On the use of evolutionary time series analysis for segmenting paleoclimate data

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Abstract

Recent studies propose that different dynamical systems, such as climate, ecological and financial systems, among others, present critical transition points named as tipping points (TPs). Climate TPs can severely affect millions of lives on Earth so that an active scientific community is working on finding early warning signals. This paper deals with the development of a time series segmentation algorithm for paleoclimate data in order to find segments sharing common statistical patterns. The proposed algorithm uses a clustering-based approach for evaluating the solutions and six statistical features, most of which have been previously considered in the detection of early warning signals in paleoclimate TPs. Due to the limitations of classical statistical methods, we propose the use of a genetic algorithm to automatically segment the series, together with a method to compare the segmentations. The final segments provided by the algorithm are used to construct a prediction model, whose promising results show the importance of segmentation for improving the understanding of a time series.

Key words: Time series segmentation, genetic algorithms, clustering, paleoclimate data, tipping points, abrupt climate change

1. Introduction

In contrast to the famous statement of Linnaeus (1751) “natura non facit saltus” (or nature makes no leaps), it has been proven that some points of no return, thresholds and phase changes are widespread in nature and these are often non linear [1]. Such events can be rarely anticipated and some of them can have detrimental consequences on Earth’s climate and large-scale impacts on human and ecological systems. This increases the imperious necessity of studying, analysing and developing techniques for characterising them in order to construct reliable early warning systems. Although the human being have influenced their local environment for millennia, e.g. reducing biodiversity, it is now, since the industrial revolution, that truly global changes are being noticed [2, 3]. Examples that are currently receiving attention include the potential collapse of the Atlantic thermohaline circulation, the dieback of the Amazon rainforest or the decay of the Greenland ice sheet [1]. Formally, a climate “tipping point” (TP, also known as “little things can make a big difference”) occurs when a small change in forcing triggers a strongly nonlinear response in the internal dynamics of part of the climate system, qualitatively changing its future state.

The critical relevance of early TPs detection has produced a growing attention of the scientific community. Lenton differentiates between several types of TPs, and presents some indicators that can help to detect them, such as the increase of auto-correlation of the series values [4]. In [5], more concrete techniques regarding data processing and indicators are presented. They study a bank of methods using only simulated ecological data, concluding in concordance with the literature that there is no unique best indicator for identifying an upcoming transition. They also conclude that all the methods require specific data-treatment. Up to our knowledge, all previous works tackle the TP detection with statistical methods trying to select (by trial and error) the method (and the time-window) most suitable to detect those transitions. They require an intensive data preprocessing that includes, for instance, the use of Gaussian filters or rolling windows that introduce extra parameters (such as the width of the Gaussian function or size of the window) that need to be optimised [4, 5]. The main limitation behind these methods is that different TPs require specific treatment, which is specifically the objective that this paper tries to tackle.

Time series segmentation is a research field, aiming to provide a compact representation of the time series values, dividing it into segments and using an abstract representation of each segment. It is very important for time series representation and time series mining [6, 7, 8] and is commonly used as a pre-processing step for different mining tasks [8, 9, 10, 11] (e.g. clustering, classification or motif detection) and for data compressing [12, 13]. In this way, segmentation algorithms have been used in many different fields, such as paleoecological problems [14], phoneme recognition [15] or paleontological climate [16]. Some recent works have proposed the use of algebraic segmentation for the specific case of short-term [17, 18]
and short nonstationary time series [19].

In the context of time series segmentation, this paper deals with climate time series segmentation to effectively analyse Dansgaard–Oeschger (DO) events (i.e. the consequence of different tipping points occurred in the past). The main objective of the paper is to improve the ability to understand paleo-records and identify abrupt climate changes (in this case DO events). TPs have been studied using predefined and fixed time-windows, or specifying different windows for each one (as it has been suggested that there are different types of TPs, represented by different time spans and statistical features). In other words, it is well-known that there exist a nonlinear trigger (known as a TP) which anticipates abrupt climate changes, but the way to characterise and detect these triggers is uncertain to this point, mainly because the nature of each TP is usually different from the rest (they occur in different time spans and there exist different statistical features that help to detect them). Motivated by this hypothesis, the main aim for using a segmentation algorithm is to automatically detect and exploit these differences between the TPs. The combined use of an evolutionary segmentation algorithm, a clustering method and a set of features previously linked to TP detection would ideally serve two different objectives: 1) Group similar patterns according to the set of defined features (which should help to detect TPs) and 2) find the best time span to detect each TP (this search is guided by a measure of the compactness of the clustering). Note that given the nature of both the evolutionary algorithm and fitness function these two objectives are achieved jointly, optimising the segmentation so that the differentiation between the clusters is maximal (according to a set of predefined features which have been linked to TP detection, which would guide the search towards clusters composed of TPs against clusters of characteristics radically different to the ones found in TPs).

A preliminary study in this direction was presented in [16], where an evolutionary segmentation algorithm was applied to analyse the segments preceding TPs, using the quotient of the number of segments divided by the sum of squared error of the clustering as the fitness function. This paper extends the previous work with the following contributions:

- The evaluation of the segmentations obtained by the different versions of the algorithm has been automated using two different metrics, the former comparing the segmentation to an ideal segmentation and the latter evaluating the stability of the algorithm. These evaluation metrics are used as a post-evolution analysis of the TP detection capability of each version of the algorithm.

- Moreover, this automatic evaluation method has been used to perform a battery of experiments comparing the results obtained by a total of 10 different cluster validity indexes used as fitness functions. The new fitness functions improve the results of [16] to a great extent. These metrics, which measure cluster compression, are used to guide the algorithm to different solutions, which are later compared.

