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Maternal DNA Methylation Regulates Early Trophoblast Development

Graphical Abstract

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In Brief

Branco et al. dissect the role of DNA methylation in mouse trophoblast development through genome-wide profiling of methylation-deficient mutants. DNA methylation marks carried over from the oocyte play a major role in trophoblast development and cell adhesion, which is partly dependent on silencing of the Polycomb gene Scml2.

Highlights

- Oocyte-derived DNA methylation is an important regulator of GSE66049 trophoblast transcription
- DNA methylation controls trophoblast cell adhesion
- Silencing of Polycomb gene Scml2 is necessary for normal trophoblast development
- Accession Numbers

Maternal DNA Methylation Regulates Early Trophoblast Development

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SUMMARY

Critical roles for DNA methylation in embryonic development are well established, but less is known about its roles during trophoblast development, the extraembryonic lineage that gives rise to the placenta. We dissected the role of DNA methylation in trophoblast development by performing mRNA and DNA methylation profiling of Dnmt3a/3b mutants. We find that oocyte-derived methylation plays a major role in regulating trophoblast development but that imprinting of the key placental regulator Ascl2 is only partially responsible for these effects. We have identified several methylation-regulated genes associated with trophoblast differentiation that are involved in cell adhesion and migration, potentially affecting trophoblast invasion. Specifically, trophoblast-specific DNA methylation is linked to the silencing of Scml2, a Polycomb Repressive Complex 1 protein that drives loss of cell adhesion in methylation-deficient trophoblast. Our results reveal that maternal DNA methylation controls multiple differentiation-related and physiological processes in trophoblast via both imprinting-dependent and -independent mechanisms.

Fertilization marks the start of a cascade of rapid epigenetic changes which, coupled to an intricate network of signaling and transcriptional events, ultimately lead from a totipotent zygote to a myriad of differentiated tissues that comprise the embryo as well as supporting extraembryonic tissues. DNA methylation plays essential roles during this time, mainly by mediating silencing of specific genes and transposable elements. Importantly, while genome-wide DNA methylation erasure occurs after fertilization, key genomic regions are kept methylated, including imprinting control regions (ICRs) and murine intracisternal A-par-

ticle (IAP) retrotransposons ([Lane et al., 2003; Smith et al., 2012](#page-12-0)). This epigenetic reprogramming phase is followed by de novo DNA methylation post-implantation, which helps to establish and cement tissue-specific expression programs, thereby driving cell differentiation and organogenesis.

In mammals, three DNA methyltransferases (DNMTs) are responsible for establishing and maintaining DNA methylation profiles: DNMT1 is mainly involved in the maintenance of methylation patterns during replication, whereas DNMT3A and DNMT3B have de novo methylation activity. Mouse knockout (KO) models have shown that all three enzymes are essential for correct embryonic development: *Dnmt1* and *Dnmt3b* KOs are embryonic lethal ([Li et al., 1992; Okano et al., 1999\)](#page-12-1), whereas *Dnmt3a* KO mice die postnatally [\(Okano et al., 1999\)](#page-12-2). The combined double KO (DKO) of *Dnmt3a* and *Dnmt3b* has a more severe phenotype than either single KO, with embryos dying at around embryonic day 10.5 (E10.5) [\(Okano et al., 1999](#page-12-2)). Importantly, conditional deletion of *Dnmt3a* in the oocyte is sufficient to halt embryonic development at E10.5 [\(Kaneda et al., 2004](#page-11-0)), showing that maternal methylation is critical for developmental progression. Maternal KO of *Dnmt3l*, a catalytically inactive cofactor that interacts with DNMT3A for the establishment of DNA methylation in the oocyte, displays a very similar phenotype [\(Bourc'his et al., 2001](#page-11-1)).

While the role of DNA methylation in the development of the embryo is well established ([Auclair et al., 2014](#page-11-2)), its importance in the development of the extraembryonic trophoblast lineage remains less clear. Notably, trophoblast tissues are largely hypomethylated when compared with embryonic tissues, in particular at repeat elements, and embryos derived by nuclear transfer from ESCs lacking all three active DNMTs can contribute to extraembryonic tissues when aggregated with wild-type (WT) embryos [\(Sakaue et al., 2010\)](#page-12-3). On the other hand, conceptuses from *Dnmt3l*-null mothers display morphogenic defects across all layers of the placenta ([Arima et al., 2006\)](#page-11-3). To date, comprehensive molecular characterization of the DNA methylation and gene expression alterations linked to these defects is lacking. While they have been largely attributed to the loss of imprinted gene expression, some methylation marks outside of imprinting are also carried over from the oocyte to the blastocyst stage [\(Smallwood et al., 2011\)](#page-12-4), but have not been functionally

Figure 1. Oocyte Methylation Is a Major Regulator of Trophoblast Gene Expression

(A) Females carrying floxed (f) alleles for *Dnmt3a* and *Dnmt3b* as well as a *Zp3*-driven Cre transgene were crossed to heterozygous males, yielding four different genotypes collectively referred to as mDKO, due to the absence of methylation in the oocyte; conceptuses from females without the *Zp3*-Cre transgene were used as controls.

(B) Maternal deletion of *Dnmt3a/3b* results in trophoblast defects at E9.5 (top) characterized by loss of adhesion of TGCs (arrowheads), with no apparent difference in phenotype between different post-zygotic genotypes. In contrast, DKO embryos are more severely affected than DHet embryos (bottom). Images are not on the same scale.

(C) H&E staining of paraffin-embedded sections shows that *Dnmt3a* mKO trophoblast lacks the labyrinthine layer that is otherwise seen developing in WT trophoblast (marked by an asterisk); the TGC layer is less dense in *Dnmt3a* KO trophoblast, possibly due to cell adhesion defects. ch, chorion; epc, ectoplacental cone.

(D) Hierarchical clustering of mRNA-seq data from E7.5 EPCs reveals segregation of mDKO and Ctrl genotypes but no further differentiation of individual mDKO genotypes.

(E) mRNA-seq expression values for examples of deregulated genes common to all mDKO genotypes (top), and genes controlled by post-zygotic DNA methylation (bottom). Error bars represent SD. See also Figure S1.

explored. Indeed, of 1,329 CpG islands (CGIs) that are hypermethylated in oocytes relative to sperm, only 23 are associated with known ICRs ([Kobayashi et al., 2012](#page-12-5)). In this study, we have performed mRNA sequencing (mRNA-seq) and whole-genome bisulfite sequencing (BS-seq) on trophoblast from *Dnmt3a/3b* KO mice. We show that the prevailing phenotype is explained by the absence of maternal methylation marks. However, failure to establish correct imprinted gene expression does not explain all observed transcriptional changes. Our data suggest that maternal DNA methylation plays critical roles in the control of cell adhesion in trophoblast giant cells (TGCs) and in the formation of syncytiotrophoblast (SynT).

RESULTS

Absence of Oocyte DNA Methylation Leads to Cell

To study the role of DNA methylation in trophoblast development, we used female mice carrying conditional alleles for both *Dnmt3a* and *Dnmt3b* ([Dodge et al., 2005; Kaneda et al., 2004\)](#page-11-4), as well as a *Zp3-Cre* transgene; these were crossed to double heterozygous males, i.e., *Dnmt3a^{+/-};Dnmt3b^{+/-}* ([Figure 1](#page-2-0)A). Deletion of *Dnmt3a* and *Dnmt3b* driven by *Zp3* expression yields oocytes that lack both enzymes (and virtually all DNA methylation) ([Kaneda et al., 2010; Shirane et al., 2013](#page-11-5)). We will therefore refer to the group of genotypes resulting from this cross collectively as maternal DKOs (mDKOs), and the individual genotypes derived from this cross as such: DHet for *Dnmt3a/+*;*Dnmt3b/+*, 3aKO for *Dnmt3a^{-/-};Dnmt3b^{-/+},* 3bKO for *Dnmt3a^{-/+}; Dnmt3b/*, and DKO for *Dnmt3a/*;*Dnmt3b/*. To generate a control cohort, we made a separate cross using females without the *Zp3-Cre* transgene ([Figure 1](#page-2-0)A). For simplicity, we will refer to these genotypes as control (Ctrl) genotypes and will not distinguish between the various combinations of WT homozygous and heterozygous alleles generated by this cross.

We first dissected conceptuses at E9.5 for morphological characterization. As previously described [\(Okano et al., 1999\)](#page-12-2), DKO embryos were severely developmentally delayed, with few defined somites and open neural tube, among other defects, whereas other genotypes exhibited less pronounced abnormalities ([Figure 1B](#page-2-0)). However, to our surprise trophoblast tissues showed a very consistent phenotype across all genotypes of the mDKO cohort, with no obvious additional defects being observed in DKO trophoblast over DHet trophoblast ([Figure 1B](#page-2-0)). The most prominent characteristic of these tissues was a reduction in the adhesion of TGCs that make up the outermost lining of the implantation site, as these cells were easily dissociated from the remaining tissue when compared with control trophoblast. Maternal deletion of *Dnmt3a* was sufficient to produce the same phenotype. Histological analysis of *Dnmt3a* maternal knockout (mKO) trophoblast at E9.5 revealed defects similar to those described for *Dnmt3l* mKO trophoblast [\(Arima et al.,](#page-11-3) [2006; Bourc'his et al., 2001\)](#page-11-3). Namely, *Dnmt3a* mKOs had a defect in chorio-allantoic fusion that in turn led to a failure in development of the labyrinthine layer, which can be made out in WT trophoblast by finger-like invaginations of the allantoic mesoderm into the chorionic ectoderm layer forming welldefined fetal blood spaces ([Figure 1C](#page-2-0)). In addition, while the TGC layer appeared enlarged, this was mostly a result of reduced tissue density, as there was a notable increase in extracellular space in this layer ([Figures 1C](#page-2-0) and S1A). It is possible that such spacing only becomes apparent as a result of the histological preparation, but given that WT and mKO tissues were processed in parallel and embedded in the same paraffin block, this is likely to be an expression of the loss of cell adhesion seen in dissected tissues. Cellular adhesion plays key roles in placental development, regulating trophoblast inva-siveness into the maternal decidua ([Harris et al., 2009; Suther](#page-11-6)[land et al., 1993\)](#page-11-6). Importantly, dysregulation of adhesion molecules is commonly observed in placental disorders, including pre-eclampsia and intrauterine growth restriction (Harris et al., 2009; Pollheimer and Knöfler, 2005). It is therefore plausible that the developmental arrest of methylation-deficient conceptuses is due, at least in part, to alterations in the cellular adhesion profile of trophoblast cells.

