# Package 'abseqR'

November 4, 2025

Type Package

**Title** Reporting and data analysis functionalities for Rep-Seq datasets of antibody libraries

Version 1.29.0

**Description** AbSeq is a comprehensive bioinformatic pipeline for the analysis of sequencing datasets generated from antibody libraries and abseqR is one of its packages. abseqR empowers the users of abseqPy (https://github.com/malhamdoosh/abseqPy) with plotting and reporting capabilities and

allows them to generate interactive HTML reports for the convenience of viewing and sharing with other researchers. Additionally, abseqR extends abseqPy to compare multiple repertoire analyses and perform further downstream analysis on its output.

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**Encoding** UTF-8

LazyData true

**Depends** R (>= 3.5.0)

Imports ggplot2, RColorBrewer, circlize, reshape2, VennDiagram, plyr, flexdashboard, BiocParallel (>= 1.1.25), png, grid, gridExtra, rmarkdown, knitr, vegan, ggcorrplot, ggdendro, plotly, BiocStyle, stringr, utils, methods, grDevices, stats, tools, graphics

VignetteBuilder knitr

RoxygenNote 6.1.0

Collate 'accessors-AbSeq.R' 'AbSeqCRep.R' 'util.R' 'distributions.R' 'upstreamAnalysis.R' 'productivityAnalysis.R' 'primerAnalysis.R' 'diversityAnalysis.R' 'annotationAnalysis.R' 'abundanceAnalysis.R' 'plotter.R' 'AbSeqRep.R' 'abseqReport.R' 'statistics.R' 'pairwise.R'

**SystemRequirements** pandoc (>= 1.19.2.1)

URL https://github.com/malhamdoosh/abseqR

BugReports https://github.com/malhamdoosh/abseqR/issues

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+, AbSeqCRep, AbSeqCRep-method

Combines 2 AbSeqCRep objects together for comparison

#### **Description**

Combines 2 AbSeqCRep objects together for comparison

## Usage

```
## S4 method for signature 'AbSeqCRep,AbSeqCRep'
e1 + e2
```

#### **Arguments**

e1 AbSeqCRep.e2 AbSeqCRep.

#### Value

AbSeqCRep object. Calling abseqR's functions on this object will always result in a comparison.

# See Also

abseqReport returns a list of AbSeqReps

```
# Use example data from abseqR as abseqPy's output, substitute this
# with your own abseqPy output directory
abseqPyOutput <- tempdir()
file.copy(system.file("extdata", "ex", package = "abseqR"), abseqPyOutput, recursive=TRUE)
samples <- abseqReport(file.path(abseqPyOutput, "ex"), report = 0)
# The provided example data has PCR1, PCR2, and PCR3 samples contained within
# pcr12 and pcr13 are instances of AbSeqCRep
pcr12 <- samples[["PCR1"]] + samples[["PCR2"]]
pcr13 <- samples[["PCR1"]] + samples[["PCR3"]]</pre>
```

```
# all_S is also an instance of AbSeqCRep
all_S <- pcr12 + pcr13

# you can now call the report function on this object
# report(all_S)  # uncomment this line to execute report</pre>
```

+, AbSeqCRep, AbSeqRep-method

Combines a AbSeqCRep object with a AbSeqRep object together for comparison

### **Description**

Combines a AbSeqCRep object with a AbSeqRep object together for comparison

# Usage

```
## S4 method for signature 'AbSeqCRep,AbSeqRep'
e1 + e2
```

#### **Arguments**

- e1 AbSeqCRep. e2 AbSeqRep.
- Value

AbSeqCRep object. Calling abseqR's functions on this object will always result in a comparison.

#### See Also

abseqReport returns a list of AbSeqReps

```
# Use example data from abseqR as abseqPy's output, substitute this
# with your own abseqPy output directory
abseqPyOutput <- tempdir()
file.copy(system.file("extdata", "ex", package = "abseqR"), abseqPyOutput, recursive=TRUE)
samples <- abseqReport(file.path(abseqPyOutput, "ex"), report = 0)
# The provided example data has PCR1, PCR2, and PCR3 samples contained within
# pcr12 is an instance of AbSeqCRep
pcr12 <- samples[["PCR1"]] + samples[["PCR2"]]
# pcr3 is instance of AbSeqRep
pcr3 <- samples[["PCR3"]]
# pcr123 is an instance of AbSeqCRep
pcr123 <- pcr12 + pcr3</pre>
```

```
# you can now call the report function on this object
# report(pcr123)  # uncomment this line to execute report
```

+, AbSeqRep, AbSeqCRep-method

Combines a AbSeqRep object with a AbSeqCRep object together for comparison

#### **Description**

Combines a AbSeqRep object with a AbSeqCRep object together for comparison

# Usage

```
## S4 method for signature 'AbSeqRep,AbSeqCRep'
e1 + e2
```

#### **Arguments**

e1 AbSeqRep.e2 AbSeqCRep.

#### Value

AbSeqCRep object. Calling abseqR's functions on this object will always result in a comparison.

#### See Also

abseqReport returns a list of AbSeqReps

```
# Use example data from abseqR as abseqPy's output, substitute this
# with your own abseqPy output directory
abseqPyOutput <- tempdir()
file.copy(system.file("extdata", "ex", package = "abseqR"), abseqPyOutput, recursive=TRUE)
samples <- abseqReport(file.path(abseqPyOutput, "ex"), report = 0)
# The provided example data has PCR1, PCR2, and PCR3 samples contained within
# pcr1 is an instance of AbSeqRep
pcr1 <- samples[["PCR1"]]
# pcr23 is instance of AbSeqCRep
pcr23 <- samples[["PCR2"]] + samples[["PCR3"]]
# pcr123 is an instance of AbSeqCRep
pcr123 <- pcr1 + pcr23
# you can now call the report function on this object
# report(pcr123) # uncomment this line to execute report</pre>
```

+,AbSeqRep,AbSeqRep-method

Combines 2 AbSeqRep objects together for comparison

# **Description**

Combines 2 AbSeqRep objects together for comparison

# Usage

```
## S4 method for signature 'AbSeqRep,AbSeqRep'
e1 + e2
```

#### **Arguments**

- e1 AbSeqRep object.
- e2 AbSeqRep object.

#### Value

AbSeqCRep object. Calling abseqR's functions on this object will always result in a comparison.

#### See Also

abseqReport returns a list of AbSeqReps

```
# Use example data from abseqR as abseqPy's output, substitute this
# with your own abseqPy output directory
abseqPyOutput <- tempdir()
file.copy(system.file("extdata", "ex", package = "abseqR"), abseqPyOutput, recursive=TRUE)
samples <- abseqReport(file.path(abseqPyOutput, "ex"), report = 0)
# The provided example data has PCR1, PCR2, and PCR3 samples contained within
# pcr1 and pcr2 are instances of AbSeqRep
pcr1 <- samples[["PCR1"]]
pcr2 <- samples[["PCR2"]]
# pcr12 is an instance of AbSeqCRep
pcr12 <- pcr1 + pcr2
# you can now call the report function on this object
# report(pcr12) # uncomment this line to execute report</pre>
```

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.abundanceAnalysis

Conducts abundance analysis

# Description

Conducts abundance analysis

# Usage

```
.abundanceAnalysis(abundanceDirectories, abunOut, sampleNames,
  combinedNames, mashedNames, skipDgene = FALSE, .save = TRUE)
```

#### **Arguments**

abundanceDirectories

list type. List of sample directories

abunOut string type. Output directory

sampleNames vector type. 1-1 correspondence with abundanceDirectories

combinedNames string type. Title "combined" sample names

mashedNames string type. File "mashed" names - avoid special chars

skipDgene logical type. Skip D gene plots?
. save logical type. Save ggplot as Rdata

# Value

None

.abundancePlot Abundance distribution

# Description

Abundance distribution

# Usage

```
.abundancePlot(files, sampleNames, outputDir, skipDgene = FALSE,
    .save = TRUE)
```

#### **Arguments**

files list type. list of files in abundance directory sampleNames vector type. 1-1 correspondance to files

outputDir string type.

skipDgene logical type. Skip D germline abundance plot if TRUE.

