Heritability of Magnetoencephalography Phenotypes Among Patients With Genetic Generalized Epilepsy and Their Siblings

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**Abbreviations**

AE absence epilepsy
AED antiepileptic drugs
CAE childhood absence epilepsy
FWE familywise error correction
GGE genetic generalized epilepsy
GSWD generalized spike-wave discharges
GTCS generalized tonic clonic seizures only
ICC intraclass correlation
JAE juvenile absence epilepsy
JME juvenile myoclonic epilepsy
MEG magnetoencephalography
SNR signal-to-noise ratio
Abstract

Objective
To assess whether neuronal signals in patients with genetic generalized epilepsy (GGE) are heritable, we examined magnetoencephalography (MEG) resting-state recordings in patients and their healthy siblings.

Methods
In a prospective, cross-sectional design, we investigated source-reconstructed power and functional connectivity in patients, siblings and controls. We analyzed 5 minutes of cleaned and awake data without epileptiform discharges in six frequency bands (1-40 Hz). We further calculated intraclass correlations (ICC) to estimate heritability for the imaging patterns within families.

Results
Compared with controls (n = 45), patients with GGE (n = 25) showed widespread increased functional connectivity (theta to gamma frequency bands) and power (delta to gamma frequency bands) across the spectrum. Siblings (n = 18) fell between the levels of patients and controls. Heritability of the imaging metrics was observed in regions, where patients strongly differed from controls, mainly in beta frequencies, but also for delta and theta power. Network connectivity in GGE was heritable in frontal, central and inferior parietal brain areas and power in central, temporo-parietal, and subcortical structures. Presence of generalized spike-wave activity during recordings and medication were associated with the network patterns, whereas other clinical factors such as age of onset, disease duration or seizure control were not.
Conclusion

Metrics of brain oscillations are well suited to characterize GGE and likely relate to genetic factors rather than the active disease or treatment. High power and connectivity levels cosegregated in patients with GGE and healthy siblings, predominantly in the beta band, representing an endophenotype of GGE.
Introduction

Idiopathic/genetic generalized epilepsy (GGE) is a common epilepsy syndrome accounting for 15-20% of all epilepsies. Different seizure types can occur, ranging from absences, myoclonic and generalized tonic-clonic seizures. For GGE, a polygenic background is presumed. So far, gene discovery has been scarce despite high heritability and large-scale collaborative efforts. Thus, it is of high interest to seek for subclinical traits of the syndrome (endophenotypes), which reflect the genetic background of the disease and cosegregate in families with affected individuals. Candidate markers for GGE have been proposed, such as cognitive functioning, frontal lobe and hippocampal morphology, hippocampal function, and functional network topology. Further, juvenile myoclonic epilepsy (JME) patients and their siblings have shown increased activations of the motor system during cognitive tasks. However, increased brain connectivity and power has also been found in absence of cognitive load in a mixed GGE cohort and in widespread regions. It is less clear to which extent observed findings merely reflect disease activity, effects of seizure burden or treatment. Also, various methodological approaches hinder reproducibility and comparability of functional network studies. Given that characteristics of spontaneous brain oscillations at rest are heritable, the studies of unaffected siblings may help to disentangle genetic factors from secondary disease effects.

This study set out to assess whether imaging metrics based on oscillatory neural activity and measured by magnetoencephalography (MEG) during resting-state could represent an endophenotype of GGE.
Methods

Standard protocol approvals, registrations, and patient consents
This study was approved by the local Ethics Committee of the Medical Faculty of the University of Tübingen and conducted in compliance with the principles of the Declaration of Helsinki. All participants provided written informed consent.

Recruitment
Patients with GGE and their siblings were consecutively recruited through the clinical database of the Department of Neurology, University Hospital of Tübingen, between 2013 and 2019. Advertisement and recruitment of controls was conducted in the local area. All patients were diagnosed with GGE according to the recent International League Against Epilepsy classification\(^2\). At the time of the study, siblings and controls never had experienced seizures, were free of any neurological and psychiatric diseases and did not take any medication.

