# Package 'brainGraph' 

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## Type Package

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Title Graph Theory Analysis of Brain MRI Data
Description A set of tools for performing graph theory analysis of brain MRI data. It works with data from a Freesurfer analysis (cortical thickness, volumes, local gyrification index, surface area), diffusion tensor tractography data (e.g., from FSL) and resting-state fMRI data (e.g., from DPABI). It contains a graphical user interface for graph visualization and data exploration, along with several functions for generating useful figures.

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BugReports https://groups.google.com/forum/?hl=en\#! forum/brainGraph-help
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'plot_global.R' 'plot_group_means.R' 'plot_rich_norm.R' 'plot_vertex_measures.R' 'random_graphs.R' 'rich_club.R' 'robustness.R' 's_core.R' 'set_brainGraph_attributes.R' 'small_world.R' 'spatial_dist.R' 'update_brainGraph_gui.R' 'utils.R' 'utils_matrix.R' 'vertex_roles.R' 'vulnerability.R' 'write_brainnet.R' 'zzz.R'
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apply_thresholds Threshold additional set of matrices

## Description

apply_thresholds thresholds an additional set of matrices (e.g., FA-weighted matrices for DTI tractography) based on the matrices that have been returned from create_mats. This ensures that the same connections are present in both sets of matrices.

## Usage

apply_thresholds(sub.mats, group.mats, W.files, inds)

## Arguments

| sub.mats | List (length equal to number of thresholds) of numeric arrays (3-dim) for all <br> subjects |
| :--- | :--- |
| group.mats | List (equal to number of thresholds) of lists (equal to number of groups) of <br> numeric matrices for group-level data |
| W.files | Character vector of the filenames of the files with connectivity matrices |
| inds | List (length equal to number of groups) of integers; each list element should be <br> a vector of length equal to the group sizes |

## Details

The argument W.files accepts the same formats as A.files; see create_mats for details.

## Value

List containing:
W
A 3-d array of the raw connection matrices
W. norm. sub List of 3-d arrays of the normalized connection matrices for all given thresholds
W. norm.mean List of lists of numeric matrices averaged for each group

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## Examples

```
## Not run:
    W.mats <- apply_thresholds(A.norm.sub, A.norm.mean, f.W, inds)
## End(Not run)
```

    Atlas Helpers Atlas helper functions
    
## Description

guess_atlas tries to determine which atlas is being used based on the data; i.e., the number of vertices/regions.
as_atlas and create_atlas converts/coerces an object to a data.table, or creates one, that is compatible with brainGraph.

## Usage

guess_atlas(x)
as_atlas(object)
create_atlas(regions, coords, lobes, hemis, other = NULL)

## Arguments

| x, object | An object to test or convert to an atlas data.table |
| :--- | :--- |
| regions | Character vector of region names |
| coords | Numeric matrix of spatial coordinates; must have 3 columns |
| lobes | Character or factor vector of lobe membership |
| hemis | Character or factor vector of hemisphere membership. There should probably <br> not be more than 3 unique elements (for left, right, and bi-hemispheric regions) |
| other | A named list of vectors with other data. The names of the list will become <br> column names in the return object. |

## Value

guess_atlas - Character string; either the matched atlas or NA
as_atlas and create_atlas return a data.table that conforms to other atlases in the package, or exits with an error.

## Guessing the atlas from an object

There are several valid inputs to guess_atlas:
data.table The atlas will be guessed based on the number of columns (subtracting by 1 if a "Study ID" column is present). This is the same behavior as for data. frame objects, as well.
igraph The vertex count
brainGraph If there is a atlas graph-level attribute, it will return that. Otherwise, the vertex count.
matrix,array The number of rows, which should equal the number of columns if the input is a connectivity matrix.

Note that this will only work properly for atlases that are currently in the package. If you are using a custom atlas and you receive errors, please open an issue on GitHub.

## Coercing to an atlas

There are several things as_atlas tries to do to make it work without error:

- Coerce the object to data. table
- Add a column of integers named index
- Change columns named ' $x$ ', ' $y$ ', or ' $z$ ' to have .mni at the end
- Convert the lobe and hemi columns to be factors


## Examples

```
my_atlas <- data.frame(name=paste('Region', 1:10), x.mni=rnorm(10),
    y.mni=rnorm(10), z.mni=rnorm(10),
    lobe=rep(c('Frontal', 'Parietal', 'Temporal', 'Occipital', 'Limbic'), 2),
    hemi=c(rep('L', 5), rep('R', 5)))
my_atlas2 <- as_atlas(my_atlas)
str(my_atlas)
str(my_atlas2)
regions <- paste('Region', 1:10)
xyz <- matrix(rnorm(30), nrow=10, ncol=3)
lobe <- rep(c('Frontal', 'Parietal', 'Temporal', 'Occipital', 'Limbic'), 2)
hemi <- c(rep('L', 5), rep('R', 5))
other <- list(network=rep(c('Default mode', 'Task positive'), 5))
my_atlas <- create_atlas(regions, xyz, lobe, hemi, other)
str(my_atlas)
```


## Description

set_brainGraph_attr is a convenience function that sets a number of graph, vertex, and edge attributes for a given graph object. Specifically, it calculates measures that are common in MRI analyses of brain networks.

## Usage

```
set_brainGraph_attr(g, type = c("observed", "random"),
    use.parallel = TRUE, A = NULL, xfm.type = c("1/w", "-log(w)",
    "1-w", "-log10(w/max(w))", "-log10(w/max(w)+1)"),
    clust.method = "louvain")
xfm.weights(g, xfm.type = c("1/w", "-log(w)", "1-w", "-log10(w/max(w))",
    "-log10(w/max(w)+1)"), invert = FALSE)
```


## Arguments

| g | A graph object |
| :--- | :--- |
| type | Character string indicating the type of graphs. Default: observed |
| use.parallel | Logical indicating whether to use foreach. Default: TRUE <br> Numeric matrix; the (weighted) adjacency matrix, which can be used for faster <br> calculation of local efficiency. Default: NULL |
| A | Character string specifying how to transform the weights. Default: $1 / \mathrm{w}$ |
| clust.method | Character string indicating which method to use for community detection. De- <br> fault: 'louvain' |
| invert | Logical indicating whether or not to invert the transformation. Default: FALSE |

## Details

Including type='random' in the function call will reduce the number of attributes calculated. It will only add graph-level attributes for: clustering coefficient, characteristic path length, rich club coefficient, global efficiency, and modularity.

## Value

A graph object with the following attributes:
Graph-level Density, connected component sizes, diameter, \# of triangles, transitivity, average path length, assortativity, global \& local efficiency, modularity, vulnerability, hub score, rich-club coefficient, \# of hubs, edge asymmetry
Vertex-level Degree, strength; betweenness, eigenvector, and leverage centralities; hubs; transitivity (local); k-core, s-core; local \& nodal efficiency; color (community, lobe, component); membership (community, lobe, component); gateway and participation coefficients, within-module degree $z$-score; vulnerability; and coordinates ( $\mathrm{x}, \mathrm{y}$, and z )
Edge-level Color (community, lobe, component), edge betweenness, Euclidean distance (in mm ), weight (if weighted)
xfm.weights returns the same graph object, with transformed edge weights plus a graph attribute (xfm. type) recording the method of transformation

## Negative edge weights

If there are any negative edge weights in the graph, several of the distance-based metrics will not be calculated, because they can throw errors which is undesirable when processing a large dataset. The metrics are: local and nodal efficiency, diameter, characteristic path length, and hubness.

## Transforming edge weights

For distance-based measures, it is important to transform the edge weights so that the strongest connections are re-mapped to having the lowest weights. Then you may calculate e.g., the shortest path length which will include the strongest connections.
xfm.type allows you to choose from 5 options for transforming edge weights when calculating distance-based metrics (e.g., shortest paths). There is no "best-practice" for choosing one over the other, but the reciprocal is probably most common.

1/w reciprocal (default)
$-\log (w)$ the negative (natural) logarithm
1-w subtract weights from 1
$-\log 10(w / \max (w))$ negative (base-10) log of normalized weights
$-\log 10(w / \max (w)+1)$ same as above, but add 1 before taking the $\log$
To transform the weights back to original values, specify invert=TRUE.

## Community detection

clust.method allows you to choose from any of the clustering (community detection) functions available in igraph. These functions begin with cluster_; the function argument should not include this leading character string. There are a few possibilities, depending on the value and the type of input graph:

1. By default, louvain is used, calling cluster_louvain
2. Uses spinglass if there are any negative edges and/or the selected method is spinglass
3. Uses walktrap if there are any negative edge weights and any other method (besides spinglass) is selected
4. Automatically transforms the edge weights if edge_betweenness is selected and the graph is weighted, because the algorithm considers edges as distances

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## See Also

components, diameter, centr_betw, betweenness, centr_eigen, transitivity, distances, assortativity, coreness, communities, knn

```
Bootstrapping Bootstrapping for global graph measures
```


## Description

Perform bootstrapping to obtain groupwise standard error estimates of a global graph measure.
The plot method returns two ggplot objects: one with shaded regions based on the standard error, and the other based on confidence intervals (calculated using the normal approximation).

## Usage

```
brainGraph_boot(densities, resids, R = 1000, measure = c("mod",
```

    "E.global", "Cp", "Lp", "assortativity", "strength", "mod.wt",
    "E.global.wt"), conf = 0.95, .progress = getOption("bg.progress"),
    xfm.type \(=c(" 1 / w ", "-\log (w) ", " 1-w ", "-\log 10(w / \max (w)) "\),
    " \(-\log 10(w / \max (w)+1) "))\)
    \#\# S3 method for class 'brainGraph_boot'
summary (object, ...)
\#\# S3 method for class 'brainGraph_boot'
plot(x, ..., alpha = 0.4)

## Arguments

| densities | Numeric vector of graph densities to loop through |
| :--- | :--- |
| resids | An object of class brainGraph_resids (the output from get.resid) |
| R | Integer; the number of bootstrap replicates. Default: 1e3 |
| measure | Character string of the measure to test. Default: mod |
| conf | Numeric; the level for calculating confidence intervals. Default: 0.95 |
| .progress | Logical indicating whether or not to show a progress bar. Default: getOption('bg.progress' ) |
| xfm.type | Character string specifying how to transform the weights. Default: 1/w |
| object, x | A brainGraph_boot object |
| $\ldots$ | Unused |
| alpha | A numeric indicating the opacity for the confidence bands |

## Details

The confidence intervals are calculated using the normal approximation at the $100 \times$ conf\% level (by default, 95\%).
For getting estimates of weighted global efficiency, a method for transforming edge weights must be provided. The default is to invert them. See xfm. weights.

## Value

brainGraph_boot - an object of class brainGraph_boot containing some input variables, in addition to a list of boot objects (one for each group).
plot - list with the following elements:
se A ggplot object with ribbon representing standard error
ci A ggplot object with ribbon representing confidence intervals

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## See Also

boot, boot.ci
Other Group analysis functions: GLM, Mediation, NBS, brainGraph_permute, mtpc
Other Structural covariance network functions: IndividualContributions, Residuals, brainGraph_permute, corr.matrix, import_scn, plot_volumetric

## Examples

```
## Not run:
boot.E.global <- brainGraph_boot(densities, resids.all, 1e3, 'E.global')
## End(Not run)
```

Brain Atlases Coordinates for data from brain atlases

## Description

Datasets containing spatial coordinates for: the original AAL atlases, the newer AAL2 atlases, Freesurfer atlases, Brainsuite, Craddock200, Dosenbach160, Harvard-Oxford, and LONI probabilistic brain atlas. In addition to coordinates, there are indices for the major lobes and hemispheres of the brain, the class variable (for Destrieux atlases), functional networks (for Dosenbach, Power, and Gordon atlases; plus the Yeo network labels for the Brainnetome atlas).

## Usage

aal116
aal90
aal2.120
aal2.94
destrieux
destrieux.scgm
dk
dk.scgm
dkt
dkt.scgm
brainsuite
craddock200
dosenbach160
hoa112
lpba40
hcp_mmp1.0
power264
brainnetome
gordon333

## Format

A data frame with 90 or 116 (for the original AAL atlases), 94 or 120 (for the newer AAL2 atlases), 148 or 162 (for Destrieux), 68 or 82 (for DK), 62 or 76 (for DKT), 74 (Brainsuite), 200 (Craddock), 160 (Dosenbach), 112 (Harvard-Oxford), 40 (LONI), 246 (Brainnetome), 360 (HCP), 264 (Power), or 333 (Gordon) observations on (some of) the following 19 variables:
name a character vector of region names
x.mni a numeric vector of x-coordinates (in MNI space)
$y$.mni a numeric vector of $y$-coordinates (in MNI space)
z.mni a numeric vector of z-coordinates (in MNI space)
lobe a factor with some of levels Frontal Parietal Temporal Occipital Insula Limbic Cingulate
SCGM Cerebellum (for aal116 and aal2.120) and Brainstem (for craddock200)
hemi a factor with levels $L R$ and $B$ (for dosenbach160)
index a numeric vector
name. full a character vector of full region names, for the DK and DKT atlases
class a factor with levels G G_and_S S, for the Destrieux atlases
network (dosenbach160) a factor with levels default fronto-parietal cingulo-opercular sensorimotor cerebellum occipital
gyrus (brainnetome) Abbreviated names of gyri/regions (including subcortical), with 24 unique values
gyrus.full (brainnetome) Full names of gyrus
subregion (brainnetome) Abbreviated names of subregions (including subdivisions of subcortical gray matter)
subregion. full (brainnetome) Full names of subregion
Yeo_7network (brainnetome) Factor with 8 levels consisting of SCGM plus the 7 networks from Yeo et al.
Yeo_17network (brainnetome) Factor with 18 levels consisting of SCGM plus the 17 networks from Yeo et al.
area (HCP) a factor with 23 cortical areas
Anatomy (power264) Full region/gyrus names for the Power atlas; contains 53 unique regions Brodmann (power264) Integer values for Brodmann areas

## Note

Use of the HCP parcellation is subject to the terms at https://balsa.wustl. edu/WN56. In particular: "I will acknowledge the use of WU-Minn HCP data and data derived from WU-Minn HCP data when publicly presenting any results or algorithms that benefitted from their use."

Region names in the gordon333 atlas were chosen to match those of the hcp_mmp1.0 atlas. Many were determined from the coordinates (using FSL's atlasquery), while the rest were entered manually by me. The lobe values were matched to the HCP atlas, as well.

## Source

https://neuroimaging-core-docs.readthedocs.io/en/latest/pages/atlases.html

## References

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brainGraph Default options for brainGraph

## Description

brainGraph is a package for performing graph theory analysis of brain MRI data.

## Package options

brainGraph uses the following options to configure behavior:

- bg. subject_id: character string specifying the name your project/study uses as a subject identifier. All imported data (e.g., covariates tables) MUST have a column matching this. One possible alternative is 'participant_id', recommended by BIDS. Default: 'Study. ID'
- bg.group: character string specifying the name your project/study uses as a group identifier. All imported data (e.g., covariates tables) MUST have a column matching this. One possible alternative is 'group', recommended by BIDS. Default: 'Group'
- bg.session: character string specifying the name your project/study uses as a "time" or session identifier, in the case of longitudinal studies. All imported data (e.g., covariates tables) MUST have a column matching this. One possible alternative is 'session_id', recommended by BIDS. Default: 'Time'
- bg. progress: logical indicating whether to show progress bars for functions that provide the option. Default: TRUE
- bg. ncpus: integer indicating the number of cores to use for parallel operations. Only used if you have not already registered a parallel backend (see Chapter 5 of the User Guide or https: //github.com/cwatson/brainGraph/blob/master/README.md for examples). Default: 2L
brainGraph-methods brainGraph generic methods


## Description

These functions are S3 generics for various brainGraph-defined objects.
groups returns the "Group" graph attribute for each graph or observation in the object.
region. names is a generic method for extracting region names from various brainGraph objects. These are generally convenience functions.
nregions is a generic method for extracting the number of regions from various brainGraph objects.

```
Usage
    ## S3 method for class 'brainGraphList'
    groups(x)
    ## S3 method for class 'corr_mats'
    groups(x)
    region.names(object)
    ## S3 method for class 'data.table'
    region.names(object)
    nregions(object)
```


## Arguments

$$
x \text {, object An object }
$$

## Details

For a data.table, region. names assumes that it contains a factor column named region.

```
brainGraphList Create a list of brainGraph graphs
```


## Description

make_brainGraphList creates a brainGraphList object, a list containing a set of graphs for all subjects (or group-average graphs) in a study at a specific threshold (or density), in addition to some graph-level attributes common to those graphs.
The [ method will let you subset/slice the graphs for individual subjects and/or groups.
as_brainGraphList coerces a list of graphs to a brainGraphList object. It is assumed that certain metadata attributes - threshold, package version, atlas, imaging modality, edge weighting, and whether they are random graphs - are identical for all graphs in the list.

## Usage

make_brainGraphList(x, atlas, type = c("observed", "random"), level = c("subject", "group", "contrast"), set.attrs = TRUE, modality $=$ NULL, weighting $=$ NULL, threshold $=$ NULL,
gnames $=$ NULL, ...)
\#\# S3 method for class 'array'
make_brainGraphList(x, atlas, type = c("observed",
"random"), level = c("subject", "group", "contrast"), set.attrs = TRUE, modality = NULL, weighting = NULL,

```
    threshold = NULL, gnames = NULL, grpNames = NULL, subnet = NULL,
    mode = "undirected", weighted = NULL, diag = FALSE,
    .progress = getOption("bg.progress"), ...)
## S3 method for class 'corr_mats'
make_brainGraphList(x, atlas = x$atlas,
    type = "observed", level = "group", set.attrs = TRUE,
    modality = NULL, weighting = NULL, threshold = x$densities,
    gnames = names(x$r.thresh), grpNames = gnames, mode = "undirected",
    weighted = NULL, diag = FALSE,
    .progress = getOption("bg.progress"), ...)
## S3 method for class 'brainGraphList'
x[i, g = NULL, drop = TRUE]
## S3 method for class 'brainGraphList'
print(x, ...)
is.brainGraphList(x)
## S3 method for class 'brainGraphList'
nobs(object, ...)
as_brainGraphList(g.list, type = c("observed", "random"),
    level = c("subject", "group", "contrast"))
```


## Arguments

x
atlas Character string specifying the brain atlas
type Character string indicating the type of graphs. Default: observed
level Character string indicating whether the graphs are subject-, group-, or contrastspecific. Default: ' subject'
set.attrs Logical indicating whether to assign all graph-, vertex-, and edge-level attributes (via set_brainGraph_attr). Default: TRUE
modality Character string indicating imaging modality (e.g. 'dti'). Default: NULL
weighting Character string indicating how the edges are weighted (e.g., 'fa', 'pearson', etc.). Default: NULL
threshold Integer or number indicating the threshold used when "sparsifying" the connectivity matrix (if any). Default: NULL
gnames Character vector of graph names (e.g., study IDs if level=' subject ' ). Default: NULL
... Other arguments passed to set_brainGraph_attr
grpNames Character (or factor) vector of group names. If level == 'group ', then you do not need to include this argument (the group names will be the same as gnames). Default: NULL)

| subnet | Integer or character vector indicating the vertices to keep, if you are interested in working with a subset of an atlas. By default, all vertices are used. |
| :---: | :---: |
| mode | Character string defining how the matrix should be interpreted. Default: 'undirected' |
| weighted | Logical specifying whether to create a weighted network |
| diag | Logical indicating whether to include the diagonal of the connectivity matrix. Default: FALSE |
| . progress | Logical indicating whether to print a progress bar. Default: getOption('bg.progress') |
| i | Integer, character, or logical vector for subsetting by subject, or by group (if x\$level='group') |
| g | Integer, character, or logical vector for subsetting by group (if $x \$$ level= 'subject') |
| drop | If TRUE (the default), then return only the list of graphs; otherwise, subset the graphs and return the entire object |
| object | A brainGraphList object |
| g.list | List of graph objects |

## Details

In addition to creating the initial igraph graphs from the connectivity matrices, then attributes will be calculated and assigned for each graph via set_brainGraph_attr if set. attrs=TRUE. Other arguments can be passed to that function. You may display a progress bar by setting . progress=TRUE.
This object can be considered comparable to a 4-D NIfTI file, particularly that returned by FSL's TBSS "prestats" step since that file contains the FA volumes for all study subjects.
To convert an object with 3 "levels" (i.e., subject-level lists from an older brainGraph version), see the code in the Examples below.

## Value

make_brainGraphList returns an object of class brainGraphList with elements:

| threshold | The specified threshold/density |
| :--- | :--- |
| version | The versions of R, igraph, and brainGraph used when creating the graphs |
| atlas | The atlas common to all the graphs |
| modality The imaging modality (if supplied) <br> weighting A string indicating what edge weights represent (if applicable) <br> graphs <br> A named list of brainGraph graphs; the names correspond to the individual <br> graphs' Study IDs  <br> [ - A brainGraphList object (if drop=FALSE) or a list of graphs  |  |

## Subsetting/extracting

The first index is for subsetting the individual graphs. The second index is for subsetting by group membership and requires that the graphs have a Group graph attribute. When both are included, the first index cannot have length or numeric value greater than the number of remaining subjects after subsetting by group.
If the indexing vector(s) is (are) character, the vector(s) must contain one (or more) of the subject or group names. If logical, its length must equal the number of subjects or groups.

