Title: Clustering in non-parametric multivariate analyses.

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Keywords: Non-parametric multivariate; divisive clustering; flat clustering; SIMPROF; cophenetic correlation; cophenetic distance

Abstract: Non-parametric multivariate analyses of complex ecological datasets are widely used. Following appropriate pre-treatment of the data inter-sample resemblances are calculated using appropriate measures. Ordination and clustering derived from these resemblances are used to visualise relationships among samples (or variables). Hierarchical agglomerative clustering with group-average (UPGMA) linkage is often the clustering method chosen. Using an example dataset of zooplankton densities from the Bristol Channel and Severn Estuary, UK, a range of existing and new clustering methods are applied and the results compared. Although the examples focus on analysis of samples, the methods may also be applied to species analysis. Dendrograms derived by hierarchical clustering are compared using cophenetic correlations, which are also used to determine optimum in flexible beta clustering. A plot of cophenetic correlation against original dissimilarities reveals that a tree may be a poor representation of the full multivariate information.UNCTREE is an unconstrained binary divisive clustering algorithm in which values of the ANOSIM R statistic are used to determine (binary) splits in the data, to form a dendrogram. A form of flat clustering, K-R clustering, uses a combination of ANOSIM R and Similarity Profiles (SIMPROF) analyses to determine the optimum value of k, the number of groups into which samples should be clustered, and the sample membership of the groups. Robust outcomes from the application of such a range of differing techniques to the same resemblance matrix, as here, result in greater confidence in the validity of a clustering approach.
Dear Sandy,

It gives me great pleasure to submit for consideration for publication in the journal our latest manuscript, entitled “Clustering in non-parametric multivariate analyses”, quite out of character for Bob in the brevity of its title but, we believe, an important advance in terms of ecological methods for complex data.

Obviously, as required by the journal, all authors are happy to declare that they agree to this submission. I believe that it is all in order, and look forward to hearing from you in due course.

Actually, Bob asked me to submit this while he is on a much needed and hard-earned vacation, and in the course of doing so I nominated him as the corresponding author. I then found that I couldn’t complete the submission as a key step was unavailable to me as a non-corresponding author. I tried twice before I realised this. Thus elsewhere there are two versions of this manuscript in the system, which need to be cleared. Hopefully that will be sorted once he returns. In the meantime, I’m happy to correspond and hope you like the manuscript.

Yours sincerely,

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HIGHLIGHTS

Dendrograms may be poor representations of inter-sample dissimilarities

ANOSIM R and SIMPROF are combined to generate new methods of clustering

UNCTREE is a binary divisive clustering algorithm

$k$-R clustering is a flat clustering algorithm

Robustness of clustering is assessed by applying different methods to example data
Clustering in non-parametric multivariate analyses.

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ABSTRACT

Non-parametric multivariate analyses of complex ecological datasets are widely used. Following appropriate pre-treatment of the data inter-sample resemblances are calculated using appropriate measures. Ordination and clustering derived from these resemblances are used to visualise relationships among samples (or variables). Hierarchical agglomerative clustering with group-average (UPGMA) linkage is often the clustering method chosen. Using an example dataset of zooplankton densities from the Bristol Channel and Severn Estuary, UK, a range of existing and new clustering methods are applied and the results compared. Although the examples focus on analysis of samples, the methods may also be applied to species analysis. Dendrograms derived by hierarchical clustering are compared using cophenetic correlations, which are also used to determine optimum $\beta$ in flexible beta clustering. A plot of cophenetic correlation against original dissimilarities reveals that a tree may be a poor representation of the full multivariate information.

UNCTREE is an unconstrained binary divisive clustering algorithm in which values of the ANOSIM $R$ statistic are used to determine (binary) splits in the data, to form a dendrogram. A form of flat clustering, $k$-R clustering, uses a combination of ANOSIM $R$ and Similarity Profiles (SIMPROF) analyses to determine the optimum value of $k$, the number of groups into which samples should be clustered, and the sample membership of the groups. Robust outcomes from the application of such a range of differing techniques to the same resemblance matrix, as here, result in greater confidence in the validity of a clustering approach.

KEYWORDS: Non-parametric multivariate; divisive clustering; flat clustering; SIMPROF; cophenetic correlation; cophenetic distance
1. Introduction

Field et al. (1982) described a robust non-parametric multivariate strategy for the analysis of biological assemblage data, such as the abundance or biomass of taxa in samples. Collins and Williams (1982) present one of the first applications of the strategy, to plankton data from the Bristol Channel and Severn Estuary. In essence the strategy, expanded and clarified by Clarke (1993), is to display patterns among samples determined by appropriate resemblance measures (Clarke et al., 2006) using clustering and ordination, and to analyse these patterns using a range of hypothesis tests and associated analyses, primarily based on ranked resemblances. Additional analyses are constantly added to the framework. Clarke et al. (2008) described a method for divisive clustering constrained by thresholds in explanatory variables, Linkage Trees, and Similarity Profiles analysis (SIMPROF) which tests for multivariate structure within groups of samples. The latter was further discussed by Somerfield and Clarke (2013) in the context of species (r-mode) analysis.