MEG recording
Resting-state data were measured in supine position (275 channels system, CTF Inc., Vancouver, Canada) in the MEG center of the University of Tübingen (585.9Hz sampling rate). Participants underwent 30 minutes of continuous recording in order to have sufficient data after exclusion of segments with generalized spike-wave discharges (GSWD). All participants were instructed to relax, keep their eyes closed, not to fall asleep and not to think of anything in particular.
Individual head anatomy

A sagittal high-resolution T1-weighted image was acquired for all participants (3D-MPRAGE, repetition time = 2.3 s, echo time = 3.03 ms, flip angle = 8°, voxel size = 1 x 1 x 1 mm), either on a Siemens Magnetom Trio 3T scanner (Siemens AG, Erlangen, Germany) equipped with a 12-channel head coil (11/45 controls, 5/25 patients) or at the Siemens Magnetom prisma 3T system (Siemens, AG, Erlangen, Germany) with a 64-channel head coil (34/45 controls, 18/18 siblings, 20/25 patients). Detailed description of processing methods and references can be found elsewhere. In brief, individual cortical surfaces were reconstructed using FreeSurfer (https://surfer.nmr.mgh.harvard.edu/) and further subjected to SUMA (https://afni.nimh.nih.gov/download/). SUMA decimated each participant’s cortical surface to 1002 common vertices per hemisphere. The surface was resampled using the ‘fsaverage’ template (FreeSurfer) and SUMA (ld = 10). Additionally, six subcortical nuclei (bilateral amygdala, hippocampus, thalamus, caudate, putamen and pallidum) were reconstructed based on the ‘fsaverage’ template. Each region was converted to surfaces and spatially normalized to MNI space (DARTEL; SPM12; https://www.fil.ion.ucl.ac.uk/spm/software/spm12/) using CAT12 DARTEL template (http://www.neuro.uni-jena.de/cat/). Thus, this procedure eventually yielded 2338 vertices for each participant and point-for-point anatomical correspondence for cortical and subcortical regions. Finally, the individual cortical mesh was realigned to the CTF sensor space using the fiducial positions recorded during the MEG session. A volume conduction head model was constructed for the MEG source analysis using the ‘single shell’ method implemented in Fieldtrip.
MEG data processing and source analysis

Preprocessing and further analysis steps were performed using Fieldtrip (https://www.fieldtriptoolbox.org/) running in Matlab (version 9.0, R2016a, Mathworks Inc.) as described and referenced elsewhere\(^{20}\). In short, data were preprocessed (Butterworth band-pass filter 1-70 Hz, line-noise removal), downsampled (150 Hz) and cut into epochs of 10 s length. Trials with GSWD were manually marked and excluded from the further analysis including one trial preceding and one trial after the event (±10 seconds). Each trial was visually inspected and trials with artefacts were manually removed (e.g. movements, excessive muscle activity, sensor jumps). We used independent component analysis (ICA) to detect and manually reject cardiac and eye-movement artefacts. All trials were again reviewed and vigilance was rated according to sleep scoring criteria of the American Academy of Sleep Medicine (AASM). 30 trials of cleaned and awake data (300 seconds) per participant were randomly selected for source analysis. We performed spectral analysis on the MEG sensor data using a multitaper fast Fourier time-frequency transformation approach with frequency-dependent Discrete Prolate Spheroidal Sequences (DPSS) tapers for six frequency bands (delta: 2 ± 2 Hz, theta: 6 ± 2Hz, alpha 10 ± 2 Hz, beta1 16 ± 4 Hz, beta2 25 ± 4 Hz and gamma 40 ± 8 Hz). Power and the cross-spectral density were derived from the Fourier transformed sensor-level data. We used beamforming (Dynamic Imaging of Coherent Sources, DICS) to project the data to the source space. For each vertex point of the individual cortical mesh, the lead field matrix was calculated and an adaptive spatial filter was separately applied for each frequency band (regularization: lambda = 5%). Power was computed for each source position. The coherency coefficient, which quantifies phase synchrony between two signals, was estimated between all pairs of sources (n = 2338). We then investigated the absolute imaginary part of coherency to reduce contributions to the connectivity estimate, which are due to potential field spread\(^{21}\). In sum, a symmetrical, weighted and undirected individual functional connectivity matrix was constructed for each
frequency band. We averaged the weights of each vertex to estimate the overall connection strength of a vertex. To get an overall indicator of metrics, we also averaged connectivity and power across all vertices, yielding one global value per participant.

