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Single Cell Methylomes Identify Neuronal Subtypes and Regulatory Elements in Mammalian Cortex

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Abstract

The mammalian brain contains diverse neuronal types, yet we lack single-cell epigenomic assays able to identify and characterize them. DNA methylation is a stable epigenetic mark that distinguishes cell types and marks regulatory elements. We generated >6,000 methylomes from single neuronal nuclei and used them to identify 16 mouse and 21 human neuronal subpopulations in the frontal cortex. CG and non-CG methylation exhibited cell type-specific distributions and we identified regulatory elements with differential methylation across neuron types. Methylation signatures identified a layer 6 excitatory neuron subtype and a unique human parvalbumin-expressing inhibitory neuron subtype. We observed stronger cross-species conservation of regulatory elements in inhibitory compared with excitatory neurons. Single-nucleus methylomes

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Supplementary Materials:

Materials and Methods

Supplementary texts

Figures S1–S17

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expand the atlas of brain cell types and identify regulatory elements that drive conserved brain cell diversity.

Mammalian neuron types are identified by their structure, electrophysiology and connectivity (1). The difficulty of scaling traditional cellular and molecular assays to whole neuronal populations has prevented comprehensive analysis of brain cell types. Sequencing mRNA transcripts from single cells or nuclei has identified cell types with unique transcriptional profiles in the mouse (2, 3) and human brain (4). However, these methods are restricted to RNA signatures, which are influenced by the environment. Epigenomic marks, such as DNA methylation (mC), are cell-type specific and developmentally regulated, yet stable across individuals and over the lifespan (5–7). We theorized that epigenomic profiles utilizing single-cell DNA methylomes could enable the identification of neuron subtypes in the mammalian brain.

During postnatal synaptogenesis, neurons accumulate substantial DNA methylation at non-CG sites (mCH) and reconfigure patterns of CG methylation (mCG, (7)). Patterns of mCG and mCH at gene bodies, promoters and enhancers are specific to neuronal types (5–8). Gene body mCH is more predictive of gene expression than mCG or chromatin accessibility (5). Because mCH is modulated over large domains, single neuron methylomes with sparse coverage can be used to accurately estimate mCH levels for over 90% of the genome by using coarse-grained bins (100kb) (Fig. S1). Whereas single cell RNA-Seq mainly informs about highly expressed transcripts, single neuron methylome sequencing assays any gene or non-gene region long enough to have sufficient coverage.

We developed a protocol for single nucleus methylcytosine sequencing (snmC-seq) and applied it to neurons from young adult mouse (8 week) and human (25 year) frontal cortex (FC) (Fig. 1A, (9)). snmC-seq provides a high rate of read mapping compared with published protocols (10–12) and allows multiplex reactions for large-scale cell type classification (Fig. S2,(9)). Like other bisulfite sequencing techniques (13), snmC-seq measures the sum of 5-methyl- and 5-hydroxymethylcytosines. Single neuronal nuclei labeled with anti-NeuN antibody were isolated by fluorescence-activated cell sorting (FACS) from human FC and from dissected superficial, middle and deep layers of mouse FC. We generated methylomes from 3,377 mouse and 2,784 human neurons with an average of 1.4 (1.8) million stringently filtered reads per cell in mouse (human), covering 4.7% (5.7%) of the genome (Fig. 1B–C, Table S1–2).

We calculated the mCH level for each neuron in nonoverlapping 100kb bins across the genome followed by dimensionality reduction and visualization using t-Stochastic Neighbor Embedding (t-SNE, (14)). The 2-dimensional tSNE representation was largely invariant over a wide range of experimental and analysis parameters (Fig. S3). A substantially similar tSNE representation was obtained using CG methylation levels in 100 kb bins, suggesting snmC-seq could be effective for cell-type classification of non-brain tissues without high levels of mCH (Fig. S3F).

