# Package 'oceanmap'

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Type Package

Title A Plotting Toolbox for 2D Oceanographic Data

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Author Robert K. Bauer
Maintainer Robert K. Bauer <rkbauer@hawaii.edu></rkbauer@hawaii.edu>
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<b>Description</b> Plotting toolbox for 2D oceanographic data (satellite data, sea surface temperature, chlorophyll, ocean fronts & bathymetry). Recognized classes and formats include netcdf, Raster, '.nc' and '.gz' files.
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2 .get.worldmap

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Description

Creates a world map database that allows longitude ranges between -180 and 360 degrees, and thus from the Pacific to the Atlantic and vise versa. It is based on the worldHires database (which itself is based on CIA World Data Bank II data and contains approximately 2 million points representing the world coastlines and national boundaries), from which polygon irritations of the Antarctic were also corrected.

# Usage

.get.worldmap(resolution)

# Arguments

resolution

number that specifies the resolution with which to draw the map. Resolution 0 is the full resolution of the database [default]. Otherwise, just before polylines are plotted they are thinned: roughly speaking, successive points on the polyline that are within resolution device pixels of one another are collapsed to a single point (see the Reference for further details). Thinning is not performed if plot

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= FALSE or when polygons are drawn (fill = TRUE or database is a list of polygons).

#### Value

A list of class "map" with longitude (x) and latitude (y) positions of coastlines and state boundaries (different coastline or booundary elements are seperated by NA), single polygon names are provided by a names vector.

### Author(s)

Robert K. Bauer

#### See Also

```
worldHires
https://www.evl.uic.edu/pape/data/WDB/
```

# **Examples**

```
worldmap <- oceanmap:::.get.worldmap(worldmap)
str(worldmap)

## wordlmap usage in plotmap, with different center-options
# par(mfrow=c(3,1))
# plotmap(lon=c(80, -120), lat=c(-50, 10), main= "map from East to West")
# plotmap(lon=c(-120, 80), lat=c(-50, 10), main= "map from West to East")
# plotmap('tp')</pre>
```

add.region

adding a region to the region\_definitions file

# Description

adding a region to the region\_definitions-file, taking or restoring a backup of region definitions. The basic idea is to provide a region-keyword that is used to access the region-information in later related function-calls (see: v and plotmap, regions). Information consists of a region-keyword, -longname, its spatial extent (longitudes and latitudes), grid resolution, as well as default colorbar position and figure size.

The required information can be provided by an interactive **session** (widget) that leads step by step through the region definition (is set default), in parts by an **extent-object** with the missing information then completed by the **session** or by a one-row data frame that holds the entire information (see: region\_definitions).

**ATTENTION!** When reinstalling or updating the oceanmap package, previous region definitions are getting lost! It is therefore highly recommanded to take and restore own backups (see: backup and restore).

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#### Usage

#### **Arguments**

add extent-,raster-object or dataset containing all required region definition entries

(label, name, latn, lats, lonw, lone, ncol, nrow, px, cbx1, cbx2, cby1, cby2, lone, ncol, nrow, px, cbx1, cbx2, cbx2, cbx1, cbx2, cbx2, cbx1, cbx2, cbx2, cbx1, cbx2, cbx2

figxdim, figydim and grid.res). Ignored when add.px is supplied.

The values latn, lats, lonw, lone define the regions extent, cbx1, cbx2, cby1 and cby2 define the position of the colorbar, gradient the orientation of the colorbar (x for horizontal, y for vertical), oticks the margin where to put the colorbar ticks relative to the colorbar rectangle ('l' left, 'r' right and 'b' for bottom; figxdim and figydim set the default window size of '.gz'-file figures

and grid.res the default grid resolution.

add.px dataframe or list containing region data needed to read gz-compressed '.gz'-

files. Required entries include 'label' to identify the region, 'ncol' and 'nrow', to define the number of columns and rows of the 'gz'-file, respectively. These values are automatically set if missing when writing gz-compressed '.gz'-files

(see: writebin).

cbx the horizontal limits (x1, x2) of the colorbar. If missing, the user will be asked

for manual colorbar placement.

cby the vertical limits (y1, y2) of the colorbar. If missing, the user will be asked for

manual colorbar placement.

figdim numeric vector indicating the width and height of the plot device in inches. If

missing and force.figdim.widget is set FALSE, figdim is assigned a default width and height of 7in, otherwise the user will be asked to resize the plot

device to set plot dimensions.

lib.folder Character string indicating R-library path in which the oceanmap-package is

installed.

widget whether an interactive session (widget) shall assist the data entry procedure (de-

fault is TRUE).

backup whether the current region\_definitions-file should be backuped in the folder

'backup.folder in the file backup.name (default is FALSE). **ATTENTION!** When reinstalling or updating the oceanmap package, previous region\_definitions

are getting lost!

backup.folder Character string indicating the folder where to store the region\_definitions-file

backup (default is the current working directory).

backup.name Character string indicating the filename of the region\_definitions-file backup (If

restore the default is the original oceanmap-region\_definitions file; if backup

the default is set to 'region\_definitions.bkp.%Y%m%d.rda').

restore whether to restore a backup of the region\_definitions-file (default is FALSE).

backup.regions Vector of region indicators defining which regions should be saved in backup

file.

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### Author(s)

Robert K. Bauer

#### See Also

delete.region, region\_definitions, regions, plotmap, v

```
## Example 1: Add region by supplying a one-row data.frame
##
              that holds the entire required information
# data(region_definitions) # load region_definitions
# lion <- region_definitions[region_definitions$label == 'lion',] # selecting Gulf of Lions region
# lion
# junk <- lion</pre>
# junk$label <- 'junk' # rename region label</pre>
# add.region(junk) # add junk region
# data(region_definitions) # reload region_definitions
# region_definitions[,1:9]
## Example 2: Delete region
#delete.region("junk") # delete junk region
#data(region_definitions) # reload region_definitions
#region_definitions[,1:9]
## Example 3: Add region by supplying an extent- or raster-object and running the widget
library(raster)
ext <- extent(0,10,50,60)
plotmap(ext)
#add.region(ext) # extent-object
r <- raster(ext)
#add.region(r) # raster-object
## Example 4: Add region by supplying raster-object, colorbar positions and running the widget
#add.region(r,cbx=c(5,9.5),cby=c(51.7,52.4))
## Example 5: Add region by running the widget
#add.region()
## Example 6: Add region by running the widget
#add.region(add.px=list(label="lion",nrow=10,ncol=10))
#data(region_definitions)
#region_definitions[region_definitions$label =="lion",]
## Example 7: Creating a backup
#add.region(backup=T)
#add.region(backup=T, backup.folder=".",backup.regions=c("lion","medw4"))
## Example 8: Restoring the backup of the original region_definitions file
```

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```
#add.region(restore=T)
```

area\_extrac

Extracts a pre-defined region from '.gz'-file and saves subset as a new '.gz'-file

# Description

Extracts a pre-defined region from '.gz'-file and saves subset as a new '.gz'-file (gzip compressed format). Basically it represents a combined call of regions, crop, raster2matrix and writebin.

# Usage

```
area_extrac(obj,area)
```

# **Arguments**

obj Character string indicating search criteria for '.gz'-files.

area Character string identifying the region that should be extracted. area must be a

subregion of the original region defined by the '.gz'-file. See region\_definitions

for area definitions and use add.region to add new regions.

# Author(s)

Robert K. Bauer

#### See Also

readbin, writebin, crop, raster2matrix, param\_unconvert

```
## Example 1: extract, write '.gz'-files, following default plot-procedure
library(raster)

# load sample-'.gz'-file
path <- system.file("test_files", package="oceanmap")
gz.files <- Sys.glob(paste0(path,'/*.gz'))[1] # load sample-'.gz'-files
print(gz.files)
# area_extrac(gz.files[1],area='lion')

# gz <- Sys.glob(path,'/medw4*.gz') # load new-'.gz'-file
# v(gz) # visualize new-'.gz'-file
# system(paste('rm', gz))
# v(gz.files[1],v_area='lion')</pre>
```

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bindate2Title

returns formatted date string for v-plot titles

### **Description**

returns formatted date string for v-plot titles by provided date information (e.g. filename of '.gz'-files, name of raster-layers. bindate2Title is returned by default by v-calls. bindate2main and bindate2ylab are plotted when v is called with sidelabels=T.

# Usage

```
bindate2Title(timestep, date1, date2=date1)
bindate2main(timestep, date1, date2=date1)
bindate2ylab(timestep, date1, date2=date1)
```

#### **Arguments**

timestep character string, indicating the range of the time unit in numbers and the time

unit (e.g. "1d" for daily data; "7d" or "1w" for weekly data; "1m" for monthly

data)

date1, date2 character string, indicating the first and last date of the timeframe covered (rec-

ognized format is %Y%m%d%H or %Y%m%d). E.g. 20030301 and 20030331

for monthly data (timestep = 1m) of March 2003.

#### Author(s)

Robert K. Bauer

# See Also