**Statistical analysis of imaging metrics**

Group differences in power and connectivity were assessed using Permutation Analysis of Linear Models (https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/PALM), a nonparametric statistical tool. In order to allow permutation inference in presence of possible dependence structures among related participants, we applied multi-level block permutation. Blocks of exchangeable units were defined and shuffled as a whole (families and single unrelated participants). Observations within a block were rearranged among themselves (observations within families). We carried out single $t$-contrasts instead of an overall $F$-test, which would have only allowed limited exchangeability of the data given related family structures (patients/siblings) and unrelated controls. Based on previous work, we hypothesized increased network levels in patients with GGE compared with controls and thus, ran one-sided comparisons (controls < siblings, siblings < patients, controls < patients). Groups were contrasted vertex-based and on a global level, respectively, and for each frequency band separately. For each comparison, a general linear model was fit for every permutation, with imaging metrics as dependent variables. Group association and age constituted the predictors. Sex was initially included as additional predictor in the model but did not change any of the main results and was not further considered in the analyses. The data was permuted 5000 times. An estimate of the empirical distribution of the $t$-statistics under the null hypothesis was constructed, from which the $p$-values were generated. In the vertex-based analysis, we corrected for multiple comparisons on cluster level using threshold-free cluster enhancement (TFCE). $P$-values were familywise error (FWE)-corrected within each group contrast and indicated as $-\log_{10} p$ with a significance threshold of 1.3 ($= p < 0.05$). Effect sizes (Cohen’s
$d$) for vertex-based and global group comparisons were derived from the $t$-values of the linear models. $d$ is therefore adjusted for age effects. An effect size of $d = 0.2$ is considered to be small, $d = 0.5$ intermediate and $d = 0.8$ large\textsuperscript{24}.

**Heritability of imaging patterns**

We explored the extent to which imaging phenotypes are heritable and quantified this using intraclass correlation (ICC) through linear mixed-effects modelling\textsuperscript{25}. ICC values were estimated based on the random effect components of a mixed model, which allows the incorporation of confounding effects. Here, a mixed model was constructed for power and connectivity, respectively, as dependent variables, family membership as random effect alongside group (patients versus siblings) and age as subject-level covariates (fixed-effects).

ICC(1,1) was computed based on the variances of the random effect (family) and the total random effect variance (family and residual variance)\textsuperscript{25} using R (nlme package; https://CRAN.R-project.org/package=nlme) and restricted maximum-likelihood estimation (REML). ICC ranges from 0 to 1, where an ICC close to 1 indicates correlated connectivity and power levels for patients-siblings pairs in a family. A low ICC means that family affiliation is not relevant and thus, genetic contribution unlikely. In total, 14 GGE-families contributed to the ICC estimations. We performed a regional resampling of the vertex-level metrics using the Desikan-Killiany atlas\textsuperscript{26} to improve the signal-to-noise ratio (SNR) and calculated ICCs for 80 anatomically defined cortical and subcortical regions. Finally, we investigated whether heritability estimates of imaging metrics are particularly high in brain areas, where patients show stronger differences from controls. To this end, effect sizes (Cohen’s $d$) from the vertex-wise group comparisons were averaged for each anatomical region and related to the ICC maps using Spearman’s rank correlation for each frequency band and metric. Higher positive correlations of effect sizes and ICC-values imply genetic contribution to disease-related patterns in GGE.
Data availability

All relevant data including power and connectivity results and ICC estimations are available from the corresponding author upon request. Raw imaging data is not publicly available due to data protection regulations.
Results

Participants

28 patients with GGE, 21 siblings and 50 controls underwent resting-state measurements. We excluded participants due to technical problems during the acquisition (n = 5), movement artifacts (n = 4) or sleep (n = 2), leaving datasets of 25 patients, 18 siblings (related to 15 patients) and 45 controls for further analysis. Raw data from six patients were also used in a previous study. Anatomical MRI scans were visually rated as normal in all controls, siblings and in 22 patients. Three patients had unspecific findings (two uncomplicated cysts, single unspecific white matter lesion). Demographics and clinical details are described in table 1. Family membership and GGE syndromes are indicated in figure 3A and figure 4A. The groups were comparable for age (analysis of variance, \( p = 0.95 \)) and sex (chi-square, \( p = 0.84 \)).

