tclust: An R Package for a Trimming Approach to Cluster Analysis

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Abstract

This introduction to the R package tclust is a (slightly) modified version of Fritz et al. (2012), published in the Journal of Statistical Software.

Outlying data can heavily influence standard clustering methods. At the same time, clustering principles can be useful when robustifying statistical procedures. These two reasons motivate the development of feasible robust model-based clustering approaches. With this in mind, an R package for performing non-hierarchical robust clustering, called tclust, is presented here. Instead of trying to “fit” noisy data, a proportion \( \alpha \) of the most outlying observations is trimmed. The tclust package efficiently handles different cluster scatter constraints. Graphical exploratory tools are also provided to help the user make sensible choices for the trimming proportion as well as the number of clusters to search for.

Keywords: Model-based clustering, trimming, heterogeneous clusters.

1. Introduction to robust clustering and tclust

Methods for cluster analysis attempt to detect homogeneous clusters with large heterogeneity among them. As happens with other (non-robust) statistical procedures, clustering methods may be heavily influenced by even a small fraction of outlying data. For instance, two or more clusters might be joined artificially, due to outlying observations, or “spurious” non-informative clusters may be composed of only a few outlying observations (see, e.g., García-Escudero and Gordaliza 1999; García-Escudero et al. 2010). Therefore, the application of robust methods in this context is very advisable, especially in fully automatic clustering (unsupervised learning) problems. Certain relations between cluster analysis and robust methods (Rocke and Woodruff 2002; Hardin and Rocke 2004; García-Escudero et al. 2003; Woodruff and Reiners 2004) also motivate interest in robust clustering techniques. For example, robust clustering techniques can be used to handle “clusters” of highly concentrated outliers which are especially dangerous in (non-robust) estimation. García-Escudero et al. (2010) provides a recent survey of robust clustering methods.

The tclust package for the R environment for statistical computing (R Development Core Team 2010) implements different robust non-hierarchical clustering algorithms where trimming plays a key role. This package is available at http://CRAN.R-project.org/package=tclust.
When trimming allows the removal of a fraction $\alpha$ of the “most outlying” data, the strong influence of outlying observations can be avoided. This trimming approach to clustering has been introduced in Cuesta-Albertos et al. (1997), Gallegos (2002), Gallegos and Ritter (2005) and García-Escudero et al. (2008). Trimming also serves to identify potentially interesting anomalous observations.

Trimming is not a new concept in statistics. For instance, the trimmed mean for one-dimensional data removes a proportion $\alpha/2$ each of the largest and smallest observations before computing the mean. However, it is not straightforward to extend this philosophy to cluster analysis, because most of these problems are of multivariate nature. Moreover, it is often the case that “bridge points” lying between clusters ought to be trimmed. Instead of forcing the statistician to define the regions to be trimmed in advance, the procedures implemented in `tclust` take the whole data structure into account in order to decide which parts of the sample should be discarded. By considering this type of trimming, these procedures are even able to trim outlying bridge points. The “self-trimming” philosophy behind these procedures is exactly the same as adopted by some well-known high breakdown-point methods (see, e.g., Rousseeuw and Leroy 1987).

As a first example of this trimming approach, let us consider the trimmed $k$-means method introduced in Cuesta-Albertos et al. (1997). The function `tkmeans` from the `tclust` package implements this method. In the following example, this function is applied to a bivariate data set based on the Old Faithful geyser called `geyser2` that accompanies the `tclust` package. The code given below creates Figure 1.

```r
R > library("tclust")
R > data("geyser2")
R > clus <- tkmeans (geyser2, k = 3, alpha = 0.03)
R > plot (clus)
```

In the data set `geyser2`, we are searching for $k = 3$ clusters and a proportion $\alpha = 0.03$ of the data is trimmed. The clustering results are shown in Figure 1. Among this 3% of trimmed data, we can see 6 anomalous “short followed by short” eruptions lengths. Notice that an observation situated between the clusters is also trimmed.

The package presented here adopts a “crisp” clustering approach, meaning that each observation is either trimmed or fully assigned to a cluster. In comparison, mixture approaches estimate a cluster pertinence probability for each observation. Robust mixture alternatives have also been proposed where noisy data is tried to be fitted through additional mixture components. For instance, package `mclust` (Fraley and Raftery 2012; Banfield and Raftery 1993; Fraley and Raftery 1998) and the Fortran program `emmix` (McLachlan 1999; McLachlan and Peel 2000) implement such robust mixture fitting approaches. Mixture fitting results can be easily converted into a “crisp” clustering result by converting the cluster pertinence probabilities into 0-1 probabilities. Contrary to these mixture fitting approaches, the procedures implemented in the `tclust` package simply remove outlying observations and do not intend to fit them at all. Package `tlemix` (see Neytchev et al. 2012; Neykov et al. 2007) also implements a closely related trimming approach. As described in Section 3, the `tclust` package focuses on offering adequate cluster scatter matrix constraints leading to a wide range of clustering procedures depending on the chosen constraint, and avoiding the occurrence of spurious non-interesting clusters.
The outline of the paper is as follows: In Section 2 we briefly review the so-called “spurious outliers” model and show how to derive two different clustering criteria from it. Different constraints on the cluster scatter matrices and their implementation in the `tclust` package are addressed in Section 3. Section 4 presents the numerical output returned by this package. Section 5 provides some brief comments concerning the algorithms implemented, and a comparison of `tclust` and several other robust clustering approaches are given in Section 6. Section 7 shows some graphical outputs that help advise the choice of the number of clusters and the trimming proportion. Other useful plots summarizing the robust clustering results are shown in Section 8. Finally, Section 9 applies the `tclust` package to a well-know real data set.
2. Trimming and the spurious outliers model

