# Package 'hte'

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## Description

The **hte** package ...

### Author(s)

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2 Bernoulli\_filter

#### **Description**

Filter the individual patient data with given forces of infection, recovery rates, and test characteris-

#### Usage

```
Bernoulli_filter(data, lambda, gamma, theta)
Bfilter(data, theta)
```

#### **Arguments**

data data set

lambda unit-specific force of infection gamma unit-specific recovery rate

theta list of parameters

#### **Details**

Bernoulli\_filter runs a Bernoulli filter, updating the expected prevalence. It returns the log likelihood, occupancy, and expected prevalences.

Bfilter uses an alternative algorithm.

#### Value

Bernoulli\_filter returns a tibble containing the expected prevalences (for both isolated and un-isolated patients), unit occupancy, force of infection, and conditional log likelihood for each unit at each event time. The sum of the log likelihood column (logLik) is the log likelihood of the data

Bfilter returns the log likelihood of the furnished data.

```
library(tidyverse)
set.seed(626292345)

fake_data |>
    arrange(patient, time) -> dat

theta <- list(
    lambda=c(out=0, A=0.01, B=0.1, C=0.2, D=0.3, E=0.5),
    gamma=c(out=0.1, A=0.1, B=0.1, C=0.1, D=0.1, E=0.1),</pre>
```

coal\_last 3

```
p0=0.2,
  isol_factor=0.1,
  alpha=0.05,
  beta=0.2
)
dat |>
  Bfilter(theta) -> ll1
sum(111)
dat |>
  Bernoulli_filter(
   lambda=theta$lambda,
    gamma=theta$gamma,
   theta
  ) -> f
f |> filter(logLik!=0) |> pull(logLik) -> 112
sum(112)
f
f |>
  select(unit,time,prev_i,prev_u) |>
  pivot_longer(c(prev_i,prev_u)) |>
  group_by(unit) |>
  ## prevalence is not estimated outside the hospital
  filter(!all(is.na(value))) |>
  ungroup() |>
  ggplot(aes(x=time,color=name,y=value))+
  geom_line(alpha=0.5)+
  scale_color_manual(values=c(prev_i="blue",prev_u="red"))+
  facet_grid(unit~.,labeller=label_both)+
  labs(y="prevalence")+
  theme_bw()
```

coal\_last

Coalesce with last

#### Description

Fills NA with previous non-NA.

#### Usage

```
coal_last(x)
```

## Arguments

Х

vector

fake\_data

fake\_data

Fake hospital movement, testing, and isolation data

#### **Description**

A simulated outbreak

#### **Details**

```
Data are generated using the default settings of simuldat.
```

The data were generated by:

```
set.seed(339613584)
simuldat(verbose=TRUE) -> fake_data
save(fake_data,file="fake_data.rda")
```

#### See Also

More on simulated data: simuldat()

```
library(tidyverse)
library(lubridate)
## Examine the data:
fake_data
## Verify certain conditions hold:
stopifnot(
  `admission condition violation`=fake_data |>
    group_by(patient,visit) |> slice_head() |>
    filter(event!="admit") |> nrow()==0,
  `discharge condition violation`=fake_data |>
    group_by(patient,visit) |> slice_tail() |>
    filter(event!="discharge", event!="stop") |> nrow()==0,
  `unit violation`=fake_data |> filter(is.na(unit)) |> nrow()==0,
  `event violation`=fake_data |> filter(is.na(event)) |> nrow()==0
)
fake_data |>
  mutate(
   time=as.numeric(
      as.duration(
        interval(date, start="2000-01-01T00:00:00+0000")
      units="day"
  ) -> fake_data
```

