

Package: do (via r-universe)

August 2, 2024

Type Package

Title Data Operator

Version 2.0.0.0

Description Flexibly convert data between long and wide format using just two functions: `reshape_toLong()` and `reshape_toWide()`.

Author Jing Zhang, Zhi Jin

Maintainer Jing Zhang<zj391120@163.com>

License GPL-3

Encoding UTF-8

Imports data.table, plyr, tmcn, methods, tidyclear, reshape2, tidyr, rvest, xml2, crayon, httr, usethis, desc, utils, rstudioapi, stringr, openxlsx

RoxygenNote 7.1.1

URL <https://github.com/yikeshu0611/do>

BugReports <https://github.com/yikeshu0611/do/issues>

Suggests knitr, rmarkdown

VignetteBuilder knitr

Depends R (>= 4.1)

Repository <https://yikeshu0611.r-universe.dev>

RemoteUrl <https://github.com/yikeshu0611/do>

RemoteRef HEAD

RemoteSha 7d91dc33676d7e1d84a30650c23691ffdea49f0e

Contents

add_biocViews	4
all_children	4
Apriori.Basket	5
as.data.frame	6

as.data.frame.rules	6
as.transactions	7
attr_href	7
c.xml_nodeset	8
cat_n	8
character.nms	9
chinese_utf8	9
CIGI	10
cnOS	10
columntrans	11
col_split	12
common	12
compare	13
complete.data	14
CTD2	14
decrease	15
delete_left	16
delete_up	16
deparse0	17
desc2df	17
detach2	18
dump.it	18
dup.connect	19
duplicate	19
duplicated_all	20
duplicated_last	21
equal_length	21
exec	22
expand	23
factor.nms	23
file.dir	24
file.name	24
fmt	25
formal_dir	26
get_names	26
give_names	27
Grepl	28
has_children	29
increase	29
inner_Add_Symbol	30
insertglue	30
install_Rversion	31
is.dir	31
is.linux	32
is.mac	32
is.windows	33
join	33
keep	34

knife	35
last	35
last_column	36
last_row	36
left	37
left_equal	37
legal	38
load_extdata	39
mid	39
mirror.current	40
mirror.set	40
mirror.speed	41
model.data	41
NA.col.prob	42
NA.col.sums	42
NA.row.prob	43
NA.row.sums	43
NA.whole.prob	44
NA.whole.sums	45
names_n	45
Nchar	46
numeric.nms	46
package_all	47
paste0_columns	47
pipe	48
rd2df	48
read_R	48
Replace	49
Replace0	50
Replace_ex	50
replicate	51
reshape_toLong	52
reshape_toWide	52
reshape_towide	53
reverse	54
right	55
right_equal	56
rm_all	56
rm_nchar	57
row.freq	57
select	58
seq_range	59
show_function	60
split_expand	60
table_NA	61
take_out	61
Target	62
Trim	63

unique_no.NA	63
unlibray	64
upper.dir	64
write_xlsx	65
%==%	66
%+%	66
%s=%	67

Index 68

add_biocViews *Add biocViews Field to Description File*

Description

Add biocViews Field to Description File

Usage

```
add_biocViews(value = "", overwrite = TRUE)
```

Arguments

value	package names
overwrite	logical, TRUE is defaulted

all_children *Extract all children nodes*

Description

Extract all children nodes

Usage

```
all_children(x, res = "do not change", i = 1)
```

Arguments

x	one or more documents, nodes, or node sets.
res	omit. do not make any change.
i	must be 1

Value

nodeset

Examples

```
txt='<div class="activityBody postBody thing">
  <p>
    <a href="/forum/conversation" class="mqPostRef">(22)</a>
    where?
  </p>
  <p>
    stays
    <b>disappears</b>
    <a>disappears</a>
    <span>disappears</span>
    stays
  </p>
</div>'
library(xml2)
html=read_html(txt)

all_children(html)
```

Apriori.Basket

Convert vector to sparse matrix

Description

Convert vector or dataframe to sparse matrix.

Usage

```
Apriori.Basket(x, sep = ";", dup.delete = FALSE)
```

Arguments

x	a vector
sep	one separator
dup.delete	whether to delete duplicated values in the same row, default is FALSE

Value

a sparse matrix

Examples

```
# convert a vector to sparse matrix
g=c('a,b,a,,','a,b,c,d','d,c,f,g,h')
Apriori.Basket(x=g,sep = ',')

# convert a dataframe to sparse matrix
library(data.table)
```

```
df=fread(text = '
t1 t2 t3
a NA d
g a j')
Apriori.Basket(x=df,sep = ',')
```

as.data.frame *Transform to dataframe rules object or calibrate object*

Description

Transform to dataframe rules object or calibrate object

Usage

```
## S3 method for class 'rules'
as.data.frame(x, row.names = NULL, optional = FALSE, ...)

## S3 method for class 'calibrate'
as.data.frame(x, row.names = NULL, optional = FALSE, ...)
```

Arguments

x	data with rules class for package 'arules'
row.names	ignore
optional	ignore
...	ignore

Value

a dataframe

as.data.frame.rules *Transform rules to dataframe*

Description

Transform rules to dataframe

Usage

```
## S3 method for class 'rules'
as.data.frame(x, row.names = NULL, optional = FALSE, ...)
```

Arguments

x	data with rules class for package 'arules'
row.names	ignore
optional	ignore
...	ignore

Value

a dataframe

as.transactions	<i>Transform to transactions</i>
-----------------	----------------------------------

Description

Transform to transactions

Usage

```
as.transactions(x)
```

Arguments

x	dataframe or matrix
---	---------------------

Value

a transaction data

attr_href	<i>Get hypertext reference attributes</i>
-----------	-------------------------------------------

Description

Get hypertext reference attributes

Usage

```
attr_href(x)
```

Arguments

x	A document (from read_html()), node set (from html_elements()), node (from html_element()), or session (from session()).
---	--------------------------------------------------------------------------------------------------------------------------

Value

hypertext reference attributes

c.xml_nodeset	<i>Comine xml_nodeset</i>
---------------	---------------------------

Description

Comine xml_nodeset

Usage

```
## S3 method for class 'xml_nodeset'  
c(...)
```

Arguments

... one or more xml_nodeset

Value

xml_nodeset

cat_n	<i>print vector by lines</i>
-------	------------------------------

Description

print vector by lines

Usage

```
cat_n(x, n = 3, ind = 0)
```

Arguments

x one vector
n number of element in each line, default is 3
ind indentation, default is 0

Value

print vector by lines

Examples

```
cat_n(1:10)  
cat_n(1:10, ind=3)
```

character.nms	<i>Return character names in matrix or dataframe</i>
---------------	------------------------------------------------------

Description

Return character names in matrix or dataframe

Usage

```
character.nms(df)
```

Arguments

df dataframe or matrix

Value

character names vectors

chinese_utf8	<i>UTF8 Code for Chinese</i>
--------------	------------------------------

Description

UTF8 Code for Chinese

Usage

```
chinese_utf8(x)
```

Arguments

x chinese characters

Value

an expression with UTF8 code.

