

Epilepsy Detection from Multi-Channel EEG Using Cross-Recurrence Quantification Analysis and Machine Learning

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1 Introduction

Epilepsy is a neurological disorder, which generates recurrent, unprovoked seizures. Seizures are the result of abnormal neuronal brain activity which subsequently lead into disturbances in the behavior, sensation, the consciousness and the movement of the affected subject. Information gathered by World Health Organization (WHO), report that epilepsy affects around 50 million people worldwide, placing it as one of the most common neurological conditions globally.

Epilepsy has its origin in diverse and various causes depending on many factors. The generation of epileptic activity can be an impact of identifiable structural, genetic, infectious, metabolic, or immune-related abnormalities, while for almost 50% of the cases, their trigger still remains unknown[53]. Depending on the brain regions involved, a seizure can have several classifications. It can be classified as focal (if its origin is a specific area) or a generalized one (involving both hemispheres). Clinically, seizures are characterized by high variability, from short term lapses in awareness, up to convulsive episodes, and the fact of their unpredictable occurrence has severe impact in patient's quality of life.

For the diagnosis and monitoring of epilepsy, electroencephalography (EEG) is widely utilized, as it offers a non invasive technique for recording brain's electrical activity using surface electrodes. EEG signals contain temporal information that reflects the dynamic interactions of neuronal populations. During the seizure events, the presence of characteristic patterns such as spikes, sharp waves, or rhythmic discharges often appear, distinguishing this kind of activity from the normal background rhythms. As a result, EEG analysis holds a central role both in clinical diagnosis and for research related to this application domain.

Recent advances in signal processing combined with machine learning have greatly improved the ability EEG data analysis. Techniques like time-frequency decomposition, nonlinear dynamics, recurrence analysis, and deep neural networks can offer new solutions for automated extraction of complex spatial and temporal features from EEG recordings. The ultimate goal of these approaches is to support clinicians by providing objective, data-driven tools for seizure detection, prediction, and classification, in order to contribute to better patient care and personal treatments.

In summary, epilepsy represents a major public health challenge due to its prevalence, variability, and social consequences. Understanding the electrophysiological mechanisms that drive seizure generation and developing reliable methods for automatic EEG analysis remains a crucial research direction in modern neuroscience and biomedical engineering.

Recurrence Quantification Analysis (RQA) and Cross-Recurrence Quantification Analysis (CRQA) are nonlinear methods for the analysis of nonstationary time series, such as EEG signals. They offer the quantification of the recurring patterns in phase space trajectories [19, 20]. Introduced by Trulla et al.[19] (directly built on quantifying recurrence plots[24]) and expanded by Webber and Zbilut[20], RQA measures metrics like recurrence rate, determinism, and laminarity to capture dynamic system behavior. Thomasson et al.[21] in their work, demonstrated RQA's applicability on EEG data, mentioning the robustness it shows in accordance to noise and nonstationarity. Marwan et al.[22] further advanced recurrence plot techniques, emphasizing on developing a confidence measure of RQA in detecting dynamic transitions. Works like these, serve as a foundation of applying RQA and CRQA on EEG studies in various conditions such as epilepsy, cognitive disorders and others. XXX

2 Related Work

Frolov et al.[1] proposed an approach to analyze frequency based multiplex brain networks using recurrence quantification analysis (RQA) on EEG data, and demonstrated the way that recurrence-based synchronization indices can effectively capture both within-frequency (intralayer) and cross-frequency (interlayer) functional connectivity during cognitive tasks. Their work showed that RQA is particularly suitable for analyzing non-stationary EEG signals and revealed important insights about the evolution of functional connectivity patterns during cognitive tasks. In addition the dataset used in this research are openly available in a Figshare repository.

Núñez et al. [16] worked with resting-state EEG recordings from subjects with mild cognitive impairment(MCI), Alzheimer’s disease(AD), and healthy ground truth controls in order to detect frequency based changes into their brain dynamics. By blending wavelet based Kullback–Leibler divergence (KLD) for capturing non-stationarity, and two RQA metrics(*entropy of the recurrence point density* and the *median of the recurrence point density*) insights have been extracted related to neurodegeneration presence. Research’s findings show that MCI and AD are presenting notable changes in the recurrence structure and non-stationarity of EEG signals, and more specifics on the theta and beta frequency bands. Therefore, recurrence based dynamics show a capability as potential biomarkers for monitoring and detecting early Alzheimer’s disease and its progression.

MCI has also investigated by Timothy et al.[25], where researchers have focused on the classification of MCI using EEG signals and combining RQA and CRQA methods. Analysis has been performed on both resting-state (eyes closed) and task-based (short-term memory) EEG data, focusing on complexity (via RQA) and synchronization (via CRQA) features. Their results indicate that MCI patients exhibit lower complexity and higher inter- and intra-hemispheric synchronization compared to healthy controls, particularly during memory tasks. The study also proposes a novel feature space approach using RQA and CRQA measures, achieving high classification accuracy (91.7%) under task conditions.

Fan and Chou [15] have also proposed an approach for real-time epileptic seizure detection using as a method the analysis of temporal synchronization patterns of EEG signals with recurrence networks and spectral graph theory. Recurrence plots were used for the modeling of the EEG dynamics, extracting graph theory’s features for quantifying the synchronization. Results showed high sensitivity of 98.48% and low latency (6 seconds) for detecting seizure on the CHB-MIT dataset, performing better than other RQA measures.

Heunis and co-authors[23] have utilized resting state EEG and RQA in order to distinguish individuals of ages 0-18 of two categories; ASD(autism spectrum disorder) and typically developing. RQA features were extracted and tested on various linear and nonlinear classifiers achieving 92.9% classification accuracy with nonlinear SVM classifier.

Author in [6], investigated changes related to aging in brain sensorimotor systems using RQA and theta-band functional connectivity in EEG signals. In the study a VR experimental paradigm was utilized with auditory stimulus across different age groups(young and elder subjects). Key findings include that elder subjects present decreased EEG complexity during motor preparation stages as measured by RQA metrics (ΔRR and ΔRTE), and had increased theta band functional connectivity highlighting the potential of RQA in detecting age related biomarkers that were not detectable using standalone signal spectral analysis.

Guglielmo et al. [7] utilized RQA features extracted by EEG signals for the purpose of classification of cognitive performance during mental arithmetic tasks. They used frontal and parietal EEG signals and analyzed them, from 36 participants by extracting six RQA metrics (*recurrence*

rate, determinism, laminarity, entropy, maximum diagonal line length and average diagonal line length) from four electrodes (F7, Pz, P4, Fp1). Afterwards by applying machine learning classifiers (SVM, Random Forest, and Gradient Boosting) and they reached accuracy of classification above 0.85, showing the potential that RQA holds for generalizing on nonlinear dynamics.

Mihajlović [33] studied the discriminative efficiency of traditional spectral features in comparison to RQA-derived nonlinear metrics for the cognitive effort classification purposes. Utilizing a 4-channel wearable EEG headset, data was recorded while subjects perform tasks having variable cognitive load such as relaxation, math, reading. The key finding was that while spectral features alone often yielded higher classification accuracy, RQA features such *recurrence rate, determinism ratio* were consistently ranked among the most important features for discrimination task. A conjunction of a hybrid model using both spectral and RQA features achieved the best overall performance, showing the complementary nature of the methods in brain dynamics exploration.

Yang and co-authors [17], examined stereo electroencephalography (sEEG) recordings of 10 patients with refractory focal epilepsy for analyzing dynamical differences among discrete epileptic phases/states (inter-ictal, pre-ictal, and ictal) and regions. Using recurrence plots and CRQA, they identified epileptogenic channels with longer diagonal structures in RPs, which is a sign of more deterministic and recurrent dynamics. Their findings point out that the synchronization among the epileptogenic channels strengthened while seizures events occur, suggesting that these regions dominate the network's dynamics.

Lopes et al. [8] have proposed a combinatorial framework by mixing RQA with dynamic functional network (dFN) analysis, applying it to both MEG and stereo EEG data. The methodology they described is split into five steps: data segmentation, functional network inference, distance computation alongside networks, recurrence plot construction and finally RQA. The study demonstrated that functional networks in epilepsy patients recur more quickly than in healthy controls, suggesting RQA on dFNs could play the role of a potential biomarker. For the EEG dataset investigation, they have showed that the pre-ictal networks shown higher recurrence rates than post-ictal periods, with the τ -recurrence rate (RR_τ) proving particularly effective for seizure detection.

Rangaprakash [18] have proposed an application of RQA for the study of brain connectivity using multichannel EEG signals. In its work, a new CRQA-based feature was proposed (Correlation between Probabilities of Recurrence (CPR)), a nonlinear and non-parametric phase synchronization technique. Afterwards it was utilized for the analysis of functional connectivity in epilepsy subjects during eyes-open/eyes-closed conditions. The results demonstrated that CPR outperformed other known traditional linear methods on distinguishing seizure and pre-seizure states, identifying epileptic foci, and differentiating alongside eyes-open and eyes-closed conditions.

In another study which demonstrates the effectiveness of RQA in analyzing EEG signals for epilepsy detection, Gruszczyńska et al.[11] applied RQA on such signals in order to distinguish epileptic from healthy patients using recordings from frontal and temporal lobe electrodes (Fp1, Fp2, T3, T4). In their findings they have showed that the epileptic signals present more periodic dynamics in comparison to healthy controls, by producing higher values of RQA parameters such as determinism, laminarity, and longest diagonal line. The combination of RQA with Principal Component Analysis for dimensionality reduction and visualization, achieved 86.8% classification accuracy with SVM. Authors also demonstrated RQA's capability to identify pathological patterns in EEG signals without the requirement of seizure events during recording which have bad impact on the subject's health.

Another study utilizing advanced nonlinear analysis techniques for neural correlation investigation to cognitive functions [12] used *stereoelectroencephalography (sEEG)* combined alongside RQA

for the examination of the relationship of the DMN and empathy. Correlations have been detected relating specific RQA metrics (mean diagonal line length, entropy of diagonal line lengths, trapping time) and empathy scores, particularly within DMN subsystems.

Regarding epilepsy diagnosis, authors in [13] proposed a new framework utilizing the combination of RQA with genetic algorithms and Bayesian classifiers for identifying corresponding biomarkers for seizure detection. They utilized five distance norms (e.g., Euclidean, Mahalanobis) and multiple thresholds for extracting recurrence features from EEG signals, achieving 100% classification accuracy. More specific, the *transitivity* feature has shown capability of a highly discriminative biomarker, performing better compared to traditional linear methods.

Ngamga et al.[14] studied the performance achieved of RQA and Recurrence Network (RN) measures in identifying pre-seizure states from multi-day, multi-channel intracranial EEG (iEEG) recordings of epilepsy patients. Results highlighted the correlation among RQA measures (determinism, laminarity, and mean recurrence time) in detecting seizure precursors, while RN measures (average shortest path length and network transitivity) provided complementary but not so consistent insights than using the application of RQA measures alone.

Gao et al.[45] examined the application of RQA in the domain of automated epilepsy detection. Authors utilized a hybrid scheme combining nonlinear features(related to Approximate Entropy(ApEn) and RQA metrics) from the publicly available Bonn EEG dataset[46] with a deep learning classifier. Their key finding was that while ApEn and RQA features alone could achieve good classification accuracy, their performance was increased when used as input features for a Convolutional Neural Network (CNN). By constructing this hybrid approach, classification accuracy rised on 99.26% for distinguishing ictal from inter-ictal and healthy EEG signals, demonstrating the potential of the synergy among traditional metrics and modern deep learning architectures.

Researchers in [3], have applied RQA on resting-state fMRI data from TgF344-AD rats(a transgenic rat model which will eventually develop Alzheimer's disease) and their healthy-control counterparts wild-type rats(WT), in order to detect early stage biomarkers for the disease. By analyzing Default Mode-Like Network (DMLN) using RQA metrics(*entropy*, *recurrence rate*, *determinism* and *average diagonal line length*) changes have been detected in regions of the basal forebrain, hippocampal fields (CA1, CA3), and visual cortices (V1, V2). Also on the study's findings include reduced predictability in WT rats with aging, while AD rats exhibited less decline in predictability, suggesting some unknown yet countereacting mechanisms. This study highlights RQA's sensitivity for nonlinear dynamics in preclinical AD and the code used is also publicly available.

