Package 'ICAMS'

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Title In-depth Characterization and Analysis of Mutational Signatures

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```
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Description Analysis and visualization of experimentally elucidated mutational
      signatures -- the kind of analysis and visualization in Boot et al.,
      "In-depth characterization of the cisplatin mutational signature in
      human cell lines and in esophageal and liver tumors", 2018,
      <a href="https://genome.cshlp.org/content/28/5/654.short">https://genome.cshlp.org/content/28/5/654.short</a>. ICAMS has functions
      to read in variant call files (VCFs) and to collate the corresponding
      catalogs of mutational spectra and to analyze and plot catalogs of
      mutational spectra and signatures. Handles both "counts-based" and
      "density-based" catalogs of mutational spectra or signatures.
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2 all.abundance

R topics documented:

	all.abundance	2
	as.catalog	3
	CatalogRowOrder	3
	CollapseCatalog	4
	FindDelMH	4
	GetVAF	6
	ICAMS	7
	MutectVCFFilesToCatalog	10
	MutectVCFFilesToCatalogAndPlotToPdf	10
	PlotCatalog	11
	PlotCatalogToPdf	12
		13
	ReadAndSplitStrelkaSBSVCFs	14
	ReadCatalog	14
	ReadStrelkaIDVCFs	15
	revc	16
	StrelkaIDVCFFilesToCatalog	16
	StrelkaIDVCFFilesToCatalogAndPlotToPdf	17
	StrelkaSBSVCFFilesToCatalog	17
	StrelkaSBSVCFFilesToCatalogAndPlotToPdf	18
	TranscriptRanges	19
	TransformCatalog	20
	VCFsToDBSCatalogs	21
	VCFsToIDCatalogs	21
	VCFsToSBSCatalogs	22
	WriteCatalog	22
Index		24
all.a	abundance K-mer abundances.	

Description

An R list with one element each for BSgenome.Hsapiens.1000genomes.hs37d5, and BSgenome.Hsapiens.UCSC.hg38. Each element is in turn a sub-list keyed by exome, transcript, and genome. Each element of the sub list is keyed by the number of rows in the catalog class (as a string, e.g. "78", not 78). The keys are: 78 (DBS78Catalog), 96 (SBS96Catalog), 136 (DBS136Catalog), 144 (DBS144Catalog), 192 (SBS192Catalog), and 1536 (SBS1536Catalog). So, for example to get the exome abundances for SBS96 catalogs for BSgenome.Hsapiens.UCSC.hg38 exomes one would reference

all.abundance[["BSgenome.Hsapiens.UCSC.hg38"]][["exome"]]["96"] or all.abundance\$BSgenome.Hsapiens.UCSC.hg38\$exome\$"96". The value of the abundance is an integer vector with the K-mers as names and the values being the number of the K-mer.

Usage

all.abundance

as.catalog 3

Format

An object of class list of length 2.

as.catalog	.frame.
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Description

Create a catalog from a numeric matrix or numeric data. frame.

Usage

```
as.catalog(object, ref.genome = NULL, region = "unknown",
  catalog.type = "counts", abundance = NULL)
```

Arguments

object A numeric matrix or numeric data. frame. This object must have rownames to

denote the mutation types. See CatalogRowOrder for more details.

ref.genome A ref.genome argument as described in ICAMS.

region A character string designating a region, one of genome, transcript, exome,

unknown; see ICAMS.

catalog.type One of "counts", "density", "counts.signature", "density.signature".

abundance Optional, only needed when ref. genome is not one of the two human reference

genomes included in ICAMS. The abundance should contain the counts of dif-

ferent source sequences for mutations.

See ICAMS:::abundance.3bp.exome.unstranded.GRCh37 for an example.

Value

A catalog as described in ICAMS.

CatalogRowOrder Standard order of row names in a catalog.	
---	--

Description

This data is designed for those who need to create their own catalogs from formats not supported by this package. The rownames denote the mutation types. For example, for SBS96 catalogs, the rowname AGAT represents a mutation from AGA > ATA.

Usage

```
catalog.row.order
```

Format

A list of character vectors indicating the standard orders of row names in catalogs.

4 FindDelMH

Note

In ID (insertion and deletion) catalogs, deletion repeat sizes range from 0 to 5+, but for plotting and end-user documentation deletion repeat sizes range from 1 to 6+.

CollapseCatalog

"Collapse" a catalog.

Description

- 1. Take a mutational spectrum or signature catalog that is based on a fined-grained set of features (for example, single-nucleotide substitutions in the context of the preceding and following 2 bases).
- 2. Collapse it to a catalog based on a coarser-grained set of features (for example, single-nucleotide substitutions in the context of the immediately preceding and following bases).

Collapse192CatalogTo96 Collapse an SBS 192 catalog to an SBS 96 catalog.

Collapse1536CatalogTo96 Collapse an SBS 1536 catalog to an SBS 96 catalog.

