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**Technical Report** 

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# Combinatorial quantification of 5mC and 5hmC at individual CpG dyads and the transcriptome in single cells reveals modulators of DNA methylation maintenance fidelity

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Alex Chialastri  $0^{1,2}$ , Saumya Sarkar<sup>1,2</sup>, Elizabeth E. Schauer  $0^{1,2}$ , Shyl Lamba<sup>3</sup> & Siddharth S. Dey  $0^{1,2,4}$ 

Inheritance of 5-methylcytosine from one cell generation to the next by DNA methyltransferase 1 (DNMT1) plays a key role in regulating cellular identity. While recent work has shown that the activity of DNMT1 is imprecise, it remains unclear how the fidelity of DNMT1 is tuned in different genomic and cell state contexts. Here we describe Dyad-seq, a method to quantify the genome-wide methylation status of cytosines at the resolution of individual CpG dinucleotides to find that the fidelity of DNMT1-mediated maintenance methylation is related to the local density of DNA methylation and the landscape of histone modifications. To gain deeper insights into methylation/demethylation turnover dynamics, we first extended Dyad-seq to quantify all combinations of 5-methylcytosine and 5-hydroxymethylcytosine at individual CpG dyads. Next, to understand how cell state transitions impact maintenance methylation, we scaled the method down to jointly profile genome-wide methylation levels, maintenance methylation fidelity and the transcriptome from single cells (scDyad&T-seq). Using scDyad&T-seq, we demonstrate that, while distinct cell states can substantially impact the activity of the maintenance methylation machinery, locally there exists an intrinsic relationship between DNA methylation density, histone modifications and DNMT1-mediated maintenance methylation fidelity that is independent of cell state.

Inheritance of the epigenetic mark DNA methylation (5-methylcytosine, 5mC) during cell division is critical to ensure that cellular identity is transmitted from mother to daughter cells. While inheritance of DNA methylation is primarily performed by the maintenance DNA methyltransferase 1 (DNMT1) protein by copying methylated cytosines

in a CpG sequence context (5mCpG) from the old to new DNA strand, recent work has suggested that DNMT1 displays imprecise maintenance activity<sup>1-5</sup>. However, it remains unclear if the fidelity of DNMT1 varies at different genomic regions as well as when cells transition from one state to another. For example, one of the most dramatic illustrations

<sup>1</sup>Department of Chemical Engineering, University of California Santa Barbara, Santa Barbara, CA, USA. <sup>2</sup>Department of Bioengineering, University of California Santa Barbara, Santa Barbara, Santa Barbara, Santa Barbara, Santa Barbara, CA, USA. <sup>4</sup>Neuroscience Research Institute, University of California Santa Barbara, Santa Barbara, CA, USA. <sup>(4)</sup>Neuroscience Research Institute, University of California Santa Barbara, Santa Barbara, Santa Barbara, CA, USA.



Fig. 1 | Schematic and validation of M-M-Dyad-seq. a, The schematic shows that, in M-M-Dyad-seq, methylated cytosines at CpG dyads are detected using MspJI digestion followed by nucleobase conversion to interrogate the methylation status of the cytosine on the opposing strand of the dyad.
b, M-M-Dyad-seq-based quantification of genome-wide DNA methylation levels of wild-type mES cells grown under SL conditions in the absence (–) or presence (+) of the DNMT1 inhibitor decitabine for 24 h, and *Uhrf1*-knockout mES cells grown under SL conditions. The violin plots are made over 21,268 genomic bins.
c, M-M-Dyad-seq-based quantification of genome-wide 5mCpG maintenance

of this differential DNMT1 activity is the genome-wide erasure of 5mC while methylation is maintained at imprinted loci during mammalian preimplantation embryogenesis<sup>3</sup>. Further, we recently showed that the genome-wide loss of 5mC during mouse and human preimplantation development is heterogeneous, with a subset of cells experiencing global passive demethylation, where 5mC is not faithfully copied to the newly synthesized DNA strands<sup>6</sup>. Therefore, more generally, it remains unknown how cell-to-cell heterogeneity in DNMT1 maintenance methylation activity impacts cellular phenotypes. Investigating these questions remain a challenge due to a lack of experimental techniques. While the methylation status of CpG dinucleotides, a readout of DNMT1 maintenance activity, can be investigated using hairpin-bisulfite sequencing or extensions of this method, where complementary DNA strands are physically linked, these techniques typically have low efficiency and are challenging to scale down to a single-cell resolution<sup>7-9</sup>. Further, physically linking the two opposing strands using a hairpin prevents direct investigation of 5mC on one strand and the oxidized derivative 5-hydroxymethylcytosine (5hmC) on the other strand of a single DNA molecule. In this Technical Report, therefore, to investigate how the chromatin landscape and cellular state tune DNMT1 maintenance methylation fidelity and DNA methylation dynamics, we have developed a new technology (Dyad-seq) that integrates enzymatic detection of modified cytosines with traditional nucleobase conversion techniques to quantify all combinations of methylation levels of wild-type mES cells grown under SL conditions in the absence (–) or presence (+) of the DNMT1 inhibitor decitabine for 24 h, and *Uhrf1*knockout mES cells grown under SL conditions. 5mCpG maintenance is defined as the percentage of CpG dyads that are symmetrically methylated. The violin plots are made over 16,838 genomic bins. **d**, M-M-Dyad-seq-based quantification of genome-wide 5mCpHpG maintenance methylation levels of wild-type mES cells grown under SL conditions in the absence (–) or presence (+) of the DNMT1 inhibitor decitabine for 24 h, and *Uhrf1*-knockout mES cells grown under SL conditions. The violin plots are made over 2,404 genomic bins.

5mC and 5hmC at individual CpG dyads. Finally, we have scaled down Dyad-seq and integrated it with simultaneous quantification of the transcriptome from the same cell to gain deeper insights into how DNA methylation and DNMT1-mediated maintenance methylation regulate gene expression.

#### Results

# Dyad-seq can quantify all combinations of 5mC and 5hmC at individual CpG dyads

To address the above questions, we developed four different versions of Dyad-seq in bulk, two where 5mC is selected on one strand through the digestion of DNA with MspJI followed by the interrogation of 5mC or 5hmC on the CpG site of the opposing strand (M-M-Dyad-seq and M-H-Dyad-seq, respectively), and two where 5hmC is selected on one strand through the digestion of DNA with AbaSI followed by the interrogation of 5mC or 5hmC on the CpG site of the opposing strand (H-M-Dyad-seq and H-H-Dyad-seq, respectively)<sup>6,10</sup> (Figs. 1a and 2a and Extended Data Fig. 1a). Briefly, after digestion of DNA with either MspJI or AbaSI, the bottom strand of the fragmented molecules are captured by ligation to a double-stranded adapter containing the corresponding overhang, a sample barcode, a unique molecule identifier (UMI) and a PCR amplification sequence. We found that 92.5% of mapped reads had the correct orientation of adapter ligation, consistent with observations we made in previous methods developed

by us (Extended Data Fig. 1b-e)<sup>6,10,11</sup>. This ligation specifically allows the bottom strand of DNA to be interrogated for cytosine modifications using downstream cytosine conversion reactions and amplification. Next, to detect unmodified or methylated cytosines on the opposing DNA strand (that is, interrogated DNA strand), samples are either treated enzymatically with Tet2, T4-phage  $\beta$ -glucosyltransferase (T4-BGT) and APOBEC3A or with sodium bisulfite to convert unmodified cytosines to uracils while methylated cytosines remain unchanged (M-M-Dyad-seq and H-M-Dyad-seq)<sup>12</sup>. While the methylated cytosines detected on the bottom strand in these two methods include both 5mC and 5hmC, similar to other bisulfite-based sequencing methods, as the percentage of methylated cytosines that are 5hmC in most cell types is typically low and less than 10%, it is a reasonable approximation to call these methylated cytosines as 5mC<sup>13,14</sup>. Further, slight modification to the enzymatic conversion reaction by adding T4-BGT and APOBEC3A only results in the detection of 5hmC on the opposing DNA strand (M-H-Dyad-seq and H-H-Dyad-seq)<sup>15-19</sup>. The bottom strand of the adapter is unaffected by these conversion reactions as it is devoid of cytosines<sup>20</sup>. Next, extension using a random nonamer primer is used to incorporate part of the Illumina read 2 adapter sequence, and the resulting molecules are then PCR amplified and sequenced on an Illumina platform. From the sequencing data, the location of the methylated or hydroxymethylated cytosine on the nonamplified strand, detected by the endonuclease MspJI or AbaSI, can be inferred on the basis of its distance from the adapter, while the methylation/ hydroxymethylation status of the opposing CpG site, as well as other cytosines on this strand, can be determined directly from the sequencing results of the conversion reaction (Fig. 1a and Extended Data Fig. 1a). A summary of the DNA modifications recognized by the restriction enzymes, and the cytosine conversion reactions performed on the interrogated strand in Dyad-seq, is provided in Extended Data Fig. 1f. Thus, Dyad-seq not only enables measurement of the percentage of 5mC or 5hmC at a single-base resolution, similar to that obtained from bisulfite sequencing-based approaches, but also enables quantification of the percentage of 5mC or 5hmC maintenance at individual CpG dyads. Finally, M-H-Dyad-seq and H-M-Dyad-seq allow for the direct detection of two different epigenetic marks at individual CpG dyads, measurements that are not possible with hairpin bisulfitebased techniques.

To validate M-M-Dyad-seq, we first compared mouse embryonic stem (mES) cells grown with or without decitabine, a cytosine analog known to directly inhibit DNMT1 activity<sup>21</sup>. Treatment with decitabine for 24 h resulted in a global loss of DNA methylation as well as a dramatic reduction in 5mCpG maintenance, quantified as the fraction of CpG sites that are symmetrically methylated, demonstrating that M-M-Dyad-seq can be used to measure genome-wide DNA methylation levels and the fidelity of DNMT1-mediated maintenance methylation (Fig. 1b,c). Similarly, we performed M-M-Dyad-seq in a *Uhrf1*-knockout E14 cell line to find that both the genome-wide DNA methylation and

Fig. 2 | Dyad-seq variants enable genome-wide detection of all combinations of 5mC and 5hmC at individual CpG dinucleotides. a, A schematic describing four different versions of Dyad-seq. M-M-Dyad-seq profiles 5mC on one strand and if the cytosine on the opposing strand of the dyad is methylated or not. M-H-Dyad-seq profiles 5mC on one strand and if the cytosine on the opposing strand of the dyad is hydroxymethylated or not. H-H-Dyad-seq profiles 5hmC on one strand and if the cytosine on the opposing strand of the dyad is hydroxymethylated or not. H-M-Dyad-seq profiles 5hmC on one strand and if the cytosine on the opposing strand of the dyad is hydroxymethylated or not. H-M-Dyad-seq profiles 5hmC on one strand and if the cytosine on the opposing strand of the dyad is methylated or not. **b**, 5mCpG maintenance, quantified as the percentage of CpG dinucleotides that are symmetrically methylated, is shown for mES cells grown under different conditions. M-M-Dyad-seq is used to estimate 5mCpG maintenance. The violin plots are made over 23,766 genomic bins. **c**, M-H-Dyad-seq shows the percentage of 5mC that are paired with 5hmC at CpG dyads. The violin plots are made over 2,519 genomic bins. **d**, H-H-Dyad-seq shows the percentage of 5hmC that are

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5mCpG maintenance percentage were substantially lower than in wild-type E14 cells, consistent with the known mechanism of UHRF1 in recruiting DNMT1 to hemimethylated DNA to enable maintenance of DNA methylation (Fig. 1b,c). In addition, CpHpG maintenance methylation was very low in all experimental conditions, consistent with the known preference of DNMT1 to maintain methylation only at CpG sites in mammalian cells (Fig. 1d)<sup>3</sup>. Overall, these results, together with additional benchmarking experiments/analysis and comparison to bisulfite sequencing, hairpin-bisulfite sequencing and scMspJI-seq (a strand-specific method we recently developed for quantifying DNA methylation), show that M-M-Dyad-seq can be used to quantitatively measure DNA methylation and 5mCpG maintenance levels across the genome (Fig. 1b–d, Extended Data Figs. 1g,h, 2 and 3 and Supplementary Note 1)<sup>6,22-26</sup>.

