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Fuzzy Clustering for Exploratory Analysis of EEG Event-Related Potentials

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Abstract—We introduce an analysis method for electroencephalography (EEG) data, focused on Event-Related Potentials (ERPs). Our approach is unsupervised and makes use of a fuzzy clustering algorithm based on the possibilistic framework, and includes a data-driven noise and artifact rejection phase. Our contribution provides a general analysis tool, applicable to any ERP data set, which can uncover the data set’s internal structure. The fuzzy clustering algorithm is the core of our method, since its fine-grained membership grades how much a sample belongs to a given cluster, making the method applicable even when groups have a certain overlap. Prior to the clustering step, we apply weights to the feature vectors, optimizing them in order to enhance the variance within the dataset, and we extract time-window interval based features inspired by interval arithmetic. We apply the data processing workflow to the analysis a set of ERPs recorded during an emotional Go/NoGo task. We evaluate the performance of the unsupervised analysis by computing a measure based on the clusterization rate of trials in different experimental conditions. The results on the studied data set show that the proposed method obtains a difference of clusterization rate of 69% in Go vs. NoGo trials, when weights and interval-features are applied to the data, improving previous work not including weights and interval-features which had a rate of 31%. Furthermore, when compared with the standard Fuzzy c-means, our proposed possibilistic clustering algorithm outperforms it in terms of clusterization rate. We also examine the effect of pre-processing the data with Independent Component Analysis and removing noise-related components, and observe that this does not improve significantly the obtained results. These findings demonstrate that our proposed method provides a valuable data processing workflow robust to EEG artifacts and able to produce a clustering that is coherent with the experimental conditions represented in the ERP dataset.

Index Terms—EEG, ERP, possibilistic clustering, unsupervised, interval features

I. INTRODUCTION

The brain’s electrical activity is related to motor and cognitive tasks in complex ways. Shedding light on the causal links between tasks and observed regularities in brain activity can obviously benefit research in psychology and related disciplines. Direct brain interfaces that use brain activity as a control signal are also starting to spread in several fields, including virtual reality, advanced prosthetics, and rehabilitation. Their efficacy is crucially dependent on the ability to analyze electrical signals from the brain. This work presents a contribution to the task of discovering such links.

One of the most widely applied experimental methods for investigating human brain activity is scalp electroencephalography (EEG). This technique is non-invasive, because it uses external electrodes which are placed at standard locations on the scalp. The exposition to specific stimulations (e.g., visual, auditory) causes brain activation in the form of transient electric potentials which can be measured with scalp electrodes and averaged over repeated trials [1]. The resulting EEG signals that are triggered by a specific event are called Event-Related Potentials (ERPs).

The analysis of EEG data can be used to infer behavioral and personality data. As the EEG signal is typically high-dimensional and noisy, the conventional analysis techniques for EEG data rely significantly on the experimenter’s ability to discard outliers and observe peaks and waves that are associated with different neural processes. The usual approach to stimulus-related EEG recordings is the calculation of Event-Related Potentials (ERPs), i.e., the averaging of multiple repetitions of a trial in the same experimental conditions. In fact, the assumption for ERP analysis is that the brain always reacts with the same activation as a response to a given stimulus, but the signal recorded with EEG also includes many other components, given by concurrent brain processes, noise, muscular artifacts, etc. Averaging the repeated trials allows to make the activation (the evoked potential) emerge and become observable. Neuroscientists have discovered that different stimuli cause specific waves or components in the evoked potentials, and therefore ERPs can be used to understand the brain processes that are triggered by a certain stimulus [2]. On the other hand, the averaging of many trials can cause a significant loss of information, as the noise (e.g. muscular artifacts caused by eye blinks or movements) can have an amplitude several order of magnitudes larger than the signal. To address this, approaches based on single-trial analyses [3] have been used, with the goal of including the contribution of the entire EEG recording [4].

In this study, we define an unsupervised processing workflow based on fuzzy clustering, in order to model differences and similarities between samples in a dataset, using a data-driven feature-extraction strategy. Our goal is to build a general-purpose unsupervised analysis workflow for Event-
Related Potentials. Most conventional EEG analysis pipelines have a number of processing steps requiring expert evaluation of the results. For example, the widely used Independent Component Analysis relies on an expert inspection of the components identified by the algorithm in order to decide whether they are signal or artifact. We propose an alternative approach, which aims to extract exploratory information from an EEG (ERP) dataset, minimizing human expert intervention. Our method does not need a specialist to evaluate the artifact removal, since it can extract information despite the presence of noise. The clusters that are obtained with our technique are a way for the EEG researcher to obtain a partition of the given ERP set which was not known, possible groupings that were not previously studied or observed. To avoid the limitations of ERPs described above, we consider the original EEG signal (the whole set of trials) as the starting point of our analysis pipeline and implement a noise-rejection method to remove outlier epochs in our data processing workflow. Our method is not an alternative to (semi-)supervised approaches, but rather constitutes a possible first level of analysis, which can be followed by the application of other methods targeted, for instance, at classification. The results of the clustering can inform the algorithm design. To avoid the limitations of ERPs described above, we consider the original EEG signal (the whole set of trials) as the starting point of our analysis pipeline and implement a noise-rejection method to remove outlying artifacts in our data processing workflow.