- The best segmentation obtained has been used to devise a classification model able to recognise abrupt climate changes based on the characteristics of the previous segment. This model can be used to better understand the characteristics of potential early warning signals of TPs.

- Some improvements have been included in the algorithm, such as the binary coding (which alleviates the computational load of the algorithm) and a restriction of the minimum segment size.

The rest of the paper is organised as follows. Section 2 presents the segmentation algorithm, while Section 3 presents a proposal for segmentation comparison. Section 4 discusses the experimental results and the last section depicts the conclusions and future research.

2. Segmentation Algorithm

Given a univariate time series \( Y = \{y_n\}_{n=1}^N \), our objective is to divide the values of \( y_n \) into \( m \) consecutive subsets
or segments. These segments should present a homogeneous behaviour regarding the values of \( y_n \). This is done by partitioning the time indexes \( (n = 1, \ldots, N) \) into segments: \( S_1 = \{y_1, \ldots, y_{s_1}\}, S_2 = \{y_{s_1}, \ldots, y_{s_2}\}, \ldots, S_m = \{y_{s_m}, \ldots, y_{s_m}\} \), where \( i \)'s are the cut points and are subscripted in ascending order \((t_1 < t_2 < t_m-1)\). The cut points belong to two segments (the one before and the one after to analyse consistently the transitions of the time series). The cut points \( t_i, i = 1, \ldots, m-1 \), are the parameters to be determined by the algorithm. As done in [23], we extend this setting by trying to group the segments into \( k \) different classes or clusters \((k < m)\), where \( k \) is a parameter defined by the user. In this way, each segment, \( s_j \), will be associated to a class label: \( \{s_1, z_1\}, \ldots, \{s_m, z_m\} \), where \( z_i, i = 1, \ldots, m \), is the class label of the \( i \)-th segment and takes values in a set of \( k \) different labels, \( z_i \in [C_1, \ldots, C_k] \).

2.1. Summary of the algorithm

The Genetic Algorithm considered in this paper can be included in the area of time series segmentation [7, 24, 25, 26, 9, 10, 11, 17, 13]. Our main objective is to devise an unsupervised methodology to identify time series with similar statistical behaviour [23]. This is a necessary step to study the characteristics of these time series and be able to analyse the temporal transitions between different states (i.e., types of segments) or to construct a prediction model with a dynamic-window [16].

In the algorithm, each possible segmentation is represented as an array of binary values (chromosome representation), where a value of 1 represents a cut point. The evolution starts from a population of randomly generated segmentations. Mutation and crossover operators are applied to explore and exploit the search space. This procedure is repeated during \( g \) generations. To evaluate a solution (or segmentation), we select a set of statistics to be calculated for each segment. Then, similar segments are grouped using a clustering process. A pseudocode of the segmentation algorithm is included in Figure 1. A detailed description of each step of the GA are defined in the following subsections.

### Evolutionary time series segmentation algorithm

- **Require:** Time series.
- **Ensure:** Best segmentation of the time series.
- 1. Generate a random population of \( P \) time series segmentation solutions.
- 2. Evaluate all segmentations of the initial population by using a set of statistics, a clustering algorithm and a predefined fitness function \( f \).
- 3. while not Stop Condition do
- 4. Store a copy of the best solution.
- 5. Select parents from current population.
- 6. Generate offspring - apply crossover and mutation to construct new candidate segmentations.
- 7. Evaluate the fitness of the offspring.
- 8. Merge parents and offspring.
- 9. Replace current population with selected segmentations of the previous union.
- 10. end while
- 11. Compare with the ideal segmentation.
- 12. return Resulting segmentation.

Figure 1: Main steps for the GA. Note that in this case the stop condition is controlled by a parameter \( g \) associated to the number of generations of the evolutionary algorithm.

### 2.2. Chromosome representation

As stated before, each individual chromosome consists of an array of binary values, where the length of the chromosome is the time series length, \( N \). Each position \( c_i \) stores whether the time index \( t_i \) of the time series represents a cut point for the evaluated solution. In this sense, for a given segment \( s_j \), delimited by the cut points \( t_{i-1} \) and \( t_i \) \((t_{i-1} < t_i)\), the corresponding chromosome values will be \( c_{i-1} = 1, c_i = 1 \) and \( c_l = 0, \forall l \neq i, l < t_i \).

### 2.3. Initial population

In order to initialise the population of \( P \) segmentations solutions, an average segment length has to be specified by the user \((\bar{m})\). Taking into account that the cut points belong to two segments, the number of cut points will be \( \bar{m} = \lceil \frac{N}{g} \rceil \), so the chromosomes are binary arrays where \( \bar{m} \) random positions are 1s and the rest are 0s. In order to check that time segments are not overly small we include the constraint that no consecutive cut points are allowed (setting then a minimum length of 3 points per segment).

### 2.4. Fitness evaluation

As stated before, the main objective of the algorithm is not only to segment the time series but also to find time patterns with similar behaviour (whether they represent TP or not). Because of this, a clustering process is applied to group similar segments. The evaluation of the quality of a segmentation solution consists of three different steps: extracting the characteristics of the segments, applying a clustering process and measuring the quality of this clustering.

#### 2.4.1. Extracting segment characteristics

Given that the segments in a chromosome can have different length, an approach is designed to project all the segments into the same dimensional space. Six statistical metrics are considered and measured for all chromosome segments (note that each segment is an univariate time series). Then, the similarities between segments can be calculated in the 6-dimensional space. Consider \( s_j \) as a segment fulfilling the previously stated conditions (i.e., \( s_j \) is a segment delimited by the cut points \( t_{i-1} \) and \( t_i \), where the segment length is \( t_i - t_{i-1} + 1 \)). The mapping is done by the function \( f : \mathbb{R}^{(t_i-t_{i-1}+1)} \rightarrow \mathbb{R}^6 \), in the following way:

\[
f(s_j) = (S_j^2, y_1, y_2, a_1, MSE, AC_j)
\]

where the different characteristics are defined as:

1. Variance \((S_j^2)\): It measures the variability of the segment:

\[
S_j^2 = \frac{1}{t_i - t_{i-1} + 1} \sum_{i=t_{i-1}}^{t_i} (y_i - \overline{y}_s)^2,
\]

where \( y_i \) are the time series values of the segment, and \( \overline{y}_s \) is the average value of the segment.

