# Package 'hydrographr'

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```
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Maintainer Maria Üblacker <maria.ueblacker@igb-berlin.d>
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```

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'download_test_data.R'
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'get_segment_neighbours.R'
'get_tile_id.R'
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'reclass_raster.R'
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'snap_to_network.R'
'snap_to_subc_segment.R'

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crop\_to\_extent

Crop raster to extent

# **Description**

This function crops an input raster layer directly on disk, i.e. the input layer does not need to be loaded into R. The raster .tif is cropped to a polygon border line if a vector layer (cutline source) is provided, otherwise if a bounding box is provided (xmin, ymin, xmax, ymax coordinates or a spatial object from which to extract a bounding box), the raster is cropped to the extent of the bounding box. At least a cutline source (vector\_layer) or a bounding box (bounding\_box) should be provided. The output is always written to disk, and can be optionally loaded into R as a SpatRaster (terra package) object (using read = TRUE).

## Usage

```
crop_to_extent(
  raster_layer,
  vector_layer = NULL,
  bounding_box = NULL,
  out_dir,
  file_name,
  read = TRUE,
  quiet = TRUE
)
```

## **Arguments**

raster\_layer character. Full path to the input raster .tif layer.

vector\_layer character. Full path to a vector layer that is used as a cutline data source (similar

to a mask operation).

bounding\_box numeric vector of the coordinates of the corners of a bounding box (xmin, ymin,

xmax, ymax), SpatRaster, SpatVector, or other spatial object.

out\_dir character. The directory where the output will be stored.

file\_name character. Name of the cropped output raster .tif file.

read logical. If TRUE, the cropped raster .tif layer gets read into R. If FALSE, the

layer is only stored on disk. Default is TRUE.

quiet logical. If FALSE, the standard output will be printed. Default is TRUE.

#### Value

The function returns always a .tif raster file written to disk. Optionally, a SpatRaster (terra object) can be loaded into R with read = TRUE.

# Author(s)

Yusdiel Torres-Cambas

download\_test\_data

## **Examples**

download\_test\_data

Download test data

## Description

Download the test data of the package, which includes all Hydrography90m and species point observation data for a small geographic extent, to test the functions.

The test data is available at https://drive.google.com/file/d/1kYNWXmtVm6X7MZLISOePGpvxB1pk1scD/view?usp=share\_and can be automatically downloaded and unzipped with this function to a desired path.

# Usage

```
download_test_data(download_dir = ".")
```

## **Arguments**

download\_dir character. The directory where the files will be downloaded. Default location is the working directory.

#### **Details**

Downloads the test data for the Hydrography90m dataset

#### Author(s)

Afroditi Grigoropoulou

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#### References

Amatulli, G., Garcia Marquez, J., Sethi, T., Kiesel, J., Grigoropoulou, A., Üblacker, M. M., Shen, L. Q., and Domisch, S.: Hydrography90m: a new high-resolution global hydrographic dataset, Earth Syst. Sci. Data, 14, 4525–4550, https://doi.org/10.5194/essd-14-4525-2022, 2022.")

Amatulli G., Garcia Marquez J., Sethi T., Kiesel J., Grigoropoulou A., Üblacker M., Shen L. & Domisch S. (2022-08-09). Hydrography90m: A new high-resolution global hydrographic dataset. IGB Leibniz-Institute of Freshwater Ecology and Inland Fisheries. dataset. https://doi.org/10.18728/igb-fred-762.1

## **Examples**

```
# Download the test data to the current working directory
download_test_data()

# Download the data to a specific (existing) directory
download_test_data("path/to/your/directory")
```

download\_tiles

Download files of the Hydrography90m dataset

# Description

The function downloads files of the Hydrography90m dataset, available at https://public.igb-berlin.de/index.php/s/agciopgzX. The files will be stored in the folder architecture of the above domain. Multiple regular tile or regional unit files can be requested in a single call of the function. The tile or regional unit IDs can be obtained using the functions "get\_tile\_id" and "get\_regional\_unit\_id" respectively.

#### Usage

```
download_tiles(
  variable,
  file_format = "tif",
  tile_id = NULL,
  reg_unit_id = NULL,
  global = FALSE,
  download_dir = "."
)
```

## **Arguments**

variable character vector of variable names.

file\_format character. Format of the requested file ("tif" or "gpkg").

tile\_id character vector. The IDs of the requested tiles.

reg\_unit\_id character vector. The IDs of the requested regional units.

global logical. If TRUE, the global extent file is downloaded. Default is FALSE.

download\_dir character. The directory where the files will be downloaded. Default is the working directory.

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## Author(s)

Afroditi Grigoropoulou

## References

Amatulli G., Garcia Marquez J., Sethi T., Kiesel J., Grigoropoulou A., Üblacker M., Shen L. & Domisch S. (2022-08-09) Hydrography90m: A new high-resolution global hydrographic dataset. IGB Leibniz-Institute of Freshwater Ecology and Inland Fisheries. dataset. https://doi.org/10.18728/igb-fred-762.1

## **Examples**

extract\_from\_gpkg

Extract values from the stream order .gpkg files.

## **Description**

The function reads the attribute table of the stream network GeoPackage file (.gpkg) stored on disk and extracts the data for one or more (or all) input sub-catchment (i.e. stream segment) IDs. The output is a data.table, and only the output is loaded into R.