mRNA Profiling Reveals Genes Controlled by Oocyte

To gain deeper insights into these hypomethylation-induced changes, we profiled the transcriptome of mDKO and Ctrl trophoblasts. To compare structurally similar tissues and identify primary, causative aberrations in the gene expression patterns, we used E7.5 ectoplacental cones (EPCs), at which point mDKO conceptuses were visually indistinguishable from Ctrl conceptuses. We performed mRNA-seq of three EPCs from each genotype group. Interestingly, hierarchical clustering of the data mirrored our phenotypic observations: Ctrl EPCs clustered together, away from a large group of mDKO EPCs, with no individual genotype therein being discernible from the others ([Figure 1](#page-2-0)D). Accordingly, we identified 368 differentially expressed (DE) genes between Ctrl and DHet EPCs, whereas comparison of DHet with 3aKO or 3bKO genotypes yielded only 4 and 6 DE genes, respectively. This strongly suggests that nearly all transcriptional effects are due to oocyte methylation deficiency. Interestingly, we did identify 45 DE genes between DKO and DHet EPCs. As previously described for embryonic tissues [\(Auclair et al., 2014; Borgel et al., 2010; Hackett](#page-11-2) [et al., 2012](#page-11-2)), these include germline-specific genes such as *Dazl*, *Rhox2a*, and *Tuba3b* [\(Figure 1E](#page-2-0)). These genes are repressed by de novo methylation, as post-zygotic deletion of *Dnmt3a/3b* is sufficient to cause their upregulation (Figure S1B).

To focus on the group of oocyte methylation-dependent genes, we selected 137 genes that passed stringent criteria for differential expression in all mDKO genotypes when compared with Ctrl EPCs (Table S1). These "mDKO DE genes" were made up of 39 upregulated and 98 downregulated genes, and showed very similar expression levels in all mDKO genotypes (Figure S1C), suggesting that they are solely dependent on oocyte methylation and not on the presence of DNMTs postfertilization. Concordantly, post-zygotic deletion of *Dnmt3a/3b* has no effect on the expression of mDKO DE genes (Figure S1B). Among the mDKO DE genes were transcription factors and markers relevant to trophoblast development, such as *Cdx2*, *Tpbpa*, and *Pcdh12* ([Figure 1E](#page-2-0), RT-qPCR validation in Figure S1B). Given that *Cdx2* is a key trophoblast stem cell (TSC) transcription factor, we asked whether the TSC niche was affected in *Dnmt3a* mKO mutants by performing immunofluorescence for CDX2 on sections of E7.5 conceptuses. We found that CDX2 depletion only occurred in the EPC and that the extraem-

bryonic (chorionic) ectoderm, which harbors the TSC niche, remained unaffected (Figure S1D), suggesting that loss of CDX2 is unrelated to deregulation of the stem cell compartment, but critically affects the diploid core of the EPC. Key markers of placental development were also affected, including the spongiotrophoblast marker *Tpbpa* and the glycogen cell precursor marker *Pcdh12* ([Figures 1](#page-2-0)E and S1B). Notably, TGC markers such as *Prl3d1*/*Pl1* were unaffected (Figure S1B), suggesting that the observed phenotypic alterations are not due to major defects in the differentiation of this placental cell type.

Loss of Asc/2 Imprinting Only Partially Explains
Alterations in mDKO Trophoblast

Oocyte methylation controls several ICRs that are essential for maternal regulation of imprinted genes, which are important for both embryo and trophoblast development. As previously sug-gested [\(Arima et al., 2006](#page-11-3)), maternal effects on trophoblast development may therefore be a result of loss of specific imprinted genes. To test this hypothesis, we first identified known methylation-dependent imprinted genes within our mDKO DE gene list. Of 79 imprinted genes (paternal and maternal) in our mRNA-seq data, 59 were robustly expressed in EPCs from at least one of the genotypes. However, only five were consistently altered in all mDKO genotypes: *Zrsr1*, *Cd81*, *Ascl2*, *Phlda2*, and *Cdkn1c* ([Figure 2A](#page-4-0)). *Zrsr1* (also known as *U2af1-rs1*) was upregulated in mDKO EPCs but is unlikely to be involved in the phenotype of mDKO conceptuses, as mice with paternal disomy of chromosome 11 (where *Zrsr1* lies) are viable [\(Cattanach and](#page-11-7) [Kirk, 1985\)](#page-11-7). *Cd81*, *Cdkn1c*, *Phlda2*, and *Ascl2*, which are all part of the same imprinting cluster, were all robustly downregulated in mDKO EPCs, as lack of oocyte methylation leads to activation of the non-coding transcript *Kcnq1ot1* on the maternal allele, which is known to drive silencing of genes in its vicinity [\(Peters and Robson, 2008\)](#page-12-6). *Cd81* KO mice are viable ([Maecker](#page-12-7) [and Levy, 1997\)](#page-12-7), and both *Cdkn1c* and *Phlda2* KO placentas are enlarged and show an expansion of the spongiotrophoblast layer, which is a very different phenotype from that observed in mDKOs ([Frank et al., 2002; Takahashi et al., 2000\)](#page-11-8). However, trophoblast from maternal KO of *Ascl2* (also known as *Mash2*) has a severe phenotype mainly characterized by a lack of spongiotrophoblast formation, which leads to embryonic lethality at around E10 ([Guillemot et al., 1994](#page-11-9)). Given the similarity in phenotype timing to the mDKO trophoblast, as well as the downregulation of spongiotrophoblast marker *Tpbpa* observed in both models, we decided to test whether *Ascl2* downregulation was driving the transcriptional changes seen in mDKO EPCs. For this purpose, we used an *Ascl2-lacZ* knockin mouse line [\(Tanaka](#page-12-8) [et al., 1999](#page-12-8)) to generate mKO conceptuses of *Ascl2*. We first performed histological analysis at E9.5, which revealed that *Ascl2* mKOs had a reduced or absent labyrinthine layer despite having completed chorio-allantoic fusion, and had an enlarged TGC layer (Figure S2A), as previously described [\(Guillemot et al.,](#page-11-9) [1994\)](#page-11-9). However, unlike *Dnmt3a* mKO trophoblast, the TGC layer expansion did not involve a significant increase in extracellular space (Figure S2B), inferring that TGC cell adhesion is largely intact in *Ascl2* mKO mutants. We then isolated E7.5 EPCs from *Ascl2* WT and mKO conceptuses for RT-qPCR analyses. We confirmed that *Ascl2* repression leads to *Tpbpa* downregulation, but also found drastic downregulation of *Cdx2* and *Pcdh12*,

Figure 2. Ascl2 Depletion Does Not Drive the Bulk of Gene Expression Alterations in mDKO EPCs

(A) mRNA-seq data reveal deregulated maternally controlled imprinted genes in DHet EPCs (red), whereas a paternally controlled imprinted gene is unchanged (*Igf2*, blue).

(B) RT-qPCR data from *Ascl2* WT and mKO E7.5 EPCs shows that some key genes deregulated in mDKO EPCs are driven by *Ascl2* downregulation (*p < 0.05, **p < 0.01, ***p < 0.001; t test). Error bars represent SD.

(C) Differential expression analysis from mRNA-seq of *Ascl2* mKO EPCs reveals that only a minority of *Dnmt3* mDKO DE genes are explained by *Ascl2* repression.

(D) Log2 fold change in expression between WT and *Ascl2* or *Dnmt3* mutant EPCs for each grouping of genes defined in (C). *Ascl2*-dependent genes display similar expression changes in *Dnmt3* mDKO EPCs, whereas *Dnmt3* unique DE genes are unchanged in *Ascl2* mKO EPCs. See also Figure S2.

similar to that seen in *Dnmt3* mDKO EPCs [\(Figure 2](#page-4-0)B). We then extended this analysis by performing mRNA-seq on control and *Ascl2* mKO EPCs. We found that, while 43 genes were commonly deregulated between *Ascl2* mKO and *Dnmt3* mDKO EPCs, there were 94 DE genes that were unique to the *Dnmt3* mDKOs [\(Figure 2](#page-4-0)C). Surprisingly, we also found 216 genes seemingly only deregulated in *Ascl2* mKOs. However, when we analyzed the expression of these genes in *Dnmt3* mDKO EPCs, we found that they displayed expression changes very similar to those seen in *Ascl2* mKOs [\(Figure 2D](#page-4-0)), but had not passed our stringent criteria for differential expression calling. Importantly, genes deregulated only in the *Dnmt3* mDKO EPCs did not display substantial changes in expression in *Ascl2* mKOs [\(Figure 2](#page-4-0)D), demonstrating that these are indeed *Ascl2*-independent effects. Our data suggest that the majority of transcriptional alterations in mDKO trophoblast are independent of imprinting of a key regulator of placental development. While we cannot completely rule out that the combined loss of imprinting at other loci may drive the gene expression changes seen in mDKOs, it is likely that maternally derived methylation marks outside of ICRs play a major role in trophoblast gene regulation.

