Package ‘depmap’

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Type Package

Title Cancer Dependency Map Data Package

Version 1.16.0

Description The depmap package is a data package that accesses datasets from the Broad Institute DepMap cancer dependency study using ExperimentHub. Datasets from the most current release are available, including RNAI and CRISPR-Cas9 gene knockout screens quantifying the genetic dependency for select cancer cell lines. Additional datasets are also available pertaining to the log copy number of genes for select cell lines, protein expression of cell lines as measured by reverse phase protein lysate microarray (RPPA), Transcript Per Million (TPM) data, as well as supplementary datasets which contain metadata and mutation calls for the other datasets found in the current release. The 19Q3 release adds the drug_dependencies dataset, that contains cancer cell line dependency data with respect to drug and drug-candidate compounds. The 20Q2 release adds the proteomic dataset that contains quantitative profiling of proteins via mass spectrometry. This package will be updated on a quarterly basis to incorporate the latest Broad Institute DepMap Public cancer dependency datasets. All data made available in this package was generated by the Broad Institute DepMap for research purposes and not intended for clinical use. This data is distributed under the Creative Commons license (Attribution 4.0 International (CC BY 4.0)).

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achilles

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achilles  achilles_22Q2

Description

The ‘achilles’ dataset contains the metadata for DepMap’s cancer models/cell lines, specific to Project Achilles’ 22Q2 CRISPR screens. This dataset can be loaded into the R environment with the 'depmap_achilles' function.

Usage

achilles
achilles

Format

A data frame with 1036 rows and 6 variables:

- **depmap_id**: cancer cancer cell line foreign key (i.e. "ACH-00001")
- **cell_line**: CCLE name of cancer cell line (i.e. "184A1_BREAST")
- **achilles_n_replicates**: Number of replicates used in Achilles CRISPR screen passing QC
- **cell_line_NNMD**: Difference in the means of positive and negative controls normalized by the standard deviation of the negative control distribution
- **culture_medium**: Medium used to grow cell line
- **cas9_activity**: Percentage of cells remaining GFP negative on days 12-14 of cas9 activity assay as measured by FACs

Details

This data represents the ‘Achilles_metadata.csv’ file taken from the 22Q2 [Broad Institute](https://depmap.org/portal/download/) release.

Change log

- 22Q1: Initial dataset
- 22Q2: "culture_type" column was removed

Source

DepMap, Broad (2022)

References


Examples

```
## Not run:
depmap_achilles()

## End(Not run)
```
Description

The `copyNumber` dataset contains the 22Q2 inferred copy number data for genes and cancer cell lines. This dataset includes data from 25368 genes, 1754 cell lines, 35 primary diseases and 38 lineages. The columns of `copyNumber` are: `depmap_id`, a foreign key corresponding to the cancer cell lineage, `cell_line` containing the common CCLE name of the cancer cell lines, `gene` containing both the HUGO gene name of the knockdown gene along with entrez ID#, `gene_name` containing only the HUGO gene name, `entrez_id` containing only the entrez ID#, and `log_copy_number` containing the numerical dependency score values for each pair of genes and cell lines. This dataset can be loaded into the R environment with the `depmap_copyNumber` function.

Usage

`copyNumber`

Format

A data frame with 44799888 rows (cell lines) and 6 variables.

- **depmap_id**: Cancer cell line foreign key (i.e. "ACH-00001")
- **gene**: HUGO symbol (e.g. "SAP25") and Entrez ID# (e.g. 100316904)
- **gene_name**: HUGO symbol (e.g. "SAP25")
- **entrez_id**: Entrez ID# (e.g. 100316904)
- **log_copy_number**: numerical log fold change in copy number for a given gene and cell line
- **cell_line**: CCLE name of cancer cell line (i.e. "184A1_BREAST")

Details

This data represents the `CCLE_gene_cn.csv` file taken from the 22Q2 [Broad Institute](https://depmap.org/portal/download/) cancer dependency study. The derived dataset found in the `depmap` package features the addition of a foreign key `depmap_id` found in the first column of this dataset, which was added from the `metadata` dataset. This dataset has been converted to a long format tibble. Variables names from the original dataset were converted to lower case, put in snake case, and abbreviated where feasible.

