# Package 'ASCAT'

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ascat.loadData

2 ascat.asmultipcf

	ascat.metrics
	ascat.plotAdjustedAscatProfile
	ascat.plotAscatProfile
	ascat.plotGenotypes
	ascat.plotNonRounded
	ascat.plotRawData
	ascat.plotSegmentedData
	ascat.plotSunrise
	ascat.predictGermlineGenotypes
	ascat.prepareHTS
	ascat.prepareTargetedSeq
	ascat.runAscat
	ascat.synchroniseFiles
Index	21

ascat.asmultipcf

Allele-specific segmentation of multiple samples

#### Description

This segmentation function should only be used if part of the breakpoints are expected to be shared between samples, e.g. due to a common ancestry.

#### Usage

```
ascat.asmultipcf(
  ASCATobj,
  ascat.gg = NULL,
  penalty = 70,
  out.dir = ".",
  wsample = NULL,
  selectAlg = "exact",
  refine = TRUE,
  seed = as.integer(Sys.time())
)
```

### Arguments

ASCATobj	an ASCAT object
ascat.gg	germline genotypes (NULL if germline data is available)
penalty	penalty of introducing an additional ASPCF breakpoint (expert parameter, don't adapt unless you know what you are doing)
out.dir	directory in which output files will be written. Can be set to NA to not write PCFed files.
wsample	Vector of length length(ASCATobj\$samples). Can be used to assign different weights to samples, for example to account for differences in sequencing quality. (Default = NULL)
selectAlg	Set to "exact" to run the exact algorithm, or "fast" to run the heuristic algorithm. (Default = "exact")

ascat.aspcf 3

refine Logical. Should breakpoints be refined on a per sample base? Otherwise each

breakpoint is assumed to be present in each sample. (Default = TRUE)

seed A seed to be set when subsampling SNPs for X in males (optional, default=as.integer(Sys.time())).

#### **Details**

This function saves the results in in [sample].LogR.PCFed.txt and [sample].BAF.PCFed.txt

#### Value

output: ascat data structure containing:

- 1. Tumor\_LogR data matrix
- 2. Tumor\_BAF data matrix
- 3. Tumor\_LogR\_segmented: matrix of LogR segmented values
- 4. Tumor\_BAF\_segmented: list of BAF segmented values; each element in the list is a matrix containing the segmented values for one sample (only for probes that are germline homozygous)
- 5. Germline\_LogR data matrix
- 6. Germline\_BAF data matrix
- 7. SNPpos: position of all SNPs
- 8. ch: a list containing vectors with the indices for each chromosome (e.g. Tumor\_LogR[ch[[13]],] will output the Tumor\_LogR data of chromosome 13
- 9. chr: a list containing vectors with the indices for each distinct part that can be segmented separately (e.g. chromosome arm, stretch of DNA between gaps in the array design)

ascat.aspcf

ascat.aspcf

#### **Description**

run ASPCF segmentation

#### Usage

```
ascat.aspcf(
  ASCATobj,
  selectsamples = 1:length(ASCATobj$samples),
  ascat.gg = NULL,
  penalty = 70,
  out.dir = ".",
  out.prefix = "",
  seed = as.integer(Sys.time())
)
```

#### **Arguments**

```
ASCATobj an ASCAT object
selectsamples a vector containing the sample number(s) to PCF. Default = all
ascat.gg germline genotypes (NULL if germline data is available)
penalty penalty of introducing an additional ASPCF breakpoint (expert parameter, don't adapt unless you know what you're doing)
```

4 ascat.correctLogR

out.dir directory in which output files will be written. Can be set to NA to not write

PCFed files.

out.prefix prefix for output file names

 $seed \qquad \qquad A \ seed \ to \ be \ set \ when \ subsampling \ SNPs \ for \ X \ in \ males \ (optional, \ default=as.integer(Sys.time())).$ 

