Developmental Genome-Wide DNA Methylation Asymmetry Between
Mouse Placenta and Embryo
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Running title: Embryonic & Placental DNAme Asymmetry
<b>Summary statement:</b> The kinetics of asymmetric DNA methylation acquisition of embryo-
placenta differentially methylated regions is not a stepwise process occurring in embryonic and
extraembryonic lineages, but a prompt progression in the early post-implanted conceptus.

## 27 ABSTRACT

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29 In early embryos, DNA methylation is remodelled to initiate the developmental program but for 30 mostly unknown reasons, methylation marks are acquired unequally between embryonic and placental cells. To better understand this, we generated high-resolution DNA methylation maps 31 32 of mouse mid-gestation (E10.5) embryo and placenta. We uncovered specific subtypes of 33 differentially methylated regions (DMRs) that contribute directly to the developmental 34 asymmetry existing between mid-gestation embryonic and placental DNA methylation patterns. 35 We show that the asymmetry occurs rapidly during the acquisition of marks in the post-36 implanted conceptus (E3.5-E6.5), and that these patterns are long-lasting across subtypes of DMRs throughout prenatal development and in somatic tissues. We reveal that at the peri-37 38 implantation stages, the *de novo* methyltransferase activity of DNMT3B is the main driver of 39 methylation marks on asymmetric DMRs, and that DNMT3B can largely compensate for lack of 40 DNMT3A in the epiblast and extraembryonic ectoderm, whereas DNMT3A can only partially 41 palliate in the absence of DNMT3B. However, as development progresses and as DNMT3A 42 becomes the principal de novo methyltransferase, the compensatory DNA methylation

43 mechanism of DNMT3B on DMRs becomes less effective.

### 44 **INTRODUCTION**

45 Throughout the eutherian mammalian gestation, the placenta plays an essential role in 46 mediating maternal-embryonic exchanges of gas, nutrients and waste, and also provides the 47 developing embryo with a protective layer against adverse environmental exposures and the 48 maternal immune system (Rossant and Cross, 2001). These unique placental functions are 49 orchestrated by several distinct trophoblast cell subtypes organized in separate layers (Cross, 50 2000). The initial steps of lineage specialization of both placental and embryonic cells occur 51 promptly following fertilization during the first few embryonic cleavages as DNA methylation 52 marks are being reprogrammed (Morgan et al., 2005).

53 DNA methylation is an epigenetic mechanism that is critical in the determination of 54 lineage-specific differentiation and development, and is mainly recognized for its involvement in 55 processes such as transcriptional repression, genomic imprinting and X-inactivation (Bestor, 56 2000). DNA methylation marks are mediated by the action of DNA methyltransferases 57 (DNMTs). Establishment of new or de novo DNA methylation patterns required for cell lineage 58 determination during development is mediated by DNMT3A and DNMT3B, with cofactor 59 DNMT3L, (Okano et al., 1999, Li, 2002), whereas DNMT1 maintains heritable DNA 60 methylation patterns during cellular divisions (Leonhardt et al., 1992, Lei et al., 1996). These 61 enzymes are critical, as deletion of *Dnmt3b* or *Dnmt1* is embryonic lethal, while *Dnmt3a*-62 deficient offspring die shortly after birth (Li et al., 1992, Okano et al., 1999). During 63 gametogenesis, the acquisition of genome-wide and allele-specific methylation patterns (i.e. 64 genomic imprinting) in both oocytes and sperm is essentially due to the activity of DNMT3A 65 (Kaneda et al., 2004, Kato et al., 2007). Following fertilization, a reprogramming wave removes 66 most methylation signatures across the genome, except for imprinted regions, some types of 67 repeat sequences, as well as imprinted-like sequences, to trigger the developmental program 68 (Hirasawa et al., 2008, Howell et al., 2001, McGraw et al., 2015). Then, during the peri-69 implantation process, DNA methylation profiles are re-acquired in a sex-, cell- and tissue-70 specific manner across most parts of the genome by the combined action of DNMT3A and 71 DNMT3B. In the early stages of the *de novo* methylation wave (E4.5-E7.5), the expression of 72 Dnmt3b is more robust than Dnmt3a in the epiblast and embryonic-derived cells (Watanabe et 73 al., 2002, Auclair et al., 2014, Smith et al., 2017), with the relative expression of Dnmt3b and 74 Dnmt3a being considerably reduced in the extraembryonic ectoderm (ExE) and trophoblast

75 lineages (Smith et al., 2017, Senner et al., 2012). This discrepancy in Dnmt3a and Dnmt3b 76 expression levels coincides with the initiation of divergent DNA methylation acquisition 77 between the trophoblast and the inner cell mass of the blastocyst (Monk, 1987, Nakanishi et al., 78 2012, Oda et al., 2013, Santos et al., 2002, Fulka et al., 2004, Guo et al., 2014, Smith et al., 79 2014), a difference that becomes extremely apparent by E6.5, as the epiblast has acquired most 80 of its global DNA methylation compared to the lower-methylation state of the ExE (Auclair et 81 al., 2014, Smith et al., 2017). This divergence is a common feature across mammalian placenta, 82 as a heterogeneous and lower-methylation state compared to somatic tissues and other cell types 83 is constantly observed (Smith et al., 2017, Decato et al., 2017, Schroeder et al., 2013, Chatterjee 84 et al., 2016).

85 Although the functional role of reduced methylation levels observed across the placental genome is still not fully understood, studies suggest that it may activate transposable elements 86 87 that are typically silenced in other tissues (Chuong, 2013). DNA methylation plays an important 88 role in suppressing retrotransposons in mammalian cells, for which the activity has been 89 associated with genomic instability and disease development (Church et al., 2009, Slotkin and 90 Martienssen, 2007). Following the *de novo* methylation wave, in embryonic-derived cells from 91 the inner cell mass, transposable elements acquire higher levels of DNA methylation causing 92 transcriptional silencing, whereas in the trophectoderm-derived cells that will form the placenta, 93 these transposable elements are maintained in a relaxed methylation state and preferentially 94 expressed (Price et al., 2012, Okahara et al., 2004, Warren et al., 2015). The low-methylation 95 levels on these elements contributed to the evolution and diversification of the placenta function 96 through the regulation of gene expression by providing placenta-specific enhancers, cryptic 97 promoters and other cis-regulatory elements (Haig, 2012, Mi et al., 2000, Cohen et al., 2011, Xie 98 et al., 2013, Macaulay et al., 2011, Emera and Wagner, 2012).

