Package 'pomp'

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pomp-package

Inference for partially observed Markov processes

Description

The **pomp** package provides facilities for inference on time series data using partially-observed Markov process (POMP) models. These models are also known as state-space models, hidden Markov models, or nonlinear stochastic dynamical systems. One can use **pomp** to fit nonlinear, non-Gaussian dynamic models to time-series data. The package is both a set of tools for data analysis and a platform upon which statistical inference methods for POMP models can be implemented.

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Data analysis using pomp

pomp provides algorithms for:

- 1. Simulation of stochastic dynamical systems; see simulate.
- 2. Particle filtering (AKA sequential Monte Carlo or sequential importance sampling); see pfilter and wpfilter.
- 3. The iterated filtering methods of Ionides et al. (2006, 2011, 2015); see mif2.
- 4. The nonlinear forecasting algorithm of Kendall et al. (2005); see nlf.
- 5. The particle MCMC approach of Andrieu et al. (2010); see pmcmc.
- 6. The probe-matching method of Kendall et al. (1999, 2005); see probe match.
- 7. Synthetic likelihood a la Wood (2010); see probe.
- 8. A spectral probe-matching method (Reuman et al. 2006, 2008); see spect_match.
- 9. Approximate Bayesian computation (Toni et al. 2009); see abc.
- 10. The approximate Bayesian sequential Monte Carlo scheme of Liu & West (2001); see bsmc2.
- 11. Ensemble and ensemble adjusted Kalman filters; see kalman.
- 12. Simple trajectory matching; see traj_match.

The package also provides various tools for plotting and extracting information on models and data.

Structure of the package

pomp algorithms are arranged into several levels. At the top level, estimation algorithms estimate model parameters and return information needed for other aspects of inference. Elementary algorithms perform common operations on POMP models, including simulation, filtering, and application of diagnostic probes; these functions may be useful in inference, but they do not themselves perform estimation. At the lowest level, workhorse functions provide the interface to basic POMP model components. Beyond these, pomp provides a variety of auxiliary functions for manipulating and extracting information from 'pomp' objects, producing diagnostic plots, facilitating reproducible computations, and so on.

Implementing a model

The basic structure at the heart of the package is the 'pomp object'. This is a container holding a time series of data (possibly multivariate) and a model. The model is specified by specifying some or all of its basic model components. One does this using the basic component arguments to the pomp constructor. One can also add, modify, or delete basic model components "on the fly" in any **pomp** function that accepts them.

Documentation and examples

The package contains a number of examples. Some of these are included in the help pages. In addition, several pre-built POMP models are included with the package. Tutorials and other documentation, including a package FAQ, are available from the package website.

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Useful links

- pomp homepage: https://kingaa.github.io/pomp/
- Report bugs to: https://github.com/kingaa/pomp/issues
- Frequently asked questions: https://kingaa.github.io/pomp/FAQ.html
- User guides and tutorials: https://kingaa.github.io/pomp/docs.html
- pomp news: https://kingaa.github.io/pomp/blog.html

Citing pomp

Execute citation("pomp") to view the correct citation for publications.

Author(s)

Aaron A. King

References

A. A. King, D. Nguyen, and E. L. Ionides. Statistical inference for partially observed Markov processes via the R package **pomp**. *Journal of Statistical Software* **69**(12), 1–43, 2016. doi:10.18637/jss.v069.i12. An updated version of this paper is available on the **pomp** package website.

See the package website for more references, including many publications that use pomp.

See Also

Useful links:

- https://kingaa.github.io/pomp/
- Report bugs at https://github.com/kingaa/pomp/issues/

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

More on **pomp** workhorse functions: dinit(), dmeasure(), dprior(), dprocess(), emeasure(), flow(), partrans(), rinit(), rmeasure(), rprior(), rprocess(), skeleton(), vmeasure(), workhorses

More on **pomp** estimation algorithms: abc(), bsmc2(), estimation_algorithms, mif2(), nlf, pmcmc(), probe_match, spect_match

More on **pomp** elementary algorithms: elementary_algorithms, kalman, pfilter(), probe(), simulate(), spect(), trajectory(), wpfilter()

6 abc

abc

Approximate Bayesian computation

Description

The approximate Bayesian computation (ABC) algorithm for estimating the parameters of a partially-observed Markov process.

Usage

```
## S4 method for signature 'data.frame'
abc(
  data,
  . . . ,
 Nabc = 1,
 proposal,
  scale,
  epsilon,
  probes,
  params,
  rinit,
  rprocess,
  rmeasure,
 dprior,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
abc(
  data,
  ...,
 Nabc = 1,
 proposal,
  scale,
 epsilon,
 probes,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'probed_pomp'
abc(data, ..., probes, verbose = getOption("verbose", FALSE))
## S4 method for signature 'abcd_pomp'
abc(
  data,
  . . . ,
 Nabc,
```

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```
proposal,
scale,
epsilon,
probes,
verbose = getOption("verbose", FALSE)
```

Arguments

data either a data frame holding the time series data, or an object of class 'pomp',

i.e., the output of another \boldsymbol{pomp} calculation. Internally, data will be coerced to

an array with storage-mode double.

... additional arguments are passed to pomp. This allows one to set, unset, or modify

basic model components within a call to this function.

Nabc the number of ABC iterations to perform.

proposal optional function that draws from the proposal distribution. Currently, the pro-

posal distribution must be symmetric for proper inference: it is the user's responsibility to ensure that it is. Several functions that construct appropriate proposal

function are provided: see MCMC proposals for more information.

scale named numeric vector of scales.

epsilon ABC tolerance.

probes a single probe or a list of one or more probes. A probe is simply a scalar- or

vector-valued function of one argument that can be applied to the data array of a 'pomp'. A vector-valued probe must always return a vector of the same size. A number of useful probes are provided with the package: see basic probes.

params optional; named numeric vector of parameters. This will be coerced internally

to storage mode double.

rinit simulator of the initial-state distribution. This can be furnished either as a C

snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator

to its default. For more information, see rinit specification.

rprocess simulator of the latent state process, specified using one of the rprocess plugins.

Setting rprocess=NULL removes the latent-state simulator. For more informa-

tion, see rprocess specification for the documentation on these plugins.

rmeasure simulator of the measurement model, specified either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically

loaded library. Setting rmeasure=NULL removes the measurement model simu-

lator. For more information, see rmeasure specification.

dprior optional; prior distribution density evaluator, specified either as a C snippet,

an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. For more information, see prior specification. Setting dprior=NULL resets the prior distribution to its default, which is a flat improper

prior.

verbose logical; if TRUE, diagnostic messages will be printed to the console.

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Running ABC

abc returns an object of class 'abcd_pomp'. One or more 'abcd_pomp' objects can be joined to form an 'abcList' object.

Re-running ABC iterations

To re-run a sequence of ABC iterations, one can use the abc method on a 'abcd_pomp' object. By default, the same parameters used for the original ABC run are re-used (except for verbose, the default of which is shown above). If one does specify additional arguments, these will override the defaults.

Continuing ABC iterations

One can continue a series of ABC iterations from where one left off using the continue method. A call to abc to perform Nabc=m iterations followed by a call to continue to perform Nabc=n iterations will produce precisely the same effect as a single call to abc to perform Nabc=m+n iterations. By default, all the algorithmic parameters are the same as used in the original call to abc. Additional arguments will override the defaults.

Methods

The following can be applied to the output of an abc operation:

abc repeats the calculation, beginning with the last state

continue continues the abc calculation

plot produces a series of diagnostic plots

traces produces an mcmc object, to which the various coda convergence diagnostics can be applied

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Author(s)

Edward L. Ionides, Aaron A. King

References

- J.-M. Marin, P. Pudlo, C. P. Robert, and R. J. Ryder. Approximate Bayesian computational methods. *Statistics and Computing* **22**, 1167–1180, 2012. doi:10.1007/s1122201192882.
- T. Toni and M. P. H. Stumpf. Simulation-based model selection for dynamical systems in systems and population biology. *Bioinformatics* **26**, 104–110, 2010. doi:10.1093/bioinformatics/btp619.
- T. Toni, D. Welch, N. Strelkowa, A. Ipsen, and M. P. H. Stumpf. Approximate Bayesian computation scheme for parameter inference and model selection in dynamical systems. *Journal of the Royal Society Interface* **6**, 187–202, 2009. doi:10.1098/rsif.2008.0172.

accumvars 9

See Also

```
More on methods based on summary statistics: basic_probes, nlf, probe(), probe_match, spect(), spect_match

More on pomp estimation algorithms: bsmc2(), estimation_algorithms, mif2(), nlf, pmcmc(), pomp-package, probe_match, spect_match

More on Markov chain Monte Carlo methods: pmcmc(), proposals

More on Bayesian methods: bsmc2(), dprior(), pmcmc(), prior_spec, rprior()
```

accumvars

accumulator variables

Description

Latent state variables that accumulate quantities through time.

Details

In formulating models, one sometimes wishes to define a state variable that will accumulate some quantity over the interval between successive observations. **pomp** provides a facility to make such features more convenient. Specifically, if a is a state-variable named in the pomp's accumvars argument, then for each interval $(t_k, t_{k+1}), k = 0, 1, 2, \ldots, a$ will be set to zero at prior to any rprocess computation over that interval. Deterministic trajectory computation is handled slightly differently: see flow. For examples, see sir and the tutorials on the package website.

See Also

sir

More on implementing POMP models: Csnippet, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

Examples

10 accumvars

```
// gamma noise, mean=dt, variance=(sigma^2 dt)
      dW = rgammawn(sigma,dt);
      // compute the transition rates
      rate[0] = mu*pop;
                           // birth into susceptible class
      rate[1] = (iota+Beta*I*dW/dt)/pop; // force of infection
      rate[2] = mu;
                            // death from susceptible class
      rate[3] = gamma;
                            // recovery
                            // death from infectious class
      rate[4] = mu;
      rate[5] = mu;
                            // death from recovered class
      // compute the transition numbers
      trans[0] = rpois(rate[0]*dt); // births are Poisson
      reulermultinom(2,S,&rate[1],dt,&trans[1]);
      reulermultinom(2,I,&rate[3],dt,&trans[3]);
      reulermultinom(1,R,&rate[5],dt,&trans[5]);
      // balance the equations
      S += trans[0]-trans[1]-trans[2];
     I += trans[1]-trans[3]-trans[4];
     R += trans[3]-trans[5];
    "),
    delta.t=1/52/20
    rinit=Csnippet("
     double m = pop/(S_0+I_0+R_0);
      S = nearbyint(m*S_0);
     I = nearbyint(m*I_0);
      R = nearbyint(m*R_0);
  "),
  paramnames=c("mu","pop","iota","gamma","Beta","sigma",
    "S_0","I_0","R_0"),
  statenames=c("S","I","R"),
  params=c(mu=1/50,iota=10,pop=50e6,gamma=26,Beta=400,sigma=0.1,
    S_0=0.07, I_0=0.001, R_0=0.93)
  ) -> ew1
ew1 |>
  simulate() |>
 plot(variables=c("S","I","R"))
## A simple SIR model that tracks cumulative incidence.
ew1 |>
  pomp(
    rprocess=euler(
      Csnippet("
      int nrate = 6;
      double rate[nrate];
                            // transition rates
      double trans[nrate];  // transition numbers
      double dW;
      // gamma noise, mean=dt, variance=(sigma^2 dt)
```

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```
dW = rgammawn(sigma,dt);
      // compute the transition rates
      rate[0] = mu*pop;
                           // birth into susceptible class
      rate[1] = (iota+Beta*I*dW/dt)/pop; // force of infection
      rate[2] = mu;  // death from susceptible class
      rate[3] = gamma;
                            // recovery
      rate[4] = mu;
                             // death from infectious class
                             // death from recovered class
      rate[5] = mu;
      // compute the transition numbers
      trans[0] = rpois(rate[0]*dt); // births are Poisson
      reulermultinom(2,S,&rate[1],dt,&trans[1]);
      reulermultinom(2,I,&rate[3],dt,&trans[3]);
      reulermultinom(1,R,&rate[5],dt,&trans[5]);
      // balance the equations
      S += trans[0]-trans[1]-trans[2];
      I += trans[1]-trans[3]-trans[4];
     R += trans[3]-trans[5];
     H += trans[3];
                              // cumulative incidence
    "),
    delta.t=1/52/20
    rmeasure=Csnippet("
      double mean = H*rho;
      double size = 1/tau;
      reports = rnbinom_mu(size, mean);
  "),
  rinit=Csnippet("
     double m = pop/(S_0+I_0+R_0);
      S = nearbyint(m*S_0);
     I = nearbyint(m*I_0);
      R = nearbyint(m*R_0);
     H = 0;
  "),
  paramnames=c("mu","pop","iota","gamma","Beta","sigma","tau","rho",
    "S_0","I_0","R_0"),
  {\tt statenames=c("S","I","R","H"),}
  params=c(mu=1/50,iota=10,pop=50e6,gamma=26,
    Beta=400, sigma=0.1, tau=0.001, rho=0.6,
    S_0=0.07, I_0=0.001, R_0=0.93)
  ) -> ew2
ew2 |>
  simulate() |>
  plot()
## A simple SIR model that tracks weekly incidence.
ew2 |>
  pomp(accumvars="H") -> ew3
```

12 as.data.frame

```
ew3 |>
  simulate() |>
  plot()
```

as.data.frame

Coerce to data frame

Description

All **pomp** model objects can be recast as data frames. The contents of the resulting data frame depend on the nature of the object.

Usage

```
## S3 method for class 'pomp'
as.data.frame(x, ...)
## S3 method for class 'pfilterd_pomp'
as.data.frame(x, ...)
## S3 method for class 'probed_pomp'
as.data.frame(x, ...)
## S3 method for class 'kalmand_pomp'
as.data.frame(x, ...)
## S3 method for class 'bsmcd_pomp'
as.data.frame(x, ...)
## S3 method for class 'pompList'
as.data.frame(x, ...)
## S3 method for class 'pfilterList'
as.data.frame(x, ...)
## S3 method for class 'abcList'
as.data.frame(x, ...)
## S3 method for class 'mif2List'
as.data.frame(x, ...)
## S3 method for class 'pmcmcList'
as.data.frame(x, ...)
## S3 method for class 'wpfilterd_pomp'
as.data.frame(x, ...)
```

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Arguments

x any R object.

... additional arguments to be passed to or from methods.

Details

When object is a simple 'pomp' object, as(object, "data.frame") or as.data.frame(object) results in a data frame with the times, observables, states (if known), and interpolated covariates (if any).

When object is a 'pfilterd_pomp' object, coercion to a data frame results in a data frame with the same content as for a simple 'pomp', but with conditional log likelihood and effective sample size estimates included, as well as filtering means, prediction means, and prediction variances, if these have been computed.

When object is a 'probed_pomp' object, coercion to a data frame results in a data frame with the values of the probes computed on the data and on simulations.

When object is a 'kalmand_pomp' object, coercion to a data frame results in a data frame with prediction means, filter means and forecasts, in addition to the data.

When object is a 'bsmcd_pomp' object, coercion to a data frame results in a data frame with samples from the prior and posterior distribution. The .id variable distinguishes them.

When object is a 'wpfilterd_pomp' object, coercion to a data frame results in a data frame with the same content as for a simple 'pomp', but with conditional log likelihood and effective sample size estimates included.

as_pomp as.pomp

as_pollip

Description

```
Coerce to a 'pomp' object
```

Usage

```
as_pomp(object, ...)
```

Arguments

object the object to be coerced additional arguments

14 basic_components

basic_components

Basic POMP model components.

Description

Mathematically, the parts of a POMP model include the latent-state process transition distribution, the measurement-process distribution, the initial-state distribution, and possibly a prior parameter distribution. Algorithmically, each of these corresponds to at least two distinct operations. In particular, for each of the above parts, one sometimes needs to make a random draw from the distribution and sometimes to evaluate the density function. Accordingly, for each such component, there are two basic model components, one prefixed by a 'r', the other by a 'd', following the usual R convention.

Details

In addition to the parts listed above, **pomp** includes two additional basic model components: the deterministic skeleton, and parameter transformations that can be used to map the parameter space onto a Euclidean space for estimation purposes. There are also basic model components for computing the mean and variance of the measurement process conditional on the latent-state process.

There are thus altogether twelve **basic model components**:

- 1. rprocess, which samples from the latent-state transition distribution,
- 2. dprocess, which evaluates the latent-state transition density,
- 3. rmeasure, which samples from the measurement distribution,
- 4. emeasure, which computes the conditional expectation of the measurements, given the latent states.
- vmeasure, which computes the conditional covariance matrix of the measurements, given the latent states,
- 6. dmeasure, which evaluates the measurement density,
- 7. rprior, which samples from the prior distribution,
- 8. dprior, which evaluates the prior density,
- 9. rinit, which samples from the initial-state distribution,
- 10. dinit, which evaluates the initial-state density,
- 11. skeleton, which evaluates the deterministic skeleton,
- 12. partrans, which evaluates the forward or inverse parameter transformations.

Each of these can be set or modified in the pomp constructor function or in any of the **pomp** elementary algorithms or estimation algorithms using an argument that matches the basic model component. A basic model component can be unset by passing NULL in the same way.

Help pages detailing each basic model component are provided.

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See Also

workhorse functions, elementary algorithms, estimation algorithms.

More on implementing POMP models: Csnippet, accumvars, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

basic_probes

Useful probes for partially-observed Markov processes

Description

Several simple and configurable probes are provided with in the package. These can be used directly and as templates for custom probes.

Usage

```
probe_mean(var, trim = 0, transform = identity, na.rm = TRUE)
probe_median(var, na.rm = TRUE)
probe_var(var, transform = identity, na.rm = TRUE)
probe_sd(var, transform = identity, na.rm = TRUE)
probe_period(var, kernel.width, transform = identity)
probe_quantile(var, probs, ...)
probe_acf(
  var,
  lags,
  type = c("covariance", "correlation"),
  transform = identity
)
probe_ccf(
  vars,
  lags,
  type = c("covariance", "correlation"),
  transform = identity
)
probe_marginal(var, ref, order = 3, diff = 1, transform = identity)
probe_nlar(var, lags, powers, transform = identity)
```

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Arguments

var, vars character; the name(s) of the observed variable(s).

trim the fraction of observations to be trimmed (see mean).

transform transformation to be applied to the data before the probe is computed.

na.rm if TRUE, remove all NA observations prior to computing the probe.

kernel.width width of modified Daniell smoothing kernel to be used in power-spectrum com-

putation: see kernel.

probs the quantile or quantiles to compute: see quantile.

. . . additional arguments passed to the underlying algorithms.

lags In probe_ccf, a vector of lags between time series. Positive lags correspond to

x advanced relative to y; negative lags, to the reverse.

In probe_nlar, a vector of lags present in the nonlinear autoregressive model that will be fit to the actual and simulated data. See Details, below, for a precise

description.

type Compute autocorrelation or autocovariance?

ref empirical reference distribution. Simulated data will be regressed against the

values of ref, sorted and, optionally, differenced. The resulting regression coefficients capture information about the shape of the marginal distribution. A

good choice for ref is the data itself.

order order of polynomial regression.

diff order of differencing to perform.

powers the powers of each term (corresponding to lags) in the the nonlinear autoregres-

sive model that will be fit to the actual and simulated data. See Details, below,

for a precise description.

Value

A call to any one of these functions returns a probe function, suitable for use in probe or probe_objfun. That is, the function returned by each of these takes a data array (such as comes from a call to obs) as input and returns a single numerical value.

Author(s)

Daniel C. Reuman, Aaron A. King

References

B.E. Kendall, C.J. Briggs, W.W. Murdoch, P. Turchin, S.P. Ellner, E. McCauley, R.M. Nisbet, and S.N. Wood. Why do populations cycle? A synthesis of statistical and mechanistic modeling approaches. *Ecology* **80**, 1789–1805, 1999. doi:10.2307/176658.

S. N. Wood Statistical inference for noisy nonlinear ecological dynamic systems. *Nature* **466**, 1102–1104, 2010. doi:10.1038/nature09319.

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See Also

More on methods based on summary statistics: abc(), nlf, probe(), probe_match, spect(), spect_match

betabinomial

Beta-binomial distribution

Description

Density and random generation for the Beta-binomial distribution with parameters size, mu, and theta.

Usage

```
rbetabinom(n = 1, size, prob, theta)
dbetabinom(x, size, prob, theta, log = FALSE)
```

Arguments

n	integer; number of random variates to generate.
size	size parameter of the binomial distribution
prob	mean of the Beta distribution
theta	Beta distribution dispersion parameter
x	vector of non-negative integer quantiles

Details

log

A variable X is Beta-binomially distributed if $X \sim \operatorname{Binomial}(n, P)$ where $P \sim \operatorname{Beta}(\mu, \theta)$. Using the standard (a, b) parameterization, $a = \mu \theta$ and $b = (1 - \mu) \theta$.

logical; if TRUE, return logarithm(s) of probabilities.

Value

rbetabinom	Returns a vector of length \boldsymbol{n} containing random variates drawn from the Betabinomial distribution.
dbetabinom	Returns a vector (of length equal to the number of columns of x) containing the probabilities of observing each column of x given the specified parameters (size, prob, theta).

C API

An interface for C codes using these functions is provided by the package. Visit the package homepage to view the **pomp** C API document.

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See Also

More on implementing POMP models: Csnippet, accumvars, basic_components, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

blowflies

Nicholson's blowflies.

Description

blowflies is a data frame containing the data from several of Nicholson's classic experiments with the Australian sheep blowfly, *Lucilia cuprina*.

Usage

```
blowflies1(

P = 3.2838,

delta = 0.16073,

N0 = 679.94,

sigma.P = 1.3512,

sigma.d = 0.74677,

sigma.y = 0.026649

)

blowflies2(

P = 2.7319,

delta = 0.17377,

N0 = 800.31,

sigma.P = 1.442,

sigma.d = 0.76033,

sigma.y = 0.010846

)
```

Arguments

Р	reproduction parameter
delta	death rate
NØ	population scale factor
sigma.P	intensity of e noise
sigma.d	intensity of eps noise
sigma.v	measurement error s.d.

blowflies 19

Details

blowflies1() and blowflies2() construct 'pomp' objects encoding stochastic delay-difference equation models. The data for these come from "population I", a control culture. The experiment is described on pp. 163–4 of Nicholson (1957). Unlimited quantities of larval food were provided; the adult food supply (ground liver) was constant at 0.4g per day. The data were taken from the table provided by Brillinger et al. (1980).

The models are discrete delay equations:

$$R(t+1) \sim \text{Poisson}(PN(t-\tau) \exp{(-N(t-\tau)/N_0)}e(t+1)\Delta t)$$

 $S(t+1) \sim \text{Binomial}(N(t), \exp{(-\delta\epsilon(t+1)\Delta t)})$
 $N(t) = R(t) + S(t)$

where e(t) and $\epsilon(t)$ are Gamma-distributed i.i.d. random variables with mean 1 and variances $\sigma_P^2/\Delta t$, $\sigma_d^2/\Delta t$, respectively. blowflies1 has a timestep (Δt) of 1 day; blowflies2 has a timestep of 2 days. The process model in blowflies1 thus corresponds exactly to that studied by Wood (2010). The measurement model in both cases is taken to be

$$y(t) \sim \text{NegBin}(N(t), 1/\sigma_y^2)$$

i.e., the observations are assumed to be negative-binomially distributed with mean N(t) and variance $N(t) + (\sigma_u N(t))^2$.

Default parameter values are the MLEs as estimated by Ionides (2011).

Value

blowflies1 and blowflies2 return 'pomp' objects containing the actual data and two variants of the model.

References

A.J. Nicholson. The self-adjustment of populations to change. *Cold Spring Harbor Symposia on Quantitative Biology* **22**, 153–173, 1957. doi:10.1101/SQB.1957.022.01.017.

Y. Xia and H. Tong. Feature matching in time series modeling. *Statistical Science* **26**, 21–46, 2011. doi:10.1214/10sts345.

E.L. Ionides. Discussion of "Feature matching in time series modeling" by Y. Xia and H. Tong. *Statistical Science* **26**, 49–52, 2011. doi:10.1214/11sts345c.

S. N. Wood Statistical inference for noisy nonlinear ecological dynamic systems. *Nature* **466**, 1102–1104, 2010. doi:10.1038/nature09319.

W.S.C. Gurney, S.P. Blythe, and R.M. Nisbet. Nicholson's blowflies revisited. *Nature* **287**, 17–21, 1980. doi:10.1038/287017a0.

D.R. Brillinger, J. Guckenheimer, P. Guttorp, and G. Oster. Empirical modelling of population time series: The case of age and density dependent rates. In: G. Oster (ed.), *Some Questions in Mathematical Biology* vol. 13, pp. 65–90, American Mathematical Society, Providence, 1980. doi:10.1007/9781461413448_19.

20 bsflu

See Also

```
More examples provided with pomp: childhood_disease_data, compartmental_models, dacca(), ebola, gompertz(), ou2(), pomp_examples, ricker(), rw2(), verhulst()

More data sets provided with pomp: bsflu, childhood_disease_data, dacca(), ebola, parus
```

Examples

```
plot(blowflies1())
plot(blowflies2())
```

bsflu

Influenza outbreak in a boarding school

Description

An outbreak of influenza in an all-boys boarding school.

Details

Data are recorded from a 1978 flu outbreak in a closed population. The variable 'B' refers to boys confined to bed on the corresponding day and 'C' to boys in convalescence, i.e., not yet allowed back to class. In total, 763 boys were at risk of infection and, over the course of the outbreak, 512 boys spent between 3 and 7 days away from class (either in bed or convalescent). The index case was a boy who arrived at school from holiday six days before the next case.

References

Anonymous. Influenza in a boarding school. British Medical Journal 1, 587, 1978.

See Also

compartmental models

```
More data sets provided with pomp: blowflies, childhood_disease_data, dacca(), ebola, parus
```

Examples

```
if (require(tidyr) && require(ggplot2)) {
  bsflu |>
    gather(variable,value,-date,-day) |>
    ggplot(aes(x=date,y=value,color=variable))+
    geom_line()+
    labs(y="number of boys",title="boarding school flu outbreak")+
    theme_bw()
}
```

bsmc2 21

bsmc2

The Liu and West Bayesian particle filter

Description

Modified version of the Liu & West (2001) algorithm.

Usage

```
## S4 method for signature 'data.frame'
bsmc2(
  data,
  . . . ,
  Nρ,
  smooth = 0.1,
  params,
  rprior,
  rinit,
  rprocess,
  dmeasure,
  partrans,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
bsmc2(data, ..., Np, smooth = 0.1, verbose = getOption("verbose", FALSE))
```

Arguments

data

either a data frame holding the time series data, or an object of class 'pomp', i.e., the output of another **pomp** calculation. Internally, data will be coerced to an array with storage-mode double.

• • •

additional arguments are passed to pomp. This allows one to set, unset, or modify basic model components within a call to this function.

Np

the number of particles to use. This may be specified as a single positive integer, in which case the same number of particles will be used at each timestep. Alternatively, if one wishes the number of particles to vary across timesteps, one may specify Np either as a vector of positive integers of length

```
length(time(object,t0=TRUE))
```

or as a function taking a positive integer argument. In the latter case, Np(k) must be a single positive integer, representing the number of particles to be used at the k-th timestep: Np(0) is the number of particles to use going from timezero(object) to time(object)[1], Np(1), from timezero(object) to time(object)[1], and so on, while when T=length(time(object)), Np(T) is the number of particles to sample at the end of the time-series.

22 bsmc2

smooth Kernel density smoothing parameter. The compensating shrinkage factor will be sqrt(1-smooth^2). Thus, smooth=0 means that no noise will be added to parameters. The general recommendation is that the value of smooth should be chosen close to 0 (e.g., shrink ~ 0.1). optional; named numeric vector of parameters. This will be coerced internally params to storage mode double. optional; prior distribution sampler, specified either as a C snippet, an R funcrprior tion, or the name of a pre-compiled native routine available in a dynamically loaded library. For more information, see prior specification. Setting rprior=NULL removes the prior distribution sampler. rinit simulator of the initial-state distribution. This can be furnished either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator to its default. For more information, see rinit specification. rprocess simulator of the latent state process, specified using one of the rprocess plugins. Setting rprocess=NULL removes the latent-state simulator. For more information, see rprocess specification for the documentation on these plugins. dmeasure evaluator of the measurement model density, specified either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting dmeasure=NULL removes the measurement density evaluator. For more information, see dmeasure specification. optional parameter transformations, constructed using parameter_trans. partrans Many algorithms for parameter estimation search an unconstrained space of parameters. When working with such an algorithm and a model for which the parameters are constrained, it can be useful to transform parameters. One should supply the partrans argument via a call to parameter_trans. For more information, see parameter trans. Setting partrans=NULL removes the parameter transformations, i.e., sets them to the identity transformation. verbose logical; if TRUE, diagnostic messages will be printed to the console.

Details

bsmc2 uses a version of the original algorithm (Liu & West 2001), but discards the auxiliary particle filter. The modification appears to give superior performance for the same amount of effort.

Samples from the prior distribution are drawn using the rprior component. This is allowed to depend on elements of params, i.e., some of the elements of params can be treated as "hyperparameters". Np draws are made from the prior distribution.

Value

An object of class 'bsmcd_pomp'. The following methods are avaiable:

plot produces diagnostic plots

as.data.frame puts the prior and posterior samples into a data frame

bsplines 23

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Author(s)

Michael Lavine, Matthew Ferrari, Aaron A. King, Edward L. Ionides

References

J. Liu and M. West. Combining parameter and state estimation in simulation-based filtering. In A. Doucet, N. de Freitas, and N. J. Gordon, (eds.), *Sequential Monte Carlo Methods in Practice*, pp. 197–224. Springer, New York, 2001. doi:10.1007/9781475734379_10.

See Also

```
More on Bayesian methods: abc(), dprior(), pmcmc(), prior_spec, rprior()

More on full-information (i.e., likelihood-based) methods: mif2(), pfilter(), pmcmc(), wpfilter()

More on sequential Monte Carlo methods: cond_logLik(), eff_sample_size(), filter_mean(), filter_traj(), kalman, mif2(), pfilter(), pmcmc(), pred_mean(), pred_var(), saved_states(), wpfilter()

More on pomp estimation algorithms: abc(), estimation_algorithms, mif2(), nlf, pmcmc(),
```

bsplines

B-spline bases

pomp-package, probe_match, spect_match

Description

These functions generate B-spline basis functions. bspline_basis gives a basis of spline functions. periodic_bspline_basis gives a basis of periodic spline functions.

Usage

```
bspline_basis(x, nbasis, degree = 3, deriv = 0, names = NULL, rg = range(x))
periodic_bspline_basis(
    x,
    nbasis,
    degree = 3,
    period = 1,
    deriv = 0,
    names = NULL
)
```

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Arguments

X	Vector at which the	spline functions are to	be evaluated.
---	---------------------	-------------------------	---------------

nbasis The number of basis functions to return.

degree Degree of requested B-splines.

deriv The order of the derivative required.

names optional; the names to be given to the basis functions. These will be the column-

names of the matrix returned. If the names are specified as a format string (e.g., "basis%d"), sprintf will be used to generate the names from the column number. If a single non-format string is specified, the names will be generated by paste-ing name to the column number. One can also specify each column name explicitly by giving a length-nbasis string vector. By default, no column-

names are given.

rg numeric of length 2; range of the B-spline basis. To be properly specified, we

must have rg[1] < rg[2].

period The period of the requested periodic B-splines.

Value

 $\verb|bspline_basis| Returns a matrix with length(x) rows and nbasis columns. Each column$

contains the values one of the spline basis functions.

periodic_bspline_basis

Returns a matrix with length(x) rows and nbasis columns. The basis functions returned are periodic with period period.

If deriv>0, the derivative of that order of each of the corresponding spline basis functions are returned.

C API

Access to the underlying C routines is available: see the **pomp** C API document. for definition and documentation of the C API.