CIGI *Download data from CIGI database*

Description

Download data from CIGI database

Usage

CIGI_download(url)

CIGI_download_file(url)

CIGI_download_url(url)

Arguments

url url link of data

Value

downloaded data

Examples

```
CIGI_download('https://cigi-data.nci.nih.gov/Public/')
```

```
# download file by url
```

```
CIGI_download_file(url='https://cgci-data.nci.nih.gov/Public/BLGSP/biospecimen/BCCA/CGCI-BLGSP_BiospecimenSupp')
```

```
# get all downloaded urls
```

```
CIGI_download_url(url='https://cigi-data.nci.nih.gov/Public/')
```

cnOS *Chinese operating system Whether the computer is Chinese operating system*

Description

Chinese operating system Whether the computer is Chinese operating system

Usage

cnOS()

Value

logical

Examples

```
cnOS()
```

columntrans

Change data type

Description

Change data type

Usage

```
factor.it(x) <- value  
factor.it(x, value)  
numeric.it(x, value)  
numeric.it(x) <- value
```

Arguments

x	dataframe
value	column names

Value

factor or numeric columns in a dataframe

Examples

```
str(mtcars)  
factor.it(mtcars,c("cyl", "vs", "am", "gear"))  
factor.it(mtcars)=c("cyl", "vs", "am", "gear")  
str(mtcars)  
  
numeric.it(mtcars,c("cyl", "vs", "am", "gear"))  
numeric.it(mtcars)=c("cyl", "vs", "am", "gear")  
str(mtcars)
```

col_split	<i>Split A Vector into Columns</i>
-----------	------------------------------------

Description

Split A Vector into Columns

Usage

```
col_split(x, split, reg_expr, colnames, cat = TRUE)
```

Arguments

x	a vector
split	one or more characters. Split exactly
reg_expr	character. Split by regular expressions
colnames	optional. Column names for outcome
cat	logical, whether to show message

Value

A dataframe with several columns.

Examples

```
x=c('1a2', '3a4', '4a4')
col_split(x,split='a')
col_split(x = x,reg_expr = '[a-z]')

#two splits
df=data.frame(result=c('A, B-C',
                      'A, C-D',
                      'E, F-G'))
col_split(x = df[,1],split = c(',', '-'))
```

common	<i>Find Common Objects from Vectors</i>
--------	-----------------------------------------

Description

Find Common Objects from Vectors

Usage

```
common(...)
```

Arguments

... must be several vectors

Value

common objects

Examples

```
x1=c('a','e','d')
x2=c('a','c','e')
x3=c('a','e','j','d')
common(x1,x2,x3)
```

compare

Compare two vectors

Description

Compare two vectors

Usage

equal(a, b)

over(a, b)

lower(a, b)

Arguments

a one vector

b the other vector

Value

the compared object

Examples

```
equal(letters,c('a','b'))
over(1:10,5)
over(1:10,5)
```

<code>complete.data</code>	<i>Complete data</i>
----------------------------	----------------------

Description

Removing rows with NA in dataframe or matrix. Removing NA atomic.

Usage

```
complete.data(x)
```

Arguments

x dataframe or matrix or atomic

Value

complete data

Examples

```
x=c(1,NA,2)
complete.data(x)

x=data.frame(a=c(1,NA))
complete.data(x)
```

CTD2	<i>Download data from CTD2 database</i>
------	-----------------------------------------

Description

Download data from CTD2 database

Usage

```
CTD2_download_dir(url)

CTD2_download_file(url)

CTD2_download_url(url)
```

Arguments

url url link of data

Value

downloaded data

Examples

```
CTD2_download_dir('https://ctd2-data.nci.nih.gov/Public/')

# download files by file url
CTD2_download_file(url='https://ctd2-data.nci.nih.gov/Public/Broad/CTRPv1.0_2013_pub_Cell_154_1151/CTRPv1.0_2013_pub_Cell_154_1151/')

# get file url
CTD2_download_url(url='https://ctd2-data.nci.nih.gov/Public/Broad/CTRPv1.0_2013_pub_Cell_154_1151/')
```

decrease	<i>decrease character</i>
----------	---------------------------

Description

decrease character

Usage

```
decrease(chr)
```

Arguments

chr one character vector

Value

decreased vector

Examples

```
set.seed(2020)
x=rnorm(20)
decrease(x)
```

delete_left	<i>Delete and Move Left the rest Values</i>
-------------	---------------------------------------------

Description

Delete and Move Left the rest Values

Usage

```
delete_left(x, delete)
```

Arguments

x	dataframe or matrix
delete	one delete object

Value

dataframe or matrix

Examples

```
a=c(1,NA,7,NA)
b=c(NA,2,2,7)
d=c(1,NA,40,7)
df=data.frame(a,b,d)
delete_left(x=df,NA)
```

delete_up	<i>Delete and Move Up the Rest Values</i>
-----------	-------------------------------------------

Description

Delete and Move Up the Rest Values

Usage

```
delete_up(x, delete)
```

Arguments

x	dataframe or matrix
delete	one delete object

Value

dataframe or matrix

Examples

```
a=c(1,NA,7,NA)
b=c(NA,2,2,7)
d=c(1,NA,40,7)
df=data.frame(a,b,d)

delete_up(x = df,delete = NA)
```

deparse0	<i>substitutue, deparse and paste</i>
----------	---------------------------------------

Description

substitutue, deparse and paste

Usage

```
deparse0(x)
```

Arguments

x one object

Value

character

Examples

```
deparse0(j)
```

desc2df	<i>Convert package description file to dataframe</i>
---------	------------------------------------------------------

Description

Convert package description file to dataframe

Usage

```
desc2df(desc)
```

Arguments

desc description file path

Value

One dataframe with column names of field

detach2	<i>Detach package</i>
---------	-----------------------

Description

Detach package

Usage

```
detach2(x)
```

Arguments

x one package name, if missing, detach all packages

Value

detach one package

dump.it	<i>Create dump matrix for a vector</i>
---------	----------------------------------------

