

Package: PerseusR (via r-universe)

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Title Perseus R Interop

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Description Enables the interoperability between the Perseus platform for omics data analysis (Tyanova et al. 2016) <doi:10.1038/nmeth.3901> and R. It provides the foundation for developing and running Perseus plugins implemented in R by providing all required input and output handling, including data and parameter parsing as described in Rudolph and Cox 2018 <doi:10.1101/447268>.

Depends R (>= 3.3.0)

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Imports utils, plyr, methods, XML, Biobase, stringr

Suggests testthat, roxygen2, knitr, rmarkdown

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annotCols	<i>Get annotation columns</i>
-----------	-------------------------------

Description

Get annotation columns

Usage

```
annotCols(mdata)
```

Arguments

mdata	matrixData
-------	------------

See Also

Other matrixData basic functions: [annotCols<-](#), [annotRows<-](#), [annotRows](#), [description<-](#), [description](#), [imputeData<-](#), [imputeData](#), [main<-](#), [main](#), [matrixData-class](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData<-](#), [qualityData](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a','b','c')),
  annotRows=data.frame(x=factor(c('1','1'))))
annotCols(mdata)
```

annotCols<-	<i>Set annotation columns</i>
-------------	-------------------------------

Description

Set annotation columns

Usage

```
annotCols(mdata) <- value
```

Arguments

mdata	matrixData
value	value

See Also

Other matrixData basic functions: [annotCols](#), [annotRows<-](#), [annotRows](#), [description<-](#), [description](#), [imputeData<-](#), [imputeData](#), [main<-](#), [main](#), [matrixData-class](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData<-](#), [qualityData](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a','b','c')),
  annotRows=data.frame(x=factor(c('1','1'))))
value <- data.frame(d=c('d', 'e', 'f'))
annotCols(mdata) <- value
```

annotRows	<i>Get annotation rows</i>
-----------	----------------------------

Description

Get annotation rows

Usage

```
annotRows(mdata)
```

Arguments

mdata matrixData

See Also

Other matrixData basic functions: [annotCols<-](#), [annotCols](#), [annotRows<-](#), [description<-](#), [description](#), [imputeData<-](#), [imputeData](#), [main<-](#), [main](#), [matrixData-class](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData<-](#), [qualityData](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a', 'b', 'c')),
  annotRows=data.frame(x=factor(c('1', '1'))))
annotRows(mdata)
```

annotRows<-	<i>Set annotation rows</i>
-------------	----------------------------

Description

Set annotation rows

Usage

```
annotRows(mdata) <- value
```

Arguments

mdata matrixData
value value

See Also

Other matrixData basic functions: [annotCols<-](#), [annotCols](#), [annotRows](#), [description<-](#), [description](#), [imputeData<-](#), [imputeData](#), [main<-](#), [main](#), [matrixData-class](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData<-](#), [qualityData](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a','b','c')),
  annotRows=data.frame(x=factor(c('1','1'))))
value <- data.frame(y=factor(c('2','2')))
annotRows(mdata) <- value
```

as.ExpressionSet.matrixData

Coerces a MatrixData into an ExpressionSet

Description

Coerces a MatrixData object into an ExpressionSet object

Usage

```
as.ExpressionSet.matrixData(mdata)
```

Arguments

mdata a [matrixData](#) object

Details

function to convert a [matrixData](#) [ExpressionSet](#)

Value

returns an [ExpressionSet](#) object

Examples

```
mD <- matrixData(
  main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(b=c('a','b','c')),
  annotRows=data.frame(x=factor(c('1','1'))))

eSet <- as(mD, "ExpressionSet")
print(eSet)
```

```
as.matrixData.ExpressionSet
```

Coerces an ExpressionSet into a MatrixData

Description

function to convert an [ExpressionSet](#) object into a [matrixData](#)

Usage

```
as.matrixData.ExpressionSet(ExpressionSet)
```

Arguments

ExpressionSet an [ExpressionSet](#) object

Value

returns a [matrixData](#) object

Examples

```
eSet <- eSet <- Biobase::ExpressionSet(matrix(1:10, ncol = 2))
mD <- as(eSet, "matrixData")
print(mD)
```

```
boolParamValue            Bool parameter value
```

Description

Extract the value chosen in an BoolParam

Usage

```
boolParamValue(parameters, name)
```

Arguments

parameters The parameters object (see [parseParameters](#))
name The name of the parameter