#### **Details**

This function can be easily parallelised by controlling the selectsamples parameter it saves the results in LogR\_PCFed[sample]\_[segment].txt and BAF\_PCFed[sample]\_[segment].txt

#### Value

output: ascat data structure containing:

- 1. Tumor\_LogR data matrix
- 2. Tumor BAF data matrix
- 3. Tumor\_LogR\_segmented: matrix of LogR segmented values
- 4. Tumor\_BAF\_segmented: list of BAF segmented values; each element in the list is a matrix containing the segmented values for one sample (only for probes that are not germline homozygous)
- 5. Germline\_LogR data matrix
- 6. Germline\_BAF data matrix
- 7. SNPpos: position of all SNPs
- 8. ch: a list containing vectors with the indices for each chromosome (e.g. Tumor\_LogR[ch[[13]],] will output the Tumor\_LogR data of chromosome 13
- 9. chr: a list containing vectors with the indices for each distinct part that can be segmented separately (e.g. chromosome arm, stretch of DNA between gaps in the array design)

ascat.correctLogR ascat.correctLogR

#### **Description**

Corrects logR of the tumour sample(s) with genomic GC content (replication timing is optional)

#### Usage

```
ascat.correctLogR(ASCATobj, GCcontentfile = NULL, replictimingfile = NULL)
```

#### **Arguments**

ASCATobj an ASCAT object

GCcontentfile File containing the GC content around every SNP for increasing window sizes replictimingfile

File containing replication timing at every SNP for various cell lines (optional)

#### **Details**

Note that probes not present in the GC content file will be lost from the results

#### Value

ASCAT object with corrected tumour logR

ascat.GCcorrect 5

ascat.GCcorrect ascat.GCcorrect

#### **Description**

Function kept for backward compatibility, please use ascat.correctLogR instead

#### Usage

```
ascat.GCcorrect(ASCATobj, GCcontentfile = NULL)
```

#### **Arguments**

ASCATobj an ASCAT object

GCcontentfile File containing the GC content around every SNP for increasing window sizes

ascat.getAlleleCounts Obtain allele counts for a given set of loci through external program alleleCounter.

#### **Description**

Obtain allele counts for a given set of loci through external program alleleCounter.

#### Usage

```
ascat.getAlleleCounts(
  seq.file,
  output.file,
  loci.file,
  min.base.qual = 20,
  min.map.qual = 35,
  allelecounter.exe = "alleleCounter",
  ref.fasta = NA
)
```

#### **Arguments**

seq.file A BAM/CRAM alignment file on which the counter should be run.

output.file The file where output should go.

loci.file A file with SNP loci.

min.base.qual The minimum base quality required for it to be counted (optional, default=20).

min.map.qual The minimum mapping quality required for it to be counted (optional, default=35).

allelecounter.exe

A pointer to where the alleleCounter executable can be found (optional, default

points to \$PATH).

ref. fasta A FASTA file for CRAM processing (optional).

#### Author(s)

sd11, tl

ascat.getBAFsAndLogRs Obtain BAF and LogR from the allele counts.

#### **Description**

Obtain BAF and LogR from the allele counts.

#### Usage

```
ascat.getBAFsAndLogRs(
  samplename,
  tumourAlleleCountsFile.prefix,
  normalAlleleCountsFile.prefix,
  tumourLogR_file,
  tumourBAF_file,
 normalLogR_file,
  normalBAF_file,
 alleles.prefix,
 gender,
  genomeVersion,
  chrom_names = c(1:22, "X"),
 minCounts = 20,
 BED_file = NA,
 probloci_file = NA,
  seed = as.integer(Sys.time())
)
```

