

# Package ‘seqCAT’

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**Title** High Throughput Sequencing Cell Authentication Toolkit

**Version** 1.2.1

**Description** The seqCAT package uses variant calling data (in the form of VCF files) from high throughput sequencing technologies to authenticate and validate the source, function and characteristics of biological samples used in scientific endeavours.

**Depends** R (>= 3.5), GenomicRanges (>= 1.26.4), VariantAnnotation(>= 1.20.3)

**Imports** dplyr (>= 0.5.0), GenomeInfoDb (>= 1.13.4), ggplot2 (>= 2.2.1), grid (>= 3.5.0), IRanges (>= 2.8.2), lazyeval (>= 0.2.0), scales (>= 0.4.1), S4Vectors (>= 0.12.2), stats, SummarizedExperiment (>= 1.4.0), tidyr (>= 0.6.1), utils

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**calculate\_similarity** *SNV profile similarity calculations*

---

**Description**

Calculate the similarity statistics for SNV profile comparisons.

**Usage**

```
calculate_similarity(data, similarity = NULL, a = 1, b = 5)
```

**Arguments**

- data**            The input SNV data dataframe.
- similarity**      Optional dataframe to add results to.
- a**              Similarity score parameter a (integer).
- b**              Similarity score parameter b (integer).

**Details**

This function calculates various summary statistics and sample similarities for a given profile comparison dataframe. It returns a small dataframe with the overall similarity score (whose parameters ‘a’ and ‘b’ can be adjusted in the function call), total SNV data, the concordance of the data and the sample names in question. This dataframe can also be given to the function, in which case it will simply add another row for the current samples, facilitating downstream aggregate analyses.

**Value**

A dataframe with summary statistics.

## Examples

```
# Load test data
data(test_comparison)

# Calculate similarities
similarity <- calculate_similarity(test_comparison)

# Add another row of summary statistics
calculate_similarity(test_comparison, similarity = similarity)
```

compare\_many

*Comparisons of many SNV profiles*

## Description

Overlap and compare genotypes in many SNV profiles.

## Usage

```
compare_many(many, one = NULL, a = 1, b = 5)
```

## Arguments

many	SNV profiles to be compared (list of GRanges objects).
one	SNV profile to be compared to all others (GRanges object).
a	Similarity score parameter a (integer).
b	Similarity score parameter b (integer).

## Details

This is a function that compares all the combinations of the SNV profiles input to it, either in a one-to-many or many-to-many manner. It returns both a dataframe containing summary statistics for all unique combinations and a list of dataframes with all the performed comparisons, for easy re-use and downstream analyses of said comparisons.

## Value

A list of summary statistics and comparisons.

## Examples

```
# Load test data
data(test_profile_1)
data(test_profile_2)

# Perform many-to-many comparisons
profiles <- list(test_profile_1, test_profile_2)
comparisons <- compare_many(profiles)

# View aggregate similarities
## Not run: comparisons[[1]])

# View data of first comparison
## Not run: head(comparisons[[2]][[1]])
```

`compare_profiles`      *Binary SNV profile comparisons*

## Description

Overlap and compare genotypes in two SNV profiles.

## Usage

```
compare_profiles(profile_1, profile_2, mode = "intersection")
```

## Arguments

- |                        |   |
|------------------------|---|
| <code>profile_1</code> | The first SNV profile (GRanges object).                     |
| <code>profile_2</code> | The second SNV profile (GRanges object).                    |
| <code>mode</code>      | Merge profiles using "union" or "intersection" (character). |

## Details

This is a function for finding overlapping variants in two different SNV profiles (stored as GenomicRanges objects), followed by comparing the genotypes of the overlapping variants. The "compare\_overlaps" function calls the "add\_metadata" function twice in succession in order to merge the metadata for the two profiles (supplied as GRanges objects), returns the results as a dataframe, compares the genotypes of the overlapping variants using the "compare\_genotypes" function and, finally, returns the final dataframe with all variant overlaps and their similarity.

## Value

A dataframe.

## Examples

```
# Load test data
data(test_profile_1)
data(test_profile_2)

# Compare the two profiles
comparison <- compare_profiles(test_profile_1, test_profile_2)
```

`create_profile`      *SNV profile creation*

## Description

Create an SNV profile from data in a VCF file.

