# Package 'PCAWG7'

January 25, 2022

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 ${\tt CancerTypes}$ 

Return a character vector of some common cancer types

### Description

Return a character vector of some common cancer types

### Usage

CancerTypes()

### **Examples**

CancerTypes()[1:5]

exposure

PCAWG7 SigProfiler signature assignments (numbers of mutations due to each signature in each tumor)

### Description

PCAWG7 SigProfiler signature assignments (numbers of mutations due to each signature in each tumor)

### Usage

exposure

### **Format**

A list with the elements:

**PCAWG** A list with the elements:

SBS96 Strand-agnostic single-base substitutions in trinucleotide context.

DBS78 Strand-agnostic doublet-base substitutions.

**ID** Strand-agnostic indels. These are signature assignments for the PCAWG platinum genomes.

**TCGA** A list with the elements:

SBS96 As above.

exposure.stats 3

**ID** As above. These are signature assignments for the TCGA exomes.

**other.genome** A list with the element:

**SBS96** As above. This contains signature assignments for non-TCGA genomes.

**other.exome** A list with the element:

**SBS96** As above. This contains signature assignments for non-TCGA exomes.

#### **Source**

```
Files of https://www.synapse.org/#!Synapse:syn12009743, 2019 Oct 09, populated by data-raw/sig.profiler.sures/load.package.variable.exposure.R.
```

#### **Examples**

SBS96.exposure <- exposure\$PCAWG\$SBS96

exposure.stats

Exposure statistics from the PCAWG7 paper

### **Description**

Exposure statistics from the PCAWG7 paper

### Usage

```
exposure.stats
```

#### **Format**

A list with one element, PCAWG, which has the sub-elements SBS96, DBS78, ID with statistics for the corresponding mutation types by cancer type. I.e. each element has a sub-element for each cancer type, and this element is a data.frame with one row for each signature and columns mean.of.those.present (the mean number of mutations for those tumors that have the mutation) and proportion.present (the proportion of tumors in which the signature is present).

#### **Source**

Computed from other package variables using GatherPCAWG7ExposureStatsSBS96.

```
exposure.stats$PCAWG$SBS96$`Biliary-AdenoCA`[1:3, ]
```

#### **Description**

Translate aliquot IDs (e.g. e0fccaf5-925a-41f9-b87c-cd5ee4aecb59) to "SP" IDs (e.g. SP1682)

### Usage

```
map_aliquot_ID_to_SP_ID(aliquot.ids)
```

### **Arguments**

aliquot.ids Character vector of aliquot IDs.

#### **Details**

If there are aliquot IDs that cannot be matched to any "SP" IDs, return NA with a warning.

#### Value

Character vector of corresponding "SP" IDs. If a corresponding aliquot ID cannot be found, then return return NA with a warning.

### Note

This function is mainly designed to translate the file names of PCAWG consensus callsets for SNV/Indel (https://dcc.icgc.org/api/v1/download?fn=/PCAWG/consensus\_snv\_indel/final\_consensus\_snv\_indel\_passor

### **Examples**

```
 \begin{array}{lll} aliquot.ids &<- c("e0fccaf5-925a-41f9-b87c-cd5ee4aecb59", "foo") \\ SP.ids &<- map\_aliquot\_ID\_to\_SP\_ID(aliquot.ids) \\ \end{array}
```

```
map_SP_ID_to_tumor_type
```

Given PCAWG "SP" IDs (e.g. SP123958) return either the "full" IDs (Kidney-ChRCC::SP123958) or just the tumor type (Kidney-ChRCC)

### **Description**

Given PCAWG "SP" IDs (e.g. SP123958) return either the "full" IDs (Kidney-ChRCC::SP123958) or just the tumor type (Kidney-ChRCC)

#### Usage

```
map_SP_ID_to_tumor_type(SP.IDs, merge = TRUE)
```

PCAWG.sample.id 5

### **Arguments**

SP. IDs A character vector of PCAWG "SP" IDs.

merge If TRUE return a parallel vector of <tumor\_type>::<SP\_ID>; otherwise just <tu-

mor\_type>.

#### **Details**

Fails with an "subscript out of bounds" error if any of the elements of SP. IDs is unknown.

### **Examples**

```
map_SP_ID_to_tumor_type(c("SP123958", "SP43633"))
map_SP_ID_to_tumor_type(c("SP123958", "SP43633"), merge = FALSE)
```

PCAWG.sample.id

Vectors of the PCAWG tumor\_wgs\_icgc\_specimen\_ids

### **Description**

Note that the PCAWG7 spectra catalogs have 2 sample ids that were blacklisted after the mutational signature analysis was underway. The blacklisted samples are SP116419 and SP116883, which are in PCAWG.sample.id\$black.

#### Usage

```
PCAWG.sample.id
```

### **Format**

A list with the elements:

```
white Whitelisted IDsgrey Greylisted IDsblack Blacklisted IDs
```

#### Source

```
https://dcc.icgc.org/api/v1/download?fn=/PCAWG/data\_releases/latest/release\_may2016. v1.4.with\_consensus\_calls.tsv, 2019 Oct 09
```

```
PCAWG.white.ids <- PCAWG.sample.id$white
```

PCAWG7

PCAWG.sample.sheet

PCAWG sample sheet which contains various sample information

#### **Description**

PCAWG sample sheet which contains various sample information

### Usage

```
PCAWG.sample.sheet
```

#### **Format**

A data table with the following columns:

- · donor\_unique\_id
- donor\_wgs\_exclusion\_white\_gray
- submitter\_donor\_id
- icgc\_donor\_id
- dcc\_project\_code
- · aliquot\_id
- submitter\_specimen\_id
- icgc\_sample\_id
- dcc\_specimen\_type
- library\_strategy

#### **Source**

```
https://dcc.icgc.org/api/v1/download?fn=/PCAWG/data_releases/latest/pcawg_sample_sheet.v1.4.2016-09-14.tsv, 2019 Oct 15
```

### **Examples**

```
aliquot.ids <- PCAWG.sample.sheet$aliquot_id</pre>
```

PCAWG7

PCAWG7: A package of data from paper 'Repertoire of Mutational Signatures in Human Cancer'

### Description

This is a data package with 2 main package variables: exposure and spectra.