#### Arguments

```
samplename
                  String, name of the sample.
tumourAlleleCountsFile.prefix
                  Prefix of the allele counts files for the tumour (e.g. "Tumour_alleleFrequencies_chr").
normalAlleleCountsFile.prefix
                  Prefix of the allele counts files for the normal (e.g. "Normal_alleleFrequencies_chr").
tumourLogR_file
                  File where LogR from the tumour will be written.
tumourBAF_file File where BAF from the tumour will be written.
normalLogR_file
                  File where LogR from the normal will be written.
normalBAF_file File where BAF from the normal will be written.
alleles.prefix Prefix path to the allele data (e.g. "G1000_alleles_chr")
                  Gender information, either 'XX' (=female) or 'XY' (=male).
gender
                  Genome version, either 'hg19' or 'hg38'.
genomeVersion
                  A vector with allowed chromosome names (optional, default=c(1:22,'X')). Do
chrom_names
                  not set it to paste0('chr',c(1:22,'X')) if data is 'chr'-based.
                  Minimum depth, in normal samples, required for a SNP to be considered (op-
minCounts
                  tional, default=20).
```

ascat.loadData 7

BED\_file A BED file for only looking at SNPs within specific intervals (optional, de-

fault=NA).

probloci\_file A file (chromosome <tab> position; no header) containing specific loci to ignore

(optional, default=NA).

seed A seed to be set when randomising the alleles (optional, default=as.integer(Sys.time())).

#### Author(s)

dw9, sd11, tl

ascat.loadData

ascat.loadData

#### **Description**

Function to read in SNP array data

#### Usage

```
ascat.loadData(
   Tumor_LogR_file,
   Tumor_BAF_file,
   Germline_LogR_file = NULL,
   Germline_BAF_file = NULL,
   chrs = c(1:22, "X", "Y"),
   gender = NULL,
   sexchromosomes = c("X", "Y"),
   genomeVersion = NULL,
   isTargetedSeq = F
)
```

#### Arguments

```
Tumor_LogR_file
```

file containing logR of tumour sample(s)

Tumor\_BAF\_file file containing BAF of tumour sample(s)

Germline\_LogR\_file

file containing logR of germline sample(s), NULL

Germline\_BAF\_file

file containing BAF of germline sample(s), NULL

chrs a vector containing the names for the chromosomes (e.g. c(1:22,"X"))

gender a vector of gender for each cases ("XX" or "XY"). Default = all female ("XX")

sexchromosomes a vector containing the names for the sex chromosomes. Default = c("X","Y") genomeVersion a string (either 'hg19' or 'hg38') so nonPAR coordinates on X can be stored,

NULL

isTargetedSeq a boolean indicating whether data come from a targeted sequencing experiment.

Default = F

8 ascat.metrics

#### **Details**

germline data files can be NULL - in that case these are not read in

#### Value

ascat data structure containing:

- 1. Tumor LogR data matrix
- 2. Tumor\_BAF data matrix
- 3. Tumor\_LogR\_segmented: placeholder, NULL
- 4. Tumor\_BAF\_segmented: placeholder, NULL
- 5. Germline\_LogR data matrix
- 6. Germline\_BAF data matrix
- 7. SNPpos: position of all SNPs
- 8. ch: a list containing vectors with the indices for each chromosome (e.g. Tumor\_LogR[ch[[13]],] will output the Tumor\_LogR data of chromosome 13
- 9. chr: a list containing vectors with the indices for each distinct part that can be segmented separately (e.g. chromosome arm, stretch of DNA between gaps in the array design)
- 10. chrs: a vector containing chromosome names
- 11. samples: a vector containing sample name(s)
- 12. gender: a vector of gender for each cases ("XX" or "XY"). Default = NULL: all female ("XX")
- 13. sexchromosomes: a vector containing names of sex chromosomes
- 14. X\_nonPAR: a vector of two values (start and stop) to define where the nonPAR region is on X
- 15. isTargetedSeq: boolean indicating whether data come from a targeted sequencing experiment
- 16. failedarrays: placeholder, NULL

ascat.metrics

Function to extract different metrics from ASCAT profiles.

#### **Description**

Function to extract different metrics from ASCAT profiles.