## Usage

```
create_profile(vcf_file, sample, output_file, filter_depth = 10,
               python = FALSE)
```

### Arguments

vcf_file	The VCF file from which the profile will be created (path).
sample	The sample in the VCF for which a profile will be created (character).
output_file	The output file with the SNV profile (path).
filter_depth	Remove variants below this sequencing depth (integer).
python	Extract variants using Python instead of R (boolean).

### Details

This function creates a SNV profile from a given VCF file by extracting the variants that pass the filtering criterias. It can either be performed using R, or by the `create_profile.py` function included (which requires that Python is installed, along with the PyVCF package). Profile creation is performed to facilitate and accelerate the cell authentication procedures, which is especially relevant when more than one pairwise comparison will be performed on the same sample.

### Value

Does not return any data object, but outputs results to `output_file` (to save computational time from having to repeatedly create profiles).

### Examples

```
# Path to the test VCF file
vcf_file = system.file("extdata", "test.vcf.gz", package = "seqCAT")

# Create SNV profiles
## Not run:
create_profile(vcf_file, "sample1", "profile1.txt")
create_profile(vcf_file, "sample1", "profile1.txt", filter_depth = 15)
create_profile(vcf_file, "sample1", "profile1.txt", python = TRUE)

## End(Not run)
```

### Description

Filter variants on sequencing depth.

### Usage

```
filter_variants(data, filter_depth = 10)
```

### Arguments

data	The dataframe containing the variant data to be filtered.
filter_depth	Threshold for variant depth (integer; default 10).

## Details

This is a function for filtering variants on sequencing depth. Variants with a depth lower than 10 are removed by default, but can be changed in the function call.

## Value

A data frame containing the filtered variants.

## Examples

```
# Load test comparisons
data(test_comparison)

# Filter variants
filt_1 <- filter_variants(test_comparison)
filt_2 <- filter_variants(test_comparison, filter_depth = 20)
```

**list\_variants**            *List known variants*

## Description

List known variants present in SNV profiles

## Usage

```
list_variants(profiles, known_variants)
```

## Arguments

profiles	The SNV profiles to analyse (list)
known_variants	The known variants to look for (dataframe)

## Details

This is a function for listing known variants present in SNV profiles. Input is a list of profiles and a dataframe of known variants, containing at least the genomic locations ("chr" and "pos"). Any additional columns will be retained.

## Value

A dataframe containing the known variant genotypes in each profile.

## Examples

```
# Load test data
data(test_profile_1)
data(test_profile_2)

# Create some variants to analyse
known_variants <- data.frame(chr = 1, pos = 16229, gene = "DDX11L1")

# List the known variants in each profile
```

```
profiles <- list(test_profile_1, test_profile_2)
known_variants <- list_variants(profiles, known_variants)
```

---

**plot\_heatmap***Plot similarity heatmap*

---

**Description**

Plot a heatmap of similarities from many-to-many SNV profile comparisons.

**Usage**

```
plot_heatmap(similarities, cluster = TRUE, annotate = TRUE,
             annotate_size = 9, legend = TRUE, legend_size = c(36, 8),
             limits = c(0, 50, 90, 100), text_size = 14, colour = "#1954A6")
```

**Arguments**

<code>similarities</code>	The long-format dataframe containing the data.
<code>cluster</code>	Cluster the samples based on similarity (boolean).
<code>annotate</code>	Annotate each cell with the score (boolean).
<code>annotate_size</code>	Text size of the annotations (numeric).
<code>legend</code>	Show a legend for the colour gradient (boolean).
<code>legend_size</code>	Height and width of the legend (vector of two integers).
<code>limits</code>	The limits for the colour gradient (vector of four integers).
<code>text_size</code>	Text size for axes, labels and legend (numeric).
<code>colour</code>	The main colour to use for the gradient (character).

**Details**

This function creates publication-ready plots of heatmaps for many-to-many sample comparisons, taking a long-format dataframe containing the summary statistics of each comparison as input.

**Value**

A ggplot2 graphical object.

**Examples**

```
# Load test similarities
data(test_similarities)

# Plot a similarity heatmap
heatmap <- plot_heatmap(test_similarities)
```

**plot\_impacts**      *Plot SNV impact distribution*

### Description

Plot SNV impact distributions for a binary SNV profile comparison.