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1. Note that the first and last points of the chromosome are considered as cut points.
2. Skewness ($\gamma_1$): It represents the (vertical) asymmetry of the distribution of the series values in the segment with respect to the arithmetic mean:

$$\gamma_1 = \frac{1}{t_i-t_{i-1}} \sum_{i=1}^{t_i} \frac{(y_i - \bar{y})^3}{S_s^3},$$

where $S_s$ is the standard deviation of the $s$-th segment.

3. Kurtosis ($\gamma_2$): This statistic is related to the degree of concentration that the values present around the mean of the distribution:

$$\gamma_2 = \frac{1}{t_i-t_{i-1}} \sum_{i=1}^{t_i} \frac{(y_i - \bar{y})^4}{S_s^4} - 3. \quad (4)$$

4. Slope of a linear regression of the points of the segment ($a_s$): A linear model is constructed for every segment trying to achieve the best linear approximation of the points of the time series in the evaluated segment. The slope is a measure of the general tendency of the segment:

$$a_s = \frac{S_s^{xy}}{(S_s^y)^2}, \quad (5)$$

where, for the $s$-th segment, $S_s^{xy}$ is the covariance between the time indexes, $t_i$ and the time series values, $y_i$; and $S_s^y$ is the standard deviation of the time values. Covariance $S_s^{xy}$ is defined by:

$$S_s^{xy} = \frac{1}{t_i-t_{i-1}} \sum_{i=1}^{t_i} (i - T_s) \cdot (y_i - \bar{y}_s). \quad (6)$$

5. Mean Squared Error ($MSE_s$): Considering the same linear model than the one used for the slope, we measure the error ($MSE_s$) of this linear fitting:

$$MSE_s = S_s^2 \cdot (1 - r_s^2), \quad (7)$$

where $r_s^2 = \frac{S_s^{xy}}{S_s^x \cdot S_s^y}$.

6. Autocorrelation coefficient ($AC_s$): This a measure of the correlation between the current values of the time series and the previous ones:

$$AC_s = \frac{\sum_{i=1}^{t_i} (y_i - \bar{y}_s) \cdot (y_{i+1} - \bar{y}_s)}{S_s^2}.$$

2.4.2. Clustering process: k-means

A clustering process is applied to group similar segments (taking into account the six selected statistical measures). For simplicity, the algorithm chosen for the clustering step is the well-known k-means. Before the clustering algorithm, note that a normalisation of the values of the segment metrics is conducted, as the distance from each segment to its centroid strongly depends on the range of values of each metric.

In the classic k-means, the initial centroids are randomly chosen from the set of patterns. Instead, we have developed a deterministic process to select these centroids which ensures that a chromosome will always have the same fitness. First, we choose the feature with the maximum standard deviation. The first initial centroid will be the segment with the highest value for this feature. The second one will be the segment with the highest Euclidean distance from the first centroid. The third centroid will be that which is farthest from both, and so on. This assures a deterministic initialisation, at the same time that the initial centroids are as far as possible from each other, favouring centroid diversity.

2.4.3. Measuring the quality of the clustering process

The last step of the evaluation of the chromosome is to measure how well the segments are grouped (compactness of the clustering). It is clear that different clustering algorithms usually lead to different clusters or reveal different clustering structures. In this sense, the problem of objectively and quantitatively evaluating the clustering results is particularly important, and this is known in the literature as cluster validation. There are two different testing criteria for this purpose [27, 28]: external criteria and internal criteria. When a clustering is evaluated based on the clustered data it is called internal evaluation. In external evaluation, the clustering is evaluated using, for example, known class labels. Based on these concepts, the internal criteria evaluation metrics will be a suitable option for the evolution, because the GA is not given a priori information of the segments to be found. Note that the segments metrics are normalised at this step as well. We have considered ten different metrics:

- **Sum of squared errors (SSE):** The simplest error measure is the sum of squared errors (considering errors as the distance from each point to their centroid), i.e.:  
  \[
  SSE = \frac{1}{N} \sum_{i=1}^{k} \sum_{c \in C_i} d(x, c)^2, \quad (9)
  \]

  where $k$ is the number of clusters, $c_i$ is the centroid of cluster $C_i$, and $d(x, c_i)$ is the Euclidean distance between pattern $x$ and centroid $c_i$. This function does not prevent clusters to fall very close in the clustering space. As this index has to be minimised, the fitness will be defined as $f = \frac{1}{1+SSE}$.

- **Normalised sum of squared errors (NSSE):** One of the disadvantages with the SSE measure is that it is sensitive to the number of segments in the solution (this index being generally lower as the number of segments decreases). Because of this, a more consistent approach would be to divide the error by the number of segments:
  \[
  NSSE = \frac{1}{k} \sum_{i=1}^{k} \sum_{c \in C_i} d(x, c)^2 \quad (10)
  \]

  The fitness in this case will be also defined as $f = \frac{1}{1+NSSE}$.