Author(s)

Aaron A. King

See Also

More on interpolation: covariates, lookup()

Examples

```
x <- seq(0,2,by=0.01)
y <- bspline_basis(x,degree=3,nbasis=9,names="basis")
matplot(x,y,type='l',ylim=c(0,1.1))
lines(x,apply(y,1,sum),lwd=2)
x <- seq(-1,2,by=0.01)</pre>
```

childhood_disease_data

```
y <- periodic_bspline_basis(x,nbasis=5,names="spline%d")
matplot(x,y,type='l')</pre>
```

childhood_disease_data

Historical childhood disease incidence data

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Description

LondonYorke is a data frame containing the monthly number of reported cases of chickenpox, measles, and mumps from two American cities (Baltimore and New York) in the mid-20th century (1928–1972).

ewmeas and ewcitmeas are data frames containing weekly reported cases of measles in England and Wales. ewmeas records the total measles reports for the whole country, 1948–1966. One questionable data point has been replaced with an NA. ewcitmeas records the incidence in seven English cities 1948–1987. These data were kindly provided by Ben Bolker, who writes: "Most of these data have been manually entered from published records by various people, and are prone to errors at several levels. All data are provided as is; use at your own risk."

References

W. P. London and J. A. Yorke, Recurrent outbreaks of measles, chickenpox and mumps: I. Seasonal variation in contact rates. *American Journal of Epidemiology* **98**, 453–468, 1973. doi:10.1093/oxfordjournals.aje.a121575.

See Also

```
compartmental models, bsflu
```

```
More data sets provided with pomp: blowflies, bsflu, dacca(), ebola, parus

More examples provided with pomp: blowflies, compartmental_models, dacca(), ebola, gompertz(),
ou2(), pomp_examples, ricker(), rw2(), verhulst()
```

Examples

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```
plot(
    cases~time,
    data=LondonYorke,
    subset=disease=="mumps"&town=="New York",
    type='l',col="blue",main="mumps, New York",
    bty='l'
    )

plot(reports~time,data=ewmeas,type='l')

plot(reports~date,data=ewcitmeas,subset=city=="Liverpool",type='l')
```

coef

Extract, set, or alter coefficients

Description

Extract, set, or modify the estimated parameters from a fitted model.

Usage

```
## S4 method for signature 'listie'
coef(object, ...)

## S4 method for signature 'pomp'
coef(object, pars, transform = FALSE, ...)

## S4 replacement method for signature 'pomp'
coef(object, pars, transform = FALSE, ...) <- value

## S4 method for signature 'objfun'
coef(object, ...)

## S4 replacement method for signature 'objfun'
coef(object, pars, transform = FALSE, ...) <- value</pre>
```

Arguments

object an object of class 'pomp', or of a class extending 'pomp'
... ignored or passed to the more primitive function

pars optional character; names of parameters to be retrieved or set.

transform logical; perform parameter transformation?

value numeric vector or list; values to be assigned. If value = NULL, the parameters are unset.

Details

```
coef allows one to extract the parameters from a fitted model.
coef(object, transform=TRUE) returns the parameters transformed onto the estimation scale.
coef(object) <- value sets or alters the coefficients of a 'pomp' object.
coef(object, transform=TRUE) <- value assumes that value is on the estimation scale, and applies the "from estimation scale" parameter transformation from object before altering the coefficients.</pre>
```

See Also

```
Other extraction methods: cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_mean(), pred_var(), saved_states(), spy(), states(), summary(), time(), timezero(), traces()
```

Description

Simple SIR-type models implemented in various ways.

Usage

```
sir(
  gamma = 26,
 mu = 0.02,
 iota = 0.01,
 beta1 = 400,
 beta2 = 480,
 beta3 = 320,
 beta_sd = 0.001,
  rho = 0.6,
  k = 0.1,
 pop = 2100000,
  S_0 = 26/400,
  I_0 = 0.001,
 R_0 = 1 - S_0 - I_0
  t0 = 0,
  times = seq(from = t0 + 1/52, to = t0 + 4, by = 1/52),
  seed = 329343545,
  delta.t = 1/52/20
)
sir2(
 gamma = 24,
 mu = 1/70,
```

```
iota = 0.1,
beta1 = 330,
beta2 = 410,
beta3 = 490,
rho = 0.1,
k = 0.1,
pop = 1e+06,
S_0 = 0.05,
I_0 = 1e-04,
R_0 = 1 - S_0 - I_0,
t0 = 0,
times = seq(from = t0 + 1/12, to = t0 + 10, by = 1/12),
seed = 1772464524
)
```

Arguments

gamma recovery rate

mu death rate (assumed equal to the birth rate)

iota infection import rate

beta1, beta2, beta3

seasonal contact rates

beta_sd environmental noise intensity

rho reporting efficiency

k reporting overdispersion parameter (reciprocal of the negative-binomial size pa-

rameter)

pop overall host population size

S_0, I_0, R_0 the fractions of the host population that are susceptible, infectious, and recov-

ered, respectively, at time zero.

t0 zero time

times observation times

seed seed of the random number generator

delta.t Euler step size

Details

sir() producees a 'pomp' object encoding a simple seasonal SIR model with simulated data. Simulation is performed using an Euler multinomial approximation.

sir2() has the same model implemented using Gillespie's algorithm.

In both cases the measurement model is negative binomial: reports is distributed as a negative binomial random variable with mean equal to rho*cases and size equal to 1/k.

This and similar examples are discussed and constructed in tutorials available on the package website.

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Value

These functions return 'pomp' objects containing simulated data.

See Also

```
More examples provided with pomp: blowflies, childhood_disease_data, dacca(), ebola, gompertz(), ou2(), pomp_examples, ricker(), rw2(), verhulst()
```

Examples

```
po <- sir()
plot(po)
coef(po)

po <- sir2()
plot(po)
plot(simulate(window(po,end=3)))
coef(po)

po |> as.data.frame() |> head()
```

conc

Concatenate

Description

Internal methods to concatenate objects into useful listie.

Usage

```
## S4 method for signature 'Pomp'
conc(...)
## S4 method for signature 'Pfilter'
conc(...)
## S4 method for signature 'Abc'
conc(...)
## S4 method for signature 'Mif2'
conc(...)
## S4 method for signature 'Pmcmc'
conc(...)
```

Details

Not exported.

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concat

Concatenate

Description

Concatenate two or more 'pomp' objects into a list-like 'listie'.

Usage

```
## S3 method for class 'Pomp'
c(...)
concat(...)
```

Arguments

... elements to be recursively combined into a 'listie'

Details

concat applied to one or more 'pomp' objects or lists of 'pomp' objects converts the list into a 'listie'. In particular, concat(A,B,C) is equivalent to do.call(c,unlist(list(A,B,C))).

Examples

```
gompertz(sigma=2,tau=1) -> g
Np <- c(low=100,med=1000,high=10000)
lapply(
    Np,
    \((np)) pfilter(g,Np=np)
) |>
    concat() -> pfs

pfs
coef(pfs)
logLik(pfs)
eff_sample_size(pfs)
cond_logLik(pfs)

pfs |> plot()
```

cond_logLik 31

cond_logLik

Conditional log likelihood

Description

The estimated conditional log likelihood from a fitted model.

Usage

```
## S4 method for signature 'kalmand_pomp'
cond_logLik(object, ..., format = c("numeric", "data.frame"))
## S4 method for signature 'pfilterd_pomp'
cond_logLik(object, ..., format = c("numeric", "data.frame"))
## S4 method for signature 'wpfilterd_pomp'
cond_logLik(object, ..., format = c("numeric", "data.frame"))
## S4 method for signature 'bsmcd_pomp'
cond_logLik(object, ..., format = c("numeric", "data.frame"))
## S4 method for signature 'pfilterList'
cond_logLik(object, ..., format = c("numeric", "data.frame"))
```

Arguments

object result of a filtering computation

... ignored

format of the returned object

Details

The conditional likelihood is defined to be the value of the density of

$$Y(t_k)|Y(t_1),\ldots,Y(t_{k-1})$$

evaluated at $Y(t_k) = y_k^*$. Here, $Y(t_k)$ is the observable process, and y_k^* the data, at time t_k .

Thus the conditional log likelihood at time t_k is

$$\ell_k(\theta) = \log f[Y(t_k) = y_k^* | Y(t_1) = y_1^*, \dots, Y(t_{k-1}) = y_{k-1}^*],$$

where f is the probability density above.

Value

The numerical value of the conditional log likelihood. Note that some methods compute not the log likelihood itself but instead a related quantity. To keep the code simple, the cond_logLik function is nevertheless used to extract this quantity.

When object is of class 'bsmcd_pomp' (i.e., the result of a bsmc2 computation), cond_logLik returns the conditional log "evidence" (see bsmc2).

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See Also

```
More on sequential Monte Carlo methods: bsmc2(), eff_sample_size(), filter_mean(), filter_traj(), kalman, mif2(), pfilter(), pmcmc(), pred_mean(), pred_var(), saved_states(), wpfilter()

Other extraction methods: coef(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_mean(), pred_var(), saved_states(), spy(), states(), summary(), time(), timezero(), traces()
```

continue

Continue an iterative calculation

Description

Continue an iterative computation where it left off.

Usage

```
## S4 method for signature 'abcd_pomp'
continue(object, ..., Nabc = 1)

## S4 method for signature 'pmcmcd_pomp'
continue(object, ..., Nmcmc = 1)

## S4 method for signature 'mif2d_pomp'
continue(object, ..., Nmif = 1)
```

Arguments

object	the result of an iterative pomp computation
• • •	additional arguments will be passed to the underlying method. This allows one to modify parameters used in the original computations.
Nabc	positive integer; number of additional ABC iterations to perform
Nmcmc	positive integer; number of additional PMCMC iterations to perform
Nmif	positive integer; number of additional filtering iterations to perform

See Also

```
mif2 pmcmc abc
```

covariates 33

Description

Incorporating time-varying covariates using lookup tables.

Usage

```
## S4 method for signature 'numeric'
covariate_table(..., order = c("linear", "constant"), times)
## S4 method for signature 'character'
covariate_table(..., order = c("linear", "constant"), times)
repair_lookup_table(table, t, order)
```

Arguments

• • •	numeric vectors or data frames containing time-varying covariates. It must be possible to bind these into a data frame.
order	the order of interpolation to be used. Options are "linear" (the default) and "constant". Setting order="linear" treats the covariates as piecewise linear functions of time; order="constant" treats them as right-continuous piecewise constant functions.
times	the times corresponding to the covariates. This may be given as a vector of (non-decreasing, finite) numerical values. Alternatively, one can specify by name which of the given variables is the time variable.
table	a 'covartable' object created by a call to covariate_table
t	numeric vector; times at which interpolated values of the covariates in table are required.

Details

If the 'pomp' object contains covariates (specified via the covar argument), then interpolated values of the covariates will be available to each of the model components whenever it is called. In particular, variables with names as they appear in the covar covariate table will be available to any C snippet. When a basic component is defined using an R function, that function will be called with an extra argument, covars, which will be a named numeric vector containing the interpolated values from the covariate table.

An exception to this rule is the prior (rprior and dprior): covariate-dependent priors are not allowed. Nor are parameter transformations permitted to depend upon covariates.

repair_lookup_table applies lookup at the provided values of t and returns the resulting lookup table. If order is unsupplied, the interpolation-order from table is preserved. repair_lookup_table should be considered experimental: its interface may change without notice.

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Value

covariate_table returns a lookup table suitable for inclusion of covariates in a 'pomp' object. Specifically, this is an object of class 'covartable'.

repair_lookup_table returns a lookup table with entries at the provided values of t.

Extrapolation

If t is outside the range of the lookup table, the values will be extrapolated, and a warning will be issued. The type of extrapolation performed will be constant or linear according to the order flag used when creating the table.

See Also

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

More on interpolation: bsplines, lookup()

covmat

Estimate a covariance matrix from algorithm traces

Description

A helper function to extract a covariance matrix.

Usage

```
## S4 method for signature 'pmcmcd_pomp'
covmat(object, start = 1, thin = 1, expand = 2.38, ...)
## S4 method for signature 'pmcmcList'
covmat(object, start = 1, thin = 1, expand = 2.38, ...)
## S4 method for signature 'abcd_pomp'
covmat(object, start = 1, thin = 1, expand = 2.38, ...)
## S4 method for signature 'abcList'
covmat(object, start = 1, thin = 1, expand = 2.38, ...)
## S4 method for signature 'probed_pomp'
covmat(object, ...)
```

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Arguments

object	an object extending 'pomp'
start	the first iteration number to be used in estimating the covariance matrix. Setting thin > 1 allows for a burn-in period.
thin	factor by which the chains are to be thinned
expand	the expansion factor
	ignored

Value

When object is the result of a pmcmc or abc computation, covmat(object) gives the covariance matrix of the chains. This can be useful, for example, in tuning the proposal distribution.

When object is a 'probed_pomp' object (i.e., the result of a probe computation), covmat(object) returns the covariance matrix of the probes, as applied to simulated data.

See Also

```
MCMC proposals.
```

```
Other extraction methods: coef(), cond_logLik(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_mean(), pred_var(), saved_states(), spy(), states(), summary(), time(), timezero(), traces()
```

Csnippet	C snippets	

Description

Accelerating computations through inline snippets of C code

Usage

```
Csnippet(text)
```

Arguments

text character; text written in the C language

Details

pomp provides a facility whereby users can define their model's components using inline C code. C snippets are written to a C file, by default located in the R session's temporary directory, which is then compiled (via R CMD SHLIB) into a dynamically loadable shared object file. This is then loaded as needed.

Note to Windows and Mac users

By default, your R installation may not support R CMD SHLIB. The package website contains installation instructions that explain how to enable this powerful feature of R.

General rules for writing C snippets

In writing a C snippet one must bear in mind both the *goal* of the snippet, i.e., what computation it is intended to perform, and the *context* in which it will be executed. These are explained here in the form of general rules. Additional specific rules apply according to the function of the particular C snippet. Illustrative examples are given in the tutorials on the package website.

- C snippets must be valid C. They will embedded verbatim in a template file which will then be compiled by a call to R CMD SHLIB. If the resulting file does not compile, an error message will be generated. Compiler messages will be displayed, but no attempt will be made by **pomp** to interpret them. Typically, compilation errors are due to either invalid C syntax or undeclared variables.
- 2. State variables, parameters, observables, and covariates must be left undeclared within the snippet. State variables and parameters are declared via the statenames or paramnames arguments to pomp, respectively. Compiler errors that complain about undeclared state variables or parameters are usually due to failure to declare these in statenames or paramnames, as appropriate.
- 3. A C snippet can declare local variables. Be careful not to use names that match those of state variables, observables, or parameters. One must never declare state variables, observables, covariates, or parameters within a C snippet.
- 4. Names of observables must match the names given given in the data. They must be referred to in measurement model C snippets (rmeasure and dmeasure) by those names.
- 5. If the 'pomp' object contains a table of covariates (see above), then the variables in the covariate table will be available, by their names, in the context within which the C snippet is executed.
- 6. Because the dot '.' has syntactic meaning in C, R variables with names containing dots ('.') are replaced in the C codes by variable names in which all dots have been replaced by underscores ('_').
- 7. The headers 'R.h' and 'Rmath.h', provided with R, will be included in the generated C file, making all of the R C API available for use in the C snippet. This makes a great many useful functions available, including all of R's statistical distribution functions.
- 8. The header 'pomp.h', provided with **pomp**, will also be included, making all of the **pomp** C API available for use in every C snippet.
- 9. Snippets of C code passed to the globals argument of pomp will be included at the head of the generated C file. This can be used to declare global variables, define useful functions, and include arbitrary header files.

Linking to precompiled libraries

It is straightforward to link C snippets with precompiled C libraries. To do so, one must make sure the library's header files are included; the globals argument can be used for this purpose. The shlib.args argument can then be used to specify additional arguments to be passed to R CMD SHLIB. FAQ 3.7 gives an example.

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C snippets are salted

To prevent collisions in parallel computations, a 'pomp' object built using C snippets is "salted" with the current time and a random number. A result is that two 'pomp' objects, built on identical codes and data, will **not** be identical as R objects, though they will be functionally identical in every respect.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

See Also

spy

More on implementing POMP models: accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

dacca

Model of cholera transmission for historic Bengal.

Description

dacca constructs a 'pomp' object containing census and cholera mortality data from the Dacca district of the former British province of Bengal over the years 1891 to 1940 together with a stochastic differential equation transmission model. The model is that of King et al. (2008). The parameters are the MLE for the SIRS model with seasonal reservoir.

```
dacca(
  gamma = 20.8,
  eps = 19.1,
  rho = 0,
  delta = 0.02,
  deltaI = 0.06,
  clin = 1,
  alpha = 1,
  beta_trend = -0.00498,
  logbeta = c(0.747, 6.38, -3.44, 4.23, 3.33, 4.55),
  logomega = log(c(0.184, 0.0786, 0.0584, 0.00917, 0.000208, 0.0124)),
  sd_beta = 3.13,
```

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```
tau = 0.23,

S_0 = 0.621,

I_0 = 0.378,

Y_0 = 0,

R1_0 = 0.000843,

R2_0 = 0.000972,

R3_0 = 1.16e-07
```

Arguments

gamma	recovery rate
eps	rate of waning of immunity for severe infections
rho	rate of waning of immunity for inapparent infections
delta	baseline mortality rate
deltaI	cholera mortality rate
clin	fraction of infections that lead to severe infection
alpha	transmission function exponent
beta_trend	slope of secular trend in transmission
logbeta	seasonal transmission rates
logomega	seasonal environmental reservoir parameters
sd_beta	environmental noise intensity
tau	measurement error s.d.
S_0	initial susceptible fraction
I_0	initial fraction of population infected
Y_0	initial fraction of the population in the Y class
R1_0, R2_0, R3_0	initial fractions in the respective R classes

Details

Data are provided courtesy of Dr. Menno J. Bouma, London School of Tropical Medicine and Hygiene.

Value

dacca returns a 'pomp' object containing the model, data, and MLE parameters, as estimated by King et al. (2008).

References

A.A. King, E.L. Ionides, M. Pascual, and M.J. Bouma. Inapparent infections and cholera dynamics. *Nature* **454**, 877-880, 2008. doi:10.1038/nature07084.

design 39

See Also

More examples provided with **pomp**: blowflies, childhood_disease_data, compartmental_models, ebola, gompertz(), ou2(), pomp_examples, ricker(), rw2(), verhulst()

More data sets provided with pomp: blowflies, bsflu, childhood_disease_data, ebola, parus

Examples

```
# takes too long for R CMD check
po <- dacca()
plot(po)
## MLE:
coef(po)
plot(simulate(po))</pre>
```

design

Design matrices for pomp calculations

Description

These functions are useful for generating designs for the exploration of parameter space.

profile_design generates a data-frame where each row can be used as the starting point for a profile likelihood calculation.

runif_design generates a design based on random samples from a multivariate uniform distribu-

slice_design generates points along slices through a specified point.

sobol_design generates a Latin hypercube design based on the Sobol' low-discrepancy sequence.

```
profile_design(
    ...,
    lower,
    upper,
    nprof,
    type = c("runif", "sobol"),
    stringsAsFactors = getOption("stringsAsFactors", FALSE)
)

runif_design(lower = numeric(0), upper = numeric(0), nseq)

slice_design(center, ...)

sobol_design(lower = numeric(0), upper = numeric(0), nseq)
```

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Arguments

... In profile_design, additional arguments specify the parameters over which to

profile and the values of these parameters. In slice_design, additional numeric

vector arguments specify the locations of points along the slices.

lower, upper named numeric vectors giving the lower and upper bounds of the ranges, respec-

tively.

nprof The number of points per profile point.

type the type of design to use. type="runif" uses runif_design. type="sobol"

uses sobol_design;

stringsAsFactors

should character vectors be converted to factors?

nseq Total number of points requested.

center center is a named numeric vector specifying the point through which the slice(s)

is (are) to be taken.

Details

The Sobol' sequence generation is performed using codes from the **NLopt** library by S. Johnson.

Value

profile_design returns a data frame with nprof points per profile point.

runif_design returns a data frame with nseq rows and one column for each variable named in lower and upper.

slice_design returns a data frame with one row per point. The 'slice' variable indicates which slice the point belongs to.

sobol_design returns a data frame with nseq rows and one column for each variable named in lower and upper.

Author(s)

Aaron A. King

References

- S. Kucherenko and Y. Sytsko. Application of deterministic low-discrepancy sequences in global optimization. *Computational Optimization and Applications* **30**, 297–318, 2005. doi:10.1007/s10589-00546151.
- S.G. Johnson. The **NLopt** nonlinear-optimization package. https://github.com/stevengj/nlopt/.
- P. Bratley and B.L. Fox. Algorithm 659 Implementing Sobol's quasirandom sequence generator. *ACM Transactions on Mathematical Software* **14**, 88–100, 1988. doi:10.1145/42288.214372.
- S. Joe and F.Y. Kuo. Remark on algorithm 659: Implementing Sobol' quasirandom sequence generator. *ACM Transactions on Mathematical Software* **29**, 49–57, 2003. doi:10.1145/641876.641879.

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Examples

```
## Sobol' low-discrepancy design
plot(sobol_design(lower=c(a=0,b=100),upper=c(b=200,a=1),nseq=100))
## Uniform random design
plot(runif_design(lower=c(a=0,b=100),upper=c(b=200,a=1),100))
## A one-parameter profile design:
x \leftarrow profile_design(p=1:10,lower=c(a=0,b=0),upper=c(a=1,b=5),nprof=20)
dim(x)
plot(x)
## A two-parameter profile design:
x \leftarrow profile_design(p=1:10, q=3:5, lower=c(a=0, b=0), upper=c(b=5, a=1), nprof=200)
dim(x)
plot(x)
## A two-parameter profile design with random points:
x \leftarrow \texttt{profile\_design(p=1:10,q=3:5,lower=c(a=0,b=0),upper=c(b=5,a=1),nprof=200,type="runif")}
dim(x)
plot(x)
## A single 11-point slice through the point c(A=3,B=8,C=0) along the B direction.
x \leftarrow slice_design(center=c(A=3,B=8,C=0),B=seq(0,10,by=1))
dim(x)
plot(x)
## Two slices through the same point along the A and C directions.
x <- slice_design(c(A=3,B=8,C=0),A=seq(0,5,by=1),C=seq(0,5,length=11))
dim(x)
plot(x)
```

dinit

dinit workhorse

Description

Evaluates the initial-state density.

```
## S4 method for signature 'pomp'
dinit(
  object,
    ...,
  params = coef(object),
  t0 = timezero(object),
  x,
```

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```
log = FALSE
)
```

Arguments

object an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the **pomp** inference algorithms. additional arguments are ignored. . . . a npar x nrep matrix of parameters. Each column is treated as an independent params parameter set, in correspondence with the corresponding column of x. t0 the initial time, i.e., the time corresponding to the initial-state distribution. an array containing states of the unobserved process. The dimensions of x are nvars x nrep x ntimes, where nvars is the number of state variables, nrep is the number of replicates, and ntimes is the length of times. One can also pass x as a named numeric vector, which is equivalent to the nrep=1, ntimes=1 case. if TRUE, log probabilities are returned. log

Value

dinit returns a 1-D numerical array containing the likelihoods (or log likelihoods if log=TRUE). By default, t0 is the initial time defined when the 'pomp' object ws constructed.

See Also

Specification of the initial-state distribution: dinit_spec

More on **pomp** workhorse functions: dmeasure(), dprior(), dprocess(), emeasure(), flow(), partrans(), pomp-package, rinit(), rmeasure(), rprior(), rprocess(), skeleton(), vmeasure(), workhorses

Description

Specification of the initial-state distribution density evaluator, dinit.

Details

To fully specify the unobserved Markov state process, one must give its distribution at the zero-time (t0). One specifies how to evaluate the log probability density function for this distribution using the dinit argument. As usual, this can be provided either as a C snippet or as an R function. In the former case, bear in mind that:

1. The goal of a this snippet is computation of a log likelihood, to be put into a variable named loglik.

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2. In addition to the state variables, parameters, and covariates (if any), the variable t, containing the zero-time, will be defined in the context in which the snippet is executed.

General rules for writing C snippets can be found here.

If an R function is to be used, pass

```
dinit = f
```

to pomp, where f is a function with arguments that can include the time t, any or all of the model state variables, parameters, and covariates. As usual, f may take additional arguments, provided these are passed along with it in the call to pomp. f must return a single numeric value, the log likelihood.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

See Also

dinit

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

dmeasure

dmeasure workhorse

Description

dmeasure evaluates the probability density of observations given states.

```
## S4 method for signature 'pomp'
dmeasure(
  object,
    ...,
  y = obs(object),
  x = states(object),
  times = time(object),
  params = coef(object),
  log = FALSE
)
```

dmeasure_spec

Arguments

object	an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the pomp inference algorithms.
• • •	additional arguments are ignored.
У	a matrix containing observations. The dimensions of y are nobs x ntimes, where nobs is the number of observables and ntimes is the length of times.
х	an array containing states of the unobserved process. The dimensions of x are nvars x nrep x ntimes, where nvars is the number of state variables, nrep is the number of replicates, and ntimes is the length of times. One can also pass x as a named numeric vector, which is equivalent to the nrep=1, ntimes=1 case.
times	a numeric vector (length ntimes) containing times. These must be in non-decreasing order.
params	a npar x nrep matrix of parameters. Each column is treated as an independent parameter set, in correspondence with the corresponding column of x.
log	if TRUE, log probabilities are returned.

Value

dmeasure returns a matrix of dimensions nreps x ntimes. If d is the returned matrix, d[j,k] is the likelihood (or log likelihood if log = TRUE) of the observation y[,k] at time times[k] given the state x[,j,k].

See Also

Specification of the measurement density evaluator: dmeasure_spec

More on **pomp** workhorse functions: dinit(), dprior(), dprocess(), emeasure(), flow(), partrans(), pomp-package, rinit(), rmeasure(), rprior(), rprocess(), skeleton(), vmeasure(), workhorses

dmeasure_spec	dmeasure specification

Description

Specification of the measurement model density function, dmeasure.

Details

The measurement model is the link between the data and the unobserved state process. It can be specified either by using one or both of the rmeasure and dmeasure arguments.

Suppose you have a procedure to compute the probability density of an observation given the value of the latent state variables. Then you can furnish

```
dmeasure = f
```

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to **pomp** algorithms, where f is a C snippet or R function that implements your procedure.

Using a C snippet is much preferred, due to its much greater computational efficiency. See Csnippet for general rules on writing C snippets. The goal of a *dmeasure* C snippet is to fill the variable lik with the either the probability density or the log probability density, depending on the value of the variable give_log.

In writing a dmeasure C snippet, observe that:

- 1. In addition to the states, parameters, covariates (if any), and observables, the variable t, containing the time of the observation will be defined in the context in which the snippet is executed.
- 2. Moreover, the Boolean variable give_log will be defined.
- 3. The goal of a dmeasure C snippet is to set the value of the lik variable to the likelihood of the data given the state, if give_log == 0. If give_log == 1, lik should be set to the log likelihood.

If dmeasure is to be provided instead as an R function, this is accomplished by supplying

```
dmeasure = f
```

to pomp, where f is a function. The arguments of f should be chosen from among the observables, state variables, parameters, covariates, and time. It must also have the arguments \dots , and \log . It can take additional arguments via the userdata facility. f must return a single numeric value, the probability density (or \log probability density if \log = TRUE) of y given x at time t.

Important note

It is a common error to fail to account for both log = TRUE and log = FALSE when writing the dmeasure C snippet or function.

Default behavior

If dmeasure is left unspecified, calls to dmeasure will return missing values (NA).

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

See Also

dmeasure

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

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Examples

```
## We start with the pre-built Ricker example:
ricker() -> po
## To change the measurement model density, dmeasure,
## we use the 'dmeasure' argument in any 'pomp'
## elementary or estimation function.
## Here, we pass the dmeasure specification to 'pfilter'
## as an R function.
po |>
 pfilter(
    dmeasure=function (y, N, phi, ..., log) {
     dpois(y,lambda=phi*N,log=log)
    },
   Np=100
 ) -> pf
## We can also pass it as a C snippet:
po |>
 pfilter(
    dmeasure=Csnippet("lik = dpois(y,phi*N,give_log);"),
    paramnames="phi",
    statenames="N",
   Np=100
 ) -> pf
```

dprior

dprior workhorse

Description

Evaluates the prior probability density.

Usage

```
## S4 method for signature 'pomp'
dprior(object, ..., params = coef(object), log = FALSE)
```

Arguments

object	an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the pomp inference algorithms.
	additional arguments are ignored.
params	a npar x nrep matrix of parameters. Each column is treated as an independent parameter set, in correspondence with the corresponding column of x.
log	if TRUE, log probabilities are returned.

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Value

The required density (or log density), as a numeric vector.

See Also

```
Specification of the prior density evaluator: prior_spec
```

```
More on pomp workhorse functions: dinit(), dmeasure(), dprocess(), emeasure(), flow(), partrans(), pomp-package, rinit(), rmeasure(), rprior(), rprocess(), skeleton(), vmeasure(), workhorses
```

More on Bayesian methods: abc(), bsmc2(), pmcmc(), prior_spec, rprior()

dprocess

dprocess workhorse

Description

Evaluates the probability density of a sequence of consecutive state transitions.

Usage

```
## S4 method for signature 'pomp'
dprocess(
   object,
    ...,
   x = states(object),
   times = time(object),
   params = coef(object),
   log = FALSE
)
```

Arguments

object	an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the pomp inference algorithms.
	additional arguments are ignored.
x	an array containing states of the unobserved process. The dimensions of x are nvars x nrep x ntimes, where nvars is the number of state variables, nrep is the number of replicates, and ntimes is the length of times. One can also pass x as a named numeric vector, which is equivalent to the nrep=1, ntimes=1 case.
times	a numeric vector (length ntimes) containing times. These must be in non-decreasing order.
params	a npar x nrep matrix of parameters. Each column is treated as an independent parameter set, in correspondence with the corresponding column of x.
log	if TRUE, log probabilities are returned.

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Value

dprocess returns a matrix of dimensions nrep x ntimes-1. If d is the returned matrix, d[j,k] is the likelihood (or the log likelihood if log=TRUE) of the transition from state x[,j,k-1] at time times[k-1] to state x[,j,k] at time times[k].

See Also

Specification of the process-model density evaluator: dprocess_spec

More on **pomp** workhorse functions: dinit(), dmeasure(), dprior(), emeasure(), flow(), partrans(), pomp-package, rinit(), rmeasure(), rprior(), rprocess(), skeleton(), vmeasure(), workhorses

dprocess_spec

dprocess specification

Description

Specification of the latent state process density function, dprocess.

Details

Suppose you have a procedure that allows you to compute the probability density of an arbitrary transition from state x_1 at time t_1 to state x_2 at time $t_2 > t_1$ under the assumption that the state remains unchanged between t_1 and t_2 . Then you can furnish

```
dprocess = f
```

to pomp, where f is a C snippet or R function that implements your procedure. Specifically, f should compute the *log* probability density.

Using a C snippet is much preferred, due to its much greater computational efficiency. See Csnippet for general rules on writing C snippets. The goal of a *dprocess* C snippet is to fill the variable loglik with the log probability density. In the context of such a C snippet, the parameters, and covariates will be defined, as will the times t_1 and t_2. The state variables at time t_1 will have their usual name (see statenames) with a "_1" appended. Likewise, the state variables at time t_2 will have a "_2" appended.

If f is given as an R function, it should take as arguments any or all of the state variables, parameter, covariates, and time. The state-variable and time arguments will have suffices "_1" and "_2" appended. Thus for example, if var is a state variable, when f is called, var_1 will value of state variable var at time t_1, var_2 will have the value of var at time t_2. f should return the *log* likelihood of a transition from x1 at time t1 to x2 at time t2, assuming that no intervening transitions have occurred.

To see examples, consult the demos and the tutorials on the package website.

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Note

It is not typically necessary (or even feasible) to define dprocess. In fact, no current **pomp** inference algorithm makes use of dprocess. This functionality is provided only to support future algorithm development.

Default behavior

By default, dprocess returns missing values (NA).

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

See Also

dprocess

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

ebola

Ebola outbreak, West Africa, 2014-2016

Description

Data and models for the 2014–2016 outbreak of Ebola virus disease in West Africa.

