Altered Neocortical Gene Expression, Brain Overgrowth and Functional Over-Connectivity in Chd8 Haploinsufficient Mice

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Abstract
Truncating CHD8 mutations are amongst the highest confidence risk factors for autism spectrum disorder (ASD) identified to date. Here, we report that Chd8 heterozygous mice display increased brain size, motor delay, hypertelorism, pronounced hypoactivity, and anomalous responses to social stimuli. Whereas gene expression in the neocortex is only mildly affected at midgestation, over 600 genes are differentially expressed in the early postnatal neocortex. Genes involved in cell adhesion and axon guidance are particularly prominent amongst the downregulated transcripts. Resting-state functional MRI identified increased synchronized activity in cortico-hippocampal and auditory-parietal networks in Chd8 heterozygous mutant mice, implicating altered connectivity as a potential mechanism underlying the behavioral phenotypes. Together, these data suggest that altered brain growth and diminished expression of important neurodevelopmental genes that
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regulate long-range brain wiring are followed by distinctive anomalies in functional brain connectivity in Chd8−/− mice. Human imaging studies have reported altered functional connectivity in ASD patients, with long-range under-connectivity seemingly more frequent. Our data suggest that CHD8 haploinsufficiency represents a specific subtype of ASD where neuropsychiatric symptoms are underpinned by long-range over-connectivity.

Key words: ASD, autism, axon guidance, behavior, CHD8, chromatin remodelling, cortex, functional connectivity, gene expression, macrocephaly, mouse

Introduction

Autism spectrum disorder (ASD) is diagnosed on the basis of socio-communicative deficits and repetitive, perseverative behaviors with restricted interests (APA 2013). ASD is frequently associated with comorbidities like hypersensitivity to sensory stimuli, seizures, and anxiety (Tavassoli et al. 2014; Croen et al. 2015; Jeste and Tuchman 2015). The phenotypic and genetic heterogeneity of ASD has hampered the elucidation of the molecular mechanisms that may underlie specific behavioral symptoms. However, the recent identification of de novo, likely gene disrupting (LGD) mutations that show highly significant associations with autism (Neale et al. 2012; Talkowski et al. 2012b; O’Roak et al. 2012b; O’Roak et al. 2014) provides an opportunity to phenotype and molecularly characterize genetically defined ASD subtypes. Exome sequencing studies of several thousand simplex families detected de novo, LGD mutations in the CHD8 (Chromodomain Helicase DNA binding factor 8) gene (Neale et al. 2012; Talkowski et al. 2012; O’Roak et al. 2012b; lossiof et al. 2014; O’Roak et al. 2014). Patients with CHD8 mutations are characterized by a high incidence of autism, macrocephaly, facial dysmorphism, motor delay and hypotonia, intellectual disability, and gastrointestinal problems (Bernier et al. 2007; Stierman et al. 2016; Stessman et al. 2017). CHD8 encodes an ATP-dependent chromatin remodeling protein of the chromodomain helicase DNA binding family (Yuan et al. 2007; Thompson et al. 2008). The recruitment of CHD8 to gene promoters in mouse and human neural progenitors is strongly associated with transcriptional activation, while CHD8 knock-down in these cells results in the reduced expression of many ASD-associated genes (Sugathan et al. 2014; Cotney et al. 2015). Three groups recently described Chd8−/− mouse models (Katayama et al. 2016; Gompers et al. 2017; Platt et al. 2017). Megalencephaly, subtle but wide-spread transcriptional changes and behavioral anomalies were found in all these Chd8−/− mouse lines. Individual studies reported attenuated expression of neural genes and derepression of REST (Katayama et al. 2016), alterations in striatal neurotransmission (Platt et al. 2017) and a developmental RNA splicing phenotype (Gompers et al. 2017). Understanding the contribution of each of these mechanisms to the ASD phenotype remains a major challenge.

Altered brain connectivity, characterized by local over-connectivity and long-range under-connectivity, has been hypothesized to underpin some of the neuropsychiatric phenotypes observed in ASD (Belmonte et al. 2004; Just et al. 2004). Resting-state functional MRI (rsfMRI) studies in ASD patients have provided evidence for reduced long-range synchronization in spontaneous brain activity (reviewed in Picci et al. 2016). Increased long-range connectivity has also been reported in a subset of cases (Di Martino et al. 2014), consistent with the phenotypic heterogeneity of ASD. Thus, the exact nature of aberrant functional connectivity in ASD may depend on the specific underlying aetiology.

Similar rsfMRI studies in ASD mouse models may help bridge the gap between ASD models and the human condition (Liska and Gozzi 2016). As one example, homozygous Ctnnap2 mouse mutants exhibit hyperconnectivity of the default mode network (Liska et al. 2017), a phenotype often observed in idio-pathic ASD patients (Cherkassky et al. 2006) and recapitulating analogous clinical observations in humans with CNTNAP2 mutations (Scott-Van Zeeland et al. 2010).

In the present study we generated a novel Chd8−/− mouse model. We report behavioral anomalies, macrocephaly and functional over-connectivity in cortico-hippocampal networks in these mice that are prefigured by dysregulation of the cortical transcriptome in the early postnatal period.

Methods

Chd8 Gene Targeting

A 14.84 kb genomic DNA fragment was subcloned from C57BL/6 BAC clone (RP23: 318M20) into pSP72 (Promega). This fragment encompassed a 9.45 kb 5’ long homology arm (LA) and a 4.4 kb 3’ short homology arm (SA). The targeting construct was generated by inserting a loxP/FRT-PGK-gb2-Neo cassette 214 bp 3’ of exon 3 (inGenious Targeting Laboratory [iTL], Ronkonkoma, NY, USA). An additional single loxP site containing a BclI restriction site for Southern blot screening was inserted 5’ of exon 3. The final targeting construct of 18.8 kb was linearized by NotI digestion and electroporated into C57BL/6 ES cells. G418-resistant clones were selected, screened by PCR and Southern blot for successful homologous recombination. Five clones with successful recombination were identified (Supplementary Fig. S1) and 2 clones (124 and 254) were injected into Balb/c blastocysts (iTL). Resulting chimaeras were bred with Flpe deleter mice on a C57BL/6 background to excise the neo cassette and produce Chd8fl/fl−/− mice (Supplementary Fig. S1). Chd8fl/fl−/− mice were then crossed with ß-actinCre mice (Lewandoski and Martin 1997) to generate a Chd8 null allele (Chd8−/−). β-actinCre;Chd8−/− mice were crossed with C57BL/6J mice to remove the Cre transgene and establish a Chd8−/− line.

