Package 'LESYMAP'

June 12, 2017

Title Leions to Symptom Mapping in R

Date 2017-06-07 Description LESYMAP maps the specific brain areas responsible for cognitive deficits by taking a series of lesion maps and a vector of behavioral scores. Both univariate (t-test, Brunner-Munzel,regression) and multivariate (sparse canonical corelations) tests are available. LESYMAP is built to run both real and simulated lesion-to-symptom mapping analyses. License Apache License 2.0 Encoding UTF-8 LazyData true Depends R (>= 3.0),ANTsR Imports ANTsRCore,graphics,ImPerm,nparcomp,Rcpp,stats,utils Remotes stnava/ANTsR,stnava/ANTsRCore LinkingTo Rcpp, RcppArmadillo NeedsCompilation yes RoxygenNote 6.0.1.9000 R topics documented: .createFolds BM BMfast BMfast BMfast BMfast2 checkAntsInput checkFilenameHeaders checkImageList checkMask getLesionLoad	Version 0.0.0.9003			
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getLesionSize	createFolds BM BMfast BMfast2 checkAntsInput checkFilenameHeaders checkImageList checkMask			

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getUniqueLesionPatches
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lsm_BMfast
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lsm_regres
lsm_regresfast
lsm_regresPerm
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optimize_SCCANsparseness
print.lesymap
regresfast
save.lesymap
simulateBehavior

.createFolds

createFolds

Description

Used to create folds for k-fold validation

Usage

```
.createFolds(y, k = 10, list = TRUE, returnTrain = FALSE)
```

Arguments

y split sample by balancing y

k number of folds

list logical whether to return folds in a list

 $\verb"returnTrain" logical whether to return training indices (T) or the test samples (F)$

Author(s)

Caret Package

BM 3

ВМ

Massive Brunner-Munzel tests

Description

Takes a binary matrix of voxels and a vector of behavior and runs Brunner-Munzel tests on each voxel. This function is not compiled and is slow.

Usage

```
BM(lesmat, behavior)
```

Arguments

lesmat matrix of voxels
behavior vector of behavior

Value

Returned list with:

- statistic statistical values
- dof degrees of freedom

Author(s)

Dorian Pustina

Examples

```
set.seed(123)
lesmat = matrix(rbinom(200,1,0.5), ncol=2)
set.seed(123)
behavior = rnorm(100)
result = BM(lesmat, behavior)
```

BMfast

Fast Brunner-Munzel tests (v1)

Description

Takes a binary matrix of voxels and a vector of behavior and runs Brunner-Munzel tests on each voxel. This is a fast function, but may produce infinite values for perfectly separated group. Use BMfast2 which avoids this problem.

```
BMfast(X, y)
```

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Arguments

X binary matrix ov voxlels (columns) for all subjects (rows)

y vector of behavioral scores.

Value

List with two vectors: - statistic - BM values - dfbm - degrees of freedom

Author(s)

Dorian Pustina

BMfast2

Fast Brunner-Munzel tests (v2)

Description

Takes a binary matrix of voxels and a vector of behavior and runs Brunner-Munzel tests on each voxel. This is a fast function that corrects for infinite values with a similar approach as the nparcomp package.

Usage

```
BMfast2(X, y, computeDOF = TRUE)
```

Arguments

X binary matrix ov voxlels (columns) for all subjects (rows)

y vector of behavioral scores.

computeDOF (true) chooses whether to compute degrees of freedom. Set to false to save time

during permutations.

Value

List with two vectors: - statistic - BM values - dfbm - degrees of freedom

Author(s)

checkAntsInput 5

checkAntsInput checkAntsInput

Description

Function to check a variable whether is composed of an antsImage, list of antsImages, or simply filenames. If none of the above, an error is returned.

Usage

```
checkAntsInput(input, checkHeaders = F)
```

Arguments

```
input the variable to be checked checkHeaders make sure all images have the same headers
```

Value

Type of variable (antsImage, antsImageList, antsFiles) or error if variable cannot be established.

Author(s)

Dorian Pustina

Examples

```
## Not run:
    files = Sys.glob('/data/jag/nifti/*.nii.gz')
    myimagelist = imageFileNames2ImageList(files)
    checkAntsInput(myimagelist) # returns 'antsImageList'
    checkAntsInput(antsFiles) # returns 'antsFiles'
    checkAntsInput(myimagelist[[1]]) # returns 'antsImage'
## End(Not run)
```

checkFilenameHeaders

checkFilenameHeaders

Description

Function to check that all filenames in a vector point to existing files with the same resolution, orientation, size, and origin.

```
checkFilenameHeaders(files, showError = T)
```

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Arguments

files character vector of filenames

showError logical whether to show an error (True) or to return a boolean instead. Returned

values are True=pass,False=Fail

Value

logical if the test was successful or not

Author(s)

Dorian Pustina

checkImageList checkImageList

Description

Function to check that all antsImages in a list have the same orientation, origin, and resolution. The function stops with an error if one of the images has unusual headers. This behavior can be overcome by setting showError=F, and using the returned status (True=pass, False=fail) to make decisions outside this function.

