Package 'RHawkes'

October 12, 2022

Type Package
Title Renewal Hawkes Process
Version 1.0
Date 2022-5-4
Description The renewal Hawkes (RHawkes) process (Wheatley, Filimonov, and Sornette, 2016 <doi:10.1016 j.csda.2015.08.007="">) is an extension to the classical Hawkes self-exciting point process widely used in the modelling of clustered event sequence data. This package provides functions to simulate the RHawkes process with a given immigrant hazard rate function and offspring birth time density function, to compute the exact likelihood of a RHawkes process using the recursive algorithm proposed by Chen and Stindl (2018) <doi:10.1080 10618600.2017.1341324="">, to compute the Rosenblatt residuals for goodness-of-fit assessment, and to predict future event times based on observed event times up to a given time. A function implementing the linear time RHawkes process likelihood approximation algorithm proposed in Stindl and Chen (2021) <doi:10.1007 s11222-021-10002-0=""> is also included.</doi:10.1007></doi:10.1080></doi:10.1016>
License GPL (>= 2)
Depends R ($>= 2.10$), IHSEP
NeedsCompilation no
Author Feng Chen [aut, cre] (https://orcid.org/0000-0001-8627-9337), Tom Stindl [ctb] (https://orcid.org/0000-0001-8627-9337)
Maintainer Feng Chen <feng.chen@unsw.edu.au></feng.chen@unsw.edu.au>
Repository CRAN
Date/Publication 2022-05-05 14:40:09 UTC
R topics documented:
RHawkes-package damllRH EM1partial

2 RHawkes-package

	EM2partial		
	mllRH1		
	mllRH2		
	pred.den		
	pred.haz		
	sim.pred		
	sim.pred1		
	simRHawkes		
	simRHawkes1		
	tms		
Index			23
RHawkes-package Renewal Hawkes Process			

Description

The renewal Hawkes (RHawkes) process (Wheatley, Filimonov, and Sornette, 2016 <doi:10.1016/j.csda.2015.08.007>) is an extension to the classical Hawkes self-exciting point process widely used in the modelling of clustered event sequence data. This package provides functions to simulate the RHawkes process with a given immigrant hazard rate function and offspring birth time density function, to compute the exact likelihood of a RHawkes process using the recursive algorithm proposed by Chen and Stindl (2018) <doi:10.1080/10618600.2017.1341324>, to compute the Rosenblatt residuals for goodness-of-fit assessment, and to predict future event times based on observed event times up to a given time. A function implementing the linear time RHawkes process likelihood approximation algorithm proposed in Stindl and Chen (2021) <doi:10.1007/s11222-021-10002-0> is also included.

Details

The DESCRIPTION file:

Package: RHawkes Type: Package

Title: Renewal Hawkes Process

Version: 1.0 Date: 2022-5-4

Authors@R: c(person(given="Feng", family="Chen", role = c("aut", "cre"), email = "feng.chen@unsw.edu.au", come Description: The renewal Hawkes (RHawkes) process (Wheatley, Filimonov, and Sornette, 2016 <doi:10.1016/j.csda

License: GPL (>=2)

Depends: R (>= 2.10), IHSEP

NeedsCompilation: no

Author: Feng Chen [aut, cre] (https://orcid.org/0000-0002-9646-3338), Tom Stindl [ctb] (https://orcid.org/0000-0002-9646-3338)

Maintainer: Feng Chen <feng.chen@unsw.edu.au>

damllRH 3

Index of help topics:

EM1partial Partial EM algorithm for the RHawkes process,

version 1

EM2partial Partial EM algorithm for the RHawkes process,

version 2

RHawkes-package Renewal Hawkes Process

damllRH Dynamically approxomated minus loglikelihood of

a RHawkes model

mllRH Minus loglikelihood of a RHawkes model mllRH1 Minus loglikelihood of a RHawkes model with

parent probabilities

mllRH2 Minus loglikelihood of a RHawkes model with

Rosenblatt residuals

pred.den RHawkes predictive density function pred.haz RHawkes predictive hazard function quake An RHawkes earthquake data set

sim.pred Simulate a fitted RHawkes process model sim.pred1 Simulate a fitted RHawkes process model for

prediction purposes

simRHawkes Simulate a renewal Hawkes (RHawkes) process simRHawkes1 Simulate a renewal Hawkes (RHawkes) process tms mid-price change times of the AUD/USD exchange

rate

Author(s)

NA

Maintainer: NA

damllRH

Dynamically approxomated minus loglikelihood of a RHawkes model

Description

Calculates an apprximation to the minus loglikelihood of a RHawkes model with given immigration hazard function μ , offspring birth time density function h and branching ratio η relative to event times tms on interval [0, cens].

