Comprehensive multi-omics integration identifies differentially active enhancers during human brain development with clinical relevance

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Abstract

Background: Non-coding regulatory elements (NCREs), such as enhancers, play a crucial role in gene regulation, and genetic aberrations in NCREs can lead to human disease, including brain disorders. The human brain is a complex organ that is susceptible to numerous disorders; many of these are caused by genetic changes, but a multitude remain currently unexplained. Understanding NCREs acting during brain development has the potential to shed light on previously unrecognized genetic causes of human brain disease. Despite immense community-wide efforts to understand the role of the non-coding genome and NCREs, annotating functional NCREs remains challenging.

Methods: Here we performed an integrative computational analysis of virtually all currently available epigenome data sets related to human fetal brain.

Results: Our in-depth analysis unravels 39,709 differentially active enhancers (DAEs) that show dynamic epigenomic rearrangement during early stages of human brain development, indicating likely biological function. Many of these DAEs are linked to clinically relevant genes, and functional validation of selected DAEs in cell models and zebrafish confirms their role in gene regulation. Compared to enhancers without dynamic epigenomic rearrangement, DAEs are subjected to higher sequence constraints in humans, have distinct sequence characteristics and are bound by a distinct transcription factor landscape. DAEs are enriched for GWAS loci for brain-related traits and for genetic variation found in individuals with neurodevelopmental disorders, including autism.

Conclusion: This compendium of high-confidence enhancers will assist in deciphering the mechanism behind developmental genetics of human brain and will be relevant to uncover missing heritability in human genetic brain disorders.

Keywords: Clinical genetics, Non-coding genome, Enhancer, Gene regulatory elements, Meta-analysis, Data integration, Computational analysis, Mendelian disorders, Human brain development
Background

Non-coding regulatory elements (NCREs), such as enhancers, play a pivotal role in gene regulation [1, 2]. Enhancers ensure correct spatio-temporal gene expression, and it is increasingly recognized that genetic aberrations disturbing enhancer function can lead to human disease, including brain disorders [3–6]. Such non-coding genetic variants are expected to explain a considerable fraction of so-called missing heritability (e.g., the absence of a genetic diagnosis despite a high genetic clinical suspicion). These developments are pushing genetic diagnostic investigations to shift from whole-exome sequencing to whole genome sequencing, and the number of potentially pathogenic non-coding variants found in patients is expected to rise [4]. It is therefore of urgent clinical interest to understand where functionally relevant non-coding sequences are located in the human genome, as this will help to interpret the effects on health and disease.

Despite tremendous progress over the last decades, our understanding of the underlying mechanisms of enhancer biology remains limited due to challenges in annotating functional enhancers genome-wide. Large-scale community-driven efforts [7–11] and an uncountable plethora of individual studies have produced a vast amount of epigenome data sets, such as profiles of histone modifications, chromatin accessibility, and chromatin interactions for different human tissues and cell types, that can be used to predict putative enhancers at a large scale. More recently, new technologies such as massively parallel reporter assays and CRISPR-Cas9-based screens have entered the stage [12–14], providing novel means to directly test the functionality of non-coding regions. In addition, computational prediction algorithms [15, 16], trained on epigenome and experimental data, are improving the capability to predict functional sequences and the effects of variants in these regions.

One of the inherent problems with this increasing amount of data is the difficulty in keeping track of individual data sets and the ability to integrate data from various sources. Usually, individual studies focus on a limited number of cell types or tissues and compare their findings to a small number of previously published data sets. Although this is a logical step, it does not leverage the potential to fine-tune enhancer predictions which integrating all available enhancer data could have. This is illustrated by our previous findings that the overlap between individual enhancer predictions from several studies tends to be quite poor [4]. This is likely caused by heterogeneity of starting biological samples, limitations of current technologies, and differences in data analysis. Although the first two are difficult to change, analyzing these data in a similar way could avoid some of the noise and difference generated by data analysis.

Here we undertook such an integrative effort, focusing on human brain development (Fig. 1A, Additional file 1: Fig. S1). We retrieved virtually all previously published putative enhancers for brain (from PubMed and enhancer databases, n ≈ 1.6 million putative enhancers) (Additional file 2: Table S1) [9, 11, 21–31], and performed an integrative analysis of relevant available epigenome data sets (n = 494) [9, 10, 19, 31–35] (Additional file 3: Table S2), after reanalyzing the data. Using this approach, we identify around 200 thousand putative critical regions (pCRs) in reported brain enhancers, of which around 40 thousand show dynamic epigenomic rearrangement during fetal brain development, indicating switching on and off of regulatory elements during development. We thus refer to these regions as differentially active enhancers (DAEs). Compared to their non-variable counterparts (nDAEs), DAEs have a higher level of sequence constraint, regulate genes that are expressed during fetal brain development and are associated with brain developmental processes. DAEs are enriched for binding sites of brain-relevant transcription factors, brain-related GWAS loci and are regulating disease-relevant Online Mendelian Inheritance in Man (OMIM) genes. We validate a selected number of DAEs using in vitro reporter assays and CRISPRi in cell lines, and reporter assays during zebrafish development. Together, this provides an easily accessible and comprehensive resource of NCREs that are likely functional during human brain development.

Methods

Data visualization

To visualize enhancers and epigenome data, we used the UCSC Genome Browser (https://genome.ucsc.edu/). To generate UCSC Genome Browser Tracks, aligned reads were converted to bedgraph using genomeCoverageBed, after which the bedGraphToBigWig tool from the UCSC Genome Browser was used to create a bigwig file [36, 37]. All enhancer regions, enhancer-gene interactions, and topologically associating domain (TAD) coordinates were uploaded directly as bed files. Other plots were drawn using R packages and Figs. 1, 2, 3, 4, 5, and 6 and Additional file 1: Figures S1-S8 were assembled in Adobe Illustrator [47]. Additional file 2: Table S1, Additional file 3: Table S2, Additional file 4: Table S3, Additional file 5: Table S4, Additional file 6: Table S5, Additional file 7: Table S6, Additional file 8: Table S7, Additional file 9: Table S8, Additional file 10: Table S9, Additional file 11: Table S10, Additional file 12: Table S11, Additional file 13: Table S12, Additional file 14: Table S13 and Additional file 15: Table S14 were exported as text or Excel files.
Data collection and processing

Collection of putative brain enhancers
To generate a comprehensive set of putative brain enhancers active during fetal brain development, we scrutinized PubMed and various enhancer databases (last assessed: April 2019), including amongst others EnhancerAtlas, the FANTOM5 Project, and the Vista Enhancer database [9, 11, 21–31]. This resulted in 1,595,292 putative enhancers (Additional file 2: Table S1). Enhancers with identical coordinates were deduplicated and the unique regions were used to determine putative critical regions (pCRs), reasoning that overlapping parts of a putative enhancer obtained from different sources might point to functional relevant regions of that putative enhancer. If there is any overlap between coordinates of putative enhancers derived from two or more databases, the pCRs were defined as maximum overlapping regions present in those databases using the BEDtools suite (mergeBed, intersectBed, genomeCoverageBed, and groupBy sub-commands) (version 2.30.0) [36]. Putative enhancers that were only present in one of the input sources were also included in the pCRs (Fig. 1A, step 1), as it cannot be excluded that these putative enhancers are biologically relevant. pCRs with length less than 50 bp and more than 1000 bp were excluded. To avoid any overlap with gene promoters, enhancers located within 2 kb upstream or 1 kb downstream of a transcriptional start site (TSS) (Ensembl GRCh37.p13 Release 102) were excluded using intersectBed. Following this procedure, we identified a total of 202,462 pCRs which were used for downstream analyses. Next, we excluded 299 pCRs that were not covered by sufficient amounts of epigenome data (less than 10 reads in at least two samples (see section on defining DAEs)), resulting in a final number of 202,163 pCRs (Additional file 4: Table S3). GREAT web interface was used (version 4.0.1.1) (http://broadinstitute.github.io/picard/). These reads were used to define differentially active enhancers (DAEs).

Defining differentially active enhancers (DAEs)
We assumed that pCRs with high variability in different epigenome data (dynamic epigenomic rearrangement) across different developmental stages are more likely to be functional than other pCRs. To determine this variability, the number of overlapping reads (for each epigenome mark) with pCRs was counted using the multiBamCov sub-command of BEDtools and a matrix was generated that included enhancers as rows and epigenome features as columns. Epigenome features were from different brain regions and developmental stages. In total, 299 pCRs with less than 10 reads were excluded, leaving 202,163 pCRs for this analysis. Subsequently, the raw read count matrix was normalized using TMM-normalization [49] and the normalized count matrix was used to define DAEs across different developmental stages and brain regions of a given epigenome data using edgeR (version 3.32.1) [50]. Since there were different developmental stages (time-point factor) and brain regions (brain part factor) in each epigenome data, a design matrix was generated for each factor separately. A limited number of samples without biological replicates were grouped together with other samples based on high correlation (Pearson correlation; r > 0.89). The DAEs were defined based on each design matrix using a generalized linear model and quasi-likelihood F-tests. In order to define the final DAE list, DAEs identified from at least two epigenome data-specific matrices were pooled. In total, this resulted in 39,709 DAEs (FDR adjusted p value < 0.05). The remaining 162,454 pCRs that did not show variability were considered as nDAEs (Additional file 4: Table S3).

