

Package ‘listarrays’

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Type Package

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Description A toolbox for R arrays. Flexibly split, bind, reshape, modify, subset and name arrays.

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<https://t-kalinowski.github.io/listarrays/>

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Author Tomasz Kalinowski [aut, cre]

Maintainer Tomasz Kalinowski <kalinowskit@gmail.com>

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array2	<i>Make or reshape an array with C-style (row-major) semantics</i>
--------	--

Description

These functions reshape or make an array using C-style, row-major semantics. The returned array is still R's native F-style, (meaning, the underlying vector has been reordered).

Usage

```
array2(data, dim = length(data), dimnames = NULL)
```

```
matrix2(...)
```

```
dim2(x) <- value
```

```
set_dim2(...)
```

Arguments

data	what to fill the array with
dim	numeric vector of dimensions
dimnames	a list of dimnames, must be the same length as dims
...	passed on to set_dim()
x	object to set dimensions on (array or atomic vector)
value	a numeric (integerish) vector of new dimensions

Details

Other than the C-style semantics, these functions behave identically to their counterparts (array2() behaves identically to array(), `dim2<-`() to `dim<-`()). set_dim2() is just a wrapper around set_dim(...,order = "C").

See examples for a drop-in pure R replacement to reticulate::array_reshape()

Examples

```

array(1:4, c(2,2))
array2(1:4, c(2,2))

# for a drop-in replacement to reticulate::array_reshape
array_reshape <- listarrays::array_reshape
array_reshape(1:4, c(2,2))

```

bind_as_dim	<i>Bind arrays along a specified dimension</i>
-------------	--

Description

bind_as_* introduces a new dimension, such that each element in list_of_arrays corresponds to one index position along the new dimension in the returned array. bind_on_* binds all elements along an existing dimension, (meaning, the returned array has the same number of dimensions as each of the arrays in the list).

Usage

```

bind_as_dim(list_of_arrays, which_dim)

bind_as_rows(...)

bind_as_cols(...)

bind_on_dim(list_of_arrays, which_dim)

bind_on_rows(...)

bind_on_cols(...)

```

Arguments

list_of_arrays	a list of arrays. All arrays must be of the same dimension. NULL's in place of arrays are automatically dropped.
which_dim	Scalar integer specifying the index position of where to introduce the new dimension to introduce. Negative numbers count from the back. For example, given a 3 dimensional array, -1, is equivalent to 3, -2 to 2 and -3 to 1.
...	Arrays to be bound, specified individually or supplied as a single list

Details

bind_*_rows() is a wrapper for the common case of bind_*_dim(X, 1). bind_*_cols() is a wrapper for the common case of bind_*_dim(X, -1).

Value

An array, with one additional dimension.

Examples

```
list_of_arrays <- replicate(10, array(1:8, dim = c(2,3,4)), FALSE)

dim(list_of_arrays[[1]])

# bind on a new dimension
combined_as <- bind_as_rows(list_of_arrays)
dim(combined_as)
dim(combined_as)[1] == length(list_of_arrays)

# each element in `list_of_arrays` corresponds to one "row"
# (i.e., one entry in along the first dimension)
for(i in seq_along(list_of_arrays))
  stopifnot(identical(combined_as[i,,], list_of_arrays[[i]]))

# bind on an existing dimension
combined_on <- bind_on_rows(list_of_arrays)
dim(combined_on)
dim(combined_on)[1] == sum(sapply(list_of_arrays, function(x) dim(x)[1]))
identical(list_of_arrays[[1]], combined_on[1:2,,])
for (i in seq_along(list_of_arrays))
  stopifnot(identical(
    list_of_arrays[[i]], combined_on[ (1:2) + (i-1)*2,,]
  ))

# bind on any dimension
combined <- bind_as_dim(list_of_arrays, 3)
dim(combined)
for(i in seq_along(list_of_arrays))
  stopifnot(identical(combined[, ,i], list_of_arrays[[i]]))
```

 DIM

Helpers for working with 1-d arrays

Description

DIM() is to dim() as NROW() is to nrow(). That is, it is identical to dim() in most cases except if the input is a bare atomic vector with no dim attribute, in which case, the length of the vector is returned instead of NULL.

DROP first calls base::drop and then completely removes the dim attribute if the result is a 1-d array

Usage

DIM(x)

DROP(x)

Arguments

`x` an R vector, potentially with a dim attribute

Value

For DIM, the dim attribute, or if that's not found, then `length(x)`

For DROP an array with 2 or more axes, or a vector with no dim attributes.

