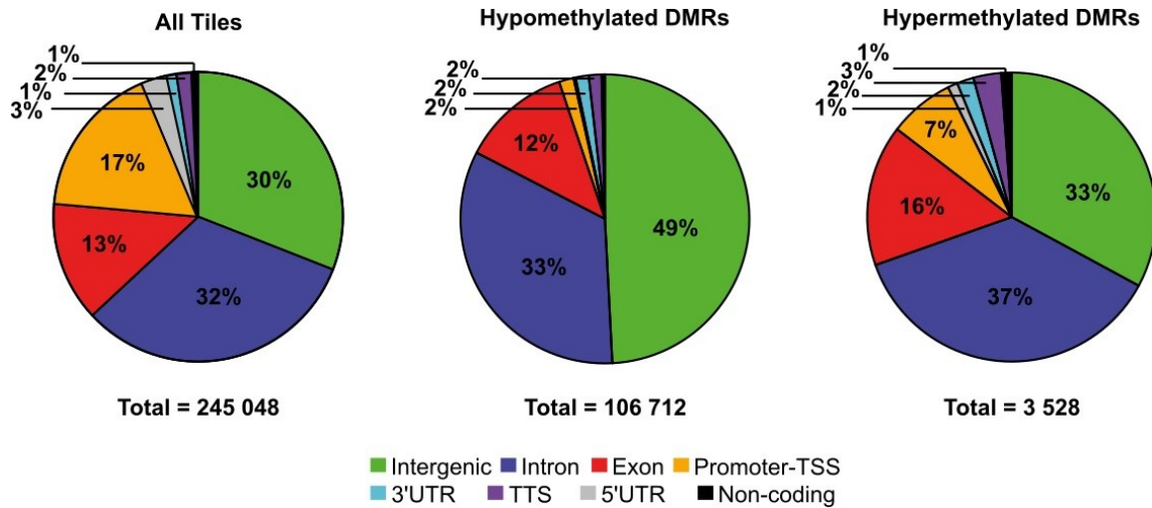
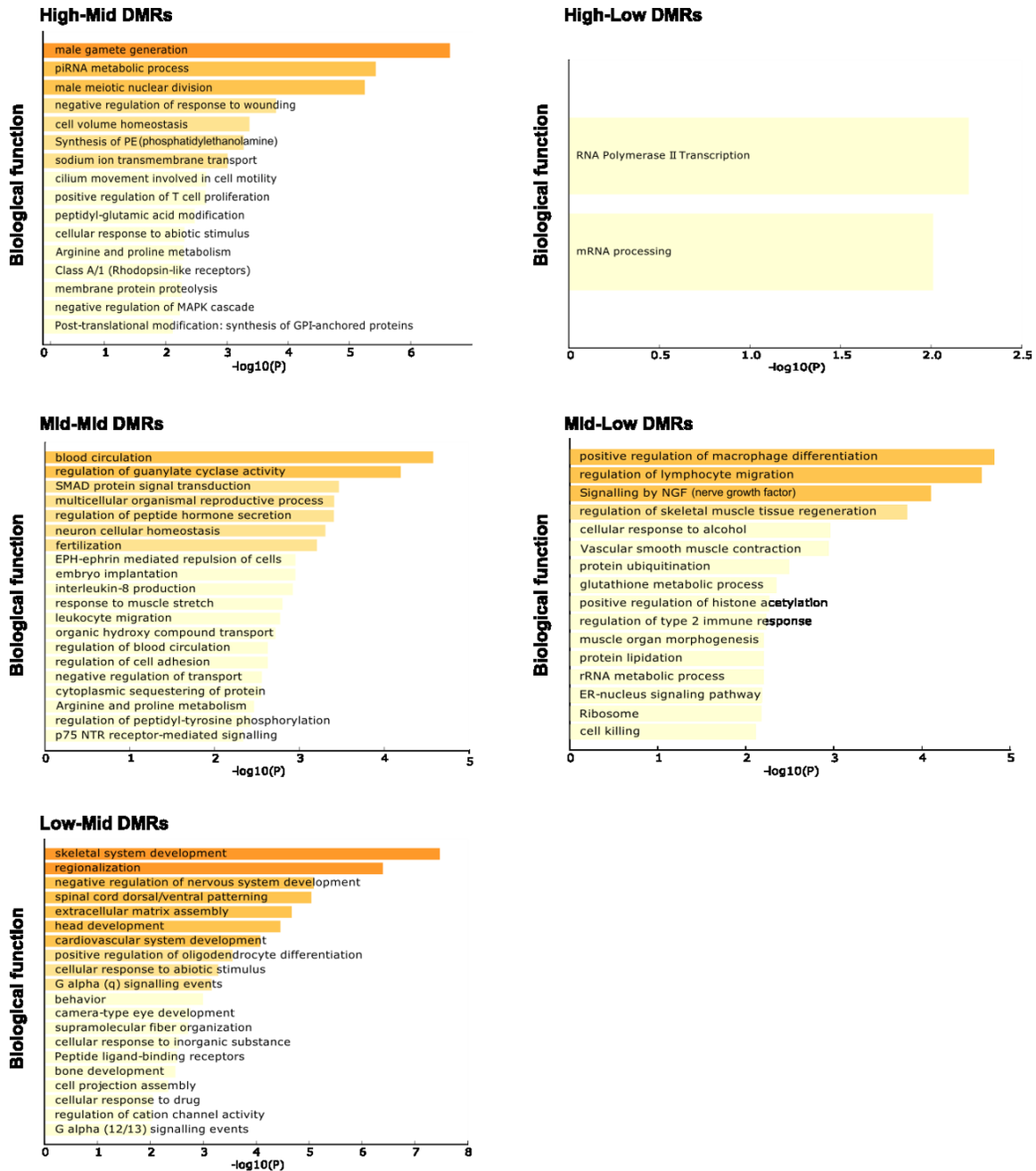


