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Resection of high frequency oscillations predicts seizure outcome in the individual patient

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High frequency oscillations (HFOs) are recognized as biomarkers for epileptogenic brain tissue. A remaining challenge for epilepsy surgery is the prospective classification of tissue sampled by individual electrode contacts. We analysed long-term invasive recordings of 20 consecutive patients who subsequently underwent epilepsy surgery. HFOs were defined prospectively by a previously validated, automated algorithm in the ripple (80–250 Hz) and the fast ripple (FR, 250–500 Hz) frequency band. Contacts with the highest rate of ripples co-occurring with FR over several five-minute time intervals designated the HFO area. The HFO area was fully included in the resected area in all 13 patients who achieved seizure freedom (specificity 100%) and in 3 patients where seizures recurred (negative predictive value 81%). The HFO area was only partially resected in 4 patients suffering from recurrent seizures (positive predictive value 100%, sensitivity 57%). Thus, the resection of the prospectively defined HFO area proved to be highly specific and reproducible in 13/13 patients with seizure freedom, while it may have improved the outcome in 4/7 patients with recurrent seizures. We thus validated the clinical relevance of the HFO area in the individual patient with an automated procedure. This is a prerequisite before HFOs can guide surgical treatment in multicentre studies.

The treatment of choice in patients with drug-resistant focal epilepsy is the surgical resection or disconnection of the epileptogenic zone (EZ). The EZ is defined as the region indispensable for generating seizures. Recent studies have pointed to the high frequency oscillation (HFO) recorded in intracranial EEG (iEEG) as a new indicator for the EZ. HFOs are generally viewed as spontaneous EEG patterns in the frequency range between 80–500 Hz that consist of at least four oscillations that clearly stand out of the background activity. HFOs are differentiated into “ripples” (80–250 Hz) and “fast ripples” (FRs, 250–500 Hz). Interictal HFOs have proven more specific in localizing the seizure onset zone (SOZ) than spikes and have presented a good association with the post-surgery outcome in epilepsy patients. While most studies to date only report group results with mean HFO rates in SOZ electrodes exceeding mean rates in non-SOZ electrodes, surgical decisions require the precise classification of cortex sampled by individual electrodes.

To facilitate HFO analysis, several algorithms for their automated or semi-automated detection have been proposed. As an advance, we apply here a fully automated definition of HFOs, which we have previously optimized on visual markings in a separate large dataset. While we do not aim to outperform visual marking, we provide a prospective definition of a clinically relevant HFO. We define as HFO area the electrode contacts with the HFO rate exceeding the 95% percentile of the HFO distribution and we focus on ripples co-occurring with FR (FRandR) as a new entity. We quantify the reproducibility of the HFO area over multiple time intervals from different nights of pre-operative monitoring. We evaluate the clinical relevance of HFO in individual electrodes by comparing the HFO area with the resected area (RA) and then by predicting seizure outcome in the individual patient.

Results

We included 20 patients (Table 1) of whom 9 with mesial temporal lobe epilepsy (TLE) and 11 with extratemporal epilepsy (ETE). Complete seizure freedom (ILAE 1 - International League Against Epilepsy, class 1) was reached in 13 patients, while 7 patients suffered from recurrent seizures (seizure-free rate = 65%).

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The HFO area predicted seizure outcome in the individual patient. Automatic HFO detection yielded ripples (median amplitude 27.3 µVpp, interquartile range 16.0–41.5 µVpp) and FR (median amplitude 9.2 µVpp, interquartile range 7.5–14.5 µVpp). For each individual patient, we determined whether the HFO area was fully or partly resected and whether seizure freedom was achieved (Table S1). The mean follow-up for poor outcome (29 ± 15 mo) was longer than for good outcome (23 ± 11 mo) but the two distributions are indistinguishable (Wilcoxon rank sum test, p = 0.25). When defining the HFO area by FRandR (FRandR area, Fig. 1), we achieved specificity 100%, CI [78–100%], which was more specific in predicting seizure freedom than FR and ripples separately (Table 2). The mean follow-up for poor outcome (29 ± 15 mo) was longer than for good outcome (23 ± 11 mo) but the two distributions are indistinguishable (Wilcoxon rank sum test, p = 0.25).

Test-retest reliability of the spatial distribution of HFO rates. To evaluate the reproducibility of the spatial distribution of HFO, we calculated the HFO rates during several periods of slow wave sleep for 18 patients. To illustrate the procedure, the test-retest HFO analysis for Patient 1 is presented in Fig. 3. First, we quantified the HFO rates for different nights (Fig. 3A). We then computed the scalar product for all pairs of HFO vectors within and between nights and tested significance against a random distribution (Fig. 3B).

Across the group of 18 patients (Table 1), the percentage of significant scalar products had median of 97.5% (lower quartile 67%) when taking pairs of all 5 minutes intervals, and a median of 100% (lower quartile 100%) when taking pairs of entire nights. Over the intervals of TLE patients, the median was >98% irrespective of outcome. Over the intervals of ETE patients, the median was 97% for patients with good outcome and only 36% for patients with poor outcome.

