Package 'skitools'

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```
Title Various mskilab R utilties
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Description R miscellaneous utilities for basic data manipulation, debugging,
      viz, lsf management, and common mskilab tasks.
Imports gplots,
     hwriter,
      plotrix,
      S4Vectors,
      RColorBrewer,
      methods,
      reshape2,
      IRanges,
      tools,
      Biostrings,
      DT,
      GenomeInfoDb
Depends R (>= 3.1.0),
      GenomicRanges,
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```

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5 .igv_host 85 Index **86** .igv_host .igv_host

Description

several settings for igv host

Usage

```
.igv_host(h)
```

Author(s)

Marcin Imielinski

af af

Description

computes prob density of af over a set of n mutation calls using alt allele count, total counts, cn, and provided grid.size

as per landau, carter et al

altc, totc, and cn are of length n, purity is length 1

Usage

```
af(altc, totc, grid.size = 0.01, verbose = F)
```

Author(s)

6 allele_multiplicity

af2 af2

Description

computes 2D probability density af over a set of n mutation counts two samples, output columns correspond to x and rows correspond to y and dimnames correspond to amounts

```
as per landau, carter et al
altc1, totc1, altc2, totc2 are of length n
```

Usage

```
af2(altc1, totc1, altc2, totc2, grid.size = 0.01, verbose = F,
   animate = NA)
```

Author(s)

Marcin Imielinski

allele_multiplicity allele_multiplicity

Description

Given individuals file (with either Purity value or a bsp_participant_id or Tumor_scan_name columns) and maf file annotated with absolut copy number (maf\$cn.tot) and column \$patient for firehose id.

You can provide maf without cn.tot annotations latter case will pull absolute segs using bsp_participant_id in ind and annotate, but may take a few minutes to pull.

Usage

```
allele_multiplicity(ind, maf, abs.seg = NULL, verbose = TRUE)
```

Author(s)

bisort 7

bisort bisort

Description

"bisorts" matrix according to rows and columns (and optionally removes empty rows, ie with no nonzero)

Usage

```
bisort(A, drop = F)
```

Arguments

A matrix to sort

drop logical flag whether to drop empty rows (=FALSE)

Author(s)

Marcin Imielinski

border border

Description

orders rows of a logical / binary matrix treating each row as binary number with digits encoded as TRUE / FALSE values of entries

Usage

```
border(B, na.rm = TRUE)
```

Arguments

B input matrix logical format, or castable to logical

na.rm removes NA

Value

B resorted using "binary" order

8 bsub_cmd

brewer.master

brewer.master

Description

Makes a lot of brewer colors using an "inexhaustible" brewer palette ie will not complain if number of colors requested is too high.

Yes - this technically violates the "grammar of graphics", but meant for quick and dirty use.

Usage

```
brewer.master(n, palette = "Accent")
```

Arguments

n TODO

palette

character specifyign pallette to start with (options are: Blues, BuGn, BuPu, GnBu, Greens Greys, Oranges, OrRd, PuBu, PuBuGn, PuRd, Purples, RdPu, Reds, YlFn, YlFnBu, YlOrBr, YlOrRd, BrBg, PiYG, PRGn, PuOr, RdBu, RdGy, RdYlBu, RdYlGn, Spectral, Accent, Dark2, Paired, Pastel1, Pastel2, Set2, Set3)

Value

length(n) character vector of colors

Author(s)

Marcin Imielinski

bsub_cmd

bsub_cmd

Description

Makes bsub command that wraps shell command "cmd" to send to queue "queue" redirebmccting output / error etc streams to path prefixed by "jname", optional_args: maximum memory requirements "mem", "jlabel" job label b

Usage

```
bsub_cmd(cmd, queue, jname = NULL, jlabel = NULL, jgroup = NULL,
  mem = NULL, group = "cgafolk", cwd = NULL, mc.cores = NULL,
  deadline = F)
```

camerplot 9

Arguments

cmd	length n vector of shell commands, optionally named, one per job
queue	optional length ${\bf n}$ or length 1 character specifying queue to send jobs to (default hour)
jname	optional length n character specifying names of jobs, this will be the root of the output files generated by the job
jlabel	optional length n character specifying labels of jobs, this the string
jgroup	optional length n character specifying job group name
mem	length n or length 1 integer specifying GB of memory to be used by jobs
group	character specifying job group (default cgafolk)
cwd	character specifying which working directory to launch jobs from (default is current working directory of R session)
mc.cores	length n or 1 integer specifying how many cores to assign to each job
deadline	logical flag whether to send jobs to deadline queue

Value

character vector of bsub commands, which can run using system or dumped to a shell script

Author(s)

Marcin Imielinski

|--|

Description

plots the results of CAMERA in limma package

Usage

```
cameraplot(camera.res, gene.sets, voom.res, design, contrast = ncol(design),
  title = "Camera Gene set notch plot", cex.space = 1,
  col.axis = alpha("gray20", 0.8), col.ramp = c("blue", "red"),
  cex.slab = 1, cex.glab = 1, lwd.notch = 1, tick.w = 0.1,
  max.genes = 10, text.shift = 0.5, height.wf = 0.1, min.corr = 0.1,
  min.dist = 10, max.gene.sets = 20, gtext.shift = 0.2)
```

Arguments

camera.res	output of camera from limma
gene.sets	gene set input to camera (named list of indices into the voom.res gene expression matrix) $$
voom.res	output of voom from limma
design	design matrix input to camera
cex.space	label spacing expansion factor (use if labels get too crowded

10 capitalize

col.axis	axis color character
col.ramp	ramp from lowest to highest expression to phenotype correlation (default blue, red)
cex.slab	set label cex
cex.glab	gene label cex
lwd.notch	notch thickness
max.genes	max genes to draw in "leading edge" of gene set
text.shift	amount to shift text from notches (>0, <1)
height.wf	height of the topmost correlation waterfall plot
min.corr	minimal abs(correlation) value for leading edge definition
text.shift	minimal distance between labels

Author(s)

Marcin Imielinski

Description

Capitalize first letter of each character element of vector "string"

Usage

```
capitalize(string, un = FALSE)
```

Arguments

string character vector to capitalize
un logical flag whether to uncapitalize (=FALSE)

Value

character vector of strings with capitalized values

ccf 11

 ccf

Description

computes fuzzy histogram of ccf across a set of n mutation calls using alt allele count, total counts, cn, and purity, and provided grid.size

as per landau, carter et al

altc, totc, and cn are of length n, purity is length

Usage

```
ccf(altc, totc, cn, purity, grid.size = 0.01, verbose = F)
```

Author(s)

Marcin Imielinski

charToDec

charToDec

Description

converts character vector to byte vector in decimal representation

Usage

charToDec(c)

Arguments

С

character vector

Value

length(c) integer vector of byte representation of c

Author(s)

12 chunk

chr2num

Convert from chrXX to numeric format

Description

Convert from chrXX to numeric format

Usage

```
chr2num(x, xy = FALSE)
```

Arguments

x factor, Rle or character vector with chromosome names

xy Flag to convert M to 25, Y to 24 and X to 23. Default FALSE

Value

character vector with xy=FALSE, or numeric vector with xy=TRUE

chron chron

Description

Repeat a command periodically, e.g. every 10 seconds

Usage

```
chron(expr, period = 5)
```

chunk

chunk

Description

takes same input as seq (from, to, by, length.out) and outputs a 2 column matrix of indices corresponding to "chunks"

Usage

```
chunk(from, to = NULL, by = 1, length.out = NULL)
```

Arguments

from integer where to begin sequence

to integer to end sequence by interval to space sequence

length.out number of desired chunks, i.e. nrows of output matrix

clock 13

Value

2 column matrix of indices, each row representing chunk

Author(s)

Marcin Imielinski

clock

clock times expression

Usage

```
clock(expr)
```

Arguments

expr

R code to eval while suppressing all errors

Author(s)

Marcin Imielinski

clone_cluster

clone_cluster

Description

determines "clone membership" using CCF threshold for n variants, given alt read count, total read count, and purity

first fits k component mixture model (k pre-specified) to CCF histogram .. mixture model can also be given as input returns cluster centers and membership probabilities for each mutation

 $mu \ k \ vector \ of means \ sigma \ k \ vector \ of sigma \ p.cluster \ n \ x \ k \ matrix \ of cluster probabilities \ lambda \ k \ vector \ of cluster membership frequencies return \ n \ x \ k \ matrix \ of probabilities that CCF>ccf.thresh for each variant$

altc, totc, and cn are of length n, purity is length 1

Usage

```
clone_cluster(altc, totc, cn, purity, thresh = 0.95, k = 2,
    mix.model = NULL, grid.size = 0.01, verbose = F, nsamp = 10000)
```

Author(s)

14 coloredData

col.slice

col.slice

Description

Hacked operator for subsetting columns of data.frames, DataFrames, data.tables, GRanges using a vector of regexps

df

Usage

```
df %!% ...
```

Arguments

df

data.frame

regex

string to match or number in which case that column is returned (same behavior

for data.table)

Value

slices of data.frame matching regex

Author(s)

Marcin Imielinski

coloredData

coloredData

Description

S4 class for data with colors used by heatmap.plus

simple object with data (e.g. vector or matrix of categorical, real numbers) + a colormap colormap is a (named) vector mapping factor levels / unique values in data into colors, or otherwise assigning a color range to numeric data.