Examples

```
x <- 1:3
dim(x)
dim(array(x))

DIM(x)
DIM(array(x))

x <- array(1:3)
str(drop(x))
str(DROP(x))
```

drop_dimnames

Drop dimnames

Description

A pipe-friendly wrapper for `dim(x) <- NULL` and `dimnames(x) <- NULL` or, if `which_dim` is not `NULL`, `dimnames(x)[which_dim] <- list(NULL)`

Usage

```
drop_dimnames(x, which_dim = NULL, keep_axis_names = FALSE)
```

```
drop_dim(x)
```

```
drop_dim2(x)
```

Arguments

`x` an object, potentially with dimnames

`which_dim` If `NULL` (the default) then all dimnames are dropped. If integer vector, then dimnames only at the specified dimensions are dropped.

`keep_axis_names`

TRUE or FALSE, whether to preserve the axis names when dropping the dimnames

`expand_dims`*Expand the shape of an array*

Description

This is the inverse operation of `base::drop()`. It is analogous to python's `numpy.expand_dims()`, but vectorized on `which_dim`.

Usage

```
expand_dims(x, which_dim = -1L)
```

Arguments

`x` an array. Bare vectors are treated as 1-d arrays.

`which_dim` numeric. Desired index position of the new axis or axes in the returned array. Negative numbers count from the back. Can be any length. Throws a warning if any duplicates are provided.

Value

the array `x` with new dim

Examples

```
x <- array(1:24, 2:4)
dim(x)
dim(expand_dims(x))
dim(expand_dims(x, 2))
dim(expand_dims(x, c(1,2)))
dim(expand_dims(x, c(1,-1)))
dim(expand_dims(x, 6)) # implicitly also expands dims 4,5
dim(expand_dims(x, 4:6))

# error, implicit expansion with negative indexes not supported
try(expand_dims(x, -6))

# supply them explicitly instead
dim(expand_dims(x, -(4:6)))
```

extract_dim	<i>Extract with [on a specified dimension</i>
-------------	--

Description

Extract with [on a specified dimension

Usage

```
extract_dim(X, which_dim, idx, drop = NULL, depth = Inf)
```

```
extract_rows(X, idx, drop = NULL, depth = Inf)
```

```
extract_cols(X, idx, drop = NULL, depth = Inf)
```

Arguments

X	Typically, an array, but any object with a [method is accepted (e.g., dataframe, vectors)
which_dim	A scalar integer or character, specifying the dimension to extract from
idx	A numeric, boolean, or character vector to perform subsetting with.
drop	Passed on to [. If NULL (the default), then drop is omitted from the argument, and the default is used (defaults to TRUE for most objects, including arrays)
depth	Scalar number, how many levels to recurse down if X is a list of arrays. Set this if you want to explicitly treat a list as a vector (that is, a one-dimensional array). (You can alternatively set a dim attribute with dim<- on the list to prevent recursion)

Examples

```
# extract_rows is useful to keep the same code path for arrays of various sizes
X <- array(1:8, c(4, 3, 2))
y <- c("a", "b", "c", "d")
(Y <- onehot(y))

extract_rows(X, 2)
extract_rows(Y, 2)
extract_rows(y, 2)

library(zeallot)
c(X2, Y2, y2) %<-% extract_rows(list(X, Y, y), 2)
X2
Y2
y2
```

<code>map_along_dim</code>	<i>Apply a function across subsets along an array dimension</i>
----------------------------	---

Description

`map_along_dim(X, dim, func)` is a simple wrapper around `split_along_dim(X, dim) %>% map(func)`. It is conceptually and functionally equivalent to `base::apply()`, with the following key differences:

- it is guaranteed to return a list (`base::apply()` attempts to simplify the output to an array, sometimes unsuccessfully, making the output unstable)
- it accepts the compact lambda notation `~.x` just like in `purrr::map` (and `modify_along_dim()`)

Usage

```
map_along_dim(X, .dim, .f, ...)
```

```
map_along_rows(X, .f, ...)
```

```
map_along_cols(X, .f, ...)
```

