Unsupervised Detection of High-Frequency Oscillations Using Time-Frequency Maps and Computer Vision

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High-frequency oscillations >80 Hz (HFOs) have unique features distinguishing them from spikes and artifactual components that can be well-evidenced in the time-frequency representations. We introduce an unsupervised HFO detector that uses computer-vision algorithms to detect HFO landmarks on two-dimensional (2D) time-frequency maps. To validate the detector, we introduce an analytical model of the HFO based on a sinewave having a Gaussian envelope, for which analytical equations in time-frequency space can be derived, allowing us to establish a direct correspondence between common HFO detection criteria in the time domain with the ones in the frequency domain, used by the computer-vision detection algorithm. The detector identifies potential HFO events on the time-frequency representation, which are classified as true HFOs if criteria regarding the HFO’s frequency, amplitude, and duration are met. The detector is validated on simulated HFOs according to the analytical model, in the presence of noise, with different signal-to-noise ratios (SNRs) ranging from −9 to 0 dB. The detector’s sensitivity was 0.64 at an SNR of −9 dB, 0.98 at −6 dB, and >0.99 at −3 dB and 0 dB, while its positive prediction value was >0.95, regardless of the SNR. Using the same simulation dataset, our detector is benchmarked against four previously published HFO detectors. The F-measure, a combined metric that takes into account both sensitivity and positive prediction value, was used to compare detection algorithms. Our detector surpassed the other detectors at −6, −3, and 0 dB and had the second best F-score at −9 dB SNR after the MNI detector (0.77 vs. 0.83). The ability to detect HFOs in clinical recordings has been tested on a set of 36 intracranial electroencephalogram (EEG) channels in six patients, with 89% of the detections being validated by two independent reviewers. The results demonstrate that the unsupervised detection of HFOs based on their 2D features in time-frequency maps is feasible and has a performance comparable or better than the most used HFO detectors.

Keywords: electroencephalogram (EEG), high-frequency oscillations, time-frequency maps, computer vision, signal detection
INTRODUCTION

A pioneering study by Fisher et al. (1992) on five epileptic patients implanted with subdural electrodes showed a significant increase of spectral power above 35 Hz at the beginning of epileptic seizures exhibiting electrodecremental patterns and hypothesized that high-frequency (HF) recordings may be useful in localizing the seizure onset (Fisher et al., 1992).

In the years that followed, many studies revealed that HF oscillations (HFOs) in the 80–250-Hz frequency range, also known as “ripples,” can be identified in the hippocampal and parahippocampal regions of rodents (Buzsáki et al., 1992; Ylinen et al., 1995), primates (Skaggs et al., 2007), and humans (Bragin et al., 1999; Matsumoto et al., 2013). These studies identified HFOs in healthy subjects or in epileptic patients performing visual or motor tasks (Matsumoto et al., 2013), while other studies found an increased number of HFOs in brain regions that are part of the epileptogenic network; therefore, the HFO was considered to be a potential epilepsy biomarker (Urrestarazu et al., 2007; Jacobs et al., 2009; Brázdil et al., 2010; Kerber et al., 2013; Geertsema et al., 2015). However, in spite of the various existing models of HFO generation (Fink et al., 2015; Helling et al., 2015), it is still a matter of debate how to distinguish physiological and pathological HFOs (Engel et al., 2009; Waldman et al., 2018).

Currently, the gold standard of marking HFOs is still based on visual analysis of band-pass filtered signals, as described in detail by Jacobs et al. (2009). It implies the filtering of the electroencephalogram (EEG) signal in the 80–250-Hz frequency range (“ripple” band) or 250–500 Hz (“fast ripple” band), followed by a visual search of portions of the filtered EEG signal that have “clearly visible” higher amplitudes than the rest of the signal. At least four oscillations have to be counted in each HFO (Jacobs et al., 2009; Zijlmans et al., 2017). The main drawback of this method derives from the fact that the human eye cannot accurately evaluate a complex mix of frequencies by looking at the filtered signal, as the high-amplitude low-frequency components of the spectrum may obscure low-amplitude HFOs. Moreover, it has been previously shown that interictal spikes and sharp transients, when filtered in the HFO frequency range of 80–250 Hz, may result in HFO-like oscillations, leading to false HFO detections, and it was suggested that HFOs should be identified using time-frequency analysis (Bénar et al., 2010; Bénar et al., 2010). The distinctive signature of true HFOs is considered to consist in “blobs” of more or less regular shapes that are detached from the time axis and whose centroid is located above 80 Hz (Bénar et al., 2010).

The aims of this report are to: (1) come up with a definition of HFOs in the time-frequency domain, referring to the presence of blobs having centroids located above 80 Hz, by introducing an analytical model that translates the one-dimensional (1D) time-domain definition into the 2D time-frequency domain; and (2) design a detector to identify HFO’s signature in 2D time-frequency domain using computer-vision algorithms.

Most automatic HFO detector tool kits use the time-frequency analysis solely for the visual validation of HFO candidates which have been identified using a time-domain detection method (Crépon et al., 2010; Burnos et al., 2014; Amiri et al., 2016; Fedele et al., 2016; Liu et al., 2016). Therefore, HFOs that are not part of the candidates will not be identified in the time-frequency analysis step, resulting in “misses” or “false negatives.” Visual analysis of time-frequency maps provides access to detailed features of the HFOs, allowing for a better discrimination of the true HFOs from the artifactual components (Bénar et al., 2010). We aim at duplicating this complex process of visual detection or validation of the HFOs using time-frequency maps in a computer-based image analysis algorithm, to be used as a high-performance HFO detector. In our study, we provide an unsupervised detection algorithm that combines time-frequency analysis and computer vision to replicate the “visual analysis” of the time-frequency representation. Our algorithm performs the detection directly in the time-frequency space, reducing the number of potential misses. To avoid the limitations and subjectivity of visual markings and the absence of a “ground truth” in real EEG recordings confirmed by time-frequency representations, we validate our automatic detector on simulated EEG signals that contain a mix of discrete HFO events and pink noise (Miyakoshi et al., 2013), with various signal-to-noise ratios (SNRs). We compare the performance of our detector with four previously published detection algorithms (Staba et al., 2002; Gardner et al., 2007; Crépon et al., 2010; Zelmann et al., 2012) using their implementation available in RIPPLELAB (Navarrete et al., 2016). We also validate the detector on intracranial EEG signals recorded during stereo-EEG procedures.