## Note

If the input is a corr_mats object, and the extent of the 3-D array is greater than 1 , then only the first will be converted to a graph.

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## See Also

Other Graph creation functions: Creating_Graphs_GLM, Creating_Graphs, make_ego_brainGraph

## Examples

```
## Not run:
# Create a list, one for each threshold
g <- vector('list', length(thresholds))
for (i in seq_along(thresholds)) {
    g[[i]] <- make_brainGraphList(A.norm.sub[[i]], thresholds[i], atlas,
        covars.dti$Study.ID, covars.dti$Group, modality='dti', weighting='fa')
}
## End(Not run)
## Not run:
# Subset the first 10 subjects, irrespective of group
my.bgl[1:10]
# Return object for only 'Control' subjects
my.bgl[, 'Control']
# Return object with graphs from groups 1 and 3
my.bgl[g=c(1, 3), drop=FALSE]
# Subset the first 10 subjects of group 2
my.bgl[1:10, 2]
## End(Not run)
## Not run:
## Convert old version single-subject graph lists
## g[[1]] is group 1, g[[1]][[1]] is threshold 1, g[[1]][[1]][[1]] is subj. 1
kNumThresholds <- length(g[[1]])
g.l <- vector('list', kNumThresholds)
for (i in seq_len(kNumThresholds)) {
    g.l[[i]] <- as_brainGraphList(do.call(Map, c(c, g))[[i]])
}
## End(Not run)
```


## Description

brainGraph_permute draws permutations from linear model residuals to determine the significance of between-group differences of a global or vertex-wise graph measure. It is intended for structural covariance networks (in which there is only one graph per group), but can be extended to other types of data.

## Usage

brainGraph_permute(densities, resids, $N=5000$, perms $=$ NULL, auc = FALSE, level = c("graph", "vertex", "other"), measure = c("btwn.cent", "coreness", "degree", "eccentricity", "clo.cent", "communicability", "ev.cent", "lev.cent", "pagerank", "subg.cent", "E.local", "E.nodal", "knn", "Lp", "transitivity", "vulnerability"), .function = NULL)
\#\# S3 method for class 'brainGraph_permute'
summary (object, measure = object\$measure, alternative = c("two.sided", "less", "greater"), alpha = 0.05, p.sig $=c(" p ", \quad " p . f d r "), \ldots)$
\#\# S3 method for class 'brainGraph_permute'
plot(x, measure = x\$measure,
alternative = c("two.sided", "less", "greater"), alpha = 0.05, p.sig = c("p", "p.fdr"), ptitle = NULL, ...)

## Arguments

| densities | Numeric vector of graph densities |
| :--- | :--- |
| resids | An object of class brainGraph_resids (the output from get.resid) |
| N | Integer; the number of permutations (default: 5e3) |
| perms | Numeric matrix of permutations, if you would like to provide your own (default: <br> NULL) |
| auc | Logical indicating whether or not to calculate differences in the area-under-the- <br> curve of metrics (default: FALSE) |
| level | A character string for the attribute "level" to calculate differences (default: graph) |
| measure | A character string specifying the vertex-level metric to calculate, only used if <br> level=' vertex' (default: btwn. cent). For the summary method, this is to <br> focus on a single graph-level measure (since multiple are calculated at once). |
| .function | A custom function you can pass if level=' other' |
| object, x | A brainGraph_permute object (output by brainGraph_permute). |


| alternative | Character string, whether to do a two- or one-sided test. Default: 'two.sided' |
| :--- | :--- |
| alpha | Numeric; the significance level. Default: 0.05 |
| p.sig | Character string specifying which p-value to use for displaying significant re- <br> sults (default: p) |
| $\ldots$ | Unused |
| ptitle | Character string specifying a title for the plot (default: NULL) |

## Details

If you would like to calculate differences in the area-under-the-curve (AUC) across densities, then specify auc=TRUE.

There are three possible "levels":

1. graph Calculate modularity (Louvain algorithm), clustering coefficient, characteristic path length, degree assortativity, and global efficiency.
2. vertex Choose one of: centrality metrics (betweenness, closeness, communicability, eigenvector, leverage, pagerank, subgraph); k-core; degree; eccentricity; nodal or local efficiency; k-nearest neighbor degree; shortest path length; transitivity; or vulnerability.
3. other Supply your own function. This is useful if you want to calculate something that I haven't hard-coded. It must take as its own arguments: g (a list of lists of igraph graph objects); and densities (numeric vector).

## Value

An object of class brainGraph_permute with input arguments in addition to:
DT A data table with permutation statistics
obs.diff A data table of the observed group differences
Group Group names
The plot method returns a list of ggplot objects

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## See Also

Other Group analysis functions: Bootstrapping, GLM, Mediation, NBS, mtpc
Other Structural covariance network functions: Bootstrapping, IndividualContributions, Residuals, corr.matrix, import_scn, plot_volumetric

## Examples

```
## Not run:
myResids <- get.resid(lhrh, covars)
myPerms <- shuffleSet(n=nrow(myResids$resids.all), nset=1e3)
out <- brainGraph_permute(densities, m, perms=myPerms)
out <- brainGraph_permute(densities, m, perms=myPerms, level='vertex')
out <- brainGraph_permute(densities, m, perms=myPerms,
    level='other', .function=myFun)
## End(Not run)
```

centr_betw_comm Calculate communicability betweenness centrality

## Description

centr_betw_comm calculates the communicability betweenness of the vertices of a graph. The centrality for vertex $r$ is

$$
\omega_{r}=\frac{1}{C} \sum_{p} \sum_{q} \frac{\left(e^{\mathbf{A}}\right)_{p q}-\left(e^{\mathbf{A}+\mathbf{E}(r)}\right)_{p q}}{\left(e^{\mathbf{A}}\right)_{p q}}
$$

where $C=(n-1)^{2}-(n-1)$ is a normalization factor.

## Usage

centr_betw_comm(g, A = NULL)

## Arguments

g
An igraph graph object
A
Numeric matrix; the adjacency matrix of the input graph. Default: NULL

## Value

A numeric vector of the centrality for each vertex

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## References

Estrada, E. and Higham, D.J. and Hatano N. (2009) Communicability betweenness in complex networks. Physica A, 388, 764-774. https://dx.doi.org/10.1016/j.physa.2008.11.011

## See Also

Other Centrality functions: centr_lev

```
    centr_lev Calculate a vertex's leverage centrality
```


## Description

Calculates the leverage centrality of each vertex in a graph.

## Usage

centr_lev (g, $\mathrm{A}=\mathrm{NULL})$

## Arguments

g
An igraph graph object
A
Numeric matrix; the adjacency matrix of the input graph. Default: NULL

## Details

The leverage centrality relates a vertex's degree with the degree of its neighbors. The equation is:

$$
l_{i}=\frac{1}{k_{i}} \sum_{j \in N_{i}} \frac{k_{i}-k_{j}}{k_{i}+k_{j}}
$$

where $k_{i}$ is the degree of the $i^{t h}$ vertex and $N_{i}$ is the set of neighbors of $i$. This function replaces $N a N$ with $N A$ (for functions that have the argument na.rm).

## Value

A vector of the leverage centrality for all vertices.

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## References

Joyce, K.E. and Laurienti P.J. and Burdette J.H. and Hayasaka S. (2010) A new measure of centrality for brain networks. PLoS One, 5(8), e12200. https://dx.doi.org/10.1371/journal.pone. 0012200

## See Also

Other Centrality functions: centr_betw_comm

```
check_sID Test if an object is a character vector of numbers
```


## Description

check_sID is a convenience function to test if a vector (typically the subject ID column in a data.table) is a character vector of numbers, a factor vector of numbers, or a numeric vector. If so, it will zero-pad the variable to have equal width.
pad_zeros pads a vector with zeros to avoid issues with ordering a column of integers or integers converted to character.

## Usage

check_sID(x)
pad_zeros(x)

## Arguments

x
pad_zeros accepts either a vector (numeric or character) or a single integer. check_sID accepts a character, numeric, or factor vector

## Details

This function is meant to avoid issues that arise when sorting a vector of numbers that have been converted to character. For example, import_scn automatically reads in the first column (with FreeSurfer outputs this is the column of subject IDs) as a character variable. If the subject IDs had been all numbers/integers, then sorting (i.e., setting the key in a data.table) would be incorrect: e.g., it might be '1', '10', '2', ...

If " $x$ " is a numeric vector, then the resultant string width will be determined by $\max (x)$ or $x$ itself if the input is a single integer. For example, if $x=10$, it will return ' $01^{\prime}, ~, ~ 02 ', \ldots, 10$ '. If " $x$ " is a character vector, then the output's string width will be $\max (n \operatorname{char}(x))$. For example, if $x$ includes both ' 1 ' and ' 1000 ', it will return '0001', etc.

## Value

check_sID returns either the input vector or a character vector padded with 0
A character vector with zero-padded values

## Examples

```
pad_zeros(10) # '01' '02' ... '10'
x <- c(1, 10, 100)
pad_zeros(x) # '001' '010' '100'
x <- as.character(x)
pad_zeros(x) # '001' '010' '100'
```

coeff_var Calculate coefficient of variation

## Description

coeff_var is a S3 generic that calculates the coefficient of variation, defined as

$$
C V(x)=\frac{\operatorname{sd}(x)}{\operatorname{mean}(x)}
$$

## Usage

```
coeff_var(x, na.rm = FALSE, ...)
## Default S3 method:
coeff_var(x, na.rm = FALSE, ...)
```


## Arguments

| x | Numeric vector, matrix, or array |
| :--- | :--- |
| na.rm | Logical indicating whether NA values should be stripped when calculating sums. <br> Default: FALSE |
| $\ldots$ | Unused |

## Details

If x is a matrix, it will calculate the CV for each column. If x is a 3 D array, it will calculate the coefficient of variation for each row-column combination. If the input dimensions are $n \times n \times r$, a matrix with size $n \times n$ will be returned.

## Value

A numeric vector or matrix

```
communicability Calculate communicability
```


## Description

communicability calculates the communicability of a network, a measure which takes into account all possible paths (including non-shortest paths) between vertex pairs.

## Usage

communicability(g, weights $=$ NULL)

## Arguments

| g | An igraph graph object |
| :--- | :--- |
| weights | Numeric vector of edge weights; if NULL (the default), and if the graph has edge <br> attribute weight, then that will be used. To avoid using weights, this should be |
| NA. |  |

## Details

The communicability $G_{p q}$ is a weighted sum of the number of walks from vertex $p$ to $q$ and is calculated by taking the exponential of the adjacency matrix $A$ :

$$
G_{p q}=\sum_{k=0}^{\infty} \frac{\left(\mathbf{A}^{k}\right)_{p q}}{k!}=\left(e^{\mathbf{A}}\right)_{p q}
$$

where $k$ is walk length.
For weighted graphs with $D=\operatorname{diag}\left(d_{i}\right)$ a diagonal matrix of vertex strength,

$$
G_{p q}=\left(e^{\mathbf{D}^{-1 / 2} \mathbf{A} \mathbf{D}^{-1 / 2}}\right)_{p q}
$$

## Value

A numeric matrix of the communicability

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## References

Estrada, E. and Hatano, N. (2008) Communicability in complex networks. Physical Review E. 77, 036111. https://dx.doi.org/10.1103/PhysRevE.77.036111

Crofts, J.J. and Higham, D.J. (2009) A weighted communicability measure applied to complex brain networks. J. R. Soc. Interface. 6, 411-414. https://dx.doi.org/10.1098/rsif. 2008.0484
contract_brainGraph Contract graph vertices based on brain lobe and hemisphere

## Description

Create a new graph after merging vertices within specified groups. By default, groups are brain lobe and hemisphere membership.

## Usage

contract_brainGraph(g, vgroup = "lobe.hemi")

## Arguments

## g

A brainGraph graph object
vgroup Character string; the name of the vertex attribute to use when contracting the graph. Default: 'lobe.hemi'

## Details

The size vertex-level attribute of the resultant graph is equal to the number of vertices in each group. The $\mathrm{x}-, \mathrm{y}$-, and z -coordinates of the new graph are equal to the mean coordinates of the vertices per group. The new edge weights are equal to the number of inter-group connections of the original graph.

## Value

A new brainGraph graph object with vertex-level attributes representing the mean spatial coordinates, and vertex- and edge-level attributes of color names

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## See Also

contract

```
cor.diff.test
Calculate the p-value for differences in correlation coefficients
```


## Description

Given two sets of correlation coefficients and sample sizes, this function calculates and returns the $z$-scores and $p$-values associated with the difference between correlation coefficients. This function was adapted from https://stackoverflow.com/a/14519007/3357706.

## Usage

cor.diff.test(r1, r2, n, alternative = c("two.sided", "less", "greater"))

## Arguments

| $r 1, r 2$ | Numeric (vector or matrix) of correlation coefficients for both groups |
| :--- | :--- |
| n | Integer vector; number of observations for both groups |
| alternative | Character string, whether to do a two- or one-sided test. Default: 'two.sided' |

## Value

A list with elements $p$ and $z$, the $p$-values and $z$-scores for the difference in correlations.

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## Examples

```
## Not run:
kNumSubjs <- summary(covars$Group)
corr.diffs <- cor.diff.test(corrs$R[, , 1], corrs$R[, , 2], kNumSubjs)
edge.diffs <- t(sapply(which(corr.diffs$p < .05), function(x)
    mapply('[[',
                            dimnames(corr.diffs$p),
                    arrayInd(x, dim(corr.diffs$p)))
                    ))
```

\#\# End(Not run)
corr.matrix Calculate correlation matrix and threshold

## Description

corr.matrix calculates the correlation between all column pairs of a given data frame, and thresholds the resultant correlation matrix based on a given density (e.g., 0.1 if you want to keep only the $10 \%$ strongest correlations). If you want to threshold by a specific correlation coefficient (via the thresholds argument), then the densities argument is ignored.
The plot method will plot "heat maps" of the correlation matrices.

## Usage

```
corr.matrix(resids, densities, thresholds = NULL, what = c("resids",
    "raw"), exclude.reg = NULL, type = c("pearson", "spearman"),
    rand = FALSE)
## S3 method for class 'corr_mats'
x[i, g = NULL]
## S3 method for class 'corr_mats'
plot(x, mat.type = c("thresholded", "raw"),
    thresh.num = 1L, ordered = TRUE, order.by = "lobe",
    graphs = NULL, grp.names = NULL, legend.title = NULL, ...)
## S3 method for class 'corr_mats'
region.names(object)
## S3 method for class 'corr_mats'
nregions(object)
```


## Arguments

| resids | An object of class brainGraph_resids (the output from get.resid) |
| :---: | :---: |
| densities | Numeric vector indicating the resultant network densities; keeps the top $X \%$ of correlations |
| thresholds | Numeric; absolute correlation value to threshold by (default: NULL) |
| what | Character string indicating whether to correlate the residuals or the raw structural MRI values (default: 'resids') |
| exclude.reg | Character vector of regions to exclude (default: NULL) |
| type | Character string indicating which type of correlation coefficient to calculate (default: 'pearson') |
| rand | Logical indicating whether the function is being called for permutation testing; not intended for general use (default: FALSE) |
| x, object | A corr_mats object |
| i | Integer for subsetting by density/threshold |
| g | Integer, character, or logical for subsetting by group |
| mat.type | Character string indicating whether to plot raw or thresholded (binarized) matrices. Default: 'raw' |
| thresh.num | Integer specifying which threshold to plot (if mat.type='thresholded'). Default: 1L |
| ordered | Logical indicating whether to order the vertices by some grouping. Default: TRUE |
| order.by | Character string indicating how to group vertices. Default: ' lobe ' |
| graphs | A brainGraphList object containing graphs with the vertex-level attribute of interest. Default: NULL |
| grp.names | Character vector specifying the names of each group of vertices. Default: NULL |
| legend.title | Character string for the legend title. Default is to leave blank |
|  | Unused |

## Details

If you wish to exclude regions from your analysis, you can give the indices of their columns with the exclude. reg argument.
By default, the Pearson correlation coefficients are calculated, but you can return Spearman by changing the type argument.

## Value

A corr_mats object containing the following components:
R, P Numeric arrays of correlation coefficients and P-values. The length of the 3rd dimension equals the number of groups
$r$.thresh A list of 3-d binary arrays indicating correlations that are above a certain threshold. The length of the list equals the number of groups, and the length of the 3rd dimension equals the number of thresholds/densities.
thresholds Numeric matrix of the thresholds supplied. The number of columns equals the number of groups.
what Residuals or raw values
exclude.reg Excluded regions (if any)
type Pearson or Spearman
atlas The brain atlas used
densities $\quad$ Numeric vector; the densities of the resulting graphs, if you chose to threshold each group to have equal densities.

## Plotting correlation matrices

There are several ways to control the plot appearance. First, you may plot the "raw" correlations, or only those of the thresholded (binarized) matrices. Second, you may order the vertices by a given vertex attribute; by default, they will be ordered by lobe, but you may also choose to order by, e.g., network (for the dosenbach160 atlas) or by community membership. In the latter case, you need to pass a brainGraphList object to the graphs argument; each graph in the object must have a vertex attribute specified in order.by. Finally, you can control the legend text with grp. names.

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## See Also

rcorr
Other Structural covariance network functions: Bootstrapping, IndividualContributions, Residuals, brainGraph_permute, import_scn, plot_volumetric

## Examples

```
## Not run:
myResids <- get.resid(lhrh, covars)
corrs <- corr.matrix(myResids, densities=densities)))
## End(Not run)
## Not run:
corrs <- corr.matrix(myResids, densities)
plot(corrs, order.by='comm', graphs=g.list, grp.names='Community')
## End(Not run)
```


## Description

count_homologous counts the number of edges between homologous regions in a brain graph (e.g. between L and R superior frontal).
count_inter counts the number of edges between and within all vertices in one group (e.g. lobe, hemi, network, etc.).

## Usage

```
count_homologous(g)
count_inter(g, group = c("lobe", "hemi", "network", "class", "gyrus",
    "Yeo_7network", "Yeo_17network", "area", "Brodmann"))
```


## Arguments

```
g A brainGraph graph object
group Character string specifying which grouping to calculate edge counts for. De-
    fault:'lobe'
```


## Value

count_homologous - a named vector of the edge ID's connecting homologous regions count_inter - a data. table of total, intra-, and inter-group edge counts

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## Examples

```
## Not run:
g1.lobecounts <- count_inter(g[[1]][[N]], 'lobe')
## End(Not run)
```


## Description

create_mats creates arrays from connection matrices (e.g.,fdt_network_matrix from FSL or ROICorrelation.txt from DPABI). You may choose to normalize these matrices by the waytotal or region size (tractography), or not at all.

## Usage

```
create_mats(A.files, modality = c("dti", "fmri"), divisor = c("none",
    "waytotal", "size", "rowSums"), div.files = NULL,
    threshold.by = c("consensus", "density", "mean", "consistency", "raw"),
    mat.thresh = 0, sub.thresh = 0.5, inds = list(seq_along(A.files)),
    algo = c("probabilistic", "deterministic"), P = 5000, ...)
```


## Arguments

| A.files | Character vector of the filenames with connection matrices |
| :---: | :---: |
| modality | Character string indicating data modality (default: dti) |
| divisor | Character string indicating how to normalize the connection matrices; either 'none' (default), 'waytotal', 'size', or 'rowSums' (ignored if modality equals fmri) |
| div.files | Character vector of the filenames with the data to normalize by (e.g. a list of waytotal files) (default: NULL) |
| threshold.by | Character string indicating how to threshold the data; choose density, mean, or consistency if you want all resulting matrices to have the same densities (default: consensus) |
| mat.thresh | Numeric (vector) for thresholding connection matrices (default: 0) |
| sub.thresh | Numeric (between 0 and 1) for thresholding by subject numbers (default: 0.5) |
| inds | List (length equal to number of groups) of integers; each list element should be a vector of length equal to the group sizes |
| algo | Character string of the tractography algorithm used (default: 'probabilistic'). Ignored if modality is fmri. |
| P | Integer; number of samples per seed voxel (default: 5000) |
|  | Arguments passed to symmetrize |

## Value

A list containing:
A
A 3-d array of the raw connection matrices
A. norm
A 3-d array of the normalized connection matrices
A.bin A list of 3-d arrays of binarized connection matrices, one array for each threshold
A.bin.sums A list of 3-d arrays of connection matrices, with each entry signifying the number of subjects with a connection present; the number of list elements equals the length of mat. thresh, and the extent of the arrays equals the number of groups
A.inds A list of arrays of binarized connection matrices, containing 1 if that entry is to be included
A. norm. sub List of 3-d arrays of the normalized connection matrices for all given thresholds
A. norm.mean List of 3-d arrays of connection matrices averaged for each group

## Connection matrix files

The A.files argument is mandatory and may be specified in a few ways:

1. A character vector of the filenames (preferably with full path).
2. A single character string specifying the directory in which all connectivity matrices are located. This will load all files in the directory.
3. A named list in which the names match the arguments to list.files. This will load all files in path that match the pattern argument, if present, and will load all files in child directories if recursive=TRUE. See examples below.