Usage

```
checkImageList(imgList, showError = T, binaryCheck = F)
```

Arguments

imgList list of antsImages

showError boolean indicating whether to show the exact error and interrupt the function

(TRUE, default), or don't show the error and return the check status (FALSE).

The returned values when showError=F are T=passed or F=Failed.

binaryCheck boolean, check if images are binary (0/1 values). Useful when checking masks

or lesions. This check slows the output of the function.

Value

True if list has images with same headers, otherwise False.

Author(s)

checkMask 7

Examples

```
## Not run:
files = Sys.glob('/data/jag/nifti/*.nii.gz')
myimagelist = imageFileNames2ImageList(files)
checkImageList(myimagelist) # no value returned
checkImageList(lesions, showError=F) # True returned
myimagelist[[4]] = cropIndices(myimagelist[[4]], c(1,1,1), c(20,20,20))
checkImageList(myimagelist) # error on image 4
## End(Not run)
```

checkMask

checkMask

Description

Function to check if mask is in the same space as inputs

Usage

```
checkMask(lesions.list, mask)
```

Arguments

```
lesions.list list of antsImages or character vector of filenames
mask antsImage of mask to check
```

Value

Nothing is returned, function stops with error if mask is not in the same space as images in lesions.list

Author(s)

Dorian Pustina

getLesionLoad

getLesionLoad

Description

Computes lesion loads from a series of images. A parcellation image (or simple mask) is required to define the regions from which to compute the lesion load.

```
getLesionLoad(lesions.list, parcellation, label = NA, mask = NA,
binaryCheck = F, keepAllLabels = F, minSubjectPerLabel = "10%")
```

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Arguments

lesions.list list of antsImages or filenames. Must be binary (0 and 1 values). parcellation ansImage or filename of the parcellated volumes. A parcellation is an image brain regions showned as with integer values (i.e. ,1,2,3,...). label (default=NA) you can ask to get output for a specific label in the parcellation volume (i.e., label=122). (default=NA) if this mask is specified (antsImage or filename) lesioned voxels mask outside the mask are ignored. This is not a good choice, but in case you need it its there. (default=FALSE) check whether lesion maps are binary (0/1). Will output an binaryCheck error if lesion files are not binary. keepAllLabels (default=FALSE) by default labels are removed if affected in just few subjects. Setting this to TRUE will keep all labels. minSubjectPerLabel minimum number of subjects a parcel must be lesioned to keep and return it.

Value

• outputMatrix of lesion loads between 0 and 1. 1 means 100% lesioned. Each column is a single parcel and each row a single subject. Parcel numbers are placed as column names.

Author(s)

Dorian Pustina

Examples

```
lesydata = file.path(find.package('LESYMAP'),'extdata')
filenames = Sys.glob(file.path(lesydata, 'lesions', '*.nii.gz'))
lesions = imageFileNames2ImageList(filenames[1:10])
parcellation = antsImageRead(
file.path(lesydata,'template', 'Parcellation_403areas.nii.gz'))
lesload = getLesionLoad(lesions, parcellation)
```

getLesionSize

getLesionSize

Description

Compute lesion sizes from a list of antsImages.

Usage

```
getLesionSize(lesions.list)
```

Arguments

lesions.list List of antsImages. For proper measurement, images must be binary.

Value

vector of lesion sizes

Author(s)

Dorian Pustina

```
getUniqueLesionPatches
```

Unique Lesion Patches

Description

Compute unique patches of voxels with the same pattern of lesions in all subjects. Useful to understand the number of patterns that will be analyzed in a lesion dataset. A patch is a group of voxels, not necessarily close to each other, which have the same identical lesion pattern.

Usage

```
getUniqueLesionPatches(lesions.list, mask = NA, returnPatchMatrix = F,
    thresholdPercent = 0.1, binaryCheck = F, showInfo = T)
```

Arguments

lesions.list list of antsImages (faster) or filenames (slower)

mask

(default=NA) a mask image to restrict the search for patches. Will be automatically calculated if not provided. Normally the mask restricts the search only to voxels lesioned in >10% of subejets. To set this proportion use thresholdPercent.

returnPatchMatrix

(default=FALSE) logical, should the matrix of patches be returned. This is used in lesymap to run the analyses.

thresholdPercent

(default=0.1) voxels with lesions in less than this proportion of subjects will not be considered. I.e., 0.1 = 10%.

binaryCheck (default=FALSE) set this to TRUE to verify that maps are binary.

showInfo (default=TRUE) logical indicating whether to display information.

Value

List of objects named as follows:

- patchimg antsImage with every voxel assigned a patch number
- patchimg.samples antsImage mask of one representative voxel for each patch. Can be used to extract the patchmatrix.
- patchimg.size antsImage with the patch size at every voxel
- patchimg.mask antsImage of full mask with all voxels. Can be used to put back results in combination with patchindx.
- patchindx vector of patch membership for each voxel. Can be used to put back results in an image.

