Package ‘riverdist’

July 18, 2017

Type Package
Title River Network Distance Computation and Applications
Version 0.15.0
Date 2017-07-18
Author Matt Tyers [aut, cre]
Maintainer Matt Tyers <matt.tyers@alaska.gov>
Description Reads river network shape files and computes network distances. Also included are a variety of computation and graphical tools designed for fisheries telemetry research, such as minimum home range, kernel density estimation, and clustering analysis using empirical k-functions with a bootstrap envelope. Tools are also provided for editing the river networks, meaning there is no reliance on external software.
License GPL-2
Depends R (>= 2.14.0)
Imports rgdal (>= 0.9-1), sp (>= 1.0-15), methods
Suggests knitr, rmarkdown, testthat
VignetteBuilder knitr
LazyData TRUE
URL https://cran.r-project.org/package=riverdist
BugReports http://github.com/mbtyers/riverdist/issues
RoxygenNote 5.0.1
NeedsCompilation no
Repository CRAN
Date/Publication 2017-07-18 19:21:54 UTC

R topics documented:

riverdist-package ......................................................... 3
abstreams ................................................................. 5
R topics documented:

- abstreams0 ................................................. 6
- addcumuludist .......................................... 6
- addverts .................................................... 7
- buildlookup ............................................. 8
- buildsegroutes ......................................... 9
- calculateconnections .................................. 10
- checkbraided ........................................... 11
- checkbraidedTF ......................................... 12
- cleanup .................................................... 13
- cleanup_verts .......................................... 14
- connectsegs ............................................ 15
- detectroute .............................................. 16
- dissolve .................................................. 17
- fakefish ................................................... 18
- fakefish_density ....................................... 19
- Gulk ....................................................... 20
- highlightseg ........................................... 20
- homerange ............................................... 21
- homerange-class ....................................... 22
- homerangeoverlap ..................................... 23
- isflowconnected ....................................... 24
- Kenai1 .................................................... 25
- Kenai2 .................................................... 25
- Kenai3 .................................................... 26
- kfunc ....................................................... 26
- KilleyW .................................................... 28
- Koyukuk0 .................................................. 28
- Koyukuk1 .................................................. 29
- Koyukuk2 .................................................. 29
- line2network ........................................... 30
- line98 ...................................................... 31
- makeriverdensity ...................................... 31
- mapbyname ............................................... 33
- matbysurveylist ....................................... 34
- mouthdist ............................................... 35
- mouthdistbysurvey .................................... 36
- pdist ....................................................... 38
- pdisttot .................................................... 38
- plot.homerange ......................................... 39
- plot.riverdensity ...................................... 39
- plot.rivernetwork ....................................... 40
- plot homerangeoverlap ................................ 42
- plotmatbysurveylist .................................. 43
- plotriverdensitypoints ................................ 44
- plotseq .................................................... 45
- points2segvert ......................................... 46
- removeduplicates ....................................... 47
- removemicrosegs ....................................... 48
Description

Reads river network shape files and computes network distances. Also included are a variety of
computation and graphical tools designed for fisheries telemetry research, such as minimum home
range, kernel density estimation, and clustering analysis using empirical k-functions with a boot-
strap envelope. Tools are also provided for editing the river networks, meaning there is no reliance
on external software.
The riverdist package provides tools for distance calculation along a river network. The river network is imported from a projected shapefile. Spatial point data may be imported from a shapefile as well, or directly from coordinates.

Some basic formatting of the river shapefile may be necessary. If available, formatting in a geographic information system (GIS) prior to importing into R is recommended (projecting, spatial trimming to the study area, and possibly dissolving river segments), but the riverdist package and its dependencies also include tools for accomplishing the necessary formatting within R.

Author(s)

Matt Tyers
Maintainer: Matt Tyers <matt.tyers@alaska.gov>

Description

A complex river network object, a subset of the streams in the Absaroka-Beartooth Wilderness.

Usage

data(abstreams)

Format

A river network object, see rivernetwork
abstreams0  Dataset: A-B Streams 0

Description
An unusably messy river network object, included for the purpose of testing river network editing functions.

Usage
data(abstreams0)

Format
A river network object, see rivernetwork

addcumuldist  Add Cumulative Distance to a River Network

Description
Adds a vector of cumulative distances to a river network. Called internally.

Usage
addcumuldist(rivers)

Arguments
rivers  The river network object to use.

Value
Returns an object of class "rivernetwork" containing all spatial and topological information. See rivernetwork-class.

Author(s)
Matt Tyers

Examples
Gulk1 <- addcumuldist(rivers=Gulk)
addverts

**Add Vertices To Maintain a Minimum Distance Between Vertices**

**Description**

In certain cases, such as when there is a lake within a river system, there may be long, straight lines in a river network with vertices only at either end. In these cases, any point data along these stretches would be snapped to the vertices at either end. This function automatically adds equally-spaced vertices along the straight line, according to a specified minimum allowable distance between vertices.

**Usage**

```r
addverts(rivers, mindist = 500)
```

**Arguments**

- `rivers`: The river network object to use.
- `mindist`: The minimum distance to use between vertices. Defaults to 500.

**Value**

A new river network object with the specified segments connected (see `rivernetwork`)

**Note**

This function is called within `cleanup`, which is recommended in most cases.

**Author(s)**

Matt Tyers

**See Also**

- `line2network`

**Examples**

```r
data(Kenai3)
Kenai3split <- addverts(Kenai3,mindist=200)

zoomtoseg(seg=c(47,74,78), rivers=Kenai3)
points(Kenai3$lines[[74]]) # vertices before adding

zoomtoseg(seg=c(47,74,78), rivers=Kenai3split)
points(Kenai3split$lines[[74]]) # vertices after adding
```
**Description**

Adds lookup tables for distance computation, dramatically reducing computation time. It may take some time to calculate, particularly in a braided network.

**Usage**

```r
buildlookup(rivers)
```

**Arguments**

- `rivers` The river network object to use

**Value**

A rivernetwork object, with a new list element, `$distlookup`, a list of three matrices. Element `[i, j]` of each matrix corresponds to the route between segment `i` and `j`. The `$distlookup$distlookup$distlookup[middist]` matrix gives the total distance of the "middle" of each route (between the starting and ending segments"), and the `$distlookup$distlookup$starttop` and `$distlookup$distlookup$endtop` matrices have value TRUE, FALSE, or NA if the segments at the beginning or end of the route are connected to the rest of the route at the top of the coordinate matrix, bottom of the coordinate matrix, or if the route is contained to just one segment, respectively. (See `rivernetwork`.)

**Note**

This will add three `n` by `n` matrices to the river network object, which will be very large if the river network has many segments.

This function is called within `cleanup`, which is recommended in most cases. It is also called within `buildsegroutes`, and will add lookup tables by default if there are fewer than 400 segments in the river network.

This function can still be called in the presence of a braided network, but all resulting distances used in subsequent analyses will be the shortest route.

If segment routes (`$segroutes`) are not present, this function may take a very long time to run.

**Author(s)**

Matt Tyers

**Examples**

```r
data(abstreams)
abstreams1 <- buildlookup(abstreams)
```
**buildsegroutes**

---

**Build Segment Routes**

**Description**

Adds the travel routes from the mouth (lowest point) of a river network to each segment, and (optionally) distance lookup tables. This greatly reduces the time needed to detect routes, making distance calculation much more efficient, particularly in the case of multiple distance calculations.

**Usage**

```r
buildsegroutes(rivers, lookup = NULL, verbose = FALSE)
```

**Arguments**

- **rivers**: The river network object to use
- **lookup**: Whether to build lookup tables as well. This may take some time, but will result in even faster distance computation in analyses (see `buildlookup`). Because of the object size returned, this may not be advisable in a large river network (more than a few hundred segments). Accepts `TRUE` or `FALSE`, and defaults to `NULL`. If the default value is accepted, lookup tables will be built if the river network has 400 segments or fewer.
- **verbose**: Whether or not to print the segment number the function is currently building a route for (used for error checking). Defaults to `FALSE`.

**Value**

A rivernetwork object, with a new list element, `Dsegroutes`, which gives the route from the mouth to each rivernetwork segment. Optionally, it may add `Ddistlookup`, distance lookup tables for even faster distance computation. (See `rivernetwork`.)

**Note**

In the event of braiding (multiple channels), it is likely that there will be differences in the routes detected. If this is the case, building routes will likely result in a shorter and more efficient route. Regardless, extreme caution is always advised in the event of braiding.

The mouth segment and vertex must be specified (see `setmouth`).

This function is called within `cleanup`, which is recommended in most cases.

**Author(s)**

Matt Tyers
calculateconnections

**Examples**

data(abstreams)
plot(x=abstreams)
abstreams1 <- abstreams
abstreams1$segroutes <- NULL  # taking out the $segroutes component

# before

tstart <- Sys.time()
detectroute(start=120, end=111, rivers=abstreams1)
Sys.time() - tstart

# after

tstart <- Sys.time()
detectroute(start=120, end=111, rivers=abstreams1)
Sys.time() - tstart

calculateconnections  *Calculate the Connectivity Matrix for a River Network*

**Description**

Calculates the connectivity matrix for a river network, during import and editing. Called internally.

**Usage**

calculateconnections(lines, tolerance)

**Arguments**

- **lines**  
  A list of coordinate matrices, each corresponding to a line segment.

- **tolerance**  
  The spatial tolerance for establishing connectivity.

**Value**

A matrix with topological information. See the $connections element of the rivernetwork-class.

**Author(s)**

Matt Tyers

**Examples**

Gulk_connections <- calculateconnections(lines=Gulk$lines, tolerance=Gulk$tolerance)
checkbraided

Description
Detects braiding (multiple flow channels between two locations) within a river network object. Braiding can either be checked for in the route between two segments, or in the river network as a whole.

Usage
```r
checkbraided(rivers, startseg = NULL, endseg = NULL, progress = TRUE)
```

Arguments
- **rivers**: The river network object to check.
- **startseg**: Starting segment of a route to investigate. If this and endseg are NULL, the full river network will be checked.
- **endseg**: Starting segment of a route to investigate. If this and startseg are NULL, the full river network will be checked.
- **progress**: Whether to show the progress bar. Defaults to TRUE.

Note
This function is called within `cleanup`, which is recommended in most cases.

Author(s)
Matt Tyers

Examples
```r
data(Gulk)
plot(x=Gulk)
checkbraided(rivers=Gulk)

data(KilleyW)
plot(x=KilleyW)
checkbraided(rivers=KilleyW)

Kenai3.subset <- trimriver(trimto=c(22,2,70,30,15,98,96,89,52,3), rivers=Kenai3)
plot(x=Kenai3.subset)

checkbraided(startseg=1, endseg=7, rivers=Kenai3.subset)
checkbraided(startseg=1, endseg=5, rivers=Kenai3.subset)
```
checkbraidedTF  

**Check for Braiding in a River Network**

**Description**

Detects braiding (multiple flow channels between two locations) within a river network object, and returns a logical value for specifying braiding within a river network object.

**Usage**

`checkbraidedTF(rivers, toreturn = "rivers", progress = TRUE)`

**Arguments**

- `rivers`: The river network object to check.
- `toreturn`: Specifying `toreturn = "rivers"` (the default) will return a river network object with a value of TRUE or FALSE assigned to the $braided element of the river network object. Specifying `toreturn = "logical"` will just return TRUE if braiding is detected or FALSE if no braiding is detected. Specifying `toreturn = "routes"` will return the first two differing routes detected, which may be useful in identifying where the problem lies.
- `progress`: Whether to show the progress bar. Defaults to TRUE.

**Note**

This function is called within `cleanup`, which is recommended in most cases.

**Author(s)**

Matt Tyers

**Examples**

```r
data(Gulk, KilleyW)
Gulk <- setmouth(seg=1, vert=1, rivers=Gulk)
plot(x=Gulk)
checkbraidedTF(rivers=Gulk, toreturn="logical")

KilleyW <- setmouth(seg=1, vert=288, rivers=KilleyW)
plot(x=KilleyW)
checkbraidedTF(rivers=KilleyW, toreturn="logical")
checkbraidedTF(rivers=KilleyW, toreturn="routes")

KilleyW.1 <- checkbraidedTF(rivers=KilleyW, toreturn="rivers")
str(KilleyW.1)
```
**cleanup**

**Interactive Cleanup of a River Network**

**Description**

This is the recommended function to use for cleanup of a river network. It calls all available river network editing functions in appropriate sequence, detecting which are needed or recommended, and prompts user input wherever necessary.

Currently, it automatically calls `removeduplicates`, prompts the user whether to run `dissolve`, automatically runs `removemicrosegs` and `splitsegments` if needed, provides user prompts for `addverts` and `setmouth`, detects if segments are unconnected and provides user prompts for `removeunconnected` or `connectsegs`, automatically runs `checkbraidedTF`, and prompts the user whether to run `buildsegroutes` if no braiding is detected.

**Usage**

```
cleanup(rivers)
```

**Arguments**

- `rivers` The river network object to use

**Value**

A new river network object, see `rivernetwork`

**Author(s)**

Matt Tyers

**See Also**

`line2network`

**Examples**

```r
data(abstreams0,Koyukuk0,Kenai1)

# abstreams_fixed <- cleanup(abstreams0)
# Koyukuk <- cleanup(Koyukuk0)
# Kenai <- cleanup(Kenai1)
```
cleanup_verts | Interactive Cleanup of the Vertices of Individual Segments

Description
A trial version of a function for deep-cleaning a river network.