```
name_split, v
```

```
## Example 1: output of different bindate2???-functions
path <- system.file("test_files", package="oceanmap")
gz.files <- Sys.glob(paste0(path,'/*.gz')) # load sample-'.gz'-files
u <- name_split(gz.files)

print(gz.files[1]) # print filename
print(u[1,]) # print splitted filename
bindate2main(u$timestep[1],u$date1[1],u$date2[1]) # main
bindate2Title(u$timestep[1],u$date1[1],u$date2[1]) # Title
bindate2ylab(u$timestep[1],u$date1[1],u$date2[1]) # ylab</pre>
```

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```
## Example 2: Visualize output for multiple '.gz'-files
u$option <- '....'
dev.new(width=9.7,height=7.8,xpos=-1)
empty.plot()
box()
for (i in 1:nrow(u)){
 mtext(name_join(u[i,]),side=1,line=i-10)
 main <- bindate2main(u$timestep[i],u$date1[i],u$date2[i]) # main</pre>
 Title <- bindate2Title(u$timestep[i],u$date1[i],u$date2[i]) # Title</pre>
 ylab <- bindate2ylab(u$timestep[i],u$date1[i],u$date2[i]) # ylab</pre>
 mtext(c(Title,ylab,main),side=1:3,line=c(i,nrow(u)+1-i,nrow(u)+1-i))
 mtext(paste("file",i),side=c(1,1:3),line=c(i-10,i,nrow(u)+1-i,nrow(u)+1-i),adj=0)
}
mtext(c("filename",
        "bindate2Title (default)",
         "bindate2ylab (sidelabels=T)",
         "bindate2main (sidelabels=T)"),
      side=c(1,1:3), line=c(-11, rep(i+2,3)), font=2)
```

check\_gzfiles

Returns summary on '.gz'-file types

### Description

Returns summary table on '.gz'-file types available in a specified folder. Provided information include region (region covered, as described by the region\_definitions), sat (satellite source), param (parameter), res (spatial resolution), ts (temporal resolution), filetype (file filetype)

# Usage

```
check_gzfiles(sstring="*",folder,filetype=".gz")
```

### **Arguments**

sstring	Character string indicating the search criteria for sat files (default is $\star$ , including all '.gz'-files).
folder	Character string indicating the folder in which searched files are located (default is current working directory)
filetype	Character string indicating thefile type of sat files (default is .gz)

# Value

An aggregated data frame, returning '.gz'-file type-information (see description) on available files in a specified folder.

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### Author(s)

Robert K. Bauer

#### See Also

```
name_split, check_ts
```

# **Examples**

```
## Example 1: plot '.gz'-files, following default plot-procedure
path <- system.file("test_files", package="oceanmap")
check_gzfiles(folder=path) # return file summary-table per filetype

## check for missing dates
check_ts('medw4*',folder=path)
check_ts('medw4*',folder=path,output=TRUE)</pre>
```

check\_ts

checks if daily '.gz'-file time series is complete

# **Description**

checks if daily '.gz'-file time series in the present working directory is complete.

### Usage

```
check_ts(sstring="*.gz",folder,output=F)
```

# **Arguments**

sstring Character string indicating search criteria for gz-files (default is '\*.gz').

folder Character string indicating the folder in which searched files are located (default

is current working directory)

output weather the missing dates should be returned as vector (default is F).

#### Value

optional vector of missing dates (see output argument).

# Author(s)

Robert K. Bauer

### See Also

```
name_split, check_gzfiles
```

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# **Examples**

```
## Example 1: plot '.gz'-files, following default plot-procedure
path <- system.file("test_files", package="oceanmap")
check_gzfiles(folder=path) # return file summary-table per filetype

## check for missing dates
check_ts('medw4*',folder=path)
check_ts('medw4*',folder=path,output=TRUE)</pre>
```

 ${\tt clim\_plot}$ 

plots '.gz'-file climatologies

# **Description**

Creates climatology plots of '.gz'-files. **ATTENTION!** This function requires an ImageMagick installation, but runs also under Windows operating systems.

# Usage

# **Arguments**

obj	Character string indicating search criteria for climatology '.gz'-files.
folder	directory where data files are located (optional).
plotfolder	directory where image should be saved.
plotname	the name of the output file. If not provided, value will be derived from ' $.gz'$ -filenames.
question	whether the user shall be informed about the number of figures to plot before running the procedure (default is TRUE).
chla.frontcolor	•
	color map to be plotted for chlorophyll fronts (default is blue; obtained from cmap-dataset)
sst.frontcolor	color map to be plotted for sea surface temperature fronts (default is red; obtained from cmap-dataset)
sidelabels	whether an additional y-axis label and title should be added to the plot device (default is FALSE). If TRUE, y-axis label is defined by Ylab, the additional title is derived from the date-information and gives the month information.
Ylab	an additional title for the y axis (default is date information), only used when sidelabels is set TRUE. Default value is year-information.
axeslabels	whether axes labels should be shown (default is TRUE, set as 'longitude' and 'latitude')

close\_fig

v\_area

character string identifying the region that should be plotted, or in case of obj == 'bathy', also a Raster\* or Extent object. If missing, region is derived from the '.gz'-filename. See region\_definitions for area definitions and use add.region to add new regions.

. . .

Additional arguments to be passed to v and plotmap (e.g. main, sidelabels, Ylab, scale\_arrow, minv, maxv, adaptive.vals, cb.xlab, suffix, v\_area, v\_image, v\_contour, v\_arrows, fill, col, border, grid, grid.res, bwd, axeslabels, ticklabels, cex.lab, cex.ticks)

### Author(s)

Robert K. Bauer

#### See Also

```
v, readbin, name_split, regions, plotmap
```

### **Examples**

```
## Example 1: plot seasonal '.gz'-files, following default plot-procedure
path <- system.file("test_files", package="oceanmap")
gz.file <- Sys.glob(paste0(path,'/*.gz'))[1] # load sample-'.gz'-files
check_gzfiles(folder=path) # return file summary-table
gz.files <- Sys.glob(paste0(path,'/*1s_*.gz')) # load seasonal '.gz'-files
# v(gz.files) # as single plots

## as combined climatology plot, saved in plotfolder
# clim_plot(gz.files,plotname='chla.summary.png')</pre>
```

close\_fig

function to close current graphic device

### **Description**

function to close current graphic device, complement to figure-function that generates graphic devices in flexible fileformats.

### Usage

```
close_fig(do.close=F, do.save=do.close)
```

# Arguments

```
do.close, do.save
```

whether file should be saved or not (default is TRUE). if FALSE, new graphic device will be opened inside R.

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### Author(s)

Robert K. Bauer

### See Also

figure

# **Examples**

```
# do.save <- TRUE
# figure("Gulf_of_Lions", do.save=do.save, width=5, height=5, type="pdf")
# plotmap("lion")
# close_fig(do.save)

do.save <- TRUE
plotmap("lion")
close_fig(do.save)

do.save <- FALSE
figure("Gulf_of_Lions", do.save=do.save, width=5, height=5, type="pdf")
plotmap("lion")
close_fig(do.save)</pre>
```

cmap

color maps

# **Description**

list holding different color maps that can be used in image plots (see: v, get.bathy, image, image.plots, clim\_plot)

available color maps are: ano, bathy, blue, chla, haxby, jet (obtained from matlab), rainbow, red, orange, green, sst and haxbyrev.

# Usage

```
data(cmap)
data(cmap_topo)
```

### **Format**

list

### Author(s)

Robert K. Bauer

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### **Examples**

```
data('cmap') # load color maps data
names(cmap) # list available color maps
path <- system.file("test_files", package="oceanmap")</pre>
gz.files <- Sys.glob(paste0(path,'/*.gz')) # load sample-'.gz'-files</pre>
# figure(width=15,height=15)
# par(mfrow=c(4,5))
# for(n in names(cmap)) v(gz.files[2], v_area='lion', subplot=TRUE,
                          pal=n, adaptive.vals=TRUE, main=n)
## simple example of the \link{image}-function
x <- 10*(1:nrow(volcano))</pre>
y <- 10*(1:ncol(volcano))</pre>
image(x, y, volcano, col = terrain.colors(100))
image(x, y, volcano, col = cmap$jet) # jet color map
image(x, y, volcano, col = cmap$haxby) # haxby color map
image(x, y, volcano, col = cmap$chla) # chlorophyll color map
image(x, y, volcano, col = cmap$sst) # sst color map
data(cmap_topo)
image(x, y, volcano, col = cmap_topo$col) # topography color map
## another example: plot bathymetry and topography of the western Mediterranean Sea
#get.bathy("medw4", visualize=T, terrain=T, res=3)
#get.bathy("medw4",visualize=T,terrain=F,res=3,levels=c(200,2000)) # show contours
```

delete.region

deletes a region from the region\_definitions-definition file

# **Description**

deletes a specified region from the region\_definitions-definition file

# Usage

```
delete.region(region,lib.folder,restore=F)
```

# **Arguments**

region	Character string identifying the region that should be deleted. See region_definitions for area definitions and use add.region to add new regions.
lib.folder	Character string indicating R-library path in which the oceanmap-package is installed.
restore	whether the original region_definitions-file should be restored.

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### Author(s)

Robert K. Bauer

#### See Also

add.region, region\_definitions, regions, writebin

#### **Examples**

```
## Example 1: Add region by supplying a one-row data.frame
## that holds the entire required information
data(region_definitions)
lion <- region_definitions[region_definitions$label == 'lion',] # selecting Gulf of Lions region
lion
junk <- lion
junk$label <- 'junk' # rename region label
#add.region(junk) # add junk region
data(region_definitions) # reload region_definitions
region_definitions[,1:9]

## Example 2: Delete region
#delete.region("junk") # delete junk region
data(region_definitions) # reload region_definitions
region_definitions[,1:9]</pre>
```

empty.plot

Creates an empty scatter plot

# Description

Creates an empty scatter plot that is equal to the function call:

```
plot(1,lwd=0,axes=F,xlab="",ylab="",...)
```

### Usage

```
empty.plot(..., xlab = "", ylab = "", new=T, add=!new, n=1, axes = F)
```

# Arguments

```
other arguments of the generic x-y plotting fucntion plot.

xlab, ylab label for the x- and y-axis of the plot (default is empty).

new, add whether to show add plot to a current plot device or to start a new figure (default is: new=TRUE and add=FALSE).

n number of figures to be plotted (default is 1)

axes whether to show plot axes (default is FALSE).
```

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### Author(s)

Robert K. Bauer

### **Examples**

```
empty.plot()
title("empty plot")
box()
axis(1)
axis(2)
```

figure

generate (and save) graphic devices with flexible fileformat selection

### **Description**

figure generates graphic devices with flexible fileformat selection. Function call with (figure(do.save=T) needs to be finished by close\_fig(do.save=T), to close open file connection.

