Noninvasive Mapping of Ripple Onset Predicts Outcome in Epilepsy Surgery

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Objective: Intracranial electroencephalographic (icEEG) studies show that interictal ripples propagate across the brain of children with medically refractory epilepsy (MRE), and the onset of this propagation (ripple onset zone [ROZ]) estimates the epileptogenic zone. It is still unknown whether we can map this propagation noninvasively. The goal of this study is to map ripples (ripple zone [RZ]) and their propagation onset (ROZ) using high-density EEG (HD-EEG) and magnetoencephalography (MEG), and to estimate their prognostic value in pediatric epilepsy surgery.

Methods: We retrospectively analyzed simultaneous HD-EEG and MEG data from 28 children with MRE who underwent icEEG and epilepsy surgery. Using electric and magnetic source imaging, we estimated virtual sensors (VSs) at brain locations that matched the icEEG implantation. We detected ripples on VSs, defined the virtual RZ and virtual ROZ, and estimated their distance from icEEG. We assessed the predictive value of resecting virtual RZ and virtual ROZ for postsurgical outcome. Interictal spike localization on HD-EEG and MEG was also performed and compared with ripples.

Results: We mapped ripple propagation in all patients with HD-EEG and in 27 (96%) patients with MEG. The distance from icEEG did not differ between HD-EEG and MEG when mapping the RZ (26–27mm, p = 0.6) or ROZ (22–24mm, p = 0.4). Resecting the virtual ROZ, but not virtual RZ or the sources of spikes, was associated with good outcome for HD-EEG (p = 0.016) and MEG (p = 0.047).

Interpretation: HD-EEG and MEG can map interictal ripples and their propagation onset (virtual ROZ). Noninvasively mapping the ripple onset may augment epilepsy surgery planning and improve surgical outcome of children with MRE.

Epilepsy surgery is the therapy of choice for children with medically refractory epilepsy (MRE). The goal of epilepsy surgery is to resect the epileptogenic zone (EZ), the brain area indispensable for the generation of seizures.1 However, identifying this zone is challenging.2,3 In ~25% of patients, noninvasive techniques cannot formulate a clear hypothesis regarding the EZ.2,4 For them, an invasive evaluation is recommended, which consists of long-term intracranial electroencephalography (icEEG) with macroelectrodes directly implanted into or onto the brain. The main purpose of icEEG is to record seizures and trace the seizure onset zone (SOZ), the most logical EZ estimator.1 Nonetheless, seizures are unpredictable; their recording can be time-consuming and come at the
expense of substantial resources. The availability of interictal biomarkers that estimate the EZ noninvasively without the need to wait for unpredictable seizures is paramount.

Several studies investigated interictal high-frequency oscillations (HFOs) as an alternative to seizures. HFOs are recorded both invasively with icEEG and noninvasively with EEG and/or magnetoencephalography (MEG). HFOs are classified into ripples (>80Hz) and fast ripples (>250Hz). Fast ripples are closely linked to epileptogenicity but can be too focal to be captured noninvasively. In contrast, ripples can be recorded with conventional icEEG and detected using EEG and MEG, although less frequently. Nonetheless, the presurgical value of ripples has been long debated, because they are seen over large areas (called ripple zones [RZs]), which may also encompass nonepileptogenic regions that generate physiological ripples.

Two research groups independently showed that ripples propagate across icEEG electrodes in patients with MRE, and the onset of this propagation (ripple onset zone [ROZ]) is more predictive of surgical outcome compared to areas of spread. This prompts the value of mapping ripples before surgery and raises the question of whether noninvasive techniques can map this propagation and identify the ROZ. Using MEG and high-density EEG (HD-EEG), virtual sensors (VSs) can be reconstructed at desired brain locations to increase the signal-to-noise ratio (SNR), facilitating the noninvasive identification of ripples. Nonetheless, it remains unknown whether a noninvasive implantation of VSs can capture the propagation phenomenon and define an epilepsy biomarker (the ROZ) that so far is known exclusively on invasive icEEG, which currently constitutes the technique of choice for recording ripples and their propagation. Such a virtual implantation could potentially augment presurgical planning and outcomes of epilepsy surgery.

Here, we aim to map noninvasively the spatiotemporal propagation of interictal ripples across the brain of children with MRE using a virtual implantation and to assess the prognostic value of resecting the noninvasively localized areas that initiate this propagation. We hypothesize that areas that initiate the ripple propagation can be identified noninvasively and their proximity to resection predicts outcome. To test our hypothesis, we reconstructed VSs in children with MRE, detected ripples and their onset on VSs, and compared them with the icEEG-defined RZ and ROZ as well as with resection and outcome. VSs were placed to match the icEEG location to enable direct comparison between the proposed noninvasive approach and the invasive benchmark.

