

# Package ‘SMITIDvisu’

June 14, 2019

**Type** Package

**Title** Visualize Data for Host and Viral Population from 'SMITIDstruct'  
using HTMLwidgets

**Version** 0.0.6

**Description** Visualisation tools for 'SMITIDstruct' package.  
Allow to visualize host timeline, transmission tree, index diversities  
and variant graph using HTMLwidgets. It mainly using D3JS javascript framework.

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**Suggests** SMITIDstruct, knitr, shiny, testthat

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SMITIDvisu-package	<i>Visualize Data for Host and Viral Population from SMITIDstruct using HTMLwidgets</i>
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**Description**

Visualisation tools for SMITIDstruct package. Allow to visualize host timeline, transmission tree, index diversities and variant graph using HTMLwidgets. It mainly using D3JS, noUiSlider and FileSaver javascript libraries.

**Details**

Package:	SMITIDvisu
Type:	Package
Version:	0.0.6
Date:	2019-06-14
License:	GPL (>=3)

**Author(s)**

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**Examples**

```
library(SMITIDvisu)
demo.SMITIDvisu.run()
```

---

`createRainbowColors`    *createRainbowColors* Create a list of colors for each value v

---

**Description**

`createRainbowColors` Create a list of colors for each value v

**Usage**

```
createRainbowColors(v)
```

**Arguments**

v                    a vector of characters

**Value**

a list of value=color

---

`demo.SMITIDvisu.run`    *demo.SMITIDvisu.run*

---

**Description**

run a demo to visualize data

**Usage**

```
demo.SMITIDvisu.run()
```

df2geojson

*df2geojson*

---

**Description**

Transform a data frame into a string formatted in GeoJSON

**Usage**

```
df2geojson(df, multipleValuesByTime = c())
```

**Arguments**

**df** Data frame to convert in GeoJSON. It must contain at least columns 'id', 'time', 'X' and 'Y'. Additional columns will be added as features' properties.

**multipleValuesByTime** Vector of strings indicating the df columns names which can contain several values by time.

**Value**

a geojson string

**Examples**

```
library(SMITIDvisu)
data(transmissiontree)
geojson <- df2geojson(tt.events, multipleValuesByTime = c('infectedby', 'probabilities'))
```

---

hostline*A host infomation over time*

---

**Description**

kind of host time line

**Usage**

```
data("hostline")
```

**Format**

A data frame with 8 observations on the following 5 variables.

level a character vector  
 label a character vector  
 ID a character vector  
 timestart a character vector  
 timeend a character vector

**Examples**

```
data(hostline)
print(hostline)
```

---

maptt	<i>maptt</i>
-------	--------------

---

**Description**

Display a Transmission Tree over a map.

**Usage**

```
maptt(data, multipleValuesByTime = c(), circleRadius = 6,
  defaultNodeColor = "steelblue", nodeColorByState = list(),
  moveEdgeColor = "steelblue", color1 = "green", color2 = "red",
  nbColors = 10, minWeight = 0, maxWeight = 1, weight1 = 0,
  weight2 = 1, autoFocus = TRUE, keepOldFeatures = TRUE,
  optionsControl = TRUE, gradientControl = TRUE, legend = TRUE,
  width = NULL, height = NULL, elementId = NULL)
```

**Arguments**

data	Either a data frame that will be converted to a GeoJSON collection, or a string describing a valid GeoJSON collection. The data frame must contain at least columns 'id', 'time', 'X' and 'Y'. It can contain columns 'infectedby', 'probabilities'. Additional columns will be added as properties, but will do nothing in this implementation of maptt. See the 'df2geojson' function for more informations.
multipleValuesByTime	Vector of strings indicating the df columns names which can contain several values by time. Typically, you would use 'c('infectedby','probabilities')' if you have these values.
circleRadius	Numeric value specifying the radius of the nodes in pixels.