After validating the technique, we next applied Dyad-seq to an in vitro model of epigenetic reprogramming by transitioning mES cells cultured in serum-containing medium supplemented with leukemia inhibitory factor (LIF) (denoted by 'SL') to a serum-free medium (basal medium) containing LIF and two inhibitors, GSK3i (CHIR99021) and MEKi (PD0325901) (denoted by '2iL') (Fig. 2b-g)<sup>27</sup>. These two states of mES cell are interconvertible, with SL mES cells displaying high levels of methylation that become hypomethylated in 2iL<sup>24</sup>. After transitioning SL mES cells to 2iL for 48 h, we similarly observed a global loss of 5mCpG that is accompanied with a corresponding reduction in DNMT1-mediated maintenance methylation activity (Fig. 2b, f). These genome-wide changes were also observed over different functional elements in the genome (Supplementary Fig. 1a-n). Further, these results were consistent across replicates with robust differences between the experimental culture conditions (Extended Data Fig. 4a). Similarly, when we bin the genome into 100 kb regions, we observed that, in more than 95% of the bins, reduced 5mCpG in the 2iL condition is associated with reduced maintenance methylation, consistent with previous observations that passive demethylation is the main contributor to this demethylation process (Extended Data Fig. 4b)<sup>28-30</sup>. To further investigate the role of different modes of demethylation during this global erasure of the methylome, we transitioned SL mES cells to different medium conditions for 48 h and performed all four variants of Dyad-seq (Fig. 2b-g and Extended Data Fig. 4c,d). In the basal medium containing neither of the two inhibitors or LIF (denoted by 'No'), cells spontaneously differentiated with a rapid increase in both the absolute levels of 5mCpG as well as DNMT1-mediated maintenance methylation (Fig. 2b, f and Extended Data Fig. 4c). In the case where LIF alone (denoted by 'BL') or a combination of LIF and GSK3i (denoted by 'GL') were added to the basal medium, we observed limited changes in both the absolute levels of 5mCpG and maintenance methylation activity (Fig. 2b, f and Extended Data Fig. 4c). In contrast, basal medium containing LIF and MEKi (denoted by 'ML') induced an even larger decrease in both the global levels of 5mCpG and DNMT1-mediated maintenance methylation fidelity than 2iL (Fig. 2b,f and Extended

paired with 5hmC at CpG dyads. The violin plots are made over 2,496 genomic bins. **e**, H-M-Dyad-seq shows the percentage of 5hmC that are paired with 5mC at CpG dyads. The violin plots are made over 2,506 genomic bins. **f**, Genomewide 5mCpG levels quantified using M-M-Dyad-seq. The violin plots are made over 22,468 genomic bins. **g**, Genome-wide 5hmCpG levels quantified using M-H-Dyad-seq. The violin plots are made over 2,599 genomic bins. In **b** and **f**, the violin plots are made using 100 kb bins, and in **c**-**e** and **g**, the violin plots are made using 1 Mb bins. In **b**-**g**, mES cells are grown under SL conditions before being transitioned to the following conditions for 48 h: No, basal medium; BL, basal medium with LIF; SL, serum-containing medium with LIF; GL, basal medium with LIF and GSK3i; 2iL, basal medium with LIF, GSK3i and MEKi; ML, basal medium with LIF and MEKi. **h**, A heatmap of differentially expressed genes with a putative role in regulating DNMT1-mediated maintenance methylation fidelity. The data in this figure are based on three independent biological replicates. Data Fig. 4c). Interestingly, although the maintenance methylation fidelity decreased in both ML and 2iL conditions compared to SL and No, M-H-Dyad-seq showed that the fraction of 5mC-containing dyads that were paired with 5hmC on the opposing strand were similar for No, SL, 2iL and ML (Fig. 2c). Similarly, the genome-wide levels of 5hmC were also mostly unchanged between these different conditions (Fig. 2g and Extended Data Fig. 4d). Overall, these results highlight that the global gain or loss of methylation in this system is not dependent on the

activity of Ten-eleven translocation (TET) family proteins but is closely linked to the fidelity of DNMT1-mediated methylation maintenance.

As we observed that the transition of SL mES cells to the No condition was associated with an increase in both 5mCpG and DNMT1-mediated maintenance methylation, while the 2iL and ML conditions were associated with a decrease in both 5mCpG and DNMT1-mediated maintenance methylation, we hypothesized that quantifying the transcriptome in these different conditions could



potentially be used to identify new factors that are involved in regulating the DNA maintenance methylation machinery. For example, 2iL and ML conditions involve the inhibition of the MAPK/ERK pathway, which has previously been shown to reduce protein levels of DNMT1 in multiple systems, including during the transition of mES cells from SL to 2iL<sup>30-32</sup>. Therefore, we performed RNA sequencing (RNA-seq) on all conditions and, as expected, found each condition to be transcriptionally distinct (Extended Data Fig. 4e, f). As DNMT1 displayed reduced maintenance methylation fidelity in the ML and 2iL conditions, but an increase in the No condition, we reasoned that putative genes involved in tuning maintenance methylation could be identified as those that are upregulated or downregulated in ML and 2iL when compared to No, but are expressed at intermediate levels in SL, GL and BL conditions. Using this combinatorial criteria, we identified 61 differentially expressed genes, 39 of which were highly expressed in the 2iL and ML conditions with enrichment in pathways associated with pluripotency, negative cell cycle regulation and blastocyst development, while 22 genes were highly expressed in the No condition with enrichment in pathways associated with the negative regulation of ERK1 and ERK2 cascade and mesenchymal cell differentiation (Fig. 2h and Extended Data Fig. 4g,h)<sup>33</sup>. Notably, our screen identified developmental pluripotency associated 3 (Dppa3) as one of the hits that is highly expressed in the ML and 2iL condition (Fig. 2h and Extended Data Fig. 4f). Previous studies have found that ectopic expression of DPPA3 leads to global hypomethylation, while Dppa3 knockout leads to global hypermethylation<sup>29,34</sup>. Further, DPPA3 has even been shown to directly bind the PHD domain of ubiquitin like with PHD and ring finger domains 1 (UHRF1), a critical partner of DNMT1 necessary for 5mCpG maintenance, and displaces it from chromatin, thus inhibiting methylation maintenance<sup>29</sup>. The identification of a previously well-characterized factor involved in DNA methylation maintenance suggests that several of the other 60 genes identified in our screen, by combining Dyad-seq with RNA-seq, could potentially reveal novel regulators of DNA methylation maintenance fidelity and pluripotency.

We also used these experiments to further investigate the dynamics and relationship between DNA methylation and DNA hydroxymethylation at individual CpG dyads. While M-M-Dyad-seq showed, as expected, that a majority of 5mC existed as symmetrically methylated dyads instead of hemimethylated dyads, H-H-Dyad-seq showed that 5hmC was found to infrequently occur as symmetrically hydroxymethylated dvads (Fig. 2b.d). In contrast to this, H-M-Dvad-seq showed that 5hmC sites had high levels of 5mC on the CpG site of the opposing DNA strand, which showed similar trends to the global levels of 5mC among conditions (Fig. 2e, f and Extended Data Figs. 3d and 4c). This observation is in agreement with single-molecule fluorescence resonance energy transfer experiments, which, while lacking locus-specific information, globally identified that approximately 60% of 5hmC sites exist in a 5hmC/5mC dyad state in mES cell<sup>35</sup>. These measurements strongly suggest that in mES cells TET proteins hydroxymethylate only one of the two 5mC sites in a symmetrically methylated dyad and do not sequentially convert both 5mC to 5hmC. This result is consistent with previous in vitro work showing that TET proteins have a stronger binding affinity for symmetrically methylated dyads over hemimethylated dyads, and a crystal structure that indicates that the nonreactive cytosine is not involved in protein-DNA contacts<sup>36,37</sup>. In summary, these experiments show that the combination of the four Dyad-seq variants can provide insights into DNA methylation/hydroxymethylation turnover at a single-base and genome-wide resolution that was previously not possible.

# Fidelity of maintenance methylation is linked to the local density of the methylome and histone marks

As the genome-wide methylation levels were strongly related to the overall maintenance methylation activity for mES cells grown under different conditions, we next explored how methylation levels were linked to DNMT1-mediated maintenance methylation fidelity for different genomic regions within the same cell state (Fig. 2b, f). For example, as has been shown previously, we find that compared to other genomic loci within the same cell state, intracisternal A particles (IAPs) display high 5mCpG levels in SL as well as in the globally hypomethylated 2iL and ML conditions (Fig. 3a)<sup>28,38</sup>. Interestingly, these elevated 5mCpG levels at IAPs are associated with higher maintenance methylation activity, suggesting that high levels of methylation locally could be linked to increased DNMT1-mediated maintenance methylation fidelity (Fig. 3b). More generally, we find that genomic regions with increasing methylation levels are correlated with higher maintenance methylation fidelity, with a more pronounced effect in regions with higher CpG density, such as CpG islands (Fig. 3c,d and Extended Data Fig. 5a,b). Consistent with recent findings, our results from M-M-Dyad-seq suggest that there exists a tight coupling and a positive feedback loop between local methylation density and DNMT1-mediated maintenance methylation activity<sup>2</sup>.

Based on this discovery that 5mCpG maintenance can be tuned by the methylation levels at different genomic loci, we hypothesized that the genome-wide landscape of histone modifications could also impact DNMT1-mediated maintenance methylation. We found that highly methylated genomic regions are associated with high 5mCpG maintenance, independent of the histone modification at those loci (Fig. 3e). Surprisingly, however, we discovered that, for regions that are lowly methylated, the presence of a particular histone mark can dramatically alter the maintenance methylation fidelity, and that these observations were not influenced by the activity of TET1 (Fig. 3e and Extended Data Figs. 5c-k and 6a-m). For example, we found that regions enriched for the repressive mark H3K9me2 are associated with higher maintenance methylation fidelity than a randomly selected bin at similar methylation levels (Fig. 3e and Extended Data Fig. 5d). This is consistent with previous observations that UHRF1 can specifically bind H3K9me2 with high affinity, providing a mechanistic rationale for the recruitment of DNMT1 and higher maintenance seen in these regions<sup>39-41</sup>. Interestingly, we identify that enhancers marked by H3K4me1 or H3K27ac and active promoters/enhancers marked by H3K9ac also have increased DNMT1-mediated maintenance methylation fidelity (Fig. 3e and Extended Data Fig. 5e,h,k,l). In contrast, H3K4me3, a mark found at active gene promoters, had lower levels of 5mCpG maintenance than the average genome-wide maintenance (Fig. 3e and Extended Data Fig. 5j,l)<sup>42</sup>. Overall, these results demonstrate that, for regions of the genome that are lowly methylated, histone modifications can substantially alter maintenance methylation fidelity from approximately 40% to over 70% of the dyads being symmetrically methylated.

## scDyad&T-seq enables joint profiling of the methylome,

maintenance methylation and transcriptome from single cells In addition to the dramatic heterogeneity we observed in maintenance methylation across different genomic contexts, we next wanted to understand how cell-to-cell variability in DNMT1-mediated maintenance methylation fidelity could impact cellular phenotypes within a population. Addressing this question has been challenging as current hairpin bisulfite techniques have not successfully been applied to single cells. To overcome this limitation, we scaled down M-M-Dyad-seq to a single-cell resolution (termed as scDyad-seq). As proof of concept, cells treated with decitabine for 24 h experienced a global loss of DNA methylation and a dramatic reduction in the fraction of CpG dyads that were symmetrically methylated, while showing no changes in 5mCpHpG maintenance, as expected (Extended Data Fig. 7a,b). Further, to directly link heterogeneity in DNA methylation and DNMT1-mediated maintenance methylation fidelity to gene expression variability, we extended the method to simultaneously also capture the transcriptome (scDyad&T-seq) using an amplification strategy we recently developed<sup>43</sup>. Briefly, single cells are sorted into 384-well plates and the messenger RNA (mRNA) is converted to cDNA, as described previously, using poly-A primers with overhangs



#### **Fig. 3** | **DNA methylation maintenance activity is linked to the local density of the methylome and the distribution of histone modifications. a**, The bar plots show 5mCpG levels estimated using M-M-Dyad-seq at various repetitive elements for mES cells after 48 h in the indicated media conditions. The dots indicate independent biological replicates (*n* = 3). IAP, intracisternal A particles; LINE, long interspersed nuclear elements; SINE, short interspersed nuclear elements; LTR, long terminal repeats. b, The bar plots show 5mCpG maintenance fidelity estimated using M-M-Dyad-seq at various repetitive elements for mES cells grown under different conditions for 48 h. IAPs are more resistant to demethylation than other genomic regions and show higher 5mCpG maintenance levels. The dots indicate independent biological replicates (*n* = 3). **c**, A box plot of 5mCpG maintenance levels in 1 kb genomic bins over varying

absolute methylation levels. One-kilobase regions in which at least five unique

CpG dyads are detected are included. The number of bins in each category is denoted above each box plot. Data from mES cells grown in SL and profiled using M-M-Dyad-seq are shown. **d**, 5mCpG maintenance levels at CpG islands separated on the basis of absolute methylation levels. 'Methylated' indicates CpG islands with 5mCpG levels higher than 20%, 'mixed' indicates 5mCpG levels between 10% and 20%, and 'unmethylated' indicates CpG islands with 5mCpG levels lower than 10%. Data from mES cells grown in SL and profiled using M-M-Dyad-seq are shown. The dots indicate independent biological replicates (*n* = 3). **e**, A heatmap of 5mCpG maintenance percent in SL-grown mES cells at genomic regions enriched for various histone marks. The numbers within parentheses indicate the total number of regions analyzed in the meta-region. 5mCpG maintenance percent is not reported as only a small number of bins within H3K9me3 peaks were found to have low 5mCpG levels. N/A, not applicable.



