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corr

corr

Description

Plot the correlation of the true data against the fitted resimulated data.

Usage

```
corr(sim)
```

Arguments

`sim` The dataframe or list produced by the `'runtsir'` function.

derivative	<i>derivative</i>
------------	-------------------

Description

This function computes an 8 point derivative.

Usage

```
derivative(X, Y)
```

Arguments

X	The variable to differentiate with respect to.
Y	The function / vector to differentiate.

epitimes	<i>epitimes</i>
----------	-----------------

Description

The times at which we declare a new outbreak has started based on the threshold parameter.

Usage

```
epitimes(data, threshold, epi.length = 3)
```

Arguments

data	The inputted data frame with the cases vector. This is the same data you put into runsir.
threshold	The required number of cases observed to declare it an outbreak.
epi.length	The required duration (in 52/IP weeks) to declare it an outbreak.

<code>estpars</code>	<i>estpars</i>
----------------------	----------------

Description

This function computes the set up to run the TSIR model, i.e. reconstructs susceptible and estimates beta and alpha. This can be plugged into `simulatetsir`.

Usage

```
estpars(
  data,
  xreg = "cumcases",
  IP = 2,
  seasonality = "standard",
  regtype = "gaussian",
  sigmamax = 3,
  family = "gaussian",
  link = "identity",
  userYhat = numeric(),
  alpha = NULL,
  sbar = NULL,
  printon = F
)
```

Arguments

<code>data</code>	The data frame containing cases and interpolated births and populations.
<code>xreg</code>	The x-axis for the regression. Options are 'cumcases' and 'cumbirths'. Defaults to 'cumcases'.
<code>IP</code>	The infectious period in weeks. This should be the same as your timestep. Defaults to 2 weeks.
<code>seasonality</code>	The type of contact to use. Options are standard for 52/IP point contact or schoolterm for just a two point on off contact or none for a single contact parameter. Defaults to standard.
<code>regtype</code>	The type of regression used in susceptible reconstruction. Options are 'gaussian', 'lm' (linear model), 'spline' (smooth.spline with 2.5 degrees freedom), 'lowess' (with $f = 2/3$, $iter = 1$), 'loess' (degree 1), and 'user' which is just a user inputted vector. Defaults to 'gaussian' and if that fails then defaults to loess.
<code>sigmamax</code>	The inverse kernel width for the gaussian regression. Default is 3. Smaller, stochastic outbreaks tend to need a lower sigma.
<code>family</code>	The family in the GLM regression. One can use any of the GLM ones, but the options are essentially 'poisson' (with <code>link='log'</code>), 'gaussian' (with <code>link='log'</code> or 'identity'), or 'quasipoisson' (with <code>link='log'</code>). Default is 'gaussian'.

link	The link function used with the glm family. Options are link='log' or 'identity'. Default is 'identity'. to include some bayesian approaches. For 'bayesglm' we use a gaussian prior with mean 1e-4.
userYhat	The inputted regression vector if regtype='user'. Defaults to NULL.
alpha	The mixing parameter. Defaults to NULL, i.e. the function estimates alpha.
sbar	The mean number of susceptibles. Defaults to NULL, i.e. the function estimates sbar.
printon	Whether to show diagnostic prints or not, defaults to FALSE.

Examples

```
## Not run:
require(kernlab)
London <- twentymeas[["London"]]
parms <- estpars(London)
names(parms)
sim <- simulatetsir(London,parms=parms,inits.fit=FALSE)
plotres(sim)

## End(Not run)
```

jagsfilter

jagsfilter

Description

Used internally to filter jags results to give just the inference well use.

Usage

```
jagsfilter(mcmcresults)
```

Arguments

mcmcresults is the input from the jags model.

logcorr	<i>logcorr</i>
---------	----------------

Description

Plot the correlation of the true data against the fitted resimulated data.

Usage

```
logcorr(sim)
```

Arguments

sim	The dataframe or list produced by the 'runsir' function.
-----	--

maxthreshold	<i>maxthreshold</i>
--------------	---------------------

Description

A function used to optimize the threshold parameter to give the best fit to the data. Optimizes the fit based on R squared.

