Explainable Clustering Applied to the Definition of Terrestrial Biomes

Mohamed Redha Sidoumou\(^1\)\(^a\), Alisa Kim\(^2\), Jeremy Walton\(^3\)\(^b\), Douglas I. Kelley\(^4\)\(^c\), Robert J. Parker\(^5\)\(^d\) and Ranjini Swaminathan\(^6\)\(^e\)

\(^1\)Amazon Web Services, U.K.
\(^2\)Amazon Web Services, Germany
\(^3\)Met Office Hadley Centre for Climate Science and Services, Exeter, U.K.
\(^4\)UK Centre for Ecology and Hydrology, Wallingford, U.K.
\(^5\)National Centre for Earth Observation, Space Park Leicester, University of Leicester, U.K.
\(^6\)National Centre for Earth Observation, Department of Meteorology, University of Reading, U.K.

\{redsid, alisakim\}@amazon.com, jeremy.walton@metoffice.gov.uk, douglas.i.kelley@gmail.com, rjp23@leicester.ac.uk, r.swaminathan@reading.ac.uk

Keywords: Explainable AI, Explainability, Biomes, Clustering, Segmentation

Abstract: We present an explainable clustering approach for use with 3D tensor data and use it to define terrestrial biomes from observations in an automatic, data-driven fashion. Our approach allows us to use a larger number of features than is feasible for current empirical methods for defining biomes, which typically rely on expert knowledge and are inherently more subjective than our approach. The data consists of 2D maps of geophysical observation variables, which are rescaled and stacked to form a 3D tensor. We adapt an image segmentation algorithm to divide the tensor into homogeneous regions before partitioning the data using the k-means algorithm. We add explainability to the classification by approximating the clusters with a compact decision tree whose size is limited. Preliminary results show that, with a few exceptions, each cluster represents a biome which can be defined with a single decision rule.

1 INTRODUCTION

Natural environment data are complex, noisy and challenging to analyse. However, interesting results—see, for example (Ben-Dor et al., 1999)—can be obtained by the application of data clustering, which aims to group similar data points according to some chosen measure. Clustering is an example of unsupervised learning, in which algorithms find structures and relationships in the data without making use of labels applied to the data, or with any preconceptions of patterns in the data. Many clustering algorithms have been developed such as k-means (Hartigan and Wong, 1979) and hierarchical clustering (Johnson, 1967). However, before computational clustering methods existed, experts used empirical methods to categorise their data, relying on their domain-specific knowledge and expertise.

Terrestrial biomes are constructs that cluster together similar geographical areas. More specifically, a biome is defined as a community of plants with similar functions which form in response to a shared climate. Examples of biomes include grassland, tropical rainforest and desert. Biome distributions affect life on Earth and represent helpful constructs for the organization of knowledge about ecosystems. Clustering is a natural choice for the characterization of biomes. Experts have used empirical approaches using data derived from precipitation and temperature to make such constructs, leading to the Köppen-Geiger (KG) tree model (Peel et al., 2007), built using rules to decompose the terrestrial map into distinct biomes. These methods rely on expert assessment and interpretation of observational data to assess future environmental change. While this has led to a plethora of biome representations, very few of these biome maps target the comparatively coarser scale vegetation cover changes that are associated with numerical Dynamic Global Vegetation Models (DGVMs) and
the climate models which incorporate them. Modelling studies are often forced to re-interpret biome maps designed for very different ecological purposes. In addition, the specific motivation for an ecological study can affect which variables are selected, causing this to be less objective.

An alternative approach is to use expert-led decisions to partition a small number of bioclimatic variables into biomes (Prentice et al., 2011; Sato et al., 2021). While these are custom-made for climate model resolutions, they are restricted to a small number of variables—for example, (Prentice et al., 2011) uses between two and five variables, depending on the biome of interest—due to limits on the number of features which can be assessed by human experts. Biome definitions can also cut across areas of dense bioclimatic space, suggesting biome boundaries and expert assessment for biome definitions do not necessarily align.