fake\_data 5

```
fake_data |>
 group_by(patient, visit) |>
 summarize(dur=max(time)-min(time)) |>
 ungroup() |>
 group_by(patient) |>
 summarize(dur=sum(dur)) |>
 ggplot(aes(x=log10(dur)))+
 geom_histogram(bins=20)+
 labs(title="total duration of hospitalization")+
 theme_bw()
fake_data |>
 group_by(patient,visit) |>
 summarize(dur=max(time)-min(time)) |>
 ungroup() |>
 ggplot(aes(x=log10(dur)))+
 geom_histogram(bins=40)+
 labs(title="duration of hospital visit")+
 theme_bw()
fake_data |>
 filter(
    event!="test",
    event!="isolate",
   event!="release",
    event!="stop"
 group_by(patient, visit) |>
 arrange(time) |>
 mutate(dur=lead(time)-time) |>
 ungroup() |>
 filter(unit!="out",!is.na(dur)) |>
 ggplot(aes(x=log10(dur),fill=unit,group=unit))+
 geom_histogram(aes(y=after_stat(density)),bins=40)+
 facet_grid(unit~.,scales="free_y")+
 labs(title="duration of stay by unit")+
 theme_bw()
fake_data |>
 group_by(patient) |>
 summarize(ntest=sum(event=="test")) |>
 ungroup() |>
 ggplot(aes(x=ntest))+
 geom_histogram(binwidth=1,center=0)+
 labs(title="number of tests per patient")+
 theme_bw()
fake_data |>
 arrange(time) |>
 mutate(
   dn=case_when(
     event=="admit"~1L,
```

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```
event=="discharge"~-1L,
     TRUE~0L
   ),
   occ=cumsum(dn)
 ) |>
 ggplot(aes(x=date,y=occ))+
 geom_step()+
 labs(title="hospital occupancy")+
 theme_bw()
fake_data |>
 arrange(time) |>
 select(date,test.result=result,isol,infected) |>
 pivot_longer(c(test.result,isol,infected)) |>
 filter(!is.na(value)) |>
 ggplot(aes(x=date,y=value,color=name))+
 geom_point()+
 geom_smooth()+
 guides(color="none")+
 labs(
    title="infection and isolation status, test results",
   y=""
 )+
 facet_grid(name~.)+
 theme_bw()
fake_data |>
 filter(event=="test") |>
 mutate(
   interval=cut(time,breaks=72,ordered=TRUE)
 ) |>
 select(interval,time,infected,isol,result) |>
 pivot_longer(c(infected,isol,result)) |>
 group_by(name,interval) |>
 summarize(
   time=mean(time),
   prev=mean(value),
   n=n()
 ) |>
 ungroup() |>
 ggplot(aes(x=time,y=prev,group=name,fill=name))+
 geom_col(position="dodge")+
 labs(title="infection, isolation, and detection through time")+
 theme_bw()+
 theme(axis.text.x=element_text(angle=90))
fake_data |>
 filter(event=="test") |>
 select(infected,result) |>
 count(infected,result) |>
 group_by(infected) |>
 mutate(prob=n/sum(n)) |>
 ungroup()
```

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Independent infection model

#### **Description**

Model under which each patient's trajectory of infection is independent, conditional on the force of infection

#### Usage

```
indep_homog_filter(params, data)
indep_homog_objfun(params, data, est = character(0))
indep_unit_spec_filter(params, data)
indep_unit_spec_objfun(params, data, est = character(0))
```

#### **Arguments**

params named vector of parameters

data patient movement, isolation, and testing data

est names of parameters to estimate

#### **Details**

indep\_homog\_filter runs a Bernoulli filter for the independent model with a global lambda and gamma.

indep\_homog\_objfun is a stateful objective function for the independent model with a global lambda and gamma.

indep\_unit\_spec\_filter runs a Bernoulli filter for the independent model with a unit specific lambda, one gamma for inside the hospital, and one gamma outside the hospital.

indep\_unit\_spec\_objfun is a stateful objective function for the independent model with unit specific lambda, one gamma for inside the hospital, and one gamma outside the hospital.

#### See Also

More on the independent infection model: simuldat()

More on stateful objective functions: stobfun(), transmission

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```
library(tidyverse)
library(lubridate)
library(pomp)
library(hte)
set.seed(626292345)
fake_data |>
  ## filter out tests prior to day 250
  filter(
    event!="test" | time < 250,
    time < 1000
  ) |>
  arrange(patient,time) -> dat
indep_homog_objfun(
  params=c(
    lambda=0.1,gamma=0.01,p0=0.1,
    isol_factor=0.2,alpha=0.02,beta=0.1
  est=c("lambda", "gamma"),
  data=dat
) -> f
optim(
  par=log(c(0.1,0.01)),
  fn=f,
  control=list(reltol=1e-3)
) -> out
f(out$par)
coef(f)
indep_homog_objfun(
  params=coef(f),
  est=c("lambda"),
  data=dat
) -> g
log.lambda \leftarrow seq(log(0.001), to=log(1), length=10)
plot(exp(log.lambda),sapply(log.lambda,g),log='x')
theta <- coef(f)
indep_unit_spec_objfun(
  params=c(
    lambda=setNames(rep.int(theta["lambda"],6),unique(dat$unit)),
    gamma.out=unname(theta["gamma"]),
    theta[c("p0","gamma","alpha","beta","isol_factor")]
  ),
  est=c(
    "lambda.A", "lambda.B", "lambda.C", "lambda.D", "lambda.E", "lambda.out",
    "gamma", "gamma.out"
```