- **Calinski and Harabasz index (CH):** This index has been found to be one of the best performing ones for adjusting
the value of \( k \). It is defined as:

\[
CH = \frac{\text{Tr}(S_B) \cdot (N - k)}{\text{Tr}(S_W) \cdot (k - 1)},
\]

(11)

where \( N \) is the number of patterns, and \( \text{Tr}(S_B) \) and \( \text{Tr}(S_W) \) are the trace of the between and within-class scatter matrices, respectively. Note that the value of \( k \) will be fixed in our algorithm. As this index has to be maximised, the fitness will be defined as \( f = CH \).

- **Davies-Bouldin index (DB):** This index also attempts to maximize the between-cluster distance while minimizing the distance from the cluster centroids to the rest of points. It is calculated as follows:

\[
DB = \frac{1}{k} \sum_{i=1}^{k} \max_{i \neq j} \frac{d_i + d_j}{d(c_i, c_j)}.
\]

(12)

where \( d_i \) is the average distance of all elements in cluster \( C_i \) to centroid \( c_i \), and \( d(c_i, c_j) \) is the distance between centroids \( c_i \) and \( c_j \). As this index has to be minimised, the fitness will be defined as \( f = \frac{1}{1 + DB} \).

- **Dunn index (DU) and variants:** The Dunn index attempts to identify clusters that are compact and well-separated. In this case, the distance between two clusters is defined as \( d(C_i, C_j) = \min_{x \in C_i, y \in C_j} d(x, y) \), that is, the minimum distance between a pair of points \( x \) and \( y \) belonging to \( C_i \) and \( C_j \). Furthermore, we define the diameter \( \text{diam}(C_i) \) of cluster \( C_i \) as the maximum distance between two of its members, such as: \( \text{diam}(C_i) = \max_{x, y \in C_i} d(x, y) \). Then, the Dunn index is constructed as:

\[
DU = \min_{i=1, \ldots, k} \left( \min_{j \neq i} \left( \frac{\delta(C_i, C_j)}{\text{diam}(C_i)} \right) \right).
\]

(13)

The Dunn index has been found to be sensitive to noise, but this disadvantage can be avoided by considering different definitions of cluster distance or cluster diameter. In this paper, we consider four different definitions of the Dunn index (corresponding to different definitions of the distance between clusters and the cluster diameter). For example, as suggested in [27], the cluster diameter can be computed as:

\[
\text{diam}(C_i) = \frac{1}{N_{C_i}(N_{C_i} - 1)} \sum_{x, y \in C_i} d(x, y),
\]

(14)

where \( N_{C_i} \) is the number of patterns belonging to cluster \( C_i \). This cluster diameter estimation has been found to be more robust in the presence of noise and as such, it has been included in the experiments (DU fitness function). Moreover, as said, other variants exist for this index. We selected the three variants that reported the best results in [28], referred to as GD33, GD43 and GD53, which correspond to the following variations of \( \delta(C_i, C_j) \):

\[
\delta^3(C_i, C_j) = \frac{1}{N_{C_i}N_{C_j}} \sum_{x \in C_i, y \in C_j} d(x, y),
\]

(15)

\[
\delta^5(C_i, C_j) = d(C_i, C_j),
\]

(16)

\[
\delta^7(C_i, C_j) = \frac{1}{N_{C_i} + N_{C_j}} \left( \sum_{x \in C_i} d(x, C_j) + \sum_{y \in C_j} d(y, C_i) \right),
\]

(17)

where \( C_i \) represents the mean of the cluster. The term \( \text{diam}(C_i) \) is defined in this case as:

\[
\text{diam}(C_i) = \frac{2}{N_{C_i}} \sum_{x \in C_i} d^p(x, C_i),
\]

(18)

where \( d^p(x, C_i) \) is the Point Symmetry-Distance between object \( x \) and the cluster \( C_i \), defined in [29].

Since all the variations of this index have to be maximised, the fitness will be defined as \( f = DU \).

- **Silhouette index (SI):** For this index, the data cohesion is computed based on the distance between all the points in the same cluster, and the separation is measured using the nearest neighbour distance. It is defined as:

\[
SI = \frac{1}{N} \sum_{i=1}^{k} \sum_{x \in C_i} \left( b(x, C_i) - a(x, C_i) \right),
\]

(19)

where \( a \) and \( b \) correspond respectively to the intra-cluster and inter-cluster distances and are defined as follows:

\[
a(x, C_i) = \frac{1}{N_{C_i}} \sum_{y \in C_i} d(x, y),
\]

(20)

\[
b(x, C_i) = \min_{C_j \neq i} \left( \frac{1}{N_{C_j}} \sum_{y \in C_j} d(x, y) \right).
\]

(21)

- **COP index (COP):** The cluster cohesion is estimated, for this index, using the distance from the points to their cluster centroid and the furthest neighbour distance. This index is defined as:

\[
COP = \frac{1}{N} \sum_{i=1}^{k} \frac{\sum_{x \in C_i} d(x, C_i)}{\min_{y \in C_i} \max_{x \in C_i} d(x, y)}.
\]

(22)

### 2.5. Selection and replacement processes

All individuals will be considered for reproduction and generation of offspring, promoting a greater diversity because all individuals are potential parents. After the application of the genetic operators, the offspring and the parent population are joined and a replacement process is performed by roulette wheel selection. The selection probability for each individual chromosome is calculated from its fitness value. The roulette wheel process is repeated as many times as the population size minus one, and the last place is kept for the best segmentation of the previous generation, thus being an elitist algorithm. As can be seen, the selection process promotes diversity, while the replacement process promotes elitism.
2.6. Crossover Operator

The algorithm includes a crossover operator, whose main function is to perform an exploitation of the existing solutions. For each parent individual, the crossover operator is applied with a given probability $p_c$. The operator randomly selects a different parent and a time index. Then, the left and right parts of the selected chromosomes are interchanged with respect to the time index. After this step, the algorithm checks if the solutions verify the minimum length per segment constraint. If not, the crossover operator is repeated using the same parents, until a maximum of three attempts is reached. If no feasible solution is produced, a different parent individual is chosen. This process is repeated until the crossover ends successfully.