Fig. 4 | scDyad&T-seq enables joint profiling of the methylome, maintenance methylation fidelity and the transcriptome from the same cell. a, Schematic of scDyad&T-seq. By scaling down M-M-Dyad-seq and combining it with the capture of mRNA, scDyad&T-seq can simultaneously quantify DNA methylation levels, maintenance methylation levels and the transcriptome from the same cell in a high-throughput format. Black, solid orange and dotted orange lines correspond to genomic DNA, mRNA and amplified RNA, respectively. **b**, The coverage of CpG dinucleotides that provide information on maintenance methylation (5mCpG dyad coverage), and coverage of CpG sites that enable

that contain a cell-specific barcode, UMI, 5' Illumina adapter and T7 promoter (Fig. 4a)<sup>43</sup>. Following this, the next steps are similar to M-M-Dyad-seq-based digestion and ligation where the genomic DNA within the reaction mix is glucosylated, digested with the restriction enzyme MspJI and ligated to Dyad-seq double-stranded adapters that contain a cell-specific barcode, UMI and cytosine-depleted PCR handle. Thereafter, as the genomic DNA and cDNA within individual cells are uniquely tagged, the samples are pooled and subjected to in vitro transcription (IVT). IVT only amplifies the T7 promoter containing cDNA molecules, and these amplified RNA molecules are pulled down using biotinylated poly-A oligonucleotides and magnetic streptavidin beads, as described previously<sup>43</sup>. The amplified RNA molecules are then used to prepare Illumina libraries, enabling quantification of the transcriptome of single cells<sup>43,44</sup>. The flowthrough from the bead separation contains unamplified Dyad-seq adapter ligated genomic

quantification of absolute methylation levels (CpG coverage), together with the number of unique transcripts detected in individual cells. Note that 5mCpG dyad coverage and CpG coverage are derived from nonoverlapping regions of the same sequencing read; therefore, the total number of CpG sites interrogated in a cell can be obtained by summing the 5mCpG dyad coverage and CpG coverage. **c**, The distribution of CpG sites that are detected by different methods (scDyad&T-seq, scMspJI-seq and scBS-seq) over genomic regions of varying CpG density<sup>6,26</sup>. The green curve shows the distribution of CpG sites over 5 kb bins in the mouse genome.

DNA molecules that are then subjected to nucleobase conversion, random linear amplification and PCR, similar to that described above for M-M-Dyad-seq, to quantify the methylome and the methylation status of CpG dyads on a genome-wide scale from the same single cells (Fig. 4a).

We first applied scDyad&T-seq to serum-grown mES cells to detect up to 75,835 unique transcripts per cell, and the methylation status of up to 1,118,393 CpG sites per cell, together with the additional detection of the maintenance methylation status of up to 203,620 CpG dyads per cells (with an average of 25,066 unique transcripts per cell (5,825 genes per cell), covering the methylation status of 328,967 CpG sites on average per cell and the maintenance methylation status of an additional 51,650 CpG dyads on average per cell) (Fig. 4b and Extended Data Fig. 7c–h). In comparison, while scM&T-seq, a method that captures the methylome and transcriptome from the same cell, covered



Fig. 5 | scDyad&T-seq can directly correlate heterogeneity in the methylome and DNMT1-mediated maintenance methylation fidelity to transcriptional variability in single cells. a, A heatmap comparing 5mCpG maintenance over individual chromosomes in mES cells computed using scDyad&T-seq with the strand bias metric that can be estimated from techniques such as scMspJI-seq from the same single cells. The heatmap shows that the 5mCpG maintenance estimated from scDyad&T-seq displays increased sensitivity in quantifying strand-specific DNA methylation compared to the strand bias metric obtained from scMspJI-seq<sup>6</sup>. The transcriptional group that individual cells belong to (top) and their genome-wide 5mCpG methylation levels (bottom) are also reported. b, Example of strand-specific methylation of three cells, P7L3.67, P7L4.78 and P7L3.69, from **a**. For cell P7L3.67, similar levels of 5mCpG detected on the plus and minus strand of each chromosome by the enzyme MspJI is in agreement with the high levels of 5mCpG maintenance quantified using scDyad&T-seq. In contrast, cells P7L4.78 and P7L3.69 show very similar levels of 5mCpG maintenance

more CpG sites and genes per cell, these libraries were proportionally sequenced at a much higher depth than the scDyad&T-seq libraries (Extended Data Fig. 7i)<sup>45</sup>. Further, as scM&T-seq requires physical separation of genomic DNA and mRNA before barcoding, amplification and

computed using scDyad&T-seq but display substantial differences when MspJIbased quantification is used to estimate strand-specific methylation, indicating the limited accuracy of the strand bias score used in scMspJI-seq. A low Pearson's correlation indicates deviations from a strand bias score of 0.5. The color of the data points indicates the 5mCpG maintenance percentage of individual chromosomes estimated using scDyad&T-seq. **c**, DNA methylation and 5mCpG maintenance levels at different genomic regions stratified by Stadler et al. as fully methylated regions (FMR), lowly methylated regions (LMR) and unmethylated regions (UMR)<sup>53</sup>. The data points represent 101 individual cells. **d**, A UMAP visualization of serum-grown mES cells based on the single-cell transcriptomes obtained from scDyad&T-seq. **e**, 5mCpG levels in regions marked by specific histone modifications. **f**, 5mCpG maintenance levels in regions marked by specific histone modifications. In **e** and **f**, cells are grouped on the basis of the transcriptional clusters shown in **d**. The data in **e** and **f** are based on 42 Nan<sup>HI</sup> and 55 Nan<sup>Lo</sup> cells.

library preparation, this method is more challenging to scale up to sequence thousands of cells per day and, unlike scDyad&T-seq, cannot capture the methylation status of individual CpG dyads in single cells. Finally, the distribution of CpG and GpC sites covered by scDyad&T-seq

was similar to their distribution in the genome and that detected by single-cell bisulfite sequencing, suggesting that scDyad&T-seq can capture the genome-wide methylome and maintenance methylation landscape in single cells (Fig. 4c and Extended Data Fig. 7j)<sup>26</sup>.

We further compared scDyad&T-seq to scMspJI-seq, a method we recently developed for strand-specific quantification of 5mC<sup>6</sup>. While scMspJI-seq does not have the resolution of individual CpG dyads, it can be used to estimate the extent of asymmetry in DNA methylation between two strands of DNA over a large genomic region<sup>6</sup>. We previously quantified this strand-specific DNA methylation using a metric known as strand bias, defined as the number of methylated cytosines on the plus strand divided by the total number of methylated cytosines on both DNA strands, with deviations from a score of 0.5 indicating asymmetric DNA methylation between the two strands of DNA<sup>6</sup>. Therefore, we directly compared the individual-CpG-dyad (or 5mCpG maintenance) resolution afforded by scDyad&T-seq to the strand bias score that can be obtained from both scDyad&T-seq as well as scMspJI-seq (as both these methods employ the same restriction enzyme MspJI). Overall, we found agreement between both measurements, with high 5mCpG maintenance associated with no strand bias (Fig. 5a,b and Extended Data Fig. 8a). Notably, however, we found that the strand bias score in general can only identify cells that experience high levels of strand-specific asymmetry in DNA methylation and, unlike the individual-CpG-dyad resolution of scDyad&T-seq, is not able to capture the wide gradient in strand-specific DNA methylation in single cells (Fig. 5a). Further, this limited sensitivity when using the strand bias score in scMspJI-seq possibly also arises from a lack of allele-specific resolution of this technique. For example, while low levels of 5mCpG maintenance (mean 5mCpG maintenance, 25.4%) in cell P7L3.69 was associated with strand bias scores that substantially deviated from 0.5 as expected, cell P7L4.78 displaying similarly low levels of 5mCpG maintenance (mean 5mCpG maintenance, 30.7%) showed very little strand bias (Fig. 5b). Similarly, we next used scDyad&T-seq to test the performance of a computational method for estimating maintenance methylation from nucleobase conversion-based 5mC sequencing data<sup>46</sup>. While the computational metric performed reasonably well for cells that displayed low 5mCpG maintenance, its accuracy was lower for cells with high levels of 5mCpG maintenance (Spearman correlation,  $\rho$  = 0.42; Extended Data Fig. 8b). Together these results demonstrate that scDyad&T-seq is a highly sensitive method for assessing the maintenance methylation status at individual CpG dyads in single cells, and when combined with the readout of absolute levels of methylation and the transcriptome from the same cell, this method could potentially allow us to directly link DNA methylation dynamics and heterogeneity to cellular phenotypes.

## Distinct cell states display varied maintenance methylation and methylome landscapes

After establishing that scDyad&T-seq can accurately quantify maintenance methylation fidelity, in addition to DNA methylation and the transcriptome in single cells, we next attempted to estimate the

Fig. 6 | Serum to 2iL transition of mES cells involves a transient loss of methylation maintenance fidelity and is associated with the emergence of two distinct 2iL transcriptional populations. a, b, Genome-wide methylation (a) and maintenance methylation levels (b) of individual mES cells cultured in SL or in 2iL conditions for 3, 6 or 10 days. c, Genome-wide 5mCpG methylation and 5mCpG maintenance levels of single cells as they transition from SL to 2iL conditions. Cells transition from highly methylated and highly maintained to a lowly methylated and lowly maintained or lowly methylated and highly maintained state. d, As cells transition from SL to 2iL conditions, they are classified as lowly (mC<sup>L0</sup>) or highly (mC<sup>H1</sup>) methylated and lowly (Mnt<sup>L0</sup>) or highly (Mnt<sup>H1</sup>) maintained on the basis of hierarchical clustering, as described in Extended Data Fig. 9a,b. e,f, Genomewide methylation (e) and maintenance methylated in mES cells grown in long-term cell-to-cell heterogeneity in the methylome and its relationship to transcriptional variability and cellular phenotypes. Surprisingly, even when considering regions of similar 5mC content in the genome based on previous bulk measurements, we observed substantial heterogeneity in both the methylation levels and 5mCpG maintenance fidelity between individual cells (Fig. 5c). Furthermore, we also noted that the relationship of higher 5mCpG maintenance fidelity at regions of the genome with higher methylation density is preserved at the single-cell level (Figs. 3c-e and 5c and Extended Data Fig. 5a,b). To systematically correlate heterogeneity in the epigenome to gene expression variability and the occurrence of distinct cell states, we first used the transcriptome to identify two subpopulations in the serum-grown mES cells-one high in NANOG, REX1 and ESRRB (referred to as NANOG high or 'Nan<sup>Hi</sup>') and one low in the expression of these genes (referred to as NANOG low or 'Nan<sup>Lo'</sup>) (Fig. 5d, Extended Data Fig. 8c and Supplementary Table 1). While these two well-established subpopulations in serum-grown mES cells are known to be transcriptionally heterogeneous with bimodal expression of key pluripotency genes, how these cell states are linked to the methylome and DNMT1-mediated maintenance methylation fidelity remains less well studied<sup>45,47</sup>. We found that cells in the Nan<sup>Hi</sup> state are globally hypomethylated with reduced genome-wide 5mCpG maintenance compared to the Nan<sup>Lo</sup> cell state (Mann–Whitney U test,  $P = 8 \times 10^{-6}$ and  $P = 3.6 \times 10^{-4}$ , respectively), suggesting that DNMT1-mediated maintenance methylation fidelity could play an important role in tuning the cell state in serum-grown mES cells (Fig. 5e,f).