```
ebolaModel(
    country = c("GIN", "LBR", "SLE"),
    data = NULL,
    timestep = 1/8,
    nstageE = 3L,
    R0 = 1.4,
    rho = 0.2,
    cfr = 0.7,
    k = 0,
    index_case = 10,
    incubation_period = 11.4,
    infectious_period = 7
)
```

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Arguments

country ISO symbol for the country (GIN=Guinea, LBR=Liberia, SLE=Sierra Leone).

data if NULL, the situation report data (WHO Ebola Response Team 2014) for the

appropriate country or region will be used. Providing a dataset here will override

this behavior.

timestep duration (in days) of Euler timestep for the simulations.

nstageE integer; number of incubation stages.

R0 basic reproduction ratio rho case reporting efficiency

cfr case fatality rate

k dispersion parameter (negative binomial size parameter)

index_case number of cases on day 0 (2014-04-01)

incubation_period, infectious_period

mean duration (in days) of the incubation and infectious periods.

Details

The data include monthly case counts and death reports derived from WHO situation reports, as reported by the U.S. CDC. The models are described in King et al. (2015).

The data-cleaning script is included in the R source code file 'ebola.R'.

Model structure

The default incubation period is supposed to be Gamma distributed with shape parameter nstageE and mean 11.4 days and the case-fatality ratio ('cfr') is taken to be 0.7 (cf. WHO Ebola Response Team 2014). The discrete-time formula is used to calculate the corresponding alpha (cf. He et al. 2010).

The observation model is a hierarchical model for cases and deaths:

$$p(R_t, D_t|C_t) = p(R_t|C_t)p(D_t|C_t, R_t).$$

Here, $p(R_t|C_t)$ is negative binomial with mean ρC_t and dispersion parameter 1/k; $p(D_t|C_t,R_t)$ is binomial with size R_t and probability equal to the case fatality rate cfr.

References

A.A. King, M. Domenech de Cellès, F.M.G. Magpantay, and P. Rohani. Avoidable errors in the modelling of outbreaks of emerging pathogens, with special reference to Ebola. *Proceedings of the Royal Society of London, Series B* **282**, 20150347, 2015. doi:10.1098/rspb.2015.0347.

WHO Ebola Response Team. Ebola virus disease in West Africa—the first 9 months of the epidemic and forward projections. *New England Journal of Medicine* **371**, 1481–1495, 2014. doi:10.1056/NEJMoa1411100.

D. He, E.L. Ionides, and A.A. King. Plug-and-play inference for disease dynamics: measles in large and small populations as a case study. *Journal of the Royal Society Interface* **7**, 271–283, 2010. doi:10.1098/rsif.2009.0151.

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See Also

```
More data sets provided with pomp: blowflies, bsflu, childhood_disease_data, dacca(), parus

More examples provided with pomp: blowflies, childhood_disease_data, compartmental_models, dacca(), gompertz(), ou2(), pomp_examples, ricker(), rw2(), verhulst()
```

Examples

```
# takes too long for R CMD check
if (require(ggplot2) && require(tidyr)) {
    ebolaWA2014 |>
        pivot_longer(c(cases,deaths)) |>
        ggplot(aes(x=date,y=value,group=name,color=name))+
        geom_line()+
        facet_grid(country~.,scales="free_y")+
        theme_bw()+
        theme(axis.text=element_text(angle=-90))
}

plot(ebolaModel(country="SLE"))
plot(ebolaModel(country="GIN"))
plot(ebolaModel(country="LBR"))
```

 eff_sample_size

Effective sample size

Description

Estimate the effective sample size of a Monte Carlo computation.

```
## S4 method for signature 'bsmcd_pomp'
eff_sample_size(object, ..., format = c("numeric", "data.frame"))
## S4 method for signature 'pfilterd_pomp'
eff_sample_size(object, ..., format = c("numeric", "data.frame"))
## S4 method for signature 'wpfilterd_pomp'
eff_sample_size(object, ..., format = c("numeric", "data.frame"))
## S4 method for signature 'pfilterList'
eff_sample_size(object, ..., format = c("numeric", "data.frame"))
```

Arguments

object result of a filtering computation

... ignored

format of the returned object

Details

Effective sample size is computed as

$$\left(\sum_{i} w_{it}^{2}\right)^{-1},$$

where w_{it} is the normalized weight of particle i at time t.

See Also

```
More on sequential Monte Carlo methods: bsmc2(), cond_logLik(), filter_mean(), filter_traj(), kalman, mif2(), pfilter(), pmcmc(), pred_mean(), pred_var(), saved_states(), wpfilter()

Other extraction methods: coef(), cond_logLik(), covmat(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_mean(), pred_var(), saved_states(), spy(), states(), summary(), time(), timezero(), traces()
```

Description

In **pomp**, elementary algorithms perform POMP model operations. These operations do not themselves estimate parameters, though they may be instrumental in inference methods.

Details

There are six elementary algorithms in **pomp**:

- simulate which simulates from the joint distribution of latent and observed variables,
- pfilter, which performs a simple particle filter operation,
- wpfilter, which performs a weighted particle filter operation,
- probe, which computes a suite of user-specified summary statistics on actual and simulated data,
- spect, which performs a power-spectral density function computation on actual and simulated data,
- trajectory, which iterates or integrates the deterministic skeleton according to whether the latter is a (discrete-time) map or a (continuous-time) vectorfield.

Help pages detailing each elementary algorithm component are provided.

emeasure 53

See Also

basic model components, workhorse functions, estimation algorithms.

More on **pomp** elementary algorithms: kalman, pfilter(), pomp-package, probe(), simulate(), spect(), trajectory(), wpfilter()

emeasure

emeasure workhorse

Description

Return the expected value of the observed variables, given values of the latent states and the parameters.

Usage

```
## S4 method for signature 'pomp'
emeasure(
  object,
    ...,
  x = states(object),
  times = time(object),
  params = coef(object)
)
```

Arguments

object	an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the pomp inference algorithms.
	additional arguments are ignored.
х	an array containing states of the unobserved process. The dimensions of x are nvars x nrep x ntimes, where nvars is the number of state variables, nrep is the number of replicates, and ntimes is the length of times. One can also pass x as a named numeric vector, which is equivalent to the nrep=1, ntimes=1 case.
times	a numeric vector (length $ntimes$) containing times. These must be in non-decreasing order.
params	a npar x nrep matrix of parameters. Each column is treated as an independent parameter set, in correspondence with the corresponding column of x .

Value

emeasure returns a rank-3 array of dimensions nobs x nrep x ntimes, where nobs is the number of observed variables.

54 emeasure_spec

See Also

Specification of the measurement-model expectation: emeasure_spec

More on **pomp** workhorse functions: dinit(), dmeasure(), dprior(), dprocess(), flow(), partrans(), pomp-package, rinit(), rmeasure(), rprior(), rprocess(), skeleton(), vmeasure(), workhorses

emeasure_spec

emeasure specification

Description

Specification of the measurement-model conditional expectation, emeasure.

Details

The measurement model is the link between the data and the unobserved state process. Some algorithms require the conditional expectation of the measurement model, given the latent state and parameters. This is supplied using the emeasure argument.

Suppose you have a procedure to compute this conditional expectation, given the value of the latent state variables. Then you can furnish

```
emeasure = f
```

to **pomp** algorithms, where f is a C snippet or R function that implements your procedure.

Using a C snippet is much preferred, due to its much greater computational efficiency. See Csnippet for general rules on writing C snippets.

In writing an emeasure C snippet, bear in mind that:

- 1. The goal of such a snippet is to fill variables named E_y with the conditional expectations of observables y. Accordingly, there should be one assignment of E_y for each observable y.
- 2. In addition to the states, parameters, and covariates (if any), the variable t, containing the time of the observation, will be defined in the context in which the snippet is executed.

The demos and the tutorials on the package website give examples.

It is also possible, though less efficient, to specify emeasure using an R function. In this case, specify the measurement model expectation by furnishing

```
emeasure = f
```

to pomp, where f is an R function. The arguments of f should be chosen from among the state variables, parameters, covariates, and time. It must also have the argument f must return a named numeric vector of length equal to the number of observable variables. The names should match those of the observable variables.

Default behavior

The default emeasure is undefined. It will yield missing values (NA).

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Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

See Also

emeasure

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

estimation_algorithms Parameter estimation algorithms for POMP models.

Description

pomp currently implements the following algorithms for estimating model parameters:

- iterated filtering (IF2)
- particle Markov chain Monte Carlo (PMCMC)
- approximate Bayesian computation (ABC)
- probe-matching via synthetic likelihood
- · nonlinear forecasting
- power-spectrum matching
- Liu-West Bayesian sequential Monte Carlo
- Ensemble and ensemble-adjusted Kalman filters

Details

Help pages detailing each estimation algorithm are provided.

See Also

basic model components, workhorse functions, elementary algorithms.

More on **pomp** estimation algorithms: abc(), bsmc2(), mif2(), nlf, pmcmc(), pomp-package, probe_match, spect_match

56 eulermultinom

eulermultinom

Eulermultinomial and gamma-whitenoise distributions

Description

pomp provides a number of probability distributions that have proved useful in modeling partially observed Markov processes. These include the Euler-multinomial family of distributions and the the Gamma white-noise processes.

Usage

```
reulermultinom(n = 1, size, rate, dt)
deulermultinom(x, size, rate, dt, log = FALSE)
eeulermultinom(size, rate, dt)
rgammawn(n = 1, sigma, dt)
```

Arguments

n	integer; number of random variates to generate.
size	scalar integer; number of individuals at risk.
rate	numeric vector of hazard rates.
dt	numeric scalar; duration of Euler step.
X	matrix or vector containing number of individuals that have succumbed to each death process.
log	logical; if TRUE, return logarithm(s) of probabilities.
sigma	numeric scalar; intensity of the Gamma white noise process.

Details

If N individuals face constant hazards of death in K ways at rates r_1, r_2, \ldots, r_K , then in an interval of duration Δt , the number of individuals remaining alive and dying in each way is multinomially distributed:

$$(\Delta n_0, \Delta n_1, \dots, \Delta n_K) \sim \text{Multinomial}(N; p_0, p_1, \dots, p_K),$$

where $\Delta n_0 = N - \sum_{k=1}^K \Delta n_k$ is the number of individuals remaining alive and Δn_k is the number of individuals dying in way k over the interval. Here, the probability of remaining alive is

$$p_0 = \exp(-\sum_k r_k \Delta t)$$

and the probability of dying in way k is

$$p_k = \frac{r_k}{\sum_j r_j} (1 - p_0).$$

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In this case, we say that

```
(\Delta n_1, \ldots, \Delta n_K) \sim \text{Eulermultinom}(N, r, \Delta t),
```

where $r = (r_1, \dots, r_K)$. Draw m random samples from this distribution by doing

```
dn <- reulermultinom(n=m, size=N, rate=r, dt=dt),</pre>
```

where r is the vector of rates. Evaluate the probability that $x = (x_1, \dots, x_K)$ are the numbers of individuals who have died in each of the K ways over the interval $\Delta t = dt$, by doing

```
deulermultinom(x=x,size=N,rate=r,dt=dt).
```

Bretó & Ionides (2011) discuss how an infinitesimally overdispersed death process can be constructed by compounding a multinomial process with a Gamma white noise process. The Euler approximation of the resulting process can be obtained as follows. Let the increments of the equidispersed process be given by

```
reulermultinom(size=N,rate=r,dt=dt).
```

In this expression, replace the rate r with $r \Delta W/\Delta t$, where $\Delta W \sim \mathrm{Gamma}(\Delta t/\sigma^2,\sigma^2)$ is the increment of an integrated Gamma white noise process with intensity σ . That is, ΔW has mean Δt and variance $\sigma^2 \Delta t$. The resulting process is overdispersed and converges (as Δt goes to zero) to a well-defined process. The following lines of code accomplish this:

```
dW <- rgammawn(sigma=sigma,dt=dt)
dn <- reulermultinom(size=N,rate=r,dt=dW)
dn <- reulermultinom(size=N,rate=r*dW/dt,dt=dt).</pre>
```

He et al. (2010) use such overdispersed death processes in modeling measles and the "Simulation-based Inference" course discusses the value of allowing for overdispersion more generally.

For all of the functions described here, access to the underlying C routines is available: see below.

Value

or

reulermultinom returns a length(rate) by n matrix. Each column is a different random draw. Each row contains the numbers of individuals that have succumbed to the corresponding process.

deulermultinom returns a vector (of length equal to the number of columns of x). This contains the probabilities of observing each column of x given the specified parameters (size, rate, dt).

eeulermultinom returns a length(rate)-vector containing the expected number of individuals to have succumbed to the corresponding process.

rgammawn returns a vector of length n containing random increments of the integrated Gamma white noise process with intensity sigma.

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C API

An interface for C codes using these functions is provided by the package. Visit the package homepage to view the **pomp C** API document.

Author(s)

Aaron A. King

References

C. Bretó and E. L. Ionides. Compound Markov counting processes and their applications to modeling infinitesimally over-dispersed systems. *Stochastic Processes and their Applications* **121**, 2571–2591, 2011. doi:10.1016/j.spa.2011.07.005. D. He, E.L. Ionides, and A.A. King. Plug-and-play inference for disease dynamics: measles in large and small populations as a case study. *Journal of the Royal Society Interface* **7**, 271–283, 2010. doi:10.1098/rsif.2009.0151.

See Also

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

Examples

```
## Simulate 5 realizations of Euler-multinomial random variable:
dn <- reulermultinom(5,size=100,rate=c(a=1,b=2,c=3),dt=0.1)
dn

## Compute the probability mass function at each of the 5 realizations:
deulermultinom(x=dn,size=100,rate=c(1,2,3),dt=0.1)

## Compute the expectation of an Euler-multinomial:
eeulermultinom(size=100,rate=c(a=1,b=2,c=3),dt=0.1)

## An Euler-multinomial with overdispersed transitions:
dt <- 0.1
dW <- rgammawn(sigma=0.1,dt=dt)
reulermultinom(5,size=100,rate=c(a=1,b=2,c=3),dt=dW)</pre>
```

filter_mean 59

filter_mean

Filtering mean

Description

The mean of the filtering distribution

Usage

```
## S4 method for signature 'kalmand_pomp'
filter_mean(object, vars, ..., format = c("array", "data.frame"))
## S4 method for signature 'pfilterd_pomp'
filter_mean(object, vars, ..., format = c("array", "data.frame"))
```

Arguments

object result of a filtering computation

vars optional character; names of variables

... ignored

format of the returned object

Details

The filtering distribution is that of

$$X(t_k)|Y(t_1) = y_1^*, \dots, Y(t_k) = y_k^*,$$

where $X(t_k)$, $Y(t_k)$ are the latent state and observable processes, respectively, and y_t^* is the data, at time t_k .

The filtering mean is therefore the expectation of this distribution

$$E[X(t_k)|Y(t_1) = y_1^*, \dots, Y(t_k) = y_k^*].$$

See Also

```
More on sequential Monte Carlo methods: bsmc2(), cond_logLik(), eff_sample_size(), filter_traj(), kalman, mif2(), pfilter(), pmcmc(), pred_mean(), pred_var(), saved_states(), wpfilter()

Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_traj(), forecast(), logLik, obs(), pred_mean(), pred_var(), saved_states(), spy(), states(), summary(), time(), timezero(), traces()
```

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filter_traj

Filtering trajectories

Description

Drawing from the smoothing distribution

Usage

```
## S4 method for signature 'pfilterd_pomp'
filter_traj(object, vars, ..., format = c("array", "data.frame"))
## S4 method for signature 'listie'
filter_traj(object, vars, ..., format = c("array", "data.frame"))
## S4 method for signature 'pmcmcd_pomp'
filter_traj(object, vars, ...)
```

Arguments

object result of a filtering computation

vars optional character; names of variables

... ignored

format of the returned object

Details

The smoothing distribution is the distribution of

$$X(t_k)|Y(t_1) = y_1^*, \dots, Y(t_n) = y_n^*,$$

where $X(t_k)$ is the latent state process and $Y(t_k)$ is the observable process at time t_k , and n is the number of observations.

To draw samples from this distribution, one can run a number of independent particle filter (pfilter) operations, sampling the full trajectory of *one* randomly-drawn particle from each one. One should view these as *weighted* samples from the smoothing distribution, where the weights are the *likelihoods* returned by each of the pfilter computations.

One accomplishes this by setting filter.traj = TRUE in each pfilter computation and extracting the trajectory using the filter_traj command.

In particle MCMC (pmcmc), the tracking of an individual trajectory is performed automatically.

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See Also

```
More on sequential Monte Carlo methods: bsmc2(), cond_logLik(), eff_sample_size(), filter_mean(), kalman, mif2(), pfilter(), pmcmc(), pred_mean(), pred_var(), saved_states(), wpfilter()

Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), forecast(), logLik, obs(), pred_mean(), pred_var(), saved_states(), spy(), states(), summary(), time(), timezero(), traces()
```

flow

flow workhorse

Description

Compute the flow generated by a deterministic vectorfield or map.

Usage

```
## S4 method for signature 'pomp'
flow(
  object,
    ...,
  x0,
  t0 = timezero(object),
  times = time(object),
  params = coef(object),
  verbose = getOption("verbose", FALSE)
)
```

Arguments

object	an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the pomp inference algorithms.
	Additional arguments are passed to the ODE integrator (if the skeleton is a vectorfield) and are ignored if it is a map. See ode for a description of the additional arguments accepted by the ODE integrator. By default, this is the parameter vector stored in object (see coef).
x0	an array with dimensions nvar x nrep giving the initial conditions of the trajectories to be computed.
t0	the time at which the initial conditions are assumed to hold. By default, this is the zero-time (see timezero).
times	a numeric vector (length ntimes) containing times at which the itineraries are desired. These must be in non-decreasing order with times[1]>t0. By default, this is the full set of observation times (see time).
params	a npar x nrep matrix of parameters. Each column is treated as an independent parameter set, in correspondence with the corresponding column of x.
verbose	logical; if TRUE, diagnostic messages will be printed to the console.

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Details

In the case of a discrete-time system (map), flow iterates the map to yield trajectories of the system. In the case of a continuous-time system (vectorfield), flow uses the numerical solvers in **deSolve** to integrate the vectorfield starting from given initial conditions.

Value

flow returns an array of dimensions nvar x nrep x ntimes. If x is the returned matrix, x[i,j,k] is the i-th component of the state vector at time times[k] given parameters params[,j].

Accumulator variables

When there are accumulator variables (as determined by the accumvars argument), their handling in the continuous-time (vectorfield) case differs from that in the discrete-time (map) case. In the latter, accumulator variables are set to zero at the beginning of each interval (t_k, t_{k+1}) , $k = 0, 1, 2, \ldots$ over which flow computation is required. In the former, the flow computation proceeds over the entire set of intervals required, and accumulator variables are then differenced. That is, the value a_k of accumulator variable a at times t_k , $k = 1, 2, \ldots$ will be $A_k - A_{k-1}$, where A_k is the solution of the corresponding differential equation at t_k .

See Also

```
More on pomp workhorse functions: dinit(), dmeasure(), dprior(), dprocess(), emeasure(), partrans(), pomp-package, rinit(), rmeasure(), rprior(), rprocess(), skeleton(), vmeasure(), workhorses
```

More on methods for deterministic process models: skeleton(), skeleton_spec, traj_match, trajectory()

forecast

Forecast mean

Description

Mean of the one-step-ahead forecasting distribution.

```
forecast(object, ...)
## S4 method for signature 'kalmand_pomp'
forecast(object, vars, ..., format = c("array", "data.frame"))
## S4 method for signature 'pfilterd_pomp'
forecast(object, vars, ..., format = c("array", "data.frame"))
```

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Arguments

object result of a filtering computation

... ignored

vars optional character; names of variables

format of the returned object

See Also

```
Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), logLik, obs(), pred_mean(), pred_var(), saved_states(), spy(), states(), summary(), time(), timezero(), traces()
```

gompertz

Gompertz model with log-normal observations.

Description

gompertz() constructs a 'pomp' object encoding a stochastic Gompertz population model with log-normal measurement error.

Usage

```
gompertz(
  K = 1,
  r = 0.1,
  sigma = 0.1,
  tau = 0.1,
  X_0 = 1,
  times = 1:100,
  t0 = 0
)
```

Arguments

K	carrying capacity
r	growth rate
sigma	process noise intensity
tau	measurement error s.d.
X_0	value of the latent state variable X at the zero time
times	observation times
t0	zero time

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Details

The state process is

$$X_{t+1} = K^{1-S} X_t^S \epsilon_t,$$

where $S=e^{-r}$ and the ϵ_t are i.i.d. lognormal random deviates with variance σ^2 . The observed variables Y_t are distributed as

$$Y_t \sim \text{Lognormal}(\log X_t, \tau).$$

Parameters include the per-capita growth rate r, the carrying capacity K, the process noise s.d. σ , the measurement error s.d. τ , and the initial condition X_0 . The 'pomp' object includes parameter transformations that log-transform the parameters for estimation purposes.

Value

A 'pomp' object with simulated data.

See Also

More examples provided with **pomp**: blowflies, childhood_disease_data, compartmental_models, dacca(), ebola, ou2(), pomp_examples, ricker(), rw2(), verhulst()

Examples

```
plot(gompertz())
plot(gompertz(K=2,r=0.01))
```

hitch

Hitching C snippets and R functions to pomp_fun objects

Description

The algorithms in **pomp** are formulated using R functions that access the basic model components (rprocess, dprocess, rmeasure, dmeasure, etc.). For short, we refer to these elementary functions as "workhorses". In implementing a model, the user specifies basic model components using functions, procedures in dynamically-linked libraries, or C snippets. Each component is then packaged into a 'pomp_fun' objects, which gives a uniform interface. The construction of 'pomp_fun' objects is handled by the hitch function, which conceptually "hitches" the workhorses to the user-defined procedures.

```
hitch(
...,
templates,
obsnames,
statenames,
paramnames,
```

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```
covarnames,
PACKAGE,
globals,
cfile,
cdir = getOption("pomp_cdir", NULL),
on_load,
shlib.args,
compile = TRUE,
verbose = getOption("verbose", FALSE)
)
```

Arguments

. . .

named arguments representing the user procedures to be hitched. These can be functions, character strings naming routines in external, dynamically-linked libraries, C snippets, or NULL. The first three are converted by hitch to 'pomp_fun' objects which perform the indicated computations. NULL arguments are translated to default 'pomp_fun' objects. If any of these procedures are already 'pomp_fun' objects, they are returned unchanged.

templates

named list of templates. Each workhorse must have a corresponding template. See pomp:::workhorse_templates for a list.

obsnames, statenames, paramnames, covarnames

character vectors specifying the names of observable variables, latent state variables, parameters, and covariates, respectively. These are only needed if one or more of the horses are furnished as C snippets.

PACKAGE

optional character; the name (without extension) of the external, dynamically loaded library in which any native routines are to be found. This is only useful if one or more of the model components has been specified using a precompiled dynamically loaded library; it is not used for any component specified using C snippets. PACKAGE can name at most one library.

globals

optional character or C snippet; arbitrary C code that will be hard-coded into the shared-object library created when C snippets are provided. If no C snippets are used, globals has no effect.

cfile

optional character variable. cfile gives the name of the file (in directory cdir) into which C snippet codes will be written. By default, a random filename is used. If the chosen filename would result in over-writing an existing file, an error is generated.

cdir

optional character variable. cdir specifies the name of the directory within which C snippet code will be compiled. By default, this is in a temporary directory specific to the R session. One can also set this directory using the pomp_cdir global option.

on_load

optional character or C snippet: arbitrary C code that will be executed when the C snippet file is loaded. If no C snippets are used, on_load has no effect.

shlib.args

optional character variables. Command-line arguments to the R CMD SHLIB call that compiles the C snippets. One can, for example, specify libraries against which the C snippets are to be linked. In doing so, take care to make sure the

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11 1	ate header files are available to the C snippets, e.g., using the globals t. See Csnippet for more information.
logical; ineeded.	f FALSE, compilation of the C snippets will be postponed until they are
logical. played.	Setting verbose=TRUE will cause additional information to be dis-

Value

compile

verbose

hitch returns a named list of length two. The element named "funs" is itself a named list of 'pomp_fun' objects, each of which corresponds to one of the horses passed in. The element named "lib" contains information on the shared-object library created using the C snippets (if any were passed to hitch). If no C snippets were passed to hitch, lib is NULL. Otherwise, it is a length-3 named list with the following elements:

name The name of the library created.

dir The directory in which the library was created. If this is NULL, the library was created in the session's temporary directory.

src A character string with the full contents of the C snippet file.

Author(s)

Aaron A. King

See Also

```
pomp, spy
```

kalman

Ensemble Kalman filters

Description

The ensemble Kalman filter and ensemble adjustment Kalman filter.

```
## $4 method for signature 'data.frame'
enkf(
   data,
    ...,
   Np,
   params,
   rinit,
   rprocess,
   emeasure,
   vmeasure,
```

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```
verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
enkf(data, ..., Np, verbose = getOption("verbose", FALSE))
## S4 method for signature 'kalmand_pomp'
enkf(data, ..., Np, verbose = getOption("verbose", FALSE))
## S4 method for signature 'data.frame'
eakf(
  data,
  . . . ,
 Nρ,
  params,
  rinit,
  rprocess,
  emeasure,
  vmeasure,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
eakf(data, ..., Np, verbose = getOption("verbose", FALSE))
```

Arguments

data either a data frame holding the time series data, or an object of class 'pomp',

i.e., the output of another **pomp** calculation. Internally, data will be coerced to

an array with storage-mode double.

.. additional arguments are passed to pomp. This allows one to set, unset, or modify

basic model components within a call to this function.

Np integer; the number of particles to use, i.e., the size of the ensemble.

params optional; named numeric vector of parameters. This will be coerced internally

to storage mode double.

rinit simulator of the initial-state distribution. This can be furnished either as a C

snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator

to its default. For more information, see rinit specification.

rprocess simulator of the latent state process, specified using one of the rprocess plugins.

Setting rprocess=NULL removes the latent-state simulator. For more informa-

tion, see rprocess specification for the documentation on these plugins.

emeasure the expectation of the measured variables, conditional on the latent state. This

can be specified as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting emeasure=NULL removes the emeasure component. For more information, see emeasure specifi-

cation.

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vmeasure the covariance of the measured variables, conditional on the latent state. This

can be specified as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting vmeasure=NULL removes the vmeasure component. For more information, see vmeasure specifi-

cation.

verbose logical; if TRUE, diagnostic messages will be printed to the console.

Value

An object of class 'kalmand_pomp'.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Author(s)

Aaron A. King

References

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- J.L. Anderson. An ensemble adjustment Kalman filter for data assimilation. *Monthly Weather Review* **129**, 2884–2903, 2001. doi:10.1175/15200493(2001)129<2884:AEAKFF>2.0.CO;2.
- G. Evensen. *Data assimilation: the ensemble Kalman filter*. Springer-Verlag, 2009. doi:10.1007/9783642037115.

See Also

kalmanFilter

```
More on sequential Monte Carlo methods: bsmc2(), cond_logLik(), eff_sample_size(), filter_mean(), filter_traj(), mif2(), pfilter(), pmcmc(), pred_mean(), pred_var(), saved_states(), wpfilter()
```

More on **pomp** elementary algorithms: elementary_algorithms, pfilter(), pomp-package, probe(), simulate(), spect(), trajectory(), wpfilter()

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kalmanFilter	Kalman filter

Description

The basic Kalman filter for multivariate, linear, Gaussian processes.

Usage

```
kalmanFilter(object, X0 = rinit(object), A, Q, C, R, tol = 1e-06)
```

Arguments

object	a pomp object containing data;
X0	length-m vector containing initial state. This is assumed known without uncertainty.
Α	$m \times m$ latent state-process transition matrix. $E[X_{t+1} X_t] = A.X_t$.
Q	$m \times m$ latent state-process covariance matrix. $Var[X_{t+1} X_t] = Q$
С	$n \times m$ link matrix. $E[Y_t X_t] = C.X_t$.
R	$n \times n$ observation process covariance matrix. $Var[Y_t X_t] = R$
tol	numeric; the tolerance to be used in computing matrix pseudoinverses via singular-value decomposition. Singular values smaller than tol are set to zero.

Details

If the latent state is X, the observed variable is Y, $X_t \in \mathbb{R}^m$, $Y_t \in \mathbb{R}^n$, and

$$X_t \sim \text{MVN}(AX_{t-1}, Q)$$

 $Y_t \sim \text{MVN}(CX_t, R)$

where MVN(M, V) denotes the multivariate normal distribution with mean M and variance V. Then the Kalman filter computes the exact likelihood of Y given A, C, Q, and R.

Value

A named list containing the following elements:

```
object the 'pomp' object A, Q, C, R as in the call filter.mean E[X_t|y_1^*, \ldots, y_t^*] pred.mean E[X_t|y_1^*, \ldots, y_{t-1}^*] forecast E[Y_t|y_1^*, \ldots, y_{t-1}^*] cond.logLik f(y_t^*|y_1^*, \ldots, y_{t-1}^*) logLik f(y_1^*, \ldots, y_T^*)
```

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See Also

```
enkf, eakf
```

Examples

```
if (require(dplyr)) {
  gompertz() -> po
  po |>
   as.data.frame() |>
   mutate(
     logY=log(Y)
   ) |>
    select(time,logY) |>
   pomp(times="time",t0=0) |>
   kalmanFilter(
     X0=c(logX=0),
      A=matrix(exp(-0.1),1,1),
      Q=matrix(0.01,1,1),
     C=matrix(1,1,1),
      R=matrix(0.01,1,1)
   ) -> kf
  po |>
   pfilter(Np=1000) -> pf
  kf$logLik
  logLik(pf) + sum(log(obs(pf)))
}
```

listie listie

Description

List-like objects.