## Usage

```
extract_from_gpkg(
  data_dir,
  subc_id,
  subc_layer,
  var_layer,
  out_dir = NULL,
  file_name = NULL,
  n_cores = NULL,
  quiet = TRUE
)
```

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## **Arguments**

data_dir	character. Path to the directory containing all input data.
subc_id	a numeric vector of sub-catchment IDs or "all". If "all", the attribute table is extracted for all the stream segments of the input .gpkg layer. The stream segment IDs are the same as the sub-catchment IDs. A vector of the sub-catchment IDs can be acquired from the extract_ids() function, by sub-setting the resulting data.frame.
subc_layer	character. Full path to the sub-catchment ID .tif layer
var_layer	character vector of .gpkg files on disk, e.g. "order_vect_point_h18v04.gpkg".
out_dir	character. The directory where the output will be stored. If the out_dir is specified, the attribute tables will be stored as .csv files in this location, named after their input variable vector files (e.g. "/path/to/stats_order_vect_point_h18v04.csv"). If NULL, the output is only loaded in R and not stored on disk.
file_name	character. Name of the .csv file where the output table will be stored. out_dir should also be specified for this purpose.
n_cores	numeric. Number of cores used for parallelization, in case multiple .gpkg files are provided to var_layer. If NULL, available cores - 1 will be used.
quiet	logical. If FALSE, the standard output will be printed. Default is TRUE.

# Author(s)

Afroditi Grigoropoulou, Jaime Garcia Marquez, Maria M. Üblacker

# References

```
https://grass.osgeo.org/grass82/manuals/v.in.ogr.html
```

```
# Download test data into temporary R folder
# or define a different directory
my_directory <- tempdir()</pre>
download_test_data(my_directory)
# Define path to the directory containing all input data
test_data <- paste0(my_directory, "/hydrography90m_test_data")</pre>
# Define sub-catchment ID layer
subc_raster <- paste0(my_directory, "/hydrography90m_test_data",</pre>
                  "/subcatchment_1264942.tif")
# Extract the attribute table of the file order_vect_59.gpkg for all the
# sub-catchment IDs of the subcatchment_1264942.tif raster layer
attribute_table <- extract_from_gpkg(data_dir = test_data,</pre>
                                      subc_id = "all",
                                      subc_layer = subc_raster,
                                      var_layer = "order_vect_59.gpkg",
                                      n\_cores = 1)
```

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```
# Show the output table
attribute_table
```

extract\_ids

Extract sub-catchment and/or basin IDs

# **Description**

Extracts the ID value of the basin and/or sub-catchment raster layer at a given point location.

# Usage

```
extract_ids(
  data,
  lon,
  lat,
  id = NULL,
  basin_layer = NULL,
  subc_layer = NULL,
  quiet = TRUE
)
```

# **Arguments**

data	a data.frame or data.table with lat/lon coordinates in WGS84.
lon	character. The name of the column with the longitude coordinates.
lat	character. The name of the column with the latitude coordinates.
id	character. The name of a column containing unique IDs for each row of "data" (e.g., occurrence or site IDs).
basin_layer	character. Full path to the basin ID .tif layer.
subc_layer	character. Full path to the sub-catchment ID .tif layer.
quiet	logical. If FALSE, the standard output will be printed. Default is TRUE.

## **Details**

For the extraction of a value at a given point location from the basin and/or sub-catchment raster layer of the Hydrography90m dataset, the GDAL function 'gdallocationinfo' is used. The point locations have to be defined by coordinates of the WGS84 reference system. The function can also be used to extract any value from a given raster layer with a WGS84 projection, such as e.g. environmental information that is stored in the input raster file.

# Note

Duplicated rows will be removed.

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#### Author(s)

Afroditi Grigoropoulou, Maria Üblacker

#### References

https://gdal.org/programs/gdallocationinfo.html

## **Examples**

```
# Download test data into the temporary R folder
# or define a different directory
my_directory <- tempdir()</pre>
download_test_data(my_directory)
# Load occurrence data
species_occurrence <- read.table(paste0(my_directory,</pre>
                                          "/hydrography90m_test_data",
                                          "/spdata_1264942.txt"),
                                  header = TRUE)
# Define full path to the basin and sub-catchments raster layer
basin_raster <- paste0(my_directory,</pre>
                      "/hydrography90m_test_data/basin_1264942.tif")
subc_raster <- paste0(my_directory,</pre>
                     "/hydrography90m_test_data/basin_1264942.tif")
# Extract basin and sub-catchment IDs from the Hydrography90m layers
hydrography90m_ids <- extract_ids(data = species_occurrence,</pre>
                                   lon = "longitude",
                                   lat = "latitude",
                                    id = "occurrence_id",
                                   subc_layer = subc_raster,
                                   basin_layer = basin_raster)
# Show the output table
hydrography90m_ids
```

extract\_zonal\_stat

Calculate zonal statistics

# **Description**

Calculate zonal statistics based on one or more environmental variable raster .tif layers. This function can be used to aggregate data across a set (or all) sub-catchments. The sub-catchment raster (.tif) input file is stored on disk. The output is a data.table which is loaded into R.

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# Usage

```
extract_zonal_stat(
  data_dir,
  subc_id,
  subc_layer,
  var_layer,
  out_dir = NULL,
  file_name = NULL,
  n_cores = NULL,
  quiet = TRUE
)
```

# Arguments

data_dir	character. Path to the directory containing all input data.
subc_id	Vector of sub-catchment IDs or "all". If "all", zonal statistics are calculated for all sub-catchments of the given sub-catchment raster layer. A vector of the sub-catchment IDs can be acquired from the extract_ids() function, and by subsetting the resulting data.frame.
subc_layer	character. Full path to the sub-catchment ID .tif layer.
var_layer	character vector of variable raster layers on disk, e.g. "slope_grad_dw_cel_h00v00.tif". Variable names should remain intact in file names, even after file processing, i.e., slope_grad_dw_cel should appear in the file name. The files should be cropped to the extent of the sub-catchment layer to speed up the computation.
out_dir	character. The directory where the output will be stored. If the out_dir and file_name are specified, the output table will be stored as a .csv file in this location. If they are NULL, the output is only loaded in R and not stored on disk.
file_name	character. Name of the .csv file where the output table will be stored. out_dir should also be specified for this purpose.
n_cores	numeric. Number of cores used for parallelization, in case multiple .tif files are provided to var_layer.
quiet	logical. If FALSE, the standard output will be printed. Default is TRUE.

# Author(s)

Afroditi Grigoropoulou, Jaime Garcia Marquez, Maria M. Üblacker

# References