Literally hundreds of clustering methods exist, some of them operating on resemblance matrices whilst others are based on the original data (Legendre and Legendre, 2012). Everitt (1980) and Cormack (1971) give excellent and readable reviews, while Clifford and Stephenson (1975) is another well-established text from an ecological viewpoint. To cope with this variety a widely adopted approach has been to use a single technique that has been found to be of widespread utility in ecological studies while recommending the need to perform a cluster analysis in conjunction with a range of other techniques (e.g. ordination, statistical testing) to obtain balanced and reliable conclusions (Clarke et al., 2014).

Hierarchical clustering with group-average linking, based on sample similarities or dissimilarities such as the Bray-Curtis coefficient, has proved a useful technique in many ecological studies over the past half-century. As with clustering methods in general, it is appropriate for delineating groups of sites with distinct community structure. It is an agglomerative method. Agglomerative methods are bottom-up and ‘see’ only the nearby points throughout much of the process. When reaching the top of the dendrogram no possibility of taking a different view, of the main merged groups that have formed, remains. Binary divisive methods, however, are potentially advantageous for some clustering situations. They take a top-down view of the samples, so that the initial binary splits should (in theory) be better able to respect any major groupings in the data, since these are found first. However, as with all hierarchical methods, once a sample has been placed within one initial group it cannot jump to another at a later stage. Whilst divisive methods have the potential to produce marginally better solutions in practice, there is a counterbalancing downside to their
algorithms, in that they can be computationally intensive and complex (Gower, 1967), so iterative
approaches are generally required. The agglomerative approach, in contrast, is simple and entirely
determined, requiring nothing more than simple numerical operations based on values of
resemblance measures.

The purpose of this paper is to compare and discuss methods for, and associated with, cluster
analysis in a non-parametric multivariate framework. We look at some existing methods and
consider how their success, in conserving the inter-sample patterns in the underlying resemblance
matrix, may be assessed and compared. We go on to introduce new clustering methods and
compare their results. For the purpose of the paper we restrict ourselves to examples based on
analyses among samples, though it should be remembered that clustering of variables (taxa,
functional groups, OTUs, environmental measurements) is often entirely appropriate following
suitable pre-treatment of the data (Somerfield and Clarke, 2013).

2. Material and methods

2.1 Hierarchical agglomerative clustering

The most commonly used clustering techniques are hierarchical agglomerative methods. These
usually take a resemblance matrix (Clarke et al., 2006) as their starting point and successively fuse
the samples into groups, and the groups into larger clusters, starting with the highest mutual
similarities then lowering the similarity level at which groups are formed, ending when all samples
are in a single cluster. The result of a hierarchical clustering is generally presented as a tree diagram
or dendrogram. There is no firm convention for which way up a dendrogram should be portrayed
(increasing or decreasing resemblance values) or even whether the tree can be placed on its side,
but we will refer to the x axis as representing the full set of samples and the y axis defining a
resemblance level at which two samples or groups are considered to have fused. Neither is there
anything sacrosanct about the ordering of samples along the x axis, with the exception of constraints
imposed by the grouping structure among samples at higher levels in the tree.

2.2. Linkage options

Within hierarchical agglomerative clustering a range of linkage/sorting/joining options are defined
which determine how resemblances between samples and groups of samples are recalculated
following fusion of samples into a group. For single linkage (also called nearest-neighbour joining)
the dissimilarity of groups A and B, $\delta_{A,B}$, is the minimum across all dissimilarities between pairs of
samples with the first in A and the second in B. The dissimilarity of a group C to two merged groups
A and B, \( \delta_{CAB} \), is therefore just the minimum of \( \delta_{CA} \) and \( \delta_{CB} \). For complete linkage (also called
farthest-neighbour joining), \( \delta_{CAB} \) is the maximum of \( \delta_{CA} \) and \( \delta_{CB} \). In group-average linkage \( \delta_{AB} \) is the
simple (unweighted) average over all dissimilarities from A to B pairs, leading to the acronym
UPGMA, Unweighted Pair Group Method with Arithmetic mean. When A and B are of different sizes,
it follows that, under UPGMA, \( \delta_{CAB} \) is a weighted average of \( \delta_{CA} \) and \( \delta_{CB} \), e.g. giving more weight to
\( \delta_{CA} \) if there are more samples in A than B. Somewhat confusingly, the simple average of \( \delta_{CA} \) and \( \delta_{CB} \)
is then referred to as weighted linkage, WPGMA, since it weights the original dissimilarities between
samples in C and those in the combined group A and B unequally.