METHODS

Description of the Workflow for High-Frequency Oscillation Detection

The detection algorithm flowchart is shown in Figure 1 and is described in detail in the following sections.

Analytical Modeling of High-Frequency Oscillations

Simulated signals, based on a general model of HFOs superimposed on spikes or sharp transients (Urrestarazu et al., 2007), are introduced for assessing the detector’s performance.

The spikes are considered to have a Gaussian shape, characterized by the spike amplitude ($A_{spike}$) and the standard deviation ($\sigma_{spike}$). In the time domain, it can be written as:

$$\text{Spike}(t) = A_{spike} e^{-\frac{t^2}{2\sigma_{spike}^2}} \quad (1)$$

The HFO is modeled as a sinewave with a Gaussian envelope:

$$\text{HFO}(t) = A_{HFO} \cdot e^{-\frac{t^2}{2\sigma_{HFO}^2}} \cdot \cos(2\pi \cdot f_0 \cdot t) \quad (2)$$

where $A_{HFO}$ and $\sigma_{HFO}$ are the amplitude and the standard deviation of the Gaussian envelope, and $f_0$ is the frequency of the sinewave.

Combining (1) and (2), we obtain the formula in the time domain of an HFO superimposed on a spike:

$$x(t) = \text{Spike}(t) + \text{HFO}(t) \quad (3)$$
FIGURE 1 | Flowchart of detection algorithm.
In the frequency domain, the Fourier transform of the HFO’s Gaussian envelope is also a Gaussian:

$$\mathcal{F}\{e^{-\frac{t^2}{2\sigma_{HFO}^2}}\} = \sqrt{2\pi}\sigma_{HFO}e^{-2\pi^2\sigma_{HFO}^2 f^2}$$  \hfill (4)

The modulation theorem (Papoulis, 1962):

$$\mathcal{F}\{\cos(2\pi f_0 t) \cdot g(t)\}(f) = \frac{1}{2} [G(f - f_0) + G(f + f_0)]$$  \hfill (5)

allows us to derive the equation for the frequency spectrum of the HFO:

$$HFO(f) = A_{HFO}\sqrt{\frac{\pi}{2}}\sigma_{HFO}e^{-2\pi^2\sigma_{HFO}^2(f - f_0)^2}$$  \hfill (6)

The Fourier transform of the spike and HFO signal described in the time domain by Equation (3) is therefore:

$$y(t) = A_{\text{spike}}\sqrt{\frac{\pi}{2}}\sigma_{\text{spike}}e^{-2\pi^2\sigma_{\text{spike}}^2t^2}$$

$$+ A_{HFO}\sqrt{\frac{\pi}{2}}\sigma_{HFO}e^{-2\pi^2\sigma_{HFO}^2(f - f_0)^2}$$  \hfill (7)

Figure 2 visually illustrates a simulated signal based on our model (A) and its frequency spectrum (B). The spike has an amplitude of $A_{\text{spike}} = 2.5\text{ a.u.}$ and 50-ms duration at full-width half-maximum (FWHM). The HFO has a frequency of $f_0 = 100\text{ Hz}$, while its envelope has an amplitude of $A_{HFO} = 1\text{ a.u.}$, 40-ms duration, and four oscillations, both computed at FWHM. To make the model more realistic, we have added a $\Delta t = 15\text{ ms}$ delay to the HFO with respect to the spike, by replacing $t$ in Equation (2) with $t + \Delta t$. One can clearly see in both Equation (7) and its plot in Figure 2B that the spectral components of the spike are represented by a Gaussian centered on the origin, whereas that for the HFO, there is a Gaussian centered at $f_0$.

**Time-Frequency Maps**

The 1D analysis in the frequency domain alone has a number of limitations. First, in the context of EEG signal analysis and visual markings of HFOs, it has been previously shown that notch filtering (Kirácz et al., 2016) and high-pass filtering of spikes and sharp transients may produce artifacts that can be easily mistaken as HFOs (Bénar et al., 2010). A second limitation relates to the fact that low-frequency components of the spectrum have a higher amplitude than the HF components, thus restricting the visual analysis to oscillations that are close to the high-pass filtering frequency, which are shadowing higher-frequency lower-amplitude oscillations that may be also present in the EEG signal.

The most commonly used method for computing time-frequency maps is by employing a continuous wavelet transform, which is defined as:

$$x(s, \tau) = \frac{1}{\sqrt{\|\psi\|}} \int_{-\infty}^{\infty} x(t) \bar{\psi} \left( \frac{t - \tau}{s} \right) dt$$  \hfill (8)

where $\psi$ is the mother wavelet, $s$ is the scale of the mother wavelet, and $\tau$ is the translation of the mother wavelet.

The wavelet transform has been intensively used to study a wide range of neurological diseases (Ahmadlou et al., 2012a,b; Sankari et al., 2012) and can be implemented in applications such as single-unit isolation (Ortiz-Rosario et al., 2015), seizure detection (Li et al., 2016), and brain computer interfaces (Hsu, 2011).