Spectral analysis of ripples co-occurring with FR. In order to identify the spectral signature of FRandR, we selected the bipolar channel with highest FRandR rate included in the HFO area of each patient. For each detected FRandR, we computed the instantaneous power spectrum and detected peaks and troughs in the HFO range (80–500 Hz). Out of 6182 detected events, 45% exhibited two distinct peaks in ripple and FR band with the second peak at median frequency 240 Hz (Fig. 4A). 28% had only one peak in the HFO range (median frequency 184 Hz, Fig. 4B), and 27% did not show a prominent peak >100 Hz in the HFO range. No significant difference was observed between TLE and ETE channels. Since the algorithm does not use the power spectrum to detect a single event, the finding of a prominent spectral peak indicates that an HFO is a distinct spectral entity.

Discussion

Building on a prospective definition of HFO, our algorithm classified tissue sampled by each individual electrode as epileptogenic or normal as validated by the prediction of seizure outcome.

FRandR predict seizure freedom in the individual patient. As a main strength of our study, the resection of FRandR provided a high specificity in predicting seizure freedom (13/13 patients, 100%) with narrow confidence intervals (75–100%). Such high specificity not only generalizes the value of FRandR across different types of patients, but still holds true at the level of the individual patient. Different from other studies that considered only HFO summed across the SOZ13,23, we based our analysis on post-surgical seizure freedom, which is a prerequisite to guide epilepsy surgery.

In all seizure free patients with ETE (patients 10–16) the RA was limited to 1–2 cm² whereas the SOZ extended also to eloquent cortex. While the SOZ was therefore not fully resected in all ETE patients, the HFO area as defined by FRandR was always fully resected. Thus, compared to the SOZ, FRandR was a more specific marker of the EZ in the individual patient.

FRandR in patients with recurrent seizures. Among the 7 patients with recurrent seizures, FRandR were fully resected in 3 FN and not fully resected in 4 TP, resulting in a sensitivity of 57% CI [18–90%]. Among the 3 FN (15%), in patient 9 the HFO area was fully included in the RA but seizures recurred, which may reflect the limited coverage of the implanted electrodes. Patients 7 and 8 were seizure free under antiepileptic drugs but missed the intake of one dosage in noncompliance and consequently suffered a single recurrent seizure, so their seizure outcome was rated ILAE 3. In these cases, HFO guidance was at least consistent with routine surgical planning. There were 4 patients where the HFO area was not fully resected and who suffered from recurrent seizures (TP). In patient 17 the RA was much larger than the 1–2 cm² mentioned above and almost the entire right frontal cortex was resected. While parts of the HFO area were indeed included in this resection, the resection of the HFO area was not complete. In patient 18, our analysis identified the HFO area over occipital contacts ipsilateral to the resection, which included parietal cortex. Patient 19 was implanted with a parietal 4 × 8 electrode grid and a temporal 2 × 8 temporal grid covering multiple cavernoma. The HFO area included the parietal cavernoma but not the temporal cavernoma. The surgical choice was to resect only the temporal cavernoma. In patient 20, the resection included the seizure onset area in the posterior margin of the left occipito-temporal gyrus but the...
Automated detection predicts seizure outcome. Our automated analysis provides a prospective definition of the HFO area, which was then subjected to an unbiased evaluation of its clinical relevance. In contrast, expert observers’ HFO analysis suffers from time-consuming visual marking and lacks reproducibility. In our study, the high reproducibility of the spatial HFO distribution within and between nights confirms the reliability of the HFO analysis, but it may even provide a level of confidence for its diagnostic power. The most rigorous validation for HFO analysis is its comparison with post-surgical outcome. To date, only a few studies correlated automated HFO analysis with patients’ outcome. Our algorithm used here was trained with data from different epilepsy centres and marked by different expert viewers. As a major advance over previous studies, we applied here a fully automated procedure. With minimal clinical information, expert observers’ HFO analysis suffers from time-consuming visual marking and lacks reproducibility. In contrast, our automated analysis provides a prospective definition of the HFO area, which was then subjected to an unbiased evaluation of its clinical relevance. In our study, the high reproducibility of the spatial HFO distribution within and between nights confirms the clinical value of HFO as a reliable biomarker for epileptogenic tissue. Interestingly, ETE patients with poor outcome stood out in the low reproducibility of their HFO distribution. The test-retest analysis thus not only shows the reliability of the HFO analysis, but it may even provide a level of confidence for its diagnostic power.

