# Package 'simIDM'

December 11, 2023

Type Package

Title Simulating Oncology Trials using an Illness-Death Model

Version 0.1.0

Description Based on the illness-death model a large number of clinical trials with oncology endpoints progression-free survival (PFS) and overall survival (OS) can be simulated, see Meller, Beyersmann and Rufibach (2019) <doi:10.1002/sim.8295>. The simulation set-up allows for random and event-driven censoring, an arbitrary number of treatment arms, staggered study entry and drop-out. Exponentially, Weibull and piecewise exponentially distributed survival times can be generated. The correlation between PFS and OS can be calculated.

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URL https://github.com/insightsengineering/simIDM/

BugReports https://github.com/insightsengineering/simIDM/issues

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# ${\sf R}$ topics documented:

simIDM-package
addStaggeredEntry
assert_intervals
assert_positive_number
avgHRExpOS
avgHRIntegExpOS
censoringByNumberEvents
corPFSOS
corTrans
empSignificant
estimateParams
ExpHazOS
exponential_transition
ExpQuantOS
ExpSurvOS
ExpSurvPFS
expvalOSInteg
expvalPFSInteg
getCensoredData
getClinicalTrials
getDatasetWideFormat
getEventsAll
getInit
getNumberEvents
getOneClinicalTrial
getOneToTwoRows
getPCWDistr
getPWCHazard
getResults
getSimulatedData
getSumPCW
getTarget
getTimePoint
getWaitTimeSum
haz
logRankTest
log_p11
negLogLik
PCWInversionMethod
piecewise_exponential
prepareData

simIDM-package	
SIIIIDWI-package	•

weibull transition	
WeibSurvPFS	
WeibSurvOS	
survTrans	
survPFSOS	
survPFS	
survOS	
PWCsurvPFS	
pwA	38

# Description

simIDM simulates a survival multistate model that jointly models PFS and OS.

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# See Also

# Useful links:

- https://github.com/insightsengineering/simIDM/
- Report bugs at https://github.com/insightsengineering/simIDM/issues

4 addStaggeredEntry

addStaggeredEntry Staggered Study Entry

### **Description**

This function adds staggered study entry times to a simulated data set with illness-death model structure.

# Usage

```
addStaggeredEntry(simData, N, accrualParam, accrualValue)
```

### **Arguments**

simData (data.frame)

simulated data frame containing entry and exit times at individual study time

scale. See getSimulatedData() for details.

N (int)

number of patients.

accrualParam (string)

possible values are 'time' or 'intensity'.

accrualValue (number)

specifies the accrual intensity. For accrualParam equal time, it describes the length of the accrual period. For accrualParam equal intensity, it describes the number of patients recruited per time unit. If accrualValue is equal to 0, all

patients start at calendar time 0 in the initial state.

# Value

This returns a data set containing a single simulated study containing accrual times, i.e. staggered study entry. This is a helper function of getSimulatedData().

```
 \begin{array}{l} simData <- \ data.frame( \\ id = c(1, 1, 2, 3), \ from = c(0, 1, 0, 0), \ to = c(1, 2, "cens", 2), \\ entry = c(0, 3, 0, 0), \\ exit = c(3, 5.3, 5.6, 7.2), \ censTime = c(6.8, 6.8, 5.6, 9.4) \\ ) \\ addStaggeredEntry(simData, 3, accrualParam = "time", accrualValue = 5) \\ \end{array}
```

assert\_intervals 5

assert\_intervals

Assertion for vector describing intervals

# **Description**

We define an intervals vector to always start with 0, and contain unique ordered time points.

### Usage

```
assert_intervals(x, y)
```

# **Arguments**

x what to check.

y (count)

required length of y.

#### Value

Raises an error if x is not an intervals vector starting with 0.

# **Examples**

```
assert_intervals(c(0, 5, 7), 3)
```

assert\_positive\_number

Assertion for Positive Number

# Description

Assertion for Positive Number

### Usage

```
assert_positive_number(x, zero_ok = FALSE)
```

#### **Arguments**

x what to check.

zero\_ok (flag)

whether x can be zero or not.

# Value

Raises an error if x is not a single positive (or non-negative) number.

6 avgHRExpOS

### **Examples**

```
assert_positive_number(3.2)
assert_positive_number(0, zero_ok = TRUE)
```

avgHRExp0S

Average OS Hazard Ratio from Constant Transition Hazards

### **Description**

Average OS Hazard Ratio from Constant Transition Hazards

### Usage

```
avgHRExpOS(transitionByArm, alpha = 0.5, upper = Inf)
```

# Arguments

```
transitionByArm
```

(list)

transition parameters for each treatment group. See exponential\_transition(),

piecewise\_exponential() and weibull\_transition() for details.

alpha (number)

assigns weights to time points, where values higher than 0.5 assign greater

weight to earlier times and values lower than 0.5 assign greater weight to later

times.

upper (number)

upper (time) limit of integration.

# Value

This returns the value of the average hazard ratio.

```
transitionTrt <- exponential_transition(h01 = 0.18, h02 = 0.06, h12 = 0.17) transitionCtl <- exponential_transition(h01 = 0.23, h02 = 0.07, h12 = 0.19) transitionList <- list(transitionCtl, transitionTrt) avgHRExpOS(transitionByArm = transitionList, alpha = 0.5, upper = 100)
```

avgHRIntegExpOS 7

avgHRIntegExp	OS Helper Function for a	/gHRExpOS()

# Description

It is an integrand of the form OS hazard function with intensities h01, h02, h12 at time point t multiplied with a weighted product of the two OS Survival functions at t (one for intensities h0 and one for h1).

# Usage

```
avgHRIntegExpOS(x, h01, h02, h12, h0, h1, alpha)
```

# **Arguments**

x	(numeric) variable of integration.
h01	(positive number) transition hazard for 0 to 1 transition.
h02	(positive number) transition hazard for 0 to 2 transition.
h12	(positive number) transition hazard for 1 to 2 transition.
h0	(list) transition parameters for the first treatment group.
h1	(list) transition parameters for the second treatment group.
alpha	<pre>(number) weight parameter, see avgHRExpOS().</pre>

### Value

This returns the value of the integrand used to calculate the average hazard ratio for constant transition hazards, see avgHRExpOS().

```
h0 <- list(h01 = 0.18, h02 = 0.06, h12 = 0.17)
h1 <- list(h01 = 0.23, h02 = 0.07, h12 = 0.19)
avgHRIntegExpOS(x = 5, h01 = 0.2, h02 = 0.5, h12 = 0.7, h0 = h0, h1 = h1, alpha = 0.5)
```

 ${\tt censoringByNumberEvents}$ 

Event-driven censoring.

# **Description**

This function censors a study after a pre-specified number of events occurred.