Usage

```
maxthreshold(
  data,
  nsim = 2,
  IP = 2,
  method = "deterministic",
  inits.fit = FALSE,
  parms,
  thresholdmin = 2,
  thresholdmax = 20,
  printon = FALSE
)
```

Arguments

data	The time, cases, births, pop data frame.
nsim	The number of simulations to do.
IP	The infectious period, which should be the time step of the data.
method	The forward simulation method used, i.e. deterministic, negbin, pois.
inits.fit	Whether or not to fit initial conditions as well. Defaults to FALSE here. This parameter is more necessary in more chaotic locations.

parms	The estimated parameters from estpars or mcmcestpars.
thresholdmin	The minimum number of cases to be considered an outbreak.
thresholdmax	The max number of cases to be considered an outbreak.
printon	A T/F statement to print the progress.

Examples

```
require(kernlab)
Mold <- twenty meas[["Mold"]]
plotdata(Mold)
## Not run:
parms <- estpars(data=Mold,alpha=0.97)
tau <- maxthreshold(data=Mold,parms=parms,
  thresholdmin=8,thresholdmax=12,inits.fit=FALSE)
res <- simulatetsir(data=Mold,parms=parms,
  epidemics='break',threshold=tau,method='negbin',inits.fit=FALSE)
plotres(res)

## End(Not run)
```

mcmcestpars

mcmcestpars

Description

This function computes the set up to run the TSIR model, i.e. reconstructs susceptible and estimates beta and alpha using MCMC computations. Used the same way as estpars.

Usage

```
mcmcestpars(
  data,
  xreg = "cumcases",
  IP = 2,
  regtype = "gaussian",
  sigmax = 3,
  seasonality = "standard",
  userYhat = numeric(),
  update.iter = 10000,
  n.iter = 30000,
  n.chains = 3,
  n.adapt = 1000,
  burn.in = 100,
  sbar = NULL,
  alpha = NULL,
  printon = F
)
```

Arguments

<code>data</code>	The data frame containing cases and interpolated births and populations.
<code>xreg</code>	The x-axis for the regression. Options are 'cumcases' and 'cumbirths'. Defaults to 'cumcases'.
<code>IP</code>	The infectious period in weeks. Defaults to 2 weeks.
<code>regtype</code>	The type of regression used in susceptible reconstruction. Options are 'gaussian', 'lm' (linear model), 'spline' (smooth.spline with 2.5 degrees freedom), 'lowess' (with $f = 2/3$, $iter = 1$), 'loess' (degree 1), and 'user' which is just a user inputted vector. Defaults to 'gaussian' and if that fails then defaults to loess.
<code>sigmamax</code>	The inverse kernel width for the gaussian regression. Default is 3. Smaller, stochastic outbreaks tend to need a lower sigma.
<code>seasonality</code>	The type of contact to use. Options are standard for 52/IP point contact or schoolterm for just a two point on off contact or none for a single contact parameter. Defaults to standard.
<code>userYhat</code>	The inputted regression vector if <code>regtype='user'</code> . Defaults to NULL.
<code>update.iter</code>	Number of MCMC iterations to use in the update aspect. Default is 10000.
<code>n.iter</code>	Number of MCMC iterations to use. Default is 30000.
<code>n.chains</code>	Number of MCMC chains to use. Default is 3.
<code>n.adapt</code>	Adaptive number for MCMC. Default is 1000.
<code>burn.in</code>	Burn in number. Default is 100.
<code>sbar</code>	The mean number of susceptibles. Defaults to NULL, i.e. the function estimates sbar.
<code>alpha</code>	The mixing parameter. Defaults to NULL, i.e. the function estimates alpha.
<code>printon</code>	Whether to show diagnostic prints or not, defaults to FALSE.

mcmctsir

mcmctsir

Description

This function runs the TSIR model using a MCMC estimation. The susceptibles are still reconstructed in the same way as the regular tsir model, however beta, alpha, and sbar (or whatever combination you enter) are estimated using rjargs.

