

Package ‘SMITIDstruct’

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Title Data Structure and Manipulations Tool for Host and Viral Population

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Description Statistical Methods for Inferring Transmissions of Infectious Diseases from deep sequencing data (SMITID).

It allow sequence-space-time host and viral population data storage, indexation and querying.

License GPL (>= 2) | file LICENSE

LazyData true

BuildVignettes true

NeedsCompilation no

Biarch true

URL <https://informatique-mia.inra.fr/biosp/anr-smtid-project>,
<https://gitlab.paca.inra.fr/SMITID/structR>

BugReports <https://gitlab.paca.inra.fr/SMITID/structR/issues>

Depends methods, utils, grDevices (>= 3.0.0), graphics (>= 3.0.0), R (>= 3.3.0)

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ImportsNote BioC (>= 3.0), Recommended: Biostrings

Suggests testthat (>= 2.0)

Collate 'Class-Host.R' 'Class-ViralPop.R' 'Methods-Host.R'
'Methods-ViralPop.R' 'Methods-time.R' 'SMITIDstruct.R' 'demo.R'
'diversity.R' 'index.R'

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SMITIDstruct-package *Data Structure and Manipulation Tool for Host and Viral Population*

Description

Statistical Methods for Inferring Transmissions of Infectious Diseases from deep sequencing data (SMITID). It allow sequence-space-time host and viral population data storage, indexation and querying.

Details

Package: SMITIDstruct
Type: Package
Version: 0.0.5
Date: 2019-06-14
License: GPL (>=2)

The SMITIDstruct package contains functions and methods for manipulating Host and Viral population genetico-space-time data.

Author(s)

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See Also

[demo.SMITIDstruct.run](#)

Examples

```
## Run a simulation
library("SMITIDstruct")
demo.SMITIDstruct.run()
```

addcode

*addcode***Description**

add a code event to an another

Usage

```
addcode(code, code.add)
```

Arguments

code	an existing code
code.add	the code to add

Value

merge of the two code

addHost

*addHost***Description**

add an Host to a HostSet

Usage

```
addHost(lhost, id)
```

Arguments

lhost	a hostSet Object
id	a character of host ID

Value

a HostSet of host object with there ID

Examples

```
lhost <- list()
lhost <- addHost(lhost,"42")
```

`addIndex`

addIndex

Description

add to an index a new eventcode

Usage

`addIndex(index, id_host, time, code)`

Arguments

<code>index</code>	an index
<code>id_host</code>	an host index in HostSet
<code>time</code>	a time
<code>code</code>	an event code

Value

the index updated (add a row or update one)

`addViralObs`

addViralObs

Description

load Viral pop observation in Host object

Usage

`addViralObs(lhost, lvpop)`

Arguments

<code>lhost</code>	a HostSet
<code>lvpop</code>	a ViralPopSet

Value

`lhost` update with viral population observed

<code>alleleCount</code>	<i>alleleCount</i>
--------------------------	--------------------

Description

count allele at each position

Usage

```
alleleCount(mat, seq.char = c("A", "T", "G", "C"))
```

Arguments

<code>mat</code>	a genomique seq list as matrix by row
<code>seq.char</code>	allele alphabet

Value

a matrix, each row as a unique seq and col as allele count by position

<code>concatViralPop</code>	<i>concatViralPop</i>
-----------------------------	-----------------------

Description

concat several Viral population in one ViralPop object

Usage

```
concatViralPop(lvpop, lid)
```

Arguments

<code>lvpop</code>	a ViralPop Set
<code>lid</code>	vector of viralpop id to concat

Value

a ViralPop object with ID concatenation from all IDs and time at 0.

`createAViralPop`*createAViralPop*

Description

Create a new ViralPop object

Usage

```
createAViralPop(host_id, obs_time, seq, id_seq = "seq_ID",
                 seq_value = "seq", prop = "prop", compact = FALSE)
```

Arguments

host_id	host ID which viral pop is observed
obs_time	time of the observation (numeric or date)
seq	a data.frame of sequences ID, sequences and counts
id_seq	column name containing the sequences ID
seq_value	column name containing the sequences
prop	column name containing the count of each sequences
compact	boolean, default FALSE, if TRUE will try group identicals sequences (not implemented yet)

`createHost`*createHost*

Description

create a list of Host class object

Usage

```
createHost(list_host)
```

Arguments

list_host	a character vector of host ID
-----------	-------------------------------