Instantiate coloredData

Usage

```
coloredData(data, colormap, upright = T)
```

Author(s)

dcast2

dcast2

dcast.data.table but allows vector arguments for value.var,

Description

if value.var is a vector then will combine the right hand side column names with each element of value.var in a merged cast table

Usage

```
dcast2(data, formula, ..., value.var = NULL, fun.aggregate = function(x) if (length(x) \le 1) x[1] else paste(x, collapse = ","), sep = "_")
```

Author(s)

Marcin Imielinski

ddd

ddd

Description

shortcut to gr2dt

Usage

ddd(x)

dedup

dedup

Description

relabels duplicates in a character vector with .1, .2, .3 (where "." can be replaced by any user specified suffix)

relabels duplicates in a character vector with .1, .2, .3 (where "." can be replaced by any user specified suffix)

Usage

```
dedup(x, suffix = ".")
dedup(x, suffix = ".")
```

16 dfstring

Arguments

x input vector to dedup

suffix suffix separator to use before adding integer for dups in x

x input vector to dedup

suffix suffix separator to use before adding integer for dups in x

Value

```
length(x) vector of input + suffix separator + integer for dups and no suffix for "originals" length(x) vector of input + suffix separator + integer for dups and no suffix for "originals"
```

Author(s)

Marcin Imielinski

Marcin Imielinski

dev.all.off

dev.all.off

Description

kills all plot devices

Usage

```
dev.all.off()
```

Author(s)

Marcin Imielinski

dfstring

dfstring

Description

"tuple" style chraacter representation of a table, key name1 = value1, name2 = value2 either as a single line or many lines useful for quick eyeballing of tabular data

Usage

```
dfstring(df, oneline = TRUE, binary = FALSE, sep1 = "; ", sep2 = ", ")
```

Arguments

1.0	1
df	data.frame input

oneline logical flag whether to print on one line (=TRUE)

sep1 first level separator (=;) i.e. between rows

sep2 second level separator (=,) i.e. between columns

dirr 17

Value

character vector of string representation

Author(s)

Marcin Imielinski

dirr dirr

Description

a variant of dir that gsubs pattern from normal output of dir to name output vector eg dirr(path, '.txt') will return dir output with .txt removed eg dirr(path, '.txt', '.rds') will return dir output with .txt subbed with .rds

Usage

```
dirr(x, pattern = NULL, rep = "", full = TRUE, ...)
```

Arguments

x character of path to run dir on
pattern character pattern to limit files to and to replace with rep
rep character pattern to replace filenames with
full whether to return full path
... additional arguments to dir

Value

named vector of file paths, named by file names in dir gsub-stripped with pattern

Author(s)

Marcin Imielinski

discordant.pairs Label discordant read pairs

Description

Labels read pairs discordant based on whether they have (1) ++ or – strand orientation (2) "-" strand read start is not greater than dmin or less than dmin ahead of "+" strand read on same chromosome

Usage

```
discordant.pairs(pairs, inter.only = F, dmin = 50, dmax = 500)
```

Note

need to merge with gr.isdisc

18 dmix

dmix
dmix

Description

generates data frame of density points in a provided range for a provided mix of k densities of a singlen family useful for plugging into downstream plotting (eg ggplot 2)

"..." variables depend on density function, arguments should be provided as they would to the corresponding R function (ie with respect to vectorization)

if collapse = TRUE then the density will be summed according to the mixing parameter yielding a single density (ie a fuzzy histogram) summarizing the mixing distribution

Usage

```
dmix(dens = "dnorm", xlim = NULL, n = 500, alpha = NULL, plot = F,
  fill = T, collapse = F, ...)
```

Arguments

dens	character specifying R density function, the possibilities include (with arguments shown alongside the density names) dnorm: mean, sd dbinom: size, prob dmultinom: size, prob dgamma: shape, rate dbeta: shape1, shape2
xlim	length 2 vector specifying plot bounds (=NULL)
n	integer number of points to draw distribution over (=500)
alpha	length(k) numeric vector specifying mixing probability
plot	logical flag specifying whether to draw the plot (=FALSE)
fill	logical flag specifying whether to fill the colored plots (=TRUE)
collapse	collapse logical flag whether to collapse the mixture components into a single mixture)
	additional density specific arguments each with vectorized values of length k, where k is the number of desired mixture componetns, see dens arugment)

Value

if plot == TRUE then ggplot2 object of plot, otherwise data.frame of data points with fields id specifying thee mixture id, x = t

Author(s)

dplot 19

Description

Plots dotplot of grouped data

Usage

```
dplot(y, group, ylab = "", xlab = "", log = F, dotsize = NULL,
  binwidth = 0.02, title = NULL, ylim = NULL, text.size = NULL)
```

Arguments

У	numeric vector of data
group	length(y) vector of catageories
ylab	y axis label (="')
xlab	x axis label (="')
log	logical flag whether to plot y axis in log format (=FALSE)
dotsize	integer dot size to plot with, as function of 0.02 category width plot real estate (= NULL)
binwidth	numeric binwidth of histogram in units of data quantiles (= NULL)
title	character title of plot (=")
ylim	y limits of plot (= NULL)
text.size	text size of legend (= NULL)

Author(s)

Marcin Imielinski

dtgr	Convert data.table to GRanges	

Description

Takes as input a data.table which must have the fields: start, end, strand, seqnames. All of the remaining fields are added as meta data to the GRanges

Usage

```
dtgr(dt)
```

Arguments

dt data.table to convert to GRanges

20 file.dir

Value

```
GRanges object of length = nrow(dt)
```

Examples

```
## Not run: r <- dtgr(data.table(start=1, seqnames="X", end=2, strand='+'))
```

elcycles

elcycles

Description

enumerates all elementary cycles in a graph via igraph library A is either an adjacency matrix or igraph object

Usage

```
elcycles(A)
```

Arguments

Α

adjacency matrix

Value

list with fields: \$cycles = list of vertices in elementary cycles \$cycles.eix = list of edges in elementary cycles, where edges are numbered according to the 1D index of adj matrix A

Author(s)

Marcin Imielinski

file.dir

file.dir

Description

grabs file.dirs from list of paths

Usage

```
file.dir(paths)
```

Arguments

paths

character vector of full paths

Value

character vector of just file.names

file.name 21

Author(s)

Marcin Imielinski

file.name

file.name

Description

parses filenames from character vector of paths

Usage

```
file.name(paths)
```

Arguments

paths

character vector of full paths

Value

character vector of just file names

Author(s)

Marcin Imielinski

fisher.combined

fisher.combined

Description

Computes fisher combined p value for a matrix of p values where the columns correspond to individual (independent) tests rows correspond to hypotheses.

Usage

```
fisher.combined(Ps)
```

Arguments

Ps

n x k matrix of p values from k different independent tests

Value

length n numeric vector of p values

Author(s)

22 fisher.plot

fisher.pairwise

fisher.pairwise

Description

Performs fisher test on cols of matrix / df x vs cols of matrix / df y

returns list with ncol(x) by ncol(y) matrices p and ratio denoting the p value and odds ratio of the result of the fisher test on col i of x and col j of y

If y is not provided, will correlate rows of x with themselves.

Usage

```
fisher.pairwise(x, y = x)
```

Arguments

- x n x k1 data frame of categorical data on k1 variables
- y n x k2 optional data frame of categorical data on k2 variables (= x)

Value

list with field \$p and \$or correspodning to k1 x k2 matrices of p values and odds ratios for each pair of tests

Author(s)

Marcin Imielinski

fisher.plot

Plots fisher contingency table with p value

Description

Plots fisher contingency table with p value

Usage

```
fisher.plot(0)
```

Arguments

0 observed matrix of counts

fready 23

fready

fread with name cleaning

Description

calls fread while cleaning names using provided or default pattern and sub

Usage

```
fready(..., pattern = "\\W+", sub = "_{-}")
```

Arguments

pattern

character (default \W)

sub

character to sub in names (default _)

Value

data.table

fuckr

fuckr

Description

 \dots what you feel when R is getting on your nerves. Toggles options(error =) to enable / disable debugging mode.

toggles options error recover / NULL

Usage

fuckr()

Author(s)

24 gatk_callvariants

func_code func_code

Description

Produces (simple) R code calling function named "func" with args in list "argv", prepending with source() call to directories in the vector "sources" if specified.

NOTE: args in ... can be lists or vectors consisting of numerical values or characters. Lists can have named fields. These will be assigned in a "hard coded" way in the Rcode, so these should be ideally scalars or pretty short vectors / lists.

For code to run properly, the names of "argv" must correspond to argument names of "func", or if the list has unnamed fields then they must be ordered in the order of the function args.

Useful for dumping tmp code files for farming when there are many arguments being passed

Usage

```
func_code(func, sources = c(), ...)
```

Arguments

func function to call sources files to source

... additional arguments to function which should contain numerical or character

vectors or lists of such vectors

Value

character data of source file containing call to function with arguments hard coded

Author(s)

Marcin Imielinski

gatk_callvariants

Description

Outputs variant calling pipeline on a set of whole genome (or whole exome) bams to shell script and prints instructions on how to execute from command line.

Usage

```
gatk_callvariants(bams, outdir = "./", run = gsub("[^\\w+]", "_",
   as.character(Sys.time()), perl = T), queue = "hour", memlimit = 2,
   sep = "/", skipclean = T, chunk = 250000, runtype = "wg",
   dry = FALSE, dcov = 200, bsub = FALSE, vqsr.memlimit = memlimit)
```

gatk_haplotypecaller 25

Arguments

bams bams to run variant calling on outdir out directory to output to

run name to give run (=timestamp) queue LSF queue to run on (='week')

memlimit memory limit in GB

skipclean logical flag whether to skipclean (=TRUE)

chunk integer chunk of variants to run each section on (=250e3)

runtype character run type (='wg')

dry logical flag whether to do dry run (=FALSE)

dcov parameter to GATK (=200)

bsub whether to run on LSF vs local (=FALSE)

Author(s)

Marcin Imielinski

```
gatk_haplotypecaller gatk_haplotypecaller
```

Description

calls haplotype caller on a set of input bams and a given set of targets, outputting to target.dir intervals are given as GRanges

Usage

```
gatk_haplotypecaller(outdir, bams, intervals = NULL,
  dbsnp = Sys.getenv("GATK.DBSNP"), hg = Sys.getenv("GATK.FASTA"),
  genome = "hg19", outroot = "out", stand_call_conf = 30,
  stand_emit_conf = 2, min_mapq = 20, other.args = "", run = F,
  verbose = T, write_bam = F, oncotate = T)
```

Arguments

outdir out directory to output to
bams bams to run variant calling on

intervals GRanges intervals to run haplotype caller on dbSNP path (=Sys.getenv('GATK.DBSNP'))

hg genome fasta location (=Sys.getenv('GATK.FASTA'))

genome name of genome build (='hg19')
outroot prefix to give output files (='out')

stand_call_conf

confidence for calls (=30)

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stand_emit_conf

confience for emission (=30)

min_mapq minimum mapping quality (=20)

run logical flag whether to run immediately or just return character vector of com-

mand (= FALSE)

verbose logical flag (=TRUE)

write_bam logical flag whether to write the bam (=FALSE)

oncotate logical flag whether to oncotate output into MAF files (=TRUE)

other.arg other arguments to give to haplotype caller (=")

Value

character vector of command(s) (only if run = FALSE), otherwise just runs command with system call.