We next explored how histone modifications impact 5mCpG maintenance in the context of different cell states. Consistent with bulk findings, H3K9me2/3-, H3K4me1-, H3K27ac- and H3K9ac-enriched genomic regions showed higher maintenance methylation fidelity while H3K4me3-marked regions showed reduced maintenance methylation fidelity (Fig. 5e,f and Extended Data Fig. 8d–g). Interestingly though, across all the histone modifications investigated, cells in the Nan<sup>Hi</sup> state consistently showed reduced methylation levels and correspondingly lower 5mCpG maintenance compared to Nan<sup>Lo</sup> cells, suggesting an intrinsic and mechanistic relationship between DNA methylation density and DNMT1-mediated maintenance methylation fidelity that is independent of cell state. Finally, these results also demonstrate that the cell state could play a key role in impacting the DNA methylation landscape (Fig. 5e, f).

# SL to 2iL demethylation is associated with heterogeneous maintenance methylation activity and transcriptionally distinct subpopulations

To further investigate how maintenance methylation fidelity is influenced as cells transition from one state to another, we applied scDyad&T-seq to mES cells that were switched from serum-containing medium (SL) to 2iL medium for 3, 6 or 10 days (2iL-D3, 2iL-D6 and 2iL-D10, respectively). While the transition from SL to 2iL resulted in dramatic erasure of DNA methylation and loss of maintenance methylation

2iL culture (denoted as 'SmC high') and at all other genomic regions (denoted as 'Other loci'), with each point denoting a single cell<sup>24</sup>. Data in these panels are based on 101 SL cells and 168 2iL-D3, 133 2iL-D6, 151 2iL-D10 cells. **g,h**, A UMAP visualization of SL- and 2iL-grown cells based on the single-cell transcriptomes obtained from scDyad&T-seq, and classified by culture conditions (**g**) or by transcriptome-based clustering (**h**). **i**, The expression levels of select genes and transposable elements, such as DPPA3, KHDC3, RLTR45 and RLTR45-int, that were found to be highly expressed in the 2iL-2 population. **j**,**k**, Genome-wide methylation (**j**) and maintenance methylation levels (**k**) of single cells in different transcriptional clusters shown in **h**. **I**, The percentage of 2iL-1 and 2iL-2 cells in the four groups classified on the basis of the genome-wide methylation and maintenance methylation levels. The numbers within parentheses indicate the total number of cells in the transcriptional clusters 2iL-1 and 2iL-2. activity, we surprisingly also found that the epigenetic reprogramming to the 2iL state is highly heterogeneous with a fraction of cells still retaining high levels of methylation and 5mCpG maintenance even after 10 days in 2iL (Fig. 6a–c). Using hierarchical clustering, cells were classified as either highly or lowly methylated (mC<sup>Hi</sup> and mC<sup>Lo</sup>), and highly or lowly maintained (Mnt<sup>Hi</sup> and Mnt<sup>Lo</sup>), leading to four distinct categories of epigenetic states (Extended Data Fig. 9a,b). Superimposing the time-course data on these epigenetic states shows that cells generally start off in a highly methylated and highly maintained state, with passive demethylation thereafter resulting in the loss of



5mC till they reach a lowly methylated and lowly maintained state. Surprisingly, a fraction of cells subsequently move toward a lowly methylated but highly maintained state to establish a globally hypomethylated genomic landscape that is maintained at high fidelity (Fig. 6c, d). Consistent with this result, we found that regions previously identified as retaining high methylation in the 2iL state also showed higher methylation in our data and, correspondingly, displayed higher maintenance methylation fidelity at all time points, including in SL-grown cells<sup>24</sup> (Fig. 6e, f). These results again highlight the intrinsic relationship between the local methylation density and maintenance methylation fidelity that is conserved through transitions in cell states (Figs. 3e, 5c,e,f and 6e,f). Further, these highly methylated regions were previously shown to correlate with H3K9me3 peaks, which are found at similar genomic regions for both SL- and 2iL-grown mES cells<sup>5,24,48</sup>. This, together with our previous observation that H3K9me2/3-marked regions show high 5mCpG maintenance fidelity, suggests that the relationship between methylation density and maintenance methylation fidelity across cell states in this system could be coupled to H3K9me2/3, and that the global impairment to the 5mCpG maintenance machinery in 2iL-grown mES cells is partially restored at H3K9me3-enriched regions (Fig. 3e and Extended Data Fig. 5i).

We next analyzed the gene expression patterns from the same cells to find that most cells in 2iL were transcriptionally similar and distinct from SL-grown cells, as expected (Extended Data Fig. 9c). Further, we found that the 2iL cells did not strongly separate by the time spent in culture, suggesting that the transcriptional reprogramming is quickly activated once cells are transitioned from SL to 2iL conditions (Fig. 6g and Extended Data Fig. 9c-g)<sup>49</sup>. Beyond the two broad transcriptional groups of SL-like and 2iL-like cells, further clustering revealed four distinct populations, with two clusters transcriptionally similar to SL-grown cells-Nanog low like-cells and Nanog high like-cells (referred to as 'Nan<sup>Lo-LC'</sup> and 'Nan<sup>Hi-LC'</sup>, respectively)-that primarily included serum-grown mES cells, and two similar to 2iL cells (referred to as '2iL-1' and '2iL-2') (Fig. 6h). Interestingly, we found that a small group of cells from the 2iL-D3 condition cluster with the Nan<sup>Lo-LC</sup> cell population. Previous work has shown that SL-grown cells in the Nan<sup>Lo</sup> state are less likely to survive the transition to 2iL when compared to those in the Nan<sup>Hi</sup> state<sup>50</sup>. This low survival rate of cells in the Nan<sup>Lo</sup> state could possibly arise due to an epigenetic barrier, and comparison of the 2iL-D3 cells that cluster with the SL-like cells shows that this group is characterized by high 5mCpG maintenance and correspondingly high methylation levels relative to the 2iL-D3 cells that successfully transition and cluster with the 2iL-like cells (Extended Data Fig. 10a,b). Further, consistent with recent observations, we find that the 2iL-D3 cells that cluster with the SL-like cells express low levels of Pou5f1 (Extended Data Fig. 10c)<sup>50</sup>. Finally, these cells also express *Sox1*, suggesting that they begin to differentiate toward the neuroectoderm lineage but do not survive long-term culture in 2iL conditions and are not present by day 10 (Fig. 6g,h and Extended Data Fig. 10d).

In contrast, cells that successfully transition their transcriptional program to 2iL-like cells could be split into two distinct clusters, 2iL-1 and 2iL-2, with a small group of genes and transposable elements that are differentially expressed between these two subpopulations (Extended Data Fig. 10e). Notably, we found that cells in the 2iL-2 cluster expressed Dppa3 at higher levels than the 2iL-1 cluster (Fig. 6i). Consistent with the known role of DPPA3 as an inhibitor of maintenance methylation, we found that 2iL-2 cells displayed dramatically lower DNMT1-mediated maintenance methylation fidelity, resulting in genome-wide demethylation compared to 2iL-1 cells (Fig. 6j,k)<sup>29</sup>. These results highlight the underlying intrinsic relationship between DNA methylation levels and maintenance methylation fidelity across the genome and how cell states preferentially occupy specific regions within the methylome density versus maintenance methylation fidelity landscape (Extended Data Fig. 10f,g). Further, the globally hypomethylated state in the 2iL-2 population correlates with higher expression of endogenous retroviral

elements RLTR45 and RLTR45-int, as well as higher expression of *Khdc3* (also known as *Filia*), a factor known to be involved in safeguarding genomic integrity in mES cells and preimplantation embryos<sup>51,52</sup> (Fig. 6i). Taken together, these observations suggest that reduced DNMT1-mediated maintenance methylation fidelity in the 2iL-2 population results in genome-wide DNA demethylation, an associated loss of retroviral silencing, and genome instability. Finally, we observed that, between the two 2iL subpopulations, the 2iL-2 cluster is dominated by cells in the mC<sup>Lo</sup> and Mnt<sup>Lo</sup> state and is weakly biased toward cells that have been cultured longer in 2iL conditions (Fig. 6l and Extended Data Fig. 10h). Similar to differences in the cell state transition potential between Nan<sup>Lo</sup> and Nan<sup>Hi</sup> serum-grown cells, additional work in the future will be required to investigate how differences in the maintenance methylation fidelity and DNA methylation landscape between 2iL-1 and 2iL-2 subpopulations impact their differentiation and cell fate potential.

#### Discussion

In summary, Dyad-seq is a generalized genome-wide approach for profiling all combinations of 5mC and 5hmC at individual CpG dyads. Using M-M-Dyad-seq, we discovered that DNMT1-mediated maintenance methylation fidelity is directly tied to local methylation levels, and for regions of the genome that have low methylation, specific histone marks can substantially modulate the maintenance methylation activity. Further, when combined with RNA-seq, we identified well-characterized factors, such as DPPA3, as well as other putative factors that are potentially involved in regulating the maintenance methylation fidelity of DNMT1. Similarly, using H-M-Dyad-seq and H-H-Dyad-seq, we found that 5hmCs are more commonly duplexed with 5mC in mES cells. These results highlight that variants of Dyad-seq, such as H-M-Dyad-seg and M-H-Dyad-seg, enable measurements that are currently not possible with hairpin bisulfite techniques, thereby providing deeper insights into the regulation of methylation and demethylation pathways in mammalian systems. To understand how changes in maintenance methylation fidelity during cell state transitions reprogram the methylome and transcriptome, we developed scDyad&T-seq to simultaneously quantify genome-wide methylation levels, maintenance methylation and mRNA from the same cells. Applying scDyad&T-seq to mES cells transitioning from SL to 2iL, we found that the epigenetic reprogramming was highly heterogeneous, with the emergence of four distinct cell populations that dramatically differ in 5mCpG maintenance, DNA methylation levels and gene expression. These results showed that, in addition to cell identity, DNA methylation levels and histone modifications are closely tied to how faithfully the DNA methylation maintenance machinery copies methylated cytosines from one cell generation to another. Overall, scDyad-seq is an enhancement over both scMspJI-seq and single-cell bisulfite sequencing techniques, enabling high-resolution quantification of both genome-wide 5mC levels and maintenance methylation in thousands of single cells, and when extended to scDyad&T-seq, the method can also be used to simultaneously obtain the transcriptome from the same cells (Figs. 4a,b and 5a,b and Extended Data Figs. 8a,b and 10i,j).

#### **Online content**

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41594-024-01291-w.

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## Methods

#### **Cell culture**

All cells were maintained in incubators at 37 °C and 5% CO<sub>3</sub>, mES cell line ES-E14TG2a (E14) was cultured on tissue culture plates coated with gelatin (Millipore Sigma, ES-006-B) with medium containing high-glucose Dulbecco's modified Eagle medium (Gibco, 10569044), 1% nonessential amino acid (Gibco, 11140050), 1% GlutaMAX (Gibco, 35050061), 1× penicillin-streptomycin (Gibco, 15140122) and 15% stem cell qualified serum (Millipore Sigma, ES-009-B). The medium was frozen in aliquots and used thereafter for a maximum of 2 weeks after thawing while storing it at 4 °C. Once thawed, 1 μl of β-mercaptoethanol (Gibco, 21985023) and 1 µl of LIF (Gibco, A35933) were added for every 1 ml of thawed medium. Cells were washed with 1× Dulbecco's phosphate-buffered saline (DPBS: Gibco, 14190250), and the medium was changed daily. Cells were routinely passaged 1:6 once they reached 75% confluency using 0.25% trypsin-EDTA (Gibco, 25200056). E14 cells grown under these conditions also describe the SL experimental group. Uhrf1-knockout E14 cells were cultured under the same SL conditions described above. For fluorescence-activated cell sorting (FACS), a single-cell suspension was made using 0.25% trypsin-EDTA. The trypsin was then inactivated using serum-containing medium. Afterwards, the cells were washed with 1× DPBS before being passed through a cell strainer and sorted for single cells into 384-well plates. Cell sorting was performed using gates set for forward and back scatter, doublet discrimination was performed using forward scatter width and forward scatter height, and propidium iodide was used to distinguish between live and dead cells.

K562 cells were grown in RPMI (Gibco, 61870036) with 10% serum (Gibco, 10437028) and 1× penicillin–streptomycin. When the culture reached a density of approximately 1 million cells  $ml^{-1}$ , they were split and resuspended at a density of 200,000 cells  $ml^{-1}$ . Cells were washed and FACS sorted as described for E14 cells.

#### **Decitabine treatment**

E14 mES cells were cultured as described above. Upon passage of the E14 cells, SL medium was supplemented with 0.05  $\mu$ M of decitabine. After 24 h, cells were collected using 0.25% trypsin–EDTA. The trypsin was then inactivated using serum-containing medium. The cells were washed with 1× DPBS and then resuspended in 200  $\mu$ l of DPBS. Genomic DNA was extracted using the DNeasy kit (Qiagen, 69504) according to the manufacturer's recommendations.