Mice

Experimental mice were produced by Chd8−/− × C57BL/6J crosses, taking care to equalize paternal or maternal inheritance of the Chd8 null allele, especially for behavioral experiments. For genotyping, genomic DNA was extracted using Proteinase K digestion or the HotSHOT method (Truett et al. 2000). Genotyping reactions were then performed for the presence of Chd8 wildtype and null alleles using the following primer pair: FW: CCC ACA TCA AGT long homology arm (LA) and a 4.4 kb 3’ short homology arm (SA). The targeting construct was generated by inserting a loxP/FRT-PGK-gb2-Neo cassette 214 bp 3’ of exon 3 (inGenious Targeting Laboratory [iTL], Ronkonkoma, NY, USA). An additional single loxP site containing a BclI restriction site for Southern blot screening was inserted 5’ of exon 3. The final targeting construct of 18.8 kb was linearized by NotI digestion and electroporated into C57BL/6 ES cells. G418-resistant clones were selected, screened by PCR and Southern blot for successful homologous recombination. Five clones with successful recombination were identified (Supplementary Fig. S1) and 2 clones (124 and 254) were injected into Balb/c blastocysts (iTL). Resulting chimaeras were bred with Flpe deleter mice on a C57BL/6J background to excise the neo cassette and produce Chd8fl/fl−/− mice (Supplementary Fig. S1). Chd8fl/fl−/− mice were then crossed with ß-actinCre mice (Lewandoski and Martin 1997) to generate a Chd8 null allele (Chd8−/−). β-actinCre;Chd8−/− mice were crossed with C57BL/6J mice to remove the Cre transgene and establish a Chd8−/− line.

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Western Blot

Telencephalic vessels were dissected from E12.5 embryos and total cell culture preparations were lysed in 8 M urea, 1% CHAPS, 50 mM Tris (pH 7.9) containing protease inhibitors (PMSF, Pepstatin A, Leupeptin, Aprotinin, Roche) and a phosphatase inhibitor cocktail (Sigma). Samples were loaded (10 μg total protein per lane) onto a Mini-PROTEAN precast gel (Bio-Rad) and resolved using gel electrophoresis. Protein was transferred to a nitrocellulose membrane (Bio-Rad) which was then blocked in 5% nonfat milk powder (Bio-Rad) and 1% bovine serum albumin (BSA, Sigma) in TBS with 0.1% Tween-20 (TBST), followed by incubation with anti-Chd8 primary antibody (rabbit anti-Chd8 N-terminal, Bethyl Laboratories (cat#: A301-224 A), 1:5000) in 3% nonfat milk powder and 1% BSA in TBST overnight at 4 °C. After washing, the membrane was incubated with HRP-conjugated secondary antibody (Millipore), HRP detected with Clarity ECL reagent (Bio-Rad) and the membrane imaged using a Bio-Rad ChemiDoc system. The same membrane was subsequently incubated with anti-GAPDH primary antibody (rabbit anti-GAPDH, Abcam (cat#: ab9485), 1:40000) overnight at 4 °C and probed with HRP-conjugate and imaged as before. Relative protein quantity was calculated using Bio-Rad ImageLab software.

X-ray Computed Tomography

Fixed heads from adult (26–27 days old) Chd8−/− and Chd8+/+ mice (n = 7 of each from 2 different litters) were scanned using a GE Locus SP microCT scanner. The specimens were immobilized using cotton gauze and scanned to produce 28 μm voxel size volumes, using a X-ray tube voltage of 80kVp and a tube current of 80mA. An aluminum filter (0.05 mm) was used to adjust the energy distribution of the X-ray source. Reconstructions of computer tomography scans, images and measurements were done in MicroView 2.5.0 software (Parallax Innovations, ON, Canada). Each 3D landmark point was recorded, twice for each sample, using the 3D point recording built-in tool within the same software, with the operator blind to the genotypes. The distances between the landmarks were normalized for each sample to the average of the wild-type littermates. Graphics of the plotted data and statistical analysis were performed using GraphPad Prism version 6.0 h for Mac OS X (GraphPad Software, La Jolla, CA USA, www.graphpad.com). Unpaired Student t-tests were applied to analyze the variation between the 2 groups, for every distance between 2 specific 3D landmark points. 3D coordinate locations of a total of 22 biological relevant cranial landmarks were chosen based on a landmark list for adult mouse skull (Hill et al. 2009).

Behavioral Assessments

Mice for behavioral testing were housed, marked for identification and behaviors assessed essentially as described in Whittaker et al. (2017a). Different batches of mice were used for (1) recording pup USVs and spontaneous motor behaviors, and (2) adult behaviors (9–12 weeks of age at the start of testing; 19–22 weeks of age at the end of testing). For adult behaviors, tests were carried out in the following order: rotarod, grip strength, open field, self-grooming, marble burying, adult social investigation, 3 chamber social approach, light/dark test, olfactory habituation/dishabituation and Morris water maze.

General Activity Measurements

General activity was measured using a running wheel paradigm. Mice were housed individually under a 12 h:12 h light-dark cycle (lights on at 8 am; lights off at 8 pm) in a light-, air-, temperature-controlled ventilated cabinet (Arrowmigh, Hereford, UK). Running-wheel cages were equipped with an infrared sensor (Bilaney consultant Ltd, Sevenoaks, UK) connected to a computer. Data were collected in 1-min bins using ClockLab software (Actimetrics, Inc, Wilmette, IL, USA). Mice were continuously monitored undisturbed from the day they were placed in the running wheel cages and their general activity during the light versus dark phase were compared over the first 7 days.

Structural MRI

After completion of adult behavioral tests, mice were terminally anesthetized and intracardially perfused. Samples were processed, imaged and analyzed as previously described (Whittaker et al. 2017b).

Resting-State fMRI

rsfMRI experiments were performed on 15–18 weeks old mice (n = 23 Chd8−/−; n = 19 Chd8+/+). Animals were prepared for imaging as previously described (Ferrari et al. 2012; Sforazzini et al. 2016). Briefly, mice were anesthetized using isoflurane (5% induction), intubated and artificially ventilated (2% maintenance). Blood pressure was monitored continuously by cannulating the left femoral artery, also allowing for terminal arterial blood sampling. Administration of isoflurane was ceased after surgery and substituted with halothane (0.75%). Functional data acquisition commenced 45 min after isoflurane cessation. To rule out possible genotype-dependent differences in anesthesiain sensitivity we continuously recorded 2 independent readouts previously shown to be linearly correlated with anesthesiain depth: arterial blood pressure and amplitude of cortical BOLD signal fluctuations (Steffey et al. 2003; Liu et al. 2011; Zhan et al. 2014). Arterial blood pressure (P = 0.79; Supplementary Fig. S5A) and the amplitude of BOLD signal fluctuations of motor cortex (P = 0.56; Supplementary Fig. S5B) did not significantly differ between Chd8−/− mice and littermate controls, eliminating a confounding contribution of anesthesia. In vivo images were obtained using a 7.0 T MRI scanner (Bruker Biospin, Milan), as previously described (Liska et al. 2017). Signal transmission and reception were achieved using a 72 mm birdcage transmit coil and a 4-channel solenoide coil. For each session, high-resolution anatomical images were acquired using a fast spin echo sequence based on the following parameters: repetition time (TR)/echo time (TE) 5500/60 ms, matrix 192 × 192, field of view 2 × 2 cm2, 24 coronal slices, and slice thickness 0.5 mm. Cocentred BOLD rsfMRI time series were acquired using an echo planar imaging (EPI) sequence with the following parameters: TR/TE 1200/15 ms, flip angle 30°, matrix 100 × 100, field of views 2 × 2 cm2, 24 coronal slices, slice thickness 0.5 mm, 500 volumes, and 10 min total acquisition time. Raw MRI data, templates, and code employed to generate functional maps are available by contacting AG.

