Neuron

Chd2 Is Necessary for Neural Circuit Development and Long-Term Memory

Highlights

- . CHD2 is widely expressed in the embryonic and mature brain
- Gene expression is broadly altered by Chd2^{+/-}
- Chd2^{+/-} mice have deficits in neuron proliferation, synaptic function and memory
- Interneuron transplantation rescues memory problems in Chd2^{+/-} mice

Authors

Young J. Kim, Sattar Khoshkhoo, Jan C. Frankowski, ..., Sunyoung Lee, Ye Emily Wu, Robert F. Hunt

Correspondence

yjkim13@uci.edu (Y.J.K.), robert.hunt@uci.edu (R.F.H.)

In Brief

Kim et al. show that loss-of-function mutation of the chromatin regulator gene Chd2 produces deficits in neuron proliferation, synaptic function, and memory in mice.





Cell²ress



Chd2 Is Necessary for Neural Circuit Development and Long-Term Memory

Young J. Kim,^{1,*} Sattar Khoshkhoo,^{1,2} Jan C. Frankowski,¹ Bingyao Zhu,¹ Saad Abbasi,¹ Sunyoung Lee,^{1,3} Ye Emily Wu,⁴ and Robert F. Hunt^{1,5,6,*}

¹Department of Anatomy & Neurobiology, University of California, Irvine, Irvine, CA 92697, USA

²Department of Neurology, Brigham and Women's Hospital and Massachusetts General Hospital, Harvard University, Boston, MA 02115. USA

³Department of Integrative Biology and Physiology, University of California, Los Angeles, Los Angeles, CA 90095, USA

⁴Department of Biological Chemistry and Department of Neurobiology, David Geffen School of Medicine, University of California, Los Angeles, Los Angeles, CA 90095, USA

⁵Twitter: @hunt_lab ⁶Lead Contact

*Correspondence: yjkim13@uci.edu (Y.J.K.), robert.hunt@uci.edu (R.F.H.)

https://doi.org/10.1016/j.neuron.2018.09.049

SUMMARY

Considerable evidence suggests loss-of-function mutations in the chromatin remodeler CHD2 contribute to a broad spectrum of human neurodevelopmental disorders. However, it is unknown how CHD2 mutations lead to impaired brain function. Here we report mice with heterozygous mutations in Chd2 exhibit deficits in neuron proliferation and a shift in neuronal excitability that included divergent changes in excitatory and inhibitory synaptic function. Further in vivo experiments show that Chd2+/mice displayed aberrant cortical rhythmogenesis and severe deficits in long-term memory, consistent with phenotypes observed in humans. We identified broad, age-dependent transcriptional changes in Chd2^{+/-} mice, including alterations in neurogenesis, synaptic transmission, and disease-related genes. Deficits in interneuron density and memory caused by Chd2+/- were reproduced by Chd2 mutation restricted to a subset of inhibitory neurons and corrected by interneuron transplantation. Our results provide initial insight into how Chd2 haploinsufficiency leads to aberrant cortical network function and impaired memory.

INTRODUCTION

Exome sequencing studies have now identified hundreds of gene mutations carrying an increased risk for neurodevelopmental disorders (Allen et al., 2013; De Rubeis et al., 2014; Iossifov et al., 2014; Deciphering Developmental Disorders Study, 2015). Of these, mutations in genes encoding chromatin regulators have emerged as a common risk factor, with *CHD2*, a member of the chromodomain helicase DNA-binding (CHD) family of

proteins, being frequently affected (Carvill et al., 2013; Carvill et al., 2015). CHD2 is an ATP-dependent chromatin-remodeling factor with poorly understood function in the developing or adult brain. In human, CHD2 haploinsufficiency is associated with intellectual disability, a variety of catastrophic childhood epilepsies, autism spectrum disorder, and photosensitivity (Neale et al., 2012; Rauch et al., 2012; Chénier et al., 2014; Galizia et al., 2015), with large phenotypic variability among affected individuals. In mice, heterozygous deletion of Chd2 results in histological abnormalities of heart, muscle, lung, liver, kidney, spleen, and bone (Marfella et al., 2006, 2008; Harada et al., 2012; Kulkarni et al., 2008). Initial studies in brain indicate that Chd2 knockdown during the peak of embryonic neurogenesis promotes production of neurons from neural progenitors, possibly depleting the precursor pool (Shen et al., 2015). However, unlike other CHD family members in which recent progress using animal models has led to important mechanistic insights about behavioral phenotypes and cellular pathways (Durak et al., 2016; Katayama et al., 2016; Gompers et al., 2017; Platt et al., 2017), there is essentially nothing known about brain defects that arise as a consequence of Chd2 haploinsufficiency.

Understanding how mutations in chromatin remodeling genes impact brain function may reveal new opportunities for targeted therapies. This led us to generate a *Chd2* mutant mouse line and investigate the effect of *Chd2* haploinsufficiency on the developing and mature mouse brain. Our findings support the hypothesis that *Chd2* has a critical function in forebrain neurogenesis *in vivo* and the generation of GABAergic interneurons in particular. We also found evidence of a functional role for *Chd2* in cortical circuit physiology and long-term spatial memory.

RESULTS

CHD2 Is Expressed in Neurons and Oligodendrocytes

We first determined the expression profile of CHD2 in wild-type (WT) C57BL/6J mice at postnatal day 30 (P30) (n = 6 mice)



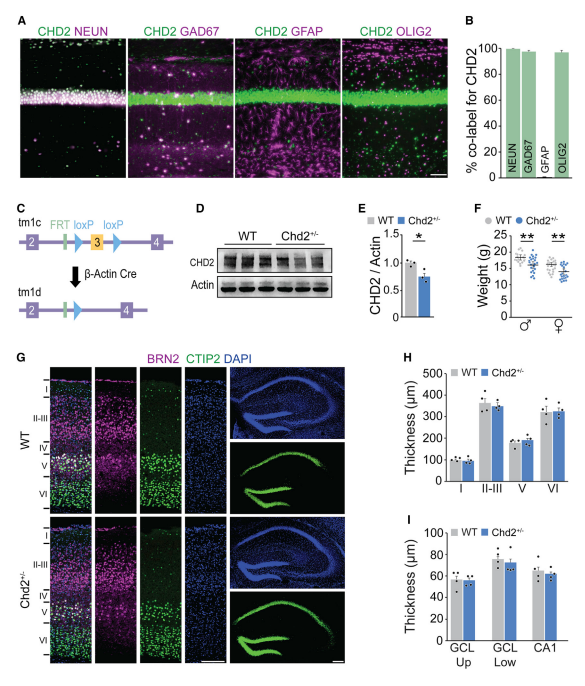


Figure 1. Generation of Chd2+/- Mice

- (A) At P30, CHD2 (green) co-labeled with NEUN, GAD67, and OLIG2, but not GFAP (all in magenta).
- (B) Quantification of CHD2 expression in each cell type (n = 3 mice per marker).
- (C) Schematic of the conditional allele for Chd2 (tm1c). Cre deletes the floxed exon 3 of Chd2 to generate a frameshift mutation (tm1d).
- (D) Western blot analysis for CHD2 and Actin protein.
- (E) At P30, western blot analysis showed a reduction of CHD2 protein in brain of Chd2+/- (p = 0.027, n = 3 mice per genotype).
- (F) Mean body weight of male and female Chd2*/- mice was reduced compared to WT littermates (n = 17-23 mice per genotype).
- (G) Immunostaining for BRN2 (magenta), CTIP2 (green), and DAPI (blue) in somatosensory cortex (left) and hippocampus (right) at P30.
- (H) Thickness of individual cell layers in somatosensory cortex (SS Ctx) was not altered by $Chd2^{+/-}$ (n = 4 mice per genotype).
- (I) Width of granule cell layer (GCL) or CA1 pyramidal cell layer (CA1) was not altered by $Chd2^{+/-}$ (n = 4 mice per genotype).

Error bars, SEM; *p < 0.05; **p < 0.01; scale bars, 75 μ m in (A) and 150 μ m in (G). See also Figure S1.

(Figure 1). Immunofluorescence experiments revealed CHD2 was widely expressed throughout the young adult mouse brain, with especially strong expression in olfactory bulb, neocortex, hippocampus, and cerebellum (Figures S1A and S1B). These observations are consistent with RNA expression patterns of Chd2 detected by in situ hybridization (Gene Paint; https://gp3.mpg.de/results/chd2). To determine whether CHD2 expression was limited to a specific brain cell type, we further evaluated coronal sections from hippocampus of P30 WT mice (n = 11 mice). We found that CHD2 co-localized in nearly all mature neurons (NEUN-positive, 99.8% ± 0.1%), GABAergic interneurons (GAD67-positive, 97.7% ± 0.39%), and oligodendrocytes (OLIG2-positive, 97.6% ± 0.95%). Conversely, CHD2 was not expressed in the majority of GFAP-positive astrocytes (0.4% ± 0.4%) (Figures 1A and 1B). CHD2 co-localized with DAPI, consistent with its role in chromatin regulation (Figure S1C).

Generation of Chd2+/- Mice

In humans, CHD2 mutations are typically loss of function (Carvill et al., 2015). To establish a mouse line with a heterozygous lossof-function mutation in Chd2 (i.e., Chd2+/- mice), we crossed transgenic mice containing loxP-flanked exon 3 of Chd2 (i.e., Chd2^{tm1c(EUCOMM)Hmgu} mice) with a β-actin Cre line (Figure 1C; Figures S1D-S1F). Mice with heterozygous deletion in Chd2 showed approximately half the expression of CHD2 protein in brain as compared to WT littermates (Figures 1D and 1E). Chd2+/- mice of both sexes were viable and fertile but had reduced body weight compared to WT littermates (Male: WT: $18.5 \pm 0.8 \text{ g}$, n = 17 mice, $Chd2^{+/-}$: $16.2 \pm 0.9 \text{ g}$, n = 23 mice; two-tailed t test; p = 0.001; Female: WT: 16.4 ± 0.4 g, n = 22mice, $Chd2^{+/-}$: 14.2 ± 0.7 g, n = 22 mice; two-tailed t test; p = 0.0004) (Figure 1F). $Chd2^{+/-}$ mice also exhibited mild lordokyphosis by 2 months of age (Figure S1H). This phenotype was not apparent in WT littermates and is consistent with reports of scoliosis in humans with Chd2 mutations (Chénier et al., 2014: Kulkarni et al., 2008).

We first asked whether a reduction in Chd2 disrupts cytoarchitecture of the cortex. Analysis of the layer-specific markers BRN2 (layer II/III), CTIP2 (layer V and VI), and DAPI revealed no gross alterations to lamination at P30 (Figures 1G-1I), and analysis of NEUN immunostaining showed no obvious laminar disorganization in somatosensory cortex or hippocampus (Figure S1I). No significant differences were detected in the thickness of somatosensory neocortex or dorsal hippocampus between genotypes (Figure 1H; Figure S1J). Likewise, we did not observe any differences in the width of the granule cell or CA1 pyramidal cell layers within hippocampus of Chd2+/mice compared to WT littermates (Figure 1I; Figure S1K). Finally, we evaluated WT and Chd2+/- mice for cellular phenotypes of epilepsy in dentate gyrus. Analysis of Timm's staining revealed no mossy fiber sprouting into the inner molecular layer of the dentate gyrus (Mann-Whitney rank-sum test, p = 1.00, U-statistic = 162.00) (Figure S1L). We also found no change in the intensity of GFAP immunostaining or density of GFAP+ cells between genotypes (Figures S1M-S1P). Hence, our results show Chd2 haploinsufficiency does not substantially disrupt cortical cytoarchitecture.

Reduced Numbers of GABAergic Interneurons in Chd2*/- Mice

Because CHD2 was co-expressed with nearly all GABAergic interneurons, we asked whether numbers of individual GABA cell populations were altered in Chd2+/- mice. At P30, we found ~20% decrease in the density of GAD67-expressing cells in somatosensory cortex (WT: 302.3 ± 12.2 GAD67+ cells/mm², $Chd2^{+/-}$: 239.8 ± 11.3 GAD67+ cells/mm²; n = 6 mice per genotype, p = 0.03) and hippocampus CA1 (WT: 175.6 ± 5.9 GAD67+ cells/mm², n = 5 mice; $Chd2^{+/-}$: 140.8 ± 2.8 GAD67+ cells/mm², n = 6 mice; p = 0.0003) (Figure 2). We also found \sim 10% decrease in the density of NEUN+/GAD67- cells (putative excitatory neurons) in somatosensory cortex (WT: 1,938 ± 45 cells/mm², $Chd2^{+/-}$: 1,791 ± 17 cells/mm²; n = 4 mice per genotype, p = 0.03) (Figure S2), suggesting that $Chd2^{+/-}$ does not selectively alter interneuron number, but no change was found in hippocampus CA1 (WT: 4,235 \pm 205 cells/mm², Chd2^{+/-}: 4,534 \pm 192 cells/mm²; n = 4 mice per genotype, p = 0.3). Immunostaining analysis for GABAergic subtypes revealed decreased density of cells expressing parvalbumin (PV; 15% decrease in somatosensory cortex and 22% in CA1), somatostatin (SST; 11% decrease in somatosensory cortex and 31% in CA1), and reelin (12% decrease in CA1) (Figure 2). The density of cells expressing calretinin (CR) or vasoactive intestinal peptide (VIP) was not significantly different between genotypes.

One hypothesis that could explain fewer GABA neurons in $Chd2^{+/-}$ mutants is that $Chd2^{+/-}$ may favor the production of oligodendrocytes over interneurons, which derive from the same progenitor regions of the ventral telencephalon (Petryniak et al., 2007). To test this, we quantified the number of OLIG2+ cells at P30. However, we found no difference in OLIG2+ cell densities in somatosensory cortex (WT: 659.2 \pm 33.5 cells/mm²; $Chd2^{+/-}$: 662.7 \pm 26.1 cells/mm²; $Chd2^{+/-}$: 513.8 \pm 25.0 cells/mm², $Chd2^{+/-}$: 513.8 \pm 25.0 cells/mm², $Chd2^{+/-}$: 513.8 \pm 25.0 cells/mm², $Chd2^{+/-}$ mice; $Chd2^{+/-}$ mice; $Chd2^{+/-}$ mice in oligodendrogenesis in $Chd2^{+/-}$ mice.

Chd2^{+/−} Disrupts Cell Proliferation in the Developing Forebrain

Interneuron density in the adult nervous system is largely determined by the size of the precursor pool in the embryo (Southwell et al., 2012). To investigate whether Chd2 has a role in GABAergic neurogenesis, we crossed Chd2+/mice with a GAD67-GFP knockin reporter labeling nearly all GABAergic interneurons (Tamamaki et al., 2003). CHD2 expression was observed throughout the embryonic brain of WT mice at embryonic day 14.5 (E14.5; n = 12 mice) (Figures 3A-3C). In dorsal telencephalon, and in agreement with a previous report (Shen et al., 2015), CHD2 was expressed at the highest levels in the cortical plate, where it co-localized with the majority of TBR1+ cells, as well as the ventricular zone (VZ) and subventricular zone (SVZ), where CHD2 co-localized with KI67+ mitotically active precursors and PAX6+ radial glia (Figure 3A; Figure S4). In contrast, CHD2 expression was low or absent in the intermediate zone (IZ) and marginal zone (MZ) at this stage and rarely overlapped with TBR2+ intermediate progenitors. In ventral telencephalon, CHD2 was expressed in the medial and

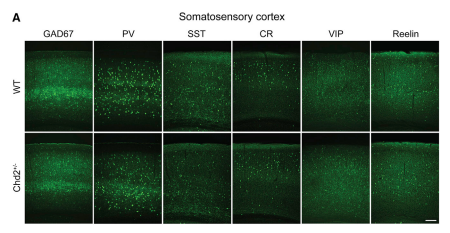
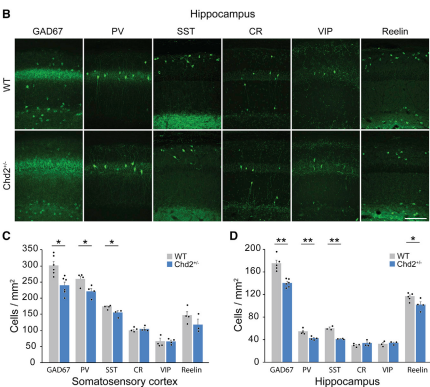


Figure 2. *Chd2*^{+/-} Mice Exhibit Decreased Density of GABAergic Interneurons

(A and B) Immunostaining of coronal sections through somatosensory neocortex (A) and CA1 region of hippocampus (B) for GAD67, PV, SST, CR, VIP, and reelin at P30.