#### Usage

```
ascat.metrics(ASCAT_input_object, ASCAT_output_object)
```

#### **Arguments**

ASCAT\_input\_object

R object generated by the ascat.aspcf function and given to the ascat.runAscat function.

ASCAT\_output\_object

R object generated by the ascat.runAscat function.

#### Value

A dataframe (one sample per line) with the following metrics (as columns):

sex - Sex information as provided.

tumour\_mapd - Median Absolute Pairwise Difference (MAPD) in tumour logR track.

normal\_mapd - Median Absolute Pairwise Difference (MAPD) in normal logR track (should be NA

```
without matched normals and 0 for sequencing data).
GC correction before - logR/GC correlation before correction.
GC_correction_after - logR/GC correlation after correction.
RT_correction_before - logR/RT correlation before correction.
RT_correction_after - logR/RT correlation after correction.
n_het_SNP - Number of heterozygous SNPs.
n_segs_logR - Number of segments in the logR track.
n_segs_BAF - Number of segments in the BAF track.
n_segs_logRBAF_diff - Difference between number of segments in the logR versus BAF track.
frac_homo - Fraction of homozygous (<0.1 | >0.9) probes in tumour.
purity - Purity estimate.
ploidy - Ploidy estimate.
goodness of fit - Goodness of fit.
size intermediate segments - Total size of (unrounded) segments in the X.45-X.55 range.
size_odd_segments - Total size of segments with an odd (1/3/5/+) CN (either nMajor or nMinor).
n_segs - Number of copy-number segments.
segs size - Total size of all segments.
n_segs_1kSNP - Number of segments per 1k heterozygous SNPs.
homdel_segs - Number of segments with homozygous deletion.
homdel_largest - largest segment with homozygous deletion.
homdel_size - Total size of segments with homozygous deletion.
homdel_fraction - Fraction of the genome with homozygous deletion.
LOH - Fraction of the genome with LOH (ignoring sex chromosomes).
mode_minA - Mode of the minor allele (ignoring sex chromosomes).
mode_majA - Mode of the major allele (ignoring sex chromosomes).
WGD - Whole genome doubling event (ignoring sex chromosomes).
GI - Genomic instability score (ignoring sex chromosomes).
```

#### Author(s)

tl

```
{\tt ascat.plotAdjustedAscatProfile}
```

ascat.plotAdjustedAscatProfile

#### Description

Function plotting the "adjusted" (with realistic chromosome sizes) rounded/unrounded ASCAT profiles over all chromosomes.

#### Usage

```
ascat.plotAdjustedAscatProfile(
   ASCAT_output_object,
   REF,
   y_limit = 5,
   plot_unrounded = F,
   png_prefix = ""
)
```

10 ascat.plotAscatProfile

#### **Arguments**

ASCAT\_output\_object

R object generated by the ascat.runAscat function.

REF Can be either "hg19" or "hg38" for standard human genome or a data.frame with

three columns: chrom, start and end.

y\_limit Optional parameter determining the size of the y axis in the profile (default=5). plot\_unrounded Optional parameter to define whether rounded (default) or unrounded profile (set

to TRUE) should be plotted.

png\_prefix Optional parameter to add a prefix to png name (can be also used to set a path).

#### Value

Plot showing the adjusted (rounded/unrounded) ASCAT profile of the sample

```
ascat.plotAscatProfile
```

ascat.plotAscatProfile

#### **Description**

Function plotting the rounded ASCAT profiles over all chromosomes

#### Usage

```
ascat.plotAscatProfile(
  n1all,
  n2all,
  heteroprobes,
  ploidy,
  rho,
  goodnessOfFit,
  nonaberrant,
  y_limit = 5,
  ch,
  lrr,
  bafsegmented,
  chrs
)
```

#### **Arguments**

n1all copy number major allele n2all copy number minor allele

heteroprobes probes with heterozygous germline

ploidy ploidy of the sample
rho purity of the sample
goodnessOfFit estimated goodness of fit

nonaberrant boolean flag denoting non-aberrated samples

ascat.plotGenotypes 11

y\_limit Optional parameter determining the size of the y axis in the nonrounded plot and