Despite their shortcomings, these empirical approaches have the merit of being interpretable—that is, the reasons for the classification of each biome can be understood. Whilst interpretability is not mandatory to validate and use a particular clustering method, it is a feature which appeals to experts, offering a way to understand the results and learn from the findings. For this reason, we seek a clustering methodology with an application to terrestrial biomes which, in addition to being objective, automatic and data-driven, is also interpretable.

2 RELATED WORK

The biome map of Olson and his colleagues (Olson et al., 2001) is perhaps the most commonly used in DGVMs—e.g., (Sellar et al., 2019; Forrest et al., 2020). This is a meta-analysis of previously published ecosystem maps which have been grouped into biomes in consultation with regional experts and global ecologists. The map, commissioned by the World Wide Fund for Nature, is designed specifically for the global coordination of regional and local conservation efforts—a very different purpose to global land surface model evaluation. The Olson biome map provides more detail than can be represented by DGVMs or climate models, and as a result, global vegetation studies have aggregated Olson biomes (in a fashion which is largely inconsistent).

The KG tree model is more targeted to global biome distributions, and has been considered as the standard method for biome classification (Kottek et al., 2006). KG uses temperature and precipitation measurements with decisions based on expert opinion. This type of expert-based model tends to generalise well but can be prone to bias arising from the prior experience of those designing it. The definition of a set of subjective classes is required before the model is defined. In (Thornthwaite, 1948) the authors mitigate the KG model’s simplicity by using variables that are related to moisture and temperature values (as opposed to direct use of precipitation and temperature), although expert-related biases will still be present because of the subjective nature of the classification approach.

There have been a few attempts to define biomes automatically from data. Among them, (Netzel and Stepinski, 2016) represents variables as a time-series of mean monthly observations. The time-series data are then encoded as dissimilarity matrices using dynamic time warping dissimilarity functions or Euclidean distances which take every pair of values from the time series to produce the matrix. Finally, a clustering algorithm is applied to the dissimilarity matrix to create the clusters. Clustering algorithms used include hierarchical clustering with Ward linkage (Ward, 1963) and k-medoids (Kaufman and Rousseeuw, 1987). Finally, (Netzel and Stepinski, 2016) use information theory to deduce that clustering methods are superior to heuristic approaches such as the KG model. (DeSantis et al., 2020) encode observational data as tensors to create a spatial-temporal link between data. A discrete wavelet transform is employed to generate coefficients which are then used in a clustering algorithm. Several scales are used, and for each the authors note the different clusters produced along with their variations.

These clustering methods produce biomes that are not easily achievable by empirical methods because they take many more features into account and usually optimise a well-defined objective function. However, they lack interpretability, a concept often requested by domain knowledge experts because it can help them update their knowledge or assess the limits and potential of interpretable methods.

3 DATA

To determine the distribution of biomes, we used a mixture of land surface (e.g. tree cover, population density) and climate (e.g. mean annual precipitation, mean annual dry days) variables—see Table 1 for details. We chose the set of variables to reflect the properties of large-scale climate and vegetation distributions. The selection of variables used consists of typical parameters that are either measurable from observations or which can be modelled by DGVMs.
This allows our approach to be applied in the future to biome characterization using data from climate models (in particular, we are interested in using it for output from Earth system models—see, for example (Sellar et al., 2019)—which are climate models that explicitly model the movement of carbon through the Earth system).

Table 1: Variables names and their descriptions

<table>
<thead>
<tr>
<th>Variable name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>MADD</td>
<td>Mean annual dry days (i.e seasonality of rainfall)</td>
</tr>
<tr>
<td>MAP</td>
<td>Mean annual precipitation</td>
</tr>
<tr>
<td>MAT</td>
<td>Mean annual temperature</td>
</tr>
<tr>
<td>MTCM</td>
<td>Mean minumum temperature of the coldest month</td>
</tr>
<tr>
<td>MTWM</td>
<td>Mean maximum temperature of the warmest month</td>
</tr>
<tr>
<td>MaxWind</td>
<td>Mean Maximum windspeed</td>
</tr>
<tr>
<td>PopDen</td>
<td>Population density</td>
</tr>
<tr>
<td>DirectRadiation</td>
<td>Direct downwards shortwave radiation</td>
</tr>
<tr>
<td>DiffuseRadiation</td>
<td>Diffuse downwards shortwave radiation</td>
</tr>
<tr>
<td>TreeCover</td>
<td>Tree coverage</td>
</tr>
<tr>
<td>Crop</td>
<td>Cropland area</td>
</tr>
<tr>
<td>nonTreeCover</td>
<td>i.e grasses</td>
</tr>
<tr>
<td>Pasture</td>
<td>Pasture area</td>
</tr>
<tr>
<td>Urban</td>
<td>Urban area</td>
</tr>
<tr>
<td>BurntArea</td>
<td>Burnt area</td>
</tr>
</tbody>
</table>