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- npatches number of unique patches in the image
- nvoxels total number of lesioned voxels in patchimg.mask
- patchvoxels vector of voxel count for each patch
- patchvolumes vector of volume size for each patch
- patchmatrix matrix of patches. This is used in lesymap to save time when running repetitive analyses.

Author(s)

Dorian Pustina

Examples

```
lesydata = file.path(find.package('LESYMAP'), 'extdata')
filenames = Sys.glob(file.path(lesydata, 'lesions', '*.nii.gz'))
patchinfo = getUniqueLesionPatches(filenames[1:10]) # slower
lesions = imageFileNames2ImageList(filenames[1:10])
patchinfo = getUniqueLesionPatches(lesions) # faster
```

```
lesyload_mricron lesyload_mricron
```

Description

Function to load data from a previous analysis in MRIcron/npm in a ready format for use in lesymap

Usage

```
lesyload_mricron(valfile, imageFolder = NA, returnFilenames = F,
   checkHeaders = T, showInfo = T)
```

Arguments

valfile

mricron filename with extention *.val. The function will search for images in the same folder where valfile is located, unless you specify imageFolder. If any of the files listed in the .val file are not found in the folder, an error will be displayed.

 $\verb|imageFolder| (default=NA) folder to look for the image files$

returnFilenames

(default=FALSE) By default the function will load the images in memory to speed up things in lesymap. This may require too much RAM memory in some cases, and you may want to use filenames instead, which requires less memory but is slower in lesymap.

checkHeaders (default=TRUE) Headers will be checked to make sure all images have the same dimension/origin/resolution, etc.

showInfo (default=TRUE) show information upon successful load

Value

List with the following information lesions - list of antsImages or vector of filenames behavior - vector of behavioral scores

Author(s)

Dorian Pustina

lesymap

Lesion to Symptom Mapping

Description

Lesymap uses univariate and multivariate methods to map functional regions of the brain that, when lesioned, cause specific cognitive deficits. All is required is a set of Nifti images with the lesion of each subject and the vector of behavioral scores. Lesions must be already registered in template space, use 'antsRegistration' or other ANTs tools to achieve this. Lesymap will check that lesions are in the same space before running. By default, voxels with identical lesion patterns are grouped together in unique patches, and analysis are run on patches. Patch-based mapping decreases the number of multiple comparisons and speeds up the analyses. Multivariate mapping is performed using an optimized version of sparse canonical correlations (SCCAN).

Usage

```
lesymap(lesions.list, behavior, mask = NA, patchinfo = NA, method = "BM",
   correctByLesSize = "none", multipleComparison = "fdr",
   pThreshold = 0.05, flipSign = F, minSubjectPerVoxel = "10%",
   nperm = 1000, saveDir = NA, binaryCheck = F, noPatch = F,
   showInfo = T, ...)
```

Arguments

lesions.list	list of antsImages, or a ve	ector of filenames,	or a single antsImage	with 4 dimen-
	cione			

behavior vector of behavioral scores or filename pointing to a file with a single column of

numbers.

mask (default=NA) binary image to select the area where analysis will be performed.

If not provided will be computed automatically by thresholding the average le-

sion map at minSubjectPerVoxel.

patchinfo (default=NA) an object obtained with getUniqueLesionPatches or from a previ-

ous analyses. Useful for repetitive analysis to save time and avoid the computa-

tion of patches each time.

method what tests to run, one of 'BM' (default), 'BMfast', 'ttest', 'welch', 'regres',

'regresfast', 'regresPerm', 'sccan', 'sccanRaw'.

BM - Brunner-Munzel non parametric test, also called the Generalized Wilcoxon Test. The BM test is the same test used in the npm/Mricron software. See (see

Rorden (2007)).

BMfast - ultrafast Brunner-Munzel with compiled code. BMfast can be combined with multipleComparison='FWERperm' to perform permutation based thresholding in a short time.

ttest - Regular single tailed t-test. Variances of groups are assumed to be equal. This is the test used in the voxbo software. Relies on t.test function in R. It is assumed that 0 voxels are healthy, i.e., higher behavioral scores. See the alternative parameter for inverted cases. (see Bates (2003)).

welch - t-test that assumes unequal variance between groups. Relies on t.test function in R.

regres - linear model between voxel values and behavior. Uses the lm function in R. This is equivalent to a t-test, but is useful when voxel values are continuous. To model the effect of covariates use the "regresfast" method

regresfast - ultrafast linear regressions with compiled code. This method allows setting covariates. If covariates are specified the effect of each voxel will be estimated with the formula:

```
behavior ~ voxel + covar1 + covar2 + ...
```

This method allows multiple comparison correction with permutation based methods "FWERperm" and "clusterPerm". If these corrections are required and covariates are specified, the effect of each voxel is established with the Freedman-Lane method (see Winkler (2014)).

