# Package 'neuroimaGene'

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Title Transcriptomic Atlas of Neuroimaging Derived Phenotypes

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Type Package

Version 0.1.2

Description Contains functions to query and visualize the Neuroimaging features associated with genetically regulated gene expression (GReX). The primary utility, neuroimaGene(), relies on a list of user-defined genes and returns a table of neuroimaging features (NIDPs) associated with each gene. This resource is designed to assist in the interpretation of genome-wide and transcriptome-wide association studies that evaluate brain related traits. Bledsoe (2024) <doi:10.1016 j.ajhg.2024.06.002="">. In addition there are several visualization functions that generate summary plots and 2-dimensional visualizations of regional brain measures. Mowinckel (2020).</doi:10.1016>
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Contents
anno

2 anno

	check_db																			
	fs_anno																			3
	listNIDPs																			4
	neuroimaGene																			5
	neuro_vis																			
	ng_vignette .							 												7
	plot_gnNIDP																			
	plot_gns																			9
	plot_nidps																			9
Index																				11

anno

NIDP annotation data

## **Description**

annotation data for all NIDPs taken from the UKbiobank

#### **Format**

A data.table with 10 columns and 3935 rows:

gwas\_phenotype character: UKB-derived neuroimaging derived phenotype (NIDP)

**modality** character: MRI neuroimaging modality **atlas** character: neuroimaging cortical atlas

side character: right or left hemisphere or midline/whole brain

primary character: primary cortical regionsecondary character: secondary cortical regionregion character: named region of the brain

measurement character: morphology measurement

fMRI\_node\_1 character: fMRI node 1
fMRI\_node\_2 character: fMRI node 2

NIDP character: user-friendly name for each NIDP

# Value

This script has no return. This is a documentation file for the annotation dataset for all neuroimaging derived phenotypes.

#### **Source**

Bledsoe, X. (2024) A transcriptomic atlas of the human brain reveals genetically determined aspects of neuropsychiatric health

check\_db 3

check\_db

Check neuroimaGene database downloaded

#### **Description**

Check if the NeuroimaGene database exists in the proper location prior to running the query and prompt user to download if not.

## Usage

```
check_db(timeout = 900)
```

## **Arguments**

timeout

time to spend downloading the NeuroimaGene database in seconds (default = 900)

#### Value

no return value, called to give information on status of neuroimaGene database and prompt user the user to download if resource file is missing.

#### **Examples**

```
check_db(timeout = 600)
```

fs\_anno

NIDP freesurfer annotation data

## **Description**

Freesurfer names for select cortical and subcortical NIDPs

# Usage

```
data(fs_anno)
```

#### **Format**

A data.table with 7 columns and 890 rows:

gwas\_phenotype character: UKB-derived neuroimaging derived phenotype

atl character: neuroimaging cortical atlas

hemisphere character: right or left hemisphere or midline/whole brain

secondary character: secondary cortical region

**fs\_name** character: freesurfer name **label** character: label name for the region

atlas character: freesurfer name for neuroimaging cortical atlas

4 listNIDPs

## Value

This script has no return. This is a documentation file for the annotation dataset for all neuroimaging derived phenotypes in the fsbrain package.

## **Source**

Bledsoe, X. (2024) A transcriptomic atlas of the human brain reveals genetically determined aspects of neuropsychiatric health

listNIDPs List NIDPs

## **Description**

Supplies a list of all NIDPs by name for any given modality or atlas

## Usage

```
listNIDPs(modality = NA, atlas = NA, filename = NA, verbose = FALSE)
```

## **Arguments**

modality	Neuroimaging modality. Defaults to NA; see README for additional options
atlas	Neuroimaging parcellation atlas for NIDP query. Defaults to NA; see README for additional options $$
filename	optional filename for writing data to a table
verbose	print runtime messages to R console. Default to FALSE

## Value

a list of NIDP names satisfying the required criteria

# **Examples**

```
dk_names <- listNIDPs(modality = 'T1', atlas = 'Desikan')</pre>
```

neuroimaGene 5

neuroimaGene	Main neuroimaGene query	
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# Description

Using a user supplied vector of genes (Ensembl ID's or HUGO names), this function queries the NeuroimaGene resource for all statistically significant GReX-NIDP associations and returns all findings as a data.table.