simuldat

simuldat

#### **Description**

simuldat simulates data representing the flow of a body of patients through a hospital over a specified window of time.

simul\_patient simulates a single patient's history of movement, testing, isolation, and infection.

#### Usage

```
simuldat(
      nbeds = c(50, 60),
      arrival = 20,
      window = c("1999-12-31T23:59:59+0000", "2003-01-01T00:00:00+0000"),
     units = list(A = list(shape = 10, scale = 2/10), B = list(shape = 5, scale = 0.5/5), C
         = list(shape = 1, scale = 0.5/1), D = list(shape = 1, scale = 3/1), E = list(shape =
              0.2, scale = 8/0.2), out = list(shape = 0.5, scale = 300/0.5)),
       visits = list(size = 0.5, mu = 9),
       uperv = list(size = 1, mu = 0.5),
      min_dur = 1/24,
       testing_freq = c(A = NA, B = NA, C = 1/7, D = 1/7, E = 1/7, out = NA),
       isolation = list(on = 1/50, off = 1/50),
      infection = list(lambda = c(A = 0.01, B = 0.02, C = 0.001, D = 0.5, E = 0.1, out = 0.001, D = 0.0
              0.05), gamma = 0.01, p0 = 0.1, isol_factor = 0.2),
       alpha = 0.02,
       beta = 0.1,
       verbose = getOption("verbose", TRUE)
```

```
)
simul_patient(
  patient,
  t0,
  tf,
  units,
  visits,
  uperv,
 min_dur,
  testing_freq,
  isolation,
  infection,
  alpha,
 beta
)
indep_infect(lambda, gamma, p0, isol_factor, times, loc, isol)
```

#### **Arguments**

nbeds upper and lower bounds on number of beds

arrival Poisson arrival rate of new patients

window window of simulation

units a named list with one entry per unit. Each entry is itself a list with the parameters

of the Gamma-distribution for the duration of stay in the unit.

visits list containing parameters of a negative binomial distribution for the number of

visits per patient.

uperv list containing parameters for a negative binomial distribution for the number of

units visited per visit.

min\_dur minimum duration of stay in any unit

testing\_freq named numeric vector of unit-specific testing frequencies.

isolation list containing parameters of the isolation model

infection parameters of the infection model

alpha, beta false positive and negative testing error rates

verbose run-time information?
patient patient name or number

t0, tf initial and final times of patient itinerary

lambda force of infection gamma recovery rate

p0 initial probability of infection

isol\_factor reduction in susceptibility due to isolation

times times at which status is reported loc location of patient at each time

isol isolation status

#### **Details**

Simulate hospital movement, testing, and isolation data.