Patients and Methods

Patients
We retrospectively reviewed patients with MRE who underwent epilepsy surgery at Boston Children’s Hospital (BCH) between June 2011 and June 2018. We included patients with (1) preoperative HD-EEG and MEG, (2) long-term icEEG, (3) ≥1-year follow-up, (4) postimplantation computerized tomography (CT), and (5) preoperative and postoperative magnetic resonance imaging (MRI). Patients were excluded if <5 minutes of good-quality high-frequency data (≥80Hz) were available for HD-EEG or MEG. The study protocol received approval by the institutional review board of BCH (IRB-P00022114), which waived the need for informed consent due to the retrospective nature.

Simultaneous HD-EEG/MEG Recordings
HD-EEG/MEG recordings were conducted at the MEG Laboratory of Athinoula Martinos Center for Biomedical Imaging (Charlestown, MA) in a 3-layer magnetically shielded room (Imedco, Hägendorf, Switzerland) with a whole-head 306-channel MEG system (Neuromag VectorView, Elekta, Helsinki, Finland). HD-EEG was recorded with 70-channel electrode caps (EASYCAP, Herrsching, Germany) plus 2 temporal electrodes (T1/T2). More details about the protocol can be found in previous studies. Data were recorded for 10 to 12 sessions (4 minutes each; sampling rate: 600, 1000, or 2000Hz). We analyzed sessions regarded as containing interictal activity by the attending epileptologist regardless of the patient’s vigilance state. Data quality was checked using standard (1–70Hz, 10 s/page) and HFO display settings (80–200Hz, 4 s/page). Segments were regarded as good quality in the high-frequency range when showing regular low-amplitude background with HFO display settings. Given the low SNR of scalp EEG and MEG signals for frequencies greater than 200Hz, no fast ripples were considered, and ripples were limited to frequencies less than 200Hz. Finally, we selected epochs free of artifacts, high-frequency noise, and technical disruptions.

icEEG Recordings
icEEG was recorded with subdural (10mm distance) and/or depth electrodes (3–5mm interdistance; Ad-Tech, Racine, WI) using XLTEK NeuroWorks (Natus, Pleasanton, CA). Per our institution’s clinical practice, pediatric epileptologists reviewed each patient’s icEEG daily and extracted multiple 5- to 10-minute segments containing interictal epileptiform activity. We retrospectively reviewed these segments and
selected 5 to 10 minutes of data. This duration was shown to be robust to possible HFO propagation changes across time. Channels with continuous artifacts were excluded.

The location of icEEG contacts was determined on patients’ presurgical MRI coregistered with postimplantation CT (Fig 1A) using Brainstorm. To account for brain shift that occurs after electrocorticographic implantation, subdural electrodes were projected onto the cortical surface (reconstructed via FreeSurfer). When both subdural and depth electrodes were implanted, depth electrodes were also adjusted to compensate for brain shift.

**VS Reconstruction**

VSs were placed to match the icEEG contacts (see Fig 1B). VS time series were reconstructed for HD-EEG and MEG separately (see Fig 1C).

**Beamformer Analysis.** We extracted cortical surfaces from the preoperative MRIs via Freesurfer and constructed realistic head models using OpenMEEG (3-layer boundary elementary model). Source space included the entire brain volume.
We used linearly constrained minimum variance beamformer, implemented in Brainstorm, to estimate brain activity within the source space. Data covariance was computed from 200-millisecond windows around each spike, filtered between 80 and 200Hz. For noise covariance, we used 10 seconds of broadband data (1–300Hz) without epileptiform activity. A source activation map (covering all volume points) was generated for each time point.

**Virtual Sensors.** Beamformer output was used to reconstruct the activity of selected brain locations (VSs), where the icEEG electrodes had been implanted during the patient’s phase 2 evaluation. For each icEEG electrode, we delineated nonoverlapping regions of interest that included the closest volume points surrounding the electrode’s center, up to 5 or 10mm for depth or subdural electrode, respectively (see Fig 1B). Finally, we reconstructed each VS’s time series by computing its mean activation (mean across volume points) for HD-EEG and MEG separately (see Fig 1C).

**Ripple Detection** Automated ripple detection was performed on icEEG and VSs. Given the low SNR of HD-EEG and MEG signals greater than 200Hz, detection on VSs was performed between 80 and 200Hz using an envelope threshold of 4 standard deviations (SDs), which is lower than on icEEG (5 SDs). The detector identifies as ripples only events showing an “island” in the time–frequency plane (Fig 2). We analyzed ripples independently from other features (eg, overlap with spikes, amplitude, or morphology) that may discriminate pathological from physiological events, because no well-established methods exist to this purpose, particularly for scalp EEG/MEG. In addition, the importance of ripple propagation in MRE is only known from icEEG studies, where all ripples were considered, independently from other features.

We computed ripple rates for each icEEG electrode (ripples/min) and normalized them with respect to their maximum per patient. Similarly, we computed normalized ripple rates for VSs (HD-EEG and MEG separately), which we will refer to as “virtual ripple rate.”