For EEG signal analysis, the Morlet wavelet is perhaps the most commonly used and was found to be particularly useful in HFO analysis (Le Van Quyen and Bragin, 2007). As it has been successfully used in previous studies for both visual validation of manual markings (Bénar et al., 2010; Crépon et al., 2010; Alkawadri et al., 2014; Navarrete et al., 2016) and automatic HFO detectors (Zelmann et al., 2010), we decided to use it in the current study.

The Morlet mother wavelet (Figure 3A) has been initially introduced for the analysis of geoseismic signals (Goupillaud
et al., 1984) and is a particular case of the more general Gabor wavelet, defined by Equation (9), in which $2\pi \cdot f_0 = 5$ (Kumar and Foufoula-Georgiou, 1997). Le Van Quyen et al. (2001) defined the number of cycles as $n_{co} = \omega_{\sigma} = 2\pi \cdot f_0$.

\[ \psi(t) = e^{\frac{-t^2}{2\sigma^2}} \cdot \cos(2\pi \cdot f_0 \cdot t) \]  

(9)

The frequency spectrum of the Morlet wavelet is shown in Figure 3B. Within the continuous wavelet transform, the Morlet wavelets are scaled so that their envelope’s standard deviation fits the duration of various oscillations within the EEG signal.

The time-frequency representation of the analytical signal defined by Equations (1–3) and plotted in Figure 2A containing a prototype HFO superimposed on a spike is shown in Figure 3C. One has to note the appearance of the HFO as a blob that is isolated from the low-frequency components of the spike. A cross section of the time-frequency map, performed at a time corresponding to the peak of the HFO (Figure 3D), shows a spectral density in agreement with the analytical form plotted in Figure 2B. The presence of blobs and their characteristics (amplitude, centroid, FWHM) will be used in the subsequent analysis as the signature of true HFOs.

**Computer Vision for Automatic High-Frequency Oscillation Detection**

Our automatic HFO detector is based on time-frequency maps that are computed for a specified interval (1 s) of EEG signal (Figure 4A) by performing a continuous wavelet transform. Consecutive windows have a 200-ms overlap that ensures HFO candidates located at the window edges will still be detected in at least one window. A computer-vision algorithm processes each time-frequency map (Figures 4B,D) as an image, attempting to identify blobs that correspond to the presence of the HFOs. This approach aims to automate the visual analysis of time-frequency representations that would be performed by a human. So, in this context, computer vision means all processing steps performed on the time-frequency images that lead to the identification of HFO candidates.

We suppress structures that are lighter than their surroundings and that are connected to the image border (Soille, 2004) (this step is useful to discard any wavelet filtering artifacts located at the extremities of the time window and to eliminate low-frequency components of the signal). Otsu’s method allows us to compute a threshold which is further used to convert the image to black and white (Otsu, 1979). We use the black-and-white image to perform blob analysis and detection (Figures 4C,E). The blob analysis is performed using the “vision.BlobAnalysis” function in Matlab’s Computer Vision Toolbox. This function analyzes the black-and-white image and, for each white blob resulting after image binarization, computes the blob centroid coordinates, width and height in pixels. Knowing the exact dimensions of the image and the screen’s resolution, we can determine the exact location in time, frequency, and the amplitude of each detected blob in the original time-frequency representation (Figure 4F).
Detection Validation Criteria

Blobs detected in the previous step are considered HFO candidates, which are validated as HFOs using the following criteria:

1. $80 < \text{HFO}_{\text{frequency}} < 250 \text{ Hz}$ (ripples) or $250 < \text{HFO}_{\text{frequency}} < 500 \text{ Hz}$ (fast ripples)
2. $\text{HFO}_{\text{duration}} > N_{\text{osc}} \times \frac{1}{\text{HFO}_{\text{freq}}}$
3. $\text{HFO}_{\text{amplitude}} > 3.5 \times C_{\text{thresh}}$ (adapted from Alkawadri et al., 2014)

where $\text{HFO}_{\text{frequency}} \rightarrow$ blob's centroid frequency; $\text{HFO}_{\text{amplitude}} \rightarrow$ blob's centroid amplitude (wavelet coefficients); $\text{HFO}_{\text{duration}} \rightarrow$ width of the rectangle which contains the blob; $N_{\text{osc}} = 4.5 \rightarrow$ minimum number of oscillations at $\text{HFO}_{\text{frequency}}$ (adapted from Haegelen et al., 2013, and Jacobs et al., 2010); $C_{\text{thresh}} \rightarrow$
mean of wavelet coefficients in the 80–100-Hz range, computed on time intervals where the wavelet coefficients are less than the amplitude threshold.

**Image Intensity Thresholds**

The steps described in previous paragraphs only take into account one intensity threshold of the time-frequency image. As the amplitude of various frequencies within the EEG signal shows large variations, it is required to use a variable image intensity threshold to be able to detect the HFOs in the upper frequency range of the spectrum, which would otherwise be shadowed by the higher amplitudes of the lower-frequency components within the spectrum (Figures 4B,D).

Our algorithm starts with an initial threshold computed as the root mean square (RMS) of the EEG signal for the selected channel, computed during the first 15 s. This step ensures the adaptation of the algorithm to the different EEG amplitudes which are found on different channels.

The threshold is then decreased exponentially starting from an initial threshold using a geometric progression with a common ratio of 0.8:

\[
\text{thresh}_i = 0.8^{i-1} \cdot \text{thresh}_0
\]

where \( \text{thresh}_i \) is the \( i \)-th term of the geometric progression and \( \text{thresh}_0 \) is the initial threshold, computed as the RMS of the first 15 s of the EEG signal.

For each image intensity threshold, we repeat the previously described steps. For the decreasing thresholds, the detected HFOs are considered identical and not marked if the rectangle of the newly detected HFO already contains the centroid of a previously detected HFO. In this way, we avoid false large markings which may occur due to the fact that two nearby blobs may become one larger blob at lower thresholds.