Collapse144CatalogTo78 Collapse a DBS 144 catalog to a DBS 78 catalog.

Usage

```
Collapse192CatalogTo96(catalog)
Collapse1536CatalogTo96(catalog)
Collapse144CatalogTo78(catalog)
```

Arguments

catalog

A catalog as defined in ICAMS.

Value

A catalog as defined in ICAMS.

FindDelMH

Return the length of microhomology at a deletion.

Description

Return the length of microhomology at a deletion.

Usage

```
FindDelMH(context, deleted.seq, pos, trace = 0)
```

FindDelMH 5

Arguments

context The deleted sequence plus ample surrounding sequence on each side (at least as

long as del. sequence).

deleted.seq The deleted sequence in context.

pos The position of del. sequence in context.

trace If > 0, cat various messages.

Details

This function is primarily for internal use, but we export it to document the underlying logic.

Example:

GGCTAGTT aligned to GGCTAGAACTAGTT with a deletion represented as:

```
GGCTAGAACTAGTT
GG-----CTAGTT GGCTAGTT GG[CTAGAA]CTAGTT
```

Presumed repair mechanism leading to this:

GGCTAGAACTAGTT CCGATCTTGATCAA

=>

GGCTAG TT
CC GATCAA

=>

GGCTAGTT CCGATCAA

Variant-caller software can represent the same deletion in several different, but completely equivalent, ways.

```
GGCTAGTT GGCTAGTT GGC[TAGAAC]TAGTT

* --- * ---

GGCT-----AGTT GGCTAGTT GGCT[AGAACT]AGTT

** -- ** --

GGCTA-----GTT GGCTAGTT GGCTA[GAACTA]GTT

*** - *** -

GGCTAG----TT GGCTAGTT GGCTAG[AACTAG]TT

*** ***
```

6 GetVAF

A deletion in a *repeat* can also be represented in several different ways. A deletion in a repeat is abstractly equivalent to microhomology that spans the entire deleted sequence. For example;

```
GACTAGCTAGTT

GACTAGTT GACTAGTT GACTA[GCTA]GTT

*** -*** -

is really a repeat

GACTAG---TT GACTAGTT GACTAG[CTAG]TT

**** ----

GACT---AGTT GACTAGTT GACT[AGCT]AGTT

** --** --
```

This function only flags this case with a -1 return; it does not figure out the repeat extent.

This function finds:

- 1. The maximum match of undeleted sequence to the left of the deletion that is identical to the right end of the deleted sequence, and
- 2. The maximum match of undeleted sequence to the right of the deletion that is identical to the left end of the deleted sequence.

The microhomology sequence is the concatenation of items (1) and (2).

Value

The length of the maximum microhomology of del. sequence in context.

GetVAF

Extract the VAFs (variant allele frequencies) from a VCF file.

Description

Extract the VAFs (variant allele frequencies) from a VCF file.

Usage

```
GetStrelkaVAF(vcf)
GetMutectVAF(vcf)
```

Arguments

vcf

Said VCF as a data.frame.

Value

A vector of VAFs, one for each row of vcf.

ICAMS 7

ICAMS	ICAMS: In-depth Characterization and Analysis of Mutational Signatures

Description

Analysis and visualization of experimentally elucidated mutational "In-depth characterization of the cisplatin mutational signature in human cell lines and in esophageal and liver tumors", *Genome Research* 2018, https://genome.cshlp.org/content/28/5/654.short. ICAMS has functions to read in variant call files (VCFs) and to collate the corresponding catalogs of mutational spectra and to analyze and plot catalogs of mutational spectra and signatures. Handles both "counts-based" and "density-based" catalogs of mutational spectra or signatures. ICAMS can read in VCFs generated by Strelka or Mutect, and collate the mutations into "catalogs" of mutational spectra. ICAMS can create and plot catalogs of mutational spectra or signatures for single base substitutions (SBS), double base substitutions (DBS), and small insertions and deletions (ID). It can also read and write these catalogs.

Catalogs

A key data type in ICAMS is a "catalog" of mutation counts, of mutation densities, or of mutational signatures.

Catalogs are R S3 objects of class matrix and one of several additional classes that specify the types of the mutations represented in the catalog (e.g. SBS96, ID, etc, ...). The possible additional classes are one of SBS96Catalog, SBS192Catalog, SBS1536Catalog, DBS78Catalog, DBS144Catalog, DBS136Catalog, IndelCatalog. as.catalog is the main constructor.