Value

a HostSet of host object with there ID

Examples

```
lh <- seq(1,30,1)
lhost <- createHost(lh)
```

createIndex*createIndex***Description**

create an index of time id_host and event code

Usage

```
createIndex(hostlist)
```

Arguments

hostlist a Hostset

Value

a data.frame with TIME, ID_HOST and EVENTCODE as columns

demo.SMITIDstruct.run *demo.SMITIDstruct.run***Description**

run a demo to load HostSet, ViralPopSet and index

Usage

```
demo.SMITIDstruct.run()
```

diversity.pDistance *diversity.pDistance***Description**

diversity calculation using Mean Pairwise Distance

Usage

```
diversity.pDistance(vpop)
```

Arguments

vpop a ViralPop object

Value

result

<code>diversity.sfs</code>	<i>diversity.sfs</i>
----------------------------	----------------------

Description

Allele frequency spectrum or Site frequency spectra : the distribution of alternative allele frequencies across all sites of genetic sequences

Usage

`diversity.sfs(vpop)`

Arguments

`vpop` a viralPop class

Value

the site frequency spectra

<code>getCov</code>	<i>getCov</i>
---------------------	---------------

Description

get Host(s) covariates

Usage

`getCov(lhost, id = NA)`

Arguments

`lhost` a HostSet
`id` a vector of host id (default NA : all lhost)

Value

a data.frame

getDate	<i>getDate</i>
---------	----------------

Description

Converte timestamp to Date (string)

Usage

```
getDate(time, format = "%Y-%m-%dT%H:%M:%S")
```

Arguments

time	a timestamp or vector of
format	Date format output (default %Y-%m-%dT%H:%M:%S)

Value

time as string date

getDiversity.pDistance	<i>getDiversity.pDistance</i>
------------------------	-------------------------------

Description

get pairwise distance of an host over viral population observed

Usage

```
getDiversity.pDistance(host, lvpop)
```

Arguments

host	an Host object
lvpop	a ViralPopSet object

Value

a data.frame with col as time of observation and p_distance

getDiversity.sfs *getDiversity.sfs*

Description

get Allele Frequency Spectrum or Site Frequency spectra for observed viral pop of an host

Usage

`getDiversity.sfs(host, lvpop)`

Arguments

<code>host</code>	an Host object
<code>lvpop</code>	an ViralPopSet object

Value

a list indexed by time that contains allele.time and count

getInfosByHostAndTime *getInfosByHostAndTime*

Description

get hosts informations, status, infectedby, coordinates and time

Usage

`getInfosByHostAndTime(index, lhost)`

Arguments

<code>index</code>	an index
<code>lhost</code>	a hosts list

Value

a data.frame with colnames (id, time, infectedby, status, probabilities, X ,Y)

getStates

getStates

Description

get Host(s) states

Usage

`getStates(lhost, id = NA)`

Arguments

<code>lhost</code>	a HostSet
<code>id</code>	a vector of host id (default NA : all lhost)

Value

a data.frame

getTimeLine

getTimeLine

Description

get the time line of an host

Usage

`getTimeLine(lhost, id)`

Arguments

<code>lhost</code>	a hostSet
<code>id</code>	a host ID

Value

a data.frame

`getTimestamp`*getTimestamp***Description**

Get the timestamp of Date

Usage

```
getTimestamp(date, format = "%Y-%m-%dT%H:%M:%S")
```

Arguments

- | | |
|---------------------|---|
| <code>date</code> | a date (as string) or vector of |
| <code>format</code> | the date format (default %Y-%m-%dT%H:%M:%S) |

Value

timestamp of the date(s)

`getTransmissionTree`*getTransmissionTree***Description**

get a transmission tree as a data.frame

Usage

```
getTransmissionTree(lhost, id = NA)
```

Arguments

- | | |
|--------------------|---|
| <code>lhost</code> | a hostSet |
| <code>id</code> | a vector of hosts ids (default NA : all host) |

Value

a data.frame as source|target|time in columns

Examples

```
path = system.file("extdata", "data-simul/", package="SMITIDstruct")
lhost <- list()
lhost <- loadTree(lhost,paste(path,"/tree.txt",sep=''))
print(getTransmissionTree(lhost))
```

Host	<i>Class Host</i>
------	-------------------

Description

Spatio-temporal information about Host.

