# Package ‘simstudy’ 

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Description Simulates data sets in order to explore modeling techniques or better understand data generating processes. The user specifies a set of relationships between covariates, and generates data based on these specifications. The final data sets can represent data from randomized control trials, repeated measure (longitudinal) designs, and cluster randomized trials. Missingness can be generated using various mechanisms (MCAR, MAR, NMAR).
License GPL-3
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BugReports https://github.com/kgoldfeld/simstudy/issues
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addColumns Add columns to existing data set

## Description

Add columns to existing data set

## Usage

addColumns(dtDefs, dtOld, envir = parent.frame())

## Arguments

dtDefs name of definitions for added columns
dt0ld name of data table that is to be updated
envir Environment the data definitions are evaluated in. Defaults to base::parent.frame.

## Value

an updated data.table that contains the added simulated data

## Examples

```
    # New data set
    def <- defData(varname = "xNr", dist = "nonrandom", formula = 7, id = "idnum")
    def <- defData(def, varname = "xUni", dist = "uniform", formula = "10;20")
    dt <- genData(10, def)
    # Add columns to dt
    def2 <- defDataAdd(varname = "y1", formula = 10, variance = 3)
    def2 <- defDataAdd(def2, varname = "y2", formula = .5, dist = "binary")
    def2
    dt <- addColumns(def2, dt)
    dt
```

    addCompRisk Generating single competing risk survival variable
    
## Description

Generating single competing risk survival variable

## Usage

addCompRisk( dtName, events, timeName, censorName = NULL, eventName = "event",
typeName = "type",
keepEvents = FALSE,
idName = "id"
)

## Arguments

| dtName | Name of complete data set to be updated |
| :--- | :--- |
| events | Vector of column names that include time-to-event outcome measures |
| timeName | A string to indicate the name of the combined competing risk time-to-event out- <br> come that reflects the minimum observed value of all time-to-event outcomes. |
| censorName | The name of a time-to-event variable that is the censoring variable. Must be one <br> of the "events" names. Defaults to NULL. |


| eventName | The name of the new numeric/integer column representing the competing event <br> outcomes. If censorName is specified, the integer value for that event will be 0. <br> Defaults to "event", but will be ignored if timeName is NULL. |
| :--- | :--- |
| typeName | The name of the new character column that will indicate the event type. The <br> type will be the unique variable names in survDefs. Defaults to "type", but will <br> be ignored if timeName is NULL. |
| keepEvents | Indicator to retain original "events" columns. Defaults to FALSE. |
| idName | Name of id field in existing data set. |

## Value

An updated data table

## Examples

```
d1 <- defData(varname = "x1", formula = .5, dist = "binary")
d1 <- defData(d1, "x2", .5, dist = "binary")
dS <- defSurv(varname = "reinc", formula = "-10 - 0.6*x1 + 0.4*x2", shape = 0.3)
dS <- defSurv(dS, "death", "-6.5 + 0.3*x1 - 0.5*x2", shape = 0.5)
dS <- defSurv(dS, "censor", "-7", shape = 0.55)
dd <- genData(10, d1)
dd <- genSurv(dd, dS)
addCompRisk(dd, c("reinc","death", "censor"), timeName = "time",
    censorName = "censor", keepEvents = FALSE)
```


## Description

Add a single column to existing data set based on a condition

## Usage

addCondition(condDefs, dtOld, newvar, envir = parent.frame())

## Arguments

| condDefs | Name of definitions for added column |
| :--- | :--- |
| dt0ld | Name of data table that is to be updated |
| newvar | Name of new column to add |
| envir | Environment the data definitions are evaluated in. Defaults to base::parent.frame. |

## Value

An updated data.table that contains the added simulated data

## Examples

```
# New data set
    def <- defData(varname = "x", dist = "categorical", formula = ".33;.33")
    def <- defData(def, varname = "y", dist = "uniform", formula = "-5;5")
    dt <- genData(1000, def)
    # Define conditions
    defC <- defCondition(
        condition = "x == 1", formula = "5 + 2*y-. 5*y^2",
        variance = 1, dist = "normal"
    )
    defC <- defCondition(defC,
        condition = "x == 2",
        formula = "3 - 3*y + y^2", variance = 2, dist = "normal"
    )
    defC <- defCondition(defC,
        condition = "x == 3",
        formula = "abs(y)", dist = "poisson"
    )
    # Add column
    dt <- addCondition(defC, dt, "NewVar")
    # Plot data
    library(ggplot2)
    ggplot(data = dt, aes(x = y, y = NewVar, group = x)) +
    geom_point(aes(color = factor(x)))
```

    addCorData
    
## Description

Add correlated data to existing data.table

## Usage

```
    addCorData(
        dtOld,
        idname,
        mu,
        sigma,
        corMatrix = NULL,
        rho,
        corstr = "ind",
        cnames = NULL
    )
```


## Arguments

| dt0ld | Data table that is the new columns will be appended to. |
| :--- | :--- |
| idname | Character name of id field, defaults to "id". |
| mu | A vector of means. The length of mu must be nvars. |
| sigma | Standard deviation of variables. If standard deviation differs for each variable, <br> enter as a vector with the same length as the mean vector mu. If the standard <br> deviation is constant across variables, as single value can be entered. |
| corMatrix | Correlation matrix can be entered directly. It must be symmetrical and posi- <br> tive semi-definite. It is not a required field, if a matrix is not provided, then a <br> structure and correlation coefficient rho must be specified. |
| rho | Correlation coefficient, -1 <= rho <= 1. Use if corMatrix is not provided. |
| corstr | Correlation structure of the variance-covariance matrix defined by sigma and <br> rho. Options include "ind" for an independence structure, "cs" for a compound <br> symmetry structure, and "ar1" for an autoregressive structure. |
| cnames | Explicit column names. A single string with names separated by commas. If no <br> string is provided, the default names will be V\#, where \# represents the column. |

## Value

The original data table with the additional correlated columns

## Examples

```
def <- defData(varname = "xUni", dist = "uniform", formula = "10;20", id = "myID")
def <- defData(def,
    varname = "xNorm", formula = "xUni * 2", dist = "normal",
    variance = 8
)
dt <- genData(250, def)
mu <- c(3, 8, 15)
sigma <- c(1, 2, 3)
```

```
dtAdd <- addCorData(dt, "myID",
    mu = mu, sigma = sigma,
    rho = .7, corstr = "cs"
)
dtAdd
    round(var(dtAdd[, .(V1, V2, V3)]), 3)
    round(cor(dtAdd[, .(V1, V2, V3)]), 2)
    dtAdd <- addCorData(dt, "myID",
        mu = mu, sigma = sigma,
        rho = .7, corstr = "ar1"
)
round(cor(dtAdd[, .(V1, V2, V3)]), 2)
corMat <- matrix(c(1, .2, .8, .2, 1, .6, .8, .6, 1), nrow = 3)
dtAdd <- addCorData(dt, "myID",
        mu = mu, sigma = sigma,
        corMatrix = corMat
)
round(cor(dtAdd[, .(V1, V2, V3)]), 2)
```

    addCorFlex
    
## Description

Create multivariate (correlated) data - for general distributions

## Usage

addCorFlex (
$d t$,
defs,
rho $=0$,
tau $=$ NULL,
corstr = "cs",
corMatrix = NULL,
envir = parent.frame()
)

## Arguments

dt
Data table that will be updated.
defs
Field definition table created by function defDataAdd.
rho
Correlation coefficient, $-1<=$ rho $<=1$. Use if corMatrix is not provided.

| tau | Correlation based on Kendall's tau. If tau is specified, then it is used as the <br> correlation even if rho is specified. If tau is NULL, then the specified value of <br> rho is used, or rho defaults to 0. |
| :--- | :--- |
| corstr | Correlation structure of the variance-covariance matrix defined by sigma and <br> rho. Options include "cs" for a compound symmetry structure and "ar1" for an <br> autoregressive structure. Defaults to "cs". |
| corMatrix | Correlation matrix can be entered directly. It must be symmetrical and posi- <br> tive semi-definite. It is not a required field; if a matrix is not provided, then a <br> structure and correlation coefficient rho must be specified. |
| envir | Environment the data definitions are evaluated in. Defaults to base::parent.frame. |

## Value

data.table with added column(s) of correlated data

## Examples

```
defC <- defData(
    varname = "nInds", formula = 50, dist = "noZeroPoisson",
    id = "idClust"
)
dc <- genData(10, defC)
#### Normal only
dc <- addCorData(dc,
    mu = c(0, 0, 0, 0), sigma = c(2, 2, 2, 2), rho = .2,
    corstr = "cs", cnames = c("a", "b", "c", "d"),
    idname = "idClust"
)
di <- genCluster(dc, "idClust", "nInds", "id")
defI <- defDataAdd(
    varname = "A", formula = "-1 + a", variance = 3,
    dist = "normal"
)
defI <- defDataAdd(defI,
    varname = "B", formula = "4.5 + b", variance = .5,
    dist = "normal"
)
defI <- defDataAdd(defI,
    varname = "C", formula = "5*c", variance = 3,
    dist = "normal"
)
defI <- defDataAdd(defI,
    varname = "D", formula = "1.6 + d", variance = 1,
    dist = "normal"
)
#### Generate new data
```

```
    di <- addCorFlex(di, defI, rho = 0.4, corstr = "cs")
```

    \# Check correlations by cluster
    for (i in 1:nrow(dc)) \{
    print(cor(di[idClust == i, list(A, B, C, D)]))
    \}
    \# Check global correlations - should not be as correlated
    \(\operatorname{cor}(d i[, \operatorname{list}(A, B, C, D)])\)
    addCorGen

Create multivariate (correlated) data - for general distributions

## Description

Create multivariate (correlated) data - for general distributions

## Usage

```
addCorGen(
    dtOld,
    nvars = NULL,
    idvar = "id",
    rho = NULL,
    corstr = NULL,
    corMatrix = NULL,
    dist,
    param1,
    param2 = NULL,
    cnames = NULL,
    method = "copula",
)
```


## Arguments

dt0ld The data set that will be augmented. If the data set includes a single record per id, the new data table will be created as a "wide" data set. If the original data set includes multiple records per id, the new data set will be in "long" format.
nvars The number of new variables to create for each id. This is only applicable when the data are generated from a data set that includes one record per id.
idvar String variable name of column represents individual level id for correlated data.
rho Correlation coefficient, $-1<=$ rho $<=1$. Use if corMatrix is not provided.
corstr Correlation structure of the variance-covariance matrix defined by sigma and rho. Options include "cs" for a compound symmetry structure and "ar1" for an autoregressive structure.

| corMatrix | Correlation matrix can be entered directly. It must be symmetrical and posi- <br> tive semi-definite. It is not a required field; if a matrix is not provided, then a <br> structure and correlation coefficient rho must be specified. |
| :--- | :--- |
| dist | A string indicating "normal", "binary", "poisson" or "gamma". |
| param1 | A string that represents the column in dtOld that contains the parameter for the <br> mean of the distribution. In the case of the uniform distribution the column <br> specifies the minimum. |
| param2 | A string that represents the column in dtOld that contains a possible second <br> parameter for the distribution. For the normal distribution, this will be the vari- <br> ance; for the gamma distribution, this will be the dispersion; and for the uniform <br> distribution, this will be the maximum. |
| cnames | Explicit column names. A single string with names separated by commas. If no <br> string is provided, the default names will be V\#, where \# represents the column. |
| method | Two methods are available to generate correlated data. (1) "copula" uses the <br> multivariate Gaussian copula method that is applied to all other distributions; <br> this applies to all available distributions. (2) "ep" uses an algorithm developed <br> by Emrich and Piedmonte (1991). |
| I. | May include additional arguments that have been deprecated and are no longer <br> used. |

## Value

Original data.table with added column(s) of correlated data

## References

Emrich LJ, Piedmonte MR. A Method for Generating High-Dimensional Multivariate Binary Variates. The American Statistician 1991;45:302-4.

## Examples

```
# Wide example
def <- defData(varname = "xbase", formula = 5, variance = .4, dist = "gamma", id = "cid")
def <- defData(def, varname = "lambda", formula = ". 5 + . 1*xbase", dist = "nonrandom", link = "log")
dt <- genData(100, def)
addCorGen(
    dtOld = dt, idvar = "cid", nvars = 3, rho = .7, corstr = "cs",
    dist = "poisson", param1 = "lambda"
)
# Long example
def <- defData(varname = "xbase", formula = 5, variance = .4, dist = "gamma", id = "cid")
def2 <- defDataAdd(
    varname = "p", formula = "-3+.2*period + .3*xbase",
```

```
    dist = "nonrandom", link = "logit"
)
dt <- genData(100, def)
dtLong <- addPeriods(dt, idvars = "cid", nPeriods = 3)
dtLong <- addColumns(def2, dtLong)
addCorGen(
    dtOld = dtLong, idvar = "cid", nvars = NULL, rho = .7, corstr = "cs",
    dist = "binary", param1 = "p"
)
```

addMarkov

Add Markov chain

## Description

Generate a Markov chain for n individuals or units by specifying a transition matrix.

