# Package 'Sysrecon'

July 21, 2025

```
Description In the past decade, genome-scale metabolic reconstructions have widely been
     used to comprehend the systems biology of metabolic pathways within an organism.
     Different GSMs are constructed using various techniques that
     require distinct steps, but the input data, information conversion and software tools
     are neither concisely defined nor mathematically or programmatically formulated
     in a context-specific manner. The tool that quantitatively and qualitatively specifies
     each reconstruction steps and can generate a template list of reconstruction steps dynamically
     selected from a reconstruction step reservoir, constructed based on all available published papers.
License GPL-3
Imports ape, dplyr, ggplot2, ggtree, magrittr, methods, patchwork,
     plyr, RColorBrewer, rlang, SnowballC, stats, stringr, tm, utils
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2 contentTypes

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# Description

A data frame contains the labels and groups of the contents of metabolic reconstruction

# Usage

 ${\tt contentTypes}$ 

# **Format**

label the contents of the metabolic reconstructiongroup the classification of the labels

conversionMatrix 3

	conversionMatrix	A data frame contains the details of the transformation of the metabolic reconstruction
--	------------------	---

## **Description**

A data frame contains the details of the transformation of the metabolic reconstruction

#### Usage

conversionMatrix

#### **Format**

A data frame contains 93 rows and 67 variables:

MarkerWords the key words of the steps of metabolic reconstruction

ThresholdValue the values filter the steps after the mapping

**Steps** the steps of metabolic reonstruction

Group the classification of the steps

**SpeciesName** 1 means the output of Species name.

**TaxonID** 1 means the output of Taxon ID.

**NCBI** 1 means the output of NCBI.

**Uniprot** 1 means the output of Uniprot.

**KEGG** 1 means the output of KEGG.

MetaCyc 1 means the output of MetaCyc.

**PubChem** 1 means the output of PubChem.

**Brenda** 1 means the output of Brenda.

**Download** 1 means the output of Download.

**GeneticInformation** 1 means the output of Genetic information.

**ProteinInformation** 1 means the output of Protein information.

**GenomeSequence** 1 means the output of Genome sequence.

**ProteinSequence** 1 means the output of Protein sequence.

**MetabolicFunctionInformation** 1 means the output of Metabolic function information.

**Metabolites** 1 means the output of Metabolites.

**Cofactor** 1 means the output of Cofactor.

**Nucleotides** 1 means the output of Nucleotides.

AminoAcid 1 means the output of Amino acid.

MolecularWeight 1 means the output of Molecular weight.

**DryWeight** 1 means the output of Dry weight.

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**MetabolicReaction** 1 means the output of Metabolic reaction.

**TerminalMetabolite** 1 means the output of Terminal metabolite.

**Secretion** 1 means the output of Secretion.

**BiomassReaction** 1 means the output of Biomass reaction.

**DemandReaction** 1 means the output of Demand reaction.

**SinkReaction** 1 means the output of Sink reaction.

**GapReaction** 1 means the output of Gap reaction.

**SpontaneousReaction** 1 means the output of Spontaneous reaction.

**Extracellular And Periplasmic Transport Reactions** 1 means the output of Extracellular and periplasmic transport reactions reaction.

**ExchangeReaction** 1 means the output of Exchange reaction.

Intracellular Transport Reaction 1 means the output of Intracellular transport reaction.

**ReactionFlux** 1 means the output of Reaction flux.

**GPR** 1 means the output of GPR.

**BlastComparison** 1 means the output of Blast comparison.

**Homology** 1 means the output of Homology.

**HomologousGene** 1 means the output of Homology gene.

StoichiometricMatrix 1 means the output of Stoichiometric matrix.

**Knockout** 1 means the output of Knockout.

**TargetReaction** 1 means the output of Target reaction.

**Restrictions** 1 means the output of Restrictions.

**GrowthConditions** 1 means the output of Growth conditions.

MinORMax 1 means the output of min | max.

**ReconstructionData** 1 means the output of Reconstruction data.

**FVA** 1 means the output of FVA.

MetabolicFlux 1 means the output of Metabolic flux.

**Statistics** 1 means the output of Metabolic Statistics.

NeutralMolecularFormula 1 means the output of Neutral molecular formula.

ChargedMolecularFormula 1 means the output of Charged molecular formula.

**LiteratureDataCollection** 1 means the output of Literature data collection.

**ConservationOfMassAndCharge** 1 means the output of Conservation of mass and charge.

**GibbsFreeEnergy** 1 means the output of Gibbs free energy.

**CellCompartmentInformation** 1 means the output of Cell compartment information.

**SubsystemInformation** 1 means the output of Subsystem information.

**MetaboliteIdentification** 1 means the output of Metabolite Identification.

**Unite** 1 means the output of Unite.

**ManualPlanning** 1 means the output of Manual planning.

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**Coefficient** 1 means the output of Coefficient.

**ScatterPlot** 1 means the output of Scatter plot.

**TestReport** 1 means the output of Test Report.

**TypeIIIPath** 1 means the output of Type III path.

**SBML** 1 means the output of SBML file.

Mat 1 means the output of Mat file.

**Excel** 1 means the output of Excel file.

 ${\it conversion Types} \qquad {\it A data frame contains the labels and groups of the transformation of }$ 

metabolic reconstruction

#### **Description**

A data frame contains the labels and groups of the transformation of metabolic reconstruction

#### Usage

conversionTypes

#### Format

A data frame contains 63 rows and 2 variables:

**label** the transformation of the metabolic reconstruction

group the classification of the labels

draw\_conversion\_tree

#### **Description**

Drawing of metabolic process matrix diagram and phylogenetic tree diagram

#### Usage

draw\_conversion\_tree(infomatrix, Matrix, stepTypes, conversionTypes)

#### **Arguments**

infomatrix Matrix generated using the words2steps function

Matrix The matrix about the step or transformation or databases and tools used in the

metabolic reconstruction

stepTypes Grouping information for reconstruction processes

conversionTypes

Grouping information for conversion content

draw\_step\_tree

## Value

a plot

# **Examples**