Description

Create dump matrix for a vector

Usage

```
dump.it(..., include.name = TRUE)
```

Arguments

... one vector
include.name logical, default is TRUE, wether to include name of variable

Value

a dump matix contains 0 and 1

Examples

```
x=c('a','b','c','a','a')
dump.it(x)
dump.it(mtcars$am)
dump.it(mtcars[, 'am'])
```

dup.connect	<i>Connect Duplicated Values</i>
-------------	----------------------------------

Description

Connect Duplicated Values

Usage

```
dup.connect(data, id, dup.var)
```

Arguments

data	dataframe or matrix
id	id column names or indexes
dup.var	duplicated column names or indexes

Value

dataframe contains id and duplicated values

Examples

```
dup.connect(data = mtcars, id = 'am', dup.var = 'cyl')
dup.connect(data = mtcars,
            id = c('am', 'gear'),
            dup.var = c('cyl', 'qsec'))
```

duplicate	<i>find all duplicated ones</i>
-----------	---------------------------------

Description

find all duplicated ones

Usage

```
duplicate(x, column_between = FALSE, row_between = FALSE,
          row_in = FALSE, column_in = FALSE, every = FALSE)
```

Arguments

x can be number, strings, vectors, dataframe or matrix
 column_between TRUE is to detect the duplicated rows.
 row_between TRUE is to detect the duplicated columns.
 row_in TRUE is to detect the duplicated values in each row.
 column_in TRUE is to detect the duplicated values in each column.
 every TRUE is to detect the duplicated values in dataframe or matrix.

Value

TRUE as duplicated, FALSE as not

Examples

```
x <- c('a', 'b', 'a', 1, 1, 2)
duplicate(x)
a = data.frame(k=c(1, 1, 2, 1, 4),
l=c(1, 1, 'a', 5, 6))
duplicate(a, row_in = TRUE)
duplicate(a, row_between = TRUE)
duplicate(a, column_between = TRUE)
duplicate(a, column_in = TRUE)
```

duplicated_all

Determine All Duplicate Elements

Description

Determine All Duplicate Elements

Usage

```
duplicated_all(x)
```

Arguments

x character

Value

logical value

Examples

```
x=c(1, 3, 2, 1, 2)
duplicated(x)
duplicated_all(x)
```

duplicated_last	<i>Determine Duplicate Elements in the Last Position</i>
-----------------	----------------------------------------------------------

Description

Determine Duplicate Elements in the Last Position

Usage

```
duplicated_last(x)
```

Arguments

x	character
---	-----------

Value

logical value

Examples

```
x=c(1,3,2,1,2)
duplicated(x)
duplicated_last(x)
```

equal_length	<i>Equal Length</i>
--------------	---------------------

Description

Equal Length

Usage

```
equal_length(x, suffix = " ", nchar, colname = FALSE, rowname = FALSE)
```

Arguments

x	can be number, strings, vectors, dataframe or matrix.
suffix	suffix
nchar	maximum length
colname	a logical value, default is FALSE
rowname	a logical value, default is FALSE

Value

equal length results

Examples

```
a=c(123,1,24,5,1.22554)
equal_length(a,0)

df = data.frame(
  a=c(12,1,1.23),
  b=c('a', 'abcd', 'd')
)
equal_length(x = df,suffix = 'x')

equal_length(x = df,suffix = 0,nchar =5)
```

exec	<i>execute string command This command just execute in the parent frame.</i>
------	------------------------------------------------------------------------------

Description

execute string command This command just execute in the parent frame.

Usage

```
exec(string, envir = parent.frame())
```

Arguments

string	one string
envir	the environment in which sting is to be evaluated.

Value

execute string command

Examples

```
a=2
exec('a = 1')
a
```

expand	<i>Expand Data by Weight</i>
--------	------------------------------

Description

Expand Data by Weight

Usage

```
expand(x, weight)
```

Arguments

x	dataframe or matrix
weight	weight column names or index

Value

expanded data

Examples

```
df=data.frame(v=c(1,2,3),
              x=c(7,8,9),
              n=c(2,3,4))
expand(x = df,weight = 3)
expand(x = df,weight = 'n')
```

factor.nms	<i>Return factor names in matrix or dataframe</i>
------------	---------------------------------------------------

Description

Return factor names in matrix or dataframe

Usage

```
factor.nms(df)
```

Arguments

df	dataframe or matrix
----	---------------------

Value

factor names vectors

file.dir	<i>up level directory</i>
----------	---------------------------

Description

up level directory

Usage

```
file.dir(path, end.slash = TRUE, extension = TRUE)
```

Arguments

path	path of file
end.slash	logical. Whether to end with slash
extension	logical. whether file name include extension

Value

upper directory

file.name	<i>Extract file name</i>
-----------	--------------------------

Description

Extract file name

Usage

```
file.name(..., extension = TRUE)
```

Arguments

...	one or more file path
extension	whether include extension, default is TRUE

Value

file names

Examples

```
file.name('f:/dir/1.txt')
file.name('f:/dir/1.txt', 'f:/dir/1.txt')
file.name('f:/dir/1.txt', 'f:/dir/1.txt', 'f:/dir/')

```

 fmt *Formatting Replacement*

Description

Formatting Replacement

Usage

```
fmt(x, ...)
```

Arguments

x	format with slash number and one space, which is like "/1 ". Number means replacement order.
...	values to be passed into x

Value

replaced string

Examples

```
'whwdzg, ykybnfg'
fmt('/ hwdzg, ykybnfg',
    'w')
fmt('/ h/ dzg, ykybnfg',
    'w')
fmt('/1 h/ dzg, ykybnfg',
    'w')
fmt('/1 h/ dzg, ykybnfg',
    'w', '-w-')

fmt('/ h/1 dzg, ykybnfg',
    'w', '-w-')

fmt('/1 h/0 dzg, ykybnfg',
    'w', '-w-')

'|w|' |>
  fmt(x = '/ h/ dzg, ykybnfg')

'|w|' |>
  fmt(x = '/ h/ dzg, ykybnfg',
      '-w-')

'|w|' |>
  fmt(x = '/ h/1 dzg, ykybnfg',
      '-w-')
```

formal_dir	<i>formal directory</i>
------------	-------------------------

Description

formal directory

Usage

```
formal_dir(dir, end.slash = FALSE)
```

Arguments

dir	one directory
end.slash	logical

Value

formed directory

get_names	<i>Get Names of Object</i>
-----------	----------------------------

Description

Return the names of input. For example: if you input a, you will get 'a'.

