

# Package: CoNI (via r-universe)

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**Type** Package

**Title** Correlation Guided Network Integration (CoNI)

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**Description** Integrates two numerical omics data sets from the same samples using partial correlations. The output can be represented as a network, bipartite graph or a hypergraph structure. The method used in the package refers to Klaus et al (2021) <[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)>.

**License** GPL-3

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**Suggests** kableExtra (>= 1.3.2), knitr (>= 1.31), rmarkdown (>= 2.6)

**VignetteBuilder** knitr

**SystemRequirements** python3

**Repository** <https://jmanuelmk.r-universe.dev>

**RemoteUrl** <https://github.com/jmanuelmk/coni>

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---

assign\_colorsAnnotation

*Assing Colors to Class*

---

### Description

This function assigns two color columns (color name and rgb) to an annotation data frame according to a column named 'Class' or 'class'

### Usage

```
assign_colorsAnnotation(AnnotationDf, col = "Class")
```

**Arguments**

AnnotationDf    Annotation data frame that contains a factor variable to use to assign colors  
col             Column with factor variable that will be used to assign colors

**Value**

The input data.frame with two extra columns specifying the colors for all vertexes according to their respective vertex-class

---

Chow\_GeneExpData    *Chow gene expression data*

---

**Description**

Chow gene expression data

**Author(s)**

Jose Manuel Monroy <no1ozz@gmail.com>

**References**

[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)

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Chow\_MetaboliteData    *Chow metabolite data*

---

**Description**

Chow metabolite data

**Author(s)**

Jose Manuel Monroy <no1ozz@gmail.com>

**References**

[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)

---

Compare\_Triplets      *Compare triplets*

---

### Description

Compare vertexFeature-vertexFeature-edgeFeature between two treatments, that is, find the shared triplets between two different CoNI runs.

### Usage

```
Compare_Triplets(
  Treat1_path,
  Treat2_path,
  OutputName = "Shared_Genes_and_Edges_Treat1vsTreat2.csv"
)
```

### Arguments

Treat1_path	TableForNetwork_file1 (file generated by CoNI) with path for Treatment 1
Treat2_path	TableForNetwork_file2 (file generated by CoNI) with path for Treatment 2
OutputName	Output file name with path

### Value

A data.frame with the shared triplets (vertex1 vertex2 edge\_feature) between two CoNI runs

### Examples

```
#For an example see the vignette
```

---

Compare\_VertexClasses\_sharedEdgeFeatures  
*Table VertexClass pairs of shared Edge Features*

---

### Description

Compare VertexClass pairs of the shared Edge Features of two treatments (e.g., lipid-class-pairs per shared gene)

### Usage

```
Compare_VertexClasses_sharedEdgeFeatures(
  Treat1_path,
  Treat2_path,
  OutputName = "Shared_Genes_and_Edges_Treat1vsTreat2.csv",
  Treat1Name = "Treat1",
  Treat2Name = "Treat2"
)
```

**Arguments**

Treat1_path	TableForNetwork_file (file generated by CoNI) with path of Treatment 1
Treat2_path	TableForNetwork_file (file generated by CoNI) with path of Treatment 2
OutputName	Output file name with path
Treat1Name	Name of treatment one, default Treat1
Treat2Name	Name of treatment one, default Treat2

**Value**

A data.frame with all possible vertex-class pairs and their numbers per edge-feature and treatment.

**Examples**

```
#For an example see the vignette
```

---

CoNI *Correlation guided Network Integration*

---

**Description**

CoNI is the main function of Correlation guided Network Integration (CoNI). Input data should come from two sources (e.g., gene expression and metabolite expression), and it should come from the same samples. It calculates all pairwise correlations of the second data input elements and the partial correlations of these pairwise elements with respect to the elements of the first data input. Both data inputs can be prefiltered to include only those elements that significantly correlate. The first data input can be prefiltered to keep just low variance elements ( $\text{var} < 0.5$ ). A Steiger test is used to identify significant changes between the correlation and partial correlation values. Results can be visually represented in a Network.

