

# Package ‘SIMMS’

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**Type** Package

**Title** Subnetwork Integration for Multi-Modal Signatures

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**Description** Algorithms to create prognostic biomarkers using biological genesets or networks.

**License** GPL-2

**LazyLoad** yes

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## Description

Algorithms to create prognostic biomarkers using biological networks

## Details

Package:	SIMMS
Type:	Package
License:	GPL-2
LazyLoad:	yes

## Author(s)

Syed Haider, Michal Grzadkowski & Paul C. Boutros

## Examples

```
options("warn" = -1);

# get data directory
data.directory <- get.program.defaults(networks.database = "test")[["test.data.dir"]];
```

```
# initialise params
output.directory <- ".";
data.types <- c("mRNA");
feature.selection.datasets <- c("Breastdata1");
training.datasets <- c("Breastdata1");
validation.datasets <- c("Breastdata2");
feature.selection.p.thresholds <- c(0.5);
feature.selection.p.threshold <- 0.5;
learning.algorithms <- c("backward", "forward", "glm");
top.n.features <- 5;

# compute network HRs for all the subnet features
derive.network.features(
  data.directory = data.directory,
  output.directory = output.directory,
  data.types = data.types,
  feature.selection.datasets = feature.selection.datasets,
  feature.selection.p.thresholds = feature.selection.p.thresholds,
  networks.database = "test"
);

# preparing training and validation datasets.
# Normalisation & patientwise subnet feature scores
prepare.training.validation.datasets(
  data.directory = data.directory,
  output.directory = output.directory,
  data.types = data.types,
  p.threshold = feature.selection.p.threshold,
  feature.selection.datasets = feature.selection.datasets,
  datasets = unique(c(training.datasets, validation.datasets)),
  networks.database = "test"
);

# create classifier assessing univariate prognostic power of subnetwork modules (Train and Validate)
create.classifier.univariate(
  data.directory = data.directory,
  output.directory = output.directory,
  feature.selection.datasets = feature.selection.datasets,
  feature.selection.p.threshold = feature.selection.p.threshold,
  training.datasets = training.datasets,
  validation.datasets = validation.datasets,
  top.n.features = top.n.features
);

# create a multivariate classifier (Train and Validate)
create.classifier.multivariate(
  data.directory = data.directory,
  output.directory = output.directory,
  feature.selection.datasets = feature.selection.datasets,
  feature.selection.p.threshold = feature.selection.p.threshold,
  training.datasets = training.datasets,
  validation.datasets = validation.datasets,
```

```

learning.algorithms = learning.algorithms,
top.n.features = top.n.features
);

# (optional) plot Kaplan-Meier survival curves and perform sensitivity analysis
if (FALSE){
  create.survivalplots(
    data.directory = data.directory,
    output.directory = output.directory,
    training.datasets = training.datasets,
    validation.datasets = validation.datasets,
    top.n.features = top.n.features,
    learning.algorithms = learning.algorithms,
    survtime.cutoffs = c(5),
    KM.plotting.fun = "create.KM.plot",
    resolution = 100
  );
}

```

**calculate.meta.survival***Fit a meta-analytic Cox proportional hazards model to a single feature***Description**

Takes a meta-analysis data object and fits a Cox proportional hazards model (possibly with adjustment for some specific covariates) by median-dichotomizing patients within each individual dataset.

**Usage**

```
calculate.meta.survival(feature.name, expression.data, survival.data,
  rounding = 3, other.data = NULL, data.type.ordinal = FALSE,
  centre.data = "median")
```

**Arguments**

- |                                |  |
|--------------------------------|--|
| <code>feature.name</code>      | Character indicate what feature (gene/probe/etc.) should be extracted for analysis                             |
| <code>expression.data</code>   | A list where each component is an expression matrix (patients = columns, genes = rows) for a different dataset |
| <code>survival.data</code>     | A list where each component is an object of class Surv   |
| <code>rounding</code>          | How many digits after the decimal place to include   |
| <code>other.data</code>        | A list of other covariates to be passed to the Cox model (all elements in this list are used)                  |
| <code>data.type.ordinal</code> | Logical indicating whether to treat this datatype as ordinal. Defaults to FALSE                                |

`centre.data` A character string specifying the centre value to be used for scaling data. Valid values are: 'median', 'mean', or a user defined numeric threshold e.g. '0.3' when modelling methylation beta values. This value is used for both scaling as well as for dichotomising data for estimating univariate betas from Cox model. Defaults to 'median'

### Value

Returns a vector containing the HR, p-value, n, and 95% confidence limits of the HR (see `fit.coxmodel()` for details)