regresPerm - linear model between voxel values and behavior. The p-value of each individual voxel us established by permuting voxel values. The lmPerm package is used for this purpose. Note, these permutations do not correct for multiple comparisons, they only establish voxel-wise p-values.

chisq-chi-square test between voxel values and behavior. The method is used when behavioral scores are binary (i.e. presence of absence of deficit). Relies on the chisq.test R function. By default this method corrects individual voxel p-values with the Yates method (the same approach offered in the Voxbo software).

chisqPerm - chi-square tests. P-values are established through permutation tests instead of regular statistics. Relies on the chisq.test R function.

sccan - sparse canonical correlations (NEW). Multivariate method that considers all voxels at once. By default, lesymap will run a lengthy procedure to determine the optimal sparseness value (how extensive the results should be). You can set optimizeSparseness=FALSE if you want to skip this optimization. The search for optimal sparsness provides a cross-validated correlation measure that shows how well the sparseness value can predict new patients. If this predicive correlation is below significance (i.e., below pThreshold), the entire solution will be ignored and a NULL result will be returned. Lesymap returns normalized (0-1) voxel weights converted to positive; you can use rawStat=TRUE to retain the original voxel weights. Note that both lesion and behavior data are scale and centered before running SCCAN (hardcoded in lsm_sccan). You must apply the same scaling if you were to predict behavioral scores with the obtained voxel weights.

correctByLesSize

whether to correct for lesion size in the analysis. Options are "none", "voxel", "behavior":

- "none": (default) no correction
- "voxel": divide voxel values by 1/sqrt(lesionsize). This is the method used in Mirman (2015) and Zhang (2014). This correction works only with 'regres' methods. Two sample comparisons (t-tests and Brunner-Munzel) use binary voxels and will ignore this correction.
- "behavior": residualize behavioral scores by removing the effect of lesion size. This works on all methods, but is more agressive on results.

multipleComparison

(default='fdr') method to adjust p-values. Standard methods include "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr". (see p.adjust)

Permutation methods include:

"FWERperm" (permutation based family-wise threshold) is enabled with methods 'BMfast' and 'regresfast'. In this case, many analysis are run with permuted behavioral scores, and the peak score is recorded each time (see Winkler 2014). The optimal threshold is established at 95th percentile of this distribution (or whatever pThreshold you choose). You can choose to use as reference another voxel lower in the ranks by specifying another 'v' value (i.e., lesymap(..., v=10) will record the 10th highest voxel).

"clusterPerm" (permutation based cluster correction) is enabled for 'regresfast'. It records the maximal cluster size from many random permutations of the behavior score and sets a cluster threshold based on that distribution. You must select pThreshold (voxel-wise, default=0.05) and clusterPermThreshold (cluster-wise, default 0.05) to achieve optimal with this method.

pThreshold (default=0.05) threshold statistics at this p-value (after corrections or permuta-

tions)

flipSign logical (default=FALSE), invert the sign in the statistics image.

minSubjectPerVoxel

(default='10%') remove voxels/patches with lesions in less than X subjects. Value can be speficifed as percentage ('10%') or exact number of subjects (10).

nperm (default=1000) number of permutations to run when necessary.

saveDir (default=NA) save results in the specified folder.

binaryCheck logical (default=FALSE), make sure the lesion matrix is 0/1.

noPatch logical (default=FALSE), if True avoids using patch information and will ana-

lyze all voxels. It will take longer and results will be worse due to more multiple comparison corrections. This argument is ignored when performing SCCAN

analyses.

showInfo logical (default=TRUE), display time-stamped info messages

arguments that will be passed down to other functions (i.e., sparsness=0.045)

Details

Several other parameters can be specified to lesymap() which will be passed to other called fuctions. Here are some examples:

permuteNthreshold - (default=9) for Brunner-Munzel tests only. Voxels lesioned in less than this number of subjects will undergo permutation-based p-value estimation. Useful because the BM test is not valid when comparing groups with N < 9. Note, permuted BM tests currently require the package 'nparcomp'.

clusterPermThreshold - threshold used to find the optimal cluster size when using 'cluster-Perm' multiple comparison correction.

alternative - (default='greater') for two sample tests (ttests and BM). By default LESYMAP computes single tailed p-values assuming that non-lesioned 0 voxels have higher behavioral scores. You can specify the opposite relationship with alternative='less' or compute two tailed p-values with alternative='two.sided'.

covariates - (default=NA) enabled for method = 'regresfast'. This will allow to model the effect of each voxel in the context of other covariates, i.e., formula "behavior ~ voxel + covar1 + covar2 +

...".

I,.e., lesymap(lesions,behavior, method='regresfast', covariates=cbind(lesionsize, age)). If you choose permutation based thresholding with covariates, lesymap will use the Freedman-Lane method for extracting the unique effect of each voxel (see Winkler 2014, Freedman 1983)

template - antsImage or filename used for plotting the results if a saving directory is specified (see saveDir)

v - (default=1) which voxel to record for permutation based thresholding. Normally the peak voxel is used (1), but other voxels can be recorded. See Mirman 2017 for this approach.