<code>defaultNodeColor</code>	String indicating the default color of nodes, if their status doesn't match with any color. Colors can be specified in hex.
<code>nodeColorByState</code>	List of strings, indicating the color scheme for each node state.
<code>moveEdgeColor</code>	String indicating the color of the edges representing the move of a node.
<code>color1</code>	String indicating the color corresponding to the <code>minWeight</code> value.
<code>color2</code>	String indicating the color corresponding to the <code>maxWeight</code> value.
<code>nbColors</code>	Number of colors for the color scheme using a gradient between <code>color1</code> and <code>color2</code> . These colors will be used to represent the infection edges according to the infection probability. If no probability is used, the edge will use <code>color2</code> . Three intervals are created : <code>color1</code> will be used for the probabilities between <code>minWeight</code> and <code>weight1</code> . Colors between <code>color1</code> and <code>color2</code> will be used for probabilities between <code>weight1</code> and <code>weight2</code> . <code>color2</code> will be used for probabilities between <code>weight2</code> and <code>maxWeight</code> . This setting can be modified directly on the map if ' <code>gradientControl</code> ' is activated.
<code>minWeight</code>	Minimal weight.
<code>maxWeight</code>	Maximal weight.
<code>weight1</code>	Lowest weight for the color scheme. This setting can be modified directly on the map if ' <code>gradientControl</code> ' is activated.
<code>weight2</code>	Greatest weight for the color scheme. This setting can be modified directly on the map if ' <code>gradientControl</code> ' is activated.
<code>autoFocus</code>	Boolean indicating if the map should focus at the displayed features at each time. This setting can be toggled directly on the map if ' <code>optionsControl</code> ' is activated.
<code>keepOldFeatures</code>	Boolean indicating if old features should be displayed or not. Features are considered "old" if their last 'time' is prior to the current time displayed. This setting can be toggled directly on the map if ' <code>optionsControl</code> ' is activated.
<code>optionsControl</code>	Boolean indicating if the options control should be displayed or not
<code>gradientControl</code>	Boolean indicating if the gradient control should be displayed or not
<code>legend</code>	Boolean indicating if the legend should be displayed or not
<code>width</code>	Numeric width for the area in pixels.
<code>height</code>	Numeric height for the area in pixels.
<code>elementId</code>	The element ID where the map is displayed

### Examples

```
library(SMITIDvisu)
data(transmissiontree)

maptt(tt.events, multipleValuesByTime = c('infectedby', 'probabilities'))

# In this example:
```

```

# - values lower than 20 will be yellow ;
# - values between 20 and 25 will use colors between yellow and red ;
# - values greater than 25 will be red.
maptt(tt.events,
  multipleValuesByTime = c('infectedby', 'probabilities'),
  color1 = 'yellow',
  color2 = 'red',
  nbColors = 10,
  minWeight = 0,
  maxWeight = 30,
  weight1 = 20,
  weight2 = 25
)

```

---

mstCompute

*compute the minimum spanning tree*


---

### Description

compute the minimum spanning tree of a matrix representing edges between nodes (of a graph)

### Usage

```
mstCompute(mat)
```

### Arguments

mat                    weighted matrix representing nodes connection (edges weight)

### Value

a matrix with 1 if nodes are linked, 0 otherwise.

---

mstVariant

*mstVariant*


---

### Description

Draw Variants genotypes distances as a graph using Minimum Spanning Tree algorithm.

### Usage

```
mstVariant(mat, prop, node.prop = NULL, width = NULL, height = NULL,
  elementId = NULL)
```

**Arguments**

mat	a distance matrix between sequence of variants (interger distance no floating values)
prop	a data.frame for variants sequences proportions and count (see details)
node.prop	list of variants with proportions and time (default NULL)
width	numeric width for the area in pixels.
height	numeric hieght for the area in pixels.
elementId	the element ID where is draw

**Details**

**mat** is a simple distance matrix with interger values, row and lines contain a unique identifier of each variant sequences. **prop** is a data.frame where each row is a variant sequence, it have to contain in columns factor "ID", "proportion" and "count". "ID" is a unique identifier matching matrix value identifier, "proportion" is the proportions of the variant sequence and "count" the number of variant sequence in a varions set. **node.prop** is a list with name that matching **mat** identifier and **prop** "ID". Each list element contains a subvector time (Julian or timestamp) and value (proportions). That allow to draw variants proportions over time.