### Author(s)

Paul C. Boutros

### Examples

```
data.directory <- get.program.defaults()[[ "test.data.dir" ]];
data.types <- c("mRNA");
x1 <- load.cancer.datasets(
  datasets.to.load = c('Breastdata1'),
  data.types = data.types,
  data.directory = data.directory
);
x2 <- calculate.meta.survival(
  feature.name = "1000_at",
  expression.data = x1$all.data[[data.types[1]]],
  survival.data = x1$all.survobj
);
```

## calculate.network.coefficients

*Calculate Cox statistics for input dataset*

### Description

Function to compute hazard ratios for the genes in pathway-derived networks, by aggregating input datasets into one training cohort. The hazard ratios are computed for each pair by calculating the HR of each gene independently and as an interaction (i.e.  $y = \text{HR}(A) + \text{HR}(B) + \text{HR}(A:B)$ )

### Usage

```
calculate.network.coefficients(data.directory = ".",
  output.directory = ".", training.datasets = NULL,
  data.types = c("mRNA"), data.types.ordinal = c("cna"),
  centre.data = "median", subnets.file.flattened = NULL,
  truncate.survival = 100, subset = NULL)
```

## Arguments

**data.directory** Path to the directory containing datasets as specified by **training.datasets**  
**output.directory** Path to the output folder where intermediate and results files will be saved  
**training.datasets** A vector containing names of training datasets  
**data.types** A vector of molecular datatypes to load. Defaults to c('mRNA')  
**data.types.ordinal** A vector of molecular datatypes to be treated as ordinal. Defaults to c('cna')  
**centre.data** A character string specifying the centre value to be used for scaling data. Valid values are: 'median', 'mean', or a user defined numeric threshold e.g. '0.3' when modelling methylation beta values. This value is used for both scaling as well as for dichotomising data for estimating univariate betas from Cox model. Defaults to 'median'  
**subnets.file.flattened** File containing all the binary interactions derived from pathway-derived networks  
**truncate.survival** A numeric value specifying survival truncation in years. Defaults to 100 years which effectively means no truncation  
**subset** A list with a Field and Entry component specifying a subset of patients to be selected whose annotation Field matches Entry

## Value

Returns a list of matrices for each of the data types. Matrices contain nodes HR/P, edges HR and edges P.

## Author(s)

Syed Haider & Paul C. Boutros

## Examples

```
options("warn" = -1);
program.data <- get.program.defaults(networks.database = "test");
data.directory <- program.data[["test.data.dir"]];
subnets.file.flattened <- program.data[["subnets.file.flattened"]];
coef.nodes.edges <- calculate.network.coefficients(
  data.directory = data.directory,
  output.directory = ".",
  training.datasets = c("Breastdata1"),
  data.types = c("mRNA"),
  subnets.file.flattened = subnets.file.flattened
);
```

---

```
calculate.sensitivity.stats  
    Computes sensitivity measures
```

---

**Description**

Computes sensitivity measures: TP, FP, TN, FN, Sensitivity, Specificity, Accuracy

**Usage**

```
calculate.sensitivity.stats(all.data = NULL)
```

**Arguments**

all.data      A data matrix containing predicted and real risk groups

**Value**

A vector containing TP, FP, TN, FN, Sensitivity, Specificity, Accuracy

**Author(s)**

Syed Haider

---

```
centre.scale.dataset    Centre and scale a data matrix
```

---

**Description**

Centre and scale a data matrix. Scaling is done on each column separately

**Usage**

```
centre.scale.dataset(x = NULL, centre.data = "median")
```

**Arguments**

x      A sample by feature data matrix

centre.data      A character string specifying the centre value to be used for scaling data. Valid values are: 'median', 'mean', or a user defined numeric threshold e.g. '0.3' when modelling methylation beta values. This value is used for both scaling as well as for dichotomising data for estimating univariate betas from Cox model. Defaults to 'median'

**Value**

A centred and scaled data matrix

**Author(s)**

Syed Haider

**Examples**

```
tmp <- matrix(data = rnorm(100, 10, 2), nrow = 20);
tmp.scaled.median <- centre.scale.dataset(x = tmp);
tmp.scaled.mean <- centre.scale.dataset(x = tmp, centre.data = "mean");
tmp.scaled.custom <- centre.scale.dataset(x = tmp, centre.data = 0.3);
```

`create.classifier.multivariate`

*Trains and tests a multivariate survival model*

**Description**

Trains a model on training datasets. Predicts the risk score for all the training & datasets, independently. This function also predicts the risk score for combined training datasets cohort and validation datasets cohort. The risk score estimation is done by multivariate models fit by `fit.survivalmodel`. The function also predicts risk scores for each of the `top.n.features` independently.

