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Structural brain abnormalities in the common epilepsies assessed in a worldwide ENIGMA study

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Progressive functional decline in the epilepsies is largely unexplained. We formed the ENIGMA-Epilepsy consortium to understand factors that influence brain measures in epilepsy, pooling data from 24 research centres in 14 countries across Europe, North and South America, Asia, and Australia. Structural brain measures were extracted from MRI brain scans across 2149 individuals with epilepsy, divided into four epilepsy subgroups including idiopathic generalized epilepsies (n = 367), mesial temporal lobe epilepsies with hippocampal sclerosis (MTLE; left, n = 415; right, n = 339), and all other epilepsies in aggregate (n = 1026), and compared to 1727 matched healthy controls. We ranked brain structures in order of greatest differences between patients and controls, by meta-analysing effect sizes across 16 subcortical and 68 cortical brain regions. We also tested effects of duration of disease, age at onset, and age-by-diagnosis interactions on structural measures. We observed widespread patterns of altered subcortical volume and reduced cortical grey matter thickness. Compared to controls, all epilepsy groups showed lower volume in the right thalamus

(Cohen's d = -0.24 to -0.73; $P < 1.49 \times 10^{-4}$), and lower thickness in the precentral gyri bilaterally (d = -0.34 to -0.52; $P < 4.31 \times 10^{-6}$). Both MTLE subgroups showed profound volume reduction in the ipsilateral hippocampus (d = -1.73 to -1.91, $P < 1.4 \times 10^{-19}$), and lower thickness in extrahippocampal cortical regions, including the precentral and paracentral gyri, compared to controls (d = -0.36 to -0.52; $P < 1.49 \times 10^{-4}$). Thickness differences of the ipsilateral temporopolar, parahippocampal, entorhinal, and fusiform gyri, contralateral pars triangularis, and bilateral precuneus, superior frontal and caudal middle frontal gyri were observed in left, but not right, MTLE (d = -0.29 to -0.54; $P < 1.49 \times 10^{-4}$). Contrastingly, thickness differences of the ipsilateral pars opercularis, and contralateral transverse temporal gyrus, were observed in right, but not left, MTLE (d = -0.27 to -0.51; $P < 1.49 \times 10^{-4}$). Lower subcortical volume and cortical thickness associated with a longer duration of epilepsy in the all-epilepsies, all-other-epilepsies, and right MTLE groups (beta, b < -0.0018; $P < 1.49 \times 10^{-4}$). In the largest neuroimaging study of epilepsy to date, we provide information on the common epilepsies that could not be realistically acquired in any other way. Our study provides a robust ranking of brain measures that can be further targeted for study in genetic and neuropathological studies. This worldwide initiative identifies patterns of shared grey matter reduction across epilepsy syndromes, and distinctive abnormalities between epilepsy syndromes, which inform our understanding of epilepsy as a network disorder, and indicate that certain epilepsy syndromes involve more widespread structural compromise than previously assumed.

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Abbreviations: ENIGMA = Enhancing Neuro Imaging Genetics through Meta-Analysis; IGE = idiopathic generalized epilepsy; MTLE-L/R = mesial temporal lobe epilepsy with left/right hippocampal sclerosis

Introduction

Epilepsy is a prevalent neurological disorder, comprising many different syndromes and conditions, affecting 0.6–1.5% of the population worldwide (Bell *et al.*, 2014). Approximately one-third of affected individuals do not respond to antiepileptic drug therapy (French, 2007). Alternative treatment options may not be appropriate (Englot *et al.*, 2011), and are not always effective (Téllez-Zenteno *et al.*, 2005; Englot *et al.*, 2011). The identification of shared biological disease pathways may help elucidate diagnostic and prognostic biomarkers and therapeutic targets, which, in turn, could help to optimize individual treatment (Pitkänen *et al.*, 2016). However, disease biology remains unexplained for most cases—especially in commonly occurring epilepsies.

Epilepsy is a network disorder typically involving widespread structural alterations beyond the putative epileptic focus (Bernhardt et al., 2015; Vaughan et al., 2016). Hippocampal sclerosis is a common pathological substrate of mesial temporal lobe epilepsy (MTLE), but extrahippocampal abnormalities are also frequently observed in MTLE, notably in the thalamus (Keller and Roberts, 2008; Coan et al., 2014; Alvim et al., 2016) and neocortex (Keller and Roberts, 2008; Bernhardt et al., 2009b, 2010; Blanc et al., 2011; Labate et al., 2011; Vaughan et al., 2016). Neocortical abnormalities are also reported in idiopathic generalized epilepsies (IGE) (Bernhardt et al., 2009a), and many childhood syndromes (O'Muircheartaigh et al., 2011; Vollmar et al., 2011; Ronan et al., 2012; Overvliet et al., 2013). Thus, common epilepsies may be characterized by shared disturbances in distributed cortico-subcortical

brain networks (Berg *et al.*, 2010), but the pattern, consistency and cause of these disturbances, and how they relate to functional decline (Vlooswijk *et al.*, 2010; Bernasconi, 2016; Nickels *et al.*, 2016), are largely unknown.

Currently, we lack reliable data from large cross-sectional neuroimaging, brain tissue, or biomarker studies in the common epilepsies. Brain tissue is not available from large cohorts of patients: common forms of epilepsy are often unsuitable for surgical treatment, so biopsied tissues are simply unavailable in sufficient numbers for research into disease biology. Brain-wide post-mortem studies also require extensive effort for comprehensive analysis. MRI offers detailed information on brain structure, but MRI measures from groups of individuals with and without epilepsy are not always consistent. For example, MTLE is associated with hippocampal sclerosis in up to 70% of brain MRI scans (Blümcke et al., 2013). However, the effects of laterality, and the extent of extrahippocampal grey matter loss are inconsistently reported in studies of left versus right MTLE (Kemmotsu et al., 2011; Liu et al., 2016). Similarly, abnormalities of the basal ganglia, hippocampus, lateral ventricles, and neocortex have all been reported in IGE (Betting et al., 2006), but most alterations are non-specific, and visual inspection of clinical MRI in IGE is typically normal (Woermann et al., 1998). Genomewide association studies (GWAS) have identified genetic variants associated with complex epilepsies by 'lumping' different epilepsy types together (International League Against Epilepsy Consortium on Complex Epilepsies, 2014), but MRI studies are typically of smaller scale, and have not widely explored whether distinct epilepsy syndromes share common structural abnormalities.

There are many sources of inconsistency in previously reported MRI findings. First, epileptic seizures and syndromes are diverse; classifications are often revised and contested (Berg *et al.*, 2010; Scheffer *et al.*, 2017). Second, most cross-sectional brain imaging studies are based on small samples (typically <50 cases), limiting the power to detect subtle group differences (Button *et al.*, 2013). Third, variability in scanning protocols, image processing, and statistical analysis may affect the sensitivity of brain measures across studies.

The Enhancing Neuro Imaging Genetics through Meta-Analysis (ENIGMA) Consortium was formed to address these issues (Bearden and Thompson, 2017). ENIGMA is a global initiative, combining large samples with coordinated image processing, and integrating genomic and MRI data across hundreds of research centres worldwide. Prior ENIGMA studies have identified genetic variants associated with variations in brain structure (Stein *et al.*, 2012; Hibar *et al.*, 2015, 2017*a*; Adams *et al.*, 2016), and have reliably characterized patterns of brain abnormalities in schizophrenia (van Erp *et al.*, 2016), major depression (Schmaal *et al.*, 2016), obsessive compulsive disorder (Boedhoe *et al.*, 2017), attention deficit hyperactivity disorder (Hoogman *et al.*, 2017), and many other brain illnesses (Thompson *et al.*, 2017). Large-scale, collaborative

initiatives such as ENIGMA may improve our understanding of epilepsy, helping clinicians make more informed decisions and provide personalized treatment strategies (Ben-Menachem, 2016). Thus, we formed the Epilepsy Working Group of ENIGMA ('ENIGMA-Epilepsy') to apply coordinated, well-powered studies of imaging and genetic data in epilepsy.