### Usage

```
plot_impacts(comparison, legend = TRUE, annotate = TRUE,
             annotate_size = 9, text_size = 14, palette = c("#0D2D59", "#1954A6"))
```

### Arguments

comparison	The SNV profile comparison to be plotted.
legend	Show the legend (boolean).
annotate	Annotate each category (boolean).
annotate_size	Text size for annotations (numeric).
text_size	Text size for axes, ticks and legend (numeric).
palette	Colour palette for filling of bars (character vector).

### Details

This function creates publication-ready plots of the impact distribution from a binary dataset comparison across the matched/mismatched SNVs.

### Value

A ggplot2 graphical object.

### Examples

```
# Load test comparison data
data(test_comparison)

# Plot the impact distribution
impacts <- plot_impacts(test_comparison)
```

**plot\_variant\_list**      *Plot known variants list*

### Description

Plot a genotype grid from a list of known variants

### Usage

```
plot_variant_list(variant_list, legend = TRUE, legend_size = 22,
                  text_size = 14, palette = c("#4e8ce4", "#a6c6f2", "#999999", "#cccccc"))
```

**Arguments**

variant_list	The data containing the variants (dataframe)
legend	Show a legend for the genotype colours (boolean)
legend_size	Size of the legend (numeric).
text_size	Text size for axes and legend (numeric).
palette	Nucleotide colour palette (4-element character vector)

**Details**

This function creates publication-ready plots from lists of known variants, taking a dataframe containing all the genotypes (on "A1/A2" format) for each sample (columns) and variant (row names).

**Value**

A ggplot2 graphical object.

**Examples**

```
# Load test variant list
data(test_variant_list)

# Plot each variant's genotype per sample
genotype_grid <- plot_variant_list(test_variant_list)
```

---

**read\_cosmic***Read COSMIC SNV data*

---

**Description**

Read COSMIC cell line-specific mutational data.

This function lists the available cell lines in the provided CosmicCLP\_MutantExport.tsv.gz file, and take about half the time it takes to read the full file with the read\_cosmic function, making it useful for just seeing if your particular cell line is listed in COSMIC or not.

**Usage**

```
read_cosmic(file_path, cell_line)

list_cosmic(file_path)
```

**Arguments**

file_path	The CosmicCLP_MutantExport.tsv.gz file (path).
cell_line	The cell line to be investigated (character).

**Details**

This function reads the "CosmicCLP\_MutantExport.tsv.gz" file obtained from COSMIC and returns a GRanges object with all the listed mutations for the specified cell line, which can then be used in downstream profile comparisons. Only non-duplicated (gene-level) SNVs are included in COSMIC profiles.

**Value**

A GRanges object with COSMIC SNVs.

A vector of cell line names

**Examples**

```
# Path to COSMIC test data
file <- system.file("extdata",
  "subset_CosmicCLP_MutantExport.tsv.gz",
  package = "seqCAT")

# Read COSMIC test data for HCT116 cell line
cosmic_hct116 <- read_cosmic(file, "HCT116")
file <- system.file("extdata",
  "subset_CosmicCLP_MutantExport.tsv.gz",
  package = "seqCAT")
cell_lines <- list_cosmic(file)
```

**read\_profile***Read SNV profile***Description**

Read SNV profiles for use in downstream comparisons.

**Usage**

```
read_profile(file, sample_name, remove_mt = TRUE)
```

**Arguments**

<code>file</code>	The SNV profile to be read (path).
<code>sample_name</code>	The sample of the SNV profile (character).
<code>remove_mt</code>	Remove or keep mitochondrial variants (boolean).

**Details**

This is a function for reading SNV profiles created from VCF files. The data is returned as a GenomicRanges object, suitable for merging of metadata.

**Value**

A GRanges object.

**Examples**

```
# Path to test data
profile = system.file("extdata",
  "test_profile_1.txt.gz",
  package = "seqCAT")

# Read test profile
profile_1 <- read_profile(profile, "sample1")
```

**Description**

The \*seqCAT\* package provides a number of functions for performing evaluation, characterisation and authentication of biological samples through analysis of high throughput sequencing data.

**Description**

Overlapping and compared variants from "sample1" and "sample2" originating from the example.vcf file included in the inst/extdata directory, for use in unit tests.