**Usage**

```
create.classifier.multivariate(data.directory = ".",
                               output.directory = ".", feature.selection.datasets = NULL,
                               feature.selection.p.threshold = 0.05, training.datasets = NULL,
                               validation.datasets = NULL, top.n.features = 25, models = c("1",
                               "2", "3"), learning.algorithms = c("backward", "forward"),
                               alpha.glm = c(1), k.fold.glm = 10, seed.cv.glm = 51214,
                               cores.glm = 1)
```

**Arguments**

`data.directory` Path to the directory containing datasets as specified by `feature.selection.datasets`, `training.datasets`, `validation.datasets`

`output.directory` Path to the output folder where intermediate and results files will be saved

`feature.selection.datasets` A vector containing names of datasets used for feature selection in function `derive.network.features()`

`feature.selection.p.threshold` One of the P values that were used for feature selection in function `derive.network.features()`. This function does not support vector of P values as used in `derive.network.features()` for performance reasons

<b>training.datasets</b>	A vector containing names of training datasets
<b>validation.datasets</b>	A vector containing names of validation datasets
<b>top.n.features</b>	A numeric value specifying how many top ranked features will be used for univariate survival modelling
<b>models</b>	A character vector specifying which of the models ('1' = N+E, '2' = N, '3' = E) to run
<b>learning.algorithms</b>	A character vector specifying which learning algorithm to be used for model fitting and feature selection. Defaults to c('backward', 'forward'). Available options are: c('backward', 'forward', 'glm')
<b>alpha.glm</b>	A numeric vector specifying elastic-net mixing parameter alpha, with range alpha raning from [0,1]. 1 for LASSO (default) and 0 for ridge. For multiple values of alpha, most optimal value is selected through cross validation on training set
<b>k.fold.glm</b>	A numeric value specifying k-fold cross validation if glm was chosen in learning.algorithms
<b>seed.cv.glm</b>	A numeric value specifying seed for k-fold cross validation if glm was chosen in learning.algorithms
<b>cores.glm</b>	An integer value specifying number of cores to be used for glm if it was chosen in learning.algorithms

**Value**

The output files are stored under `output.directory/output/`

**Author(s)**

Syed Haider & Vincent Stimper

**Examples**

```
# see package's main documentation
```

---

`create.classifier.univariate`

*Trains and tests a univariate (per subnetwork module) survival model*

---

**Description**

Trains a model on training datasets. Predicts the risk score for all the training & datasets, independently. This function also predicts the risk score for combined training datasets cohort and validation datasets cohort. The risk score estimation is done by multivariate models fit by `fit.survivalmodel`. The function also predicts risk scores for each of the `top.n.features` independently.

**Usage**

```
create.classifier.univariate(data.directory = ".",
  output.directory = ".", feature.selection.datasets = NULL,
  feature.selection.p.threshold = 0.05, training.datasets = NULL,
  validation.datasets = NULL, top.n.features = 25, models = c("1",
  "2", "3"))
```

**Arguments**

**data.directory** Path to the directory containing datasets as specified by `feature.selection.datasets`, `training.datasets`, `validation.datasets`

**output.directory** Path to the output folder where intermediate and results files will be saved

**feature.selection.datasets** A vector containing names of datasets used for feature selection in function `derive.network.features()`

**feature.selection.p.threshold** One of the P values that were used for feature selection in function `derive.network.features()`. This function does not support vector of P values as used in `derive.network.features()` for performance reasons

**training.datasets** A vector containing names of training datasets

**validation.datasets** A vector containing names of validation datasets

**top.n.features** A numeric value specifying how many top ranked features will be used for univariate survival modelling

**models** A character vector specifying which of the models ('1' = N+E, '2' = N, '3' = E) to run

**Value**

The output files are stored under `output.directory/output/`

**Author(s)**

Syed Haider

**Examples**

```
# see package's main documentation
```

---

create.KM.plot	<i>Plots Kaplan-meier survival curve for a given risk grouping &amp; survival params</i>
----------------	--

---

**Description**

A generic method to plot KM curves

**Usage**

```
create.KM.plot(riskgroup = NULL, survtime = NULL, survstat = NULL,
               file.name = NULL, main.title = "", resolution = 100)
```

**Arguments**

riskgroup	A vector containing dichotomized risk groups
survtime	A vector containing survival time of the samples
survstat	A vector containing survival status of the samples
file.name	A string containing full qualified path of the output tiff file
main.title	A string specifying main title of the image
resolution	A numeric value specifying resolution of the tiff image of KM survival curves. Defaults to 100

**Value**

The KM survival curves are stored under `output.dir/graphs/`

**Author(s)**

Syed Haider

---

create.sensitivity.plot	<i>Plots sensitivity analysis for class label dichotomization at supplied survtime cutoffs</i>
-------------------------	--

---

**Description**

A method to compute sensitivity, specificity and accuracy at all the survtime cutoff steps provided

**Usage**

```
create.sensitivity.plot(riskscore = NULL, riskgroup = NULL,
                        survtime = NULL, survstat = NULL, survtime.cutoffs = c(seq(5, 10,
                        1)), output.directory = ".", file.stem = NULL, main.title = "",
                        resolution = 100)
```

