MALDITOFSpectraPA

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MALDITOFSpectraPA is a set of 22 R functions for processing and analysis of MALDI_TOF spectra applied to the taxonomic classification of bacteria. Different functions have been very useful for the development of this package, some coming from multivariate statistical analysis packages and others from spectrum import and processing packages. The functions of this package can be grouped into 6 classes, as shown below.

- **1 Metadata processing:** AgFilter, PMetaData, MergeMetaData, DropCMetadata
- **2 MALDI_TOF Spectrum processing:** CopyRenameXml, XmlNCleaning, SlowXml26Cleaning, Peak_Detector, RowColumnSelector
- **3 Clustering of MALDI_TOF Spectra and cluster validation:** BHclus, Phyclus, PhyclusVar, Vizclus, Dendrogram_pairComp, OptClusters, VisualOptClusters, PointClusterVal
- 4 Correlograms for MALDI_TOF spectra: Visual_CorrDistM
- 5 Principal Component Analysis and clustering of MALDI TOF spectra: PCA Clus, SPCA
- 6 Multidimensional scaling and clustering of MALDI TOF spectra. MDS Clus, SMDS

These functions are organized in the following 8 R scripts.

source("MetaDProcessing.R"), source("DProcessing.R"), source("SProcessing.R"), source("HClus_dendrograms.R"), source("ClusVal.R"), source("VisualDM_Correlograms.R") source("S_PCA_Clus.R"), source("DClus_MS.R")

and depend on the following 18 R packages referred below,

library("MALDIquant"), library("MALDIquantForeign"), library("matrixStats"), library("stringr") library("readxl"), library("(tidyverse"), library("dplyr"), library("readxl"), library("factoextra") library("fpc"), library("cluster"), library("ggplot2"), library("FactoMineR"), library("corrplot") library("gppubr"), library("pvclust"), library("ape"). library("dendextend")

The functions were included in the R package MaldiTOFSpectraPA_0.1.0 (Source and binary packages delivered together with this document).

The development of this small set of functions has two basic objectives: 1) Accelerating as much as possible the processing of spectra and metadata contained in csv, xlsx and mzXML files and directories. 2) developing pipelines that connect these processing functions to a few of the most known R functions for clustering, PCA, Classic Multidimensional Scaling, correlograms and dendrogram building. Most of the arguments of these functions come directly from the functions they use internally, and, in many cases, are the most relevant ones used by these functions. Thus, probably by adjusting only the values of these arguments, the desired graphs and data displays can be obtained. The rest of the arguments of these internal functions take either default values, or values set according to the specific characteristics of these data or the concrete examples taken from the literature.

In this document we describe the general characteristics of each function. This description of each function is structured in four sections. In section @param, the arguments of each function are presented as well as the values they take and the default values. In the sections identified as @return and @examples, as their names indicate, the returned values and concrete examples of use of each function are shown. The last section, called source, contains links to documents that were very useful for the development of the corresponding function. Below the description of each function, some examples of its use are shown in much more detail.

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1. Metadata processing

1.1. AgFilter

AgFilter is an internal function for column selection ("select", default value) or column aggregation based on "means", "medians", "max" and "min" of scores.

1.2. PMetaData

PMetaData extracts from csv files the taxonomic identifications of the isolates and transforms them according to the values of the scores

```
    @param Dirp: folder (and path) where the csv files are located,
    @param fileout: name of the output file containing the Maldi identifiers and species names,
    @param fu: this parameter determines whether only one of the scores (y, default value=2) is considered, or instead, the mean, median, minimum or maximum of the scores corresponding to each of the isolates, "means", "medians", "select", "max" and "min",
    @param id: determine whether unidentified isolates are labeled with "NRI" (default value) or removed,
    @param sc. columns containing the scores
    @param sp. Names: columns containing the names of the identified species
    dataframe (and csv file) with three columns: Maldi identifiers, isolate identifications and the corresponding scores
    @examples Rn<-PMetaData("Bees project 2019-2020"), Rn<-PMetaData("Bees project 2019-2020", fileout="expMeans.csv", fu="means")</li>
    Rn<-PMetaData("Bees project 2019-2020", fileout="expMin.csv", fu="min"), Rn<-PMetaData("Bees project 2019-2020", fileout="expMin.csv", fu="min"), Rn<-PMetaData("Bees project 2019-2020", fileout="expMax.csv", fu="max")</li>
```

1.3. MergeMetaData

It merges two dataframes, one is the output of the function PMetaData1 and the other comes from a xlsx file. The merging is done through the Maldi code.

```
    @param df:a dataframe which is an output of PMetaData1,
    @param Dp: folder containing the metadata files,
    @param filein: xlsx files for metadata
    @param fileout: csv file for both joined dataframes,
    @param keyCode: key code used for Dataframe merging, "Maldi_code" (Default value)
    @return both dataframes joined into one dataframe and exported as an csv file
    @examples dfm<-MergeMetaData(Rn, "Bees project 2019-2020", "Bees metadata.xlsx", "Filered_Meta.csv")</li>
```

1.4. DropCMetadata

DropCMetadata removes previously chosen variables from the metadata csv files

```
@param dm: dataframe,
@param fileout: csv file for both joined dataframes
@return output dataframe and csv file
@examples dfc<-DropCMetadata(dfm,"DCFilered_Meta.csv", c("Date d'analyse","Date de récolte"))</pre>
```

2. MALDI_TOF Spectrum processing

2.1. CopyRenameXml

CopyRenameXml is a function aimed at copying and renaming the mzXML files not being in the folders: "BTS", "BTS_Validation", "CTL", "Autocalibration"

```
@param Dirp: folder containing the mzXML files,
@param newdir: the folder to which the mzXML files are copied
@examples CopyRenameXml("Bees project 2019-2020", "newdata")
```

2.2. XmINCleaning

XmINCleaning copies all mzXML files with at least n lines to a new folder or delete those with less than n lines, (the mzXML files have 26 lines), this way incomplete or truncated files are removed...the number of characters of line n=18 is also verified in order to detect other defective files.

Note: it's recommended to run this function with op="delete" to clean the CopyRenameXml output directory

2.3. SlowXml26Cleaning

SlowXml26Cleaning copies all mzXML files with 26 lines to a new folder or delete those with less than 26 lines, slow **and** old version of XmlNCleaning

2.4. Peak_Detector

Peak_Detector is a pipeline of two stages for 1) importing and processing of Maldi_Tof spectra and 2) peak detection:

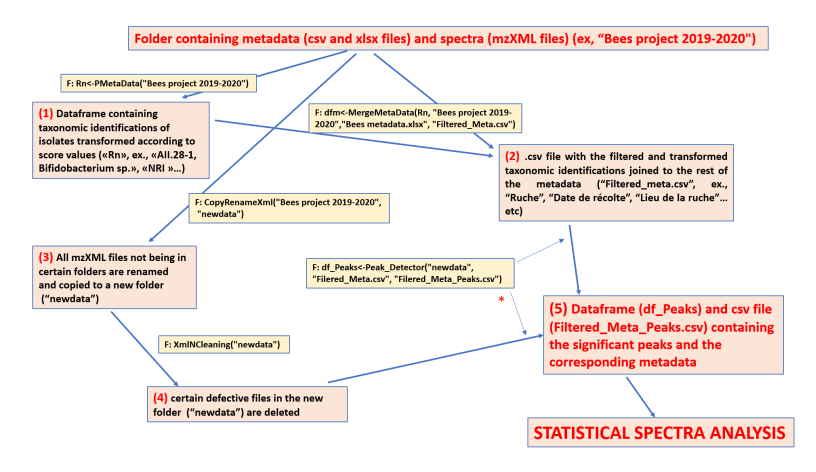
Stage 1) import, check of quality, transformation, smoothing, baseline removing, normalization, and alignment of Maldi_Tof spectra.

Stage 2) peak detection, binning, filtering and merging with metadata.

```
    @param
    percolumn: taxonomic identification of isolates
```

```
@param
           t: spectrum transformation (default value, t="sqrt"), "log",
           smooth: smoothing method (default value smooth="SavitzkyGolay"), "MovingAverage", "WMovingAverage",
@param
           baseline: baseline removing method, (default value, baseline="SNIP"), "TopHat", "ConvexHull",
@param
           normalization: calibration or normalization algorithms (default value, normalization="TIC"), "PQN",
@param
@param
           Iter: number of iterations for baseline removing (default value, Iter=100)
           SN_R: signal_to_noise ratio (default value, SN_R=2),
@param
           minFreq: the minimum peak frequency for spectrum selection (default value, minFreq=0.25),
@param
@param
           align: Boolean parameter to indicate whether or not spectrum alignment is performed (default value, align=TRUE)
@return
           merged dataframes and csv file
@examples df_Peaks<-Detect_Peaks("newdata", "Filered_Meta.csv", "Filered_Meta_Peaks.csv"),
           df_Peaks_0<-Detect_Peaks("newdata", "Filered_Meta.csv", "Filered_Meta_Peaks_0.csv", minFreq=0, align=FALSE)
           df_PeaksTH<-Detect_Peaks("newdata", "Filered_Meta.csv", "Filered_Meta_Peaks.csv", baseline="TopHat")
```

note: Peak_Detector produces the dataframe (and csv file) that serves as starting point for the analysis stage. df_Peaks, (exported as Filtered_Peaks_Meta.csv), see figure below



This figure shows the workflow for importing and processing MALDI_TOF spectra and metadata. The folders, csv files and dataframes generated in each of the stages of this workflow are compressed together with the scripts in the zip files. The df_Peaks dataframe groups all significant spectrum peaks and variables used in the analysis stage. The salmon-colored boxes represent the data (and metadata) and the yellow ones, the functions. *note: Peak_Detector must be executed after folder cleaning with XmlNCleaning (in the example above, the name of this folder is "newdata")

2.5. RowColumnSelector

RowColumnSelector selects rows and columns that meet certain conditions determined by a categorical variable.

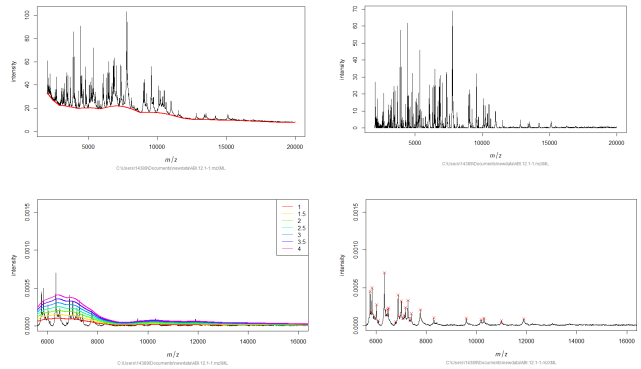
@param df_m: dataframe containing peaks and metadata

@param varCat1: selected categorical variable

@param value: chosen level of varCat1

@param ni,nf: first and last columns corresponding to categorical variables

@examples filtered<-RowColumnSelector(df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus")



This figure shows some of the stages performed by the Peak_Detector pipeline, baseline estimating and deleting, signal-to-noise ratio and significant peak estimates

3. Clustering of MALDI_TOF spectra and cluster validation

3.1. BHclus

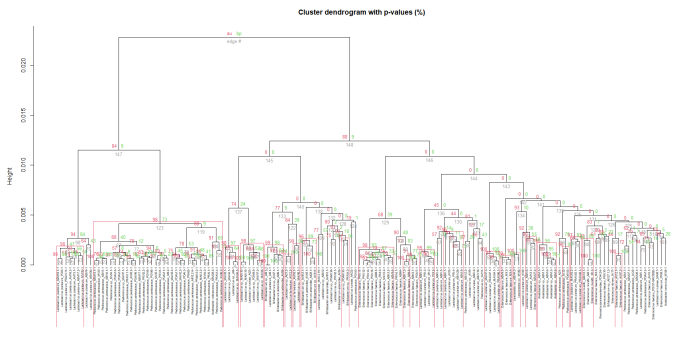
It's a function wrapper of pvclust for building dendrograms labeled with bootstrap probabilities for isolates chosen according to a categorical variable

```
@param
            df m: dataframe containing peaks and metadata
            meth: hierarchical clustering algorithm ("ward2", default), other values: "average", "ward.D", "single", "complete",
@param
           "mcquitty", "median" or "centroid"
           dist: distance ("euclidean", defaul), "maximum", "manhattan", "canberra", "binary" "correlation", "uncentered",
@param
                  "abscor"
            varCat1: categorical variable, "Genre", "Taxonomie", "Nutrition", "Ruche"...
@param
            value: level of catVar1, examples: "Lactobacillus" ("Genre"), Taxonomie ("Pediococcus pentosaceus"), "Erica cinerea"
@param
                 ("Nutrition"),...
           nb: number of bootstrap iterations (nb=100, default value)
@param
@param
            fig: boolean variable to indicate output figure
@return
            output cluster and figure
@examples dft<-BHclus(df_Peaks, varCat1="Genre", value="All", nb=500), dft<-BHclus(df_Peaks, varCat1="Genre",
              value="Lactobacillus", nb=500)
              dft<-BHclus(df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus"),
              dft<-BHclus(df_Peaks, meth="complete", dist="canberra",varCat1="Taxonomie", value="Pediococcus
              pentosaceus")
```

source: https://academic.oup.com/bioinformatics/article/22/12/1540/207339 https://www.rdocumentation.org/packages/pvclust/versions/2.2-0/topics/pvclust https://www.rdocumentation.org/packages/stats/versions/3.2.1/topics/dist