. save logical type. Save Rdata ggplot item

# Value

None

# **Description**

Plots alignment quality vs:

- · mismatches
- gaps
- bitscore
- percent identity
- subject start

# Usage

 $. \verb| alignQualityHeatMaps(abundanceDirectory, sampleName)|\\$ 

### **Arguments**

```
abundanceDirectory
```

character type. fully qualified path to abundance directory

sampleName character type. sample name

#### Value

list of ggplotly heatmaps

.allPrimerNames

Collect primer names from FASTA

# Description

Collect primer names from FASTA

# Usage

.allPrimerNames(primerFile)

#### **Arguments**

primerFile string type. Path to primer file

# Value

vector of primer names as seen in primerFile

10 .aminoAcidPlot

.am	ir	۱∩∆	Ci	dΕ	lar

Plots amino acid composition logo

#### **Description**

Plots amino acid composition logo

# Usage

```
.aminoAcidBar(df, scale, region, germ = "")
```

# **Arguments**

df dataframe

scale logical. scale to proportion?
region string. which region is this
germ string. V germline family

#### Value

ggplot2 object

.aminoAcidPlot

Composition logo plot

# Description

Plots 2 kinds: scaled and unscaled composition logos

# Usage

```
.aminoAcidPlot(compositionDirectory, outdir, sampleName,
  regions = c("FR1", "CDR1", "FR2", "CDR2", "FR3", "CDR3", "FR4"),
    .save = TRUE)
```

# **Arguments**

compositionDirectory

string type. string type.

outdir string type. sampleName string type.

regions logical type. vector of FR/CDR regions to plot

. save logical type. save ggplot object

#### Value

none

```
.analyzeUpstreamValidity
```

Plots the validity of upstream sequences

# Description

Plots the distribution of valid, faulty, and missing start codon in IGV germlines (repeated for gene and family levels).

#### Usage

```
.analyzeUpstreamValidity(upstreamDirectories, upstreamOut, expectedLength,
  upstreamLengthRange, sampleNames, combinedNames, mashedNames,
  .save = TRUE)
```

### **Arguments**

upstreamDirectories

list type. List of sample directories

upstreamOut string type. Output directory

expectedLength int type. Expected length of upstream sequences. (i.e. upstream\_end - up-

stream\_start + 1). If this is infinite, no plots will be generated.

 ${\tt upstreamLengthRange}$ 

string type. start\_end format

sampleNames vector type. 1-1 with upstream directories

combinedNames string type. Title friendly "combined" sample names string type. File friendly "mashed-up" sample names

. save logical type. Save Rdata?

#### Value

None

.annotAnalysis  ${\it A}$ 

Annotation analysis

#### **Description**

Annotation analysis

#### **Usage**

```
.annotAnalysis(annotDirectories, annotOut, sampleNames, mashedNames,
    .save = TRUE)
```

#### **Arguments**

annotDirectories

list type. List of sample directories

annotOut string type. Output directory

sampleNames vector type. 1-1 with annotDirectories

mashedNames string type. File output "mashed" sample names

. save logical type. Saves ggplot object

# Value

none

.asRepertoireAlignLen Accessor for alignlen slot

# Description

Accessor for alignlen slot

# Usage

```
.asRepertoireAlignLen(object, collapse = " - ")
```

# Arguments

object AbSeqRep object

collapse character type, collapse the range using this string.

# Value

character type. If collapse is a string, then the ranges are represented as 'start - end' in a string, if collapse is NULL, returns a character vector of length two, denoting the start and end value respectively.

.asRepertoireBitscore 13

.asRepertoireBitscore Accessor for bitscore slot

# Description

Accessor for bitscore slot

### Usage

```
.asRepertoireBitscore(object, collapse = " - ")
```

# **Arguments**

object AbSeqRep object

collapse character type, collapse the range using this string.

#### Value

character type. If collapse is a string, then the ranges are represented as 'start - end' in a string, if collapse is NULL, returns a character vector of length two, denoting the start and end value respectively.

.asRepertoireChain Acce

Accessor for chain slot

# Description

Accessor for chain slot

#### Usage

```
.asRepertoireChain(object)
```

#### **Arguments**

object

AbSeqRep object

# Value

character type, the chain type of this sample

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.asRepertoireDir

Accessor for the outdir slot

# Description

Accessor for the outdir slot

# Usage

```
.asRepertoireDir(object)
```

# **Arguments**

object

AbSeqRep object

#### Value

character type, the output directory of this object

 $. \, as {\tt RepertoireList}$ 

Accessor for AbSeqCRep's list of AbSeqRep objects

# Description

Accessor for AbSeqCRep's list of AbSeqRep objects

# Usage

```
.asRepertoireList(object)
```

# Arguments

object

AbSeqCRep object

# Value

list type, list of AbSeqRep objects that together, compose this AbSeqCRep object.

.asRepertoireName 15

.asRepertoireName

Accessor for the name slot

# Description

Accessor for the name slot

# Usage

.asRepertoireName(object)

# Arguments

object

AbSeqRep object

#### Value

character type, the sample name of this object.

# Description

Accessor for the primer3end slot

# Usage

```
.asRepertoirePrimer3(object)
```

# Arguments

object

AbSeqRep object

# Value

character type, the FASTA file name for primer 3' end sequences

.asRepertoirePrimer5 Accessor for the primer5 end slot

# Description

Accessor for the primer5end slot

# Usage

```
.asRepertoirePrimer5(object)
```

# Arguments

object AbSeqRep object

#### Value

character type, the FASTA file name for primer 5' end sequences

 $. as {\tt RepertoireQueryStart}$ 

 $Accessor for \ {\it qstart} \ slot$ 

# **Description**

Accessor for qstart slot

# Usage

```
.asRepertoireQueryStart(object, collapse = " - ")
```

# **Arguments**

object AbSeqRep object

collapse character type, collapse the range using this string.

# Value

character type. If collapse is a string, then the ranges are represented as 'start - end' in a string, if collapse is NULL, returns a character vector of length two, denoting the start and end value respectively.

.asRepertoireSubjectStart

Accessor for sstart slot

# **Description**

Accessor for sstart slot

# Usage

```
.asRepertoireSubjectStart(object, collapse = " - ")
```

# Arguments

object AbSeqRep object

collapse character type, collapse the range using this string.

#### Value

character type. If collapse is a string, then the ranges are represented as 'start - end' in a string, if collapse is NULL, returns a character vector of length two, denoting the start and end value respectively.