(C and D) Quantification of each subtype marker shows $Chd2^{+/-}$ mice had decreased density of cells expressing GAD67, PV, and SST in somatosensory cortex (C) and decreased density of cells expressing GAD67, PV, SST, and reelin in hippocampus (D) compared to WT littermates (n = 3–6 mice per genotype).

Error bars, SEM; $^*p < 0.05$; $^{**}p < 0.01$; scale bars, 150 μ m in (A) and (B). See also Figures S2 and S3.



of KI67-expressing cells in VZ/SVZ of $Chd2^{+/-}$ mice (WT: 8,087.5 ± 934.8 cells/mm², n = 4 mice; $Chd2^{+/-}$: 5,150 ± 180.3 cells/mm²; n = 3 mice; p = 0.04) (Figures 3D and 3E). In MGE, we also observed an ~35% decrease in the density of KI67-expressing cells in VZ/SVZ of $Chd2^{+/-}$ mice (WT: 8,910.5 ± 279.5 cells/mm², n = 5 mice; $Chd2^{+/-}$: $7,108.3 \pm 361.2 \text{ cells/mm}^2$; n = 3 mice; p = 0.04). Further analysis at E14.5 revealed a subsequent decrease in NKX2.1-expressing progenitors in MGE (WT: $17,929.2 \pm 717.2 \text{ cells/mm}^2$, n = 6 mice; $Chd2^{+/-}$: 15,000 ± 365.5 cells/ mm^2 , n = 5 mice; p = 0.008) and reduced density of GAD67-GFP+ neurons in cortex of Chd2+/- mice, as compared to WT littermates (WT: 3,122.5 ± 162.9 cells/mm², n = 4 mice; $Chd2^{+/-}$: $2,450.5 \pm 39.9 \text{ cells/mm}^2$, n = 6 mice; p = 0.02) (Figures 3F-3I). Finally, we asked whether Chd2+/- altered caspase-3 mediated interneuron cell death, which peaks around P7 in mice (Southwell et al., 2012). At P7, we found a decrease in the density of GAD67-GFP+ cells in somatosensory cortex (WT: 781.1 ± 50.78

cells/mm²; $Chd2^{+/-}$: 643.3 ± 40.21 cells/mm²; n = 5 mice per genotype; p = 0.04), but the density of cells expressing caspase-3 was not different between genotypes (Figures 3J–3L). We conclude that Chd2 haploinsufficiency disrupts cell proliferation and neurogenesis in the developing forebrain, but not developmentally programmed cell death.

caudal ganglionic eminence (MGE and CGE), progenitor domains that generate nearly all cortical GABAergic interneurons (Figures 3B and 3C; Figure S4). At this stage, CHD2 strongly co-localized with Kl67 and NKX2.1 but only partially co-localized with GAD67-GFP+ cells in MGE. CHD2 was not detected in tangentially migrating interneurons in neocortex (Figure 3A). By demonstrating progenitor domain-specific expression of CHD2 during cortical development, our results suggest that CHD2 plays a role in cell proliferation, terminal differentiation, and maturation of cortical principal neurons and GABAergic interneurons.

To determine whether forebrain precursors were altered by $Chd2^{+/-}$, we performed a series of immunostaining studies at E14.5. In cortex, we found \sim 35% decrease in the density

Differential Gene Expression in Chd2+/- Mice

As a chromatin modifier, *Chd2* likely plays an important role in global transcriptional regulation, but its exact function in brain is unknown (Marfella and Imbalzano, 2007; Tyagi et al., 2016). Having established that CHD2 is expressed in both the embryonic forebrain and mature neurons, we next sought to identify transcriptional changes resulting from *Chd2* haploinsufficiency

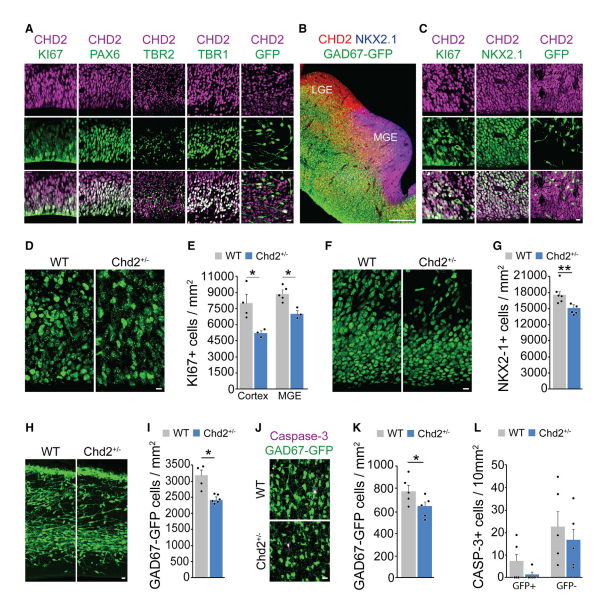


Figure 3. Chd2 Regulates Proliferation of Neural Progenitors in Developing Forebrain

- (A) Immunostaining for CHD2 (magenta) and KI67, PAX6, TBR2, TBR1, and GAD67-GFP (all green) in neocortex at E14.5.
- (B) Immunostaining for CHD2 (red), NKX2.1 (blue), and GAD67-GFP (green) in ventral telencephalon shows CHD2 expression in GABAergic progenitor domains at E14.5.
- (C) At E14.5, CHD2 (magenta) co-labeled with KI67 and NKX2.1, but not GAD67-GFP (all in green), in MGE. Lower-magnification images of dorsal and ventral telencephalon are shown in Figure S4.
- (D) Immunostaining for KI67 (green) in MGE of a WT and $Chd2^{+/-}$ mouse at E14.5.
- (E) At 14.5, the density of KI67⁺ cells was reduced in MGE and cortex of Chd2^{+/-} mice (n = 3–5 mice per genotype).
- (F) Immunostaining for NKX2.1 (green) in MGE of a WT and Chd2+/- mouse at E14.5.
- (G) At E14.5, the density of NKX2.1+ GABA progenitors was reduced in MGE of Chd2+/- mice (n = 5-6 mice per genotype).
- (H) Immunostaining for GAD67-GFP (green) in cortex of a WT and Chd2+/- mouse at E14.5
- (I) At E14.5, the density of GAD67-GFP progenitors was reduced in cortex of $Chd2^{+/-}$ mice (n = 4-6 mice per genotype).
- (J) Immunostaining for GAD67-GFP (green) and caspase-3 (magenta) in somatosensory neocortex of a WT and $Chd2^{+/-}$ mouse at P7.
- (K and L) At P7, immunostaining analysis revealed a reduction in GAD67-GFP progenitors (K), but the density of cells expressing caspase-3 (CASP-3) was not different between genotypes (n = 5 mice per genotype) (L).

Error bars, SEM; *p < 0.05; **p < 0.01; scale bars, 10 μ m in (A)–(C), 200 μ m in (B), and 20 μ m in (D), (F), (H), and (J). See also Figure S4.

in an unbiased way. To do this, we performed polyA+ RNA sequencing on tissue micro-dissected from neocortex and MGE of developing embryos at E13.5 as well as hippocampus

in adult mice at P45 (Figure 4A). We tested for differential expression (DE) across 24,062 genes expressed in our datasets in 9 WT mice and 7 $Chd2^{+/-}$ littermates. At significance cutoffs

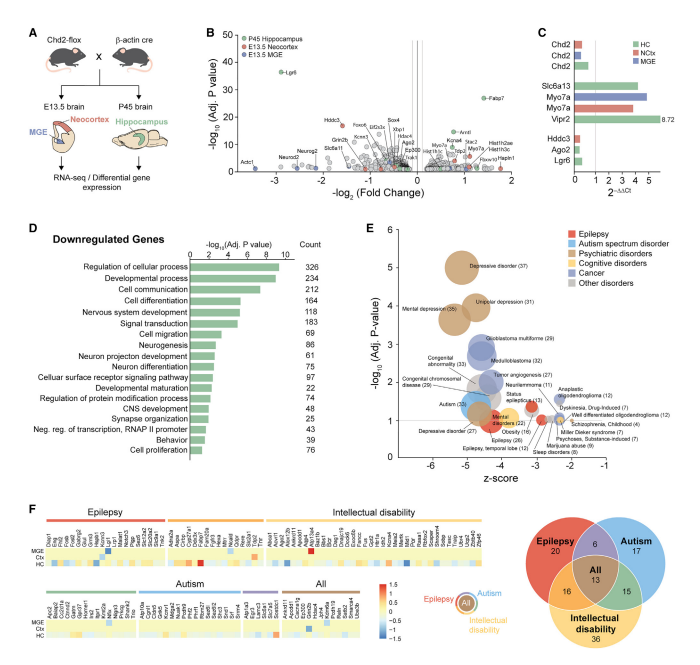


Figure 4. Chd2 Haploinsufficiency Disrupts Genes Necessary for Early Cortical Development and Synaptic Function

- (A) Schematic showing the experimental approach used for RNA sequencing.
- (B) Volcano plot displaying genes that were significantly differentially expressed (Adj. p < 0.1) in $Chd2^{+/-}$ mice (n = 3 mice per genotype).
- (C) qRT-PCR validation of DE genes predicted by RNA sequencing.
- (D) Gene ontology for downregulated genes at P45. Complete gene ontology is provided in Table S3.
- (E) Plot of Z-score by $-\log_{10}$ of the adjusted p value for human diseases identified by disease ontology. The complete disease ontology is provided in Table S5.
- (F) Heatmaps of genes associated with childhood epilepsy, intellectual disability, and autism spectrum disorder that were differentially expressed in E13.5 MGE, E13.5 neocortex, and/or P45 hippocampus.

corresponding to Benjamini and Hochberg adjusted p value < 0.1, we found 44 (E13.5 MGE), 14 (E13.5 neocortex), and 652 (P45 hippocampus) DE genes in $Chd2^{+/-}$ mice (Figure 4B; Table S2). In addition, we further validated expression changes of a randomly selected group of DE genes via qRT-PCR (Figure 4C). In all three groups, Chd2 was downregulated (E13.5

MGE: $2^{-\Delta\Delta Ct}$ = 0.32, p = 0.04; E13.5 neocortex: $2^{-\Delta\Delta Ct}$ = 0.36, p = 0.04; P45 hippocampus: $2^{-\Delta\Delta Ct}$ = 0.62, p = 0.04) (Figure 4C).

Analysis of Gene Ontology (GO) terms revealed brain regionspecific dysregulation in $Chd2^{+/-}$ mice compared to WT littermates. For this analysis, we expanded the list of DE genes to a significance cutoff of p value < 0.05 for E13.5 datasets, which

increased sample sizes to 1,416 (E13.5 neocortex) and 622 (E13.5 MGE) DE genes; adjusted p value < 0.1 (p < 0.0045) was used for P45 hippocampus samples. At E13.5, DE genes showed strong enrichment for annotations associated with nervous system development, neuron differentiation, and neurogenesis (Table S3). Similar enrichment was observed for DE genes in hippocampus at P45, with numerous additional annotations associated with synapse organization, transcriptional regulation, and behavior (Figure 4D; Table S3). In particular, we found robust differential expression of transcription factors related to forebrain neurogenesis (e.g., Neurod2, Neurog2, and Sox4), RNA silencers (e.g., Ago2), global regulators of the epigenome (e.g., Ep300 and Smarca4), cell adhesion molecules (e.g., Cdh4 and Pcdh17), and ion channels (e.g., Kcna4 and Cacnb1). However, most striking was the broad range of up- and downregulated genes related to neuronal activity and synaptic plasticity in P45 hippocampus (Table S4). Of note, the majority of differentially expressed synaptic molecules are involved in glutamatergic synaptic function (e.g., Grina, Shank2, and Homer1). In agreement with previous reports that chromatin remodeling proteins act as regulators of Wnt signaling pathways (Kwan et al., 2016), we found downregulation of numerous Wnt pathway genes in Chd2+/- mice, such as Wnt7a, Wnt7b, Fzd1, Lrp1, Ctnnd2, Apc2, and Apcdd1. However, we noted little overlap in the specific DE genes found in our data compared to prior studies on Chd8 mutation (Durak et al., 2016; Katavama et al., 2016; Gompers et al., 2017; Platt et al., 2017), suggesting that CHD subfamily members have distinct functional roles in brain.

A number of chromatin remodelers, including Chd2, have been implicated in human neurodevelopmental disorders (Marfella and Imbalzano, 2007; Carvill et al., 2015; Kwan et al., 2016; Tyaqi et al., 2016; Mastrototaro et al., 2017; Sokpor et al., 2017). Therefore, we next asked whether Chd2+/- altered genes implicated in human disease. Analysis of Disease Ontology (DO) terms identified DE genes for annotations associated with epilepsy and autism spectrum disorders as well as numerous other neurological diseases, such as mental health disorders, congenital disease, and cancer (Figure 4E; Table S5). In particular, we found 123 DE genes associated with intellectual disability (e.g., Cacna1g, Ctnnd2, and Hexa), childhood epilepsy (e.g., Hdac4, Pcdh19, Reln, and Slc6a1), and/or autism spectrum disorders (e.g., Grin2b, Ktm2a, and Shank2) (Figure 4F), some of which are strong candidate "risk" genes while others are relatively uncharacterized. Altogether, our results indicate that Chd2+/- leads to a broad dysregulation of genes involved in neurogenesis, synapse organization, and disease-related pathways.

Chd2+/- Disrupts Excitatory and Inhibitory Synaptic **Function in Hippocampus**

The observation that Chd2^{+/-} drastically alters genes involved in synapse organization led us to investigate whether there is a change in neuronal or synaptic function in Chd2+/- mice. To do this, we examined electrophysiological properties of CA1 pyramidal neurons at P30-P35 using patch-clamp recordings in acute hippocampal slices (Figure 5). Compared to WT animals, pyramidal neurons from Chd2+/- mice showed an increase in action potential firing at current injections exceeding 100 pA

(genotype: $F_{(1,25)} = 2.06$; p = 0.164), current step: $F_{(1,16)} =$ 279.22; p < 0.001; genotype by current step interaction: $F_{(1.16)} = 4.53$; p < 0.001; n = 13 cells from 3 WT mice, n = 14 cells from 4 Chd2+/- mice; two-way repeated-measures ANOVA) (Figures 5A and 5B). Consistent with greater firing capacity in CA1 pyramidal neurons, we also found a significant decrease in spike adaptation at 2x spike threshold in recorded neurons (WT: 1.61 \pm 0.13, $Chd2^{+/-}$: 1.3 \pm 0.07; p = 0.03; Mann Whitney U), but all other intrinsic electrophysiological properties examined were not significantly different between treatment groups (Figure 5C; Table S6).

