

# Package ‘glmmBUGS’

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**Type** Package

**Title** Generalised Linear Mixed Models with BUGS and JAGS

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**Depends** R (>= 3.0.0)

**Imports** abind, methods, MASS, sp, graphics, stats

**Suggests** spdep

**Enhances** nlme, diseasemapping, geoRglm, R2WinBUGS, R2OpenBUGS, BRugs,  
R2jags, rjags

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**Description** Automates running Generalized Linear Mixed Models, including  
spatial models, with WinBUGS, OpenBUGS and JAGS. Models are specified with  
formulas, with the package writings model files, arranging unbalanced data  
in ragged arrays, and creating starting values. The model is re-parameterized,  
and functions are provided for converting model outputs to the original  
parameterization.

**SystemRequirements** Compiling vignette requires JAGS  
(<http://mcmc-jags.sourceforge.net>)

**License** GPL

**RoxygenNote** 5.0.1

**NeedsCompilation** no

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<b>addSpatial</b>	<i>Calculate adjacency values for WinBUGS</i>
-------------------	---

---

**Description**

Put an adjacency object in a ragged array

**Usage**

```
addSpatial(map, raggedArray, effect = NULL, prefix=NULL)
```

**Arguments**

<b>map</b>	a spatialPolygonsDataFrame object, or an nb object or a list of two vectors, adj and num
<b>raggedArray</b>	the result from <code>winBugsRaggedArray</code>
<b>effect</b>	a character vector listing the effect names
<b>prefix</b>	Character string to be appended to variable names

**Details**

Computes the values need by the `car.normal` distribution in WinBUGS. This function is called by `glmmBUGS` when a spatial argument is provided, `addSpatial` is usually not called by a user.

**Value**

The ragged array is returned, with the following additional elements

num	a vector of the number of neighbours of each region
adj	a vector containing the neighbours
weights	a vector of ones, the same length as adj
NregionSpatial	where 'region' is replaced by the name of the effect. The number of regions.

**Author(s)**

Patrick Brown

**References**

Also see the geoBUGS manual

**Examples**

```
## Not run:

# get a winbugs model and data ready, without a spatial effect
data(ontario)

forBugs = glmmBUGS(formula=observed + logExpected ~ 1,
  effects="CSDUID",   family="poisson",
  data=data.frame(ontario))

# now add a spatial effect.
# first, compute the adjacency matrix
# if region ID's are stored as factors, make sure to convert
# them to characters rather than the default of converting them
# to integers
library(diseasemapping)
data(popdata)
popDataAdjMat = poly2nb(popdata, row.names=as.character(popdata[["CSDUID"]]) )

data(popDataAdjMat)

# add the adjacency matrix to the ragged array
raggedWithSpatial = addSpatial(popDataAdjMat, forBugs$ragged, "CSDUID")

# write a new bugs model with a spatial effect
writeBugsModel("model.bug", "CSDUID", NULL, c("count", "expected"),
  "poisson", spatial="CSDUID")
startingValues = forBugs$startingValues
source("getInits.R")

library(R2WinBUGS)
popResult = bugs(raggedWithSpatial, getInits,
  parameters.to.save = names(getInits()), model.file="model.bug",
```

```
n.chain=3, n.iter=1000, n.burnin=100, n.thin=10, program="winbugs")
## End(Not run)
```

**binToBinom***Convert Bernoulli observations to Binomial***Description**

Combines multiple Bernoulli observations with the same covariates into one Binomial response

**Usage**

```
binToBinom(obs, covariates)
```

**Arguments**

obs	logical vector of observations
covariates	Data frame or matrix of covariates

**Value**

A data frame with one row for each unique value for the covariates, including the covariates and the following additional columns:

y	Number of positive observations for the corresponding covariate values
N	Total number of observations for these covariates

**Author(s)**

Patrick Brown

**Examples**

```
thedata = data.frame(sex = rep(c("m", "f"), 10), age=rep(c(20,30), c(10, 10)))
y = rbinom(dim(thedata)[1], 1, 0.5)
bindata = binToBinom(y, thedata)
bindata$zeros = bindata$N - bindata$y
glm(as.matrix(bindata[,c("y", "zeros")]) ~ sex, data=bindata, family=binomial)
```

---

**checkChain***Plot an MCMC run*

---

## Description

Makes time series plots of the parameters (not the random effects) of an MCMC run.