Usage

```
mcmctsir(
  data,
  xreg = "cumcases",
  IP = 2,
  nsim = 100,
  regtype = "gaussian",
```



```

    sigmamax = 3,
    userYhat = numeric(),
    update.iter = 10000,
    n.iter = 30000,
    n.chains = 3,
    n.adapt = 1000,
    burn.in = 100,
    method = "deterministic",
    epidemics = "cont",
    pred = "forward",
    seasonality = "standard",
    inits.fit = FALSE,
    threshold = 1,
    sbar = NULL,
    alpha = NULL,
    add.noise.sd = 0,
    mul.noise.sd = 0,
    printon = F
  )

```

Arguments

<code>data</code>	The data frame containing cases and interpolated births and populations.
<code>xreg</code>	The x-axis for the regression. Options are 'cumcases' and 'cumbirths'. Defaults to 'cumcases'.
<code>IP</code>	The infectious period in weeks. Defaults to 2 weeks.
<code>nsim</code>	The number of simulations to do. Defaults to 100.
<code>regtype</code>	The type of regression used in susceptible reconstruction. Options are 'gaussian', 'lm' (linear model), 'spline' (smooth.spline with 2.5 degrees freedom), 'lowess' (with $f = 2/3$, $iter = 1$), 'loess' (degree 1), and 'user' which is just a user inputted vector. Defaults to 'gaussian' and if that fails then defaults to loess.
<code>sigmamax</code>	The inverse kernel width for the gaussian regression. Default is 3. Smaller, stochastic outbreaks tend to need a lower sigma.
<code>userYhat</code>	The inputted regression vector if <code>regtype='user'</code> . Defaults to NULL.
<code>update.iter</code>	Number of MCMC iterations to use in the update aspect. Default is 10000.
<code>n.iter</code>	Number of MCMC iterations to use. Default is 30000.
<code>n.chains</code>	Number of MCMC chains to use. Default is 3.
<code>n.adapt</code>	Adaptive number for MCMC. Default is 1000.
<code>burn.in</code>	Burn in number. Default is 100.
<code>method</code>	The type of next step prediction used. Options are 'negbin' for negative binomial, 'pois' for poisson distribution, and 'deterministic'. Defaults to 'deterministic'.
<code>epidemics</code>	The type of data splitting. Options are 'cont' which doesn't split the data up at all, and 'break' which breaks the epidemics up if there are a lot of zeros. Defaults to 'cont'.

pred	The type of prediction used. Options are 'forward' and 'step-ahead'. Defaults to 'forward'.
seasonality	The type of contact to use. Options are standard for 52/IP point contact or schoolterm for just a two point on off contact or none for a single contact parameter. Defaults to standard.
inits.fit	Whether or not to fit initial conditions using simple least squares as well. Defaults to FALSE. This parameter is more necessary in more chaotic locations.
threshold	The cut off for a new epidemic if epidemics = 'break'. Defaults to 1.
sbar	The mean number of susceptibles. Defaults to NULL, i.e. the function estimates sbar.
alpha	The mixing parameter. Defaults to NULL, i.e. the function estimates alpha.
add.noise.sd	The sd for additive noise, defaults to zero.
mul.noise.sd	The sd for multiplicative noise, defaults to zero.
printon	Whether to show diagnostic prints or not, defaults to FALSE.

plotbeta	<i>plotbeta</i>
----------	-----------------

Description

Plots the inferred beta with confidence intervals (when they can be calculated)

Usage

```
plotbeta(dat)
```

Arguments

dat	the list produced from the runsir, mcmctsir, and simulatetsir function.
-----	---

plotbreaks	<i>plotbreaks</i>
------------	-------------------

Description

Plots the cases data with a line whenever the forward simulation is seeded using the real data.

Usage

```
plotbreaks(data, threshold)
```

Arguments

data	Data frame with the cases vector.
threshold	The epidemic threshold, i.e. the number of cases required to spark a new outbreak in the model.

plotcases	<i>plotcases</i>
-----------	------------------

Description

Plots just the cases data.

Usage

```
plotcases(data)
```

Arguments

data	The data frame with cases.
------	----------------------------

plotcomp	<i>plotcomp</i>
----------	-----------------

Description

Plots just the comparison of the forward simulation fit to the data.

Usage

```
plotcomp(sim, errtype = "95", max.plot = 10)
```

Arguments

sim	is list produced by runsir or memctsir
errtype	is the type of error bands to show. Defaults to '95' for 95 percent CI, the other option is 'sd' to standard deviation.
max.plot	the number of individual stochastic simulations to plot. Defaults to 10.

plotdata	<i>plotdata</i>
----------	-----------------

Description

Plots the cases data as well as birth and population dynamics.