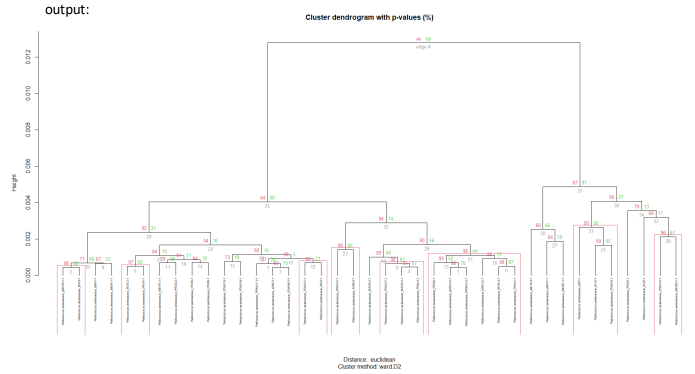
dft<-BHclus(df_Peaks, varCat1="Genre", value="All", nb=500)

output:



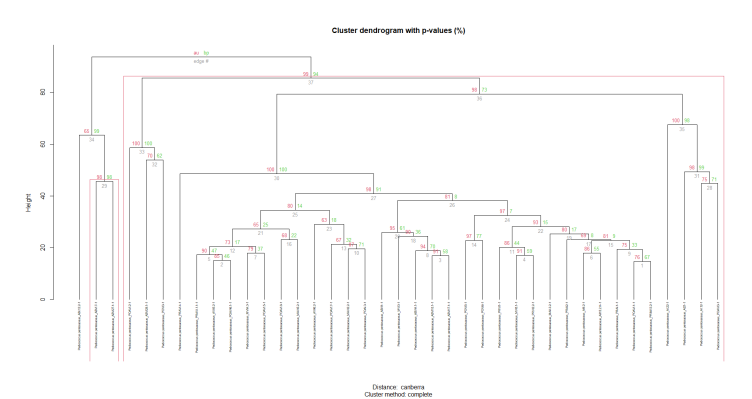
Distance: euclidean

dft<-BHclus(df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus")



dft<-BHclus(df_Peaks, meth="complete", dist="canberra",varCat1="Taxonomie", value="Pediococcus pentosaceus")

output:



3.2. Phyclus

Phyclus builds different types of dendrograms for isolates chosen according to a categorical variable

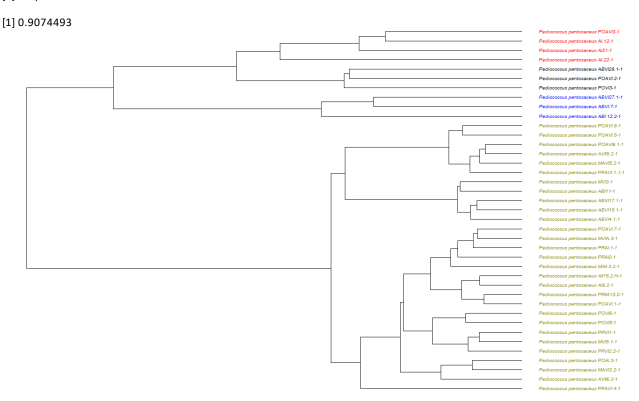
```
@param
           df_m: dataframe containing peaks and metadata
@param
           meth: hierarchical clustering algorithm ("ward2", default), other values: "average", "ward.D", "single", "complete",
                 "mcquitty", "median" or "centroid"
@param
           dist: distance ("euclidean", defaul), "euclidean", "maximum", "manhattan", "canberra", "binary" "minkowski"
           varCat1: categorical variable, example: "Genre", "Taxonomie", "Nutrition", "Ruche"...
@param
@param
           value: level of catVar1, example: "Lactobacillus" ("Genre"), Taxonomie ("Pediococcus pentosaceus"), "Erica cinerea"
           ("Nutrition"),...
@param
           nc: number of clusters (nc=4, default value)
           dendrogram: dendrogram type ("phylogram", default value), "cladogram", "unrooted", "fan" and "radial"
@param
@return
           dataframe and figure
@examples c<-Phyclus(df_Peaks, varCat1="Taxonomie",value="Pediococcus pentosaceus"), c<-Phyclus(df_Peaks,
            varCat1="Nutrition", value="Erica cinerea")
            c<-Phyclus(df Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus", dendogram="cladogram")
```

source: https://www.rdocumentation.org/packages/stats/versions/3.6.2/topics/hclust https://www.rdocumentation.org/packages/ape/versions/5.4-1

c<-Phyclus(df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus")

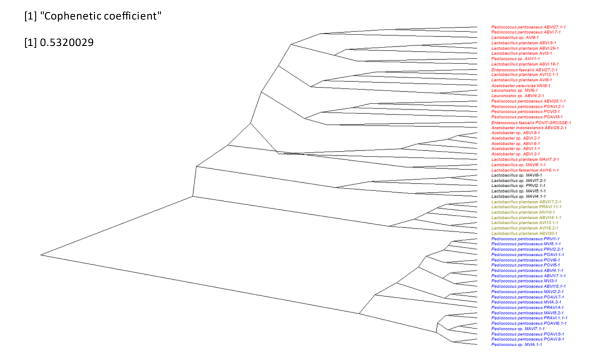
output:

[1] "Cophenetic coefficient"



c<-Phyclus(df_Peaks, varCat1="Ruche",value="VI", dendrogram="cladogram")

output:

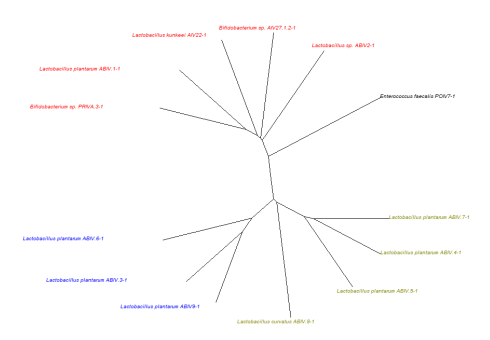


c<-Phyclus(df_Peaks, varCat1="Ruche", value="IV", dendrogram="unrooted")

output:

[1] "Cophenetic coefficient"



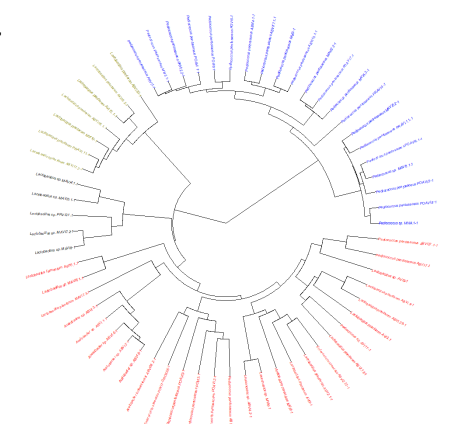


$c < -Phyclus (df_Peaks, varCat1 = "Nutrition", value = "Erica cinerea", dendrogram = "fan")$

output

[1] "Cophenetic coefficient"

[1] 0.5320029



c<-Phyclus(df_Peaks, dendrogram="fan")

meth="complete",

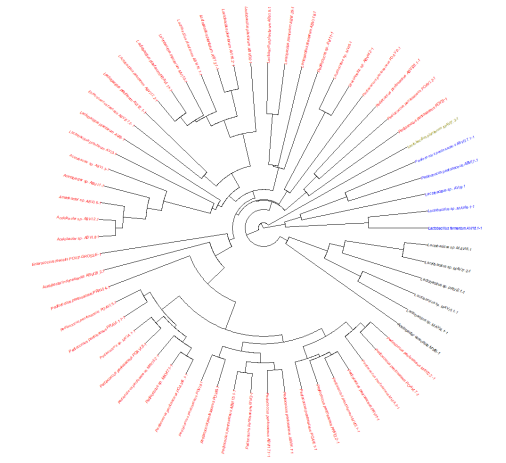
varCat1="Nutrition",value="Erica

cinerea"

output:

[1] "Cophenetic coefficient"

[1] 0.9295954



3.3. PhyclusVar

Phyclus Var builds different types of dendrograms for isolates selected and categorized based on varCat2 and varCat1 levels

```
@param
            df_m: dataframe containing peaks and metadata
@param
            meth: hierarchical clustering algorithm ("ward2", default value), other values: "average", "ward.D", "single", "complete",
                  "mcquitty", "median" or "centroid"
            dist: distance ("euclidean", default value), "euclidean", "maximum", "manhattan", "canberra", "binary" "minkowski"
@param
            varCat1: categorical variable, examples: "Taxonomie", "Genre", "Date.d.analyse", "Origine", "Ruche", "Nutrition",
@param
                   "Date.de.récolte", "Lieu.de.la.ruche"
            value: level of catVar1, examples: "Lactobacillus" ("Genre"), Taxonomie("Pediococcus pentosaceus"), "Erica cinerea"
@param
            ("Nutrition"),...
@param
           varCat2: categorical variable
@param
            nc: number of clusters (nc=4, default value)
@param
            dendrogram: dendrogram type ("phylogram", default value), "cladogram", "unrooted", "fan" and "radial"
@return
            output cophenetic coefficient and figure
@examples dft<-PhyclusVar(df_Peaks, varCat1="Genre", value="Lactobacillus", varCat2="Nutrition"),
             dft<-PhyclusVar(df Peaks, varCat1="Taxonomie", value="Lactobacillus plantarum", varCat2="Ruche")
             dft<-PhyclusVar(df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus", varCat2="Nutrition",
             dendrogram="cladogram"),
             dft<-PhyclusVar(df_Peaks, varCat1="Taxonomie", value="All", varCat2="Date.de.récolte", dendrogram="fan")
             dft<-PhyclusVar(df_Peaks, varCat1="Taxonomie", value="Lactobacillus plantarum", varCat2="Ruche", dendrogram="fan"),
             dft<-PhyclusVar(df_Peaks, varCat1="Taxonomie", value="Lactobacillus plantarum", varCat2="Ruche",
             dendrogram="unrooted")
```

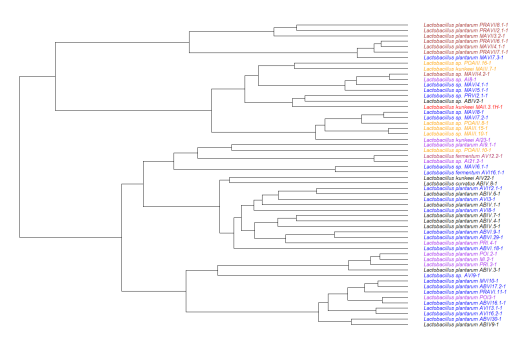
source: https://www.rdocumentation.org/packages/stats/versions/3.6.2/topics/hclust https://www.rdocumentation.org/packages/ape/versions/5.4-1

dft<-PhyclusVar(df Peaks, varCat1="Genre", value="Lactobacillus", varCat2="Nutrition")

output:

[1] "Cophenetic coefficient"

[1] 0.6082436

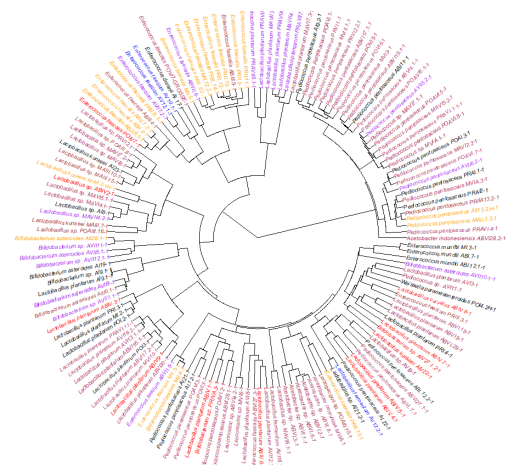


dft<-PhyclusVar(df_Peaks, varCat1="Taxonomie", value="All", varCat2="Date.de.récolte", dendrogram="fan")

output

[1] "Cophenetic coefficient"

[1] 0.39633



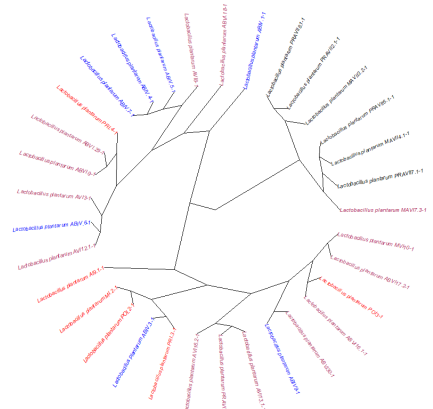
 $dft < -Phyclus Var (df_Peaks, var Cat 1 = "Taxonomie", value = "Lactobac illus plantarum", var Cat 2 = "Ruche", var Cat 2 = "Ruche",$

dendrogram="radial")

output:

[1] "Cophenetic coefficient"