Change log

- 19Q1: Initial dataset consisted of a data frame with 37907473 rows (cell lines) and 6 variables representing 23299 genes, 1604 cell lines, 38 primary diseases and 33 lineages.
- 19Q2: adds 23 cell lines and 1 lineage
- 19Q3: adds 3263 genes, 30 cell lines and removes 2 primary diseases. Now a dataframe with 45670234 rows and 6 variables
- 19Q4: adds 77 genes, 25 cell lines, 0 primary diseases and 2 lineages
- 20Q1: adds 31 cell lines
- 20Q2: adds 32 cell lines
- 20Q3: no change
- 20Q4: removes 77 genes, adds 8 cell lines and 1 lineage
- 21Q1: removes 13 cell lines
- 21Q2: adds 2 cell lines
- 21Q3: removes 1 cell line and 1 lineage
- 21Q4: removes 194 genes and adds 9 cell lines
- 22Q1: removes 2000 genes, adds 4 cell lines and 1 lineage
- 22Q2: adds 12 cell lines and removes 2 primary diseases and 8 lineages

Source

DepMap, Broad (2020)

References


Examples

```r
## Not run:
depmap_copyNumber()

## End(Not run)```
Description

The ‘crispr’ dataset contains the 22Q2 genetic effect of CRISPR-Cas9 gene knockout of select genes in various cancer cell lines. This dataset includes data from 17386 genes, 1086 cell lines, 31 primary diseases and 28 lineages. The columns of ‘crispr’ are: ‘depmap_id’ a foreign key corresponding to the cancer cell lineage, ‘cell_line’ containing the common CCLE name of the cancer cell lines, ‘gene’ containing both the HUGO gene name of the knockout gene along with entrez ID#, ‘gene_name’ containing only the HUGO gene name, ‘entrez_id’ containing only the entrez ID# and ‘dependency’ which contains the numerical dependency score values for each pair of genes and cell lines. This dataset can be loaded into R environment with the ‘depmap_crispr’ function. NOTE: the ‘crispr’ dataset has replaced the ‘rnai’ dataset as the primary supported measurement of cancer dependency data as of the 19Q4 release.

Usage