**Spatiotemporal Propagation of Ripples** To characterize the spatiotemporal propagation of ripples and identify the ROZ (see Fig 2), we followed our previously described methodology, which automatically (1) finds propagation sequences, defined as at least 3 temporally overlapping ripples in neighboring contacts (<30mm apart); and (2) detects onset ripples of each sequence (occurring within 10 milliseconds from propagation onset).

For each patient, we computed the rate of onset ripples of each icEEG electrode and normalized it. Similarly, we computed normalized rates of onset ripples per VS (for HD-EEG and MEG separately), which we call "virtual onset ripple rate." The icEEG electrodes with a normalized rate of onset ripples greater than 0.8 defined the intracranial ROZ as the brain tissue within 10mm from their center (Fig 3A, red volume). Similarly, intracranial RZ was defined by icEEG electrodes with a normalized ripple rate greater than 0.8 (independently from propagation analysis).

**Comparison with icEEG** We assessed the ability of VSs to localize RZ or ROZ by comparing the location of VSs showing ripples or onset ripples with the icEEG-defined ripple zones. We quantified the performance of the virtual ripple rate to identify the VSs within the intracranial RZ (see Fig 3A) by estimating the area under the curve (AUC) of the receiver operating characteristic (ROC) curves (built for each patient and modality). Similarly, we built ROC curves on the virtual onset ripple rate to assess its ability to localize the intracranial ROZ. We averaged ROC curves for RZ and ROZ separately and identified the optimal operating point (through Youden index). VSs were regarded as belonging to RZ or ROZ when their rate was above the optimal cutoff. The median distance of these VSs from the icEEG-defined RZ or ROZ (D<sub>icEEG</sub>) was calculated per patient (see Fig 3A) separately for HD-EEG and MEG.

**Resection and Postsurgical Outcome** We coregistered preoperative and postoperative MRIs using Brainstorm and defined the resection volume (see Fig 3B, C). For each VS, we calculated their distance from resection (D<sub>RES</sub>) as the Euclidean distance of their center from the closest resection margin (see Fig 3B, C). VSs were considered resected when D<sub>RES</sub> was ≤10mm. Considering a mean gyral width of 11 to 21mm, we defined concordance, across the whole study, as a 10mm distance or less (which can be interpreted as a measure of gyral-width concordance).

Postsurgical outcome was evaluated using Engel classification based on the most recent follow-up at least 12 months after surgery and dichotomized into good (Engel IA–D) and poor (Engel class ≥ II).

**Outcome Prediction** To test whether the virtual mapping of RZ and ROZ helps surgical planning, we evaluated whether this could classify the tissue around each VS as epileptogenic or not.
FIGURE 2: Ripple propagation on virtual sensors (VSs). Examples are shown of ripple propagation on magnetoencephalography (MEG) VSs (top) and electroencephalographic (EEG) VSs (bottom). Each scenario shows the ripple propagation in (1) the time domain (left), where ripples are seen on adjacent VSs with a certain temporal latency from the onset (red dashed line; VSs showing onset ripples, i.e., within 10 milliseconds from the onset, are highlighted in blue); (2) the time–frequency domain (middle), where ripples are seen as an island in the spectral content within the ripple frequency band (80–200Hz); and (3) the spatiotemporal domain (right), where the VSs involved in the propagation are displayed on the patient’s magnetic resonance imaging and color coded by their temporal latency from the onset. F = female; M = male; yo = years old.
For each patient, we defined virtual RZ or virtual ROZ by identifying the VS(s) with high rates of ripples or onset ripples. The optimal threshold to identify these VSs was not known a priori. Thus, we varied the threshold from 0 to 100% of the patient’s maximum rate (5% steps). The resection percentage of the obtained virtual zone was computed as the percentage of its VSs that were resected. We considered a zone to be resected when most of it was surgically removed (resection percentage > 50%). Good outcome following resection was regarded as true positive (TP) prediction and poor outcome following missed resection as true negative (TN) prediction. After estimating positive predictive value (PPV = TP/[TP + false positive (FP)]), negative predictive value (NPV = TN/[TN + false negative (FN)]), and prediction accuracy ([TP + TN]/[TP + TN + FP + FN]) for all thresholds, virtual RZ and virtual ROZ were defined as those providing the highest predictive performance. Median DRES of virtual RZ and virtual ROZ were computed per patient.

**Spike Localization and SOZ**

We compared virtual ripple zones with the localization of spikes (which are the standard approach in HD-EEG/MEG source localization for epilepsy) and the icEEG-defined SOZ.

Spike localization was performed by an experienced reader (C.P.) on HD-EEG and MEG independently. Each individual spike was localized using an equivalent current dipole (as is common practice for HD-EEG/MEG at our institution) following the same methodology described in our previous study. Spike average was not applied to avoid merging discharges with similar scalp topography generated by different sources. Dipoles with a goodness of fit less than 60% were discarded.

For each dipole, we estimated DRES as well as distance from virtual RZ and virtual ROZ (as distance from the center of their closest VS), regarding them as overlapping when closer than 10mm. For each patient, we estimated the overall overlap (as percentage) with virtual
RZ, virtual ROZ, and resection. Finally, we automatically identified dipole clusters as groups of at least 5 dipoles within 10mm distance and identified the anatomical location (frontal, central, parietal, temporal, occipital) of the preponderant cluster.