The mammalian cortex arises from a conserved developmental program that adds excitatory neuron classes in an inside-out fashion, progressing from deep layers (L5,L6) to middle (L4)

and superficial layers (L2/3) (1). Inhibitory interneurons arise from distinct progenitors in the ganglionic eminences and migrate transversely to their cortical locations (15). We used mCH patterns to identify a conservative and unbiased clustering of nuclei for each species (9). Cluster robustness was validated by shuffling, down-sampling and comparison to density-based clustering (Fig. S3–4, (9, 16)). In addition, clustering was not significantly associated with experimental factors (e.g. batches, $q > 0.1$, chi-squared test, Fig. S5).

We applied identical clustering parameters to mouse and human cortical neuron mCH data and identified 16 mouse and 21 human neuron clusters (Fig. 2A–D). Assuming an inverse relationship between gene body mCH (average mCH across annotated genic region) and gene expression (7), we annotated each cluster on the basis of depletion of mCH at known cortical glutamatergic or GABAergic neuron markers (e.g. *Satb2*, *Gad1*, *Slc6a1*), cortical layer markers (e.g. *Cux2*, *Rorb*, *Deptor*, *Tle4*), or inhibitory neuron subtype markers (e.g. *Pvalb*, *Lhx6*, *Adarb2*) (1, 15, 17) (Fig. 2E–F, Fig. S6–7). For most clusters, mCH depletion at multiple marker genes (Fig. S6–7) allowed us to assign cluster labels indicating the putative cell type. For example, we found a cluster of mouse neurons with ultra-low mCH at *Rorb* (Fig. 2E, S6), a known marker of L4 and L5a excitatory pyramidal cells (17). Combining this information with markers such as *Deptor* (Fig. S6), which marks L5 but not L4 neurons, we labeled the cluster by species and layer, e.g. mL4 for mouse L4. Similarly, we used classical markers for inhibitory neurons such as *Pvalb* to label corresponding clusters, e.g. mPv for putative mouse *Pvalb*+ fast-spiking interneurons (15). We confirmed the accuracy of these classifications by comparing to layer dissected cortical neurons (Fig. S8A–B) and co-clustering with high-coverage methylC-seq data from purified populations of PV+ and VIP+ (5), as well as SST+ inhibitory neurons (Fig. S8C). Aggregated single neuron methylomes showed consistent mCH and mCG profiles compared to bulk methylomes of matching cell populations (Fig. S8D–E, S15F). Neuronal cluster classification for each of the major cell subtypes in mouse and human cortex based on single nuclei methylomes (Fig. 2G, S9A–B) was in good agreement with annotations based on single cell RNA-seq (2–4). Gene body mCH was anti-correlated with expression levels for corresponding clusters (Fig. S9C–E), validating our mCH marker-gene based annotation.

We found a greater diversity of excitatory neurons in deep layers compared to superficial layers for both mouse and human (Fig. 2). In both species, we identified one neuronal cluster for cortical L2/3 (mL2/3, hL2/3) and L4 (mL4, hL4), whereas L5 and L6 contained seven clusters in mouse (mL5, mL6 and mDL, where DL denotes deep layer neurons) and ten clusters in human (hL5, hL6 and hDL). Mouse L5 excitatory clusters (mL5-1, mL5-2) were hypomethylated at *Deptor* and *Bcl6*, which mark cortical L5a and L5b, respectively (Fig. S6, (18)). L6 excitatory clusters included subtypes with low mCH at the L6 excitatory neuron marker *Tle4* (mL6-1, mL6-2; hL6-1, hL6-2, hL6-3, (1)). Interestingly, several deep layer neuron clusters (mDL-2, hDL-1, hDL-2, hDL-3) were not hypomethylated at *Tle4*. We identified marker genes for each neuron type based on cell type-specific mCH depletion (Table S3) (9). While many marker genes were either classically established (1, 17) or recently identified neuron-type markers (Fig. S10A–B, (2–4)), we identified a significant number with no prior association to neuronal cell types (Fig. S10D–E, Table S3). mCH signature genes were hypomethylated in homologous clusters in mouse and human, with a

few notable exceptions. For example, the mouse L5a marker *Deptor* showed no specificity for human L5 neurons (Fig. S6–7, S10A–B).