```r
crispr
```

Format

A data frame with 18881196 rows (cell lines) and 6 variables:

- **depmap_id** Cancer cell line foreign key (i.e. "ACH-00001")
- **gene** HUGO symbol (e.g. "SAP25") and Entrez ID# (e.g. 100316904)
- **gene_name** HUGO symbol (e.g. "SAP25")
- **entrez_id** Entrez ID# (e.g. 100316904)
- **dependency** numerical dependency score of given gene and cell line
- **cell_line** CCLE name of cancer cell line (i.e. "184A1_BREAST")

Details

This data represents the ‘Achilles_gene_effect.csv’ file taken from the 22Q2 [Broad Institute](https://depmap.org/portal/download/) cancer depenedency study. The derived dataset found in the ‘depmap’ package features the addition of a foreign key ‘depmap_id’ found in the first column of this dataset, which was added from the ‘metadata’ dataset. This dataset has been converted to a long format tibble. Variables names from the original dataset were converted to lower case, put in snake case, and abbreviated where feasible.

Change log

- 19Q1: Initial dataset consisted of a data frame with 9839772 rows (cell lines) and 6 variables representing 17634 genes, 558 cell lines, 26 primary diseases and 28 lineages.
- 19Q2: adds 5 cell lines, 1 primary disease and 1 lineage
- 19Q3: adds 699 genes, 62 cell lines and 1 primary disease. Now a dataset with 11458125 rows and 6 variables.
- 19Q4: adds 0 genes, 64 cell lines, 1 primary disease and 0 lineages.
- 20Q1: adds 50 cell lines.
- 20Q2: adds 30 cell lines.
- 20Q3: adds 20 cell lines, 1 primary disease and 1 lineages.
- 20Q4: adds 19 cell lines, 1 primary disease and 2 lineages
- 21Q1: no change
- 21Q2: removes 474 genes and adds 182 cell lines 1 primary disease and 1 lineage
- 21Q3: removes 252 genes and adds 42 cell lines
- 21Q4: removes 7 genes and 22 cell lines
- 22Q1: adds 16 cell lines and 1 lineage
- 22Q2: adds 16 cell lines and removes 1 primary diseases and 3 lineages.

Source
DepMap, Broad (2020)

References

Examples
## Not run:

defmap_crispr()

## End(Not run)
Description

The depmap package is a data package that accesses datasets from the Broad Institute DepMap cancer dependency study using ExperimentHub. Datasets from the most current release are available, including RNAI and CRISPR-Cas9 gene knockout screens quantifying the genetic dependency for select cancer cell lines. Additional datasets are also available pertaining to the log copy number of genes for select cell lines, protein expression of cell lines as measured by reverse phase protein lysate microarray (RPPA), 'Transcript Per Million' (TPM) data, chemical dependency (drug_sensativity), mass spectrometry proteomic quantification (proteomic) as well as supplementary datasets which contain metadata and mutation calls for the other datasets found in the current release. This package will be updated on a quarterly basis to incorporate the latest Broad Institute DepMap Public cancer dependency datasets. All data made available in this package was generated by the Broad Institute DepMap for research purposes and not intended for clinical use. This data is distributed under the Creative Commons license (Attribution 4.0 International (CC BY 4.0)).

Details

See the package vignettes and respective man pages for details.

Description

Returns the depmap release

Usage

depmap_release()

Value

character(1) with the depmap release
The ‘drug_sensitivity’ dataset contains the 19Q3 replicate collapsed logfold change values relative to DMSO, corrected for experimental confounders using ComBat. This dataset contains information referring to 4686 compounds, 578 cell lines, 23 primary diseases and 25 lineages. This dataset is part of the SIGMA Re-purposing release which contains small molecule viability datasets generated using the Broad Re-purposing Library and the PRISM multiplexed cell-line viability assay. The columns of ‘drug_sensitivity’ are: ‘depmap_id’ a foreign key corresponding to the cancer cell lineage, ‘cell_line’ the common CCLE name of the cancer cell lines, ‘compound’ the synonym for the drug compound, and ‘dependency’ which contains the numerical dependency score values for each pair of genes and cell lines. Compounded metadata has also been added.

Usage

drug_sensitivity

Format

A data frame with 2708508 rows (cell lines) and 14 variables:

- **depmap_id**: Cell line foreign key (i.e. "ACH-000956")
- **cell_line**: Name of cancer cell line (i.e. "22RV1_PROSTATE")
- **compound**: Drug compound name (i.e. BRD-A00077618-236-07-6::2.5::HTS)
- **dependency**: Numerical dependency score of a compound on a cell line
- **broad_id**: Broad ID for compound (i.e. BRD-A00077618-236-07-6)
- **name**: Standard chemical name (i.e. 8-bromo-cGMP)
- **dose**: Dose of compound
- **screen_id**: Broad ID for compound (i.e. HTS)
- **moa**: Mechanism of action (i.e. PKA activator)
- **target**: Molecular target of compound (i.e. PRKG1)
- **disease_area**: Anatomical target (i.e. hematologic malignancy)
- **indication**: Prescribed for disease (i.e. acute myeloid leukemia (AML))
- **smiles**: Simplified molecular-input line-entry specification
- **phase**: Clinical phase
Details
This data originates from the ‘primary_replicate_collapsed_logfold_change.csv’ file taken from the 19Q3 [Broad Institute](https://depmap.org/portal/download/) cancer dependency study. The derived dataset found in the ‘depmap’ package features the addition of a foreign key ‘depmap_id’ found in the first column of this dataset, which was added from the ‘metadata’ dataset. This dataset has been converted to a long format tibble. Variables names from the original dataset were converted to lower case, put in snake case, and abbreviated where feasible. Note: compound metadata was added to this dataset, consisting of 10 new features derived from the file ‘primary-screen-replicate-collapsed-treatment-info.csv’ from the [Broad Institute](https://depmap.org/portal/download/) website. The Drug sensitivity data remains from 19Q3, however the version was bumped to 21Q2 to distinguish between the different datasets.

Change log
- 19Q3: Initial dataset consisted of a data frame with 2708508 rows (cell lines) and 6 variables representing 686 compounds, 578 cell lines, 23 primary diseases and 25 lineages.
- 19Q4: no change, no further releases are scheduled at this time.
- 20Q1: no change, no further releases are scheduled at this time.
- 20Q2: no change, no further releases are scheduled at this time.
- 20Q3: no change, no further releases are scheduled at this time.
- 20Q4: no change, no further releases are scheduled at this time.
- 21Q1: no change, no further releases are scheduled at this time.
- 21Q2: Drug sensitivity data combined with compound metadata, added 10 new features from the file primary-screen-replicate-collapsed-treatment-info.csv
- 21Q3: no change, no further releases are scheduled at this time.
- 21Q4: no change, no further releases are scheduled at this time.
- 22Q1: no change, no further releases are scheduled at this time.
- 22Q2: no change, no further releases are scheduled at this time.

Source
DepMap, Broad (2021)

References
Description

The `gene_summary` dataset contains the 22Q1 gene essentiality probabilities for select genes. This dataset can be loaded into the R environment with the `depmap_gene_summary` function.

Usage

gene_summary

Format

A data frame with 69746 rows (cell lines) and 7 variables:

- `entrez_id` Entrez ID# (e.g. 100316904)
- `gene_name` HUGO symbol (e.g. "SAP25")
- `dataset` which dataset this probability derives
- `dependent_cell_lines` number of dependent cell lines
- `cell_lines_with_data` number of cell lines with relevant dependency data
- `strongly_selective` Gene knockout is selective (not pan-lethal)
- `common_essential` common essential gene dependency

Details

This data represents the 'Gene Dependency Profile Summary.csv' file taken from the 22Q1 [Broad Institute](https://depmap.org/portal/api/download/gene_dep_summary) release.

Change log

- 22Q1: Initial dataset
- 22Q2: no change, no further releases are scheduled at this time.

Source

DepMap, Broad (2022)

References


Examples