In cluster analysis, the choice of a suitable number of clusters is one of the most important questions. Different methods for determining the optimal number have been proposed, included randomized algorithms based on Chebyshev polynomial approximation [5]. In our approach, we choose a number of clusters that is 2 to 3 times the expected number and then merge overlapping clusters. The merging procedure relies on measuring overlap with a fuzzy version of the Jaccard index. This lets our method determine a suitable number of clusters with a data-driven approach.

Previous work on this framework was presented in [6] as a conference paper. Here we present a significantly updated treatment which includes a number of new methods. In particular:

- we apply weights to the sample vectors and we optimize those weights in order to maximize the variability of the features within the dataset,
- we define interval-valued features computed on time windows of different lengths in order to improve the comparison of signals.

The data processing workflow is outlined in Fig. 1. The remainder of this article is structured as follows: Section II presents the state of the art and Sections III to V introduce the theoretical tools that we are using: Graded Possibilistic Clustering, the Jaccard Index and Weighted Interval-valued Features. The dataset that we consider as a case study is presented in Section VI. The way the data processing workflow is applied to the example dataset is the subject of Section VII. The remaining Sections VIII–IX present the results, discussions and conclusions.

It should be noted that in this article we apply the proposed method to a specific set of ERPs (from an experimental paradigm based on an emotional Go/NoGo task), but the method can be applied without significant changes to the analysis of any ERP-dataset.

II. STATE OF THE ART

Electroencephalography is the most widely used method to record human brain activity with high temporal resolution in a non-invasive way in clinical and research contexts, and recently in consumer-grade wearable devices. Despite being inexpensive and easy to use, EEG is characterized by a high noise-to-signal ratio, caused by artifacts of different origin (e.g., muscular, electromagnetic, or caused by the poor adherence of the electrodes on the scalp giving too high impedance). Most EEG studies include in their method a manual phase of artifact detection and rejection of the samples that are too noisy: in case of ERP studies, the signals obtained in multiple trials are averaged, and the resulting waves are examined in order to identify peaks associated to specific brain processes. The cognitive processes in the brain are caused by the coordinated activation of an extremely high number of interconnected neuron, whose synchronous combined electrical fields are strong enough to be measured on the scalp. Therefore, ERPs can be seen as the brain response to a specific stimulus (e.g. visual or auditory). An ERP waveform’s duration is generally not greater than one second from the stimulus onset, and along its time course one can observe positive and negative peaks or deflections, which can be considered as the combination of a number of components, positive or negative, each with a specific onset. Neuroscientists have discovered a number of components that are associated to particular classes of stimuli, and such components are characterized by a specified latency, shape, and positive or negative magnitude [2].

Most classical ERP results are obtained via supervised methods, and rely heavily on the experimenter’s ability for artifact removal and for the interpretation of the wave components.
associated with the studied neural processes. Machine learning techniques have been recently applied to these problems [7]. This includes methods such as Echo State Networks [8], multilayer perceptron neural networks [9], feature-extraction based on time intervals [10], or dimension reducing methods such as Independent Component Analysis [11]–[13]. It should be noted that such approaches can be used to address specific classification problems and have the disadvantage of being dependent of the choice of a set of features, designed by the experimenter in order to capture a certain aspect of the data. As a consequence, the human supervision is moved to a different step of the analysis, opening the door to biased learning and over-fitting of the model on the training data.

### III. GRADED POSSIBILISTIC CLUSTERING

To obtain a reliable clustering of EEG signals, which are noisy and highly variable, we employ a fuzzy clustering approach; in particular we focus on a robust clustering method that is rooted in the Possibilistic approach of Krishnapuram and Keller [14], [15]. The **Graded Possibilistic Clustering** method [16] is a central clustering model which inherits outlier rejection properties from the possibilistic model, while at the same time avoiding their well-known issues related to convergence and to overlapping clusters, similarly to methods which include probabilistic constraint (as in Fuzzy c-means [17], [18], and Deterministic Annealing [19], [20]). In the current work, we introduce a new version of the algorithm, called Graded Possibilistic Clustering model (GPC-II), which is an improvement on the one presented in [16].

Let $X$ be a set of $n$ observations (or instances) $x_i$, for $l \in \{1, \ldots, n\}$, and let $C$ be a set of $c$ fuzzy clusters denoted $C_1, \ldots, C_c$. The clusters are represented via their centroids $y_j$, for $j \in \{1, \ldots, c\}$. We associate to each cluster a fuzzy cluster indicator (or membership) function $u_{lj} \in [0, 1] \subset \mathbb{R}$ to a given observation $x_i$.

We start by defining the **free membership** of an observation $x_i$ to cluster number $j$ as $v_{lj} = e^{-d_{lj}/\beta_j}$, where $d_{lj}$ denotes the distance between the $j$-th centroid and the observation $x_i$. The coefficient $\beta_j$ is a cluster-specific parameter of the model that plays a role in how data is represented by influencing cluster width.