[1] 0.7178863



3.4. Vizclus

Vizclus computes clustering statistics and has other options for dendrogram visualization

```
df_m: dataframe containing peaks and metadata
@param
            meth: hierarchical clustering algorithm ("ward2", default value), other values: "average", "ward.D", "single",
                  "complete", "mcquitty", "median" or "centroid"
@param
            dist: distance ("euclidean", default value), "euclidean", "maximum", "manhattan", "canberra", "binary" "minkowski"
@param
           varCat1: categorical variable for choosing isolates, examples: "Taxonomie", "Genre",
                   "Date.d.analyse", "Origine", "Ruche", "Nutrition", "Date.de.récolte", "Lieu.de.la.ruche"
@param
            value: level of catVar1, examples: "Lactobacillus" ("Genre"), Taxonomie ("Pediococcus pentosaceus"), "Erica cinerea"
                 ("Nutrition"),...
@param
            nc: number of clusters (nc=4, default value)
@param
            dendrogram: dendrogram and factor map
            output figures and statistics
@return
@examples Lt<-Vizclus(df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus"),
            Lt<-Vizclus(df_Peaks,varCat1="Ruche", value="VI")
            Lt<-Vizclus(df_Peaks,varCat1="Nutrition", value="Caraway", nc=2),
           Lt<-Vizclus(df_Peaks,varCat1="Genre", value="Lactobacillus", nc=4)
            Lt<-Vizclus(df_Peaks,varCat1="Nutrition", value="All"),
           Lt<-Vizclus(df_Peaks,varCat1="Nutrition", value="Caraway", nc=3, graph="fm")
   source: https://www.rdocumentation.org/packages/stats/versions/3.6.2/topics/hclust
           https://www.rdocumentation.org/packages/factoextra/versions/1.0.7/topics/fviz
           http://www.sthda.com/english/wiki/beautiful-dendrogram-visualizations-in-r-5-must-known-methods-
           unsupervised-machine-learning
```

Lt<-Vizclus(df Peaks,varCat1="Genre", value="Lactobacillus", nc=4)

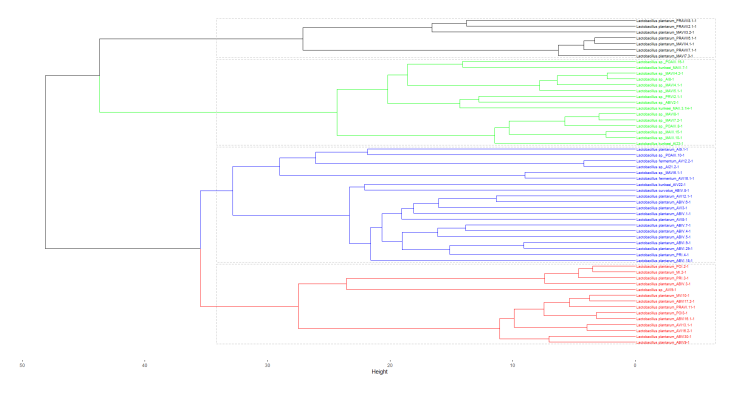
Output:

```
[1] "clustering vector: cluster assignment to each isolate"
 Lactobacillus plantarum_ABIV.1-1 Lactobacillus plantarum_ABIV.3-1 Lactobacillus plantarum_ABIV.4-1
                 1
                                     2
                                                        1
 Lactobacillus plantarum_ABIV.5-1 Lactobacillus plantarum_ABIV.6-1 Lactobacillus plantarum_ABIV.7-1
                 1
                                     1
                                                        1
  Lactobacillus curvatus_ABIV.8-1 Lactobacillus plantarum_ABVI.18-1 Lactobacillus plantarum_ABVI.29-1
                 1
                                     1
                                                        1
 Lactobacillus plantarum_ABVI.9-1 Lactobacillus kunkeei_MAII.3.1H-1
                                                                       Lactobacillus sp._MAIII.10-1
   Lactobacillus sp._MAIII.15-1 Lactobacillus kunkeei_MAIII.7-1 Lactobacillus plantarum_MI.2-1
                 3
                                     3
                                                        2
   Lactobacillus sp._POAIII.10-1 Lactobacillus sp._POAIII.16-1
                                                                Lactobacillus sp. POAIII.8-1
                                     3
  Lactobacillus plantarum_POI.2-1 Lactobacillus plantarum_PRAVI.11-1 Lactobacillus plantarum_PRI.3-1
                 2
                                     2
  Lactobacillus plantarum_PRI.4-1
                                     Lactobacillus sp._AI21.2-1 Lactobacillus fermentum_AV12.2-1
                 1
                                     1
                                                        1
 Lactobacillus plantarum_AVI12.1-1 Lactobacillus fermentum_AVI16.1-1 Lactobacillus plantarum_AVI16.2-1
                 1
                                     1
                                                        2
  Lactobacillus plantarum_AVI3-1
                                   Lactobacillus plantarum_AVI8-1
                                                                       Lactobacillus sp._AVI9-1
                 1
                                     1
```

.....shortened output.....

```
[1] "The size of each cluster" cl
1 2 3 4
20 14 15 7
[1] "Cophenetic coeficient"
```

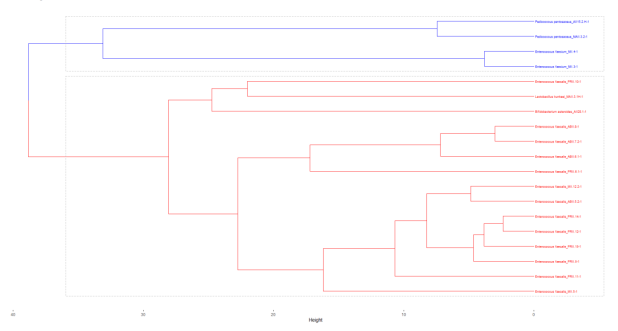
Cluster Dendrogram



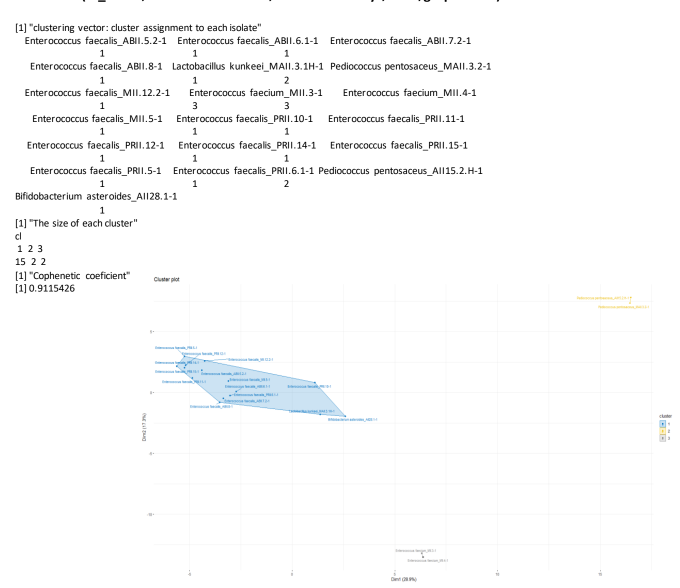
Lt<-Vizclus(df_Peaks,varCat1="Nutrition", value="Caraway", nc=2)

Output:

```
[1] "clustering vector: cluster assignment to each isolate"
  Enterococcus faecalis_ABII.5.2-1 Enterococcus faecalis_ABII.6.1-1 Enterococcus faecalis_ABII.7.2-1
                                      1
                                                          1
   Enterococcus faecalis_ABII.8-1 Lactobacillus kunkeei_MAII.3.1H-1 Pediococcus pentosaceus_MAII.3.2-1
                                      1
  Enterococcus faecalis_MII.12.2-1
                                      Enterococcus faecium_MII.3-1
                                                                       Enterococcus faecium_MII.4-1
   Enterococcus faecalis_MII.5-1
                                   Enterococcus faecalis_PRII.10-1 Enterococcus faecalis_PRII.11-1
  Enterococcus faecalis_PRII.12-1
                                   Enterococcus faecalis_PRII.14-1 Enterococcus faecalis_PRII.15-1
                  1
                                      1
                                                          1
   Enterococcus faecalis_PRII.5-1 Enterococcus faecalis_PRII.6.1-1 Pediococcus pentosaceus_AII15.2.H-1
                  1
                                                          2
Bifidobacterium asteroides_AII28.1-1
[1] "The size of each cluster"
cl
1 2
15 4
[1] "Cophenetic coeficient"
[1] 0.9115426
```



Lt<-Vizclus(df_Peaks,varCat1="Nutrition", value="Caraway", nc=3,graph="fm")



Lt<-Vizclus(df_Peaks,varCat1="Nutrition", value="All")

Output:

```
[1] "clustering vector: cluster assignment to each isolate"
   Pediococcus pentosaceus AI.12-1
                                        Enterococcus durans AI.17-1
                                                                       Pediococcus pentosaceus AI.22-1
   Enterococcus mundtii_ABI.12.1-1 Pediococcus pentosaceus_ABI.12.2-1
                                                                            Enterococcus mundtii_ABI.7-1
                                       1
                                                           1
  Enterococcus faecalis_ABII.5.2-1
                                   Enterococcus faecalis_ABII.6.1-1
                                                                     Enterococcus faecalis_ABII.7.2-1
                                                           2
   Enterococcus faecalis_ABII.8-1
                                   Enterococcus faecalis_ABIII.5-1 Lactobacillus plantarum_ABIV.1-1
                                                           1
  Lactobacillus plantarum_ABIV.3-1
                                     Lactobacillus plantarum_ABIV.4-1
                                                                       Lactobacillus plantarum_ABIV.5-1
                  1
                                       1
                                                           1
  Lactobacillus plantarum_ABIV.6-1
                                     Lactobacillus plantarum_ABIV.7-1
                                                                        Lactobacillus curvatus_ABIV.8-1
                                       1
                                                           1
      Acetobacter sp._ABVI.1-1 Lactobacillus plantarum_ABVI.18-1
                                                                         Acetobacter sp._ABVI.2-1
                  1
  Lactobacillus plantarum_ABVI.29-1
                                          Acetobacter sp._ABVI.3-1
                                                                         Acetobacter sp._ABVI.6-1
                  1
                                                           1
  Pediococcus pentosaceus_ABVI.7-1
                                           Acetobacter sp._ABVI.8-1
                                                                     Lactobacillus plantarum_ABVI.9-1
                  1
   Enterococcus faecium_ABVII.5-1
                                     Enterococcus faecalis_ABVII.6-1
                                                                     Lactobacillus kunkeei_MAII.3.1H-1
                  1
 Pediococcus pentosaceus_MAII.3.2-1
                                         Lactobacillus sp._MAIII.10-1
                                                                         Lactobacillus sp._MAIII.15-1
   Lactobacillus kunkeei_MAIII.7-1
                                     Lactobacillus plantarum_MI.2-1
                                                                        Enterococcus mundtii_MI.3-1
                  3
                                      1
  Enterococcus faecalis_MII.12.2-1
                                      Enterococcus faecium_MII.3-1
                                                                        Enterococcus faecium_MII.4-1
                  2
    Enterococcus faecalis_MII.5-1
                                       Pediococcus sp._MVIA.1-1 Pediococcus pentosaceus_MVIA.3-1
            .....shortened output.....
[1] "The size of each cluster"
1 2 3 4
```

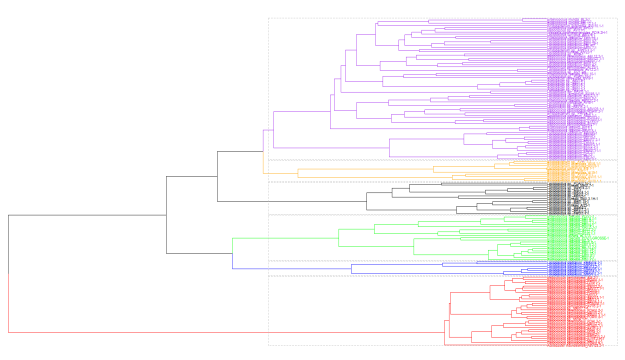
cl

75 28 15 32

[1] "Cophenetic coeficient"

[1] 0.39633

Cluster Dendrogram



3.5. Dendrogram_pairComp

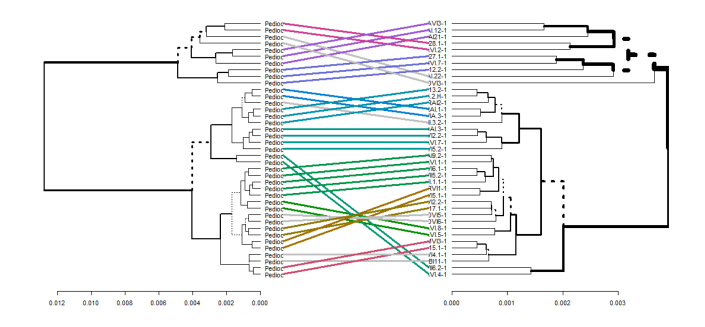
Dendrogram_pairComp computes pairwise dendrogram alignments and correlations

