

Cluster Analysis in R

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Introduction

- What is Clustering?
 - Clustering is the classification of data objects into similarity groups (clusters) according to a defined distance measure.
 - It is used in many fields, such as machine learning, data mining, pattern recognition, image analysis, genomics, systems biology, etc.
 - Machine learning typically regards data clustering as a form of unsupervised learning.
- Why Clustering and Data Mining in R?
 - Efficient data structures and functions for clustering
 - Reproducible and programmable
 - Comprehensive set of clustering and machine learning libraries
 - Integration with many other data analysis tools
- Useful Links
 - Cluster Task Views
 - Machine Learning Task Views
 - UCR Manual

Data Preprocessing

Data Transformations

Choice depends on data set!

- Center and standardize
 1. Center: subtract from each value the mean of the corresponding vector
 2. Standardize: divide by standard deviation
 - Result: $Mean = 0$ and $STDEV = 1$
- Center and scale with the `scale()` function
 1. Center: subtract from each value the mean of the corresponding vector
 2. Scale: divide centered vector by their *root mean square (rms)*:

$$x_{rms} = \sqrt{\frac{1}{n-1} \sum_{i=1}^n x_i^2}$$

- Result: $Mean = 0$ and $STDEV = 1$
- Log transformation
- Rank transformation: replace measured values by ranks
- No transformation

Distance Methods

List of most common ones!

- Euclidean distance for two profiles X and Y :

$$d(X, Y) = \sqrt{\sum_{i=1}^n (x_i - y_i)^2}$$

- **Disadvantages:** not scale invariant, not for negative correlations
- Maximum, Manhattan, Canberra, binary, Minowski, ...
- Correlation-based distance: $1-r$
 - Pearson correlation coefficient (PCC):

$$r = \frac{n \sum_{i=1}^n x_i y_i - \sum_{i=1}^n x_i \sum_{i=1}^n y_i}{\sqrt{(\sum_{i=1}^n x_i^2 - (\sum_{i=1}^n x_i)^2)(\sum_{i=1}^n y_i^2 - (\sum_{i=1}^n y_i)^2)}}$$

- * **Disadvantage:** outlier sensitive
- Spearman correlation coefficient (SCC)
 - * Same calculation as PCC but with ranked values!

There are many more distance measures

- If the distances among items are quantifiable, then clustering is possible.
- Choose the most accurate and meaningful distance measure for a given field of application.
- If uncertain then choose several distance measures and compare the results.

Cluster Linkage

Clustering Algorithms

Hierarchical Clustering

Overview of algorithm

1. Identify clusters (items) with closest distance
2. Join them to new clusters
3. Compute distance between clusters (items)
4. Return to step 1

Hierarchical clustering: agglomerative Approach

Hierarchical Clustering with Heatmap

- A heatmap is a color coded table. To visually identify patterns, the rows and columns of a heatmap are often sorted by hierarchical clustering trees.
- In case of gene expression data, the row tree usually represents the genes, the column tree the treatments and the colors in the heat table represent the intensities or ratios of the underlying gene expression data set.

Hierarchical Clustering Approaches

1. Agglomerative approach (bottom-up)
 - R functions: `hclust()` and `agnes()`
2. Divisive approach (top-down)
 - R function: `diana()`

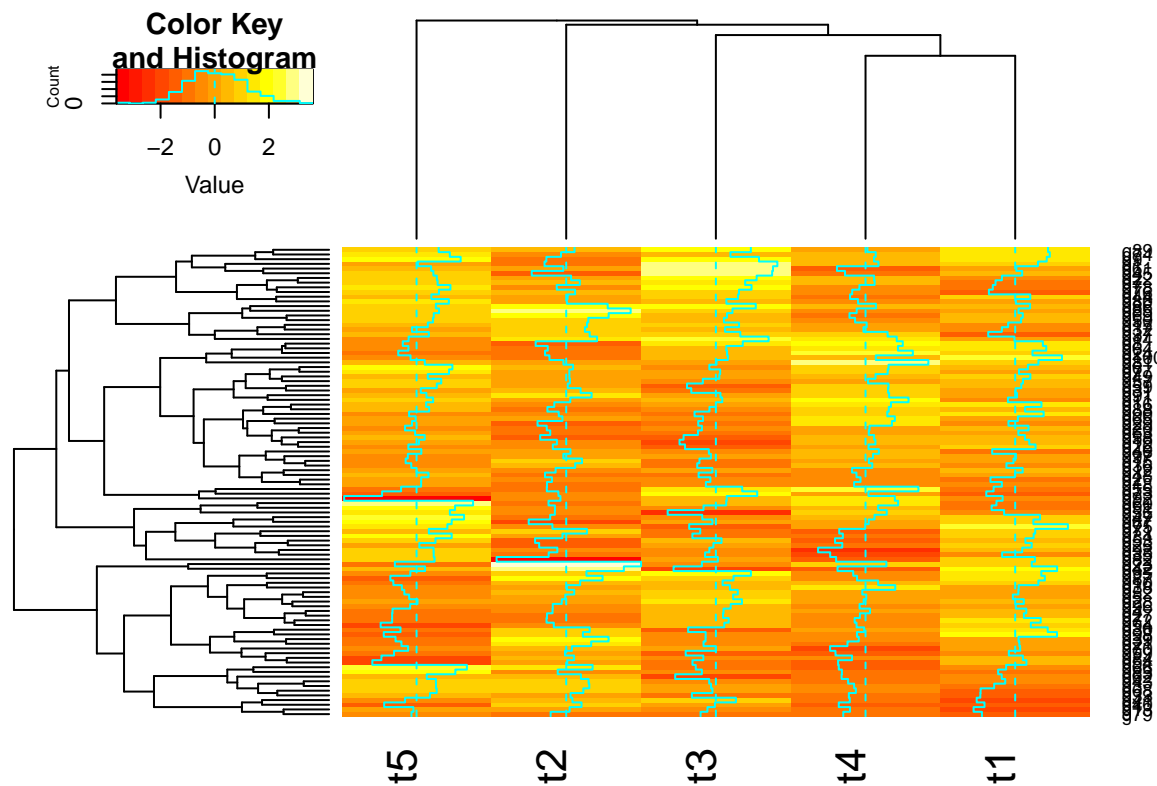
Tree Cutting to Obtain Discrete Clusters