The **total membership mass** for the same observation is then defined as:

$$\zeta_l = \sum_{j=1}^{c} v_{lj}. \quad (1)$$

The actual membership of observation $x_i$ to cluster $j$ is written as a function of the above quantities as:

$$u_{lj} = \frac{v_{lj}}{\zeta_l} \quad (2)$$

In this equation we employ as a normalizer the **generalized partition function**:

$$Z_l = \zeta_l^\alpha = \left( \sum_{j=1}^{c} v_{lj} \right)^\alpha, \quad \alpha \in [0, 1] \subset \mathbb{R} \quad (3)$$

In the expression for the generalized partition function, $\alpha$ is a parameter which controls the “possibility level” of the GPC-II model. The value $\alpha = 1$ corresponds to a totally probabilistic model, while $\alpha = 0$ results in a totally possibilistic one, without normalization. The intermediate values $0 < \alpha < 1$ allow for the grading of this model.

The following equation represents the relation between the GPC-II cluster centroids and the membership vectors:

$$y_j = \frac{\sum_{l=1}^{c} u_{lj} x_l}{\sum_{l=1}^{c} u_{lj}}. \quad (4)$$

The GPC-II model is trained by a Picard iteration of Eqs. 2 and 4 after an initial step of random initialization of centroids. It should be noted that, for $\alpha = 1$, the model has the same representation properties as Deterministic Annealing [19], [20], where the solution of a regularized minimization problem with constraints yields

$$u_{lj} = \frac{e^{-d_{lj}/\beta_j}}{\sum_{h=1}^{c} e^{-d_{lh}/\beta_h}}, \quad (5)$$

equivalent to (2) when $\alpha = 1$. In turn, this is equivalent to Fuzzy c-means, up to a non-linear transformation, as proved in [21]. When $\alpha = 0$, they are equivalent to those of Possibilistic c-means [15]) which is designed for robust clustering, since its clusters are not influenced by each other or by outliers. This comes at the cost of difficult convergence and collapsing clusters. The most interesting behavior is therefore obtained for $0 < \alpha < 1$. Whenever $\alpha$ is positive, there is a degree of competition between clusters which improves convergence, as in probabilistic models. On the other hand, memberships eventually vanish for points sufficiently far away from the centroids, as in the possibilistic case, for noise insensitivity. This trade off is under user control, and depends in a complex way on the nature of the data (dimensionality, density, clustering structure, distribution isotropy, distribution uniformity...), so it is usually selected by grid search.

Deterministic annealing can be implemented in the Graded Possibilistic Clustering model obtaining a variation named DAGPC-II. To do so, the model parameters $\beta_j$ are decomposed as $\beta_j = \beta b_j$, where the new parameter $\beta$ is the basis for the deterministic annealing procedure (small at the start of the algorithm and increasing each time the Picard iteration converges) and $b_j$ are the **relative** cluster scales. The values for the parameters $b_j$ can be obtained with heuristics such as the ones proposed in [15]. This makes it possible to for the Deterministic Annealing Graded Possibilistic Clustering method to benefit from the powerful optimization technique proposed in [19], [20]: after starting from clusters that fully overlap, it performs a hierarchical clustering by progressively splitting clusters as $\beta$ increases. “Natural” aggregations are therefore discovered.

### IV. MEASURING OVERLAP BETWEEN FUZZY CLUSTERINGS

The procedure presented requires a measure of overlap between clusters. Many approaches are available in the literature; a concise survey and an experimental comparison is presented in [22].

Several interesting indices to compare two clusterings are centered around counts of concordance patterns on pairs of data points, where any given pair can be in the same cluster...
in both clusterings, in the same cluster in only one of the two clusterings, or in different clusters in both clusterings. Methods for fuzzifying such indices are usually based on replacing counts with combinations of fuzzy cluster memberships [23]–[25].

In this study we adopt the Jaccard index, defined as follows:

$$J(A, B) = \frac{|A \cap B|}{|A \cup B|},$$

(6)

This measure is less sensitive to irrelevant matches (pairs in different clusters in both clusterings) than, for instance, the Rand index [26].

In the case of fuzzy clusters, we can use the membership function to express the fact that an element belongs to a cluster, i.e. a real quantity in the interval [0, 1]. In order to define the Jaccard index (Eq. (6)) in the fuzzy case, we adopt the following definitions of cardinality for the intersection and the union of fuzzy clusters:

$$|C_l \cap C_m| = \sum_{j=1}^{N} \min(u_{jl}, u_{jm}),$$

(7)

$$|C_l \cup C_m| = \sum_{j=1}^{N} \max(u_{jl}, u_{jm}),$$

(8)

where $N$ is the total number of samples. The resulting fuzzy Jaccard index is still a real value in the interval [0, 1] even for possibilistic partitions, provided that the total membership of any data point does not exceed 1. With the definitions provided (Eq. (1) to (3)), this is true in realistic situations.