Author(s)

Marcin Imielinski

gatk_oncotate gatk_oncotate

Description

Makes shell script to oncotate variants outputted from GATK UG run in directory "gatk.dir", outputs instructions how to run the script.

Usage

```
gatk_oncotate(gatk.dir, jname = "gatk.oncotate", mem = 3, queue = "week",
...)
```

Arguments

gatk.dir output directory containing GATK UG output

jname job name to run jobs with mem max memory in GB (=3)

queue queue to run on

Author(s)

gc_content 27

gc_content	Get GC content	from reference genome
gc_content	dei de comeni	jiom rejerence genome

Description

Uses BSgenome package to compute gc content for a collection of segments in seg data frame (\$chr, \$start, \$end or \$chr, \$pos1, \$pos2 or \$chr, \$begin, \$end) Returns vector of gc content of length nrow(segs).

Usage

```
gc_content(segs, bs_genome)
```

Arguments

segs	Segment	data	frame 1	to pull	gc from
3053	Segment	autu	manic (io puii	SCHOIL

bs_genome A BSgenome object. Perhaps BSgenome.Hsapiens.UCSC.hg19::Hsapiens

gr.all	gr.all	

Description

Return a GRanges that holds interavals for all of HG19

Usage

```
gr.all(unmap = FALSE, M = FALSE, Y = TRUE)
```

Arguments

unmap	[default F] Optinally add a "unmapped" chr
М	[default F] Include mitochondrial chr
Υ	[default T] Include Y chr

Value

GRanges object with one element per chromosome

28 gr.isdisc

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gr		1	9	r	П	1	n	
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Check if reads are clipped

Description

Returns a logical vector of length of the input GRanges that that classifies a read as clipped or not. The user can specify a cutoff value for how many bases need to be clipped.

Usage

```
gr.isclip(gr, clip.cutoff = 10)
```

Arguments

gr	Granges OR	data.table	that has	cigar	field and	aname field

clip.cutoff Minimum number of bases that are clipped to call the reads as clipped

Value

logical of length of input, denoting whether that read is part of a clipped read pair.

		٠			٠		
gr		7	C	М	1	C	\sim
51	٠	_	J	u	_	J	·

Checks if reads are discordant

Description

Returns a logical vector denoting if a read is discordant. There is only a minimum absolute isize, and any read below this is not considered discordant. This will return logicals based on read pairs

Usage

```
gr.isdisc(gr, isize = 1000, unmap.only = FALSE)
```

Arguments

gr	Granges OR data.table that has isize field and qname field
isize	Minimum insert size required to call dis <cordant. 1000<="" default="" td=""></cordant.>
-	

unmap.only Find only pairs with an unmapped read

Value

logical vector of length of input, denoting each read as discordant or not

gr.mincov 29

or		mincov	
<u>ہ</u>	٠	III T I I C C V	

Return windows with minimal coverage

Description

Takes a set of GRanges and removes any ranges that don't have a minimal coverage value. If you give it a GRangesList, you will get back an unlisted GRanges.

Usage

```
gr.mincov(gr, min.cov = 2, buffer = 0, ignore.strand = TRUE,
    pintersect = FALSE)
```

Arguments

gr GRanges to filter

min.cov Minimum number of overlaps to keep. Default 2

buffer Add a buffer to the ranges when computing overlaps. Default 0

ignore.strand Ignore the strand when comparing overlaps. Default TRUE

pintersect Force the pintersect option for gr.findoverlaps

Value

GRanges

gr.	. peal	ks
-----	--------	----

Find peaks in a GRanges over a given meta-data field

Description

Finds "peaks" in an input GRanges with value field y. first piles up ranges according to field score (default = 1 for each range) then finds peaks. If peel > 0, then recursively peels segments contributing to top peak, and recomputes nextpeak "peel" times if peel>0, bootstrap controls whether to bootstrap peak interval nbootstrap times if id.field is not NULL will peel off with respect to unique (sample) id of segment and not purely according to width if FUN preovided then will complex aggregating function of piled up values in dijoint intervals prior to computing "coverage" (FUN must take in a single argument and return a scalar) if id.field is not NULL, AGG.FUN is a second fun to aggregate values from id.field to output interval

Usage

```
gr.peaks(gr, field = "score", minima = FALSE, peel = 0, id.field = NULL,
bootstrap = TRUE, na.rm = TRUE, pbootstrap = 0.95, nbootstrap = 10000,
FUN = NULL, AGG.FUN = sum, peel.gr = NULL, score.only = FALSE,
verbose = peel > 0)
```

30 gr.readfilter

Arguments

GRanges with some meta-data field to find peaks on gr field character field specifying metadata to find peaks on, default "score, can be NULL in which case the count is computed minima logical flag whether to find minima or maxima id.field character denoting field whose values specifyx individual tracks (e.g. samples) bootstrap logical flag specifying whether to bootstrap "peel off" to statistically determine peak boundaries na.rm remove NA from data pbootstrap quantile of bootstrap boundaries to include in the robust peak boundary estimate (i.e. essentially specifies confidence interval) **FUN** function to apply to compute score for a single individual AGG. FUN function to aggregate scores across individuals

Examples

nboostrap

```
## outputs example gene rich hotspots from example_genes GRanges
pk = gr.peaks(example_genes)

## now add a numeric quantity to example_genes and compute
## peaks with respect to a numeric scores, e.g. "exon_density"
example_genes$exon_density = example_genes$exonCount / width(example_genes)
pk = gr.peaks(example_genes, field = 'exon_density')

## can quickly find out what genes lie in the top peaks by agggregating back with
## original example_genes
pk[1:10] %$% example_genes[, 'name']
```

number of bootstraps to run

gr.readfilter Filter reads by average PHRED score Defines a cutoff score for the mean PHRED quality of a read in a GRanges.

Description

Filter reads by average PHRED score Defines a cutoff score for the mean PHRED quality of a read in a GRanges.

Usage

```
gr.readfilter(gr, cutoff = "+")
```

Arguments

gr GRanges or data.table of reads that has a qname and qual field cutoff cutoff score for mean PHRED quality. Default "+"

gr.reduce 31

Value

GRanges or data.table where reads have mean quality score >= cutoff

gr.reduce

Minimal overlaps for GRanges/GRangesList

Description

Takes any number of GRanges or GRangesList and reduces them to the minimal set of overlapping windows, ignoring strand (optional). Can also collapse only within levels of a meta data field "by"

Usage

```
gr.reduce(..., by = NULL, ignore.strand = TRUE, span = FALSE)
```

Arguments

... GRanges or GRangesList

Details

Will populate output with metadata of first row of input contributing the reduced output range.

Value

GRanges

gr.refactor

gr.refactor

Description

Takes a pile of ranges gr and new seqnames "sn" (either of length 1 or of length(gr)) and returns a gr object with the new seqnames and same widths and new start coordinates. These coordinates are determined by placing each gr on the corresponding chromosome at 1 + gap after previous gr (or at 1)

Usage

```
gr.refactor(gr, sn, gap = 0, rev = FALSE)
```

Arguments

gr	GRanges to refactor
sn	character vector of new seqnames
gap	Default 0
rev	Default FALSE

32 gr.tostring

gr.round

Round a set of GRanges to another set "rounds" a set of query ranges Q to a given interval set S using the following rule: 1) If q in Q is partially / fully within S then return intersection of q with S. 2) If q intersects multiple ranges in S and up = F then return the "first" range, otherwise the last range 3) If q in Q is fully outside of S (ie fully inside not S) then return the start-1 (if up = F) or end+1 (if up = F) of the matching range in not S

Description

Round a set of GRanges to another set "rounds" a set of query ranges Q to a given interval set S using the following rule: 1) If q in Q is partially / fully within S then return intersection of q with S. 2) If q intersects multiple ranges in S and up = F then return the "first" range, otherwise the last range 3) If q in Q is fully outside of S (ie fully inside not S) then return the start-1 (if up = T) or end+1 (if up = F) of the matching range in not S

Usage

```
gr.round(Q, S, up = TRUE, parallel = FALSE)
```

Arguments

Q	Query GRanges (strand is ignored)
S	Subject GRanges (strand is ignored)
up	[default TRUE] See description.
parallel	[default FALSE] If TRUE, assumes Q and S are same length and this analysis is only performed between the corresponding Q and S pairs.

Value

Rounded GRanges

Examples

```
## Not run: query <- GRanges(1, IRanges(c(100,110),width=201), seqinfo=Seqinfo("1", 500)) subject <- GRanges(1, IRanges(c(160,170),width=201), seqinfo=Seqinfo("1", 500)) gr.round(query, subject) ## End(Not run)
```

gr.tostring gr.tostring

Description

dumps out a quick text representation of a gr object (ie a character vector)

gr2gatk 33

Usage

```
gr.tostring(gr, places = 2, interval = 1e+06, unit = "MB",
    prefix = "chr")
```

Arguments

gr GRanges

places Number of decimal places. Default 2

interval Default 1e6
unit Default "MB"
prefix Default "chr"

Value

text representation of input

gr2gatk

Dump GRanges to GATK file

Description

Dumps gr object into gatk intervals in file path "file"

Usage

```
gr2gatk(gr, file, add.chr = F)
```

Arguments

gr GRanges

file file

 ${\it add.chr} \qquad \qquad {\it Flag\ to\ add\ "chr"\ to\ seqnames.\ Default\ FALSE}$

Value

returns 0 if completed

34 grl.span

gr2grl

gr2grl Quick way to make grl from list of indices into a GRanges gr

Description

gr2grl Quick way to make grl from list of indices into a GRanges gr

Usage

```
gr2grl(gr, ix)
```

Arguments

gr GRanges to split ix vector to split on

grl.filter

Filters GRangesList to only include ranges in the specified window

Description

(this is different from

Usage

```
grl.filter(grl, windows)
```

Details

does not return list in necessarily same order

grl.span

grl.span

Description

Returns GRanges object representing the left / right extent of each GRL item. In case of "chimeric" GRL items (ie that map to two chromosomes) there are two options: (1) specify "chr" chromosome as argument to subset GRL's that are on that chromosome, and compute GRL extents from this, any GRL full outside of that chromosome will get a 0 width GRL (2) (default) allow chimeric GRL items to get an extent that is with respect to the first chromosome in that GRL

Usage

```
grl.span(grl, chr = NULL, ir = FALSE, keep.strand = TRUE)
```

grl.split 35

Arguments

grl	GRangesList to query
chr	[Default NULL]
ir	[Default FALSE]
keep.strand	[Default TRUE]

Details

If a grl item contains ranges that lie on different chromosomes, then corresponding grange will have chromosome "NA" and IRange(0,0)

grl.split	grl.split
gi 1.3piit	gii.