## Usage

```
checkChain(chain, parameters=NULL, oneFigure=TRUE)
```

## Arguments

- |            |   |
|------------|---|
| chain      | The result from <a href="#">restoreParams</a> , or the sims.array component of a <a href="#">bugs</a> call.   |
| parameters | Vector of character strings giving names of parameters to plot. Default is all parameters with names starting with either "beta", "intercept", or "SD". |
| oneFigure  | if TRUE, use <code>par(mfrow=c(a,b))</code> to put all plots on the same device. Otherwise create a new device for each plot.                           |

## Value

Plots are produced, nothing is returned

## Author(s)

Patrick Brown

## See Also

[restoreParams](#), [summaryChain](#)

## Examples

```
thechain = list(beta = array(1, c(10, 3,4),
dimnames = list(NULL, NULL, paste("beta[", 1:4, "]", sep=""))),
intercept = matrix(1, 10, 3))

checkChain(thechain)
```

**cholInvArray***Precision matrices to variance matrices for Winbugs output***Description**

Given an array containing simulations from the posterior of a precision matrix, each individual precision matrix is converted to variances, covariances, and correlations.

**Usage**

```
cholInvArray(x, prefix = "T", chol=FALSE)
```

**Arguments**

- |        |  |
|--------|--|
| x      | An array of winbugs output, with precision matrix entries of the form "T[1,3]" |
| prefix | The name of the precision matrix in winbugs, the "T" in "T[1,2 ]"              |
| chol   | If TRUE, the cholesky decomposition is returned instead of the inverse         |

**Details**

Inverts the matrices with the cholesky decomposition, but operating on all matrices simultaneously using array arithmetic.

**Value**

An array with the third dimension's precision matrix entries changed to

- "sdT[i,i]" for the standard deviation of component i
- "covT[i,j]" for the covariance between i and j
- "corrT[i,j]" for the correlations between i and j

**Examples**

```
# create a random positive definite matrix by
# generating a lower triangle
N=4
lmat = diag(runif(N, 1, 10))
thetri = lower.tri(lmat)
lmat[thetri] = rnorm(sum(thetri), 0, 2)
# precmat = solve(lmat %*% t(lmat))
precmat = solve(lmat %*% t(lmat))

# put this matrix into an array
precarray = array(c(precmat), dim=c(1,1,length(precmat)))
dimnames(precarray) = list(NULL, NULL,
  paste("T[", rep(1:N, N), ", ", rep(1:N, rep(N,N)), "]"),sep=""))
# invert it with cholInvArray and the solve function
```

```
cholInvArray(precarray)[1,1,]
# the off diagonals of solve(precmat) should be
# the covT elements of cholInvArray(precarray)
solve(precmat)
# the standard deviations in cholInvArray(precarray) should be the
# root of the diagonals of solve(precmat)
sqrt(diag(solve(precmat)))
```

---

**getDesignMatrix***Computes a design matrix from factors and interactions*

---

## Description

Converts all factors and interactions to indicator variables, suitable for passing to WinBUGS.

## Usage

```
getDesignMatrix(formula, data, effects = NULL)
```

## Arguments

formula	A formula object specifying the fixed effects for the model
data	A data frame containing the covariates and factors for random effects
effects	A vector of character strings containing the grouping levels, from most general to most specific

## Details

The most populous level of a factor is made the baseline.

## Value

A matrix containing the covariates, the response(s), and the random effect factors. Also attributes

covariates	A list giving the covariates which apply at each level, suitable for passing to <a href="#">winBugsRaggedArray</a>
response	A vector of character strings giving the responses

## Author(s)

Patrick Brown

## See Also

[winBugsRaggedArray](#), [glmmBUGS](#)

## Examples

```
library(nlme)
data(Muscle)
muscleDesign = getDesignMatrix(conc ~ length, data=Muscle, effects="Strip" )
attributes(muscleDesign)$covariates
attributes(muscleDesign)$response
```

`getRaggedSeq`

*Get one sequence for a ragged array*

## Description

This function is called by [winBugsRaggedArray](#)

### Usage

```
getRaggedSeq(data)
```

### Arguments

data	a data frame with two columns
------	-------------------------------

### Value

The ragged sequence

### Author(s)

Patrick Brown <patrick.brown@utoronto.ca>

### See Also

[winBugsRaggedArray](#)

`getStartingValues`

*Extract starting values for an MCMC chain from glmmPQL results*

## Description

Parameter estimates and random effect predictions are extracted from a glmmPQL model fit, and formatted to correspond to the levels in the supplied ragged array.