Most clusters associated with classical cell type markers, but the identity of some clusters such as mDL-2 was less clear. We found that mDL-2 shares 24 marker genes with mL6-2, while 93 marker genes distinguish these clusters (Table S3). To validate the distinction between the two cell types, we selected a shared marker (*Sulfl*) and one unique to mL6-2 (*Tle4*) and performed double *in situ* RNA hybridization experiments in mouse FC (Fig. S11). The result confirmed the mCH-based prediction of a substantial proportion of L6 neurons expressing *Sulfl* but not *Tle4*, which likely correspond to mDL-2 (Fig. S11A–D). The proportion of L6 neurons expressing both *Sulfl* and *Tle4* likely represents a subset of mL6-2 (Fig. S11A–D). *Tle4*-expressing neurons in somatosensory cortex project to thalamus while *Sulfl* is expressed by both cortico-thalamic and cortico-cortical projecting neurons (18), suggesting neurons in cluster mDL-2 may have different projection targets compared to clusters showing hypomethylation of *Tle4* (e.g. mL6-2). We also observed extensive overlap of *in situ* hybridization signals using probes for a classical inhibitory neuron marker gene, *Pvalb*, and *Adgra3*, a predicted mCH signature of PV inhibitory neurons (Fig. S11E–G), further validating the specificity of marker prediction using mCH.

We paired homologous mouse and human neuron clusters using the correlation of mCH levels at homologous genes and found expanded neuronal diversity in the human FC compared with mouse (Fig. 2H, S12A, (19)). Multiple human neuron clusters showed homology to mouse L5a excitatory neurons (mL5-1), L6a pyramidal neuron (mL6-2), or to VIP, PV, and SST inhibitory neurons (Fig. 2H). We found unique gene-specific mCH pattern and super-enhancer-like mCG signatures in a potential human specific inhibitory population (hPv-2, Fig. S12B, S16H (9)).

Though we detected substantial mCH in all human and mouse neurons, cell types varied over a wide range in terms of their genome-wide mCH level (1.3–3.4% in mouse, 2.8–6.6% in human) (Fig. S13A–F). The sequence context of mCH was similar across all neuron types and consistent with previous reports (Fig. S13I,J, (5, 7)). Interestingly, global and gene specific mCH differences were found in PV and SST inhibitory neurons located in different cortical layers (Fig. S14, (9)). Genes with low mCH in superficial layer PV+ neurons are enriched in functional annotations including neurogenesis, axon guidance functions and synapse part (Fig. S14F–H, (9)), suggesting layer-specific epigenetic regulation of synaptic functions in inhibitory neurons.

A key advantage of single cell methylome analysis is the ability to obtain regulatory information from the vast majority of the genome (>97%, (19)) not directly assessed by RNA-Seq. By pooling reads from all neurons in each cluster, we could find statistically significant differentially methylated regions with low mCG in specific neuronal populations (CG-DMRs), which are reliable markers for regulatory elements (5). We found 575,524 mouse (498,432 human) CG-DMRs with average size of 263.6 bp (282.8 bp), covering 5.8% (5.0%) of the genome (Fig. 3A and S15A, Table S5–6). Most CG-DMRs (73.2% in mouse, 68.6% in human) are located >10 kb from the nearest annotated transcription start site (Fig. S15B–E). mPv and mVip CG-DMRs showed strongest overlap with ATAC-seq peaks and

putative enhancers identified from purified PV⁺ and VIP⁺ populations, respectively (Fig. S15G–H,(9, 20)). Hierarchical clustering of mCG levels at CG-DMRs grouped neuron-types by cortical layer and inhibitory neuron subtypes (Fig.S15I–J). Thus, neuron-type classification is supported by the epigenomic state of regulatory sequences.

