# RImaGen

August 20, 2016

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## Description

Downsample an image.

## Usage

```
downsample(img, subFactor = 1, method = c("FLIRT", "SUBSAMP2"),
  bin = FALSE, thresh = 0.9)
```

## Arguments

img	An image of class nifti.	
subFactor	Downsampling factor.	
method	Downsampling method.	
bin	logical. Set to TRUE for downsampling of binary images or masks.	
thresh	Treshhold for re-binarisation of downsampled binary images and masks. threshold near 1 (say 0.9) is conservative according to FLIRT FAQ.	A

2 getSNPresults

#### Value

A downsampled image.

getSNPfdr

Get False Discovery Rates.

### Description

Get False Discovery Rates.

## Usage

```
getSNPfdr(p.value, eff.no.tests)
```

## Arguments

p. value A vector of raw p-values.

#### Value

A summary of FDR q-values.

 ${\tt getSNPresults}$ 

Get vGWAS results for selected SNPs.

## Description

Get vGWAS results for selected SNPs.

## Usage

```
getSNPresults(meh, sel.snps)
```

## Arguments

meh vGWAS results.

sel.snps character vector of SNP names.

#### Value

A named list of results for each SNP.

performVGWAS 3

performVGWAS	Perform voxelwise genome-wide association study.

### Description

Loads and prepares the data, runs preliminary (if necessary) and full analyses. Then visualises the results.

## Usage

```
performVGWAS(genePath, niiFiles, niiIDs, ref.imgPath, maskPath, subFactor,
  top.range, covar = character(), errorCovariance = numeric(),
  out.subFactor = subFactor, matPath = NULL, outPath = NULL,
  force.snps = NULL, useModel = MatrixEQTL::modelLINEAR, log.cutoff = 4,
  eff.no.tests = 275575, sampleSize = NULL, randomSample = FALSE,
  visualise = TRUE, saveNIfTI = TRUE, uncut = TRUE,
  mockPath.flatROIs = NULL, mockPath.pre = NULL)
```

### **Arguments**

genePath	Path to a directory containing plink files with genomic data.
niiFiles	Paths to files containing imaging data.
niiIDs	Subject IDs for each image. Shuffle for permutation tests.
ref.imgPath	Path to a reference image.
maskPath	Path to an image mask.
subFactor	Downsampling factor.
top.range	Range of top SNPs to be analysed and visualised.
covar	Covariates matrix with subject IDs as column names.
errorCovarianc	e
	Covariance matrix for the error term. Set to numeric() for multiple of identity (default, most cases).
out.subFactor	Output images downsampling factor. Default: equal to subFactor.
matPath	Path to convolution matrix for coregistration of results to the reference image. Set to NULL if in the same space.
outPath	Path to output directory.
force.snps	character vector of SNPs forced to be analysed even if not passing the quality control.
useModel	Regression model to use.
log.cutoff	Negative log p-value cutoff value for results visualisation.
eff.no.tests	Effective number of tests (SNPs) for p-value correction.
sampleSize	Subject sample size.
randomSample	logical. Set to TRUE for subject sample randomisation.
visualise	logical. Set to FALSE to disable results visualisation for bigger analyses.
saveNIfTI	logical. Set to FALSE to disable saving nifti results for bigger analyses.
uncut	logical. Set to FALSE to disable full maps generation.

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mockPath.flatROIs

Path to input file with saved flatROIs object for vGWAS mocking. Mock parameters should match original parameters. Useful for registering results to different standard spaces and resolutions, selecting alternative SNPs for the full analysys, of using different genetic models.

mockPath.pre

Path to input file with saved preliminary analysis results object for vGWAS mocking. Mock parameters should match original parameters. Useful for registering results to different standard spaces and resolutions, selecting alternative SNPs for the full analysis, or using different genetic models. NULL for real analyses.

#### Value

A list containing all the resulting statistical parametric maps.

readFlatR0Is

Read flattened ROIs from a list of NIfTI files using a cluster of choice.

#### **Description**

Read flattened ROIs from a list of NIfTI files using a cluster of choice.

#### Usage

```
readFlatROIs(paths, ids, subFactor = 0, mask = NULL, method = c("FLIRT",
   "SUBSAMP2"), nCores = getOption("mc.cores", detectCores(logical = FALSE)),
   clType = "PSOCK", makeCluster = TRUE, userCluster = NULL)
```

#### **Arguments**

paths A list of paths to the NIfTI files.

ids A list of subject IDs.
subFactor Downsampling factor.

mask Image mask (a nifti object or an array).

method Downsampling method.

nCores number of cores to use. Default: all physical cores available

clType cluster type. Default: PSOCK

makeCluster logical. TRUE if a cluster should be made (default).

userCluster Cluster to use specified by the user. If NULL and makeCluster is FALSE, a default

cluster will be used.

#### Value

Array of chosen voxel intensities for all the subjects.

readSNPs 5

readSNPs Read SNPs performing genome quality control.	
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#### **Description**

Read SNPs performing genome quality control.