### Arguments

<code>riskscore</code>	A vector containing predicted risk scores
<code>riskgroup</code>	A vector containing dichotomized risk groups
<code>survtime</code>	A vector containing survival time of the samples
<code>survstat</code>	A vector containing survival status of the samples
<code>survtime.cutoffs</code>	A vector containing cutoff time points used to dichotomize patients into low- and high-risk groups
<code>output.directory</code>	Path to the output folder where intermediate and results files will be saved
<code>file.stem</code>	A string containing base name for image and text files produced by this method
<code>main.title</code>	A string specifying main title of the image
<code>resolution</code>	A numeric value specifying resolution of the tiff image of KM survival curves. Defaults to 100

### Value

The sensitivity analysis plots are stored under `output.directory/graphs/`. The sensitivity analysis results are stored under `output.directory/output/`

### Author(s)

Syed Haider

`create.survivalplots` *Plots Kaplan-meier survival curves*

### Description

Plots Kaplan-meier survival curves for all the training & datasets, independently as well as combined training datasets cohort and validation datasets cohort. The function also plots KM survival curves for each of the `top.n.features` independently.

### Usage

```
create.survivalplots(data.directory = ".", output.directory = ".",
  training.datasets = NULL, validation.datasets = NULL,
  top.n.features = 25, learning.algorithms = c("backward", "forward"),
  truncate.survival = 100, survtime.cutoffs = c(seq(5, 10, 1)),
  main.title = FALSE, KM.plotting.fun = "create.KM.plot",
  plot.univariate.data = FALSE, plot.multivariate.data = TRUE,
  resolution = 100)
```

## Arguments

**data.directory** Path to the directory containing datasets as specified by `training.datasets`,  
**validation.datasets**  
**output.directory** Path to the output folder where intermediate and results files were saved  
**training.datasets** A vector containing names of training datasets  
**validation.datasets** A vector containing names of validation datasets  
**top.n.features** A numeric value specifying how many top ranked features will be used for univariate survival modelling  
**learning.algorithms** A character vector specifying which learning algorithm to be used for model fitting and feature selection. Defaults to `c('backward', 'forward')`. Available options are: `c('backward', 'forward', 'glm')`  
**truncate.survival** A numeric value specifying survival truncation in years. Defaults to 100 years which effectively means no truncation  
**survtime.cutoffs** A vector containing survival cutoff time points to be used for dichotomization of patients into risk groups for sensitivity analysis  
**main.title** A logical to specify plot's main title. Defaults to FASLE  
**KM.plotting.fun** A string containing the name of the method to use for plotting KM curves. Defaults to `create.KM.plot`  
**plot.univariate.data** Logical to indicate whether to plot univariate results for all subnetworks. Default to FALSE  
**plot.multivariate.data** Logical to indicate whether to plot multivariate results for all subnetworks. Defaults to TRUE  
**resolution** A numeric value specifying resolution of the png images of KM survival curves. Defaults to 100

## Value

The KM survival curves are stored under `output.directory/graphs/`

## Author(s)

Syed Haider

## Examples

```
# see package's main documentation
```

create.survobj

*Utility function for loading meta-analysis lists***Description**

Create Surv objects from an annotation-matrix with handling for different time units.

**Usage**

```
create.survobj(annotation = NULL, truncate.survival = 100)
```

**Arguments**

- |                   |  |
|-------------------|--|
| annotation        | A patient annotation matrix (patients = rows) with (at least) columns for surv-time, survstat, and survtime.unit     |
| truncate.survival | A numeric value specifying survival truncation in years. Defaults to 100 years which effectively means no truncation |

**Value**

Returns an object of class Surv

**Author(s)**

Paul C. Boutros

**Examples**

```
annotation.file <- paste(
  get.program.defaults()[[ "test.data.dir" ]],
  "/Breastdata2/patient_annotation.txt", sep = ""
);
annotation <- read.table(
  annotation.file,
  header = TRUE,
  row.names = 1,
  sep = "\t"
);

# select the appropriate survtime and survstat variable for this dataset
annotation$survstat    <- annotation[, 'e.dfs'];
annotation$survtime     <- annotation[, 't.dfs'];
annotation$survtime.unit <- annotation[, 't.dfs.unit'];

# only keep samples with survival data
annotation <- annotation[!is.na(annotation$survstat) & !is.na(annotation$survstat),];
```

---

```
surv.obj <- create.survobj(annotation = annotation);
```

---

**derive.network.features***Derive univariate features from pathway-derived networks***Description**

This function fits Cox model to features as well as interaction between features. The coefficients of features are subsequently used to compute impact score of each of the pathway-derived networks.