1. Node height in tree
2. Number of clusters
3. Search tree nodes by distance cutoff

Examples

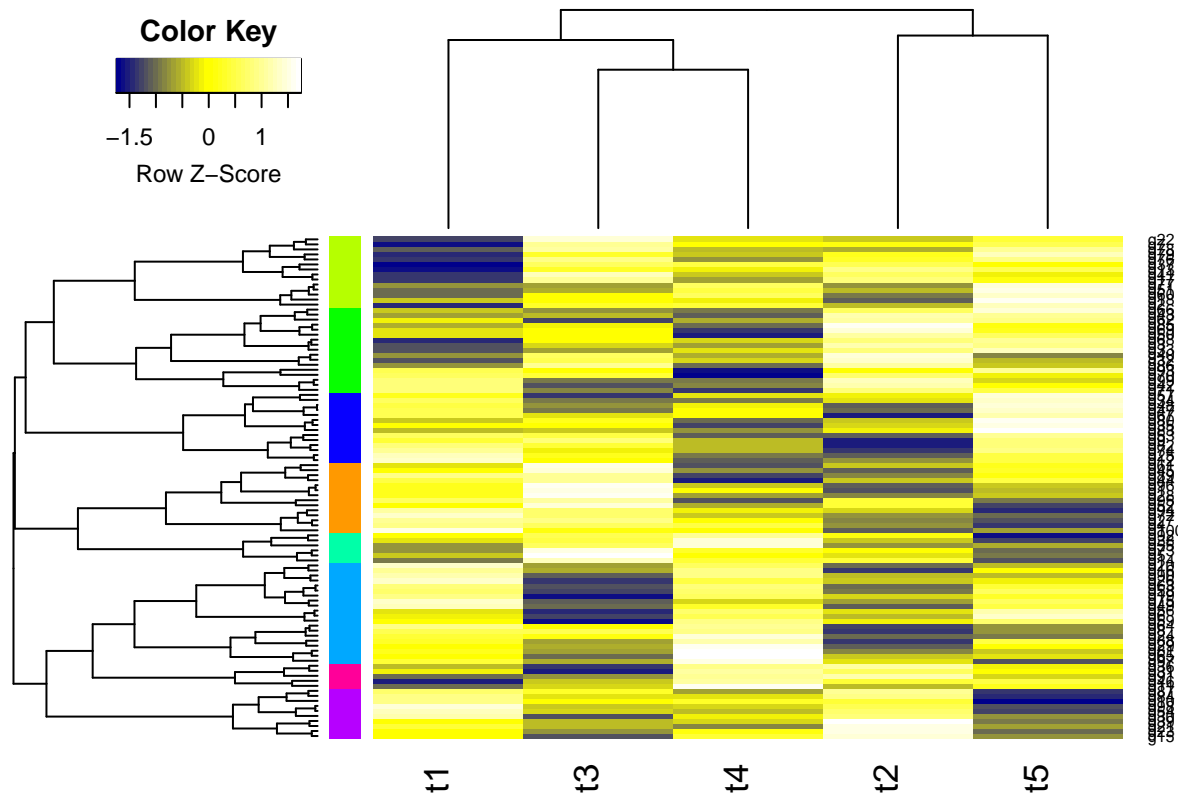
Using `hclust` and `heatmap.2`

```
library(gplots)
y <- matrix(rnorm(500), 100, 5, dimnames=list(paste("g", 1:100, sep=""), paste("t", 1:5, sep="")))
heatmap.2(y) # Shortcut to final result
```



Stepwise Approach with Tree Cutting

```
## Row- and column-wise clustering
hr <- hclust(as.dist(1-cor(t(y), method="pearson")), method="complete")
hc <- hclust(as.dist(1-cor(y, method="spearman")), method="complete")
## Tree cutting
mycl <- cutree(hr, h=max(hr$height)/1.5); mycolhc <- rainbow(length(unique(mycl)), start=0.1, end=0.9);
## Plot heatmap
mycol <- colorpanel(40, "darkblue", "yellow", "white") # or try redgreen(75)
heatmap.2(y, Rowv=as.dendrogram(hr), Colv=as.dendrogram(hc), col=mycol, scale="row", density.info="none")
```



K-Means Clustering

Overview of algorithm

1. Choose the number of k clusters
2. Randomly assign items to the k clusters
3. Calculate new centroid for each of the k clusters
4. Calculate the distance of all items to the k centroids
5. Assign items to closest centroid
6. Repeat until clusters assignments are stable

Examples

```
km <- kmeans(t(scale(t(y))), 3)
km$cluster
```

##	g1	g2	g3	g4	g5	g6	g7	g8	g9	g10	g11	g12	g13	g14	g15	g16	g17	g18	g19	g20
##	1	3	3	1	1	2	3	2	3	1	2	3	3	1	2	2	1	3	2	2
##	g21	g22	g23	g24	g25	g26	g27	g28	g29	g30	g31	g32	g33	g34	g35	g36	g37	g38	g39	g40
##	1	3	1	2	3	2	1	1	2	2	2	1	3	3	3	2	1	2	1	1
##	g41	g42	g43	g44	g45	g46	g47	g48	g49	g50	g51	g52	g53	g54	g55	g56	g57	g58	g59	g60
##	1	1	3	3	3	2	3	3	2	3	3	1	3	1	2	2	3	2	2	3
##	g61	g62	g63	g64	g65	g66	g67	g68	g69	g70	g71	g72	g73	g74	g75	g76	g77	g78	g79	g80
##	1	3	2	2	2	3	3	3	1	1	3	1	1	3	2	3	3	3	3	3
##	g81	g82	g83	g84	g85	g86	g87	g88	g89	g90	g91	g92	g93	g94	g95	g96	g97	g98	g99	g100
##	2	3	3	1	1	1	1	2	1	2	2	1	3	1	1	1	2	1	1	2

Fuzzy C-Means Clustering

- In contrast to strict (hard) clustering approaches, fuzzy (soft) clustering methods allow multiple cluster memberships of the clustered items (Hathaway, Bezdek, and Pal 1996).
- This is commonly achieved by assigning to each item a weight of belonging to each cluster.
- Thus, items at the edge of a cluster, may be in a cluster to a lesser degree than items at the center of a cluster.
- Typically, each item has as many coefficients (weights) as there are clusters that sum up for each item to one.

Examples

Fuzzy Clustering with fanny

```
library(cluster) # Loads the cluster library.
fanny <- fanny(y, k=4, metric = "euclidean", memb.exp = 1.2)
round(fanny$membership, 2)[1:4,]
```

```
##      [,1] [,2] [,3] [,4]
## g1 0.78 0.03 0.08 0.11
## g2 0.03 0.91 0.03 0.03
## g3 0.07 0.89 0.02 0.02
## g4 0.03 0.02 0.88 0.07
```

```
fanny$clustering
```

##	g1	g2	g3	g4	g5	g6	g7	g8	g9	g10	g11	g12	g13	g14	g15	g16	g17	g18	g19	g20
##	1	2	2	3	1	4	1	4	1	3	4	2	2	1	3	4	1	2	4	4
##	g21	g22	g23	g24	g25	g26	g27	g28	g29	g30	g31	g32	g33	g34	g35	g36	g37	g38	g39	g40
##	3	1	3	4	2	4	3	1	4	3	4	1	2	2	2	3	1	4	3	3
##	g41	g42	g43	g44	g45	g46	g47	g48	g49	g50	g51	g52	g53	g54	g55	g56	g57	g58	g59	g60
##	2	3	2	1	1	2	4	2	4	1	2	1	2	3	4	4	4	4	4	4
##	g61	g62	g63	g64	g65	g66	g67	g68	g69	g70	g71	g72	g73	g74	g75	g76	g77	g78	g79	g80
##	1	2	4	4	4	2	4	2	1	3	2	3	1	4	4	1	4	1	2	1
##	g81	g82	g83	g84	g85	g86	g87	g88	g89	g90	g91	g92	g93	g94	g95	g96	g97	g98	g99	g100
##	4	2	2	3	1	1	3	4	1	4	1	3	2	3	1	3	3	3	3	4

Principal Component Analysis (PCA)

Principal components analysis (PCA) is a data reduction technique that allows to simplify multidimensional data sets to 2 or 3 dimensions for plotting purposes and visual variance analysis.