Lombardi et al.[5] investigated the nonlinear properties in fMRI BOLD signals during a working memory task in schizophrenic patients and healthy controls. They have attempted by using RQA, to analyze recurrence plots for quantifying determinism, trapping time, and maximal vertical line length in functionally relevant brain clusters. Outcome revealed differences in the dynamics between the two groups, and more specific in working memory and DMN areas. While their work have focused on fMRI, the methodology can be adapted also into EEG signals, which can offer a higher resolution for capturing rapid neural dynamics.

Kang et al. [2], in their study explore the dynamics and functional connectivity of the Default Mode Network (DMN) in schizophrenia, applying RQA-CRQA on resting-state fMRI data. Findings include decreased *determinism* between specific DMN regions (vMPFC-posterios cingulate and vMPFC-precuneus) in first-episode schizophrenia patients, as a signal of disturbed predictability of functional interactions. Moreover, their results achieve to correctly classify using SVM(support vector machine) schizophrenia patients from healthy controls with 77% classification accuracy.

In their research, Pentari et al.[9] have applied CRQA to resting-state fMRI data for examining

the dynamic functional connectivity on patients with neuropsychiatric systemic lupus erythematosus (NPSLE). Results contain the fact that CRQA metrics, such as determinism, appear more sensitive than conventional static functional connectivity methods in order to identify aberrant connectivity patterns that correlated with visuomotor performance. The study focused on 16 frontoparietal regions and found that CRQA could detect both increased and decreased connectivity in NPSLE patients compared against the healthy controls. Building on these findings, Pentari et al.[10] subsequently expanded the investigation to whole brain network analysis in a larger cohort. In this study they demonstrate the capability of CRQA to integrate multiple recurrence metrics for revealing both hyperconnectivity in parietal regions (angular gyrus and superior parietal lobule) and hypoconnectivity in medial temporal structures (hippocampus and amygdala).

In addition there have been works where simulated data have been used in conjunction with RQA. Lameu et al.[4], investigated burst phase synchronization in neural networks using RQA. They analyzed two network types; a small-world network and a network of networks (to mimic better the real human brain), using coupled Rulkov maps to model bursting neurons. By applying RQA, they identified synchronized neuron groups and quantified their sizes during synchronization transitions. The study showed that RQA measures (*recurrence rate, laminarity inspired*(custom feature), *and average structure size*) complement traditional order parameters by revealing localized synchronization patterns, such as the formation and growth of synchronized clusters. Kashyap and Keilholz[26] conducted a comprehensive comparison between simulated brain network models (BNMs) and real rs-fMRI data using dynamic analysis techniques, including Recurrence Quantification Analysis (RQA). In the study they employed two BNMs, the Kuramoto oscillator model and the Firing Rate model, for simulating the whole-brain activity, which was then compared to human rs-fMRI data. Among the compared dynamic analysis methods, RQA was proved particularly effective in distinguishing between the models and empirical data, demonstrating that RQA metrics (*recurrence rate, entropy, and average diagonal length*) could robustly separate the empirical data from simulations.

Shalbaf et al. [28] investigated the synchronization of EEG signals between frontal and temporal regions during propofol anesthesia using *Order Patterns Cross Recurrence Analysis* (OPCR). Their study introduced a novel index, *Order Pattern Laminarity* (OPL), for the quantification of neuronal synchronization and compared its performance with the traditional Bispectral Index (BIS). The results demonstrated that OPL correlated more strongly with propofol concentration ($P_k = 0.9$) and exhibited faster response times to transient changes in consciousness compared to BIS. Additionally, OPL showed lower variability at the point of loss of consciousness (LOC), suggesting its robustness as a measure of anesthetic depth. This work highlights the potential of recurrence-based methods (e.g., CRQA) for analyzing brain network dynamics under anesthesia, particularly in noisy, non-stationary EEG data.

2.1 RQA relevant patents utilizing EEG modality

The application of RQA utilizing EEG modality on the biomedical field, shows an increasing interest not only in academic research, but also in commercial and clinical applications, as we can inspect on recent patent filings. Reviewing these documents can reveal industrial viable solutions being developed for real-time, embedded systems.

Becker et al.[42] describes on patent (US20080234597A1) a monitoring device and method for creating an assessment of the depth of anesthesia or coma characterizing an individual subject. Authors analyzes neuronal EEG data and uses RQA to compute a complexity parameter that

Table 1: Comparison among the retrieved studies using recurrence analysis

#	Reference	Modality	Analysis Methods	Network Type
1	Frolov et al. (2020)	EEG	RQA, CRQA	Multiplex functional networks
2	Kang el al. (2023)	fMRI	RQA, CRQA	DMN, schizophrenia
3	Rezaei el al. (2023)	fMRI	RQA	Default model-like network, AD
4	Lameu et al. (2018)	—	RQA	Small-world & cluster network
5	Lombardi et al. (2014)	fMRI	RQA	schizophrenia, working memory
6	Pitsik E. (2025)	EEG	RQA	aging
7	Guglielmo et al. (2022)	EEG	RQA	cognitive tasks
8	Lopes et al. (2020)	sEEG, MEG	RQA	epilepsy
9	Pentari et al. (2022)	fMRI	RQA, CRQA	NPSLE
10	Pentari et al. (2023)	fMRI	CRQA	NPSLE
11	Gruszczyńska et al. (2019)	EEG	RQA	epilepsy
12	Mo et al. (2022)	sEEG	RQA	DMN, epilepsy
13	Palanisamy et al. (2024)	EEG	RQA	epilepsy
14	Ngamga et al. (2016)	EEG	RQA,RN	epilepsy
15	Fan and Chou (2019)	EEG	RQA,RN	epilepsy, seizure detection
16	Nunez et al. (2020)	EEG	RQA	AD
17	Yang et al. (2019)	sEEG	RQA,CRQA	epilepsy
18	Rangaprakash (2014)	EEG	CPR(CRQA-based)	epilepsy
19	Heunis et al. (2018)	rsEEG	RQA	autism spectrum disorder
20	Timothy et al. (2017)	EEG	RQA-CRQA	MCI
21	Kashyap et al. (2019)	fMRI	RQA	distinguish BNMs
22	Shalbaf et al. (2014)	EEG	CRQA(OPL)	Anesthesia depth monitoring
23	Mihajlović. (2019)	EEG	RQA	cognitive tasks

quantitatively reflects the level of consciousness. In the device’s core, a buffer is utilized for storing time-series data and an analysis circuit performs RQA by reconstructing phase-space trajectories, calculating recurrence plots, and extracting determinism-based complexity measures. This makes possible monitoring the depth of anesthesia level in real-time and can be utilized in clinical applications for anesthesia control during surgery or even in long term coma assessment.

Patent US20250195894A1 [39], entitled “Systems and Methods for Seizure Detection and Closed-Loop Neurostimulation,” Inventors proceed in an alternative calculation of RQA measures which entirely bypass the construction of the recurrence plot matrix(RP). They achieve this, by not creating the traditional RP in order to extract certain metrics from, but by calculating them, *on-the-fly*; dynamically accumulating the lengths of diagonal lines as each new data point is processed. This method offers the following advantages:

1. **Memory Efficiency:** It does not require the large $N \times N$ comparison matrix, reducing memory usage by approximately 88%.
2. **Computational Efficiency:** It avoids the expensive read/write cycles associated with managing the large matrix, reducing processing time by approximately 30% per channel.

In addition, the patent by [40], titled ”An EEG signal classification model based on genetic

algorithm and random forest”, presents a framework for EEG signal classification. The inventors propose a hybrid model consisting of three key stages:

1. **Feature Extraction:** The method employs a multi-modal feature extraction strategy. Among other features, it explicitly includes *RQA* metrics from the EEG signal, alongside other traditional time-domain/frequency features.
2. **Feature Optimization:** A genetic algorithm (GA) is then utilized for feature selection. Inventors use binary encoding for representing chromosomes, where each bit corresponds to the selection (1) or rejection (0) of a specific feature from the large extracted pool. The aim of this procedure is to optimize the feature subset in order to have maximum discriminative power.
3. **Classification:** The optimized feature subset is fed into a Random Forest classifier for a final prediction.

The patent claims that this integrated approach, validated on a public dataset, yields a superior classification accuracy compared to existing methods at the time of filing, while also demonstrating robustness through cross-validation.

Another patent[41] (CN106512206B) describes an implantable closed-loop deep brain stimulation (DBS) system that uses electrophysiological signals (specific deep brain local field potentials (LFPs) and ECG signals) for monitoring the states of a human sleeping and adjusting various stimulation parameters in real time. A device acquires ECG and deep brain signals for feature extraction in the domains of time and frequency, while calculating complexity measures. RQA metrics such recurrence rate, determinism, entropy and laminarity are utilized alongside other complexity and spectral features to classify sleep states and trigger appropriate stimulation responses. Features are then utilized for detection of sleep stages and for emergency detection/alerts (f.e, cardiac arrest or abnormal excitation).

3 The CHB-MIT EEG Database

The **CHB-MIT Scalp EEG Database**[47] is a public collection of EEG recordings from pediatric subjects that includes intractable seizures. All the recordings of the database were collected at the Children’s Hospital Boston. The dataset contains multiple cases (patients), each with long-term scalp EEG signals recorded using the international 10–20 system. Table 2 provides a summary of the demographic information of the database’s subjects. This particular database is used in many studies for testing algorithms on epileptic seizure detection and epileptic research in general.

3.1 Dataset Description

The database includes recordings from male and female patients, with their ages in the range of 1.5 to 22 years old. Most subjects are children, a fact reflecting the pediatric nature of the dataset. This demographic diversity provides a representative sample for studying epileptic activity across different developmental stages.

The total recording duration spans approximately 982 hours, segmented into 664 different EDF files, where each recording contains 1 hour of data (though durations vary from 1 to over 4 hours). EEG signals were recorded using 23 scalp electrodes (according to the international 10–20 system),

Table 2: Demographic information for patients in the CHB-MIT Scalp EEG Database.

Case	Gender	Age (years)
chb01	F	11
chb02	M	11
chb03	F	14
chb04	M	22
chb05	F	7
chb06	F	1.5
chb07	F	14.5
chb08	M	3.5
chb09	F	10
chb10	M	3
chb11	F	12
chb12	F	2
chb13	F	3
chb14	F	9
chb15	M	16
chb16	F	7
chb17	F	12
chb18	F	18
chb19	F	19
chb20	F	6
chb21	F	13
chb22	F	9
chb23	F	6

but in some files there are extra channels records, such as ECG and EMG references. The sampling rate of the recordings is 256 Hz with 16-bit resolution. Seizure events are annotated by domain experts, providing a time-stamp for seizure onsets and offsets.