```
p1 <- draw_conversion_tree(matrixProcessConversion, conversionMatrix,
    stepTypes, conversionTypes)
```

draw\_step\_tree

draw\_step\_tree

# Description

Drawing of metabolic process matrix diagram and phylogenetic tree diagram

# Usage

```
draw_step_tree(infomatrix, Matrix, stepTypes, contentTypes)
```

# Arguments

infomatrix Matrix generated using the words2steps function

Matrix The matrix about the steps or transformation or the databases and tools of the

metabolic reconstruction.

stepTypes Grouping information for metabolic processes contentTypes Grouping information for metabolic content

#### Value

a plot

# **Examples**

```
p1 <- draw_step_tree(matrixProcess, stepsMatrix, stepTypes, contentTypes)</pre>
```

get\_term\_matrix 7

get\_term\_matrix

get\_term\_matrix

# Description

```
get\_term\_matrix
```

# Usage

```
get_term_matrix(data)
```

# **Arguments**

data

information of data text

## Value

words matrix

# Examples

```
wordsMatrix <- get_term_matrix(text)</pre>
```

inputTxt

inputTxt

# Description

A variable containing the contents of the metabolic reconstruction:

# Usage

inputTxt

## **Format**

A data frame with characters:

V1 the contents of the metabolic reconstruction

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map\_word\_to\_step

map\_word\_to\_step

#### **Description**

```
map_word_to_step
```

#### Usage

```
map_word_to_step(wordsMatrix, stepsMatrix)
```

#### **Arguments**

wordsMatrix The word matrix generated by the function TermDocumentMatrix in the tm

package.

stepsMatrix Manually constructed metabolic process matrix.

#### Value

Specific metabolic process matrix based on text content mapping.

## **Examples**