```
https://grass.osgeo.org/grass82/manuals/r.univar.html
```

# See Also

report\_no\_data to check the defined NoData value. set\_no\_data to define a NoData value.

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## **Examples**

```
# Download test data into the temporary R folder
# or define a different directory
my_directory <- tempdir()</pre>
download_test_data(my_directory)
# Define full path to the sub-catchment ID .tif layer
subc_raster <- paste0(my_directory, "/hydrography90m_test_data",</pre>
                        "/subcatchment_1264942.tif")
# Define the directory where the output will be stored
output_folder <- paste0(my_directory, "/hydrography90m_test_data/output")</pre>
# Create output folder if it doesn't exist
if(!dir.exists(output_folder)) dir.create(output_folder)
# Calculate the zonal statistics for all sub-catchments for two variables
stat <- extract_zonal_stat(data_dir = paste0(my_directory,</pre>
                                               "/hydrography90m_test_data"),
                            subc_id = c(513837216, 513841103,
                                        513850467, 513868394,
                                        513870312),
                            subc_layer = subc_raster,
                            var_layer = c("spi_1264942.tif",
                                           "sti_1264942.tif"),
                            out_dir = output_folder,
                            file_name = "zonal_statistics.csv",
                            n_{cores} = 2)
# Show output table
stat
```

get\_catchment\_graph

Get catchment from graph

## **Description**

Subset the network graph by extracting the upstream, downstream or entire catchment, for one or multiple stream segments. The function will return either one or more data.tables or graph objects for each input stream segment. Note that the stream segment and sub-catchment IDs are identical, and for consistency, we use the term "subc\_id".

By switching the mode to either "in", "out" or "all", only the upstream, downstream or all connected segments will be returned.

## Usage

```
get_catchment_graph(
   g,
   subc_id = NULL,
   outlet = FALSE,
```

```
mode = NULL,
as_graph = FALSE,
n_cores = 1,
max_size = 1500
)
```

#### **Arguments**

g igraph object. A directed graph.

subc\_id numeric vector of a single or multiple IDs, e.g (c(ID1, ID2, ID3, ...). The sub-

catchment (equivalent to stream segment) IDs for which to delineate the upstream drainage area. If empty, then outlets will be used as sub-catchment IDs (with outlet = TRUE). Note that you can browse the entire network online at https://geo.igb-berlin.de/maps/351/view and to left hand side, select the "Stream

segment ID" layer and click on the map to get the ID. Optional.

outlet logical. If TRUE, the outlets of the given network graph will be used as addi-

tional input subc\_ids. Outlets will be identified internally as those stream segments that do not have any downstream connected segment. Default is FALSE.

mode character. One of "in", "out" or "all". "in" returns the upstream catchment, "out"

returns the downstream catchment (all catchments that are reachable from the

given input segment), and "all" returns both.

as\_graph logical. If TRUE, the output will be a new graph or a list of new graphs with

the original attributes. If FALSE, the output will be a new data.table or a list of

data.tables. List objects are named after the subc\_ids. Default is FALSE.

n\_cores numeric. Number of cores used for parallelization in the case of multiple stream

segments / outlets. Default is 1. Currently, the parallelization process requires copying the data to each core. In case the graph is very large, and many segments are used as an input, setting n\_cores to a higher value can speed up the computation. This comes however at the cost of possible RAM limitations and even slower processing since the large data will be copied to each core. Hence

consider testing with  $n_{cores} = 1$  first. Optional.

max\_size numeric. Specifies the maximum size of the data passed to the parallel back-end

in MB. Default is 1500 (1.5 GB). Consider a higher value for large study areas

(more than one  $20^{\circ}x20^{\circ}$  tile). Optional.

#### Value

A graph or data.table that reports all subc\_ids. In case of multiple input segments, the results are stored in a list.

#### Author(s)

Sami Domisch

## References

Csardi G, Nepusz T: The igraph software package for complex network research, InterJournal, Complex Systems 1695. 2006. https://igraph.org

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## **Examples**

```
# Download test data into the temporary R folder
# or define a different directory
my_directory <- tempdir()</pre>
download_test_data(my_directory)
# Load stream network as a graph
my_graph <- read_geopackage(gpkg = paste0(my_directory,</pre>
                                           "/hydrography90m_test_data",
                                           "/order_vect_59.gpkg"),
                            import_as = "graph")
# Pick a random subc_id
subc_id = "513855877"
# Get the upstream catchment as a graph
g_up <- get_catchment_graph(g = my_graph, subc_id = subc_id, mode = "in",</pre>
                             outlet = FALSE, as_graph = TRUE, n_cores = 1)
# Get the downstream segments as a data.table,
g_down <- get_catchment_graph(g = my_graph, subc_id = subc_id, mode = "out",</pre>
                               outlet = FALSE, as_graph = FALSE, n_cores = 1)
# Get the catchments of all outlets in the study area as a graph
g_all <- get_catchment_graph(g = my_graph, mode = "in", outlet = TRUE,</pre>
                              as_graph = TRUE, n_cores = 1)
```

get\_distance

Calculate euclidean or along the network distance between points

## **Description**

Calculate euclidean or along the network distance between points. To calculate the distance along the network, point coordinates need to be snapped to the stream network using the function snap\_to\_network or snap\_to\_subc\_segment.

# Usage