.as RepertoireUpstream  $\ Accessor for the \ {\it upstream } slot$ 

# **Description**

Accessor for the upstream slot

# Usage

.asRepertoireUpstream(object)

# Arguments

object AbSeqRep object

#### Value

character type

18 .calculateDInd

.boxPlot

Creates a box plot

# Description

Creates a box plot

# Usage

```
.boxPlot(dataframes, sampleNames, plotTitle, xlabel = "", ylabel = "",
   subs = "")
```

# **Arguments**

dataframes list type. List of sample dataframes sampleNames vector type. 1-1 with dataframes plotTitle string type xlabel string type ylabel string type subs string type

#### Value

ggplot2 object

 $. \verb|calculateDInd|$ 

Calculates the "standard" diversity indices

# Description

Calculates the "standard" diversity indices

# Usage

```
.calculateDInd(df)
```

# Arguments

```
clonotype dataframe. Vegan format: + + | S.1| S.2| S.3 | S.4 | ... | (each species should have its own column) + + | v1 | v2 | v3 | .... | (each species' count values are placed in the corresponding column) + + | v1 | v2 | v3 | .... |
```

# Value

dataframe with the column headers: shannon , simpson.con , simpson.inv , simpson.gini , renyi.0 , renyi.1 , renyi.2 , renyi.1nf , hill.0 , hill.1 , hill.2 , hill.1nf

renyi.0 => species richness renyi.1 => shannon entropy renyi.2 => inv.gini renyi.Inf => min.entropy finally: hill\_a = exp(renyi\_a)

.calculateDiversityEstimates

Calculates Lower Bound Estimates for unseen species and Common Diversity Indices from clonotype tables

# **Description**

Employ common techniques to calculate LBE for unseen species and commonly used diversity indices

### Usage

```
.calculateDiversityEstimates(diversityDirectories, diversityOut,
    sampleNames)
```

# Arguments

diversityDirectories

list type. List of directories to diversity dir

diversityOut string type. Output directory

sampleNames vector type. 1-1 with diversityDirectories sample names

# Value

None

.canonicalizeTitle

Convert file names to human friendly text

# Description

Convert file names to human friendly text

# Usage

```
.canonicalizeTitle(str)
```

20 .checkVert

# **Arguments**

str

string type

#### Value

string

.capitalize

Helper function to capitalize the first letter of str

# Description

Helper function to capitalize the first letter of str

# Usage

```
.capitalize(str)
```

# **Arguments**

str

string type

#### Value

string, str capitalized

.checkVert

Checks if abseqPy has a metadata line that suggests the orientation

# Description

Checks if abseqPy has a metadata line that suggests the orientation

# Usage

```
.checkVert(filename)
```

# Arguments

filename

csv filename

# Value

True if CSV metadata says "plot vertically"

.cloneDistHist 21

.cloneDistHist	Marginal histogram of clonotypes (blue for shared, grey for total). The y axis is scaled by sqrt (but it doesn't really matter anyway, since we're
	stripping away the y-ticks)

# **Description**

Marginal histogram of clonotypes (blue for shared, grey for total). The y axis is scaled by sqrt (but it doesn't really matter anyway, since we're stripping away the y-ticks)

# Usage

```
.cloneDistHist(df.original, otherClones, lim.min, flip)
```

#### **Arguments**

df.original dataframe with all clones

otherClones clones from the other dataframe

lim.min x-axis minimum limit

flip logical type

# Value

ggplot2 object

 $. \verb|cloneDistMarginal| & \textit{Marginal density graph of clonotypes (blue for shared, grey for total, and the property of the pr$ 

purple for exclusive clones)

# **Description**

Marginal density graph of clonotypes (blue for shared, grey for total, purple for exclusive clones)

#### **Usage**

```
.cloneDistMarginal(df.original, otherClones, lim.min, flip)
```

# **Arguments**

df.original dataframe with all clones

otherClones clones from the other dataframe

lim.min x-axis minimum limit

flip logical type

#### Value

ggplot2 object

22 .collateReports

.clonotypeAnalysis

Comprehensive clonotype analyses

#### **Description**

Comprehensive clonotype analyses

#### Usage

```
.clonotypeAnalysis(diversityDirectories, clonotypeOut, sampleNames,
  mashedNames, .save = TRUE)
```

#### **Arguments**

diversityDirectories

list type. List of directories to diversity dir

clonotypeOut string type. Output directory

sampleNames vector type. 1-1 with diversityDirectories

mashedNames string type. Prefix for ooutput files using "mashed-up"

. save logical type. Save ggplot object?

#### Value

Nothing

.collateReports Collate all HTML reports into a single directory and cretate an entry

index.html file that redirects to all collated HTML files

# **Description**

Collate all HTML reports into a single directory and cretate an entry index.html file that redirects to all collated HTML files

# Usage

```
.collateReports(reports, individualSamples, outputDirectory)
```

# **Arguments**

 $reports \hspace{1cm} list/vector \ type. \ Collection \ of \ strings \ that \ are \ path(s) \ to \ <sample>\_report.html \\ individual Samples$ 

list type. list of AbSeqRep objects. Used to extract filtering information and % read counts.

outputDirectory

string type. Where should the report be placed.

.commonPrimerNames 23

#### Value

Nothing

.commonPrimerNames Collect the intersection of all primer names within a collection of primer files

# **Description**

Collect the intersection of all primer names within a collection of primer files

#### Usage

```
.commonPrimerNames(primerFiles)
```

# **Arguments**

primerFiles list / vector type. Collection of primer files

# Value

vector type. Vector of primerNames that are present in ALL primerFiles. NULL if there's no intersection at all

.correlationTest

Conducts pearson and spearman correlation analysis on dataframe

# **Description**

Conducts pearson and spearman correlation analysis on dataframe

# Usage

```
. {\tt correlationTest}({\tt df})
```

# Arguments

df

dataframe with at least the following 2 columns: + + + | prop.x | prop.y | + + + | ..... | .... | + + where prop.x and prop.y are normalized counts (i.e. frequencies) of the clones They may contain 0 in a column to denote it being missing from sample x or y.

#### Value

named list of pearson, pearson.p, spearman, spearman.p

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.distanceMeasure

Computes the distance between pariwise samples

#### **Description**

Computes the distance between pariwise samples

#### Usage

.distanceMeasure(df)

# **Arguments**

df

dataframe with at least the following 2 columns: +—+ | prop.x | prop.y | +—+ | ..... | .... | +—+ where prop.x and prop.y are normalized counts (i.e. frequencies) of the clones They may contain 0 in a column to denote it being missing from sample x or y.

#### Value

named list of bray.curtis, jaccard, and morisita.horn

.diversityAnalysis

Title Diversity analysis

# Description

Title Diversity analysis

#### **Usage**

```
.diversityAnalysis(diversityDirectories, diversityOut, sampleNames,
  mashedNames, .save = TRUE)
```

#### Arguments

diversityDirectories

list type. List of directories to diversity dir

diversityOut string type. Output directory

sampleNames vector type. 1-1 with diversityDirectories

mashedNames string type. Prefix for output files using "mashed-up" sample names

. save logical type. Save ggplot object?

#### Value

None

.emptyPlot 25

.emptyPlot

Creates and returns an empty plot

# Description

Creates and returns an empty plot

# Usage

```
.emptyPlot()
```

#### Value

empty ggplot2 object

 $. \\ find \\ Repertoires$ 

Given a directory = <abseqPy\_outputdir>/RESULT\_DIR/, returns the directories (repositories) in 'directory'. That is, will not return any sample\_vs\_sample directories. This is done by asserting that a 'repository' must have an (analysis.params) file, and a summary.txt file.

# Description

A sample\_vs\_sample directory will not have these files.

# Usage

```
.findRepertoires(directory)
```

# **Arguments**

directory

string. Path up until <abseqPy\_outputdir>/RESULT\_DIR/

#### Value

vector of strings that are samples in 'directory', note, this is NOT a full path, but just the sample/repertoire name itself

.generateAllSpectratypes

Generates all FR/CDR spectratypes

#### **Description**

Generates all FR/CDR spectratypes

#### Usage

```
.generateAllSpectratypes(diversityDirectories, diversityOut, sampleNames,
  mashedNames, .save = TRUE)
```

#### **Arguments**

diversityDirectories

list type. List of directories to diversity dir

diversityOut string type. Output directory

sampleNames vector type. 1-1 with diversityDirectories

mashedNames string type. Prefix for output files using "mashed-up" sample names

. save logical type. Save ggplot object?

