The genomic landscape across 474 surgically accessible epileptogenic human brain lesions

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Understanding the exact molecular mechanisms involved in the aetiology of epileptogenic pathologies with or without tumour activity is essential for improving treatment of drug-resistant focal epilepsy. Here, we characterize the landscape of somatic genetic variants in resected brain specimens from 474 individuals with drug-resistant focal epilepsy using deep whole-exome sequencing (>350×) and whole-genome genotyping. Across the exome, we observe a greater number of somatic single-nucleotide variants in low-grade epilepsy-associated tumours (7.92 ± 5.65 single-nucleotide variants) than in brain tissue from malformations of cortical development (6.11 ± 4 single-nucleotide variants) or hippocampal sclerosis (5.1 ± 3.04 single-nucleotide variants). Tumour tissues also had the largest number of likely pathogenic variant carrying cells. Low-grade epilepsy-associated tumours had the highest proportion of samples with one or more somatic copy-number variants (24.7%), followed by malformations of cortical development (5.4%) and hippocampal sclerosis (4.1%). Recurring somatic whole chromosome duplications affecting Chromosome 7 (16.8%), chromosome 5 (10.9%), and chromosome 20 (9.9%) were observed among low-grade epilepsy-associated tumours. For germline variant-associated malformations of cortical development genes such as TSC2, DEPDC5 and PTEN, germline single-nucleotide variants were frequently identified within large loss of heterozygosity regions, supporting the recently proposed ‘second hit’ disease mechanism in these genes. We detect somatic variants in 12 established lesional epilepsy genes and demonstrate exome-wide statistical support for three of these in the aetiology of low-grade epilepsy-associated tumours (e.g. BRAF) and malformations of cortical development (e.g. SLC35A2 and MTOR). We also identify novel significant associations for PTPN11 with low-grade epilepsy-associated tumours and NRAS Q61 mutated protein with a complex malformation of cortical development characterized by polymicrogyria and nodular heterotopia. The variants identified in NRAS are known from cancer studies to lead to hyperactivation of NRAS, which can be targeted pharmacologically. We identify large recurrent 1q21–q44 duplication including AKT3 in association with focal cortical dysplasia type 2a with hyaline astrocytic inclusions, another rare and possibly under-recognized brain lesion. The clinical-genetic analyses showed that the numbers of somatic single-nucleotide variant across the exome and the fraction of affected cells were positively correlated with the age at seizure onset and surgery in individuals with low-grade epilepsy-associated tumours. In summary, our comprehensive genetic screen sheds light on the genome-scale landscape of genetic variants in epileptic brain lesions, informs the design of gene panels for clinical diagnostic screening and guides future directions for clinical implementation of epilepsy surgery genetics.
Genomic landscape of epileptogenic lesions

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Introduction
Drug-resistant epilepsies due to focal brain lesions represent a huge health burden and challenge for everyday clinical practice. Neurosurgical resection strategies have proven helpful in carefully selected patients, especially for brain lesions visible through MRI and confirmed by histopathology diagnosis. The most common types of epilepsy-associated brain lesions comprise hippocampal sclerosis (HS), low-grade epilepsy-associated developmental brain tumours (LEAT) such as ganglioglioma (GG) and dysembryoplastic neuroepithelial tumours (DNET), and malformations of cortical development (MCD) such as focal cortical dysplasia (FCD). Overall, these three most common categories accounted for >80% of the almost 10000 patients who underwent epilepsy surgery and were collected in the European Epilepsy Brain Bank. Genetic factors have been associated with many common and rare epilepsies. Until recently, specific genes with variants of large effects have mainly been discovered in rare and severe forms of paediatric epilepsy that typically do not show structural abnormalities on MRI. In the past decade, >15 genes have been associated with somatic variants in epilepsy-associated MCD and LEAT. In contrast and to the best of our knowledge, the burden of somatic variants has not been systematically evaluated yet for HS. For MCD, somatic variants in genes encoding proteins of the canonical PI3K-AKT-MTOR pathway and germline loss-of-function variants—with or without co-occurring somatic losses of heterozygosity—in genes encoding proteins of the GATOR complex, a negative regulator of the PI3K-AKT-MTOR pathway, with or without co-occurring somatic losses of heterozygosity—in genes encoding proteins of the GATOR complex, a negative regulator of the PI3K-AKT-MTOR pathway, have been associated with FCD type 2 and hemimegalencephaly (HME). In addition, the SLC35A2 gene has been associated with the recently discovered mild MCD with oligodendroglial hyperplasia in the epilepsy (MOGHE) disease entity. For LEAT, somatic variants that affect the RAS-RAF-MAPK pathway (i.e. BRAF and FGFR1) play a major role, with the BRAF variant V600E reported in 18–56% of GG and pathogenic variants in FGFR1 in 58–82% of DNET. There is also emerging evidence for somatic variants in the RAS-RAF-MAPK pathway underlying MCD, with recent case studies reporting...
variants in KRAS associated with epilepsy-associated tumours and malformations.23–25

The search for actionable treatment targets in drug-resistant focal epilepsies pushed genetic studies of epilepsy-associated brain lesions into an emerging field. Most current studies use ultra-deep targeted sequencing with coverages >1000x to identify low allelic fraction disease-causing single-nucleotide variants (SNV) with high sensitivity and specificity. Due to the high coverage and correlated high sequencing cost, current studies typically prioritized targeted over whole-exome sequencing (WES).16,26 Nevertheless, this approach has been very successful, leading to the discovery of 15 le- sional epilepsy-associated genes.5,14–16 However, such candidate gene approaches did not interrogate the mutational signature for a structural lesion across the whole exome or genome, which is a typ- ical study design in cancer research.27,28 As a result, the mutation rate and genetic architecture across epileptic brain lesion categories have not been systematically described. Subsequently, it is unclear whether MCD are single gene or oligogenic somatic disorders and whether somatic variation contributes to the development of HS. In addition, the role of genome-wide postzygotic copy-number variants (CNV) in different epileptic lesions has not been investigated. Recent evidence suggests that low-frequency mosaicism of variants in germline epilepsy-associated ion-channel encoding genes (e.g. SCN8A) can cause epilepsy in rodent models.29 However, it is unclear whether somatic variants in these genes can cause focal epilepsy in humans. It is also unclear whether variants in the 15 recently identi- fied genes5,14–16 represent the most common genetic cause of leional epilepsy as most previous studies were gene panel candidate studies or small-scale exome-wide studies, underpowered for statistical gene burden analyses across the exome.16,26

Here, we explored the exome-wide somatic variant burden and the genome-wide CNV burden of the three most common categories of epileptogenic brain lesions (i.e. LEAT, MCD and HS). We ana- lysed differential variant burden across lesion categories, statistically identified novel epileptogenic pathology-associated genes, explored the role of driver mutations observed in cancer and refined genotype–phenotype associations. The results of our study will shed light on the genetic architecture for a lot of epilepto- genic brain lesions and inform the design of diagnostic genetic tests with reliably high sensitivity and specificity.