Details

Object can be created by calling ...

rdname Host-class

Slots

ID Host identifier

coordinates Host coordinates in time (as sf)

states Host States/Status (dob, Inf...)

sources data.frame of time and host id who infected this host

offsprings data.frame of time and host id who has been contaminated by this host

ID_V_POP data.frame of time and index of Viral population Observation

covariates data.frame of time, covariate and value of this host.

is.juliendate	<i>is.juliendate</i>
---------------	----------------------

Description

Chek if a numeric is not a timestamp

Usage

```
is.juliendate(time)
```

Arguments

time	a numeric
------	-----------

Value

TRUE if time is a julien day, otherwise FALSE

is.StringDate *is.StringDate*

Description

Check if a string represent a date

Usage

`is.StringDate(date)`

Arguments

`date` a string or a vector of string (without NA)

Value

TRUE if date contains date format

is.timestamp *is.timestamp*

Description

Check if a numeric represent a timestamp

Usage

`is.timestamp(time)`

Arguments

`time` a numeric

Value

TRUE if time \geq 1971

<code>isInCode</code>	<i>isInCode</i>
-----------------------	-----------------

Description

check a code contains a specific code

Usage

```
isInCode(code, thecode)
```

Arguments

<code>code</code>	list of code to test
<code>thecode</code>	the real code

Value

TRUE if code contain thecode otherwise FALSE

<code>loadCoords</code>	<i>loadCoords</i>
-------------------------	-------------------

Description

Load Hosts states

Usage

```
loadCoords(lhost, dfCoords, id = "ID")
```

Arguments

<code>lhost</code>	a HostSet
<code>dfCoords</code>	a data.frame with host ID, time and longitude latitude values
<code>id</code>	colname for host ID

Value

`lhost` updated

Examples

```
path = system.file("extdata", "data-simul/", package="SMITIDstruct")
lhost <- list()
lhost <- loadTree(lhost,paste(path,"/tree.txt",sep=''))
coords <- read.table(file=paste(path,"/hosts_coords.txt",sep=''), header=TRUE, check.names=FALSE)
lhost <- loadCoords(lhost,coords)
```

`loadCovs`*loadCovs*

Description

Load Hosts covariates

Usage

```
loadCovs(lhost, dfCovs, id = "ID", colCovs)
```

Arguments

<code>lhost</code>	a HostSet
<code>dfCovs</code>	a data.frame with host ID in rows and covariates in columns
<code>id</code>	colname for host ID
<code>colCovs</code>	colnames of covariates columns

Value

`lhost` updated with covariates

`loadHost`*loadHost*

Description

load host object from a file

Usage

```
loadHost(file = "host.txt")
```

Arguments

<code>file</code>	a file containing hosts data
-------------------	------------------------------

Value

a list of Host object (HostSet) include Class-Host.R

loadStates

*loadStates***Description**

Load Hosts states

Usage

```
loadStates(lhost, dfStates, id = "ID", colStates)
```

Arguments

lhost	a HostSet
dfStates	a data.frame with host ID and states in columns and time as value
id	colname for host ID
colStates	colnames of States columns

Value

lhost updated

Examples

```
path = system.file("extdata", "data-simul/", package="SMITIDstruct")
lhost <- list()
class(lhost) <- "hostSet"
lhost <- loadTree(lhost,paste(path,"/tree.txt",sep=''))
obs <- read.table(paste(path,"/obs.txt",sep=''),header=TRUE, check.names=FALSE)
obs.states <- c(colnames(obs[-grep("ID|Tobs.*", colnames(obs))]))
lhost <- loadStates(lhost, obs, colStates=obs.states)
```

loadTree

*loadTree***Description**

load sources and offsprings from file

Usage

```
loadTree(lhost = list(), file = "tree.txt", source = "ID-source",
receptor = "ID-receptor", tinf = "Tinf", weight = "Weight")
```

Arguments

lhost	a HostSet
file	a file containing tree data
source	column name for source ID
receptor	column name for receptor ID
tinf	column name for infection Time
weight	column name of infection weight

Value

the lhost param update with sources and offsprings

Examples

```
path = system.file("extdata", "data-simul/", package="SMITIDstruct")
lhost <- list()
class(lhost) <- "hostSet"
lhost <- loadTree(lhost,paste(path,"/tree.txt",sep=''))
```

loadTreeDF

*loadTreeDF***Description**

load sources and offsprings from a data.frame

Usage

```
loadTreeDF(lhost = list(), df = data.frame(), source = "ID-source",
           receptor = "ID-receptor", tinf = "Tinf", weight = "Weight")
```