Value

The selected boolean

Examples

```
tmp <- tempfile(fileext = ".xml")
write('<BoolParam Name="test_bool">\n<Value>>false</Value>\n</BoolParam>', file=tmp)
parameters <- parseParameters(tmp)
boolParamValue(parameters, "test_bool")
```

create_annotRows	<i>Create annotation rows</i>
------------------	-------------------------------

Description

Create the annotation rows data.frame from the list of comment rows parsed from the input file and the main columns indicator

Usage

```
create_annotRows(commentRows, isMain)
```

Arguments

commentRows	list of comment rows
isMain	logical array indicating all main columns

See Also

used by [read.perseus](#)

description	<i>Get column description</i>
-------------	-------------------------------

Description

Get column description

Usage

```
description(mdata)
```

Arguments

mdata	matrixData
-------	------------

See Also

Other matrixData basic functions: [annotCols<-](#), [annotCols](#), [annotRows<-](#), [annotRows](#), [description<-](#), [imputeData<-](#), [imputeData](#), [main<-](#), [main](#), [matrixData-class](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData<-](#), [qualityData](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a','b','c')),
  annotRows=data.frame(x=factor(c('1','1'))),
  description=c('aaa', 'bbb', 'ccc'))
description(mdata)
```

description<- *Set column description*

Description

Set column description

Usage

```
description(mdata) <- value
```

Arguments

mdata	matrixData
value	value

See Also

Other matrixData basic functions: [annotCols<-](#), [annotCols](#), [annotRows<-](#), [annotRows](#), [description](#), [imputeData<-](#), [imputeData](#), [main<-](#), [main](#), [matrixData-class](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData<-](#), [qualityData](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a','b','c')),
  annotRows=data.frame(x=factor(c('1','1'))))
value <- c('aaa', 'bbb', 'ccc')
description(mdata) <- value
```

imputeData	<i>Get imputation of main data frame</i>
------------	--

Description

Get imputation of main data frame

Usage

```
imputeData(mdata)
```

Arguments

mdata	matrixData
-------	------------

See Also

Other matrixData basic functions: [annotCols<-](#), [annotCols](#), [annotRows<-](#), [annotRows](#), [description<-](#), [description](#), [imputeData<-](#), [main<-](#), [main](#), [matrixData-class](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData<-](#), [qualityData](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a', 'b', 'c')),
  annotRows=data.frame(x=factor(c('1', '1'))),
  imputeData=data.frame(impute=c('False', 'True', 'False')))
imputeData(mdata)
```

imputeData<-	<i>Set imputation of main data frame</i>
--------------	--

Description

Set imputation of main data frame

Usage

```
imputeData(mdata) <- value
```

Arguments

mdata	matrixData
value	value

See Also

Other matrixData basic functions: [annotCols<-](#), [annotCols](#), [annotRows<-](#), [annotRows](#), [description<-](#), [description](#), [imputeData](#), [main<-](#), [main](#), [matrixData-class](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData<-](#), [qualityData](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a','b','c')),
  annotRows=data.frame(x=factor(c('1','1'))),
  imputeData=data.frame(impute=c('False', 'True', 'False')))
value <- data.frame(impute=c('True', 'True', 'True'))
imputeData(mdata) <- value
```

infer_perseus_annotation_types

Infer Perseus type annotation row from DataFrame column classes

Description

Infer Perseus type annotation row from DataFrame column classes

Usage

```
infer_perseus_annotation_types(df, typeMap)
```

Arguments

df	The data.frame
typeMap	A list with elements 'Perseus' and 'R'. The ordering determines the mapping

Value

A vector with perseus type annotations

See Also

Based on [mapvalues](#)

```
initialize,matrixData-method
      matrixData initializer
```

Description

Initializes the annotCols data frame to have the same number of rows as the main data. This might not be the cleanest solution.