# Usage

```
censoringByNumberEvents(data, eventNum, typeEvent)
```

# **Arguments**

data (data.frame)

illness-death data set in 1rowPatient format.

eventNum (int)

number of events.

typeEvent (string)

type of event. Possible values are PFS and OS.

#### Value

This function returns a data set that is censored after eventNum of typeEvent-events occurred.

```
transition1 <- weibull_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6, p01 = 0.8, p02 = 0.9, p12 = 1)
transition2 <- weibull_transition(h01 = 1, h02 = 1.3, h12 = 1.7, p01 = 1.1, p02 = 0.9, p12 = 1.1)

simStudy <- getOneClinicalTrial(
    nPat = c(20, 20), transitionByArm = list(transition1, transition2),
    dropout = list(rate = 0.3, time = 10),
    accrual = list(param = "time", value = 7)
)
simStudyWide <- getDatasetWideFormat(simStudy)
censoringByNumberEvents(data = simStudyWide, eventNum = 20, typeEvent = "PFS")</pre>
```

corPFSOS 9

corPFS0S

Correlation of PFS and OS event times for data from the IDM

#### **Description**

Correlation of PFS and OS event times for data from the IDM

# Usage

```
corPFSOS(
  data,
  transition,
 bootstrap = TRUE,
 bootstrap_n = 100,
  conf_level = 0.95
)
```

# Arguments

(data.frame) data

in the format produced by getOneClinicalTrial().

transition (TransitionParameters object)

specifying the assumed distribution of transition hazards. Initial parameters

for optimization can be specified here. See exponential\_transition() or

weibull\_transition() for details.

(flag) bootstrap

if TRUE computes confidence interval via bootstrap.

bootstrap\_n (count)

number of bootstrap samples.

conf\_level (proportion)

confidence level for the confidence interval.

#### Value

The correlation of PFS and OS.

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)</pre>
data <- getClinicalTrials(</pre>
 nRep = 1, nPat = c(100), seed = 1234, datType = "1rowTransition",
 transitionByArm = list(transition), dropout = list(rate = 0.5, time = 12),
 accrual = list(param = "intensity", value = 7)
)[[1]]
corPFSOS(data, transition = exponential_transition(), bootstrap = FALSE)
## Not run:
corPFSOS(data, transition = exponential_transition(), bootstrap = TRUE)
```

10 empSignificant

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

corTrans

Correlation of PFS and OS event times for Different Transition Models

#### **Description**

Correlation of PFS and OS event times for Different Transition Models

### Usage

```
corTrans(transition)
```

#### **Arguments**

transition (TransitionParameters)

see exponential\_transition(), weibull\_transition() or piecewise\_exponential()

for details.

#### Value

The correlation of PFS and OS.

### **Examples**

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6) corTrans(transition)
```

empSignificant

Empirical Significance for a List of Simulated Trials

# Description

This function computes four types of empirical significance — PFS, OS, at-least (significant in at least one of PFS/OS), and joint (significant in both PFS and OS) — using the log-rank test. Empirical significance is calculated as the proportion of significant results in simulated trials, each ending when a set number of PFS/OS events occur. Critical values for PFS and OS test significance must be specified. If trials simulate equal transition hazards across groups (H0), empirical significance estimates type I error; if they simulate differing transition hazards (H1), it estimates power.

# Usage

```
empSignificant(simTrials, criticalPFS, criticalOS, eventNumPFS, eventNumOS)
```

estimateParams 11

# Arguments

```
simTrials (list)
simulated trial data sets, see getClinicalTrials().

criticalPFS (positive number)
critical value of the log-rank test for PFS.

criticalOS (positive number)
critical value of the log-rank test for OS.

eventNumPFS (integer)
number of PFS events required to trigger PFS analysis.

eventNumOS (integer)
number of OS events required to trigger OS analysis.
```

#### Value

This returns values of four measures of empirical significance.

# **Examples**

```
transition1 <- exponential_transition(h01 = 0.06, h02 = 0.3, h12 = 0.3)
transition2 <- exponential_transition(h01 = 0.1, h02 = 0.4, h12 = 0.3)
simTrials <- getClinicalTrials(
    nRep = 50, nPat = c(800, 800), seed = 1234, datType = "1rowPatient",
    transitionByArm = list(transition1, transition2), dropout = list(rate = 0.5, time = 12),
    accrual = list(param = "intensity", value = 7)
)
empSignificant(
    simTrials = simTrials, criticalPFS = 2.4, criticalOS = 2.2,
    eventNumPFS = 300, eventNumOS = 500
)</pre>
```

estimateParams Estimate Parameters of the Multistate Model Using Clinical Trial Data

# Description

Estimate Parameters of the Multistate Model Using Clinical Trial Data

#### Usage

```
estimateParams(data, transition)
```

12 ExpHazOS

# Arguments

data (data.frame) in the format produced by getOneClinicalTrial().

transition (TransitionParameters object)

specifying the assumed distribution of transition hazards. Initial parameters for optimization can be specified here. See exponential\_transition() or unitable transition() for details

weibull\_transition() for details.

#### **Details**

This function estimates parameters for transition models using clinical trial data. The transition object can be initialized with starting values for parameter estimation. It uses stats::optim() to optimize the parameters.

#### Value

Returns a TransitionParameters object with the estimated parameters.

#### **Examples**

```
transition <- exponential_transition(h01 = 2, h02 = 1.4, h12 = 1.6)
simData <- getOneClinicalTrial(
    nPat = c(30), transitionByArm = list(transition),
    dropout = list(rate = 0.3, time = 12),
    accrual = list(param = "time", value = 1)
)
# Initialize transition with desired starting values for optimization:
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
estimate <- estimateParams(simData, transition)</pre>
```

ExpHaz0S

OS Hazard Function from Constant Transition Hazards

# **Description**

OS Hazard Function from Constant Transition Hazards

#### **Usage**

```
ExpHazOS(t, h01, h02, h12)
```

# **Arguments**

```
t (numeric)
study time-points.
h01 (positive number)
```

transition hazard for 0 to 1 transition.

exponential\_transition 13

h02	(positive number)
	transition bazard for

transition hazard for 0 to 2 transition.

h12 (positive number)

transition hazard for 1 to 2 transition.

#### Value

This returns the value of the OS hazard function at time t.

# **Examples**

```
ExpHazOS(c(1:5), 0.2, 1.1, 0.8)
```

```
exponential_transition
```

Transition Hazards for Exponential Event Times

# Description

This creates a list with class TransitionParameters containing hazards, time intervals and Weibull rates for exponential event times in an illness-death model.

# Usage

```
exponential_transition(h01 = 1, h02 = 1, h12 = 1)
```

# **Arguments**

positive nur	ıber)
	positive num

transition hazard for 0 to 1 transition.

h02 (positive number)

transition hazard for 0 to 2 transition.

h12 (positive number)

transition hazard for 1 to 2 transition.