99 Despite the evident distinction between embryonic and placental DNA methylation 100 levels, it remains unclear how, when and where methylation levels are acquired unequally across 101 these genomes. To better understand the developmental dynamics of epigenetic asymmetry that 102 exists between embryo and placenta, we first generated high-resolution maps of DNA 103 methylation marks using Reduced Representation Bisulfite Sequencing (RRBS) at mid-gestation, 104 when the mouse placenta is first considered mature (E10.5), to identify embryo-placenta 105 differentially methylated regions (DMRs). Then, using publicly available DNA methylation data sets and computational analyses, we defined how various categories of DMRs are established in early stages and maintained throughout development. In addition, we outlined the contribution of *Dnmt3a* and *Dnmt3b* in the acquisition and maintenance of embryonic and extraembryonic specific DMR patterns.

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### 111 METHODS

## 112 Animals and Sample Collection

Female C57BL/6 (8-10 week-old) were purchased from Harlan Sprague-Dawley Laboratories (Indianapolis, IN) and were mated with male C57BL/6 (4 months of age). Following natural mating, embryos and placental tissues were collected at E10.5 (presence of vaginal plug at E0.5). Samples were frozen immediately in liquid nitrogen and stored at -80 °C until analyzed.

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# 119 DNA Methylation Analyses

120 RRBS libraries were generated as published protocols (Boyle et al., 2012, Gu et al., 2011) 121 with our specifications (Magnus et al., 2014, McGraw et al., 2015, Legault et al., 2019). 500 ng 122 of extracted DNA (Qiagen) from placenta (male n=2, female n=2) and embryo (male n=2, 123 female n=2) samples was *MspI* digested, adaptor ligated and PCR amplified (multiplex). 124 Multiplexed samples were pooled and 100 bp paired-end sequenced (HiSeq-2000, Illumina). The 125 data analyses were done according to the pipeline established at the McGill Epigenomics 126 Mapping and Data Coordinating Centers (Magnus et al., 2014, McGraw et al., 2015) that include 127 BSMAP and methylKit. Specific parameters were chosen including 100 bp step-wise tiling 128 windows, containing a minimum of 2 CpGs per tile and a minimum 15× CpG coverage of each 129 tile per sample. The methylation level of a 100-bp tile was the result of all CpG C/T read counts 130 within the tile after coverage normalization between samples, and the methylation level reported 131 for a sample on autosomal chromosomes was the average methylation level across replicates. 132 Significant DNA methylation changes were designated as  $\pm \geq 20\%$  average differences between 133 groups of replicates and a q-value < 0.01 using the logistic regression function of methylKit 134 (Akalin et al., 2012). Direct comparisons between DNA methylation averages were done using 135 Wilcoxon-Mann-Whitney test in R. Gene ontology (GO) terms and pathway analyses for

136 RefSeqs associated to promoter-TSS were conducted using Metascape gene annotation and137 analysis resource (Zhou et al., 2019).

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# 139 Sequencing Data

Publicly available DNA methylation data sets (Auclair et al., 2014, Decato et al., 2017, Smith et al., 2014, Smith et al., 2017, Whidden et al., 2016, Hon et al., 2013) were analyzed using a custom script to intersect single CpG site methylation calls from these data sets within defined 100bp tiles associated to the embryo-placenta DMRs categories

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### 145 **RESULTS**

### 146 Increased Fluctuation in Genome-Wide DNA Methylation Levels in Placental Cells

147 To identify the overall epigenetic asymmetry that exists between placental and embryonic 148 genomes during mouse in utero development, we first established genome-wide DNA 149 methylation profiles using RRBS (McGraw et al., 2015, Magnus et al., 2014, Legault et al., 150 2019) of embryos and their corresponding placentas at mid-gestation (E10.5), the developmental 151 stage at which the mouse placenta is considered mature (Cross et al., 1994). Using this approach, 152 we quantified the DNA methylation profiles of ~1.8 million CpG sites in each sample (embryo 153 n=4, placenta n=4). We found that the accumulation of CpG methylation was very distinct 154 between the placenta and the embryo. In the placenta, we detected a greater proportion of CpGs 155 in the 0-50% methylation range and a lower proportion of CpGs in the 80-100% methylation 156 range (Figure 1A). Although we observed that a large proportion of CpGs within the examined 157 regions had no methylation marks in both the embryo and the placenta, interestingly, most CpGs 158 with partial methylation (20-50%) in placenta showed high methylation levels (>80%) in 159 embryos (Figure 1B). The divergence in global DNA methylation profiles conferred a high 160 degree of clustering between both tissue types (Figure S1). These results are consistent with 161 previous studies indicating lower levels of overall DNA methylation in extraembryonic tissues 162 (Price et al., 2012, Schroeder et al., 2013, Smith et al., 2017), reviewed in (Robinson and Price, 163 2015).

164 To further investigate the dynamics of DNA methylation between placental and 165 embryonic genomes and enable direct comparison of precise regions, we segmented the genome 166 of autosomal chromosomes into 100bp non-overlapping genomic windows (*tiles*; see methods 167 section). After removal of sex chromosomes, we identified 245 048 unique sequenced tiles 168 (referred to as All-tiles) containing 896 820 common CpGs between all placenta and embryo 169 samples and with a minimum of 15x sequencing depth. We observed a strong reduction in the 170 fraction of highly methylated tiles (80-100%) in the placenta compared to the embryo (Figure 171 1C), which correlated with a sharp increase in the number of placenta tiles in the 0-20%, 20-40%172 and 40-60% methylation range. Globally, we found that the average DNA methylation level 173 across all placental tiles was significantly lower compared to all embryo tiles (27% vs 45%, 174 p<0.0001) (Figure 1C). This overall epigenetic disparity in embryo and placenta DNA 175 methylation levels was especially noticeable when we mapped the methylation mean of All-tiles 176 with respect to regions surrounding the transcription start sites (TSS) (Figure 1D). When we 177 focused on a specific chromosome section (e.g., chr 7, 35Mb) (Figure 1E), we observed that 178 genomic segments with high DNA methylation levels in embryos were predominantly 179 hypomethylated in placenta. We also observed that gene or CpG island (CGI) poor regions had 180 consistently high methylation levels in embryos and lower methylation levels in the placenta. 181 Together, these results indicate that the mid-gestation placenta has very distinctive global DNA 182 methylation profiles compared to the embryo, and that this lower level of global DNA 183 methylation (i.e., DNA hypomethylation) across the placental genome is due to a significant 184 lower number of highly methylated ( $\geq 80\%$ ) genomic regions. We can also conclude that despite 185 the cellular heterogeneity in E10.5 placental and embryonic tissues, the vast majority of the 186 conceptus possesses specific genomic regions with either low (0-20%) or high (80-100%) levels 187 of methylation. However, the placenta genome presents an increased number of regions having a 188 broader distribution of DNA methylation (20-80%), revealing a greater diversity in methylation 189 levels across placental cell types compared to embryonic cell types.

