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DNA methylation dynamics of the human preimplantation embryo

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Abstract

In mammals, cytosine methylation is predominantly restricted to CpG dinucleotides and stably distributed across the genome, with local, cell type-specific regulation directed by DNA binding factors1-3. This comparatively static landscape dramatically contrasts the events of fertilization, where the paternal genome is globally reprogrammed. Paternal genome demethylation includes the majority of CpGs, though methylation is maintained at several notable features4-7. While these
dynamics have been extensively characterized in the mouse, only limited observations are available in other mammals, and direct measurements are required to understand the extent to which early embryonic landscapes are conserved. We present genome-scale DNA methylation maps of human preimplantation development and embryonic stem cell (ESC) derivation, confirming a transient state of global hypomethylation that includes most CpGs, while sites of persistent maintenance are primarily restricted to gene bodies. While most features share similar dynamics to mouse, maternally contributed methylation is divergently targeted to species-specific sets of CpG island (CGI) promoters that extend beyond known Imprint Control Regions (ICRs). Retrotransposon regulation is also highly diverse and transitions from maternally to embryonically expressed, species-specific elements. Together, our data confirm that paternal genome demethylation is a general attribute of early mammalian development that is characterized by distinct modes of epigenetic regulation.

We generated genome-scale methylation maps of human preimplantation using reduced representation bisulfite sequencing (RRBS) to accommodate minimal DNA inputs. We thawed and screened morphologically normal cleavage stage embryos and blastocysts to represent early and late preimplantation, two replicates of pooled, matched Inner Cell Mass (ICM) and Trophoderm (TE), motile sperm from four unrelated, healthy donors, and fetal tissues (Extended Data Fig. 1). To estimate the time, extent, and targets of global remethylation, we generated derivation time series of three human ESC lines, collecting the primary outgrowth, first, and fifth passage per line. On average, replicates showed high reproducibility and captured 1,753,958 CpGs of methylation data at 10× coverage (Extended Data Fig. 2).

We noted two distinguishable architectures for DNA methylation across this time series: somatic-like CpG density-dependent bimodality in sperm, ESCs, and fetal tissues, and extensive CpG density-independent hypomethylation in preimplantation embryos (Fig. 1a). The substantial intermediate methylation in sperm reflects disparate repetitive element regulation, though non-repetitive sequences still fit the somatic paradigm (Fig. 1b). Almost no hypermethylated CpGs persist into cleavage, with residual methylation diminishing further into the blastocyst indicating that the embryonic landscape is rapidly established, prior to the 3rd embryonic division (Fig. 1b, Extended Data Fig. 2f). Without access to human epiblasts, in vivo characterization of global remethylation is unavailable, but cross-species comparison between mouse epiblast and human ESCs suggest that human ESCs are a reasonably proxy for postimplantation pluripotency (Extended Data Fig. 3). Notably, within primary ESC outgrowths, global remethylation is nearly complete, including for intermediately methylated repetitive elements apparent in sperm (Fig. 1b).

Despite predominant hypomethylation, erasure is not the de facto fate of all loci. Ten dynamic patterns emerged by k-means clustering of non-repetitive, 100 bp genomic tiles. 45% of sperm hypermethylated tiles retain some methylation over preimplantation, and 23% display high enough levels in cleavage embryos to be biparentally inherited and maintained (Fig. 1c). Local maintenance is significantly weighted to gene bodies: only 31% of 40,486 sperm hypermethylated intergenic tiles are ≥0.2 methylated in cleavage, compared to 57% or 59% of exons or introns, respectively (Fig. 1d, e). Frequently, gene body methylation...
extends for tens to hundreds of kilobases within a single gene (Fig. 1f). Sites of retained embryonic methylation suggest residual DNA methyltransferase activity within a phase where maintenance appears otherwise impeded\textsuperscript{12}.

We incorporated recently published RNA-seq data to interpret the relationship between DNA methylation and expression\textsuperscript{13} (Extended Data Table 1). Despite global hypomethylation, the canonical negative correlation between promoter methylation and gene expression extends to preimplantation, though the overall range in promoter methylation is contracted (Extended Data Fig. 4a). Few demethylated promoters are transcribed, suggesting that promoter demethylation largely reflects the global trend (Extended Data Fig. 4b). However, demethylated promoters are more frequently upregulated than repressed and include $POU5F1$, whose embryonic induction is essential for development\textsuperscript{14} (Extended Data Fig. 4c, d). Thus, our preimplantation data support models where distinct mechanisms may regulate global versus targeted reprogramming\textsuperscript{15}.

The global DNA methylation dynamics of the human embryo closely mirror those of mouse, with sharp transitions both into and out of preimplantation\textsuperscript{7,16} (Fig. 2a). We investigated the behavior of orthologous exons within our full mouse timeline, predicated on their human dynamics. Surprisingly, simply sorting mouse exons according to the dynamics of their human orthologues recapitulated similar trends in sperm and over preimplantation (Fig. 2b). In mouse, demethylated exons are erased early, with moderate passive depletion over cleavage (Fig. 2c, d). Alternatively, exons that maintain methylation behave similarly in mouse and are hypermethylated in both gametes (Fig. 2c, d). Intronic dynamics recapitulated these trends, but only after repetitive elements were removed from methylation estimates (Extended Data Fig. 5). Thus, both species pass through an equivalent global reprogramming with highly similar kinetics, while many orthologous regions maintain methylation that decays passively.

Transient, maternally-inherited monoallelic methylation has been previously observed in mouse\textsuperscript{5-7}. To identify candidate loci in human without access to oocytes, we searched for regions that are significantly more methylated in preimplantation embryos than in sperm, since this methylation would likely be of maternal origin (Extended Data Fig. 6a, Methods). Using these criteria, we identify 5,265 candidate maternal DMRs, including most canonical ICRs. We clustered these regions by their resolution in ESCs and found that most are preimplantation-specific and either hyper or hypomethylated in somatic tissue (Fig. 3a). The location and CpG density of DMR tiles depends on their resolution, with somatically hypomethylated DMRs substantially enriched for CGI-containing promoters, while somatically hypermethylated are more likely to be intragenic and distributed further downstream (Fig. 3a, Extended Data Fig. 6b,c). To confirm that these signatures represent true monoallelic, imprint-like maintenance, we generated RRBS libraries of two unrelated, single blastocysts and identified CpGs that could be assigned to each allele, nearly all of which were monoallelically methylated (Fig. 3b, Extended Data Fig. 7).