mDKO-Affected Genes Are Involved in Trophoblast

Development and Addressed Correlation
We decided to focus on the group of 94 genes that were affected in *Dnmt3* mDKO but not *Ascl2* mKO EPCs. Gene ontology (GO) analysis revealed that this *Dnmt3*-specific gene list was enriched for genes involved in signal transduction (e.g., *Ephb2*, *Stk10*, *Pik3ap1*, *Ptpn3*) and the regulation of guanosine triphosphatases (GTPases) (e.g., *Asap1*, *Rgs3*, *Arhgef4*, *Rasa4*) (Table S2), whereas no such enrichment was seen in *Ascl2*-specific genes (Table S3). GTPases control many key cellular processes, including focal adhesion and migration/invasion ([Menke and](#page-12-9) [Giehl, 2012](#page-12-9)), which is consistent with the cellular adhesion defect observed in mDKO trophoblast. Notably, numerous other genes involved in cell adhesion were found to be deregulated in mDKO, such as *Itga7*, *Flnc*, *Dbnl*, and *Plxna1*. Although the GO term ''cell adhesion'' did not reach significance in our analysis, we noted that many deregulated genes with known roles in cell adhesion and migration lack this annotation (e.g., *Asap1*, *Rasa4*, *Srgap3*, *Spry1*). Acquisition of an invasive phenotype is a key component of the differentiation process of TGCs ([Hemberger et al., 2003;](#page-11-10) [Hunkapiller et al., 2011\)](#page-11-10). In line with this, *Ephb2*, a receptor tyrosine kinase that activates Rho family GTPases and is involved in the formation of secondary TGCs ([El-Hashash and Kimber,](#page-11-11) [2006\)](#page-11-11), was downregulated in mDKO EPCs. Other deregulated genes involved in trophoblast differentiation included *Gata3* ([Ral](#page-12-10)[ston et al., 2010\)](#page-12-10), *Gjb5* ([Kibschull et al., 2014](#page-12-11)), *Dlx3* [\(Morasso](#page-12-12) [et al., 1999](#page-12-12)), and *Alkbh1/2700073G19Rik* [\(Pan et al., 2008\)](#page-12-13). This prompted us to ask whether mDKO-affected genes were generally associated with trophoblast differentiation. Genes annotated with GO terms associated with trophoblast or placental development were not significantly enriched (although we again found this annotation to be incomplete). However, five out of six tested genes showing upregulation in mDKO EPCs were found to increase in expression during differentiation in vivo [\(Figure 3](#page-5-0)A), and similar results were obtained during in vitro differentiation of TSCs (Figure S3). To expand on these observations using an annotation-independent approach, we examined the behavior of mDKO-affected genes in expression data from E9.5 WT TGCs [\(Sher et al., 2013](#page-12-14)) and found that most of these genes are indeed differentially expressed relative to WT E7.5 EPCs [\(Fig](#page-5-0)[ure 3](#page-5-0)B). Moreover, their expression in mDKO E7.5 EPCs was well correlated with the expression profile of E9.5 TGCs [\(Figure 3B](#page-5-0)), indicative of precocious TGC differentiation in mDKO trophoblast. Although some TGC markers (such as *Prl3d1*) are unchanged in mDKO EPCs (Figure S1B), it is likely that deregulation of these TGC-associated genes affects the function of this cell population.

Trophoblast invasion and placental development critically depend on epithelial-to-mesenchymal transition (EMT) ([Kokki](#page-12-15)[nos et al., 2010; Parast et al., 2001; Sutherland, 2003\)](#page-12-15). Notably, apart from signal transduction and GTPase regulators, other mDKO-deregulated genes with known roles in EMT included

Figure 3. mDKO DE Genes Are Associated with Trophoblast Differentiation

(A) Several *Dnmt3*-specific genes affected in mDKO EPCs become upregulated during trophoblast development, as revealed by RT-qPCR of E7.5 and E9.5 WT trophoblast.

(B) Expression of mDKO DE genes in E9.5 TGCs relative to E7.5 EPCs. mRNA-seq data from E9.5 TGCs ([Sher et al., 2013\)](#page-12-14) were plotted against our mRNA-seq data for Ctrl (left) or mDKO (right) E7.5 EPCs. Genes upregulated in mDKO EPCs (green) have increased expression in TGCs relative to WT E7.5 EPCs. Similarly, mDKO downregulated genes (red) tend to have lower expression in TGCs. When plotted against mDKO EPC data, the expression of mDKO DE genes is much more comparable with that seen in TGCs.

(C) RT-qPCR of EMT-associated genes in E9.5 trophoblast reveals that mDKO trophoblasts do not exhibit a classic EMT phenotype, although *Cdh1* is downregulated.

Error bars represent SD. *p < 0.05, **p < 0.01, ***p < 0.001; t test comparing E9.5 with E7.5 (A), or ANOVA with post hoc tests comparing control (Ctrl) and mDKO genotypes (C). See also Figure S3.

Mmp15 ([Tao et al., 2011\)](#page-12-16) and *Grhl2* [\(Cieply et al., 2012\)](#page-11-12). However, genes widely involved in EMT (e.g., *Cdh1*, *Snai1*, *Zeb1*) were not significantly altered in our E7.5 mRNA-seq data. To test whether an overt EMT phenotype expressed itself at a later developmental stage, we measured the expression of key EMTassociated genes in E9.5 Ctrl and mDKO trophoblast [\(Figure 3](#page-5-0)C). While we did observe a prominent decrease in the classic epithelial marker *Cdh1* (E-cadherin), this was unexpectedly accompanied by a comparable reduction in the mesenchymal marker *Cdh2* (N-cadherin). Similarly, while EMT is commonly driven by the expression of *Snai1/2* (*Snail* and *Slug*), *Twist1/2*, and *Zeb1/2*, we found either no change or a robust decrease in the expression of these genes [\(Figure 3C](#page-5-0)). These results indicate that the cell adhesion changes observed in mDKO trophoblast do not display the characteristics of a classic EMT and may be driven by independent or non-canonical pathways.

Contrary to what has been previously suggested ([Sakaue](#page-12-3) [et al., 2010\)](#page-12-3), these data show that oocyte methylation, directly or indirectly, regulates genes that are important for trophoblast differentiation and function. Notably, genes involved in signal transduction pathways that control adhesion and migration are particularly affected.

Methylation-Deficient TSCs Display Cell Adhesion

The fact of the state of the link between DNA methylation and cellular adhesion, we cultured TSCs null for all three active DNA methyltransferases (*Dnmt1*, *Dnmt3a*, and *Dnmt3b*) and

compared them with WT TSCs; both being derived from nuclear transfer embryos ([Sakaue et al., 2010\)](#page-12-3). We observed that WT cells grew as flat epithelial colonies with sharp boundaries as is characteristic of TSCs, whereas triple-knockout (TKO) TSCs did not form distinct colonies but instead exhibited a much more disorganized morphology and a notable loss of discrete colony margins [\(Figure 4](#page-6-0)A). In WT TSCs, E-cadherin delineated cell-cell junctions within colonies, whereas in TKO TSCs such junctions were less frequent and instead fibroblast-like cytoplasmic protrusions were visible, suggestive of an increased migratory capacity. Furthermore, using an assay to measure cell adhesion to tissue culture wells, we found that TKO TSCs were less adherent than WT cells on uncoated wells [\(Figure 4B](#page-6-0)). Interestingly, the difference could be rescued by laminin, suggesting that the adhesion defect can be overcome by changing the ECM/substrate composition. These differences were not due to cell differentiation, as TKO TSCs retained expression of key TSC markers (Figures S4 and [6G](#page-8-0)). In addition, TSCs derived from *Ascl2* mKO trophoblast do not display any morphological abnormalities compared with their WT counterparts (A.B.B. and L.L., unpublished data), ruling out a role for the loss of *Ascl2* imprinting in the TKO TSC phenotype.

To test whether loss of cell adhesion in TKO TSCs was driven by deregulation of the same genes identified in mDKO trophoblast, we plotted the expression changes of mDKO DE genes in TKO TSCs using published microarray data ([Figure 4](#page-6-0)C) ([Sa](#page-12-3)[kaue et al., 2010\)](#page-12-3). We found that mDKO upregulated genes are also derepressed in TKO TSCs, supporting a role for DNA

Figure 4. Loss of DNA Methylation Drives a Cell Adhesion Defect in TSCs

(A) E-cadherin staining of WT and TKO TSCs highlights that TKO TSCs do not display the typical epithelial colony morphology of ESCs, appearing to exhibit reduced cell adhesion and increased migration.

(B) TKO TSCs also have reduced adhesion capacity to cell culture wells, but only in the absence of the extracellular matrix component laminin.

(C) Gene expression differences (log2 fold change) between TKO and WT ESCs or TSCs for all mDKOderegulated genes. *Dnmt3* upregulated genes are derepressed in TKO TSCs (but not ESCs), consistent with a direct role of DNA methylation in their regulation, whereas downregulated genes are unchanged.

(D) RT-qPCR on TKO TSCs shows expression changes consistent with those seen in mDKO trophoblast (*Scml2* and *Spry1*) and raised expression of mesenchymal markers (*Cdh2* and *Snai2*). Bar plot error bars represent SD. **p < 0.01 , ***p < 0.001; t test comparingWT with TKO TSCs (B and D), or one-sample t test (C). See also Figure S4.

and widespread hypomethylation across several genomic features, including gene promoters, gene bodies, and LINE1 retrotransposons [\(Figure 5](#page-7-0)A). IAP retrotranspo-

methylation in their silencing. In contrast, mDKO downregulated genes were unaffected, suggesting that they are likely indirect effects that occur in the context of trophoblast differentiation. Interestingly, in contrast to TKO TSCs, TKO ESCs showed no changes in the expression of mDKO upregulated genes [\(Fig](#page-6-0)[ure 4C](#page-6-0)), indicating that TSC-specific transcription factors are required to activate these genes. In line with this hypothesis, we found that mDKO upregulated genes (but not downregulated genes) were enriched for sites bound by both ELF5 and TFAP2C $(p = 1.3 \times 10^{-10})$ ([Latos et al., 2015\)](#page-12-17).

Using RT-qPCR on WT and TKO TSCs we confirmed the derepression of *Scml2*, which was the top mDKO upregulated gene [\(Figure 4](#page-6-0)D). Equally concordant with the in vivo data was the downregulation of *Spry1*, which is a major EGFR signaling regulator that controls cell adhesion and migration [\(Mekkawy et al.,](#page-12-18) [2014\)](#page-12-18). We also detected an increase in the expression of the mesenchymal markers *Cdh2* and *Snai2* ([Figure 4](#page-6-0)D), although this was not accompanied by significant changes in *Cdh1* or *Snai1* (Figure S4), suggesting that, similarly to the mDKO trophoblast, canonical EMT is not involved in the loss of cell adhesion in TKO TSCs.

