Package 'gpaExample'

July 4, 2024

Type	Package
	Example data for the GPA package (Genetic analysis incorporating Pleiotropy and Annotation)
Versio	on 1.16.0
Deper	nds R (>= $4.0.0$)
Date	2020-02-24
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Descr	iption
	Example data for the GPA package, consisting of the p-values of 1,219,805 SNPs for five psychiatric disorder GWAS from the psychiatric GWAS consortium (PGC), with the annotation data using genes preferentially expressed in the central nervous system (CNS).
Licen	se $GPL (>= 2)$
URL	http://dongjunchung.github.io/GPA/
LazyI	Data FALSE
biocV	iews ExperimentData, Homo_sapiens_Data, SNPData
Needs	Compilation no
git_ur	thttps://git.bioconductor.org/packages/gpaExample
git_br	ranch RELEASE_3_19
git_la	st_commit bf7e7c8
git_la	st_commit_date 2024-04-30
Repos	sitory Bioconductor 3.19
Date/l	Publication 2024-07-04
Con	tents
	exampleData
Index	4

2 exampleData

exampleData

PGC GWAS Dataset and Annotation Dataset

Description

This is the PGC GWAS dataset and annotation dataset used in Chung et al. (2013).

Usage

data(exampleData)

Format

A list with two matrices as elements. Two matrices of sizes 1,219,805 x 5 and 1,219,805 x 1, exampleData\$pval and exampleData\$ann, contain the p-values of 1,219,805 SNPs for five psychiatric disorder GWAS (ADHD, ASD, BPD, MDD, SCZ) from the psychiatric GWAS consortium (PGC) and the binary annotation data using genes preferentially expressed in the central nervous system (CNS), respectively.

Details

Five columns of the matrix exampleData\$pval correspond to attention deficit-hyperactivity disorder (ADHD), autism spectrum disorder (ASD), bipolar disorder (BPD), major depressive disorder (MDD), and schizophrenia (SCZ). Detailed information about these data sets is provided in Cross-Disorder Group of the Psychiatric Genomics Consortium (2013a, 2013b). Summary statistics of these five psychiatric disorders were downloaded from the section for cross-disorder analysis at the Psychiatric Genomics Consortium (PGC) website (https://pgc.unc.edu/Sharing.php). exampleData\$ann provides annotation data using genes preferentially expressed in the central nervous system (CNS) (Lee et al., 2012; Raychaudhuri et al., 2010), where the entries corresponding to SNPs within 50-kb of the genes from the CNS set were set to be one and zero otherwise. See the vignette of R package GPA and Chung et al. (2013) for more details.

Source

Cross-Disorder Group of the Psychiatric Genomics Consortium (2013a), "Genetic relationship between five psychiatric disorders estimated from genome-wide SNPs", *Nature Genetics*, 45:984–994.

Cross-Disorder Group of the Psychiatric Genomics Consortium (2013b), "Identification of risk loci with shared effects on five major psychiatric disorders: a genome-wide analysis", *Lancet*, 381:1371–1379.

Lee SH, DeCandia TR, Ripke S, Yang J, Sullivan PF, et al. (2012), "Estimating the proportion of variation in susceptibility to schizophrenia captured by common SNPs", *Nature Genetics*, 44:247–250.

Raychaudhuri S, Korn JM, McCarroll SA, Altshuler D, Sklar P, et al. (2010), "Accurately assessing the risk of schizophrenia conferred by rare copy-number variation affecting genes with brain function", *PLoS Genetics*, 6:e1001097.

exampleData 3

References

Chung D, Yang C, Li C, Gelernter J, and Zhao H (2013), "GPA: A statistical approach to prioritizing GWAS results by integrating pleiotropy and annotation", To appear in PLoS Genetics.

Examples

data(exampleData)
head(exampleData)

Index

$*\ datasets$

exampleData, 2

 $\verb|exampleData|, 2$