Here, in the largest analysis of structural brain abnormalities in epilepsy to date, we ranked effect sizes for 16 subcortical and 68 cortical brain regions in 2149 individuals with epilepsy and 1727 healthy controls, using harmonized image processing, quality control, and meta-analysis. First, we grouped all epilepsies together, to determine whether biologically distinct syndromes show robust, common structural deficits. Second, we assessed a wellcharacterized form of epilepsy: MTLE with hippocampal sclerosis, analysing patients with left- and right-sided hippocampal sclerosis as independent groups. Third, we examined another major set of epilepsy syndromes: IGE. Finally, we studied all remaining epilepsies as a combined subgroup, to understand the relative contributions of IGE, MTLE-L, MTLE-R, and all other syndromes on shared patterns of structural compromise. We tested how age at scan, age of onset, and epilepsy duration affected brain structural measures. Based on existing neuroimaging (Gotman et al., 2005; Bernhardt et al., 2009a; Liu et al., 2016), neurophysiological (Gotman et al., 2005), neuropathological (Thom et al., 2009), and genetic data (International League Against Epilepsy Consortium on Complex Epilepsies, 2014), we predicted that (i) biologically distinct epilepsy syndromes would exhibit shared patterns of structural abnormalities; (ii) MTLEs with left or right hippocampal sclerosis would show distinct patterns of hippocampal and extrahippocampal structural deficits; and (iii) IGEs would also display subcortical volume and cortical thickness differences, compared to healthy controls.

Materials and methods

Each centre received approval from their local institutional review board or ethics committee. Written informed consent was provided according to local requirements (Supplementary Table 1).

Experimental design

Participants

Twenty-four cross-sectional samples from 14 countries were included in the study, totalling 2149 people with epilepsy and 1727 research centre-matched healthy control subjects (Fig. 1 and Table 1). The locations, dates, and periods of participant recruitment are provided in Supplementary Table 1. An epilepsy specialist assessed seizure and syndrome classifications at each centre, using International League Against Epilepsy terminology (Berg *et al.*, 2010). Participants were aged 18–55.

To test for shared and syndrome-specific structural alterations, analyses included one group combining all epilepsies ('all-epilepsies'; n = 2149), and four stratified subgroups: (i) left MTLE with left hippocampal sclerosis (MTLE-L: n = 415; (ii) right MTLE with right hippocampal sclerosis (MTLE-R; n = 339); (iii) IGE (n = 367); and (iv) all other epilepsies (n = 1028). Supplementary Table 2 lists all syndromic diagnoses included in the aggregate 'all-epilepsies' group. For the MTLE subgroups, we included anyone with the typical electroclinical constellation (Berg et al., 2010), and a neuroradiologically-confirmed diagnosis of unilateral hippocampal sclerosis on clinical MRI. Participants were included in the IGE subgroup if they presented with tonic-clonic, absence or myoclonic seizures with generalized spike-wave discharges on EEG. Participants were included in the 'all-other-epilepsies' subgroup if they were diagnosed with non-lesional MTLE (43.3%), occipital (1.67%), frontal (8.78%), or parietal lobe epilepsy (0.84%), focal epilepsies not otherwise specified (37.03%), or another unclassified syndrome (8.37%; Supplementary Table 2). We excluded participants with a progressive disease (e.g. Rasmussen's encephalitis), malformations of cortical development, tumours or previous neurosurgery.

MRI data collection and processing

Structural T₁-weighted MRI brain scans were collected at the 24 participating centres. Scanning details are provided in

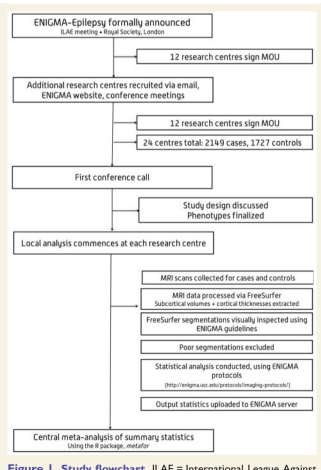


Figure I Study flowchart. ILAE = International League Against Epilepsy; MOU = memorandum of understanding.

Supplementary Table 3. T₁-weighted images from cases and controls were analysed at each site using FreeSurfer 5.3.0, for automated analysis of brain structure (Fischl, 2012). Volumetric measures were extracted for 12 subcortical grey matter regions (six left and six right, including the amygdala, caudate, nucleus accumbens, pallidum, putamen, and thalamus), the left and right hippocampi, and the left and right lateral ventricles. Cortical thickness measures were extracted for 34 left-hemispheric grey matter regions, and 34 right-hemispheric grey matter regions (68 total; Supplementary Table 4). Visual inspections of subcortical and cortical segmentations were conducted following standardized ENIGMA protocols (http://enigma.usc.edu), used in prior genetic studies of brain structure (Stein et al., 2012; Hibar et al., 2015, 2017a; Adams et al., 2016), and large-scale case-control studies of neuropsychiatric illnesses (Schmaal et al., 2015, 2016; Hibar et al., 2016; van Erp et al., 2016; Boedhoe et al., 2017). Analysts were blind to participants' diagnoses. Each analyst was instructed to execute a series of standardized bash scripts, identifying participants with volumetric or thickness measures greater or less than 1.5 times the interquartile range as outliers. Outlier data were then visually inspected, by overlaying the participant's cortical segmentations on their whole-brain anatomical images. If the blinded local analyst judged any structure as inaccurately segmented, that structure was omitted from the analysis. The Supplementary material provides further information.

Statistical analysis

Participant demographics

All research centres tested for differences in age between individuals with epilepsy and controls using an unpaired, two-tailed *t*-test in the *R* statistics package (https://www.r-project. org). Each centre also tested for sex differences between individuals with epilepsy and controls using a chi-squared test in SPSS Statistics package (IBM Corp., Version 21.0).

Meta-analytical group comparisons

Each research centre tested for case-versus-control differences using multiple linear regressions (via the *lm* function implemented in R), where a binary indicator of diagnosis (0 = healthy control, 1 = person with epilepsy) was the predictor of interest, and the volume or thickness of a specified brain region was the outcome measure. We calculated effect size estimates across all brain regions using Cohen's d, adjusting for age, sex and intracranial volume (ICV). ICV is a reliable, indirect measure of head size (Hansen et al., 2015), used as a covariate in other large-scale ENIGMA collaborations (Schmaal et al., 2015, 2016; Hibar et al., 2016; van Erp et al., 2016; Boedhoe et al., 2017). Cohen's d effect sizes and regression beta coefficients were pooled across centres using a random-effects, restricted maximum likelihood method of meta-analysis via the R package, metafor (Viechtbauer, 2010). The Supplementary material provides additional details.

Meta-analytical regression with clinical variables

Each centre conducted a series of linear regressions, testing the association between subcortical volume or cortical thickness, and: (i) age at onset of epilepsy; and (ii) duration of epilepsy.

Table I ENIGMA - Epilepsy Working Group demographics, including age (in years), mean age at onset of epilepsy (in years), mean duration of illness (in years), sex, and case-control breakdown for participating sites