Usage

```
plotdata(data)
```

Arguments

data	The dataframe with time, cases, births, and pop.
------	--

plotforward	<i>plotforward</i>
-------------	--------------------

Description

Plots the forward simulation from the TSIR model

Usage

```
plotforward(dat, inverse = F)
```

Arguments

dat	the list produced from the runsir, mcmtsir, and simulatetsir function.
inverse	a TRUE or FALSE option to plot the forward simulate negative (TRUE) or positive (FALSE). Defaults to FALSE.

plotLLE	<i>plotLLE</i>
---------	----------------

Description

Function to plot the Local Lyapunov Exponents. The output is of class ggplot2 so you can add standard ggplot2 options to it if desired.

Usage

```
plotLLE(LLE)
```

Arguments

LLE	The output from TSIR_LLE
-----	--------------------------

Examples

```
## Not run:
require(kernlab)
require(ggplot2)
require(kernlab)
London <- twentymeas$London
## just analyze the biennial portion of the data
London <- subset(London, time > 1950)

## define the interval to be 2 weeks
IP <- 2
```

```

## first estimate parameters from the London data
parms <- estpars(data=London, IP=2, regtype='gaussian',family='poisson',link='log')

## look at beta and alpha estimate
plotbeta(parms)

## simulate the fitted parameters
sim <- simulatetsir(data=London,parms=parms,IP=2,method='deterministic',nsim=2)

## now lets predict forward 200 years using the mean birth rate,
## starting from rough initial conditions
times <- seq(1965,2165, by = 1/ (52/IP))
births <- rep(mean(London$births),length(times))
S0 <- parms$sbar
I0 <- 1e-5*mean(London$pop)

pred <- predicttsir(times=times,births=births,
                    beta=parms$contact$beta,alpha=parms$alpha,
                    S0=S0,I0=I0,
                    nsim=50,stochastic=T)

## take the last 10 years
pred <- lapply(pred, function(x) tail(x, 52/IP * 20) )

## now compute the Lyapunov Exponent for the simulate and predicted model

simLE <- TSIR_LE(
  time=sim$res$time,
  S=sim$sim$mean,
  I=sim$res$mean,
  alpha=sim$alpha,
  beta=sim$contact$beta,
  IP=IP
)

predLE <- TSIR_LE(
  time=pred$I$time,
  S=pred$S$X3,
  I=pred$I$X3,
  alpha=parms$alpha,
  beta=parms$contact$beta,
  IP=IP
)

simLE$LE
predLE$LE

simLLE <- TSIR_LLE(simLE)
predLLE <- TSIR_LLE(predLE)

plotLLE(simLLE)
plotLLE(predLLE)

```

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

plotregression	<i>plotregression</i>
----------------	-----------------------

Description

Plots the cumulative cases - cumulative births data and regression fit

Usage

```
plotregression(dat)
```

Arguments

dat the list produced from the runtsir, mcmctsir, and simulatetsir function.

plotres	<i>plotres</i>
---------	----------------

Description

Plots diagnostics and results of the runtsir function.

Usage

```
plotres(dat, max.plot = 10)
```

Arguments

dat the list produced from the runtsir, mcmctsir, and simulatetsir function.
max.plot the number of individual stochastic simulations to plot. Defaults to 10.

plotrho	<i>plotrho</i>
---------	----------------

Description

Plots the inferred reporting rate, rho

Usage

```
plotrho(dat)
```

Arguments

dat the list produced from the runsir, mcmetsir, and simulatetsir function.

plotsbar	<i>plotsbar</i>
----------	-----------------

Description

Plots the profile log likelihood calculation for inferred sbar

Usage

```
plotsbar(dat)
```

Arguments

dat the list produced from the runsir, mcmetsir, and simulatetsir function.

predicttsir	<i>predicttsir</i>
-------------	--------------------

Description

function to predict incidence and susceptibles using the tsir model. This is different than simulatetsir as you are inputting parameters as vectors. The output is a data frame I and S with mean and confidence intervals of predictions.