**Examples**

```
library(SMITIDvisu)
data(st)
mstV <- mstVariant(st.dist113_all,st.prop113_all, st.listTimeProp113)

## export as standalone html file
htmlwidgets::saveWidget(mstV, "mstVariant.html")
browseURL("mstVariant.html")
```

---

mstVariantProxy	<i>mstVariantProxy</i>
-----------------	------------------------

---

**Description**

get mstVariantProxy

**Usage**

```
mstVariantProxy(mstVid, session = shiny::getDefaultReactiveDomain())
```

**Arguments**

mstVid	widget instance identifier
session	shiny session



## Examples

```
## Not run:
library(SMITIDvisu)
## server.R
mstVariantProxy <- mstVaraintProxy("mstvariantoutput")

## End(Not run)
```

---

SMITIDvisu-shiny      *Shiny bindings for visualisation widgets*

---

## Description

Output and render functions for using visualisation widgets within Shiny applications and interactive Rmd documents.

## Usage

```
mapttOutput(outputId, width = "100%", height = "400px")
renderMaptt(expr, env = parent.frame(), quoted = FALSE)
mstVariantOutput(outputId, width = "100%", height = "600px")
rendermstVariant(expr, env = parent.frame(), quoted = FALSE)
timeLineOutput(outputId, width = "100%", height = "400px")
renderTimeLine(expr, env = parent.frame(), quoted = FALSE)
transmissionTreeOutput(outputId, width = "100%", height = "500px")
renderTransmissionTree(expr, env = parent.frame(), quoted = FALSE)
```

## Arguments

outputId	output variable to read from
width, height	Must be a valid CSS unit (like "100%", "400px", "auto") or a number, which will be coerced to a string and have "px" appended.
expr	An expression that generates a networkD3 graph
env	The environment in which to evaluate expr.
quoted	Is expr a quoted expression (with quote())? This is useful if you want to save an expression in a variable.

---

st	<i>A SMITIDstruct package variable.</i>
----	---

---

**Description**

A SMITIDstruct package variable from simul-chain as a list. The list is a set of HostSet, ViralPopset and an index

**Usage**

```
data("st")
```

**Format**

The format is: List of 3 \$ :List of 21 .. ..- attr(\*, "class")= chr "HostSet" \$ :List of 20 .. ..- attr(\*, "class")= chr "ViralPopSet" \$ :'data.frame': 79 obs. of 3 variables: ..\$ TIME : chr [1:79] "0" "0" "1.26" "1.35" ... ..\$ ID\_HOST : chr [1:79] "1" "2" "2" "2" ... ..\$ EVENTCODE: chr [1:79] "000011" "000110" "001000" "001000" ...

**Examples**

```
data(st)
## maybe str(st) ; plot(st) ...
```

---

st.dist113_2	<i>Distance matrix of observed variants sequences of a host 113 at time 2 from simulation.</i>
--------------	--

---

**Description**

Levenshtein Distance matrix with rows and cols label as sequences ID.

**Usage**

```
data("st")
```

**Format**

The format is: num [1:23, 1:23] 0 1 1 1 2 1 1 1 1 2 ...

**Examples**

```
data(st)
```

---

st.dist113_all	<i>Distance matrix of observed variants sequences of a host 113 at time 2, 3 and 4 from simulation.</i>
----------------	---

---

**Description**

Levenshtein Distance matrix with rows and cols label as sequences ID. Unique sequence variants observed on host 113 at time 2, 3 and 4 from a simulation.

**Usage**

```
data("st")
```

**Format**

The format is: num [1:51, 1:51] 0 1 1 1 1 1 3 1 1 1 ...