K562 cells were cultured as described above. Upon passage, the medium was supplemented with 0.6  $\mu$ M of decitabine or dimethyl sulfoxide (as a control). After 24 h, the cells were washed and single-cell FACS was performed as described above.

#### Transition from SL to 2iL and other medium conditions

E14 mES cells were cultured in SL conditions as described above. Upon passage, cells were resuspended in the following media depending on the condition studied. Commercial 2i medium containing LIF (Millipore, SF016-200) was used for BL, GL, 2iL and ML experiments. For 2iL, all components were used according to the manufacturer's recommendations. For GL and ML conditions, only the GSK3B inhibitor or MEK1/2 inhibitor was added, respectively. For the BL condition, no inhibitors were added. For the No condition, commercial 2i medium without LIF (Millipore, SF002-100) was used with no inhibitors added. After 24 h, the cells were washed with 1× DPBS and the medium was changed. Forty-eight hours after the initial medium switch, the cells were collected using 0.25% trypsin-EDTA, quenched using serum-containing media, washed in 1× DPBS and finally resuspended in 1× DPBS. The sample was then split in half. One half was resuspended in 200 µl of DPBS for genomic DNA extraction, as described above. The other half was resuspended in 500 µl of TRIzol reagent (Invitrogen, 15596018), and total RNA was extracted according to the manufacturer's recommendations. Experiments for each condition were performed in triplicate.

#### Dyad-seq adapters

The double-stranded Dyad-seq adapters are designed to be devoid of cytosines on the bottom strand. They contain a PCR sequence, a 4 bp UMI and a 10 bp cell-specific barcode. For Dyad-seq variants that use MspJI as a restriction enzyme (M-M-Dyad-seq and M-H-Dyad-seq), the adapters contain a random 4 bp 5' overhang and their general design is shown below.

Top oligo: 5'-NNNN [10 bp barcode] HHWHCCAAACCCACT ACACC-3'

Bottom oligo: 5'-GGTGTAGTGGGTTTGGDWDD [10 bp barcode]-3' The sequences of the 10 bp cell-specific barcode for scDyad&T-seq are provided in Supplementary Table 2. For bulk Dyad-seq, cell-specific barcodes were used as sample-specific barcodes.

For the M-M-Dyad-seq experiments described in the manuscript, a prototype of the above design was used consisting of a 3 bp UMI and an 8 bp sample-specific barcode (Supplementary Table 3).

Top oligo: 5'-NNNN [8 bp barcode] HHHCCAAACCCACTACACC-3' Bottom oligo: 5'-GGTGTAGTGGGTTTGGDDD [8 bp barcode]-3'

The sequences of the 8 bp sample-specific barcode for M-M-Dyad-seq are provided in Supplementary Table 3.

For Dyad-seq variants that use AbaSI as a restriction enzyme (H-H-Dyad-seq and H-M-Dyad-seq), the adapters contain a random 2 bp 3' overhang as shown below:

Top oligo: 5'-[10 bp barcode] HHWHCCAAACCCACTACACC-3'

Bottom oligo: 5'-GGTGTAGTGGGTTTGGDWDD [10 bp barcode] NN-3'

The sequences of the 10 bp sample-specific barcode used for H-M-Dyad-seq and H-H-Dyad-seq are provided in Supplementary Table 4.

Note that all adapters described above are unphosphorylated. The protocol for annealing the top and bottom strands to generate double-stranded adapters is described previously in scAba-seq<sup>10</sup>.

#### Synthetic oligonucleotides

To test the cutting activity of MspJI in different CpG dinucleotide modification contexts, we designed the following double-stranded DNA molecules:

Top oligo: this oligo is designed to have a defined sequence with a central CpG where the cytosine is methylated (underlined in the sequence below) followed by an UMI and an amplification handle devoid of cytosines.

5'-GAGGGCAAAGAAGATTTCCAATAATCAGAAGACCTCGGC TCCCGTTTAGTTACGTAGACATTCGCGAGCGTAGGCATCGTTGGA CTGCATACACAGTGATGAGTTWDDDWDDDWDDDWDDD WDDDGGTTAAGGTGGTAATGGATG-3'

Bottom oligo: these oligos are designed to have a defined sequence with a central CpG where the cytosine is either unmethylated, methylated or hydroxymethylated (underlined in the sequence below), followed by an UMI and an amplification handle devoid of cytosines.

5'-AACTCATCACTGTGTATGCAGTCCAACGATGCCTACGCTCG CGAATGTCTACGTAACTAAACGGGAGCCGAGGTCTTCTGATTA TTGGAAATCTTCTTTGCCCTCWDDDWDDDWDDDWDDD WDDDGGTTAAGGTGGTAATGGATG-3'

For experiments involving the bottom oligo where the central cytosine is hydroxymethylated, after annealing the top and bottom strands, the double-stranded DNA is treated with T4-BGT to glucosylate the central hydroxymethylated cytosine on the bottom strand.

#### **Bulk Dyad-seq**

For all bulk Dyad-seq experiments, 100 ng of purified genomic DNA was resuspended in 20  $\mu$ l of glucosylation mix (1× CutSmart buffer (NEB, B7204S), 2.5× UDP–glucose and 10 U T4-BGT (NEB, M0357L)) and the samples were incubated at 37 °C for 16 h. Afterwards, 10  $\mu$ l of protease mix (100  $\mu$ g protease (Qiagen, 19155) and 1× CutSmart buffer) was added to each sample, and the samples were heated to 50 °C for

5 h, 75 °C for 20 min, and 80 °C for 5 min. After this, processing of the samples differed on the basis of the version of Dyad-seq used.

For M-M-Dyad-seq and M-H-Dyad-seq, 10  $\mu$ l of MspJI digestion mix (2 U MspJI, 1× enzyme activator solution and 1× CutSmart buffer) was added to each sample and the samples were heated to 37 °C for 5 h, and 65 °C for 20 min. Next, 1  $\mu$ l of barcoded 1  $\mu$ M double-stranded adapter was added. Then, 9  $\mu$ l of ligation mix (1.11× T4 ligase reaction buffer, 4.44 mM ATP (NEB, P0756L) and 2,000 U T4 DNA ligase (NEB, M0202M)) was added to each sample, and the samples were incubated at 16 °C for 16 h.

For H-M-Dyad-seq and H-H-Dyad-seq, 10  $\mu$ l of AbaSI digestion mix (10 U AbaSI (NEB, R0665S) and 1× CutSmart buffer) was added to each sample and the samples were incubated at 25 °C for 2 h, and then heated to 65 °C for 20 min. Next, 1  $\mu$ l of barcoded 1  $\mu$ M double-stranded adapter was added. Then 9  $\mu$ l of ligation mix (1.11× T4 ligase reaction buffer, 4.44 mM ATP (NEB, P0756L) and 2,000 U T4 DNA ligase (NEB, M0202M)) was added to each sample, and the samples were incubated at 16 °C for 16 h.

After ligation, up to three barcoded libraries of the same type were pooled and all Dyad-seq versions were subjected to a  $1 \times$  AMPure XP bead cleanup (Beckman Coulter, A63881) and eluted in 40 µl of water.

M-M-Dyad-seq and H-M-Dyad-seq samples were then concentrated to a volume of 28  $\mu$ l and subjected to nucleobase conversion using the NEBNext enzymatic methyl-seq conversion module (NEB, E7125S) according to the manufacturer's recommendations except for performing the final elution step in 40  $\mu$ l of water. For M-H-Dyad-seq and H-H-Dyad-seq samples, nucleobase conversion was performed using the NEBNext enzymatic methyl-seq conversion module similar to what has been described previously<sup>16</sup>. Briefly, samples were first concentrated to a volume of 17  $\mu$ l. Then, 4  $\mu$ l of formamide (Sigma-Aldrich, F9037-100ML) was added and the samples were heated to 85 °C for 10 min before being quenched on ice. APOBEC nucleobase conversion was performed as described by the manufacturer except for two minor changes. Samples were incubated at 37 °C for 16 h, and the final elution step was performed using 40  $\mu$ l of water.

To all Dyad-seq versions, the nucleobase converted samples were subjected to one round of linear amplification. To do this, 9 µl of amplification mix was added (5.56× NEBuffer 2.1 (NEB, B7202S), 2.22 mM dNTPs (NEB, NO447L) and 2.22 µM linear amplification 9-mer: 5'-GCCTTGGCACCCGAGAATTCCANNNNNNNN-3') and the samples were heated to 95 °C for 45 s before being guenched on ice. Once cold, 100 U of high-concentration Klenow DNA polymerase (3'-5' Exo-) (Fisher Scientific, 50-305-912) was added. Then samples were quickly vortexed, centrifuged and then incubated at 4 °C for 5 min, followed by an increase of 1 °C every 15 s at a ramp rate of 0.1 °C s<sup>-1</sup> till the samples reach 37 °C, which was then held for an additional 1.5 h. Afterwards a 1.1× AMPure XP bead cleanup was performed, and the samplers were eluted in 40 µl of water before being concentrated down to 10 µl. The entire sample was then used in a linear PCR reaction by adding 15 µl of PCR mix (1.67× high-fidelity PCR mix (NEB, M0541L) and 0.67 µM Extended **RPI primer: 5'-AATGATACGGCGACCACCGAGATCTACACGTTC** AGAGTTCTACAGTCCGACGATCGGTGTAGTGGGTTTGG-3') and performing PCR as follows: initial denaturing at 98 °C for 30 s, followed by 16 cycles of 98 °C for 10 s, 59 °C for 30 s and 72 °C for 30 s, and a final extension step at 72 °C for 1 min. Next, 5 µl of the linear PCR product was amplified further in a standard Illumina library PCR reaction, incorporating a uniquely indexed i7 primer. The remaining linear PCR product was stored at -20 °C. To the final sequencing library, two 0.825× AMPure XP bead cleanups were performed with a final elution volume of 15 µl in water. The libraries were then quantified on an Agilent Bioanalyzer and Qubit fluorometer. Finally, libraries were subjected to paired-end 150 bp Illumina sequencing on a HiSeq platform.

#### **Bulk RNA-seq**

Total RNA was extracted using TRIzol (Ambion, 15596018). Fifty nanograms of total RNA was heated to  $65 \,^{\circ}$ C for 5 min and returned to ice. Thereafter, it was combined with 9 µl of reverse transcription mix (20 U RNAseOUT (Invitrogen, 10777-019), 1.11× first-strand buffer, 11.11 mM dithiothreitol, 0.56 mM dNTPs (NEB, N0447S), 100 U Superscript II (Invitrogen, 18064-071) and 25 ng of barcoded reverse transcription primer), and the sample was incubated at 42 °C for 75 min, 4 °C for 5 min and 70 °C for 10 min. Each replicate received a different barcoded reverse transcription primer. The reverse transcription primers used here have been described previously in Wangsanuwat et al.<sup>54</sup>. Afterwards, 50 µl of second-strand synthesis mix (1.2× second-strand buffer (Invitrogen, 10812-014), 0.24 mM dNTPs (NEB, N0447S), 4 U E. coli DNA Ligase (Invitrogen, 18052-019), 15 UE. coli DNA Polymerase I (Invitrogen, 18010-025) and 0.8 U RNase H (Invitrogen, 18021-071)) was added to each sample, and the samples were incubated at 16 °C for 2 h. The barcoded replicates were then pooled, and a 1× AMPure XP bead (Beckman Coulter, A63881) cleanup was performed, eluting in 30 µl of water, which was subsequently concentrated to 6.4 µl. The molecules were amplified with IVT and an Illumina sequencing library was prepared as described in CEL-seq2 (ref. 44). Libraries were sequenced on an Illumina HiSeq platform obtaining 150 bp reads from both ends.