Functional Connectivity Analyses

To allow for T1 equilibration effects, the first 20 volumes of rsfMRI data were removed. The time series were then despiked, corrected for motion and spatially normalized to an in-house mouse brain template (Sforazzini et al. 2014). Normalized data had a spatial resolution of 0.1042 × 0.1042 × 0.5 mm3 (192 × 192 × 24 matrix). Mean ventricular signal (averaged rsfMRI time course within a reference ventricular mask) and head motion traces
were regressed out of each time series. No genotype-dependent differences were observed in ventricular volume, as measured by the dimensions of individual ventricular masks. All rsfMRI time series were then spatially smoothed (full width at half maximum of 0.6 mm) and band-pass filtered using a frequency window of 0.01–0.1 Hz.

To identify brain regions displaying genotype-dependent differences in functional connectivity in an unbiased manner, we calculated global rsfMRI connectivity maps for all subjects, as described previously in detail (Liska et al. 2015, 2017). A previously described seed-based approach was then used to examine between-group differences in the intensity and scope of long-range rsfMRI correlation networks (Sforazzini et al. 2016).

**Tissue Collection and Processing**

Pups were weighed and sacrificed, while embryos were collected by dissection in ice-cold PBS, excess PBS drained and whole embryos weighed.Brains were then dissected from the skull in ice-cold PBS and cut below the brain stem, immediately drained on paper towels using a slotted spoon and wet weights determined using a fine scale. Brain weights were normalized to body weight and group differences were calculated using unpaired student’s t-test. Brains were postfixed in 4% PFA at 4°C for 24 h, dehydrated and paraffin embedded. Serial coronal sections were cut at 10 μm such that each slide contained 3 consecutive sections.

**Immunohistochemistry**

Sections were rehydrated using standard protocols and heated in 10 mM Sodium Citrate solution (pH 6). Endogenous peroxidases were blocked by incubating in 3% H2O2 and 10% MeOH in PBS for 15 min. Sections were permeabilised in 0.2% Triton X-100 (Sigma-Aldrich) in PBS (PBTr2) for 5 min and blocked using 10% heat-inactivated normal goat serum and 2% gelatin in PBTr2 for 1 h. Sections were incubated in 5% G5 in PBTr2 containing primary antibody (rabbit anti-phosphohistone 3B (Cell Signaling (#9701), 1/100) overnight at 4°C. After incubation with primary antibody, sections were incubated in biotinylated anti-rabbit immunoglobulin secondary antibody (Dako (#E0432), 1/ primary antibody, sections were incubated in biotinylated anti- (9701), 1/100) overnight at 4°C. After incubation with primary antibody, sections were incubated in biotinylated anti-rabbit immunoglobulin secondary antibody (Dako (#9701), 1/200) in 5% goat serum in PBTr2. Samples were washed in PBS and incubated with avidin/biotin complex (ABC, Vector) in PBS for 1 h. Sections were developed using 0.025% DAB and 0.03% H2O2 in PBS for 10 min, counterstaining using Ehrlich’s Hematoxylin solution and mounted in DPX (Sigma-Aldrich). Images were acquired on a Nikon 80i microscope equipped with a Nikon 5 M pixel Nikon DS digital camera. Images were processed using Adobe Photoshop and Illustrator.

**RNA Sequencing**

RNA was isolated from microdissected cortices at E12.5 (both hemispheres) and P5 (one hemisphere) and reverse transcribed (n = 3 per experimental group). cDNA was end-repaired, adaptor-ligated, and A-tailed. Paired-end sequencing was performed on the Illumina HiSeq 4000 platform. Quality of the raw sequencing data was checked using FastQC version 0.11.2 (Andrews 2010; available at: http://www.bioinformatics. babraham.ac.uk/projects/fastqc) and trimming of adaptor sequences was performed using Trim Galore! version 0.4.1 (Krueger 2012; available at: http://www.bioinformatics. babraham.ac.uk/projects/trim_galore/). Reads were aligned to the mouse genome (GRCm38.p4) using Tophat version 2.1.0 and aligned reads were counted using FeatureCounts version 1.5.0 (Kim et al. 2013; Liao et al. 2014). Differential expression testing was performed using DESeq2 version 1.10.1, as previously described (Love et al. 2014). Gene ontology analysis and functional classification was performed using DAVID with all detected DEGs below a 0.05 FDR (Huang et al. 2009). Putative regulatory transcription factors were determined with EnhancerDB using the “ENCODE and ChEA Consensus TFs from ChIP-X” database with all DEGs below a 0.05 FDR (Chen et al. 2013). The R package ggplot2 version 2.1.0 was used to generate volcano plots. The list of ASD-associated genes used for overlap with P5 DEGs was obtained from the SFARI Gene Scoring module (https://gene.sfari.org/autdb/HG_Home.do). RNA-seq data have been deposited into GEO, accession number GSE81103.

**Gene Expression Enrichment Analysis**

In total, 4345 gene expression images corresponding to 4082 unique genes were downloaded from the Allen Institute’s Mouse Brain Atlas coronal expression dataset (Lein et al. 2007). The coronal expression dataset is limited by the fact that it only partially covers the genome and was defined in a biased manner (Ng et al. 2009). However, it offers higher data quality and resolution especially in lateral cortical areas including auditory cortex, compared with the sagittal dataset. This dataset, obtained via situ hybridization, consisted of 3D spatial expression images aligned to a single reference model and summarized over the whole mouse brain at a 200 μm isotropic resolution. Specifically, the gene expression energies, defined by the Allen Institute as the sum of expressing pixel intensities divided by the sum of all pixels in a subdivision, were obtained. Expression data were obtained from adult (P56) male C57Bl/6.

Mean gene expression energies were extracted under a set of Allen Institute-defined segmentations, resulting in vectors of expression values for each gene that describe their spatial expression patterns. To account for differences in probe affinities that subsequently affect the total expression levels reported in the images, expression values for each gene were further normalized by dividing by the total expression (summed over regions) for that gene.

Next, we extracted rsfMRI time series data under the aforementioned set of segmentations. This was achieved by aligning, via ANTS (Avants et al. 2009; available at: http://hdl.handle.net/10380/3113), the Allen Institute’s average 2-photon microscopy
Methods

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We focused on the pair of regions that showed strongest over-connectivity under these parcellations and determined candidate genes by choosing those that had the highest normalized expression in both regions, compared with all other genes. Candidates that were in the top 20% of genes for both regions were passed through an enrichment analysis via the GoRilla tool (Eden et al. 2009; http://cbl-gorilla.cs.technion.ac.il/). Another set of candidates were obtained by ranking each gene by the sum of their normalized expression values in both regions (equivalent to the L1 distance); this set also favors genes with high expression in one of the 2 regions, along with both regions. For enrichment analyses of ranked lists of genes, GoRilla automatically thresholds the list independently for each GO term so that the optimal enrichment is found (Eden et al. 2009).