Usage

```
get_names(...)
```

Arguments

...	any type of data object
-----	-------------------------

Value

names of object

Examples

```
a=c(1,2,3)
get_names(a,mtcars)
```

give_names	<i>change vector, dataframe or matrix names</i>
------------	-------------------------------------------------

Description

change vector, dataframe or matrix names

Usage

```
give_names(data, ...)  
  
## S3 method for class 'character'  
give_names(data, ...)  
  
## S3 method for class 'numeric'  
give_names(data, ...)  
  
## S3 method for class 'logical'  
give_names(data, ...)  
  
## S3 method for class 'list'  
give_names(data, ...)  
  
## S3 method for class 'data.frame'  
give_names(data, ..., row = FALSE)  
  
## S3 method for class 'matrix'  
give_names(data, ..., row = FALSE)
```

Arguments

data	one vector, list, dataframe or matrix
...	one or more names
row	logical, whether the names is row names. Default is FALSE

Value

names changed data

Grep1

Judge for Included Character

Description

Judge for Included Character

Usage

```
Grep1(pattern, x)
```

Arguments

pattern	one or more vectors
x	one or more vectors

Details

,

Value

a matrix with logical words

Examples

```
a=c('abcd', 'agj', 'abcu')

# Grep1 for one vector
pat1='b'
Grep1(pat1,a)

# Grep1 for two vectors
pat2=c('c', 'd')
Grep1(pat2,a)

# use %or% in pattern
pat3=c('a%or%c', 'd')
Grep1(pat3,a)

# use %and% in pattern
pat4=c('a%and%c', 'd')
Grep1(pat4,a)
```

has_children	<i>Wether children nodes exist</i>
--------------	------------------------------------

Description

Wether children nodes exist

Usage

```
has_children(...)
```

Arguments

... one or more documents, nodes, or node sets.

Value

logical value

Examples

```
txt='<div class="activityBody postBody thing">
  <p>
    <a href="/forum/conversation" rel="post" >(22)</a>
    where?
  </p>
  <p>
    stays
    <b>disappears</b>
    <a>disappears</a>D
    <span>disappears</span>
    stays
  </p>
</div>'
library(xml2)
html=read_html(txt)
has_children(html)
```

increase	<i>increase character</i>
----------	---------------------------

Description

increase character

Usage

```
increase(chr)
```

Arguments

chr one vector

Value

increased vector

Examples

```
set.seed(2020)
x=rnorm(20)
increase(x)
```

inner_Add_Symbol *Concatenate Strings*

Description

Concatenate vectors by adding a symbol.

Usage

```
inner_Add_Symbol(x, symbol = "+")
```

Arguments

x vectors
symbol default is '+'

Value

a concatenated string

Examples

```
inner_Add_Symbol(c('a','b'))
inner_Add_Symbol(c('a','b'),"$")
inner_Add_Symbol(c('a','b'),"")
```

insertglue *glue*

Description

glue

Usage

```
insertglue()
```

install_Rversion	<i>Install contributed packages by R version</i>
------------------	--------------------------------------------------

Description

Install contributed packages by R version

Usage

```
install_Rversion(..., platform, Rversion = NULL, lib = ".")
```

Arguments

...	one or more package
platform	windows or mac
Rversion	version of R
lib	path

Value

contributed packages

is.dir	<i>Whether file path is directory</i>
--------	---------------------------------------

Description

Whether file path is directory

Usage

```
is.dir(...)
```

Arguments

...	one or more file path
-----	-----------------------

Value

logical

`is.linux`*operation system*

Description

operation system

Usage`is.linux()`**Value**

logical

Examples`is.linux()`

`is.mac`*operation system*

Description

operation system

Usage`is.mac()`**Value**

logical

Examples`is.mac()`

is.windows	<i>operation system</i>
------------	-------------------------

Description

operation system

Usage

```
is.windows()
```

Value

logical

Examples

```
is.windows()
```

join	<i>Join two dataframes together</i>
------	-------------------------------------

Description

Join two dataframes by the same id column.

Usage

```
join_inner(x, y, by = NULL)
```

```
join_full(x, y, by = NULL)
```

```
join_left(x, y, by = NULL)
```

```
join_right(x, y, by = NULL)
```

```
join_out(x, y, by = NULL)
```

Arguments

x	one dataframe
y	the other dataframe
by	the id name in x and y dataframe

Details

`join_inner()`, `join_full()`, `join_left()`, `join_right()` and `join_out()` are five functions to join two dataframes together. They are based on package 'data.table', so they are more efficient and fast.

Value

one joined dataframe.

Examples

```
df1=data.frame(x=rep(c('b','a','c'),each=3),
               y=c(1,3,6),
               v=1:9)

df2=data.frame(x=c('c','b','e'),
               v=8:6,
               foo=c(4,2,1))
join_inner(df1,df2,'x')
join_full(df1,df2,'x')
join_left(df1,df2,'x')
join_right(df1,df2,'x')
join_out(df1,df2,'x')
```

keep

Keep objects

Description

Keep objects

Usage

```
keep(..., envir = .GlobalEnv)
```

Arguments

...	one or more objects
envir	environment, default is global

Examples

```
a <- 1
b <- 2
d <- 4
keep(a)
```

knife	<i>Knife characters</i>
-------	-------------------------

Description

Knife characters

Usage

```
knife_left(x, n)
```

```
knife_right(x, n)
```

Arguments

x	one character
---	---------------

n	number
---	--------

Examples

```
knife_left(123,2)  
knife_right(123,2)
```

last	<i>Select character from last</i>
------	-----------------------------------

Description

Select character from last

Usage

```
last(x, n)
```

Arguments

x	vector
---	--------

n	If missing, the last element will be used.
---	--------------------------------------------

Value

last element

Examples

```
letters |> last()  
letters |> last(1:2)
```

last_column	<i>Select dataframe column from last</i>
-------------	------------------------------------------

Description

Select dataframe column from last

Usage

```
last_column(x, n)
```

Arguments

x	dataframe
n	If missing, the last element will be used.

Value

last column

Examples

```
mtcars |> last_column()  
mtcars |> last_column(1:2)
```

last_row	<i>Select dataframe row from last</i>
----------	---------------------------------------

Description

Select dataframe row from last

Usage

```
last_row(x, n)
```

Arguments

x	dataframe
n	If missing, the last element will be used.