Basic Steps

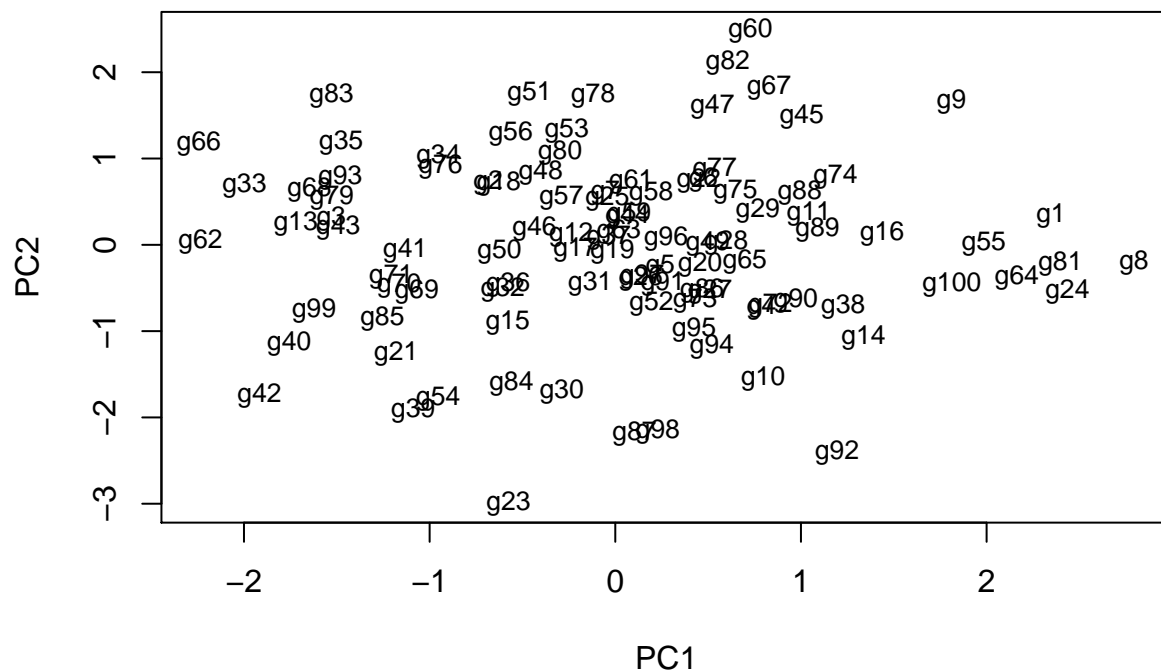
- Center (and standardize) data
- First principal component axis
 - Across centroid of data cloud
 - Distance of each point to that line is minimized, so that it crosses the maximum variation of the data cloud
- Second principal component axis
 - Orthogonal to first principal component
 - Along maximum variation in the data
- First PCA axis becomes x-axis and second PCA axis y-axis
- Continue process until the necessary number of principal components is obtained

Example

```
pca <- prcomp(y, scale=T)
summary(pca) # Prints variance summary for all principal components
```

```
## Importance of components%s:
##          PC1      PC2      PC3      PC4      PC5
## Standard deviation    1.1055 1.0346 1.0148 0.9312 0.9003
## Proportion of Variance 0.2444 0.2141 0.2059 0.1734 0.1621
## Cumulative Proportion 0.2444 0.4585 0.6644 0.8379 1.0000
```

```
plot(pca$x, pch=20, col="blue", type="n") # To plot dots, drop type="n"
text(pca$x, rownames(pca$x), cex=0.8)
```



and 2nd principal components explain x% of variance in data.

1st

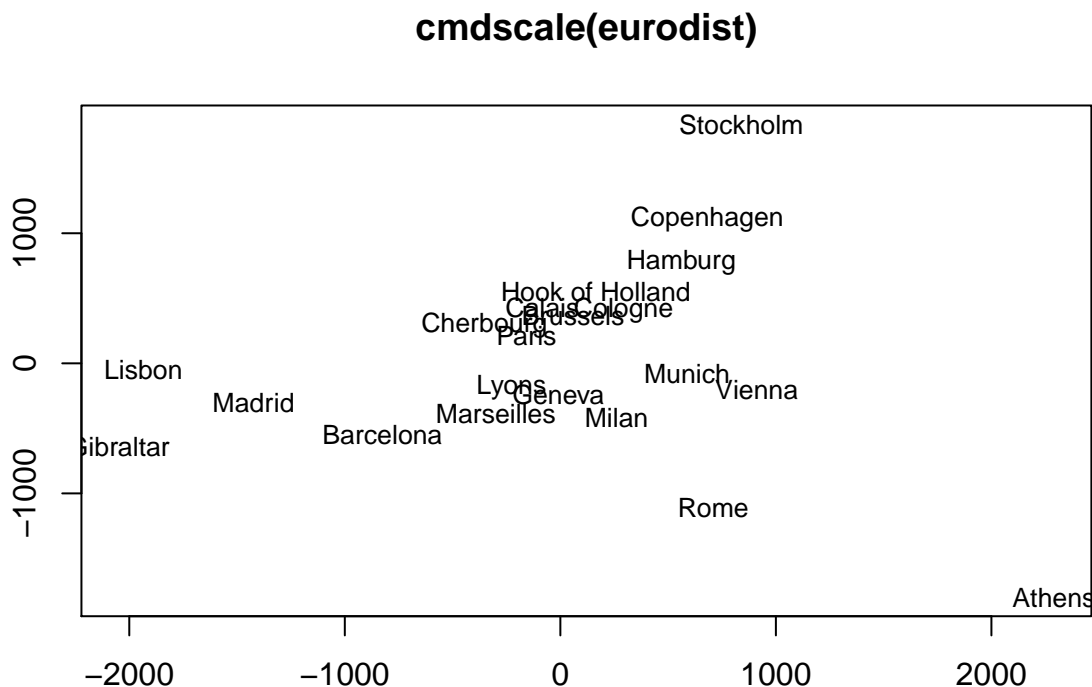
Multidimensional Scaling (MDS)

- Alternative dimensionality reduction approach
- Represents distances in 2D or 3D space
- Starts from distance matrix (PCA uses data points)

Example

The following example performs MDS analysis with `cmdscale` on the geographic distances among European cities.

```
loc <- cmdscale(eurodist)
plot(loc[,1], -loc[,2], type="n", xlab="", ylab="", main="cmdscale(eurodist)")
text(loc[,1], -loc[,2], rownames(loc), cex=0.8)
```



Biclustering

Finds in matrix subgroups of rows and columns which are as similar as possible to each other and as different as possible to the remaining data points.

Unclustered —————> Clustered

Similarity Measures for Clusters

- Compare the numbers of identical and unique item pairs appearing in cluster sets
- Achieved by counting the number of item pairs found in both clustering sets (a) as well as the pairs appearing only in the first (b) or the second (c) set.
- With this a similarity coefficient, such as the Jaccard index, can be computed. The latter is defined as the size of the intersect divided by the size of the union of two sample sets: $a/(a+b+c)$.

- In case of partitioning results, the Jaccard Index measures how frequently pairs of items are joined together in two clustering data sets and how often pairs are observed only in one set.
- Related coefficient are the Rand Index and the Adjusted Rand Index. These indices also consider the number of pairs (d) that are not joined together in any of the clusters in both sets.