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# 191 Embryonic and Placental DNA Methylation Divergences Across Genomic Features

To explain the developmentally divergent methylated states between the mouse embryo and the placenta, we next sought to precisely determine the genomic features revealing DNA methylation differences. We defined differentially methylated regions (DMRs) as 100bp genomic segments showing a significant difference of methylation levels between embryonic and placental samples with an absolute mean methylation difference of 20% or higher (McGraw et al., 2015, Shaffer et al., 2015, Piche et al., 2019). Using these conditions, we screened the 245

198 048 unique tiles common between all samples and identified 110 240 DMRs (~45% of All-tiles; 199 Supplementary Table 1) with tissue- and/or developmental-specific DNA methylation variations 200 between the embryo and the placenta (Figure2A; random subset of 20,000 DMRs shown). 201 Consistent with our findings (Figure 1), the majority (96.8%; n=106 712) of DMRs had lower 202 DNA methylation in the placenta (referred to as Hypo-DMRs) and only a small proportion 203 (3.2%; n=3 528) showed increased methylation levels (referred to as Hyper-DMRs) compared to 204 the embryo (Figure S2). For Hypo-DMRs, tiles mainly overlapped (96%; n=102 982) with 205 intergenic, intron, exon and promoter-TSS regions (Figure S2). For each of the genomic feature 206 categories of Hypo-DMRs, the average DNA methylation levels in the placenta were essentially 207 half of those present in the embryo (Figure 2B). As for DMRs with higher methylation levels in 208 the placenta (Hyper-DMRs), we noticed that the vast majority of these tiles (83%; n=2 929) had 209 very low methylation levels in the embryo (<20%) (Figure 2A-B). Most of Hyper-DMRs 210 overlapped intergenic, intron, exon and promoter-TSS regions (93%; n=3 272, Figure S2). We 211 then investigated the relationship between gene expression and DMR DNA methylation levels. 212 Highly expressed genes in placental tissue such as Syna (Syncytin A) (Mi et al., 2000) showed 213 overall lower methylation levels in the placenta when compared to the embryo (Figure 2D, 214 smoothed representation) (Hansen et al., 2012). Similar observations were made in the gene body 215 of Atf6b (Activating Transcription Factor 6 Beta), a gene implicated in the transcriptional 216 downregulation of Pgf (Placental Growth Factor) in response to endoplasmic reticulum stress in 217 pathological placentas (Mizuuchi et al., 2016). As for genes predominantly expressed in the 218 embryo, Mir219a-2/Mir219b, a brain-specific non-coding microRNA, showed higher 219 methylation in the placenta (Figure 2D). It was recently shown that *Mir219* was abnormally 220 expressed during pregnancy in amnion retrieved specifically from obese women (Nardelli et al., 221 2014) and, interestingly, obesity has been suggested to cause epigenetic changes (Lillycrop and 222 Burdge, 2011). Another example of placenta Hyper-DMRs is Sox6 (SRY-Box 6), which is 223 implicated in the terminal differentiation of muscle (Figure 2D) (Kamachi and Kondoh, 2013). 224 These observations point to an association between tissue-specific DNA methylation levels of 225 DMRs and biological functions. Gene ontology enrichment analyses further support that Hypo-226 DMRs were strongly associated to germline functions and reproduction (e.g., male and female227 gamete generation, reproduction, piRNA metabolic process, meiotic cell cycle, germ cell 228 development) (Figure 2C). As for Hyper-DMRs, top biological processes were mostly associated

with developmental and differentiation processes (*e.g.*, regionalization, embryo development, pattern specification process, skeletal system, head development) (Figure 2C). Fittingly, the biological functions were completely divergent between Hypo- and Hyper-DMRs. Altogether, these results denote that Hypo- and Hyper-DMRs are present across genomic features between mid-gestation embryo and placenta, and that these DNA methylation divergences are implicated in promoting/repressing specific processes during embryonic and placental development.

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### 5 Presence of Distinctive DMR Categories between Mid-Gestation Embryo and Placenta

237 Amongst DMRs, our analyses also suggest the presence of particular DMR categories 238 based on their level of DNA methylation in the embryo and placenta (Figure 2A). By defining 239 subsets of DMRs and establishing their dynamic properties between tissues, we might better 240 understand the genome-wide asymmetry in DNA methylation levels observed between the 241 embryo and the placenta. To do so, we first clustered DMR-associated tiles in 6 different 242 categories based on their range of low, mid and high methylation level (Low; <20%, Mid;  $\geq 20$  to 243 <80%, High;  $\geq$ 80%) in the embryo and the placenta, and followed the DMR category transitions 244 between both tissues. We observed that DMRs with High-levels of methylation in the embryo 245 overlapped with a large proportion of DMRs that showed Mid-levels of methylation in the 246 placenta (Figure 3A,B; High-Mid n=72 715), whereas only a fraction corresponded to DMRs 247 with Low-levels of methylation in the placenta (Figure 3A,B; High-Low n=1 889). As for DMRs 248 with Mid-levels of methylation in embryo, the largest part remained in that same Mid-levels 249 category in the placenta (Figure 3A; Mid-Mid n=23 640). Nonetheless, a portion of these 250 embryonic Mid-levels DMRs were directed to DMRs with either Low- (Figure 3A; Mid-Low 251 n=9 055) or High- (Figure 3A; Mid-High n= 21) levels of methylation in the placenta. Finally, 252 DMRs with Low-levels of methylation in the embryo all showed Mid-levels of methylation in 253 the placenta (Figure 3A; Low-Mid n=2 920). Thus, when we subdivide our DMRs into 254 distinctive categories, we uncover that embryonic cells possess a large proportion of DMRs of 255 High-level ( $\geq$ 80%) of DNA methylation, which remain potentially static in embryonic tissues, 256 whereas the vast majority of these DMRs have inferior and wide-range DNA methylation levels 257 within placental tissue.

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#### 259 Specific Genomic Features Associated with DMR Categories