#### Value

List with elements hazards, intervals, weibull\_rates and family (exponential).

```
exponential_transition(1, 1.6, 0.3)
```

14 ExpSurvOS

ExpQuantOS Quantile function for OS survival function induced by an illness-demodel	ath
---	-----

# Description

Quantile function for OS survival function induced by an illness-death model

# Usage

```
ExpQuantOS(q = 1/2, h01, h02, h12)
```

# Arguments

q	(numeric) quantile(s) at which to compute event time ( $q = 1 / 2$ corresponds to median).
h01	(numeric vector) constant transition hazards for 0 to 1 transition.
h02	(numeric vector) constant transition hazards for 0 to 2 transition.
h12	(numeric vector) constant transition hazards for 1 to 2 transition.

# Value

This returns the time(s) t such that the OS survival function at t equals q.

# **Examples**

```
ExpQuantOS(1 / 2, 0.2, 0.5, 2.1)
```

ExpSurv0S	
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OS Survival Function from Constant Transition Hazards

# Description

OS Survival Function from Constant Transition Hazards

# Usage

```
ExpSurvOS(t, h01, h02, h12)
```

**ExpSurvPFS** 15

# **Arguments**

t (numeric) study time-points. h01 (positive number) transition hazard for 0 to 1 transition. h02 (positive number) transition hazard for 0 to 2 transition. h12

(positive number)

transition hazard for 1 to 2 transition.

#### Value

This returns the value of OS survival function at time t.

# **Examples**

```
ExpSurvOS(c(1:5), 0.2, 0.4, 0.1)
```

ExpSurvPFS

PFS Survival Function from Constant Transition Hazards

# **Description**

PFS Survival Function from Constant Transition Hazards

# Usage

```
ExpSurvPFS(t, h01, h02)
```

# **Arguments**

t (numeric) study time-points. h01 (positive number)

transition hazard for 0 to 1 transition.

h02 (positive number)

transition hazard for 0 to 2 transition.

### Value

This returns the value of PFS survival function at time t.

```
ExpSurvPFS(c(1:5), 0.2, 0.4)
```

16 expvalPFSInteg

expvalOSInteg

*Helper Function for Computing E(OS*^2)

# Description

Helper Function for Computing E(OS^2)

# Usage

```
expvalOSInteg(x, transition)
```

### **Arguments**

x (numeric)

variable of integration.

transition (TransitionParameters)

see exponential\_transition(), weibull\_transition() or piecewise\_exponential()

for details.

#### Value

Numeric results of the integrand used to calculate E(OS^2).

# **Examples**

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6) expvalOSInteg(0.4, transition)
```

expvalPFSInteg

*Helper Function for Computing E(PFS*^2)

# Description

Helper Function for Computing E(PFS^2)

# Usage

```
expvalPFSInteg(x, transition)
```

#### **Arguments**

x (numeric)

variable of integration.

transition (TransitionParameters)

see exponential\_transition(), weibull\_transition() or piecewise\_exponential()

for details.

17 getCensoredData

#### Value

Numeric results of the integrand used to calculate E(PFS^2).

#### **Examples**

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
expvalPFSInteg(0.4, transition)
```

getCensoredData

Helper function for censoringByNumberEvents

### **Description**

Helper function for censoringByNumberEvents

#### Usage

```
getCensoredData(time, event, data)
```

### **Arguments**

time (numeric) event times. event (numeric) event indicator. data

(data.frame)

data frame including patient id id, recruiting time recruitTime and individual

censoring time censTimeInd.

# Value

This function returns a data frame with columns: event time, censoring indicator, event indicator and event time in calendar time.

```
transition1 < -weibull\_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6, p01 = 0.8, p02 = 0.9, p12 = 1)
transition2 <- weibull_transition(h01 = 1, h02 = 1.3, h12 = 1.7, p01 = 1.1, p02 = 0.9, p12 = 1.1)
simStudy <- getOneClinicalTrial(</pre>
 nPat = c(20, 20), transitionByArm = list(transition1, transition2),
 dropout = list(rate = 0.3, time = 10),
 accrual = list(param = "time", value = 7)
)
simStudyWide <- getDatasetWideFormat(simStudy)</pre>
simStudyWide$censTimeInd <- 5 - simStudyWide$recruitTime</pre>
NotRecruited <- simStudyWide$id[simStudyWide$censTimeInd < 0]
censoredData <- simStudyWide[!(simStudyWide$id %in% NotRecruited), ]</pre>
getCensoredData(time = censoredData$OStime, event = censoredData$OSevent, data = censoredData)
```

18 getClinicalTrials

getClinicalTrials

Simulation of a Large Number of Oncology Clinical Trials

# **Description**

Simulation of a Large Number of Oncology Clinical Trials

# Usage

```
getClinicalTrials(nRep, ..., seed = 1234, datType = "1rowTransition")
```

# **Arguments**

nRep	(int) number of simulated trials.
•••	parameters transferred to $getOneClinicalTrial()$ , see $getOneClinicalTrial()$ for details.
seed	(int) random seed used for this simulation.
datType	<pre>(string) possible values are 1rowTransition and 1rowPatient.</pre>

#### Value

This function returns a list with nRep simulated data sets in the format specified by datType. See getDatasetWideFormat() getOneClinicalTrial() for details.

```
transition1 <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
transition2 <- exponential_transition(h01 = 1, h02 = 1.3, h12 = 1.7)
getClinicalTrials(
   nRep = 10, nPat = c(20, 20), seed = 1234, datType = "1rowTransition",
   transitionByArm = list(transition1, transition2), dropout = list(rate = 0.5, time = 12),
   accrual = list(param = "intensity", value = 7)
)</pre>
```

getDatasetWideFormat 19

getDatasetWideFormat	Conversion of a Data Set from One Row per Transition to One Row per Patient
----------------------	---

# Description

Conversion of a Data Set from One Row per Transition to One Row per Patient

#### Usage

```
getDatasetWideFormat(data)
```

#### **Arguments**

data (data.frame)

data frame containing entry and exit times of an illness-death model. See getSimulatedData() for details.

#### **Details**

The output data set contains the following columns:

- id (integer): patient id.
- trt integer): treatment id.
- PFStime (numeric): event time of PFS event.
- CensoredPFS (logical): censoring indicator for PFS event.
- PFSevent (logical): event indicator for PFS event.
- OStime (numeric): event time of OS event.
- CensoredOS (logical): censoring indicator for OS event.
- OSevent (logical): event indicator for OS event.
- recruitTime (numeric): time of recruitment.
- OStimeCal (numeric): OS event time at calendar time scale.
- PFStimeCal (numeric): PFS event time at calendar time scale.

#### Value

This function returns a data set with one row per patient and endpoints PFS and OS.