**Table 1.** Patient characteristics. Abbreviations: C = central; depth = depth electrode; ETE = extratemporal lobe epilepsy; F = frontal; FCD = focal cortical dysplasia; FR = fast ripple; FRandR = FR occurring together with ripple; grid = grid electrode; HFO = high frequency oscillation; HS = hippocampus sclerosis; ILAE = International League Against Epilepsy; MTL = mesial temporal lobe; L = left; Lat = lateral; Les = lesionectomy; O = occipital; P = parietal; Pr = precentral; R = right; sAHE = selective amygdala-hippocampectomy; strip = strip electrode; T = temporal; TLE = mesial temporal lobe epilepsy. Test-retest intervals/nights: percentage of scalar products between pairs of intervals/nights exceeding the significance threshold (see Methods: Test-retest reliability of the spatial distribution of HFO). Over the intervals of TLE patients, the median was >98% irrespective of outcome. Over the intervals of ETE patients, the median was 97% for patients with good outcome and only 36% for patients with poor outcome.

| Patient | Age, Gender | Histology/Pathology | Epilepsy | Electrode placement | Type of electrodes | Surgery | Nights | Intervals | Test-retest intervals | Test-retest nights | Ripple area resected? | FR area resected? | FRandR area resected? | Outcome (ILAE) | Postoperative follow-up (months) |
|---------|-------------|---------------------|----------|---------------------|-------------------|---------|--------|----------|-------------------|-------------------|-------------------|----------------|----------------------|----------------|-----------------------------|
| 1       | 25, M       | HS and gliosis      | TLE      | MTL, L, R           | 5 depth, 1 strip \(4 \times 1\), 1 strip \(6 \times 1\) | sAHE, Les | 4      | 28       | 0.99   | 1.00             | N                 | N                 | Y              | Y                    | 1              | 12                          |
| 2       | 33, M       | Glioma              | TLE      | MTL, L, R           | 8 depth            | sAHE, Les | 2      | 13       | 1.00   | 1.00             | Y                 | Y                 | Y              | Y                    | 1              | 29                          |
| 3       | 20, F       | HS                  | TLE      | MTL, L, R           | 5 depth            | sAHE, Les | 5      | 39       | 0.78   | 1.00             | N                 | Y                 | Y              | Y                    | 1              | 13                          |
| 4       | 20, F       | HS                  | TLE      | MTL, L, R           | 8 depth            | sAHE, Les | 6      | 34       | 1.00   | 1.00             | N                 | Y                 | Y              | Y                    | 1              | 41                          |
| 5       | 40, M       | HS                  | TLE      | MTL, L, R           | 8 depth            | sAHE, Les | 5      | 35       | 0.94   | 1.00             | Y                 | Y                 | Y              | Y                    | 1              | 14                          |
| 6       | 48, M       | HS                  | TLE      | MTL, L, R           | 8 depth            | sAHE, Les | 5      | 35       | 0.98   | 1.00             | Y                 | Y                 | Y              | Y                    | 1              | 11                          |
| 7       | 25, M       | HS                  | TLE      | MTL, L, R           | 8 depth            | sAHE, Les | 1      | 1        | —      | —                | Y                 | Y                 | Y              | Y                    | 3              | 42                          |
| 8       | 21, F       | HS                  | TLE      | MTL, L, R           | 8 depth            | sAHE, Les | 2      | 16       | 0.67   | 1.00             | Y                 | Y                 | Y              | Y                    | 3              | 15                          |
| 9       | 52, M       | HS                  | TLE      | MTL, L, R           | 8 depth            | sAHE, Les | 2      | 12       | 1.00   | 1.00             | Y                 | Y                 | Y              | Y                    | 5              | 46                          |
| 10      | 37, M       | FCD 2b              | ETE      | Pr, R               | 1 grid \(8 \times 4\), 2 strips \(4 \times 1\) | Les | 1      | 6        | 1.00   | —                | N                 | Y                 | Y              | Y                    | 1              | 36                          |
| 11      | 36, M       | FCD 2b              | ETE      | F, R                | 1 grid \(8 \times 8\), 1 depth | Les | 3      | 19       | 1.00   | 1.00             | Y                 | Y                 | Y              | Y                    | 1              | 37                          |
| 12      | 49, M       | Ganglioglioma       | ETE      | T-Lat, L            | 1 grid \(8 \times 4\), 1 depth | Les | 4      | 25       | 0.53   | 0.67             | Y                 | N                 | Y              | Y                    | 1              | 25                          |
| 13      | 17, M       | FCD 1a              | ETE      | P, R                | 1 grid \(8 \times 8\), 1 depth | Les | 2      | 16       | 0.75   | 1.00             | N                 | N                 | Y              | Y                    | 1              | 25                          |
| 14      | 46, F       | FCD 1b              | ETE      | P, R                | 2 grids \(8 \times 2\), 1 strip \(6 \times 1\), 1 strip \(4 \times 1\), 1 depth | Les | 2      | 13       | 0.97   | 1.00             | Y                 | Y                 | Y              | Y                    | 1              | 10                          |
| 15      | 31, F       | Gliosis             | ETE      | T-Lat, L            | 1 grid \(8 \times 4\), 2 strips \(4 \times 1\) | Les | 4      | 28       | 0.32   | 0.33             | Y                 | N                 | Y              | Y                    | 1              | 25                          |
| 16      | 17, F       | FCD 2a              | ETE      | F, Pr, C, L         | 1 grid \(8 \times 4\), 1 depth | Les | 2      | 17       | 1.00   | 1.00             | N                 | Y                 | Y              | Y                    | 1              | 19                          |
| 17      | 30, M       | FCD 2a              | ETE      | F, R                | 2 grids \(8 \times 2\), 1 depth | Les | 1      | 1        | —      | —                | Y                 | Y                 | N              | 5                    | 45                          |
| 18      | 40, M       | FCD 2a              | ETE      | P, R                | 2 strips \(6 \times 1\), 1 depth | Les | 1      | 5        | 1.00   | —                | N                 | Y                 | N              | 5                    | 30                          |
| 19      | 38, M       | Cavernoma           | ETE      | T, P                | 1 grid \(8 \times 4\), 1 grid \(8 \times 2\) | Les | 2      | 13       | 0.36   | 0.00             | N                 | N                 | N              | 6                    | 11                          |
| 20      | 17, M       | FCD 3               | ETE      | O, T                | 1 grid \(8 \times 2\) | Les | 4      | 29       | 0.23   | 1.00             | N                 | N                 | N              | N                    | 5              | 16                          |
and minimal human monitoring of data quality, our automated algorithm analysed the iEEG by detecting HFOs and by rate thresholding to delineate the HFO area, which was then validated against clinical outcome.