Given that maternal DMRs in mouse are also enriched for CGIs, we next examined the conservation of targeted loci between species (Fig. 3c). We found that 795 and 293 CGIs behave as transient DMRs in human and mouse, respectively, with substantially more
resolving to hypermethylation in human (Fig. 3d and Extended Data Table 2). Notably, Human DMRs resolving to hypomethylation are more likely to be annotated as CGIs in mouse than those resolving to hypermethylation (Extended Data Fig. 6d,e). We restricted our comparison to DMRs that share CGI status in both species and found that maternally contributed, preimplantation-specific DMRs are notably divergent, with only 7.5% found in human equivalently regulated in mouse. No obvious trend distinguished shared from species-specific signatures, though several, such as the somatic promoter of DNMT1, indicate conserved regulatory utility\(^{17}\) (Extended Data Fig. 8a). The disparity of maternal methylation targeting contrasts true ICRs, which are generally conserved\(^{18}\) (Fig. 3e, Extended Data Fig. 8b). Moreover, hypomethylation-resolving DMRs specific to one species are constitutively hypomethylated in the other, suggesting these signatures are frequently and diversely targeted (Fig. 3f).

We next investigated species-specific repetitive elements, incorporating RNA-seq data to interpret DNA methylation’s role in their regulation\(^{13}\). In human sperm, repetitive elements are frequently incompletely methylated or hypomethylated\(^{11}\). Long Terminal Repeat-containing elements (LTRs) are unexpectedly bimodal, with only a fraction hypermethylated and most displaying gametic escape that persists over preimplantation (Extended Data Fig. 9a). Long Interspersed Nuclear Elements (LINEs) are generally highly methylated in sperm, demethylated in the early embryo, with some partial remethylation in hESCs and complete hypermethylation in somatic cells (Extended Data Fig. 9b). Finally, Short Interspersed Nuclear Elements (SINEs), which represent the majority of human genome repetitive content, exhibit a uniform global behavior (Extended Data Table 3). Though intermediately methylated in sperm, their dynamics over preimplantation were otherwise similar to intergenic sequences in general (Extended Data Fig. 9c). Alternatively, diverse LTR and LINE subfamilies are dynamic, providing several examples where DNA methylation appears to coordinate specific regulatory transitions (Extended Data Tables 4 and 5).

We found that the bimodality of LTR methylation is explained by species-specific ERV1 elements. Alternatively, the ERVK family generally maintained high methylation levels, similar to observations in mouse and suggesting conserved, constitutive targeting\(^{4}\) (Fig. 4a, Extended Data Fig. 9d-g). After fertilization, expression sharply transitioned from a MalR dominated early cleavage state resembling the oocyte to one composed of ERV1 and ERVK elements in the blastocyst and ESCs (Fig. 4b). We were surprised to find that some ERV1s appear to be induced later, following global remethylation in ESCs, indicating a transition in the specific subfamilies that are expressed. Transcripts present early in preimplantation are generally from gametically hypomethylated ERV1 elements that are downregulated prior to de novo methylation (Fig. 4c, d). Moreover, for these elements, methylation and expression are negatively correlated, indicating discriminatory targeting for even extremely related sequences (Extended Data Fig. 9h). In contrast, the LTR7 subfamily is hypermethylated in sperm, rapidly demethylated, and upregulated in the blastocyst and hESCs (Fig. 4c, d). LTRs are rarely dynamic outside of this early versus late preimplantation axis: either they are already expressed in the oocyte and silenced later or are induced following demethylation and remain expressed in ESCs. Notably, this latter dynamic includes a limited number of recently emergent, unrelated subfamilies\(^{19-21}\).
Compared to LTRs, LINEs maintain higher methylation levels and only the primate-specific L1PA phylogeny is dynamically expressed (Fig. 5a, Extended Data Fig. 9c). As the only actively transposing lineage in humans, L1PA subfamilies emerged as a linear phylogeny. We found that the human-specific L1HS and its two closest ancestors, L1PA2 and L1PA3, are demethylated early, while older elements maintain higher embryonic methylation (Fig. 5b). Correspondingly, nearly all embryonic transcription could be attributed to these three youngest subfamilies (Fig. 5c, Extended Data Fig. 10a). Given the homology between subfamilies, we searched for sequence composition changes that may explain the preimplantation-specific escape of younger elements. We aligned 5′ UTRs of full-length L1PA7 to L1HS and compared sequences that demarcated demethylated L1PA3-descended elements from constitutively targeted ancestors. The largest discrete difference corresponds to an ∼130 bp deletion found within the L1PA3 lineage itself that separates older elements from the L1HS progenitor L1PA3a. Intriguingly, the presence or absence of this region isolates two disparately regulated subpopulations and marks the transition to embryonic expression (Fig. 5d, Extended Data Fig. 10b,c). This adaptation may represent a specific moment in the evolutionary progression of the L1PAs when emerging elements evaded a seemingly sequence-directed, repressive mechanism. Whether older subfamilies retain expression or transposition potential without active silencing, or if this signature reflects a vestigial, host genome adaptation that is no longer required, remains to be investigated.

We present basepair resolution maps of DNA methylation as it is dynamically reconfigured during human early development. These data identify a set of transient, maternally contributed methylation at CGI promoters, the resolution of which suggest independent modes of acquisition: male germline specific protection against methylation or de novo targeting in the oocyte of otherwise canonically unmethylated CGIs. Both are common and show poor conservation compared to classic ICRs, indicating that short-lived, parent-specific signatures are less evolutionarily constrained than those persisting after implantation. We find that repetitive element regulation is strikingly diverse in human, far more so than in mouse, with gametically hypomethylated LTR subfamilies present in the oocyte and early embryo and others sharply demethylated and induced embryonically. For LINEs, the stepwise phylogeny within the primate-specific L1PA lineage pinpoints a specific, adaptive transition. Because L1HS elements remains transpositionally active, including somatically in numerous cancers, the targeting of epigenetic silencing machinery during preimplantation may be relevant in identifying the root cause of their aberrant induction later. Understanding the regulatory principles inherent to the early embryo will improve continuing efforts to evaluate complex traits with unclear modes of epigenetic inheritance. Future work to characterize the mechanisms that impose these diversely targeted embryonic methylation patterns will illuminate their contribution to normal human development and disease.
Methods

IRB Approval

Harvard University institutional review board (IRB) and Embryonic Stem Cell Research Oversight (ESCRRO) Committee approval was obtained for both the collection and experimental use of surplus embryos resulting from infertility treatment and donated for research.

The Harvard University and Beth Israel Deaconess Medical Center institutional review boards both determined that collection, preparation, and experiments using discarded human gametes was not human subjects research and therefore did not require a full IRB review.