SOZ was defined by the contact(s) showing the earliest change associated with clinical seizures during icEEG monitoring (independently from this study). We calculated the distance of each VS from SOZ (D_{SOZ}), regarding them as overlapping when \( \leq 10 \) mm apart.

**Statistical Analysis**

Wilcoxon signed rank test was used for paired comparisons between modalities and Wilcoxon rank sum for nonpaired comparisons between outcome groups. Nonparametric tests were used because variables were not normally distributed or variables were of skewed distribution.

Wilcoxon test was used to test association between resection of virtual RZ, virtual ROZ, or dipoles and outcome. Phi coefficient (\( \phi \)) was calculated for both HD-EEG and MEG to identify the earliest change associated with clinical seizures during icEEG monitoring (independently from this study). The distance of each VS from SOZ (D_{SOZ}) was estimated as a measure of association strength.

We considered \( p \leq 0.05 \) to be significant. Results are reported as median (interquartile range). MATLAB R2018a (MathWorks, Natick, MA) was used for statistics.

**Results**

**Patient Cohort**

Twenty-eight children with MRE (age at surgery = 12.5 years) were included. Thirty-six patients met the inclusion criteria; 8 were excluded because of high-frequency noise (>80Hz) on HD-EEG or MEG. Table 1 summarizes patients’ characteristics. For icEEG, we analyzed 5.2 minutes (5.0–6.3) and 100 (87–121) contacts per patient without difference between outcomes (\( p = 0.2 \)). Subdural electrodes were implanted in 11 patients, depth electrodes in 6 patients, and both types in 11 patients, with a total of 72% of contacts in the gray matter. For HD-EEG and MEG, we analyzed an average of 9.6 and 11 minutes of artifact-free data per patient.

Average interval between HD-EEG/MEG was 6 (3–13) months.

Seventeen patients (61%) had good postsurgical outcome. An average of 1.26% (0.9–1.88%) of the brain was resected in our cohort without difference between outcomes (\( p = 0.08 \); see Tables 1 and 2). No differences were seen in sex, age at surgery and epilepsy onset, follow-up period, and pathology between outcomes (see Table 2).

**Comparison with icEEG**

**Ripple Occurrence and Spatiotemporal Propagation.** Ripples on VSs (HD-EEG and MEG) were detected in all patients. Ripple rates were lower on VSs than icEEG (\( p < 0.001 \)), with EEG VSs recording higher rates than MEG VSs (0.9 vs 0.4 ripples/min, \( p < 0.001 \); Table 3). Ripple propagation was observed in all patients using icEEG and EEG VSs, and in 27 patients (96%) using MEG VSs. Table 3 reports the spatiotemporal characteristics of propagation per modality. We detected more ripple propagation on icEEG than EEG VSs and MEG VSs (15 vs 5.4 vs 2.4 propagations/min, \( p < 0.001 \)).

Frequency of icEEG ripples was positively correlated with their amplitude (\( p < 0.001 \), \( R = 0.2 \)); because MEG and HD-EEG are more sensitive to high-amplitude than low-amplitude ripples, this correlation may explain the slightly higher frequencies on VSs.

The propagation spatial extent was the longest on EEG VSs (411mm or 6 sensors per propagation) followed by MEG VSs and icEEG (32 and 36mm or 5 sensors; see Table 3). Temporal extent was longer on icEEG than VSs for both HD-EEG and MEG (86 vs 27 vs 22 milliseconds; see Table 3). Propagation onsets encompassed more channels when recorded by VSs than icEEG: 4 VSs per sequence for HD-EEG and MEG compared to 1 electrode for icEEG.

**RZ and ROZ.** ROC curve analysis showed that virtual ripple rate was able to identify the icEEG-defined RZ with median AUC of 0.76 (0.57–0.79) and 0.65 (0.54–0.79) for HD-EEG and MEG, respectively (\( p = 0.24 \); Fig 4A).

AUC for HD-EEG and MEG was greater than 0.5 in 86% and 89% of our cohort, respectively. D_{icEEG} of the RZ defined by VSs did not differ between HD-EEG and MEG (\( p = 0.6 \), 27 vs 26mm; see Fig 4A), nor did their sensitivity (HD-EEG, 71% [9–100]; MEG, 50% [0–100]; \( p = 0.67 \)) or specificity to the icEEG RZ (HD-EEG, 63% [51–81]; MEG, 69% [50–78]; \( p = 0.77 \)).