### Usage

```
getStartingValues(pql, ragged, prefix=NULL, reparam=NULL)
```

## Arguments

pql	output from the <code>glmmPQLstrings</code> function
ragged	a ragged array, from <code>winBugsRaggedArray</code>
prefix	string to append to object names
reparam	vector of random effect names, subtract covariates at this level from the intercept.

## Details

This function produces a list suitable for passing to `startingFunction` to generate random starting values for use with `bugs`. If `ragged` has a spatial component, starting values for a spatial random effect will also be computed.

## Value

A list of vectors, one for each set of parameters or random effects, and a list of estimated standard deviations.

## Author(s)

Patrick Brown <patrick.brown@utoronto.ca>

## See Also

`glmmPQLstrings`, `startingFunction`, `bugs`, `glmmBUGS`

## Description

Creates ragged arrays, writes a model file, and generates sensible starting estimates.

## Usage

```
glmmBUGS(formula, data, effects, modelFile = "model.txt",
initFile = "getInits.R",
family = c("bernoulli", "binomial", "poisson", "gaussian"),
spatial=NULL, spatialEffect = NULL,
reparam=NULL, prefix=NULL, priors=NULL,
brugs=length(grep("unix|linux",
.Platform$OS.type,
ignore.case=TRUE)))
```

## Arguments

formula	A formula for the fixed effects portion of the model
data	A data frame containing the response, covariates, and group membership
effects	A vector of character strings containing the grouping levels, from most general to most specific
modelFile	File for saving the bugs model
initFile	File for saving the function for generating initial values
family	distribution of responses
spatial	For Markov Random Field models, a polygons or adjacency matrix. For Geostatistical models, a SpatialPoints objects, a matrix or data frame with columns "x" and "y", or a vector of complex numbers.
spatialEffect	spatial variable from data
reparam	vector of random effect names, subtract covariates at this level from the intercept.
prefix	string to append to object names
priors	List or vector where names refer to parameters and elements are prior distributions, for example <code>list(SDsite="dunif(0,10)")</code> .
brugs	compatiblity with OpenBUGS, using the inprod function in place of inprod2, defaults to FALSE on windows and TRUE on unix platforms.

## Details

Consider the following model, where  $Y_{ijk}$  is the number of absences from individual k from class j in school k.

$$\begin{aligned}
 Y_{ijk} &\sim Poisson(\mu_i) \\
 \log(\mu_i) &= \delta age_{ijk}\beta + classSize_{ij}\alpha + schoolCategory_i\gamma + U_i + V_{ij} \\
 U_i &\sim N(0, \sigma^2) \\
 V_{ij} &\sim N(0, \nu^2)
 \end{aligned}$$

Here there are covariates which apply to each of the three levels, and random effects at the school and class level. If data is a data frame with one line per individual, the following would implement this model:

```
glmmBUGS(data, effects=c("school", "class"), covariates = list(school="schoolCategory", class="class", observations = "absences"), family="poisson")
```

To aid in convergence, the bugs model is actually the following:

$$\log(\mu_i) = age_{ijk}\beta + V_{ij}$$

$$V_{ij} \sim N(U_i + classSize_{ij}\alpha, \nu^2)$$

$$U_i \sim N(\delta + schoolCategory_i\gamma, \sigma^2)$$

and the function `restoreParams` subtracts the means from the random effects to restore the original set of equations.

glmmBUGS calls the following functions:

`getDesignMatrix` to convert factors and interactions to indicator variables and find which covariates apply at which levels

`winBugsRaggedArray` to prepare the ragged array

`glmmPQLstrings` estimate starting values

`writeBugsModel` to create a model file

`getStartingValues` to extract starting values from the glmmPQL result

`startingFunction` to write a function to generate random starting values

Type `glmmBUGS` on the R command line to see the source code, it provides a good summary of the roles of the various functions in the `glmmBUGS` package.