20 getEventsAll

#### **Examples**

```
transition1 <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
transition2 <- exponential_transition(h01 = 1, h02 = 1.3, h12 = 1.7)
transition3 <- exponential_transition(h01 = 1.1, h02 = 1, h12 = 1.5)
simData <- getOneClinicalTrial(
    nPat = c(30, 20, 30), transitionByArm = list(transition1, transition2, transition3),
    dropout = list(rate = 0, time = 12),
    accrual = list(param = "time", value = 0)
)
getDatasetWideFormat(simData)</pre>
```

getEventsAll

Number of recruited/censored/ongoing Patients.

#### **Description**

Number of recruited/censored/ongoing Patients.

#### Usage

```
getEventsAll(data, t)
```

#### **Arguments**

```
data (data.frame)
illness-death data set in 1rowPatient format.

t (numeric)
study time-point.
```

#### Value

This function returns number of recruited patients, number of censored and number of patients under observations.

```
transition1 <- weibull_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6, p01 = 0.8, p02 = 0.9, p12 = 1)
transition2 <- weibull_transition(h01 = 1, h02 = 1.3, h12 = 1.7, p01 = 1.1, p02 = 0.9, p12 = 1.1)

simStudy <- getOneClinicalTrial(
    nPat = c(20, 20), transitionByArm = list(transition1, transition2),
    dropout = list(rate = 0.6, time = 10),
    accrual = list(param = "time", value = 0)
)
simStudyWide <- getDatasetWideFormat(simStudy)
getEventsAll(data = simStudyWide, t = 1.5)</pre>
```

getInit 21

getInit

Retrieve Initial Parameter Vectors for Likelihood Maximization

#### **Description**

Retrieve Initial Parameter Vectors for Likelihood Maximization

#### Usage

```
getInit(transition)
## S3 method for class 'ExponentialTransition'
getInit(transition)
## S3 method for class 'WeibullTransition'
getInit(transition)
```

# **Arguments**

transition

(ExponentialTransition or WeibullTransition) containing the initial parameters. See exponential\_transition() or weibull\_transition() for details.

#### Value

The numeric vector of initial parameters for likelihood maximization.

#### Methods (by class)

- getInit(ExponentialTransition): for the Exponential Transition Model
- getInit(WeibullTransition): for the Weibull Transition Model

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
getInit(transition)
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
getInit(transition)
transition <- weibull_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6, p01 = 2, p02 = 2.5, p12 = 3)
getInit(transition)</pre>
```

22 getOneClinicalTrial

getNumberEvents

Helper Function for trackEventsPerTrial

# **Description**

Helper Function for trackEventsPerTrial

#### Usage

```
getNumberEvents(event, time, t)
```

# **Arguments**

```
event (numeric)
event indicator.

time (numeric)
event times.

t (numeric)
study time-point.
```

#### Value

This function returns the number of events occurred until time t.

# **Examples**

```
event <- c(0, 1, 1, 1, 0)
time <- c(3, 3.4, 5, 6, 5.5)
getNumberEvents(event = event, time = time, t = 5)
```

getOneClinicalTrial

Simulation of a Single Oncology Clinical Trial

#### **Description**

This function creates a data set with a single simulated oncology clinical trial with one row per transition based on an illness-death model. Studies with an arbitrary number of treatment arms are possible.

# Usage

```
getOneClinicalTrial(
  nPat,
  transitionByArm,
  dropout = list(rate = 0, time = 12),
  accrual = list(param = "time", value = 0)
)
```

getOneToTwoRows 23

### Arguments

nPat (integer)
numbers of patients per treatment arm.

transitionByArm

(list)
transition parameters for each treatment group. See exponential\_transition(),
piecewise\_exponential() and weibull\_transition() for details.

dropout dropout (list)
specifies drop-out probability. See getSimulatedData() for details. Can be
specified either as one list that should be applied to all treatment groups or a
separate list for each treatment group.

accrual accrual (list)

specifies accrual intensity. See addStaggeredEntry() for details. Can be specified either as one list that should be applied to all treatment groups or a separate list for each treatment group.

#### Value

This returns a data frame with one simulated clinical trial and multiple treatment arms. See getSimulatedData() for the explanation of the columns. The column trt contains the treatment indicator. This is a helper function of getClinicalTrials().

# Examples

```
transition1 <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
transition2 <- exponential_transition(h01 = 1, h02 = 1.3, h12 = 1.7)
transition3 <- exponential_transition(h01 = 1.1, h02 = 1, h12 = 1.5)
getOneClinicalTrial(
   nPat = c(30, 20, 30), transitionByArm = list(transition1, transition2, transition3),
   dropout = list(rate = 0, time = 12),
   accrual = list(param = "time", value = 0)
)</pre>
```

getOneToTwoRows

Transitions from the Intermediate State to the Absorbing State

### Description

This function creates transition entry and exit times from the intermediate state to the absorbing state for an existing data frame containing the exit times out of the initial state.

# Usage

```
getOneToTwoRows(simDataOne, transition)
```

24 getPCWDistr

# **Arguments**

simDataOne (data.frame)

a data frame containing all patients with transitions into the intermediate state.

See getSimulatedData() for details.

transition (TransitionParameters)

transition parameters comprising hazards, corresponding intervals and weibull\_rates, see exponential\_transition(), piecewise\_exponential() and weibull\_transition()

for details.

#### Value

This returns a data frame with one row per patient for the second transition, i.e. the transition out of the intermediate state. This is a helper function of getSimulatedData().

#### **Examples**

```
simDataOne <- data.frame( id = c(1:3), to = c(1, 1, 1), from = c(0, 0, 0), entry = c(0, 0, 0), \\ exit = c(3, 5.6, 7.2), censTime = c(6.8, 5.9, 9.4) ) \\ transition <- exponential_transition(1, 1.6, 0.3) \\ getOneToTwoRows(simDataOne, transition)
```

getPCWDistr

Piecewise Exponentially Distributed Event Times

# **Description**

This returns event times with a distribution resulting from piece-wise constant hazards using the inversion method.

#### Usage

```
getPCWDistr(U, haz, pw, t_0)
```

# **Arguments**

U	(numeric) uniformly distributed random variables.
haz	(numeric) piecewise constant hazard.
рw	(numeric) time intervals for the piecewise constant hazard.
t_0	(numeric)

the starting times.

getPWCHazard 25

# Value

This returns a vector with event times.

# **Examples**

```
getPCWDistr(U = runif(3), haz = c(1.1, 0.5, 0.4), pw = c(0, 7, 10), t_0 = c(0, 1, 4.2))
```

getPWCHazard

Piecewise Constant Hazard Values

# Description

This returns piecewise constant hazard values at specified time points.

# Usage

```
getPWCHazard(haz, pw, x)
```

# Arguments

haz (numeric)

piecewise constant input hazard.

pw (numeric)

time intervals for the piecewise constant hazard.

x (numeric)

time-points.

# Value

Hazard values at input time-points.