```r
## Not run:
depmap_gene_summary()

## End(Not run)
```

### Description

The `metadata` dataset contains the metadata about cell lines in the 22Q2 Broad Institute DepMap release, which includes mapping between 'depmap_id' and 'cell_line' name for cancer cell lines. This dataset does not contain any data from the Achilles screen nor dependency data, but contains the metadata from the other datasets pertaining to the 22Q1 DepMap release, for 1840 cell lines, 0 genes, 33 primary diseases and 30 lineages. The columns of `metadata` are: 'depmap_id', 'stripped_cell_line_name', 'cell_line', 'aliases', 'cosmic_id', 'sanger_id', 'WTSI_master_cell_ID', 'primary_disease', 'subtype_disease', 'sub_subtype_disease', 'gender', 'source'. This dataset can be loaded into the R environment with the `depmap_metadata` function.

### Usage

`metadata`

### Format

A data frame with 1829 rows (cell lines) and 22 variables:

- **depmap_id** Cancer cell line primary key (i.e. "ACH-00001")
- **stripped_cell_line_name** Name of stripped cell line
- **cell_line** CCLE name of cancer cell line (i.e. "184A1_BREAST")
- **cell_line_name** Abbreviated name of cancer cell line (i.e. "NIH:OVCAR-3")
- **aliases** Aliases of cancer cell line
- **cosmic_id** Catalogue Of Somatic Mutations In Cancer ID number (e.g. 905933)
- **sex** Sex of tissue sample
- **source** Source of tissue sample
- **culture_type** Culture type of tissue sample
- **RRID** Resource Identification Portal ID
- **sample_collection_site** Site of sample collection (AML), M3 (Promyelocytic)
- **primary_or_metastasis** Primary cancer cell line or metastatic
- **primary_disease** Primary Disease (e.g. cancer type)
- **subtype_disease** Subtype Disease (e.g. Acute Myelogenous Leukemia)
metadata

age  Age of individual sample of cell line was derived
sanger_id  Sanger ID (eg. 2201)
WTSI_master_cell_ID  Wellcome Trust Sanger Institute ID (eg. 1369)
additional_info  Additional information about samples
lineage  Lineage of cancer cell line
lineage_subtype  Subtype of lineage of cancer cell line
lineage_sub_subtype  Subtype of subtype of Lineage of cancer cell line
lineage_molecular_subtype  Molecular type of Lineage of cancer cell line
model_manipulation  Culture model manipulation details
model_manipulation_details  Culture model manipulation details
patient_id  Patient id
parent_patient_id  Parent patient id
Cellosaurus_NCIt_disease  Cellosaurus NCIt disease
Cellosaurus_NCIt_id  Cellosaurus NCIt_id
Cellosaurus_NCIt_id  Cellosaurus NCIt_id

Details

This data represents the 'sample_info.csv' file taken from the 22Q2 [Broad Institute](https://depmap.org/portal/download/) cancer dependency study. This dataset features the a primary key 'depmap_id' which is a unique ID given to each cell line and is found in the first column of this dataset. The 'depmap_id' attribute is used as a foreign key in all other datasets in the package. This dataset has been converted to a long format tibble. This dataset does not contain any expression or dependency data but rather contains the metadata for all cancer cell lines used in the depmap project. Variables names were converted to lower case, put in snake case, and abbreviated where feasible (e.g. "Sanger ID” was changed to "sanger_id").

Change log

- 19Q1: Initial dataset consisted of data frame with 1677 rows (cell lines) and 9 variables, representing 0 genes, 1677 cell lines, 38 primary diseases and 33 lineages
- 19Q2: adds 37 new cell lines, 1 primary disease and 1 lineage. This version of the metadata dataset contains 6 variables not found in previous versions, relating the the Achilles metadata: 'Achilles_n_replicates', 'cell_line_NNMD', 'culture_type', 'culture_medium', and 'cas9_activity'.
- 19Q3: adds 30 cell lines, 2 primary diseases and 2 lineages
- 19Q4: adds 42 cell lines, 0 primary diseases and 3 lineages
- 20Q1: adds 19 cell lines, 'gender' was changed to 'sex', 'age', 'primary_or_metastasis' and 'sample_collection_site” were added
- 20Q2: adds 30 cell lines and 1 lineage
- 20Q3: adds new column 'WTSI_master_cell_ID'
- 20Q4: adds 6 cell lines and 1 lineage. Adds column ‘cell_line_name’
- 21Q1: removes 1 cell line
- 21Q2: adds 3 cell lines
- 21Q3: adds 1130 cell lines, 8 primary diseases and 8 lineages