#### Value

Nothing

.generateDelayedReport

Generates report for all samples in 'compare'

# Description

This function is needed because we are delaying the generation of reports until after all threads/processes have joined. There's currently an issue with rmarkdown::render() in parallel execution, see: https://github.com/rstudio/rmarkdown

# Usage

.generateDelayedReport(root, compare, interactivePlot)

# Arguments

root string, project root directory.

compare vector of strings, each string is a comparison defined by the user (assumes that

this value has been checked).

interactivePlot

logical, whether or not to plot interactive plotly plots.

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#### Value

a named list of samples, each an AbSeqRep object found in "root"

.generateReport

Generates HTML report from AbSeqRep and AbSeqCRep ojects

# **Description**

Generates HTML report from AbSeqRep and AbSeqCRep ojects

#### Usage

```
.generateReport(object, root, outputDir, interactivePlot = TRUE,
    .indexHTML = "#")
```

# **Arguments**

object AbSeqCRep type.

root string type. Root directory of the sample(s)

outputDir string type. The path where the HTML will be generated

interactivePlot

logical type. Interactive or not

. indexHTML character type. The back button will redirect to this link. This is typically used

to redirect users back to index.html page

# Value

path (including HTML name) where the report (HTML file) was saved to

.getLineTypes Helper function to return line types by importance based on provided CD/Fs regions

# **Description**

In the aesthetics of diversity plots (rarefaction, recapture, and duplication), the line types should emphasise the most important antibody region, they're ranked in ascending order of: "FR4", "FR1", "FR2", "FR3", "CDR1", "CDR2", "CDR3", "V".

# Usage

```
.getLineTypes(regions)
```

28 .hmFromMatrix

#### **Arguments**

regions a list/vector of strings (regions)

# Value

vector of strings, each corresponding to the appropriate line type for regions.

.getTotal

Get total number of samples (n)

# **Description**

Often enough, the CSV values supplied do not contain raw counts but percentages (so this value will let us know exactly the sample size).

# Usage

```
.getTotal(filename)
```

# **Arguments**

filename csv filename

# Value

string, sample size.

.hmFromMatrix

Plots a plotly heatmap from provided matrix

# **Description**

Plots a plotly heatmap from provided matrix

#### Usage

```
.hmFromMatrix(m, title, xlabel = "", ylabel = "")
```

# Arguments

m matrix type
title character type
xlabel character type
ylabel character type

#### Value

list with keys: static and interactive (ggplot2 object and plotly object respectivelyb)

.inferAnalyzed 29

.inferAnalyzed

Returns all samples found under sampleDirectory

#### **Description**

Returns all samples found under sampleDirectory

#### Usage

```
.inferAnalyzed(sampleDirectory)
```

#### **Arguments**

```
sampleDirectory
```

string, path to sample directory.

#### Value

un-normalized path to all samples under sampleDirectory

.loadMatrixFromDF

Given a dataframe with the columns "from", "to", and value.var, return a symmetric matrix (with diagonal values = diag). I.e. a call to isSymmetric(return\_value\_of\_this\_function) will always be TRUE.

# **Description**

Given a dataframe with the columns "from", "to", and value.var, return a symmetric matrix (with diagonal values = diag). I.e. a call to isSymmetric(return\_value\_of\_this\_function) will always be TRUE.

#### Usage

```
.loadMatrixFromDF(dataframe, value.var, diag, unidirectional = TRUE)
```

# Arguments

dataframe	dataframe with 3 required columns, namely: +
	+   from   to   value.var     +
	parameter
value.var	the column to use as the matrix value
diag	what should the diagonal values be if the dataframe doesn't provide them
unidirectional	logical type. If the dataframe provided has the reverse pairs (i.e. a from-to pair AND a to-from pair with the save values in the value.var column), then this should be FALSE. Otherwise, this function will flip the from-to columns to generate a symmetric dataframe (and hence, a symmetric matrix).

.pairwiseComparison

#### Value

a symmetric matrix with rownames(mat) == colnames(mat) The diagonal values are filled with diag if the dataframe itself doesn't have diagonal data

.loadSamplesFromString

Loads AbSeqCRep or AbSeqRep objects from a list of sampleNames

# Description

Loads AbSeqCRep or AbSeqRep objects from a list of sampleNames

# Usage

.loadSamplesFromString(sampleNames, root, warnMove = TRUE)

# **Arguments**

sampleNames vector, singleton or otherwise root string type. root directory

warnMove logical type. Warning message ("message" level, not "warning" level) if the

directory has been moved?

#### Value

AbSeqRep or AbSeqCRep object depending on sampleNames

.pairwiseComparison Conduct all vs all pairwise comparison analyses

#### **Description**

Conduct all vs all pairwise comparison analyses

# Usage

.pairwiseComparison(dataframes, sampleNames, outputPath, .save = TRUE)

# Arguments

dataframes list of dataframes

sampleNames 1-1 vector corresponding to dataframes

outputPath string . save logical

.plotCirclize 31

# Value

nothing

.plotCirclize

V-J association plot

# **Description**

V-J association plot

# Usage

```
.plotCirclize(sampleName, path, outputdir)
```

# Arguments

sampleName string type

path string type. Path to \_vjassoc.csv

outputdir string type

#### Value

None

.plotDist

Bar plotter

# Description

Plots barplot for all sample in dataframes. If length(sampleNames) == 1, then the bars will also have y-values (or x if horizontal plot) labels on them. Use 'perc' to control if the values are percentages.

# Usage

```
.plotDist(dataframes, sampleNames, plotTitle, vert = TRUE, xlabel = "",
  ylabel = "", perc = TRUE, subs = "", sorted = TRUE,
  cutoff = 15, legendPos = "right")
```

32 .plotDiversityCurves

#### **Arguments**

dataframes list type. List of dataframes

sampleNames vector type. 1-1 correspondence to dataframes.

plotTitle string type.

vert boolean type. True if the plot should be vertical

xlabel string type ylabel string type

perc boolean type. True if data's axis is a percentage proportion (instead of 0-1) only

used if length(sampleNames) == 1

subs string type

sorted boolean type. True if bar plot should be sorted in descending order

cutoff int type. Number of maximum ticks to show (x on vert plots, y on hori plots).

legendPos string type. Where to position legend (see ggplot's theme())

#### Value

ggplot2 object

#### **Description**

Plots rarefaction, recapture, and de-dup plots for specified region

# Usage

```
.plotDiversityCurves(region, diversityDirectories, sampleNames,
    mashedNames, diversityOut, .save = TRUE)
```

#### **Arguments**

region string type. One of: "cdr", "cdr\_v", and "fr". "cdr" means CDR1-3, "cdr\_v"

means CDR3 and V only, and finally "fr" means FR1-4.

diversityDirectories

list type. List of directories to diversity dir

sampleNames vector type. 1-1 with diversityDirectories

mashedNames string type. Prefix for output files using "mashed-up"

diversityOut string type. Output directory sample names

. save logical type. Save ggplot object?

#### Value

Nothing

.plotDuplication 33

ication Duplication level plot	
--------------------------------	--

# **Description**

bins singletons, doubletons, and higher order clonotypes into a line plot

#### Usage

```
.plotDuplication(files, sampleNames, regions = c("CDR3", "V"))
```

#### **Arguments**

files list type. List of strings to \_cdr\_v\_duplication.csv pathname sampleNames vector type. Vector of strings each representing sample names

regions vector type. Which regions to include in the plot. Default = c("CDR3", "V")

#### Value

ggplot2 object

.plotErrorDist	Plots the error distribution for each region: CDRs, FRs, IGV, IGD,
	and IGJ

# **Description**

Plots the distribution of indels (gaps), indels in out-of-frame sequences, and the distribution of mismatches for CDRs, FRs, IGV, IGD, and IGJ.