![Figure 2: Representation of the crossover operator: A time index is chosen as cut point for the two parents, resulting in the offspring.](image)

2.7. Mutation Operator

Two mutation operators are included in the GA with the aim of reducing the dependency on the initial population and for escaping from local optima. The probability $p_m$ of performing any mutation is decided by the user. When a mutation is to be performed, the type of perturbation is randomly selected from the following two: 1) add or remove (with the same probability) a given number of cut points; and 2) move a given number of cut points towards the left or the right (with the same probability).

For all the mutations, the number of cut points to be mutated is decided by the user parameter ($m_p$) as a percentage of the current number of cut points. When moving cut points to the right or to the left, each selected cut point is randomly pushed towards the previous or the following cut point (with the constraint that it has to maintain the minimum segment length). If the mutation does not maintain this constraint, the same procedure used in the crossover is considered.

2.8. Parameters of the algorithm

For the sake of clarity this section includes a summary of all the parameters involved in the proposed segmentation algorithm. Concerning the clustering step, the two parameters are $k$ (the number of clusters to be found) and $i_t$ (number of iterations for $k$-means). With respect to the evolutionary aspect of the algorithm we have a wider range of parameters: $g$ (number of generations), $P$ (population size), $p_m$ (mutation probability), $p_c$ (crossover probability), $s$ (seed), $m_p$ (percentage of cut points to be mutated), $s_l$ (initial mean segment length) and $f$ (fitness function).

3. Automatic evaluation of the segmentation

In order to evaluate the results of the algorithm, two evaluation metrics are proposed. These measures analyse both the homogeneity of cluster assignment with respect to the DO events and the robustness of the results obtained from different seeds. They are not included in the fitness function, serving only as an automatic way of evaluating the quality of the segmentation, avoiding the intervention of the expert. Both indexes compare two different clustering partitions:

1. **Rand index (RI):** This metric is particularly useful for data clustering evaluation [30]. It is related to the accuracy or correct classification rate (since it measures the correlation between the obtained clusters and the ideal ones), but is applicable even when class labels are not available, as in our case. A set $Y = \{y_n\}_{n=1}^N$ is given (in our case, the time series), and two clustering partitions of $Y$ are to be compared: $X = \{X_1, \ldots, X_l\}$ and $Z = \{Z_1, \ldots, Z_l\}$. For a given segmentation, the partitions are defined in the following way: $X_l$ is a set containing every $y_i$ that is either in $X_l$ or in $C_l$, i.e., the partitions are based on the label assigned to each time series value $y_i$ from the current segmentation. The following two numbers are defined: $a$ (number of pairs in $Y$ that are in the same set in $X$ and $Z$) and $b$ (number of pairs in $Y$ that are in different sets in $X$ and $Z$). Then, the Rand index is defined as: $RI = (a + b)/(\binom{N}{2})$. This metric has a value between 0 and 1, with 0 indicating that the two partitions do not agree on any pair of points and 1 indicating that they are exactly the same.

2. **Adjusted rand index (ARI):** It is a corrected version of the RI [31] trying to fix some of its known problems, e.g., the expected value of the RI for two random partitions does not take a constant value and it approaches its upper limit of unity as the number of clusters increases. ARI values range from $-1$ to +1, yielding negative values if the index is less than the expected one. The detailed formulation can be found in [31].

In order to evaluate the segmentation returned by the algorithm, we compare it with an ideal segmentation\(^2\). These ideal segmentations have been designed by examining the literature about Dansgaard-Oeschger (DO) events, which are associated to TPs. The ideal segmentation for NGRIP has been included in Figure 3. In the Figures, the onsets of the DO events (in a first approximation, we do not consider the error margin) reported in [22] are represented by vertical lines and the segments covering the precursor period to the DO events are delimited by the slope closest to the corresponding onset. The closer the segmentation returned by the GA is to this ideal segmentation, the better the segmentation. To perform this comparison, RI and ARI indexes will be used (ARI_Ideal and RI_Ideal).

\(^2\)It is a hypothetically ideal segmentation, based on the available data. The hypothesis is that the onset of the DO events is detected from combined analysis of benthic sediment and ice core data [32]. Those data do not always agree, which could result in part of the error margin. The timing method contributes the rest of the error.
Given that the wishful ideal segmentation would be binary (non DO event or DO event) and the segmentation returned by the GA can have a value of $k > 2$, we need to binarise the segmentation of the GA (i.e. decide which clusters represent the DO events and which not). Preliminary experiments [16] revealed that DO events were usually identified by two different sequences of clusters, so we evaluated $\text{ARI}_{\text{Ideal}}$ and $\text{RI}_{\text{Ideal}}$ for all possible combinations of one or two clusters. The final value was the maximum $\text{ARI}_{\text{Ideal}}$ and $\text{RI}_{\text{Ideal}}$ values of all these combinations.

Additionally, the stability of the GA was estimated by comparing the 30 segmentations from the 30 different runs. This was done by averaging $\text{RI}$ and $\text{ARI}$ comparing all possible pairs of segmentations ($\text{ARI}_{\text{Seeds}}$ and $\text{RI}_{\text{Seeds}}$).

### 4. Experimental setting

As stated before, we study and analyse Dansgaard–Oeschger (DO) events (i.e. the result of different tipping points occurred in the past). To do so, the paleoclimate datasets chosen for this study are the Greenland Ice Sheet Project Two (GISP2) and the North Greenland Ice Core Project (NGRIP) $\delta^{18}O$ ice core data [21, 22]. The $\delta^{18}O$ water isotope record is used as a proxy for past atmospheric temperature variations. We focus on the 20-yr resolution $\delta^{18}O$ isotope records from both drilling sites. The dataset is pre-processed by computing a 5-point average to reduce short-term fluctuations within the data. In this way, the time series considered is $y_{i,j}^{\text{avg}} = \frac{1}{5} \sum_{l=j}^{j+4} y_i$.