#### scDyad&T-seq

Four microliters of Vapor-Lock (QIAGEN, 981611) was manually dispensed into each well of a 384-well plate using a 12-channel pipette. All downstream dispensing into 384-well plates was performed using the Nanodrop II liquid handling robot (BioNex Solutions). To each well, 100 nl of uniquely barcoded reverse transcription primers (7.5 ng  $\mu$ l<sup>-1</sup>) containing six-nucleotide UMI was added. The reverse transcription primers used here were previously described in Grun et al. with the exception that a UMI length of 6 was used<sup>55</sup>. Next, 100 nl of lysis buffer (0.175% IGEPAL CA-630, 1.75 mM dNTPs (NEB, N0447S), 1:1,250,000 ERCC RNA spike-in mix (Ambion, 4456740) and 0.19 URNase inhibitor (Clontech, 2313A)) was added to each well. Single cells were sorted into individual wells of a 384-well plate using FACS and stored at -80 °C. To begin processing, plates were heated to 65 °C for 3 min and returned to ice. Next, 150 nl of reverse transcription mix (0.7 URNAseOUT (Invitrogen, 10777-019), 2.33× first-strand buffer, 23.33 mM dithiothreitol and 3.5 U Superscript II (Invitrogen, 18064-071)) was added to each well, and the plates were incubated at 42 °C for 75 min, 4 °C for 5 min and 70 °C for 10 min. Thereafter, 1.5 µl of second-strand synthesis mix (1.23× second-strand buffer (Invitrogen, 10812-014), 0.25 mM dNTPs (NEB, N0447S). 0.14 UE. coli DNA Ligase (Invitrogen, 18052-019). 0.56 UE. coli DNA Polymerase I (Invitrogen, 18010-025) and 0.03 U RNase H (Invitrogen, 18021-071)) was added to each well, and the plates were incubated at 16 °C for 2 h. Following this step, 650 nl of protease mix (6 µg protease (Qiagen, 19155) and 3.85× NEBuffer 4 (NEB, B7004S)) was added to each well, and the plates were heated to 50 °C for 15 h, 75 °C for 20 min and 80 °C for 5 min. Next, 500 nl of glucosylation mix (1 U T4-BGT (NEB, M0357L), 6× UDP-glucose and 1× NEBuffer 4) was added to each well, and the plates were incubated at 37 °C for 16 h. Thereafter, 500 nl of protease mix (2 µg protease and 1× NEBuffer 4) was added to each well, and the plates were incubated at 50 °C for 3 h, 75 °C for 20 min and 80 °C for 5 min. Next, 500 nl of MspJI endonuclease mix (1× NEBuffer 4, 8× enzyme activator solution and 0.1 U MspJI (NEB, R0661L)) was added to each well, and the plates were incubated at 37 °C for 4.5 h and then heated to 65 °C for 25 min. To each well, 280 nl of uniquely barcoded 250 nM unphosphorylated double-stranded Dyad-seq adapters were added. Next, 720 nl of ligation mix (1.39× T4 ligase reaction buffer, 5.56 mM ATP (NEB, P0756L) and 140 U T4 DNA ligase (NEB, M0202M)) was added to each well, and the plates were incubated at 16 °C for 16 h. After ligation, uniquely barcoded reaction wells were pooled using a multichannel pipette, and the oil phase was discarded. The aqueous phase was incubated for 30 min with 1× AMPure XP beads (Beckman Coulter, A63881) and then subjected to standard bead cleanup with the DNA eluted in 30 µl of water. After vacuum concentrating the elute to 6.4 µl, IVT was performed as previously described in the scAba-seq

The supernatant from the RNA enrichment process contains unamplified barcoded scDyad-seg DNA molecules. A 1× AMPure XP bead cleanup was performed by incubating the samples with beads for 30 min and eluting in 40 µl of water. Samples were then concentrated to 28 µl, and nucleobase conversion was performed as described above for bulk M-M-Dvad-seq. Samples were then subjected to four rounds of linear amplification. The first round was the same as described for bulk Dyad-seq. In subsequent rounds, samples were first heated to 95 °C for 45 s before being guenched on ice. Once cold, 5 µl of amplification mix was added (1× NEBuffer 2.1 (NEB, B7202S), 2 mM dNTPs (NEB, NO447L), 2 µM linear amplification 9-mer and 10 U of high-concentration Klenow DNA polymerase (3'-5' Exo-) (Fisher Scientific, 50-305-912)). Samples were then quickly vortexed and centrifuged, and the same thermocycler conditions were used as in the first round of linear amplification. After four rounds of linear amplification, sequencing libraries were prepared the same way as described for bulk Dyad-seq. Finally, 150 bp paired-end Illumina sequencing was performed on a HiSeq platform. Sequencing metrics and associated metadata for each cell can be found in Supplementary Data 1.

scDyad-seq is performed similar to scDyad&T-seq, except the initial reverse transcription and second-strand synthesis steps are replaced with the equivalent volume of 1× NEBuffer 4. In addition, as the transcriptome is not captured, IVT is not performed and steps involving RNA enrichment and processing are omitted.

#### ChIP-seq data processing

The following published serum-grown E14 chromatin immunoprecipitation (ChIP) datasets were used in this study (GEO accessions): GSM1000123 (H3K9ac), GSE74055 (H3K9me1 and H3K27ac), GSE23943 (H3K4me3, H3K9me3, H3K27me3 and H3K36me3), GSE77420 (H3K9me2), GSM611192 (TET1 C terminus) and GSM611194 (TET1 N terminus). ATAC-seq data were obtained from GSM3399494. For all these datasets, the processed data file was downloaded from GEO and further processed if needed. For GSE74055, the bigwigCompare tool on Galaxy (version 2.1.1.20160309.6) was used for 1 kb bins to identify enriched regions compared to the input data, with bins with a log<sub>2</sub> enrichment score greater than 2 being considered enriched regions. For GSE23943, peak calling was performed using MACS2 on Galaxy, and the resulting narrow peaks file was used as enriched regions. For GSE77420, the enrichment score for H3K9me2 in serum-grown conditions was compared to the input score within 2 kb bins. Regions were considered enriched if the H3K9me2 score was greater than the input score for both replicates. For all samples, contiguous enriched regions were combined into a single region. When applicable, enriched regions were converted from mm9 to mm10 using the UCSC genome browser LiftOver tool.

#### **Bulk RNA-seq analysis**

Bulk RNA-seq data were processed as described previously with the following modification<sup>43</sup>. Reads were mapped to the RefSeq gene model based on the mouse genome release mm10, along with the set of 92 ERCC spike-in molecules (Ambion, 4456740).

DESeq2 was used for normalization and differential gene expression calling<sup>56</sup>. Gene expression differences between each condition were evaluated using adaptive shrinkage to adjust the log fold change observed<sup>57</sup>. For differential gene expression calling an adjusted *P* value cutoff of 0.01 and a shrunken log fold change cutoff of 0.75 was used. For visualization and clustering, variance stabilizing transformation was performed and batch effects from different reverse transcription primer barcodes were removed using the removeBatchEffect function in the LIMMA package<sup>58</sup>.

#### **Dyad-seq analysis**

Dyad-seq provides information on methylation or hydroxymethylation levels as well as information on 5mCpG or 5hmCpG maintenance levels. These two outputs of Dyad-seq were analyzed separately. To quantify 5mCpG maintenance levels, read 1 was trimmed to 86 nucleotides, and then exact duplicates were removed using Clumpify from BBTools. Next, reads containing the correct PCR amplification sequence and correct barcode were extracted. These reads were then trimmed using the default settings of TrimGalore. For mapping, Bismark was used in conjunction with Bowtie2 v2.3.5 to map to the mm10 build of the mouse genome<sup>46</sup>. For experiments using K562 cells, the hg19 build of the human genome was used. After mapping, Bismark was used to further deduplicate samples on the basis of UMI, cell barcode and mapping location. For libraries that were prepared using MspJI, a custom Perl script was used to identify 5mC positions on the basis of the cutting preference of MspJI, and the methylation status of the opposing cytosine in a CpG or CpHpG dyad context was inferred from the nucleobase conversion. For libraries that were prepared using AbaSI, a custom Perl script was used to identify 5hmC positions on the basis of the cutting preference of AbaSI, and the methylation status of the opposing cytosine in a CpG dyad context was inferred from the nucleobase conversion. To quantify absolute methylation or hydroxymethylation levels, the cell barcode and UMI were transferred from read 1 to read 2. Read 1 was trimmed using TrimGalore in paired-end mode. The 5' end of read 1 was clipped by 20 bases and the 3' end of read 2 was hard clipped 34 bases after detection of the PCR amplification sequence CCACATCACC-CAAACC to remove potential bias arising from enzymatic digestion and to avoid recounting unmethylated, methylated or hydroxymethylated cytosines detected at CpG dyads. The 5' end of read 2 was clipped by nine bases to minimize potential bias arising from the linear amplification random 9-mer primer. Similarly, the 3' end of read 1 was also hard clipped nine bases after the Illumina adapter was detected. Each read was mapped separately to mm10 using Bismark, and both the resulting sam files were deduplicated further using UMI, cell barcode and mapping location. The bismark methylation extractor tool was then used to extract the methylation status of detected cytosines. Next. a custom Perl code was used to demultiplex detected cytosines to the respective single cells on the basis of the associated cell barcode. Thereafter, for cytosines detected in a CpG context, information from read 1 and read 2 was merged. Then, using UMIs, duplicate cytosine coverage resulting from overlapping paired-end reads or generated during the random priming step were deduplicated. Custom codes for analyzing Dyad-seq data and the accompanying documentation is provided with this work (Supplementary Software or ref. 59). In all figures, 5mCpG percent over genomic bins or loci is calculated as the number of methylated CpG sites detected in a region divided by the total number of CpG sites detected in that region. Similarly, 5mCpG maintenance percent over genomic bins or loci is calculated as the number of symmetrically methylated CpG dyads detected in a region divided by the total number of CpG dyads detected in that region. Cells for which fewer than 25,000 CpG sites were covered were discarded from downstream DNA methylation analysis. To cluster cells on the basis of the methylome, hierarchical clustering was used and the optimal number of clusters was assigned using silhouette scores. Supplementary Data 1 provides statistics of the methylome and transcriptome quantified in individual cells.

For the strand-independent methylation model shown in Extended Data Fig. 1g,h, the genome-wide DNA methylation data from M-M-Dyad-seq was used to compute the probability of DNA methylation at each CpG site in the genome. Thereafter, assuming strand independence, the probability that a CpG dyad is symmetrically methylated, hemimethylated or symmetrically unmethylated is calculated. Results from this background model are compared to the experimental data that provide the methylation status of individual CpG dyads in the genome. CpG sites with a coverage of greater than 10 were included in this analysis.

Note that, for all the box plots shown in the figures, extended data figures and Supplementary Fig. 1, the median is shown as the center line. Hinges depict the first and third quartile, and the whisker ends extend no further than 1.5 times the interquartile range. Datapoints outside these values are shown directly on the plot or shown in the violin plot distributions.

#### scDyad&T-seq gene expression analysis

Read 2 was trimmed using the default settings of TrimGalore. After trimming, STARsolo (STAR aligner version 2.7.8a) was used to map the reads to mm10 using the gene annotation file from Ensembl. The reads were again mapped to mm10 using the transposable elements annotation file described in TEtranscripts<sup>60</sup>. Transcripts with the same UMI were deduplicated, and genes or transposable elements that were not detected in at least one cell were removed from any downstream analysis. The combined counts from genes and transposable elements for each cell were considered the expression profile of that cell and were used in downstream analysis.

The standard analysis pipeline in Seurat (version 3.1.5) was used for single-cell RNA expression normalization and analysis<sup>61</sup>. Cells containing more than 500 genes and more than 2,000 unique transcripts were used for downstream analysis. The default NormalizeData function was used to log normalize the data. The top 1,000 most variable genes were used for making principal components, and the elbow method was used to determine the optimal number of principal components for clustering. Uniform Manifold Approximation and Projection (UMAP)-based clustering was performed by running the following functions: FindNeighbors, FindClusters and RunUMAP. To identify differentially expressed genes, the FindAllMarkers or FindMarkers function was used. The Wilcoxon rank sum test was used to classify a gene as differentially expressed, requiring a natural log fold change of at least 0.1 and an adjusted *P* value of less than 0.05.

#### **Reporting summary**

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

#### Data availability

Sequencing data have been deposited in the Gene Expression Omnibus (GEO) database accession code GEO GSE197501. Source data are provided with this paper.

#### **Code availability**

Custom codes for analyzing Dyad-seq data and the accompanying documentation are provided with this work (Supplementary Software or via GitHub at https://github.com/alexchialastri/scDyad-T-seq (ref. 59)).

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#### **Author contributions**

Conceptualization, A.C. and S.S.D.; methodology, A.C. and S.S.D.; investigation, A.C., S.S., E.E.S. and S.L.; formal analysis, A.C.; writing—original draft, A.C.; writing—review and editing, A.C. and S.S.D.; funding acquisition, S.S.D.; resources, S.S.D; supervision, S.S.D.

#### **Competing interests**

A.C. and S.S.D. are co-inventors on a patent describing Dyad-seq. The remaining authors declare no competing interests.

#### **Additional information**

**Extended data** is available for this paper at https://doi.org/10.1038/s41594-024-01291-w.

**Supplementary information** The online version contains supplementary material available at https://doi.org/10.1038/s41594-024-01291-w.