260 We next aimed to determine if the genomic distribution of embryo-placenta DMR 261 categories was associated with distinct genomic features. First, by classifying by genomic 262 annotations, we observed that DMRs with reduced methylation levels in the placenta (High-Mid, 263 High-Low, Mid-Low) were prevailingly found in intergenic regions (>50% of tiles) (Figure 3C), 264 whereas DMR categories with equivalent (Mid-Mid) or superior methylation level (Mid-High, 265 Low-Mid) in the placenta were more frequent in genic associated regions (>50% of tiles). 266 However, divergence between DMR categories was observed when we performed ontology 267 analyses on promoter regions (Figure S3), as each DMR subtype clearly showed distinct 268 biological functions. For example, High-Mid DMRs top functions were related to genes (e.g., 269 Aszl, Tnp1, Piwil2) associated with male gamete generation, piRNA metabolic process and male 270 meiotic nuclear division, whereas those of the Low-Mid category were mostly associated with 271 developmental aspects (e.g., skeletal system development, regionalization, nervous system 272 development). Since we know that in placenta, activation of retrotransposon-derived genes is 273 interrelated with low DNA methylation levels (Macaulay et al., 2011, Reiss et al., 2007, Cohen 274 et al., 2011), we next assessed how DMR categories overlapped with major types of 275 retrotransposons (LINE; long interspersed nuclear elements, SINE; short interspersed nuclear 276 element, and LTR; long terminal repeats). Out of the DMR categories, those with High-levels of 277 methylation in the embryo and either Mid- or Low-levels in the placenta (High-Mid and High-278 Low) showed the most enriched overlap with retrotransposons, with 92% and 96% respectively 279 (Figure 3D). DMR categories associated with gain of methylation in the placenta (Mid-High and 280 Low-Mid) showed the least overlap with retrotransposons, especially the Low-Mid subtype. This 281 highlights that during the *de novo* acquisition of DNA methylation patterns, DMRs with High-282 levels of methylation in the embryo and lower levels in the placenta are almost exclusively 283 within retrotransposons-associated sequences, whereas DMRs with Low-levels of methylation in 284 the embryo and higher DNA methylation in the placenta are preferentially outside 285 retrotransposons-associated sequences. Finally, we investigated the proximity of the DMR 286 categories in regards to CpG rich (CpG islands; CGI), neighbouring (shore; < 2kb away from 287 CGIs, shelf; 2-4kb away from CGIs) and distant (open sea; > 4kb away from CGIs) regions 288 (Figure 3E). We observed that most DMR categories are depleted from CGIs and are mostly 289 found in open sea regions. In contrast, ~60% of tiles in Low-Mid DMRs overlapped with CGI, 290 shores and shelves, revealing that the acquisition of *de novo* methylation for these genomic

fragments in the extraembryonic lineage preferentially targets sequences inside or surroundingCGIs.

Altogether, these results indicate that the asymmetry within DMR categories, based on their methylation levels in the embryo and the placenta, can be associated to specific biological functions and genomic-derived features (e.g., CpG, retrotransposon contents). This is particularly apparent for DMRs with High-levels in the embryo (High-Mid, High-Low) and those with higher methylation levels in the placenta (Low-Mid).

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# DMR Categories are Established During the De Novo Methylation Wave and Maintained Throughout Development

301 To gain insights into the kinetics of lineage-specific DMR establishment between mid-302 gestation embryo and placenta, as well as their status during development, we assessed the levels 303 of methylation associated with tiles for each DMR category as a function of their developmental 304 stage. Publicly available sequencing data (Smith et al., 2012, Whidden et al., 2016) were 305 analyzed using our custom script to generate 100bp tiles, and the DNA methylation levels for 306 each tile were calculated. In E3.5 blastocysts, when the mouse genome is mostly depleted from 307 DNA methylation marks, we observed low global DNA methylation levels (average <20%) for 308 all DMR subtypes, with similar median levels between the committed cells of the inner cell mass 309 (ICM) and trophectoderm lineages (Figure 4). DNA methylation levels tended to be higher in the 310 trophectoderm for regions falling in the Mid-High DMRs, although measurements are based on 311 very few DMRs for this specific category (n=21, Figure 3B, Mid-High). Since global DNA 312 methylation is re-acquired in the next few subsequent developmental stages, we then asked 313 whether the contrast in DNA methylation levels associated with the various DMR categories at 314 mid-gestation would already be present between E6.5 epiblast and extraembryonic ectoderm 315 (ExE) cell lineages, layers that are mostly composed of homogeneous and undifferentiated cell 316 populations. For all DMR categories, DNA methylation levels in the E6.5 epiblast and ExE 317 already showed similar pattern trends to those observed in the E10.5 embryo and placenta 318 (Figure 4, Table S2). Interestingly, for all DMR categories we observed slightly higher DNA 319 methylation levels in E6.5 ExE compared to E10.5 placenta (Figure 4, Table S2). However, 320 when measured in the subsequent developmental stages (E10.5, E11.5, E15 and E18), levels 321 associated with each category stabilized and remained in the same methylation range. This is true

322 for all categories, except for the Mid-High DMRs where methylation levels are reduced by  $\sim 20\%$ 323 at E18. As for the E6.5 epiblast DNA methylation profiles, they closely matched those observed 324 in the E10.5 embryo, except again for the Mid-High DMRs. For each of the DMR subtypes, the 325 DNA methylation levels were largely stable and persisted across other time points (E6.5, E10.5, 326 E11.5) in embryonic cells. Although no public DNA methylation data were available for whole 327 embryos at later stages, when we overlapped tiles associated with E10.5 DMR categories with 328 data from differentiated tissues of adult mice, the global DNA methylation profiles closely 329 matched those for the majority of DMR categories (High-Mid, High-Low, Mid-Mid and Mid-330 Low) (Figure 4, Figure S4). We conclude that the various DMR categories observed at E10.5 are 331 established during the embryonic and extraembryonic lineage-specification processes occurring 332 during the peri-implantation wave of *de novo* methylation, and that these DNA methylation landscapes are widely retained throughout embryo and placenta development. Furthermore, for 333 334 most DMR categories, the DNA methylation levels observed in developing embryos are long-335 lasting and conserved throughout somatic cell differentiation.

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## 337 Dnmt3a- or Dnmt3b-Deficiency Alters Proper Establishment of DMR-Associated Patterns

338 Since the combined activity of DNMT3A and DNMT3B is essential for proper 339 establishment of DNA methylation profiles and normal development, we next sought to define 340 the contribution of each enzyme in the *de novo* establishment of DNA methylation in subtypes of 341 DMRs in embryonic and extraembryonic cell lineages. To do so, we used DNA methylation data 342 from publicly available datasets (whole genome bisulfite sequencing and RRBS) at E6.5 343 (epiblast and ExE) and E8.5 (embryo) with various inactive forms of *Dnmts*. First, when we 344 overlapped our All-tiles subset, we observed a substantial reduction ( $p \square 0.0001$ ) in average DNA 345 methylation in absence of DNMT3A (37.81%) or DNMT3B (31.48%) in the E6.5 epiblast 346 compared to wild-type (44.81%), whereas in the E6.5 ExE such a comparable loss was only associated with a *Dnmt3b*-deficiency (wt: 34.38% vs *Dnmt3b*<sup>-/-</sup>: 19.33%, p $\Box$  0.0001) (Figure S5). 347 348 Similarly, in the E8.5 embryo, a significant reduction in average DNA methylation level was 349 measured with lack of Dnmt3a or Dnmt3b expression compared to wild-type (wt: 47.5%;  $Dnmt3a^{--1}$ : 43.6%;  $Dnmt3b^{--1}$ : 34%, p $\square$ 0.0001) (Figure S5). For most DMR subtypes, absence of 350 351 DNMT3A caused modest or no reduction on overall DNA methylation levels in E6.5 epiblast 352 and ExE (Figure 5A and S6, Table S3). Intriguingly, *Dnmt3a*-deficiency caused a slight gain of