V. WEIGHTED INTERVAL-VALUED FEATURES

Our strategy for comparing ERP signals is based on the following considerations: Event-related peaks might occur at slightly different moment during the signal’s time course, and not all instants of the signals are equally meaningful. Therefore we use a kernel smoothing procedure based on a weight-mask, and we extract temporal, interval-valued features that summarize the data while accounting for this source of uncertainty.

A.Interval-valued features

Comparing ERPs and estimating their similarity can be challenging because of differences in the onset time of measured potentials. We address this issue by extracting interval-valued features based on time windows.

A similar approach was used in [10]. Here we employ a lower number of features which appears to be sufficient, in order to avoid model over-fitting.

Each signal is represented by means of multiple features, each of which summarizes a specific time window as an interval.

The time window features are extracted as follows: Along a signal vector $x \in \mathbb{R}^L$, we select onset points $a_j$ at regular intervals such that $a_{j+1} - a_j = R$. Starting from each onset point, we select offset points $b_{jk}$ such that $b_{jk} - a_j = 2^{k+1}$, for all the possible values of $k$. This procedure identifies time windows: $I_{jk} = [a_j, b_{jk}]$ of length $2^{k+1}$ and for each window we define the features:

$$M_{jk} = \max(x(i) \mid a_j \leq i \leq b_{jk}),$$

(9)

$$m_{jk} = \min(x(i) \mid a_j \leq i \leq b_{jk}),$$

(10)

namely the maximum and the minimum values of the signal vector in the time window. This way, the signal vector $x$ is represented by the vector formed by all the time window features $[m_{jk}, M_{jk}]$. An example of the obtained windows can be observed in Fig. 2.

B. Parameter tuning

To take into account the relative importance of each feature, in comparing two signals they are multiplied by suitable weights. These weights are convex, i.e., their sum is constrained to one, and their value is selected according to the following procedure.

Since the number of parameters equals the number of features (minus 1 due to the convexity constraint), we also introduce a method to optimize their value. The resulting weights can be obtained offline.

Let $x_l \in \mathbb{R}^L$ for $l \in \{1, \ldots, N\}$ be the ERP samples and $w \in \mathbb{R}^L$ a vector of weights such that

$$\sum_{j=1}^{L} w_j = 1.$$ (11)

(convexity of weights). Then we can define a weighted similarity matrix $K \in M_{N,N}(\mathbb{R})$ whose entry $(i,j)$ is $k(x_i, x_j, w)$ such that:

$$k(x_i, x_j, w) = \exp\left(-\sum_{l=1}^{L} w_l(x_i(t) - x_j(t))^2\right).$$ (12)

This similarity is naturally related to the Euclidean distance used in clustering.

We observe that

$$0 < k(x_i, x_j, w) \leq 1 \quad \forall x_i, \forall x_j, \forall w$$ (13)
Given a starting point \( w \in \mathbb{R}^L \),

repeat

1. \( \Delta w := -\nabla f (w) \).
2. Update. \( w := w + \eta \Delta w \).
3. Normalize: \( w := w/|w| \).

until stopping criterion is satisfied.

![Fig. 3. Gradient descent algorithm](image)

(normal similarity). Furthermore \( K \) is a symmetric matrix and all its diagonal entries are equal to \( 1 \). This implies that \( K \) has all real non-negative eigenvalues. Let \( \lambda_i \) be the \( i \)-th eigenvalue if they are sorted in descending order. Then the trace of the matrix satisfies the relation:

\[
N = \operatorname{tr}(K) = \sum_{i=1}^{N} \lambda_i . \tag{14}
\]

This, along with normality of the similarity measure as per Eq. (13), implies that the largest eigenvalue \( \lambda_1 \) or spectral radius satisfies \( 1 \leq \lambda_1 \leq N \).

As known in the field of spectral clustering [27], the rank of a normal similarity matrix approximates the number of clusters in the data. Specifically, the case \( \lambda_1 = 1 \) corresponds to \( L \) singleton clusters, and the case \( \lambda_1 = N \) corresponds to one cluster including all the samples; the remaining values correspond to intermediate cases.

Therefore, by minimizing the spectral radius over the weight vector \( w \), we attain the solution with maximal diversity, i.e., with the highest separation of clusters. Note that convexity (11) prevents trivial solutions.

To minimize the spectral radius without computing the eigendecomposition, we minimize an upper bound. In general, \( \lambda_1 \) satisfies the following:

\[
\min(s_i) \leq \lambda_1 \leq \max(s_i) , \tag{15}
\]

where \( s_i \) is the sum of the \( i \)-th row of \( K \). Therefore, to minimize the spectral radius we can minimize the objective function \( F(w) = \max_j(s_j) \). To this aim we use an iterative gradient descent algorithm, illustrated in Fig. 3. In this procedure we use the following objective function gradient:

\[
\frac{\partial}{\partial w_r} F(w) = \sum_{i=1}^{N} \frac{\partial}{\partial w_r} k(x_i, x_j, w)
\]

\[
= \sum_{i=1}^{N} -k(x_i, x_j, w)(x_i(r) - x_j(r)), \tag{16}
\]

for \( j = \arg\max(s_i) \).