These results further support a critical role for DNA methylation in the regulation of cell adhesion in trophoblast.

A Subset of mDKO Upregulated Genes Are Controlled by

We next sought to identify which genes were likely to be regulated through the direct action of DNA methylation. To link gene expression changes with alterations in DNA methylation patterns, we performed whole-genome BS-seq of Ctrl, DHet, and DKO E7.5 EPCs. As expected, DKO EPCs showed deep sons were largely resistant to demethylation, consistent with DNMT1 being sufficient to maintain DNA methylation at these sites [\(Arand et al., 2012](#page-11-13)).

DHet EPCs displayed a slight reduction in DNA methylation across most genomic features [\(Figure 5](#page-7-0)A). To test whether this was solely due to a loss of oocyte methylation, we compared our data with published data on CGI methylation in oocytes and blastocysts [\(Kobayashi et al., 2012](#page-12-5)). Most CGIs that are partially resistant to demethylation during pre-implantation development are methylated in oocytes (Figure S5) [\(Smallwood et al., 2011\)](#page-12-4). We therefore asked whether these reprogramming-resistant CGIs were more likely to be hypomethylated in DHet EPCs when compared with de novo methylated CGIs. Our analysis shows that, overall, both groups of CGIs undergo a similar loss of methylation in DHet EPCs [\(Figures 5B](#page-7-0) and S5). In contrast, maternally methylated ICRs displayed a markedly more pronounced loss of methylation [\(Figure 5B](#page-7-0)). Therefore, it appears that the global hypomethylation seen in EPCs is a result of haploinsufficiency of *Dnmt3a* and/or *Dnmt3b* during the de novo methylation phase. However, the lack of correlation between the post-zygotic genotype and expression of mDKO DE genes (Figure S1C) argues against a major role of haploinsufficiency in driving the mDKO phenotype.

To exclude these small genome-wide differences in DNA methylation, we performed an unbiased search for differentially methylated regions (DMRs) between DHet and Ctrl EPCs that displayed at least a 20% methylation difference. We identified 6,685 DMRs, nearly all of which (96.6%) involved a loss of methylation in DHet EPCs and were seemingly not enriched at CGIs, promoters, gene bodies, or placental enhancers

Figure 5. DNA Methylation Differences Are Associated with DE Genes

(A) DNA methylation profiling by BS-seq shows that DHet EPCs have a slight genome-wide reduction of DNA methylation.

(B) CGIs methylated (>25%) in Ctrl EPCs were separated into reprogramming-resistant (>25% methylation in blastocyst; [Kobayashi et al., 2012\)](#page-12-5) or de novo methylated (<15% methylation in blastocyst). Methylation levels in Ctrl and DHet EPCs (left) and the relative methylation change between the two (right) shows that both subsets are hypomethylated to a similar extent in DHet EPCs; maternally methylated ICRs (blue) undergo more extensive methylation loss. (C) BS-seq profiles of methylation levels at example loci containing DHet DMRs (highlighted in green) that are associated with genes displaying altered gene expression.

See also Figure S5.

predicted from the mouse ENCODE project. Around 67% of DMRs had more than 80% methylation in WT oocytes ([Kobaya](#page-12-5)[shi et al., 2012](#page-12-5)), whereas only 46% of randomly generated regions passed the threshold, suggesting an important contribution of oocyte methylation to these DMRs. We identified 59 DMRs lying within 20 kb of mDKO DE genes (excluding *Ascl2*-dependent effects), covering 35 out of 94 DE genes (see examples in [Figure 5C](#page-7-0)). Surprisingly, these DMRs were associated with both upregulated ($n = 14$) and downregulated $(n = 21)$ genes. To further probe which gene expression alterations were likely to be directly regulated by DNA methylation, we asked which genes were also affected in TKO TSCs. Notably, 6 out of 14 DMR-associated genes that were upregulated in mDKO EPCs were also more than 2-fold upregulated in TKO TSCs. Five of these genes (*Dst*, *Plekha6*, *Stk10*, *Ptpre*, and *Plcb4*) encoded an actin-binding protein, a phospholipid-binding adaptor, a protein kinase, a protein tyrosine phosphatase, and a phospholipase that are implicated in cell adhesion and signal transduction.

These data reveal that a number of oocyte-dependent DMRs are linked to genes deregulated in mDKO EPCs, a subset of which appear to be directly controlled by DNA methylation in a trophoblast-specific manner.

Sc*ml2* Is Hypomethylated in mDKO EPCs and Affects
Trophoblast Differentiation and Adhesion

Trophoblast Differentiation and Adhesion One of the DMRs we identified overlapped *Scml2* ([Figure 5](#page-7-0)C), which was the most upregulated gene in mDKO EPCs, and was also highly upregulated in TKO TSCs ([Figure 4D](#page-6-0)). SCML2 is a non-canonical member of the Polycomb Repressive Complex 1 (PRC1), which plays important epigenetic roles in the establishment of the male germline [\(Hasegawa et al., 2015;](#page-11-14) [Luo et al., 2015\)](#page-11-14). We therefore investigated whether *Scml2* silencing was important for trophoblast development and adhesion. Firstly, we confirmed that *Scml2* upregulation in mDKO EPCs was driven by the maternal deletion of *Dnmt3a* [\(Fig](#page-8-0)[ure 6](#page-8-0)A). We then validated the DMR associated with *Scml2* using Sequenom MassARRAY, which showed a specific loss of methylation in DHet and DKO EPCs near an intragenic tran-scription start site (TSS) ([Figure 6](#page-8-0)B). This alternative promoter overlaps a CGI (highlighted in [Figure 5](#page-7-0)C) that is 95% methylated in WT oocytes (losing all methylation in *Dnmt3l*-null

Figure 6. Scml2 Is Controlled by DNA Methylation and Affects SynT Formation and Cell Adhesion

(A) RT-qPCR of *Dnmt3a* mKO EPCs confirms that *Scml2* is controlled by oocyte methylation.

(B) Methylation analysis by Sequenom MassARRAY in E7.5 male EPCs, confirming the DMR at an intragenic TSS of *Scml2*. Each data point may include more than one CpG from the amplicon, as indicated on the x axis.

(C) RT-qPCR analysis of TSCs grown in FGF+ (TSC conditions) or FGF (differentiation conditions) medium for 6 days, with or without *Scml2* overexpression. (D) Expression of *Syna* is reduced in *Dnmt3a* mKO EPCs, whereas markers of SynT-II *Synb* and *Cebpa* are unaffected.

(E) E-cadherin staining of two independent *Scml2* knockout clones from TKO TSCs shows a rescue of the morphological alterations seen in TKO TSCs.

(F) *Scml2* KO on TKO TSCs also rescues the defect in cell adhesion to cell culture wells in the absence of laminin.

(G) RT-qPCR analysis of TKO *Scml2* KO clones shows maintained expression of the TSC marker *Cdx2*; the expression of genes involved in cell adhesion is not rescued upon *Scml2* deletion.

Error bars represent SD. *p < 0.05, **p < 0.01, ***p < 0.001; t test comparing WT and *Dnmt3a* mKO EPCs (A and D) or *Scml2*-expressing TSCs versus vector control (C), or ANOVA with post hoc tests comparing Ctrl with DHet/DKO (B) or TKO TSC lines with WT TSCs. See also Figure S6.

oocytes) and 34% methylated in blastocysts [\(Kobayashi et al.,](#page-12-5) [2012\)](#page-12-5), supporting the notion that *Scml2* methylation levels are carried over from the oocyte through the pre-implantation phase. Interestingly, maintenance of methylation and silencing of *Scml2* are specific to the trophoblast compartment, as methylation in the epiblast is lost and *Scml2* is expressed (Figure S6A). This pattern is also clearly observed when comparing ES and TS cells (Figure S6A). *Scml2* lies on the X chromosome, but while *Scml2* is unmethylated in X-containing sperm ([Ko](#page-12-5)[bayashi et al., 2012](#page-12-5)), no placental *Scml2* expression is expected from the paternal allele in females due to imprinted X inactivation in mouse trophoblast. Accordingly, we found no difference in *Scml2* expression levels between male and female EPCs within Ctrl or mDKO genotypes. Furthermore, X inactivation appeared unaffected in female DHet EPCs, as *Xist* expression was unchanged. Importantly, the X-linked nature of *Scml2* makes its epigenetic regulation distinguishable from a genomic imprinting mechanism.

To test the effect of *Scml2* expression on trophoblast morphology and development, we overexpressed *Scml2* in TSCs (*Scml2* expression levels in Figure S6C). When cells were grown under stem cell conditions (fibroblast growth factor [FGF]+), we found no significant differences in morphology or cell adhesion (Figure S6B) upon overexpression of *Scml2*. There were also no detectable differences in the expression of key TSC markers (Figure S6C). However, after induction of TSC differentiation by removal of FGF from the medium, *Scml2* impaired the expression of Syncytin A (*Syna*), a marker of SynT-I (which interfaces the maternal blood), while no effect was seen on other differentiation markers, including markers of TGCs and SynT-II ([Figure 6C](#page-8-0) and data not shown). This suggests that *Scml2* silencing is important for SynT-I formation in WT trophoblast. Accordingly, *Dnmt3a* mKO trophoblast also displayed reduced *Syna* expression at E7.5 but no alterations in SynT-II markers *Synb* and *Cebpa* ([Figure 6](#page-8-0)D). While we found that the latter markers were markedly reduced at E9.5, this reflects the absence of the developing labyrinthine layer observed in these mutants. However, the early change seen in *Syna* expression at E7.5 occurs prior to the morphological establishment of SynT and the labyrinth, suggesting that it is a direct effect of methylation deficiency. In contrast, while *Ascl2* mKO trophoblast also displays a reduction in *Syna* expression at E9.5 due to a reduction of the labyrinthine layer [\(Oh-McGinnis et al., 2011](#page-12-19)), no difference is detected at E7.5 (Figure S6D).