**Correspondence and requests for materials** should be addressed to Siddharth S. Dey.

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 $\label{eq:constraint} Extended \, Data \, Fig. \, 1 | \, See \, next \, page \, for \, caption.$ 

## **Technical Report**

**Extended Data Fig. 1** | **Overview of Dyad-seq. (a)** In H-H-Dyad-seq, the schematic shows that hydroxymethylated cytosines at CpG dyads are detected using AbaSI digestion followed by the appropriate nucleobase conversion to interrogate the hydroxymethylation status of the cytosine on the opposing strand of the dyad. For a schematic of M-M-Dyad-seq, see Fig. 1a. (b) Schematic shows that following restriction enzyme digestion and adapter ligation, methylated cytosines can be quantified by sequencing the 'direct' or 'indirect' capture molecules. (c) Bar plots show the percentage of reads that lead to quantification of 5mC by MspJI in M-M-Dyad-seq and scMspJI-seq. (d) Bar plots show the percentage of 'direct' and 'indirect' reads that lead to unique detection of 5mC by MspJI in M-M-Dyad-

seq and scMspJI-seq. (e) Nucleotide composition downstream of MspJI cut site obtained from sequenced unique direct capture reads in M-M-Dyad-seq. (f) Summary of DNA modifications recognized by restriction enzymes and the cytosine conversion reactions performed in different variants of Dyad-seq. (g) Distribution of the percentage of symmetrical methylation at individual CpG dyads in a background model where DNA strands methylate and demethylate independently. (h) Distribution of the percentage of symmetrical methylation at individual CpG dyads obtained from experimental M-M-Dyad-seq data. In panels (g,h), only CpG sites with a coverage of greater than 10 were included in the analysis.



**Extended Data Fig. 2** | **Genomic and epigenomic contexts associated with quantification of the methylome in M-M-Dyad-seq. (a)** Genomic nucleotide composition in the vicinity of symmetrically methylated CpG dyads detected by M-M-Dyad-seq. (b) Genomic nucleotide composition in the vicinity of asymmetrically methylated CpG dyads detected by M-M-Dyad-seq. (c) Schematic of synthetic molecules with different combinations of DNA modifications at the central CpG dyad. All other nucleotides in these molecules are unmodified. The amplification handles are cytosine depleted. (**d**) Ratio of the top to bottom cytosine detected by M-M-Dyad-seq for different combinations of DNA modifications at the central CpG dyad (hemimethylated, symmetrically methylated and hemi-hydroxy methylated CpG dyads). Data is represented as mean values and the error bars represent standard deviation. Points represent individual measurements from two independent replicates.



Extended Data Fig. 3 | See next page for caption.

Extended Data Fig. 3 | M-M-Dyad-seq can accurately quantify DNA methylation levels and DNMT1-mediated maintenance methylation activity on a genome-wide scale. (a) Number of unique CpG sites detected as a function of sequencing depth for all four variants of Dyad-seq (M-M-Dyad-seq, M-H-Dyadseq, H-H-Dyad-seq, and H-M-Dyad-seq). (b) M-M-Dyad-seq shows similar 5mCpG maintenance levels as hairpin bisulfite sequencing in SL cultured mES cells<sup>8</sup>. Each point corresponds to a 100 kb bin the genome. (c) Genome-wide 5mCpG levels quantified by M-M-Dyad-seq correlates well with results obtained from bisulfite sequencing for SL cultured mESCs (Pearson r = 0.94, Spearman  $\rho = 0.94$ )<sup>24</sup>. Each point corresponds to a 100 kb bin of the genome. (d) Genome-wide 5mCpG levels quantified by H-M-Dyad-seq correlates well with results obtained from bisulfite sequencing for SL cultured mESCs (Pearson r = 0.90, Spearman  $\rho = 0.90$ )<sup>24</sup>. Each point corresponds to a 100 kb bin of the genome. (e) Ratio of coverage of CpG sites in M-M-Dyad-seq to that in whole-genome bisulfite sequencing (WGBS) over genomic regions of varying 5mCpG levels. (f) Relative coverage of CpG sites within ATAC-seq peaks in M-M-Dyad-seq and whole-genome bisulfite sequencing<sup>25</sup>. (**g**-**j**) Pie chart shows that the distribution of 5mCpG sites detected over promoters, 5' UTRs, exons, introns and 3' UTRs is similar between M-M-Dyad-seq (panels g,h) and other techniques, such as scMspJI-seq (panel i) and whole-genome bisulfite sequencing (panel j)<sup>6,26</sup>. (**k**) Violin plots show the distribution of DNA methylation over 3' UTRs, 5' UTRs, promoters, exons, introns, and CpG islands in M-M-Dyad-seq and whole-genome bisulfite sequencing. The data are plotted over 15,616 regions in M-M-Dyad-seq and 16,718 regions in WGBS for 3' UTRs, 29,516 regions in M-M-Dyad-seq and 31,090 regions in WGBS for 5' UTRs, 35,125 regions in M-M-Dyad-seq and 35,415 regions in WGBS for promoters, 170,205 regions in M-M-Dyad-seq and 217,098 regions in WGBS for exons, 188,606 regions in M-M-Dyad-seq and 203,123 regions in WGBS for introns, and 15,700 regions in M-M-Dyad-seq and 15,882 regions in WGBS for CpG islands. Data in these panels corresponds to SL cultured mESCs.



Extended Data Fig. 4 | See next page for caption.

Extended Data Fig. 4 | Global DNA methylation and transcriptome reprogramming of mESCs transitioned from SL to different media conditions after 48 hours. (a) Principal component analysis (PCA) of genomewide DNA methylation and DNA maintenance methylation levels of mESCs grown in different culture conditions for 48 hours. M-M-Dyad-seq was performed on three independent replicates for each culture condition. (b) Loss of DNA methylation after culturing mESCs in 2iL conditions for 48 hours is associated with a reduction in 5mCpG maintenance levels. Each point represents genomic tilling of 100 kb. (c) Genome-wide 5mCpG levels quantified using H-M-Dyad-seq for mESCs grown under different conditions. The violin plots are made over 2,599 genomic bins. (d) Genome-wide 5hmCpG levels quantified using H-H-Dyad-seq for mESCs grown under different conditions. The violin plots are made over 2,605 genomic bins. (e) The first two principal components show distinct transcriptomes of mESCs grown under different conditions. Bulk RNA-seq was performed in triplicate. (f) Heatmap of the expression level of genes related to *de novo* methylation, maintenance methylation, and demethylation pathways. (g,h) Gene pathway enrichment analysis for differentially expressed genes (hypergeometric test with Benjamini-Hochberg correction) was performed using Metascape<sup>33</sup>. Panel (g) shows gene sets associated with specific pathways that are highly expressed in 2iL and ML conditions, lowly expressed in No, and not differentially expressed in the No condition, lowly expressed in 2iL and ML, and not differentially expressed across SL, BL, and GL.



Extended Data Fig. 5 | See next page for caption.

Extended Data Fig. 5 | Local density of the methylome and specific histone marks alter 5mCpG maintenance activity. (a) Box plot of 5mCpG maintenance levels in 1 kb genomic bins categorized based on the number of CpG sites in the bin and the absolute methylation levels. 'Low 5mC' indicates methylation levels lower than 20%, 'Medium 5mC' indicates methylation levels between 20% and 80%, and 'High 5mC' indicates methylation levels greater than 80%. N.D. stands for "Not detected". 1 kb regions in which at least 5 unique CpG dyads are detected were included in this panel. The number of bins in each category is denoted above each boxplot. Data in this panel corresponds to mESCs grown in SL condition and profiled using M-M-Dyad-seq. (b) 5mCpG maintenance levels at fully methylated regions (FMR), lowly methylated regions (LMR), and unmethylated regions (UMR) as stratified by Stadler et al.<sup>53</sup>. Data in this panel corresponds to mESCs grown in SL condition and profiled using M-M-Dyad-seq. Dots indicate independent biological replicates (*n* = 3). (**c**-**k**) Box plots of 5mCpG maintenance levels as a function of absolute 5mCpG levels at individual loci enriched for a histone mark ('I') or a meta-region ('M') containing all enriched loci corresponding to a histone mark for all 1kb bins in the genome (c), H3K9me2 (d), H3K9ac (e), H3K36me3 (f), H3K27me3 (g), H3K27ac (h), H3K9me3 (i), H3K4me3 (j), H3K4me1 (k). Distributions for the meta-regions were obtained using bootstrapping, where resampling was performed 1,000 times per histone mark. Blue dots indicate average values found in genome-wide 1kb bins (same as data presented in panel (c)). (I) Standard error for the meta-regions in panels (c-k) and Fig. 3e.



Extended Data Fig. 6 | Genome-wide distribution of 5mCpG over various features and the distribution of 5hmC over TET1 binding sites. (a-j) Panels show the distribution of DNA methylation over 1kb bins in the genome (a), various histone modifications, such as H3K9me2 (b), H3K9ac (c), H3K36me3 (d), H3K27me3 (e), H3K27ac (f), H3K9me3 (g), H3K4me3 (h), H3K4me1 (i), and TET1binding sites (j). (k) Box plot shows the percentage of 5hmC that is paired with 5mC at CpG dyads as a function of absolute 5mCpG levels at individual loci ('T) or a meta-region ('M') (**I**,**m**) Box plot shows the percentage of 5hmC that is paired with 5mC at CpG dyads as a function of absolute 5mCpG levels at individual loci enriched for TET1 binding ('I') or a meta-region ('M') containing all enriched loci corresponding to TET1 binding sites. Distributions for the meta-regions were obtained using bootstrapping, where resampling was performed 1,000 times. Blue dots indicate average values found in genome-wide 1kb bins. M-H-Dyad-seq data was used in the analysis of panels (k-m).



Extended Data Fig. 7 | See next page for caption.

## **Technical Report**

**Extended Data Fig. 7 | scDyad&T-seq enables combined measurement of DNA methylation levels, 5mCpG maintenance levels and the transcriptome from the same cell. (a)** Genome-wide methylation and 5mCpG maintenance levels of single K562 cells treated with (+) or without (-) 0.6 μM Decitabine for 24 hours. (**b**) 5mCpHpG maintenance levels of single K562 cells treated with (+) or without (-) 0.6 μM Decitabine for 24 hours. Data is based on 134 cells treated with Decitabine and 143 cells not treated with Decitabine. (**c**) Number of genes detected as a function of transcriptome-derived reads sequenced per cell in scDyad&T-seq. Data corresponds to SL grown single E14 cells. (**d**) Coverage of CpG sites as a function of genomic DNA-derived reads sequenced per cell in scDyad&T-seq. Data corresponds to SL grown single E14 cells. (**e**,**f**) Coverage of individual CpG sites in scDyad&T-seq. Data corresponds to SL grown single E14 cells. (**g**,**h**) Coverage of individual CpG sites in M-M-Dyad-seq. Data corresponds to SL grown E14 mESCs. (**i**) Coverage of CpG sites and the number of genes detected per cell in scDyad&T-seq (blue) and scM&T-seq (red)<sup>45</sup>. The diameter of each circle corresponds to the read depth at which a cell was sequenced. Data in this panel corresponds to SL cultured E14 mESCs. (**j**) Distribution of CpG sites that are detected by different methods (scDyad&T-seq, scMspJI-seq and scBSseq) over genomic regions of varying GpC density.



Extended Data Fig. 8 | See next page for caption.

Extended Data Fig. 8 | Serum grown mESCs contain two distinct transcriptomic subpopulations with Nanog high cells exhibiting decreased DNA methylation and 5mCpG maintenance levels across a broad range of histone modifications. (a) Comparison of chromosome-wide 5mCpG strand bias scores, estimated using techniques such as scMspJI-seq, to 5mCpG maintenance percent estimated using scDyad&T-seq. The color of the data points correspond to the absolute methylation levels estimated using scDyad&Tseq. (b) Comparison of genome-wide concordance of methylation calls to 5mCpG maintenance percent estimated using scDyad&T-seq for single cells<sup>46</sup>. Concordance is defined as the fraction of reads (with at least 5 CpG sites covered) where 90% or more of the sites are methylated. The color of the data points correspond to the absolute methylation levels estimated using scDyad&T-seq. (c) Expression level of pluripotency related genes NANOG, REX1, and ESRRB in the two transcriptional clusters (NANOG high ('Nan<sup>Hir</sup>) and NANOG low ('Nan<sup>Lor</sup>)) identified in Fig. 5d using scDyad&T-seq for serum grown mESCs. (d,e) DNA methylation levels at genomic regions with 0-40% 5mCpG (d) and 40–100% 5mCpG (e) marked by different histone modifications. (f,g) 5mCpG maintenance percent at genomic regions with 0-40% 5mCpG (f) and 40–100% 5mCpG (g) marked by different histone modifications. From bulk measurements (see Fig. 3e), regions were previously categorized as less than (panels (d,f)) or greater than (panels (e,g)) 40% methylated. Data in panels (d-f) are based on 42 Nan<sup>Hi</sup> and 55 Nan<sup>Lo</sup> cells.