As for all non-stochastic gradient descent heuristics, for a sufficiently small step size \( \eta \) this procedure will asymptotically converge to a local minimum and may benefit both from multiple restarts, for better quality minima, and from adaptive step size strategies, for faster convergence [28]. However, the problem was experimentally found to be simple, so neither of these improvements was needed.

A total of nineteen volunteer participants (14 males, 5 females, mean age 28 (SD = 6.69)) participated in experimental study performed at the University of Lausanne (Switzerland). All had normal or corrected-to-normal vision, none reported a history of sustained head injury, and all were naïve to the behavioral task. The participants provided written consent for their participation to the experiment approved by the local ethical committee, in line with the Declaration of Helsinki [29]. Continuous EEG was recorded using 60 scalp Ag/AgCl active electrodes (ActiveTwo MARK II Biosemi EEG System, BioSemi B.V., Amsterdam, The Netherlands) mounted on a headcap (NeuroSpec Quick Cap) and referenced to the linked earlobes. Electrophysiological signals were sampled at 2048 Hz with lower cutoff at 0.05 Hz and upper cut-off at 200 Hz, 24 bit resolution (DC amplifiers and software by Biosemi, USA). Electrode impedances were kept below 5 kΩ for all recordings. Vertical and horizontal ocular movements were also recorded using two pairs of bipolar electrodes.

**B. Behavioral task**

The participants performed a behavioral task based on an emotional Go/NoGo paradigm [30]. The experimental procedure, including stimulus presentation and registration of the responses was implemented using the software the E-Prime software (Psychology Software Tools, Inc., Sharpsburg, PA 15215-2821, USA). This software was used to send triggers to the EEG acquisition system via a parallel port connection. The stimuli consisted in pictures showing a face (displayed at the center of the computer screen). The faces were either displaying an emotion (happiness, fear, anger, or sadness) or neutral. In the task studied here, the “Go”-cues were the neutral faces, for which the participants were instructed to respond as fast as possible by pressing a button, while the “NoGo”-cues were the emotional faces. In the trials with “NoGo”-cues, the participants were instructed to withhold response. The stimulus presentation lasted 500 ms per trial and was followed by a blank screen displaying only fixation cross (+) for a duration of 1000 ms, during which the possible button press was recorded. The experimental task was composed by four blocks, and each of them consisted of 30 “Go/NoGo” trials in pseudo-randomized order. In each block, 20 trials presented “Go”-cues and 10 “NoGo”-cues.

**VII. DATA PROCESSING WORKFLOW**

For each participant, the data processing includes the following phases.

**A. Pre-processing**

The EEG data files are imported in EEGLAB [31] and for each participant they are re-referenced with respect to the mastoidal electrodes M1 and M2. We derived two datasets for the subsequent analyses. The first dataset (non-ICA dataset) is formed by raw data files, as explained above. The second dataset (ICA dataset) is formed by raw data preprocessed with...
Independent Component Analysis (ICA) [2] for artifact correction: blink, saccade and eyelid artifact components are set to zero, based on their respective shape and topography [32]. The ICA dataset is also downsampled to 1024 Hz with anti-aliasing filter, prior to the ICA step. Then, for both datasets we apply an IIR Butterworth Band-Pass filter (called “Basic Filter for continuous EEG” in EEGLAB). The Half-Power threshold parameters are \(0.1\) Hz for High-Pass and \(27.6\) Hz for Low-Pass. Then the files are exported as EDF (European Data Format). The remainder of the data processing workflow is formulated for the non-ICA dataset.

B. Segments extraction

The filtered EDF data files are processed with a custom Python script. They are split into vectors corresponding to trials, using the triggers present in the EDF files. We exclude the trials in which the participant gave a wrong response (press the button for a “NoGo” cue, or not press it for a “Go” cue), which leave, on average, 108 (out of 120) trials per participant. We consider a time course of 600 ms for each vector, starting at the beginning of the trial (the instant when the visual stimulus is presented). This short duration is motivated by necessity to avoid muscular artifacts produced by the participant moving to press the button. For each signal vector, we apply a baseline correction by subtracting the average amplitude of the signal in the 200 ms preceding the trial onset. Since the sample rate is \(2048\) Hz, a duration of \(600\) ms corresponds to vectors in \(\mathbb{R}^{1228}\), which are the input to our analysis. The phases of the EEG pre-processing and epoching are displayed in Fig. 4.

C. Smoothing

In order to remove low-amplitude noise and artifact, we regularize the signals with the application of the 1-dimensional Anisotropic Diffusion filter by Perona & Malik [33]. The algorithm is applied to each signal vector, with 1000 iterations. The Anisotropic Diffusion parameter \(\kappa\) is set to \(30\); this parameter controls conduction as a function of gradient, and was chosen heuristically within the recommended range in order to allow smoothing but still preserve deflections and peaks in the signals. The parameter \(\Delta t\), which controls the speed of diffusion, was set to \(0.33\). This is recommended by the authors of the used implementation of the algorithm, in order to preserve numerical stability [34].