Consent Process

All embryos used in this study had previously been donated and stored at Harvard University. Couples donating surplus embryos for the purpose of research signed an extensive consent form at the time of their donation. These consent forms were approved by the Harvard University IRB. The authors did not have access to any identifying personal health information.

Gametes were obtained from patients who signed a consent form authorizing the experimental use of discarded gametes. These consent forms were scanned and stored in the patient's electronic medical record. The authors did not have access to any identifying personal health information.

Human sperm collection and preparation

Semen samples were collected from five healthy patients between the ages of 30-34 undergoing an evaluation for infertility. Each male was a non-smoker with a body mass index < 30 kg/m^2. Semen samples were collected by masturbation after 2 days of abstinence. A semen analysis was performed by an experienced andrologist confirming a normal sperm concentration (> 20 million/ml), normal motility (>50% motile), and normal morphology using the Kruger strict criteria (≥4% normal forms).

A PureCeption gradient solution (PureCeption 100% Isotonic Solution, Quinn's Advantage Medium with HEPES, In-Vitro Fertilization Inc) was prepared to purify each sperm sample and remove somatic cell contaminants. The gradient consisted of two layers of 1ml of PureCeption: 90% and 47% in a 15ml conical tube.

2ml of the semen sample was placed on top of the gradient. The gradient was centrifuged at 1100 rpm for 20 minutes and the supernatant removed. 1ml of sperm washing medium (Quinn's Sperm Washing Medium, In Vitro Fertilization Inc) was used to re-suspend the pellet. The sample was then centrifuged at 750 rpm for 10 minutes. The supernatant was removed. 0.1ml of the remaining pellet of sperm was transferred to a 1.7ml SafeSeal Microcentrifuge Tube (Sorenson Bioscience) and placed immediately at minus 80 degrees Celsius.
Human embryo thawing

Excess human embryos created via in vitro fertilization for the treatment of infertility were previously donated by patients undergoing assisted reproduction and stored in liquid nitrogen at minus 196 degrees Celsius.

Embryo culture dishes were set up using 60mm culture dishes (BD Falcon) and eight 30 microliter drops of Global embryo culture media (LifeGlobal) plus 15% Plasmanate (Talecris) overlaid with 10ml of oil (SAGE). Rinse dishes were also set up using 2.5ml of embryo culture media plus 15% Plasmanate. These dishes were equilibrated overnight at 37 degrees Celsius, 5% CO₂.

Both cleavage stage embryos and blastocysts were thawed using the Quinn's Advantage Embryo Thaw Kit (SAGE). This kit contains three solutions – 0.5M sucrose, 0.2M sucrose, and diluent. The straw or the vial containing the cryopreserved embryos was placed in a water bath at 30 degrees Celsius for 2 minutes. The embryos were then expelled from the straw by removing the heatsealed end or transferred from the vial using a Pasteur pipette to a clean tissue culture dish on a heated stage. The embryos were located and thawed according to the manufacturer's instructions.

Each embryo was rinsed and then placed in a single drop in the embryo culture dishes described above. Embryos were cultured in a humidified atmosphere at 37 degrees Celsius and 5% CO₂ in air.

Embryo evaluation

Cleavage stage embryos were evaluated 2-4 hours after the thawing process. Blastocysts were evaluated 18-24 hours after the thawing process. Embryos were evaluated using a Nikon Eclipse 80i microscope and images of each embryo were obtained at 40× using the Hamilton Thorne Clinical Laser Software. These images were then independently evaluated by two senior embryologists. Survival and quality were determined and an embryo was only included in this study if both embryologists agreed upon viability.

Human embryo collection

Single viable embryos were passed through several rounds of additional defined KSOM media (Millipore) under mineral oil prior to an Acidic Tyrode's Solution (Sigma) wash to dissolve the zona pellucida, somatic cellular debris, and additional sperm. Single embryos were then rinsed in clean media drops before pooling of the embryos, assessment for the absence of contaminants and snap freezing in minimal volume.

ICM/TE isolation of human and mouse embryos

Single human and mouse embryos exhibiting a clear ICM were isolated and the zona pellucida carefully removed to preserve the integrity of the blastocoel. They were then oriented using standard micromanipulation equipment (Narishige) and oriented with a clear plane available to separate the ICM from the expanded TE dissected using a Hamilton Thorne XYClone laser with 300 μs pulsing at 100% intensity. Short pulses progressed over the cleavage plane until the ICM and TE compartments were cleanly separated, at which
point both pools were separated for collection in clean microdrops, serially washed and snap frozen in minimal volume.

**Derivation of new hESC lines**

Human embryo culture and hESC derivations were carried out as previously described\(^2^6\). Briefly, cleavage stage human embryos were thawed using Quinn's Advantage Thaw Kit (SAGE) and cultured in Global medium (LifeGlobal) supplemented with 15% Plasmanate (Talecris) for 2-3 days until the blastocyst stage. For hESC derivation, the visible ICM was separated from the blastocyst by exposing the trophectoderm cells to 20-30 cell lethal laser pulses from a Xyclone laser (Hamilton Thorne Biosciences). The isolated ICM was then plated on a layer of gamma-irradiated mouse embryonic fibroblasts (MEFs) in derivation media consisting of KO-DMEM (Life Technologies), 15% KO-SR (Life Technologies), 2.5% Fetal Bovine Serum (FBS) (Hyclone), 2mM Glutamax, 1% non-essential amino acids, 50units/ml penicillin and 50μg/ml streptomycin (Life Technologies), 0.055mM β-mercaptoethanol (Life Technologies), 10ng/ml bFGF (Millipore). Ten to twelve days after ICM plating, the embryonic stem cell outgrowth (passage 0) was mechanically dispersed with half of the outgrowth plated onto a new MEF feeder layer for hESC line establishment (passage 1), and half used for methylation analysis. The human ESC colonies that resulted following plating were continuously mechanically passaged. The pluripotency of the lines was confirmed by staining for pluripotency markers and by in vitro differentiation into the 3 germ layers. The lines were registered with the Harvard University ESCRO Committee as HUES 71, HUES 72 and HUES 73.