```
## S4 method for signature 'listie' x[i, j, ..., drop = TRUE]
```

load 71

load

Loading and unloading shared-object libraries

Description

pompLoad and pompUnload cause compiled codes associated with object to be dynamically linked or unlinked, respectively. solibs<- is a helper function for developers of packages that extend **pomp**.

Usage

```
## S4 method for signature 'pomp'
pompLoad(object, ...)
## S4 method for signature 'pomp'
pompUnload(object, ...)
## S4 replacement method for signature 'pomp'
solibs(object, ...) <- value
## S4 method for signature 'objfun'
pompLoad(object, ...)
## S4 method for signature 'objfun'
pompUnload(object, ...)</pre>
```

Arguments

object

an object of class 'pomp', or extending this class.

Details

When C snippets are used in the construction of a 'pomp' object, the resulting shared-object library is dynamically loaded (linked) before each use, and unloaded afterward.

solibs<- prepends the 'lib' generated by hitch to the 'solibs' slot of a 'pomp' object.

logLik

Log likelihood

Description

Extract the estimated log likelihood (or related quantity) from a fitted model.

72 logLik

Usage

```
logLik(object, ...)
## S4 method for signature 'listie'
logLik(object, ...)
## S4 method for signature 'pfilterd_pomp'
logLik(object)
## S4 method for signature 'wpfilterd_pomp'
logLik(object)
## S4 method for signature 'probed_pomp'
logLik(object)
## S4 method for signature 'kalmand_pomp'
logLik(object)
## S4 method for signature 'pmcmcd_pomp'
logLik(object)
## S4 method for signature 'bsmcd_pomp'
logLik(object)
## S4 method for signature 'objfun'
logLik(object)
## S4 method for signature 'spect_match_objfun'
logLik(object)
## S4 method for signature 'nlf_objfun'
logLik(object, ...)
```

Arguments

object fitted model object

... ignored

Value

numerical value of the log likelihood. Note that some methods compute not the log likelihood itself but instead a related quantity. To keep the code simple, the logLik function is nevertheless used to extract this quantity.

When object is of 'pfilterd_pomp' class (i.e., the result of a wpfilter computation), logLik retrieves the estimated log likelihood.

When object is of 'wpfilterd_pomp' class (i.e., the result of a wpfilter computation), logLik retrieves the estimated log likelihood.

logmeanexp 73

When object is of 'probed_pomp' class (i.e., the result of a probe computation), logLik retrieves the "synthetic likelihood".

When object is of 'kalmand_pomp' class (i.e., the result of an eakf or enkf computation), logLik retrieves the estimated log likelihood.

When object is of 'pmcmcd_pomp' class (i.e., the result of a pmcmc computation), logLik retrieves the estimated log likelihood as of the last particle filter operation.

When object is of 'bsmcd_pomp' class (i.e., the result of a bsmc2 computation), logLik retrieves the "log evidence".

When object is of 'spect_match_objfun' class (i.e., an objective function constructed by spect_objfun), logLik retrieves minus the spectrum mismatch.

When object is an NLF objective function, i.e., the result of a call to nlf_objfun, logLik retrieves the "quasi log likelihood".

See Also

```
Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), obs(), pred_mean(), pred_var(), saved_states(), spy(), states(), summary(), time(), timezero(), traces()
```

logmeanexp

The log-mean-exp trick

Description

logmeanexp computes

$$\log \frac{1}{n} \sum_{i=1}^{n} e^{x_i},$$

avoiding over- and under-flow in doing so. It can optionally return an estimate of the standard error in this quantity.

Usage

```
logmeanexp(x, se = FALSE, ess = FALSE)
```

Arguments

x numeric

se logical; give approximate standard error?

ess logical; give effective sample size?

Details

When se = TRUE, logmeanexp uses a jackknife estimate of the variance in log(x).

When ess = TRUE, logmeanexp returns an estimate of the effective sample size.

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Value

log(mean(exp(x))) computed so as to avoid over- or underflow. If se = TRUE, the approximate standard error is returned as well. If ess = TRUE, the effective sample size is returned also.

Author(s)

Aaron A. King

Examples

```
# takes too long for R CMD check
## an estimate of the log likelihood:
ricker() |>
   pfilter(Np=1000) |>
   logLik() |>
   replicate(n=5) -> ll
logmeanexp(ll)
## with standard error:
logmeanexp(ll,se=TRUE)
## with effective sample size
logmeanexp(ll,ess=TRUE)
```

lookup

Lookup table

Description

Interpolate values from a lookup table

Usage

```
lookup(table, t)
```

Arguments

table a 'covartable' object created by a call to covariate_table

t numeric vector; times at which interpolated values of the covariates in table

are required.

Value

A numeric vector or matrix of the interpolated values.

Extrapolation

If t is outside the range of the lookup table, the values will be extrapolated, and a warning will be issued. The type of extrapolation performed will be constant or linear according to the order flag used when creating the table.

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See Also

More on interpolation: bsplines, covariates

тсар	Monte Carlo adjusted profile

Description

Given a collection of points maximizing the likelihood over a range of fixed values of a focal parameter, this function constructs a profile likelihood confidence interval accommodating both Monte Carlo error in the profile and statistical uncertainty present in the likelihood function.

Usage

```
mcap(logLik, parameter, level = 0.95, span = 0.75, Ngrid = 1000)
```

Arguments

logLik numeric; a vector of profile log likelihood evaluations.

parameter numeric; the corresponding values of the focal parameter.

level numeric; the confidence level required.

span numeric; the loess smoothing parameter.

Ngrid integer; the number of points to evaluate the smoothed profile.

Value

mcap returns a list including the loess-smoothed profile, a quadratic approximation, and the constructed confidence interval.

Author(s)

Edward L. Ionides

References

E. L. Ionides, C. Bretó, J. Park, R. A. Smith, and A. A. King. Monte Carlo profile confidence intervals for dynamic systems. *Journal of the Royal Society, Interface* **14**, 20170126, 2017. doi:10.1098/rsif.2017.0126.

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melt Melt

Description

Convert arrays, lists, and other objects to data frames.

Usage

```
## S4 method for signature 'ANY'
melt(data, ...)
## S4 method for signature 'array'
melt(data, ...)
## S4 method for signature 'list'
melt(data, ..., level = 1)
```

Arguments

data object to convert
... ignored
level integer; level of recursion

Details

melt converts its first argument to a data frame. It is a simplified version of the melt command provided by the no-longer maintained **reshape2** package.

An array can be melted into a data frame. In this case, the data frame will have one row per entry in the array.

A list can be melted into a data frame. This operation is recursive. A variable will be appended to distinguish the separate list entries.

mif2 Iterated filtering: maximum likelihood by iterated, perturbed Bayes maps

Description

An iterated filtering algorithm for estimating the parameters of a partially-observed Markov process. Running mif2 causes the algorithm to perform a specified number of particle-filter iterations. At each iteration, the particle filter is performed on a perturbed version of the model, in which the parameters to be estimated are subjected to random perturbations at each observation. This extra variability effectively smooths the likelihood surface and combats particle depletion by introducing diversity into particle population. As the iterations progress, the magnitude of the perturbations is diminished according to a user-specified cooling schedule. The algorithm is presented and justified in Ionides et al. (2015).

mif2 77

Usage

```
## S4 method for signature 'data.frame'
mif2(
  data,
  . . . ,
 Nmif = 1,
  rw.sd,
  cooling.type = c("geometric", "hyperbolic"),
  cooling.fraction.50,
  params,
  rinit,
  rprocess,
  dmeasure,
  partrans,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
mif2(
  data,
  . . . ,
 Nmif = 1,
  rw.sd,
  cooling.type = c("geometric", "hyperbolic"),
  cooling.fraction.50,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pfilterd_pomp'
mif2(data, ..., Nmif = 1, Np, verbose = getOption("verbose", FALSE))
## S4 method for signature 'mif2d_pomp'
mif2(
  data,
  . . . ,
 Nmif,
  rw.sd,
  cooling.type,
  cooling.fraction.50,
  verbose = getOption("verbose", FALSE)
)
```

Arguments

data

either a data frame holding the time series data, or an object of class 'pomp', i.e., the output of another **pomp** calculation. Internally, data will be coerced to

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an array with storage-mode double.

additional arguments are passed to pomp. This allows one to set, unset, or modify . . . basic model components within a call to this function.

Nmif The number of filtering iterations to perform.

> specification of the magnitude of the random-walk perturbations that will be applied to some or all model parameters. Parameters that are to be estimated should have positive perturbations specified here. The specification is given using the rw_sd function, which creates a list of unevaluated expressions. The latter are evaluated in a context where vector of observation times is visible (as 'time'). The expression ivp(s) can be used in this context as shorthand for

ifelse(time==time[1],s,0).

Likewise, ivp(s,lag) is equivalent to

ifelse(time==time[lag],s,0).

See below for some examples.

The perturbations that are applied are normally distributed with the specified s.d. If parameter transformations have been supplied, then the perturbations are applied on the transformed (estimation) scale.

cooling.type, cooling.fraction.50

specifications for the cooling schedule, i.e., the manner and rate with which the intensity of the parameter perturbations is reduced with successive filtering iterations. cooling. type specifies the nature of the cooling schedule. See below (under "Specifying the perturbations") for more detail.

the number of particles to use. This may be specified as a single positive integer, in which case the same number of particles will be used at each timestep. Alternatively, if one wishes the number of particles to vary across timesteps, one may specify Np either as a vector of positive integers of length

length(time(object,t0=TRUE))

or as a function taking a positive integer argument. In the latter case, Np(k) must be a single positive integer, representing the number of particles to be used at the k-th timestep: Np(0) is the number of particles to use going from timezero(object) to time(object)[1], Np(1), from timezero(object) to time(object)[1], and so on, while when T=length(time(object)), Np(T) is the number of particles to sample at the end of the time-series.

optional; named numeric vector of parameters. This will be coerced internally to storage mode double.

simulator of the initial-state distribution. This can be furnished either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator to its default. For more information, see rinit specification.

simulator of the latent state process, specified using one of the rprocess plugins. Setting rprocess=NULL removes the latent-state simulator. For more information, see rprocess specification for the documentation on these plugins.

Np

rw.sd

params

rinit

rprocess

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dmeasure evaluator of the measurement model density, specified either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting dmeasure=NULL removes the measurement density

evaluator. For more information, see dmeasure specification.

partrans optional parameter transformations, constructed using parameter_trans.

Many algorithms for parameter estimation search an unconstrained space of parameters. When working with such an algorithm and a model for which the parameters are constrained, it can be useful to transform parameters. One should supply the partrans argument via a call to parameter_trans. For more information, see parameter_trans. Setting partrans=NULL removes the parameter

transformations, i.e., sets them to the identity transformation.

verbose logical; if TRUE, diagnostic messages will be printed to the console.

Value

Upon successful completion, mif2 returns an object of class 'mif2d_pomp'.

Number of particles

If Np is anything other than a constant, the user must take care that the number of particles requested at the end of the time series matches that requested at the beginning. In particular, if T=length(time(object)), then one should have Np[1]==Np[T+1] when Np is furnished as an integer vector and Np(0)==Np(T) when Np is furnished as a function.

Methods

The following methods are available for such an object:

continue picks up where mif2 leaves off and performs more filtering iterations.

logLik returns the so-called *mif log likelihood* which is the log likelihood of the perturbed model, not of the focal model itself. To obtain the latter, it is advisable to run several pfilter operations on the result of a mif2 computatation.

coef extracts the point estimate

eff_sample_size extracts the effective sample size of the final filtering iteration

Various other methods can be applied, including all the methods applicable to a pfilterd_pomp object and all other **pomp** estimation algorithms and diagnostic methods.

Specifying the perturbations

The rw_sd function simply returns a list containing its arguments as unevaluated expressions. These are then evaluated in a context in which the vector of observation times is defined (as 'time'). This allows for easy specification of the structure of the perturbations that are to be applied. For example,

```
rw_sd(
    a=0.05,
    b=ifelse(time==time[1], 0.2, 0),
    c=ivp(0.2),
```

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```
d=ifelse(time==time[13], 0.2, 0),
    e=ivp(0.2, lag=13),
    f=ifelse(time<23, 0.02, 0),
    g=ifelse(time>=23 & time<50, 0.02, 0),
    h=ivp(0.1,lags=3:8)
)</pre>
```

results in random perturbations of parameter a with s.d. 0.05 at every time step, while parameters b and c both get perturbations of s.d. 0.2 only before the first observation (i.e., at the zero-time). Parameters d and e, by contrast, get perturbations of s.d. 0.2 only before the thirteenth observation. Parameter f gets a random perturbation of size 0.02 before every observation falling before t=23, while g gets perturbed before all observations that fall in the interval $23 \le t < 50$. Finally, the magnitude of the perturbation of parameter h is applied before the third through eighth observations.

On the m-th IF2 iteration, prior to time-point n, the d-th parameter is given a random increment normally distributed with mean 0 and standard deviation $c_{m,n}\sigma_{d,n}$, where c is the cooling schedule and σ is specified using rw_sd, as described above. Let N be the length of the time series and α =cooling.fraction.50. Then, when cooling.type="geometric", we have

$$c_{m,n} = \alpha^{\frac{n-1+(m-1)N}{50N}}.$$

When cooling.type="hyperbolic", we have

$$c_{m,n} = \frac{s+1}{s+n+(m-1)N},$$

where s satisfies

$$\frac{s+1}{s+50N} = \alpha.$$

Thus, in either case, the perturbations at the end of 50 IF2 iterations are a fraction α smaller than they are at first.

Re-running IF2 iterations

To re-run a sequence of IF2 iterations, one can use the mif2 method on a 'mif2d_pomp' object. By default, the same parameters used for the original IF2 run are re-used (except for verbose, the default of which is shown above). If one does specify additional arguments, these will override the defaults.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Author(s)

Aaron A. King, Edward L. Ionides, Dao Nguyen

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References

E.L. Ionides, D. Nguyen, Y. Atchadé, S. Stoev, and A.A. King. Inference for dynamic and latent variable models via iterated, perturbed Bayes maps. *Proceedings of the National Academy of Sciences* **112**, 719–724, 2015. doi:10.1073/pnas.1410597112.

See Also

```
More on full-information (i.e., likelihood-based) methods: bsmc2(), pfilter(), pmcmc(), wpfilter()

More on sequential Monte Carlo methods: bsmc2(), cond_logLik(), eff_sample_size(), filter_mean(),
filter_traj(), kalman, pfilter(), pmcmc(), pred_mean(), pred_var(), saved_states(),
wpfilter()
```

More on **pomp** estimation algorithms: abc(), bsmc2(), estimation_algorithms, nlf, pmcmc(), pomp-package, probe_match, spect_match

More on maximization-based estimation methods: nlf, probe_match, spect_match, traj_match

nlf

Nonlinear forecasting

Description

Parameter estimation by maximum simulated quasi-likelihood.

Usage

```
## S4 method for signature 'data.frame'
nlf_objfun(
 data,
  est = character(0),
  lags,
  nrbf = 4,
  ti,
  tf,
  seed = NULL,
  transform.data = identity,
  period = NA,
  tensor = TRUE,
  fail.value = NA_real_,
  params,
  rinit,
  rprocess,
  rmeasure,
  verbose = getOption("verbose")
)
## S4 method for signature 'pomp'
```

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```
nlf_objfun(
 data,
 est = character(0),
 lags,
 nrbf = 4,
  ti,
  tf,
  seed = NULL,
  transform.data = identity,
 period = NA,
  tensor = TRUE,
  fail.value = NA,
 verbose = getOption("verbose")
)
## S4 method for signature 'nlf_objfun'
nlf_objfun(
 data,
  ...,
 est,
 lags,
 nrbf,
  ti,
  tf,
  seed = NULL,
 period,
  tensor,
  transform.data,
 fail.value,
 verbose = getOption("verbose", FALSE)
)
```

Arguments

data	either a data frame holding the time series data, or an object of class 'pomp', i.e., the output of another pomp calculation. Internally, data will be coerced to an array with storage-mode double.
	additional arguments are passed to pomp. This allows one to set, unset, or modify basic model components within a call to this function.
est	character vector; the names of parameters to be estimated.
lags	A vector specifying the lags to use when constructing the nonlinear autoregressive prediction model. The first lag is the prediction interval.
nrbf	integer scalar; the number of radial basis functions to be used at each lag.
ti, tf	required numeric values. NLF works by generating simulating long time series from the model. The simulated time series will be from ti to tf, with the same sampling frequency as the data. ti should be chosen large enough so that

transient dynamics have died away. tf should be chosen large enough so that sufficiently many data points are available to estimate the nonlinear forecasting model well. An error will be generated unless the data-to-parameter ratio exceeds 10 and a warning will be given if the ratio is smaller than 30.

seed

integer. When fitting, it is often best to fix the seed of the random-number generator (RNG). This is accomplished by setting seed to an integer. By default, seed = NULL, which does not alter the RNG state.

transform.data

optional function. If specified, forecasting is performed using data and model simulations transformed by this function. By default, transform.data is the identity function, i.e., no transformation is performed. The main purpose of transform.data is to achieve approximately multivariate normal forecasting errors. If the data are univariate, transform.data should take a scalar and return a scalar. If the data are multivariate, transform.data should assume a vector input and return a vector of the same length.

period

numeric; period=NA means the model is nonseasonal. period > 0 is the period of seasonal forcing. period <= 0 is equivalent to period = NA.

tensor

logical; if FALSE, the fitted model is a generalized additive model with time mod period as one of the predictors, i.e., a gam with time-varying intercept. If TRUE, the fitted model is a gam with lagged state variables as predictors and time-periodic coefficients, constructed using tensor products of basis functions of state variables with basis functions of time.

fail.value

optional numeric scalar; if non-NA, this value is substituted for non-finite values of the objective function. It should be a large number (i.e., bigger than any legitimate values the objective function is likely to take).

params

optional; named numeric vector of parameters. This will be coerced internally to storage mode double.

rinit

simulator of the initial-state distribution. This can be furnished either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator to its default. For more information, see rinit specification.

rprocess

simulator of the latent state process, specified using one of the rprocess plugins. Setting rprocess=NULL removes the latent-state simulator. For more information, see rprocess specification for the documentation on these plugins.

rmeasure

simulator of the measurement model, specified either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rmeasure=NULL removes the measurement model simulator. For more information, see rmeasure specification.

verbose

logical; if TRUE, diagnostic messages will be printed to the console.

Details

Nonlinear forecasting (NLF) is an 'indirect inference' method. The NLF approximation to the log likelihood of the data series is computed by simulating data from a model, fitting a nonlinear autoregressive model to the simulated time series, and quantifying the ability of the resulting fitted model to predict the data time series. The nonlinear autoregressive model is implemented as a

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generalized additive model (GAM), conditional on lagged values, for each observation variable. The errors are assumed multivariate normal.

The NLF objective function constructed by nlf_objfun simulates long time series (nasymp is the number of observations in the simulated times series), perhaps after allowing for a transient period (ntransient steps). It then fits the GAM for the chosen lags to the simulated time series. Finally, it computes the quasi-likelihood of the data under the fitted GAM.

NLF assumes that the observation frequency (equivalently the time between successive observations) is uniform.

Value

nlf_objfun constructs a stateful objective function for NLF estimation. Specfically, nlf_objfun returns an object of class 'nlf_objfun', which is a function suitable for use in an optim-like optimizer. In particular, this function takes a single numeric-vector argument that is assumed to contain the parameters named in est, in that order. When called, it will return the negative log quasilikelihood. It is a stateful function: Each time it is called, it will remember the values of the parameters and its estimate of the log quasilikelihood.

Periodically-forced systems (seasonality)

Unlike other **pomp** estimation methods, NLF cannot accommodate general time-dependence in the model via explicit time-dependence or dependence on time-varying covariates. However, NLF can accommodate periodic forcing. It does this by including forcing phase as a predictor in the nonlinear autoregressive model. To accomplish this, one sets period to the period of the forcing (a positive numerical value). In this case, if tensor = FALSE, the effect is to add a periodic intercept in the autoregressive model. If tensor = TRUE, by contrast, the fitted model includes time-periodic coefficients, constructed using tensor products of basis functions of observables with basis functions of time.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Important Note

Since **pomp** cannot guarantee that the *final* call an optimizer makes to the function is a call *at* the optimum, it cannot guarantee that the parameters stored in the function are the optimal ones. Therefore, it is a good idea to evaluate the function on the parameters returned by the optimization routine, which will ensure that these parameters are stored.

Warning! Objective functions based on C snippets

If you use C snippets (see Csnippet), a dynamically loadable library will be built. As a rule, **pomp** functions load this library as needed and unload it when it is no longer needed. The stateful objective functions are an exception to this rule. For efficiency, calls to the objective function do not execute

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pompLoad or pompUnload: rather, it is assumed that pompLoad has been called before any call to the objective function. When a stateful objective function using one or more C snippets is created, pompLoad is called internally to build and load the library: therefore, within a single R session, if one creates a stateful objective function, one can freely call that objective function and (more to the point) pass it to an optimizer that calls it freely, without needing to call pompLoad. On the other hand, if one retrieves a stored objective function from a file, or passes one to another R session, one must call pompLoad before using it. Failure to do this will typically result in a segmentation fault (i.e., it will crash the R session).

Author(s)

Stephen P. Ellner, Bruce E. Kendall, Aaron A. King

References

S.P. Ellner, B.A. Bailey, G.V. Bobashev, A.R. Gallant, B.T. Grenfell, and D.W. Nychka. Noise and nonlinearity in measles epidemics: combining mechanistic and statistical approaches to population modeling. *American Naturalist* **151**, 425–440, 1998. doi:10.1086/286130.

B.E. Kendall, C.J. Briggs, W.W. Murdoch, P. Turchin, S.P. Ellner, E. McCauley, R.M. Nisbet, and S.N. Wood. Why do populations cycle? A synthesis of statistical and mechanistic modeling approaches. *Ecology* **80**, 1789–1805, 1999. doi:10.2307/176658.

B.E. Kendall, S.P. Ellner, E. McCauley, S.N. Wood, C.J. Briggs, W.W. Murdoch, and P. Turchin. Population cycles in the pine looper moth (*Bupalus piniarius*): dynamical tests of mechanistic hypotheses. *Ecological Monographs* **75** 259–276, 2005. doi:10.1890/034056.

See Also

```
optim subplex nloptr
```

More on **pomp** estimation algorithms: abc(), bsmc2(), estimation_algorithms, mif2(), pmcmc(), pomp-package, probe_match, spect_match

More on methods based on summary statistics: abc(), basic_probes, probe(), probe_match, spect(), spect_match

More on maximization-based estimation methods: mif2(), probe_match, spect_match, traj_match

Examples

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```
plot(simulate(m1))
```

objfun

}

Objective functions

Description

Methods common to **pomp** stateful objective functions

Important Note

Since **pomp** cannot guarantee that the *final* call an optimizer makes to the function is a call *at* the optimum, it cannot guarantee that the parameters stored in the function are the optimal ones. Therefore, it is a good idea to evaluate the function on the parameters returned by the optimization routine, which will ensure that these parameters are stored.

Warning! Objective functions based on C snippets

If you use C snippets (see Csnippet), a dynamically loadable library will be built. As a rule, pomp functions load this library as needed and unload it when it is no longer needed. The stateful objective functions are an exception to this rule. For efficiency, calls to the objective function do not execute pompLoad or pompUnload: rather, it is assumed that pompLoad has been called before any call to the objective function. When a stateful objective function using one or more C snippets is created, pompLoad is called internally to build and load the library: therefore, within a single R session, if one creates a stateful objective function, one can freely call that objective function and (more to the point) pass it to an optimizer that calls it freely, without needing to call pompLoad. On the other hand, if one retrieves a stored objective function from a file, or passes one to another R session, one must call pompLoad before using it. Failure to do this will typically result in a segmentation fault (i.e., it will crash the R session).

obs obs

Description

Extract the data array from a 'pomp' object.

Usage

```
## S4 method for signature 'pomp'
obs(object, vars, ..., format = c("array", "data.frame"))
## S4 method for signature 'listie'
obs(object, vars, ..., format = c("array", "data.frame"))
```

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Arguments

```
object an object of class 'pomp', or of a class extending 'pomp' vars names of variables to retrieve ... ignored format format of the returned object
```

See Also

```
Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, pred_mean(), pred_var(), saved_states(), spy(), states(), summary(), time(), timezero(), traces()
```

ou2

Two-dimensional discrete-time Ornstein-Uhlenbeck process

Description

ou2() constructs a 'pomp' object encoding a bivariate discrete-time Ornstein-Uhlenbeck process with noisy observations.

Usage

```
ou2(
   alpha_1 = 0.8,
   alpha_2 = -0.5,
   alpha_3 = 0.3,
   alpha_4 = 0.9,
   sigma_1 = 3,
   sigma_2 = -0.5,
   sigma_3 = 2,
   tau = 1,
   x1_0 = -3,
   x2_0 = 4,
   times = 1:100,
   t0 = 0
)
```

Arguments

```
alpha_1, alpha_2, alpha_3, alpha_4 entries of the \alpha matrix, in column-major order. That is, alpha_2 is in the lower-left position. 
 sigma_1, sigma_2, sigma_3 entries of the lower-triangular \sigma matrix. sigma_2 is the entry in the lower-left position.
```

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tau measurement error s.d. $x1_0, x2_0$ latent variable values at time t0 times vector of observation times t0 the zero time

Details

If the state process is $X(t) = (X_1(t), X_2(t))$, then

$$X(t+1) = \alpha X(t) + \sigma \epsilon(t),$$

where α and σ are 2x2 matrices, σ is lower-triangular, and $\epsilon(t)$ is standard bivariate normal. The observation process is $Y(t) = (Y_1(t), Y_2(t))$, where $Y_i(t) \sim \text{normal}(X_i(t), \tau)$.

Value

A 'pomp' object with simulated data.

See Also

More examples provided with **pomp**: blowflies, childhood_disease_data, compartmental_models, dacca(), ebola, gompertz(), pomp_examples, ricker(), rw2(), verhulst()

Examples

```
po <- ou2()
plot(po)
coef(po)
x <- simulate(po)
plot(x)
pf <- pfilter(po,Np=1000)
logLik(pf)</pre>
```

parameter_trans

parameter transformations

Description

Equipping models with parameter transformations to facilitate searches in constrained parameter spaces.

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Usage

```
parameter_trans(toEst, fromEst, ...)

## S4 method for signature 'NULL,NULL'
parameter_trans(toEst, fromEst, ...)

## S4 method for signature 'pomp_fun,pomp_fun'
parameter_trans(toEst, fromEst, ...)

## S4 method for signature 'Csnippet,Csnippet'
parameter_trans(toEst, fromEst, ..., log, logit, barycentric)

## S4 method for signature 'character,character'
parameter_trans(toEst, fromEst, ...)

## S4 method for signature 'function,function'
parameter_trans(toEst, fromEst, ...)
```

Arguments

toEst, fromEst procedures that perform transformation of model parameters to and from the

estimation scale, respectively. These can be furnished using C snippets, R func-

tions, or via procedures in an external, dynamically loaded library.

... ignored.

log names of parameters to be log transformed.

logit names of parameters to be logit transformed.

barycentric names of parameters to be collectively transformed according to the log barycen-

tric transformation. **Important note:** variables to be log-barycentrically trans-

formed *must be adjacent* in the parameter vector.

Details

When parameter transformations are desired, they can be integrated into the 'pomp' object via the partrans arguments using the parameter_trans function. As with the other basic model components, these should ordinarily be specified using C snippets. When doing so, note that:

1. The parameter transformation mapping a parameter vector from the scale used by the model codes to another scale, and the inverse transformation, are specified via a call to

```
\verb|parameter_trans(toEst,fromEst)|
```

2. The goal of these snippets is the transformation of the parameters from the natural scale to the estimation scale, and vice-versa. If p is the name of a variable on the natural scale, its value on the estimation scale is T_p. Thus the toEst snippet computes T_p given p whilst the fromEst snippet computes p given T_p.

3. Time-, state-, and covariate-dependent transformations are not allowed. Therefore, neither the time, nor any state variables, nor any of the covariates will be available in the context within which a parameter transformation snippet is executed.

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These transformations can also be specified using R functions with arguments chosen from among the parameters. Such an R function must also have the argument '...'. In this case, toEst should transform parameters from the scale that the basic components use internally to the scale used in estimation. fromEst should be the inverse of toEst.

Note that it is the user's responsibility to make sure that the transformations are mutually inverse. If obj is the constructed 'pomp' object, and coef(obj) is non-empty, a simple check of this property is

```
x <- coef(obj, transform = TRUE)
obj1 <- obj
coef(obj1, transform = TRUE) <- x
identical(coef(obj), coef(obj1))
identical(coef(obj1, transform=TRUE), x)</pre>
```

One can use the log and logit arguments of parameter_trans to name variables that should be log-transformed or logit-transformed, respectively. The barycentric argument can name sets of parameters that should be log-barycentric transformed.

Note that using the log, logit, or barycentric arguments causes C snippets to be generated. Therefore, you must make sure that variables named in any of these arguments are also mentioned in paramnames at the same time.

The logit transform is defined by

$$logit(\theta) = log \frac{\theta}{1 - \theta}.$$

The log barycentric transformation of variables $\theta_1, \dots, \theta_n$ is given by

logbarycentric
$$(\theta_1, \dots, \theta_n) = \left(\log \frac{\theta_1}{\sum_i \theta_i}, \dots, \log \frac{\theta_n}{\sum_i \theta_i}\right)$$
.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

See Also

partrans

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

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parmat

Create a matrix of parameters

Description

parmat is a utility that makes a vector of parameters suitable for use in **pomp** functions.

Usage

```
parmat(params, ...)
## S4 method for signature 'numeric'
parmat(params, nrep = 1, ..., names = NULL)
## S4 method for signature 'array'
parmat(params, nrep = 1, ..., names = NULL)
## S4 method for signature 'data.frame'
parmat(params, nrep = 1, ...)
```

Arguments

params named numeric vector or matrix of parameters.
... additional arguments, currently ignored.
nrep number of replicates (columns) desired.
names optional character; column names.

Value

parmat returns a matrix consisting of nrep copies of params.

Author(s)

Aaron A. King

Examples

```
# takes too long for R CMD check
## generate a bifurcation diagram for the Ricker map
p <- parmat(coef(ricker()),nrep=500)
p["r",] <- exp(seq(from=1.5,to=4,length=500))
trajectory(
   ricker(),
   times=seq(from=1000,to=2000,by=1),
   params=p,
   format="array"
) -> x
```

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```
matplot(p["r",],x["N",,],pch='.',col='black',
    xlab=expression(log(r)),ylab="N",log='x')
```

partrans

partrans workhorse

Description

Performs parameter transformations.

Usage

```
## S4 method for signature 'pomp'
partrans(object, params, ..., dir = c("fromEst", "toEst"))
## S4 method for signature 'objfun'
partrans(object, ...)
```

Arguments

object	an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the pomp inference algorithms.
params	a npar x nrep matrix of parameters. Each column is treated as an independent parameter set, in correspondence with the corresponding column of x .
	additional arguments are ignored.
dir	the direction of the transformation to perform.

Value

If dir=fromEst, the parameters in params are assumed to be on the estimation scale and are transformed onto the natural scale. If dir=toEst, they are transformed onto the estimation scale. In both cases, the parameters are returned as a named numeric vector or an array with rownames, as appropriate.

See Also

```
Specification of parameter transformations: parameter_trans
```

```
More on pomp workhorse functions: dinit(), dmeasure(), dprior(), dprocess(), emeasure(), flow(), pomp-package, rinit(), rmeasure(), rprior(), rprocess(), skeleton(), vmeasure(), workhorses
```

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parus

Parus major population dynamics

Description

Size of a population of great tits (Parus major) from Wytham Wood, near Oxford.

Details

Provenance: Global Population Dynamics Database dataset #10163. (NERC Centre for Population Biology, Imperial College (2010) The Global Population Dynamics Database Version 2. https://www.imperial.ac.uk/cpb/gpdd2/). Original source: McCleer and Perrins (1991).

References

R. McCleery and C. Perrins. Effects of predation on the numbers of Great Tits, *Parus major*. In: C.M. Perrins, J.-D. Lebreton, and G.J.M. Hirons (eds.), *Bird Population Studies*, pp. 129–147, Oxford. Univ. Press, 1991. doi:10.1093/oso/9780198577300.003.0006.

See Also

More data sets provided with **pomp**: blowflies, bsflu, childhood_disease_data, dacca(), ebola

Examples

```
# takes too long for R CMD check
parus |>
  pfilter(Np=1000, times="year", t0=1960,
     params=c(K=190,r=2.7,sigma=0.2,theta=0.05,N.0=148),
     rprocess=discrete_time(
       function (r, K, sigma, N, ...) {
         e <- rnorm(n=1,mean=0,sd=sigma)
        c(N = exp(log(N)+r*(1-N/K)+e))
       },
       delta.t=1
     rmeasure=function (N, theta, ...) {
       c(pop=rnbinom(n=1, size=1/theta, mu=N+1e-10))
     dmeasure=function (pop, N, theta, ..., log) {
       dnbinom(x=pop,mu=N+1e-10,size=1/theta,log=log)
     partrans=parameter_trans(log=c("sigma","theta","N_0","r","K")),
     paramnames=c("sigma","theta","N_0","r","K")
   ) -> pf
pf |> logLik()
```

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```
pf |> simulate() |> plot()
```

pfilter

Particle filter

Description

A plain vanilla sequential Monte Carlo (particle filter) algorithm. Resampling is performed at each observation.