Description

splits GRL's with respect to their seqnames and strand (default), returning new grl whose items only contain ranges with a single seqname / strand

Usage

```
grl.split(grl, seqname = TRUE, strand = TRUE, values = c())
```

Arguments

grl	GRangesList to split
seqname	Default TRUE
strand	Default TRUE

values columns of values field in grl

Details

can also split by arbitrary (specified) genomic ranges value fields

```
gsea_leading_edge gsea_leading_edge
```

Description

Draws gsea plot for an input gene set and outputs leading edge

Usage

```
gsea_leading_edge(gene.set, sig, draw.plot = T, cex = 1, asp = 2,
  eps = 1e-16, name = "")
```

Author(s)

36 heatmap.plus

gstring

gstring

Description

quick function to parse gr from character vector IGV / UCSC style strings of format gr1;gr2;gr3 where each gr is of format chr:start-end[+/-]

Usage

```
gstring(...)
```

headf

headf

Description

"head" +/- grep vector of files

Usage

```
headf(x, n = 5, grep = NULL)
```

Arguments

x vector of iles

grep string to grep in files (=NULL)

Author(s)

Marcin Imielinski

heatmap.plus

heatmap.plus

Description

Additional features: allows several label tracks on top, bottom, left, and right, with separate top and bottom legend frames tohouse each allows use of coloredData in tracks

hets 37

Usage

```
heatmap.plus(x, Rowv = NULL, Colv = if (symm) "Rowv" else NULL,
bar = FALSE, show.rdend = TRUE, show.cdend = TRUE, topColAttr = NULL,
bottomColAttr = NULL, leftRowAttr = NULL, rightRowAttr = NULL,
leg.args = NULL, dim.heatmap = c(4, 4), distfun = dist,
hclustfun = hclust, reorderfun = function(d, w) reorder(d, w), add.expr,
symm = FALSE, revC = identical(Colv, "Rowv"), scale = c("row", "column",
"none"), na.rm = TRUE, margins = c(1, 1), cexRow = 0.2 + 1/log10(nr),
cexCol = 0.2 + 1/log10(nc), labRow = NULL, add.grid = F,
col.grid = "gray", lwd.grid = 1, size.legend.panel = 0.4,
size.feature.panel = 0.2, col = topo.colors(100), optimal.leaf = T,
return.clust = F, labCol = NULL, main = NULL, xlab = NULL,
ylab = NULL, keep.dendro = FALSE, las.col = 2,
verbose = getOption("verbose"), ...)
```

hets

hets

Description

generates allele fraction at all possible hets at sites specified by vcf (eg hapmap) input for tumor and normal

Usage

```
hets(tum.bam, norm.bam = NULL, out.file,
  vcf.file = "/cga/meyerson/home/marcin/DB/dbSNP/hapmap_3.3.b37.vcf",
  chunk.size1 = 1000, chunk.size2 = 100, mc.cores = 1, verbose = T,
  na.rm = TRUE, filt.norm = T)
```

html_link

html_link

Description

returns text with html link

Usage

```
html_link(href, text = NULL)
```

Arguments

href character vector of paths to link

text text to display

Value

character vector of html link text

38 igv

Author(s)

Marcin Imielinski

Description

makes a open and close html tag with optional text inside and optional (named) vector of name=value pairs contained inside of opening tag

Usage

```
html_tag(tag, text = NULL, collapse = " ", ...)
```

Arguments

tag character vector of tags (without brackets)

text text to put inside tags

collapse how to collapse tags (=newline)

Value

character vector of html

Author(s)

Marcin Imielinski

igv igv

Description

Controls IGV on localhost (or specified host, separate from where R session is running). Igv application must be running and listening to a specified port. Then if you configure this port via environment variables (IGV_HOST, IGV_PORT) in the current R session then you can apply the following usages

igv(fn) ## sends any given file(s) into igv (eg .bam, .wig, .bed) igv(loci = cool.loci) ## plots the windows specified as GRanges or IGV-parsable strings (eg gene name) igv(gr = cool.gr) ## sends granges object to igv session, Note: currently requires the ability to write to a public_html that is web viewable by computer running IGV igv(snapshot = fn) ## sends current screen to file igv(new = TRUE) ## refreshes current session igv(reset = TRUE) ## resets connections, sometimes useful if IGV not responding

If alternate file paths are present on server (where R is runing) and computer running IGV, then can specify gsub.paths variable which is a list of length 2 vectors specifying how to convert file paths from arguments given to igv to ones that can be loaded locally.

39 igv.loci

Usage

```
igv(paths = NULL, gr = NULL, loci = NULL, snapshot = NULL,
 track.view = NULL, new = FALSE, reset = FALSE, wkspace = "PanLungWGS",
 host = Sys.getenv("IGV_HOST"), mac = !grepl("(^cga)|(node\\d+)", host),
 rawpaths = FALSE, sort.locus = NULL, gsub.paths = list(),
 port = Sys.getenv("IGV_PORT"))
```

Arguments

paths	file paths to display on current igv session (=NULL)
gr	GRanges or GRAngesList of numeric genomic data or interval genomic annotations to send to IGV session, if gr has field \$score then data will be dumped to .bw otherwise to .bed or .gff (=NULL)
loci	GRanges or IGV parsable string specifying what window(s) on genome to view (=NULL)
snapshot	file path to store snapshot in (has to be interpretable on file system where IGV is running)
track.view	command for setting the track display mode ("expand", "squish" or "collapse")
new	logical flag whether to start new IGV session
reset	logical flag whether to reset connection between R and IGV (useful if IGV non responsive)
host	character specifying host where IGV is running
mac	logical flag specifying whether host is a local "mac" (ie then apply gsub.paths) otherwise keep paths the same
gsub.paths	list of length 2 vectors specifying gsub args to apply to filenames when mac = TRUE
port	integer specifying port where IGV is running

Author(s)

Marcin Imielinski

igv.loci	igv.loci Wrapper for igv function to dump table + screenshots for individual GRanges loci that have sample column (default Tumor_Sample_Barcode) that is a key into ind data.table where columns matching col.string are fetched and plotted IGV host and port are
	taken from environment variables

Description

Wrapper for igv function to visualize a given (single) mutation in a maf file

Usage

```
igv.loci(mut, ind, out.path, sample.key = "Tumor_Sample_Barcode",
 sleep = 30, window = 400, host = Sys.getenv("IGV_HOST"),
 port = Sys.getenv("IGV_PORT"), overwrite = FALSE, snapshots = TRUE,
 verbose = TRUE)
```

40 img_link

Author(s)

Marcin Imielinski

img.html img.html

Description

takes img.paths and dumps out html with imgs +/- names

can be dumped into a file for showing many images into a single page alternative to img_link for "embedding images"

Usage

```
img.html(paths, text = names(paths), height = 1024, width = 768,
header = 1)
```

Arguments

paths vector of (relative) paths to embed in html

text optional text label to put above embedded images (default = names(paths))

Value

character vector of img tags

Author(s)

Marcin Imielinski

img_link img_link

Description

Returns vector of html image links to files "file" with text "caption"

if embed = T, then will make img link, and additional arguments may be supplied to image tag (eg height, width)

Usage

```
img_link(file, caption = NULL, embed = F, ...)
```

Arguments

file vector of (relative) image file paths to link to caption character vector of captions to add (= ")

embed logical flag whether to imbed images instead of returning links (=FALSE)

... additional parameters to embed in tag (e.g. height and width)

import.ucsc 41

Value

character vector of links (<a> tags) or image tags (or <embed) to dump into an html document

Author(s)

Marcin Imielinski

import.ucsc

import.ucsc

Description

wrapper around rtracklayer import that (1) handles "as" formats (2) has additional flag chrsub to sub in 'chr' in selection, and then sub it out of the output

Usage

```
import.ucsc(con, selection = NULL, text, chrsub = TRUE, verbose = FALSE,
  as = NULL, ...)
```

ind2sub

ind2sub

Description

MATLAB style ind2sub function in R physical essence. Provides the 2D row / column index for a given 1D query

Usage

```
ind2sub(dim, ind, byrow = F)
```

Arguments

dim dimensions of matrix to query

ind 1D index

byrow whether to calculate indices by row or column (= FALSE)

Value

length(ind) x 2 matrix of row and column index pairs corresponding to input ind in dim "dim" matrix

Author(s)

42 install.packages.github

```
install.packages.bioc install.packages.bioc
```

Description

shortcut to install bioconductor packages

Usage

```
install.packages.bioc(pkg)
```

Arguments

pkg

character vector of package names to install

Author(s)

Marcin Imielinski

```
in stall. packages. github \\ in stall. packages. github
```

Description

shortcut to install github packages

Usage

```
install.packages.github(pkg, username, branch)
```

Arguments

pkg

character vector of package names to install

Author(s)

is.dup 43

is.dup is.dup

Description

labels which vectors in x are part of a dup returns logical TRUE if vector is part of a dup

Note: this is a twist on "duplicated" which only returns TRUE if a given element is a duplicate (i.e. duplicated () is FALSE for the original version for the duplicate, while is.dup() will be TRUE for that element)

x can be vector or matrix

Usage

```
is.dup(x)
```

Arguments

Y

vector or matrix to check

Value

logical vector of length(x) or nrow(x)

Author(s)

Marcin Imielinski

ldim

Description

returns dimensions of all objects contained in list

ldim

Usage

ldim(1)

Arguments

1

list

Author(s)

44 list.expr

levapply	levapply
----------	----------

Description

Applies FUN locally to levels of x and returns vector of length() (eg can do a "local" order within levels)

Usage

```
levapply(x, by, FUN = "order")
```

Arguments

x input vector of data

by length(x) vector of categorical labels

FUN function that takes a length k vector and outputs a length k vector, used for

processing each "level" of by

Value

length(x) vector of outputs, the results of applying FUN to each "by" defined level of x

Author(s)

Marcin Imielinski

	list.expr	list.expr
--	-----------	-----------

Description

Takes a character or numeric vector and makes an expression for re-creating that character in source code

Usage

```
list.expr(x)
```

Arguments

x input vector

Value

character vector of command to create the input vector

Author(s)

loud 45

loud loud

Description

Runs a system command but prints a message with the output

Usage

```
loud(x)
```

maf2vcf

maf2vcf

Description

Dumps maf data frame to VCF file "fn"

Usage

```
maf2vcf(maf, fn)
```

Arguments

maf maf data.frame fn output file

Author(s)

Marcin Imielinski

mafcount

mafcount

Description

Returns base counts for reference and alternative allele for an input tum and norm bam and maf data frame or GRAnges specifying substitutions

Usage

```
mafcount(tum.bam, norm.bam = NULL, maf, chunk.size = 100, verbose = T,
    mc.cores = 1, ...)
```

Details

maf is a single width GRanges describing variants and field 'ref' (or 'Reference_Allele'), 'alt' (or 'Tum_Seq_Allele1') specifying reference and alt allele. maf is assumed to have width 1 and strand is ignored.