## Usage

addMarkov(
dd,
transMat,
chainLen, wide = FALSE, id = "id",
pername = "period",
varname = "state",
widePrefix = "S",
start0lab = NULL,
trimvalue $=$ NULL
)

## Arguments

dd
transMat Square transition matrix where the sum of each row must equal 1. The dimensions of the matrix equal the number of possible states.
chainLen Length of each chain that will be generated for each chain; minimum chain length is 2.
wide Logical variable (TRUE or FALSE) indicating whether the resulting data table should be returned in wide or long format. The wide format includes all elements of a chain on a single row; the long format includes each element of a chain in its own row. The default is wide $=$ FALSE, so the long format is returned by default.

| id | Character string that represents name of "id" field. Defaults to "id". |
| :--- | :--- |
| pername | Character string that represents the variable name of the chain sequence in the <br> long format. Defaults "period", |
| varname | Character string that represents the variable name of the state in the long format. <br> Defaults to "state". <br> Character string that represents the variable name prefix for the state fields in <br> the wide format. Defaults to "S". |
| widePrefix | Character string that represents name of the integer field containing starting state <br> (State 0) of the chain for each individual. If it is NULL, starting state defaults to |
| start0lab | 1. Default is NULL. <br> Integer value indicating end state. If trimvalue is not NULL, all records after the |
| frimvalue instance of state = trimvalue will be deleted. |  |

## Value

A data table with n rows if in wide format, or n by chainLen rows if in long format.

## Examples

```
def1 <- defData(varname = "x1", formula = 0, variance = 1)
def1 <- defData(def1, varname = "x2", formula = 0, variance = 1)
def1 <- defData(def1,
    varname = "S0", formula = ".6;.3;.1",
    dist = "categorical"
)
dd <- genData(20, def1)
# Transition matrix P
P <- t(matrix(c(
    0.7, 0.2, 0.1,
    0.5, 0.3, 0.2,
    0.0, 0.7, 0.3
),
nrow = 3
))
d1 <- addMarkov(dd, P, chainLen = 3)
d2 <- addMarkov(dd, P, chainLen = 5, wide = TRUE)
d3 <- addMarkov(dd, P, chainLen = 5, wide = TRUE, start0lab = "S0")
d4 <- addMarkov(dd, P, chainLen = 5, start0lab = "S0", trimvalue = 3)
```


## Description

Add multi-factorial data

## Usage

addMultiFac(dtOld, nFactors, levels = 2, coding = "dummy", colNames = NULL)

## Arguments

| dtOld | data.table that is to be modified |
| :--- | :--- |
| nFactors | Number of factors (columns) to generate. |
| levels | Vector or scalar. If a vector is specified, it must be the same length as nFatctors. <br> Each value of the vector represents the number of levels of each corresponding <br> factor. If a scalar is specified, each factor will have the same number of levels. <br> The default is 2 levels for each factor. |
| coding | String value to specify if "dummy" or "effect" coding is used. Defaults to <br> "dummy". |
| colNames | A vector of strings, with a length of $n$ nFactors. The strings represent the name <br> for each factor. |

## Value

A data.table that contains the added simulated data. Each new column contains an integer.

## Examples

```
defD <- defData(varname = "x", formula = 0, variance = 1)
DT <- genData(360, defD)
DT <- addMultiFac(DT, nFactors = 3, levels = c(2, 3, 3), colNames = c("A", "B", "C"))
DT
DT[, .N, keyby = .(A, B, C)]
DT <- genData(300, defD)
DT <- addMultiFac(DT, nFactors = 3, levels = 2)
DT[, .N, keyby = .(Var1, Var2, Var3)]
```

addPeriods

## Description

Create longitudinal/panel data

## Usage

```
addPeriods(
    dtName,
    nPeriods = NULL,
    idvars = "id",
    timevars = NULL,
    timevarName = "timevar",
    timeid = "timeID",
    perName = "period",
    periodVec = NULL
)
```


## Arguments

| dtName | Name of existing data table |
| :--- | :--- |
| nPeriods | Number of time periods for each record |
| idvars | Names of index variables (in a string vector) that will be repeated during each <br> time period |
| timevars | Names of time dependent variables. Defaults to NULL. |
| timevarName | Name of new time dependent variable |
| timeid | Variable name for new index field. Defaults to "timevar" |
| perName | Variable name for period field. Defaults to "period" |
| periodVec | Vector of period times. Defaults to NULL |

## Details

It is possible to generate longitudinal data with varying numbers of measurement periods as well as varying time intervals between each measurement period. This is done by defining specific variables in the data set that define the number of observations per subject and the average interval time between each observation. nCount defines the number of measurements for an individual; mInterval specifies the average time between intervals for a subject; and vInterval specifies the variance of those interval times. If mInterval is not defined, no intervals are used. If vInterval is set to 0 or is not defined, the interval for a subject is determined entirely by the mean interval. If vInterval is greater than 0 , time intervals are generated using a gamma distribution with specified mean and dispersion. If either nPeriods or timevars is specified, that will override any $\boldsymbol{n C o u n t}$, mInterval, and vInterval data.
periodVec is used to specify measurement periods that are different the default counting variables. If periodVec is not specified, the periods default to $0,1, \ldots n-1$, with $n$ periods. If periodVec is specified as $c\left(x \_1, x_{-} 2, \ldots x \_n\right)$, then $x_{-} 1, x_{-} 2, \ldots x_{-} n$ represent the measurement periods.

## Value

An updated data.table that that has multiple rows per observation in dtName

## Examples

```
tdef <- defData(varname = "T", dist = "binary", formula = 0.5)
tdef <- defData(tdef, varname = "Y0", dist = "normal", formula = 10, variance = 1)
tdef <- defData(tdef, varname = "Y1", dist = "normal", formula = "Y0 + 5 + 5* T", variance = 1)
tdef <- defData(tdef, varname = "Y2", dist = "normal", formula = "Y0 + 10 + 5 * T", variance = 1)
dtTrial <- genData(5, tdef)
dtTrial
dtTime <- addPeriods(dtTrial,
    nPeriods = 3, idvars = "id",
    timevars = c("Y0", "Y1", "Y2"), timevarName = "Y"
)
dtTime
# Varying # of periods and intervals - need to have variables
# called nCount and mInterval
def <- defData(varname = "xbase", dist = "normal", formula = 20, variance = 3)
def <- defData(def, varname = "nCount", dist = "noZeroPoisson", formula = 6)
def <- defData(def, varname = "mInterval", dist = "gamma", formula = 30, variance = .01)
def <- defData(def, varname = "vInterval", dist = "nonrandom", formula = .07)
dt <- genData(200, def)
dt[id %in% c(8, 121)]
dtPeriod <- addPeriods(dt)
dtPeriod[id %in% c(8, 121)] # View individuals 8 and 121 only
```

addSynthetic

Add synthetic data to existing data set

## Description

This function generates synthetic data from an existing data.table and adds it to another (simstudy) data.table.

## Usage

addSynthetic(dtOld, dtFrom, vars = NULL, id = "id")

## Arguments

dtOld data.table that is to be modified
dtFrom Data table that contains the source data
vars A vector of string names specifying the fields that will be sampled. The default is that all variables will be selected.
id
A string specifying the field that serves as the record id. The default field is "id".

## Details

Add synthetic data

## Value

A data.table that contains the added synthetic data.

## Examples

```
### Create fake "real" data set - this is the source of the synthetic data
d <- defData(varname = "a", formula = 3, variance = 1, dist = "normal")
d <- defData(d, varname = "b", formula = 5, dist = "poisson")
d <- defData(d, varname = "c", formula = 0.3, dist = "binary")
d <- defData(d, varname = "d", formula = "a + b + 3*c", variance = 2, dist = "normal")
### Create synthetic data set from "observed" data set A (normally this
### would be an actual external data set):
A <- genData(1000, d)
### Generate new simstudy data set (using 'def')
def <- defData(varname = "x", formula = 0, variance = 5)
S <- genData(120, def)
### Create synthetic data from 'A' and add to simulated data in 'S'
S <- addSynthetic(dtOld = S, dtFrom = A, vars = c("b", "d"))
```

betaGetShapes Convert beta mean and precision parameters to two shape parameters

## Description

Convert beta mean and precision parameters to two shape parameters

## Usage

betaGetShapes(mean, precision)

## Arguments

mean
The mean of a beta distribution
precision
The precision parameter (phi) of a beta distribution

## Details

In simstudy, users specify the beta distribution as a function of two parameters - a mean and precision, where $0<$ mean $<1$ and precision $>0$. In this case, the variance of the specified distribution is (mean) $)^{*}(1-$ mean $) /(1+$ precision $)$. The base R function rbeta uses the two shape parameters to specify the beta distribution. This function converts the mean and precision into the shape 1 and shape 2 parameters.

## Value

A list that includes the shape parameters of the beta distribution

## Examples

```
set.seed(12345)
mean <- 0.3
precision <- 1.6
rs <- betaGetShapes(mean, precision)
c(rs$shape1, rs$shape2)
vec <- rbeta(1000, shape1 = rs$shape1, shape2 = rs$shape2)
(estMoments <- c(mean(vec), var(vec)))
(theoryMoments <- c(mean, mean * (1 - mean) / (1 + precision)))
(theoryMoments <- with(rs, c(
    shape1 / (shape1 + shape2),
    (shape1 * shape2) / ((shape1 + shape2)^2 * (1 + shape1 + shape2))
)))
```

blockDecayMat Create a block correlation matrix

## Description

The function genBlockMat() generates correlation matrices that can accommodate clustered observations over time where the within-cluster between-individual correlation in the same time period can be different from the within-cluster between-individual correlation across time periods. The matrix generated here can be used in function addCorGen().

## Usage

blockDecayMat(ninds, nperiods, rho_w, r, pattern = "xsection", nclusters = 1)

## Arguments

ninds $\quad$ The number of units (individuals) in each cluster in each period.
nperiods The number periods that data are observed.
rho_w The within-period/between-individual correlation coefficient between -1 and 1 .
r
The decay parameter if correlation declines over time, and can have values of "exp" or "prop". See details.
pattern A string argument with options "xsection" (default) or "cohort".
nclusters An integer that indicates the number of matrices that will be generated.

## Details

Two general decay correlation structures are currently supported: a *cross-sectional* exchangeable structure and a *closed cohort* exchangeable structure. In the *cross-sectional* case, individuals or units in each time period are distinct. In the *closed cohort* structure, individuals or units are repeated in each time period. The desired structure is specified using pattern, which defaults to "xsection" if not specified.
This function can generate correlation matrices of different sizes, depending on the combination of arguments provided. A single matrix will be generated when nclusters == 1 (the default), and a list of matrices of matrices will be generated when nclusters $>1$.
If nclusters $>1$, the length of ninds will depend on if sample sizes will vary by cluster and/or period. There are three scenarios, and function evaluates the length of ninds to determine which approach to take:

- if the sample size is the same for all clusters in all periods, ninds will be a single value (i.e., length =1).
- if the sample size differs by cluster but is the same for each period within each cluster each period, then ninds will have a value for each cluster (i.e., length = nclusters).
- if the sample size differs across clusters and across periods within clusters, ninds will have a value for each cluster-period combination (i.e., length $=$ nclusters $\times$ nperiods). This option is only valid when pattern = "xsection".

In addition, rho_w and $r$ can be specified as a single value (in which case they are consistent across all clusters) or as a vector of length nclusters, in which case either one or both of these parameters can vary by cluster.
See vignettes for more details.

## Value

A single correlation matrix of size nvars $x$ nvars, or a list of matrices of potentially different sizes with length indicated by nclusters.

A single correlation matrix or a list of matrices of potentially different sizes with length indicated by nclusters.

## References

Li et al. Mixed-effects models for the design and analysis of stepped wedge cluster randomized trials: An overview. Statistical Methods in Medical Research. 2021;30(2):612-639. doi:10.1177/0962280220932962

## See Also

blockExchangeMat and addCorGen

## Examples

```
blockDecayMat(ninds = 4, nperiods = 3, rho_w = .8, r = .9)
blockDecayMat(ninds = 4, nperiods = 3, rho_w = .8, r = .9, pattern = "cohort")
blockDecayMat(ninds = 2, nperiods = 3, rho_w = . 8, r = .9, pattern = "cohort", nclusters=2)
```

```
blockDecayMat(ninds = c(2, 3), nperiods = 3, rho_w = c(.8,0.7), r = c(.9,.8),
    pattern = "cohort", nclusters=2)
blockDecayMat(ninds = c(2, 3, 4, 4, 2, 1), nperiods = 3, rho_w = .8, r = .9, nclusters=2)
```


## blockExchangeMat Create a block correlation matrix with exchangeable structure

## Description

The function blockExchangeMat generates exchangeable correlation matrices that can accommodate clustered observations over time where the within-cluster between-individual correlation in the same time period can be different from the within-cluster between-individual correlation across time periods. The matrix generated here can be used in function addCorGen.