As SCML2 was not sufficient to drive cell adhesion defects, we asked whether it was necessary for the phenotype, in combination with other methylation-dependent expression changes. We therefore used CRISPR/Cas9 gene editing to delete *Scml2* in TKO TSCs (Figure S6E). Strikingly, both *Scml2* KO clones that were established displayed a distinct morphology from the parental TKO TSCs, closely resembling the WT colony arrangement [\(Figure 6E](#page-8-0)). TKO *Scml2* KO clones also showed a restored ability to attach to cell culture wells comparable with WT levels ([Figure 6](#page-8-0)F). This was not due to changes in cell differentiation state, as cells maintained expression of TSC markers and did not show upregulation of differentiation markers [\(Figure 6](#page-8-0)G and data not shown). Loss of SCML2 did not rescue the rise in *Cdh2* expression seen in TKO TSCs ([Figure 6](#page-8-0)G). We then asked whether loss of SCML2 was rescuing the cell adhesion defect via modulation of some of the other DE genes identified in mDKO trophoblast. None of the nine genes tested (e.g., *Spry1* and *Itga7*) showed significant differences between TKO TSCs and the TKO *Scml2* KO clones [\(Figure 6G](#page-8-0)), suggesting that *Scml2* acts through an independent pathway. Impairment of cell adhesion in TKO TSCs may therefore require both SCML2-dependent pathways together with other methylation-dependent alterations seen in vivo. This is in line with SCML2 overexpression being insufficient to drive cell adhesion changes (Figure S6B).

Our phenotypic, molecular, and functional analyses show that, contrary to previous reports [\(Sakaue et al., 2010](#page-12-3)), DNA methylation is essential for early trophoblast development. In particular, our work demonstrates that maternal methylation (and its maintenance during pre-implantation development) plays a major regulatory role in trophoblast differentiation and function. This encompasses the need for controlling imprinted genes (*Ascl2*), but also non-imprinted genes, as demonstrated by the specific example of *Scml2*, which we have shown to affect *Syna* expression and cell adhesion. Moreover, methylation at the *Scml2* promoter is specifically maintained in the EPC and lost in the epiblast (Figure S6A), highlighting the need for targeting DNA methylation to the trophoblast compartment at specific loci. Loss of imprinting at methylation-dependent loci other than *Ascl2* is also known to have an impact on placental function at later developmental stages [\(Tunster et al., 2013](#page-12-20)), further emphasizing the importance of maternal methylation.

Although non-imprinted oocyte methylation marks have been generally associated with brain- and testis-linked genes ([Rut](#page-12-21)[ledge et al., 2014](#page-12-21)), we have identified critical trophoblast genes that are regulated through maternal DNA methylation (directly or indirectly). Deregulated genes were enriched for signal transduction and regulators of Ras and Rho family GTPases, which are implicated in cell adhesion and migration. Along with the decreased *Cdh1* expression seen in E9.5 DHet trophoblast [\(Fig](#page-5-0)[ure 3C](#page-5-0)), these alterations are in line with the cellular adhesion phenotype seen in mDKO TGCs. Interestingly, links between DNA methylation and EMT have been described in human trophoblast cell lines [\(Chen et al., 2013a; 2013b\)](#page-11-15). Epigenetic regulation of mouse trophoblast migration and invasion has also been described through the action of the histone lysine demethylase LSD1 ([Zhu et al., 2014](#page-12-22)). Similar to some of our observations, LSD1 depletion in TSCs leads to early differentiation onset, which disrupts their epithelial morphology and increases cell migration and invasion. However, we only found one gene (*Reep6*) in common between mDKO DE genes and LSD1-regulated genes, suggesting that separate epigenetic mechanisms act on different pathways to regulate the crucial processes of cell adhesion and control of invasive behavior in trophoblast.

We uncovered *Scml2* as a putative methylation-controlled gene that is kept silent in the trophoblast lineage to allow for appropriate control of cell adhesion and *Syna* expression. The relatively low methylation levels of *Scml2* in Ctrl EPCs suggest that other mechanisms aid in its silencing. We also cannot completely exclude the possibility that *Scml2* expression is indirectly controlled by DNA methylation. In the male germline, SCML2 regulates PRC1-dependent ubiquitination of histone H2A either positively or negatively, in a context- and targetdependent manner [\(Hasegawa et al., 2015](#page-11-14)). SCML2 may therefore regulate the expression of genes involved in the control of cell adhesion and migration, as well as of *Syna*. Notably, based on our *Scml2* overexpression experiment, hypomethylation of *Scml2* is not sufficient to drive the adhesion defects, implicating maternal DNA methylation more widely in the regulation of cell adhesion and migration, as discussed above. Interestingly, human SCML2 interacts with SFMBT2 [\(Zhang et al., 2013\)](#page-12-23), another PcG protein whose ortholog in mouse is essential for placental development [\(Miri et al., 2013](#page-12-24)). *Sfmbt2* is a paternally expressed imprinted gene, but silencing of the maternal allele is independent of DNA methylation [\(Okae et al., 2012](#page-12-25)). However, the epigenetic control of *SCML2* during human pre-implantation development appears to differ from that of the mouse, as the syntenic region to the DMR that we identified (also at a

TSS and CGI in the human) is largely unmethylated in human oocytes and morulae (Figure S6F) [\(Guo et al., 2014\)](#page-11-16). Differences in placental morphology and organization may ultimately be driven by such gene-regulatory changes throughout evolution. Notably, the X chromosome is particularly enriched in trophoblast-associated genes and has been argued to be an important driver of placental evolution and speciation ([Hemberger, 2002\)](#page-11-17). Along with this argument, it is interesting to note that *Scml2* is X-linked exclusively in eutherians.

Our study highlights a critical and widespread role of oocyte methylation in the development of the placenta. Although we cannot completely rule out that other imprinted genes are involved in the early mDKO phenotype we describe, their known placental roles [\(Tunster et al., 2013\)](#page-12-20) and our data strongly suggest that additional, non-imprinted methylation marks also control trophoblast development. Interestingly, when we measured human oocyte and morula methylation at regions syntenic to our mouse DMRs, we found that CpGs within these regions are generally hypermethylated when compared with the rest of the genome (Figure S6G). It is tempting to speculate that conservation of methylation at these sites is relevant for human placental development, and that oocyte methylation evolved to play a major role in the trophoblast lineage.

All experimental procedures were performed under licenses by the Home Office (UK) in accordance with the Animals (Scientific Procedures) Act 1986. Mice carrying conditional deletions of both *Dnmt3a* and *Dnmt3b* ([Okano](#page-12-2) [et al., 1999\)](#page-12-2), or of *Dnmt3a* alone, and with or without a *Zp3-Cre* transgene were crossed by natural mating. Female mice heterozygous for an *Ascl2 lacZ* allele ([Tanaka et al., 1999](#page-12-8)) were crossed to WT males. For immunofluorescence, E7.5 implantation sites were fixed with 4% paraformaldehyde (PFA) and processed as for routine paraffin histology. For RNA and DNA isolation, trophoblast and epiblast tissues were dissected from E7.5 or E9.5 conceptuses and snap-frozen.

Tissue Culture and Scml2 Overexpression/Knockout

Blastocyst-derived TS-EGFP cells (a kind gift from Dr. Janet Rossant) or WT/ TKO TSCs derived from nuclear transfer embryos ([Sakaue et al., 2010\)](#page-12-3) were cultured under routine conditions (20% fetal bovine serum, 1 mM Na-pyruvate, penicillin/streptomycin, 50 µM 2-mercaproethanol, 25 ng/ml basic FGF (Sigma) and 1 μ g/ml heparin in RPMI 1640, with 70% of the medium pre-conditioned on embryonic feeder cells). For *Scml2* overexpression, the open reading frame of the isoform that starts at the intragenic CGI-associated TSS was cloned via Gateway cloning (Life Technologies) into a PiggyBac vector and sequence-verified. This construct or the empty vector were co-transfected with a PiggyBac transposase plasmid using Lipofectamine 2000 (Life Technologies) or Fugene 6 (Promega), according to the manufacturer's instructions. Integrants were selected with $5 \mu g/ml$ blasticidin S, after which cells were switched to TS cell differentiation medium (unconditioned medium without basic FGF or heparin) for the indicated times. For CRISPR/Cas9-mediated deletion of *Scml2*, TKO TSCs were co-transfected with pCAG-cGFPblasticidin and pCas9.2A.GFP/Puro ([Ran et al., 2013](#page-12-26)) harboring guide RNAs flanking exon 3 (Figure S6D): GTTCATCCCTAGGCAATTAT, CAGGGA TGTTTGCAACGTGC. After 48 hr of blasticidin selection, single cells were sorted by flow cytometry and left to grow for 10–14 days before genotyping (see Supplemental Experimental Procedures, Primers).

Histology and Immunofluorescence

Histology and Immunofluorescence Paraffin sections were deparaffinized with Histo-Clear and dehydrated through an ethanol series, followed either by standard H&E staining or antigen retrieval by boiling slides for 30 min (in 1 mM EDTA, 0.05% Tween 20, pH 8) and cooling at room temperature for 20 min. After blocking with 1% BSA overnight, sections were incubated with a rabbit monoclonal anti-CDX2 antibody (EPR2764Y, Novus Biologicals, 1:250 dilution) for 2 hr. Secondary detection was done with an AlexaFluor 488 anti-rabbit antibody (Life Technologies, 1:500 dilution). TSCs were fixed with 4% PFA, permeabilized with Triton X-100, blocked as above, and labeled with a mouse anti-CDH1 antibody (BD 610181, BD Biosciences, 1:400 dilution).

Adhesion of TSCs to cell-culture wells was performed using the Vybrant cell adhesion assay kit (Life Technologies) according to the manufacturer's instructions, with the following details/modifications: 10⁵ cells were plated per well of a 96-well tissue culture plate, either uncoated or coated with laminin, and left to attach for 2 hr in serum-free RPMI medium.