To establish specificity of the enriched gene ontology terms, we performed equivalent analyses between 10 randomly chosen regional pairs (DG, PAL, P-sat, MY-sat-DG, PAL-GU, P-sat-LZ, PAAS-VERM, ANO-HEM, MY-mot-MB). The number of returned GO terms was generally low and did not show an enrichment for neuronal development terms.

Statistical Analysis

Data are reported as mean ± standard error of the Mean (SEM) and graphs show all individual data points where feasible. Significant P-values are reported in the results section and figure legends provide full details of all relevant statistical parameters including group sizes. Statistical analyses were performed either with SPSS (Version 22, IBM, Armonk, USA) or GraphPad Prism (Version 6, GraphPad Software, La Jolla, CA, USA). All analyses were performed blind to genotype.

Behavior

Data were analyzed using either a between-subjects ANOVA or a 2-way repeated measures ANOVA, as appropriate. If there was no statistically significant sex difference, data were pooled. When the appropriate ANOVA showed a significant effect for a particular task, student’s t-tests were used as post hoc analyses, as there were only 2 groups for comparison. Cohort details can be found in the methods, group sizes are stated in the figure legend.

Proliferation

Phosphohistone 3B-positive cells lining the ventricular surface of the dorsal cortex were counted and normalized to the length of ventricular surface. These were quantified on both sides of the brain in 3 consecutive sections and averaged to calculate the number of phosphohistone 3B-positive cells per μm of ventricular surface in the dorsal cortex. Group differences were calculated using unpaired student’s t-test.

μCT Analysis

Each 3D landmark point was recorded twice for each sample and distances between landmark points normalized to the average of the wildtype controls. Group differences for distances between 2 specific 3D landmark points were calculated using unpaired student t-test.

MRI Analyses

Processing of raw data is described in detail in the relevant method sections. For structural MRI, significant differences were determined between groups for the 159 different regions in the brain. Voxelwise comparisons were made between mutants and littermate controls, group differences were calculated using unpaired student’s t-test and multiple comparisons were controlled for using a false discovery rate (FDR < 0.15). Exact P-values can be found in Supplementary Table S1.

For rsfMRI studies, group-level differences in connectivity distributions were calculated using 2-tailed student’s t-test (P < 0.05, family-wise error cluster-corrected, with cluster-defining threshold of t ≥ 2.06, P < 0.05) and multiple comparisons were controlled for using an FDR < 0.05.

RNA-seq

Processing of raw data and differential expression testing is described in Methods. Multiple comparisons were controlled for using an FDR < 0.05. Exact P-values and FDR adjusted P-values for all differentially expressed genes (DEGs) are listed in Supplementary Tables S2 and S3.

Results

A mouse line with a conditional Chd8 allele was produced through homologous recombination in C57Bl/6 embryonic stem cells (Supplementary Fig. S1A, B). Chd8fl/+ mice were crossed with the ubiquitously expressing β-actinCre line (Lewandoski and Martin 1997) to generate Chd8fl/+ mice (Supplementary Fig. S1C). Cre-mediated deletion of loxP-flanked (flox) exon 3 results in an early frameshift and termination of translation at amino acid 419, predicted to produce a protein that lacks all functional domains, equivalent to nonsense and frameshift mutations terminating CHD8 at amino acids 62 and 747 in patients (Barnard et al. 2015).

Quantitative RT-PCR (qRT-PCR) on RNA isolated from E12.5 and P5 neocortices using primers spanning the exon 3/4 boundary showed Chd8 expression reduced by 64% (P = 0.006) and 52% (P = 0.01), respectively (Supplementary Fig. S1D). CHD8 protein levels were reduced by 51% in Chd8fl/fl mice compared with controls (Supplementary Fig. S1E, F), validating our Chd8fl/+ mice as a suitable model for CHD8 haplinsufficiency. Importantly, we found no evidence for a truncated protein product of 419aa (45 kDa) that may have resulted from translation of any mutant transcript (Supplementary Fig. S1E). qRT-PCR analysis at E12.5 with primers spanning the exon 1/2 boundary (upstream of the recombination event) revealed reduced Chd8 expression of 52% (Supplementary Fig. S1G), indicating that the mutant transcript is most likely subject to nonsense-mediated decay.

Chd8 Heterozygous Mice Have Specific Craniofacial and Structural Brain Phenotypes

Humans with truncating mutations in a single CHD8 allele often present with macrocephaly (64%) and distinct craniofacial phenotypes (89%), which include hypertelorism (wide-set eyes, 67%) (Bernier et al. 2014; Stessman et al. 2017). We characterized the cranioskeleton of Chd8fl/+ mice by μCT to ask whether these phenotypes were also present in Chd8fl/+ mice (Fig. 1A–D). The interorbital distance (landmarks 8–9, Fig. 1C,D) was...
significantly wider in Chd8+/− mice compared with controls, indicative of a hyperteloric phenotype (P = 0.0273; Fig. 1C,D,F). In addition, the anterior–posterior length of the interparietal bone (landmarks 4–5) is increased in Chd8+/− animals ("P = 0.0025, Fig. 1A,B,E), suggestive of more wide-spread craniofacial anomalies associated with Chd8 haploinsufficiency.

To examine whether structural brain abnormalities were present in Chd8+/− mice, their brains were compared with Chd8+/+ littermates by high-resolution MRI (Fig 1G). Total brain volume was increased by 2.7% in Chd8+/− mice (476 vs. 463 mm3, P = 0.048, FDR = 15%, Fig. 1H). Accordingly, several brain regions, including cortical areas, hippocampus and parts of the cerebellum showed volumetric increases (Fig. 3G,H, Supplementary Table S1). Structural alterations of these brain areas have been implicated in autism (Blatt 2012; Ecker 2016; Donovan and Basson 2017) providing potential neural substrates for the autism phenotype associated with ChD8 haploinsufficiency in humans.

**Chd8+/− Mice Show Abnormal Activity Levels and Differences in Social Interaction**

We next assessed Chd8+/− mice in a number of behavioral tests to ask whether they exhibited any signs of socio-communicative deficits, repetitive behaviors or cognitive inflexibility, representing core ASD-like behaviors in humans.

Chd8 heterozygous pups displayed signs of delayed motor development in the first 2 weeks after birth. Chd8+/− pups took slightly longer than wildtype littermates to develop an effective righting reflex over time ("P = 0.014; Fig. 2A). Correspondingly, Chd8+/− pups spent more time engaged in unsuccessful attempts to turn over on their stomachs as measured during the spontaneous motor behavior observations (P6: P = 0.0312, P8: P = 0.0354; Fig. 2B). Once they were able to move around the cage, mutant pups spent on average more time in locomotion than wildtype littermates suggestive of hyperactivity ("P = 0.009; Fig. 2C).