Usage

```
## S4 method for signature 'data.frame'
pfilter(
 data,
  . . . ,
 Nρ,
  params,
  rinit,
  rprocess,
  dmeasure,
  pred.mean = FALSE,
  pred.var = FALSE,
  filter.mean = FALSE,
  filter.traj = FALSE,
 save.states = c("no", "filter", "prediction", "weighted", "unweighted", "FALSE",
    "TRUE"),
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
pfilter(
  data,
  ...,
 Nρ,
  pred.mean = FALSE,
  pred.var = FALSE,
  filter.mean = FALSE,
  filter.traj = FALSE,
 save.states = c("no", "filter", "prediction", "weighted", "unweighted", "FALSE",
    "TRUE"),
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pfilterd_pomp'
pfilter(data, ..., Np, verbose = getOption("verbose", FALSE))
```

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```
## S4 method for signature 'objfun'
pfilter(data, ...)
```

Arguments

data either a data frame holding the time series data, or an object of class 'pomp',

i.e., the output of another \boldsymbol{pomp} calculation. Internally, data will be coerced to

an array with storage-mode double.

additional arguments are passed to pomp. This allows one to set, unset, or modify

basic model components within a call to this function.

Np the number of particles to use. This may be specified as a single positive integer, in which case the same number of particles will be used at each timestep. Al-

ternatively, if one wishes the number of particles to vary across timesteps, one may specify Np either as a vector of positive integers of length

length(time(object,t0=TRUE))

or as a function taking a positive integer argument. In the latter case, Np(k) must be a single positive integer, representing the number of particles to be used at the k-th timestep: Np(0) is the number of particles to use going from timezero(object) to time(object)[1], Np(1), from timezero(object) to time(object)[1], and so on, while when T=length(time(object)), Np(T)

is the number of particles to sample at the end of the time-series.

params optional; named numeric vector of parameters. This will be coerced internally

to storage mode double.

rinit simulator of the initial-state distribution. This can be furnished either as a C

snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator

to its default. For more information, see rinit specification.

rprocess simulator of the latent state process, specified using one of the rprocess plugins.

Setting rprocess=NULL removes the latent-state simulator. For more informa-

tion, see rprocess specification for the documentation on these plugins.

dmeasure evaluator of the measurement model density, specified either as a C snippet, an

R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting dmeasure=NULL removes the measurement density

evaluator. For more information, see dmeasure specification.

pred.mean logical; if TRUE, the prediction means are calculated for the state variables and

parameters.

pred.var logical; if TRUE, the prediction variances are calculated for the state variables

and parameters.

filter.mean logical; if TRUE, the filtering means are calculated for the state variables and

parameters.

filter.traj logical; if TRUE, a filtered trajectory is returned for the state variables and pa-

rameters. See filter_traj for more information.

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save.states

character; If save.states="no" (the default), information on the latent states is not saved. If save.states="filter", the state-vector for each filtered particle $X_{n,j}^F$ at each time n is saved. If save.states="prediction", the state-vector for each prediction particle $X_{n,j}^P$ at each time n is saved, along with the corresponding weight $w_{n,j} = f_{Y_n|X_n}(y^*|X_{n,j}^P;\theta)$. The options "unweighted", "weighted", TRUE, and FALSE are deprecated and will issue a warning if used, mapping to the new values for backward compatibility. The options "unweighted" and TRUE are synonymous with "filter"; the option "weighted" is synonymous with "prediction"; the option FALSE is synonymous with "no". To retrieve the saved states, apply saved_states to the result of the pfilter computation.

verbose

logical; if TRUE, diagnostic messages will be printed to the console.

Value

An object of class 'pfilterd_pomp', which extends class 'pomp'. Information can be extracted from this object using the methods documented below.

Methods

```
logLik the estimated log likelihood

cond_logLik the estimated conditional log likelihood

eff_sample_size the (time-dependent) estimated effective sample size

pred_mean, pred_var the mean and variance of the approximate prediction distribution

filter_mean the mean of the filtering distribution

filter_traj retrieve one particle trajectory. Useful for building up the smoothing distribution.

saved_states retrieve saved states

as.data.frame coerce to a data frame

plot diagnostic plots
```

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Author(s)

Aaron A. King

References

M.S. Arulampalam, S. Maskell, N. Gordon, and T. Clapp. A tutorial on particle filters for online nonlinear, non-Gaussian Bayesian tracking. *IEEE Transactions on Signal Processing* **50**, 174–188, 2002. doi:10.1109/78.978374.

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A. Bhadra and E.L. Ionides. Adaptive particle allocation in iterated sequential Monte Carlo via approximating meta-models. *Statistics and Computing* **26**, 393–407, 2016. doi:10.1007/s11222-0149513x.

See Also

```
More on pomp elementary algorithms: elementary_algorithms, kalman, pomp-package, probe(), simulate(), spect(), trajectory(), wpfilter()

More on sequential Monte Carlo methods: bsmc2(), cond_logLik(), eff_sample_size(), filter_mean(), filter_traj(), kalman, mif2(), pmcmc(), pred_mean(), pred_var(), saved_states(), wpfilter()

More on full-information (i.e., likelihood-based) methods: bsmc2(), mif2(), pmcmc(), wpfilter()
```

Examples

```
pf <- pfilter(gompertz(),Np=1000) ## use 1000 particles

plot(pf)
logLik(pf)
cond_logLik(pf) ## conditional log-likelihoods
eff_sample_size(pf) ## effective sample size
logLik(pfilter(pf)) ## run it again with 1000 particles

## run it again with 2000 particles
pf <- pfilter(pf,Np=2000,filter.mean=TRUE,filter.traj=TRUE,save.states="filter")
fm <- filter_mean(pf) ## extract the filtering means
ft <- filter_traj(pf) ## one draw from the smoothing distribution
ss <- saved_states(pf,format="d") ## the latent-state portion of each particle

as(pf,"data.frame") |> head()
```

plot

pomp plotting facilities

Description

Diagnostic plots.

Usage

```
## S4 method for signature 'pomp_plottable'
plot(
    x,
    variables,
    panel = lines,
    nc = NULL,
    yax.flip = FALSE,
    mar = c(0, 5.1, 0, if (yax.flip) 5.1 else 2.1),
    oma = c(6, 0, 5, 0),
```

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```
axes = TRUE,
)
## S4 method for signature 'Pmcmc'
plot(x, ..., pars)
## S4 method for signature 'Abc'
plot(x, ..., pars, scatter = FALSE)
## S4 method for signature 'Mif2'
plot(x, ..., pars, transform = FALSE)
## S4 method for signature 'probed_pomp'
plot(x, y, ...)
## S4 method for signature 'spectd_pomp'
 Х,
  ...,
 max.plots.per.page = 4,
 plot.data = TRUE,
 quantiles = c(0.025, 0.25, 0.5, 0.75, 0.975),
 quantile.styles = list(lwd = 1, lty = 1, col = "gray70"),
 data.styles = list(lwd = 2, lty = 2, col = "black")
)
## S4 method for signature 'bsmcd_pomp'
plot(x, pars, thin, ...)
## S4 method for signature 'probe_match_objfun'
plot(x, y, ...)
## S4 method for signature 'spect_match_objfun'
plot(x, y, ...)
```

Arguments

x	the object to plot
variables	optional character; names of variables to be displayed
panel	function of the form $panel(x, col, bg, pch, type,)$ which gives the action to be carried out in each panel of the display.
nc	the number of columns to use. Defaults to 1 for up to 4 series, otherwise to 2.
yax.flip	logical; if TRUE, the y-axis (ticks and numbering) should flip from side 2 (left) to 4 (right) from series to series.
mar, oma	the par mar and oma settings. Modify with care!
axes	logical; indicates if x- and y- axes should be drawn

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ignored or passed to low-level plotting functions

pars names of parameters.

scatter logical; if FALSE, traces of the parameters named in pars will be plotted against ABC iteration number. If TRUE, the traces will be displayed or as a scatterplot.

transform logical; should the parameter be transformed onto the estimation scale?

y ignored

max.plots.per.page

positive integer; maximum number of plots on a page

plot.data logical; should the data spectrum be included?

quantiles numeric; quantiles to display

quantities numeric, quantities to display

quantile.styles

list; plot styles to use for quantiles

data.styles list; plot styles to use for data

thin integer; when the number of samples is very large, it can be helpful to plot a

random subsample: thin specifies the size of this subsample.

pmcmc

. . .

The particle Markov chain Metropolis-Hastings algorithm

Description

The Particle MCMC algorithm for estimating the parameters of a partially-observed Markov process. Running pmcmc causes a particle random-walk Metropolis-Hastings Markov chain algorithm to run for the specified number of proposals.

Usage

```
## S4 method for signature 'data.frame'
pmcmc(
  data,
  . . . ,
 Nmcmc = 1,
  proposal,
 Nρ,
  params,
  rinit,
  rprocess,
  dmeasure,
  dprior,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
pmcmc(
```

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```
data,
 Nmcmc = 1,
  proposal,
 Np,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pfilterd_pomp'
pmcmc(
  data,
  . . . ,
 Nmcmc = 1,
 proposal,
 Np,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pmcmcd_pomp'
pmcmc(data, ..., Nmcmc, proposal, verbose = getOption("verbose", FALSE))
```

Arguments

data

either a data frame holding the time series data, or an object of class 'pomp', i.e., the output of another **pomp** calculation. Internally, data will be coerced to an array with storage-mode double.

. . .

additional arguments are passed to pomp. This allows one to set, unset, or modify basic model components within a call to this function.

Nmcmc

The number of PMCMC iterations to perform.

proposal

optional function that draws from the proposal distribution. Currently, the proposal distribution must be symmetric for proper inference: it is the user's responsibility to ensure that it is. Several functions that construct appropriate proposal function are provided: see MCMC proposals for more information.

Np

the number of particles to use. This may be specified as a single positive integer, in which case the same number of particles will be used at each timestep. Alternatively, if one wishes the number of particles to vary across timesteps, one may specify Np either as a vector of positive integers of length

```
length(time(object,t0=TRUE))
```

or as a function taking a positive integer argument. In the latter case, Np(k) must be a single positive integer, representing the number of particles to be used at the k-th timestep: Np(0) is the number of particles to use going from timezero(object) to time(object)[1], Np(1), from timezero(object) to time(object)[1], and so on, while when T=length(time(object)), Np(T) is the number of particles to sample at the end of the time-series.

params

optional; named numeric vector of parameters. This will be coerced internally to storage mode double.

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rinit simulator of the initial-state distribution. This can be furnished either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator to its default. For more information, see rinit specification. rprocess simulator of the latent state process, specified using one of the rprocess plugins. Setting rprocess=NULL removes the latent-state simulator. For more information, see rprocess specification for the documentation on these plugins. dmeasure evaluator of the measurement model density, specified either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting dmeasure=NULL removes the measurement density evaluator. For more information, see dmeasure specification. dprior optional; prior distribution density evaluator, specified either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. For more information, see prior specification. Setting dprior=NULL resets the prior distribution to its default, which is a flat improper prior. verbose logical; if TRUE, diagnostic messages will be printed to the console.

Value

An object of class 'pmcmcd_pomp'.

Methods

The following can be applied to the output of a pmcmc operation:

pmcmc repeats the calculation, beginning with the last state

continue continues the pmcmc calculation

plot produces a series of diagnostic plots

filter_traj extracts a random sample from the smoothing distribution

traces produces an mcmc object, to which the various coda convergence diagnostics can be applied

Re-running PMCMC Iterations

To re-run a sequence of PMCMC iterations, one can use the pmcmc method on a 'pmcmc' object. By default, the same parameters used for the original PMCMC run are re-used (except for verbose, the default of which is shown above). If one does specify additional arguments, these will override the defaults.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Author(s)

Edward L. Ionides, Aaron A. King, Sebastian Funk

References

C. Andrieu, A. Doucet, and R. Holenstein. Particle Markov chain Monte Carlo methods. *Journal of the Royal Statistical Society, Series B* **72**, 269–342, 2010. doi:10.1111/j.14679868.2009.00736.x.

See Also

```
More on pomp estimation algorithms: abc(), bsmc2(), estimation_algorithms, mif2(), nlf, pomp-package, probe_match, spect_match
```

```
More on sequential Monte Carlo methods: bsmc2(), cond_logLik(), eff_sample_size(), filter_mean(), filter_traj(), kalman, mif2(), pfilter(), pred_mean(), pred_var(), saved_states(), wpfilter()
```

More on full-information (i.e., likelihood-based) methods: bsmc2(), mif2(), pfilter(), wpfilter()

More on Markov chain Monte Carlo methods: abc(), proposals

More on Bayesian methods: abc(), bsmc2(), dprior(), prior_spec, rprior()

pomp-class

The basic pomp class

Description

The basic class implementing a POMP model with data

See Also

pomp_constructor

pomp_constructor

Constructor of the basic pomp object

Description

This function constructs a 'pomp' object, encoding a partially-observed Markov process (POMP) model together with a uni- or multi-variate time series. As such, it is central to all the package's functionality. One implements the POMP model by specifying some or all of its *basic components*. These comprise:

rinit which samples from the distribution of the state process at the zero-time;

dinit which evaluates the density of the state process at the zero-time;

rprocess the simulator of the unobserved Markov state process;

dprocess the evaluator of the probability density function for transitions of the unobserved Markov state process;

rmeasure the simulator of the observed process, conditional on the unobserved state;

dmeasure the evaluator of the measurement model probability density function;

emeasure the expectation of the measurements, conditional on the latent state;

vmeasure the covariance matrix of the measurements, conditional on the latent state;

rprior which samples from a prior probability distribution on the parameters;

dprior which evaluates the prior probability density function;

skeleton which computes the deterministic skeleton of the unobserved state process;

partrans which performs parameter transformations.

The basic structure and its rationale are described in the *Journal of Statistical Software* paper, an updated version of which is to be found on the package website.

Usage

```
pomp(
  data,
  . . . ,
  times,
  t0,
  rinit,
  dinit,
  rprocess,
  dprocess,
  rmeasure,
  dmeasure,
  emeasure,
  vmeasure,
  skeleton,
  rprior,
  dprior,
  partrans,
  covar,
  params,
  accumvars,
  obsnames,
  statenames,
  paramnames,
  covarnames,
  nstatevars,
  PACKAGE,
  globals,
  on_load,
  userdata,
  cdir = getOption("pomp_cdir", NULL),
  cfile,
```

```
shlib.args,
compile = TRUE,
verbose = getOption("verbose", FALSE)
)
```

Arguments

data either a data frame holding the time series data, or an object of class 'pomp',

i.e., the output of another ${\bf pomp}$ calculation. Internally, data will be coerced to

an array with storage-mode double.

... additional arguments will generate an error.

times the sequence of observation times. times must indicate the column of obser-

vation times by name or index. The time vector must be numeric and non-

decreasing.

to The zero-time, i.e., the time of the initial state. This must be no later than the

time of the first observation, i.e., t0 <= times[1].

rinit simulator of the initial-state distribution. This can be furnished either as a C

snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator

to its default. For more information, see rinit specification.

dinit evaluator of the initial-state density. This can be furnished either as a C snip-

pet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting dinit=NULL removes this basic component.

For more information, see dinit specification.

rprocess simulator of the latent state process, specified using one of the rprocess plugins.

Setting rprocess=NULL removes the latent-state simulator. For more informa-

tion, see rprocess specification for the documentation on these plugins.

dprocess evaluator of the probability density of transitions of the unobserved state pro-

cess. Setting dprocess=NULL removes the latent-state density evaluator. For

more information, see dprocess specification.

rmeasure simulator of the measurement model, specified either as a C snippet, an R func-

tion, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rmeasure=NULL removes the measurement model simu-

lator. For more information, see rmeasure specification.

dmeasure evaluator of the measurement model density, specified either as a C snippet, an

R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting dmeasure=NULL removes the measurement density

evaluator. For more information, see dmeasure specification.

emeasure the expectation of the measured variables, conditional on the latent state. This

can be specified as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting emeasure=NULL removes the emeasure component. For more information, see emeasure specifi-

cation.

vmeasure the covariance of the measured variables, conditional on the latent state. This

can be specified as a C snippet, an R function, or the name of a pre-compiled

native routine available in a dynamically loaded library. Setting vmeasure=NULL removes the vmeasure component. For more information, see vmeasure specification.

skeleton

optional; the deterministic skeleton of the unobserved state process. Depending on whether the model operates in continuous or discrete time, this is either a vectorfield or a map. Accordingly, this is supplied using either the vectorfield or map fnctions. For more information, see skeleton specification. Setting skeleton=NULL removes the deterministic skeleton.

rprior

optional; prior distribution sampler, specified either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. For more information, see prior specification. Setting rprior=NULL removes the prior distribution sampler.

dprior

optional; prior distribution density evaluator, specified either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. For more information, see prior specification. Setting dprior=NULL resets the prior distribution to its default, which is a flat improper prior.

partrans

optional parameter transformations, constructed using parameter_trans.

Many algorithms for parameter estimation search an unconstrained space of parameters. When working with such an algorithm and a model for which the parameters are constrained, it can be useful to transform parameters. One should supply the partrans argument via a call to parameter_trans. For more information, see parameter_trans. Setting partrans=NULL removes the parameter transformations, i.e., sets them to the identity transformation.

covar

optional covariate table, constructed using covariate_table.

If a covariate table is supplied, then the value of each of the covariates is interpolated as needed. The resulting interpolated values are made available to the appropriate basic components. See the documentation for covariate_table for details.

params

optional; named numeric vector of parameters. This will be coerced internally to storage mode double.

accumvars

optional character vector; contains the names of accumulator variables. See accumvars for a definition and discussion of accumulator variables.

obsnames

optional character vector; names of the observables. It is not usually necessary to specify obsnames since, by default, these are read from the names of the data variables.

statenames

optional character vector; names of the latent state variables. It is typically only necessary to supply statenames when C snippets are in use. See also nstatevars.

paramnames

optional character vector; names of model parameters. It is typically only necessary to supply paramnames when C snippets are in use.

covarnames

optional character vector; names of the covariates. It is not usually necessary to specify covarnames since, by default, these are read from the names of the covariates.

nstatevars optional integer scalar; If C snippets or native routines are used, one can specify the number of state variables with this argument. By default, nstatevars = length(statenames). **PACKAGE** optional character; the name (without extension) of the external, dynamically loaded library in which any native routines are to be found. This is only useful if one or more of the model components has been specified using a precompiled dynamically loaded library; it is not used for any component specified using C snippets. PACKAGE can name at most one library. globals optional character or C snippet; arbitrary C code that will be hard-coded into the shared-object library created when C snippets are provided. If no C snippets are used, globals has no effect. on_load optional character or C snippet: arbitrary C code that will be executed when the C snippet file is loaded. If no C snippets are used, on_load has no effect. userdata optional list; the elements of this list will be available to basic model components. This allows the user to pass information to the basic components outside of the usual routes of covariates (covar) and model parameters (params). See userdata for information on how to use this facility. cdir optional character variable. cdir specifies the name of the directory within which C snippet code will be compiled. By default, this is in a temporary directory specific to the R session. One can also set this directory using the pomp_cdir global option. cfile optional character variable. cfile gives the name of the file (in directory cdir) into which C snippet codes will be written. By default, a random filename is used. If the chosen filename would result in over-writing an existing file, an error is generated. optional character variables. Command-line arguments to the R CMD SHLIB call shlib.args that compiles the C snippets. One can, for example, specify libraries against which the C snippets are to be linked. In doing so, take care to make sure the appropriate header files are available to the C snippets, e.g., using the globals argument. See Csnippet for more information. logical; if FALSE, compilation of the C snippets will be postponed until they are compile needed. logical; if TRUE, diagnostic messages will be printed to the console. verbose

Details

Each basic component is supplied via an argument of the same name. These can be given in the call to pomp, or to many of the package's other functions. In any case, the effect is the same: to add, remove, or modify the basic component.

Each basic component can be furnished using C snippets, R functions, or pre-compiled native routine available in user-provided dynamically loaded libraries.

Value

The pomp constructor function returns an object, call it P, of class 'pomp'. P contains, in addition to the data, any elements of the model that have been specified as arguments to the pomp constructor

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function. One can add or modify elements of P by means of further calls to pomp, using P as the first argument in such calls. One can pass P to most of the **pomp** package methods via their data argument.

Note

It is not typically necessary (or indeed feasible) to define all of the basic components for any given purpose. However, each **pomp** algorithm makes use of only a subset of these components. When an algorithm requires a basic component that has not been furnished, an error is generated to let you know that you must provide the needed component to use the algorithm.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Author(s)

Aaron A. King

References

A. A. King, D. Nguyen, and E. L. Ionides. Statistical inference for partially observed Markov processes via the R package **pomp**. *Journal of Statistical Software* **69**(12), 1–43, 2016. doi:10.18637/jss.v069.i12. An updated version of this paper is available on the **pomp** package website.

See Also

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

pomp_examples

pre-built pomp examples

Description

Examples of pomp objects containing models and data.

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Details

pomp includes a number of pre-built examples of pomp objects and data that can be analyzed using **pomp** methods. These include:

blowflies Data from Nicholson's experiments with sheep blowfly populations

blowflies1() A pomp object with some of the blowfly data together with a discrete delay equation model.

blowflies2() A variant of blowflies1.

bsflu Data from an outbreak of influenza in a boarding school.

dacca() Fifty years of census and cholera mortality data, together with a stochastic differential equation transmission model (King et al. 2008).

ebolaModel() Data from the 2014 West Africa outbreak of Ebola virus disease, together with simple transmission models (King et al. 2015).

gompertz() The Gompertz population dynamics model, with simulated data.

LondonYorke Data on incidence of several childhood diseases (London and Yorke 1973)

ewmeas Measles incidence data from England and Wales

ewcitmeas Measles incidence data from 7 English cities

ou2() A 2-D Ornstein-Uhlenbeck process with simulated data

parus Population censuses of a Parus major population in Wytham Wood, England.

ricker The Ricker population dynamics model, with simulated data

rw2 A 2-D Brownian motion model, with simulated data.

sir() A simple continuous-time Markov chain SIR model, coded using Euler-multinomial steps, with simulated data.

sir2() A simple continuous-time Markov chain SIR model, coded using Gillespie's algorithm, with simulated data.

verhulst() The Verhulst-Pearl (logistic) model, a continuous-time model of population dynamics, with simulated data

See also the tutorials on the package website for more examples.

References

Anonymous. Influenza in a boarding school. British Medical Journal 1, 587, 1978.

A.A. King, E.L. Ionides, M. Pascual, and M.J. Bouma. Inapparent infections and cholera dynamics. *Nature* **454**, 877-880, 2008. doi:10.1038/nature07084.

A.A. King, M. Domenech de Cellès, F.M.G. Magpantay, and P. Rohani. Avoidable errors in the modelling of outbreaks of emerging pathogens, with special reference to Ebola. *Proceedings of the Royal Society of London, Series B* **282**, 20150347, 2015. doi:10.1098/rspb.2015.0347.

W. P. London and J. A. Yorke, Recurrent outbreaks of measles, chickenpox and mumps: I. Seasonal variation in contact rates. *American Journal of Epidemiology* **98**, 453–468, 1973. doi:10.1093/oxfordjournals.aje.a121575.

A.J. Nicholson. The self-adjustment of populations to change. *Cold Spring Harbor Symposia on Quantitative Biology* **22**, 153–173, 1957. doi:10.1101/SQB.1957.022.01.017.

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See Also

More examples provided with **pomp**: blowflies, childhood_disease_data, compartmental_models, dacca(), ebola, gompertz(), ou2(), ricker(), rw2(), verhulst()

pomp_fun

The "pomp_fun" class

Description

Definition and methods of the 'pomp_fun' class.

Usage

```
## S4 method for signature 'missing'
pomp_fun(
  slotname = NULL,
 obsnames = character(0),
  statenames = character(0),
  paramnames = character(0),
  covarnames = character(0),
)
## S4 method for signature 'function'
pomp_fun(f, proto = NULL, slotname = NULL, ...)
## S4 method for signature 'character'
pomp_fun(
  f,
 PACKAGE = NULL,
 obsnames = character(0),
  statenames = character(0),
  paramnames = character(0),
  covarnames = character(0),
  slotname = NULL,
)
## S4 method for signature 'Csnippet'
pomp_fun(
  f,
  slotname = NULL,
  libname = NULL,
  obsnames = character(0),
  statenames = character(0),
  paramnames = character(0),
```

pred_mean

```
covarnames = character(0),
  Cname,
  ...
)

## S4 method for signature 'pomp_fun'
pomp_fun(f, ...)
```

Arguments

f A function or the name of a native routine.

proto optional string; a prototype against which f will be checked.

PACKAGE optional; the name of the dynamically-loadable library in which the native func-

tion f can be found.

Details

The 'pomp_fun' class implements a common interface for user-defined procedures that can be defined in terms of R code or by compiled native routines.

Author(s)

Aaron A. King

See Also

pomp

pred_mean

Prediction mean

Description

The mean of the prediction distribution

Usage

```
## S4 method for signature 'kalmand_pomp'
pred_mean(object, vars, ..., format = c("array", "data.frame"))
## S4 method for signature 'pfilterd_pomp'
pred_mean(object, vars, ..., format = c("array", "data.frame"))
```

Arguments

object result of a filtering computation
vars optional character; names of variables

... ignored

format of the returned object

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Details

The prediction distribution is that of

$$X(t_k)|Y(t_1) = y_1^*, \dots, Y(t_{k-1}) = y_{k-1}^*,$$

where $X(t_k)$, $Y(t_k)$ are the latent state and observable processes, respectively, and y_k^* is the data, at time t_k .

The prediction mean is therefore the expectation of this distribution

$$E[X(t_k)|Y(t_1)=y_1^*,\ldots,Y(t_{k-1})=y_{k-1}^*].$$

See Also

More on sequential Monte Carlo methods: bsmc2(), cond_logLik(), eff_sample_size(), filter_mean(), filter_traj(), kalman, mif2(), pfilter(), pmcmc(), pred_var(), saved_states(), wpfilter()

Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_var(), saved_states(), spy(), states(), summary(), time(), timezero(), traces()

pred_var

Prediction variance

Description

The variance of the prediction distribution

Usage

```
## S4 method for signature 'pfilterd_pomp'
pred_var(object, vars, ..., format = c("array", "data.frame"))
```

Arguments

object result of a filtering computation

vars optional character; names of variables

ignored

... ignored

format format of the returned object

Details

The prediction distribution is that of

$$X(t_k)|Y(t_1) = y_1^*, \dots, Y(t_{k-1}) = y_{k-1}^*,$$

where $X(t_k)$, $Y(t_k)$ are the latent state and observable processes, respectively, and y_k^* is the data, at time t_k .

The prediction variance is therefore the variance of this distribution

$$Var[X(t_k)|Y(t_1) = y_1^*, \dots, Y(t_{k-1}) = y_{k-1}^*].$$

prior_spec

See Also

```
More on sequential Monte Carlo methods: bsmc2(), cond_logLik(), eff_sample_size(), filter_mean(), filter_traj(), kalman, mif2(), pfilter(), pmcmc(), pred_mean(), saved_states(), wpfilter()

Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_mean(), saved_states(), spy(), states(), summary(), time(), timezero(), traces()
```

print

Print methods

Description

These methods print their argument and return it *invisibly*.

Usage

```
## S4 method for signature 'unshowable'
print(x, ...)

## S4 method for signature 'listie'
print(x, ...)

## S4 method for signature 'pomp_fun'
print(x, ...)
```

Arguments

```
x object to print
... ignored
```

prior_spec

prior specification

Description

Specification of prior distributions via the rprior and dprior components.

prior_spec 113

Details

A prior distribution on parameters is specified by means of the rprior and/or dprior arguments to pomp. As with the other basic model components, it is preferable to specify these using C snippets. In writing a C snippet for the prior sampler (rprior), keep in mind that:

- 1. Within the context in which the snippet will be evaluated, only the parameters will be defined.
- 2. The goal of such a snippet is the replacement of parameters with values drawn from the prior distribution.
- 3. Hyperparameters can be included in the ordinary parameter list. Obviously, hyperparameters should not be replaced with random draws.

In writing a C snippet for the prior density function (dprior), observe that:

- Within the context in which the snippet will be evaluated, only the parameters and give_log will be defined.
- 2. The goal of such a snippet is computation of the prior probability density, or the log of same, at a given point in parameter space. This scalar value should be returned in the variable lik. When give_log == 1, lik should contain the log of the prior probability density.
- 3. Hyperparameters can be included in the ordinary parameter list.

General rules for writing C snippets can be found here.

Alternatively, one can furnish R functions for one or both of these arguments. In this case, rprior must be a function that makes a draw from the prior distribution of the parameters and returns a named vector containing all the parameters. The only required argument of this function is

Similarly, the dprior function must evaluate the prior probability density (or log density if log == TRUE) and return that single scalar value. The only required arguments of this function are . . . and log.

Default behavior

By default, the prior is assumed flat and improper. In particular, dprior returns 1 (0 if log = TRUE) for every parameter set. Since it is impossible to simulate from a flat improper prior, rprocess returns missing values (NAs).

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

See Also

dprior rprior

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom,

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parameter_trans(), pomp-package, pomp_constructor, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

More on Bayesian methods: abc(), bsmc2(), dprior(), pmcmc(), rprior()

Examples

```
# takes too long for R CMD check
## Starting with an existing pomp object:
verhulst() |> window(end=30) -> po
## We add or change prior distributions using the two
## arguments 'rprior' and 'dprior'. Here, we introduce
## a Gamma prior on the 'r' parameter.
## We construct 'rprior' and 'dprior' using R functions.
<| oq
  bsmc2(
     rprior=function (n_0, K0, K1, sigma, tau, r0, r1, ...) {
      c(
        n_0 = n_0
         K = rgamma(n=1,shape=K0,scale=K1),
         r = rgamma(n=1, shape=r0, scale=r1),
        sigma = sigma,
         tau = tau
      )
     },
     dprior=function(K, K0, K1, r, r0, r1, ..., log) {
      p \leftarrow dgamma(x=c(K,r), shape=c(K0,r0), scale=c(K1,r1), log=log)
      if (log) sum(p) else prod(p)
     params=c(n_0=10000,K=10000,K0=10,K1=1000,
       r=0.9, r0=0.9, r1=1, sigma=0.5, tau=0.3),
     Np=1000
   ) -> B
## We can also pass them as C snippets:
po |>
  bsmc2(
     rprior=Csnippet("
       K = rgamma(K0,K1);
        r = rgamma(r0,r1);
     ),
     dprior=Csnippet("
        double lik1 = dgamma(K,K0,K1,give_log);
        double lik2 = dgamma(r,r0,r1,give_log);
        lik = (give_log) ? lik1+lik2 : lik1*lik2;"
     ),
     paramnames=c("K","K0","K1","r","r0","r1"),
     params=c(n_0=10000,K=10000,K0=10,K1=1000,
       r=0.9,r0=0.9,r1=1,sigma=0.5,tau=0.3),
     Np=10000
```

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```
## The prior is plotted in grey; the posterior, in blue.
plot(B)

B |>
    pmcmc(Nmcmc=100,Np=1000,proposal=mvn_diag_rw(c(r=0.01,K=10))) -> Bb

plot(Bb,pars=c("loglik","log.prior","r","K"))
```

probe

Probes (AKA summary statistics)

Description

Probe a partially-observed Markov process by computing summary statistics and the synthetic likelihood.