Usage

```
damllRH(tms, cens, par, q=0.999, qe=0.999,
    h.fn = function(x, p) dexp(x, rate = 1 / p),
    mu.fn = function(x, p) {
        exp(dweibull(x, shape = p[1], scale = p[2], log = TRUE) -
        pweibull(x, shape = p[1], scale = p[2], lower.tail = FALSE, log.p = TRUE))
    },
```

4 damlIRH

```
H.fn = function(x, p) pexp(x, rate = 1 / p),
Mu.fn = function(x, p) {
  -pweibull(x, shape = p[1], scale = p[2], lower.tail = FALSE, log.p = TRUE)
},
keepB=FALSE,
H.inv=function(x,p)qexp(x,rate=1/p) )
```

Arguments

tms	A numeric vector, with values sorted in ascending order. Event times to fit the RHawkes point process model.
cens	A numericl scalar. The censoring time.
par	A numeric vector containing the parameters of the model, in order of the immigration parameters, in $\mu(.)$, offspring distribution parameters, in $h(.)$, and lastly the branching ratio $\eta(.)$.
q	A numeric scalar in (0,1] and close to 1, which controls how far we look back when truncating the distribution of the most recent immigrant.
qe	A numeric scalar in (0,1] and close to 1, which controls how to truncation is used in the offspring birth time distribution.
h.fn	A (vectorized) function. The offspring birth time density function.
mu.fn	A (vectorized) function. The immigrant waiting time hazard function.
H.fn	A (vectorized) function. Its value at t gives the integral of the offspring birth time density function from 0 to t .
Mu.fn	A (vectorized) function. Its value at t gives the integral of the immigrant waiting time hazard function from 0 to t.
keepB	A boolean scalar, indicating whether the looking back values B_i should be part of the output or not.
H.inv	A (vectorized) function, giving the inverse function of the integral of the excitation.

Value

A scalar giving the value of the (approximate) negative log-likelihood, when keepB is FALSE (the default); A list with components mll, whhich contains the value of the negative log-likelihood, Bs, which gives the look-back order of the truncation of the distribution of the last immigrant, and Bes, which gives the look-forward order in determining how far into the future the excitation effect is allowed to last.

Author(s)

Feng Chen <feng.chen@unsw.edu.au>

```
## Not run:
## earthquake times over 96 years
```

EM1partial 5

```
data(quake);
tms <- sort(quake$time);</pre>
# add some random noise to the simultaneous occurring event times
tms[213:214] <- tms[213:214] +
                    sort(c(runif(1, -1, 1)/(24*60), runif(1, -1, 1)/(24*60)))
## calculate the minus loglikelihood of an RHawkes with some parameters
## the default hazard function and density functions are Weibull and
## exponential respectively
mllRH(tms, cens = 96*365.25, par = c(0.5, 20, 1000, 0.5))
damllRH(tms, cens = 96*365.25, par = c(0.5, 20, 1000, 0.5), q=1, qe=1)
## calculate the MLE for the parameter assuming known parametric forms
## of the immigrant hazard function and offspring density functions.
system.time(est <- optim(c(0.5, 20, 1000, 0.5),
                        mllRH, tms = tms, cens = 96*365.25,
                        mu.fn=function(x,p)p[1]/p[2]*(x/p[2])^(p[1]-1),
                        Mu.fn=function(x,p)(x/p[2])^p[1],
                        control = list(maxit = 5000, trace = TRUE),
                        hessian = TRUE)
system.time(est1 <- optim(c(0.5, 20, 1000, 0.5),
                        function(p){
                            if(any(p<0)||p[4]<0||p[4]>=1)
                                return(Inf);
                            damllRH(tms = tms, cens = 96*365.25,
                                    mu.fn=function(x,p)p[1]/p[2]*(x/p[2])^(p[1]-1),
                                    Mu.fn=function(x,p)(x/p[2])^p[1],
                                    par=p,q=0.999999,qe=0.999999)
                        },
                        control = list(maxit = 5000, trace = TRUE),
                        hessian = TRUE)
## point estimate by MLE
est$par
est1$par
## standard error estimates:
diag(solve(est$hessian))^0.5
diag(solve(est1$hessian))^0.5
## End(Not run)
```

EM1partial

Partial EM algorithm for the RHawkes process, version 1

Description

Calculates the RHawkes model parameters via a partial Expectation-Maximization (EM1) algorithm of Wheatley, Filimonov and Sornette (2016).

