# Package ‘SomaticCancerAlterations’

March 23, 2017

**Type** Package  
**Title** Somatic Cancer Alterations  
**Version** 1.10.0  
**Author** Julian Gehring (EMBL Heidelberg)  
**Maintainer** Julian Gehring <jg-bioc@gmx.com>  
**Imports** GenomicRanges, exomeCopy, stringr, IRanges, S4Vectors  
**Depends** R (>= 3.0.0)  
**Suggests** testthat, ggbio, ggplot2, knitr  
**VignetteBuilder** knitr  
**Description** Collection of somatic cancer alteration datasets  
**License** GPL-3  
**LazyLoad** yes  
**biocViews** ExperimentData, Genome, CancerData, Project1000genomes, NCI  
**Encoding** UTF-8  
**NeedsCompilation** no

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**Description**

A collection of studies with somatic cancer alterations.
Details

The `SomaticCancerAlterations` package provides a collection of the mutational calls for different cancer studies, aiming for a tight integration with R and Bioconductor. At the moment, this covers somatic single nucleotide variants (SNVs) and indels for several TCGA studies. Over time, this collection will be updated to include additional studies.

The Cancer Genome Atlas (TCGA) is a consortium effort to analyze a variety of tumor types at multiple levels, including gene expression, methylation, copy number alterations and somatic mutations [http://cancergenome.nih.gov]. Most of this data is publically available, and offers a rich resource in understanding cancer-related data sets and experiments [https://wiki.nci.nih.gov/display/TCGA/TCGA+Home].

Author(s)

Julian Gehring (EMBL Heidelberg), with suggestions and contributions from Bernd Fischer

Maintainer: Julian Gehring <julian.gehring@embl.de>

References

https://tcga-data.nci.nih.gov/tcga/
https://tcga-data.nci.nih.gov/docs/publications/

See Also

SomaticCancerAlterations-functions
SomaticCancerAlterations-data

Description

Data sets with studies of somatic cancer alterations.

Details

Currently, the data sets include somatic mutations (SNVs and indels) of the publically available TCGA studies. Further studies will be included in the future.

For details on the TCGA mutation calls, please see the TCGA documentation https://wiki.nci.nih.gov/display/TCGA/TCGA+Data+Primer. Please note that the mutational calling was performed by different centers. The original '*.maf' files can be obtained from the TCGA data portal https://tcga-data.nci.nih.gov/tcgafiles/ftp_auth/distro_ftpusers/anonymous/tumor/.

After importing, the data is automatically processed to improve consistency across studies and remove common artifacts in the annotation. The most obvious changes are:

- Only studies cleared for unrestricted usage are included (see http://cancergenome.nih.gov/publications/publicationsguidelines).
- Only studies processed by the Broad Instutute are considered at the moment.
• All variants located on the mitochondrium are mapped to the sequence identifier 'MT' (note that the original files contain both 'MT' and 'M' mixed).
• All locations are now consistent with the 1000genomes reference sequence (NCBI37 coordinates).

About the TCGA data:
“All data generated by The Cancer Genome Atlas (TCGA) Research Network are made open to the public through the Data Coordinating Center and the TCGA Data Portal.” For details on the usage of the data, please have a look at http://cancergenome.nih.gov/abouttcga/policies/publicationguidelines.

Value
A 'GRanges' object for each study. Each row corresponds to a somatic variant reported in the respective study, with the coordinates referring to the location on the reference genome.

References
https://tcga-data.nci.nih.gov/tcga/

See Also
SomaticCancerAlterations-functions
SomaticCancerAlterations-package

Examples
all_datasets = scaListDatasets()
grl = scaLoadDatasets(all_datasets[1])
Arguments

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<tr>
<td>names</td>
<td>Character vector with data set name(s) to load. The names are the same as returned from <code>scaListDatasets()</code></td>
</tr>
<tr>
<td>merge</td>
<td>Logical. Should the results be a merged <code>GRanges</code> object with data of all studies (TRUE), or a <code>GenomicRangesList</code> with one list element per dataset (FALSE [default]).</td>
</tr>
<tr>
<td>x</td>
<td>GRanges object</td>
</tr>
<tr>
<td>binSize</td>
<td>Integer with the bin size used for computing the density of variants.</td>
</tr>
<tr>
<td>chrs</td>
<td>Character vector with the chromosomes/seqlevels to include (by default, all chromosomes are used.)</td>
</tr>
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Details

The `sca_load_dataset` function makes it easy to load the data of a study directly into a variable. The `mutationDensity` function computes the density of events in a binned-manner.

Value

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<td>scaMetadata</td>
<td>A data frame summarizing the available data sets, with rows corresponding to data sets.</td>
</tr>
<tr>
<td>scaListDatasets</td>
<td>A character vector with all available study names.</td>
</tr>
<tr>
<td>sca_load_dataset</td>
<td>A GenomicRangesList or GRanges objects, depending on the <code>merge</code> argument.</td>
</tr>
<tr>
<td>mutationDensity</td>
<td>A GRanges object, with ranges corresponding to bins along the genome. The columns 'counts' and 'density' refer to the number and the fraction (given per 1Mbp) of alterations, respectively.</td>
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References

https://tcga-data.nci.nih.gov/tcga/
https://wiki.nci.nih.gov/display/TCGA/TCGA+Data+Primer
https://tcga-data.nci.nih.gov/tcgafiles/ftp_auth/distro_ftpusers/anonymous/tumor/

See Also

SomaticCancerAlterations-data

Examples

```r
meta_data = scaMetadata()
head(meta_data)

all_datasets = scaListDatasets()
head(all_datasets)

grl = scaLoadDatasets(all_datasets[1], merge = FALSE)
gr = scaLoadDatasets(all_datasets[1], merge = TRUE)
```
Description

Functions to help with common analysis tasks.

Usage

ncbi2hg(x)

hg2ncbi(x)

seqchr(x)

Arguments

x  A GRanges object

Details

These functions provide useful wrappers for converting between genomic coordinate systems and classes.

Value

ncbi2hg, hg2ncbi

A GRanges object.

seqchr  A character vector with the 'seqnames'.

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