We obtained TreeCover and nonTreeCover from the MODIS (Moderate Resolution Imaging Spectroradiometer) Vegetation Continuous Fields (VCF) collection 6 fractional tree cover (Dimiceli et al., 2015) regrided as per (Kelley et al., 2019). Although (Adzhar et al., 2021) have shown VCF underestimates tree cover in landscapes which have a mixture of tree and grass, VCF is the only readily available global tree cover product not based on tile aggregation. We obtained Urban, PopDen, Crop and Pasture from the History Database of the Global Environment, Version 3.1 (HYDEv3.1) (Klein Goldewijk et al., 2011). We selected the Global Fire Emissions Database, Version 4.1 (GFEDv4.1) for BurntArea (Van Der Werf et al., 2017), and version 4.01 of the Climatic Research Unit Time Series high resolution gridded dataset (CRU TS v4.01) (Harris and Jones, 2017) for MADD and MAP. (Kelley et al., 2021a) have demonstrated that MADD is likely the best proxy for rainfall distribution controls on vegetation distribution. We used the approach in (Davis et al., 2017) to calculate DirectRadiation and DiffuseRadiation from CRU TS v4.03 monthly cloud cover (Harris and Jones, 2017). MTWM and MTCM, also from CRU TS v4.03, were used to measure heat and frost stress. We take MaxWind from the CRU-NCEP (National Centers for Environmental Prediction) data (Harris, 2019).

We largely pre-processed variables in the same way as (Kelley et al., 2021a), and regridded to a N96 (1.25° x 1.875° spatial resolution) climate model grid. We also aggregated the Olson biome map in the same way as was done in (Kelley et al., 2021a), but to an N96 grid instead of a 0.5° x 0.5° grid.

We retain correlation between variables and do not perform feature selection because this would risk injecting bias. This is not a standard practice in machine learning problems, which suggests treating correlated features first. However, correlation has different effects in different algorithms. For example, standard decision trees algorithms such as CART (Classification and Regression Trees) (Breiman et al., 2017) and C4.5 (Quinlan, 2014) are immune to it and the resulted trees will not be affected. To put correlation into context, we need to analyse its behaviour in clustering and particularly its effect on our k-means with the Euclidean distance metric approach.

K-means algorithm minimises the variance within clusters (Lloyd, 1982). As a result, it produces clusters that are near spherical (or hyper-spherical in higher dimensions). To better achieve this, we applied rescaling to our variables to ensure ellipses become more spherical. Another implication of using k-means without any dimensionality reduction is that correlated features will have a larger impact on cluster formation than uncorrelated features. A variant of k-means uses the Mahalanobis distance (Mahalanobis, 1936) between data points and cluster distributions instead of the Euclidean distance between data points. The Mahalanobis distance is the difference between data points and clusters variances, leading to correlation-invariant results. We however do not use this variant because latent features are not necessarily all equal when defining clusters and we want to keep their influence proportional to the number of observed features they are related to.

4 EXPLAINABLE CLUSTERING METHODOLOGY

We seek to develop a method that can define biomes from features in an interpretable way. The goal of interpretability here is to be able to compare with the empirical biome definitions produced by experts using methods such as (Olson et al., 2001). It also allows an evaluation of the rule-based biome definition and contrasts it with less explainable methods. Our
approach is composed of the following steps:

1. Rescale the data and stack it into a tensor.
2. Use Felzenszwalb’s segmentation algorithm (Felzenszwalb and Huttenlocher, 2004) on the tensor data to decompose it into regions. This step helps to produce smoother biomes and alleviates the effects of outliers without removing them.
3. Transform the newly formed segments into a tabular dataset $D$ where each segment represents a row of data. The values of each row are the median values of the pixels’ segments.
4. Use k-means to cluster $D$ into $k$ clusters.
5. Produce a decision tree taking the inputs as the dataset $D$ and the labels as the $k$ cluster identifiers (or pseudo-labels) with a set of hyper-parameters allowing a good compromise between accuracy and interpretability.

