Package 'FGRepo'

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Type Package
Title Functional Genomics Repository for POST-GWAS Analysis
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Author Alireza Ani [aut, cre], Ahmad Vaez [aut]
Maintainer Alireza Ani <a.ani@umcg.nl></a.ani@umcg.nl>
Depends R (>= 3.5)
Description A collection of datasets essential for functional genomic analysis. Gene names, gene positions, cytoband information, sourced from Ensembl and phenotypes association graph prepared from GWAScatalog are included. Data is available in both GRCh37 and 38 builds. These datasets facilitate a wide range of genomic studies, including the identification of genetic variants, exploration of genomic features, and post-GWAS functional analysis.
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LazyDataCompression xz
RoxygenNote 7.3.2
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cytoband_grch37

Cytoband information (GRCh37)

Description

A dataset containing cytoband information based on GRCh37.

Usage

```
cytoband_grch37
```

Format

A data frame with 4 columns:

chr chromosome

start start position

end end position

band band name

Source

```
Ensembl, https://www.ensembl.org
```

References

Harrison, PW, et al. (2024). Ensembl 2024. Nucleic Acids Research, 52, Pages D891–D899. doi:10.1093/nar/gkad1049

Examples

```
data(cytoband_grch37)
```

cytoband_grch38

Cytoband information (GRCh38)

Description

A dataset containing cytoband information based on GRCh38.

Usage

```
cytoband_grch38
```

gene_names_grch37

Format

A data frame with 4 columns:

chr chromosome
start start position

end end positionband band name

Source

```
Ensembl, https://www.ensembl.org
```

References

Harrison, PW, et al. (2024). Ensembl 2024. Nucleic Acids Research, 52, Pages D891–D899. doi:10.1093/nar/gkad1049

Examples

```
data(cytoband_grch38)
```

gene_names_grch37

Gene Names (GRCh37)

Description

A dataset containing gene names based on GRCh37.

Usage

```
gene_names_grch37
```

Format

A data frame with 6 columns:

id Gene identifier

name Gene name

chr Chromosome

start Start position

end End position

type Gene type

Source

```
Ensembl, https://www.ensembl.org
```

gene_names_grch38

References

Harrison, PW, et al. (2024). Ensembl 2024. Nucleic Acids Research, 52, Pages D891–D899. doi:10.1093/nar/gkad1049

Examples

```
data(gene_names_grch37)
```

gene_names_grch38

Gene Names (GRCh38)

Description

A dataset containing gene names based on GRCh38.

Usage

```
gene_names_grch38
```

Format

A data frame with 6 columns:

id Gene identifier

name Gene name

chr Gene name

start Gene name

end Gene name

type Gene name

Source

```
Ensembl, https://www.ensembl.org
```

References

```
Harrison, PW, et al. (2024). Ensembl 2024. Nucleic Acids Research, 52, Pages D891–D899. doi:10.1093/nar/gkad1049
```

Examples

```
data(gene_names_grch38)
```

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GWASCat_graph

GWAScatalog graph

Description

A graph object of GWAScatalog data. The association between variants and phenotypes are indicated.

Usage

GWASCat_graph

Format

An object of igraph class.

Source

```
GWAS Catalog, https://www.ebi.ac.uk/gwas
```

References

Sollis, E, et al. (2022). The NHGRI-EBI GWAS Catalog: knowledgebase and deposition resource. Nucleic Acids Research, 51, Pages D977–D985. doi:10.1093/nar/gkac1010

Examples

```
data(GWASCat_graph)
```

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