```
getPWCHazard(c(1, 1.2, 1.4), c(0, 2, 3), c(1, 4, 6))
```

26 getResults

getResults	Format Results of Parameter Estimation for Different Transition Models

# **Description**

Format Results of Parameter Estimation for Different Transition Models

# Usage

```
getResults(transition, res)
## S3 method for class 'ExponentialTransition'
getResults(transition, res)
## S3 method for class 'WeibullTransition'
getResults(transition, res)
```

# **Arguments**

# Value

Returns a TransitionParameters object with parameter estimates.

# Methods (by class)

- getResults(ExponentialTransition): for the Exponential Transition Model
- getResults(WeibullTransition): for the Weibull Transition Model

```
results <- c(1.2, 1.5, 1.6)
getResults(exponential_transition(), results)
results <- c(1.2, 1.5, 1.6)
getResults(exponential_transition(), results)
results <- c(1.2, 1.5, 1.6, 2, 2.5, 1)
getResults(weibull_transition(), results)</pre>
```

getSimulatedData 27

getSimulatedData

Simulate Data Set from an Illness-Death Model

# **Description**

This function creates a single simulated data set for a single treatment arm. It simulates data from an illness-death model with one row per transition and subject.

### Usage

```
getSimulatedData(
   N,
   transition = exponential_transition(h01 = 1, h02 = 1, h12 = 1),
   dropout = list(rate = 0, time = 12),
   accrual = list(param = "time", value = 0)
)
```

### Arguments

N (int)

number of patients.

transition (TransitionParameters)

transition parameters comprising hazards, corresponding intervals and weibull\_rates, see exponential\_transition(), piecewise\_exponential() and weibull\_transition()

for details.

dropout (list)

specifies drop-out probability. Random censoring times are generated using exponential distribution. dropout\$rate specifies the drop-out probability per dropout\$time time units. If dropout\$rate is equal to 0, then no censoring is

applied.

accrual (list)

specifies accrual intensity. See addStaggeredEntry() for details.

#### **Details**

The output data set contains the following columns:

- id (integer): patient id.
- from (numeric): starting state of the transition.
- to (character): final state of the transition.
- entry (numeric): entry time of the transition on the individual time scale.
- exit (numeric): exit time of the transition on the individual time scale.
- entryAct (numeric): entry time of the transition on study time scale.
- exitAct (numeric): exit time of the transition on study time scale.
- censAct (numeric): censoring time of the individual on study time scale.

28 getSumPCW

# Value

This returns a data frame with one row per transition per individual.

# **Examples**

```
getSimulatedData(
  N = 10,
  transition = exponential_transition(h01 = 1, h02 = 1.5, h12 = 1),
  dropout = list(rate = 0.3, time = 1),
  accrual = list(param = "time", value = 5)
)
```

getSumPCW

Sum of Two Piecewise Constant Hazards

# Description

This returns the sum of two piecewise constant hazards per interval.

# Usage

```
getSumPCW(haz1, haz2, pw1, pw2)
```

# Arguments

haz1	(numeric) first summand (piecewise constant hazard).
haz2	(numeric) second summand (piecewise constant hazard).
pw1	(numeric) time intervals of first summand.
pw2	(numeric) time intervals of second summand.

# Value

List with elements hazards and intervals for the sum of two piecewise constant hazards.

```
getSumPCW(c(1.2, 0.3, 0.6), c(1.2, 0.7, 1), c(0, 8, 9), c(0, 1, 4))
```

getTarget 29

getTarget

Generate the Target Function for Optimization

#### **Description**

Generate the Target Function for Optimization

# Usage

```
getTarget(transition)
## S3 method for class 'ExponentialTransition'
getTarget(transition)
## S3 method for class 'WeibullTransition'
getTarget(transition)
```

# Arguments

transition (TransitionParameters)

specifying the distribution family. See exponential\_transition() or weibull\_transition()

for details.

#### **Details**

This function creates a target function for optimization, computing the negative log-likelihood for given parameters, data, and transition model type.

# Value

Function that calculates the negative log-likelihood for the given parameters.

#### Methods (by class)

- getTarget(ExponentialTransition): for the Exponential Transition Model
- getTarget(WeibullTransition): for the Weibull Transition Model

```
transition <- exponential_transition(2, 1.3, 0.8)
simData <- getOneClinicalTrial(
   nPat = c(30), transitionByArm = list(transition),
   dropout = list(rate = 0.8, time = 12),
   accrual = list(param = "time", value = 1)
)
params <- c(1.2, 1.5, 1.6) # For ExponentialTransition
data <- prepareData(simData)
transition <- exponential_transition()</pre>
```

30 getTimePoint

```
fun <- getTarget(transition)</pre>
fun(params, data)
transition <- exponential_transition(2, 1.3, 0.8)</pre>
simData <- getOneClinicalTrial(</pre>
  nPat = c(30), transitionByArm = list(transition),
  dropout = list(rate = 0.8, time = 12),
  accrual = list(param = "time", value = 1)
)
params <- c(1.2, 1.5, 1.6)
data <- prepareData(simData)</pre>
transition <- exponential_transition()</pre>
target <- getTarget(transition)</pre>
target(params, data)
transition <- weibull_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6, p01 = 2, p02 = 2.5, p12 = 3)
simData <- getOneClinicalTrial(</pre>
  nPat = c(30), transitionByArm = list(transition),
  dropout = list(rate = 0.8, time = 12),
  accrual = list(param = "time", value = 1)
)
params \leftarrow c(1.2, 1.5, 1.6, 0.8, 1.3, 1.1)
data <- prepareData(simData)</pre>
transition <- weibull_transition()</pre>
target <- getTarget(transition)</pre>
target(params, data)
```

getTimePoint

*Time-point by which a specified number of events occurred.* 

# **Description**

This returns the study time-point by which a specified number of events (PFS or OS) occurred.

# Usage

```
getTimePoint(data, eventNum, typeEvent, byArm = FALSE)
```

#### **Arguments**

data (data.frame)

illness-death data set in 1rowPatient format.

eventNum (int)

number of events.

typeEvent (string)

type of event. Possible values are PFS and OS.

byArm (logical)

if TRUE time-point per treatment arm, else joint evaluation of treatment arms.

getWaitTimeSum 31

# Value

This returns the time-point by which eventNum of typeEvent-events occurred.

#### **Examples**