We give details about each step of our approach in the following sections.

4.1 Data pre-processing

Our dataset consists of 15 global land surface variables on a latitude-longitude N96 climate model grid whose dimensions are 180 (longitude) by 142 (latitude). We recalculate each variable and then stack them at each grid cell to produce an image whose pixels have 15 channels. This can be viewed as a 3D tensor whose dimensions are longitude, latitude and channel.

4.2 Processing outliers

Some pixels may contain values that could be considered as outliers, or noise resulting from errors associated with the data or other conditions. If we cluster such pixels into groups, neighbouring pixels will belong to different clusters which will result in a grained distribution—for example, tiny clusters represented by individual pixels within a large cluster. Figure 1 represents such phenomena. To alleviate the effects of outliers, we use an image segmentation-based approach. Although our data are not images in the conventional sense, they have an analogous structure—i.e. our data points have a two-dimensional spatial relationship in which proximity and location is significant.

Image segmentation methods decompose images into regions whose pixels have similar characteristics. The pixels within every segment are rather similar but
not identical. They also tend to be localized (i.e. pixels which are far apart are unlikely to be part of the same segment).

We chose Felzenszwalb’s (Felzenszwalb and Huttenlocher, 2004) algorithm as a segmentation that does not require a fixed number of segments in advance (unlike SLIC superpixels (Achanta et al., 2012) for example). Felzenszwalb is a graph-based algorithm that ignores low variabilities in high variability regions, resulting in segments which appear more natural. There is no restriction on the shape of the segments it produces. While images encoded as matrices typically have three channels (e.g. RGB, or HSL values), there are no restrictions on the number of channels this algorithm can work with.

Applying Felzenszwalb’s algorithm with the parameters $\text{scale} = 0.4, \text{sigma} = 0.5, \text{minsize} = 3$ produces 1663 segments (see left hand side of Figure 1). The $\text{scale}$ parameter controls the size of the segments—larger values produce larger segments. $\text{minsize}$ is the minimum number of pixels which each segment can contain, and $\text{sigma}$ is a pre-processing parameter that controls the dimensions of the Gaussian kernel. Close inspection reveals that certain small non-terrestrial (water) regions were mistaken for noise due to their small size. A second pass of the segmentation method grouped these non-terrestrial regions into one segment, which we then removed (along with segments corresponding to Antarctica) since these regions have no terrestrial biomes of interest. The right hand side of Figure 1 shows the 1216 segments which remained after this removal; comparing the two sides of the figure shows the improvement in the resolution of the coastline arising from the removal of the non-terrestrial regions.

$15$ values characterise each segment—that is, the median of each of the $15$ variables within the segment’s boundary.

\[v_k(\text{segment}_i) = \text{median}(x_{ki})\]  

where $v_k$ is variable $k$’s value for segment $i$, $j$ is a point index within its segment, and $x_{ki}$ is the value of variable $k$ of data point $j$ belonging to segment $i$.

### 4.3 Clustering the map via k-means

K-means is a distance-based algorithm that groups points with similar feature values by assigning each point to the cluster with the nearest centroid. It finds centroids that minimise the sum of the squared Euclidean distance between each data point and its associated centroid:

\[
\argmin_{\mu} \sum_{i=0}^{m} ||x_i - \mu||^2
\]  

Figure 3: A plot of gap value against cluster count. This is one of the methods used to determine a suitable cluster size (see text).

where $x_i$ are the vectors representing data points $i$ and $\mu_i$ are the centroid vectors associated with data point $i$.

We used k-means with all $15$ variables for our analysis. We acknowledge that some of these variables are correlated as they are influenced by the same underlying physical processes and that such correlations might determine cluster associations. However, we did not see any evidence that the semantic integrity of our clusters was significantly impacted by these correlations and so we chose to retain all $15$ variables.