Arguments

lhost	a HostSet
df	a data.frame containing tree data
source	column name for source ID
receptor	column name for receptor ID
tinf	column name for infection Time
weight	infection links probability

Value

the lhost param update with sources and offsprings

loadViralObs*loadViralObs***Description**

load a ViralPop object

Usage

```
loadViralObs(id, time, file)
```

Arguments

<code>id</code>	host pathogen ID
<code>time</code>	time of the observation (numeric or Date)
<code>file</code>	a fasta file

Value

a new ViralPop object

loadViralPop*loadViralPop***Description**

Load all ViralPop observed in the file.obs

Usage

```
loadViralPop(directory, listFiles, listCol = list(id = "id", timeObs =
"time", filename = "filename"), file.extension = "fasta")
```

Arguments

<code>directory</code>	path where is data
<code>listFiles</code>	a dataframe with host ID, time observation and file name (filename.fasta)
<code>listCol</code>	a list of listFiles colomns names ("id", "timeObs", "filename")
<code>file.extension</code>	genotype file extension

Value

a vector of VirlaPop object

Examples

```
path = system.file("extdata", "data-simul/", package="SMITIDstruct")
files <- list.files(path, pattern = ".*.fasta" ,full.names=FALSE)
lfileinfo <- sapply(files,function(x){return(substr(x,1,nchar(x)-6))})
splitFiles <- strsplit(lfileinfo, "_");
listF <- cbind(data.frame(matrix(unlist(splitFiles),nrow=length(splitFiles), byrow=TRUE),
                           stringsAsFactors = FALSE), names(splitFiles))
colnames(listF) <- c("id", "time", "filename")
lvpop <- loadViralPop(path,listF)
```

loadViralPopSet

loadViralPopSet

Description

load a list of viral populations

Usage

```
loadViralPopSet(lvpop = list(), list)
```

Arguments

lvpop	a viralPopSet (default new one)
list	a list (see details)

Details

The list have to be on this format: list\$HOST_ID\$TIME\$list\$seq_id \$seq \$prop A list indexed by host ID, follow by a list indexed by time (of observation). The last list contains an array of seq_ID (sequence ID), an array of seq (sequence as characters), and an array of the count of seq. example : '\$HOST_42'\$'2014-01-01T00:00:00'\$seq_ID ["SEQ_1","SEQ_2"] '\$HOST_42'\$'2014-01-01T00:00:00'\$seq ["ACGT","TGCA"] '\$HOST_42'\$'2014-01-01T00:00:00'\$seq_ID ["46","6"]

mergeCode

mergeCode

Description

merge a list of event code

Usage

```
mergeCode(listcode)
```

Arguments

listcode	a list of event code*
----------	-----------------------

Value

a code

plotDiversity.pDistance	<i>plotDiversity.pDistance</i>
-------------------------	--------------------------------

Description

plot Mean Pairwise Distance for an host viralpop over time

Usage

```
plotDiversity.pDistance(host, lvpop)
```

Arguments

host	an Host object
lvpop	a ViralPopSet object

plotDiversity.sfs	<i>plotDiversity.sfs</i>
-------------------	--------------------------

Description

plot Allele frequency spectrum for an host viralpop over time

Usage

```
plotDiversity.sfs(host, lvpop)
```

Arguments

host	an Host object
lvpop	a ViralPopSet object

setStates	<i>setStates</i>
-----------	------------------

Description

set hosts states from a data.frame

Usage

```
setStates(lhost, dfStates, colStates = c(id = "ID", time = "time", states  
= "value"))
```

Arguments

lhost	a HostSet
dfStates	a data.frame with host ID and states and time in columns
colStates	vector of the columns name, id, time and states

Value

the HostSet updated

simulateStates	<i>simulateStates</i>
----------------	-----------------------

Description

simulate states from sources infection

Usage

```
simulateStates(lhost)
```

Arguments

lhost	a HostSet
-------	-----------

Value

lhost update with states from sources time ~

ViralPop

Class ViralPop

Description

Viral population data containing genotypes

Slots

ID Host identifier
time Observation time as numeric since 1970/01/01
size Qt of variants
names list of variants id with same sequence
genotypes all variants genotypes (as DNAStringSet)
proportions proportions of each variants

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