Total n	134	161	127	96	96	42	229	130	66	45	138	167	197	4	307	174	277	174	80	71	689	911	71	3876
'Other' cases	26	38 0	09	0 (4)	36	∞	29	47	0	25	0	29	22	9	195	45	104	98	21	91	09	37	7	1028
IGE	12	> <u>@</u>	∞	44	2	91	0	0	0	5	39	0 (3)	32	0 (4)	36	0	36	25	0	0	40	40	=	367
MTLE-R cases	8 22	3 2	0 (4)	0	0	0	27	5	25	0	0	36	0 (4)	0 (2)	6	38	=	13	œ	0	84	0 (2)	15	339
MTLE-L cases	01	<u> </u>	=	0	9	0	61	œ	27	(I) 0	0	17	2	0 (3)	0	45	œ	22	4	0	107	0 (3)	25	415
Total cases	56	79	83	48	47	24	113	09	52	31	39	115	96	15	240	128	159	146	43	36	291	82	28	2149
Total controls	78	112	4	48	49	<u>&</u>	911	2	47	4	66	52	101	79	29	46	8 –	28	37	35	398	34	13	1727
Female	28	94	49	34	28	<u>&</u>	71	37	35	12	21	29	20	7	135	71	93	70	22	24	183	47	20	1228
Female	14 6	49	24	34	30	6	29	30	24	œ	09	29	54	91	33	21	62	12	91	25	249	70	4	949
Duration of illness (Mean ± years)	- 00 00	17.9 ± 12.93	19.02 ± 12.77	$\textbf{14.81} \pm \textbf{9.91}$	$\textbf{16.84} \pm \textbf{13.18}$	14.45 ± 11.14		18.93 ± 10.88	22.68 ± 14.28	$\textbf{14.27} \pm \textbf{8.06}$	$\textbf{14.13} \pm \textbf{12.81}$	$\textbf{17.64} \pm \textbf{10.51}$	20.67 ± 11.23	$\textbf{8.33} \pm \textbf{9.99}$	$\textbf{9.35} \pm \textbf{11.23}$	16.05 ± 11.32	$\textbf{16.43} \pm \textbf{12.7}$	10.18 ± 12.65	18.8 ± 15.36	15.03 ± 12.53	27.96 ± 12.54	14.34 ± 10.94	11.76 \pm 8.78	$\textbf{17.42} \pm \textbf{11.99}$
Age of onset (Mean ± SD)		17.9 ± 11.49	$\textbf{14.46} \pm \textbf{10.13}$	$\textbf{13.56} \pm \textbf{5.18}$	$\textbf{17.04} \pm \textbf{11.09}$	$\textbf{17.32} \pm \textbf{10.8}$		$\textbf{17.03} \pm \textbf{13.7}$	14.51 ± 11.8	$\textbf{12.69} \pm \textbf{8.02}$	28.12 ± 17.86	$\textbf{18.07} \pm \textbf{11.72}$	$\textbf{13.22} \pm \textbf{8.2}$	$\textbf{23.13} \pm \textbf{7.55}$	24 ± 13.22	$\textbf{16.48} \pm \textbf{9.72}$	$\textbf{16.96} \pm \textbf{11.27}$	28.23 ± 17.98	19.32 ± 14.77	16.26 ± 11.33	12.07 ± 9.52	$\textbf{12.58} \pm \textbf{8.13}$	17.04 ± 12.2	$\textbf{17.63} \pm \textbf{11.47}$
Age cases (Mean ± SD)	30.48 ± 10.13	33.28 ± 10.59	33.79 ± 9.9	28.42 ± 8.06	33.58 ± 11.07	31.13 ± 10.74	30.42 ± 10.13	36.2 ± 9.97	37.46 ± 10.69	28 ± 7.77	26.23 ± 7.49	36.77 ± 9.52	33.2 ± 8.9	31.47 ± 11.33	33.35 ± 11.21	$\textbf{32.53} \pm \textbf{9.92}$	33.23 ± 9.66	38.08 ± 15.91	37.67 ± 11.79	$\textbf{31.47} \pm \textbf{11.81}$	39.98 ± 10.25	28.36 ± 10.26	28.79 ± 9.06	$\textbf{34.36} \pm \textbf{10.65}$
Age controls (Mean ± SD)	32.5 ± 9.39	34.73 ± 10.61	$\textbf{26.64} \pm \textbf{4.34}$	$\textbf{28.04} \pm \textbf{8.16}$	34.82 ± 11.38	35.33 ± 12.27	30.48 ± 9.39	34.75 ± 9.36	31.7 ± 9.24	35.29 ± 8.48	42.26 ± 14.97	33.13 ± 5.99	$\textbf{31.68} \pm \textbf{8.4}$	28.73 ± 8.29	25.16 ± 1.55	30.74 ± 7.38	30.1 ± 10.36	39.35 ± 20.26	36.89 ± 15.1	33.2 ± 12.29	34.39 ± 10.45	28.47 ± 5.25	31.54 ± 6.99	$\textbf{33.31} \pm \textbf{9.91}$
Site name	Bern	BRI	Brussels	CUBRIC	EKUT_A	EKUT_B	EPICZ	EPIGEN_3.0	EPIGEN_1.5	Florence	Greifswald	IDIBAPS-HCP	KCL_CNS	KCL_CRF	Kuopio	ZΣ	NYU	RMH	UCSD	UNAM	UNICAMP	UNIMORE	XMU	Combined

Also provided is the total number of MTLE cases with left hippocampal sclerosis, MTLE cases with right hippocampal sclerosis, IGE and all-other-epilepsies ('other') cases per site. Research centres with fewer than five participants for a given phenotype are marked as '0' for that phenotype, with the original sample size noted in parentheses. $SD = standard\ deviation$.

All centres tested for interactions between diagnosis of epilepsy (including syndrome groups) and age at time of scan. Beta values representing the unstandardized slopes of each regression were extracted for each analysis. Sex and ICV were included as covariates in all secondary analyses.

Correction for multiple comparisons

We conducted four independent regressions (one case versus control regression, and three regressions with clinical variables) across 84 regions of interest, adjusting the statistical significance threshold to $P_{\rm thresh} < 1.49 \times 10^{-4}$ to correct for 336 comparisons. To account for correlations between tests, we also applied a less conservative adjustment for false discovery rate (FDR), using the Benjamini and Hochberg method (Benjamini and Hochberg, 1995). For clarity, we report only P-values significant after stringent Bonferroni correction; FDR-adjusted P-values are summarized in the Supplementary material.

Power analyses

Across all regions of interest, we calculated the sample sizes necessary to achieve 80% power to detect case-control differences, given the observed effect sizes at each region of interest, based on two-tailed *t*-tests, using G^*Power Version 3.1. For each region of interest, we also estimated N_{80} : the total number of samples required, per group, to achieve 80% power to detect group differences using a *t*-test at the threshold of P < 0.05 (two-tailed).

Results

Participant demographics

The sample size-weighted mean age across all epilepsy samples was 34.4 (range: 26.2–40) years, and the weighted mean age of healthy controls was 33.3 (range: 25.2–42.3) years. The weighted mean age at onset of epilepsy and duration of epilepsy were 17.6 (range: 12.1–28.2) years and 17.4 (range: 8.3–28) years, respectively. Females comprised 57% of the total epilepsy sample (range: 34–75% by individual sample), and 53% of the controls (range: 31–71% by individual sample). Case-control differences in age were observed at 8 of 24 research centres, and case-control differences in sex were observed at 2 of 24 research centres (Supplementary Table 5); hence, age and sex were included as covariates in all group comparisons.

Volumetric findings

Compared to controls, the aggregate all-epilepsies group exhibited lower volumes in the left $(d = -0.36; P = 1.31 \times 10^{-6})$ and right thalamus $(d = -0.37; P = 7.67 \times 10^{-14})$, left $(d = -0.35; P = 3.04 \times 10^{-7})$ and right hippocampus $(d = -0.34; P = 6.63 \times 10^{-10})$, and the right pallidum $(d = -0.32; P = 8.32 \times 10^{-9})$. Conversely, the left $(d = 0.29; P = 2.14 \times 10^{-12})$ and right $(d = 0.27; P = 3.73 \times 10^{-15})$ lateral ventricles were enlarged across all epilepsies when compared to controls (Table 2 and Fig. 2A). A supplementary analysis of all-epilepsies,

excluding individuals with hippocampal sclerosis or other lesions, revealed similar patterns of volume loss in the right thalamus and pallidum, and bilaterally enlarged ventricles; however, volume differences were not observed in the hippocampus (Supplementary Table 6).

The MTLE-L subgroup showed lower volumes in the left hippocampus $(d=-1.73;\ P=1.35\times 10^{-19})$, left $(d=P=2.19\times 10^{-11})$ and right thalamus $(d=-0.46;\ P=8.12\times 10^{-5})$, left putamen $(d=-0.39;\ P=1.07\times 10^{-6})$, and right pallidum $(d=-0.45;\ P=5.48\times 10^{-7})$. As in the overall group comparison, we observed larger left $(d=0.47;\ P=1.96\times 10^{-7})$ and right lateral ventricles $(d=0.36;\ P=8.95\times 10^{-5})$ in MTLE-L patients relative to controls (Table 2 and Fig. 2B).