## Value

Returns a list with the ragged array, from `winBugsRaggedArray`, and the list of starting values from `getStartingValues`. Writes a model file and an initial value function. Note that the initial value function in `initFile` will look for an object called `startingValues`, which does not exist as this is part of a list. Either create `startingValues <- result$startingValues` or edit `initFile`.

## Warning

You are strongly encouraged to modify the model file and the initial value function file prior to using them.

## Note

glmmBUGS uses the `inprod2` function, which isn't implemented in OpenBUGS, the model file will have to be modified for use with OpenBUGS.

## Author(s)

Patrick Brown, <patrick.brown@utoronto.ca>

## References

"Handling unbalanced datasets" in the "Tricks: Advanced Use of the BUGS Language" section of the bugs manual, at <http://www.openbugs.net/Manuals/Tricks.html>

## See Also

`winBugsRaggedArray`, `glmmPQLstrings`, `writeBugsModel`, `getStartingValues`, `startingFunction`, `bugs`

## Examples

```

library(nlme)
data(Muscle)

muscleRagged = glmmBUGS(conc ~ length, data=Muscle,
effects="Strip", family="gaussian",
modelFile='model.bug', reparam='Strip')
startingValues = muscleRagged$startingValues

## Not run:
# run with winbugs
source("getInits.R")
require('R2WinBUGS')
muscleResult = bugs(muscleRagged$ragged, getInits,
parameters.to.save = names(getInits()),
model.file="model.bug", n.chain=3, n.iter=1000,
n.burnin=100, n.thin=10, program="winbugs",
working.directory=getwd()
)

# a jags example
require('R2jags')
muscleResultJags = jags(
  muscleRagged$ragged, getInits, parameters.to.save = names(getInits()),
  model.file="model.bug", n.chain=3, n.iter=1000,
  n.burnin=100, n.thin=10,
  working.directory=getwd())

muscleParamsJags = restoreParams(
  muscleResultJags$BUGSoutput,
  muscleRagged$ragged)
checkChain(muscleParamsJags)

## End(Not run)

data(muscleResult)

muscleParams = restoreParams(muscleResult, muscleRagged$ragged)
summaryChain(muscleParams)
checkChain(muscleParams)

# a spatial example
## Not run:
library(diseasemapping)

data('popdata')
data('casedata')

```

```

model = getRates(casedata, popdata, ~age*sex)
ontario = getSMR(popdata, model, casedata)
ontario = ontario@data[,c("CSDUID", "observed", "logExpected")]

popDataAdjMat = spdep::poly2nb(popdata,
row.names=as.character(popdata[["CSDUID"]])))

data('popDataAdjMat')
data('ontario')

forBugs = glmmBUGS(formula=observed + logExpected ~ 1,
effects="CSDUID", family="poisson", spatial=popDataAdjMat,
spatialEffect="CSDUID",
data=ontario)

startingValues = forBugs$startingValues

source("getInits.R")
# find patrick's OpenBUGS executable file
if(Sys.info()['user'] == 'patrick') {
  obExec = system(
    "find /store/patrick/ -name OpenBUGS",
    TRUE)
  obExec = obExec[length(obExec)]
} else {
  obExec = NULL
}

bugsResult = bugs(forBugs$ragged, getInits,
parameters.to.save = names(getInits()),
model.file="model.bug", n.chain=3, n.iter=50, n.burnin=10,
n.thin=2,
program="winbugs", debug=T,working.directory=getwd())

data('ontarioResult')

ontarioParams = restoreParams(ontarioResult, forBugs$ragged)

ontarioSummary = summaryChain(ontarioParams)

# posterior probability of having 10x excess risk
postProb = apply(ontarioParams$FittedRCSDUID, 3, function(x) mean(x>log(10)) )
hist(postProb)

ontario = mergeBugsData(popdata, ontarioSummary)

spplot(ontario, "FittedRateCSDUID.mean")

ontario = mergeBugsData(ontario, postProb, newcol="postProb", by.x="CSDUID")
spplot(ontario, "postProb")

