

Package ‘ctsem’

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Type Package

Title Continuous Time Structural Equation Modelling

Version 3.10.6

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Description Hierarchical continuous (and discrete) time state space modelling, for linear and nonlinear systems measured by continuous variables, with limited support for binary data. The subject specific dynamic system is modelled as a stochastic differential equation (SDE) or difference equation, measurement models are typically multivariate normal factor models.

Linear mixed effects SDE's estimated via maximum likelihood and optimization are the default. Nonlinearities, (state dependent parameters) and random effects on all parameters are possible, using either max likelihood / max a posteriori optimization (with optional importance sampling) or Stan's Hamiltonian Monte Carlo sampling.

See <<https://github.com/cdriveraus/ctsem/raw/master/vignettes/hierarchicalmanual.pdf>>

for details. See <https://osf.io/preprints/psyarxiv/4q9ex_v2> for a detailed tutorial.

Priors may be used. For the conceptual overview of the hierarchical Bayesian linear SDE approach,

see <https://www.researchgate.net/publication/324093594_Hierarchical_Bayesian_Continuous_Time_Dynamic_Modeling>.

Exogenous inputs may also be included, for an overview of such possibilities see <https://www.researchgate.net/publication/328221807_Understanding_the_Time_Course_of_Interventions_with_Continuous_Time_Dynamic_Models> .

<<https://cdriver.netlify.app/>> contains some tutorial blog posts.

License GPL-3

Depends R (>= 4.2.0), Rcpp (>= 0.12.16)

URL <https://github.com/cdriveraus/ctsem>

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ctsem-package	<i>ctsem</i>
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Description

ctsem is an R package for continuous time structural equation modelling of panel ($N > 1$) and time series ($N = 1$) data, using either a frequentist or Bayesian approach, or middle ground forms like maximum a posteriori.

The general workflow begins by specifying a model using the `ctModel` function, in which the type of model is also specified. Then the model is fit to data using `ctStanFit`. The `ctFit` function which allows for fitting using the OpenMx / SEM form, as described in the original JSS ctsem paper, can now be found in the ctsemOMX package. The omx forms are no longer in development and for most purposes, the newer stan based forms are more robust and flexible. For examples, see `ctStanFit`. For citation info, please run `citation('ctsem')`.

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References

<https://www.jstatsoft.org/article/view/v077i05>

Driver, C. C., & Voelkle, M. C. (2018). Hierarchical Bayesian continuous time dynamic modeling. *Psychological Methods*. Advance online publication. <http://dx.doi.org/10.1037/met0000168>

Stan Development Team (2018). RStan: the R interface to Stan. R package version 2.17.3. <http://mc-stan.org>

#' @keywords internal

See Also

Useful links:

- <https://github.com/cdriveraus/ctsem>

AnomAuth

AnomAuth

Description

A dataset containing panel data assessments of individuals Anomia and Authoritarianism.

Format

data frame with 2722 rows, 14 columns. Column Y1 represents anomia, Y2 Authoritarianism, dTx the time interval for measurement occasion x.

Source

See [doi:10.1037/a0027543](https://doi.org/10.1037/a0027543) for details.

ctACF

Continuous Time Autocorrelation Function (ctACF)

Description

This function computes an approximate continuous time autocorrelation function (ACF) for data containing multiple subjects and/or variables.

Usage

```
ctACF(  
  dat,  
  varnames = "auto",  
  ccfnames = "all",  
  idcol = "id",  
  timecol = "time",  
  plot = TRUE,  
  timestep = "auto",  
  time.max = "auto",  
  nboot = 100,  
  scale = FALSE,  
  center = FALSE,  
  ...  
)
```

Arguments

<code>dat</code>	The input data in data frame or data table format.
<code>varnames</code>	Character vector of variable names in the data to compute the ACF for. 'auto' uses all columns that are not time / id.
<code>ccfnames</code>	Character vector of variable names in the data to compute cross correlation for. 'all' uses all variables in varnames, NA uses none.
<code>idcol</code>	The name of the column containing subject IDs (default is 'id').
<code>timecol</code>	The name of the column containing time values (default is 'time').
<code>plot</code>	A logical value indicating whether to create a plot (default is TRUE).
<code>timestep</code>	The time step for discretizing data. 'auto' to automatically determine the timestep based on data distribution (default is 'auto'). In this case the timestep is computed as half of the median for time intervals in the data.
<code>time.max</code>	The maximum time lag to compute the ACF (default is 10). If 'auto', is set to 10 times the 90th percentile interval in the data.
<code>nboot</code>	The number of bootstrap samples for confidence interval estimation (default is 100).
<code>scale</code>	if TRUE, scale variables based on within-subject standard deviation.
<code>center</code>	if TRUE, center variables based on within-subject mean.
<code>...</code>	additional arguments (such as <code>demean=FALSE</code>) to pass to the <code>stats::acf</code> function.

Details

This function computes the continuous time ACF by discretizing the data and then performing bootstrapped ACF calculations to estimate the confidence intervals. It can create ACF plots with confidence intervals if 'plot' is set to TRUE.

Value

If 'plot' is TRUE, the function returns a ggplot object of the ACF plot. If 'plot' is FALSE, it returns a data table with ACF estimates and confidence intervals.

See Also

[ctDiscretiseData](#)

Examples

```
data.table::setDTthreads(1) #ignore this line
# Example usage:
head(ctstantestdat)
ctACF(ctstantestdat, varnames=c('Y1'), idcol='id', timecol='time', nboot=5)
```

ctACFresiduals	<i>Calculate Continuous Time Autocorrelation Function (ACF) for Standardized Residuals of ctsem fit.</i>
----------------	--

Description

This function takes a fit object from ctsem and computes the continuous time autocorrelation function (ACF) on the standardized residuals.

Usage

```
ctACFresiduals(fit, ...)
```

Arguments

<code>fit</code>	A fitted model object generated by the ctsem package.
<code>...</code>	Additional arguments to be passed to the <code>ctACF</code> function.

Details

This function first extracts the standardized residuals from the fit object using the `ctStanKalman` function. Then, it calculates the continuous time ACF for these residuals and returns the results as a data table.

Value

A data table containing the continuous time ACF estimates for standardized residuals.

See Also

[ctStanKalman](#)

Examples

```
data.table::setDTthreads(1) #ignore this line
# Example usage:
ctACFresiduals(ctstantestfit, varnames='Y1',nboot=5)
```

ctAddSamples	<i>Sample more values from an optimized ctstanfit object</i>
--------------	--

Description

Sample more values from an optimized ctstanfit object

Usage

```
ctAddSamples(fit, nsamples, cores = 2)
```

Arguments

fit	fit object
nsamples	number of samples desired
cores	number of cores to use

Value

fit object with extra samples

Examples

```
## Not run:
newfit <- ctAddSamples(ctstantestfit, 10, 1)

## End(Not run)
```

ctCheckFit	<i>Visual model fit diagnostics for ctsem fit objects.</i>
------------	--

Description

Visual model fit diagnostics for ctsem fit objects.

Usage

```
ctCheckFit(
  fit,
  data = TRUE,
  postpred = TRUE,
  priorpred = FALSE,
  statepred = FALSE,
  residuals = FALSE,
  by = fit$ctstanmodelbase$timeName,
```



```

TIpredNames = fit$ctstanmodelbase$TIpredNames,
nsamples = 30,
covplot = FALSE,
corr = TRUE,
combinevars = NA,
fastcov = FALSE,
lagcovplot = FALSE,
aggfunc = mean,
aggregate = FALSE,
groupbysplit = FALSE,
byNA = TRUE,
lag = 0,
smooth = TRUE,
k = 4,
breaks = 4,
entropy = FALSE,
reg = FALSE,
verbose = 0,
indlines = 30
)

```

Arguments

fit	ctStanFit object.
data	Include empirical data in plots?
postpred	Include post predictive (conditional on estimated parameters and covariates) distribution data in plots?
priorpred	Include prior predictive (conditional on priors) distribution data in plots?
statepred	Include one step ahead (conditional on estimated parameters, covariates, and earlier data points) distribution data in plots?
residuals	Include one step ahead error (conditional on estimated parameters, covariates, and earlier data points) in plots?
by	Variable name to split or plot by. 'time', 'LogLik', and 'WhichObs' are also possibilities.
TIpredNames	Since time independent predictors do not change with time, by default observations after the first are ignored. For observing attrition it can be helpful to set this to NULL, or when the combinevars argument is used, specifying different names may be useful.
nsamples	Number of samples (when applicable) to include in plots.
covplot	Splits variables in the model by the 'by' argument, according to the number of breaks (breaks argument), and shows the covariance (or correlation) for the different data sources selected, as well as the differences between each pair.
corr	Turns the covplot into a correlation plot. Usually easier to make sense of visually.

combinevars	Can be a list of (possibly new) variable names, where each named element of the list contains a character vector of one or more variable names in the fit object, to combine into the one variable. By default, the mean is used, but see the aggfunc argument. The combinevars argument can also be used to ensure that only certain variables are plotted.
fastcov	Uses base R cov function for computing covariances. Not recommended with missing data.
lagcovplot	Logical. Output lagged covariance type plots?
aggfunc	Function to use for aggregation, if needed.
aggregate	If TRUE, duplicate observation types are aggregated over using aggfunc. For example, if by = 'time' and there are 8 time points per subject, but breaks = 2, there will be 4 duplicate observation types per 'row' that will be collapsed. In most cases it is helpful to not collapse.
groupbysplit	Logical. Affects variable ordering in covariance plots. Defaults to FALSE, grouping by variable, and within variable by split.
byNA	Logical. Create an extra break for when the split variable is missing?
lag	Integer vector. lag = 1 creates additional variables for plotting, prefixed by 'lag1_', containing the prior row of observations for that subject.
smooth	For bivariate plots, use a smoother for estimation?
k	Integer denoting number of knots to use in the smoothing spline.
breaks	Integer denoting number of discrete breaks to split variables by (when covariance plotting).
entropy	Still in development.
reg	Logical. Use regularisation when estimating covariance matrices? Can be necessary / faster for some problems.
verbose	Logical. If TRUE, shows optimization output when estimating covariances.
indlines	Integer number of individual subject lines to draw per data type.

Value

Nothing. Just plots.

Examples

```
ctCheckFit(ctstantestfit)
```

ctChisqTest	<i>Chi Square test wrapper for ctStanFit objects.</i>
-------------	---

Description

Chi Square test wrapper for ctStanFit objects.

Usage

```
ctChisqTest(fit1, fit2)
```

Arguments

fit1	One of the fits to be compared (better fit is assumed as base for comparison)
fit2	Second fit to be compared

Value

Numeric probability

Examples

```
df <- data.frame(id=1, time=1:length(sunspot.year), Y1=sunspot.year)

m1 <- ctModel(type='dt', LAMBDA=diag(1),MANIFESTVAR=0)
m2 <- ctModel(type='dt', LAMBDA=diag(1),MANIFESTVAR=0,DRIFT = .9)

f1 <- ctStanFit(df,m1,cores=1)
f2 <- ctStanFit(df,m2,cores=1)

ctChisqTest(f1,f2)
```

ctCollapse	<i>ctCollapse Easily collapse an array margin using a specified function.</i>
------------	---

Description

ctCollapse Easily collapse an array margin using a specified function.

Usage

```
ctCollapse(inarray, collapsemargin, collapsefunc, plyr = TRUE, ...)
```

Arguments

inarray	Input array of more than one dimension.
collapsemargin	Integers denoting which margins to collapse.
collapsefunc	function to use over the collapsing margin.
plyr	Whether to use plyr.
...	additional parameters to pass to collapsefunc.

Examples

```
testarray <- array(rnorm(900,2,1),dim=c(100,3,3))
ctCollapse(testarray,1,mean)
```

ctDeintervalise	<i>ctDeintervalise</i>
-----------------	------------------------

Description

Converts intervals in ctsem long format data to absolute time

Usage

```
ctDeintervalise(datalong, id = "id", dT = "dT", startoffset = 0)
```

Arguments

datalong	data to use, in ctsem long format (attained via function ctWideToLong)
id	character string denoting column of data containing numeric identifier for each subject.
dT	character string denoting column of data containing time interval preceding observations in that row.
startoffset	Number of units of time to offset by when converting.

ctDensity	<i>ctDensity</i>
-----------	------------------

Description

Wrapper for base R density function that removes outliers and computes 'reasonable' bandwidth and x and y limits. Used for ctsem density plots.

Usage

```
ctDensity(x, bw = "auto", plot = FALSE, ...)
```

Arguments

x numeric vector on which to compute density.
 bw either 'auto' or a numeric indicating bandwidth.
 plot logical to indicate whether or not to plot the output.
 ... Further args to density.

Examples

```
y <- ctDensity(exp(rnorm(80)))
plot(y$density, xlim=y$xlim, ylim=y$ylim)

#### Compare to base defaults:
par(mfrow=c(1,2))
y=exp(rnorm(10000))
ctdens<-ctDensity(y)
plot(ctdens$density, ylim=ctdens$ylim, xlim=ctdens$xlim)
plot(density(y))
```

ctDiscretiseData *Discretise long format continuous time (ctsem) data to specific timestep.*

Description

Extends and rounds timing information so equal intervals, according to specified timestep, are achieved. NA's are inserted in other columns as necessary, any columns specified by TDpredNames or TIpredNames have zeroes rather than NA's inserted (because some estimation routines do not tolerate NA's in covariates).

Usage

```
ctDiscretiseData(
  dlong,
  timestep,
  timecol = "time",
  idcol = "id",
  TDpredNames = NULL,
  TIpredNames = NULL
)
```

Arguments

dlong Long format data
 timestep Positive real value to discretise
 timecol Name of column containing absolute (not intervals) time information.
 idcol Name of column containing subject id variable.
 TDpredNames Vector of column names of any time dependent predictors
 TIpredNames Vector of column names of any time independent predictors

Value

long format ctsem data.

Examples

```
long <- ctDiscretiseData(dlong=ctstantestdat, timestep = .1,
  TDpredNames=c('TD1'), TIpredNames=c('TI1', 'TI2', 'TI3'))
```

ctDocs

Get documentation pdf for ctsem

Description

Get documentation pdf for ctsem

Usage

```
ctDocs()
```

Value

Nothing. Opens a pdf when run interactively.

Examples

```
ctDocs()
```

ctExample1

ctExample1

Description

Simulated example dataset for the ctsem package

Format

100 by 17 matrix containing containing ctsem wide format data. 6 measurement occasions of leisure time and happiness and 5 measurement intervals for each of 100 individuals.

ctExample1TIpred	<i>ctExample1TIpred</i>
------------------	-------------------------

Description

Simulated example dataset for the ctsem package

Format

100 by 18 matrix containing containing ctsem wide format data. 6 measurement occasions of leisure time and happiness, 1 measurement of number of friends, and 5 measurement intervals for each of 100 individuals.

ctExample2	<i>ctExample2</i>
------------	-------------------

Description

Simulated example dataset for the ctsem package

Format

100 by 18 matrix containing containing ctsem wide format data. 8 measurement occasions of leisure time and happiness, 7 measurement occasions of a money intervention dummy, and 7 measurement intervals for each of 50 individuals.

ctExample2level	<i>ctExample2level</i>
-----------------	------------------------

Description

Simulated example dataset for the ctsem package

Format

100 by 18 matrix containing ctsem wide format data. 8 measurement occasions of leisure time and happiness, 7 measurement occasions of a money intervention dummy, and 7 measurement intervals for each of 50 individuals.