Gallegos (2002) and Gallegos and Ritter (2005) propose the “spurious outliers model” as a probabilistic framework for robust crisp clustering. Let \( f(\cdot; \mu, \Sigma) \) denote the probability density function of the \( p \)-variate normal distribution with mean \( \mu \) and covariance matrix \( \Sigma \). The “spurious-outlier model” is defined through “likelihoods” like

\[
\prod_{j=1}^{k} \prod_{i \in R_j} f(x_i; \mu_j, \Sigma_j) \prod_{i \in R_0} g_i(x_i)
\]

(1)

with \( \{R_0, ..., R_k\} \) being a partition of the set of indices \( \{1, 2, ..., n\} \) such that \#\( R_0 = \lceil n\alpha \rceil \). \( R_0 \) are the indices of the “non-regular” observations generated by other (not necessarily normal) probability density functions \( g_i \). “Non-regular” observations can be clearly considered as “outliers” if we assume certain sensible assumptions for the \( g_i \) (see details in Gallegos 2002; Gallegos and Ritter 2005). Under these assumptions, the search of a partition \( \{R_0, ..., R_k\} \) with \#\( R_0 = \lceil n\alpha \rceil \), vectors \( \mu_j \) and positive definite matrices \( \Sigma_j \) maximizing (1) can be simplified to the same search (of a partition, vectors and positive definite matrices) by just maximizing

\[
\sum_{j=1}^{k} \sum_{i \in R_j} \log f(x_i; \mu_j, \Sigma_j).
\]

(2)

Notice that observations \( x_i \) with \( i \in R_0 \) are not taken into account in (2). Maximizing (2) with \( k = 1 \) yields the Minimum Covariance Determinant (MCD) estimator (Rousseeuw 1985). Unfortunately, the direct maximization of (2) is not a well-defined problem when \( k > 1 \). It is easy to see that (2) is unbounded without any constraint on the cluster scatter matrices \( \Sigma_j \). The \texttt{tclust} function from the \texttt{tclust} package approximately maximizes (2) under different cluster scatter matrix constraints which will be shown in Section 3.

The maximization of (2) implicitly assumes equal cluster weights. In other words, we are ideally searching for clusters with equal sizes. The function \texttt{tclust} provides this option by setting the argument \texttt{equal.weights = TRUE}. The use of this option does not guarantee that all resulting clusters exactly contain the same number of observations, but the method hence prefers this type of solutions.

Alternatively, different cluster sizes or cluster weights can be considered by searching for a partition \( \{R_0, ..., R_k\} \) (with \#\( R_0 = \lceil n\alpha \rceil \)), vectors \( \mu_j \), positive definite matrices \( \Sigma_j \) and weights \( \pi_j \in [0, 1] \) maximizing

\[
\sum_{j=1}^{k} \sum_{i \in R_j} (\log \pi_j + \log f(x_i; \mu_j, \Sigma_j)).
\]

(3)

The (default) option \texttt{equal.weights = FALSE} is used in this case. Again, the scatter matrices also have to be constrained such that the maximization of (3) becomes a well-defined problem. Note that (3) simplifies to (2) when assuming \texttt{equal.weights = TRUE} and all weights are equally set to \( \pi_j = 1/k \).
equal.weights = TRUE | equal.weights = FALSE
--- | ---
restr = "eigen" | $k$-means Cuesta-Albertos et al. (1997) García-Escudero et al. (2008)
restr = "deter" | Gallegos (2002) This work
restr = "sigma" | Friedman and Rubin (1967) Gallegos and Ritter (2005) This work

Table 1: Clustering methods handled by tclust. Names in cursive letters are untrimmed ($\alpha = 0$) methods.

3. Constraints on the cluster scatter matrices

As already mentioned, the function tclust implements different algorithms aimed at approximately maximizing (2) and (3) under different types of constraints which can be applied on the scatter matrices $\Sigma_j$. The type of constraint is specified by the argument restr of the tclust function. Table 1 gives an overview of the different clustering approaches implemented by the tclust function depending on the chosen type of constraint.

Imposing constraints is compulsory because maximizing (2) or (3) without any restriction is not a well-defined problem. Notice that an almost degenerated scatter matrix $\Sigma_j$ would cause trimmed log-likelihoods (2) and (3) to tend to infinity. This issue can cause a robust clustering algorithm of this type to end up finding “spurious” clusters almost lying in lower-dimensional subspaces. Moreover, the resulting clustering solutions might heavily depend on the chosen constraint. The strength of the constraint is controlled by the argument restr.fact $\geq 1$ in the tclust function. The larger restr.fact is chosen, the looser is the restriction on the scatter matrices, allowing for more heterogeneity among the clusters. On the contrary, small values of restr.fact close to 1 imply very “equally scattered” clusters. This idea of constraining cluster scatters to avoid spurious solutions goes back to Hathaway (1985), who proposed it in mixture fitting problems.

Also arising from the spurious outlier model, other types of constraints have recently been introduced by Gallegos and Ritter (2009, 2010). These (closely related) constraints also serve to avoid degeneracy of trimmed likelihoods but they are not implemented in the current version of the tclust package.

3.1. Constraints on the eigenvalues

Based on the eigenvalues of the cluster scatter matrices, a scatter similarity constraint may be defined. With $\lambda_i(\Sigma_j)$ as the eigenvalues of the cluster scatter matrices $\Sigma_j$ and

$$M_n = \max_{j=1,\ldots,k} \max_{l=1,\ldots,p} \lambda_l(\Sigma_j)$$
$$m_n = \min_{j=1,\ldots,k} \min_{l=1,\ldots,p} \lambda_l(\Sigma_j)$$

as the maximum and minimum eigenvalues, the restriction restr = "eigen" constrains the ratio $M_n/m_n$ to be smaller or equal than a fixed value restr.fact $\geq 1$. A theoretical study of the properties of this approach with equal.weights = FALSE can be found in García-Escudero et al. (2008).