**Limitations of rate thresholding.** In principle, rate thresholding could compromise the HFO area as a guide to the EZ. For example, if the EZ were not sampled by the electrodes at all, the HFO area would appear at

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**Figure 1.** Automated HFO analysis in patient 4. (A) Presurgical MRI shows right hippocampal sclerosis (HS). (B) MRI after the implantation of 8 depth electrodes sampling the mesial temporal structures. (C) MRI after right selective amygdala-hippocampectomy, resulting in long-term seizure freedom. (D) Example of a ripple co-occurring with a FR (FRandR, green line). (E,F,G) The HFO types differ in their distribution over channels. Given the neuroradiological diagnosis of this patient, we present here the electrode contacts located in the mesial temporal structures. Note that several channels have FRandR rate = 0. Channels with rates above the 95 percentile (black horizontal line) were taken to define the HFO area. Channels inside the resected area (RA) are shown in red, channels outside the RA in blue. The ripple area was not resected (E). The resection of the FRandR area (G, channel AR1-AR2) led to seizure freedom. Implantation sites: AL amygdala left; AR amygdala right; EL entorhinal cortex left; ER entorhinal cortex right; HL anterior hippocampus left; HR anterior hippocampus right; PL posterior hippocampus left; PR posterior hippocampus left.
a spurious location outside the EZ. Given the quality of our surgical planning, this did not occur in our patient series. As another limitation, the size of the HFO area will depend on the proportion of electrodes covering the EZ. This was, however, not a relevant source of error in our patient series, where the HFO area always provided an estimate of the EZ that seemed reasonable to the surgeon. It has to be kept in mind that HFO analysis is not intended to replace routine surgical planning and overrule the surgeon's decision but that it might provide added value in ambiguous cases.

FR and R are more specific than FR and ripple separately. The frequency bands for ripple (80–250 Hz) and FR (250–500 Hz) are often used in the literature and we therefore designed our algorithm to detect HFO patterns separately in these two bands. FRs are considered more specific for epileptogenicity than ripples because of their close relation to the SOZ and to the seizure outcome. Surprisingly, the co-occurrence of FRs and ripples was not evaluated before. Here we demonstrated that the resection of FR and R co-occurring with ripples was more specifically associated with seizure freedom than the resection of ripples and FRs separately.

In the instantaneous power spectrum, FR and R had at least one prominent peak around 200 Hz (median bimodal FR and R: 240 Hz, median unimodal FR and R: 184 Hz). In the high-pass filtered data (>250 Hz) the contribution of this peak matched the criteria for FR detection. In about half of the detected events (bimodal), there was a simultaneous occurrence of a ripple peak and a FR peak, suggesting the presence of two separate spectral entities. In the other half of the detected events (unimodal), a major spectral contribution in the ripple band spread over to the FR band. Our recording setup and our prospective definition of HFO includes both bimodal and unimodal events for the prediction of surgery outcome.

|               | Ripple | FR  | FR and R |
|---------------|--------|-----|----------|
| Specificity [%] | 54     | 69  | 100      |
| Sensitivity [%] | 43     | 29  | 57       |
| Negative Predictive Value NPV [%] | 64     | 64  | 81       |
| Positive Predictive Value PPV [%] | 33     | 33  | 100      |
| Accuracy [%] | 50     | 55  | 85       |

Table 2. HFO area and seizure outcome. Defining the HFO area by FR co-occurring with ripples was more specific in predicting seizure freedom than FR and ripples separately. False Negative cases may reflect the limited coverage of the implanted electrodes. Abbreviations: FN = False Negative; FP = False Positive; FR = fast ripple; FR and R = FR co-occurring with ripple; NPV = Negative Predictive Value; PPV = Positive Predictive Value; TN = True Negative; TP = True Positive; N = number of patients.