methylation (wt: 76.9% vs  $Dnmt3a^{-/-}$ : 81.5%, p $\Box$ 0.0001) level in the E6.5 epiblast for High-Mid 353 354 DMRs, suggesting a potential overcompensation effect by DNMT3B in the epiblast on this 355 specific DMR subtype. In comparison, *Dnmt3b*-depletion led to a substantial loss of average 356 methylation levels in all DMR categories in the E6.5 epiblast and ExE, except for epiblast High-357 Mid DMRs (Figure 5A, Table S3) for which a compensatory mechanism provided high levels of methylation (wt: 76.9% vs Dnmt3b<sup>-/-</sup>: 71.6%). The compensatory mechanism on High-Mid 358 359 DMRs in response to Dnmt3b-deficiency was ineffective in the E6.5 ExE. When we focused on 360 promoter-TSS for each DMR categories, we observed again that loss of methylation for these 361 regulatory regions was principally associated with *Dnmt3b*-deficiency (Figure 5B). The 362 promoter-TSS regions associated with High-Mid DMRs retained relatively high methylation levels for either Dnmt3a<sup>-/-</sup> or Dnmt3b<sup>-/-</sup> epiblast samples, demonstrating a robust and 363 364 compensatory de novo methylation mechanism. To further underline the impact of DNMT3A or 365 DNMT3B on the *de novo* methylation of DMRs, we measured methylation levels for DMRs 366 selected from our gene enrichment analyses (Fgb, P2rx7, Pcyt2, Etnppl, Ralgds, Lrp5) and other 367 genomic segments covered by multiple tiles (*Cxxc1*, *Fcgrt*, *Lamp5*, *Mbd1*, *Lphn1*, *Pick1*, *Irf1*) 368 (Figure S7 & S8) in Dnmt3a- or Dnmt3b-deficient epiblast and ExE. In line with our global 369 observations, for most DMR-associated tiles, a Dnmt3b-deficiency in E6.5 epiblast or ExE 370 caused a more severe loss of methylation compared to lack of Dnmt3a. However, for some 371 regions in the E6.5 epiblast (e.g., *Pick1*, *Irf1*), *Dnmt3a<sup>-/-</sup>* methylation levels were inferior to those of *Dnmt3b<sup>-/-</sup>*. Noteworthy, for most DMRs, lack of co-factor *Dnmt3l* affected DNA methylation 372 373 levels to greater extent than *Dnmt3a*-deficiency (Figure S6). Overall, we show that of DNMT3A 374 and DNMT3B participate in the establishment of the asymmetric methylation patterns associated 375 with the various DMR categories in both embryonic and extraembryonic cells, with DNMT3B 376 being the principal contributor in both cell lineages during the *de novo* reprogramming wave as it 377 can compensate almost entirely for the absence of DNMT3A.

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# 379 Decline in Compensatory DNA Methylation Mechanisms in Response to Dnmt3a- or Dnmt3b380 Deficiency.

As embryonic development progresses from E6.5 to E8.5, our data suggest that lack of DNMT3A or DNMT3B further reduces global methylation levels of DMRs (High-Mid, High-Low, Mid-Low and Mid-Mid), evoking that absence of either enzymatic activity causes an

384 additive and extended effect (Figure 5A-B). To further define the robustness in compensatory 385 mechanism between DNMT3A or DNMT3B in the establishment and maintenance of DNA 386 methylation on DMRs categories, we focused on High-Mid promoter-TSS- associated DMRs as 387 they have the highest DNA methylation levels and require the most *de novo* methyltransferase 388 activity. Methylation differences of  $\geq 20\%$  between wild-type and *Dnmt3a*- or *Dnmt3b*-deficient 389 samples were considered as regions showing substantial lack of compensation. In agreement 390 with our results (Figure 5A-B), we observed that at E6.5, DNMT3B can compensate almost 391 entirely for DNMT3 Aloss by maintaining methylation levels on most promoters-TSS associated 392 tiles in the epiblast (410/437 = 93.8%) and ExE (558/602 = 92.7%), whereas DNMT3A can only 393 partially alleviate the lack of DNMT3B in the epiblast (315/410 = 78.6%) and ExE (244/524 =394 46.6%) (Figure 6A-B). As embryonic cell lineages development evolves between E6.5 and E8.5, 395 global methylation levels increase on promoter-TSS of High-Mid DMRs. However, we detected 396 a sharp decline in the compensatory mechanism to rescue the *de novo* and/or maintenance of 397 methylation levels of promoter-TSS DMRs in the Dnmt3a- (524/627 = 80.9%) or Dnmt3b-398 (256/622 = 41.2%) deficient E8.5 embryos. This is further highlighted when we focus on a 399 subgroup of 314 promoter-TSS associated tiles overlapping all of 9 data sets (Figure 6C), 400 including gene promoters (e.g., Asz1, Catsper1, Ccdc42, Dmrtb1, Piwil2, Rpl10l, Sox30, Sycp1, 401 Tnp1, Ttll1, Zfn42) related to our top enriched biological functions (i.e., piRNA, gamete 402 generation and germ cells, meiotic nuclear division). Although E8.5 Dnmt3a- or Dnmt3b-403 deficient ExE data was not available, our data suggest that greater compensation failure would 404 also be observed in this tissue. Thus, embryonic-placental DMRs need the combined action of 405 DNMT3A DNMT3B to both establish and maintain proper asymmetric levels during early 406 development as compensatory DNA methylation mechanisms during the de novo wave of 407 methylation decline overtime and fail to overcome methyltransferase shortage in later 408 development stages.

409

### 410 **DISCUSSION**

With recent breakthroughs in high-throughput sequencing, we now have a better understanding of the dynamic of DNA methylation erasure occurring in early cleavage stage embryos following fertilization. However, the discrepancies in acquisition of genome-wide DNA methylation patterns in early post-implantation embryonic and extraembryonic cell lineages 415 remain to be methodically delineated. To address this issue and further our understanding of the 416 DNA methylation asymmetry that guides the developmental trajectory of embryonic and 417 extraembryonic cell lineages, we established genome-wide DNA methylation profiles of mouse 418 embryo and placenta at mid-gestation, and analyzed various publicly available developmental 419 stage specific embryo and placenta DNA methylation datasets. Using this strategy, we uncovered 420 that 45% of the genomic regions analyzed differ in DNA methylation status ( $\geq 20\%$ ) between 421 mid-gestation embryo and placenta, and that these DMRs can be further divided into categories 422 based on their levels of DNA methylation (Low; <20%, Mid;  $\geq$ 20 to <80%, High;  $\geq$ 80%) in the 423 embryo and placenta. We show that the embryo and placenta acquire specific DMR categories 424 during the early stage of the *de novo* DNA methylation wave, and that these DMRs persist 425 throughout prenatal development, as well as into somatic adult tissues. Furthermore, we show 426 that Dnmt3b primarily drives the divergence in DNA methylation levels associated with these 427 specific DMRs and that *de novo* methyltransferase activity of *Dnmt3b* can almost entirely 428 compensate for lack of *Dnmt3a* in the asymmetric establishment of embryonic and placental 429 DMRs, but that Dnmt3a can only partially palliate the absence of Dnmt3b. However, with 430 developmental progression, this compensatory DNA methylation mechanism becomes less 431 effective.