Next, we examined miniature excitatory and inhibitory synaptic inputs onto CA1 pyramidal neurons in the presence of 1 μM tetrodotoxin (TTX). Compared to WT littermates, pyramidal neurons in Chd2+/- mice exhibited a significant increase in miniature excitatory postsynaptic current (mEPSC) amplitudes without a change in event frequency (n = 14 cells from 4 WT mice, n = 16 cells from 4 $Chd2^{+/-}$ mice; Figures 5D-5F). This was accompanied by significantly faster decay kinetics in Chd2+/mice; 10%-90% rise time was not different between genotypes. (Figures 5G-5I). In contrast, Chd2+/- mice exhibited a decrease in miniature inhibitory postsynaptic current (mIPSC) frequency without a change in mIPSC amplitude or event kinetics (n = 20 cells from 3 WT mice, n = 21 cells from 3 $Chd2^{+/-}$ mice; Figures 5J-50). Fewer mIPSCs onto CA1 pyramidal neurons could be explained by a loss of GABAergic synapses or a change in GABA release probability in Chd2+/- mice (Hirsch et al., 1999). Therefore, we evaluated evoked IPSC responses to paired stimuli applied to stratum radiatum, a commonly used measure of presynaptic short-term plasticity. However, we did not observe a change in paired-pulse ratio (PPR) of evoked IPSCs between genotypes, suggesting that presynaptic GABA release is unaltered in Chd2+/- mice (Figure S5). Taken together, our findings indicate that Chd2+/- alters neuronal excitability and has differential effects on glutamatergic versus GABAergic synaptic transmission in hippocampus.

Chd2+/- Mice Exhibit Changes in Cortical **Rhythmogenesis**

A change in the normal balance between excitation and inhibition has the capacity to disrupt naturally occurring oscillations in the brain. Thus, we implanted electroencephalogram (EEG) electrodes into somatosensory neocortex of freely behaving Chd2+/mice and WT littermates (n = 5 mice per genotype) and recorded intracranial EEG > 1 week later at P40-P65 (Figure 6). Analyses of the local field potential revealed clear episodes of each frequency band in WT mice and Chd2+/- littermates, indicating that these rhythms remain intact in the mutants (Figures 6A and 6B). We found that Chd2+/- mice exhibited a significant increase in the alpha (8-13 Hz) and gamma (30-70 Hz) frequency ranges compared to WT littermates (Figure 6C), suggesting that cortical synchrony is increased by Chd2 haploinsufficiency. Notably, elevations in resting alpha or gamma power have been observed in human epilepsies (Willoughby et al., 2003; Vaudano et al., 2017), autism spectrum disorders (Orekhova et al., 2007; Cornew et al., 2012), and mouse models featuring deficits in interneuron function (Korotkova et al., 2010; Carlén et al., 2012; Del Pino et al., 2013; Howard et al., 2014; Cho et al.,

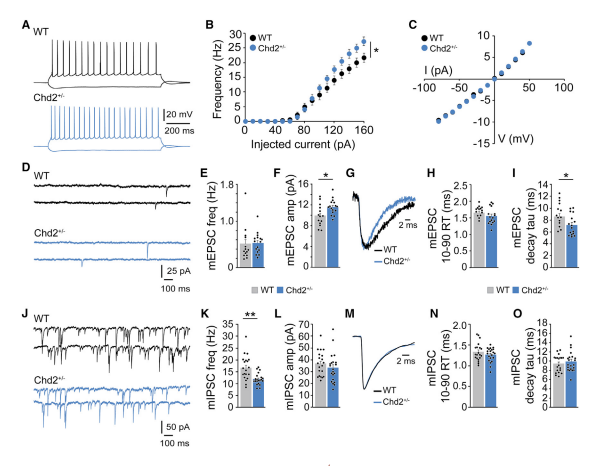


Figure 5. Changes in Neuronal Excitability and Synaptic Function in Chd2+/- Mice

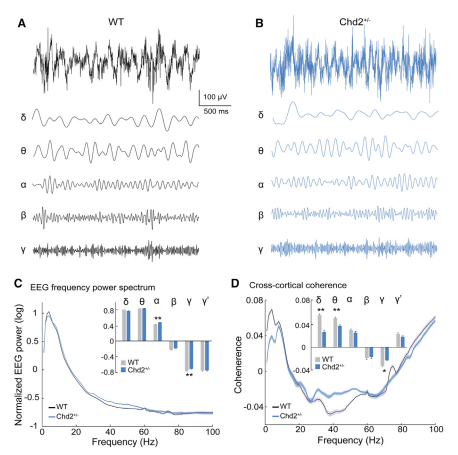
(A) Voltage responses to hyperpolarizing (-80 pA) and depolarizing (+140 pA) current pulses in CA1 pyramidal neurons from a WT (black) and Chd2^{+/-} mouse (blue).

- (B) Plot of action potential firing frequency (Hz) as a function of current intensity shows increased firing in pyramidal neurons of Chd2+/- mice.
- (C) Summary current-voltage plot shows R_{input} was not different between groups. Intrinsic electrophysiological properties are summarized in Table S6.
- (D) Miniature EPSCs (mEPSCs) recorded from pyramidal neurons in a WT (black) and Chd2+/- mouse (blue).
- (E and F) At P30-P35, mEPSC frequency was unchanged (E), but mEPSC amplitudes were increased (F) in pyramidal neurons of Chd2+/- mice.
- (G) Averaged mEPSCs recorded from a pyramidal neuron in a WT (black) and Chd2+/- mouse (blue).
- (H and I) At P30-P35, mEPSC 10%-90% rise time (RT) was unchanged (H), but mEPSC decay time constant was decreased (I) in pyramidal neurons of $Chd2^{+/-}$ mice.
- (J) Miniature IPSCs (mIPSCs) recorded from pyramidal neurons in a WT (black) and Chd2+/- mouse (blue).
- (K and L) At P30–P35, mIPSC frequency was decreased (K), but mIPSC amplitudes were unchanged (L) in pyramidal neurons of Chd2*/- mice.
- (M) Averaged mIPSCs recorded from a pyramidal neuron in a WT (black) and Chd2+/- mouse (blue).
- (N and O) At P30–P35, mIPSC 10%–90% rise time (RT) (N) and decay time constant (tau) (O) were not different between genotypes.
- Error bars, SEM; *p < 0.05; **p < 0.01. See also Figure S5.

2015). Examination of the mean normalized power spectra for all other frequency ranges revealed no differences between genotypes (Figure 6C). Further analysis revealed differences in cross-cortical coherence (a measure of phase and amplitude synchrony between hemispheres) in delta, theta, and gamma frequency ranges (Figure 6D). Next, we performed 24 hr/7 day EEG monitoring to test whether $Chd2^{+/-}$ mice develop spontaneous electrographic seizures. In 7 days of continuous monitoring at P40–P65, we did not observe any overt convulsive seizures in WT or $Chd2^{+/-}$ mice (n = 6 mice per genotype). Altogether, these findings indicate that Chd2 haploinsufficiency leads to a dysregulation of neural oscillations and synchrony in cortex.

Chd2+/- Disrupts Long-Term Memory

Memory requires changes in neuronal gene expression that are coordinated, at least in part, by epigenetic mechanisms and chromatin regulation (Levenson and Sweatt, 2005; Vogel-Ciernia et al., 2013; Mews et al., 2017). To assess long-term memory in $Chd2^{+/-}$ mice, we used the well-characterized object location and recognition memory paradigms (Vogel-Ciernia and Wood, 2014) (Figure 7; Figures S6A and S6B). During a 10 min habituation phase in the open field arena, the total distance traveled was not different between groups, but $Chd2^{+/-}$ mice spent more time in the center region of the arena, as compared to WT littermates (Figures S6C–S6E; Table S7). In the object location task, WT mice exhibited



increased exploration of the object that had been moved (Figures 7A-7C). In contrast, Chd2+/- mice explored both objects equally (Figures 7A-7C) and displayed a significantly lower discrimination index compared to WT controls (Figures 7B) and 7C). Similarly, in the object recognition task, WT mice exhibited increased exploration of the novel object (Figures 7D-7F), but Chd2+/- mice explored both objects equally (Figures 7D-7F) and displayed a significantly lower discrimination index compared to WT controls (Figures 7E and 7F). There was no difference in the time spent exploring the objects (Figures S6F and S6G), suggesting that the poor performance of Chd2+/- mice was not due to disinterest in the objects and that locomotor activity during training and test phases was comparable between groups (Figure S6E). Chd2-conditional mutant mice with Chd2 haploinsufficiency only in inhibitory interneurons (Nkx2.1-Cre; Chd2^{loxP/+}; Ai14-tdTomato mice) showed similar impairments in interneuron density and memory behaviors (Figure S7). We conclude that Chd2+/- mice exhibit severe deficits in long-term spatial and recognition memory, and this is due, at least in part, to a reduction in the number of cortical interneurons.

MGE Transplantation Rescues Spatial Memory Problems in *Chd2**/- Mice

In hippocampus, deficits in interneuron number or function have been implicated in a wide range of cognitive disorders, such as

Figure 6. Abnormal Rhythmogenesis in *Chd2*+/- Mice

(A and B) Example of each frequency band isolated from the local field potential (LFP) in a WT (A) and $Chd2^{+/-}$ (B) mouse.

(C) Normalized EEG power spectra. Inset shows mean power for each frequency band (n = 5 mice per genotype).

(D) Cross-cortical coherence across the EEG power spectra. Inset shows mean coherence for each frequency band (n = 5 mice per genotype). Error bars, SEM; *p < 0.05; **p < 0.01.

Alzheimer's disease (Satoh et al., 1991), intellectual disability (Opperman et al., 2017), and epilepsy (de Lanerolle et al., 1989). With this in mind, we next asked whether transplantation of inhibitory interneurons is sufficient to "rescue" memory deficits observed in $Chd2^{+/-}$ animals. To do this, MGE progenitors were obtained from E13.5 β-actin:GFP donor mice, and 5×10^4 cells were injected bilaterally into hippocampus of neonatal recipients at P5 (Figure 8A). We performed polyA+ RNA sequencing and DE analysis between MGE and neocortex micro-dissected from E13.5 embryos to confirm that the grafted cells expressed genes highly enriched in MGE-derived progenitors (Olig2, Dlx5, Lhx6, Pou3f4,

and Gad1) (Figure 8B; Table S8). By 45 days after transplant (DAT), MGE-GFP cells migrated away from the injection and were dispersed throughout hippocampal subfields (n = 3 mice per genotype) (Figure 8C: Figures S8A and S8B), GFP-labeled cells expressed the interneuron markers PV (WT: 18% ± 4%; $Chd2^{+/-}$: 24% ± 6%) and SST (WT: 35% ± 6%; $Chd2^{+/-}$: 28% \pm 2%) but did not express VIP (<1% both genotypes) (Figure 8D). No differences in marker expression were detected between genotypes (Figure S8C). Finally, we assessed the effect of MGE transplantation on long-term memory. Replication with a second, independent cohort of WT and Chd2+/- mice showed that Chd2+/- mutants had deficits in both object location and recognition memory tasks (Figures 8E and 8F). Chd2+/- mice that received MGE grafts exhibited improvement in the novel object location task, but not the object recognition assay; MGE transplantation had no effect on memory in WT animals. These findings suggest that increasing the number of inhibitory interneurons is sufficient to restore deficits in long-term spatial memory of Chd2+/- mice, but recognition memory remains impaired.

DISCUSSION

Our results provide the first comprehensive molecular and physiological analysis of *Chd2* haploinsufficiency. Using a new mouse model of *Chd2* mutation, we found alterations in neural progenitor proliferation in the embryo as well as defects

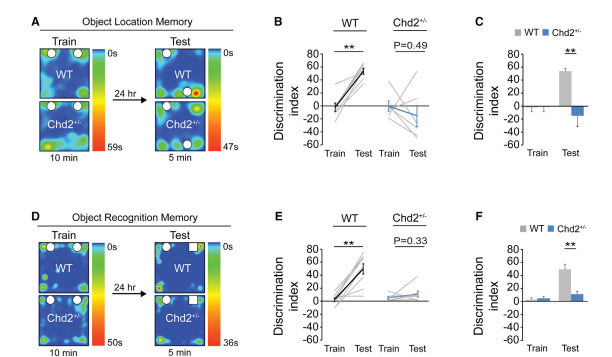


Figure 7. Chd2+/- Mice Exhibit Deficits in Long-Term Memory

(A) Heatmap showing the location of WT and $Chd2^{+/-}$ littermates during the entire training and testing phases of the object location memory (OLM) assay. (B and C) Discrimination index during training and testing phases of OLM (n = 7 WT mice and n = 6 $Chd2^{+/-}$ mice) for each individual mouse (B) and group means (C).

(D) Heatmap showing the location of WT and $Chd2^{+/-}$ littermates during the entire training and testing phases of the object recognition memory (ORM) assay. (E and F) Discrimination index during training and testing phases of ORM (n = 9 WT mice and n = 7 $Chd2^{+/-}$ mice) for each individual mouse (E) and group means (F).

Error bars, SEM; **p < 0.01. See also Figures S6 and S7.

in synaptic transmission, cortical synchrony, and hippocampal-dependent memory behavior in adult mice. In agreement with these findings, transcriptome analysis revealed broad changes in the expression of genes involved in chromatin regulation, neurogenesis, and synaptic transmission. Transplantation of MGE-derived interneurons rescued deficits in interneuron number and hippocampal-dependent spatial memory. While Chd2+/- mice exhibited changes in excitation-inhibition activity at the synaptic and network levels, we did not observe overt spontaneous seizures. However, many individuals with CHD2 haploinsufficiency also do not have epilepsy, and CHD2 mutations have been identified in patients with autism (O'Roak et al., 2014) and intellectual disability without seizures (Hamdan et al., 2014). Overall, these observations demonstrate that Chd2+/- mice reproduce many, but not all, of the key behavioral phenotypes observed in humans with Chd2 mutations. Careful phenotyping of additional patients will help better define the phenotypic spectrum of this disorder in human.

An emerging literature on chromatin biology is revealing that genes encoding CHD proteins are critical regulators of brain development, from cell proliferation to differentiation (Marfella et al., 2006, 2008; Marfella and Imbalzano, 2007; Shen et al., 2015; Durak et al., 2016; Katayama et al., 2016; Kwan et al., 2016; Tyagi et al., 2016; Mastrototaro et al., 2017; Sokpor

et al., 2017; Gompers et al., 2017; Platt et al., 2017; Meganathan et al., 2017). Whereas most studies have focused on cell signaling pathways and/or behavioral phenotypes, our results demonstrate a potential link between chromatin regulation and behavior via alterations in synaptic transmission and cortical network synchrony. Though CHD2 is located in cell nuclei, we found that Chd2 mutations perturb neural circuit function in at least two distinct ways in the mature brain depending on the system. In glutamatergic circuits, which have a prominent role in hippocampal-dependent learning and memory, we observed transcriptional changes related to pre- and post-synaptic domains that lead to an increase in glutamatergic transmission and faster event kinetics. In contrast, reduced progenitor proliferation predominates in GABAergic circuits, and this leads to fewer inhibitory interneurons and a reduction in synaptic inhibition in hippocampus. The balance of excitatory-inhibitory synaptic interactions is essential for the formation of neural oscillations (Atallah and Scanziani, 2009) and memory (Lim and Goldman, 2013), deficits we observed in $Chd2^{+/-}$ mice.