# Usage

```
.plotErrorDist(productivityDirectories, prodOut, sampleNames,
  combinedNames, mashedNames, .save = TRUE)
```

#### **Arguments**

productivityDirectories

list type. List of directories

prod0ut string type. Output directory

sampleNames vector type. 1-1 with productivity directories

combinedNames string type. Title friendly "combined" sample names string type. File friendly "mashed-up" sample names

. save logical type. Save Rdata?

#### Value

None

.plotIGVErrors

Plots the error distribution for IGV germlines

#### **Description**

Plots the distribution of in-frame unproductive, out-of-frame unproductive, and productive IGV germlines.

#### Usage

```
.plotIGVErrors(productivityDirectories, prodOut, sampleNames,
  combinedNames, mashedNames, .save = TRUE)
```

# **Arguments**

productivityDirectories

list type. List of directories

prod0ut string type. Output directory

sampleNames vector type. 1-1 with productivity directories

combinedNames string type. Title friendly "combined" sample names string type. File friendly "mashed-up" sample names

. save logical type, save Rdata?

#### Value

None

.plotIGVUpstreamLenDist

Plot IGV family distribution for a given upstreamLengthRange

# Description

Given an upstream length range, plot the distributions of IGV family without showing their actual lengths. If their actual lengths matter, refer to .plotIGVUpstreamLenDistDetailed.

#### Usage

```
.plotIGVUpstreamLenDist(upstreamDirectories, upstreamOut,
   upstreamLengthRange, lengthType, sampleNames, combinedNames, mashedNames,
   .save = TRUE)
```

#### **Arguments**

upstreamDirectories

list type. List of sample directories

upstreamOut string type. Output directory

upstreamLengthRange

The range of upstream sequences to be included in this plot. This is usually determined by abseqPy and the format should be as follows: "min\_max", e.g.:

1\_15 means range(1, 15) inclusive.string type.

lengthType string type. "" (the empty string) denotes everything and "\_short" denotes a short

sequence. abseqPy dictates this because it's used for locating the files.

sampleNames vector type. 1-1 with upstream directories

combinedNames string type. Title friendly "combined" sample names string type. File friendly "mashed-up" sample names

. save logical type. Save Rdata?

#### Value

None

.plotIGVUpstreamLenDistDetailed

Plots the detailed length distribution for IGV families

# Description

A boxplot for each IGV families showing the IQR of upstream lengths. In contrast to .plotIGVUpstreamLenDist, which only shows the distribution of IGV families over upstreamLengthRange.

# Usage

```
.plotIGVUpstreamLenDistDetailed(upstreamDirectories, upstreamOut,
   upstreamLengthRange, lengthType, sampleNames, combinedNames, mashedNames,
   .save = TRUE)
```

# Arguments

upstreamDirectories

list type. List of sample directories

upstreamOut string type. Output directory

upstreamLengthRange

The range of upstream sequences to be included in this plot. This is usually determined by abseqPy and the format should be as follows: "min\_max", e.g.: 1\_15 means range(1, 15) inclusive.string type.

36 .plotPrimerIGVStatus

lengthType string type. "" (the empty string) denotes everything and "\_short" denotes a short

sequence. abseqPy dictates this because it's used for locating the files.

sampleNames vector type. 1-1 with upstream directories

combinedNames string type. Title friendly "combined" sample names string type. File friendly "mashed-up" sample names

. save logical type. Save Rdata?

#### Value

None

.plotPrimerIGVStatus Plots, for a given category and pend, the primer IGV indelled dis-

tribution in a bar plot

# **Description**

Plots the abundace of indelled primers relative to IGV germlines

## Usage

```
.plotPrimerIGVStatus(primer, pend, category, primerDirectories,
  sampleNames, primerOut, combinedNames, mashedNames, .save = TRUE)
```

# **Arguments**

primer string, primer name

pend string, either 3 or 5 (primer end)

category string, either "all", "productive", or "outframe"

primerDirectories

string type. Path to primer analysis directory

sampleNames vector type. 1-1 with primerDirectories

primerOut string type. output directory

combinedNames string type. Title friendly "combined" sample names string type. File friendly "mashed-up" sample names

. save logical type. Save Rdata?

#### Value

None

.plotPrimerIntegrity 37

.plotPrimerIntegrity  $Plots\ the\ distribution\ of\ primer\ integrity\ for\ a\ given\ {\it category\ and\ 5'}\ or\ 3'\ {\it pend}$ 

#### **Description**

Plots the distribution of primer integrity for a given category and 5' or 3' pend

## Usage

```
.plotPrimerIntegrity(primerIntegrity, pend, category, primerDirectories,
  sampleNames, primerOut, combinedNames, mashedNames, .save = TRUE)
```

#### **Arguments**

primerIntegrity

string. One of "stopcodon", "integrity", "indelled", "indel\_pos"

pend string, either 3 or 5 (primer end)

category string, either "all", "productive", or "outframe"

primerDirectories

string type. Path to primer analysis directory

sampleNames vector type. 1-1 with primerDirectories

primerOut string type. output directory

combinedNames string type. Title friendly "combined" sample names string type. File friendly "mashed-up" sample names

. save logical type. Save Rdata?

## Value

None

.plotRarefaction Rarefaction plot

## **Description**

Plots the number of unique clonotypes (on the y-axis) drawn from a sample size on the x axis. The number of unique clonotypes is averaged over 5 repeated rounds.

```
.plotRarefaction(files, sampleNames, regions = c("CDR3", "V"))
```

.plotRecapture

## Arguments

files list type. A list of files consisting of path to samples

sampleNames vector type. A vector of strings, each being the name of samples in files

regions vector type. A vector of strings, regions to be included. Defaults to c("CDR3",

"V")

#### Value

ggplot2 object

.plotRecapture

Plots capture-recapture

# Description

Plots the percent of recapture clonotypes (on the y-axis) drawn from a repeated (with replacement) sample size on the x axis. The percentage of recaptured clonotypes is averaged over 5 recapture rounds.

## Usage

```
.plotRecapture(files, sampleNames, regions = c("CDR3", "V"))
```

# Arguments

files list type. List of \_cdr\_v\_recapture.csv.gz files.

sampleNames vector type. A vector of strings each representing the name of samples in files.

regions vector type. A vector of strings, regions to be included in the plot. defaults to

c("CDR3", "V")

#### Value

ggplot2 object

.plotSamples 39

.plotSamples	Monolith AbSeq Plot function - the "driver" program	

# Description

Monolith AbSeq Plot function - the "driver" program

# Usage

```
.plotSamples(sampleNames, directories, analysis, outputDir, primer5Files,
    primer3Files, upstreamRanges, skipDgene = FALSE)
```

## **Arguments**

sampleNames	vector type. Vector of sample names in strings
directories	vector type. Vector of directories in strings, must be 1-1 with sampleNames
analysis	vector / list type. What analysis to plot for. If sampleNames or directories is > 1 (i.e. AbSeqCRep), then make sure that it's an intersection of all analysis conducted by the repertoires, otherwise, it wouldn't make sense
outputDir	string type. Where to dump the output
primer5Files	vector / list type. Collection of strings that the sample used for primer5 analysis. If sample N doesn't have a primer 5 file, leave it as anthing but a valid file path.
primer3Files	vector / list type. Collection of strings that the sample used for primer 3 analysis. If sample N doesn't have a primer 3 file, leave it as anthing but a valid file path.
upstreamRanges	list type. Collection of "None"s or vector denoting upstreamStart and upstreamEnd for each sample.
skipDgene	logical type. Whether or not to skip D gene distribution plot

#### Value

none

|--|--|--|

# Description

Plots length distribution

```
.plotSpectratype(dataframes, sampleNames, region, title = "Spectratype",
   subtitle = "", xlabel = "Length(AA)", ylabel = "Distribution",
   showLabel = FALSE)
```

# **Arguments**

dataframes list type. List of dataframes.

sampleNames vector type. 1-1 correspondance with dataframes

region string type. Region that will be displayed in the plot title. This specifies which

region this spectratype belongs to. If not supplied, a default (start, end) range

will be displayed instead

title string type. Ignored if region is specified.

subtitlestring typexlabelstring typeylabelstring type

showLabel bool type. Show geom\_text? - Ignored if samples > 1

#### Value

ggplot2 object

.plotUpstreamLength

Plot upstream distribution

## **Description**

Plot upstream distribution

#### Usage

```
.plotUpstreamLength(upstreamDirectories, upstreamOut, expectedLength,
  upstreamLengthRange, sampleNames, combinedNames, mashedNames,
  .save = TRUE)
```

#### **Arguments**

upstreamDirectories

list type. List of sample directories

upstreamOut string type. Output directory

expectedLength int type. Expected length of upstream sequences. (i.e. upstream\_end - up-

stream start + 1).

