OncodriveFML Documentation

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BBGLab

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ONCODRIVEFML

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ONCODRIVEFML

Distinguishing the driver mutations from somatic mutations in a tumor genome is one of the major challenges of cancer research. This challenge is more acute and far from solved for non-coding mutations. OncodriveFML is a method designed to analyze the pattern of somatic mutations across tumors in both coding and non-coding genomic regions to identify signals of positive selection, and therefore, their involvement in tumorigenesis. We described the method and illustrated its usefulness to identify protein coding genes, promoters, untranslated regions, intronic splice regions, and lncRNAs-containing driver mutations in several malignancies in Mularoni et al., Genome Biology 2016.

To use OncodriveFML check its website or download the source code from our git repository.

OncodriveFML is a project developed by the Barcelona Biomedical Genomics Lab.

We are a research group integrated in the Institute for Research Biomedicine in Barcelona, which is part of the Barcelona Institute of Science and Technology. Our lab is located at the Barcelona Science Park.

Our main research interest is the computational study of cancer at the genomic level.

Check the README file to find infomation about licensing and installation.

Run the example for a quick check of the installation.

TWO

HOW IT WORKS

This section will try to give an overview of how OncodriveFML carries on the analysis.

2.1 The command line interface

By typing oncodrivefml -h you will have a brief description of how to use OncodriveFML:

Options:

- -i, --input MUTATIONS_FILE Variants file [required] (see format)
- -e, --elements ELEMENTS_FILE Genomic elements to analyse [required] (see format)
- -o, --output OUTPUT_FOLDER Output folder. Default to regions file name without extensions.
- -c, --configuration CONFIG_FILE Configuration file. Default to 'oncodrivefml_v2.conf' in the current folder if exists or to ~/.config/bbglab/oncodrivefml_v2.conf if not.
- --samples-blacklist SAMPLES_BLACKLIST Remove these samples when loading the input file.
- --signature SIGNATURE File with the signatures to use

See *details about the command line interface* to find more information about this option.

- -signature-correction [wglwx] Correct the computed signutares by genomic or exomic signtures. Only valid for human genomes (hg19 and hg38)
 - wg: correction using whole genome counts
 - wx: correction using whole exome counts

See details about the command line interface to find more information about this option.

no-indels	Discard indels in your analysis
cores INTEGER	Cores to use. Default: all
seed INTEGER	Set up an initial random seed to have reproducible results
debug	Show more progress details
version	Show the version and exit.
-h,help	Show this message and exit.

2.2 The files

2.2.1 Input files

OncodriveFML makes use of three files:

Variants Also named as input. This file contains the observed mutations for the analysis.

Regions File containing the regions for the analysis. Only mutations that fall in these regions are analysed and only the genomic positions defined in this file are used for the simulation.

You can define your own regions file based on your criteria. You can check an example of a regions file downloading our example.

Warning: It is not recommended to mix coding and non-coding regions in your regions file. In fact this will likely produce artifacts in the results as coding and non-coding regions of the genome have a very different functional impact scores. A good set of genomic regions should include elements that share biological functions (e.g. CDS, UTRs, promoters, enhancers, etc.).

Check the formats for the input files.

Configuration The configuration file is also a key part of the run, and understanding how to adapt it to your needs is important. Check *this section* to find more details about it.

2.2.2 Output files

Find information about the output output files section.

2.3 Workflow

- 1. The first thing that is done by OncodriveFML is to load the configuration file.
- 2. The output is checked. The default behaviour is that OncodriveFML creates an output folder in the current directory with the same name as the elements file (without extension).

If an output is provided and it exists and is a folder, OncodriveFML checks whether a file with the expected output name exits and, if so, it does not run. Otherwise, it assumes it is a path name an uses that as output.

Note: If the output does not exits, OncodriveFML only computes the tsv file with the results and skips the plots.

- 3. The regions file is loaded, and a tree with the intervals is created. This tree is used to find which mutations fall in the regions being analysed.
- 4. Loads the mutations file and keeps only the ones that fall into the regions being analysed.
- 5. Computes the signature (see the signature section), if not provided as an external file.
- 6. Analyses each region separately (only the ones that have mutations). In each region the analysis is as follow:
 - 1. Computes the score of each of the observed mutations.
 - 2. Simulates the same number of mutations in the segments of the region under analysis. Save the scores of each of the simulated mutations. The simulation is done several times.

- 3. Applies a predefined function to the observed scores and to each of the simulated groups of scores. Counts how many times the simulated value is higher than, or equal to, the observed.
- 4. From these counts, computes a P-value by dividing the counts by the number of simulations performed.

You can find more details in the *analysis section*.

- 7. Joins the results and performs a multiple test correction. The multiple test correction is only done for regions with mutations from at least two samples.
- 8. Creates the *output files*.
- 9. Checks that the output file does not contain missing or repeated genomic regions.

THREE

FILES

3.1 File formats

Note: All the files can be compressed using GZIP (extension ".gz"), BZIP2 (extension ".bz2") or LZMA (extension ".xz")

3.1.1 Input file format

The variants file is a text file with, at least, 5 columns separated by a tab character (the header is required, but the order of the columns can change):

- Column CHROMOSOME: Chromosome. A number between 1 and 22 or the letter X or Y (upper case)
- Column POSITION: Mutation position. A positive integer.
- Column REF: Reference allele¹.
- Column ALT: Alternate allele¹.
- Column SAMPLE: Sample identifier. Any alphanumeric string.
- Column CANCER_TYPE: Cancer type. Any alphanumeric string. Optional.
- Column SIGNATURE: User defined signature categories. Any alphanumeric string. Optional.

Mutations are expected to be in the positive strand.

Note: OncodriveFML, although reading the SAMPLE column, it does not perform a per-sample analysis.

A **by-sample** option can be enabled in the configuration file, in which only one mutation per sample is included in the analysis. More details in the *configuration section*.

