr. 5:12,950,000-13,350,000).

C) Boxplots of mCG for all h1ddm1 heterochromatic TEs (red), beginning with WT (green, left) and ending with ddm1 F5 (dark orange, right). Box depicts the interquartile range and whiskers extend to 1.5X interquartile range; horizontal line shows the median value.

D) F2 h1ddm1 per-TE mCG for all TEs >2 kb as a function of H3K9me2 (expressed as log2 ChIP/input) in the parental genotype, h1.

E) Heatmaps of per-TE mCG Pearson's correlation (color bar) for mCG vs. mCG, mCHG, and mCHH across h1ddm1 generations (x-axes, from left to right respectively). Biological replicates are indicated with a grey bar over columns/rows (one replicate for the F10, two for all other generations). Note increased correlation between CG and non-CG methylation in later generations.

F) Density plot of per-CG methylation for all CG sites with coverage >10 in heterochromatic TEs for the indicated genotypes.

G) 2D density plots of per-CG methylation for all CG sites in heterochromatic TEs for F4 vs. F5 generations (except for h1ddm1, where F2 vs. F3 is shown to illustrate the correlation between intermediate values across generations at a stage when mCG is changing the most). Darker color indicates more data points.

H) Density plots of mCG per-bisulfite-read with >=3 CG sites per read from heterochromatic TEs.

I) Genome browser view of bisulfite reads for WT (top) and h1ddm1 (bottom); Chr.1:14,918,800-14,918,900. Only (+) strand mapping reads are shown for clarity.

Figure 2. Intermediate h1ddm1 mCG is heritable.

A) Boxplots of F5 mCG for all heterochromatic TEs with leaf values (L) displayed next to sperm (S) in the indicated genotypes.

B-C) Average mCG in sperm and leaf for the indicated generations and genotypes plotted relative to the start and end sites of heterochromatic TEs, as in Fig. 1A.
D) Genome browser view of 500 kb of heterochromatin (Chr 5: 11,050,000-11,550,000) highlighting the similarities between sperm and leaf h1ddm1 mCG. Source tissue is either leaf (L) or sperm (S). All non-h1ddm1 genotypes shown are F5 generation.

E) 2D density plots comparing per-CG methylation at sites with coverage >10 in F4 sperm vs. F4 leaf (top) or F5 sperm (bottom) across heterochromatic loci. Pearson’s r is shown. Darker color indicates more data points.

F) Density plots of sperm per-CG methylation for all CG sites with coverage >5 in heterochromatic TEs.

G) Density plots of sperm mCG per-bisulfite-read with ≥3 CG sites per read.

H) Screenshot of per-read methylation from F4 sperm of WT and h1ddm1 at Chr.3: 12,425,889-12,426,499. Only (+) strand is shown for clarity.

Figure 3. Promoter methylation is quantitatively associated with TE expression.

A) Histograms of replicate-averaged expression levels (log2 transformed transcript-per-million, TPM) of ddm1 de-repressed heterochromatic transcripts with expression above 2.5 TPM (n=796).

B) Boxplots of expression data from (A) showing biological replicates.

C) DNA methylation averages of indicated cytosine context (y-axis) plotted from transcription start and end sites of TEs that are similarly expressed in ddm1 and h1ddm1 (qval >0.05 based on likelihood ratio test (Pimentel et al., 2017) and ddm1 average expression level >2.5 TPM, n = 284).

D) Genome browser view of interval showing depletion of DNA methylation at a TE gene promoter and corresponding expression (promoter region highlighted in box). Cytosine context noted in grey.

E) Methylation averages as in (C) of TEs downregulated in h1ddm1 (qval <0.05 and ddm1 average expression level >2.5 TPM, n = 210).

F) Genome browser view of interval showing promoter mCG at a TE gene promoter and corresponding repression of expression in h1ddm1 but not ddm1 (promoter region highlighted in box).

G) Expression level per ddm1 derepressed TE >2 kb (n=146) and corresponding h1ddm1 F3 values as a function of their respective promoter mCG, where the promoter region is +/- 500 bp of TSS.

H) Methylation averages of indicated context (y-axis) plotted from TE TSS (TSS=0 on y-axis; left, categories as from (Osakabe et al., 2021); right, categories determined in this study).
I) Boxplots of TE expression for published categories (Osakabe et al., 2021, left; neutral: 414, inactive: 381) and categories determined in this study (right, neutral: 450, inactive: 253).

J) ddm1 H2A.W (log2 ChIP/input) as a function of ddm1 expression level per TE.