The same options apply to div.files as well.

## Thresholding methods

The argument threshold. by has 4 options:

1. consensus Threshold based on the raw (normalized, if selected) values in the matrices. If this is selected, it uses the sub. thresh value to perform "consensus" thresholding.
2. density Threshold the matrices to yield a specific graph density (given by the mat. thresh argument).
3. mean Keep only connections for which the cross-subject mean is at least 2 standard deviations higher than the threshold (specified by mat. thresh)
4. consistency Threshold based on the coefficient of variation to yield a graph with a specific density (given by mat. thresh). The edge weights will still represent those of the input matrices. See Roberts et al. (2017) for more on "consistency-based" thresholding.
5. raw Threshold each subject's matrix individually, irrespective of group membership. Ignores sub.thresh.

The argument mat. thresh allows you to choose a numeric threshold, below which the connections will be replaced with 0 ; this argument will also accept a numeric vector. The argument sub. thresh will keep only those connections for which at least $X \%$ of subjects have a positive entry (the default is 0.5 , or $50 \%$ ).

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## References

Roberts, JA and Perry, A and Roberts, G and Mitchell, PB and Breakspear, M (2017) Consistencybased thresholding of the human connectome. NeuroImage. 145, 118-129. https://dx.doi.org/ 10.1016/j.neuroimage.2016.09.053

## Examples

```
## Not run:
thresholds <- seq(from=0.001, to=0.01, by=0.001)
fmri.mats <- create_mats(f.A, modality='fmri', threshold.by='consensus',
    mat.thresh=thresholds, sub.thresh=0.5, inds=inds)
dti.mats <- create_mats(f.A, divisor='waytotal', div.files=f.way,
    mat.thresh=thresholds, sub.thresh=0.5, inds=inds)
# Specify a directory and filename pattern
conn_files <- list(path='~/data', pattern='.*fdt_network_matrix')
dti.mats <- create_mats(conn_files, ...)
## End(Not run)
```

Creating_Graphs Create a brainGraph object

## Description

make_brainGraph is the main creation function for creating a brainGraph graph object. This is simply an igraph graph object with additional attributes (at all levels). Several of the graph-level attributes serve the purpose of providing metadata on how the connectivity matrices/networks were created.
make_brainGraph.bg_mediate creates a graph only for vertex-level analyses.
make_empty_brainGraph creates an empty undirected brainGraph object with vertex count equal to the atlas specified; i.e., it creates a graph with 0 edges. Typically used to present results from an analysis in which edges don't make sense (e.g., GLM comparing differences in a vertex-level attribute).

## Usage

make_brainGraph(x, atlas, type = c("observed", "random"),
level = c("subject", "group", "contrast"), set.attrs = TRUE,
modality $=$ NULL, weighting $=$ NULL, threshold $=$ NULL, ...)
\#\# S3 method for class 'igraph'
make_brainGraph(x, atlas, type = c("observed",
"random"), level = c("subject", "group", "contrast"),
set.attrs = TRUE, modality = NULL, weighting = NULL,
threshold $=$ NULL, name $=$ NULL, Group $=$ NULL, subnet $=$ NULL,.. )

```
## S3 method for class 'matrix'
make_brainGraph(x, atlas, type = c("observed",
    "random"), level = c("subject", "group", "contrast"),
    set.attrs = TRUE, modality = NULL, weighting = NULL,
    threshold = NULL, name = NULL, Group = NULL, subnet = NULL,
    mode = "undirected", weighted = NULL, diag = FALSE, ...)
## S3 method for class 'bg_mediate'
make_brainGraph(x, atlas = x$atlas,
    type = "observed", level = "contrast", set.attrs = FALSE,
    modality = NULL, weighting = NULL, threshold = NULL, ...)
is.brainGraph(x)
## S3 method for class 'brainGraph'
summary(object, print.attrs = c("all", "graph",
    "vertex", "edge", "none"), ...)
make_empty_brainGraph(atlas, type = c("observed", "random"),
    level = c("subject", "group", "contrast"), modality = NULL,
    weighting = NULL, threshold = NULL, name = NULL, Group = NULL,
    ...)
```


## Arguments

| x |  |
| :--- | :--- |
| atlas |  |
| type | An igraph graph object, numeric matrix, or bg_mediate object |
| level | Character string specifying the brain atlas <br> Character string indicating the type of graphs. Default: observed <br> Character string indicating whether the graphs are subject-, group-, or contrast- <br> specific. Default: 'subject' |
| set.attrs | Logical indicating whether to assign all graph-, vertex-, and edge-level attributes <br> (via set_brainGraph_attr). Default: TRUE |
| modality | Character string indicating imaging modality (e.g. 'dti'). Default: NULL <br> Character string indicating how the edges are weighted (e.g., 'fa', 'pearson', <br> etc.). Default: NULL |
| weighting | Integer or number indicating the threshold used when "sparsifying" the connec- <br> tivity matrix (if any). Default: NULL |
| Arguments passed to set_brainGraph_attr |  |
| name | Character string indicating subject ID or group/contrast name, depending on the |
| Group | level. Default: NULL <br> Character string indicating group membership. Default: NULL |
| subnet | Integer or character vector indicating the vertices to keep, if you are interested <br> in working with a subset of an atlas. By default, all vertices are used. |
| mode | Character string defining how the matrix should be interpreted. Default: 'undirected ' |
| weighted | Logical specifying whether to create a weighted network |


| diag | Logical indicating whether to include the diagonal of the connectivity matrix. <br> Default: FALSE |
| :--- | :--- |
| object | A brainGraph object |
| print.attrs | Character string indicating whether or not to list the object's attributes (default: <br> all) |

## Value

A brainGraph graph object with additional graph-, vertex-, and edge-level attributes (see below).
The method for bg_mediate returns a brainGraph_mediate object, which has extra attributes:
Graph mediator, treat, outcome, nobs
Vertex b?.acme, p?.acme, b?.ade, p?.ade, b?.prop, p?.prop, b.tot, p.tot
make_empty_brainGraph - An empty brainGraph graph object

## Graph-level attributes

Graph-level attributes added are:
version The $R$, brainGraph, and igraph package versions used to create the graph
date The creation date, from as.POSIXct
atlas Character string denoting the brain atlas used
type Character string specifying whether this is an observed or random graph
modality The imaging modality; you can choose anything you like, but the summary.brainGraph knows about dti, fmri, thickness, area, and volume
weighting What edge weights represent; you can choose anything you like, but summary. brainGraph knows about fa, sld (streamline density, tractography), pearson, spearman, kendall, and partial (partial correlation coefficient)
threshold Numeric indicating the threshold used to create the final connectivity matrix (if any)
name Character string specifying the study ID or group/contrast name, depending on the level argument
Group Character string specifying the experimental group that the given subject belongs to, or if it is a group-level graph
subnet Integer vector, if subnet was specified in the call

## Vertex attributes

Vertex-level attributes added are:
name The names of the brain regions in the network
lobe The names of the major brain lobes for each vertex
hemi The names of the hemisphere for each vertex (either 'L', 'R', or 'B')
lobe.hemi The lobe-hemisphere combination (represented as an integer vector)
class The tissue class (if applicable)
network The network (if the atlas is dosenbach160)
$\mathbf{x}, \mathbf{y}, \mathbf{z}$ The spatial coordinates of the (centers-of-mass) brain regions in MNI space
x.mni,y.mni,z.mni Same as above
color.lobe,color.class,color.network Colors for vertices of their respective membership
circle.layout Integer vector indicating the order (going counter-clockwise from the top) for circular layouts

## Edge attributes

Edge-level attributes added are:
color.lobe,color.class,color.network Correspond to the vertex attribute of the same name. Intergroup edges will be colored gray

## Specifying a subnetwork

You can create a graph for a subset of an atlas's regions with the subnet argument. This can either be a numeric or character vector. If the input object (either a matrix or an igraph graph) has fewer rows/columns or vertices, respectively, than the atlas then the subnet graph attribute will also be added to the return object. This may occur if, for example, you use make_auc_brainGraph on graphs that were initially created from subnetworks.

## See Also

Other Graph creation functions: Creating_Graphs_GLM, brainGraphList, make_ego_brainGraph

## Examples

```
## Not run:
bg <- make_brainGraph(A, 'dkt', modality='dti', weighting='fa',
    mode='undirected', diag=FALSE, weighted=TRUE)
## End(Not run)
```


## Description

These methods create a brainGraphList with attributes specific to the results of brainGraph_GLM, mtpc , or NBS. The graphs element of the returned object will contain one graph for each contrast.

## Usage

```
## S3 method for class 'bg_GLM'
make_brainGraphList(x, atlas = x$atlas,
    type = "observed", level = "contrast", set.attrs = FALSE,
    modality = NULL, weighting = NULL, threshold = NULL,
    gnames = x$con.name, ...)
    ## S3 method for class 'mtpc'
    make_brainGraphList(x, atlas = x$atlas,
        type = "observed", level = "contrast", set.attrs = FALSE,
        modality = NULL, weighting = NULL, threshold = NULL,
        gnames = x$con.name, ...)
    ## S3 method for class 'NBS'
make_brainGraphList(x, atlas, type = "observed",
    level = "contrast", set.attrs = TRUE, modality = NULL,
    weighting = NULL, threshold = NULL, gnames = x$con.name,
    mode = "undirected", weighted = TRUE, diag = FALSE, ...)
```


## Arguments

x
atlas Character string specifying the brain atlas to use
type Character string indicating the type of graphs. Default: observed
level Character string indicating whether the graphs are subject-, group-, or contrastspecific. Default: ' subject'
set.attrs Logical indicating whether to assign all graph-, vertex-, and edge-level attributes (via set_brainGraph_attr). Default: TRUE
modality Character string indicating imaging modality (e.g. 'dti'). Default: NULL
weighting Character string indicating how the edges are weighted (e.g., 'fa', 'pearson', etc.). Default: NULL
threshold Integer or number indicating the threshold used when "sparsifying" the connectivity matrix (if any). Default: NULL
gnames Character vector of graph names (e.g., study IDs if level=' subject '). Default: NULL
... Other arguments passed to set_brainGraph_attr
mode Character string defining how the matrix should be interpreted. Default: 'undirected'
weighted Logical specifying whether to create a weighted network
diag Logical indicating whether to include the diagonal of the connectivity matrix. Default: FALSE

## Value

A brainGraphList object, with a graph object for each contrast with additional attributes:

Graph name (contrast name), outcome (the outcome variable), alpha (the significance level); for MTPC: tau.mtpc, S.mtpc, S.crit, A.crit

Vertex size2 (t-statistic); size (the t-stat transformed for visualization purposes); p (equal to $1-p$ ); $p . f d r$ (equal to $1-p_{F D R}$, the FDR-adjusted p-value); effect.size (the contrast of parameter estimates for t-contrasts; the extra sum of squares for F-contrasts); se (the standard error of gamma); A.mtpc, sig (binary indicating whether A.mtpc > A.crit) (for MTPC)
make_brainGraphList. NBS returns graphs with additional attributes:
Vertex comp (integer vector indicating connected component membership), p.nbs ( P value for each component)

Edge $\quad$ stat (the test statistic for each connection), $p$ (the P -value)

## Note

Only valid for vertex-level and NBS analyses.

## See Also

```
brainGraph_GLM,mtpc,NBS
```

Other Graph creation functions: Creating_Graphs, brainGraphList, make_ego_brainGraph

```
edge_asymmetry Calculate an asymmetry index based on edge counts
```


## Description

Calculate an asymmetry index, a ratio of intra-hemispheric edges in the left to right hemisphere of a graph for brain MRI data.

## Usage

edge_asymmetry(g, level = c("hemi", "vertex"), A = NULL)

## Arguments

g An igraph graph object
level Character string indicating whether to calculate asymmetry for each region, or the hemisphere as a whole (default: 'hemi')

A
Numeric matrix; the adjacency matrix of the input graph. Default: NULL

## Details

The equation is:

$$
A=\frac{E_{l h}-E_{r h}}{0.5 \times\left(E_{l h}+E_{r h}\right)}
$$

where $l h$ and $r h$ are left and right hemispheres, respectively. The range of this measure is $[-2,2]$ (although the limits will only be reached if all edges are in one hemisphere), with negative numbers indicating more edges in the right hemisphere, and a value of 0 indicating equal number of edges in each hemisphere.
The level argument specifies whether to calculate asymmetry for each vertex, or for the whole hemisphere.

## Value

A data table with edge counts for both hemispheres and the asymmetry index; if level is vertex, the data table will have vcount $(\mathrm{g})$ rows.

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

```
efficiency Calculate graph global, local, or nodal efficiency
```


## Description

This function calculates the global efficiency of a graph or the local or nodal efficiency of each vertex of a graph.

## Usage

efficiency(g, type = c("local", "nodal", "global"), weights = NULL, xfm = FALSE, xfm.type = NULL, use.parallel = TRUE, $A=$ NULL, D = NULL)

## Arguments

| g | An igraph graph object |
| :--- | :--- |
| type | Character string; either local, nodal, or global. Default: local |
| weights | Numeric vector of edge weights; if NULL (the default), and if the graph has edge <br> attribute weight, then that will be used. To avoid using weights, this should be |
|  | NA. |
| xfm | Logical indicating whether to transform the edge weights. Default: FALSE |
| xfm.type | Character string specifying how to transform the weights. Default: 1/w |
| use.parallel | Logical indicating whether or not to use foreach. Default: TRUE |
| A | Numeric matrix; the adjacency matrix of the input graph. Default: NULL |
| D | Numeric matrix; the graph's "distance matrix" |

## Details

Local efficiency for vertex $i$ is:

$$
E_{l o c a l}(i)=\frac{1}{N} \sum_{i \in G} E_{\text {global }}\left(G_{i}\right)
$$

where $G_{i}$ is the subgraph of neighbors of $i$, and $N$ is the number of vertices in that subgraph.
Nodal efficiency for vertex $i$ is:

$$
E_{\text {nodal }}(i)=\frac{1}{N-1} \sum_{j \in G} \frac{1}{d_{i j}}
$$

Global efficiency for graph $G$ with $N$ vertices is:

$$
E_{\text {global }}(G)=\frac{1}{N(N-1)} \sum_{i \neq j \in G} \frac{1}{d_{i j}}
$$

where $d_{i j}$ is the shortest path length between vertices $i$ and $j$. Alternatively, global efficiency is equal to the mean of all nodal efficiencies.

## Value

A numeric vector of the efficiencies for each vertex of the graph (if type is local|nodal) or a single number (if type is global).

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## References

Latora, V. and Marchiori, M. (2001) Efficient behavior of small-world networks. Phys Rev Lett, 87.19, 198701. https://dx.doi.org/10.1103/PhysRevLett.87. 198701

Latora, V. and Marchiori, M. (2003) Economic small-world behavior in weighted networks. Eur Phys J B, 32, 249-263. https://dx.doi.org/10.1140/epjb/e2003-00095-5

## Description

brainGraph_GLM specifies and fits a General Linear Model (GLM) at each vertex for a given vertex measure (e.g. degree) or at the graph-level (e.g., global efficiency). Given a contrast matrix or list of contrast(s), and contrast type (for t - or F-contrast(s), respectively) it will calculate the associated statistic(s) for the given contrast(s).

The summary method prints the results, only for which $p<\alpha$, where alpha comes from the bg_GLM object. "Simple" P-values are used by default, but you may change this to the FDR-adjusted or permutation P -values via the function argument p . sig. You may also choose to subset by contrast.
The plot method plots the GLM diagnostics (similar to that of plot.lm). There are a total of 6 possible plots, specified by the which argument; the behavior is the same as in plot. 1m. Please see the help for that function.
The [ method allows you to select observations (i.e., rows of $X$ and $y$ ) and independent variables (i.e., columns of X) from a bg_GLM object.

## Usage

```
brainGraph_GLM(g.list, covars, measure, contrasts, con.type = c("t",
    "f"), outcome = NULL, X = NULL, con.name = NULL,
    alternative = c("two.sided", "less", "greater"), alpha = 0.05,
    level = c("vertex", "graph"), permute = FALSE,
    perm.method = c("freedmanLane", "terBraak", "smith", "draperStoneman",
    "manly", "stillWhite"), part.method = c("beckmann", "guttman",
    "ridgway"), N = 5000, perms = NULL, long = FALSE, ...)
## S3 method for class 'bg_GLM'
print(x, ...)
## S3 method for class 'bg_GLM'
summary(object, p.sig = c("p", "p.fdr", "p.perm"),
    contrast = NULL, alpha = object$alpha, digits = max(3L,
    getOption("digits") - 2L), print.head = TRUE, ...)
## S3 method for class 'bg_GLM'
plot(x, region = NULL, which = c(1L:3L, 5L),
    ids = TRUE, ...)
## S3 method for class 'bg_GLM'
x[i, j]
```


## Arguments

| g.list <br> covars <br> measure | A brainGraphList object <br> A data. table of covariates |
| :--- | :--- |
| contrasts | Character string of the graph measure of interest <br> Numeric matrix (for T statistics) or list of matrices (for F statistics) specifying <br> the contrast(s) of interest; if only one contrast is desired, you can supply a vector <br> (for T statistics) |
| con.type | Character string; either ' $t$ ' or ' $f$ ' (for t or F-statistics). Default: ' $t$ ' <br> outcome |
| Character string specifying the name of the outcome variable, if it differs from <br> the graph metric (measure) |  |
| $X$ | Numeric matrix, if you wish to supply your own design matrix. Ignored if <br> outcome $!=$ measure. |


| con. name | Character vector of the contrast name(s); if contrasts has row/list names, those will be used for reporting results |
| :---: | :---: |
| alternative | Character string, whether to do a two- or one-sided test. Default: 'two.sided' |
| alpha | Numeric; the significance level. Default: 0.05 |
| level | Character string; either vertex (default) or graph |
| permute | Logical indicating whether or not to permute group labels. Default: FALSE |
| perm.method | Character string indicating the permutation method. Default: 'freedmanLane' |
| part.method | Character string; the method of partitioning the design matrix into covariates of interest and nuisance. Default: 'beckmann' |
| N | Integer; number of permutations to create. Default: 5 e 3 |
| perms | Matrix of permutations, if you would like to provide your own. Default: NULL |
| long | Logical indicating whether or not to return all permutation results. Default: FALSE |
|  | Arguments passed to brainGraph_GLM_design |
| object, x | A bg_GLM object |
| p.sig | Character string specifying which P -value to use for displaying significant results (default: p) |
| contrast | Integer specifying the contrast to plot/summarize; defaults to showing results for all contrasts |
| digits | Integer specifying the number of digits to display for P-values |
| print.head | Logical indicating whether or not to print only the first and last 5 rows of the statistics tables (default: TRUE) |
| region | Character string specifying which region's results to plot; only relevant if level='vertex ' . Default: NULL |
| which | Integer vector indicating which of the 6 plots to print to the plot device. Default: $c(1: 3,5)$ |
| ids | Logical indicating whether to plot subject ID's for outliers. Otherwise plots the integer index |
| i | Integer/character vector; the observation number(s) or row names to select or remove |
| j | Integer/character vector; the design matrix column number(s) or names to select or remove |

## Details

The measure argument will be the graph- or vertex-level measure of interest. Often, this will serve as the model's outcome (or dependent, or response) variable; i.e., the variable typically denoted by $y$ in GLMs. In other cases, you may wish to choose some other variable as the outcome; e.g., IQ, age, etc. Then you could test for a direct association between the network measure and outcome of interest, or test for another association while adjusting for the network metric. For these applications, you must provide the variable name via the outcome argument. This is analogous to -evperdat in FSL's PALM and to --pvr in FreeSurfer.

## Value

An object of class bg_GLM containing some input-specific variables (level, outcome, measure, con.type, contrasts, con.name, alt, alpha, permute, perm.method, part.method, $N$ ) in addition to:

DT. Xy A data table from which the design matrices are created and the outcome variable, for all regions.
X A named numeric matrix or a 3D array of the design matrix. Rownames are subject IDs, column names are predictor variables, and dimnames along the 3rd dimension are region names (if applicable). This is a 3D array only if outcome != measure and level == 'vertex'.
y
A named numeric matrix of the outcome variable. Rownames are Study IDs and column names are regions. There will be multiple columns only if outcome == measure and level == 'vertex'.
DT A data table with an entry for each vertex (region) containing statistics of interest
removed. subs A named integer vector in which the names are subject ID's of those removed due to incomplete data (if any). The integers correspond to the row number in the input covars table.
runX If outcome != measure and level == 'vertex', this will be a character vector of the regions for which the design matrix is invertible. Otherwise, it is NULL.
runY Character vector of the regions for which the outcome variable has 0 variability. For example, if level='vertex' and measure='degree', some regions may be disconnected or have the same degree for all subjects.
atlas $\quad$ Character string of the atlas used (guessed based on the vertex count).
perm A list containing: null.dist (the null distribution of maximum statistics), thresh (the statistic value corresponding to the $100 \times(1-\alpha)$ th \% percentile of the null distribution)

The plot method returns a list of ggplot objects (if installed) or writes the plots to a PDF in the current directory named bg_GLM_diagnostics.pdf

A bg_GLM object with the specified row(s) selected or removed from both $X$ and $y$, and column(s) selected/removed from $X$

## Design matrix

The GLM's design matrix will often be identical to the model matrix associated with 1 m objects (if "dummy" coding, the default, is used) and is created from the input data. table and arguments passed to brainGraph_GLM_design. The first column should have the name of getOption('bg. subject_id') and its values must match the name graph-level attribute of the input graphs. The covariates table must be supplied even if you provide your own design matrix $X$. If level='vertex' and outcome == measure, there will be a single design for all regions but a separate model for each region (since the graph measure varies by region). If level='vertex' and outcome != measure, there will be a separate design (and, therefore, a separate model) for each region even though the outcome is the same in all models.