Connectivity analysis

Compared with controls, patients with GGE showed increased functional connectivity in most of the frequency bands studied. Global connectivity (figure 1A) was higher in the theta (\( t_{(67)} = 2.17, p = 0.011, d = 0.54 \)), alpha (\( t_{(67)} = 2.36, p = 0.016, d = 0.59 \)), beta1 (\( t_{(67)} = 3.35, p = 0.0004, d = 0.84 \)), beta2 (\( t_{(67)} = 2.40, p = 0.023, d = 0.60 \)), and gamma band (\( t_{(67)} = 3.17, p = 0.008, d = 0.79 \)), but not in delta (\( t_{(67)} = 0.88, p = 0.152, d = 0.22 \)). Vertex-based comparisons showed widespread bilateral increases across the frequency spectrum (figure 1B). Strongest effects were observed in the beta1 frequency band with a focus on left-hemispheric temporal, frontal, central and parietal regions. Mesio-frontal regions were also pronounced in alpha, beta2 and gamma frequency bands and postcentral regions mainly in the theta band. Connectivity of siblings statistically fell between patients and healthy controls (figure 1A). Global connectivity of patients was higher than in siblings for alpha (\( t_{(40)} = 1.67, p = 0.047, d \)
but there were no significant differences in the remaining frequency bands (delta: $t_{(40)} = 1.38$, $p = 0.098$, $d = 0.43$; theta: $t_{(40)} = 1.19$, $p = 0.247$, $d = 0.37$; beta1: $t_{(40)} = 0.90$, $p = 0.205$, $d = 0.28$). Siblings did not significantly differ from controls (delta: $t_{(60)} = -0.8$, $p = 0.444$, $d = -0.21$; theta: $t_{(60)} = 0.49$, $p = 0.223$, $d = 0.14$; alpha: $t_{(60)} = 0.45$, $p = 0.267$, $d = 0.13$; beta1: $t_{(60)} = 1.59$, $p = 0.101$, $d = 0.44$, beta2: $t_{(60)} = 0.46$, $p = 0.250$, $d = 0.13$; gamma: $t_{(60)} = 0.749$, $p = 0.520$, $d = 0.21$). On a vertex level (figure 1C) and after correction for multiple comparisons, siblings differed from patients with GGE in alpha and beta2 frequency bands ($p_{FWE} < 0.05$), but not from controls ($p_{FWE} > 0.05$).

Power analysis

Patients with GGE had higher power than controls in all frequency bands studied, in the global (figure 2A, delta: $t_{(67)} = 3.15$, $p = 0.002$, $d = 0.79$; theta: $t_{(67)} = 3.73$, $p = 0.0002$, $d = 0.93$; alpha: $t_{(67)} = 4.22$, $p = 0.0002$, $d = 1.05$; beta1: $t_{(67)} = 5.44$, $p = 0.0002$, $d = 1.36$, beta2: $t_{(67)} = 4.24$, $p = 0.0002$, $d = 1.06$; gamma: $t_{(67)} = 3.60$, $p = 0.0006$, $d = 0.90$) and vertex-based analysis ($p_{FWE} < 0.05$, figure 2B). Differences were focused on occipital-parietal and temporal regions. Also, hippocampal (alpha, beta1, beta2) and subcortical structures such as thalamus and putamen (beta1) showed higher power. Patients also exhibited higher global power than siblings (figure 2A) in alpha ($t_{(40)} = 2.40$, $p = 0.027$, $d = 0.74$), beta1 ($t_{(40)} = 1.34$, $p = 0.024$, $d = 0.41$) and gamma ($t_{(40)} = 1.74$, $p = 0.048$, $d = 0.54$), but not in delta ($t_{(40)} = 1.96$, $p = 0.123$, $d = 0.61$), theta ($t_{(40)} = 1.60$, $p = 0.189$, $d = 0.50$), beta2 ($t_{(40)} = 0.68$, $p = 0.096$, $d = 0.21$) frequency bands. In the vertex-based comparison, patients with GGE had higher power than siblings in all frequency bands but theta (figure 2C), mainly in occipital regions. Global power in siblings was higher than in controls (figure 2A) in beta1 ($t_{(60)} = 2.53$, $p = 0.044$, $d = 0.71$) and beta2 bands ($t_{(60)} = 2.62$, $p = 0.044$, $d = 0.73$), but not in the remaining frequency bands (delta: $t_{(60)} = 0.16$, $p = 0.191$, $d = 0.04$; theta: $t_{(60)} = 1.28$, $p =
0.108, \( d = 0.36; \) alpha: \( t_{(60)} = 0.85, p = 0.246, d = 0.24; \) gamma: \( t_{(60)} = 1.02, p = 0.193, d = 0.29). \) Differences at vertices did not reach statistical significance \( (p_{\text{FWE}} > 0.05). \)