432 Our results indicate that the kinetics of DNA methylation acquisition leading to specific 433 embryo-placenta DMR categories is not a stepwise process occurring throughout cell fate 434 decisions and patterning of embryonic and extraembryonic lineages, but a prompt progression in 435 the early post-implanted conceptus. These results are in line with studies reflecting that the 436 initiation of asymmetric DNA methylation levels begins within the trophoblast and the inner cell 437 mass of the blastocyst (Santos et al., 2002), and becomes highly evident by the time the epiblast 438 acquires its initial global DNA methylation patterns at E6.5 during the *de novo* reprogramming wave (Auclair et al. 2014, Smith et al. 2017). Our results show that the acquisition period 439 440 between E4.5 and E6.5 is particularly key to establishing asymmetric DNA methylation patterns 441 associated with mid-gestation embryo-placenta DMR classes, and that these DMR associated-442 patterns are long-lasting across stages of prenatal development. Since we studied cell populations 443 derived from embryonic and placental tissues, we cannot dismiss the prevalence of cell-to-cell 444 heterogeneity in the acquisition kinetics of specific DMR patterns. Despite this concern, our 445 results indicate that embryonic cells have a large body of DMRs with High-levels ( $\geq 80\%$ ) and

446 Low-levels (<20%) of methylation that remain static across development, as well as in somatic 447 cell types (High-Mid & High-Low DMRs), whereas compared to the embryo, the placental 448 DMRs have overall lower methylation levels and are more broadly distributed amongst 449 methylation levels, which has also been shown in other studies where genome-wide DNA 450 methylation profiles of placental cells were compared to other tissues and specific cell types 451 (Schroeder et al., 2013; Chatterjee et al., 2016; Smith et al., 2017). Lower methylation levels in 452 the placenta have been associated with reduced *de novo* methyltransferase activity in the ExE 453 during the *de novo* reprogramming wave (Fulka et al., 2004, Santos et al., 2002). Nevertheless, 454 we observe methylation level peaks for all DMR categories within the ExE at E6.5 before levels 455 stabilize at E10.5. As of now, the implication of these methylation level peaks on future 456 regulation mechanisms and methylation profiles is unknown. Moreover, it remains to be 457 determined whether global reduction in DNA methylation marks on DMRs between the E6.5 and 458 E10.5 extraembryonic cell lineages is stochastic or targeted, and whether it occurs through 459 passive or active mechanisms.

460 We also observed an enrichment of retrotransposons (i.e., LINE, SINE, LTR) in DMRs 461 especially in those with High-levels of methylation in the embryo and lower-levels in the 462 placenta, whereas DMRs with Low-levels of methylation in the embryo and higher levels in the 463 placenta are preferentially outside retrotransposons-associated sequences. Earlier findings of 464 Chapman et al. (Chapman et al., 1984) revealed that repeat regions in the placenta appear to lack 465 tight control of their methylation patterns, perhaps indicating that maintaining methylation, and 466 therefore repression of these elements for genome stability and integrity, is not critical given the 467 relatively short lifespan of this organ. However, we do observe Mid-range methylation levels 468 (20-80%) in the placenta for a substantial portion of DMRs that are associated with 469 retrotransposons, revealing that specific genomic regions associated with repetitive elements do 470 need tight regulation in extraembryonic cell lineages for proper development. These results are in 471 line with findings exposing that the deletion of genome-defense gene Tex19.1 leads to the de-472 repression of LINE1 and compromises placental development, suggesting that disparities 473 between retrotransposon suppression and genome-defense mechanisms might contribute to 474 placenta dysfunction and disease (Reichmann et al., 2013).

475 DNMT3A and DNMT3B are required to establish proper methylation profiles on the 476 embryonic genome during the *de novo* reprogramming wave, as both methyltransferase enzymes 477 have redundant, but also specific functions. However, the activity of DNMT3B is the main 478 contributor in the acquisition of profiles in epiblast cells, and especially commands methylation 479 on CGIs associated with developmental genes (Auclair et al., 2014). Auclair et al. highlighted 480 that in the absence of DNMT3B, DNMT3A was not able to counterbalance, leading to the loss of 481 promoter-CGI methylation and gain of expression of germline genes (e.g., Sycp1, Sycp2, Mael, 482 *Rp1101, Dmrtb1*) in somatic cells of the embryo. Here, we show that the promoter of these 483 germline genes, associated with meiotic and piRNA processes as well as other genes with similar 484 biological functions, are highly enriched in High-Mid DMRs. We also reveal that for this 485 particular set of High-Mid DMRs, mainly deprived of CGIs, DNMT3A is able to substantially 486 counterbalance the lack of DNMT3B in the E6.5 epiblast and establish global DNA methylation 487 levels, whereas in the ExE, this compensatory mechanism is rather inefficient. Globally, we 488 show that DNMT3B is much more potent at compensating than DNMT3A in both the epiblast 489 and ExE for all DMR categories, indicating that DNMT3B is the main *de novo* enzyme driving 490 asymmetric DNA methylation patterns between the embryo and placenta. Although 491 compensatory mechanisms have been observed in *Dnmt3a* or *Dnmt3b*-depleted conceptuses, we 492 still do not fully understand the process as Dnmt3a and Dnmt3b have cell lineage specific 493 expression during the peri-implantation developmental period. As development progresses, we 494 observed that the compensation mechanism in *Dnmt3a* or *Dnmt3b*-deficient embryos remains 495 apparent at E8.5 on most DMR categories, but is less effective as the methylation gaps increase 496 compared to wild-type. This correlates with a developmental period where DNMT3A is now the 497 main *de novo* methyltransferase enzyme in both the embryo and placenta (Okano et al., 1999, 498 Watanabe et al., 2002), and with a compensatory activity of DNMT3A being less effective. This 499 suggests that DNMT3B activity is critical to ensure proper establishment of DNA methylation 500 asymmetry between the embryonic and extraembryonic cell lineages during the de novo reprogramming wave, but that DNMT3A is required during the developmental progression to 501 502 safeguard methylation levels on DMR categories.