¹ The alleles consist on a single letter or a set of letters using A, C, G or T (upper case). Single Nucleotide Variants are indentified because both, REF and ALT contain only one letter. In Multi-Nucleotide Variants REF and ALT columns contain a set of letters of the same length. Insertions use – in the REF and a set of letters as ALT while deletions contain the set of deleted characters in the REF and – in the ALT columns.

3.1.2 Regions file format

The regions file is a text file with, at least, 4 columns separated by a tab character (header is required):

- Column CHROMOSOME]: Chromosome. A number between 1 and 22 or the letter X or Y (upper case)
- Column START: Start position. A positive integer.
- Column END: End position. A positive integer.
- Column ELEMENT: Element identifier. Can appear multiple times if the element is divided in *segments*.

Important: Analysis is perform element-wise. One single element can have multiple *segments* (even if you do not provide an identifier for them).

It is also important that different segments of the same element do not overlap.

Optional columns are:

- Column STRAND: Strand: + for positive, for negative, . for unknown.
- Column SEGMENT: Segment identifier. Optional column.
- Column SYMBOL: Symbol, a different identifier for the element that will also be printed in the output file. Optional column.

3.1.3 Signature file format

The signature file is a JSON file, where pairs of key-values represent the changes and the probabilities of those changes.

Changes are represented as AAA>C (reference triplet, > and alternate).

See the bgsignature package for more information on how to create such signatures.

3.1.4 Output file format

OncodriveFML generates a tabulated file with the results with the extension ".tsv.gz". It is compressed with gzip.

Check the *output section* to find a detailed description regarding the output.

CONFIGURATION

The method behaviour can be modified through a configuration file.

Warning: Using the command line interface overwrites some setting in the configuration file. Check how the command line interface changes the configuration in the *command line interface* section.

Check the $oncodrivefml_v2.conf.template$ that is included in the package to find an example of the configuration file.

This section will explain each of the parameters in the configuration file:

4.1 Genome

```
[genome]
# Build of the reference genome
build = 'hg19'
```

The genome section makes reference to the reference genome used by OncodriveFML.

The reference genome has been obtained from http://hgdownload.cse.ucsc.edu/downloads.html.

Currently, only HG19 and HG38 are fully supported. Use build = 'hg19' or build = 'hg38' to use any of them.

There is a partial support for other genomes. The support is only partial because the values for the position and alterations of the stops in the these genomes have not been computed yet. If you want to run OncodriveFML with any of these genomes, make sure you do not use the stop method for the indels (*ref*). In addition, signature correction cannot be performed.

Warning: Make sure that the scores file you use is compatible with your reference genome. For human reference genomes, we have been using CADD scores

4.2 Signature

```
[signature]
# Choose the method to calculate the trinuclotide singature:
method = 'complement'
# Choose the classifier (categorical value for the signature):
classifier = 'SAMPLE'
# None: do not correct (comment the option)
# normalize_by_sites = ''
```

The signature represents the probability of a certain nucleotide to mutate taking into account its context¹.

The signature can be configured using the following parameters:

method Method used to compute the signature. Options are:

- method = 'none': all changes have equal probability. This approach is recommended for small datasets.
- method = 'complement': use a 96 matrix with the signatures complemented.
- method = 'full': use a 192 matrix with all the possible signatures.
- method = 'bysample': equivalent to method = 'complement' and classifier =
 'SAMPLE'
- method = 'file': use a precomputed signature. This option requires to add the path to the file as path = '/path/to/file'. Note that this option *can* be overwritten by the *-signature option* in the command line interface.

The precomputed signature can be obtained using

the bgsignature package.

classifier The signature by default is computed using the whole dataset. However, you can group the mutations in categories that correspond to any of the values in SAMPLE, CANCER_TYPE and SIGNATURE columns (if provided).

If a file with the signature is provided, and that signature has also been computed using groups, the same classifier must be specified.

normalize_by_sites Compute a normalization of the signature. This option appears commented because it *is* overridden by the *-signature-correction option* of the command line interface.

If you provide an external file with the signature, it will never be corrected, regardless of the value of this option.

The recommended approach is to compute your own signature (e.g. using the bgsignature package) and pass it to OncodriveFML.

¹ Previous and posterior nucleotides

4.3 Score

The score section is used to know which scores are going to be used.

```
[score]
# Path to score file
file = "/path/to/scores/file"
# Format of the file
format = 'tabix'
# Column that has the chromosome
chr = 0
# If the chromosome has a prefix like 'chr'. Example: chrX chr1 ...
chr_prefix = ''
# Column that has the position
pos = 1
# Column that has the reference allele
ref = 2
# Column that has the alternative allele
alt = 3
# Column that has the score value
score = 5
```

The scores should be a file that for a given position, in a given chromosome, gives a value to every possible alteration.

Some of the parameters in this section are optional, while others are mandatory.

file It is a string and represents the path to the scores file.

format Indicates the format of the file. Options are:

- format = 'tabix' indicates that the file is a tab separated file compressed with bgzip. This means that a .tbi index file should be present in the same location.
- format = 'pack' is a binary format we have implemented to reduce the file size. It is only available for specific scores.

Thus, if you want to use your own file, use the tabix format.

- chr Column in the file where the chromosome is indicated.
- chr_prefix When querying the tabix file for a specif chromosome OncodriveFML only uses the number of the chromosome or 'X' or 'Y'. If the tabix file requires a prefix before the chromosome, use this option. For instance, if the chromosomes in the tabix file are labeled as chr1, chr2, ..., chrY, set this option to: chr_prefix = 'chr'. If this is not the case, use an empty string: chr_prefix = ''.
- pos Column that indicates the position of the scored alteration in the chromosome.
- ref Column that contains the reference allele. It is optional.
- **alt** Column that contains the alternate allele. It is optional. If is not specified, it is assumed that the 3 possible changes have the same score.
- score Column that contains the score.
- **element** Column that contains the element identifier. It is optional. If it is provided and the value does not match with the one from the regions, these scores are discarded.

