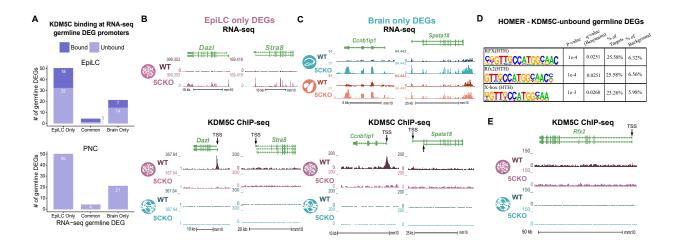
- <sup>1</sup> Supplement Erosion of somatic tissue identity with loss of the
- 2 X-linked intellectual disability factor KDM5C

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Supplementary Figure 1: KDM5C's catalytic activity promotes long-term silencing of germline genes via DNA methylation. A. Left - Bigwigs of representative histone 3 lysine 4 trimethylation (H3K4me3) ChIP-seq peaks at two germline genes in the wild-type (WT) and Kdm5c-KO (5CKO) adult amygdala. Right - Average H3K4me3 at the transcripiton start site (TSS) of all germline-enriched genes in wild-type (dark red) and Kdm5c-KO (light red) amygdala. B. Left - Bigwigs of representative histone 3 lysine 4 dimethylation (H3K4me2) ChIP-seq peaks at representative germline genes in wild-type and Kdm5c-KO EpiLCs. Right - Average H3K4me2 at the TSS of all germline-enriched genes in wild-type (dark purple) and Kdm5c-KO (light purple) EpiLCs. C. Diagram of embryonic stem cell (ESC) to epiblast-like cell (EpiLC) differentiation protocol and collection time points for RNA and protein. D. Real time quantitative PCR (RT-qPCR) of Kdm5c RNA expression, calculated in comparision to TBP expression ( $2^{-deltaCT}$ ). \* p < 0.05, \*\* p < 0.01, \*\*\* p < 0.001, Welch's t-test. E. KDM5C protein expression normalized to DAXX. Quantified intensity using ImageJ (artificial units - au). Right representative lanes of Western blot for KDM5C and DAXX. \* p < 0.05, \*\* p < 0.01, \*\*\* p < 0.001, Welch's t-test F. XXX G. XXX.