## Contrasts and statistics

Either t- or F-contrasts can be calculated (specified by con.type). Multiple t-contrasts can be specified by passing a multi-row matrix to contrasts. Multiple F-contrasts can be specified by passing a list of matrices; all matrices must have the same number of columns. All F-contrasts are necessarily two-sided; t-contrasts can be any direction, but only one can be chosen per function call. If you choose con. type=" $f$ ", the calculated effect size is represented by the ESS ("extra sum of squares"), the additional variance explained for by the model parameters of interest (as determined by the contrast matrix). The standard error for F-contrasts is the sum of squared errors of the full model.

## Non-parametric permutation tests

You can calculate permutations of the data to build a null distribution of the maximum statistic which corrects for multiple testing. To account for complex designs, the design matrix must be partitioned into covariates of interest and nuisance; the default method is the Beckmann method. The default permutation strategy is that of Freedman \& Lane (1983), and is the same as that in FSL's randomise. See randomise.

## Note

The [ method is used when calculating studentized residuals and other "leave-one-out" diagnostics, and typically should not be called directly by the user.

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## See Also

plot.lm
Other GLM functions: GLM design, GLM fits, mtpc
Other Group analysis functions: Bootstrapping, Mediation, NBS, brainGraph_permute, mtpc

## Examples

```
## Not run:
conmat <- matrix(c(0, 0, 0, 1), nrow=1)
rownames(conmat) <- 'Control > Patient'
res.lm <- brainGraph_GLM(g[[6]], covars=covars.all[tract == 1],
    measure='strength', contrasts=conmat, alt='greater', permute=TRUE, long=TRUE)
## End(Not run)
## Not run:
## Save objects and then to multipage PDF
lmPlots <- plot(x)
ggsave('lmPlots.pdf', lmPlots)
## Save all the GLM sub-objects from MTPC analysis
res.mtpc <- mtpc(...)
```

```
glmPlots <- lapply(res.mtpc$res.glm, plot, which=1:6)
ml <- marrangeGrob(glmPlots, nrow=1, ncol=1)
ggsave('glmPlots.pdf', ml, width=8.5, height=11)
## End(Not run)
```

GLM basic info Extract basic information from a bg_GLM object

## Description

These functions return the terms, term labels, model formula, "case names", "variable names", region names, and number of observations for a bg_GLM object. The term labels are used for ANOVA tables.

## Usage

```
## S3 method for class 'bg_GLM'
nobs(object, ...)
## S3 method for class 'bg_GLM'
terms(x, ...)
## S3 method for class 'bg_GLM'
formula(x, ...)
## S3 method for class 'bg_GLM'
labels(object, ...)
## S3 method for class 'bg_GLM'
case.names(object, ...)
## S3 method for class 'bg_GLM'
variable.names(object, ...)
## S3 method for class 'bg_GLM'
region.names(object)
## S3 method for class 'bg_GLM'
nregions(object)
```


## Arguments

... Unused
$x$, object A bg_GLM object

## Value

terms returns a named integer list in which the names are the term labels and the list elements are the column(s) of the design matrix for each term. nobs returns an integer. The other functions return character vectors.

## Note

formula returns a character string, not a formula object.

```
GLM design Create a design matrix for linear model analysis
```


## Description

brainGraph_GLM_design takes a data.table of covariates and returns a design matrix to be used in linear model analysis.

## Usage

brainGraph_GLM_design(covars, coding = c("dummy", "effects", "cell.means"), factorize = TRUE, binarize = NULL, int = NULL, mean.center = FALSE, center.how = c("all", "within-groups"), center.by = getOption("bg.group"))

## Arguments

covars A data. table of covariates
coding Character string indicating how factor variables will be coded. Default: ' dummy '
factorize Logical indicating whether to convert character columns into factor. Default: TRUE
binarize Character vector specifying the column name(s) of the covariate(s) to be converted from type factor to numeric. Default: NULL
int Character vector specifying the column name(s) of the covariate(s) to test for an interaction. Default: NULL
mean. center Logical indicating whether to mean center non-factor variables. Default: FALSE
center how Character string indicating whether to use the grand mean or groupwise means. Default: 'all'
center.by Character string indicating which grouping variable to use for calculating means (if applicable). Default: 'Group'

## Details

There are three different ways to code factors: dummy, effects, or cell-means (chosen by the argument coding). Effects coding is sometimes referred to as deviation coding. Dummy coding is the default when calling lm. To understand the difference between these, see Chapter 8 of the User Guide.

## Value

A numeric matrix. Rownames are subject ID's and column names are the variable names. There will be additional attributes recording the coding, factorize, and mean. center function arguments. There will also be attributes for binarize and int if they are not NULL, and center. how and center.by if mean. center=TRUE.

## Character variables

The default behavior is to convert all character columns (excluding the Study ID column and any that you list in the binarize argument) to factor variables. To change this, set factorize=FALSE. So, if your covariates include multiple character columns, but you want to convert Scanner to binary instead of a factor, you may still specify binarize='Scanner' and get the expected result. binarize will convert the given factor variable(s) into numeric variable(s), which is performed before centering (if applicable).

## Centering

The argument mean. center will mean-center (i.e., subtract the mean of from each variable) any non-factor variables (including any dummy/indicator covariates). This is done after "factorizing" and "binarizing". If center. how= 'all', then the "grand mean" will be used; otherwise, the groupwise means will be used. The grouping variable is determined by center.by and is by default 'Group'.

## Interactions

int specifies which variables should interact with one another. This argument accepts both numeric/continuous (e.g., Age) and factor variables (e.g., Sex). All interaction combinations will be generated: if you supply 3 variables, all two-way and the single three-way interaction will be generated. This variable must have at least two elements; it is otherwise ignored. It is generally recommended that centering be performed when including interaction terms.

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## See Also

Other GLM functions: GLM fits, GLM, mtpc

## Examples

```
## Not run:
# Recreate design matrix when "outcome == measure"
DT <- res.glm$DT.Xy[region == levels(region)[1L],
    !c('region', res.glm$outcome),
    with=FALSE]
X <- do.call(brainGraph_GLM_design, c(list(covars=DT),
                                    attributes(res.glm$X)[-c(1L, 2L)]))
all.equal(X, res.glm$X)
```

```
## End(Not run)
```

GLM fits Fit design matrices to one or multiple outcomes

## Description

These are the "base" model-fitting functions that solve the least squares problem to estimate model coefficients, residuals, etc. for brain network data.
fastLmBG_t and fastLmBG_f calculate contrast-based statistics for T or F contrasts, respectively. It accepts any number of contrasts (i.e., a multi-row contrast matrix).

## Usage

fastLmBG(X, $Y, Q R=q r . d e f a u l t(X), Q=q r \_Q 2(Q R, n=n, p=p)$, $R=q r_{2} R 2(Q R, p), n=\operatorname{dim}(X)[1 L], p=Q R \$ r a n k, n y=\operatorname{dim}(Y)[2 L]$, $\mathrm{dfR}=\mathrm{n}-\mathrm{p}, \mathrm{XtXinv}=\operatorname{inv}(\mathrm{QR}))$
fastLmBG_3d(X, Y, runX, $\mathrm{QR}=\operatorname{qr}(X[$, , runX, drop $=F A L S E])$, $\mathrm{Q}=$ lapply (QR, qr_Q2, $\mathrm{n}=\mathrm{n}, \mathrm{p}=\mathrm{p}), \mathrm{R}=\operatorname{lapply}\left(\mathrm{QR}, \mathrm{qr} \_\mathrm{R} 2, \mathrm{p}\right)$, $\mathrm{n}=\operatorname{dim}(\mathrm{X})[1 \mathrm{~L}], \mathrm{p}=\mathrm{QR}[[1 \mathrm{~L}]] \$$ rank, ny $=$ length $(\mathrm{runX}), \mathrm{dfR}=\mathrm{n}-$ $\mathrm{p}, \mathrm{XtXinv}=\operatorname{inv}(\mathrm{QR}))$
fastLmBG_3dY(X, Y, runX, QR = qr(X[, , runX, drop = FALSE]), $\mathrm{Q}=$ lapply (QR, qr_Q2, $\mathrm{n}=\mathrm{n}, \mathrm{p}=\mathrm{p}), \mathrm{R}=$ lapply(QR, qr_R2, p), $\mathrm{n}=\operatorname{dim}(\mathrm{X})[1 \mathrm{~L}], \mathrm{p}=\mathrm{QR}[[1 \mathrm{~L}]] \$ r a n k$, ny $=$ length (runX), $\mathrm{dfR}=\mathrm{n}-$ $\mathrm{p}, \mathrm{XtXinv}=\operatorname{inv}(\mathrm{QR})$ )
fastLmBG_3dY_1p(X, Y, runX, $Q R=\operatorname{rr}(X[, \quad, \quad r u n X, d r o p=F A L S E])$, $\mathrm{Q}=\operatorname{lapply}\left(\mathrm{QR}, \operatorname{qr} \_\mathrm{Q} 2, \operatorname{diag}(1 \mathrm{~L}, \mathrm{n}, 1 \mathrm{~L}), \mathrm{n}, 1 \mathrm{~L}\right), \mathrm{R}=\operatorname{lapply}(\mathrm{QR}$, function( $r$ ) $r$ \$qr[1L]), $n=\operatorname{dim}(X)[1 L], p=1 L$, ny = length(runX), $\mathrm{dfR}=\mathrm{n}-1 \mathrm{~L}, \mathrm{XtXinv}=\operatorname{inv}(\mathrm{QR})$ )
fastLmBG_t(fits, contrasts, alternative = c("two.sided", "less",
"greater"), alpha = NULL)
fastLmBG_f(fits, contrasts, rkC = NULL, nC = length(contrasts))

## Arguments

$X \quad$ Design matrix or 3D array of design matrices
Y Numeric matrix; there should be 1 column for each outcome variable (so that in a graph-level analysis, this is a column matrix)
$Q R, Q, R \quad$ The $Q R$ decomposition(s) and $Q$ and $R$ matrix(es) of the design matrix(es). If $X$ is a 3D array, these should be lists

| $\mathrm{n}, \mathrm{p}, \mathrm{ny}, \mathrm{dfR}$ | Integers; the number of observations, model rank, number of regions/outcome <br> variables, and residual degrees of freedom |
| :--- | :--- |
| XtXinv | Numeric matrix or array; the inverse of the cross-product of the design ma- <br> trix(es) |
| runX | Character vector of the regions for which the design matrix is not singular |
| fits | List object output by one of the model fitting functions (e.g., fastLmBG) |
| contrasts | Numeric matrix (for T statistics) or list of matrices (for F statistics) specifying <br> the contrast(s) of interest; if only one contrast is desired, you can supply a vector <br> (for T statistics) |
| alternative | Character string, whether to do a two- or one-sided test. Default: 'two. sided' |
| alpha | Numeric; the significance level. Default: 0.05 |
| rkC, nc | Integers; the rank of the contrast matrix and number of contrasts, respectively <br> (for F contrasts) |

## Value

A list with elements

| coefficients | Parameter estimates |
| :--- | :--- |
| rank | Model rank |
| df.residual | Residual degrees of freedom |
| residuals | Model residuals |
| sigma | The residual standard deviation, or root mean square error (RMSE) |
| fitted.values | Model fitted values |
| qr | The design matrix QR decomposition(s) |
| cov.unscaled | The "unscaled covariance matrix" |

fastLmBG_t - A multidimensional array with the third dimension equaling the number of contrasts; each matrix contains the contrast of parameter estimates, standard error of the contrast, T-statistics, P -values, FDR -adjusted P -values, and confidence intervals (if alpha is given)
fastLmBG_f - A numeric matrix with columns for the effect size, standard error, F statistic, Pvalues, and FDR-adjusted P -values

## Parameter estimation

These functions use the $Q R$ decomposition to calculate the least squares solution which is the same as the base 1 m function. If we substitute $X=Q R$ in the standard normal equations, the equation to be solved reduces to

$$
X^{T} X \hat{\beta}=X^{T} y \Rightarrow R \hat{\beta}=Q^{T} y
$$

Since R is an upper-triangular matrix, we can use the backsolve function which is a bit faster than solve. In some cases, the fastLmBG* functions are about as fast or faster (particularly when X is not permuted) as one in which the normal equations are solved directly; additionally, using the $Q R$ method affords greater numerical stability.

## Different scenarios

There are a few different scenarios for fitting models of the data, with a separate function for each:
fastLmBG The main function for when there is a single design matrix $X$ and any number of outcome variables $Y$.
fastLmBG_3d Fits models when there is a different design matrix $X$ for each region and a single outcome variable $Y$, which in this case will be a column matrix.
fastLmBG_3dY Fits models when there is both a different design matrix $X$ and outcome variable $Y$ for each region. Occurs under permutation for the Freedman-Lane, ter Braak, and StillWhite methods.
fastLmBG_3dY_1p Fits models when there is both a different design and outcome variable for each region, and also when $X$ is a rank-1 matrix (i.e., it has 1 column). Only occurs under permutation with the Still-White method if there is a single regressor of interest.

In the last case above, model coefficients are calculated by simple (i.e., non-matrix) algebra.

## Improving speed/efficiency

Speed/efficiency gains will be vast for analyses in which there is a single design matrix $X$ for all regions, there are multiple outcome variables (i.e., vertex-level analysis), and the permutation method chosen does not permute $X$. Specifically, these are Freedman-Lane, ter Braak, and Manly methods. Therefore, the QR decomposition, the $Q$ and $R$ matrices, and the "unscaled covariance matrix" (which is $\left(X^{T} X\right)^{-1}$ ) only need to be calculated once for the entire analysis. Other functions (e.g., lm.fit) would recalculate these for each permutation.
Furthermore, this (and the other model fitting functions in the package) will likely only work in models with full rank. I sacrifice proper error checking in favor of speed, but hopefully any issues with the model will be identified prior to the permutation step. Finally, the number of observations, model rank, number of outcome variables, and degrees of freedom will not change and therefore do not need to be recalculated (although these probably amount to a negligible speed boost).
In case there are multiple design matrices, or the permutation method permutes the design, then the QR decomposition will need to be calculated each time anyway. For these cases, I use more simplified functions qr_Q2 and qr_R2 to calculate the $Q$ and $R$ matrices, and then the fitted values, residuals, and residual standard deviation are calculated at the same time (whereas lm.fit and others would calculate these each time).

## Contrast-based statistics

The contrast of parameter estimates, $\gamma$, for T contrasts is

$$
\gamma=C \hat{\beta}
$$

where $C$ is the contrast matrix with size $k \times p$ (where $k$ is the number of contrasts) and $\hat{\beta}$ is the matrix of parameter estimates with size $p \times r$ (where $r$ is the number of regions). For F contrasts, the effect size is the extra sum of squares and is calculated as

$$
\gamma\left(C\left(X^{T} X\right)^{-1} C^{T}\right)^{-1} \gamma^{T}
$$

The standard error of a T contrast is

$$
\sqrt{\hat{\sigma}\left(X^{T} X\right)^{-1}}
$$

where $\hat{\sigma}$ is the residual standard deviation of the model and the second term is the unscaled covariance matrix. The standard error for F contrasts is simply the residual sum of squares. P -values and FDR-adjusted P-values (across regions) are also calculated. Finally, if $\alpha$ is provided for T contrasts, confidence limits are calculated.

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## See Also

randomise
Other GLM functions: GLM design, GLM, mtpc

```
GLM influence measures
```

Influence measures for a bg_GLM object

## Description

These functions compute common (leave-one-out) diagnostics for the models in a bg_GLM object.

## Usage

```
## S3 method for class 'bg_GLM'
rstandard(model, type = c("sd.1", "predictive"), ...)
## S3 method for class 'bg_GLM'
rstudent(model, ...)
## S3 method for class 'bg_GLM'
hatvalues(model, ...)
## S3 method for class 'bg_GLM'
cooks.distance(model, ...)
dffits.bg_GLM(model)
## S3 method for class 'bg_GLM'
dfbeta(model, ...)
## S3 method for class 'bg_GLM'
dfbetas(model, ...)
    covratio.bg_GLM(model)
    ## S3 method for class 'bg_GLM'
    influence(model, do.coef = TRUE, region = NULL, ...)
```


## Arguments

| model | A bg_GLM object |
| :--- | :--- |
| type | The type of standardized residuals. Default: 'sd. $1^{\prime}$ |
| $\ldots$ | Unused |
| do.coef | Logical indicating whether to calculate dfbeta |
| region | Character string of the region(s) to return results for. Default is to calculate for <br> all regions |

## Details

The influence method calculates all diagnostics present in lm.influence and influence.measures, consisting of the following functions:
rstandard Standardized residuals. Choosing type='predictive' returns leave-one-out cross validation residuals. The "PRESS" statistic can be calculated as colSums(resids.p^2)
rstudent Studentized residuals
hatvalues The leverage, or the diagonal of the hat/projection matrix
cooks.distance Cook's distance
dffits.bg_GLM The change in fitted values when deleting observations
dfbeta The change in parameter estimates (coefficients) when deleting observations
dfbetas The scaled change in parameter estimates
covratio.bg_GLM The covariance ratios, or the change in the determinant of the covariance matrix of parameter estimates when deleting observations

## Value

Most influence functions return a numeric matrix in which rownames are Study ID's and column names are regions. dfbeta and dfbetas return a numeric array in which each column is a parameter estimate and the 3rd dimension is for each region. influence returns a list with class infl.bg_GLM and elements:
infmat Numeric array (like dfbeta) with DFBETAs, DFFITs, covratios, Cook's distance, and hat values
is.inf Logical array of the same data as infmat; values of TRUE indicate the subject-variable-region combination is an outlier value
f The model formula
sigma The leave-one-out residual standard deviation
wt.res Model residuals

## Outlier values

Each variable has a different criterion for determining outliers. In the following: $x$ is the influence variable (for DFBETA, the criterion applies to all DFBETAs); $k$ is the number of columns of the design matrix; dfR is the residual degrees of freedom; and n is the number of observations.

DFBETAs If $|x|>1$
DFFITs If $|x|>3 \sqrt{k / d f R}$
covratio If $|1-x|>(3 k / d f R)$
cook If $F_{k, d f R}(x)>0.5$
hat If $x>3 k / n$
The return object of influence has a print method which will list the subjects/variables/regions for which an outlier was detected.

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## See Also

GLM

## Description

These functions compute the log-likelihood and Akaike's An Information Criterion (AIC) of a bg_GLM object. See logLik. lm and extractAIC for details.

## Usage

\#\# S3 method for class 'bg_GLM'
$\operatorname{logLik}($ object, REML $=$ FALSE,... )
\#\# S3 method for class 'bg_GLM'
extractAIC(fit, scale $=0, k=2, \ldots$ )

## Arguments

object, fit
REML
... Unused
scale
Should be left at its default
k
A bg_GLM object

Logical indicating whether to return the restricted log-likelihood. Default: FALSE

Numeric; the weight of the equivalent degrees of freedom

## Details

The functions AIC and BIC will also work for bg_GLM objects because they each call logLik.

## Value

logLik returns an object of class logLik with several attributes. extractAIC returns a numeric vector in which the first element is the equivalent degrees of freedom and the remaining are the AIC's for each region

```
GLM statistics Extract model fit statistics from a bg_GLM object
```


## Description

These functions extract or calculate model fit statistics of a bg_GLM object. These can be found in the output from summary. lm.