Usage

```
predicttsir(times, births, beta, alpha, S0, I0, nsim, stochastic)
```

Arguments

times	The time vector to predict the model from. This assumes that the time step is equal to IP
births	The birth vector (of length length(times) or a single element) where each element is the births in that given (52/IP) time step
beta	The length(52/IP) beta vector of contact.
alpha	A single numeric which acts as the homogeneity parameter.
S0	The starting initial condition for S. This should be greater than one, i.e. not a fraction.
I0	The starting initial condition for I. This should be greater than one, i.e. not a fraction.
nsim	The number of simulations to perform.
stochastic	A TRUE / FALSE argument where FALSE is the deterministic model, and TRUE is a negative binomial distribution.

Examples

```
## Not run:
require(kernlab)
require(ggplot2)
require(kernlab)
require(tsiR)
London <- twentymeas$London

London <- subset(London, time > 1950)

IP <- 2
## first estimate paramters from the London data
parms <- estpars(data=London, IP=2, regtype='gaussian')

plotbeta(parms)

## now lets predict forward 20 years using the mean birth rate,
## starting from rough initial conditions
births <- min(London$births)
times <- seq(1965,1985, by = 1/ (52/IP))
S0 <- parms$sbar
I0 <- 1e-5*mean(London$pop)

pred <- predicttsir(times=times,births=births,
                   beta=parms$contact$beta,alpha=parms$alpha,
                   S0=S0,I0=I0,
                   nsim=50,stochastic=T)

## plot this prediction
ggplot(pred$I,aes(time,mean))+geom_line()+geom_ribbon(aes(ymin=low,ymax=high),alpha=0.3)
```



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

residual.births	<i>residuals.births</i>
-----------------	-------------------------

Description

computes the residuals for when X is the cumulative births. Used internally.

Usage

```
residual.births(rho, Yhat, Y)
```

Arguments

rho	The reporting rate, used to get units correct.
Yhat	The fitted regression line.
Y	The cumulative cases.

residual.cases	<i>residuals.cases</i>
----------------	------------------------

Description

Computes the residuals for when X is the cumulative cases. Used internally.

Usage

```
residual.cases(Yhat, Y)
```

Arguments

Yhat	The fitted regression line.
Y	The cumulative births.

 runtsir

runtsir

Description

This function runs the TSIR model.

Usage

```
runtsir(
  data,
  xreg = "cumcases",
  IP = 2,
  nsim = 10,
  regtype = "gaussian",
  sigmamax = 3,
  userYhat = numeric(),
  alpha = NULL,
  sbar = NULL,
  family = "gaussian",
  link = "identity",
  method = "deterministic",
  inits.fit = FALSE,
  epidemics = "cont",
  pred = "forward",
  threshold = 1,
  seasonality = "standard",
  add.noise.sd = 0,
  mul.noise.sd = 0,
  printon = F,
  fit = NULL,
  fittype = NULL
)
```

Arguments

data	The data frame containing cases and interpolated births and populations.
xreg	The x-axis for the regression. Options are 'cumcases' and 'cumbirths'. Defaults to 'cumcases'.
IP	The infectious period in weeks. Defaults to 2 weeks.
nsim	The number of simulations to do. Defaults to 100.
regtype	The type of regression used in susceptible reconstruction. Options are 'gaussian', 'lm' (linear model), 'spline' (smooth.spline with 2.5 degrees freedom), 'lowess' (with $f = 2/3$, $iter = 1$), 'loess' (degree 1), and 'user' which is just a user inputted vector. Defaults to 'gaussian' and if that fails then defaults to loess.