6 EM1partial

Usage

Arguments

cens A scalar. The censoring time. A numeric vector containing the parameters of the model, in order of the immigration parameters $\mu(.)$, offspring parameters $h(.)$ and lastly the branching ratio $\eta(.)$. The maximum number of iterations to perform. tol The algorithm stops when the difference between the previous iteration and current iteration parameters sum is less than tol. h.fn A (vectorized) function. The offspring density function. mu.fn A (vectorized) function. The immigration hazard function. H.fn A (vectorized) function. Its value at t gives the integral of the offspring density function from 0 to t. Mu.fn A (vectorized) function. Its value at t gives the integral of the immigrant hazard function from 0 to t.	tms	A numeric vector, with values sorted in ascending order. Event times to fit the RHawkes point process model.
gration parameters $\mu(.)$, offspring parameters $h(.)$ and lastly the branching ratio $\eta(.)$. maxiter The maximum number of iterations to perform. tol The algorithm stops when the difference between the previous iteration and current iteration parameters sum is less than tol. h.fn A (vectorized) function. The offspring density function. mu.fn A (vectorized) function. The immigration hazard function. H.fn A (vectorized) function. Its value at t gives the integral of the offspring density function from 0 to t. Mu.fn A (vectorized) function. Its value at t gives the integral of the immigrant hazard	cens	A scalar. The censoring time.
The algorithm stops when the difference between the previous iteration and current iteration parameters sum is less than tol. h.fn A (vectorized) function. The offspring density function. mu.fn A (vectorized) function. The immigration hazard function. H.fn A (vectorized) function. Its value at t gives the integral of the offspring density function from 0 to t. Mu.fn A (vectorized) function. Its value at t gives the integral of the immigrant hazard	pars	gration parameters $\mu(.)$, offspring parameters $h(.)$ and lastly the branching ratio
rent iteration parameters sum is less than tol. h.fn A (vectorized) function. The offspring density function. mu.fn A (vectorized) function. The immigration hazard function. H.fn A (vectorized) function. Its value at t gives the integral of the offspring density function from 0 to t. Mu.fn A (vectorized) function. Its value at t gives the integral of the immigrant hazard	maxiter	The maximum number of iterations to perform.
 mu.fn A (vectorized) function. The immigration hazard function. H.fn A (vectorized) function. Its value at t gives the integral of the offspring density function from 0 to t. Mu.fn A (vectorized) function. Its value at t gives the integral of the immigrant hazard 	tol	
 H. fn A (vectorized) function. Its value at t gives the integral of the offspring density function from 0 to t. Mu.fn A (vectorized) function. Its value at t gives the integral of the immigrant hazard 	h.fn	A (vectorized) function. The offspring density function.
function from 0 to t. Mu.fn A (vectorized) function. Its value at t gives the integral of the immigrant hazard	mu.fn	A (vectorized) function. The immigration hazard function.
	H.fn	
	Mu.fn	
logg.fn A (vectorized) function. The log of the immigrant distribution function.	logg.fn	A (vectorized) function. The log of the immigrant distribution function.

Value

iterations	The number of iterations until convergence
diff	The absolute sum of the difference between the final two parameter estimates
pars	The parameter estimates from the EM algorithm

Author(s)

Feng Chen <feng.chen@unsw.edu.au> Tom Stindl <t.stindl@unsw.edu.au>

EM2partial 7

Examples

```
## Not run:
## simulated data
tms <- sort(runif(100,0,100))
## the slower version of the EM algorithms on simulated data with default
## immigrant hazard function
## and offspring density
system.time( est1 <- EM1partial(tms, 101, c(2,1,0.5,1)) )
## End(Not run)</pre>
```

EM2partial

Partial EM algorithm for the RHawkes process, version 2

Description

Calculates the RHawkes model parameters via a partial Expectation-Maximization (EM2) algorithm of Wheatley, Filimonov and Sornette (2016).

Usage

Arguments

tms	A numeric vector, with values sorted in ascending order. Event times to fit the RHawkes point process model.
cens	A scalar. The censoring time.
pars	A numeric vector containing the parameters of the model, in order of the immigration parameters $\mu(.)$, offspring parameters $h(.)$ and lastly the branching ratio $\eta(.)$.

maxiter The maximum number of iterations to perform.

8 mllRH

tol	The algorithm stops when the difference between the previous iteration and current iteration parameters sum is less than tol.
h.fn	A (vectorized) function. The offspring density function.
mu.fn	A (vectorized) function. The immigration hazard function.
H.fn	A (vectorized) function. Its value at t gives the integral of the offspring density function from 0 to τ .
Mu.fn	A (vectorized) function. Its value at t gives the integral of the immigrant hazard function from 0 to t .
logg.fn	A (vectorized) function. The log of the immigrant distribution function.