The processing steps described above were implemented in Matlab R2016a using the “vision.BlobAnalysis” function in the Computer Vision Toolbox. As the Computer Vision Toolbox requires a license, we have provided a compiled executable version of our detector at https://bitbucket.org/cristidonos/hfodetector. The algorithm performs a detection on a 1-s window signal at 15 different image intensity thresholds in ~1.5–2 s on a laptop (Asus Q324U, Intel® Core™ i7 7500U CPU, 16GB RAM).

**Simulation Data**

Three hundred HFO events were simulated using the previously described formula [Equation (2)]. The HFO amplitude was normalized to 1, while the frequency was randomly distributed in the 80–250-Hz range. The HFO duration at FWHM was frequency dependent, so that it will be 5, 6, or 7 oscillations long at the HFO frequency in the simulation dataset. We will refer to the 300 s continuous signal having 2 kHz sampling rate and one HFO every second as our simulated HFO signal.

The choice of pink noise was motivated by the resemblance with the frequency spectrum of real EEG recordings, as well as their previous usage in previously published HFO detection studies (Miyakoshi et al., 2013). Three hundred seconds of pink noise with a sampling rate of 2 kHz were generated. The amplitude of the noise signal was normalized to ±1 au. We will further refer to this signal as our simulated noise signal.

Four additional signals were obtained by performing a weighted summation of the simulated HFO and the simulated noise signals so that constant SNRs of −9, −6, −3, and 0 dB were obtained. The combined signal (\( \text{Sig}_{\text{combined}} \)) was obtained by multiplying the pink noise (\( \text{noise} \)) by a factor \( f \) so that the desired SNR (\( \text{SNR}_{\text{dB}} \)) is obtained, according to the formula:

\[
\text{Sig}_{\text{combined}} = \text{Sig}_{\text{HFO}} + f \cdot \text{noise}
\]

where \( f = 2^{\frac{\text{SNR}_{\text{dB}}}{10}} \cdot \frac{\text{RMS}(\text{Sig}_{\text{HFO}})}{\text{RMS}(\text{noise})} \) and \( \text{SNR}_{\text{dB}} \in \{-9; -6; -3; 0\} \).

The simulation dataset and a Matlab GUI of our detection software are made available for testing at https://bitbucket.org/cristidonos/hfodetector.

**Comparison With Other High-Frequency Oscillation Detectors**

Four previously published automatic detectors were used to perform HFO detection on the same simulation dataset using their implementation in RIPPLELAB (Navarrete et al., 2016). As named in RIPPLELAB, the four detectors are Hilbert (HIL) (Crépon et al., 2010), MNI detector (MNI) (Zelmann et al., 2010), Short Time Energy (STE) (Staba et al., 2002), and Short Line Length (SLL) (Gardner et al., 2007). For all four detectors in RIPPLELAB, we set the frequency range to 80–250 Hz and the epoch duration to 300 s. Two additional parameters were tuned for MNI and STE, and we will discuss the reasoning in the Results section. All remaining parameters were used with their default values, recommended by the original papers describing the algorithms and successfully used for similar comparisons on manually annotated EEG data (Zelmann et al., 2010; Navarrete et al., 2016; Roehri et al., 2017). For all detectors, we computed the number of true (TPs) and false positives (FPs), the positive prediction value (PPV), the sensitivity (Sens), and the F-measure. F-measure is defined as \( F = \frac{2 \cdot \text{PPV} \cdot \text{Sens}}{\text{PPV} + \text{Sens}} \) (or \( \frac{TP}{TP + FP} \)), where FN is the number of false negatives.

**Intracranial Electroencephalogram Data**

The detection algorithm’s performance was also evaluated on intracranial EEG data recorded during slow-wave sleep (Jacobs et al., 2008; Alkawadri et al., 2014) in six patients undergoing stereo-EEG presurgical evaluation for drug-resistant epilepsy in our epilepsy center using the methodology described in Donos et al. (2017). For each patient, we selected a total of six bipolar EEG traces recorded on three contacts located in epileptogenic areas and three contacts located in non-epileptogenic areas, and we performed automatic detection of HFOs on a 5-min EEG segment. The detections were visually inspected independently by two experienced reviewers (CD, AB) based on a combined display of the raw EEG trace, the 80–300-Hz filtered EEG trace, and the time-frequency representation of the signal. Detections that did not meet the standard inclusion criteria (Jacobs et al., 2009, 2012; Zijlmans et al., 2017), including the appearance as blob separated from the time axis on time-frequency maps (Bénar
et al., 2010; Roehri et al., 2017) were rejected as FPs. The inter-rater agreement was quantified by Cohen’s kappa coefficient (Cohen, 1960).

RESULTS
Detector Evaluation
The automatic detection algorithm was run on the simulation dataset. Figure 5 shows a simulated HFO at 88 Hz, superimposed on the noise signal with SNRs of 0 (Figure 5A), −3 (Figure 5B), −6 (Figure 5C), and −9 dB (Figure 5D). The corresponding time-frequency representations are provided in Figures 5E–H.

If multiple candidate events were detected within some limits of tolerance in both time (50 ms) and frequency (5 Hz), they were merged and considered as a single event. Detections within the tolerance limits were compared to the known time and frequency location of each simulated HFO and were considered TPs, while other detections were considered FPs. Whenever multiple detections within the tolerance limits occurred due to detection window overlap and the associated baseline changes, they were considered as unique events.

For our detector, the number of TPs ranged between 193 at −9 dB SNR and 299 at −6, −3, and 0 dB SNRs for the mixed simulated HFOs and noise signals. A small number of FPs were detected at all SNRs, with a maximum of 15 FPs at −6 dB SNR. Figure 6 shows the time and frequency distribution of TPs and FPs for all SNRs pooled together. Our detector successfully detected simulated HFOs that cover the whole 80–250-Hz frequency range. The time and frequency localization errors of each simulated HFO were computed as the difference between the ground truth simulated values and the detection values. The average time localization errors and the average frequency localization errors had similar values regardless of the SNR: ~5 ms and ~0.8 Hz, respectively (Table 1).