Early insight into the function of CHD2 in brain indicated expression of repressor element 1-silencing transcription factor (REST), a master regulator of neuronal development, was positively correlated with CHD2 (Shen et al., 2015), which may lead to alterations in radial glia self-renewal and neurogenesis when disrupted. Our RNA sequencing analysis did not confirm

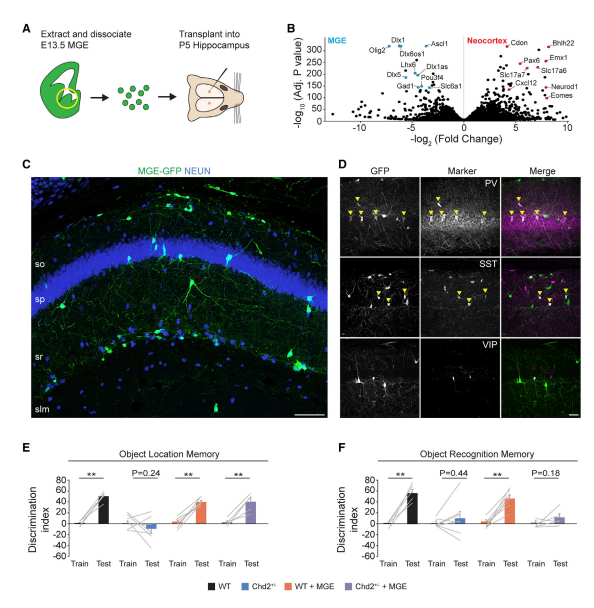


Figure 8. MGE Transplantation Rescues Hippocampal Memory Problems in Chd2+/- Mice

- (A) Schematic showing the experimental approach used for MGE transplantation.
- (B) Volcano plot of differentially expressed genes between E13.5 MGE and neocortex (n = 3 mice per genotype).
- (C) Hippocampus of a Chd2*/- mouse (45 DAT) labeled for NEUN (blue) and transplanted GFP-labeled inhibitory neurons (green). Abbreviations: so, stratum oriens; sp, stratum pyramidale; sr, stratum radiatum; slm, stratum lacunosum-moleculare.
- (D) At 45 DAT, GFP-labeled cells (green) co-expressed PV and SST but did not express VIP (all in magenta).

(E and F) Discrimination index during training and testing phases of OLM (E) and ORM (F) assays shows MGE transplantation rescues spatial memory deficits in $Chd2^{+/-}$ mice (n = 5-7 mice per treatment group).

Arrowheads, co-labeled cells; error bars, SEM; **p < 0.01; scale bars, 100 µm (C) and 50 µm (D). See also Figure S8.

downregulation of REST in any brain region examined. This may be related to differences between in utero knockdown via Chd2 shRNA delivery to embryonic ventricles and transgenic mutation; transcriptional changes in REST following in utero knockdown of Chd8 also could not be replicated by germline mutation (Durak et al., 2016; Katayama et al., 2016; Gompers et al., 2017; Platt et al., 2017). One of the main differences between Chd2+/- and other mouse models of Chd mutation (e.g., Chd8+/-) is that Chd2+/- mice have a severe interneuron deficit whereas no

such deficit has been reported in any of the mouse models of Chd8+/- (Durak et al., 2016; Katayama et al., 2016; Gompers et al., 2017; Platt et al., 2017). Indeed, a recent report suggests that Chd2 may co-associate with Nkx2.1 at cis-regulatory elements to regulate gene expression programs in presumptive interneurons derived from human embryonic stem cells (Meganathan et al., 2017). We found evidence of altered neurogenesis in Chd2+/- mice, notably a reduction in precursor pools of both excitatory and inhibitory neurons in the embryonic forebrain.

Nevertheless, a clear disruption of cortical cytoarchitecture was not apparent in mutant animals, consistent with the absence of structural brain abnormalities among many people with *CHD2* mutations (Chénier et al., 2014). It is possible that *Chd2* also plays a role in the development of non-neuronal cell types, such as oligodendrocytes or astrocytes. Our experiments primarily focused on analysis of neural progenitors, because we did not observe co-localization of CHD2 in GFAP+ cells or a change in OLIG2+ cell densities in *Chd2*+/- mice. A detailed and direct analysis of individual neuronal and glial cell types should clarify the developmental source or molecular mechanisms driving *Chd2*-related pathology. Nevertheless, our findings suggest that cell-type specification and migration of neural progenitors into the cortex remains largely intact in *Chd2*+/- mice.

Hundreds of rare gene mutations are now associated with neurodevelopmental disorders, many of which have devastating behavioral consequences that cannot be managed with available treatment options. Although discovery of new diseaseassociated gene mutations has been expanding rapidly, a major challenge in the field is to identify mechanisms underlying each of these rare disorders so that new, disease-specific therapies can be developed. Our results suggest interventions targeting cell proliferation and/or synaptic signaling may be particularly promising therapeutic candidates in Chd2 haploinsufficiency. Given that DO analysis revealed 16 DE genes associated with obesity and the strong link between altered metabolism and seizures (Scharfman, 2015), metabolic pathways may be a promising disease target as well. Indeed, some patients with CHD2 mutation found improvement with ketogenic diet (Chd2 Support and Research Group; https://www.facebook.com/ groups/1462485137354985/), and there is evidence of improved memory performance with this treatment in aging mice (Newman et al., 2017). While further studies of the neuroanatomical and physiological changes associated with Chd2 mutations are necessary before precise therapies can be designed and translated to the clinic, our results are an important step toward understanding how mutations in Chd2 impact brain development and function.

STAR*METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- CONTACT FOR REAGENT AND RESOURCE SHARING
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
 - Mice
- METHOD DETAILS
 - O Experimental Design
 - Western Blot
 - Immunostaining
 - Cell Quantification
 - Timm's Staining
 - O RNA Sequencing
 - O Differential Expression Analysis
 - O Quantitative (q) RT-PCR
 - Slice Electrophysiology

- Electroencephalography (EEG)
- Object Location and Recognition Memory Assays
- MGE Transplantation
- QUANTIFICATION AND STATISTICAL ANALYSIS
- DATA AND SOFTWARE AVAILABILITY

SUPPLEMENTAL INFORMATION

Supplemental Information includes eight figures and eight tables and can be found with this article online at https://doi.org/10.1016/j.neuron.2018.09.049.

ACKNOWLEDGMENTS

This work was supported by funding from the Lennox-Gastaut Syndrome Foundation; National Institutes of Health grants NINDS R00-NS085046, R01-NS096012, and T32-NS045540; UCI SURP and UROP Fellowship Programs; UCI Center for Autism Research and Translation (CART), and UCLA QCBio Collaboratory. We thank Haico van Attikum, Martijn Luijsterburg, and Wouter Wiegant for generously sharing western blot protocols; Daniel Vogt and John Rubenstein for kindly sharing GAD67-GFP mice; Sarah Smith and Kimberly Gonzalez for genotyping; Alex Lee with assistance with genotyping immunostaining; Jenny Wu and the UCI Genomics High-Throughput Facility for sequence alignment of RNA sequencing; and Daniel Vogt, Kei Igarashi, and Scott Baraban for helpful discussions and comments on earlier versions of this manuscript.

AUTHOR CONTRIBUTIONS

Y.J.K. contributed to the design, execution, and analysis of experiments and wrote the manuscript. S.K., S.A., J.C.F., B.Z., and S.L. contributed to the execution and analysis of experiments and helped edit the manuscript. Y.E.W. contributed to analysis of RNA-seq experiments. R.F.H. contributed to the concept, design, execution, and analysis of experiments and funding and wrote the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: October 10, 2017 Revised: July 31, 2018 Accepted: September 26, 2018 Published: October 18, 2018

REFERENCES

Allen, A.S., Berkovic, S.F., Cossette, P., Delanty, N., Dlugos, D., Eichler, E.E., Epstein, M.P., Glauser, T., Goldstein, D.B., Han, Y., et al.; Epi4K Consortium; Epilepsy Phenome/Genome Project (2013). De novo mutations in epileptic encephalopathies. Nature *501*, 217–221.

Atallah, B.V., and Scanziani, M. (2009). Instantaneous modulation of gamma oscillation frequency by balancing excitation with inhibition. Neuron 62, 566–577.

Cain-Hom, C., Splinter, E., van Min, M., Simonis, M., van de Heijning, M., Martinez, M., Asghari, V., Cox, J.C., and Warming, S. (2017). Efficient mapping of transgene integration sites and local structural changes in Cre transgenic mice using targeted locus amplification. Nucleic Acids Res. 45, e62.

Carlén, M., Meletis, K., Siegle, J.H., Cardin, J.A., Futai, K., Vierling-Claassen, D., Rühlmann, C., Jones, S.R., Deisseroth, K., Sheng, M., et al. (2012). A critical role for NMDA receptors in parvalbumin interneurons for gamma rhythm induction and behavior. Mol. Psychiatry *17*, 537–548.

Carvill, G.L., Heavin, S.B., Yendle, S.C., McMahon, J.M., O'Roak, B.J., Cook, J., Khan, A., Dorschner, M.O., Weaver, M., Calvert, S., et al. (2013). Targeted resequencing in epileptic encephalopathies identifies de novo mutations in CHD2 and SYNGAP1. Nat. Genet. 45, 825–830.

Carvill, G., Helbig, I., and Mefford, H. (2015). CHD2-related neurodevelopmental disorders. In GeneReviews®, R.A. Pagon, M.P. Adam, H.H. Ardinger, S.E. Wallace, A. Amemiya, L.J.H. Bean, T.D. Bird, N. Ledbetter, H.C. Mefford, R.J.H. Smith, and K. Stephens, eds. (University of Washington, Seattle), [Internet].

Chénier, S., Yoon, G., Argiropoulos, B., Lauzon, J., Laframboise, R., Ahn, J.W., Ogilvie, C.M., Lionel, A.C., Marshall, C.R., Vaags, A.K., et al. (2014). CHD2 haploinsufficiency is associated with developmental delay, intellectual disability, epilepsy and neurobehavioural problems. J. Neurodev. Disord. 6, 9.

Cho, K.K., Hoch, R., Lee, A.T., Patel, T., Rubenstein, J.L., and Sohal, V.S. (2015). Gamma rhythms link prefrontal interneuron dysfunction with cognitive inflexibility in Dlx5/6(+/-) mice. Neuron *85*, 1332–1343.

Cornew, L., Roberts, T.P., Blaskey, L., and Edgar, J.C. (2012). Resting-state oscillatory activity in autism spectrum disorders. J. Autism Dev. Disord. *42*, 1884–1894.

de Lanerolle, N.C., Kim, J.H., Robbins, R.J., and Spencer, D.D. (1989). Hippocampal interneuron loss and plasticity in human temporal lobe epilepsy. Brain Res. 495, 387–395.

De Rubeis, S., He, X., Goldberg, A.P., Poultney, C.S., Samocha, K., Cicek, A.E., Kou, Y., Liu, L., Fromer, M., Walker, S., et al.; DDD Study; Homozygosity Mapping Collaborative for Autism; UK10K Consortium (2014). Synaptic, transcriptional and chromatin genes disrupted in autism. Nature 515, 209-215

Deciphering Developmental Disorders Study (2015). Large-scale discovery of novel genetic causes of developmental disorders. Nature *519*, 223–228.

Del Pino, I., García-Frigola, C., Dehorter, N., Brotons-Mas, J.R., Alvarez-Salvado, E., Martínez de Lagrán, M., Ciceri, G., Gabaldón, M.V., Moratal, D., Dierssen, M., et al. (2013). Erbb4 deletion from fast-spiking interneurons causes schizophrenia-like phenotypes. Neuron *79*, 1152–1168.

Durak, O., Gao, F., Kaeser-Woo, Y.J., Rueda, R., Martorell, A.J., Nott, A., Liu, C.Y., Watson, L.A., and Tsai, L.H. (2016). Chd8 mediates cortical neurogenesis via transcriptional regulation of cell cycle and Wnt signaling. Nat. Neurosci. 19, 1477–1488

Galizia, E.C., Myers, C.T., Leu, C., de Kovel, C.G., Afrikanova, T., Cordero-Maldonado, M.L., Martins, T.G., Jacmin, M., Drury, S., Krishna Chinthapalli, V., et al.; EuroEPINOMICS CoGIE Consortium (2015). CHD2 variants are a risk factor for photosensitivity in epilepsy. Brain *138*, 1198–1207.

Gompers, A.L., Su-Feher, L., Ellegood, J., Copping, N.A., Riyadh, M.A., Stradleigh, T.W., Pride, M.C., Schaffler, M.D., Wade, A.A., Catta-Preta, R., et al. (2017). Germline Chd8 haploinsufficiency alters brain development in mouse. Nat. Neurosci. 20, 1062–1073.

Hamdan, F.F., Srour, M., Capo-Chichi, J.M., Daoud, H., Nassif, C., Patry, L., Massicotte, C., Ambalavanan, A., Spiegelman, D., Diallo, O., et al. (2014). De novo mutations in moderate or severe intellectual disability. PLoS Genet. *10*, e1004772.

Harada, A., Okada, S., Konno, D., Odawara, J., Yoshimi, T., Yoshimura, S., Kumamaru, H., Saiwai, H., Tsubota, T., Kurumizaka, H., et al. (2012). Chd2 interacts with H3.3 to determine myogenic cell fate. EMBO J. 31, 2994–3007.

Hirsch, J.C., Agassandian, C., Merchán-Pérez, A., Ben-Ari, Y., DeFelipe, J., Esclapez, M., and Bernard, C. (1999). Deficit of quantal release of GABA in experimental models of temporal lobe epilepsy. Nat. Neurosci. *2*, 499–500.

Howard, M.A., Rubenstein, J.L., and Baraban, S.C. (2014). Bidirectional homeostatic plasticity induced by interneuron cell death and transplantation in vivo. Proc. Natl. Acad. Sci. USA 111, 492–497.

Hunt, R.F., Scheff, S.W., and Smith, B.N. (2009). Posttraumatic epilepsy after controlled cortical impact injury in mice. Exp. Neurol. *215*, 243–252.

Hunt, R.F., Girskis, K.M., Rubenstein, J.L., Alvarez-Buylla, A., and Baraban, S.C. (2013). GABA progenitors grafted into the adult epileptic brain control seizures and abnormal behavior. Nat. Neurosci. *16*, 692–697.

lossifov, I., O'Roak, B.J., Sanders, S.J., Ronemus, M., Krumm, N., Levy, D., Stessman, H.A., Witherspoon, K.T., Vives, L., Patterson, K.E., et al. (2014). The contribution of de novo coding mutations to autism spectrum disorder. Nature *515*, 216–221.

Jensen, E.C. (2013). Quantitative analysis of histological staining and fluorescence using ImageJ. Anat. Rec. (Hoboken) 296, 378–381.

Katayama, Y., Nishiyama, M., Shoji, H., Ohkawa, Y., Kawamura, A., Sato, T., Suyama, M., Takumi, T., Miyakawa, T., and Nakayama, K.I. (2016). CHD8 haploinsufficiency results in autistic-like phenotypes in mice. Nature *537*, 675–679.

Khoshkhoo, S., Vogt, D., and Sohal, V.S. (2017). Dynamic, cell-type-specific roles for GABAergic interneurons in a mouse model of optogenetically inducible seizures. Neuron *93*, 291–298.