**Usage**

```
derive.network.features(data.directory = ".", output.directory = ".",
  data.types = c("mRNA"), data.types.ordinal = c("cna"),
  centre.data = "median",
  feature.selection.fun = "calculate.network.coefficients",
  feature.selection.datasets = NULL,
  feature.selection.p.thresholds = c(0.05), truncate.survival = 100,
  networks.database = "default", subset = NULL, ...)
```

**Arguments**

- data.directory** Path to the directory containing datasets as specified by `feature.selection.datasets`
- output.directory** Path to the output folder where intermediate and results files will be saved
- data.types** A vector of molecular datatypes to load. Defaults to `c('mRNA')`
- data.types.ordinal** A vector of molecular datatypes to be treated as ordinal. Defaults to `c('cna')`
- centre.data** A character string specifying the centre value to be used for scaling data. Valid values are: 'median', 'mean', or a user defined numeric threshold e.g. '0.3' when modelling methylation beta values. This value is used for both scaling as well as for dichotomising data for estimating univariate betas from Cox model. Defaults to 'median'
- feature.selection.fun** Name of the function to be used to estimate network coefficients. Defaults to `'calculate.network.coefficients'`
- feature.selection.datasets** A vector containing names of training datasets to be used to compute cox statistics
- feature.selection.p.thresholds** A vector containing P values to be used as threshold for including features into overall impact score of a network

<code>truncate.survival</code>	A numeric value specifying survival truncation in years. Defaults to 100 years which effectively means no truncation
<code>networks.database</code>	Name of the pathway networks database. Default to NCI PID/Reactome/Biocarta i.e "default"
<code>subset</code>	A list with a Field and Entry component specifying a subset of patients to be selected from each dataset whose annotation Field matches Entry
<code>...</code>	other params to be passed on to user-defined method for estimating coefficients of network features

### Value

The output files are stored under `data.directory/output/`

### Author(s)

Syed Haider

### Examples

```
options("warn" = -1);

# get data directory
data.directory <- get.program.defaults(networks.database = "test")[[["test.data.dir"]]];

# initialise params
output.directory <- ".";
data.types <- c("mRNA");
feature.selection.datasets <- c("Breastdata1");
feature.selection.p.thresholds <- c(0.05);

# estimate network coefficients for all the subnet features
derive.network.features(
  data.directory = data.directory,
  output.directory = output.directory,
  data.types = data.types,
  feature.selection.fun = "calculate.network.coefficients",
  feature.selection.datasets = feature.selection.datasets,
  feature.selection.p.thresholds = feature.selection.p.thresholds,
  networks.database = "test"
);
```

---

dichotomize.dataset     *Dichotomize a single dataset*

---

### Description

Split a dataset into two groups by median-dichotomization

### Usage

```
dichotomize.dataset(x, split.at = "median")
```

### Arguments

- |          |  |
|----------|--|
| x        | A vector of values to be dichotomized  |
| split.at | An character string or a numeric value that is be used to dichotomize. Valid values are: 'median', 'mean', or a user defined numeric threshold. Defaults to 'median' |

### Value

A vector of the data dichotomized onto a 0/1 (low/high) scale.

### Author(s)

Syed Haider & Paul C. Boutros

### Examples

```
tmp <- rnorm(100);
tmp.groups.median <- dichotomize.dataset(tmp);
tmp.groups.mean <- dichotomize.dataset(tmp, split.at = "mean");
tmp.groups.custom <- dichotomize.dataset(tmp, split.at = 0.3);
```

---

---

dichotomize.meta.dataset     *Dichotomize and unlist a meta-analysis list*

---

### Description

Takes a meta-analysis list (and possibly extra data) and dichotomizes based on a specific gene, then returns the unlisted data to the caller.

## Usage

```
dichotomize.meta.dataset(feature.name, expression.data, survival.data,
  other.data = NULL, data.type.ordinal = FALSE,
  centre.data = "median")
```

## Arguments

<code>feature.name</code>	Character indicate what feature (gene/probe/etc.) should be extracted for analysis
<code>expression.data</code>	A list where each component is an expression matrix (patients = columns, genes = rows) for a different dataset
<code>survival.data</code>	A list where each component is an object of class <code>Surv</code>
<code>other.data</code>	A list of other covariates to be unlisted in the final output (all elements in this list are used)
<code>data.type.ordinal</code>	Logical indicating whether to treat this datatype as ordinal. Defaults to FALSE
<code>centre.data</code>	A character string specifying the centre value to be used for scaling data. Valid values are: 'median', 'mean', or a user defined numeric threshold e.g. '0.3' when modelling methylation beta values. This value is used for both scaling as well as for dichotomising data for estimating univariate betas from Cox model. Defaults to 'median'

## Details

NB: `other.data` handling of missing components (i.e. those present in only some datasets) has not been debugged (but may work regardless).

## Value

Returns a list containing components groups (after dichotomization), survtime (in the units of the input data), and survstat. Additional vectors are unlisted from `other.data` if that parameter is not `NULL`.

