Package 'mSigAct'

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Title Mutational Signature Activity Analysis ('mSigAct')

devtools,

```
Version 3.0.0
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Description Analyze the ``activities" of mutational signatures in one or more mutational spectra.
     'mSigAct' stands for mutational Signature Activity. mSigAct uses a maximum likelihood approach
     to estimate (conservatively) whether there is evidence that a particular set of mutational
     signatures is present in a spectrum. It can also determine a *minimal* subset of signatures
     needed to plausibly reconstruct an observed spectrum.
License GPL-3
URL https://github.com/steverozen/mSigAct
BugReports https://github.com/steverozen/mSigAct/issues
Encoding UTF-8
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Depends R (>= 4.0),
RoxygenNote 7.2.3
biocViews
Imports cosmicsig,
     dplyr,
     gtools,
     ICAMS (>= 3.0.6),
     nloptr,
     mSigTools,
     PCAWG7 (>= 0.1.3),
     philentropy,
     quadprog,
     stats,
     sets,
     tibble,
     utils
Remotes github::steverozen/ICAMS@v3.0.6-branch,
     github::steverozen/PCAWG7@v0.1.3-branch
Suggests BSgenome. Hsapiens. 1000 genomes. hs37d5,
```

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```
htmlwidgets, ipc, knitr, profvis, rmarkdown, testthat (>= 2.1.0), usethis
```

R topics documented:

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cossim

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Cosine similarity with useful argument types

Description

Cosine similarity with useful argument types

Usage

```
cossim(v1, v2)
```

Arguments

v1 A vector or single-column matrix v2 A vector or single-column matrix

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DefaultManyOpts

Set default options for many functions, especially nloptr

Description

Set default options for many functions, especially nloptr

Usage

```
DefaultManyOpts(likelihood.dist = "neg.binom", spectra = NULL)
```

Arguments

likelihood.dist

The probability distribution used to calculate the likelihood, can be either "multi-nom" (multinomial distribution) or "neg.binom" (negative binomial distribution).

spectra

The catalog (matrix) to analyze. This could be an ICAMS catalog or a numerical matrix

Value

A list with the following elements

global.opts A sub-list with several options for nloptr, q.v., for the global optimization phase.

local.opts A sub-list with several options for nloptr, q.v., for the local optimization phase.

nbinom.size Only appearing if likelihood.dist = "neg.binom". The dispersion parameter for the negative binomial distribution; smaller is more dispersed. See NegBinomial.

trace If > 0 print progress messages.

global_eval_f The objective function for the global optimization phase.

 $local_eval_f$ The objective function for the local optimization phase.

local_eval_g_ineq The inequality constraint function for the local optimization phase.

likelihood.dist The probability distribution used to calculate the likelihood.

```
my.opts <- DefaultManyOpts()
my.opts$trace <- 10</pre>
```

```
deprecated_SparseAssignActivity
```

Find known signatures that can most sparsely reconstruct each spectrum in a catalog.

Description

Find known signatures that can most sparsely reconstruct each spectrum in a catalog.

Usage

```
deprecated_SparseAssignActivity(
  spectra,
 sigs,
  output.dir,
 max.level = 5,
 p.thresh = 0.05,
 m.opts = DefaultManyOpts(),
 num.parallel.samples = 5,
 mc.cores.per.sample = min(20, 2^max.level),
 progress.monitor = NULL,
  seed = NULL,
 max.subsets = 1000,
 drop.low.mut.samples = TRUE,
 use.sig.presence.test = FALSE
)
```

Arguments

```
The spectra (multiple spectra) to be reconstructed.
spectra
                 A numerical matrix, possibly an ICAMS catalog.
sigs
output.dir
                 Directory path to save the output file.
                 The maximum number of signatures to try removing.
max.level
                 If the p value for a better reconstruction with a set of signatures (as opposed to
p.thresh
                 without that set of signatures) is > than this argument, then we can use exposures
                 without this set.
m.opts
                 See DefaultManyOpts.
num.parallel.samples
                 The (maximum) number of samples to run in parallel. On Microsoft Windows
                 machines it is silently changed to 1. Each sample in turn can require multiple
                 cores, as governed by mc.cores.per.sample.
mc.cores.per.sample
                 The maximum number of cores to use for each sample. On Microsoft Windows
```

machines it is silently changed to 1.

