Package 'rpostgisLT'

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Title Managing Animal Movement Data with 'PostGIS' and R

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Description The 'rpostgisLT' package develops the integration of R and 'PostGIS' for managing movement trajectories. The focus is on streamlining the work flow for biologists to store and process animal trajectories in 'PostGIS' and analyze them in R, thus utilizing the strengths of both software. The package relies on 'Itraj' objects from the R package 'adehabitatLT', and provides the analogous 'pgtraj' data structure in 'PostGIS', with all functions to create and manage 'pgtraj' data, and convert from and to both formats ('pgtraj' in 'PostGIS', 'Itraj' in R).
SystemRequirements PostgreSQL with PostGIS extension
Depends R (>= 3.3.0), DBI, RPostgreSQL, rpostgis (>= 1.0.2), adehabitatLT (>= 0.3.12)
License GPL (>= 3)
URL https://github.com/mablab/rpostgisLT
BugReports https://github.com/mablab/rpostgisLT/issues
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VignetteBuilder knitr
R topics documented:
as_pgtraj

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Description

as_pgtraj

as_pgtraj populates a traj schema from the data provided in relocations_table. If the provided schema doesn't exist, it will be created. On successful data input, as_pgtraj creates two database views for each pgtraj. These views are named parameters_<pgtraj_name>, step_geometry_<pgtraj_name> and described more in detail in the package vignette.

Imports location data from a database table into a 'traj' schema.

The time zone of the pgtraj is set to the local time zone of the user.

Usage

```
as_pgtraj(conn, relocations_table, schema = "traj", pgtrajs = "pgtraj",
animals = "animal", bursts = NULL, relocations, timestamps = NULL,
rids = "rid", srid = NULL, note = NULL, clauses = NULL,
info_cols = NULL, info_table = NULL, info_rids = NULL)
```

Arguments

conn	Connection object created with RPostgreSQL
relocations_ta	ble
	String. Name of the schema and table that stores the relocations, e.g. c("schema", "relocations")
schema	String. Name of the schema that stores or will store the pgtraj data model (Default = "traj").
pgtrajs	String. Name of the pgtraj or name of the field that stores the pgtraj names.
animals	String. Name of the animal or name of the field that stores the animal names.
bursts	String. (Optional) name of the burst or name of the field that stores the burst names. If not given, each animal will have one burst.
relocations	String. Name of the field that contains the relocations in relocations_table. Relocations can be provided either as X,Y coordinates or PostGIS geometry. In both cases all relocations in the 'relocations_table' have to have the same projection.
timestamps	String. Name of the field in relocations_table that contains the timestamps. If NULL, Type I trajectory is assumed.
rids	String. Name of the field in relocations_table that contains the numeric IDs of relocations. If timestamps = NULL, relocations will be sorted by the ascending numeric IDs in this field.
srid	Integer. Optional SRID (spatial reference ID) of (x,y) coordinates provided for relocations. Ignored if relocations is a geometry type.
note	String. Comment on the pgtraj. The comment is only used in the database and not transferred into an ltraj.

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clauses	character, additional SQL to append to modify data selected from relocations_table. Must begin with WHERE, and cannot contain ORDER BY or LIMIT clauses.
info_cols	String. Optional character vector of column names of additional information on relocations (replicating "infolocs" from the adehabitatLT object ltraj).
info_table	Character vector of c("schema", "table") holding the info_cols. If info_cols are in relocations_table, leave NULL.
info_rids	String. Column name of unique integer ID in info_table to join with rids. If info_cols are in relocations_table, leave NULL.

Details

Opening and closing connections have to be done manually by the user. However, the function checks if the provided connection is still valid.

Value

TRUE on success

Author(s)

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References

https://cran.r-project.org/web/packages/adehabitatLT/vignettes/adehabitatLT.pdf

See Also

Section on traj data model in the package vignette.

Examples

```
## Not run:
as_pgtraj(conn,
        relocations_table = c("example_data", "relocations_plus"),
        schema = "traj_t4",
        pgtrajs = "id",
        animals = "animal",
        bursts = "burst",
        relocations = "geom",
        timestamps = "time",
        rids = "gid")
## End(Not run)
## Not run:
as_pgtraj(conn,
        relocations_table = c("example_data","relocations_plus"),
        schema = "traj_t4",
        pgtrajs = "id",
        animals = "animal",
        bursts = "burst",
        relocations = c("x","y"),
        timestamps = "time",
        rids = "gid")
```

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```
## End(Not run)
```

ltraj2pgtraj

Export ltraj object from R into database.

Description

ltraj2pgtraj creates a new traj schema or uses an existing one and exports an ltraj to the database. The time zone and projection information stored in the ltraj is transferred to the database.

Usage