**Examples**

```
data(st)
```

---

st.listTimeProp113	<i>List of variants ID with subvector for time and value.</i>
--------------------	---

---

**Description**

A list indexed by variants sequences ID. Each element contain a time and value vector for time of observation and proportions observed at this time.

**Usage**

```
data("st")
```

**Examples**

```
data(st)
```

---

st.prop113_2	<i>Variants proportions and count for host 113 at time 2 from simulation.</i>
--------------	---

---

**Description**

A data.frame with label "ID", "proportion" and "count" for an host 113 at time 2 from simulation. Each row is a sequence.

**Usage**

```
data("st")
```

**Format**

A data frame with 23 observations on the following 3 variables.

ID a character vector  
proportion a numeric vector  
count a numeric vector

**Examples**

```
data(st)
```

---

st.prop113_all	<i>Variants proportions and count for an host 113 at time 2, 3 and 4 from simulation.</i>
----------------	---

---

**Description**

A data.frame with label "ID", "proportion" and "count" for an host 113 at time 2, 3 and 4 from simulation. Each row is a sequence.

**Usage**

```
data("st")
```

**Format**

A data frame with 51 observations on the following 3 variables.

ID a character vector  
proportion a numeric vector  
count a numeric vector

**Examples**

```
data(st)
```

---

timeLine	<i>timeLine</i>
----------	-----------------

---

## Description

Draw a host time line. Time use timestamp or Date in ISO format.

## Usage

```
timeLine(data, title, color = NULL, width = NULL, height = NULL,  
         elementId = NULL)
```

## Arguments

data	a data.frame that represent hosts status in time with ID, status and time in columns
title	a title as character
color	list of color for timeline elements
width	numeric width for the area in pixels.
height	numeric height for the area in pixels.
elementId	the element ID where is draw

## Examples

```
library(SMITIDvisu)  
data(hostline)  
tl <- timeLine(hostline,  
               title="Example host 113",  
               color=list("infected"="red", "offspring"="green",  
                           "alive"="blue", "inf"="orange",  
                           "dead"="black", "Obs"="purple"))  
  
## export as standalone html file  
htmlwidgets::saveWidget(tl, "timeline.html")  
browseURL("timeline.html")
```

---

timeLineProxy	<i>timeLineProxy</i> get an instance of a timeline
---------------	--

---

**Description**

timeLineProxy get an instance of a timeline

**Usage**

```
timeLineProxy(tlid, session = shiny::getDefaultReactiveDomain())
```

**Arguments**

tlid	a timeline instance id
session	shiny session

**Value**

an object of class timeline\_proxy

**Examples**

```
## Not run:
## server.R
## output server variable
output$timeline <- renderTimeLine({
  timeline(data.frame(), "")
})
## ui.R
timelineOutput("timeline")
## server.R
tlproxy <- timeLineProxy("timeline")

## End(Not run)
```

---

transmissionTree	<i>transmissionTree</i>
------------------	-------------------------

---

**Description**

Draw a transmission tree over the time. Time use timestamp or Date in ISO format ("

**Usage**

```
transmissionTree(nodes, edges, nodes.color = NULL, width = NULL,
  height = NULL, elementId = NULL)
```

**Arguments**

nodes	a data.frame that represent hosts status in time with ID, status and time in columns
edges	a data.frame that represent transmission link between hosts (pathogens) with ID, source, target and time in columns
nodes.color	a list of color for nodes status "status"="color"
width	numeric width for the area in pixels.
height	numeric height for the area in pixels.
elementId	the element ID where is draw

**Examples**

```
library(SMITIDvisu)
data(transmissiontree)
tt <- transmissionTree(tt.nodes,tt.edges, nodes.color = list("default"="black","Inf"="red"))

## export as standalone html file
htmlwidgets::saveWidget(tt, "transTree.html")
browseURL("transTree.html")
```