# Usage

```
neuroimaGene(
  gene_list,
  modality = "T1",
  atlas = "Desikan",
  mtc = "BH",
  nidps = NA,
  filename = NA,
  verbose = FALSE,
  vignette = FALSE
)
```

# Arguments

gene_list	List of genes to work as inputs. There are no defaults.
modality	Neuroimaging modality for NIDP query. Defaults to T1 structural NIDPs. Other common options include 'dMRI' for diffusion MRI imaging and 'fMRI' for fucntional MRI image results.
atlas	Neuroimaging parcellation atlas for NIDP query. Defaults to Desikan structural atlas. Common T1 atlases include the 'DKT', and 'Destrieux' cortical atlases and 'Subcortex' for freesurfer parcellation of subrotical regions. See package documentation or vignette for full list.
mtc	Statistical multiple testing correction for NIDP query. This defaults to 'BH' for the Benjamini Hochberg False Discovery Rate. Other options include 'BF' for the bonferroni family wise error rate and 'nom' for nominal findings at pvalue $\leq 0.05$ .
nidps	optional user defined vector of target NIDPs to query. Specific NIDP names can be obtained from the listNIDPnames() function. Use of this parameter overrides the multiple testing correction, returning all nominally significant findings.
filename	optional user defined path/filename to which the script will write the nueroima-Gene output data table.
verbose	print runtime messages to R console. Default to FALSE
vignette	use for building vignette on installation. Default to FALSE

neuro\_vis

#### Value

a neuroimaGene object: data table with all significant associations between the user provided genes and the UKB NIDPs satisfying the multiple testing correction and atlas/modality/name filters.

#### **Examples**

```
gene_list <- c('TRIM35', 'PROSER3', 'EXOSC6', 'PICK1', 'UPK1A', 'ESPNL', 'ZIC4')
ng <- neuroimaGene(gene_list, atlas = NA, mtc = 'BH', vignette = TRUE)</pre>
```

neuro\_vis

2D visualization plot of a neuroimaGene object

## **Description**

Generates a 2D visualization plot of the neuroimaGene object. Neuroimaging regions are defined by the atlas parameter and colored according to the magnitude and direction of the aggregate effect from each gene in the NeuroimaGene object. Colors can be defined by the user.

# Usage

```
neuro_vis(
  ng_obj,
  atlas = "Desikan",
  lowcol = "red2",
  midcol = "white",
  highcol = "royalblue2",
  title = NA
)
```

## Arguments

ng_obj	NeuroimaGene object produced by neuroimaGene() function
atlas	desired atlas for visualization. Desikan (default), Subcortex, DKT, Destrieux.
lowcol	color for low end of Zscore spectrum. Default is dark red
midcol	color for middle of Zscore spectrum. Default is white
highcol	color for top end of Zscore spectrum. Default is blue4
title	optional title tag for the plot

#### Value

class: ggplot object depicting 2D visualization of the NIDPs from the neuroimaGene object portrayed on the brain and shaded by mean effect size.

ng\_vignette 7

#### **Examples**

```
gene_list <- c('TRIM35', 'PROSER3', 'EXOSC6', 'PICK1', 'UPK1A', 'ESPNL', 'ZIC4')
ng <- neuroimaGene(gene_list, atlas = NA, mtc = 'BH', vignette = TRUE)
neuro_vis(ng, atlas = 'DKT')</pre>
```

ng\_vignette

NeuroimaGene object for vignette illustration

## **Description**

NeuroimaGene data table containing select associations used in the package vignette.

#### Usage

```
data(ng_vignette)
```

#### **Format**

A data.table with 6 columns and 3824 rows:

gene character: ENSEMBL Gene ID
gene\_name character: HUGO gene name

gwas\_phenotype character: neuroimaging derived phenotype

**training\_model** character: JTI derived tissue gene expression model **zscore** numeric: normalized effect size of GReX on NIDP morphology

mod\_BHpval character: Benjamini Hochberg corrected pvalue corrected by modality

#### Value

This script has no return. This is a documentation file for the neuroimaGene data subset required to build the vignette.