## Author(s)

Syed Haider & Paul C. Boutros

## Examples

```
data.directory <- get.program.defaults()[["test.data.dir"]];
data.types <- c("mRNA");
x1 <- load.cancer.datasets(
  datasets.to.load = c('Breastdata1'),
  data.types = data.types,
  data.directory = data.directory
);
x2 <- dichotomize.meta.dataset(
```

```
feature.name = "1000_at",
expression.data = x1$all.data[[data.types[1]]],
survival.data = x1$all.survobj
);
```

**fit.coxmodel***Fit a Cox proportional hazards model***Description**

Fit a Cox model (possibly with some linear adjustments) and return key statistics about the fit.

**Usage**

```
fit.coxmodel(groups, survobj, stages = NA, rounding = 3,
other.data = NULL, data.type.ordinal = FALSE)
```

**Arguments**

<code>groups</code>	Grouping of patients (passed directly to coxph, so factors & continuous variables are okay)
<code>survobj</code>	An object of class Surv (from the survival package) – patient ordering needs to be identical as for groups
<code>stages</code>	DEPRECATED! Use <code>other.data</code> instead.
<code>rounding</code>	How many digits of precision should be returned?
<code>other.data</code>	A data-frame (or matrix?) of variables to be controlled in the Cox model. If null, no adjustment is done. No interactions are fit.
<code>data.type.ordinal</code>	Logical indicating whether to treat this datatype as ordinal. Defaults to FALSE

**Value**

A list containing two elements. `cox.stats` containing a vector or matrix: HR, lower 95% CI of HR, upper 95% CI of HR, P-value (for groups), number of samples (total with group assignments, although some may not be included in fit for other reasons so this is an upper-limit). `cox.obj` containing coxph model object

**Author(s)**

Syed Haider & Paul C. Boutros

## Examples

```
survtime <- sample(seq(0.1,10,0.1), 100, replace = TRUE);
survstat <- sample(c(0,1), 100, replace = TRUE);
survobj <- Surv(survtime, survstat);
groups <- sample(c('A','B'), 100, replace = TRUE);
fit.coxmodel(
  groups = as.factor(groups),
  survobj = survobj
);
```

---

**fit.interaction.model** *Cox model two features separately and together*

---

## Description

Using a meta-analysis dataset take two features and Cox model them separately and together and extract HRs and p-values.

## Usage

```
fit.interaction.model(feature1, feature2, expression.data, survival.data,
  data.type.ordinal = FALSE, centre.data = "median")
```

## Arguments

feature1	String indicate what feature (gene/probe/etc.) should be extracted for analysis
feature2	String indicate what feature (gene/probe/etc.) should be extracted for analysis
expression.data	A list where each component is an expression matrix (patients = columns, features = rows) for a different dataset
survival.data	A list where each component is an object of class Surv
data.type.ordinal	Logical indicating whether to treat this datatype as ordinal. Defaults to FALSE
centre.data	A character string specifying the centre value to be used for scaling data. Valid values are: 'median', 'mean', or a user defined numeric threshold e.g. '0.3' when modelling methylation beta values. This value is used for both scaling as well as for dichotomising data for estimating univariate betas from Cox model. Defaults to 'median'

## Details

The interaction model compares cases where feature1 and feature2 concord (both high or both low) to those where they do not. That is, the model is  $y = x_1 + x_2 + (x_1 == x_2)$  and not the typical  $y = x_1 + x_2 + x_1 \cdot x_2$

**Value**

Returns a vector of six elements containing (HR,P) pairs for feature1, feature2, and the interaction

**Author(s)**

Syed Haider & Paul C. Boutros

**Examples**

```
data.dir <- get.program.defaults()["test.data.dir"];
data.types <- c("mRNA");
x1 <- load.cancer.datasets(
  datasets.to.load = c('Breastdata1'),
  data.types = data.types,
  data.directory = data.dir
);
x2 <- fit.interaction.model(
  feature1 = "1000_at",
  feature2 = "2549_at",
  expression.data = x1$all.data[[data.types[1]]],
  survival.data = x1$all.survobj
);
```

**fit.survivalmodel**      *Trains a multivariate survival model*

**Description**

Trains a multivariate survival model and conducts feature selection using both backward elimination and forward selection, independently. TO BE DEPRECATED AND HAS BEEN REPLACED BY `create.classifier.multivariate`

**Usage**

```
fit.survivalmodel(data.directory = ".",
  output.directory = ".",
  feature.selection.datasets = NULL,
  feature.selection.p.threshold = 0.05,
  training.datasets = NULL,
  top.n.features = 25,
  models = c("1", "2", "3"))
```