```
get_distance(
  data,
  lon,
  lat,
  id,
  basin_id = NULL,
  basin_layer = NULL,
  stream_layer = NULL,
  distance = "both",
  n_cores = 1,
  quiet = TRUE
)
```

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#### **Arguments**

data	a data.frame or data.table with lat/lon coordinates in WGS84.
lon	character. The name of the column with the longitude coordinates.
lat	character. The name of the column with the latitude coordinates.
id	character. The name of a column containing unique IDs for each row of "data" (e.g., occurrence or site IDs).
basin_id	character. The name of the column with the basin IDs. If NULL and distance is set to 'network' or 'both', the basin IDs will be extracted automatically. Default is NULL.
basin_layer	character. Full path to the basin ID .tif layer. Needs to be defined to calculate the distance along the network.
stream_layer	character. Full path of the stream network .gpkg file. Needs to be defined to calculate the distance along the network.
distance	character. One of "euclidean", "network", or "both". If "euclidean", the euclidean distances between all pairs of points are calculated. If "network", the shortest path along the network between all pairs of points is calculated. (see "Details" for more information). If method is set to "both", both distance measures are calculated. Distances are given in meters. Default is "both".
n_cores	numeric. Number of cores used for parallelization. Default is 1.
quiet	logical. If FALSE, the standard output will be printed. Default is TRUE.

#### **Details**

To calculate the euclidian distance between all pair of points the function uses the v.distance command of GRASS GIS, which has been set up to produce a square matrix of distances. The calculation of distances along the stream network has been implemented with the command v.net.allpairs of GRASS GIS. The along the network distance calculation is done for all pair points located within the same basin. If the points are located in different basins the function can be run in parallel (i.e., each core for the distance calculations of all points within one basin). The distance between points located in different basins is zero because they are not connected through the network.

## Value

If distance='euclidean', a distance matrix, in meters, of the euclidean distances between all the pairs of points (object of class data.frame). If distance='network', a data.frame with three columns: from\_id, to\_id, dist. The 'dist' column includes the distance, in meters, of the shortest path along the network from the node from\_id to the node to\_id. If distance='both', a list containing both objects is returned.

#### Author(s)

Afroditi Grigoropoulou, Maria M. Üblacker, Jaime Garcia Marquez

#### References

https://grass.osgeo.org/grass82/manuals/v.net.allpairs.html https://grass.osgeo.org/grass82/manuals/v.distance.html

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#### See Also

snap\_to\_network to snap the data points to the next stream segment within a given radius and/or a given flow accumulation threshold value. snap\_to\_subc\_segment to snap the data points to the next stream segment of the sub-catchment the data point is located. extract\_ids to extract basin and sub-catchment IDs.

```
# Download test data into the temporary R folder
# or define a different directory
my_directory <- tempdir()</pre>
download_test_data(my_directory)
# Load occurrence data
species_occurrence <- read.table(paste0(my_directory,</pre>
                             "/hydrography90m_test_data/spdata_1264942.txt"),
                               header = TRUE)
basin_rast <- paste0(my_directory,</pre>
                      "/hydrography90m_test_data/basin_1264942.tif")
# Define full path to the sub-catchment raster layer
subc_rast <- paste0(my_directory,</pre>
                     "/hydrography90m_test_data/subcatchment_1264942.tif")
# Define full path to the vector file of the stream network
stream_vect <- paste0(my_directory,</pre>
                       "/hydrography90m_test_data/order_vect_59.gpkg")
# Automatically extract the basin and sub-catchment IDs and
# snap the data points to the stream segment
snapped_coordinates <- snap_to_subc_segment(data = species_occurrence,</pre>
                                              lon = "longitude",
                                              lat = "latitude",
                                              id = "occurrence_id",
                                              basin_layer = basin_rast,
                                              subc_layer = subc_rast,
                                              stream_layer = stream_vect,
                                              n_{cores} = 2)
# Show head of output table
head(snapped_coordinates)
# Get the euclidean distance and the distance along the network between all
# pairs of points
distance_table <- get_distance(data = snapped_coordinates,</pre>
                                lon = "lon_snap",
                                lat = "lat_snap",
                                id = "occurrence_id",
                                basin_id = "basin_id",
                                basin_layer = basin_rast,
                                stream_layer = stream_vect,
                                distance = "network")
```

get\_regional\_unit\_id

```
# Show table
distance_table
```

```
get_regional_unit_id Get Hydrography90m regional unit IDs
```

# Description

Identifies the IDs of the regional units within the Hydrography90m data in which the input points are located. The IDs are required to then download the data using download\_tiles(). Input is a point data frame.

# Usage

```
get_regional_unit_id(data, lon, lat, quiet = TRUE)
```

## **Arguments**

a data.frame or data.table with lat/lon coordinates in WGS84.

lon character. The name of the column with the longitude coordinates.

character. The name of the column with the latitude coordinates.

## Author(s)

Afroditi Grigoropoulou

# References

```
https://gdal.org/programs/gdallocationinfo.html
```

```
get_segment_neighbours
```

Get stream segment neighbours

# **Description**

For each segment, reports those upstream, downstream, or up-and downstream segments that are connected to one or multiple input segments within a specified neighbour order, with the option to summarize attributes across these segments. Note that the stream segment and sub-catchment IDs are identical, and for consistency, we use the term "subc\_id".

This function can also be used to create the connectivity table for Marxan by using var\_layer="length" and attach\_only=TRUE. The resulting table reports the connectivity from each segment, along with the stream length for all connected segments.

## Usage