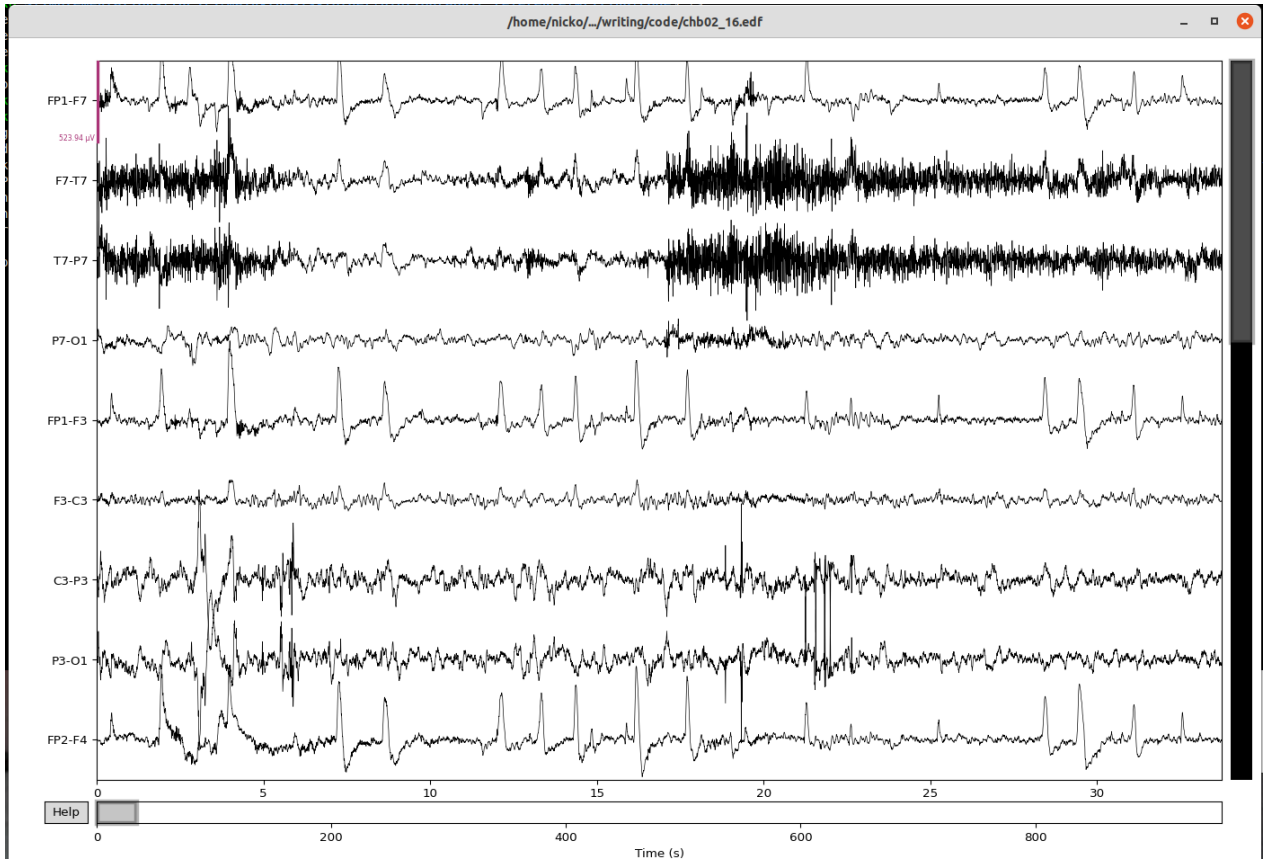


Figure 1: EEG recording visualized utilizing Python-MNE, for a patient of the dataset.

4 Filtering

Signals of electroencephalogram, in most cases carry not only the desired signal but also added noise and artifacts from physiological (eye blinks, muscle or cardiac activity) and non-physiological sources (f.e. powerline interference). In general, artifacts in a recording consist of all the non-neural signals that are mixed with the pure EEG data.

Preprocessing is required in order to increase the quality of the signal(signal-to-noise ratio), to boost the performance of further examinations in later pipelines in domains like brain-computer interfaces (BCIs) or clinical diagnostics[29, 30, 31, 32].

Some of the common preprocessing techniques are:

- **Filtering** (f.e, Butterworth, Chebyshev) to remove unwanted frequency bands.
- **Regression methods** used for removing ocular artifacts with help of reference channels.
- **Blind Source Separation (BSS)** (f.e, ICA, CCA), decompose and isolate neural activity from artifacts.

- **Wavelet/EMD-based methods** for non-stationary artifact removal.

Also hybrid approaches (f.e, wavelet-ICA) exist, combining multiple techniques for improved artifact rejection. The choice of method is related on computational constraints, artifact type, and the satisfaction of real-time processing needs, if any.

Effective preprocessing is a key, that ensures reliable feature extraction for later analysis.

Among these, wavelet-based methods have been proposed and used for effective EEG denoising based on the non stationary nature of brain signals. Transient artifacts appearing in EEGs in varying frequency patterns are the reason that traditional linear filtering methods struggle to handle them without introducing distortions. Wavelet transforms on the other hand, decompose the signal into time-frequency representations using scalable, localized basis functions called wavelets, enabling the analysis: the signal is broken down into approximation (low-frequency) and detail (high-frequency) coefficients across decomposition levels. Artifacts, such as ocular blinks or EMG bursts, manifest as sparse, high-amplitude coefficients that can be selectively silenced using thresholds (f.e, hard or soft), followed by reconstruction—thus removing noise while preserving neural transients like epileptic spikes [55].

4.1 Evaluation metrics for EEG denoising in the CHB-MIT dataset

In order to evaluate the performance of different wavelet-based filters for EEG denoising, quantitative metrics have been computed over entire recordings per channel and filter configuration. Metrics used were:

- **Signal-to-Noise Ratio (SNR, dB)**: Measure of the ratio between the power of the clean signal and the power of the noise. Higher values indicate better noise suppression while preserving the structure of the signal.
- **Root Mean Square Error (RMSE, μV)**: This metric quantifies the average deviation between the denoised and reference signal in microvolts. Lower values indicate closer similarity to the original signal.
- **Normalized RMSE (NRMSE, %)**: RMSE normalized by the dynamic range of the reference signal expressed as a percentage. Lower values represent better performance.
- **Correlation Coefficient**: Pearson's correlation among the denoised and reference signal, providing an assessment of similarity of the two waveforms. Observed values close to 1 indicate high waveform preservation.
- **Percent Root-mean-square Difference (PRD, %)**: Quantification of the relative distortion introduced by the denoising process. Lower values indicate less distortion.

For each one of the filters configuration, metrics were computed channel-wise and then averaged across all channels in order to calculate their global performance score.

4.2 Filter Selection Criteria

The selection of the optimal filter was based on a multi-criteria ranking strategy, where:

1. Metrics where *higher* values indicate better performance (**SNR**, **Correlation**) were ranked in descending order.

2. Metrics where *lower* values indicate better performance (**RMSE**, **NRMSE**, **PRD**) were ranked in ascending order.

3. The ranks from all metrics were averaged to obtain an overall performance rank for each filter.

The filter with the lowest average rank was considered the best compromise between noise reduction and signal fidelity. According to this evaluation, the **SYM8, level 4, threshold 0.5, hard thresholding** filter presented the highest overall performance, exhibiting:

- the highest SNR values,
- one of the lowest RMSE and the lowest NRMSE value,
- the highest correlation coefficient,
- and the lowest PRD.

This indicates that the chosen filter effectively suppressed noise while preserving the morphological features of the EEG signal, making it the most suitable choice for subsequent analysis. The results of the benchmarking those metrics in 5 EDF recording files which include seizures are presented in the following figure.

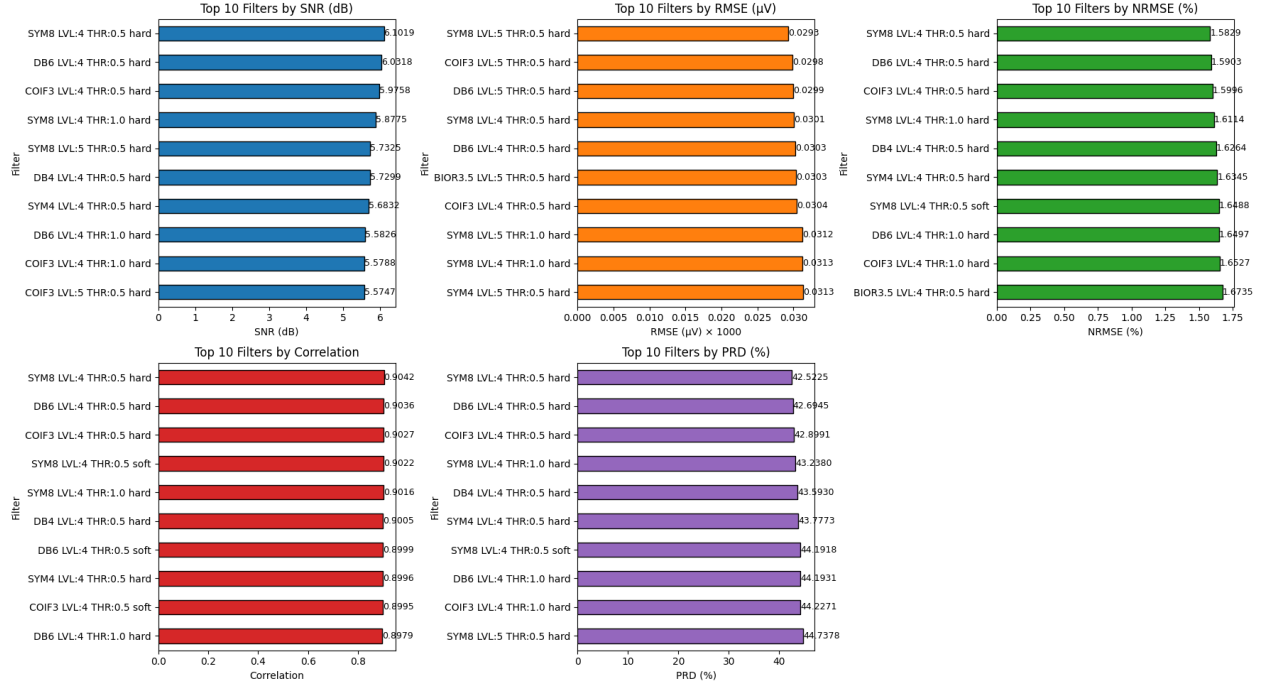


Figure 2: The top 10 filter configurations per EEG metric. RMSE values scaled.

In the following figures we present a visualization using the recording named *chb01_03.edf* comparing the original 10 first EEG channels against the filtered ones with the respective wavelets filters.

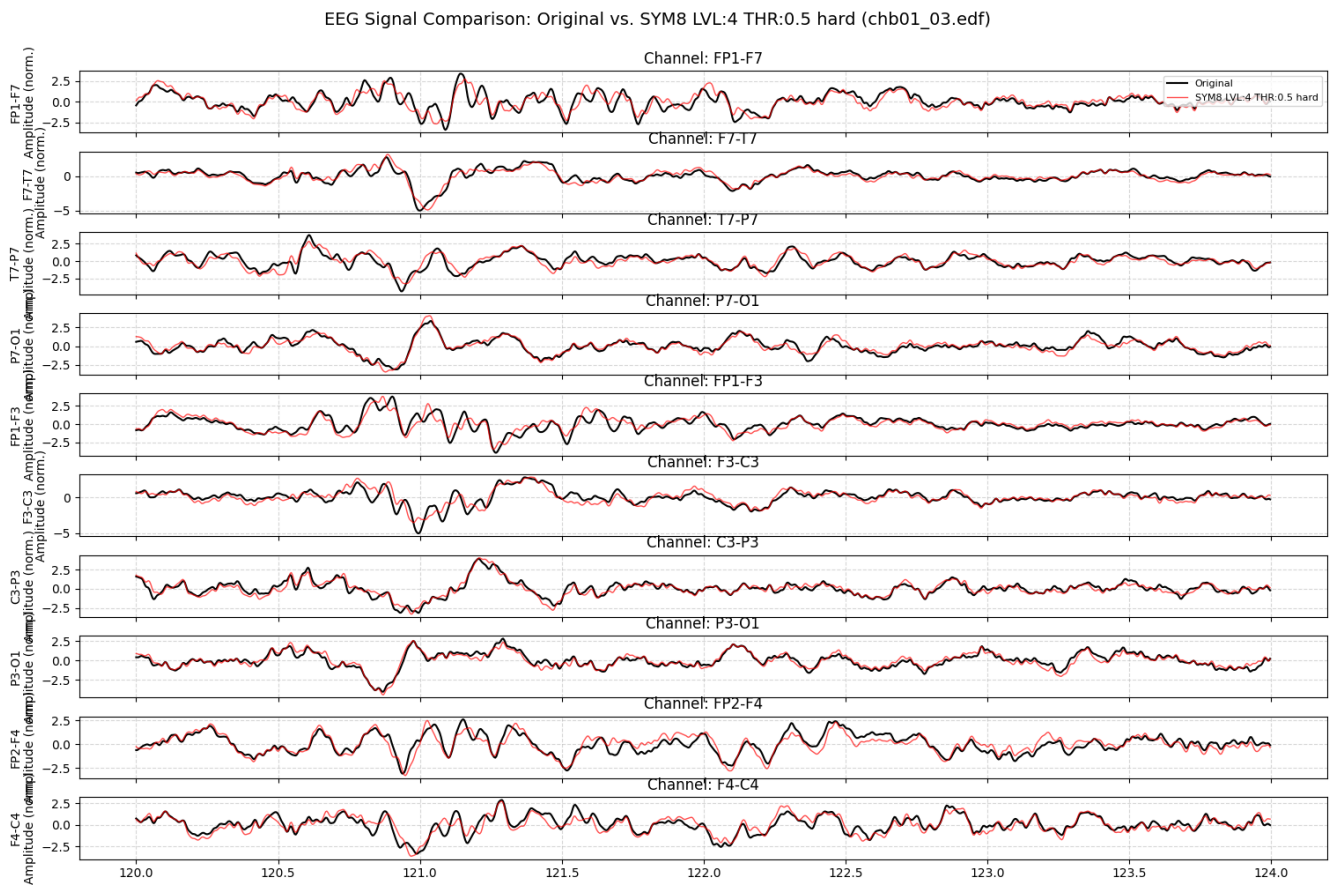


Figure 3

EEG Signal Comparison: Original vs. DB6 LVL:4 THR:0.5 hard (chb01_03.edf)