```
progress.monitor
```

Function called at the start of each new level (number of signatures to try excluding). Must take named arguments value and detail, and no others. Designed for a AsyncProgress progress bar function.

seed

Random seed; set this to get reproducible results. (The numerical optimization is in two phases; the first, global phase might rarely find different optima depending on the random seed.)

max.subsets

This argument provides a way to heuristically limit the amount of time spent by this function. Larger values of this argument will tend to allow longer running times. The algorithm successively tries to remove all subsets of 1 signature, 2 signatures, 3 signatures, etc., down to max.level. (Not every subset is tested at each level; if a subset was already found to be necessary the algorithm does not test supersets of that subset.) If at any level the algorithm needs to test more than max.subsets this function will not proceed.

drop.low.mut.samples

Whether to exclude low mutation samples from the analysis. If TRUE, samples with SBS total mutations less than 100, DBS or ID total mutations less than 25 will be dropped.

use.sig.presence.test

Whether to use signature presence test first to filter out those signatures that are not needed in the reconstruction of the spectrum.

Value

A list with the elements:

- proposed.assignment: The most sparse set of signatures that can plausibly explain spectra.
- proposed.reconstruction: The reconstruction based on sparse assignment.
- reconstruction.distances: Various distances and similarities between spectra and proposed.reconstruction.
- all.tested: All tested possible ways to reconstruct each sample in spectra.
- alt.solutions: A tibble showing all the alternative solutions that are statistically as good as the proposed.assignment that can plausibly reconstruct spectra.
- time.for.assignment: Value from system.time for running SparseAssignActivity for each sample in spectra.
- error.messages: Only appearing if there are errors running SparseAssignActivity.

The elements proposed.assignment, proposed.reconstruction, reconstruction.distances, all.tested, time.for.assignment will be NULL if the algorithm could not find the optimal reconstruction or there are errors coming out for all samples.

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```
p.thresh = 0.05 / ncol(sigs.to.use),
num.parallel.samples = 2,
mc.cores.per.sample = 30,
seed = 2561)
## End(Not run)
```

ExposureProportions

Return the proportions of tumors of a given cancer type that have a particular signature

Description

Return the proportions of tumors of a given cancer type that have a particular signature

Usage

```
ExposureProportions(
  mutation.type,
  cancer.type,
  all.sigs = NULL,
  drop.sigs.no.info = TRUE,
  must.include = character(),
  must.include.prop = 0.1
)
```

Arguments

```
mutation.type
                 A character string, one of "SBS96", "SBS192", "ID", "DBS78".
cancer.type
                 A character string. For some common cancer types, see CancerTypes for more
                 details.
                 An optional matrix of known signatures, with column names being signatures
all.sigs
                 ids. Only used to drop signatures not present in all.sigs.
drop.sigs.no.info
                 If TRUE, drop signatures not present in the column names of all.sigs.
must .include A character vector of signature IDs that must be included, even if they have
                 not previously been observed in that cancer type. The associated proportion is
                 specified by must.include.prop.
must.include.prop
                 The value used for the expected proportion of signatures in must.include
                 but not previously observed in the given cancer.type.
```

Value

A numerical vector of the proportion of tumors of type cancer.type with each signature for those signatures observed in cancer.type. The names are the signature ids.

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Examples

MAPAssignActivity Find Maximum A Posteriori (MAP) assignment of signature exposures that explain multiple spectra

Description

This function also can do sparse assignment by specifying use.sparse.assign = TRUE.

Usage

```
MAPAssignActivity(
 spectra,
  sigs,
  output.dir,
  sigs.presence.prop = NULL,
 max.level = 5,
  p.thresh = 0.05,
 m.opts = DefaultManyOpts(),
 num.parallel.samples = 5,
 mc.cores.per.sample = min(20, 2^max.level),
 progress.monitor = NULL,
 seed = NULL
 max.subsets = 1000,
 use.sparse.assign = FALSE,
 use.forward.search = FALSE,
  drop.low.mut.samples = TRUE,
 use.sig.presence.test = FALSE,
  save.files = TRUE
)
```

without this set.