The MTLE-R subgroup showed lower volumes across a number of regions in the right hemisphere only, including the hippocampus (d = -1.91; $P = 6.36 \times 10^{-37}$), thalamus (d = -0.73; $P = 1.6 \times 10^{-12}$), and pallidum (d = -0.45; $P = 3.96 \times 10^{-7}$), together with increased volumes of the left (d = 0.39; $P = 1.52 \times 10^{-6}$) and right lateral ventricles (d = 0.44; $P = 6.57 \times 10^{-12}$) compared to controls (Table 2 and Fig. 2C).

The IGE subgroup showed lower volumes in the right thalamus (d = -0.4; $P = 3.6 \times 10^{-6}$) compared to controls (Table 2 and Fig. 2D).

The all-other-epilepsies subgroup showed lower volumes in the right thalamus (d = -0.31; $P = 7.9 \times 10^{-11}$) and the right pallidum (d = -0.24; $P = 8.1 \times 10^{-5}$) compared to controls. The all-other-epilepsies subgroup also showed significant enlargements of the left (d = 0.33; $P = 5.1 \times 10^{-7}$) and right amygdala (d = 0.22; $P = 1.46 \times 10^{-4}$), and the left (d = 0.2; $P = 1.2 \times 10^{-5}$) and right lateral ventricles (d = 0.21; $P = 4.62 \times 10^{-6}$) compared to controls (Table 2 and Fig. 2E).

All volume differences can be visualized using the interactive ENIGMA-Viewer tool (Zhang *et al.*, 2017), at http://enigma-viewer.org/ENIGMA_epilepsy_subcortical.html (Supplementary material). Volume differences significant after FDR adjustment can also be visualized at http://enigma-viewer.org/ENIGMA_epilepsy_subcortical_fdr.html (Supplementary Tables 26–30).

Cortical thickness findings

The all-epilepsies group showed reduced thickness of cortical grey matter across seven regions bilaterally, including the left (d = -0.38; $P = 1.82 \times 10^{-18}$) and right precentral gyri (d = -0.4; $P = 8.85 \times 10^{-20}$), left (d = -0.32; $P = 2.11 \times 10^{-15}$) and right caudal middle frontal gyri (d = -0.31; $P = 2.09 \times 10^{-9}$), left (d = -0.31; $P = 2.05 \times 10^{-6}$) and right paracentral gyri (d = -0.32; $P = 2.19 \times 10^{-9}$), left (d = -0.19; $P = 1.29 \times 10^{-4}$) and right pars triangularis (d = -0.2; $P = 4.25 \times 10^{-8}$), left (d = -0.28; $P = 1.51 \times 10^{-7}$) and right superior frontal gyri (d = -0.27; $P = 4.49 \times 10^{-6}$), left (d = -0.19; $P = 1.05 \times 10^{-5}$) and right transverse temporal gyri (d = -0.18; $P = 2.81 \times 10^{-5}$), and left (d = -0.23; $P = 9.87 \times 10^{-5}$) and right

Table 2 Effect size differences between epilepsy cases and healthy controls (Cohen's d) for the mean volume of subcortical structures, controlling for age, sex and intracranial volume

CI = confidence interval; LH = left hemisphere; RH = right hemisphere; SE = standard error; l^2 = heterogeneity index; N_{80} = number of subjects required in each group to yield 80% power to detect significant group differences (l^2 < 0.05, two-tailed). Uncorrected P-values are reported. Subcortical structures that failed to survive Bonferroni correction (l^2 < 1.49 × 10⁻⁴) are not reported (see 'Materials and methods' section for statistical threshold determination). See Supplementary material for a full list of volume differences with adjustment for false discovery rate (FDR).

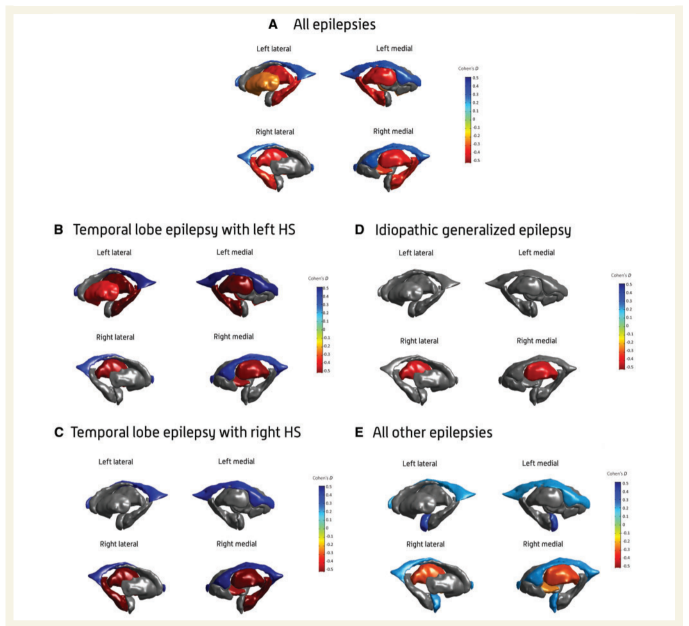


Figure 2 Subcortical volume findings. Cohen's d effect size estimates for case-control differences in subcortical volume, across the (A) allepilepsies, (B) mesial temporal lobe epilepsies with left hippocampal sclerosis (HS; MTLE-L), (C) mesial temporal lobe epilepsies with right hippocampal sclerosis (MTLE-R), (D) idiopathic generalized epilepsies (IGE), and (E) all-other-epilepsies groups. Cohen's d effect sizes were extracted using multiple linear regressions, and pooled across research centres using random-effects meta-analysis. Subcortical structures with Pvalues $< 1.49 \times 10^{-4}$ are shown in heatmap colours; strength of heat map is determined by the size of the Cohen's d (d < 0 = blue, d > 0 = yellow/red). Image generated using MATLAB, with annotations added using Adobe Photoshop. An interactive version of this figure is available online, via 'ENIGMA-Viewer': http://enigma-viewer.org/ENIGMA_epilepsy_subcortical.html. See Supplementary material for guidelines on how to use the interactive visualization.

supramarginal gyri (d = -0.22; $P = 5.24 \times 10^{-5}$). The allepilepsies group also showed unilaterally thinner right cuneus (d = -0.2; $P = 9.68 \times 10^{-8}$), right pars opercularis $(d = -0.18; P = 6.48 \times 10^{-7})$, right precuneus (d = -0.28; $P = 2.7 \times 10^{-5}$), and left entorhinal gyrus (d = -0.26; $P = 2.04 \times 10^{-5}$), compared to healthy controls (Table 3 and Fig. 3A). Supplementary analysis in a non-lesional epilepsy subgroup revealed a similar pattern of cortical thickness differences compared to controls, suggesting that the

changes observed in our main analysis were not driven by the inclusion of patients with hippocampal sclerosis or other common lesions (Supplementary Table 7).