Figure 2. The HFO area exceeds the resected area in the patient with recurrent seizures (patient 17). (A) Coverage of the implanted electrodes. (B) Reconstruction of surgical resection. Red electrode contacts denote the seizure onset zone (SOZ). The HFO area (contacts filled yellow) is smaller than the SOZ. (C) The resected area (RA) does not comprise all channels of the HFO area. The patient suffers from recurrent seizures. Full resection of the HFO area might have achieved seizure freedom.

FR and R are more specific than FR and ripple separately. The frequency bands for ripple (80–250 Hz) and FR (250–500 Hz) are often used in the literature and we therefore designed our algorithm to detect HFO patterns separately in these two bands. FRs are considered more specific for epileptogenicity than ripples because of their close relation to the SOZ and to the seizure outcome. Surprisingly, the co-occurrence of FRs and ripples was not evaluated before. Here we demonstrated that the resection of FR and R co-occurring with ripples was more specifically associated with seizure freedom than the resection of ripples and FRs separately.
It remains open whether ripples and FR (or very high frequency oscillations) reflect distinct biological processes. HFO, and in particular FR, are thought to express abnormal connectivity patterns reflecting a local imbalance between excitatory and inhibitory circuitry in the epileptogenic zone. One direction of further investigation of the generative processes in humans might be the combination of macro- and micro-electrode recordings. Such a combination of electrodes probing different spatial extent might shed light on the complex spatio-temporal synchronization patterns and neuronal networks that generate HFO.

**Generalizability.** The results of our algorithm suggest that the information provided by prospectively defined HFOs could contribute to surgery planning in cases where the extent of surgical resection can be adapted. As an example, the exact extent of focal cortical dysplasia (FCD) is sometimes challenging to detect in MRI and remains difficult to delineate. It is in these patients where complementary electrophysiological markers such as HFO may be useful.

In general, several methods are currently discussed to improve surgery outcome. Promising are, e.g. strategies based on patient’s clinical characteristics. While HFO are spatio-temporally highly localized, the analysis of functional connectivity between electrode sites aims to identify the nodes with higher seizure likelihood, both during surgery and during the preoperative period. Given the two different spatio-temporal scales in iEEG analysis, the two approaches appear complementary and their combination might add to the understanding of the underlying pathophysiology.

In our study, the resection of the prospectively defined HFO area proved to be highly specific and reproducible in all patients with seizure freedom, while it may have improved the outcome in four patients with recurrent seizures.
Figure 4. Spectral analysis of FRandR events. We provide here two examples of FRandR events with bimodal (A) and unimodal (B) frequency spectrum, respectively. From top to bottom: the raw data, the signal filtered in the ripple band (80–250 Hz), the signal filtered in the FR band (250–500 Hz), the Stockwell time-frequency transform, the instantaneous power spectrum averaged over the duration of the FR. The duration of the FRandR is highlighted in red, the FR in bold. (A) A blob appears in the time-frequency domain at latency 135.53 s and central frequency 350 Hz in the FR spectral range. This explains the event in the FR band filtered signal. (B) A blob appears in the time-frequency domain at latency 68.25 s and central frequency 190 Hz in the ripple spectral range. Energy of the broad peak in the ripple band spreads to higher frequencies and appears as an event in the FR band filtered signal.

seizures. The HFO area was defined by a fully automated algorithm. We thus validated the HFO area against outcome in the individual patient, which is a prerequisite before HFOs can guide surgical treatment in multicentre studies.
Methods

Patient selection. We included all consecutive patients with drug-resistant focal epilepsy in our centre that 1) underwent invasive EEG recordings with subdural and/or depth electrodes from March 2012 to April 2016 as part of their presurgical evaluation, 2) consequently underwent resective surgery, and 3) were followed-up for > 1 year after surgery in case of seizure freedom. The postsurgical seizure outcome was determined in follow-up visits and classified according to the International League Against Epilepsy (ILAE)40.

Ethics statement. Collection of personal patient data and retrospective scientific workup was approved by the research ethics committee (Kantonale Ethikkommission KEK-ZH-Nr. 2012-0212) and collection of patients' written informed consent was waived. The study was performed in accordance with the relevant guidelines and regulations.

Electrode types and implantation sites. Subdural strip and grid electrodes as well as depth electrodes were placed according to the findings of the non-invasive presurgical evaluation (Fig. 1A,B).

In TLE patients, depth electrodes (1.3 mm diameter, 8 contacts of 1.6 mm length, spacing between contacts centres 5 mm, ADTech®, www.adtechmedical.com) were implanted stereotactically into the amygdala, the hippocampal head and the entorhinal and perirhinal cortex bilaterally.

In ETE patients, a combination of depth and subdural grid and strip electrodes (contact diameter 4 mm with a 2.3 mm exposure, spacing between contact centers 10 mm, ADTech®) was placed after craniotomy. Post-implantation MR images were used to locate each contact anatomically along the electrode trajectory.

Surgical planning. The decision for resection surgery was based on non-invasive investigations as well as on intracranial investigations1. The results of HFO analysis were not used for surgical planning.