4.4 Statistic

The statistic section is related to the configuration of the analysis

```
[statistic]
# Mathematical method to use to compare observed and simulated values
method = 'amean'
# Do not use/use MNP mutations in the analysis
discard_mnp = False
# Compute the observed values using only 1 mutation per sample
# per_sample_analysis = 'max'
# Minimum sampling
sampling = 100000
# Maximum sampling
sampling_max = 1000000
# Sampling chunk (in millions)
sampling_chunk = 100
# Minimum number of observed (if not reached, keeps computing)
sampling_min_obs = 10
    [[indels]]
    # Include/exclude indels from your analysis
   include = True
    # Method used to simulate indels
   method = 'max'
    # Number of consecutive times the indel appears to consider it falls in a_
\rightarrow repetitive region
   max\_consecutive = 7
```

There a different parameters you can configure:

method Represents the type of operation that is applied to observed and simulated scores before comparing them. Options are:

- method = 'amean': arithmetic mean
- method = 'gmean': geometric mean

discard_mnp Indicates whether to include or not MNP mutations in the analysis.

- discard_mnp = False: include them
- discard_mnp = True: discard them
- **per_sample_analysis** In some cases, you might be interested in performing the analysis per sample. This means that all the mutations that come from the same sample are reduced to a single score. This score can be computed as:
 - per_sample_analysis = 'max': maximum score
 - per_sample_analysis = 'amean': arithmetic mean
 - per_sample_analysis = 'gmean': geometric mean

Comment this option if you are not interested in this type of analysis.

OncodriveFML includes a few more parameters that are related to how many simulations are performed.

sampling Represents the minimum number of simulations to be performed.

sampling_max Represents the maximum number of simulations to be performed.

sampling_chunk Represents the maximum size (in millions) that a single process can handle. This value is used to keep the memory usage within certain limits.

Note: With a value of 100, each process takes less than 4 GB of RAM. We have not considered the memory taken by the main process.

sampling_min_obs Represents the minimum number of observations². When it is reached, no more simulations are performed.

4.4.1 Indels

The indels subsection of statistic contains the configuration for the analysis of indels.

```
[[indels]]
# Include/exclude indels from your analysis
include = True
# Method used to simulate indels
method = 'max'
# Number of consecutive times the indel appears to consider it falls in a_
→repetitive region
max_consecutive = 7
# Indels longer than this size will be discarded
max_size = 20
```

OncodriveFML accepts various parameters related to the indels:

include Indicates whether to include indels in the analysis or not.

- include = True: include indels in the analysis
- include = False: exclude indels from the analysis

This option *is* overridden by the *-no-indels flag* of the command line interface.

method Indicates how to simulate the indels.

- method = 'max': simulates the indels as a set of substitutions. Indels that are simulated as substitutions³ follow the same signature patter as the mutatinal signature.
- method = 'stop': simulates indels as stops. See more infomation of this option *below*.

Check the analysis of indels section to find more details.

- **max_consecutive** OncodriveFML discards indels that fall in repetitive regions. OncodriveFML considers that an indel is in a repetitive region when the same sequence of the indel appears consecutively in a genomic element a certain number of times (or even more). The maximum number of consecutive repetitions can be set with the max_consecutive option. OncodriveFML will not discard any indel due to repetitive regions if you set max_consecutive = 0.
- **max_size** Indels with a length bigger than this value are automatically discarded by the analysis, as they are assumed to be sequencing error or other artifacts.

 $^{^{2}}$ An observation is counted when a simulated value, after applying the function in method to the simulated scores, is higher than the result of applying the same function to the observed scores.

³ All indels are simulated as substitutions when method = 'max'. Indels that are in-frame are also simulated as substitutions when method = 'stop'.

Configuring indels as stops

Attention: This feature is experimental and results might be biased.

As explained in the *analysis section* OncodriveFML can be configured to simulate indels as stops.

This option should be used with care as it gives a lot of weight to the indels.

To enable this option, a number of parameters needs to be modified or added to the configuration file.

The *indels section* of the configuration file, you need to change the method to method = 'stop' and add the following parameters:

- gene_exomic_frameshift_ratio Indicates which mutations influence the *probabilities* for frameshift indels and substitutions.
 - gene_exomic_frameshift_ratio = False: the probabilities are taken from the mapped mutations discarding those whose length is multiple of 3.
 - gene_exomic_frameshift_ratio = True: probabilities are taken from the observed mutations rate in each region.
- **stops_function** The *observed* score of an indel that is computed with the method = 'stop' option is related to the score of the stops in its gene. You can decide how this relation is by choosing a function that is applied to all stops scores in the gene.
 - stops_function = 'mean': associates the indel to a value that is equal to the *mean* of all stop scores in the gene
 - stops_function = 'median': associates the indel to a value that is equal to the *median* of all stop scores in the gene
 - stops_function = 'random': associates the indel to a value that is a *random value between the maximum and the minimum* of all stop scores in the gene
 - stops_function = 'random_choice': associates the indel to a value that is a *random value* between all the possible stop scores in the gene
- minimum_number_of_stops When analysing a certain gene, OncodriveFML gets all the scores associated with the mutations that produce a stop in that gene. minimum_number_of_stops indicates the minimum number of stops that a gene is required. If the minimum is not satisfied, OncodriveFML uses the maximum possible score.

Attention: These parameters must also be adjusted for each scores file.

4.5 Settings

To configure the system where the analysis is performed OncodriveFML includes the setting section:

```
[settings]
# Number of cores to use in the analysis
cores = 6
# Random seed
seed = 1234
```

Use the cores option to indicate how many cores to use. You can comment this option in order to use all the available cores.