## Usage

blockExchangeMat ( ninds, nperiods, rho_w, rho_b = 0, rho_a = NULL, pattern = "xsection", nclusters = 1
)

## Arguments

| ninds | The number of units (individuals) in each cluster in each period. |
| :--- | :--- |
| nperiods | The number periods that data are observed. |
| rho_w | The within-period/between-individual correlation coefficient between -1 and 1. |
| rho_b | The between-period/between-individual correlation coefficient between -1 and <br> 1. |
| rho_a | The between-period/within-individual auto-correlation coefficient between -1 <br> and 1. |
| pattern | A string argument with options "xsection" (default) or "cohort". |
| nclusters | An integer that indicates the number of matrices that will be generated. |

## Details

Two general exchangeable correlation structures are currently supported: a *cross-sectional* exchangeable structure and a *closed cohort* exchangeable structure. In the *cross-sectional* case, individuals or units in each time period are distinct. In the *closed cohort* structure, individuals or units are repeated in each time period. The desired structure is specified using pattern, which
defaults to "xsection" if not specified. rho_a is the within-individual/unit exchangeable correlation over time, and can only be used when xsection = FALSE.
This function can generate correlation matrices of different sizes, depending on the combination of arguments provided. A single matrix will be generated when nclusters $==1$ (the default), and a list of matrices of matrices will be generated when nclusters $>1$.

If nclusters $>1$, the length of ninds will depend on if sample sizes will vary by cluster and/or period. There are three scenarios, and function evaluates the length of ninds to determine which approach to take:

- if the sample size is the same for all clusters in all periods, ninds will be a single value (i.e., length $=1$ ).
- if the sample size differs by cluster but is the same for each period within each cluster each period, then ninds will have a value for each cluster (i.e., length = nclusters).
- if the sample size differs across clusters and across periods within clusters, ninds will have a value for each cluster-period combination (i.e., length $=$ nclusters $\times$ nperiods). This option is only valid when pattern = "xsection".

In addition, rho_w, rho_b, and rho_a can be specified as a single value (in which case they are consistent across all clusters) or as a vector of length nclusters, in which case any or all of these parameters can vary by cluster.

See vignettes for more details.

## Value

A single correlation matrix or a list of matrices of potentially different sizes with length indicated by nclusters.

## References

Li et al. Mixed-effects models for the design and analysis of stepped wedge cluster randomized trials: An overview. Statistical Methods in Medical Research. 2021;30(2):612-639. doi:10.1177/0962280220932962

## See Also

blockDecayMat and addCorGen

## Examples

```
blockExchangeMat(ninds \(=4\), nperiods \(=3\), rho_w \(=.8\) )
blockExchangeMat(ninds \(=4\), nperiods \(=3\), rho_w \(=.8\), rho_b \(=0.5\) )
blockExchangeMat(ninds = 4, nperiods = 3, rho_w = .8, rho_b \(=0.5\), rho_a \(=0.7\),
    pattern = "cohort")
blockExchangeMat(ninds \(=2\), nperiods \(=3\), rho_w \(=.8\), rho_b \(=0.5\), rho_a \(=0.7\),
    nclusters = 3, pattern = "cohort")
blockExchangeMat(ninds = c(2, 3), nperiods = 3, rho_w = .8, rho_b = 0.5, rho_a = 0.7,
    nclusters = 2, pattern="cohort")
blockExchangeMat(ninds \(=c(2,3,4,4,2,1)\), nperiods \(=3\), rho_w \(=.8\), rho_b \(=0.5\),
    nclusters = 2)
```


## defCondition Add single row to definitions table of conditions that will be used to add data to an existing definitions table

## Description

Add single row to definitions table of conditions that will be used to add data to an existing definitions table

## Usage

defCondition(
dtDefs = NULL,
condition,
formula,
variance $=0$,
dist = "normal",
link = "identity"
)

## Arguments

| dtDefs | Name of definition table to be modified. Null if this is a new definition. |
| :--- | :--- |
| condition | Formula specifying condition to be checked |
| formula | An R expression for mean (string) |
| variance | Number |
| dist | Distribution. For possibilities, see details |
| link | The link function for the mean, see details |

## Value

A data.table named dtName that is an updated data definitions table

## See Also

distributions

## Examples

```
# New data set
def <- defData(varname = "x", dist = "noZeroPoisson", formula = 5)
def <- defData(def, varname = "y", dist = "normal", formula = 0, variance = 9)
dt <- genData(10, def)
# Add columns to dt
```

```
    defC <- defCondition(
        condition = "x == 1", formula = "5 + 2*y",
        variance = 1, dist = "normal"
    )
    defC <- defCondition(defC,
        condition = "x <= 5 & x >= 2", formula = "3 - 2*y",
        variance = 1, dist = "normal"
    )
    defC <- defCondition(defC,
        condition = "x >= 6", formula = 1,
        variance = 1, dist = "normal"
    )
    defC
    # Add conditional column with field name "z"
    dt <- addCondition(defC, dt, "z")
    dt
```

    defData Add single row to definitions table
    
## Description

Add single row to definitions table

## Usage

```
defData(
    dtDefs = NULL,
    varname,
    formula,
    variance = 0,
    dist = "normal",
    link = "identity",
    id = "id"
)
```


## Arguments

| dtDefs | Definition data.table to be modified |
| :--- | :--- |
| varname | Name (string) of new variable |
| formula | An R expression for mean (string) |
| variance | Number |

dist Distribution. For possibilities, see details
link The link function for the mean, see details
id A string indicating the field name for the unique record identifier

## Details

The possible data distributions are: normal, binary, binomial, poisson, noZeroPoisson, uniform, categorical, gamma, beta, nonrandom, uniformInt, negBinomial, exponential, mixture, trtAssign, clusterSize, custom.

## Value

A data.table named dtName that is an updated data definitions table

## See Also

distributions

## Examples

```
extVar <- 2.3
def <- defData(varname = "xNr", dist = "nonrandom", formula = 7, id = "idnum")
def <- defData(def, varname = "xUni", dist = "uniform", formula = "10;20")
def <- defData(def,
    varname = "xNorm", formula = "xNr + xUni * 2", dist = "normal",
    variance = 8
)
def <- defData(def,
    varname = "xPois", dist = "poisson", formula = "xNr - 0.2 * xUni",
    link = "log"
)
def <- defData(def, varname = "xCat", formula = "0.3;0.2;0.5", dist = "categorical")
def <- defData(def,
    varname = "xGamma", dist = "gamma", formula = "5+xCat",
    variance = 1, link = "log"
)
def <- defData(def,
    varname = "xBin", dist = "binary", formula = "-3 + xCat",
    link = "logit"
)
def <- defData(def,
    varname = "external", dist = "nonrandom",
    formula = "xBin * log(..extVar)"
)
def
```


## Description

Add single row to definitions table that will be used to add data to an existing data.table

## Usage

```
    defDataAdd(
        dtDefs = NULL,
        varname,
        formula,
        variance = 0,
        dist = "normal",
        link = "identity"
    )
```


## Arguments

dtDefs Name of definition table to be modified. Null if this is a new definition.
varname $\quad$ Name (string) of new variable
formula An R expression for mean (string)
variance Number
dist Distribution. For possibilities, see details
link The link function for the mean, see details

## Value

A data.table named dtName that is an updated data definitions table

## See Also

[distributions]

## Examples

```
# New data set
def <- defData(varname = "xNr", dist = "nonrandom", formula = 7, id = "idnum")
def <- defData(def, varname = "xUni", dist = "uniform", formula = "10;20")
dt <- genData(10, def)
# Add columns to dt
```

```
    def2 <- defDataAdd(varname = "y1", formula = 10, variance = 3)
    def2 <- defDataAdd(def2, varname = "y2", formula = .5, dist = "binary")
    def2
    dt <- addColumns(def2, dt)
    dt
```

    defMiss Definitions for missing data
    
## Description

Add single row to definitions table for missing data

## Usage

```
    defMiss(
        dtDefs = NULL,
        varname,
        formula,
        logit.link = FALSE,
        baseline = FALSE,
        monotonic = FALSE
    )
```


## Arguments

| dtDefs | Definition data.table to be modified |
| :--- | :--- |
| varname | Name of variable with missingness |
| formula | Formula to describe pattern of missingness |
| logit.link | Indicator set to TRUE when the probability of missingness is based on a logit <br> model. |
| baseline | Indicator is set to TRUE if the variable is a baseline measure and should be <br> missing throughout an entire observation period. This is applicable to repeated <br> measures/longitudinal data. |
| monotonic | Indicator set to TRUE if missingness at time t is followed by missingness at all <br> follow-up times $>\mathrm{t}$. |

Value
A data.table named dtName that is an updated data definitions table

## See Also

genMiss, genObs

## Examples

```
def1 <- defData(varname = "m", dist = "binary", formula = . 5)
def1 <- defData(def1, "u", dist = "binary", formula = .5)
def1 <- defData(def1, "x1", dist = "normal", formula = "20*m + 20*u", variance = 2)
def1 <- defData(def1, "x2", dist = "normal", formula = "20*m + 20*u", variance = 2)
def1 <- defData(def1, "x3", dist = "normal", formula = "20*m + 20*u", variance = 2)
dtAct <- genData(1000, def1)
defM <- defMiss(varname = "x1", formula = .15, logit.link = FALSE)
defM <- defMiss(defM, varname = "x2", formula = ".05 + m * 0.25", logit.link = FALSE)
defM <- defMiss(defM, varname = "x3", formula = ".05 + u * 0.25", logit.link = FALSE)
defM <- defMiss(defM, varname = "u", formula = 1, logit.link = FALSE) # not observed
defM
# Generate missing data matrix
missMat <- genMiss(dtName = dtAct, missDefs = defM, idvars = "id")
missMat
# Generate observed data from actual data and missing data matrix
dtObs <- genObs(dtAct, missMat, idvars = "id")
dtObs
```

defRead Read external csv data set definitions

## Description

Read external csv data set definitions

## Usage

defRead(filen, id = "id")

## Arguments

filen String file name, including full path. Must be a csv file.
id string that includes name of id field. Defaults to "id"

## Value

A data.table with data set definitions

## See Also

[distributions]

## Examples

```
    # Create temporary external "csv" file
    test1 <- c(
    "varname,formula,variance,dist,link",
    "nr,7, 0,nonrandom,identity",
    "x1,.4, 0,binary,identity",
    "y1,nr + x1 * 2,8,normal,identity",
    "y2,nr - 0.2 * x1,0,poisson, log"
    )
    tfcsv <- tempfile()
    writeLines(test1, tfcsv)
    # Read external csv file stored in file "tfcsv"
    defs <- defRead(tfcsv, id = "myID")
    defs
    unlink(tfcsv)
    # Generate data based on external definition
    genData(5, defs)
```

    defReadAdd Read external csv data set definitions for adding columns
    
## Description

Read external csv data set definitions for adding columns

## Usage

defReadAdd(filen)

## Arguments

filen String file name, including full path. Must be a csv file.

## Value

A data.table with data set definitions

## See Also

[distributions]

## Examples

```
# Create temporary external "csv" files
    test1 <- c(
    "varname, formula, variance,dist,link",
    "nr,7, 0,nonrandom,identity"
)
tfcsv1 <- tempfile()
writeLines(test1, tfcsv1)
test2 <- c(
    "varname, formula, variance, dist,link",
    "x1,.4, 0,binary,identity",
    "y1,nr + x1 * 2,8,normal,identity",
    "y2,nr - 0.2 * x1,0,poisson, log"
)
tfcsv2 <- tempfile()
writeLines(test2, tfcsv2)
# Generate data based on external definitions
defs <- defRead(tfcsv1)
dt <- genData(5, defs)
dt
# Add additional data based on external definitions
defs2 <- defReadAdd(tfcsv2)
dt <- addColumns(defs2, dt)
dt
unlink(tfcsv1)
unlink(tfcsv2)
```

defReadCond

## Description

Read external csv data set definitions for adding columns

## Usage

defReadCond(filen)

## Arguments

filen String file name, including full path. Must be a csv file.