**Isolation of mouse E6.5 epiblast and extraembryonic ectoderm**

Isolation of E6.5 epiblast and extraembryonic ectoderm was adapted from Ref. 27. Hormone-primed C57Bl6/J mice were euthanized 6 days after mating with 129S1/SvIMJ males. Deciduae were removed from the uterine horn and the full embryo extruded and placed under mineral oil in KSOM media using a pulled glass capillary. Residual maternal contaminants were removed by continuous mouth pipetting, after which the epiblast and extraembryonic ectoderm were bisected using an obliquely cut flame drawn glass capillary and the respective tissues segregated in separate KSOM drops. Visceral endoderm was removed from either epiblast or extraembryonic ectoderm by incubation in 0.5% trypsin/2.5% pancreatin (Sigma) dissolved in PBS for 20 minutes at 4°C, after which they were returned to KSOM medium drops and incubated for an additional 5 minutes at room temperature. Using a glass capillary pulled to a diameter slightly less than that of the embryo, visceral endoderm was removed by rapid aspiration and expulsion. Cleaned epiblast or extraembryonic ectoderm tissue were then serially washed through several additional drops of KSOM prior to pooling and snap freezing at minimal volume.

**Library Preparation and Sequencing**

RRBS libraries were generated as described and sequenced on an Illumina Genome Analyzer II before alignment and analysis\(^7\). The sequencing reads were aligned to the Human Genome Build 19 (hg19) for human samples and Mouse Genome Build 37 (mm9) for mouse samples using a custom computational pipeline taking into account the strain background for mouse samples. The data set was supplemented with mouse early


\(^{7}\) Smith et al. Nature, Author manuscript; available in PMC 2015 January 31.
development methylation profiles from Ref. 7 and human fetal somatic methylation profiles from the NIH Epigenomics Roadmap Project. Human ESC lines H1, H9, HUES64, and HUES6 were used for comparison with our newly derived hESC lines. Sample quality was assessed by looking at coverage numbers (i.e. number of loci present and coverage of loci) and similarity between biological replicates using pearson correlation, euclidean distance, and visual inspection of methylation histograms (see Extended Data Figs. 1 and 8)

**Estimating methylation levels**

The methylation level of each sampled cytosine was estimated as the number of reads reporting a C, divided by the total number of reads reporting a C or T. Single CpG methylation levels were limited to those CpGs that had at least 10-fold coverage. For 100bp tiles, reads for all the CpGs that were covered more than 5-fold within the tile were pooled and used to estimate the methylation level as described for single CpGs. The CpG density for a given single CpG is the number of CpGs 50bp up- and downstream of that CpG. The CpG density for a 100bp tile is the average of the CpG density for all single CpGs used to estimate methylation level in the tile.

The methylation level reported for a sample is the average methylation level across replicates. A replicate will contribute to the average only if it meets the coverage criteria within the replicate. Technical replicates were averaged before contributing to the sample average.

**Genomic features**

For mouse, high density CpG promoters (HCP), intermediate density CpG promoters (ICP), low density CpG promoters (LCP), CpG island, and DMR annotations were taken from Ref. 28. LINE, LTR, and SINE annotations were downloaded from the UCSC browser (mm9) RepeatMasker tracks. Gene annotations were downloaded from the UCSC browser (mm9) refseq track. Promoters (TSSs) are defined as 1kb up and downstream of the TSS and are parsed from refseq annotation. Promoters for all isoforms are included. Enhancer annotations were taken from Table S1 from Ref. 29. Corresponding human annotations were downloaded from the UCSC browser for hg19. Phastcons (Placental, PCE) coordinates were downloaded from the UCSC browser phastcons hg19 track. Human imprinting control regions were taken from Ref. 30.

In each case, the methylation level of an individual feature is estimated by pooling read counts for all CpGs within the feature that are covered greater than 5-fold, and levels are only reported if a feature contains at least 5 CpGs with such coverage (in contrast to 100bp tiles where no minimum number of CpGs is required). A tile is annotated as a genomic feature if any portion of the tile overlaps with the feature and thus, may be annotated by more than one feature (e.g. the same can be annotated as both a promoter and a gene).

**Gene Expression Analysis**

Raw RNA-seq data was downloaded from the gene expression omnibus from accession GSE36552 for human oocytes, zygotes, 2-cell, 8-cell, morulae, late blastocyst, hESC passage 0 and hESC passage 10, taken from Ref. 13. For all samples but morulae and late...
blastocyst, data for all single cells were pooled before alignment. Morulae and late blastocyst samples were pooled according to their respective embryos. Duplicate reads were removed before alignment. Alignment was performed using TOPHAT against human genome assembly 19 with default settings. Cufflinks was used for quantification and statistical tests of significant change using default settings.

**Retrotransposon expression analysis**

Alignment was performed using BWA against human genome assembly 19 with default settings. Repeat subfamily FPKM is the sum of the number of reads that align to each repeat element for the subfamily divided by the genome coverage of the subfamily in kilobases and normalized by the total number of reads in the sample. Repeat subfamily FPM is the same as FPKM without normalizing to the subfamily’s genome coverage. Samtools was used to find candidate repeat alignments and the CIGAR string was parsed to determine if the read overlapped with the repeat element. One L1PA6 element (chr2: 49454725-49460932) contained an unusual tall, short peak of read density and was excluded from analysis.

The L1PA consensus plot for expression was produced by using the consensus coordinates for the element from the UCSC genome browser to align the reads from the element’s alignment to the consensus sequence. Read count is the read coverage at each position in the consensus sequence divided by the total number of reads in the sample. Note that we did not use a multiple sequence alignment of the repetitive elements so this is not a fully accurate representation of expression over the consensus. For example, if the element has an insertion compared to the consensus sequence, then reads that overlap the insertion will contribute to read density on the consensus. The genomic sequence density plot was generated in a similar manner.

**Orthology between human and mouse**

The 46 mammals multiple sequence alignment downloaded from the UCSC browser was used to find orthologous regions from hg19 to mm9. For 100 bp tiles, the methylation for the corresponding region in mouse was used for comparison regardless of the length of the corresponding region. For genomic features, the methylation level of the corresponding region in mouse was used unless the mouse coordinates overlapped a mouse annotation of the same type. In the latter case, the methylation level of the corresponding feature was used instead. The 60 mammals multiple sequence alignment downloaded from the UCSC browser for mm10 was used to find orthologous regions from mouse to human by first translating the mm9 genomic feature coordinates to mm10, and then following the same procedure above.

RRBS selectively enriches for a consistent fraction of CpG dense genomic fragments within a given species, and as such provides genome-scale information versus genome-wide. In human and mouse, the coverage for CGIs is 87.4% and 89.6%, for Exons, 11.3% and 8.8%, and for Introns, 26.4% and 13.4%, respectively. In general, for both species, most features are captured at similar frequencies, but far more SINEs are captured in human than in mouse. Of the CpG islands that are captured by RRBS in human, 82.7% align to the mouse genome, and 53.9% align and share CGI status in mouse. Out of the loci that aligned to the mouse genome, mouse RRBS captured 93.6% of shared CGIs and 72.9% of aligned regions.
not annotated as CGIs in mouse. 85.2% of RRBS-captured human exons align to mouse exons and, of these, 52.7% are captured by mouse RRBS. For introns, 86.1% of human introns align to mouse introns and, of these, 34.2% are captured in mouse.