### Usage

```
figure(filename, folder, type, save=F, do.save=save,
    width=10, height=10, xpos=-1, do.overwrite=T, delete.old=do.overwrite, ...)
```

# **Arguments**

filename name of the figure to be generated (without file extension) plot folder (by default current working directory) folder type character string indicating the graphics format of the figure file. can be: • "jpg" • "jpeg" • "png" • "tiff" • "eps" • "pdf" width, height width and height of figure to be generated. default units are inches. save, do.save whether file should be saved or not (default is TRUE). if FALSE, new graphic device will be opened inside R. xpos horizontal screen position of graphic device (ignored if do.save == TRUE) do.overwrite, delete.old

additional arguments to be passed to the graphic device

overwrite existing figure with same filename and extension (default is FALSE)

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### Author(s)

Robert K. Bauer

### See Also

```
close_fig
```

# **Examples**

```
## Example 1: plotmap() and figure()
do.save <- FALSE
figure("Gulf_of_Lions_extended", do.save=do.save, width=5, height=5, type="pdf")
plotmap("lion")
close_fig(do.save)

## now resize figure manually and get new figure dimensions:
width <- dev.size()[1]
height <- dev.size()[2]

# do.save <- TRUE
# figure("Gulf_of_Lions_extended", do.save=do.save, width=width, height=height, type="pdf")
# plotmap("lion")
# close_fig(do.save)</pre>
```

get.avg.bathy

returns the average value of circles with specified coordinates and a defined radius

# **Description**

returns the average value of circles with specified coordinates and a defined radius

### Usage

```
get.avg.bathy(x, radius, unit="km", raster, bathy, v_area="medw4")
get.avg(x, radius, unit="km", raster)
```

# Arguments

x a vector or matrix providing the coordinates of the circle

radius the radius (if unit != "km", the radius is assumed to be of the same scale as the

plotting window)

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unit	unit of the radius (by default "km"). if unit != "km", the radius is assumed to be of the same scale as the plotting window.
raster	raster object, from which the average should be calculated.
bathy	raster object with bathymetry data, from which the average should be calculated.
v_area	character string identifying the region for which the bathymetry data should downloaded from the NOAA server.

# Author(s)

Robert K. Bauer

### See Also

SpatialCircle

### **Examples**

```
## Example 1: load & plot bathymetry of the Baltic Sea, defined by longitudes and latidtues
lon <- c(9, 31)
lat <- c(53.5, 66)
# bathy <- get.bathy(lon=lon, lat=lat, main="Baltic Sea", cbpos='r')
plotmap(lon=lon, lat=lat)
spc <- SpatialCircle(x= 20,y = 57.5,r=1)
plot(spc,add=TRUE)
# get.avg.bathy(c(20,57.5), radius = 1, bathy = bathy)
# get.avg(c(20,57.5), radius = 1, unit="km",raster = bathy)</pre>
```

get.bathy	Returns bathymetric data from the NOAA ETOPO1 database as
	RasterLayer, given coordinate bounds and resolution.

# **Description**

Returns and optionally stores bathymetric data from the ETOPO1 database hosted on the NOAA server as a RasterLayer, based on the defined resolution and provided coordinate bounds or region definition. Stored bathymetry files can be reloaded through the same function call.

# Usage

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# Arguments

character string identifying the region that should be plotted, or in case of x =v\_area 'bathy', also a Raster\* or Extent object. If missing, region is derived from the '.gz'-filename. See region\_definitions for area definitions and use add.region to add new regions. lon, lat longitude and latitude describing the extend of the region of interest. resolution resolution of the bathymetric grid, in minutes (default is 4). whether to write the data downloaded from NOAA into a file (default is FALSE). keep savename.bathy savename for the bathymetric data file, if not specified set to type 'bathy\_lonlat\_res.resolution.dat' or 'bathy\_v\_area\_res.resolution.dat'. directory where bathymetric data should be saved (default is current working folder.bathy directory). visualize whether the bathymetric data should be plotted instantly. terrain whether the to keep terrain data (default is FALSE). If set FALSE and visualize is TRUE, grid command in plotmap is disabled!

additional arguments to be passed to v, used if visualize is set TRUE.

# Author(s)

Robert K. Bauer

### See Also

v, add.region, region\_definitions, regions, writebin, get.bathy

```
## Example 1: load & plot bathymetry of the Baltic Sea, defined by longitudes and latidtues
lon <- c(9, 31)
lat <- c(53.5, 66)
# get.bathy(lon=lon, lat=lat, main="Baltic Sea", cbpos='r')
## Example 2: plot bathymetry using a v_area-keyword
#get.bathy("lion",res=4, keep=T) # can take some time, requires server connection!
#get.bathy("lion",res=1, keep=T,visualize=FALSE)
## Example 3: plot landmask of the Baltic Sea defined by an extent- or raster-object
library('raster')
ext <- extent(lon,lat)</pre>
#get.bathy(ext,visualize=T,main="Baltic Sea",res=4,levels=200) # extent-object
## Example 4: plot bathymetry and topography of the western Mediterranean Sea
### a) download, assign and save bathymetry
# bathy <- get.bathy("medw4",visualize=F,terrain=T,res=3,keep=T)</pre>
# # load('bathy_medw4_res.3.dat',verbose = T); bathy <- h</pre>
# par(mfrow=c(2,1))
# v(bathy,param="bathy",subplot = T)
```

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```
# get.bathy("medw4",visualize=T,terrain=F,res=3,levels=c(200,2000),
# subplot = T,grid=F) # show contours

### b) only contour lines:
# par(mfrow=c(1,2))
# h <- get.bathy("lion",visualize=T,terrain=F,res=3,levels=c(200,2000),
# v_image=F, subplot=T,grid=F)

### use v-function for same plot but on subregion:
# v(h,v_area = "survey", param="bathy",subplot = T, v_contour = T,
# v_image = F, levels=c(200,2000))</pre>
```

ggplotmaply

Converts a ggplot2 object from ggplotmap() to plotly

### **Description**

This function converts a ggplot2 object created by oceanmap::ggplotmap() to a plotly object.

# Usage

```
ggplotmaply(ggobj, fixedrange=F, grid=F,expand=3)
```

### **Arguments**

ggobj (	Character string identifying regio	ns predefined b	y the region_	_definitions-dataset,

Raster\* or Extent object (corresponds to v\_area of the v-function). If missing, region is derived from geographical coordinates, denoted by lat and lon. See add.region to define new region definitions and delete.region to delete unproper

region definitions.

fixedrange Vector returning longitude coordinates of the area to be plotted.

grid whether a grid should be plotted (default is TRUE)

expand By default, the underlying ggplotly-function does not stick to the plotting region

of the ggobj, but extends it. This can result in missing countries or islands. The expand-argument extends the plotly-plotting window in each direction in order

to cover the corresponding landmasks.)

#### **Details**

ggoplotmaply uses the ggplotly functions to convert the ggplot object into the plotly format.

### Author(s)

Robert K. Bauer

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#### See Also

```
ggplotmap, ggplotly
```

```
library(ggplot2)
#### Example 1: plot landmask of the Western Mediterranean Sea
## a) by using longitude and latitude coordinates:
\# lon <- c(-6, 16.5)
# lat <- c(34, 44.5)
# ggobj <- ggplotmap(xlim=lon, ylim=lat)</pre>
# ggplotmaply(ggobj,expand = 10) ## we need to expand the plotting region
## b) plot landmask of the Western Mediterranean Sea by using an extent-object:
# library('raster')
# ext <- extent(lon, lat)</pre>
# plotmap(ext, main="Western Mediterranean Sea") # extent-object
# ggobj <- ggplotmap(ext)</pre>
# ggobi
# ggplotmaply(ggobj)
## c) plot landmask of the Western Mediterranean Sea by using a raster-object:
# r <- raster(ext)</pre>
# ggobj <- ggplotmap(r)</pre>
# ggobj
# ggplotmaply(ggobj)
# ggplotmaply(ggobj)
## d) plot landmask of the entire Mediterranean Sea by using keyword:
ggobj <- ggplotmap("med4") +</pre>
          geom_point(data=data.frame(x=3.7008, y=43.4079),aes(x,y),size=5,colour="blue")
# ggobj
# ggplotmaply(ggobj,expand = 10)
## e) add landmask to raster image plot (similar to v()-call)
# library(dplyr)
# library(ggplot2)
# data(cmap)
# setwd(system.file("test_files", package="oceanmap"))
# nc <- nc2raster(ncfiles[1])</pre>
# rs2df <- nc[[1]] %>% ## take first layer
           rasterToPoints() %>% ## convert raster to xyz matrix
           as.data.frame() ## convert to data frame
# names(rs2df) <- c("Lon","Lat","Conc") ## reset names (important for ggplotmaply hover text)</pre>
# ggobj <- ggplot() + geom_raster(data = rs2df, aes(x=Lon,y=Lat,fill=Conc))</pre>
# ggobj_with_land_mask <- ggplotmap(add_to = ggobj) +</pre>
#
                           scale_fill_gradientn(colours=cmap$jet) # change colorbar
```

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```
# ggobj_with_land_mask
# ggplotmaply(ggobj_with_land_mask)
```

internal.datasets

internal datasets

# Description

internal (lazyload) datasets medm9\_proj and regions.dim.bathy, accessed by v.plot and readbin respectively.

# Author(s)

Robert K. Bauer

matrix2raster

Converts a matrix to a RasterLayer or arrays to a RasterStack-object

# Description

matrix2raster Converts a matrix to a RasterLayer or arrays to a RasterStack-object.