Value

iterations	The number of iterations until convergence
diff	The absolute sum of the difference between the final two parameter estimates
pars	The parameter estimates from the EM algorithm

Author(s)

Feng Chen <feng.chen@unsw.edu.au> Tom Stindl <t.stindl@unsw.edu.au>

Examples

```
## Not run:
## simulated data
tms <- sort(runif(100,0,100))
## the quicker version on simulated data with default immigrant hazard function
## and offspring density
system.time( est2 <- EM2partial(tms, 101, c(2,1,0.5,1)) )
## End(Not run)</pre>
```

mllRH

Minus loglikelihood of a RHawkes model

Description

Calculates the minus loglikelihood of a RHawkes model with given immigration hazard function μ , offspring density function h and branching ratio η for event times tms on interval [0, cens].

Usage

```
mllRH(tms, cens, par,
    h.fn = function(x, p) dexp(x, rate = 1 / p),
    mu.fn = function(x, p) {
        exp(dweibull(x, shape = p[1], scale = p[2], log = TRUE) -
        pweibull(x, shape = p[1], scale = p[2], lower.tail = FALSE, log.p = TRUE))
```

mllRH 9

```
},
H.fn = function(x, p) pexp(x, rate = 1 / p),
Mu.fn = function(x, p) {
  -pweibull(x, shape = p[1], scale = p[2], lower.tail = FALSE, log.p = TRUE)
})
```

Arguments

tms	A numeric vector, with values sorted in ascending order. Event times to fit the RHawkes point process model.
cens	A scalar. The censoring time.
par	A numeric vector containing the parameters of the model, in order of the immigration parameters $\mu(.)$, offspring parameters $h(.)$ and lastly the branching ratio $\eta(.)$.
h.fn	A (vectorized) function. The offspring density function.
mu.fn	A (vectorized) function. The immigration hazard function.
H.fn	A (vectorized) function. Its value at t gives the integral of the offspring density function from 0 to t .
Mu.fn	A (vectorized) function. Its value at t gives the integral of the immigrant hazard function from 0 to t.

Value

The value of the negative log-likelihood.

Author(s)

Feng Chen <feng.chen@unsw.edu.au> Tom Stindl <t.stindl@unsw.edu.au>

```
## Not run:
## earthquake times over 96 years
data(quake);
tms <- sort(quake$time);</pre>
# add some random noise to the simultaneous occurring event times
tms[213:214] \leftarrow tms[213:214] +
                    sort(c(runif(1, -1, 1)/(24*60), runif(1, -1, 1)/(24*60)))
## calculate the minus loglikelihood of an RHawkes with some parameters
## the default hazard function and density functions are Weibull and
## exponential respectively
mllRH(tms, cens = 96*365.25, par = c(0.5, 20, 1000, 0.5))
## calculate the MLE for the parameter assuming known parametric forms
## of the immigrant hazard function and offspring density functions.
system.time(est <- optim(c(0.5, 20, 1000, 0.5),
                        mllRH, tms = tms, cens = 96*365.25,
                        control = list(maxit = 5000, trace = TRUE),
```

10 mlIRH1

```
hessian = TRUE)
)

## point estimate by MLE
est$par

## standard error estimates:
diag(solve(est$hessian))^0.5

## End(Not run)
```

mllRH1

Minus loglikelihood of a RHawkes model with parent probabilities

Description

Calculates the minus loglikelihood of a RHawkes model with given immigration hazard function μ , offspring density function h and branching ratio η for event times tms on interval [0,cens]. The same as ml1RH although this version also returns the parent probabilities.

Usage

```
mllRH1(tms, cens, par,
    h.fn = function(x, p) dexp(x, rate = 1/p),
    mu.fn = function(x, p) {
        exp(dweibull(x, shape = p[1], scale = p[2], log = TRUE) -
        pweibull(x, shape = p[1], scale = p[2], lower.tail = FALSE, log.p = TRUE))
        },
        H.fn = function(x, p) pexp(x, rate = 1/p),
        Mu.fn = function(x, p) {
        -pweibull(x, shape = p[1], scale = p[2], lower.tail = FALSE, log.p = TRUE)
        })
```

Arguments

tms	A numeric vector, with values sorted in ascending order. Event times to fit the RHawkes point process model.
cens	A scalar. The censoring time.
par	A numeric vector containing the parameters of the model, in order of the immigration parameters $\mu(.)$, offspring parameters $h(.)$ and lastly the branching ratio $\eta(.)$.
h.fn	A (vectorized) function. The offspring density function.
mu.fn	A (vectorized) function. The immigration hazard function.
H.fn	A (vectorized) function. Its value at t gives the integral of the offspring density function from 0 to t .
Mu.fn	A (vectorized) function. Its value at t gives the integral of the immigrant hazard function from 0 to t.