**Heritability of imaging patterns**

ICC values, which represent heritability estimations of MEG-derived patterns, strongly correlated with beta1 connectivity differences of patients against controls \( (r_s = 0.59, p = 1.88 \text{e}^{-08}) \) and negatively to beta2 levels \( (r_s = -0.39, p = 4.23 \text{e}^{-04}) \). The correlation of ICC and connectivity differences in the remaining frequency bands did not reach significance \( (\text{delta}: r_s = -0.01, p = 0.900; \text{theta}: r_s = -0.10, p = 0.364; \text{alpha}: r_s = 0.14, p = 0.223; \text{gamma}: r_s = 0.18, p = 0.120). \) ICC values for power were significantly associated with GGE contrast maps in all frequency bands except alpha and gamma \( (\text{delta}: r_s = 0.60, p = 8.6 \text{e}^{-09}; \text{theta}: r_s = 0.46, p = 2.5 \text{e}^{-05}; \text{alpha}: r_s = 0.18, p = 0.109; \text{beta1}: r_s = 0.59, p = 1.47 \text{e}^{-08}; \text{beta2}: r_s = 0.34, p = 0.002; \text{gamma}: r_s = 0.15 p = 0.17). \) Below, we only report results for ICC maps with positive and significant correlations with the GGE phenotype \( (p < 0.05), \) implying genetic contribution to disease-related patterns \( (\text{figure 3B and figure 4B}). \) ICC values for connectivity in beta1 frequency band were highest in rostral and caudal anterior cingulate, orbitofrontal, paracentral, entorhinal and inferior parietal regions \( (ICC > 0.4) \) and lower in temporal regions and subcortical nuclei \( (ICC > 0.2) \) \( (\text{figure 3B}). \) ICC estimates were generally higher for power than connectivity and peaked in beta frequency bands in temporal, subcortical and parietal regions such as lingual gyrus and cuneus as well as postcentral gyrus \( (ICC > 0.5). \) ICC maps for delta and theta frequency bands showed similar patterns but generally lower ICC values \( (\text{ICCs up to } \sim0.5) \) \( (\text{figure 4B}). \)

**Clinical variables and imaging findings**

We further carried out secondary analyses to evaluate the relation of clinical variables with brain oscillations. We investigated whether networks of GGE patients with GSWD during the
MEG recordings differ from patients without GSWD. Although trials containing GSWD (±10 seconds of data) were rejected in all analyses, patients with GSWD in the recording had higher connectivity in the delta frequency band (global: $t_{(22)} = 2.95$, $p = 0.004$, $d = 1.26$; vertex-based: $p < 0.05$, **figure 5A**). There was also a tendency of higher power across frequency bands in those patients, with significant differences for delta (global: $t_{(22)} = 2.55$, $p = 0.010$, $d = 1.09$; vertex-based: $p < 0.05$) and beta1 frequency bands (global: $t_{(22)} = 2.23$, $p = 0.019$, $d = 0.96$, vertex-based: $p < 0.05$) (**figure 5A**). Mean global power and connectivity for patients without GSWD during the recording remained higher for patients than controls and never dropped below the mean levels of siblings (data not shown). Patients taking two or more drugs at the study date had lower connectivity in beta1 frequency band than patients taking less than two drugs (global: $t_{(21)} = -2.04$, $p = 0.029$, $d = -0.98$; vertex-based: $p < 0.05$, **figure 5C**). The effect was stronger when accounting for the presence of GSWD during MEG recordings. There were no differences for power in any of the frequency band studied (all $p > 0.05$). Other clinical factors such as age of onset, disease duration or seizure control, were not associated with global and vertex-based connectivity or power (all $p > 0.05$).
Discussion

We assessed power and phase-based connectivity during rest in patients with GGE and their healthy siblings to evaluate endophenotypic potential of electrophysiological metrics. Patients with GGE showed bilateral, widespread and highly increased power and connectivity compared with controls. Asymptomatic siblings presented with intermediate levels between patients and controls, particularly in beta frequencies, suggesting genetic background as major driver for those patterns.

We expanded and replicated previous work of ours using MEG data of mixed GGE cohorts15, 16, confirming strong interictal network differences in GGE as quantified with the investigated imaging metrics. Both, global and local measures, have been reliable in source-space analyses using the same processing pipeline20. Increased power in GGE is a previously reported finding27, 28, but M/EEG connectivity studies on source level are scarce15, 16. High beta band connectivity in GGE was the most consistent finding across the studies, followed by increases in theta, alpha and gamma bands. Epileptic seizures commonly involve pathological synchronization29. Also during the interictal state, patients have had higher liability to synchronize30, 31.