The k-means clustering algorithm requires the number of clusters (that is, $k$) as one of the hyperparameters. The selection of a value for this parameter requires some consideration. For example, we could set it to the number of biomes used in an empirical characterization approach for comparative purposes, or to a higher or lower value to obtain a more generalized or more granular result. We employed three different evaluation metrics to estimate the optimal number of clusters: the Distortion Coefficient, Gap values and the Silhouette score. We show an illustration of how we employed one of the metrics (Gap values) in Figure 3. Based on this metric, we determined that a number between $10$ and $15$ (clusters) would optimally represent our data. We found that the Distortion Coefficient and the Silhouette scores were in agreement with this range, suggesting a comparable number of clusters.

We finally chose to employ $11$ clusters for defining biomes in this approach – a number consistent with both expectations from domain-expertise and the evaluation metrics we employed.

### 4.4 Tree-based interpretability

Machine learning models that are not easy to explain in simple terms require the application of specific extra methods, including SHAP (SHapley Additive exPlanations) (Lundberg and Lee, 2017) and
LIME (Local Interpretable Model-agnostic Explanations) (Ribeiro et al., 2016). Models which are more explainable (for example, a decision tree with a comparatively low depth) do not need methods such as these to be applied. Such models include decision trees, linear regressions and naive Bayes. These models belong to the supervised learning family. Explainable models in clustering and unsupervised learning in general are very scarce.

Decision rules provide an explainable way to define categories. Each category is bounded by a set of conditions, generally Boolean. A number of algorithms exist that can generate decision rules for a given problem. (Chavent, 1998) builds decision rules directly for clustering purposes on unlabelled data. It uses a similarity measure to decide on dividing clusters of data by finding the optimal splitting value on features. The process is similar to the splits used in decision trees. The method builds hyper-cubic shaped clusters but does not put a restriction on the number of rules or the number of features used. This method optimises the decisions at each level instead of optimising the whole structure of the clusters. (Dasgupta et al., 2020) uses first k-means to build clusters, and then builds a minimum set of decision rules on top of these clusters to approximate them. The simpler the rules, and the smaller the set, the more explainable the clustering will be. Another method of building explainable clustering is to approximate k-means clusters with a decision tree while restricting the size of the tree via its hyper-parameters. We use this approach. Since decision boundaries in decision trees are made of vertical and horizontal lines and k-means clusters boundaries are spherical-like, we would not expect them to be exactly equivalent. Instead, we choose hyper-parameters that allow a good compromise between interpretability and equivalence. We define two models as equivalent if they assign the same label (or cluster) to a given data point—i.e., if the models agree on the classification of all the data points. We adopt the following algorithm to generate interpretable decision trees:

1. Generate k clusters using k-means (see §4.3). Results are presented in figure 4.
2. Use the resultant clusters identifiers as pseudo labels for the data.
3. Build a decision tree using:
   \[ \text{min\_samples\_per\_node}=30, \text{max\_depth}=9 \]
4. Recursively trim the resulting tree if the predicted classes do not change in sibling leaves

It is worth noting that there is a set of decision trees that can approximate the clustering results while producing explainable rules. The above algorithm generates one among several possibilities. Tweaking the hyper-parameters such as \text{min\_samples\_per\_nodes} and \text{max\_depth} will produce slightly different trees. We note that the algorithms we used to build the trees are (Breiman et al., 2017) and C4.5 (Quinlan, 2014), other clustering or tree-based algorithms will produce different trees. (Dasgupta et al., 2020) describe a k-means clustering result along with a possible decision tree approximation. Our aim here, however, is to explore the effect of the addition of interpretability to k-means clustering, in particular whether it enhances the comparison with results from empirical classification methods for terrestrial biomes. To better satisfy interpretability, we have two goals that will guide us during the tree’s construction. The first one is to produce short rules, which is equivalent to the decision tree length. The second aim is to reduce the number of rules per cluster to as close to one as possible. These goals will help deliver the definition of biomes in a fashion which is more accessible to the domain knowledge experts who can examine, strip, augment, or even discard them. Such rules will also be comparable in structure to the empirical rules from models such as KG tree or expert knowledge such as (Olson et al., 2001).