```
matrixProcess <- map_word_to_step(wordsMatrix, stepsMatrix)</pre>
```

matrixProcess

A data frame produced by the function map\_word\_to\_step

#### **Description**

A data frame produced by the function map\_word\_to\_step

#### Usage

matrixProcess

#### **Format**

A data frame contains 67 variables

degree the number of the steps used in a article.

SpeciesName -1 means the input of Species name. 1 means the output of Species name.

**TaxonID** -1 means the input of Taxon ID. 1 means the output of Taxon ID.

**KEGG** -1 means the input of KEGG reaction. 1 means the output of KEGG reaction.

NCBI -1 means the input of NCBI gene. 1 means the output of NCBI gene.

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Uniprot -1 means the input of Uniprot gene. 1 means the output of Uniprot.

**MetaCyc** -1 means the input of MetaCyc gene. 1 means the output of MetaCyc.

**PubChem** -1 means the input of PubChem molecular formula. 1 means the output of PubChem molecular formula.

**Brenda** identify whether input the Brenda. -1 means the input of Brenda. 1 means the output of Brenda.

Chebi -1 means the input of Chebi information. 1 means the the output of Chebi information.

**WebGCM** -1 means the input of Web GCM. 1 means the the output of Web GCM.

**SpontaneousReaction** -1 means the input of spontaneous reaction. 1 means the output of spontaneous reaction.

**ExtracellularAndPeriplasmicTransportReactions** -1 means the input of Extracellular and periplasmic transport reactions. 1 means the output of Extracellular and periplasmic transport reactions.

**ExchangeReaction** -1 means the input of exchange reaction. 1 means the the output of exchange reaction.

**MissingExchangeReaction** -1 means the input of missing exchange reaction. 1 means the the output of missing exchange reaction.

**IntracellularTransportReaction** -1 means the input of intracellular transport reaction. 1 means the the output of intracellular transport reaction.

Gene -1 means the input of gene. 1 means the the output of gene.

**Protein** -1 means the input of protein. 1 means the the output of protein.

**Knockout** -1 means the input of knockout. 1 means the the output of knockout.

**StoichiometricMatrix** -1 means the input of stoichiometric matrix. 1 means the the output of stoichiometric matrix.

**ObjectiveReaction** -1 means the input of objective reaction. 1 means the output of objective reaction.

**Constraints** -1 means the input of constraints. 1 means the the output of constraints.

**Secretion** -1 means the input of secretion. 1 means the output of secretion.

**Mutisecretion** -1 means the input of mutisecretion. 1 means the output of mutisecretion.

RichMedia -1 means the input of rich media. 1 means the the output of rich media.

**GenomeSequence** -1 means the input of genome sequence. 1 means the output of genome sequence.

**ProteomeSequence** -1 means the input of proteome sequence. 1 means the the output of proteome sequence.

**AminoAcidWeight** -1 means the input of amino acid weight. 1 means the the output of amino acid weight.

**AminoAcidMolecularWeight** -1 means the input of amino acid molecular weight. 1 means the the output of amino acid molecular weight.

**NucleotideWeight** -1 means the input of nucleotide weight. 1 means the the output of nucleotide weight.

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**NucleotideMolecularWeight** -1 means the input of nucleotide molecular weight. 1 means the the output of nucleotide molecular weight.

DryWeight -1 means the input of dry weight. 1 means the the output of dry weight.

**BiomassReaction** -1 means the input of biomass reaction. 1 means the the output of biomass reaction.

**DemandReaction** -1 means the input of demand reaction. 1 means the the output of demand reaction.

SinkReaction -1 means the input of sink reaction. 1 means the the output of sink reaction.

**GapReaction** -1 means the input of gap reaction. 1 means the the output of gap reaction.

MinORMax -1 means the input of gap reaction. 1 means the the output of gap reaction.

**GeneticInformation** -1 means the input of genetic information. 1 means the output of genetic information.

**MetabolicFunction** -1 means the input of metabolic function. 1 means the the output of metabolic function.

**Metabolites** -1 means the input of metabolites. 1 means the output of metabolites.

**BiomassMetabolites** -1 means the input of biomass metabolites. 1 means the the output of biomass metabolites.

**MetabolicReaction** -1 means the input of metabolic reaction. 1 means the the output of metabolic reaction.

**ReconstructionData** -1 means the input of reconstruction data. 1 means the the output of reconstruction data.

**DataStatistics** -1 means the input of data statistics. 1 means the the output of data statistics.

**NeutralMolecularFormula** -1 means the input of neutral molecular formula. 1 means the the output of neutral molecular formula.

**ChargedMolecularFormula** -1 means the input of charged molecular formula. 1 means the the output of charged molecular formula.

**ConservationOfMassAndCharge** -1 means the input of conservation of mass and charge. 1 means the the output of conservation of mass and charge.

**GibbsFreeEnergyInformation** -1 means the input of gibbs free energy information. 1 means the the output of gibbs free energy information.

**Mass\_ChargeConservationAssessment** -1 means the input of mass-charge conservation assessment. 1 means the the output of conservation assessment.

**CellularCompartment** -1 means the input of cellular compartment. 1 means the the output of cellular compartment.

**SubsystemInformation** -1 means the input of subsystem information. 1 means the the output of subsystem information.

**IdentifiersInKEGG** -1 means the input of identifiers in KEGG. 1 means the output of identifiers in KEGG.

**IdentifiersInMetaCyc** -1 means the input of identifiers in MetaCyc. 1 means the output of identifiers in MetaCyc.

**UniformIdentifier** -1 means the input of uniform identifier. 1 means the the output of uniform identifier.

matrixProcessConversion 11

**Coefficient** -1 means the input of coefficient. 1 means the the output of coefficient.

**ScatterPlot** -1 means the input of scatter plot. 1 means the the output of scatter plot.

**TerminalMetabolite** -1 means the input of terminal metabolite. 1 means the the output of terminal metabolite.

**TypeIIIPathway** -1 means the input of Type III pathway. 1 means the output of Type III pathway.

NetworkGap -1 means the input of network gap. 1 means the the output of network gap.

**Growth** -1 means the input of growth. 1 means the the output of growth.

BlockReaction -1 means the input of block reaction. 1 means the the output of block reaction.