We inferred transcription factors (TFs) that play roles in neuron-type specification by identifying enriched TF-binding DNA sequence motifs in CG-DMRs (Fig. 3B–C and Fig. S15K). We identified known transcriptional regulators and observed that several TF-binding motifs were enriched in human but depleted in mouse CG-DMRs in homologous clusters (Fig. 3C). The binding motif of Nuclear Factor 1 (NF1) was enriched in CG-DMRs for two human inhibitory neuron subtypes (hVip-2, hNdnf) but was depleted in homologous mouse clusters (mVip, mNdnf-2), suggesting a specific involvement of NF1 in human inhibitory neuron specification. Thus, although the TF regulatory circuits governing tissue types are conserved between mouse and human (21), fine-grained distinctions between neuronal cell types may be shaped by species-specific TF activity.

Super-enhancers are clusters of regulatory elements, marked by large domains of mediator binding and/or the enhancer histone mark H3K27ac, that control genes with cell-type specific roles (22). Extended regions of depleted mCG (large CG-DMRs) are also reliable markers of super-enhancers (Fig. S16A–C, (9, 23)). Therefore, we used our neuron type specific methylomes to predict super-enhancers for each mouse and human neuron type (Fig. 16D–I, Table S7–8). For example, super-enhancer activity was indicated by a large CG-DMRs at *Bcl11b* (*Ctip2*) in a subset of deep layer neurons (Fig. S16F–G), and supported by broad H3K27ac enrichment in mouse excitatory neurons (Fig. S16F). Super-enhancers overlap with key regulatory genes in the associated cell type, such as *Prox1* in VIP⁺ and NDNF⁺ neurons (Fig. S16H–I).

Global mCH and mCG levels were correlated between homologous clusters across mouse and human (Pearson $r = 0.698$ for mCH, and $r = 0.803$ for mCG, $p < 0.005$), suggesting evolutionary conservation of cell type specific regulation of mC (Fig. 4A, S13G–H). Examining 12,157 orthologous gene pairs, we found stronger correlation of gene body mCH between homologous clusters in mouse and human (median Spearman $r = 0.236$; Fig. 4B,C) than between different cell types within the same species ($r = -0.050$, mouse; $r = -0.068$, human). We found shared and species-specific CG-DMRs for homologous clusters based on sequence conservation (liftover; Fig. S17A–B). Cross-species correlation of mCG at CG-DMRs was significantly greater for inhibitory compared to excitatory neurons ($p < 0.001$ Wilcoxon rank sum test, Fig. 4D and S17C,(9)). Greater sequence conservation at inhibitory neuron CG-DMRs could partly explain the greater regulatory conservation (Wilcoxon rank sum test, $p < 0.001$, Fig. 4E). Sequence conservation was only observed within 1kb of the center of inhibitory neuron CG-DMRs, but did not extend to the flanking regions (Fig. S17G). These results support conservation of neuron type-specific DNA methylation, with greater conservation of inhibitory compared with excitatory neuron regulatory elements.

Single cell methylomes contain rich information enabling high-throughput neuron type classification, marker gene prediction and identification of regulatory elements. Applying a uniform experimental and computational pipeline to mouse and human allowed unbiased

comparison of neuronal epigenomic diversity in the two species. The expanded neuronal diversity in human, revealed by DNA methylation patterns, is consistent with more complex human neurogenesis, such as the presence of outer radial glia and the potential dorsal origin of certain interneuron subtypes (15, 24, 25). Further immunological, physiological and functional experiments are needed to characterize the DNA methylation based neuronal populations defined by our study. Single neuron epigenomic profiling allowed the identification of regulatory elements with neuron type specific activity outside of protein-coding regions of the genome. We expect that the single nucleus methylome approach can be applied to studies of disease, drug exposure or cognitive experience, to examine the role of cell type-specific epigenomic alteration in neurological/neuropsychiatric disorders.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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One Sentence Summary

Single-nucleus cytosine DNA methylomes distinguish neuron types and predict conserved gene regulatory elements in mouse and human cortex.