Value

The following objects are typically found in the returned list:

- stat.img statistical map
- pval.img p-values map
- zmap.img zscore map
- mask.img mask used for the analyses
- average.img map of all lesions averaged. Map is produced only if no mask is defined.
- callinfo list of details of how you called lesymap
- perm.vector the values obtained from each permutation
- perm.clusterThreshold threshold computed for cluster thresholding
- perm.FWERthresh threshold computed for FWERperm thresholding
- patchinfo list of variables describing patch information:
 - patchimg antsImage with the patch number each voxels belongs to
 - patchimq.samples antsImage mask with a single voxel per patch
 - patchimg.size antsImage with the patch size at each voxel
 - patchimg.mask the mask within which the function will look for patches
 - npatches number of unique patches in the image
 - nvoxels total number of lesioned voxels in mask
 - patchvoxels vector of voxel count for each patch
 - patchvolumes vector of volume size for each patch
 - patchmatrix the lesional matrix, ready for use in analyses. Matrix has size NxP (N=number of subjects, P=number of patches)

Author(s)

Dorian Pustina

Examples

```
lesydata = file.path(find.package('LESYMAP'),'extdata')
filenames = Sys.glob(file.path(lesydata, 'lesions', 'Subject*.nii.gz'))
behavior = Sys.glob(file.path(lesydata, 'behavior', 'behavior.txt'))
template = antsImageRead(
   Sys.glob(file.path(lesydata, 'template', 'ch2.nii.gz')))
lsm = lesymap(filenames, behavior, method = 'BMfast')
plot(template, lsm$stat.img, window.overlay = range(lsm$stat.img))
## Not run:
```

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```
# Same analysis with SCCAN
lsm = lesymap(filenames, behavior, method = 'sccan',
sparseness=0.045, optimizeSparseness=FALSE)
plot(template, lsm$stat.img, window.overlay = range(lsm$stat.img))
save.lesymap(lsm, saveDir='/home/dp/Desktop/SCCANresults')
## End(Not run)
```

 lsm_BM

lsm_BM

Description

Lesion to symptom mapping performed on a prepared matrix. Brunner-Munzel tests are performed using each column of the matrix to split the behavioral scores in two groups.

Usage

```
lsm_BM(lesmat, behavior, permuteNthreshold = 9, nperm = 10000,
alternative = "greater", showInfo = TRUE, ...)
```

Arguments

lesmat binary matrix (0/1) of voxels (columns) and subjects (rows). behavior vector of behavioral scores. permuteNthreshold (default=9) Voxels lesioned in less than this number will undergo permutation based thresholding. See Medina et al 2010. nperm Number of permutations to perform when needed. (default="greater") It is assumed that healthy voxels (0) have greater behavioral alternative scores. If your data follow an inverted relationship choose "less" or "two.sided". display info messagges when running the function. showInfo other arguments received from lesymap. . . .

Value

List of objects returned:

- statistic vector of statistical values
- pvalue vector of pvalues
- zscore vector of zscores

Author(s)

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Examples

```
set.seed(123)
lesmat = matrix(rbinom(200,1,0.5), ncol=2)
set.seed(123)
behavior = rnorm(100)
result = lsm_BM(lesmat, behavior)
```

lsm_BMfast

lsm_BMfast

Description

Lesion to symptom mapping performed on a prepared matrix. Brunner-Munzel tests are performed using each column of the matrix to split the behavioral scores in two groups. This function relies on a compiled version for fast processing.

Usage

```
lsm_BMfast(lesmat, behavior, permuteNthreshold = 9, alternative = "greater",
  statOnly = F, nperm = 1000, FWERperm = F, v = 1, pThreshold = 0.05,
  showInfo = F, ...)
```

Arguments

. . .

binary matrix (0/1) of voxels (columns) and subjects (rows). lesmat vector of behavioral scores. behavior permuteNthreshold (default=9) Voxels lesioned in less than this number will undergo permutation based thresholding. See Medina et al 2010. (default="greater") It is assumed that healthy voxels (0) have greater behavioral alternative scores. If your data follow an inverted relationship choose "less" or "two.sided". statOnly logical (default=FALSE), skips some computations, don't use unless you know it's effects (default=1000) Number of permutations to perform when needed. nperm logical (default=FALSE) whether to perform permutation based FWER thresh-**FWERperm** olding. (default=1) which voxel to record at each permutation, first or other voxels (i.e., v=10 for 10 highest voxel) pThreshold (default=0.05) what threshold to use for FWER display info messagges when running the function. showInfo other arguments received from lesymap.

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Value

List of objects returned:

- statistic vector of statistical values
- pvalue vector of pvalues
- zscore vector of zscores
- perm. vector (optional) vector of permuted statistics
- perm.FWERthresh (optional) permutation threshold established from the distribution of perm.vector

Author(s)

Dorian Pustina

Note on zscores quorm gives same values as MRIcron and relies on the normal distribution. however, we are computing t-scores, and should have relied on that distribution, which is the t-score itself.

Examples