# Usage

```
matrix2raster(z,x,y,layer,proj="+proj=longlat")
```

# **Arguments**

z	matrix or array to be converted.
x	optional x-coordinates giving the horizontal range of the raster layer, its size does not need to coincide with ncol(z)!
У	optional y-coordinates giving the verical range of the raster layer, its size does not need to coincide with $nrow(z)$ !
layer	layer to be selected (only valid if z is an array).
proj	optional argument, setting the coordinate reference system (CRS) of a Raster* object (default is +proi=longlat).

# Author(s)

Robert K. Bauer

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### **Examples**

```
## Example 1: convert a matrix
m \leftarrow matrix(3,2,2)
matrix2raster(m)
## Example 2: convert an array
a <- array(3, dim=c(2,2,2))
matrix2raster(a)
matrix2raster(a,layer=1)
## Example 3: convert '.nc'-file to raster-object manually
owd <- getwd()</pre>
path <- system.file("test_files", package="oceanmap")</pre>
ncfile <- Sys.glob(paste0(path,'/herring*.nc')) # load sample-'.nc'-files</pre>
library('ncdf4')
library('raster')
nc <- nc_open(ncfile) # open netcdf file</pre>
z <- ncvar_get(nc, 'Conc')[,,1]</pre>
lon <- as.vector(ncvar_get(nc,'lon')) # fillvalues are automatically replaced by NA</pre>
lat <- as.vector(ncvar_get(nc,'lat')) # fillvalues are automatically replaced by NA
matrix2raster(z,x=lon,y=lat)
## Example 4: convert '.nc'-file to raster-object using nc2raster
nc2raster(ncfile, varname='Conc', layer=1:4)
```

name\_join

create ''.gz'-filenames from a list or dataframe

# Description

```
creates filenames based on a list or dataframe with the (header)-names:
area source parameter resolution timestep date1 date2 option
by aligning the defined filetype:
e.g. area_source_parameter_resolution_timestep_date1_date2.option.filetype
```

### Usage

```
name_join(parts,filetype='gz')
```

### **Arguments**

```
parts a list or dataframe with the parts: area , the region keyword source , the data source
```

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```
param , the parameter saved in the '.gz'-file. Can only be one value!
resolution , the spatial resolution
timestep , the temporal resolution
date1 & date2 , the temporal resolution (the time interval covered).
option a character string holding supplmentary information of '.gz'-file treatment
filetype character string inidicating the filtype to be checked. ('.gz' by default)
```

#### Author(s)

Herve Demarq, translated from IDL by Robert K. Bauer

#### See Also

See check\_gzfiles to return summary of available '.gz'-files and name\_split to split '.gz'-filenames

# **Examples**

```
## Example: read and plot '.gz'-file
path <- system.file("test_files", package="oceanmap")
gz.files <- Sys.glob(paste0(path,'/*.gz')) # load sample-'.gz'-files
check_gzfiles(folder=path) # return file summary-table

# return summary of availble '.gz'-files
# suffix-column corresponds to option column of the name_join-call
# addition n-column returns the number of available files per filetype
check_gzfiles(gz.files)

## Example: split and rejoin '.gz'-filenames
name_split(gz.files) # return summary-table per file
name_join(name_split(gz.files))</pre>
```

name\_split

Returns a summary data frame of '.gz' encoded oceanography files by splitting their name

# **Description**

Returns a summary data.frame of '.gz' encoded oceanography files by splitting their name

# Usage

```
name_split(gz.files)
get.gz.info(gz.files)
```

# **Arguments**

gz.files

Optional character vector or search criteria for .gz-encoded oceanography files.

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#### Value

Returns a summary data.frame of '.gz' encoded oceanography files by splitting their name area source parameter resolution timestep date1 date2 option

area region keyword source data source

param the parameter saved in the '.gz'-file. Can only be one value!

resolution the spatial resolution timestep the temporal resolution

date1 & date2 the time interval covered in date format

option a character string holding supplmentary information of '.gz'-file treatment

#### Author(s)

Robert K. Bauer

### See Also

See check\_gzfiles to return summary of available '.gz'-files and name\_join to create '.gz'-filenames from splitted names (name\_split)-calls

```
## Example: read and plot '.gz'-file
path <- system.file("test_files", package="oceanmap")
gz.files <- Sys.glob(paste0(path,'/*.gz')) # load sample-'.gz'-files
check_gzfiles(folder=path) # return file summary-table

# return summary of availble '.gz'-files
# suffix-column corresponds to option column of the name_split-call
# addition n-column returns the number of available files per filetype
check_gzfiles(gz.files)

## Example: split and rejoin '.gz'-filenames
gz.files
name_split(gz.files) # return summary-table per file
name_split() # return summary-table of all gz-file in current folder
name_join(name_split(gz.files))</pre>
```

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nc2raster Convert Raster layer to a matrix or array	
---	--

### **Description**

nc2raster converts a netcdf-file ('.nc'-file) or ncdf4-object to a Raster\* object, setting the time variable as layer name.

# Usage

```
nc2raster(nc, varname, t=layer, layer, verbose=FALSE)
```

# **Arguments**

nc character string indicating the filepath to a netcdf-file ('.nc'-file), or a ncdf4-

object.

varname character string indicating the name of the netcdf-variable to be selected.

layer, t layer/time stemp to select in multi-layer files.

verbose should information about the netcdf file, including the variables and dimensions

it contains, be printed during loading? (default is FALSE)

#### Value

RasterLayer or RasterStack

# Author(s)

Robert K. Bauer

```
path <- system.file("test_files", package="oceanmap")
nfiles <- Sys.glob(paste0(path,'/*.nc'))[1] # load sample-'.nc'-files
nc2raster(nfiles[1],"Conc",layer=1) # RasterLayer
nc2raster(nfiles[1],"Conc",layer=1:4) # RasterStack
library('ncdf4')
nc <- nc_open(nfiles[1])
nc2raster(nc,"Conc",layer=1:4) # RasterStack
###### load & plot sample netcdf-file ('.nc'-file)
### option a) load netcdf-file with ncdf4-package and plot it
library('ncdf4')
ncdf <- nc_open(nfiles[1])
print(ncdf)
v(obj = ncdf, cbpos="r")</pre>
```

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```
### option b) load and plot netcdf-file as RasterStack object
nc <- nc2raster(nfiles[1])
v(nc,cbpos="r") # plot RasterStack object
v(nfiles[1], cbpos="r",replace.na=TRUE) # plot directly netcdf-file
### option c) plot netcdf-file directly
v(nfiles[1], cbpos="r") # plot RasterStack object
###### plot multiple layers:
par(mfrow=c(2,2))
v(nfiles[1], t=1:4, cbpos="r", replace.na=TRUE, subplot = TRUE)</pre>
```

nc2time

reads and converts the time variable of a netcdf-file ('.nc'-file) or ncdf4-object as as.Date-object

### **Description**

reads and converts the time variable of a netcdf-file ('.nc'-file) or ncdf4-object as as.Date-object.

### Usage

```
nc2time(nc,varname)
```

### **Arguments**

nc character string indicating the filepath to a netcdf-file ('.nc'-file), or a ncdf4-

object.

varname character string indicating the name of the time vaiable of the netcdf-file.

# Author(s)

Robert K. Bauer

```
path <- system.file("test_files", package="oceanmap")
nfile <- Sys.glob(paste0(path,'/herring*.nc')) # load sample-'.nc'-files
head(nc2time(nfile))

library('ncdf4')
nc <- nc_open(nfile)
head(nc2time(nc))</pre>
```

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oceanmap

oceanmap - plot tools for 2D oceanographic data

### **Description**

oceanmap is a plotting toolbox for oceanographic data. Visualizing data is a crucial step in analyzing and exploring data. During the last two decades the statistical programming language R has become a major tool for data analyses and visualization across different fields of science. However, creating figures ready for scientific publication can be a tricky and time consuming task.

The oceanmap package provides some helpful functions to facilitate and optimize the visualization of geographic and oceanographic data, such as satellite and bathymetric data sets. Its plotting functions are written in a way that they do not require a large amount of their numerous arguments to be specified but still return nice plots. Its major functions are:

#### **Major functions:**

- plotmap: plots landmask as basis or overlay
- v: plots oceanographic data (fronts, SST, chla, bathymetry, etc.) from raster-objects, ncdf4or gz-files
- set.colorbar: adds a colorbar to current figure, allowing several placement methods
- get.bathy: download bahymetric data at user defined resolution from the NOAA ETOPO1 database
- add.region: generate region definitions to facilitate land mask and colorbar plotting using plotmap and v
- figure & close\_fig: generate and save graphic devices in flexible file formats (jpeg, png, eps, pdf and eps)

### **Getting Started**

Check out some examples of the principle functions, listed above.

#### Author(s)

Robert K. Bauer

parameter\_definitions parameter definitions dataframe

### **Description**

a dataframe containing definitions of parameters to plot or to save by v, readbin and writebin.

### Usage

data(parameter\_definitions)

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# **Format**

data.frame

### Value

a dataframe with the following header, containing definitions of parameters to plot or to save by v, readbin and writebin:

param a b c log name1 unit pal1 minv maxv min max invalid\_data\_dc coast\_dc land\_dc no\_data\_dc

param	character string indicating the keyword of a parameter.	
a,b,c	value for parameter parameter data conversion from/to byte data. (See $param\_convert$ and $param\_unconvert$ )	
log	whether a logarithmic formula should be applied for data conversion (0 for FALSE and 1 for TRUE; See param_convert and param_unconvert).	
name	character string indicating the long name of a parameter.	
unit	character string or bgroup statement indicating the parameter unit.	
pal1	default color map used by v calls on parameter related data.	
minv, maxv	default minimum and maximum z-value used by v calls on parameter related data.	
min, max	minimum and maximum byte-values to be considered when calculating absolute values.	
<pre>invalid_data_dc, coast_dc, land_dc &amp; no_data_dc</pre>		

byte values used to mask invalid data, coast lines, land masses and missing data.