**Figure S1. Hierarchical dendrogram showing clustering of E10.5 embryo and placenta samples according to genome-wide DNA methylation profiles. Samples *a*, *c*: females; Samples *b*, *d*: males.**

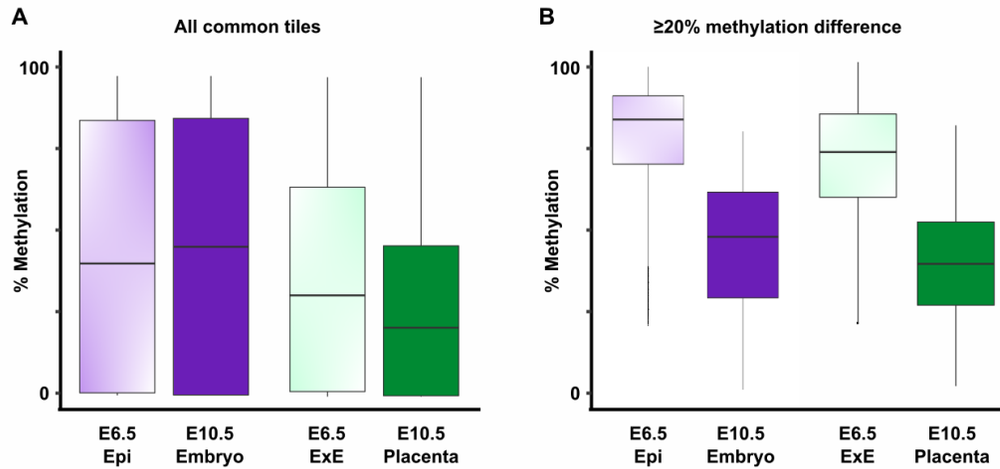


**Figure S2. Pie charts representing genomic annotation of Hypo- and Hyper-DMRs found between E10.5 embryo and placenta.** Proportion of all sequenced tiles (All-tiles; n=245 048) common to embryo and placenta, Hypo-DMRs (n=106 712) and Hyper-DMRs (n=3 528), found in intergenic and genic regions (exons, introns, promoters-TSS, 3' and 5' untranslated regions, and transcription termination sites, non-coding).

