Package 'xcoredata'

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Type Package Title data package for xcore Version 1.13.0 **Description** Provides data to use with xcore package. **Depends** R (>= 4.2)Imports ExperimentHub (>= 2.2.0), utils (>= 4.2.0) **Suggests** BiocGenerics (>= 0.40.0), data.table (>= 1.14.2), GenomeInfoDb (>= 1.30.0), GenomicRanges (>= 1.46.1), IRanges (>= 2.28.0), knitr (>= 1.37), rmarkdown (>= 2.11), Matrix (>= 1.3.4), stringr (>= 1.4.0), S4Vectors (>= 0.32.3), TxDb.Hsapiens.UCSC.hg38.knownGene (>= 3.14.0), xcore License GPL-2 **Encoding** UTF-8 RoxygenNote 7.1.2 VignetteBuilder knitr biocViews ExperimentHub, ExperimentData, Homo_sapiens_Data git_url https://git.bioconductor.org/packages/xcoredata git_branch devel git_last_commit aa6c0d6 git_last_commit_date 2025-04-15 **Repository** Bioconductor 3.22 Date/Publication 2025-07-15 Author Maciej Migdał [aut, cre] (ORCID: <https://orcid.org/0000-0002-8021-7263>), Bogumił Kaczkowski [aut] (ORCID: <https://orcid.org/0000-0001-6554-5608>)

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chip_atlas_meta ChIP-Atlas metadata

Description

Metadata associated with chip_atlas_promoters.

Usage

chip_atlas_meta

Format

A data.frame with 13891 rows and 5 columns.

id Character giving internal experiment ID.

tf Character giving transcription factor name.

tf_dbd Character giving transcription factor DNA binding domain family, as per CIS BP database.

biotype Character giving experiment biological origin.

study Character giving study ID.

Examples

chip_atlas_meta()

chip_atlas_promoters_f5

ChIP-Atlas FANTOM5 promoters intersection matrix

Description

An intersection matrix describing overlaps between ChIP-Atlas's ChIP-seq tracks and promoters_f5. To find overlapping regions promoters were extended by 500bp in both directions.

Usage

chip_atlas_promoters_f5

entrez2fantom

Format

A Matrix with 209911 rows and 13891 columns. Row names corresponds to promoters names, column names are formatted as TranscriptionFactor_Origin_Cell_ExperimentID (eg. PARK7_Neural_SH-SY5Y_DRX000550, MLL-AF6_Blood_ML-2_DRX001460).

Examples

chip_atlas_promoters_f5()

entrez2fantom ENTREZ IDs to FANTOM5 core promoters names mappings

Description

Vector mapping ENTREZ IDs to FANTOM5 core promoters names.

Usage

entrez2fantom

Format

A named character vector of length 14214.

Examples

entrez2fantom()

promoters_f5 Prom

Promoters GenomicRanges object

Description

FANTOM5's hg38 promoters annotated with nearest features in GENCODE ver. 38 annotation and UCSC hg38 knownGene annotation ver. 3.13.0.

Usage

promoters_f5

Format

A GenomicRanges object of length 209911, with 11 metadata columns:

name Promotor name.

score Numeric vector.

gene_type_gencode Gene type of associated gene as defined by GENCODE annotation.

ENTREZID ENTREZ ID of associated gene as defined by all three annotation sources. Prevalence of annotations: UCSC > GENCODE > FANTOM5.

SYMBOL Gene symbol of associated gene as defined by all three annotation sources. Prevalence of annotations: UCSC > GENCODE > FANTOM5.

Examples

promoters_f5()

promoters_f5_core Core promoters GenomicRanges object

Description

Core promoters selected from promoters_f5. Selection criteria were GENCODE confirmation and ENCODE ROADMAP confirmation. Further for each gene single promoter with highest FAN-TOM5 score was selected.

Usage

promoters_f5_core

Format

A GenomicRanges object of length 14191, with 16 metadata columns:

name Promotor name.

score Numeric vector.

gene_type_gencode Gene type of associated gene as defined by GENCODE annotation.

- **ENTREZID** ENTREZ ID of associated gene as defined by all three annotation sources. Prevalence of annotations: UCSC > GENCODE > FANTOM5.
- **SYMBOL** Gene symbol of associated gene as defined by all three annotation sources. Prevalence of annotations: UCSC > GENCODE > FANTOM5.

Examples

promoters_f5_core()

remap_meta

ReMap2020 metadata

Description

Metadata associated with remap_promoters.

Usage

remap_meta

Format

A data.table with 5798 rows and 6 columns.

id Character giving internal experiment ID.

tf Character giving transcription factor name.

tf_dbd Character giving transcription factor DNA binding domain family, as per CIS BP database.

biotype Character giving experiment biological origin.

study Character giving study ID.

condition Character specifiying experiment conditions or treatment.

Examples

remap_meta()

remap_promoters_f5 ReMap2020 and FANTOM5 promoters intersection matrix

Description

An intersection matrix describing overlaps between ReMap2020's ChIP-seq tracks and promoters_f5. To find overlapping regions promoters were extended by 500bp in both directions.

Usage

remap_promoters_f5

Format

A Matrix with 209911 rows and 5728 columns. Row names corresponds to promoters names, column names are formatted as ExperimentID.TranscriptionFactor.Biotype.

Examples

remap_promoters_f5()

symbol2fantom

Description

Vector mapping gene symbols to FANTOM5 core promoters names.

Usage

symbol2fantom

Format

A named character vector of length 14222.

Examples

symbol2fantom()

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