The PPVs were above 0.95 for all SNRs. Sensitivity and F-measure were 0.64 and 0.77 at −9 dB SNR, but they both increased to more than 0.97 for the remaining SNRs.

The FPs were either simulated HFOs detected outside the time and frequency tolerance limits, either random oscillations in the pink noise that matched the HFO detection criteria by chance.

The performance metrics of our detector are shown in detail in Table 1 and are compared to four other detection algorithms implemented in RIPPLELAB in Figure 7. Results in Table 1 and Figure 7 are obtained for simulation dataset.

The detector is able to detect HFO events with virtually any frequency by adjusting the frequency criterion (1) described in the Detection Validation Criteria section. In the Supplementary Material, we have validated the detector with an additional dataset of simulated HFOs in the fast ripple band (250–500 Hz), and we obtained similar sensitivity results. The ability to detect even higher-frequency events can be accommodated by adjusting the time and frequency intervals, as well as the resolution of the time-frequency maps, such that the blobs associated with HFOs having the minimal number of oscillations will have a minimal area that will allow the computer-vision algorithm to properly operate.

Comparison With Other Automatic Detectors
We used the same evaluation scheme as described in Detector Evaluation based on comparing detections to the ground truth of the simulated HFO parameters.

MNI and STE almost failed to detect HFOs with their default parameters. The MNI detector had the initial baseline duration set to 125 s, and using this value, the detection sensitivity and F-measure were <0.05 at all SNRs. After decreasing the baseline duration to 15 s, the detection greatly improved. STE requires
FIGURE 6 | Distribution of high-frequency oscillation (HFO) detections in time and frequency for the simulation dataset. True positive (TP) and false positive (FP) are represented in blue and orange in the scatterplot and in the histograms.

TABLE 1 | Detection performance of our detector on the simulation dataset across various SNRs.

| SNR | −9 dB | −6 dB | −3 dB | 0 dB | HFOs only | Noise only |
|-----|-------|-------|-------|------|-----------|-----------|
| Number of simulated HFOs | 300  | 300  | 300  | 300  | 300       | 0         |
| TP  | 193   | 295   | 299   | 299  | 299       | 0         |
| FP  | 7     | 15    | 6     | 2    | 0         | 7         |
| PPV | 0.965 | 0.952 | 0.980 | 0.993| 1.000     | 0         |
| Sensitivity | 0.643 | 0.983 | 0.997 | 0.987 | 0.997 | 0 |
| F-measure | 0.772 | 0.967 | 0.988 | 0.986 | 0.997 | 0 |
| Mean time localization error | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | - |
| SD time localization error | 0.004 | 0.003 | 0.003 | 0.003 | 0.003 | - |
| Mean frequency localization error | −0.787| −0.832| −0.824| −0.824 | −0.824 | - |
| SD frequency localization error | 1.281 | 1.162 | 1.156 | 1.156 | 1.156 | - |

FP, false positive; HFO, high-frequency oscillation; PPV, positive prediction value; SNR, signal-to-noise ratio; TP, true positive.

explicit definition of the minimum number of oscillations for HFO events. Since all our simulated HFOs had five, six, or seven oscillations, we set the minimum number of oscillations to 4.5, the same value we use for our detector. Additionally, the MNI detector did not detect any HFOs in the absence of noise (SimHFOs). A possible explanation is that the MNI detector relies on a baseline period for detection, which in the absence of noise was maybe too similar to the HFO simulations which were targeted for detection.

Our detector and MNI surpassed all other detectors in terms of sensitivity, F-measure, and number of TP at all SNRs. However, MNI performed slightly better at −9 dB in terms of TPs (234 vs. 193) but also had a significantly larger number of FPs, ranging between 28 vs. 7 at −9 dB and 34 vs. 2 at 0 dB. Consequently, our detector reached higher PPVs at all SNRs and higher F-measures for −6, −3, and 0 dB.

The HIL and SLL detectors had almost similar TP and sensitivities, but overall, we would say that SLL performed better due to the fact that SLL had a sensitivity of 0.44 at −9 dB SNR, which increased to 0.69 at 0 dB SNR. HIL had 0 sensitivity at −9 dB but quickly increased to 0.07, 0.5, and 0.77 at −6, −3, and 0 dB, respectively. Similar increase trend was observed for F-measure: SLL was more consistent across various SNRs, but HIL reached 0.87 at 0 dB SNR. STE had its highest sensitivity of 0.29 and its highest F-measure of 0.17 at 0 dB. While investigating the causes of such sensitivities, we observed that STE’s performance can be improved by reducing the detection threshold from 5 to 2.5 standard deviations, represented by the “RMS SD_threshold” field in RippleLab. However, for the purpose of our simulation, it is important to use the detection parameters already optimized and validated in real EEG data by the authors of each detection algorithm to be able to properly compare performances between various methods.

Our detector had a few FPs at all SNRs, reaching a maximum of 15 at −6 dB. MNI had 28 FPs at −9 and −6 dB SNR. MNI had 34 FPs at −3 and 0 dB, and in the noise only signal. STE had only 34 FPs in the noise only signal. In other cases, no more than 1–2 FPs were detected, regardless of the SNRs. As a consequence, all detectors have a PPV equal or very close to 1.

Performance on Real Electroencephalogram Data

A total of 173 HFOs were detected across all 36 EEG channels recorded intracranially in six patients diagnosed with
FIGURE 7 | Comparison of detection performance in the simulation dataset. SimHFOs is the signal containing only simulated high-frequency oscillation (HFO), while SimNoise is the signal containing only the pink noise.
FIGURE 8 | High-frequency oscillation (HFO) with a particular morphology, simultaneous with an epileptic spike, detected on intracranial electroencephalogram (EEG). The panels show raw EEG signal (A), filtered EEG in 80–500-Hz frequency range (B), and the time-frequency representation (C).