Increased network connectivity and power was also observed in siblings without active epilepsy6. Intraclass correlations substantiate heritability in regions, where patients with GGE strongly differed from controls, specifically for beta1 band connectivity and both, beta1 and beta2 power. In delta and theta power, heritability estimates corresponded to the phenotype of patients, but effect sizes and ICC values were comparably low. Increased beta2 connectivity patterns in patients compared with controls were less concordant within families.
Elevated MEG connectivity was mostly heritable in anterior cingulum, orbito- and superior frontal regions. In GGE, direct evidence for a genetically determined functional dysregulation in frontal cortex is limited, but has been documented for prefrontal and cingulate morphology. Altered structural network integration of the frontal cortex potentially leads to cognitive impairments in JME patients and similarly, in their healthy siblings. Furthermore, network power and connectivity was heritable in central brain regions. Hyperactivations in motor systems has been suggested as endophenotypes of JME during cognitive fMRI, but also for a mixed GGE cohort during resting-state. We complement these findings by studying much faster neuronal oscillations using MEG and without vascular confounds. Our results point to more globally increased network levels in GGE and siblings, irrespective of a task involved. This includes increased power patterns in mesio-temporal cortices and in subcortical structures with a strong genetic substrate. ICC estimates reached levels of up to 0.85, particularly in beta frequencies, indicating that familial and, thus, likely genetic factors explain the majority of the observed variance. Thalamocortical circuits are critical for GGE and structural alterations in the thalamus have been linked to subcortico-cortico hub organization in GGE. Using MEG, it is debatable whether signals of deep brain structures can be captured. Yet, recent work has used simultaneous intracerebral and MEG recordings in epilepsy patients and demonstrated detectability of signals generated in mesial temporal lobe structures as well as thalamic activity at the surface. In line with our results, mesio-temporal task-based activations co-segregated in JME patients and their healthy siblings, along with changes in hippocampal morphology. The role of increased occipital power in our GGE cohort is less clear. Low-density EEG-data and microstructural alterations in fronto-occipital white matter association tracts point to the involvement of occipital areas in GGE. In our study, patients and siblings mostly differed in occipital power, suggesting that this finding may not be due to genetic factors but to other yet unknown disease-related effects. Overall, we consolidate findings of earlier GGE-siblings studies using...
(f)MRI and add evidence for resting-state trait heritability in extended brain networks at higher temporal resolution.

Electrophysiological studies in twins suggest strong genetic influence on brain oscillations relative to environmental factors, particularly for wideband power ($h^2 \approx 0.5-0.8$)\textsuperscript{18, 37}. Heritability for connectivity on the cortical source-level has been lower than for power and highest in alpha and beta frequencies (10-20%)\textsuperscript{19}. Lower reproducibility of connectivity metrics may, at least partly, explain a lower heritability estimate compared to power. Similarly, connectivity in alpha and beta bands have been more repeatable than in other frequency bands.\textsuperscript{20} Network alterations characterizing GGE were captured in brain rhythms of different frequencies, but imaging phenotypes of siblings and patients with GGE most strongly correlated in beta frequencies. Beta band oscillations are classically related to sensory and motor processing\textsuperscript{38} as well as long-distance synchronization\textsuperscript{39}. Interestingly, GABAergic processes are presumably involved in the generation of MEG beta oscillations as shown with endogenous GABA-concentrations in humans\textsuperscript{40}. GABAA receptor gene variants constitute GGE disease risk\textsuperscript{5, 41} and one of those candidates, the $GABA\text{R}2$ gene, has been linked to EEG beta band activity\textsuperscript{42, 43}. Moreover, recently discovered genetic markers in GGE have been enriched in the frontal cortex, specifically in the dorso-lateral prefrontal cortex\textsuperscript{5}. Genetic signals have further converged on the inferior temporal lobe, angular gyrus, cingulum and subcortical tissue, but less strongly\textsuperscript{5}. Our ICC-maps show substantial spatial correspondence to those findings, emphasizing the functional relevance of imaging resting-state markers as investigated in our study. Previously, oscillatory activity has been successfully utilized to detect psychiatric liability genes\textsuperscript{43} and EEG coherence has served as an endophenotype for alcohol use disorders\textsuperscript{44}. Yet, the functional role of spectral perturbations is not fully understood. Here, we only can speculate about molecular changes occurring within specific networks in the brain, such as an excitation-inhibition imbalance as
putative key factor in epilepsy. Through coupling mechanisms, the coordination of neuronal spike timing might be affected across networks in sum leading to increased oscillatory amplitudes.\textsuperscript{45}

Disease duration and age of onset did not correlate with the imaging patterns in our GGE cohort. These clinical variables do not necessarily describe disease severity and the lack of a significant association with the imaging findings may support the notion of a genetic imaging trait. Patients showing epileptic discharges during the MEG recording had higher delta connectivity and delta and beta power increases at rest (after careful exclusion of segments with GSWD $\pm$ 10 seconds of data). These patterns had a spatial profile with a temporal and central focus. GSWD typically have a frequency around 3 Hz and it is possible that we captured network dynamics around the on- or offset, evolving during a considerable time span\textsuperscript{31}. Only nine patients in our study showed GSWD and at least five of them had experienced seizures within the past year. However, seizure control was not associated with significantly different network patterns in our cohort and might not be a sensitive marker in a rather well controlled cohort. Similarly, persistent GSWD do not necessarily have an effect on long-term seizure prognosis\textsuperscript{46}. Moreover, patients with higher medication load had lower connectivity in the beta\textsubscript{1} frequency band. Since only six patients received more than two antiepileptic agents at the time of the study, we can only speculate about a network down-regulation through antiepileptic treatment. Normalizing effects of antiepileptic drugs on background synchronization has been demonstrated before\textsuperscript{47}. Further investigations are needed to confirm our exploratory results and study drug-specific effects.