- 21Q4: removes 1119 cell lines, 8 primary diseases and 8 lineages
- 22Q1: adds 4 cell lines. The features relating to Achilles metadata have been removed and put into their own dataset: ‘Achilles_n_replicates’, ‘cell_line_NNMD’, ‘culture_type’, ‘culture_medium’, and ‘cas9_activity’.
- 22Q2: adds 11 cell lines and removes 2 primary diseases and 30 lineages. The feature ‘culture_type’ has been removed and columns "model_manipulation", "model_manipulation_details", "patient_id", "parent_depmap_id", "Cellosaurus_NCIt_disease", "Cellosaurus_NCIt_id" and "Cellosaurus_issues" have been added.

Source
DepMap, Broad (2020)

References

Examples
```r
## Not run:
depmap metadata()
## End(Not run)
```

---

**mutationCalls**  
**mutationCalls_22Q2**

Description
The ‘mutationCalls’ dataset contains merged the 22Q2 mutation calls (for coding region, germline filtered) and includes data from 18784 genes, 1771 cell lines, 33 primary diseases and 30 lineages. This dataset can be considered the metadata data set for mutations and does not contain any dependency data. This dataset can be loaded into the R environment with the ‘depmap_mutationCalls’ function.
**Usage**

`mutationCalls`

**Format**

A data frame with 1235466 rows and 32 variables:

- `depmap_id` `depmap_id`
- `gene_name` Hugo Symbol denotes a unique and meaningful name for each gene (e.g. SAP25)
- `entrez_id` Gene ID for NCBI Entrez gene database, (e.g. 100316904)
- `ncbi_build` NCBI Build (i.e. reference genome)
- `chromosome` Chromosome
- `start_pos` Gene start position
- `end_pos` Gene end position
- `strand` Strand location of gene
- `var_class` Variant Classification
- `var_type` Variant Type
- `ref_allele` Reference Allele
- `alt_allele` Tumor Seq Allele1
- `dbSNP_RS` Single Nucleotide Polymorphism Database (dbSNP) reference cluster
- `dbSNP_val_status` dbSNP Val Status
- `genome_change` Genome Change
- `annotation_transcript` Annotation Transcript
- `cDNA_change` change in cDNA
- `codon_change` Codon Change
- `protein_change` Protein_Change
- `is_deleterious` Status of gene knockout on cell lineage
- `is_tcga_hotspot` isTCGAhotspot
- `tcga_hsCnt` TCGAhCnt
- `is_cosmic_hotspot` isCOSMIChotspot
- `cosmic_hsCnt` COSMIChsCnt
- `ExAC_AF` ExAC_AF
- `CGA_WES_AC` CGA_WES_AC
- `sanger_WES_AC` SangerWES_AC
- `RNAseq_AC` RNAseq_AC
- `HC_AC` HC.AC
- `RD_AC` RD.AC
- `WGS_AC` WGS.AC
- `var_annotation` Variant_annotation
Details

This data represents the ‘CCLE_mutations.csv’ file taken from the 22Q2 [Broad Institute](https://depmap.org/portal/download/) cancer dependency study. The derived dataset found in the ‘depmap’ package features the addition of a foreign key ‘depmap_id’ found in the first column of this dataset, which was added from the ‘metadata’ dataset. This dataset has been converted to a long format tibble. Variables names from the original dataset were converted to lower case, put in snake case, and abbreviated where feasible.

Change log

- 19Q1: Initial dataset for package consisted of dataframe with 1243145 rows and 35 variables representing 18755 genes, 1601 cell lines, 37 primary diseases and 33 lineages
- 19Q2: adds 30 cell lines, 1 primary disease and 1 lineage. This version has different columns than the previous version: the variable "VA_WES_AC" is no longer present in this dataset. Some minor alterations to the original file were made. The first column of the original dataset, (itemIDSample number) was removed, as this column was only the row number and did not serve any unique identifying purpose.
- 19Q3: adds 1 gene, 25 cell lines and removes 1 primary disease
- 19Q4: adds 1 gene, 10 cell lines, 0 primary diseases and 2 lineages
- 20Q1: adds 4 genes, 31 cell lines, 1 lineage
- 20Q2: adds 44 cell lines, 1 lineage
- 20Q3: no change
- 20Q4: removes 13 genes, adds 8 cell lines and 1 lineage. Columns ‘tumor_sample_barcode’ and ‘sanger_recalib_WES_AC’ were removed
- 21Q1: removes 11 genes and 2 cell lines
- 21Q2: removes 1 genes and adds 3 cell lines
- 21Q3: removes 3 genes, 4 cell lines and 1 lineage
- 21Q4: adds 9 cell lines
- 22Q1: adds 4 cell lines and 1 lineage. The variable ‘tumor_seq_allele1’ was renamed ‘alt_allele’
- 22Q2: adds 12 cell lines and removes 2 primary diseases and 8 lineages.

Source

DepMap, Broad (2020)

References

Examples