Conceptually, a catalog has one of the following types, which are indicated in the attribute catalog. type:

- 1. Matrix of mutation counts (one column per sample), representing (counts-based) mutational spectra (catalog.type = "counts").
- 2. Matrix of mutation densities, i.e. mutations per occurrences of source sequences (one column per sample), representing (density-based) mutational spectra (catalog.type = "density").
- 3. Matrix of mutational signatures, which are similar to spectra. However where spectra consist of counts or densities of mutations in each mutation class (e.g. ACA > AAA, ACA > AGA, ACA > ATA, ACC > AAC, ...), signatures consist of the proportions of mutations in each class (with all the proportions summing to 1). A mutational signature can be based on either:
 - mutation counts (a "counts-based mutational signature", catalog.type = "counts.signature"), or
 - mutation densities (a "density-based mutational signature", catalog.type = "density.signature").

Catalogs also have the attribute abundance, which contains the counts of different source sequences for mutations. For example, for SBSs in trinucleotide context, the abundances would be the counts of each trinucleotide in the human genome, exome, or in the transcribed region of the genome. See below under TransformCatalog for more information. Abundances logically depend on the species in question and on the part of the genome being analyzed.

In ICAMS functions these can sometimes be inferred from catalog class attribute and function arguments region, ref.genome, and catalog.type. Otherwise they can be provided as an abundance argument. See all.abundance for examples.

Possible values for region are the strings genome, transcript, exome, and unknown; transcript includes entire transcribed regions, i.e. the introns as well as the exons.

8 ICAMS

If you need to create a catalog from a source other than this package (i.e. other than with ReadCatalog or StrelkaSBSVCFFilesToCatalog, MutectVCFFilesToCatalog, etc.), then use as.catalog.

Creating catalogs from variant call files (VCF files)

- 1. StrelkaSBSVCFFilesToCatalog creates 3 SBS catalogs (96, 192, 1536) and 3 DBS catalogs (78, 136, 144) from the Strelka SBS VCFs.
- 2. StrelkaIDVCFFilesToCatalog creates ID (indel) catalog from the Strelka ID VCFs.
- 3. MutectVCFFilesToCatalog creates 3 SBS catalogs (96, 192, 1536), 3 DBS catalogs (78, 136, 144) and ID (indel) catalog from the Mutect VCFs.

Plotting catalogs

The PlotCatalog functions plot mutational spectra for **one** sample or plot **one** mutational signature.

The PlotCatalogToPdf functions plot catalogs of mutational spectra or of mutational signatures to a PDF file.

Wrapper functions to create catalogs from VCFs and plot the catalogs to PDF files

- 1. StrelkaSBSVCFFilesToCatalogAndPlotToPdf creates all type of SBS and DBS catalogs from Strelka SBS VCFs and plots the catalogs.
- StrelkaIDVCFFilesToCatalog creates an ID (indel) catalog from Strelka ID VCFs and plot it.
- 3. MutectVCFFilesToCatalog creates all types of SBS, DBS, and ID catalogs from Mutect VCFs and plots the catalogs.

The ref.genome (reference genome) argument

Many functions take the argument ref.genome.

In order to create a mutational spectrum catalog, ICAMS needs to know the sequence context of the mutations in the VCF file. For this, ICAMS needs the reference genome sequence that matches the VCF file. The ref.genome argument provides this.

ref.genome can be either

- 1. A variable from the Bioconductor BSgenome package that contains a particular reference genome, for example BSgenome. Hsapiens. 1000genomes. hs37d5.
- 2. The strings "hg38" or "GRCh38" are shorthand for BSgenome. Hsapiens. UCSC. hg38, and the strings "hg19" or "GRCh37" are shorthand for BSgenome. Hsapiens. 1000genomes. hs37d5.

The Bioconductor BSgenome package and two human genomes genomes from BSgenome are "imported" by ICAMS and therefore should be installed when ICAMS is installed. The two genomes that are installed as dependencies are:

- BSgenome. Hsapiens. 1000genomes. hs37d5
- BSgenome.Hsapiens.UCSC.hg38

Any other needed reference genomes must be installed separately by the user. Use available.genomes() to get the list of available genomes. Further instructions are at https://bioconductor.org/packages/release/bioc/html/BSgenome.html.

Use of ICAMS with other reference genomes is restricted to catalog. type of counts or counts. signature unless the user also creates the necessary abundance vectors. See all.abundance.

ICAMS 9

Writing catalogs to files

The WriteCatalog functions write a catalog to a file.

Reading catalogs

The ReadCatalog functions read a file that contains a catalog in standardized format.

Transforming catalogs

The TransformCatalog function transforms catalogs of mutational spectra or signatures to account for differing abundances of the source sequence of the mutations in the genome.

For example, mutations from ACG are much rarer in the human genome than mutations from ACC simply because CG dinucleotides are rare in the genome. Consequently, there are two possible representations of mutational spectra or signatures. One representation is based on mutation counts as observed in a given genome or exome, and this approach is widely used, as, for example, at https://cancer.sanger.ac.uk/cosmic/signatures, which presents signatures based on observed mutation counts in the human genome. We call these "counts-based spectra" or "counts-based signatures".