## Value

A data.table with data set definitions

## See Also

[distributions]

## Examples

```
# Create temporary external "csv" files
    test1 <- c(
    "varname,formula,variance,dist,link",
    "x,0.3;0.4;0.3,0,categorical,identity"
)
tfcsv1 <- tempfile()
writeLines(test1, tfcsv1)
test2 <- c(
    "condition,formula,variance,dist,link",
    "x == 1, 0.4,0,binary,identity",
    "x == 2, 0.6,0,binary,identity",
    "x >= 3, 0.8,0,binary,identity"
)
    tfcsv2 <- tempfile()
    writeLines(test2, tfcsv2)
    # Generate data based on external definitions
    defs <- defRead(tfcsv1)
    dt <- genData(2000, defs)
    dt
    # Add column based on
    defsCond <- defReadCond(tfcsv2)
    dt <- addCondition(defsCond, dt, "y")
    dt
    dt[, mean(y), keyby = x]
    unlink(tfcsv1)
    unlink(tfcsv2)
```

    defRepeat
    
## Description

Add multiple (similar) rows to definitions table

## Usage

```
defRepeat(
        dtDefs = NULL,
        nVars,
        prefix,
        formula,
        variance = 0,
        dist = "normal",
        link = "identity",
        id = "id"
    )
```


## Arguments

| dtDefs | Definition data.table to be modified |
| :--- | :--- |
| nVars | Number of new variables to define |
| prefix | Prefix (character) for new variables |
| formula | An R expression for mean (string) |
| variance | Number or formula |
| dist | Distribution. For possibilities, see details |
| link | The link function for the mean, see details |
| id | A string indicating the field name for the unique record identifier |

## Details

The possible data distributions are: 'r paste 0 (.getDists(),collapse = ", ")'.

## Value

A data.table named dtName that is an updated data definitions table

## See Also

[distributions]

## Examples

```
def <- defRepeat(
    nVars = 4, prefix = "g", formula = "1/3;1/3;1/3",
    variance = 0, dist = "categorical"
)
def <- defData(def, varname = "a", formula = "1;1", dist = "trtAssign")
def <- defRepeat(def, 8, "b", formula = "5 + a", variance = 3, dist = "normal")
def <- defData(def, "y", formula = "0.10", dist = "binary")
```

def
defRepeatAdd Add multiple (similar) rows to definitions table that will be used to add data to an existing data.table

## Description

Add multiple (similar) rows to definitions table that will be used to add data to an existing data.table

## Usage

```
    defRepeatAdd(
        dtDefs = NULL,
        nVars,
        prefix,
        formula,
        variance = 0,
        dist = "normal",
        link = "identity",
        id = "id"
    )
```


## Arguments

dtDefs Definition data.table to be modified
nVars Number of new variables to define
prefix Prefix (character) for new variables
formula An R expression for mean (string)
variance Number or formula
dist Distribution. For possibilities, see details
link The link function for the mean, see details
id A string indicating the field name for the unique record identifier

## Details

The possible data distributions are: 'r paste0(.getDists(),collapse = ", ")'.

## Value

A data.table named dtName that is an updated data definitions table

## See Also

[distributions]

## Examples

```
    def <- defRepeatAdd(
        nVars = 4, prefix = "g", formula = "1/3;1/3;1/3",
        variance = 0, dist = "categorical"
)
def <- defDataAdd(def, varname = "a", formula = "1;1", dist = "trtAssign")
def <- defRepeatAdd(def, 8, "b", formula = "5 + a", variance = 3, dist = "normal")
def <- defDataAdd(def, "y", formula = "0.10", dist = "binary")
def
```

    defSurv Add single row to survival definitions
    
## Description

Add single row to survival definitions

## Usage

```
defSurv(
    dtDefs = NULL,
    varname,
    formula = 0,
    scale = 1,
    shape = 1,
    transition = 0
    )
```


## Arguments

| dtDefs | Definition data.table to be modified |
| :--- | :--- |
| varname | Variable name |
| formula | Covariates predicting survival |
| scale | Scale parameter for the Weibull distribution. |
| shape | The shape of the Weibull distribution. Shape $=1$ for an exponential distribution |
| transition | An integer value indicating the starting point for a new specification of the haz- <br> ard function. It will default to 0 (and must be 0 ) for the first instance of a <br> "varname". |

## Value

A data.table named dtName that is an updated data definitions table

## Examples

```
    # Baseline data definitions
    def <- defData(varname = "x1", formula = .5, dist = "binary")
    def <- defData(def, varname = "x2", formula = .5, dist = "binary")
    def <- defData(def, varname = "grp", formula = .5, dist = "binary")
    # Survival data definitions
    sdef <- defSurv(
    varname = "survTime", formula = "1.5*x1",
    scale = "grp*50 + (1-grp)*25", shape = "grp*1 + (1-grp)*1.5"
)
sdef <- defSurv(sdef, varname = "censorTime", scale = 80, shape = 1)
sdef
    # Baseline data definitions
    dtSurv <- genData(300, def)
    # Add survival times
    dtSurv <- genSurv(dtSurv, sdef)
    head(dtSurv)
```

    delColumns Delete columns from existing data set
    
## Description

Delete columns from existing data set

## Usage

delColumns(dtOld, vars)

## Arguments

dtold Name of data table that is to be updated.
vars Vector of column names (as strings).

## Value

An updated data.table without vars.

## Examples

```
    # New data set
    def <- defData(varname = "x", dist = "noZeroPoisson", formula = 7, id = "idnum")
    def <- defData(def, varname = "xUni", dist = "uniformInt", formula = "x-3;x+3")
    dt <- genData(10, def)
    dt
    # Delete column
    dt <- delColumns(dt, "x")
    dt
```

    distributions Distributions for Data Definitions
    
## Description

This help file describes the distributions used for data creation in simstudy.

## Arguments

| formula | Desired mean as a Number or an R expression for mean as a String. Variables <br> defined via defData() and variables within the parent environment (prefixed <br> with . .) can be used within the formula. Functions from the parent environment <br> can be used without a prefix. |
| :--- | :--- |
| variance | Number. Default is 0. |
| link | String identifying the link function to be used. Default is identity. |

## Details

For details about the statistical distributions please see stats:: distributions, any non-statistical distributions will be explained below. Required variables and expected pattern for each distribution can be found in this table:

| name | formula | format | variance |
| :--- | :--- | :--- | :--- |
| beta | mean | String or Number | dispersion value |
| binary | probability for 1 | String or Number | NA |
| binomial | probability of success | String or Number | number of trials |
| categorical | probabilities | p_1;p_2; . ;p_n | category labels: $a ; b ; c, 50 ; 130 ; 20$ |
| exponential | mean (lambda) | String or Number | NA |
| gamma | mean | String or Number | dispersion value |
| mixture | formula | x_1 lp_1 + x_2lp_2 $\ldots$ x_nl p_n | NA |
| negBinomial | mean | String or Number | dispersion value |
| nonrandom | formula | String or Number | NA |
| normal | mean | String or Number | variance |

link identity identity identity identity identity identity NA identity NA NA

| noZeroPoisson | mean | String or Number | NA |
| :--- | :--- | :--- | :--- |
| poisson | mean | String or Number | NA |
| trtAssign | ratio | $r_{-} 1 ; r_{-} 2 ; \ldots ; r_{-} n$ | stratification |
| uniform | range | from; to | NA |
| uniformInt | range | from; to | NA |

identity
identity
identity
NA
NA

## Mixture

The mixture distribution makes it possible to mix to previously defined distributions/variables. Each variable that should be part of the new distribution $x_{-} 1, \ldots, X_{-} n$ is assigned a probability p_1, . . , p_n. For more information see rdatagen.net.

## Examples

```
ext_var <- 2.9
def <- defData(varname = "external", formula = "3 + log(..ext_var)", variance = .5)
def
genData(5, def)
```

gammaGetShapeRate | Convert gamma mean and dispersion parameters to shape and rate |
| :--- |
| parameters |

## Description

Convert gamma mean and dispersion parameters to shape and rate parameters

## Usage

gammaGetShapeRate(mean, dispersion)

## Arguments

mean The mean of a gamma distribution
dispersion The dispersion parameter of a gamma distribution

## Details

In simstudy, users specify the gamma distribution as a function of two parameters - a mean and dispersion. In this case, the variance of the specified distribution is (mean^2)*dispersion. The base R function rgamma uses the shape and rate parameters to specify the gamma distribution. This function converts the mean and dispersion into the shape and rate.

## Value

A list that includes the shape and rate parameters of the gamma distribution

## Examples

```
    set.seed(12345)
    mean <- 5
    dispersion <- 1.5
    rs <- gammaGetShapeRate(mean, dispersion)
    c(rs$shape, rs$rate)
    vec <- rgamma(1000, shape = rs$shape, rate = rs$rate)
    (estMoments <- c(mean(vec), var(vec)))
    (theoryMoments <- c(mean, mean^2 * dispersion))
    (theoryMoments <- c(rs$shape / rs$rate, rs$shape / rs$rate^2))
```

    genCatFormula Generate Categorical Formula
    
## Description

Create a semi-colon delimited string of probabilities to be used to define categorical data.

## Usage

genCatFormula(..., $\mathrm{n}=0$ )

## Arguments

$$
\begin{array}{ll}
\ldots & \text { one or more numeric values to be concatenated, delimited by ";". } \\
\mathrm{n} & \text { Number of probabilities (categories) to be generated - all with equal probability. }
\end{array}
$$

## Details

The function accepts a number of probabilities or a value of $n$, but not both.
If probabilities are passed, the string that is returned depends on the nature of those probabilities. If the sum of the probabilities is less than 1 , an additional category is created with the probability $1-\operatorname{sum}($ provided probabilities). If the sum of the probabilities is equal to 1 , then the number of categories is set to the number of probabilities provided. If the sum of the probabilities exceeds one (and there is more than one probability), the probabilities are standardized by dividing by the sum of the probabilities provided.

If $n$ is provided, $n$ probabilities are included in the string, each with a probability equal to $1 / n$.

## Value

string with multinomial probabilities.

## Examples

```
genCatFormula(0.25, 0.25, 0.50)
genCatFormula(1 / 3, 1 / 2)
genCatFormula(1, 2, 3)
genCatFormula(n = 5)
```

genCluster $\quad$ Simulate clustered data

## Description

Simulate data set that is one level down in a multilevel data context. The level "2" data set must contain a field that specifies the number of individual records in a particular cluster.

## Usage

genCluster(dtClust, cLevelVar, numIndsVar, level1ID, allLevel2 = TRUE)

## Arguments

$\left.\begin{array}{ll}\begin{array}{l}\text { dtClust } \\ \text { cLevelVar } \\ \text { numIndsVar }\end{array} & \begin{array}{l}\text { Name of existing data set that contains the level "2" data } \\ \text { Variable name (string) of cluster id in dtClust }\end{array} \\ \text { Variable name (string) of number of observations per cluster in dtClust. Can } \\ \text { also be a single integer value that will be used for all clusters. }\end{array}\right\}$

## Value

A simulated data table with level "1" data

## Examples

```
gen.school <- defData(
    varname = "s0", dist = "normal",
    formula = 0, variance = 3, id = "idSchool"
)
gen.school <- defData(gen.school,
    varname = "nClasses",
    dist = "noZeroPoisson", formula = 3
)
dtSchool <- genData(3, gen.school) #'
dtSchool
dtClass <- genCluster(dtSchool,
    cLevelVar = "idSchool",
    numIndsVar = "nClasses", level1ID = "idClass"
)
dtClass
dtClass <- genCluster(dtSchool,
```

```
        cLevelVar = "idSchool",
        numIndsVar = 3, level1ID = "idClass"
    )
    dtClass
```

    genCorData Create correlated data
    
## Description

## Create correlated data

```
Usage
    genCorData(
        n,
        mu,
        sigma,
        corMatrix = NULL,
        rho,
        corstr = "ind",
        cnames = NULL,
        idname = "id"
    )
```


## Arguments

| n | Number of observations |
| :--- | :--- |
| mu | A vector of means. The length of mu must be nvars. |
| corMatrix | Standard deviation of variables. If standard deviation differs for each variable, <br> enter as a vector with the same length as the mean vector mu. If the standard <br> deviation is constant across variables, as single value can be entered. |
| rorrelation matrix can be entered directly. It must be symmetrical and posi- |  |
| tive semi-definite. It is not a required field; if a matrix is not provided, then a |  |
| structure and correlation coefficient rho must be specified. |  |

## Value

A data.table with n rows and the $\mathrm{k}+1$ columns, where k is the number of means in the vector mu.

