Package ‘acnr’

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Description

This data package contains SNP array data from different types of copy-number regions. These regions were identified manually by the authors of the package and may be used to generate realistic data sets with known truth.

Details

Package: acnr
Type: Package
Title: Annotated Copy-Number Regions
Version: 0.2.2
Date: 2014-09-08
Author: Morgane Pierre-Jean and Pierre Neuvial
Maintainer: Morgane Pierre-Jean <morgane.pierrejean@genopole.cnrs.fr>
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Author(s)

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getMinorMajorCopyNumbers

Get minor and major copy number labels from region annotation labels

Description

Get minor and major copy number labels from region annotation labels

Usage

getMinorMajorCopyNumbers(region)
Arguments

region A character value, the annotation label for a copy number region. Should be encoded as ",(C1,C2)\), where

C1 denotes the minor copy number, that is, the smallest of the two parent-specific copy numbers

C2 denotes the minor copy number, that is, the smallest of the two parent-specific copy numbers

Value

A matrix with \text{length(region)} rows and two columns: C1 and C2, as described above.

References


Examples

dat <- loadCnRegionData(dataSet="GSE29172_H1395", tumorFraction=1)
regions <- unique(dat$region)
getMinorMajorCopyNumbers(regions)

Description

The GEO GSE11976 data set is a dilution series from the Illumina HumanCNV370v1 chip type (Staaf et al, 2008).

Format

A data frame with 770668 observations of 7 variables:

c total copy number (not log-scaled)

b allelic ratios in the diluted tumor sample (after TumorBoost)

genotype germline genotypes

region a character value, annotation label for the region. Should be encoded as "(C1,C2)\), where

C1 denotes the minor copy number and C2 denotes the major copy number. For example,
(1,1) Normal
(0,1) Hemizygous deletion
(0,0) Homozygous deletion
(1,2) Single copy gain
(0,2) Copy-neutral LOH
(2,2) Balanced two-copy gain
(1,3) Unbalanced two-copy gain
(0,3) Single-copy gain with LOH

cellularity A numeric value between 0 and 1, the percentage of tumor cells in the sample.


Details

These data have been processed from the files available at http://cbbp.thep.lu.se/~markus/software/BAFsegmentation/ using scripts that are included in the ‘inst/preprocessing/GSE11976’ directory of this package.

Examples

dat <- loadCnRegionData("GSE11976_CRL2324")
unique(dat$region)

--------------------------------------------------------------------------------

GSE13372_HCC1143 Annotated copy-number regions from the GEO GSE13372 data set.

--------------------------------------------------------------------------------

Description

The GEO GSE13372 data set is from the Affymetrix GenomeWideSNP_6 chip type. We have extracted one tumor/normal pair corresponding to the breast cancer cell line HCC1143. For consistency with the other data sets in the package the tumor and normal samples are labeled according to their tumor cellularity, that is, 100

Format

A data frame with 205842 observations of 7 variables:

c total copy number (not log-scaled)
b allelic ratios in the diluted tumor sample (after TumorBoost)
genotype germline genotypes
bT allelic ratios in the diluted tumor sample (before TumorBoost)
bN allelic ratios in the matched normal sample
region a character value, annotation label for the region. Should be encoded as "(C1,C2)", where C1 denotes the minor copy number and C2 denotes the major copy number. For example,
(1,1) Normal
(0,1) Hemizygous deletion
(0,0) Homozygous deletion
(1,2) Single copy gain
(0,2) Copy-neutral LOH
(2,2) Balanced two-copy gain
(1,3) Unbalanced two-copy gain
(0,3) Single-copy gain with LOH

**genotype** the (germline) genotype of SNPs. By definition, rows with missing genotypes are interpreted as non-polymorphic loci (a.k.a. copy number probes).

**cellularity** A numeric value between 0 and 1, the percentage of tumor cells in the sample.

**Details**

These data have been processed from the files available from GEO using scripts that are included in the 'inst/preprocessing/GSE13372' directory of this package. This processing includes normalization of the raw CEL files using the CRMAv2 method implemented in the aroma.affymetrix package.

**Source**


**References**


**Examples**

```r
dat <- loadCnRegionData("GSE13372_HCC1143")
unique(dat$region)
```
Annotated copy-number regions from the GEO GSE29172 (and GSE26302) data sets.