Materials and methods

Study cohorts

Snap-frozen surgical brain tissue samples obtained from 474 indivi- duals with epilepsy-associated brain lesions (223 MCD, 154 LEAT and 97 HS) were retrieved from the Cleveland Clinic Epilepsy Center biorepository (n = 154) and European Epilepsy Brain Bank consortium (n = 320). All studies were performed following institutional guidelines and regulations regarding research involving human subjects and approved by the ethics review boards of the Cleveland Clinic and the University of Erlangen, Germany. All pa- tients underwent a comprehensive presurgical evaluation followed by a discussion at a multidisciplinary patient management confer- ence where the surgical strategy was approved. Histopathological review of all resected brain tissue was performed and interpreted by board-certified clinical neuropathologists at the Cleveland Clinic in all patients and followed by a detailed re-review by experi- enced neuropathologists (I.B. and R.C.) using the International League against Epilepsy (ILAE) classification scheme for FCD,30 HS31 and the 4th edition of the WHO classification of tumours of the central nervous system.30 Characteristics of our study cohorts are detailed in Table 1. Genomic DNA was extracted from fresh-frozen brain tissue for all 474 individuals. The DNeasy Blood and Tissue Kit (Qiagen) was used for DNA extraction from fresh-frozen brain sam- ples according to the manufacturer’s protocol.

Somatic and germline SNV calling

Deep WES (350x) of all samples in this study was performed using Agilent SureSelect Human All Exon V7 enrichment and paired-end reads (151 bp) Illumina sequencing. All paired-end FASTQ files were aligned to the GRCh37/hg19 human reference genome, including the hs37d5 decay sequence using BWA-MEM,31 following GATK (Genome Analysis Toolkit) best practices.32 We used MuTect233 to generate a panel of normals (PoN) from 24 additional surgically re- moved and sequenced brain samples to exclude sequencing and alignment artefacts. According to GATK best practices guidelines,34 these 24 brain samples were best suited for the PoN on the basis of their technical properties (i.e. sequenced with the same method- ology, platform and in the same batch as the study samples). All 24 samples included in the PoN underwent surgery for epilepsy and were histopathologically confirmed as non-lesional or due to an external insult (glial scar or encephalitis, detailed in Supplementary Table 1) and thus had a low likelihood of carrying overgrowth disorder or cancer-driver variants that are predominant- ly involved in MCD or tumours. We then called somatic SNV,

Table 1 Study cohort

<table>
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<th>Total</th>
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The bold highlighted values represent the numbers of individuals with one of the three main classes of epilepsy-associated brain lesions. GCF = Cleveland Clinic Foundation; EEEB = European Epilepsy Brain Bank; mMCD = mild MCD; NOS = not otherwise specified; PMG = polymicrogyria; NH = nodular heterotopia; MVNT = multinodular vacuolated neuronal tumour.
insertions and deletion polymorphisms (indels) using MuTect2 in conjunction with the PoN. We only retained somatic variants with a maximum minor allele frequency (MAF) $< 10^{-5}$ in the Genome Aggregation Database (gnomAD). Variants within segmental duplication regions or non-diploid regions were removed. Low-quality calls flagged by MuTect2 with ‘t_lod_fstar’, ‘str_contraction’ and ‘triallelic_site’ were removed. Finally, we excluded somatic indels within RepeatMasker or simple repeat regions.

We then used MosaicForecast to perform read-based phasing and identify high-confidence somatic mosaic variants from all called variants. MosaicForecast is a machine-learning-based method optimized to detect somatic mutations without a matched reference tissue. Briefly, the method trains a random forest model on read-based features from phased variants and has been shown to reliably identify somatic mosaic mutations with variant allelic fractions (VAF) as low as 2%. Therefore, the method has the sensitivity to detect variants present in as few as 4% of cells in a given sample of brain tissue. After all training and filtration, we identified 217,560 putative somatic mosaic SNV across all samples. We then filtered all exonic or splice site variants with a VAF between 0.02 and 0.40 and an alternative read depth ≥ 3. Only non-recurring mutations within the cohort were considered, except for variants in 15 genes that have been previously associated with epileptogenic lesions: MTOR, SLC35A2, AKT3, PIK3CA, RHEB, TSC1, TSC2, NPRL2, NPRL3, DEPDC5, Pten, Braf, Fgrf1, Myb and MYBL1. Last, to ensure that no potentially meaningful variants in established lesional epilepsy genes were mistakenly filtered out, we used the Integrated Genomics Viewer (IGV) browser to visually examine all somatic variants in established epileptic lesion genes called by MuTect2-PoN, but judged as potential artefacts by MosaicForecast. However, such variants were not included in the somatic variant enrichment analyses and were only included when reporting the final yield from our screen.

Finally, we used GATK’s HaplotypeCaller (GATK v.4.1.9.0) following GATK best practices to identify germline variants in lesional epilepsy-associated genes in negative regulators shown to act through a germline or two-hit mechanism (i.e. TSC1, TSC2, DEPDC5, NPRL2, NPRL3 and PTEN). We also used GATK’s HaplotypeCaller to identify somatic VAF > 0.30 in other lesional epilepsy-associated genes (i.e. MTOR, SLC35A2, AKT3, PIK3CA, RHEB, BRAF, FGFR1, MYB and MYBL1). HaplotypeCaller may have missed the resulting variants were filtered for variants with: (i) phred quality score (QUAL) < 30; (ii) genotype type quality (GQ) < 99; (iii) sample read depth (DP) ≥ 30; (iv) max DP < 1000; and (v) GATK truth sensitivity tranche ≥ 99.5% for SNV and ≥ 95% for indels and were not included in the somatic variant enrichment analyses (See ‘Materials and methods’ section ‘Exome-wide statistical identification of genes associated with epileptogenic brain lesions’).