Data acquisition. Intracranial data was acquired at 4000 Hz sampling frequency with an ATLAS recording system (0.5–1000 Hz pass-band, Neuralynx, www.neuralynx.com) and down-sampled to 2000 Hz for HFO analysis. In addition, scalp EEG according to the 10–20 system, with minor adaptations in order to avoid the surgical scalp lesions, and the submental electromyogram (EMG) were recorded. The iEEG was recorded against a common intracranial reference and then transformed to bipolar channels for further analysis.

Data selection. From each night recording we selected up to six intervals of five minutes of interictal slow-wave sleep that promotes low muscle activity and high HFO rates14,41. Sleep scoring was performed based on scalp EEG, electro-oculogram, EMG and video recordings. We chose time intervals at least three hours apart from epileptic seizures in order to eliminate the influence of seizure activity on our analysis. We excluded all electrode contacts where electrical stimulation evoked motor or language responses (Table S1). In TLE patients, we included only the 3 most mesial bipolar channels. The amount of nights and intervals varied across patients (Table 1).

Prospective definition of HFOs. HFOs were defined prospectively by an automated detector, which we had previously trained, tested and validated to detect visually marked events in datasets from the Montreal Neurological Institute22.

The detector incorporates information from both time and frequency domain and operates in two stages. In the first stage – baseline detection – baseline segments with low oscillatory activity were identified by the Stockwell entropy distribution17,42. The distribution of the envelope of baseline segments was used to define an amplitude threshold42. The second stage – HFO detection – was conducted separately for ripples (band-pass 80–240 Hz, stopband 70 Hz and 250 Hz, FIR equiripple filter with stopband attenuation 60 dB) and FRs (band-pass 250–490 Hz, stopband 240 Hz and 500 Hz). Events with the envelope of the filtered signal exceeding the amplitude threshold for at least 20 ms/10 ms were labelled as ripples/FR (Fig. 1D). The algorithm then identified FRs overlapping with a ripple, which we defined as a third type of HFO: FR co-occurring with ripples (FRandR). As an advantage over other automatic detectors, this algorithm reports several channels where no HFO occur (Fig. 1G).

The code is freely available at the HFO detectors repository on github (https://github.com/HFO-detect/HFO-detect-matlab). The iEEG data with the markings of our HFO events is freely available at CRCNS.org http://dx.doi.org/10.6080/K06Q1VD5.

Definition of the HFO area by rate thresholding. We analysed the spatial distribution of HFO rates across channels in each patient. The ensemble of those channels whose rates exceeded the rate threshold (95 percentile of the HFO rate distribution) was defined as the HFO area (Fig. 1E–G). The 95 percentile was computed across channels in each patient. The ensemble of those channels whose rates exceeded the rate threshold (95 percentile of the HFO rate distribution) was defined as the HFO area (Fig. 1E–G). The 95 percentile was computed by the Matlab function prctile.m. To compare HFO types, we identified an HFO area for all three types of HFOs (ripples, FRs and FRandR).

Test-retest reliability of the spatial distribution of HFO. We quantified the test-retest reliability of the distribution of HFO rates over the ensemble of slow wave sleep 5 minutes intervals (Fig. 3A). For each interval pair we computed the normalized scalar product of the spatial distribution of the HFO rates. The scalar product is ~1 for highly overlapping spatial distributions of HFO rate, and lower otherwise. To test the magnitude of the true scalar product against chance, we constructed a distribution of scalar products by randomly permuting (N = 5000) the order of channels for each interval. The true value of the scalar product was considered statistically significant if it exceeded the 97.5 percentile of the distribution23 (Fig. 3B). Next, we took the mean HFO rate for each night and tested reproducibility over nights (Fig. 3C).
Clinical validation of HFO against seizure outcome. We evaluated whether the HFO area was included in the resection area to quantify the predictive value of the HFO area with respect to seizure outcome (Table 1). Automated HFO detection and analysis were blind to clinical information. Following 13, we defined as true positive (TP) a patient where the HFO area was not fully located within the RA, i.e. at least one channel of the HFO area was not resected, and the patient suffered from recurrent seizures (ILAE 2–6). We defined as false positive (FP) a patient where the HFO area was not fully located inside the RA but who achieved seizure freedom (ILAE 1). We defined as false negative (FN) a patient where the HFO area was fully located within the RA but who suffered from recurrent seizures. We defined as true negative (TN) a patient where the HFO area was fully located inside the RA and who became seizure free. The positive predictive value was calculated as PPV = TP/(TP + FP), negative predictive value as NPV = TN/(TN + FN), sensitivity = TP/(TP + FN), specificity = TN/(TN + FP), and accuracy = (TP + TN)/N. Estimates of the 95% confidence intervals (CI) used the binomial method for sample size <10 and the asymptotic method for sample size >10 based on the normal approximation. Statistical significance was established at p < 0.05.