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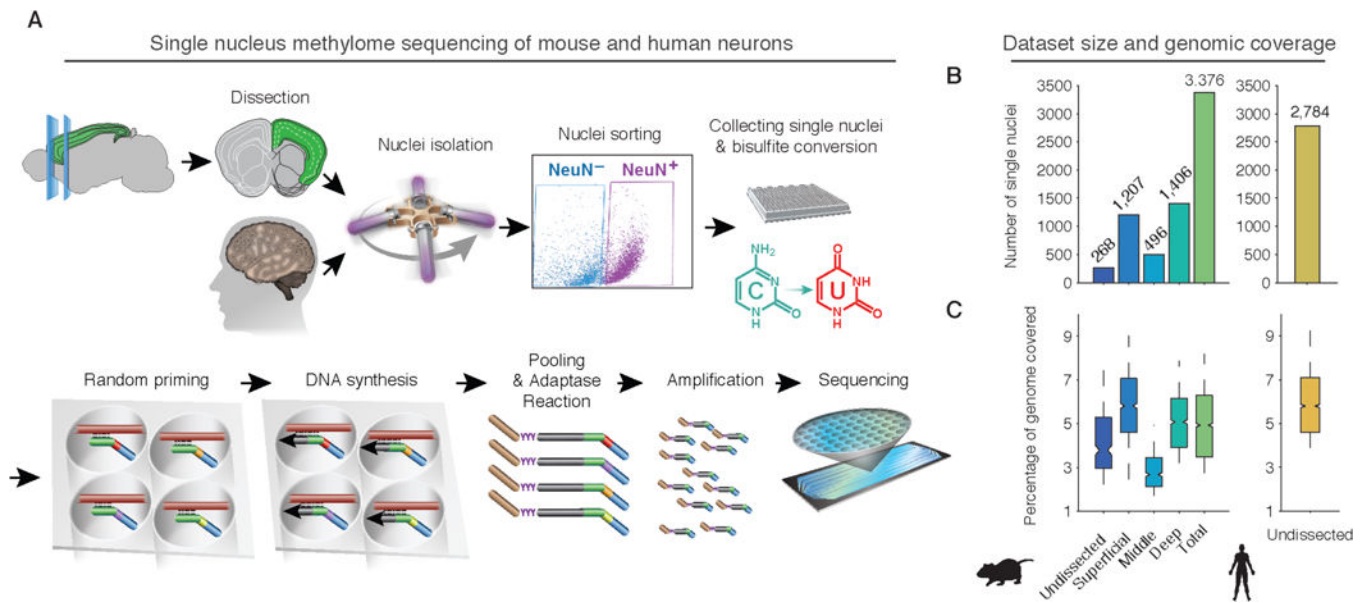


Fig. 1. High-throughput single nucleus methylome sequencing (snmC-seq) of mouse and human frontal cortex (FC) neurons
 (A) Workflow of snmC-seq. (B,C) Number of single neuron methylomes (B) and distribution of genomic coverage per dataset (C).