**Usage**

```
data(test_comparison)
```

**Format**

A dataframe with 51 rows and 39 columns:

**chr** chromosome  
**pos** SNV position  
**DP.sample\_1** total variant depth, sample 1  
**AD1.sample\_1** allelic depth, allele 1, sample 1  
**AD2.sample\_1** allelic depth, allele 2, sample 1  
**A1.sample\_1** allele 1, sample 1  
**A2.sample\_1** allele 2, sample 1  
**warnings.sample\_1** warnings from variant calling, sample 1  
**DP.sample\_2** total variant depth, sample 2  
**AD1.sample\_2** allelic depth, allele 1, sample 2  
**AD2.sample\_2** allelic depth, allele 2, sample 2  
**A1.sample\_2** allele 1, sample 2  
**A2.sample\_2** allele 2, sample 2  
**warnings.sample\_2** warnings from variant calling, sample 2  
**sample\_1** name, sample 1  
**sample\_2** name, sample 2  
**match** status of genotype comparison  
**rsID** mutation ID  
**gene** associated gene  
**ENSGID** ensembl gene ID

**ENSTID** ensembl transcript ID  
**REF** reference allele  
**ALT** alternative allele  
**impact** putative variant impact  
**effect** variant effect  
**feature** transcript feature  
**biotype** transcript biotype

**test\_profile\_1**      *SNV profile 1*

### Description

SNV profile in GRanges format from "sample1", originating from the test\_profile\_1.txt in the inst/extdata directory, for use in unit tests.

### Usage

```
data(test_profile_1)
```

### Format

A GRanges object with 383 elements and 17 metadata columns:

**rsID** mutation ID, if available  
**gene** associated gene  
**ENSGID** ensembl gene ID  
**ENSTID** ensembl transcript ID  
**REF** reference allele  
**ALT** alternative allele  
**impact** putative variant impact  
**effect** variant effect  
**feature** transcript feature  
**biotype** transcript biotype  
**DP** total variant depth  
**AD1** allelic depth, allele 1  
**AD2** allelic depth, allele 2  
**A1** allele 1  
**A2** allele 2  
**warnings** warnings from variant calling  
**sample** sample name

---

test_profile_2	<i>SNV profile 2</i>
----------------	----------------------

---

## Description

SNV profile in GRanges format from "sample2", originating from the test\_profile\_2.txt in the inst/extdata directory, for use in unit tests.

## Usage

```
data(test_profile_2)
```

## Format

A GRanges object with 382 elements and 17 metadata columns:

**rsID** mutation ID, if available  
**gene** associated gene  
**ENSGID** ensembl gene ID  
**ENSTID** ensembl transcript ID  
**REF** reference allele  
**ALT** alternative allele  
**impact** putative variant impact  
**effect** variant effect  
**feature** transcript feature  
**biotype** transcript biotype  
**DP** total variant depth  
**AD1** allelic depth, allele 1  
**AD2** allelic depth, allele 2  
**A1** allele 1  
**A2** allele 2  
**warnings** warnings from variant calling  
**sample** sample name

**test\_profile\_3**      *SNV profile 3*

### Description

SNV profile in GRanges format from "sample3", originating from the test\_profile\_3.txt in the inst/extdata directory, for use in unit tests.

### Usage

```
data(test_profile_3)
```

### Format

A GRanges object with 99 elements and 9 metadata columns:

**rsID** mutation ID, if available

**REF** reference allele

**ALT** alternative allele

**DP** total variant depth

**AD1** allelic depth, allele 1

**AD2** allelic depth, allele 2

**A1** allele 1

**A2** allele 2

**sample** sample name

**test\_similarities**      *Collated similarities object*

### Description

Collated similarities of multiple sample comparisons from "sample1" and "sample" from the example.vcf file, for use in unit tests.

### Usage

```
data(test_similarities)
```

### Format

A dataframe with 3 rows and 6 columns:

**sample\_1** name of sample 1

**sample\_2** name of sample 2

**overlaps** the number of overlaps for the comparison

**matches** the number of matches for the comparison

**concordance** the concordance of the profiles

**similarity\_score** the similarity score of the profiles

---

**test\_variant\_list**      *Modified variant list object*

---

### Description

A variant list object from the ‘list\_variants‘ function, where the row names have been defined as "chr: pos (gene)" and the corresponding columns removed, for use in plotting.

### Usage

```
data(test_variant_list)
```

### Format

A dataframe with 2 rows and 2 columns:

**sample1** the genotypes of sample1

**sample2** the genotypes of sample2

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