ASCAT profile. Default=5

ch a list containing c vectors, where c is the number of chromosomes and every

vector contains all probe numbers per chromosome

1rr (unsegmented) log R, in genomic sequence (all probes), with probe IDs

bafsegmented B Allele Frequency, segmented, in genomic sequence (only probes heterozygous

in germline), with probe IDs

chrs a vector containing the names for the chromosomes (e.g. c(1:22,"X"))

#### Value

plot showing the ASCAT profile of the sample

ascat.plotGenotypes

#### **Description**

ascat.plotGenotypes

#### Usage

```
ascat.plotGenotypes(ASCATobj, title, Tumor_BAF_noNA, Hom, ch_noNA)
```

#### Arguments

ASCATobj an ASCAT object title main title of the plot

Tumor\_BAF\_noNA B-allele frequencies of the tumour sample with removed NA values

Hom Boolean vector denoting homozygous SNPs

ch\_noNA vector of probes per chromosome (NA values excluded)

#### Value

plot showing classified BAF per sample, with unused SNPs in green, germline homozygous SNPs in blue and all others in red

```
ascat.plotNonRounded \quad \textit{ascat.plotNonRounded}
```

#### Description

Function plotting the unrounded ASCAT copy number over all chromosomes

#### Usage

```
ascat.plotNonRounded(
  ploidy,
  rho,
  goodnessOfFit,
  nonaberrant,
  nAfull,
  nBfull,
  y_limit = 5,
  bafsegmented,
  ch,
  lrr,
  chrs
)
```

#### **Arguments**

ploidy	ploidy of the sample
rho	purity of the sample
goodnessOfFit	estimated goodness of fit
nonaberrant	boolean flag denoting non-aberrated samples
nAfull	copy number major allele
nBfull	copy number minor allele
y_limit	Optional parameter determining the size of the y axis in the nonrounded plot and ASCAT profile. Default=5
bafsegmented	B Allele Frequency, segmented, in genomic sequence (only probes heterozygous in germline), with probe IDs
ch	a list containing c vectors, where c is the number of chromosomes and every vector contains all probe numbers per chromosome
lrr	(unsegmented) log R, in genomic sequence (all probes), with probe IDs
chrs	a vector containing the names for the chromosomes (e.g. c(1:22,"X"))

#### Value

plot showing the nonrounded copy number profile, using base plotting function

ascat.plotRawData 13

ascat.plotRawData as

ascat.plotRawData

#### **Description**

Plots SNP array data

#### Usage

```
ascat.plotRawData(
  ASCATobj,
  img.dir = ".",
  img.prefix = "",
  logr.y_values = c(-2, 2)
)
```

#### **Arguments**

```
ASCATobj an ASCAT object (e.g. data structure from ascat.loadData)
```

img.dir directory in which figures will be written

img.prefix prefix for figure names

logr.y\_values define Y min and max values for logR track (optional; default: c(-2,2))

#### Value

Produces png files showing the logR and BAF values for tumour and germline samples

```
ascat.plot Segmented Data\\
```

ascat.plotSegmentedData

#### **Description**

plots the SNP array data before and after segmentation

#### Usage

```
ascat.plotSegmentedData(
  ASCATobj,
  img.dir = ".",
  img.prefix = "",
  logr.y_values = c(-2, 2)
)
```

#### Arguments

```
ASCATobj an ASCAT object (e.g. from ascat.aspcf) img.dir directory in which figures will be written
```

img.prefix prefix for figure names

logr.y\_values define Y min and max values for logR track (optional; default: c(-2,2))