Value

last row

Examples

```
mtcars |> last_row()
mtcars |> last_row(1:2)
```

left	<i>Truncate Characters from the Left</i>
------	------------------------------------------

Description

Truncate Characters from the Left

Usage

```
left(x, n)
```

Arguments

x	can be number, strings, vectors, dataframe or matrix.
n	length

Value

substring

Examples

```
left("abcd", 3)
left(c("abc", "gjh"), 2)
df = data.frame(
  a = c(123, 234, 456),
  b = c("abc", "bcd", "hjk")
)
left(df, 2)
```

left_equal	<i>Compare two characters from left Much useful for arguments input. Case is ignored.</i>
------------	-------------------------------------------------------------------------------------------

Description

Compare two characters from left Much useful for arguments input. Case is ignored.

Usage

```
left_equal(x1, x2)
```

Arguments

x1 one character
x2 the other character

Value

logical

Examples

```
left_equal('o', 'OK')  
left_equal('ok', 'O')  
left_equal('ok', 'Ok')
```

legal	<i>Check legal character Whether the character is legal for names in dataframe or formula</i>
-------	-----------------------------------------------------------------------------------------------

Description

Check legal character Whether the character is legal for names in dataframe or formula

Usage

```
legal(...)
```

Arguments

... one or more string

Value

logical, TRUE means legal.

Examples

```
legal('a', 'b', 'a b')
```

load_extdata	<i>Load external data from R package</i>
--------------	------------------------------------------

Description

Load external data from R package

Usage

```
load_extdata(package, file)
```

Arguments

package	one package name
file	one file name

Value

path of data

mid	<i>Truncate Characters from the Inside</i>
-----	--------------------------------------------

Description

Truncate Characters from the Inside

Usage

```
mid(x, start, n = 1e+11)
```

Arguments

x	can be number, strings, vectors, dataframe or matrix.
start	starting position
n	length, n can be less than zero

Value

substring

Examples

```
mid("abcd",3,1)
mid(c("abc","gjh"),2,2)
df = data.frame(
  a = c(123,234,456),
  b = c("abc","bcd","hjk")
)
mid(df,2,1)
mid(df,2,-2)
```

mirror.current	<i>Current mirror</i>
----------------	-----------------------

Description

Current mirrors of CRAN and Bioconductor

Usage

```
mirror.current()
```

Value

a list contains CRAN and Bioconductor mirror

Examples

```
mirror.current()
```

mirror.set	<i>Set mirror</i>
------------	-------------------

Description

Set mirror

Usage

```
mirror.set(cran, bioc)
```

Arguments

cran	one cran mirror
bioc	one bioconductor mirror

mirror.speed	<i>Test speed of mirror</i>
--------------	-----------------------------

Description

Test speed of mirror

Usage

```
mirror.speed(min.second = 0.2, cran = TRUE, bioc = TRUE)
```

Arguments

min.second	the minium second time to visit the mirror web page
cran	logical, whether to test CRAN mirrors. Default is TRUE
bioc	logical, whether to test bioconductor mirrors. Default is TRUE

Value

repositories which visiting time is minus the minium second.

model.data	<i>Extract data of model</i>
------------	------------------------------

Description

Extract data of model

Usage

```
model.data(fit)
```

```
model.y(fit)
```

```
model.x(fit)
```

Arguments

fit	fitted results
-----	----------------

Value

dataframe in the model

Examples

```
fit <- lm(mpg~vs+am+poly(qsec,2),data=mtcars)
head(model.data(fit))
model.y(fit)
model.x(fit)
```

NA.col.prob	<i>Proportion of missing value by column</i>
-------------	----------------------------------------------

Description

NA is treated as missing value.

Usage

```
NA.col.prob(data)
```

Arguments

data must be dataframe or matrix

Value

proportion of missing value by column

Examples

```
df = data.frame(x=rep(c(1,NA,2,NA,6,NA),10),
                y=rep(c(1,NA,2),20))
NA.col.prob(df)
```

NA.col.sums	<i>Sum of missing value by column</i>
-------------	---------------------------------------

Description

NA is treated as missing value.

Usage

```
NA.col.sums(data)
```

Arguments

data must be dataframe or matrix

Value

sum of missing value by column

Examples

```
df = data.frame(x=rep(c(1,NA,2,NA,6,NA),10),
               y=rep(c(1,NA,2),20))
NA.col.sums(df)
```

NA.row.prob	<i>Proportion of missing value by row</i>
-------------	-------------------------------------------

Description

NA is treated as missing value.

Usage

```
NA.row.prob(data)
```

Arguments

data must be dataframe or matrix

Value

proportion of missing value by row

Examples

```
df = data.frame(x=rep(c(1,NA,2,NA,6,NA),10),
               y=rep(c(1,NA,2),20))
NA.row.prob(df)
```

NA.row.sums	<i>Sum of missing value by row</i>
-------------	------------------------------------

Description

NA is treated as missing value.

Usage

```
NA.row.sums(data)
```

Arguments

data must be dataframe or matrix

Value

sum of missing value by row

Examples

```
df = data.frame(x=rep(c(1,NA,2,NA,6,NA),10),
                y=rep(c(1,NA,2),20))
NA.row.sums(df)
```

NA.whole.prob	<i>Proportion of missing value in the whole dataframe</i>
---------------	-----------------------------------------------------------

Description

NA is treated as missing value.

Usage

```
NA.whole.prob(data)
```

Arguments

data must be dataframe or matrix

Value

proportion of missing value in the whole dataframe

Examples

```
df = data.frame(x=rep(c(1,NA,2,NA,6,NA),10),
                y=rep(c(1,NA,2),20))
NA.whole.prob(df)
```

NA.whole.sums	<i>Sum of missing value in the whole dataframe</i>
---------------	----------------------------------------------------

Description

NA is treated as missing value.

Usage

```
NA.whole.sums(data)
```

Arguments

data must be dataframe or matrix

Value

sum of missing value in the whole dataframe

Examples

```
df = data.frame(x=rep(c(1,NA,2,NA,6,NA),10),
                y=rep(c(1,NA,2),20))
NA.whole.sums(df)
```

names_n	<i>Names with different letters</i>
---------	-------------------------------------

Description

Names with different letters

Usage

```
names_n(df, most = NULL, least = NULL)
```

Arguments

df dataframe or matrix
most names with at most different letters, which means \leq
least names with at least different letters, which means \geq

Value

names

Nchar *Number of Characters*

Description

Number of Characters

Usage

Nchar(x)

Arguments

x can be number, strings, vectors, dataframe or matrix.