Sometimes a shapefile contains errors that are not obvious at an initial check, typically vertices that
should not be there.

This function steps through each segment in sequence, and allows the user to interactively remove
vertices.

Usage
cleanup_verts(rivers, startwith = 1)

Arguments
rivers The river network object to use
startwith The segment (number) to start with, defaulting to 1.

Value
A new river network object, see rivernetwork

Note
Stepping through a large and messy river network can be time-consuming. To resume a cleanup ses-
tion, use the startwith= argument and the last returned river network. For example, if rivers1 <-
cleanup_verts(rivers) were initially called and the user selected "save & close" at segment 100, cleanup can be resumed
by calling rivers2 <- cleanup_verts(rivers1, startwith=100).

Author(s)
Matt Tyers

See Also
line2network

Examples
data(abstreams0, Koyukuk0, Kenai1)

# abstreams_fixed1 <- cleanup_verts(abstreams0)
# Koyukuk <- cleanup(Koyukuk0)
# Kenai <- cleanup(Kenai1)
connectsegs

**Connect Segments**

**Description**

Provides a method to manually connect unconnected segments within a river network. The nearest endpoint (or vertex) of the second segment is added as a new vertex to the first, and the network topology is then updated.

**Usage**

```r
collectsegs(connect, connectto, nearestvert = TRUE, rivers, calcconnections = TRUE)
```

**Arguments**

- `connect`: The segment(s) to connect to the network. Typically, this is the segment that is disconnected from the rest of the river network. A vector of segments may be used.
- `connectto`: The segment(s) to connect it (them) to. Typically, this is a segment that is connected to the rest of the river network. A vector of segments may be used, corresponding to that used in `connect`.
- `nearestvert`: Whether to connect at the nearest vertex and split the segment (FALSE), or connect at the nearest endpoint (TRUE). Defaults to TRUE. A vector may be used, corresponding to those used in `connect` and `connectto`.
- `rivers`: The river network object to use.
- `calcconnections`: Whether to recalculate all connections. Defaults to TRUE. Setting to FALSE is not recommended unless many connections are to be made, in which case connections can be calculated afterward.

**Value**

A new river network object with the specified segments connected (see `rivernetwork`)

**Note**

This function is called within `cleanup`, which is recommended in most cases.

**Author(s)**

Matt Tyers

**See Also**

`line2network`
Examples

data(Koyukuk0)
plot(Koyukuk0, ylim=c(1930500,1931500), xlim=c(194900,195100))
topologydots(Koyukuk0, add=TRUE)

Koyukuk0.1 <- connectsegs(connect=21, connectto=20, rivers=Koyukuk0)
plot(Koyukuk0.1, ylim=c(1930500,1931500), xlim=c(194900,195100))
topologydots(Koyukuk0.1, add=TRUE)

# or the vector version
zoomtoseg(seg=21:23, rivers=Koyukuk0)
Koyukuk0.2 <- connectsegs(connect=c(20,21,22), connectto=c(21,22,23),
nearestvert=c(FALSE,FALSE,TRUE), rivers=Koyukuk0)
zoomtoseg(seg=21:23, rivers=Koyukuk0.2)
topologydots(Koyukuk0.2, add=TRUE)

detectroute.jpg

detectroute

Detect Route

Description

Called internally within riverdistance. Detects the sequential route from one river network segment to another.

Usage

detectroute(start, end, rivers, verbose = FALSE, stopiferror = TRUE,
algorithm = NULL)

Arguments

- **start**: Segment number of the start of the route
- **end**: Segment number of the end of the route
- **rivers**: The river network object to use
- **verbose**: Whether or not to print all routes being considered (used for error checking). Defaults to FALSE.
- **stopiferror**: Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, detectroute() will return NA. Defaults to TRUE.
- **algorithm**: Which route detection algorithm to use. If set to NULL (the default), the function will automatically make a selection. Choices are:
  - Setting algorithm="sequential" will be quite slow, and may give inaccurate results in the event of braiding. This algorithm returns the first complete route detected, which may not be the shortest. This algorithm is not recommended in almost all cases, but is retained as an option for certain checks. It will not be used unless specified.
Setting algorithm="Dijkstra" will be much faster, and will return the shortest route in the event of braiding. If braiding is present or unknown, this will be the algorithm automatically chosen.

Setting algorithm="segroutes" will be the fastest of all, but will only return results in a non-braided network. This will be the algorithm automatically selected if segment routes are present - see buildsegroutes.

Value

A vector of segment numbers corresponding to the ordered route.

Author(s)

Matt Tyers

Examples

data(Gulk)
plot(x=Gulk, cex=1)
detectroute(start=6, end=14, rivers=Gulk)

tstart <- Sys.time()
detectroute(start=120, end=111, rivers=abstreams, algorithm="sequential")
tend <- Sys.time()
tend - tstart

data(abstreams)
tstart <- Sys.time()
detectroute(start=120, end=111, rivers=abstreams, algorithm="Dijkstra")
tend <- Sys.time()
tend - tstart

tstart <- Sys.time()
detectroute(start=120, end=111, rivers=abstreams, algorithm="segroutes")
tend <- Sys.time()
tend - tstart

dissolve

Dissolve

Description

Acts like a spatial dissolve within a GIS environment. Simplifies a river network object by combining "runs" of segments with no other connections.

Usage

dissolve(rivers)
Arguments

rivers The river network object to use

Value

A new river network object with segments combined

Note

This function is called within cleanup, which is recommended in most cases.

Author(s)

Matt Tyers

See Also

line2network

Examples

data(Kenai2)
plot(x=Kenai2)

Kenai2dissolve <- dissolve(rivers=Kenai2)
plot(x=Kenai2dissolve)

---

fakefish

Dataset: Fakefish

Description

A set of observations of Fakefish on the Gulkana River and its tributaries.

Usage

data(fakefish)

Format

A data frame
fakefish_density

Details

- x. X-coordinate of observation (Alaska Albers Equal Area). Note that the locations do not align with the river network object.
- y. Y-coordinate of observation
- seg. River segment (with x- and y-coordinates snapped to river network object)
- vert. River vertex
- fish.id. Numeric identifier for each fish (individual fish were observed more than once)
- flight. Numeric identifier for each telemetry flight
- flight.date. Date of each telemetry flight

See Also

Gulk

fakefish_density  Dataset: Fakefish Density

Description

An object created by riverdensity, describing the density of Fakefish points in the Gulkana River during ten surveys.

Usage

data(fakefish_density)

Format

A river density object, see riverdensity, plotriverdensity, riverdensity-class

Details

Intended for plotting using plotriverdensity.
Gulkna River

**Description**

A stretch of Gulkana River and tributaries.

**Usage**

data(Gulk)

**Format**

A river network object, see `rivernetwork`.

```
highlightseg(seg, rivers, cex = 0.8, lwd = 3, add = FALSE, color = FALSE, ...)
```

**Arguments**

- `seg`: A vector of segments to highlight
- `rivers`: The river network object to use
- `cex`: The character expansion factor to use for segment labels
- `lwd`: The line width to use for highlighted segments
- `add`: Whether to add the highlighted segments to an existing plot (TRUE) or call a new plot (FALSE). Defaults to FALSE.
- `color`: Whether to display segment labels as the same color as the segments. Defaults to FALSE.
- `...`: Additional plotting arguments (see `par`)

**Author(s)**

Matt Tyers
homerange

Examples

```r
data(Kenai3)
plot(Kenai3)
highlightseg(seg=c(10,30,68),rivers=Kenai3)
```

homerange  Home Range

Description

Returns the minimum observed home range for multiple observations of each individual fish.

Usage

```r
homerange(unique = NULL, survey = NULL, seg, vert, rivers, map = FALSE,
  algorithm = NULL, main = NULL, ...)
```

Arguments

- `unique`  
  A vector of unique identifiers for each fish. If the default (NULL) is used, the function will assume all observations come from a single individual.

- `survey`  
  A vector of survey identifiers for each fish. This argument is not needed for home range calculation, but can affect plotting (see `plot.homerange`).

- `seg`  
  A vector of river locations (segment component).

- `vert`  
  A vector of river locations (vertex component).

- `rivers`  
  The river network object to use.

- `map`  
  Deprecated, use `plot.homerange` for plotting instead. Originally, whether to produce sanity-check maps of observed locations and calculated home range for each fish.

- `algorithm`  
  Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See `detectroute` for more details.

- `main`  
  Deprecated, use `plot.homerange` for plotting instead. Originally, plot title, if `map` is set to TRUE. If unspecified, the unique ID will be used for the title.

- `...`  
  Deprecated, use `plot.homerange` for plotting instead. Originally, additional plotting arguments, if `map` is set to TRUE.

Value

An object of the `homerange-class`. The `ranges` element is a data frame with two columns: $ID$ is a list of unique fish (as specified by `unique=`), and `$range` is calculated minimum home range, in the units of the coordinate system (this will likely be meters). The other elements are used for plotting, see `homerange-class` for more details.
homerange-class

The "homerange" Class

A class that holds information computed from the `homerange` function. Contains all information for plotting in `plot.homerange`.

Elements

- ranges: Object of class "data.frame". Contains a column of the identifiers for each individual, and a column of the associated home ranges.
- subseg_n: List of the number of times each subsegment was traveled. The first level of the list corresponds to individual, the second level to river segment.
- subseg_length: List of lengths of each subsegment.
- seg, vert, unique, rivers: All inputs from the original `homerange` call.

Author(s)

Matt Tyers
Description

Returns matrices describing the overlap of the minimum observed home range for multiple observations of each individual fish.

Usage

```
homerangeoverlap(x)
```

Arguments

- `x` An object returned from `homerange`.

Value

A list of three matrices, with `either` giving the distances represented by the union of home ranges of each pair of individuals, and `both` giving the distances represented by the intersection of home ranges of each pair of individuals. Element `prop_both` gives the proportion of overlap, defined as intersection/union.

Author(s)

Matt Tyers

See Also

- `homerange`, `plot.homerange`, `plothomerangeoverlap`

Examples

```
data(Gulk, fakefish)
ranges <- with(fakefish, homerange(unique=fish.id, survey=flight, seg=seg, vert=vert, rivers=Gulk))
ranges

# 19 plots will be produced, recommend calling par(mfrow=c(4,5))
plot(ranges)
plot(ranges,cumulative=TRUE,label=TRUE)

homerangeoverlap(ranges)

plothomerangeoverlap(ranges)
with(fakefish, riverpoints(seg=seg, vert=vert, rivers=Gulk))
```
isflowconnected  

*Check Flow-Connectedness*

**Description**

Checks to see if two segments are flow-connected. Called internally within `riverdirection` and `upstream`.

**Usage**

`isflowconnected(seg1, seg2, rivers, stopiferror = TRUE, algorithm = NULL)`

**Arguments**

- **seg1**: First input segment
- **seg2**: Second input segment
- **rivers**: The river network object to use
- **stopiferror**: Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return `NA` in the appropriate entry. Defaults to TRUE. See `detectroute`.
- **algorithm**: Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See `detectroute` for more details.

**Value**

Logical TRUE if the two segments are flow-connected, FALSE if they are not

**Note**

The river mouth must be specified (see `setmouth`).

**Author(s)**

Matt Tyers

**Examples**

```r
data(Gulk)
plot(Gulk)

Gulk <- setmouth(seg=1, vert=1, rivers=Gulk)

isflowconnected(seg1=13, seg2=14, rivers=Gulk)
isflowconnected(seg1=13, seg2=1, rivers=Gulk)
```
**Kenai1**  

Dataset: Kenai River 1

**Description**

A first pass at a messy river network object.

**Usage**

`data(Kenai1)`

**Format**

A river network object, see `rivernetwork`

**See Also**

Kenai2, Kenai3

---

**Kenai2**  

Dataset: Kenai River 2

**Description**

A second pass at a messy river network object. In this iteration of cleanup, several non-connected segments have been removed.

**Usage**

`data(Kenai2)`

**Format**

A river network object, see `rivernetwork`

**See Also**

Kenai1, Kenai3
Kenai3  

*Dataset: Kenai River 3*

**Description**

A third pass at a messy river network object. In this iteration of cleanup, several non-connected segments have been removed, and several series of segments have been dissolved into single segments.

**Usage**

```r
data(Kenai3)
```

**Format**

A river network object, see `rivernetwork`

**See Also**

`Kenai1`, `Kenai2`

---

**kfunc**  

*Plotting K-functions for a Set of Surveys*

**Description**

Plots K-functions for locations in each of a set of surveys. In this implementation, this can be interpreted as the proportion of additional fish within a given distance. This will increase as a function of distance, and may provide evidence of clustering or dispersion features, particularly if the envelope is used.