Example:

Jaccard index for cluster sets

The following imports the `cindex()` function and computes the Jaccard Index for two sample clusters.

```
source("http://faculty.ucr.edu/~tgirke/Documents/R_BioCond/My_R_Scripts/clusterIndex.R")
library(cluster); y <- matrix(rnorm(5000), 1000, 5, dimnames=list(paste("g", 1:1000, sep=""), paste("t", 1:5, sep="")))
ci <- cindex(clV1=clV1, clV2=clV2, self=FALSE, minSZ=1, method="jaccard")
ci[2:3] # Returns Jaccard index and variables used to compute it

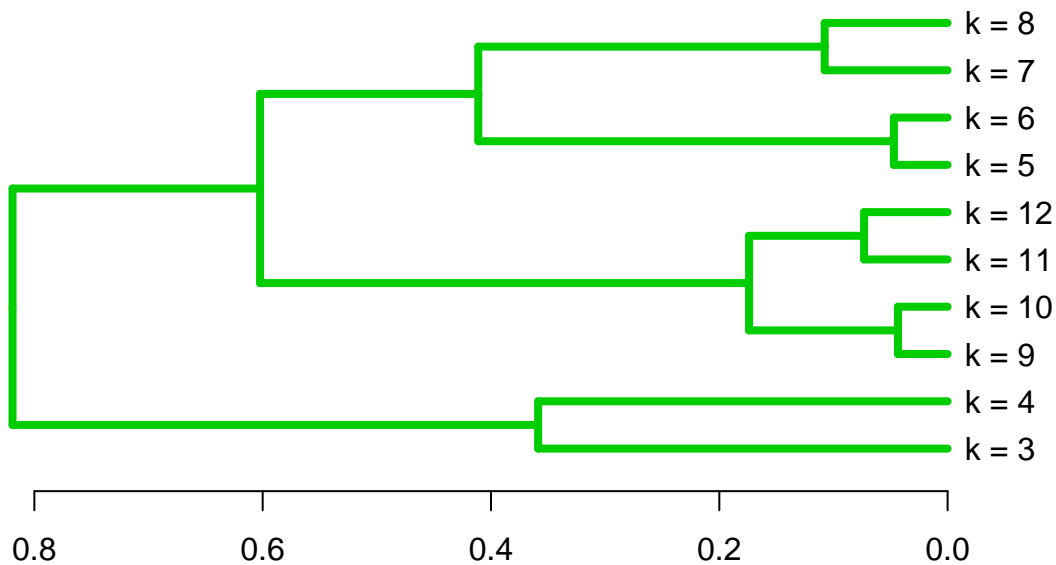
## $variables
##      a      b      c
## 5073 8762 8423
##
## $Jaccard_Index
## [1] 0.2279181
```

Clustering cluster sets with Jaccard index

The following example shows how one can cluster entire cluster result sets. First, 10 sample cluster results are created with Clara using k-values from 3 to 12. The results are stored as named clustering vectors in a list object. Then a nested sapply loop is used to generate a similarity matrix of Jaccard Indices for the clustering results. After converting the result into a distance matrix, hierarchical clustering is performed with `hclust`.

```
clVlist <- lapply(3:12, function(x) clara(y[1:30, ], k=x)$clustering); names(clVlist) <- paste("k", "=", 3:12)
d <- sapply(names(clVlist), function(x) sapply(names(clVlist), function(y) cindex(clV1=clVlist[[x]], clV2=clVlist[[y]])))
hv <- hclust(as.dist(1-d))
plot(as.dendrogram(hv), edgePar=list(col=3, lwd=4), horiz=T, main="Similarities of 10 Clara Clustering Results")
```


Similarities of 10 Clara Clustering Results for k: 3–12



- Remember: there are many additional clustering algorithms.
- Additional details can be found in the Clustering Section of the R/Bioconductor Manual.

Clustering Exercises

Data Preprocessing

Scaling

```
## Sample data set
set.seed(1410)
y <- matrix(rnorm(50), 10, 5, dimnames=list(paste("g", 1:10, sep=""),
                                             paste("t", 1:5, sep="")))
dim(y)
```

```
## [1] 10 5
```

```
## Scaling
yscaled <- t(scale(t(y))) # Centers and scales y row-wise
apply(yscaled, 1, sd)
```

```
## g1 g2 g3 g4 g5 g6 g7 g8 g9 g10
## 1 1 1 1 1 1 1 1 1 1
```

Distance Matrices

Euclidean distance matrix

```
dist(y[1:4,], method = "euclidean")
```

```
##           g1           g2           g3
## g2 4.793697
## g3 4.932658 6.354978
## g4 4.033789 4.788508 1.671968
```

Correlation-based distance matrix

Correlation matrix

```
c <- cor(t(y), method="pearson")
as.matrix(c)[1:4,1:4]
```

```
##           g1           g2           g3           g4
## g1 1.00000000 -0.2965885 -0.00206139 -0.4042011
## g2 -0.29658847 1.0000000 -0.91661118 -0.4512912
## g3 -0.00206139 -0.9166112 1.00000000 0.7435892
## g4 -0.40420112 -0.4512912 0.74358925 1.0000000
```

Correlation-based distance matrix

```
d <- as.dist(1-c)
as.matrix(d)[1:4,1:4]
```

```
##           g1           g2           g3           g4
## g1 0.0000000 1.296588 1.0020614 1.4042011
## g2 1.296588 0.000000 1.9166112 1.4512912
## g3 1.002061 1.916611 0.0000000 0.2564108
## g4 1.404201 1.451291 0.2564108 0.0000000
```

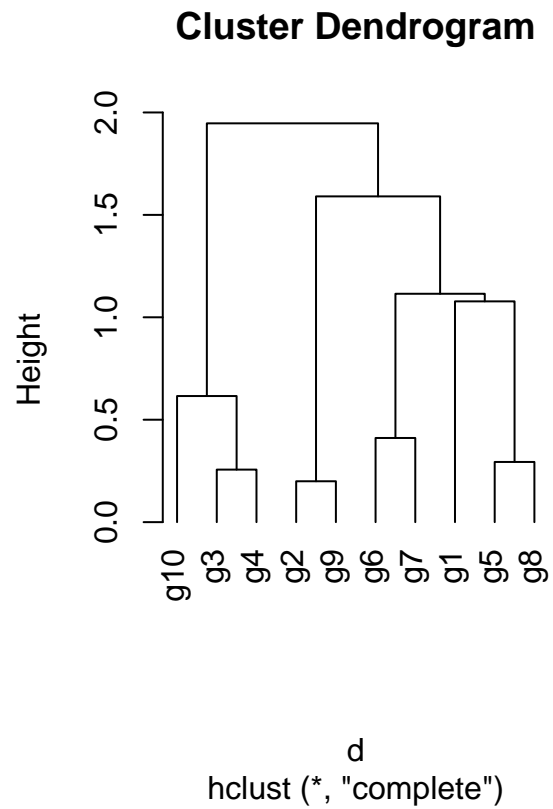
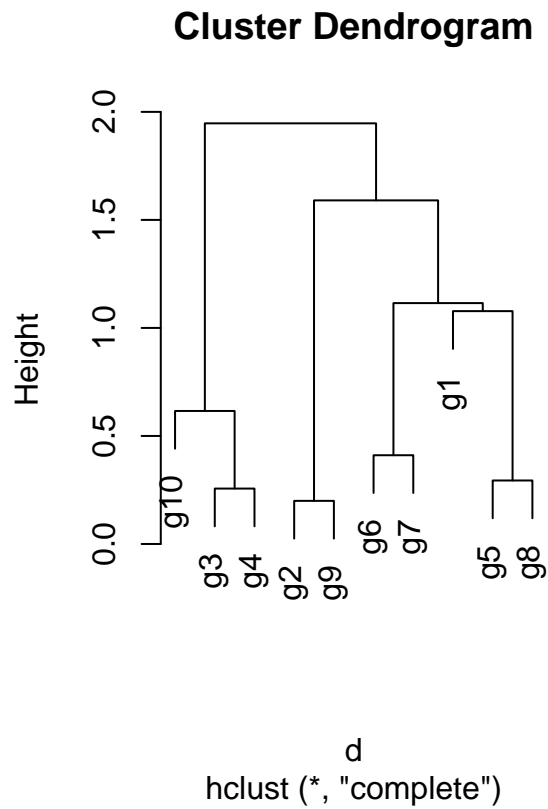
Hierarchical Clustering with hclust