Value

number of characters in each location

Examples

```
Nchar("abcd")
Nchar(c("abc", "gjh"))
df = data.frame(
  a = c(1, 12, 12.3),
  b = c("a", "ab", "abc")
)
Nchar(df)
```

numeric.nms *Return numeric names in matrix or dataframe*

Description

Return numeric names in matrix or dataframe

Usage

numeric.nms(df)

Arguments

df dataframe or matrix

Value

numeric names vectors

package_all	<i>Get all functions in one package</i>
-------------	-----------------------------------------

Description

Get all functions in one package

Usage

```
package_all(x)
```

Arguments

x	package
---	---------

Value

all functions in one package

paste0_columns	<i>Paste Columns Together</i>
----------------	-------------------------------

Description

Paste each column in a dataframe together.

Usage

```
paste0_columns(df, collapse = ",")
```

Arguments

df	a dataframe
collapse	collapse, default is comma

Value

a character

Examples

```
df=data.frame(a=c(1,2,30),
              b=c('x','y','z'))
paste0_columns(df)

df=data.frame(a=c(1,2,30),b=c('x','y','z'),c=c(1,7,8))
paste0_columns(df)
```

pipe	<i>pipe</i>
------	-------------

Description

pipe

Usage

pipe()

rd2df	<i>Convert package Rd file under man directory into dataframe</i>
-------	-------------------------------------------------------------------

Description

Convert package Rd file under man directory into dataframe

Usage

rd2df(pkg)

Arguments

pkg	source package path unzip from "tar.gz" file
-----	----------------------------------------------

Value

one dataframe

read_R	<i>Read R file</i>
--------	--------------------

Description

Read R file

Usage

read_R(R)

Arguments

R	path of R file
---	----------------

Value

one vector of R command with names of R file

Replace

Replace

Description

There are two methods in this function. You can use `replace` many objects to one by `from` and `to`. `pattern` can be used to one object replaced by the other one.

Usage

```
Replace(data, from, to, pattern, ignore.case = FALSE)
```

Arguments

<code>data</code>	can be number, strings, vectors, dataframe or matrix.
<code>from</code>	replaced strings
<code>to</code>	replacements
<code>pattern</code>	like <code>from:to</code>
<code>ignore.case</code>	logical, whether to ignore case

Value

replaced data

Examples

```
Replace(data = 232, from = 2, to = 1)
Replace(data = c(232, 'a4b'),
        from = c(2, '.*4'), to = 1,
        pattern = c('a:e', 'b:h'))
df = data.frame(
  a = c(232, 452),
  b = c("nba", "cba")
)
Replace(data = df,
        from = 2, to = 1,
        pattern = c('a:e', 'b:h'))
```

Replace0	<i>Replaced by Empty</i>
----------	--------------------------

Description

Replaced by Empty

Usage

```
Replace0(data, ...)
```

Arguments

data	can be number, strings, vectors, dataframe or matrix.
...	replaced stings

Value

replaced data

Examples

```
Replace0(data = 232,2)
Replace0(data = c(232,'a4b'),2,'.*4')

df = data.frame(
  a = c(232, 452),
  b = c("nba", "cba")
)
Replace0(data = df, 2,'a')
```

Replace_ex	<i>Replace Exactly</i>
------------	------------------------

Description

Replace Exactly

Usage

```
Replace_ex(x, from, to, pattern)
```

Arguments

x vector, dataframe or matrix
 from replaced stings
 to replacements
 pattern a special pattern, see examples for detail

Value

replaced data

Examples

```
a=c(1,2,3,1,4)
Replace_ex(x = a, from = c(1,2), to=5)
Replace_ex(x=a, pattern = c('1:5', '2:5'))
Replace_ex(x=a, pattern = '[12]:5')
```

```
a=data.frame(v=c(1,2,3,2,4),
             b=c(7,8,9,4,6))
Replace_ex(x = a, from = c(1,2), to=5)
Replace_ex(x=a, pattern = c('1:5', '2:5'))
```

 replicate

Replicate Each Elements of Vectors

Description

Replicate Each Elements of Vectors

Usage

```
rep_n(x, each)
```

```
rep_character(x, each)
```

Arguments

x vectors
 each one or more numbers for replication

Value

replicated vectors

Examples

```
rep_n(c('ab', 'cde', 'k', 'op'), 5)
rep_n(c('ab', 'cde', 'k', 'op'), c(4, 6))
rep_n(c('ab', 'cde', 'k', 'op'), c(1, 2, 3, 4))

rep_character(c('ab', 'cde', 'k', 'op'), 5)
rep_character(c('ab', 'cde', 'k', 'op'), c(4, 6))
rep_character(c('ab', 'cde', 'k', 'op'), c(1, 2, 3, 4))
```

reshape_toLong	<i>Convert Wide Data to Long</i>
----------------	----------------------------------

Description

It is easy to convert wide data to long in this function. Be careful, id must be unique. prefix, suffix and var.names can be used together.

Usage

```
reshape_toLong(data, prefix = NULL, suffix = NULL, var.names = NULL)
```

Arguments

data	wide data
prefix	prefix of value variables
suffix	suffix of value variables
var.names	names of value variables, do.value will be created as the name of value column

Value

long data

reshape_toWide	<i>Reshape to Wide Format</i>
----------------	-------------------------------

Description

Reshape to Wide Format

Usage

```
reshape_toWide(
  data,
  key = NULL,
  value = NULL,
  prefix = NULL,
  suffix = NULL,
  sep = "_"
)
```

Arguments

data	long data
key	column names for key, which can be one or more
value	column names for exchange, which can be one or more
prefix	column names for prefix, which can be one or more
suffix	column names for suffix, which can be one or more
sep	seperation

Value

A wide data.

reshape_towide	<i>Reshape to Wide Format</i>
----------------	-------------------------------

Description

Reshape to Wide Format

Usage

```
reshape_towide(data, id, col_change, prefix, suffix, sep = "_")
```

Arguments

data	long data
id	column names for id, which can be one or more
col_change	column names for exchange, which can be one or more
prefix	column names for prefix, which can be one or more
suffix	column names for suffix, which can be one or more
sep	seperation

Value

A wide data.