Alternatively, mutational spectra or signatures can be represented as mutations per source sequence, for example the number of ACT > AGT mutations occurring at all ACT 3-mers in a genome. We call these "density-based spectra" or "density-based signatures".

This function can also transform spectra based on observed genome-wide counts to "density"-based catalogs. In density-based catalogs mutations are expressed as mutations per source sequences. For example, a density-based catalog represents the proportion of ACCs mutated to ATCs, the proportion of ACGs mutated to ATGs, etc. This is different from counts-based mutational spectra catalogs, which contain the number of ACC > ATC mutations, the number of ACG > ATG mutations, etc.

This function can also transform observed-count based spectra or signatures from genome to exome based counts, or between different species (since the abundances of source sequences vary between genome and exome and between species).

Collapsing catalogs

The CollapseCatalog functions

- 1. Take a mutational spectrum or signature catalog that is based on a fined-grained set of features (for example, single-nucleotide substitutions in the context of the preceding and following 2 bases).
- 2. Collapse it to a catalog based on a coarser-grained set of features (for example, single-nucleotide substitutions in the context of the immediately preceding and following bases).

Data

- 1. CatalogRowOrder Standard order of rownames in a catalog. The rownames of encode the type of each mutation. The rownames denote the mutation types. For example, for SBS96 catalogs, the rowname AGAT represents a mutation from AGA > ATA.
- 2. TranscriptRanges Transcript ranges and strand information for a particular reference genome.

MutectVCFFilesToCatalog

Create SBS, DBS and Indel catalogs from Mutect VCF files

Description

Create 3 SBS catalogs (96, 192, 1536), 3 DBS catalogs (78, 136, 144) and Indel catalog from the Mutect VCFs specified by files

Usage

```
MutectVCFFilesToCatalog(files, ref.genome, trans.ranges = NULL,
  region = "unknown")
```

Arguments

files Character vector of file paths to the Mutect VCF files.

ref.genome A ref.genome argument as described in ICAMS.

trans.ranges a data.table which contains transcript range and strand information. Please

refer to TranscriptRanges for more details.

region A character string designating a genomic region; see as.catalog and ICAMS.

Details

This function calls VCFsToSBSCatalogs, VCFsToDBSCatalogs and VCFsToIDCatalogs

Value

A list of 3 SBS catalogs (one each for 96, 192, and 1536), 3 DBS catalogs (one each for 78, 136, and 144) and ID catalog. If trans.ranges = NULL, SBS 192 and DBS 144 catalog will not be generated. Each catalog has attributes added. See as .catalog for more details.

Note

SBS 192 and DBS 144 catalogs include only mutations in transcribed regions.

 ${\tt MutectVCFFilesToCatalogAndPlotToPdf}$

Create SBS, DBS and Indel catalogs from Mutect VCF files and plot them to PDF

Description

Create 3 SBS catalogs (96, 192, 1536), 3 DBS catalogs (78, 136, 144) and Indel catalog from the Mutect VCFs specified by files and plot them to PDF

Usage

```
MutectVCFFilesToCatalogAndPlotToPdf(files, ref.genome,
   trans.ranges = NULL, region = "unknown", output.file)
```

PlotCatalog 11

Arguments

files Character vector of file paths to the Mutect VCF files.

ref.genome A ref.genome argument as described in ICAMS.

trans.ranges a data.table which contains transcript range and strand information. Please refer to TranscriptRanges for more details.

region A character string designating a genomic region; see as . catalog and ICAMS.

output.file The name of the PDF file to be produced.

Details

This function calls MutectVCFFilesToCatalog and PlotCatalogToPdf

Value

A list of 3 SBS catalogs (one each for 96, 192, and 1536), 3 DBS catalogs (one each for 78, 136, and 144), Indel catalog and their graphs plotted to PDF with specified file name. If trans.ranges = NULL, SBS 192 and DBS 144 catalog will not be generated and plotted. Each catalog has attributes added. See as.catalog for more details.

Note

SBS 192 and DBS 144 catalogs include only mutations in transcribed regions.

PlotCatalog Plot one spectrum or signature.
--

Description

Plot the spectrum of **one** sample or plot **one** signature. The type of graph is based on one attribute("catalog.type") of the input catalog. You can first use TransformCatalog to get different types of catalog and then do the plotting.

Usage

```
PlotCatalog(catalog, plot.SBS12 = NULL, cex = NULL, grid = NULL,
    upper = NULL, xlabels = NULL)
```

Arguments

catalog	A catalog as defined in ICAMS with attributes added. See as.catalog for more details.
plot.SBS12	Only meaningful for class SBS192Catalog; if TRUE, generate an abbreviated plot of only SBS without context, i.e. C>A, C>G, C>T, T>A, T>C, T>G each on transcribed and untranscribed strands, rather than SBS in trinucleotide context, e.g. ACA > AAA, ACA > AGA,, TCT > TAT,
cex	A numerical value giving the amount by which mutation class labels, mutation counts(if it exists), y axis and its labels, x axis labels and its annotations(if it exists), sample name and legend(if it exists) should be magnified relative to the default. Only implemented for SBS96Catalog, SBS192Catalog and DBS144Catalog.