 ${\tt upstreamLengthRange}$ 

string type. start\_end format

sampleNames vector type. 1-1 with upstream directories

combinedNames string type. Title friendly "combined" sample names string type. File friendly "mashed-up" sample names

. save logical type. Save Rdata?

#### Value

None

.plotUpstreamLengthDist

Plot upstream sequence length distribution for upstream sequences (5'UTR or secretion signal) for a given upstreamLengthRange

#### **Description**

Given an upstream length range, plot the distribution of upstream sequence lengths.

## Usage

```
.plotUpstreamLengthDist(upstreamDirectories, upstreamOut,
    upstreamLengthRange, lengthType, sampleNames, combinedNames, mashedNames,
    .save)
```

## **Arguments**

upstreamDirectories

list type. List of sample directories

upstreamOut string type. Output directory

upstreamLengthRange

The range of upstream sequences to be included in this plot. This is usually determined by abseqPy and the format should be as follows: "min\_max", e.g.:

1\_15 means range(1, 15) inclusive.string type.

lengthType string type. "" (the empty string) denotes everything and "\_short" denotes a short

sequence. abseqPy dictates this because it's used for locating the files.

sampleNames vector type. 1-1 with upstream directories

combinedNames string type. Title friendly "combined" sample names

mashedNames string type. File friendly "mashed-up" sample names

. save logical type. Save Rdata?

#### Value

None

.prodDistPlot

.primerAnalysis Cond	lucts primer specificity analysis
----------------------	-----------------------------------

## **Description**

Conducts primer specificity analysis

#### Usage

```
.primerAnalysis(primerDirectories, primer5Files, primer3Files, primerOut,
    sampleNames, combinedNames, mashedNames, .save = TRUE)
```

## **Arguments**

primerDirectories

string type. Path to primer analysis directory

primer5Files vector / list type. 5' end primer files primer3Files vector / list type. 3' end primer files

primerOut string type. output directory

sampleNames vector type. 1-1 with primerDirectories

combinedNames string type. Title friendly "combined" sample names string type. File friendly "mashed-up" sample names

. save logical type. Save Rdata?

#### Value

None

.prodDistPlot	Plots a distribution plot for different productivity analysis files
---------------	---

# Description

A wrapper for plotDist

```
.prodDistPlot(productivityDirectories, sampleNames, title, reg,
  outputFileName, region, .save = TRUE)
```

.productivityAnalysis 43

#### **Arguments**

productivityDirectories

vector type. directories where all productivity csv files lives (usually <sample-

name>/productivity/)

sampleNames vector type. title string type.

reg string type. Regular expression to find the right files for this particular distribu-

tion plot

outputFileName string type. Vector of file names to save in the order of regions

region string type. Most of the dist plots are regional based. use "" if no regions are

involved

. save logical type. Save Rdata?

#### Value

None

.productivityAnalysis Conducts productivty analysis

#### **Description**

Conducts productivty analysis

#### Usage

```
.productivityAnalysis(productivityDirectories, prodOut, sampleNames,
  combinedNames, mashedNames, .save = TRUE)
```

## **Arguments**

productivityDirectories

list type. List of directories

prod0ut string type. Output directory

sampleNames vector type. 1-1 with productivity directories

combinedNames string type. Title friendly "combined" sample names string type. File friendly "mashed-up" sample names

. save logical type. Save Rdata

## Value

None

.readSummary

.productivityPlot

Summary of productivity

#### **Description**

Shows the percentage of 1. productivity, 2. non-functional + reason for being unproductive, i.e. "Stop Codon" or "Out of frame" or "Stop & Out"

## Usage

```
.productivityPlot(dataframes, sampleNames)
```

#### Arguments

dataframes list type. List of sample dataframes sampleNames vector type. 1-1 with dataframes

#### Value

ggplot2 object

.readSummary

Return value specifed by key from AbSeq's summary file

#### **Description**

Return value specifed by key from AbSeq's summary file

## Usage

.readSummary(sampleRoot, key)

#### **Arguments**

sampleRoot

sample's root directory. For example, /path/to/<outputdir>/reports/<sample\_name>.

key

character type. Possible values are (though they might change)

- · RawReads
- · AnnotatedReads
- · FilteredReads
- · ProductiveReads

#### Value

value associated with key from summary file. "NA" (in string) if the field is not available refer to util.R for the key values

regionAnalysis 45

.regionAnalysis

Title Shows varying regions for a given clonotype defined by its CDR3

#### **Description**

Title Shows varying regions for a given clonotype defined by its CDR3

#### Usage

```
.regionAnalysis(path, sampleName, top = 15)
```

## **Arguments**

path

string type. Path to diversity folder where <sampleName>\_clonotype\_diversity\_region\_analysis.csv.gz

is located

sampleName

string type

top

int type. Top N number of clones to analyze

#### Value

ggplot2 object

.reportLBE

Reports abundance-based (Lower bound) diversity estimates using the

Vegan package

# Description

Reports abundance-based (Lower bound) diversity estimates using the Vegan package

#### Usage

```
.reportLBE(df)
```

## **Arguments**

df

#### Value

46 .scatterPlot

.saveAs

Saves ggplot object as a Rdata file.

## **Description**

It's a convinient function that does the check and saves at the same time, for brevity within other areas of the code (to eliminate repeated if checks).

#### Usage

```
.saveAs(.save, filename, plot)
```

## **Arguments**

. save logical type. Whether or not we should save.

filename string.

plot ggplot object.

#### Value

nothing

.scatterPlot

Title Creates a scatter plot

#### **Description**

Title Creates a scatter plot

## Usage

```
.scatterPlot(df1, df2, name1, name2, cloneClass)
```

## **Arguments**

df1 dataframe for sample 1
 df2 dataframe for sample 2
 name1 string type, Sample 1 name
 name2 string type. Sample 2 name

cloneClass string type. What region was used to classify clonotypes - appears in title. For

example, CDR3 or V region

#### Value

ggplot2 object

.scatterPlotComplex 47

.scatterPlotComplex Creates a complex scatter plot

#### **Description**

Creates a complex scatter plot

#### Usage

```
.scatterPlotComplex(df.union, df1, df2, name1, name2, cloneClass)
```

#### **Arguments**

df.union a 'lossless' dataframe created by intersecting sample1 and sample2's dataframes.

It should contain NAs where clones that appear in one sample doesn't appear in

the other. For example:

df1 dataframe for sample 1
 df2 dataframe for sample 2
 name1 string type, Sample 1 name
 name2 string type. Sample 2 name

cloneClass string type. What region was used to classify clonotypes - appears in title. For

example, CDR3 or V region

this plotting techique was shamelessly plagarised from https://github.com/mikessh/vdjtools/blob/master/s.