D. Artifact rejection

A simple artifact rejection method based on peak-to-peak amplitude is applied. The peak-to-peak amplitude is computed for each signal vector, and then we compute its median value and the median absolute deviation (MAD) across all vectors. A signal vector is rejected if the difference between its peak-to-peak amplitude and the median exceeds twice the MAD. This procedure rejects on average \(14\) signal vectors per participant. The remaining vectors for each participant are considered in the remaining part of the analysis. We denote the set of valid data as \(X \subset \mathbb{R}^{1228}\). Fig. 5 shows an example of the application of artifact rejection.

E. Extraction of the weighted interval-valued features

We compute a weight vector for each participant according to the procedure described in Section V-B. The iterative gradient descent is run for 700 steps with constant step size \(\eta = 10^{-6}\), leading to convergence of the objective function. The resulting mean weight vector is displayed in Fig. 6. As the weights attain their largest values (corresponding to the largest variance among the signals) in the initial and final part
This leads to selecting meaningful information in the evoked potential, we exclude
of the epochs’ duration, which are not believed to convey the average of the weight vectors for all participants in a compact form.

of the epochs’ duration, which are not believed to convey meaningful information in the evoked potential, we exclude the first and last 75ms of each epoch from the remaining part of the analysis.

The participant signal vectors are then pointwise multiplied with the obtained weight vector in order to amplify the differences among them. They are furthermore multiplied by a fixed constant of 1000, in order to avoid numerical errors caused by approximations in the remainder of the data processing.

Then interval-valued features are extracted as described in Section V-A, with a step length \( r \) corresponding to 200ms. This leads to selecting 74 time windows for each signal vector (compared to the original length of the signal vectors that is 1228).

**F. Possibilistic clustering**

The Graded Possibilistic Clustering algorithm with deterministic annealing DAGPC-II (Section III) is applied to the data. We set the parameter \( \alpha \) to 0.85 and the initial number of clusters to \( c_0 = 7 \). This number is obtained by multiplying the expected number of clusters by a factor between 2 and 3. The clustering algorithm outputs, for each cluster \( C_j \), its centroid \( y_j \in \mathbb{R}^{|X|} \) and its membership vector \( u_j \in \mathbb{R}^{|X|} \). Its components express the membership of each sample to the cluster.

**G. Singleton clusters rejection**

Singleton clusters are those modeling only a single sample. We identify them by selecting the clusters whose total membership differs from the average membership across all the clusters by more than 1.5 standard deviations. If there is only one signal vector whose maximum cluster membership is in correspondence of the singleton cluster, then the vector is removed and the clustering algorithm of VII-F is applied again, reducing the number of required clusters by one.

**H. Merging of overlapping clusters**

We apply the fuzzy Jaccard index, defined in Section IV, to determine the closeness of each pair of clusters. For each participant we compute (independently) the distribution of the Jaccard indices for each pair of clusters. This will be in general a multimodal distribution that can be seen as a mixture of normal distributions. The rightmost mode includes pairs of “neighbor” clusters, which are very similar to each other, since their mutual Jaccard index is close to one. The exploration of the dataset made us conclude that a cut-off point \( J_0 = 0.7 \) can be heuristically chosen as a threshold above which clusters have a large overlap and can be merged. This threshold value is applied for each participant: two clusters are merged if their Jaccard index is above \( J_0 \). To merge, we compute a new centroid as the average of the two previous clusters’ centroids weighted by the membership values of their elements, and then new memberships are computed with respect to the new centroid in the same fashion as done in the clustering step of Section VII-F. The procedure continues iteratively recomputing the Jaccard indices at each step, until no more neighboring clusters can be merged.

**VIII. Results**

The analyses and results presented in this study are limited to the signal recorded at the electrode site ‘Fz’. While the full dataset is not publicly accessible, since this was not part of the study protocol, we have included a summary of the data as Supplemental Material. For each participant, the described analysis workflow gave a number of clusters between 2 and 5. An example is displayed in Fig. 7.

The clusters could not definitely be associated to different experimental conditions (Go/NoGo condition, type of emotion). This suggests that high cognitive functions such as affective discrimination are not associated with simple wave
components of the brain signal and might instead require the choice of higher level features or specific indices, or the study of a larger number of EEG locations.

A. Clusterization rate

In order to assess the validity and interest of the obtained clusters, we measure the correspondence between the result of the analysis and the “Go” and “NoGo” experimental conditions. This is done by defining, for each participant, a measure called clusterization rate. The clusterization rate of the “Go” trials (respectively, the “NoGo”) trials is defined as the fraction of “Go” (respectively, “NoGo”) trials characterized by a high membership to one of the identified clusters. This is defined as the number of samples with a membership above the 95-th percentile of the cluster membership distribution for one of the clusters, divided by the total number of “Go” trials for the participant (respectively, “NoGo”). For most participants, the clusterization rate was higher in “Go” trials than in “NoGo” trials (Fig. 8).

Since the results of the data processing workflow are dependent on the random seed initialization for the clustering step, we have repeated the same analysis for 20 different values of the random seed, in order to evaluate the robustness of the result.