**Somatic CNV and CNN-LOH calling**

A total of 688,032 single-nucleotide polymorphisms (SNPs) were genotyped for all samples of this study using the Global Screening Array with Multi-disease drop-in (GSA-MD v.1.0; Illumina). The SNP data were used to detect somatic CNV using MoChA. MoChA is available as a bcftools extension that uses phased VCF files with 8 allele frequency (BAF) and log R ratios (LRR) to identify somatic CNV and copy-number neutral loss of heterozygosity (CNN-LOH). MoChA uses a three-state hidden Markov model to capture somatic-CNV-induced deviations in allelic balance ($\Delta$BAF) at heterozygous sites. After affine-normalization and GC wave-correction, the BAF and LRR values were transformed as described elsewhere. Autosomes and sex chromosomes were separately considered. Before phasing with Eagle v.2.48 we generated a list of variants that were excluded from modelling by both Eagle and MoChA using the following parameters: (i) segmental duplications with low divergence (<2%); (ii) high levels of missingness (>2%); (iii) variants with excess heterozygosity ($P < 10^{-6}$); and (iv) variants that unexpectedly correlate with sex ($P < 10^{-6}$). After somatic CNV calling, we removed samples on the basis of the following quality control (QC) parameters: (i) cell rates <0.97 and (ii) baf_auto > 0.3. We then removed variants on the basis of the following QC parameters: (i) lod_baf_phase < 10 unless the somatic CNV was larger than 5 Mb (or 10 Mb if they span the centromere); (ii) CNV calls flagged as germline CNV; and (iii) possible constitutional duplications with length > 10 Mb and either LRR > 0.35 or LRR > 0.2 and $\Delta$BAF > 0.16 or length < 10 Mb and either LRR > 0.2 or LRR > 0.1 and $\Delta$BAF > 0.1. The estimation of the allelic fraction for each somatic CNV or CNN-LOH by MoChA has been detailed elsewhere. All somatic CNV and CNN-LOH were examined visually by plotting the LRR, BAF and phased BAF values with MoChA.

**Variant annotation and assessment of deleteriousness**

We applied different strategies for the identified CNV and SNV to assess the likelihood of a deleterious effect on disease-relevant loci or genes. We used ANNOVAR with custom databases to perform variant annotation. The deleteriousness of SNV used to characterize the genetic architecture of the different brain lesion categories was assessed on the basis of two filters: (i) variant type and population frequency; and (ii) predicted variant deleteriousness. The frequency filter was based on the variant not being present at an AF > 10^{-5} in the gnomAD database. From the remaining variants, we selected only variants with a high-confidence prediction to be deleterious using the following criteria: (i) loss-of-function (LoF) variants ranked in the top 1% most deleterious variants in the human genome according to the Combined Annotation Dependent Depletion score (CADD, scaled CADD phred score ≥ 20) found in established lesional epilepsy genes or highly LoF-intolerant genes (genes with a gnomAD LoF Observed/Expected Upper bound Fraction LOEUF ≤ 0.35); and (ii) missense variants ranked in the top 1% most deleterious variants in the human genome (scaled CADD phred score > 20) in established lesional epilepsy genes or found in missense-constrained sites (MPC score ≥ 2) of missense intolerant genes (gnomAD missense Z-score > 3.09, corresponding to P < 10^{-5}). The supporting aligned reads of all reported variants were visually inspected using the IGV browser.

Somatic CNV were annotated with triplosensitivity, haploinsufficient and lesonal epilepsy genes. Somatic CNV or CNN-LOH that had a size > 1 Mb were considered deleterious. Deleterious CNV that overlapped genes associated with lesional epilepsies (i.e. MTOR, SLC35A2, AKT3, PIK3CA, RHEB, TSC1, TSC2, NPRL2, NPRL3, DEPDC5, Pten, Braf, Fgrf1, Myb and MYBL1) were considered pathogenic and are reported in Fig. 2C.

**Exome-wide statistical identification of genes associated with epileptogenic brain lesions**

To detect genes under evolutionary mutational selection, we used a Poisson-based dN/dS model using dNdScv. This model tests the normalized ratio of non-synonymous (missense, nonsense and splicing) over background (synonymous) mutations while
correcting for sequence composition and mutational signatures. This method has been shown to reliably identify genes under positive selection in cancer and normal tissues. A global false discovery rate adjusted \( P \)-value \( q \leq 0.1 \) was used to identify statistically significant non-synonymous variant-enriched genes and an unadjusted \( P \leq 0.005 \) to identify nominally significant non-synonymous variant-enriched genes. Genes that showed a suggestive enrichment of non-synonymous variants were only considered when their biological role and known disease associations matched the associated phenotype. To improve the quality of our results, the reads for all variants in genes identified by dNdScv were visually inspected using the IGV browser to assess variant quality. Low-quality variants were flagged as potential artefacts, and genes with only low-quality variants were excluded.

**Post hoc genetics-informed histopathological review**

In addition to the evaluation on enrolment, we performed an in-depth post hoc histopathological review of all mutation-positive samples in the context of their genetic diagnosis. All slides and formalin-fixed paraffin-embedded tissue blocks were retrieved from the laboratory archives. Following microscopic review of the H&E stained sections, additional immunohistochemical stainings recommended by the ILAE for a comprehensive neuropathologic workup of epilepsy surgery brain tissue were added to the review where necessary.

**Statistical analysis**

All statistical analyses and filtering were done using R. We used the Chi-square, Fisher’s exact or Wilcoxon rank-sum test where appropriate. Correlation statistics were generated with Pearson’s correlation.

**Data availability**

All somatic variant calls are available from the supplementary material. All code is available at https://github.com/LalResearchGroup. Additional data are available from the corresponding author on reasonable request.

**Results**

**Cohort overview**

We extracted DNA from snap-frozen brain tissue samples of 474 individuals with drug-resistant epilepsy who underwent epilepsy surgery and were diagnosed with histopathologically confirmed HS, MCD or LEAT (Table 1). HS and LEAT samples primarily originated from the temporal lobe, whereas MCD samples were primarily derived from extratemporal resections and had the largest proportion of multilobar involvement (Table 2). The average age at seizure onset for MCD patients was significantly earlier than that of HS and LEAT (MCD = 6.3 years, HS = 13.73 years, LEAT = 13.14 years; Wilcoxon rank-sum test: MCD versus HS \( P = 1.17 \times 10^{-7} \), MCD versus LEAT \( P = 2.27 \times 10^{-10} \)). Conversely, the average age at surgery for HS patients was significantly later than that of MCD and LEAT (HS = 39.13 years, MCD = 18.48 years, LEAT = 21.82 years; Wilcoxon rank-sum test: HS versus MCD \( P = 1.02 \times 10^{-14} \), HS versus LEAT \( P = 9.59 \times 10^{-15} \), MCD versus LEAT \( P = 0.17 \)). Last, individuals with HS had a longer duration of epilepsy before undergoing surgery, followed by individuals with MCD, and with LEAT (HS = 25.81, MCD = 12.24, LEAT = 8.53 years; Wilcoxon rank-sum test: HS versus MCD \( P = 4.33 \times 10^{-13} \), HS versus LEAT \( P = 6.63 \times 10^{-19} \), MCD versus LEAT \( P = 1.31 \times 10^{-9} \)).