```
{
set.seed(123)
lesmat = matrix(rbinom(200,1,0.5), ncol=2)
set.seed(123)
behavior = rnorm(100)
result = lsm_BMfast(lesmat, behavior)
}
```

lsm_chisq

lsm_chisq

Description

Lesion to symptom mapping performed on a prepared matrix. The behavior must be a binary vector. Chi square tests are performed at each voxel. By default the Yates correction is performed, use correct=FALSE if you need to disable it. The behavior must be a binary vector. Exact p-values can be obtained with permutation based estimatins.

Usage

```
lsm_chisq(lesmat, behavior, YatesCorrect = TRUE, runPermutations = F,
   nperm = 2000, showInfo = TRUE, ...)
```

Arguments

```
lesmat binary matrix (0/1) of voxels (columns) and subjects (rows).
behavior vector of behavioral scores (must be binary.

YatesCorrect (default=T) logical whether to use Yates correction.
```

runPermutations

logical (default=FALSE) whether to use permutation based p-value estimation.

lsm_regres

```
nperm (default=2000) The number of permutations to run.
showInfo display info messagges when running the function.
... other arguments received from lesymap.
```

Value

List of objects returned:

- statistic vector of statistical values
- pvalue vector of pvalues
- zscore vector of zscores

Author(s)

Dorian Pustina

Examples

```
{
set.seed(123)
lesmat = matrix(rbinom(200,1,0.5), ncol=2)
set.seed(1234)
behavior = rbinom(100,1,0.5)
result = lsm_chisq(lesmat, behavior)
}
```

lsm_regres

lsm_regres

Description

Lesion to symptom mapping performed on a prepared matrix. Regressions are performed between behavior and each column in the lesmat matrix.

Usage

```
lsm_regres(lesmat, behavior)
```

Arguments

lesmat matrix of voxels (columns) and subjects (raws).
behavior vector of behavioral scores.

Value

List of objects returned:

- statistic vector of statistical values
- pvalue vector of pvalues
- zscore vector of zscores

lsm_regresfast 19

Author(s)

Dorian Pustina

Examples

```
{
set.seed(123)
lesmat = matrix(rbinom(200,1,0.5), ncol=2)
set.seed(123)
behavior = rnorm(100)
result = lsm_regres(lesmat, behavior)
}
```

lsm_regresfast

lsm_regresfast

Description

Lesion to symptom mapping performed on a prepared matrix. Regressions are performed between behavior and each column of the lesmat matrix. Fast function based on compiled code.

Usage

```
lsm_regresfast(lesmat, behavior, covariates = NA, FWERperm = F,
   nperm = 1000, v = 1, pThreshold = 0.05, clusterPerm = F, mask = NA,
   voxindx = NA, samplemask = NA, clusterPermThreshold = 0.05,
   showInfo = T, ...)
```

Arguments

lesmat	matrix of voxels (columns) and subjects (rows).	
behavior	vector of behavioral scores.	
covariates	(default=NA) vector of matrix of covariates.	
FWERperm	logical (default=FALSE) whether to run permutation based FWER thresholding.	
nperm	Number of permutations to perform when needed.	
V	(default=1) what voxel to record for FWER thresholding.	
pThreshold	(default=0.05) Voxel-wise threshold.	
clusterPerm	logical (default=FALSE), whether to perform permutation based cluster thresholding.	
mask	(default=NA) antsImage reference mask used for cluster computations.	
voxindx	(default=NA) indices of voxels to put in mask	
samplemask	(default=NA) antsImage used to extract voxels back in a matrix.	
clusterPermThreshold		
	(default=0.05) threshold for cluster selection after obtaining cluster size distrubution.	
showInfo	display info messagges when running the function.	
	other arguments received from lesymap.	

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Value

List of objects returned:

- statistic vector of statistical values
- pvalue vector of pvalues
- zscore vector of zscores
- perm.vector (optional) vector of permuted statistics
- perm.FWERthresh (optional) permutation threshold established from the distribution of perm.vector
- perm.clusterThreshold (optional) permutation threshold established from the distribution of perm.vector

Author(s)

Dorian Pustina

Examples

```
{
set.seed(123)
lesmat = matrix(rbinom(200,1,0.5), ncol=2)
set.seed(123)
behavior = rnorm(100)
result = lsm_regresfast(lesmat, behavior)
}
```

lsm_regresPerm

lsm_regresPerm

Description

Lesion to symptom mapping performed on a prepared matrix. Regressions are performed between behavior and each column in the lesmat matrix. This function relies on the lmPerm package to run. The number of permutations required to reach stable p-values is established automatically. For this reason, the user cannot specify a predefined number of permutations.

Usage