`ctExample3`*ctExample3*

Description

Simulated example dataset for the ctsem package

Format

1 by 399 matrix containing containing ctsem wide format data. 100 observations of variables Y1 and Y2 and 199 measurement intervals, for 1 subject.

`ctExample4`*ctExample4*

Description

Simulated example dataset for the ctsem package

Format

20 by 79 matrix containing 20 observations of variables Y1, Y2, Y3, and 19 measurement intervals dTx, for each of 20 individuals.

`ctExtract`*Extract samples from a ctStanFit object*

Description

Extract samples from a ctStanFit object

Usage

```
ctExtract(  
  object,  
  subjectMatrices = FALSE,  
  cores = 2,  
  nsamples = "all",  
  subjects = "all"  
)
```


Arguments

object	ctStanFit object, samples may be from Stan's HMC, or the importance sampling approach of ctsem.
subjectMatrices	Calculate subject specific system matrices?
cores	Only used if subjectMatrices = TRUE . For faster computation use more cores.
nsamples	either 'all' or an integer denoting number of random samples to extract.
subjects	either 'all', or an integer vector denoting subjects to extract.

Value

Array of posterior samples.

Examples

```
e = ctExtract(ctstantestfit)
```

ctFit

ctFit function placeholder

Description

For the original ctsem OpenMx functionality, the package ctsemOMX should be loaded.

Usage

```
ctFit(...)
```

Arguments

... arguments to pass to ctFit, if ctsemOMX is loaded.

Value

message or fit object.

Examples

```
data(AnomAuth)
AnomAuthmodel <- ctModel(LAMBDA = matrix(c(1, 0, 0, 1), nrow = 2, ncol = 2),
  Tpoints = 5, n.latent = 2, n.manifest = 2, MANIFESTVAR=diag(0, 2), TRAITVAR = NULL)
AnomAuthfit <- ctFit(AnomAuth, AnomAuthmodel)
```

 ctFitAuto

ctFitAuto

Description

Fit a ctStan model with automatic parameter selection

Usage

```
ctFitAuto(
  m,
  dat,
  DRIFT = TRUE,
  DIFFUSION = TRUE,
  fast = FALSE,
  initialRestrictions = NA,
  individuals = FALSE,
  groupFreeThreshold = 0.5,
  cores = 2,
  ...
)
```

Arguments

<code>m</code>	ctStan model object without time independent predictors.
<code>dat</code>	Data in long format
<code>DRIFT</code>	Logical, if TRUE, off diagonal drift parameters in the model are tested for inclusion
<code>DIFFUSION</code>	Logical, if TRUE, off diagonal diffusion parameters in the model are tested for inclusion
<code>fast</code>	Logical, if TRUE, do not compute uncertainty hessian / samples in individual level models.
<code>initialRestrictions</code>	Alternative to the DRIFT / DIFFUSION arguments – specify explicitly which parameters should be fixed initially, vector of integers based on the <code>\$setup\$matsetup</code> element of the ctStanFit object, which gives the parameter numbers. Primarily for internal use.
<code>individuals</code>	Logical, if TRUE, fit individual level models and determine a group model based on the <code>groupFreeThreshold</code> argument.
<code>groupFreeThreshold</code>	Numeric, threshold for group model structure – if a parameter improves fit in this proportion of individuals or greater, it is freed for all individuals.
<code>cores</code>	Number of CPU cores to use
<code>...</code>	Additional arguments passed to ctStanFit

Details

This function is used to automatically select parameters in a ctStan model. Any specified DRIFT / DIFFUSION matrix off diagonals are only included if they significantly improve the likelihood, based on an estimated likelihood ratio test (relying on the Hessian).

Value

A ctStan fit object

Examples

```
## Not run:
testmodel <- ctstantestfit$ctstanmodelbase
testmodel$pars$TI1_effect <- NULL
testmodel$n.TIpred <- 0
testmodel$TIpredNames <- NULL
testfit <- ctFitAuto(testmodel, dat = ctstantestdat, DRIFT = TRUE, DIFFUSION = TRUE)
summary(testfit)

## End(Not run)
```

ctFitAutoGroupModel *ctFitAutoGroupModel*

Description

Fit a ctStan model with automatic parameter selection for multiple subjects

Usage

```
ctFitAutoGroupModel(
  m,
  dat,
  cores,
  DRIFT = TRUE,
  DIFFUSION = TRUE,
  groupFreeThreshold = 0.5,
  ...
)
```

Arguments

m	ctStan model object without time independent predictors.
dat	Data in long format
cores	Number of CPU cores to use
DRIFT	Logical, if TRUE, off diagonal drift parameters in the model are tested for inclusion

```

DIFFUSION      Logical, if TRUE, off diagonal diffusion parameters in the model are tested for
                inclusion
groupFreeThreshold
                Numeric, threshold for group free parameter selection. Default is .5
...
                Additional arguments passed to ctStanFit

```

Details

This function is used to automatically select parameters in a ctStan model. Any specified DRIFT / DIFFUSION matrix off diagonals are only included if they significantly improve the likelihood, based on an estimated likelihood ratio test (relying on the Hessian). Subjects are fit one by one, and a group model is determined based on the groupFreeThreshold parameter – when the proportion of subjects with a parameter free is above this threshold, the parameter is freed in the group model.

Value

A list containing a list of ctStan fit objects for each subject, and a group model

Examples

```

## Not run:
testmodel <- ctstantestfit$ctstanmodelbase
testmodel$pars$TI1_effect <- NULL
testmodel$n.TIpred <- 0
testmodel$TIpredNames <- NULL
testfit <- ctFitAutoGroupModel(testmodel,
dat = ctstantestdat, cores=2, DRIFT = TRUE, DIFFUSION = TRUE)
ctModelLatex(testfit$groupModel)
lapply(testfit$fits,function(x) print(ctStanContinuousPars(x)$DRIFT))

## End(Not run)

```

ctFitCovCheck	<i>Visual model-fit diagnostics for ctsem fits, optionally split by the mean of a chosen observed variable.</i>
---------------	---

Description

Visual model-fit diagnostics for ctsem fits, optionally split by the mean of a chosen observed variable.

Usage

```
ctFitCovCheck(fit, cor = TRUE, plot = TRUE, splitby = NULL)
```

Arguments

fit	A 'ctStanFit' object.
cor	Logical; if 'TRUE' correlations are analysed instead of covariances.
plot	Logical; if 'TRUE' (default) a 'ggplot2' object (or list of plots) is returned. If 'FALSE', the raw 'data.table' with diagnostics is returned.
splitby	Optional character string giving the variable name on which to split subjects.

Value

Either a plot/list of plots (default, given by 'ctFitCovCheckPlot') or a 'data.table'.

Examples

```
ctFitCovCheck(ctstantestfit, cor = TRUE, plot = TRUE)

gg=ctFitCovCheck(ctstantestfit, cor = FALSE, plot = TRUE, splitby = "TI1")
print(gg[[1]]+
  ggplot2::ggtitle("Covariance Check for Y1 with all other manifest variables, split by TI1"))
```

ctFitCovCheckPlot *ctFitCovCheckPlot*

Description

Plot the results of ctFitCovCheck.

Usage

```
ctFitCovCheckPlot(x, maxlag = 10, vars = NA, splitvar = NA, cor = FALSE, ...)
```

Arguments

x	Output from ctFitCovCheck.
maxlag	Maximum lag to plot.
vars	Optional character vector of variable names to plot. If 'NA', all variables are plotted.
splitvar	Optional character string specifying a variable to split the plot by.
cor	Logical; if 'TRUE', plot correlations instead of covariances. (label change)
...	not used.

Value

ggplot object.

Examples

```
## Not run:
ctFitCovCheckPlot(ctFitCovCheck(ctstantestfit,cor=TRUE,plot=FALSE),maxlag=3)

## End(Not run)
```

ctFitMultiModel

Fit and summarise a list of ctsem models

Description

Fit and summarise a list of ctsem models

Usage

```
ctFitMultiModel(
  mlist,
  datalong,
  prefix = "",
  type = "ct",
  cores = 2,
  summaryOutput = TRUE,
  saveFits = TRUE,
  summaryArgs = list(),
  cv = FALSE,
  cvArgs = list(),
  ...
)
```

Arguments

mlist	Named list of models
dalong	ctsem long format data
prefix	prefix for output files.
type	'ct' for continuous time or 'dt' for discrete time
cores	number of cpu cores to use
summaryOutput	Generate summary output into ctSummary folder? Large datasets can take some time.
saveFits	Save fit objects to working directory?
summaryArgs	Additional arguments for ctSummarise.
cv	Perform k-fold cross validation?
cvArgs	Additional arguments for ctLOO function used for cross validation.
...	Additional arguments for ctStanFit.

Value

List containing a named list of model fits (`$fits`), and a compare object (`$compare`)

Examples

```
## Not run:
sunspots<-data.frame(id=1,
  time=do.call(seq,(lapply(attributes(sunspot.year)$tsp,function(x) x))),
  sunspots=sunspot.year)

ssmodel1 <- ctModel(type='omx', manifestNames='sunspots', Tpoints=3,
  latentNames=c('ss_level', 'ss_velocity'),
  LAMBDA=matrix(c( 1, 'ma1| log(1+(exp(param)))' ), nrow=1, ncol=2),
  DRIFT=matrix(c(0, 'a21 | -log(1+exp(param))', 1, 'a22'), nrow=2, ncol=2),
  MANIFESTMEANS=matrix(c('m1|param * 10 + 44'), nrow=1, ncol=1),
  MANIFESTVAR=diag(0,1), #As per original spec
  CINT=matrix(c(0, 0), nrow=2, ncol=1),
  DIFFUSION=matrix(c(0, 0, 0, "diffusion"), ncol=2, nrow=2))

ssmodel2 <- ssmodel1
ssmodel2$LAMBDA[2] <- 0

fits<-ctFitMultiModel(list(m1=ssmodel1,m2=ssmodel2),datalong = sunspots,
  summaryOutput = FALSE, saveFits = FALSE, cores=1, cv=TRUE, cvArgs=list(folds=5))
print(fits$compare)

## End(Not run)
```

ctGenerate

ctGenerate

Description

This function generates data according to the specified ctsem model object.

Usage

```
ctGenerate(
  ctmodelobj,
  n.subjects = 100,
  burnin = 0,
  dtmean = 1,
  logdtsd = 0,
  dtmat = NA,
  wide = FALSE
)
```

Arguments

ctmodelobj	ctsem model object from ctModel .
n.subjects	Number of subjects to output.
burnin	Number of initial time points to discard (to simulate stationary data)
dtmean	Positive numeric. Average time interval (delta T) to use.
logdtsd	Numeric. Standard deviation for variability of the time interval.
dtmat	Either NA, or numeric matrix of n.subjects rows and Tpoints-1 columns, containing positive numeric values for all time intervals between measurements. If not NA, dtmean and logdtsd are ignored.
wide	Logical. Output in wide format?

Details

Covariance related matrices are treated as Cholesky factors. TRAITDPREDCOV and TIPREDCOV matrices are not accounted for, at present. The first 1:n.TDpred rows and columns of TD-PREDCOV are used for generating tdpreds at each time point.

Examples

```
#generate data for 2 process model, each process measured by noisy indicator,
#stable individual differences in process levels.

generatingModel<-ctModel(Tpoints=8,n.latent=2,n.TDpred=0,n.TIpred=0,n.manifest=2,
  MANIFESTVAR=diag(.1,2),
  LAMBDA=diag(1,2),
  DRIFT=matrix(c(-.2,-.05,-.1,-.1),nrow=2),
  TRAITVAR=matrix(c(.5,.2,0,.8),nrow=2),
  DIFFUSION=matrix(c(1,.2,0,4),2),
  CINT=matrix(c(1,0),nrow=2),
  T0MEANS=matrix(0,ncol=1,nrow=2),
  T0VAR=diag(1,2))

data<-ctGenerate(generatingModel,n.subjects=15,burnin=10)
```

ctIntervalise

Converts absolute times to intervals for wide format ctsem panel data

Description

Converts absolute times to intervals for wide format ctsem panel data

Usage

```

ctIntervalise(
  datawide,
  Tpoints,
  n.manifest,
  n.TDpred = 0,
  n.TIpred = 0,
  imputedefs = F,
  manifestNames = "auto",
  TDpredNames = "auto",
  TIpredNames = "auto",
  digits = 5,
  mininterval = 0.001,
  individualRelativeTime = TRUE,
  startoffset = 0
)

```

Arguments

datawide	Wide format data, containing absolute time measurements, to convert to interval time scale. See ctLongToWide to easily convert long format data.
Tpoints	Maximum number of discrete time points (waves of data, or measurement occasions) for an individual in the input data structure.
n.manifest	number of manifest variables per time point in the data.
n.TDpred	number of time dependent predictors in the data structure.
n.TIpred	number of time independent predictors in the data structure.
imputedefs	if TRUE, impute time intervals based on the measurement occasion (i.e. column) they are in, if FALSE (default), set related observations to NA. FALSE is recommended unless you are certain that the imputed value (mean of the relevant time column) is appropriate. Noise and bias in estimates will result if wrongly set to TRUE.
manifestNames	vector of character strings giving variable names of manifest indicator variables (without <code>_Tx</code> suffix for measurement occasion).
TDpredNames	vector of character strings giving variable names of time dependent predictor variables (without <code>_Tx</code> suffix for measurement occasion).
TIpredNames	vector of character strings giving variable names of time independent predictor variables.
digits	How many digits to round to for interval calculations.
mininterval	set to lower than any possible observed measurement interval, but above 0 - this is used for filling NA values where necessary and has no impact on estimates when set in the correct range. (If all observed intervals are greater than 1, <code>mininterval=1</code> may be a good choice)
individualRelativeTime	if TRUE (default), the first measurement for each individual is assumed to be taken at time 0, and all other times are adjusted accordingly. If FALSE, new

columns for an initial wave are created, consisting only of observations which occurred at the earliest observation time of the entire sample.

`startoffset` if 0 (default) uses earliest observation as start time. If greater than 0, all first observations are NA, with distance of `startoffset` to first recorded observation.

Details

Time column must be numeric!

Examples

```
wideexample <- ctLongToWide(datalong = ctstantestdat, id = "id",
  time = "time", manifestNames = c("Y1", "Y2"),
  TDpredNames = "TD1", TIpredNames = c("TI1", "TI2", "TI3"))

#Then convert the absolute times to intervals, using the Tpoints reported from the prior step.
wide <- ctIntervalise(datawide = wideexample, Tpoints = 10, n.manifest = 2,
  n.TDpred = 1, n.TIpred = 3, manifestNames = c("Y1", "Y2"),
  TDpredNames = "TD1", TIpredNames = c("TI1", "TI2", "TI3") )

print(wide)
```

ctKalman

ctKalman

Description

Outputs predicted, updated, and smoothed estimates of manifest indicators and latent states, with covariances, for specific subjects from data fit with `ctStanFit`, based on either the mode (if optimized) or mean (if sampled) of parameter distribution.