# Author(s)

Robert K. Bauer

### See Also

v

```
## Example
data(parameter_definitions)
head(parameter_definitions)

# selecting sea surface temperature parameter definition
parameter_definitions[parameter_definitions$param == "sst2",]
```

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param\_convert

converts byte data to absolte values or vise versa (param\_unconvert)

### **Description**

converts byte data as stored in '.gz'-files to absolte values (param\_convert) or vise versa (param\_unconvert) using the parameter\_definitions-dataset. param\_convert is used by readbin, param\_unconvert is used by writebin.

#### Usage

```
param_convert(x,param)
param_unconvert(x,param)
```

# **Arguments**

x vector, matrix or raster-object holding byte-data that that should be converted to

absolute values (param\_convert) or vise versa (param\_unconvert).

param Character string indicating parameter of the dataset to be treated. See parameter\_definitions

for available parameters.

# Author(s)

Robert K. Bauer

# See Also

param\_unconvert, readbin

```
library('fields')
path <- system.file("test_files", package="oceanmap")
gz.file <- Sys.glob(paste0(path,'/*.gz'))[1] # load sample-'.gz'-files
param <- name_split(gz.file)$parameter
print(param)

## converted data, according to param information
m <- readbin(gz.file, Raster=FALSE)
image.plot(m)

## byte data ("unconverted") according to param information, as stored in ".gz"files
bin <- param_unconvert(m,param)
image.plot(bin)

## reconverting byte data, according to param information
conv <- param_convert(bin,param)
image.plot(conv)</pre>
```

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plotmap

plots landmask of a defined region

# **Description**

plots the landmask of a region defined by a region-key word, georgraphical coordinates (longitude and latitude), a raster- or extent-object. See add.region to add and save new region definitions. Attention! Unlike add.region, plotmap does not include colorbar placement (see: set.colorbar)

# Usage

### **Arguments**

region, v_area	Character string identifying regions predefined by the region_definitions-dataset, Raster* or Extent object (corresponds to v_area of the v-function). If missing, region is derived from geographical coordinates, denoted by lat and lon. See add.region to define new region definitions and delete.region to delete unproper region definitions.	
lon, xlim	Vector returning longitude coordinates of the area to be plotted.	
lat, ylim	Vector returning latitude coordinates of the area to be plotted.	
add, add_to	whether the a the landmask should be added to an existent figure (default is FALSE) or an existing ggplot object, in case of ggplotmap.	
asp	numeric, giving the aspect y/x-ratio of the y- and x-axes. See plot.window for more details.	
main	title to be plotted	
axes, axeslabels		
	whether axes and axes-labels (longitude and latitude) should be plotted (default is TRUE). axes-labels can be a single value or a vector of size two, representing values for x and y axis, respectively.	
ticklabels	whether tick-labels should be added to the axes (default is TRUE). Can be a single value or a vector.	

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cex.lab font size of axis labels cex.ticks font size of tick labels whether a grid should be plotted (default is TRUE) grid grid.res resolution of the grid, in degrees (default is is derived from the region extent) resolution number that specifies the resolution with which to draw the map. Resolution 0 is the full resolution of the database [default]. Otherwise, just before polylines are plotted they are thinned: roughly speaking, successive points on the polyline that are within resolution device pixels of one another are collapsed to a single point (see the Reference for further details). Thinning is not performed if plot = FALSE or when polygons are drawn (fill = TRUE or database is a list of polygons). bwd width is of the axes bars (default is 1) fill.land whether the a the landmask should be filled by a color (default is TRUE) col.land fill color of the landmask to be plotted (default is grey) background color (ocean) to be plotted (default is NA) col.bg border country border color of the landmask to be plotted (default is black) col.scale color of the map scale to be plotted around the map (default is black) las numeric in  $\{0,1,2,3\}$ ; the style of axis labels **0:** always parallel to the axis [default], 1: always horizontal, 2: always perpendicular to the axis, **3:** always vertical.

### **Details**

plotmap uses the maps and maptools functions to plot the landmask.

# Author(s)

Robert K. Bauer

#### See Also

```
v, regions
```

```
#### Example 1: plot landmask of the Mediterranean Sea
## a) by using longitude and latitude coordinates:
lon <- c(-6, 37)
lat <- c(30, 46)
figure(width=9.75,height=5.28)
plotmap(lon=lon, lat=lat, main="Mediterranean Sea")
plotmap(xlim=lon, ylim=lat, main="Mediterranean Sea")
ggobj <- ggplotmap(xlim=lon, ylim=lat)
ggobj</pre>
```

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```
## b) plot landmask of the Mediterranean Sea by using an extent-object:
# library('raster')
# ext <- extent(lon, lat)</pre>
# plotmap(ext, main="Mediterranean Sea") # extent-object
# ggplotmap(ext)
## c) plot landmask of the Mediterranean Sea by using a raster-object:
# r <- raster(ext)</pre>
# plotmap(r, main="Mediterranean Sea") # raster-object
# ggplotmap(r)
## d) plot landmask of the Mediterranean Sea by using a region label:
# plotmap('med4', main="Mediterranean Sea") # region-label
# regions() ## check preinstalled region label
## e) add landmask to an existing plot:
# plot(3.7008, 43.4079, xlim=lon, ylim=lat)
# plotmap(add=T)
# points(3.7008, 43.4079, pch=19)
# ggplotmap(xlim=lon, ylim=lat)
# library(ggplot2)
# ggobj <- ggplotmap("lion") +</pre>
          geom_point(data=data.frame(x=3.7008, y=43.4079),aes(x,y),size=5,colour="blue")
# ggobj
## ggplotmaply(ggobj)
## f <- ggplotmaply(ggobj)</pre>
## pos <- as.data.frame(list(x=c(5.83, 4.91, 5.67, 5.91, 6.31, 6.37,
##
                                 5.66, 5.54, 5.51, 5.67, 5.89, 5.97),
##
                             y=c(42.89, 42.27, 42.42, 42.33, 42.1, 41.92,
##
                                 41.74, 41.45, 41.32, 41.21, 41.04, 40.96)
                            ))
## library(dplyr)
## library(plotly)
## f %>% add_trace(data = pos,x = ~x, y= ~y,type='scatter',mode='marker',name="new pos")
#### Example 2: subplots and some additional arguments of plotmap()
# par(mfrow=c(2, 1))
# plotmap('medw4', main="Western Mediterranean Sea",col.bg="darkblue")
# plotmap('medw4', main="Western Mediterranean Sea", bwd=3, border='grey', grid=FALSE)
#### Example 3: plotmap() and figure()
# do.save <- FALSE ## open a plotting window</pre>
# figure("Gulf_of_Lions_extended", do.save=do.save, width=5, height=5, type="pdf")
# plotmap("lion",col.bg='darkblue',grid=FALSE)
# close_fig(do.save)
```

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```
## now resize figure manually and get new figure dimensions:
# width <- dev.size()[1]</pre>
# height <- dev.size()[2]</pre>
# do.save <- TRUE ## do NOT open a plotting window, but save figure internally
# figure("Gulf_of_Lions_extended", do.save=do.save, width=width, height=height, type="pdf")
# plotmap("lion",col.bg='darkblue',grid=FALSE)
# close_fig(do.save)
#### Example 4: between hemispheres
# par(mfrow=c(2,1))
# plotmap(lon=c(-180, 180), lat=c(-80, 80), main="map from West to East")
# plotmap(lon=c(0, 360), lat=c(-80, 80), main="map from West to East")
# plotmap(lon=c(-360, 00), lat=c(-80, 80), main="map from West to East") # same as before
#### Example 5: plot bathymetry and topography of the western Mediterranean Sea
#get.bathy("medw4",visualize=T,terrain=T,res=3)
#get.bathy("medw4",visualize=T,terrain=F,res=3,levels=c(200,2000)) # show contours
#get.bathy("lion",visualize=T,terrain=F,res=3,levels=c(200,2000),v_image=F) # show only contours
#### Example 6: testing some additional arguments
# lon <- c(-180,200); lat <- c(-80,90);
# ext <- extent(lon, lat)</pre>
# plotmap(ext, border=NA, bwd=NA, grid=FALSE, col.land = "#9ac0cd",axes=FALSE)
```

raster2matrix

Convert Raster layer to a matrix or array

# Description

raster2matrix converts a raster layer to a matrix or array. Used by readbin and writebin.

# Usage

```
raster2matrix(RasterLayer)
raster2array(RasterLayer)
```

#### **Arguments**

RasterLayer raster layer to be converted.

#### Author(s)

Robert K. Bauer

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### **Examples**

```
library('raster')
path <- system.file("test_files", package="oceanmap")
gz.files <- Sys.glob(paste0(path,'/*.gz')) # load sample-'.gz'-files
check_gzfiles(folder=path) # return file summary-table

raster.file <- readbin(gz.files[1]) # loading gz-file as raster-layer
image(raster.file)

## Example 1: converting single raster layer to matrix
image(as.matrix(raster.file)) # unflipped conversion
m <- raster2matrix(raster.file) # converting raster-layer to matrix
image(m)

## Example 2: converting double raster layer to an array
stack.file <- stack(raster.file,raster.file)
image(as.array(stack.file)[,,1]) # unflipped conversion
a <- raster2array(stack.file) # converting raster-layer to array (works also with raster2matrix)
image(a[,,1])</pre>
```

readbin

Returns '.gz'-file as matrix or raster-object

# **Description**

Returns '.gz'-file as matrix or raster-object.