**Usage**

```
CoNI(  
  edgeD,  
  vertexD,  
  outputDir = "./CoNIOutput/",  
  saveRaw = TRUE,  
  outputNameRaw = "CoNIOutput",  
  onlySgRes = FALSE,  
  multipleTAdj = TRUE,  
  padjustvertexD = TRUE,  
  correlateDFs = TRUE,  
  filter_highVarianceEdge = TRUE,  
  splitedgeD = TRUE,  
  split_number = 2,  
  delPrevious = FALSE,  
  delIntermediaryFiles = TRUE,
```

```

iteration_start = 1,
numCores = NULL,
verbose = TRUE,
more_coef = FALSE,
edgeDname = "edgeD",
vertexDname = "vertexD",
saveFiles = TRUE
)

```

## Arguments

edgeD	Object to use as first data input (e.g., protein expression)
vertexD	Object to use as second data input (e.g., metabolite expression)
outputDir	Output Directory where results are stored
saveRaw	logical. If TRUE the raw output of CoNI is saved in the output directory (outputDir)
outputNameRaw	Name for the raw output file if saved
onlySgRes	logical. If TRUE CoNI output is filtered and only significant results are kept
multipleTAdj	logical. If TRUE it will filter results after adjustment of multiple testing
padjustvertexD	logical. If TRUE vertexD is filtered according to the significant adjusted p-value of its pairwise correlations
correlatedDFs	logical. If TRUE the elements that significantly correlate of vertexD are correlated with the elements of edgeD. Only the elements that significantly correlate are kept
filter_highVarianceEdge	logical. If TRUE features of edgeD with high variance are filtered out
splitedgeD	logical. If TRUE edgeD will be split in n subsets for the computation (some instances n+1). Keep as TRUE unless the data input is small
split_number	Number of parts to split the elements of edgeD
delPrevious	logical. If TRUE previous files of a previous run are deleted
delIntermediaryFiles	logical. If TRUE the output file of every iteration is deleted and only a single file with all results is kept
iteration_start	Iteration start for CoNI. Useful if run is interrupted as one can restart from the last iteration
numCores	Cores assigned for parallelization
verbose	logical. If TRUE output in the console is more verbose
more_coef	logical. If TRUE it will include the partial correlation of edge and vertex Features
edgeDname	File name extension for the edge features that significantly correlate with at least one vertex feature. This file will be read if the function is called again with the same input and with delPrevious=FALSE

vertexDname      File name extension for the vertex features that are involved in at least one significant correlation. This file will be read if the function is called again with the same input and with delPrevious=FALSE

saveFiles        logical. If FALSE CoNI function will not save any file to disk

### Value

CoNI returns a data.frame with the correlation coefficients of the vertex-pairs, the partial correlation coefficients for every triplet, and the pvalue of the Steiger tests

### Examples

```
#Run CoNI

#Load gene expression - Toy dataset of two treatments
data(GeneExpToy)
#Samples in rows and genes in columns
GeneExp <- as.data.frame(t(GeneExpToy))
hfd_gene <- GeneExp[1:8,] #high fat diet
chow_gene<- GeneExp[9:nrow(GeneExp),] #chow diet
#Load metabolite expression - Toy dataset of two treatments
data(MetaboExpToy)
MetaboExp <- MetaboExpToy
hfd_metabo <- MetaboExp[11:18,] #high fat diet
chow_metabo <- MetaboExp[1:10,] #chow diet
#Match row names both data sets
rownames(hfd_metabo)<-rownames(hfd_gene)
rownames(chow_metabo)<-rownames(chow_gene)

#Run CoNI with tiny example and no significance testing
#High fat diet
#For big datasets it is recommended to set splitedgeD to TRUE
#and split_number should be adjusted accordingly
#See vignette for an example
#Running CoNI with only a tiny dataset

CoNIResultsHFD <- CoNI(hfd_gene,hfd_metabo,
                      numCores = 2,
                      onlySgRes = FALSE,
                      filter_highVarianceEdge=FALSE,
                      padjustvertexD = FALSE,
                      correlateDFs = FALSE,
                      edgeDname="HFD_genes",
                      vertexDname = "HFD_metabolites",
                      saveFiles = FALSE,
                      splitedgeD = FALSE,
                      outputDir = "./")
```

---

CoNIResultsHFDToy      *Toy data HFD results*

---

**Description**

Toy data HFD results

**Author(s)**

Jose Manuel Monroy <nolozz@gmail.com>

**References**

[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)

---

CoNIResults\_Chow      *CoNI Results Chow*

---

**Description**

CoNI Results Chow

**Author(s)**

Jose Manuel Monroy <nolozz@gmail.com>

**References**

[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)

---

CoNIResults\_HFD      *CoNI Results HFD*

---

**Description**

CoNI Results HFD

**Author(s)**

Jose Manuel Monroy <nolozz@gmail.com>

**References**

[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)



---

createBipartiteGraph    *Bipartite Network*

---

### Description

This function creates a simple bipartite graph, it shows the driver and linker features as nodes.