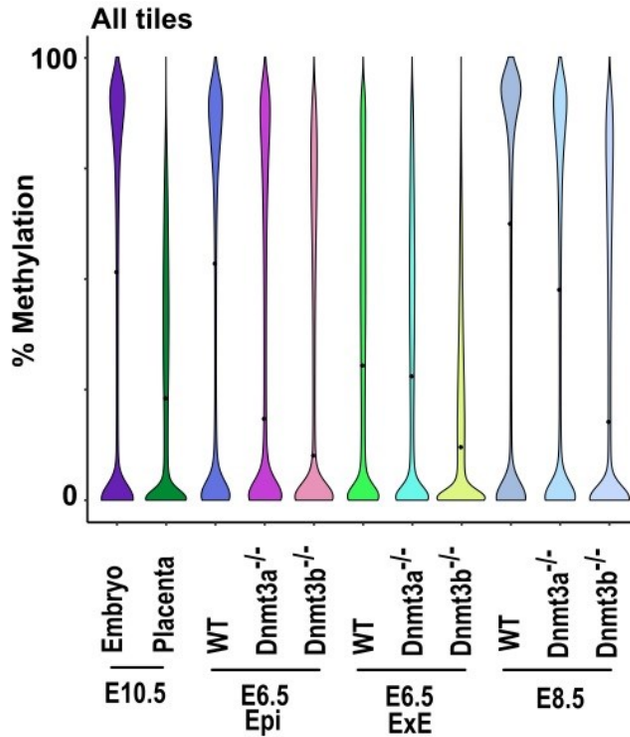


**Figure S3. Summary of biological functions associated with E10.5 embryo-placenta DMRs subtypes.** Number of tiles in promoter-TSS regions used as input for each methylation category: High-Low: 38; High-Mid: 749; Mid-Lo: 477; Mid-Mid: 509 and Low-Mid: 235. No enrichment in biological functions found for Mid-High: 1.