Figure 4. *h1ddm1* rescues diverse *ddm1* phenotypes.

A) Boxplot illustrating mCHG hypermethylation in *ddm1* (F4) gene-body methylated genes is partially ameliorated in F4 *h1ddm1* (n=561).

B) Genome browser view of *ddm1* mCHG hypermethylation and its reduction in *h1ddm1* (F4) at AT4G09130.

C) Plants at 4 weeks post-germination, genotypes are indicated. All genotypes are F4 except *h1ddm1* is F7.

D) Boxplots of plant measurements conducted on F4 plants of indicated genotypes. Sample numbers per genotype are as follows: WT: 43; *h1*: 38; *ddm1*: 35; *h1ddm1*: 49.

E) Boxplots of mCG in genes hypomethylated in *ddm1* (n=529, p<1.0x10^-13, Fisher exact test). *ddm1* F10 obtained from (Ito et al., 2015).

Figure 5. Estimation of per-TE de novo and maintenance mCG rates in *h1ddm1*.

A) Diagram illustrating the relationship between *de novo* methylation (δ), maintenance methylation failure (ε), and the methylation states of a given CG site. U = unmethylated, M = methylated, and H = hemimethylated.

B) The average mCG of all TEs>2 kb with WT mCG>0.65 across *h1ddm1* generations (generation 1 = *h1ddm1* F2) fits an exponential decay. Solid line indicates the fit without including WT methylation values as the initial data point (generation 0); dashed line represents the WT+*h1ddm1* fit.

C) Scatterplot of δ vs. ε at those TEs fit by an exponential (n=1617). Green lines indicate the δ and ε required to achieve the indicated steady-state methylation level, M*.

D) Scatterplot of modeled steady-state methylation (M*) vs. *h1ddm1* mCG at F11.

E) Plot of changes in methylation per cell cycle for a hypothetical TE with 100 CG sites beginning with 90% methylation and with average *h1ddm1* δ and ε values (0.021 and 0.033, respectively). Dotted line illustrates the number of changes at steady-state (0.64).
F) Different δ and ε regimes produce a variety of mCG dynamics in h1ddm1. Quintiles of δ and ε were used to subset the modeled TEs and fit their respective average mCG. Generation 0 = WT, generation 1 = h1ddm1 F2, etc.

G) Genome browser views of representative TEs from each of the 4 groups in (F). Numbers listed next to curves in (F) correspond to browser views shown.

Figure 6. Steady-state mCG in h1ddm1 depends on DRM2.

A) Average mCG is plotted for all heterochromatic TEs for indicated genotypes around TE start and stop sites (TEs >250 bp). h1ddm1, h1ddm1cmt2, and h1ddm1cmt3 resemble one another at F7, whereas mCG is much reduced in h1ddm1drm2.

B) Boxplots of per-TE mCG over seven generations for h1ddm1 and compound h1ddm1 methyltransferase mutants (h1ddm1cmt3 F2 is absent; please see text).

C) Scatterplot comparing h1ddm1drm2 F7 to h1ddm1 F7 average mCG of previously modeled TEs (n=1619), with blue points highlighting the TEs significantly hypomethylated in h1ddm1cmt3 relative to h1ddm1 (n=86, p-val <0.01, Fisher exact test).

D) Boxplots comparing mCG in modeled TEs that are either hypomethylated in h1ddm1cmt3 (red) or control TEs that are not significantly hypomethylated (black).

E-F) Genome browser view illustrating mCG (E) and mCHH (F) at TE with CMT3-dependent mCG.

G) Boxplots comparing mCHH levels as in (D). Note the near complete depletion of mCHH in h1ddm1cmt3 but not -cmt2 at CMT3-depndent TEs, indicating CMT3 is required for DRM2 activity at these loci.

Figure 7. Estimation of mCG rates in compound h1ddm1 methyltransferase and met1 mutants.

A-C) Comparison of average modeled TE mCG decay fits (colored by genotype) to h1ddm1 (black dashed lines).

D) Density plots of modeled ε for individual TEs.

E) Scatterplot showing h1ddm1 ε as a function of parental h1 H3K9me2.

F) Density plots of modeled δ (see Methods regarding the low-δ peaks).

G) Boxplots of the change in δ (left) or ε (right) for the indicated genotypes. Orange horizontal lines are at y=0.
H) Boxplots of modeled steady-state methylation ($M^*$) for the indicated mutants.