Korotkova, T., Fuchs, E.C., Ponomarenko, A., von Engelhardt, J., and Monyer, H. (2010). NMDA receptor ablation on parvalbumin-positive interneurons impairs hippocampal synchrony, spatial representations, and working memory. Neuron *68*, 557–569.

Kulkarni, S., Nagarajan, P., Wall, J., Donovan, D.J., Donell, R.L., Ligon, A.H., Venkatachalam, S., and Quade, B.J. (2008). Disruption of chromodomain helicase DNA binding protein 2 (CHD2) causes scoliosis. Am. J. Med. Genet. A. 146A. 1117–1127.

Kwan, V., Unda, B.K., and Singh, K.K. (2016). Wnt signaling networks in autism spectrum disorder and intellectual disability. J. Neurodev. Disord. 8, 45.

Levenson, J.M., and Sweatt, J.D. (2005). Epigenetic mechanisms in memory formation. Nat. Rev. Neurosci. 6, 108–118.

Lim, S., and Goldman, M.S. (2013). Balanced cortical microcircuitry for maintaining information in working memory. Nat. Neurosci. *16*, 1306–1314.

Love, M.I., Huber, W., and Anders, S. (2014). Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol. 15, 550.

Luijsterburg, M.S., de Krijger, I., Wiegant, W.W., Shah, R.G., Smeenk, G., de Groot, A.J.L., Pines, A., Vertegaal, A.C.O., Jacobs, J.J.L., Shah, G.M., and van Attikum, H. (2016). PARP1 links CHD2-mediated chromatin expansion and H3.3 deposition to DNA repair by non-homologous end-joining. Mol. Cell 61, 547–562.

Marfella, C.G., and Imbalzano, A.N. (2007). The Chd family of chromatin remodelers. Mutat. Res. 618, 30–40.

Marfella, C.G., Ohkawa, Y., Coles, A.H., Garlick, D.S., Jones, S.N., and Imbalzano, A.N. (2006). Mutation of the SNF2 family member Chd2 affects mouse development and survival. J. Cell. Physiol. 209, 162–171.

Marfella, C.G., Henninger, N., LeBlanc, S.E., Krishnan, N., Garlick, D.S., Holzman, L.B., and Imbalzano, A.N. (2008). A mutation in the mouse Chd2 chromatin remodeling enzyme results in a complex renal phenotype. Kidney Blood Press. Res. *31*, 421–432.

Mastrototaro, G., Zaghi, M., and Sessa, A. (2017). Epigenetic mistakes in neurodevelopmental disorders. J. Mol. Neurosci. *61*, 590–602.

Meganathan, K., Lewis, E.M.A., Gontarz, P., Liu, S., Stanley, E.G., Elefanty, A.G., Huettner, J.E., Zhang, B., and Kroll, K.L. (2017). Regulatory networks specifying cortical interneurons from human embryonic stem cells reveal roles for CHD2 in interneuron development. Proc. Natl. Acad. Sci. USA *114*, E11180–E11189.

Mews, P., Donahue, G., Drake, A.M., Luczak, V., Abel, T., and Berger, S.L. (2017). Acetyl-CoA synthetase regulates histone acetylation and hippocampal memory. Nature *546*, 381–386.

Neale, B.M., Kou, Y., Liu, L., Ma'ayan, A., Samocha, K.E., Sabo, A., Lin, C.F., Stevens, C., Wang, L.S., Makarov, V., et al. (2012). Patterns and rates of exonic de novo mutations in autism spectrum disorders. Nature 485, 242–245.

Newman, J.C., Covarrubias, A.J., Zhao, M., Yu, X., Gut, P., Ng, C.P., Huang, Y., Haldar, S., and Verdin, E. (2017). Ketogenic diet reduces midlife mortality and improves memory in aging mice. Cell Metab. *26*, 547–557.e8.

O'Roak, B.J., Stessman, H.A., Boyle, E.A., Witherspoon, K.T., Martin, B., Lee, C., Vives, L., Baker, C., Hiatt, J.B., Nickerson, D.A., et al. (2014). Recurrent de novo mutations implicate novel genes underlying simplex autism risk. Nat. Commun. 5, 5595.

Opperman, K.J., Mulcahy, B., Giles, A.C., Risley, M.G., Birnbaum, R.L., Tulgren, E.D., Dawson-Scully, K., Zhen, M., and Grill, B. (2017). The HECT family ubiquitin ligase EEL-1 regulates neuronal function and development. Cell Rep. 19, 822–835.

Orekhova, E.V., Stroganova, T.A., Nygren, G., Tsetlin, M.M., Posikera, I.N., Gillberg, C., and Elam, M. (2007). Excess of high frequency electroencephalogram oscillations in boys with autism. Biol. Psychiatry *62*, 1022–1029.

Petryniak, M.A., Potter, G.B., Rowitch, D.H., and Rubenstein, J.L. (2007). Dlx1 and Dlx2 control neuronal versus oligodendroglial cell fate acquisition in the developing forebrain. Neuron *55*, 417–433.

Platt, R.J., Zhou, Y., Slaymaker, I.M., Shetty, A.S., Weisbach, N.R., Kim, J.A., Sharma, J., Desai, M., Sood, S., Kempton, H.R., et al. (2017). Chd8 mutation leads to autistic-like behaviors and impaired striatal circuits. Cell Rep. *19*, 335–350.

Rauch, A., Wieczorek, D., Graf, E., Wieland, T., Endele, S., Schwarzmayr, T., Albrecht, B., Bartholdi, D., Beygo, J., Di Donato, N., et al. (2012). Range of genetic mutations associated with severe non-syndromic sporadic intellectual disability: an exome sequencing study. Lancet *380*, 1674–1682.

Satoh, J., Tabira, T., Sano, M., Nakayama, H., and Tateishi, J. (1991). Parvalbumin-immunoreactive neurons in the human central nervous system are decreased in Alzheimer's disease. Acta Neuropathol. *81*, 388–395.

Scharfman, H.E. (2015). Neuroscience. Metabolic control of epilepsy. Science 347, 1312–1313.

Scheel, J.R., Garrett, L.J., Allen, D.M., Carter, T.A., Randolph-Moore, L., Gambello, M.J., Gage, F.H., Wynshaw-Boris, A., and Barlow, C. (2003). An inbred 129SvEv GFPCre transgenic mouse that deletes loxP-flanked genes in all tissues. Nucleic Acids Res. 31, e57.

Shen, T., Ji, F., Yuan, Z., and Jiao, J. (2015). CHD2 is required for embryonic neurogenesis in the developing cerebral cortex. Stem Cells 33, 1794–1806.

Skarnes, W.C., Rosen, B., West, A.P., Koutsourakis, M., Bushell, W., Iyer, V., Mujica, A.O., Thomas, M., Harrow, J., Cox, T., et al. (2011). A conditional knockout resource for the genome-wide study of mouse gene function. Nature 474. 337–342.

Sokpor, G., Xie, Y., Rosenbusch, J., and Tuoc, T. (2017). Chromatin remodeling BAF (SWI/SNF) complexes in neural development and disorders. Front. Mol. Neurosci. 10, 243.

Southwell, D.G., Paredes, M.F., Galvao, R.P., Jones, D.L., Froemke, R.C., Sebe, J.Y., Alfaro-Cervello, C., Tang, Y., Garcia-Verdugo, J.M., Rubenstein, J.L., et al. (2012). Intrinsically determined cell death of developing cortical interneurons. Nature *491*, 109–113.

Tamamaki, N., Yanagawa, Y., Tomioka, R., Miyazaki, J., Obata, K., and Kaneko, T. (2003). Green fluorescent protein expression and colocalization with calretinin, parvalbumin, and somatostatin in the GAD67-GFP knock-in mouse. J. Comp. Neurol. *467*, 60–79.

Tauck, D.L., and Nadler, J.V. (1985). Evidence of functional mossy fiber sprouting in hippocampal formation of kainic acid-treated rats. J. Neurosci. 5. 1016–1022.

Tyagi, M., Imam, N., Verma, K., and Patel, A.K. (2016). Chromatin remodelers: we are the drivers!! Nucleus 7, 388–404.

Vaudano, A.E., Ruggieri, A., Avanzini, P., Gessaroli, G., Cantalupo, G., Coppola, A., Sisodiya, S.M., and Meletti, S. (2017). Photosensitive epilepsy is associated with reduced inhibition of alpha rhythm generating networks. Brain *140*. 981–997.

Vogel-Ciernia, A., and Wood, M.A. (2014). Examining object location and object recognition memory in mice. Curr. Protoc. Neurosci. 69, 1–17.

Vogel-Ciernia, A., Matheos, D.P., Barrett, R.M., Kramár, E.A., Azzawi, S., Chen, Y., Magnan, C.N., Zeller, M., Sylvain, A., Haettig, J., et al. (2013). The neuron-specific chromatin regulatory subunit BAF53b is necessary for synaptic plasticity and memory. Nat. Neurosci. *16*, 552–561.

Vogt, D., Hunt, R.F., Mandal, S., Sandberg, M., Silberberg, S.N., Nagasawa, T., Yang, Z., Baraban, S.C., and Rubenstein, J.L.R. (2014). Lhx6 directly regulates Arx and CXCR7 to determine cortical interneuron fate and laminar position. Neuron 82, 350–364.

Walter, W., Sánchez-Cabo, F., and Ricote, M. (2015). GOplot: an R package for visually combining expression data with functional analysis. Bioinformatics *31*, 2912–2914.

Willoughby, J.O., Fitzgibbon, S.P., Pope, K.J., Mackenzie, L., Medvedev, A.V., Clark, C.R., Davey, M.P., and Wilcox, R.A. (2003). Persistent abnormality detected in the non-ictal electroencephalogram in primary generalised epilepsy. J. Neurol. Neurosurg. Psychiatry 74, 51–55.

Young, M.D., Wakefield, M.J., Smyth, G.K., and Oshlack, A. (2010). Gene ontology analysis for RNA-seq: accounting for selection bias. Genome Biol. 11, R14.

Yu, G., Wang, L.-G., Yan, G.-R., and He, Q.-Y. (2015). DOSE: an R/Bioconductor package for disease ontology semantic and enrichment analysis. Bioinformatics 31, 608–609.

STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies	· · · · · · · · · · · · · · · · · · ·	
β-actin, mouse (1:10,000, WB)	Sigma-Aldrich	Cat# A1978, RRID: AB_476692
Chd2, rat (1:300-1:500, IF; 1:1,000, WB)	Millipore	Cat# MABE873 RRID: AB_2737347
CR, rabbit (1:1,000, IF)	Millipore	Cat# AB5054, RRID: AB_2068506
Gad67, mouse (1:1,000, IF)	Millipore	Cat# MAB5406, RRID: AB_2278725
GFAP, mouse (1:500, IF)	Millipore	Cat# MAB3402, RRID: AB_94844
GFP, chicken (1:1,000, IF)	Aves Labs	Cat# GFP-1020, RRID: AB_10000240
Ki-67, mouse (1:500, IF)	BD Biosciences	Cat# 556003, RRID: AB_396287
NeuN, mouse (1:500, IF)	R&D Systems	Cat# MAB377, RRID: AB_2298767
Nkx2.1, rabbit (1:500, IF)	Abcam	Cat# ab76013, RRID: AB_1310784
Dlig2, rabbit (1:300, IF)	Millipore	Cat# AB9610, RRID: AB_570666
Pax6, mouse (1:500, IF)	BioLegend	Cat# 901301, RRID: AB_2565003
PV, mouse (1:500, IF)	Sigma-Aldrich	Cat# P3088, RRID: AB_477329
Reelin, mouse (1:500, IF)	Millipore	Cat# MAB5364, RRID: AB_2179313
SST, goat (1:200, IF)	Santa Cruz Biotechnology	Cat# sc-7819, RRID: AB_2302603
Гbr1, rabbit (1:500, IF)	Abcam	Cat# ab31940, RRID: AB_2200219
Tbr2, rabbit (1:500, IF)	Abcam	Cat# ab23345, RRID: AB_778267
Tuj1, mouse (1:1,000, IF)	BioLegend	Cat# 801201, RRID: AB_2313773
/IP, rabbit (1:200, IF)	ImmunoStar	Cat# 20077 RRID: AB_572270
Caspase-3, rabbit (1:500, IF)	Cell Signaling Technology	Cat# 9661S, RRID: AB_2341188
3rn2, mouse (1:500, IF)	Santa Cruz	Cat# sc-393324, RRID: AB_2737345
Ctip2, rat (1:500, IF)	Abcam	Cat# ab18465 RRID: AB_10015215
Chemicals, Peptides, and Recombinant Proteins		
Bicuculine	Sigma-Aldrich	14343
Kynurenic acid	Sigma-Aldrich	K3375
TX	Tocris	1078
Critical Commercial Assays		
Direct-zol RNAMiniPrep Plus kit	Zymo Research	Cat# R2063
NEBNext Ultra RNA Library Prep Kit for Illumina	NEB	Cat# E7530S
Deposited Data		
RNA sequencing data	This paper	GEO: GSE112196
Experimental Models: Organisms/Strains		
Nouse: Chd2tm1c(EUCOMM)Hmgu	This paper	N/A
Mouse: B6N.FVB-Tmem163Tg(ACTB-cre)2Mrt/CjDswJ	The Jackson Laboratory	Jax #:003376
Mouse: GAD67-GFP	Tamamaki et al., 2003	N/A
Mouse: C57BL/6J-Tg(Nkx2-1-cre)2Sand/J	The Jackson Laboratory	Jax #:008661
Mouse: B6;129S6-Gt(ROSA)26Sortm14(CAG-tdTomato)Hze/J	The Jackson Laboratory	Jax #:007908
Mouse: C57BL/6-Tg(CAG-EGFP)131Osb/LeySopJ	The Jackson Laboratory	Jax#:006567
Mouse: CD-1 IGS Mouse	Charles River Laboratories	Charles River#: 022
Dligonucleotides		
Chd2(tm1c) forward primer: AAGGCGCATAACGATACCAC	Skarnes et al., 2011	N/A
Chd2(tm1c) reverse primer: CCGCCTACTGCGACTATAGAGA	Skarnes et al., 2011	N/A

(Continued on next page)

Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Chd2(tm1d) forward primer: AAGGCGCATAACGATACCAC	Skarnes et al., 2011	N/A
Chd2(tm1d) reverse primer: ACTGATGGCGAGCTCAGACC	Skarnes et al., 2011	N/A
GFP forward primer: ACGTAAACGGCCACAAGTTC	Tamamaki et al., 2003	N/A
GFP reverse primer: AAGTCGTGCTTCATGTG	Tamamaki et al., 2003	N/A
Cre forward primer: CCGGGCTGCCACGACCAA	Scheel et al., 2003	N/A
Cre reverse primer: GGCGCGGCAACACCATTTTT	Scheel et al., 2003	N/A
Chd2_1 qpcr forward: AGGAGGTCAAATCGAAGCAGA	PrimerBank - MGH-PGA	N/A
Chd2_1 qpcr reverse: GCCTCTTCTTTTCGGACTCCC	PrimerBank - MGH-PGA	N/A
Chd2_2 qpcr forward: CTTCTGAGAGCCAATCAGAGTC	PrimerBank - MGH-PGA	N/A
Chd2_2 qpcr reverse: CATCAGCTATCCGTTCCTT	PrimerBank - MGH-PGA	N/A
Vipr2 qpcr forward: GACCTGCTACTGCTGGTTG	PrimerBank - MGH-PGA	N/A
Vipr2 qpcr reverse: CAGCTCTGCACATTTTGTCTCT	PrimerBank - MGH-PGA	N/A
Hddc3 qpcr forward: CCGCTCACAAACACCGACA	PrimerBank - MGH-PGA	N/A
Hddc3 qpcr reverse: GCGGCCTGTAACACCACAA	PrimerBank - MGH-PGA	N/A
Myo7a qpcr forward: AGGGGGACTATGTATGGATGGA	PrimerBank - MGH-PGA	N/A
Myo7a qpcr reverse: ATGTGCGTGGCATTCTGAGG	PrimerBank - MGH-PGA	N/A
Slc6a13 gpcr forward: CAGTACACCAACCAGGGAGG	PrimerBank - MGH-PGA	N/A
Slc6a13 qpcr reverse: GCCAGGACAACGATGTAGTAGA	PrimerBank - MGH-PGA	N/A
	PrimerBank - MGH-PGA	N/A
Ago2 aper reverse: CCAGAGGTATGGCTTCCTTCA	PrimerBank - MGH-PGA	N/A
Ago2 qpcr reverse: CCAGAGGTATGGCTTCCTTCA		N/A
Lgr6 qpcr forward: GAGGACGGCATCATGCTGTC	PrimerBank - MGH-PGA	
Lgr6 qpcr reverse: GCTCCGTGAGGTTGTTCATACT	PrimerBank - MGH-PGA	N/A
Actb qpcr forward: GGCTGTATTCCCCTCCATCG	PrimerBank - MGH-PGA	N/A
Actb qpcr reverse: CCAGTTGGTAACAATGCCATGT	PrimerBank - MGH-PGA	N/A
Software and Algorithms		
pClamp 10.7	Molecular Devices	https://www.moleculardevices.com/ products/axon-patch-clamp-system/ acquisition-and-analysis-software/ pclamp-software-suite
MiniAnlysis 6.0.7	Synaptosoft	https://www.moleculardevices.com/ products/axon-patch-clamp-system/ acquisition-and-analysis-software/ pclamp-software-suite
Sigmaplot 13.1	Systat	http://www.synaptosoft.com/ MiniAnalysis/
SYSTAT 13.1	SYStat	https://systatsoftware.com/
MATLAB	MathWorks	https://www.mathworks.com/ products/matlab.html; RRID: SCR_001622
DESeq2	Love et al., 2014	RRID: SCR_015687
ANY-maze	ANY-maze	http://anymaze.co.uk/
Goseq	Young et al., 2010	RRID: SCR_001875
DOSE	Yu et al., 2015	https://bioconductor.org/packages/ release/bioc/html/DOSE.html
Goplot	Walter et al., 2015	http://wencke.github.io/
pheatmap	N/A	https://cran.r-project.org/web/ packages/pheatmap/index.html
Other		



CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Robert F. Hunt (robert.hunt@uci.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice

Mice were maintained in standard housing conditions on a 12h light/dark cycle with food and water provided *ad libitum*. All protocols and procedures followed the guidelines of the University Laboratory Animal Resources at the University of California, Irvine and adhered to National Institutes of Health Guidelines for the Care and Use of Laboratory Animals. To generate *Chd2*-flox mice (i.e., Chd2^{tm1c(EUCOMM)Hmgu} mice), we obtained sperm with tm1c conditional allele from The Centre for Phenogenomics and re-derived the transgenic line on a C57BL/6J background (Jackson Laboratories cat. no. 000664) at the UCI Transgenic Mouse Facility. A full description of the targeting of *Chd2* can be found at International Mouse Phenotyping Consortium (https://www.mousephenotype.org/data/genes/MGI:2448567). *Chd2*-flox mice were then crossed for three generations to WT C57BL/6J mice and the offspring of the third generation was crossed to an ACTB-Cre line (Jackson Laboratories cat. no. 019099) to generate *Chd2*+/- mice. In some experiments, *Chd2*-flox mice were mated with a *Nkx2.1*-Cre (Jackson Laboratories cat. no. 008661), an Ai14 tdTomato reporter (Jackson Laboratories cat. no. 007908) or a hemizygous glutamic acid decarboxylase - enhanced green fluorescence protein (GAD67-GFP) knockin line maintained on a CD-1 background (Tamamaki et al., 2003). To obtain embryos for MGE transplantation, male β-actin:GFP mice (Jackson Laboratories cat. no. 006567) were crossed to CD-1 females (Charles River, cat no. 022). Experiments were performed on male and female littermates between E14.5 and P75.

METHOD DETAILS

Experimental Design

Experiments were performed on male and female littermates between E14.5 and P75. Animals were randomly allocated to experimental groups, and the experimenter was blinded to genotype of the animals until the experiment was complete. No data or animals were excluded from analysis. Memory behavior assays were replicated using a separate, independent cohort of WT and $Chd2^{+/-}$ littermates. No other replication studies were performed.

Western Blot

Western blot was performed on P30 mice as previously described (Luijsterburg et al., 2016). Nuclear protein extracts from whole brain were prepared by homogenizing the tissue in subcellular fractionation buffer (20 mM HEPES (pH7.4), 74.55 mM KCl, 95.21 mM MgCl2, 292.24 mM EDTA, 380.35 mM EGTA, 1 mM DTT and 1x Protease inhibitor cocktail). Proteins were separated by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and transferred to PVDF membranes. Protein expression was analyzed by immunoblotting. Primary antibodies and dilutions are provided in the Key Resources table. Secondary antibodies were IRDye 800CW (LI-COR) and CF770 (Biotium). Gels were imaged and quantified using a LI-COR Odyssey infrared imaging scanning system.

Immunostaining

Mice were transcardially perfused with 4% paraformaldehyde (PFA) and free-floating vibratome sections (50 μm) were processed using standard immunostaining procedures (Hunt et al., 2013). For E14.5 embryos, brains were dissected and drop-fixed overnight in 4% PFA. For PAX6, KI67, and TBR2 staining, sections were pretreated with an antigen unmasking solution (10mM Sodium Citrate, pH 6.0) for 60 min at room temperature. All antibodies have been previously used for immunostaining analysis in brain. Primary antibodies and dilutions are provided in the Key Resources table. Secondary antibodies were: Alexa 488, Alexa 546, Alexa 594 and Alexa 647 (Invitrogen). Sections were then mounted on charged slides (Superfrost plus; Fisher Scientific) with Vectashield that contained DAPI or Aqua Mount. Images were obtained with a Leica DM6 epifluorescence microscope or an Olympus FV3000 laser-scanning confocal microscope. Brightness and contrast were adjusted manually using Adobe Photoshop; z stacks were generated using Olympus, Leica or Helicon Focus 6 software.

Cell Quantification

Fluorescently labeled sections ($50 \mu m$) were imaged using a Leica DM6 microscope with a x10 or x20 objective or Olympus FV3000 confocal microscope with a x20 or x40 objective and counted using ImageJ, as described previously (Hunt et al., 2013). All cells that expressed a subtype maker were counted in every sixth coronal section (that is, $300 \mu m$ apart). Four to six sections were analyzed per animal and the values averaged to obtain a mean cell density (cells / mm^2). Intensity measurements were analyzed within regions of interest and a range of threshold limit was applied using ImageJ according to a previous protocol (Jensen, 2013).

Timm's Staining

Animals were perfused transcardially with 0.37% sodium sulfide solution in 0.1 M NaHPO4, followed by 4% PFA and stored overnight in 4% PFA solution. Brains were then sectioned at $50~\mu m$ on a vibratome and every sixth section mounted on charged slides (Superfrost Plus; Fisher Scientific). Sections were dried overnight and treated according to previous protocols using Timm's stain to reveal mossy fibers and Nissl counterstained with cresyl violet to visualize cell bodies (Tauck and Nadler, 1985; Hunt et al., 2009). To semiquantitatively assess the presence or absence of mossy fiber sprouting into the inner molecular layer of the dentate gyrus, Timm scores from 0 (little to no sprouting) to 3 (robust mossy fiber sprouting) (Tauck and Nadler, 1985; Hunt et al., 2009) were assigned to six randomly chosen sections from the dorsal portion of the hippocampus and averaged for each animal. Images were taken with a Leica MZ10F stereoscope.

RNA Sequencing

Total RNA was isolated from freshly dissected tissue using the Direct-zol RNAMiniPrep Plus kit (Zymo Research) according to the manufacturer instructions. RNA quantity and quality were then determined for each sample using NanoDrop (ThermoFisher), Qubit RNA BR Assay Kit (ThermoFisher), and a TapeStation (Agilent). Samples were then converted to cDNA using the NEBNext Ultra RNA Library Preparation Kit (NEB) with poly(A) selection. Paired-end sequencing was performed on the Illumina HiSeq 4000 platform by GENEWIZ Next Generation Sequencing facility (South Plainfield, NJ). Quality of the raw sequencing reads was accessed using FASTQC and adaptors were trimmed with Trimmomatic. Reads for each library (32.7 \pm 4.6 million per replicate, n = 18 replicates) were then aligned using TopHat2 (version 2.1.1) and the mouse genome index mm10 generated from iGenome UCSC mm10 FASTQ genome sequence.

Differential Expression Analysis

Quantification and differential expression of the annotated mouse genes were performed using DESeq2 as previously described (Love et al., 2014) using three biological replicates for each dataset. Differences were considered statistically significant with Benjamini and Hochberg (BH) adjusted p value < 0.10 and p values < 0.05. GO analysis and functional classification were performed separately on up- or downregulated genes using the R package *goseq* (Young et al., 2010), with corrected gene length and expressed genes as the background test set. Significance was set at Benjamini and Hochberg adjusted p value < 0.1 or p value < 0.05. DO analysis of functional enrichment based on human diseases in the DisGeNET database (http://www.disgenet.org/) was performed separately on up- or downregulated genes using the R package *DOSE* (Yu et al., 2015) and expressed genes as the background test set. Heatmaps and bubble plot were generated with the R packages *pheatmap* and *GOplot*. For disease gene plots in Figure 4F, risk genes for epilepsy, intellectual disability and/or autism were determined based on the following databases: SFARI Human Gene Module (https://gene.sfari.org/database/human-gene), Citizens United Against Epilepsy (CURE) Foundation Epilepsy Genetics Initiative (https://www.cureepilepsy.org/egi), DECIPHER v9.17 (https://decipher.sanger.ac.uk) and ID Gene Database Project (http://gfuncpathdb.ucdenver.edu/iddrc/iddrc/home.php). We found a significant disruption in *Tmem163* of *Chd2*+/- mice (Table S2), which is likely the result of Cre insertion into intron 4 of *Tmem163* in the ACTB-Cre line (Cain-Hom et al., 2017). Since we analyzed three female and three male samples by chance for the comparison of WT and *Chd2*+/- MGE, we also found several sex-specific genes with significant and large changes (Table S2).

Quantitative (q) RT-PCR

Total RNA was extracted with Direct-zol RNA MiniPrep Plus (Zymo Research) and reverse-transcription performed using iScript cDNA Synthesis Kit (Bio-Rad) according to the manufacturer instructions. The resulting cDNA was subjected to qPCR analysis with the Applied Biosystems ViiA 7 Real-Time PCR System using SsoAdvanced Universal Inhibitor-Tolerant SYBR Green Supermix (Bio-Rad cat no.1725016) and gene specific primers (Key Resources table). Reactions were repeated in triplicates. Relative expression levels were calculated using the $2^{-\Delta\Delta CT}$ method using *Actb* as an endogenous control gene.

Slice Electrophysiology

Coronal brain slices (300 μ m thickness) were prepared from WT and $Chd2^{+/-}$ littermates at P30-P35. Slices were submerged in the recording chamber and continuously perfused with oxygenated ACSF (32 –34°C) containing (in mM): 124 NaCl, 3 KCl, 1.25 NaH₂PO₄-H₂O, 2 MgSO₄-7H₂O, 26 NaHCO₃, 10 dextrose, and 2 CaCl₂ (pH 7.2–7.4, 300–305 mOsm/kg). Whole-cell patch-clamp recordings were performed at x40 using an upright, fixed-stage microscope (Olympus BX51WI) equipped with infrared, differential interference contrast (IR-DIC) and epifluorescence optics. For current-clamp and voltage-clamp recordings of EPSCs, patch pipettes (3–5 M Ω) were filled with an internal solution containing (in mM): 140 K⁺ gluconate, 1 NaCl, 5 EGTA, 10 HEPES, 1 MgCl₂, 1 CaCl₂, 3 KOH, and 2 ATP, pH 7.25. For voltage-clamp recordings of IPSCs, patch pipettes (3–5 M Ω) were filled with an internal solution containing (in mM): 140 CsCl, 11 EGTA, 10 HEPES, 1 MgCl₂, 2 NaATP, 0.5 NaGTP and 1.25 QX-314, pH 7.2. Recordings were obtained with a Multiclamp 700B amplifier, filtered at 4 kHz, and recorded to pClamp 10.7 software (Clampfit; Axon Instruments). For current-clamp experiments, cells were held at -70 mV, and electrophysiological properties were measured in response to a series of long (1000 ms) hyperpolarizing and depolarizing current-injections (10 pA steps; range: -80 pA to 160 pA). Voltage-clamp recordings were examined at a holding potential of -70 mV. Glutamatergic currents were measured in the presence of 30uM bicuculine, GABAergic currents were measured in the presence of 1 mM kynurenic acid and 1 uM TTX was added to the bath to isolate miniature

PSCs. A concentric bipolar stimulating electrode made of platinum–iridium wire (125 μ m diameter; FHC) was used to apply paired stimuli to CA1 stratum radiatum (100 μ s duration, 100ms interstimulus interval). Stimulus intensity was set at 1.5 x threshold, where threshold was defined as the stimulus required for evoking an IPSC in 50% of the trials. Ten consecutive responses were obtained at 0.1 Hz, averaged and the ratio of the amplitude of the second evoked IPSC to the first (elPSC2/elPSC1) was calculated to obtain the paired-pulse ratio (PPR). Series resistance was uncompensated and monitored throughout the recordings. Data were only used for analysis if the series resistance remained < 20 M Ω and changed by \leq 20% during the recordings. Recordings were not corrected for a liquid junction potential. Resting membrane potentials were measured immediately after breakthrough by temporarily removing the voltage clamp and monitoring voltage. Data analysis was performed using pClamp 10.7, MiniAnalysis 6.0.7 (Synaptosoft), Microsoft excel or Sigmaplot 13 programs. A 2 min sample recording per cell was used for measuring event frequency, amplitude, 10%–90% rise time, and decay time constant. Events characterized by a typical fast rising phase and exponential decay phase were manually detected using MiniAnalysis. The threshold for event detection was currents with amplitudes greater than three times the root mean square (RMS) noise level.