## **Technical Report**

Extended Data Fig. 9 | Epigenetic and transcriptional reprogramming of mESCs transitioning from SL to 2iL conditions. (a) Hierarchical clustering based on genome-wide 5mCpG levels show that cells transitioning from SL to 2iL conditions can be classified into two major groups – a 5mCpG low (mC<sup>Lo</sup>) or a 5mCpG high (mC<sup>Hi</sup>) state. (b) Hierarchical clustering based on genome-wide 5mCpG maintenance levels show that cells transitioning from SL to 2iL conditions can be classified into two major groups – a lowly maintained (Mnt<sup>Hi</sup>) or a highly maintained (Mnt<sup>Hi</sup>) state. (c) UMAP visualization of cells transiting from SL to 2iL conditions, based on the single-cell transcriptomes obtained from scDyad&T-seq, shows that cells can be classified into two broad transcriptional clusters.

The cluster names, 2iL-like and SL-like were assigned based on expression of key marker genes in mESCs grown in 2iL or SL conditions, respectively. (d) Expression of key pluripotency genes that have previously been shown to be similar between SL and 2iL culture<sup>49</sup>. (e) Expression of genes known to be transcribed at higher levels in 2iL mESCs compared to those grown in SL conditions<sup>49</sup>. (f) Expression of genes known to be transcribed at higher levels in 2iL culture<sup>49</sup>. (g) Bar plot shows the percentage of 2iL and SL grown mESCs that are assigned to the 2iL-like or SL-like transcriptional clusters. The number in the parenthesis indicates the total number of cells in that cluster.



Extended Data Fig. 10 | See next page for caption.

## **Technical Report**

Extended Data Fig. 10 | scDyad&T-seq directly relates transcriptional cell identity to demethylation dynamics in single cells transitioning from SL to 2iL. (a,b) Genome-wide DNA methylation (panel a) and 5mCpG maintenance (panel b) levels of 2iL-D3 cells in the two broad transcriptional groups – 2iLlike and SL-like – described in Extended Data Fig. 9c. (c,d) Expression levels of the pluripotency marker POU5F1 (also known as OCT4) (panel c) and early neuroectoderm lineage marker SOX1 (panel d) in 2iL-D3 cells. (e) Heatmap of differentially expressed genes between the 2iL-1 and 2iL-2 population. (f,g) Absolute DNA methylation levels *vs.* the corresponding 5mCpG maintenance levels within 100 kb bins for cells in sub-population 2iL-1 (panel f) or 2iL-2 (panel g). (h) Bar plot shows how cells cultured in the 2iL condition for different number of days are distributed between the 2iL-1 and 2iL-2 sub-populations. The number in the parenthesis indicates the total number of cells in that sub-population. (i) Panel shows the coverage of CpG dinucleotides providing information on 5mCpG maintenance (dyad coverage), and the coverage of CpG sites providing information on the absolute levels of DNA methylation in single cells (CpG coverage). The color of the data points indicate the total number of unique transcripts detected in single cells grown in SL and 2iL conditions. (j) Heatmap of 5mCpG maintenance for individual chromosomes in single cells indicates increased sensitivity in quantifying DNMT1-mediated maintenance fidelity and demethylation compared to the strand bias score obtained from methods such as scMspJI-seq. The panel also shows the culture conditions and genome-wide 5mCpG methylation levels for the same cells.

# nature portfolio

Corresponding author(s): Siddharth S. Dey

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		Our web collection on <u>statistics for biologists</u> contains articles on many of the points above.

## Software and code

Policy information about availability of computer code

Data collection Next-generation sequencing libraries were sequenced on an Illumina Hiseq 4000 and Illumina NextSeq 500 platform.

#### Data analysis Dyad-seq analysis

Dyad-seq provides information on methylation or hydroxymethylation levels as well as information on 5mCpG or 5hmCpG maintenance levels. These two outputs of Dyad-seq were analyzed separately. To quantify 5mCpG maintenance levels, read 1 was trimmed to 86 nucleotides, and then exact duplicates were removed using Clumpify from BBTools (from BBMap version 38.76). Next, reads containing the correct PCR amplification sequence and correct barcode were extracted. These reads were then trimmed using the default settings of TrimGalore (version 0.6.4\_dev). For mapping, Bismark (version 0.21.0) was used in conjunction with Bowtie2 (version 2.3.5) to map to the mm10 build of the mouse genome (Krueger et al., Bioinformatics 2011). For experiments using K562 cells, the hg19 build of the human genome was used. After mapping, Bismark was used to further deduplicate samples based on UMI, cell barcode and mapping location. For libraries that were prepared using MspJI, a custom Perl script was used to identify 5mC positions based on the cutting preference of MspJI, and the methylation status of the opposing cytosine in a CpG or CpHpG dyad context was inferred from the nucleobase conversion. For libraries that were prepared using AbaSI, a custom Perl script was used to identify 5hmC positions based on the cutting preference of AbaSI, and the methylation status of the opposing cytosine in a CpG dyad context was inferred from the nucleobase conversion. To quantify absolute methylation or hydroxymethylation levels, the cell barcode and UMI were transferred from read 1 to read 2. Read 1 was trimmed using TrimGalore in pairedend mode. The 5' end of read 1 was clipped by 20 bases and the 3' end of read 2 was hard clipped 34 bases after detection of the PCR amplification sequence CCACATCACCCAAACC to remove potential bias arising from enzymatic digestion and to avoid recounting unmethylated, methylated or hydroxymethylated cytosines detected at CpG dyads. The 5' end of read 2 was clipped by 9 bases to minimize potential bias arising from the linear amplification random 9-mer primer. Similarly, the 3' end of read 1 was also hard clipped 9 bases after the Illumina adapter was detected. Each read was mapped separately to mm10 using Bismark, and both the resulting sam files were deduplicated further using UMI, cell barcode and mapping location. The bismark methylation extractor tool was then used to extract the methylation status of detected cytosines. Next, a custom Perl code was used to demultiplex detected cytosines to the respective single cells based on the associated cell barcode. Thereafter, for cytosines detected in a CpG context, information from read 1 and read 2 were merged. Then, using

UMIs, duplicate cytosine coverage resulting from overlapping paired-end reads or generated during the random priming step were deduplicated. Custom codes for analyzing Dyad-seq data and the accompanying documentation is provided with this work (Supplementary Software and github.com/alexchialastri/scDyad-T-seq [commit id 49415639a3db50dea9c2e616fa12423261f74518]). In all figures, 5mCpG percent over genomic bins or loci is calculated as the number of methylated CpG sites detected in a region divided by the total number of CpG sites detected in that region. Similarly, 5mCpG maintenance percent over genomic bins or loci is calculated as the number of CpG dyads detected in that region. Cells for which less than 25,000 CpG sites were covered were discarded from downstream DNA methylation analysis. To cluster cells based on the methylome, hierarchical clustering was used and the optimal number of clusters was assigned using silhouette scores. Supplementary File 1 provides statistics of the methylome and transcriptome quantified in individual cells.

scDyad&T-seq gene expression analysis

Read 2 was trimmed using the default settings of TrimGalore. After trimming, STARsolo (STAR aligner version 2.7.8a) was used to map the reads to mm10 using the gene annotation file from Ensembl. The reads were again mapped to mm10 using the transposable elements annotation file described in TEtranscripts (Jin et al., Bioinformatics 2015). Transcripts with the same UMI were deduplicated and genes or transposable elements that were not detected in at least one cell were removed from any downstream analysis. The combined counts from genes and transposable elements for each cell was considered the expression profile of that cell and was used in downstream analysis. The standard analysis pipeline in Seurat (version 3.1.5) was used for single-cell RNA expression normalization and analysis (Stuart et al., Cell 2019). Cells containing more than 500 genes and more than 2,000 unique transcripts were used for downstream analysis. The default NormalizeData function was used to log normalize the data. The top 1,000 most variable genes were used for making principal components and the elbow method was used to determine the optimal number of principle components for clustering. UMAP-based clustering was performed by running the following functions: FindNeighbors, FindClusters, and RunUMAP. To identify DEGs, the FindAllMarkers or FindMarkers function was used. The Wilcoxon rank sum test was used to classify a gene as differentially expressed, requiring a natural log fold change of at least 0.1 and an adjusted p-value of less than 0.05.

Chip-seq data processing

The following published serum grown E14 ChIP datasets were used in this study (GEO accessions): GSM1000123 (H3K9ac), GSE74055 (H3K9me1 and H3K27ac), GSE23943 (H3K4me3, H3K9me3, H3K27me3, and H3K36me3), GSE77420 (H3K9me2), GSM611192 (TET1 C terminus) and GSM611194 (TET1 N terminus). ATAC-seq data was obtained from GSM3399494. For all these datasets, the processed data file was downloaded from GEO and further processed if needed. For GSE74055, the bigwigCompare tool on Galaxy (version 2.1.1.20160309.6) was used for 1 kb bins to identify enriched regions compared to the input data, with bins with a log2 enrichment score greater than 2 being considered enriched regions. For GSE23943, peak calling was performed using MACS2 on Galaxy, and the resulting narrow peaks file was used as enriched regions. For GSE77420, the enrichment score for H3K9me2 in serum grown conditions was compared to the input score within 2 kb bins. Regions were considered enriched if the H3K9me2 score was greater than the input score for both replicates. For all samples, contiguous enriched regions were combined into a single region. When applicable, enriched regions were converted from mm9 to mm10 using the UCSC genome browser LiftOver tool (web application https://genome.ucsc.edu/cgi-bin/hgLiftOver).

Bulk RNA-seq analysis

Bulk RNA-seq data was processed as described previously with the following modification. Reads were mapped to the RefSeq gene model based on the mouse genome release mm10, along with the set of 92 ERCC spike-in molecules (Ambion, 4456740) using the aln tool of BWA (version 0.7.15-r1140).

DESeq2 (version 1.26.0) was used for normalization and differential gene expression calling. Gene expression differences between each condition were evaluated using adaptive shrinkage to adjust the log fold change observed. For differential gene expression calling an adjusted p-value cutoff of 0.01 and a shrunken log fold change cutoff of 0.75 was used. For visualization and clustering, variance stabilizing transformation was performed and batch effects from different reverse transcription primer barcodes were removed using the removeBatchEffect function in the LIMMA package (version 3.42.2).

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio guidelines for submitting code & software for further information.

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Data generated in this manuscript can be openly accessed at Gene Expression Omnibus (GEO). Accession Code GEO: GSE197501. (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE197501).

The following GEO datasets, previously published/generated by other research groups, were used in this study: GSM1000123, GSE74055, GSE23943, GSE77420, GSM611192, GSM611194 and GSM3399494.

All figures are associated with raw data, with the exception of Figure 1a, 2a, 4a and Extended Data Figure 1a,b,f, 2c.

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# Life sciences study design

Sample size	No statistical calculations were performed to estimate sample size. Sample sizes were chosen to robustly estimate the quantities described in the main manuscript. For example, downsampling showed that the results were robust to sample size.
Data exclusions	All data was included except data from single cells that failed to sequence. This includes experiments described in Figures 1-6, and Extended Data Figures 1-10, and Supplementary Figure 1.
Replication	Bulk experiments were performed in triplicate. The replicates for all other experiments are provided in the main manuscript. All replication attempts were successful.
Randomization	Single mESCs and K562 cells were randomly obtained by FACS sorting. No other randomization was performed as this is not relevant to this study.
Blinding	Blinding was not relevant to this study as no manual assessment of experiments was performed. All data obtained from experiments were assessed in the same way using the analysis pipelines and scripts described above and in the main manuscript.

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## Eukaryotic cell lines

Policy information about <u>cell lines</u>	
Cell line source(s)	Mouse embryonic stem cell line ES-E14TG2a (E14, ATCC Number: CRL-1821) and the Uhrf1 KO E14 line were obtained from the Clark lab at the University of California, Los Angeles. The K562 cell line (ATCC Number: CCL-243) was obtained from the van Oudenaarden lab at Hubrecht Institute.
Authentication	The cell lines were not authenticated
Mycoplasma contamination	Cell lines were tested negative for mycoplasma.
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