**Derivation Sliders: Comparison of hESCs to mouse pre and post implantation pluripotent tissues**

Regions with a methylation difference >0.1 between mouse ICM and mouse epiblast were used to assess the similarity of a sample to mouse ICM vs mouse epiblast. For each sample, a region scores as mouse ICM if its methylation level is more similar to mouse ICM than mouse epiblast and vice versa. Regions with methylation values that are equidistant from mouse ICM and mouse epiblast are excluded. If the slider is viewed as going from 0 (mouse ICM) to 1 (mouse epiblast), then the position of a sample is simply the proportion of regions that scored mouse ICM over the total number of regions that contribute a score.

**100 bp clustering and feature enrichment**

100 bp tiles were clustered using k-means clustering. Clusters were designated as hypermethylated (>= 0.5) or hypomethylated (<0.5) in sperm according to the cluster center. A tile and a feature were designated as overlapping if there was an overlap of 1bp or more between them. Feature enrichment scores are the negative log of the p-value calculated using the hypergeometric distribution.

**Identification of gametic differentially methylated regions in human**

Regions that have low methylation in sperm and some methylation in the early embryo likely represent instances of maternal methylation if the assumption that there is no *de novo* methylation over the cleavage divisions is true. A region is considered a maternally methylated differentially methylated region if: 1) it is significant after a two sample t-test between sperm and blastocyst with equal variance after correction for multiple hypothesis testing (q value < 0.05 using the Storey method, Ref. 31) it has a methylation difference >=0.2 higher in blastocyst than in sperm using sample means, and 3) it has a mean methylation level >=0.2 for 8 cell. These criteria were applied to both 100 bp tiles and to CGIs.

**Identification of human SNPs**

SNPs in human were downloaded from the 1000 genomes project (ftp://ftp-trace.ncbi.nih.gov/1000genomes/ftp/release/20110521/). SNPs that are not trackable by RRBS (C/T or A/G) and positions that are not covered in an *in silico* digest of hg19, i.e. covered by RRBS, were removed. The ratio [reference count/(reference count + alternative count)] was calculated for each SNP in the single human blastocyst samples and SNPs with ratios <0.2 or ratios >0.8 were removed since they likely represent homozygous alleles in the sample. We used the resulting genotypes to facilitate parent-of-origin methylation tracking.
**Parent-of-origin methylation tracking**

Reads were segregated into either the reference or alternative allele, and CpG methylation levels were called in the same manner described above. SNP normalized methylation values (Extended Data Fig. 8) are the average of the methylation values derived from each haplotype.

**L1PA Sequence Alignment**

All LINE L1HS and L1PA2-7 elements $\geq 5,500$ bp were taken from the human genome and aligned using Muscle with the following parameters: `muscle --maxiters 2 --diags`. Methylation levels are from full-length elements that are captured by RRBS. The 132 bp insert sequences common to elements L1PA3b and older were similarly aligned for Extended Data Fig. 11c.

**Extended Data**

![Extended Data Figure 1. Isolation of human preimplantation embryos for DNA methylation mapping](image)
a. Three replicates of D6 embryos, ranging in inputs from 3 to 5 embryos, were thawed, screened for proper morphology, independently isolated from the zona pellucida and pooled prior to RRBS profiling. Embryos are displayed prior to purification.

b. Two replicates of 18 and 19 human D3 cleavage stage embryos were thawed, screened for proper morphology, assessed for embryonic stage/cell number, and purified from the zona pellucida prior to RRBS profiling. Embryos are displayed prior to purification.

c. Cell numbers from thawed cleavage stage embryos ranged from 4-11 cells per embryo with a median of 8 (± 1.6 standard deviation) cells. Within each replicate, only 3 embryos demonstrated onset of compaction at the time of collection. Red line signifies the median, boxes and whiskers the 25th/75th and 2.5th/97.5th percentiles, respectively.

Extended Data Figure 2. Assembly of a genome wide DNA methylation time series through human early development and over ESC derivation

a. Summary of RRBS libraries generated, with number of biological replicates (n), number CpGs captured at 1x, 5x, and 10x, mean/median methylation values for 100 bp tiles estimated from CpGs covered at ≥5x, and mean Euclidean distance and Pearson correlation across biological replicates for these tiles.

b. Pearson correlation matrix for sperm, early embryonic, and fetal tissue samples.
c. Clustering of gametic, somatic, and preimplantation methylation profiles segregate according to their global DNA methylation landscape, with Sperm/Somatic tissue forming an adult methylation cluster that contrasts the unique epigenetic landscape present in preimplantation embryos.

d. Summary of RRBS libraries generated for ESC derivation, with number of biological replicates (n), number CpGs captured at 1×, 5×, and 10×, mean/median methylation values for 100 bp tiles estimated from CpGs covered at ≥5×, and mean Euclidean distance and Pearson correlation across biological replicates for these tiles. hESC ref refers to a reference collection of previously assayed ESC lines as part of the NIH Roadmap Epigenomics Project (Methods). Human ICM/TE were isolated through laser-assisted microdissection.

e. Pearson correlation matrix for human samples used to model ESC derivation. A consistent signature is rapidly acquired by the outgrowth stage (p0) and stably maintained over additional passages.

f. Methylation histograms for 100 bp tiles for human blastocysts and dissected ICM/TE tissue show minimal global difference, which is also observed when comparing previously assayed, immunosurgically purified mouse ICM to mechanically dissected ICM and TE.

g. Boxplots of the change in methylation (Δ methylation) for 100 bp tiles from cleavage to the blastocyst stage show passive demethylation of DNA methylation, particularly for regions that exhibit the highest methylation levels at this stage. Red line signifies the median, boxes and whiskers the 25th/75th and 2.5th/97.5th percentiles, respectively.

Extended Data Figure 3. Characterization of ESC derivation to in vivo pluripotent tissues in mouse

a. Global methylation histograms of 100 bp tiles for human ICM and p5 ESCs (rows) compared against mouse preimplantation and postimplantation embryos (ICM, E6.5 Epi), as well as with extraembryonic ectoderm (ExE) (columns), demonstrate the rapid acquisition of an epiblast-like, somatic methylation pattern upon ESC derivation in human.

b. Regions that discriminate mouse ICM from E6.5 epiblast were used to assign human ICM and ESCs to an equivalent in vivo pluripotent state for orthologous features. The position
along the axis from preimplantation (ICM) to postimplantation (Epi) pluripotency represents the proportion of regions in a set that resemble one state versus the other. For all feature sets, human ESCs rapidly establish an epiblast identity, maintaining this signature from the outgrowth stage over ensuing passages. *hESC Ref* refers to a reference collection of previously assayed ESC lines as part of the NIH Roadmap Epigenomics Project (*Methods*).