4.5 Results

Having applied the methodology described in (§§4.1–4.4) to generate explainable biomes, we present our results and compare them to those from the empirical biome attributions. Upon a partial application of our method, without selecting the hyper-parameters that account for short and small numbers of rules and tree trimming, the resultant decision tree (called TreeA) approximates k-means clustering with an agreement of 95%. Although the decision tree gives a very good representation of the original clusters, it uses around
80 rules to define all the biomes. This is a comparatively large number, representing a challenge for the usability of the classification scheme, and for comparisons with the results from the empirical characterization approach. Therefore, we compromise agreement with the clusters and usability by setting values for the hyper-parameters in the decision tree algorithm, defining the minimum number of samples per leaf `min_samples_per_nodes` to 30, limiting the tree depth. We also recursively trim the tree, using the following algorithm:

Order pairs of sister leaves based on length for each pair of sister leaves a,b:
   if majority_class(a) == majority_class(b):
      c=merge(a,b)
      d=find_sister_leaf(c)
      insert pair(c,d) in list of sister leaves keeping the order
end

Applying the algorithm with the hyper-parameter values mentioned above and using recursive trimming produces decision tree `TreeB` which is in 85% agreement with the original clusters. However, the tree contains 14 leaves, which makes it considerably smaller than `TreeA`—see Figure 5. In a trimmed tree all biomes (except three) are explained by a single leaf and thus a single rule. This supports the hypothesis that most biome definitions can be defined in an explainable way. We then re-clustered the regions of the map according to `TreeB`. In this tree, leaves are numbered from 0 to 10. Leaves are assigned to biomes according to the index of the maximum value in the leaf's segments array—for example, a leaf with segments array of `[2, 55, 3, 0, 1, 0, 4, 4, 1, 1, 0]` would represent biome 1.

5 DISCUSSION

By comparing our results to the empirical biomes defined by experts in the literature, we can observe a number of biomes that are in accordance with our results such as the tropical rainforest, the tundra, the taiga, the desert, the steppe, the savanna and the mixed forest with some variations on their boundaries. However, we also observe a couple of extra biomes, one of which is mainly present in the Indian subcontinent, while the other is a semi-arid biome similar to desert. The biome present in the Indian subcontinent seems to be influenced by agriculture and suggests an anthropogenically-induced change. Further analysis of these results is required to compare the boundaries of the discovered biomes to those derived by empirical methods. Figure 7 shows a comparison between Olson’s biomes (Olson et al., 2001) and the biomes derived from the explainable clustering methodology. Typically there is a strong overlap between our derived biomes and those from Olson (for example, the desert biome has a 92% overlap, tropical forest has 87%). Interestingly, when there is some disagreement, a biome is still often attributed as belonging to a similar biome (e.g. boreal forest and tundra) suggesting the boundary between biomes is somewhat fluid. Applying an image segmentation algorithm helped in making biomes more continuous in space, thus ignoring localized small variations. An examination of the decision rules and t-distributed stochastic neighbour embedding (t-SNE) visualizations of the biomes reveals the following:

- Most biomes require only one rule to define them.
- Two leaves do not have a clear majority decision.
- Three biomes (biomes number 4, 6 and 9) each require two rules to be defined.
Figure 7: A comparison of biomes obtained from explainable clusters against Olson’s biomes (Olson et al., 2001). The numbers show the % of clustering biomes that fall in each Olson biome. In biome names, trop = tropical, med = Mediterranean, frst = forest, wood = woodland, shrub = shrubland, scrb = scrubland, grass = grassland, bl = broadleaf. The color of each cell denotes the type of biome (green = forest, blue = woodland, purple = grassy, orange = barren); color intensity is proportional to the value in the cell.

Biome 6 (represented by the pink leaves in Figure 5) covers a large part of the map. Most of its region is governed by a single rule covering 185 segments from the map, whilst the second rule covers only 25 segments. The 25 segments appear to be close to the borders with other biomes.