In the 3-chamber sociability test, adult Chd8+/− mice spent significantly more time in the chamber with the novel age- and sex-matched conspecific mouse than in the other chambers, indicative of normal sociability (Fig. 2D). Interestingly, rather than displaying sociability deficits, mutant mice spent slightly, but significantly more time in the chamber containing the mouse, compared with controls ("P = 0.029; Fig. 2D). Chd8+/− mice also spent more time investigating conspecific mice in a reciprocal social interaction test ("P = 0.015; Fig. 2E). A quantitative olfactory habituation/dishabituation test revealed an increased interest in an odor with social significance (urine) in Chd8+/− mice compared with controls ("P = 0.03, ***P = 0.0002; Fig. 2F). No difference in the time spent investigating a non-social (banana) odor was observed, implying an increased interest specifically in social cues and an otherwise normal capacity for odor discrimination (Fig. 2F).

Examination of these animals in the open field arena revealed a marked hypoactivity in Chd8+/− mice (**P = 10−9; Fig. 2G,H). The hypoactive phenotype was also observed in mutant mice in their homecage environment by measuring activity on a running wheel over a 1-week period ("P = 0.019, Fig. 2I). The open field test did not show any evidence of anxiety in these mice, that is, an increased reluctance to enter the inner, most exposed area of an open field arena (Fig. 2J). This was confirmed in the light/dark box test that showed no difference between wildtype and mutant mice (Fig. 2K). Forelimb grip strength was slightly but significantly reduced in mutant mice ("P = 0.045 [males] "P = 0.042 [females]; Fig. 2L) but Chd8+/− mice showed normal motor abilities on the revolving rotarod, indicating that a reduced capacity to perform motor tasks was unlikely to be the cause of the hypoactive phenotype (Fig. 2M). No evidence of repetitive behaviors was observed by assessing marble burying and self-grooming behaviors (Fig. 2N, O). In fact, mutants showed slightly delayed marble burying behavior, most likely due to their general hypoactivity ("P = 0.04, ***P = 0.0004; Fig. 2N).

Spatial learning abilities and cognitive flexibility were assessed in the hippocampus-dependent Morris water maze test. Chd8+/− mice performed normally in the learning part of this test (Fig. 2P). In a reversal paradigm, these mice were also indistinguishable from wildtype littermates, implying normal cognitive, spatial learning abilities and flexibility (Fig. 2P). Finally, no differences in the number of ultrasonic vocalizations (USVs) of pups separated from the nest were recorded, indicating no obvious communication deficits (Fig. 2Q).

As male mice from this adult behavioral cohort were used for structural MRI analyses (Fig. 1G,H) we were able to correlate their brain volume with specific behaviors. We observed significant inverse correlations between activity in the open field and hippocampal volume ("P = 0.012; Supplementary Fig. S3A), as well as cortical volume ("P = 0.036; Supplementary Fig. S2B). Overall brain volume and hypoactivity showed a weaker correlation ("P = 0.092; Supplementary Fig. S3C) hinting at a degree of specificity for cortical and hippocampal regions.

In summary, Chd8+/− mice displayed no socio-communicative deficits, but rather exhibited a heightened interest in social cues. No evidence for perseverative and repetitive behaviors were observed. Chd8+/− pups showed evidence for hyperactivity and delayed motor development while adult Chd8+/− mice exhibited a hypoactive phenotype, which was significantly correlated with overgrowth in cortical and hippocampal regions.

**Chd8 Haploinsufficiency Causes General Growth Delay but Postnatal Brain Overgrowth**

To determine whether the brain overgrowth phenotype was already present at early postnatal stages when developmental delay was evident, we measured body and brain weights from birth. Chd8−/− pups showed significant growth retardation from postnatal day 3 onwards and into early adulthood (Fig. 3A). Brain and body weight were well correlated in both wildtype and heterozygous mice at P35 (r2 = 0.25, P = 0.0004 and r2 = 0.28, P = 0.005, respectively), with Chd8 mutants displaying higher brain weights compared with their wildtype littermate controls with equivalent body weight (Fig. 3B). A group-wise comparison confirmed the significant increase in normalized brain weight in Chd8+/− mice compared with wildtype littermates (20.4% increase, ***P < 0.0001; Fig. 3C). At P7, normalized brain weights were already significantly larger in Chd8+/− pups compared with wildtype littermate controls (9.3%, ***P = 0.0009) with more subtle differences between the groups observed at P0 (6.7%, P = 0.01).

Together, these analyses suggested that subtle, but cumulative differences in brain growth over time may be responsible for small increases in brain size. Indeed, we did not detect any significant differences in cortical ventricular zone (VZ) proliferation as measured by phosphohistone H3 immunostaining at E12.5, E16.5, or P0 (Fig. 3E,F). However, subtle increases in progenitor proliferation in the VZ cannot be completely ruled out.