Other linkage options have been suggested. One is the flexible beta method of Lance and Williams
(1967), in which \( \delta_{CAB} = (1 - \beta) [(\delta_{CA} + \delta_{CB})/2] + \beta \delta_{AB} \). Only negative values of \( \beta \) in the range (-1, 0)
make much sense in theory, the effect of including the \( \delta_{AB} \) term then being to make the merged AB
group more likely to join with the group C, the further A and B themselves are from each other. That
is, there will be a tendency to merge loosely bound samples or groups with each other, leaving
tightly bound groups separate. Lance and Williams (1967) suggest the use of \( \beta = -0.25 \), for which the
flexible beta has affinities with Gower’s median method (Gower 1967). If \( \beta = 0 \), \( \delta_{CAB} = (\delta_{CA}+\delta_{CB})/2 \),
which is the WPMGA method given above, also known as McQuitty’s (1967) linkage.

Within a non-parametric multivariate analytical framework it might be expected that a linkage
option that is a function only of the ranks in the underlying resemblance matrix would be preferred.
Single linkage does this, but experience shows that it leads to ‘chaining’ in the resulting dendrogram,
with samples continuously joined to the next most similar sample without forming discrete clusters.
Complete linkage, conversely, tends to result in starkly separated, compact clusters. Group average
linkage will find a seemingly reasonable balance between the two. In order to choose between
linkage methods and their associated dendrograms a more objective means than simple visual
comparison of dendrograms is clearly needed.

2.3. Cophenetic correlation

One objective approach is provided by cophenetic correlation, which is a (Pearson) matrix correlation
between each original dissimilarity and the (vertical) distance through a dendrogram to the common
node of the corresponding pair of samples (Jain and Dubes, 1988). If the y-axis of the dendrogram is
a dissimilarity scale then, naturally, these vertical distances are also dissimilarities. A dendrogram is a
good representation of the dissimilarity matrix, therefore, if the cophenetic correlation is close to 1.
As such the correlation may be seen as a way to compare different dendrograms, to assess the
performance of different analysis choices starting from the same dissimilarity matrix. In particular, the correlation may also be used to determine $\beta$ for the flexible beta method, computing a range of values and choosing that which maximises the cophenetic correlation.

2.4. Binary divisive clustering

In hierarchical agglomerative clustering, samples start in separate groups and are successively merged until, at some level of similarity, all are considered to belong to a single group. Hierarchical divisive clustering does the converse operation: samples start in a single group and are divided into two sub-groups, which may be of quite unequal size, each of those being further sub-divided into two (i.e. binary division), and so on. Ultimately, all samples become singleton groups unless (preferably) some criterion is applied to stop further sub-division of any specific group. Clarke et al. (2008) describe such a clustering method (LINKTREE), which derives a binary divisive dendrogram from a resemblance matrix. Divisions are constrained by thresholds on individual explanatory variables, and the criterion for assessing the ‘best’ division at each step is to choose the one which maximises the ANOSIM $R$ statistic, defined as the difference between the average of the rank dissimilarities between the (two) groups and within the groups. This is suitably scaled by a divisor of $M/2$, where $M = n(n-1)/2$ is the total number of dissimilarities calculated between all the $n$ samples currently being split. This divisor ensures that $R$ takes its maximum value of 1 when the two groups are perfectly separated, defined as all between-group dissimilarities being larger than any within-group ones. $R$ will be approximately zero when there is no separation of groups at all, but this will never occur in this context since the groups are chosen to maximise the value of $R$. There is an important point not to be missed here: $R$ is in no way being used as a test statistic, the reason for its development (Clarke and Green, 1988). Instead, its value provides a pure measure of separation of groups of points represented by the high-dimensional structure of the resemblances (here perhaps Bray-Curtis, but any coefficient can be used with $R$, including Euclidean distance). This provides a universal scaling of between vs. within group dissimilarities/distances (whatever their measurement scale) through their reduction to simple ranks. A stopping rule is provided naturally here by the SIMPROF test: if there is no demonstrable structure within a group, i.e. the null hypothesis for a SIMPROF test cannot be rejected, then that group is not further sub-divided (Clarke et al., 2008).

The constrained divisive LINKTREE approach (typically used to ‘explain’ a biotic clustering by a series of inequalities on individual environmental variables) permits examination of only a relatively small number of the possible ways of splitting a single group into two sub-groups. A division is only considered if one of the constraining variables takes larger values for all the samples in one sub-
group than the other. This reduces the number of choices at each step to a maximum of $p(n-1)$, where $p$ is the number of constraining variables and $n$ the number of samples in the group currently under division, and full calculation for all valid choices is straightforward. In contrast, an unconstrained divisive approach needs, for each of the current groups, a full (binary) flat clustering, namely computation of some optimality criterion (here the maximisation of ANOSIM $R$) for all possible binary divisions of that group of $n$ samples. There are $2^{n-1} - 1$ possibilities and for even quite modest $n$ (say $>25$) evaluating all of them quickly becomes prohibitive. This necessitates an iterative search procedure (a variant of a standard “$k$-means” algorithm; MacQueen 1967), starting from an arbitrary allocation of samples to the two sub-groups. In turn, each sample is then re-allocated to the opposite group if this increases $R$, or left where it is if that would reduce $R$, and that process continues until an entire loop has passed in which no sample changes its allocation. Such a converged solution will typically vary depending on the starting configuration, so the whole iteration must be repeated many times from different starting configurations. The ‘best’ of the divisions from these different random restarts is then selected as likely, though not guaranteed, to be the optimal solution. The binary divisive procedure (UNCTREE) starts with all samples in a single group, and if a SIMPROF test provides evidence that the group has structure which can be further examined, a search is made for an optimal split of those samples into two groups, maximising $R$. This could produce anything from splitting off a singleton sample through to an even balance of the sub-group sizes. The SIMPROF test is then repeated for each sub-group and this may justify a further split, again based on maximising $R$, but now calculated having re-ranked the dissimilarities in that sub-group. The process repeats until SIMPROF cannot justify further binary division on any branch: groups of two are therefore never split as there is no sensible SIMPROF test for two samples (Clarke et al., 2008).