Usage

```
## S4 method for signature 'data.frame'
probe(
  data,
  ...,
  probes,
  nsim,
  seed = NULL,
  params,
  rinit,
  rprocess,
  rmeasure,
  verbose = getOption("verbose", FALSE)
## S4 method for signature 'pomp'
probe(
  data,
  probes,
  nsim,
  seed = NULL,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'probed_pomp'
probe(
```

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```
data,
...,
probes,
nsim,
seed = NULL,
verbose = getOption("verbose", FALSE)
)

## S4 method for signature 'probe_match_objfun'
probe(data, ..., seed, verbose = getOption("verbose", FALSE))

## S4 method for signature 'objfun'
probe(data, ..., seed = NULL)
```

Arguments

nsim

data	either a data frame holding the time series data, or an object of class 'pomp',
	i.e., the output of another pomp calculation. Internally, data will be coerced to

an array with storage-mode double.

... additional arguments are passed to pomp. This allows one to set, unset, or modify

basic model components within a call to this function.

probes a single probe or a list of one or more probes. A probe is simply a scalar- or

vector-valued function of one argument that can be applied to the data array of a 'pomp'. A vector-valued probe must always return a vector of the same size. A number of useful probes are provided with the package: see basic probes.

the number of model simulations to be computed.

seed optional integer; if set, the pseudorandom number generator (RNG) will be ini-

tialized with seed. The RNG will be restored to its original state afterward.

params optional; named numeric vector of parameters. This will be coerced internally

to storage mode double.

rinit simulator of the initial-state distribution. This can be furnished either as a C

snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator

to its default. For more information, see rinit specification.

rprocess simulator of the latent state process, specified using one of the rprocess plugins.

Setting rprocess=NULL removes the latent-state simulator. For more informa-

tion, see rprocess specification for the documentation on these plugins.

rmeasure simulator of the measurement model, specified either as a C snippet, an R func-

tion, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rmeasure=NULL removes the measurement model simu-

lator. For more information, see rmeasure specification.

verbose logical; if TRUE, diagnostic messages will be printed to the console.

Details

probe applies one or more "probes" to time series data and model simulations and compares the

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results. It can be used to diagnose goodness of fit and/or as the basis for "probe-matching", a generalized method-of-moments approach to parameter estimation.

A call to probe results in the evaluation of the probe(s) in probes on the data. Additionally, nsim simulated data sets are generated (via a call to simulate) and the probe(s) are applied to each of these. The results of the probe computations on real and simulated data are stored in an object of class 'probed_pomp'.

When probe operates on a probe-matching objective function (a 'probe_match_objfun' object), by default, the random-number generator seed is fixed at the value given when the objective function was constructed. Specifying NULL or an integer for seed overrides this behavior.

Value

probe returns an object of class 'probed_pomp', which contains the data and the model, together with the results of the probe calculation.

Methods

The following methods are available.

plot displays diagnostic plots.

summary displays summary information. The summary includes quantiles (fractions of simulations with probe values less than those realized on the data) and the corresponding two-sided p-values. In addition, the "synthetic likelihood" (Wood 2010) is computed, under the assumption that the probe values are multivariate-normally distributed.

logLik returns the synthetic likelihood for the probes. NB: in general, this is not the same as the likelihood.

as.data.frame coerces a 'probed_pomp' to a 'data.frame'. The latter contains the realized values of the probes on the data and on the simulations. The variable .id indicates whether the probes are from the data or simulations.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Author(s)

Daniel C. Reuman, Aaron A. King

References

B.E. Kendall, C.J. Briggs, W.W. Murdoch, P. Turchin, S.P. Ellner, E. McCauley, R.M. Nisbet, and S.N. Wood. Why do populations cycle? A synthesis of statistical and mechanistic modeling approaches. *Ecology* **80**, 1789–1805, 1999. doi:10.2307/176658.

S. N. Wood Statistical inference for noisy nonlinear ecological dynamic systems. *Nature* **466**, 1102–1104, 2010. doi:10.1038/nature09319.

See Also

```
More on pomp elementary algorithms: elementary_algorithms, kalman, pfilter(), pomp-package, simulate(), spect(), trajectory(), wpfilter()
```

More on methods based on summary statistics: abc(), $basic_probes$, nlf, $probe_match$, $spect_match$

probe_match

Probe matching

Description

Estimation of parameters by maximum synthetic likelihood

Usage

```
## S4 method for signature 'data.frame'
probe_objfun(
  data,
  est = character(0),
  fail.value = NA,
  probes,
  nsim,
  seed = NULL,
  params,
  rinit,
  rprocess,
  rmeasure,
  partrans,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
probe_objfun(
  data,
  est = character(0),
  fail.value = NA,
 probes,
  nsim,
  seed = NULL,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'probed_pomp'
probe_objfun(
```

```
data,
  est = character(0),
  fail.value = NA,
  probes,
  nsim,
  seed = NULL,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'probe_match_objfun'
probe_objfun(
  data,
  . . . ,
  est.
  fail.value,
  seed = NULL,
  verbose = getOption("verbose", FALSE)
)
```

Arguments

data either a data frame holding the time series data, or an object of class 'pomp',

i.e., the output of another **pomp** calculation. Internally, data will be coerced to

an array with storage-mode double.

.. additional arguments are passed to pomp. This allows one to set, unset, or modify

basic model components within a call to this function.

est character vector; the names of parameters to be estimated.

fail.value optional numeric scalar; if non-NA, this value is substituted for non-finite values

of the objective function. It should be a large number (i.e., bigger than any

legitimate values the objective function is likely to take).

probes a single probe or a list of one or more probes. A probe is simply a scalar- or

vector-valued function of one argument that can be applied to the data array of a 'pomp'. A vector-valued probe must always return a vector of the same size.

A number of useful probes are provided with the package: see basic probes.

nsim the number of model simulations to be computed.

seed integer. When fitting, it is often best to fix the seed of the random-number

generator (RNG). This is accomplished by setting seed to an integer. By default,

seed = NULL, which does not alter the RNG state.

params optional; named numeric vector of parameters. This will be coerced internally

to storage mode double.

rinit simulator of the initial-state distribution. This can be furnished either as a C

snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator

to its default. For more information, see rinit specification.

rprocess simulator of the latent state process, specified using one of the rprocess plugins.

Setting rprocess=NULL removes the latent-state simulator. For more informa-

tion, see rprocess specification for the documentation on these plugins.

rmeasure simulator of the measurement model, specified either as a C snippet, an R func-

tion, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rmeasure=NULL removes the measurement model simu-

lator. For more information, see rmeasure specification.

partrans optional parameter transformations, constructed using parameter_trans.

Many algorithms for parameter estimation search an unconstrained space of parameters. When working with such an algorithm and a model for which the parameters are constrained, it can be useful to transform parameters. One should supply the partrans argument via a call to parameter_trans. For more information, see parameter_trans. Setting partrans=NULL removes the parameter

transformations, i.e., sets them to the identity transformation.

verbose logical; if TRUE, diagnostic messages will be printed to the console.

Details

In probe-matching, one attempts to minimize the discrepancy between simulated and actual data, as measured by a set of summary statistics called *probes*. In **pomp**, this discrepancy is measured using the "synthetic likelihood" as defined by Wood (2010).

Value

probe_objfun constructs a stateful objective function for probe matching. Specifically, probe_objfun returns an object of class 'probe_match_objfun', which is a function suitable for use in an optim-like optimizer. In particular, this function takes a single numeric-vector argument that is assumed to contain the parameters named in est, in that order. When called, it will return the negative synthetic log likelihood for the probes specified. It is a stateful function: Each time it is called, it will remember the values of the parameters and its estimate of the synthetic likelihood.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Important Note

Since **pomp** cannot guarantee that the *final* call an optimizer makes to the function is a call *at* the optimum, it cannot guarantee that the parameters stored in the function are the optimal ones. Therefore, it is a good idea to evaluate the function on the parameters returned by the optimization routine, which will ensure that these parameters are stored.

Warning! Objective functions based on C snippets

If you use C snippets (see Csnippet), a dynamically loadable library will be built. As a rule, pomp functions load this library as needed and unload it when it is no longer needed. The stateful objective functions are an exception to this rule. For efficiency, calls to the objective function do not execute pompLoad or pompUnload: rather, it is assumed that pompLoad has been called before any call to the objective function. When a stateful objective function using one or more C snippets is created, pompLoad is called internally to build and load the library: therefore, within a single R session, if one creates a stateful objective function, one can freely call that objective function and (more to the point) pass it to an optimizer that calls it freely, without needing to call pompLoad. On the other hand, if one retrieves a stored objective function from a file, or passes one to another R session, one must call pompLoad before using it. Failure to do this will typically result in a segmentation fault (i.e., it will crash the R session).

Author(s)

Aaron A. King

References

B.E. Kendall, C.J. Briggs, W.W. Murdoch, P. Turchin, S.P. Ellner, E. McCauley, R.M. Nisbet, and S.N. Wood. Why do populations cycle? A synthesis of statistical and mechanistic modeling approaches. *Ecology* **80**, 1789–1805, 1999. doi:10.2307/176658.

S. N. Wood Statistical inference for noisy nonlinear ecological dynamic systems. *Nature* **466**, 1102–1104, 2010. doi:10.1038/nature09319.

See Also

```
optim subplex nloptr
```

More on methods based on summary statistics: abc(), basic_probes, nlf, probe(), spect(), spect_match

More on **pomp** estimation algorithms: abc(), bsmc2(), estimation_algorithms, mif2(), nlf, pmcmc(), pomp-package, spect_match

More on maximization-based estimation methods: mif2(), nlf, spect_match, traj_match

Examples

```
gompertz() -> po

## A list of probes:
plist <- list(
    mean=probe_mean("Y",trim=0.1,transform=sqrt),
    sd=probe_sd("Y",transform=sqrt),
    probe_marginal("Y",ref=obs(po)),
    probe_acf("Y",lags=c(1,3,5),type="correlation",transform=sqrt),
    probe_quantile("Y",prob=c(0.25,0.75),na.rm=TRUE)
)

## Construct the probe-matching objective function.
## Here, we just want to estimate 'K'.</pre>
```

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```
probe_objfun(probes=plist,nsim=100,seed=5069977,
    est="K") -> f
## Any numerical optimizer can be used to minimize 'f'.
if (require(subplex)) {
  subplex(fn=f,par=0.4,control=list(reltol=1e-5)) -> out
} else {
 optim(fn=f,par=0.4,control=list(reltol=1e-5)) -> out
## Call the objective one last time on the optimal parameters:
f(out$par)
coef(f)
## There are 'plot' and 'summary' methods:
f |> as("probed_pomp") |> plot()
f |> summary()
## One can convert an objective function to a data frame:
f |> as("data.frame") |> head()
f |> as("probed_pomp") |> as("data.frame") |> head()
f |> probe() |> plot()
## One can modify the objective function with another call
## to 'probe_objfun':
f |> probe_objfun(est=c("r","K")) -> f1
optim(fn=f1,par=c(0.3,0.3),control=list(reltol=1e-5)) \rightarrow out
f1(out$par)
coef(f1)
```

proposals

MCMC proposal distributions

Description

Functions to construct proposal distributions for use with MCMC methods.

Usage

```
mvn_diag_rw(rw.sd)
mvn_rw(rw.var)
```

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```
mvn_rw_adaptive(
  rw.sd,
  rw.var,
  scale.start = NA,
  scale.cooling = 0.999,
  shape.start = NA,
  target = 0.234,
  max.scaling = 50
)
```

Arguments

rw.sd

named numeric vector; random-walk SDs for a multivariate normal random-walk proposal with diagonal variance-covariance matrix.

rw.var

square numeric matrix with row- and column-names. Specifies the variance-covariance matrix for a multivariate normal random-walk proposal distribution.

scale.start, scale.cooling, shape.start, target, max.scaling

parameters to control the proposal adaptation algorithm. Beginning with MCMC iteration scale.start, the scale of the proposal covariance matrix will be adjusted in an effort to match the target acceptance ratio. This initial scale adjustment is "cooled", i.e., the adjustment diminishes as the chain moves along. The parameter scale.cooling specifies the cooling schedule: at n iterations after scale.start, the current scaling factor is multiplied with scale.cooling^n. The maximum scaling factor allowed at any one iteration is max.scaling. After shape.start accepted proposals have accumulated, a scaled empirical covariance matrix will be used for the proposals, following Roberts and Rosenthal (2009).

Value

Each of these calls constructs a function suitable for use as the proposal argument of pmcmc or abc. Given a parameter vector, each such function returns a single draw from the corresponding proposal distribution.

Author(s)

Aaron A. King, Sebastian Funk

References

G.O. Roberts and J.S. Rosenthal. Examples of adaptive MCMC. *Journal of Computational and Graphical Statistics* **18**, 349–367, 2009. doi:10.1198/jcgs.2009.06134.

See Also

More on Markov chain Monte Carlo methods: abc(), pmcmc()

pStop

pStop, pWarn, pMess

Description

Custom error, warning, and message functions.

Usage

```
pStop(..., who = -1L)
pStop_(...)
pWarn(..., who = -1L)
pWarn_(...)
pMess(..., who = -1L)
pMess_(...)
```

Arguments

who

• • •

integer or character. If who is an integer, it is passed to sys.call to retrieve the name of the calling function. One can also pass the name of the calling function in who. In either case, the name of the calling function is included in

the message.

message

 ${\tt reproducibility_tools} \ \ \textit{Tools for reproducible computations}$

Description

Archiving of computations and control of the random-number generator.

Usage

```
bake(
  file,
  expr,
  seed = NULL,
  kind = NULL,
  normal.kind = NULL,
  dependson = NULL,
```

```
info = FALSE,
  timing = TRUE,
  dir = getOption("pomp_archive_dir", getwd())
)
stew(
  file,
  expr,
  seed = NULL,
 kind = NULL,
  normal.kind = NULL,
  dependson = NULL,
  info = FALSE,
  timing = TRUE,
 dir = getOption("pomp_archive_dir", getwd())
freeze(
  expr,
  seed = NULL.
 kind = NULL,
 normal.kind = NULL,
 envir = parent.frame(),
 enclos = if (is.list(envir) || is.pairlist(envir)) parent.frame() else baseenv()
)
append_data(
  data,
  file,
 overwrite = FALSE,
 dir = getOption("pomp_archive_dir", getwd())
)
```

Arguments

file

Name of the archive file in which the result will be stored or retrieved, as appropriate. For bake, this will contain a single object and hence be an RDS file (extension 'rds'); for stew, this will contain one or more named objects and hence be an RDA file (extension 'rda'); for append_data, this will be a CSV file.

expr

Expression to be evaluated.

seed, kind, normal.kind

optional. To set the state and of the RNG. The default, seed = NULL, will not change the RNG state. seed should be a single integer. See set.seed for more information.

dependson

arbitrary R object (optional). Variables on which the computation in expr depends. A hash of these objects will be archived in file, along with the results of evaluation expr. When bake or stew are called and file exists, the hash

of these objects will be compared against the archived hash; recomputation is forced when these do not match. The dependencies should be specified as unquoted symbols: use a list if there are multiple dependencies. See the note below about avoiding using 'pomp' objects as dependencies.

info logical. If TRUE, the "ingredients" of the calculation are returned as a list. In

the case of bake, this list is the "ingredients" attribute of the returned object. In the case of stew, this list is a hidden object named ".ingredients", located in the

environment within which stew was called.

timing logical. If TRUE, the time required for the computation is returned. This is

returned as the "system.time" attribute of the returned object.

dir Directory holding archive files; by default, this is the current working directory.

This can also be set using the global option pomp_archive_dir. If it does not

exist, this directory will be created (with a message).

envir the environment in which expr is to be evaluated. May also be NULL, a list, a

data frame, a pairlist or an integer as specified to sys.call.

enclos relevant when envir is a (pair)list or a data frame. Specifies the enclosure, i.e.,

where R looks for objects not found in envir. This can be NULL (interpreted as

the base package environment, baseenv()) or an environment.

data data frame

overwrite logical; if TRUE, data are written to file, replacing any existing contents. If

FALSE, the data is appended to the existing contents of file.

Details

On cooking shows, recipes requiring lengthy baking or stewing are prepared beforehand. The bake and stew functions perform analogously: an computation is performed and archived in a named file. If the function is called again and the file is present, the computation is not executed. Instead, the results are loaded from the archive. Moreover, via their optional seed argument, bake and stew can control the pseudorandom-number generator (RNG) for greater reproducibility. After the computation is finished, these functions restore the pre-existing RNG state to avoid side effects.

The freeze function doesn't save results, but does set the RNG state to the specified value and restore it after the computation is complete.

Both bake and stew first test to see whether file exists. If it does, bake reads it using readRDS and returns the resulting object. By contrast, stew loads the file using load and copies the objects it contains into the user's workspace (or the environment of the call to stew).

If file does not exist, then both bake and stew evaluate the expression expr; they differ in the results that they save. bake saves the value of the evaluated expression to file as a single object. The name of that object is not saved. By contrast, stew creates a local environment within which expr is evaluated; all objects in that environment are saved (by name) in file. bake and stew also store information about the code executed, the dependencies, and the state of the random-number generator (if the latter is controlled) in the archive file. Re-computation is triggered if any of these things change.

Value

bake returns the value of the evaluated expression expr. Other objects created in the evaluation of expr are discarded along with the temporary, local environment created for the evaluation.

The latter behavior differs from that of stew, which returns the names of the objects created during the evaluation of expr. After stew completes, these objects are copied into the environment in which stew was called.

freeze returns the value of evaluated expression expr. However, freeze evaluates expr within the parent environment, so other objects created in the evaluation of expr will therefore exist after freeze completes.

bake and stew store information about the code executed, the dependencies, and the state of the random-number generator in the archive file. In the case of bake, this is recorded in the "ingredients" attribute (attr(., "ingredients")); in the stew case, this is recorded in an object, ".ingredients", in the archive. This information is returned only if info=TRUE.

The time required for execution is also recorded. bake stores this in the "system.time" attribute of the archived R object; stew does so in a hidden variable named .system.time. The timing is obtained using system.time.

append_data returns a data frame containing the new contents of file, invisibly.

Avoid using 'pomp' objects as dependencies

Note that when a 'pomp' object is built with one or more C snippets, the resulting code is "salted" with a random element to prevent collisions in parallel computations. As a result, two such 'pomp' objects will never match perfectly, even if the codes and data used to construct them are identical. Therefore, avoid using 'pomp' objects as dependencies in bake and stew.

Compatibility with older versions

With **pomp** version 3.4.4.2, the behavior of bake and stew changed. In particular, older versions did no dependency checking, and did not check to see whether expr had changed. Accordingly, the archive files written by older versions have a format that is not compatible with the newer ones. When an archive file in the old format is encountered, it will be updated to the new format, with a warning message. **Note that this will overwrite existing archive files!** However, there will be no loss of information.

Author(s)

Aaron A. King

Examples

```
## Not run:
  bake(file="example1.rds",{
    x <- runif(1000)
    mean(x)
})

bake(file="example1.rds",{
    x <- runif(1000)
    mean(x)
})

bake(file="example1.rds",{</pre>
```

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```
a <- 3
   x <- runif(1000)
   mean(x)
 })
 a <- 5
 b <- 2
 stew(file="example2.rda",
   dependson=list(a,b),{
     x <- runif(10)</pre>
      y <- rnorm(n=10,mean=a*x+b,sd=2)
    })
 plot(x,y)
 set.seed(11)
 runif(2)
 freeze(runif(3), seed=5886730)
 runif(2)
 freeze(runif(3), seed=5886730)
 runif(2)
 set.seed(11)
 runif(2)
 runif(2)
 runif(2)
## End(Not run)
```

resample

Resample

Description

Systematic resampling.

Usage

```
systematic_resample(weights, Np = length(weights))
```

Arguments

weights numeric; vector of weights.

Np integer scalar; number of samples to draw.

Value

A vector of integers containing the indices of the resample.

ricker 129

ricker

Ricker model with Poisson observations.

Description

ricker is a 'pomp' object encoding a stochastic Ricker model with Poisson measurement error.

Usage

```
ricker(r = \exp(3.8), sigma = 0.3, phi = 10, c = 1, N_0 = 7)
```

Arguments

r	intrinsic growth rate
sigma	environmental process noise s.d.
phi	sampling rate
С	density dependence parameter
N_0	initial condition

Details

The state process is $N_{t+1} = rN_t \exp(-cN_t + e_t)$, where the e_t are i.i.d. normal random deviates with zero mean and variance σ^2 . The observed variables y_t are distributed as $Poisson(\phi N_t)$.

Value

A 'pomp' object containing the Ricker model and simulated data.

See Also

More examples provided with **pomp**: blowflies, childhood_disease_data, compartmental_models, dacca(), ebola, gompertz(), ou2(), pomp_examples, rw2(), verhulst()

Examples

```
po <- ricker()
plot(po)
coef(po)
simulate(po) |> plot()

# takes too long for R CMD check

## generate a bifurcation diagram for the Ricker map
p <- parmat(coef(ricker()),nrep=500)
p["r",] <- exp(seq(from=1.5,to=4,length=500))
trajectory(
    ricker(),
    times=seq(from=1000,to=2000,by=1),
    params=p,
    format="array"</pre>
```

rinit rinit

```
) -> x
matplot(p["r",],x["N",,],pch='.',col='black',
    xlab=expression(log(r)),ylab="N",log='x')
```

rinit rinit workhorse

Description

Samples from the initial-state distribution.

Usage

```
## S4 method for signature 'pomp'
rinit(object, ..., params = coef(object), t0 = timezero(object), nsim = 1)
```

Arguments

object	an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the pomp inference algorithms.
	additional arguments are ignored.
params	a npar x nrep matrix of parameters. Each column is treated as an independent parameter set, in correspondence with the corresponding column of x.
t0	the initial time, i.e., the time corresponding to the initial-state distribution.
nsim	optional integer; the number of initial states to simulate per column of params.

Value

rinit returns an nvar x nsim*ncol(params) matrix of state-process initial conditions when given an npar x nsim matrix of parameters, params, and an initial time t0. By default, t0 is the initial time defined when the 'pomp' object ws constructed.

See Also

```
_spec of the initial-state distribution: rinit_spec
```

```
More on pomp workhorse functions: dinit(), dmeasure(), dprior(), dprocess(), emeasure(), flow(), partrans(), pomp-package, rmeasure(), rprior(), rprocess(), skeleton(), vmeasure(), workhorses
```

rinit_spec rinit specification

Description

Specification of the initial-state distribution simulator, rinit.

Details

To fully specify the unobserved Markov state process, one must give its distribution at the zero-time (t0). One does this by furnishing a value for the rinit argument. As usual, this can be provided either as a C snippet or as an R function. In the former case, bear in mind that:

- 1. The goal of a this snippet is the construction of a state vector, i.e., the setting of the dynamical states at time t_0 .
- 2. In addition to the parameters and covariates (if any), the variable t, containing the zero-time, will be defined in the context in which the snippet is executed.
- 3. **NB:** The statenames argument plays a particularly important role when the rinit is specified using a C snippet. In particular, every state variable must be named in statenames. **Failure to follow this rule will result in undefined behavior.**

General rules for writing C snippets can be found here.

If an R function is to be used, pass

```
rinit = f
```

to pomp, where f is a function with arguments that can include the initial time t0, any of the model parameters, and any covariates. As usual, f may take additional arguments, provided these are passed along with it in the call to pomp. f must return a named numeric vector of initial states. It is of course important that the names of the states match the expectations of the other basic components.

Note that the state-process rinit can be either deterministic (as in the default) or stochastic. In the latter case, it samples from the distribution of the state process at the zero-time, t0.

Default behavior

By default, pomp assumes that the initial distribution is concentrated on a single point. In particular, any parameters in params, the names of which end in "_0" or ".0", are assumed to be initial values of states. When the state process is initialized, these are simply copied over as initial conditions. The names of the resulting state variables are obtained by dropping the suffix.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

rmeasure

See Also

rinit

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

Examples

```
## Starting with an existing pomp object
verhulst() -> po
## we add or change the initial-state simulator,
## rinit, using the 'rinit' argument in any 'pomp'
## elementary or estimation function (or in the
## 'pomp' constructor itself).
## Here, we pass the rinit specification to 'simulate'
## as an R function.
po |>
  simulate(
    rinit=function (n_0, ...) {
     c(n=rpois(n=1,lambda=n_0))
    }
 ) -> sim
## We can also pass it as a C snippet:
po |>
  simulate(
    rinit=Csnippet("n = rpois(n_0);"),
    paramnames="n_0",
    statenames="n"
  ) -> sim
```

rmeasure

rmeasure workhorse

Description

Sample from the measurement model distribution, given values of the latent states and the parameters.

rmeasure_spec 133

Usage

```
## $4 method for signature 'pomp'
rmeasure(
  object,
    ...,
  x = states(object),
  times = time(object),
  params = coef(object)
)
```

Arguments

object	an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the pomp inference algorithms.
	additional arguments are ignored.
х	an array containing states of the unobserved process. The dimensions of x are nvars x nrep x ntimes, where nvars is the number of state variables, nrep is the number of replicates, and ntimes is the length of times. One can also pass x as a named numeric vector, which is equivalent to the nrep=1, ntimes=1 case.
times	a numeric vector (length ${\tt ntimes}$) containing times. These must be in non-decreasing order.
params	a npar x nrep matrix of parameters. Each column is treated as an independent parameter set, in correspondence with the corresponding column of x.

Value

rmeasure returns a rank-3 array of dimensions nobs x nrep x ntimes, where nobs is the number of observed variables.

See Also

Specification of the measurement-model simulator: rmeasure_spec

More on **pomp** workhorse functions: dinit(), dmeasure(), dprior(), dprocess(), emeasure(), flow(), partrans(), pomp-package, rinit(), rprior(), rprocess(), skeleton(), vmeasure(), workhorses

rmeasure_spec rmeasure specification

Description

Specification of the measurement-model simulator, rmeasure.

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Details

The measurement model is the link between the data and the unobserved state process. It can be specified either by using one or both of the rmeasure and dmeasure arguments.

Suppose you have a procedure to simulate observations given the value of the latent state variables. Then you can furnish

```
rmeasure = f
```

to **pomp** algorithms, where f is a C snippet or R function that implements your procedure.

Using a C snippet is much preferred, due to its much greater computational efficiency. See Csnippet for general rules on writing C snippets.

In writing an rmeasure C snippet, bear in mind that:

- 1. The goal of such a snippet is to fill the observables with random values drawn from the measurement model distribution. Accordingly, each observable should be assigned a new value.
- 2. In addition to the states, parameters, and covariates (if any), the variable t, containing the time of the observation, will be defined in the context in which the snippet is executed.

The demos and the tutorials on the package website give examples.

It is also possible, though far less efficient, to specify rmeasure using an R function. In this case, specify the measurement model simulator by furnishing

```
rmeasure = f
```

to pomp, where f is an R function. The arguments of f should be chosen from among the state variables, parameters, covariates, and time. It must also have the argument f must return a named numeric vector of length equal to the number of observable variables.

Default behavior

The default rmeasure is undefined. It will yield missing values (NA).

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

See Also

rmeasure

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rprocess_spec, skeleton_spec, transformations, userdata, vmeasure_spec

rprior 135

Examples

```
## We start with the pre-built Ricker example:
ricker() -> po
## To change the measurement model simulator, rmeasure,
## we use the 'rmeasure' argument in any 'pomp'
## elementary or estimation function.
## Here, we pass the rmeasure specification to 'simulate'
## as an R function.
po |>
  simulate(
    rmeasure=function (N, phi, \dots) {
     c(y=rpois(n=1,lambda=phi*N))
    }
 ) -> sim
## We can also pass it as a C snippet:
po |>
 simulate(
    rmeasure=Csnippet("y = rpois(phi*N);"),
    paramnames="phi",
    statenames="N"
  ) -> sim
```

rprior

rprior workhorse

Description

Sample from the prior probability distribution.

Usage

```
## S4 method for signature 'pomp'
rprior(object, ..., params = coef(object))
```

Arguments

an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the **pomp** inference algorithms.
 additional arguments are ignored.
 a npar x nrep matrix of parameters. Each column is treated as an independent parameter set, in correspondence with the corresponding column of x.

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Value

A numeric matrix containing the required samples.

See Also

```
Specification of the prior distribution simulator: prior_spec
```

```
More on pomp workhorse functions: dinit(), dmeasure(), dprior(), dprocess(), emeasure(), flow(), partrans(), pomp-package, rinit(), rmeasure(), rprocess(), skeleton(), vmeasure(), workhorses
```

More on Bayesian methods: abc(), bsmc2(), dprior(), pmcmc(), prior_spec

rprocess

rprocess workhorse

Description

rprocess simulates the process-model portion of partially-observed Markov process.

Usage

```
## S4 method for signature 'pomp'
rprocess(
   object,
    ...,
   x0 = rinit(object),
   t0 = timezero(object),
   times = time(object),
   params = coef(object)
)
```

Arguments

object	an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the pomp inference algorithms.
	additional arguments are ignored.
x0	an nvar x nrep matrix containing the starting state of the system. Columns of x0 correspond to states; rows to components of the state vector. One independent simulation will be performed for each column. Note that in this case, params must also have nrep columns.
t0	the initial time, i.e., the time corresponding to the state in $x0$.
times	a numeric vector (length ntimes) containing times. These must be in non-decreasing order.
params	a npar x nrep matrix of parameters. Each column is treated as an independent parameter set, in correspondence with the corresponding column of $x0$.

Details

When rprocess is called, t0 is taken to be the initial time (i.e., that corresponding to x0). The values in times are the times at which the state of the simulated processes are required.

Value

rprocess returns a rank-3 array with rownames. Suppose x is the array returned. Then

```
dim(x)=c(nvars,nrep,ntimes),
```

where nvars is the number of state variables (=nrow(x0)), nrep is the number of independent realizations simulated (=ncol(x0)), and ntimes is the length of the vector times. x[,j,k] is the value of the state process in the j-th realization at time times[k]. The rownames of x will correspond to those of x0.

See Also

Specification of the process-model simulator: rprocess_spec

More on **pomp** workhorse functions: dinit(), dmeasure(), dprior(), dprocess(), emeasure(), flow(), partrans(), pomp-package, rinit(), rmeasure(), rprior(), skeleton(), vmeasure(), workhorses

rprocess_spec

rprocess specification

Description

Specification of the latent state process simulator, rprocess.

Usage

```
onestep(step.fun)
discrete_time(step.fun, delta.t = 1)
euler(step.fun, delta.t)
gillespie(rate.fun, v, hmax = Inf)
gillespie_hl(..., .pre = "", .post = "", hmax = Inf)
```

Arguments

step. fun a C snippet, an R function, or the name of a native routine in a shared-object library. This gives a procedure by which one simulates a single step of the latent

state process.

delta.t positive numerical value; for euler and discrete_time, the size of the step to

take

rate.fun a C snippet, an R function, or the name of a native routine in a shared-object

library. This gives a procedure by which one computes the event-rate of the

elementary events in the continuous-time latent Markov chain.

integer matrix; giving the stoichiometry of the continuous-time latent Markov

process. It should have dimensions nvar x nevent, where nvar is the number of state variables and nevent is the number of elementary events. v describes the changes that occur in each elementary event: it will usually comprise the values 1, -1, and 0 according to whether a state variable is incremented, decremented, or unchanged in an elementary event. The rows of v may be unnamed or named. If the rows are unnamed, they are assumed to be in the same order as the vector of state variables returned by rinit. If the rows are named, the names of the state variables returned by rinit will be matched to the rows of v to ensure a correct mapping. If any of the row names of v cannot be found among the state

variables or if any row names of v are duplicated, an error will occur.

hmax maximum time step allowed (see below)

... individual C snippets corresponding to elementary events

.pre, .post C snippets (see Details)

Discrete-time processes

If the state process evolves in discrete time, specify rprocess using the discrete_time plug-in. Specifically, provide

```
rprocess = discrete_time(step.fun = f, delta.t),
```

where f is a C snippet or R function that simulates one step of the state process. The former is the preferred option, due to its much greater computational efficiency. The goal of such a C snippet is to replace the state variables with their new random values at the end of the time interval. Accordingly, each state variable should be over-written with its new value. In addition to the states, parameters, covariates (if any), and observables, the variables t and dt, containing respectively the time at the beginning of the step and the step's duration, will be defined in the context in which the C snippet is executed. See Csnippet for general rules on writing C snippets. Examples are to be found in the tutorials on the package website.

If f is given as an R function, its arguments should come from the state variables, parameters, covariates, and time. It may also take the argument 'delta.t'; when called, the latter will be the timestep. It must also have the argument '...'. It should return a named vector of length equal to the number of state variables, representing a draw from the distribution of the state process at time t+delta.t conditional on its value at time t.

Continuous-time processes

If the state process evolves in continuous time, but you can use an Euler approximation, implement rprocess using the euler plug-in. Specify

```
rprocess = euler(step.fun = f, delta.t)
```

in this case. As before, f can be provided either as a C snippet or as an R function, the former resulting in much quicker computations. The form of f will be the same as above (in the discrete-time case).