Identifying chromatin interactions

Enhancer-gene interactions
In order to define enhancer-gene interaction, published HiC data from 3 human fetal brains, for cortical plate (CP) and germinal zone (GZ) at gestation weeks 17–18 were used [30]. This data provides 10 kb resolution bins for gene loop interactions and 40 kb resolution for

FASTQ files using our analysis pipeline (Additional file 1: Fig. S1). First, adaptor contamination was removed using Trim Galore (version 0.6.5) (https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/), and trimmed data were aligned to the GRCh37/hg19 human genome using Bowtie2 aligner (version 2.4.2) (with --very-sensitive parameter) [48]. Only properly paired and uniquely mapped reads, with mapping quality more than 30 (MAPQ ≥ 30), were kept followed by removing any possible duplicated reads using Picard’s MarkDuplication (version 4.0.1.1) (http://broadinstitute.github.io/picard/). These reads were used to define differentially active enhancers (DAEs).

Epigenome data
Epigenome data were collected from the Roadmap Epigenomics Consortium, ENCODE, PsychENCODE, and other studies (Additional file 3: Table S2). Epigenome data sets used for integration included histone modifications (H3K27ac, H3K27me3, H3K4me1, H3K4me2, H3K4me3) and chromatin accessibility (ATAC-seq and DNase-seq) from different brain regions and different human developmental stages (Fig. 1A, step 1). To avoid any possible confounding biases because of the various pipelines used in different studies, we reanalyzed the raw
TADs. Pre-calculated significant interactions were intersected with pCRs (DAEs and nDAEs) using intersectBed to define gene-enhancer interaction for both CP and GZ separately. Out of the 202,163 pCRs, 41,041 pCRs engaged in 101,366 interactions in CP, and 41,085 pCRs had 100,521 interactions in GZ. Enhancer-gene interactions locating within the same TAD were considered for downstream analyses (almost 80% of all interactions were intra-TAD). We only included protein-coding and lncRNA genes in our analysis. To determine enhancer-enhancer interactions in Additional file 1: Fig. S7A, we also intersected HiC data with pCRs, focusing on interactions between DAEs and both DAEs and nDAEs.

In addition to HiC, we employed other enhancer-gene interaction predictions including JEME (http://yiplab.cse.cuhk.edu.hk/jeme/) [51], ENCODE (https://ernstlab.biolchem.ucla.edu/roadmaplinking/) [52], FOCs (http://acgt.cs.tau.ac.il/focs/download.html) [53], and GeneHancer (downloaded from UCSC table browser; hg19; updated 2019) [54]. These databases apply statistical models on different types of omics data to predict enhancer-gene interactions. We collected fetal brain enhancer-gene predictions from JEME and ENCODE and all brain-related enhancer-gene predictions from FOCs and GeneHancer, as the latter two resources do not specify fetal-specific interactions. In addition, we used H3K27ac HiChIP derived chromatin interactions from several postnatal brain regions [55] cell type-specific chromatin conformation capture data from PLAC-seq experiments in postnatal brain tissue [56] and enhancer-gene interaction predictions generated by the Activity-by-contact (ABC) model (https://github.com/broadinstitute/ABC-Enhancer-Gene-Prediction) [57]. We performed the ABC model by fixing the length of pCRs to 500 bps from the center (250 bps from each side). The enhancer activity was then determined considering DNase, and H3K27ac samples, and gene expression data from fetal brain [10] using default settings of the “run.neighborhoods.py” function. The ABC score was calculated by integrating the fetal HiC data and enhancer activity defined using the default settings of the “predict.py” function and adjusting “--hic_type bedpe,” “--hic_resolution 10000” flags, and ignoring “--cellType” flag.

Intersections between the pCRs and each of these predictions were considered as enhancer-gene interaction (Additional file 6: Table S5). The coordinates of the HiChIP interactions were lifted over to hg19 before intersecting with pCRs.

Functional enrichment analysis

Enhancer sequence characteristics analysis

To determine whether different DNA sequence features distinguish different enhancer groups and whether there is any association between these features and functional prediction, we considered the following features: (i) the non-coding essential regulation (ncER) score (i https://github.com/TentiLab/ncER_datasets/; updated 06-03-2019) [38]; (ii) GC content, as determined by the GContent R packages based on BSeq. Haploinsufficiency scores [43]; (vii) probability of loss-of-function intolerance (pLI) score [42]. The overlaps between DNA sequence features and enhancer coordinates were defined using intersectBed. As assessed enhancers (e.g., pCRs) varied in length between 50 and 1000 bp, and the abovementioned scores were given either at the nucleotide level or in certain bins (depending on the given scores from the individual resources), we calculated the mean value for each enhancer and used this in group comparisons. For gene-specific scores (e.g., pLI), we plotted the scores of the genes linked to the enhancers. Statistical significant differences between groups were determined using Wilcoxon signed rank test in R.

Gene expression correlation

To compare gene expression levels of enhancer target genes between different groups, various transcriptome data were collected. This included transcriptome data from different brain regions and developmental stages, and also various control data from other fetal tissues from the Roadmap Epigenomics Consortium, ENCODE project, Allen human brain atlas, and other studies (Additional file 7: Table S6) [7, 10, 18–20]. Raw data (FASTQ) was quality controlled and adaptors and other contaminants were removed using Trim Galore (version 0.6.5), reads were mapped to the GRCh37/hg19 human genome assembly using STAR aligner (version 2.7) [58], and gene counts were obtained using htseq-count (version 0.12.4) [59]. Gene expression levels were normalized based on fragments per kilobase of transcript per million mapped reads (FPKM). To correlate enhancers to gene expression, enhancer-gene interactions were derived from the HiC data or the alternative enhancer-gene predictions as described above. Gene expression levels were plotted and statistical comparison was performed, between expression levels of subgroups, using Wilcoxon signed rank test in R. We also compared genes linked to DAEs and nDAEs by HiC, to the three trajectory gene groups from BrainVar [60]. For this, we first found the overlap between genes interacting with DAEs/nDAEs using HiC-CP/GZ and each of the three trajectory groups (e.g., falling, rising, and constant genes). We then determined the odds ratio between
DAE and nDAE linked genes for each of the three gene trajectories, and used Fisher’s exact test to determine significance.

**Gene ontology analysis**

For functional enrichment analysis, we used GREAT [17], Enrichr [61], and Metascape [62]. GREAT was used via the web interface (version 4.0.4) (http://great.stanford.edu/public/html/) using the following settings: basal plus extension, proximal 5 kb upstream and 1 kb downstream, plus distal up to 100 kb, including curated regulatory domains, and either whole genome or all pCRs as background, as indicated in the tabs of Additional file 5: Table S4. The – log 10 p value was used to rank GREAT enrichment. Enrichr and Metascape were also used via the web interface (https://maayanlab.cloud/Enrichr/; https://metascape.org/gp/index.html#/main/step1), using the default settings and the whole genome set as background. All outputs of p value, adjusted p value (q value), and combined score (which is the estimation of significance based on the combination of Fisher’s exact test p value and z score deviation from the expected rank) for Enrichr and of LogP, enrichment, z score, and log(q value) for Metascape are reported in Additional file 5: Table S4, Additional file 8: Table S7 and Additional file 10: Table S9.

**Transcription factor binding enrichment**

We used LOLA [63] using default settings to assess binding of known transcription factors to DAEs and nDAEs (Fig. 3). We used motifs from the JASPAR motif database (using reference genome GRCh37/hg19 and LOLAjaspar database core), to test the TF enrichment in DAEs and nDAEs, using all pCRs as background. The mean rank index (a combination of p value, odds ratio, from Fisher’s exact test and the raw number of overlapping regions), was used to rank the known motifs. To display TF enrichment in Fig. 3C and F, we re-scaled the rank between 0 and 100 using the rescale () R function. To further identify motifs across the different relative DAE or nDAE bins and distinguish motifs in the central versus peripheral parts of the enhancers (Fig. 3G, H), we split the 100 relative bins into 20 groups of 5 consecutive bins and performed motif enrichment analysis using HOMER (version 4.11) [64], using function “findMotifs-Genome.pl” and all pCRs as background. A p value ≤ 0.01 was considered to select significantly enriched motifs.

**Transposable element enrichment**

The RepeatMask (GRCh37/hg19, updated 20-02-2020) was downloaded from the UCSC table browser and joined to the pCRs. To determine enrichment of transposable elements in brain enhancers, we followed a strategy previously used when investigating active enhancers in human embryonic stem cells [65]. The number of overlaps of each type of repeat (n_overlaps) with all pCRs (n) was used to calculate the relative frequency (f_all = n_overlaps/n). Multiplication of the relative frequency with the number of regions (n_test, e.g., DAE, nDAE) in any tested group yields the expected frequency (E). This number was compared with the actual observed frequency in the subgroups (f(test) × (n_overlap, test)/n_test = O) to calculate the observed versus expected ratio (O/E). We considered repeats with O/E < 0.5 as depleted, or O/E > 1 as enriched. For the subsequent data interpretation we only focused on transposable elements that were present multiple times (n_overlap > 15) in all pCRs (Additional file 9: Table S8).