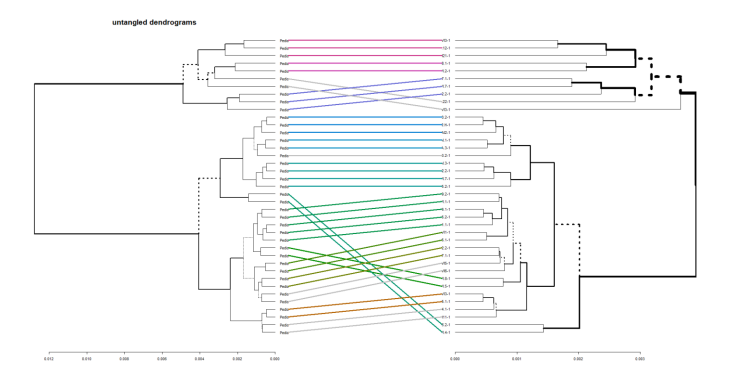
```
@param
           df m: dataframe containing peaks and metadata
@param
           methCorr: dendrogram correlation method ("cophenetic", default), others: "baker", "common nodes", "FM index"
                     meth1, meth2...meth6: hierarchical clustering algorithms "ward2", "average", "ward.D", "single",
@param
                     "complete", "mcquitty", "median" or "centroid"
           dist1, dist2, ...dist6: distances "euclidean", "maximum", "manhattan", "canberra", "binary" "minkowski"
@param
@param
           varCat1: categorical variable for choosing isolates, examples: "Taxonomie",
           "Genre", "Date.d.analyse", "Origine", "Ruche", "Nutrition", "Date.de.récolte", "Lieu.de.la.ruche"
           value: value of catVar1, examples: "Lactobacillus" ("Genre"), Taxonomie("Pediococcus pentosaceus"), "Erica cinerea"
@param
                  ("Nutrition"),...
@return
           untangled and tangled dendrograms, dendrogram correlation matrix and statistics
@examples mc<-Dendrogram pairComp(df Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus"),
           mc<-Dendrogram pairComp(df Peaks, varCat1="Nutrition", value="Marrubium vulgare")
   source: https://cran.r-project.org/web/packages/dendextend/vignettes/dendextend.html#:~:text=The%20dendextend
          %20package%20offers%20a,its%20branches%2C%20nodes%20and%20labels.
          https://www.rdocumentation.org/packages/dendextend/versions/1.14.0
          https://www.r-graph-gallery.com/340-custom-your-dendrogram-with-dendextend.html
          https://academic.oup.com/bioinformatics/article/31/22/3718/240978
          https://cran.rstudio.com/web/packages/dendextend/vignettes/Cluster Analysis.html
          https://www.datanovia.com/en/lessons/comparing-cluster-dendrograms-in-r/
          https://rdrr.io/cran/dendextend/man/untangle.html
```

mc<-Dendrogram_pairComp(df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus")

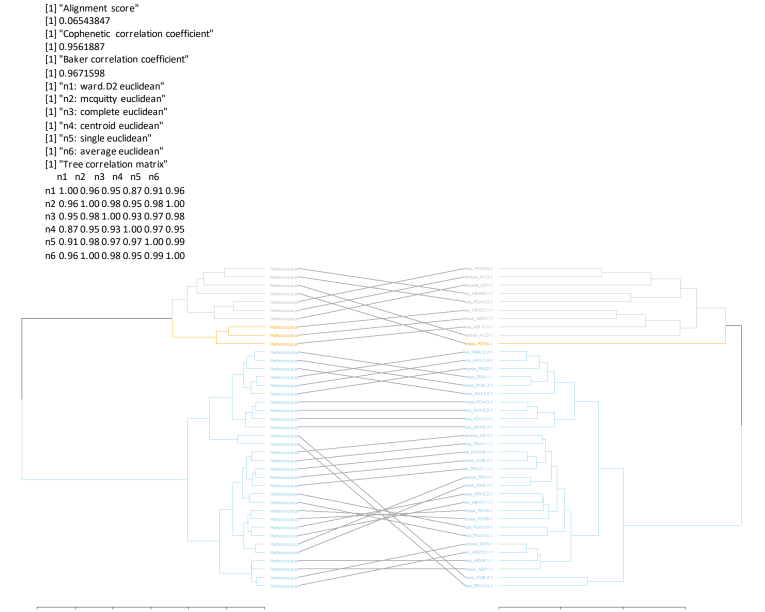
output:

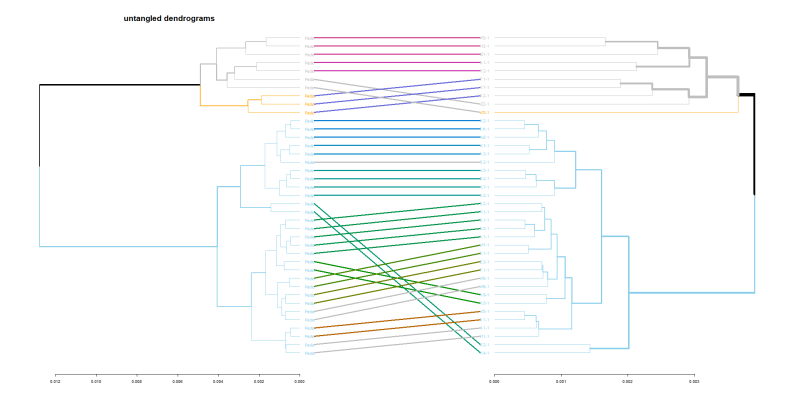
[1] "Alignment score" [1] 0.06543847 [1] "Cophenetic correlation coefficient" [1] 0.9561887 [1] "Baker correlation coefficient" [1] 0.9671598 [1] "n1: ward.D2 euclidean" [1] "n2: mcquitty euclidean" [1] "n3: complete euclidean" [1] "n4: centroid euclidean" [1] "n5: single euclidean" [1] "n6: average euclidean" [1] "Tree correlation matrix" n1 n2 n3 n4 n5 n6 n1 1.00 0.96 0.95 0.87 0.91 0.96 n2 0.96 1.00 0.98 0.95 0.98 1.00 n3 0.95 0.98 1.00 0.93 0.97 0.98 n4 0.87 0.95 0.93 1.00 0.97 0.95 n5 0.91 0.98 0.97 0.97 1.00 0.99 n6 0.96 1.00 0.98 0.95 0.99 1.00





mc<-Dendrogram_pairComp(df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus", graph="ClusterDends")





3.6. OptClusters

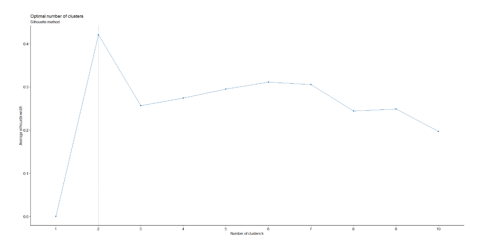
OptClusters, is a wrapper of several functions to visualize and compute optimal clusters for different clustering and evaluation methods

```
@param
           df_m: dataframe containing peaks and metadata
@param
           meth: clustering algorithms "kmeans", (default value), other values: "pam" or "hclust"
           dist: distances "euclidean", (default value), "maximum", "manhattan", "canberra", "binary" "minkowski"
@param
           varCat1: categorical variable for choosing isolates, examples: "Taxonomie", "Genre", "Date.d.analyse"
@param
                  ,"Origine","Ruche", "Nutrition", "Date.de.récolte", "Lieu.de.la.ruche"
            value: level of catVar1, examples: "Lactobacillus" ("Genre"), Taxonomie ("Pediococcus pentosaceus"), "Erica cinerea"
@param
               ("Nutrition"),...
           minc: minimal number of clusters (minc=2, default value)
@param
@param
           maxc: maximal number of clusters (maxc=10, default value)
@param
           ind: methods to evaluate clustering algorithms: "total within sum of squares", "average silhouette width" and "gap
            statistics"
@param
           nb: number of bootstrap samples (nb=100, default value)
@return
           figures and statistics
@examples OptClusters(df_Peaks, varCat1="Taxonomie", value="Enterococcus faecalis"),
           OptClusters(df_Peaks, varCat1="Taxonomie", value="All")
           OptClusters(df Peaks, meth="pam", varCat1="Taxonomie", value="All", ind="gap statistics"),
           OptClusters(df Peaks, meth="hclust", varCat1="Taxonomie", value="All", ind="gap statistics")
  source: https://rpubs.com/pg2000in/OptimumClusters
          http://rstudio-pubs-static.s3.amazonaws.com/265632 3ad9e0b981244e15887677f8dffb39a0.html#
          using-30-different-indices
          https://www.datanovia.com/en/lessons/determining-the-optimal-number-of-clusters-3-must-know-methods/
```

https://www.rdocumentation.org/packages/factoextra/versions/1.0.7/topics/fviz nbclust

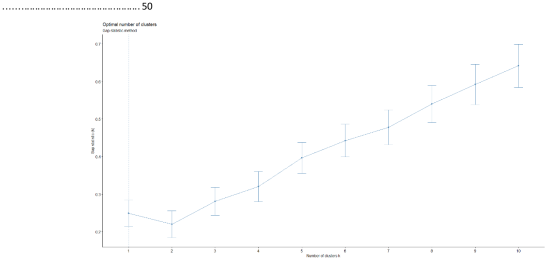
$OptClusters (df_Peaks, varCat1="Taxonomie", \ value="Enterococcus faecalis")$

Output:



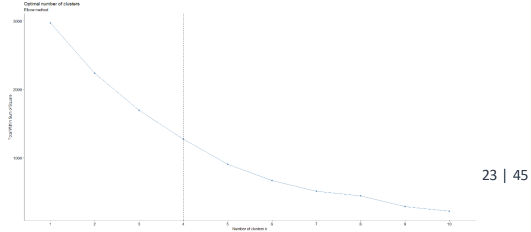
OptClusters(df_Peaks, varCat1="Taxonomie", value="Enterococcus faecalis", ind="gap statistics") Output:

Clustering k = 1,2,..., K.max (= 10): .. done Bootstrapping, b = 1,2,..., B (= 50) [one "." per sample]:



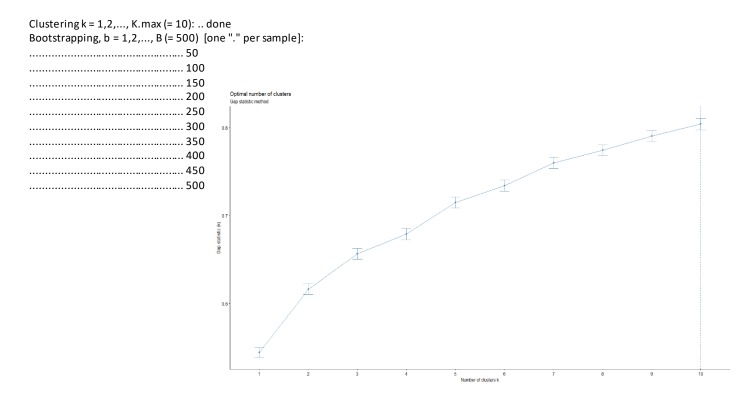
$OptClusters \ (df_Peaks, varCat1="Taxonomie", value="Enterococcus faecalis", ind="total within sum of squares")$

Output:



OptClusters(df_Peaks, meth="pam", varCat1="Taxonomie", value="All", ind="gap statistics", nb=500)

Output:



3.7. VisualOptClusters

visualization and statistics for MALDI_TOF spectra clustering validation

```
@param
            df_m: dataframe containing peaks and metadata
@param
            meth: clustering algorithms "hclust", (default value), other values: "kmeans", "pam", "clara", "fanny",
                  "hclust", "agnes", "diana"
@param
            dist: distances "euclidean", (default value), "maximum", "manhattan", "canberra", "binary"
                 "minkowski"
            meth2: hc methods "ward.D2", (default value), average", "ward.D", "single", "complete", "mcquitty",
@param
                  "median" or "centroid"
@param
            varCat1: categorical variable for choosing isolates, examples: "Taxonomie", "Genre",
                    "Date.d.analyse", "Origine", "Ruche", "Nutrition", "Date.de.récolte", "Lieu.de.la.ruche"
            value: level of catVar1, examples: "Lactobacillus" ("Genre"), Taxonomie ("Pediococcus
@param
                   pentosaceus"), "Erica cinerea" ("Nutrition"),...
@param
            nc: number of clusters, nc=3, (default value)
@return
             figures and statistics
@examples
               s<-VisualOptClusters(df_Peaks, varCat1="Genre", value="Enterococcus"),
               s<-VisualOptClusters(df_Peaks, meth="pam", varCat1="Genre", value="Lactobacillus", nc=2)
               s<-VisualOptClusters(df_Peaks, meth="hclust", meth2="average", dist="pearson",
                varCat1="Genre", value="All", nc=5),
               s<-VisualOptClusters(df_Peaks, meth="kmeans", dist="euclidean", varCat1="Genre",
```

value="AII", nc=3)

source: https://www.rdocumentation.org/packages/factoextra/versions/1.0.7/topics/fviz_cluster https://afit-r.github.io/kmeans_clustering http://rstudio-pubs-static.s3.amazonaws.com/ 265632_3ad9e0b981244e15887677f8dffb39a0.html#using-30-different-indices

s<-VisualOptClusters(df_Peaks, varCat1="Genre", value="Enterococcus")

output:

\$widths

	cluster neighbor	sil_v	vidth					
Enterococcus	faecalis_ABIII.5-1	1	3 0.46934828					
Enterococcus	faecalis_PRII.14-1	1	3 0.46005835					
Enterococcus	faecalis_PRII.12-1	1	3 0.45779439					
Enterococcus	faecalis_PRII.5-1	1	3 0.43660605					
Enterococcus	faecalis_PRII.15-1	1	3 0.43493818					
Enterococcus	faecalis_ABVII.6-1	1	3 0.42491011					
Enterococcus	faecalis_ABII.5.2-1	1	3 0.42264275					
Enterococcus	faecalis_MII.12.2-1	1	3 0.38646783					
Enterococcus	faecalis_PRII.11-1	1	3 0.35187800					
Enterococcus	faecalis_ABII.6.1-1	1	3 0.33850791					
Enterococcus	faecalis_ABII.7.2-1	1	3 0.32897691					
Enterococcus	faecalis_ABII.8-1	1	3 0.31944142					
Enterococcus	faecalis_POIV7-1	1	3 0.31295360					
Enterococcus	faecalis_ABIII4-1	1	3 0.29358295					
Enterococcus	faecalis_MII.5-1	1	3 0.24999632					
Enterococcus	faecalis_PRII.6.1-1	1	3 0.23297577					
Enterococcus	faecium_AVII7.2-1	1	3 0.19485022					
Enterococcus	faecium_AV10.1-1	1	3 0.18642428					
Enterococcus	faecium_AV12.1-1	1	3 0.17833360					
shortened output								