Genomic DNA and RNA were isolated from the same samples using the Allprep DNA/RNA Kit (Qiagen). RNA was DNase-treated with the DNA-free kit (Life Technologies). DNA was bisulfite-converted using the Imprint DNA Modification Kit (Sigma).

RT-aPCR

For RT-qPCR analysis, total RNA was reverse transcribed by random priming using the RevertAid First Strand cDNA Synthesis Kit (Thermo Scientific). qPCR was performed using Brilliant III Ultra-Fast SYBR Green qPCR Master Mix (Agilent Technologies) or Mesa Blue qPCR MasterMix Plus (Eurogentec) using gene-specific primers (see Supplemental Experimental Procedures).

PCR was performed on bisulfite-converted DNA using HotStarTaq DNA Polymerase (Qiagen) and target-specific primers (see Supplemental Experimental Procedures); samples were processed using the "T" Cleavage MassCLEAVE Reagent Kit (Agena Bioscience) and subjected to MALDI-TOF analysis using the MassARRAY system, according to the manufacturer's instructions.

mar Seq Library Generation and Sequencing
mRNA was purified from 45–400 ng of total RNA using a Dynabeads mRNA Purification Kit (Life Technologies). For each genotype, three strand-specific libraries were generated from single EPCs using the ScriptSeg v2 RNA-Seg Library Preparation Kit (Epicentre), according to the manufacturer's instructions. Indexed libraries were pooled and sequenced on an Illumina HiSeq 2000 or 2500 platform using 50 bp (*Dnmt3* libraries) or 100 bp (*Ascl2* libraries) single-end reads. Reads were trimmed using ''Trim Galore!'' and mapped to the NCBIM37 genome assembly using TopHat ([Trapnell et al., 2009](#page-12-27)).

Whole-genome BS-seq libraries were prepared from 25–70 ng of genomic DNA, using a post-bisulfite adaptor tagging protocol, as previously described ([Peat et al., 2014](#page-12-28)). In brief, first-strand synthesis was performed on bisulfiteconverted DNA using biotin-tagged random primers containing part of the Illumina-compatible 5' adaptor sequence. After biotin capture using Dynabeads (Life Technologies), second-strand synthesis was performed using random primers containing part of the Illumina-compatible 3' adaptor sequence. DNA was then eluted from the beads and libraries amplified using Phusion (New England Biolabs). Libraries were sequenced on an Illumina HiSeq 2500 platform using 100-bp paired-end reads. Reads were trimmed using ''Trim Galore!'' and mapped to the NCBIM37 genome assembly using bismark ([Krueger and Andrews, 2011](#page-12-29)).

All data analyses were performed using Seqmonk and/or R scripts. Differential gene expression analysis was performed with DEseq ([Anders and Huber,](#page-11-18) [2010\)](#page-11-18), using a 1% false discovery rate cut-off and minimum fold change of 2. GO analysis was performed using topGO. To measure DNA methylation at given genomic features, methylation calls from both strands at CpG sites were pooled; regions that had at least three CpGs covered by at least five reads were selected and the average CpG methylation value per region calculated. Promoters were defined as -1 kb to $+500$ bp relative to each TSS; CGI

annotation was from [Illingworth et al. \(2008\).](#page-11-19) For DMR detection, methylation calls from running windows containing five CpGs within 1 kb were pooled; significant differences were determined using a Fisher test and a Benjamini-Hochberg corrected p value cut-off of 0.01. Significantly different windows within 200 bp were merged into DMRs, and only DMRs with a methylation difference larger than 20% were kept. LiftOver was used to identify regions of synteny to the identified DMRs in human. Of 4,084 syntenic regions identified, 353 were covered by at least five reads in human oocyte and morula reduced representation bisulfite sequencing data ([Guo et al., 2014](#page-11-16)) and used for further analysis.

ACCESSION NUMBERS

The accession number for the RNA-seq and BS-seq data reported in this paper is GEO: GSE66049.

Supplemental Information includes Supplemental Experimental Procedures, six figures, and three tables and can be found with this article online at [http://dx.doi.org/10.1016/j.devcel.2015.12.027.](http://dx.doi.org/10.1016/j.devcel.2015.12.027)

AUTHOR CONTRIBUTIONS

M.R.B., W.D., M.H., and W.R. designed the study and experiments. M.R.B. performed histology/immunofluorescence, RT-qPCR, TSC experiments, and bioinformatics analyses. M.K. performed RNA-seq, BS-seq, RT-qPCR, and Sequenom experiments. V.P. performed CRISPR/Cas9 experiments. A.B.B. and L.L. isolated tissue from *Ascl2* KOs. M.C. and E.F. prepared histological sections. W.D. isolated tissue from *Dnmt3* KOs. M.H. performed TSC experiments. S.J.C. interpreted RNA-seq and RT-qPCR data. M.R.B. wrote the manuscript with all other authors.

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REFERENCES

[Anders, S., and Huber, W. \(2010\). Differential expression analysis for sequence](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref1) [count data. Genome Biol.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref1) *11*, R106.

[Arand, J., Spieler, D., Karius, T., Branco, M.R., Meilinger, D., Meissner, A.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref2) [Jenuwein, T., Xu, G., Leonhardt, H., Wolf, V., et al. \(2012\). In vivo control of](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref2) [CpG and non-CpG DNA methylation by DNA methyltransferases. PLoS](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref2) Genet. *8*[, e1002750](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref2).

[Arima, T., Hata, K., Tanaka, S., Kusumi, M., Li, E., Kato, K., Shiota, K., Sasaki,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref3) [H., and Wake, N. \(2006\). Loss of the maternal imprint in Dnmt3Lmat](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref3)-[/](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref3)- [mice](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref3) [leads to a differentiation defect in the extraembryonic tissue. Dev. Biol.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref3) *297*, [361–373](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref3).

[Auclair, G., Guibert, S., Bender, A., and Weber, M. \(2014\). Ontogeny of CpG](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref4) [island methylation and specificity of DNMT3 methyltransferases during embry](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref4)[onic development in the mouse. Genome Biol.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref4) *15*, 545.

Borgel, J., Guibert, S., Li, Y., Chiba, H., Schübeler, D., Sasaki, H., Forné, T., [and Weber, M. \(2010\). Targets and dynamics of promoter DNA methylation](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref5) [during early mouse development. Nat. Genet.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref5) *42*, 1093–1100.

[Bourc'his, D., Xu, G.L., Lin, C.S., Bollman, B., and Bestor, T.H. \(2001\). Dnmt3L](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref6) [and the establishment of maternal genomic imprints. Science](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref6) *294*, 2536–2539.

[Cattanach, B.M., and Kirk, M. \(1985\). Differential activity of maternally and](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref7) [paternally derived chromosome regions in mice. Nature](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref7) *315*, 496–498.

[Chen, Y., Wang, K., and Leach, R. \(2013a\). 5-Aza-dC treatment induces](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref8) [mesenchymal-to-epithelial transition in 1st trimester trophoblast cell line](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref8) [HTR8/SVneo. Genes. Cells.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref8) *432*, 116–122.

[Chen, Y., Wang, K., Qian, C.-N., and Leach, R. \(2013b\). DNA methylation is](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref9) [associated with transcription of Snail and Slug genes. Biochem. Biophys.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref9) [Res. Commun.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref9) *430*, 1083–1090.

[Cieply, B., Riley, P., Pifer, P.M., Widmeyer, J., Addison, J.B., Ivanov, A.V.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref10) [Denvir, J., and Frisch, S.M. \(2012\). Suppression of the epithelial-mesenchymal](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref10) [transition by Grainyhead-like-2. Cancer Res.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref10) *72*, 2440–2453.

[Dodge, J.E., Okano, M., Dick, F., Tsujimoto, N., Chen, T., Wang, S., Ueda, Y.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref11) [Dyson, N., and Li, E. \(2005\). Inactivation of Dnmt3b in mouse embryonic fibro](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref11)[blasts results in DNA hypomethylation, chromosomal instability, and sponta](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref11)[neous immortalization. J. Biol. Chem.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref11) *280*, 17986–17991.

[El-Hashash, A.H.K., and Kimber, S.J. \(2006\). PTHrP induces changes in cell](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref12) [cytoskeleton and E-cadherin and regulates Eph/Ephrin kinases and](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref12) [RhoGTPases in murine secondary trophoblast cells. Dev. Biol.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref12) *290*, 13–31.

[Frank, D., Fortino, W., Clark, L., Musalo, R., Wang, W., Saxena, A., Li, C.-M.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref13) [Reik, W., Ludwig, T., and Tycko, B. \(2002\). Placental overgrowth in mice lack](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref13)[ing the imprinted gene Ipl. Proc. Natl. Acad. Sci. USA](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref13) *99*, 7490–7495.

[Guillemot, F., Nagy, A., Auerbach, A., Rossant, J., and Joyner, A.L. \(1994\).](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref14) [Essential role of Mash-2 in extraembryonic development. Nature](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref14) *371*, [333–336](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref14).

[Guo, H., Zhu, P., Yan, L., Li, R., Hu, B., Lian, Y., Yan, J., Ren, X., Lin, S., Li, J.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref15) [et al. \(2014\). The DNA methylation landscape of human early embryos. Nature](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref15) *511*[, 606–610.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref15)

[Hackett, J.A., Reddington, J.P., Nestor, C.E., Dunican, D.S., Branco, M.R.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref16) [Reichmann, J., Reik, W., Surani, M.A., Adams, I.R., and Meehan, R.R.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref16) [\(2012\). Promoter DNA methylation couples genome-defence mechanisms to](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref16) [epigenetic reprogramming in the mouse germline. Development](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref16) *139*, 3623– [3632](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref16).

[Harris, L.K., Jones, C.J.P., and Aplin, J.D. \(2009\). Adhesion molecules in hu](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref17)[man trophoblast - a review. II. extravillous trophoblast. Placenta](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref17) *30*, 299–304.