503

### 504 **CONCLUSION**

505 We demonstrate that asymmetry between embryo and placenta DNA methylation patterns 506 occurs rapidly during *de novo* acquisition of methylation marks in the early post-implanted 507 conceptus, and that these patterns are long-lasting across subtypes of DMRs. We also reveal that

508 at the peri-implantation stages, *de novo* methyltransferase activity of DNMT3B is the main 509 provider of asymmetric methylation marks on DMRs, and that it largely palliates the lack of 510 DNMT3A in the epiblast and ExE. However, as development progresses, DNMT3A becomes the 511 principal *de novo* methyltransferase by mid-gestation, and DNMT3B methyltransferase activity 512 is less effective at promoting compensation. These results further underline why embryos 513 developing without DNMT3B have severe DNA methylation defects and die at mid-gestation, 514 whereas those without DNMT3A only die postnatally. Further investigation is required to 515 determine the molecular mechanisms controlling the precise *de novo* acquisition of long-lasting 516 methylation marks on specific DMR subtypes in the embryonic and extraembryonic cell 517 lineages, and how errors in this process could lead to abnormal development and diseases.

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523

### 524 **COMPETING INTERESTS**

525 No competing interests declared

526

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532

# 533 DATA AVAILABILITY

534 The data from this study have been submitted to the Gene Expression Omnibus (#GSE95610).

535 Reviewer link:

536	https://www	.ncbi.nlm.n	ih.gov/	geo/q	uery/acc.c	gi?token=	ghsnmc	octlgilc	j&acc=GSE956	10

537

- 538 Table S1. Annotation and DNA methylation values associated with DMRs.
- 539 Table S2. DNA methylation levels for DMR categorises across development.
- Table S3. DNA methylation levels for DMR categories in various *Dnmt* knockout mouse
   models.
- 542 Table S4. Number of 100bp tiles in each category of public data that overlap with Embryo-
- 543 Placenta DMRs (related to Figure 4; 5A; S4; S5; S6)
- 544

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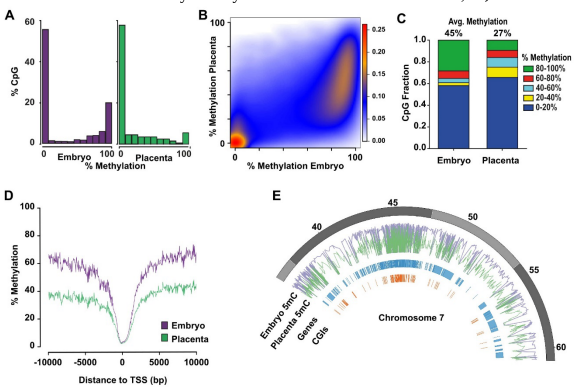
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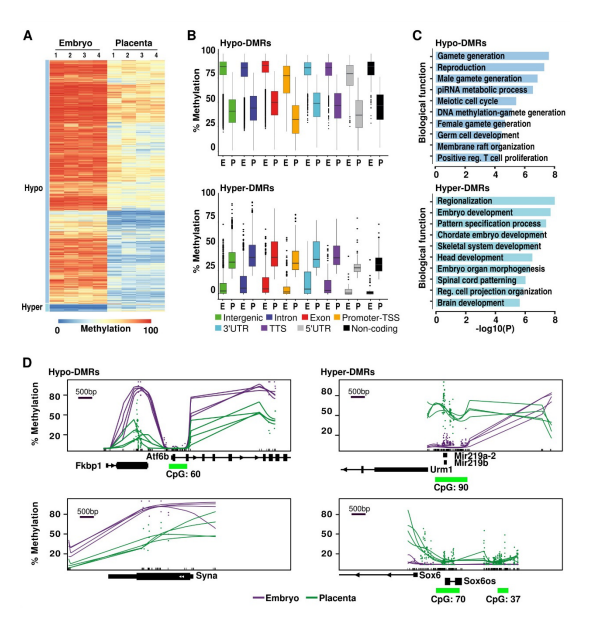




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**Figure 1. Distinctive patterns of genome-wide DNA methylation accumulation between** mid-gestation embryo and placenta. Analyses of genome-wide DNA methylation sequencing results for embryo (n=4) and placenta (n=4). A) Density histograms showing the distribution of CpG methylation levels for embryo (*purple*) and placenta (*green*). B) Pairwise comparison of CpG methylation between embryo and placenta. Density increases from blue to red. C) CpG fraction of 100 bp tiles within 0-20%, 20-40%, 40-60%, 60-80% and 80-100% ranges in embryo

and placenta. Shown above bars, is the average CpG methylation for each unique tile represented
in graph. D) DNA methylation means surrounding the transcription start site (TSS) for All-tiles
in each experimental group. E) Circle plot showing methylation average of embryo (*purple*) and
placenta (*green*) across a portion of chromosome 7.

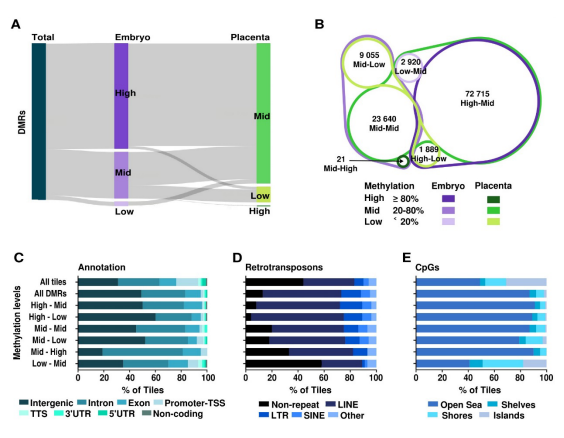






754 and placenta (n=4) samples. A) Heatmap representation of DNA methylation levels for the 755 20 000 tiles with the most variable levels between embryo and placenta (DNA methylation 756 variance >20% and P<0.05). B) Box-plots representing DNA methylation distribution in embryo 757 and placenta for the different genomic annotation regions (intergenic, intron, exon, promoter-758 TSS, 3'UTR, TTS, 5'UTR and non-coding). C) Summary of biological functions associated with 759 promoter regions in Hypo- and Hyper-DMRs. D) Examples of smoothed methylation profiles 760 (BSmooth tool) in regions with lower methylation profiles (Atf6b, and Syna) or higher 761 methylation profiles (Mir219a-2/Mir219b, and Sox6) in placenta.







765 Figure 3. Distinct genomic features in DMRs based on their methylation status in embryo and placenta. DMR analysis by methylation levels in embryo and placenta. A) Sankey diagram 766 767 dividing DMRs by methylation levels in embryo (*purple*) and placenta (green) for the associated 768 tiles. High :  $\geq$ 80% methylation, Mid (intermediate) :  $\geq$ 20% - <80% methylation , Low : <20% 769 methylation. B) Venn diagram showing the proportion of tiles in the different DMR categories 770 based on DNA methylation levels between embryo and placenta. C), D) and E) Analysis of all 771 tiles, all the DMRs, as well as the 6 different DMR categories based on levels of DNA 772 methylation in embryo and placenta for : C) Genomic annotations, D) Main retrotransposons and 773 E) Proximity of CpG rich regions. Neighboring CpG dense regions were defined as shore; up to 774 2kb away from CGIs, shelf; 2-4kb away from CGIs, and open sea; >4kb away from CGIs.