This study has limitations. We found alterations for both, power and imaginary part of coherency, a phase-based metric. Power and phase characterize different aspects of neural signals, however, both, physiological and non-physiological coupling between those
characteristics has been noted\textsuperscript{45}. In particular, high values in phase-based connectivity require a temporally stable phase relationship of signals, which depends on the SNR and eventually, on signal power. In our study, the topography for GGE power differences had a posterior focus and was distinct from connectivity maps with a more frontal emphasis. Recent work suggests that spontaneous MEG networks can be decoupled into anterior and posterior states, both connected to the posterior cingulate cortex, likely reflecting functional specialization\textsuperscript{48}. Given the different effects of power and connectivity on the MEG-signal topography, it is very likely that both measures captured independent features. Furthermore, we cannot assess the mechanisms of genetic control over oscillatory markers. Methodological aspects may also affect heritability estimates. For example, higher sensitivity of connectivity measures to noise might be a reason for generally lower effect sizes than for power and weaker ICC-values in our study. Accordingly, higher statistical power is needed for significant connectivity differences between siblings and controls or patients, given the intermediate levels of the siblings. The relatively small number of 14 families that were available for the study also limits the precise estimation of the ICCs. Finally, we cannot discern specific effects for the GGE subsyndromes due to the small sample size. Yet, multiple GGE subtypes occur within the same families\textsuperscript{49} and overlap of genetic risk factors has been suggested\textsuperscript{5}. Network phenotypes as assessed in our study could reflect shared pathophysiological features across the syndromes, such as the occurrence of GSWD. Interestingly, GSWD have been more frequently observed in unaffected first-degree relatives of patients with GGE than in the general population\textsuperscript{50}, which again points to genetic contributions to network function in GGE.

We propose that increased interictal MEG power and connectivity in fronto-central and temporo-parietal cortical regions are a hallmark of GGE. These network features are likely driven by genetic factors and not by the presence or absence of the active disease or clinical
confounds. Siblings without epilepsy had similarly increased network levels during rest, predominantly in beta frequencies. We show that power and phase-based connectivity are heritable and may serve as markers to link imaging with genetics in epilepsy.
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Disclosures

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### Appendix 1. Authors

<table>
<thead>
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<th>Location</th>
<th>Contribution</th>
</tr>
</thead>
<tbody>
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<tr>
<td>Name</td>
<td>Affiliation</td>
<td>Role in the study</td>
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<td>Designed and conceptualized the study, project supervision, interpreted the results; revised the manuscript for intellectual content</td>
</tr>
</tbody>
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References


# Tables

## Table 1 Study population

<table>
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<tr>
<th></th>
<th>Patients</th>
<th>Siblings</th>
<th>Controls</th>
</tr>
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<tbody>
<tr>
<td>Total (n)</td>
<td>25</td>
<td>18</td>
<td>45</td>
</tr>
<tr>
<td>Female, n (%)</td>
<td>16 (64)</td>
<td>10 (55)</td>
<td>28 (62)</td>
</tr>
<tr>
<td>Age (y), median (IQR)</td>
<td>25 (22-37)</td>
<td>26 (22-42)</td>
<td>25 (23-35)</td>
</tr>
<tr>
<td>Positive family history of epilepsy / seizures, n (%)</td>
<td>12 (48)</td>
<td>7 (39) &lt;sup&gt;a&lt;/sup&gt;</td>
<td>0 (0)</td>
</tr>
<tr>
<td>GSWD during MEG recordings, n (%)</td>
<td>9 (36)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>GSWD in routine / longterm EEG, n (%)</td>
<td>22 (88)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Seizure free &gt; 12 months, n (%)</td>
<td>16 (64)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Drugs at measurement, mean (range)</td>
<td>1.2 (0-3)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Epilepsy syndrome, n (%)</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>CAE</td>
<td>5 (20)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>JAE</td>
<td>6 (24)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>JME</td>
<td>5 (20)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>GTCS</td>
<td>4 (16)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>GGE</td>
<td>5 (20)</td>
<td>-</td>
<td>-</td>
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<tr>
<td>Age of onset (y), median (IQR)</td>
<td>15 (10-17)</td>
<td>-</td>
<td>-</td>
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<tr>
<td>Disease duration (y), median (IQR)</td>
<td>17 (8-24)</td>
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Abbreviations: CAE = childhood absence epilepsy; GSWD = generalized spike-wave discharges; GTCS = generalized tonic clonic seizures only; GGE = genetic generalized epilepsy (unclassified); IQR = interquartile range; JAE = juvenile absence epilepsy; JME = juvenile myoclonic epilepsy; y = years. <sup>a</sup> Positive family history beyond the related index patient with GGE.
Figures