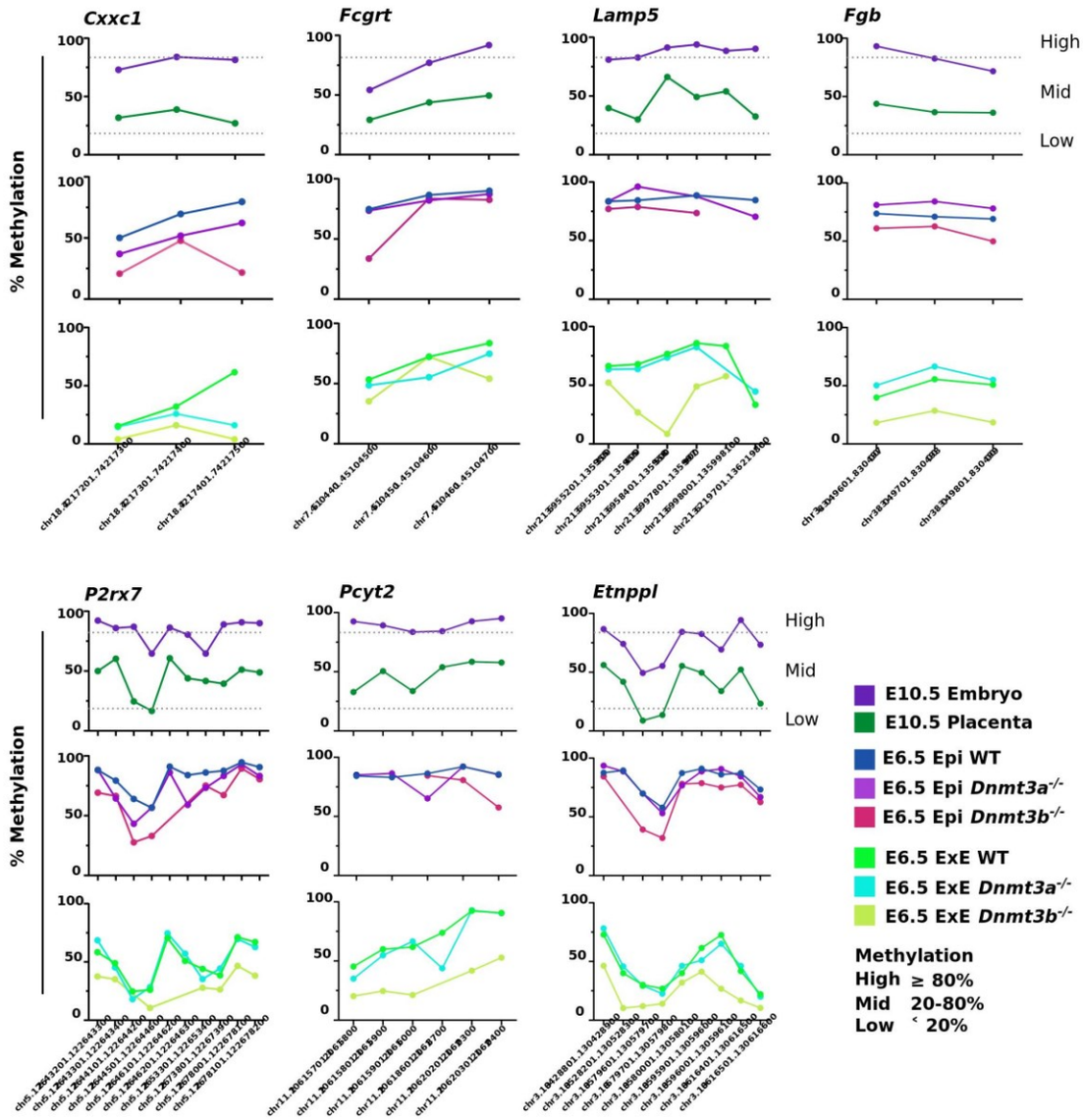




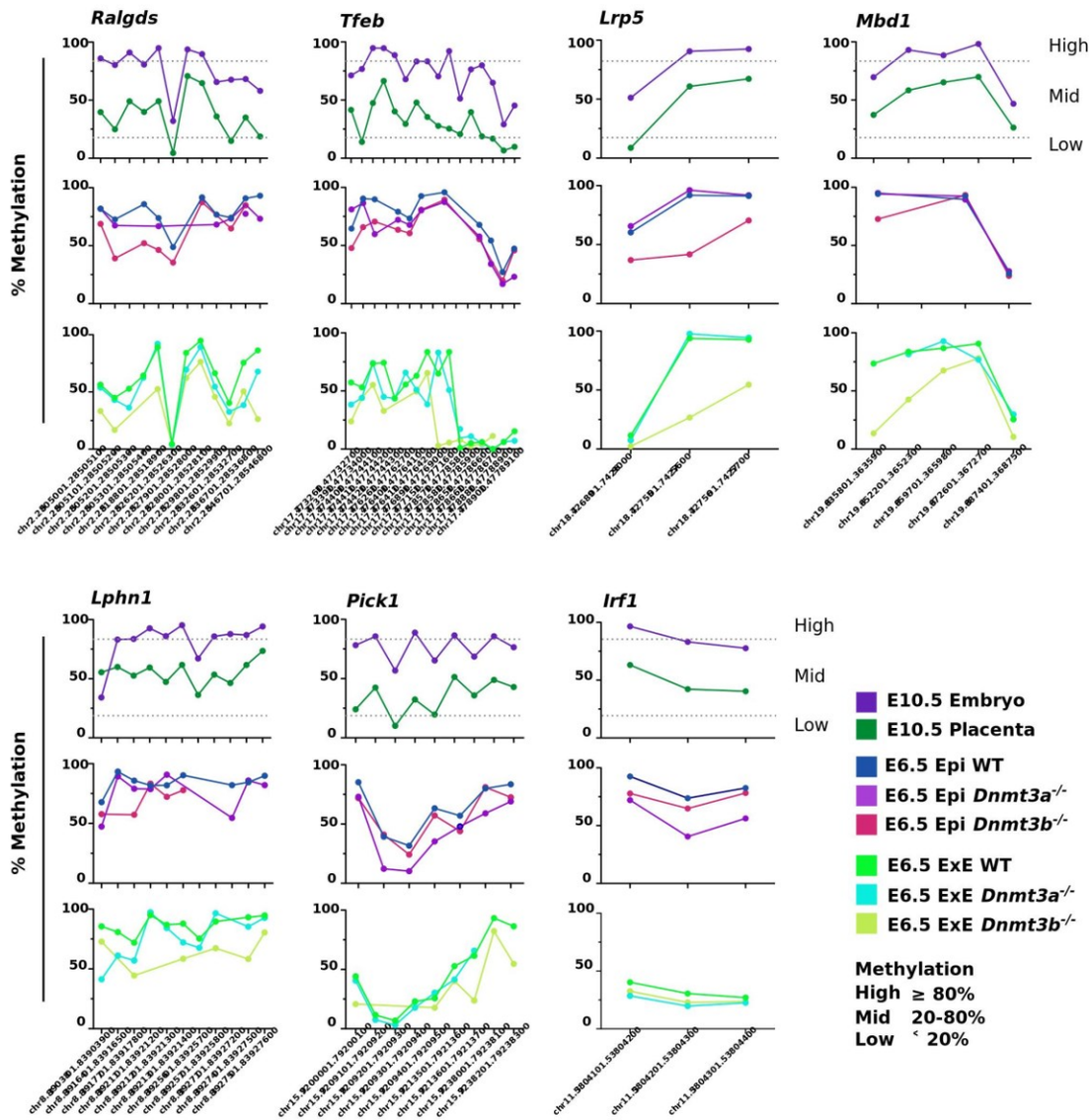
**Figure S5. DNA methylation profiles at E6.5 and E10.5.** **A)** Box plots representing the DNA methylation distribution and median values of overlapping common tiles between E6.5 ExE (Smith et al. 2014) and E10.5 placenta (All-tiles), as well as E6.5 epiblast (Smith et al. 2014) and E10.5 embryo (All-tiles). E6.5 ExE vs E10.5 placenta; 35.66% vs 25.79%,  $p < 0.0001$ ;  $n = 188\,098$  tiles, E6.5 epi vs E10.5 embryo; 43.2% vs 43.65%;  $p < 0.0001$ ;  $n = 200\,581$  tiles. **B)** Box plots representing the DNA methylation distribution and median values of overlapping common tiles between E6.5 ExE (Smith et al. 2014) and E10.5 placenta (All-tiles) (70.22% vs 37.49%;  $p < 0.0001$ ;  $n = 45\,967$ ), as well as E6.5 epiblast (Smith et al. 2014) and E10.5 embryo (All-tiles) (77.80% vs 43.87%;  $p < 0.0001$ ;  $n = 11\,183$ ), having more than 20% higher DNA methylation at E6.5.