```r
## Not run:
depmap_mutationCalls()

## End(Not run)
```

---

Description

The 'proteomic' dataset contains the 20Q2 quantitative profiling of proteins via mass spectrometry from the Gygi lab. This dataset contains 12399 proteins tested in 375 cell lines, including 24 primary diseases and 27 lineages. The columns of this dataset are: 'depmap_id', a foreign key corresponding to the cancer cell lineage, 'cell_line' the common CCLE name of the cancer cell lines, 'gene_name' containing the HUGO gene name and 'entrez_id' containing only the entrez ID# and 'protein_expression' which contains the normalized protein expression for cancer cell lines. This dataset can be loaded into R environment with the 'depmap_proteomic' function.

Usage

`proteomic`

Format

A data frame with 24963776 rows (cell lines) and 12 variables:

- `depmap_id` Cell line foreign key (i.e. "ACH-000956")
- `cell_line` Name of cancer cell line (i.e. "22RV1_PROSTATE")
- `gene_name` HUGO symbol (e.g. "TSPAN6")
- `entrez_id` Ensembl ID (e.g. ENSG00000044574)
- `protein_expression` normalized protein expression
- `protein` protein name with TenPx (e.g. MDAMB468_BREAST_TenPx01)
- `protein_id` Protein ID (e.g. sp|P55011|S12A2_HUMAN)
- `desc` Description (e.g. S12A2_HUMAN Solute carrier family 12 member 2)
- `group_id` Group ID
- `uniprot` Uniprot ID (e.g. S12A2_HUMAN)
- `uniprot_acc` Uniprot accession ID (e.g. P55011)
- `TenPx` TenPx number (e.g. TenPx01)
Details

This data originates from the ‘protein_quant_current_normalized.csv’ file taken from the 20Q2 [Broad Institute](https://depmap.org/portal/download/) cancer dependency study. The derived dataset found in the ‘depmap’ package features the addition of a foreign key ‘depmap_id’ found in the first column of this dataset, which was added from the ‘metadata’ dataset. This dataset has been converted to a long format tibble. Variables names from the original dataset were converted to lower case, put in snake case, and abbreviated where feasible.

Change log

- 20Q2: Initial dataset consisted of a data frame with 24963776 rows (cell lines) and 12 variables
- 20Q3: no change, no further releases are scheduled at this time.
- 20Q4: no change, no further releases are scheduled at this time.
- 21Q1: no change, no further releases are scheduled at this time.
- 21Q2: no change, no further releases are scheduled at this time.
- 21Q3: no change, no further releases are scheduled at this time.
- 21Q4: no change, no further releases are scheduled at this time.
- 22Q1: no change, no further releases are scheduled at this time.
- 22Q2: no change, no further releases are scheduled at this time.

Source

DepMap, Broad (2020)

References


Examples