```
ltraj2pgtraj(conn, ltraj, schema = "traj", pgtraj = NULL, note = NULL,
  overwrite = FALSE, infolocs = TRUE)
```

Arguments

conn	A connection object.
ltraj	An object of class ltraj.
schema	Character. Name of the schema that stores or will store the pgtraj data model.
pgtraj	Character. Name of the new pgtraj. Defaults to the name of the provided $ltraj$.
note	Character. A note that will be stored with the pgtraj in the database.
overwrite	Logical. Use if a pgtraj with the same name as the provided ltraj already exists in the database: If TRUE, the existing pgtraj is deleted and the provided ltraj is inserted. If FALSE, the function exits. Note that overwrite requires an exact match among the pgtraj names otherwise it is ignored.
infolocs	Logical. Whether to write infolocs to database.

Value

TRUE on success.

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See Also

as_pgtraj to create a pgtraj with data already stored in the database.

Examples

```
## Not run:
    # create pgtraj from ltraj "ibex" in schema "traj_t2"
    ltraj2pgtraj(conn, ibex, "traj_t2")
## End(Not run)
```

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Description

pgtraj2ltraj imports a single pgtraj from a database into an ltraj object.

Usage

```
pgtraj2ltraj(conn, pgtraj, schema = "traj")
```

Arguments

conn Connection object created with RPostgreSQL

pgtraj String. Name of the pgtraj.

schema String. Name of the schema that stores or will store the pgtraj data model.

Value

an ltraj object

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Examples

```
## Not run:
    # create ltraj from pgtraj named "ibex" in schema "traj_t2"
    ibex<-pgtraj2ltraj(conn, "ibex", "traj_t2")
## End(Not run)</pre>
```

pgtrajDrop

Delete a pgtraj from a traj schema.

Description

pgtrajDrop deletes a pgtraj from a traj schema.

Usage

```
pgtrajDrop(conn, pgtraj, schema = "traj", full_clean = TRUE)
```

Arguments

conn Connection object created with RPostgreSQL

pgtraj String. Name of the pgtraj.

schema String. Name of the schema that stores the traj data model.

full_clean String. Whether to delete all unused rows in all tables associated with the pgtraj

(and any other previously deleted pgtrajs). Recommended, but may take a long

time to run in schemas with many large pgtraj's.

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Value

TRUE on success

Author(s)

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Examples

```
## Not run:
    # drop "ibex" pgtraj in schema "traj"
    pgtrajDrop(conn, "ibex")
## End(Not run)
```

pgtrajSchema

Check pgtraj schema.

Description

Checks if the provided schema is a valid pgtraj schema, and creates one if it does not exist.

Usage

```
pgtrajSchema(conn, schema = "traj")
```

Arguments

conn Connection object created with RPostgreSQL.

schema Character string. Name of the schema that stores or will store the pgtraj data

model.

Details

Creates a schema to store pgtrajs in the database by calling a SQL script from ./sql/traj_schema.sql. The schema name defaults to traj. If a schema with the provided name already exists in the database, it checks if it contains all the required tables. The function does not attempt to repair the schema if all pgtraj tables are not present (e.g. because some were manually deleted). In this case, a new pgtraj schema needs to be created, or the existing schema needs to be deleted and recreated.

The function has its own standalone transaction control.

Value

TRUE if the schema exists (whether it was already available or was successfully created).

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Examples

```
## Not run:
    # Check (or create) pgtraj schema with name "traj_1"
    pgtrajSchema(conn, "traj_1")
## End(Not run)
```

pgtrajVacuum

pgtrajVacuum VACUUM on a pgtraj schema.

Description

Performs a VACUUM (garbage-collect and optionally analyze) on all the tables of a traj schema.

Usage

```
pgtrajVacuum(conn, schema = "traj", full = FALSE, verbose = FALSE,
  analyze = TRUE)
```

Arguments

conn	Connection object created with RPostgreSQL
schema	String. Name of the schema that stores or will store the pgtraj data model.
full	Logical. Whether to perform a "full" vacuum, which can reclaim more space, but takes much longer and exclusively locks the table.
verbose	Logical. Whether to print a detailed vacuum activity report for each table.
analyze	Logical. Whether to update statistics used by the planner to determine the most efficient way to execute a query (default to TRUE).

Value

TRUE on success.

Author(s)

```
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```

Examples

```
## Not run:
    # Vacuum analyze all tables in pgtraj schema with default name "traj"
    pgtrajVacuum(conn)
## End(Not run)
```

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rpostgisLT

Integration of ltraj (adehabitatLT) and pgtraj (PostGIS).

Description

rpostgisLT

Details

The 'rpostgisLT' package develops the integration of R and PostGIS for managing movement trajectories. The focus is on streamlining the workflow for biologists to store and process animal trajectories in PostGIS and analyze them in R, thus utilizing the strengths of both software. The package relies on 'ltraj' objects from the R package 'adehabitatLT', and provides the analogous 'pgtraj' data structure in PostGIS, with all functions to create and manage 'pgtraj' data, and convert from and to both format ('pgtraj' in PostGIS, 'ltraj' in R). For a list of documented functions, use library(help = "rpostgisLT")

Author(s)

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