46 maf_classify

maflite

maflite

Description

take maf data frame and returns columns corresponding to "maflite" format https://confluence.broadinstitute.org/display/QMafliteformat

Usage

```
maflite(maf)
```

Arguments

maf

maf.data.frame

Value

data.frame in maf.lite format

Author(s)

Marcin Imielinski

maf_classify

 $maf_classify$

Description

Re-classifies oncotated variants

Usage

```
maf_classify(maf)
```

Arguments

maf

maf data.frame

Value

variant categories

Author(s)

maf_coding 47

Description

Scans "Variant_Classification" field in maf and outputs TRUE if variant is in protein coding region (includes synonymous)

Usage

```
maf_coding(maf, inclusive = T)
```

Arguments

maf

maf data.frame

Value

logical vector specifying whether row satisfies the criterion

Author(s)

Marcin Imielinski

maf_disp	maf_disp	
----------	----------	--

Description

Quick display of rows of data.frame holding contents of Oncotated maf file

Usage

```
maf_disp(maf, flavor = NULL, sorted = F, show.pat = TRUE,
  extra_cols = NULL, gene = NULL, pat = NULL)
```

Arguments

maf	data.frame with Oncotated maf columns
flavor	character specifying 'functional' or 'validation' flavors, which correspond to special column slices of maf data.frame (= NULL)
sorted	logical flag whether to return output sorted by gene, variant classification, uniprot site, then patient (=FALSE)
show.pat	logical flag whether to show patient (=TRUE)
gene	character vector of Hugo_Symbol to subset on (=NULL)
pat	character vector of Tumor_Sample_Barcodes to subset on (=NULL)
extra_col	character vector of additional columns to include (=NULL)

48 maf_genic

Value

character vector or sliced data.frame

Author(s)

Marcin Imielinski

maf_exonic

maf_exonic

Description

Scans "Variant_Classification" field in maf and outputs TRUE if variant is exonic

Usage

```
maf_exonic(maf, inclusive = T)
```

Arguments

maf

maf data.frame

Value

logical vector specifying whether maf row satisfies the criterion

Author(s)

Marcin Imielinski

maf_genic

 maf_genic

Description

Scans "Variant_Classification" field in maf and outputs TRUE if variant is genic

Usage

```
maf_genic(maf, inclusive = T)
```

Arguments

maf

maf data.frame

Value

logical vector specifying whether maf row satisfies the criterion

Author(s)

maf_indel 49

maf_indel

 maf_indel

Description

Scans "Variant_Classification" field in maf and outputs TRUE if variant is a indel

Usage

```
maf_indel(maf, inclusive = T)
```

Arguments

maf

maf data.frame

Value

logical vector specifying whether maf row satisfies the criterion

Author(s)

Marcin Imielinski

maf_nonyn

 maf_nonyn

Description

Scans "Variant_Classification" field in maf and outputs TRUE if variant is non-synonymous

Usage

```
maf_nonsyn(maf, inclusive = T)
```

Arguments

maf

maf data.frame

Value

logical vector specifying whether maf row satisfies the criterion

Author(s)

50 maf_sub

 ${\sf maf_ns}$

 maf_ns

Description

Scans "Variant_Classification" field in maf and outputs TRUE if variant is non-synonymous

Usage

```
maf_ns(maf, inclusive = T)
```

Arguments

maf

maf data.frame

Value

logical vector specifying whether maf row satisfies the criterion

Author(s)

Marcin Imielinski

maf_sub

 maf_sub

Description

Scans "Variant_Classification" field in maf and outputs TRUE if variant is a substitution

Usage

```
maf_sub(maf, inclusive = T)
```

Arguments

maf

maf data.frame

Value

logical vector specifying whether maf row satisfies the criterion

Author(s)

maf_syn 51

maf_syn maf_syn

Description

Scans "Variant_Classification" field in maf and outputs TRUE if variant is synonymous

Usage

```
maf_syn(maf, inclusive = T)
```

Arguments

maf

maf data.frame

Value

logical vector specifying whether maf row satisfies the criterion

Author(s)

Marcin Imielinski

maf_to_simple

maf_to_simple

Description

Dumps maf files to "simple" format for input into oncotator, adds dummy ref_allele and tum_allele1 cols if not provided

Usage

```
maf_to_simple(maf, filename, genome)
```

Arguments

maf data.frame filename output file

genome An BSgenome object (was "build genome build (='hg19')")

Author(s)

52 match.bs

maf_truncating

maf_truncating

Description

Scans "Variant_Classification" field in maf and outputs TRUE if variant is truncating

Usage

```
maf_truncating(maf, inclusive = T)
```

Arguments

maf

maf data.frame

Value

logical vector specifying whether maf row satisfies the criterion

Author(s)

Marcin Imielinski

match.bs

Identify matches between query and dictionary

Description

Wrapper around matchPdict to identify matches between a query string query and dictionary dict (both BString objects or subclasses)

Usage

```
match.bs(query, dict, midpoint = FALSE)
```

Arguments

query Query dict Dictionary

midpoint Flag for output the coordinates of the match as the location, where the midpoint

of the dict string matches the given query. Default FALSE

Value

a vector of indices of length width(query) that contains indices of the (starting) dictionary in the query string

mmatch 53

mmatch mmatch

Description

match rows of matrix A to matrix B

Usage

```
mmatch(A, B, dir = 1)
```

Arguments

```
A query matrix k1 x n

B subject matrix k2 x n

dir 1
```

Value

length k1 vector specifying first row of B matching row i of A

Author(s)

Marcin Imielinski

modix modix

Description

```
Takes integer input ix and projects on to 1-based modulus over base l ie modix(1, 5) \rightarrow 1, modix(5, 5) \rightarrow 5, modix(6, 5) \rightarrow 1

Takes integer input ix and projects on to 1-based modulus over base l ie modix(1, 5) \rightarrow 1, modix(5, 5) \rightarrow 5, modix(6, 5) \rightarrow 1
```

Usage

```
modix(ix, 1)
modix(ix, 1)
```

Arguments

ix	input indices to apply module
1	base of ix
ix	input indices to apply module
1	base of ix

54 more

Value

```
((ix-1) \bmod 1) - 1
((ix-1) \bmod 1) - 1
```

Author(s)

Marcin Imielinski

Marcin Imielinski

morder

morder

Description

matrix order wrt columns .. ie ordering rows matrix based on left to right ordering of columns (if MARGIN = 1) OR ordering columns of matrix based on top to bottom ordering of rows (if MARGIN = 2)

Usage

```
morder(A, orient = 1)
```

Arguments

A matrix of values

orient integer orientation, if 1 will do row-wise ordering, otherwise column ordering

(=1)

Value

input matrix with rows and columns ordered

Author(s)

Marcin Imielinski

 $\quad \text{more} \quad$

more

Description

```
"more" +/- grep vector of files
```

Usage

```
more(x, grep = NULL)
```

mtable 55

Arguments

x vector of iles

grep string to grep in files (=NULL)

Author(s)

Marcin Imielinski

mtable

mtable

Description

tabulates unique rows values for matrix / data frame

Usage

```
mtable(mat)
```

Arguments

mat

input matrix

Value

unique rows of mat, with additional column \$count on the left

Author(s)

Marcin Imielinski

muffle

muffle Runs code returning NULL is there is any error

Usage

```
muffle(code, ...)
```

Arguments

code R code to eval while suppressing all errors

... additional tryCatch arguments

Value

output of evaluated R code or NULL if error

Author(s)

56 mutclusters

Description

unlists a list of vectors, matrices, data frames into a n x k matrix whose first column specifies the list item index of the entry and second column specifies the sublist item index of the entry and the remaining columns specifies the value(s) of the vector or matrices.

force.cbind = T will force concatenation via 'cbind' force.rbind = T will force concatenation via 'rbind'

Usage

```
munlist(x, force.rbind = F, force.cbind = F, force.list = F)
```

Arguments

X	list of vectors, matrices, or data frames
force.rbind	logical flag to force concatenation via rbind (=FALSE), otherwise will guess
force.cbind	logical flag to force concatenation via cbind (=FALSE), otherwise will guess
force.list	logical flag to force concatenation via unlist (=FALSE), otherwise will guess

Value

data.frame of concatenated input data with additional fields \$ix and \$iix specifying the list item and within-list index from which the given row originated from

Author(s)

Marcin Imielinski9

Description

Returns genes with a degree of mutaiton clustering (e.g. ranked by how many k>k.thresh clusters with d<d.thresh pairwise distance, or the largest cluster with those characteristics)

if max.cluster = TRUE returns maximum size cluster in gene where either all (method == complete) or at least one (method single) mutation pair is within distance d

if max.cluster = F then returns number of clusters of mutations of count greater than k within a distance d per gene

eg d = 0, k = 1, will give the number of unique sites with more than 1 perfectly recurrent mutation per gene

Clustering is by default using single-linkage agglomerative clustering, but any method that is input to helust can be used

mutpairsd 57

Usage

```
mutclusters(maf, d = 0, k = 1, method = "single", max.cluster = TRUE)
```

Arguments

maf data.frame

d max distance threshold in amino acid space

k minimum number of mutations in returned clusters

method character specifying "single" or "complete" linkage clustering of mutations

max.cluster logical flag whether to return the gene with the largest cluster (if TRUE) or the

most number of clusters (if FALSE) (=TRUE)

Value

genes ranked by numbers of cluster or max.cluster size

Author(s)

Marcin Imielinski

mutpairsd	mutpairsd
-----------	-----------

Description

Takes maf data.frame and outputs table that lists how many pairs there are <= distance d in amino acid space

Usage

```
mutpairsd(maf, d = 0)
```

Arguments

maf data.frame

d distance threshold in amino acid space

Value

clusters of mutations with how many pairs of variants supporting

Author(s)

58 nz

mutrate_window

mutrate_window

Description

Computes mutation rates along k gene "windows" along an ordered list "genes" of genes.