#### Value

infection status vector

#### See Also

More on simulated data: fake\_data

More on the independent infection model: independent

```
library(tidyverse)
library(lubridate)
## Examine the data:
fake_data
## Verify certain conditions hold:
stopifnot(
  `admission condition violation`=fake_data |>
   group_by(patient,visit) |> slice_head() |>
    filter(event!="admit") |> nrow()==0,
 `discharge condition violation`=fake_data |>
   group_by(patient,visit) |> slice_tail() |>
   filter(event!="discharge",event!="stop") |> nrow()==0,
  `unit violation`=fake_data |> filter(is.na(unit)) |> nrow()==0,
  `event violation`=fake_data |> filter(is.na(event)) |> nrow()==0
)
fake_data |>
 mutate(
   time=as.numeric(
     as.duration(
        interval(date, start="2000-01-01T00:00:00+0000")
     ),
     units="day"
 ) -> fake_data
fake_data |>
 group_by(patient, visit) |>
 summarize(dur=max(time)-min(time)) |>
 ungroup() |>
 group_by(patient) |>
 summarize(dur=sum(dur)) |>
 ggplot(aes(x=log10(dur)))+
 geom_histogram(bins=20)+
 labs(title="total duration of hospitalization")+
```

```
theme_bw()
fake_data |>
 group_by(patient,visit) |>
 summarize(dur=max(time)-min(time)) |>
 ungroup() |>
 ggplot(aes(x=log10(dur)))+
 geom_histogram(bins=40)+
 labs(title="duration of hospital visit")+
 theme_bw()
fake_data |>
 filter(
    event!="test",
   event!="isolate",
   event!="release",
   event!="stop"
 ) |>
 group_by(patient,visit) |>
 arrange(time) |>
 mutate(dur=lead(time)-time) |>
 ungroup() |>
 filter(unit!="out",!is.na(dur)) |>
 ggplot(aes(x=log10(dur),fill=unit,group=unit)) +
 geom_histogram(aes(y=after_stat(density)),bins=40)+
 facet_grid(unit~.,scales="free_y")+
 labs(title="duration of stay by unit")+
 theme_bw()
fake_data |>
 group_by(patient) |>
 summarize(ntest=sum(event=="test")) |>
 ungroup() |>
 ggplot(aes(x=ntest))+
 geom_histogram(binwidth=1,center=0)+
 labs(title="number of tests per patient")+
 theme_bw()
fake_data |>
 arrange(time) |>
 mutate(
   dn=case_when(
     event=="admit"~1L,
     event=="discharge"~-1L,
     TRUE~0L
   ),
   occ=cumsum(dn)
 ) |>
 ggplot(aes(x=date,y=occ))+
 geom_step()+
 labs(title="hospital occupancy")+
 theme_bw()
```

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```
fake_data |>
 arrange(time) |>
 select(date,test.result=result,isol,infected) |>
 pivot_longer(c(test.result,isol,infected)) |>
 filter(!is.na(value)) |>
 ggplot(aes(x=date,y=value,color=name))+
 geom_point()+
 geom_smooth()+
 guides(color="none")+
 labs(
    title="infection and isolation status, test results",
   y=""
 facet_grid(name~.)+
 theme_bw()
fake_data |>
 filter(event=="test") |>
 mutate(
    interval=cut(time,breaks=72,ordered=TRUE)
 ) |>
 select(interval,time,infected,isol,result) |>
 pivot_longer(c(infected,isol,result)) |>
 group_by(name,interval) |>
 summarize(
    time=mean(time),
   prev=mean(value),
   n=n()
 ) |>
 ungroup() |>
 ggplot(aes(x=time,y=prev,group=name,fill=name))+
 geom_col(position="dodge")+
 labs(title="infection, isolation, and detection through time")+
 theme_bw()+
 theme(axis.text.x=element_text(angle=90))
fake_data |>
 filter(event=="test") |>
 select(infected,result) |>
 count(infected,result) |>
 group_by(infected) |>
 mutate(prob=n/sum(n)) |>
 ungroup()
```

stobfun

Stateful objective functions

#### **Description**

Convenience functions for constructing and working with stateful objective functions ('stobfun'-class objects).

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#### Usage

```
stobfun(
  embed,
  params,
  est = character(0),
  log = character(0),
  logit = character(0),
  objfun,
  data,
  ...
)

filterfun(embed, params, filtfun, data, ...)

transf_fns(log = character(0), logit = character(0), est = character(0))

embedding(...)