## Usage

```
## S3 method for class 'bg_GLM'
coef(object, ...)
## S3 method for class 'bg_GLM'
confint(object, parm, level = 0.95, ...)
## S3 method for class 'bg_GLM'
fitted(object, ...)
## S3 method for class 'bg_GLM'
residuals(object, type = c("response", "partial"), ...)
## S3 method for class 'bg_GLM'
deviance(object, ...)
    coeff_determ(object, adjusted = FALSE)
    ## S3 method for class 'bg_GLM'
    df.residual(object, ...)
    ## S3 method for class 'bg_GLM'
    sigma(object, ...)
    ## S3 method for class 'bg_GLM'
    vcov(object, ...)
    coeff_table(object, CI = FALSE, level = 0.95)
    ## S3 method for class 'bg_GLM'
    anova(object, region = NULL, ...)
```


## Arguments

object
...
parm
level
type
adjusted
CI
region

A bg_GLM object
Unused
Vector of parameters to calculate confidence intervals for. Default is to use all parameters
The confidence level. Default: 0.95
Character string specifying the type of residuals to return. Default: 'response ' Logical indicating whether to calculate the adjusted R-squared. Default: FALSE
Logical indicating whether to include confidence intervals of parameter estimates in the coefficient summary table. Default: FALSE
Character vector indicating the region(s) to calculate ANOVA statistics for. Default: NULL (use all regions)

## Details

These mimic the same functions that operate on lm objects, and include:
coef Regression coefficients (parameter estimates)
confint Confidence intervals (by default, 95\%) for parameter estimates
fitted Fitted (mean) values; i.e., the design matrix multiplied by the parameter estimates, $X \hat{\beta}$
residuals Model residuals; i.e., the response/outcome variable minus the fitted values. Partial residuals can also be calculated
deviance Model deviance, or the residual sum of squares
coeff_determ Calculate the coefficient of determination (or $R^{2}$ ), adjusted or unadjusted
df.residual Residual degrees of freedom
sigma Residual standard deviation, sometimes called the root mean squared error ( $R M S E$ )
vcov Variance-covariance matrix of the model parameters
coeff_table returns model coefficients, standard errors, T-statistics, and P-values for all model terms and regions in a bg_GLM object. This is the same as running summary ( $x$ )\$coefficients for a lm object.

## Value

A named numeric vector, matrix, or array, depending on the function:
coef Matrix in which rownames are parameter names and column names are regions fitted, residuals

Matrix in which rownames are Study ID's and column names are regions. If type='partial', an array is returned in which columns are terms and the 3rd dimension are regions
deviance, coeff_determ, sigma
Numeric vector with elements for each region
df.residual Single integer; the degrees of freedom
confint, vcov, coeff_table
Numeric array; the extent of the third dimension equals the number of regions
anova returns a list of tables of class anova

## ANOVA tables

The anova method calculates the so-called Type III test statistics for a bg_GLM object. These standard ANOVA statistics include: sum of squares, mean squares, degrees of freedom, F statistics, and P-values. Additional statistics calculated are: $\eta^{2}$, partial $\eta^{2}, \omega^{2}$, and partial $\omega^{2}$ as measures of effect size.

## Note

sigma - The denominator is not the number of observations, but rather the model's residual degrees of freedom.
When calculating partial residuals, the parameter estimates are not re-calculated after removing one of the model terms.

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## See Also

GLM, Anova

Graph Data Tables Create a data table with graph global and vertex measures

## Description

graph_attr_dt is a helper function that takes a brainGraphList or a list of graphs and creates a data. table of global measures for each graph. Each row will be for a different graph.
vertex_attr_dt is a helper function that creates a data.table in which each row is a vertex and each column is a different network measure (degree, centrality, etc.).

## Usage

graph_attr_dt(bg.list)
vertex_attr_dt(bg.list)

## Arguments

bg.list A brainGraphList object, or a list of graph objects

## Value

A data.table

```
See Also
    graph_attr,graph_attr_names
    vertex_attr,vertex_attr_names,graph_from_data_frame
```

    Graph Distances Calculate Euclidean distance of edges and vertices
    
## Description

edge_spatial_dist calculates the Euclidean distance of an igraph graph object's edges. The distances are in mm and based on MNI space. These distances are NOT along the cortical surface, so can only be considered approximations, particularly concerning inter-hemispheric connections. The input graph must have atlas as a graph-level attribute.
vertex_spatial_dist calculates, for each vertex of a graph, the average Euclidean distance across all of that vertex's connections.

## Usage

edge_spatial_dist(g)
vertex_spatial_dist(g)

## Arguments

g An igraph graph object

## Value

edge_spatial_dist - a numeric vector with length equal to the edge count of the input graph, consisting of the Euclidean distance (in mm ) of each edge
vertex_spatial_dist - a named numeric vector with length equal to the number of vertices, consisting of the average distance (in mm ) for each vertex

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## References

Alexander-Bloch, A.F. and Vertes, P.E. and Stidd, R. et al. (2013) The anatomical distance of functional connections predicts brain network topology in health and schizophrenia. Cerebral Cortex, 23, 127-138. https://dx.doi.org/10.1093/cercor/bhr388

## Description

hubness calculates the "hubness" (see reference) of the vertices in a graph. These are vertices which meet at least two of the following four criteria:

1. Have high degree/strength
2. Have high betweenness centrality
3. Have low clustering coefficient
4. Have low average path length

For each criterion, "high" or "low" means "in the top 20\%" across all vertices. Vertices meeting any of the criteria get a value of 1 for that metric; these are summed to yield the hubness score which ranges from 0-4. As in the reference article, vertices with a score of 2 or higher are to be considered hubs, although that determination isn't made in this function.

## Usage

hubness(g, xfm.type = g\$xfm.type, weights = NULL, prop.keep = 0.2)

## Arguments

g An igraph graph object
xfm.type Character string specifying how to transform the weights. Default: 1/w
weights Numeric vector of edge weights; if NULL (the default), and if the graph has edge attribute weight, then that will be used. To avoid using weights, this should be NA.
prop.keep Numeric (between 0 and 1) indicating the proportion of vertices to consider as having a high score. Default: 0.2 (20\%)

## Value

A numeric vector with the vertices' hubness score

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## References

van den Heuvel, M.P. and Mandl, R.C.W. and Stam, C.J. and Kahn, R.S. and Pol, H.E.H. (2010) Aberrant frontal and temporal complex network structure in schizophrenia: a graph theoretical analysis. The Journal of Neuroscience, 30(47), 15915-15926. https://dx.doi.org/10.1523/ JNEUROSCI.2874-10. 2010
import_scn Import data for structural connectivity analysis

## Description

Given a directory, atlas name, and imaging modality/structural metric, this function imports data for structural connectivity analysis. It expects files containing a table of region-wise structural MRI measures (e.g., mean cortical thickness), with one file for each hemisphere. The first column of all files should contain the subject $I D$; the column name will be changed to the value of getOption('bg.subject_id').

## Usage

import_scn(datadir, atlas, modality = "thickness", exclude.subs = NULL, custom.atlas $=$ NULL)

## Arguments

datadir The path name of the directory containing the data files
atlas Character string specifying the atlas in use. For a custom atlas, please specify 'custom', and provide the name to the custom. atlas argument
modality The structural imaging measure (default: 'thickness')
exclude.subs Vector indicating the subjects to exclude, if any (default: NULL)
custom.atlas Character string specifying the name of the R object for the atlas in use, if atlas=' custom' was also supplied (default: NULL)

## Details

The files should have specific names; the second in the following list is only required for atlases/parcellations that include subcortical gray matter (e.g., dk. scgm).

- \$\{parcellation\}_\$\{hemi\}_\$\{modality\}.csv for cortical volume, thickness, surface area, or local gyrification index (LGI). Here, \$\{parcellation\} can be aparc, aparc.DKTatlas40, or aparc.a2009s. For example, for cortical thickness with the Desikan-Killiany atlas, the filename should be aparc_lh_thickness.csv. If you are using a custom atlas, see the Note below. The $\$\{$ hemi $\}$ variable is either lh or rh. Finally, $\$\{$ modality\} should be either volume, thickness, area, or lgi.
- asegstats.csv for SCGM volume


## Value

A list containing:
atlas Character string
modality Character string
lhrh A data.table of structural MRI measures for both hemispheres

```
aseg A data.table of structural MRI measures for subcortical gray matter, if appli-
    cable
subs.excluded Vector of subject ID's that were excluded
subs.missing Vector of subject ID's that are not present in both the cortical and subcortical
    tables (if applicable)
```


## Note

When using a custom atlas, the name of the atlas's data.table should match the \$\{parcellation\} portion of the filename (specification shown above). Furthermore, it must conform to the output of Freesurfer's aparcstats2table (and asegstats2table, if applicable). Otherwise, please contact me for inclusion of a different data type.
The subject ID column will be zero-padded (to the left) to avoid issues when the variable is numeric; this ensures that all ID's will have the same number of characters and sorting will be done properly.

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## See Also

Other Structural covariance network functions: Bootstrapping, IndividualContributions, Residuals, brainGraph_permute, corr.matrix, plot_volumetric

## Examples

```
## Not run:
    raw_data <- import_scn('/home/cwatson/data', atlas='dkt',
                exclude.subs=c('con07', 'con23', 'pat15'))
## End(Not run)
```

IndividualContributions
Approaches to estimate individual network contribution

## Description

loo calculates the individual contribution to group network data for each subject in each group using a "leave-one-out" approach. The residuals of a single subject are excluded, and a correlation matrix is created. This is compared to the original correlation matrix using the Mantel test.
aop calculates the individual contribution using an "add-one-patient" approach. The residuals of a single patient are added to those of a control group, and a correlation matrix is created. This is repeated for all individual patients and each patient group.
The summary method prints the group/region-wise means and standard deviations.
The plot method is only valid for regional contribution estimates, and plots the average regional contribution for each vertex/region.

## Usage

```
loo(resids, corrs, level = c("global", "regional"))
aop(resids, corrs, level = c("global", "regional"), control.value = 1L)
## S3 method for class 'IC'
summary(object, region = NULL, digits = max(3L,
        getOption("digits") - 2L), ...)
## S3 method for class 'IC'
plot(x, plot.type = c("mean", "smooth", "boxplot"),
    region = NULL, ids = TRUE, ...)
```


## Arguments

| resids | An object of class brainGraph_resids (the output from get.resid) |
| :--- | :--- |
| corrs | List of lists of correlation matrices (as output by corr.matrix). <br> Character string; the level at which you want to calculate contributions (either <br> global or regional) |
| control.value | Integer or character string specifying the control group (default: 1L) |
| object, x | A IC object <br> Character vector specifying which regions' IC's to print. Only relevant if method= 'Leave <br> one out ' |
| digits | Integer specifying the number of digits to display for P-values |
| $\ldots$ | Unused <br> plot.type |
| Character string indicating the type of plot; the default is to plot the mean (along <br> with standard errors) |  |
| ids | Logical indicating whether to plot Study ID's for outliers. Otherwise plots the <br> integer index |

## Value

A data. table with columns for

| Study.ID | Subject identifier |
| :--- | :--- |
| Group | Group membership |
| region | If level='regional' |
| IC, RC | The value of the individual/regional contributions |

## Note

For aop, it is assumed by default that the control group is the first group.

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## References

Saggar, M. and Hosseini, S.M.H. and Buno, J.L. and Quintin, E. and Raman, M.M. and Kesler, S.R. and Reiss, A.L. (2015) Estimating individual contributions from group-based structural correlations networks. NeuroImage, 120, 274-284. https://dx.doi.org/10.1016/j.neuroimage. 2015.07.006

## See Also

Other Structural covariance network functions: Bootstrapping, Residuals, brainGraph_permute, corr.matrix, import_scn, plot_volumetric

## Examples

```
## Not run:
IC <- loo(resids.all, corrs)
RC <- loo(resids.all, corrs, level='regional')
## End(Not run)
## Not run:
IC <- aop(resids.all, corrs)
RC <- aop(resids.all, corrs, level='regional')
## End(Not run)
```

Inverse $\quad$ Calculate the inverse of the cross product of a design matrix

## Description

inv is a S3 generic that calculates the inverse of the cross product of a design matrix, also referred to as the "unscaled covariance matrix".
pinv calculates $M^{+}=\left(M^{T} M\right)^{-1} M^{T}$ for full (column) rank matrices. However, it does not verify the matrix's rank.

## Usage

```
inv(x, ...)
## S3 method for class 'matrix'
inv(x, y = NULL, transpose = FALSE, ...)
## S3 method for class 'array'
inv(x, y = NULL, transpose = FALSE, ...)
## S3 method for class 'qr'
inv(x, p = x$rank, ...)
```

```
## S3 method for class 'list'
inv(x, p = x[[1L]]$rank, r = length(x),
    vnames = dimnames(x[[1L]]$qr)[[2L]], nms = names(x), ...)
pinv(x)
```


## Arguments

x
... Unused
y A numeric matrix or vector (for the matrix and array methods). If supplied, this will be multiplied by $x$ before the inverse is calculated. Default: NULL
transpose Logical. If FALSE (the default), take the cross product of the arguments. If TRUE, use tcrossprod
$\mathrm{p} \quad$ The rank of the original matrix
$r \quad$ The number of design matrices; i.e., the length of the input list
vnames $\quad$ Character vector of the design matrix's variable names
nms The region names; i.e., the names of the input list

## Details

If $x$ is a matrix, the Cholesky decomposition of the cross product is calculated (or using tcrossprod if transpose=TRUE), and the inverse is calculated from that result. That is,

$$
\begin{gathered}
\operatorname{inv}(X)=\left(X^{T} X\right)^{-1} \\
\operatorname{inv}(X, \operatorname{transpose}=T R U E)=\left(X X^{T}\right)^{-1} \\
\operatorname{inv}(X, y)=\left(X^{T} y\right)^{-1}
\end{gathered}
$$

If $x$ is a 3-dimensional array, then the inverse will be calculated for each matrix along the 3rd dimension, with the same input arguments for each.
Finally, there is a method for objects with class qr, and lists of QR decomposition objects.

## Value

A numeric matrix or array
pinv returns the input matrix's pseudoinverse

## Note

These methods should only be used on full-rank matrices, as there is no error checking being performed.

## Description

Given a list of brainGraphList objects, this function will calculate the area under the curve (AUC) across all thresholds/densities for each subject or group.

## Usage

make_auc_brainGraph(g.list, g.attr = NULL, v.attr = NULL, norm = FALSE)

## Arguments

g.list A list of brainGraphList objects
g.attr A character vector of graph attribute name(s). Default: NULL
v.attr A character vector of vertex attribute name(s). Default: NULL
norm Logical indicating whether to normalize threshold values to be between 0 and 1 (inclusive). Default: FALSE

## Details

If the elements of the input list do not have a threshold element (or if it is NULL) then the AUC will be calculated on the interval $[0,1]$ (inclusive). This has the same effect as specifying norm=TRUE in the function call.

## Value

A brainGraphList object with one graph for each subject

## Examples

```
## Not run:
g.auc <- make_auc_brainGraph(g.fa, g.attr='E.global.wt')
## End(Not run)
```

make_ego_brainGraph Create a graph of the union of multiple vertex neighborhoods

## Description

This function accepts multiple vertices, creates graphs of their neighborhoods (of order 1), and returns the union of those graphs.

## Usage

make_ego_brainGraph(g, vs)

## Arguments

g
An igraph graph object
vs Either a character or integer vector (vertex names or indices, respectively) for the vertices of interest

## Value

An igraph graph object containing the union of all edges and vertices in the neighborhoods of the input vertices; only the vertex attribute name will be present

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## See Also

ego
Other Graph creation functions: Creating_Graphs_GLM, Creating_Graphs, brainGraphList

## Examples

```
## Not run:
subg <- make_ego_brainGraph(g1[[N]], c(24, 58))
subg <- make_ego_brainGraph(g1[[N]], c('lPCUN', 'rPCUN'))
## End(Not run)
```

```
make_intersection_brainGraph
```

Create the intersection of graphs based on a logical condition

## Description

Returns a graph object with vertices that meet certain criteria. By default, only vertices that meet these criteria for all input graphs will be retained.

## Usage

make_intersection_brainGraph(..., subgraph, keep.all.vertices = FALSE)

## Arguments

$$
\begin{array}{ll}
\ldots . & \text { Graph objects or lists of graph objects } \\
\text { subgraph } & \text { Character string specifying an equation (logical condition) for the vertices to } \\
\text { subset }
\end{array} \begin{aligned}
& \text { keep.all.vertices } \\
& \\
& \\
& \\
& \\
& \text { Logical indicating whether to keep all vertices that meet the criteria in at least } 1 \\
& \text { inperault: FALSE }
\end{aligned}
$$

## Details

If no vertices meet criteria for all input graphs, then an igraph graph object with 0 vertices is returned. If keep.all.vertices=TRUE, this is essentially performing a union of vertex sets that meet the criteria. In any case, the return graph will have 0 edges.

## Value

An igraph graph object

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## Examples

```
## Not run:
res.mtpc <- mtpc(g, covars, ...)
g.mtpc <- make_glm_brainGraph(res.mtpc, atlas)
## All vertices with a significant MTPC result for all contrasts:
g.mtpc.int <- make_intersection_brainGraph(g.mtpc, subgraph='sig == 1')
## Return graphs with vertices with degree > 0 for each group separately
tapply(g.list, groups(g.list), make_intersection_brainGraph,
    subgraph='degree > 0')
```

```
## End(Not run)
```

Matrix utilities Matrix/array utility functions

## Description

These functions are utility/helper functions when working with matrices or arrays.
diag_sq is a pared-down version of diag for square matrices. It does not return any dimnames, does not check if $x$ is a square matrix, and it cannot be used to create a matrix with a given value along the diagonal. Meant to be used in code that is called repeatedly (thousands of times).
get_thresholds calculates the threshold values that would result in a specific graph density. These depend, necessarily on the values in the matrix themselves.
qr . array will calculate the QR decomposition for each matrix in a 3D array.
qr_Q2 and qr_R2 are simplified versions of qr. Q and qr.R.
symm_mean returns a symmetric matrix in which the off-diagonal elements $A[i, j]$ and $A[j, i]$ are set to the mean of the values in the input matrix.
symmetrize will symmetrize a numeric matrix (or each matrix in an array) by assigning to the off-diagonal elements either the max (default), min, or average of $\{A(i, j), A(j, i)\}$.

## Usage

```
colMax(x, n = dim(x)[1L])
colMaxAbs(x, n = dim(x)[1L])
colMin(x, n = dim(x)[1L])
diag_sq(x, n = dim(x)[1L], inds = 1L + 0L:(n - 1L) * (n + 1L))
get_thresholds(x, densities, emax = dim(x)[1L] * (dim(x)[1L] - 1L)/2,
    ...)
is_binary(x)
## S3 method for class 'array'
qr(x, ...)
qr_Q2(QR, y = diag(1, n, p), n = dim(QR$qr)[1L], p = QR$rank)
qr_R2(QR, p = QR$rank)
symm_mean(x)
```

```
symmetrize(x, ...)
\#\# S3 method for class 'matrix'
symmetrize(x, symm.by = c("max", "min", "avg"), ...)
\#\# S3 method for class 'array'
symmetrize(x, symm.by = c("max", "min", "avg"), ...)
```


## Arguments

$x \quad$ Numeric matrix or array (the latter, for qr.array and symmetrize.array)
$\mathrm{n}, \mathrm{p} \quad$ Integer; the number of rows or rank (respectively) of the input matrix or QR decomposition
inds Vector-based indices of the diagonal
densities Numeric vector of densities
emax Integer; the maximum number of edges
Arguments passed to either sort (for get_thresholds) or qr.default (for qr.array). For the former, this will typically only be decreasing=TRUE, if that is the desired behavior
QR A qr object
y A numeric matrix with 1 along the diagonal, of the same size as the input matrix (i.e., QR\$qr)
symm.by Character string; how to create symmetric off-diagonal elements. Default: max

## Details

Given a vector of densities, get_thresholds returns the numeric values that will result in graphs of the given densities after thresholding by those values. In the Examples section, the thresholds should result in graphs with densities of $5,15, \ldots, 55$ percent.

## Value

diag_sq returns an unnamed numeric vector with the values along the diagonal of the input matrix get_thresholds returns a numeric vector of the thresholds
is_binary returns a logical of length 1
qr.array returns a list in which each element is the QR decomposition of each matrix along x's 3rd dimension

## Examples

```
x <- matrix(runif(25 * 25), 25, 25)
x <- symmetrize(x)
diag(x) <- 0
densities <- seq(0.05, 0.55, by=0.1)
threshes <- get_thresholds(x, densities)
## Verify that the densities are correct
```

```
graphs <- lapply(threshes, function(th) {
    graph_from_adjacency_matrix(x * (x > th), mode='undirected',
                            diag=FALSE, weighted=TRUE)
    })
sapply(graphs, graph.density)
```

```
mean_distance_wt Calculate weighted shortest path lengths
```


## Description

Calculate graph or vertex average shortest path lengths. For vertices, this is just the row means of the distance matrix. For the graph-level, it is the overall mean of the distance matrix.

## Usage

mean_distance_wt(g, level = c("graph", "vertex"), weights = NULL, xfm = FALSE, xfm.type $=$ NULL, $D=N U L L)$

## Arguments

| g | An igraph graph object |
| :--- | :--- |
| level | Character string indicating whether to calculate vertex- or graph-level shortest <br> path length. Default: 'graph' |
| weights | Numeric vector of edge weights; if NULL (the default), and if the graph has edge <br> attribute weight, then that will be used. To avoid using weights, this should be <br> NA. |
| xfm | Logical indicating whether to transform the edge weights. Default: FALSE |
| xfm.type | Character string specifying how to transform the weights. Default: $1 / \mathrm{w}$ |
| D | Numeric matrix; the graph's "distance matrix" |

## Details

By default, edge weights are not transformed (e.g., inverted). However, if set to TRUE, then the input graph must have a graph-level attribute called 'xfm.type' or you must supply a value in the function call. If you supply a distance matrix (the $D$ argument), it is not necessary to transform edge weights, as it is assumed the the distance matrix was calculated from a graph with transformed edge weights already.