Arguments

<code>X</code>	an R array	
<code>.dim</code>	which dimension to map along. Passed on to <code>split_along_dim()</code> , and accepts all the same inputs. Valid inputs include	<ul style="list-style-type: none"> • positive integers (index position(s) of dimension), • negative integers (index positions(s) of dimensions, counting from the back), or • character vector (corresponding to array dimnames)
<code>.f</code>	A function, string of a function name, or <code>purrr</code> style compact lambda syntax (e.g. <code>~.x + 1</code>)	
<code>...</code>	passed on to <code>.f()</code>	

Value

An R list

Examples

```
X <- matrix2(letters[1:15], ncol = 3)

apply(X, 1, function(x) paste(x, collapse = "")) # simplifies to a vector
map_along_dim(X, 1, ~paste(.x, collapse = ""))   # returns a list

identical(
```



```

map_along_rows(X, identity),
map_along_dim(X, 1, identity)) # TRUE

identical(
  map_along_cols(X, identity),
  map_along_dim(X, -1, identity)) # TRUE

```

modify_along_dim	<i>Modify an array by mapping over 1 or more dimensions</i>
------------------	---

Description

This function can be thought of as a version of `base::apply()` that is guaranteed to return a object of the same dimensions as it was input. It also generally preserves attributes, as it's built on top of `[<-`.

Usage

```

modify_along_dim(X, which_dim, .f, ...)

modify_along_rows(X, .f, ...)

modify_along_cols(X, .f, ...)

```

Arguments

<code>X</code>	An array, or a list of arrays
<code>which_dim</code>	integer vector of dimensions to modify at
<code>.f</code>	a function or formula defining a function(same semantics as <code>purrr::map()</code>). The function must return either an array the same shape as it was passed, a vector of the same length, or a scalar, although the type of the returned object does not need to be the same as was passed in.
<code>...</code>	passed on to <code>.f()</code>

Value

An array, or if `X` was a list, a list of arrays of the same shape as was passed in.

Examples

```

x <- array(1:6, 1:3)
modify_along_dim(x, 3, ~mean(.x))
modify_along_dim(x, 3, ~.x/mean(.x))

```

ndim	<i>Length of DIM()</i>
------	------------------------

Description

Returns the number of dimensions, or 1 for an atomic vector.

Usage

```
ndim(x)
```

Arguments

x	a matrix or atomic vector
---	---------------------------

onehot_with_decoder	<i>Convert vector to a onehot representation (binary class matrix)</i>
---------------------	--

Description

Convert vector to a onehot representation (binary class matrix)

Usage

```
onehot_with_decoder(y, order = NULL, named = TRUE)
```

```
onehot(y, order = NULL, named = TRUE)
```

```
decode_onehot(
  Y,
  classes = colnames(Y),
  n_classes = ncol(Y) %||% length(classes)
)
```

```
onehot_decoder(Y, classes = colnames(Y), n_classes = length(classes))
```

Arguments

y	character, factor, or numeric vector
order	NULL, FALSE, or a character vector. If NULL (the default), then levels are sorted with <code>sort()</code> . If FALSE, then levels are taken in order of their first appearance in y. If a character vector, then order must contain all levels found in y.
named	if the returned matrix should have column names
Y	a matrix, as returned by <code>onehot()</code> or similar.

classes	A character vector of class names in the order corresponding to Y's onehot encoding. Typically, colnames(Y). if NULL, then the decoder returns the column number.
n_classes	The total number of classes expected in Y. Used for input checking in the returned decoder, also, to reconstruct the correct dimensions if the passed in Y is missing dim() attributes.

Value

A binary class matrix

See Also

[keras::to_categorical](#)

Examples

```
if(require(zeallot)) {
  y <- letters[1:4]
  c(Y, decode) %<-% onehot_with_decoder(y)
  Y
  decode(Y)
  identical(y, decode(Y))
  decode(Y[2,,drop = TRUE])
  decode(Y[2,,drop = FALSE])
  decode(Y[2:3,])

  rm(Y, decode)
}

# more peicemeal functions
Y <- onehot(y)
decode_onehot(Y)

# if you need to decode a matrix that lost colnames,
# make your own decoder that remembers classes
my_decode <- onehot_decoder(Y)
colnames(Y) <- NULL
my_decode(Y)
decode_onehot(Y)

# factor and numeric vectors also accepted
onehot(factor(letters[1:4]))
onehot(4:8)
```

seq_along_dim	<i>Sequence along a dimension</i>
---------------	-----------------------------------

Description

Sequence along a dimension

Usage

```
seq_along_dim(x, which_dim)
```

```
seq_along_rows(x)
```

```
seq_along_cols(x)
```

Arguments

x a dataframe, array or vector. For `seq_along_rows`, and `seq_along_cols` sequence along the first and last dimensions, respectively. Atomic vectors are treated as 1 dimensional arrays (i.e., `seq_along_rows` is equivalent to `seq_along` when `x` is an atomic vector or list).

which_dim a scalar integer or character string, specifying which dimension to generate a sequence for. Negative numbers count from the back.