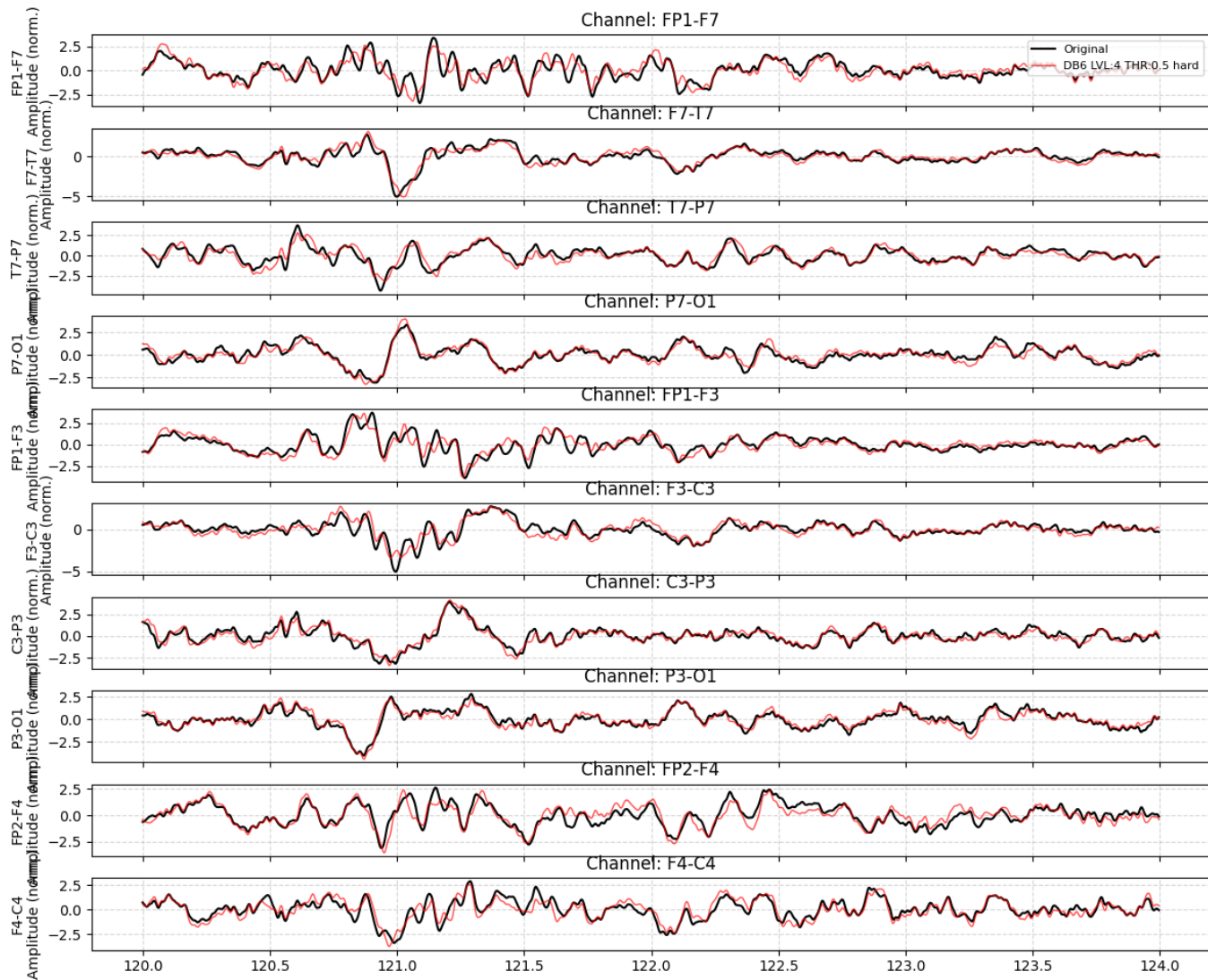


Figure 4

5 Phase space reconstruction of EEG signals

In order to analyze multi-channel EEG's nonlinear dynamical system, the reconstruction of the underlying phase space is required, based on the scalar measurements of each channel. According to Takens' embedding theorem [34], a time series $x(t)$ can be embedded in an m -dimensional space using time-delay coordinates:

$$\vec{y}(t) = [x(t), x(t + \tau), x(t + 2\tau), \dots, x(t + (m - 1)\tau)] \quad (1)$$

where m is the embedding dimension and τ is the time delay. The critical challenge lies in determining the appropriate values for these parameters to faithfully reconstruct the system's dynamics without distortion.

5.1 Determination of Embedding Parameters

The reconstruction of the phase space from a single time series $x(t)$ requires the specification of two parameters: the time delay τ and the embedding dimension m . These two parameters determine how the reconstruction will represent and how close it will reveal the underlying dynamics without distortion.

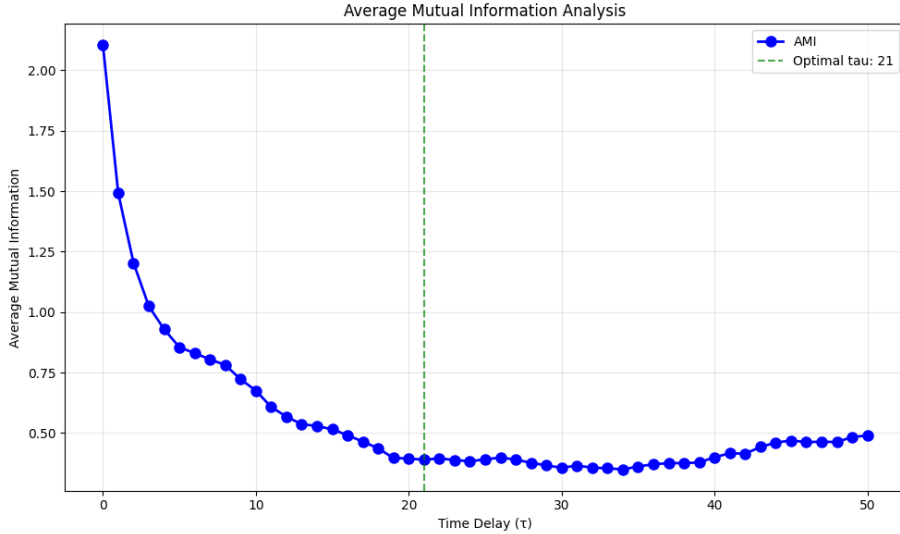


Figure 5: Calculation of τ using AMI for a sample EEG channel. The first minimum of the AMI function (green dashed line) is chosen to become the optimal τ to ensure independence between delay coordinates.

5.1.1 Calculation of time delay τ utilizing mutual information

The time delay τ can be estimated by applying the *Average Mutual Information* (AMI) method, a concept which was first introduced by Fraser and Swinney [36]. In contrast to linear autocorrelation, mutual information has the ability to capture both linear and nonlinear dependencies among the original time series $x(t)$ and its delayed version $x(t + \tau)$.

The mutual information $I(\tau)$ between $x(t)$ and $x(t + \tau)$ is defined as:

$$I(\tau) = \sum_{x(t), x(t+\tau)} P(x(t), x(t + \tau)) \log_2 \left(\frac{P(x(t), x(t + \tau))}{P(x(t)) P(x(t + \tau))} \right)$$

where $P(\cdot)$ denotes probability.

The optimal time delay τ is chosen as the value at which $I(\tau)$ reaches its *first minimum*. This value indicates a good compromise between independence (too small τ) and irrelevance (too large τ) of the coordinates in the embedding vector.

5.1.2 Estimating embedding dimension m using false nearest neighbors approach

When the embedding dimension m is too small, the phase space becomes *projected* rather than properly *embedded*. This projection can create artificial neighborhoods where points appear to be close due to geometrical constraints of the space rather than their actual dynamical similarity. These are named as *false nearest neighbors*. An example of such an occurrence can be observed in Figure 7.

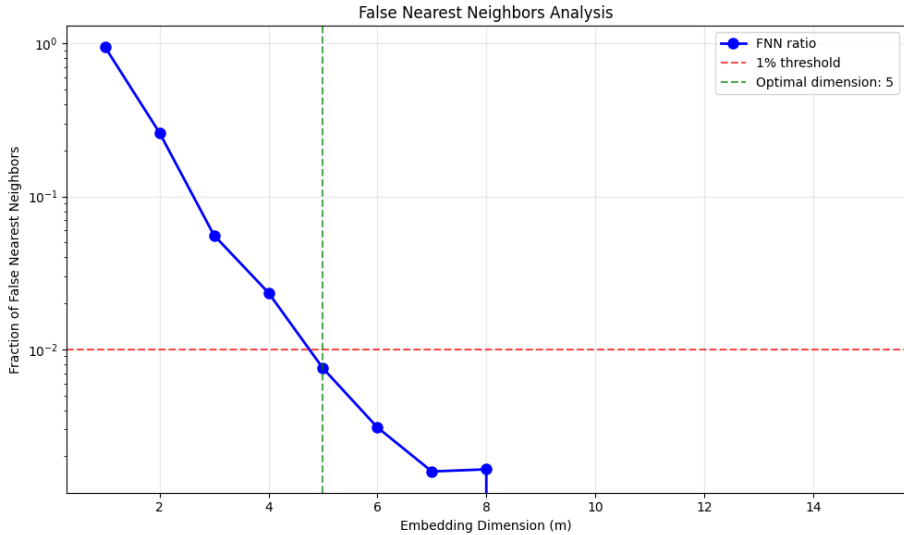


Figure 6: Calculation of the embedding dimension using the FNN scheme.

Mathematically, two points \vec{y}_i and \vec{y}_j are false neighbors if their distance increases significantly when embedded in higher dimension:

$$\frac{\|\vec{y}_i^{(m+1)} - \vec{y}_j^{(m+1)}\|}{\|\vec{y}_i^{(m)} - \vec{y}_j^{(m)}\|} > R_{\text{tol}} \quad (2)$$

where R_{tol} is a tolerance threshold (typically 10–15).

The False Nearest Neighbors(FNN) method [35] provides a systematic approach in order to determine the minimal sufficient embedding dimension. The method's steps are:

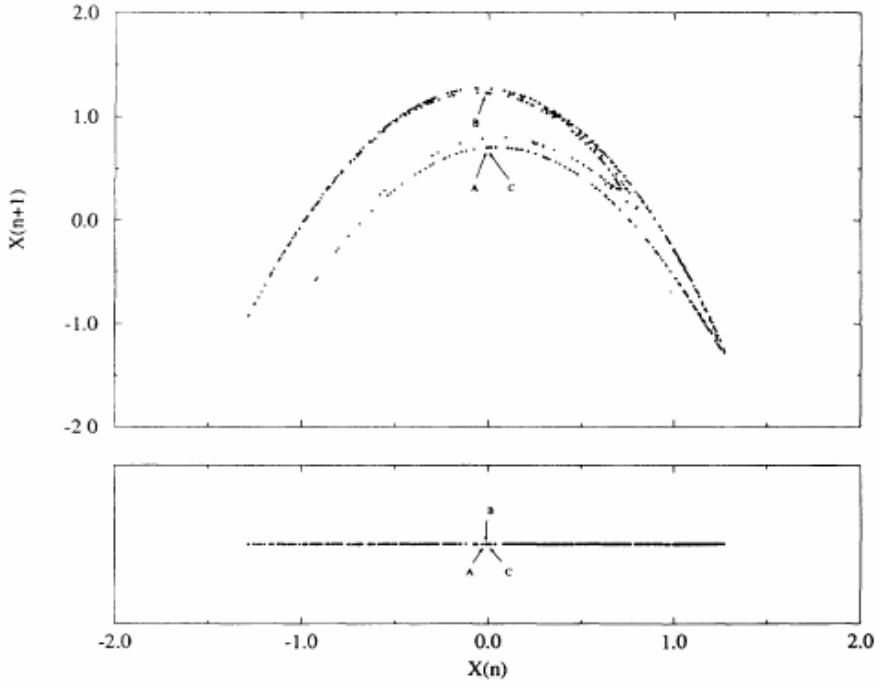


Figure 7: Schematic illustration of false neighbors. In insufficient embedding dimension (down), points A and B appear neighbors due to projection. When proper embedding is employed(up), their true separation is revealed.