**Usage**

```r
kfunc(seg, vert, survey = NULL, rivers, lwd = 2, envelope = TRUE,
envreps = 1000, envcol = "grey80", envborder = NA, maxdist = NULL,
xlab = "Distance", ylab = "% within", showN = TRUE,
whichplots = NULL, returnoutput = FALSE, ...)
```

**Arguments**

- `seg`  
  A vector of river locations (segment)
- `vert`  
  A vector of river locations (vertex)
- `survey`  
  A vector of survey IDs corresponding to the values of `seg` and `vert`. Defaults to NULL. If this argument is used, K-functions will be calculated for each unique survey, and separate plots will be produced.
- `rivers`  
  The river network object to use
**kfunc**

lwd Line width used for plotting. Defaults to 2.

envelope Whether to construct and display a 95 percent confidence envelope (see note.) Defaults to TRUE if survey is specified, and is automatically FALSE otherwise.

enreps Number of bootstrap replicates to use for envelope calculation. Defaults to 1000.

envcol Color to use for envelope plotting. Defaults to "grey80".

envborder Border color to use for envelope plotting. Defaults to NA, which will result in no border being plotted.

maxdist Maximum distance (x-axis value) for plotting. The default value (NULL) will result in an appropriate value being chosen.

xlab X-coordinate label for plotting

ylab Y-coordinate label for plotting

showN Whether to show the sample size for each survey in each plot title. Defaults to TRUE.

whichplots A vector of plots to produce, if multiple plots are produced. For example, specifying whichplot=c(2,3,4) will result in only the second, third, and fourth plots of the sequence being produced. Accepting the default (NULL) will result in all plots being produced.

returnoutput Whether to return output instead of producing a plot. Defaults to FALSE.

... Additional plotting parameters.

**Note**

K-function envelopes for each survey are constructed by bootstrapping all within-survey distances, that is, the distances between all individuals within each survey, for all surveys. This results in a confidence envelope under the assumption that spacing is independent of survey; therefore a survey K-function outside the envelope provides evidence of clustering or dispersal in that survey that is outside the typical range. An envelope is not available if only one survey is plotted.

A K-function above the envelope for a given distance range provides evidence of a greater number of individuals than expected at that distance range (clustering); A K-function below the envelope for a given distance range provides evidence of a smaller number of individuals than expected at that distance range (dispersal).

This function is distance-computation intensive, and will be extremely slow-running if a river network is used that does not have segment routes and/or distance lookup tables for fast distance computation. See buildsegroutes and/or buildlookup for more information.

**Author(s)**

Matt Tyers

**Examples**

data(Hulk, fakefish)

# # 10 plots will be created - recommend calling
# # par(mfrow=c(3,4))
kfunc(seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk, survey=fakefish$flight, 
enreps=100, maxdist=200000)

### This shows relatively high amounts of clustering for surveys 1 and 8, 
### and relatively high amounts of dispersal in surveys 5 and 6.

### plotting the survey locations that led to this calculation, for comparison

### 10 plots will be created - recommend calling
### par(mfrow=c(3,4))
for(i in 1:10) {
  plot(x=Gulk, segmentnum=FALSE, color=FALSE, main=i)
  riverpoints(seg=fakefish$seg[fakefish$flight==i], 
    vert=fakefish$vert[fakefish$flight==i], rivers=Gulk, col=2, pch=15)
}

---

**KilleyW**  
**Dataset: Killey River, West Channel**

**Description**

A messy and braided section of the Kenai River network - actually a subset of Kenai3.

**Usage**

data(KilleyW)

**Format**

A river network object, see rivernetwork

---

**Koyukuk0**  
**Dataset: Koyukuk River 0**

**Description**

An unusably messy river network object, included for the purpose of testing river network editing functions.

**Usage**

data(Koyukuk0)

**Format**

A river network object, see rivernetwork


**Koyukuk1**

**See Also**

Koyukuk1, Koyukuk2

---

**Koyukuk1**

*Dataset: Koyukuk River 1*

**Description**

A first pass at a messy river network object. The way it was dissolved in ArcGIS makes the endpoints appear disconnected to line2network and the topologies do not work.

**Usage**

`data(Koyukuk1)`

**Format**

A river network object, see `rivernetwork`

**See Also**

Koyukuk2

---

**Koyukuk2**

*Dataset: Koyukuk River 2*

**Description**

A second pass at a messy river network object, with topologies fixed from Koyukuk1.

**Usage**

`data(Koyukuk2)`

**Format**

A river network object, see `rivernetwork`

**See Also**

Koyukuk1
line2network

Create a River Network Object from a Shapefile

Description

Uses readOGR in package 'rgdal' to read a river shapefile, and establishes connectivity of segment endpoints based on spatial proximity.

Usage

```r
line2network(path = ".", layer, tolerance = 100, reproject = NULL,
             supplyprojection = NULL)
```

Arguments

- `path` : File path, default is the current working directory.
- `layer` : Name of the shapefile, without the .shp extension.
- `tolerance` : Snapping tolerance of segment endpoints to determine connectivity. Default is 100, therefore care should be exercised when working with larger units of distance, such as km.
- `reproject` : A valid Proj.4 projection string, if the shapefile is to be re-projected. Re-projection is done using spTransform in package 'sp'.
- `supplyprojection` : A valid Proj.4 projection string, if the input shapefile does not have the projection information attached.

Value

Returns an object of class "rivernetwork" containing all spatial and topological information. See rivernetwork-class.

Note

Since distance can only be calculated using projected coordinates, line2network() will generate an error if a non-projected input shapefile is detected. To resolve this, the shapefile can be re-projected in a GIS environment, or using reproject=, shown in the second example below.

Author(s)

Matt Tyers
Examples

```r/filepath <- system.file("extdata", package="riverdist")
Gulk_UTM5 <- line2network(path=filepath, layer="Gulk_UTM5")
plot(Gulk_UTM5)

# Re-projecting in Alaska Albers Equal Area projection:
AKalbers <- "+proj=aea +lat_1=55 +lat_2=65 +lat_0=50 +lon_0=-154
+x_0=0 +y_0=0 +datum=NAD83 +units=m +no_defs +ellps=GRS80 +towgs84=0,0,0"
Gulk_AKalbers <- line2network(path=filepath, layer="Gulk_UTM5", reproject=AKalbers)
plot(Gulk_AKalbers)
```

---

**Dataset: Line 98 of Kenai River 1 (Long-Lat)**

**Description**

A matrix of coordinates in longitude-latitude, used to illustrate coordinate transformation. Coordinates come from arbitrary line number 98 in the Kenai River 1 shapefile, rendered in long-lat.

**Usage**

data(line98)

**Format**

A matrix of values

---

**makeriverdensity**

*Calculate Kernel Density Using River Distance*

**Description**

Uses spatial point data (segment and vertex) to calculate a kernel density object to use in the output class plotting method, `plot.riverdensity`. Scaled kernel density is calculated at approximately regularly-spaced locations, with spacing specified by the user.

If an argument is used in the survey field, kernel densities will be calculated for each unique value of survey, resulting in a separate plot for each.

The purpose of this function is to generate a kernel density object to plot using plot(), see `plot.riverdensity`.
Usage

makeriverdensity(seg, vert, rivers, survey = NULL, kernel = "gaussian", bw = NULL, resolution = NULL)

Arguments

seg A vector of river locations (segment)
vert A vector of river locations (vertex)
rivers The river network object to use
survey A vector of survey IDs corresponding to the values of seg and vert. If this argument is used, kernel densities will be calculated for each unique survey, and separate plots will be produced.
kernl The type of density kernel to use. Allowed types are "gaussian" (normal) and "rect" (rectangular, giving simple density). Defaults to "gaussian".
bw The kernel bandwidth to use. If kernel is set to "gaussian", this provides the standard deviation of the gaussian (normal) kernel to use. If kernel is set to "rect", this provides the half-width of the rectangular kernel, or the distance to use in simple density. Accepting the default (NULL) will result in the function determining a value to use, based on the total length of the river network and the value of the resolution argument.
resolution The approximate spacing of the river locations used for kernel density calculation. Accepting the default (NULL) will result in the function determining a value to use, based on the total length of the river network.

Value

A river density object, see riverdensity-class.

Note

It is likely that calculation will be very slow. Use of this function with a river network for which segment routes has not yet been calculated is not recommended.

This function is distance-computation intensive, and may be slow-running if a river network is used that does not have segment routes and/or distance lookup tables for fast distance computation. See buildsegroutes and/or buildlookup for more information.

Author(s)

Matt Tyers

See Also

plot.riverdensity, plotriverdensitypoints
mapbyname

Examples

data(Gulk, fakefish)

Gulk_dens <- makeriverdensity(seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk, survey=fakefish$flight.date)

# # 10 plots will be created, recommend calling par(mfrow=c(2,5))
plot(x=Gulk_dens)

__________________________

mapbyname  

Map Segments by Name

__________________________

Description

Provides a check that river network segments were appropriately named.

Usage

mapbyname(rivers, scale = TRUE, cex = 0.6, ...)

Arguments

rivers  
The river network object to use. Function checks segment names contained in the river network object.
scale  
Whether or not to give x- and y-axes the same scale
cex  
Global character expansion factor for plotting
...

Additional plotting arguments (see par)

Author(s)

Matt Tyers

Examples

data(Gulk)
str(Gulk)

Gulk$names <- c("Gulkana River","Trib 1","West Fork","Gulkana River","Trib 1", "West Fork","Trib 2","West Fork","Twelvemile Creek","Gulkana River", "Middle Fork","Gulkana River","Middle Fork","Hungry Hollow")

str(Gulk)

mapbyname(rivers=Gulk)
**matbysurveylist**

*Generate List of Distance Matrix Between Observations, for All Individuals*

**Description**

Returns a list of matrices, each giving the river distance, direction, or upstream travel distance between all observations of one unique fish. This function is principally intended for producing an object to plot in `plotmatbysurveylist`.

**Usage**

```
matbysurveylist(unique, survey, seg, vert, rivers, indiv = NULL, 
method = "upstream", flowconnected = FALSE, net = FALSE, 
stopiferror = TRUE, algorithm = NULL)
```

**Arguments**

- `unique` A vector of unique identifiers for each fish.
- `survey` A vector of identifiers for each survey. It is recommended to use a numeric or date format (see `as.Date`) to preserve survey order.
- `seg` A vector of river locations (segment component).
- `vert` A vector of river locations (vertex component).
- `rivers` The river network object to use.
- `indiv` A vector of unique individuals to use. Accepting the default (NULL) will result in a matrix being returned for all unique individuals.
- `method` Which general method to use. Setting `method="distance"` will compute distance for each pair of observation, setting `method="direction"` will compute direction between each pair of observation, and setting `method="upstream"` will compute directional (upstream) distance between each pair of observation. Defaults to "upstream".
- `flowconnected` Optional parameter to pass to the distance or direction calculation. Defaults to FALSE.
- `net` Optional parameter to pass to the distance or direction calculation. Defaults to FALSE.
- `stopiferror` Optional parameter to pass to the distance or direction calculation. Defaults to TRUE.
- `algorithm` Optional parameter to pass to the distance or direction calculation. Defaults to NULL.

**Value**

A list with each element corresponding to a unique fish. Each list element is the output from either `riverdistancematbysurvey`, `riverdirectionmatbysurvey`, or `upstreammatbysurvey`.
mouthdist

Note

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see buildsegroutes).

Author(s)

Matt Tyers

See Also

riverdistance, riverdirection, upstream, riverdistancematbysurvey, riverdirectionmatbysurvey, upstreammatbysurvey, plotmatbysurveylist

Examples

data(Gulk, smallset)
matabysurveylist <- matabysurveylist(unique=smallset$id, survey=smallset$flight, seg=smallset$seg, vert=smallset$vert, rivers=Gulk)
plotmatabysurveylist(matabysurveylist)
plotmatabysurveylist(matabysurveylist,type="confint")
plotmatabysurveylist(matabysurveylist,type="dotplot")

data(fakefish)
# matabysurveylist <- matabysurveylist(unique=fakefish$fish.id, survey=fakefish$flight,
#   seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)
# plotmatabysurveylist(matabysurveylist)

mouthdist Distance From Mouth

Description

Calculates distance from river locations (given as vectors of segment and vertex) and the specified mouth of the river network. The mouth must first be specified (see setmouth).

Usage

mouthdist(seg, vert, rivers, stopiferror = TRUE, algorithm = NULL)

Arguments

seg Vector of segments
vert Vector of vertices
rivers The river network object to use
stopiferror Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See detectroute.
Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See `detectroute` for more details.

### Value

Distance (numeric)

### Note

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see `buildsegroutes`).

### Author(s)

Matt Tyers

### Examples

```r
data(Gulk)

# Mouth must be specified
Gulk$mouth$mouth.seg <- 1
Gulk$mouth$mouth.vert <- 1

mouthdist(seg=4, vert=40, rivers=Gulk)
mouthdist(seg=c(4,5), vert=c(40,20), rivers=Gulk)
```

### Description

Calculates distance from the mouth of a river network to all observations of each individual (given as segment and vertex) and the specified mouth of the river network. The mouth must first be specified (see `setmouth`). Returns a matrix of distances, with a row for each unique individual and a column for each survey.

A plotting method is provided for the output; see `plotseq`.

### Usage

```r
mouthdistbysurvey(unique, survey, seg, vert, rivers, logical = NULL, stopiferror = TRUE, algorithm = NULL)
```
Arguments

- **unique**: A vector of identifiers for each fish.
- **survey**: A vector of identifiers for each survey. It is recommended to use a numeric or date format (see `as.Date`) to preserve survey order.
- **seg**: A vector of river locations (segment).
- **vert**: A vector of river coordinates (vertex).
- **rivers**: The river network object to use.
- **logical**: A boolean vector that can be used for subsetting - if used, `mouthdistbysurvey()` will only return distances in which a specified condition is met.
- **stopiferror**: Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See `detectroute`.
- **algorithm**: Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See `detectroute` for more details.

Value

A vector of river network distances (numeric), with each row corresponding to a unique fish and each column corresponding to a unique survey. Values of NA indicate the individual not being located during the survey in question.

Note

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see `buildsegroutes`).