#### Value

png files showing raw and segmented tumour logR and BAF

#### Description

ascat.plotSunrise

#### Usage

```
ascat.plotSunrise(d, psi_opt1, rho_opt1, minim = T)
```

#### **Arguments**

d distance matrix for a range of ploidy and tumour percentage values

psi\_opt1 optimal ploidy rho\_opt1 optimal purity

minim when set to true, optimal regions in the sunrise plot are depicted in blue; if set

to false, colours are inverted and red corresponds to optimal values (default:

TRUE)

#### Value

plot visualising range of ploidy and tumour percentage values

```
as cat. predict Germline Genotypes \\ as cat. predict Germline Genotypes
```

#### **Description**

predicts the germline genotypes of samples for which no matched germline sample is available

#### Usage

```
ascat.predictGermlineGenotypes(
   ASCATobj,
   platform = "AffySNP6",
   img.dir = ".",
   img.prefix = ""
)
```

#### Arguments

ASCATobj an ASCAT object platform used array platform

img.dir directory in which figures will be written

img.prefix prefix for figure names

ascat.prepareHTS 15

#### **Details**

Currently possible values for platform:

AffySNP6 (default)

Custom10k

IlluminaASA

IlluminaGSAv3

Illumina109k

IlluminaCytoSNP

IlluminaCytoSNP850k

Illumina610k

Illumina660k

Illumina700k

Illumina1M

Illumina2.5M

IlluminaOmni5

Affy10k

Affy100k

Affy250k\_sty

Affy250k\_nsp

AffyOncoScan

AffyCytoScanHD

HumanCNV370quad

HumanCore12

HumanCoreExome24

HumanOmniExpress12

Illumina Omni Express Exome

#### Value

predicted germline genotypes

ascat.prepareHTS

Extract both logR and BAF values from sequencing data

#### **Description**

Method derived from the Battenberg package (https://github.com/Wedge-lab/battenberg).

#### Usage

```
ascat.prepareHTS(
tumourseqfile,
normalseqfile,
tumourname,
normalname,
allelecounter_exe,
alleles.prefix,
loci.prefix,
gender,
genomeVersion,
```

16 ascat.prepareHTS

```
nthreads = 1,
      tumourLogR_file = NA,
      tumourBAF_file = NA,
      normalLogR_file = NA,
      normalBAF_file = NA,
      minCounts = 10,
      BED_file = NA,
      probloci_file = NA,
      chrom_names = c(1:22, "X"),
      min_base_qual = 20,
      min_map_qual = 35,
      ref.fasta = NA,
      skip_allele_counting_tumour = F,
      skip_allele_counting_normal = F
    )
Arguments
    tumourseqfile
                     Full path to the tumour BAM/CRAM file.
    normalseqfile
                     Full path to the normal BAM/CRAM file.
    tumourname
                      Identifier to be used for tumour output files.
                      Identifier to be used for normal output files.
    normalname
    allelecounter_exe
                      Path to the allele counter executable.
    alleles.prefix Prefix path to the allele data (e.g. "G1000_alleles_chr").
    loci.prefix
                      Prefix path to the loci data (e.g. "G1000_loci_chr").
                      Gender information, either 'XX' (=female) or 'XY' (=male).
    gender
    genomeVersion
                      Genome version, either 'hg19' or 'hg38'.
                      The number of parallel processes for getting allele counts (optional, default=1).
    nthreads
    tumourLogR_file
                      Path to the tumour logR output (optional, paste0(tumourname, "_tumourLogR.txt")).
    tumourBAF_file Path to the tumour BAF output (optional, paste0(tumourname," tumourBAF.txt")).
    normalLogR_file
                      Path to the normal logR output (optional, paste0(tumourname, "_normalLogR.txt")).
    normalBAF_file Path to the normal BAF output (optional, paste0(tumourname, "_normalBAF.txt")).
                      Minimum depth required in the normal for a SNP to be considered (optional,
    minCounts
                      default=10).
    BED_file
                      A BED file for only looking at SNPs within specific intervals (optional, de-
                      fault=NA).
                      A file (chromosome <tab> position; no header) containing specific loci to ignore
    probloci_file
                      (optional, default=NA).
```