**CHD8 Controls the Expression of ASD-Associated Axon Guidance Genes in the Early Postnatal Neocortex**

To gain insights into the transcriptional programs that may underlie the subtle brain overgrowth and abnormal behaviors...
Figure 1. Hypertelorism and mild megalencephaly in Chd8<sup>+/−</sup> mice. (A–D) Representative lateral (A,B) and dorsal (C,D) μCT views of 3D reconstructed skulls from mice with the indicated genotypes. Landmarks from 1 to 15 are indicated by yellow dots. Scale bars = 2 mm. (E,F) Graphs for measurements between indicated landmarks, normalized to average measurements from corresponding wildtype littermates. Mean ± SEM; landmarks 4–5: *P* = 0.0025, *t* = 3.797; landmarks 8–9: *P* = 0.0273, *t* = 2.512; df = 12, student’s *t*-test, *n* = 7 per genotype. (G) High-resolution 7 T structural MRI coronal images of Chd8<sup>+/−</sup> brains from posterior (top left) to anterior (bottom right) are shown. Absolute volumetric differences in size, relative to wildtype controls are colored according to the scale on the left. Effect size is measured in units of standard deviation. Some regions with enlarged volumes are labeled as follows: CbVI = cerebellar lobule VI, MC = motor cortex, EcC = ectorhinal cortex, EnC = entorhinal cortex, HC = hippocampus, CC = cingulate cortex, FC = frontal association cortex. (H) Absolute volumes (mm<sup>3</sup>) are plotted for whole brain, neocortex, and several other brain regions for the different genotypes as indicated. #FDR < 0.15, student’s *t*-tests: brain: *P* = 0.0484, *t* = 2.096; cortex: *P* = 0.0055, *t* = 3.093; entorhinal cortex: *P* = 0.011, *t* = 2.788; hippocampus: *P* = 0.0091, *t* = 2.873 primary motor cortex: *P* = 0.0126, *t* = 2.727; cingulate cortex: *P* = 0.0074, *t* = 2.965; frontal cortex: *P* = 0.0154, *t* = 2.639; frontal association cortex: *P* = 0.0238, *t* = 2.438; df = 21, Chd8<sup>+/−</sup>: *n* = 11, Chd8<sup>++</sup>: *n* = 12. Individual volumes and volume differences for all brain regions are listed in Supplementary Table S1.
The development of the righting reflex in pups at the indicated postnatal days. Pups failing to right by the end of the 60 s test period were given a score of 60 s. Note the significant delay in the acquisition of the full righting reflex response in Chd8−/− animals compared with littermate controls. Mean ± SEM; *P = 0.014 (one-way repeated-measures ANOVA: f1,72 = 6.36, between-subjects effect). (B) The duration, in seconds, spent on their back (curling) as recorded during the analysis of spontaneous movements during USV recordings. Note that Chd8−/− mice spend significantly more time curling at P6 and P8 compared with littermate controls. Mean ± SEM, P6: *P = 0.032, P8: *P = 0.0354 (one-way repeated-measures ANOVA: f1,72 = 12.64, P = 0.001 [between-subjects effect], with student’s t-test as post hoc analysis) (p6: df = 72, t = 2.197, P8: df = 72, t = 2.145). (C) The duration, in seconds, pups spent in locomotion as recorded during the analysis of spontaneous movements during USV recordings. At P12 Chd8−/− animals spent significantly more time in locomotion as compared with littermate controls. Mean ± SEM; *P = 0.009 (one-way repeated-measures ANOVA: f1,72 = 7.33, P = 0.008 [between-subjects effect], with student’s t-test as post hoc analysis t = 2.687). (D) The duration, in seconds, spent in each chamber of the 3-chamber sociability test. All mice spent a significantly higher proportion of time in the chamber with the age- and sex-matched stranger conspecific mouse compared with the other chambers. Mean ± SEM; *P = 0.029 (between-subjects ANOVA: f1,53 = 5.033). (E) Duration, in seconds, of social investigation over a 3-min period. Social investigation was defined as the combined total duration of head, body and anogenital sniffing of a conspecific mouse. Mean ± SEM; *P = 0.015 (between-subjects ANOVA: f1,6 = 5.307). (F) Graph plotting the performance in the olfactory habituation/dishabituation test. Mean ± SEM; *P = 0.03, **P = 0.0002 (repeated-measures ANOVA: f2,85,145.23 = 9.24, P = 0.00002, with student’s t-test as post hoc analysis “df = 53, t = 4.04, df = 53, t = 2.23”). (G) Representative ethovision tracks of a Chd8−/− (+/−) and Chd8−/− (+/+) animal plotting their movements during the 10-min open field task. (H) The total distance traveled in the outer part of the open field arena over a 10-min time-period. Mean ± SEM; **P = 2 × 10−6 (between-subjects ANOVA: f1,53 = 52.72). (I) The total activity counts per 12 h period on running wheels in the homecage during 7 days of dark-phase recording. Mean ± SEM; *P = 0.019 (repeated-measures ANOVA: f1,17 = 9.12, between-subjects effect). (J) The percentage of time spent in the center of the open field arena during the 10-min test. Mean ± SEM (between-subjects ANOVA: f1,53 = 0.007, P = 0.93). (K) The percentage of time spent in the light chamber during the 5 min light/dark test. Mean ± SEM (between-subjects ANOVA: f1,15 = 0.824, P = 0.368). (L) The average of 3 measurements of forelimb grip strength on a Linton Grip Strength meter. Mean ± SEM, males: *P = 0.045 (between-subjects ANOVA: f1,29 = 4.371) females: *P = 0.042 (between-subjects ANOVA: f1,12 = 4.677). (M) The mean latency of mice to fall from the rotarod. Mean ± SEM (repeated-measures ANOVA: f1,64,102 = 0.620, P = 0.540). (N) The average number of marbles buried, out of a maximum of 12, within a 30-min time period. Mean ± SEM; *P = 0.04, **P = 0.0004, (repeated-measures ANOVA: f3,66,265 = 4.70, P = 0.002, with student’s t-test as post hoc analysis “df = 53, t = 2.12, “df = 53, t = 3.79). (O) The duration, in seconds, mice spent self-grooming during the 10-min self-grooming test. Mean ± SEM (between-subjects ANOVA: f1,51 = 1.21, P = 0.28). (P) Graph plotting the average distance swum for 4 trials daily over 8 consecutive training days to find the hidden platform (hidden trials), followed by 6 training days where the location of the platform was reversed (reversal trials). Mean ± SEM (repeated-measures ANOVA: f8,761,714 = 1.064, P = 0.388). (Q) The mean number of ultrasonic vocalizations per minute on indicated postnatal days. Mean ± SEM (repeated-measures ANOVA: f1,72 = 0.76, P = 0.39).
observed in Chd8<sup>−/−</sup> mice, we performed RNA-seq analysis on dissected neocortical tissue at 2 stages: (1) At E12.5, when Chd8 expression peaks (Durak et al. 2016) and neural progenitor cells predominate; and (2) At P5, when many developmental processes with relevance for ASD aetiology, such as axon growth and guidance and synaptogenesis, are under way.

Surprisingly, only 5 genes, including Chd8, showed significant (FDR < 0.05) differential expression in Chd8<sup>−/−</sup> embryos at E12.5 in this experiment (Fig. 4A, Supplementary Table S2). By contrast, 649 DEGs (FDR < 0.05) were identified in the P5 neocortex, with over two-thirds of these genes downregulated (Fig. 4B, Supplementary Table S3).

Comparing all DEGs from the P5 dataset with the SFARI autism gene list identified 56 shared genes, representing a highly significant enrichment of ASD-associated genes in the DEG set (P = 1.06 × 10<sup>−10</sup> [OR = 2.87]; Fig. 4C, Supplementary Table S4). Almost all (53/56 = 95%) of these ASD-associated genes were downregulated (Supplementary Table S4). We also overlapped our gene set with high confidence (SFARI categories 1 and 2) ASD candidates (P = 3.26 × 10<sup>−4</sup> [OR = 4.61]; Fig. 4D). Nine genes, representing 16% of all SFARI categories 1 and 2 genes, were present in our DEG set at P5. All of these high confidence ASD candidate genes were downregulated (Supplementary Table S4).