## Value

Numeric vector (if level='vertex') of each vertex's shortest path length, or a single number for the graph-level average

## Description

brainGraph_mediate performs simple mediation analyses in which a given graph- or vertex-level measure (e.g., weighted global efficiency) is the mediator $M$. The outcome (or dependent/response)
variable $Y$ can be a neuropsychological measure (e.g., $I Q$ ) or can be a disease-specific metric (e.g., recovery time).
bg_to_mediate converts the results into an object of class mediate. In brainGraph, it is only used for the summary.mediate method, but you can similarly use its output for the plot.mediate method.

## Usage

brainGraph_mediate(g.list, covars, mediator, treat, outcome, covar.names, level = c("graph", "vertex"), control.value = 0, treat.value = 1, int = FALSE, boot = TRUE, boot.ci.type = c("perc", "bca"), $\mathrm{N}=1000$, conf.level $=0.95$, long = FALSE, ...)
\#\# S3 method for class 'bg_mediate'
summary (object, mediate = FALSE, region = NULL,
digits $=\max (3 \mathrm{~L}$, getOption("digits") - 2L), ...)
bg_to_mediate(x, region = NULL)

## Arguments

| g.list | A brainGraphList object |
| :---: | :---: |
| covars | A data table containing covariates of interest. It must include columns for getOption('bg.subject_id'), treat, outcome, and covar. names. |
| mediator | Character string; the name of the graph measure acting as the mediating variable |
| treat | Character string; the treatment variable (e.g., Group) |
| outcome | Character string; the name of the outcome variable of interest |
| covar.names | Character vector of the column name(s) in covars to include in the models as pre-treatment covariate(s). |
| level | Character string; either vertex (default) or graph |
| control.value | Value of treat to be used as the control condition. Default: 0 |
| treat.value | Value of treat to be used as the treatment condition. Default: 1 |
| int | Logical indicating whether or not to include an interaction of the mediator and treatment. Default: FALSE |
| boot | Logical indicating whether or not to perform bootstrapping. This should always be done. Default: TRUE |


| boot.ci.type | Character string; which type of CI's to calculate. Default: perc |
| :---: | :---: |
| N | Integer; the number of bootstrap samples to run. Default: 1e3 |
| conf.level | Numeric between 0 and 1 ; the level of the CI's to calculate. Default: 0.95 for the 2.5 and 97.5 percentiles) |
| long | Logical indicating whether or not to return all bootstrap samples. Default: FALSE |
|  | Other arguments passed to brainGraph_GLM_design (e.g., binarize) (unused in the summary method) |
| object | A bg_mediate object |
| mediate | Logical indicating whether or not to use the summary method from mediate (default: FALSE). If TRUE, only a single region can be printed. |
| region | Character string specifying which region's results to summarize; only relevant if level=' vertex' (default: NULL) |
| digits | Integer specifying the number of digits to display for P-values |
| x | Object output from brainGraph_mediate |

## Details

This code was adapted closely from mediate in the mediation package, and the procedure is exactly the same as theirs (see the references listed below). If you use this function, please cite their work.

## Value

An object of class bg_mediate with elements:

| level | Either graph or vertex. |
| :---: | :---: |
| removed.subs | A character vector of Study.ID's removed due to incomplete data |
| X.m, X.y | Design matrix and numeric array for the model with the mediator as the outcome variable (X.m) and for the model with the mediator as an additional predictor (X.y), respectively |
| y.m, y.y | Outcome variables for the associated design matrices above. y.m will be a matrix of size \# subj. X \# regions |
| res.obs | A data. table of the observed values of the point estimates. |
| res.ci | A data.table of the confidence intervals for the effect estimates. |
| res.p | A data. table of the two-sided p-values for the effect estimates |
| boot | Logical, the boot argument. |
| boot.ci.type | Character string indicating which type of bootstrap confidence intervals were calculated. |
| res.boot | A data. table with $N$ rows of the bootstrap results for all effects. |
| treat | Character string of the treatment variable. |
| mediator | Character string of the mediator variable. |
| outcome | Character string of the outcome variable. |


| covariates | Returns NULL; not used in this package. |
| :--- | :--- |
| INT | Logical indicating whether the models included an interaction between treat- <br> ment and mediator. |
| conf.level | The confidence level. |
| control.value | The value of the treatment variable used as the control condition. |
| treat.value | The value of the treatment variable used as the treatment condition. |
| nobs | Integer; the number of observations in the models. |
| sims | Integer; the number of bootstrap replications. |
| covar.names | The pre-treatment covariate names. |
| bg_to_mediate returns an object of class mediate |  |

Note
As of brainGraph v2.0.0, this function has been tested only for a treatment (independent) variable $X$ being a factor (e.g., disease group, old vs. young, etc.). If your treatment variable has more than 2 levels, then you must explicitly specify the levels you would like to compare; otherwise, the baseline and first levels are taken to be the control and treatment values, respectively. Be aware that these are 0 indexed; that is, if you have 3 groups and you would like the treatment group to be the 3rd, you should specify as either the group's character string or as treat. value=2.
Allowing for treatment-mediator interaction (setting int=TRUE) currently will only work properly if the mediator is a continuous variable; since the mediator is always a graph metric, this should always be the case.

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## References

Tingley, D. and Yamamoto, T. and Hirose, K. and Keele, L. and Imai, K. (2014) mediation: R package for causal mediation analysis. Journal of Statistical Software, 59(5), 1-38. https://dx. doi.org/10.18637/jss.v059.i05
Imai, K. and Keele, L. and Yamamoto, T. (2010) Identification inference, and sensitivity analysis for causal mediation effects. Statistical Science, 25(1), 51-71. https://dx.doi.org/10.1214/ 10-STS321

Imai, K. and Keele, L. and Tingley, D. (2010) A general approach to causal mediation analysis. Psychological Methods, 15(4), 309-334. https://dx.doi.org/10.1037/a0020761
Imai, K. and Keele, L. and Tingley, D. and Yamamoto, T. (2011) Unpacking the black box of causality: learning about causal mechanisms from experimental and observational studies. American Political Science Review, 105(4), 765-789. https://dx.doi.org/10.1017/S0003055411000414

Imai, K. and Yamamoto, T. (2013) Identification and sensitivity analysis for multiple causal mechanisms: revisiting evidence from framing experiments. Political Analysis, 21(2), 141-171. https:
//dx.doi.org/10.1093/pan/mps040

## See Also

## mediate

Other Group analysis functions: Bootstrapping, GLM, NBS, brainGraph_permute, mtpc

## Examples

```
## Not run:
med.EglobWt.FSIQ <- brainGraph_mediate(g[[5]], covars.med, 'E.global.wt',
    'Group', 'FSIQ', covar.names=c('age', 'gender'), N=1e4)
med.strength.FSIQ <- brainGraph_mediate(g[[5]], covars.med, 'strength',
    'Group', 'FSIQ', covar.names=c('age', 'gender'), level='vertex')
## End(Not run)
```

mtpc Multi-threshold permutation correction

## Description

Applies the multi-threshold permutation correction (MTPC) method to perform inference in graph theory analyses of brain MRI data.

Plot the statistics from an MTPC analysis, along with the maximum permuted statistics. The output is similar to Figure 11 in Drakesmith et al. (2015).

## Usage

```
mtpc(g.list, thresholds, covars, measure, contrasts, con.type = c("t",
    "f"), outcome = NULL, con.name = NULL, level = c("vertex",
    "graph"), clust.size = 3L, perm.method = c("freedmanLane",
    "terBraak", "smith", "draperStoneman", "manly", "stillWhite"),
    part.method = c("beckmann", "guttman", "ridgway"), N = 500L,
    perms = NULL, alpha = 0.05, res.glm = NULL, long = TRUE, ...)
## S3 method for class 'mtpc'
summary(object, contrast = NULL, digits = max(3L,
    getOption("digits") - 2L), print.head = TRUE, ...)
## S3 method for class 'mtpc'
plot(x, contrast = 1L, region = NULL,
    only.sig.regions = TRUE, show.null = TRUE, caption.stats = FALSE,
    ...)
## S3 method for class 'mtpc'
nobs(object, ...)
## S3 method for class 'mtpc'
```

```
terms(x, ...)
## S3 method for class 'mtpc'
formula(x, ...)
## S3 method for class 'mtpc'
labels(object, ...)
## S3 method for class 'mtpc'
case.names(object, ...)
## S3 method for class 'mtpc'
variable.names(object, ...)
## S3 method for class 'mtpc'
df.residual(object, ...)
## S3 method for class 'mtpc'
region.names(object)
## S3 method for class 'mtpc'
nregions(object)
```


## Arguments

| g.list | A list of brainGraphList objects for all thresholds |
| :---: | :---: |
| thresholds | Numeric vector of the thresholds applied to the raw connectivity matrices. |
| covars | A data. table of covariates |
| measure | Character string of the graph measure of interest |
| contrasts | Numeric matrix (for T statistics) or list of matrices (for F statistics) specifying the contrast(s) of interest; if only one contrast is desired, you can supply a vector (for T statistics) |
| con.type | Character string; either ' $t$ ' or ' f ' (for t or F-statistics). Defaul |
| outcome | Character string specifying the name of the outcome variable, if it differs from the graph metric (measure) |
| con. name | Character vector of the contrast name(s); if contrasts has row/list names, those will be used for reporting results |
| level | Character string; either vertex (default) or graph |
| clust.size | Integer indicating the size of "clusters" (i.e., consecutive thresholds for which the observed statistic exceeds the null) (default: 3L) |
| perm.method | Character string indicating the permutation method. Default: 'freedmanLane' |
| part.method | Character string; the method of partitioning the design matrix into covariates of interest and nuisance. Default: 'beckmann' |
| $N$ | Integer; number of permutations to create. Default: 5e3 |
| perms | Matrix of permutations, if you would like to provide your own. Default: NULL |


| alpha |  |
| :--- | :--- |
| res.glm | Numeric; the significance level. Default: 0.05 <br> A list of bg_GLM objects, as output by a previous run of mtpc. Useful if you want <br> to change the cluster size without re-running all of the GLM's and permutations <br> (default: NULL) |
| long | Logical indicating whether or not to return all permutation results. Default: <br> FALSE |
| Other arguments passed to brainGraph_GLM and/or brainGraph_GLM_design |  |
| object, x | A mtpc object |
| contrast | Integer specifying the contrast to plot/summarize; defaults to showing results <br> for all contrasts |
| digits | Integer specifying the number of digits to display for P-values |
| print.head | Logical indicating whether or not to print only the first and last 5 rows of the <br> statistics tables (default: TRUE) |
| region | Character string specifying which region's results to plot; only relevant if level=' vertex' <br> Default: NULL |
| only.sig.regions |  |
| Logical indicating whether to plot only significant regions (default: TRUE) |  |

## Details

This is a multi-step procedure: (steps 3-4 are the time-consuming steps)

1. Apply thresholds $\tau$ to the networks, and compute network metrics for all networks and thresholds. (already done beforehand)
2. Compute test statistics $S_{o b s}$ for each threshold. (done by brainGraph_GLM)
3. Permute group assignments and compute test statistics for each permutation and threshold. (done by brainGraph_GLM)
4. Build a null distribution of the maximum statistic across thresholds (and across brain regions) for each permutation. (done by brainGraph_GLM)
5. Determine the critical value, $S_{\text {crit }}$ from the null distribution of maximum statistics.
6. Identify clusters where $S_{o b s}>S_{c r i t}$ and compute the AUC for these clusters (denoted $A_{M T P C}$ ).
7. Compute a critical AUC $\left(A_{\text {crit }}\right)$ from the mean of the supra-critical AUC's for the permuted tests.
8. Reject $H_{0}$ if $A_{M T P C}>A_{\text {crit }}$.

## Value

An object of class mtpc with some input arguments plus the following elements:

| X, qr, cov.unscaled |  |
| :---: | :---: |
|  | Design matrix, QR decomposition, and unscaled covariance matrix, if the design is the same across thresholds |
| contrasts | The contrast matrix or list of matrices |
| con. name | Contrast names |
| removed.subs | Named integer vector of subjects with incomplete data |
| atlas | The atlas of the input graphs |
| rank,df.residual |  |
|  | The model rank and residual degrees of freedom |
| res.glm | List with length equal to the number of thresholds; each list element is the output from brainGraph_GLM |
| DT | A data.table for all thresholds, combined from the outputs of brainGraph_GLM |
| stats | A data.table containing S.mtpc (the max. observed statistic), tau.mtpc (the threshold of the max. observed statistic), S.crit (the critical statistic value), and A.crit (the critical AUC) |
| null.dist | Numeric array with $N$ columns and number of rows equal to the number of thresholds. The 3rd dimension is for each contrast. Each element of the array is the maximum statistic for that permutation, threshold, and contrast combination. |
| perm.order | Numeric matrix; the permutation set applied for all thresholds (each row is a separate permutation) |

The plot method returns a trellis object or a list of ggplot objects

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## References

Drakesmith, M. and Caeyenberghs, K. and Dutt, A. and Lewis, G. and David, A.S. and Jones, D.K. (2015) Overcoming the effects of false positives and threshold bias in graph theoretical analyses of neuroimaging data. NeuroImage, 118, 313-333. https://dx.doi.org/10.1016/j.neuroimage. 2015.05.011

## See Also

Other Group analysis functions: Bootstrapping, GLM, Mediation, NBS, brainGraph_permute Other GLM functions: GLM design, GLM fits, GLM

## Examples

```
## Not run:
diffs.mtpc <- mtpc(g.list=g.norm, thresholds=thresholds, N=N,
    covars=covars.dti, measure='E.nodal.wt', coding='effects',
    contrasts=c(0, 0, 0, 0, -2), alt='greater',
    binarize=c('Sex', 'Scanner'), con.name='Group 1 > Group 2')
sig.regions <- diffs.mtpc$DT[A.mtpc > A.crit]
```

```
## End(Not run)
## Not run:
mtpcPlots <- plot(mtpc.diffs)
## Arrange plots into 3x3 grids
ml <- marrangeGrob(mtpcPlots, nrow=3, ncol=3)
ggsave('mtpc.pdf', ml)
## End(Not run)
```


## Description

Calculates the network-based statistic (NBS), which allows for family-wise error (FWE) control over network data, introduced for brain MRI data by Zalesky et al. Requires a three-dimensional array of all subjects' connectivity matrices and a data. table of covariates, in addition to a contrast matrix or list. A null distribution of the largest connected component size is created by fitting a GLM to permuted data. For details, see GLM.

## Usage

```
NBS(A, covars, contrasts, con.type = c("t", "f"), X = NULL,
    con.name = NULL, p.init = 0.001, perm.method = c("freedmanLane",
    "terBraak", "smith", "draperStoneman", "manly", "stillWhite"),
    part.method = c("beckmann", "guttman", "ridgway"), N = 1000,
    perms = NULL, symm.by = c("max", "min", "avg"),
    alternative = c("two.sided", "less", "greater"), long = FALSE, ...)
## S3 method for class 'NBS'
summary(object, contrast = NULL, digits = max(3L,
    getOption("digits") - 2L), ...)
## S3 method for class 'NBS'
nobs(object, ...)
## S3 method for class 'NBS'
terms(x, ...)
## S3 method for class 'NBS'
formula(x, ...)
## S3 method for class 'NBS'
labels(object, ...)
## S3 method for class 'NBS'
```

```
case.names(object, ...)
## S3 method for class 'NBS'
variable.names(object, ...)
## S3 method for class 'NBS'
df.residual(object, ...)
## S3 method for class 'NBS'
nregions(object)
```


## Arguments

| A | Three-dimensional array of all subjects' connectivity matrices |
| :--- | :--- |
| covars | A data.table of covariates |
| contrasts | Numeric matrix (for T statistics) or list of matrices (for F statistics) specifying <br> the contrast(s) of interest; if only one contrast is desired, you can supply a vector <br> (for T statistics) |
| con.type | Character string; either ' t ' or ' f ' (for t or F-statistics). Default: ' t ' |
| X |  |

## Details

When printing a summary, you can include arguments to printCoefmat.

## Value

An object of class NBS with some input arguments in addition to:

X
The design matrix
removed. subs Character vector of subject ID's removed due to incomplete data (if any)
T.mat 3-d array of (symmetric) numeric matrices containing the statistics for each edge
p.mat 3-d array of (symmetric) numeric matrices containing the P -values
components List containing data tables of the observed and permuted connected component sizes and P -values
rank, df.residual, qr, cov. unscaled
The rank, residual degrees of freedom, QR decomposition, and unscaled covariance matrix of the design matrix

## Note

It is assumed that the order of the subjects in covars matches that of the input array A. You will need to ensure that this is the case. Prior to v3.0.0, the covars table was sorted by Study.ID before creating the design matrix.

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## References

Zalesky, A. and Fornito, A. and Bullmore, E.T. (2010) Network-based statistic: identifying differences in brain networks. NeuroImage, 53(4), 1197-1207. https://dx.doi.org/10.1016/j. neuroimage. 2010.06.041

## See Also

Other Group analysis functions: Bootstrapping, GLM, Mediation, brainGraph_permute, mtpc

## Examples

```
## Not run:
max.comp.nbs <- NBS(A.norm.sub[[1]], covars.dti, N=5e3)
## End(Not run)
```


## Description

plot.brainGraph plots a graph in which the spatial layout of the nodes is important. The network itself is plotted over a brain MRI slice from the MNI152 template by default (when mni=TRUE).

## Usage

\#\# S3 method for class 'brainGraph'
plot(x, plane = c("axial", "sagittal", "circular"), hemi = c("both", "L", "R"), mni = TRUE, subgraph = NULL, main $=$ NULL, subtitle $=$ "default", label $=$ NULL, side $=1$, line $=-2$, adj $=0.025$, cex $=2.5$, col = "white", font $=2$, show.legend $=$ FALSE, rescale $=$ FALSE, asp $=0, \ldots$ )

## Arguments

X
plane Character string indicating which orientation to plot. Default: 'axial'
hemi Character string indicating which hemisphere to plot. Default: 'both'
mni Logical indicating whether or not to plot over a slice of the brain. Default: TRUE
subgraph Character string specifying a logical condition for vertices to plot. Default: NULL
main Character string; the main title. Default: NULL
subtitle Character string; the subtitle. Default: 'default'
label Character string specifying text to display in one corner of the plot (e.g., 'A. '). Default: NULL
side Label placement. Default: 1 (bottom)
line Which margin line to place the text.
adj If side=1, a value closer to 0 places the text closer to the left margin. Default: 0.025
cex Amount of character expansion of the label text. Default: 2.5
col Label font color. Default: 'white'
font Integer specifying the font type. Default: 2 (bold face)
show. legend Logical indicating whether or not to show a legend. Default: FALSE
rescale Logical, whether to rescale the coordinates. Default: FALSE
asp Numeric constant; the aspect ratio. Default: 0
... Other parameters (passed to plot.igraph). See plot.common for details.

## Selecting specific vertices to display

With the argument subgraph, you can supply a simple logical expression specifying which vertices to show. For example, 'degree $>10$ ' will plot only vertices with a degree greater than 10. Combinations of $A N D$ (i.e., \&) and $O R$ (i.e., |) are allowed. This requires that any vertex attribute in the expression must be present in the graph; e.g., $\mathrm{V}(\mathrm{g}) \$$ degree must exist.

## Title, subtitle, and label

By default, a title (i.e., text displayed at the top of the figure) is not included. You can include one by passing a character string to main, and control the size with cex.main. A subtitle (i.e., text at the bottom), is included by default and displays the number of vertices and edges along with the graph density. To exclude this, specify subtitle=NULL. A "label" can be included in one corner of the figure (for publications). For example, you can choose label='A. ' or label='a)'. Arguments controlling the location and appearance can be changed, but the default values are optimal for bottom-left placement. See mtext for more details. The label-specific arguments are:
side The location. 1 is for bottom placement.
line If side=1 (bottom), a negative number places the text above the bottom of the figure; a higher number could result in the bottom part of the text to be missing. This can differ if plane='circular', in which case you may want to specify a positive number.
adj Seems to be the percentage away from the margin. So, for example, adj=0.1 would place the text closer to the center than the default value, and adj=0.5 places it in the center.
cex The degree of "character expansion". A value of 1 would not increase the text size.
col The text color.
font The font type. The default font=2 is bold face. See par for details.

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## See Also

Other Plotting functions: Plotting GLM graphs, plot.brainGraphList, plot_brainGraph_gui, plot_brainGraph_multi

## Examples

```
## Not run:
plot(g[[1]], hemi='R')
plot(g[[1]], subgraph='degree > 10 | btwn.cent > 50')
## Place label in upper-left
plot(g.ex, label='A)', side=3, line=-2.5)
## End(Not run)
```


## Description

The plot method will write a PDF file containing plots for all graphs in the given object.