This paper considers three different objectives, which are explored in this section: 1) Test our segmentation algorithm which provides a set of clustered time segments; 2) analyse the result of this clustering studying the clusters that best represent TPs; and finally 3) study whether future TPs can be predicted via a classification approach.

The source code in Matlab for the methodology developed in this paper will be made available upon acceptance of the manuscript.

#### 4.1. Experimental setting

It is well-known that GAs usually involve adjusting a notable set of parameters. However, their search dynamics can easily adapt to different problems, resulting in a performance that is negligibly affected by minor changes in the parameters. The GA was configured with the following parameters: the number of individuals of the population is $P = 100$. The crossover probability is $p_c = 0.8$ and the mutation probability $p_m = 0.2$. The percentage of cut points to be mutated ($m_p$) is the integer part of the 20% of the number of cut points, and the average segment length for the initialisation is $\bar{l} = 4$. The maximum number of generations is set to $g = 100$, and the $k$-means clustering process is allowed a maximum of 20 iterations ($it$). These parameters were optimised by a trial and error procedure, although the algorithm showed very robust performance to their values. The most important parameters for the final performance of the algorithm were $\bar{l}$ and $k$.

We performed different experiments considering the 10 different fitness functions presented in Section 2.4.3. The value of $k = 5$ for the $k$-means algorithm has been chosen given the previous good results in the NGRIP data [33]. This result indicates that the concept and nature of DO events is too complex to only consider a binary approach. The climate system exhibits a dynamical behaviour with intrinsic variability hence a binary approach is not able to encompass all features present within a DO event, being $k = 5$ a reasonable choice. Moreover, the method can group several DO events together and is still a useful tool to better understand the behaviour of DO events. It is important to recall that the algorithm estimates the optimal segments without any prior information of the DO events. The only information given to the algorithm is the time series and the statistics to use for the clustering (to validate whether the statistics proposed in the literature are useful for characterising paleoclimate TPs and DO events in general). The best segmentation returned by the GA in the last generation is analysed by an evaluation metric based on the onsets of the DO events. This evaluation metric will be presented in the next subsection.

Finally, since GAs are stochastic optimisation algorithms with a embedded random number generator, their results can be different depending on the seed. To obtain more robust results, the algorithm is run several times with different seeds. For each dataset, the GA was run 30 times with different seeds.

#### 4.2. Segmentation results

The results of these experiments for the two datasets are included in Table 1. As can be seen, the fitness function that results in the best set of segmentations for both datasets is the $CH$ index, which achieves very promising performance. However, as indicated by the $\text{ARI}_{\text{Seeds}}$ and $\text{RI}_{\text{Seeds}}$ metrics, it presents a low stability. In this sense, both $DB$ and $DU$ present an acceptable trade-off between all evaluation metrics.

To properly analyse the evolution process of our algorithm, we have compared the initial and final solutions using the seed that provides the best results with the $CH$ fitness function, resulting in the following performance: For NGRIP the initial solution obtained 0.371 and 0.812 for $\text{ARI}_{\text{Ideal}}$ and $\text{RI}_{\text{Ideal}}$, respectively, and the final solution obtained 0.565 and 0.855. For GISP2, the initial solution resulted in a 0.394 of $\text{ARI}_{\text{Ideal}}$ and a 0.781 of $\text{RI}_{\text{Ideal}}$, while the final solution obtained 0.576 and 0.864. This performance demonstrates the capability of the proposed algorithm to capture statistical features of different time segments and cluster together similar patterns, specially in relation to paleoclimate data.

The segmentations obtaining the highest $\text{ARI}_{\text{Ideal}}$ metric for the $CH$ fitness function (with a value of 0.565 and 0.576 for NGRIP and GISP2, respectively), along with a representation of the 18 DO events can be seen in Figures 4 and 5. The segments have been coloured according to their cluster assignment. Moreover, Figures 6 and 7 include the individual cut points obtained by the algorithm. Note that those segments belonging to the same cluster in Figures 6 and 7 are grouped together in Figures 4 and 5. For the sake of visualisation, Figure 8 represents a zoom of Figure 6. The clusters related to the DO events are $C_2$ and $C_3$ in both time series. If one compares the NGRIP segmentation to the ideal one in Figure 3, it can be seen that almost all DO events are correctly segmented by the algorithm ($C_2$ and
C4 segments are usually close to the DO onset) and that there are very few “false positives” labels (C2 and C4 segments are not found in a non DO event part of the series). For the case of NGRIP, there are four events that are not detected: 2, 9, 13 and 16. In the case of GISP2 there are only two events that have not been detected: 9 and 13.

The clustering space of the best segmentations produced can be analysed in Figures 9 and 10. These Figures confirm that there are obvious differences between the two clusters associated to the DO events (C2 and C4), mainly from the values of the $MS_E$ and $a_j$ metrics.