**Extended Data Figure 4. Inverse correlation between expression and promoter methylation is retained during human preimplantation**

*a*. Scatterplots of oocyte, preimplantation embryo and ESC derivation gene expression compared to promoter methylation display a canonical negative correlation, even during preimplantation where the range of promoter methylation values is contracted by global hypomethylation.

*b*. Box plots of gene expression values for genes significantly upregulated by ≥2 fold from oocyte to 8 cell compared to non-dynamic genes and categorized by promoter methylation dynamics. Genes that are both demethylated and upregulated are associated with induction from a silenced state, while those that are demethylated but not upregulated display only basal level transcription that is significantly lower than observed in promoters that are not demethylated. Bold line signifies the median, boxes and whiskers the 25th/75th and 2.5th/97.5th percentiles, respectively.
c. Gene expression dynamics following fertilization for hypermethylated sperm promoters demethylated ≥0.5 by the cleavage stage compared to the rest of promoters (Other). 123 of 541 (22%) demethylated promoters demonstrate significant upregulation (≥2-fold) compared to only 13.6% of other promoters. Moreover, the ratio of upregulated to downregulated genes in the demethylated set substantially favors zygotic activation, while other promoters include more downregulated maternal transcripts (Odds Ratio = 1.877, p = 1.344 × 10⁻⁸, hypergeometric test).

d. RNA-seq track of the pluripotency promoting, zygotically induced gene *POU5F1*, whose promoter is demethylated from 0.59 in sperm to 0.02 in cleavage, concurrent with its transcriptional induction.

![Extended Data Figure 5](image)

Extended Data Figure 5. Local retention of DNA methylation is similar for introns over human and mouse preimplantation

a. Introns are clustered according to their dynamics in human and the methylation of orthologous regions are tracked in mouse. Divergence is predominantly restricted to intermediately methylated features in human sperm that are generally hypermethylated in mouse. ∆ methylation heatmap displays the difference in methylation values between equivalent preimplantation timepoints, with ESCs in human serving as a proxy for comparison to the E6.5 epiblast in mouse. Deviation is most apparent for intermediately
methylated human sperm introns, where they are less methylated than in mouse. RMSK included: repeat masker annotated regions included.

b. When repetitive elements are removed from the calculation of intron methylation, the apparent divergence between mouse and human values is lost. Methylation and Δ methylation heatmaps are as in a. Gray denotes missing values (m.v.) where estimates for intronic methylation were exclusively derived from repetitive elements. RMSK-free: repeat masker annotated regions excluded.

c. Violin plots of the two main dynamics (maintained vs demethylated) for sperm hypermethylated introns over human and mouse preimplantation after repetitive elements are removed. As observed for orthologous exons, regions that retain high methylation throughout human preimplantation are conserved, hypermethylated in both mouse gametes, and display maintained regulation as early as the zygote stage.

Extended Data Figure 6. Genomic characterization of transient maternally contributed imprint-like regions

a. Heatmap of 100 bp tiles in mouse preimplantation identified using the same criteria as applied to human (Methods). This criteria, which assumes limited de novo methylation, identifies 2,044 tiles in mouse where methylation is ≥0.2 in both 8 cell and the ICM, there is ≥0.2 methylation difference between the ICM and sperm, and this difference is significant via t test, (q-value < 0.05). 89% of those tiles that are captured in the mouse oocyte are

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monoallelically inherited and show significant differences between the gametes by t-test, providing an empirical upper bound on the False Discovery Rate for this strategy when applied to human of ≤0.11, assuming the underlying principles of imprint regulation are the same as in mouse.

b. The proportion of 100 bp tiles, classified according to their resolution in ESCs, for each genomic feature presented in Figure 3a.

c. Cumulative density function (CDF) plot of the distance to the nearest annotated TSS for CGI DMRs that resolve to hypomethylation, intermediate/variable methylation, or hypermethylation. There is a discrepancy in genomic location between those that resolve to hypomethylation, of which a sizable fraction are in the TSS, and those that do not, which are generally enriched further downstream.

d. Boxplots of CpG density for CGI DMRs that resolve to hypomethylation, intermediate/variable methylation, or hypermethylation paired with comparable non-DMR CGIs (Somatic). Those resolving to hypomethylation have higher CpG densities than those that resolve to intermediate/variable or hypermethylation, but have slightly lower CpG density than non-DMR, constitutively hypomethylated CGIs. Alternatively, while CGIs that resolve to hypermethylation show a lower CpG density than other DMRs, they show higher density than non-DMR hypermethylated islands, suggesting some level of protection against deamination as an attribute of their uniquely hypomethylated status in the male germline.

e. Pie charts of cross species alignment and CGI status of human CGI DMRs into mouse. Those that resolve to hypomethylation are more often conserved in mouse and more frequently retain their CGI status, whereas those resolving to hypermethylation are less conserved. Moreover, intermediate/variable and hypermethylation-resolving regions that do align are less frequently retained as CGIs suggesting that hypomethylation specific to the male germline is insufficient to protect these regions from progressive deamination over time. 368, 166, and 260 CGIs comprise the hypo, intermediate/variable, and hyper methylation sets, respectively.
Extended Data Figure 7. Generation of single blastocyst libraries confirm the monoallelic behavior of putative maternal DMRs

a. Summary of two single blastocyst RRBS libraries. Number of CpGs captured at 1×, 5×, and 10×, mean/median methylation values for 100 bp tiles estimated from CpGs covered at ≥5×, and mean Euclidean distance and Pearson correlation when single blastocyst replicates are compared to the pooled blastocyst timepoint.

b. Histograms of DNA methylation for 100 bp regions captured for each single blastocyst replicate.

c. The ratio of reference allele to alternative allele for single nucleotide polymorphisms (SNPs) called as heterozygous in each blastocyst replicate.

d. For the 4,492 and 5,118 SNPs that were considered as heterozygous within each single blastocyst, 10,068 and 11,415 single CpGs could be assigned to alleles. Scatterplots depict untracked methylation values for these CpGs against their normalized methylation values, which are the average of their monoallelic methylation states.
Extended Data Figure 8. The somatic promoter of *DNMT1* is maternally methylated in human and mouse

**a.** Plots of single CpG methylation for *DNMT1*, including a CGI over the somatic promoter that behaves as a transient, preimplantation-specific DMR in both human and mouse. In mouse, hypermethylation of this island corresponds to its transcriptional readthrough and exclusion as part of an oocyte-specific isoform (*Dnmt1-o*) that is not annotated in human. Annotated CGIs and species conservation tracks are included for reference below.