Usage

```
ctKalman(
  fit,
  timerange = "asdata",
  timestep = "auto",
  subjects = fit$standata$idmap[1, 1],
  removeObs = FALSE,
  plot = FALSE,
  standardisederrors = FALSE,
  realid = TRUE,
  ...
)
```

Arguments

fit	fit object as generated by <code>ctStanFit</code> .
timerange	Either 'asdata' to just use the observed data range, or a numeric vector of length 2 denoting start and end of time range, allowing for estimates outside the range of observed data. Ranges smaller than the observed data are ignored.
timestep	Either 'asdata' to just use the observed data (which also requires 'asdata' for timerange) or a positive numeric value indicating the time step to use for interpolating values. Lower values give a more accurate / smooth representation, but take a little more time to calculate.
subjects	vector of integers denoting which subjects (from 1 to N) to plot predictions for.
removeObs	Logical or integer. If TRUE, observations (but not covariates) are set to NA, so only expectations based on parameters and covariates are returned. If a positive integer N, every N observations are retained while others are set NA for computing model expectations – useful for observing prediction performance forward further in time than one observation.
plot	Logical. If TRUE, plots output instead of returning it. See <code>plot.ctKalmanDF</code> (Stan based fit) for the possible arguments.
standardisederrors	if TRUE, also include standardised error output (based on covariance per time point).
realid	use original (not necessarily integer sequence) subject id's? Otherwise use integers 1:N.
...	additional arguments to pass to <code>plot.ctKalmanDF</code> .

Value

Returns a list containing matrix objects `etaprior`, `etaupd`, `etasmooth`, `y`, `yprior`, `yupd`, `ysmooth`, `prederror`, `time`, `loglik`, with values for each time point in each row. `eta` refers to latent states and `y` to manifest indicators - `y` itself is thus just the input data. Covariance matrices `etapriorcov`, `etaupdcov`, `etasmoothcov`, `ypriorcov`, `yupdcov`, `ysmoothcov`, are returned in a row * column * time array. Some outputs are unavailable for `ctStan` fits at present. If `plot=TRUE`, nothing is returned but a plot is generated.

Examples

```
#Basic
ctKalman(ctstantestfit, timerange=c(0,60), plot=TRUE)

#Multiple subjects, y and yprior, showing plot arguments
plot1<-ctKalman(ctstantestfit, timerange=c(0,60), timestep=.1, plot=TRUE,
  subjects=2:3,
  kalmanvec=c('y','yprior'),
  errorvec=c(NA,'ypriorcov')) #'auto' would also have achieved this

#modify plot as per normal with ggplot
print(plot1+ggplot2::coord_cartesian(xlim=c(0,10)))
```

```
#or generate custom plot from scratch:#'
k=ctKalman(ctstantestfit, timerange=c(0,60), timestep=.1, subjects=2:3)
library(ggplot2)
ggplot(k[k$Element %in% 'yprior',],
  aes(x=Time, y=value, colour=Subject, linetype=Row)) +
  geom_line() +
  theme_bw()
```

ctLongToWide	<i>ctLongToWide Restructures time series / panel data from long format to wide format for ctsem analysis</i>
--------------	--

Description

ctLongToWide Restructures time series / panel data from long format to wide format for ctsem analysis

Usage

```
ctLongToWide(
  datalong,
  id,
  time,
  manifestNames,
  TDpredNames = NULL,
  TIpredNames = NULL
)
```

Arguments

datalong	dataset in long format, including subject/id column, observation time (or change in observation time, with 0 for first observation) column, indicator (manifest / observed) variables, any time dependent predictors, and any time independent predictors.
id	character string giving column name of the subject/id column
time	character string giving column name of the time columnn
manifestNames	vector of character strings giving column names of manifest indicator variables
TDpredNames	vector of character strings giving column names of time dependent predictor variables
TIpredNames	vector of character strings giving column names of time independent predictor variables

Details

Time column must be numeric

See Also[ctIntervalise](#)**Examples**

```

wideexample <- ctLongToWide(datalong = ctstantedat, id = "id",
  time = "time", manifestNames = c("Y1", "Y2"),
  TDpredNames = "TD1", TIpredNames = c("TI1", "TI2", "TI3"))

#Then convert the absolute times to intervals, using the Tpoints reported from the prior step.
wide <- ctIntervalise(datawide = wideexample, Tpoints = 10, n.manifest = 2,
  n.TDpred = 1, n.TIpred = 3, manifestNames = c("Y1", "Y2"),
  TDpredNames = "TD1", TIpredNames = c("TI1", "TI2", "TI3") )

print(wide)

```

ctLOO

*K fold cross validation for ctStanFit objects***Description**

K fold cross validation for ctStanFit objects

Usage

```

ctLOO(
  fit,
  folds = 10,
  cores = 2,
  parallelFolds = FALSE,
  tol = 1e-05,
  subjectwise = ifelse(length(unique(fit$standata$subject)) >= folds, TRUE, FALSE),
  keepfirststobs = FALSE,
  leaveOutN = NA,
  refit = TRUE,
  casewiseApproximation = FALSE
)

```

Arguments

fit	ctStanfit object
folds	Number of cross validation splits to use – 10 folds implies that the model is re-fit 10 times, each time to a data set with 1/10 of the observations randomly removed.
cores	Number of processor cores to use.
parallelFolds	compute folds in parallel or use cores to finish single folds faster. parallelFolds will use folds times as much memory.

tol	tolerance for optimisation of refitted samples, can generally be more relaxed than the tolerance used for fitting initially.
subjectwise	drop random subjects instead of data rows?
keepfirstobs	do not drop first observation (more stable estimates)
leaveOutN	if a positive integer is given, the folds argument is ignored and instead the folds are calculated by leaving out every Nth row from the data when fitting. Leaving 2 out would result in 3 folds (starting at rows 1,2,3), each containing one third of the data.
refit	if FALSE, do not optimise parameters for the new data set, just compute the likelihoods etc from the original parameters
casewiseApproximation	if TRUE, use a bootstrapped gradient contributions approach to approximate the cross validation parameters – much faster but less reliable.

Value

list

Examples

```
ctLOO(ctstantestfit)
```

 ctModel

Define a ctsem model

Description

This function is used to specify a continuous time structural equation model, which can then be fit to data with function `ctStanFit`.

Usage

```
ctModel(
  LAMBDA,
  type = "omx",
  n.manifest = "auto",
  n.latent = "auto",
  Tpoints = NULL,
  manifestNames = "auto",
  manifesttype = rep(0, nrow(LAMBDA)),
  latentNames = "auto",
  id = "id",
  time = "time",
  silent = FALSE,
  T0VAR = "auto",
```

```

T0MEANS = "auto",
MANIFESTMEANS = "auto",
MANIFESTVAR = "diag",
DRIFT = "auto",
CINT = 0,
DIFFUSION = "auto",
n.TDpred = "auto",
TDpredNames = "auto",
n.TIpred = "auto",
TIpredNames = "auto",
tipredDefault = TRUE,
TRAITVAR = NULL,
T0TRAITEFFECT = NULL,
MANIFESTTRAITVAR = NULL,
TDPREDMEANS = "auto",
TDPREDEFFECT = "auto",
T0TDPREDCOV = "auto",
TDPREDVAR = "auto",
TRAITTDPREDCOV = "auto",
TDTIPREDCOV = "auto",
TIPREDMEANS = "auto",
TIPREDEFFECT = "auto",
T0TIPREDEFFECT = "auto",
TIPREDVAR = "auto",
PARS = NULL,
startValues = NULL
)

```

Arguments

LAMBDA	n.manifest*n.latent loading matrix relating latent to manifest variables, with latent processes 1:n.latent along the columns, and manifest variables 1:n.manifest in the rows.
type	character string. If 'omx' (default) configures model for maximum likelihood fitting with ctFit, using OpenMx. If 'ct' or 'dt' configures either continuous ('ct') or discrete ('dt') time model for Bayesian fitting with ctStanFit, using Stan.
n.manifest	Number of manifest indicators per individual at each measurement occasion / time point. Manifest variables are included as the first element of the wide data matrix, with all the 1:n.manifest manifest variables at time 1 followed by those of time 2, and so on.
n.latent	Number of latent processes.
Tpoints	For type='omx' only. Number of time points, or measurement occasions, in the data. This will generally be the maximum number of time points for a single individual, but may be one extra if sample relative time intervals are used, see ctIntervalise.
manifestNames	n.manifest length vector of manifest variable names as they appear in the data

	structure, without any <code>_Tx</code> time point suffix that may be present in wide data. Defaults to Y1, Y2, etc.
manifesttype	n.manifest length vector of manifest variable types, defaults to 0 for continuous vars, 1 for binary vars is also possible.
latentNames	n.latent length vector of latent variable names (used for naming parameters, defaults to eta1, eta2, etc).
id	character string denoting column name containing subject identification variables. id data may be of any form, though will be coerced internally to an integer sequence rising from 1.
time	character string denoting column name containing timing data. Timing data must be numeric.
silent	Suppress all output to console.
T0VAR	lower triangular n.latent*n.latent cholesky matrix of latent process initial variance / covariance. "auto" freely estimates all parameters.
T0MEANS	n.latent*1 matrix of latent process means at first time point, T0. "auto" freely estimates all parameters.
MANIFESTMEANS	n.manifest*1 matrix of manifest intercept parameters. "auto" frees all parameters.
MANIFESTVAR	lower triangular n.manifest*n.manifest cholesky matrix of variance / covariance between manifests at each measurement occasion (i.e. measurement error / residual). "auto" freely estimates variance parameters, and fixes covariances between manifests to 0. "free" frees all values, including covariances.
DRIFT	n.latent*n.latent DRIFT matrix of continuous auto and cross effects, relating the processes over time. "auto" freely estimates all parameters.
CINT	n.latent * 1 matrix of latent process intercepts, allowing for non 0 asymptotic levels of the latent processes. Generally only necessary for additional trends and more complex dynamics.
DIFFUSION	lower triangular n.latent*n.latent cholesky matrix of diffusion process variance and covariance (latent error / dynamic innovation). "auto" freely estimates all parameters.
n.TDpred	Number of time dependent predictor variables in the dataset.
TDpredNames	n.TDpred length vector of time dependent predictor variable names, as they appear in the data structure, without any <code>_Tx</code> time point suffix that may appear in wide data. Default names are TD1, TD2, etc.
n.TIpred	Number of time independent predictors. Each TIpredictor is inserted at the right of the data matrix, after the time intervals.
TIpredNames	n.TIpred length vector of time independent predictor variable names, as they appear in the data structure. Default names are TI1, TI2, etc.
tipredDefault	Logical. TRUE sets any parameters with unspecified time independent predictor effects to have effects estimated, FALSE fixes the effect to zero unless individually specified.
TRAITVAR	For type='omx' only. Either NULL, if no trait / unobserved heterogeneity effect, or lower triangular n.latent*n.latent cholesky matrix of trait variance / covariance across subjects. "auto" freely estimates all parameters.

T0TRAITEFFECT	For type='omx' only. Either NULL, if no trait / individual heterogeneity effect, or lower triangular $n.latent * n.latent$ cholesky matrix of initial trait variance / covariance. "auto" freely estimates all parameters, if the TRAITVAR matrix is specified.
MANIFESTTRAITVAR	For type='omx' only. Either NULL (default) if no trait variance / individual heterogeneity in the level of the manifest indicators, otherwise a lower triangular $n.manifest * n.manifest$ variance / covariance matrix. Set to "auto" to include and free all parameters - but identification problems will arise if TRAITVAR is also set.
TDPREDMEANS	For type='omx' only. $(n.TDpred * (Tpoints - 1))$ rows * 1 column matrix of time dependent predictor means. If 'auto', the means are freely estimated. Otherwise, the means for the Tpoints observations of your first time dependent predictor are followed by those of TDpred 2, and so on.
TDPREDEFFECT	$n.latent * n.TDpred$ matrix of effects from time dependent predictors to latent processes. Effects from 1:n.TDpred columns TDpredictors go to 1:n.latent rows of latent processes. "auto" freely estimates all parameters.
T0TDPREDCOV	For type='omx' only. $n.latent$ rows * $(Tpoints * n.TDpred)$ columns covariance matrix between latents at T0 and time dependent predictors. Default of "auto" restricts covariance to 0, which is consistent with covariance to other time points. To freely estimate parameters, specify either 'free', or the desired matrix.
TDPREDVAR	For type='omx' only. lower triangular $(n.TDpred * Tpoints)$ rows * $(n.TDpred * Tpoints)$ columns variance / covariance cholesky matrix for time dependent predictors. "auto" (default) freely estimates all parameters.
TRAITTDPREDCOV	For type='omx' only. $n.latent$ rows * $(n.TDpred * Tpoints)$ columns covariance matrix of latent traits and time dependent predictors. Defaults to zeroes, assuming predictors are independent of subjects baseline levels. When predictors depend on the subjects, this should instead be set to 'free' or manually specified. The Tpoints columns of the first predictor are followed by those of the second and so on. Covariances with the trait variance of latent process 1 are specified in row 1, process 2 in row 2, etc. "auto" (default) sets this matrix to zeroes, (if both traits and time dependent predictors exist, otherwise this matrix is set to NULL, and ignored in any case).
TDTIPREDCOV	For type='omx' only. $(n.TDpred * Tpoints)$ rows * $n.TIpred$ columns covariance matrix between time dependent and time independent predictors. "auto" (default) freely estimates all parameters.
TIPREDMEANS	For type='omx' only. $n.TIpred * 1$ matrix of time independent predictor means. If 'auto', the means are freely estimated.
TIPREDEFFECT	For type='omx' only. $n.latent * n.TIpred$ effect matrix of time independent predictors on latent processes. "auto" freely estimates all parameters and generates starting values. TIPREDEFFECT parameters for type='stan' are estimated by default on all subject level parameters, to restrict this, manually edit the model object after creation.
T0TIPREDEFFECT	For type='omx' only. $n.latent * n.TIpred$ effect matrix of time independent predictors on latents at T0. "auto" freely estimates all parameters, though note that under the default setting of stationary for ctFit, this matrix is ignored as the effects are determined based on the overall process parameters.

TIPREDVAR	For type='omx' only. lower triangular $n.TIpred * n.TIpred$ Cholesky decomposed covariance matrix for all time independent predictors. "auto" (default) freely estimates all parameters.
PARS	for types 'ct' and 'dt' only. May be of any structure, only needed to contain extra parameters for certain non-linear models.
startValues	For type='omx' only. A named vector, where the names of each value must match a parameter in the specified model, and the value sets the starting value for that parameter during optimization. If not set, random starting values representing relatively stable processes with small effects and covariances are generated by ctFit. Better starting values may improve model fit speed and the chance of an appropriate model fit.