#### Usage

```
readSNPs(plinkFiles, call.rate.cutoff = 0.95, maf.cutoff = 0.1,
  hwe.pval = 5.7e-07, min.group = 7, subjects = NULL, force.snps = NULL,
  forcedOnly = FALSE, outPath = NULL)
```

#### **Arguments**

plinkFiles Paths to the plink files. call.rate.cutoff Call rate cutoff value. maf.cutoff Minor allele frequency cutoff value. Hardy-Weinberg equilibrium p-value cutoff value. hwe.pval Minimum subjects per genotype group. Note that it can be only theoretically min.group derived from MAF under H-W equilibrium assumption. Set to 0 to rely solely on the HWE test. Vector of subject IDs to read data for. subjects character vector of SNPs forced to be analysed even if not passing the quality force.snps control. forcedOnly logical. If TRUE, only the SNPs of interest will be analysed, regardless of their

data quality.

outPath A character string giving the base filename for optional output of QC-ed data.

The extensions .bed, .bim, and .fam are appended to this string to give the file-

names of the three output files.

## Value

SnpMatrix of SNPs passing the quality control

report2mni	Convert snp report to MNI coordinates	
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## Description

Convert snp report to MNI coordinates

#### Usage

```
report2mni(path)
```

#### **Arguments**

path Path to report file.

6 toMni

RImaGen: An R package for voxelwise genome-wide association studies.

#### **Description**

This package allows you to perform vGWAS analyses as described in Stein et al., 2010, but faster.

selectSNPs

Select SNPs by name from a SlicedData object.

### Description

Select SNPs by name from a SlicedData object.

## Usage

```
selectSNPs(sel.snps, snpData)
```

### **Arguments**

sel.snps character vector of SNP names.

snpData SlicedData object containing SNP data.

#### Value

Array containing selected SNP data.

toMni

Convert voxel coordinates to MNI coordinates.

### **Description**

Convert voxel coordinates to MNI coordinates.

## Usage

```
toMni(v, res = 2)
```

## Arguments

v A 3D vector of voxel coordinates.

res Resolution in mm.

#### Value

MNI coordinates.

vGWAS 7

vGWAS	Perform voxelwise genome-wide association analysis.	

#### **Description**

Checks for associations prepared SlicedData objects containing flattened image ROIs, genomic data and covariates. Wrapper for Matrix\_eQTL\_engine.

### Usage

```
vGWAS(snpData, voxelData, cvrt, useModel, prescan,
errorCovariance = numeric(), pvalue.hist = FALSE)
```

#### **Arguments**

snpData SlicedData object containing selected SNPs.

voxelData SlicedData object containing flat voxel ROIs.

cvrt SlicedData object containing covariates.

useModel Regression model to use.

prescan logical. TRUE for preliminary analysis allowing "winning SNP" ranking.

errorCovariance

numeric. The error covariance matrix. Use numeric() for homoskedastic inde-

pendent errors.

pvalue.hist logical, numeric or "qq-plot". Defines whether and how the distribution of

p-values is recorded in the returned object. Set to FALSE for a faster analysis. To record information for a histogram set to the number of bins. To record information for a qq-plot, set to qq-plot. Use plot on the returned object.

#### Value

vGWAS results.

visualiseSNP	Visualise by-voxel statistics for a chosen SNP, e.g that SNP's p-value per voxel.
	per voxei.

#### **Description**

Visualise by-voxel statistics for a chosen SNP, e.g that SNP's p-value per voxel.

#### Usage

```
visualiseSNP(snp.name, results, mask, ref.img, pv.range = NULL,
  log.cutoff = 0, plot = TRUE, title = snp.name, titleSize = 2,
  beforeTitle = "-log10 p-values by\n", afterTitle = "", outPath = NULL,
  matPath = NULL, coordsOnly = FALSE)
```

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#### **Arguments**

snp.name Chosen SNP name.

results A named list of results for each SNP.

mask Image mask (a nifti object or an array).

ref.img Reference image.

pv.range P-value range for scale.

log.cutoff Negative log p-value cutoff value.

plot logical. Set to TRUE for automated plotting of the results.

title Plot title.
titleSize Title font size.

beforeTitle Text to print before the title.

afterTitle Text to print after the title.

outPath Output path. If NULL (default) no plot files will be saved.

matPath Path to convolution matrix for coregistration of results to the reference image.

Set to NULL if in the same space.

coordsOnly logical. Set to TRUE to return coordinates of the best voxel only.

#### Value

A list containing the results visualisation image and a vector of coordinates for the max. value voxel.

visualiseVox Visualise by-voxel statistics for each voxel, e.g winning SNP p-value per voxel.

#### **Description**

Visualise by-voxel statistics for each voxel, e.g winning SNP p-value per voxel.

#### Usage

```
visualiseVox(result, mask, ref.img, pv.range = NULL, log.cutoff = 0,
plot = TRUE, title = "winning SNP", titleSize = 2,
beforeTitle = "-log10 p-values by\n", afterTitle = "", outPath = NULL,
matPath = NULL, coordsOnly = FALSE)
```

#### **Arguments**

result Vector of target statistics.

mask Image mask.
ref.img Reference image.
pv.range P-value range for scale.

log.cutoff Negative log p-value cutoff value.

plot logical. Set to TRUE for automated plotting of the results.

visualise Vox 9

title Plot title. titleSize Title font size.

beforeTitle Text to print before the title.

afterTitle Text to print after the title.#'

outPath Output path. If NULL (default) no plot files will be saved.

matPath Path to convolution matrix for coregistration of results to the reference image.

Set to NULL if in the same space.

coordsOnly logical. Set to TRUE to return coordinates of the best voxel only.

#### Value

A list containing the results visualisation image and a vector of coordinates for the max. value voxel.

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