The resulting groupings may be plotted in a dendrogram, but it is not readily apparent what values on the y axis should be assigned to the various divisions. Unlike hierarchical agglomerative clustering, in which fusions occur at increasing levels of similarity, each division in UNCTREE is determined by a maximum value of $R$, which may be higher or lower than values at other levels in the same analysis. There are two sensible approaches here. The first is to simply plot each level on an arbitrary scale with equal spacing between levels and the level of subsequent divisions set so that they sum to 100. This A% scale allows the plotting of the dendrogram, though gives little sense of the relative importance of the divisions. An alternative is to use the B% scale introduced by Clarke et al. (2008) for LINKTREE, calculating the average of between-group dissimilarities for each division and dividing this by the maximum value it can take for a perfect split ($R = 1$) on the first division, and then multiplying by 100 to give a positive scale which never exceeds 100. This scale measures how
well separated the two groups of samples are in the current division relative to the maximum separation possible in the first division.

2.5. K-R clustering (non-hierarchical)

Another major class of clustering techniques is non-hierarchical, referred to above as flat clustering. The desired number of clusters (k) must be specified in advance, and an iterative search attempts to divide the samples in an optimal way into k groups, in one operation rather than incrementally. The classic method is k-means clustering, which seeks to minimise within-group sums of squares about the k group centroids. This idea, appropriate only for Euclidean distance matrices, can again be generalised to apply to any resemblance measure, e.g. Bray-Curtis, by maximising ANOSIM R, which measures (non-parametrically) the degree of overall separation of the k groups, formed from the ranks in the full resemblance matrix. By analogy with k-means clustering, the principle of maximising R to obtain a k-group division of the samples is referred to as k-R clustering, and it again involves an iterative search, from several different random starting allocations of samples to the k groups.

The k-group solution will not, of course, simply split one of the groups in the (k – 1)-group solution: there could be a wide-scale rearrangement of many of the points into different groups. A widely perceived disadvantage of the k-means idea is the need to specify k before entering the routine. Another, if it is re-run for many different k values, is the absence of a convenient visualisation of the clustering structure for differing values of k, analogous to the hierarchy of a dendrogram. The output of the solution is simply a factor denoting which samples belong to the same groups. This has tended to restrict its use to cases where there is a clear a priori idea of the approximate number of groups required, perhaps for operational reasons (e.g. in a quality classification system). However, the SIMPROF test can also come to the rescue here, to provide an objective choice of k. Starting from a low value for k (say 2) the two groups produced by k-R clustering are tested for evidence of within-group structure by SIMPROF. If either of the tests are significant, the routine increments k (to 3), finds the 3-group solution and retests those groups by SIMPROF. The procedure is repeated until a value for k is reached in which none of the k groups generates significance in their SIMPROF test, and the process terminates with that group structure as the best solution. (This will not, in general, correspond to the maximum R when these optima for each k are compared across all possible k; R must increase to its maximum of 1 as k approaches n, the number of samples.)

A variation of this flat-clustering procedure, rather than based on the R statistic, utilises the concept of group-average linking met earlier, though still in a non-parametric setting. For a pre-specified number of groups (k), each stage of the iteration process involves removing each sample in turn and
then allocating it to one of the other \((k - 1)\) groups currently defined, or returning it to its original group. In \(k-R\) clustering it is re-allocated to the group which produces the highest \(R\) value for the resulting full set of groups. In the group-average rank variation, the sample is re-allocated to the group to which it has the best rank similarity, defined as the average of the pair-wise rank similarities between it and all members of that group (or all remaining members, in the case of its original group). The process is then iterated until it converges and repeated a fair number of times from different random starting allocations to groups, as before. The choice of \(k\) can use the same SIMPROF procedure as previously.

2.6. Data

The data used here are from Collins and Williams (1982). Full details of the sampling method are given in Collins and Williams (1981). In brief, the set of samples used here were collected in April 1974 from 57 sites (numbered 1-58, site 30 was not sampled) in the Bristol Channel and Severn Estuary, UK (Fig. 1) by means of double-oblique plankton hauls. Holoplanktonic taxa were identified and counted in the samples, and abundances were converted to densities (numbers/m\(^3\)) using standard conversion factors. As in Collins and Williams (1982), data were then fourth-root transformed and inter-sample pairwise resemblances were calculated using the Bray-Curtis similarity measure.