```
get_segment_neighbours(
   g,
   subc_id = NULL,
   var_layer = NULL,
   stat = NULL,
   attach_only = FALSE,
   order = 5,
   mode = "in",
   n_cores = 1,
   max_size = 1500
)
```

## **Arguments**

g	igraph object. A directed graph.
subc_id	numeric vector of the input sub-catchment IDs (=stream segment IDs) for which to search the connected segments.
var_layer	character vector. One or more attributes (variable layers) of the input graph that should be reported for each output segment_id ("to_stream"). Optional.
stat	one of the functions mean, median, min, max, sd (without quotes). Aggregates (or summarizes) the variables for the neighbourhood of each input segment ("stream", e.g., the average land cover in the next five upstream segments or sub-catchments). Default is NULL.
attach_only	logical. If TRUE, the selected variables will be only attached to each segment without any further aggregation. Default is FALSE.
order	numeric. The neighbouring order as in igraph::ego. Order = 1 would be immediate neighbours of the input sub-catchment IDs, order = 2 would be the order 1 plus the immediate neighbours of those sub-catchment IDs in order 1, and so on.

mode character. One of "in", "out", or "all". "in" returns only upstream neighbouring segments, "out" returns only the downstream segments, and "all" returns both.

n\_cores numeric. Number of cores used for parallelization in the case of multiple stream

segments / outlets. Default is 1. Currently, the parallelization process requires copying the data to each core. In case the graph is very large, and many segments are used as an input, setting n\_cores to a higher value can speed up the computation. This comes however at the cost of possible RAM limitations and even slower processing since the large data will be copied to each core. Hence

consider testing with  $n_{cores} = 1$  first. Optional.

max\_size numeric. Specifies the maximum size of the data passed to the parallel back-end

in MB. Default is 1500 (1.5 GB). Consider a higher value for large study areas

(more than one 20°x20° tile). Optional.

## Author(s)

Sami Domisch

#### References

Csardi G, Nepusz T: The igraph software package for complex network research, InterJournal, Complex Systems 1695. 2006. https://igraph.org

```
# Download test data into the temporary R folder
# or define a different directory
my_directory <- tempdir()</pre>
download_test_data(my_directory)
# Load the stream network as graph
my_graph <- read_geopackage(gpkg= paste0(my_directory,</pre>
                                          "/hydrography90m_test_data",
                                          "/order_vect_59.gpkg"),
                            import_as = "graph")
# Get the upstream segment neighbours in the 5th order
# and report the length and source elevation
# for the neighbours of each input segment
get_segment_neighbours(g = my_graph, subc_id = subc_id,
                       order = 5, mode = "in", n_cores = 1,
                       var_layer = c("length", "source_elev"),
                       attach_only = TRUE)
# Get the downstream segment neighbours in the 5th order
# and calculate the median length and source elevation
# across the neighbours of each input segment
get_segment_neighbours(g = my_graph, subc_id = subc_id,
                       order = 2, mode ="out", n_cores = 1,
                       var_layer = c("length", "source_elev"),
                       stat = median)
```

get\_tile\_id

get\_tile\_id

Get the Hydrography90m regular tile ID

## **Description**

Identifies the ids of the tiles within the Hydrography90m data in which the given points are located. The IDs are required to then download the data using download\_tiles(). Input is a point data frame.

## Usage

```
get_tile_id(data, lon, lat)
```

## **Arguments**

data a data.frame or data.table with lat/lon coordinates in WGS84.

lon character. The name of the column with the longitude coordinates.

lat character. The name of the column with the latitude coordinates.

## Author(s)

Afroditi Grigoropoulou

```
get_upstream_catchment
```

Calculate upstream basin

# Description

Calculates the upstream basin of a given point, considering the point as the outlet.

# Usage

```
get_upstream_catchment(
  data,
  id,
  lon,
  lat,
  direction_layer = NULL,
  out_dir = NULL,
  n_cores = NULL,
  quiet = TRUE
)
```

# **Arguments**

(e.g., occurrence or site IDs).  lon character. The name of the column with the longitude coordinates.  lat character. The name of the column with the latitude coordinates.  direction_layer  character. Full path to raster file with the direction variable.  out_dir Full path to the directory where the output(s) will be stored. To identify the upstream catchment the output file name includes the site id.  n_cores  numeric. Number of cores used for parallelization. If NULL, available cores - 1 will be used.	data	a data.frame or data.table with lat/lon coordinates in WGS84, which have been snapped to the stream network. The snapping can be done using the function 'snap_to_network'.
lat character. The name of the column with the latitude coordinates.  direction_layer  character. Full path to raster file with the direction variable.  out_dir  Full path to the directory where the output(s) will be stored. To identify the upstream catchment the output file name includes the site id.  n_cores  numeric. Number of cores used for parallelization. If NULL, available cores - 1 will be used.	id	character. The name of a column containing unique IDs for each row of "data" (e.g., occurrence or site IDs).
character. Full path to raster file with the direction variable.  out_dir  Full path to the directory where the output(s) will be stored. To identify the upstream catchment the output file name includes the site id.  n_cores  numeric. Number of cores used for parallelization. If NULL, available cores - 1 will be used.	lon	character. The name of the column with the longitude coordinates.
character. Full path to raster file with the direction variable.  out_dir  Full path to the directory where the output(s) will be stored. To identify the upstream catchment the output file name includes the site id.  n_cores  numeric. Number of cores used for parallelization. If NULL, available cores - 1 will be used.		
upstream catchment the output file name includes the site id.  n_cores  numeric. Number of cores used for parallelization. If NULL, available cores - 1 will be used.	un cecion_raye	
will be used.	out_dir	Full path to the directory where the output(s) will be stored. To identify the upstream catchment the output file name includes the site id.
gui et logical If EALSE the standard output will be printed. Default is TDIE	n_cores	•
dufet logical. If FALSE, the standard output will be printed. Default is TROE.	quiet	logical. If FALSE, the standard output will be printed. Default is TRUE.

# Author(s)

Jaime Garcia Marquez, Afroditi Grigoropoulou, Maria M. Üblacker

## References

```
https://grass.osgeo.org/grass82/manuals/r.water.outlet.html https://grass.osgeo.org/grass82/manuals/r.region.html
```

#### See Also

snap\_to\_network to snap the data points to the next stream segment within a given radius and/or a given flow accumulation threshold value. snap\_to\_subc\_segment to snap the data points to the next stream segment within the sub-catchment the point is located. extract\_ids to extract basin and sub-catchment IDs.