**Somatic SNV profiles differ across the major categories of epilepsy-associated brain lesions**

Deep WES achieved an average mean target coverage of 364\( \times \) across all 474 samples. DNA quality and experimental performance across pathology groups were similar, with no differences in mean coverage among the three phenotype groups (Fig. 1A). On average, the number of somatic SNV carried per sample was higher in LEAT (7.92 ± 5.65) and MCD (6.11 ± 4) than in HS (5.1 ± 3.04; Fig. 1B). From a total of 3078 somatic SNV, 172 SNV affecting 141 individuals were bioinformatically classified as potentially deleterious. These included 128 missense variants and 44 protein-truncating variants. Overall, a higher proportion of LEAT (48.1%, 74/154) had at least one potentially deleterious SNV, compared to MCD (26.9%, 60/223) and HS (6.2%, 6/97; Fig. 1C). The average VAF of somatic SNV was 1.18-fold higher in LEAT (average VAF = 0.078) than in MCD (average VAF = 0.066) and 1.26-fold higher than in HS (average VAF = 0.062). MCD and HS had similar VAF distributions (Fig. 1D). When considering only potentially deleterious variants, the average VAF in LEAT (average deleterious VAF = 0.108) was 1.63-fold higher than in MCD (average deleterious VAF = 0.066) and 1.22-fold higher than in HS (average deleterious VAF = 0.088; Fig. 1D). Overall, MCD and HS showed similar somatic SNV profiles, although MCD carried a nominally significant higher number of somatic SNV per sample and a larger proportion of samples with at least one deleterious missense SNV more than HS (Fig. 1B and C).

**Somatic CNV profiles differ across the major categories of epilepsy-associated brain lesions**

After QC, we identified a total of 153 large CNV and CNN-LOH across 56 samples: 105 duplications (85 whole chromosome gains), 27 deletions (5 whole chromosome losses) and 21 CNN-LOH. LEAT had the highest proportion of samples with one or more somatic CNV or CNN-LOH (38/154, 24.7%), followed by MCD (12/223, 5.4%) and HS (4/97, 4.1%; Fig. 1E). LEAT were also enriched for CNV duplications and deletions compared to MCD and HS, whereas CNN-LOH were not distributed differently across all three categories. Among LEAT, somatic duplications were the most frequently observed type of CNV, with 71% (74/105) of CNV or CNN-LOH-positive samples carrying one or more somatic duplications. Somatic whole chromosome duplications were the major type of duplications among LEAT (82/101 LEAT gain CNV, 81.2%; Fig. 1E). Conversely, somatic CNN-LOH were predominantly among MCD and HS (8/12 CNV-positive MCD, 66.7%; 3/4 CNV-positive HS, 75%).

Most somatic CNV-positive LEAT had multiple somatic CNV (27/38, 71%), whereas only two (2/12, 16.7%) CNV-positive MCD had multiple somatic CNV or CNN-LOH (Fig. 1F). No CNV-positive HS sample carried multiple somatic CNV or CNN-LOH. The highest average VAF of somatic CNV and CNN-LOH was identified in LEAT (average VAF = 0.125), followed by MCD (average VAF = 0.058) and HS (average VAF = 0.028; Fig. 1G). Last, LEAT samples carried larger somatic CNV and CNN-LOH (average size = 108.6 Mbp) than MCD (average size = 59.6 Mbp) and HS (average size = 25.3 Mbp; Fig. 1H). Overall, LEAT showed the most distinct CNV profile; they had a larger proportion of samples with CNV or CNN-LOH, were enriched for duplications and deletions, tended to carry more CNV per sample and had larger CNV affecting a higher number of cells.
The number of somatic SNV and the fraction of mutated tumour cells are associated with age at seizure onset and age at surgery in LEAT

For LEAT, we identified positive correlations between the number of somatic SNV per sample and later age at seizure onset and older age at surgery (Pearson’s correlation; LEAT total somatic SNV and age at surgery: \( r = 0.26, P = 1.16 \times 10^{-3} \); LEAT total somatic SNV and age at onset: \( r = 0.22, P = 9 \times 10^{-3} \)). Similarly, we also identified positive correlations between higher somatic VAF (a proxy marker for the number of mutated cells) and later age at seizure onset and older age at surgery for LEAT (Pearson’s correlation; LEAT somatic VAF and age at surgery: \( r = 0.16, P = 1.47 \times 10^{-3} \); LEAT somatic VAF and age at onset: \( r = 0.18, P = 1.44 \times 10^{-5} \)). This signal was driven primarily by the somatic SNV VAF, with the VAF of the CNV and CNN-LOH having little effect (Pearson’s correlation; LEAT somatic SNV VAF and age at surgery: \( r = 0.11, P = 5.55 \times 10^{-3} \)). This signal was driven primarily by the somatic SNV VAF, with the VAF of the CNV and CNN-LOH having little effect (Pearson’s correlation; LEAT somatic SNV VAF and age at surgery: \( r = 0.11, P = 5.55 \times 10^{-3} \)).

Unbiased somatic variant burden analysis confirms previously reported and discovers novel lesional epilepsy-associated genes

We performed the first exome-wide burden analysis for a lesional epilepsy cohort (HS, MCD and LEAT) to identify genes under positive mutational selection with significant enrichment of somatic SNV in any of the three main lesion categories. In line with previous results from candidate gene studies, we identified four genes with significant somatic SNV burden for LEAT and MCD and no genetic associations with HS (Supplementary Table 2 and Fig. 2A). For LEAT, the gene with the most significant exome-wide somatic SNV burden was \( \text{BRAF} \), which represents the most well-established previously reported LEAT gene and is primarily associated with GG.\(^5,54\) The second gene identified with a significant variant enrichment was \( \text{PTPN11} \), which has not been previously associated with LEAT. The \( \text{PTPN11} \) gene encodes Protein Tyrosine Phosphatase Non-Receptor Type 11, an upstream regulator of the RAS/MAPK and mTOR signalling pathways. Activating mutations in \( \text{PTPN11} \) have been shown to cause Noonan syndrome (a congenital developmental syndrome), play a role in tumorigenesis and affect the development of white matter microstructure in humans.\(^5,57\) For
MCD, our burden analysis identified SLC35A2 and MTOR, which have been previously reported in multiple candidate gene studies or descriptive exome analysis screens as the two most frequently mutated genes in mild MOGHE and FCD type 2, respectively. However, we are the first to use an unbiased statistical approach to show that variants in these genes are a major genetic cause of MCD entities.

Identification of additional lesional epilepsy genes with suggestive statistical support and high biological plausibility

The previous statistical analysis could identify and validate the most frequently mutated lesional epilepsy genes that remained significant after controlling for the false discovery rate ($q \leq 0.1$). Next, we investigated variants in genes with suggestive variant enrichment ($P < 5 \times 10^{-3}$) for association with LEAT or MCD by considering additional evidence criteria (see ‘Materials and methods’ section and Supplementary Table 2). We also screened our cohort for variants in 579 high-confidence cancer-driver sites, previously identified in 53 genes (Supplementary Table 3). The genes AKT3, NRAS and PIK3CA had nominally significant mutational burden and/or variants in cancer-driving sites and met additional disease-association criteria for a role in the aetiology of MCD (Supplementary Tables 2 and 3). AKT3 and PIK3CA are well-established MCD genes and the identified variants have been molecularly characterized as activating variants.