Virtual onset ripple rate showed an AUC of 0.70 (0.35–0.85) and 0.64 (0.45–0.84) for HD-EEG and MEG, respectively (\( p = 0.8 \); see Fig 4B). AUC for HD-EEG and MEG was greater than 0.5 in 68% and 61% of our cohort. D_{icEEG} of the ROZ defined by the VSs did not differ between HD-EEG and MEG (\( p = 0.4 \), 24 and 22mm; see Fig 4B), nor did their sensitivity (HD-EEG, 61% [0–100]; MEG, 100% [0–100]; \( p = 0.85 \)) or specificity to the icEEG ROZ (HD-EEG, 76% [58–87]; MEG, 70% [53–81]; \( p = 0.19 \)). Average extent of VSs with high ripple rate and onset ripple rate was 11.3 and 6.3 cm, respectively, when estimated via HD-EEG, and 7 and 5.3 cm via MEG.

**Overlap with Resection and Outcome**

The virtual ROZ estimated via EEG VSs was closer to resection in good than in poor outcomes (\( p = 0.0026 \), \( D_{RES} = 8 \) vs 18mm); this was not found for the virtual RZ (see Table 2).
| #/Sex | Epilepsy Age, yr | Onset, yr Side | MRI Findings | icEEG Type (contacts, n) | icEEG Location | Spike Clusters, HD-EEG/MEG, n* | Res Lobe Vol, % | Engel (f/u, mo) |
|-------|-----------------|---------------|--------------|---------------------------|----------------|-----------------------------|----------------|----------------|
| 1/M   | 10              | 4             | R            | Normal                    |                |                            |                |                |
| 2/F   | 7               | 3             | L            | FCD (T and Ins)            |                |                            |                |                |
| 3/F   | 9               | 0.3           | L            | Hippocampal sclerosis      |                |                            |                |                |
| 4/F   | 13              | 10            | L            | Normal                    |                |                            |                |                |
| 5/M   | 17              | 9             | L            | Tumor (T)                 |                |                            |                |                |
| 6/M   | 2               | 0.33          | R            | TSC (multifocal)          |                |                            |                |                |
| 7/F   | 8               | 4             | R            | FCD (P)                   |                |                            |                |                |
| 8/M   | 18              | 8             | L            | FCD (mesial T, periventricular) |                |                            |                |                |
| 9/F   | 18              | 15            | L            | Normal                    |                |                            |                |                |
| 10/M  | 15              | 4             | L            | Normal                    |                |                            |                |                |
| 11/M  | 16              | 4             | L            | Normal (mild gliosis)      |                |                            |                |                |
| 12/F  | 9               | 1             | R            | Low-grade neoplasm (Fr)   |                |                            |                |                |
| 13/F  | 18              | 4             | L            | FCD (Fr)                  |                |                            |                |                |
| 14/F  | 14              | 6             | L            | FCD (mesial P)            |                |                            |                |                |
| 15/M  | 13              | 8             | L            | Encephalomalacia (P, superior T) |                |                            |                |                |
| 16/M  | 12              | 7             | L            | FCD (T)                   |                |                            |                |                |
| 17/M  | 13              | 0             | L            | Infarct (MCA territory)   |                |                            |                |                |
| 18/M  | 22              | 5             | L            | FCD (C P)                 |                |                            |                |                |
| 19/M  | 11              | 1             | L            | FCD (mesial T)            |                |                            |                |                |
| 20/M  | 16              | 5             | L            | FCD                        |                |                            |                |                |
| 21/M  | 10              | 7             | L            | Polymicrogyria (Fr, P)    |                |                            |                |                |
| 22/F  | 9               | 8             | L            | FCD (Fr)                  |                |                            |                |                |
| 23/F  | 7               | 4             | R            | FCD (Fr operculum)        |                |                            |                |                |
| 24/F  | 15              | 3             | L            | None                      |                |                            |                |                |
| 25/F  | 7               | 6             | L            | FCD (posterior Fr)        |                |                            |                |                |
| 26/F  | 18              | 3             | L            | FCD (Fr)                  |                |                            |                |                |
| 27/F  | 4               | 0.5           | R            | None                      |                |                            |                |                |
| 28/M  | 10              | 5             | L            | None                      |                |                            |                |                |

*Spine source localization: lobe containing main dipole cluster.

ant = anterior; CP = central; DE = depth electrodes; EEG = electroencephalography; F = female; f/u = follow-up; FCD = focal cortical dysplasia; Fr = frontal; HD = high-density; icEEG = intra cranial EEG; IE = interhemispheric; Ins = insula; L = center; M = male; MCA = middle cerebral artery; MEG = magnetoencephalography; MRI = magnetic resonance imaging; O = occipital; P = parietal; post = posterior; R = right; Res = resection; SE = subdural electrodes; T = temporal; TSC = tuberous sclerosis complex; Vol = volume.
Fig 4C; \( p = 0.074, 13 \text{ vs } 18\text{mm})\). Resection percentages of virtual ROZ \( (p = 0.009)\) and virtual RZ \( (p = 0.03)\) estimated via EEG VSs were higher in good than in poor outcomes (see Fig 4D; \( p = 0.009, p = 0.03)\). Resection percentages of virtual ROZ estimated via MEG VSs was higher in good than in poor outcomes \( (p = 0.025)\).
whereas this was not the case for the virtual RZ ($p = 0.12$). $D_{RES}$ of both virtual ROZ and virtual RZ estimated via MEG VSs did not differ between outcome groups ($p = 0.0887$, $p = 0.082$).