## Examples

```
mu <- c(3, 8, 15)
sigma <- c(1, 2, 3)
corMat <- matrix(c(1, .2, .8, .2, 1, .6, .8, .6, 1), nrow = 3)
dtcor1 <- genCorData(1000, mu = mu, sigma = sigma, rho = .7, corstr = "cs")
dtcor2 <- genCorData(1000, mu = mu, sigma = sigma, corMatrix = corMat)
dtcor1
dtcor2
round(var(dtcor1[, .(V1, V2, V3)]), 3)
round(cor(dtcor1[, .(V1, V2, V3)]), 2)
round(var(dtcor2[, .(V1, V2, V3)]), 3)
round(cor(dtcor2[, .(V1, V2, V3)]), 2)
```

genCorFlex

Create multivariate (correlated) data - for general distributions

## Description

Create multivariate (correlated) data - for general distributions

## Usage

genCorFlex(n, defs, rho = 0, tau = NULL, corstr = "cs", corMatrix = NULL)

## Arguments

| n | Number of observations |
| :--- | :--- |
| defs | Field definition table created by function 'defData'. All definitions must be <br> scalar. Definition specifies distribution, mean, and variance, with all caveats for <br> each of the distributions. (See defData). |
| rho | Correlation coefficient, $-1<=$ rho $<=1$. Use if corMatrix is not provided. |
| tau | Correlation based on Kendall's tau. If tau is specified, then it is used as the <br> correlation even if rho is specified. If tau is NULL, then the specified value of <br> rho is used, or rho defaults to 0. |
| cormatrix | Correlation structure of the variance-covariance matrix defined by sigma and <br> rho. Options include "cs" for a compound symmetry structure and "ar1" for an <br> autoregressive structure. Defaults to "cs". |
|  | Correlation matrix can be entered directly. It must be symmetrical and positive <br> semi-definite. It is not a required field; if a matrix is not provided, then a struc- <br> ture and correlation coefficient rho must be specified. This is only used if tau is |
| not specified. |  |

## Value

data.table with added column(s) of correlated data

## Examples

```
    ## Not run:
    def <- defData(varname = "xNorm", formula = 0, variance = 4, dist = "normal")
    def <- defData(def, varname = "xGamma1", formula = 15, variance = 2, dist = "gamma")
    def <- defData(def, varname = "xBin", formula = 0.5, dist = "binary")
    def <- defData(def, varname = "xUnif1", formula = "0;10", dist = "uniform")
    def <- defData(def, varname = "xPois", formula = 15, dist = "poisson")
    def <- defData(def, varname = "xUnif2", formula = "23;28", dist = "uniform")
    def <- defData(def, varname = "xUnif3", formula = "100;150", dist = "uniform")
    def <- defData(def, varname = "xGamma2", formula = 150, variance = 0.003, dist = "gamma")
    def <- defData(def, varname = "xNegBin", formula = 5, variance = .8, dist = "negBinomial")
    dt <- genCorFlex(1000, def, tau = 0.3, corstr = "cs")
    cor(dt[, -"id"])
    cor(dt[, -"id"], method = "kendall")
    var(dt[, -"id"])
    apply(dt[, -"id"], 2, mean)
    ## End(Not run)
```

    genCorGen Create multivariate (correlated) data - for general distributions
    
## Description

Create multivariate (correlated) data - for general distributions

## Usage

```
genCorGen(
    n,
    nvars,
    params1,
    params2 = NULL,
    dist,
    rho,
    corstr,
    corMatrix = NULL,
    wide = FALSE,
    cnames = NULL,
    method = "copula",
    idname = "id"
)
```


## Arguments

| n | Number of observations |
| :---: | :---: |
| nvars | Number of variables |
| params1 | A single vector specifying the mean of the distribution. The vector is of length 1 if the mean is the same across all observations, otherwise the vector is of length nvars. In the case of the uniform distribution the vector specifies the minimum. |
| params2 | A single vector specifying a possible second parameter for the distribution. For the normal distribution, this will be the variance; for the gamma distribution, this will be the dispersion; and for the uniform distribution, this will be the maximum. The vector is of length 1 if the mean is the same across all observations, otherwise the vector is of length nvars. |
| dist | A string indicating "binary", "poisson" or "gamma", "normal", or "uniform". |
| rho | Correlation coefficient, $-1<=$ rho $<=1$. Use if corMatrix is not provided. |
| corstr | Correlation structure of the variance-covariance matrix defined by sigma and rho. Options include "cs" for a compound symmetry structure and "ar1" for an autoregressive structure. |
| corMatrix | Correlation matrix can be entered directly. It must be symmetrical and positive semi-definite. It is not a required field; if a matrix is not provided, then a structure and correlation coefficient rho must be specified. |
| wide | The layout of the returned file - if wide = TRUE, all new correlated variables will be returned in a single record, if wide $=$ FALSE, each new variable will be its own record (i.e. the data will be in long form). Defaults to FALSE. |
| cnames | Explicit column names. A single string with names separated by commas. If no string is provided, the default names will be V\#, where \# represents the column. |
| method | Two methods are available to generate correlated data. (1) "copula" uses the multivariate Gaussian copula method that is applied to all other distributions; this applies to all available distributions. (2) "ep" uses an algorithm developed by Emrich and Piedmonte (1991). |
| idname | Character value that specifies the name of the id variable. |

## Value

data.table with added column(s) of correlated data

## References

Emrich LJ, Piedmonte MR. A Method for Generating High-Dimensional Multivariate Binary Variates. The American Statistician 1991;45:302-4.

## Examples

```
set.seed(23432)
lambda <- c(8, 10, 12)
genCorGen(100, nvars = 3, params1 = lambda, dist = "poisson", rho = .7, corstr = "cs")
genCorGen(100, nvars = 3, params1 = 5, dist = "poisson", rho = . 7, corstr = "cs")
```

```
genCorGen(100, nvars = 3, params1 = lambda, dist = "poisson", rho = . 7, corstr = "cs", wide = TRUE)
genCorGen(100, nvars = 3, params1 = 5, dist = "poisson", rho = . 7, corstr = "cs", wide = TRUE)
genCorGen(100,
    nvars = 3, params1 = lambda, dist = "poisson", rho = .7, corstr = "cs",
    cnames = "new_var"
)
genCorGen(100,
    nvars = 3, params1 = lambda, dist = "poisson", rho = .7, corstr = "cs",
    wide = TRUE, cnames = "a, b, c"
)
```


## genCorMat $\quad$ Create a correlation matrix

## Description

Create a correlation matrix

## Usage

genCorMat(nvars, cors = NULL, rho = NULL, corstr = "cs", nclusters = 1)

## Arguments

nvars number of rows and columns (i.e. number of variables) for correlation matrix. It can be a scalar or vector (see details).
cors vector of correlations.
rho Correlation coefficient, $-1<=r h o<=1$. Use if corMatrix is not provided. It can be a scalar or vector (see details).
corstr Correlation structure. Options include "cs" for a compound symmetry structure, "ar1" for an autoregressive structure of order 1, "arx" for an autoregressive structure that has a general decay pattern, and "structured" that imposes a prescribed pattern between observation based on distance (see details).
nclusters An integer that indicates the number of matrices that will be generated.

## Details

This function can generate correlation matrices randomly or deterministically, depending on the combination of arguments provided. A single matrix will be generated when nclusters $==1$ (the default), and a list of matrices of matrices will be generated when nclusters $>1$.
If the vector 'cors' is specified with length 'nvars - 1' then 'corstr' must be "structured". If 'cors' is specified with length 'choose(nvars, 2)' then 'corstr' should not be specified as "structured". In this case the 'cors' vector should be interpreted as the lower triangle of the correlation matrix, and is specified by reading down the columns. For example, if $\mathbf{C M}$ is the correlation matrix and nvars $=3$, then $\operatorname{CM}[2,1]=\operatorname{CM}[1,2]=\operatorname{cors}[1], \operatorname{CM}[3,1]=\operatorname{CM}[1,3]=\operatorname{cors}[2]$, and $\operatorname{CM}[3,2]=\operatorname{CM}[2,3]$ $=\operatorname{cors}[3]$.

If the vector cors and rho are not specified, random correlation matrices are generated based on the specified corstr. If the structure is "arx", then a random vector of length nvars - 1 is randomly generated and sorted in descending order; the correlation matrix will be generated base on this set of structured correlations. If the structure is not specified as "arx" then a random positive definite of dimensions nvars x nvars with no structural assumptions is generated.

If cors is not specified but rho is specified, then a matrix with either a "cs" or "ar1" structure is generated.

If nclusters $>1$, nvars can be of length 1 or nclusters. If it is of length 1 , each cluster will have correlation matrices with the same dimension. Likewise, if nclusters > 1, rho can be of length 1 or nclusters. If length of rho is 1 , each cluster will have correlation matrices with the same autocorrelation.

## Value

A single correlation matrix of size nvars $x$ nvars, or a list of matrices of potentially different sizes with length indicated by nclusters.

## Examples

```
genCorMat(nvars = 3, cors = c(.3, -.2, .1))
genCorMat(nvars = 3)
genCorMat(nvars = 4, c(.3, -.2, .1, .2, .5, .2))
genCorMat(4)
genCorMat(nvars = 4, cors = c(.3, .2, .1), corstr = "structured")
genCorMat(nvars = 4, corstr = "arx")
genCorMat(nvars = 4, rho = .4, corstr = "cs")
genCorMat(nvars = 4, rho = .4, corstr = "ar1")
genCorMat(nvars = c(3, 2, 5), rho = c(.4, .8, .7), corstr = "ar1", nclusters = 3)
```

genData Calling function to simulate data

## Description

Calling function to simulate data

## Usage

genData(n, dtDefs = NULL, id = "id", envir = parent.frame())

## Arguments

n
the number of observations required in the data set.
dtDefs
id
name of definitions data.table/data.frame. If no definitions are provided a data set with ids only is generated.
The string defining the id of the record. Will override previously set id name with a warning (unless the old value is 'id'). If the id attribute in dtDefs is NULL will default to 'id'.
envir Environment the data definitions are evaluated in. Defaults to base::parent.frame.

## Value

A data.table that contains the simulated data.

## Examples

```
genData(5)
genData(5, id = "grpID")
def <- defData(
    varname = "xNr", dist = "nonrandom", formula = 7,
    id = "idnum"
)
def <- defData(def,
    varname = "xUni", dist = "uniform",
    formula = "10;20"
)
def <- defData(def,
    varname = "xNorm", formula = "xNr + xUni * 2",
    dist = "normal", variance = 8
)
def <- defData(def,
    varname = "xPois", dist = "poisson",
    formula = "xNr - 0.2 * xUni", link = "log"
)
def <- defData(def,
    varname = "xCat", formula = "0.3;0.2;0.5",
    dist = "categorical"
)
def <- defData(def,
    varname = "xGamma", dist = "gamma", formula = "5+xCat",
    variance = 1, link = "log"
)
def <- defData(def,
    varname = "xBin", dist = "binary", formula = "-3 + xCat",
    link = "logit"
)
def
genData(5, def)
```

genDummy Create dummy variables from a factor or integer variable

## Description

Create dummy variables from a factor or integer variable

## Usage

genDummy (dtName, varname, sep = ".", replace = FALSE)

## Arguments

| dtName | Data table with column |
| :--- | :--- |
| varname | Name of factor |
| sep | Character to be used in creating new name for dummy fields. Valid characters <br> include all letters and "_. Will default to ".". If an invalid character is provided, <br> it will be replaced by default. |
| replace | If replace is set to TRUE (defaults to FALSE) the field referenced varname will <br> be removed. |

## Examples

```
# First example:
def <- defData(varname = "cat", formula = ".2;.3;.5", dist = "categorical")
def <- defData(def, varname = "x", formula = 5, variance = 2)
dx <- genData(200, def)
dx
dx <- genFactor(dx, "cat", labels = c("one", "two", "three"), replace = TRUE)
dx <- genDummy(dx, varname = "fcat", sep = "_")
dx
# Second example:
dx <- genData(15)
dx <- trtAssign(dtName = dx, 3, grpName = "arm")
dx <- genDummy(dx, varname = "arm")
dx
```

```
    genFactor Create factor variable from an existing (non-double) variable
```


## Description

Create factor variable from an existing (non-double) variable

## Usage

genFactor (dtName, varname, labels = NULL, prefix = "f", replace = FALSE)

## Arguments

dtName Data table with columns.
varname $\quad$ Name of field(s) to be converted.
labels Factor level labels. If not provided, the generated factor levels will be used as the labels. Can be a vector (if only one new factor or all factors have the same labels) or a list of character vectors of the same length as varname.
prefix By default, the new field name will be a concatenation of " $f$ " and the old field name. A prefix string can be provided.
replace If replace is set to TRUE (defaults to FALSE) the field referenced varname will be removed.

## Examples

```
# First example:
def <- defData(varname = "cat", formula = ".2;.3;.5", dist = "categorical")
def <- defData(def, varname = "x", formula = 5, variance = 2)
dx <- genData(200, def)
dx
dx <- genFactor(dx, "cat", labels = c("one", "two", "three"))
dx
# Second example:
dx <- genData(10)
dx <- trtAssign(dtName = dx, 2, grpName = "studyArm")
dx <- genFactor(dx, varname = "studyArm", labels = c("control", "treatment"), prefix = "t_")
dx
```


## Description

Formulas for additive linear models can be generated with specified coefficient values and variable names.

## Usage

genFormula(coefs, vars)

## Arguments

coefs A vector that contains the values of the coefficients. Coefficients can also be defined as character for use with double dot notation. If length(coefs) $==$ length(vars), then no intercept is assumed. Otherwise, an intercept is assumed.
vars A vector of strings that specify the names of the explanatory variables in the equation.