Large somatic chromosomal alterations in lesional epilepsies frequently include established lesional epilepsy-associated genes

We identified several somatic CNV patterns and recurring regions across the lesion categories represented in our cohort. In LEAT specifically, the most frequently recurring duplicated regions were whole chromosome duplications of chromosome 7 (17/101, 16.8%), chromosome 5 (11/101, 10.9%) and chromosome 20 (10/...
We also identified a recurring CNV gain of the q-arm of chromosome 1 in our MCD cohort ($n=2$). Duplications of chromosome 1q have been previously identified in cases of macrocephaly, megalencephaly, polymicrogyria and FCD. The most frequently recurring regions of CNN-LOH mapped to the longer q-arm of chromosome 22 (5/21, 23.8%) and chromosome 16p13.3 (4/21, 19%; Fig. 3A). These regions were most prevalent in MCD, but were also identified in HS or LEAT. A single recurring region of CNN-LOH was unique to HS, which mapped to the q-arm of chromosome 19 (19q13.42–13.43).

To pinpoint the putative causative genes, we investigated whether the identified CNV and CNN-LOH overlapped with previously established lesional epilepsy genes. We observed that 68.4% of LEAT, 83.3% of MCD and 50% of HS with a somatic CNV or CNN-LOH carried at least one chromosomal alteration affecting one of the 15 established lesional epilepsy genes (chromosome 7) overlapped with the most frequently mutated gene in lesional epilepsies (BRAF). Additionally, the two most frequently recurring regions of CNN-LOH included DEPDC5 (22q) or TSC2 (16p13.3), both established lesional epilepsy genes that require a somatic second hit such as a CNN-LOH to cause disease.

**Table 2 Clinical characteristics of the study cohort**

<table>
<thead>
<tr>
<th></th>
<th>HS ($n=97$)</th>
<th>MCD ($n=223$)</th>
<th>LEAT ($n=154$)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sex</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>46 (47.4%)</td>
<td>107 (48%)</td>
<td>92 (59.7%)</td>
</tr>
<tr>
<td>Female</td>
<td>51 (52.6%)</td>
<td>116 (52%)</td>
<td>62 (40.3%)</td>
</tr>
<tr>
<td><strong>Age of onset (years)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average</td>
<td>$13.73 \pm 13.25$</td>
<td>$6.32 \pm 8.61$</td>
<td>$13.17 \pm 12.98$</td>
</tr>
<tr>
<td>Median [IQR]</td>
<td>9.5 [2, 22]</td>
<td>3 [0.5, 9.25]</td>
<td>8.75 [4.25, 18]</td>
</tr>
<tr>
<td><strong>Age at surgery (years)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average</td>
<td>$39.13 \pm 14.15$</td>
<td>$18.5 \pm 14.65$</td>
<td>$21.8 \pm 16.09$</td>
</tr>
<tr>
<td><strong>Duration of epilepsy (years)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average</td>
<td>$25.81 \pm 15.26$</td>
<td>$12.23 \pm 10.89$</td>
<td>$8.49 \pm 9.57$</td>
</tr>
<tr>
<td><strong>Lateralization</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Right</td>
<td>45 (52.3%)</td>
<td>132 (61.1%)</td>
<td>83 (54.6%)</td>
</tr>
<tr>
<td>Left</td>
<td>41 (47.7%)</td>
<td>83 (38.4%)</td>
<td>69 (45.4%)</td>
</tr>
<tr>
<td>Both</td>
<td>0</td>
<td>1 (0.5%)</td>
<td>0</td>
</tr>
<tr>
<td><strong>Localization</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Temporal</td>
<td>97 (100%)</td>
<td>48 (21.8%)</td>
<td>114 (74.5%)</td>
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<tr>
<td>Right</td>
<td>45</td>
<td>28</td>
<td>63</td>
</tr>
<tr>
<td>Left</td>
<td>41</td>
<td>18</td>
<td>50</td>
</tr>
<tr>
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<td>2</td>
<td>1</td>
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<tr>
<td>Frontal</td>
<td>0</td>
<td>96 (43.6%)</td>
<td>17 (11.1%)</td>
</tr>
<tr>
<td>Right</td>
<td>0</td>
<td>60</td>
<td>8</td>
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<td>35</td>
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</tr>
<tr>
<td>Not specified</td>
<td>0</td>
<td>1</td>
<td>0</td>
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<tr>
<td>Parietal</td>
<td>0</td>
<td>11 (5%)</td>
<td>8 (5.2%)</td>
</tr>
<tr>
<td>Right</td>
<td>0</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Left</td>
<td>0</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Not specified</td>
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<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Occipital</td>
<td>0</td>
<td>6 (2.7%)</td>
<td>3 (2%)</td>
</tr>
<tr>
<td>Right</td>
<td>0</td>
<td>5</td>
<td>1</td>
</tr>
<tr>
<td>Left</td>
<td>0</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Multiple Lobes</td>
<td>0</td>
<td>59 (26.8%)</td>
<td>11 (7.2%)</td>
</tr>
<tr>
<td>Right</td>
<td>0</td>
<td>34</td>
<td>6</td>
</tr>
<tr>
<td>Left</td>
<td>0</td>
<td>24</td>
<td>5</td>
</tr>
<tr>
<td>Both</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><strong>Follow-up time (months)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average + SD</td>
<td>31.88 ± 29.4</td>
<td>37.71 ± 34.37</td>
<td>27.2 ± 19.28</td>
</tr>
<tr>
<td>Surgical outcome at last follow-up</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Engel I</td>
<td>47 (81%)</td>
<td>120 (63.8%)</td>
<td>104 (82.5%)</td>
</tr>
<tr>
<td>Engel II</td>
<td>9 (15.5%)</td>
<td>20 (10.6%)</td>
<td>12 (9.5%)</td>
</tr>
<tr>
<td>Engel III-IV</td>
<td>2 (3.5%)</td>
<td>48 (25.6%)</td>
<td>10 (8%)</td>
</tr>
</tbody>
</table>

IQR = Interquartile range, SD = standard deviation.

101, 9.9%; Fig. 3A). We also identified a recurring CNV gain of the q-arm of chromosome 1 in our MCD cohort ($n=2$). Duplications of chromosome 1q have been previously identified in cases of macrocephaly, megalencephaly, polymicrogyria and FCD. The most frequently recurring regions of CNN-LOH mapped to the longer q-arm of chromosome 22 (5/21, 23.8%) and chromosome 16p13.3 (4/21, 19%; Fig. 3A). These regions were most prevalent in MCD, but were also identified in HS or LEAT. A single recurring region of CNN-LOH was unique to HS, which mapped to the q-arm of chromosome 19 (19q13.42–13.43).