12 PlotCatalogToPdf

grid	A logical value indicating whether to draw grid lines. Only implemented for SBS96Catalog.
upper	A logical value indicating whether to draw horizontal lines and the names of major mutation class on top of graph. Only implemented for SBS96Catalog.
xlabels	A logical value indicating whether to draw x axis labels. Only implemented for SBS96Catalog.

Value

```
invisible(TRUE)
```

Note

The sizes of repeats involved in deletions range from 0 to 5+ in the mutational-spectra and signature catalog rownames, but for plotting and end-user documentation they range from 1 to 6+.

tCatalogToPdf Plot catalog to a PDF file.

Description

Plot catalog to a PDF file. The type of graph is based on one attribute("catalog.type") of the input catalog. You can first use TransformCatalog to get different types of catalog and then do the plotting.

Usage

```
PlotCatalogToPdf(catalog, file, plot.SBS12 = NULL, cex = NULL,
   grid = NULL, upper = NULL, xlabels = NULL)
```

Arguments

catalog	A catalog as defined in ICAMS with attributes added. See as.catalog for more details.
file	The name of the PDF file to be produced.
plot.SBS12	Only meaningful for class SBS192Catalog; if TRUE, generate an abbreviated plot of only SBS without context, i.e. C>A, C>G, C>T, T>A, T>C, T>G each on transcribed and untranscribed strands, rather than SBS in trinucleotide context, e.g. ACA > AAA, ACA > AGA,, TCT > TAT, There are 12 bars in the graph.
cex	A numerical value giving the amount by which mutation class labels, mutation counts(if it exists), y axis and its labels, x axis labels and its annotations(if it exists), sample name and legend(if it exists) should be magnified relative to the default. Only implemented for SBS96Catalog, SBS192Catalog and DBS144Catalog.
grid	A logical value indicating whether to draw grid lines. Only implemented for SBS96Catalog.
upper	A logical value indicating whether to draw horizontal lines and the names of major mutation class on top of graph. Only implemented for SBS96Catalog.
xlabels	A logical value indicating whether to draw x axis labels. Only implemented for SBS96Catalog.

Value

invisible(TRUE)

Note

The sizes of repeats involved in deletions range from 0 to 5+ in the mutational-spectra and signature catalog rownames, but for plotting and end-user documentation they range from 1 to 6+.

ReadAndSplitMutectVCFs

Read and split Mutect VCF files.

Description

Read and split Mutect VCF files.

Usage

ReadAndSplitMutectVCFs(files)

Arguments

files

Character vector of file paths to the Mutect VCF files.

Value

A list with 3 in-memory VCFs and two left-over VCF-like data frames with rows that were not incorporated into the first 3 VCFs, as follows:

- 1. SBS VCF with only single base substitutions.
- 2. DBS VCF with only doublet base substitutions as called by Mutect.
- 3. ID VCF with only small insertions and deletions.
- 4. other.subs VCF like data.frame with rows for coordinate substitutions involving 3 or more nucleotides, e.g. ACT > TGA or AACT > GGTA.
- 5. multiple.alternative.alleles VCF like data.frame with rows for variants with multiple alternative alleles, for example ACT mutated to both AGT and ACT at the same position.

See Also

MutectVCFFilesToCatalog

14 ReadCatalog

ReadAndSplitStrelkaSBSVCFs

Read and split Strelka SBS VCF files.

Description

Read and split Strelka SBS VCF files.

Usage

ReadAndSplitStrelkaSBSVCFs(files)

Arguments

files Character vector of file paths to the Strelka SBS VCF files.

Value

A list of 3 in-memory objects as follows:

- 1. SBS.vcfs List of data.frames of pure SBS mutations no DBS or 3+BS mutations.
- 2. DBS.vcfs List of data.frames of pure DBS mutations no SBS or 3+BS mutations.
- 3. ThreePlus List of data.tables with the key CHROM, LOW.POS, HIGH.POS. containing rows that that in the input that did not represent SBSs or DBSs.

See Also

StrelkaSBSVCFFilesToCatalog

ReadCatalog $Read\ catalog.$

Description

Read a catalog in standardized format from path.

Usage

```
ReadCatalog(file, ref.genome, region, catalog.type, strict = TRUE)
```

Arguments

file	Path to a catalog on disk in the standardized format.
ref.genome	A ref. genome argument as described in ICAMS.
region	region A character string designating a genomic region; see as $\mbox{\it catalog}$ and $\mbox{\it ICAMS}.$
catalog.type	One of "counts", "density", "counts.signature", "density.signature".
strict	If TRUE, do additional checks on the input, and stop if the checks fail.