I-K) Scatterplots of modeled steady-state methylation ($M^*$) vs. mCG at F7 for the indicated genotypes, with Pearson’s r shown.

L) Scatterplot of $\delta$ vs. $\varepsilon$ in h1ddm1drm2 TEs, with differently colored points indicating the different extremes possible in this mutant, grouped per quintile combination.

M) Color-matched exponential decay fits to average mCG of the quintile combinations shown in (L). Generation 0 = WT, generation 1 = h1ddm1drm2 F2, etc.

N) Density plots depicting the distributions of $\delta$ for the indicated genotypes in either MET1-dependent (left, n=271) or -independent (right, n=1258) modeled TEs (see Methods). The 5.5% of modeled TEs not in these categories were excluded from this analysis. Red dashed lines indicates the estimated false-positive methylation rate for the met1 and h1met1 data (see text).

Supplemental Figure Legends

Supplemental Figure 1.

A-B) Average fractional h1ddm1 mCHG (A) and mCHH (B) is plotted for all heterochromatic TEs >250bp for indicated genotypes around TE start and stop sites, as in Fig. 1A.

C) Average ddm1 mCG per generation (per replicate) from F2, beginning with WT, to highlight the steep drop in mCG that does not match that of the gradual h1ddm1 pattern (note differences in y-axes values with panel D). Points are overlayed with a LOESS curve, smoothed with default ggplot stat_smooth settings (span=0.75, polynomial degree=2), with 95% confidence interval shown in grey.

D-F) Average h1ddm1 mCG (D), mCHG (E), and mCHH (F) in heterochromatic TEs plotted as single points per generation (per replicate) overlayed with a LOESS curve as in (C).

G) Average h1ddm1 mCG at well-positioned (red, left panels) or poorly-positioned (cyan, right panels) nucleosomes in heterochromatic TEs (Lyons & Zilberman, 2017). Generation of h1ddm1 methylation data is indicated on the right.

H) Density plots of per-CG methylation for all CG sites with coverage > 10 in heterochromatic TEs in all h1ddm1 generations assayed.
I) Pairwise per-TE mCG Pearson’s correlations for indicated h1ddml generations represented as a heatmap.

**Supplemental Figure 2.**

A-B) Line plots of average h1ddml mCHG (A) and mCHH (B) at all heterochromatic TE s for the indicated genotypes around TE start and stop sites. Leaf is shown at the top, sperm at the bottom.

**Supplemental Figure 3.**

A) Scatterplot of WT H3K9me2 vs. WT H3K9me1 at ddm1 transcribed loci illustrating that most upregulated ddm1 transcripts are in regions with heterochromatic histone modifications.

B) Scatterplot of ddm1 vs. h1ddml average expression levels for heterochromatic transcripts. Color indicates whether transcript was found to be significantly changed (q-val <0.05, likelihood ratio test per (Pimentel et al., 2017)) in h1ddml.

C) Boxplots of expression levels of euchromatic loci upregulated in ddm1 for the indicated genotypes, with biological replicates shown individually.

D) Scatterplot of Col-0 x ddm1 vs. ddm1 F1 average expression levels for TE genes as defined in (Osakabe et al., 2021) using TPM from reanalysis performed in this study. Color indicates classification of TE gene as inactive (red) or neutral (black) based on source data from (Osakabe et al., 2021).

E) Scatterplot of Col-0 x ddm1 vs. ddm1 F1 average expression levels for TE genes using TPM from reanalysis performed in this study. Red color (inactive) indicates that a transcript is both significantly downregulated at qval <0.05 and (ddm1 / Col-0 x ddm1 F1) >4.

**Supplemental Figure 4.**

A) Plants at 4 weeks post-germination, genotypes are indicated. All genotypes shown are F4, except h1ddml is F14.

B) Scatterplot of genic mCG in ddm1 vs. WT illustrating genes that are significantly hypomethylated in ddm1 (n=529, right panel; p-val<1 x 10^{-13}, Fisher exact test).

**Supplemental Figure 5.**

A) Scatterplot of h1ddml δ vs. mCG in ddm1 x Col-0 F1 as compared to average mCG of ddm1 and WT (the expected level if mCG of the ddm1 chromosomes is unchanged from ddm1 levels in ddm1 x Col-0 F1) at TE genes. mCG is calculated as indicated on the x-axis.
B-C) Scatterplots of *h1ddm1* δ as a function of F11 mCHG (B) and mCHH (C). Pearson’s r is shown for both plots.