Spectral analysis of ripples co-occurring with FR. For each patient we selected the bipolar channel with highest FRandR rate included in the HFO area. We computed the Stockwell transform of the signal in an interval of 0.6 s around each detected FRandR. We extracted the averaged instantaneous spectrum over the duration of the detected FR and detected peaks and troughs in the HFO frequency band [80 500] Hz after least-squares fitting maximum likelihood estimation with gaussian distribution (four gaussians, lsqcurvefit.m in Matlab). We defined bimodal events as events with a trough between 150 and 250 Hz, and at least one peak above 200 Hz. We defined unimodal events as events with a single peak >100 Hz that is detectable in the ripple band.

References
1. Ryvlin, P. & Rheims, S. Predicting epilepsy surgery outcome. Current opinion in neurology 29, 182–188, https://doi.org/10.1097/WCO.0000000000000366 (2016).
2. Rosemow, F. & Lüders, H. Presurgical evaluation of epilepsy. Brain: a journal of neurology 124, 1683–1700 (2001).
3. Jacobs, J. et al. High-frequency oscillations (HFOs) in clinical epilepsy. Progress in neurobiology 98, 302–315, https://doi.org/10.1016/j.pneurobio.2012.03.001 (2012).
4. Urestarazu, E., Chander, R., Dubeau, F. & Gotman, J. Interictal high-frequency oscillations (100–500 Hz) in the intracerebral EEG of epileptic patients. Brain: a journal of neurology 130, 2354–2366, https://doi.org/10.1093/brain/awm149 (2007).
5. Zijlmans, M. et al. High-frequency oscillations as a new biomarker in epilepsy. Annals of neurology 71, 169–178, https://doi.org/10.1002/ana.22548 (2012).
6. Zijlmans, M. et al. How to record high-frequency oscillations in epilepsy: A practical guideline. Epilepsia (2017).
7. Bragin, A., Engel, J., Wilson, C. L., Fried, I. & Buzsáki, G. High-frequency oscillations in human brain. Hippocampus 9, 137–142, https://doi.org/10.1002/(sici)1098-1063(1999)9:2<137::aid-hipo3>3.0.co;2-0 (1999).
8. Jacob, J. et al. Interictal high-frequency oscillations (80–500 Hz) are an indicator of seizure onset area independent of spikes in the human epileptic brain. Epilepsia 49, 1893–1907, https://doi.org/10.1111/j.1528-1167.2008.01656.x (2008).
9. Wu, J. Y. et al. Removing interictal fast ripples on electrocorticography linked with seizure freedom in children. Neurology 75, 1686–1694, https://doi.org/10.1212/WNL.0b013e3181f73c7d (2010).
10. Akiyama, T. et al. Focal resection of fast ripples on extraoperative intracranial EEG improves seizure outcome in pediatric epilepsy. Epilepsia 52, 1802–1811, https://doi.org/10.1111/j.1528-1167.2011.03199.x (2011).
11. Jacob, J. et al. High-frequency electroencephalographic oscillations correlate with outcome of epilepsy surgery. Annals of neurology 67, 209–220, https://doi.org/10.1002/ana.21847 (2010).
12. van Klompenburg, M. A. et al. Residual fast ripples in the intraoperative corticogram predict epilepsy surgery outcome. Neurology 85, 120–128, https://doi.org/10.1212/WNL.0000000000001727 (2015).
13. Cimbalník, J., Kucewicz, M. T. & Worrell, G. Interictal high-frequency oscillations in focal human epilepsy. Current opinion in neurology 29, 175–181, https://doi.org/10.1097/WCO.0000000000000302 (2016).
14. Staba, R. J. et al. High-frequency oscillations recorded in human medial temporal lobe during sleep. Annals of neurology 56, 188–195, https://doi.org/10.1002/ana.20164 (2004).
15. Crepon, B. et al. Mapping interictal oscillations greater than 200 Hz recorded with intracranial macroelectrodes in human epilepsy. Brain: a journal of neurology 133, 33–45, doi:awp277 [pii] 10.1093/brain/awp277 (2010).
16. Gardner, A. B., Worrell, G. A., Marsh, E., Dlugos, D. & Litt, B. Human and automated detection of high-frequency oscillations in clinical intracranial EEG recordings. Clinical neurophysiology: official journal of the International Federation of Clinical Neurophysiology 118, 1134–1143, https://doi.org/10.1016/j.clinph.2008.12.019 (2007).
17. Zelmann, R. et al. A comparison between detectors of high frequency oscillations. Clinical neurophysiology: official journal of the International Federation of Clinical Neurophysiology 123, 106–116, https://doi.org/10.1016/j.clinph.2011.06.006 (2012).
18. Birot, G., Kachenoura, A., Albera, L., Bénar, C. & Wendling, F. Automatic detection of fast ripples. Journal of Neuroscience Methods 213, 236–249, https://doi.org/10.1016/j.jneumeth.2012.12.013 (2013).
19. Burnos, S. et al. Human intracranial high frequency oscillations (HFOs) detected by automatic time-frequency analysis. PloS One 9, e94381, https://doi.org/10.1371/journal.pone.0094381 (2014).