Usage

```
## S4 method for signature 'matrixData'
initialize(.Object, ...)
```

Arguments

.Object	Initialized object
...	Additional arguments

```
intParamValue      Int parameter value
```

Description

Extract the value chosen in an IntParam

Usage

```
intParamValue(parameters, name)
```

Arguments

parameters	The parameters object (see parseParameters)
name	The name of the parameter

Value

The selected number

Examples

```
tmp <- tempfile(fileext = ".xml")
write('<IntParam Name="test_int">\n<Value>2</Value>\n</IntParam>', file=tmp)
parameters <- parseParameters(tmp)
intParamValue(parameters, "test_int")
```

main	<i>Get main columns</i>
------	-------------------------

Description

Gets the main columns (main matrix) of a `matrixData` object as a `data.frame` object

Usage

```
main(mdata)
```

Arguments

mdata	matrixData
-------	------------

See Also

Other `matrixData` basic functions: [annotCols<-](#), [annotCols](#), [annotRows<-](#), [annotRows](#), [description<-](#), [description](#), [imputeData<-](#), [imputeData](#), [main<-](#), [matrixData-class](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData<-](#), [qualityData](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a', 'b', 'c')),
  annotRows=data.frame(x=factor(c('1', '1'))))
main(mdata)
```

main<-	<i>Set main columns</i>
--------	-------------------------

Description

Set main columns

Usage

```
main(mdata) <- value
```

Arguments

mdata	matrixData
value	value

See Also

Other matrixData basic functions: [annotCols<-](#), [annotCols](#), [annotRows<-](#), [annotRows](#), [description<-](#), [description](#), [imputeData<-](#), [imputeData](#), [main](#), [matrixData-class](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData<-](#), [qualityData](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a','b','c')),
  annotRows=data.frame(x=factor(c('1','1'))))
value<-data.frame(c=c(0,0,0), d=c(1,1,1))
main(mdata) <- value
```

matrixData

matrixData constructor

Description

matrixData constructor

Usage

```
matrixData(...)
```

Arguments

... main, annotCols, annotRows, description, imputeData, qualityData

See Also

Other matrixData basic functions: [annotCols<-](#), [annotCols](#), [annotRows<-](#), [annotRows](#), [description<-](#), [description](#), [imputeData<-](#), [imputeData](#), [main<-](#), [main](#), [matrixData-class](#), [names](#), [matrixData-method](#), [qualityData<-](#), [qualityData](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a','b','c')),
  annotRows=data.frame(x=factor(c('1','1'))),
  description=c('aaa', 'bbb', 'ccc'),
  imputeData=data.frame(impute=c('False', 'True', 'False')),
  qualityData=data.frame(quality=c('0', '1', '0')))
```

matrixData-class	<i>MatrixData</i>
------------------	-------------------

Description

MatrixData

Slots

main Main expression data.frame.
 annotCols Annotation Columns data.frame.
 annotRows Annotation Rows data.frame.
 description Column descriptions.
 imputeData Imputation data.frame.
 qualityData Quality values data.frame.

See Also

Other matrixData basic functions: [annotCols<-](#), [annotCols](#), [annotRows<-](#), [annotRows](#), [description<-](#), [description](#), [imputeData<-](#), [imputeData](#), [main<-](#), [main](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData<-](#), [qualityData](#)

MatrixDataCheck	<i>MatrixDataCheck: a function to check the validity of an object as a perseus data frame</i>
-----------------	---

Description

Check perseus compatibility of an object

Usage

```
MatrixDataCheck(object, ...)
```

Default S3 method:
 MatrixDataCheck(object = NULL, main, annotationRows,
 annotationCols, descriptions, imputeData, qualityData, all_colnames, ...)

S3 method for class 'matrixData'
 MatrixDataCheck(object, ...)

S3 method for class 'list'
 MatrixDataCheck(object, ...)

S3 method for class 'ExpressionSet'
 MatrixDataCheck(object, ...)

Arguments

object	object to check consistency with perseus data frames
...	additional arguments passed to the respective method
main	Main Data frame
annotationRows	Rows containing annotation information
annotationCols	Columns containing annotation information
descriptions	Descriptions of all the columns
imputeData	Is imputed or not
qualityData	quality number
all_colnames	The colnames to be used

Value

a logical indicating the validity of the object (or series of objects) as a perseus DF or the string of errors

NULL

NULL

NULL

Examples

```
require(PerseusR)

mat <- matrixData(
  main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a', 'b', 'c')),
  annotRows=data.frame(x=factor(c('1', '1'))))

MatrixDataCheck(mat)
```

multiChoiceParamValue *Multiple choice value*

Description

Extract the value selected in a MultiChoiceParam.

Usage

```
multiChoiceParamValue(parameters, name)
```

Arguments

parameters	The parameters object (see parseParameters)
name	The name of the parameter

Value

The string representing the value

Examples