### Usage

```
createBipartiteGraph(TableNetwork, colorVertexTable, incidenceMatrix = FALSE)
```

### Arguments

TableNetwork    TableForNetwork\_file (file generated by CoNI) with path

colorVertexTable

Table specifying the colors for the vertex features. The first column should contain the names matching the features of the vertex Data and another column should specify the colors (column name: Colors).

incidenceMatrix

logical. If TRUE it returns a hypergraph incidence matrix instead of a bipartite graph

### Value

An igraph object for a bipartite graph or a hypergraph incidence matrix to represent ResultsCoNI. Basic network statistics are included in the bipartite graph. See generate\_network function for details or consult the igraph package

### Examples

```
#See vignette for an example
```

---

create\_edgeFBarplot    *Vertex-class pairs profile of one shared edge feature*

---

### Description

This function will create a barplot from the output of Compare\_VertexClasses\_sharedEdgeFeatures for a specific shared Edge Feature (e.g., a shared gene).

**Usage**

```
create_edgeFBarplot(
  CompTreatTable,
  edgeF,
  treat1 = "Treatment1",
  treat2 = "Treatment2",
  factorOrder = NULL,
  col1 = "red",
  col2 = "blue",
  EdgeFeatureType = "Edge Feature",
  xlb = "Vertex-Class Pairs",
  ylb = "Number of pairs",
  szaxisTxt = 12,
  szaxisTitle = 12
)
```

**Arguments**

CompTreatTable	Output of Compare_VertexClasses_sharedEdgeFeatures
edgeF	Edge feature present in output of Compare_VertexClasses_sharedEdgeFeatures
treat1	Name of treatment one, default Treatment1. It should match the column names of the output of Compare_VertexClasses_sharedEdgeFeatures
treat2	Name of treatment one, default Treatment2. It should match the column names of the output of Compare_VertexClasses_sharedEdgeFeatures
factorOrder	A list specifying the order of the treatments.
col1	Color for Treatment 1
col2	Color for Treatment 2
EdgeFeatureType	Type of Edge Feature (e.g., Gene)
xlb	Name for x-axis
ylb	Name for the y-axis
szaxisTxt	Size axis text
szaxisTitle	Size axis titles

**Value**

A ggplot object for a barplot. The barplot shows the vertex-class pairs profile of a single shared edge feature between two treatments

**Examples**

```
data(VertexClassesSharedGenes_HFDvsChow)
create_edgeFBarplot(CompTreatTable = VertexClassesSharedGenes_HFDvsChow,
  edgeF = "Lilr4b",
  treat1 = "HFD",
  treat2 = "Chow",
```

```
factorOrder = c("HFD", "Chow"),
EdgeFeatureType = "Gene")
```

---

create\_GlobalBarplot *Vertex-class pairs profile of shared features*

---

### Description

This function will create a barplot from the output of Compare\_VertexClasses\_sharedEdgeFeatures using all shared Edge Features (e.g., genes).

### Usage

```
create_GlobalBarplot(
  CompTreatTable,
  treat1 = "Treatment1",
  treat2 = "Treatment2",
  factorOrder = NULL,
  col1 = "red",
  col2 = "blue",
  maxpairs = 1,
  xlb = "Vertex-Class Pairs",
  ylb = "Number of pairs",
  szggrepel = 3.5,
  nudgex = 0.5,
  nudgex = 0.5,
  szaxisTxt = 12,
  szaxisTitle = 12
)
```