**Arguments**

<code>data.directory</code>	Path to the directory containing datasets as specified by <code>feature.selection.datasets</code> , <code>training.datasets</code>
<code>output.directory</code>	Path to the output folder where intermediate and results files will be saved

```

feature.selection.datasets
    A vector containing names of datasets used for feature selection in function
    derive.network.features()
feature.selection.p.threshold
    One of the P values that were used for feature selection in function derive.network.features().
    This function does not support vector of P values as used in derive.network.features()
    for performance reasons
training.datasets
    A vector containing names of training datasets to be used to train multivariate
    survival model
top.n.features A numeric value specifying how many top ranked features will be used to train
    the multivariate survival model
models A character vector specifying which models ('1' = N+E, '2' = N, '3' = E) to run

```

**Value**

The output files are stored under `output.directory/output/`

**Author(s)**

Syed Haider

**See Also**

`create.classifier.multivariate`

**Examples**

```
# see package's main documentation
```

**get.adjacency.matrix** *A utility function to convert tab delimited networks file into adjacency matrices*

**Description**

A utility function to convert tab-delimited networks file into adjacency matrices

**Usage**

```
get.adjacency.matrix(subnets.file = NULL)
```

**Arguments**

<b>subnets.file</b>	A tab-delimited file containing networks. New networks start with a new line         with '#' at the begining of network name and subsequent lines contain a binary         interaction per line
---------------------	--

**Value**

A list of adjacency matrices

**Author(s)**

Syed Haider

**Examples**

```
subnets.file <- get.program.defaults()[["subnets.file"]];
all.adjacency.matrices <- get.adjacency.matrix(subnets.file);
```

---

get.chisq.stats      *Applies survdiff function*

---

**Description**

Applies survdiff on different prognoses groups and computes Logrank P using chisquare statistics.

**Usage**

```
get.chisq.stats(groups, survobj)
```

**Arguments**

groups	Grouping of patients (passed directly to survdiff, so factors & continuous variables are okay)
survobj	An object of class Surv (from the survival package) – patient ordering needs to be identical as for groups

**Value**

A vector containing: Chisq, degrees of freedom (DOF) and Logrank P-value.

**Author(s)**

Syed Haider

## Examples

```
survtime <- sample(seq(0.1,10,0.1), 100, replace = TRUE);
survstat <- sample(c(0,1), 100, replace = TRUE);
survobj <- Surv(survtime, survstat);
groups <- sample(c('A','B'), 100, replace = TRUE);
get.chisq.stats(
  groups = as.factor(groups),
  survobj = survobj
);
```

**get.program.defaults**    *A utility function to return the inst/ directory of the installed package and other default settings*

## Description

A utility function to return the inst/ directory of the installed package to get the test datasets and other program related data contents

## Usage

```
get.program.defaults(networks.database = "default")
```

## Arguments

networks.database	Name of the pathway networks database. Default to NCI PID/Reactome/Biocarta i.e "default"
-------------------	--

## Value

Returns a list of paths to the input directories/files where the contents of this package are installed

## Author(s)

Syed Haider

## Examples

```
program.data <- get.program.defaults();
```

---

load.cancer.datasets *Load all cancer meta-analysis datasets*

---

## Description

Returns a list of lists containing all cancer meta-analysis datasets

## Usage

```
load.cancer.datasets(tumour.only = TRUE, with.survival.only = TRUE,  
    truncate.survival = 100, datasets.to.load = "all",  
    data.types = c("mRNA"), datasets.file = "datasets.txt",  
    data.directory = ".", verbose = FALSE, subset = NULL)
```

## Arguments

tumour.only      Logical indicating if we should only load tumour samples (TRUE, the default)  
with.survival.only      Logical indicating if we should only load samples with survival data (TRUE, the default)  
truncate.survival      A numeric value specifying survival truncation in years. Defaults to 100 years which effectively means no truncation  
datasets.to.load      A vector of datasets to be loaded. If 'all', then all available datasets are loaded  
data.types      A vector of molecular datatypes to load. Defaults to c('mRNA')  
datasets.file      A file in data.directory containing a listing of all usable datasets  
data.directory      A directory containing all data-files to be loaded  
verbose      Logical indicating whether or not status messages should be given  
subset      A list with a Field and Entry component specifying a subset of patients to be selected whose annotation Field matches Entry

## Value

Returns a meta-analysis list of lists

## Author(s)

Syed Haider & Paul C. Boutros

## Examples

```
data.dir <- get.program.defaults()["test.data.dir"];
x1 <- load.cancer.datasets(
  datasets.to.load = c('Breastdata1'),
  data.types = c("mRNA"),
  data.directory = data.dir
);
```

### **make.matrix**

*Utility function used by get.adjacency.matrix()*

## Description

Utility function used by `get.adjacency.matrix()`

## Usage

```
make.matrix(vertices, interactions)
```

## Arguments

<code>vertices</code>	Comma separated list of nodes
<code>interactions</code>	Comma separated list of edges

## Value

Returns adjacency matrix

## Author(s)

Syed Haider

## Examples

```
x1 <- make.matrix("a,b,c", "a:b,b:c");
```

---

pred.survivalmodel      *Apply a multivariate survival model to validation datasets*