(VDJTools) with minor modifications

#### Value

```
ggplot2 object
```

```
.secretionSignalAnalysis
```

Secretion signal analysis

# Description

Generates all the required plots for Secretion signal analysis. This includes upstream length distributions and upstream sequence validity.

```
.secretionSignalAnalysis(secDirectories, secOut, sampleNames,
  combinedNames, mashedNames, upstreamRanges, .save = TRUE)
```

#### **Arguments**

secDirectories list type. Secretion signal directories where files are located

secOut string type. Where to dump output sampleNames vector type. 1-1 with secDirectories combinedNames string type. Title friendly string

mashedNames string type. File name friendly string

upstreamRanges list type. Upstream ranges for each sample. If length(secDirectories) > 1, the

plots will only be generated for upstream ranges that are present in ALL sam-

ples. (i.e. the intersection)

. save logical type, save Rdata?

#### Value

none

.substituteStringInFile

Substitutes the first occurance of 'key' with 'value' in 'filename'

## **Description**

Substitutes the first occurance of 'key' with 'value' in 'filename'

## Usage

```
.substituteStringInFile(filename, key, value, fixed = FALSE)
```

## **Arguments**

filename character type
key character type
value character type
fixed logical type

## Value

None

.summarySE 49

.summarySE	Summary of dataframe	

#### **Description**

Gives count, mean, standard deviation, standard error of the mean, and confidence interval (default 95%).

adapted from http://www.cookbook-r.com/Graphs/Plotting\_means\_and\_error\_bars\_(ggplot2)/#Helper functions

#### Usage

```
.summarySE(data = NULL, measurevar, groupvars = NULL, na.rm = FALSE,
  conf.interval = 0.95, .drop = TRUE)
```

#### **Arguments**

data a data frame.

measurevar the name of a column that contains the variable to be summariezed groupvars a vector containing names of columns that contain grouping variables

na.rm a boolean that indicates whether to ignore NA's

conf.interval the percent range of the confidence interval (default is 95%)

.drop logical.

#### Value

dataframe

.topNDist	Title Clonotype table	

# Description

Title Clonotype table

# Usage

```
.topNDist(dataframes, sampleNames, top = 10)
```

#### **Arguments**

dataframes list type. List of dataframes.

sampleNames vector type. vector of strings representing sample names should have one-to-one

correspondence with dataframes

top int type. Top N clonotypes to plot

50 .UTR5Analysis

## Value

None

.UTR5Analysis

5' UTR analysis

# Description

Generates all the required plots for 5' UTR analysis. This includes upstream length distributions and upstream sequence validity.

# Usage

```
.UTR5Analysis(utr5Directories, utr5Out, sampleNames, combinedNames,
  mashedNames, upstreamRanges, .save = TRUE)
```

## **Arguments**

utr5Directories

list type. 5UTR directories where files are located

utr50ut string type. Where to dump output

sampleNames vector type. 1-1 with utr5Directories

combinedNames string type. Title friendly string

mashedNames string type. File name friendly string

upstreamRanges list type. Upstream ranges for each sample. If length(utr5Directories) > 1, the

plots will only be generated for upstream ranges that are present in ALL sam-

ples. (i.e the intersection)

. save logical type, save Rdata?

## Value

none

vennIntersection 51

.vennIntersection	Title Creates Venndiagram for clonotype intersection

#### **Description**

Title Creates Venndiagram for clonotype intersection

## Usage

```
.vennIntersection(dataframes, sampleNames, outFile, top = Inf)
```

## Arguments

dataframes list type. List of sample dataframes. Only accepts 2 - 5 samples. Warning

message will be generated for anything outside of the range

sampleNames vector type. 1-1 with dataframes outFile string type. Filename to be saved as

top int type. Top N cutoff, defaults to ALL clones if not specified

#### Value

Nothing

AbSeqCRep-class S4 class - AbSeqCompositeRepertoire analysis object

## **Description**

AbSeqCRep is a collection of AbSeqRep S4 objects. This S4 class contains multiple samples(repertoires) and it can be "combined" with other samples by using the + operator to create an extended AbSeqCRep object. This value, in turn, can be used as the first argument to report which generates a comparison between all samples included in the + operation.

Users do not manually construct this class, but rather indirectly obtain this class object as a return value from the + operation between two AbSeqRep objects, which are in turn, obtained indirectly from abseqReport and report functions. It is also possible to obtain this class object by + (adding) AbSeqCRep objects.

#### Value

AbSeqCRep

#### Slots

repertoires list of AbSeqRep objects.

52 AbSeqRep-class

#### See Also

AbSeqRep

#### **Examples**

```
# Use example data from abseqR as abseqPy's output, substitute this
# with your own abseqPy output directory
abseqPyOutput <- tempdir()
file.copy(system.file("extdata", "ex", package = "abseqR"), abseqPyOutput, recursive=TRUE)
samples <- abseqReport(file.path(abseqPyOutput, "ex"), report = 0)
# The provided example data has PCR1, PCR2, and PCR3 samples contained within
# pcr12 and pcr13 are instances of AbSeqCRep
pcr12 <- samples[["PCR1"]] + samples[["PCR2"]]
pcr13 <- samples[["PCR1"]] + samples[["PCR3"]]
# all_S is also an instance of AbSeqCRep
all_S <- pcr12 + pcr13</pre>
```

AbSeqRep-class

S4 class - AbSeqRepertoire analysis object

## **Description**

The AbSeqRep object contains all metadata associated with the AbSeq (python backend) run conducted on it. This S4 class represents a single sample(repertoire) and it can be "combined" with other samples by using the + operator to create an AbSeqCRep object. This value, in turn, can be used as the first argument to report which generates a comparison between all samples included in the + operation.

Users do not manually construct this class, but rather indirectly obtain this class object as a return value from the abseqReport and report functions.

#### Value

AbSeqRep

#### Slots

```
f1 character. Path to FASTA/FASTQ file 1.f2 character. Path to FASTA/FASTQ file 2.
```

chain character. Type of chain, possible values:

- hv
- 1v
- kv
- klv

each representing Heavy, Lambda and Kappa respectively.

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task character. Type of analysis conducted, possible values:

- all
- · annotate
- · abundance
- · diversity
- · productivity
- fastqc
- · primer
- 5utr
- rsasimple
- seqlen
- · secretion
- seqlenclass

name character. Name of analysis.

bitscore numeric. Part of filtering criteria: V gene bitscore filter value.

qstart numeric. Part of filtering criteria: V gene query start filter value.

sstart numeric. Part of filtering criteria: V gene subject start filter value.

alignlen numeric. Part of filtering criteria: V gene alignment length filter value.

clonelimit numeric. Number of clones to export into csv file. This is only relevant in -t all or -t
 diversity where clonotypes are exported into <outdir>/<name>/diversity/clonotypes

detailedComposition logical. Plots composition logo by IGHV families if set to true, otherwise, plots logos by FR/CDRs.

log character. Path to log file.

merger character. Merger used to merge paired-end reads.

fmt character. File format of file1 and (if present) file2. Possible values are FASTA or FASTQ.

sites character. Path to restriction sites txt file. This option is only used if -t rsasimple

primer5end ANY. Path to 5' end primer FASTA file.

primer3end ANY. Path to 3' end primer FASTA file.

trim5 numeric. Number of nucleotides to trimd at the 5' end;

trim3 numeric. Number of nucleotides to trimd at the 3' end;

outdir character. Path to output directory

primer5endoffset numeric. Number of nucleotides to offset before aligning 5' end primers in primer5end FASTA file.

threads numeric. Number of threads to run.

upstream character. Index (range) of upstream nucleotides to analyze. This option is only used if -t 5utr or -t secretion.

seqtype character. Sequence type, possible values are either dna or protein.

database character. Path to IgBLAST database.

actualqstart numeric. Query sequence's starting index (indexing starts from 1). This value overrides the inferred query start position by AbSeq.