sigmamax	The inverse kernel width for the gaussian regression. Default is 3. Smaller, stochastic outbreaks tend to need a lower sigma.
userYhat	The inputted regression vector if regtype='user'. Defaults to NULL.
alpha	The mixing parameter. Defaults to NULL, i.e. the function estimates alpha.
sbar	The mean number of susceptibles. Defaults to NULL, i.e. the function estimates sbar.
family	The family in the GLM regression. One can use any of the GLM ones, but the options are essentially 'poisson' (with link='log'), 'gaussian' (with link='log' or 'identity'), or 'quasipoisson' (with link='log'). Default is 'gaussian'.
link	The link function used with the glm family. Options are link='log' or 'identity'. Default is 'identity'.
method	The type of next step prediction used. Options are 'negbin' for negative binomial, 'pois' for poisson distribution, and 'deterministic'. Defaults to 'deterministic'.
inits.fit	Whether or not to fit initial conditions using simple least squares as well. Defaults to FALSE. This parameter is more necessary in more chaotic locations.
epidemics	The type of data splitting. Options are 'cont' which doesn't split the data up at all, and 'break' which breaks the epidemics up if there are a lot of zeros. Defaults to 'cont'.
pred	The type of prediction used. Options are 'forward' and 'step-ahead'. Defaults to 'forward'.
threshold	The cut off for a new epidemic if epidemics = 'break'. Defaults to 1.
seasonality	The type of contact to use. Options are standard for 52/IP point contact or schoolterm for just a two point on off contact, or none for a single contact parameter. Defaults to standard.
add.noise.sd	The sd for additive noise, defaults to zero.
mul.noise.sd	The sd for multiplicative noise, defaults to zero.
printon	Whether to show diagnostic prints or not, defaults to FALSE.
fit	Now removed but gives a warning.
fitype	Now removed but gives a warning.

Examples

```
require(kernlab)
London <- twentymeas[["London"]]
## Not run:
plotdata(London)
res <- runtsir(data=London,method='pois',nsim=10, IP=2,inits.fit=FALSE)
plotres(res)

## End(Not run)
```

 simulatetsir

simulatetsir

Description

This function just simulates the forward prediction given the data and a parms list generated from estpars or mcmcestpars.

Usage

```

simulatetsir(
  data,
  nsim = 100,
  IP = 2,
  parms,
  method = "deterministic",
  epidemics = "cont",
  pred = "forward",
  threshold = 1,
  inits.fit = FALSE,
  add.noise.sd = 0,
  mul.noise.sd = 0
)

```

Arguments

data	The data frame containing cases and interpolated births and populations.
nsim	The number of simulations to do. Defaults to 100.
IP	The infectious period. Defaults to 2.
parms	Either the parameters estimated by estpars or mcmcestpars, or a list containing beta, rho, Z, sbar, alpha, X, Y, Yhat, contact, alphalow, alphahigh, loglik, pop vectors.
method	The type of next step prediction used. Options are 'negbin' for negative binomial, 'pois' for poisson distribution, and 'deterministic'. Defaults to 'deterministic'.
epidemics	The type of data splitting. Options are 'cont' which doesn't split the data up at all, and 'break' which breaks the epidemics up if there are a lot of zeros. Defaults to 'cont'.
pred	The type of prediction used. Options are 'forward' and 'step-ahead'. Defaults to 'forward'.
threshold	The cut off for a new epidemic if epidemics = 'break'. Defaults to 1.
inits.fit	Whether or not to fit initial conditions using simple least squares as well. Defaults to FALSE. This parameter is more necessary in more chaotic locations.
add.noise.sd	The sd for additive noise, defaults to zero.
mul.noise.sd	The sd for multiplicative noise, defaults to zero.

tsiRdata	<i>tsiRdata</i>
----------	-----------------

Description

A function to take in time cases births and pop vectors (of any lengths) and interpolate them using the given infectious period.

Usage

```
tsiRdata(time, cases, births, pop, IP = 2)
```

Arguments

time	The time vector.
cases	The cases vector.
births	The births vector.
pop	The population vector.
IP	The infectious period (in weeks) to discretize to. Defaults to 2.

TSIR_LE	<i>TSIR_LE</i>
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Description

A function to calculate the Lyapunov Exponent (LE) from the TSIR model

Usage

```
TSIR_LE(time, S, I, alpha, beta, IP)
```

Arguments

time	The time vector from the data or simulated data
S	The S output from the simulated or predicted TSIR model
I	The I output from the simulated or predicted TSIR model
alpha	The homogeneity parameter from the simulated or predicted TSIR model
beta	The inferred contact rate, use <code>beta = contact\$beta</code> where <code>contact</code> is an output from <code>runtsir</code> or <code>simulatetsir</code>
IP	The generation interval of the pathogen (in weeks)