**Disease-relevance enrichment**

The Online Mendelian inheritance in Man (OMIM) gene list (updated 28-09-2020) was downloaded using biomaRt R package [66] from Ensembl GRCh37.p13 Release 101. The GWAS catalog (GRCh37/hg19, updated 17-03-2021) was downloaded from the UCSC table browser. The GWAS catalog was manually filtered to keep brain-related studies and their variants with p value ≤ 9e−08 (Additional file 12: Table S11). Stratified LD score regression analysis was performed by implementing the full baseline model to calculate enrichment (https://github.com/bulik/litätsc/wiki [67, 68]. Annotation and LD score files were created using the “make_annot.py” and “ldsc.py” functions, respectively. Partitioning heritability was performed using the “ldsc.py” script considering default parameters with “--h2” flag. We obtained GWAS summary statistics for several brain-related traits including Alzheimer’s disease [69], anorexia nervosa [70], anxiety [71], attention deficit hyperactivity disorder [72], autism spectrum disorder [73], bipolar disorder and schizophrenia [74], epilepsy [75], insomnia [76], intelligence [77], major depressive disorder [78], neuroticism [79, 80], obsessive compulsive disorder / Tourette syndrome [81], Parkinson’s disease [82], and schizophrenia [83] (Additional file 12: Table S11). Z-scores were used to calculate the p values which were corrected for multiple hypothesis testing using the Benjamini-Hochberg method. For CNV analysis, we retrieved pre-processed published data from Brandler et al. (their supplemental Table 9: de_novo_SVs sheet, and their supplemental Table 7: Primary CR Trans and Replication CR Trans sheets) [44] and Monlong et al. (cnv-PopSV-Epilepsy-198affected-301controls-5 kb.tsv.gz file in https://figshare.com/s/20dfdedcc471e465185) [45]. For SNV analysis of the ASD simplex families, we collected de novo variants from supplemental Table 1 of Zhou et al [46]. Autism genes were collected from the SFARI Gene database (http://gene.sfari.org/database/
human-gene) [84]. The overlap between enhancer regions (DAE and nDAE) and each data set was determined using intersectBed. The odds ratio and p value between DAE and nDAE was calculated using fisher.test () R function. The Haldane–Anscombe correction was used to adjust the odds ratio.

Distribution of features across enhancer bins
To investigate the distribution of enrichment of different features (ncER score, GC content, phastCons score, and epigenome data) across enhancers, we divided the enhancer regions into 10 bp bins and calculated the relative scores (the median value for ncER score, GC content, phastCons score) and the number of reads (for epigenome data) for each bin. As the enhancers under investigation differed in size between 50 and 1000 bp, to make enrichments between enhancers comparable, we re-scaled each enhancer bin. To this end, we calculated a relative position between 1 and 100 for each bin of each enhancer, where 1 is the first bin, and 100 is the last bin of each individual enhancer. We then plotted the distribution of each feature across all these re-scaled enhancer bins.

DAE clustering analysis
The matrix of DAEs was used to determine the pattern of epigenome data through different developmental stages. To determine the optimal clustering algorithm, we used clValid R package which simultaneously compares multiple clustering algorithms (hierarchical, kmeans, model-based, pam, and clara). Based on this, the pam algorithm (which is similar to k-means but more robust to noise and outliers) was selected to cluster DAEs using the spearman distance and ward.D^2 method. To define the optimal number of clusters, we used fviz_nbclust and NbClust R packages which compute different indices by bootstrapping (n = 1000). The predicted number of clusters was tested using the silhouette R package to examine whether the clustering performed correctly. This approach resulted in 2 clusters for DAEs and epigenome features at 8–12 PCW, 3 clusters for 13–18 PCW, and 2 clusters for > 18 PCW, for each of CP and GZ, respectively. For each cluster, we determined the gene expression of protein-coding genes interacting with the DAEs from each cluster, as obtained from published RNA-seq data sets. Significant differences in expression levels between different clusters were determined using the Wilcoxon signed rank test in R. Also, target genes linked to each cluster were used for functional enrichment analysis using Enrichr [61], as described under gene ontology analysis (Additional file 10: Table S9).

Enhancer cell type specificity and their dynamics in adult brain
To determine cell type specificity of enhancers, we compared DAEs and nDAEs to recently described cell type-specific regulatory elements from two studies on adult brain (obtained from Supplementary Data Set 4 (data lifted over to hg19) of Corces et al. [55] and Supplementary Table 5 of Nott et al [56]) and a study of fetal brain (obtained from Supplementary file 4 of Domcke et al., specificity scores for top 10000 regions [85]). We used bedtools to intersect DAEs or nDAEs and different cell type-specific regulatory elements. For all DAEs and nDAEs linked to target genes in CP and GZ by HiC, we compared dynamics of H3K27ac levels in both fetal and adult samples, using H3K27ac data from Li et al. [86]. Clustering analysis was performed as described under “DAE clustering analysis” above. Gene ontology analysis for each defined cluster was performed using Enrichr, as described above.

Experimental validation
Cell culture
HEK293 LTV cells (Cell Biolabs) were cultured in DMEM medium (Gibco), supplemented with 10% FBS at 37 °C, 5% CO2. Human neural stem cells (NSCs) (Gibco) were cultured in NSC medium (KnockOut DMEM-F12 (Gibco), 2 mM L-glutamine (Gibco), 20 ng/ml bFGF (Peprotech), 20 ng/ml EGF (Peprotech), 2% StemPro Neural supplement (Gibco), 100 U/ml penicillin and 100 μg/ml streptomycin), as previously described [87].

Enhancer activity in STARR-seq reporter plasmids
For experimental validation in Fig. 1G, we randomly selected 22 DAEs that showed interaction with a target gene by HiC, and of which the target gene was expressed in neural stem cells, as indicated from our previously generated RNA-seq data (GSE137129 [87]). DAEs were amplified from genomic DNA and cloned into the STARR-seq plasmid (kind gift of A.Stark) [88] as previously described [65]. For the additional tested enhancer deletions (Additional file 1: Fig. S4), the obtained STARR-seq plasmids containing IRF2BPL, CHD2, and MACF1 enhancers were modified by site-directed mutagenesis to remove regions with high or low ncER score. The following regions were deleted: IRF2BPL (chr14: 77422484–77422514); CHD2 (ncER1 chr15: 93363603-93363640, ncER3 chr15: 93363780-93363790); MACF1 (ncER1 chr1: 39598824-39598844, ncER2 chr1:39598744-39598754). The regions with low ncER score at the 5’ and 3’ ends (80–100 bp) of IRF2BPL, CHD2, and MACF1 enhancers were excluded by Gibson assembly. Primer sequences are provided in Additional file 15: Table S14. HEK293 and NSC were transfected with STARR-seq plasmid containing enhancer regions using polyethyleneimine (PEI, Sigma) or
Lipofectamine™ Stem Transfection Reagent (Thermo Scientific) respectively. Spike-in of a pmCherry-N1 plasmid (Clontech) was used as a transfection control. Twenty-four hours post transfection cells were collected, stained with Hoechst dye and the enhancer activity was measured by FACS analysis (20,000 cells per sample). GFP-positive cells within the mcherry-positive population were quantified to assess enhancer activity compared to an empty STARR-seq vector. Two independent transfection experiments were performed, each in duplicates. Statistical analysis was performed using a one-way ANOVA test followed by multiple comparison test (Fisher’s LSD test). Calculations were conducted in GraphPad Prism (version 8).

dCas9-KRAB-MeCP2 silencing of active enhancers in NSC
We selected DAEs linked to CHD2, CAD, and TRAK1 and designed for each DAE two targeting gRNAs (primer sequences are given in Additional file 15: Table S14). gRNAs were cloned into a pgRGFP plasmid (Addgene #82695, a kind gift of Allan Mullen) [89]. NSCs were co-transfected with dCas9-KRAB-MeCP2 (Addgene #110824, kind gift of Alejandro Chavez and George Church) [90] and the two gRNAs/DAE and collected for RNA isolation 48 h post transfection. Transfection efficiency was estimated by FACS analysis (78–92% GFP-positive cells detected). RNA was isolated using TRI reagent (Sigma) followed by cDNA preparation using iSCRIPT cDNA synthesis kit (Bio-Rad). Fold change in gene expression (ΔΔct method) was evaluated by qPCR (iTaq universal SYBR Green Supermix) (Sigma), performed in CFX96RTS thermal cycler (Bio-Rad), as previously described [87]. TBP expression was used as housekeeping normalization control. Statistical analysis was performed using a one-way ANOVA test followed by multiple comparison test (Fisher’s LSD test). Calculations were conducted in GraphPad Prism (version 8).

Zebrafish studies
Zebrafish (Danio rerio) were raised and maintained under standard conditions [91]. Adult and larval fish were kept on a 14 h/10 h light–dark cycle at 28°C. Larvae were kept in HEPES-buffered E3 medium. Media was refreshed daily, and at 24 hpf, 0.003% 1-phenyl 2-thiourea (PTU) was added to prevent pigmentation. All zebrafish experiments were performed in compliance with Dutch animal welfare legislation. Selected DAEs used in the in vitro experiments were transferred by Gibson assembly between the AscI and PacI site of a E1b-GFP-Tol2 enhancer assay plasmid (a kind gift of Ramon Birnbaum) [92] containing an E1b minimal promoter followed by GFP, using the following transfer primers: Transfer_fw: 5′-AGATGGGCCCCTCGGGTAGAGCATGCACCGG-3′ and Transfer_rv: 5′-TCGAGAACTCTTTAATGGGCCCAGATTCCGTGCA-3′. Constructs were injected into zebrafish embryos using standard procedures, together with Tol2 mRNA to facilitate genomic integration. At least 50 embryos were injected per construct in at least two different injection experiments. GFP expression was observed and annotated at 1, 2, and 3 dpf by a fluorescent Leica M165FC stereomicroscope (Additional file 14: Table S13). Images were analyzed using ImageJ (FIJI). An enhancer was considered active when at least 30% of the larvae showed consistent GFP expression.