---

## Description

Predicts the risk score for all the training & datasets, independently. This function also predicts the risk score for combined training datasets cohort and validation datasets cohort. The risk score estimation is done by multivariate models fit by `fit.survivalmodel`. The function also predicts risk scores for each of the `top.n.features` independently. TO BE DEPRECATED AND HAS BEEN REPLACED BY `create.classifier.multivariate`

## Usage

```
pred.survivalmodel(data.directory = ".", output.directory = ".",
  feature.selection.datasets = NULL,
  feature.selection.p.threshold = 0.05, training.datasets = NULL,
  validation.datasets = NULL, top.n.features = 25, models = c("1",
  "2", "3"), write.risk.data = TRUE)
```

## Arguments

`data.directory` Path to the directory containing datasets as specified by `feature.selection.datasets`, `training.datasets`, `validation.datasets`

`output.directory` Path to the output folder where intermediate and results files will be saved

`feature.selection.datasets` A vector containing names of datasets used for feature selection in function `derive.network.features()`

`feature.selection.p.threshold` One of the P values that were used for feature selection in function `derive.network.features()`. This function does not support vector of P values as used in `derive.network.features()` for performance reasons

`training.datasets` A vector containing names of training datasets

`validation.datasets` A vector containing names of validation datasets

`top.n.features` A numeric value specifying how many top ranked features will be used for univariate survival modelling

`models` A character vector specifying which of the models ('1' = N+E, '2' = N, '3' = E) to run

`write.risk.data` A toggle to control whether risk scores and patient risk groups should be written to file

**Value**

The output files are stored under `output.directory/output/`

**Author(s)**

Syed Haider

**See Also**

`create.classifier.multivariate`

**Examples**

```
# see package's main documentation
```

**prepare.training.validation.datasets**  
*Prepare training and validation datasets*

**Description**

Computes per-patient pathway-derived network impact scores across all input datasets, independently

**Usage**

```
prepare.training.validation.datasets(data.directory = ".",
                                    output.directory = ".", data.types = c("mRNA"),
                                    data.types.ordinal = c("cna"), min.ordinal.threshold = c(cna = 3),
                                    centre.data = "median", p.threshold = 0.5,
                                    feature.selection.datasets = NULL, datasets = NULL,
                                    truncate.survival = 100, networks.database = "default",
                                    write.normed.datasets = TRUE, subset = NULL)
```

**Arguments**

`data.directory` Path to the directory containing datasets as specified by `datasets`

`output.directory`

Path to the output folder where intermediate and results files will be saved

`data.types` A vector of molecular datatypes to load. Defaults to `c('mRNA')`

`data.types.ordinal`

A vector of molecular datatypes to be treated as ordinal. Defaults to `c('cna')`

min.ordinal.threshold	A named vector specifying minimum percent threshold for each ordinal data type to be used prior to estimating coefficients. Coefficient for features not satisfying minimum threshold will not be estimated, and set to 0. Defaults to cna threshold as 3 percent
centre.data	A character string specifying the centre value to be used for scaling data. Valid values are: 'median', 'mean', or a user defined numeric threshold e.g. '0.3' when modelling methylation beta values. This value is used for both scaling as well as for dichotomising data for estimating univariate betas from Cox model. Defaults to 'median'
p.threshold	Cox P value threshold to be applied for selecting features (e.g. genes) which will contribute to patient risk score estimation. Defaults to 0.5
feature.selection.datasets	A vector containing names of datasets used for feature selection in function derive.network.features()
datasets	A vector containing names of all the datasets to be later used for training and validation purposes
truncate.survival	A numeric value specifying survival truncation in years. Defaults to 100 years which effectively means no truncation
networks.database	Name of the pathway networks database. Default to NCI PID/Reactome/Biocarta i-e "default"
write.normed.datasets	A toggle to control whether processed mRNA and survival data should be written to file
subset	A list with a Field and Entry component specifying a subset of patients to be selected whose annotation Field matches Entry

## Value

The output files are stored under output.directory/output/

## Author(s)

Syed Haider

## Examples

```
# get data directory
data.directory <- get.program.defaults()[[ "test.data.dir" ]];

# initialise params
output.directory <- ".";
data.types <- c("mRNA");
feature.selection.datasets <- c("Breastdata1");
training.datasets <- c("Breastdata1");
```

```
validation.datasets <- c("Breastdata1", "Breastdata2");

# preparing training and validation datasets.
# Normalisation & patientwise subnet feature scores
prepare.training.validation.datasets(
  data.directory = data.directory,
  output.directory = output.directory,
  data.types = data.types,
  feature.selection.datasets = feature.selection.datasets,
  datasets = unique(c(training.datasets, validation.datasets)),
  networks.database = "test"
);
```

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