Electroencephalography (EEG)

Male Chd2+/- mice and age-matched male WT littermates were housed together before and after EEG surgeries. Local field potential recordings were obtained at 2kHz using an EEG monitoring system (Pinnacle Technologies) as previously described (Hunt et al., 2013; Khoshkhoo et al., 2017). Briefly, mice were anesthetized with ketamine and xylazine (10 mg/kg and 1 mg/kg i.p.), and sterile, stainless steel screw recording electrodes were placed epidurally through burr holes in the skull using surface head-mount EEG hardware (Pinnacle Technologies). The EEG recording electrodes were implanted at (in millimeters relative to the bregma): -1.25AP, ± 1.5ML; reference electrode was implanted at -5.0 AP, 0 ML (in cerebellum), and a ground electrode was implanted at -3.5 AP, -3.0 ML. Electrodes were cemented in place with a fast-acting adhesive and dental acrylic. Two wires were laid on the shoulder muscles for electromyographic (EMG) recording. Animals were allowed to recover for at least 7 days before experiments were initiated and then monitored for 7-10 days (24 h/day). EEG recordings for each mouse were visualized in 1 hr time windows and carefully inspected for any abnormal electrographic signals or seizures. This process was repeated twice for each animal. All analyses were performed using custom-written and built-in MATLAB (MathWorks) functions. To generate EEG power plots, total EEG power for each frequency interval was calculated in 10 min bins using the MATLAB function bandpower and averaged over 1 hr intervals. Then, the output was normalized by the total area under the curve (for frequencies < 100 Hz, excluding 59-61 Hz) for each mouse. To analyze changes in baseline power 60 hr of continuous EEG recording, hours 12 through 72, was included for each mouse in order reduce the effect of arousal and environmental noise in our analysis. Two-way ANOVA was used to assess the effect of genotype, frequency, and genotype-frequency on baseline EEG power followed by a Tukey-Kramer multiple comparisons test to compare EEG power for each frequency band. To assess inter-hemispheric synchronization in each frequency interval, coherence between the EEG signals from the two hemispheres was calculated using the MATLAB function mscohere in 10 min bins, and averaged over 1 hr. A total of 60 hr of continuous EEG recording was included in this analysis for each mouse. Two-way ANOVA was used to assess the effect of genotype, frequency, and genotype-frequency on inter-hemispheric coherence followed by a Tukey-Kramer multiple comparisons test to compare inter-hemispheric coherence in mutant and WT mice for each frequency band.

Object Location and Recognition Memory Assays

Each object location task consisted of a habituation phase, training phase, and testing phase and were performed according to a previous protocol (Vogel-Ciernia et al., 2013; Vogel-Ciernia and Wood, 2014). Mice were handled for 2-5 min on 5 consecutive days before testing. On day 1, animals were habituated individually to the open field arena. Mice were placed in the center of a 40 cm × 40 cm × 35 cm open field arena with a vertical marking strip for 10 min under dim overhead lighting conditions (45 lux). For the training session (day 2), two identical objects were placed in the open field, 1 cm from the back wall and mice were placed in the center of the opposite wall. Animals were allowed to explore each object for 10 min. The arena and objects were cleaned with 70% (v/v) EtOH (OLM) or 1% acetic acid (ORM) between trials. A retention test was performed 24 hr after the training session (day 3). For OLM, one object was placed in a different location. For ORM, one object was exchanged for a new object that was different in size, shape, and texture. The objects used were Falcon 50mL conical centrifuge Tubes (Fisher, Cat no. 14-432-22) filled with beach sand, 3D printed cube and a 75mL glass flask. For detailed description of the experimental setups, schematics are shown in Figures S6A,B and S7C,E. All behavioral assays were conducted between 2pm and 6pm during the light phase of the light/dark cycle (lights off at 8pm; lights on at 8am). Mouse identities were coded, and all behaviors were performed using a video tracking system and analyzed using ANY-maze software by investigators who were blind to the genotype and treatment of the animals. A mouse was scored as exploring an object when its head was oriented toward the object within a distance of 1 cm or when the nose was touching the object. The relative exploration time was recorded and expressed by a discrimination index (DI = $[t_{novel} - t_{familiar}]/[t_{novel} + t_{familiar}]/[t$ t_{familiar}] × 100) where t represents time. Mean exploration times were calculated and the discrimination indexes between treatment groups were compared. To diminish bias, control and mutant littermates were evaluated on the same day in the same arena, and the location of the novel object was counterbalanced across experiments and groups.

MGE Transplantation

Ventricular and subventricular layers of the MGE were harvested from E13.5 GFP+ embryos. The time point at which the sperm plug was detected was considered E0.5. Embryonic MGE explants were dissected in Leibovitz L-15 medium, mechanically dissociated by

repeated pipetting and concentrated by centrifugation (3 min at 1,000 g). Concentrated cell suspensions (\sim 10 3 cells nI $^{-1}$) were front loaded into beveled glass micropipettes (\sim 50- μ m tip diameter, Wiretol 5 μ I, Drummond Scientific) and bilaterally injected (5 × 10 4 cells per injection) into dorsal hippocampus of WT or $Chd2^{+/-}$ littermates at P5 as described previously (Hunt et al., 2013; Vogt et al., 2014).

QUANTIFICATION AND STATISTICAL ANALYSIS

Samples were randomly collected across litters and processed blind to genotype. All analyses were performed with SYSTAT 13.1 software and assessed for normality (Shapiro-Wilk) and variance (Brown-Forsythe). Mossy fiber sprouting scores and intrinsic electrophysiological properties were assessed by nonparametric Mann-Whitney U test on ranks. All other data were compared by two-tailed t test, one-way ANOVA for multiple comparisons or by two-way repeated-measures ANOVA. A Tukey's post hoc test was performed when appropriate. Sample sizes for behavior assays were determined by power analyses using 20% as a value for β (1 – β = power or 80%), 0.05 as a value for α and an expected 30% difference between groups; post hoc power for each behavior assay is indicated in Table S7. No statistical methods were used to predetermine sample size in other experiments. Sample sizes can be found within results and/or figure legends, and individual data points are shown for each quantification. For slice electrophysiology studies, experimental data were averaged across neurons (i.e., n = neurons). In all other studies, experimental data were averaged across animals (i.e., n = mice). Data are expressed as mean \pm standard error (SEM) and significance was set at p < 0.05. For differential gene expression analysis, significance was set at FDR < 0.10 and p < 0.05.

DATA AND SOFTWARE AVAILABILITY

All RNA-seq data are available in Tables S1 and S2; raw counts from RNA-sequencing are available on the Gene Expression Omnibus, GEO: GSE112196. Data that support the findings of this study are available from the corresponding authors upon reasonable request.

Supplemental Information

Chd2 Is Necessary

for Neural Circuit Development

and Long-Term Memory

Young J. Kim, Sattar Khoshkhoo, Jan C. Frankowski, Bingyao Zhu, Saad Abbasi, Sunyoung Lee, Ye Emily Wu, and Robert F. Hunt

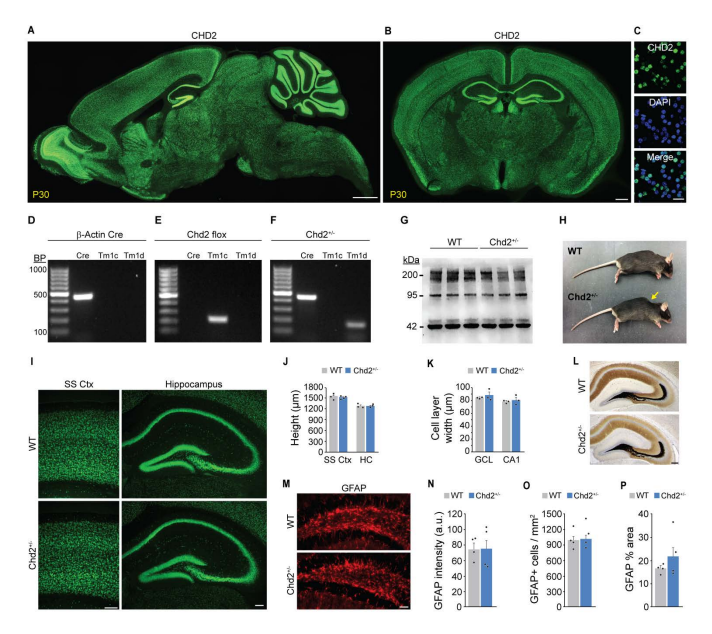


Figure S1. CHD2 expression in brain, related to Figure 1. A, B. Immunostaining for CHD2 (green) at P30 in a sagittal section (A) and coronal section (B) of wild-type mice. C. Immunostaining for CHD2 (green) and DAPI (blue) in mouse neocortex at P30. CHD2 co-localized in most, but not all, cell nuclei. D-F. Genotyping for *tm1c*, *tm1d* and *Cre* primer sets in a β-actin Cre mouse (D), *Chd2*-flox mouse (E) and *Chd2*+/- mouse (F). Only the *Chd2*+/- mutant contained the *tm1d* allele, indicating Cre-mediated recombination occurred in this animal. G. Whole western blot for the image shown in Figure 1D. H. Two month old male WT and *Chd2*+/- littermates. *Chd2*+/- mice exhibited prominent curvature of the spine (yellow arrow). I. NEUN immunostaining (green) of somatosensory cortex (left panels) and hippocampus (right panels) at P30. J. Height of somatosensory cortex (SS Ctx) or hippocampus (HC) was not altered by *Chd2*+/- (SS Ctx: P= 0.24, n=3 WT, n=4 *Chd2*+/-; HC: P= 0.98; n=3 mice per genotype). K. Width of granule cell layer (GCL) or CA1 pyramidal cell layer (CA1) was not altered by *Chd2*+/- (GCL: P= 0.51; CA1: P= 0.68; n=3 mice per genotype). L. Timm's stain (brown) and Nissl counter stain (blue) did not reveal an overt change in mossy fiber projections between WT and *Chd2*+/- mice (n = 3 mice per genotype). M. Immunostaining for GFAP (red) at P30 in WT and *Chd2*+/- mice (n=4 mice per genotype). N-P. Quantification of GFAP immunostaining. Error bars, s.e.m.; scale bars, 500 μm in A and B, 50 μm in C and M, 150 μm in I and 200 μm in J.

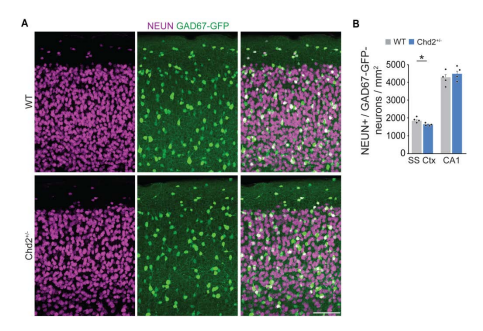


Figure S2. Density of putative excitatory neurons are reduced in $Chd2^{+/-}$ mice, related to Figure 2. A. Immunostaining for NEUN (magenta) and GAD67-GFP (green) in somtatosensory neocortex in WT (top panels) and $Chd2^{+/-}$ mice (bottom panels). **B.** Quantification of NEUN-positive/GAD67-GFP-negative cell density in SS Ctx and in CA1 pyramidal cell layer (n=4 mice per genotype). Error bars, s.e.m.; * p < 0.05; scale bar, 100 μ m.

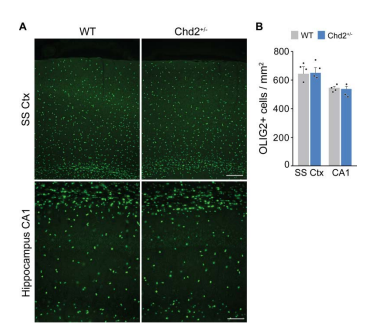


Figure S3. Density of OLIG2+ cells is not altered in $Chd2^{+/-}$ mice, related to Figure 2. A. Immunostaining for OLIG2 in somtatosensory neocortex (SS Ctx, top panels) and hippocampus CA1 (bottom panels) in WT and $Chd2^{+/-}$ mice. B. Quantification of OLIG2+ cell density in SS Ctx and CA1 (n=3-4 mice per genotype). Error bars, s.e.m.; scale bars, 150 μ m in SS Ctx and 75 μ m in Hippocampus CA1.

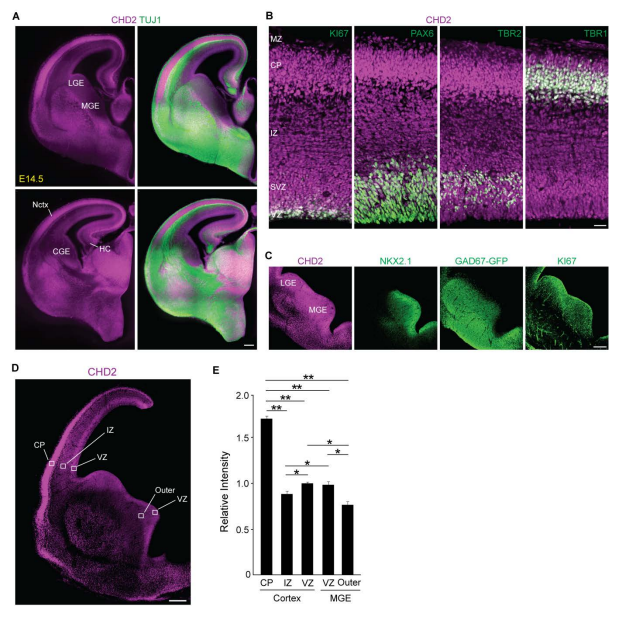


Figure S4. CHD2 expression in E14.5 brain, related to Figure 3. A. Immunostaining for CHD2 (magenta) and neuronal TUJ1 (green) in two sections of embryonic brain. B. In dorsal forebrain, immunostaining for CHD2 (magenta) and KI67, PAX6, TBR2 and TBR1 (all in green). C. In ventral forebrain, immunostaining for CHD2 (magenta) and NKX2.1, GAD67-GFP, and KI67 (all in green). D. Immunostaining for CHD2 at E14.5 E. Quantification of relative CHD2 intensity by subregion of the embryo. Nctx: Neocortex; HC: Hippocampus, LGE: lateral ganglionic eminence; MGE: medial ganglionic eminence; CGE: caudal ganglionic eminence; MZ: marginal zone; CP: cortical plate; IZ: intermediate zone; SVZ: subventricular zone; VZ: ventricular zone. Error bars, s.e.m.; * p < 0.05, ** p < 0.01; scale bars, 200 μm in A, 50 μm in B and 200 μm in C and D.

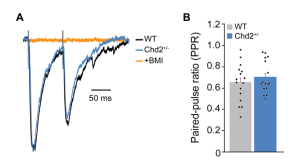


Figure S5. Pre-synaptic GABA release is not altered in $Chd2^{+/-}$ mice, related to Figure 5. A. Representative responses of CA1 pyramidal neurons to paired stimulation of the stratum radiatum in WT (black) and $Chd2^{+/-}$ (blue). Recordings were performed in the presence of 1mM kynurenic acid to block glutamate receptors. Response of the same WT recording in the presence of 30μ M bicuculline methiodine is also shown (orange). B. Quantification of PPRs shows no difference between WT and $Chd2^{+/-}$ littermates (WT: 0.65 ± 0.04 , n= 16 cells from 3 mice, $Chd2^{+/-}$: 0.71 ± 0.04 , n= 16 cells from 3 mice, P=0.39, two-tailed t-test). Error bars, s.e.m.

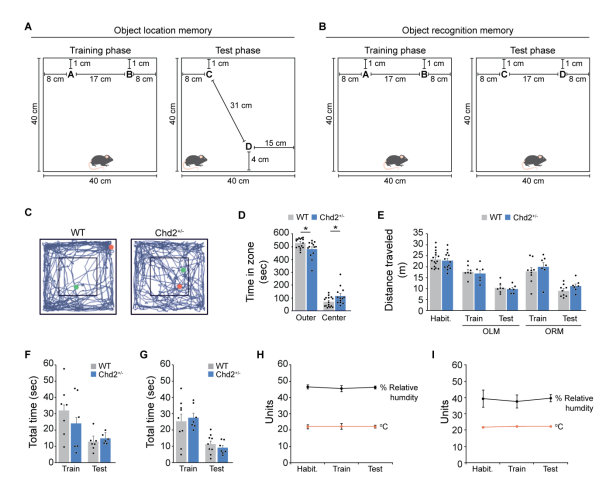


Figure S6. Object location and recognition memory tasks and experimental design, related to Figure 7. A, B. Schematics showing measurements for the context and object placement in Object Location Memory (OLM) and Object Recognition Memory (ORM) assays. **C.** Tracking plot of the path taken by a WT and $Chd2^{+/-}$ mouse during the entire 10 minute Open Field Test (OFT) period. Center region is outlined by the inner square. Green dot indicates starting position and red dot indicates stop. **D.** Time spent in the outer and center of the OFT apparatus (n=13-16 mice per genotype). **E.** Total distance traveled in during each phase of testing (n=13-16 mice per genotype). **F, G.** Total time spent exploring the objects during the each phase of the OLM assay (n=7 WT, n=6 $Chd2^{+/-}$) (**F**) and ORM assay (n=9 WT, n=7 $Chd2^{+/-}$) (**G**) . **H, I.** Relative humidity (%) and temperature (°C) during each phase of the OLM assay (**H**) and ORM assay (**I**). Error bars, s.e.m.; * p < 0.05.