The command line **--cores** option *can* override this value.

Note: OncodriveFML works on shared memory systems using the multiprocessing module.

The seed option can be used to fix the random seed, to get reproducible results.

The command line **--seed** option *can* override this value.

ANALYSIS

This sections explains how OncodriveFML compute the scores for the observed mutations and how mutations are simulated.

The analysis is done for each element independently. The same number of observed mutations is simulated within the element, taking only the positions indicated in the regions file.

5.1 Observed

5.1.1 Single Nucleotide Polymorphism (SNP)

SNP mutations are the simplest to compute. To score them, OncodriveFML get the score for the corresponding alteration in the position of the mutation.

If there is not a score for that particular change, the mutation is ignored¹.

5.1.2 Multi Nucleotide Polymorphism (MNP)

MNP mutations are considered as set of SNPs. The observed value is the maximum value of all the changes produced by the MNP.

MNPs are ignored¹ when none of the changes it introduces has a score.

5.1.3 Insertion or deletion (INDEL)

Indels are scored in two different ways: as substitutions or as stops.

As substitutions Indels that fall in non-coding regions or in-frame indels in coding regions are considered as a set of substitutions. Similarly to MNP mutations, the changes produced by the indel are computed as a set of SNPs mutation and OncodriveFML assigns the indel the maximum score of those changes. In an insertion, the reference genome is compared with the indel. In a deletion, the reference genome is compared with itself but shifted a number of position equal to the length of the indel. Only the changes produced in the length of the indel are considered.

Note: If none of the changes produced by the indel has a score, the indel is ignored¹.

¹ When an observed mutation is ignored it means that it cannot be assigned a score, and thus it does not contribute to the observed scores and in the simulation the number of mutations simulated is one less for that region.

As stops Indels can be scored as stops in the analysis of coding regions and if their length is not a multiple of 3. In coding regions, a frameshift indel might cause, somewhere in the gene, a stop. This is why OncodriveFML can use this approach. The way OncodriveFML scores this type of indels is taking all the stop scores² in the gene under analysis and applying a user defined function to them. In some cases, OncodriveFML can infer a value for the scores of the stops using the mean score of all mutations in the gene. See the *configuration of indel* section for further information.

Attention: This feature is experimental. Thus, it is only available for hg19 and hg38 genomes, and it needs to be manually set up in the configuration using the configuration file.

Indels with a length higher than 20 nucleotides are ignored¹. This value can be configured in the *configuration file*.

5.2 Simulated

The same number of mutations that are observed and have a score are simulated.

To perform the simulation two arrays are computed:

- One contains the scores of all possible changes to be simulated.
- The other array contains the probabilities of each of those changes.

Using the probability array, a random sampling of the scores array is done to obtain the simulated scores.

5.2.1 Probabilities

The probability array is computed taking into account different parameters.

If only substitutions are simulated, either because the analysis excludes indels or because they are simulated as substitutions, the probabilities are:

$$p = p_{subs} * \frac{\sum_{s} p_s * f_s}{n_{substitutions}}$$

where s represents each of the signatures found in the gene in the observed mutations, p_s is the probability of a particular mutation to occur given the s signature, $n_{substitutions}$ is the total number of substitutions, and f_s is the relative frequency of a particular signature s in the gene.

However, if you are not using any signature (see *singature configuration*):

$$p = p_{subs}/n_{substitutions}$$

where $n_{substitutions}$ is the amount of substitutions in the gene.

However, if you configure indels to be analysed as *stops* things are slightly more complex. Substitution are simulated as explained above, as well as in frame indels. However, there is also a chance that a the score of one stop is selected.

The probability associated to any of the stop scores is:

$$p = \frac{1}{n_{stops}} * p_{frameshiftindel}$$

² The package BgData includes the precomputed position and alteration of the stops for the HG19 genome build. OncodriveFML makes use of it.

where $p_{frameshiftindel} + p_{subs} = 1$, and n_{stops} is the number of stop scores for that gene.

 $p_{frameshiftindel}$ represents the probability of simulating a frameshift indel in that gene, and p_{subs} represents the probability of simulating a substitution.

The probability of simulating a frameshift indel, also, depends on whether you are analysing using the whole cohort percentages or only the mutations observed in each gene.

- When using *exomic frameshift probabilities* OncodriveFML computes how many indels you observe, and how many of those fall into the region you are analysing (which should be coding). Among the mapped indels OncodriveFML distinguishes between frameshift and in-frame indels. The ratio of frameshift indels against the total amount of mutations is used to compute *p*_{frameshiftindel}.
- When using the probabilities taken from the gene:

 $p_{frameshiftindel} = \frac{n_{observed frameshiftindels}}{n_{observed mutations}}$

where $n_{observed frame shift indels}$ is the number of observed frameshift indels and $n_{observed mutations}$ is the number of observed mutations.

SIGNATURE

The signature is an array that assigns a probability to a single nucleotide mutation taking into account its context¹. It represents the chance of a certain mutation to occur within a context.

Check the different options for the signature in the *configuration file*. In short, you can choose between not using any signature, using your own signature or computing the signature from the mutations file. Additionally, signatures can be grouped into different categories (such as the sample).

The signature is computed count all the Single Nucleotide Polymorphisms in the input file, taking into account their context. The counts are used to compute a frequency $f_i = \frac{m_i}{M}$ where $M = \sum_j m_j$, and m_i represent the number of times that the mutation *i* with its context¹ has been observed.

Optionally, the signature can be corrected taking into account the frequency of trinucleotides in the reference genome. OncodriveFML introduces this feature because the distribution of triplets is not expected to be constant. When using the command line interface, OncodriveFML does this correction automatically according to the value passed in the flag --signature-correction (you can list all the options *using the help*).

Important: Signature correction is done using precomputed counts of whole genome and whole exome of HG19 reference genome.