The MTLE-L and MTLE-R subgroups showed distinct patterns of cortical thickness reductions when compared to healthy controls (Table 3, Fig. 3B and C). In MTLE-R, lower cortical thickness was reported across four motor regions, including the left $(d = -0.51; P = 7.67 \times 10^{-7})$ and right paracentral gyri (d = -0.42; $P = 6.24 \times 10^{-11}$),

Table 3 Effect size differences between epilepsy cases and healthy controls (Cohen's d) for the mean thickness of cortical structures, controlling for age, sex and intracranial volume

	ad koulair	Conen's d	1		j)	-		of	of cases
									controls	
Caudal middle frontal gyrus (LH)	MTLE-L	-0.403	0.07	-5.789	-0.538 to -0.2663	7.07×10^{-9}	13.807	86	1344	412
	All epilepsies	-0.319	0.04	-7.935	-0.397 to -0.24	2.11×10^{-15}	17.112	156	1650	2061
	All other epilepsies	-0.291	0.045	-6.425	-0.38 to -0.202	1.32×10^{-10}	0	197	1447	0001
Caudal middle frontal gyrus (RH)	MTLE-L	-0.441	0.087	-5.089	-0.611 to -0.271	3.61×10^{-7}	39.444	82	1348	412
	All epilepsies	-0.307	0.051	-5.991	-0.407 to -0.206	2.09×10^{-9}	46.443	168	1653	2059
	All other epilepsies	-0.212	0.045	-4.699	-0.301 to -0.124	2.62×10^{-6}	0	350	1451	866
Cuneus (RH)	All other epilepsies	-0.234	0.045	-5.186	-0.323 to -0.146	2.15×10^{-7}	0	288	1449	966
	All epilepsies	-0.204	0.038	-5.333	-0.279 to -0.129	9.68 ×10 ⁻⁸	11.423	379	1651	2057
Entorhinal gyrus (LH)	MTLE-L	-0.445	0.072	-6.158	-0.5865 to -0.303	7.35×10^{-10}	0	8	1102	303
	All epilepsies	-0.264	0.062	-4.261	-0.385 to -0.142	2.04×10^{-5}	56.648	227	1402	1724
Fusiform gyrus (LH)	MTLE-L	-0.359	0.069	-5.183	-0.494 to -0.223	2.19×10^{-7}	13.465	123	1339	412
Lateral occipital gyrus (RH)	All other epilepsies	-0.211	0.045	-4.659	-0.299 to -0.122	3.18×10^{-6}	2.50×10^{-3}	354	1450	266
Lingual gyrus (RH)	All other epilepsies	-0.180	0.045	-3.972	-0.268 to -0.091	7.12×10^{-5}	1.25×10^{-2}	491	1450	966
Paracentral gyrus (LH)	MTLE-R	-0.505	0.102	-4.944	-0.705 to -0.305	7.67×10^{-7}	52.283	63	1292	338
	MTLE-L	-0.426	0.099	-4.313	-0.62 to -0.232	1.61×10^{-5}	53.165	88	1344	412
	All epilepsies	-0.311	0.065	-4.748	-0.439 to -0.182	2.05×10^{-6}	67.476	164	1650	2061
	All other epilepsies	-0.257	0.045	-5.680	-0.346 to -0.168	1.34×10^{-8}	0	239	1447	0001
Paracentral gyrus (RH)	MTLE-R	-0.421	0.064	-6.538	-0.548 to -0.295	6.24×10^{-11}	0.407	06	1296	338
	MTLE-L	-0.378	0.075	-5.021	-0.526 to -0.231	5.14×10^{-7}	23.536	Ξ	1348	412
	All other epilepsies	-0.351	0.045	-7.733	-0.44 to -0.262	1.05×10^{-14}	3.43×10^{-3}	129	1451	866
	All epilepsies	-0.315	0.053	-5.983	-0.418 to -0.212	2.19×10^{-9}	49.261	091	1654	2059
Parahippocampal gyrus (LH)	MTLE-L	-0.3	0.073		-0.444 to -0.1572	3.95×10^{-5}	19.366	176	1335	410
Pars opercularis (RH)	MTLE-R	-0.271	0.071	-3.8	-0.411 to -0.131	1.45×10^{-4}	12.105	215	1295	338
	All epilepsies	-0.177	0.036	-4.976	-0.247 to -0.107	6.48×10^{-7}	2.624	503	1652	2059
Pars triangularis (LH)	All epilepsies	-0.192	0.05	-3.828	-0.2897 to -0.094	1.29×10^{-4}	44.414	427	1650	2060
Pars triangularis (RH)	MTLE-L	-0.285	90.0	-4.738	-0.403 to -0.167	2.16×10^{-6}	0	195	1346	412
	All epilepsies	-0.199	0.036	-5.48	-0.27 to -0.128	4.25×10^{-8}	4.66	398	1652	2058
	All other epilepsies	-0.210	0.045	-4.650	-0.299 to -0.122	3.32×10^{-6}	2.58×10^{-3}	357	1449	866
Precentral gyrus (LH)	MTLE-L	-0.466	0.081	-5.755	-0.625 to -0.307	8.64×10^{-9}	31.602	74	1339	412
	MTLE-R	-0.415	60.0	-4.596	-0.592 to -0.238	4.31×10^{-6}	40.044	93	1287	338
	All epilepsies	-0.384	0.044	-8.768	-0.469 to -0.298	1.82×10^{-18}	27.649	108	1645	2058
	All other epilepsies	-0.375	0.046	-8.237	-0.464 to -0.286	1.76×10^{-16}	5.59×10^{-3}	13	1442	266
	IGE	-0.342	0.071	-4.78	-0.482 to -0.201	1.75×10^{-6}	0.003	136	1043	297
Precentral gyrus (RH)	MTLE-R	-0.52	980.0	-6.073	-0.687 to -0.352	1.25×10^{-9}	33.288	09	1293	337
	MTLE-L	-0.492	0.078	-6.335	-0.6436 to -0.339	2.37×10^{-10}	26.33	99	1345	412
	All epilepsies	-0.399	0.044	-9.102	-0.485 to -0.313	8.85×10^{-20}	27.929	00	1649	2054
	JGE	-0.39	0.072	-5.442	-0.531 to -0.25	5.27×10^{-8}	0.005	105	1044	295
	All other epilepsies	-0.348	0.045	-7.672	-0.437 to -0.259	1.70×10^{-14}	0	131	1448	966
Precuneus (LH)	MTLE-L	-0.536	0.135	-3.965	-0.801 to -0.271	7.35×10^{-5}	75.18	26	1343	412
	All other epilepsies	-0.178	0.047	-3.819	-0.27 to -0.087	1.34×10^{-4}	4.474	497	1446	866

Table 3 Continued

Structure	Phenotype	Cohen's d	SE	Z score	95% CI	P-value	12	N 80	Number of controls	Number of cases
Precuneus (RH)	MTLE-L	-0.473	0.104	-4.558	-0.676 to -0.27	5.16 × 10 ⁻⁶	57.498	72	1348	412
	All epilepsies	-0.275	990.0	-4.197	-0.404 to -0.147	2.70×10^{-5}	809'29	209	1654	2055
	All other epilepsies	-0.238	0.053	-4.471	-0.343 to -0.134	7.78×10^{-6}	22.378	279	1451	994
Superior frontal gyrus (LH)	MTLE-L	-0.411	90:0	-6.804	-0.529 to -0.292	1.02×10^{-11}	0	94	1343	412
	All epilepsies	-0.283	0.054	-5.251	-0.389 to -0.177	1.51×10^{-7}	51.773	197	1649	2059
	All other epilepsies	-0.243	0.059	-4.138	-0.358 to -0.128	3.51×10^{-5}	34.545	267	1446	666
Superior frontal gyrus (RH)	MTLE-L	-0.365	90:0	-6.051	-0.483 to -0.246	1.44×10^{-9}	0	611	1345	412
	All epilepsies	-0.269	0.059	-4.588	-0.385 to -0.154	4.49×10^{-6}	59.483	218	1650	2058
	All other epilepsies	-0.235	0.052	-4.489	-0.337 to -0.132	7.15×10^{-6}	20.049	286	1448	266
Superior parietal gyrus (LH)	All other epilepsies	-0.224	0.045	-4.954	-0.313 to -0.136	7.27×10^{-7}	0.001	314	1444	966
Superior parietal gyrus (RH)	All other epilepsies	-0.220	0.045	-4.864	-0.309 to -0.131	1.15×10^{-6}	0.002	326	1450	266
Supramarginal gyrus (LH)	All epilepsies	-0.232	90:0	-3.894	-0.348 to -0.115	9.87×10^{-5}	59.391	293	9091	1965
Supramarginal gyrus (RH)	All epilepsies	-0.223	0.055	-4.045	-0.331 to -0.115	5.24×10^{-5}	52.895	317	1597	1971
	All other epilepsies	-0.206	0.047	-4.418	-0.297 to -0.115	9.95×10^{-6}	0	371	1395	196
Temporal pole (LH)	MTLE-L	-0.315	890.0	-4.649	-0.447 to -0.182	3.33×10^{-6}	10.901	091	1341	410
Transverse temporal gyrus (LH)	MTLE-R	-0.312	0.073	-4.249	-0.456 to -0.168	2.15×10^{-5}	15.614	163	1289	338
	All epilepsies	-0.192	0.044	-4.406	-0.278 to -0.107	1.05×10^{-5}	28.178	427	1647	2061
Transverse temporal gyrus (RH)	All epilepsies	-0.182	0.044	-4.188	-0.267 to -0.097	2.81×10^{-5}	27.918	475	1654	2059
	All other epilepsies	-0.18	0.045	-3.982	-0.269 to -0.091	6.84×10^{-5}	0.012	486	1451	866