Results
Integrative data analysis identifies differentially active regions during fetal brain development
We started our analysis by collecting relevant fetal brain epigenome data sets and previously published putative enhancers (Additional file 2: Table S1, Additional file 3: Table S2). Epigenome data sets included ChIP-seq for various histone modifications, DNase- and ATAC-seq data from various developmental time points and anatomical regions of human fetal brain, generated by several independent studies, including Roadmap, PsychENCODE, and other publications [9, 10, 19, 31–35, 86]. All primary data were reanalyzed using identical computational pipelines, and in total we processed 494 data sets. Scrutinizing through previously published literature on enhancers in brain and neuronal cell types, we collected 1,595,292 putative brain enhancers (Additional file 2: Table S1). These included enhancers retrieved from various enhancer databases, such as VISTA, FANTOM, and EnhancerAtlas, enhancer predictions from the PsychENCODE consortium, human accelerated regions, ultra-conserved regions, and others [9, 11, 21–31]. We first analyzed the overlap between the different putative enhancers and found only a small overlap between enhancer predictions from different studies (Additional file 2: Table S1). We reasoned that if different enhancer prediction methods used in the individual studies identified the same enhancers that only differ by the exact location or length, by merging the overlaps between different studies we could identify functional relevant parts of enhancers. We thus proceeded to determine putative critical regions (pCRs), by determining the unifying overlaps between all putative enhancers (Fig. 1A, step 1). In this analysis, we kept those putative enhancers that were only found in a single study, merged the overlaps of multiple studies and eliminated those regions that were located within 2 kb upstream and 1 kb downstream of a transcriptional start site (TSS) or which had <10 reads in epigenome data (see “Methods”). This resulted in 202,163 pCRs, with a total length of 93 Mb, an average size of 460 bps, and
Fig. 1 (See legend on next page.)
most pCRs located between 5 and 50 kb away from the closest gene TSS (Additional file 1: Fig. S2A, B).

We assumed that enhancers that have functional relevant roles during brain development would show dynamic changes in the levels of histone modifications and chromatin accessibility correlating with their function. To investigate this, we next intersected all pCRs with all epigenome data sets from different time points of fetal brain development and calculated the read count for each pCR region. After TMM-normalization, we performed differential accessibility analysis (for ATAC-seq and DNase data) and generated differential histone modification profiles (for H3K27ac, H3K27me3, H3K4me1, H3K4me2, H3K4me3) using edgeR [50]. This resulted in 39,709 pCRs that showed a high variability for these features across developmental time points (Fig. 1B, Additional file 1: Fig. S2C, Additional file 4: Table S3, see “Methods”) which we refer to as differentially active enhancers (DAEs). In contrast, the remaining 162,454 pCRs showed a more constant epigenome pattern and we thus refer to them as not-differentially active enhancers, nDAEs (Fig. 1B, Additional file 1: Fig. S2C, Additional file 4: Table S3).

Gene ontology analysis using GREAT [17] showed that DAEs were significantly enriched for terms related to brain development, including processes such as forebrain neuron fate commitment, dorsal/ventral axon guidance, and spinal cord development (Fig. 1B, Additional file 5: Table S4). nDAEs appeared to be enriched for more general terms, including various chromatin modifications and receptor-mediated endocytosis (Fig. 1B, Additional file 5: Table S4).

To have a more specific view about the genes regulated by these pCRs, we next linked DAEs and nDAEs to their target genes, using different resources, which either link enhancer to gene promoters by direct chromatin interaction as determined by chromatin conformation capture techniques (HiC [30], HiChIP [55], PLAC-seq [56]) or by predicting enhancer-gene interactions using statistical models and correlation between gene expression, omics data, and epigenome features (JEME, FOCS, GeneHancer, ENCODE, Activity-by-contact (ABC) method) [51–54, 57] (Additional file 6: Table S5). Since only a limited number of interactions between DAEs or nDAEs and target genes identified by these different methods were supported by >2 of the available resources (Additional file 1: Fig. S3A), and as most interactions and target genes were predicted by the HiC data (Additional file 1: Fig. S3B), we focused on these HiC predicted target gene interactions for the remainder of the analysis. These HiC data were generated from post conceptional week (PCW) 17–18 human brains [30] and were available for the germinal zone (GZ) (containing primarily mitotically active neural progenitors), and the cortical and subcortical plate (CP) (consisting primarily of post-mitotic and migrating neurons). Enhancer-promoter interactions derived from these HiC data do not exclude the fact that the identified DAEs and nDAEs interact with or regulate other genes at other developmental time points or in other cell types, for which at this moment no specific enhancer-promoter predictions are available.

Taking only those enhancer-promoter interactions that occurred in the same topological associated domain (TAD) into account, we found that from all DAEs, 6858 and 6883 for CP and GZ, respectively, interacted with promoters of protein-coding genes or lincRNAs, of which the majority of interactions occur with protein-coding genes. Similarly, 27,004 and 27,161 nDAEs interacted with target genes in CP and GZ, respectively, with a similar distribution between protein-coding and lincRNA genes in CP (upper panel) and GZ (lower panel).
In total, DAEs interacted with 5946 and 6085 protein-coding and lincRNA genes in CP and GZ, respectively, of which 3841 genes were shared between both CP and GZ (Fig. 1C). The majority of these genes (86%) also had interactions with nDAEs (Fig. 1D). We next integrated available gene expression data from fetal and adult brain (Additional file 7: Table S6) and found that genes that interacted with a DAE had a significantly higher gene expression compared to those genes not interacting with a DAE, at various regions and stages of fetal development but not in adult brain (12 PCW: CP genes p-value = 0.002671, GZ genes p value = 5.111e−05; 15–17 PCW: CP genes p value = 0.003251, GZ genes p value = 0.003813; 17 PCW: CP genes p value = 0.002533, GZ genes p value = 0.001813); 81 years: CP genes p value = 0.1377, GZ genes p value = 0.2641; fetal sources mean: CP genes p value = 0.0002696, GZ genes p value = 0.00046; DAE fetal sources mean vs DAE 81 years: CP genes p value = 0.04744, GZ genes p value = 0.01525; nDAE fetal sources mean vs nDAE 81 years: CP genes p value = 0.781, GZ genes p value = 0.4904, wilcox.test) (Fig. 1E, Additional file 1: Fig. S3E). Similar observations were made when using the alternative, not HiC-based enhancer-promoter predictions (Additional file 1: Fig. S3F). In line with earlier findings [93], we find that the more enhancers a gene is interacting with, the higher the gene expression is, and this was also true for the DAEs (Fig. 1F). A recent study determined gene expression trajectories in the dorsolateral prefrontal cortex during pre- and postnatal development. This study identified constant, rising and falling genes, that showed respectively similar, increased or decreased gene expression levels upon development [60]. In line with the earlier gene expression findings, we found that the odds ratio between DAE and nDAE linked genes (as determined by HiC) was significantly higher (odds ratio = 1.183, p value = 0.0008 for GZ; odds ratio = 1.198, p value = 0.0004 for CP, Fisher’s exact test) for falling genes, that showed higher gene expression levels in pre-natal RNA-seq samples (Additional file 1: Fig. S3G,H).

Finally, to validate that DAEs can function as enhancers, we selected 22 DAEs linked to genes that are expressed in human neural stem cells (NSCs), cloned them in an enhancer reporter plasmid [88] and tested their enhancer activity in cell transfection experiments. Upon transfection in NSCs, 18 out of 22 tested sequences showed significantly increased percentage of GFP+ cells compared to control (normalized for transfection efficiency using an mCherry spiked-in control), confirming enhancer activity (Fig. 1G). Transfecting the same plasmids in non-neural HEK cells showed less pronounced activity. This indicates that 81.8% of the tested DAEs had a measurable enhancer activity using this assay in an in vitro neural cell type. Of note, this does not exclude activity of those 4 DAEs that do not show enhancer activity in NSCs, in other cell types during fetal brain development.

We conclude that an integrative data analysis of virtually all previously reported brain enhancers identifies a set of DAEs which are associated with a brain developmental gene ontology, increased gene expression in fetal brain and display enhancer activity in vitro.

Multi-gene-interacting enhancers regulate genes implicated in multiple cellular processes and have distinguishing sequence characteristics

In order to understand the biological function of DAEs and nDAEs in more detail, we further characterized these two groups. When determining the number of genes that each DAE is interacting with, we found that the majority of DAEs interact with 1 or 2 genes; but, in addition, a considerable fraction of DAEs also interact with more than 2 genes (19.7 % for CP, 19.4% for GZ) (Fig. 2A), and the same was found for nDAEs (Fig. 2B). When comparing the enrichment of biological processes for the genes that interact by HiC with these multi-gene-interacting DAEs using Enrichr, we found that these genes were enriched for broader developmental and metabolic processes. However, genes that interact with DAEs that only regulate single genes were enriched for more specific brain-related terms, such as “neuron differentiation” and “neuron migration” (Additional file 8: Table S7). Similar results were obtained using GREAT and Metascape analysis, where multi-gene-interacting DAEs for example were enriched in mouse phenotypes associated with “early lethality,” whereas DAEs associated with only a single gene were enriched for “regulation of neural precursor cell proliferation” (Additional file 8: Table S7).