This type of constraint limits the relative size of the axes of the equidensity ellipsoids defined through the obtained $\Sigma_j$ when assuming normality. This way we are simultaneously controlling the relative group sizes and also the deviation from sphericity in each cluster.
Setting `equal.weights = TRUE`, `restr = "eigen"` and `restr.fact = 1` implies the most constrained case. In this case, the `tclust` function tries to solve the trimmed $k$-means problem as introduced by Cuesta-Albertos et al. (1997). This problem simplifies to the well-known $k$-means clustering criterion when no trimming is done (i.e., $\alpha = 0$). The `tkmeans` function directly implements this most constrained application of the `tclust` function.

### 3.2. Constraints on the determinants

Another way of restricting cluster scatter matrices is constraining their determinants. Thus, if

$$M_n = \max_{j=1, \ldots, k} |\Sigma_j| \quad \text{and} \quad m_n = \min_{j=1, \ldots, k} |\Sigma_j|$$

are the maximum and minimum determinants, we attempt to maximize (2) or (3) by constraining the ratio $M_n/m_n$ to be smaller or equal than a fixed value `restr.fact`. This is done in the function `tclust` by using the option `restr = "deter"`.

Now, this type of constraint limits the relative volumes of the mentioned equidensity ellipsoids, but not the cluster shapes. The use of this type of constraint is particularly advisable when affine equivariance is required because this property is satisfied when `restr = "deter"`.

The untrimmed case $\alpha = 0$, `restr = "deter"` and `restr.fact = 1` was already outlined in Maronna and Jacovkis (1974), as the only sensible way to avoid (Mahalanobis distance modified) $k$-means type algorithms to return clusters of a few almost collinear observations. The possibility of trimming data was also considered in Gallegos (2002) who implicitly assumed $|\Sigma_1| = \ldots = |\Sigma_k|$ (and so `restr.fact = 1`). The package presented here extends her approach to more general cases (`restr.fact > 1`).

### 3.3. Equal scatter matrices

Among the methods considered, `tclust` also implements a stronger type of constraint by setting `restr = "sigma"` which forces all cluster scatter matrices to be the same: $\Sigma_1 = \ldots = \Sigma_k$. This is known as the “determinantal” criterion and it goes back to Friedman and Rubin (1967). The trimmed version of this approach was introduced by Gallegos and Ritter (2005). The argument `restr.fact` is ignored when applying this type of constraint.

### 3.4. Example

In this example, we examine the influence of different constraints by applying the function `tclust` to the so-called `M5data` data set. This data set, which accompanies the `tclust` package, has been generated following the simulation scheme M5 introduced in García-Escudero et al. (2008). Thus it is a bivariate mixture of three simulated gaussian components with very different scatters and a clear overlap between two of these components. A 10% proportion of outliers is also added in the outer region of the bounding rectangle enclosing the three gaussian components. See Figure 2 for a graphical representation and García-Escudero et al. (2008) for more details on the structure of this `M5data` data set. Executing the following code yields Figure 3.

```r
R > data("M5data")
R > x <- M5data[, 1:2]
```
Figure 2: A scatter plot of the M5data data set. Different symbols are used for the data points generated by each of the three bivariate normal components and “◦” for the added outliers.

R > res.a <- tclust (x, k = 3, alpha = 0.1, restr.fact = 1, restr = "eigen", + equal.weights = TRUE)
R > res.b <- tclust (x, k = 3, alpha = 0.1, restr.fact = 1, restr = "sigma", + equal.weights = TRUE)
R > res.c <- tclust (x, k = 3, alpha = 0.1, restr.fact = 1, restr = "deter", + equal.weights = TRUE)
R > res.d <- tclust (x, k = 3, alpha = 0.1, restr.fact = 50, restr = "eigen", + equal.weights = FALSE)
R > plot (res.a, main = "/r")
R > plot (res.b, main = "/r")
R > plot (res.c, main = "/r")
R > plot (res.d, main = "/r")
Figure 3: Results of the clustering processes for the M5data data set for different constraints on the cluster scatter matrices and the parameters $\alpha = 0.1$ and $k = 3$. Different colors and symbols represent each observation’s individual cluster assignment.

Although different constraints are imposed, we are searching for $k = 3$ clusters and the trimming proportion is set to $\alpha = 0.1$ in all the cases. Note that only the clustering procedure introduced in García-Escudero et al. (2008), shown in Figure 3,(d), with a sufficiently large value of `restr.fact` approximately returns the three original clusters in spite of the very different cluster scatters and different cluster sizes. Moreover, this clustering procedure adequately handles the severe overlap of two clusters. The value `restr.fact = 50` has been chosen in this case because the eigenvectors of the covariance matrices of the three gaussian components satisfy restriction (4) for this value. Due to their underlying assumptions, the other three clustering methods (trimmed $k$-means in Figure 3,(a), Gallegos and Ritter (2005) in (b), Gallegos (2002) in (c)) return rather similarly structured clusters. In fact, we found spherical clusters in (a), clusters with the same scatter matrix in (b) and clusters with the same cluster scatter matrix determinant in (c). The M5data is perhaps a very “extreme” sit-
Figure 4: Applying tclust with $k = 3$ and $\alpha = 0$ on a simulated data set which originally consists of 2 clusters when equal.weights = FALSE.

4. Numerical output

The function tclust returns an S3 object containing the cluster centers $\mu_j$ by columns ($\text{centers}$), scatter matrices $\Sigma_j$ as an array ($\text{cov}$), the weights ($\text{weights}$), the number of observations in each group ($\text{size}$) and the maximum value found for the trimmed log-likelihood objective function (2) or (3) ($\text{obj}$). The vector $\text{cluster}$ provides the cluster assignment of each observation, whereas an artificial cluster “0” (without location and scatter information) is introduced which holds all trimmed data points.