CI = confidence interval; LH = left hemisphere; RH = right hemisphere; SE = standard error; l^2 = heterogeneity index; N_{80} = number of subjects required in each group to yield 80% power to detect significant group differences (l^2 = heterogeneity index; l^2 = heterogeneity index; l^2 = heterogeneity index; l^2 = number of subjects required index; l^2 = heterogeneity index;

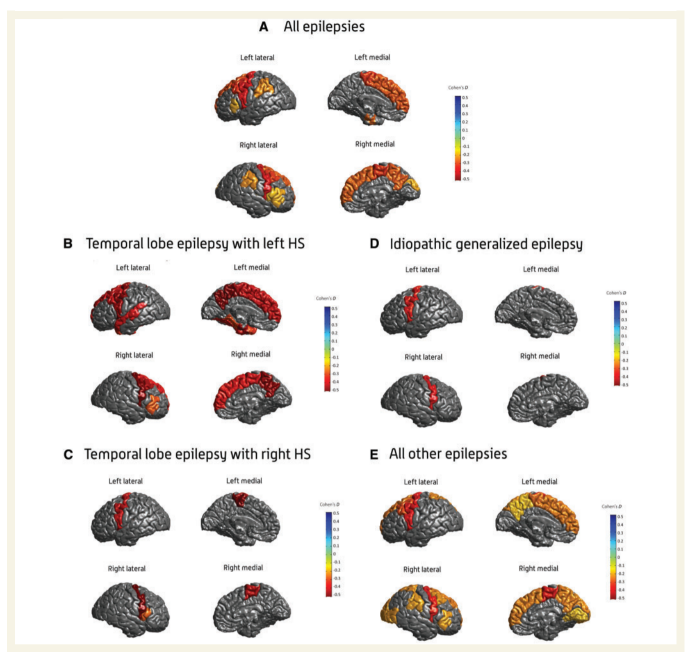


Figure 3 Cortical thickness findings. Cohen's d effect size estimates for case-control differences in cortical thickness, across the (**A**) all-epilepsies, (**B**) mesial temporal lobe epilepsies with left hippocampal sclerosis (MTLE-L), (**C**) mesial temporal lobe epilepsies with right hippocampal sclerosis (MTLE-R), (**D**) idiopathic generalized epilepsies (IGE), and (**E**) all-other-epilepsies groups. Cohen's d effect sizes were extracted using multiple linear regressions, and pooled across research centres using random-effects meta-analysis. Cortical structures with P-values $< 1.49 \times 10^{-4}$ are shown in heatmap colours; strength of heat map is determined by the size of the Cohen's d (d < 0 = blue, d > 0 = yellow/red). Image generated using MATLAB with annotations added using Adobe Photoshop. An interactive version of this figure is available online, via 'ENIGMA-Viewer': http://enigma-viewer.org/ENIGMA_epilepsy_cortical.html. See Supplementary material for guidelines on how to use the interactive visualization. HS = hippocampal sclerosis.

and the left (d = -0.42; $P = 4.31 \times 10^{-6}$) and right precentral gyri (d = -0.52; $P = 1.25 \times 10^{-9}$). The MTLE-R subgroup also showed thickness changes in the left transverse temporal gyrus (d = -0.31; $P = 2.15 \times 10^{-5}$), and right pars opercularis (d = -0.27; $P = 1.45 \times 10^{-4}$) (Table 3 and Fig. 3C). By contrast, in MTLE-L, lower thickness was observed across six regions of the motor cortex, including the left (d = -0.43; $P = 1.61 \times 10^{-5}$) and right

paracentral gyri $(d = -0.38; P = 5.14 \times 10^{-7})$, left $(d = -0.47; P = 8.64 \times 10^{-9})$ and right precentral gyri $(d = -0.49; P = 2.37 \times 10^{-10})$, and left $(d = -0.54; P = 7.35 \times 10^{-5})$ and right precuneus $(d = -0.47; P = 5.16 \times 10^{-6})$. The MTLE-L group also showed thickness changes across five regions of the frontal cortex, including the left $(d = -0.41; P = 1.02 \times 10^{-11})$ and right superior frontal gyri $(d = -0.37; P = 1.44 \times 10^{-9})$, left $(d = -0.4; P = 1.44 \times 10^{-9})$, left $(d = -0.4; P = 1.44 \times 10^{-9})$, left $(d = -0.4; P = 1.44 \times 10^{-9})$, left $(d = -0.4; P = 1.44 \times 10^{-9})$, left $(d = -0.4; P = 1.44 \times 10^{-9})$, left $(d = -0.4; P = 1.44 \times 10^{-9})$, left $(d = -0.4; P = 1.44 \times 10^{-9})$, left $(d = -0.4; P = 1.44 \times 10^{-9})$, left $(d = -0.4; P = 1.44 \times 10^{-9})$, left $(d = -0.4; P = 1.44 \times 10^{-9})$, left $(d = -0.4; P = 1.44 \times 10^{-9})$

 $P = 7.07 \times 10^{-9}$) and right caudal middle frontal gyri $(d = -0.44; P = 3.61 \times 10^{-7})$, and the right *pars triangularis* $(d = -0.29; P = 2.16 \times 10^{-6})$. In MTLE-L, thickness alterations were also observed in four regions of the temporal cortex, including the left temporopolar cortex $(d = -0.32; P = 3.33 \times 10^{-6})$, left parahippocampal gyrus $(d = -0.3; P = 3.95 \times 10^{-5})$, left entorhinal gyrus $(d = -0.45; P = 7.35 \times 10^{-10})$, and left fusiform gyrus $(d = -0.36; P = 2.19 \times 10^{-7})$ (Table 3 and Fig. 3B).

The IGE subgroup showed reduced thickness in the left $(d = -0.34; P = 1.75 \times 10^{-6})$ and right precentral gyri $(d = -0.39; P = 5.27 \times 10^{-8})$, when compared to healthy controls (Table 3 and Fig. 3D).

The all-other-epilepsies subgroup showed lower thickness across six cortical regions bilaterally, including the left $(d = -0.38; P = 1.76 \times 10^{-16})$ and right precentral gyri $(d = -0.35; P = 1.7 \times 10^{-14}), \text{ left } (d = -0.26; P = 1.34 \times 10^{-14})$ 10^{-8}) and right paracentral gyri (d = -0.35; $P = 1.1 \times$ 10^{-14}), left $(d = -0.29; P = 1.32 \times 10^{-10})$ and right caudal middle frontal gyri (d = -0.21; $P = 2.62 \times 10^{-6}$), left (d = -0.22; $P = 7.27 \times 10^{-7}$) and right superior parietal gyri (d = -0.22; $P = 1.15 \times 10^{-6}$), left (d = -0.24; P = 3.51 $\times 10^{-5}$) and right superior frontal gyri (d = -0.23; $P = 7.15 \times 10^{-6}$), and the left $(d = -0.18; P = 1.34 \times 10^{-6})$ 10^{-4}) and right precuneus (d = -0.24; $P = 7.78 \times 10^{-6}$) compared to controls. The all-other-epilepsies group also showed unilaterally reduced thickness in six right hemispheric regions, including the cuneus (d = -0.23; P = 2.15 $\times 10^{-7}$), lateral occipital gyrus (d = -0.21; $P = 3.18 \times 10^{-7}$) 10^{-6}), pars triangularis (d = -0.21; $P = 3.32 \times 10^{-6}$), supramarginal gyrus (d = -0.21; $P = 9.95 \times 10^{-6}$), transverse temporal gyrus (d = -0.18; $P = 6.84 \times 10^{-5}$), and lingual gyrus (d = -0.18; $P = 7.12 \times 10^{-5}$), compared to controls (Table 3 and Fig. 3E).