We next asked whether DAEs that regulate single or multiple genes could have distinguishing DNA sequence characteristics, which could support their presumed distinct functional roles. To answer this, we focused on scores that provide some weight based on the underlying sequences: non-coding essential regulation (ncER) score [38], GC content [39], and phastcons score [39]. The ncER scores were recently established using a machine learning model [38], taking functional, mutational, and structural features into account, including sequence constraint in the human population, and provides a score where 0 is non-essential and 1 is putative-essential. We observe that DAEs that interact with 3 or more genes have a significantly higher ncER percentile compared to DAEs that interact with only 1 gene (Fig. 2C). This might reflect their biological function regulating multiple genes, resulting in a higher tendency to be constraint. A similar trend was observed for GC content, where DAEs interacting with more than one gene had a significantly
Fig. 2 (See legend on next page.)

Fig. 2: (a) Number of DAE and nDAE per gene per DAE. (b) Number of DAE and nDAE per gene per nDAE. (c) Enhancer type and enriched type analysis. (d) Enhancer type and enriched type analysis. (e) Enhancer type and enriched type analysis. (f) Enhancer type and enriched type analysis. (g) Density plot of LoF Score. (h) O/E Ratio All nDAE, O/E Ratio All DAE, O/E Ratio DAE_CP, O/E Ratio DAE_GZ.

Enhancer Type:
- DAE
- nDAE

Enriched Type:
- All Regions
- Linked to genes
- Linked to OMIM genes
higher GC content, whereas for the phastcons score, an indicator of multi-species conservation, differences were not significant (Fig. 2C). Similar observations were made for nDAEs (Fig. 2D). Higher GC content has also been observed in more broadly active enhancers in the immune system [94] and might be explained by binding of broadly active transcription factors (TFs) to GC-rich motifs [95].

Sequence characteristics distinguish DAEs from nDAEs

Given the differences in gene ontology between DAE and nDAE linked genes (Fig. 1B) and the differences in ncER score and CG content between enhancers that regulate single versus multiple genes (Fig. 2C, D), we next asked whether there are differences between these scores in DAEs and nDAEs, and whether any potential difference would be influenced by gene interactions that these regulatory elements have. We observed a significantly higher ncER percentile, CG content, and phastcons score when comparing all DAEs to nDAEs (Fig. 2E). Interestingly, some of these scores further increased, when only considering those DAEs and nDAEs that interact with target genes (as determined by HiC). This increased even further when only considering interacting target genes that are associated with known Online Mendelian Inheritance in Man (OMIM) phenotypes. Similar observations were made when using the Orion [40] and CADD scores [41] (Fig. 2E) that similarly depletation of variation in the human population and likelihood of deleteriousness of a given nucleotide based on integration of various annotations into account, respectively. Again, DAEs scored significantly higher for Orion and CADD scores than nDAEs, emphasizing the potentially biological important role of DAEs during brain development. Genes that are essential in humans are generally depleted of loss-of-function alleles, and this is reflected by a higher probability of loss-of-function intolerance (pLI) score [42]. When we plotted the median pLI of genes linked to DAEs, or to nDAEs, genes linked to DAEs scored significantly higher (Fig. 2F).

Finally, a recent study determined loss-of-function tolerance scores for non-coding sequences, by using machine learning and structural variants from whole genome sequencing, including homozygous enhancer deletions [43]. Using this analysis, we observed that DAEs were more likely to be intolerant to loss-of-function, whereas nDAEs were more often tolerant to loss-of-function (Fig. 2G). Again, when only considering those interactions linked to known target genes, scores further improved, in favor of DAEs.

We and others previously showed that functional enhancers can be enriched for transposable elements (TEs), some of which can be human specific [65, 96–98]. We thus asked whether DAEs and nDAEs showed a similar TE enrichment, and whether any TEs could distinguish both groups (Fig. 2H, Additional file 9: Table S8). nDAEs showed a small enrichment for various LTR-containing TEs (e.g., LTR75B, LTR60, LTR36). Compared to nDAEs, DAEs were mainly enriched for CG-rich repeat sequences, and a number of LTR repeats, such as Harlequin-int, HERVS71-int, and HERV K3-int. Enrichment of the latter LTR repeats was not seen when only considering gene-interacting DAEs. The MER130 repeat family was previously shown to be enriched near critical genes for the development of the mouse neocortex and suggested to be co-opted for developmental enhancers of these genes [99]. Interestingly, MER130 repeats were enriched in all DAEs, but this enrichment was lost when only assessing DAEs that interact with genes, which made it difficult to further investigate the role of MER130 in human brain regulation. Compared to our previous findings in human embryonic stem cells (ESCs) [65], the overall TE enrichment in enhancers in brain was markedly different.
Fig. 3 (See legend on next page.)
with none of the TEs enriched in active enhancers in ESC showing enrichment at brain enhancers. This could indicate that different TEs co-opted into the regulatory landscape acquired different tissue-specific roles during evolution.

Together this indicates that by investigating unbiased variability in epigenome marks over putative brain enhancers across developmental time points, DAEs and nDAEs can be identified which are associated with different gene ontologies, show different enrichments, have different sequence characteristics, and are distinctively linked to disease-relevant genes.

**DAEs and nDAEs are enriched for distinct transcription factor binding sites**

As the merging of pCRs and subsequent variability calling identified DAEs with distinct sequence characteristics, we next wondered whether we could further zoom in into each of the DAEs, to identify functional relevant nucleotides. To this end, we again made use of the ncER, CG, and phastcons scores, assuming that the functional relevant nucleotides in each DAE might be those that have higher scores. As the identified DAEs varied in size between 50 and 1000 bps, we first split each DAE into 10 bp bins and assigned the median ncER, CG, and phastcons scores to each bin. To be able to compare the score distribution within each bin between all DAEs, we re-scaled each DAE to a relative bin position from 1 to 100 (see “Methods” for details). Strikingly, the mean of ncER, CG, and phastcons scores were highest between bins 40 and 60 (Fig. 3A). To exclude that this was an artifact from the bin-rescaling, we plotted the mean distribution for the same scores also for DAEs that had an identical length and found similar results (Additional file 1: Fig. S4A). We next calculated the number of reads from all epigenome data sets and plotted the log2 enrichment over the same relative DAE bin positions. We found that ATAC-seq, DNase-seq, H3K27ac, H3K4me1, and H3K4me2 signals (all associated with enhancers) again were most enriched between bins 40 and 60, whereas signal for H3K4me3 (of which high levels are associated with promoters and lower levels are found at enhancers) and H3K27me3 (associated with repressed chromatin) showed a broader distribution (Fig. 3B), and this holds true for all developmental time points assessed. This suggests that on average, the center of the DAEs most likely contains the functional relevant sequences, and given the increased chromatin accessibility at those locations, this could indicate binding of functionally relevant TFs in these central regions.

To investigate this further, we first performed TF enrichment analysis using **Locus Overlap Analysis (LOLA)** [63], on both full length DAEs, as well as on only the central DAE parts between bins 40 and 60 (ncER subset). LOLA performs enrichment analysis based on genomic regions and tests the overlap of the query regions with a core reference database assembled from public data, including amongst others ChIP-seq data from ENCODE [100]. We found a similar enrichment of TF binding sites between full length and central parts of DAEs (Fig. 3C, Additional file 9: Table S8), and between all DAEs and those interacting with target genes in CP and GZ. The most enriched TFs at DAEs according to LOLA included well-known TFs with essential roles for brain development. This includes amongst others ETS1, a widely studied TF with functions in different biological systems which was previously shown to be necessary for radial glia formation in vertebrates [101] and FGF-dependent patterning of anterior-posterior compartments in the central nervous system of *Ciona* (a marine invertebrate that is a well-suited model to study cell fate specification in chordates) [102]; YY1, a crucial TF which is involved in both gene activation and repression [103], mediating enhancer-promoter interactions [104] and of which mutations cause a neurodevelopmental disorder [105]; and CTCF, a master regulator of chromatin...
Fig. 4 (See legend on next page.)
structure, of which de novo mutations cause intellectual disability [106]. We next repeated the same analysis for nDAEs (Fig. 3D–F, Additional file 9: Table S8). Similar to our observations for DAEs, nDAEs had higher ncERs, CG content, and conservation at the central part, with those regions being enriched for enhancer associated histone marks, but showed less variability over time. When performing TF enrichment analysis using LOLA, we observed a different and less specific set of TFs enriched at nDAEs compared to DAEs. Also, enrichment was lower at those nDAEs that were interacting with target genes. Again similar enrichment was found in the central part compared to the whole nDAEs. Enriched TFs for nDAEs included amongst others FOXL1, a transcriptional repressor that regulates central nervous system development [107]; the LIM homeodomain TF LHX3, that is essential for pituitary and nervous system development [108, 109]; and FOXA2, which plays a role in midbrain dopaminergic neurons [110, 111] (Fig. 3F). Shared TFs enriched both at DAEs and nDAEs included SP1, loss of which in astrocytes impacts on neurons in the cortex and hippocampus of mice [112]; MAFB, a basic leucine zipper TF that plays a role in hindbrain development [113–115] and postnatal brain development [116, 117]; and ZEB1 which is required for neuronal differentiation [118, 119].