Figure 1 Group-level connectivity differences

A Violin plots show individual data points, the density of the data, group means and standard errors of the means for the global imaginary part of coherency in each frequency band (controls n = 45, siblings n = 18, patients with GGE n = 25). Asterisks denote statistical significance at *p < 0.05 and **p < 0.001 for permutation-based group comparisons. B The plot highlights vertices with significantly higher connectivity values in patients with GGE (n = 25) than in controls (n = 45) and C higher connectivity values in patients with GGE (n = 25) than in siblings (n = 18). The colour scale indicates –log10 p with a cutoff of 1.3 (corresponding to p < 0.05, FWE corrected). In all analyses, age was included as covariate of no interest.
Figure 2 Group-level power differences

A Violin plots show individual data points, the density of the data, group means and standard errors of the means for global power in each frequency band (controls n = 45, siblings n = 18, GGE patients n = 25). Asterisks denote statistical significance at *p < 0.05 and **p < 0.001 for permutation-based group comparisons. For visualization purposes, power data was log10-transformed. B The plot highlights cortical and below, subcortical vertices with significantly higher power values in GGE patients (n = 25) than in controls (n = 45). C The plot shows vertices with significantly higher connectivity values in GGE patients (n = 25) than in siblings (n = 18). The colour scale indicates -log10 p with a cutoff of 1.3 (corresponding to p < 0.05, FWE corrected). In all analyses, age was included as covariate of no interest.
Figure 3 Connectivity within families related to GGE imaging patterns

A Individual global connectivity values of patients with GGE and siblings are plotted with regard to their family membership (columns of data within frequency bands). Data of GGE patients without a corresponding sibling are not shown. GGE syndromes of patients in each family are indicated on the x-axis. CAE and JAE are referred to absence epilepsies (AE). B shows colour-coded heritability estimates (ICC values) per region based on the Desikan-Kiliany atlas with small to large group-level differences between GGE patients and controls (Cohen’s $d > 0.2$). Only ICC-maps with a positive and significant correlation with averaged effect sizes are shown ($p < 0.05$). The colour coding indicates the strength of ICC values in those regions for connectivity differences. ICC estimates were derived from random effect components of mixed models for each region, taking group and age effects into account. A large ICC indicates correlated imaging patterns for patients-siblings pairs in a family ($n = 14$) and thus heritability of the metrics. Cortical regions are displayed in the left column and subcortical regions separately shown in the right column of the plot.
Figure 4 Power within families related to GGE imaging patterns

A Individual global power values of patients with GGE and siblings are plotted with regard to their family membership (columns of data within frequency bands). Data of GGE patients without a corresponding sibling are not shown. GGE syndromes of patients in each family are indicated on the x-axis. CAE and JAE are referred to absence epilepsies (AE). For visualization purposes, power data was log10-transformed. B shows colour-coded heritability estimates (ICC values) per region based on the Desikan-Killiany atlas\textsuperscript{26} with small to large group-level differences between GGE patients and controls (Cohen’s $d > 0.2$). Only ICC-maps with a positive and significant correlation with averaged effect sizes are shown ($p < 0.05$). The colour coding indicates the strength of ICC values in those regions for power differences. ICC estimates were derived from random effect components of mixed models for each region, taking group and age effects into account. A large ICC indicates correlated imaging patterns for patients-siblings pairs in a family ($n = 14$) and thus heritability of the metrics. Cortical regions are displayed in the left column and subcortical regions separately shown in the right column of the plot.
Figure 5 Clinical variables and imaging metrics

Vertex-plots highlight cortical vertices with higher A connectivity and B power values in GGE patients with generalized spike-wave discharges (GSWD) (n = 9) than patients without GSWD (n = 16) during the MEG recordings. These effects were present after exclusion of trials containing GSWD ±10 seconds of data and corrected for age effects. C The plot shows significantly lower connectivity in GGE patients taking two or more antiepileptic drugs (n = 6) than patients taking less than two drugs (n = 19) at the study date. Age and presence of GSWD was included as covariate of no interest. Colour scales indicate -log10 p with a cutoff of 1.3 (corresponding to p < 0.05, FWE corrected).
Heritability of Magnetoencephalography Phenotypes Among Patients With Genetic Generalized Epilepsy and Their Siblings
Christina Stier, Adham Elshahabi, Yiwen Li Hegner, et al.
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