Usage

```
mutrate_window(genes, maf, cov, window = 100)
```

Author(s)

Marcin Imielinski

mut_genecluster

mut_genecluster

Description

Greedy divisive clustering of genes based on mutation rates along a provided order (eg order of gene expression)

Outputs a list of gene clusters (each list a character vector of gene symbols)

Usage

```
mut_genecluster(genes, maf, cov, p.thresh = 0.05, min.cluster.size = 2,
bonferonni = TRUE)
```

Author(s)

Marcin Imielinski

nz

nz

Description

outputs the nonzero entries of a vector or array

Usage

```
nz(x, zero = 0, full = FALSE, matrix = TRUE)
```

Arguments

```
\mathsf{k} length(x)
```

zero integer specifying what to use as the "zero" value in the input (=0)

pad 59

Value

data.frame of row id col id value pairs

Author(s)

Marcin Imielinski

pad pad

Description

pads an (integer) vector with ${\bf k}$ places below and above its lowest and highest value (by default, clips at 0)

useful for querying around specific entires of vector, matrix, data.frame, GRanges ewtc

Usage

```
pad(x, k, clip = T)
```

Arguments

x integer vector to pad

k window around each entry to pad

clip logical flag whether to clip elements below 0 (=TRUE)

Value

"padded" integer vector of unique entires with entries in k window around each input included

Author(s)

Marcin Imielinski

parsesnpeff parsesnpeff

Description

parses vcf file containing SnpEff annotations on Strelka calls

Usage

```
parsesnpeff(vcf, id = NULL)
```

Arguments

vcf path to vcf id id of case

60 pindel

Value

GRanges object of all variants and annotations

Author(s)

Kevin Hadi

phist phist

Description

Quick plotlyhistogram

Usage

```
phist(expr, data = data.frame(), ...)
```

pindel

Author(s)

Marcin Imielinski

pindel

Description

calls pindel on a set of input bams and a given set of targets, outputting to target.dir intervals are given as GRanges

Usage

```
pindel(outdir, bams, intervals = NULL, isizes = NULL,
  hg = Sys.getenv("GATK.FASTA"), genome = "hg19", outroot = "out",
  run = F, verbose = T, write_bam = F, oncotate = F, threads = 1,
  window.size = 10, other.args = "")
```

Arguments

outdir out directory to output to bams vector of input bams

intervals GRanges of to run on (=NULL)

isizes integer insert.size to use

hg genome fasta location (=Sys.getenv('GATK.HG19'))

genome genome build (='hg19')

outroot prefix to give output files (='out')

plop 61

run logical flag whether to run immediately or just return character vector of com-

mand (= FALSE)

verbose logical flag

write_bam logical flag whether to write the bam (=FALSE)

oncotate logical flag whether to oncotate output into MAF files (=TRUE)

threads number of threads to use

window.size integer window size to use (=10)

other.args other args to add (=")

Value

character vector of command(s) (only if run = FALSE), otherwise just runs command with system call.

Author(s)

Marcin Imielinski

plop plop

Description

grabs file and plops into public_html (or Sys.getenv('PLOP.DIR') if defined)

prefix will be added to left of file name (can include firectory)

if fn is list then prefix is expanded to unlisted fn

Useful for inspecting a specific subset of analysis files eg when debugging.

Usage

```
plop(fn, prefix = NULL, force = NULL)
```

Arguments

fn character vector of filenames to "plop" into ~public_html/prefix (Sys.getenv('PLOP.DIR')

is used as alternative if defined)

prefix character prefix to add to filenames after plopping (can include subdirectories

which can be inspected)

Author(s)

62 pmGSEA

```
plot_multiplicity plot_multiplicity
```

Description

Plots allele fractions and colors in an individual based on multiplicity call stratified by total copy number value "individual" is a list or data frame with field \$Individual_Id, \$Purity

maf is a maf file annotated with multiplicity (ie the output of allele_multiplicity) with additional fields \$cn.tot, \$mult, \$mult.p

Usage

```
plot_multiplicity(individual, maf, plot.reads = F)
```

Author(s)

Marcin Imielinski

pmGSEA

poor mans GSEA

Description

```
pmGSEA "poor man's GSEA " ***
```

Given a gene.set (character vector) or gene.sets (list of character vectors) and given a named vector of significance values or table of significant genes (sig.table) (if table then significance column is \$p or first column) identifies gene sets that have significant negative deviation of a "signed K-S" statistic vs uniform distribution (ie have p values significantly clustering towards zero) ie are significantly enriched in genes showing positive selection.

if positive.selection = F, will identify sets with significantly positive deviation of a "signed K-S" statistic (ie have p values significantly clustering towards 1) these are sets showing significant negative selection.

All p-values are computed against a distribution of signed K-S statistic obtained through permutation using random gene sets of the same size chosen from sig.table

Will adaptively perform permutations between minperms and maxperms using following rule of thumb: if there are <PERM.THRESH permutations with greater than (lower.tail = F) or less than (lower.tail = T) score than observed score, then will compute additional perms

*** actually not much poorer than the original GSEA, basically a reimplementation of Mootha et al Nat Gen 2002

Usage

```
pmGSEA(gene.sets, sig.table, min.perms = 100, max.perms = 1e+05,
positive.selection = T, length.filter = F, length.range = c(5, 50),
plot.hist = F, verbose = F, bootstrap = T, rank.test = F)
```

ppdf 63

Arguments

gene.sets	a named list of character vectors, each list item is a gene set, i.e. a character vector of genes
sig.table	named vector of \boldsymbol{p} values from an analysis e.g. mutSig, the names of the genes are
min.perms	minimum number of permutations to do in the adaptive permutation test
max.perms	maximum number of permutations to do in the adaptive permutation test
length.range	length 2 integer vector specifying min and max gene set size to score after intersection with genes in sig.table default: c(5,50)

Author(s)

Marcin Imielinski

Description

sends quick plot to ~public_html/plot.pdf. If PPDF.DIR env variable defined, then will send to that directory (i.e. instead of public_html) Useful for doing quick standard plots to a static location which one views through tabs in Chrome or other web browser.

Usage

```
ppdf(expr, filename = "plot.pdf", height = 10, width = 10, cex = 1,
   title = NULL, byrow = TRUE, dim = NULL, cex.title = 1, ...)
```

Arguments

expr	Plotting expression eg plot(runif(1000), runif(1000))
filename	filename under ~/public_html/ or Sys.getenv('PPDF.DIR') to dump plots to (='plot.pdf')
height	integer pixel height of plot (=1000)
width	integer pixel width of plot (=1000)
cex	expansion factor of plot from "default size" either length 1 scalar or length 2 vector specifying height and width expansion (=1)
title	title to add to plot (=")
dim	length 2 integer vector, if expr contains multiple plot calls then will output to matrix of plots with specified dim (=NULL)
cex.title	character expansion factor to title)
	additional arguments to pdf()

Author(s)

64 qhost

Description

sends quick plot to ~public_html/plot.png. If PPNG.DIR env variable defined, then will send to that directory (i.e. instead of public_html) Useful for doing quick standard plots to a static location which one views through tabs in Chrome or other web browser.

Usage

```
ppng(expr, filename = "plot.png", height = 1000, width = 1000,
  dim = NULL, cex = 1, title = NULL, cex.pointsize = min(cex),
  cex.title = 1, ...)
```

Arguments

expr	Plotting expression eg plot(runif(1000), runif(1000))
filename	filename under ~/public_html/ or Sys.getenv('PPNG.DIR') to dump plots to (='plot.png')
height	integer pixel height of plot (=1000)
width	integer pixel width of plot (=1000)
dim	length 2 integer vector, if expr contains multiple plot calls then will output to matrix of plots with specified dim (=NULL)
cex	expansion factor of plot from "default size" either length 1 scalar or length 2 vector specifying height and width expansion (=1)
title	title to add to plot (="')
cex.title	character expansion factor to title)
	additional arguments to png()

Author(s)

Marcin Imielinski

Description

Tabulates per host cluster load

Usage

```
qhost(full = FALSE, numslots = TRUE)
```

qq_pval 65

qq_pval qq plot given input p values	
--------------------------------------	--

Usage

```
qq_pval(obs, highlight = c(), exp = NULL, lwd = 1, bestfit = T,
  col = NULL, col.bg = "black", pch = 18, cex = 1, conf.lines = T,
  max = NULL, max.x = NULL, max.y = NULL, qvalues = NULL,
  label = NULL, plotly = FALSE, annotations = list(), gradient = list(),
  titleText = "", subsample = NA, ...)
```

Arguments

obs	vector of pvalues to plot, names of obs can be intepreted as labels
highlight	optional arg specifying indices of data points to highlight (ie color red)
lwd	integer, optional, specifying thickness of line fit to data
pch	integer dot type for scatter plot
cex	integer dot size for scatter plot
conf.lines	logical, optional, whether to draw 95 percent confidence interval lines around x-y line
max	numeric, optional, threshold to max the input p values
label	character vector, optional specifying which data points to label (obs vector has to be named, for this to work)
plotly	toggles between creating a pdf (FALSE) or an interactive html widget (TRUE)
annotations	named list of vectors containing information to present as hover text (html widget), must be in same order as obs input
gradient	named list that contains one vector that color codes points based on value, must bein same order as obs input
titleText	title for plotly (html) graph only
samp	integer, optional specifying how many samples to draw from input data (default NULL)

Author(s)

Marcin Imielinski, Eran Hodis, Zoran Gajic

Description

Tabulates cluster usage (qstat()) or if full = TRUE flag given will dump out all running jobs in a data.table

Usage

```
qstat(full = FALSE, numslots = TRUE)
```

66 queues

query_lsf_out	query_lsf_out
quer y_IST_out	query_isj_out

Description

parses "out" and "err" files of jobs with jname root to identify exit status and error codes of jobs

Usage

```
lsf_query(dir = NULL, jname = NULL, detailed = F, mc.cores = 1)
```

Arguments

dir character specifyhing directory where .out and .err files are located

jname character vector names of jobs (as specified in bsub_cmd

detailed logical flag specifying whether to return "detailed" information

mc.cores integer specifying how many cores to use to parse the output data

Value

data.frame of job info

Author(s)

Marcin Imielinski

|--|--|--|

Description

Lists all available queues

Usage

queues()

quickSig 67

quickSig quickSig

Description

Quick implementation of Mike / Gaddy binomial / poisson model. Requires category based coverage output (*.per_gene.coverage.txt file) computed during mutsig preprocess step. (run mutsig_preprocess setting the following additional flags: P.output_per_gene_coverage = true; P.output_per_gene_mutation_c = true; P.simplified_gene_sample_coverage_table = false; P.simplified_gene_sample_mutation_counts_table = false;)

Computes context-category specific mutation rates either across whole cohort or within strata of gene-patient categories.