## Value

A string that represents the desired formula

## Examples

```
genFormula(c(.5, 2, 4), c("A", "B", "C"))
genFormula(c(.5, 2, 4), c("A", "B"))
genFormula(c(.5, "..x", 4), c("A", "B", "C"))
genFormula(c(.5, 2, "..z"), c("A", "B"))
changeX <- c(7, 10)
genFormula(c(.5, 2, changeX[1]), c("A", "B"))
genFormula(c(.5, 2, changeX[2]), c("A", "B"))
genFormula(c(.5, 2, changeX[2]), c("A", "B", "C"))
newForm <- genFormula(c(-2, 1), c("A"))
def1 <- defData(varname = "A", formula = 0, variance = 3, dist = "normal")
def1 <- defData(def1, varname = "B", formula = newForm, dist = "binary", link = "logit")
set.seed(2001)
dt <- genData(500, def1)
summary(glm(B ~ A, data = dt, family = binomial))
```


## genMarkov Generate Markov chain

## Description

Generate a Markov chain for n individuals or units by specifying a transition matrix.

```
Usage
    genMarkov(
        n,
        transMat,
        chainLen,
        wide = FALSE,
        id = "id",
        pername = "period",
        varname = "state",
        widePrefix = "S",
        trimvalue = NULL,
        startProb = NULL
    )
```


## Arguments

n
transMat
chainLen Length of each chain that will be generated for each chain; minimum chain length is 2.
wide Logical variable (TRUE or FALSE) indicating whether the resulting data table should be returned in wide or long format. The wide format includes all elements of a chain on a single row; the long format includes each element of a chain in its own row. The default is wide = FALSE, so the long format is returned by default.
id Character string that represents name of "id" field. Defaults to "id".
pername Character string that represents the variable name of the chain sequence in the long format. Defaults "period",
varname Character string that represents the variable name of the state in the long format. Defaults to "state".
widePrefix Character string that represents the variable name prefix for the state fields in the wide format. Defaults to "S".
trimvalue Integer value indicating end state. If trimvalue is not NULL, all records after the first instance of state $=$ trimvalue will be deleted .
startProb A string that contains the probability distribution of the starting state, separated by a ";". Length of start probabilities must match the number of rows of the transition matrix.

## Value

A data table with n rows if in wide format, or n by chainLen rows if in long format.

## Examples

\# Transition matrix $P$
$P<-t$ (matrix $(c)$
$0.7,0.2,0.1$,
$0.5,0.3,0.2$,
$0.0,0.1,0.9$
), nrow $=3$, ncol = 3))
d1 <- genMarkov( $n=10$, transMat $=P$, chainLen $=5$ )
$\mathrm{d} 2<-\operatorname{genMarkov}(\mathrm{n}=10$, transMat $=P$, chainLen $=5$, wide $=$ TRUE)
d3 <- genMarkov(
$\mathrm{n}=10$, transMat $=P$, chainLen $=5$,
pername = "seq", varname = "health",
trimvalue $=3$
)
genMiss Generate missing data

## Description

Generate missing data

## Usage

genMiss( dtName, missDefs, idvars, repeated = FALSE, periodvar = "period", envir = parent.frame()
)

## Arguments

| dtName | Name of complete data set |
| :--- | :--- |
| missDefs | Definitions of missingness |
| idvars | Index variables |
| repeated | Indicator for longitudinal data |
| periodvar | Name of variable that contains period |
| envir | parent.frame() by default, allows functionality with double-dot notation |

## Value

Missing data matrix indexed by idvars (and period if relevant)

## See Also

```
defMiss,genObs
```


## Examples

```
def1 <- defData(varname = "m", dist = "binary", formula = .5)
def1 <- defData(def1, "u", dist = "binary", formula = . 5)
def1 <- defData(def1, "x1", dist = "normal", formula = "20*m + 20*u", variance = 2)
def1 <- defData(def1, "x2", dist = "normal", formula = "20*m + 20*u", variance = 2)
def1 <- defData(def1, "x3", dist = "normal", formula = "20*m + 20*u", variance = 2)
dtAct <- genData(1000, def1)
defM <- defMiss(varname = "x1", formula = .15, logit.link = FALSE)
defM <- defMiss(defM, varname = "x2", formula = ".05 + m * 0.25", logit.link = FALSE)
defM <- defMiss(defM, varname = "x3", formula = ".05 + u * 0.25", logit.link = FALSE)
defM <- defMiss(defM, varname = "u", formula = 1, logit.link = FALSE) # not observed
defM
# Generate missing data matrix
missMat <- genMiss(dtAct, defM, idvars = "id")
missMat
# Generate observed data from actual data and missing data matrix
dtObs <- genObs(dtAct, missMat, idvars = "id")
dtObs
```

genMixFormula Generate Mixture Formula

## Description

Generates a mixture formula from a vector of variable names and an optional vector of probabilities.

## Usage

genMixFormula(vars, probs = NULL, varLength = NULL)

## Arguments

vars Character vector/list of variable names.
probs Numeric vector/list of probabilities. Has to be same length as vars or NULL. Probabilities will be normalized if the sum to $>1$.
varLength If vars is of length one and varLength is set to any integer $>0$, vars will be interpreted as array of length varLength and all elements will used in sequence.

## Value

The mixture formula as a string.

## Examples

```
genMixFormula(c("a", "..b[..i]", "c"))
genMixFormula(c("a", "..b", "c"), c(.2, .5, .3))
# Shorthand to use external vectors/lists
genMixFormula("..arr", varLength = 5)
```

```
genMultiFac Generate multi-factorial data
```


## Description

Generate multi-factorial data

## Usage

```
genMultiFac(
    nFactors,
    each,
    levels = 2,
    coding = "dummy",
    colNames = NULL,
    idName = "id"
)
```


## Arguments

| nFactors | Number of factors (columns) to generate. <br> each <br> Number of replications for each combination of factors. Must be specified. |
| :--- | :--- |
| levels | Vector or scalar. If a vector is specified, it must be the same length as nFatctors. <br> Each value of the vector represents the number of levels of each corresponding <br> factor. If a scalar is specified, each factor will have the same number of levels. <br> The default is 2 levels for each factor. <br> coding |
| String value to specify if "dummy" or "effect" coding is used. Defaults to <br> "dummy". |  |
| colNames | A vector of strings, with a length of nFactors. The strings represent the name <br> for each factor. |
| idName | A string that specifies the id of the record. Defaults to "id". |

## Value

A data.table that contains the added simulated data. Each column contains an integer.

## Examples

```
genMultiFac(nFactors = 2, each = 5)
genMultiFac(nFactors = 2, each = 4, levels = c(2, 3))
genMultiFac(
        nFactors = 3, each = 1, coding = "effect",
        colNames = c("Fac1", "Fac2", "Fac3"), id = "block"
    )
```

genNthEvent

Generate event data using longitudinal data, and restrict output to time until the nth event.

## Description

Generate event data using longitudinal data, and restrict output to time until the nth event.

## Usage

genNthEvent(dtName, defEvent, nEvents = 1, perName = "period", id = "id")

## Arguments

| dtName | name of existing data table |
| :--- | :--- |
| defEvent | data definition table (created with defDataAdd) that determines the event gener- <br> ating process. |
| nEvents | maximum number of events that will be generated (the nth event). |
| perName | variable name for period field. Defaults to "period" |
| id | string representing name of the id field in table specified by dtName |

## Value

data.table that stops after "nEvents" are reached.

## Examples

```
defD <- defData(
    varname = "effect", formula = 0, variance = 1,
    dist = "normal"
)
defE <- defDataAdd(
    varname = "died", formula = "-2.5 + 0.3*period + effect",
    dist = "binary", link = "logit"
)
```

d <- genData(1000, defD)
d <- addPeriods(d, 10)
$\mathrm{dx}<-$ genNthEvent( d , defEvent $=$ defE, nEvents $=3$ )

## genObs $\quad$ Create an observed data set that includes missing data

## Description

Create an observed data set that includes missing data

## Usage

genObs(dtName, dtMiss, idvars)

## Arguments

| $d t$ Name | Name of complete data set |
| :--- | :--- |
| $d t M i s s$ | Name of missing data matrix |
| idvars | Index variables that cannot be missing |

## Value

A data table that represents observed data, including missing data

## See Also

defMiss, genMiss

## Examples

```
def1 <- defData(varname = "m", dist = "binary", formula = .5)
def1 <- defData(def1, "u", dist = "binary", formula = .5)
def1 <- defData(def1, "x1", dist = "normal", formula = "20*m + 20*u", variance = 2)
def1 <- defData(def1, "x2", dist = "normal", formula = "20*m + 20*u", variance = 2)
def1 <- defData(def1, "x3", dist = "normal", formula = "20*m + 20*u", variance = 2)
dtAct <- genData(1000, def1)
defM <- defMiss(varname = "x1", formula = .15, logit.link = FALSE)
defM <- defMiss(defM, varname = "x2", formula = ".05 + m * 0.25", logit.link = FALSE)
defM <- defMiss(defM, varname = "x3", formula = ".05 + u * 0.25", logit.link = FALSE)
defM <- defMiss(defM, varname = "u", formula = 1, logit.link = FALSE) # not observed
defM
# Generate missing data matrix
missMat <- genMiss(dtAct, defM, idvars = "id")
```

missMat
\# Generate observed data from actual data and missing data matrix
dtObs <- genObs(dtAct, missMat, idvars = "id")
dtObs
genOrdCat Generate ordinal categorical data

## Description

Ordinal categorical data is added to an existing data set. Correlations can be added via correlation matrix or rho and corstr.

## Usage

genOrdCat
dtName, adjVar = NULL, baseprobs, catVar = "cat", asFactor = TRUE, idname = "id", prefix = "grp", rho = 0, corstr = "ind", corMatrix = NULL, npVar = NULL, npAdj = NULL
)

## Arguments

dtName Name of complete data set
adjVar Adjustment variable name in dtName - determines logistic shift. This is specified assuming a cumulative logit link.
baseprobs Baseline probability expressed as a vector or matrix of probabilities. The values (per row) must sum to $<=1$. If rowSums(baseprobs) $<1$, an additional category is added with probability 1 - rowSums(baseprobs). The number of rows represents the number of new categorical variables. The number of columns represents the number of possible responses - if an particular category has fewer possible responses, assign zero probability to non-relevant columns.
catVar Name of the new categorical field. Defaults to "cat". Can be a character vector with a name for each new variable defined via baseprobs. Will be overridden by prefix if more than one variable is defined and length(catVar) $=1$.

| asFactor | If asFactor $==$ TRUE (default), new field is returned as a factor. If asFactor $==$ <br> FALSE, new field is returned as an integer. |
| :--- | :--- |
| idname | Name of the id column in dtName. |
| prefix | A string. The names of the new variables will be a concatenation of the prefix <br> and a sequence of integers indicating the variable number. |
| corstr | Correlation coefficient, $-1<$ rho < 1. Use if corMatrix is not provided. <br> Correlation structure of the variance-covariance matrix defined by sigma and <br> rho. Options include "ind" for an independence structure, "cs" for a compound <br> symmetry structure, and "ar1" for an autoregressive structure. |
| corMatrix | Correlation matrix can be entered directly. It must be symmetrical and positive <br> definite. It is not a required field; if a matrix is not provided, then a structure and <br> correlation coefficient rho must be specified. (The matrix created via rho and <br> corstr must also be positive definite.) |
| npVar | Vector of variable names that indicate which variables are to violate the propor- <br> tionality assumption. |
| npAdj | Matrix with a row for each npVar and a column for each category. Each value <br> represents the deviation from the proportional odds assumption on the logistic <br> scale. |

## Value

Original data.table with added categorical field.