Hierarchical clustering with complete linkage and basic tree plotting

```
hr <- hclust(d, method = "complete", members=NULL)
names(hr)
```

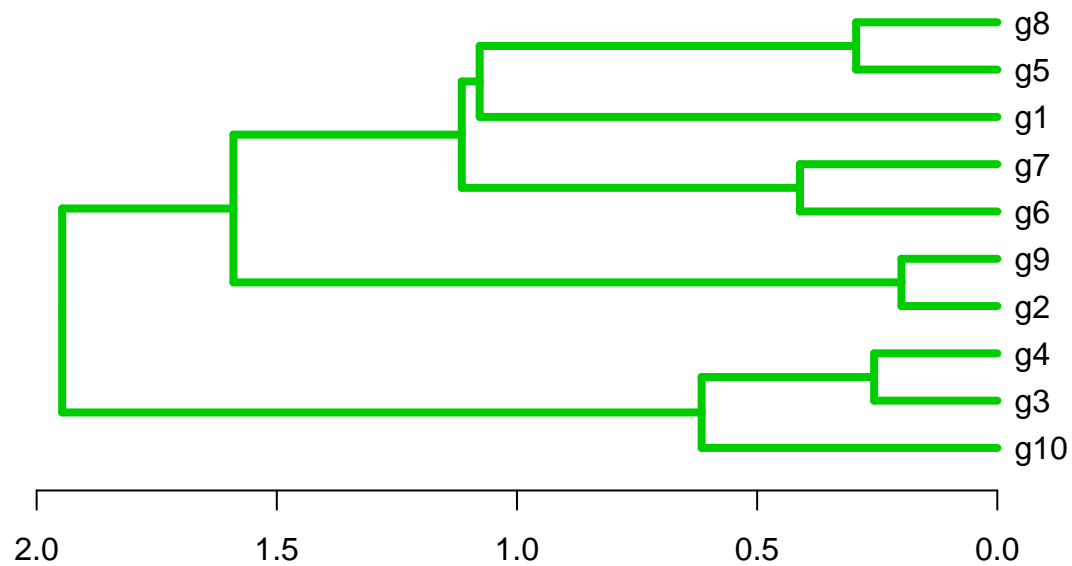
```
## [1] "merge"      "height"     "order"      "labels"     "method"     "call"
## [7] "dist.method"
```

```
par(mfrow = c(1, 2)); plot(hr, hang = 0.1); plot(hr, hang = -1)
```



Tree plotting I

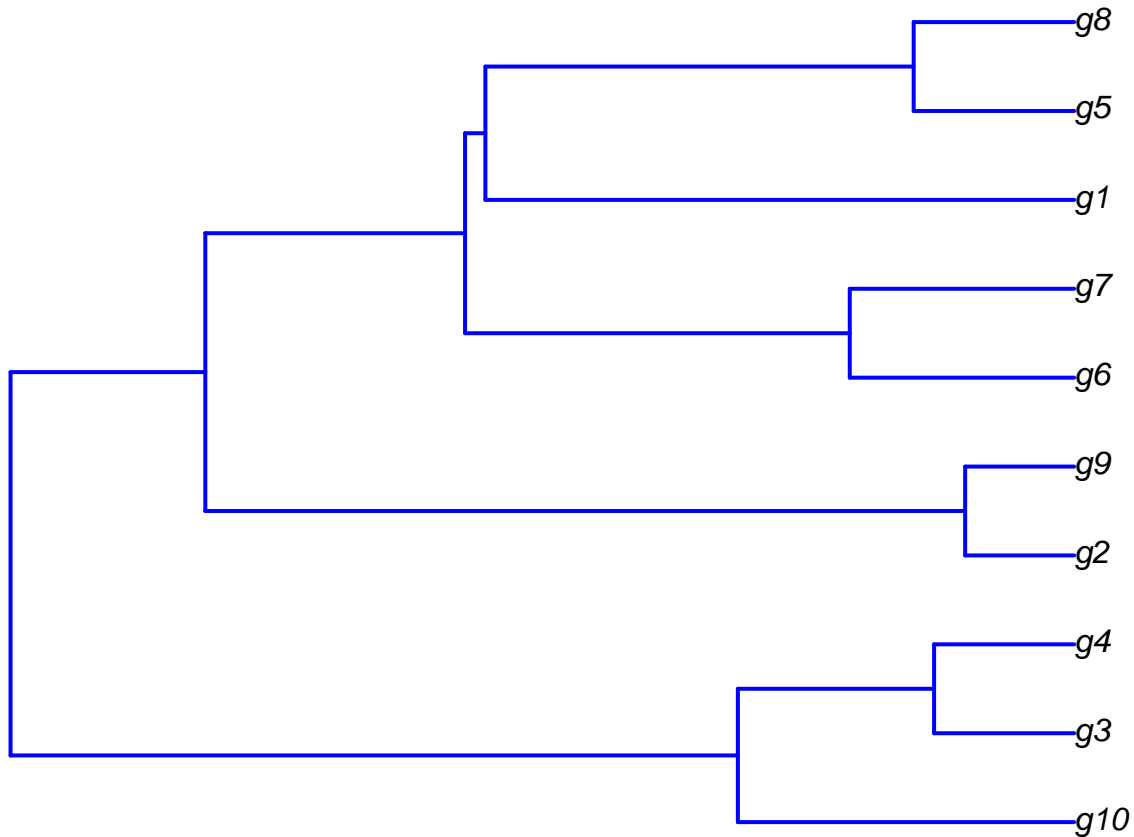
```
plot(as.dendrogram(hr), edgePar=list(col=3, lwd=4), horiz=T)
```



Tree plotting II

The **ape** library provides more advanced features for tree plotting

```
library(ape)
plot.phylo(as.phylo(hr), type="p", edge.col=4, edge.width=2,
           show.node.label=TRUE, no.margin=TRUE)
```



Tree Cutting

Accessing information in *hclust* objects

```
hr
```

```
##
## Call:
## hclust(d = d, method = "complete", members = NULL)
##
## Cluster method   : complete
## Number of objects: 10
```

```
## Print row labels in the order they appear in the tree
hr$labels[hr$order]
```

```
## [1] "g10" "g3"  "g4"  "g2"  "g9"  "g6"  "g7"  "g1"  "g5"  "g8"
```

Tree cutting with `cutree`

```
mycl <- cutree(hr, h=max(hr$height)/2)
mycl[hr$labels[hr$order]]
```