## Usage

```
## S3 method for class 'brainGraphList'
plot(x, plane, hemi, filename.base,
        diffs = FALSE, ...)
```


## Arguments

x
plane Character string indicating which orientation to plot. Default: 'axial'
hemi Character string indicating which hemisphere to plot. Default: 'both'
filename.base Character string specifying the base of the filename
diffs Logical, indicating whether edge differences should be highlighted. Default: FALSE
... Other parameters (passed to plot.brainGraph)

## Details

You can choose to highlight edge differences between subsequent list elements; in this case, new/different edges are colored pink. This is useful mostly for a list of group-level graphs.

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## See Also

Other Plotting functions: Plotting GLM graphs, plot.brainGraph, plot_brainGraph_gui, plot_brainGraph_multi

## Description

These methods are convenience functions for plotting a graph based on results from GLM-based analyses (i.e., brainGraph_GLM, brainGraph_mediate, mtpc, NBS). There are several default arguments which differ depending on the input object.

## Usage

```
    ## S3 method for class 'brainGraph_NBS'
    plot(x, alpha = 0.05,
        subgraph = paste("p.nbs >", 1 - alpha), vertex.label = NA,
        vertex.color = "color.comp", edge.color = "color.comp",
        subtitle = NULL, main = paste0("NBS: ", x$name), cex.main = 2, ...)
    ## S3 method for class 'brainGraph_GLM'
    plot(x, p.sig = c("p", "p.fdr", "p.perm"),
        subgraph = NULL, main = paste0(x$outcome, ": ", x$name),
        subtitle = NULL, cex.main = 2, ...)
    ## S3 method for class 'brainGraph_mtpc'
    plot(x, subgraph = "sig == 1",
        main = paste0(x$outcome, ": ", x$name), subtitle = NULL,
        cex.main = 2, ...)
    ## S3 method for class 'brainGraph_mediate'
    plot(x, subgraph = "p.acme > 0.95",
        main = sprintf("Effect of \"%s\" on\n\"%s\"\nmediated by \"%s\"",
        x$treat, x$outcome, x$mediator), subtitle = NULL, cex.main = 1, ...)
```


## Arguments

x
alpha Numeric; the significance level. Default: 0.05
subgraph Character string specifying the condition for subsetting the graph.
vertex.label Character vector of the vertex labels to be displayed.
vertex.color Character string specifying the vertex attribute to color the vertices by.
edge.color Character string specifying the edge attribute to color the edges by.
subtitle Character string; the subtitle. Default: 'default'
main Character string; the main title. Default: NULL
cex.main Numeric; the scaling factor for text size; see par
... Other arguments passed to plot.brainGraph
p.sig Character string indicating which p-value to use for determining significance (default: p)

## Details

The default arguments are specified so that the user only needs to type plot ( $x$ ) at the console, if desired. For all methods, the plot's subtitle will be omitted.

## NBS

By default, a subgraph will be plotted consisting of only those vertices which are part of a significant connected component. Vertex/edge colors will correspond to connected component membership. Vertex names will be omitted. Finally, the plot title will contain the contrast name.

## brainGraph_GLM

By default, a subgraph will be plotted consisting of only those vertices for which $p<\alpha$. It will also include a plot title with the outcome measure and contrast name.

## mtpe

By default, a subgraph will be plotted consisting of only those vertices for which $A_{m t p c}>A_{\text {crit }}$. It will also include a plot title with the outcome measure and contrast name.

## brainGraph_mediate

By default, a subgraph will be plotted consisting of only those vertices for which $P_{a c m e}<\alpha$. It will also include a plot title with the treatment, mediator, and outcome variable names.

## See Also

Other Plotting functions: plot.brainGraphList, plot.brainGraph, plot_brainGraph_gui, plot_brainGraph_multi

```
plot_brainGraph_gui GUI for plotting graphs overlaid on an MNI152 image or in a circle
```


## Description

This function creates a GUI for plotting graphs over an image from the MNI152 template. It gives the user control over several plotting parameters. Also possible is a circular plot (in addition to the axial and sagittal views). It is necessary for the graphs to have an atlas attribute, and several vertexand edge-level attributes (set by set_brainGraph_attr).

## Usage

plot_brainGraph_gui()

## See Also

Other Plotting functions: Plotting GLM graphs, plot.brainGraphList, plot.brainGraph, plot_brainGraph_multi
plot_brainGraph_multi Save PNG of one or three views for all graphs in a brainGraphList

## Description

plot_brainGraph_multi writes a PNG file to disk containing three views (columns) of 1 or more brainGraph objects (from left-to-right): left sagittal, axial, and right sagittal. The number of rows in the figure will equal the number of graphs to plot.
slicer writes a PNG file to disk containing a single view (i.e., either sagittal, axial, or circular) of all brainGraph objects in the input list/brainGraphList.

## Usage

plot_brainGraph_multi(g.list, filename = "orthoview.png", subgraph $=$ NULL, main $=$ NULL, label $=$ NULL, cex.main $=1, \ldots$ )
slicer(g.list, nrows, ncols, plane = "axial", hemi = "both", filename = "all.png", main = NULL, cex.main = 1, ...)

## Arguments

| g.list | A brainGraphList or a list of brainGraph objects |
| :---: | :---: |
| filename | Character string of the filename of the PNG to be written. |
| subgraph | A vector or list of character strings to (optionally) subset the graph(s), possibly by multiple conditions |
| main | A vector or list of character strings to be placed in the main title of the center (axial) plot for each graph |
| label | A vector or list of character strings to be placed in one of the corners of the left plot (sagittal) in each row |
| cex.main | Numeric specifying the level of character expansion for the plot titles. Default: 1 (no expansion) |
|  | Other arguments passed to plot.brainGraph |
| nrows | Integer; the number of rows in the figure |
| ncols | Integer; the number of columns in the figure |
| plane | Character string indicating which orientation to plot. Default: 'axial' |
| hemi | Character string indicating which hemisphere to plot. Default: 'both' |

## Details

Whether the first input is a brainGraphList object or a list of brainGraph objects, all graphs in the object will be displayed in the figure. For plot_brainGraph_multi, this may be undesirable if you have more than 4 or 5 graphs in one object. You can choose fewer by using simple subsetting operations (see Examples below).

## Using subgraphs, titles, and labels

There are three arguments that can differ for each graph to be displayed. Each follows the same "rules". If you would like the same value applied to all graphs, you can specify a character string. If you would like a different value for each group, you must supply a vector or list with length equal to the number of graphs. If its length is less than the number of graphs, values will be recycled. To "skip" applying a value to one (or more) graph(s), you can use the NULL value only within a list (see the Examples below).
subgraph Can be used to apply one or more conditions for subsetting the graph(s).
main Controls the main plot title, which appears in the axial view along with each graph's name attribute. Depending on the level of the brainGraphList, this will either be a Study ID, Group name, or contrast name.
label Can be used to print a text label in a corner for each group/graph. For example, you can print a letter if you will refer to, e.g., "Figure 1A", "Figure 1B", etc.

## Note

All other arguments (passed to plot.brainGraph) will be applied to all graphs. For example, if you include vertex. label=NA in the function call, vertex labels will be omitted for all graphs.

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## See Also

Other Plotting functions: Plotting GLM graphs, plot.brainGraphList, plot.brainGraph, plot_brainGraph_gui

## Examples

```
## Not run:
## "g.hubs" contains 2 groups; apply same subset to both
plot_brainGraph_multi(g.hubs, filename='Figure01_hubs.png',
    subgraph='N > 0', vertex.color='color.lobe', vertex.size=15,
    show.legend=TRUE, vertex.label.cex=1.5)
## Single group, different subgraphs for both plots
## "g" is a "brainGraphList" object
gg <- g[rep(1, 3), drop=FALSE]
plot_brainGraph_multi(gg, filename='group1_5-6-7core.png',
    vertex.color='color.lobe', edge.color='color.lobe', vertex.label=NA,
    subgraph=as.list(paste('coreness >', 5:7)),
    main=as.list(paste('k-core', 5:7)))
```

```
\#\# Apply different subset for groups 1 \& 3; no subset for group 2
plot_brainGraph_multi(g, groups \(=1: 3\), vertex.label=NA,
    subgraph=list('degree > 5', NULL, 'degree > 4'))
\#\# End(Not run)
```

plot_global Plot global graph measures across densities

## Description

Create a faceted line plot of global graph measures across a range of graph densities, calculated from a list of brainGraphList objects. This requires that the variables of interest are graph-level attributes of the input graphs.

## Usage

```
    plot_global(g.list, xvar = c("density", "threshold"), vline = NULL,
        level.names = "default", exclude = NULL, perms = NULL,
        alt = "two.sided")
```


## Arguments

g.list List of brainGraphList objects; the length of this list should equal the number of thresholds/densities in the study
xvar A character string indicating whether the variable of interest is "density" or "threshold" (e.g. with DTI data)
vline $\quad$ Numeric of length 1 specifying the $x$-intercept if you would like to plot a vertical dashed line (e.g., if there is a particular density of interest). Default: NULL
level.names Character vector of variable names, which are displayed as facet labels. If you do not want to change them, specify NULL. By default, they are changed to preset values.
exclude $\quad$ Character vector of variables to exclude. Default: NULL
perms A data.table of permutation group differences
alt Character vector of alternative hypotheses; required if perms is provided, but defaults to "two.sided" for all variables

## Details

You can choose to insert a dashed vertical line at a specific density/threshold of interest, rename the variable levels (which become the facet titles), exclude variables, and include a brainGraph_permute object of permutation data to add asterisks indicating significant group differences.

## Value

Either a trellis or ggplot object

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>
plot_rich_norm Plot normalized rich club coefficients against degree threshold

## Description

Returns a line plot of the normalized rich club coefficient. Optionally, can include a shaded region demarcating the rich_core cutoff (if you supply a list of graph objects to the g argument).

## Usage

plot_rich_norm(rich.dt, facet.by = c("density", "threshold"), densities, alpha $=0.05$, fdr $=$ TRUE, g.list $=$ NULL, smooth $=$ TRUE)

## Arguments

| rich.dt | A data.table with rich-club coefficients |
| :--- | :--- |
| facet.by | A character string indicating whether the variable of interest is "density" or <br> "threshold" (e.g. with DTI data) |
| densities | A numeric vector of the densities to plot |
| alpha | The significance level. Default: 0.05 |
| fdr | A logical, indicating whether or not to use the FDR-adjusted p-value for deter- <br> mining significance. Default: TRUE |
| g.list | A list brainGraphList objects; required if you want to plot a shaded region <br> demarcating the rich_core |
|  | Logical indicating whether or not to plot a smooth curve when data from mul- <br> tiple subjects (per group) are present. Default: TRUE. Ignored for group-level <br> data. |

## Value

A trellis or ggplot object

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## See Also

Other Rich-club functions: Rich Club, rich_club_attrs

## Examples

```
## Not run:
plot_rich_norm(rich.dt, facet.by='density', densities[N:(N+1)], g=g)
## End(Not run)
```

```
plot_vertex_measures Plot vertex-level graph measures at a single density or threshold
```


## Description

Creates boxplots of a single vertex-level graph measure at a single density or threshold, grouped by the variable specified by group.by and optionally faceted by another variable (e.g., lobe or network).

## Usage

plot_vertex_measures(g.list, measure, facet.by = NULL, group.by = getOption("bg.group"), type = c("violin", "boxplot"), show.points = FALSE, ylabel = measure, ...)

## Arguments

| g.list | A brainGraphList or a list of brainGraph objects |
| :--- | :--- |
| measure | A character string of the graph measure to plot |
| facet.by | Character string indicating the variable to facet by (if any). Default: NULL |
| group.by | Character string indicating which variable to group the data by. Default: getOption('bg.group ' ) |
| type | Character string indicating the plot type. Default: 'violin' |
| show.points | Logical indicating whether or not to show individual data points (default: FALSE) |
| ylabel | A character string for the y-axis label |
| $\ldots$ | Arguments passed to geom_boxplot or geom_violin |

## Value

A trellis or ggplot object

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## Examples

```
## Not run:
p.deg <- plot_vertex_measures(g[[1]], facet.by='network', measure='degree')
## End(Not run)
```


## plot_volumetric Plot group distributions of volumetric measures for a given brain region

## Description

This function takes a "tidied" dataset of cortical volumetric measures (thickness, volume, LGI, etc.) and plots a histogram or violin plot for 1 or more groups, and of 1 or more brain regions.

## Usage

plot_volumetric(dat, regions, type = c("violin", "histogram"), all.vals = TRUE, modality = c("thickness", "volume", "lgi", "area"))

## Arguments

dat A data table of volumetric data; needs columns for 'Group', 'region', and 'value'
regions A vector of character strings or integers of the brain region(s) to plot; if integer, the region(s) is/are chosen from the input data table based on the index
type A character string indicating the plot type; either 'histogram' or 'violin'
all.vals A logical indicating whether or not to plot horizontal lines for all observations (only valid for 'violin' plots) (default: TRUE)
modality A character string indicating the type of volumetric measure ('thickness', 'volume', 'lgi', or 'area')

## Value

A trellis or ggplot object

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## See Also

Other Structural covariance network functions: Bootstrapping, IndividualContributions, Residuals, brainGraph_permute, corr.matrix, import_scn

## Description

analysis_random_graphs performs the steps needed for doing typical graph theory analyses with brain MRI data if you need to generate equivalent random graphs. This includes calculating small world parameters and normalized rich club coefficients.
sim. rand.graph.par simulates $N$ simple random graphs with the same clustering (optional) and degree sequence as the input. Essentially a wrapper for sample_degseq (or, if you want to match by clustering, sim.rand.graph.clust) and make_brainGraph. It uses foreach for parallel processing.
sim.rand.graph.clust simulates a random graph with a given degree sequence and clustering coefficient. Increasing the max.iters value will result in a closer match of clustering with the observed graph.
sim. rand.graph. hqs generates a number of random covariance matrices using the Hirschberger-Qi-Steuer (HQS) algorithm, and create graphs from those matrices.

## Usage

```
analysis_random_graphs(g.list, level = g.list[[1L]]$level, N = 100L,
    savedir = ".", ...)
    sim.rand.graph.par(g, level = c("subject", "group"), N = 100L,
        clustering = FALSE, rewire.iters = max(10 * ecount(g), 10000L),
        cl = g$transitivity, max.iters = 100L, ...)
    sim.rand.graph.clust(g, rewire.iters = 10000, cl = g$transitivity,
    max.iters = 100)
sim.rand.graph.hqs(resids, level = c("subject", "group"), N = 100L,
    weighted = TRUE, r.thresh = NULL, ...)
```


## Arguments

| g.list | List of brainGraphList objects; the length of this list should equal the number <br> of thresholds/densities in the study <br> Character string indicating whether the graphs are subject-, group-, or contrast- <br> specific. Default: ' subject ' |
| :--- | :--- |
| level | Integer; the number of random graphs to simulate. Default: 100 |
| savedir | Character string specifying the directory in which to save the generated graphs. <br> Default: current working directory |
| $\ldots$ | Other arguments passed to make_brainGraph |
| g | A graph object |


| clustering <br> rewire.iters | Logical; whether or not to control for clustering. Default: FALSE <br> Integer; number of rewiring iterations for the initial graph randomization. De- <br> fault: le4 |
| :--- | :--- |
| cl | The clustering measure. Default: transitivity |
| max.iters | The maximum number of iterations to perform; choosing a lower number may <br> result in clustering that is further away from the observed graph's. Default: 100 |
| resids | A brainGraph_resids object, a data.table of residuals, or a numeric matrix |
| weighted | Logical indicating whether to create weighted graphs. If true, a threshold must <br> be provided. |
| r.thresh | Numeric value for the correlation threshold, if weighted=FALSE. |

## Details

analysis_random_graphs does the following:

1. Generate $N$ random graphs for each graph and density/threshold
2. Write graphs to disk in savedir. Read them back into $R$ and combine into lists; then write these lists to disk. You can later delete the individual .rds files afterwards.
3. Calculate small world parameters, along with values for a few global graph measures that may be of interest.
4. Calculate normalized rich club coefficients and associated p-values.

If you do not want to match by clustering, then simple rewiring of the input graph is performed (the number of rewires equaling the larger of 1e4 and $10 \times m$, where $m$ is the graph's edge count).
sim.rand.graph.hqs - The first step is to create the observed covariance of residuals (or whatever matrix/data.table is provided). Then random covariance matrices are created with the same distributional properties as the observed matrix, they are converted to correlation matrices, and finally graphs from these matrices. By default, weighted graphs will be created in which the edge weights represent correlation values. If you want binary matrices, you must provide a correlation threshold.

## Value

analysis_random_graphs returns a list containing:
rich A data table containing normalized rich-club coefficients and p-values
small A data table with small-world parameters
rand A data table with some global graph measures for all random graphs generated
sim. rand.graph. par - a list of $N$ random graphs with some additional vertex and graph attributes
sim. rand.graph.clust - A single igraph graph object
sim.rand.graph.hqs - A list of random graphs from the null covariance matrices

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## References

Bansal, S. and Khandelwal, S. and Meyers, L.A. (2009) Exploring biological network structure with clustered random networks. BMC Bioinformatics, 10, 405-421. https://dx.doi.org/10.1186/ 1471-2105-10-405
Hirschberger M., Qi Y., Steuer R.E. (2007) Randomly generating portfolio-selection covariance matrices with specified distributional characteristics. European Journal of Operational Research. 177, 1610-1625. https://dx.doi.org/10.1016/j.ejor.2005.10.014

## See Also

```
small.world
rewire,sample_degseq,keeping_degseq
transitivity
```

Other Random graph functions: Rich Club

## Examples

```
## Not run:
rand_all <- analysis_random_graphs(g.norm, 1e2,
    savedir='/home/cwatson/dti/rand', clustering=F)
## End(Not run)
## Not run:
rand1 <- sim.rand.graph.par(g[[1]][[N]], N=1e3)
rand1.cl <- sim.rand.graph.par(g[[1]][[N]], N=1e2,
    clustering=T, max.iters=1e3)
## End(Not run)
```

randomise GLM non-parametric permutation testing

## Description

randomise and randomise_3d perform non-parametric permutation testing for analyses in which there is a single or multiple design matrix per region, respectively. In the latter case, X should be a 3D array.
partition partitions a full design matrix into separate matrices of covariates of interest and nuisance covariates based on a given contrast and partition method.

## Usage

partition(M, contrast, part.method = c("beckmann", "guttman", "ridgway"))
randomise(perm.method, part.method, $N$, perms, $X, y$, contrasts, ctype, $n C$, skip $=$ NULL, $n=\operatorname{dim}(X)[1 L], p=q r . \operatorname{default}(X) \$ r a n k$,

```
    ny \(=\operatorname{dim}(\mathrm{y})[2 \mathrm{~L}], \operatorname{dfR}=\mathrm{n}-\mathrm{p})\)
randomise_3d(perm.method, part.method, N, perms, X, y, contrasts, ctype,
    \(\mathrm{nC}, \operatorname{runX}=\operatorname{dimnames}(X)[[3 \mathrm{~L}]], \mathrm{n}=\operatorname{dim}(\mathrm{X})[1 \mathrm{~L}], \mathrm{p}=\mathrm{qr} . \operatorname{default}(\mathrm{X}[\), ,
    1L]) \$rank, ny = length(runX), dfR = \(n\) - \(p\) )
```


## Arguments

| M | Numeric matrix or array of the full design matrix(es) |
| :---: | :---: |
| contrast | For partition, a numeric matrix with 1 or more rows (for T and F contrasts, respectively) representing a single contrast. |
| part.method | Character string; the method of partitioning the design matrix into covariates of interest and nuisance. Default: 'beckmann' |
| perm.method | Character string indicating the permutation method. Default: 'freedmanLane ' |
| N | Integer; number of permutations to create. Default: 5e3 |
| perms | Matrix of permutations, if you would like to provide your own. Default: NULL |
| X | Numeric matrix or 3D array of the design matrix(es) |
| y | Numeric matrix of outcome variables, with 1 column per region, or a single column if there is a different design matrix per region |
| contrasts | Numeric matrix (for T statistics) or list of matrices (for F statistics) specifying the contrast(s) of interest; if only one contrast is desired, you can supply a vector (for T statistics) |
| ctype | The contrast type |
| nC | Integer; the number of contrasts |
| skip | Integer vector indicating which (if any) contrasts to skip. Only used by NBS. |
| $n, p, n y, d f R$ | Integers for the number of observations, design matrix columns (its rank), number of regions/outcome variables, and residual degrees of freedom, respectively |
| runX | Character vector of regions with non-singular designs |

## Value

partition returns a list containing:
Mp Numeric array; the combined partitioned arrays
$X \quad$ Numeric array for the covariates of interest
Z Numeric array for the nuisance covariates
$\mathrm{eCm} \quad$ The effective contrast, equivalent to the original, for the partitioned model $[\mathrm{X}, \mathrm{Z}]$ and considering all covariates
eCx Same as eCm, but considering only $X$
A numeric array with dimensions $n_{y} \times N \times n_{c}$; the number of rows equals number of regions/outcome variables, number of columns equals N , and the 3 rd dimension is the number of contrasts

## Model partitioning

Consider the matrix formulation of the general linear model:

$$
\mathbf{Y}=\mathbf{M} \psi+\epsilon
$$

where $Y$ is the vector of outcomes, $M$ is the full design matrix (including nuisance covariates), $\psi$ is the vector of parameter estimates, and $\in$ is the vector of error terms. In a permutation framework, algorithms are applied differently depending on the presence/absence of nuisance covariates; thus the model is separated depending on the contrast of interest:

$$
\mathbf{Y}=\mathbf{X} \beta+\mathbf{Z} \gamma+\epsilon
$$

where $\mathbf{X}$ contains covariates of interest, $\mathbf{Z}$ contains nuisance covariates, and $\beta$ and $\gamma$ are the associated parameter estimates.
The manner of partitioning depends on the method. For example, for the guttman method, X is formed from the columns of contrast that have non-zero entries.