### Arguments

CompTreatTable	Output of Compare_VertexClasses_sharedEdgeFeatures
treat1	Name of treatment one, default Treatment1. It should match the column names of the output of Compare_VertexClasses_sharedEdgeFeatures
treat2	Name of treatment one, default Treatment2. It should match the column names of the output of Compare_VertexClasses_sharedEdgeFeatures
factorOrder	A list specifying the order of the treatments.
col1	Color for Treatment 1
col2	Color for Treatment 2
maxpairs	If number of class-vertex-pairs > maxpairs, display number pairs on top of bar
xlb	Name for x-axis
ylb	Name for the y-axis
szggrepel	Size ggrepel labels

nudgey	Nudge y ggrepel
nudgex	Nudge x ggrepel
szaxisTxt	Size axis text
szaxisTitle	Size axis title

**Value**

A ggplot object for a barplot. The barplot shows the vertex-class pairs profile of all shared edge features between treatments

**Examples**

```
data(VertexClassesSharedGenes_HFDvsChow)
create_GlobalBarplot(CompTreatTable = VertexClassesSharedGenes_HFDvsChow,
  treat1 = "HFD",
  treat2 = "Chow",
  factorOrder = c("HFD", "Chow"),
  col1="red",
  col2 ="blue",
  maxpairs = 1,
  szggrepel = 6,
  szaxisTxt = 15,
  szaxisTitle = 15,
  xlb = "Metabolite-pair classes")
```

---

```
create_stackedGlobalBarplot_perTreatment
  Stacked Global Barplot (One treatment)
```

---

**Description**

This function will create a stacked barplot from the output of `Compare_VertexClasses_sharedEdgeFeatures` using all shared Edge Features (e.g., genes) between two treatments.

**Usage**

```
create_stackedGlobalBarplot_perTreatment(
  CompTreatTable,
  treat = NULL,
  xlb = "Vertex-Class Pairs",
  ylb = "Number of pairs",
  max_pairsLegend = 2,
  mx.overlaps = Inf,
  szggrepel = 6,
  force = 0.1,
  szTitle = 12,
  szaxisTxt = 12,
```

```

    szaxisTitle = 12,
    ylim = NULL
)

```

### Arguments

CompTreatTable	Output of Compare_VertexClasses_sharedEdgeFeatures
treat	Name of treatment to display. It should match the column name in the output of Compare_VertexClasses_sharedEdgeFeatures
xlb	Name for x-axis
ylb	Name for y-axis
max_pairsLegend	If number of Edge Features $\geq$ max_pairsLegend, display number of Edge Features as label with ggrepel
mx.overlaps	Max number of overlaps ggrepel
szggrepel	Size ggrepel labels
force	Repelling force for ggrepel labels
szTitle	Size Title
szaxisTxt	Size axis text
szaxisTitle	Size axis titles
ylim	Optional y-limits of the plot

### Value

A ggplot object to create a stacked barplot. The stacked barplot shows the vertex-class pairs profile of all shared edge features but restricted to a single treatment. Every bar consists of multiple edge features (stacked) that are represented with different colors

### Examples

```

data(VertexClassesSharedGenes_HFDvsChow)
create_stackedGlobalBarplot_perTreatment(CompTreatTable = VertexClassesSharedGenes_HFDvsChow,
                                         treat = "HFD",
                                         max_pairsLegend = 9,
                                         xlb = "Metabolite-class-pairs")

```

---

find\_localControllingFeatures

*Find local controlling features*

---

### Description

This function applies for a selected subnetwork a binomial test using the frequency of appearance of an edge feature and the total number of edge features. The probability corresponds to  $1/n_{df}$ , where  $n_{df}$  corresponds to the total number of edge features in the network. The selected subnetwork corresponds to the second level neighborhood of a specific node. The test is applied to all possible second level neighborhoods in the network.

**Usage**

```
find_localControllingFeatures(ResultsCoNI, network, padjust = TRUE)
```

**Arguments**

ResultsCoNI	The output of CoNI (after p-adjustment)
network	Network created with the function generate_network
padjust	logical. Filter output based on adjusted p values

**Value**

Returns a data.frame with the results of the binomial tests. Significant results correspond to local controlling features