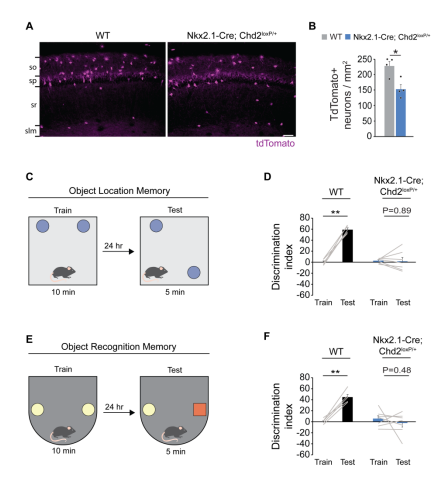


Figure S7. Mice with *Chd2* haploinsufficiency only in inhibitory interneurons have memory deficits, related to Figure 7. A. Immunostaining for tdTomato (magenta) in hippocampus CA1 of WT and *Nkx2.1*-Cre; *Chd2*^{loxP/+} mice. Abbreviations: so, stratum oriens; sp, stratum pyramidale; sr, stratum radiatum; slm, stratum lacunosum-moleculare **B.** Quantification of tdTomato+ cell density in WT and *Chd2*^{+/-} mice (n=4 mice per genotype). **C.** Schematic of OLM assay. **D.** Discrimination index during training and testing phases of OLM (n=5 WT, n=7 *Nkx2.1*-Cre; *Chd2*^{loxP/+}). **E.** Schematic of ORM assay. **F.** Discrimination index during training and testing phases of ORM (n=5 WT, n=7 *Nkx2.1*-Cre; *Chd2*^{loxP/+}). Error bars, s.e.m.; * p < 0.05, ** p < 0.01; scale bar, 50 μm.

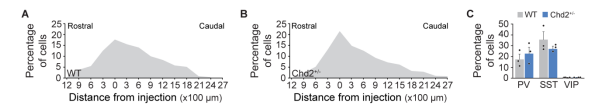


Figure S8. Characterization of MGE transplantations, related to Figure 8. A, B. Distribution of transplanted MGE cells expressing GFP 45 DAT into WT (**A**) and $Chd2^{+/-}$ mice (**B**) (n= 3 mice per genotype). **C.** Quantification of interneuron marker expression in GFP-labeled cells (n=3 mice per genotype). Error bars, s.e.m.

Table S4. Differentially expressed synaptic plasticity genes in P45 hippocampus, related to Figure 4

Cellular domain	Gene	Description	baseMean	log2FoldChange	IfcSE	stat	pvalue	padj
Presynaptic proteins	Trim9	E3 ubiquitin ligase	9878.0758	-0.297617138	0.060249	-4.93977	7.82E-07	0.000143
	Caskin1	Adaptor protein	8660.9239	-0.326542884	0.070994	-4.59957	4.23E-06	0.000581
	Ncs1	Calcium sensor	13869.236	-0.236470481	0.057108	-4.14079	3.46E-05	0.003259
	Snph	Synaptic vesicle protein	16771.881	-0.296922917	0.073095	-4.06216	4.86E-05	0.004272
	Stxbp3a	Synaptic vesicle protein	556.96647	0.422053648	0.109217	3.864357	0.000111	0.007569
	Epha5	Receptor tyr kinase	4788.0206	0.308838248	0.087976	3.510497	0.000447	0.021129
	Rims4	Synaptic vesicle protein	491.71387	-0.386983206	0.118691	-3.26044	0.001112	0.040598
	Map1a	Microtubule-assoc protein	52381.892	-0.235485447	0.074369	-3.16646	0.001543	0.051325
	Epn1	Accessory protein	14793.831	-0.206113977	0.06697	-3.0777	0.002086	0.062486
	Bsn	Cytomatrix protein	20288.517	-0.3015999	0.098672	-3.0566	0.002239	0.066231
	Cacnb1	L-VGCC	6046.6011	-0.19026523	0.06255	-3.04179	0.002352	0.06859
	Nos1ap	Adaptor protein	4164.6464	-0.201999846	0.070247	-2.87557	0.004033	0.093233
	Cnrip1	Cannabinoid receptor	2259.5062	0.222272497	0.078268	2.839899	0.004513	0.099877
Transporters	Slc6a13	GABA transporter	128.76955	1.079248407	0.291063	3.707956	0.000209	0.012216
	Glul	Glutamine synthatase	41533.456	-0.232575348	0.067694	-3.43569	0.000591	0.02565
	Slc6a1	GABA transporter	12452.841	-0.209124539	0.069143	-3.02451	0.00249	0.071906
Neurotransmiters and	Htr1a	Serotonin receptor	1355.6582	-0.682404756	0.120615	-5.65771	1.53E-08	5.26E-06
peptide receptors	Sstr4	SST receptor	1491.9177	-0.44376754	0.082167	-5.40082	6.63E-08	1.80E-05
	Grm3	mGlu receptor	1166.7208	0.387702432	0.0976	3.972356	7.12E-05	0.00552
	Htr5b	Serotonin receptor	241.97736	-0.591174889	0.154531	-3.8256	0.00013	0.008505
	Adra2c	Adrenergic receptor	1453.4627	-0.337620838	0.089403	-3.77638	0.000159	0.009882
	Adra2a	Adrenergic receptor	382.96341	-0.513859802	0.141234	-3.63835	0.000274	0.014642
	Gabrg2	GABAA receptor	4832.4349	0.286843503	0.087874	3.264262	0.001097	0.040236
	Adra1d	Adrenergic receptor	4206.8036	-0.28301214	0.089959	-3.14601	0.001655	0.053896
	Vipr1	VIP receptor	811.1855	-0.288108864	0.095605	-3.01353	0.002582	0.072902
	Grik5	Kainate receptor	11781.728	-0.217421338	0.072266	-3.00864	0.002624	0.073342
	Vipr2	VIP receptor	21.244918	1.480979896	0.516428	2.86774	0.004134	0.094547
	Grina	NMDA receptor	16333.148	-0.163432572	0.057141	-2.86018	0.004234	0.095766

Postsynaptic proteins	Synpo	Actin-associated protein	7486.7657	-0.265907101	0.062478	-4.25602	2.08E-05	0.002205
	Kcnj4	Inward rect. potassium channel	2545.1107	0.366215527	0.091037	4.022726	5.75E-05	0.004791
	Nptx1	Neuronal pentraxin	27241.162	-0.277001789	0.069611	-3.97928	6.91E-05	0.005472
	Dlgap4	Guanylate kinase	9285.0882	-0.248774457	0.062707	-3.96724	7.27E-05	0.005545
	Sdc4	Cell surface proteoglycan	1806.3254	-0.30958312	0.078192	-3.95927	7.52E-05	0.005671
	Homer1	PSD scaffold	3348.8022	-0.382856923	0.097512	-3.92624	8.63E-05	0.006318
	Slitrk2	Integral membrane protein	955.87629	0.389953905	0.099275	3.928018	8.56E-05	0.006318
	Neurl1a	E3 ubiquitin ligase	8815.3489	-0.274762814	0.072054	-3.81331	0.000137	0.00882
	Ncald	Calcium sensor	15079.737	0.225981212	0.061546	3.671759	0.000241	0.013411
	Ctnnd2	Adhesive junction protein	6450.3146	-0.243849179	0.068653	-3.55193	0.000382	0.018855
	Dapk1	Ca2+/calmodulin-dep kinase	6285.0713	-0.342000179	0.097872	-3.49435	0.000475	0.02202
	Shank2	PSD scaffold	6031.9254	-0.292767867	0.085053	-3.44219	0.000577	0.025194
	Jph4	Junctional complex protein	21437.564	-0.212417659	0.06292	-3.37602	0.000735	0.030102
	Farp1	Rho/ArhGEF protein	2007.4847	-0.293744956	0.087318	-3.3641	0.000768	0.03108
	Ap2b1	Adaptor protein	15474.326	-0.207002967	0.06339	-3.26557	0.001092	0.040236
	Sdcbp	Adaptor protein	6974.9246	0.223613741	0.069269	3.228196	0.001246	0.044444
	Dlgap3	Guanylate kinase	14731.433	-0.214398761	0.068118	-3.14745	0.001647	0.05381
	Rps6ka2	Protein kinase	1804.8713	-0.227100184	0.075376	-3.01289	0.002588	0.072902
	Bai1	Adhesion-GPCR	19742.357	-0.200291765	0.066973	-2.99065	0.002784	0.075537
	Prkcg	Protein kinase	28000.929	-0.214661078	0.071905	-2.98532	0.002833	0.076147
	Mdga1	Glycoprotein	3799.5167	-0.322846088	0.108329	-2.98022	0.00288	0.077139
	Ppfibp2	Protein-tyrosine phosphatase	287.38425	0.493817131	0.169158	2.919273	0.003508	0.086856
	Baiap2	Adaptor protein	14401.212	-0.234383326	0.080664	-2.90567	0.003665	0.088
	Nlgn3	Neuroligin	2939.9086	-0.225606427	0.078374	-2.87859	0.003995	0.092531
	Nos1ap	Adaptor protein	4164.6464	-0.201999846	0.070247	-2.87557	0.004033	0.093233
Ion channels	Kcna4	A-VGKC	963.4687	0.743731603	0.103635	7.176449	7.15E-13	1.03E-09
	Cacna1i	T-VGCC	2974.1655	-0.566857707	0.111742	-5.07294	3.92E-07	8.42E-05
	Cacna1g	T-VGCC	2230.5431	-0.515222895	0.112826	-4.56652	4.96E-06	0.000655
	Kcnv1	V-VGKC	1368.9607	0.389275901	0.089412	4.353709	1.34E-05	0.001461
	Kcnj4	Inward rectifying K+ channel	2545.1107	0.366215527	0.091037	4.022726	5.75E-05	0.004791
	Cacnb1	L-VGCC	6046.6011	-0.19026523	0.06255	-3.04179	0.002352	0.06859

Immediate-early								
respoonse	Ptgs2	Inducible cyclooxygenase	941.65501	0.698005372	0.114525	6.094801	1.10E-09	6.32E-07
	Fosb	bZIP protein; AP-1 comp	377.72767	-0.742871317	0.143018	-5.19425	2.06E-07	5.02E-05
	Irs2	Insulin receptor	2516.7345	-0.465831764	0.098665	-4.72134	2.34E-06	0.000379
	Xbp1	Transcription factor	2194.3227	-0.465420752	0.099447	-4.68008	2.87E-06	0.000426
	Fosl2	Transcription factor	1308.1252	-0.553990218	0.119239	-4.64606	3.38E-06	0.000491
	Nr4a2	Nuclear receptor	1871.2012	-0.487850093	0.110374	-4.41997	9.87E-06	0.001147
	Srf	Transcription factor	3312.4911	-0.25607372	0.069949	-3.66088	0.000251	0.013822
	Ntf3	Neurotrophic factor	272.35458	0.573812056	0.173484	3.307583	0.000941	0.036448
	Egr3	Transcription factor	4337.7776	-0.38582253	0.117391	-3.28663	0.001014	0.038044
	Klf4	Transcription factor	112.67039	-0.733373112	0.231522	-3.16762	0.001537	0.051321
LTP/LTD	Ppp1r14a	Protein phosphotase	718.64766	0.62244396	0.148952	4.178834	2.93E-05	0.002944
	Crtc1	Transcription factor	11689.59	-0.239543022	0.072022	-3.32595	0.000881	0.034784
	Prkcg	Protein kinase	28000.929	-0.214661078	0.071905	-2.98532	0.002833	0.076147
Extracellular matrix	C1ql2	Complement protein	3843.0189	0.435208914	0.086265	5.04502	4.53E-07	9.29E-05
	Mmp14	Matrix metalloproteinase	520.20192	0.531655053	0.112275	4.735283	2.19E-06	0.000358
	Cdh4	Cadherin	384.64011	-0.596291704	0.139207	-4.28348	1.84E-05	0.001977
	Pcdh17	Protocadherin	3183.0857	-0.255019461	0.068645	-3.71505	0.000203	0.011996
	Cdh5	Cadherin	284.00028	-0.570990198	0.154212	-3.70262	0.000213	0.012357
	Pcdh9	Protocadherin	3588.3639	0.268392374	0.078583	3.415407	0.000637	0.026918
	Tnr	Glycoprotein	3965.5614	-0.302667156	0.091959	-3.29133	0.000997	0.03761
	Pcdh19	Protocadherin	2608.9379	-0.264672254	0.087786	-3.01498	0.00257	0.072889
	Cdhr3	Cadherin	136.55122	0.771942617	0.266725	2.894154	0.003802	0.090008
	Reln	Glycoprotein	2444.9309	-0.205371212	0.072329	-2.83941	0.00452	0.099877
Other, related	Dab2ip	Ras GAP	4567.9536	-0.381692402	0.075143	-5.07955	3.78E-07	8.26E-05
	Dbi	Diazepam binging protein	2437.8684	0.479412768	0.109691	4.370574	1.24E-05	0.001373
	Eif4ebp2	Transcription factor	748.50468	-0.396431535	0.101243	-3.91564	9.02E-05	0.006465
	Panx2	Innexin, gap junction	5255.58	-0.2652065	0.07085	-3.74322	0.000182	0.010907
	Celsr2	Cadherin	11405.93	-0.258293433	0.083587	-3.0901	0.002001	0.060789

Table S6. Electrophysiological properties of CA1 pyramidal neurons, related to Figure 5.

Electrophysiological property	WT	Chd2+/-	U-statistic	P-value
Resting potential (mV)	-63.538 ± 0.627	-64.143 ± 0.543	U=74.5	0.432
Input resistance (M Ω)	137.846 ± 9.943	146.250 ± 8.446	U=77.0	0.512
Time constant, tau (ms)	22.106 ± 1.542	20.429 ± 0.924	U=70.00	0.32
Spike threshold (mV)	-43.847 ± 0.717	-44.996 ± 0.368	U=62.00	0.167
Spike amplitude (mV)	96.199 ± 2.286	92.808 ± 3.237	U=82.00	0.68
Spike half-width (ms)	1.157 ± 0.0642	1.044 ± 0.03	U=65.00	0.215
AHP amplitude (mV)	10.002 ± 0.694	9.825 ± 0.700	U=91	1
Spike Adaptation at 2x threshold	1.609 ± 0.126	1.303 ± 0.0651	U=46.00	0.031
Spike Acomodation at 2x threshold (mV)	11.945 ± 1.704	10.993 ± 1.424	U=89.00	0.942
Sag amplitude (mV)	3.296 ± 0.264	2.879 ± 0.168	U=70.00	0.32