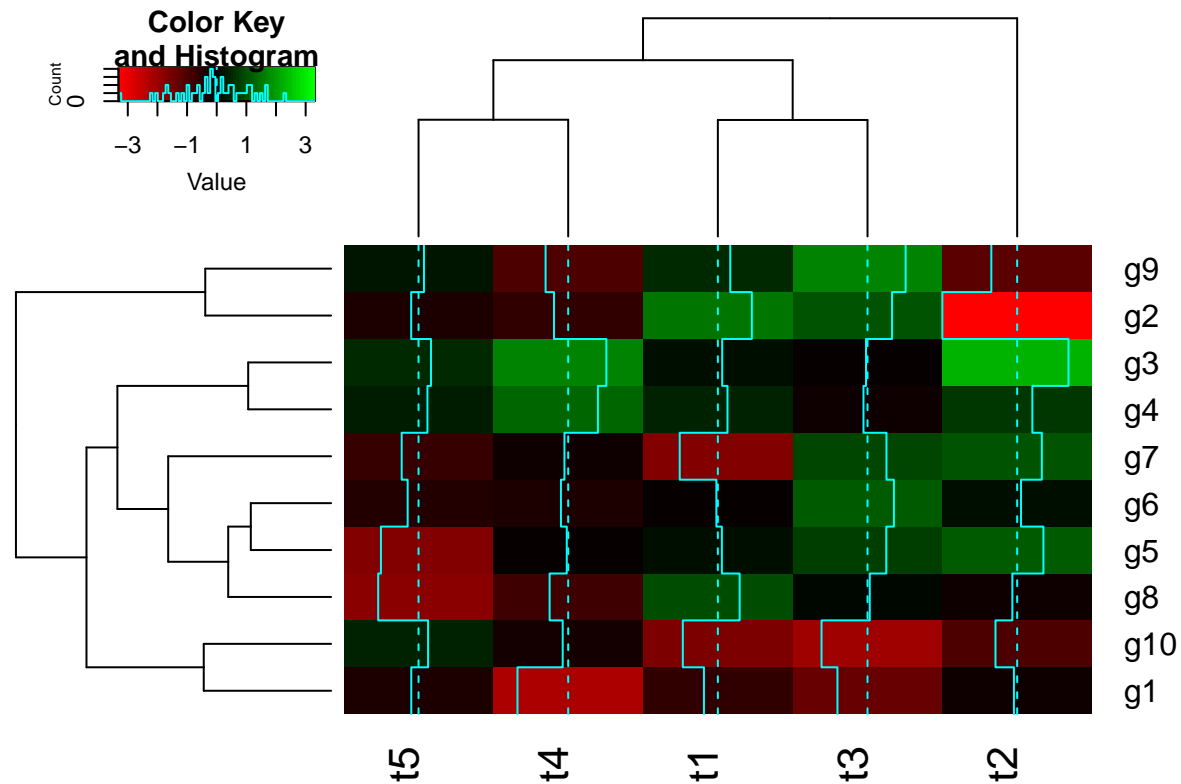
```
## g10 g3 g4 g2 g9 g6 g7 g1 g5 g8
##    3  3  3  2  2  5  5  1  4  4
```

Heatmaps

With `heatmap.2`

All in one step: clustering and heatmap plotting

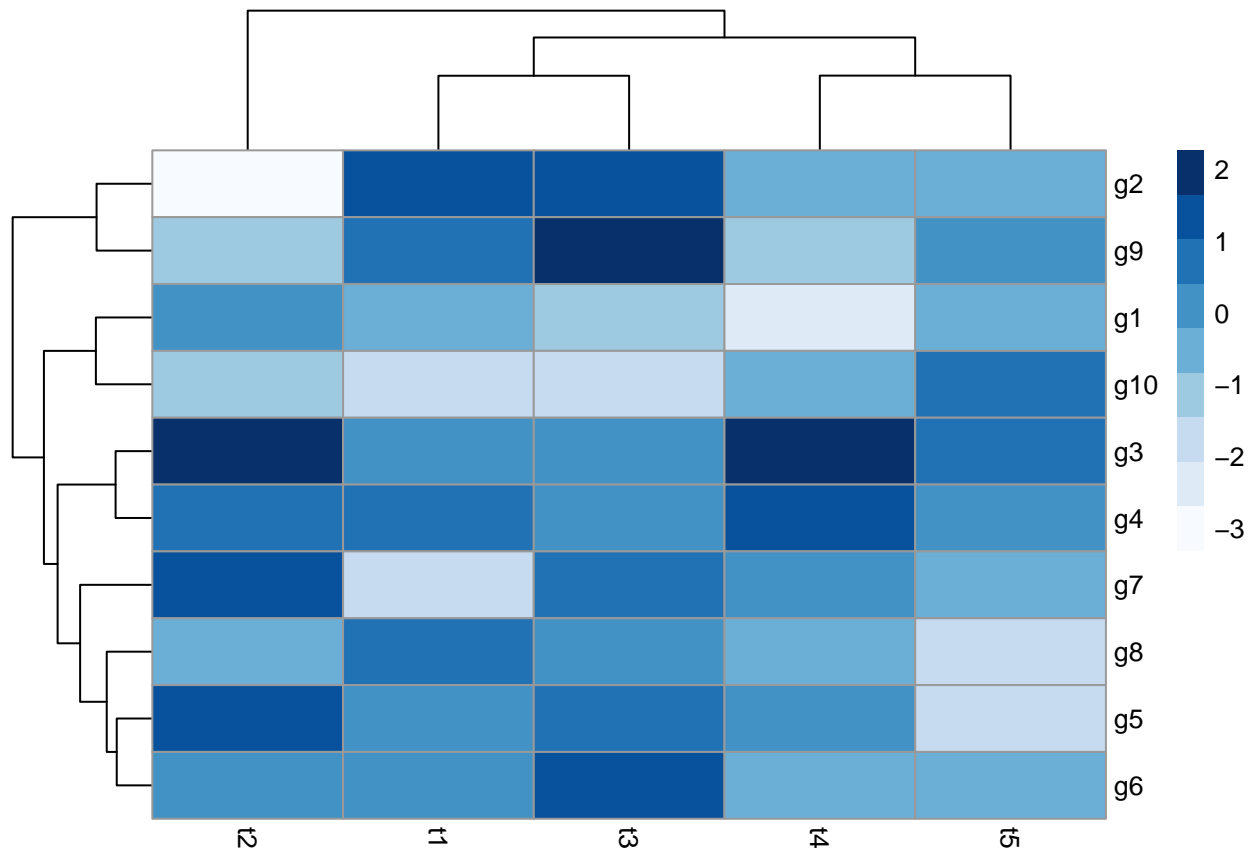
```
library(gplots)
heatmap.2(y, col=redgreen(75))
```



With `pheatmap`

All in one step: clustering and heatmap plotting

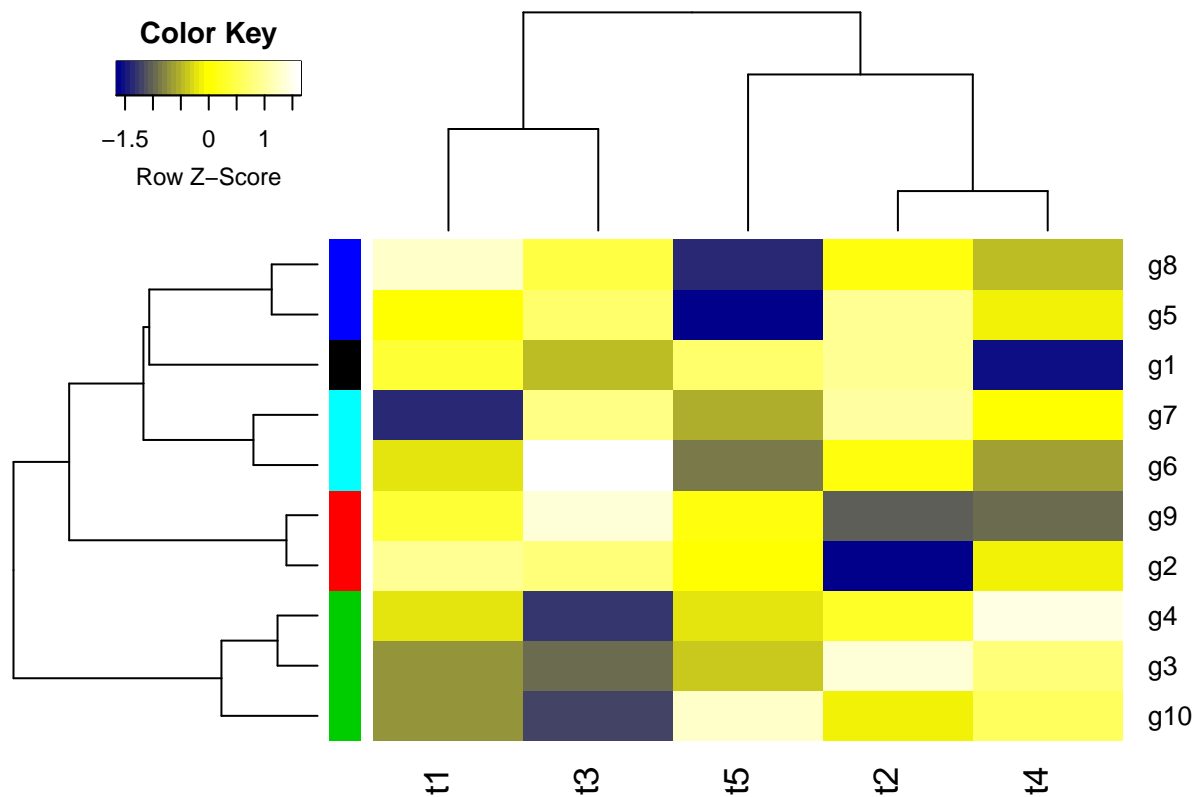
```
library(pheatmap); library("RColorBrewer")
pheatmap(y, color=brewer.pal(9,"Blues"))
```