DISCUSSION

The HFOs are thought to be a correlate of tissue epileptogenicity (Urrestarazu et al., 2007; Jacobs et al., 2009, 2012; Brázdil et al., 2010; Kerber et al., 2013). Being considered a potential biomarker of epileptogenic tissue, HFOs drew a lot of attention from the scientific community. Based on the gold standard of HFO identification, which is the band-pass filtering combined with visual analysis, a large number of automatic HFO detectors emerged (Staba et al., 2002; Gardner et al., 2007; Crépon et al., 2010; Zelmann et al., 2010; Dümpelmann et al., 2012; Gliske et al., 2015). However, all of the detectors mentioned above rely, at least to some extent, on the optimization of the detection against some visually marked HFOs. The subjectivity of visual markings, quantified by the Cohen's kappa coefficient (Cohen, 1960), is reflected by the direct comparison of four automatic detectors (Zelmann et al., 2012), a study in which the median kappa coefficient varied in the 0.05–0.97 range for the four detectors. Indeed, this limitation is also acknowledged by Dümpelmann et al. (2012), which shows that the automatically detected HFOs partially overlap with the visual markings, but nevertheless, the overall HFO rates across channels reflect the patient's epileptogenicity. Considering all of the above, we believe...
that a standardized model of HFOs is much needed, a model
which can be described through mathematical equations and that
can be further used to validate automatic detection algorithms.
In this study, we provide such an analytical model of the HFO,
together with a practical way of identifying the HFOs from real
EEG data in an unsupervised manner.

The use of simulated HFOs, whose time localization, duration,
and frequency are known, considered to represent the ground
truth for evaluating our detection algorithm allows us to
overcome the layer of uncertainty caused by the subjectivity of
manual HFO markings in real EEG data, for which the ground
truth is unknown. Our algorithm showed high sensitivity and
F-measure across various SNRs, ranging from 0.64 and 0.77 at
−9 dB to 0.997 and 0.995 at 0 dB, respectively. This is a clear
proof that our detection algorithm is able to find the HFOs that
match the HFO guidelines in terms of frequency and number
of oscillations, provided by previous studies (Jacobs et al., 2010;
Alkawadri et al., 2014). As a second validation step, we used
our HFO detector on intracranial data recorded from patients
undergoing stereo-EEG presurgical evaluation for drug-resistant
epilepsy. Detections were visually validated by two independent
expert reviewers, resulting in 89% of detections being visually
confirmed as HFOs. We relied on the fact that our detector had
an insignificant number of FPs on simulated data and assumed
a similar number of FPs on real EEG data as well. This was
indirectly confirmed by the visual validation of the detections and
the HFO rates, which were 1.1 events per minute, in agreement
other studies (Alkawadri et al., 2014).

The direct comparison on the same simulated dataset between
our detector and four other automatic detectors showed that our
method provides better sensitivities and F-measures, which are
robust in respect to SNR changes. Interestingly, the STE and
MNI detectors failed to generally detect the simulated HFOs
with their default parameters, which were optimized based on
visual markings of real EEG data. While their performance on
simulated data was improved by changing some of their
parameters, it is very likely that such changes will affect their
performance on real EEG. The remaining algorithms, HIL and
SLL detectors, had moderate performance across all SNRs.

An interesting observation is that the simulated pink noise
contains some “HFO-like” events, which are within the HFO
frequency range and meet the other detection criteria defined in
the Methods section. The presence of HFO-like events in noise,
without electrophysiological substrate, may explain to some
extent the modest inter-rater agreements and modest sensitivities
that were previously reported in HFO studies (Zelmann et al.,
2010; D Humphelmann et al., 2012; Spring et al., 2017).

The four detectors that we compare ours with use indirect
ways of detecting and quantifying oscillations in the time domain
(line length) or across a frequency range which is biased toward
its low edge of the spectrum. These detectors do not heavily
rely on the mathematical properties of oscillations, instead, they
eventually use empirically derived features tuned to maximize the
match between the detections and the visual markings of HFO.
A new dataset of simulated HFOs, generated by an innovative
method combining an autoregressive model for simulating EEG
and discrete wavelet transforms for extracting HFOs from visual
markings, was made available by Roehri et al. (2017). However,
this dataset also lacks the information about HFO frequencies.
Nevertheless, this simulation dataset is used for comparing
Delphos, a new detection algorithm with the four other detection
algorithms in RIPPLELAB. While such detection algorithms
proved to be useful in delineating the epileptogenic areas (Jacobs
et al., 2008, 2009; Zeljms et al., 2011; Burnos et al., 2014
Fedele et al., 2016), it is still a matter debate how to differentiate
pathological from physiological HFOs (Engel et al., 2009; Jefferys
et al., 2012; Fink et al., 2015). Using our automatic detector,
the HFO analysis and the quest for separating pathological and
physiological HFOs can extend to another dimension, by using
the frequency information.