Author(s)

Matt Tyers

See Also

- `plotseq`

Examples

```r
data(Gulk, fakefish)

seqbysurvey <- mouthdistbysurvey(unique=fakefish$fish.id, survey=fakefish$flight.date, seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)
seqbysurvey
plotseq(seqbysurvey)
```
### pdist

**Pythagorean Distance**

**Description**

Pythagorean distance between two points. Called internally.

**Usage**

```r
pdist(p1, p2)
```

**Arguments**

- `p1` X-Y coordinates of point 1
- `p2` X-Y coordinates of point 2

**Value**

Distance (numeric)

**Author(s)**

Matt Tyers

**Examples**

```r
point1 <- c(1,3)
point2 <- c(4,7)
pdist(point1, point2)
```

### pdisttot

**Total Pythagorean Distance**

**Description**

Total Pythagorean distance of a sequence of points. Called internally.

**Usage**

```r
pdisttot(xy)
```

**Arguments**

- `xy` A matrix of X-Y coordinates of the sequence of points.
plot.homerange

Value
Distance (numeric)

Author(s)
Matt Tyers

Examples
points <- matrix(c(1:10), nrow=5, ncol=2, byrow=FALSE)
pdisttot(xy=points)

plot.homerange  Plot Home Range

Description
Plotting method for home range, the minimum observed home range for multiple observations of each individual fish.

Usage
## S3 method for class 'homerange'
plot(x, cumulative = FALSE, lwd = 3, maxlwd = 10,
    col = 4, pch = 21, label = FALSE, main = NULL, ...)

Arguments

x  An object returned from homerange.
 cumulative  Whether to plot travel as cumulative, with line thickness depending on the number of times a given region was traveled by a given individual. Defaults to FALSE.
 lwd  The line width for plotting homerange, or minimum line width if cumulative is TRUE. Defaults to 3.
 maxlwd  The maximum line width if cumulative is TRUE. Defaults to 10.
 col  The line color to use. Defaults to “blue”.
 pch  The point character to use for individual points. Defaults to open circles, the color of lines.
 label  Whether to add survey labels to individual points, if used in homerange. Defaults to FALSE.
 main  Plot title. If the default NULL is used, plots will be titled according to unique individual.
 ...  Additional plotting parameters, see plot.rivernetwork.
plot.riverdensity

Description

Produces a kernel density plot from a kernel density object created by `makeriverdensity`.

If the kernel density object includes densities from multiple surveys, a new plot will be created for each survey.

Densities can be displayed using either line widths, color, or both.

The relative densities that are displayed in the plot are calculated according to the form (density/maxdensity)^pwr, with the value of pwr set by the `pwr` argument. Setting `pwr` to a value less than 1 allows smaller values to be more visible on the plot.

Usage

```r
## S3 method for class 'riverdensity'
plot(x, whichplots = NULL, points = TRUE,
     bycol = TRUE, bylwd = TRUE, maxlwd = 10, pwr = 0.7, scalebyN = TRUE,
     ramp = "grey", lwd = 1, linecol = "black", denscol = "black",
     alpha = 1, dark = 1, showN = TRUE, main = NULL, xlab = "",
     ylab = "", add = FALSE, scalebar = TRUE, ...)
```

Arguments

- **x**: A river density object created by `makeriverdensity`.
- **whichplots**: A vector of plots to produce, if multiple plots are produced. For example, specifying `whichplot=c(2,3,4)` will result in only the second, third, and fourth plots of the sequence being produced. Accepting the default (NULL) will result in all plots being produced.
- **points**: Whether to add the points used for density calculation. Defaults to TRUE.
- **bycol**: Whether to use a color ramp to show densities. Defaults to TRUE.
- **bylwd**: Whether to use line thickness to show densities. Defaults to TRUE.
- **maxlwd**: The maximum line width to use if `bylwd` is set to TRUE. Defaults to 10.
- **pwr**: The power to use in the nonlinear transformation calculating the relative density values to be displayed (see above.) Defaults to 0.7.
- **scalebyN**: Whether to display relative density values scaled by sample size. Specifying `scalebyN=TRUE` will show larger density values associated with surveys with more points, and may be more appropriate for displaying total density. Specifying `scalebyN=FALSE` will allow surveys with smaller sample sizes to be plotted with similar density values as those with larger sample sizes, and may be more appropriate for displaying relative density. Defaults to TRUE.
- **ramp**: The color ramp used to display densities if `bycol` is set to TRUE. Allowed values are "grey" (or "gray"), "red", "green", "blue", "heat", "stoplight", and "rainbow". Defaults to "grey".
- **lwd**: The line width to use for background lines if `bylwd` is set to TRUE, or all lines if `bylwd` is set to FALSE. Defaults to 1.
- **linecol**: The line color to use for background lines if `bylwd` is set to TRUE. Defaults to "black".
- **denscol**: The line color to use for showing density if `bylwd` is set to TRUE. Defaults to "black".
- **alpha**: The opacity value for lines. This could potentially allow multiple density plots to be overlayed with different colors.
- **dark**: A color-saturation adjustment, with values in [0,1]. A value of 1 uses the true colors, and a value less than 1 will render the colors as slightly darker (less saturated), which may be appear better. Defaults to 1.
- **showN**: Whether to automatically include the number of points used as part of the plot title(s).
- **main**: Plot title(s), either given as a single text string which is repeated if multiple plots are produced, or a vector of text strings (one for each plot produced). If multiple plots are produced (resulting from multiple surveys), accepting the default (NULL) will result in each unique value of survey being used as the plot title, along with the sample size if `showN` is set to TRUE.
- **xlab**: X-axis label
- **ylab**: Y-axis label
- **add**: Whether to produce a new plot (FALSE), or add to an existing plot (TRUE). Defaults to FALSE.
Scalebar

Whether to add a scale bar to plot(s). Defaults to TRUE.

Additional plotting parameters.

Author(s)

Matt Tyers

See Also

makeriverdensity, plotriverdensitypoints

Examples

data(Gulk, fakefish)

Gulk_dens <- makeriverdensity(seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk, survey=fakefish$flight.date)

# # 10 plots will be created, recommend calling par(mfrow=c(2,5))
plot(x=Gulk_dens)

plot.rivernetwork

Plotting a River Network

Description

S3 plotting method for the rivernetwork-class. Produces a map of all river segments of a river network object.

Usage

## S3 method for class 'rivernetwork'
plot(x, segmentnum = TRUE, offset = TRUE, lwd = 1,
     cex = 0.6, scale = TRUE, color = TRUE, empty = FALSE, linecol = 1,
     xlab = "", ylab = "", ...)
empty

Creates an empty plot if set to TRUE. Suppresses differentiation by line type if color==FALSE, and suppresses segment number labels. Defaults to FALSE.

linecol

Line color to use if empty is TRUE or color is FALSE. Defaults to black.

xlab

Label for X-axis (defaults to "")

ylab

Label for Y-axis (defaults to "")

... Additional plotting arguments (see par)

Note

This function is intended to provide basic visual checks for the user, not for any real mapping.

Author(s)

Matt Tyers

Examples

data(Gulk)
plot(x=Gulk)

Description

Produces a plot of the overlap of the minimum observed home range for multiple observations of each individual fish, with line thickness illustrating the respective number of individuals’ home-ranges represented.

Usage

plot homerangeoverlap(x, lwd = 3, maxlwd = 10, col = 4, ...)

Arguments

x

An object returned from homerange.

lwd

Minimum line width to use, defaults to 3.

maxlwd

Maximum line width to use, defaults to 10.

col

Line color to use, defaults to "blue".

... Additional plotting parameters, see plot.rivernetwork.

Author(s)

Matt Tyers
plotmatbysurveylist

Plot Upstream Distance Between Observations of All Individuals

Description

Produces a matrix of plots (boxplots are default), with plot \([i,j]\) giving the distribution of upstream distances from observation \(i\) to observation \(j\), for all individuals.

Usage

\[
\text{plotmatbysurveylist(matbysurveylist, type = "boxplot", showN = TRUE, ...)}
\]

Arguments

- **matbysurveylist**: A list of distance matrices returned from `matbysurveylist`.
- **type**: If type is set to "boxplot", boxplots will be produced for each cell. If type is set to "confint", lines denoting an approximate 95 percent confidence interval for the mean will be produced instead. If type is set to "dotplot", a jittered dotplot will be produced for each cell, which will be the most appropriate if sample sizes are small. Defaults to "boxplot".
- **showN**: Whether to display the sample size for each cell. Defaults to TRUE.
- **...**: Additional plotting arguments.

Note

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see `buildsegroutes`).
plotriverdensitypoints

Author(s)

Matt Tyers

See Also

upstream, upstreammatbysurvey

Examples

data(Gulk, smallset)
matbysurveylist <- matbysurveylist(unique=smallset$id, survey=smallset$flight, seg=smallset$seg, vert=smallset$vert, rivers=Gulk)
plotmatbysurveylist(matbysurveylist)
plotmatbysurveylist(matbysurveylist,type="confint")
plotmatbysurveylist(matbysurveylist,type="dotplot")

data(fakefish)
# matbysurveylist <- matbysurveylist(unique=fakefish$fish.id, survey=fakefish$flight, 
# seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)
# plotmatbysurveylist(matbysurveylist)

plotriverdensitypoints

Plot Points Used for Kernel Density

Description

Plots the points used to calculate a kernel density object in makeriverdensity.

This function is intended as a visual check that a sufficient resolution was used.

Usage

plotriverdensitypoints(riverdensity)

Arguments

riverdensity A river density object created by makeriverdensity.

Author(s)

Matt Tyers

See Also

makeriverdensity, plot.riverdensity
Examples

data(Gulk, fakefish)

Gulk_dens <- makeriverdensity(seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)
plotriverdensitypoints(riverdensity=Gulk_dens)

plotseq

Plot Sequence of Observations

Description

Plots the sequence of observations or movements of each individual (given as segment and vertex). This function is primarily intended for use with mouthdistbysurvey, but will also work with riverdistanceseq and upstreamseq.

Usage

plotseq(seqbysurvey, type = "boxplot", xlab = "", ylab = "", main = "",
cex.axisx = 0.8, lowerbound = NULL, upperbound = NULL,
boundtype = "negative", surveysareDates = F, ...)

Arguments

seqbysurvey A matrix returned from mouthdistbysurvey, riverdistanceseq, or upstreamseq.
type The type of plot to generate. Options are "boxplot", "dotplot", "boxline", or "dotline". Defaults to "boxplot".
xlab X-axis label
ylab Y-axis label
main Plot title
cex.axisx Character expansion factor for X-axis labels
lowerbound An optional vector of lower survey bounds
upperbound An optional vector of upper survey bounds
boundtype Method of plotting survey bounds. Options are "positive", "negative" (default), and "lines".
surveysareDates If surveys are in Date format (see as.Date), a value of TRUE allows the x-coordinates points to be spaced apart according to date, not equidistantly. Defaults to FALSE. Any formatting of the survey variable must be done within the original call to mouthdistbysurvey, riverdistanceseq, or upstreamseq. Dates must already be formatted as dates, or in the form "YYYY-MM-DD" or "YYYY/MM/DD".

... Additional plotting parameters
Note

Plots are intended as descriptive only. Any ANOVA-like inference that is suggested from these plots is strongly discouraged. The user is instead advised to use a mixed-effects model or some other inferential tool that accounts for repeated-measures and/or temporal autocorrelation.

Author(s)

Matt Tyers

Examples

data(Gulk, fakefish)

```r
x <- mouthdistbysurvey(unique=fakefish$id, survey=fakefish$flight.date, seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)
```

```r
plotseq(seqbysurvey=x)
plotseq(seqbysurvey=x, type="boxline")
plotseq(seqbysurvey=x, type="dotplot")
plotseq(seqbysurvey=x, type="dotline")
```

```r
plotseq(seqbysurvey=x, type="dotline", surveysareDates=TRUE)
```

```r
from_upstreamseq <- upstreamseq(unique=fakefish$id, survey=fakefish$flight, seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)
plotseq(seqbysurvey=from_upstreamseq)
```

---

pointshp2segvert  

Convert a Point Shapefile to River Locations

Description

This function reads a point shapefile and determines the closest vertex in the river network to each point of XY data, returning a data frame with river locations, defined as segment numbers and vertex numbers, along with the data table read from the input shapefile.

Usage

```r
pointshp2segvert(path = ".", layer, rivers)
```

Arguments

- **path**  
  File path, default is the current working directory.
- **layer**  
  Name of the shapefile, without the .shp extension.
- **rivers**  
  The river network object to use.
Value
A data frame of river locations, with segment numbers in `Dseg`, vertex numbers in `Dvert`, snapping distances in `Dsnapdist`, and the remaining columns corresponding to the data table in the input point shapefile.

Note
If the input shapefile is detected to be in a different projection than the river network, the input shapefile will be re-projected before conversion to river locations.

Author(s)
Matt Tyers

Examples
```r
filepath <- system.file("extdata", package="riverdist")

fakefish_UTM5 <- pointshp2segvert(path=filepath, layer="fakefish_UTM5", rivers=Gulk)
head(fakefish_UTM5)

plot(x=Gulk)
points(fakefish_UTM5$x, fakefish_UTM5$y)
riverpoints(seg=fakefish_UTM5$seg, vert=fakefish_UTM5$vert, rivers=Gulk, pch=16, col=2)
```

Description
Removes duplicated line segments, which can sometimes exist within a shapefile.