Examples

```

## Not run:
require(kernlab)
require(ggplot2)
require(kernlab)
London <- twentymeas$London
## just analyze the biennial portion of the data
London <- subset(London, time > 1950)

## define the interval to be 2 weeks
IP <- 2

## first estimate paramters from the London data
parms <- estpars(data=London, IP=2, regtype='gaussian',family='poisson',link='log')

## look at beta and alpha estimate
plotbeta(parms)

## simulate the fitted parameters
sim <- simulatetsir(data=London,parms=parms,IP=2,method='deterministic',nsim=2)

## now lets predict forward 200 years using the mean birth rate,
## starting from rough initial conditions
times <- seq(1965,2165, by = 1/ (52/IP))
births <- rep(mean(London$births),length(times))
S0 <- parms$sbar
I0 <- 1e-5*mean(London$pop)

pred <- predicttsir(times=times,births=births,
                    beta=parms$contact$beta,alpha=parms$alpha,
                    S0=S0,I0=I0,
                    nsim=50,stochastic=T)

## take the last 10 years
pred <- lapply(pred, function(x) tail(x, 52/IP * 20) )

## now compute the Lyapunov Exponent for the simulate and predicted model

simLE <- TSIR_LE(
  time=sim$res$time,
  S=sim$simS$mean,
  I=sim$res$mean,
  alpha=sim$alpha,
  beta=sim$contact$beta,
  IP=IP
)

predLE <- TSIR_LE(
  time=pred$I$time,
  S=pred$S$X3,
  I=pred$I$X3,
  alpha=parms$alpha,

```

```

beta=parms$contact$beta,
IP=IP
)

simLE$LE
predLE$LE

## End(Not run)

```

TSIR_LLE

TSIR_LLE

Description

A function to calculate the Local Lyapunov Exponent (LLE) from the TSIR model

Usage

```
TSIR_LLE(LE, m = 1)
```

Arguments

LE	The output of TSIR_LE to pass the Jacobian elements
m	The window to sweep the time-varying Jacobian elements. Defaults to one.

Examples

```

## Not run:
require(kernlab)
require(ggplot2)
require(kernlab)
London <- twentymeas$London
## just analyze the biennial portion of the data
London <- subset(London, time > 1950)

## define the interval to be 2 weeks
IP <- 2

## first estimate parameters from the London data
parms <- estpars(data=London, IP=2, regtype='gaussian',family='poisson',link='log')

## look at beta and alpha estimate
plotbeta(parms)

## simulate the fitted parameters
sim <- simulatetsir(data=London,parms=parms,IP=2,method='deterministic',nsim=2)

```

```

## now lets predict forward 200 years using the mean birth rate,
## starting from rough initial conditions
times <- seq(1965,2165, by = 1/ (52/IP))
births <- rep(mean(London$births),length(times))
S0 <- parms$sbar
I0 <- 1e-5*mean(London$pop)

pred <- predicttsir(times=times,births=births,
                    beta=parms$contact$beta,alpha=parms$alpha,
                    S0=S0,I0=I0,
                    nsim=50,stochastic=T)

## take the last 10 years
pred <- lapply(pred, function(x) tail(x, 52/IP * 20) )

## now compute the Lyapunov Exponent for the simulate and predicted model

simLE <- TSIR_LE(
  time=sim$res$time,
  S=sim$simS$mean,
  I=sim$res$mean,
  alpha=sim$alpha,
  beta=sim$contact$beta,
  IP=IP
)

predLE <- TSIR_LE(
  time=pred$I$time,
  S=pred$S$X3,
  I=pred$I$X3,
  alpha=parms$alpha,
  beta=parms$contact$beta,
  IP=IP
)

simLE$LE
predLE$LE

simLLE <- TSIR_LLE(simLE)
predLLE <- TSIR_LLE(predLE)

plotLLE(simLLE)
plotLLE(predLLE)

## End(Not run)

```


Description

twentymeas is a list containing 20 dataframes with cases, births, populations. Each dataframe is a 22 year time series at biweekly (i.e. IP=2) intervals.

Usage

```
data("twentymeas")
```

Source

From Bryan Grenfell

Examples

```
names(twentymeas)
london <- twentymeas[["London"]]
plotdata(london)
```

vlayout

vlayout

Description

the function just breaks up the plot area into a grid. Called internally.

Usage

```
vlayout(x, y)
```

Arguments

x is the x location of the plot
y is the y lcoation of the ploy

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