MetabolicFlux -1 means the input of metabolic flux. 1 means the the output of metabolic flux.

ModelPredictCorrectly 1 means the output of model PredictCorrectly.

**ModelGrowingTooFast** -1 means the input of model growing too fast. 1 means the the output of model growing too fast.

SBML -1 means the input of SBML file. 1 means the the output of SBML file.

Mat -1 means the input of Mat file. 1 means the the output of Mat file.

Excel -1 means the input of Excel file. 1 means the the output of Excel file.

matrixProcessConversion

A data frame produced by the function map\_word\_to\_step

### **Description**

A data frame produced by the function map\_word\_to\_step

#### Usage

matrixProcessConversion

#### **Format**

A data frame contains 64 variables

**degree** the number of the steps used in a article.

**SpeciesName** 1 means the output of Species name.

**TaxonID** 1 means the output of Taxon ID.

**NCBI** 1 means the output of NCBI.

**Uniprot** 1 means the output of Uniprot.

**KEGG** 1 means the output of KEGG.

**MetaCyc** 1 means the output of MetaCyc.

**PubChem** 1 means the output of PubChem.

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**Brenda** 1 means the output of Brenda.

**Download** 1 means the output of Download.

**GeneticInformation** 1 means the output of Genetic information.

**ProteinInformation** 1 means the output of Protein information.

**GenomeSequence** 1 means the output of Genome sequence.

**ProteinSequence** 1 means the output of Protein sequence.

**MetabolicFunctionInformation** 1 means the output of Metabolic function information.

Metabolites 1 means the output of Metabolites.

**Cofactor** 1 means the output of Cofactor.

Nucleotides 1 means the output of Nucleotides.

AminoAcid 1 means the output of Amino acid.

Molecular Weight 1 means the output of Molecular weight.

**DryWeight** 1 means the output of Dry weight.

**MetabolicReaction** 1 means the output of Metabolic reaction.

**TerminalMetabolite** 1 means the output of Terminal metabolite.

**Secretion** 1 means the output of Secretion.

**BiomassReaction** 1 means the output of Biomass reaction.

**DemandReaction** 1 means the output of Demand reaction.

**SinkReaction** 1 means the output of Sink reaction.

**GapReaction** 1 means the output of Gap reaction.

**SpontaneousReaction** 1 means the output of Spontaneous reaction.

**Extracellular And Periplasmic Transport Reactions** 1 means the output of Extracellular and periplasmic transport reactions reaction.

**ExchangeReaction** 1 means the output of Exchange reaction.

**Intracellular Transport Reaction** 1 means the output of Intracellular transport reaction.

**ReactionFlux** 1 means the output of Reaction flux.

**GPR** 1 means the output of GPR.

**BlastComparison** 1 means the output of Blast comparison.

**Homology** 1 means the output of Homology.

HomologousGene 1 means the output of Homology gene.

**StoichiometricMatrix** 1 means the output of Stoichiometric matrix.

**Knockout** 1 means the output of Knockout.

**TargetReaction** 1 means the output of Target reaction.

**Restrictions** 1 means the output of Restrictions.

**GrowthConditions** 1 means the output of Growth conditions.

**MinORMax** 1 means the output of min | max.

**ReconstructionData** 1 means the output of Reconstruction data.

matrixProcessFile 13

**FVA** 1 means the output of FVA.

MetabolicFlux 1 means the output of Metabolic flux.

Statistics 1 means the output of Metabolic Statistics.

**NeutralMolecularFormula** 1 means the output of Neutral molecular formula.

ChargedMolecularFormula 1 means the output of Charged molecular formula.

**LiteratureDataCollection** 1 means the output of Literature data collection.

**ConservationOfMassAndCharge** 1 means the output of Conservation of mass and charge.

GibbsFreeEnergy 1 means the output of Gibbs free energy.

**CellCompartmentInformation** 1 means the output of Cell compartment information.

**SubsystemInformation** 1 means the output of Subsystem information.

**MetaboliteIdentification** 1 means the output of Metabolite Identification.

**Unite** 1 means the output of Unite.

**ManualPlanning** 1 means the output of Manual planning.

**Coefficient** 1 means the output of Coefficient.

**ScatterPlot** 1 means the output of Scatter plot.

**TestReport** 1 means the output of Test Report.

**TypeIIIPath** 1 means the output of Type III path.

**SBML** 1 means the output of SBML file.

Mat 1 means the output of Mat file.

**Excel** 1 means the output of Excel file.

matrixProcessFile

A data frame produced by the function map\_to\_word in the function vizProcess

#### **Description**

A data frame produced by the function map\_to\_word in the function vizProcess

### Usage

matrixProcessFile

#### **Format**

A data frame contains with 3 variables

**step** the steps used in the metabolic reconstruction

degree the number that steps used in the metbolic reconstructoin occur in an article

**step\_ID** the order of the steps used in the metabolic reconstruction

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slotsFunction

slotsFunction

## **Description**

slotsFunction

## Usage

slots(variablesFile)

#### **Arguments**

variablesFile A data frame contains three different variables.

stepsMatrix

A data frame contains the details of the steps of the metabolic reconstruction

#### **Description**

A data frame contains the details of the steps of the metabolic reconstruction

## Usage

stepsMatrix

#### **Format**

A data frame with 93 rows and 70 variables:

MarkerWords the key words of the steps of metabolic reconstruction

Threshold Value the values filter the steps after the mapping

**Steps** the steps of metabolic reonstruction

**Group** the classification of the steps

SpeciesName -1 means the input of Species name. 1 means the output of Species name.

**TaxonID** -1 means the input of Taxon ID. 1 means the output of Taxon ID.

**KEGG** -1 means the input of KEGG reaction. 1 means the output of KEGG reaction.

NCBI -1 means the input of NCBI gene. 1 means the output of NCBI gene.

Uniprot -1 means the input of Uniprot gene. 1 means the output of Uniprot.

MetaCyc -1 means the input of MetaCyc gene. 1 means the output of MetaCyc.

**PubChem** -1 means the input of PubChem molecular formula. 1 means the output of PubChem molecular formula.

stepsMatrix 15