```r
## Not run:
depmap_proteomic()

## End(Not run)```
rnai

rnai_19Q3

Description

The ‘rnai’ dataset contains the 19Q3 cancer dependency of select cancer cell lines for genes found by RNAi gene knockdown. This dataset includes data from 17309 genes, 712 cancer cell lines, 31 primary diseases and 31 lineages. The columns of ‘rnai’ are: ‘depmap_id’, a foreign key corresponding to the cancer cell lineage, ‘cell_line’ containing the common CCLE name of the cancer cell lines, ‘gene’ containing both the HUGO gene name of the knockdown gene along with entrez ID#, ‘gene_name’ which only contains HUGO gene name, ‘entrez_id’ which contains only the entrez ID# and ‘dependency’ which contains the numerical dependency score values for each pair of genes and cell lines. This dataset can be loaded into the R environment with the ‘depmap_rnai’ function.

Usage

rnai

Format

A data frame with 12324008 rows (cell lines) and 6 variables:

- depmap_id  cancer cancer cell line foreign key (i.e. "ACH-00001")
- cell_line  CCLE name of cancer cell line (i.e. "184A1_BREAST")
- gene  HUGO symbol (e.g. "SAP25") and Entrez ID# (e.g. 100316904)
- gene_name  HUGO symbol (e.g. "SAP25")
- entrez_id  Entrez ID# (e.g. 100316904)
- dependency  numerical dependency score of a gene for a cell line

Details

This data represents the ‘D2_combined_genetic_dependency_scores’ file taken from the 19Q3 [Broad Institute](https://depmap.org/portal/download/) cancer dependency study. The derived dataset found in the ‘depmap’ package features the addition of a foreign key ‘depmap_id’ found in the first column of this dataset, which was added from the ‘metadata’ dataset. This dataset has been converted to a long format tibble. Variables names from the original dataset were converted to lower case, put in snake case, and abbreviated where feasible.

Change log

- 19Q1: Initial dataset consisted of a data frame with 12324008 rows (cell lines) and 6 variables representing 17309 genes, 711 cancer cell lines, 30 primary diseases and 31 lineages.
- 19Q2: adds 1 cell line
- 19Q3: adds 1 primary disease
- 19Q4: no change, no further releases are scheduled at this time. NOTE: as of this release, the crispr dataset replaces the rnai dataset for dependency measurements in all future releases.

- 20Q1: no change, this dataset is superceded by the CRISPR dependency screens and no further releases are scheduled.

Source
DepMap, Broad (2019)

References


Examples
```r
## Not run:
depmap_rnai()
## End(Not run)
```

RPPA

RPPA_19Q3

Description
The ‘RPPA’ dataset contains the 19Q3 CCLE Reverse Phase Protein Array (RPPA) cellular model expression data. This dataset includes data from 214 genes, 899 cancer cell lines, 28 primary diseases and 28 lineages. The columns of ‘RPPA’ are: ‘depmap_id’, a foreign key corresponding to the cancer cell lineage, ‘cell_line’ which contains the common CCLE name of the cancer cell lines, ‘gene’ which contains the knockdown gene expression, ‘antibody’ containing the name of knocked down gene and ‘expression’ containing numerical protein expression data. This dataset can be loaded into R environment with the ‘depmap_RPPA’ function.

Usage
RPPA
RPPA

Format

A data frame with 192386 rows and 4 variables:

- **depmap_id** cancer cell line foreign key (i.e. "ACH-000001")
- **cell_line** CCLE name of cancer cell line (i.e. "NIHOVCAR3_OVARY")
- **antibody** Name of antibody targeting protein (i.e. "14-3-3_beta")
- **expression** Observed expression via RPPA of protein coding genes

Details

This data represents the ‘CCLE_RPPA_20181003.csv’ file taken from the 19Q3 [Broad Institute](https://depmap.org/portal/download/) cancer dependency study. The derived dataset found in the ‘depmap’ package features the addition of a foreign key ‘depmap_id’ found in the first column of this dataset, which was added from the ‘metadata’ dataset. This dataset has been converted to a long format tibble. Variables names from the original dataset were converted to lower case, put in snake case, and abbreviated where feasible.