## Examples

```
# Ordinal Categorical Data ----
def1 <- defData(
    varname = "male",
    formula = 0.45, dist = "binary", id = "idG"
)
def1 <- defData(def1,
    varname = "z",
    formula = "1.2*male", dist = "nonrandom"
)
def1
## Generate data
set.seed(20)
dx <- genData(1000, def1)
probs <- c(0.40, 0.25, 0.15)
dx <- genOrdCat(dx,
    adjVar = "z", idname = "idG", baseprobs = probs,
    catVar = "grp"
)
```

```
dx
# Correlated Ordinal Categorical Data ----
baseprobs <- matrix(c(
    0.2, 0.1, 0.1, 0.6,
    0.7, 0.2, 0.1, 0,
    0.5, 0.2, 0.3, 0,
    0.4, 0.2, 0.4, 0,
    0.6, 0.2, 0.2, 0
),
nrow = 5, byrow = TRUE
)
set.seed(333)
dT <- genData(1000)
dX <- genOrdCat(dT,
        adjVar = NULL, baseprobs = baseprobs,
    prefix = "q", rho = .125, corstr = "cs", asFactor = FALSE
)
dX
dM <- data.table::melt(dX, id.vars = "id")
dProp <- dM[, prop.table(table(value)), by = variable]
dProp[, response := c(1:4, 1:3, 1:3, 1:3, 1:3)]
data.table::dcast(dProp, variable ~ response,
    value.var = "V1", fill = 0
)
# proportional odds assumption violated
d1 <- defData(varname = "rx", formula = "1;1", dist = "trtAssign")
d1 <- defData(d1, varname = "z", formula = "0 - 1.2*rx", dist = "nonrandom")
dd <- genData(1000, d1)
baseprobs <- c(.4, .3, .2, .1)
npAdj <- c(0, 1, 0, 0)
dn <- genOrdCat(
    dtName = dd, adjVar = "z",
    baseprobs = baseprobs,
    npVar = "rx", npAdj = npAdj
)
```

    genSpline Generate spline curves
    
## Description

Generate spline curves

## Usage

```
genSpline(
    dt,
    newvar,
    predictor,
    theta,
    knots = c(0.25, 0.5, 0.75),
    degree = 3,
    newrange = NULL,
    noise.var = 0
)
```


## Arguments

| dt | data.table that will be modified |
| :--- | :--- |
| newvar | Name of new variable to be created |
| predictor | Name of field in old data.table that is predicting new value |
| theta | A vector or matrix of values between 0 and 1. Each column of the matrix rep- <br> resents the weights/coefficients that will be applied to the basis functions de- <br> termined by the knots and degree. Each column of theta represents a separate <br> spline curve. |
| knots | A vector of values between 0 and 1, specifying quantile cut-points for splines. <br> Defaults to c(0.25, 0.50, 0.75$).$ |
| degree | Integer specifying polynomial degree of curvature. <br> newrange |
| Range of the spline function, specified as a string with two values separated <br> by a semi-colon. The first value represents the minimum, and the second value <br> represents the maximum. Defaults to NULL, which sets the range to be between <br> 0 and 1. |  |
| noise.var | Add to normally distributed noise to observation - where mean is value of spline <br> curve. |

## Value

A modified data.table with an added column named newvar.

## Examples

```
ddef <- defData(varname = "age", formula = "0;1", dist = "uniform")
theta1 <- c(0.1, 0.8, 0.6, 0.4, 0.6, 0.9, 0.9)
knots <- c(0.25, 0.5, 0.75)
viewSplines(knots = knots, theta = theta1, degree = 3)
```

```
set.seed(234)
dt <- genData(1000, ddef)
dt <- genSpline(
    dt = dt, newvar = "weight",
    predictor = "age", theta = theta1,
    knots = knots, degree = 3,
    noise.var = . 025
)
dt
```

    genSurv Generate survival data
    
## Description

Survival data is added to an existing data set.

## Usage

```
    genSurv(
        dtName,
        survDefs,
        digits = 3,
        timeName = NULL,
        censorName = NULL,
        eventName = "event",
        typeName = "type",
        keepEvents = FALSE,
        idName = "id"
    )
```


## Arguments

| dtName | Name of data set |
| :--- | :--- |
| survDefs | Definitions of survival |
| digits | Number of digits for rounding |
| timeName | A string to indicate the name of a combined competing risk time-to-event out- <br> come that reflects the minimum observed value of all time-to-event outcomes. <br> Defaults to NULL, indicating that each time-to-event outcome will be included <br> in dataset. |
| censorName | The name of a time to event variable that is the censoring variable. Will be <br> ignored if timeName is NULL. |


| eventName | The name of the new numeric/integer column representing the competing event <br> outcomes. If censorName is specified, the integer value for that event will be 0. <br> Defaults to "event", but will be ignored if timeName is NULL. |
| :--- | :--- |
| typeName | The name of the new character column that will indicate the event type. The <br> type will be the unique variable names in survDefs. Defaults to "type", but will <br> be ignored if timeName is NULL. |
| keepEvents | Indicator to retain original "events" columns. Defaults to FALSE. |
| idName | Name of id field in existing data set. |

## Value

Original data table with survival time

## Examples

```
# Baseline data definitions
def <- defData(varname = "x1", formula = .5, dist = "binary")
def <- defData(def, varname = "x2", formula = .5, dist = "binary")
def <- defData(def, varname = "grp", formula = .5, dist = "binary")
# Survival data definitions
sdef <- defSurv(
    varname = "survTime", formula = "1.5*x1",
    scale = "grp*50 + (1-grp)*25", shape = "grp*1 + (1-grp)*1.5"
)
sdef <- defSurv(sdef, varname = "censorTime", scale = 80, shape = 1)
sdef
# Baseline data definitions
dtSurv <- genData(300, def)
# Add survival times
dtSurv <- genSurv(dtSurv, sdef)
head(dtSurv)
```

genSynthetic Generate synthetic data

## Description

Synthetic data is generated from an existing data set

## Usage

genSynthetic(dtFrom, $\mathrm{n}=\mathrm{nrow}(\mathrm{dtFrom})$, vars $=$ NULL, $\mathrm{id}=$ "id")

## Arguments

dtFrom Data table that contains the source data
n
Number of samples to draw from the source data. The default is number of records that are in the source data file.
vars A vector of string names specifying the fields that will be sampled. The default is that all variables will be selected.
id A string specifying the field that serves as the record id. The default field is "id".

## Value

A data table with the generated data

## Examples

```
### Create fake "real" data set
d <- defData(varname = "a", formula = 3, variance = 1, dist = "normal")
d <- defData(d, varname = "b", formula = 5, dist = "poisson")
d <- defData(d, varname = "c", formula = 0.3, dist = "binary")
d <- defData(d, varname = "d", formula = "a + b + 3*c", variance = 2, dist = "normal")
A <- genData(100, d, id = "index")
### Create synthetic data set from "observed" data set A:
def <- defDataAdd(varname = "x", formula = " 2*b + 2*d", variance = 2)
S <- genSynthetic(dtFrom = A, n = 120, vars = c("b", "d"), id = "index")
S <- addColumns(def, S)
```

iccRE Generate variance for random effects that produce desired intra-class coefficients (ICCs) for clustered data.

## Description

Generate variance for random effects that produce desired intra-class coefficients (ICCs) for clustered data.

## Usage

iccRE(ICC, dist, varTotal = NULL, varWithin = NULL, lambda = NULL, disp = NULL)

## Arguments

| ICC | Vector of values between 0 and 1 that represent the target ICC levels |
| :--- | :--- |
| dist | The distribution that describes the outcome data at the individual level. Possible <br> distributions include "normal", "binary", "poisson", or "gamma" |
| varTotal | Numeric value that represents the total variation for a normally distributed model. <br> If "normal" distribution is specified, either varTotal or varWithin must be speci- <br> fied, but not both. |
| varWithin | Numeric value that represents the variation within a cluster for a normally dis- <br> tributed model. If "normal" distribution is specified, either varTotal or varWithin <br> must be specified, but not both. |
| lambda | Numeric value that represents the grand mean. Must be specified when distribu- <br> tion is "poisson" or "negative binomial". |
| disp | Numeric value that represents the dispersion parameter that is used to define a <br> gamma or negative binomial distribution with a log link. Must be specified when <br> distribution is "gamma". |

## Value

A vector of values that represents the variances of random effects at the cluster level that correspond to the ICC vector.

## References

Nakagawa, Shinichi, and Holger Schielzeth. "A general and simple method for obtaining R2 from generalized linear mixed-effects models." Methods in ecology and evolution 4, no. 2 (2013): 133142.

## Examples

```
targetICC <- seq(0.05, 0.20, by = .01)
iccRE(targetICC, "poisson", lambda = 30)
iccRE(targetICC, "binary")
iccRE(targetICC, "normal", varTotal = 100)
iccRE(targetICC, "normal", varWithin = 100)
iccRE(targetICC, "gamma", disp = .5)
iccRE(targetICC, "negBinomial", lambda = 40, disp = .5)
```

logisticCoefs Determine intercept, treatment/exposure and covariate coefficients that can be used for binary data generation with a logit link and a set of covariates

## Description

This is an implementation of an iterative bisection procedure that can be used to determine coefficient values for a target population prevalence as well as a target risk ratio, risk difference, or AUC. These coefficients can be used in a subsequent data generation process to simulate data with these desire characteristics.

## Usage

logisticCoefs(
defCovar, coefs, popPrev,
rr = NULL,
$r d=N U L L$,
auc $=$ NULL,
tolerance $=0.001$,
sampleSize $=1 \mathrm{e}+05$,
trtName = "A"
)

## Arguments

defCovar A definition table for the covariates in the underlying population. This tables specifies the distribution of the covariates.
coefs A vector of coefficients that reflect the relationship between each of the covariates and the log-odds of the outcome.
popPrev The target population prevalence of the outcome. A value between 0 and 1 .
rr
The target risk ratio, which must be a value between 0 and 1/popPrev. Defaults to NULL.
rd The target risk difference, which must be between -(popPrev) and (1-popPrev). Defaults to NULL
auc The target AUC, which must be a value between 0.5 and 1.0 . Defaults to NULL.
tolerance The minimum stopping distance between the adjusted low and high endpoints. Defaults to 0.001 .
sampleSize The number of units to generate for the bisection algorithm. The default is $1 \mathrm{e}+05$. To get a reliable estimate, the value should be no smaller than the default, though larger values can be used, though computing time will increase.
trtName If either a risk ratio or risk difference is the target statistic, a treatment/exposure variable name can be provided. Defaults to "A".

## Details

If no specific target statistic is specified, then only the intercept is returned along with the original coefficients. Only one target statistic (risk ratio, risk difference or AUC) can be specified with a single function call; in all three cases, a target prevalence is still required.

## Value

A vector of parameters including the intercept and covariate coefficients for the logistic model data generating process.

## References

Austin, Peter C. "The iterative bisection procedure: a useful tool for determining parameter values in data-generating processes in Monte Carlo simulations." BMC Medical Research Methodology 23, no. 1 (2023): 1-10.

## Examples

```
## Not run:
d1 <- defData(varname = "x1", formula = 0, variance = 1)
d1 <- defData(d1, varname = "b1", formula = 0.5, dist = "binary")
coefs <- log(c(1.2, 0.8))
logisticCoefs(d1, coefs, popPrev = 0.20)
logisticCoefs(d1, coefs, popPrev = 0.20, rr = 1.50, trtName = "rx")
logisticCoefs(d1, coefs, popPrev = 0.20, rd = 0.30, trtName = "rx")
logisticCoefs(d1, coefs, popPrev = 0.20, auc = 0.80)
## End(Not run)
```

mergeData Merge two data tables

## Description

Merge two data tables

## Usage

mergeData(dt1, dt2, idvars)

## Arguments

## dt1

Name of first data.table
dt2 Name of second data.table
idvars Vector of string names to merge on

## Value

A new data table that merges dt 2 with dt1

## Examples

```
def1 <- defData(varname = "x", formula = 0, variance = 1)
def1 <- defData(varname = "xcat", formula = ".3;.2", dist = "categorical")
def2 <- defData(varname = "yBin", formula = 0.5, dist = "binary", id = "xcat")
def2 <- defData(def2, varname = "yNorm", formula = 5, variance = 2)
dt1 <- genData(20, def1)
dt2 <- genData(3, def2)
dtMerge <- mergeData(dt1, dt2, "xcat")
dtMerge
```

negbinomGetSizeProb Convert negative binomial mean and dispersion parameters to size and prob parameters

## Description

Convert negative binomial mean and dispersion parameters to size and prob parameters

## Usage

negbinomGetSizeProb(mean, dispersion)

## Arguments

mean The mean of a gamma distribution
dispersion The dispersion parameter of a gamma distribution

## Details

In simstudy, users specify the negative binomial distribution as a function of two parameters - a mean and dispersion. In this case, the variance of the specified distribution is mean + (mean^2)*dispersion. The base R function rnbinom uses the size and prob parameters to specify the negative binomial distribution. This function converts the mean and dispersion into the size and probability parameters.

## Value

A list that includes the size and prob parameters of the neg binom distribution

## Examples

```
set.seed(12345)
mean <- 5
dispersion <- 0.5
sp <- negbinomGetSizeProb(mean, dispersion)
c(sp$size, sp$prob)
vec <- rnbinom(1000, size = sp$size, prob = sp$prob)
(estMoments <- c(mean(vec), var(vec)))
(theoryMoments <- c(mean, mean + mean^2 * dispersion))
(theoryMoments <- c(sp$size * (1 - sp$prob) / sp$prob, sp$size * (1 - sp$prob) / sp$prob^2))
```

```
simstudy-deprecated Deprecated functions in simstudy
```


## Description

These functions are provided for compatibility with older versions of simstudy only, and will be defunct in the future.