An interactive 3D visualization of these results is available via the ENIGMA-Viewer tool (Zhang *et al.*, 2017), at http://enigma-viewer.org/ENIGMA_epilepsy_cortical.html (Supplementary material). Cortical thickness differences significant after FDR adjustment can also be visualized at http://enigma-viewer.org/ENIGMA_epilepsy_cortical_fdr.html (Supplementary Tables 31–35).

Duration of illness, age at onset, and age-by-diagnosis effects on brain abnormalities

A secondary analysis identified significant associations between duration of epilepsy and several affected brain regions in the all-epilepsies, MTLE-R, and all-other-epilepsies groups. In the all-epilepsies group, duration of epilepsy negatively associated with volume measures in the left hippocampus (b = -8.32; $P = 8.16 \times 10^{-13}$), left (b = -13.58; $P = 3.52 \times 10^{-15}$), and right thalamus (b = -12.25; $P = 1.58 \times 10^{-13}$), and right pallidum (b = -2.67; $P = 1.78 \times 10^{-7}$), in addition to bilateral thickness measures in the left (b = -0.003; $P = 2.99 \times 10^{-11}$) and right pars

triangularis (b = -0.002; $P = 4.24 \times 10^{-9}$), left (b = -0.003; $P = 1.61 \times 10^{-15}$) and right caudal middle frontal gyri $(b = -0.003; P = 1.65 \times 10^{-17}), \text{ left } (b = -0.003; P = 1.77)$ $\times 10^{-13}$) and right supramarginal gyri (b = -0.003; $P = 2.58 \times 10^{-19}$), left $(b = -0.003; P = 5.84 \times 10^{-12})$ and right precentral gyri (b = -0.003; $P = 2.54 \times 10^{-24}$), left $(b = -0.004; P = 1.94 \times 10^{-12})$ and right superior frontal gyri $(b = -0.003; P = 4.65 \times 10^{-11})$, left (b = -0.004; $P = 1.05 \times 10^{-10}$) and right transverse temporal gyri $(b = -0.003; P = 8.24 \times 10^{-10})$, and left (b = -0.002; $P = 5.22 \times 10^{-6}$) and right paracentral gyri (b = -0.002; $P = 5.63 \times 10^{-6}$). Duration of epilepsy also negatively associated with unilateral thickness measures in the right precuneus (b = -0.003; $P = 6.03 \times 10^{-21}$), right pars opercularis (b = -0.003; $P = 5.59 \times 10^{-13}$), and right cuneus $(b = -0.002; P = 1.1 \times 10^{-9}; Supplementary Table 8).$ In the MTLE-R subgroup, duration of epilepsy negatively associated with volume measures in the right hippocampus $(b = -22.42; P = 1.1 \times 10^{-7})$, and the right thalamus $(b = -18.11; P = 1.84 \times 10^{-5})$, and thickness measures in the left transverse temporal gyrus (b = -0.007; P = 8.39 \times 10⁻⁵; Supplementary Table 8). In the all-other-epilepsies subgroup, duration of epilepsy negatively associated with bilateral thickness measures in the left (b = -0.003; $P = 3.39 \times 10^{-7}$) and right caudal middle frontal gyri $(b = -0.003; P = 6.91 \times 10^{-8}), left (b = -0.003; P = 1.36 \times 10^{-8})$ 10^{-9}) and right superior frontal gyri (b = -0.003; P = 3.16 $\times 10^{-7}$), and the left (b = -0.003; P = 3.17 $\times 10^{-5}$) and right precuneus (b = -0.003; $P = 5.01 \times 10^{-9}$), in addition to unilateral thickness measures in the right precentral gyrus $(b = -0.004; P = 1.16 \times 10^{-12})$, right cuneus $(b = -0.003; P = 8.57 \times 10^{-8})$, right pars triangularis $(b = -0.003; P = 5.16 \times 10^{-7})$, and right supramarginal gyrus (b = -0.003; $P = 2.24 \times 10^{-7}$). Duration of epilepsy also showed a positive association with the size of the left lateral ventricle in the all-other-epilepsies group (b = 13.6; $P = 1.17 \times 10^{-5}).$

In the all-epilepsies group, age at onset of epilepsy negatively associated with thickness measures in the left $(b = -0.003; P = 2.66 \times 10^{-15})$ and right superior frontal gyri $(b = -0.003; P = 9.77 \times 10^{-10})$, left (b = -0.003; $P = 2.78 \times 10^{-9}$) and right pars triangularis (b = -0.003; $P = 6.51 \times 10^{-7}$), right pars opercularis (b = -0.003; P = -0.003) 5.4×10^{-14}), left transverse temporal gyrus (b = -0.003; $P = 1.03 \times 10^{-8}$), and right cuneus (b = -0.001; $P = 4.9 \times 10^{-6}$). In the all-other-epilepsies subgroup, age at onset negatively correlated with thickness measures in the left (b = -0.003; $P = 3.21 \times 10^{-8}$) and right superior frontal gyri $(b = -0.002; P = 1.18 \times 10^{-4}),$ left $(b = -0.002; P = 8.42 \times 10^{-6})$ and right precuneus $(b = -0.002; P = 7.23 \times 10^{-5})$, right pars triangularis $(b = -0.003; P = 2.53 \times 10^{-5})$, and right supramarginal gyrus $(b = -0.002; P = 2.38 \times 10^{-6})$. Age at onset also positively associated with the size of the right lateral ventricle in the all-other-epilepsies subgroup (b = 57.73; $P = 1.62 \times 10^{-7}).$

Age at onset negatively associated with other regional volumetric and thickness measures in the all-epilepsies, IGE, MTLE-L, MTLE-R, and all-other-epilepsies groups, but these associated areas showed no significant structural differences in the primary case-control analysis (Table 1 and Supplementary Table 8).

There were no interaction effects between age and syndromic diagnosis in the all-epilepsies, MTLE-L, MTLE-R, IGE, or all-other-epilepsies groups.

Power analyses for detection of case-control differences

In our sample of 2149 individuals with epilepsy and 1727 healthy controls, we had 80% power to detect Cohen's d effect sizes as small as d = 0.091 at the standard alpha level of P < 0.05 (two-tailed), and 80% power to detect Cohen's d effect sizes as small as d = 0.149 at the study's stringent Bonferroni-corrected threshold of $P < 1.49 \times 10^{-4}$.

 N_{80} , the number of cases and controls required to achieve 80% power to detect group differences using a two-tailed *t*-test at P < 0.05, ranged from $N_{80} = 6$, to detect group effects in the right hippocampus in our MTLE-R group, to $N_{80} = 503$, to detect group effects in the right pars opercularis in our 'all epilepsies' group (Tables 2 and 3).

Discussion

In the largest coordinated neuroimaging study of epilepsy to date, we identified a series of quantitative imaging signatures—some shared across common epilepsy syndromes, and others characteristic of selected, specific epilepsy syndromes. Our sample of 2149 individuals with epilepsy and 1727 controls provided 80% power to detect differences as small as d = 0.091 (P < 0.05, two-tailed), allowing us to identify subtle, consistent brain abnormalities that are typically undetectable on visual inspection, or overlooked using smaller case-control designs. This international collaboration addresses prior inconsistencies in the field of epilepsy neuroimaging, providing a robust, *in vivo* map of structural aberrations, upon which future studies of disease mechanisms may expand.

In the first of five cross-sectional MRI analyses, we investigated a diverse aggregation of epilepsy syndromes, putative causes, and durations of disease. This all-epilepsies group exhibited shared, diffuse brain structural differences across several regions including the thalamus, pallidum, precentral, paracentral, and superior frontal cortices. With the exception of hippocampal volume and entorhinal thickness differences (Supplementary material), these structural alterations were not driven by any specific syndrome or dataset (Supplementary Figs 3 and 7). Our findings suggest a common neuroanatomical signature of epilepsy across a wide spectrum of disease types, complementing recent evidence for shared genetic susceptibility to a wide spectrum

of epilepsies (International League Against Epilepsy Consortium on Complex Epilepsies, 2014). Some structural and genetic pathways may be shared across syndromes, despite the heterogeneity of epilepsy and seizure types. This shared MRI signature underpins the contemporary shift towards the study of epilepsies as network phenomena (Caciagli *et al.*, 2014).