Examples

```
### Frequentist example:
### impulse and level change time dependent predictor
### example from Driver, Oud, Voelkle (2015)
data('ctExample2')
tdpredmodel <- ctModel(n.manifest = 2, n.latent = 3, n.TDpred = 1,
  Tpoints = 8, manifestNames = c('LeisureTime', 'Happiness'),
  TDpredNames = 'MoneyInt',
  latentNames = c('LeisureTime', 'Happiness', 'MoneyIntLatent'),
  LAMBDA = matrix(c(1,0, 0,1, 0,0), ncol = 3), TRAITVAR = "auto")

tdpredmodel$TRAITVAR[3, ] <- 0
tdpredmodel$TRAITVAR[, 3] <- 0
tdpredmodel$DIFFUSION[, 3] <- 0
tdpredmodel$DIFFUSION[3, ] <- 0
tdpredmodel$T0VAR[3, ] <- 0
tdpredmodel$T0VAR[, 3] <- 0
tdpredmodel$CINT[3] <- 0
tdpredmodel$T0MEANS[3] <- 0
tdpredmodel$TDPREDEFFECT[3, ] <- 1
tdpredmodel$DRIFT[3, ] <- 0

### Bayesian example:
model <- ctModel(type='ct',
  n.latent=2, latentNames=c('eta1','eta2'),
  n.manifest=2, manifestNames=c('Y1','Y2'),
  n.TDpred=1, TDpredNames='TD1',
  n.TIpred=3, TIpredNames=c('TI1','TI2','TI3'),
  LAMBDA=diag(2))
```

Description

Performs a coverage check analysis by generating data from a model, fitting it multiple times with different fit arguments, and plotting the results.

Usage

```
ctModelCoverage_check(
  initialData,
  fitting_model,
  niter,
  fit_args,
  cores = 10,
  plot_every = max(c(10, cores))
)
```

Arguments

initialData	An initial dataset to fit to determine 'true' parameters for further generation.
fitting_model	A ctModel object used for fitting the data
niter	Number of iterations to run
fit_args	Named list of fit argument sets to test (e.g., list(boot = list(optimcontrol = list(bootstrapUncertainty = TRUE)), hess = list(optimcontrol = list(bootstrapUncertainty = FALSE))))
cores	Number of cores to use for parallel processing
plot_every	Print plots every n iterations (default = 10)

Value

A list containing the results data.table and final plots

ctModelHigherOrder *Raise the order of a ctsem model object of type 'omx'.*

Description

Raise the order of a ctsem model object of type 'omx'.

Usage

```
ctModelHigherOrder(
  ctm,
  indices,
  diffusion = TRUE,
  crosseffects = FALSE,
  cint = FALSE,
  explosive = FALSE
)
```

Arguments

ctm	ctModel
indices	Vector of integers, which latents to raise the order of.
diffusion	Shift the diffusion parameters / values to the higher order?
crosseffects	Shift cross coupling parameters of the DRIFT matrix to the higher order?
cint	shift continuous intercepts to higher order?
explosive	Allow explosive (non equilibrium returning) processes?

Value

extended ctModel

Examples

```
om <- ctModel(LAMBDA=diag(1,2),DRIFT=0,
  MANIFESTMEANS=0,type='omx',Tpoints=4)

om <- ctModelHigherOrder(om,1:2)
print(om$DRIFT)

m <- ctStanModel(om)
print(m$pars)
```

ctModelLatex	<i>Generate and optionally compile latex equation of subject level ctsem model.</i>
--------------	---

Description

Generate and optionally compile latex equation of subject level ctsem model.

Usage

```
ctModelLatex(
  x,
  matrixnames = TRUE,
  digits = 3,
  linearise = class(x) %in% "ctStanFit",
  textsize = "normalsize",
  folder = tempdir(),
  filename = paste0("ctsemTex", as.numeric(Sys.time())),
  tex = TRUE,
  equationonly = FALSE,
  compile = TRUE,
  open = TRUE,
  includeNote = TRUE,
```

```

    minimal = FALSE,
    savepng = FALSE
  )

```

Arguments

x	ctsem model object or ctStanFit object.
matrixnames	Logical. If TRUE, includes ctsem matrix names such as DRIFT and DIFFUSION under the matrices.
digits	Precision of decimals for numeric values.
linearise	Logical. Show the linearised normal approximation for subject parameters and covariate effects, or the raw parameters?
textsize	Standard latex text sizes – tiny scriptsize footnotesize small normalsize large Large LARGE huge Huge. Useful if output overflows page.
folder	Character string specifying folder to save to, defaults to temporary directory, use "." for working directory.
filename	filename, without suffix, to output .tex and .pdf files too.
tex	Save .tex file? Otherwise latex is simply returned within R as a string.
equationonly	Logical. If TRUE, output is only the latex relevant to the equation, not a compileable document.
compile	Compile to .pdf? (Depends on tex = TRUE)
open	Open after compiling? (Depends on compile = TRUE)
includeNote	Include text describing matrix transformations and subject notation? triangular matrices (which results in a covariance or Cholesky matrix) is shown – the latter is a more direct representation of the model, while the former is often simpler to convey.
minimal	if TRUE, outputs reduced form version displaying matrix dimensions and equation structure only.
savepng	Logical. If TRUE, renders the equation as a png file instead of a pdf, viewing the png in RStudio viewer when available.

Value

character string of latex code. Side effects include saving a .tex, .pdf, and displaying the pdf.

Examples

```

ctmodel <- ctModel(type='ct',
  n.latent=2, n.manifest=1,
  manifestNames='sunspots',
  latentNames=c('ss_level', 'ss_velocity'),
  LAMBDA=matrix(c( 1, 'ma1' ), nrow=1, ncol=2),
  DRIFT=matrix(c(0, 1, 'a21', 'a22'), nrow=2, ncol=2, byrow=TRUE),
  MANIFESTMEANS=matrix(c('m1'), nrow=1, ncol=1),
  CINT=matrix(c(0, 0), nrow=2, ncol=1),
  DIFFUSION=matrix(c(

```

```

0, 0,
0, "diffusion"), ncol=2, nrow=2, byrow=TRUE))

l=ctModelLatex(ctmodel, compile=FALSE, open=FALSE)
cat(l)

```

ctPlotArray

Plots three dimensional y values for quantile plots

Description

1st margin of \$Y sets line values, 2nd sets variables, 3rd quantiles.

Usage

```

ctPlotArray(
  input,
  grid = FALSE,
  add = FALSE,
  colvec = "auto",
  lwdvec = "auto",
  ltyvec = "auto",
  typevec = "auto",
  plotcontrol = list(ylab = "Array values", xaxs = "i"),
  legend = TRUE,
  legendcontrol = list(),
  polygon = TRUE,
  polygonalpha = 0.1,
  polygoncontrol = list(steps = 25)
)

```

Arguments

input	list containing 3 dimensional array to use for Y values, \$y and vector of corresponding x values \$x.
grid	Logical. Plot with a grid?
add	Logical. If TRUE, plotting is overlayed on current plot, without creating new plot.
colvec	color vector of same length as 2nd margin.
lwdvec	lwd vector of same length as 2nd margin.
ltyvec	lty vector of same length as 2nd margin.
typevec	type vector of same length as 2nd margin.
plotcontrol	list of arguments to pass to plot.
legend	Logical. Draw a legend?
legendcontrol	list of arguments to pass to legend .

polygon Logical. Draw the uncertainty polygon?
 polygonalpha Numeric, multiplier for alpha (transparency) of the uncertainty polygon.
 polygoncontrol list of arguments to pass to [ctPoly](#)

Value

Nothing. Generates plots.

Examples

```

#'
input<-ctStanTIPreEffects(ctstancodefit, plot=FALSE, whichpars='CINT',
  nsamples=10, nsubjects=10)

ctPlotArray(input=input)

```

ctPoly *Plots uncertainty bands with shading*

Description

Plots uncertainty bands with shading

Usage

```
ctPoly(x, y, ylow, yhigh, steps = 20, ...)
```

Arguments

x x values
 y y values
 ylow lower limits of y
 yhigh upper limits of y
 steps number of polygons to overlay - higher integers lead to smoother changes in transparency between y and yhigh / ylow.
 ... arguments to pass to `polygon()`

Value

Nothing. Adds a polygon to existing plot.

Examples

```

plot(0:100, sqrt(0:100), type='l')
ctPoly(x=0:100, y=sqrt(0:100),
  yhigh=sqrt(0:100) - runif(101),
  ylow=sqrt(0:100) + runif(101),
  col=adjustcolor('red', alpha.f=.1))

```

ctPostPredData	<i>Create a data.table to compare data generated from a ctsem fit with the original data.</i>
----------------	---

Description

This function allows for easy comparison of data generated from a fitted ctsem model with the original data used to fit the model. It provides options to include residuals in the comparison.

Usage

```
ctPostPredData(fit, residuals = F)
```

Arguments

fit	A fitted ctsem model.
residuals	If set to TRUE, includes residuals in the comparison.

Value

A data table containing the comparison between generated and original data.

See Also

Other ctsem functions for model fitting and analysis.

Examples

```
data_comparison <- ctPostPredData(ctstantestfit)
```

ctPostPredPlots	<i>Create diagnostic plots to assess the goodness-of-fit for a ctsem model.</i>
-----------------	---

Description

This function generates a set of diagnostic plots to assess the goodness-of-fit for a fitted ctsem model.

Usage

```
ctPostPredPlots(fit)
```

Arguments

fit	A fitted ctsem model.
-----	-----------------------

Details

The function calculates various statistics and creates visualizations to evaluate how well the generated data matches the original data used to fit the model. The plots included are as follows: - A scatter plot comparing observed values and the median of generated data. - A plot showing the proportion of observed data outside the 95 - A density plot of the proportion of observed data greater than the generated data. - A time series plot of the proportion of observed data greater than generated data.

Value

a list of ggplot2 plots.

Examples

```
print(ctPostPredPlots(ctstantestfit))
```

 ctPredictTIP

ctPredictTIP

Description

Outputs the estimated effect of time independent predictors (covariate moderators) on the expected observations.

Usage

```
ctPredictTIP(
  sf,
  tipreds = "all",
  subject = 1,
  timestep = "auto",
  doDynamics = TRUE,
  plot = TRUE,
  quantiles = c(0.16, 0.5, 0.84),
  discreteTimeQuantiles = c(0.025, 0.5, 0.975),
  showUncertainty = TRUE,
  TIPvalues = NA
)
```

Arguments

sf A fitted ctStanFit object from the ctsem package.

tipreds A character vector specifying which time independent predictors to use. Default is 'all', which uses all time independent predictors in the model.

subject	An integer value specifying the internal ctsem subject ID (mapping visible under myfit\$setup\$tidmap) for which predictions are made. This is relevant only when time dependent predictors are also included in the model.
timestep	A numeric value specifying the time step for predictions. Default is 'auto', which tries to automatically determine an appropriate time step.
doDynamics	A logical value indicating whether to plot the effects of time independent predictors on the dynamics of the system. Default is TRUE. Can be problematic for systems with many dimensions.
plot	A logical value indicating whether to ggplot the results instead of returning a data.frame of predictions. Default is TRUE.
quantiles	A numeric vector specifying the quantiles of the time independent predictors to plot. Default is 1SD either side and the median, c(.32,.5,.68).
discreteTimeQuantiles	a numeric vector of length 3 specifying the quantiles of the discrete time points to plot, when showUncertainty is TRUE.
showUncertainty	A logical value indicating whether to plot the uncertainty of the predictions. Default is TRUE.
TIPvalues	An nvalue * nTIpred numeric matrix specifying the fixed values for each time independent predictor effect to plot. Default is NA, which instead relies on the quantiles specified in the quantiles argument.

Details

This function estimates the effects of covariate moderators on the expected process and observations for a specified subject in a dynamic system. The covariate moderators are defined at the specified quantiles, and their effects on the trajectory are plotted or returned as a data frame.

Value

If plot is TRUE, a list of ggplot objects showing the estimated effects of covariate moderators. Otherwise, a data frame with the predictions.

Examples

```
# Example usage:
ctPredictTIP(ctstantestfit, tipreds='all', doDynamics=FALSE, plot=TRUE)
```

ctResiduals

Extract Standardized Residuals from a ctsem Fit

Description

This function takes a fit object from the ctsem package and extracts the standardized residuals.

Usage

```
ctResiduals(fit)
```

Arguments

`fit` A fitted model object generated by the ctsem package.

Details

This function uses the [ctStanKalman](#) function to calculate the standardized residuals and then extracts and formats them as a data table. The standardized residuals represent the differences between the observed and predicted values, divided by the standard errors of the observations.

Value

A data table containing the standardized residuals for each subject and time point.

See Also

[ctStanKalman](#)

Examples

```
data.table::setDTthreads(1) #ignore this line
# Example usage:
residuals <- ctResiduals(ctstantestfit)
```

ctStanContinuousPars *ctStanContinuousPars*

Description

Returns the continuous time parameter matrices of a ctStanFit fit object

Usage

```
ctStanContinuousPars(  
  fit,  
  calcfunc = quantile,  
  calcfuncargs = list(probs = 0.5),  
  timeinterval = 1  
)
```

Arguments

fit	fit object from ctStanFit
calcfunc	Function to apply over samples, must return a single value. By default the median over all samples is returned using the quantile function, but one might also be interested in the mean or sd , for instance.
calcfuncargs	A list of additional parameters to pass to calcfunc. For instance, with the default of calcfunc = quantile, the probs argument is needed to ensure only a single value is returned.
timeinterval	time interval for discrete time parameter matrix computation.

Examples

```
#posterior median over all subjects (also reflects mean of unconstrained pars)
ctStanContinuousPars(ctstantestfit)
```

ctStanDiscretePars *ctStanDiscretePars*

Description

Calculate model implied regressions for a sequence of time intervals (if ct) or steps (if dt) based on a ctStanFit object, for specified subjects. Wrap with print() when used inside for loops!

Usage

```
ctStanDiscretePars(
  ctstanfitobj,
  subjects = "popmean",
  times = seq(from = 0, to = 10, by = 0.1),
  nsamples = 200,
  observational = FALSE,
  standardise = FALSE,
  cov = FALSE,
  plot = FALSE,
  cores = 2,
  ...
)
```

Arguments

ctstanfitobj	model fit from ctStanFit
subjects	Either 'popmean', to use the population mean parameter, or a vector of integers denoting which subjects.

times	Numeric vector of positive values, discrete time parameters will be calculated for each. If the fit object is a discrete time model, these should be positive integers.
nsamples	Number of samples from the stanfit to use for plotting. Higher values will increase smoothness / accuracy, at cost of plotting speed. Values greater than the total number of samples will be set to total samples.
observational	Logical. If TRUE, outputs expected change in processes *conditional on observing* a 1 unit change in each – this change is correlated according to the DIFFUSION matrix. If FALSE, outputs expected regression values – also interpretable as an independent 1 unit change on each process, giving the expected response under a 1 unit experimental impulse.
standardise	Logical. If TRUE, output is standardised according to expected total within subject variance, given by the asymDIFFUSIONcov matrix.
cov	Logical. If TRUE, covariances are returned instead of regression coefficients.
plot	Logical. If TRUE, ggplots output using <code>ctStanDiscreteParsPlot</code> instead of returning output.
cores	Number of cpu cores to use for computing subject matrices. If subject matrices were saved during fitting, not used.
...	additional plotting arguments to control <code>ctStanDiscreteParsPlot</code>

Details

If `plot=TRUE`, the function will return a `ggplot2` object (and hence needs to be printed if intended to display within a loop). This can be modified by the various `ggplot2` functions, or displayed using `print(x)`.