[Hasegawa, K., Sin, H.-S., Maezawa, S., Broering, T.J., Kartashov, A.V.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref18) [Alavattam, K.G., Ichijima, Y., Zhang, F., Bacon, W.C., Greis, K.D., et al.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref18) [\(2015\). SCML2 establishes the male germline epigenome through regulation](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref18) [of histone H2A ubiquitination. Dev. Cell](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref18) *32*, 574–588.

[Hemberger, M. \(2002\). The role of the X chromosome in mammalian extra em](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref19)[bryonic development. Cytogenet. Genome Res.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref19) *99*, 210–217.

[Hemberger, M., Nozaki, T., Masutani, M., and Cross, J.C. \(2003\). Differential](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref20) [expression of angiogenic and vasodilatory factors by invasive trophoblast gi](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref20)[ant cells depending on depth of invasion. Dev. Dyn.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref20) *227*, 185–191.

[Hunkapiller, N.M., Gasperowicz, M., Kapidzic, M., Plaks, V., Maltepe, E.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref21) [Kitajewski, J., Cross, J.C., and Fisher, S.J. \(2011\). A role for Notch signaling](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref21) [in trophoblast endovascular invasion and in the pathogenesis of pre](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref21)[eclampsia. Development](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref21) *138*, 2987–2998.

[Illingworth, R., Kerr, A., Desousa, D., Jørgensen, H., Ellis, P., Stalker, J.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref22) [Jackson, D., Clee, C., Plumb, R., Rogers, J., et al. \(2008\). A novel CpG island](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref22) [set identifies tissue-specific methylation at developmental gene loci. PLoS](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref22) Biol. *6*[, e22.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref22)

[Kaneda, M., Okano, M., Hata, K., Sado, T., Tsujimoto, N., Li, E., and Sasaki, H.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref23) [\(2004\). Essential role for de novo DNA methyltransferase Dnmt3a in paternal](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref23) [and maternal imprinting. Nature](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref23) *429*, 900–903.

[Kaneda, M., Hirasawa, R., Chiba, H., Okano, M., Li, E., and Sasaki, H. \(2010\).](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref24) [Genetic evidence for Dnmt3a-dependent imprinting during oocyte growth ob](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref24)[tained by conditional knockout with Zp3-Cre and complete exclusion of](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref24) [Dnmt3b by chimera formation. Genes Cells](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref24) *15*, 169–179.

[Kibschull, M., Colaco, K., Matysiak-Zablocki, E., Winterhager, E., and Lye, S.J.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref25) [\(2014\). Connexin31.1 \(Gjb5\) deficiency blocks trophoblast stem cell differenti](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref25)[ation and delays placental development. Stem Cells Dev.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref25) *23*, 2649–2660.

[Kobayashi, H., Sakurai, T., Imai, M., Takahashi, N., Fukuda, A., Yayoi, O., Sato,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref26) [S., Nakabayashi, K., Hata, K., Sotomaru, Y., et al. \(2012\). Contribution of intra](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref26)[genic DNA methylation in mouse gametic DNA methylomes to establish](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref26) [oocyte-specific heritable marks. PLoS Genet.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref26) *8*, e1002440.

[Kokkinos, M.I., Murthi, P., Wafai, R., Thompson, E.W., and Newgreen, D.F.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref27) [\(2010\). Cadherins in the human placenta—epithelial-mesenchymal transition](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref27) [\(EMT\) and placental development. Placenta](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref27) *31*, 747–755.

[Krueger, F., and Andrews, S.R. \(2011\). Bismark: a flexible aligner and methyl](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref28)[ation caller for Bisulfite-Seq applications. Bioinformatics](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref28) *27*, 1571–1572.

[Lane, N., Dean, W., Erhardt, S., Hajkova, P., Surani, A., Walter, J., and Reik, W.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref29) [\(2003\). Resistance of IAPs to methylation reprogramming may provide a](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref29) [mechanism for epigenetic inheritance in the mouse. Genesis](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref29) *35*, 88–93.

[Latos, P.A., Sienerth, A.R., Murray, A., Senner, C.E., Muto, M., Ikawa, M.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref30) [Oxley, D., Burge, S., Cox, B.J., and Hemberger, M. \(2015\). Elf5-centered tran](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref30)[scription factor hub controls trophoblast stem cell self-renewal and differenti](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref30)[ation through stoichiometry-sensitive shifts in target gene networks. Genes](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref30) Dev. *29*[, 2435–2448](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref30).

[Li, E., Bestor, T.H., and Jaenisch, R. \(1992\). Targeted mutation of the DNA](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref31) [methyltransferase gene results in embryonic lethality. Cell](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref31) *69*, 915–926.

[Luo, M., Zhou, J., Leu, N.A., Abreu, C.M., Wang, J., Anguera, M.C., de Rooij,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref32) [D.G., Jasin, M., and Wang, P.J. \(2015\). Polycomb protein SCML2 associates](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref32) [with USP7 and counteracts histone H2A ubiquitination in the XY chromatin](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref32) [during male meiosis. PLoS Genet.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref32) *11*, e1004954.

[Maecker, H.T., and Levy, S. \(1997\). Normal lymphocyte development but de](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref33)[layed humoral immune response in CD81-null mice. J. Exp. Med.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref33) *185*, 1505– [1510.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref33)

[Mekkawy, A.H., Pourgholami, M.H., and Morris, D.L. \(2014\). Human Sprouty1](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref34) [suppresses growth, migration, and invasion in human breast cancer cells.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref34) [Tumour Biol.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref34) *35*, 5037–5048.

[Menke, A., and Giehl, K. \(2012\). Regulation of adherens junctions by Rho](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref35) [GTPases and p120-catenin. Arch. Biochem. Biophys.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref35) *524*, 48–55.

[Miri, K., Latham, K., Panning, B., Zhong, Z., Andersen, A., and Varmuza, S.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref36) [\(2013\). The imprinted polycomb group gene Sfmbt2 is required for trophoblast](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref36) [maintenance and placenta development. Development](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref36) *140*, 4480–4489.

[Morasso, M.I., Grinberg, A., Robinson, G., Sargent, T.D., and Mahon, K.A.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref37) [\(1999\). Placental failure in mice lacking the homeobox gene Dlx3. Proc. Natl.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref37) [Acad. Sci. USA](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref37) *96*, 162–167.

[Oh-McGinnis, R., Bogutz, A.B., and Lefebvre, L. \(2011\). Partial loss of Ascl2](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref38) [function affects all three layers of the mature placenta and causes intrauterine](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref38) [growth restriction. Dev. Biol.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref38) *351*, 277–286.

[Okae, H., Hiura, H., Nishida, Y., Funayama, R., Tanaka, S., Chiba, H.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref39) [Yaegashi, N., Nakayama, K., Sasaki, H., and Arima, T. \(2012\). Re-investigation](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref39) [and RNA sequencing-based identification of genes with placenta-specific im](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref39)[printed expression. Hum. Mol. Genet.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref39) *21*, 548–558.

[Okano, M., Bell, D.W., Haber, D.A., and Li, E. \(1999\). DNA methyltransferases](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref40) [Dnmt3a and Dnmt3b are essential for de novo methylation and mammalian](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref40) [development. Cell](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref40) *99*, 247–257.

[Pan, Z., Sikandar, S., Witherspoon, M., Dizon, D., Nguyen, T., Benirschke, K.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref41) [Wiley, C., Vrana, P., and Lipkin, S.M. \(2008\). Impaired placental trophoblast](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref41) [lineage differentiation in Alkbh1\(-/-\) mice. Dev. Dyn.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref41) *237*, 316–327.

[Parast, M.M., Aeder, S., and Sutherland, A.E. \(2001\). Trophoblast giant-cell](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref42) [differentiation involves changes in cytoskeleton and cell motility. Dev. Biol.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref42) *230*[, 43–60](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref42).

[Peat, J.R., Dean, W., Clark, S.J., Krueger, F., Smallwood, S.A., Ficz, G., Kim,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref43) [J.K., Marioni, J.C., Hore, T.A., and Reik, W. \(2014\). Genome-wide bisulfite](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref43) [sequencing in zygotes identifies demethylation targets and maps the contribu](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref43)[tion of TET3 oxidation. Cell Rep.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref43) *9*, 1990–2000.

[Peters, J., and Robson, J.E. \(2008\). Imprinted noncoding RNAs. Mamm.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref44) Genome *19*[, 493–502](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref44).

Pollheimer, J., and Knö[fler, M. \(2005\). Signalling pathways regulating the inva](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref45)[sive differentiation of human trophoblasts: a review. Placenta](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref45) *26* (*Suppl A*), [S21–S30](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref45).

[Ralston, A., Cox, B.J., Nishioka, N., Sasaki, H., Chea, E., Rugg-Gunn, P.J.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref46) [Guo, G., Robson, P., Draper, J.S., and Rossant, J. \(2010\). Gata3 regulates](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref46) [trophoblast development downstream of Tead4 and in parallel to Cdx2.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref46) [Development](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref46) *137*, 395–403.

[Ran, F.A., Hsu, P.D., Wright, J., Agarwala, V., Scott, D.A., and Zhang, F. \(2013\).](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref47) [Genome engineering using the CRISPR-Cas9 system. Nat Protoc.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref47) *8*, 2281– [2308.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref47)

[Rutledge, C.E., Thakur, A., O'Neill, K.M., Irwin, R.E., Sato, S., Hata, K., and](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref48) [Walsh, C.P. \(2014\). Ontogeny, conservation and functional significance of](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref48) [maternally inherited DNA methylation at two classes of non-imprinted genes.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref48) [Development](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref48) *141*, 1313–1323.

[Sakaue, M., Ohta, H., Kumaki, Y., Oda, M., Sakaide, Y., Matsuoka, C.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref49) [Yamagiwa, A., Niwa, H., Wakayama, T., and Okano, M. \(2010\). DNA methyl](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref49)[ation is dispensable for the growth and survival of the extraembryonic line](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref49)[ages. Curr. Biol.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref49) *20*, 1452–1457.