#### Source

Bledsoe, X. (2024) A transcriptomic atlas of the human brain reveals genetically determined aspects of neuropsychiatric health

8 plot\_gnNIDP

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Gene by NIDP summary plot of neuroimaGene object

## **Description**

Generate overview plot of the neuroimagene object according to gene/NIDP pair

# Usage

```
plot_gnNIDP(
   ng_obj,
   maxNidps = 20,
   maxGns = 15,
   title = NA,
   shortnames = TRUE,
   verbose = FALSE
)
```

# Arguments

ng_obj	NeuroimaGene object
maxNidps	maximum number of NIDPs to visualize. default=20
maxGns	maximum number of genes to visualize. default=15
title	optional title tag for the plot
shortnames	optional boolean tag for simplified names. Default to TRUE
verbose	print runtime messages to R console. Default to FALSE

## Value

a ggplot class heatmap showing tissue models per NIDP/Gene pair

# **Examples**

```
gene_list <- c('TRIM35', 'PROSER3', 'EXOSC6', 'PICK1', 'UPK1A', 'ESPNL', 'ZIC4')
ng <- neuroimaGene(gene_list, atlas = NA, mtc = 'BH', vignette = TRUE)
plot_gnNIDP(ng)</pre>
```

plot\_gns 9

plot\_gns

Gene summary plot of neuroimaGene object

## **Description**

Generates an overview plot of the neuroimaGene object according to each gene input.

# Usage

```
plot_gns(ng_obj, maxGns = 15, title = NA, verbose = FALSE)
```

## **Arguments**

ng\_obj NeuroimaGene Object
maxGns maximum number of genes to visualize. default=15
title optional title tag for the plot
verbose print runtime messages to R console. Default to FALSE

#### Value

a ggplot class plot detailing NIDPs per gene, colored by brain measure type

#### **Examples**

```
gene_list <- c('TRIM35', 'PROSER3', 'EXOSC6', 'PICK1', 'UPK1A', 'ESPNL', 'ZIC4')
ng <- neuroimaGene(gene_list, atlas = NA, mtc = 'BH', vignette = TRUE)
plot_gns(ng)</pre>
```

plot\_nidps

NIDP summary plot of NeuroimaGene object

## **Description**

Generate overview plot of the neuroimagene object according to nidps

## Usage

```
plot_nidps(
  ng_obj,
  maxNidps = 30,
  title = NA,
  shortnames = TRUE,
  mag = TRUE,
  verbose = FALSE
)
```

10 plot\_nidps

# Arguments

ng\_obj NeuroimaGene Object

maxNidps maximum number of NIDPs to visualize. default=30

title optional title tag for the plot

shortnames optional boolean tag for simplified names. Default to TRUE

mag boolean to present effect sizes by magnitude rather than as a vector. Default to

**TRUE** 

verbose print runtime messages to R console. Default to FALSE

## Value

a ggplot class object detailing mean effect size magnitude per NIDP, colored by brain region

## **Examples**

```
gene_list <- c('TRIM35', 'PROSER3', 'EXOSC6', 'PICK1', 'UPK1A', 'ESPNL', 'ZIC4')
ng <- neuroimaGene(gene_list, atlas = NA, mtc = 'BH', vignette = TRUE)
plot_nidps(ng)</pre>
```

# **Index**

```
\ast installation
    check_db, 3
*\ neuroimaging
    neuro_vis, 6
    neuroimaGene, 5
    plot_gnNIDP, 8
    plot_gns, 9
    plot_nidps, 9
*\ reference
    listNIDPs, 4
anno, 2
check_db, 3
fs_anno, 3
listNIDPs, 4
neuro_vis, 6
neuroimaGene, 5
ng\_vignette, 7
plot_gnNIDP, 8
plot_gns, 9
plot\_nidps, 9
```