mllRH2

Value

m11	minus log-likelihood
log.p	parent probabilities
n	number of events

Author(s)

Feng Chen <feng.chen@unsw.edu.au> Tom Stindl <t.stindl@unsw.edu.au>

See Also

ml1RH

Examples

```
tmp <- mllRH1(sort(runif(1000,0,1000)), 1001, c(2,1,0.5,1))
for(i in 1:tmp$n)
  cat(exp(tmp$log.p[i*(i - 1)/2 + 1:i]), "\n")</pre>
```

mllRH2

Minus loglikelihood of a RHawkes model with Rosenblatt residuals

Description

Calculates the minus loglikelihood of a RHawkes model with given immigration hazard function μ , offspring density function h and branching ratio η for event times tms on interval [0,cens]. The same as mllRH although this version also returns the Rosenblatt residuals.

Usage

Arguments

tms A numeric vector, with values sorted in ascending order. Event times to fit the

RHawkes point process model.

cens A scalar. The censoring time.

Par A numeric vector containing the parameters of the model, in order of the immigration parameters $\mu(.)$, offspring parameters h(.) and lastly the branching ratio

 $\eta(.)$.

12 mlIRH2

h.fn	A (vectorized) function. The offspring density function.
mu.fn	A (vectorized) function. The immigration hazard function.
H.fn	A (vectorized) function. Its value at t gives the integral of the offspring density function from 0 to t .
Mu.fn	A (vectorized) function. Its value at t gives the integral of the immigrant hazard function from 0 to t.

Details

Calculate the RHawkes point process Rosenblatt residuals

Value

mll	minus log-likelihood	
U	Rosenblatt residual of observed event time	
n	number of events	

Author(s)

Feng Chen <feng.chen@unsw.edu.au> Tom Stindl <t.stindl@unsw.edu.au>

See Also

mllRH

```
## Not run:
tmp <- mllRH2(sort(runif(1000,0,1000)),1001,c(2,1,0.5,1))
par(mfrow=c(1,2))
qqunif<-function(dat,...){
   dat<-sort(as.numeric(dat));
   n<-length(dat);
   pvec<-ppoints(n);
   plot(pvec,dat,xlab="Theoretical Quantiles",
        ylab="Sample Quantiles",main="Uniform Q-Q Plot",...)
}
qqunif(tmp$U)
acf(tmp$U)
ks.test(tmp$U,"punif")
## End(Not run)</pre>
```

pred.den 13

n	red	_ ^	Δr

RHawkes predictive density function

Description

Calculates the predictive density of the next observed event time after the censoring time cens based on observations over the interval [0,cens].

Usage

```
pred.den(x, tms, cens, par,
    h.fn = function(x, p) dexp(x, rate = 1 / p),
    mu.fn = function(x, p) {
        exp(dweibull(x, shape = p[1], scale = p[2], log = TRUE) -
        pweibull(x, shape = p[1], scale = p[2], lower.tail = FALSE, log.p = TRUE))},
        H.fn = function(x, p) pexp(x, rate = 1 / p),
        Mu.fn = function(x, p) {
        -pweibull(x, shape = p[1], scale = p[2], lower.tail = FALSE, log.p = TRUE)
      })
```

Arguments

X	A scalar. The amount of time after the censoring time cens.
tms	A numeric vector, with values sorted in ascending order. The event times to fit the RHawkes point process model.
cens	A scalar. The censoring time.
par	A numeric vector. Contains the parameters of the model, in order of the immigration parameters $\mu(.)$, offspring parameters $h(.)$ and lastly the branching ratio $\eta(.)$.
h.fn	A (vectorized) function. The offspring density function.
mu.fn	A (vectorized) function. The immigration hazard function.
H.fn	A (vectorized) function. Its value at t gives the integral of the offspring density function from 0 to t .
Mu.fn	A (vectorized) function. Its value at t gives the integral of the immigrant hazard function from 0 to t .

Value

The predictive density of the next event evaluated at x.

Author(s)

Feng Chen <feng.chen@unsw.edu.au> Tom Stindl <t.stindl@unsw.edu.au>

14 pred.haz

Examples

pred.haz

RHawkes predictive hazard function

Description

Calculates the predictive hazard function of the next observed event time after the censoring time cens based on observations over the interval [0,cens].