As LOLA analysis considers a single shared base pair being sufficient for regions to count as overlapping, this analysis could not distinguish well between TFs specifically enriched at the central part of DAEs and nDAEs relative to the flanking regions. We therefore further investigated which TFs motifs were specifically enriched at the central parts versus other parts of DAEs and nDAEs, using motif enrichment analysis with HOMER [64], a motif discovery algorithm, which identifies regulatory elements that are specifically enriched in the query set relative to background. We first split the 100 relative bins into 20 groups of 5 consecutive bins each and determined the significantly enriched TF motifs \((p \leq 0.01)\) for each of these 20 bin groups (Additional file 9: Table S8). Amongst the enriched motifs, we found back, amongst others, the motifs for the TFs enriched using the LOLA analysis, validating these findings (Additional file 9: Table S8). When plotting the number of significant motifs \((p \leq 0.01)\) per bin group and the number of target sequences with those motifs, we found that bins located in the central part of both DAEs and nDAEs had both the highest numbers of significant TF motifs and the highest number of target sequences (Additional file 1: Fig. S4B,C). As most enriched motifs were found in multiple bins, and there can be multiple TF bindings sites of the same TF within the same enhancer, we next focused on only those TF motifs which were not equally enriched in all 20 bin groups \((n = 251\) for DAEs and \(n = 218\) for nDAEs). For both DAEs and nDAEs, we again found most motif enrichment in the central enhancer part, with DAEs being more enriched than nDAEs (Fig. 3G, H, Additional file 1: Fig. S4D,E). Amongst the most enriched TF motifs at the center of DAEs were motifs for the proneural basic helix-loophelix transcription factors NEUROG2, ATOH1, and NEUROD1 that promote neurogenesis [120–122], OLIG1, a marker of oligodendrocytes [123] that also regulates the neuron-glial switch during earlier embryonic development [124, 125], TCF4, that is necessary for neuronal migration and the correct development of the cerebral cortex [126] and loss of which is associated with intellectual disability [127], and NF1 that regulates neuronal and glial differentiation and is causative of neurofibromatosis type 1 when mutant [128] (Fig. 3G). Enriched TF motifs at the central part of nDAEs are involved in more ubiquitous processes and include mainly activator protein 1 (AP-1), a heterodimer composed of members of the JUN (including JUNB), FOS (including FOSL2, FRA1, FRA2), ATF (including ATR3, BAFT), and MAF family that regulates a wide variety of cellular processes in response to a wide range of extracellular cues [129] (Fig. 3H). Together this indicates that on average the central part of brain enhancers (both DAE and nDAEs) contains relevant but partially distinct TF binding sites and might be enriched for functional relevant sequences, which can be further fine-mapped using ncER scores and other sequence characteristics. To test this directly, we selected three DAEs, linked to IRF2BP1, CHD2, and MACF1, that showed activity in reporter assays in NSCs (Fig. 1G) and deleted 10–30 bp of those regions that had the highest ncER scores in those enhancers. Upon transfection of
Fig. 5 (See legend on next page.)

**a**

- **OMIM phenotype**
  - Mental retardation
  - Deafness
  - Developmental and epileptic encephalopathy
  - Intellectual developmental disorder
  - Cerebellar ataxia
  - Cardiomyopathy
  - Charcot–Marie–Tooth disease
  - Neurodevelopmental disorder
  - Immunodeficiency
  - Nephrotic syndrome
  - Retinitis pigmentosa
  - Citrullinemia
  - Usher syndrome
  - Spastic paraplegia
  - Epilepsy
  - Prostate cancer
  - Leukemia
  - Left ventricular noncompliance
  - Encephalopathy
  - Diabetes mellitus
  - Cataract
  - Lymphoma
  - Microcephaly
  - Amyotrophic lateral sclerosis
  - Mitochondrial complex I deficiency
  - Microphthalmia
  - Spermatogenic failure
  - Long QT syndrome
  - Colorectal cancer
  - Glutaric aciduria type 1
  - Ehlers–Danlos syndrome
  - Leber congenital amaurosis
  - Lissencephaly
  - Hypoplasia
  - Thyroid dysgenesis
  - Concerns with muscular dystrophy
  - Combined oxidative phosphorylation deficiency
  - Hypothyroidism
  - Pancreatic ductal adenocarcinoma
  - Alport syndrome
  - Alzheimers disease

**b**

- **GWAS phenotype**
  - Cortical surface area (min-P)
  - Subcortical brain region volumes
  - Nucleus accumbens volume
  - Multiple sclerosis and systolic blood pressure (genotype)
  - Thalamus-circulate cortex volume
  - Cortical thickness
  - Corpus callosum volume
  - Hippocampus volume
  - Nigrostriatal volume
  - Whole grey matter microstructure (white matter)
  - Whole grey matter microstructure (white matter)
  - Cognitive performance
  - Cognitive aspects of educational attainment
  - Cognitive performance (MTAG)
  - Whole grey matter microstructure (radial diffusivity)
  - Whole grey matter microstructure (radial diffusivity)
  - Hipocampus volume
  - Depression (broad)

**c**

- **Regions**
  - **pvalue**
    - de novo: 0.003**
    - inherited CNVs: 0.208

**d**

- **GABRD enhancer**
  - chr1: 1,839,541 - 1,963,600
  - **Deletion**
  - **RNA-seq**

**e**

- **Regions**
  - **pvalue**
    - DAE_CP: 0.157
    - DAE_GZ: 0.021*

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**Legend**

- **OMIM phenotype**
- **GWAS phenotype**
- **Regions**
- **pvalue**
these mutant DAEs, we observed a significantly reduced enhancer activity for IRF2BPL and CHD2, but not for MACF1 (Additional file 1: Fig. S4F). Deleting regions with a lower ncER score did not affect enhancer activity. Together, this indicates that integrative analysis, variability analysis during development, and sequence characteristics can identify functional relevant nucleotides in brain enhancers.

**DAEs show temporal epigenome dynamics during human brain development**

To further understand the dynamics of enhancer regulation, we subdivided DAEs interacting with genes in GZ and CP by performing clustering analysis on all available epigenome data sets, at different developmental stages (between 8 and 12 PCW, 13–18 PCW, and > 18 PCW) (Fig. 4A, Additional file 10: Table S9). At 8–12 PCW, we found two clusters for both GZ and CP that showed relatively constant enrichments over time, with the first cluster (red) showing a higher enrichment for all epigenome features available for that developmental stage, compared to the second cluster (green). No statistically significant differences in gene expression levels between genes linked to both clusters were found. Genes associated with cluster 1 DAEs in CP were enriched for gene ontology terms related to neuronal differentiation, whereas cluster 2 was dominated by processes in the Golgi. Likewise, for GZ, genes associated with cluster 1 seemed to be associated with more specific biological functions, whereas processes associated with cluster 2 showed more broad involvements (Fig. 4A, Additional file 10: Table S9).

At 13–18 PCW, three clusters emerged in both GZ and CP (Fig. 4B, Additional file 10: Table S9). Whereas cluster 3 (green) showed relatively low levels of epigenome marks similar to cluster 2 at 8–12 PCW, cluster 1 (red) and cluster 2 (blue) showed higher epigenome enrichments. Both cluster 1 and 2 showed similar levels of H3K27ac, but mainly diverged from each other on the levels of H3K4me3. Cluster 2 was strongly enriched for processes involved in neural system development both in CP and GZ. Gene ontology of genes associated with cluster 1 (red) which showed higher H3K4me3 levels, showed enrichment for insulin-like growth factor receptor signaling and immune cell-related processes in CP. Insulin-like growth factors are important for neuronal survival and neurogenesis [130]. As high levels of H3K4me3 have also been found at enhancers in blood cells [131], possibly stabilizing their transcription, it is tempting to speculate that part of this cluster reflects enhancers active in hematopoietic cells from the developing vasculature [132] and microglia (brain tissue macrophages) that are invading the brain at these developmental time points [133]. In GZ, cluster 1 was associated with phosphatidylinositol 3-kinase signaling, which is important for commitment of neural progenitor cells [134, 135].

Finally, at > 18 PCWs, we found two clusters of DAEs, of which cluster 1 (red) was marked by higher levels of epigenome marks (Fig. 4C, Additional file 10: Table S9). In CP, genes associated with this cluster were enriched for carboxylation processes and insulin-like growth factor receptor signaling. Genes associated with the second cluster (green) were again more enriched for broad developmental processes, including the Golgi system. In GZ, genes associated with cluster 1 (red) were amongst others involved in DNA damage repair. Indeed, alterations in this pathway can lead to reduced proliferation of neural progenitor cells leading to microcephaly [136, 137]. Cluster 2 (green) in GZ was associated with terms related to neurodevelopment and organ morphogenesis.

Together, this shows that temporal epigenomic rearrangement in DAEs is reflected in regulating the expression level of genes that are important in developmental and cell type-specific processes.