1. For each point in dimension m , identify its nearest neighbor.
2. Embed the data in dimension $m + 1$.
3. Calculate the relative distance increase between each point and its former neighbor.
4. If the increase exceeds predetermined thresholds, the neighbor point is classified as a false neighbor
5. The optimal m is the smallest dimension where the fraction of false neighbors drops below an acceptable level (typically 1–5%)

Both relative and absolute criteria are included in the algorithm:

$$\text{Relative: } \frac{\|\bar{y}_i^{(m+1)} - \bar{y}_j^{(m+1)}\|}{\|\bar{y}_i^{(m)} - \bar{y}_j^{(m)}\|} > R_{\text{tol}} \quad (3)$$

$$\text{Absolute: } \|\bar{y}_i^{(m+1)} - \bar{y}_j^{(m+1)}\| > A_{\text{tol}} \cdot \sigma_x \quad (4)$$

where σ_x is the standard deviation of the time series.

When optimal parameters τ and m have been determined for a given signal, the phase space can be reconstructed according to Takens' theorem. This reconstruction provides the geometric picture of the underlying dynamics.

Figure 8 presents the reconstructed phase space for a normal EEG segment from channel 'Fp1-F7' from CHB-MIT's chb24_01.edf data.

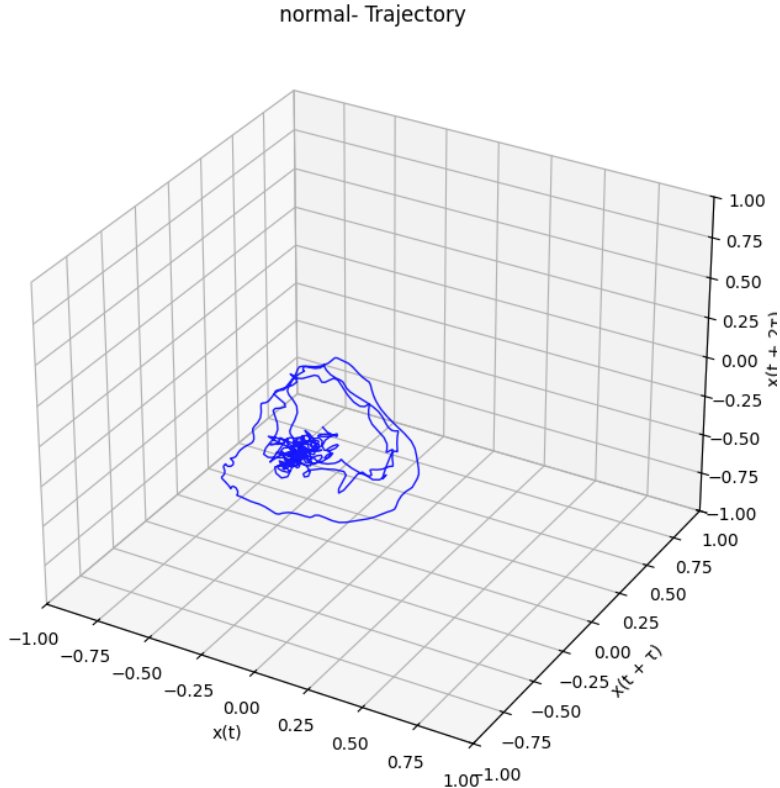


Figure 8: 3D phase space reconstruction for a 3-second EEG segment's channel.

6 Recurrence Quantification Analysis (RQA)

Having reconstructed the phase space trajectory of the EEG signals, the next step is to analyze its dynamical properties. Recurrence Quantification Analysis is a powerful nonlinear method that provides precisely this functionality by quantifying the number and duration of recurrences of a dynamical system to its previous states [37]. The core of this quantification process is the *Recurrence Plot (RP)*, a visualization which denotes the times at which the phase space trajectory revisits approximately the same area. In most non trivial cases, a phase space does not have a dimension (two or three) which allows a direct visualization, so for higher dimensional phase spaces the only solution is a projection into a two or three dimensional space. However, RP enables the examination of a higher-dimensional phase space trajectory via its two-dimensional representation of its recurrences.

6.1 The Recurrence Plot (RP)

RP is a symmetric, two-dimensional matrix that visualizes the recurrences of states. For a reconstructed trajectory $\vec{y}(t)$ of length N , the recurrence matrix \mathbf{R} is defined as:

$$R_{i,j} = \Theta(\varepsilon - \|\vec{y}(i) - \vec{y}(j)\|), \quad i, j = 1, \dots, N \quad (5)$$

where:

- $\Theta(\cdot)$ is the Heaviside step function ($\Theta(x) = 0$ if $x < 0$, and $\Theta(x) = 1$ otherwise),
- ε is a predefined distance threshold (radius),
- $\|\cdot\|$ is a norm.

By interpreting the RP, several metrics can be extracted for further analysis.

6.2 Key RQA Metrics and Their Interpretation

RQA provides a set of metrics that can quantify the number and the duration of the recurrences of a dynamical system. These metrics are categorized by those which are based on diagonal structures, which relate to the predictability and deterministic nature of the system, and those that are based on vertical structures, which can capture laminar states or chaos-chaos transitions.

The definitions of the core RQA metrics, as implemented in tools like the utilized PyRQA, are as follows [44]:

Recurrence Rate (RR) The recurrence rate is the simplest measure, defined as the density of recurrence points in the RP. It corresponds to the probability that a state recurs and is analogous to the correlation sum.

$$RR = \frac{1}{N^2} \sum_{i,j=1}^N R_{i,j}$$

Determinism (DET) Determinism quantifies the percentage of recurrence points that form diagonal lines. Diagonal lines are a signature of deterministic dynamics, where segments of

the trajectory run in parallel for some time. A higher DET indicates a more predictable, deterministic system.

$$DET = \frac{\sum_{l=l_{\min}}^N l P(l)}{\sum_{l=1}^N l P(l)}$$

where $P(l)$ is the histogram of diagonal line lengths l , and l_{\min} is the minimum line length (typically 2).

Laminarity (LAM) Laminarity measures the percentage of recurrence points that form vertical lines. Vertical lines indicate states that do not change or change very slowly for a period (laminar states). It can detect chaos-chaos transitions or intermittency.

$$LAM = \frac{\sum_{v=v_{\min}}^N v P(v)}{\sum_{v=1}^N v P(v)}$$

where $P(v)$ is the histogram of vertical line lengths v , and v_{\min} is the minimum line length.

Ratio (RATIO) The ratio is a measure of complexity, calculated as the ratio between DET and RR. It can be sensitive to transitions between order and chaos.

$$RATIO = \frac{N^2 \sum_{l=l_{\min}}^N l P(l)}{\left(\sum_{l=1}^N l P(l) \right)^2}$$

Average Diagonal Line Length (L) This metric represents the average time that two segments of the trajectory remain close, providing an estimate of the mean prediction time.

$$L = \frac{\sum_{l=l_{\min}}^N l P(l)}{\sum_{l=l_{\min}}^N P(l)}$$

Trapping Time (TT) Trapping time is the average length of vertical lines, quantifying the mean time the system remains trapped in a specific state (laminarity in time).

$$TT = \frac{\sum_{v=v_{\min}}^N v P(v)}{\sum_{v=v_{\min}}^N P(v)}$$

Longest Diagonal Line (L_{\max}) The length of the longest diagonal line in the RP is related to the Lyapunov exponent of the system. A shorter L_{\max} suggests a faster divergence of trajectories, which is a hallmark of chaos.

$$L_{\max} = \max(\{l_i \mid i = 1, \dots, N_l\})$$

Divergence (DIV) Divergence is the inverse of L_{\max} . It is related to the Kolmogorov-Sinai entropy and the sum of the positive Lyapunov exponents, providing a measure of how quickly nearby trajectories diverge.

$$DIV = \frac{1}{L_{\max}}$$

Longest Vertical Line (V_{\max}) The length of the longest vertical line is another indicator of the system's laminar behavior.

$$V_{\max} = \max(\{v_i \mid i = 1, \dots, N_v\})$$

Entropy (ENTR) The Shannon entropy of the probability distribution $p(l)$ of the diagonal line lengths. It reflects the complexity of the deterministic structure in the system. A higher ENTR indicates a more complex and less periodic dynamics.

$$ENTR = - \sum_{l=l_{\min}}^N p(l) \ln p(l), \quad \text{where } p(l) = \frac{P(l)}{\sum_{l=l_{\min}}^N P(l)}$$

Trend (TREND) Trend quantifies the paling of the RP towards its edges, which can be caused by non-stationarity in the data (e.g., a slow drift in the mean of the signal). It is calculated as the slope of the linear regression of the local recurrence rate RR_i over the distance from the main diagonal.

$$TREND = \frac{\sum_{i=1}^{\tilde{N}} (i - \tilde{N}/2)(RR_i - \langle RR_i \rangle)}{\sum_{i=1}^{\tilde{N}} (i - \tilde{N}/2)^2}$$

where \tilde{N} is the number of diagonals parallel to the Line of Identity (LOI) that are considered, and RR_i is the recurrence rate in the i -th diagonal.

These metrics, when applied to EEG signals, allow for the characterization of the brain's dynamic states. For example, in an epileptic seizures occurrence, often higher determinism (DET), laminarity (LAM) or recurrence rate(RR) is observed compared to the more stochastic and complex inter-ictal states. This fact makes RQA metrics a viable solution for identifying pathological patterns.

6.3 Cross-Recurrence Quantification Analysis (CRQA)

While Recurrence Quantification Analysis (RQA) is powerful for analyzing the dynamics of a single system, many real-world phenomena, including brain activity, involve the interaction between multiple subsystems. Cross-Recurrence Quantification Analysis (CRQA) extends the concepts of RQA to analyze the coupling, synchronization and degree of similarity that the dynamics between two different systems present[?].

6.3.1 The Cross-Recurrence Plot (CRP)

The foundation of CRQA is the Cross-Recurrence Plot (CRP). For two reconstructed phase space trajectories $\vec{x}(i)$ from system X and $\vec{y}(j)$ from system Y , both of length N , the cross-recurrence matrix is defined as:

$$CR_{i,j} = \Theta(\varepsilon - \|\vec{x}(i) - \vec{y}(j)\|), \quad i, j = 1, \dots, N \quad (6)$$

Unlike the standard RP, which is symmetric about the main diagonal (Line of Identity, LOI), the CRP is generally *not symmetric*. This asymmetry can reveal directional relationships or leader-follower dynamics alongside the two systems.

6.3.2 Interpretation of CRQA Metrics

The same quantitative measures defined for RQA (Section 6.2) can be applied to the CRP, but their interpretation shifts from describing *self-similarity* to describing *coupling* and *interaction*:

- **Cross-Recurrence Rate (CRR):** The probability that the state of system X at time i is close to the state of system Y at time j . A high CRR indicates overall similar states between the two systems.
- **Cross-Determinism (CDET):** The percentage of recurrent points in the CRP that form diagonal lines. Diagonal lines occur when the two systems follow a similar path in phase space for some time. **This is a crucial metric for epilepsy detection**, as it quantifies the transient synchronization between different brain regions. A seizure often manifests as increased CDET between channels in the epileptogenic zone.
- **Cross-Laminarity (CLAM):** Measures the laminarity between the two systems, indicating when one system gets trapped in a state while the other changes.
- **Average Diagonal Line Length (L)** in the CRP estimates the mean time that the two systems remain synchronized or follow a similar trajectory.

6.3.3 Why CRQA for Multi-Channel EEG?

Applying CRQA to pairs of EEG channels is particularly well-suited for epilepsy detection for several reasons:

- **Synchronization Detection:** Epileptic seizures are characterized by abnormal, excessive synchronization of neuronal populations. CRQA directly quantifies this synchronization in the phase space.
- **Nonlinear and Non-stationary:** CRQA does not assume linearity or stationarity, making it robust for analyzing the complex, transient dynamics of EEG signals.
- **Directional Insights:** While not explored in all analyses, the potential asymmetry of the CRP can, in principle, help identify the propagation path of a seizure.
- **Focus on Interaction:** It moves beyond analyzing individual channels in isolation to directly measure the dynamic interplay between different brain regions, which is often where the pathology lies.