Regarding spikes, the resection percentage, as well as $D_{RES}$, did not differ between good and poor outcomes (see Fig 4D, E) for HD-EEG ($p = 0.72$, $p = 0.42$) and MEG ($p = 0.83$, $p = 0.96$). Additionally, in good outcome patients (proof of successful resection), both virtual RZ and virtual ROZ were closer to resection compared to spike localization for both HD-EEG ($p = 0.0016$, $p < 0.001$) and MEG ($p = 0.003$, $p = 0.001$). Table 4 reports distances and overlap between spike sources and ripple zones.

Finally, virtual RZ presented a $D_{SOZ}$ of 14 and 16mm for MEG and HD-EEG, respectively ($p = 0.01$), and overlap with SOZ of 33% (19–45%) and 29% (13–40%, $p = 0.0095$). Virtual ROZ showed a $D_{SOZ}$ of 13 and 14mm for MEG and HD-EEG, respectively ($p = 0.68$), and overlap of 33% (15–49%) and 30% (15–50%, $p = 0.64$).

### Outcome Prediction

At the individual patient level, resecting the virtual ROZ predicted good outcome (see Table 4), with PPV of 91%, NPV of 59%, and accuracy of 71% when estimated via EEG VSs ($p = 0.016$), and PPV of 83%, NPV of 60%, and accuracy of 70% when estimated via MEG VSs ($p = 0.047$). In contrast, no association was found for the virtual RZ estimated via EEG VSs ($p = 0.125$), whereas a weak association was found with MEG VSs ($p = 0.054$; Table 4).

When looking at spikes, resecting most of dipoles (resection percentage > 50%) did not predict outcome for HD-EEG ($p = 0.72$) or MEG ($p = 0.67$).

### Discussion

We present for the first time the noninvasive mapping of interictal ripple propagation in children with MRE using electric and magnetic source imaging. We previously showed ripple spatiotemporal propagation on icEEG, demonstrating the prognostic value of its onset generator (ROZ) for pediatric epilepsy surgery compared to areas of spread.21 This earlier study, followed by Otárula et al,22 demonstrated that ictal HFOs are not isolated events on icEEG, but pathophysiological activity organized in networks. Here, we present evidence that this epileptogenic phenomenon of ripple propagation can be captured noninvasively through a virtual implantation (reconstructed using HD-EEG or MEG) and show its value as an outcome predictor in children with MRE.
FIGURE 4: Results from validation of virtual zones versus invasive electroencephalography (EEG) and resection. Significant differences ($p < 0.05$) are marked with asterisks. (A, B) Validation of the virtual mapping of ripples (A) and onset ripples (B) against the benchmark given by intracranial EEG (iEEG). Each scenario shows the boxplots of the (1) area under the curve (AUC) obtained from each patient using high-density (HD)-EEG virtual sensors (VSs; orange) and magnetoencephalography (MEG) VSs (blue); and (2) median distance of VSs from intracranial ripple zone (RZ) or intracranial ripple onset zone (ROZ; DicEEG) in centimeters. Differences between modalities were not statistically significant (N.S.; $p > 0.05$). No difference was observed when we differentiated DicEEG for the icEEG contacts in the gray and white matter ($p > 0.1$). (C, D) Boxplots of the distance from resection ($D_{DRES}$; C) and resection percentage (D) for virtual RZ (in green) and virtual ROZ (in red) in good versus poor outcome (solid vs dashed box) patients. Each scenario shows boxplots for both HD-EEG VSs and MEG VSs. (E, F) Boxplots of $D_{DRES}$ (E) and resection percentage (F) for interictal spikes in good versus poor outcome (solid vs dashed box) patients. Each scenario shows boxplots for both HD-EEG VSs and MEG VSs.
data indicate that ripple propagation, as captured by VSs, reflects the hierarchical epileptogenic organization seen on icEEG;21 mapping the ripple onset generator (virtual ROZ) estimates the EZ better than mapping all ripple generators independently from propagation (virtual RZ). This derives from our main findings: (1) EEG VSs and MEG VSs, placed over specific brain areas, can capture ripples and their propagation onset; (2) areas generating ripples (RZ) and initiating their propagation (ROZ) are localized via VSs with an accuracy that is similar between HD-EEG and MEG; and (3) resecting the virtual ROZ, but not virtual RZ or spike sources, predicts good outcome (seizure freedom) in pediatric epilepsy surgery.