A vector containing the names of chromosomes to be considered (optional, de-

Minimum base quality required for a read to be counted (optional, default=20). Minimum mapping quality required for a read to be counted (optional, de-

FASTA file used for generating CRAMs (optional, default=NA).

chrom\_names

min\_base\_qual

min\_map\_qual

ref.fasta

fault=c(1:22,'X')).

fault=35).

```
skip_allele_counting_tumour
```

Flag, set to TRUE if tumour allele counting is already complete (files are expected in the working directory on disk; optional, default=FALSE).

```
skip_allele_counting_normal
```

Flag, set to TRUE if normal allele counting is already complete (files are expected in the working directory on disk; optional, default=FALSE).

#### Author(s)

sd11, tl

```
ascat.prepareTargetedSeq
```

Method to extract a curated list of SNPs covered by a targeted sequencing experiment.

#### **Description**

From a complete set of loci (alleles.prefix), this method will keep SNPs falling into the targeted design (based on BED\_file) and check allele counts in normal samples (listed in Worksheet). The cleaned list of loci/allele files will be located under Workdir/alleleData/Cleaned/.

#### Usage

```
ascat.prepareTargetedSeq(
 Worksheet,
 Workdir,
 alleles.prefix,
 BED_file,
 allelecounter_exe,
 genomeVersion,
 nthreads = 1,
 minCounts = 10,
  is_chr_based = F,
  chrom_names = c(1:22, "X"),
 min_base_qual = 20,
 min_map_qual = 35,
 ref.fasta = NA,
 plotQC = T
)
```

#### **Arguments**

Worksheet A table with the following columns: Patient\_ID, Normal\_ID, Normal\_file and

Gender. Must contain one single normal per patient. Normal\_file can either be BAMs/CRAMs or paths to pre-computed (zipped) alleleCounts (e.g. "sample\_alleleCounts\_chr"). Gender must either be XX (females) or XY (males).

Workdir The folder where output should go (will be created if it doesn't exist).

alleles.prefix Prefix path to the allele data (e.g. "G1000\_alleles\_chr").

BED\_file A BED file for only looking at SNPs within specific intervals. Must fit with the

design used for targeted sequencing.

18 ascat.runAscat

```
allelecounter_exe
                  Path to the allele counter executable.
                  Genome version, either 'hg19' or 'hg38'.
genomeVersion
nthreads
                  The number of parallel processes to speed up the process (optional, default=1).
minCounts
                  Minimum depth required in the normal for a SNP to be considered (optional,
                  default=10).
                  A boolean indicating whether data is 'chr'-based (e.g. 'chr1' instead of '1';
is_chr_based
                  optional, default=F).
                  A vector containing the names of chromosomes to be considered (optional,
chrom_names
                  default=c(1:22,'X')). Do not set it to paste0('chr',c(1:22,'X')) if data is 'chr'-
                  Minimum base quality required for a read to be counted (optional, default=20).
min_base_qual
                  Minimum mapping quality required for a read to be counted (optional, de-
min_map_qual
                  fault=35).
ref.fasta
                  FASTA file used for generating CRAMs (optional, default=NA).
                  A boolean to generate QC reports as PNGs (optional, default=T).
plotQC
```

ascat.runAscat ascat.runAscat

#### **Description**

ASCAT main function, calculating the allele-specific copy numbers

#### Usage

```
ascat.runAscat(
 ASCATobj,
 gamma = 0.55,
 pdfPlot = F,
 y_limit = 5,
 circos = NA,
 min_ploidy = 1.5,
 max_ploidy = 5.5,
 min_purity = 0.1,
 max_purity = 1.05,
 rho_manual = NA,
  psi_manual = NA,
  img.dir = ".",
  img.prefix = ""
 write_segments = F
)
```