If you have a procedure that allows you, given the value of the state process at any time, to simulate it at an arbitrary time in the future, use the onestep plug-in. To do so, specify

```
rprocess = onestep(step.fun = f).
```

Again, f can be provided either as a C snippet or as an R function, the former resulting in much quicker computations. The form of f should be as above (in the discrete-time or Euler cases).

Size of time step

The simulator plug-ins discrete_time, euler, and onestep all work by taking discrete time steps. They differ as to how this is done. Specifically,

- onestep takes a single step to go from any given time t1 to any later time t2 (t1 <= t2).
 <p>Thus, this plug-in is designed for use in situations where a closed-form solution to the process exists.
- 2. To go from t1 to t2, euler takes n steps of equal size, where

```
n = ceiling((t2-t1)/delta.t).
```

3. discrete_time assumes that the process evolves in discrete time, where the interval between successive times is delta.t. Thus, to go from t1 to t2, discrete_time takes n steps of size exactly delta.t, where

```
n = floor((t2-t1)/delta.t).
```

Exact (event-driven) simulations

If you desire exact simulation of certain continuous-time Markov chains, an implementation of Gillespie's algorithm (Gillespie 1977) is available, via the gillespie and gillespie_hl plug-ins. The former allows for the rate function to be provided as an R function or a single C snippet, while the latter provides a means of specifying the elementary events via a list of C snippets.

A high-level interface to the simulator is provided by gillespie_hl. To use it, supply

```
rprocess = gillespie_hl(..., .pre = "", .post = "", hmax = Inf)
```

to pomp. Each argument in ... corresponds to a single elementary event and should be a list containing two elements. The first should be a string or C snippet; the second should be a named integer vector. The variable rate will exist in the context of the C snippet, as will the parameter, state variables, covariates, and the time t. The C snippet should assign to the variable rate the corresponding elementary event rate.

The named integer vector specifies the changes to the state variables corresponding to the elementary event. There should be named value for each of the state variables returned by rinit. The arguments .pre and .post can be used to provide C code that will run respectively before and after the elementary-event snippets. These hooks can be useful for avoiding duplication of code that performs calculations needed to obtain several of the different event rates.

Here's how a simple birth-death model might be specified:

```
gillespie_hl(
    birth=list("rate = b*N;",c(N=1)),
    death=list("rate = m*N;",c(N=-1))
)
```

In the above, the state variable N represents the population size and parameters b, m are the birth and death rates, respectively.

To use the lower-level gillespie interface, furnish

```
rprocess = gillespie(rate.fun = f, v, hmax = Inf)
```

to pomp, where f gives the rates of the elementary events. Here, f may be an R function of the form

```
f(j, x, t, params, ...)
```

When f is called, the integer j will be the number of the elementary event (corresponding to the column the matrix v, see below), x will be a named numeric vector containing the value of the state process at time t and params is a named numeric vector containing parameters. f should return a single numerical value, representing the rate of that elementary event at that point in state space and time.

Here, the stoichiometric matrix v specifies the continuous-time Markov process in terms of its elementary events. It should have dimensions nvar x nevent, where nvar is the number of state variables and nevent is the number of elementary events. v describes the changes that occur in each elementary event: it will usually comprise the values 1, -1, and 0 according to whether a state variable is incremented, decremented, or unchanged in an elementary event. The rows of v should have names corresponding to the state variables. If any of the row names of v cannot be found among the state variables or if any row names of v are duplicated, an error will occur.

It is also possible to provide a C snippet via the rate. fun argument to gillespie. Such a snippet should assign the correct value to a rate variable depending on the value of j. The same variables will be available as for the C code provided to gillespie_hl. This lower-level interface may be preferable if it is easier to write code that calculates the correct rate based on j rather than to write a snippet for each possible value of j. For example, if the number of possible values of j is large and the rates vary according to a few simple rules, the lower-level interface may provide the easier way of specifying the model.

When the process is non-autonomous (i.e., the event rates depend explicitly on time), it can be useful to set hmax to the maximum step that will be taken. By default, the elementary event rates will be recomputed at least once per observation interval.

Default behavior

The default rprocess is undefined. It will yield missing values (NA) for all state variables.

rw2

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

See Also

rprocess

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, skeleton_spec, transformations, userdata, vmeasure_spec

rw2

Two-dimensional random-walk process

Description

rw2 constructs a 'pomp' object encoding a 2-D Gaussian random walk.

Usage

```
rw2(x1_0 = 0, x2_0 = 0, s1 = 1, s2 = 3, tau = 1, times = 1:100, t0 = 0)
```

Arguments

x1_0, x2_0	initial conditions (i.e., latent state variable values at the zero time t0)
s1, s2	random walk intensities
tau	observation error s.d.
times	observation times
t0	zero time

Details

The random-walk process is fully but noisily observed.

Value

A 'pomp' object containing simulated data.

See Also

More examples provided with **pomp**: blowflies, childhood_disease_data, compartmental_models, dacca(), ebola, gompertz(), ou2(), pomp_examples, ricker(), verhulst()

Examples

```
if (require(ggplot2)) {
  rw2() |> plot()

  rw2(s1=1,s2=1,tau=0.1) |>
    simulate(nsim=10,format="d") |>
    ggplot(aes(x=y1,y=y2,group=.id,color=.id))+
    geom_path()+
    guides(color="none")+
    theme_bw()
}
```

rw_sd

rw_sd

Description

Specifying random-walk intensities.

Usage

```
rw_sd(...)
```

Arguments

... Specification of the random-walk intensities (as standard deviations).

Details

See mif2 for details.

See Also

mif2

sannbox 143

	sannbox	Simulated annealing with box constraints.	
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Description

A straightforward implementation of simulated annealing with box constraints.

Usage

```
sannbox(par, fn, control = list(), ...)
```

Arguments

par	Initial values for the parameters to be optimized over.
fn	A function to be minimized, with first argument the vector of parameters over which minimization is to take place. It should return a scalar result.
control	A named list of control parameters. See 'Details'.
	ignored.

Details

The control argument is a list that can supply any of the following components:

trace Non-negative integer. If positive, tracing information on the progress of the optimization is produced. Higher values may produce more tracing information.

fnscale An overall scaling to be applied to the value of fn during optimization. If negative, turns the problem into a maximization problem. Optimization is performed on fn(par)/fnscale.

parscale A vector of scaling values for the parameters. Optimization is performed on par/parscale and these should be comparable in the sense that a unit change in any element produces about a unit change in the scaled value.

maxit The total number of function evaluations: there is no other stopping criterion. Defaults to 10000.

temp starting temperature for the cooling schedule. Defaults to 1.

tmax number of function evaluations at each temperature. Defaults to 10.

candidate.dist function to randomly select a new candidate parameter vector. This should be a function with three arguments, the first being the current parameter vector, the second the temperature, and the third the parameter scaling. By default, candidate.dist is

sched cooling schedule. A function of a three arguments giving the temperature as a function of iteration number and the control parameters temp and tmax. By default, sched is

```
function(k,temp,tmax) temp/log(((k-1)%/%tmax)*tmax+exp(1)).
```

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Alternatively, one can supply a numeric vector of temperatures. This must be of length at least maxit.

lower,upper optional numeric vectors. These describe the lower and upper box constraints, respectively. Each can be specified either as a single scalar (common to all parameters) or as a vector of the same length as par. By default, lower=-Inf and upper=Inf, i.e., there are no constraints.

Value

sannbox returns a list with components:

counts two-element integer vector. The first number gives the number of calls made to fn. The second number is provided for compatibility with optim and will always be NA.

convergence provided for compatibility with optim; will always be 0.

final.params last tried value of par.

final.value value of fn corresponding to final.params.

par best tried value of par.

value value of fn corresponding to par.

Author(s)

Daniel Reuman, Aaron A. King

See Also

trajectory matching, probe matching, spectrum matching, nonlinear forecasting.

saved_states

Saved states

Description

Retrieve latent state trajectories from a particle filter calculation.

Usage

```
## S4 method for signature 'pfilterd_pomp'
saved_states(object, ..., format = c("list", "data.frame"))
## S4 method for signature 'pfilterList'
saved_states(object, ..., format = c("list", "data.frame"))
```

Arguments

object result of a filtering computation

... ignored

format character; format of the returned object (see below).

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Details

When one calls pfilter with save.states="filter" or save.states="prediction", the latent state vector associated with each particle is saved. This can be extracted by calling saved_states on the 'pfilterd.pomp' object. If the filtered particles are saved, these particles are unweighted, saved after resampling using their normalized weights. If the argument save.states="prediction" was used, the particles correspond to simulations from rprocess, and their corresponding unnormalized weights are included in the output.

Value

According to the format argument, the saved states are returned either as a list or a data frame.

If format="data.frame", then the returned data frame holds the state variables and (optionally) the unnormalized log weight of each particle at each observation time. The .id variable distinguishes particles.

If format="list" and pfilter was called with save.states="unweighted" or save.states="TRUE", the returned list contains one element per observation time. Each element consists of a matrix, with one row for each state variable and one column for each particle. If pfilter was called with save.states="weighted", the list itself contains two lists: the first holds the particles as above, the second holds the corresponding unnormalized log weights. In particular, it has one element per observation time; each element is the vector of per-particle log weights.

See Also

```
More on sequential Monte Carlo methods: bsmc2(), cond_logLik(), eff_sample_size(), filter_mean(), filter_traj(), kalman, mif2(), pfilter(), pmcmc(), pred_mean(), pred_var(), wpfilter()

Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_mean(), pred_var(), spy(), states(), summary(), time(), timezero(), traces()
```

show

Show methods

Description

Display the object, according to its class.

```
## S4 method for signature 'unshowable'
show(object)

## S4 method for signature 'listie'
show(object)

## S4 method for signature 'rprocPlugin'
show(object)
```

simulate simulate

```
## S4 method for signature 'onestepRprocPlugin'
show(object)
## S4 method for signature 'discreteRprocPlugin'
show(object)
## S4 method for signature 'eulerRprocPlugin'
show(object)
## S4 method for signature 'gillespieRprocPlugin'
show(object)
## S4 method for signature 'pomp_fun'
show(object)
## S4 method for signature 'partransPlugin'
show(object)
## S4 method for signature 'covartable'
show(object)
## S4 method for signature 'skelPlugin'
show(object)
## S4 method for signature 'vectorfieldPlugin'
show(object)
## S4 method for signature 'mapPlugin'
show(object)
```

simulate

Simulations of a partially-observed Markov process

Description

simulate generates simulations of the state and measurement processes.

```
## S4 method for signature 'missing'
simulate(
  object,
  nsim = 1,
  seed = NULL,
   ...,
  times,
```

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```
t0,
  params,
  rinit,
  rprocess,
  rmeasure,
  format = c("pomps", "arrays", "data.frame"),
  include.data = FALSE,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'data.frame'
simulate(
 object,
  nsim = 1,
  seed = NULL,
  . . . ,
  times,
  t0,
  params,
  rinit,
 rprocess,
  rmeasure,
  format = c("pomps", "arrays", "data.frame"),
  include.data = FALSE,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
simulate(
 object,
  nsim = 1,
  seed = NULL,
  format = c("pomps", "arrays", "data.frame"),
  include.data = FALSE,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'objfun'
simulate(object, nsim = 1, seed = NULL, ...)
```

Arguments

object optional; if present, it should be a data frame or a 'pomp' object.

nsim The number of simulations to perform. Note that the number of replicates will

be nsim times ncol(params).

seed optional integer; if set, the pseudorandom number generator (RNG) will be ini-

tialized with seed. The RNG will be restored to its original state afterward.

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... additional arguments are passed to pomp.

times the sequence of observation times, times must indicate the column of obser-

vation times by name or index. The time vector must be numeric and non-

decreasing.

to The zero-time, i.e., the time of the initial state. This must be no later than the

time of the first observation, i.e., $t0 \le times[1]$.

params a named numeric vector or a matrix with rownames containing the parameters

at which the simulations are to be performed.

rinit simulator of the initial-state distribution. This can be furnished either as a C

snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator

to its default. For more information, see rinit specification.

rprocess simulator of the latent state process, specified using one of the rprocess plugins.

Setting rprocess=NULL removes the latent-state simulator. For more informa-

tion, see rprocess specification for the documentation on these plugins.

rmeasure simulator of the measurement model, specified either as a C snippet, an R func-

tion, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rmeasure=NULL removes the measurement model simu-

lator. For more information, see rmeasure specification.

format in which to return the results.

format = "pomps" causes the results to be returned as a single "pomp" object, if params is a vector, or a list of "pomp" objects, if params is a matrix with more than one column. Each of these will be identical to object except in that the latent states and observations will have been replaced by their simulated values.

format = "arrays" causes the results to be returned as a list of two arrays. The "states" element will contain the simulated state trajectories in a rank-3 array with dimensions nvar x (ncol(params)*nsim) x ntimes. Here, nvar is the number of state variables and ntimes the length of the argument times. The "obs" element will contain the simulated data, returned as a rank-3 array with dimensions nobs x (ncol(params)*nsim) x ntimes. Here, nobs is the number

of observables.

format = "data.frame" causes the results to be returned as a single data frame containing the time, states, and observations. An ordered factor variable, '.id',

distinguishes one simulation from another.

include.data if TRUE, the original data and covariates (if any) are included (with .id = "data").

This option is ignored unless format = "data.frame".

verbose logical; if TRUE, diagnostic messages will be printed to the console.

Value

A single "pomp" object, a "pompList" object, a named list of two arrays, or a data frame, according to the format option.

If params is a matrix, each column is treated as a distinct parameter set. In this case, if nsim=1, then simulate will return one simulation for each parameter set. If nsim>1, then simulate will yield nsim simulations for each parameter set. These will be ordered such that the first

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ncol(params) simulations represent one simulation from each of the distinct parameter sets, the second ncol(params) simulations represent a second simulation from each, and so on.

Adding column names to params can be helpful.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Author(s)

Aaron A. King

See Also

More on **pomp** elementary algorithms: elementary_algorithms, kalman, pfilter(), pomp-package, probe(), spect(), trajectory(), wpfilter()

skeleton

skeleton workhorse

Description

Evaluates the deterministic skeleton at a point or points in state space, given parameters. In the case of a discrete-time system, the skeleton is a map. In the case of a continuous-time system, the skeleton is a vectorfield. NB: skeleton just evaluates the deterministic skeleton; it does not iterate or integrate (see flow and trajectory for this).

```
## S4 method for signature 'pomp'
skeleton(
   object,
    ...,
   x = states(object),
   times = time(object),
   params = coef(object)
)
```

skeleton_spec

Arguments

object	an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the pomp inference algorithms.
	additional arguments are ignored.
х	an array containing states of the unobserved process. The dimensions of x are nvars x nrep x ntimes, where nvars is the number of state variables, nrep is the number of replicates, and ntimes is the length of times. One can also pass x as a named numeric vector, which is equivalent to the nrep=1, ntimes=1 case.
times	a numeric vector (length ntimes) containing times. These must be in non-decreasing order.
params	a npar x nrep matrix of parameters. Each column is treated as an independent parameter set, in correspondence with the corresponding column of x.

Value

skeleton returns an array of dimensions nvar x nrep x ntimes. If f is the returned matrix, f[i,j,k] is the i-th component of the deterministic skeleton at time times[k] given the state x[,j,k] and parameters params[,j].

See Also

Specification of the deterministic skeleton: skeleton_spec

```
More on pomp workhorse functions: dinit(), dmeasure(), dprior(), dprocess(), emeasure(), flow(), partrans(), pomp-package, rinit(), rmeasure(), rprior(), rprocess(), vmeasure(), workhorses
```

More on methods for deterministic process models: flow(), skeleton_spec, traj_match, trajectory()

eleton_spec skeleton specification

Description

Specification of the deterministic skeleton.

Usage

```
vectorfield(f)
map(f, delta.t = 1)
```

Arguments

f	procedure for evaluating the deterministic skeleton This can be a C snippet, an R function, or the name of a native routine in a dynamically linked library.
delta.t	positive numerical value; the size of the discrete time step corresponding to an application of the map

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Details

The skeleton is a dynamical system that expresses the central tendency of the unobserved Markov state process. As such, it is not uniquely defined, but can be both interesting in itself and useful in practice. In **pomp**, the skeleton is used by trajectory and traj_objfun.

If the state process is a discrete-time stochastic process, then the skeleton is a discrete-time map. To specify it, provide

```
skeleton = map(f, delta.t)
```

to pomp, where f implements the map and delta.t is the size of the timestep covered at one map iteration.

If the state process is a continuous-time stochastic process, then the skeleton is a vectorfield (i.e., a system of ordinary differential equations). To specify it, supply

```
skeleton = vectorfield(f)
```

to pomp, where f implements the vectorfield, i.e., the right-hand-size of the differential equations.

In either case, f can be furnished either as a C snippet (the preferred choice), or an R function. General rules for writing C snippets can be found here. In writing a skeleton C snippet, be aware that:

- 1. For each state variable, there is a corresponding component of the deterministic skeleton. The goal of such a snippet is to compute all the components.
- 2. When the skeleton is a map, the component corresponding to state variable x is named Dx and is the new value of x after one iteration of the map.
- 3. When the skeleton is a vectorfield, the component corresponding to state variable x is named Dx and is the value of dx/dt.
- 4. As with the other C snippets, all states, parameters and covariates, as well as the current time, t, will be defined in the context within which the snippet is executed.
- 5. **NB:** When the skeleton is a map, the duration of the timestep will **not** be defined in the context within which the snippet is executed. When the skeleton is a vectorfield, of course, no timestep is defined. In this regard, C snippets for the skeleton and rprocess components differ.

The tutorials on the package website give some examples.

If f is an R function, its arguments should be taken from among the state variables, parameters, covariates, and time. It must also take the argument '...'. As with the other basic components, f may take additional arguments, provided these are passed along with it in the call to pomp. The function f must return a numeric vector of the same length as the number of state variables, which contains the value of the map or vectorfield at the required point and time.

Masking of map

Other packages (most notably the **tidyverse** package **purrr**) have functions named 'map'. Beware that, if you load one of these packages after you load **pomp**, the **pomp** function map described here will be masked. You can always access the **pomp** function by calling pomp::map.

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Default behavior

The default skeleton is undefined. It will yield missing values (NA) for all state variables.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

See Also

skeleton

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, transformations, userdata, vmeasure_spec

More on methods for deterministic process models: flow(), skeleton(), traj_match, trajectory()

Examples

```
## Starting with an existing pomp object,
## e.g., the continuous-time Verhulst-Pearl model,
verhulst() -> po
## we add or change the deterministic skeleton
## using the 'skeleton' argument in any 'pomp'
## elementary or estimation function
## (or in the 'pomp' constructor itself).
## Here, we pass the skeleton specification
## to 'trajectory' as an R function.
## Since this is a continuous-time POMP, the
## skeleton is a vectorfield.
po |>
  trajectory(
    skeleton=vectorfield(
      function(r, K, n, ...) {
        c(n=r*n*(1-n/K))
      }
    ),
    format="data.frame"
  ) -> traj
## We can also pass it as a C snippet:
po |>
  traj_objfun(
```

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```
skeleton=vectorfield(Csnippet("Dn=r*n*(1-n/K);")),
    paramnames=c("r","K"),
    statenames="n"
  ) -> ofun
ofun()
## For a discrete-time POMP, the deterministic skeleton
## is a map. For example,
gompertz() -> po
po |>
 traj_objfun(
    skeleton=map(
     Csnippet("
        double dt = 1.0;
        double s = exp(-r*dt);
       DX = pow(K,(1-s))*pow(X,s);"
     ), delta.t=1
    ),
    paramnames=c("r","K"),
    statenames=c("X")
 ) -> ofun
ofun()
```

spect

Power spectrum

Description

Power spectrum computation and spectrum-matching for partially-observed Markov processes.

```
## S4 method for signature 'data.frame'
spect(
  data,
    ...,
  vars,
  kernel.width,
  nsim,
  seed = NULL,
  transform.data = identity,
  detrend = c("none", "mean", "linear", "quadratic"),
  params,
```

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```
rinit,
  rprocess,
  rmeasure,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
spect(
 data,
  . . . ,
  vars,
  kernel.width,
  nsim,
  seed = NULL,
  transform.data = identity,
 detrend = c("none", "mean", "linear", "quadratic"),
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'spectd_pomp'
spect(
  data,
  . . . ,
  vars,
 kernel.width,
 nsim,
  seed = NULL,
  transform.data,
  detrend,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'spect_match_objfun'
spect(data, ..., seed, verbose = getOption("verbose", FALSE))
## S4 method for signature 'objfun'
spect(data, ..., seed = NULL)
```

Arguments

data	either a data frame holding the time series data, or an object of class 'pomp', i.e., the output of another pomp calculation. Internally, data will be coerced to an array with storage-mode double.
• • •	additional arguments are passed to pomp. This allows one to set, unset, or modify basic model components within a call to this function.
vars	optional; names of observed variables for which the power spectrum will be

optional; names of observed variables for which the power spectrum will be computed. By default, the spectrum will be computed for all observables.

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kernel.width width parameter for the smoothing kernel used for calculating the estimate of

the spectrum.

nsim number of model simulations to be computed.

seed optional; if non-NULL, the random number generator will be initialized with this

seed for simulations. See simulate.

transform.data function; this transformation will be applied to the observables prior to estima-

tion of the spectrum, and prior to any detrending.

detrend de-trending operation to perform. Options include no detrending, and subtrac-

tion of constant, linear, and quadratic trends from the data. Detrending is applied

to each data series and to each model simulation independently.

params optional; named numeric vector of parameters. This will be coerced internally

to storage mode double.

rinit simulator of the initial-state distribution. This can be furnished either as a C

snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator

to its default. For more information, see rinit specification.

rprocess simulator of the latent state process, specified using one of the rprocess plugins.

Setting rprocess=NULL removes the latent-state simulator. For more informa-

tion, see rprocess specification for the documentation on these plugins.

rmeasure simulator of the measurement model, specified either as a C snippet, an R func-

tion, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rmeasure=NULL removes the measurement model simu-

lator. For more information, see rmeasure specification.

verbose logical; if TRUE, diagnostic messages will be printed to the console.

Details

spect estimates the power spectrum of time series data and model simulations and compares the results. It can be used to diagnose goodness of fit and/or as the basis for frequency-domain parameter estimation (spect.match).

A call to spect results in the estimation of the power spectrum for the (transformed, detrended) data and nsim model simulations. The results of these computations are stored in an object of class 'spectd_pomp'.

When spect operates on a spectrum-matching objective function (a 'spect_match_objfun' object), by default, the random-number generator seed is fixed at the value given when the objective function was constructed. Specifying NULL or an integer for seed overrides this behavior.

Value

An object of class 'spectd_pomp', which contains the model, the data, and the results of the spect computation. The following methods are available:

plot produces some diagnostic plots

summary displays a summary

logLik gives a measure of the agreement of the power spectra

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Author(s)

Daniel C. Reuman, Cai GoGwilt, Aaron A. King

References

D.C. Reuman, R.A. Desharnais, R.F. Costantino, O. Ahmad, J.E. Cohen. Power spectra reveal the influence of stochasticity on nonlinear population dynamics. *Proceedings of the National Academy of Sciences* **103**, 18860-18865, 2006. doi:10.1073/pnas.0608571103.

D.C. Reuman, R.F. Costantino, R.A. Desharnais, J.E. Cohen. Color of environmental noise affects the nonlinear dynamics of cycling, stage-structured populations. *Ecology Letters* **11**, 820-830, 2008. doi:10.1111/j.14610248.2008.01194.x.

See Also

More on methods based on summary statistics: abc(), basic_probes, nlf, probe(), probe_match, spect_match

More on **pomp** elementary algorithms: elementary_algorithms, kalman, pfilter(), pomp-package, probe(), simulate(), trajectory(), wpfilter()

spect_match

Spectrum matching

Description

Estimation of parameters by matching power spectra

```
## S4 method for signature 'data.frame'
spect_objfun(
   data,
   ...,
   est = character(0),
   weights = 1,
   fail.value = NA,
   vars,
   kernel.width,
   nsim,
```

```
seed = NULL,
  transform.data = identity,
  detrend = c("none", "mean", "linear", "quadratic"),
  params,
  rinit,
  rprocess,
  rmeasure,
 partrans,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
spect_objfun(
 data,
  est = character(0),
 weights = 1,
  fail.value = NA,
  vars,
  kernel.width,
 nsim,
  seed = NULL,
  transform.data = identity,
 detrend = c("none", "mean", "linear", "quadratic"),
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'spectd_pomp'
spect_objfun(
 data,
  ...,
  est = character(0),
 weights = 1,
  fail.value = NA,
  vars,
  kernel.width,
  nsim,
  seed = NULL,
  transform.data = identity,
 detrend,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'spect_match_objfun'
spect_objfun(
  data,
  . . . ,
  est,
```

```
weights,
fail.value,
seed = NULL,
verbose = getOption("verbose", FALSE)
```

Arguments

data either a data frame holding the time series data, or an object of class 'pomp',

i.e., the output of another pomp calculation. Internally, data will be coerced to

an array with storage-mode double.

... additional arguments are passed to pomp. This allows one to set, unset, or modify

basic model components within a call to this function.

est character vector; the names of parameters to be estimated.

weights optional numeric or function. The mismatch between model and data is mea-

sured by a weighted average of mismatch at each frequency. By default, all frequencies are weighted equally. weights can be specified either as a vector (which must have length equal to the number of frequencies) or as a function of frequency. If the latter, weights(freq) must return a nonnegative weight for

each frequency.

fail.value optional numeric scalar; if non-NA, this value is substituted for non-finite values

of the objective function. It should be a large number (i.e., bigger than any

legitimate values the objective function is likely to take).

vars optional; names of observed variables for which the power spectrum will be

computed. By default, the spectrum will be computed for all observables.

kernel.width width parameter for the smoothing kernel used for calculating the estimate of

the spectrum.

nsim the number of model simulations to be computed.

seed integer. When fitting, it is often best to fix the seed of the random-number

generator (RNG). This is accomplished by setting seed to an integer. By default,

seed = NULL, which does not alter the RNG state.

transform.data function; this transformation will be applied to the observables prior to estima-

tion of the spectrum, and prior to any detrending.

detrending operation to perform. Options include no detrending, and subtrac-

tion of constant, linear, and quadratic trends from the data. Detrending is applied

to each data series and to each model simulation independently.

params optional; named numeric vector of parameters. This will be coerced internally

to storage mode double.

rinit simulator of the initial-state distribution. This can be furnished either as a C

snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator

to its default. For more information, see rinit specification.

rprocess simulator of the latent state process, specified using one of the rprocess plugins.

Setting rprocess=NULL removes the latent-state simulator. For more informa-

tion, see rprocess specification for the documentation on these plugins.

rmeasure simulator of the measurement model, specified either as a C snippet, an R func-

tion, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rmeasure=NULL removes the measurement model simu-

lator. For more information, see rmeasure specification.

partrans optional parameter transformations, constructed using parameter_trans.

Many algorithms for parameter estimation search an unconstrained space of parameters. When working with such an algorithm and a model for which the parameters are constrained, it can be useful to transform parameters. One should supply the partrans argument via a call to parameter_trans. For more information, see parameter_trans. Setting partrans=NULL removes the parameter

transformations, i.e., sets them to the identity transformation.

verbose logical; if TRUE, diagnostic messages will be printed to the console.

Details

In spectrum matching, one attempts to minimize the discrepancy between a POMP model's predictions and data, as measured in the frequency domain by the power spectrum.

spect_objfun constructs an objective function that measures the discrepancy. It can be passed to any one of a variety of numerical optimization routines, which will adjust model parameters to minimize the discrepancies between the power spectrum of model simulations and that of the data.

Value

spect_objfun constructs a stateful objective function for spectrum matching. Specifically, spect_objfun returns an object of class 'spect_match_objfun', which is a function suitable for use in an optim-like optimizer. This function takes a single numeric-vector argument that is assumed to contain the parameters named in est, in that order. When called, it will return the (optionally weighted) L^2 distance between the data spectrum and simulated spectra. It is a stateful function: Each time it is called, it will remember the values of the parameters and the discrepancy measure.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Important Note

Since **pomp** cannot guarantee that the *final* call an optimizer makes to the function is a call *at* the optimum, it cannot guarantee that the parameters stored in the function are the optimal ones. Therefore, it is a good idea to evaluate the function on the parameters returned by the optimization routine, which will ensure that these parameters are stored.

Warning! Objective functions based on C snippets

If you use C snippets (see Csnippet), a dynamically loadable library will be built. As a rule, **pomp** functions load this library as needed and unload it when it is no longer needed. The stateful objective

functions are an exception to this rule. For efficiency, calls to the objective function do not execute pompLoad or pompUnload: rather, it is assumed that pompLoad has been called before any call to the objective function. When a stateful objective function using one or more C snippets is created, pompLoad is called internally to build and load the library: therefore, within a single R session, if one creates a stateful objective function, one can freely call that objective function and (more to the point) pass it to an optimizer that calls it freely, without needing to call pompLoad. On the other hand, if one retrieves a stored objective function from a file, or passes one to another R session, one must call pompLoad before using it. Failure to do this will typically result in a segmentation fault (i.e., it will crash the R session).

References

D.C. Reuman, R.A. Desharnais, R.F. Costantino, O. Ahmad, J.E. Cohen. Power spectra reveal the influence of stochasticity on nonlinear population dynamics. *Proceedings of the National Academy of Sciences* **103**, 18860-18865, 2006. doi:10.1073/pnas.0608571103.

D.C. Reuman, R.F. Costantino, R.A. Desharnais, J.E. Cohen. Color of environmental noise affects the nonlinear dynamics of cycling, stage-structured populations. *Ecology Letters* **11**, 820-830, 2008. doi:10.1111/j.14610248.2008.01194.x.

See Also

```
spect optim subplex nloptr
```

More on **pomp** estimation algorithms: abc(), bsmc2(), estimation_algorithms, mif2(), nlf, pmcmc(), pomp-package, probe_match

More on methods based on summary statistics: abc(), basic_probes, nlf, probe(), probe_match, spect()

More on maximization-based estimation methods: mif2(), nlf, probe_match, traj_match

Examples

```
ricker() |>
    spect_objfun(
        est=c("r", "sigma", "N_0"),
        partrans=parameter_trans(log=c("r", "sigma", "N_0")),
        paramnames=c("r", "sigma", "N_0"),
        kernel.width=3,
        nsim=100,
        seed=5069977
        ) -> f

f(log(c(20,0.3,10)))
    f |> spect() |> plot()

if (require(subplex)) {
        subplex(fn=f,par=log(c(20,0.3,10)),control=list(reltol=1e-5)) -> out
} else {
        optim(fn=f,par=log(c(20,0.3,10)),control=list(reltol=1e-5)) -> out
} f(out$par)
```

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```
f |> summary()

f |> spect() |> plot()
```

spy

Spy

Description

Peek into the inside of one of **pomp**'s objects.

Usage

```
## S4 method for signature 'pomp'
spy(object)
```

Arguments

object

the object whose structure we wish to examine

See Also

Csnippet

```
Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_mean(), pred_var(), saved_states(), states(), summary(), time(), timezero(), traces()
```

Examples

```
ricker() |> spy()
sir() |> spy()
sir2() |> spy()
```

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states

Latent states

Description

Extract the latent states from a 'pomp' object.

Usage

```
## S4 method for signature 'pomp'
states(object, vars, ..., format = c("array", "data.frame"))
## S4 method for signature 'listie'
states(object, vars, ..., format = c("array", "data.frame"))
```

Arguments

object an object of class 'pomp', or of a class extending 'pomp' vars names of variables to retrieve ... ignored format format of the returned object

See Also

```
Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_mean(), pred_var(), saved_states(), spy(), summary(), time(), timezero(), traces()
```

summary

Summary methods

Description

Display a summary of a fitted model object.

```
## S4 method for signature 'probed_pomp'
summary(object, ...)
## S4 method for signature 'spectd_pomp'
summary(object, ...)
## S4 method for signature 'objfun'
summary(object, ...)
```

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Arguments

```
object a fitted model object
... ignored or passed to the more primitive function
```

See Also

```
Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_mean(), pred_var(), saved_states(), spy(), states(), time(), timezero(), traces()
```

time

Methods to extract and manipulate the obseration times

Description

Get and set the vector of observation times.