**Brenda** identify whether input the Brenda. -1 means the input of Brenda. 1 means the output of Brenda.

**Chebi** -1 means the input of Chebi information. 1 means the the output of Chebi information.

**WebGCM** -1 means the input of Web GCM. 1 means the the output of Web GCM.

**SpontaneousReaction** -1 means the input of spontaneous reaction. 1 means the the output of spontaneous reaction.

**Extracellular And Periplasmic Transport Reactions** -1 means the input of Extracellular and periplasmic transport reactions. 1 means the output of Extracellular and periplasmic transport reactions.

**ExchangeReaction** -1 means the input of exchange reaction. 1 means the the output of exchange reaction.

**MissingExchangeReaction** -1 means the input of missing exchange reaction. 1 means the the output of missing exchange reaction.

**IntracellularTransportReaction** -1 means the input of intracellular transport reaction. 1 means the the output of intracellular transport reaction.

**Gene** -1 means the input of gene. 1 means the the output of gene.

**Protein** -1 means the input of protein. 1 means the the output of protein.

**Knockout** -1 means the input of knockout. 1 means the the output of knockout.

**StoichiometricMatrix** -1 means the input of stoichiometric matrix. 1 means the the output of stoichiometric matrix.

**ObjectiveReaction** -1 means the input of objective reaction. 1 means the the output of objective reaction.

**Constraints** -1 means the input of constraints. 1 means the the output of constraints.

**Secretion** -1 means the input of secretion. 1 means the the output of secretion.

Mutisecretion -1 means the input of mutisecretion. 1 means the output of mutisecretion.

RichMedia -1 means the input of rich media. 1 means the the output of rich media.

**GenomeSequence** -1 means the input of genome sequence. 1 means the the output of genome sequence.

**ProteomeSequence** -1 means the input of proteome sequence. 1 means the the output of proteome sequence.

**AminoAcidWeight** -1 means the input of amino acid weight. 1 means the the output of amino acid weight.

**AminoAcidMolecularWeight** -1 means the input of amino acid molecular weight. 1 means the the output of amino acid molecular weight.

**NucleotideWeight** -1 means the input of nucleotide weight. 1 means the the output of nucleotide weight.

**NucleotideMolecularWeight** -1 means the input of nucleotide molecular weight. 1 means the the output of nucleotide molecular weight.

**DryWeight** -1 means the input of dry weight. 1 means the the output of dry weight.

**BiomassReaction** -1 means the input of biomass reaction. 1 means the the output of biomass reaction.

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**DemandReaction** -1 means the input of demand reaction. 1 means the the output of demand reaction.

SinkReaction -1 means the input of sink reaction. 1 means the the output of sink reaction.

**GapReaction** -1 means the input of gap reaction. 1 means the the output of gap reaction.

**MinORMax** -1 means the input of gap reaction. 1 means the the output of gap reaction.

**GeneticInformation** -1 means the input of genetic information. 1 means the the output of genetic information.

**MetabolicFunction** -1 means the input of metabolic function. 1 means the output of metabolic function.

**Metabolites** -1 means the input of metabolites. 1 means the output of metabolites.

**BiomassMetabolites** -1 means the input of biomass metabolites. 1 means the the output of biomass metabolites.

**MetabolicReaction** -1 means the input of metabolic reaction. 1 means the the output of metabolic reaction.

**ReconstructionData** -1 means the input of reconstruction data. 1 means the the output of reconstruction data.

DataStatistics -1 means the input of data statistics. 1 means the the output of data statistics.

**NeutralMolecularFormula** -1 means the input of neutral molecular formula. 1 means the the output of neutral molecular formula.

**ChargedMolecularFormula** -1 means the input of charged molecular formula. 1 means the the output of charged molecular formula.

**ConservationOfMassAndCharge** -1 means the input of conservation of mass and charge. 1 means the the output of conservation of mass and charge.

**GibbsFreeEnergyInformation** -1 means the input of conservation of gibbs free energy information. 1 means the output of gibbs free energy information.

**Mass\_ChargeConservationAssessment** -1 means the input of mass-charge conservation assessment. 1 means the the output of mass-charge conservation assessment.

**CellularCompartment** -1 means the input of cellular compartment. 1 means the the output of cellular compartment.

**SubsystemInformation** -1 means the input of subsystem information. 1 means the the output of subsystem information.

**IdentifiersInKEGG** -1 means the input of subsystem information. 1 means the the output of subsystem information.

**IdentifiersInMetaCyc** -1 means the input of identifiers in MetaCyc. 1 means the output of identifiers in MetaCyc.

**UniformIdentifier** -1 means the input of uniform identifier. 1 means the output of uniform identifier.

Coefficient -1 means the input of coefficient. 1 means the the output of coefficient.

**ScatterPlot** -1 means the input of scatter plot. 1 means the the output of scatter plot.

**TerminalMetabolite** -1 means the input of terminal metabolite. 1 means the the output of terminal metabolite.

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**TypeIIIPathway** -1 means the input of Type III pathway. 1 means the the output of Type III pathway.

NetworkGap -1 means the input of network gap. 1 means the the output of network gap.

**Growth** -1 means the input of growth. 1 means the the output of growth.

BlockReaction -1 means the input of block reaction. 1 means the the output of block reaction.

MetabolicFlux -1 means the input of metabolic flux. 1 means the the output of metabolic flux.

**ModelPredictCorrectly** -1 means the input of model predict correctly. 1 means the output of model predict correctly.

**ModelGrowingTooFast** -1 means the input of model growing too fast. 1 means the the output of model growing too fast.

**SBML** -1 means the input of SBML file. 1 means the the output of SBML file.

Mat -1 means the input of Mat file. 1 means the the output of Mat file.

**Excel** -1 means the input of Excel file. 1 means the the output of Excel file.

stepTypes	A data frame contains the labels and groups of the steps of metabolic reconstruction

#### Description

A data frame contains the labels and groups of the steps of metabolic reconstruction

#### Usage

stepTypes

#### Format

A data frame contains 93 rows and 2 variables:

label the steps of the metabolic reconstruction

**group** the classfication of the labels

Sysrecon Sysrecon

|--|

# Description

Input the txt and output the visualization of the steps, transformation and databases and tools.

## Usage