3. Results and specific discussion

3.1. Hierarchical agglomerative clustering

Collins and Williams (1982) presented cluster analyses derived from the Bray-Curtis similarities using hierarchical agglomerative clustering with group-average sorting. The resulting dendrogram (Fig. 2A) divides the samples into 4 groups; once each of the main groups has formed it remains separate from other groups over a relatively large drop in similarity. Collins and Williams (1982) selected the four groups determined at a 55% similarity level and characterised these as true estuarine (sites 1-8, 10, 12), estuarine and marine (9, 11, 13-27, 29), euryhaline marine (28, 31, 33-35, 42-44, 47-50, 53-55) and stenohaline marine (32, 36-41, 45, 46, 51, 52, 56-58). It is not clear from the dendrogram alone whether there is any natural sequence of community change across the four main clusters (implicit in the designations true estuarine, estuarine and marine, euryhaline marine, stenohaline marine). For example, the stenohaline marine group could just as correctly have been rotated to lie between the estuarine and marine and euryhaline marine groups. In fact, there is a strong (and
more-or-less continuous) gradient of community change across the region, associated with changing
salinity levels.

More importantly, there is a degree of subjectivity in deciding the level of similarity at which
apparent divisions among groups of samples make sense. A visual inspection of the dendrogram
(Fig. 2A) suggests a sensible separation is at 55%, dividing the samples into 4 groups. An equally
justifiable decision could have been made to slice the dendrogram at the 50% level (Fig. 2A), giving 3
groups of samples. While there are methods available to decide at what level of similarity a division
of a dendrogram may be justified, it is not always the case that dividing all groups of samples at the
same level of similarity makes sense. It was to address these issues in an objective and robust
statistical manner that Clarke et al. (2008) initially developed Similarity Profiles (SIMPROF) analysis.
According to the terminology of Somerfield and Clarke (2013), the appropriate test here is Type 1
SIMPROF, and carrying out such a test at each node in the resulting dendrogram does indeed show
that the samples divide into 4 internally-homogeneous groups (Fig. 2B). It is convenient to represent
all splits down to single points, but the grey dashed lines indicate divisions where SIMPROF provides
no support for that sub-structure.

3.2. Linkage options and cophenetic correlations

This dendrogram has a cophenetic correlation of 0.797. A corresponding dendrogram constructed
using single linkage gives a correlation of 0.633, and with complete linkage a value of 0.722. Thus
the cophenetic correlations provide objective evidence for the guidance based on experience, that
group-average linkage usually provides a better view of the inter-sample relationships than analyses
based on other linkage options.

It is instructive to see how reliable this view may be. A plot of cophenetic distances (the vertical
distance to the first shared node between pairs of points; Jain and Dubes, 1988) against the
underlying dissimilarities (Fig. 3) shows that the overall pattern of dissimilarities are generally poorly
retained in the cluster analysis. Although, as indicated by the cophenetic correlation, increasing
dissimilarity tends to result in increasing distance (itself a dissimilarity on the same measurement
scale), there is also a strongly increasing trend in variability with the larger cophenetic distances
representing very wide ranges of dissimilarities in the dendrogram.

An analysis of cophenetic correlations for a range of β (Fig. 4) shows no compelling evidence for
preferring Lance and Williams’ (1967) suggestion of a value of -0.25, for these data at least. In fact,
the optimum value of β, -0.04, is very close to zero suggesting that WPMGA is an appropriate linkage
choice. The cophenetic correlation from the resulting dendrogram is 0.793, slightly less than that from UPMGA, and the analyses differ primarily in the placement of one sample, 23, from the estuarine and marine group to the euryhaline marine group. Empirical evidence suggests that an optimum flexible beta solution is usually inferior to group average linkage, perhaps as a result of the failure of WPGMA-type solutions to weight $\delta_{CA}$ and $\delta_{CB}$ appropriately under averaging, for groups A and B of very different sizes.

3.3. Binary divisive clustering (UNCTREE)

The tree diagram which results from unconstrained binary divisive clustering of the Bray-Curtis resemblances for the 57 Bristol Channel zooplankton samples is given in Fig. 5, showing the two alternative scalings (A% and B%) of the y-axis (Fig. 5A and 5B). As with the comparative agglomerative clustering (Fig. 2) it is convenient to represent all splits down to single points, but the grey dashed lines indicate divisions where SIMPROF provides no support for that sub-structure. Visual comparison of the divisive and agglomerative trees (e.g. Fig. 2B and Fig. 5B) is not particularly easy, though they have been manually rotated to aid this (a dendrogram is only defined down to arbitrary rotations of its branches, in the manner of a ‘mobile’). Clearly, however, only four groups have been identified by the SIMPROF tests in both cases. The group constitutions have much in common, though they are not identical. This is more readily seen from Fig. 6A & B, which use a non-metric MDS plot to represent the community sample relationships in 2-d ordination space. It is clear that only sites 9, 23 and 24 change groups between the two hierarchical clustering methods and these all appear at the edges of their groups in both plots, which are thus reassuringly consistent (bear in mind also that a 2-d MDS plot gives only an approximation to the true sample relationships in higher dimensions, the MDS stress of 0.11 here being low but not negligible).