B. Weight optimizations and interval features

In order to assess the contribution to the result given by the weight optimization procedure (Section V-B) and the interval-valued features selection (Section V-A), we have performed the same analysis switching off one or both optimizations. The results for DAGPC-II are summarized in Table I. The mean differences in each row of the table are significant (Welch Two Sample t-test, all p-values < 10^{-3}). We can observe that, even though all four versions of the analysis gave a significant difference in clusterization rate between “Go” and “NoGo” trials, the use of both the weight optimization procedure and the interval feature selection gave the best result in terms of relative difference in clusterization rate between the two classes (69%). In this version of the analysis, we observed that the average clusterization rates for the “Go” trials were larger than the corresponding clusterization rates for the “NoGo” trials for 16 participants (Fig. 8a).

It can also be noticed that the internal variability of the clusterization rate values for the “Go” trials was smaller than for the “NoGo” trials (Fig. 8b). This is reasonable, since the “Go” trials all had neutral facial expression as a cue. This can explain more similar ERPs, which in turn would be associated with a high membership to one of the identified clusters. Conversely, the “NoGo” trials corresponded to stimuli presenting different facial expressions and therefore tended to be more different. This is also compatible with the fact that, if we inspect the data, we generally observe larger inner variability among the “NoGo” trials compared to the Go trials (Fig. 9).

C. Comparison with different algorithms

For comparison purposes, we have run numerical experiment replacing our proposed clustering algorithm DAGPC-II (Section III) with the Fuzzy c-means (FCM) algorithm [35].

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**Fig. 8.** Comparison of the clusterization rates of “Go” and “NoGo” trials for all participants (after the application of weight optimization procedure and the calculation of the interval-values features). (a) Scatter plot of the average values over 20 different random seed initializations (each point corresponds to one participant). For 10 participants (out of 19), the clusterization rate of the “Go” trials is higher than the one of the “NoGo” trials. This reflects the fact that the signal vectors associated to “Go” trials tend to be more similar to each other, while the “NoGo” trials are characterized by higher variability which escapes the clusters. (b) Average value and standard error of the clusterization rate of “Go” and “NoGo” trials for each participant over the same 20 different random seed initializations.
TABLE I
DAGPC-II algorithm. Comparison of clusterization rates. Each row presents the average clusterization rates for the “Go” and “NoGo” trials in the specified condition for the Non-ICA and the ICA datasets

<table>
<thead>
<tr>
<th>Method</th>
<th>Clustering rate</th>
<th># participants with</th>
<th>Non-ICA dataset</th>
<th>ICA dataset</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weights</td>
<td></td>
<td></td>
<td>Go trials</td>
<td>NoGo trials</td>
</tr>
<tr>
<td>Yes Yes</td>
<td>0.198</td>
<td>0.117</td>
<td>16 out of 19</td>
<td>0.229</td>
</tr>
<tr>
<td>Yes No</td>
<td>0.183</td>
<td>0.132</td>
<td>13 out of 19</td>
<td>0.213</td>
</tr>
<tr>
<td>No Yes</td>
<td>0.189</td>
<td>0.119</td>
<td>16 out of 19</td>
<td>0.169</td>
</tr>
<tr>
<td>No No</td>
<td>0.189</td>
<td>0.144</td>
<td>13 out of 19</td>
<td>0.173</td>
</tr>
</tbody>
</table>

TABLE II
Fuzzy c-means algorithm. Comparison of clusterization rates. Each row presents the average clusterization rates for the “Go” and “NoGo” trials in the specified condition for the Non-ICA and the ICA datasets

<table>
<thead>
<tr>
<th>Method</th>
<th>Clustering rate</th>
<th># participants with</th>
<th>Non-ICA dataset</th>
<th>ICA dataset</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weights</td>
<td></td>
<td></td>
<td>Go trials</td>
<td>NoGo trials</td>
</tr>
<tr>
<td>Yes Yes</td>
<td>0.286</td>
<td>0.154</td>
<td>15 out of 19</td>
<td>0.319</td>
</tr>
<tr>
<td>Yes No</td>
<td>0.212</td>
<td>0.135</td>
<td>13 out of 19</td>
<td>0.309</td>
</tr>
<tr>
<td>No Yes</td>
<td>0.269</td>
<td>0.163</td>
<td>15 out of 19</td>
<td>0.313</td>
</tr>
<tr>
<td>No No</td>
<td>0.211</td>
<td>0.156</td>
<td>15 out of 19</td>
<td>0.294</td>
</tr>
</tbody>
</table>

Fig. 9. Mean ERP signals for participant #12721 grouped in Go and NoGo trials. The shaded areas represent the instantaneous standard deviation. It can be observed that the variability is larger among the NoGo trial, compared to the Go trials. This is compatible with a higher clusterization rate for the Go trials.