Sometimes, (2) and (3) maximize with some clusters remaining empty (see Figure 4). In this
case, only information on the non-empty groups is returned. Notice that, if we are searching for \( k \) clusters, empty clusters can be found when a clustering solution for a number of clusters strictly smaller than \( k \) attains a higher value for (2) or (3) than the best solution found with \( k \) clusters. In this case, artificial empty clusters may be defined by considering sufficiently remote centers \( \mu_j \) and scatter matrices \( \Sigma_j \) satisfying the specific constraints that are assigned to these empty clusters. They are chosen such that \( f(\cdot; \mu_j, \Sigma_j) \) gives almost null density to all the observations in the sample. These artificially added centers and scatter matrices are not returned as output by the \texttt{tclust} function and a warning is issued. For instance, let us consider the following code

R > set.seed (10)  
R > x <- rbind (rmvnorm (200, c (0, 0), diag (2)),  
R + rmvnorm (200, c (5, 0), diag (2)))  
R > clus <- tclust (x, k = 3, alpha = 0, restr.fact = 1)  
Warning messages:  
1: In .tclust.warn(warnings, ret) :  
  1 empty cluster has been detected - try reducing k.  
...  
R > plot (clus)

Although we are searching for \( k = 3 \) clusters, Figure 4 and the issued warning show that only 2 clusters are found. Notice that \( k = 2 \) is surely a more sensible choice for the number of clusters than \( k = 3 \) for this generated data set. Therefore, the detection of empty clusters, or clusters with few data points, can be helpful, providing valuable tools for making sensible choices for \( k \) as we will see in Section 7. On the other hand, the detection of empty clusters is very unlikely to happen when the argument \texttt{equal.weights = TRUE} is provided in the call to \texttt{tclust}.

5. Algorithms

The maximization of (2) or (3) considering different cluster scatter matrix constraints is not straightforward because of the combinatorial nature of the associated maximization problems. The algorithm presented in García-Escudero \textit{et al.} (2008) can be adapted to approximately solve all these problems. The methods implemented in \texttt{tclust} could be seen as Classification EM algorithms (Schroeder 1976; Celeux and Govaert 1992), whereas a certain type of “concentration” steps (see the fast-MCD algorithm in Rousseeuw and Van Driessen 1999) is also applied. In fact, the concentration steps applied by the package \texttt{tclust} can be considered as an extension of those applied by the batch-mode \( k \)-means algorithm (Steinhaus 1956; Forgy 1965). It can be seen that the target function always increases throughout the application of concentration steps, whereas several random start configurations are needed in order to avoid ending trapped in local maxima. Therefore, \texttt{nstart} random initializations and \texttt{iter.max} concentration steps are considered. The probability that the algorithm converges close to the global optimum maximizing (2) or (3) increases with larger values of \texttt{nstart} and \texttt{iter.max}. The drawback of high values of \texttt{nstart} and \texttt{iter.max} obviously is the increasing computational effort.

In the concentration step, the centers and scatter matrices are updated by considering the cluster sample means and cluster sample covariance matrices. New cluster assignments are
Figure 5: PAM’s clustering results for the geyser2 data with 3-medoids denoted by the symbol “×” in (a). PAM’s clustering results for a modified geyser2 data set in (b) and when applying tkmeans with $k = 3$ and $\alpha = 0.03$ in (c).

obtained by gathering the “closest” observations to the new centers. Mahalanobis distances, based on the computed cluster sample covariance matrices, are used in order to decide which are the closest observations to each center. If needed, in the updating step, the cluster sample covariance matrices are modified as little as possible but in such a way that they satisfy the desired constraints (García-Escudero et al. 2008). The main idea behind these “constrained” concentration steps is to replace the eigenvalues of the sample covariance matrices by optimally truncated eigenvalues, which satisfy the desired constraint. A more detailed description of the algorithm applied by tclust and the way the restrictions are forced onto the cluster scatter matrices can be found in Fritz et al. (2011).

6. Comparison with other robust clustering proposals

In this section, we briefly compare the performance of the clustering procedures implemented in the tclust package with respect to other robust clustering proposals in the literature.

The Partitioning Around Medoids (PAM) clustering method (Kaufman and Rousseeuw 1990) has been proposed as a robust alternative to k-means clustering. It can be seen that the effect of the 6 anomalous “short followed by short” eruptions lengths in the lower left corner of Figure 1 do not affect the position of the $k$-medoid centers (see Figure 5,(a)) too much, nor the resulting clusters. However, in Figure 5,(b), we see that the clustering results with $k = 3$ are strongly affected when moving these 6 anomalous points toward a more distant position. On the other hand, in Figure 5,(c), we can see that these outlying data points do not affect the trimmed $k$-means based clustering at all once that they are trimmed.

In fact, only one single outlier placed in a very remote position is able to completely break down the PAM method (García-Escudero and Gordaliza 1999). This also happens when applying emmix, which has a breakdown point of zero (Hennig 2004). The emmix approach is able to obtain appropriate clustering results for the two data sets made of mixtures of
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Figure 6: A data set made up of three elliptical $t$ components is shown in (a) and three asymmetrical $t$ components in (d). The associated clustering results when applying the tclust function with $k = 3$ and $\alpha = 0$ are shown in (b) and (e) and with $k = 3$ and $\alpha = 0.05$ in (c) and (f).

symmetric and asymmetric $t$ components as those shown in Figure 6. These two data sets contain three main clusters with some distant observations in the heavy tails of these $t$ components, which would be considered as outliers when assuming normality. When applying the classification EM algorithm without trimming to these data sets, we are not able to find the three cluster structures and two main clusters are artificially joined together. However, when considering $\alpha = 0.05$, tclust perfectly avoids the harmful effect of the observations in the tails and still discovers the three clusters. In fact, almost all non-trimmed observations are correctly clustered in both, symmetric and asymmetric, cases. Any $\alpha > 0$ discarding the most outlying observations would give similar results. Moreover, it may be seen that the shape of the elliptical clusters are essentially discovered in the case of the symmetric-elliptical $t$ components. In this example, we see that applying tclust to data sets including non-normally distributed components as those in Figure 6 may result in proper clustering solutions, this however cannot be guaranteed if the underlying distributions differ too much from normality. The closely related tlemix package allows to consider other non-normal models by taking advantage of the flexibility provided by the FlexMix package (Leisch 2004). On the other hand, tclust focuses on normally distributed components and on the implementation of appropriate cluster scatter matrix constraints while tlemix does not. The tlemix mainly controls the minimum number of observations in a cluster.