Amongst the upregulated gene set, the most significant KEGG pathways, molecular functions and biological processes were related to protein transport, the ribosome and oxidative phosphorylation, whereas the downregulated gene set included categories related to cell adhesion, axonal guidance and calcium signaling pathways (Fig. 4E, F, Supplementary Fig. S2A, Supplementary Tables S5–10). Identification of potential regulatory transcription factors was performed using Enrichr, which found over-representation of Suz12 targets in the downregulated gene set (Supplementary Fig. S2B). Suz12 is a component
Figure 4. Gene expression changes in Chd8-deficient neocortices. (A) Volcano plot of RNA-seq data from embryonic (E12.5) Chd8<sup>−/−</sup> neocortex. Each point represents an individual gene and all genes differentially expressed in Chd8<sup>−/−</sup> samples with an FDR of 0.05 are highlighted in red. All differentially regulated genes are listed in Supplementary Table S2. (B) Volcano plot indicating differentially expressed genes (DEGs) detected by RNA-seq in P5 Chd8<sup>−/−</sup> neocortex. All differentially regulated genes are listed in Supplementary Table S3. (C) Venn diagram showing extent of overlap between P5 DEGs and ASD-associated genes (categories 1–5 and S) in the SFARI gene database. Enrichment was calculated using Fisher’s exact test for count data. SFARI genes that overlap with P5 DEGs are listed by category in Supplementary Table S4. (D) Pie chart showing the proportion of high confidence ASD candidate genes (categories 1–2) that are found in the P5 DEG set. Enrichment was calculated using Fisher’s exact test for count data. (E, F) Results of gene set enrichment analysis using the DAVID knowledgebase on the P5 DEG set (FDR < 0.05). The 5 most significant Gene Ontology terms in the Biological Processes category are shown for upregulated DEGs (E) and downregulated DEGs (F), respectively. The 5 most significant Gene Ontology terms in the molecular function and pathways categories and the 4 most over-represented transcription factors identified by Enrichr analysis are shown in Supplementary Figure S2. A comprehensive list of all significant Gene Ontology terms in the biological processes, molecular functions and pathways categories is given in Supplementary Tables S5–10.

of the Polycomb repressor complex 2 (PRC2) and is required for both histone methyl transferase and gene silencing activities of PRC2 (Cao and Zhang 2004). The observation that Suz12 targets are over-represented in the downregulated gene set offers a potential mechanistic explanation for the downregulation of some of these genes. None of the genes that encode PRC2 subunits like Suz12 were differentially expressed at P5, excluding the possibility that increased PRC2 gene expression at this stage of development was responsible for repression of Suz12 target genes in Chd8 mutants.

**Chd8<sup>−/−</sup>** Mice Exhibit Over-Connectivity in Cortical and Hippocampal Networks

The significant enrichment of cell adhesion and axonal guidance genes in the downregulated gene set at P5 led us to hypothesize that long-range connectivity might be altered in Chd8 heterozygous neocortices. To test this hypothesis, we performed rsfMRI to probe functional brain connectivity in mature brain networks. Synchronous fluctuations in blood-oxygen-level dependent (BOLD) signals in different brain regions are used as an indication of them being functionally connected. A regionally unbiased analysis for long-range connectivity changes revealed hotspots for increased connectivity in Chd8<sup>−/−</sup> mice compared with wildtype littersmate controls, which included the entorhinal, retrosplenial, auditory cortical and posterior hippocampal areas (t-test, P < 0.05 FEW cluster-corrected, with cluster-defining threshold t<sub>k</sub> > 2.06, P < 0.05; orange areas in Fig. 5A). This analysis suggested that hyperconnected areas were predominantly located on the left side of the brain. A reanalysis of these results without the use of cluster correction revealed the presence of foci with increased connectivity also on the right side, mirroring the effects observed on the left (dark red areas in Fig. 5A). Interhemispheric mapping of rsfMRI connectivity strength in previously characterized rsfMRI network systems of the mouse brain (Sforazzini et al. 2014), revealed increased cortical connectivity in auditory regions (P < 0.05, student’s t-test, uncorrected), although the effect did not survive FDR correction (q = 0.05) for multiple comparison across the rsfMRI networks probed. We next used a seed-based approach to specifically probe regions with altered connectivity to these hotspots to reveal the brain networks affected. Most strikingly, this revealed a reciprocal increase in connectivity between ventral hippocampus and auditory cortical regions in Chd8 mutant mice (t-test, P < 0.05 FEW cluster-corrected, with cluster-defining threshold t<sub>k</sub> > 2.06, P < 0.05; Fig. 5B,C). Seed placement in the auditory cortex revealed increased connectivity of this region with both cingulate and entorhinal cortices (Fig. 5B), whereas a hippocampal seed uncovered strengthened long-range connectivity with somatosensory and visual cortices (Fig 5C).

Previous studies suggested that neurodevelopmental genes show correlated expression patterns in brain areas that are connected (French and Pavlidis 2011). To test if the same applies to our model we harnessed gene expression data contained within the Allen brain atlas to identify genes that are highly expressed in hippocampal and auditory areas. Using expression data from the Allen brain atlas, we determined genes with high relative expression in both the hippocampal CA2 region and auditory areas (Supplementary Fig. S4A, B; Supplementary Table S11; see Methods for details). Enriched genes preferentially clustered into GO term categories relating
to synaptic development and function, axonal structure and neuron projections (Supplementary Table S11). To determine if any of these highly expressed genes are likely contributing to a developing process linked to the functional connectivity phenotype, we compared them with the DEG set from our P5 RNAseq experiment. This analysis revealed several axon guidance and cell adhesion genes that are preferentially expressed in CA2 and auditory areas at adult stages and whose expression is dysregulated at P5 (e.g., Cdh2, Cdh11, Cdk5r1, Epha4, Fat3, Nrcam, Robo1; Supplementary Table S11).

Taken together, two independent experimental approaches at different time points identified specific axon guidance and cell adhesion genes, further strengthening our hypothesis that dysregulation of these genes may be important for the functional connectivity phenotype and provides a solid platform for future detailed analyses.

We conclude that early postnatal gene expression changes prefigure abnormal functional connectivity in Chd8 heterozygous mice. Our findings suggest that abnormalities in specific cortical–hippocampal circuits involved in sensory processing may underlie some of the unique anomalous behaviors observed in Chd8+/- mice, and by extension, the neuropsychiatric symptoms in patients with CHD8 mutations.

Discussion

Here we identify a crucial developmental role for Chd8 in regulating axon guidance gene expression in the early postnatal period and, for the first time, associate Chd8 haploinsufficiency with functional over-connectivity of specific brain areas. A recent rsfMRI study involving over 150 male probands with an ASD diagnosis and nearly 200 typically developing individuals.
described over-connectivity between sensory cortices and subcortical structures as a central feature in ASD (Cerilani et al. 2015). It will be very important to determine whether these specific functional connectivity abnormalities are present in patients with CHD8 mutations.

While we cannot at this stage establish a direct causal relationship between transcriptional, connectivity and behavioral phenotypes, our data suggest that functionally altered connectivity of sensory cortical areas in Chd8 mutant mice underpins behavioral phenotypes. In concordance with previous studies (Katayama et al. 2016; Gompers et al. 2017), our data suggest that moderate expression changes of many genes, rather than severe disruption of few genes, cooperate to give rise to phenotypic changes in Chd8−/− mice. A total of 21 axon guidance genes are downregulated in the early postnatal period in our Chd8−/− mice (Supplementary Table S6). Therefore, experimental validation of a causal link will be challenging as multiple axon guidance pathways may contribute to the functional connectivity and behavioral phenotypes.