```
write('<MultiChoiceParam Name="test_multi">\n<Value>\n<Item>1</Item>\n<Item>2</Item>\n</Value>\n
<Values>\n<Item>A</Item>\n<Item>B</Item>\n</Values>\n</MultiChoiceParam>', file='tmp.xml')
parameters <- parseParameters("tmp.xml")
multiChoiceParamValue(parameters, "test_multi")
```

names,matrixData-method

Get names

Description

Get the column names of main and annotation columns.

Usage

```
## S4 method for signature 'matrixData'
names(x)
```

Arguments

x	matrixData
---	------------

See Also

Other matrixData basic functions: [annotCols<-](#), [annotCols](#), [annotRows<-](#), [annotRows](#), [description<-](#), [description](#), [imputeData<-](#), [imputeData](#), [main<-](#), [main](#), [matrixData-class](#), [matrixData](#), [qualityData<-](#), [qualityData](#)

names.matrixData	<i>Column names of main and annotation columns</i>
------------------	--

Description

Column names of main and annotation columns

Usage

```
## S3 method for class 'matrixData'  
names(x)
```

Arguments

x	matrixData
---	------------

parseParameters	<i>Parse parameters</i>
-----------------	-------------------------

Description

Parse parameters from the parameters xml file.

Usage

```
parseParameters(paramFile)
```

Arguments

paramFile	Parameters xml file
-----------	---------------------

Examples

```
tmp <- tempfile(fileext = ".xml")  
write('<IntParam Name="test_int">\n<Value>2</Value>\n</IntParam>', file=tmp)  
parameters <- parseParameters(tmp)
```

qualityData	<i>Get quality values of main data frame</i>
-------------	--

Description

Get quality values of main data frame

Usage

```
qualityData(mdata)
```

Arguments

mdata	matrixData
-------	------------

See Also

Other matrixData basic functions: [annotCols<-](#), [annotCols](#), [annotRows<-](#), [annotRows](#), [description<-](#), [description](#), [imputeData<-](#), [imputeData](#), [main<-](#), [main](#), [matrixData-class](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData<-](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a','b','c')),
  annotRows=data.frame(x=factor(c('1','1'))),
  qualityData=data.frame(quality=c('1', '1', '1')))
qualityData(mdata)
```

qualityData<-	<i>Set quality values of main data frame</i>
---------------	--

Description

Set quality values of main data frame

Usage

```
qualityData(mdata) <- value
```

Arguments

mdata	matrixData
value	value

See Also

Other matrixData basic functions: [annotCols<-](#), [annotCols](#), [annotRows<-](#), [annotRows](#), [description<-](#), [description](#), [imputeData<-](#), [imputeData](#), [main<-](#), [main](#), [matrixData-class](#), [matrixData](#), [names](#), [matrixData-method](#), [qualityData](#)

Examples

```
mdata <- matrixData(main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(c=c('a','b','c')),
  annotRows=data.frame(x=factor(c('1','1'))),
  qualityData=data.frame(quality=c('1', '1', '1')))
value <- data.frame(quality=c('0', '0', '0'))
qualityData(mdata) <- value
```

read.perseus.default *Read Perseus matrix files*

Description

Read the custom Perseus matrix file format *.txt into R.

Usage

```
read.perseus.default(con, check = TRUE, additionalMatrices = FALSE)
```

```
read.perseus.as.list(con, check = TRUE)
```

```
read.perseus.as.matrixData(con, check = TRUE,
  additionalMatrices = FALSE)
```

```
read.perseus.as.ExpressionSet(con, check = TRUE)
```

```
read.perseus(con, check = TRUE, additionalMatrices = FALSE)
```

Arguments

con	A connection object or the path to input file
check	Logical indicating whether to check for the validity of the exported object (slightly slower)
additionalMatrices	Logical indication whether to write out quality and imputation matrices in perseus format

Value

Defaults to a [matrixData](#) object.