#### **Details**

There are also PDF plots of the signatures in data-raw/plots/.

There are also several functions for handling PCAWG identifiers:

```
* map_SP_ID_to_tumor_type
```

- \* SampleIDToCancerType
- \* SplitPCAWGMatrixByTumorType
- \* SplitMatrixBySampleType

The reference for the data is

Alexandrov, L.B., Kim, J., Haradhvala, N.J. et al. The repertoire of mutational signatures in human cancer. Nature 578, 94-101 (2020). doi: 10.1038/s4158602019433.

SampleIDToCancerType Split out the cancer type from the sample ID for PCAWG IDs

### Description

Split out the cancer type from the sample ID for PCAWG IDs

### Usage

SampleIDToCancerType(PCAWGID)

### Arguments

PCAWGID

A character vector of PCAWG IDs of the form <cancer.type>::<sample.id>.

### Value

A character vector parallel to PCAWGID containing only the <cancer.type> strings.

```
cancer.type <- SampleIDToCancerType("Biliary-AdenoCA::SP117655")</pre>
```

<sup>\*</sup> map\_aliquot\_ID\_to\_SP\_ID

spectra

PCAWG7 mutational spectra (catalogs)

#### **Description**

PCAWG7 mutational spectra (catalogs)

#### Usage

spectra

#### **Format**

A list with the elements:

SBS96 Deprecated.

DBS78 Deprecated.

**PCAWG** A list with the elements:

SBS96 Strand-agnostic single-base substitutions in trinucleotide context.

SBS192 Single-base substitutions in transcripts based on the sense strand.

SBS1536 Strand-agnostic single-base substitutions in pentanucleotide context.

DBS78 Strand-agnostic doublet-base substitutions.

**ID** Strand-agnostic indels.

**TCGA** A list with the same elements as the PCAWG element.

**other.genome** A list with the same elements as the PCAWG element but with ID omitted.

other.exome A list with the same elements as the PCAWG element but with ID omitted.

#### **Source**

Files below https://www.synapse.org/#!Synapse:syn11801889, 2019 Oct 09. Populated by data-raw/spectra/load.package.variable.specra.R.

### **Examples**

```
SBS96.spectra <- spectra$PCAWG$SBS96
```

 ${\tt SplitMatrixBySampleType}$ 

Split an exposure matrix or spectrum matrix into a list of matrices, each for a single sample type

### **Description**

Split an exposure matrix or spectrum matrix into a list of matrices, each for a single sample type

#### Usage

```
SplitMatrixBySampleType(M, sample.type)
```

### **Arguments**

M A numerical matrix or data frame or ICAMS catalog in which columns are sam-

ples (e.g. tumors) and rows are either mutational signatures (for exposures) or mutation types (for spectra), and, each element is the number of mutations due

to a given mutational signature or mutation type in a single sample

sample.type A character or numeric vector, each element of which indicates a particular sam-

ple type.

#### Value

Invisibly, the list of exposure or spectrum matrices created by splitting M by sample.type.

### **Examples**

```
ff <- matrix(1, nrow=3, ncol = 2)
colnames(ff) <- c("sample1", "sample2")
xx <- SplitMatrixBySampleType(ff, c("sample.type.x", "sample.type.y"))
xx</pre>
```

SplitPCAWGMatrixByTumorType

Extract tumor type from column names and return the input matrix split by tumor type based on the PCAWG <tumor\_type>::<sample\_id> convention

### **Description**

Extract tumor type from column names and return the input matrix split by tumor type based on the PCAWG <tumor\_type>::<sample\_id> convention

### Usage

```
SplitPCAWGMatrixByTumorType(M)
```

### **Arguments**

М

A numerical matrix or data frame or ICAMS catalog in which columns are samples (e.g. tumors) and rows are either mutational signatures (for exposures) or mutation types (for spectra), and each element is the number of mutations due to a given mutational signature or mutation type in a single sample. The column names must be of the the form <cancer.type>::<sample.ID>.

#### Value

Invisibly, the list of exposure matrices or ICAMS catalogs created by splitting matrix by the tumor type encoded in the column names.

```
mm <- SplitPCAWGMatrixByTumorType(spectra$PCAWG$DBS78)</pre>
```

### Description

Translate TCGA (The Cancer Genome Atlas) IDs to ICGC (International Cancer Genome Consortium) IDs

### Usage

```
TCGA_ID_to_ICGC_ID(tcga.ids)
```

### Arguments

tcga.ids Character vector of TCGA IDs.

#### **Details**

If there are TCGA IDs that cannot be matched to any ICGC IDs, return NA with a warning.

### Value

Character vector of corresponding ICGC IDs. If a corresponding ICGC ID cannot be found, then return NA with a warning.

```
tcga.ids <- c("TCGA-AA-A01V", "foo", "TCGA-CA-6717", "bar")
icgc.ids <- TCGA_ID_to_ICGC_ID(tcga.ids)
icgc.ids <- icgc.ids[nzchar(icgc.ids)]</pre>
```

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