**Chd8−/− Mice as a Model for Human CHD8 Haploinsufficiency Syndrome**

CHD8 is one of the highest confidence ASD-associated genes to emerge from recent exome sequencing studies (Neale et al. 2012; Talkowski et al. 2012; O’Roak et al. 2012a; Bernier et al. 2014; Iossifov et al. 2014). We therefore expected Chd8−/− mice to present with robust, autism-associated behaviors. Chd8−/− mice displayed delayed motor development and distinctive behavioral anomalies that featured a heightened interest in social cues, but surprisingly did not include repetitive and perseverative behaviors or communication deficits.

In agreement with other published studies (Katayama et al. 2016; Gompers et al. 2017; Platt et al. 2017) we did not observe repetitive behaviors in Chd8−/− mice. While Katayama et al. reported increased persistence following directional reversal in the T-maze forced alteration test, suggestive of perseverative behaviors, we did not find such evidence in the Morris water maze test. This may be due to the higher complexity of decision making in the Morris water maze compared with the binary choice required by the T-maze. In addition, Chd8−/− mice consistently did not show any evidence for perseverative behaviors in the marble burying test (Fig. 2N; Gompers et al. 2017; Platt et al. 2017), suggesting that any perseverative behaviors in Chd8 mutants may be subtle or task-specific. Chd8−/− mice show an apparent heightened interest in social cues, indicating that altering the CHD8 gene dosage during development can impact socially motivated behaviors (Fig. 2E). An increased duration of contacts in the social investigation test was also seen in two other published behavioral analyses of Chd8 heterozygous mouse models (Katayama et al. 2016; Platt et al. 2017). Katayama et al. additionally described a reduced duration of active social contacts in Chd8 mutants, although all test groups showed evidence for high levels of anxiety, a known behavioral confound. Katayama et al. (2016) and Platt et al. (2017) further reported normal sociability but minor deficits in social novelty in the 3-chamber social approach task in Chd8−/− mice.

Despite not observing typical ASD-like behaviors, we did detect a delay in early motor development in Chd8−/− mice (Fig. 2A,B). There is a growing body of evidence suggesting that delayed motor milestones in toddlers predate and predict the emergence and severity of language deficits in later life (Bedford et al. 2016; Chinello et al. 2016). Of note, the only available longitudinal case reports in the literature also describe early motor delay in both patients with CHD8 haploinsufficiency (Merner et al. 2016; Stolerman et al. 2016).

A key characteristic of autism is restricted behaviors or interests, which often manifest as hyper- or hyporeactivity to sensory input or unusual interest in sensory stimuli, for example, excessive smilling or touching of objects (Constantino and Charman 2016). One may speculate that the excessive smilling of social cues and the increased duration of social contacts observed in our Chd8−/− mice may be indicative of behavioral abnormalities in these domains.

**Dysregulation of the Cortical Transcriptome in Chd8 Heterozygous Mice**

Gene expression analysis showed little evidence for transcriptional dysregulation at midembryonic stages, but revealed disruption of key developmental processes involved in establishing brain connectivity in the early postnatal neocortex. These data are in agreement with a recent study where Gompers and colleagues only found a handful of genes differentially expressed in bulk forebrain at E12.5 and E14.5, while detecting subtle, more widespread changes perinatally (E17.5: 89 DEGs, P0: 35 DEGs; FDR < 0.05) and more pronounced dysregulation at adult stages (295 DEGs; FDR < 0.05) (Gompers et al. 2017).

Many of the transcripts that were dysregulated in the early postnatal period are themselves ASD-associated genes and were predominantly downregulated. Our gene expression studies therefore provided strong evidence that a variety of genes, pathways and developmental processes implicated in ASD might be dysregulated by Chd8 haploinsufficiency.

An expanding number of ASD risk genes have roles in axon guidance, synapse development and plasticity (Bourguerón 2015). We detected significant enrichment of genes in these functional categories in our downregulated gene set, including the major Slit protein receptors Rob1 and Rob2, EphA4 and 5, and cell adhesion molecules such as L1CAM and Cad2, 5, 8, and 11 (Supplementary Tables S3 and S6). Similarly, Gompers et al. found enrichment for axon growth and guidance factors amongst downregulated genes in their M3 module (Gompers et al. 2017). Moreover, Sugathan et al. showed enrichment for genes associated with the GO terms “cell adhesion,” “axon guidance,” and “neuron differentiation” amongst downregulated genes in CHD8-deficient human iPSC-derived neural progenitors. This suggests that these important developmental gene sets are regulated by CHD8 in both mouse and human cells (Sugathan et al. 2014).

In sum, our data identify early postnatal development as a key stage at which transcriptional changes caused by Chd8 heterozygosity may precipitate ASD-related phenotypes. They further indicate that Chd8 heterozygosity defines a transcriptional program characterized by diminished expression of key neurodevelopmental regulators that are predicted to affect cellular functions essential for the appropriate wiring of the brain.

**Increased Functional Connectivity in Sensory Networks**

Significantly, functional connectivity was altered in the adult brain of Chd8−/− mice. Our rsfMRI analysis found evidence for over-connectivity between sensory regions in the neocortex and limbic cortical regions. Most notably, the auditory cortex showed a global increase in functional connectivity that involved connections to other cortical areas and reciprocal strengthening of connectivity to the ventral hippocampus. It seems likely that altered connectivity is the consequence of
some of the disrupted brain wiring pathways uncovered by our RNA-seq experiments. Encouragingly, the expression of several axon guidance and cell adhesion genes, which are dysregulated at P5, is enriched in hippocampus and auditory cortex (Supplementary Table S11); nevertheless, this hypothesis will require further in-depth scrutiny. More importantly, it will be critical to investigate whether these connectivity changes are pertinent to any of the behavioral anomalies in CHD8 heterozygous mice or the ASD phenotype in patients with CHD8 haploinsufficiency. The over-connectivity in networks involving the auditory cortex and the hippocampus is intriguing. Auditory processing deficits in ASD are well documented and range from a lack of lateralization to a general delay in network maturation (Brunet et al. 1992; Edgar et al. 2015), although the functional behavioral consequences of these deficits are not clear. Furthermore, over-responsivity to sensory stimuli is frequently observed in ASD patients, can affect all sensory modalities and appears to be positively correlated with the severity of autistic traits (reviewed in Tavassoli et al. 2014; Sinclair et al. 2017). Although a definitive causal relationship is difficult to establish, it has been hypothesized that sensory over-responsivity may trigger compensatory and avoidance behaviors that promote the emergence of core behavioral autism traits (Marco et al. 2011). In support, tactile hypersensitivity during critical developmental periods has been shown to underlie anxiety and social deficits in a number of genetic ASD mouse models (Orefice et al. 2016). Whether this would be equally the case for other sensory modalities and a general mechanistic feature in the behavioral etiology of ASD remains an open question.

Supplementary Material
Supplementary material is available at Cerebral Cortex online.

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