This counts might be similar for other human genomes but ensure that correction is not done genomes of other species. Check the command line and configuration file.

More complex signatures (e.g. using only mutations that map to the regions under analysis, or normalizing by the frequency of trinucleotides in specific regions of the genome) can be computed using the bgsignature package and passed to OncodriveFML via the configuration file.

¹ The context is formed by the previous and posterior nucleotides.

SEVEN

OUTPUT

OncodriveFML generates 3 output files:

- A .tsv.gz with the analysis results
- A .png image with the most significant genes labeled.
- A . html interactive plot which can be used to search for specific genes.

The plots are only generated if the --output option is not passed or is an existing directory.

7.1 Naming

All the 3 files generated by OncodriveFML have the same name. They only differ in the extension. The name given to the files is the same as the name of the mutations file followed by -oncodrivefml and the extension.

7.2 The .tsv file

This tabulated file is the most important (as the others are just plots using the data in this one) and contains the results of the analysis.

In the file, the following columns can be found:

index Gene ID from Ensembl

MUTS number of mutations found in the dataset for that gene

MUTS_RECURRENCE number of mutations that do not occur in the same position

SAMPLES number of mutated samples in the gene

- **P_VALUE** times that the observed value is higher than or equal to the expected value, divided by the number of randomizations
- Q_VALUE pvalue corrected using the Benjamini/Hochberg correction (for samples with at least 2 samples_mut)
- **P_VALUE_NEG** times that the observed value is lower than or equal to the expected value, divided by the number of randomizations
- Q_VALUE_NEG pvalue_neg corrected using the Benjamini/Hochberg correction (for samples with at least 2
 samples_mut)
- SNP number of mutations that are Single Nucleotide Polymorphisms
- MNP number of mutations that are Multi Nucleotide Polymorphisms (two or more)

INDELS number of mutations that are insertions or deletions

SYMBOL HGNC Symbol

7.3 The plots

Both plots (.png and .html) represent the same. They are similar to Q-Q plots where in the Y axis the -log10 of the computed P-values are represented (sorted) and in the X axis the -log10 of the expected P-values are reported (sorted).

The expected P-values represent the null distribution: -log 10(i/N) where $i \in [1, N]$ and N represents the number of computed P-values.

Note: The P-values of OncodriveFML are always > 0, even when all the simulated functional impact scores are lower than the observed functional impact score. In this case, a pseudocount is added.

The genomic elements that have a lighter color in the plot are the ones for which the number of mutated sample does not reach the minimum required to perform the multiple test correction.

All the genomic regions above the red line in the plot represent those with a Q-value below 0.1. The ones between the green line and the red line are the ones with a Q-value between 0.25 and 0.1.

BEHIND THE SCENES

This section will point out some parts which might be interesting if you are running OncodriveFML yourself.

8.1 Command line interface

The command line interface of OncodriveFML overwrites some of the parameters in the configuration file.

Warning: This overwrite is performed regardless the parameter is set or not in the configuration file.

The flag --no-indels also affects the *indels configuration parameters*. Particularly, it has effect on the include option. The use of this flag discards the analysis of indels by setting include = False.

Using the --signature of the command line, set the signature configuration to method = "file" and path
= "<provided path>"

Note: Signatures provided as an external file are not normalized.

The *table below* shows the effects of the **--signature-correction** flag in the *signature configuration*:

Value	Effect in signature
wg	<pre>normalize_by_sites = 'whole_genome'</pre>
WX	<pre>normalize_by_sites = 'whole_exome'</pre>

Table 1: Effects of -signature-correction

Note: This option does not have any impact if signatures are passed with the --signature option.

8.2 BgData

OncodriveFML uses external data retrieved using the BgData package. You can download and check this data yourself. If you want to use different data, you can download the source code and modify the code to use your own data.

8.2.1 Reference genome

As March 2017 BgData includes three reference genomes: HG18, HG19 and HG38.

```
bgdata datasets/genomereference/hg38
```

If you want to use a different genome, you need to modify the code in the oncodrivefml.signature module.

8.2.2 Gene stops

OncodriveFML also uses a tabix file that contains the positions and the alterations of the gene stops.

```
bgdata datasets/genestops/hg38
```

NINE

CAVEATS

Signature computation is performed using all mutations in your input file, not only the ones that map to the region of interest.

If the scores files lacks scores for some positions or certain alterations, OncodriveFML ignores them.

If, for any reason, your signatures lack certain triplets (probability equal to 0) that are the only ones present in certain region, OncodriveFML will not compute a P-value for that region.

OncodriveFML statistical power is limited by the number of simulations performed in each regions. You can increase the number of simulations, but be aware that the time cost is exponential.

Indels do not contribute to the signatures. You can simulate indels as substitutions and perform the simulations taking the signatures into account, but be aware that the signatures are not calculated considering indels.

Depending on the values of sampling_min_obs and sampling_chunk in the configuration file the number of simulations performed for a particular genomic element can differ.

TEN

ONCODRIVEFML

10.1 oncodrivefml package

10.1.1 Subpackages

oncodrivefml.executors package

Submodules

oncodrivefml.executors.bymutation module

oncodrivefml.executors.bysample module

oncodrivefml.executors.element module

oncodrivefml.executors.sig2probs module

```
class oncodrivefml.executors.sig2probs.GroupSignature(signature, classifier)
Bases: oncodrivefml.executors.sig2probs.SubstitutionProbs
```

add_background(change)

add_observed(mutation)

property probs

property size

class oncodrivefml.executors.sig2probs.**NoSignature** Bases: oncodrivefml.executors.sig2probs.SubstitutionProbs

 $\verb+add_background(\mathit{change})$

property probs

property size

class oncodrivefml.executors.sig2probs.SubstitutionProbs(signature)
 Bases: object

add_background(change)

add_observed(mutation)

property probs

property size

oncodrivefml.executors.sig2probs.build(signature=None, classifier=None)

Module contents

10.1.2 Submodules

10.1.3 oncodrivefml.config module

This module contains code related with the configuration file (see Configuration).