The widely used mclust package considers a uniformly distributed component for explaining outlying data points. As we can see, this uniform component successfully accommodates the 10% “background noise” as seen in Figure 7,(a). However, it is not able to cope with
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Figure 7: Clustering results for two simulated data sets when applying \texttt{mclust} with $k = 2$ in (a) and (c) and \texttt{tclust} with $k = 2$ and $\alpha = 0.1$ in (b) and (d).

A more structured noise pattern like the “helix” in Figure 7, (c) which also accounts for 10% of the data, although the information of a 10% contamination level was passed to \texttt{mclust}. Alternatively, the \texttt{tclust} package with $k = 2$ and $\alpha = 0.1$ properly discovers the outlying data points without trying to fit them.

Since groups of outliers may be considered as further clusters, it could be argued that robust clustering problems can always be solved by increasing the number of groups we are searching for. However, as explained in García-Escudero et al. (2010), this is not necessarily the best strategy. Firstly, sometimes the researcher fixes the number of clusters in advance, not being aware of the presence of a small amount of outlying observations. Secondly, it could lead to a quite large number of clusters when very scattered outliers are present in the data set.

A clear limitation of \texttt{tclust} is that it is not applicable on high-dimensional data sets, as the method in its current definition definitely needs a data set containing more observations than dimensions.
7. Selecting the number of groups and the trimming size

Perhaps one of the most complex problems when applying cluster analysis is the choice of the number of clusters, \( k \). In some cases one might have an idea of the number of clusters in advance, but usually \( k \) is completely unknown. Moreover, in the approach proposed here, the trimming proportion \( \alpha \) has also to be chosen without knowing the true contamination level.

As we will see through the following example, the choices for \( k \) and \( \alpha \) are related problems that should be addressed simultaneously. It is important to see that a particular trimming level implies a specific number of clusters and vice versa. This dependency can be explained as entire clusters tend to be trimmed completely when increasing \( \alpha \). On the other hand, when choosing \( \alpha \) too low, groups of outliers might form new spurious clusters and thus it appears that the number of clusters found in the data set is higher. Moreover, the simultaneous choice of \( k \) and \( \alpha \) depends on the type of clusters we are searching for and on the allowed differences between cluster sizes. These two aspects can be controlled by the choice of arguments `restr` and `restr.fact`.

To demonstrate the relation between \( \alpha \), \( k \) and `restr.fact`, let us consider `restr = "eigen"` and the data set in Figure 8 which could either be interpreted as a mixture of three components (a) or a mixture of two components (b) with a 10% outlier proportion. Both clustering solutions shown in Figure 8 are perfectly sensible and the final choice of \( \alpha \) and \( k \) only depends on the value given to `restr.fact`. The code used to obtain Figure 8 is the following:

```R
R > sigma1 <- diag (2) ## EigenValues: 1, 1
R > sigma2 <- diag (2) * 8 - 2 ## EigenValues: 8, 4
R > sigma3 <- diag (2) * 50 ## EigenValues: 50, 50
R > mixt <- rbind (
  + rmvnorm (360, mean = c (0.0, 0), sigma = sigma1),
  + rmvnorm (540, mean = c (5.0, 10), sigma = sigma2),
  + rmvnorm (100, mean = c (2.5, 5), sigma = sigma3))
R > plot (tclust (mixt, k = 3, alpha = 0.00, restr.fact = 50))
R > plot (tclust (mixt, k = 2, alpha = 0.05, restr.fact = 12))
```

Considering `sigma1` and `sigma2`, the quotient of the largest and smallest eigenvalue is 8, whereas the maximal quotient is 50 if we consider `sigma1`, `sigma2` and `sigma3`. Thus `restr.fact = 8` would allow to consider two clusters while `restr.fact = 50` would also allow to assume three groups there. Although the proportion of “contaminated” data is equal to 10%, the trimming level must be reduced to 5%, when considering \( k = 2 \), because the third (more scattered) gaussian component partially overlaps with the other two components.

Let us assume first that `restr` and `restr.fact` have been fixed in advance by the researcher who applies the robust clustering method. Even with this information and assuming \( \alpha = 0 \), choosing the appropriate number of clusters is not an easy task. The careful monitoring of the maximum value attained by log-likelihoods like those in (2) and (3) while changing \( k \) has traditionally been applied as a method for choosing the number of clusters when \( \alpha = 0 \). Moreover Bryant (1991) stated that the use of “weighted” log-likelihoods (3) is preferred to the use of log-likelihoods assuming equal weights (2). Notice that increasing \( k \) always causes the maximized log-likelihood (2) to increase too, and this could lead to “overestimate” the appropriate number of clusters (see García-Escudero et al. 2011).
In this trimming framework, let us consider $L^\Pi_{\text{restr.fact}}(\alpha, k)$ as the maximum value reached by (3) for each combination of a given set of values for $k$ and $\alpha$. García-Escudero et al. (2011) propose to monitor the “classification trimmed likelihoods” functionals

$$(\alpha, k) \mapsto L^\Pi_{\text{restr.fact}}(\alpha, k)$$

while altering $\alpha$ and $k$, which yields an exploratory graphical tool for making sensible choices for parameters $\alpha$ and $k$. In fact, it is proposed to choose the number of clusters as the smallest value of $k$ such that

$$L^\Pi_{\text{restr.fact}}(\alpha, k + 1) - L^\Pi_{\text{restr.fact}}(\alpha, k)$$

is (close to) 0 except for small values of $\alpha$. Once the number of clusters is fixed, a good choice for the trimming level is the first $\alpha_0$ such that (5) is (close to) 0 for every $\alpha \geq \alpha_0$. Although we are convinced that monitoring the classification trimmed likelihoods functionals is very informative, no theoretical statistical procedures are available yet for determining when (5) can be formally considered as “close to 0”.