Value

a vector of integers `1:nrow(x)`, safe for use in for loops and vectorized equivalents.

Examples

```
for (r in seq_along_rows(mtcars[1:4,]))
  print(mtcars[r,])

x <- 1:3
identical(seq_along_rows(x), seq_along(x))
```

set_as_rows	<i>Reshape an array to send a dimension forward or back</i>
-------------	---

Description

Reshape an array to send a dimension forward or back

Usage

```
set_as_rows(X, which_dim)
```

```
set_as_cols(X, which_dim)
```

Arguments

`X` an array

`which_dim` scalar integer or string, which dim to bring forward. Negative numbers count from the back

This is a powered by `base::aperm()`.

Value

a reshaped array

See Also

`base::aperm()` `set_dim()` `keras::array_reshape()`

Examples

```
x <- array(1:24, 2:4)
y <- set_as_rows(x, 3)

for (i in seq_along_dim(x, 3))
  stopifnot( identical(x[, ,i], y[i, ,]) )
```

set_dim

Reshape an array

Description

Pipe friendly `dim<-()`, with option to pad to necessary length. Also allows for filling the array using C style row-major semantics.

Usage

```
set_dim(
  x,
  new_dim,
  pad = getOption("listarrays.autopad_arrays_with", NULL),
  order = c("F", "C"),
  verbose = getOption("verbose")
)
```

Arguments

x	A vector or array to set dimensions on
new_dim	The desired dimensions (an integer(ish) vector)
pad	The value to pad the vector with. NULL (the default) performs no padding.
order	whether to use row-major (C) or column major (F) style semantics. The default, "F", corresponds to the default behavior of R's dim<-(), while "C" corresponds to the default behavior of reticulate::array_reshape(), numpy, reshaping semantics commonly encountered in the python world.
verbose	Whether to emit a message if padding. By default, FALSE.

Value

Object with dimensions set

See Also

set_dim2(), `dim<-`() , reticulate::array_reshape()

Examples

```

set_dim(1:10, c(2, 5))
try( set_dim(1:7, c(2, 5)) ) # error by default, just like `dim<-`()
  set_dim(1:7, c(2, 5), pad = 99)
  set_dim(1:7, c(2, 5), pad = 99, order = "C") # fills row-wise

y <- x <- 1:4
# base::dim<- fills the array column wise
dim(x) <- c(2, 2)
x

# dim2 will fill the array row-wise
dim2(y) <- c(2, 2)
y

identical(x, set_dim(1:4, c(2,2)))
identical(y, set_dim(1:4, c(2,2), order = "C"))

## Not run:
py_reshaped <- reticulate::array_reshape(1:4, c(2,2))
storage.mode(py_reshaped) <- "integer" # reticulate coerces to double
identical(y, py_reshaped)
# if needed, see listarrays::array_reshape() for
# a drop-in pure R replacement for reticulate::array_reshape()

## End(Not run)

```



```

# if the array already has axis names, those are used when possible
nx <- set_dimnames(x, paste0("axis", 1:3))
dimnames(nx)
dimnames(set_dimnames(nx, list(axis2 = c("x", "y", "z"))))
dimnames(set_dimnames(nx, c("x", "y", "z"), which_dim = "axis2"))

# pass NULL to drop all dimnames, or just names along a single dimension
nx2 <- set_dimnames(nx, c("x", "y", "z"), which_dim = "axis2")
nx2 <- set_dimnames(nx2, LETTERS[1:4], which_dim = "axis3")
dimnames(nx2)
dimnames(set_dimnames(nx2, NULL))
dimnames(set_dimnames(nx2, NULL, 2))
dimnames(set_dimnames(nx2, NULL, c(2, 3)))
# to preserve an axis name and only drop the dimnames, wrap the NULL in a list()
dimnames(set_dimnames(nx2, list(NULL)))
dimnames(set_dimnames(nx2, list(NULL), 2))
dimnames(set_dimnames(nx2, list(axis2 = NULL)))
dimnames(set_dimnames(nx2, list(axis2 = NULL, axis3 = NULL)))
dimnames(set_dimnames(nx2, list(NULL), 2:3))