## Permutation methods

The permutation methods can be split into 2 groups, depending on which part of the model they permute. For full details, see Winkler et al., 2014.

Permute Y Freedman-Lane, Manly, and ter Braak
Permute X Smith, Draper-Stoneman, and Still-White
Depending on the size of the data, it may be faster to use a method that permutes $Y$ instead of $X$. For example, in NBS with dense matrices (more than 400-500 edges), it will be somewhat faster to use the "Smith" method compared to "Freedman-Lane". If using brainGraph_GLM, the number of vertices follows the same relationship.
Furthermore, all methods except Still-White include the Z (nuisance covariate) matrix when calculating the permuted statistics.

## References

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Residuals Linear model residuals in structural covariance networks

## Description

get. resid runs linear models across brain regions listed in a data. table (e.g., cortical thickness), adjusting for variables in covars (e.g. age, sex, etc.), and calculates the externally Studentized (or leave-one-out) residuals.

The [ method reorders or subsets residuals based on a given numeric vector. However, this is used in bootstrap and permutation analysis and should generally not be called directly by the user.
The summary method prints the number of outliers per region, and the number of times a given subject was an outlier (i.e., across regions).

The plot method lets you check the model residuals for each brain region in a structural covariance analysis. It shows a qqplot of the studentized residuals, as output from get. resid.

## Usage

get.resid(dt.vol, covars, method = c("comb.groups", "sep.groups"), use.mean $=$ FALSE, exclude.cov $=$ NULL, atlas $=$ NULL, ...)
\#\# S3 method for class 'brainGraph_resids'
$\mathrm{x}[\mathrm{i}, \mathrm{g}=\mathrm{NULL}]$
\#\# S3 method for class 'brainGraph_resids'
summary (object, region $=$ NULL,
outlier.thresh = 2, ...)
\#\# S3 method for class 'brainGraph_resids'
plot(x, region = NULL, outlier.thresh = 2, cols = FALSE, ids = TRUE, ...)
\#\# S3 method for class 'brainGraph_resids'
nobs(object, ...)

```
## S3 method for class 'brainGraph_resids'
case.names(object, ...)
## S3 method for class 'brainGraph_resids'
groups(x)
## S3 method for class 'brainGraph_resids'
region.names(object)
## S3 method for class 'brainGraph_resids'
nregions(object)
```


## Arguments

dt.vol A data. table containing all the volumetric measure of interest (i.e., the object lhrh as output by import_scn)
covars A data.table of the covariates of interest
method Character string indicating whether to test models for subject groups separately or combined. Default: comb.groups
use.mean Logical should we control for the mean hemispheric brain value (e.g. mean LH/RH cortical thickness). Default: FALSE
exclude.cov Character vector of covariates to exclude. Default: NULL
atlas Character string indicating the brain atlas
... Arguments passed to brainGraph_GLM_design (optional)
x, object A brainGraph_resids object
i Numeric vector of the indices
g Character string indicating the group. Default: NULL
region Character vector of region(s) to focus on; default behavior is to show summary for all regions
outlier.thresh Number indicating how many standard deviations above/below the mean indicate an outlier. Default: 2
cols Logical indicating whether to color by group. Default: FALSE
ids Logical indicating whether to plot subject ID's for outliers. Otherwise plots the integer index

## Details

You can choose to run models for each of your subject groups separately or combined (the default) via the method argument. You may also choose whether to include the mean, per-hemisphere structural measure in the models. Finally, you can specify variables that are present in covars which you would like to exclude from the models. Optional arguments can be provided that get passed to brainGraph_GLM_design.
If you do not explicitly specify the atlas name, then it will be guessed from the size of your data. This could cause problems if you are using a custom atlas, with or without the same number of regions as a dataset in the package.

## Value

get.resid - an object of class brainGraph_resids with elements:
data A data.table with the input volume/thickness/etc. data as well as the covariates used in creating the design matrix.
$X \quad$ The design matrix, if using default arguments. If use.mean=TRUE then it will be a named list with a separate matrix for the left and right hemispheres. If method='sep.groups', a nested named list for each group and hemisphere.
method The input argument method
use.mean The input argument use.mean
resids.all The "wide" data.table of residuals
Group Group names
atlas The atlas name
summary.brainGraph_resids returns a list with two data tables, one of the residuals, and one of only the outlier regions
The plot method returns a trellis object or a list of ggplot objects

## Note

It is assumed that $d t . v o l$ was created using import_scn. In older versions, there were issues when the Study ID was specified as an integer and was not "zero-padded". This is done automatically by import_scn, so if you are using an external program, please be sure that the Study ID column is matched in both dt . vol and covars.

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## See Also

influence.measures, qqnorm
Other Structural covariance network functions: Bootstrapping, IndividualContributions, brainGraph_permute, corr.matrix, import_scn, plot_volumetric

## Examples

```
## Not run:
myresids <- get.resids(lhrh, covars)
residPlots <- plot(myresids, cols=TRUE)
## Save as a multi-page PDF
ml <- marrangeGrob(residPlots, nrow=3, ncol=3)
ggsave('residuals.pdf', ml)
## End(Not run)
```

Rich Club Rich club calculations

## Description

rich_club_coeff calculates the rich club of a graph, returning the rich-club coefficient, $\phi$, and the subgraph of rich club vertices.
rich_club_all is a wrapper for rich_club_coeff that calculates the rich-club coefficient for all degrees present in the graph. It returns a data. table with the coefficients and vertex and edge counts for each successive rich club.
rich_club_norm will (optionally) generate a number of random graphs, calculate their rich club coefficients $(\phi)$, and return $\phi_{\text {norm }}$ of the graph of interest, which is the observed rich-club coefficient divided by the mean across the random graphs.
rich_core finds the boundary of the rich core of a graph, based on the decreasing order of vertex degree. It also calculates the degree that corresponds to that rank, and the core size relative to the total number of vertices in the graph.

## Usage

rich_club_coeff(g, k = 1, weighted = FALSE, $A=N U L L)$
rich_club_all(g, weighted = FALSE, A = NULL)
rich_club_norm(g, N = 100, rand = NULL, ...)
rich_core(g, weighted $=$ FALSE, $A=$ NULL)

## Arguments

| g | An igraph graph object |
| :--- | :--- |
| k | Integer; the minimum degree for including a vertex. Default: 1 |
| weighted | Logical indicating whether or not edge weights should be used. Default: FALSE |
| A | Numeric matrix; the adjacency matrix of the input graph. Default: NULL |
| N | Integer; the number of random graphs to generate. Default: 100 |
| rand | A list of igraph graph objects, if random graphs have already been generated. |
|  | Default: NULL |
| $\ldots$ | Other parameters (passed to sim. rand.graph.par) |

## Details

If random graphs have already been generated, you can supply a list as an argument.
For weighted graphs, the degree is substituted by a normalized weight:

$$
\operatorname{ceiling}\left(A / w_{\min }\right)
$$

where $w_{\text {min }}$ is the minimum weight (that is greater than 0 ), and ceiling () is the ceiling function that rounds up to the nearest integer.

## Value

rich_club_coeff - a list with components:
phi The rich club coefficient, $\phi$.
graph A subgraph containing only the rich club vertices.
Nk , Ek The number of vertices/edges in the rich club graph.
rich_club_all-a data.table with components:
$k \quad$ A vector of all vertex degrees present in the original graph
phi The rich-club coefficient
Nk, Ek The number of vertices/edges in the rich club for each successive $k$
rich_club_norm - a data table with columns:

| k | Sequence of degrees |
| :---: | :---: |
| rand | Rich-club coefficients for the random graphs |
| orig | Rich-club coefficients for the original graph. |
| norm | Normalized rich-club coefficients. |
| p | P -values based on the distribution of rand |
| p.fdr | The FDR-adjusted P-values |
| density | The observed graph's density |
| threshold, Group, name |  |
| rich_core - a data table with columns: |  |
| density | The density of the graph. |
| rank | The rank of the boundary for the rich core. |
| k.r | The degree/strength of the vertex at the boundary. |
| core.size | The size of the core relative to the graph size. |
| weighted | Whether or not weights were used |

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## References

Zhou, S. and Mondragon, R.J. (2004) The rich-club phenomenon in the internet topology. IEEE Comm Lett, 8, 180-182. https://dx.doi.org/10.4018/978-1-59140-993-9.ch066
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Colizza, V. and Flammini, A. and Serrano, M.A. and Vespignani, A. (2006) Detecting rich-club ordering in complex networks. Nature Physics, 2, 110-115. https://dx.doi.org/10.1038/ nphys209
Ma, A and Mondragon, R.J. (2015) Rich-cores in networks. PLoS One, 10(3), e0119678. https: //dx.doi.org/10.1371/journal.pone. 0119678

## See Also

Other Rich-club functions: plot_rich_norm, rich_club_attrs
Other Random graph functions: Random Graphs
rich_club_attrs Assign graph attributes based on rich-club analysis

## Description

Assigns vertex- and edge-level attributes based on the results of a rich-club analysis, based on a range of vertex degrees in which the rich-club coefficient was determined to be significantly greater than that of a set of random graphs (see rich_club_norm).

## Usage

rich_club_attrs(g, deg.range = NULL, adj.vsize = FALSE)

## Arguments

$\begin{array}{ll}\mathrm{g} & \text { An igraph graph object } \\ \text { deg. range } & \begin{array}{l}\text { Numeric vector of the range of degrees indicating inclusion in the rich-club; if } \\ \text { the default } N U L L, \text { it will be from } 1 \text { to the maximum degree in the graph }\end{array} \\ \text { adj.vsize } & \begin{array}{l}\text { Logical indicating whether to adjust vertex size proportional to degree. Default: } \\ \text { FALSE }\end{array}\end{array}$

## Details

Vertices which are in the rich club will be assigned an attribute rich, taking on a binary value. Their colors (attribute color.rich) will be either red or gray. Their sizes (attribute size.rich) will either be 10 or will be proportional to their degree.
Edge attribute type.rich takes on three values: rich-club (if it connects two rich-club vertices), feeder (if it connects a rich- to a non-rich-club vertex), and local (if it connects two non-rich-club vertices). The color. rich attribute is red, orange, or green. Edge sizes (size.rich) will be largest for rich-club connections, then smaller for feeder, and smallest for local.

## Value

An igraph graph object with additional attributes:

| rich | Binary indicating membership in the rich-club |
| :--- | :--- |
| type.rich | Edge attribute indicating the type of connection |
| color.rich | Edge and vertex attributes |
| size.rich | Edge and vertex attributes |

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## See Also

Other Rich-club functions: Rich Club, plot_rich_norm

## Examples

```
## Not run:
g <- rich_club_attrs(g, rich.dt[density == densities[N] & p.fdr < .01,
    range(k)])
## End(Not run)
```

```
robustness Analysis of network robustness
```


## Description

This function performs a "targeted attack" of a graph or a "random failure" analysis, calculating the size of the largest component after edge or vertex removal.

## Usage

```
robustness(g, type = c("vertex", "edge"), measure = c("btwn.cent",
    "degree", "random"), N = 1000)
```


## Arguments

| g | An igraph graph object |
| :--- | :--- |
| type | Character string; either 'vertex ' or 'edge ' removals. Default: vertex |
| measure | Character string; sort by either 'btwn.cent' or 'degree ', or choose 'random' <br> Default: 'btwn.cent' |
| N | Integer; the number of iterations if 'random' is chosen. Default: 1 e 3 |

## Details

In a targeted attack, it will sort the vertices by either degree or betweenness centrality (or sort edges by betweenness), and successively remove the top vertices/edges. Then it calculates the size of the largest component.

In a random failure analysis, vertices/edges are removed in a random order.

Value
Data table with elements:
type $\quad$ Character string describing the type of analysis performed
measure The input argument measure
comp.size The size of the largest component after edge/vertex removal
comp.pct Numeric vector of the ratio of maximal component size after each removal to the observed graph's maximal component size
removed.pct Numeric vector of the ratio of vertices/edges removed
Group Character string indicating the subject group, if applicable

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## References

Albert, R. and Jeong, H. and Barabasi, A. (2000) Error and attack tolerance of complex networks. Nature, 406, 378-381. https://dx.doi.org/10.1038/35019019

## Description

small.world calculates the normalized characteristic path length and clustering coefficient based on observed and random graphs, used to calculate the small-world coefficient $\sigma$.

## Usage

small.world(g.list, rand)

## Arguments

g.list A brainGraphList object or list of graphs
rand List of (lists of) equivalent random graphs (output from sim.rand.graph.par)

## Value

A data.table with the following components:
density The range of density thresholds used.
$\mathrm{N} \quad$ The number of random graphs that were generated.
Lp,Lp. rand, Lp.norm
The observed, average random, and normalized characteristic path length.
Cp, Cp. rand, Cp. norm
The observed, average random, and normalized clustering coefficient.
sigma The small-world measure of the graph.

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## References

Watts, D.J. and Strogatz S.H. (1998) Collective dynamics of "small-world" networks. Nature, 393, 440-442. https://dx.doi.org/10.1038/30918
s_core $\quad$ Calculate the s-core of a network

## Description

Calculates the $s$-core decomposition of a network. This is analogous to the $k$-core decomposition, but takes into account the strength of vertices (i.e., in weighted networks). If an unweighted network is supplied, then the output of the function coreness is returned.

## Usage

s_core(g, W = NULL)

## Arguments

g
An igraph graph object
W Numeric matrix of edge weights (default: NULL)

## Details

The $s$-core consists of all vertices $i$ with $s_{i}>s$, where $s$ is some threshold value. The $s_{0}$ core is the entire network, and the threshold value of the $s_{n}$ core is

$$
s_{n-1}=\min _{i} s_{i}
$$

for all vertices $i$ in the $s_{n-1}$ core.
Note that in networks with a wide distribution of vertex strengths, in which there are almost as many unique values as there are vertices, then several separate cores will have a single vertex. See the reference provided below.

## Value

Integer vector of the vertices' $s$-core membership

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## References

Eidsaa, M and Almaas, E. (2013) s-core network decomposition: a generalization of k-core analysis to weighted networks. Physical Review E, 88, 062819. https://dx.doi.org/10.1103/ PhysRevE. 88.062819

## See Also

coreness

Vertex Roles Gateway coefficient, participation coefficient, and within-mod degree z-score

## Description

gateway_coeff calculates the gateway coefficient of each vertex, based on community membership.
part_coeff calculates the participation coefficient of each vertex, based on community membership.
within_module_deg_z_score is a measure of the connectivity from a given vertex to other vertices in its module/community.

## Usage

```
gateway_coeff(g, memb, centr = c("btwn.cent", "degree", "strength"),
    A = NULL)
    part_coeff(g, memb, A = NULL)
    within_module_deg_z_score(g, memb, A = NULL)
```


## Arguments

| g | An igraph graph object |
| :--- | :--- |
| memb | A numeric vector of membership indices of each vertex |
| centr | Character string; the type of centrality to use in calculating GC (default: btwn. cent) |
| A | Numeric matrix; the adjacency matrix of the input graph. Default: NULL |

## Details

The gateway coefficient $G_{i}$ of vertex $i$ is:

$$
G_{i}=1-\sum_{S=1}^{N_{M}}\left(\frac{\kappa_{i S}}{\kappa_{i}}\right)^{2}\left(g_{i S}\right)^{2}
$$

where $\kappa_{i S}$ is the number of edges from vertex $i$ to vertices in module $S$, and $\kappa_{i}$ is the degree of vertex $i . N_{M}$ equals the number of modules. $g_{i i}$ is a weight, defined as:

$$
g_{i S}=1-\kappa_{i S}^{-} \overline{i_{i S}^{-}}
$$

where

$$
\kappa_{i S}^{-}=\frac{\kappa_{i S}}{\sum_{j} \kappa_{j S}}
$$

for all nodes $j$ in node $i$ 's module, and

$$
c_{i S}^{-}=c_{i S} / \max \left(c_{n}\right)
$$

The participation coefficient $P_{i}$ of vertex $i$ is:

$$
P_{i}=1-\sum_{s=1}^{N_{M}}\left(\frac{\kappa_{i s}}{\kappa_{i}}\right)^{2}
$$

where $\kappa_{i s}$ is the number of edges from vertex $i$ to vertices in module $s$, and $\kappa_{s}$ is the degree of vertex i. $N_{M}$ equals the number of modules.

As discussed in Guimera et al., $P_{i}=0$ if vertex $i$ is connected only to vertices in the same module, and $P_{i}=1$ if vertex $i$ is equally connected to all other modules.
The within-module degree $z$-score is:

$$
z_{i}=\frac{\kappa_{i}-\bar{\kappa}_{s_{i}}}{\sigma_{\kappa_{s_{i}}}}
$$

where $\kappa_{i}$ is the number of edges from vertex $i$ to vertices in the same module $s_{i}, \bar{\kappa}_{s_{i}}$ is the average of $\kappa$ over all vertices in $s_{i}$, and $\sigma_{\kappa_{s_{i}}}$ is the standard deviation.

## Value

A vector of the participation coefficients, within-module degree z-scores, or gateway coefficients for each vertex of the graph.

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## References

Vargas, E.R. and Wahl, L.M. (2014) The gateway coefficient: a novel metric for identifying critical connections in modular networks. Eur Phys J B, 87, 161-170. https://dx.doi.org/10.1140/ epjb/e2014-40800-7
Guimera, R. and Amaral, L.A.N. (2005) Cartography of complex networks: modules and universal roles. Journal of Statistical Mechanics: Theory and Experiment, 02, P02001. https://dx.doi. org/10.1088/1742-5468/2005/02/P02001

## Description

Variance inflation factors for bg_GLM objects

## Usage

vif.bg_GLM(mod, ...)

## Arguments

mod A bg_GLM object
... Unused

## Value

A named array of VIFs; names of the 3rd dimension are regions

```
vulnerability Calculate graph vulnerability
```


## Description

This function calculates the vulnerability of the vertices of a graph. Here, vulnerability is considered to be the proportional drop in global efficiency when a given vertex is removed from the graph. The vulnerability of the graph is considered the maximum across all vertices.

## Usage

vulnerability (g, use.parallel $=$ TRUE, weighted $=$ FALSE)

## Arguments

$\begin{array}{ll}\mathrm{g} & \text { An igraph graph object } \\ \text { use.parallel } & \text { Logical indicating whether or not to use foreach (default: TRUE) } \\ \text { weighted } & \text { Logical indicating whether weighted efficiency should be calculated (default: } \\ & \text { FALSE) }\end{array}$

## Value

A numeric vector of length equal to the vertex count of $g$

## Author(s)

Christopher G. Watson, <cgwatson@bu. edu>

## References

Latora, V. and Marchiori, M. (2005) Variability and protection of infrastructure networks. Physical Review E, 71, 015103. https://dx.doi.org/10.1103/physreve.71.015103

## See Also

efficiency

```
write_brainnet Write fles to be used for visualization with BrainNet Viewer
```


## Description

Write the . node and . edge files necessary for visualization with the BrainNet Viewer software.

## Usage

write_brainnet(g, vcolor = "none", vsize = "constant", edge.wt $=$ NULL, file.prefix $=$ "")

## Arguments

g The igraph graph object of interest
vcolor Character string indicating how to color the vertices (default: 'none')
vsize $\quad$ Character string indicating what size the vertices should be; can be any vertexlevel attribute (default: 'constant')
edge.wt Character string indicating the edge attribute to use to return a weighted adjacency matrix (default: NULL)
file.prefix Character string for the basename of the .node and .edge files that are written

## Details

For the .node file, there are 6 columns:

- Columns 1-3: Vertex x-, y-, and z-coordinates
- Column 4: Vertex color
- Column 5: Vertex size
- Column 6: Vertex label

The .edge file is the graph's associated adjacency matrix; a weighted adjacency matrix can be returned by using the edge.wt argument.

## Author(s)

Christopher G. Watson, [cgwatson@bu.edu](mailto:cgwatson@bu.edu)

## References

Xia, M. and Wang, J. and He, Y. (2013). BrainNet Viewer: a network visualization tool for human brain connectomics. PLoS One, 8(7), e68910. https://dx.doi.org/10.1371/journal.pone. 0068910

## Examples

```
## Not run:
write_brainnet(g, vcolor='community', vsize='degree', edge.wt='t.stat')
## End(Not run)
```


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