Examples

```
data.table::setDTthreads(1) #ignore this line
ctStanDiscretePars(ctstantestfit, times=seq(.5, 4, .1),
  plot=TRUE, indices='CR')

#modify plot
require(ggplot2)
g=ctStanDiscretePars(ctstantestfit, times=seq(.5, 4, .1),
  plot=TRUE, indices='CR')
g= g+ labs(title='Cross effects')
print(g)
```

ctStanDiscreteParsPlot

ctStanDiscreteParsPlot

Description

Plots the output from `ctStanDiscretePars`, for model implied regression strengths at specified times for continuous time models fit with `ctStanFit`.

Usage

```
ctStanDiscreteParsPlot(
  x,
  indices = "all",
  quantiles = c(0.025, 0.5, 0.975),
  latentNames = "auto",
  ylab = "Coefficient",
  xlab = "Time interval",
  ylim = NA,
  facets = NA,
  splitSubjects = TRUE,
  colour = "Effect",
  title = "auto",
  polygonalpha = 0.1,
  ggcode = FALSE
)
```

Arguments

x	list object returned from <code>ctStanDiscretePars</code> .
indices	Either a string specifying type of plot to create, or an n by 2 matrix specifying which indices of the output matrix to plot. 'AR' specifies all diagonals, for discrete time autoregression parameters. 'CR' specifies all off-diagonals, for discrete time cross regression parameters. 'all' plots all AR and CR effects at once.
quantiles	numeric vector of length 3, with values between 0 and 1, specifying which quantiles to plot. The default plots 95% credible intervals and the posterior median at 50%.
latentNames	Vector of character strings denoting names for the latent variables. 'auto' just uses eta1 eta2 etc.
ylab	y label.
xlab	x label.
ylim	Custom ylim.
facets	May be 'Subject' or 'Effect'.
splitSubjects	if TRUE, subjects are plotted separately, if FALSE they are combined.
colour	Character string denoting how colour varies. 'Effect' or 'Subject'.
title	Character string. 'auto' generates automatically, NULL can be used to disable title.
polygonalpha	Numeric between 0 and 1 to multiply the alpha of the fill.
ggcode	if TRUE, returns a list containing the data.table to plot, and a character string that can be evaluated (with the necessary arguments such as ylab etc filled in). For modifying plots.

Value

A ggplot2 object. This can be modified by the various ggplot2 functions, or displayed using `print(x)`.

Examples

```
data.table::setDTthreads(1) #ignore this line
x <- ctStanDiscretePars(ctstantestfit)
ctStanDiscreteParsPlot(x, indices='CR')

#to modify plot:
g <- ctStanDiscreteParsPlot(x, indices='CR') +
  ggplot2::labs(title='My ggplot modification')
print(g)
```

ctStanFit

ctStanFit

Description

Fits a ctsem model specified via [ctModel](#) with type either 'ct' or 'dt'.

Usage

```
ctStanFit(
  datalong,
  ctstanmodel,
  stanmodeltext = NA,
  iter = 1000,
  intoverstates = TRUE,
  binomial = FALSE,
  fit = TRUE,
  intoverpop = "auto",
  sameInitialTimes = FALSE,
  stationary = FALSE,
  plot = FALSE,
  derrind = NA,
  optimize = TRUE,
  optimcontrol = list(),
  nlcontrol = list(),
  nopriors = NA,
  priors = FALSE,
  chains = 2,
  cores = ifelse(optimize, getOption("mc.cores", 2L), "maxneeded"),
  inits = NULL,
  compileArgs = list(),
  forcerecompile = FALSE,
  saveCompile = TRUE,
  savescores = FALSE,
  savesubjectmatrices = FALSE,
  saveComplexPars = FALSE,
```

```

  gendata = FALSE,
  control = list(),
  verbose = 0,
  vb = FALSE,
  ...
)

```

Arguments

<code>datalong</code>	long format data containing columns for subject id (numeric values, 1 to max subjects), manifest variables, any time dependent (i.e. varying within subject) predictors, and any time independent (not varying within subject) predictors.
<code>ctstanmodel</code>	model object as generated by <code>ctModel</code> with <code>type='ct'</code> or <code>'dt'</code> , for continuous or discrete time models respectively.
<code>stanmodeltext</code>	already specified Stan model character string, generally leave NA unless modifying Stan model directly. (Possible after modification of output from fitting with argument <code>fit=FALSE</code>)
<code>iter</code>	used when <code>optimize=FALSE</code> . number of iterations, half of which will be devoted to warmup by default when sampling.
<code>intoverstates</code>	logical indicating whether or not to integrate over latent states using a Kalman filter. Generally recommended to set TRUE unless using non-gaussian measurement model.
<code>binomial</code>	Deprecated. Logical indicating the use of binary rather than Gaussian data, as with IRT analyses. This now sets <code>intoverstates = FALSE</code> and the <code>manifesttype</code> of every indicator to 1, for binary.
<code>fit</code>	If TRUE, fit specified model using Stan, if FALSE, return stan model object without fitting.
<code>intoverpop</code>	if 'auto', set to TRUE if optimizing and FALSE if using hmc. if TRUE, integrates over population distribution of parameters rather than full sampling. Allows for optimization of non-linearities and random effects, via state expansion.
<code>sameInitialTimes</code>	if TRUE, include an empty observation for every subject that has no observation at the earliest observation time of the dataset. This ensures that the TOMEANS occurs for every subject at the same time, rather than just at the earliest observation for that subject. Important when modelling trends over time, age, etc.
<code>stationary</code>	Logical. If TRUE, T0VAR and T0MEANS input matrices are ignored, the parameters are instead fixed to long run expectations. More control over this can be achieved by instead setting parameter names of T0MEANS and T0VAR matrices in the input model to 'stationary', for elements that should be fixed to stationarity.
<code>plot</code>	if TRUE, for sampling, a Shiny program is launched upon fitting to interactively plot samples. May struggle with many (e.g., > 5000) parameters. For optimizing, various optimization details are plotted – in development.
<code>derrind</code>	deprecated, latents involved in dynamic error calculations are determined automatically now.

optimize	if TRUE, use <code>stanoptimis</code> function for maximum a posteriori / importance sampling estimates, otherwise use the HMC sampler from Stan, which is (much) slower, but generally more robust for complex individual differences.
optimcontrol	list of parameters sent to <code>stanoptimis</code> governing optimization / importance sampling.
nlcontrol	List of non-linear control parameters. <code>maxtimestep</code> must be a positive numeric, specifying the largest time span covered by the numerical integration. The large default ensures that for each observation time interval, only a single step of exponential integration is used. When <code>maxtimestep</code> is smaller than the observation time interval, the integration is nested within an Euler like loop. Smaller values may offer greater accuracy, but are slower and not always necessary. Given the exponential integration, linear model elements are fit exactly with only a single step.
nopriors	deprecated, use <code>priors</code> argument. logical. If TRUE, any priors are disabled – sometimes desirable for optimization.
priors	if TRUE, priors are included in computations, otherwise specified priors are ignored.
chains	used when <code>optimize=FALSE</code> . Number of chains to sample, during HMC or post-optimization importance sampling. Unless the <code>cores</code> argument is also set, the number of chains determines the number of cpu cores used, up to the maximum available minus one. Irrelevant when <code>optimize=TRUE</code> .
cores	number of cpu cores to use. Either 'maxneeded' to use as many as available minus one, up to the number of chains, or a positive integer. If <code>optimize=TRUE</code> , more cores are generally faster.
inits	either character string 'optimize, NULL, or vector of (unconstrained) parameter start values, as returned by the <code>rstan</code> function <code>rstan::unconstrain_pars</code> , or the parameter values found in a <code>ctsem</code> fit object <code>myfit\$stanfit\$rawest</code> (or <code>\$rawposterior</code>) for instance.
compileArgs	List of arguments to pass to <code>stan_model</code> for compilation of the Stan model.
forcerecompile	logical. For development purposes. If TRUE, stan model is recompiled, regardless of apparent need for compilation.
saveCompile	if TRUE and compilation is needed / requested, writes the stan model to the parent frame as <code>ctsem.compiled</code> (unless that object already exists and is not from <code>ctsem</code>), to avoid unnecessary recompilation.
savescores	Logical. If TRUE, output from the Kalman filter is saved in output. For datasets with many variables or time points, will increase file size substantially.
savesubjectmatrices	Logical. If TRUE, subject specific matrices are saved – only relevant when either time dependent predictors or individual differences are used. Can increase memory usage dramatically in large models, and can be computed after fitting using <code>ctExtract</code> or <code>ctStanSubjectPars</code> .
saveComplexPars	Logical. If TRUE, also save rowwise output of any complex parameters specified, i.e. combinations of parameters, functions and states.

gendata	Logical – If TRUE, uses provided data for only covariates and a time and missingness structure, and generates random data according to the specified model / priors. Generated data is in the \$Ygen subobject after running extract on the fit object. For datasets with many manifest variables or time points, file size may be large. To generate data based on the posterior of a fitted model, see ctStanGenerateFromFit .
control	Used when optimize=FALSE. List of arguments sent to stan control argument, regarding warmup / sampling behaviour. Unless specified, values used are: list(adapt_delta = .8, adapt_window=2, max_treedepth=10, adapt_init_buffer=2, stepsize = .001)
verbose	Integer from 0 to 2. Higher values print more information during model fit – for debugging.
vb	Logical. Use variational Bayes algorithm from stan? Only kind of working, not recommended.
...	additional arguments to pass to stan function.

Examples

```
#generate a ctStanModel relying heavily on defaults
model<-ctModel(type='ct',
  latentNames=c('eta1','eta2'),
  manifestNames=c('Y1','Y2'),
  MANIFESTVAR=diag(.1,2),
  TDpredNames='TD1',
  TIpredNames=c('TI1','TI2','TI3'),
  LAMBDA=diag(2))

fit<-ctStanFit(ctstantestdat, model,priors=TRUE)

summary(fit)

plot(fit,wait=FALSE)

#### extended examples

library(ctsem)
set.seed(3)

# Data generation (run this, but no need to understand!) -----

Tpoints <- 20
nmanifest <- 4
nlatent <- 2
nsubjects<-20

#random effects
age <- rnorm(nsubjects) #standardised
cint1<-rnorm(nsubjects,2,.3)+age*.5
cint2 <- cint1*.5+rnorm(nsubjects,1,.2)+age*.5
```

```

tdpredeffect <- rnorm(nsubjects,5,.3)+age*.5

for(i in 1:nsubjects){
  #generating model
  gm<-ctModel(Tpoints=Tpoints,n.manifest = nmanifest,n.latent = nlatent,n.TDpred = 1,
    LAMBDA = matrix(c(1,0,0,0, 0,1,.8,1.3),nrow=nmanifest,ncol=nlatent),
    DRIFT=matrix(c(-.3, .2, 0, -.5),nlatent,nlatent),
    TDPREDMEANS=matrix(c(rep(0,Tpoints-10),1,rep(0,9)),ncol=1),
    TDPREDEFFECT=matrix(c(tdpredeffect[i],0),nrow=nlatent),
    DIFFUSION = matrix(c(1, 0, 0, .5),2,2),
    CINT = matrix(c(cint1[i],cint2[i]),ncol=1),
    T0VAR=diag(2,nlatent,nlatent),
    MANIFESTVAR = diag(.5, nmanifest))

  #generate data
  newdat <- ctGenerate(ctmodelobj = gm,n.subjects = 1,burnin = 2,
    dtmat<-rbind(c(rep(.5,8),3,rep(.5,Tpoints-9))))
  newdat[, 'id'] <- i #set id for each subject
  newdat <- cbind(newdat,age[i]) #include time independent predictor
  if(i ==1) {
    dat <- newdat[1:(Tpoints-10),] #pre intervention data
    dat2 <- newdat #including post intervention data
  }
  if(i > 1) {
    dat <- rbind(dat, newdat[1:(Tpoints-10),])
    dat2 <- rbind(dat2,newdat)
  }
}
colnames(dat)[ncol(dat)] <- 'age'
colnames(dat2)[ncol(dat)] <- 'age'

#plot generated data for sanity
plot(age)
matplot(dat[,gm$manifestNames],type='l',pch=1)
plotvar <- 'Y1'
plot(dat[dat[, 'id']==1, 'time'],dat[dat[, 'id']==1,plotvar],type='l',
  ylim=range(dat[,plotvar],na.rm=TRUE))
for(i in 2:nsubjects){
  points(dat[dat[, 'id']==i, 'time'],dat[dat[, 'id']==i,plotvar],type='l',col=i)
}

dat2[,gm$manifestNames][sample(1:length(dat2[,gm$manifestNames]),size = 100)] <- NA

#data structure
head(dat2)

# Model fitting -----
##simple univariate default model

```

```

m <- ctModel(type = 'ct', manifestNames = c('Y1'), LAMBDA = diag(1))
ctModelLatex(m)

#Specify univariate linear growth curve

m1 <- ctModel(type = 'ct',
  manifestNames = c('Y1'), latentNames=c('eta1'),
  DRIFT=matrix(-.0001,nrow=1,ncol=1),
  DIFFUSION=matrix(0,nrow=1,ncol=1),
  T0VAR=matrix(0,nrow=1,ncol=1),
  CINT=matrix(c('cint1'),ncol=1),
  T0MEANS=matrix(c('t0m1'),ncol=1),
  LAMBDA = diag(1),
  MANIFESTMEANS=matrix(0,ncol=1),
  MANIFESTVAR=matrix(c('merror'),nrow=1,ncol=1))

ctModelLatex(m1)

#fit
f1 <- ctStanFit(datalong = dat2, ctstanmodel = m1, optimize=TRUE, priors=FALSE)

summary(f1)

#plots of individual subject models v data
ctKalman(f1,plot=TRUE,subjects=1,kalmanvec=c('y','yprior'),timestep=.01)
ctKalman(f1,plot=TRUE,subjects=1:3,kalmanvec=c('y','ysmooth'),timestep=.01,errorvec=NA)

ctStanPostPredict(f1, wait=FALSE) #compare randomly generated data from posterior to observed data

cf<-ctCheckFit(f1) #compare mean and covariance of randomly generated data to observed cov
plot(cf,wait=FALSE)

### Further example models

#Include intervention
m2 <- ctModel(type = 'ct',
  manifestNames = c('Y1'), latentNames=c('eta1'),
  n.TDpred=1,TDpredNames = 'TD1', #this line includes the intervention
  TDPREDEFFECT=matrix(c('tdpredeffect'),nrow=1,ncol=1), #intervention effect
  DRIFT=matrix(-1e-5,nrow=1,ncol=1),
  DIFFUSION=matrix(0,nrow=1,ncol=1),
  CINT=matrix(c('cint1'),ncol=1),
  T0MEANS=matrix(c('t0m1'),ncol=1),
  T0VAR=matrix(0,nrow=1,ncol=1),
  LAMBDA = diag(1),
  MANIFESTMEANS=matrix(0,ncol=1),
  MANIFESTVAR=matrix(c('merror'),nrow=1,ncol=1))

#Individual differences in intervention, Bayesian estimation, covariates
m2i <- ctModel(type = 'ct',