In this thesis, CRQA is employed to compute a set of features (Table 4) for all unique pairs of EEG channels. These features capture the complex synchronization patterns that distinguish pre-ictal, ictal, and inter-ictal states, forming the basis for the subsequent machine learning classification.

7 Methodology

In this section the methodology for processing EEG data is described in order to perform CRQA to analyze epileptic and non-epileptic brain activity. The approach consists of different parts such as loading and segmenting EEG recordings, extracting non-overlapping time windows, selecting the embedding parameters and computing CRQA features for channel pairs. The methodology is implemented in Python using libraries such as `numpy`, `torch`, `pyopencl`, and `pyrqa` [43].

7.1 Data preprocessing and windowing

Prior filtererd EEG recordings are stored in NumPy array format (`.npy`), accompanied by metadata specifying the sampling frequency (f_s) and channel information (22 channels with their respective labels as FP1-F7, F7-T7, ..., FT10-T8). The time axis is computed as $t = \frac{n}{f_s}$, where n is the sample index and f_s is the sampling frequency in Hertz (Hz).

Each EEG channel is segmented into continuous regions based on predefined boundaries from CHB-MIT dataset [47] annotations, distinguishing epileptic from non-epileptic segments. Then, each segment is further divided into non-overlapping time windows of fixed size (512 samples, equivalent to 2 seconds at 256 Hz). For each segment, the number of windows is calculated by performing integer division of the segment length by the window size and discarding any incomplete windows. Each window is associated with a segment index, window index within the segment, start and end sample indices, and a label (1 for epileptic, 0 for non-epileptic).

7.2 Embedding parameters selection

The embedding dimension m and time delay τ were set to 3 and 1, respectively, following common practice in EEG analysis. This choice was also motivated by known limitations that the FNN algorithm experiences when applied to noisy and autocorrelated signals, such as EEG. For instance, [48] have shown that FNN can falsely indicate low-dimensional determinism in autocorrelated stochastic processes, while [49] showed that the effects of noise can actually lead in overestimation of the embedding dimension. In order to mitigate these effects and keep a consist methodology across a large dataset, the presented methodology adopts fixed values for the embedding parameters rather than optimizing them per recording or window. The decision to use constant values is influenced by applied precedents in the EEG literature. McSharry et al.[50] have applied with success fixed embedding parameters in their multi-channel scalp EEG seizure research, arguing that nonlinear methods must justify their complexity over simpler linear benchmarks. In our case, fixed parameters ensure consistency across the large CHB-MIT dataset and help avoid overfitting to local dynamics that may not generalize.

7.3 Threshold selection

Radius fraction R is utilized for determination of the percentage of mean diameter of the reconstructed phase space where a recurrence can occur. In order to estimate and standardize R , an exploration of its effect on CRPs and RR/DET metrics is performed. By keeping constant $\tau = 1$ and $m = 3$ CRPs are generated by selecting random patients and random recording windows of the dataset, while computing the mean channel-wise recurrence rate and mean channel-wise determinism from the 22x22 CRPs features.

Results of RR and DET are presented in the following table. The different explored values for radius fraction are set to be 0.1, 0.15, 0.20 and 0.30 for this experiment.

Table 3: Comparison of RR and DET values for different radius (R) values

Recording	R	RR (%)	DET (%)	Window
patient_24	0.10	9	77.36	Normal
patient_24	0.15	16.7	85.3	Normal
patient_24	0.20	23.8	89	Normal
patient_24	0.30	36.4	93.25	Normal
patient_24	0.10	16.35	97.6	Epileptic
patient_24	0.15	26	99	Epileptic
patient_24	0.20	35.16	99.44	Epileptic
patient_24	0.30	51.9	99.74	Epileptic
patient_10	0.10	10.6	83.5	Normal
patient_10	0.15	16.54	86.8	Normal
patient_10	0.20	22.13	88.97	Normal
patient_10	0.30	32.5	92.87	Normal
patient_10	0.10	14.94	93.6	Epileptic
patient_10	0.15	22.9	95.7	Epileptic
patient_10	0.20	30.3	96.41	Epileptic
patient_10	0.30	43.59	97.16	Epileptic
patient_8	0.10	7.08	59.26	Normal
patient_8	0.15	11.82	65.12	Normal
patient_8	0.20	16.33	67.59	Normal
patient_8	0.30	24.76	71.73	Normal
patient_8	0.10	16.4	98.3	Epileptic
patient_8	0.15	25	99.46	Epileptic
patient_8	0.20	33.1	99.60	Epileptic
patient_8	0.30	47.71	99.76	Epileptic
patient_1	0.10	18.31	96.02	Normal
patient_1	0.15	27.83	97.70	Normal
patient_1	0.20	36.61	97.89	Normal
patient_1	0.30	52.08	98.89	Normal
patient_1	0.10	20.89	96.11	Epileptic
patient_1	0.15	33.93	98.46	Epileptic
patient_1	0.20	45.61	99.22	Epileptic
patient_1	0.30	64.64	99.68	Epileptic

As it can be observed, both RR and DET increase while the radius fraction R increases for all patients/windows combinations, since by having a larger radius there are more points to be considered as recurrent in the phase space. Epileptic windows present consistently higher RR and DET values when compared with normal windows at the same R and same patients. It should be noted that there is inter-patient variability also, suggesting that optimal radius selection may benefit

from patient-specific tuning. Additionally, DET values in epileptic windows approach saturation near 100% as R increases, a fact that suggests having a moderate radius fraction (e.g., $R = 0.15\text{--}0.20$) could provide a better balance between sensitivity and specificity in CRP analysis for the determinism metric.

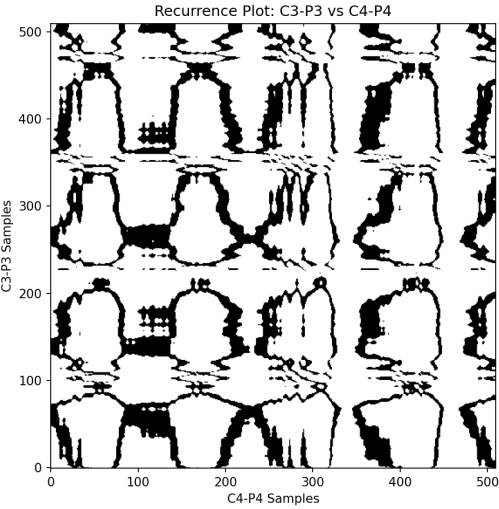
7.4 Cross Recurrence Quantification Analysis (CRQA)

CRQA quantifies the recurrent patterns between pairs of EEG channels within each time window. The `pyrqa` library is used with OpenCL acceleration for efficient computation. The process is as follows:

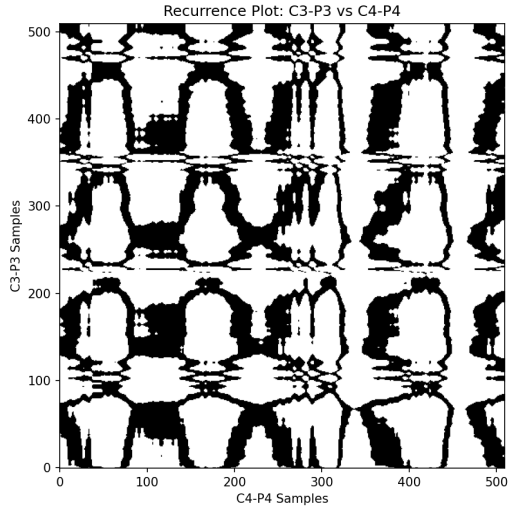
1. **Time Series Length Validation:** For each window pair, the lengths of the two time series are compared on having same length to ensure compatibility.
2. **Phase Space Reconstruction:** The time series are embedded into a phase space using the optimal τ and m , via the `TimeSeries` class in `pyrqa`.
3. **Radius Selection:** The radius for defining recurrence is computed using the Phase Space Separation (PSS) method (`pss` function). The maximum distances in the phase spaces of both channels are averaged to obtain a mean diameter, and the radius is set to 15% of this value (`radius_fraction=0.15`).
4. **CRQA Computation:** The `RQAComputation` class constructs a cross-recurrence matrix using a `FixedRadius` neighborhood, Euclidean metric, and Theiler corrector of 1. The computation yields 16 CRQA features, listed in Table 4, plus the segment label as the 17th feature.

Table 4: Quantitative measures computed by PyRQA

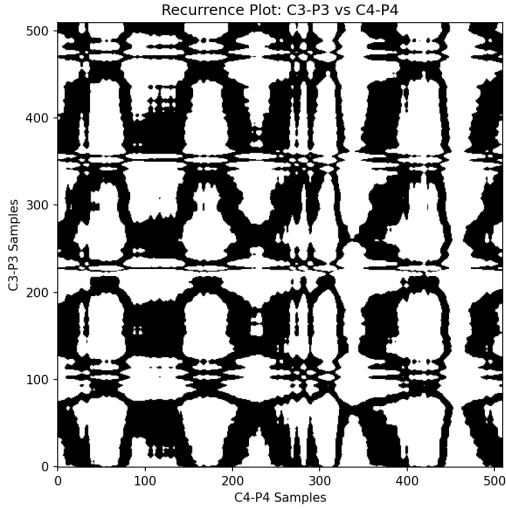
Metric	Abbreviation
Recurrence Rate	RR
Determinism	DET
Average Diagonal Line Length	L_{avg}
Longest Diagonal Line Length	L_{max}
Divergence	DIV
Entropy Diagonal Lines	H_{diag}
Laminarity	LAM
Trapping Time	TT
Longest Vertical Line Length	V_{max}
Average White Vertical Line Length	W_{avg}
Longest White Vertical Line Length	W_{max}
Longest White Vertical Line Divergence	W^{-1}_{max}
Entropy Vertical Lines	H_{vert}
Entropy White Vertical Lines	H_{wvert}
Ratio of Determinism to Recurrence Rate	DET/RR
Ratio of Laminarity to Determinism	LAM/DET



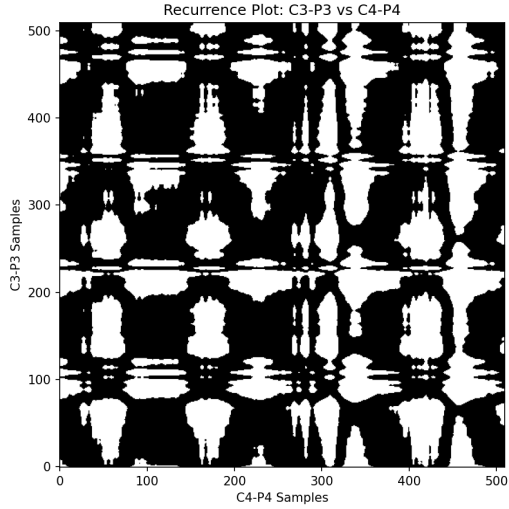
(a) radius fraction = 0.1



(b) radius fraction = 0.15



(c) radius fraction = 0.2



(d) radius fraction = 0.3

Figure 9: CRPs for the selected EEG channel pairs, for an epileptic window. (a) R = 0.1 (b) R = 0.15
(c) R = 0.2 (d) R = 0.3

7.5 Algorithm Summary

The methodology is summarized in Algorithm 1, which is describing the CRQA computation for each window and channel pair.