Usage
```r
removeduplicates(rivers)
```

Arguments
- `rivers`: The river network object to use

Value
A new river network object with duplicated segments removed, see `rivernetwork`

Author(s)
Matt Tyers
removemicrosegs

See Also

line2network

Examples

data(abstreams0)
zoomtoseg(seg=c(170,171,157),rivers=abstreams0)

abstreams1 <- removeduplicates(rivers=abstreams0)
zoomtoseg(seg=c(166,167,154),rivers=abstreams1)

removemicrosegs  Remove Segments that are Smaller than the Connectivity Tolerance

Description

Automatically detects and removes segments with total displacement (straight-line distance between endpoints) less than the connectivity tolerance. These segments do not serve any real purpose, are bypassed in routes, and cannot be dissolved.

Usage

removemicrosegs(rivers)

Arguments

rivers The river network object to use.

Value

A new river network object with the specified segments connected (see rivernetwork)

Note

This function is called within cleanup, which is recommended in most cases.

Author(s)

Matt Tyers

See Also

line2network

Examples

data(abstreams0)
abstreams1 <- removemicrosegs(abstreams0)
removeunconnected  

*Remove Unconnected Segments*

**Description**

Detects and removes segments that are not connected to the river mouth.

**Usage**

removeunconnected(rivers)

**Arguments**

- `rivers`: The river network object to use.

**Note**

This function is called within `cleanup`, which is recommended in most cases.

**Author(s)**

Matt Tyers

**Examples**

```r
data(Koyukuk2)
Koy_subset <- trimriver(trimto=c(30,28,29,3,19,27,4), rivers=Koyukuk2)
Koy_subset <- setmouth(seg=1, vert=427, rivers=Koy_subset)
plot(Koy_subset)

Koy_subset_trim <- removeunconnected(Koy_subset)
plot(Koy_subset_trim)
```

---

**riverdensity**  

*The "riverdensity" Class*

**Description**

A class that holds density information computed from point data along a river network.

**Details**

Created by `makeriverdensity` from point data and a river network. Contains all information for plotting in `plot.riverdensity`. 
Elements

densities: Object of class "list". Each list element corresponds to a unique value of survey. Each element is itself of class "list", with each element corresponding to a segment from the associated river network. Each element is a vector of class "numeric", with values equal to the scaled densities calculated at the river network vertices stored in $densverts of the associated river network segment.

dendptverts: List of vectors of class "numeric". Each list element is a vector of the vertices of the endpoints of the subsegments considered for density calculation. Each list element corresponds to a river segment from the associated river network.

densverts: List of vectors of class "numeric". Each element is a vector of the vertices of the points of the subsegments considered for density calculation, that were used for density calculation. Each list element corresponds to a river segment from the associated river network.

pointsegs: Vector of class "numeric". Defined as the segment numbers of the point data used for density calculation.

pointverts: Vector of class "numeric". Defined as the vertex numbers of the point data used for density calculation.

survey: Vector of class "numeric" or class "character". Defined as the survey identifiers associated with the point data used for density calculation.

rivers: Object of class "rivernetwork"; see rivernetwork-class.

Author(s)

Matt Tyers

riverdirection | River Direction

Description

Calculates direction of travel between two points. Only works if river mouth (lowest point) has been specified (see setmouth).

Usage

riverdirection(startseg, endseg, startvert, endvert, rivers, flowconnected = FALSE, stopiferror = TRUE, algorithm = NULL)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>startseg</td>
<td>Segment number of the start of the route</td>
</tr>
<tr>
<td>endseg</td>
<td>Segment number of the end of the route</td>
</tr>
<tr>
<td>startvert</td>
<td>Vertex number of the start of the route</td>
</tr>
<tr>
<td>endvert</td>
<td>Vertex number of the end of the route</td>
</tr>
<tr>
<td>rivers</td>
<td>The river network object to use</td>
</tr>
</tbody>
</table>
flowconnected If TRUE, only returns direction if the two input segments are flow-connected. Defaults to FALSE.

stopiferror Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See detectroute.

algorithm Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See detectroute for more details.

Value

Direction: "up", "down", or "0" (character). Returns NA if flowconnected==TRUE and the two segments are not flow-connected.

Note

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see buildsegroutes).

Author(s)

Matt Tyers

See Also

setmouth

Examples

data(Gulk)

# Mouth must be specified
Gulk$mouth$mouth.seg <- 1
Gulk$mouth$mouth.vert <- 1

plot(x=Gulk)
riverdirection(startseg=6, endseg=3, startvert=40, endvert=40, rivers=Gulk)
riverdirectionmat

Usage

riverdirectionmat(seg, vert, rivers, logical = NULL, ID = NULL,
flowconnected = FALSE, stopiferror = TRUE, algorithm = NULL)

Arguments

seg A vector of river locations (segment component).
vert A vector of river locations (vertex component).
rivers The river network object to use
logical A boolean vector that can be used for subsetting - if used, riverdirectionmat() will only return pairwise distances in which a specified condition is met.
ID a vector of observation IDs for aid in interpreting the output table
flowconnected If TRUE, only returns direction if the input segments are flow-connected. Defaults to FALSE.
stopiferror Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See detectroute.
algorithm Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See detectroute for more details.

Value

A matrix of directions (character) with rows and columns labeled by corresponding values of ID. See riverdirection for additional information.

Note

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see buildsegroutes).

Author(s)

Matt Tyers

See Also

riverversion

Examples

data(Gulk, fakefish)

# Mouth must be specified
Gulk$mouth$mouth.seg <- 1
Gulk$mouth$mouth.vert <- 1

logi1 <- (fakefish$flight.date==as.Date("2015-11-25"))
riverdirectionmatbysurvey

River Direction Matrix of All Observations of an Individual

Description

Returns a matrix of travel direction between all observations of one unique fish.

Usage

riverdirectionmatbysurvey(indiv, unique, survey, seg, vert, rivers, 
full = TRUE, flowconnected = FALSE, stopiferror = TRUE, 
algorithm = NULL)

Arguments

- indiv: The unique identifier of the fish in question.
- unique: A vector of identifiers for each fish.
- survey: A vector of identifiers for each survey. It is recommended to use a numeric or date format (see `as.Date`) to preserve survey order.
- seg: A vector of river locations (segment component).
- vert: A vector of river locations (vertex component).
- rivers: The river network object to use.
- full: Whether to return the full matrix, with NA values for missing data (TRUE), or a the subset of rows and columns corresponding to successful observations. Defaults to TRUE.
- flowconnected: If TRUE, only returns direction if the input segments are flow-connected. Defaults to FALSE.
- stopiferror: Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See `detectroute`.
- algorithm: Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See `detectroute` for more details.

Value

A matrix of directions (character), with rows and columns defined by survey. In the resulting matrix, the element with the row identified as A and column identified as B is defined as the direction traveled from survey A to survey B. Therefore, it is likely that only the upper triangle of the matrix will be of interest.
riverdirectionseq

Description

Returns a matrix of directions traveled by unique fish between sequential surveys. The mouth (lowest point) segment and vertex must be specified (see setmouth).

Usage

riverdirectionseq(unique, survey, seg, vert, rivers, logical = NULL, flowconnected = FALSE, stopiferror = TRUE, algorithm = NULL)

Arguments

unique A vector of identifiers for each fish.
survey A vector of identifiers for each survey. It is recommended to use a numeric or date format (see \texttt{as.Date}) to preserve survey order.
seg A vector of river locations (segment component).
vert A vector of river locations (vertex component).
rivers The river network object to use.
Logical A boolean vector that can be used for subsetting - if used, \texttt{riverdirectionseq()} will only return pairwise distances in which a specified condition is met.
flowconnected If TRUE, only returns direction if the input segments are flow-connected. Defaults to FALSE.

Note

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see \texttt{buildsegroutes}).

Author(s)

Matt Tyers

See Also

riverdirection

Examples

data(Gulk, fakefish)
riverdirectionmatbysurvey(indiv=1, unique=fakefish$fish.id, survey=fakefish$flight, seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)
riverdirectionmatbysurvey(indiv=1, unique=fakefish$fish.id, survey=fakefish$flight, seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk, full=FALSE)
**stopiferror**  Whether or not to exit with an error if a route cannot be found. If this is set to `FALSE` and a route cannot be found, the function will return `NA` in the appropriate entry. Defaults to `TRUE`. See `detectroute`.

**algorithm**  Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as `NULL` (the default), the function will automatically make a selection. See `detectroute` for more details.

**Value**

A data frame of directions (character), with rows defined by unique fish and columns defined by observation increment (1 to 2, 2 to 3, etc.) See `riverdirection` for additional information.

**Note**

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see `buildsegroutes`).

**Author(s)**

Matt Tyers

**See Also**

`riverdirection`

**Examples**

data(Gulk, fakefish)

# Mouth must be specified
Gulk$mouth$mouth.seg <- 1
Gulk$mouth$mouth.vert <- 1

riverdirectionseq(unique=fakefish$fish.id, survey=fakefish$flight, seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)

riverdirectionseq(unique=fakefish$fish.id, survey=fakefish$flight.date, seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)
**Usage**

```r
riverdirectiontofrom(seg1, vert1, seg2, vert2, rivers, logical1 = NULL,
logical2 = NULL, ID1 = NULL, ID2 = NULL, flowconnected = FALSE,
stopiferror = TRUE, algorithm = NULL)
```

**Arguments**

- **seg1**: First vector of river locations (segment component). These are expressed as rows in the output matrix.
- **vert1**: First vector of river locations (vertex component). These are expressed as rows in the output matrix.
- **seg2**: Second vector of river locations (segment component). These are expressed as columns in the output matrix.
- **vert2**: Second vector of river locations (vertex component). These are expressed as columns in the output matrix.
- **rivers**: The river network object to use.
- **logical1**: A boolean vector that can be used for subsetting. If used, `riverdirectiontofrom` will only return directions in which a specified condition is met for the first dataset.
- **logical2**: A boolean vector that can be used for subsetting. If used, `riverdirectiontofrom` will only return directions in which a specified condition is met for the second dataset.
- **ID1**: A vector of observation IDs for the first dataset that will be used as row names in the output matrix.
- **ID2**: A vector of observation IDs for the second dataset that will be used as column names in the output matrix.
- **flowconnected**: If `TRUE`, only returns distance if the input segments are flow-connected. Defaults to `FALSE`.
- **stopiferror**: Whether or not to exit with an error if a route cannot be found. If this is set to `FALSE` and a route cannot be found, the function will return `NA` in the appropriate entry. Defaults to `TRUE`. See `detectroute`.
- **algorithm**: Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as `NULL` (the default), the function will automatically make a selection. See `detectroute` for more details.

**Value**

A matrix of directions (character) with rows and columns labeled by corresponding values of `ID`. See `riverdirection` for additional information.

**Note**

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see `buildsegroutes`).
Author(s)

Matt Tyers

See Also

riverdirection

Examples

data(Gulk)

streamlocs.seg <- c(1,8,11)
streamlocs.vert <- c(50,70,90)
streamlocs.ID <- c(“A”,”B”,”C”)

fish.seg <- c(1,4,9,12,14)
fish.vert <- c(10,11,12,13,14)
fish.ID <- c(“fish1”,”fish2”,”fish3”,”fish4”,”fish5”)

Gulk <- setmouth(seg=1, vert=1, rivers=Gulk)

riverdirectiontofrom(seg1=streamlocs.seg, vert1=streamlocs.vert,
seg2=fish.seg, vert2=fish.vert, rivers=Gulk,
ID1=streamlocs.ID, ID2=fish.ID)

logi1 <- streamlocs.ID==”B” | streamlocs.ID==”C”
logi2 <- fish.ID==”fish3”

riverdirectiontofrom(seg1=streamlocs.seg, vert1=streamlocs.vert,
seg2=fish.seg, vert2=fish.vert, rivers=Gulk, logical1=logi1,
logical2=logi2, ID1=streamlocs.ID, ID2=fish.ID)

---

riverdistance River Distance

Description

Calculates the total river network distance between two points on the river network, given in river locations (segment and vertex).

Usage

riverdistance(startseg = NULL, endseg = NULL, startvert, endvert, rivers,
path = NULL, map = FALSE, add = FALSE, stopiferror = TRUE,
algorithm = NULL)
**riverdistance**

**Arguments**

- `startseg`: Segment number of the start of the route
- `endseg`: Segment number of the end of the route
- `startvert`: Vertex number of the start of the route
- `endvert`: Vertex number of the end of the route
- `rivers`: The river network object to use
- `path`: (optional) The vector-format route of segment numbers can also be supplied instead of the starting and ending segments.
- `map`: Whether or not to draw a sanity-check map, showing the calculated route in entirety. Defaults to FALSE.
- `add`: If `map`==TRUE, whether to add the route drawing to an existing plot (add=TRUE) or produce a new plot (add=FALSE).
- `stopiferror`: Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, `riverdistance()` will return NA. Defaults to TRUE. See `detectroute`.
- `algorithm`: Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See `detectroute` for more details.

**Value**

Total route distance, in the units of the coordinate system used (this will likely be meters).

**Note**

If a distance lookup table (`$distlookup`) is present in the river network object, accepting NULL will bypass route detection and return distance automatically, the fastest algorithm of all. This is done automatically in `buildsegroutes`, but can be called directly using `buildlookup`.

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see `buildsegroutes`).