The function ctlcurves in package tclust approximates the classification trimmed likelihoods by successively applying the tclust function for a sequence of values of $k$ and $\alpha$. A default value restr.fact = 50 is considered but, if desired, other values of restr.fact can be passed to tclust via ctlcurves too.

For instance, the following code applied to the previously simulated mixt data set

```
R > plot (ctlcurves (mixt, k = 1:4, alpha = seq (0, 0.2, by = 0.05)))
```

results in Figure 9. This figure shows that increasing $k$ from 2 to 3 is needed when $\alpha = 0$, as the objective functions value differs noticeably between $k = 2$ and $k = 3$. On the other hand, increasing $k$ from 2 to 3 is not needed anymore as the third (more scattered) “cluster” vanishes when trimming 5% of the most outlying observations. Thus, there is no discernable difference of the objective functions value with $\alpha \geq \alpha_0 = 0.05$ and $k \geq 2$. Increasing $k$ from 3 to 4 is not needed in any case.
The previously described procedure for making sensible choices for parameters $k$ and $\alpha$ requires an active role from the researcher. The type of restriction and the allowed restriction factor, which do not necessarily depend on the given data set, must be specified in advance. For instance, some specific clustering applications like “location-facilities” problems require almost spherical clusters that can be obtained by setting \texttt{restr = "eigen"} and \texttt{restr.fact} close to 1. The researcher’s decision on the restriction consequently modifies the proper determination of parameters $k$ and $\alpha$.

Due to the important role of the statement of \texttt{restr} and \texttt{restr.fact}, some general guideline for fixing them will be given here. For instance, as already commented, fixing \texttt{restr = "deter"} is recommended when only the relative cluster sizes shall be constrained, or when affine equivariance is clearly needed. On the other hand, using \texttt{restr = "eigen"} is advised when we want to simultaneously constrain relative cluster sizes and shapes.

With respect to the choice of \texttt{restr.fact}, we recommend to initially use large values when applying \texttt{ctlcurves}, thus, providing high flexibility to the clustering method. The default value \texttt{restr.fact = 50} is suggested for \texttt{ctlcurves}, as it worked well with a lot of data sets (especially if the variables have been properly standardized through the \texttt{scale} argument in the \texttt{tclust} function). The so obtained “sensible” values for $k$ and $\alpha$ and their associated clustering solutions must be explored carefully. For instance, \texttt{tclust} issues a warning when the returned clustering solution has been “artificially restricted” by the algorithm, as shown in
Section 9. This means, that the values $M_n$ and $m_n$ (see Section 3.1 and Section 3.2) derived from the returned scatter matrices satisfy $M_n/m_n = \text{restr.fact}$, because the algorithm has forced the chosen constraint, since the (unconstrained) group sample covariance matrices do not satisfy $M_n/m_n \leq \text{restr.fact}$. In this situation, if no specific constraints are required, $\text{restr.fact}$ may be increased stepwise until this warning disappears. Moreover, printing the object returned by the \texttt{ctlcurves} function points out all “artificially restricted” solutions for each computed combination of parameters $k$ and $\alpha$. In this way, if desired, we can easily search for clustering solutions which are not artificially restricted and do not contain spurious clusters. Finally, the exploratory tools in Section 8 also help to evaluate whether all these parameters are reasonably chosen.

Note that arguments \texttt{nstart} and \texttt{iter.max} may be provided in the call to \texttt{ctlcurves} and they are internally passed to function \texttt{tclust}.

The curves presented in García-Escudero et al. (2003) can be considered as precedents of those we obtain by using the \texttt{ctlcurves} function. Trimmed likelihoods have also been taken into account in Neykov et al. (2007) for choosing $k$ and $\alpha$ by using a BIC criterion.

## 8. Graphical displays

As seen in previous examples, the package \texttt{tclust} provides functions for visualizing the computed cluster assignments. One-dimensional, two-dimensional and higher-dimensional cases are visualized differently:

$p = 1$: The one-dimensional data set with the corresponding cluster assignments is displayed along the $x$-axis. Setting the argument \texttt{jitter} = \texttt{TRUE} jitters the data along the $y$-axis in order to increase the visibility of the actual data structure. Additionally, a (robust) scatter estimation of each cluster is also displayed.

$p = 2$: Tolerance ellipsoids are plotted additionally in order to visualize the estimated cluster scatter matrices.

$p > 2$: The first two Fisher’s canonical coordinates are displayed in this case, which are computed based on the estimated cluster scatter matrices. Notice that trimmed observations are not taken into account when computing these coordinates, since they have been completely discarded. The implementation of these canonical coordinates is derived from the function \texttt{discrcoord} as implemented in the package \texttt{fpc} (Hennig 2010).