```
# Download test data into temporary R folder
# or define a different directory
my_directory <- tempdir()</pre>
download_test_data(my_directory)
# Before running the function get_upstream_catchment(), snap the points to
# to the stream segment. There are multiple ways to snap the points. Here is
# one example:
# Load occurrence data
species_occurence <- read.table(paste0(my_directory,</pre>
                                        "/hydrography90m_test_data",
                                         "/spdata_1264942.txt"),
                               header = TRUE)
# Define full path to the basin and sub-catchments raster layer
basin_raster <- paste0(my_directory,</pre>
                        "/hydrography90m_test_data/basin_1264942.tif")
subc_raster <- paste0(my_directory,</pre>
                       "/hydrography90m_test_data/subcatchment_1264942.tif")
# Define full path to the vector file of the stream network
stream_vector <- paste0(my_directory,</pre>
                         "/hydrography90m_test_data/order_vect_59.gpkg")
# Automatically extract the basin and sub-catchment IDs and
# snap the data points to the stream segment
snapped_coordinates <- snap_to_subc_segment(data = species_occurence,</pre>
                                             lon = "longitude",
                                             lat = "latitude",
                                             id = "occurrence_id",
                                             basin_layer = basin_raster,
                                              subc_layer = subc_raster,
                                             stream_layer = stream_vector,
                                             n_{cores} = 2)
# Define full path to the direction .tif
direction_raster <- paste0(my_directory,</pre>
                            "/hydrography90m_test_data/direction_1264942.tif")
# Define the path for the output file(s)
output_folder <- paste0(my_directory, "/upstream_catchments")</pre>
if(!dir.exists(output_folder)) dir.create(output_folder)
# Get the upstream catchment for each point location
get_upstream_catchment(snapped_coordinates,
```

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```
lon = "lon_snap",
lat = "lat_snap",
id = "occurrence_id",
direction_layer = direction_raster,
out_dir = output_folder,
n_cores = 2)
```

merge\_tiles

Merge raster or vector objects

# **Description**

Merge multiple raster or spatial vector objects from disk to form a new raster or spatial vector object with a larger spatial extent. A directory with at least two raster .tif or spatial vector geopackage files should be provided. Depending on the input, the output is a .tif or a .gpkg file (saved under out\_dir). If read = TRUE, the output is read into R as a SpatRaster (terra package) object in case of .tif files, or as a SpatVector (terra package) object in case of .gpkg files.

# Usage

```
merge_tiles(
   tile_dir,
   tile_names,
   out_dir,
   file_name,
   name = "stream",
   read = FALSE,
   quiet = TRUE
)
```

# **Arguments**

tile_dir	character. The directory containing the raster or spatial vectors tiles, which should be merged.
tile_names	character. The names of the files to be merged, including the file extension (.tif or $.gpkg$ ).
out_dir	character. The directory where the output will be stored.
file_name	character. Name of the merged output file, including the file extension (.tif or $.gpkg$ ).
name	character. The attribute table column name of the stream segment ("stream"), sub-catchment ("ID"), basin ("ID") or outlet ("ID") column which is used for merging GeoPackages. Default is "stream".
read	logical. If TRUE, the merged layer gets read into R. If FALSE, the layer is only stored on disk. Default is FALSE.
quiet	logical. If FALSE, the standard output will be printed. Default is TRUE.

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#### Value

A .tif raster file or .gpkg spatial vector object that is always written to disk, and optionally loaded into R.

#### Author(s)

Thomas Tomiczek, Jaime Garcia Marquez, Afroditi Grigoropoulou

#### References

```
https://gdal.org/programs/gdalbuildvrt.html
https://gdal.org/programs/gdal_translate.html
https://gdal.org/programs/ogrmerge.html#ogrmerge
https://gdal.org/programs/ogr2ogr.html
```

#### **Examples**

```
# Download tiles into the temporary R folder
# or define a different directory
my_directory <- tempdir()</pre>
download_tiles(variable = "basin",
               file_format = "tif",
               tile_id = c("h22v08","h22v10"),
               download_dir = my_directory)
# Define folder containing only the tiles, which should me merged
tiles_folder <- paste0(my_directory, "/r.watershed/basin_tiles20d")</pre>
# Define output folder
output_folder <- paste0(my_directory, "/merged_tiles")</pre>
# Create output folder if it doesn't exist
if(!dir.exists(output_folder)) dir.create(output_folder)
# Merge tiles
merged_tiles <- merge_tiles(tile_dir = tiles_folder,</pre>
                             tile_names = c("h22v08", "h22v10"),
                             out_dir = output_folder,
                             file_name = "basin_merged.tif",
                             read = TRUE)
```

read\_geopackage

Read a GeoPackage file

#### **Description**

Reads an entire, or only a subset of a GeoPackage vector file from disk either as a table (data.table), as a directed graph object (igraph), a spatial dataframe (sf) or a SpatVect object (terra).

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## Usage

```
read_geopackage(
  gpkg,
  import_as = "data.table",
  layer_name = NULL,
  subc_id = NULL,
  name = "stream"
)
```

## Arguments

gpkg character. Full path of the GeoPackage file.

import\_as character. "data.table", "graph", "sf", or "SpatVect". "data.table" imports data

as a data.table. "graph" imports the layer as a directed graph (igraph object). This option is only possible for a network layer (e.g. the stream network). "sf" imports the layer as a spatial data frame (sf object). "SpatVect" imports the layer

as a SpatVector (terra object). Default is "data.table".

layer\_name character. Name of the specific data layer to import from the GeoPackage. A

specific data layer only needs to be defined if the GeoPackage contains multiple layers. To see the available layers the function st\_layers() from the R package

'sf' can be used. Optional. Default is NULL.

subc\_id numeric. Vector of the sub-catchment (or stream segment) IDs in the form of

(c(ID1, ID2, ...) for which the spatial objects or attributes of the GeoPackage

should be imported. Optional. Default is NULL.

name character. The attribute table column name of the stream segment ("stream"),

sub-catchment ("ID"), basin ("ID") or outlet ("ID") column which is used for subsetting the GeoPackage prior importing. Optional. Default is "stream".