Additionally, it includes other file realted code, specially from bgconfig.

oncodrivefml.config.load_configuration(config_file, override=None)
Load the configuration file and checks the format.

Parameters config_file – configuration file path

Returns configuration as a dict

Return type bgconfig.BGConfig

```
oncodrivefml.config.possible_extensions = ['.gz', '.xz', '.bz2', '.tsv', '.txt']
Some expected extensions
```

oncodrivefml.config.remove_extension_and_replace_special_characters (*file_path*) Modifies the name of a file by removing any extension in *possible_extensions* and replacing any character in *special_characters* for -.

Parameters file_path – path to a file

Returns file name modified

Return type str

```
oncodrivefml.config.special_characters = ['.', '_']
Some special characters
```

10.1.4 oncodrivefml.error module

```
exception oncodrivefml.error.OncodriveFMLError
Bases: Exception
```

10.1.5 oncodrivefml.indels module

This module contains all utilities to process insertions and deletions.

Currently 3 methods have been implemented to compute the impact of the indels.

1. As a set of substitutions ('max'):

The indel is treated as set of substitutions. It is used for non-coding regions

The functional impact of the observed mutation is the maximum of all the substitutions. The background is simulated as substitutions are.

2. As a stop ('stop'):

The indel is expected to produce a stop in the genome, unless it is a frame-shift indel. It is used for coding regions.

The functional impact is derived from the function impact of the stops of the gene. The background is simulated also as stops.

class oncodrivefml.indels.Indel (scores)

Bases: object

Methods to compute the impact of indels for the observed and the background

Parameters

- scores (Scores) functional impact per position
- **signature** (*dict*) see *signature*
- **signature_id** (*str*) classifier for the signatures
- method (str) identifies which method to use to compute the functional impact (see methods)
- **strand** (*str*) if the element being analysed has positive, negative or unknown strand (+,-,.)

compute_scores (*reference*, *alternation*, *initial_position*, *size*)

Compute the scores of all substitution between the reference and altered sequences

Parameters

- **reference** (*str*) sequence
- **alternation** (*str*) sequence
- initial_position (*int*) position where the indel occurs
- **size** (*int*) number of position to look

Returns Scores of the substitution in the indel. nan when it is not possible to compute a value.

Return type list

get_background_indel_scores_as_stops()

Returns Values of the stop scores of the gene

Return type list

- get_background_indel_scores_as_substitutions_without_signature() Return the values of scores of all possible substitutions :returns: list.
- get_indel_score_from_stop (mutation)

Compute the indel score as a stop

A function is applied to the values of the scores in the gene

Parameters mutation (dict) – a mutation object as in here

Returns Score value. nan if is not possible to compute it

Return type float

get_indel_score_max_of_subs(mutation)

Compute the score of an indel by treating each alteration as a substitution.

Parameters mutation (*dict*) – a mutation object as in *here*

Returns Maximum value of all substitutions

Return type float

get_mutation_sequences(mutation, size)

Get the reference and altered sequence of the indel along the window size

Parameters

- mutation (dict) a mutation object as in here
- **size** (*int*) window length

Returns Reference and alternated sequences

Return type tuple

static is_frameshift(size)

Parameters size (*int*) – length of the indel

Returns bool. Whether the size is multiple of 3 (in the frames have been enabled in the configuration)

is_in_repetitive_region(mutation)

Check if an indel falls in a repetitive region

Looking in the window with the indel in the middle, check if the same sequence of the indel appears at least a certain number of times specified in the configuration. The window where to look has twice the size of the indel multiplied by the number of times already mentioned.

Parameters mutation (*dict*) – a mutation object as in *here*

Returns Whether the indel falls in a repetitive region or not

Return type bool

not_found (mutation)

```
class oncodrivefml.indels.StopsScore(funct_type)
```

Bases: object

choose(x)

function(x)

mean(x)

median(x)

random(x)

oncodrivefml.indels.init_indels_module(indels_config)
Litigize the indels module

Initialize the indels module

Parameters indels_config (dict) - configuration of how to compute the impact of indels

10.1.6 oncodrivefml.load module

This module contains the methods used to load and parse the input files: elements and mutations

elements (dict) contains all the segments related to one element. The information is taken from the elements_file. Basic structure:

```
{ element_id:
    [
        {
            {CHROMOSOME': chromosome,
            'START': start_position_of_the_segment,
            'END': end_position_of_the_segment,
            'STRAND': strand (+ -> positive | - -> negative)
            'ELEMENT': element_id,
            'SEGMENT': segment_id,
            'SYMBOL': symbol_id
        }
    ]
}
```

mutations (dict) contains all the mutations for each element. Most of the information is taken from the mutations_file but the *element_id* and the *segment* that are taken from the **elements**. More information is added during the execution. Basic structure:

mutations_data (dict) contains the *mutations dict* and some metadata information about the mutations. Currently, the number of substitutions and indels. Basic structure:

```
{
    'data':
        {
            `mutations dict`_
        },
    'metadata':
        {
            'snp': amount of SNP mutations
            'mnp': amount of MNP mutations
            'mnp_length': total length of the MNP mutations
            'indel': amount of indels
        }
}
```

oncodrivefml.load.build_regions_tree (regions)

Generates a binary tree with the intervals of the regions

```
Parameters regions (dict) – segments grouped by elements.
```

Returns

for each chromosome, it get one IntervalTree which is a binary tree. The leafs are intervals [low limit, high limit) and the value associated with each interval is the tuple (element, segment). It can be interpreted as:

```
{ chromosome:
    (start_position, end_position +1): (element, segment)
}
```

Return type dict of IntervalTree

```
oncodrivefml.load.mutations (file, blacklist=None, metadata_dict=None, indels_max_size=None)
Parsed the mutations file
```

Parameters

- file mutations file (see OncodriveFML)
- metadata_dict (dict) dict that the function will fill with useful information
- **blacklist** (*optional*) file with blacklisted samples (see OncodriveFML). Defaults to None.
- indels_max_size (*int*, *optional*) max size of indels. Indels with logner sizes will be discarded.