Algorithm 1 Cross Recurrence Quantification Analysis (CRQA) for EEG Windows

```

1: Input: EEG windows  $\{X_{c,w}\}$  for channels  $c \in C$ , windows  $w = 1, \dots, N_w$ , where  $N_w$  = number of windows, number of electrodes  $N_e$ 
2: Output: CRQA feature matrix  $M$  of shape  $(N_w, N_e, N_e, 17)$ 
3: for each window index  $w = 1$  to  $N_w$  do
4:   for each channel pair  $(c_1, c_2) \in C \times C$  do
5:     Time Series Preparation
6:       Ensure the input time series  $X_{c_1,w}$  and  $X_{c_2,w}$  have the same length
7:       Set embedding parameters
8:       Set  $\tau = 1$ 
9:       Set  $m = 3$ 
10:      CRQA Computation
11:        Construct cross-recurrence plot for  $(X_{c_1,w}, X_{c_2,w})$  using PyRQA with:
12:          Fixed radius neighborhood  $r = 0.15 \times$  mean diameter, Euclidean metric, Theiler corrector = 1
13:          Extract 16 CRQA features: {RR, DET,  $L_{\text{avg}}$ ,  $L_{\text{max}}$ , DIV,  $H_{\text{diag}}$ , LAM, TT,  $V_{\text{max}}$ ,  $W_{\text{avg}}$ ,  $W_{\text{max}}$ ,  $W_{\text{max}}^{-1}$ ,  $H_{\text{vert}}$ ,  $H_{\text{wvert}}$ , DET/RR, LAM/DET}
14:      Feature Storage
15:        Append window label  $l_w$  to features
16:        Store features in  $M[w, c_1, c_2, :]$ 
17:      end for
18:    end for
19:  Return RQA feature matrix  $M$ 

```

7.6 Feature aggregation

The resulting RQA feature matrix has dimensions $[N_w, N_e, N_e, 17]$, where N_w is the number of windows, N_e is the number of channels, and 17 represents the 16 CRQA features plus the segment label(epileptic or normal). To summarize CRQA features across all channel pairs for each window, a mean feature matrix is computed by averaging the 16 CRQA features across all channel pairs, resulting in a final matrix (**mean_feature_matrix**) of shape $[N_w, 17]$, where the last column retains the window label. This matrix summarizes the average dynamical interactions within each window.

It should be noted that prior to the averaging, all feature metrics that presented invalid numerical values(such as *nan* or *inf*) were set to 0. This can occur each time where the recurrence rate is 0 or it has a value very close to zero, having a consequence to propagate invalid values in the set of {Determinism, Average Diagonal Line Length, Divergence, Laminarity, Trapping Time, DET/RR and LAM/DET} features computed by the PyRQA computation.

7.7 Class Imbalance in CRQA-EEG Data

Real world EEG datasets for epileptic seizure detection are by their nature imbalanced, reflecting the transient nature of seizure's occurrence in contrast to normal brain activity. In our aggregated dataset, derived from CRQA features across the multi-channel EEG recordings, the distribution is stark: 98.99% normal segments (89633 examples) against 1.01% epileptic (901 examples), yielding a ~99:1 ratio. This skew is a well known challenge in machine learning which is often referred as the "imbalanced learning problem".

As discussed by He and Garcia[54], standard classifiers, (f.e, Random Forest), can achieve misleadingly high accuracy (~99%) by over-predicting the majority class, resulting in poor recall for epileptic events. In RQA based models, this bias can minimize the effects of discriminative patterns, such as the elevated determinism (DET) or laminarity (LAM) in epileptic signals with false negatives posing ethical risks, delaying interventions, while naive oversampling (f.e, duplication) incorporates the risk of overfitting in minority RQA features.

In order to mitigate this, data-level resampling strategies can be employed, focusing on synthetic generation to enrich the minority class without discarding information gained from the majority class.

Table 5: Class distribution in the preprocessed CRQA-EEG dataset, highlighting severe imbalance.

Class	Percentage (%)	Count
Normal (0)	98.99	89,633
Epileptic (1)	1.01	901

To address the highly imbalanced dataset, three main methods can be employed and are dominant in bibliography.

Regarding the data level, methods on creating more synthetic samples belonging to the minority class exist, such as Synthetic Minority Over-sampling Technique (SMOTE). With this method it is possible to generate synthetic epileptic samples by interpolating between minority instances and their k-nearest neighbors in feature space [51]. Unlike random duplication, SMOTE promotes diversity, reducing overfitting risks in low-sample regimes.

On the classification algorithm level, specific weights and costs can be used in order to favor the minority class. For instance, many classifiers, such as SVM or tree-based models, allow for the assignment of class weights. These weights are usually set to be inversely proportional to the class frequencies, pushing the model towards paying more attention on errors made on the minority class on the training phase. In the same manner, cost-sensitive learning methods define a higher penalty for misclassifying minority class samples, optimizing for a cost function that reflects the real-world imbalance.

As a third alternative, ensemble methods leverage two or more base models to swift the skew of bias towards the majority class. An ensemble can integrate different classifiers and employ different aggregation strategies for its final classification decision, such as majority vote, weighted votes or stacking.

7.8 Stratified Downsampling Methodology for a Balanced Dataset

The dataset that has been derived by concatenating all (`mean_feature_matrices`) of every recording, presents severe class imbalance as it has been presented in Table 5. This is a common challenge for epileptic seizure detection using long and continuous EEG recordings. Initial analysis revealed a distribution of approximately 99% normal segments versus 1% epileptic segments across all the processed EEG recordings. This kind of extreme imbalance would lead into classifier bias on the favor of the majority class, having as a consequence high accuracy by simply predicting "normal" for all instances.

In order to tackle this imbalance problem, a patient-specific stratified downsampling approach has been used for the normal segments. For each `mean_feature_matrix` corresponding to each one of the 136 recordings, its normal segments have been grouped into 40 continuous splits, respecting their timing order.

The selection of $S = 40$ splits was empirically determined to achieve approximate class balance while ensuring each averaged vector represents a meaningful sample of normal brain activity.

Patient-specific processing maintains inter-patient physiological variability and avoids the data leakage between patients, which is essential for developing classification models later.

The processed data, stored as `.npy` files in a dedicated directory, serves as the foundation for all subsequent machine learning experiments described in the following sections and the class distribution can be observed in the following table.

Table 6: Class distribution after the application of normal's segments downsampling.

Class	Percentage (%)	Count
Normal (0)	49.46	5440
Epileptic (1)	50.54	5558

8 Classification - Experimental

In this section, the classification methodology which is applied on the generated dataset is described.

8.1 Classification Performance Metrics

To evaluate the performance of any classifier, several commonly applied metrics can be derived from the confusion matrix.

8.1.1 Confusion Matrix

A confusion matrix is a tabular summary of the number of correct and incorrect predictions achieved by the classifier. For a binary classification problem, it is typically represented as follows:

	Predicted Positive	Predicted Negative
Actual Positive	True Positive (TP)	False Negative (FN)
Actual Negative	False Positive (FP)	True Negative (TN)

8.1.2 Accuracy

Accuracy represents the proportion of correctly classified instances over the total number of instances:

$$\text{Accuracy} = \frac{TP + TN}{TP + TN + FP + FN} \quad (7)$$

8.1.3 Sensitivity / Recall

Sensitivity, also known as recall or true positive rate, measures the proportion of actual positives correctly identified:

$$\text{Sensitivity} = \frac{TP}{TP + FN} \quad (8)$$

8.1.4 Specificity

Specificity, also known as true negative rate, measures the proportion of actual negatives correctly identified:

$$\text{Specificity} = \frac{TN}{TN + FP} \quad (9)$$

8.1.5 Precision

Precision measures the proportion of predicted positives that are correctly identified:

$$\text{Precision} = \frac{TP}{TP + FP} \quad (10)$$

8.1.6 F1-Score

The F1-score is the harmonic mean of precision and recall, providing a single metric that balances both:

$$\text{F1-score} = 2 \cdot \frac{\text{Precision} \cdot \text{Sensitivity}}{\text{Precision} + \text{Sensitivity}} \quad (11)$$

The interpretation of these metrics provide an evaluation of classifier's performance.

Leave-One-Subject-Out (LOSO) Cross-Validation

To obtain an unbiased estimate of the classifier's ability to generalize across different individuals, the experiments conducted utilize the Leave-One-Subject-Out (LOSO) cross-validation. In LOSO, data from one subject (patient) are held out as the test set, while data from all remaining subjects are used for training. The model is always evaluated on unpresented data to it. This process is repeated iteratively until every subject has served as the test set exactly once.

The LOSO method is useful for physiological datasets, where the inter-patient variability is often observable. Training and evaluating on different subjects ensures that the classifier does not inadvertently learn subject-specific patterns that would not generalize to unseen individuals. Consequently, LOSO provides a conservative and realistic estimate of real-world performance in clinical scenarios, where new patient data must be classified without retraining the model.

For each hyperparameter configuration explored, the SVM model was trained and tested across all LOSO folds. Performance metrics (accuracy, sensitivity, specificity, and F1-score) are computed for each held-out subject and then averaged to obtain a final estimate. Using this evaluation ensures that the reported results reflect consistent behavior across subjects and are not biased by any single individual.

8.2 Support Vector Machine (SVM) Classifier

Support Vector Machines (SVMs) belong to the so-called supervised learning models used for classification and regression purposes. The idea behind the SVM algorithm is the construction of a hyperplane or a set of hyperplanes in a high-dimensional space that can be used to separate different classes. The main objective of SVM is to find the hyperplane that maximizes the margin between different classes, which eventually results into improving the generalization ability of the classifier.

Given a training dataset $\{(x_i, y_i)\}_{i=1}^N$, where $x_i \in \mathbb{R}^d$ is a feature vector and $y_i \in \{-1, 1\}$ is the class label, SVM solves the following optimization problem:

$$\min_{w,b} \frac{1}{2} \|w\|^2 \quad \text{subject to} \quad y_i(w^\top x_i + b) \geq 1, \quad i = 1, \dots, N \quad (12)$$

where w is the weight vector and b is a bias term. For non-linearly separable data, kernel functions can be used to map the input features to a higher-dimensional space where a linear separation is possible.

As shown in Figure 10, the SVM optimization problem seeks the hyperplane that maximizes the margin between classes. Hyperplane H1 fails to separate the data, where on the other hand, H2 and H3 are both valid classifiers. The SVM selects H3 though, over H2 because it achieves the maximum possible margin ($\rho_{H3} > \rho_{H2}$), leading to better generalization performance.

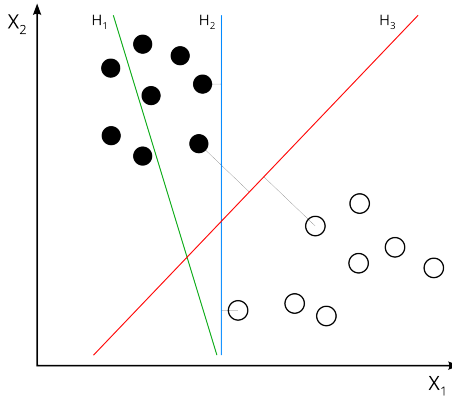


Figure 10: SVM maximization of margin among different classes

In order to evaluate the SVM classifier, an exploration of its hyperparameters was conducted to reveal their effects on the classification performance. SVMs have configurable options that can control the geometry of the decision function, the complexity of the model, the smoothness of the boundary, and the optimization procedure.

The *kernel function* defines the transformation that maps the input data into a higher-dimensional feature space, where a linear separating hyperplane may possibly exist. The following kernels were examined:

- **Polynomial Kernel (poly):**

$$K(\mathbf{x}, \mathbf{x}') = (\gamma \mathbf{x}^\top \mathbf{x}' + \text{coef0})^{\text{degree}}$$

This kernel allows highly flexible, non-linear decision boundaries whose shape is controlled by the degree of the polynomial.

- **Linear Kernel (linear):** Represents a standard linear classifier without non-linear mapping.
- **RBF Kernel (rbf):** A Gaussian kernel capable of modelling highly complex, non-linear structures.

The regularization parameter $C > 0$ is in the control of the trade-off among two quantities. The maximization of the margin and minimization of the classification error on the training data. Using a small value of C produces a wider margin among classes and a more regularized model, possibly at the cost of more misclassifications. By using a larger value of C , model enforces more strict classification of training samples by heavily penalizing misclassified points.

The parameter γ (kernel scale) controls the influence of individual training samples when using kernels such as RBF or polynomial. In the polynomial kernel, γ acts as a scaling factor on the dot product:

$$(\gamma \mathbf{x}^\top \mathbf{x}' + \text{coef0})^{\text{degree}}.$$

A small γ produces smoother, less complex decision boundaries, while a large γ leads to highly localized and complex surfaces.

For the polynomial kernel, the degree determines the maximum order of interactions between features. Higher degrees have as result more complex decision boundaries, but also increase the risk of overfitting and are computational more heavy procedures.

The coefficient `coef0` shifts the kernel function and influences how much the model relies on higher-order terms versus lower-order interactions. In polynomial kernels, tuning `coef0` can significantly modify the boundary shape.