ReadStrelkaIDVCFs 15

Details

See also WriteCatalog

Value

A catalog as an S3 object; see as.catalog.

Note

In ID (insertion and deletion) catalogs, deletion repeat sizes range from 0 to 5+, but for plotting and end-user documentation deletion repeat sizes range from 1 to 6+.

ReadStrelkaIDVCFs

Read Strelka ID (insertion and deletion) VCF files.

Description

Read Strelka ID (insertion and deletion) VCF files.

Usage

ReadStrelkaIDVCFs(files)

Arguments

files

Character vector of file paths to the VCF files.

Value

A list of vcfs from files.

Note

In ID (insertion and deletion) catalogs, deletion repeat sizes range from 0 to 5+, but for plotting and end-user documentation deletion repeat sizes range from 1 to 6+.

See Also

StrelkaIDVCFFilesToCatalog

revc

Reverse complement every string in string.vec.

Description

Reverse complement every string in string.vec.

Usage

```
revc(string.vec)
```

Arguments

string.vec

a vector of type character.

Value

A vector of type characters with the reverse complement of every string in string.vec.

StrelkaIDVCFFilesToCatalog

Create ID (indel) catalog from Strelka ID VCF files

Description

Create ID (indel) catalog from the Strelka ID VCFs specified by files

Usage

```
StrelkaIDVCFFilesToCatalog(files, ref.genome, region = "unknown")
```

Arguments

files Character vector of file paths to the Strelka ID VCF files.

ref.genome A ref.genome argument as described in ICAMS.

region A character string designating a genomic region; see as.catalog and ICAMS.

Details

This function calls VCFsToIDCatalogs

Value

An ID (indel) catalog with attributes added. See as.catalog for more details.

Note

In ID (insertion and deletion) catalogs, deletion repeat sizes range from 0 to 5+, but for plotting and end-user documentation deletion repeat sizes range from 1 to 6+.

StrelkaIDVCFFilesToCatalogAndPlotToPdf

Create ID (indel) catalog from Strelka ID VCF files and plot them to PDF

Description

Create ID (indel) catalog from the Strelka ID VCFs specified by files and plot them to PDF

Usage

```
StrelkaIDVCFFilesToCatalogAndPlotToPdf(files, ref.genome,
  region = "unknown", output.file)
```

Arguments

files Character vector of file paths to the Strelka ID VCF files.

ref.genome A ref.genome argument as described in ICAMS.

region A character string designating a genomic region; see as.catalog and ICAMS.

output.file The name of the PDF file to be produced.

Details

This function calls VCFsToIDCatalogs and PlotCatalogToPdf

Value

An ID (indel) catalog and its graph plotted to PDF with specified file name. The ID (indel) catalog has attributes added. See as.catalog for more details.

Note

In ID (insertion and deletion) catalogs, deletion repeat sizes range from 0 to 5+, but for plotting and end-user documentation deletion repeat sizes range from 1 to 6+.

StrelkaSBSVCFFilesToCatalog

Create SBS and DBS catalogs from Strelka SBS VCF files.

Description

Create 3 SBS catalogs (96, 192, 1536) and 3 DBS catalogs (78, 136, 144) from the Strelka SBS VCFs specified by files

Usage

```
StrelkaSBSVCFFilesToCatalog(files, ref.genome, trans.ranges = NULL,
  region = "unknown")
```

Arguments

files Character vector of file paths to the Strelka SBS VCF files.

ref.genome A ref.genome argument as described in ICAMS.

trans.ranges a data.table which contains transcript range and strand information. Please

refer to TranscriptRanges for more details.

region A character string designating a genomic region; see as.catalog and ICAMS.

Details

This function calls VCFsToSBSCatalogs and VCFsToDBSCatalogs.

Value

A list of 3 SBS catalogs (one each for 96, 192, and 1536) and 3 DBS catalogs (one each for 78, 136, and 144). If trans.ranges = NULL, SBS 192 and DBS 144 catalog will not be generated. Each catalog has attributes added. See as .catalog for more details.

Note

SBS 192 and DBS 144 catalog only contains mutations in transcribed regions.

StrelkaSBSVCFFilesToCatalogAndPlotToPdf

Create SBS and DBS catalogs from Strelka SBS VCF files and plot them to PDF

Description

Create 3 SBS catalogs (96, 192, 1536) and 3 DBS catalogs (78, 136, 144) from the Strelka SBS VCFs specified by files and plot them to PDF

Usage

```
StrelkaSBSVCFFilesToCatalogAndPlotToPdf(files, ref.genome,
  trans.ranges = NULL, region = "unknown", output.file)
```

Arguments

files Character vector of file paths to the Strelka SBS VCF files.

ref.genome A ref.genome argument as described in ICAMS.

trans.ranges a data.table which contains transcript range and strand information. Please

refer to TranscriptRanges for more details.

region A character string designating a genomic region; see as.catalog and ICAMS.

output.file The name of the PDF file to be produced.