#### **Arguments**

ASCATobj an ASCAT object from ascat.aspcf

gamma technology parameter, compaction of Log R profiles (expected decrease in case

of deletion in diploid sample, 100% aberrant cells; 1 in ideal case, 0.55 of Illu-

mina 109K arrays)

ascat.runAscat 19

pdfPlot	Optional flag if nonrounded plots and ASCAT profile in pdf format are desired. Default=F
y_limit	Optional parameter determining the size of the y axis in the nonrounded plot and ASCAT profile. Default=5
circos	Optional file to output the non-rounded values in Circos track format. Default=NA
min_ploidy	optional numerical parameter determining the minimum boundary of the ploidy solution search space (expert parameter, don't adapt unless you know what you're doing). Default=1.5
max_ploidy	optional numerical parameter determining the maximum boundary of the ploidy solution search space (expert parameter, don't adapt unless you know what you're doing). Default=5.5
min_purity	optional numerical parameter determining the minimum boundary of the purity solution search space (expert parameter, don't adapt unless you know what you're doing). Default=0.1
max_purity	optional numerical parameter determining the maximum boundary of the purity solution search space (expert parameter, don't adapt unless you know what you're doing). Default=1.05
rho_manual	optional argument to override ASCAT optimization and supply rho parameter (expert parameter, don't adapt unless you know what you're doing).
psi_manual	optional argument to override ASCAT optimization and supply psi parameter (expert parameter, don't adapt unless you know what you're doing).
img.dir	directory in which figures will be written
img.prefix	prefix for figure names
write_segments	Optional flag to output segments in text files (.segments_raw.txt and .segments.txt under img.dir). Default= $F$

#### **Details**

Note: for copy number only probes, nA contains the copy number value and nB = 0.

#### Value

an ASCAT output object, containing:

- 1. nA: copy number of the A allele
- 2. nB: copy number of the B allele
- 3. purity: the tumour purity of all arrays
- 4. aberrantcellfraction: the aberrant cell fraction (=tumour purity) of all arrays
- 5. ploidy: the ploidy of all arrays
- 6. failedarrays: arrays on which ASCAT analysis failed
- 7. nonaberrantarrays: arrays on which ASCAT analysis indicates that they show virtually no aberrations
- 8. segments: an array containing the copy number segments of each sample (not including failed arrays)
- 9. segments\_raw: an array containing the copy number segments of each sample without any rounding applied
- 10. distance\_matrix: distances for a range of ploidy and tumor percentage values

20 ascat.synchroniseFiles

```
ascat.synchroniseFiles
```

Synchronise SNPs across files

#### Description

Synchronise SNPs across files

#### Usage

```
ascat.synchroniseFiles(
  samplename,
  tumourLogR_file,
  tumourBAF_file,
  normalLogR_file,
  normalBAF_file
)
```

#### **Arguments**

```
samplename String, name of the sample.
tumourLogR_file File where LogR from the tumour will be read and overwritten.
tumourBAF_file File where BAF from the tumour will be read and overwritten.
normalLogR_file File where LogR from the normal will be read and overwritten.
normalBAF_file File where BAF from the normal will be read and overwritten.
```

#### Author(s)

tl

## **Index**

```
ascat.asmultipcf, 2
ascat.aspcf, 3
ascat.correctLogR, 4
ascat.GCcorrect, 5
ascat.getAlleleCounts, 5
\verb|ascat.getBAFsAndLogRs|, 6
ascat.loadData, 7
ascat.metrics, 8
ascat.plotAdjustedAscatProfile, 9
{\tt ascat.plotAscatProfile, 10}
ascat.plotGenotypes, 11
ascat.plotNonRounded, 12
ascat.plotRawData, 13
{\tt ascat.plotSegmentedData, 13}
ascat.plotSunrise, 14
ascat.predictGermlineGenotypes, 14
ascat.prepareHTS, 15
{\tt ascat.prepareTargetedSeq},\,17
ascat.runAscat, 18
ascat.synchroniseFiles, 20
```