```
transition1 <- weibull_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6, p01 = 0.8, p02 = 0.9, p12 = 1)
transition2 <- weibull_transition(h01 = 1, h02 = 1.3, h12 = 1.7, p01 = 1.1, p02 = 0.9, p12 = 1.1)

simStudy <- getOneClinicalTrial(
    nPat = c(20, 20), transitionByArm = list(transition1, transition2),
    dropout = list(rate = 0.3, time = 10),
    accrual = list(param = "time", value = 0)
)
simStudyWide <- getDatasetWideFormat(simStudy)
getTimePoint(simStudyWide, eventNum = 10, typeEvent = "OS", byArm = FALSE)</pre>
```

getWaitTimeSum

Event Times Distributed as Sum of Weibull

# Description

This returns event times with a distribution resulting from the sum of two Weibull distributed random variables using the inversion method.

#### Usage

```
getWaitTimeSum(U, haz1, haz2, p1, p2, entry)
```

# Arguments

U	(numeric) uniformly distributed random variables.
haz1	(positive number) first summand (constant hazard).
haz2	(positive number) second summand (constant hazard).
p1	(positive number) rate parameter of Weibull distribution for haz1.
p2	(positive number) rate parameter of Weibull distribution for haz2.
entry	(numeric) the entry times in the current state.

#### Value

This returns a vector with event times.

32 haz

### **Examples**

```
getWaitTimeSum(U = c(0.4, 0.5), haz1 = 0.8, haz2 = 1, p1 = 1.1, p2 = 1.5, entry = c(0, 0))
```

haz

Hazard Function for Different Transition Models

# Description

Hazard Function for Different Transition Models

# Usage

```
haz(transition, t, trans)
## S3 method for class 'ExponentialTransition'
haz(transition, t, trans)
## S3 method for class 'WeibullTransition'
haz(transition, t, trans)
## S3 method for class 'PWCTransition'
haz(transition, t, trans)
```

### **Arguments**

```
transition (ExponentialTransition or WeibullTransition)
see exponential_transition() or weibull_transition() for details.

t (numeric)
time at which hazard is to be computed.

trans (integer)
index specifying the transition type.
```

#### **Details**

The transition types are:

- 1: Transition from state 0 (stable) to 1 (progression).
- 2: Transition from state 0 (stable) to 2 (death).
- 3: Transition from state 1 (progression) to 2 (death).

#### Value

The hazard rate for the specified transition and time.

logRankTest 33

#### Methods (by class)

- haz(ExponentialTransition): for an exponential transition model.
- haz(WeibullTransition): for the Weibull transition model.
- haz(PWCTransition): for the piecewise constant transition model.

#### **Examples**

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6) haz(transition, 0.4, 2) transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6) haz(transition, 0.4, 2) transition <- weibull_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6, p01 = 2, p02 = 2.5, p12 = 3) haz(transition, 0.4, 2) transition <- piecewise_exponential( h01 = c(1, 1, 1), h02 = c(1.5, 0.5, 1), h12 = c(1, 1, 1), pw01 = c(0, 3, 8), pw02 = c(0, 6, 7), pw12 = c(0, 8, 9) ) haz(transition, 6, 2)
```

logRankTest

Log-Rank Test for a Single Trial

### **Description**

This function evaluates the significance of either PFS or OS endpoints in a trial, based on a prespecified critical value.

# Usage

```
logRankTest(data, typeEvent = c("PFS", "OS"), critical)
```

# **Arguments**

data (data.frame)

data frame containing entry and exit times of an illness-death model. See getSimulatedData()

for details.

typeEvent (string)

endpoint to be evaluated, possible values are PFS and OS.

critical (positive number)

critical value of the log-rank test.

#### Value

Logical value indicating log-rank test significance.

34 log\_p11

#### **Examples**

```
transition1 <- exponential_transition(h01 = 0.06, h02 = 0.3, h12 = 0.3)
transition2 <- exponential_transition(h01 = 0.1, h02 = 0.4, h12 = 0.3)
simTrial <- getClinicalTrials(
   nRep = 1, nPat = c(800, 800), seed = 1234, datType = "1rowPatient",
   transitionByArm = list(transition1, transition2), dropout = list(rate = 0.5, time = 12),
   accrual = list(param = "intensity", value = 7)
)[[1]]
logRankTest(data = simTrial, typeEvent = "OS", critical = 3.4)</pre>
```

log\_p11

Probability of Remaining in Progression Between Two Time Points for Different Transition Models

### **Description**

Probability of Remaining in Progression Between Two Time Points for Different Transition Models

#### Usage

```
log_p11(transition, s, t)
```

# Arguments

#### Value

This returns the natural logarithm of the probability of remaining in progression (state 1) between two time points, conditional on being in state 1 at the lower time point.

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6) log_p11(transition, 1, 3)
```

negLogLik 35

negLogLik	Compute the Negative Log-Likelihood for a Given Data Set and Transition Model

# Description

Compute the Negative Log-Likelihood for a Given Data Set and Transition Model

### Usage

```
negLogLik(transition, data)
```

# **Arguments**

#### **Details**

Calculates the negative log-likelihood for a given data set and transition model. It uses the hazard and survival functions specific to the transition model.

### Value

The value of the negative log-likelihood.

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
simData <- getOneClinicalTrial(
    nPat = c(30), transitionByArm = list(transition),
    dropout = list(rate = 0.8, time = 12),
    accrual = list(param = "time", value = 1)
)
negLogLik(transition, prepareData(simData))</pre>
```

**PCWInversionMethod** 

Single Piecewise Exponentially Distributed Event Time

# Description

This returns an event time with a distribution resulting from piece-wise constant hazards using the inversion method.

# Usage

```
PCWInversionMethod(haz, pw, LogU)
```

# **Arguments**

haz (numeric)

piecewise constant hazard.

pw (numeric)

time intervals for the piecewise constant hazard.

LogU (numeric)

transformed uniformly distributed random variables (log(1-U)).

#### Value

This returns one single event time.

# **Examples**

```
PCWInversionMethod(haz = c(1.1, 0.5, 0.4), pw = c(0, 7, 10), LogU = log(1 - runif(1)))
```

piecewise\_exponential Transition Hazards for Piecewise Exponential Event Times

# Description

This creates a list with class TransitionParameters containing hazards, time intervals and Weibull rates for piecewise exponential event times in an illness-death model.

#### Usage

```
piecewise_exponential(h01, h02, h12, pw01, pw02, pw12)
```

prepareData 37

# Arguments

h01	(numeric vector) constant transition hazards for 0 to 1 transition
h02	(numeric vector) constant transition hazards for 0 to 2 transition
h12	(numeric vector) constant transition hazards for 1 to 2 transition
pw01	(numeric vector) time intervals for the piecewise constant hazards h01
pw02	(numeric vector) time intervals for the piecewise constant hazards h02
pw12	(numeric vector) time intervals for the piecewise constant hazards h12

#### Value

List with elements hazards, intervals, weibull\_rates and family (piecewise exponential).