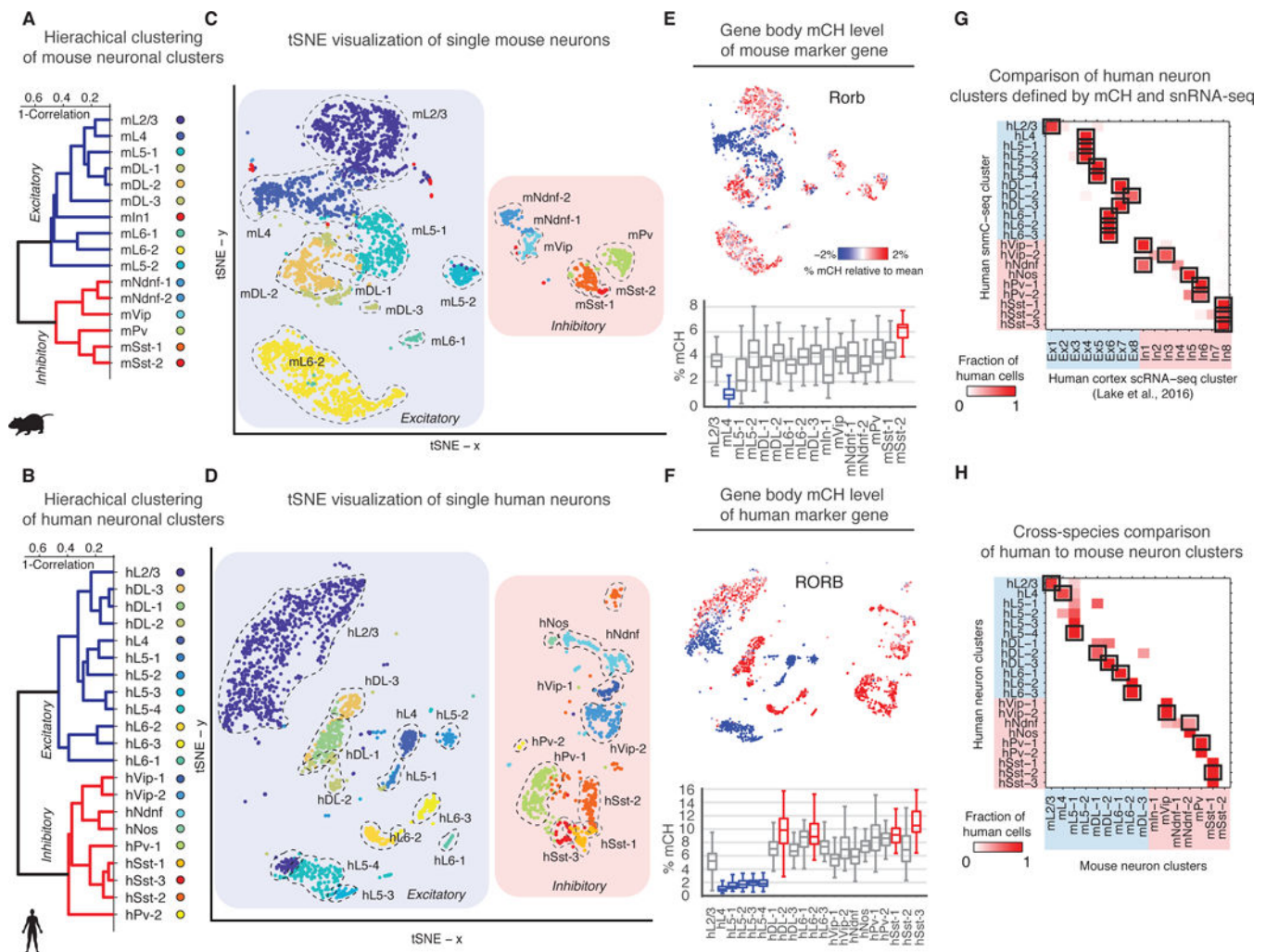


Fig. 2. Non-CG methylation (mCH) signatures identify distinct neuron populations in mouse and human FC

(A,B) Hierarchical clustering of neuron types using gene body mCH level. (C,D) Two-dimensional visualization of single neuron clusters (tSNE (9)). Mouse and human homologous clusters are labeled with similar colors. (E,F) Gene body mCH at *Rorb* for each single neuron (top), and the distribution for each cluster (bottom) with hyper/hypo-methylated clusters highlighted in red/blue. (G) Comparison of human neuron clusters defined by mCH with clusters from single nucleus RNA-Seq (4, 9). (H) Fraction of cells in each human cluster assigned to each mouse cluster based on mCH correlation at orthologous genes (9). Mutual best matches are highlighted with black rectangles.