Usage

```
pred.haz(x, tms, cens, par,
    h.fn = function(x, p) dexp(x, rate = 1 / p),
    mu.fn = function(x, p) {
        exp(dweibull(x, shape = p[1], scale = p[2], log = TRUE) -
        pweibull(x, shape = p[1], scale = p[2], lower.tail = FALSE, log.p = TRUE))
        },
        H.fn = function(x, p) pexp(x, rate = 1/p),
        Mu.fn = function(x, p) {
        - pweibull(x, shape = p[1], scale = p[2], lower.tail = FALSE, log.p = TRUE)
      })
```

Arguments

x	A scalar. The amount of time after the censoring time cens.
tms	A numeric vector, with values sorted in ascending order. The event times to fit the RHawkes point process model.
cens	A scalar. The censoring time.
par	A numeric vector. Contains the parameters of the model, in order of the immigration parameters $\mu(.)$, offspring parameters $h(.)$ and lastly the branching ratio $\eta(.)$.
h.fn	A (vectorized) function. The offspring density function.
mu.fn	A (vectorized) function. The immigration hazard function.
H.fn	A (vectorized) function. Its value at t gives the integral of the offspring density function from 0 to t .
Mu.fn	A (vectorized) function. Its value at t gives the integral of the immigrant hazard function from 0 to t .

quake 15

Value

The predictive hazard rate of the next event evaluated at x.

Author(s)

Feng Chen <feng.chen@unsw.edu.au> Tom Stindl <t.stindl@unsw.edu.au>

Examples

quake

An RHawkes earthquake data set

Description

An earthquake data set containing the earthquake occurrence times near the Japan region previously examined by Ogata (1998).

Usage

```
data(quake)
```

Format

The format is a vector of the arrival/birth times of earthquakes.

Details

Times of arrivals of earthquake occurrences in a vector in ascending order.

Source

Simulated by a call to the function simHawkes1.

```
## Not run:
data(quake)
## number of earthquake occurrences
nrow(quake)
## End(Not run)
```

sim.pred

sim.pred	Simulate a fitted RHawkes process model

Description

Simulate a fitted RHawkes process model after the censoring time cens to a future time point cens.tilde.

Usage

```
sim.pred(tms, re.dist = rweibull, par,
   par.redist = list(shape = par[1], scale = par[2]),
   h.fn = function(x, p) dexp(x, rate = 1 / p), p.ofd = par[3],
   branching.ratio = par[4], cens, cens.tilde = cens * 1.5,
   mu.fn = function(x, p) {
      exp(dweibull(x, shape = p[1], scale = p[2], log = TRUE) -
      pweibull(x, shape = p[1], scale = p[2], lower.tail = FALSE, log.p = TRUE))
})
```

Arguments

	A numeric vector, with values sorted in ascending order. Event times to fit the RHawkes point process model.
re.dist	A (vectorized) function. The immigrant renewal distribution function.
•	A numeric vector, giving the parameters of the model with the immigration parameters $\mu(.)$, offspring parameters $h(.)$ and lastly the branching ratio $\eta(.)$.
par.redist	A numeric vector. The parameters of the immigrant renewal distribution.
h.fn	A (vectorized) function. The offspring density function.
p.ofd	A (named) list. The parameters of the offspring density.
branching.ratio	
	A scalar. The branching ratio parameter.
cens	A scalar. The censoring time.
cens.tilde	A scalar. The future time that the simulation run until.
mu.fn	A (vectorized) function. The immigration hazard function.

Value

A numeric vector that contains the simulated event times from censoring time cens up until cens. tilde

Author(s)

Feng Chen <feng.chen@unsw.edu.au> Tom Stindl <t.stindl@unsw.edu.au>

sim.pred1 17

Examples

```
N <- 5; i <- 0;
data(quake); tms <- sort(quake$time);</pre>
# add some random noise the simultaneous occurring event times
tms[213:214] \leftarrow tms[213:214] +
                    sort(c(runif(1, -1, 1)/(24*60), runif(1, -1, 1)/(24*60)))
# simulate future event time based on MLE fitted RHawkes model
times <- replicate(N,
                    {cat(i<<-i+1,'\n');
                    sim.pred(tms = tms, par = c(0.314, 22.2, 1266, 0.512),
                             cens=35063)
                   })
plot(NA,NA,xlim=c(0,35063*1.5),ylim=c(0,max(lengths(times))+nrow(quake)),
     xlab="time",ylab="Sample path")
lines(c(0,quake$time),0:nrow(quake),type="s")
for(i in 1:N)
    lines(c(tail(quake$time,1),times[[i]]),nrow(quake)+0:length(times[[i]]),
          type="s", lty=2)
```

sim.pred1

Simulate a fitted RHawkes process model for prediction purposes

Description

Simulate a fitted RHawkes process model from the censoring time cens to a future time point cens.tilde, conditional on the observed event times until the censoring time.