# **Examples**

```
piecewise_exponential( h01 = c(1, 1, 1), h02 = c(1.5, 0.5, 1), h12 = c(1, 1, 1), \\ pw01 = c(0, 3, 8), pw02 = c(0, 6, 7), pw12 = c(0, 8, 9))
```

prepareData

Preparation of a Data Set to Compute Log-likelihood

## **Description**

Preparation of a Data Set to Compute Log-likelihood

## Usage

```
prepareData(data)
```

# Arguments

data (data.frame)

containing entry and exit times of an illness-death model. See getOneClinicalTrial()

for details.

38 pwA

#### **Details**

The output data set contains the following columns:

- id (integer): patient id.
- from (integer): start event state.
- to (integer): end event state.
- trans (integer): transition (1, 2 or 3) identifier
  - 1: Transition from state 0 (stable) to 1 (progression).
  - 2: Transition from state 0 (stable) to 2 (death).
  - 3: Transition from state 1 (progression) to 2 (death).
- entry (numeric): time at which the patient begins to be at risk for the transition.
- exit (numeric): time at which the patient ends to be at risk for the transition.
- status (logical): event indicator for the transition.

#### Value

This function returns a data set with one row per patient and transition, when the patient is at risk.

## **Examples**

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
simData <- getOneClinicalTrial(
   nPat = c(30), transitionByArm = list(transition),
   dropout = list(rate = 0.8, time = 12),
   accrual = list(param = "time", value = 1)
)
prepareData(simData)</pre>
```

pwA

Cumulative Hazard for Piecewise Constant Hazards

#### **Description**

Cumulative Hazard for Piecewise Constant Hazards

#### Usage

```
pwA(t, haz, pw)
```

# **Arguments**

```
t (numeric)
study time-points.

haz (numeric vector)
constant transition hazards.

pw (numeric vector)
time intervals for the piecewise constant hazards.
```

PWCsurvOS 39

# Value

This returns the value of cumulative hazard at time t.

# **Examples**

```
pwA(1:5, c(0.5, 0.9), c(0, 4))
```

PWCsurvOS

OS Survival Function from Piecewise Constant Hazards

# Description

OS Survival Function from Piecewise Constant Hazards

# Usage

```
PWCsurvOS(t, h01, h02, h12, pw01, pw02, pw12)
```

# Arguments

t	(numeric) study time-points.
h01	(numeric vector) constant transition hazards for 0 to 1 transition.
h02	(numeric vector) constant transition hazards for 0 to 2 transition.
h12	(numeric vector) constant transition hazards for 1 to 2 transition.
pw01	(numeric vector) time intervals for the piecewise constant hazards h01.
pw02	(numeric vector) time intervals for the piecewise constant hazards h02.
pw12	(numeric vector) time intervals for the piecewise constant hazards h12.

## Value

This returns the value of OS survival function at time t.

```
PWCsurvOS(1:5, c(0.3, 0.5), c(0.5, 0.8), c(0.7, 1), c(0, 4), c(0, 8), c(0, 3))
```

40 survOS

PWCsu	ırvF	FS
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PFS Survival Function from Piecewise Constant Hazards

# Description

PFS Survival Function from Piecewise Constant Hazards

# Usage

```
PWCsurvPFS(t, h01, h02, pw01, pw02)
```

# Arguments

t	(numeric) study time-points.
h01	(numeric vector) constant transition hazards for 0 to 1 transition.
h02	(numeric vector) constant transition hazards for 0 to 2 transition.
pw01	(numeric vector) time intervals for the piecewise constant hazards h01.
pw02	(numeric vector) time intervals for the piecewise constant hazards h02.

# Value

This returns the value of PFS survival function at time t.

# **Examples**

```
PWCsurvPFS(1:5, c(0.3, 0.5), c(0.5, 0.8), c(0, 4), c(0, 8))
```

surv0S

OS Survival Function for Different Transition Models

# Description

OS Survival Function for Different Transition Models

survOS 41

#### Usage

```
survOS(transition, t)
## S3 method for class 'ExponentialTransition'
survOS(transition, t)
## S3 method for class 'WeibullTransition'
survOS(transition, t)
## S3 method for class 'PWCTransition'
survOS(transition, t)
```

#### **Arguments**

```
transition (TransitionParameters)
see exponential_transition(), weibull_transition() or piecewise_exponential()
for details.

t (numeric)
time at which the value of the OS survival function is to be computed.
```

#### Value

The value of the survival function for the specified transition and time.

#### Methods (by class)

- survOS(ExponentialTransition): Survival Function for an exponential transition model.
- survOS(WeibullTransition): Survival Function for a Weibull transition model.
- survOS(PWCTransition): Survival Function for a piecewise constant transition model.

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6) survOS(transition, 0.4) transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6) survOS(transition, 0.4) transition <- weibull_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6, p01 = 2, p02 = 2.5, p12 = 3) survOS(transition, 0.4) transition <- piecewise_exponential( h01 = c(1, 1, 1), h02 = c(1.5, 0.5, 1), h12 = c(1, 1, 1), pw01 = c(0, 3, 8), pw02 = c(0, 6, 7), pw12 = c(0, 8, 9) ) survOS(transition, 0.4)
```

42 survPFS

survPFS

PFS Survival Function for Different Transition Models

## **Description**

PFS Survival Function for Different Transition Models

## Usage

```
survPFS(transition, t)
## S3 method for class 'ExponentialTransition'
survPFS(transition, t)
## S3 method for class 'WeibullTransition'
survPFS(transition, t)
## S3 method for class 'PWCTransition'
survPFS(transition, t)
```

## **Arguments**

## Value

The value of the survival function for the specified transition and time.

# Methods (by class)

- survPFS(ExponentialTransition): Survival Function for an exponential transition model.
- survPFS(WeibullTransition): Survival Function for a Weibull transition model.
- survPFS(PWCTransition): Survival Function for a piecewise constant transition model.

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
survPFS(transition, 0.4)
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
survPFS(transition, 0.4)
transition <- weibull_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6, p01 = 2, p02 = 2.5, p12 = 3)
survPFS(transition, 0.4)
transition <- piecewise_exponential(</pre>
```

survPFSOS 43

```
h01 = c(1, 1, 1), h02 = c(1.5, 0.5, 1), h12 = c(1, 1, 1),

pw01 = c(0, 3, 8), pw02 = c(0, 6, 7), pw12 = c(0, 8, 9)

)

survPFS(transition, 0.4)
```

survPFS0S

Survival Function of the Product PFS\*OS for Different Transition Models

# **Description**

Survival Function of the Product PFS\*OS for Different Transition Models

## Usage

```
survPFSOS(t, transition)
```

#### **Arguments**

t (numeric)

time at which the value of the PFS\*OS survival function is to be computed.

transition (TransitionParameters)

see exponential\_transition(), weibull\_transition() or piecewise\_exponential()

for details.

#### Value

This returns the value of PFS\*OS survival function at time t.

# **Examples**

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6) survPFSOS(0.4, transition)
```

survTrans

Survival Function for Different Transition Models

## **Description**

Survival Function for Different Transition Models

44 trackEventsPerTrial

## Usage

```
survTrans(transition, t, trans)
## S3 method for class 'ExponentialTransition'
survTrans(transition, t, trans)
## S3 method for class 'WeibullTransition'
survTrans(transition, t, trans)
```

## **Arguments**

transition (ExponentialTransition or WeibullTransition)
see exponential\_transition() or weibull\_transition() for details.

t (numeric)
time at which survival probability is to be computed.

trans (integer)
index specifying the transition type.