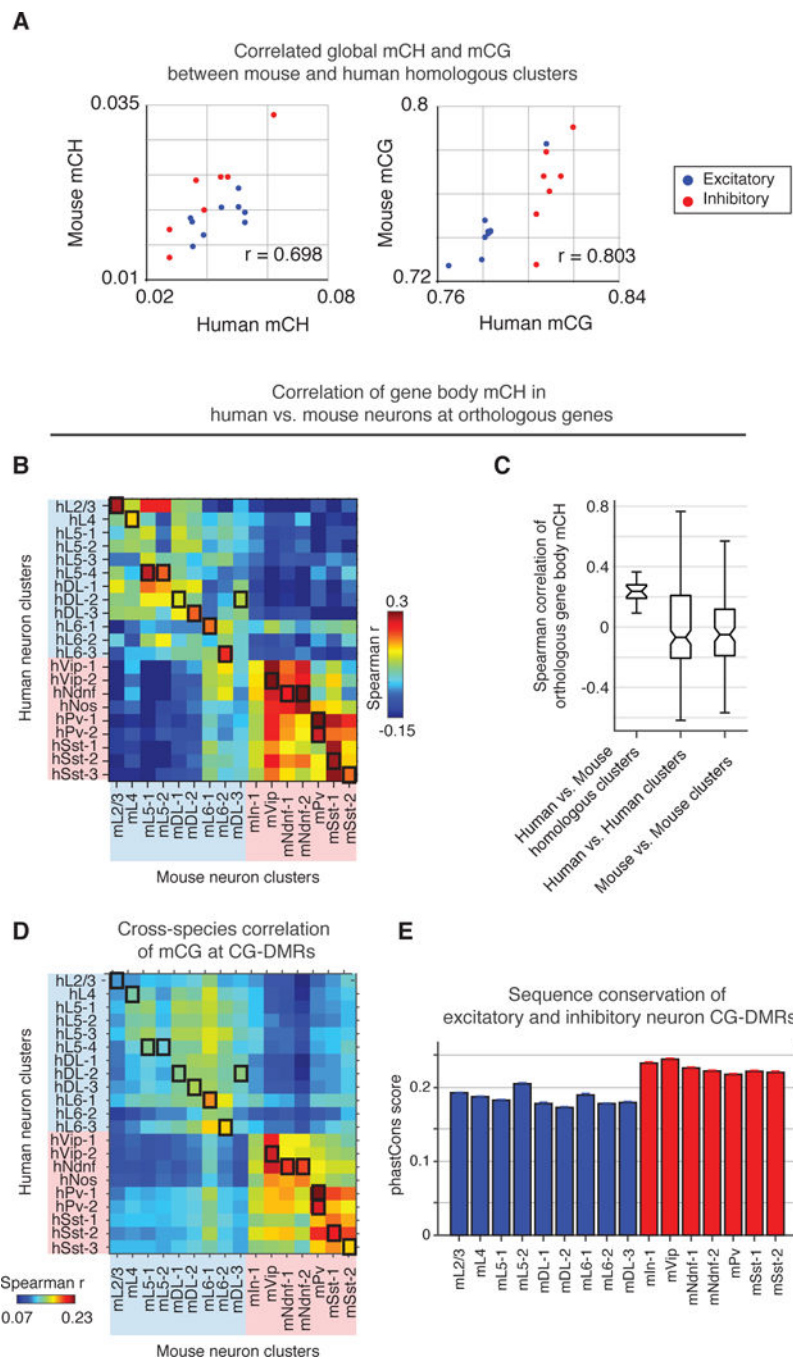


Fig. 4. Conserved gene body mCH and CG-DMRs between mouse and human

(A) Global mCH and mCG levels are strongly conserved within homologous cell types between mouse and human. (B) Cross-species correlation of gene body mCH at orthologous genes shows cell type-specific conservation. Black boxes denote homologous neuron clusters. (C) The median correlation of gene body mCH for homologous clusters is higher than the within-species correlation for distinct clusters. (D) Cross-species correlation of

mCG at neuron-type-specific CG-DMRs. (E) Sequence conservation at neuron-type-specific DMRs.

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