```

```

manifestNames = c('Y1'), latentNames=c('eta1'),
TIpredNames = 'age',
TDpredNames = 'TD1', #this line includes the intervention
TDPREDEFFECT=matrix(c('tdpredeffect||TRUE'),nrow=1,ncol=1), #intervention effect
DRIFT=matrix(-1e-5,nrow=1,ncol=1),
DIFFUSION=matrix(0,nrow=1,ncol=1),
CINT=matrix(c('cint1'),ncol=1),
T0MEANS=matrix(c('t0m1'),ncol=1),
T0VAR=matrix(0,nrow=1,ncol=1),
LAMBDA = diag(1),
MANIFESTMEANS=matrix(0,ncol=1),
MANIFESTVAR=matrix(c('merror'),nrow=1,ncol=1))

#Including covariate effects
m2ic <- ctModel(type = 'ct',
  manifestNames = c('Y1'), latentNames=c('eta1'),
  n.TIpred = 1, TIpredNames = 'age',
  n.TDpred=1,TDpredNames = 'TD1', #this line includes the intervention
  TDPREDEFFECT=matrix(c('tdpredeffect'),nrow=1,ncol=1), #intervention effect
  DRIFT=matrix(-1e-5,nrow=1,ncol=1),
  DIFFUSION=matrix(0,nrow=1,ncol=1),
  CINT=matrix(c('cint1'),ncol=1),
  T0MEANS=matrix(c('t0m1'),ncol=1),
  T0VAR=matrix(0,nrow=1,ncol=1),
  LAMBDA = diag(1),
  MANIFESTMEANS=matrix(0,ncol=1),
  MANIFESTVAR=matrix(c('merror'),nrow=1,ncol=1))

m2ic$pars$indvarying[m2ic$pars$matrix %in% 'TDPREDEFFECT'] <- TRUE

#Include deterministic dynamics
m3 <- ctModel(type = 'ct',
  manifestNames = c('Y1'), latentNames=c('eta1'),
  n.TDpred=1,TDpredNames = 'TD1', #this line includes the intervention
  TDPREDEFFECT=matrix(c('tdpredeffect'),nrow=1,ncol=1), #intervention effect
  DRIFT=matrix('drift11',nrow=1,ncol=1),
  DIFFUSION=matrix(0,nrow=1,ncol=1),
  CINT=matrix(c('cint1'),ncol=1),
  T0MEANS=matrix(c('t0m1'),ncol=1),
  T0VAR=matrix('t0var11',nrow=1,ncol=1),
  LAMBDA = diag(1),
  MANIFESTMEANS=matrix(0,ncol=1),
  MANIFESTVAR=matrix(c('merror1'),nrow=1,ncol=1))

#Add system noise to allow for fluctuations that persist in time
m3n <- ctModel(type = 'ct',
  manifestNames = c('Y1'), latentNames=c('eta1'),

```

```

n.TDpred=1,TDpredNames = 'TD1', #this line includes the intervention
TDPREDEFFECT=matrix(c('tdpredeffect'),nrow=1,ncol=1), #intervention effect
DRIFT=matrix('drift11',nrow=1,ncol=1),
DIFFUSION=matrix('diffusion',nrow=1,ncol=1),
CINT=matrix(c('cint1'),ncol=1),
T0MEANS=matrix(c('t0m1'),ncol=1),
T0VAR=matrix('t0var11',nrow=1,ncol=1),
LAMBDA = diag(1),
MANIFESTMEANS=matrix(0,ncol=1),
MANIFESTVAR=matrix(c(0),nrow=1,ncol=1))

#include 2nd latent process

m4 <- ctModel(n.manifest = 2,n.latent = 2, type = 'ct',
  manifestNames = c('Y1','Y2'), latentNames=c('L1','L2'),
  n.TDpred=1,TDpredNames = 'TD1',
  TDPREDEFFECT=matrix(c('tdpredeffect1','tdpredeffect2'),nrow=2,ncol=1),
  DRIFT=matrix(c('drift11','drift21','drift12','drift22'),nrow=2,ncol=2),
  DIFFUSION=matrix(c('diffusion11','diffusion21',0,'diffusion22'),nrow=2,ncol=2),
  CINT=matrix(c('cint1','cint2'),nrow=2,ncol=1),
  T0MEANS=matrix(c('t0m1','t0m2'),nrow=2,ncol=1),
  T0VAR=matrix(c('t0var11','t0var21',0,'t0var22'),nrow=2,ncol=2),
  LAMBDA = matrix(c(1,0,0,1),nrow=2,ncol=2),
  MANIFESTMEANS=matrix(c(0,0),nrow=2,ncol=1),
  MANIFESTVAR=matrix(c('merror1',0,0,'merror2'),nrow=2,ncol=2))

#dynamic factor model -- fixing CINT to 0 and freeing indicator level intercepts

m3df <- ctModel(type = 'ct',
  manifestNames = c('Y2','Y3'), latentNames=c('eta1'),
  n.TDpred=1,TDpredNames = 'TD1', #this line includes the intervention
  TDPREDEFFECT=matrix(c('tdpredeffect'),nrow=1,ncol=1), #intervention effect
  DRIFT=matrix('drift11',nrow=1,ncol=1),
  DIFFUSION=matrix('diffusion',nrow=1,ncol=1),
  CINT=matrix(c(0),ncol=1),
  T0MEANS=matrix(c('t0m1'),ncol=1),
  T0VAR=matrix('t0var11',nrow=1,ncol=1),
  LAMBDA = matrix(c(1,'Y3loading'),nrow=2,ncol=1),
  MANIFESTMEANS=matrix(c('Y2_int','Y3_int'),nrow=2,ncol=1),
  MANIFESTVAR=matrix(c('Y2residual',0,0,'Y3residual'),nrow=2,ncol=2))

```

Description

Either to include different data, or because you have upgraded ctsem and the internal data structure has changed.

Usage

```
ctStanFitUpdate(oldfit, data = NA, recompile = FALSE, refit = FALSE, ...)
```

Arguments

oldfit	fit object to be upgraded
data	replacement long format data object
recompile	whether to force a recompile – safer but slower and usually unnecessary.
refit	if TRUE, refits the model using the old estimates as a starting point. Only applicable for optimized fits, not sampling.
...	extra arguments to pass to ctStanFit

Value

updated ctStanFit object.

Examples

```
newfit <- ctStanFitUpdate(ctstantestfit,refit=FALSE)
```

ctStanGenerate	<i>Generate data from a ctstanmodel object</i>
----------------	--

Description

Generate data from a ctstanmodel object

Usage

```
ctStanGenerate(
  cts,
  datastruct = NA,
  is = FALSE,
  fullposterior = TRUE,
  nsamples = 200,
  paronly = FALSE,
  cores = 2
)
```

Arguments

cts	ctStanModel , or ctStanFit,object.
datastruct	long format data structure as used by ctsem. Not used if cts is a ctStanFit object.
is	If optimizing, follow up with importance sampling?
fullposterior	Generate from the full posterior or just the (unconstrained) mean?
nsamples	How many samples to generate?
parsonly	If TRUE, only return samples of raw parameters, don't generate data.
cores	Number of cpu cores to use.

Value

List containing Y, and array of nsamples by data rows by manifest variables, and llrow, an array of nsamples by data rows log likelihoods.

Examples

```
#generate and plot samples from prior predictive
priorpred <- ctStanGenerate(cts = ctstestfit,cores=2,nsamples = 50)
```

ctStanGenerateFromFit *Add a \$generated object to ctstanfit object, with random data generated from posterior of ctstanfit object*

Description

Add a \$generated object to ctstanfit object, with random data generated from posterior of ctstanfit object

Usage

```
ctStanGenerateFromFit(
  fit,
  nsamples = 200,
  fullposterior = FALSE,
  verboseErrors = FALSE,
  cores = 2
)
```

Arguments

fit	ctstanfit object
nsamples	Positive integer specifying number of datasets to generate.
fullposterior	Logical indicating whether to sample from the full posterior (original nsamples) or the posterior mean.
verboseErrors	if TRUE, print verbose output when errors in generation encountered.
cores	Number of cpu cores to use.

Value

Matrix of generated data – one dataset per iteration, according to original time and missingness structure.

Examples

```
gen <- ctStanGenerateFromFit(ctstantestfit, nsamples=3,fullposterior=TRUE,cores=1)
plot(gen$generated$Y[3,,2],type='l') #Third random data sample, 2nd manifest var, all time points.
```

ctStanKalman

Get Kalman filter estimates from a ctStanFit object

Description

Get Kalman filter estimates from a ctStanFit object

Usage

```
ctStanKalman(
  fit,
  nsamples = NA,
  pointest = TRUE,
  collapsefunc = NA,
  cores = 1,
  subjects = 1:max(fit$standata$subject),
  timestep = "asdata",
  maxtime = "asdata",
  standardisederrors = FALSE,
  subjectpars = TRUE,
  tformsubjectpars = TRUE,
  indvarstates = FALSE,
  removeObs = F,
  ...
)
```

Arguments

fit	fit object from ctStanFit .
nsamples	either NA (to extract all) or a positive integer from 1 to maximum samples in the fit.
pointest	If TRUE, uses the posterior mode as the single sample.
collapsefunc	function to apply over samples, such as mean
cores	Integer number of cpu cores to use. Only needed if savescores was set to FALSE when fitting.
subjects	integer vector of subjects to compute for.

timestep	Either a positive numeric value, 'asdata' to use the times in the dataset, or 'auto' to select a timestep automatically (resulting in some interpolation but not excessive computation).
maxtime	only relevant if timestep is not 'asdata'. Positive numeric denoting max time for computations.
standardisederrors	If TRUE, computes standardised errors for prior, upd, smooth conditions.
subjectpars	if TRUE, state estimates are not returned, instead, predictions of each subjects parameters are returned, for parameters that had random effects specified.
tformsubjectpars	if FALSE, subject level parameters are returned in raw, pre transformation form.
indvarstates	if TRUE, do not remove indvarying states from output
removeObs	Logical or integer. If TRUE, observations (but not covariates) are set to NA, so only expectations based on parameters and covariates are returned. If a positive integer N, every N observations are retained while others are set NA for computing model expectations – useful for observing prediction performance forward further in time than one observation.
...	additional arguments to collapsefunc.

Value

list containing Kalman filter elements, each element in array of iterations, data row, variables. llrow is the log likelihood for each row of data.

Examples

```
k=ctStanKalman(ctstantestfit,subjectpars=TRUE,collapsefunc=mean)
```

ctStanModel	<i>Convert a frequentist (omx) ctsem model specification to Bayesian (Stan).</i>
-------------	--

Description

Convert a frequentist (omx) ctsem model specification to Bayesian (Stan).

Usage

```
ctStanModel(ctmodelobj, type = "ct", tipredDefault = TRUE)
```

Arguments

ctmodelobj	ctsem model object of type 'omx' (default)
type	either 'ct' for continuous time, or 'dt' for discrete time.
tipredDefault	Logical. TRUE sets any parameters with unspecified time independent predictor effects to have effects estimated, FALSE fixes the effect to zero unless individually specified.

Value

List object of class `ctStanModel`, with random effects specified for any intercept type parameters (TOMEANS, MANIFESTMEANS, and or CINT), and time independent predictor effects for all parameters. Adjust these after initial specification by directly editing the `pars` subobject, so `model$pars`.

Examples

```
model <- ctModel(type='omx', Tpoints=50,
  n.latent=2, n.manifest=1,
  manifestNames='sunspots',
  latentNames=c('ss_level', 'ss_velocity'),
  LAMBDA=matrix(c( 1, 'ma1' ), nrow=1, ncol=2),
  DRIFT=matrix(c(0, 1, 'a21', 'a22'), nrow=2, ncol=2, byrow=TRUE),
  MANIFESTMEANS=matrix(c('m1'), nrow=1, ncol=1),
  # MANIFESTVAR=matrix(0, nrow=1, ncol=1),
  CINT=matrix(c(0, 0), nrow=2, ncol=1),
  DIFFUSION=matrix(c(
    0, 0,
    0, "diffusion"), ncol=2, nrow=2, byrow=TRUE))

stanmodel=ctStanModel(model)
```

 ctStanParnames

ctStanParnames

Description

Gets internal stan parameter names of a `ctStanFit` object sampled via stan based on specified substrings.

Usage

```
ctStanParnames(x, substrings = c("pop_", "popsd"))
```

Arguments

<code>x</code>	<code>ctStanFit</code> object
<code>substrings</code>	vector of character strings, parameter names of the stan model containing any of these strings will be returned. Useful strings may be 'pop_' for population means, 'popsd' for population standard deviations, or specific combinations such as 'pop_DRIFT' for the population means of temporal dynamics parameters

Value

vector of character strings.

Examples

```

sunspots<-sunspot.year
sunspots<-sunspots[50: (length(sunspots) - (1988-1924))]
id <- 1
time <- 1749:1924
datalong <- cbind(id, time, sunspots)

#setup model
ssmodel <- ctModel(type='ct', n.latent=2, n.manifest=1,
  manifestNames='sunspots',
  latentNames=c('ss_level', 'ss_velocity'),
  LAMBDA=matrix(c( 1, 'ma1 | log(1+(exp(param)))' ), nrow=1, ncol=2),
  DRIFT=matrix(c(0, 'a21 | -log(1+exp(param))', 1, 'a22'), nrow=2, ncol=2),
  MANIFESTMEANS=matrix(c('m1|param * 10 + 44'), nrow=1, ncol=1),
  MANIFESTVAR=diag(0,1), #As per original spec
  CINT=matrix(c(0, 0), nrow=2, ncol=1),
  DIFFUSION=matrix(c(0, 0, 0, "diffusion"), ncol=2, nrow=2))

#fit
ssfit <- ctStanFit(datalong, ssmodel, iter=2,
  optimize=FALSE, chains=1)
ctStanParNames(ssfit,substrings=c('pop_', 'popsd'))

```

ctStanPlotPost

ctStanPlotPost

Description

Plots prior and posterior distributions of model parameters in a ctStanModel or ctStanFit object.