#### Value

The survival probability for the specified transition and time.

## Methods (by class)

- survTrans(ExponentialTransition): for the Exponential Transition Model
- survTrans(WeibullTransition): for the Weibull Transition Model

#### **Examples**

```
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
survTrans(transition, 0.4, 2)
transition <- exponential_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6)
survTrans(transition, 0.4, 2)
transition <- weibull_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6, p01 = 2, p02 = 2.5, p12 = 3)
survTrans(transition, 0.4, 2)</pre>
```

trackEventsPerTrial Event tracking in an oncology trial.

## **Description**

Event tracking in an oncology trial.

## Usage

```
trackEventsPerTrial(data, timeP, byArm = FALSE)
```

WeibSurvOS 45

## **Arguments**

data (data.frame)
illness-death data set in 1rowPatient format.

timeP (numeric)
vector of study time-points.

byArm (logical)

if TRUE time-point per treatment arm, else joint evaluation of treatment arms.

#### Value

This function returns a data frame including number of PFS events, number of OS events, number of recruited patients, number of censored patients and number of ongoing patients at timeP.

## **Examples**

```
transition1 <- weibull_transition(h01 = 1.2, h02 = 1.5, h12 = 1.6, p01 = 0.8, p02 = 0.9, p12 = 1)
transition2 <- weibull_transition(h01 = 1, h02 = 1.3, h12 = 1.7, p01 = 1.1, p02 = 0.9, p12 = 1.1)

simStudy <- getOneClinicalTrial(
    nPat = c(20, 20), transitionByArm = list(transition1, transition2),
    dropout = list(rate = 0.3, time = 10),
    accrual = list(param = "time", value = 0)
)
simStudyWide <- getDatasetWideFormat(simStudy)
trackEventsPerTrial(data = simStudyWide, timeP = 1.5, byArm = FALSE)</pre>
```

WeibSurvOS

OS Survival Function from Weibull Transition Hazards

#### **Description**

OS Survival Function from Weibull Transition Hazards

## Usage

```
WeibSurvOS(t, h01, h02, h12, p01, p02, p12)
```

# Arguments

t	<pre>(numeric) study time-points.</pre>
h01	(positive number) transition hazard for 0 to 1 transition.
h02	(positive number) transition hazard for 0 to 2 transition.
h12	(positive number) transition hazard for 1 to 2 transition.

46 WeibSurvPFS

p01	(positive number)
	rate parameter of Weibull distribution for h01.
p02	(positive number) rate parameter of Weibull distribution for h02.
p12	(positive number)
	rate parameter of Weibull distribution for h12.

## Value

This returns the value of OS survival function at time t.

# **Examples**

```
WeibSurvOS(c(1:5), 0.2, 0.5, 2.1, 1.2, 0.9, 1)
```

WeibSurvPFS

PFS Survival Function from Weibull Transition Hazards

# Description

PFS Survival Function from Weibull Transition Hazards

# Usage

```
WeibSurvPFS(t, h01, h02, p01, p02)
```

# Arguments

t	<pre>(numeric) study time-points.</pre>
h01	(positive number) transition hazard for 0 to 1 transition.
h02	(positive number) transition hazard for 0 to 2 transition.
p01	(positive number) rate parameter of Weibull distribution for h01.
p02	(positive number) rate parameter of Weibull distribution for h02.

## Value

This returns the value of PFS survival function at time t.

```
WeibSurvPFS(c(1:5), 0.2, 0.5, 1.2, 0.9)
```

weibull\_transition 47

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Transition Hazards for Weibull Distributed Event Times

# Description

This creates a list with class TransitionParameters containing hazards, time intervals and Weibull rates for Weibull distributed event times in an illness-death model.

#### Usage

```
weibull_transition(h01 = 1, h02 = 1, h12 = 1, p01 = 1, p02 = 1, p12 = 1)
```

# Arguments

h01	(positive number) transition hazard for 0 to 1 transition
h02	(positive number) transition hazard for 0 to 2 transition
h12	(positive number) transition hazard for 1 to 2 transition
p01	(positive number) rate parameter of Weibull distribution for h01
p02	(positive number) rate parameter of Weibull distribution for h02
p12	(positive number) rate parameter of Weibull distribution for h12

## Value

List with elements hazards, intervals, weibull\_rates and family (Weibull).

```
weibull_transition(h01 = 1, h02 = 1.3, h12 = 0.5, p01 = 1.2, p02 = 1.3, p12 = 0.5)
```

# **Index**

addStaggeredEntry, 4 addStaggeredEntry(), 23, 27 assert_intervals, 5 assert_positive_number, 5 avgHRExpOS, 6 avgHRExpOS(), 7 avgHRIntegExpOS, 7	<pre>getSimulatedData(), 4, 19, 23, 24, 33 getSumPCW, 28 getTarget, 29 getTimePoint, 30 getWaitTimeSum, 31 haz, 32</pre>
censoringByNumberEvents, 8 corPFSOS, 9 corTrans, 10	log_p11, 34 logRankTest, 33 negLogLik, 35
empSignificant, 10 estimateParams, 11 ExpHazOS, 12 exponential_transition, 13 exponential_transition(), 6, 9, 10, 12, 16,	PCWInversionMethod, 36 piecewise_exponential, 36 piecewise_exponential(), 6, 10, 16, 23, 24,
expvalPFSInteg, 16  getCensoredData, 17 getClinicalTrials, 18 getClinicalTrials(), 11, 23 getDatasetWideFormat, 19 getDatasetWideFormat(), 18 getEventsAll, 20 getInit, 21 getNumberEvents, 22 getOneClinicalTrial, 22 getOneClinicalTrial(), 9, 12, 18, 37 getOneToTwoRows, 23 getPCWDistr, 24 getPWCHazard, 25 getResults, 26 getSimulatedData, 27	simIDM-package, 3 stats::optim(), 12 survOS, 40 survPFS, 42 survPFSOS, 43 survTrans, 43  trackEventsPerTrial, 44  WeibSurvOS, 45 WeibSurvPFS, 46 weibull_transition, 47 weibull_transition(), 6, 9, 10, 12, 16, 21, 23, 24, 26, 27, 29, 32, 34, 35, 41–44