```
lsm_regresPerm(lesmat, behavior)
```

Arguments

lesmat matrix of voxels (columns) and subjects (rows).
behavior vector of behavioral scores.

Value

List with vectors of statistic, pvalue, and zscore.

Author(s)

21 lsm_sccan

lsm_sccan	Sparse canonical correlations for symptom	mapping.

Description

Multivariate SCCAN adapted for lesion to symptom mapping purposes. By default an optimization routine is used to find the best sparseness value. If you specify sparseness manually, it will be validated to find the cross-validated correlation that can be obtained with that sparseness. You can skip the entire optimization/validation by choosing optimizeSparseness=FALSE. To understand SCCAN arguments, see sparseDecom2.

Usage

```
lsm_sccan(lesmat, behavior, mask, rawStat = F, showInfo = T,
  optimizeSparseness = T, tstamp = "%H:%M:%S", pThreshold = 0.05,
 mycoption = 1, robust = 1, sparseness = 0.045, nvecs = 1,
  cthresh = 150, its = 20, npermsSCCAN = 0, smooth = 0.4, ...)
```

Arguments

lesmat matrix of voxels (columns) and subjects (rows). behavior vector of behavioral scores. mask antsImage binary mask to put back voxels in image. logical (default=FALSE) whether to skip the normalization of values and threshrawStat olding at 0.1. If TRUE, the raw voxel weights will be returned. showInfo logical (default-TRUE) display messages optimizeSparseness logical (default=TRUE) whether to run the sparseness optimization routine. If false, the defau sparseness value will be used. If sparseness is manually set this flag decides if the manual sparseness will be checked with cross validations. timestamp format used in LESYMAP tstamp (default=0.05) If cross validated correlations show significance below this value pThreshold the results are considered null and an empty map is returned. mycoption (default=1) SCCAN parameter, see sparseDecom2

(ddefault=1) SCCAN parameter, see sparseDecom2 robust (default=1) SCCAN parameter, see sparseDecom2 sparseness (default=1) SCCAN parameter, see sparseDecom2 nvecs (default=150) SCCAN parameter, see sparseDecom2 cthresh (default=20) SCCAN parameter, see sparseDecom2 its npermsSCCAN (default=0) SCCAN permutations, see sparseDecom2 (default=0.4) SCCAN parameter, see sparseDecom2 smooth other arguments received from lesymap. . . .

22 lsm_ttest

Value

List of objects returned:

- statistic vector of statistical values
- pvalue vector of pvalues
- optimalSparseness (optional) optimal value found for sparseness
- CVcorrelation.stat (optional) Correlation between true and predicted score with k-fold validation using the optimal sparseness value
- CVcorrelation.pval (optional) p-value of the above correlation

Author(s)

Dorian Pustina

Examples

lsm_ttest

lsm_ttest

Description

Lesion to symptom mapping performed on a prepared matrix. T-tests are performed using each column of the matrix to split the behavioral scores in two groups. If var.equal=TRUE the Welch test is performed instead.

```
lsm_ttest(lesmat, behavior, var.equal = T, alternative = "greater", ...)
```

minSegDistance 23

Arguments

lesmat	binary matrix (0/1) of voxels (columns) and subjects (rows).
behavior	vector of behavioral scores.
var.equal	logical (default=TRUE) should the variance between groups considered equal (t-test) or unequal (Welch test).
alternative	(default='greater') Sets the expected relationship between voxel value and behavior. By default voxels with zero are not lesioned, and behavior is expected to be higher, thus alternative='greater'. If the relationship in your data is inverted, use alternative='less', and if you don't know your data, use alternative='two.sided'.
• • •	other arguments received from lesymap.

Value

List of objects returned:

- statistic vector of statistical values
- pvalue vector of pvalues
- zscore vector of zscores

Author(s)

Dorian Pustina

Examples

```
{
set.seed(123)
lesmat = matrix(rbinom(200,1,0.5), ncol=2)
set.seed(123)
behavior = rnorm(100)
result = lsm_ttest(lesmat, behavior)
}
```

minSegDistance

min Seg Distance

Description

This function computes the metric displacement between two binary masks

```
minSegDistance(manual, predict, get = "all", binarize = F, label = 1)
```

Arguments

manual manual segmentation of class antsImage, used as reference

predict other antsImage to compare to manual

get (default='all') one of 'mean', 'max', 'min', or 'all'

binarize logical (default=FALSE) whether to binarize the input images

label (default=1) integer or vector of labels to binarize I.e., label=c(2,4) means label

2 from manual, and 4 from predict will be compared.

Value

Scalar (for 'mean', 'max', 'min') or list (for 'all'). Note, results are in milimeters

Note

max = Hausdorff distance

Author(s)

Dorian Pustina

```
optimize_SCCANsparseness
```

Optimization of SCCAN sparseness

Description

Function used to optimize SCCAN sparseness for lesion to symptom mapping.

Usage