**Supplemental Figure 6.**

A) Plants at 4 weeks post-germination, genotypes are indicated such that the rightmost 3 samples are also *h1ddm1* in addition to the indicated genotype. Generation of mutants: F7; WT is F4.

B-C) Average mCHG (B) and mCHH (C) is plotted for all heterochromatic TEs for the indicated genotypes and generations around TE start and stop sites. These were selected to illustrate depletion of mCHG in the case of *h1ddm1cmt3* and the depletion of mCHH in *h1ddm1drm2* and *h1ddm1cmt2*.

D) Scatterplot depicting per-TE mCG of modeled TEs in *h1ddm1cmt3* vs. *h1ddm1* at F7. Red color indicates TEs significantly hypomethylated in *h1ddm1cmt3* (n=86, >50% reduction of mCG and p-val<0.01, Fisher exact test).

E) Boxplots comparing mCHG in modeled TEs that are either hypomethylated in *h1ddm1cmt3* (red) or control TEs that are not significantly hypomethylated (black).

F-H) Boxplots of mCG (F), mCHG (G) and mCHH (H) across the generations in TEs either mCG hypomethylated in *h1ddm1cmt3* (top) or all remaining TEs (bottom).

**Supplemental Figure 7.**

A) Scatterplots of changes in δ vs. changes in ε for the indicated genotype relative to *h1ddm1* values. Pearson’s r is shown.

B-C) Scatterplots of *h1ddm1* δ vs. mCG (implied δ) in *met1* (B) and *h1met1* (C) at modeled TEs. Pearson’s r is shown.
References


Huff, J. T., & Zilberman, D. (2014). Dnmt1-Independent CG Methylation Contributes to


Supplemental Figure 1

A

B

C

D

E

F

G

H

I

Supplemental Figure 1

A

B

C

D

E

F

G

H

I
Figure 2
Supplemental Figure 2
Figure 3

A

B

C

ddm1.H2A.W

log2(ddm1.tpm + 1)

similarly expressed TEs

E

downregulated TEs

D

F

neutral

inactive

G

H

Osakabe et al.

re-analysis

I

Osakabe et al.

re-analysis

J
Supplemental Figure 3

A

WT H3K9me2

B

\[
\text{log}_2 \text{ddm1 F3 TPM} + 1
\]

significantly different

similar

C

TPM

D

Osakabe et al.

not significantly different

similar

neutral

E

\[
\text{log}_2 \text{col x ddm1 F1 TPM} + 1
\]

inactive

neutral

re-analysis
Figure 4

A

mCG

WT
h1
ddm1 F4
h1ddm1 F4

0.00
0.01
0.02
0.03
0.04
0.05

B

mCG

WT
h1
ddm1 F4
h1ddm1 F4

genes

AT4G09130

0.0
100
200
300
400
500
600
700
800
900
1000
1100
1200
1300
1400

C

WT
h1
ddm1 F4
h1ddm1 F4

D

leaf area at 2 weeks (mm²)

final height (mm)

WT
h1
ddm1 F4
h1ddm1 F4

100
200
300
400
500
600
700
800
900
1000
1100
1200
1300
1400

E

mCG

WT
ddm1
h1ddm1

(final generation, F):

2
5
10
2
11

as.factor(variable)

as.numeric(as.character(value))

variable

WT
h1
ddm1 F4
h1ddm1 F4

true leaves at 3 weeks

final fruit
Supplemental Figure 4

A

B

not hypometh.  

hypo/hypermeth.  

WT mCG  

ddm1 mCG  

WT  h1  ddm1  h1ddm1  F14
Supplemental Figure 5

A  

B  

C  

Col-0 x ddm1 F1 mCG – (ddm1 mCG+Col-0)/2

r=0.48

r=0.66

r=0.99

h1ddm1 δ

h1ddm1 mCHG (F11)

h1ddm1 mCHH (F11)
Figure 6

**A**

![Graph A](image)

**B**

![Graph B](image)

**C**

![Graph C](image)

**D**

![Graph D](image)

**E**

![Graph E](image)

**F**

![Graph F](image)

**G**

![Graph G](image)
Supplemental Figure 7

A

h1ddm1cmt2

h1ddm1cmt3

h1ddm1drm2

h1ddm1 methyltransferase mutant delta – h1ddm1 epsilon

h1ddm1 methyltransferase epsilon – h1ddm1

B, C

δ (h1ddm1)

met1 mCG

h1met1 mCG

r = 0.28

r = 0.34

r = 0.83

r = 0.71

r = 0.68

r = 0.34