# Package 'ASCAT'

June 20, 2016

Type Package

Title Allele-Specific Copy Number Analysis of Tumours

Version 2.4.3
<b>Date</b> 2016-06-20
Author Peter Van Loo [aut, cre], Gro Nielsen [ctb], Jiqiu Cheng [ctb], Stefan Dentro [ctb], Kerstin Haase [ctb]
Maintainer Kerstin Haase < Kerstin. Haase@crick.ac.uk >
<b>Description</b> R package of ASCAT as published in http://www.ncbi.nlm.nih.gov/pubmed/20837533
<b>Depends</b> R (>= 2.13.0), RColorBrewer, grDevices, graphics, stats, utils
License GPL-3
LazyLoad yes
RoxygenNote 5.0.1
R topics documented:
ascat.GCcorrect
ascat.loadData
ascat.plotAscatProfile
ascat.plotGenotypes
ascat.plotNonRounded
ascat.plotRawData
ascat.plotSegmentedData
ascat.plotSunrise
ascat.predictGermlineGenotypes
ascat.runAscat
Index 10

2 ascat.aspcf

ascat.aspcf	ascat.aspcf		
-------------	-------------	--	--

## **Description**

run ASPCF segmentation

#### Usage

```
ascat.aspcf(ASCATobj, selectsamples = 1:length(ASCATobj$samples),
  ascat.gg = NULL, penalty = 25)
```

# **Arguments**

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)

## **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 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.GCcorrect 3

ascat.GCcorrect ascat.GCcorrect

# Description

Corrects logR of the tumour sample(s) with genomic GC content

# Usage

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

### **Arguments**

ASCATobj an ASCAT object

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

#### **Details**

Note that probes not present in the GCcontentfile will be lost from the results

#### Value

ASCAT object with corrected tumour logR

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"))
```

# **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

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

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)
```

#### **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

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

ascat.plotGenotypes 5

#### 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)
```

6 ascat.plotRawData

#### **Arguments**

ploidy ploidy of the sample
rho purity of the sample

 ${\tt goodnessOfFit} \quad estimated \ goodness \ of \ fit$ 

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

baf segmented 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

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

## Value

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

 $ascat.plotRawData \qquad \qquad ascat.plotRawData$ 

## **Description**

Plots SNP array data

## Usage

ascat.plotRawData(ASCATobj)

# Arguments

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

#### 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)
```

# Arguments

ASCATobj an ASCAT object (e.g. from ascat.aspcf)

#### 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 aberrant cell fraction

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 {\tt GermlineGenotypes} \\ as cat. predict {\tt GermlineGenotypes}
```

# **Description**

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

## Usage

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

# **Arguments**

ASCATobj an ASCAT object platform used array platform

## **Details**

Currently possible values for platform:

AffySNP6 (default)

Custom10k

Illumina109k

IlluminaCytoSNP

Illumina610k

Illumina660k

Illumina700k

 $Illumina \\ 1 \\ M$ 

Illumina2.5M

IlluminaOmni5

Affy10k

Affy100k

Affy250k\_sty

Affy250k\_nsp

AffyOncoScan

AffyCytoScanHD

HumanCNV370quad

HumanCore12

Human Core Exome 24

HumanOmniExpress12

## Value

predicted germline genotypes

ascat.runAscat 9

ascat.runAscat
ascat.runAs

## **Description**

ASCAT main function, calculating the allele-specific copy numbers

#### Usage

```
ascat.runAscat(ASCATobj, gamma = 0.55, pdfPlot = F, y_limit = 5,
  circos = NA, rho_manual = NA, psi_manual = NA)
```

# 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 Illumina 109K arrays)
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
rho_manual	optional argument to override ASCAT optimization and supply rho parameter (not recommended)
psi_manual	optional argument to override ASCAT optimization and supply psi parameter (not recommended)

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

# **Index**

```
ascat.aspcf, 2
ascat.GCcorrect, 3
ascat.loadData, 3
ascat.plotAscatProfile, 4
ascat.plotGenotypes, 5
ascat.plotNonRounded, 5
ascat.plotRawData, 6
ascat.plotSegmentedData, 7
ascat.plotSunrise, 7
ascat.predictGermlineGenotypes, 8
ascat.runAscat, 9
```