```

shuffle_rows

Shuffle along the first dimension multiple arrays in sync

Description

Shuffle along the first dimension multiple arrays in sync

Usage

```
shuffle_rows(...)
```

Arguments

... arrays of various dimensions (vectors and data.frames OK too)

Value

A list of objects passed on to ..., or if a single object was supplied, then the single object shuffled

Examples

```

x <- 1:3
y <- matrix(1:9, ncol = 3)
z <- array(1:27, c(3,3,3))

if(require(zeallot)) {
  c(xs, ys, zs) %<-% shuffle_rows(x, y, z)

  l <- lapply(seq_along_rows(y), function(r) {

```



```

    list(x = x[r], y = y[r,], z = z[r,,])
  })

  ls <- lapply(seq_along_rows(y), function(r) {
    list(x = xs[r], y = ys[r,], z = zs[r,,])
  })

  stopifnot(
    length(unique(c(l, ls))) == length(l))
}

```

split_on_dim	<i>Split an array along a dimension</i>
--------------	---

Description

Split an array along a dimension

Usage

```

split_on_dim(
  X,
  which_dim,
  f = dimnames(X)[[which_dim]],
  drop = FALSE,
  depth = Inf
)

split_on_rows(X, f = rownames(X), drop = FALSE, depth = Inf)

split_on_cols(X, f = rownames(X), drop = FALSE, depth = Inf)

split_along_dim(X, which_dim, depth = Inf)

split_along_rows(X, depth = Inf)

split_along_cols(X, depth = Inf)

```

Arguments

X	an array, or list of arrays. An atomic vector without a dimension attribute is treated as a 1 dimensional array (Meaning, atomic vectors without a dim attribute are only accepted if which_dim is 1. Names of the passed list are preserved. If a list of arrays, all the arrays must have the same length of the dimension being split.
which_dim	a scalar string or integer, specifying which dimension to split along. Negative integers count from the back. If a string, it must refer to a named dimension (e.g, one of names(dimnames(X))).

f	Specify how to split the dimension. character, integer, factor passed on to <code>base::split()</code> . Must be the same length as the dimension being split. a list of vectors Passed on to <code>base::interaction()</code> then <code>base::split()</code> . Each vector in the list must be the same length as the dimension being split. a scalar integer used to split into that many groups of equal size a numeric vector where <code>all(f<0)</code> specifies the relative size proportions of the groups being split. <code>sum(f)</code> must be 1. For example <code>c(0.2, 0.2, 0.6)</code> will return approximately a 20\ split.
drop	passed on to <code>[]</code> .
depth	Scalar number, how many levels to recurse down. Set this if you want to explicitly treat a list as a vector (that is, a one-dimensional array). (You can alternatively set dim attributes with <code>dim<-</code> on the list to prevent recursion) <code>split_along_dim(X, which_dim)</code> is equivalent to <code>split_on_dim(X, which_dim, seq_along_dim(X, wh</code>

Value

A list of arrays, or if a list of arrays was passed in, then a list of lists of arrays.

Examples

```
X <- array(1:8, c(2,3,4))
X
split_along_dim(X, 2)

# specify f as a factor, akin to base::split()
split_on_dim(X, 2, c("a", "a", "b"), drop = FALSE)

d <- c(10, 3, 3)
X <- array(1:prod(d), d)
y <- letters[1:10]
Y <- onehot(y)

# specify `f` as relative partition sizes
if(require(zeallot) && require(magrittr) && require(purrr)) {

c(train, validate, test) %<-% {
  list(X = X, Y = Y, y = y) %>%
    shuffle_rows() %>%
    split_on_rows(c(0.6, 0.2, 0.2)) %>%
    transpose()
}

str(test)
str(train)
str(validate)

}
```

```
# with with array data in a data frame by splitting row-wise
if(require(tibble))
  tibble(y, X = split_along_rows(X))
```

t.array	<i>transpose an array</i>
---------	---------------------------

Description

transpose an array

Usage

```
## S3 method for class 'array'
t(x)
```

Arguments

x	an array
---	----------

This reverses the dimensions of an array

Examples

```
x <- array(1:27, c(3,3,3))
tx <- t(x)
for (i in 1:3)
  for(j in 1:3)
    stopifnot(x[,j,i] == tx[i,j,])
```

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