---

transmissionTreeProxy *transmissionTreeProxy*

---

**Description**

get transmissionTreeProxy

**Usage**

```
transmissionTreeProxy(ttid, session = shiny::getDefaultReactiveDomain())
```

**Arguments**

ttid	widget instance identifier
session	shiny session

**Examples**

```
## Not run:
library(SMITIDvisu)
## server.R
transmissionTreeProxy <- transmissionTreeProxyProxy("transmissionTreeoutput")

## End(Not run)
```

---

tt.edges                      *Pathogen link over the time*

---

**Description**

A data.frame of all transmission links between hosts (pathogens). Five columns ID, source, target, time and weight.

**Usage**

```
data("transmissiontree")
```

**Format**

A data frame with 13 observations on the following 5 variables.

ID a numeric vector

source a character vector

target a factor with levels 113 104 116 115 111 109 105 108 106 112

time a character vector

weight a character vector

**Examples**

```
data(transmissiontree)
print(tt.edges)
```

---

tt.events                      *Data.frame of hosts events information by time. Fake data.*

---

**Description**

Fake simulated data of hosts events over the time.

**Usage**

```
data("transmissiontree")
```



**Format**

A data frame with 63 observations on the following 7 variables.

id a character vector  
time a character vector  
status a character vector  
infectedby a character vector  
probabilities a character vector  
X a numeric vector  
Y a numeric vector

**Examples**

```
data(transmissiontree)  
print(tt.events)
```

---

tt.nodes	<i>Host list with there status over the time.</i>
----------	---

---

**Description**

a data.frame of all the hosts identify by there ID. Three colums is use ID, status and time

**Usage**

```
data("transmissiontree")
```

**Format**

A data frame with 47 observations on the following 3 variables.

ID a character vector  
status a character vector  
time a character vector

**Examples**

```
data(transmissiontree)  
print(tt.nodes)
```

---

updatemstVariant	<i>updatemstVariant</i>
------------------	-------------------------

---

**Description**

update (redraw) an instance on mstVariant

**Usage**

```
updatemstVariant(mstVProxy, mat, prop, propTime = NULL)
```

**Arguments**

mstVProxy	mstVaraintProxy instance
mat	distance matrix
prop	proportions data.frame
propTime	list of each variant by time and proportions

**See Also**

[mstVariant](#)

**Examples**

```
## Not run:
library(SMITIDvisu)
data(mstVariant)
## server.R
mstVaraintProxy("mstvariantoutput") %>% updatemstVariant(st.dist,st.prop)

## End(Not run)
```

---

updateTimeLine	<i>updateTimeLine</i>
----------------	-----------------------

---

**Description**

updateTimeLine

**Usage**

```
updateTimeLine(tlProxy, data, title)
```

**Arguments**

tlProxy	a timeline proxy instance
data	new data
title	new title

**See Also**

[timeLine](#)

**Examples**

```
## Not run:
## server.R
## output server variable
output$timeline <- renderTimeLine({
  timeLine(data.frame(), "")
})
## ui.R
timeLineOutput("timeline")
## server.R
timeLineProxy("timeline") %>% updateTimeLine(newtimeline, "newId")

## End(Not run)
```

---

```
updateTransmissionTree
      updateTransmissionTree
```

---

**Description**

update (redraw) an instance of a transmissionTree

**Usage**

```
updateTransmissionTree(TTProxy, nodes, edges, options = NULL)
```

**Arguments**

TTProxy	transmissionTreeProxy instance
nodes	a data.frame that represent hosts status in time with ID, status and time in columns
edges	a data.frame that represent transmission link between hosts (pathogens) with ID, source, weight, target and time in columns
options	transmissionTree new options

**See Also**[transmissionTree](#)**Examples**

```
## Not run:  
library(SMITIDvisu)  
data(transmissionTree)  
## server.R  
transmissionTreeProxy("transmissionTreeoutput") %>% updatetransmissionTree(tt.nodes, tt.edges)  
  
## End(Not run)
```

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