## S3 method for class 'stobfun'
coef(object, ...)
```

#### **Arguments**

embed	embedding (see embedding).
params	vector of parameters
est	character: names of parameters to be estimated
log	character; names of parameters to log transform.
logit	character; names of parameters to logit transform.
objfun	underlying objective function
data	data
	When furnished to stobfun, additional arguments are passed to objfun. When furnished to embedding, arguments define th embedding. When furnished to coef, additional arguments are ignored.
filtfun	the function that actually applies the filter
object	'stobfun'-class stateful objective function

#### **Details**

objfun will be called as objfun(theta, data), where theta is the nested list constructed according to the given specifications and data is the data.

#### Value

transf\_fns returns a list of two functions. The first is the transformation to the estimation scale; the second is its inverse.

embedding returns the embedding function corresponding to the given specification.

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coef(f) returns the parameter vector corresponding to the last call of the stateful objective function f

#### Construction and usage of stateful objective functions

A stateful objective function is an ordinary function that can be used as an objective function in an optimization problem. In particular, it can be passed to optimizers such as optim, subplex, or nloptr. It is stateful in the sense that it remembers the argument with which it was last called.

To construct a stateful objective function, call the constructor function for the model of interest. The constructor function requires that you pass a vector of model parameters: this gives the default parameter values. It also requires that you pass the data and the names of the parameters that you wish to estimate. The constructor will return an object of class 'stobfun'.

Having constructed a 'stobfun' stateful objective function, you can pass this to any suitable optimizer. Once the optimizer has returned, it is important that you call the function one last time, at the parameters the optimizer has returned (see examples). This ensures that the stored parameters are those at the (putative) optimum. You can retrieve these parameters via a call to coef.

#### See Also

More on stateful objective functions: independent, transmission

transmission

Transmission model

#### **Description**

Model under which the force of infection in each unit is proportional to the prevalence of infection in that unit.

#### **Usage**

```
trans_homog_filter(params, data, tol = 1e-04, maxit = 10)

trans_homog_objfun(params, data, est = character(0), tol = 1e-04, maxit = 10)

trans_unit_spec_filter(params, data, tol = 1e-04, maxit = 10)

trans_unit_spec_objfun(
   params,
   data,
   est = character(0),
   tol = 1e-04,
   maxit = 10
)
```

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#### Arguments

params	named vector of parameters
data	patient movement, isolation, and testing data
tol	positive scalar; convergence tolerance (mean difference).
maxit	scalar integer; maximum number of fixed-point iterations. If tol is not achieved in maxit or fewer iterations, an error is generated.
est	names of parameters to estimate

#### **Details**

The basic transmission model assumes that the force of infection on an unisolated patient is

$$\lambda = b(P_u + aP_i),$$

where  $P_u$ ,  $P_i$  are the prevalences among unisolated and isolated patients, respectively. On an isolated patient, this force of infection is reduced by the factor isol\_factor.

trans\_homog\_filter runs a fixed-point Bernoulli filter for the transmission model with global b and gamma.

trans\_homog\_objfun is a stateful objective function for the transmission model with global b and gamma.

trans\_unit\_spec\_filter runs a fixed-point Bernoulli filter for the transmission model with unit-specific b.

trans\_unit\_spec\_objfun is a stateful objective function for the transmission model with unit-specific transmission rates, an out-of-hospital force of infection parameter, and recovery rates that can be different inside and outside of hospital.

#### See Also

More on stateful objective functions: independent, stobfun()

```
library(tidyverse)
library(hte)

set.seed(339613584)

fake_data |>
    ## filter out tests prior to day 250
    filter(
        event!="test" | time < 250,
        time < 1000
    ) |>
        select(-infected) |>
        arrange(patient,time) -> dat

trans_homog_objfun(
    params=c(
```

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```
a=1,b=0.1,gamma=0.01,p0=0.1,lambda.out=0.01,
   isol_factor=0.2,alpha=0.02,beta=0.1
  est=c("a","b"),
  data=dat
) -> f
f(\log(c(0.05,0.4)))
coef(f)
## Not run:
optim(
  par=log(c(0.5,0.4)),
  fn=f,
  control=list(reltol=1e-2)
) -> out
f(out$par)
coef(f)
## End(Not run)
trans_unit_spec_objfun(
  params=c(
   a=1,
   b.A=0.1,b.B=0.1,b.C=0.1,b.D=0.1,b.E=0.1,
   gamma=0.01,p0=0.1,
   lambda.out=0.1,gamma.out=0.01,
   isol_factor=0.2,alpha=0.02,beta=0.1
  ),
  data=dat
) -> f
f()
coef(f)
```

twostate

twostate

#### **Description**

Two-state Markov process with on and off rates

#### Usage

```
twostate(on, off, tf, t0 = 0)
```

### Arguments

```
on, off on and off rates
t0, tf initial and final times
```

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