Customizing heatmaps

Customizes row and column clustering and shows tree cutting result in row color bar. Additional color schemes can be found [here](#).

```
hc <- hclust(as.dist(1-cor(y, method="spearman")), method="complete")
mycol <- colorpanel(40, "darkblue", "yellow", "white")
heatmap.2(y, Rowv=as.dendrogram(hr), Colv=as.dendrogram(hc), col=mycol,
          scale="row", density.info="none", trace="none",
          RowSideColors=as.character(mycol))
```



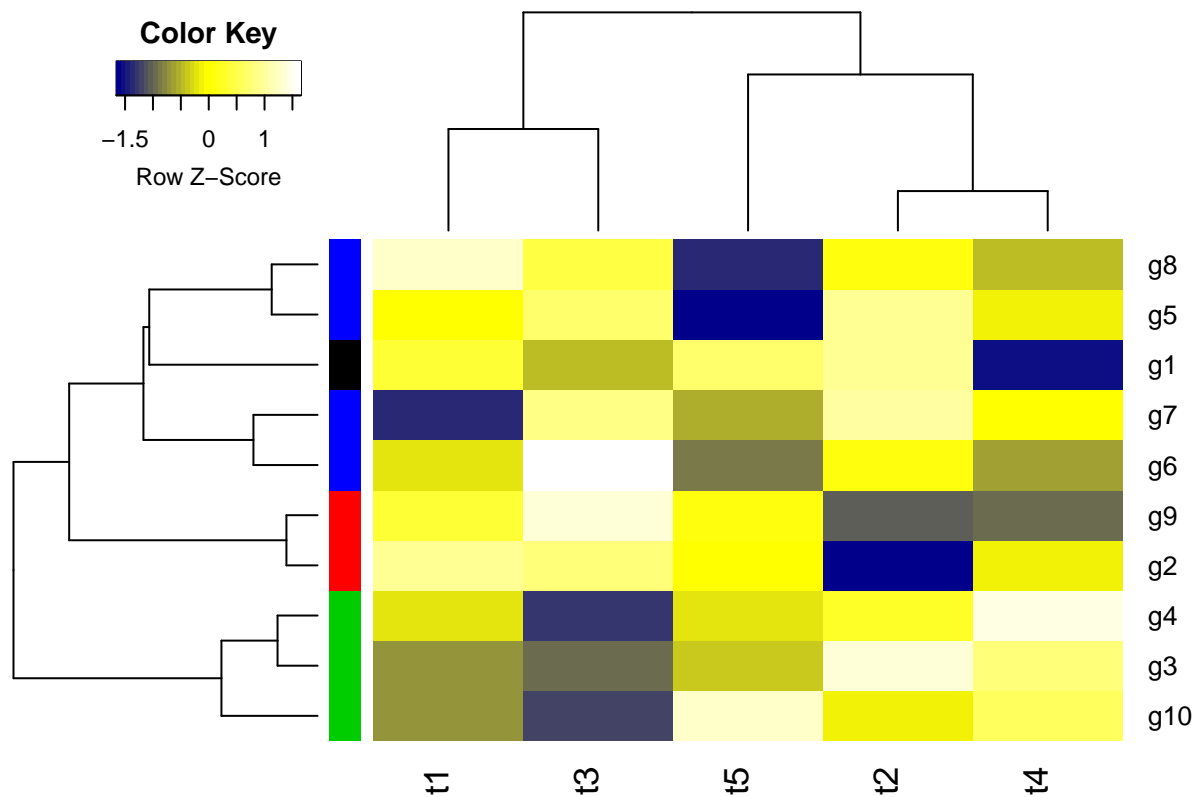
K-Means Clustering with PAM

Runs K-means clustering with PAM (partitioning around medoids) algorithm and shows result in color bar of hierarchical clustering result from before.

```
library(cluster)
pamy <- pam(d, 4)
(kmcol <- pamy$clustering)
```

```
## g1 g2 g3 g4 g5 g6 g7 g8 g9 g10
## 1 2 3 3 4 4 4 4 2 3
```

```
heatmap.2(y, Rowv=as.dendrogram(hr), Colv=as.dendrogram(hc), col=mycol,
  scale="row", density.info="none", trace="none",
  RowSideColors=as.character(kmcol))
```



K-Means Fuzzy Clustering

Performs k-means fuzzy clustering

```
library(cluster)
fanny <- fanny(d, k=4, memb.exp = 1.5)
round(fanny$membership, 2)[1:4,]
```

```
##      [,1] [,2] [,3] [,4]
## g1 1.00 0.00 0.00 0.00
## g2 0.00 0.99 0.00 0.00
## g3 0.02 0.01 0.95 0.03
## g4 0.00 0.00 0.99 0.01
```

```
fanny$clustering
```

```
##  g1  g2  g3  g4  g5  g6  g7  g8  g9  g10
##   1   2   3   3   4   4   4   4   2   3
```