#### Author(s)

Sami Domisch, Maria M.Üblacker

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```
"/order_vect_59.gpkg"))
# Read the stream network as a data.table for specific IDs
my_dt <- read_geopackage(gpkg = paste0(my_directory,</pre>
                                         "/hydrography90m_test_data",
                                         "/order_vect_59.gpkg"),
                                         subc_id = c(513833203, 513833594))
# Read the sub-catchments as a SF-object
my_sf <- read_geopackage(gpkg = paste0(my_directory,</pre>
                                         "/hydrography90m_test_data",
                                         "/sub_catchment_59.gpkg"),
                                         import_as = "sf",
                                         layer_name = "sub_catchment")
# Read a subset of sub-catchments as a SF-object
my_sf <- read_geopackage(gpkg = paste0(my_directory,</pre>
                                         "/hydrography90m_test_data",
                                         "/sub_catchment_59.gpkg"),
                                         import_as = "sf",
                                         subc_id = c(513833203, 513833594),
                                         name = "ID")
# Read the basin as SpatVect object
my_sv <- read_geopackage(gpkg = paste0(my_directory,</pre>
                                         "/hydrography90m_test_data",
                                         "/basin_59.gpkg"),
                                         import_as = "SpatVect")
```

reclass\_raster

Reclassify an integer raster layer

#### **Description**

Reclassifies an integer raster .tif layer using the r.reclass function of GRASS GIS. To reclassify the raster layer the present raster values and the new raster values have to be defined.

If the input raster layer has floating point values, you should multiply the input data by some factor (e.g. 1000) to achieve integer values, otherwise the GRASS GIS r.reclass will round the raster values down to the next integer which is not always desired.

## Usage

```
reclass_raster(
  data,
  rast_val,
  new_val,
  raster_layer,
  recl_layer,
```

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```
read = TRUE,
no_data = -9999,
type = "Int32",
compress = "DEFLATE",
quiet = TRUE
)
```

## **Arguments**

data a data.frame or data.table with the present and new raster values.

rast\_val character. The name of the column with the present raster values.

new\_val character. The name of the column with the new raster values.

raster\_layer Full path to the input raster .tif layer.

recl\_layer character. Full path of the output .tif layer of the reclassified raster file.

read logical. If TRUE, then the reclassified raster .tif layer gets read into R as a

SpatRaster (terra object). If FALSE, the layer is only stored on disk. Default is

TRUE.

no\_data numeric. The no\_data value of the new .tif layer. Default is -9999.

type character. Data type; Options are Byte, Int16, UInt16, Int32, UInt32, CInt16,

CInt32. Default is Int32.

compress Compression type: DEFLATE or LZW. Default is DEFLATE.

#### Author(s)

Maria M. Üblacker

#### References

https://grass.osgeo.org/grass82/manuals/r.reclass.html

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report\_no\_data

Report NoData value

# **Description**

This function reports the defined NoData value of a raster layer. The NoData value of a raster layer represents the absence of data. In computations the NoData value can be treated in different ways. Either the NoData value will be reported or the Nodata value will be ignored and a value is computed from the available values of a specified location.

# Usage

```
report_no_data(data_dir, var_layer, n_cores = NULL)
```

# Arguments

data\_dir character. Path to the directory containing all input data.

var\_layer character vector of variable raster layers on disk, e.g. "slope\_grad\_dw\_cel\_h00v00.tif".

n\_cores numeric. Number of cores used for parallelization, in case multiple .tif files are

provided to var\_layer.

## Author(s)

Afroditi Grigoropoulou, Maria M. Üblacker

## References

https://gdal.org/programs/gdalinfo.html

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## **Examples**

set\_no\_data

Set no data value

# **Description**

Change or set the NoData value for a raster layer. The change happens in-place, meaning that the original file is overwritten on disk.

# Usage

```
set_no_data(data_dir, var_layer, no_data, quiet = TRUE)
```

# Arguments

data_dir	character. Path to the directory containing all input data.
var_layer	character vector of variable layers on disk, e.g. c("sti_h16v02.tif", "slope_grad_dw_cel_h00v00.tif"). The original files will be overwritten.
no_data	numeric. The desired NoData value.
quiet	logical. If FALSE, the standard output will be printed. Default is TRUE.

# Author(s)

Afroditi Grigoropoulou, Maria M. Üblacker

# References

```
https://gdal.org/programs/gdal_edit.html
```

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## **Examples**

snap\_to\_network

Snap points to stream segment based on distance or flow accumulation

# **Description**

Snap points to the next stream segment within a defined radius or a minimum flow accumulation.

## Usage

```
snap_to_network(
  data,
  lon,
  lat,
  id,
  stream_layer,
  accu_layer = NULL,
  method = "distance",
  distance = 500,
  accumulation = 0.5,
  quiet = TRUE
)
```

# **Arguments**

data	a data.frame or data.table with lat/lon coordinates in WGS84.
lon	character. The name of the column with the longitude coordinates.
lat	character. The name of the column with the latitude coordinates.
id	character. The name of a column containing unique IDs for each row of "data" (e.g., occurrence or site IDs).
stream_layer	character. Full path of the stream network .tif file
accu_layer	character. Full path of the flow accumulation .tif file. Needed if the point should be snapped to the next stream segment having an accumulation value higher than the flow accumulation threshold (set by 'accumulation'). This prevents points from being snapped to small stream tributaries.

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method character. One of "distance", "accumulation", or "both". Defines if the points

are snapped using the distance or flow accumulation (see "Details" for more information). If method is set to "both" the output will contain the new coordinates

for both calculations. Default is "distance".

distance numeric. Maximum radius in meters. The points will be snapped to the next

stream within this radius. Default is 500.

accumulation numeric. Minimum flow accumulation. Points will be snapped to the next

stream with a flow accumulation equal or higher than the given value. Default

is 0.5.

quiet logical. If FALSE, the standard output will be printed. Default is TRUE.