### Table 1: Mean and standard deviation segmentation results for the different clustering indexes and time series used.

<table>
<thead>
<tr>
<th>Fitness</th>
<th>ARI_Ideal</th>
<th>RI_Ideal</th>
<th>ARI_Seeds</th>
<th>RI_Seeds</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>NGRIP time series</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DB</td>
<td>0.330 ± 0.079</td>
<td>0.791 ± 0.021</td>
<td>0.368 ± 0.079</td>
<td>0.729 ± 0.039</td>
</tr>
<tr>
<td>DU</td>
<td>0.307 ± 0.056</td>
<td>0.793 ± 0.014</td>
<td>0.381 ± 0.090</td>
<td>0.737 ± 0.042</td>
</tr>
<tr>
<td>CH</td>
<td><strong>0.429 ± 0.074</strong></td>
<td><strong>0.823 ± 0.019</strong></td>
<td>0.110 ± 0.031</td>
<td>0.602 ± 0.019</td>
</tr>
<tr>
<td>SSE</td>
<td>0.349 ± 0.068</td>
<td>0.793 ± 0.023</td>
<td>0.130 ± 0.033</td>
<td>0.613 ± 0.015</td>
</tr>
<tr>
<td>NSSE</td>
<td>0.332 ± 0.072</td>
<td>0.795 ± 0.017</td>
<td>0.126 ± 0.030</td>
<td>0.593 ± 0.023</td>
</tr>
<tr>
<td>SH</td>
<td>0.270 ± 0.050</td>
<td>0.787 ± 0.010</td>
<td>0.368 ± 0.095</td>
<td>0.712 ± 0.045</td>
</tr>
<tr>
<td>GD33</td>
<td>0.288 ± 0.062</td>
<td>0.791 ± 0.012</td>
<td>0.357 ± 0.091</td>
<td>0.731 ± 0.044</td>
</tr>
<tr>
<td>GD43</td>
<td>0.289 ± 0.059</td>
<td>0.792 ± 0.012</td>
<td>0.356 ± 0.091</td>
<td>0.726 ± 0.045</td>
</tr>
<tr>
<td>GD53</td>
<td>0.339 ± 0.063</td>
<td>0.797 ± 0.014</td>
<td>0.149 ± 0.070</td>
<td>0.573 ± 0.047</td>
</tr>
<tr>
<td>COP</td>
<td>0.394 ± 0.052</td>
<td>0.81I ± 0.020</td>
<td>0.103 ± 0.035</td>
<td>0.579 ± 0.024</td>
</tr>
<tr>
<td><strong>GISP2 time series</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DB</td>
<td>0.365 ± 0.073</td>
<td>0.788 ± 0.023</td>
<td>0.363 ± 0.082</td>
<td>0.716 ± 0.039</td>
</tr>
<tr>
<td>DU</td>
<td>0.363 ± 0.069</td>
<td>0.794 ± 0.016</td>
<td>0.360 ± 0.087</td>
<td><strong>0.718 ± 0.042</strong></td>
</tr>
<tr>
<td>CH</td>
<td><strong>0.448 ± 0.063</strong></td>
<td><strong>0.817 ± 0.023</strong></td>
<td>0.121 ± 0.028</td>
<td>0.601 ± 0.038</td>
</tr>
<tr>
<td>SSE</td>
<td>0.348 ± 0.055</td>
<td>0.788 ± 0.017</td>
<td>0.119 ± 0.030</td>
<td>0.620 ± 0.015</td>
</tr>
<tr>
<td>NSSE</td>
<td>0.354 ± 0.066</td>
<td>0.775 ± 0.025</td>
<td>0.133 ± 0.034</td>
<td>0.601 ± 0.025</td>
</tr>
<tr>
<td>SH</td>
<td>0.296 ± 0.051</td>
<td>0.775 ± 0.012</td>
<td><strong>0.374 ± 0.087</strong></td>
<td>0.702 ± 0.044</td>
</tr>
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<td>0.303 ± 0.081</td>
<td>0.689 ± 0.041</td>
</tr>
<tr>
<td>GD43</td>
<td>0.325 ± 0.079</td>
<td>0.792 ± 0.017</td>
<td>0.311 ± 0.096</td>
<td>0.698 ± 0.046</td>
</tr>
<tr>
<td>GD53</td>
<td>0.377 ± 0.073</td>
<td>0.797 ± 0.019</td>
<td>0.148 ± 0.066</td>
<td>0.577 ± 0.038</td>
</tr>
<tr>
<td>COP</td>
<td>0.385 ± 0.067</td>
<td>0.796 ± 0.021</td>
<td>0.104 ± 0.031</td>
<td>0.576 ± 0.032</td>
</tr>
</tbody>
</table>

### 4.3. Prediction experiment

As stated before, the main motivation for using a segmentation algorithm is to produce a set of time segments according to a specific criteria (in our case a set of statistical features). This segmentation will provide the expert with a predefined set of time spans to analyse and study different patterns or events (e.g. analyse time segments associated to TPs to validate the hypotheses in the literature about the increase in different statistics). In relation to paleoclimate data, tipping points have been studied using different time-windows (usually, each window is fixed depending on the DO event). In this section, we validate the hypothesis that the time segments can be helpful both for understanding the problem considered and for constructing a...
Figure 4: Best cluster assignation obtained in the NGRIP dataset after the evolutionary process using the CH clustering index.

Figure 5: Best cluster assignation obtained in the GISP2 dataset after the evolutionary process using the CH clustering index.

Figure 6: Time cuts obtained for the best cluster assignation obtained in the NGRIP dataset.

Figure 7: Time cuts obtained for the best cluster assignation obtained in the GISP2 dataset.
Finally, our proposal is to mix both sources of information (not only use the statistical features associated to the previous segment; finally, our approach considers these three sources of information).

The results have been reported in terms of two metrics, one of them specially designed to deal with imbalanced datasets:

1. The well-known Accuracy metric \( (Acc) \), that corresponds to the ratio of correctly classified patterns and measures overall performance.
2. The Geometric Mean of the sensitivities \( (GM = 100 \cdot \sqrt{S_p \cdot S_n}) \), where \( S_p \) is the sensitivity for the positive class (ratio of correctly classified patterns considering only this class) and \( S_n \) is the sensitivity for the negative one. The importance of this metric lies in the imbalanced nature of the dataset (note that we have information related to only 18 DO events).