A simple example demonstrates how the \texttt{plot} function works in different dimensions. The code:

R > geyser1 <- geyser2[, 1, drop = FALSE]
R > geyser3 <- cbind (geyser2, rnorm (nrow (geyser2)))
R > plot (tkmeans (geyser1, k = 2, alpha = 0.03), jitter = TRUE)
R > plot (tkmeans (geyser3, k = 3, alpha = 0.03))

yields Figure 10. For demonstrating the different plotting modes, we have selected one single variable from \texttt{geyser2} to obtain a one-dimensional data set (\texttt{geyser1}), and, added an additional normally distributed variable to \texttt{geyser2}, yielding a three-dimensional data set.
Figure 10: Trimmed $k$-means clustering results for $\text{geyser1}$ (one-dimensional) in (a) and for $\text{geyser3}$ (three-dimensional) in (b). These two data sets are based on $\text{geyser2}$. $k = 2$ is fixed in (a) and $k = 3$ in (b) while $\alpha = 0.03$ is fixed in both cases.

For $\text{geyser3}$. Figure 10 plots the results of the trimmed $k$-means robust clustering method for these two generated data sets.

Given a $\text{tclust}$ object, some additional exploratory graphical tools can be applied in order to evaluate the quality of the cluster assignments and the trimming decisions. This is done by applying the function $\text{DiscrFact}$.

Let $\hat{R} = \{\hat{R}_0, \hat{R}_1, ..., \hat{R}_k\}$, $\hat{\theta} = (\hat{\theta}_1, ..., \hat{\theta}_k)$ and $\hat{\pi} = (\hat{\pi}_1, ..., \hat{\pi}_k)$ be the values obtained by maximizing (2) or (3) (we set $\hat{\pi}_j = 1/k$ when maximizing (2)). $D_j(x_i; \hat{\theta}, \hat{\pi}) = \hat{\pi}_j \phi(x_i, \hat{\theta}_j)$ is a measure of the degree of affiliation of observation $x_i$ with cluster $j$. These values can be ordered as $D_1(x_i; \hat{\theta}, \hat{\pi}) \leq ... \leq D_k(x_i; \hat{\theta}, \hat{\pi})$. Thus the quality of the assignment decision of a non trimmed observation $x_i$ to the cluster $j$ with $D_k(x_i; \hat{\theta}, \hat{\pi}) = D_j(x_i; \hat{\theta}, \hat{\pi})$ can be evaluated by comparing its degree of affiliation with cluster $j$ to the best second possible assignment. That is

$$DF(i) = \log \left( \frac{D_{k-1}(x_i; \hat{\theta}, \hat{\pi})}{D_k(x_i; \hat{\theta}, \hat{\pi})} \right)$$

for $x_i$ not trimmed.

Let $x(1), ..., x(n)$ be the observations in the sample after being sorted according to their $D_k(\cdot; \hat{\theta}, \hat{\pi})$ values, i.e., $D_k(x(1); \hat{\theta}, \hat{\pi}) \leq ... \leq D_k(x(n); \hat{\theta}, \hat{\pi})$. It is not difficult to see that $x(1), ..., x([n\alpha])$ are the trimmed observations which are not assigned to any cluster. Nevertheless, it is possible to compute the degree of affiliation $D_k(x_i; \hat{\theta}, \hat{\pi})$ of a trimmed observation $x_i$ to its nearest cluster. Thus, the quality of the trimming decision on this observation can be evaluated by comparing $D_k(x_i; \hat{\theta}, \hat{\pi})$ to $D_k(x([n\alpha] + 1); \hat{\theta}, \hat{\pi})$, with $x([n\alpha] + 1)$ being the non-trimmed observation with smallest value of $D_k(\cdot; \hat{\theta}, \hat{\pi})$. That is

$$DF(i) = \log \left( \frac{D_k(x_i; \hat{\theta}, \hat{\pi})}{D_k(x([n\alpha] + 1); \hat{\theta}, \hat{\pi})} \right)$$

for $x_i$ trimmed.

Hence, discriminant factors $DF(i) \leq 0$ are obtained for every observation in the data set, whether trimmed or not.
Observations with large DF(i) values (i.e., values close to zero) indicate doubtful assignments or trimming decisions. The use of this type of discriminant factors was already suggested in Van Aelst et al. (2006) in a clustering problem without trimming. “Silhouette” plots (Rousseeuw 1987) can be used for summarizing the obtained ordered discriminant factors. Clusters in the silhouette plot with many large DF(i) values indicate the existence of not very “well-determined” clusters. The most “doubtful” assignments with DF(i) larger than a $\log(\text{threshold})$ value are highlighted by the function DiscrFact.

Figure 11 shows the result of applying the DiscrFact function to a clustering solution found for the mixt data set appearing in Figure 8. The following code is used to obtain this figure:

```r
R > clus.w <- tclust(mixt, k = 3, alpha = 0.1, restr.fact = 1, + equal.weights = TRUE)
R > discr.clus.w <- DiscrFact(clus.w, threshold = 0.1)
R > plot(discr.clus.w)
```

The choice threshold = 0.1 means that a decision on a particular observation $x_i$ is considered as doubtful, if the quality of the second best possible decision ($D_{(k-1)}(x_i; \hat{\theta}, \hat{\pi})$ or $D_{(k)}(x_{\lceil n\alpha \rceil+1}; \hat{\theta}, \hat{\pi})$ for trimmed observations) is larger than one tenth of the quality of the actually made decision ($D_{(k)}(x_i; \hat{\theta}, \hat{\pi})$).

Although Figure 9 suggests to choose $k = 2$, $k$ has been increased to 3 in order to show how such a change leads to doubtful cluster assignment decisions which can be visualized by DiscrFact. Figure 11(a) simply illustrates the cluster assignments and trimming decisions. The mentioned silhouette plot is presented in (b), whereas the doubtful decisions are marked
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Figure 12: Classification trimmed likelihoods for \( k = 1, \ldots, 4 \) and \( \alpha = 0, .025, \ldots, .3 \) when \texttt{restr.fact} = 50 for the “Swiss Bank notes” data set.

in (c). All observations with \( \text{DF}(i) \geq \log(0.1) \) are highlighted as they are plotted darker/in color. Most of the doubtful decisions are located in the overlapping area of the two artificially found clusters (highlighted symbols “×” and “+”). Some doubtfully trimmed observations (highlighted symbol “◦”) are located in the boundaries of these two clusters.