**Author(s)**

Matt Tyers

**Examples**

data(Gulk)
`riverdistance(startseg=6, endseg=14, startvert=100, endvert=200, rivers=Gulk)`
`riverdistance(startvert=100, endvert=200, path=c(6,3,4,10,11,14), rivers=Gulk)`
`riverdistance(startseg=6, endseg=14, startvert=100, endvert=200, rivers=Gulk, map=TRUE)`

# speed comparison:

data(abstrains)
riverdistancelist

Multiple River Distances

Description

Used to calculate a list of possible river distances, in the event of braiding. Calls routelist to detect a list of routes from one river location to another, and uses riverdistance to calculate the distances along those routes. Different routes are detected by randomly reordering the segment numbers of the input river network object, thus changing the internal hierarchy of segment selection.

Usage

riverdistancelist(startseg, endseg, startvert, endvert, rivers, reps = 100)

Arguments

- **startseg**: Segment number of the start of the route
- **endseg**: Segment number of the end of the route
- **startvert**: Vertex number of the start of the route
- **endvert**: Vertex number of the end of the route
- **rivers**: The river network object to use
- **reps**: Deprecated. Was the number of randomized reorderings to try.

Value

A list with two objects, `$routes` being a list of detected routes in ascending order by distance, and `$distances` being the respective distances along the routes detected.
Note

Since this function uses randomization, there is no guarantee that the list of routes will be comprehensive. Larger numbers of reps can be tried, but computation can be slow, particularly in the presence of a complex river network. It may be advantageous to use trimriver to create a smaller, more specific river network object to work with.

Author(s)

Matt Tyers

Examples

data(KilleyW)
plot(x=KilleyW)

Killey.dists <- riverdistancelist(startseg=1, endseg=16, startvert=100, endvert=25,
  rivers=KilleyW, reps=1000)
Killey.dists # 18 routes are detected.

# mapping the shortest route detected...
riverdistance(startvert=100, endvert=25, path=Killey.dists$routes[[1]], rivers=KilleyW, map=TRUE)

# mapping the shortest longest detected...
riverdistance(startvert=100, endvert=25, path=Killey.dists$routes[[18]], rivers=KilleyW, map=TRUE)

riverdistancemat  River Distance Matrix

Description

Returns a matrix of distances between every point and every other point of given river locations (segment and vertex), or of a subset.

Usage

riverdistancemat(seg, vert, rivers, logical = NULL, ID = NULL,
  stopiferror = TRUE, algorithm = NULL)

Arguments

seg A vector of river locations (segment component).
vert A vector of river locations (vertex component).
rivers The river network object to use.
logical A boolean vector that can be used for subsetting. If used, riverdistancemat will only return pairwise distances in which a specified condition is met.
ID a vector of observation IDs for aid in interpreting the output table
stopiferror  Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See detectroute

algorithm  Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See detectroute for more details.

Value

A matrix of distances (numeric) with rows and columns labeled by corresponding values of ID.

Note

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see buildsegroutes).

Author(s)

Matt Tyers

See Also

riverdistance

Examples

data(Gulk, fakefish)

logi1 <- (fakefish$flight.date==as.Date("2015-11-25"))

riverdistancemat(seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk, logical=logi1)
Arguments

- `indiv` The unique identifier of the fish in question.
- `unique` A vector of identifiers for each fish.
- `survey` A vector of identifiers for each survey. It is recommended to use a numeric or date format (see `as.Date`) to preserve survey order.
- `seg` A vector of river locations (segment component).
- `vert` A vector of river locations (vertex component).
- `rivers` The river network object to use.
- `full` Whether to return the full matrix, with NA values for missing data (TRUE), or the subset of rows and columns corresponding to successful observations. Defaults to TRUE.
- `stopiferror` Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See `detectroute`.
- `algorithm` Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See `detectroute` for more details.

Value

A matrix of distances (numeric), with rows and columns defined by survey.

Note

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see `buildsegroutes`).

Author(s)

Matt Tyers

See Also

- `riverdistance`

Examples

```r
data(Gulk, fakefish)
riverdistancematbysurvey(indiv=1, unique=fakefish$fish.id, survey=fakefish$flight, 
seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)

riverdistancematbysurvey(indiv=1, unique=fakefish$fish.id, survey=fakefish$flight, 
seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk, full=FALSE)
```
**riverdistanceseq**  
*River Distance Between Sequential Observations*

**Description**

Returns a matrix of distances traveled by unique fish, between sequential surveys. A plotting method is also provided for the output; see `plotseq`.

**Usage**

```r
riverdistanceseq(unique, survey, seg, vert, rivers, logical = NULL, stopiferror = TRUE, algorithm = NULL)
```

**Arguments**

- `unique` A vector of identifiers for each fish.
- `survey` A vector of identifiers for each survey. It is recommended to use a numeric or date format (see `as.Date`) to preserve survey order.
- `seg` A vector of river locations (segment component).
- `vert` A vector of river locations (vertex component).
- `rivers` The river network object to use.
- `logical` A boolean vector that can be used for subsetting. If used, `riverdistanceseq()` will only return pairwise distances in which a specified condition is met.
- `stopiferror` Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return `NA` in the appropriate entry. Defaults to TRUE. See `detectroute`.
- `algorithm` Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See `detectroute` for more details.

**Value**

A data frame of distances (numeric), with rows defined by unique fish and columns defined by observation increment (1 to 2, 2 to 3, etc.).

**Note**

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see `buildsegroutes`).

**Author(s)**

Matt Tyers
riverdistancetofrom

See Also

riverdistance, plotseq

Examples

data(Gulk, fakefish)
riverdistanceseq(unique=fakefish$fish.id, survey=fakefish$flight, seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)

seqbysurvey <- riverdistanceseq(unique=fakefish$fish.id, survey=fakefish$flight.date, seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)

seqbysurvey
plotseq(seqbysurvey)

riverdistancetofrom  River Distance Matrix between Two Datasets

Description

Returns a matrix of distances between each river location in two datasets, with one expressed as rows and the other expressed as columns.

Usage

riverdistancetofrom(seg1, vert1, seg2, vert2, rivers, logical1 = NULL, logical2 = NULL, ID1 = NULL, ID2 = NULL, stopiferror = TRUE, algorithm = NULL)

Arguments

seg1 First vector of river locations (segment component). These are expressed as rows in the output matrix.
vert1 First vector of river locations (vertex component). These are expressed as rows in the output matrix.
seg2 Second vector of river locations (segment component). These are expressed as columns in the output matrix.
vert2 Second vector of river locations (vertex component). These are expressed as columns in the output matrix.
rivers The river network object to use.
logical1 A boolean vector that can be used for subsetting. If used, riverdistancetofrom will only return distances in which a specified condition is met for the first dataset.
logical2 A boolean vector that can be used for subsetting. If used, riverdistancetofrom will only return distances in which a specified condition is met for the second dataset.
**ID1**  
a vector of observation IDs for the first dataset that will be used as row names in the output matrix.

**ID2**  
a vector of observation IDs for the second dataset that will be used as column names in the output matrix.

**stopiferror**  
Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See detectroute.

**algorithm**  
Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See detectroute for more details.

**Value**  
A matrix of distances (numeric) with rows and columns labeled by corresponding values of ID.

**Note**  
Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see buildsegroutes).

**Author(s)**  
Matt Tyers

**See Also**  
riverdistance

**Examples**

data(Gulk)

streamlocs.seg <- c(1,8,11)  
streamlocs.vert <- c(50,70,90)  
streamlocs.ID <- c("A","B","C")

fish.seg <- c(1,4,9,12,14)  
fish.vert <- c(10,11,12,13,14)  
fish.ID <- c("fish1","fish2","fish3","fish4","fish5")

riverdistancetofrom(seg1=streamlocs.seg, vert1=streamlocs.vert,  
                    seg2=fish.seg, vert2=fish.vert, rivers=Gulk, ID1=streamlocs.ID, ID2=fish.ID)

logi1 <- streamlocs.ID=="B" | streamlocs.ID=="C"  
logi2 <- fish.ID=="fish3"

riverdistancetofrom(seg1=streamlocs.seg, vert1=streamlocs.vert,  
                    seg2=fish.seg, vert2=fish.vert, rivers=Gulk, logical1=logi1, logical2=logi2,  
                    ID1=streamlocs.ID, ID2=fish.ID)
The "rivernetwork" Class

Description

A class that holds spatial coordinates for river networks, as well as network topology and attributes.

Details

Created by `line2network` from an input line shapefile. Contains all information for network distance calculation, plotting, etc. in the 'riverdist' package.

Plotting methods are described in `plot.rivernetwork`.

Elements

- sp: Object of class "SpatialLinesDataFrame" from package 'sp'; see `SpatialLinesDataFrame-class`. This is the original object as read by `readOGR`, and is preserved to maintain plotting capability.
- lines: Object of class "list". Each list element is a matrix of XY coordinates of the vertices of a single river segment.
- connections: Object of class "matrix", with "numeric" elements. Defined as a square matrix, with elements describing the type of connection detected between line segments.
  - A value of 1 in element \([i,j]\) indicates that the beginning of segment \(i\) is connected to the beginning of segment \(j\).
  - A value of 2 in element \([i,j]\) indicates that the beginning of segment \(i\) is connected to the end of segment \(j\).
  - A value of 3 in element \([i,j]\) indicates that the end of segment \(i\) is connected to the beginning of segment \(j\).
  - A value of 4 in element \([i,j]\) indicates that the end of segment \(i\) is connected to the end of segment \(j\).
  - A value of 5 in element \([i,j]\) indicates that segments \(i\) and \(j\) are connected at both beginning and end.
  - A value of 6 in element \([i,j]\) indicates that the beginning of segment \(i\) is connected to the end of segment \(j\), and the end of segment \(i\) is connected to the beginning of segment \(j\).
  - A value of NA in element \([i,j]\) indicates that segments \(i\) and \(j\) are not connected.
- lengths: Vector of class "numeric". Defined as the calculated total lengths of each river segment.
- names: Vector of class "character". Defined as the names of each river segment.
- mouth: Object of class "list", with two elements. Element `mouth.seg` gives the segment number of the mouth (lowest point) of the river network, and `mouth.vert` gives the vertex number.
- sequenced: "logical"; has value of TRUE if line vertices have been stored in upstream sequence using `sequenceverts`.
- tolerance: "numeric": the spatial tolerance that was used in determining river segment endpoint connectivity; see `line2network`, `splitsegments`.
units: "character": the spatial units detected from the input shapefile.

lineID: Object of class "data.frame" establishing the relationship between river segments as stored in the sp and lines elements, and is used for updating the sp element during river network editing in dissolve, splitsegments, sequenceverts, trimriver, and trimtopoints.

- rivID gives the list element number of each river segment in lines. This is the same number that is used for segment numbering in river locations.
- sp_line gives the corresponding list element in sp@lines.
- sp_seg gives the corresponding list element in sp@lines[[[]]]@Lines.

braided: "logical": Has value of TRUE if checkbraidedTF has detected braiding, FALSE if no braiding has been detected, and NA if braiding has not yet been checked.

cumuldist: List of class "numeric": Each element is a vector of cumulative distances along each river segment, beginning with 0.

segroutes: Object of class "list", with each element defined as a vector of class "numeric", describing the route from the mouth segment to the specific segment. This element only exists if buildsegroutes has been run, and can greatly speed up route and distance calculation.

distlookup: List of three matrices, of class "numeric" or "logical". Element [i,j] of each matrix corresponds to the route between segment i and j. The distlookup$middist matrix gives the total distance of the "middle" of each route (between the starting and ending segments), and the distlookup$starttop and distlookup$endtop matrices have value TRUE, FALSE, or NA if the segments at the beginning or end of the route are connected to the rest of the route at the top of the coordinate matrix, bottom of the coordinate matrix, or if the route is contained to just one segment, respectively.

Author(s)

Matt Tyers

riverpoints  

Draw Points from River Locations

Description

Add points to an active plot. Works like points but with river locations (segments and vertices) rather than xy coordinates.

Usage

riverpoints(seg, vert, rivers, pch = 1, col = 1, jitter = 0, ...)

Arguments

seg  
A vector of segments

vert  
A vector of vertices

rivers  
The river network object to use
routelist

point character, as a vector or single value
col point color, as a vector or single value
jitter maximum amount of random noise to add to “jitter” points if desired, so points do not overlap one another

Additional arguments for points

Author(s)
Matt Tyers

Examples
data(fakefish,Gulk)

plot(x=Gulk, xlim=c(862000,882000), ylim=c(6978000,6993000))
points(x=fakefish$y, y=fakefish$y, pch=16, col=2)
riverpoints(seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk, pch=15, col=4)

plot(x=Gulk, empty=TRUE)
with(fakefish, riverpoints(seg=seg, vert=vert, rivers=Gulk, pch=16, col=flight, jitter=1000))

routelist (Detect Multiple Routes)

Description
Called internally within riverdistancelist. Detects all possible routes from one river network segment to another, in the event of braiding.

Usage
routelist(startseg, endseg, rivers, reps = 100)

Arguments
startseg Segment number of the start of the route
endseg Segment number of the end of the route
rivers The river network object to use
reps Deprecated. Was used in a previous version using randomization.

Value
A list of vectors, each describing a route in segment numbers.
Note

The previous version of this function returned many possible routes using randomization - this algorithm now computes all possible routes.

Author(s)

Matt Tyers

Examples

data(KilleyW)
plot(x=KilleyW)

routelist(startseg=1, endseg=16, rivers=KilleyW, reps=1000)

sequenceverts

Store Vertices in Ascending Sequence

Description

Rearranges the vertices of a river network object so that vertices are stored sequentially moving up river for all segments (coordinates [1.] are the bottom of each segment).