Usage

```
sim.pred1(tms, par, re.dist = rweibull,
    par.redist = list(shape = par[1], scale = par[2]),
    of.dis="exp", par.ofdis = list(rate=par[3]),
    branching.ratio = par[4], cens=tail(tms,1)+mean(diff(tms))/2,
    cens.tilde = cens * 1.5,
    mu.fn = function(x, p) {
        exp(dweibull(x, shape = p[1], scale = p[2], log = TRUE) -
        pweibull(x, shape = p[1], scale = p[2], lower.tail = FALSE, log.p = TRUE))
})
```

Arguments

tms	A numeric vector, with values sorted in ascending order. Event times to fit the RHawkes point process model.
par	A numeric vector, giving the parameters of the model with the immigration parameters $\mu(.)$, offspring parameters $h(.)$ and lastly the branching ratio $\eta(.)$.
re.dist	A (vectorized) function. The function to simulate from the immigrant waiting times distribution.

18 sim.pred1

par.redist A (named) list, giving the parameters of the immigrant waiting time distribution.

of.dis A character string, for the name of the offspring birth time distribution.

par.ofdis A (named) list, giving the parameters of the offspring birth time distribution.

branching.ratio

A scalar in [0,1), the branching ratio parameter.

cens A scalar. The censoring time.

cens.tilde A scalar. The future time to run the simulation to.

A (vectorized) function. The immigration hazard function.

Value

mu.fn

A numeric vector that contains the simulated event times from censoring time cens up until cens. tilde

Author(s)

Feng Chen <feng.chen@unsw.edu.au> Tom Stindl <t.stindl@unsw.edu.au>

See Also

sim.pred.

```
N <- 5; i <- 0;
data(quake); tms <- sort(quake$time);</pre>
# add some random noise the simultaneous occurring event times
tms[213:214] \leftarrow tms[213:214] +
                    sort(c(runif(1, -1, 1)/(24*60), runif(1, -1, 1)/(24*60)))
# simulate future event time based on MLE fitted RHawkes model
times <- replicate(N,
                   {cat(i<<-i+1,'\n');
                   sim.pred1(tms = tms, par = c(0.314, 22.2, 1266, 0.512),
                              cens=35063)
                   })
plot(NA,NA,xlim=c(0,35063*1.5),ylim=c(0,max(lengths(times))+nrow(quake)),
     xlab="time",ylab="Sample path")
lines(c(0,quake$time),0:nrow(quake),type="s")
for(i in 1:N)
    lines(c(tail(quake$time,1),times[[i]]),nrow(quake)+0:length(times[[i]]),
          type="s", 1ty=2)
```

simRHawkes 19

simRHawkes	Simulate a renewal Hawkes (RHawkes) process	

Description

Simulate a renewal Hawkes (RHawkes) process with given renewal immigration distribution function, offspring density function and branching ratio.

Usage

```
simRHawkes(re.dist = rweibull, par.redist = list(shape = 1, scale = 1),
    ofspr.den = function(x, p.ofd) 1 / p.ofd * exp(-x / p.ofd),
    p.ofd = 1, branching.ratio = 0.5, cens = 1, B = 10, B0 = 50,
    max.ofspr.den = max(optimize(ofspr.den, c(0, cens), maximum = TRUE,
    p = p.ofd)$obj, ofspr.den(0, p.ofd), ofspr.den(cens, p.ofd)) * 1.1)
```

Arguments

re.dist A (vectorized) function. The immigrant renewal distribution function. par.redist A numeric vector. The parameters of the immigrant renewal distribution. A (vectorized) function. The offspring density function. ofspr.den p.ofd A numeric vector. The parameters of the offspring density. branching.ratio A scalar. The branching ratio parameter. cens A scalar. The censoring time. A scalar. Tuning parameter for simulation of further immigrants. В B0 A scalar. Tuning parameter for simulation of initial immigrants. max.ofspr.den A scalar. The maximum value of the offspring density function from 0 to cens.

Details

The function works by simulating the arrival times of immigrants according to the renewal immigration distribution. The birth times of offspring from each immigrant are then simulated according to an inhomogeneous Poisson processes with appropriate intensity functions.

Value

A numeric vector of all pooled events (immigration/birth) times of all generations 0, 1, ...

Author(s)

Feng Chen <feng.chen@unsw.edu.au> Tom Stindl <t.stindl@unsw.edu.au>

20 simRHawkes1

Examples

simRHawkes1

Simulate a renewal Hawkes (RHawkes) process

Description

Simulate a renewal Hawkes (RHawkes) process with given renewal immigration distribution function, offspring density function and branching ratio.