\$clus.avg.widths

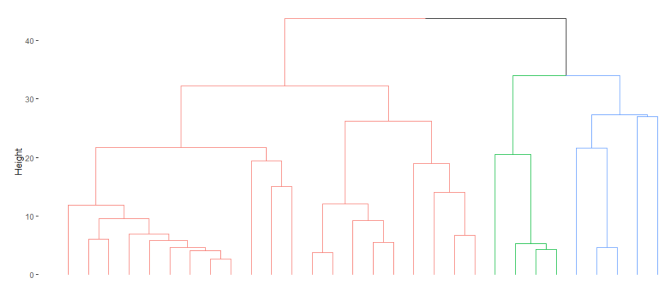
[1] 0.32418331 0.03065726 0.45669725

\$avg.width [1] 0.2929308

Clusters allhouette plot.
Average allhouette width: 0.29

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Cluster Dendrogram



s<-VisualOptClusters(df_Peaks, meth="pam", varCat1="Genre", value="All", nc=6)

output:

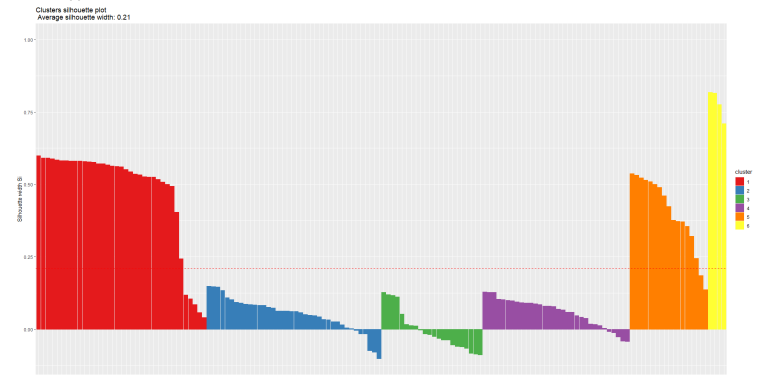
cluster size ave.sil.width 1 1 37 0.48 2 2 38 0.05 3 22 0.00 4 4 32 0.06 5 5 17 0.40 6 4 0.78

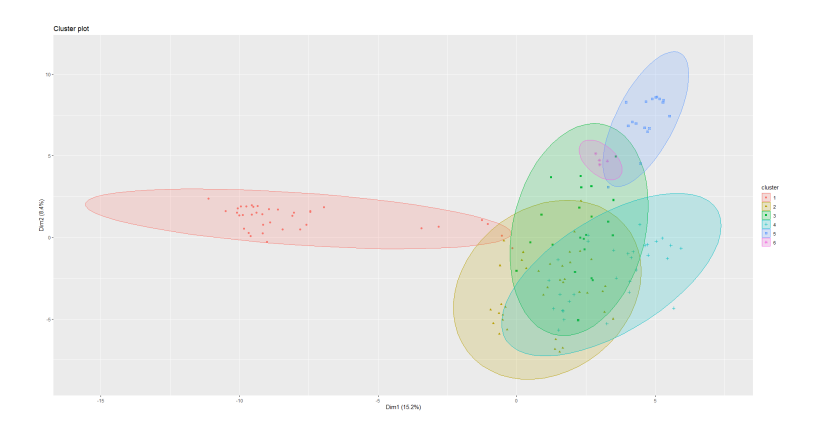
\$widths					
	cluster neighbor	sil_width			
Pediococcus	pentosaceus_POAVI.1-1	1	2 0.599407924		
Pediococcus	pentosaceus_PRAVI.1.1-1	1	2 0.591676671		
Pediococcus	pentosaceus_PRAI.1-1	1	2 0.591085652		
Pediococcus	pentosaceus_AVII5.2-1	1	2 0.588123484		
Pediococcus	pentosaceus_PRVI2.2-1	1	2 0.584962734		
Pediococcus	pentosaceus_POAVI.7-1	1	2 0.582671602		
Pediococcus	pentosaceus_PRIIA13.2-1	1	2 0.582162559		
Pediococcus	pentosaceus_MVIA.3-1	1	2 0.580898030		
Pediococcus	pentosaceus_POAVI.5-1	1	2 0.580663177		
Pediococcus	pentosaceus_AI9.2-1	1	2 0.580299720		
Pediococcus	pentosaceus_PRVI1-1	1	2 0.579086415		
Pediococcus	pentosaceus_MAVI2.2-1	1	2 0.577626216		
Pediococcus	pentosaceus_MVI5.1-1	1	2 0.577150065		
Pediococcus	pentosaceus_POVI5-1	1	2 0.572156201		
Pediococcus	pentosaceus_ABVI15.1-1	1	2 0.571275241		
Pediococcus	pentosaceus_AII15.2.H-1	1	2 0.567036446		
Pediococcus	$pentosaceus_POAVI6.1-1$	1	2 0.564319053		
Pediococcus	pentosaceus_POAI.3-1	1	2 0.562481769		
Pediococcus	pentosaceus_ABVI17.1-1	1	2 0.560626245		
Pediococcus	pentosaceus_PRAI2-1	1	2 0.552265722		
Pediococcus	pentosaceus_MVI3-1	1	2 0.544002379		
Pediococcus	pentosaceus_POVI6-1	1	2 0.535424547		
Pediococcus	pentosaceus_MAVI5.2-1	1	2 0.533393236		
Pediococcus	pentosaceus_ABVI4.1-1	1	2 0.526594660		

.....shortened output.....

\$clus.avg.widths

 $[1] \ 0.481352515 \ 0.051162303 \ -0.004511589 \ 0.058257924 \ 0.403354044 \ 0.779722190$





s<-VisualOptClusters(df_Peaks, meth="pam", varCat1="Genre", value="All", nc=3)

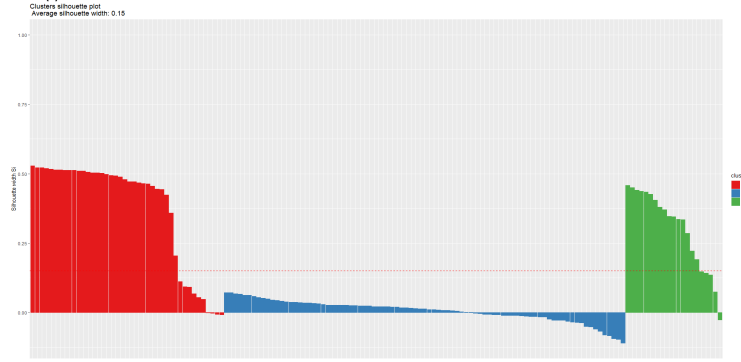
output:

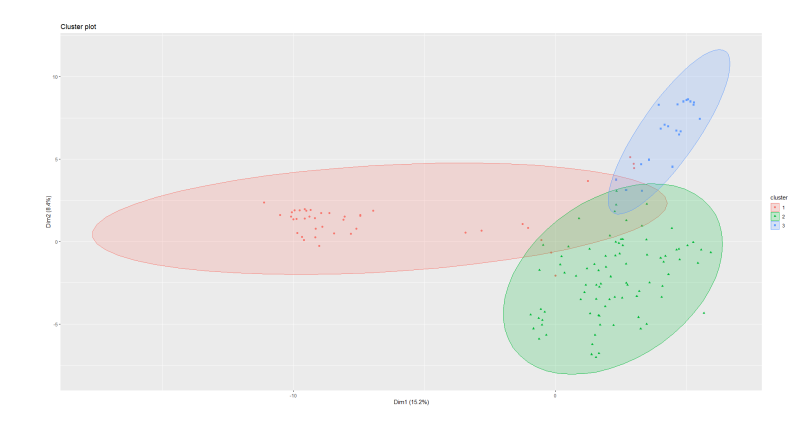
cluster neighbor sil_width Pediococcus pentosaceus_POAVI.1-1 1 2 0.5282162251 Pediococcus pentosaceus_PRAVI.1.1-1 2 0.5217488452 1 Pediococcus pentosaceus_PRAI.1-1
Pediococcus pentosaceus_AVII5.2-1 2 0.5216367308 1 3 0.5188998087 Pediococcus pentosaceus_PRVI2.2-1 1 2 0.5170117138 Pediococcus pentosaceus_PRIIA13.2-1 1 2 0.5147840466 Pediococcus pentosaceus_MVIA.3-1 1 2 0.5137295014 Pediococcus pentosaceus_POAVI.7-1 1 2 0.5133880537 Pediococcus pentosaceus_AI9.2-1 1 3 0.5128952319 Pediococcus pentosaceus_POAVI.5-1 1 2 0.5122735714 Pediococcus pentosaceus_PRVI1-1 1 2 0.5106237133 Pediococcus pentosaceus_MVI5.1-1 1 2 0.5105702308 Pediococcus pentosaceus_MAVI2.2-1 1 3 0.5056111596 Pediococcus pentosaceus_ABVI15.1-1 2 0.5033879340 Pediococcus pentosaceus_POVI5-1 2 0.5031626240 1 Pediococcus pentosaceus_AII15.2.H-1 1 2 0.5022992880 Pediococcus pentosaceus_POAVI6.1-1 3 0.4979635584 Pediococcus pentosaceus_ABVI17.1-1 2 0.4941022555 1 Pediococcus pentosaceus_POAI.3-1 3 0.4932899968 1 3 0.4896026566 Pediococcus pentosaceus PRAI2-1 Pediococcus pentosaceus_MVI3-1 1 2 0.4796409880 1 Pediococcus pentosaceus_POVI6-1 2 0.4720077534 Pediococcus pentosaceus_MAVI5.2-1 1 2 0.4718739371 Pediococcus pentosaceus_MAII.3.2-1 1 2 0.4680102940 Pediococcus pentosaceus_ABVI4.1-1 1 2 0.4649807256 Pediococcus pentosaceus POAVI.8-1 2 0.4636068257 1 Pediococcus sp._MAVI7.1-1 1 3 0.4560500196 Pediococcus pentosaceus_ABI11-1 1 2 0.4454150444shortened output......

\$clus.avg.widths

[1] 0.375931531 0.006128141 0.302342829

\$avg.width
[1] 0.1511431
sters silhouette plot





3.8. PointClusterVal

PointClusterVal is a wrapper of two functions for clustering tendency and validation statistics

```
@param
            df_m: dataframe containing peaks and metadata
            meth: clustering algorithms ("hclust", default value), other values: "kmeans", "pam", "clara", "fanny",
@param
                   "hclust", "agnes", "diana"
@param
            dist: distances, "euclidean", (default value), "maximum", "manhattan", "canberra", "binary" "minkowski"
            meth2: hc methods "ward.D2", (default value), average", "ward.D", "single", "complete", "mcquitty", "median"
@param
                   or "centroid"
            varCat1: categorical variable for choosing isolates, examples: "Taxonomie", "Genre", "Date.d.analyse",
@param
                  "Origine", "Ruche", "Nutrition", "Date.de.récolte", "Lieu.de.la.ruche"
@param
            value: level of catVar1, examples: "Lactobacillus" ("Genre"), Taxonomie ("Pediococcus pentosaceus"),
                   "Erica cinerea" ("Nutrition") ,...
@param
            varCat2: categorical variable for partitioning the set of isolates chosen by using varCat1
@return
            figures and statistics
@examples ff<-PointClusterVal (df_Peaks, varCat1="Genre", value="Lactobacillus", varCat2="Nutrition")
```

 $source: https://www.rdocumentation.org/packages/fpc/versions/2.2-8/topics/cluster.stats \\ https://www.datanovia.com/en/lessons/cluster-validation-statistics-must-know-methods/$

ff<-PointClusterVal(df_Peaks, varCat1="Genre", value="Lactobacillus", varCat2="Nutrition")

output:

```
$hopkins_stat
[1] 0.7644649
$plot
stats::hclust(d = x, method = hc_method)
Cluster method : ward.D2
Distance
           : euclidean
Number of objects: 56
[1] 56
$cluster.number
[1] 7
$cluster.size
[1] 14 5 15 4 9 2 7
$min.cluster.size
[1] 2
$noisen
[1] 0
$diameter
\hbox{\tt [1] 22.95959 19.32593 19.44587 22.41606 10.22588 9.00450 18.59128}
$average.distance
\hbox{\tt [1] 17.622715 10.696875 12.326323 18.685366 6.802126 9.004500 13.368503}
$median.distance
\hbox{\tt [1] 17.689350\ 7.063469\ 12.905385\ 21.432008\ 7.213886\ 9.004500\ 16.300675}
Sseparation
[1] 11.631167 9.989521 14.237874 15.951624 9.989521 17.598738 15.771988
$average.toother
\hbox{\tt [1]}\ 18.55982\ 17.98586\ 18.48002\ 20.32399\ 17.17040\ 21.28370\ 19.90793
$separation.matrix
    [,1] [,2] [,3] [,4] [,5] [,6] [,7]
\hbox{\tt [1,]}\ 0.00000\ 13.665972\ 14.23787\ 16.94605\ 11.631167\ 19.68021\ 17.11445
\hbox{\tt [2,] 13.66597 0.000000 15.38696 17.21943 9.989521 19.42117 18.79356}
[3,] 14.23787 15.386955 0.00000 16.79070 15.007128 18.44822 15.77199
[4,] 16.94605 17.219435 16.79070 0.00000 15.951624 17.59874 18.81842
\hbox{\tt [5,] 11.63117 9.989521 15.00713 15.95162 0.000000 18.47002 17.33686}
\hbox{\tt [6,] 19.68021 19.421169 18.44822 17.59874 18.470018 0.000000 21.06001}
[7,] 17.11445 18.793559 15.77199 18.81842 17.336860 21.06001 0.00000
$ave.between.matrix
     [,1] [,2] [,3] [,4] [,5] [,6] [,7]
[1,] 0.00000 17.80269 18.18237 20.91226 16.70647 21.59085 20.08206
[2,] 17.80269 0.00000 18.62993 19.48475 13.78734 21.12029 20.61810
\hbox{\tt [3,]}\ 18.18237\ 18.62993\ \ 0.00000\ 20.02739\ 17.03173\ 21.31666\ 19.13564
```