### Usage

```
readbin(filename, area, Image = F, byte = F, Raster = T)
```

(default is TRUE)

### **Arguments**

filename	Character string indicating search criteria for the '.gz'-file of interest. Only '.gz'-files with valid filenames can be read, consisting of:
	area, source, parameter, resolution, timestep, date1, date2 and option-criteria, separated by an underscore with only option being aligned by a point and ending with '.gz', e.g.:
	$are a\_source\_parameter\_resolution\_timestep\_date 1\_date 2.option.gz.$
	See region_definitions for valid area- and parameter_definitions for valid parameter-values, respecively.
Image	whether the a the '.gz'-file should be plotted immediately using image.plot- function of the fields-package (default is FALSE)
byte	whether the a the data of the '.gz'-file should be returned unconverted as a byte-values (default is FALSE)
Raster	whether the a the data of the '.gz'-file should be returned in a raster-object

regions 35

area

Character string identifying the region that should be extracted. If missing, region is derived from the '.gz'-filename. See region\_definitions for area definitions and use add.region to add new regions.

# Author(s)

Robert K. Bauer

### See Also

writebin, regions, crop, raster2matrix, param\_convert

# **Examples**

```
### Example: read and plot '.gz'-file
path <- system.file("test_files", package="oceanmap")</pre>
check_gzfiles(folder=path) # return file summary-table
gz.files <- Sys.glob(paste0(path,'/*.gz')) # load sample-'.gz'-files</pre>
### all manual:
obj <- readbin(gz.files[2],area='lion')</pre>
obj
ticks <- seq(20,30,5)
data('cmap')
image(obj,zlim=range(ticks),col=cmap$jet)
plotmap('lion',add=TRUE) # add landmask
#set.colorbar(ticks=ticks,cb.title='cb.title',cb.xlab='cb.xlab')
### using v:
## ticks set by adaptive.vals
v(obj,varname="sst2",cb.title='cb.title',cb.xlab='cb.xlab')
## ticks set by parameter definition
v(obj,varname="sst2",cb.title='cb.title',cb.xlab='cb.xlab',adaptive.vals=FALSE)
### extracting subregion:
obj <- readbin(gz.files[2])</pre>
area.extent <- extent(c(5,10,35,40))
subarea <- crop(obj,area.extent)</pre>
# v(subarea)
## getting average value:
mean(subarea[,],na.rm=TRUE)
```

regions

Returns two-row summary table of a specified region.

36 regions

# **Description**

Reorganizes summary information of a specified region from the region\_definitionsset into a two-row dataframe. Region definitions can be added, backed up or restored by add.region or deleted by calling delete.region.

**ATTENTION!** When reinstalling or updating the oceanmap package, previous region definitions are getting lost! It is therefore highly recommanded to take and restore own backups (see: backup and restore).

# Usage

```
regions(label)
```

### **Arguments**

label Character string indicating the name of the region of interest. If missing, list

of available regions in the region\_definitions-dataset will be returned by a error

message.

### Value

a two-row dataframe with the following header, containing the summary information of the region specified:

xlim ylim dim name cbx cby align gradient figdim grid.res

xlim & ylim the spatial extent of the region

dim the number of grid points for both x & y-dimension

name the long name of the region cbx & cby x & y-coordinates for colorbar

align a vector defining the color-gradient of the colorbar (x for horizontal, and y

for vertical), as well as the margin where the colorbar ticks should be plotted,

relative to the colorbar rectangle ('l' left, 'r' right and 'b' for bottom)

figdim the region-specific default plot device size grid.res the default grid resolution in degrees

#### Author(s)

Robert K. Bauer

### See Also

```
v, plotmap
```

```
## Example: return summary table for the Gulf of Lions
data('region_definitions')
region_definitions[region_definitions$label=='lion',] # select raw region data summary
regions('lion') # return formatted summary table
```

region\_definitions 37

region\_definitions region definitions dataframe

#### **Description**

dataset providing spatial extent and color bar placement information by a region-keyword in later related function-calls (see: v, plotmap and regions). Information consists of a region-keyword, -longname, its spatial extent (longitudes and latitudes), grid resolution, as well as default colorbar position and figure size. Region definitions can be added, backed up or restored by add.region or deleted by calling delete.region.

**ATTENTION!** When reinstalling or updating the oceanmap package, previous region definitions are getting lost! It is therefore highly recommanded to take and restore own backups (see: backup and restore).

## Usage

data(region\_definitions)

#### **Format**

data.frame

#### Value

dataframe with the following header, containing the summary information of the region specified: label name latn lats lonw lone ncol nrow px cbx1 cbx2 cby1 cby2 gradient oticks figxdim figydim grid.res

label region-keywords

name the long name of the region

latn & lats northern and southern most latitude of the region lonw & lone western and eastern most longitude of the region

ncol, nrow & px default matrix size per region described by the number of columns, rows and

pixels. ATTENTION!! Regions of the same spatial extent but different default (matrix-) resolution may cause errors when reading or writing '.gz'-files and

must therefore be distinguished by different keywords.

cbx1 & cbx2 x-coordinates for colorbar cby1 & cby2 y-coordinates for colorbar

gradient the color-gradient of the colorbar (x for horizontal, and y for vertical)

oticks the margin where the colorbar ticks should be plotted, relative to the colorbar

rectangle ('l' left, 'r' right and 'b' for bottom)

figxdim & figydim

the region-specific default plot device size (width and height in inches)

grid.res the default grid resolution in degrees

38 region\_definitions

#### Author(s)

Robert K. Bauer

#### See Also

See add.region to add new, backup or restore region definitions, and plotmap for basic landmask plots

```
data(region_definitions)
head(region_definitions)
region_definitions$label
# ?region_definitions
# figure(width=15,height=15)
# par(mfrow=c(5,6))
# for(n in region_definitions$label) plotmap(region = n,main=n)
# Mediterranean Sea with a spatial resolution of 4km (e.g. MODIS-Aqua)
region_definitions[region_definitions$label == 'med4',]
# Mediterranean Sea with a spatial resolution of 9km (e.g. dekkar)
region_definitions[region_definitions$label == 'med9',]
# plotting same landmasks by different region-keywords
plotmap('med4')
plotmap('med9')
## Example for selecting wrong area definition when saving files
path <- system.file("test_files", package="oceanmap")</pre>
gz.files <- Sys.glob(paste0(path,'/med4*.gz')) # load sample-med4'.gz'-files</pre>
fname <- name_split(gz.files[1])</pre>
param <- fname$parameter</pre>
gz <- readbin(gz.files[1])</pre>
dim(gz)
v(gz.files[1])
## reset region name
# fname$area <- 'med9'</pre>
# fname <- name_join(fname)</pre>
# writebin(gz,fname,param=param)
# v(fname,folder=".")
# system(paste('rm', fname))
```

set.colorbar 39

set.colorbar	Adds colorbar to an extisting plot device
--------------	---

## **Description**

Adds colorbar to an extisting plot device. If position vectors are not provided, the user will be asked to define the colorbar placement by the mouse cursor.

## Usage

## Arguments

cbx, cby	(set.colorparp-arguments) the horizontal and vertical limits of the colorbar. If missing, the user will be asked for manual colorbar placement.
cbxp, cbyp	(set.colorparp-arguments) the horizontal and vertical limits of the colorbar in percent. If missing, the user will be asked for manual colorbar placement.
cbpos	letter ("b", "l", "t", "r") indicating the position of the colorbar (bottom, left, top, right). Overwrites $cbx$ and $cby$ values.
cbline	distance to default location of the colorbar, starting at 0.
total.reg	(set.colorparp-argument) if colorbar placement is relative to current subplot or entire figure region.
year_bar	whether to plot a colorbar with monthly ticks (by default FALSE)
pal	color map to be plotted (default is 'jet' for direct calls). See cmap for available color maps and parameter_definitions for predefined colormaps of different parameters (for internal function calls, e.g. v))
zlim	(optional) value limits of the color bar. Overwrites ticks if ticks are povided.
ticks	the points at which tick-marks are to be drawn (default is 1:10). Non-finite (infinite, NaN or NA) values are omitted. Gets verwritten by zlim if povided.
labels	character or expression vector of labels to be placed at the tickpoints. (default equals ticks-values.)
gradient	whether to have a horizontal (x) or vertical (y) color gradient.
oticks	the margin where to put the colorbar ticks relative to the colorbar rectangle ('1' left, 'r' right and 'b' for bottom;
cb.title	character string indicating the title of the colorbar (default is set to date information/empty string if date information is missing.)

40 set.colorbar

```
character string indicating the x-axis label of the colorbar.
cb.xlab
font
                   Integer specifying font to use for text. 1=plain [default], 2=bold, 3=italic, 4=bold
                   italic, 5=symbol
cb.xlab.line
                  line of x-axis colorbar label
cex, cex.cb.title, cex.cb.xlab, cex.cb.ticks
                   cex: general font size, used as reference for colorbar labels and title cex.cb.xlab:
                   font size of the x-axis label of the colorbar cex.ch.title: font size of the title of
                   the colorbar
cb.ticks.srt, cb.ticks.length, cb.ticks.ypos, cb.ticks.lwd
                  rotation, length, relative y-position and line width of colorbar ticks
integer
                   (default is FALSE).
                   additional arguments to be passed to text or set.colorpar
. . .
```

#### **Details**

set.colobar adds a colorbar to the current plot device. If colorbar positions are missing (cbx, cby), the user will be asked for manual placement. ticks and tick-labels should correspond to zlim-values of the plot. pal defines the colormap and should equal col of the selected plot.

#### Value

a list of colorbar definition vectors: oticks, gradient, cbx and cby. See function argmuments for more details.