Details

 $This \ function \ calls \ StrelkaSBSVCFFilesToCatalog \ and \ PlotCatalogToPdf$

TranscriptRanges 19

Value

A list of 3 SBS catalogs (one each for 96, 192, and 1536), 3 DBS catalogs (one each for 78, 136, and 144) and their graphs plotted to PDF with specified file name. If trans.ranges = NULL, SBS 192 and DBS 144 catalog will not be generated and plotted. Each catalog has attributes added. See as.catalog for more details.

Note

SBS 192 and DBS 144 catalogs include only mutations in transcribed regions.

TranscriptRanges

Transcript ranges data

Description

Transcript ranges and strand information for a particular reference genome.

Usage

```
trans.ranges.GRCh37
trans.ranges.GRCh38
```

Format

A data.table which contains transcript range and strand information for a particular reference genome. colnames are chrom, start, end, strand, gene.name. It uses one-based coordinates.

Details

This information is needed to generate catalogs that depend on transcriptional strand information, for example catalogs of class SBS192Catalog.

```
trans.ranges.GRCh37: Human GRCh37.
trans.ranges.GRCh38: Human GRCh38.
```

For these two tables, only genes that are associated with a CCDS ID are kept for transcriptional strand bias analysis.

This information is needed for StrelkaSBSVCFFilesToCatalog, StrelkaSBSVCFFilesToCatalogAndPlotToPdf, MutectVCFFilesToCatalog, MutectVCFFilesToCatalogAndPlotToPdf, VCFsToSBSCatalogs and VCFsToDBSCatalogs.

Source

```
ftp://ftp.ebi.ac.uk/pub/databases/gencode/Gencode_human/release_30/GRCh37_mapping/
gencode.v30lift37.annotation.gff3.gz
ftp://ftp.ebi.ac.uk/pub/databases/gencode/Gencode_human/release_30/gencode.v30.annotation.
gff3.gz
```

20 TransformCatalog

g	between counts and density spectrum catalogs and counts ty signature catalogs.
---	--

Description

Transform between counts and density spectrum catalogs and counts and density signature catalogs.

Usage

```
TransformCatalog(catalog, target.ref.genome, target.region,
  target.catalog.type, target.catalog.abundance = NULL)
```

Arguments

catalog An SBS or DBS catalog as described in ICAMS; must **not** be an ID (indel) catalog.

target.ref.genome

A ref. genome argument as described in ICAMS.

target.region A region argument; see as.catalog and ICAMS.

target.catalog.type

A character string acting as a catalog type identifier, one of "counts", "density", "counts.signature", "density.signature"; see as.catalog.

target.catalog.abundance

Optional, only needed when target.ref.genome is not one of the two human reference genomes included in ICAMS. The abundance should contain the counts of different source sequences for mutations.

See ICAMS:::abundance.3bp.exome.unstranded.GRCh37 for an example.

Details

Only the following transformations are legal:

- 1. counts -> counts (used to transform between target.ref.genome and/or target.region)
- 2. counts -> density
- 3. counts -> (counts.signature,density.signature)
- 4. density -> counts (the semantics are to infer the genome-wide or exome-wide counts based on the densities)
- 5. density -> (counts.signature,density.signature)
- 6. counts.signature -> (counts.signature, density.signature)
- 7. density.signature -> counts.signature
- 8. density.signature -> density.signature (a null operation)
- 9. density -> density (a null operation)

Value

A catalog as defined in ICAMS.

VCFsToDBSCatalogs 21

VCFsToDBSCatalogs	Create DBS catalogs from VCFs
-------------------	-------------------------------

Description

Create a list of 3 catalogs (one each for DBS78, DBS144 and DBS136) out of the contents in list.of.DBS.vcfs. The VCFs must not contain any type of mutation other then DBSs.

Usage

```
VCFsToDBSCatalogs(list.of.DBS.vcfs, ref.genome, trans.ranges = NULL,
  region = "unknown")
```

Arguments

list.of.DBS.vcfs

List of in-memory data frames of pure DBS mutations - no SBS or 3+BS muta-

tions. The list names will be the sample ids in the output catalog.

ref.genome A ref.genome argument as described in ICAMS.

trans.ranges a data.table which contains transcript range and strand information. Please

refer to TranscriptRanges for more details.

region A character string designating a genomic region; see as . catalog and ICAMS.

Value

A list of 3 DBS catalogs, one each for 78, 144, 136: catDBS18 catDBS144 catDBS136. If trans.ranges = NULL, DBS 144 catalog will not be generated. Each catalog has attributes added. See as.catalog for more details.

Note

DBS 144 catalog only contains mutations in transcribed regions.