Change log

- 19Q1: Initial dataset consisted of a data frame with 192386 rows and 4 variables representing 214 genes, 899 cancer cell lines, 28 primary diseases and 28 lineages.
- 19Q2: no change
- 19Q3: no change, no further releases are scheduled at this time.
- 19Q4: no change, no further releases are scheduled at this time.
- 20Q1: no change, no further releases are scheduled at this time.
- 20Q2: no change, no further releases are scheduled at this time.
- 20Q3: no change, no further releases are scheduled at this time.
- 20Q4: no change, no further releases are scheduled at this time.
- 21Q1: no change, no further releases are scheduled at this time.
- 21Q2: no change, no further releases are scheduled at this time.
- 21Q3: no change, no further releases are scheduled at this time.
- 21Q4: no change, no further releases are scheduled at this time.
- 22Q1: no change, no further releases are scheduled at this time.
- 22Q2: no change, no further releases are scheduled at this time.

Source

DepMap, Broad (2019)
References


Examples

```r
## Not run:
depmap_RPPA()
## End(Not run)
```

TPM

TPM_22Q2

Description

The ‘TPM’ dataset contains the 22Q2 CCLE "Transcript Per Million" RNASeq gene expression data for protein coding genes. This dataset includes data from 19221 genes, 1406 cell lines, 33 primary diseases and 30 lineages. The columns of ‘TPM’ are: `depmap_id`, a foreign key corresponding to the cancer cell lineage, `cell_line` the common CCLE name of the cancer cell lines, `gene` containing both the HUGO gene name of the knockdown gene along with ensembl ID#, `gene_name` containing the HUGO gene name and `ensembl_id` containing only the ensembl ID# and `rna_expression` which contains the numerical protein coding gene expression change at scale (log2(TPM+1)). This dataset can be loaded into R environment with the ‘depmap_TPM’ function.

Usage

TPM

Format

A data frame with 27024726 rows (cell lines) and 6 variables:

- **depmap_id**  Cell line foreign key (i.e. "ACH-000956")
- **cell_line**  Name of cancer cell line (i.e. "22RV1_PROSTATE")
- **gene**  HUGO symbol and Ensembl ID (e.g. TSPAN6 (ENSG00000000003))
- **gene_name**  HUGO symbol (e.g. "TSPAN6")
- **ensembl_id**  Ensembl ID (e.g. ENSG00000044574)
- **rna_expression**  Log fold (log2(TPM+1)) protein expression change
Details

This data originates from the ‘CCLE_expression.csv’ file taken from the 22Q2 [Broad Institute](https://depmap.org/portal/download/) cancer dependency study. The derived dataset found in the ‘depmap’ package features the addition of a foreign key ‘depmap_id’ found in the first column of this dataset, which was added from the ‘metadata’ dataset. This dataset has been converted to a long format tibble. Variables names from the original dataset were converted to lower case, put in snake case, and abbreviated where feasible.

Change log

- 19Q1: Initial dataset consisted of a data frame with 67360300 rows (cell lines) and 6 variables representing 57820 genes, 1165 cell lines, 33 primary diseases, 32 lineages.
- 19Q2: removes 1618 genes, adds 36 cell lines, removes one primary disease and adds 1 lineage
- 19Q3: removes 37058 genes, adds 9 cell lines, removes 2 primary diseases. Now a 23164240 by 6 dataframe.
- 19Q4: 0 genes, 39 cancer cell lines, 0 primary diseases and 1 lineage
- 20Q1: adds 31 cell lines
- 20Q2: adds 34 cell lines. ‘expression’ changed to ‘rna_expression’
- 20Q3: adds 1 cell line
- 20Q4: removes 38 genes and 71 cell lines, adds 1 primary disease and 3 lineages
- 21Q1: removes 5 genes
- 21Q2: adds 3 cell lines. Additionally, a bug was fixed where the Entrez ID appeared as Ensembl_ID. This was changed for all previous versions of this dataset from 19Q3 to 21Q2
- 21Q3: removes 2 cell lines
- 21Q4: adds 12 cell lines
- 22Q1: adds 4 cell lines and 1 lineage
- 22Q2: adds 44 genes, 13 cell lines and removes 8 lineages

Source

DepMap, Broad (2020)

References


Examples

```r
## Not run:
depmap_TPM()

## End(Not run)
```
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