Similarly, biome 9 represented by the yellow leaf is defined by two rules as well. One rule governs 134 segments, while the other governs 18 segments and another 12 segments from biome 6. Both rules are characterized by a high tree coverage and adjacent intervals of mean annual temperatures. This may allow these two rules to be fused in one single rule.

Looking at the t-SNE plot in figure 6 of the decision tree, we note that biomes 1 and 4 are highly mixed with other biomes in the spatial representation while the t-SNE graph of the k-means clusters does not shows this mixtures. This suggests that these two biomes may not be easily defined by simple rules. Biome 1 which appears to represent the desert is less prone to this effect than biome 4. The latter, which to date appears to be the only biome that the decision tree approach is unable to identify. The fact that it is related to agricultural regions suggests that “non-natural” biomes may not follow the same structure as the others, and the effect of these man-made environmental changes are more complex than those represented by natural observable features. Biome 4 appears to share some characteristics with neighbouring biomes but is sufficiently different to be characterized as a new biome by k-means. However, it is not distinctive enough to be fully explainable. We note that most of the other irregularities in the decision tree-based clustering are located near the borders of biomes in data space.

We also note that decision trees that can approximate k-means are not unique. For this, a further analysis of the different ways of approximating the k-means clusters is required. For example, changing the weights of certain instances will result in a different tree with a possibility of a different interpretability level. There are also tree construction methods that are not based on CART or C4.5 such as the methods presented in (Dasgupta et al., 2020) that can applied to this use case.

6 CONCLUSION

In this work we have defined a methodology for explainable clustering which we have applied to 3D tensor data. Our motivation is the production of explainable definitions of terrestrial biomes. After rescaling
the data, we superpose them as 3D tensors and apply an image segmentation algorithm to decompose the tensor into homogeneous segments. We apply k-means clustering to the segments, and introduce explainability by approximating the clusters with a decision tree, which we subsequently trim to a manageable number of leaves.

This method allowed for the building of continuous biomes in space with one rule per biome in most cases. Three biomes required more than one rule to define them. Four of these rules could be fused into two rules thus making the clustering simpler. However, despite using all 15 features in the construction of the decision tree, the resulting rules used at most four features. This is due to underlying latent features influencing more than one correlated feature. It is worth noting that a decision tree which is relatively small and can approximate clustered data is not unique. Different methods with different hyperparameters will produce different trees. We also noticed that certain biomes that are man-made such as the Indian subcontinent biome cannot be easily defined by rules without affecting the rules of adjacent biomes. This indicates that this biome has an irregular shape (in data space) and indeed requires more than one rule to be defined. One final observation is that decision trees are not perfect in building simple rules for clustering. Because rules are constrained to be connected at different node levels, simplicity may be traded for this property. For future work, we suggest augmenting this method with a strategy for merging rules that not only merges rules when they are connected via a feature but also when features are correlated and there is a possibility for such a merge. These may lead to a loss in precision depending on the correlation level but it will improve the interpretability.

We believe that this study demonstrates the strong potential possible for advancing our understanding of Earth system science by utilising machine learning methods, such as explainable clustering. By expanding this work in the future and applying these methods to climate projections from Earth system models, we will be able to provide analyses which complement existing insights from experts about how the Earth’s biomes may alter in response to a changing climate.

DATA AVAILABILITY

The data used (Table 1) are archived at https://doi.org/10.5281/zenodo.5736407 (Kelley et al., 2021b).

ACKNOWLEDGEMENTS

This project originated as a collaboration between the Met Office and Amazon Web Services (AWS) under the AWS Machine Learning Embark program. JW is supported by the Met Office Hadley Centre Climate Programme funded by BEIS and Defra. RS and RJP are funded by the UK National Centre for Earth Observation (UKESM, NE/N018079/1). DIK was supported by the UK Natural Environment Research Council (UKESM, NE/N017951/1).

REFERENCES


Felzenszwalb, P. F. and Huttenlocher, D. P. (2004). Effi-


Harris, I. (2019). CRU JRA v1.1: A forcings dataset of climatic Research Unit (CRU) and Japanese reanalysis (JRA) data, January 1901–December 2017, University of East Anglia Climatic Research Unit, Centre for Environmental Data Analysis.