Usage

sequenceverts(rivers)

Arguments

rivers The river network object to use

Value

A new river network object (see rivernetwork)

Note

Even without calling sequenceverts, the vertices will be stored sequentially - either moving up river or down for a given segment. What sequenceverts() adds is a standardized direction.

Currently, no function in package 'riverdist' requires the vertices to be stored sequentially.

Author(s)

Matt Tyers

See Also

line2network
**setmouth**

**Examples**

```r
data(Gulk)  
Gulk <- setmouth(seg = 1, vert = 1, rivers = Gulk)  
str(Gulk)
```

```r
Gulk.dir <- sequenceverts(rivers = Gulk)  
str(Gulk.dir)
```

---

### Description

Provides a user-friendly way of specifying the segment and vertex of the mouth (lowest point) of a river network object.

### Usage

```r
setmouth(seg, vert, rivers)
```

### Arguments

- **seg**: The segment number to store for the mouth
- **vert**: The vertex number to store for the mouth
- **rivers**: The river network object to use

### Value

A new river network object (see `rivernetwork`)

### Note

The mouth segment and vertex can also be specified using direct assignment to the `mouth$seg` and `mouth$vert` components of the river network object.

This function is called within `cleanup`, which is recommended in most cases.

### Author(s)

Matt Tyers

### See Also

- `line2network`
Examples

data(Gulk)

# say we know that segment 1 is the lowest segment in this river network, but we don't know
# which end is the mouth.
showends(seg=1, rivers=Gulk)

# this means that the mouth is row 1, so we can specify this:
Gulk <- setmouth(seg=1, vert=1, rivers=Gulk)

showends

Identify Vertex Coordinates of Segment Endpoints

Description

Identifies the vertex coordinates (row numbers) of the endpoints of a given segment. The main
purpose is determining which of the endpoints is the mouth (or lowest point) of the river system.

Usage

showends(seg, rivers)

Arguments

seg The segment (number) to check
rivers The river network object to use

Note

This function is called within cleanup, which is recommended in most cases.

Author(s)

Matt Tyers

Examples

data(Gulk)

# say we know that segment 1 is the lowest segment in this river network, but we don't know
# which end is the mouth.
showends(seg=1, rivers=Gulk)

# this means that the mouth is row 1, so we can specify this:
Gulk <- setmouth(seg=1, vert=1, rivers=Gulk)
smallset

Dataset: Smallset

Description
A small set of observations of fakefish on the Gulkana River and its tributaries.

Usage
data(smallset)

Format
A data frame

Details
- x. X-coordinate of observation (Alaska Albers Equal Area). Note that the locations do not align with the river network object.
- y. Y-coordinate of observation
- seg. River segment
- vert. River vertex
- fish.id. Numeric identifier for each fish (individual fish were observed more than once)
- flight. Numeric identifier for each telemetry flight

See Also
Gulk

splitsegmentat

Split a Segment at a Specified Vertex

Description
Splits a segment at a specified vertex, creating two new segments.

Usage
splitsegmentat(seg, vert, rivers)

Arguments
- seg    The segment to split
- vert   The vertex to split it at
- rivers The river network object to use
**Value**

A new, updated river network object

**Author(s)**

Matt Tyers

**See Also**

line2network

**Examples**

```r
data(Gulk)
plot(x=Gulk)

Gulk2 <- splitsegmentat(seg=1, vert=400, rivers=Gulk)
plot(x=Gulk2)
```

---

**splitsegments**  
*Split Segments by Endpoint Proximity*

**Description**

Detects cases in which segments should be split to establish appropriate topology, and splits them. Specifically, it looks for segment endpoints intersecting (or within a tolerance of) another segment. It then splits the intersected segment at the point where the endpoint of the other segment breaks it.

**Usage**

```r
splitsegments(rivers, tolerance = NULL, splitthese = NULL, splitthemat = NULL, one2one = FALSE, append = FALSE)
```

**Arguments**

- **rivers**: The river network object to use
- **tolerance**: The spatial snapping tolerance to use for detecting intersection. If a NULL value is used (default), it will default to the tolerance that was used in river network creation in line2network.
- **splitthese**: An optional vector of target segments to split. If this argument is used, only these segments will be split. If the default (NULL) is accepted, all segments will be used.
- **splitthemat**: An optional vector of segments (endpoints) to use for splitting. If this argument is used, segments will only be split at the endpoints of these segments. If the default (NULL) is accepted, all segments will be used.
- **one2one**: Logical, indicating a one-to-one correspondence between arguments splitthese and splitthemat. Defaults to FALSE.
append Logical, indicating how to organize the output river network. If TRUE, appends newly-created segments to the end of $lines, rather than retaining original line ordering. This may be useful in retaining original line ID. Defaults to FALSE.

Value
A new, updated river network object

Note
This function is called within cleanup, which is recommended in most cases.

Author(s)
Matt Tyers

See Also
line2network

Examples
```r
data(Koyukuk1)
 topologydots(rivers=Koyukuk1)
# Segments 7, 8, 13, and 16 need to be split so topologies will work.
# Since endpoints are not in the same place, they are not detected as being connected.
plot(x=Koyukuk1)

Koyukuk1split <- splitsegments(rivers=Koyukuk1)
topologydots(rivers=Koyukuk1split)
plot(x=Koyukuk1split)

# if only segment 17 were to be split in three places
plot(x=splitsegments(rivers=Koyukuk1, splitthese=c(7,7,7), splittthemat=c(14,5,12)))

# if only segment 16 were to be split, showing behavior of append=
plot(x=splitsegments(rivers=Koyukuk1, splitthese=c(7,7,7), splittthemat=c(14,5,12), append=TRUE))
```

topologydots Check Connectivity of a River Network Object

Description
Produces a graphical check of the connectivity of a river network object. It produces a plot of the river network object, and overlays red dots at non-connected endpoints and green dots at connected endpoints.
Usage
topologydots(rivers, add = FALSE, ...)

Arguments
  rivers       The river network object to check
  add          Whether call a new plot (FALSE) or add dots to an existing plot (TRUE). Defaults to FALSE.
  ...          Additional plotting arguments (see par)

Author(s)
  Matt Tyers

See Also
  line2network

Examples
data(Gulk)
topologydots(rivers=Gulk)

trimriver       Trim a River Network Object to Specified Segments

Description
  Removes line segments from a river network object. User can specify which segments to remove (trim) or which segments to keep (trimto).

Usage
  trimriver(trim = NULL, trimto = NULL, rivers)

Arguments
  trim         Vector of line segments to remove
  trimto       Vector of line segments to keep
  rivers       The river network object

Value
  A new river network object
**Note**

Specifying segments in both trim and trimto arguments will result in an error.

**Author(s)**

Matt Tyers

**See Also**

`line2network`

**Examples**

```r
data(Kenai)
plot(x=Kenai)

Kenai.trim <- trimriver(trim=c(46,32,115,174,169,114,124,142,80), rivers=Kenai)
plot(x=Kenai.trim)

Kenai.trim2 <- trimriver(trimto=c(20,57,118,183,45,162,39,98,19), rivers=Kenai)
plot(x=Kenai.trim2)
```

---

**trimtopoints**

*Trim a River Network to a Set of X-Y Coordinates*

**Description**

Removes line segments from a river network object that are not adjacent to a set of point data, given in X-Y coordinates.

**Usage**

```r
trimtopoints(x, y, rivers, method = "snap", dist = NULL)
```

**Arguments**

- `x`: Vector of x-coordinates of point data to buffer around
- `y`: Vector of y-coordinates of point data to buffer around
- `rivers`: The river network object to use
- `method`: Three methods are available. If "snap" is specified (the default), only the closest segment to each point is retained. If "snaproute" is specified, segments are also retained that will maintain total connectivity in the resulting river network. If "buffer" is specified, all segments with endpoints or midpoints within `dist` units of the input locations are retained.
- `dist`: Distance to use for buffering, if `method=\"buffer\"`. If this is not specified, the maximum spread in the x- and y- direction will be used.
Value

A new river network object (see `rivernetwork`)

Note

If `method="buffer"`, only distances to segment endpoints and midpoints are checked, and still only whole segments are removed.

Author(s)

Matt Tyers

See Also

`line2network`

Examples

data(Koyukuk2)
x <- c(139241.0, 139416.1, 124600.1, 122226.8)
y <- c(1917577, 1913864, 1898723, 1898792)

plot(x=Koyukuk2)
points(x, y, pch=15, col=4)
legend(par("usr")[1], par("usr")[4], legend="points to buffer around", pch=15, col=4, cex=.6)

Koyukuk2.buf1 <- trimtopoints(x, y, rivers=Koyukuk2, method="snap")
plot(x=Koyukuk2.buf1)
points(x, y, pch=15, col=4)

Koyukuk2.buf2 <- trimtopoints(x, y, rivers=Koyukuk2, method="snaproute")
plot(x=Koyukuk2.buf2)
points(x, y, pch=15, col=4)

Koyukuk2.buf3 <- trimtopoints(x, y, rivers=Koyukuk2, method="buffer", dist=1000)
plot(x=Koyukuk2.buf3)
points(x, y, pch=15, col=4)

upstream

**Upstream River Distance**

Description

Calculates river network distances as +/-, defined as upriver/downriver.

Specifying `net=TRUE` will compute net upriver distance (3 river km down a tributary and then 15 river km up the mainstem will mean 12 rkm net. Otherwise the function will return 18 rkm upriver travel.)

The mouth (lowest point) segment and vertex must be specified (see `setmouth`).
**Usage**

```r
upstream(startseg, endseg, startvert, endvert, rivers, flowconnected = FALSE,
    net = FALSE, stopiferror = TRUE, algorithm = NULL)
```

**Arguments**

- `startseg` Segment number of the start of the route
- `endseg` Segment number of the end of the route
- `startvert` Vertex number of the start of the route
- `endvert` Vertex number of the end of the route
- `rivers` The river network object to use
- `flowconnected` If TRUE, only returns distance if the two input segments are flow-connected. Defaults to FALSE.
- `net` Whether to calculate net distance (net=TRUE) or total distance (net=FALSE)
- `stopiferror` Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See `detectroute`.
- `algorithm` Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See `detectroute` for more details.

**Value**

Upstream distance (numeric). Returns NA if `flowconnected` has value TRUE and the two segments are not flow-connected.

**Note**

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see `buildsegroutes`).

**Author(s)**

Matt Tyers

**See Also**

- `setmouth`

**Examples**

```r
data(Gulk)

# Mouth must be specified
Gulk$mouth$mouth.seg <- 1
Gulk$mouth$mouth.vert <- 1
```
plot(x=Gulk)
riverpoints(seg=c(6,4), vert=c(140,140), pch=16, col=2, rivers=Gulk)
upstream(startseg=6, endseg=4, startvert=140, endvert=40, rivers=Gulk, net=TRUE)
upstream(startseg=6, endseg=4, startvert=140, endvert=40, rivers=Gulk, net=FALSE)
upstream(startseg=6, endseg=4, startvert=140, endvert=40, rivers=Gulk, flowconnected=TRUE)

Description
Returns a matrix of upstream distance between every point and every other point of given river locations (segment and vertex), or of a subset. The mouth (lowest point) segment and vertex must be specified (see setmouth).

Usage
upstreammat(seg, vert, rivers, logical = NULL, ID = NULL, 
flowconnected = FALSE, net = FALSE, stopiferror = TRUE, 
algorithm = NULL)

Arguments
seg A vector of river locations (segment component).
vert A vector of river locations (vertex component).
rivers The river network object to use.
logical A boolean vector that can be used for subsetting - if used, riverdirectionseq() will only return pairwise distances in which a specified condition is met.
ID a vector of observation IDs for aid in interpreting the output table
flowconnected If TRUE, only returns distance if the input segments are flow-connected. Defaults to FALSE.
net Whether to calculate net upstream distance (net=TRUE) or total distance (net=FALSE, default). See upstream.
stopiferror Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See detectroute.
algorithm Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See detectroute for more details.

Value
A matrix of upstream distances (numeric) with rows and columns labeled by corresponding values of ID. See upstream for additional information.
Note

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see `buildsegroutes`).

Author(s)

Matt Tyers

See Also

upstream

Examples

```r
data(Gulk, fakefish)

# Mouth must be specified
Gulk$mouth$mouth.seg <- 1
Gulk$mouth$mouth.vert <- 1

logi1 <- (fakefish$flight.date==as.Date("2015-11-25"))

upstreammat(seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk, logical=logi1)
```

upstreammatbysurvey  Upstream Distance Matrix of All Observations of an Individual

Description

Returns a matrix of upstream travel distance between all observations of one unique fish.

Usage

```r
upstreammatbysurvey(indiv, unique, survey, seg, vert, rivers, full = TRUE,
flowconnected = FALSE, net = FALSE, stopiferror = TRUE,
algorithm = NULL)
```

Arguments

- `indiv` The unique identifier of the fish in question.
- `unique` A vector of identifiers for each fish.
- `survey` A vector of identifiers for each survey. It is recommended to use a numeric or date format (see `as.Date`) to preserve survey order.
- `seg` A vector of river locations (segment component).
- `vert` A vector of river locations (vertex component).
- `rivers` The river network object to use.
full Whether to return the full matrix, with NA values for missing data (TRUE), or a the subset of rows and columns corresponding to successful observations. Defaults to TRUE.

flowconnected If TRUE, only returns direction if the input segments are flow-connected. Defaults to FALSE.

net Whether to calculate net upstream distance (net=TRUE) or total distance (net=FALSE, default).

stopiferror Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See detectroute.

algorithm Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See detectroute for more details.