3.4. k-R clustering

Fig. 6C shows the optimum grouping produced by k-R clustering, superimposed on the same MDS plot as for Figs 6A and B. The SIMPROF routine has again terminated the procedure with $k=4$ groups (a to d), which are very similar to those for the two hierarchical methods, but with the three sites 9, 23 and 24 allocated to the four groups in yet a third way. This appears to be at least as convincing an allocation as for either of the hierarchical plots (though do not lose sight of the fact that the MDS itself is only an approximation to the real inter-sample resemblances). The group-average rank variation of k-R clustering produces exactly the same four groups as seen in Fig 6C. This will not always be the case, but it should be expected that these two variations will generally give closer solutions to each other than to the hierarchical methods.
3.5. Numerical comparison of methods

As already seen, agglomerative hierarchical methods with different linkage options, which result in a dendrogram with y-axis on a (dis)similarity scale, can be compared by means of a Pearson matrix correlation between their cophenetic distances (dissimilarities) and the original dissimilarity matrix. Here, as almost without exception, this demonstrated the superiority of group average (UPGMA) over single or complete linkage strategies, and also over flexible beta, albeit more marginally. However, comparison of UPGMA with the unconstrained binary divisive (UNCTREE) clustering is less straightforward, because the latter can have somewhat arbitrary scaling on the y-axis of its tree diagram. For example, for clarity of visualising the sequence of binary divisions, it is sometimes preferable to use the equi-stepped scale (A%) for divisions on the y-axis (Fig. 5A), particularly where there are reversals of direction in branch structures when using the B% scale. This does not happen in Fig. 5B, but can occur, particularly with the constrained form of binary divisive clustering (LINKTREE), if a clear division in the community samples has no constraining “explanation” in terms of any of the recorded environmental variables (Clarke et al. 2008). This is avoided by the A% scaling but y-axis values are now no longer comparable across separate branches, being dependent only on the number of samples in that section of the tree. Contrasting Figs. 5A & B, it is clear that on the A% scale even the rank order of the divisions on the y-axis is not maintained, so a non-parametric correlation (e.g. Spearman) between cophenetic y-axis distances and original dissimilarities is not a viable answer here for head-to-head comparison of these metric agglomerative and non-metric divisive solutions.

Instead, there are two possible approaches, characterisable as “home” and “away” matches. Given that both techniques have resulted in four SIMPROF groups, a “home” match (non-metric) for the UNCTREE divisive method might compare the global ANOSIM R value computed from all four groups of zooplankton samples displayed in Fig. 6A with that in Fig. 6B (this R value is computed from the original high-d dissimilarity ranks of course, and not the 2-d nMDS ordination). Not unexpectedly, this results in a “home” win, with $R = 0.880$ for the non-metric divisive UNCTREE and $R = 0.870$ for the metric agglomerative UPGMA. The “away” match for the non-metric UNCTREE is more decisive. This is to compare the cophenetic correlation for UPGMA, previously seen to be 0.797, with that for the (rank-based) divisive tree displayed with (metric) group average dissimilarities computed at every division, as if it were an agglomerative dendrogram. In other words, the y axis value for the node at the top of each group in the divisive tree is the unweighted average of all pair-wise dissimilarities between the two sub-groups of samples joined at that node. This is certainly playing a
match on the agglomerative method’s home turf but the outcome is a marginal win for the divisive method, with a (Pearson) cophenetic correlation of 0.812 with the original (metric) dissimilarities. Can such comparisons be extended to the non-hierarchical $k$-$R$ clustering method? Clearly, any concept of cophenetic distance is impossible for a flat clustering method, with no form of sequential clustering of the individual samples. But the comparison of global ANOSIM $R$ statistics for the four (slightly differing) groups identified by all three methods is perfectly viable, albeit it must be considered the “home” match for the flat clustering method, since that should have optimised the global ANOSIM $R$ statistic among the $k$ groups which result from the sequence of SIMPROF tests. And indeed, for $k$-$R$ clustering, $R = 0.884$, in comparison to $R = 0.880$ for UNCTREE and $R = 0.870$ for UPGMA.

These numerical comparisons are therefore consistent with the earlier contention of a performance ranking, with flat $>$ divisive $>$ agglomerative, though the fact that they rank in precisely the opposite order in terms of ease of calculation, and the very marginal differences seen here and in Fig. 6, suggest that such a ranking should not be taken too seriously.