## Details

- genCorOrdCat: This function is deprecated, and will be removed in the future. Use genOrdCat with asFactor $=$ FALSE instead.
- catProbs: This function is deprecated, and will be removed in the future. Use genCatFormula with the same functionality instead.

```
survGetParams Get survival curve parameters
```


## Description

Get survival curve parameters

## Usage

survGetParams(points)

## Arguments

points A list of two-element vectors specifying the desired time and probability pairs that define the desired survival curve

## Value

A vector of parameters that define the survival curve optimized for the target points. The first element of the vector represents the " $f$ " parameter and the second element represents the "shape" parameter.

## Examples

```
points <- list(c(60, 0.90), c(100, . 75), c(200, . 25), c(250, . 10))
survGetParams(points)
```

```
survParamPlot Plot survival curves
```


## Description

Plot survival curves

## Usage

survParamPlot(formula, shape, points $=$ NULL, $n=100$, scale = 1, limits $=$ NULL)

## Arguments

| formula | This is the "formula" parameter of the Weibull-based survival curve that can be <br> used to define the scale of the distribution. |
| :--- | :--- |
| shape | The parameter that defines the shape of the distribution. <br> points |
| An optional list of two-element vectors specifying the desired time and proba- |  |
| bility pairs that define the desired survival curve. If no list is specified then the |  |
| plot will not include any points. |  |
| The number of points along the curve that will be used to define the line. De- |  |
| faults to 100 . |  |

## Value

A ggplot of the survival curve defined by the specified parameters. If the argument points is specified, the plot will include them

## Examples

```
points <- list(c(60, 0.90), c(100, .75), c(200, .25), c(250, .10))
r <- survGetParams(points)
survParamPlot(r[1], r[2])
survParamPlot(r[1], r[2], points = points)
survParamPlot(r[1], r[2], points = points, limits = c(0, 100))
```


## Description

Trim longitudinal data file once an event has occurred

## Usage

trimData(dt0ld, seqvar, eventvar, idvar = "id")

## Arguments

| dt0ld | name of data table to be trimmed |
| :--- | :--- |
| seqvar | string referencing column that indexes the sequence or period |
| eventvar | string referencing event data column |
| idvar | string referencing id column |

## Value

an updated data.table removes all rows following the first event for each individual

## Examples

```
eDef <- defDataAdd(varname = "e", formula = "u==4", dist = "nonrandom")
P <- t(matrix(c(
    0.4, 0.3, 0.2, 0.1,
    0.0, 0.4, 0.3, 0.3,
    0.0, 0.0, 0.5, 0.5,
    0.0, 0.0, 0.0, 1.0
),
nrow = 4
))
dp <- genMarkov(
    n = 100, transMat = P,
    chainLen = 8, id = "id",
    pername = "period",
    varname = "u"
)
dp <- addColumns(eDef, dp)
dp <- trimData(dp, seqvar = "period", eventvar = "e", idvar = "id")
dp
```


## trtAssign Assign treatment

## Description

Assign treatment

## Usage

```
trtAssign(
    dtName,
    nTrt = 2,
    balanced = TRUE,
    strata = NULL,
    grpName = "trtGrp",
    ratio = NULL
    )
```


## Arguments

| dtName | data table |
| :--- | :--- |
| nTrt | number of treatment groups |
| balanced | indicator for treatment assignment process |
| strata | vector of strings representing stratifying variables |
| grpName | string representing variable name for treatment or exposure group |
| ratio | vector of values indicating relative proportion of group assignment |

## Value

An integer (group) ranging from 1 to length of the probability vector

## See Also

trtObserve

## Examples

```
dt <- genData(15)
dt1 <- trtAssign(dt, nTrt = 3, balanced = TRUE)
dt1[, .N, keyby = trtGrp]
dt2 <- trtAssign(dt, nTrt = 3, balanced = FALSE)
dt2[, .N, keyby = trtGrp]
def <- defData(varname = "male", formula = .4, dist = "binary")
dt <- genData(1000, def)
```

```
dt
    dt3 <- trtAssign(dt, nTrt = 5, strata = "male", balanced = TRUE, grpName = "Group")
dt3
dt3[, .N, keyby = .(male, Group)]
dt3[, .N, keyby = .(Group)]
dt4 <- trtAssign(dt, nTrt = 5, strata = "male", balanced = FALSE, grpName = "Group")
dt4[, .N, keyby = .(male, Group)]
dt4[, .N, keyby = .(Group)]
dt5 <- trtAssign(dt, nTrt = 5, balanced = TRUE, grpName = "Group")
dt5[, .N, keyby = .(male, Group)]
dt5[, .N, keyby = .(Group)]
dt6 <- trtAssign(dt, nTrt = 3, ratio = c(1, 2, 2), grpName = "Group")
dt6[, .N, keyby = .(Group)]
```

```
trtObserve Observed exposure or treatment
```


## Description

Observed exposure or treatment

## Usage

trtObserve(dt, formulas, logit.link = FALSE, grpName = "trtGrp")

## Arguments

| dt | data table |
| :--- | :--- |
| formulas | collection of formulas that determine probabilities |
| logit.link | indicator that specifies link. If TRUE, then logit link is used. If FALSE, the <br> identity link is used. |
| grpName | character string representing name of treatment/exposure group variable |

## Value

An integer (group) ranging from 1 to length of the probability vector

## See Also

```
    trtAssign
```


## Examples

```
def <- defData(varname = "male", dist = "binary", formula = .5, id = "cid")
def <- defData(def, varname = "over65", dist = "binary", formula = "-1.7 + .8*male", link = "logit")
def <- defData(def, varname = "baseDBP", dist = "normal", formula = 70, variance = 40)
dtstudy <- genData(1000, def)
dtstudy
formula1 <- c("-2 + 2*male - .5*over65", "-1 + 2*male + .5*over65")
dtObs <- trtObserve(dtstudy, formulas = formula1, logit.link = TRUE, grpName = "exposure")
dtObs
# Check actual distributions
dtObs[, .(pctMale = round(mean(male), 2)), keyby = exposure]
dtObs[, .(pctMale = round(mean(over65), 2)), keyby = exposure]
dtSum <- dtObs[, .N, keyby = .(male, over65, exposure)]
dtSum[, grpPct := round(N / sum(N), 2), keyby = .(male, over65)]
dtSum
```

trtStepWedge Assign treatment for stepped-wedge design

## Description

Assign treatment for stepped-wedge design

## Usage

```
trtStepWedge(
    dtName,
    clustID,
    nWaves,
    lenWaves,
    startPer,
    perName = "period",
    grpName = "rx",
    lag = 0,
    xrName = "xr"
)
```


## Arguments

dtName data table
clustID string representing name of column of cluster level ids
nWaves number of treatment waves

| lenWaves | the number of periods between waves |
| :--- | :--- |
| startPer | the starting period of the first wave |
| perName | string representing name of column of time periods |
| grpName | string representing variable name for treatment or exposure group <br> lag |
| integer representing length of transition period |  |

## Value

A data.table with the added treatment assignment

## See Also

trtObserve trtAssign

## Examples

```
defc <- defData(
    varname = "ceffect", formula = 0, variance = 0.10,
    dist = "normal", id = "cluster"
)
defc <- defData(defc, "m", formula = 10, dist = "nonrandom")
# Will generate 3 waves of 4 clusters each - starting 2, 5, and 8
dc <- genData(12, defc)
dp <- addPeriods(dc, 12, "cluster")
dp <- trtStepWedge(dp, "cluster",
    nWaves = 3,
    lenWaves = 3, startPer = 2
)
dp
dp <- addPeriods(dc, 12, "cluster")
dp <- trtStepWedge(dp, "cluster",
    nWaves = 2,
    lenWaves = 1, startPer = 4, lag = 3
)
dp
```

updateDef Update definition table

## Description

Updates row definition table created by function defData or defRead. (For tables created using defDataAdd and defReadAdd use updateDefAdd.) Does not modify in-place.

## Usage

```
updateDef(
        dtDefs,
        changevar,
        newformula \(=\) NULL,
        newvariance = NULL,
        newdist = NULL,
        newlink = NULL,
        remove \(=\) FALSE
)
```


## Arguments

| dtDefs | Definition table that will be modified |
| :--- | :--- |
| changevar | Name of field definition that will be changed |
| newformula | New formula definition (defaults to NULL) |
| newvariance | New variance specification (defaults to NULL) |
| newdist | New distribution definition (defaults to NULL) |
| newlink | New link specification (defaults to NULL) |
| remove | If set to TRUE, remove 'changevar'from definition (defaults to FALSE). |

## Value

The updated data definition table.

## Examples

```
# Example 1
defs <- defData(varname = "x", formula = 0, variance = 3, dist = "normal")
defs <- defData(defs, varname = "y", formula = "2 + 3*x", variance = 1, dist = "normal")
defs <- defData(defs, varname = "z", formula = "4 + 3*x - 2*y", variance = 1, dist = "normal")
defs
updateDef(dtDefs = defs, changevar = "y", newformula = "x + 5", newvariance = 2)
updateDef(dtDefs = defs, changevar = "z", newdist = "poisson", newlink = "log")
# Example 2
defs <- defData(varname = "w", formula = 0, variance = 3, dist = "normal")
defs <- defData(defs, varname = "x", formula = "1 + w", variance = 1, dist = "normal")
defs <- defData(defs, varname = "z", formula = 4, variance = 1, dist = "normal")
defs
updateDef(dtDefs = defs, changevar = "x", remove = TRUE)
updateDef(dtDefs = defs, changevar = "z", remove = TRUE)
```

\# No changes to original definition:
defs
updateDefAdd Update definition table

## Description

Updates row definition table created by functions defDataAdd and defReadAdd. (For tables created using defData or defRead use updateDef.)

## Usage

```
    updateDefAdd(
        dtDefs,
        changevar,
        newformula = NULL,
        newvariance = NULL,
        newdist = NULL,
        newlink = NULL,
        remove = FALSE
    )
```


## Arguments

| dtDefs | Definition table that will be modified |
| :--- | :--- |
| changevar | Name of field definition that will be changed |
| newformula | New formula definition (defaults to NULL) |
| newvariance | New variance specification (defaults to NULL) |
| newdist | New distribution definition (defaults to NULL) |
| newlink | New link specification (defaults to NULL) |
| remove | If set to TRUE, remove definition (defaults to FALSE) |

## Value

A string that represents the desired formula

## Examples

```
# Define original data
defs <- defData(varname = "w", formula = 0, variance = 3, dist = "normal")
defs <- defData(defs, varname = "x", formula = "1 + w", variance = 1, dist = "normal")
defs <- defData(defs, varname = "z", formula = 4, variance = 1, dist = "normal")
```

```
# Define additional columns
defsA <- defDataAdd(varname = "a", formula = "w + x + z", variance = 2, dist = "normal")
set.seed(2001)
dt <- genData(10, defs)
dt <- addColumns(defsA, dt)
dt
# Modify definition of additional column
defsA <- updateDefAdd(dtDefs = defsA, changevar = "a", newformula = "w+z", newvariance = 1)
set.seed(2001)
dt <- genData(10, defs)
dt <- addColumns(defsA, dt)
dt
```

viewBasis

Plot basis spline functions

## Description

Plot basis spline functions

## Usage

viewBasis(knots, degree)

## Arguments

knots A vector of values between 0 and 1, specifying cut-points for splines
degree Integer specifying degree of curvature.

## Value

A ggplot object that contains a plot of the basis functions. In total, there will be length(knots) + degree +1 functions plotted.

## Examples

```
knots <- c(0.25, 0. 50, 0.75)
viewBasis(knots, degree = 1)
knots <- c(0.25, 0. 50, 0.75)
viewBasis(knots, degree = 2)
knots <- c(0.25, 0. 50, 0.75)
viewBasis(knots, degree = 3)
```


## Description

Plot spline curves

## Usage

viewSplines(knots, degree, theta)

## Arguments

knots A vector of values between 0 and 1 , specifying cut-points for splines
degree Integer specifying degree of curvature.
theta A vector or matrix of values between 0 and 1. Each column of the matrix represents the weights/coefficients that will be applied to the basis functions determined by the knots and degree. Each column of theta represents a separate spline curve.

## Value

A ggplot object that contains a plot of the spline curves. The number of spline curves in the plot will equal the number of columns in the matrix (or it will equal 1 if theta is a vector).

## Examples

```
knots <- c(0.25, 0.5, 0.75)
theta1 <- c(0.1, 0.8, 0.4, 0.9, 0.2, 1.0)
viewSplines(knots, degree = 2, theta1)
theta2 <- matrix(c(
    0.1, 0.2, 0.4, 0.9, 0.2, 0.3,
    0.1, 0.3, 0.3, 0.8, 1.0, 0.9,
    0.1, 0.4, 0.3, 0.8, 0.7, 0.5,
    0.1, 0.9, 0.8, 0.2, 0.1, 0.6
),
ncol = 4
)
viewSplines(knots, degree = 2, theta2)
```


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