As before, the analysis was run 20 times with different initialization of the random seed. The resulting clusterization rates are reported in Table II. The mean differences in each row of the table are significant (Welch Two Sample t-test, all p-values < 10^{-3}), except when marked with ns. When running the analysis with FCM with weight optimization and interval features, the average clusterization rates for the “Go” trials were larger than the corresponding clusterization rates for the “NoGo” trials for 15 participants. This is a comparatively worse result with respect to that obtained with DAGPC-II, where the classification was correct for 16 participants.

D. Effect of Independent Component Analysis preprocessing
In order to assess the robustness of our proposed method to EEG artifacts, we have run the data processing workflow both on the ICA-preprocessed dataset and the non ICA-preprocessed. The results are reported in Tables I and II for the two clustering algorithms. We notice how the ICA-preprocessing does not improve the difference of clusterization rate between the “Go” and “NoGo” trials.

IX. DISCUSSION AND CONCLUSIONS
We presented a workflow for the unsupervised analysis of EEG signals using fuzzy clustering. It should be noted that the choices that we have made in the design are not bound to the specific dataset that was used as an example, and therefore make the data processing workflow, or parts thereof, applicable to any ERP dataset, requiring a minimal number of parameters to be decided. In comparison to Fuzzy c-means, our proposed algorithm DAGPC-II produced better results in most conditions, as a part of the data processing workflow in our experiments. However, the margin is thin, which is expected since the Graded Possibilistic algorithm is related to FCM, as pointed out in Section III. The advantage of DAGPC-II lies in fact in the grading, which makes it possible to change its level of “possibility”, by varying the parameter α, and thereby changing the algorithm’s reliance on local information.

ICA is very often applied as a preprocessing step for EEG data and requires an expert to assess which components are artifact-related and should therefore be removed. Since our overall goal is to define a data processing workflow that minimizes human intervention, it is relevant to remark how the application of ICA does not improve our result in terms of difference in clusterization rate between the “Go” and “NoGo” trials. This suggests that our method is robust to EEG artifact and can indeed be applied even if the signals have not been preprocessed to manually remove noise components.

The only choice in the parameters that we believe to be data-dependent is the threshold value J₀ that we used when merging similar clusters based on their fuzzy Jaccard index. We set this value J₀ = 0.7 based on features specific to our
dataset derived from the distribution of its values across all samples.

Our method could be extended to the analysis of continuous EEG signals: since the analysis workflow requires signal vectors of the same length as input, the continuous EEG would have to be pre-processed with a windowing method, in order to obtain a number of signal vectors on which to apply the clustering algorithm.

Our previous work, in which the clustering algorithm compared the signals directly by their Euclidean distances [6], has now been surpassed by the use of optimized weights and interval features, which together contributed to a better result in the index we used to assess the results of the analysis. In fact, the direct comparison of signals as vectors ignores the temporal nature of the data and can lead to the loss of higher order features and inaccurate comparisons due to time shifts and delayed onset of EEG peaks. The optimized weights contribute to enhancing the differences along the signal vectors, thereby allowing the clustering algorithm to capture this variability. Using time window interval features makes the calculation less dependent on onset delays in the evoked responses.

Our work proposes an unsupervised approach to ERP analysis. This is not to mean that unsupervised methods are intrinsically better than supervised or semi-supervised ones, but their goals are different. In fact, we see our method as a preliminary and complementary to other (semi-)supervised approaches. We set out to define a data processing workflow that can be applied in generality to the exploratory analysis of ERP datasets in order to explore their internal structure captured by clustering. We imagine that an EEG researcher will be able to use our method as a first step in the study of an ERP dataset without imposing any constraints given by pre-determined labels. Subsequently they will use the results of the clustering to choose interesting classes of samples to classify with a (semi-)supervised approach.

Brain processes at all time scales are characterized by great complexity, and require analysis techniques able to capture it [36]–[39], such as methods aimed to identify specific features of the ERPs. In future work, we plan to incorporate indices targeted to the comparison of peaks and wave components, and able to measure time delays such as those used in [40]–[43].

In addition to suitable indices, data complexity and variety may also call for more powerful analysis tools. In this respect, kernel methods [44] may provide the necessary power and flexibility. Kernel methods for one-class classification have been developed by the authors [45] and may be employed in future research to deepen our understanding of the problem. Other approaches, like Infinite Kernel Learning [46], may offer better accuracy than single-kernel learning for heterogeneous data.

The possibility of capturing higher level features related to the brain processes guiding the task will make it possible to compare the EEG recordings of different participants in an unsupervised way, grouping the participants according to their mutual similarities. It will also be important to apply normalization rules to account for inter-participant amplitude variability. As EEG hardware becomes more accessible, both in terms of cost and availability, its use becomes more common in many fields. Beyond neuroscience research, one central example is healthcare, where EEG offers an effective way to examine a patient’s brain function and can be the basis for the definition of bio-markers with a high diagnostic relevance. Other future applications lie even in field of end-user products (e.g. personal monitoring of brain function for health and well-being, Brain-Computer Interfaces). EEG analyses based on computational intelligence methods are a valuable tool to address the huge quantities of data generated by EEG devices in any field and extract meaningful patterns from them. We believe that our work contributes to tackling this challenge.

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