20. Ramantani, G. et al. Source reconstruction based on subdural EEG recordings adds to the presurgical evaluation in refractory frontal lobe epilepsy. Clinical neurophysiology: official journal of the International Federation of Clinical Neurophysiology 124, 481–491, https://doi.org/10.1016/j.clinph.2012.09.001 (2013).
21. Fedele, T. et al. Automatic detection of high frequency oscillations during epilepsy surgery predicts seizure outcome. Clinical neurophysiology: official journal of the International Federation of Clinical Neurophysiology 127, 3066–3074, https://doi.org/10.1016/j.clinph.2016.06.009 (2016).
22. Burnos, S. et al. The morphology of high frequency oscillations (HFO) does not improve delineating the epileptogenic zone. Clinical neurophysiology: official journal of the International Federation of Clinical Neurophysiology 127, 2140–2148, https://doi.org/10.1016/j.clinph.2016.01.002 (2016).
23. Höller, Y. et al. High-frequency oscillations in epilepsy and surgical outcome. A meta-analysis, Front Hum Neurosci 9, 574, https://doi.org/10.3389/fnhum.2015.00574 (2015).
24. Spring, A. M. et al. Interrater reliability of visually evaluated high frequency oscillations. Clinical neurophysiology: official journal of the International Federation of Clinical Neurophysiology 128, 433–441, https://doi.org/10.1016/j.clinph.2016.12.017 (2017).
25. Engel, J. R., Bragin, A., Staba, R. & Mody, I. High-frequency oscillations: what is normal and what is not? Epilepsy 50, 598–604, https://doi.org/10.1111/j.1528-1167.2008.01917.x (2009).
26. Braidal, M. et al. Very high frequency oscillations: Novel biomarkers of the epileptogenic zone. Annals of neurology, https://doi.org/10.1002/ana.25066 (2017).
37. Ortega, G. J., M de la Prida, L., Sola, R. G. & Pastor, J. Synchronization clusters of interictal activity in the lateral temporal cortex of
40. Wieser, H. G.
39. Sinha, N.
35. Kumar, A.
36. Antony, A. R.
41. Bagshaw, A. P., Jacobs, J., LeVan, P., Dubeau, F. & Gotman, J. Effect of sleep stage on interictal high-frequency oscillations recorded from
patients: intraoperative electrocorticographic analysis. Epilepsia
Epilepsia
81, 664–676, https://doi.org/10.1002/ana.24928 (2017).
Jehi, L. et al. Development and validation of nomograms to provide individualised predictions of seizure outcomes after epilepsy
surgery: a retrospective analysis. Lancet Neurol 14, 283–290, https://doi.org/10.1016/S1474-4422(14)70325-4 (2015).
Kumar, A. et al. Preoperative estimation of seizure control after resective surgery for the treatment of epilepsy. Seizure: the journal of the British Epilepsy Association 22, 818–826, https://doi.org/10.1016/j.seizure.2013.06.010 (2013).
Antony, A. R. et al. Functional connectivity estimated from intracranial EEG predicts surgical outcome in intractable temporal lobe epilepsy. PLoS One 8, e77796, https://doi.org/10.1371/journal.pone.0077916 (2013).
Ortega, G. J., M de la Prida, L., Sola, R. G. & Pastor, J. Synchronization clusters of interictal activity in the lateral temporal cortex of
epileptic patients: intraoperative electrocorticographic analysis. Epilepsia 49, 269–280, https://doi.org/10.1111/1528-1167.1266x (2008).
8. Coito, A. et al. Altered directed functional connectivity in temporal lobe epilepsy in the absence of interictal spikes: A high density
EEG study. Epilepsia 57, 402–411, https://doi.org/10.1111/epi.13308 (2016).
9. Sinha, N. et al. Predicting neurosurgical outcomes in focal epilepsy patients using computational modelling. Brain: a journal of
neurology 140, 319–332, https://doi.org/10.1093/brain/aww299 (2017).
Wieser, H. G. et al. ILAE Commission Report. Proposal for a new classification of outcome with respect to epileptic seizures
following epilepsy surgery. Epilepsia 42, 282–286 (2001).
11. Bagshaw, A. P., Jacobs, J., LeVan, P., Dubeau, F. & Gotman, J. Effect of sleep stage on interictal high-frequency oscillations recorded from
depth macroelectrodes in patients with focal epilepsy. Epilepsia 50, 617–628, https://doi.org/10.1111/j.1528-1167.2008.01784.x (2009).
12. Rosso, O. A. et al. Wavelet entropy: a new tool for analysis of short duration brain electrical signals. J Neurosci Methods 105, 65–75 (2001).
13. Ernst, M. D. Permutation Methods: A Basis for Exact Inference. Statist. Sin. 19, 676–685 (2004).

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Author Contributions
S.B., T.F., and E.B. analyzed data, prepared figures and tables. N.K. and T.G. treated patients and monitored
outcome. P.H. acquired data. J.S. designed and supervised the study. T.F., S.B., and J.S. wrote the article. All
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