**Cell type specificity of DAEs and nDAEs and their dynamics in adult brain**

To further investigate cell type specificity of DAEs and nDAEs, we performed two additional analyses. First, we compared DAEs and nDAEs to recently identified cell type-specific regulatory elements. A recent study used scATAC-seq to generate a human cell atlas of fetal chromatin accessibility spanning 15 organs, including fetal brain [85]. When overlapping DAEs and nDAEs to the
Fig. 6 (See legend on next page.)
most specific chromatin accessibility peaks per cell type, we found 7753 DAEs and 7946 nDAEs that overlapped with these cell type-specific chromatin accessibility peaks, including those found in several types of neurons and astrocytes (Additional file 1: Fig. S5A,B). This indicates that our bulk analysis re-identifies cell type-specific chromatin accessibility peaks which might therefore present cell type-specific enhancers, and it also shows that the bulk analysis identifies additional enhancers that are not captured by the single-cell chromatin accessibility profiles.

We next investigated how these cell type-specific enhancer might behave over time. Two recent studies determined cell type-specific regulatory elements from postnatal brain with a reasonable overlap between both studies (Additional file 1: Fig. S5C,D), by either isolating cell type-specific bulk populations from brain followed by ATAC-seq and ChiP-seq for H3K27ac and H3K4me3 [56], or by performing scATAC-seq [55]. Comparing the DAEs and nDAEs to these cell type-specific regulatory elements showed as expected that only a fraction of DAEs and nDAEs from the fetal brain analysis showed an overlap with the cell type-specific regulatory elements derived from postnatal samples. Amongst those, we found overlap with cell type-specific regulatory elements from neurons, oligodendrocytes, astrocytes, and microglia (Additional file 1: Fig. S5E,F). This indicates that despite determined from an integrative analysis of bulk samples derived during fetal brain development, a fraction of DAEs and nDAEs can be linked to cell type-specific regulatory elements which are likely to also have roles in postnatal brain. In contrast, other DAEs and nDAEs are likely having fetal-specific functions.

To further investigate the dynamics of DAEs and nDAEs in adult brain, in the second analysis, we compared H3K27ac levels obtained from both fetal and adult samples derived from a single study [86] for all DAEs and nDAEs linked to target genes in GZ and CP by HiC, and performed clustering and gene ontology analysis (Additional file 1: Fig. S6, Additional file 10: Table S9). We found that DAEs that were mainly enriched for H3K27ac in fetal samples were as expected associated with bone ontology terms related to fetal brain development, including regulation of neuron differentiation. DAEs which also showed H3K27ac enrichment in adult samples were associated with more broad physiological processes.

Together, this shows that part of DAEs and nDAEs can be linked back to cell type-specific regulatory elements despite being identified from bulk tissue analysis and that some DAEs and nDAEs are likely to also function in postnatal brain.

**DAEs regulate disease-relevant genes and are enriched for disease implicated variants**

Given our findings that DAEs are associated with genes relevant for brain development, we further investigated which disease-relevant genes are regulated by DAEs. We first focused on known disease causing genes retrieved from OMIM. We found that 1556 OMIM genes are regulated by DAEs (of which 1165 and 1166 from the interactions found in GZ and CP, respectively) (Additional file 11: Table S10). Most DAEs are linked to genes involved in mental retardation, developmental and epileptic encephalopathy, and neurodevelopmental disorders (Fig. 5A). This included genes like KMT2C, involved in Kleefstra syndrome (OMIM #617768), and GRIN2A of which heterozygous mutations cause epilepsy and speech delay (OMIM #245570). Next to genes, enhancers can also interact with other additional enhancers. Interestingly, the more additional enhancers (DAE and/or nDAE) a DAE was interacting with, the more likely the
target gene of this DAE was an OMIM gene (Additional file 1: Fig. S7A). This supports recent findings that the number of enhancers linked to a gene reflects its disease pathogenicity [138] and confirms enhancer redundancy for disease-relevant genes [139].

We next leveraged published GWAS loci for brain-related traits and disorders (Additional file 12: Table S11). When comparing the odds ratio between DAEs and nDAEs, we found that DAEs were more often enriched for various significant GWAS loci, reflecting a broad variety of both brain developmental processes (e.g., volumes of different anatomical brain regions) and neurodevelopmental disorders (e.g., mental development, autism) (Fig. 5B). Similarly, using LD score regression analysis we found enrichment of heritability for variants within DAEs, nDAEs, and pCRs, including for the trait “intelligence” (Additional file 1: Fig. S7B).

Encouraged by these findings, we next asked whether copy number variants (CNVs) or single-nucleotide variants (SNVs) at DAEs could be involved in causing genetic disease. We first leveraged previously published disease implicated CNVs. Brandler et al. performed WGS in their discovery cohort of individuals affected by an autism spectrum disorder (ASD) and unaffected individuals and reported on 135 de novo CNVs (104 deletions, 29 duplications, and 2 inversions) [44]. Of these, 25 overlapped a DAE in cases, and 8 in controls (odds ratio = 2.10, \( p = 0.144101 \)). When only considering those CNVs overlapping DAEs linked to target genes, this became 17 in cases and 1 in control for DAEs linked to CP genes (odds ratio = 11.83, \( p = 0.003003 \)) and 15 in cases and 1 in control for DAEs linked to GZ genes (odds ratio = 10.14, \( p = 0.010423 \)). For nDAEs, 36 CNVs were found in cases and 15 in controls (odds ratio = 1.63, \( p = 0.267964 \)). However, as not all these CNVs exclusively covered non-coding regions, it cannot be excluded that the observed association is due to disrupted coding genes, rather than involvement of DAEs. We therefore also assessed rare inherited deletions from the same study that did not overlap with coding exons (\( n = 213 \) in total, 175 in cases and 38 in controls). From these, 32 cases had a deletion covering a DAE, compared to two controls (odds ratio = 4.027972, \( p = 0.05119 \)). Although not significant, this might point to more deletions covering DAEs in ASD individuals but would require a larger sample size to be confirmed (Fig. 5C, Additional file 13: Table S12).

In another study, Monlong et al. [45] reported on CNVs in 198 epilepsy patients detected by WGS. They found an enrichment of rare non-coding CNVs near known epilepsy genes, with the GABRD gene showing the strongest and only nominally significant association with 4 non-coding deletions amongst the epilepsy patients. Interestingly, a 4999 bp deletion reported in that study, overlapped with a 386 bp DAE which is located ~110 kb upstream of GABRD and which interacts with its promoter (Fig. 5D). Hence, it is possible that deletion of this DAE affects GABRD expression, which might be implicated in the phenotype of that individual.

Third, we made use of de novo SNVs found in WGS from 1790 ASD simplex families [46]. We found 932 de novo variants that overlapped all DAEs in ASD individuals compared to 829 variants overlapping all DAEs in unaffected individuals (odds ratio = 1.07, \( p = 0.157 \)). We next repeated the analysis with only those DAEs that are interacting with known autism genes from the SFARI Gene database (\( n = 1003 \) genes) [84]. We found 26 cases and 11 controls with de novo variants in DAEs that interact with autism genes in CP (odds ratio = 2.249703, \( p = 0.021455 \)), whereas for DAEs interacting with autism genes in GZ, this was 20 cases and 17 controls (odds ratio = 1.11955, \( p = 0.745628 \)) (Fig. 5E, Additional file 13: Table S12). Interestingly, for each of the genes CIB2, FBRSL1, PAC2S2, KDM4B, and MYT1L, we found 2 individuals with autism with de novo variants in DAEs interacting with these genes. These variants are either absent or extremely rare in a large control cohort of gnomAD [140], possibly pointing to a role in causing the phenotype, although this will require further validation.

Together this indicates that DAEs are linked to disease-relevant genes and are enriched for GWAS loci relevant for brain-related traits and for variants linked to genetic disorders.

**CRISPRi and zebrafish experiments confirm enhancer activity of DAEs regulating genes involved in epileptic encephalopathy**

To further substantiate our findings, we validated the biological role of selected enhancers, using in vivo zebrafish transgenic reporter assays and CRISPR inhibition in human NSCs by focusing on enhancers linked to disease-relevant genes.

CHD2 belongs to the chromodomain helicase DNA-binding families of chromatin remodeling proteins, and haploinsufficiency of this gene has been associated with a developmental and epileptic encephalopathy, presenting with early onset intractable seizures, cognitive regression, intellectual disability and ASD behaviors (OMIM #615369) [141]. Around 80 kb upstream of CHD2, we found a DAE that interacts with the CHD2 promoter (Fig. 6A). In NSC reporter assays, this region showed strong enhancer activity, and this was less pronounced in non-neural HEK cells (Fig. 1G). To further study the biological relevance of this region, we first tested enhancer activity in vivo using zebrafish transgenesis. Out of the 36 analyzed zebrafish larvae, 61.1% showed GFP expression in the forebrain at 1 day post
fertilization (dpf), and this increased to 81.8% at 2dpf and 87.9% at 3dpf, indicating enhancer activity (Fig. 6B, C). Expression was also found in midbrain and hindbrain, at a slightly lower extent, in the eyes, in peripheral neurons, and in the spinal cord (Additional file 14: Table S13). GFP expression in the developing zebrafish brain correlated with in situ hybridizations of endogenous chd2 [142]. To test whether epigenome silencing of this enhancer would affect CHD2 expression, we performed CRISPR interference (CRISPRi) by targeting dCas9-KRAB-MeCP2 to the enhancer region by coexpression of gRNAs with a GFP fluorescent reporter. Transfection efficiency in these experiments, based on FACS for GFP, was 78–92%, and this resulted in around 50% reduction of CHD2 expression compared to mock cells transfected solely with dCas9-KRAB-MeCP2 (Fig. 6D). Interestingly, it was previously shown that silencing of CHD2 leads to reduced expression of REST [143]. In agreement with this, cells with reduced CHD2 expression upon CHD2 enhancer silencing showed reduced REST expression (Fig. 6E). This confirms that CHD2 is under control of the investigated DAE.