Arguments

| spectra | The spectra (multiple spectra) to be reconstructed. | |
|--------------------|---|--|
| sigs | A numerical matrix, possibly an ICAMS catalog. | |
| output.dir | Directory path to save the output file. | |
| sigs.presence.prop | | |
| | The proportions of samples that contain each signature. A numerical vector (values between 0 and 1), with names being a subset of colnames (sigs). | |
| | See ExposureProportions for more details. | |
| max.level | The maximum number of signatures to try removing. | |
| p.thresh | If the p value for a better reconstruction with a set of signatures (as opposed to without that set of signatures) is > than this argument, then we can use exposures | |

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m.opts See DefaultManyOpts.

num.parallel.samples

The (maximum) number of samples to run in parallel. On Microsoft Windows machines it is silently changed to 1. Each sample in turn can require multiple cores, as governed by mc.cores.per.sample.

mc.cores.per.sample

The maximum number of cores to use for each sample. On Microsoft Windows machines it is silently changed to 1.

progress.monitor

Function called at the start of each new level (number of signatures to try excluding). Must take named arguments value and detail, and no others. Designed for a AsyncProgress progress bar function.

seed

Random seed; set this to get reproducible results. (The numerical optimization is in two phases; the first, global phase might rarely find different optima depending on the random seed.)

max.subsets

This argument provides a way to heuristically limit the amount of time spent by this function. Larger values of this argument will tend to allow longer running times. The algorithm successively tries to remove all subsets of 1 signature, 2 signatures, 3 signatures, etc., down to max.level. (Not every subset is tested at each level; if a subset was already found to be necessary the algorithm does not test supersets of that subset.) If at any level the algorithm needs to test more than max.subsets this function will not proceed.

use.sparse.assign

Whether to use sparse assignment. If TRUE, arguments designed for Maximum A Posteriori assignment such as sigs.presence.prop will be ignored.

use.forward.search

Whether to use forward search to find the minimal number of signatures to optimally reconstruct the spectrum.

drop.low.mut.samples

Whether to exclude low mutation samples from the analysis. If TRUE, samples with SBS total mutations less than 100, DBS or ID total mutations less than 25 will be dropped.

use.sig.presence.test

Whether to use signature presence test first to filter out those signatures that are not needed in the reconstruction of the spectrum.

save.files If TRUE save several files for each input sample in a directory named after the sample.

Value

A list with the elements:

- proposed.assignment: Proposed signature assignment for spectra with the highest MAP found. If use.sparse.assign = TRUE, this will be the most sparse set of signatures that can plausibly explain spectra.
- proposed.reconstruction: Proposed reconstruction of spectra based on MAP. If use.sparse.assign = TRUE, this will be the reconstruction based on sparse assignment.
- reconstruction.distances: Various distances and similarities between spectra and proposed.reconstruction.
- all.tested: All tested possible ways to reconstruct each sample in spectra.

- alt.solutions: A tibble showing all the alternative solutions that are statistically as good as the proposed.assignment that can plausibly reconstruct spectra.
- time.for.assignment: Value from system.time for running MAPAssignActivity1 for each sample in spectra.
- error.messages: Only appearing if there are errors running MAPAssignActivity.

The elements proposed.assignment, proposed.reconstruction, reconstruction.distances, all.tested, time.for.assignment will be NULL if the algorithm could not find the optimal reconstruction or there are errors coming out for all samples.

Examples

```
## Not run:
# This is a long running example unless parallel computing is supported on your machine
indices <- grep("Lung-AdenoCA", colnames(PCAWG7::spectra$PCAWG$SBS96))</pre>
spectra <- PCAWG7::spectra$PCAWG$SBS96[, indices[1:2], drop = FALSE]</pre>
SBS96.sigs <- cosmicsig::COSMIC_v3.2$signature$GRCh37$SBS96
sigs.prop <- ExposureProportions(mutation.type = "SBS96",</pre>
                                  cancer.type = "Lung-AdenoCA")
sigs.to.use <- SBS96.sigs[, names(sigs.prop), drop = FALSE]</pre>
MAP.out <- MAPAssignActivity(spectra = spectra,
                              sigs = sigs.to.use,
                              sigs.presence.prop = sigs.prop,
                              output.dir = file.path(tempdir(), "Lung-AdenoCA"),
                              max.level = ncol(sigs.to.use) - 1,
                              p.thresh = 0.05 / ncol(sigs.to.use),
                              num.parallel.samples = 2,
                              mc.cores.per.sample = 30,
                              seed = 2561)
## End(Not run)
```

PresenceAssignActivity

Find minimal set of signatures that can explain multiple spectra by first using signature presence test

Description

Find minimal set of signatures that can explain multiple spectra by first using signature presence test

Usage

```
PresenceAssignActivity(
   spectra,
   sigs,
   output.dir,
   p.thresh = DefaultPThresh(sigs),
   m.opts = DefaultManyOpts(spectra = spectra),
   num.parallel.samples = 1,
   mc.cores.per.sample = 1,
```

```
seed = 123,
drop.low.mut.samples = FALSE,
save.files = TRUE
)
```

Arguments

spectra The spectra (multiple spectra) to be reconstructed.

sigs A numerical matrix, possibly an ICAMS catalog.