Usage

```

ctStanPlotPost(
  obj,
  rows = "all",
  npp = 6,
  priorwidth = TRUE,
  smoothness = 1,
  priorsamples = 10000,
  plot = TRUE,
  wait = FALSE,
  ...
)

```

Arguments

obj	fit or model object as generated by <code>ctStanFit</code> , <code>ctModel</code> , or <code>ctStanModel</code> .
rows	vector of integers denoting which rows of <code>obj\$setup\$popsetup</code> to plot priors for. Character string 'all' plots all rows with parameters to be estimated.
npp	Integer number of parameters to show per page.
priorwidth	if TRUE, plots will be scaled to show bulk of both the prior and posterior distributions. If FALSE, scale is based only on the posterior.
smoothness	Positive numeric – multiplier to modify smoothness of density plots, higher is smoother but can cause plots to exceed natural boundaries, such as standard deviations below zero.
priorsamples	number of samples from prior to use. More is slower.
plot	Logical, if FALSE, ggplot objects are returned in a list instead of plotting.
wait	If true, user is prompted to continue before plotting next graph. If false, graphs are plotted one after another without waiting.
...	Parameters to pass to <code>ctStanFit</code> . <code>cores = x</code> will speed things up, where x is the number of cpu cores to use.

Examples

```
ctStanPlotPost(ctstantestfit, rows=3:4)
```

ctStanPostPredict	<i>Compares model implied density and values to observed, for a ctStanFit object.</i>
-------------------	---

Description

Compares model implied density and values to observed, for a `ctStanFit` object.

Usage

```
ctStanPostPredict(
  fit,
  diffsize = 1,
  jitter = 0.02,
  wait = TRUE,
  probs = c(0.025, 0.5, 0.975),
  datarows = "all",
  nsamples = 500,
  resolution = 100,
  plot = TRUE
)
```

Arguments

fit	ctStanFit object.
diffsize	Integer > 0. Number of discrete time lags to use for data viz.
jitter	Positive numeric between 0 and 1, if TRUE, jitters empirical data by specified proportion of std dev.
wait	Logical, if TRUE and plot=TRUE, waits for input before plotting next plot.
probs	Vector of length 3 containing quantiles to plot – should be rising numeric values between 0 and 1.
datarows	integer vector specifying rows of data to plot. Otherwise 'all' uses all data.
nsamples	Number of datasets to generate for comparisons, if fit object does not contain generated data already.
resolution	Positive integer, the number of rows and columns to split plots into for shading.
plot	logical. If FALSE, a list of ggplot objects is returned.

Details

This function relies on the data generated during each iteration of fitting to approximate the model implied distributions – thus, when limited iterations are available, the approximation will be worse.

Value

If plot=FALSE, an array containing quantiles of generated data. If plot=TRUE, nothing, only plots. if plot=TRUE, nothing is returned and plots are created. Otherwise, a list containing ggplot objects is returned and may be customized as desired.

Examples

```
#'
ctStanPostPredict(ctstantestfit,wait=FALSE, diffsize=2,resolution=100)
```

ctStanSubjectPars *Extract an array of subject specific parameters from a ctStanFit object.*

Description

Extract an array of subject specific parameters from a ctStanFit object.

Usage

```
ctStanSubjectPars(fit, pointest = TRUE, cores = 2, nsamples = "all")
```

Arguments

<code>fit</code>	fit object
<code>pointest</code>	if TRUE, returns only the set of individual difference parameters based on the max a posteriori estimate (or the median if sampling approaches were used).
<code>cores</code>	Number of cores to use.
<code>nsamples</code>	Number of samples to calculate parameters for. Not used if <code>pointest=TRUE</code> .

Details

This function returns the estimates of individual parameters, taking into account any covariates and random effects.

Value

an `nsamples` by `nsubjects` by `nparams` array.

Examples

```
indpars <- ctStanSubjectPars(ctstantestfit)
dimnames(indpars)
plot(indpars[1,,'cint1'],indpars[1,,'cint2'])
```

<code>ctstantestdat</code>	<i>ctstantestdat</i>
----------------------------	----------------------

Description

Generated dataset for testing [ctStanFit](#) from `ctsem` package.

Format

matrix

<code>ctstantestfit</code>	<i>ctstantestfit</i>
----------------------------	----------------------

Description

Dummy fit for testing functions from `ctsem` package.

Format

`ctStanFit` object

ctStanTIpredEffects *Get time independent predictor effect estimates*

Description

Computes and plots combined effects and quantiles for effects of time independent predictors on subject level parameters of a ctStanFit object.

Usage

```
ctStanTIpredEffects(
  fit,
  returndifference = FALSE,
  probs = c(0.025, 0.5, 0.975),
  includeMeanUncertainty = FALSE,
  whichTIpreds = 1,
  parmatrices = TRUE,
  whichpars = "all",
  nsamples = 100,
  timeinterval = 1,
  nsubjects = 20,
  filter = NA,
  plot = FALSE
)
```

Arguments

fit	fit object from ctStanFit
returndifference	logical. If FALSE, absolute parameter values are returned. If TRUE, only the effect of the covariate (i.e. without the average value of the parameter) are returned. The former can be easier to interpret, but the latter are more likely to fit multiple plots together. Not used if parmatrices=TRUE.
probs	numeric vector of quantile probabilities from 0 to 1. Specify 3 values if plotting, the 2nd will be drawn as a line with uncertainty polygon based on 1st and 3rd.
includeMeanUncertainty	if TRUE, output includes sampling variation in the mean parameters. If FALSE, mean parameters are fixed at their median, only uncertainty in time independent predictor effects is included.
whichTIpreds	integer vector specifying which of the tipreds in the fit object you want to use to calculate effects. Unless quadratic / higher order versions of predictors have been included, selecting more than one probably doesn't make sense. If for instance a squared predictor has been included, then you can specify both the linear and squared version. The x axis of the plot (if generated) will be based off the first indexed predictor. To check what predictors are in the model, run <code>fit\$ctstanmodel\$TIpredNames</code> .

parmatrices	Logical. If TRUE (default), system matrices rather than specific parameters are referenced – e.g. 'DRIFT' instead of a parameter name like drift12.
whichpars	if parmatrices==TRUE, character vector specifying which matrices, and potentially which indices of the matrices, to plot. c('dtDRIFT[2,1]', 'DRIFT') would output for row 2 and column 1 of the discrete time drift matrix, as well as all indices of the continuous time drift matrix. If parmatrices==FALSE, integer vector specifying which of the subject level parameters to compute effects on. The integers corresponding to certain parameters can be found in the param column of the fit\$setup\$matsetup object. In either case 'all' uses all available parameters.
nsamples	Positive integer specifying the maximum number of saved iterations to use. Character string 'all' can also be used.
timeinterval	positive numeric indicating time interval to use for discrete time parameter matrices, if parmatrices=TRUE.
nsubjects	Positive integer specifying the number of subjects to compute values for. When only one TIpred is used, this specifies the number of points along the curve. Character string 'all' can also be used. Time taken for plotting is a function of nsubjects*niterations.
filter	either NA, or a length 2 vector, where the first element contains the time independent predictor index to filter by, and the second contains the comparison operator in string form (e.g. "< 3", to only calculate effects for subjects where the tipreds of the denoted index are less than 3).
plot	Logical. If TRUE, nothing is returned but instead <code>ctPlotArray</code> is used to plot the output instead.

Value

Either a three dimensional array of predictor effects, or nothing with a plot generated.

Examples

```
ctStanTIpredeffects(ctstantestfit,
  whichpars=c('CINT', 'dtDIFFUSION[2,2]'), plot=TRUE)
```

 ctStanUpdModel

Update an already compiled and fit ctStanFit object

Description

Allows one to change data and or model elements that don't require recompiling, then re fit.

Usage

```
ctStanUpdModel(fit, datalong, ctstanmodel, ...)
```

Arguments

fit	ctStanFit object
datalong	data as normally passed to <code>ctStanFit</code>
ctstanmodel	model as normally passed to <code>ctStanFit</code>
...	extra args for <code>ctStanFit</code>

ctWideNames	<i>ctWideNames sets default column names for wide ctsem datasets. Primarily intended for internal ctsem usage.</i>
-------------	--

Description

ctWideNames sets default column names for wide ctsem datasets. Primarily intended for internal ctsem usage.

Usage

```
ctWideNames(
  n.manifest,
  Tpoints,
  n.TDpred = 0,
  n.TIpred = 0,
  manifestNames = "auto",
  TDpredNames = "auto",
  TIpredNames = "auto"
)
```

Arguments

n.manifest	number of manifest variables per time point in the data.
Tpoints	Maximum number of discrete time points (waves of data, or measurement occasions) for an individual in the input data structure.
n.TDpred	number of time dependent predictors in the data structure.
n.TIpred	number of time independent predictors in the data structure.
manifestNames	vector of character strings giving column names of manifest indicator variables
TDpredNames	vector of character strings giving column names of time dependent predictor variables
TIpredNames	vector of character strings giving column names of time independent predictor variables

ctWideToLong	<i>ctWideToLong Convert ctsem wide to long format</i>
--------------	---

Description

ctWideToLong Convert ctsem wide to long format

Usage

```
ctWideToLong(
  datawide,
  Tpoints,
  n.manifest,
  n.TDpred = 0,
  n.TIpred = 0,
  manifestNames = "auto",
  TDpredNames = "auto",
  TIpredNames = "auto"
)
```

Arguments

datawide	ctsem wide format data
Tpoints	number of measurement occasions in data
n.manifest	number of manifest variables
n.TDpred	number of time dependent predictors
n.TIpred	number of time independent predictors
manifestNames	Character vector of manifest variable names.
TDpredNames	Character vector of time dependent predictor names.
TIpredNames	Character vector of time independent predictor names.

Details

Names must account for **all** the columns in the data - i.e. do not leave certain variables out just because you do not need them.

Examples

```
#create wide data
wideexample <- ctLongToWide(datalong = ctstantestdat, id = "id",
  time = "time", manifestNames = c("Y1", "Y2"),
  TDpredNames = "TD1", TIpredNames = c("TI1", "TI2", "TI3"))

wide <- ctIntervalise(datawide = wideexample, Tpoints = 10, n.manifest = 2,
  n.TDpred = 1, n.TIpred = 3, manifestNames = c("Y1", "Y2"),
  TDpredNames = "TD1", TIpredNames = c("TI1", "TI2", "TI3") )
```

```
#Then convert to long format
longexample <- ctWideToLong(datawide = wideexample, Tpoints=10,
n.manifest=2, manifestNames = c("Y1", "Y2"),
n.TDpred=1, TDpredNames = "TD1",
n.TIpred=3, TIpredNames = c("TI1", "TI2", "TI3"))

#Then convert the time intervals to absolute time
long <- ctDeintervalise(datalong = longexample, id='id', dT='dT')
head(long,22)
```

datastructure	<i>datastructure</i>
---------------	----------------------

Description

Simulated example dataset for the ctsem package

Format

2 by 15 matrix containing containing ctsem wide format data. 3 measurement occasions of manifest variables Y1 and Y2, 2 measurement occasions of time dependent predictor TD1, 2 measurement intervals dTx, and 2 time independent predictors TI1 and TI2, for 2 individuals.

inv_logit	<i>Inverse logit</i>
-----------	----------------------

Description

Maps the stan function so the same code works in R.

Usage

```
inv_logit(x)
```

Arguments

x value to calculate the inverse logit for.

Examples

```
inv_logit(-3)
```

log1p_exp	<i>log1p_exp</i>
-----------	------------------

Description

Maps the stan function so the same code works in R.

Usage

```
log1p_exp(x)
```

Arguments

x value to use.

Examples

```
log1p_exp(-3)
```

longexample	<i>longexample</i>
-------------	--------------------

Description

Simulated example dataset for the ctsem package

Format

7 by 8 matrix containing ctsem long format data, for two subjects, with three manifest variables Y1, Y2, Y3, one time dependent predictor TD1, two time independent predictors TI1 and TI2, and absolute timing information Time.

Oscillating

Oscillating

Description

Simulated example dataset for the ctsem package.

Format

200 by 21 matrix containing containing ctsem wide format data. 11 measurement occasions and 10 measurement intervals for each of 200 individuals

Source

See <https://bpspsychub.onlinelibrary.wiley.com/doi/10.1111/j.2044-8317.2012.02043.x>

plot.ctKalmanDF

Plots Kalman filter output from ctKalman.

Description

Plots Kalman filter output from ctKalman.

Usage

```
## S3 method for class 'ctKalmanDF'
plot(
  x,
  subjects = unique(x$Subject),
  kalmanvec = c("y", "yprior"),
  errorvec = "auto",
  errormultiply = 1.96,
  plot = TRUE,
  elementNames = NA,
  polygonsteps = 10,
  polygonalpha = 0.1,
  facets = "Variable",
  ...
)
```

Arguments

x	Output from <code>ctKalman</code> . In general it is easier to call <code>ctKalman</code> directly with the <code>plot=TRUE</code> argument, which calls this function.
subjects	vector of integers denoting which subjects (from 1 to N) to plot predictions for.
kalmanvec	string vector of names of any elements of the output you wish to plot, the defaults of 'y' and 'ysmooth' plot the original data, 'y', and the estimates of the 'true' value of y given all data. Replacing 'y' by 'eta' will plot latent states instead (though 'eta' alone does not exist) and replacing 'smooth' with 'upd' or 'prior' respectively plots updated (conditional on all data up to current time point) or prior (conditional on all previous data) estimates.
errorvec	vector of names indicating which kalmanvec elements to plot uncertainty bands for. 'auto' plots all possible.
errormultiply	Numeric denoting the multiplication factor of the std deviation of errorvec objects. Defaults to 1.96, for 95% intervals.
plot	if FALSE, plots are not generated and the ggplot object is simply returned invisibly.
elementNames	if NA, all relevant object elements are included – e.g. if yprior is in the kalmanvec argument, all manifest variables are plotted, and likewise for latent states if etasmooth was specified. Alternatively, a character vector specifying the manifest and latent names to plot explicitly can be specified.
polygonsteps	Number of steps to use for uncertainty band shading.
polygonalpha	Numeric for the opacity of the uncertainty region.
facets	when multiple subjects are included in multivariate plots, the default is to facet plots by variable type. This can be set to NA for no facets, or "Subject" for faceting by subject.
...	not used.

Value

A ggplot2 object. Side effect – Generates plots.

Examples

```
### Get output from ctKalman
x<-ctKalman(ctstantestfit,subjects=2,timestep=.01)

### Plot with plot.ctKalmanDF
plot(x, subjects=2)

###Single step procedure:
ctKalman(ctstantestfit,subjects=2,
  kalmanvec=c('y','yprior'),
  elementNames=c('Y1','Y2'),
  plot=TRUE,timestep=.01)
```

plot.ctStanFit *plot.ctStanFit*

Description

Plots for ctStanFit objects

Usage

```
## S3 method for class 'ctStanFit'
plot(x, types = "all", wait = TRUE, ...)
```

Arguments

x	Fit object from ctStanFit .
types	Vector of character strings defining which plots to create. 'all' plots all possible types, including: 'regression', 'kalman', 'priorcheck', 'trace', 'density', 'intervals'.
wait	Logical. Pause between plots?
...	Arguments to pass through to the specific plot functions. Bewar of clashes may occur if types='all'. For details see the specific functions generating each type of plot.