VCFsToIDCatalogs Create ID (insertion and deletion) catalog from ID VCFs

Description

Create ID (insertion and deletion) catalog from ID VCFs

Usage

```
VCFsToIDCatalogs(list.of.vcfs, ref.genome, region = "unknown")
```

Arguments

list.of.vcfs List of in-memory VCFs. The list names will be the sample ids in the output

catalog.

ref.genome A ref.genome argument as described in ICAMS.

region A character string acting as a region identifier, one of "genome", "exome".

22 WriteCatalog

Value

An S3 object containing an ID (indel) catalog with class "catalog". See as.catalog for more details.

VCFsToSBSCatalogs

Create SBS catalogs from SBS VCFs

Description

Create a list of 3 catalogs (one each for 96, 192, 1536) out of the contents in list.of.SBS.vcfs. The SBS VCFs must not contain DBSs, indels, or other types of mutations.

Usage

```
VCFsToSBSCatalogs(list.of.SBS.vcfs, ref.genome, trans.ranges = NULL,
  region = "unknown")
```

Arguments

list.of.SBS.vcfs

List of in-memory data frames of pure SBS mutations – no DBS or 3+BS muta-

tions. The list names will be the sample ids in the output catalog.

ref.genome A ref.genome argument as described in ICAMS.

trans.ranges a data.table which contains transcript range and strand information. Please

refer to TranscriptRanges for more details.

region A character string designating a genomic region; see as.catalog and ICAMS.

Value

A list of 3 SBS catalogs, one each for 96, 192, 1536: catSBS96 catSBS192 catSBS1536. If trans.ranges = NULL, SBS 192 catalog will not be generated. Each catalog has attributes added. See as.catalog for more details.

Note

SBS 192 catalogs only contain mutations in transcribed regions.

WriteCatalog

Write a catalog

Description

Write a catalog to a file.

Usage

```
WriteCatalog(catalog, file, strict = TRUE)
```

WriteCatalog 23

Arguments

catalog A catalog as defined in ICAMS; see also as.catalog.

file The path to the file to be created.

strict If TRUE, do additional checks on the input, and stop if the checks fail.

Details

See also ReadCatalog.

Note

In ID (insertion and deletion) catalogs, deletion repeat sizes range from 0 to 5+, but for plotting and end-user documentation deletion repeat sizes range from 1 to 6+.

Index

*Topic datasets	ReadStrelkaIDVCFs, 15
all.abundance, 2	revc, 16
CatalogRowOrder, 3	
TranscriptRanges, 19	$\label{eq:strelkaldvCFFilesToCatalog} StrelkaIDVCFFilesToCatalogAndPlotToPdf,$
all.abundance, 2, 7, 8	17
as.catalog, 3, 7, 8, 10–12, 14–23	StrelkaSBSVCFFilesToCatalog, 8, 14, 17,
available.genomes, 8	18, 19
	StrelkaSBSVCFFilesToCatalogAndPlotToPdf,
BSgenome, 8	8, 18, <i>19</i>
BSgenome.Hsapiens.1000genomes.hs37d5,	trans ranges (DCh27 (TranscriptDanges)
8	trans.ranges.GRCh37(TranscriptRanges), 19
BSgenome.Hsapiens.UCSC.hg38,8	
antalas naw andan (Catalas Daw Ondan) 2	trans.ranges.GRCh38(TranscriptRanges), 19
catalog.row.order (CatalogRowOrder), 3	
CatalogRowOrder, 3, 3, 9	TranscriptRanges, 9–11, 18, 19, 21, 22
Collapse144CatalogTo78	TransformCatalog, 7, 9, 11, 12, 20
(CollapseCatalog), 4 Collapse1536CatalogTo96	VCFsToDBSCatalogs, 10, 18, 19, 21
(CollapseCatalog), 4	VCFsToIDCatalogs, 10, 16, 17, 21
Collapse192CatalogTo96	VCFsToSBSCatalogs, 10, 18, 19, 22
(CollapseCatalog), 4	
CollapseCatalog, 4, 9	WriteCatalog, 9, 15, 22
corrapsecatarog, 4, 9	
data.table, 10, 11, 18, 19, 21, 22	
FindDelMH, 4	
GetMutectVAF (GetVAF), 6	
GetStrelkaVAF (GetVAF), 6	
GetVAF, 6	
ICAMS, 3, 4, 7, 10–12, 14, 16–18, 20–23 ICAMS-package (ICAMS), 7	
Totalo pueltage (Totalo), /	
MutectVCFFilesToCatalog, 8, 10, 11, 13, 19	
MutectVCFFilesToCatalogAndPlotToPdf,	
10, 19	
PlotCatalog, 8, 11	
PlotCatalogToPdf, <i>8</i> , <i>11</i> , 12, <i>17</i> , <i>18</i>	
ReadAndSplitMutectVCFs, 13	
ReadAndSplitStrelkaSBSVCFs, 14	
ReadCatalog, 8, 9, 14, 23	