**Ripple Propagation Is Captured Noninvasively by VSs**
The concept of propagation in epilepsy is typically used when interpreting seizure semiology or ictal recordings. For scalp EEG or MEG, the concept of onset is generally applied when interpreting ictal activity (SOZ) but also interictal spikes if propagation is seen.39–41 Only recently, interictal propagation was also reported for ripples on icEEG.21,22 Here, we add to that evidence, revealing this phenomenon on HD-EEG or MEG VSs as well; we found that interictal ripples define distinct spatiotemporal sequences across multiple (4–7) neighboring VSs (see Fig 2) as observed on invasive recordings, although with lower rates. Ripple onset areas on VSs showed an overall spatial concordance with icEEG, although in terms of duration, propagations appear shorter on VSs than icEEG (see Table 3), suggesting that VSs may miss some later spread, which is seen instead on icEEG. Given the lack of association between spread ripples and EZ demonstrated on icEEG,21 the absence of such activity on VSs does not hamper presurgical clinical relevance. Furthermore, ripple

|TABLE 4. Predictive Value of Removing the Virtual Zones |
|---|---|---|---|
|**Modality** | **Ripple Zone** | **Ripple Onset Zone** | |
| | EEG VSs | MEG VSs | EEG VSs | MEG VSs |
| Virtual rate threshold | 0.25 | 0.5 | 0.55 | 0.35 |
| Residual, ≥50% not resecteda | No | Yes | No | Yes |
| Outcome | | | | |
| Good, seizure-free | 5 | 12 | 10 | 7 |
| Poor, seizure recurrence | 0 | 11 | 2 | 9 |
| Total | 5 | 23 | 12 | 16 |
| PPV | 100% | 83% | 91% | 83% |
| NPV | 48% | 56% | 59% | 60% |
| Accuracy | 57% | 68% | 71% | 70% |
| Phi | 0.38 | 0.40 | 0.50 | 0.44 |
| Fisher exact test | 0.125 | 0.054 | 0.016b | 0.047b |

Comparison of ripple and spike localization

| Distance of ripple VSs from spikes, mmc | 23 (18–29) | 17 (14–25) | 15 (12–29) | 16 (12–27) |
| Distance of spikes from ripple VSs, mmd | 24 (19–30) | 30 (25–41) | 27 (18–35) | 29 (20–41) |
| Overlap with spikes | 14% (3–22%) | 17% (4–36%) | 24% (0–40%) | 14% (0–38%) |

Numbers in parentheses represent interquartile range.

aA zone was regarded as “residual” after surgery if the resection percentage was <50%.

bSignificant p value (>0.05); Fisher exact test.

cDistance of the VSs (center) from the closest spike source (dipole).

dDistance of the spike sources from the closest VS (center).

eOverlap was defined as the percentage of VSs with a distance from the closest dipole < 10 mm.

EEG = electroencephalography; MEG = magnetoencephalography; NPV = negative predictive value; Phi = Phi coefficient of association; PPV = positive predictive value; VS = virtual sensor.
propagation on VSs covers slightly larger distance than on icEEG (36–41 vs 32mm) and larger areas of onset (4 VSs vs 1 icEEG contact); this possibly reflects the lower spatial resolution of noninvasive techniques and thus lower accuracy in estimating the extent of the generators.

Our findings provide the first robust evidence that ripple propagation can be captured using virtual, rather than invasive, sensors and stimulate further methodological studies to develop automated methods to investigate ripple propagation via full-coverage VSs. Although our VSs were placed to replicate icEEG retrospectively, our findings suggest that they can be eventually placed at desired locations in a prospective way.

**HD-EEG and MEG VSs Map Ripples and Their Onset with Similar Accuracy**

We quantified the ability of VSs to map ripples (RZ) and their propagation onset (ROZ) with respect to icEEG. We observed no difference between HD-EEG and MEG performance, both showing $D_{\text{EEG}} < 3 \text{ cm} (RZ \sim 26\text{mm}, ROZ \sim 23\text{mm})$. AUCs < 0.5, observed in some patients, denote spatial discrepancy between virtual and icEEG ripple rates; this may be explained by the fact that HD-EED/MEG and icEEG recordings were not simultaneous and thus may have captured different ripple sources due to their variability across time.$^{42,43}$

We can assume that VSs over specific brain areas grant a ripple localization within the same (or most proximal) gyrus (~2–3cm) that is indicated by an invasive implantation covering the same areas. Reconstructing high-frequency activation noninvasively with 2 to 3cm accuracy could impact surgical planning, enabling the clinical team to optimize implantation options. Nonetheless, when the EZ is adjacent to eloquent areas, more accurate biomarkers are required. Overall, our data demonstrate spatial consistency between ripples on VSs and icEEG, suggesting they are expressions of the same underlying phenomenon$^{15,44}$ and expanding this notion to their propagation onset.

Regarding the comparison with spikes, although the vast majority of the virtual ripple zones did not overlap in a strict sense (<10mm apart) with spike sources, they seem to be just adjacent (15–23mm; see Table 4); preponderant spike clusters presented lobar concordance with ripples in most cases (see Table 2).