```
Sysrecon(
  inputTxt,
  stepsMatrix,
  stepTypes,
  conversionMatrix,
  conversionTypes,
  toolsMatrix,
  toolsTypes,
  contentTypes
)
```

# Arguments

inputTxt	A txt contains the methods and contents of the metabolic reconstruction in articles.
stepsMatrix	A data frame the marker words, threshold value, steps, group and other information about the metabolic reconstruction. The default file is in the data.
stepTypes	A data frame the labels and groups of the metabolic reconstructions steps. The default file is in the data.
conversionMatri	ix
	A data frame contains the marker words, threshold value, steps, group and other transformation information about the metabolic reconstruction. The default file is in the data.
conversionTypes	3
	A data frame contains the labels and groups of the metabolic reconstructions transformation. The default file is in the data.
toolsMatrix	A data frame contains the marker words, threshold value, steps, group and other information about the metabolic reconstruction databases and tools. The default file is in the data.
toolsTypes	A data frame contains the databases and the tools used in the metabolic reconstruction.
contentTypes	A data frame contains the labels and groups of the metabolic reconstructions content The default file is in the data.

# **Details**

Input takes a data.frame x with two variables v and w and returns the maximum knapsack value and which elements (rows in the data.frame).

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## Value

The pictures that visualize the steps, transformation and databases and tools of the metabolic reconstruction.

### **Examples**