In agreement with a previous study (Roehri et al., 2017),
we conclude that based on our simulated data, optimizing an
HFO detector to visual markings is not enough to ensure a
robust detection across various SNRs. Paradoxically, detectors
that generally failed to detect true oscillations, matching the
HFO detection criteria on our simulated dataset, are successful
in localizing the epileptogenic focus in real EEG data (Staba et
al., 2002; Zelmann et al., 2010). The fact that visually marked
(detected by automatic detectors tuned to visual markings) HFOs
are positively correlated to epileptogenic areas may be explained
by the high number of other types of interictal events, like spikes
and sharp-waves (Engel and da Silva, 2012; Staba, 2012; Zeljms
et al., 2012). These events, known to be associated with the
epileptogenic cortex, have an HF content that may be mistaken as
HFOs in the filtered signal (Bénar et al., 2010; Amiri et al., 2016).
Our unsupervised detector operates directly on time-frequency
representations of the EEG signal and looks for “blobs” in the
time-frequency space, an HFO landmark, as shown by Bénar et
al. (2010). Although the HFO model discussed in this paper is a
simplified one, based on HFO “atoms,” our detection algorithm
generalizes very well to complex real EEG data, in which HFOs
often appear on time-frequency representations as blobs with
irregular shape. Figure 8 shows an HFO superimposed on an
epileptic spike, centered at 237 Hz, with a duration of 18 ms, that
has a particular morphology: it starts at a frequency above 300 Hz
and decreases to roughly 220 Hz over ~30 ms. Studying the
specific morphology of HFOs may be important in distinguishing
pathological from physiological HFOs, although a recent study
on a limited subset of four morphological types demonstrated a
limited value in delineating the epileptogenic zone (Burnos et al.,
2016).

The extent to which HFOs detected on a time-frequency
map are correlated with the epileptogenic areas remains to
be determined by future clinical studies. In the meantime,
we provide additional evidence that there is an inconsistency
between the gold standard HFO detection criteria in time-
amplitude domain and the actual signal characteristics in time-
frequency domain. We come to the conclusion, just like Roehri
et al. (2017), that HFO detection criteria need to be revisited by
taking into consideration both time and frequency characteristics
of HFOs.

**CONCLUSIONS**

We provide a method for the automatic detection of HFOs
that uses a computer-vision approach for detecting the
HFO landmarks on time-frequency maps. The algorithm was validated using simulated data, based on an analytical model of the HFOs, as well as intracranial EEG. A mathematical model in time and frequency domain provides a justification of “blobs” unambiguously identifying true HFOs in time-frequency maps. Our detection algorithm was proven to have a high F-measure over a wide range of SNRs.

DATA AVAILABILITY STATEMENT

The datasets generated for this study are available on request to the corresponding author.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by Bagdasar Arseni Hospital Ethical Committee approval No 2621/03.02.2012. The patients/participants provided their written informed consent to participate in this study.

REFERENCES

Ahmadlou, M., Adeli, H., and Adeli, A. (2012a). Fractal analysis of frontal brain in major depressive disorder. Int. J. Psychophysiol. 85, 206–211. doi: 10.1016/j.ijpsycho.2012.05.001
Ahmadlou, M., Adeli, H., and Adeli, A. (2012b). Fuzzy synchronization likelihood-wavelet methodology for diagnosis of autism spectrum disorder. J. Neurosci. Methods 211, 203–209. doi: 10.1016/j.jneumeth.2012.08.020
Alkawadri, R., Gaspard, N., Gonchorova, I. I., Spencer, D. D., Gerrard, J. L., Zaveri, M., et al. (2012b). Fuzzy synchronization likelihood of SNRs. J. Neurophysiol. 108, 137–142. doi: 10.1152/jn.00872.2011
Amiri, M., Lina, J.-M., Pizzo, F., and Gotman, J. (2016). High frequency oscillations and spikes: separating real HFOs from false oscillations. Clin. Neurophysiol. 127, 187–196. doi: 10.1016/j.clinph.2015.04.290
Bénar, C. G., Chauvrière, L., Bartolomei, F., and Wendling, F. (2010). Pitfalls of high-pass filtering for detecting epileptic oscillations: a technical note on “false” ripples. Clin. Neurophysiol. 121, 301–310. doi: 10.1016/j.clinph.2009.10.019
Bragin, A., Engel, J., Wilson, C. L., Fried, I., and Buzsáki, G. (1999). High-frequency oscillations in human brain. Hippocampus 9, 137–142. doi: 10.1002/(SICI)1098-1063(1999)9:2<137::AID-HIPO5>3.0.CO;2-0
Brázdil, M., Halámek, J., Jurák, P., Daniel, P., Kuba, R., Chrastina, J., et al. (2014). Human intracranial high frequency oscillations during epilepsy surgery predicts seizure outcome. Clin. Neurophysiol. 127, 3066–3074. doi: 10.1016/j.clinph.2016.06.009
Fink, C. G., Giske, S., Catoni, N., and Stacey, W. C. (2015). Network mechanisms generating abnormal and normal hippocampal high-frequency oscillations: a computational analysis. eNeuro. 2:ENEURO.0024-15.2015
Donos, C., Mindruta, I., Maliia, M. D., Rasina, A., Ciurea, J., and Barborica, A. (2017). Co-occurrence of high-frequency oscillations and delayed responses evoked by intracranial electrical stimulation in stereo-EEG studies. Clin. Neurophysiol. 128, 1043–1052. doi: 10.1016/j.clinph.2016.11.028
Dümpelmann, M., Jacobs, I., Kerber, K. and Schulze-Bonhage, A. (2012). Automatic 20–250 Hz “ripple” high frequency oscillation detection in invasive subdural grid and strip recordings in epilepsy by a radial basis function neural network. Clin. Neurophysiol. 123, 1721–1731. doi: 10.1016/j.clinph.2012.02.072
Engel, J., Bragin, A., Staba, R., and Mody, I. (2009). High-frequency oscillations: what is normal and what is not? Epilepsia 50, 598–604. doi: 10.1111/j.1528-1167.2008.01917.x
Engel, J., and da Silva, F. L. (2012). High-frequency oscillations - where we are and where we need to go. Prog. Neurobiol. 98, 316–318. doi: 10.1016/j.pneurobio.2012.02.001
Fedele, T., van ’t Klooster, M., Burnos, S., Zweiphenning, W., van Klink, N., Leijten, F., et al. (2016). Automatic detection of high frequency oscillations during epilepsy. J. Clin. Neurophysiol. 23, 441–448. doi: 10.1016/j.jclinph.2016.01.002
Gardner, A. B., Worrell, G. A., Marsh, E., Dlugos, D., and Litt, B. (2007). Human and automated detection of high-frequency oscillations in clinical intracranial EEG recordings. Clin. Neurophysiol. 118, 1334–1343. doi: 10.1016/j.clinph.2006.12.019
Geesits, E. E., Visser, G. H., Velis, D. N., Claus, S. P., Zijlmans, M., and Kalitzin, S. N. (2015). Automated seizure onset zone approximation based on nonharmonic high-frequency oscillations in human interictal intracranial EEG. Int. J. Neural Syst. 25:1550015. doi: 10.1142/S012906571550015X
Groves, S. V., Irwin, Z. T., Davis, K. A., Sahaya, K., Chessek, C., and Stacey, W. C. (2015). Universal automated high frequency oscillation detector for real-time, long term EEG. Clin. Neurophysiol. 127, 1057–1066. doi: 10.1016/j.clinph.2015.07.016