**Figure S6. Global DNA methylation profiles in *Dnmt3a*<sup>-/-</sup> and *Dnmt3b*<sup>-/-</sup> samples.** Violin plots showing DNA methylation distribution and median values of *Dnmt3a* or *Dnmt3b* knockout mice in E6.5 epiblast (Epi), E6.5 extraembryonic ectoderm (ExE) (Smith et al. 2017) and E8.5 embryos (Auclair et al. 2014) in 100bp tiles corresponding to the 245 048 analyzed tiles (All-tiles). Overlapping tiles (minimum 15x coverage) in each dataset: E6.5 Epi WT n=227 657 ; E6.5 Epi *Dnmt3a*<sup>-/-</sup> n=160 283 ; E6.5 Epi *Dnmt3b*<sup>-/-</sup> n=145 590 ; E6.5 ExE WT n=227 926 ; E6.5 ExE *Dnmt3a*<sup>-/-</sup> n=200 801 ; E6.5 ExE *Dnmt3b*<sup>-/-</sup> n=170 164 ; E8.5 embryo WT n=209 466 ; E8.5 embryo *Dnmt3a*<sup>-/-</sup> n=209 466 ; E8.5 embryo *Dnmt3b*<sup>-/-</sup> n=207 974. See Table S3 for median and mean methylation values.



**Figure S7. *Dnmt3a*- or *Dnmt3b*-deficiency alter proper establishment of DNA methylation associated with various DMR categories within gene promoter-TSS.** DNA methylation average (%) per tile in *Dnmt3a*<sup>-/-</sup> and *Dnmt3b*<sup>-/-</sup> E6.5 epiblast and extraembryonic tissues (Smith et al. 2017) for E10.5 embryo-placenta associated DMRs. *Lrp5* (activation of *Wnt* signaling, important role in development processes); *Fgb* (encode for beta component of fibrinogen); *P2rx7* (ATP receptor); *Pcyt2* (role in the biosynthesis of phospholipid phosphatidylethanolamine); *Etnpl* (role in the biosynthesis of glycerophospholipid and metabolism); *Ralgds* (role in signaling processes).



**Figure S8. *Dnmt3a*- or *Dnmt3b*-deficiency alter proper establishment of DNA methylation associated with various DMR categories within genic regions.** DNA methylation average (%) per tile in *Dnmt3a*<sup>-/-</sup> and *Dnmt3b*<sup>-/-</sup> E6.5 epiblast and extraembryonic tissues (Smith et al. 2017) for E10.5 embryo-placenta associated DMRs. *Cxxc1* (role in regulation of gene expression and development); *Fcgrt1* (transfers IgG from mother to fetus across the placenta); *Lamp5* (role in synaptic plasticity in some GABAergic neurons); *Mbd1* (transcriptional repressor); *Lphn1* (role in cell adhesion and signal transduction); *Pick1* (role in synaptic plasticity and regulation of astrocyte morphology); *Tfeb* (role in the regulation of lysosomal genes and autophagy); *Irf1* (regulation of cellular response, including *IFN* and *IFN*-inducible-genes).