```
exam <- Sysrecon(inputTxt, stepsMatrix, stepTypes, conversionMatrix, conversionTypes,
  toolsMatrix, toolsTypes, contentTypes)</pre>
```

systemrecon-class

systemrecon-class

# Description

systemrecon-class

#### **Usage**

label

#### **Format**

An object of class character of length 72.

#### **Slots**

Taxon\_id Taxonomy ID

NCBI\_gene Gene information from NCBI

Uniprot\_gene Gene information from Uniprot

Genetic\_information Genetic information

KEGG\_reaction Reactions in KEGG

MetaCyc\_reaction Reactions in MetaCyc

Metabolic\_function Metabolic function

Metabolic\_reaction Metabolic reaction

Metabolites\_id Metabolites ID

Reconstruction\_data Reconstruction data

Genome\_sequence Genome sequence

 ${\tt Genome\_sequence\_reference}\ \ Reference\ \ of\ genome\ sequence$ 

Reconstruction\_data\_reference Reference of reconstruction data

Data\_statistics\_metabolites Data statistics of metabolites

PubChem\_molecular\_formula Molecular formula in PubChem database

Brenda\_molecular\_formula Molecular formula in Brenda database

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Neutral\_molecular\_formula Neutral molecular formula

Charged\_molecular\_formula Charged molecular formula

Chebi\_molecular\_formula Chebi molecular formula

Conservation\_of\_mass\_and\_charge Conservation of mass and charge

Web\_GCM\_Gibbs Web GCM Gibbs

Gibbs\_free\_energy\_information Information of gibbs free energy

Gene\_id Gene ID

Protein id Protein ID

Cellular\_compartment Information of cellualr compartment

Subsystem\_information Subsystem information

Uniform\_identifier\_metabolites Uniform identifiers of metabollites

Spontaneous\_reactions Spontaneous reactions

Extracellular\_and\_periplasmic\_transport\_reactions Extracellular and periplasmic transport reactions

Exchange\_reactions Exchange reactions

Intracellular\_transport\_reactions ntracellular transport reactions

Amino\_acid\_weight Amino acid weight

Amino\_acid\_molecular\_weight Amino acid\_molecular weight

Dry\_weight Dry weight

Amino\_acid\_coefficient Amino acid coefficient

Nucleotide\_coefficient Nucleotide coefficient

Nucleotide\_weight Nucleotide weight

Nucleotide\_molecular\_weight Nucleotide molecular weight

Biomass\_reactions Biomass reactions

Demand\_reactions Demand reactions

Sink\_reactions Sink reactions

Scatter\_plot\_stoichiometric\_matrix Scatter plot of stoichiometric matrix

Objective\_reaction Objective reaction

Constraints Constraints of the model

Mass\_charge\_conservation\_assessment assessment of conservation of mass and charge

Terminal\_metabolites Terminal metabolites

Gap\_reactions Gap reactions

Missing\_exchange\_reactions Missing exchange reactions

Type\_III\_pathway Type III pathway

Network\_gaps Network gaps

Biomass\_metabolites Biomass metabolites

miniaml\_or\_maxiaml miniaml or maxiaml

text 21

Metabolic\_flux\_value Value of metabolic flux

Growth Prediction of the growth of model

Secretion\_product Product of secretion

Mutisecretion Multisecretion

Rich\_media Environment of rich media

Block\_reactions Block reactions

Knockout Knockout the single gene or reaction

Model\_predict\_correctly Predict the model correctly

Model\_growing\_too\_fast Assess whether the model grow too fast

Cofactors Cofactors

FBA Flux balance analysis

GPR Gene-protein reaction

Output Output the file

Iteration Iteration

Test Test

Assessment Assessment

Identifiers\_metabolites Identifiers of metabolites

Output\_file Output the file

Elemental\_balance Elemental balance

Biomass\_metabolites\_coefficient Coefficients of biomass and metabolites

text

A list of characters from the inputTxt.

# Description

A list of characters from the inputTxt.

## Usage

text

## **Format**

An object of class character of length 1.

22 toolsMatrix

	toolsMatrix	A data frame contains the details of the databases and tools of the metabolic reconstruction
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#### **Description**

A data frame contains the details of the databases and tools of the metabolic reconstruction

## Usage

toolsMatrix

#### **Format**

A data frame contains the 93 rows and 46 varibales.

the key words of the steps of metabolic reconstruction

MarkerWoFtdsesholdValue the values filter the steps after the mapping

Steps the steps of metabolic reonstruction

**Group** the classification of the steps

UniProtKnowledgeBase 1 means the output of UniProt Knowledgebase.

**NCBI\_Gene** 1 means the output of NCBI Gene.

**KEGG\_Genes** 1 means the output of KEGG Genes.

**KEGG\_Genome** 1 means the output of KEGG Genome.

NCBI\_Protein 1 means the output of NCBI Protein.

**KEGG\_Pathway** 1 means the output of KEGG Pathway.

KEGG\_Compound 1 means the output of KEGG Compound.

**KEGG\_Reaction** 1 means the output of KEGG Reaction.

**BioCyc** 1 means the output of BioCyc.

**MetaCycCompound** 1 means the output of MetaCyc Compound.

MetaCycReaction 1 means the output of MetaCyc Reaction.

**KEGGREST** 1 means the output of KEGGREST.

**COBRA** 1 means the output of COBRA.

**RAVEN** 1 means the output of RAVEN.

**CarveMe** 1 means the output of CarveMe.

AuReMe 1 means the output of AuReMe.

MetaDraft 1 means the output of MetaDraft.

**ModelSEED** 1 means the output of ModelSEED.

PathwayTools 1 means the output of Pathway Tools.

**Merlin** 1 means the output of Merlin.

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**AGORA** 1 means the output of AGORA.

**COBRApy** 1 means the output of COBRApy.

**BLAST** 1 means the output of BLAST.

ExperimentOrLiterature 1 means the output of Experiment or literature.

**BRENDA** 1 means the output of BRENDA.

**minval** 1 means the output of minval.

ChEBI 1 means the output of ChEBI.

**PubChem Compound** 1 means the output of PubChem-compound.

**ChEMBL\_Compound** 1 means the output of ChEMBL compound.

Rhea 1 means the output of Rhea.

**pKa\_DB** 1 means the output of pKa DB.

**rsbml** 1 means the output of rsbml.

PipelinePilot 1 means the output of Pipeline Pilot.

Sybil 1 means the output of Sybil.

**BUSCA** 1 means the output of BUSCA.

**PSORT** 1 means the output of PSORT.

**PASUB** 1 means the output of PASUB.

PubChem\_Substance 1 means the output of PubChem substance.

**STRING** 1 means the output of STRING.

ManualPlanning 1 means the output of Manual planning.

**CMR** 1 means the output of CMR.

**g2f** 1 means the output of g2f.

•	contains the labels and groups of the databases and tools reconstruction
---	--

#### Description

A data frame contains the labels and groups of the databases and tools of metabolic reconstruction

#### Usage

toolsTypes

#### **Format**

**label** the databases and tools of the metabolic reconstruction **group** the classfication of the labels

24 vizTools

|--|

# Description

Create a function can visualiaze the steps used in the metabolic reconstruction.

## Usage