**Virtual Mapping of Onset Ripples Estimates the EZ Better Than All Ripples or Spikes**

In the presurgical context, it is key to evaluate an epilepsy biomarker with respect to the ground truth of good outcome after resection. We found that identifying areas of ripple onset (virtual ROZ) reflects epileptogenicity better than identifying indistinguivcely all VSs showing ripples (virtual RZ) regardless of any propagation characteristics. For the virtual RZ, we found no differences in its proximity to resection between outcomes. In contrast, virtual ROZ was closer to resection in good than poor outcome patients when estimated via EEG VSs; in addition, in good outcomes, it was closer to resection (8mm) than the virtual RZ (13mm). For MEG VSs, the resection percentage of the virtual ROZ (but not virtual RZ) was higher in good (59%) than in poor outcome patients (38%). These noninvasive findings corroborate previous icEEG data$^{21}$ that emphasized the pathological nature of onset ripples as opposed to other ripples (spread or isolated), which seem more likely to reflect physiological mechanisms. Additionally, spike localization with HD-EEG or MEG presented lower performance than virtual RZ or virtual ROZ: (1) resection percentage and $D_{\text{RES}}$ did not differ between outcomes; and (2) in good outcomes, virtual RZ and virtual ROZ were closer to resection than spikes (dipoles). Because dipoles do not estimate the spike spatial extent, we acknowledge that our reported distances from resection may represent an overestimate and warrant further investigations with distributed source modeling (DSM).$^{45}$ However, because DSM on spikes was recently reported to provide an average improvement of only ~1mm (compared to dipoles) in adults,$^{46}$ we speculate that this may not affect significantly our main findings.

**Resection of ROZ, but Not Entire RZ or Spikes, Predicts Good Outcome**

To assess clinical utility in terms of individualized care, we investigated the predictive value of resecting virtually defined zones. Removing most of the virtual ROZ predicted outcome with PPV and NPV of 83–91% and 59–60%, suggesting the prognostic value of targeting this zone during surgery. In contrast, targeting the virtual RZ, that is, regions generating any type of ripple, presented lower predictive values. Additionally, spike localization provided complementary information to the ROZ or RZ (given their low overlap) and did not carry prognostic value. We showed the superiority of mapping ripple onset compared to indistinctively locating all the sites showing ripples on VSs or localizing spikes in the traditional sense. This adds to previous efforts aimed at recognizing the most pathological ripples, which so far was mostly performed through their relationship with spikes$^{15,19}$ or morphological features.$^{47–49}$ Our findings reveal that the spatiotemporal characterization of ripples enhances their interpretation on scalp HD-EEG and MEG; evaluating whether it is possible to resect the areas initiating the
propagation on VSs provides an additional noninvasive estimate of the EZ, especially when invasive monitoring cannot be easily planned.

Although the association between outcome and virtual RZ resection did not reach significance, it showed the same predictive trend for MEG VSs ($p = 0.054$; see Table 4) and EEG VSs (see Fig 4D). This suggests that ripples on HD-EEG/MEG are less undermined by physiological counterparts than on iEEG$^{50}$ (which is more sensitive to low amplitude and spread activity); this also explains low ripple rates typically observed in noninvasive studies.$^{14,15,25}$

**Limitations**

Applicability of our approach is limited to the eligible patients; we excluded 1-stage resections and children who were not referred for HD-EEG/MEG. Generalizability to spike-negative patients and reproducibility across HD-EEG/MEG sessions need further investigation. We constructed VSs at the iEEG locations to allow direct comparison with iEEG (the technique of choice to record ripples and propagation); future studies are warranted to assess the potential of whole-brain VSs. Although automated ripple detection cannot fully exclude artifacts, our detector is particularly robust, because it rejects events showing elongated time–frequency blobs. Our analysis excluded fast ripples, because there is limited evidence of their presence on scalp EEG or MEG, and we had a 600Hz sampling rate in several recordings. In addition, some of the observed ripples may be physiological (eg, low-amplitude ripples in Fig 2); further MEG/HD-EEG investigations of ripples overlapping on spikes are warranted. Finally, the use of HD-EEG systems with a higher number of channels (eg, 256)$^{45}$ is likely to improve the localization accuracy of ripples and allow fairer comparisons with MEG.

**Conclusions**

We revealed the noninvasive mapping of interictal ripple propagation in children with MRE, using HD-EEG and MEG, and demonstrated its prognostic value. Performing a noninvasive implantation of VSs over specific brain areas allows mapping the ripple propagation onset (ROZ), an area that yields prognostic value as an EZ biomarker in epilepsy surgery. Removing areas of ripple onset, as defined by the virtual implantation, predicts good outcome better than the entire area generating ripples or spikes. This noninvasive mapping may augment surgical planning, potentially improving outcomes of pediatric epilepsy surgery.

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**Author Contributions**

E.T., S.M.S., P.L.P., and C.P. contributed to the conception and design of the study. E.T., M.A.G.M., G.N., J.R.M., P.E.G., and C.P. contributed to the acquisition and analysis of data. E.T., M.A.G.M., P.L.P., and C.P. contributed to drafting a significant portion of the manuscript and figures.

**Potential Conflicts of Interest**

Nothing to report.

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