Examples

```
df = data.frame(id = c(1,1,2,2,3,3,4,4),
               time = c(1,2,1,2,1,2,1,2),
               w = c(1,6,2,7,3,8,4,9))

reshape_towide(data = df,
              id = 'id',
              col_change = 'time',
              prefix = 'w')
df = data.frame(id = c(1,1,2,2,3,3,4,4),
               time = c(1,2,1,2,1,2,1,2),
               w = c(1,6,2,7,3,8,4,9),
               h = c(5,1,6,3,7,5,8,7),
               n = c(2,2,3,3,4,4,5,5))

reshape_towide(data = df,
              id = 'id',
              col_change = c('time','n'),
              prefix = 'w',sep = '_')

reshape_towide(data = df,
              id = 'id',
              col_change = 'time',
              prefix = c('w','h'))

reshape_towide(data = df,
              id = c('id','n'),
              col_change = 'time',
              prefix = c('w','h'))

df = data.frame(id = c(1,1,2,2,3,3,4,4),
               time = c('a','a','a','b','b','b','c','c'),
               n=c(5,5,6,6,7,7,8,8))
reshape_towide(data = df,id = 'id',col_change = 'time')
reshape_towide(data = df,id = c('id','time'),col_change = 'n')
reshape_towide(data = df,id = 'id',col_change = c('time','n'))
```

reverse

Reverse String Order

Description

Reverse String Order

Usage

```
reverse(x)
```

Arguments

x can be number, strings, vectors

Value

reversed string

Examples

```
reverse(123)
reverse(c(123, 'abc'))
```

right

Truncate Characters from the Right

Description

Truncate Characters from the Right

Usage

```
right(x, n)
```

Arguments

x can be number, strings, vectors, dataframe or matrix.
n length

Value

substring

Examples

```
right("abcd",3)
right(c("abc", "gjh"),2)
df = data.frame(
  a = c(123,234,456),
  b = c("abc", "bcd", "hjk")
)
right(df,2)
```

right_equal	<i>Compare two characters from right Much useful for arguments input. Case is ignored.</i>
-------------	--------------------------------------------------------------------------------------------

Description

Compare two characters from right Much useful for arguments input. Case is ignored.

Usage

```
right_equal(x1, x2)
```

Arguments

x1	one character
x2	the other character

Value

logical

Examples

```
right_equal('k','OK')
right_equal('ok','k')
right_equal('ok','Ok')
```

rm_all	<i>Remove all objects</i>
--------	---------------------------

Description

Remove all objects

Usage

```
rm_all()
```

Value

empty object

rm_nchar	<i>Remove elements by number of characters</i>
----------	------------------------------------------------

Description

Remove elements by number of characters

Usage

```
rm_nchar(x, least, most)
```

Arguments

x	one vector
least	least number of characters
most	most number of characters

Value

removed vector

Examples

```
x <- c('a', 'abc', 'abcd', NA)
rm_nchar(x, least = 1)
rm_nchar(x, most = 4)
rm_nchar(x, least = 1, most = 4)
```

row.freq	<i>Row Frequency</i>
----------	----------------------

Description

Row Frequency

Usage

```
row.freq(x)
```

Arguments

x	dataframe or matrix
---	---------------------

Value

data with frequency column

Examples

```
row.freq(x=mtcars[,8:11])
```

select	<i>Subset data Take subset data for</i>
--------	-----------------------------------------

Description

Subset data Take subset data for

Usage

```
select(data, i, ...)  
  
## S3 method for class 'character'  
select(data, i, ...)  
  
## S3 method for class 'numeric'  
select(data, i, ...)  
  
## S3 method for class 'logical'  
select(data, i, ...)  
  
## S3 method for class 'data.frame'  
select(data, i, j, drop = FALSE, ...)  
  
## S3 method for class 'matrix'  
select(data, i, j, drop = FALSE, ...)  
  
## S3 method for class 'list'  
select(data, i, j, drop = FALSE, ...)
```

Arguments

data	one vector, list, dataframe or matrix
i	element position for vector or list, row number for dataframe or matrix
...	ignore
j	column number for dataframe or matrix
drop	logical, whether to drop original format, default is FALSE

Value

selected data

Examples

```
x <- c('ab', 'bc', 'd')
x |> select(!grepl('a'))
x |> select(grepl('a'))
x |> select(!grepl('a'))
x |> select(grepl('a'))

x <- mtcars
x |> select(!grepl('m', ignore.case = TRUE))
x |> select(grepl('m', ignore.case = TRUE), grepl('m', ignore.case = TRUE))
x |> select(!grepl('m', ignore.case = TRUE), !grepl('m', ignore.case = TRUE))

x |> select(grepl('a') & grepl('m'))
x |> select(grepl('a|m'))
x |> select(am == 1)
```

seq_range

sequence range of one vector

Description

sequence range of one vector

Usage

```
seq_range(x, by = 1)
```

Arguments

x	one vector
by	default is 1

Value

number sequence

Examples

```
seq_range(letters)
seq_range(letters, 2)
```

show_function	<i>Show function command line in new script script will be store in your temporary directory</i>
---------------	--------------------------------------------------------------------------------------------------

Description

Show function command line in new script script will be store in your temporary directory

Usage

```
show_function(f, file = NULL)
```

Arguments

f	one function
file	file name

Value

command line in new script

split_expand	<i>Split One Column and Expand</i>
--------------	------------------------------------

Description

Split One Column and Expand

Usage

```
split_expand(data, variable, sep)
```

Arguments

data	dataframe or matrix
variable	one column name with connected values
sep	seperated symbol, which can be one or more

Value

expanded dataframe or matrix

Examples

```
df=data.frame(a=c(1,0),
              b=c('a','n'),
              cyl=c('6;6;4;4;4',
                   '6;8;'))
split_expand(data=df,variable='cyl',sep=';')
```

table_NA	<i>Count NA</i>
----------	-----------------

Description

Count NA

Usage

```
table_NA(x)
```

Arguments

x object

Value

NA and Not count

Examples

```
a <- c(1,2,3,1,NA,NA)
table_NA(a)
```

take_out	<i>Extract Some String</i>
----------	----------------------------

Description

Extract Some String

Usage

```
take_out(x, ..., type = "c")
```

Arguments

x string
 ... patterns of c('begin','after')
 type any left characters of character or list

Value

characters

Examples

```
x='abdghtyu'
take_out(x,c('a','d'),c('h','u'))
```

Target

Download data from Target database

Description

Download data from Target database

Usage

Target_download_dir(url)

Target_download_file(url)

Target_download_url(url)

Arguments

url url link of data

Value

invisible url and downloaded data

Examples

```
Target_download_dir('https://target-data.nci.nih.gov/Public/ALL/clinical/')
```

```
# download data by file url
```

```
Target_download_dir("https://target-data.nci.nih.gov/Public/OS/Brazil/gene_expression_array/METADATA/MANIFEST.")
```

```
# get file url
```

```
Target_download_dir('https://target-data.nci.nih.gov/Public/ALL/clinical/')
```