To pinpoint the putative causative genes, we investigated whether the identified CNV and CNN-LOH overlapped with previously established lesional epilepsy genes. We observed that 68.4% of LEAT, 83.3% of MCD and 50% of HS with a somatic CNV or CNN-LOH carried at least one chromosomal alteration affecting one of the 15 established lesional epilepsy genes (chromosome 7) overlapped with the most frequently mutated gene in lesional epilepsies (BRAF). Additionally, the two most frequently recurring regions of CNN-LOH included DEPDC5 (22q) or TSC2 (16p13.3), both established lesional epilepsy genes that require a somatic second hit such as a CNN-LOH to cause disease.

**Genetics-informed histopathology review reveals novel genotype–phenotype associations**

In total, we identified 19 genes of interest either through our somatic variant gene discovery analysis (SLC35A2, MTOR, BRAF, PTPN11, AKT3, NRAS, FGFR1, NF1, PIK3CA and KRAS) or previously reported as genes associated with epileptogenic brain lesions (RHEB, PTEN, TSC1, TSC2, DEPDC5, NPRL2, NPRL3, MYB and MYBL1). Across our entire cohort, we identified 151 samples that carried at least one SNV, CNV or CNN-LOH affecting one of...
these genes (31.4% of cohort; 7.2% of HS, 31.8% of MCD and 51.9% of LEAT; Fig. 2C and Supplementary Tables 4 and 5). By considering known pathogenic pathways, we identified 74 samples (64 LEAT, 9 MCD and 1 HS) with variants affecting genes in the RAS-RAF-MAPK pathway (FGFR1, PTPN11, KRAS, NRAS, NF1), 64 samples (32 MCD, 27 LEAT and five HS) with variants in genes of the PI3K-akt-MTOR pathway (PIK3CA, PTEN, AKT3, TSC1, TSC2, RHEB, MTOR) and 31 samples (19 MCD, 11 LEAT and 1 HS) with variants in genes of the GATOR1 complex (DEPDCC5, NPRL2 and NPRL3).

All mutation-positive samples underwent an in-depth post hoc histopathological review that included their genetic diagnosis (Table 3). The genes with the most variants (SNV, CNV or CNV-LOH) which had primarily homogeneous histopathology included: BRAF (52/67 in GG), SLC35A2 (17/17 in MOGHE), FGFR1 (14/14 in DNET) and MTOR (13/19 in FCD 2b or HME). A detailed summary of the genotype–phenotype analysis for the established LEAT and MCD genes is provided in the Supplementary material and Fig. 2C.

Major novel findings of our analysis include eight somatic PTPN11 SNV across seven LEAT samples (three in cancer-driving sites) and eight samples with somatic duplications covering PTPN11 (Fig. 3C). Despite a clear association with LEAT, no specific LEAT histopathological subtype was strongly associated with PTPN11 alterations; variants occurred in GG, DNET and three different glioma subtypes (Fig. 3C and Supplementary Tables 4 and 5).

Second, we observed a novel association between polymicrogyria and variants in NRAS, with 17% (2/12) of polymicrogyria samples having an activating variant in NRAS. Both variants (p.Q61K and p.Q61R) have been well recognized in the literature as cancer-driver variants (Fig. 4A).65,69 Conformational changes of the NRAS Q61 mutated protein lead to a prolonged active state (guanosine-5′-triphosphate GTP-bound) compared to the wild-type form, with variants such as Q61R having an increased affinity for GTP, a slower rate of GTP exchange and a lower rate of intrinsic GTP hydrolysis relative to wild-type (Fig. 4B).70 On microscopic re-review, both samples with NRAS somatic variants had a concordant complex MCD phenotype composed of polymicrogyria and nodular heterotopia with tumour-like glio-neuronal growth patterns (Fig. 4C–H).

Last, we identified two MCD samples with duplications of the q-arm of chromosome 1 (Table 3). Both samples had similar duplications spanning the 1q21-q44 region, including AKT3 (Fig. 5A). Histopathological review revealed that both samples with the 1q21-q44 duplication, including AKT3, had a concordant phenotype of FCD 2a with hyaline astrocytic inclusions, a rare and seldom reported epileptogenic brain lesion (Fig. 5B and C).71 This is a novel genotype–phenotype association for somatic duplications of chromosome 1q and the first genetic association reported for hyaline astrocytic inclusions.

**Discussion**

We generated deep (>350x) WES data to identify somatic SNV and whole-genome genotyping data to identify somatic CNV across surgically resected epileptogenic brain lesions from 474 individuals with focal drug-resistant epilepsy. While other studies performed targeted sequencing of lesional brain tissue from <100 individuals,16 or WES in <130 individuals,72 our study sample represents the largest cohort of epilepsy-associated brain lesions analysed through deep whole-exome analysis. Using this rich source of data, we demonstrated differential somatic variant profiles across LEAT, MCD and HS. For LEAT and MCD, we confirmed previously reported and identified novel pathology-gene associations and explored the role of genetic information in clinical characteristics and histopathological classification.

We observed a differential somatic variant burden across the three most common epileptogenic pathologies (LEAT, MCD and HS). LEAT showed a higher burden of somatic SNV and CNV than MCD or HS. This finding indicates a proliferative advantage of mutated tumour cells and conclusively shows for the first time that the genetic architecture of epilepsy-associated tumours is different from that underlying MCD or HS. We showed that increasing numbers of somatic SNV and fractions of mutated tumour cells are correlated with later age at seizure onset and older age at surgery in LEAT. This finding is in line with studies that found positive correlations between higher somatic burden and later age at diagnosis for all human genes and all cancer-associated genes across all types of cancer.73 We confirmed previously reported and discovered novel lesional epilepsy-associated genes using a combined statistical genetic and biological approach. Our study confirms that somatic variants affecting 12/15 recently established lesional epilepsy genes5,14–16 represent common genetic etiologies of lesional epilepsies (29.1%, 138/474 pathogenic variant carriers in our cohort across 12 genes). We did not observe somatic variants in RHEB, MYB and MYBL1, in which previously identified pathogenic variants are primarily structural rearrangements such as gene fusions and CNV.5,14

Our study could not provide evidence for a significant role of somatic pathogenic SNV and CNV in the aetiology of HS. We analysed 97 individuals with HS, and did not identify any genetic association with HS. However, we identified a small subset of patients with somatic variants in recognized lesional epilepsy genes or the newly described PTPN11. Whether these variants are indeed contributing to rare subtypes of HS needs to be clarified in future studies. To the best of our knowledge, this is the first large-scale exome-level investigation of genetic variants in surgically resected HS tissues.