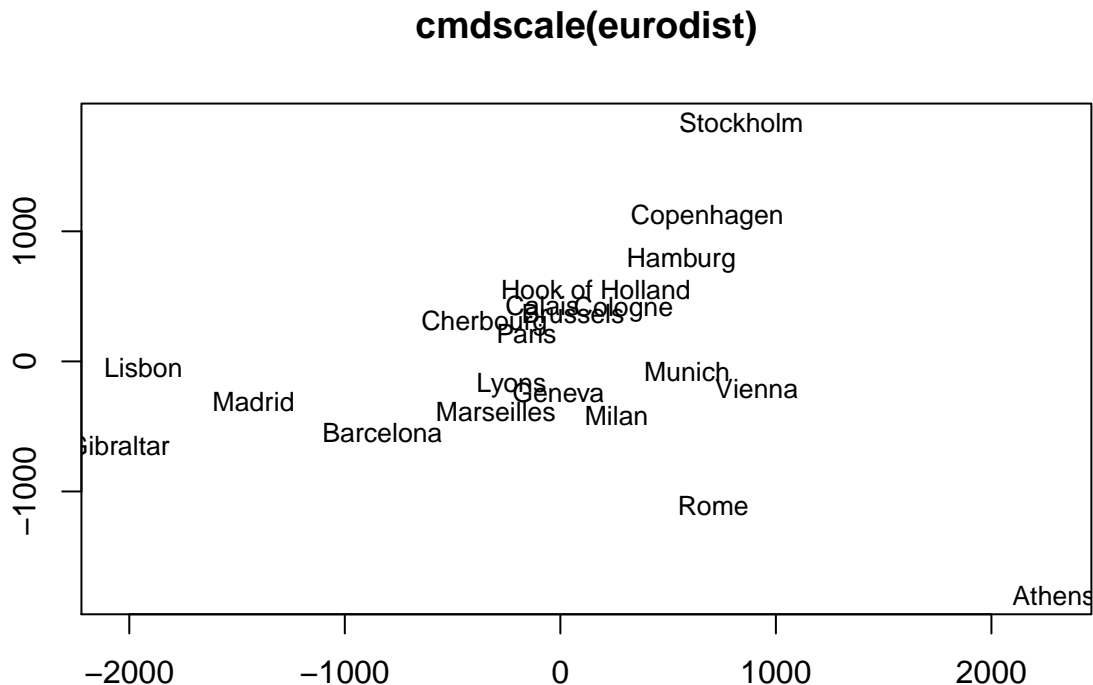
```
## Returns multiple cluster memberships for coefficient above a certain
## value (here >0.1)
fannyMA <- round(fanny$membership, 2) > 0.10
apply(fannyMA, 1, function(x) paste(which(x), collapse="_"))
```

```
##      g1      g2      g3      g4      g5      g6      g7      g8      g9      g10
##      "1"      "2"      "3"      "3"      "4"      "4"      "4"      "2_4"      "2"      "3"
```


Multidimensional Scaling (MDS)

Performs MDS analysis on the geographic distances between European cities

```
loc <- cmdscale(eurodist)
## Plots the MDS results in 2D plot. The minus is required in this example to
## flip the plotting orientation.
plot(loc[,1], -loc[,2], type="n", xlab="", ylab="", main="cmdscale(eurodist)")
text(loc[,1], -loc[,2], rownames(loc), cex=0.8)
```



Principal Component Analysis (PCA)

Performs PCA analysis after scaling the data. It returns a list with class `prcomp` that contains five components: (1) the standard deviations (`sdev`) of the principal components, (2) the matrix of eigenvectors (`rotation`), (3) the principal component data (`x`), (4) the centering (`center`) and (5) scaling (`scale`) used.

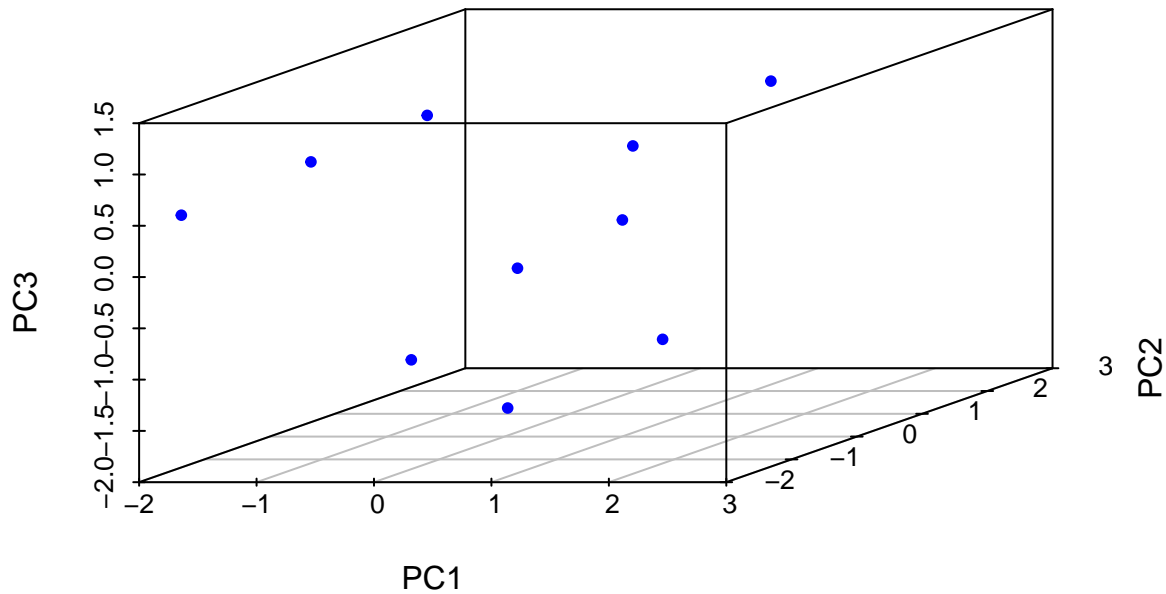
```
library(scatterplot3d)
pca <- prcomp(y, scale=TRUE)
names(pca)
```

```
## [1] "sdev"      "rotation"  "center"    "scale"     "x"
```

```
summary(pca) # Prints variance summary for all principal components.
```

```
## Importance of components%s:
##              PC1    PC2    PC3    PC4    PC5
## Standard deviation  1.3611 1.1777 1.0420 0.69264 0.4416
## Proportion of Variance 0.3705 0.2774 0.2172 0.09595 0.0390
## Cumulative Proportion 0.3705 0.6479 0.8650 0.96100 1.0000
```

```
scatterplot3d(pca$x[,1:3], pch=20, color="blue")
```



Additional Exercises

See here

Version Information

```
sessionInfo()
```

```
## R version 3.4.0 (2017-04-21)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 14.04.5 LTS
##
## Matrix products: default
## BLAS: /usr/lib/libblas/libblas.so.3.0
## LAPACK: /usr/lib/lapack/liblapack.so.3.0
##
## locale:
##  [1] LC_CTYPE=en_US.UTF-8      LC_NUMERIC=C               LC_TIME=en_US.UTF-8
##  [4] LC_COLLATE=en_US.UTF-8    LC_MONETARY=en_US.UTF-8    LC_MESSAGES=en_US.UTF-8
##  [7] LC_PAPER=en_US.UTF-8      LC_NAME=C                  LC_ADDRESS=C
## [10] LC_TELEPHONE=C            LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
##
## attached base packages:
## [1] stats      graphics  utils      datasets  grDevices  base
##
## other attached packages:
## [1] scatterplot3d_0.3-40 RColorBrewer_1.1-2  pheatmap_1.0.8       cluster_2.0.6
## [5] gplots_3.0.1         ape_4.1              ggplot2_2.2.1         BiocStyle_2.4.0
```

```
##
## loaded via a namespace (and not attached):
## [1] Rcpp_0.12.10      knitr_1.15.1      magrittr_1.5      munsell_0.4.3     colorspace_1.3-2
## [6] lattice_0.20-35   stringr_1.2.0     plyr_1.8.4        caTools_1.17.1    tools_3.4.0
## [11] parallel_3.4.0    grid_3.4.0        gtable_0.2.0      nlme_3.1-131      KernSmooth_2.23-15
## [16] gtools_3.5.0      htmltools_0.3.5   yaml_2.1.14       lazyeval_0.2.0    rprojroot_1.2
## [21] digest_0.6.12     tibble_1.3.0      bitops_1.0-6      codetools_0.2-15  evaluate_0.10
## [26] rmarkdown_1.5     gdata_2.17.0      stringi_1.1.5     compiler_3.4.0    methods_3.4.0
## [31] scales_0.4.1      backports_1.0.5
```

References

Hathaway, R J, J C Bezdek, and N R Pal. 1996. “Sequential Competitive Learning and the Fuzzy c-Means Clustering Algorithms.” *Neural Netw.* 9 (5): 787–96. <http://www.hubmed.org/display.cgi?uids=12662563>.