**Examples**

```
#Load color nodes table
data(MetColorTable)

#Assign colors according to "Class" column
MetColorTable<-assign_colorsAnnotation(MetColorTable)

#Load CoNI results
data(CoNIResultsHFDToy)

#Generate Network
#Note: Colors not visible when plotting in Igraph
HFDNetwork<-generate_network(ResultsCoNI = CoNIResultsHFDToy,
                             colorVertexNetwork = TRUE,
                             colorVertexTable = MetColorTable,
                             Class = MetColorTable,
                             outputDir = "./",
                             outputFileName = "HFD",
                             saveFiles = FALSE)

#Note: For this tiny example nothing is significant
LCG_BinomialTestTableHFD<- find_localControllingFeatures(ResultsCoNI = CoNIResultsHFDToy,
                                                         network = HFDNetwork )
LCGenes_HFD<-as.character(unique(LCG_BinomialTestTableHFD$edgeFeatures))
```

---

GeneExpToy

*Toy data gene expression*


---

**Description**

Toy data gene expression

**Author(s)**

Jose Manuel Monroy <no1ozz@gmail.com>

**References**

[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)

---

generate_network	<i>Create network</i>
------------------	-----------------------

---

**Description**

This function creates a network using as input the output of CoNI and a table specifying the colors for the nodes.

**Usage**

```
generate_network(
  ResultsCoNI,
  colorVertexNetwork = TRUE,
  colorVertexTable,
  outputDir = "./",
  outputFileName = "ResultsCoNI",
  Class = NULL,
  saveFiles = TRUE
)
```

**Arguments**

ResultsCoNI	The input of the function are the results of CoNI.
colorVertexNetwork	logical. If TRUE, the table colorVertexTable has to be provided to specify vertex colors
colorVertexTable	Table specifying the colors for the nodes (vertex features). The first column should contain the names matching the features of the vertex Data and the colors or other data can be specified in the rest of the columns
outputDir	Output directory where the network is saved as well as the file that was used to generate the network.
outputFileName	The name of the file used to create the network.
Class	Optional data frame with at least two columns, first column contains all vertex features and another column the vertex feature class (column named "Class"). Necessary for treatment comparisons based on class
saveFiles	logical. If FALSE TableForNetwork_‘outputFileName’.csv and Network_‘outputFileName’.graphml are not saved to disk

**Value**

Returns an igraph object (network) constructed from ResultsCoNI. The network includes the following network statistics

- "degree"The number of the vertex adjacent edges
- "hub\_score"The principal eigenvector of  $A \cdot t(A)$ , where A is the adjacency matrix of the graph
- "transitivity"Probability that the adjacent vertices of a vertex are connected
- "closeness"Steps required to access every other vertex from a given vertex
- "betweenness"(roughly) The number of geodesics (shortest paths) going through a vertex or an edge
- "eigen\_centrality"Takes a graph (graph) and returns the eigenvector centralities of positions v within it
- "centralized\_betweenness"The vertice-level centrality score according to the betweenness of vertices
- "centralized\_closeness"The vertice-level centrality score according to the closeness of vertices
- "centralized\_degree"The vertice-level centrality score according to the degrees of vertices

For more details see igraph package

**Examples**

```
#Generate Network

#Load color nodes table
data(MetColorTable)
#Assign colors according to "Class" column
MetColorTable<-assign_colorsAnnotation(MetColorTable)
#Load CoNI results
data(CoNIResultsHFDToy)

#Generate Network
HFDNetwork<-generate_network(ResultsCoNI = CoNIResultsHFDToy,
                             colorVertexNetwork = TRUE,
                             colorVertexTable = MetColorTable,
                             outputDir = "./",
                             outputFileName = "HFD",
                             saveFiles = FALSE)
```

---

```
getstackedGlobalBarplot_and_Grid
```

*Stacked Global Barplot Side-by-side (two treatments)*

---

**Description**

This function will create a stacked barplot from the output of Compare\_VertexClasses\_sharedEdgeFeatures using all shared Edge Features (e.g., genes) between two treatments. Results of both Treatments are side by side for better comparison.