Value

A matrix of upstream distances (numeric), with rows and columns defined by survey. In the resulting matrix, the element with the row identified as A and column identified as B is defined as the upstream distance traveled from survey A to survey B. Therefore, it is likely that only the upper triangle of the matrix will be of interest.

Note

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see buildsegroutes).

Author(s)

Matt Tyers

See Also

upstream

Examples

data(Gulk, fakefish)
upstreammatbysurvey(indiv=1, unique=fakefish$fish.id, survey=fakefish$flight, seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk)

upstreammatbysurvey(indiv=1, unique=fakefish$fish.id, survey=fakefish$flight, seg=fakefish$seg, vert=fakefish$vert, rivers=Gulk, full=FALSE)
upstreamseq

**Upstream Distance Between Sequential Observations**

**Description**

Returns a matrix of distance with direction by unique fish between sequential surveys. The mouth (lowest point) segment and vertex must be specified (see `setmouth`). A plotting method is provided for the output; see `plotseq`.

**Usage**

```r
upstreamseq(unique, survey, seg, vert, rivers, logical = NULL, 
flowconnected = FALSE, net = FALSE, stopiferror = TRUE, 
algorithm = NULL)
```

**Arguments**

- `unique`: A vector of identifiers for each fish.
- `survey`: A vector of identifiers for each survey. It is recommended to use a numeric or date format (see `as.Date`) to preserve survey order.
- `seg`: A vector of river locations (segment component).
- `vert`: A vector of river locations (vertex component).
- `rivers`: The river network object to use.
- `logical`: A boolean vector that can be used for subsetting - if used, `upstreamseq()` will only return pairwise distances in which a specified condition is met.
- `flowconnected`: If TRUE, only returns distance if the input segments are flow-connected. Defaults to FALSE.
- `net`: Whether to calculate net upstream distance (net=TRUE) or total distance (net=FALSE, default).
- `stopiferror`: Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See `detectroute`.
- `algorithm`: Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See `detectroute` for more details.

**Value**

A data frame of upstream distances (numeric), with rows defined by unique fish and columns defined by observation increment (1 to 2, 2 to 3, etc.) See `upstream` for additional information.
Note

Returns either net upstream distance (net=TRUE) or total distance (net=FALSE, default). See upstream.

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see buildsegroutes).

Author(s)

Matt Tyers

See Also

upstream, plotseq

Examples

data(Gulk, fakefish)

# Mouth must be specified
Gulk$mouth$seg <- 1
Gulk$mouth$vert <- 1

upstreamseq(unique=fakefish$fish.id, survey=fakefish$flight, seg=fakefish$seg, 
vert=fakefish$vert, rivers=Gulk)

seqbysurvey <- upstreamseq(unique=fakefish$fish.id, survey=fakefish$flight.date, seg=fakefish$seg, 
vert=fakefish$vert, rivers=Gulk)

seqbysurvey

plotseq(seqbysurvey)

Description

Returns a matrix of upstream distances between each river location in two datasets, with one expressed as rows and the other expressed as columns.

Usage

upstreamtofrom(seg1, vert1, seg2, vert2, rivers, logical1 = NULL, 
logical2 = NULL, ID1 = NULL, ID2 = NULL, net = FALSE, 
flowconnected = FALSE, stopiferror = TRUE, algorithm = NULL)
Arguments

- **seg1**: First vector of river locations (segment component). These are expressed as rows in the output matrix.
- **vert1**: First vector of river locations (vertex component). These are expressed as rows in the output matrix.
- **seg2**: Second vector of river locations (segment component). These are expressed as columns in the output matrix.
- **vert2**: Second vector of river locations (vertex component). These are expressed as columns in the output matrix.
- **rivers**: The river network object to use.
- **logical1**: A boolean vector that can be used for subsetting. If used, `upstreamtofrom` will only return upstream distances in which a specified condition is met for the first dataset.
- **logical2**: A boolean vector that can be used for subsetting. If used, `upstreamtofrom` will only return upstream distances in which a specified condition is met for the second dataset.
- **ID1**: A vector of observation IDs for the first dataset that will be used as row names in the output matrix.
- **ID2**: A vector of observation IDs for the second dataset that will be used as column names in the output matrix.
- **net**: Whether to calculate net upstream distance (TRUE) or signed total distance (FALSE). See `upstream`.
- **flowconnected**: If TRUE, only returns distance if the input segments are flow-connected. Defaults to FALSE.
- **stopiferror**: Whether or not to exit with an error if a route cannot be found. If this is set to FALSE and a route cannot be found, the function will return NA in the appropriate entry. Defaults to TRUE. See `detectroute`.
- **algorithm**: Which route detection algorithm to use ("Dijkstra", "sequential", or "segroutes"). If left as NULL (the default), the function will automatically make a selection. See `detectroute` for more details.

Value

A matrix of upstream distances (numeric) with rows and columns labeled by corresponding values of ID. See `upstream` for additional information.

Note

Building routes from the river mouth to each river network segment and/or distance lookup tables will greatly reduce computation time (see `buildsegroutes`).

Author(s)

Matt Tyers
See Also

upstream

Examples

data(Gulk)

streamlocs.sseg <- c(1,8,11)
streamlocs.vert <- c(50,70,90)
streamlocs.ID <- c("A","B","C")

fish.sseg <- c(1,4,9,12,14)
fish.vert <- c(10,11,12,13,14)
fish.ID <- c("fish1","fish2","fish3","fish4","fish5")

Gulk <- setmouth(seg=1, vert=1, rivers=Gulk)

upstreamtofrom(seg1=streamlocs.sseg, vert1=streamlocs.vert,
                seg2=fish.sseg, vert2=fish.vert, rivers=Gulk,
                ID1=streamlocs.ID, ID2=fish.ID)

logi1 <- streamlocs.ID=="B" | streamlocs.ID=="C"
logi2 <- fish.ID=="fish3"

upstreamtofrom(seg1=streamlocs.sseg, vert1=streamlocs.vert,
                seg2=fish.sseg, vert2=fish.vert, rivers=Gulk, logical1=logi1,
                logical2=logi2, ID1=streamlocs.ID, ID2=fish.ID)

whoconnected  

Check Which Segments are Connected to a Given Segment.

Description

Returns which segments are connected to a specified segment within a river network. It may be useful for error checking.

Usage

whoconnected(seg, rivers)

Arguments

seg The segment to check
rivers The river network object it belongs to

Value

A vector of segment numbers
Author(s)

Matt Tyers

Examples

data(Gulk)
plot(Gulk)
whoconnected(seg=4, rivers=Gulk)

---

xy2segvert Convert XY Coordinates to River Locations

Description

This function determines the closest vertex in the river network to each point of XY data and returns a list of river locations, defined as segment numbers and vertex numbers.

Usage

xy2segvert(x, y, rivers)

Arguments

x A vector of x-coordinates to transform
y A vector of y-coordinates to transform
rivers The river network object to use

Value

A data frame of river locations, with segment numbers in $seg, vertex numbers in $vert, and the snapping distance for each point in $snapdist.

Note

Conversion to river locations is only valid if the input XY coordinates and river network are in the same projected coordinate system. Point data in geographic coordinates can be projected using project in package 'rgdal', and an example is shown below.

Author(s)

Matt Tyers
Examples

```r
data(Gulk, fakefish)
head(fakefish)

fakefish.riv <- xy2segvert(x=fakefish$x, y=fakefish$y, rivers=Gulk)
head(fakefish.riv)

plot(x=Gulk, xlim=c(862000,882000), ylim=c(6978000,6993000))
points(fakefish$x, fakefish$y, pch=16, col=2)
riverpoints(seg=fakefish.riv$seg, vert=fakefish.riv$vert, rivers=Gulk, pch=15, col=4)
```

```r
## converting a matrix of points stored in long-lat to Alaska Albers Equal Area:
data(line98, Kenai1)
head(line98)  # note that coordinates are stored in long-lat, NOT lat-long

library(rgdal)
line98albers <- project(line98, proj="+proj=aea +lat_1=55 +lat_2=65 +lat_0=50 +lon_0=-154 +x_0=0 +y_0=0 +datum=NAD83 +units=m +no_defs +ellps=GRS80 +towgs84=0,0,0")
head(line98albers)
zoomtoseg(seg=c(162,19), rivers=Kenai1)
points(line98albers)
```

---

**zoomtoseg**

---

**Description**

Calls `plot.rivernetwork` and automatically zooms to a specified segment or vector of segments. Not intended for any real mapping - just investigating and error checking.

**Usage**

```r
zoomtoseg(seg, rivers, ...)
```

**Arguments**

- `seg` A segment or vector of segments to zoom to
- `rivers` The river network object to use
- `...` Additional plotting arguments (see `par`)

**Author(s)**

Matt Tyers
Examples

data(Kenai3)
plot(x=Kenai3)

# checking out a particularly messy region...
zoomtoseg(c(110,63), rivers=Kenai3)
Index

* Topic **datasets**
  - abstreams, 5
  - abstreams0, 6
  - fakefish, 18
  - fakefish_density, 19
  - Gulk, 20
  - Kenai1, 25
  - Kenai2, 25
  - Kenai3, 26
  - KilleyW, 28
  - Koyukuk0, 28
  - Koyukuk1, 29
  - Koyukuk2, 29
  - line98, 31
  - smallset, 73

  abstreams, 5
  abstreams0, 6
  addcumuldist, 6
  addverts, 7, 13
  as.Date, 34, 37, 46, 54, 55, 63, 64, 81, 83

  buildlookup, 8, 9, 27, 32, 59
  buildsegroutes, 8, 9, 13, 17, 22, 27, 32, 35–37, 44, 52, 53, 55–57, 59, 62–64, 66, 68, 79, 81, 82, 84, 85

  calculateconnections, 10
  checkbraided, 11
  checkbraidedTF, 12, 13, 68
  cleanup, 7–9, 11, 12, 13, 15, 18, 49, 50, 71, 72, 75
  cleanup_verts, 14
  connectsegs, 13, 15

  detectroute, 16, 21, 24, 35–37, 52–54, 56, 57, 59, 62–64, 66, 79, 80, 82, 83, 85
  dissolve, 13, 17, 68

  fakefish, 18
  fakefish_density, 19

  Gulk, 19, 20, 73

  highlightseg, 20
  homerange, 21, 22, 23, 39, 40, 43, 44
  homerange-class, 21, 22
  homerangeoverlap, 22, 23, 40, 44

  isflowconnected, 24

  Kenai1, 25, 26
  Kenai2, 25, 26
  Kenai3, 25, 26, 28
  kfunc, 26
  KilleyW, 28
  Koyukuk0, 28
  Koyukuk1, 29, 29
  Koyukuk2, 29, 29

  line2network, 29, 30, 67, 70, 71, 74–78
  line98, 31

  makeriverdensity, 31, 40–42, 45, 50
  mapbyname, 33
  mapriver (plot.rivernetwork), 42
  matbysurveylist, 34, 44
  mouthdist, 35
  mouthdistbysurvey, 36, 46

  par, 20, 33, 43, 76, 88
  pdist, 38
  pdisttot, 38
  plot, 75
  plot.homerange, 21–23, 39, 44
  plot.riverdensity, 31, 32, 40, 45, 50
  plot.rivernetwork, 39, 42, 43, 67, 88
  plot.homerangeoverlap, 22, 23, 40, 43
  plotmatbysurveylist, 34, 35, 44
  plotriverdensity, 19
  plotriverdensity (plot.riverdensity), 40
  plotriverdensitypoints, 32, 42, 45
  plotseq, 36, 37, 46, 64, 65, 83, 84
points, 68, 69
pointshp2segvert, 47
project, 87
readOGR, 30, 67
removeduplicates, 13, 48
removemicrosegs, 13, 49
removeunconnected, 13, 50
riverdensity, 19, 50
riverdensity-class, 19, 32
riverdensity-class (riverdensity), 50
riverdirection, 24, 35, 51, 53, 55–58
riverdirectionmat, 52
riverdirectionmatbysurvey, 34, 35, 54
riverdirectionseq, 55
riverdirectiontofrom, 56
riverdist (riverdist-package), 3
riverdist-package, 3
riverdistance, 16, 35, 58, 60, 62, 63, 65, 66
riverdistancelist, 60, 69
riverdistancemat, 61
riverdistancematbysurvey, 34, 35, 62
riverdistanceseq, 46, 64
riverdistancetofrom, 65
rivernetwork, 5–9, 13–15, 20, 25, 26, 28, 29, 48, 49, 67, 70, 71, 78
rivernetwork-class, 6, 10, 30, 42, 51
rivernetwork-class (rivernetwork), 67
riverpoints, 68
routelist, 60, 69
sequenceverts, 67, 68, 70
setmouth, 9, 13, 24, 35, 36, 51, 52, 55, 71, 78–80, 83
showends, 72
smallset, 73
SpatialLinesDataFrame-class, 67
splitsegmentat, 73
splitsegments, 13, 67, 68, 74
spTransform, 30
topologydots, 75
trimriver, 61, 68, 76
trimtopoints, 68, 77
upstream, 24, 35, 45, 78, 80–86
upstreammat, 80
upstreammatbysurvey, 34, 35, 45, 81
upstreamseq, 46, 83
whoconnected, 86
xy2segvert, 87
zoomtoseg, 88