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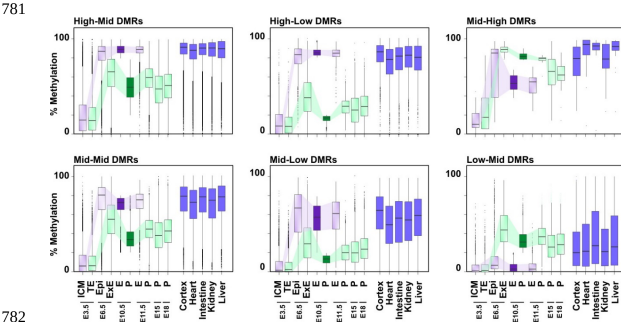
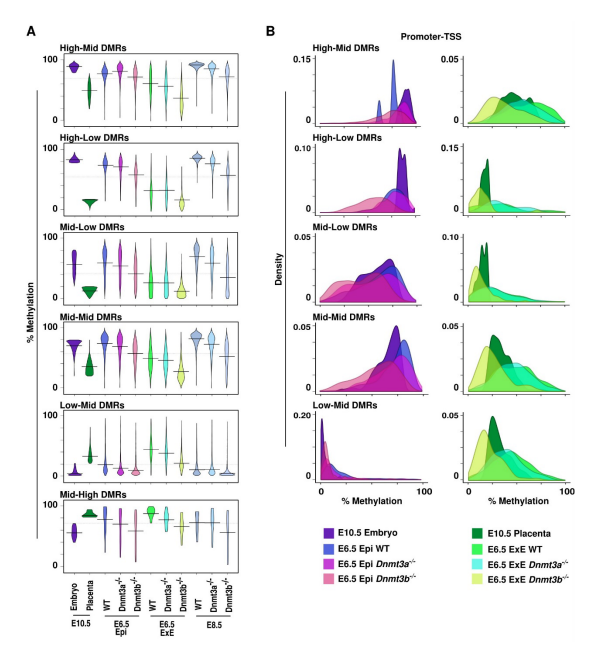




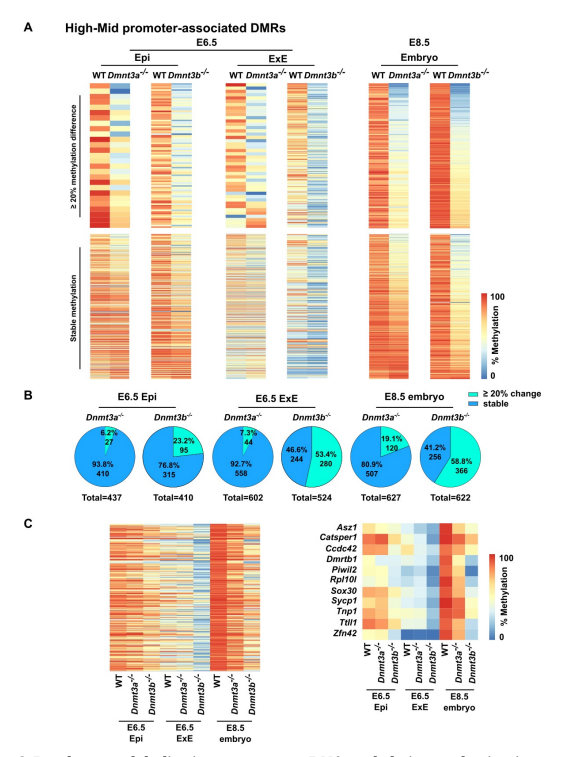
Figure 4. Dynamics of DNA methylation profiles associated with DMR categories and their evolution across embryo and placenta development. Box plots representing the DNA methylation distribution and median values for each DMR category in various developmental stages. Tiles associated with DMR categories at E10.5 were overlapped with previously published and publicly available data, and methylation levels were determined at each developmental stage (Smith et al 2012, Smith et al. 2014, Whidden et al. 2016) or in adult somatic tissues (Hon et al. 2013). ICM: inner cell mass, TE: trophectoderm, Epi: epiblast, ExE: extraembryonic ectoderm, E: embryo, Pla: placenta. See Table S2 for median and mean methylation values, and Table S4 for number of overlapping tiles analysed.





806 Figure 5. Dnmt3a- or Dnmt3b-deficiency alter proper establishment of DMR-associated 807 patterns. A) Violin plot representing DNA methylation distribution and median values of tiles 808 associated with the different DMR categories at E10.5, and their methylation levels in the 809 different tissues and genotypes at E6.5 and E8.5. B) Promoter-TSS density graphs for E10.5 embryo and E6.5 epiblast (wt,  $Dnmt3a^{-/-}$  and  $Dnmt3b^{-/-}$ ) (left panel) and for E10.5 placenta and 810 811 E6.5 extraembryonic ectoderm (wt,  $Dnmt3a^{-/-}$  and  $Dnmt3b^{-/-}$ ) (right panel). E: embryo, Epi: 812 epiblast, ExE: extraembryonic ectoderm. See Table S3 for median and mean methylation values, 813 and Table S4 for number of overlapping tiles analysed.

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817 Figure 6. Developmental decline in compensatory DNA methylation mechanism in response

to *Dnmt3a*- or *Dnmt3b*-deficiency at promoter-TSS associated with High-Mid DMRs. A) Heatmaps comparing the DNA methylation profiles of wt vs *Dnmt3a*<sup>-/-</sup> or wt vs *Dnmt3b*<sup>-/-</sup> for E6.5 epiblast, E6.5 extraembryonic ectoderm or E8.5 embryo in 100bp tiles overlapping promoter-associated DMRs that are highly methylated in the embryo ( $\geq$ 80%) and mildly methylated in the placenta ( $\geq$ 20 and <80%). Upper panel represents regions with a difference of methylation of at least 20% between wt and *Dnmt3a*<sup>-/-</sup> or wt and *Dnmt3b*<sup>-/-</sup>. Lower panel bioRxiv preprint doi: https://doi.org/10.1101/718247; this version posted August 1, 2019. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

represents stable regions (lower than 20% difference in methylation). **B**) Pie charts representing DMR numbers for each comparison. **C**) Heatmap of the methylation levels of tiles commonly represented in all 9 public data samples that overlap with promoters-associated High-Mid DMRs (n = 314 tiles) (left panel). Heatmap of the methylation level of gene associated with piRNA, gamete generation and germ cells or meiotic nuclear division that are covered in all 9 public datasets (right panel).

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