Details

This function is just a wrapper calling the necessary functions for plotting - it may be simpler in many cases to access those directly. They are: [ctStanDiscretePars](#), [ctKalman](#), [ctStanPlotPost](#), [stan_trace](#), [stan_dens](#), [stan_plot](#) rstan offers many plotting possibilities not available here, to use that functionality one must simply call the relevant rstan plotting function. Use `x$stanfit` as the stan fit object (where x is the name of your ctStanFit object). Because a ctStanFit object has many parameters, the additional argument `pars=ctStanParnames(x, 'pop_')` is recommended. This denotes population means, but see [ctStanParnames](#) for other options.

Value

Nothing. Generates plots.

Examples

```
plot(ctstantestfit, types=c('regression', 'kalman', 'priorcheck'), wait=FALSE)
```

plot.ctStanModel *Prior plotting*

Description

Plots priors for free model parameters in a ctStanModel.

Usage

```
## S3 method for class 'ctStanModel'
plot(
  x,
  rows = "all",
  wait = FALSE,
  nsamples = 1e+06,
  rawpopstd = "marginalise",
  inddifdevs = c(-1, 1),
  inddifsd = 0.1,
  plot = TRUE,
  ...
)
```

Arguments

x	ctStanModel object as generated by ctModel with type='ct' or 'dt'.
rows	vector of integers denoting which rows of ctstanmodel\$pars to plot priors for. Character string 'all' plots all rows with parameters to be estimated.
wait	If true, user is prompted to continue before plotting next graph.
nsamples	Numeric. Higher values increase fidelity (smoothness / accuracy) of density plots, at cost of speed.
rawpopstd	Either 'marginalise' to sample from the specified (in the ctstanmodel) prior distribution for the raw population standard deviation, or a numeric value to use for the raw population standard deviation for all subject level prior plots - the plots in dotted blue or red.
inddifdevs	numeric vector of length 2, setting the means for the individual differences distributions.
inddifsd	numeric, setting the standard deviation of the population means used to generate individual difference distributions.
plot	If FALSE, outputs list of GGplot objects that can be further modified.
...	not used.

Details

Plotted in black is the prior for the population mean. In red and blue are the subject level priors that result given that the population mean is estimated as 1 std deviation above the mean of the prior, or 1 std deviation below. The distributions around these two points are then obtained by marginalising over the prior for the raw population std deviation - so the red and blue distributions do not represent any specific subject level prior, but rather characterise the general amount and shape of possible subject level priors at the specific points of the population mean prior.

Examples

```
model <- ctModel(type='ct',
  manifestNames='sunspots',
  latentNames=c('ss_level', 'ss_velocity'),
  LAMBDA=matrix(c(1, 'ma1'), nrow=1, ncol=2),
  DRIFT=matrix(c(0, 1, 'a21', 'a22'), nrow=2, ncol=2, byrow=TRUE),
  MANIFESTMEANS=matrix(c('m1'), nrow=1, ncol=1),
  # MANIFESTVAR=matrix(0, nrow=1, ncol=1),
  CINT=matrix(c(0, 0), nrow=2, ncol=1),
  DIFFUSION=matrix(c(
    0, 0,
    0, "diffusion"), ncol=2, nrow=2, byrow=TRUE))

plot(model, rows=8)
```

plotctACF

Plot an approximate continuous-time ACF object from ctACF

Description

Plot an approximate continuous-time ACF object from ctACF

Usage

```
plotctACF(
  ctacfobj,
  df = "auto",
  quantiles = c(0.025, 0.5, 0.975),
  separateLearnRates = FALSE,
  reducedXlim = 1,
  estimateSpline = TRUE
)
```

Arguments

ctacfobj	object
df	df for the basis spline.
quantiles	quantiles to plot.

`separateLearnRates` if TRUE, estimate the learning rate for the quantile splines for each combination of variables. Slower but theoretically more accurate.

`reducedXlim` if non-zero, n timesteps are removed from the upper and lower end of the x range where the spline estimates are less likely to be reasonable.

`estimateSpline` if TRUE, quantile spline regression is used, otherwise the samples are simply plotted as lines and the other arguments here are not used.

Value

a ggplot object

Examples

```
data.table::setDTthreads(1) #ignore this line
# Example usage:
head(ctstantestdat)
ac=ctACF(ctstantestdat, varnames=c('Y1'), idcol='id', timecol='time', timestep=.5, nboot=5, plot=FALSE)
plotctACF(ac, reducedXlim=0)
```

sdpcor2cov

sdcor2cov

Description

Converts a lower triangular matrix with standard deviations on the diagonal and partial correlations on lower triangle, to a covariance (or cholesky decomposed covariance)

Usage

```
sdpcor2cov(mat, coronly = FALSE, cholesky = FALSE)
```

Arguments

`mat` input square matrix with std dev on diagonal and lower tri of partial correlations.

`coronly` if TRUE, ignores everything except the lower triangle and outputs correlation.

`cholesky` Logical. To return the cholesky decomposition instead of full covariance, set to TRUE.

Examples

```
testmat <- diag(exp(rnorm(5,-3,2)),5) #generate arbitrary std deviations
testmat[row(testmat) > col(testmat)] <- runif((5^2-5)/2, -1, 1)
print(testmat)
covmat <- sdpcor2cov(testmat) #convert to covariance
cov2cor(covmat) #convert covariance to correlation
```

```
standatact_specificsubjects
```

Adjust standata from ctsem to only use specific subjects

Description

Adjust standata from ctsem to only use specific subjects

Usage

```
standatact_specificsubjects(standata, subjects, timestep = NA)
```

Arguments

standata	standata
subjects	vector of subjects
timestep	ignored at present

Value

list of updated structure

Examples

```
d <- standatact_specificsubjects(ctstantestfit$standata, 1:2)
```

```
stanoptimis
```

Optimize / importance sample a stan or ctStan model.

Description

Optimize / importance sample a stan or ctStan model.

Usage

```
stanoptimis(
  standata,
  sm,
  init = "random",
  initsd = 0.01,
  estonly = FALSE,
  tol = 1e-08,
  stochastic = TRUE,
  priors = TRUE,
  carefulfit = TRUE,
```

```

bootstrapUncertainty = FALSE,
subsamplesize = 1,
parsteps = c(),
parstepsAutoModel = FALSE,
groupFreeThreshold = 0.5,
plot = FALSE,
is = FALSE,
isitersize = 1000,
isESS = 100,
finishesamples = 1000,
lproughnesstarget = 0.2,
verbose = 0,
cores = 2,
matsetup = NA,
nsubsets = 10,
stochasticTolAdjust = 1000
)

```

Arguments

standata	list object conforming to rstan data standards.
sm	compiled stan model object.
init	vector of unconstrained parameter values, or character string 'random' to initialise with random values very close to zero.
initsd	positive numeric specifying sd of normal distribution governing random sample of init parameters, if init='random' .
estonly	if TRUE, just return point estimates under \$rawest subobject.
tol	objective tolerance.
stochastic	Logical. Use stochastic gradient descent as main optimizer. Always finishes (double checks) with mize (bfgs) optimizer.
priors	logical. If TRUE, a priors integer is set to 1 (TRUE) in the standata object – only has an effect if the stan model uses this value.
carefulfit	Logical. If TRUE, priors are always used for a rough first pass to obtain starting values when priors=FALSE
bootstrapUncertainty	Logical. If TRUE, subject wise gradient contributions are resampled to estimate the hessian, for computing standard errors or initializing importance sampling.
subsamplesize	value between 0 and 1 representing proportion of subjects to include in first pass fit.
parsteps	ordered list of vectors of integers denoting which parameters should begin fixed at zero, and freed sequentially (by list order). Useful for complex models, e.g. keep all cross couplings fixed to zero as a first step, free them in second step.
parstepsAutoModel	if TRUE, determines model structure for the parameters specified in parsteps automatically. If 'group', determines this on a group level first and then a subject level. Primarily for internal ctsem use, see ?ctFitAuto.

groupFreeThreshold	threshold for determining whether a parameter is free in a group level model. If the proportion of subjects with a non-zero parameter is above this threshold, the parameter is considered free. Only used with parstepsAutoModel = 'group'.
plot	Logical. If TRUE, plot iteration details. Probably slower.
is	Logical. Use mixture importance sampling, or just return map estimates?
isitersize	Number of samples of approximating distribution per iteration of importance sampling.
isESS	target effective sample size for importance sampling. If is=TRUE, this is used to determine the number of samples to draw from the approximating distribution.
finishsamples	Number of samples to draw (either from hessian based covariance or posterior distribution) for final results computation.
lproughnesstarget	target log posterior roughness for stochastic optimizer (suggest between .05 and .4).
verbose	Integer from 0 to 2. Higher values print more information during model fit – for debugging.
cores	Number of cpu cores to use, should be at least 2.
matsetup	subobject of ctStanFit output. If provided, parameter names instead of numbers are output for any problem indications.
nsubsets	number of subsets for stochastic optimizer. Subsets are further split across cores, but each subjects data remains whole – processed by one core in one subset.
stochasticTolAdjust	Multiplier for stochastic optimizer tolerance.

Value

list containing fit elements

stanWplot	<i>Runs stan, and plots sampling information while sampling.</i>
-----------	--

Description

Runs stan, and plots sampling information while sampling.

Usage

```
stanWplot(object, iter = 2000, chains = 4, ...)
```

Arguments

object	stan model object
iter	Number of iterations
chains	Number of chains
...	All the other regular arguments to stan()

Details

On windows, requires Rtools installed and able to be found by `pkgbuild::rtools_path()`

Examples

```
library(rstan)
#### example 1
scode <- "
parameters {
  real y[2];
}
model {
  y[1] ~ normal(0, .5);
  y[2] ~ double_exponential(0, 2);
}
"

#Uncomment the following lines -- launches rscript not compatible with cran check.
#sm <- stan_model(model_code = scode)
#fit1 <- stanWplot(object = sm, iter = 100000, chains=2, cores=1)
```

`stan_checkdivergences` *Analyse divergences in a stanfit object*

Description

Analyse divergences in a stanfit object

Usage

```
stan_checkdivergences(sf, nupars = "all")
```

Arguments

<code>sf</code>	stanfit object.
<code>nupars</code>	either the string 'all', or an integer reflecting how many pars (from first to nupars) to use.

Value

A list of four matrices. `$location` and `$sd` contain the bivariate interactions of unconstrained parameters, sorted by either the relative location of any divergences, or the relative standard deviation. `$locationmeans` and `$sdmeans` collapse across the bivariate interactions to return the means for each parameter.

Examples

```

sunspots<-sunspot.year
sunspots<-sunspots[50: (length(sunspots) - (1988-1924))]
id <- 1
time <- 1749:1924
datalong <- cbind(id, time, sunspots)

#setup model
ssmodel <- ctModel(type='ct', n.latent=2, n.manifest=1,
  manifestNames='sunspots',
  latentNames=c('ss_level', 'ss_velocity'),
  LAMBDA=matrix(c( 1, 'ma1| log(1+(exp(param)))' ), nrow=1, ncol=2),
  DRIFT=matrix(c(0, 'a21 | -log(1+exp(param))', 1, 'a22'), nrow=2, ncol=2),
  MANIFESTMEANS=matrix(c('m1|param * 10 + 44'), nrow=1, ncol=1),
  MANIFESTVAR=diag(0,1), #As per original spec
  CINT=matrix(c(0, 0), nrow=2, ncol=1),
  DIFFUSION=matrix(c(0, 0, 0, "diffusion"), ncol=2, nrow=2))

#fit
ssfit <- ctStanFit(datalong, ssmodel, iter=2,
  optimize=FALSE, chains=1)

stan_checkdivergences(ssfit$stanfit$stanfit) #stan object

```

stan_reinitsf

Quickly initialise stanfit object from model and data

Description

Quickly initialise stanfit object from model and data

Usage

```
stan_reinitsf(model, data, fast = FALSE)
```

Arguments

model	stanmodel
data	standata
fast	Use cut down form for speed

Value

stanfit object

Examples

```
sf <- stan_reinitsf(ctstantestfit$stanmodel,ctstantestfit$standata)
```

 stan_unconstrainsamples

Convert samples from a stanfit object to the unconstrained scale

Description

Convert samples from a stanfit object to the unconstrained scale

Usage

```
stan_unconstrainsamples(fit, standata = NA)
```

Arguments

fit	stanfit object.
standata	only necessary if R session has been restarted since fitting model – used to reinitialize stanfit object.

Value

Matrix containing columns of unconstrained parameters for each post-warmup iteration.

Examples

```
#get data
sunspots<-sunspot.year
sunspots<-sunspots[50:(length(sunspots) - (1988-1924))]
id <- 1
time <- 1749:1924
datalong <- cbind(id, time, sunspots)

#setup model
ssmodel <- ctModel(type='ct', n.latent=2, n.manifest=1,
  manifestNames='sunspots',
  latentNames=c('ss_level', 'ss_velocity'),
  LAMBDA=matrix(c( 1, 'ma1| log(1+(exp(param)))' ), nrow=1, ncol=2),
  DRIFT=matrix(c(0, 'a21 | -log(1+exp(param))', 1, 'a22'), nrow=2, ncol=2),
  MANIFESTMEANS=matrix(c('m1|param * 10 + 44'), nrow=1, ncol=1),
  MANIFESTVAR=diag(0,1), #As per original spec
  CINT=matrix(c(0, 0), nrow=2, ncol=1),
  DIFFUSION=matrix(c(0, 0, 0, "diffusion"), ncol=2, nrow=2))

#fit
ssfit <- ctStanFit(datalong, ssmodel,
  iter=200, chains=2,optimize=FALSE, priors=TRUE,control=list(max_treedepth=4))
umat <- stan_unconstrainsamples(ssfit$stanfit$stanfit)
```

summary.ctStanFit *summary.ctStanFit*

Description

Summarise a ctStanFit object that was fit using [ctStanFit](#).

Usage

```
## S3 method for class 'ctStanFit'
summary(
  object,
  timeinterval = 1,
  digits = 4,
  parmatrices = TRUE,
  priorcheck = TRUE,
  residualcov = TRUE,
  ...
)
```

Arguments

object	fit object from ctStanFit , of class ctStanFit.
timeinterval	positive numeric indicating time interval to use for discrete time parameter calculations reported in summary.
digits	integer denoting number of digits to report.
parmatrices	if TRUE, also return additional parameter matrices – can be slow to compute for large models with many samples.
priorcheck	Whether or not to use <code>ctsem:::priorchecking</code> to compare posterior mean and sd to prior mean and sd.
residualcov	Whether or not to show standardised residual covariance. Takes a little longer to compute.
...	Additional arguments to pass to <code>ctsem:::priorcheckreport</code> , such as <code>meanlim</code> , or <code>sdlim</code> .

Value

List containing summary items.

Examples

```
summary(ctstantestfit)
```

test_isclose	<i>Tests if 2 values are close to each other</i>
--------------	--

Description

Tests if 2 values are close to each other

Usage

```
test_isclose(..., tol = 1e-08)
```

Arguments

...	values to compare
tol	tolerance

Value

Logical or testthat output.

Examples

```
test_isclose(1, 1.000001, tol=1e-4)
```

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