Yields One line from the mutations file as a dictionary. Each of the inner elements of *mutations*

From the elements and variants file, get dictionaries with the segments grouped by element ID and the mutations grouped in the same way, as well as some information related to the mutations.

Parameters

- **variants_file** mutations file (see OncodriveFML)
- **elements_file elements** file (see OncodriveFML)
- **blacklist** (*optional*) file with blacklisted samples (see OncodriveFML). Defaults to None. If the blacklist option is passed, the mutations are not loaded from a pickle file.
- **indels_max_size** (*int*, *optional*) max size of indels. Indels with logner sizes will be discarded.

Returns

mutations and elements

Elements: elements dict

Mutations: mutations data dict

Return type tuple

The process is done in 3 steps:

- load_regions()
- build_regions_tree().
- 3. each mutation (mutations ()) is associated with the right element ID

oncodrivefml.load.snp(file, blacklist=None)
Load only SNP

10.1.7 oncodrivefml.main module

10.1.8 oncodrivefml.mtc module

Module containing functions related to multiple test correction

```
oncodrivefml.mtc.multiple_test_correction (results, num_significant_samples=2)
Performs a multiple test correction on the analysis results
```

Parameters

- results (dict) dictionary with the results
- **num_significant_samples** (*int*) mininum samples that a gene must have in order to perform the correction

Returns DataFrame. DataFrame with the q-values obtained from a multiple test correction

10.1.9 oncodrivefml.oncodrivefml module

10.1.10 oncodrivefml.reference module

This module contains information related to the reference genome.

oncodrivefml.reference.change_build(*build*) Modify the default build fo the reference genome

Parameters build (*str*) – genome reference build

```
oncodrivefml.reference.get_build()
```

oncodrivefml.reference.get_ref (chromosome, start, size=1)
Gets a sequence from the reference genome

Parameters

- chromosome (*str*) chromosome
- **start** (*int*) start position where to look
- **size** (*int*) number of bases to retrieve

Returns str. Sequence from the reference genome

oncodrivefml.reference.get_ref_triplet (chromosome, start)

Parameters

- chromosome (*str*) chromosome identifier
- **start** (*int*) starting position

Returns 3 bases from the reference genome

Return type str

```
oncodrivefml.reference.ref_build = 'hg38'
Build of the Reference Genome
```

oncodrivefml.reference.reverse_complementary_sequence (seq)

10.1.11 oncodrivefml.scores module

This module contains the methods associated with the scores that are assigned to the mutations.

The scores are read from a file.

Information about the stop scores.

As of December 2016, we have only measured the stops using CADD1.0.

The stops of a gene retrieved only if there are ast least 3 stops in the regions being analysed. If not, a formula is applied to derived the value of the stops from the rest of the values.

Note: This formula was obtained using the CADD scores of the coding regions. Using a different regions or scores files will make the function to return totally nonsense values.

```
class oncodrivefml.scores.PackScoresReader(conf)
    Bases: object
```

```
BIT_TO_REF = { (0, 0, 0): '?', (0, 0, 1): 'T', (0, 1, 0): 'A', (0, 1, 1): 'C', (1,
SCORE_ALT = {'A': 'CGT', 'C': 'AGT', 'G': 'ACT', 'T': 'ACG'}
SCORE_ORDER = {'A': {'C': 0, 'G': 1, 'T': 2}, 'C': {'A': 0, 'G': 1, 'T': 2}, 'G': {'A'
STRUCT_SIZE = 6
```

get (chromosome, start, stop, *args, **kwargs)

unpack (block)

```
exception oncodrivefml.scores.ReaderError(msg)
Bases: Exception
```

```
exception oncodrivefml.scores.ReaderGetError(chr, start, end)
Bases: oncodrivefml.scores.ReaderError
```

```
class oncodrivefml.scores.ScoreValue(ref, alt, value, change)
    Bases: tuple
```

Tuple that contains the reference, the alteration, the score value and the triplets

Parameters

- **ref** (*str*) reference base
- **alt** (*str*) altered base
- **value** (*float*) score value of that substitution
- **change** (*str*) reference triplet > alt (e.g. ACG>T)

property alt

Alias for field number 1

property change

Alias for field number 3

property ref

Alias for field number 0

property value

Alias for field number 2

```
class oncodrivefml.scores.Scores (element: str, segments: list, config: dict)
Bases: object
```

Parameters

- element (*str*) element ID
- **segments** (*list*) list of the segments associated to the element
- **config** (*dict*) configuration

scores_by_pos

for each positions get all possible changes, and for each change the triplets

```
{ position:
    [
        ScoreValue(
            ref,
             alt_1,
             value,
             change
        ),
        ScoreValue(
             ref,
             alt_2,
             value,
             change
        ),
        ScoreValue(
             ref,
             alt_3,
             value,
             change
        )
    ]
```

Type dict

get_all_positions () \rightarrow List[int] Get all positions in the element

Returns list of positions

Return type list of int

get_score_by_position (*position: int*) \rightarrow List[oncodrivefml.scores.ScoreValue] Get all ScoreValue objects that are asocated with that position

Parameters position (*int*) – position

Returns list of all ScoreValue related to that positon

Return type list of ScoreValue

property stop_scores

```
class oncodrivefml.scores.ScoresTabixReader(conf)
    Bases: object
```

get (chromosome, start, stop, element=None)

oncodrivefml.scores.init_scores_module(conf, stops_required=False)

10.1.12 oncodrivefml.signature module

This module contains information related with the signature.