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fr4cut logical. The end of FR4 is marked as the end of the sequence if set to TRUE, otherwise the end of the sequence is either the end of the read itself, or trimmed to --trim3 <num>.

domainSystem character. Domain system to use in IgBLAST, possible values are either imgt or kabat.

#### See Also

abseqReport returns a list of AbSeqRep objects.

#### **Examples**

```
# this class is not directly constructed by users, but as a return
# value from the abseqReport method.

# Use example data from abseqR as abseqPy's output, substitute this
# with your own abseqPy output directory
abseqPyOutput <- tempdir()
file.copy(system.file("extdata", "ex", package = "abseqR"), abseqPyOutput, recursive=TRUE)
samples <- abseqReport(file.path(abseqPyOutput, "ex"), report = 0)</pre>
```

abseqReport

Visualize all analysis conducted by abseqPy

## Description

Plots all samples in the output directory supplied to abseqPy's --outdir or -o argument. Users can optionally specify which samples in directory should be compared. Doing so generates additional plots for clonotype comparison and the usual plots will also conveniently include these samples using additional aesthetics.

Calling this function with a valid directory will always return a named list of objects; these individual objects can be combined using the + operator to form a new comparison, in which the report function accepts as its first parameter.

#### Usage

```
abseqReport(directory, report, compare, BPPARAM)
```

#### **Arguments**

directory

string type. directory as specified in -o or --outdir in abseqPy. This tells AbSeq where to look for abseqPy's output.

report

(optional) integer type. The possible values are:

- 0 does nothing (returns named list of AbSeqRep objects)
- 1 generates plots for csv files
- 2 generates a report that collates all plots
- 3 generates interactive plots in report (default)

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each higher value also does what the previous values do. For example, report = 2 will return a named list of AbSeqRep objects, plot csv files, and generate a (non-interactive)HTML report that collates all the plots together.

compare (optional) vector of strings. From the samples in found in directory directory,

they can be selected and compared against each other. For example, to compare "sample1" with "sample2" and "sample3" with "sample4", compare should be c("sample1, sample2", "sample3, sample4"). An error will be thrown if the

samples specified in this parameter are not found in directory.

BPPARAM (optional) BiocParallel backend. Configures the parallel implementation. Refer

to BiocParallel for more information. By default, use all available cores.

#### Value

named list. List of AbSeqRep objects. The names of the list elements are taken directly from the repertoire object itself. This return value is consistent with the return value of report.

#### See Also

#### AbSeqRep

report. Analogous function, but takes input from an AbSeqRep or AbSeqCRep object instead.

#### **Examples**

```
# Use example data from abseqR as abseqPy's output, substitute this
# with your own abseqPy output directory
abseqPyOutput <- tempdir()</pre>
file.copy(system.file("extdata", "ex", package = "abseqR"), abseqPyOutput, recursive=TRUE)
### 1. The `report` parameter usage example:
# report = 0; don't plot, don't collate a HTML report, don't show anything interactive
samples <- abseqReport(file.path(abseqPyOutput, "ex"), report = 0)</pre>
# samples is now a named list of AbSeqRep objects
# report = 1; just plot pngs; don't collate a HTML report; nothing interactive
# samples <- abseqReport(file.path(abseqPyOutput, "ex"), report = 1)</pre>
# samples is now a named list of AbSeqRep objects
# report = 2; plot pngs; collate a HTML report; HTML report will NOT be interactive
# samples <- abseqReport(file.path(abseqPyOutput, "ex"), report = 2)</pre>
# samples is now a named list of AbSeqRep objects
# report = 3 (default); plot pngs; collate a HTML report; HTML report will be interactive
# samples <- abseqReport(file.path(abseqPyOutput, "ex"), report = 3)</pre>
# samples is now a named list of AbSeqRep objects
### 2. Using the return value of abseqReport:
# NOTE, often, this is used to load multiple samples from different directories
\# using abseqReport (with report = 0), then the samples are added together
# before calling the report function. This is most useful when the samples
```

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```
# live in different abseqPy output directory.
# Note that the provided example data has PCR1, PCR2, and PCR3
# samples contained within the directory
stopifnot(names(samples) == c("PCR1", "PCR2", "PCR3"))
# as a hypothetical example, say we found something
# interesting in PCR1 and PCR3, and we want to isolate them:
# we want to explicitly compare PCR1 with PCR3
pcr13 <- samples[["PCR1"]] + samples[["PCR3"]]</pre>
# see abseqR::report for more information.
# abseqR::report(pcr13)
                             # uncomment this line to run
### BPPARAM usage:
# 4 core machine, use all cores - use whatever value that suits you
# samples <- abseqReport(file.path(abseqPyOutput, "ex"),</pre>
                         BPPARAM = BiocParallel::MulticoreParam(nproc))
# run sequentially - no multiprocessing
# samples <- abseqReport(file.path(abseqPyOutput, "ex"),</pre>
                         BPPARAM = BiocParallel::SerialParam())
# see https://bioconductor.org/packages/release/bioc/html/BiocParallel.html
# for more information about how to use BPPARAM and BiocParallel in general.
```

report

Plots AbSeqRep or AbSeqCRep object to the specfied directory

## Description

Plots all samples in the object argument and saves the analysis in outputDir. Users can optionally specify which samples in object should be compared. Doing so generates additional plots for clonotype comparison and the usual plots will also conveniently include these samples using additional aesthetics.

This method is analogous to abseqReport. The only difference is that this method accepts AbSeqRep or AbSeqCRep objects as its first parameter, and the outputDir specifies where to store the result.

```
report(object, outputDir, report = 3)
## S4 method for signature 'AbSeqRep'
report(object, outputDir, report = 3)
```

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```
## S4 method for signature 'AbSeqCRep'
report(object, outputDir, report = 3)
```

#### **Arguments**

object AbSeqRep or AbSeqCRep object to plot.

outputDir string type. Directory where analysis will be saved to.

report (optional) integer type. The possible values are:

• 0 - does nothing (returns named list of AbSeqRep objects)

• 1 - generates plots for csv files

• 2 - generates a report that collates all plots

• 3 - generates interactive plots in report (default)

each value also does what the previous values do. For example, report = 2 will return a named list of AbSeqRep objects, plot csv files, and generate a (non-interactive)HTML report that collates all the plots together.

#### Value

named list. List of AbSeqRep objects. The names of the list elements are taken directly from the repertoire object itself. This return value is consistent with the return value of abseqReport.

#### See Also

abseqReport. Analogus function, but takes input from a string that signifies the output directory of abseqPy as the first arugment instead.

AbSeqRep AbSeqCRep

# Examples

```
# Use example data from abseqR as abseqPy's output, substitute this
# with your own abseqPy output directory
abseqPyOutput <- tempdir()
file.copy(system.file("extdata", "ex", package = "abseqR"), abseqPyOutput, recursive=TRUE)
samples <- abseqReport(file.path(abseqPyOutput, "ex"), report = 0)

# The provided example data has PCR1, PCR2, and PCR3 samples contained within
# We can use the + operator to combine samples, thus requesting the
# report function to compare them:
pcr12 <- samples[["PCR1"]] + samples[["PCR2"]]

# generate plots and report for this new comparison
# report(pcr12, "PCR1_vs_PCR2")

# generate plots only
# report(pcr12, "PCR1_vs_PCR2", report = 1)

# generate plots, and a non-interactive report</pre>
```

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```
# report(pcr12, "PCR1_vs_PCR2", report = 2)
# generate plots, and an interactive report
# report(pcr12, "PCR1_vs_PCR2", report = 3) # this is the default
```

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