Trim	<i>Trim</i>
------	-------------

Description

Trim

Usage

Trim(x, pattern = " ")

Trim_left(x, pattern = " ")

Trim_right(x, pattern = " ")

Arguments

x	can be vector or dataframe or matrix
pattern	one or more pattern pattern

Value

a trimmed string

unique_no.NA	<i>Unique Without NA</i>
--------------	--------------------------

Description

Unique Without NA

Usage

unique_no.NA(x)

Arguments

x	vector
---	--------

Value

unique values with no NA

Examples

```
x=c(1,2,3,1,NA)
unique(x)
unique_no.NA(x)
```

unlibray	<i>Detach package</i>
----------	-----------------------

Description

Detach package

Usage

unlibray(x)

Arguments

x one package name, if missing, detach all packages

Value

detach one package

upper.dir	<i>up level directory</i>
-----------	---------------------------

Description

up level directory

Usage

upper.dir(dir, end.slash = TRUE)

Arguments

dir present directory or file path
end.slash logical

Value

upper directory

write_xlsx	<i>Write data to Excel file write or append one or more data into one Excel file in each sheet.</i>
------------	-----------------------------------------------------------------------------------------------------

Description

Write data to Excel file write or append one or more data into one Excel file in each sheet.

Usage

```
write_xlsx(  
  ...,  
  file,  
  sheet,  
  col.names = TRUE,  
  row.names = FALSE,  
  overwrite = FALSE,  
  append = FALSE  
)
```

Arguments

...	one or more data
file	Excel file name
sheet	sheet names
col.names	logical, whether to write out column names
row.names	logical, whether to write out row names
overwrite	logical, whether to overwrite an existing file
append	logical, whether to add data to an existing file

Value

write one or more data into one Excel file

Examples

```
mtcars2 = mtcars  
write_xlsx(mtcars,mtcars2,file='mtcars')
```

%==%

Locate Accurately

Description

Locate Accurately

Usage

a %==% b

Arguments

a vector for matching
b vector for searching

Value

If length of a is one, a vector will be return. If length of a is more than one, a list for each element will be return.

Examples

```
a=c(1,2,3,4)
b=c(1,2,3,1,4,1,5,6,1,4,1)
a %==% b
```

%+%

Concatenate vectors after converting to character.

Description

Concatenate vectors after converting to character.

Usage

a %+% b

Arguments

a one R objects, to be converted to character vectors.
b one R objects, to be converted to character vectors.

Value

one vector

Examples

```
1 %+% 1
```

%s=%

Locate Similarly by grep()

Description

Locate Similarly by grep()

Usage

```
a %s=% b
```

Arguments

a	vector for matching
b	vector for searching

Value

A list contains location information.

Examples

```
1 %s=% c(1,12,3)
c(1,2) %s=% c(1,12,3)
```

Index

`%+%`, 66
`%==%`, 66
`%s=%`, 67

`add_biocViews`, 4
`all_children`, 4
`Apriori.Basket`, 5
`as.data.frame`, 6
`as.data.frame.rules`, 6
`as.transactions`, 7
`attr_href`, 7

`c.xml_nodeset`, 8
`cat_n`, 8
`character.nms`, 9
`chinese_utf8`, 9
`CIGI`, 10
`CIGI_download (CIGI)`, 10
`CIGI_download_file (CIGI)`, 10
`CIGI_download_url (CIGI)`, 10
`cnOS`, 10
`col_split`, 12
`columntrans`, 11
`common`, 12
`compare`, 13
`complete.data`, 14
`CTD2`, 14
`CTD2_download_dir (CTD2)`, 14
`CTD2_download_file (CTD2)`, 14
`CTD2_download_url (CTD2)`, 14

`decrease`, 15
`delete_left`, 16
`delete_up`, 16
`deparse0`, 17
`desc2df`, 17
`detach2`, 18
`dump.it`, 18
`dup.connect`, 19
`duplicate`, 19

`duplicated_all`, 20
`duplicated_last`, 21

`equal (compare)`, 13
`equal_length`, 21
`exec`, 22
`expand`, 23

`factor.it (columntrans)`, 11
`factor.it<- (columntrans)`, 11
`factor.nms`, 23
`file.dir`, 24
`file.name`, 24
`fmt`, 25
`formal_dir`, 26

`get_names`, 26
`give_names`, 27
`Grepl`, 28

`has_children`, 29

`increase`, 29
`inner_Add_Symbol`, 30
`insertglue`, 30
`install_Rversion`, 31
`is.dir`, 31
`is.linux`, 32
`is.mac`, 32
`is.windows`, 33

`join`, 33
`join_full (join)`, 33
`join_inner (join)`, 33
`join_left (join)`, 33
`join_out (join)`, 33
`join_right (join)`, 33

`keep`, 34
`knife`, 35
`knife_left (knife)`, 35

knife_right (knife), 35

last, 35

last_column, 36

last_row, 36

left, 37

left_equal, 37

legal, 38

load_extdata, 39

lower (compare), 13

mid, 39

mirror.current, 40

mirror.set, 40

mirror.speed, 41

model.data, 41

model.x (model.data), 41

model.y (model.data), 41

NA.col.prob, 42

NA.col.sums, 42

NA.row.prob, 43

NA.row.sums, 43

NA.whole.prob, 44

NA.whole.sums, 45

names_n, 45

Nchar, 46

numeric.it (columntrans), 11

numeric.it<- (columntrans), 11

numeric.nms, 46

over (compare), 13

package_all, 47

paste0_columns, 47

pipe, 48

rd2df, 48

read_R, 48

rep_character (replicate), 51

rep_n (replicate), 51

Replace, 49

Replace0, 50

Replace_ex, 50

replicate, 51

reshape_toLong, 52

reshape_toWide, 52

reshape_towide, 53

reverse, 54

right, 55

right_equal, 56

rm_all, 56

rm_nchar, 57

row.freq, 57

select, 58

seq_range, 59

show_function, 60

split_expand, 60

table_NA, 61

take_out, 61

Target, 62

Target_download_dir (Target), 62

Target_download_file (Target), 62

Target_download_url (Target), 62

Trim, 63

trim-left (Trim), 63

trim-right (Trim), 63

Trim_left (Trim), 63

Trim_right (Trim), 63

unique_no.NA, 63

unlibray, 64

upper.dir, 64

write_xlsx, 65