#### **Details**

The function makes use of the r.stream.snap command available in GRASS GIS to simultaneously a number of points to a stream network. A distance threshold can be specified and points will be snapped to any stream segment within this distance radius. However, to avoid snapping to small tributaries, an accumulation threshold can be used and the snapping occurs on stream segment with equal or higher accumulation threshold and within the given distance radius.

#### Value

Returns a data frame with the snapped coordinates and the sub-catchment ID of the snapped stream segment. If the sub-catchment ID is NA, no stream segment was found within the given distance (method = "distance") or no stream segment wad found within the given distance and a flow accumulation equal or higher than the given threshold (method = "accumulation"). "out-bbox" means that the provided coordinates are not within the extend (bounding box) of the provided stream network layer.

#### Note

Duplicated rows will be removed from the input data.

#### Author(s)

Maria M. Üblacker, Jaime Garcia Marquez

## References

```
https://grass.osgeo.org/grass78/manuals/addons/r.stream.snap.html
```

```
header = TRUE)
# Define full path to stream network and flow accumulation
stream_raster <- paste0(my_directory,</pre>
                      "/hydrography90m_test_data/stream_1264942.tif")
flow_raster <- paste0(my_directory,</pre>
                      "/hydrography90m_test_data/flow_1264942.tif")
# To calculate the new (snapped) coordinates for a radius and a flow
snapped_coordinates <- snap_to_network(data = species_occurrence,</pre>
                                        lon = "longitude",
                                        lat = "latitude",
                                        id = "occurrence_id",
                                        stream_layer = stream_raster,
                                        accu_layer = flow_raster,
                                        method = "both",
                                        distance = 300,
                                        accumulation = 0.8)
# Show head of output table
head(snapped_coordinates)
```

snap\_to\_subc\_segment Snap points to stream segment within the sub-catchment

# **Description**

Snaps data points to the stream segment of the sub-catchment the data point is located.

# Usage

```
snap_to_subc_segment(
  data,
  lon,
  lat,
  id,
  basin_id = NULL,
  subc_id = NULL,
  basin_layer,
  subc_layer,
  stream_layer,
  n_cores = 1,
  quiet = TRUE
)
```

## **Arguments**

data	a data.frame or data.table with lat/lon coordinates in WGS84.
lon	character. The name of the column with the longitude coordinates.
lat	character. The name of the column with the latitude coordinates.
id	character. The name of a column containing unique IDs for each row of "data" (e.g., occurrence or site IDs).
basin_id	character. The name of the column with the basin IDs. If NULL, the basin IDs will be extracted automatically. Default is NULL
subc_id	character. The name of the column with the sub-catchment IDs. If NULL, the sub-catchment IDs will be extracted automatically. Default is NULL.
basin_layer	character. Full path to the basin ID .tif layer.
subc_layer	character. Full path to the sub-catchment ID .tif layer.
stream_layer	character. Full path of the stream network .gpkg file.
n_cores	numeric. Number of cores used for parallelization. Default is 1.
quiet	logical. If FALSE, the standard output will be printed. Default is TRUE.

## **Details**

The function uses the network preparation and maintenance module of GRASS GIS (v.net), to connect a vector lines map (stream network) with a points map (occurrence/sampling points). After masking the stream segment and the sub-catchment where the target point is located, the connect operation snaps the point to the stream segment using a distance threshold. This threshold is automatically calculated as the longest distance between two points within the sub-catchment. In this way the snapping will always take place. This operation creates a new node on the vector line (i.e. stream segment) from which the new snapped coordinates can be extracted.

# Author(s)

Jaime Garcia Marquez, Maria M. Üblacker

## References

```
https://grass.osgeo.org/grass82/manuals/v.net.html
```

#### See Also

snap\_to\_network to snap the data points to the next stream segment within a given radius and/or a
given flow accumulation threshold value. extract\_ids to extract basin and sub-catchment IDs.

```
# Download test data into the temporary R folder
# or define a different directory
my_directory <- tempdir()
download_test_data(my_directory)</pre>
```

```
# Load occurrence data
species_occurence <- read.table(paste0(my_directory,</pre>
                             "/hydrography90m_test_data/spdata_1264942.txt"),
                               header = TRUE)
basin_rast <- paste0(my_directory,</pre>
                      "/hydrography90m_test_data/basin_1264942.tif")
subc_rast <- paste0(my_directory,</pre>
                     "/hydrography90m_test_data/subcatchment_1264942.tif")
# Define full path to the vector file of the stream network
stream_vect <- paste0(my_directory,</pre>
                       "/hydrography90m_test_data/order_vect_59.gpkg")
hydrography90m_ids <- extract_ids(data = species_occurence,</pre>
                                   lon = "longitude",
                                   lat = "latitude",
                                   id = "occurrence_id";
                                   subc_layer = subc_rast,
                                   basin_layer = basin_rast)
# Snap data points to the stream segment of the provided sub-catchment ID
snapped_coordinates <- snap_to_subc_segment(data = hydrography90m_ids,</pre>
                                              lon = "longitude",
                                              lat = "latitude",
                                              id = "occurrence_id"
                                              basin_id = "basin_id",
                                              subc_id = "subcatchment_id",
                                              basin_layer = basin_rast,
                                              subc_layer = subc_rast,
                                              stream_layer = stream_vect,
                                              n_{cores} = 2)
# Show head of output table
head(snapped_coordinates)
# Automatically extract the basin and sub-catchment IDs and
# snap the data points to the stream segment
snapped_coordinates <- snap_to_subc_segment(data = species_occurence,</pre>
                                              lon = "longitude",
                                              lat = "latitude",
                                              id = "occurrence_id",
                                              basin_layer = basin_rast,
                                              subc_layer = subc_rast,
                                              stream_layer = stream_vect,
                                              n_{cores} = 2)
# Show head of output table
head(snapped_coordinates)
```

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