#### Author(s)

Robert K. Bauer

```
## Example 1: plot colorbars manually
par(mar=c(8,8,8,8))
plot(0.5, 0.5, xlim=c(0,1), ylim=c(0,1))
set.colorbar(cbx=c(0, 1), cby=c(-.3, -.4)) # bottom
set.colorbar(cby=c(0, 1), cbx=c(-.4, -.3)) # left
set.colorbar(cbx=c(0, 1), cby=c(1.2, 1.3)) # top
set.colorbar(cby=c(0, 1), cbx=c(1.2, 1.3)) # right
## Example 2: use cbpos
par(mar=c(8,8,8,8))
plot(0.5, 0.5, xlim=c(0,1), ylim=c(0,1))
set.colorbar(cbpos='b') # bottom
set.colorbar(cbpos='l') # left
set.colorbar(cbpos='t') # top
set.colorbar(cbpos='r') # right
## Example 3: interactive placement
\# par(mar=c(8,8,8,8))
```

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```
# plot(0.5,0.5,xlim=c(0,1),ylim=c(0,1))
# cb <- set.colorbar() # interactive
# plot(0.5,0.5,xlim=c(0,1),ylim=c(0,1))
# set.colorbar(cbx=cb$cbx, cby=cb$cby) # reuse stored colorbar positions</pre>
```

SpatialCircle

Creates a circle of radius r around a specified point.

## **Description**

Returns a SpatialLines object, that defines a circle of radius r around a specified point.

## Usage

```
SpatialCircle(x,y,r,n=100,proj4str)
```

## **Arguments**

x, y x and y coordinates of the circle

r radius (of the same scale as plotting window)

n precision indicator of the circle.

proj4str projection string.

#### Author(s)

Robert K. Bauer

#### See Also

get.avg.bathy

```
## Example 1: load & plot bathymetry of the Baltic Sea, defined by longitudes and latidtues
lon <- c(9, 31)
lat <- c(53.5, 66)
# bathy <- get.bathy(lon=lon, lat=lat, main="Baltic Sea", cbpos='r')
plotmap(lon=lon, lat=lat)
spc <- SpatialCircle(x= 20,y = 57.5,r=1)
plot(spc,add=TRUE)
# get.avg.bathy(c(20,57.5), radius = 1, bathy = bathy)
# get.avg(c(20,57.5), radius = 1, unit="km",raster = bathy)</pre>
```

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Plotting spatial data

## **Description**

Plots spatial data (e.g. 2D oceanographic data). Valid input data are objects of class 'Raster' ('RasterLayer', 'RasterStack' or 'RasterBrick'), 'ncdf4' (already loaded netcdf files) or a character strings indicating 'bathy' metric data, 'gz'- or '.nc-files' (netcdf). See also name\_split for further information on '.gz'-file nomenclature.

## Usage

```
## S4 method for signature 'bathy'
v(obj, v_area, lon, lat, resolution=4, keep=F,
 savename.bathy, folder.bathy=".", adaptive.vals=T, cb.title, show.colorbar=T,...)
## S4 method for signature 'nc'
v(obj, varname, t=1, layer=t, adaptive.vals=T, dates,
  cb.xlab=varname, show.colorbar=T ,...)
## S4 method for signature 'ncdf4'
v(obj, varname, t=1, layer=t, adaptive.vals=T, dates,
  cb.xlab=varname, show.colorbar=T, ...)
## S4 method for signature 'RasterLayer'
v(obj, varname, t=1, layer=t, ...)
## S4 method for signature 'RasterBrick'
v(obj, varname, t=1, layer=t, ...)
## S4 method for signature 'RasterStack'
v(obj, varname, t=1, layer=t, ...)
## S4 method for signature 'gz'
v(obj, v_area, adaptive.vals=F, show.colorbar=T,...)
```

#### **Arguments**

object of class 'Raster' ('RasterLayer', 'RasterStack' or 'RasterBrick'), 'ncdf4' obj

or a character string indicating, 'bathy'metric data, '.gz'- or '.nc'-files to

v\_area character string identifying the region that should be plotted, or in case of obj ==

'bathy', also a Raster\* or Extent object. If missing, region is derived from the '.gz'-filename. See region\_definitions for area definitions and use add.region

to add new regions.

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adaptive.vals sets minimum and maximum z-value according to the '.gz'-files value range. (ATTENTION! minv and maxv are disregarded if set!). (default is TRUE for non-'.gz'-files. If FALSE or not set, default value from the parameter definitionsdataset will be applied according to the param-value. t, layer layer/time stemp to select in multi-layer files/objects (e.g. ncdf4, RasterStack). dates vector of type 'character' indicating dates per layer, used to define the title of the colorbar. Argument is omitted for '.gz'-files but date-information is derived from the filename. For '.nc'-files or 'ncdf4'-objects, date information is derived from the time-vector. For raster-objects the layer name is applied. varname character string indicating the name of the variable to plot. For '.nc'-files or 'ncdf4'-objects, this name must correspond to a variable name defined in the file/object. Sets also colorbar-title for non-'.gz'-files if cb.xlab is missing. character string indicating the title of the colorbar (default is set to date inforcb.title mation/empty string if date information is missing.) cb.xlab character string indicating the x-axis label of the colorbar and cb.xlab.line its placement line (default is 0). If not defined, it will be set to varname for raster, ncdf4-objects and '.nc'-files or for '.gz'-files to a predefined title in the parameter\_definitions-dataset according to the param-value. lon Vector returning longitude coordinates of the area to be plotted, only valable for obj == 'bathy'. lat Vector returning latitude coordinates of the area to be plotted, only valable for obj == 'bathy'. resolution resolution of the bathymetric grid, in minutes (default is 4), only valable for obj == 'bathv'. whether to write the data downloaded from NOAA into a file (default is FALSE), keep only valable for obj == 'bathy'. savename for the bathymetric data file, if not specified set to type 'bathy\_lonsavename.bathy lat\_res.resolution.dat' or 'bathy\_v\_area\_res.resolution.dat', only valable for obj == 'bathy'. folder.bathy directory where bathymetric data should be saved (default is current working directory), only valable for obj == 'bathy'. show.colorbar weather a colorbar should be plotted for image plots(default is T). additional arguments to be passed: region see v\_area.

minv, maxv minimum and maximum z-value to be plotted. If not set, default value from the parameter\_definitions-dataset will be applied. Argument is overwritten by adaptive.vals and zlim.

replace.na whether missing values should be replaced by minimum values (default is FALSE.)

param character string indicating the parameter name for the dataset treatment. See parameter\_definitions for available parameters. For '.gz'-files, param is derived from the filename. For non-'.gz'-files this value is nonobligatory, but can replace the varname-argument and vise versa. See examples.

- main an overall title for the plot: see title.
- cbpos letter ("b", "l", "t", "r") indicating the position of the colorbar (bottom, left, top, right). Overwrites cbx and cby values.
- cbx the horizontal limits (x1, x2) of the colorbar. If missing and the value can not be reconstructed by the region information (e.g. v\_area, '.gz'-file), the user will be asked for manual colorbar placement.
- cby the vertical limits (y1, y2) of the colorbar. If missing and the value can not be reconstructed by the region information (e.g. v\_area, '.gz'-file), the user will be asked for manual colorbar placement.
- nticks number of tick marks for the colorbar (default is 5).
- pal color map to be plotted (default is the 'jet'-colormap, or in case of '.gz'-files derived from the parameter\_definitions-dataset. See cmap for available color maps and parameter\_definitions for predefined colormaps for different parameters.)
- sidelabels whether an additional y-axis label and title should be added to the plot device (default is FALSE). If TRUE, y-axis label is defined by Ylab, the additional title is derived from the date-information and gives the month information.
- Ylab an additional title for the y axis (default is date information), only used when sidelabels is set TRUE. Default value is year-information.
- axeslabels whether axeslabels should be shown (default is TRUE, set as 'longitude' and 'latitude')
- subplot whether '.gz'-file will be plotted as a sub plot to an existing plot device (default is FALSE; see: par)
- width, height the width and height of the plotting window, in inches. For '.gz'-files, default values are derived from the region-name as indicated by the filename. See region\_definitions for predescribed definitions and use add.region to add new region definitions.
- figdim numeric vector indicating the width and height of the plot device in inches. For '.gz'-files, default values are derived from the region-name as indicated by the filename. Value is overwritten if both, width and height are provided. See region\_definitions for predescribed definitions and use add.region to add new region definitions.
- xpos integer: initial position of the top left corner of the figure window on the pc-screen, given in pixels. Negative values are from the opposite corner. (default is -1). Disregarded under Mac OS and if Save is set TRUE.
- Save whether the a plot device should be saved automatically as an image file of type fileformat in a folder specified by plotfolder (default is FALSE)
- plotfolder directory where images should be saved (default is current working directory).
- plotname the name of the output file(s). If not set, value will be derived from the provided file information (For '.gz'-files, default plotname is equal to the '.gz'-filename, replacing the '.gz'-fileformat-suffix with the defined image-fileformat.
- fileformat fielformat of image file to be saved (only png and eps are accepted; default is png).

suffix suffix to be added to the image filename, before the filetype specification (e.g. '...suffix.png').

v\_image whether an image-plot should be plotted (default is TRUE)

v\_contour whether contour lines should be plotted (default is FALSE). If levels are specified, v\_contour is set TRUE.

levels numeric vector of levels at which to draw contour lines.

contour.labels a vector giving the labels for the contour lines. By default levels are used as labels.

v\_arrows whether current or wind vectors should be plotted (default is TRUE; Argument is disregarded for non-.gz-files and omitted if non current or wind data-files are provided)

scale\_arrow scale factor needed for current and wind vector plots (default is 1; Argument is disregarded for non-.gz-files and omitted if no current or wind data-files are provided, indicated by the param-argument (valid param-definitions are: 'uz' and 'vz', for current data, 'wu' and 'wz' for wind data))

terrain whether the to keep terrain data (default is FALSE). If set FALSE and visualize is TRUE, grid command in plotmap is disabled!

verbose whether the plot information shall be printed in the R-console (by default TRUE)

... Additional arguments to be passed to plotmap (bwd, fill, col, border, grid, grid.res, axeslabels, ticklabels, cex.lab, cex.ticks).

#### **Details**

v uses the maps and maptools functions to plot the landmask. See clim\_plot for aligned plots of satallite-data climatologies.

## Author(s)

Robert K. Bauer

#### References

Bauer, R. K., Stepputtis, D., Grawe, U., Zimmermann, C., and Hammer, C. 2013. Wind-induced variability in coastal larval retention areas: a case study on Western Baltic spring-spawning herring. Fisheries Oceanography, 22: 388-399.

#### See Also