Outputs a significance table with the following columns for each category (if analyze.categories = TRUE) and/or category "tot" which is across all categories o.k = observed mutations of category k e.k = expected mutations of category k given background model eff.k = log(o.k / e.k) for category k p.k = p value of deviation from expectation under poisson model q.k = q value of deviation

Usage

```
quickSig(maf, cov, patients = NULL, genes = NULL, analyze.categories = F,
  remove.silent = TRUE, limit.cat = NULL, two.tailed = TRUE)
```

Author(s)

Marcin Imielinski

ra.overlaps

identifies events that are in ral that do not exist in ra2. Aside from ral and ra2, all arguments are sent to ra.overlaps

Description

identifies events that are in ra1 that do not exist in ra2. Aside from ra1 and ra2, all arguments are sent to ra.overlaps

Usage

```
ra.setdiff(ra1, ra2, ...)
```

ra.union

ra.union

Description

returns events in ra1 that are in ra2 also

Usage

```
ra.union(ra1, ra2, ...)
```

68 read.delim.cat

Description

takes in either file or data frame from dranger or snowman or path to BND / SV type vcf file and returns junctions in VCF format.

Usage

```
ra_breaks(rafile, keep.features = T, seqlengths = hg_seqlengths(),
  chr.convert = T, snowman = FALSE, swap.header = NULL,
  breakpointer = FALSE, seqlevels = NULL, force.bnd = FALSE, skip = NA,
  get.loose = FALSE)
```

Details

The default output is GRangesList each with a length two GRanges whose strands point AWAY from the break. If get.loose = TRUE (only relevant for VCF)

read.delim.cat read.delim.cat

Description

takes a vector of tab delimited file paths and concatenates them into a single data frame (takin union of identically named / numbered columns as a default)

Usage

```
read.delim.cat(paths, skip = NULL, cols = NULL, include.paths = T,
  include.index = TRUE, cores = NULL, ...)
```

Arguments

paths	length n character vector of paths to tsv files
skip	optional length n or length 1 integer specifying how many lines to skip
cols	optional character vector of which cols to keep (by default union of all columns)
include.paths	optional logical flag whether to include paths to files as column \$source.path column
include.index	optional logical flag whether to include source rownames if exist as \$source.id column
cores	optional integer specifying number of cores to use (def 1)
	additional args to read.delim

Author(s)

readRDA 69

readRDA

readRDA

Description

loads Rdata environment into a list variable and returns (to mirror RDS functionality)

Usage

```
readRDA(fn)
```

Arguments

fn

file name of .rda or .RData file

Value

object containing all the elements of the environment stored in fn

Author(s)

Marcin Imielinski

read_hg

Wrapper around BSgenome call

Description

Retreives either the BSgenome hg18 or hg19 genome by default. Requires packages BSgenome. Hsapiens. UCSC. hg19 for hg19 and BSgenome. Hsapiens. UCSC. hg19 for hg18.

Usage

```
read_hg(hg19 = T, fft = F)
```

Arguments

hg19 Logical whether to return hg18 or hg19 BSgenome. Default TRUE

fft Logical whether to return an ffTrack. Default FALSE

Details

If fft = TRUE, can also also return the hg19 ffTrack (requires that the file exists) Requires the existence of environment variable HG.FFT pointing to ffTrack .rds file..

Value

BSgenome or ffTrack of the genome

70 reorder_maf

relib relib

Description

Reload library

Usage

```
relib(lib = "Flow")
```

reorder_maf

reorder_maf

Description

Re-orders maf to comply with the TCGA MAF specifications (v2.2), tacking on all "non-standard" columns after the initial 32

Usage

```
reorder_maf(maf, include.other = TRUE)
```

Arguments

maf data.frame of MAF

include.other logical flag whether to include non-standard maf columns after the standard ones

(=TRUE)

Value

data.frame representing MAF in standard column order

Author(s)

rmix 71

Description

sample N points from a mixture of k densities of a single functional form (eg norm, beta, multinomial) where n is either an integer vector of length k denoting how many samples to be drawn from each density (in which case N = sum(n)) or n is a scalar, in which case n points are drawn from each density and N = n*k.

p = params data frame whose named columns correspond to arguments to rdens (eg \$n, \$shape1, \$shape2 for rbeta or \$n, \$mean, \$sd for rnorm) rdens = function encoding random number generator for given density, that takes as input named columns of params n = either an nrow(p) integer vector or scalar denoting how many samples to draw from each density

```
n can also be just be a column of p
useful for plotting "smears" of points
```

Output is the rbind-ed output of individual rdens calls

Usage

```
rmix(p, rdens, n = NULL)
```

Arguments

p	k x p data frame of k parameter sets of rdens density functions, each column is a parameter value, each row is a parameter setting for a mixture component
rdens	R density specific random number generator function object (eg rnorm)
n	length k or legnth 1 integer specifying number of samples to draw from each

mixture component

Author(s)

Marcin Imielinski

lice row.slice

Description

Hacked operator for subsetting rows of data.frames, DataFrames, data.tables, GRanges

df

df

Usage

```
df %~% ...
```

72 rrbind2

Arguments

df data.frame

regex string to match or number in which case that column is returned (same behavior

for data.table)

Value

slices of data.frame matching regex

Author(s)

Marcin Imielinski

rpipe

read pipe readsLines from pipe and then closes the pipe

Usage

rpipe(cmd)

rrbind2

Improved rbidn for intersecting columns of data.frames or data.tables

Description

like rbind, but takes the intersecting columns of the dfs rrbind = function(df1, df2, [df3 ... etc],)

Usage

```
rrbind2(..., union = T, as.data.table = FALSE)
```

Arguments

... list of data frames to concatenate

union if union flag is used then will take union of columns (and put NA's for columns

of df1 not in df2 and vice versa). Default TRUE

as.data.table [Default FALSE] return as a data.table

seqinfo2gr 73

seqinfo2gr

Create GRanges from Seqinfo

Description

Creates a genomic ranges from seqinfo object ie a pile of ranges spanning the genome

Usage

```
seqinfo2gr(si, strip.empty = FALSE)
```

Arguments

```
si Seqinfo object
strip.empty Don't know. Default FALSE
```

Examples

```
## Not run: si <- Seqinfo(names(hg_seqlength(), hg_seqlengths()))
seqinfo2gr(si)
## End(Not run)</pre>
```

set.comp

set.comp

Description

Compares two sets and outputs data frame with "left", "middle", "right" members

Usage

```
set.comp(s1, s2)
```

Arguments

```
s1 vector corresponding to "set 1"
s2 vector corresponding to "set 2"
```

Value

list with fields \$left, \$middle, and \$right corresponding to vectors that are in the left setdiff, intersection, right setdiff respectively

Author(s)

Bryan Hernandez

74 sparse_subset

setxor setxor

Usage

```
setxor(A, B)
```

Arguments

A vector specifying set A
B vector specifying set B

Value

elements in A or B that are not in the intersection of A and B

Author(s)

Marcin Imielinski

sortable sortable

Description

dumps sortable list for manual sorting into list.html (in public_html by default)

Usage

```
sortable(x, filename = "list.html", title = NULL)
```

sparse_subset sparse_subset

Description

given $k1 \times n$ matrix A and $k2 \times n$ matrix B returns $k1 \times k2$ matrix C whose entries ij = 1 if the set of nonzero components of row i of A is a (+/- strict) subset of the nonzero components of row j of B

Usage

```
sparse_subset(A, B, strict = FALSE, chunksize = 100, quiet = FALSE)
```

Arguments

A k1 x n matrix
B k2 x n matrix

strict logical flag whether to return strict subset (=FALSE)

chunksize integer size of rows to process from each matrix at a single iteration (=100)

quiet logical flag (=FALSE)

splot 75

Value

 $k1 \times k2$ matrix C whose entries ij = 1 if the set of nonzero components of row i of A is a (+/- strict) subset of the nonzero components of row j of B

Author(s)

Marcin Imielinski

Description

convenient formatted scatter plot with additional features as defaults, useful for fast interactive data inspection / exploration of large datasets (eg 1000s of points): - autoamtic setting of solid dots (pch = 19) - transparent colors for over plotting - automatic setting of x and y limits parametrized by "p.outlier" - quick setting of jiggle / jitter on plot - automatic fitting and plotting of regression line (fit = FALSE)

Usage

```
splot(x, y, cex = 0.4, poutlier = 0.01, col = alpha("black", 0.3),
    xlim = quantile(x, na.rm = T, prob = c(poutlier[1], 1 -
    poutlier[length(poutlier)])), ylim = quantile(y, na.rm = T, prob =
    c(poutlier[1], 1 - poutlier[length(poutlier)])), label = NULL,
    cex.label = 1, adj.label = c(1, 0.5), col.label = "black", log = "",
    jiggle = NULL, fit = FALSE, col.fit = "blue", cex.fit = 1,
    square = FALSE, pch = 19, ...)
```