Usage

```
simRHawkes1(re.dist = rweibull, par.redist = list(shape = 1, scale = 1),
    of.dis = "exp", par.ofdis = list(rate=1),
    branching.ratio = 0.5, cens = 1, B = 10, B0 = 50,
    flatten=TRUE)
```

Arguments

re.dist	A (vectorized) function. The immigrant renewal distribution function.
par.redist	A numeric vector. The parameters of the immigrant renewal distribution.
of.dis	A character string indicating the distribution for the offspring birth times, which has to be the 'distname' part of the rdistname functions for simulating postive random variables implemented in R, such as "exp", "weibull", "gamma", "lnorm", etc.
par.ofdis	A list with named elements, giving the list of parameters of the offspring distri-

branching.ratio

A scalar between 0 and 1, the branching ratio parameter.

bution, such as list(rate=1), list(shape=1,scale=1), etc.

cens A scalar. The censoring time.

B A scalar. Tuning parameter for simulation of further immigrants.

A scalar. Tuning parameter for simulation of initial immigrants.

flatten A boolean scalar, which indicates whether the output events times should be flattened into an increasing sequence of times, or not (in which case the output is the

immigrant arrival times, and the offspring birth times for different immigrants).

tms 21

Details

The function works by simulating the arrival times of immigrants according to the renewal immigration distribution. The birth times of offspring from each immigrant are then simulated according to an inhomogeneous Poisson processes with appropriate intensity functions.

Value

A numeric vector of pooled event (immigration/offspring birth) times of all generations 0, 1, ..., if flatten=TRUE; A list with two components: immitimes for immigrant arrival times, and offspringtimes for birth times of offspring due to different immigrants.

Author(s)

Feng Chen <feng.chen@unsw.edu.au> Tom Stindl <t.stindl@unsw.edu.au>

Examples

```
tms <- simRHawkes1(par.redist = list(shape = 3, scale = 1),</pre>
                   par.ofdis = list(rate=0.5), branching.ratio = 0.5,
                   cens = 50)
plot(stepfun(tms,0:length(tms)),do.points=FALSE,vertical=FALSE,xlim=c(0,50))
tms.clust <- simRHawkes1(par.redist = list(shape = 3, scale = 1),</pre>
                         par.ofdis = list(rate=0.5), branching.ratio = 0.5,
                          cens = 50,flatten=FALSE)
plot(c(0,50),c(0, 1+(nt <-length(it <- tms.clust\$immitimes))),
     type="n",xlab="time",ylab="cluster")
segments(x0 = it, y0 = -0.2, y1 = 0.2)
for(i in 1:nt)
    segments(x0 = c(it[i],it[i]+tms.clust$offspringtimes[[i]]),
             y0=i-0.2, y1=i+0.2)
abline(h=0:(nt+1),col="light gray",lty=2)
segments(x0=unlist(lapply(1:nt,function(i)c(it[i],it[i]+tms.clust$offspringtimes[[i]]))),
         y0=nt+1-0.2,y1=nt+1+0.2)
```

tms

mid-price change times of the AUD/USD exchange rate

Description

A financial data set containing the mid-price changes of the AUD/USD foreign exchange rate during the trading week from 20:00:00 GMT on Sunday 19/07/2015 to 21:00:00 GMT Friday 24/07/2015.

Usage

```
data(tms)
```

22 tms

Format

The format is a list of the arrival times of mid price changes that occur every hour in 121 non-overlapping windows.

Details

Times of arrivals of mid-price changes is listed together in ascending order.

Source

Simulated by a call to the function simHawkes1.

```
data(tms)
## number of non over-lapping hourly windows
length(tms)
```

Index

* EM algorithm	* simulations
EM1partial, 5	sim.pred, 16
EM2partial, 7	sim.pred1,17
* Hawkes process	
RHawkes-package, 2	damllRH, 3
* RHawkes	FM1+i-1 5
damllRH, 3	EM1partial, 5
EM1partial, 5	EM2partial,7
EM2partial, 7	mllRH, 8
mllRH, 8	mllRH1, 10
* Rosenblatt residual	mllRH2, 11
mllRH2, 11	,
* datasets	pred.den, 13
quake, 15	pred.haz,14
tms, 21	
* likelihood	quake, 15
damllRH, 3	Duratas (Duratas analasa)
mllRH, 8	RHawkes (RHawkes-package), 2
mllRH1, 10	RHawkes-package, 2
* package	sim.pred, 16, 18
RHawkes-package, 2	sim.pred1, 17
* parent probabilities	simRHawkes, 19
mllRH1, 10	simRHawkes1, 20
* point process	
mllRH2, 11	tms, 21
pred.den, 13	
pred.haz, 14	
* prediction	
pred.den, 13	
pred.haz, 14	
* predict	
sim.pred, 16	
sim.pred1, 17	
* renewal process	
simRHawkes, 19	
simRHawkes1, 20	
* self-exciting	
simRHawkes, 19	
simRHawkes1, 20	