```
optimize_SCCANsparseness(lesmat, behavior, mask, nfolds = 4,
   sparsenessPenalty = 0.03, lower = 0.005, upper = 0.9, tol = 0.03,
   justValidate = F, cvRepetitions = 3, mycoption = 1, robust = 1,
   sparseness = NA, nvecs = 1, cthresh = 150, its = 30,
   npermsSCCAN = 0, smooth = 0.4, sparseness.behav = -0.99, ...)
```

Arguments

lesmat lesion matrix
behavior behavior vector
mask antsImage mask

nfolds how many folds to use

sparsenessPenalty

penalty term

lower minimum searched sparseness
upper maximum searched sparseness
tol tolerance value, see optimize() in R
justValidate just check the CV of provided sparseness

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```
cvRepetitions
```

```
number of cross-validations
                standard SCCAN parameter
mycoption
robust
                standard SCCAN parameter
sparseness
                standard SCCAN parameter
                standard SCCAN parameter
nvecs
cthresh
                standard SCCAN parameter
its
                standard SCCAN parameter
npermsSCCAN SCCAN permutations
smooth
                standard SCCAN parameter
sparseness.behav
                what sparsness to use for behavior
                other arguments received from lesymap or lsm_sccan.
```

Value

List with:

```
minimum - best sparseness value
objective - minimum value of objective function
CVcorrelation - cross-validated correlation of optimal sparness
```

Author(s)

Dorian Pustina

the optimization function Will run SCCAN on each training fold, compute behavior prediction on the test fold, and finally return a cross validated correlation from entire sample end of optimfun

```
print.lesymap print.lesymap
```

Description

Funciton to display some meaningful summary when a lesymap output is called in command line.

Usage

```
## S3 method for class 'lesymap'
print(x, ...)
```

Arguments

```
x the output from a lesymap() call.... useless for compatibility with default print.
```

Author(s)

26 save.lesymap

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Fast linear regressions

Description

Takes a matrix of voxels and a vector of behavior and runs fast regressions for each voxel. Covariates can be defined (i.e. age) to find the effect of each voxel on behavior within the context of other predictive factors.

Usage

```
regresfast(X, y, covariates, hascovar = FALSE)
```

Arguments

X matrix of voxlels (columns) for all subjects (rows).

y vector of behavioral scores.

covariates matrix with one or more columns. Must be of same length as behavior. This

variable should always be set, and the next argument can tell if covariates should

be used or not.

hascovar logical to tell whether covariates should be used.

Value

List with (1) statistic, (2) n = number of subjects, (3) kxfm = degrees of freedom.

Author(s)

Dorian Pustina

save.lesymap

Save the output of lesymap.

Description

Function to save the output of lesymap.

```
save.lesymap(lsm, saveDir, infoFile = "Info.txt", template = NA,
   saveTemplate = F, savePatchImages = T, plot.alpha = 0.8,
   plot.axis = 3, plot.quality = 8, ...)
```

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Arguments

lsm	object obtained with lesymap()
saveDir	folder to save to, will be created if it doesn't exist.
infoFile	(default='Info.txt') what should be the filename of the file with information.
template	(default=NA) an antsImage to overlay the results to. If the template is provided, results will be plotted and saved as image.
saveTemplate	(default=FALSE) should the template image also be saved? Useful when passing the results to a colleague.
savePatchImag	ges
	(default=TRUE) should the patch images be saved
plot.alpha	see plot.antsImage
plot.axis	see plot.antsImage
plot.quality	see plot.antsImage
	other arguments to use for plot().

Value

Nothing is returned. Files saved include resulting maps and a descriptive file with a lot of information about the lesymap run.

Author(s)

Dorian Pustina

```
simulateBehavior Simulation of behavior scores from lesion maps
```

Description

Function simulate behavioral scores based on the lesion load of specific brain areas. Used to run simulation studies.

Usage

```
simulateBehavior(lesions.list, parcellation, label = NA, mask = NA,
errorWeight = 0.3, binaryCheck = F, exponent = 1)
```

Arguments

lesions.list	list of lesions (antsImages) or vector of filenames.
parcellation	mask or parcellation image. If a parcellation is passed, lesion load will be computed for each different label (value) in the image. Zero and non-affected labels are not returned by default. The parcellation input can be an antsImage or a character vector pointing to a file.
label	if the parcellation has multiple labels, you can select which labels to simulate behaviors for (i.e., $c(101,43)$ to simulate behavior for labels with value 101 and 43 only)

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mask to restrict the count of lesioned voxels. It is not recommended to use

a mask, because lesions should affect behavior as they are, without the user

restricting the lesions to masks defined a posteriori.

errorWeight the amount of error to be added (i.e., 0.5 means half of the simulation will be

error, the other half signal)

binaryCheck check to make sure all lesions are binary

exponent power exponent to elevate behavior in order to increase non-linearity relation-

ship with lesion load. 1 is default, and 3 is what Wang (2013) reported as lesion

load relationship with behavior.

Value

List of three objects: - behavload - a matrix of simulated behavioral scores. Each column shows simulation for a single parcel. Column names indicate the label number in the parcellation file.

- lesload same as behavload, but indicates lesions loads from which behavior was simulated.
- lesbehavCorrelation vector of correlation values (Pearson) between lesion load and simulated scores.

Author(s)

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