The algorithm considered is a C4.5 decision tree, in order to interpret the results and provide domain-knowledge. The confidence factor used for pruning is set to 0.25 and the minimum number of objects per leaf to 1. The results associated to this experiment can be seen in Table 2, where the results are provided using all the data both for training and testing the model (this experiment is referred to as Train in the table) and using a leave-one-out approach (LOO column in the table). With the purpose of extracting domain-knowledge we consider that the Train approach describes better the training data (possibly incurring in overfitting), while the LOO approach can also provide information about the generalisation of the model. The “Fixed-window” approach in Table 2 considers the three time series values preceding each segment (i.e. the prediction is performed using only three features). The “Dynamic-window” approach includes the six statistics and the cluster assignment of the segment preceding the one to be predicted (i.e. seven features). The “Combination” approach combines both sources of information, thus resulting in a set of 10 features. In the case of the fixed-window approach, different window sizes were tested, although the results were the same in all the cases. For our proposal, we checked that increasing the size of the window did not have an influence in the model.

Several conclusions can be drawn from this experiment. Firstly, it can be noted that in both cases the fixed-window approach provides a trivial solution (resulting in a model that predicts that all time segments are non DO events). In other words, none of the DO events are classified (which can be inferred by the \( GM \) result). This confirms the previous claim in the literature that all TPs cannot be predicted with the same window size and that the proposed statistics can be used as triggers for TPs. Concerning the sole use of the statistics (i.e. the “Dynamic-window” approach), the result in GISP2 for Train is more promising, although as seen it does not generalise well on unseen data, which could mean that these statistics can be a good predictor for past TPs, but they lack generalisation capabilities. However,
the combination of both sources of information (values of the time series using a fixed-window and statistics computed over a dynamic-window) is successful (specially for GISP2, which was the time series in which the GA obtained the best segmentation results). Another important result is the fact that the segmentation algorithm results in time segments and clusters that are helpful for the prediction and that a diagnostic-predictive model is feasible in the context considered.

The decision trees obtained from this experiment can be seen in Figures 11 and 12. Each node of the tree represents a condition. When the condition is met, the right part of the tree is chosen. The two classes considered are: the negative class or \( C_− \) (associated to non DO events) and the positive class or \( C_+ \) (DO events). Note that for the prediction of the class associated to segment \( s_s \) the characteristics considered are those associated to segment \( s_{s−1} \) and the three time instants before \( s_s \) (i.e., \( y_{t−1−1}, y_{t−1−2}, y_{t−1−3} \)). Note that the conditions in the decision tree are relative comparisons between the existing classes.

For both trees, the time instant previous to the beginning of the segment \( y_{t−1−1} \) is the most discriminating feature (as it is the one that appears at the top of the tree) and is also used in other nodes of the trees. In the case of NGRIP (Figure 11), the cluster of the previous segment is an important characteristic, being DOs usually preceded by \( C_5 \). The used statistics are in this case the autocorrelation and the variance. For GISP2 (Figure 12) the used statistics are kurtosis, skewness and slope. There are two possible outcomes for segment \( s_s \), it either belongs already to a DO onset segment which is found by comparison to the statistics encountered in the previous segment, or it does not belong to the DO onset, something that could
change when the next observation in time is available (i.e. the length of segment $s_{t-1}$ increases by one additional point). That could change significantly the statistical properties of the extended segment. At the addition of any point in observation the segment evaluation should be repeated. It is important to emphasize once again the duration of a segment before being characterised to one of the classes of the DO sequence that exceeds the single next time step horizon.

Table 2: Acc and GM results for the prediction of DO events using the NGRIP and GISP2 time series.

<table>
<thead>
<tr>
<th></th>
<th>NGRIP Train LOO</th>
<th>GISP2 Train LOO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Features</td>
<td>Acc</td>
<td>GM</td>
</tr>
<tr>
<td>Fixed-window</td>
<td>90.16</td>
<td>0.00</td>
</tr>
<tr>
<td>Dynamic-window</td>
<td>90.16</td>
<td>0.00</td>
</tr>
<tr>
<td>Combination</td>
<td>93.99</td>
<td>62.26</td>
</tr>
</tbody>
</table>

5. Conclusions

This work tackles the problem of time series segmentation in the context of paleoclimate time series analysis. We propose a Genetic Algorithm for time series segmentation. which also clusters the time series by using six statistic characteristics that have been found to reveal incoming tipping points (TPs). As opposed to other proposals in the literature, which design TP-specific descriptors, our algorithm considers a generic approach (which has been shown to detect most abrupt climate changes in the series considered). Our experiments also demonstrate that the development of a machine learning-based diagnostic-predictive model is feasible and leads to very promising results given that the predictive-monitoring period of the system is several decades long and that the time series contains already abrupt climate changes of the same type in its record. The above aspects open an interesting avenue for future research.

Future work includes extending the segmentation method to consider simultaneously other related time series to create a more robust and generic decision tree in order to extract domain-knowledge. Finally, other clustering strategies can be helpful for the algorithm, e.g. the DBSCAN algorithm, which provides an outlier detection framework (which could be helpful in a time series as the one considered where there could be noise and very different influencing factors).

Acknowledgments

This work has been partially subsidized by the TIN2014-54583-C2-1-R project of the Spanish Ministerial Commission of Science and Technology (MINECO), FEDER funds and the P2011-TIC-7508 project of the “Junta de Andalucía” (Spain). A. Durán-Rosal’s research has been subsidized by the FPU Pre-doctoral Program (Spanish Ministry of Education and Science), grant reference FPU14/03039. The authors M. Pérez-Ortiz and A. Durán-Rosal have contributed equally to the preparation of this paper.

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