AUTHOR CONTRIBUTIONS

CD and AB designed the study and performed the analysis. AB wrote the Matlab scripts related to the HFO model and its analysis, while CD wrote the Matlab software related to the HFO detection and its performance analysis. IM provided the clinical data. CD wrote the first draft of the manuscript. All authors contributed to manuscript revision, read, and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fnins.2020.00183/full#supplementary-material
Goupilaud, P., Grossmann, A., and Morlet, J. (1984). Cycle-octave and related transforms in seismic signal analysis. Geophysics 23, 85–102. doi: 10.1190/1.1441761

Haegele, C., Perucca, P., Châtilion, C. E., Andrade-Valença, L., Zelmann, R., Jacobs, J., et al. (2013). High-frequency oscillations, extent of surgical resection, and surgical outcome in drug-resistant focal epilepsy. Epilepsy 54, 848–857. doi: 10.1111/epi.12075

Helling, R. M., Koppert, M. M. J., Visser, G. H., and Kalitzin, S. N. (2015). Gap junctions as common cause of high-frequency oscillations and epileptic seizures in a computational cascade of neuronal mass and compartmental modeling. Int. J. Neural Syst. 25, 15050021. doi: 10.1142/S0129065715500215

Hsu, W.-Y. (2011). Continuous EEG signal analysis for asynchronous BCI application. Int. J. Neural Syst. 21, 335–350. doi: 10.1142/S0129065710002870

Jacobs, J., LeVan, P., Châtilion, C.-E., Olivier, A., Dubeau, F., and Gotman, J. (2009). High frequency oscillations in intracranial EEGs mark epileptogenicity rather than lesion type. Brain 132, 1022–1037. doi: 10.1093/brain/awn351

Jacobs, J., LeVan, P., Chander, R., Hall, J., Dubeau, F., and Gotman, J. (2008). Intercultural high-frequency oscillations (80-500 Hz) are an indicator of seizure onset areas independent of spikes in the human epileptic brain. Epilepsia 49, 1893–1907. doi: 10.1111/j.1528-1167.2008.01656.x

Jacobs, J., Staba, R., Asano, E., Otsubo, H., Wu, J. Y., Zijlman, M., et al. (2012). High-frequency oscillations (HFOs) in clinical epilepsy. Prog. Neurobiol. 98, 302–315. doi: 10.1016/j.pneurobio.2012.03.001

Jacobs, J., Zijlman, M., Zelmann, R., Châtilion, C.-E., Hall, J., Olivier, A., et al. (2010). High-frequency electroencephalographic oscillations correlate with outcome of epilepsy surgery. Ann. Neurol. 67, 209–220. doi: 10.1002/ana.21847

Jefferys, J. G. R., Menendez de la Prida, L., Wendling, F., Bragin, A., Avoli, M., Timoeve, I., et al. (2012). Mechanisms of physiological and epileptic HFO generation. Prog. Neurobiol. 98, 250–264. doi: 10.1016/j.pneurobio.2012.02.005

Kerber, K., LeVan, P., Dümpelmann, M., Fauser, S., Korinthenberg, R., Schulze-Bonhage, A., et al. (2013). High frequency oscillations mirror disease activity in patients with focal cortical dysplasia. Epilepsia 54, 1428–1436. doi: 10.1111/epi.12262

Kirac, L. B., Vollmar, C., Rémi, J., Loesch, A. M., and Noachtar, S. (2016). Notch filter artefact mimicking high frequency oscillation in epilepsy. Clin. Neurophysiol. 127, 979–981. doi: 10.1016/j.clinph.2015.05.030

Kumar, P., and Foufoula-Georgiou, E. (1997). Wavelet analysis for geophysical applications. Rev. Geophys. 35, 385–412. doi: 10.1029/97RG00427

Le Van Quyen, M., and Bragin, A. (2007). Analysis of dynamic brain oscillations: methodological advances. Trends Neurosci. 30, 365–373. doi: 10.1016/j.tins.2007.05.006

Le Van Quyen, M., Foucher, J., Lachaux, J.-P., Rodriguez, E., Lutz, A., Martinerie, J., et al. (2001). Comparison of Hilbert transform and wavelet methods for the analysis of neuronal synchrony. J. Neurosci. Methods 111, 83–98. doi: 10.1016/S0165-0270(01)00372-7

Li, J., Zhou, W., Yuan, S., Zhang, Y., Li, C., and Wu, Q. (2016). An improved method for the topographical identification and quantification of high frequency oscillations in human epileptic hippocampus and entorhinal cortex. J. Neurophysiol. 98, 1743–1752. doi: 10.1152/jn.2002.88.4.1743

Unsupervised Detection of HFOs

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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