```

```

## End(Not run)

# geostatistical example

## Not run:
library(geoRglm)
data(rongelap)

rongelap= read.table(url(
  paste("http://www.leg.ufpr.br/lib/exe/fetch.php/",
  "pessoais:paulojus:mbgbook:datasets:rongelap.txt",
  sep=""))
),header=TRUE
)

library('spdep')
coordinates(rongelap) = ~cX+cY

rongelap$logOffset = log(rongelap$time)
rongelap$site = seq(1, length(rongelap$time))

forBugs = glmmBUGS(
  formula=counts + logOffset ~ 1, family="poisson",
  data=rongelap@data, effects="site", spatial=rongelap,
  priors=list(phiSite="dgamma(100,1)"))

startingValues = forBugs$startingValues
startingValues$phi$site = 100

source("getInits.R")

rongelapResult = bugs(forBugs$ragged, getInits,
  parameters.to.save = names(getInits()),
  model.file="model.bug", n.chain=2, n.iter=20, n.burnin=4, n.thin=2,
  program="winbugs", debug=TRUE,
  working.directory=getwd())

data('rongelapResult')

rongelapParams = restoreParams(rongelapResult, forBugs$ragged)

checkChain(rongelapParams)

rongelapParams$siteGrid = CondSimuPosterior(rongelapParams, rongelap,
  gridSize=100)

rongelapSummary=summaryChain(rongelapParams)

```

```
# plot posterior probabilities of being above average
image(rongelapSummary$siteGrid$pgt0)

## End(Not run)
```

**glmmPQLstrings***An alternat interface to glmmPQL***Description**

Calls `glmmPQL` in the MASS library, with the model being specified in the same manner as `writeBugsModel`

**Usage**

```
glmmPQLstrings(effects, covariates, observations, data = NULL,
family=c("bernoulli", "binomial", "poisson", "gaussian"), ...)
```

**Arguments**

<code>effects</code>	A vector of character strings containing the grouping levels, from most general to most specific
<code>covariates</code>	A list with names corresponding to effects and each element being a vector of covariates applicable at that level
<code>observations</code>	A character string giving the column of observations, or a vector where the first element is the observations and the remaning are offsets. For binomial responses, the first element is the counts (of successes), and the second element is the total number of trials. Note this differs from <code>glmmPQL</code> and <code>glm</code> 's notation, but is consistent with WinBUGS.
<code>data</code>	A data frame containing the response, covariates, and group membership.
<code>family</code>	The distribution to use. Either using <code>glmmPQL</code> 's specifications or <code>writeBugsModel</code> further arguments to <code>glmmPQL</code>
<code>...</code>	

**Details**

This function is useful for generating starting values for an MCMC chain.

**Value**

In addition to the output from `glmmPQL`, the following are returned

`effects, covariates, observations`  
As input

**Author(s)**

Patrick Brown, patrick.brown@utoronto.ca

**See Also**

[getStartingValues](#), [glmmPQL](#)

**Examples**

```
library(nlme)
data(Muscle)
glmmPQLstrings(effects="Strip", observations="conc",
  covariates=list(observations="length") ,
  data=Muscle, family="gaussian")
```

**mergeBugsData-methods** *Merge results from BUGS into a data.frame or SPDF*

**Description**

merge the result from bugs function

**Usage**

```
## S4 method for signature 'data.frame'
mergeBugsData(
  x, bugsSummary, by.x = NULL, newcol = "mean", ...
)
## S4 method for signature 'SpatialPolygonsDataFrame'
mergeBugsData(
  x, bugsSummary, by.x = NULL, newcol = "mean", ...
)
```

**Arguments**

x	spatial polygon object i.e population data set (popdata)
bugsSummary	posterior distribution result from summaryChain function
by.x	the common term from the spatial polygon object and the bugs function result
newcol	the summary statistic that to be merged back to the data frame
...	additional arguments

**Author(s)**

Patrick Brown

**Examples**

```
if(require('diseasemapping')){
  data('popdata')
  newdata = c("3560102"=2, "3560104"=3)
  popdatatry = mergeBugsData(popdata, newdata, by.x="CSDUID")
}
```

---

muscleResult	<i>data set contains muscle result</i>
--------------	--

---

**Description**

Results from running the muscle example in [glmmBUGS](#).