**b.** Heatmap of orthologous ICR dynamics over human and mouse preimplantation. Of those that map between species and are captured by RRBS, all but one (*PEG10*) behave identically.
Extended Data Figure 9. Repetitive element regulation during human and mouse preimplantation

a. Violin plots for LTRs over human and mouse development. In human, LTRs demonstrate a bimodal distribution in sperm. Hypermethylated LTRs display a range of demethylation in the early embryo that reflects the dynamics of subfamilies. Upon ESC derivation, and within fetal tissues, LTRs become stably hypermethylated. Alternatively, during mouse preimplantation, LTRs are consistently hypermethylated in sperm and generally retain methylation over preimplantation. E6.5 Epi and E6.5 ExE refer to dissected epiblast and extraembryonic tissue from E6.5 embryos.

b. Violin plots for LINEs over human and mouse development. In human sperm, LINEs are unstably hypermethylated, with discrete populations methylated with a mean of ~0.75, ≥0.9, and a small subpopulation showing gametic escape from high methylation. Alternatively, LINEs are indiscriminately hypermethylated in mouse sperm. In both species, several populations of elements demonstrate different extents of demethylation during preimplantation, including many that retain higher levels in cleavage and only minor, passive depletion into blastocyst. Upon human ESC derivation or during mouse implantation, elements are generally remethylated, though only partially for those elements that are demethylated after fertilization. Hypermethylation is complete in fetal tissue. In human, these discrete dynamics can be attributed to the unstable methylation for L1HS-
L1PA3a subfamilies while, in mouse, the currently active L1Md_Tf and L1Md_Gf subfamilies are similarly demethylated and elements of the independently emerging L1Md_A lineage remain methylated.

c. Violin plots for SINEs highlight intermediate methylation in sperm in both species, though more so for humans. After fertilization, SINE methylation rapidly diminishes to near complete hypomethylation over preimplantation, similar to what is observed for intergenic sequence, before complete hypermethylation during ESC derivation in human or in postimplantation mouse E6.5 embryos. Taken globally, SINEs appear to be uniformly regulated regardless of subfamily, though differences in regulatory status for specific SINE elements may be reflected by their surrounding genomic context. Unfortunately, such inferences require higher genomic resolution than is currently available to distinguish the dynamics of specific integrations.

d-g. Violin plots of the four major LTR families present in mouse over the complete preimplantation timeline. ERV1 elements (d) are hypermethylated in sperm and display a range of demethylation following fertilization and prompt remethylation upon implantation. In mouse, ERVK elements (e) are emergent and largely consist of the dominating, constitutively hypermethylated IAP subfamilies. ERVL and MalR (ERVL-MalR) elements (f and g), the evolutionarily oldest mammalian LTRs, are hypermethylated in sperm and rapidly demethylated after fertilization frequently in association with their rapid zygotic induction.

h. Distribution (as boxplots) of per element expression and CpG density at different methylation levels for LTR12c demonstrates negative correlation between methylation and expression. On average, LTR12C is hypomethylated in sperm and the early embryo, but demonstrates a consistent range of values at the level of single elements, with least methylated elements contributing the most to LTR12c expression. The CpG density of these elements corresponds to their degree of hypomethylation, suggesting that escape from de novo methylation during spermatogenesis and preimplantation is maintained for specific elements over generations. Once targeted, element expression is apparently restricted and its CpG density decays correspondingly. During ESC derivation, the kinetics of LTR12c methylation is more rapid for those of lower CpG density, as evident from p0 to p5 in the ESC lines. DNA methylation in the early embryo is therefore not exclusive to the regulation of different ERV1 subfamilies, but also affects the contribution of single elements to the broader transcriptional pattern. Bold line signifies the median, boxes and whiskers the 25th/75th and 2.5th/97.5th percentiles, respectively. Expression is calculated as the number of fragments per million that align to a given element divided by its length in kb (FPKM).
Extended Data Figure 10. L1PA subfamily dynamics during human early development

a. Expression composite averaged by genomic representation for L1HS through L1PA7 from oocyte through preimplantation and ESC derivation. Dynamic expression within the L1PA phylogeny is restricted to the same subfamilies that are demethylated by cleavage. The position of each respective 5′ UTR, the functional promoter for LINEs, is highlighted in the legend. Beneath these composites is the genomic representation to the full length consensus for each annotated L1PA subfamily, which demonstrates relative equivalence of 5′ UTR representation across different subfamilies, but an increasing proportion of truncated 3′ fragments with subfamily age (Methods).

b. The frequency of CpGs within aligned L1PA subfamilies, including 5′ UTR, Orf1/2, and 3′ UTR. CpGs are primarily enriched within the 5′ UTR/promoter and become progressively CpG depleted with element age.

c. Complete composite plot of cleavage stage methylation values across aligned 5′ UTRs from L1HS through L1PA7 as in Figure 6d. The multiple sequence alignment for each subfamily to the assembled consensus is visualized below each composite, with blue corresponding to conservation, black to divergence, and white to gaps or deletions. The x-axis represents position along the 5′ UTR and a portion of ORF1 for the L1HS consensus. CpG Frequency describes the level of conservation for individual CpGs found within single elements to the consensus. The ~130 bp sequence present from L1PA7 to L1PA3b and absent from L1PA3a to L1HS is highlighted in pink, while two older sequences specific to L1PA7 are highlighted in gray.
d. Percent identity to the consensus for the extracted ~130 bp insert sequence in elements from L1PA7 through L1PA3b. Mean nucleotide identity to the consensus is 85%, with a median of 89%.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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**References**

Figure 1. Human preimplantation embryos are globally hypomethylated

a. Top. DNA methylation across 100 bp tiles for human sperm, preimplantation embryos, including the ICM and TE, ESC derivation from outgrowth to 5th passage, and somatic fetal tissues representing all germ layers. Gray highlights the average. Bottom. Boxplots of methylation at different local CpG densities. Bulls-eye signifies the median, boxes and lines the 25th/75th and 2.5th/97.5th percentiles, respectively.

b. Bar plots of 100 bp tiles segregated by non-repetitive (unique) or repetitive designation and binned by methylation status. Sp, sperm; Cl, cleavage; Bl, blastocyst; Som, somatic.