[4,] 20.91226 19.48475 20.02739 0.00000 18.71575 21.78504 22.03279 [5,] 16.70647 13.78734 17.03173 18.71575 0.00000 20.03088 19.11152 [6,] 21.59085 21.12029 21.31666 21.78504 20.03088 0.00000 22.03982 [7,] 20.08206 20.61810 19.13564 22.03279 19.11152 22.03982 0.00000

\$average.between

[1] 18.69382

\$average.within

[1] 13.08297

\$n.between

[1] 1270

\$n.within

[1] 270

\$max.diameter

[1] 22.95959

\$min.separation

[1] 9.989521

\$within.cluster.ss

[1] 4993.077

\$clus.avg.silwidths 1 2

1 2 3 4 5 6 7

 $\hbox{-0.05629018} \ 0.24330709 \ 0.27923312 \ 0.01124142 \ 0.50783301 \ 0.55037281 \ 0.29374212$

\$avg.silwidth

[1] 0.2212388

\$g2 NULL

\$g3 NULL

\$pearsongamma

[1] 0.5329142

\$dunn

[1] 0.4350914

\$dunn2

[1] 0.7378681

\$entropy

[1] 1.776377

\$wb.ratio [1] 0.6998553

\$ch

[1] 6.676386

\$cwidegap

 $\hbox{\tt [1] 17.838066 18.601795 14.075963 21.714907 7.029929 9.004500 15.560914}$

\$widestgap

[1] 21.71491

\$sindex

[1] 10.22678

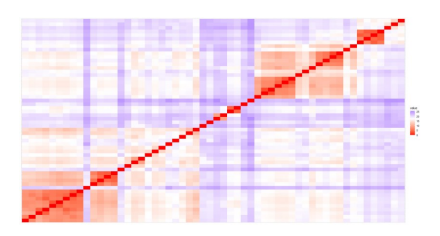
\$corrected.rand

NULL

\$vi

NULL

- [1] "external clustering validation (corrected.rand, vi)"
- [1] "Corrected Rand index"



- [1] 0.149149
- [1] "Meila variation of information (VI) index"
- [1] 2.080936

4. Correlograms for MALDI_TOF spectra

4.1. Visual CorrDistM

Visual_CorrDistM is based on functions for computing and visualizing distance matrix and correlograms

```
@param
            df_m: dataframe containing peaks and metadata
           dist: distances ("euclidean", default value), "euclidean", "maximum", "manhattan", "canberra", "binary",
@param
            "minkowski", "pearson", "spearman" or "kendall".
           varCat1: categorical variable for choosing isolates, examples: "Taxonomie", "Genre", "Date.d.analyse", "Origine",
@param
                    "Ruche", "Nutrition", "Date.de.récolte"
           value: value of catVar1 "Lactobacillus" ("Genre"), Taxonomie ("Pediococcus pentosaceus"), "Erica cinerea"
@param
                  ("Nutrition"),...
@param
           Visual: correlogram, "Corr", (default value) or distance matrix figure ("Dist") (this second option is more general
                    because it includes correlation, see argument dist)
           CorrFig: correlogram type, "circle", "square", "ellipse", "number", "pie", "shade" and "color" (default value)
@param
           Ord: correlogram arrangement methods (ord="FPC", default value), "AOE", "hclust"
@param
           layout: correlogram layout: "full", "upper", "lower" (layout="upper", default value)
@param
           pv: boolean variable including or not probability values. (pv=TRUE, default value)
@param
@param
            sig: significance level (sig="0.05", default value)
            figures and statistics
@return
@examples Visual_CorrDistM (df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus"),
              Visual_CorrDistM (df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus", VisualM="dist")
              Visual_CorrDistM (df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus", VisualM="dist",
              dist="pearson")
```

source: https://cran.r-project.org/web/packages/corrplot/vignettes/corrplot-intro.html https://www.rdocumentation.org/packages/corrplot/versions/0.84/topics/corrplot

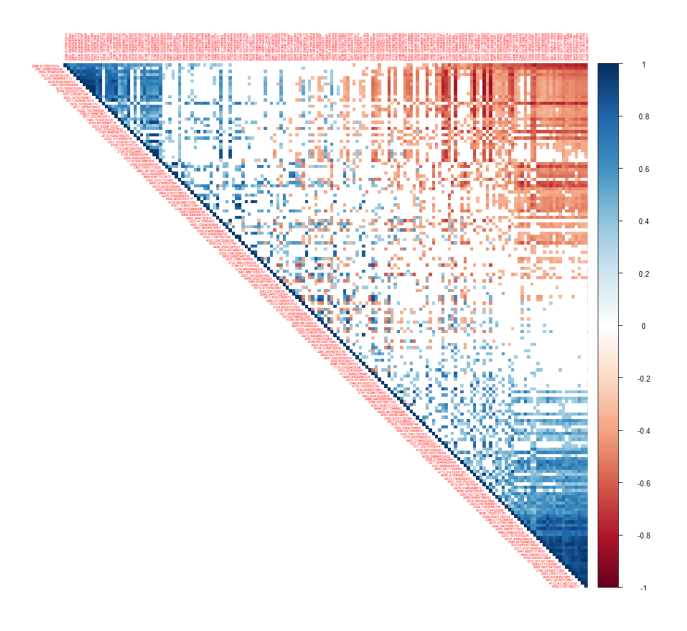
Visual_CorrDistM (df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus")

Output:

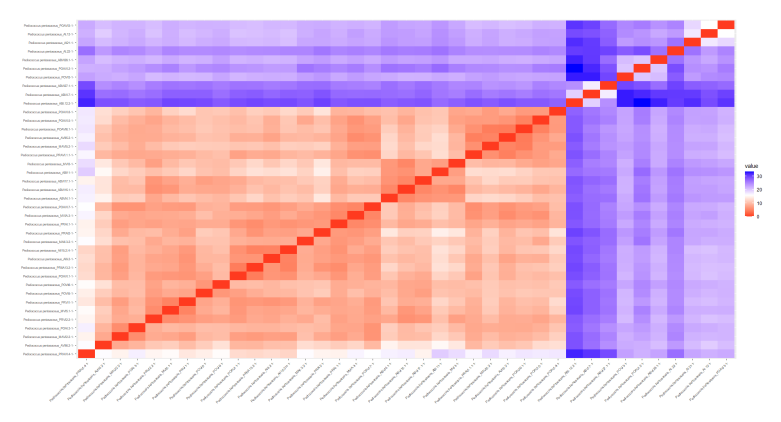
Correlation Matrix

2018.79282	2168279 2035.	60098965	176 2051.553	885901724	2067.957122	74841 2083.58086146253	
2018.79282168279	1.00	0.91	0.93	-0.29	0.10		
2035.60098965176	0.91	1.00	0.95	-0.43	-0.06		
2051.55385901724	0.93	0.95	1.00	-0.28	0.14		
2067.95712274841	-0.29	-0.43	-0.28	1.00	0.88		
2083.58086146253	0.10	-0.06	0.14	0.88	1.00		
2106.58686543244	0.94	0.90	0.94	-0.14	0.27		
2106.58686	5543244 2124.	50092568	671 2147.402	266739652	2166.009729	51332 2182.83406086648	
2018.79282168279	0.94	0.92	0.36	0.23	0.81		
2035.60098965176	0.90	0.93	0.23	0.13	0.77		
2051.55385901724	0.94	0.97	0.39	0.25	0.88		
2067.95712274841	-0.14	-0.23	0.69	0.81	0.18		
2083.58086146253	0.27	0.19	0.86	0.90	0.56		
2106.58686543244	1.00	0.97	0.49	0.38	0.90		
2205.99202292547 2223.4110890903 2242.89870280795 2258.16638713302 2261.88048340716							
2018.79282168279	0.91	0.78	-0.23	-0.14	0.15		
2035.60098965176	0.88	0.69	-0.38	-0.22	0.14		
2051.55385901724	0.96	0.84	-0.23	-0.10	0.22		

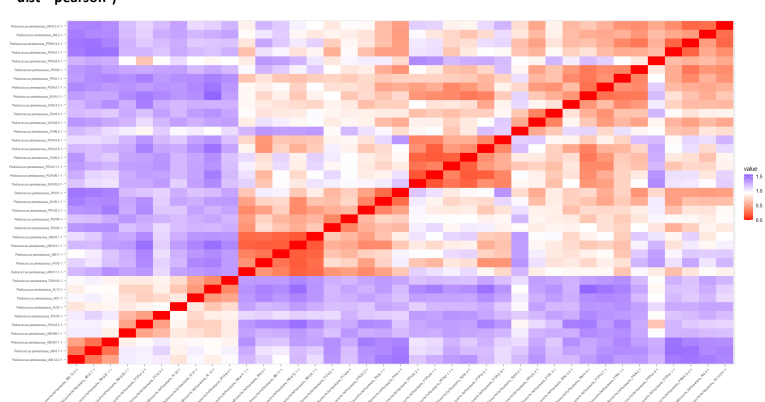
2067.95712274841	-0.19	0.07	0.87	0.49	0.18	
2083.58086146253	0.23	0.47	0.79	0.45	0.30	
2106.58686543244	0.96	0.89	-0.10	-0.06	0.21	
2271.624948	329559 2278.	057290917	749 2286.81	728075514	2306.65291	.128723 2311.37471904839
2018.79282168279	0.13	-0.33	0.61	-0.20	-0.24	
2035.60098965176	0.02	-0.36	0.64	-0.33	-0.38	
2051.55385901724	0.15	-0.28	0.57	-0.22	-0.24	
2067.95712274841	0.67	0.47	-0.55	0.66	0.79	
2083.58086146253	0.73	0.37	-0.31	0.59	0.71	
2106.58686543244	0.25	-0.24	0.52	-0.14	-0.12	
2320.093073	38399 2332.0	613581680	3 2365.017	50364438 2	393.378780	88851 2404.97257576619
2018.79282168279	0.14	0.39	-0.10	-0.40	-0.10	
2035.60098965176	0.09	0.36	-0.24	-0.43	-0.14	
2051.55385901724	0.20	0.44	-0.13	-0.32	-0.13	
2067.95712274841	0.34	0.07	0.70	0.65	-0.06	
2083.58086146253	0.45	0.31	0.66	0.54	-0.09	
2106.58686543244	0.21	0.42	0.00	-0.27	-0.12	
2411.328243	189933 2421.	047557854	101 2466.44	858585066	2478.51460)496673 2530.54564880773
2018.79282168279	0.70	0.78	0.10	-0.28	0.72	
2035.60098965176	0.68	0.76	0.08	-0.35	0.74	
2051.55385901724	0.69	0.83	0.10	-0.23	0.79	
2067.95712274841	-0.42	-0.48	0.45	0.66	-0.40	
2083.58086146253	-0.10	-0.09	0.48	0.56	-0.06	
2106.58686543244	0.70	0.75	0.23	-0.16	0.72	
shortene	ed output					



Visual_CorrDistM (df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus", VisualM="dist")



Visual_CorrDistM (df_Peaks, varCat1="Taxonomie", value="Pediococcus pentosaceus", VisualM="dist", dist="pearson")



5. Principal Component Analysis and clustering of MALDI_TOF spectra

5.1. PCA_Clus

Output:

Principal Component Analysis and clustering of Maldi_Tof spectra

```
df m: dataframe containing peaks and metadata
@param
           dist: distances, "euclidean", (default value), "euclidean", "maximum", "manhattan",
@param
                "canberra", "binary", "minkowski", "pearson", "spearman" or "kendall".
           varCat1: categorical variable for choosing isolates, examples: "Taxonomie", "Genre",
@param
                   "Date.d.analyse", "Origine", "Ruche", "Nutrition", "Date.de.récolte",
                   "Lieu.de.la.ruche"
           value: level of catVar1, examples: "Lactobacillus" ("Genre"), Taxonomie ("Pediococcus
@param
                  pentosaceus"), "Erica cinerea" ("Nutrition"),...
           meth: clustering method, ward(default value), "average", "single", "complete'
@param
           graph: visual analysis, "dendf", "dendh", (dendograms) "factorMapf", "factorMaph",
@param
                 "factorMapClus", (factor maps) (default value, graph="factorMapClus")
@param
           pc: number of principal components (pc=3, default value)
          figures and statistics
@return
@examples pc<-PCA Clus(df Peaks, varCat1="Genre", value="Lactobacillus"),
           pc<-PCA_Clus(df_Peaks, varCat1="Genre", value="Lactobacillus", graph="dendh")
           pc<-PCA Clus(df Peaks, varCat1="Genre", value="Lactobacillus", graph="dendf"),
           pc<-PCA_Clus(df_Peaks, varCat1="Genre", value="Lactobacillus", graph="factorMapf")
  source: https://www.rdocumentation.org/packages/FactoMineR/versions/2.2/topics/PCA
         https://www.rdocumentation.org/packages/FactoMineR/versions/2.2/topics/HCPC
         http://www.sthda.com/english/articles/31-principal-component-methods-in-r-practical-
         guide/117-hcpc-hierarchical-clustering-on-principal-components-essentials/
          https://rpkgs.datanovia.com/factoextra/
         http://factominer.free.fr/factomethods/hierarchical-clustering-on-principal-
          components.html
```

pc<-PCA_Clus(df_Peaks, varCat1="Genre", value="Lactobacillus")