 $\hbox{\tt output.dir} \quad \hbox{\tt Directory path to save the output file.}$

p.thresh If the p value for a better reconstruction with a set of signatures (as opposed to

without that set of signatures) is > than this argument, then we can use exposures

without this set.

num.parallel.samples

The (maximum) number of samples to run in parallel. On Microsoft Windows machines it is silently changed to 1. Each sample in turn can require multiple cores, as governed by mc.cores.per.sample.

mc.cores.per.sample

The maximum number of cores to use for each sample. On Microsoft Windows

machines it is silently changed to 1.

Random seed; set this to get reproducible results. (The numerical optimiza-

tion is in two phases; the first, global phase might rarely find different optima

depending on the random seed.)

drop.low.mut.samples

Whether to exclude low mutation samples from the analysis. If TRUE, samples with SBS total mutations less than 100, DBS or ID total mutations less than 25

will be dropped.

sample.

Value

A list with the elements:

- proposed.assignment: The proposed set of signatures that can plausibly explain spectra.
- proposed.reconstruction: The reconstruction based on proposed.assignment.
- reconstruction.distances: Various distances and similarities between spectra and proposed.reconstruction.
- time.for.assignment: Value from system.time for running PresenceAssignActivity for each sample in spectra.
- error.messages: Error messages running PresenceAssignActivity.

```
## Not run:
# This is a long running example unless parallel computing is supported on your machine
indices <- grep("Lung-AdenoCA", colnames(PCAWG7::spectra$PCAWG$SBS96))
spectra <- PCAWG7::spectra$PCAWG$SBS96[, indices[1:2], drop = FALSE]
SBS96.sigs <- cosmicsig::COSMIC_v3.2$signature$GRCh37$SBS96</pre>
```

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ReconstructSpectrum

Given signatures (sigs) and exposures (exp), return a spectrum or spectra

Description

Given signatures (sigs) and exposures (exp), return a spectrum or spectra

Usage

```
ReconstructSpectrum(sigs, exp, use.sig.names = FALSE)
```

Arguments

Signature as a matrix or data frame, with each row one mutation type (e.g. CCT > CAT or CC > TT) and each column a signature.

exp

The exposures for one or more samples as a matrix or data.frame, with each row a signature and each column a sample.

use.sig.names

If TRUE check that rownames (exp) is a subset of colnames (sigs), and use only the columns in sigs that are present in exp.

Details

Does not care or check if colSums (sigs) == 1. Error checking is minimal since this function is called often.

Value

The matrix product sigs % * % exp after some error checking.

```
SignaturePresenceTest
```

Test whether a given signature is plausibly present in a catalog.

Description

Test whether a given signature is plausibly present in a catalog.

Usage

```
SignaturePresenceTest(
   spectra,
   sigs,
   target.sig.index,
   m.opts = DefaultManyOpts(likelihood.dist = "multinom"),
   seed = NULL,
   mc.cores = 2
)
```

Arguments

| spectra | The catalog (matrix) to analyze. This could be an ${\tt ICAMS}$ catalog or a numerical matrix. | | | |
|------------------|--|--|--|--|
| sigs | A catalog of signatures from which to choose. This could be and ${\tt ICAMS}$ catalog or a numerical matrix. | | | |
| target.sig.index | | | | |
| | The index of the signature the presence of which we want to test. It can also be the signature id (e.g. "SBS22"). | | | |
| m.opts | See DefaultManyOpts. | | | |
| seed | Random seed; set this to get reproducible results. (The numerical optimization is in two phases; the first, global phase might rarely find different optima depending on the random seed.) | | | |
| mc.cores | Number of cores to use. Always silently changed to 1 on Microsoft Windows. | | | |

Value

A list of test results for each sample in spectra. Each sublist contains the following elements:

- loglh.with: The maximum log likelihood of the reconstructed spectrum using all the signatures
- loglh.without: The maximum log likelihood of the reconstructed spectrum without the target signature.
- statistic: Likelihood ratio test statistic.
- chisq.p: P-value of the likelihood ratio test. The null hypothesis is we can plausibly reconstruct the spectrum without the target signature.
- exp.with: The exposure using all the signatures which generates the maximum log likelihood loglh.with.
- exp.without: The exposure not using the target signature which generates the maximum log likelihood loglh.without.

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