**Usage**

```
getstackedGlobalBarplot_and_Grid(
  CompTreatTable,
  Treat1,
  Treat2,
  ggrep = TRUE,
  max_pairsLegend = 1,
  force = 0.1,
  mx.overlaps = Inf,
  szggrepel = 6,
  xlb = "Vertex-Class Pairs",
  ...
)
```

**Arguments**

CompTreatTable	Output of Compare_VertexClasses_sharedEdgeFeatures
Treat1	Name treatment 1 as in table CompTreatTable
Treat2	Name treatment 2 as in table CompTreatTable
ggrep	logical. If TRUE includes ggrepel labels for every bar
max_pairsLegend	If number of Edge Features $\geq$ max_pairsLegend, display number of Edge Features as ggrepel label
force	Repelling force for ggrepel labels
mx.overlaps	Max number of overlaps ggrepel
szggrepel	Size ggrepel labels
xlb	Name for x-axis
...	Other parameters for inner functions, mainly ggplot2 visual parameters

**Value**

A gtable containing stacked barplots. The barplots show the vertex-class pairs profile of all shared edge features between two treatments (one bar plot per treatment). Every bar consists of multiple edge features that are depicted with different colors

**Examples**

```
data(VertexClassesSharedGenes_HFDvsChow)
VCSGs<-VertexClassesSharedGenes_HFDvsChow
HFD_vs_Chow_stackedBarplot<-getstackedGlobalBarplot_and_Grid(VCSGs,
  Treat1 = "HFD",
  Treat2 = "Chow",
  xlb = "Metabolite-class-pairs",
  max_pairsLegend=9)

plot(HFD_vs_Chow_stackedBarplot)
```

---

```
getVertexsPerEdgeFeature
```

*Vertex Class profile per edge feature (one treatment)*

---

### Description

This function creates a barplot or barplots showing the number of vertex features per class for every shared edge feature between two treatments

### Usage

```
getVertexsPerEdgeFeature(
  CompTreatTable,
  Annotation,
  chunks = 5,
  treat = NULL,
  small = FALSE,
  ggrep = TRUE,
  xlb = "Gene",
  onlyTable = FALSE,
  szTitle = 12,
  szaxisTxt = 12,
  szaxisTitle = 12,
  ...
)
```

### Arguments

CompTreatTable	Output of Compare_VertexClasses_sharedEdgeFeatures
Annotation	Data frame that includes the rgb colors for every class. The column 'class' (or 'Class') has to be present and also the column 'ColorRgb'
chunks	To avoid a non readable dense plot the results can be spitted in multiple plots
treat	Specify the treatment for which the plot will be created. It should be one of the two treatments in the output of Compare_VertexClasses_sharedEdgeFeatures
small	logical. If only a few edge features are in the input set as TRUE. A single plot will be created
ggrep	logical. If TRUE includes ggrepel labels for every bar
xlb	x-axis label
onlyTable	logical. If TRUE a table is returned instead of a plot
szTitle	Size title
szaxisTxt	Size axis text
szaxisTitle	Size axis title
...	Other parameters for inner functions, mainly ggplot2 visual parameters

**Value**

A list of ggplot objects to create different barplots. The barplots show the number of vertex features per class for every shared edge feature between two treatments. The barplots restrict to one of the compared treatments. An alternative output is a data.frame with the number of vertex features per class and edge feature (onlyTable=TRUE)

**Examples**

```
data(VertexClassesSharedGenes_HFDvsChow)
data(MetColorTable)
#Note: No differences in example as all the Output of CoNI was kept
getVertexsPerEdgeFeature(CompTreatTable = VertexClassesSharedGenes_HFDvsChow,
                          Annotation = MetColorTable,
                          chunks = 2,
                          treat = "HFD")
```

---

```
getVertexsPerEdgeFeature_and_Grid
```

*Vertex-Class profile per edge feature Side-by-Side (two treatments)*

---

**Description**

This function creates a grid of barplots. The barplot of one side depicts the number of class vertex features per edge feature for treatment 1 and the other side the same barplot for treatment 2. Results of both Treatments are side by side for better comparison.