In MTLE, as expected, we observed hippocampal volume abnormalities ipsilateral to the patient's side of seizure onset. Neither MTLE-L nor MTLE-R showed significant contralateral hippocampal volume reductions, confirming that sporadic, unilateral MTLE is not routinely underpinned by bilateral hippocampal damage (Blümcke et al., 2013). Both MTLE groups showed extrahippocampal abnormalities in the ipsilateral thalamus and pallidum, with widespread reductions in cortical thickness, supporting a growing body of literature indicating that MTLE, as an example of a specific disease constellation in the epilepsies, is also a network disease, extending beyond the mesial temporal regions (Keller et al., 2014; de Campos et al., 2016). Disruption of this network, notably in the thalamus (Keller et al., 2015; He et al., 2017) and thalamo-temporal white matter tracts (Keller et al., 2015, 2017), may be associated with postoperative seizure outcome in MTLE.

Patients with left and right MTLE showed distinct patterns of structural abnormalities when compared to controls, resolving conflicting findings from smaller studies, some reporting an equal distribution of structural differences (Liu et al., 2016), and others indicating more diffuse abnormalities, either in left MTLE (Keller et al., 2002, 2012; Bonilha et al., 2007; Kemmotsu et al., 2011; de Campos et al., 2016) or in right MTLE (Pail et al., 2009). The structural differences observed in the present study may reflect a younger age at onset of epilepsy in left MTLE, which occurred, on average, 1.2 years earlier than those with right MTLE (Supplementary Table 20). Independent, large-scale studies of MTLE patients have confirmed a significantly earlier age at onset in left, compared to right, MTLE (Blümcke et al., 2017). Durationrelated effects were also observed in right, but not left, MTLE, pointing to possible biological distinctions between the two.

In IGE, a clinically and biologically distinct group of epilepsies typically associated with 'normal' MRI on clinical inspection (Woermann *et al.*, 1998), we identified reduced volume of the right thalamus, and thinner precentral gyri in both hemispheres, supporting prior reports of structural (Bernhardt *et al.*, 2009a), electroencephalographic, and functional (Gotman *et al.*, 2005) abnormalities in IGE. These IGE cases were considered typical by reviewing neurologists, suggesting that this common type of epilepsy is also associated with quantifiable structural brain abnormalities.

The precentral gyri, site of the primary motor cortex, showed bilateral structural deficits across all epilepsy groups (all-epilepsies, IGE, MTLE-L, MTLE-R, and all-other-epilepsies), without detectable inter-cohort or between-disease heterogeneity (Supplementary Figs 3–12).

Atrophy of the motor cortex has been linked to seizure frequency and duration of epilepsy in MTLE (Coan *et al.*, 2014); here, we observed a negative correlation between precentral (and postcentral) grey matter thickness and duration of epilepsy in the aggregate all-epilepsies group.

The right thalamus also showed evidence of structural compromise across all epilepsy cohorts, re-emphasizing the importance of the thalamus as a major hub in the epilepsy network (He et al., 2017; Jobst and Cascino, 2017). Loss of feed-forward inhibition between the thalamus and its neocortical connections may be epileptogenic (Paz and Huguenard, 2015), and thalamocortical abnormalities have previously been reported in IGE (Gotman et al., 2005; Bernhardt et al., 2009a; O'Muircheartaigh et al., 2012) and MTLE (Mueller et al., 2010; Bernhardt et al., 2012). These findings support prior 'system epilepsies' hypotheses of pathophysiology (Avanzini et al., 2012), suggesting that a broad range of common epilepsies share vulnerability within a thalamocortical structural pathway involved in, and likely affected by, seizures (Liu et al., 2003; Bernhardt et al., 2013). Given this study's cross-sectional design, we cannot determine if these are causative changes, consequences of recurrent seizures, prolonged drug treatment, or a combination of factors. The epilepsies, as a broad group, may involve progressive structural change (Caciagli et al., 2017), indicating the need for large-scale longitudinal studies.

A heterogeneous subgroup of individuals without confirmed diagnoses of IGE or MTLE with hippocampal sclerosis showed similar patterns of structural alterations to those observed in the aggregate all-epilepsies cohort. The findings included enlarged ventricles, smaller right pallidum and right thalamus, and reduced thickness across the motor and frontal cortices. Hippocampal abnormalities were not observed in this subgroup, suggesting that the patterns of reduced hippocampal grey matter observed in the aggregate group were driven by the inclusion of MTLEs with hippocampal sclerosis. Unlike the IGE, MTLE, and aggregate epilepsy cohorts, this subgroup also showed bilateral enlargement of the amygdala—a phenomenon previously reported in non-lesional localization-related epilepsies (Reves et al., 2017) and non-lesional MTLEs (Takaya et al., 2012; Coan et al., 2013). Non-lesional MTLEs formed a large proportion of this 'all-other-epilepsies' cohort (43.3%; 445 individuals), but the subgroup included many other focal and unclassified syndromes, potentially obscuring specific biological interpretations. Future, sufficiently powered studies will stratify this cohort into finer-grained subtypes to delineate syndrome-specific effects.

Despite its international scale, our study has limitations. All results were derived from cross-sectional data: we cannot distinguish between historical acute damage and progressive abnormalities. We cannot disentangle the relative contributions of environmental and treatment-related factors, including antiepileptic medications, seizure types and frequencies, disease severity, language dominance, and other initial precipitating factors. On average, duration

of epilepsy was at least 10 years; longitudinal investigations of new-onset and paediatric epilepsies will provide a more comprehensive understanding. Despite using standardized image processing protocols, quality control, and statistical techniques, some brain measures showed a wide distribution of effect sizes across research centres, which may reflect sample heterogeneity and differences in scanning protocols (Supplementary material).

We observed modest thickness differences across the majority of cortical regions; Cohen's d effect sizes ranged from small to moderate (d = 0.2-0.5), with some very small effects (d < 0.2) noted in the right pars opercularis, bilateral pars triangularis, and bilateral transverse temporal gyri of the aggregate all-epilepsies group. Other large-scale ENIGMA studies have reported similarly modest (albeit less widespread) cortical abnormalities in psychiatric illnesses including major depression (Schmaal et al., 2016) and bipolar disorder (Hibar et al., 2017b). Although epilepsy is characterized by an enduring predisposition to generate abnormal excessive or synchronous neuronal activity in the brain (Fisher et al., 2014), our findings indicate that common epilepsies are associated with widespread, but relatively subtle, structural alterations of the neocortex. Replication in independent MRI cohorts, complemented by advanced imaging modalities and large-scale gene expression datasets, will help elucidate how these cortical abnormalities relate to underlying disease processes.

Overall, in the largest neuroimaging analysis of epilepsy to date, we demonstrate a pattern of robust brain structural abnormalities within and between syndromes. Specific functional interpretations cannot be inferred from grey matter differences, but lower volume and thickness measures may reflect tissue loss, supporting recent observations that the common epilepsies cannot always be considered benign (Gaitatzis et al., 2004; Bell et al., 2016; Devinsky et al., 2017). The study provides a macroscopic neuroanatomical map upon which neuropathological work, animal models, and further gene expression studies, can expand. Our consortium plans to investigate more specific neuroanatomical traits and epilepsy phenotypes, explore sophisticated shape and sulcal measures, and eventually conduct genome-wide association analysis of brain measures, to improve our understanding and treatment of the epilepsies.

Web resources

All image processing, quality assurance, and statistical analysis protocols for this study can be downloaded from the ENIGMA website, at: http://enigma.usc.edu/ongoing/enigma-epilepsy/enigma-epilepsy-protocols/.

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Supplementary material

Supplementary material is available at Brain online.

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