### 9. Swiss Bank notes data

The well-known “Swiss Bank notes” data set includes 6 numerical measurements (six-dimensional data set) made on 100 genuine and 100 counterfeit old Swiss 1000-franc bank notes (Flury and Riedwyl 1988). The following code can be used to obtain the classification trimmed likelihoods shown in Figure 12.

\[
\text{R > data ("swissbank")}
\]
\[
\text{R > plot (ctlcurves (swissbank, k = 1:4, alpha = seq (0, 0.3, by = 0.025)))}
\]

This figure indicates the clear existence of \( k = 2 \) main clusters (“genuine” and “forged” bills). Moreover, considering the clear difference between \( \mathcal{L}_{50}^{H}(0, 3) \) and \( \mathcal{L}_{50}^{H}(0, 2) \), we can see that a further cluster, i.e., \( k = 3 \), is needed when no trimming is allowed. This extra cluster can be justified by the heterogeneity of the group of forgeries (perhaps due to the presence of different sources of forged bills).
Figure 13: Clustering results with \( k = 2 \), \( \alpha = 0.1 \) and \( \text{restr.fact} = 50 \) summarized by the use of \texttt{DiscrFact} function for the “Swiss Bank notes” data set. The threshold value is chosen in order to highlight the 7 most doubtful cluster assignments.

Considering Figure 12, the choice \( k = 2 \) and a value of \( \alpha \) close to 0.1 also seem sensible. Notice that \( \mathcal{L}_{50}^{\Pi}(\alpha, 3) \) is clearly larger than \( \mathcal{L}_{50}^{\Pi}(\alpha, 2) \) for \( \alpha < 0.1 \) while these differences are not so big when \( \alpha \geq 0.1 \). We can even see smaller differences in the classification trimmed likelihood curves when increasing \( k \) from 3 to 4. However, these differences are less significant than those previously commented. More spurious clusters can be surely found but they have less entity and importance.

Figure 13 shows the clustering results with \( k = 2 \), \( \alpha = 0.1 \) and \( \text{restr.fact} = 50 \) obtained by executing the code:

\[
\text{R} > \text{clus} <- \text{tclust} (\text{swissbank}, k = 2, \alpha = 0.1, \text{restr.fact} = 50) \\
\text{R} > \text{plot} (\text{DiscrFact (clus, threshold = 0.0001)})
\]

Notice that, in this example, we did not want to impose a specific constraint on the solution. Thus, the default parameter \( \text{restr.fact} = 50 \) has initially been used in \texttt{ctlcurves}. After choosing the combination \( \alpha = 0.10 \) and \( k = 2 \), we could try to reduce the restriction factor which resulted in a warning:

\[
\text{R} > \text{tclust} (\text{swissbank}, k = 2, \alpha = 0.1, \text{restr.fact} = 40) \\
\text{In .tclust.warn(warnings, ret):} \\
\quad \text{The result is artificially constrained due to restr.fact = 40.}
\]

Thus the choice \( \text{restr.fact} = 50 \) seems appropriate as it does not artificially restrict the result, whereas a slightly smaller restriction factor (40) does. By examining the sizes of the obtained groups, we see that no spurious groups are found with \( \text{restr.fact} = 50 \):
Figure 14: Clustering results with $k = 2$, $\alpha = .1$ and restr_fact = 50 for the “Swiss Bank notes” data set. Only the fourth and sixth variables are plotted. The 7 most doubtful decisions are rounded by a circle symbol.

R > clus$size
   [1] 95 85

We have used restr = "eigen" in this example, but restr = "deter" can be also successfully applied with smaller values of restr_fact.

We also use the function DiscrFact to summarize the obtained clustering results. The two first Fisher’s canonical coordinates derived from the final cluster assignments are plotted. The threshold value 0.0001 is chosen in order to highlight the 7 most doubtful decisions.

Finally, Figure 14 shows a scatterplot of the fourth (“Distance of the inner frame to lower border”) against the sixth variable (“Length of the diagonal”) with the corresponding cluster assignments. We use the symbols “G” for the genuine bills and “F” for the forged ones. The 7 most doubtful decisions (i.e., the observations with largest DF(i) values that were highlighted in Figure 13,(c)) are surrounded by circles in this figure. We can see that “Cluster 1” essentially includes most of the “forged” bills while “Cluster 2” includes most of the “genuine” ones. Among the trimmed observations, we can find a subset of 15 forged bills following a clearly different forgery pattern that has been previously commented by other authors (see, e.g., Flury and Riedwyl 1988; Cook 1999). These most doubtful assignments include 5 “genuine” bills that have perhaps been wrongly trimmed.

10. Conclusion

This paper presents a package called tclust for robust (non-hierarchical) clustering. The
implementation is flexible, so only the restrictions on the cluster scatters have to be changed in
order to carry out different robust clustering algorithms. Robustness is achieved by trimming
a specific proportion of observations which are identified as the “most outlying” ones.
Although this R-package implements robust clustering approaches which have already been
described in the literature, some of these approaches have been extended to provide increased
flexibility. The package also provides some graphical tools which on the one hand help to chose
appropriate parameters (ctlcurves) and on the other hand help to estimate the adequacy of
a particular clustering solution (DiscrFact).
Future work on this package focuses on implementing additional types of scatter restrictions,
making the algorithm even more flexible, and on providing numerical tools for automatically
choosing the number of clusters and the trimming proportion.

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