```
vizProcess(text, stepsMatrix, stepTypes, contentTypes)
```

# Arguments

text	The characters processed with the collapse = '.'.
stepsMatrix	A data frame contains the marker words, threshold value, steps, group and other information about the metabolic reconstruction. The default file is in the data.
stepTypes	A data frame contains the labels and groups of the metabolic reconstructions steps. The default file is in the data.
contentTypes	A data frame contains the labels and groups of the metabolic reconstructions content The default file is in the data.

# Value

The pictures that visualize the steps of the metabolic reconstruction.

# **Examples**

```
exam <- vizProcess(text, stepsMatrix, stepTypes,contentTypes)</pre>
```

|--|--|

# Description

Input the txt and output the visualization of the steps, transformation and databases and tools.

# Usage

```
vizTools(text, toolsMatrix, stepTypes, toolsTypes)
```

vizTransformation 25

#### **Arguments**

text The characters processed with the collapse = '.'.

toolsMatrix A data frame contains the marker words, threshold value, steps, group and other

information about the metabolic reconstruction databases and tools. The default

file is in the data.

stepTypes A data frame contains the labels and groups of the metabolic reconstructions

steps. The default file is in the data.

toolsTypes A data frame contains the databases and the tools used in the metabolic recon-

struction.

#### Value

The pictures that visualize the databases and tools of the metabolic reconstruction.

## **Examples**

```
exam <- vizTools(text, toolsMatrix, stepTypes, toolsTypes)</pre>
```

vizTransformation vizTransformation

## **Description**

Input the txt and output the visualization of the steps, transformation and databases and tools.

#### Usage

```
vizTransformation(text, conversionMatrix, stepTypes, conversionTypes)
```

## **Arguments**

text The characters processed with the collapse = '.'.

conversionMatrix

A data frame contains the marker words, threshold value, steps, group and other transformation information about the metabolic reconstruction. The default file

is in the data.

stepTypes A data frame contains the labels and groups of the metabolic reconstructions

steps. The default file is in the data.

conversionTypes

A data frame contains the labels and groups of the metabolic reconstructions

transformation. The default file is in the data.

#### Value

The pictures that visualize the transformation of the metabolic reconstruction.

26 wordsMatrix

## **Examples**

exam <- vizTransformation(text, conversionMatrix, stepTypes, conversionTypes)</pre>

wordsMatrix

A data frame produced by the function get\_term\_matrix

# Description

A data frame produced by the function get\_term\_matrix

# Usage

wordsMatrix

#### **Format**

A data frame contains 4 variables

freq the frequency of the word occurs in a article.

prevalent the words type that often occurs in a article.

**longest** the longest type of a word.

**shortest** the shortest type of a word.

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