The signature is a way of assigning probabilities to certain mutations that have some relation amongst them (e.g. cancer type, sample...). This relation is identified by the **signature_id**.

The classifier parameter in the *configuration* of the signature specifies which column of the mutations file (MUTATIONS_HEADER) is used as the identifier for the different signature groups. If not provided, all mutations contribute to one global signature.

The probabilities are taken only from substitutions. For them, the two bases that surround the mutated one are taken into account. This is called the triplet. For a certain mutation in a position x the reference triplet is the base in the reference genome in position x-1, the base in x and the base in the x+1. The altered triplet of the same mutation is equal for the bases in x-1 and x+1 but the base in x is the one observed in the mutation.

signature (dict)

```
{ signature_id:
    {
        (ref_triplet, alt_triplet): prob
    }
}
```

oncodrivefml.signature.collapse(counts)

oncodrivefml.signature.compute (mutations, method, classifier=None, normalize=None)

```
oncodrivefml.signature.load(file)
```

10.1.13 oncodrivefml.stats module

This modules contains different statistical methods used to compare the observed and the simulated scores

```
class oncodrivefml.stats.ArithmeticMean
    Bases: object
```

```
Dases. OD ject
```

static calc (*values*) Computes the arithmetic mean

Parameters values (list, array) - array of values

Returns mean value

Return type float

```
static calc_observed(values, observed)
```

Measure how many times the mean of the values is higher than the mean of the observed values

Parameters

- **values** (array) m x n matrix with scores (m: number of randomizations; n: number of mutations)
- **observed** (list, array) n size vector with the observed scores (n: number of mutations)

Returns

the number of times that the mean value of a randomization is greater or equal than the mean observed value

(as int) and the number of times that the mean value of a randomization is equal or lower than the mean observed value (as int).

Return type tuple

class oncodrivefml.stats.ArithmeticMeanHeteroscedasticScores Bases: object

static calc_observed(values, observed)

```
class oncodrivefml.stats.GeometricMean
```

Bases: object

The geometric mean used is not the standard.

$$\left(\prod_{i=1}^{n} (x_i+1)\right)^{1/n} - 1 = \sqrt[n]{(x_1+1)(x_2+1)\cdots(x_n+1)} - 1$$

static calc(values)

Computes the geometric mean of a set of values.

Parameters values (list, array) - set of values

Returns geometric mean (array): geometric mean by columns (if the input is a matrix)

Return type (float)

static calc_observed(values, observed)

Measure how many times the geometric mean of the values is higher than the geometric mean of the observed values

Parameters

- **values** (array) m x n matrix with scores (m: number of randomizations; n: number of mutations)
- **observed** (list, array) n size vector with the observed scores (n: number of mutations)

Returns

the number of times that the mean value of a randomization is greater or equal than the mean observed value (as int) and the number of times that the mean value of a randomization is equal or lower than the mean observed value (as int).

Return type tuple

class oncodrivefml.stats.**Maximum** Bases: object

static calc(values)

static calc_observed(values, observed)

10.1.14 oncodrivefml.store module

This module contains the methods used to store the results.

3 different types of output are available:

- tsv file
- **png** graph: uses the *tsv* file and matplotlib
- html graph: uses the *tsv* file and bokeh

```
class oncodrivefml.store.QQPlot(input_file, cutoff=True, rename_fields=None, ex-
tra_fields=None)
```

Bases: object

Parameters

- **input_file** tsv file with the data
- **cutoff** (bool) add cutoffs to the figure
- **rename_fields** (*dict*) column names from the input file can be renamed providing a dictionary {old_name : new_name}
- **extra_fields** (*list*) list of column names that want to be passed to the figure data. Need for example to search by them.

add_search_widget (fields)

Add text input for each field.

```
Parameters fields (str or list) - list of fields to do a search.
```

add_tooltip()

Adds tooltip to show the parameters of each glyph in the figure

add_tooltip_enhanced()

The tooltip is shown via JavaScript to avoid been block in areas with a high density of points

show (output_path, showit=True, notebook=False)

Show the figure

Parameters

- **output_path** file where to store the figure
- **showit** (*bool*) the figure is displayed (widgets and the like are not shown) or is fully saved. Defaults to True.
- **notebook** (bool) if is is called form a notebook or not. Defaults to False.

oncodrivefml.store.add_symbol(df)

oncodrivefml.store.eliminate_duplicates(df)

oncodrivefml.store.store_html (input_file, output_path)

Create the QQPlot and save it.

Parameters

- input_file tsv filw with the data
- **output_path** file where to store the graph
- **showit** (bool) defaults to False. See show().

```
oncodrivefml.store.store_png(input_file, output_file, showit=False)
```

Creates a figure from the resutls.

Parameters

- input_file tsv file with the results
- **output_file** file where to store the figure
- **showit** (*bool*) calls show() before returning. Defaults to False.

oncodrivefml.store.store_tsv(results, result_file)

Saves the results in a tsv file sorted by pvalue

Parameters

- **results** (DataFrame) results of the analysis
- **result_file** file where to store the results

10.1.15 oncodrivefml.utils module

This module contains some useful methods

```
oncodrivefml.utils.defaultdict_list()
```

Shortcut

```
Returns defaultdict of list
```

```
oncodrivefml.utils.executor_run (executor)
    Method to call the run method
```

Parameters executor (ElementExecutor) -

Returns run()

```
oncodrivefml.utils.exists_path(path)
```

oncodrivefml.utils.loop_logging(iterable, size=None, step=1)
Loop through an iterable object displaying messages using info()

Parameters

- iterable -
- **size** (*int*) Defaults to None.
- **step** (*int*) **Defaults** to 1.

Yields The iterable element

10.1.16 oncodrivefml.walker module

- 10.1.17 oncodrivefml.walker_cython module
- 10.1.18 oncodrivefml.walker_cython module
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