4. General discussion

In this paper we introduce some methods that use ANOSIM $R$ to determine the relative strengths of different clustering of the same dissimilarities, in order to construct group structures. There are, of course, classical equivalents of each method. For binary divisive clustering the criterion for quantifying a good binary division is clearly central. Classically (e.g. Edwards and Cavalli-Sforza, 1965), ordinary (Euclidean) distance is regarded as the relevant resemblance measure, and the total combination of sums of squares within the two groups about their group centroids, and that between the group centroids about the overall centroid (the same principles apply to one or many dimensions and two or more groups). By minimising the within-group sums of squares, that between groups is maximised, since the total sums of squares is fixed. For each group, Huygens theorem (e.g. see Anderson et al, 2008) expresses those within-group sums of squares as simply the sum of the squared Euclidean distances between every pair of points in the group, divided by that number of points. In other words, the classic criterion minimises a weighted combination of within group resemblances, defined as squared Euclidean distances. The classic flat clustering method is $k$-means clustering (MacQueen, 1967; Steinhaus, 1957), which seeks to minimise within-group sums of
squares about the $k$ group centroids (Lloyd, 1982). This is also equivalent to minimising some
weighted combination of within-group resemblances between pairs of samples, as measured by a
squared Euclidean distance coefficient. Thus the classical methods are really only appropriate in
situations where (squared) Euclidean distance is an appropriate measure of resemblance, such as
analyses of normalised environmental variables. For community analyses, appropriate measures of
resemblance such as those in the Bray-Curtis family (Clarke et al., 2006) are required, and
partitioning sums of squares is no longer a possibility. The new methods proposed here, UNCTREE
and $k$-R clustering, address the problem by maximising ANOSIM $R$, which measures (non-
parametrically) the degree of overall separation of the groups, formed from the ranks in the full
resemblance matrix. Thus they are appropriate for any data type and resemblance measure.

A ‘take-home’ message from Fig. 6 is that clustering rarely escapes a degree of arbitrariness: the
data simply may not represent clearly separated clusters. For the Bristol Channel sites, where there
certainly are plausible groups but within a more or less continuous gradation of change in plankton
communities (strongly correlated with increased salinity of the sites, Fig. 6B), different methods
must be expected to divide this continuum up in slightly different ways. It is important to remember
that, as applied here, SIMPROF tests groupings that are determined by the clustering algorithm and,
under different clustering techniques, samples can move from one group to another, both deemed
homogeneous by SIMPROF, so neither should be regarded as absolute. In spite of the above
evidence that the top-down methods can lead to marginally better solutions (notwithstanding the
vagaries of optimal search methods), use of a specific grouping from a bottom-up, group-average
hierarchy should probably be viewed operationally as on a par with that from a divisive hierarchy or
from the non-hierarchical $k$-R clustering, in either form. And, certainly, the group average (UPGMA)
approach appears uniformly superior to other linkage options in agglomerative clustering. It is
notable here that SIMPROF supports four very similar groups for all three methods shown in Fig. 6,
though this degree of consistency is not guaranteed for analyses of other sets of data. In fact,
especially in cases where a low-dimensional MDS plot is not at all reliable because of high stress, the
plurality of clustering methods may provide insight into the robustness of conclusions that can be
drawn about group structures from the (high-dimensional) resemblance matrix. Such comparisons of
differing clustering methods need to ‘start from the same place’, namely using the same
resemblance matrix, otherwise an inferred lack of a stable group structure could be due to the
differing assumptions being made about how the (dis)similarity between two samples is defined (e.g.
Bray-Curtis vs squared Euclidean distance). This is also a point to bear in mind in comparing
ordination methods: a primary difference between them is often not the way they choose to
represent high-dimensional information in lower dimensional space but how they define that higher-
dimensional information differently, in their choice of explicit or implicit resemblance measure (Clarke et al., 2006). Comparison of results from clustering methods that utilise axis scores from ordination methods (e.g. Hill, 1979; Lefkovitch, 1976; Noy-Meir, 1973) will be particularly influenced by this issue. Given that the appropriate choice of resemblance is fundamental to a successful analysis, many existing methods which use inappropriate measures, either implicitly or explicitly, are to be avoided, or at best treated with caution.

It is often recommended that cluster analysis is best used in conjunction with ordination. Indeed, that has been done here (Fig. 6) by labelling samples in the ordination plot with their group membership from cluster analyses, a simple technique to apply in either 2- or 3-d ordination space. Widely used alternatives are to draw smoothed convex hulls (contours) on a 2-d ordination, indicating the samples grouped together at different levels in the dendrogram (nested contours), or groups of samples which are not separated by SIMPROF, and so on. The implication is that both clustering and ordination provide equally valid views of the relationships among samples, so if the two methods being used in conjunction support each other then the investigator is in a position to trust both. A novel alternative view of the relationship between a cluster analysis and an ordination (Fig. 7) shows graphically the complex interrelationships between the linking of samples in the former and the placement of samples in the latter. It is readily apparent that the beguiling simplicity of a dendrogram may give a highly distorted view of true distances among objects.