Description

The GEO GSE29172 data set is a dilution series from the Affymetrix GenomeWideSNP_6 chip type. The GEO GSE26302 data set contains the experiment corresponding to the matched normal (i.e. 0% dilution).

Format

A data frame with 770668 observations of 7 variables:

- `c`: total copy number (not log-scaled)
- `b`: allelic ratios in the diluted tumor sample (after TumorBoost)
- `genotype`: germline genotypes
- `bT`: allelic ratios in the diluted tumor sample (before TumorBoost)
- `bN`: allelic ratios in the matched normal sample
- `region`: a character value, annotation label for the region. Should be encoded as "(C1,C2)", where C1 denotes the minor copy number and C2 denotes the major copy number. For example,
  - (1,1) Normal
  - (0,1) Hemizygous deletion
  - (0,0) Homozygous deletion
  - (1,2) Single copy gain
  - (0,2) Copy-neutral LOH
  - (2,2) Balanced two-copy gain
  - (1,3) Unbalanced two-copy gain
  - (0,3) Single-copy gain with LOH
- `genotype`: the (germline) genotype of SNPs. By definition, rows with missing genotypes are interpreted as non-polymorphic loci (a.k.a. copy number probes).
- `cellularity`: A numeric value between 0 and 1, the percentage of tumor cells in the sample.

Details

These data have been processed from the files available from GEO using scripts that are included in the 'inst/preprocessing/GSE29172' directory of this package. This processing includes normalization of the raw CEL files using the CRMAv2 method implemented in the aroma.affymetrix package.

Source

References


Examples

dat <- loadCnRegionData("GSE29172_H1395")
unique(dat$region)

listDataSets

List available data sets

Description

List available data sets

Usage

listDataSets()

Value

name of one of the data sets of the package, see listDataSets

Examples

listDataSets()
listTumorFractions  
*List of available tumor fractions for a data set*

**Description**
List of available tumor fractions for a data set

**Usage**
```
listTumorFractions(dataSet)
```

**Arguments**
- `dataSet`  
The name of a data set from the package, see `listDataSets`

**Value**
A numeric vector, the available tumor fractions for a data set

**Examples**
```
dataSets <- listDataSets()
fracs <- listTumorFractions(dataSets[1])
```

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loadCnRegionData  
*loadCnRegionData*

**Description**
Load real, annotated copy number data

**Usage**
```
loadCnRegionData(dataSet, tumorFraction = 1)
```

**Arguments**
- `dataSet`  
  name of one of the data sets of the package, see `listDataSets`
- `tumorFraction`  
  proportion of tumor cells in the “tumor” sample (a.k.a. tumor cellularity). See `listTumorFractions`. 

loadCnRegionData

Details

This function is a wrapper to load real genotyping array data taken from
* a dilution series from the Affymetrix GenomeWideSNP_6 chip type (Rasmussen et al, 2011), see GSE29172_H1395
* a dilution series from the Illumina HumanCNV370v1 chip type (Staaf et al, 2008), see GSE11976_CRL2324
* a tumor/normal pair from the Affymetrix GenomeWideSNP_6 chip type (Chiang et al, 2008), see GSE13372_HCC1143

Value

a data.frame containing copy number data for different types of copy number regions. Columns:

- c Total copy number
- b Allele B fraction (a.k.a. BAF)
- region a character value, annotation label for the region. Should be encoded as ")(C1,C2)", where C1 denotes the minor copy number and C2 denotes the major copy number. For example,
  - (1,1) Normal
  - (0,1) Hemizygous deletion
  - (0,0) Homozygous deletion
  - (1,2) Single copy gain
  - (0,2) Copy-neutral LOH
  - (2,2) Balanced two-copy gain
  - (1,3) Unbalanced two-copy gain
  - (0,3) Single-copy gain with LOH
- muN the (germline) genotype of SNPs. By definition, rows with missing genotypes are interpreted as non-polymorphic loci (a.k.a. copy number probes).

Author(s)

Morgane Pierre-Jean and Pierre Neuvial

Examples

```r
affyDat <- loadCnRegionData(dataSet="GSE29172_H1395", tumorFraction=1)
str(affyDat)

illuDat <- loadCnRegionData(dataSet="GSE11976_CRL2324", tumorFraction=.79)
str(illuDat)

affyDat2 <- loadCnRegionData(dataSet="GSE13372_HCC1143", tumorFraction=1)
str(affyDat2)
```
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