[1] "contribution of principal components for each cluster"

¢`1`

 v.test Mean in category Overall mean sd in category Overall sd
 p.value

 Dim.2 -4.365895
 -3.863117 -7.432051e-16
 1.831196
 4.332495 1.266031e-05

 Dim.1 -5.412847
 -5.525758 5.144694e-16
 1.603985
 4.998499 6.203042e-08

\$`2`

v.test Mean in category Overall mean sd in category Overall sd p.value

Dim. 2 6.163973 9.527227 -7.432051e-16 2.074991 4.332495 7.094185e-10

Dim. 1 -2.356219 -4.201679 5.144694e-16 1.177872 4.998499 1.846204e-02

\$'3'

v.test Mean in category Overall mean sd in category Overall sd p.value
Dim.3 6.194998 8.625274 1.298812e-15 1.903931 3.576874 5.828569e-10

\$`4`

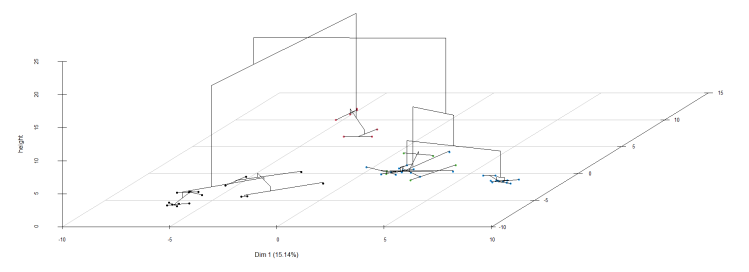
 v.test Mean in category Overall mean sd in category Overall sd
 p.value

 Dim.1 5.982709
 4.331418 5.144694e-16
 2.690896
 4.998499 2.194566e-09

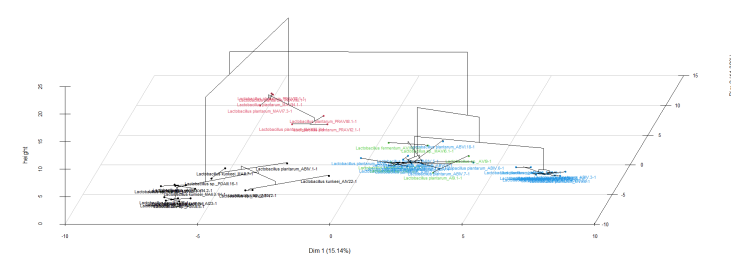
 Dim.3 -2.171633
 -1.125078 1.298812e-15
 2.226520
 3.576874 2.988339e-02

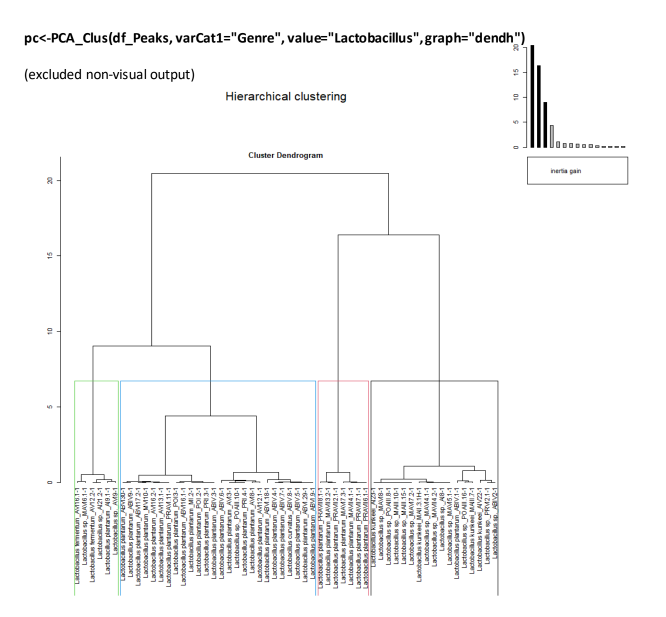
Hierarchical clustering on the factor map

cluster 1
cluster 2
cluster 3
cluster 4

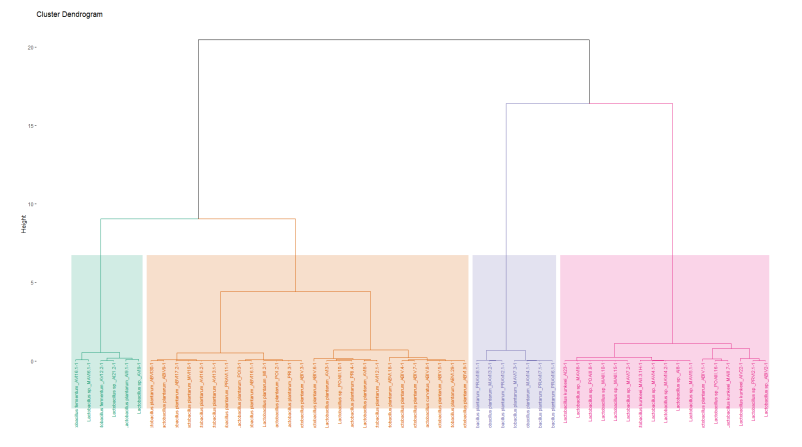




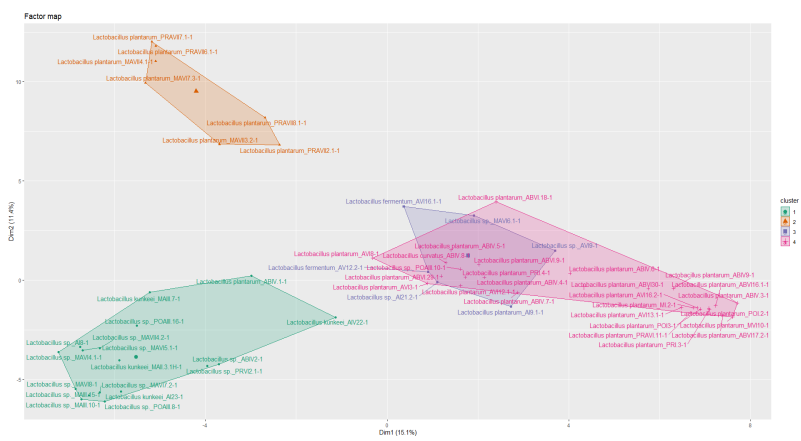




pc<-PCA_Clus(df_Peaks, varCat1="Genre", value="Lactobacillus", graph="dendf")



pc<-PCA_Clus(df_Peaks, varCat1="Genre", value="Lactobacillus", graph="factorMapf")

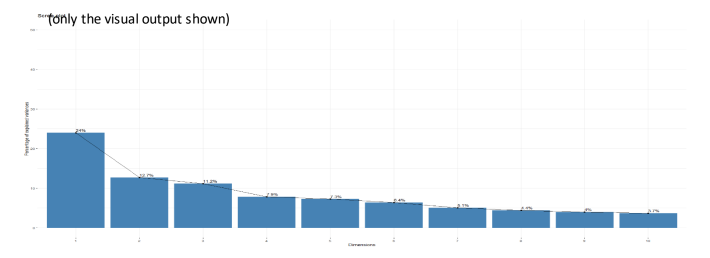


5.2. SPCA

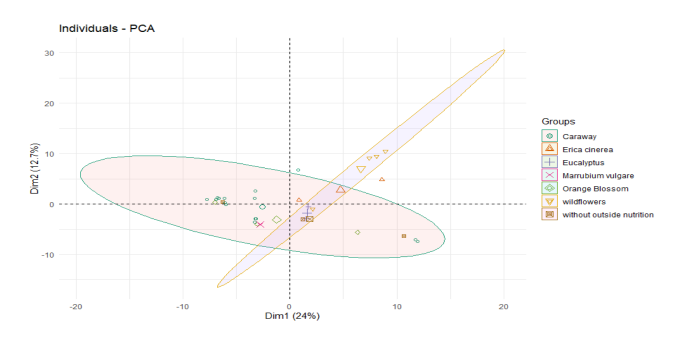
Principal Component Analysis of Maldi_Tof spectra with external clusters based on metadata information

@param df_m: dataframe containing peaks and metadata varCat1: categorical variable for choosing isolates, examples: "Taxonomie", "Genre", "Date.d.analyse", "Origine", @param "Ruche", "Nutrition", "Date.de.récolte", "Lieu.de.la.ruche" value: level of catVar1 "Lactobacillus" ("Genre"), Taxonomie ("Pediococcus pentosaceus"), "Erica cine rea" @param ("Nutrition"),... varCat2: categorical variable for partitioning the chosen isolates, Taxonomie, Genre, Date.d.analyse, Origine, @param Ruche, Nutrition, Date.de.récolte, Lieu.de.la.ruche @param contDim: graph and statistics for PC contributions (default, contDim =TRUE) @param contVar: graph and statistics for variable contributions(default, contVar=FALSE) @param contind: graph and statistics for isolate contributions (default, contind= TRUE) @examples hg<-SPCA(df_Peaks, varCat1="Genre", value="Lactobacillus", varCat2="Nutrition")

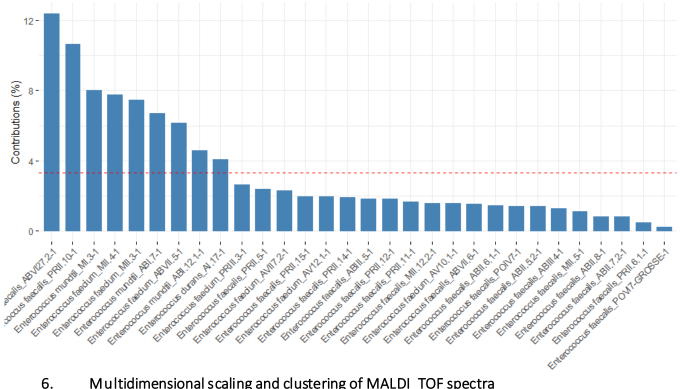
SPCA<- function(df_m, varCat1, value, varCat2, contDim=TRUE, contVar= FALSE, contInd=FALSE)



hg<-SPCA(df_Peaks, varCat1="Genre", value="Enterococcus", varCat2="Nutrition",contDim=FALSE, contVar=FALSE,contInd=TRUE) (only the visual output shown)



Contribution of individuals to Dim-1-2-3-4-5



6. Multidimensional scaling and clustering of MALDI TOF spectra

6.1. MDS_Clus

MDS Clus, function for Multidimensional scaling and kmeans-based analysis of Maldi Tof spectra

```
@param
            df_m: dataframe containing peaks and metadata
@param
            dist: distances: "euclidean", "maximum", "manhattan", "canberra", "binary" or "minkowski"
@param
            varCat1: categorical variable for choosing isolates, examples: "Taxonomie", "Genre",
             "Date.d.analyse", "Origine", "Ruche", "Nutrition", "Date.de.récolte", "Lieu.de.la.ruche"
@param value: level of the chosen categorical variable as catVar1, examples: "Lactobacillus" ("Genre"), Taxonomie("Pediococcus
               pentosaceus"), "Erica cinerea" ("Nutrition"),...
           grah: graphs: lab mdsclus, (default value) lb mds, mdsclaing
          figures and statistics
@examples g<-MDS Clus(df Peaks, varCat1="Genre", value="Lactobacillus")
```

source: https://rstudio-pubs-static.s3.amazonaws.com/274936_050c742fb3514bbaa87ce6ee2686af8c.html http://www.sthda.com/english/articles/31-principal-component-methods-in-r-practical-guide/122multidimensional-scaling-essentials-algorithms-and-r-code/ http://ugrad.stat.ubc.ca/R/library/mva/html/cmdscale.html

g<-MDS Clus(df Peaks, varCat1="Genre", value="Lactobacillus")

output:

		Dim.1	Dim.2 groups		
Lactobacillus	plantarum_	_ABIV.1-1	8.073724e-04	9.230089e-05	1
Lactobacillus	plantarum_	_ABIV.3-1	-1.626189e-03	8.995464e-04	3
Lactobacillus	nlantarum	ΔRIV 4-1	-2 157681e-04	5 663535e-04	1

```
Lactobacillus plantarum_ABIV.5-1 -7.135335e-06 5.353676e-04
                 Lactobacillus plantarum_ABIV.6-1 -9.152847e-04 1.004312e-03 3
                 Lactobacillus plantarum_ABIV.7-1 -2.614522e-04 8.268632e-04
                 Lactobacillus curvatus_ABIV.8-1 1.267334e-04 -2.178710e-04 1
                 Lactobacillus plantarum_ABVI.18-1 -2.491353e-04 -3.890558e-04
                 Lactobacillus plantarum_ABVI.29-1 7.677427e-06 4.594310e-04
                 Lactobacillus kunkeei_MAII.3.1H-1 1.531006e-03 3.411630e-04
                 Lactobacillus sp._MAIII.10-1
                                                     1.691986e-03 1.385961e-03
                 Lactobacillus sp._MAIII.15-1
                                                      1.657554e-03 1.341994e-03
                 Lactobacillus kunkeei_MAIII.7-1 7.231229e-04 -4.565799e-05
                 Lactobacillus plantarum_MI.2-1 -1.471482e-03 8.523043e-04
                 Lactobacillus sp._POAIII.10-1 -1.439868e-04 -8.438319e-04 2
                 Lactobacillus sp._POAIII.16-1 9.544077e-04 4.409873e-04 1 _{\mathrm{groups}} 1 _{\mathrm{a}} 2 _{\mathrm{a}} 3
                 Lactobacillus sp._POAIII.8-1
                                                        1.550021e-03 1.166693e-03 1
                 .....shortened out put ......
                                                                                                                                                                                             Lactobacillus sp._MAIII.10-1
                                                                                                                                                                                Lactobacillus kunkeei Al23-1--
                                                                                                                                                                        Lactobacillus sp. MAIII.15-1 Lactobacillus sc
                                                                                                                                                                                              Lactobacillus sp._MAVI8-1
                                                                    Lactobacillus plantarum_ABIV.6-1
0.001
                                                                                                            Lactobacillus plantarum_AVI12.1-1
                        Lactobacillus plantarum_ABIV.3-1 Lactobacillus plantarum_PRI.3-1 Lactobacillus plantarum_POI.2-1 * Lactobacillus plantarum_MI.2-1
                                                                                             Lactobacillus plantarum_ABIV.7-1
                                                                                                                           Lactobacillus plantarum_AVI3-1
                                                                                                                                                                               Lactobacillus sp._MAVII4.2-1
                              Lactobacillus plantarum_AVI13.1-1
                                                                                                  Lactobacillus plantarum_PRI.4-1
                                                                                                                      Lactobacillus plantarum_ABIV.5-1
                                                                                                                                                                                            Lactobacillus sp._MAVI5.1-1
                                                                                             Lactobacillus plantarum_ABIV.4-1
                                                                                                                        Lactobacillus plantarum_ABVI.9-1
                                                   Lactobacillus plantarum AVI16.2-1
                                                                                                                                                        Lactobacillus sp._POAIII.16-1
                                                                                                     Lactobacillus plantarum_ABVI.29-1
                               Lactobacillus-plantarum_ABVI30-1
                                                                                                                                                                                 Lactobacillus kunkeei MAII 3 1H-1
                                                                                                                            Lactobacillus plantarum_AVI8-1 Lactobacillus plantarum_ABIV.1-1
                              Lactobacillus plantarum POI3-1
       actobacillus plantarum_ABV116.1-1
                                                                                                                                            Lactobacillus kunkeei_MAIII.7-1
                                                                                                                Lactobacillus curvatus_ABIV.8-1
                                                                                                                                         actobacillus sp._PRVI2.1-1
                                                                                                   Lactobacillus plantarum_ABVI.18-1
                                                                                                                                              Lactobacillus kunkeei AlV22-1
                                                                                  Lactobacillus sp_AVI9-1
                                                                                                                                              Lactobacillus plantarum_MAVII3.2-1
                                                                                                          Lactobacillus sp._POAIII.10-1
                                                                                                                          Lactobacillus plantarum_PRAVII2.1-1 Lactobacillus sp._ABIV2-1
-0.001
                                                                                                                                   Lactobacillus plantarum_PRAVII8.1-1
                                                                                                                                                    Lactobacillus plantarum_MAVI7.3-1
                                                                                                  Lactobacillus sp._Al21.2-1
                                                                                                                                                                   Lactobacillus plantarum_PRAVII7.1-1
                                                                            Lactobacillus plantarum_Al9.1-1
                                                                                                      Lactobacillus fermentum_AVI16.1-1
                                                                                                                                                Lactobacillus plantarum MAVII4.1-1
-0.002
                                                                                               Lactobacillus fermentum AV12.2-1
                                                                                                                                                       Laetobacillus plantarum PRAVII6.1-1
                                                                                Lactobacillus sp._MAVI6.1-1
                         -0.002
                                                                        -0.001
                                                                                                                      0.000
                                                                                                                                                                     0.001
                                                                                                         Dim.1
```

g<-MDS_Clus(df_Peaks, varCat1="Taxonomie", value="All", nc=5)

output:

Dim.2

```
      Dim.1
      Dim.2 groups

      Pediococcus pentosaceus_AI.12-1
      2.783290e-04 6.720991e-04 5

      Enterococcus durans_AI.17-1
      1.074552e-03 2.928558e-04 5

      Pediococcus pentosaceus_AI.22-1
      3.911749e-04 1.297149e-04 5

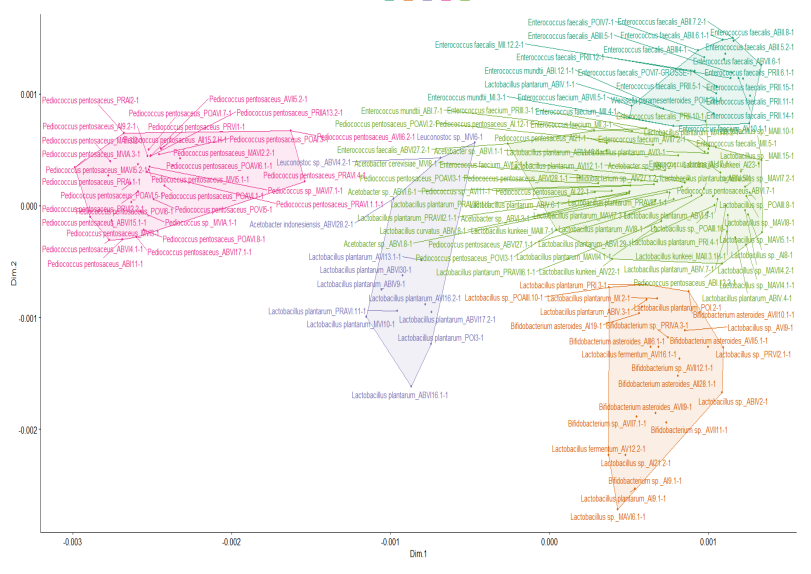
      Enterococcus mundtii_ABI.12.1-1
      7.160698e-04 8.675265e-04 1

      Pediococcus pentosaceus_ABI.12.2-1
      1.085123e-03 -5.198231e-04 5

      Enterococcus mundtii_ABI.7-1
      8.018773e-04 6.104123e-04 5
```

```
Enterococcus faecalis_ABII.5.2-1 1.105291e-03 1.356064e-03
Enterococcus faecalis_ABII.6.1-1 1.090702e-03 1.486138e-03
Enterococcus faecalis_ABII.7.2-1 1.152658e-03 1.498682e-03
Enterococcus faecalis ABII.8-1
                               1.158135e-03 1.474918e-03
Enterococcus faecalis_ABIII.5-1
                               1.146276e-03 1.190664e-03
Lactobacillus plantarum_ABIV.1-1
                                 9.803381e-04 7.566789e-04
Lactobacillus plantarum ABIV.3-1
                                 6.084417e-04 -8.424467e-04
Lactobacillus plantarum_ABIV.4-1
                                 1.166997e-03 -2.844821e-04
Lactobacillus plantarum_ABIV.5-1
                                 9.696000e-04 1.512996e-04
Lactobacillus plantarum_ABIV.6-1
                                 5.021067e-04 1.256638e-04
   .....shortened output.....
```

```
groups a 1 a 2 a 3 a 4 a 5
```



6.2. SMDS

Multidimensional scaling and external cluster-based analysis of MALDI_TOF spectra

@param df_m: dataframe containing peaks and metadata
 @param dist: distances: "euclidean" (default value), "maximum", "manhattan", "canberra", "binary" or "minkowski"
 @param varCat1: categorical variable for choosing isolates, examples: "Taxonomie",

```
"Genre", "Date.d.analyse", "Origine", "Ruche", "Nutrition", "Date.de.récolte",

"Lieu.de.la.ruche"

@param value: level of the chosen categorical variable catVar1, examples: "Lactobacillus"

("Genre"), Taxonomie("Pediococcus pentosaceus"), "Erica cinerea" ("Nutrition") ,...

@param grah: graphs: "lab_mdsGroups" (default value), "mdsGroups"

@return figures and statistics

@examples

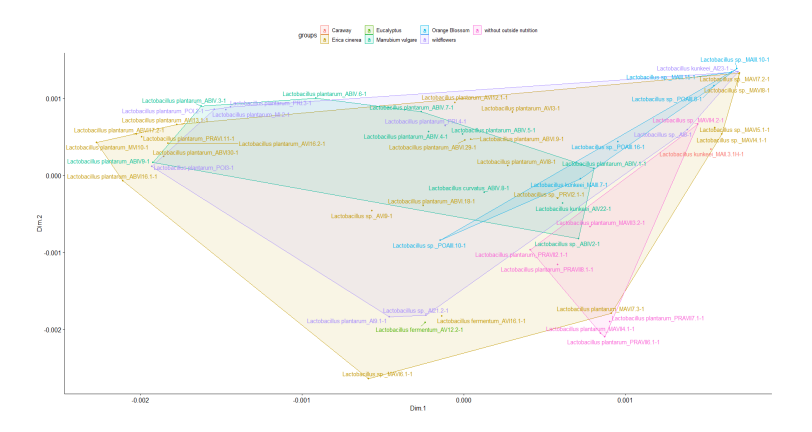
SMDS(df_Peaks, varCat1="Genre", value="Lactobacillus", varCat2="Ruche",dist="euclidean", grah="lab_mdsGroups")

SMDS(df_Peaks, varCat1="Genre", value="Lactobacillus" varCat2="Ruche",)

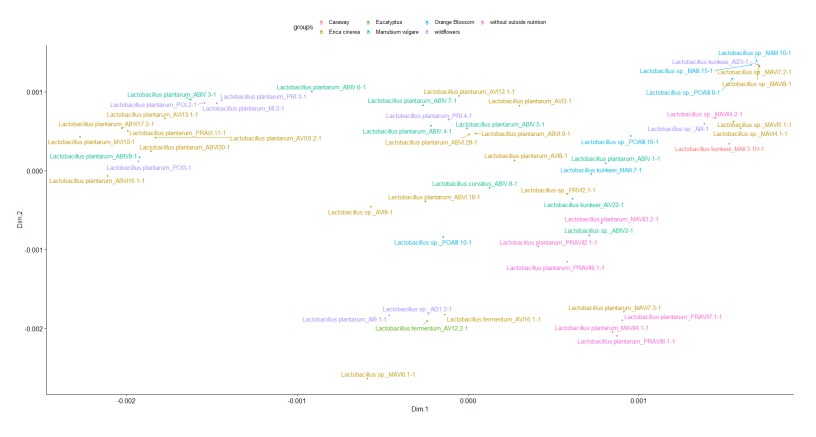
SMDS(df_Peaks, varCat1="Genre", value="Lactobacillus", varCat2="Nutrition")

SMDS(df_Peaks, varCat1="Genre", value="All",varCat2="Taxonomie")
```

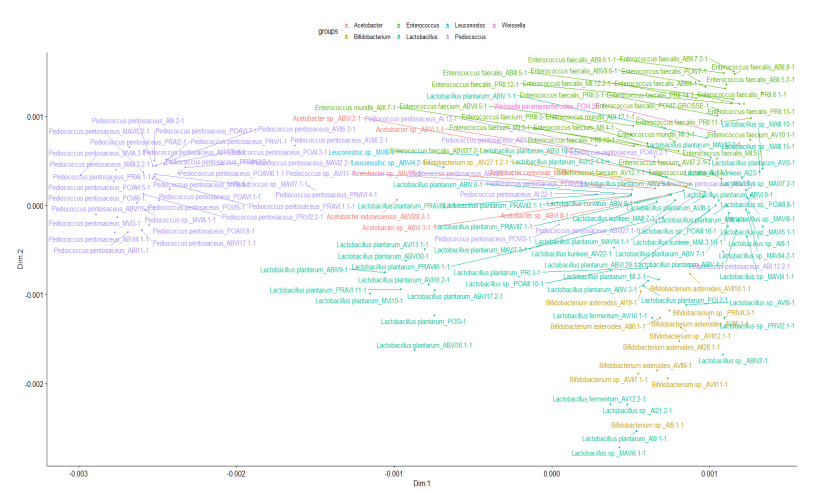
SMDS(df_Peaks, varCat1="Genre", value="Lactobacillus", varCat2="Nutrition")



SMDS(df_Peaks, varCat1="Genre", value="Lactobacillus", varCat2="Nutrition",grah="mdsGroups")



SMDS(df_Peaks, varCat1="Taxonomie", value="All", varCat2="Genre",grah="mdsGroups")



SMDS(df_Peaks, varCat1="Genre", value="All", varCat2="Taxonomie", grah="mdsGroups")

```
a Acetobacter cerevisiae a Bifidobacterium sp. a Enterococcus mundtii a Lactobacillus plantarum
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         A Pediococcus sp.
                                                                                                                                                                                                                                                                a Acetobacter indonesiensis a Enterococcus durans a Lactobacillus curvatus a Lactobacillus sp.
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                                  ediococcus pentosaceus_PRAI2-1 Pediococcus pentosaceus_POAVI.7-1
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                                ediococcus pentosaceus MPenibocccus pentosaceus POAVI.8-1
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                                Pediococcus pentosaceus_ABVI4.1-1 Pediococcus pentosaceus_ABVI17.1-1
                                                                                                                                                                                                                                                                                                                                                                      Lactobacillus plantarum ABVI30-1 Lactobacillus plantarum ABVI.9-1 Taetobacillus kunkeei AV22-1 Lactobacillus plantarum ABVI.7-1 Lactobacillus sp. AI8-1
                                                                                                            Pediococcus pentosaceus_ABI11-1
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-0.002
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                                                    -0.003
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