**Usage**

```
data(muscleResult)
```

**Format**

A list as returned by the [bugs](#) function.

**Details**

See [glmmBUGS](#) and [Muscle](#)

**Examples**

```
data(muscleResult)
```

---

ontario	<i>Ontario data on molar cancer</i>
---------	-------------------------------------

---

**Description**

Data frame showing expected and observed counts of molar cancer in Ontario

**Usage**

```
data(ontario)
```

**Format**

A data frame with 585 observations on the following 3 variables.

CSDUID factor of Ontario census subdivision ID numbers

observed Observed molar cancer cases

logExpected expected cases

**Details**

See the documentation for [glmmBUGS](#) for how this was created.

### Examples

```
data(ontario)
head(ontario)
```

ontarioResult	<i>Ontario Winbugs Results</i>
---------------	--------------------------------

### Description

Results from running Winbugs on the ontario data

### Usage

```
data(ontarioResult)
```

### Format

A list, as produced by the [bugs](#) function.

### Examples

```
data(ontarioResult)

ontarioParams = restoreParams(ontarioResult)

ontarioSummary = summaryChain(ontarioParams)
```

popDataAdjMat	<i>Data set containing an adjacency matrix</i>
---------------	--

### Description

The `popDataAdjMat` Data set contains the adjacency matrix which calculated from the [poly2nb](#) function.

### Usage

```
data('popDataAdjMat')
```

### Details

It is a adjacency matrix denoting the neighbours of Ontario census subdivisions.

## Examples

```
## Not run:
library('diseasemapping')
data('popdata')
popDataAdjMat = spdep::poly2nb(popdata,
row.names=as.character(popdata$CSDUID))

## End(Not run)
data('popDataAdjMat')
summary(popDataAdjMat)
attributes(popDataAdjMat)$region.id[1:10]
```

`restoreParams`      *Reparametrise bugs output*

## Description

Undoes the parametrisation used in [writeBugsModel](#), and gives the original names to random effect levels.

## Usage

```
restoreParams(bugsResult, ragged = NULL, extraX=NULL)
```

## Arguments

<code>bugsResult</code>	Output from <a href="#">bugs</a> , using a ragged array generated by <a href="#">winBugsRaggedArray</a> and a model generated by <a href="#">writeBugsModel</a>
<code>ragged</code>	The ragged array used to call <a href="#">bugs</a>
<code>extraX</code>	Possible extra covariates for spatial regions with no data but do have predicted spatial effects.

## Value

A list where each element is a matrix or an array. The first dimension is the number of realisations, the second the number of chains, and for vector-valued parameters and random effects, the third dimension is the length of the parameter.

If the model contains a spatial component, the result will have list entries the following:

<code>Reffect</code>	The random effect. In the case of spatial models this is the sum of the spatial and non-spatial random effects U+V.
,	
<code>ReffectSpatial</code>	The spatial random effect for each region, if any
<code>FittedReffect</code>	The predicted values on the link scale, being the random effect plus intercept and effect of covariates.

**Note**

For spatial models, one fitted rate is computed for each region in the adjacency matrix, even though some of these regions may not have spatial or non-spatial random effects simulated in the bugs model. If a spatial random effect is missing (as happens with islands), a zero is added. If a non-spatial random effect is missing (as happens when a regions does not have data), a value is simulated unconditionally from each iteration's intercept and standard deviation for that effect. Note that this does not add on the effect of possible covariates for that region. This can be added via the extraX argument.

**Author(s)**

Patrick Brown patrick.brown@utoronto.ca

**See Also**

[bugs](#)

**rongelapUTM**

*Rongelap island data*

**Description**

A SpatialPointsDataFrame containing the Rongelap data, in a UTM projection.

**Usage**

```
data("rongelapUTM")
data("rongelapResult")
```

**Details**

These coordinates were obtained by translating and rotating the original Rongelap data until all the coordinates fit into the Rongelap border given by www.gadm.org. So they are not exact.