**Usage**

```
getVertexsPerEdgeFeature_and_Grid(
  CompTreatTable,
  Treat1,
  Treat2,
  Annotation,
  chunks = 3,
  ggrep = TRUE,
  xlb = "Edge Feature",
  onlyT = FALSE,
  small = FALSE,
  ...
)
```

**Arguments**

CompTreatTable	Output of Compare_VertexClasses_sharedEdgeFeatures
Treat1	Name treatment 1 as in table CompTreatTable
Treat2	Name treatment 2 as in table CompTreatTable

Annotation	Data frame that includes the rgb colors for every class. The column 'class' (or 'Class') has to be present and also the column 'ColorRgb'
chunks	To avoid a non readable dense plot the results can be spitted in multiple plots
ggrep	logical. If TRUE includes ggrepel labels for every bar
x1b	Change the x-axis label
onlyT	logical. If TRUE a table is returned instead of a grid of plots
small	logical. If only a few edge features are in the input set as TRUE. A single plot will be created
...	Other parameters for inner functions, mainly ggplot2 visual parameters

**Value**

A gtable containing side-by-side barplots, one for each treatment, showing the number of vertex features per class for every shared edge feature

**Examples**

```
data(VertexClassesSharedGenes_HFDvsChow)
VCSGs<-VertexClassesSharedGenes_HFDvsChow
data(MetColorTable)
HFD_vs_Chow_LCP_Gene<-getVertexsPerEdgeFeature_and_Grid(VCSGs,
                                                         "HFD", "Chow",
                                                         Annotation=MetColorTable,
                                                         ggrep=FALSE,
                                                         small = FALSE,
                                                         chunks = 3,
                                                         szLegendKey=0.2)

plot(HFD_vs_Chow_LCP_Gene)
```

---

HFD\_GeneExpData

*HFD gene expression data*


---

**Description**

HFD gene expression data

**Author(s)**

Jose Manuel Monroy <no1ozz@gmail.com>

**References**

[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)

---

HFD\_MetaboliteData      *HFD metabolite data*

---

**Description**

HFD metabolite data

**Author(s)**

Jose Manuel Monroy <nolozz@gmail.com>

**References**

[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)

---

MetaboExpToy      *Toy data metabolite expression*

---

**Description**

Toy data metabolite expression

**Author(s)**

Jose Manuel Monroy <nolozz@gmail.com>

**References**

[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)

---

MetaboliteAnnotation      *Metabolite Annotation*

---

**Description**

Metabolite Annotation

**Author(s)**

Jose Manuel Monroy <nolozz@gmail.com>

**References**

[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)

MetColorTable

*Toy data annotation*

---

**Description**

Toy data annotation

**Author(s)**

Jose Manuel Monroy <no1ozz@gmail.com>

**References**

[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)

---

NetStats

*Network Statistics*

---

**Description**

This function calculates simple network statistics and returns them as a dataframe

**Usage**

```
NetStats(Network)
```

**Arguments**

Network            An Igraph network

**Value**

Returns a data.frame with nine rows with the following network statistics:

- "net\_avg\_pathL" Shortest paths between vertices
- "net\_edge\_density" Graph density, ratio of the number of edges and the number of possible edges
- "net\_transitivity" Probability that the adjacent vertices of a vertex are connected
- "net\_diameter" Length of the longest geodesic
- "net\_nodes\_first\_path\_diameter" The nodes along the first found path with the length of diameter
- "net\_eigenvalue" The eigenvalue corresponding to the centrality scores.
- "net\_centralized\_betweennessIdx" The graph level centrality index after centralizing the graph according to the betweenness of vertices

- "net\_centralized\_closenessIdx" The graph level centrality index after centralizing the graph according to the closeness of vertices
- "net\_centralized\_degreeIdx" The graph level centrality index after centralizing the graph according to the degrees of vertices

For more information on the statistics consult the igraph package.

### Examples

```
#Load color nodes table
data(MetColorTable)
#Assign colors according to "Class" column
MetColorTable<-assign_colorsAnnotation(MetColorTable)
#Load CoNI results
data(CoNIResultsHFDToy)

#Generate Network
HFDNetwork<-generate_network(ResultsCoNI = CoNIResultsHFDToy,
                             colorVertexNetwork = TRUE,
                             colorVertexTable = MetColorTable,
                             outputDir = "./",
                             outputFileName = "HFD",
                             saveFiles = FALSE)

NetStats(HFDNetwork)
```

---

plotPcorvsCor

*Correlation vs Partial correlation*

---

### Description

This function fits two linear models on standardized data and plots the results. It generates a scatter plot with two regression lines, where the slopes correspond to the correlation and partial correlation coefficients (blue for cor and red for pcor)