Most CNV (>69%) affected one of the 15 established lesional epilepsy genes.5,14 Microscopic re-review of these samples led to the identification of a novel genotype–phenotype association between duplications at q21-q44 of chromosome 1 that affects AKT3 and many additional genes and a FCD 2a phenotype with hyaline astrocytic inclusions. This is the first report of a genotype–phenotype correlation for this lesion type and expands the spectrum of MCD phenotypes associated with somatic duplications of chromosome 1q. Compared to our study, previously reported 1q duplications were larger in size and observed in individuals with larger lesions such as polymicrogyria.62,67,68 Furthermore, we identified nine cases with large somatic CNN-LOH regions overlapping lesional epilepsy genes with pathogenic mechanisms known follow the ‘two-hit’ model (TSC1, TSC2, DEPDCC5, PTEN, FGFR1 and NF1).17,74 For five of these (with the exception of TSC1), we identified an accompanying pathogenic germline SNV in the same gene. This result is in line with previous lesional epilepsy studies that identified CNN-LOH as a somatic second hit for germline variants in DEPDCC5, TSC1 and TSC2.17,75 Overall, our findings add to a growing body of evidence that somatic CNV are involved in the aetiology of epilepsy-associated brain lesions.62,76

Our genetic screen identified five LEAT samples carrying a total of six variants in PTPN11, which encodes the Protein Tyrosine Phosphatase Non-Receptor Type 11—a regulator of the RAS-RAF-MAPK signalling pathway. Five of the six PTPN11 variants identified in LEAT have been previously reported in the clinical variant database ClinVar as pathogenic variants associated with Noonan syndrome or various neoplasms (p.A72G, p.E76K, p.D61N, p.D61H,
Figure 3 Recurring somatic CNV and CNN-LOH regions often overlap with lesional epilepsy genes. (A) Recurring regions of overlap among somatic CNV and CNN-LOH. Smaller regions are identified on the basis of included cytobands, CNV or CNN-LOH affecting >80% of a chromosomal arm are identified by the chromosomal arm, and CNV affecting the entire chromosome are identified by the chromosome number. (B) Detected CNV and CNN-LOH often overlap established and newly identified lesional epilepsy genes.

Table 3 Genetic-positive samples where genetic diagnosis had an impact on histopathological review

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Lesion category</th>
<th>Original pathology</th>
<th>Genetic diagnosis</th>
<th>Inheritance (VAF)</th>
<th>Revised pathology</th>
<th>Additional information</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCD_EEBB_10</td>
<td>MCD</td>
<td>PMG</td>
<td>NRAS (p.Q61R)</td>
<td>Somatic (0.13)</td>
<td>Complex MCD</td>
<td>PMG, NH, FCD 2a and GG</td>
</tr>
<tr>
<td>MCD_EEBB_65</td>
<td>MCD</td>
<td>PMG</td>
<td>NRAS (p.Q61K)</td>
<td>Somatic (0.24)</td>
<td>Complex MCD</td>
<td>PMG, NH, FCD 2a and DNET</td>
</tr>
<tr>
<td>MCD_CCF_29</td>
<td>MCD</td>
<td>FCD (NOS)</td>
<td>TSC2 (p.C644X)</td>
<td>Germline (0.46)</td>
<td>Cortical tuber</td>
<td>FCD 2b</td>
</tr>
<tr>
<td>MCD_CCF_55</td>
<td>MCD</td>
<td>FCD (NOS)</td>
<td>TSC2 (p.F1619S)</td>
<td>Germline (0.46)</td>
<td>Cortical tuber</td>
<td>FCD 2b</td>
</tr>
<tr>
<td>MCD_CCF_56</td>
<td>MCD</td>
<td>FCD (NOS)</td>
<td>TSC2 (p.M276Lfs*61)</td>
<td>Germline (0.43)</td>
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<td>FCD 2b</td>
</tr>
<tr>
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<td>FCD 2b</td>
<td>TSC2 (p.H1135Pfs*33)</td>
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<td>FCD 2b</td>
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<td>FCD 2a</td>
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<td>Hyaline astrocytic inclusions</td>
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<td>MOGHE</td>
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</tr>
<tr>
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<td>mMCD</td>
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<td>Germline (0.46)</td>
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<td>DEPDC5 (p.P779A)</td>
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<td>FCD 1a</td>
<td>SLC35A2 (p.Q108X)</td>
<td>Somatic (0.09)</td>
<td>MOGHE</td>
<td>–</td>
</tr>
</tbody>
</table>

NOS = Not otherwise specified; PMG = polymicrogyria; NH = nodular heterotopia.
p.E139D, p.T507K), with the final variant (p.A72V) reported as having ‘conflicting interpretations of pathogenicity’ (Supplementary Table 4). Additionally, three identified variants occurred in established cancer-driver hotspots in **PTPN11** (p.A72 and p.E76). According to the American College of Medical Genetics and Genomics (ACMG) guidelines,77 this finding provides strong evidence of the pathogenicity of the identified variants (see Supplementary Table 6 for full criteria). Furthermore, our exome-wide somatic variant enrichment analysis provides the first statistical evidence for **PTPN11**, a ClinGen recognized criterion for gene to disorder association validity, therefore considering **PTPN11** as a novel disease-associated gene for LEAT.78

In addition to the evidence provided by the association result, the clinical phenotype of previously reported **PTPN11** mutations matches the phenotype of the carriers identified in our study. Heterozygous gain of function mutations in **PTPN11** cause ~50% of all Noonan syndrome cases,79 a germline overgrowth disorder where multiple cases of co-occurring epileptic tumours have been reported.55,80,81 The LEAT samples with somatic **PTPN11** SNVs had other pathogenic somatic SNVs in **TSC2**, **FGFR1**, **NF1** or **BRAF** (Supplementary Table 4). This finding suggests a modifier role for **PTPN11** in LEAT, as shown for other components of the RAS-RAF-MAPK signalling pathway.82

We also identified an individual with HS who carried a likely pathogenic somatic mutation in **PTPN11** (p.S502L; Supplementary Table 6). Our findings are in line with a recent study that reported somatic **PTPN11** mutations in two individuals with FCD type 3, an other non-LEAT epileptogenic lesion.83 Interestingly, **PTPN11** also regulates the PI3K-AKT-MTOR pathway56 and plays a role in the development of human white matter microstructure.56 Therefore, it is possible that mutations in **PTPN11** may also cause other epileptogenic lesions beyond LEAT. However, gene-disease validity criteria78 are currently only support associations with Noonan syndrome, metachondromatosis and certain cancer types.