**Source**

See the help file for [rongelap](#), or <http://www.leg.ufpr.br/doku.php/pessoais:paulojus:mbgbook:datasets>

**Examples**

```
data("rongelapUTM")
if(require("sp", quietly=TRUE)){
  plot(rongelapUTM)
}

## Not run:
rongelapBorderLL = raster::getData("GADM",
```

```

country="MHL",level=0)
library("rgdal")
rongelapBorderUTM = spTransform(rongelapBorderLL,
CRS(proj4string(rongelapUTM)))
plot(rongelapBorderUTM, add=TRUE)

## End(Not run)

rongelapUTM$logOffset = log(rongelapUTM$time)
rongelapUTM$site = seq(1, length(rongelapUTM$time))

forBugs = glmmBUGS(
formula=count + logOffset ~ 1, family="poisson",
  data=rongelapUTM@data, effects="site",
  spatial=rongelapUTM,
  priors=list(phisite="dgamma(100,1)")
  )

startingValues = forBugs$startingValues
startingValues$phi=list(site = 100)

source("getInits.R")

## Not run:
rongelapResult = bugs(forBugs$ragged, getInits,
parameters.to.save = names(getInits()),
  model.file="model.bug", n.chain=2, n.iter=20, n.burnin=4, n.thin=2,
  program="winbugs", debug=TRUE,
  working.directory=getwd())
rongelapParams = restoreParams(rongelapResult, forBugs$ragged)

## End(Not run)

data("rongelapResult")
rongelapParams = restoreParams(rongelapResult)

checkChain(rongelapParams)

rongelapSummary=summaryChain(rongelapParams)

```

**Description**

The code for the resulting function is saved in a file, to be edited and sourced in before calling WinBUGS.

**Usage**

```
startingFunction(startingValues, file = "getInits.R")
```

**Arguments**

<code>startingValues</code>	list returned from <a href="#">getStartingValues</a>
<code>file</code>	character string giving the name of the file to write to

**Details**

Given a list containing initial estimates of parameters and random effects, a text file is produced containing code for a function to generate random starting values for use with the `bugs()` function. It is intended that the file produced be checked and edited prior to use.

**Value**

A file, with the name given by the 'file' argument, is written.

**Warning**

You are strongly encouraged to edit the file to ensure the result is sensible

**Author(s)**

Patrick Brown, [patrick.brown@utoronto.ca](mailto:patrick.brown@utoronto.ca)

**See Also**

[getStartingValues](#), [bugs](#)

**Examples**

```
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##--or do  help(data=index)  for the standard data sets.
sval = list(intercept=0, beta = 1:2, Rperson = rep(0, 5), vars=list(person=1))
startingFunction(sval)
```

---

summaryChain	<i>Compute mean, standard deviation, and quantiles of an MCMC run</i>
--------------	---

---

## Description

Computes summary statistics for each parameter.

## Usage

```
summaryChain(chain, probs = c(0.005, 0.025, 0.05, 0.5))
```

## Arguments

- |       |   |
|-------|---|
| chain | The result from <a href="#">restoreParams</a> , or the sims.array component of a <a href="#">bugs</a> call. |
| probs | Quantiles for the posterior credible interval   |

## Value

A list of matrices, with rows corresponding to summary statistics and columns to parameters.

- |                   |  |
|-------------------|--|
| scalar            | Matrix for the scalar parameters   |
| ...               | One matrix for each vector valued parameter  |
| FittedRateReffect | For spatial models only, summaries on the natural scale (exponential of FittedR-effect). |

## Author(s)

Patrick Brown

## See Also

[restoreParams](#)

## Examples

```
# create a simple chain
thechain = list(beta = array(1, c(10, 3,4),
dimnames = list(NULL, NULL, paste("beta[", 1:4, "]", sep=""))),
intercept = matrix(1, 10, 3))

summaryChain(thechain)
```

---

winBugsRaggedArray      *Ragged Arrays for multilevel models in BUGS*

---

## Description

Suitable for unbalanced data.