Arguments

X	numeric vector of x data
У	numeric vector y data
cex	character inspection
poutlier	numeric value between 0 and 1 specifying quantile threshold of outliers to remove (=0.01) $$
col	character vector color (=alpha('black', 0.3))
xlim	length 2 numeric vector specifying x limits (=quantile(x, na.rm = T, prob = $c(poutlier[1], 1-poutlier[length(poutlier)])))$
ylim	<pre>length 2 numeric vector specifying y limits (=quantile(y, na.rm = T, prob = c(poutlier[1], 1-poutlier[length(poutlier)])))</pre>
log	standard plot log string
jiggle	numeric value between 0 and 1 specifying what percentage of plot area to jiggle each point (useful for overplotting) (= NULL)
fit	logical flag whether to fit a linear regression line to the data (=FALSE)
col.fit	character specifying color of linear regression fit (='blue')
cex.fit	character specifying size of text associated with linear regerssion line (=1)
square	logical flag whether to make square plot
pch	pch

76 strsplit2

Author(s)

Marcin Imielinski

standardize_segs

standardize_segs

Description

```
(data frame seg function)
```

Usage

```
standardize_segs(seg, chr = FALSE)
```

Details

Takes and returns segs data frame standardized to a single format (ie \$chr, \$pos1, \$pos2) if chr = TRUE will ensure "chr" prefix is added to chromossome(if does not exist)#'

strsplit.fwf

splits a single string according to fixed widths contained in fw (ie each components i of fw denotes the width of field i in string str

Description

splits a single string according to fixed widths contained in fw (ie each components i of fw denotes the width of field i in string str

Usage

```
strsplit.fwf(str, fw)
```

strsplit2

strsplit2

Description

Strsplit when there are two layers of separators (sep1, sep2) and one needs to extract a collapsed vector of subitem j for all items i.

Takes in a character vector and outputs a list of "separated" items

Usage

```
strsplit2(x, sep1 = ",", sep2 = " ", j = 1)
```

sub2ind 77

Arguments

X	character vector
sep1	character specifying first level separator (=',')
sep2	character specifying second level separator (=' ')
j	integer specifying which subitem to keep (=1)

Value

vector of values for subitem j

Author(s)

Marcin Imielinski

sub2ind sub.	2ind
--------------	------

Description

MATLAB style sub2ind function in R physical essence. Provides the one dim matrix index of row-column locations in matrix

```
(RIP matlab)
```

Usage

```
sub2ind(dim, r, c, byrow = F)
```

Arguments

dim	dimension of matrix to return index for
r	integer vector of row index to look up
С	length(r) integer vector of column index to look up
byrow	whether to calculate indices by row or column (= FALSE)

Value

length(r) vector of 1D indices into matrix with dim "dim"

Author(s)

78 tabstring

system.call

Wrapper to base system function to call system (e.g. perl) from R. The only benefit to this wrapper is a more controlled verbose argument.

Description

Wrapper to base system function to call system (e.g. perl) from R. The only benefit to this wrapper is a more controlled verbose argument.

Usage

```
system.call(syscall, verbose = T)
```

Arguments

syscall string containing the system call

verbose print the syscall to screen, and it's stdout

Author(s)

Jeremiah Wala < jwala@broadinstitute.org>

Examples

```
\# system.call('perl s/[0-9]+//g file1 > file2')
```

tabstring

tabstring

Description

string representation of a named vector (ie the result of tab = table(x) ie name1 (value1), name2 (value2), name3 (value3)

Usage

```
tabstring(tab, sep = ", ", sep2 = "_", dt = FALSE)
```

Arguments

tab "table" or any named(vector)

sep separator to use between table elements

Value

character representation of table

Author(s)

tailf 79

tailf

tailf

Description

```
"tail -f" +/- grep vector of files
```

Usage

```
tailf(x, n = NULL, grep = NULL)
```

Arguments

x vector of iles

grep string to grep in files (=NULL)

Author(s)

Marcin Imielinski

timestamp

timestamp

Description

returns character time stamp

Usage

timestamp()

Author(s)

Marcin Imielinski

toggle_grfo

toggle data.table vs IRanges find overlaps

Description

toggles global setting of whether to use data.table vs IRanges find overlaps machinery

Usage

```
.toggle_grfo()
```

Author(s)

80 vaggregate

ucount

ucount

Description

returns vector of same length as input with number of counts of each value in the whole list

Usage

```
ucount(x)
```

Arguments

Х

vector

Value

length(x) vector with number of instances of each item in x

Author(s)

Marcin Imielinski

vaggregate

vaggregate

Description

same as aggregate except returns named vector with names as first column of output and values as second

Note: there is no need to ever use aggregate or vaggregate, just switch to data.table

Usage

```
vaggregate(...)
```

Arguments

.. arguments to aggregate

Value

named vector indexed by levels of "by"

Author(s)

varcount 81

varcount

Description

Wrapper around applyPileups

Usage

```
varcount(bams, gr, min.mapq = 0, min.baseq = 20, max.depth = 500,
  indel = F, ...)
```

Details

takes in vector of bam paths, GRanges corresponding to sites / territories to query, and outputs a list with fields \$counts = 3D matrix of base counts (A, C, G, T, N) x sites x bams subject to mapq and baseq thresholds #'

```
(uses varbase)
... = other args go to read.bam
```

vplot vplot

Description

Quick violin plot

Usage

```
vplot(y, group = "x", facet1 = NULL, facet2 = NULL, transpose = FALSE,
  mapping = NULL, stat = "ydensity", position = "dodge", trim = TRUE,
  scale = "area", log = FALSE, count = TRUE, xlab = NULL, ylim = NULL,
  ylab = NULL, minsup = NA, scatter = FALSE, text = NULL,
  cex.scatter = 1, col.scatter = NULL, alpha = 0.3, title = NULL,
  legend.ncol = NULL, legend.nrow = NULL, vfilter = TRUE, vplot = TRUE,
  dot = FALSE, stackratio = 1, binwidth = 0.1, plotly = FALSE,
  print = TRUE)
```

Arguments

У	numeric data vector
group	length(y) vector of categories
facet1	optional length(y) vector of row categories to facet on (=NULL)
facet2	optional length(y) vector of column categories to facet on (=NULL)
transpose	logical vector whether flip row / column orientation of facets (=FALSE)
mapping	mapping of groups to colors (=NULL)
scale	scale parameter to geom_vplot (=area)

82 wfplot

log	logical flag whether to log transform (=FALSE)
count	logical flag whether to include counts in ylabels (=TRUE)
xlab	character xlabel (=NULL)
ylab	character ylabel (=NULL)
minsup	minimum support to include in a group (=NA)
scatter	logical flag whether to include scatter of points (=FALSE)
alpha	numeric vector between 0 and 1 to specify alpha transparency of points if scatter is TRUE (0.3)
title	character specifying plot title (=NULL)

Author(s)

Marcin Imielinski

wfplot	Quick waterfall plot	

Description

Quick waterfall plot

data is a numeric vector labels are text labels of the same length as data col is either (1) an unamed list of unique colors (2) a named list mapping unique labels to colors

Usage

```
wfplot(data, labels = NULL, names.arg = NULL, col = NULL, las = 2,
  cex = 1, leg.pos = NULL, ...)
```

Arguments

data		length n numeric vector to be drawn and sorted on y axis
label	S	length n character vector categorical labels of data
names	.arg	length n character vector, optional, of individual labels to be drawn verticallyon x axis
col		optional named character vector mapping unique category labels to colors
las		optional integer vector specifying orientation of labels on barplot
cex		numeric value specifying size of names.arg data labels
leg.p	os	NULL
		additional arguments to barplot

Value

plot

Author(s)

which.char 83

which.char

which.char

Description

finds the index of the character in subject (length 1 character vector) matching nchar = 1 single character query eg which.char('a', 'cat') = 2

if query has more than one char (or has length>1) then will return indices matching <any one> of the characters in any element of query

Usage

```
which.char(subject, query)
```

Arguments

subject length 1 character vector query length 1, nchar 1 character

Value

indices in subject that query appears

Author(s)

Marcin Imielinski

wij

wij

Description

Evaluates output of htmlwidget generating expression (e.g. via highcharter) and send to filename in predefined WIDGET.DIR by default plot.html

Usage

```
wij(expr, filename = "plot.html", zoom = NULL, cex = 1, force = FALSE,
  quiet = FALSE, embed = FALSE)
```

wijj

wijj

Description

Embeds widget in jupyter notebook

Usage

```
wijj(x, width = NULL, height = NULL, file = paste0("plotlyJupyterHTML/",
    digest::digest(Sys.time()), ".html"))
```

84 write.htab

Description

writes data frame (or anything castable to data frame) to pretty HTML formatted table to a static location. Useful for quick table inspection via Chrome or other web browser.

Very similar syntax to write.tab

Usage

```
write.htab(tab, file = NULL, title = NULL, footer = NULL,
highlight = NULL, row.names = TRUE, col.names = TRUE,
high.color = "yellow", row.colors = c("lightgray", "white"),
header.colors = c("#4A4A4A", "white"), data.size = 15, dt = TRUE,
force = FALSE, embed = FALSE, title.size = 15, footer.size = 20,
header.size = round(1.1 * data.size))
```

Arguments

tab	data.frame, data.table, or GRanges
file	optional output .html file (by default ~/public_html/htab.html but directory can be set using env variable HTAB.PATH)
title	title to the page (=NULL)
footer	footer to add to page (=NULL)
highlight	optional integer vector specifying what rows to highlight
row.names	logical flag whether to include row labels (=TRUE)
col.names	logical flag whether to include col labels (=TRUE)
high.color	chaoracter highlight color (= 'yellow')
row.colors	length 2 character vector to shade data rows (= c('lightgray', 'white'))
header.colors	length 2 character vector specifygin background and text for header row (= $c('\#4A4A4A', 'white'))$
data.size	integer font size in px for data, title, and footer (= 15)
title.size	integer font size in px for title (= 15)
footer.size	integer font size in px for footer (= 15)
header.size	integer font size in px for header (= 15)

Author(s)

write.tab 85

write.tab writes tab delimited no quotes without row names table (passes remaining arguments to write.table) equivalent to write.table(sep = TAB.DELIM, quote = F, row.names = F)

Usage

```
write.tab(x, ..., sep = "\t^{"}, quote = F, row.names = F)
```

Arguments

x data.frame to dump

... additional arguments to write.table

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