**Figure 4** Novel complex MCD phenotype is associated with NRAS mutations at Q61. (A) Crystal structure of the wild-type (WT) N-Ras protein. Close-up of the 61st amino acid site highlights changes in intermolecular interactions between different residues at position 61 and nearby positions T35 and Y64. (B) Overlap of the active GTP-bound 3D structures of WT, Q61K and Q61R N-Ras highlighting changes in structure due to the introduced amino acid changes that increase the protein’s affinity to GTP. (C–E) Twenty-year-old female patient with right temporal lobe epilepsy secondary to polymicrogyria seen on MRI. Histopathology examination of the surgical resection sample revealed a complex cortical malformation with nodular heterotopias shown in C, polymicrogyria (arrow in D), and a tumour-like glio-neuronal growth pattern resembling DNET in E. The arrow in E points to a dysplastic neuronal cell element. (F–H) Twenty-five-year-old male patient with left temporal lobe epilepsy and complex malformation seen on MRI. Histopathology examination of the surgical resection sample confirmed a complex cortical malformation with nodular heterotopias shown in F, polymicrogyria (arrow in G) and a tumour-like glio-neuronal growth pattern in H (arrow showing a dysplastic neuronal cell). MAP2 = immunohistochemistry using antibodies directed against microtubule-associated protein 2, NeuN = immunohistochemistry using antibodies directed against the neuronal nucleus epitope, HE = haematoxylin and eosin staining.
Our gene burden analysis identified an enrichment (Fig. 4 and Table 3); and (ii) hyaline astrocytic inclusions (arrows in B and C) next to FCD 2a.

Figure 5 Novel association between large chromosome 1q duplications that include AKT3 and type 2 FCD with hyaline astrocytic inclusions. (A) Genomic coordinates of two large patient somatic duplications at 1q21-q44. Both duplications cover the established lesional epilepsy gene AKT3, as well as additional genes of potential interest. Breakpoints of both somatic duplications are similar. (B and C) Histopathology findings were remarkably similar in both cases, showing hyaline astrocytic inclusions.

including, as of this study, LEAT. Confirming and further elucidating the role of PTPN11 in LEAT as well as other epileptic lesions will require more in-depth evaluations of large lesional epilepsy cohorts as well as comprehensive animal model.86,87

We identified three additional candidate genes for lesional epilepsy within the RAS-RAF-MAPK pathway (i.e. NF1, KRAS and NRAS), all of which had moderate levels of evidence for gene-disease association according to ACMG guidelines based on variant burden and concordance between previously reported and patient phenotypes.77 Our gene burden analysis identified an enrichment of somatic NF1 variants in LEAT. Similar to PTPN11, no particular LEAT pathology was associated with NF1 variants, although alterations of NF1 have been previously identified in GG.5,6 Interestingly, we also identified somatic variants in NF1 in four FCD 2 and one complex MCD (Fig. 2C). Although HS and FCD 3a have been described in a subset of germline neurofibromatosis type 1, our study is the first direct report of NF1 variants in bona fide MCD.86,87 We also identified an individual with sporadic meningioangiomatosis and a brain somatic cancer-driving variant in KRAS. Germline meningioangiomatosis is associated with neurofibromatosis type 2, but no causal gene for the sporadic form is known yet.88 Last, we identified two established cancer-driving SNV in NRAS in two individuals with a distinct and histopathologically concordant complex MCD phenotype composed of polymicrogyria and nodular heterotopia with tumour-like glio-neuronal growth patterns (Fig. 4). The identification of potentially pharmacologically targetable NRAS variants89 in such patients may present a promising target for precision medicine when epilepsy surgery cannot achieve successful seizure control.90

When the genetic information was disclosed for histopathology re-review for all 151 individuals carrying somatic variants in known or novel candidate genes for lesional epilepsy, we identified consistent genotype–phenotype associations not previously recognized. These included: (i) a complex MCD pattern of polymicrogyria with nodular heterotopia and low-grade glio-neuronal tumours associated with NRAS mutations (Fig. 4 and Table 3); and (ii) hyaline astrocytic inclusions91 associated with chromosomal 1q gains (Fig. 5 and Table 3). The recently defined association of MOGHE with SLC35A2 was confirmed in our case series and resulted in a histopathology reclassification of four cases previously diagnosed as mild MCD (mMCD) or FCD1a.89 Notwithstanding, mMCD and FCD1a are the most difficult and challenging differential diagnoses of MOGHE.92 There were four cases with a germline mutation in TSC2 leading to the reclassification of FCD2b to cortical tuber. However, the tuberous sclerosis complex syndrome was not documented in the clinical charts at the time of histopathological diagnosis (Table 3). Our findings exemplify the increasing impact of an integrated molecular-histopathology diagnosis also in the field of focal epilepsies, as was suggested previously92 and already introduced successfully into the clinical management of individuals with brain tumours.20,93 Last, we also observed variants in established lesional epilepsy genes, which were identified and confirmed to occur in pathologies that do not match the previously reported associated phenotype (Fig. 2C and Supplementary Table 4). These included an MTOR variant (p.R360W) in an individual with FCD1a, a germline DEPDC5 variant (p.R78P) in an individual with MOGHE, and a BRAF variant (p.I61V) in an individual with FCD2a. On the basis of the available data, we cannot definitively assess whether these variants are benign or pathogenic, thus expanding the phenotypic spectrum associated with these genes. Future larger association studies or molecular functional investigations will have to be conducted to confirm our findings.

It is important to note that our exome-wide SNV and genome-wide CNV screening design casts a wide net to elucidate the genomic architecture of the lesional epilepsies and uncover potential disease-associated genes that are typically not on hypothesis-based gene panels. However, despite a high coverage of 364x, well within the standard for recent high-impact deep-coverage
our approach is not well-powered to reliably identify somatic variants carried only by a very small number of cells within a lesion (VAF < 2%). Our study was also limited in distinguishing between somatic VAF and germline variants because of the lack of matched blood samples. Therefore, we implemented upper and lower thresholds for the allelic fraction of somatic variants that may have reduced the total number of identified somatic variants in all downstream analyses. Our variant QC using individuals with non-lesional epilepsies as the PoN may have led to the exclusion of somatic variants associated with epilepsy only. It is unlikely that the individuals with non-lesional epilepsy carried genetic factors involved in over/abnormal growth disorders (i.e. tumours and FCDs). However, we cannot rule out the possibility that our strategy may have excluded some somatic variants involved in HS. Overall, our quality filtering strategy was stringent to overcome the direct need for confirmatory sequencing analyses, but will have likely reduced the number of identified somatic variants. As such, the genetic-positive rate is likely higher than detected in our study and other studies with a diagnostic focus that include ultra-deep-coverage sequencing and secondary validation, such as amplicon sequencing or digital PCR, should be cited for diagnostic yield estimates.

Understanding the exact molecular mechanisms involved in the aetiology of epileptogenic pathologies with or without tumour activity is essential for improving treatment of drug-resistant focal epilepsy. Our study systematically shed light on the genomic landscape of the lesional epilepsies, identified four novel candidate genes for lesional epilepsy (i.e. NRAS, KRAS, NFI and PTPN11), and observed several potentially pathogenic somatic CNV. More basic and clinical-genetic research is needed to elucidate the potential suitability of post-surgery screenings after somatic variants to inform surgery outcome prediction, patient management and our general understanding of the aetiology of the various pathologies underlining focal epilepsy.

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Competing interests
The authors report no competing interests.

Supplementary material
Supplementary material is available at Brain online.

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