## Usage

```
winBugsRaggedArray(data, effects = names(data)[-length(names(data))],
covariates = NULL, observations = names(data)[length(names(data))],
returnData = FALSE,
prefix=NULL, reparam=FALSE)
```

## Arguments

<code>data</code>	A data frame containing the response, covariates, and group membership.
<code>effects</code>	A vector of character strings containing the grouping levels, from most general to most specific. Defaults to the column names of data, excluding the last column.
<code>covariates</code>	A list with names corresponding to effects and each element being a vector of covariates applicable at that level
<code>observations</code>	A character string giving the column of observations, or a vector where the first element is the observations and the remaining are offsets.
<code>returnData</code>	If true, returns the re-ordered data frame as well as the data frame
<code>prefix</code>	Character string to be appended to variable names
<code>reparam</code>	Vector of effect names, reparametrize the intercept by subtracting the mean of covariates at this level.

## Details

This function creates a list of data suitable for passing to the `bugs` function, suitable for implementation as a ragged array. The output can be passed to `getStartingValues` to manipulate the output from `glmmPQLstrings`, and to `restoreParams` to restore the original parametrisation from `bugs` output.

## Value

A list with the following components

<code>Nxx</code>	The number of levels in the most general grouping
<code>Syy</code>	Indexing sequences, one for each level. If <code>yy</code> is level n, level <code>n+1</code> has elements <code>Syy[1]</code> to <code>Syy[2]-1</code> belonging to the first category of level n.
<code>Xyy</code>	Matrix or vector of covariates belonging to level <code>yy</code> vector of observations.

**Author(s)**

Patrick Brown, <patrick.brown@utoronto.ca>

**References**

"Handling unbalanced datasets" in the "Tricks: Advanced Use of the BUGS Language" section of the bugs manual, at <http://www.openbugs.net/Manuals/Tricks.html>

**See Also**

[bugs](#)

**Examples**

```
library(nlme)
data(Muscle)
muscleRagged = winBugsRaggedArray(Muscle, effects="Strip",
observations="conc",
covariates=list(observations="length"))
```

---

writeBugsModel

*Write a bugs model file for a Generalised Linear Mixed Model*

---

**Description**

Given a list of effect groups, and the covariates associated with each level, a bugs model file is written using ragged arrays corresponding to output from [winBugsRaggedArray](#)

**Usage**

```
writeBugsModel(file, effects, covariates, observations,
family = c("bernoulli", "binomial", "poisson", "normal", "other"),
spatial = NULL, geostat=FALSE,
prefix = "", reparam=NULL, brugs=TRUE, priors=NULL)
```

**Arguments**

<code>file</code>	a character string denoting the name of the bugs model file written.
<code>effects</code>	vector of effect groups
<code>covariates</code>	A list with names corresponding to effects and each element being a vector of covariates applicable at that level
<code>observations</code>	A character string giving the column of observations, or a vector where the first element is the observations and the remaining are offsets.
<code>family</code>	Response distribution

<code>spatial</code>	name of the spatial random effect
<code>geostat</code>	Is this a geostatistical random effect? Defaults to FALSE for the Besag, York and Mollie discrete spatial variation model
<code>prefix</code>	the prefix
<code>reparam</code>	vector of random effect names, subtract covariates at this level from the intercept.
<code>brugs</code>	make the model file compatible with OpenBugs by using the <code>inprod</code> function in place of <code>inprod2</code>
<code>priors</code>	character string of prior distributions, with the name of each element referring to the parameter it is the prior for

## Details

The arguments to the function specify a generalised linear mixed model. A file containing code for a corresponding bugs model is written. The model uses ragged arrays to specify grouping factors, and includes covariates at the appropriate levels to aid in chain convergence. It is intended that the user will edit this file before its use. The prior distributions in particular may not be appropriate.

## Value

A file, suitable for passing to the `bugs` function in R2WinBUGS.

## Warning

You are strongly encouraged to modify the model file prior to using it.

## Author(s)

Patrick Brown, <patrick.brown@utoronto.ca>

## References

"Handling unbalanced datasets" in the "Tricks: Advanced Use of the BUGS Language" section of the bugs manual, at <http://www.openbugs.net/Manuals/Tricks.html>

## Examples

```
writeBugsModel("model.bug", effects="Strip", observations="conc",
               covariates=list(observations="length"),
               family="normal", priors=c(intercept="dunif(-10,10)") )

cat(scan("model.bug", "a", sep='\n'),sep='\n')
```

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