Functions

- `read.perseus.default`: Returns a list used internally to generate all other outputs
- `read.perseus.as.list`: Returns explicitly as a list
- `read.perseus.as.matrixData`: Returns explicitly as a specialized matrix data object
- `read.perseus.as.ExpressionSet`: Returns a bioconductor expression set object

Note

Limitations to column names in R still apply. Column names valid in Perseus, such as 'Column 1' will be changed to 'Column.1'

If the provided connection `con` is a character string, it will assumed to be a file path. A [connection](#) which is not seekable (see [isSeekable](#)) will be written to a temporary file. Any connection will be closed when `read.perseus` exits. `read.perseus.as.list`, `read.perseus.as.matrixData` and `read.perseus.as.ExpressionSet` are also available depending on the class desired as an output

See Also

[write.perseus](#)
[matrixData](#)

Examples

```
tmp <- tempfile(fileext = ".txt")
write('Column_1\tColumn_2\tColumn_3
#{Description}\t\t
#{Type}E\tE\tE
-1.860574\t-0.3910594\t0.2870352
NaN\t-0.4742951\t0.849998', file=tmp)
mdata <- read.perseus(tmp)
```

singleChoiceParamInd *Single choice index*

Description

Extract the index chosen in an BoolParam

Usage

```
singleChoiceParamInd(parameters, name)
```

Arguments

<code>parameters</code>	The parameters object (see parseParameters)
<code>name</code>	The name of the parameter

Value

The selected index

Examples

```
tmp <- tempfile(fileext = ".xml")
write('<SingleChoiceParam Name="test_single">\n<Value>1</Value>\n
<Values>\n<Item>A</Item>\n<Item>B</Item>\n</Values>\n</SingleChoiceParam>', file=tmp)
parameters <- parseParameters(tmp)
singleChoiceParamInd(parameters, "test_single")
```

singleChoiceParamValue

Single choice value

Description

Extract the value selected in a SingleChoiceParam.

Usage

```
singleChoiceParamValue(parameters, name)
```

Arguments

parameters	The parameters object (see parseParameters)
name	The name of the parameter

Value

The string representing the value

Examples

```
tmp <- tempfile(fileext = ".xml")
write('<SingleChoiceParam Name="test_single">\n<Value>1</Value>\n
<Values>\n<Item>A</Item>\n<Item>B</Item>\n</Values>\n</SingleChoiceParam>', file=tmp)
parameters <- parseParameters(tmp)
singleChoiceParamValue(parameters, "test_single")
```

stringParamValue	<i>String parameter value</i>
------------------	-------------------------------

Description

Extract the value chosen in an StringParam

Usage

```
stringParamValue(parameters, name)
```

Arguments

parameters	The parameters object (see parseParameters)
name	The name of the parameter

Value

The selected string

Examples

```
write('<StringParam Name="test_string">\n<Value>"test"</Value>\n</StringParam>', file='tmp.xml')
parameters <- parseParameters("tmp.xml")
stringParamValue(parameters, "test_string")
```

write.perseus	<i>write.perseus: function to generate a perseus-readable text document</i>
---------------	---

Description

Write data to a perseus text file or connection
 Write Data to file in the custom Perseus matrix file format.

Usage

```
write.perseus(object = NULL, con = NULL, ...)

## Default S3 method:
write.perseus(object = NULL, con = NULL, main,
  annotCols = NULL, annotRows = NULL, descr = NULL,
  imputeData = NULL, qualityData = NULL, ...)

## S3 method for class 'matrixData'
write.perseus(object, con, ...)
```

```
## S3 method for class 'list'
write.perseus(object, con, ...)

## S3 method for class 'data.frame'
write.perseus(object, con, annotCols = NULL, ...)

## S3 method for class 'matrix'
write.perseus(object, con, annotCols = NULL, ...)

## S3 method for class 'ExpressionSet'
write.perseus(object, con, ...)
```

Arguments

object	an expressionSet, matrixData, list or table-like object.
con	A connection object or the path to output file
...	additional arguments passed to other functions
main	a data frame containing
annotCols	a df containing columns containing metadata (about the rows)
annotRows	a df containing columns containing metadata (about the columns)
descr	a character vector that describes the columns in main and in annotCols (in that order)
imputedData	a df containing imputations – True or False of main data frame
qualityData	a df containing quality values of main data frame

Value

writes to disk a perseus-interpretable text representation of an R object

NULL

NULL

NULL

NULL

NULL

See Also

[read.perseus matrixData](#)

Examples

```
df <- matrixData(
  main=data.frame(a=1:3, b=6:8),
  annotCols=data.frame(b=c('a', 'b', 'c')),
  annotRows=data.frame(x=factor(c('1', '1'))),
  description=c('a', 'a', 'b'))
```

```
con <- textConnection('df1', 'w')  
write.perseus(df, con)  
close(con)
```


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