Bi-allelic variants in CAD cause an early infantile epileptic encephalopathy (OMIM #616457) [144] that is characterized by global developmental delay, loss of skills, therapy refractory epilepsy, brain atrophy, and dyserythropoietic anemia. We found an enhancer located in the third intron of EMILIN1, around 135 kb upstream of CAD that interacts with the CAD promoter (Fig. 6A) and which showed strong enhancer reporter activity in NSCs and only limited activity in HEK cells (Fig. 1G). Targeting this region in NSCs by CRISPRi significantly diminished gene expression of CAD to around 50% compared to mock (Fig. 6D). Similar to CHD2, in vivo reporter assays in zebrafish recapitulated in situ hybridisation results for cad [145]. From the 45 analyzed larvae, GFP expression was found in the forebrain of 88.9% larvae at 1 dpf, which remained ~85% at 2 and 3 dpf. Again, GFP expression was observed also in midbrain, hindbrain, eyes, in peripheral neurons, notochord, and spinal cord (Fig. 6B, Additional file 1: Fig. S8, Additional file 14: Table S13). These included enhancers linked to the disease genes MACF1 (OMIM #618325) and TUBB2A (OMIM #615763), of which coding pathogenic mutations cause brain malformations [148, 149], and C12orf4 (OMIM #618221) of which bi-allelic variants cause intellectual disability [150]. Together, this shows that DAEs identified in this integrative analysis show enhancer activity in vitro and in vivo and regulate, amongst others, genes linked to Mendelian disorders.

**Discussion**

Understanding the role of NCREs in development and disease still needs a significant effort at multiple levels: starting from identifying and annotating NCREs to investigating their target gene(s) and function. In the past few years, the identification and annotation of NCREs have gained a lot of attention. However, despite these developments, due to their sheer number and complex function, more studies and concerted efforts are needed to understand the role of NCREs in development and disease. Here we performed an integrative analysis of virtually all previously described putative enhancers and epigenome datasets of relevance for human brain development.

Our analysis has allowed us to first identify the intersection between previous studies and identify a list of putative NCREs. This is an important step as the different regions that were identified by previous investigations often have slightly different coordinates, length, and quality. Our putative regions are thus the commonality between all the different studies that are conducted hitherto, but at the same time keep the originality in each of them. To further specify enhancers that might...
have a biological relevance, mapping epigenomic data to these putative regions allowed us to identify around 40 thousand enhancers that display epigenomic rearrangements during human brain development. These DAEs have different sequence characteristics compared to non-variable enhancers, are bound by distinct sets of TFs, regulate disease-relevant genes, and can harbor non-coding variants that are associated with human disease. Furthermore, our integrative analysis identified a large number of enhancers linked to known disease genes and expands on the knowledge of regulation of these genes. For example, CHD2 expression regulation has so far only been known to be influenced by a highly conserved long non-coding RNA (lncRNA) referred to as CHD2 Adjacent Suppressive Regulatory RNA (CHASERR), which is located in proximity to the CHD2 TSS, and which represses Chd2 gene expression in cis [151]. It has been hypothesized that targeting CHASERR could be used to increase expression of CHD2 in haploinsufficient individuals [151], and it will be interesting to explore whether targeting the enhancer region of CHD2 that we find and validate here could be exploited as an alternative target of such a strategy. Similarly, the regulation by enhancers of other disease implicated genes that we validate here adds to the list of potential targets to find disease causing non-coding variants that disturb this regulation.

An interesting finding of our study is that by starting with putative enhancers and variability of epige-nome features over time during development, we recover DAEs and nDAEs that can be distinguished based on sequence characteristics, such as differences in GC content, the level of sequence constraint, tolerance to loss-of-function, and differential profiles of TF binding. Also, these DAEs and nDAEs seem to be associated with distinct developmental processes and result in differences in gene expression levels. It is tempting to speculate that the distinctive features between these two types of enhancers can be used to uncover key nucleotides responsible for those biological regulatory differences. It seems plausible that disturbing these functionally causative sequences could lead to altered physiology resulting in disease. Our analysis revealing GWAS loci enrichment and the link of DAEs supports this statement. We suggest that our results might help interpreting the effects of SNVs in non-coding sequences, which is at this stage not a trivial task. Our annotated database of DAE and nDAE will be instrumental to prioritize SNVs based on distinct sequence characteristics identified for these elements as well as to provide cues on potentially disturbed developmental processes based on differential temporal activity and regulatory targets of the enhancer in question. This in turn can instruct functional validation and help deciphering pathogenicity of variants. With an increasing number of whole genome sequencing data available, it is expected that more, possibly disease implicated, non-coding variants will be identified, and the need to classify those sequences in benign or pathogenic will only further increase. With more computational pathogenicity prediction tools available, such as the ncER score and outcomes of integrative analyses such as performed here that pinpoint likely functional sequences, it might become possible to further decipher the impact of these SNVs.

Conclusions
In this study, by using an integrative computational analysis of virtually all previously described putative enhancers and epigenome datasets, we identified a comprehensive compendium of likely functional enhancers that are involved human brain development and disease. By applying CRISPRi-based silencing and zebrafish enhancer reporter assays, we show that these putative regions possess enhancing characteristics. We foresee that these enhancer sequences will be instrumental in identifying disease causing variants which might explain parts of the missing heritability in the field of clinical genetics.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s13073-021-00980-1.

Additional file 1: Fig. S1: List of all putative enhancers collected Additional file 2: Table S1: Overview of all epigenome data processed in this study Additional file 3: Table S2: Overview of all RNA-seq data sets used in this study Additional file 4: Table S3: List of all pCRs, with DAEs and nDAEs indicated Additional file 5: Table S4: Functional enrichment analysis using GREAT for DAEs and nDAEs Additional file 6: Table S5: enhancer-gene predictions Additional file 7: Table S6: Overview of all RNA-seq data sets used in this study Additional file 8: Table S7: Functional enrichment analysis using GREAT, Enrichr and Metascape for multigene interacting DAEs and nDAEs Additional file 9: Table S8: TF and TE enrichment at DAEs and nDAEs Additional file 10: Table S9: DAE clusters and associated functional enrichment using Enrichr Additional file 11: Table S10: DAEs linked to OMIM genes Additional file 12: Table S11: Significant GWAS loci used in this study Additional file 13: Table S12: Calculations and p-values for gene variant associations
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- Data access from PsychENCODE was obtained for the following data sets: SynapseID: syn12033248 [9]; SynapseID: syn17092080 [86]. Data were generated as part of the PsychENCODE Consortium supported by: U01MH103339, U01MH103365, U01MH103392, U01MH103340, U01MH103346, R01MH105472, R01MH104971-04, R01MH103809, R21MH102791, R21MH105881, R21MH103877, and 5P01MH106934 awarded to: Schahram Akbarian (Icahn School of Medicine at Mount Sinai), Gregory Crawford (Duke), Stella Dracheva (Icahn School of Medicine at Mount Sinai), Peggy Farnham (USC), Mark Gerstein (Yale), Daniel Geschwind (UCLA), Thomas M. Hyde (LBBD), Andrew Jaffe (LBBD), James A. Knowles (UCSC), Patrick Sullivan (UCSC), Flora Vaccarino (Yale), Sherman Weissman (Yale), Kevin White (UCchicago) and Peter Zandi (JHU).

Authors’ contributions
SY performed primary computational analysis, with help of RD, KL, AN, and RD performed zebrafish validation experiments. SY, EM, and TSB wrote the manuscript with input from all authors. All authors read and approved the final manuscript. EM and TSB conceived and jointly supervised the work.

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Availability of data and materials
All data generated in this study is included in this published article and its supplementary files. Tables Additional file 2: Table S1, Additional file 3: Table S2 and Additional file 7: Table S6 summarizes all primary data used for the meta-analysis. Some of the primary data that were used to support the findings of this study are available from dbGaP and PsychENCODE, but restrictions apply to the availability of these data, which were used under license for the current study and so are not publicly available (third party data). The source codes and all processed data for all analysis performed in this study are available in the repositories https://github.com/syousefi87/Differentially-Active-Enhancers [152], and https://fsgshare.com/projects/Differentially-Active-Enhancers/122965 [153].

Declarations

Ethics approval and consent to participate
For zebrafish studies, no larvae older than 5 days post fertilization were used. Zebrafish were kept according to guidelines of the Erasmus MC animal welfare office and all zebrafish experiments were performed in compliance with Dutch animal welfare legislation.

Consent for publication
Not applicable

Competing interests
The authors declare that they have no competing interests.

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