In the statistical analytical framework under discussion the ordination method of choice is often nMDS, which arrives at its solution through an iterative procedure which is not guaranteed to achieve the optimal (global) solution with minimum stress. It is normal to rerun the procedure many times and to check numerically (stress values, a measure which reflects lack of agreement in the rank order of the distances among points in the ordination and the rank order of points in the resemblance matrix) and graphically (Shepard diagrams, plots of distances in the solution against original resemblances) before accepting that a nMDS plot is an accurate reflection of the true multivariate information in the underlying resemblance matrix. It is interesting that, to date, such rigour has rarely been applied to clustering. In part this reflects the numerical simplicity of widely used methods, in that there is only one solution. What is rarely questioned is the adequacy of that solution. Here we show how the adequacy of a hierarchical agglomerative clustering may be checked numerically (cophenetic correlation) and graphically (plotting cophenetic distances against dissimilarities, Fig. 3). It may surprise some to see how poor a representation of the true relationships a dendrogram or tree diagram may be, and perhaps consideration should be given to more appropriate methods. This may be particularly relevant in some areas of science where trees
are routinely used to represent relationships, such as in phylogenetics and microbial studies, especially under conditions in which a pure hierarchical structure is not mandated (e.g. in genetics, because of possibilities of recombination etc.), or in the derivation of hierarchical groupings of taxa based on taxonomic information or traits for the calculation of certain indices of relatedness which are defined in terms of branch lengths in a tree (see Somerfield et al., 2008; Warwick and Clarke, 2001).

The terms clustering and classification were originally synonymous and that is the way they still tend to be used by ecologists, but in statistical language the methods considered here are all clustering techniques. The term classification is now usually reserved for classifying unknown new samples into known prior group structures. Nearly half a century ago, Cormack (1971) warned against the indiscriminate use of cluster analysis, saying “availability of … [clustering] techniques has led to the waste of more valuable scientific time than any other ‘statistical’ innovation”. The ever larger number of clustering techniques and their increasing accessibility on modern computer systems makes this warning no less pertinent today. With that in mind it may seem rather unnecessary to consider adding to the clustering techniques available, but the thrust of this paper is to demonstrate how useful it can be to compare a range of different, but compatible, techniques when assessing the robustness of a clustering solution. And, in order to facilitate that, we have introduced two new methods: a divisive hierarchical and a non-hierarchical (k-means type) method, the latter in two variants. These are now entirely compatible with a range of other widely-employed non-parametric multivariate methods (e.g. nMDS ordination, ANOSIM tests, BEST or LINKTREE approaches to linking community structure to environmental variables; Clarke, 1993; Clarke et al., 2008, 2014) because they all start from exactly the same information, the rank orders of the entries in a defined resemblance matrix. As a result, multiple clustering methods in conjunction with ordination have, for the Bristol Channel zooplankton analyses, both directed attention to those few sites which are not firmly allocated to one of the groupings, and instilled a greater degree of confidence in the robustness of the main groupings produced.

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References


Figure legends

Fig 1. Map of the Bristol Channel and Severn Estuary showing the locations of the 57 sites from which zooplankton samples were collected in April 1974.

Fig 2. Dendrogram from hierarchical agglomerative clustering of the 57 sites using group average linking of Bray-Curtis similarities calculated from √√ transformed holoplankton densities showing: A) the three groups produced by applying an arbitrary cut at 50% similarity; B) the results of successive SIMPROF tests on groups of samples defined by nodes of the tree, beginning at the top. Only the first three tests showed significant multivariate structure in the samples below that point, so there is no evidence from SIMPROF that the detailed clustering structure (grey dashed lines) within each of the 4 main groups is interpretable.

Fig 3. Variation in the cophenetic correlation between the original resemblances from the zooplankton data and cophenetic distances from cluster analyses using flexible beta linking with different values of β. The highest correlation (0.794) is from the analysis in which β = -0.04.

Fig 4. Relationship of the original Bray-Curtis dissimilarities between pairs of samples and cophenetic distances between those same pairs of samples in the dendrogram shown in Fig 2.

Fig 5. Dendrogram from unconstrained binary divisive clustering of 57 sites maximising ANOSIM R at each binary split (UNCTREE) plotted using: A) the A% scale in which steps are arbitrarily equi-spaced; B) the B% scale which indicates the relative strength of each division. As in Fig 2B continuous lines indicate structure supported by SIMPROF and grey dashed lines indicate no interpretable structure.

Fig 6. Non-metric MDS ordination of the 57 sites derived from Bray-Curtis similarities calculated from √√ transformed holoplankton densities. Symbols indicate the groups found by SIMPROF tests for each of 3 clustering methods: A) agglomerative hierarchical with group-average linking; B) hierarchical divisive clustering; C) non-hierarchical k-R clustering. Labels are site numbers (A, C) or salinity scores (B, from 1: <26.3, ..., 9: >35.1 ppt, see Clarke et al., 2014 for full details).

Fig 7. 2-d nMDS ordination presented in a ‘3-d project’ with the dendrogram from group-average linking. Contours indicate samples joined with a similarity of 55%. Sample symbols denote SIMPROF groups.
Figure 2

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A

Bray-Curtis similarity

B

Bray-Curtis similarity

= SIMPROF test carried out

= significant structure within