“ESC” and “Som” refer to the average of these timepoints.

c. Non-repetitive 100 bp tiles are clustered via k-means into 10 dynamics. Sperm hypermethylated sequences follow three general trajectories: persistent maintenance, incomplete or complete demethylation. Other dynamics include sperm specific hypermethylation and sperm/early embryonic hypomethylation that is de novo methylated in ESCs. Finally, 3,586 tiles are hypomethylated in sperm and ESCs but methylated in embryos, representing transient imprint-like signatures.

d. Dynamics for sperm hypermethylated, non-repetitive tiles as clustered in (c). Left heatmap, per cluster average of tiles. Right heatmap, –log_{10} p value of hypergeometric enrichment for each cluster for intergenic, exonic, intronic, CGI, or TSS annotations using sperm hypermethylated regions as the background.

e. Violin plot for sperm hypermethylated intergenic (inter), exonic, and intronic features.
f. The OBSCN gene exhibits high inter- and intra-genic methylation and an unmethylated promoter in sperm and ESCs. In cleavage embryos, a 130 kb region, highlighted in blue, remains specifically methylated while the periphery is demethylated.
Figure 2. Human preimplantation dynamics are globally similar to mouse

a. Histograms of methylation changes (Δ methylation) for 100 bp tiles across human and mouse preimplantation from fertilization (Sp to Cl) through preimplantation (Cl to Bl) to global remethylation at implantation, as measured from blastocyst to ESC in human and ICM to E6.5 epiblast in mouse (Bl to ESC/Epi).

b. Exons clustered by dynamics in human with equivalent methylation values for orthologous sequences in mouse. Δ methylation heatmap displays the difference in methylation for matched timepoints.

c. Violin plots of orthologous human sperm hypermethylated exons classified as maintained vs demethylated and measured over human and mouse preimplantation.
Figure 3. Transient maternal DMRs target a divergent set of CpG island promoters

a. Heatmap of 5,265 100 bp tiles consistent with maternally contributed monoallelic methylation (Methods). Tiles are partitioned according to their hypo (≤0.2), intermediate/variable (0.2<x<0.8) or hypermethylated (≥0.8) resolution in ESCs. Feature annotations are included as separate heatmaps.

b. Boxplots of CGI DMR methylation for two independent single blastocysts, with heterozygous SNP-linked CpGs highlighted. Within each replicate, 31 and 33 CGI DMRs contain CpGs that could be assigned to parental loci. In each case, DNA methylation is restricted to only one of the two alleles. Untracked refers to the inferred methylation status prior to haplotype segregation. Red line signifies the median, boxes and whiskers the 25th/75th and 2.5th/97.5th percentiles.

c. Single CpG track of a conserved preimplantation-specific DMR in human and mouse. Human blastocyst data includes information from the pooled sample as well as for a single blastocyst replicate (purple) with allele-tracked methylation for 10 CpGs highlighted in pink and blue. Annotated CGIs are included below.

d. Resolution of CGIs that behave as maternal DMRs in human and mouse.

e. Heatmap of orthologous ICRs over human and mouse preimplantation development.
f. Orthologous hypomethylation-resolving CGID MRs in human and mouse share only 13 equivalently regulated regions. When methylation values of mouse or human specific DMRs are tracked in the alternate species, they are constitutively hypomethylated, indicating that oogenesis targets equivalent genomic features but at species-specific sequences. PreImp refers to the average value for cleavage and blastocyst in human or 8 cell and ICM in mouse.
Figure 4. LTR subfamily dynamics are divided into early and late preimplantation phases

a. Violin plots for the four LTR families present in human over early development and ESC derivation.

b. Pie charts of LTR family expression calculated as the number of fragments per million (FPM) that align to elements with in the family.

c. Mean methylation of notable ERV1, ERVK, and MalR subfamilies. Three ERV1 subfamilies are included to represent discrete dynamics: gamete/early embryonic hypomethylation (LTR12c), constitutive methylation (HERV9-INT) and rapid demethylation (LTR7). The ERVK subfamily LTR5Hs is also demethylated.

d. Expression dynamics for the same subfamilies in (c). LTR12c is expressed early and downregulated in the blastocyst. Alternatively, LTR7 is expressed throughout, but upregulated in the blastocyst and maintained in hESCs, where it accounts for the majority of ERV1 transcripts. Like LTR7, LTR5Hs is only intermediately methylated during ESC derivation and is embryonically induced. Alternatively, the ERV1 HERV9-INT remains repressed. MLT1H2 is the prevailing MalR transcribed in the oocyte and is lost after fertilization. Expression is the fragments per million that align to subfamily elements, divided by the kb annotated as the subfamily in the genome (FPKM).
Figure 5. Emergent L1PA subfamilies escape DNA methylation-based repression during preimplantation

a. Pie charts of the LINE expression divided into the L1PA subfamily, other LINE1 and LINE2 subfamilies. Total expression is calculated as the number of fragments per million (FPM) that align to family elements.

b. Mean methylation values for the most recent L1PA subfamilies. In cleavage embryos, L1HS through L1PA3 are demethylated and maintain these levels through the blastocyst.

c. Expression dynamics for the same subfamilies in (c) over preimplantation and in ESCs. The three youngest L1PA subfamilies are induced by the 8cell stage. Expression is the number of fragments per million that align to subfamily elements, divided the kb annotated as the subfamily in the genome (FPKM).

d. Composite plot of cleavage stage methylation values across aligned 5′ UTRs in L1PA subfamilies. The composite for L1PA3 is split by the presence (red) or absence (blue) of a ∼130 bp sequence that distinguishes L1PA3b from L1PA3a and demarcates methylation values between older and newer subfamilies (highlighted in pink). Multiple sequence alignment for each subfamily to the assembled consensus is below each composite, with blue corresponding to conservation, black to divergence, and white to gaps or deletions. The x-axis represents position along the L1HS 5′ UTR and a portion of ORF1. CpG Frequency describes per CpG conservation within single elements to the consensus. Two older sequences specific to L1PA7 are highlighted in gray.
e. Boxplot of L1PAmethylation in cleavage embryos, sorted by the presence of the ∼130 bp sequence for all elements and L1PA3 specifically. Preimplantation methylation is higher for elements that contain this insert. Bold line signifies the median, boxes and whiskers the 25th/75th and 2.5th/97.5th percentiles.

f. Expression composite of full-length insert deleted L1PA3a and insert containing L1PA3b subfamilies in oocyte, 8 cell and blastocyst stage embryos. Transcriptional induction is not apparent until after fertilization and is specific to L1PA3a. Read count is the read coverage normalized by total reads (Methods).