[Sher, N., Stetina Von, J.R., Bell, G.W., Matsuura, S., Ravid, K., and Orr-](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref50)[Weaver, T.L. \(2013\). Fundamental differences in endoreplication in mammals](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref50) [and Drosophila revealed by analysis of endocycling and endomitotic cells.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref50) [Proc. Natl. Acad. Sci. USA](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref50) *110*, 9368–9373.

[Shirane, K., Toh, H., Kobayashi, H., Miura, F., Chiba, H., Ito, T., Kono, T., and](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref51) [Sasaki, H. \(2013\). Mouse oocyte methylomes at base resolution reveal](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref51) [genome-wide accumulation of Non-CpG methylation and role of DNA methyl](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref51)[transferases. PLoS Genet.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref51) *9*, e1003439.

[Smallwood, S.A., Tomizawa, S.-I., Krueger, F., Ruf, N., Carli, N., Segonds-](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref52)[Pichon, A., Sato, S., Hata, K., Andrews, S.R., and Kelsey, G. \(2011\).](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref52) [Dynamic CpG island methylation landscape in oocytes and preimplantation](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref52) [embryos. Nat. Genet.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref52) *43*, 811–814.

[Smith, Z.D., Chan, M.M., Mikkelsen, T.S., Gu, H., Gnirke, A., Regev, A., and](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref53) [Meissner, A. \(2012\). A unique regulatory phase of DNA methylation in the early](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref53) [mammalian embryo. Nature](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref53) *484*, 339–344.

[Sutherland, A. \(2003\). Mechanisms of implantation in the mouse: differentia](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref54)[tion and functional importance of trophoblast giant cell behavior. Dev. Biol.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref54) *258*[, 241–251](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref54).

[Sutherland, A.E., Calarco, P.G., and Damsky, C.H. \(1993\). Developmental](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref55) [regulation of integrin expression at the time of implantation in the mouse em](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref55)[bryo. Development](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref55) *119*, 1175–1186.

[Takahashi, K., Kobayashi, T., and Kanayama, N. \(2000\). p57\(Kip2\) regulates](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref56) [the proper development of labyrinthine and spongiotrophoblasts. Mol. Hum.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref56) Reprod. *6*[, 1019–1025](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref56).

[Tanaka, M., Puchyr, M., Gertsenstein, M., Harpal, K., Jaenisch, R., Rossant, J.,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref57) [and Nagy, A. \(1999\). Parental origin-specific expression of Mash2 is estab](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref57)[lished at the time of implantation with its imprinting mechanism highly resistant](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref57) [to genome-wide demethylation. Mech. Dev.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref57) *87*, 129–142.

[Tao, G., Levay, A.K., Gridley, T., and Lincoln, J. \(2011\). Mmp15 is a direct](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref58) [target of Snai1 during endothelial to mesenchymal transformation and endo](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref58)[cardial cushion development. Dev. Biol.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref58) *359*, 209–221.

[Trapnell, C., Pachter, L., and Salzberg, S.L. \(2009\). TopHat: discovering splice](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref59) [junctions with RNA-Seq. Bioinformatics](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref59) *25*, 1105–1111.

[Tunster, S.J., Jensen, A.B., and John, R.M. \(2013\). Imprinted genes in mouse](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref60) [placental development and the regulation of fetal energy stores. Reproduction](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref60) *145*[, R117–R137.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref60)

[Zhang, J., Bonasio, R., Strino, F., Kluger, Y., Holloway, J.K., Modzelewski,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref61) [A.J., Cohen, P.E., and Reinberg, D. \(2013\). SFMBT1 functions with LSD1 to](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref61) [regulate expression of canonical histone genes and chromatin-related factors.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref61) [Genes Dev.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref61) *27*, 749–766.

Zhu, D., Hö[lz, S., Metzger, E., Pavlovic, M., Jandausch, A., Jilg, C., Galgoczy,](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref62) [P., Herz, C., Moser, M., Metzger, D., et al. \(2014\). Lysine-specific demethylase](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref62) [1 regulates differentiation onset and migration of trophoblast stem cells. Nat.](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref62) [Comms](http://refhub.elsevier.com/S1534-5807(15)00836-9/sref62) *5*, 3174.

Developmental Cell, Volume ³⁶

Supplemental Information

Maternal DNA Methylation Regulates

Early Trophoblast Development

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Figure S1 (related to Figure 1) – Gene expression alterations in Dnmt3 KO EPCs. A) The amount of extracellular space at the TGC layer was measured from histological sections (Fig. 1C) and normalised to the total area, showing that *Dnmt3a* mKO trophoblast sections show significantly increased in spacing between cells (**p<0.01, t-test). B) RT-qPCR analysis in oocyte and post-zygotic *Dnmt3* deletion E7.5 EPCs. Post-zygotic deletion was generated by crossing mice heterozygous for both *Dnmt3* genes; genotypes under each bar refer to *Dnmt3a* and *Dnmt3b*, respectively (e.g., Het;KO signifies *Dnmt3a^{+/-};Dnmt3b^{-/-})*. P-values refer to the effect of genotype as analysed by ANOVA. C) Expression differences to Ctrl EPCs across all mDKO DE genes. Log2 fold changes in each mDKO genotype relative to Ctrl EPCs are displayed as a heatmap. D) Immunofluorescence for CDX2 (green) in two sections from E7.5 wildtype or *Dnmt3a* mKO conceptuses. Sections were counterstained with DAPI (blue). Yellow staining is from autofluorescence of blood cells. In the wildtype, expression of CDX2 is visible both in the extraembryonic ectoderm (white arrowhead), where the TSC niche lies, and in the EPC (yellow arrow). In *Dnmt3a* mKO expression of CDX2 is specifically lost in the EPC, remaining high in the extraembryonic ectoderm. Error bars represent standard deviations.

Figure S2 (related to Figure 2) – Histological analysis of Ascl2 mKO E9.5 trophoblast. A) Hematoxylin and eosin staining of paraffin sections shows that *Ascl2* mKO trophoblast lacks the labyrinthine layer that is otherwise seen developing in WT trophoblast (marked by an asterisk); the TGC layer is expanded but, unlike *Dnmt3a* mKOs, does not have a pronounced reduction in tissue density when compared to WT TGCs; ch – chorion, epc – ectoplacental cone. B) The amount of extracellular space at the TGC layer was measured and normalised to the total area, showing that *Ascl2* mKO sections do not have increased spacing between cells. *p<0.05, **p<0.01 - ANOVA followed by post-hoc tests comparing *Dnmt3a* mKO with other genotypes. Error bars represent standard deviations.

Figure S3 (related to Figure 3) – RT-qPCR analysis of *Dnmt3* mDKO DE genes during TSC differentiation. TSCs were differentiated *in vitro* for 7 days. Genes with increased expression during *in vivo* differentiation (Fig. 3A) also show a differentiation-dependent profile *in vitro*.

Figure S4 (related to Figure 4) – Additional RT-qPCR analysis of TKO TSCs. TKO TSCs show no significant differences in the expression of key TSC markers (*Cdx2*, *Elf5*, *Esrrb*, *Eomes*) or the EMT markers *Cdh1* and *Snai1*. Error bars represent standard deviations.

Figure S5 (related to Figure 5) - Hypomethylation in DHet embryos is not preferentially derived from reprogramming-resistant loci. Blastocyst methylation data at CGIs (Kobayashi et al. 2012) was plotted against methylation levels in Ctrl EPCs (left) or the difference between DHet and Ctrl EPCs (right). CGIs with more than 50% methylation in oocytes are highlighted in blue and show that nearly all CGIs methylated in blastocyst are carried over from the oocyte. However, EPCs display de novo methylated CGIs as well. Hypomethylation in DHet EPCs occurs in both reprogramming-resistant and de novo methylated CGIs.

Figure S6 (related to Figure 6) - Scml2 regulation and role in early development. A) Scml2 methylation (left; amplicon CpGs 8-10) and expression (right) show asymmetry between E7.5 EPC and epiblast, and between TSCs and ESCs. B) Measurement of TSC adhesion to uncoated tissue culture wells shows that Scml2 overexpression is insufficient to drive adhesion defects. C) Additional RT-qPCRs in Scml2-overexpressing TSCs, grown under stem cell (FGF+) or differentiation (FGF-) conditions. D) Syna expression in WT and Ascl2 mKO E7.5 EPCs. E) CRISPR/Cas9 strategy for knocking out ScmI2 in TKO TSCs and corresponding genotyping PCRs. F) DNA methylation in human oocytes and morulae (Guo et al. 2014) at the syntenic region to the Scml2associated DMR; no methylation is seen in morulae. G) Comparison of CpG methylation between the whole genome ('All'), at regions of synteny with all mDKO DMRs ('DMR') and at DMR regions extended up to 400bp to include adjacent genomic regions ('Ext'). Overall, mDKO DMRs are hypermethylated in human oocytes and morulae; extending the DMRs dampens the difference, demonstrating local specificity of the hypermethylation. Bar plot error bars represent standard deviations.

SUPPLEMENTARY TABLES

Table S1 (related to Figure 1) – Differentialy expressed genes in all mDKO

genotypes versus Ctrl EPCs. Log2 fold-change and p values refer to the comparison of DHet with Ctrl EPCs. P values are corrected for multiple comparisons. Genes that are also differentially expressed in Ascl2 mKO EPCs are indicated.

Table S2 (related to Figure 2) – Gene ontology terms associated with Dnmt3 specific DE genes. The top 20 significant GO terms are shown. Benjamini-Hochberg correction of p values was applied.

Table S3 (related to Figure 2) – Gene ontology terms associated with Ascl2-

specific DE genes. The top 20 significant GO terms are shown. Benjamini-Hochberg correction of p values was applied.

SUPPLEMENTARY EXPERIMENTAL MATERIALS

Primers