### Usage

```
plotPcorvsCor(
  ResultsCoNI,
  edgeFeature,
  vertexD,
  edgeD,
  vertexFeatures = NULL,
  outputDir = "./",
  fname,
  label_edgeFeature = "Edge Feature",
  plot_to_screen = TRUE,
  height = 10,
  width = 8,
  saveFiles = FALSE
)
```

**Arguments**

ResultsCoNI	The significant results generated by CoNI
edgeFeature	The edge feature to explore e.g. Fabp2 (for a gene)
vertexD	Vertex data that was given as input to CoNI
edgeD	Edge data that was given as input to CoNI
vertexFeatures	The vertex features to include as a list. If not specified all metabolites available in combination with the edgeFeature will be used
outputDir	Output directory with path
fname	File name to save the plots
label_edgeFeature	Name for plot title e.g. Gene or Protein
plot_to_screen	logical. If TRUE plots will be outputted to the plotting screen
height	height of the plotting area for the saved file
width	width of the plotting are for the saved file
saveFiles	logical. If FALSE plot is not saved to disk

**Value**

Returns a ggplot object for a scatter plot with two regression lines. The blue line is the regression of the vertex features, and the red line is the regression of the resulting residuals after regressing each vertex feature with the edge feature. The slope of the blue line corresponds to the pearson correlation coefficient and the slope of the red line to the partial correlation coefficient

**Examples**

```
#Load gene expression - Toy dataset of two treatments
data(GeneExpToy)
#Samples in rows and genes in columns
GeneExp <- as.data.frame(t(GeneExpToy))
hfd_gene <- GeneExp[1:8,] #high fat diet
chow_gene<- GeneExp[9:nrow(GeneExp),] #chow diet

#Load metabolite expression - Toy dataset of two treatments
data(MetaboExpToy)
MetaboExp <- MetaboExpToy
hfd_metabo <- MetaboExp[11:18,] #high fat diet
chow_metabo <- MetaboExp[1:10,] #chow diet

#Match row names both data sets
rownames(hfd_metabo)<-rownames(hfd_gene)
rownames(chow_metabo)<-rownames(chow_gene)

#Load CoNI results
data(CoNIResultsHFDToy)

plotPcorvsCor(ResultsCoNI = CoNIResultsHFDToy,
              edgeFeature = "Arfrp1",
```



```

vertexFeatures = c("PC.ae.C40.2", "SM..OH..C22.1"),
vertexD = hfd_metabo,
edgeD = hfd_gene,
label_edgeFeature = "Gene",
plot_to_screen = TRUE,
height = 10,
saveFiles = FALSE)

```

---

tableLCFs_VFs	<i>Table local controlling edge features and vertex pairs</i>
---------------	---

---

### Description

This function creates a table of the local controlling edge features

### Usage

```
tableLCFs_VFs(CoNIResults, LCFs)
```

### Arguments

CoNIResults	The output of CoNI (after p-adjustment)
LCFs	Local controlling edge features as a vector

### Value

A data.frame of local controlling edge features and their respective vertex pairs, and unique vertexes.

### Examples

```

#Load CoNI results
data(CoNIResultsHFDToy)
#Note: arbitrary list of genes, not Local controlling features
tableLCFs_VFs(CoNIResultsHFDToy, c("Lilr4b", "Rps12"))

```

---

top_n_LF_byMagnitude	<i>Linker Features by magnitude of effect</i>
----------------------	---

---

### Description

This function outputs the linker features with the strongest effect on the correlation of the vertex features

### Usage

```
top_n_LF_byMagnitude(ResultsCoNI, topn = 10)
```

**Arguments**

ResultsCoNI	The output of CoNI
topn	Top n number of features to output

**Value**

Returns a data.frame, a filtered version of ResultsCoNI, showing the top n features with the strongest effect, that is, the highest difference between the partial correlation and correlation coefficient.

**Examples**

```
data(CoNIResultsHFDToy)
Top10HFD<-top_n_LF_byMagnitude(CoNIResults_HFD, topn = 10)
```

---

VertexClassesSharedGenes\_HFDvsChow  
*Toy data comparison treatments*

---

**Description**

Toy data comparison Vertex classes shared edge featuresHFD vs Chow

**Author(s)**

Jose Manuel Monroy <nolozz@gmail.com>

**References**

[doi:10.1016/j.molmet.2021.101295](https://doi.org/10.1016/j.molmet.2021.101295)

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