

Package ‘ivygapSE’

April 12, 2018

Title A SummarizedExperiment for Ivy-GAP data

Description Define a SummarizedExperiment and exploratory app for Ivy-GAP glioblastoma image, expression, and clinical data.

Version 1.0.0

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Suggests knitr

Depends R (>= 3.4.0), SummarizedExperiment

Imports shiny, survival, survminer, hwriter, plotly, ggplot2, S4Vectors, graphics, stats, utils, UpSetR

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LazyLoad yes

biocViews Transcription, Software, Visualization, Survival

RoxygenNote 6.0.1.9000

VignetteBuilder knitr

NeedsCompilation no

R topics documented:

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ivyGlimpse	<i>simple app to explore image property quantifications in relation to survival and expression</i>
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Description

simple app to explore image property quantifications in relation to survival and expression

Usage

```
ivyGlimpse()
```

Value

Side effect of starting the app only.

Examples

```
if (interactive()) print(ivyGlimpse())
```

ivySE	<i>ivySE: SummarizedExperiment for IvyGAP expression data and meta-data</i>
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Description

ivySE: SummarizedExperiment for IvyGAP expression data and metadata

Usage

```
ivySE
```

Format

SummarizedExperiment instance

Details

```
Archive: gene_expression_matrix_2014-11-25.zip
Length Date Time Name
-----
50585 03-31-2015 13:27 columns-samples.csv
86153820 10-31-2014 14:04 fpkm_table.csv
2015 11-24-2014 18:06 README.txt
1689619 10-31-2014 13:55 rows-genes.csv
-----
87896039 4 files
```

Note

Expression data retrieved from http://glioblastoma.alleninstitute.org/api/v2/well_known_file_download/305873915

Source

processed from glioblastoma.alleninstitute.org; see Note.

Examples

```
## Not run: # how it was made
ivyFpkm = read.csv("fpkm_table.csv", stringsAsFactors=FALSE,
  check.names=FALSE)
g = read.csv("rows-genes.csv", stringsAsFactors=FALSE)
library(SummarizedExperiment)
imat = data.matrix(ivyFpkm[,-1])
ivySE = SummarizedExperiment(SimpleList(fpkm=imat))
rowData(ivySE) = g
rownames(ivySE) = g$gene_symbol
col = read.csv("columns-samples.csv", stringsAsFactors=FALSE)
rownames(col) = col$rna_well_id
stopifnot(all.equal(as.character(col$rna_well_id),
  as.character(colnames(imat))))
colData(ivySE) = DataFrame(col)
colnames(ivySE) = colnames(imat)
metadata(ivySE) = list(README=readLines("README.txt"))
metadata(ivySE)$URL = "http://glioblastoma.alleninstitute.org/static/download.html"
# metadata(ivySE)$builder = readLines("build.R")
de = read.csv("tumor_details.csv", stringsAsFactors=FALSE)
metadata(ivySE)$tumorDetails = de
subbl = read.csv("sub_block_details.csv", stringsAsFactors=FALSE)
metadata(ivySE)$subBlockDetails = subbl
bamtab = read.csv("bam.csv", stringsAsFactors=FALSE)
rownames(bamtab) = as.character(bamtab$rna_well)
bamtab[colnames(ivySE),] -> bamtreo
all.equal(rownames(bamtreo), colnames(ivySE))
colData(ivySE) = cbind(colData(ivySE), bamtreo)

## End(Not run)
data(ivySE)
names(metadata(ivySE))
```

makeGeneSets

demonstration of gene set construction for ivyGlimpse app

Description

demonstration of gene set construction for ivyGlimpse app

Usage

```
makeGeneSets()
```

Value

list of gene sets with attributes facilitating dropdown construction – attr("fullTitle") is a list of strings associated with gene set elements (named list with vectors of gene symbols constituting sets of interest)

List of 4

\$ General: Ras-Raf-MEK-Erk/JNK signaling (26 genes)

: chr [1:26] "KRAS" "HRAS" "BRAF" "RAF1" ...

\$ Glioblastoma: RTK/Ras/PI3K/AKT Signaling (17 genes)

```

: chr [1:17] "EGFR" "ERBB2" "PDGFRA" "MET" ...
$ General: PI3K-AKT-mTOR signaling (17 genes)
: chr [1:17] "PIK3CA" "PIK3R1" "PIK3R2" "PTEN" ...
$ Ovarian Cancer: Putative tumor-suppressor genes in epithelial ovarian cancer (16 genes)
: chr [1:16] "DIRAS3" "RASSF1" "DLEC1" "SPARC" ...
- attr(*, "fullTitle")=List of 4
..$ glioRTK : chr "Glioblastoma: RTK/Ras/PI3K/AKT Signaling (17 genes)"
..$ pi3k : chr "General: PI3K-AKT-mTOR signaling (17 genes)"
..$ ovtumsupp: chr "Ovarian Cancer: Putative tumor-suppressor genes in epithelial ovarian cancer
(16 genes)"
..$ rasraf : chr "General: Ras-Raf-MEK-Erk/JNK signaling (26 genes)"

```

Note

Should be replaced by selections from a general catalog.

Examples

```
str(makeGeneSets())
```

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*Topic **datasets**

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