Usage

```
## S4 method for signature 'pomp'
time(x, t0 = FALSE, ...)
## S4 replacement method for signature 'pomp'
time(object, t0 = FALSE, ...) <- value
## S4 method for signature 'listie'
time(x, t0 = FALSE, ...)</pre>
```

Arguments

```
x a 'pomp' object
t0 logical; should the zero time be included?
... ignored or passed to the more primitive function
object a 'pomp' object
value numeric vector; the new vector of times
```

Details

time(object) returns the vector of observation times. time(object, t0=TRUE) returns the vector of observation times with the zero-time t0 prepended.

time(object) <- value replaces the observation times slot (times) of object with value. time(object,t0=TRUE) <- value has the same effect, but the first element in value is taken to be the initial time. The second and subsequent elements of value are taken to be the observation times. Those data and states (if they exist) corresponding to the new times are retained.

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See Also

```
Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_mean(), pred_var(), saved_states(), spy(), states(), summary(), timezero(), traces()
```

timezero

The zero time

Description

Get and set the zero-time.

Usage

```
## S4 method for signature 'pomp'
timezero(object, ...)
## S4 replacement method for signature 'pomp'
timezero(object, ...) <- value</pre>
```

Arguments

```
object an object of class 'pomp', or of a class that extends 'pomp'
... ignored or passed to the more primitive function
value numeric; the new zero-time value
```

Value

the value of the zero time

See Also

```
Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_mean(), pred_var(), saved_states(), spy(), states(), summary(), time(), traces()
```

traces 165

Description

Retrieve the history of an iterative calculation.

Usage

```
## S4 method for signature 'mif2d_pomp'
traces(object, pars, transform = FALSE, ...)
## S4 method for signature 'mif2List'
traces(object, pars, ...)
## S4 method for signature 'abcd_pomp'
traces(object, pars, ...)
## S4 method for signature 'abcList'
traces(object, pars, ...)
## S4 method for signature 'pmcmcd_pomp'
traces(object, pars, ...)
## S4 method for signature 'pmcmcd_pomp'
traces(object, pars, ...)
```

Arguments

object an object of class extending 'pomp', the result of the application of a parameter estimation algorithm

pars names of parameters

transform logical; should the traces be transformed back onto the natural scale?

... ignored or passed to the more primitive function

Details

Note that pmcmc does not currently support parameter transformations.

Value

When object is the result of a mif2 calculation, traces(object, pars) returns the traces of the parameters named in pars. By default, the traces of all parameters are returned. If transform=TRUE, the parameters are transformed from the natural scale to the estimation scale.

When object is a 'abcd_pomp', traces(object) extracts the traces as a coda::mcmc.

When object is a 'abcList', traces(object) extracts the traces as a coda::mcmc.list.

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```
When object is a 'pmcmcd_pomp', traces(object) extracts the traces as a coda::mcmc.
When object is a 'pmcmcList', traces(object) extracts the traces as a coda::mcmc.list.
```

See Also

```
Other extraction methods: coef(), cond_logLik(), covmat(), eff_sample_size(), filter_mean(), filter_traj(), forecast(), logLik, obs(), pred_mean(), pred_var(), saved_states(), spy(), states(), summary(), time(), timezero()
```

trajectory

Trajectory of a deterministic model

Description

Compute trajectories of the deterministic skeleton of a Markov process.

```
## S4 method for signature 'missing'
trajectory(
  t0,
  times,
 params,
  skeleton,
  rinit,
 ode_control = list(),
 format = c("pomps", "array", "data.frame"),
 verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'data.frame'
trajectory(
 object,
  . . . ,
  t0,
  times,
  params,
  skeleton,
  rinit,
 ode_control = list(),
 format = c("pomps", "array", "data.frame"),
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
trajectory(
```

trajectory 167

```
object,
...,
params,
skeleton,
rinit,
ode_control = list(),
format = c("pomps", "array", "data.frame"),
verbose = getOption("verbose", FALSE)
)

## S4 method for signature 'traj_match_objfun'
trajectory(object, ..., verbose = getOption("verbose", FALSE))
```

Arguments

... additional arguments are passed to pomp.

to The zero-time, i.e., the time of the initial state. This must be no later than the

time of the first observation, i.e., $t0 \le times[1]$.

times the sequence of observation times. times must indicate the column of obser-

vation times by name or index. The time vector must be numeric and non-

decreasing.

params a named numeric vector or a matrix with rownames containing the parameters

at which the simulations are to be performed.

skeleton optional; the deterministic skeleton of the unobserved state process. Depend-

ing on whether the model operates in continuous or discrete time, this is either a vectorfield or a map. Accordingly, this is supplied using either the vectorfield or map functions. For more information, see skeleton specification. Setting

skeleton=NULL removes the deterministic skeleton.

rinit simulator of the initial-state distribution. This can be furnished either as a C

snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator

to its default. For more information, see rinit specification.

ode_control optional list; the elements of this list will be passed to ode if the skeleton is a

vectorfield, and ignored if it is a map.

format the format in which to return the results.

format = "pomps" causes the trajectories to be returned as a single 'pomp' object (if a single parameter vector has been furnished to trajectory) or as a 'pompList' object (if a matrix of parameters have been furnished). In each of these, the states slot will have been replaced by the computed trajectory. Use

states to view these.

format = "array" causes the trajectories to be returned in a rank-3 array with dimensions nvar x ncol(params) x ntimes. Here, nvar is the number of state variables and ntimes the length of the argument times. Thus if x is the returned array, x[i,j,k] is the i-th component of the state vector at time times[k] given

parameters params[,j].

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format = "data.frame" causes the results to be returned as a single data frame containing the time and states. An ordered factor variable, '.id', distinguishes the trajectories from one another.

verbose logical; if TRUE, diagnostic messages will be printed to the console. object optional; if present, it should be a data frame or a 'pomp' object.

Details

In the case of a discrete-time system, the deterministic skeleton is a map and a trajectory is obtained by iterating the map. In the case of a continuous-time system, the deterministic skeleton is a vector-field; trajectory uses the numerical solvers in **deSolve** to integrate the vectorfield.

Value

The format option controls the nature of the return value of trajectory. See above for details.

See Also

```
More on pomp elementary algorithms: elementary_algorithms, kalman, pfilter(), pomp-package, probe(), simulate(), spect(), wpfilter()
```

More on methods for deterministic process models: flow(), skeleton(), skeleton_spec, traj_match

Examples

```
## The basic components needed to compute trajectories
## of a deterministic dynamical system are
## rinit and skeleton.
## The following specifies these for a simple continuous-time
## model: dx/dt = r (1+e cos(t)) x
trajectory(
  t0 = 0, times = seq(1,30,by=0.1),
  rinit = function (x0, ...) {
    c(x = x0)
  },
  skeleton = vectorfield(
    function (r, e, t, x, ...) {
      c(x=r*(1+e*cos(t))*x)
    }
 ),
  params = c(r=1,e=3,x0=1)
) -> po
plot(po,log='y')
## In the case of a discrete-time skeleton,
## we use the 'map' function. For example,
## the following computes a trajectory from
## the dynamical system with skeleton
## x \rightarrow x \exp(r \sin(\text{omega t})).
```

```
trajectory(
  t0 = 0, times=seq(1,100),
  rinit = function (x0, ...) {
    c(x = x0)
  },
  skeleton = map(
     function (r, t, x, omega, ...) {
       c(x=x*exp(r*sin(omega*t)))
     },
     delta.t=1
  ),
  params = c(r=1, x0=1, omega=4)
) -> po
plot(po)
# takes too long for R CMD check
## generate a bifurcation diagram for the Ricker map
p <- parmat(coef(ricker()),nrep=500)</pre>
p["r",] <- exp(seq(from=1.5, to=4, length=500))</pre>
trajectory(
  ricker(),
  times=seq(from=1000, to=2000, by=1),
  params=p,
   format="array"
matplot(p["r",],x["N",,],pch='.',col='black',
  xlab=expression(log(r)),ylab="N",log='x')
```

traj_match

Trajectory matching

Description

Estimation of parameters for deterministic POMP models via trajectory matching.

```
## $4 method for signature 'data.frame'
traj_objfun(
  data,
    ...,
  est = character(0),
  fail.value = NA,
  ode_control = list(),
  params,
```

```
rinit,
  skeleton,
  dmeasure,
 partrans,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
traj_objfun(
 data,
  ...,
 est = character(0),
 fail.value = NA,
 ode_control = list(),
 verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'traj_match_objfun'
traj_objfun(
 data,
  ...,
 est,
 fail.value,
 ode_control,
 verbose = getOption("verbose", FALSE)
)
```

Arguments

data	either a data frame holding the time series data, or an object of class 'pomp', i.e., the output of another pomp calculation. Internally, data will be coerced to an array with storage-mode double.
	additional arguments will modify the model structure
est	character vector; the names of parameters to be estimated.
fail.value	optional numeric scalar; if non-NA, this value is substituted for non-finite values of the objective function. It should be a large number (i.e., bigger than any legitimate values the objective function is likely to take).
ode_control	optional list; the elements of this list will be passed to ode if the skeleton is a vectorfield, and ignored if it is a map.
params	optional; named numeric vector of parameters. This will be coerced internally to storage mode double.
rinit	simulator of the initial-state distribution. This can be furnished either as a C snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator to its default. For more information, see rinit specification.
skeleton	optional; the deterministic skeleton of the unobserved state process. Depending on whether the model operates in continuous or discrete time, this is either a

vectorfield or a map. Accordingly, this is supplied using either the vectorfield or map fractions. For more information, see skeleton specification. Setting skeleton=NULL removes the deterministic skeleton.

dmeasure evaluator of the measurement model density, specified either as a C snippet, an

R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting dmeasure=NULL removes the measurement density

evaluator. For more information, see dmeasure specification.

partrans optional parameter transformations, constructed using parameter_trans.

Many algorithms for parameter estimation search an unconstrained space of parameters. When working with such an algorithm and a model for which the parameters are constrained, it can be useful to transform parameters. One should supply the partrans argument via a call to parameter_trans. For more information, see parameter_trans. Setting partrans=NULL removes the parameter

transformations, i.e., sets them to the identity transformation.

verbose logical; if TRUE, diagnostic messages will be printed to the console.

Details

In trajectory matching, one attempts to minimize the discrepancy between a POMP model's predictions and data under the assumption that the latent state process is deterministic and all discrepancies between model and data are due to measurement error. The measurement model likelihood (dmeasure), or rather its negative, is the natural measure of the discrepancy.

Trajectory matching is a generalization of the traditional nonlinear least squares approach. In particular, if, on some scale, measurement errors are normal with constant variance, then trajectory matching is equivalent to least squares on that particular scale.

traj_objfun constructs an objective function that evaluates the likelihood function. It can be passed to any one of a variety of numerical optimization routines, which will adjust model parameters to minimize the discrepancies between the power spectrum of model simulations and that of the data.

Value

traj_objfun constructs a stateful objective function for spectrum matching. Specifically, traj_objfun returns an object of class 'traj_match_objfun', which is a function suitable for use in an optim-like optimizer. In particular, this function takes a single numeric-vector argument that is assumed to contain the parameters named in est, in that order. When called, it will return the negative log likelihood. It is a stateful function: Each time it is called, it will remember the values of the parameters and its estimate of the log likelihood.

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Important Note

Since **pomp** cannot guarantee that the *final* call an optimizer makes to the function is a call *at* the optimum, it cannot guarantee that the parameters stored in the function are the optimal ones. Therefore, it is a good idea to evaluate the function on the parameters returned by the optimization routine, which will ensure that these parameters are stored.

Warning! Objective functions based on C snippets

If you use C snippets (see Csnippet), a dynamically loadable library will be built. As a rule, pomp functions load this library as needed and unload it when it is no longer needed. The stateful objective functions are an exception to this rule. For efficiency, calls to the objective function do not execute pompLoad or pompUnload: rather, it is assumed that pompLoad has been called before any call to the objective function. When a stateful objective function using one or more C snippets is created, pompLoad is called internally to build and load the library: therefore, within a single R session, if one creates a stateful objective function, one can freely call that objective function and (more to the point) pass it to an optimizer that calls it freely, without needing to call pompLoad. On the other hand, if one retrieves a stored objective function from a file, or passes one to another R session, one must call pompLoad before using it. Failure to do this will typically result in a segmentation fault (i.e., it will crash the R session).

See Also

```
optim, subplex, nloptr
```

More on methods for deterministic process models: flow(), skeleton(), skeleton_spec, trajectory() More on maximization-based estimation methods: mif2(), nlf, probe_match, spect_match

Examples

```
ricker() |>
  traj_objfun(
    est=c("r", "sigma", "N_0"),
    partrans=parameter_trans(log=c("r", "sigma", "N_0")),
    paramnames=c("r", "sigma", "N_0")
    ) -> f

f(log(c(20,0.3,10)))

if (require(subplex)) {
    subplex(fn=f,par=log(c(20,0.3,10)),control=list(reltol=1e-5)) -> out
} else {
    optim(fn=f,par=log(c(20,0.3,10)),control=list(reltol=1e-5)) -> out
}

f(out$par)

if (require(ggplot2)) {
    f |>
        trajectory(format="data.frame") |>
```

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```
ggplot(aes(x=time,y=N))+geom_line()+theme_bw()
```

transformations

Transformations

Description

}

Some useful parameter transformations.

Usage

```
logit(p)
expit(x)
log_barycentric(X)
inv_log_barycentric(Y)
```

Arguments

р	numeric; a quantity in $[0,1]$.
Х	numeric; the log odds ratio.
Χ	numeric; a vector containing the quantities to be transformed according to the log-barycentric transformation.
Υ	numeric; a vector containing the log fractions.

Details

Parameter transformations can be used in many cases to recast constrained optimization problems as unconstrained problems. Although there are no limits to the transformations one can implement using the parameter_trans facilty, **pomp** provides a few ready-built functions to implement some very commonly useful ones.

The logit transformation takes a probability p to its log odds, $\log \frac{p}{1-p}$. It maps the unit interval [0,1] into the extended real line $[-\infty,\infty]$.

The inverse of the logit transformation is the expit transformation.

The log-barycentric transformation takes a vector $X \in \mathbb{R}^n_+$, to a vector $Y \in \mathbb{R}^n$, where

$$Y_i = \log \frac{X_i}{\sum_j X_j}.$$

The transformation is not one-to-one. However, for each c>0, it maps the simplex $\{X\in R^n_+:\sum_i X_i=c\}$ bijectively onto n-dimensional Euclidean space R^n .

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The inverse of the log-barycentric transformation is implemented as inv_log_barycentric. Note that it is not a true inverse, in the sense that it takes R^n to the *unit* simplex, $\{X \in R^n_+ : \sum_i X_i = 1\}$. Thus,

```
log_barycentric(inv_log_barycentric(Y)) == Y,
but
inv_log_barycentric(log_barycentric(X)) == X
only if sum(X) == 1.
```

See Also

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, userdata, vmeasure_spec

undefined

Undefined

Description

Check for undefined methods.

Usage

```
## S4 method for signature 'pomp_fun'
undefined(object, ...)
## S4 method for signature 'skelPlugin'
undefined(object, ...)
## S4 method for signature 'rprocPlugin'
undefined(object, ...)
```

Arguments

```
object object to test.
... currently ignored.
```

Value

Returns TRUE if the **pomp** workhorse method is undefined, FALSE if it is defined, and NA if the question is inapplicable.

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userdata	Facilities for making additional information available to basic model
	components

Description

When POMP basic components need information they can't get from parameters or covariates.

Details

It can happen that one desires to pass information to one of the POMP model *basic components* (see here for a definition of this term) outside of the standard routes (i.e., via model parameters or covariates). **pomp** provides facilities for this purpose. We refer to the objects one wishes to pass in this way as *user data*.

The following will apply to every basic model component. For the sake of definiteness, however, we'll use the rmeasure component as an example. To be even more specific, the measurement model we wish to implement is

```
y1 ~ Poisson(x1+theta), y2 ~ Poisson(x2+theta),
```

where theta is a parameter. Although it would be very easy (and indeed far preferable) to include theta among the ordinary parameters (by including it in params), we will assume here that we have some reason for not wanting to do so.

Now, we have the choice of providing rmeasure in one of three ways:

- 1. as an R function,
- 2. as a C snippet, or
- 3. as a procedure in an external, dynamically loaded library.

We'll deal with these three cases in turn.

When the basic component is specified as an R function

We can implement a simulator for the aforementioned measurement model so:

```
f <- function (t, x, params, theta, ...) {
   y <- rpois(n=2,x[c("x1","x2")]+theta)
   setNames(y,c("y1","y2"))
}</pre>
```

So far, so good, but how do we get theta to this function? We simply provide an additional argument to whichever **pomp** algorithm we are employing (e.g., simulate, pfilter, mif2, abc, etc.). For example:

```
simulate(..., rmeasure = f, userdata = list(theta = 42), ...)
```

where the . . . represent other arguments.

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When the basic component is specified via a C snippet

A C snippet implementation of the aforementioned measurement model is:

```
f <- Csnippet(r"{
  double theta = *get_userdata_double("theta");
  y1 = rpois(x1+theta); y2 = rpois(x2+theta);
}")</pre>
```

Here, the call to get_userdata_double retrieves a *pointer* to the stored value of theta. Note that, by using R string literals $(r''\{\})''$ we avoid the need to escape the quotes in the C snippet text.

It is possible to store and retrieve integer objects also, using get_userdata_int.

One must take care that one stores the user data with the appropriate storage type. For example, it is wise to wrap floating point scalars and vectors with as.double and integers with as.integer. In the present example, our call to simulate might look like

```
simulate(..., rmeasure = f, userdata = list(theta = as.double(42)), ...)
```

Since the two functions get_userdata_double and get_userdata_int return pointers, it is trivial to pass vectors of double-precision and integers.

A simpler and more elegant approach is afforded by the globals argument (see below).

When the basic component is specified via an external library

The rules are essentially the same as for C snippets. typedef declarations for the get_userdata_double and get_userdata_int are given in the 'pomp.h' header file and these two routines are registered so that they can be retrieved via a call to R_GetCCallable. See the Writing R extensions manual for more information.

Setting globals

The use of the userdata facilities incurs a run-time cost. It is often more efficient, when using C snippets, to put the needed objects directly into the C snippet library. The globals argument does this. See the example below.

See Also

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, vmeasure_spec

Examples

```
## The familiar Ricker example.
## Suppose that for some reason we wish to pass 'phi'
## via the userdata facility instead of as a parameter.
## C snippet approach:
```

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```
simulate(times=1:100,t0=0,
  userdata=list(phi=as.double(100)),
  params=c(r=3.8, sigma=0.3, N.0=7),
  rprocess=discrete_time(
    step.fun=Csnippet(r"{
    double e = (sigma > 0.0) ? rnorm(0, sigma) : 0.0;
    N = r*N*exp(-N+e);
    ),
    delta.t=1
 ),
  rmeasure=Csnippet(r"{
     double phi = *get_userdata_double("phi");
     y = rpois(phi*N);}"
  ),
  paramnames=c("r","sigma"),
  statenames="N",
  obsnames="y"
) -> rick1
## The same problem solved using 'globals':
simulate(times=1:100,t0=0,
  globals=Csnippet("static double phi = 100;"),
 params=c(r=3.8,sigma=0.3,N.0=7),
  rprocess=discrete_time(
    step.fun=Csnippet(r"{
    double e = (sigma > 0.0) ? rnorm(0, sigma) : 0.0;
    N = r*N*exp(-N+e);
    ),
    delta.t=1
 ),
  rmeasure=Csnippet("
     y = rpois(phi*N);"
  paramnames=c("r","sigma"),
  statenames="N",
  obsnames="y"
) -> rick2
## Finally, the R function approach:
simulate(times=1:100,t0=0,
  userdata=list(phi=100),
  params=c(r=3.8,sigma=0.3,N_0=7),
  rprocess=discrete_time(
    step.fun=function (r, N, sigma, ...) {
      e <- rnorm(n=1,mean=0,sd=sigma)
      c(N=r*N*exp(-N+e))
    },
    delta.t=1
 ),
  rmeasure=function (phi, N, ...) {
    c(y=rpois(n=1,lambda=phi*N))
```

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```
) -> rick3
```

verhulst

Verhulst-Pearl model

Description

The Verhulst-Pearl (logistic) model of population growth.

Usage

```
verhulst(n_0 = 10000, K = 10000, r = 0.9, sigma = 0.4, tau = 0.1, dt = 0.01)
```

Arguments

n_0	initial condition
K	carrying capacity
r	intrinsic growth rate
sigma	environmental process noise s.d.
tau	measurement error s.d.
dt	Euler timestep

Details

A stochastic version of the Verhulst-Pearl logistic model. This evolves in continuous time, according to the stochastic differential equation

$$dn_t = r n_t \left(1 - \frac{n_t}{K}\right) dt + \sigma n_t dW_t.$$

Numerically, we simulate the stochastic dynamics using an Euler approximation.

The measurements are assumed to be log-normally distributed:

$$N_t \sim \text{Lognormal} (\log n_t, \tau)$$
.

Value

A 'pomp' object containing the model and simulated data. The following basic components are included in the 'pomp' object: 'rinit', 'rprocess', 'rmeasure', 'dmeasure', and 'skeleton'.

See Also

More examples provided with **pomp**: blowflies, childhood_disease_data, compartmental_models, dacca(), ebola, gompertz(), ou2(), pomp_examples, ricker(), rw2()

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Examples

```
# takes too long for R CMD check
  verhulst() -> po
  plot(po)
  plot(simulate(po))
  pfilter(po,Np=1000) -> pf
  logLik(pf)
  spy(po)
```

vmeasure

vmeasure workhorse

Description

Return the covariance matrix of the observed variables, given values of the latent states and the parameters.

Usage

```
## S4 method for signature 'pomp'
vmeasure(
  object,
    ...,
  x = states(object),
  times = time(object),
  params = coef(object)
)
```

Arguments

object	an object of class 'pomp', or of a class that extends 'pomp'. This will typically be the output of pomp, simulate, or one of the pomp inference algorithms.
	additional arguments are ignored.
x	an array containing states of the unobserved process. The dimensions of x are nvars x nrep x ntimes, where nvars is the number of state variables, nrep is the number of replicates, and ntimes is the length of times. One can also pass x as a named numeric vector, which is equivalent to the $nrep=1$, $ntimes=1$ case.
times	a numeric vector (length $ntimes$) containing times. These must be in non-decreasing order.
params	a npar x nrep matrix of parameters. Each column is treated as an independent parameter set, in correspondence with the corresponding column of x .

Value

vmeasure returns a rank-4 array of dimensions nobs x nobs x nrep x ntimes, where nobs is the number of observed variables. If v is the returned array, v[,,j,k] contains the covariance matrix at time times[k] given the state x[,j,k].

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See Also

Specification of the measurement-model covariance matrix: vmeasure_spec

More on **pomp** workhorse functions: dinit(), dmeasure(), dprior(), dprocess(), emeasure(), flow(), partrans(), pomp-package, rinit(), rmeasure(), rprior(), rprocess(), skeleton(), workhorses

vmeasure_spec

vmeasure specification

Description

Specification of the measurement-model covariance matrix, vmeasure.

Details

The measurement model is the link between the data and the unobserved state process. Some algorithms require the conditional covariance of the measurement model, given the latent state and parameters. This is supplied using the vmeasure argument.

Suppose you have a procedure to compute this conditional covariance matrix, given the value of the latent state variables. Then you can furnish

```
vmeasure = f
```

to **pomp** algorithms, where f is a C snippet or R function that implements your procedure.

Using a C snippet is much preferred, due to its much greater computational efficiency. See Csnippet for general rules on writing C snippets.

In writing a vmeasure C snippet, bear in mind that:

- 1. The goal of such a snippet is to fill variables named V_y_z with the conditional covariances of observables y, z. Accordingly, there should be one assignment of V_y_z and one assignment of V_z_y for each pair of observables y and z.
- 2. In addition to the states, parameters, and covariates (if any), the variable t, containing the time of the observation, will be defined in the context in which the snippet is executed.

The demos and the tutorials on the package website give examples.

It is also possible, though less efficient, to specify vmeasure using an R function. In this case, specify it by furnishing

```
vmeasure = f
```

to pomp, where f is an R function. The arguments of f should be chosen from among the state variables, parameters, covariates, and time. It must also have the argument f must return a square matrix of dimension equal to the number of observable variables. The row- and columnnames of this matrix should match the names of the observable variables. The matrix should of course be symmetric.

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Default behavior

The default vmeasure is undefined. It will yield missing values (NA).

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

See Also

vmeasure

More on implementing POMP models: Csnippet, accumvars, basic_components, betabinomial, covariates, dinit_spec, dmeasure_spec, dprocess_spec, emeasure_spec, eulermultinom, parameter_trans(), pomp-package, pomp_constructor, prior_spec, rinit_spec, rmeasure_spec, rprocess_spec, skeleton_spec, transformations, userdata

window Window

Description

Restrict to a portion of a time series.

Usage

```
## S4 method for signature 'pomp'
window(x, start, end, ...)
```

Arguments

```
x a 'pomp' object or object of class extending 'pomp' start, end the left and right ends of the window, in units of time ignored
```

182 workhorses

workhorses

Workhorse functions for the **pomp** *algorithms.*

Description

These functions mediate the interface between the user's model and the package algorithms. They are low-level functions that do the work needed by the package's inference methods.

Details

```
They include

rinit which samples from the initial-state distribution,

dinit which evaluates the initial-state density,

dmeasure which evaluates the measurement model density,

rmeasure which samples from the measurement model distribution,

emeasure which computes the expectation of the observed variables conditional on the latent state,

vmeasure which computes the covariance matrix of the observed variables conditional on the latent state,

dprocess which evaluates the process model density,

rprocess which samples from the process model distribution,

dprior which evaluates the prior probability density,

rprior which samples from the prior distribution,

skeleton which evaluates the model's deterministic skeleton,

flow which iterates or integrates the deterministic skeleton to yield trajectories,

partrans which performs parameter transformations associated with the model.
```

Author(s)

Aaron A. King

See Also

```
basic model components, elementary algorithms, estimation algorithms
```

```
More on pomp workhorse functions: dinit(), dmeasure(), dprior(), dprocess(), emeasure(), flow(), partrans(), pomp-package, rinit(), rmeasure(), rprior(), rprocess(), skeleton(), vmeasure()
```

wpfilter 183

wpfilter

Weighted particle filter

Description

A sequential importance sampling (particle filter) algorithm. Unlike in pfilter, resampling is performed only when triggered by deficiency in the effective sample size.

Usage

```
## S4 method for signature 'data.frame'
wpfilter(
  data,
  . . . ,
  Nρ,
  params,
  rinit,
  rprocess,
  dmeasure,
  trigger = 1,
  target = 0.5,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'pomp'
wpfilter(
  data,
  . . . ,
 Nρ,
  trigger = 1,
  target = 0.5,
  verbose = getOption("verbose", FALSE)
)
## S4 method for signature 'wpfilterd_pomp'
wpfilter(data, ..., Np, trigger, target, verbose = getOption("verbose", FALSE))
```

Arguments

data

either a data frame holding the time series data, or an object of class 'pomp', i.e., the output of another **pomp** calculation. Internally, data will be coerced to an array with storage-mode double.

• • •

additional arguments are passed to pomp. This allows one to set, unset, or modify basic model components within a call to this function.

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Np the number of particles to use. This may be specified as a single positive integer, in which case the same number of particles will be used at each timestep. Al-

ternatively, if one wishes the number of particles to vary across timesteps, one

may specify Np either as a vector of positive integers of length

length(time(object,t0=TRUE))

or as a function taking a positive integer argument. In the latter case, Np(k) must be a single positive integer, representing the number of particles to be used at the k-th timestep: Np(0) is the number of particles to use going from timezero(object) to time(object)[1], Np(1), from timezero(object) to time(object)[1], and so on, while when T=length(time(object)), Np(T)

is the number of particles to sample at the end of the time-series.

params optional; named numeric vector of parameters. This will be coerced internally

to storage mode double.

rinit simulator of the initial-state distribution. This can be furnished either as a C

snippet, an R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting rinit=NULL sets the initial-state simulator

to its default. For more information, see rinit specification.

rprocess simulator of the latent state process, specified using one of the rprocess plugins.

Setting rprocess=NULL removes the latent-state simulator. For more informa-

tion, see rprocess specification for the documentation on these plugins.

dmeasure evaluator of the measurement model density, specified either as a C snippet, an

R function, or the name of a pre-compiled native routine available in a dynamically loaded library. Setting dmeasure=NULL removes the measurement density

evaluator. For more information, see dmeasure specification.

trigger numeric; if the effective sample size becomes smaller than trigger * Np, re-

sampling is triggered.

target numeric; target power.

verbose logical; if TRUE, diagnostic messages will be printed to the console.

Details

This function is experimental and should be considered in alpha stage. Both interface and underlying algorithms may change without warning at any time. Please explore the function and give feedback via the pomp Issues page.

Value

An object of class 'wpfilterd_pomp', which extends class 'pomp'. Information can be extracted from this object using the methods documented below.

Methods

```
logLik the estimated log likelihood
cond_logLik the estimated conditional log likelihood
eff_sample_size the (time-dependent) estimated effective sample size
as.data.frame coerce to a data frame
plot diagnostic plots
```

wquant 185

Note for Windows users

Some Windows users report problems when using C snippets in parallel computations. These appear to arise when the temporary files created during the C snippet compilation process are not handled properly by the operating system. To circumvent this problem, use the cdir and cfile options to cause the C snippets to be written to a file of your choice, thus avoiding the use of temporary files altogether.

Author(s)

Aaron A. King

References

M.S. Arulampalam, S. Maskell, N. Gordon, and T. Clapp. A tutorial on particle filters for online nonlinear, non-Gaussian Bayesian tracking. *IEEE Transactions on Signal Processing* **50**, 174–188, 2002. doi:10.1109/78.978374.

See Also

```
More on pomp elementary algorithms: elementary_algorithms, kalman, pfilter(), pomp-package, probe(), simulate(), spect(), trajectory()
```

More on sequential Monte Carlo methods: bsmc2(), cond_logLik(), eff_sample_size(), filter_mean(), filter_traj(), kalman, mif2(), pfilter(), pmcmc(), pred_mean(), pred_var(), saved_states()

More on full-information (i.e., likelihood-based) methods: bsmc2(), mif2(), pfilter(), pmcmc()

wquant

Weighted quantile function

Description

Estimate weighted quantiles.

Usage

```
wquant(
    x,
    weights = rep(1, length(x)),
    probs = c(`0%` = 0, `25%` = 0.25, `50%` = 0.5, `75%` = 0.75, `100%` = 1)
)
```

Arguments

```
x numeric; a vector of data.weights numeric; vector of weights.probs numeric; desired quantiles.
```

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Details

wquant estimates quantiles of weighted data using the estimator of Harrell & Davis (1982), with improvements recommended by Andrey Akinshin (2023).

Value

wquant returns a vector containing the estimated quantiles. If probs has names, these are inherited.

Author(s)

Aaron A. King

References

F. E. Harrell and C. E. Davis. A new distribution-free quantile estimator. *Biometrika* **69**, 635–640, 1982. doi:10.1093/biomet/69.3.635.

A. Akinshin. Weighted quantile estimators. arXiv:2304.07265, 2023. doi:10.48550/arxiv.2304.07265.

Examples

```
x <- c(1,1,1,2,2,3,3,3,3,4,5,5,6,6,6)
quantile(x)
wquant(x)
wquant(c(1,2,3,4,5,6),weights=c(3,2,4,1,2,3))
wquant(c(1,2,3,4,5),c(1,0,0,1,1))</pre>
```

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