Shrinking is an optimization heuristic, that attempts to speed up the training procedure by temporarily ignoring variables that are unlikely to affect the solution. When `shrinking=False`, the full optimization problem is solved without heuristics. Disabling shrinking may lead to more stable behavior in some datasets, at the cost of slower training.

The parameter `tol` defines the tolerance for the stopping criterion of the optimization algorithm. Smaller values lead to more precise convergence but increase computational time, while larger values may speed up optimization at the cost of slightly less accurate solutions.

To address class imbalance, the SVM was configured with `class_weight='balanced'`, which automatically scales the penalty applied to each class inversely to its frequency. This ensures that minority classes contribute equally to the loss function.

Table 7: SVM Classification Results

Kernel	C	γ	Mean Acc.	Std	Mean Sens.	Std	Mean Spec.	Std	Mean F1	Std
linear	0.1	scale	0.75	0.138	0.677	0.22	0.826	0.156	0.697	0.205
linear	1.0	scale	0.759	0.139	0.68	0.22	0.824	0.159	0.698	0.205
linear	10.0	scale	0.758	0.139	0.679	0.219	0.825	0.159	0.697	0.204
rbf	0.1	scale	0.821	0.112	0.764	0.182	0.867	0.136	0.782	0.156
rbf	0.1	auto	0.821	0.112	0.764	0.182	0.867	0.136	0.782	0.156
rbf	1.0	scale	0.851	0.09	0.817	0.138	0.881	0.134	0.824	0.125
rbf	1.0	auto	0.851	0.09	0.817	0.138	0.881	0.134	0.824	0.125
rbf	10.0	scale	0.881	0.074	0.879	0.098	0.889	0.122	0.863	0.102
rbf	10.0	auto	0.881	0.074	0.879	0.098	0.889	0.122	0.863	0.102
poly	0.1	scale	0.764	0.137	0.564	0.237	0.946	0.081	0.658	0.215
poly	0.1	auto	0.764	0.137	0.564	0.237	0.946	0.081	0.658	0.215
poly	1.0	scale	0.786	0.126	0.635	0.217	0.929	0.098	0.712	0.186
poly	1.0	auto	0.786	0.126	0.635	0.217	0.929	0.098	0.712	0.186
poly	10.0	scale	0.809	0.106	0.703	0.194	0.907	0.108	0.757	0.153
poly	10.0	auto	0.809	0.106	0.703	0.194	0.907	0.108	0.757	0.153

8.3 Random Forest Classification

Random Forest is an ensemble method of learning, which constructs a number of decision trees during training and produces the final prediction utilizing the majority voting concept (for the classification) alongside all the individual trees. Each of the trees belonging in the forest, is trained using a bootstrapped subset of the training data, while at each split only a random subset of the available features is considered. By using this dual randomization, decorrelates the trees and significantly improves robustness against overfitting.

Let $\mathcal{D} = \{(\mathbf{x}_i, y_i)\}_{i=1}^N$ denote the training dataset, where $\mathbf{x}_i \in \mathbb{R}^d$ represents the feature vector and $y_i \in \{0, 1\}$ the corresponding class label. Each decision tree $h_k(\mathbf{x})$ is trained on a bootstrapped

subset $\mathcal{D}_k \subset \mathcal{D}$. The Random Forest prediction is then obtained as

$$\hat{y} = \text{mode}(h_1(\mathbf{x}), h_2(\mathbf{x}), \dots, h_K(\mathbf{x})), \quad (13)$$

where K is the total number of trees in the ensemble.

Random Forest classifiers have the ability of handling high-dimensional feature spaces, nonlinear decision boundaries, and noisy measurements without requiring strong assumptions about the underlying data distribution. Furthermore, they provide resistance to outliers and feature scaling, which is particularly useful in biomedical and physiological signals.

8.3.1 Hyperparameters

Related to the performance of a Random Forest classifier, there is a number of structural hyperparameters that control and regulate model's complexity and generalization ability. In this study, the following hyperparameters were systematically explored under a Leave-One-Subject-Out (LOSO) cross-validation method:

- **Number of Trees ($n_{\text{estimators}}$):** This parameter defines the total number of decision trees in the ensemble. Larger values generally improve classification stability at the expense of increased computational cost. The tested values were

$$n_{\text{estimators}} \in \{100, 300, 500\}.$$

- **Maximum Tree Depth (max_depth):** This parameter limits the maximum depth of each tree. Constraining tree depth prevents overly complex decision boundaries and reduces overfitting. The evaluated values were

$$\text{max_depth} \in \{\text{None}, 5, 10\}.$$

- **Minimum Samples for Node Splitting (min_samples_split):** This parameter specifies the minimum number of samples required to split an internal node. Larger values enforce smoother decision surfaces and increase regularization:

$$\text{min_samples_split} \in \{2, 5\}.$$

- **Minimum Samples in Leaf Nodes (min_samples_leaf):** This parameter constrains the minimum number of samples that must exist in a terminal node. It directly controls the granularity of leaf-level decisions and improves generalization in small-sample regimes:

$$\text{min_samples_leaf} \in \{1, 2\}.$$

- **Maximum Features per Split (max_features):** This parameter determines the number of candidate features randomly selected at each split. It introduces additional decorrelation between trees. The evaluated strategies were

$$\text{max_features} \in \{\text{sqrt}, \text{log2}\}.$$

For each hyperparameter configuration, the classifier was evaluated using a Leave-One-Subject-Out (LOSO) protocol to ensure strict subject-independent validation. Performance was quantified using accuracy, sensitivity (recall), specificity, and F1-score. Mean and standard deviation values across all subjects were computed to assess both predictive performance and inter-subject stability.

The Random Forest classifier does not require feature normalization and is resilient to monotonic feature transformations, making it particularly suitable for heterogeneous biomedical feature sets where absolute feature scaling may vary across subjects.

8.4 k-Nearest Neighbors Classification

The k-Nearest Neighbors (kNN) algorithm is a non-parametric, instance-based learning method that performs classification based on the similarity between input samples in the feature space. Unlike model-based classifiers, kNN does not explicitly construct a training model. Instead, all training samples are retained in memory, and predictions are made by identifying the k closest samples to a given test instance according to a distance metric.

Let $\mathcal{D} = \{(\mathbf{x}_i, y_i)\}_{i=1}^N$ denote the training dataset, where $\mathbf{x}_i \in \mathbb{R}^d$ represents a feature vector and $y_i \in \{0, 1\}$ the corresponding class label. For a test sample \mathbf{x} , the classifier computes the distance $d(\mathbf{x}, \mathbf{x}_i)$ between \mathbf{x} and all training samples. The set $\mathcal{N}_k(\mathbf{x})$ containing the k closest neighbors is then determined, and the predicted class label is obtained by majority voting:

$$\hat{y} = \text{mode}\{y_i \mid \mathbf{x}_i \in \mathcal{N}_k(\mathbf{x})\}. \quad (14)$$

Due to its simplicity and non-parametric nature, kNN is well suited for biomedical signal classification tasks where class boundaries may be highly nonlinear and difficult to model explicitly. The method adapts naturally to complex data distributions and performs particularly well when representative samples are available in the feature space. However, kNN is sensitive to feature scaling and noise, making proper preprocessing essential.

8.4.1 Hyperparameters

The behavior and performance of the kNN classifier are controlled by a small number of hyperparameters that regulate neighborhood size, distance computation, and voting strategy. In this study, the following hyperparameters were systematically evaluated using a grid-search framework under a Leave-One-Subject-Out (LOSO) cross-validation protocol:

- **Number of Neighbors (k):** This parameter specifies how many neighboring samples participate in the voting process. Small values of k increase sensitivity to local variations and noise, while larger values produce smoother decision boundaries:

$$k \in \{3, 5, 7, 9, 11\}.$$

- **Distance Metric:** This parameter determines how similarity between feature vectors is computed. The evaluated distance metrics were:

$$\text{metric} \in \{\text{euclidean}, \text{manhattan}\}.$$

The Euclidean distance emphasizes geometric proximity, while the Manhattan distance is more robust to outliers in high-dimensional spaces.

- **Voting Strategy (Weighting):** This parameter controls how votes from neighboring samples are weighted:

$$\text{weights} \in \{\text{uniform}, \text{distance}\}.$$

Uniform weighting assigns equal importance to all neighbors, whereas distance-based weighting assigns higher influence to closer neighbors.

- **Neighbor Search Algorithm:** This parameter determines the method used for neighbor retrieval:

$$\text{algorithm} \in \{\text{auto}, \text{ball_tree}, \text{kd_tree}\}.$$

This choice affects computational efficiency, particularly for high-dimensional feature spaces.

For each hyperparameter configuration, classifier performance was evaluated using a Leave-One-Subject-Out (LOSO) validation scheme to ensure strict subject-independent testing. Performance was assessed using accuracy, sensitivity, specificity, and F1-score, with mean and standard deviation computed across all subjects.

Since kNN relies directly on distance computations, all features were standardized prior to classification to ensure equal contribution of all feature dimensions. This normalization step is essential in heterogeneous biomedical feature sets, such as EEG-derived CRQA features or radiomic descriptors, where feature scales may differ significantly.

9 Classification Results and Comparative Analysis

This section presents the classification performance obtained using three different machine learning models: Support Vector Machines (SVM), Random Forests (RF), and k-Nearest Neighbors (kNN). All models were evaluated under a strict Leave-One-Subject-Out (LOSO) cross-validation protocol to ensure subject-independent testing. Performance was quantified using accuracy, sensitivity, specificity, and F1-score, and results are reported as mean and standard deviation across all subjects.

9.1 Overall Performance Comparison

Across all experiments, the three classifiers demonstrated distinct performance characteristics. The SVM classifier exhibited strong generalization ability, achieving consistently high accuracy and F1-scores across subjects. This behavior is attributed to its ability to construct optimal separating hyperplanes in high-dimensional feature spaces, which is particularly advantageous for CRQA-based and biomedical feature representations.

The Random Forest classifier demonstrated stable and well-balanced performance across all evaluation metrics. Its ensemble-based structure allowed it to effectively capture nonlinear feature interactions while maintaining robustness against overfitting. Additionally, Random Forest achieved competitive sensitivity and specificity values, indicating reliable discrimination between the two classes.

The kNN classifier showed competitive performance in several configurations but exhibited higher variability across subjects, as reflected by increased standard deviation values. This behavior is expected, as kNN is a distance-based, instance-driven classifier that is highly sensitive to feature scaling, class distribution, and local neighborhood structure. Despite this sensitivity, kNN achieved satisfactory performance when appropriate feature normalization and neighborhood sizes were applied.

9.2 Sensitivity–Specificity Trade-Off

An important observation across all classifiers is the inherent trade-off between sensitivity and specificity. SVM tended to favor balanced performance, achieving comparable sensitivity and specificity values. Random Forest often demonstrated slightly higher sensitivity, suggesting improved detection of positive class instances, while kNN occasionally favored specificity depending on neighborhood size and distance metric.

This trade-off is particularly important in biomedical classification tasks, where false negatives and false positives carry different clinical implications. The ability of the Random Forest and SVM classifiers to maintain stable sensitivity under LOSO validation suggests strong potential for subject-independent deployment.

9.3 Model Stability and Generalization

Model stability across subjects was assessed through the standard deviation of classification metrics. Among the three classifiers, Random Forest generally exhibited the lowest performance variance, indicating superior robustness to inter-subject variability. SVM also demonstrated stable behavior, while kNN showed the highest variance, reflecting its dependence on local sample distributions.

The LOSO protocol imposes a highly challenging validation setting, as the classifier is required to generalize to completely unseen subjects. The fact that all three models achieved consistent above-chance performance under this protocol indicates that the extracted feature representations contain discriminative subject-independent information.

9.4 Computational Considerations

From a computational standpoint, SVM required the most careful hyperparameter tuning due to the sensitivity of its regularization and kernel parameters. Random Forest incurred higher training time due to its ensemble nature but offered faster inference. In contrast, kNN required minimal training time but significantly higher inference cost, as distance computations must be performed against the entire training dataset for each test sample.

These computational characteristics further highlight the trade-offs between the three models in terms of real-time deployment potential and scalability.

9.5 Summary of Findings

In summary, SVM achieved the strongest overall generalization performance, Random Forest provided the best stability and robustness across subjects, and kNN offered a simple but less consistent alternative. The complementary strengths of these classifiers validate the robustness of the proposed feature extraction and evaluation pipeline. The agreement in performance trends across multiple classifiers further supports the reliability of the reported results.

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