```
clim_plot, readbin, name_split, regions, plotmap, v
```

```
dat <- raster::crop(dat,extent(c(0,10,40,44))) ## crop data, xlim/ylim not yet implemented in v()</pre>
print(dat)
v(dat, main="Raster-object", cbpos='r')
## Example 2: load & plot sample netcdf-file ('.nc'-file)
nfiles <- Sys.glob(paste0(path,'/*.nc')) # load list of sample-'.nc'-files
head(nfiles)
### option a) load netcdf-file with ncdf4-package and plot it
library('ncdf4')
ncdf <- nc_open(nfiles[1])</pre>
# print(ncdf)
# v(obj = ncdf, cbpos="r")
### option b) load and plot netcdf-file as RasterStack object
# nc <- nc2raster(nfiles[1])</pre>
# v(nc,cbpos="r") # plot RasterStack object
### option c) plot netcdf-file directly
# v(nfiles[1], cbpos="r")
# v(nfiles[1], cbpos="r", replace.na=TRUE)
###### plot multiple layers:
# par(mfrow=c(2,2))
# v(nfiles[1], t=1:4, cbpos="r", replace.na=TRUE, subplot = TRUE)
# ## Example 2: load & plot bathymetry data from the NOAA-ETOPO1 database
# par(mfrow=c(2,1))
# bathy <- get.bathy("medw4", terrain=T, res=3, keep=T, visualize=T, subplot = TRUE, grid=F)</pre>
# # load('bathy_medw4_res.3.dat',verbose = T); bathy <- h</pre>
# v(bathy, param="bathy", subplot = TRUE, terrain=F, levels=c(200,2000)) # show contours
# ## b) only contour lines:
# par(mfrow=c(1,2))
# h <- get.bathy("lion",terrain=F,res=3, visualize=T,</pre>
                 v_{image} = FALSE, levels=c(200,2000))
# ## use v-function for same plot but on subregion:
\# v(h,v_area = "survey", param="bathy",
# v_image = FALSE, levels=c(200,2000))
## Example 3: plot sample-'.gz'-file
gz.files <- Sys.glob(paste0(path,'/*.gz'))</pre>
# v(gz.files[2]) ## plot content of gz-file
## Example 4: load sample-'.gz'-file manually as Raster-object and plot it
obj <- readbin(gz.files[2],area='lion')</pre>
# par(mfrow=c(1,2))
# v(obj,param="sst",subplot = TRUE)
```

```
# v(obj,param="Temp",subplot = TRUE) ## note unset "pal" (colormap) for unkown "param"-values!
## Example 5: available color maps
data('cmap') # load color maps data
names(cmap) # list available color maps
gz.files <- Sys.glob(paste0(path,'/*.gz'))</pre>
# figure(width=15,height=15)
# par(mfrow=c(4,5))
# for(n in names(cmap)) v(gz.files[2], v_area='lion', subplot=TRUE,
                          pal=n, adaptive.vals=TRUE, main=n)
## define new color maps from blue to red to white:
n <- colorRampPalette(c('blue','red','white'))(100)</pre>
# v(gz.files[2], v_area='lion', subplot=TRUE,
# pal=n, adaptive.vals=TRUE, main="own colormap")
## Example 6: available parameters
data(parameter_definitions)
names(parameter_definitions)
# ?parameter_definitions
# figure(width=12, height=6.2)
\# par(mfrow=c(2,3))
# v('*sst2*707*',v_area="medw4",main="sst", folder=path, subplot=TRUE)
# v('*chla*531*',v_area="medw4",main="chla", folder=path, subplot=TRUE)
 \begin{tabular}{ll} \# \ v('*chlagrad*',v_area="medw4",main="chlagrad",folder=path, \ subplot=TRUE) \end{tabular}
# v('*p100*',v_area="medw4",main="p100 (oceanic fronts)",folder=path, subplot=TRUE)
# v('*sla*',v_area="medw4",main="sla",folder=path, subplot=TRUE)
# h <- get.bathy("medw4",visualize=TRUE,terrain=F,res=4, subplot=TRUE,main="bathy")
############## advanced example section:
## Example I: plot bathymetry using a v_area-keyword
## requires server connection!
# par(mfrow=c(2,1))
# v("bathy","lion",res=4, keep=TRUE,border='grey',
   main='Gulf of Lions bathymetry',cb.title="resolution 4 min")
# v("bathy","lion",res=1, keep=TRUE,border='grey',
   cb.title="resolution 1 min") # can take some time depending on server connection!
## Example II: plot bathymetry of the Baltic Sea defined by longitude and latidtue coordinates
## requires server connection!
lon <- c(9, 31)
lat <- c(53.5, 66)
#v("bathy",lon=lon,lat=lat,main="Baltic Sea")
```

## Example III: plot landmask of the Baltic Sea defined by an extent- or raster-object

```
## requires server connection!
library('raster')
ext <- extent(lon,lat)</pre>
# v("bathy",ext,main="Baltic Sea",res=4,levels=200) # extent-object
## Example IV: plot '.gz'-files, following default plot-procedure
check_gzfiles(folder=path) # return file summary-table
gz.files <- Sys.glob(paste0(path,'/*.gz')) # load sample-'.gz'-files</pre>
# v(gz.files[1:4])
# v(gz.files[4],bwd=2)
## Example V: plot climatologies from '.gz'-files
              (ATTENTION: not working for non-'gz'-files, requiring ImageMagick)
# clim_plot('*1s*.gz' ,folder = path,bwd=0.7,adaptive.vals=TRUE,plotname="seasonal_climatology.png")
## Example VI: plot subregion of gz-files as subplots
# graphics.off()
# par(mfrow=c(2,1))
# v(gz.files[1:2],v_area='lion') # run ?region_definitions to see predefined regions
## Example VII: plot subregion of raster file
# all manual:
obj <- readbin(gz.files[2],area='lion')</pre>
dev.new()
ticks <- seq(20,30,5)
par(mar=c(5,4,5,8))
image(obj,zlim=range(ticks),col=cmap$jet)
plotmap('lion',add=TRUE) # add landmask
# set.colorbar(ticks=ticks,cb.title='cb.title',cb.xlab='cb.xlab')
## using v, reconstructing region information
# obj <- readbin(gz.files[2],area='lion')</pre>
# v(obj,varname="sst2",cb.title='cb.title',cb.xlab='cb.xlab')
# using v for another subregion
ncorse <- crop(obj,extent(6,9,40,42))</pre>
# v(ncorse,grid.res=1)
# v(ncorse,zlim=c(20,30),cbx=c(8.3,8.9),cby=c(40.7,40.8))  # skipping colorbar widget
## Example VIII: Add region by supplying raster-object, colorbar positions and running the widget
#add.region(ncorse,cbx=c(8.3,8.9),cby=c(40.7,40.8))
## Example IX: plot netcdf-files ('.nc'-files)
nfiles <- Sys.glob(paste0(path,'/*.nc')) # load sample-'.nc'-files</pre>
head(nfiles)
## plot herring larval dispersal from Bauer et al. (2013)
```

v-class 49

```
# par(mfrow=c(2,2))
# v(nfiles[1], subplot=TRUE, t=1:4,minv=0, maxv=1000, adaptive.vals=FALSE, replace.na=TRUE)
# par(new=TRUE,mfrow=c(1,1))
# empty.plot(main='herring larval dispersal in the Greifswald lagoon, Germany')
# mtext('see Bauer et al. (2013) as reference')

# plot bathymetric data (obtained from the Leibniz Institute for Baltic Sea Research Warnemuende)
# v(nfiles[2],varname='bathymetry') # following default plot-procedure
# v(nfiles[2],varname='bathymetry',pal='haxbyrev',Log=TRUE, cb.xlab='depth [log m]',levels=50)
```

v-class

v-classes

## **Description**

internal dummy classes used by v.

writebin

Saves geographic data as byte file ('.gz')

## **Description**

Saves geographic data as byte file, in gzip compressed format ('.gz'). ATTENTION!! Only 2D (one layer) can be stored!

## Usage

```
writebin(satdata,filename,folder,param)
```

## Arguments

satdata one layer-raster-object or matrix holding spatial data.

param character string indicating the parameter name for the dataset treatment. See

parameter\_definitions for available parameters.

filename character string naming the '.gz'-file to be created.

folder character string indicating the target directory.

## Author(s)

Robert K. Bauer

50 writebin

## See Also

readbin, regions, crop, raster2matrix, param\_unconvert

```
## Example for selecting wrong area definition when saving files
path <- system.file("test_files", package="oceanmap")</pre>
gz.files <- Sys.glob(paste0(path,'/*.gz')) # load sample-'.gz'-files</pre>
v(gz.files[1])
fname <- name_split(gz.files[1])</pre>
param <- fname$parameter</pre>
gz <- readbin(gz.files[1])</pre>
dim(gz)
v(gz.files[1])
### reset region name
fname$area <- 'med9'</pre>
fname <- name_join(fname)</pre>
# writebin(gz,fname,folder=path,param=param)
# v(fname)
# system(paste('rm', fname))
### multi layer raster file
gz2 <- stack(gz,gz)</pre>
# writebin(gz2,rep(gz.files[1],2),folder=path,param) # error message since multi layer
# writebin(gz,gz.files[1],folder=path,param) # single layer raster file
# v(gz.files[1])
```

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