## Package ‘CopyNeutralIMA’

### May 2, 2024

**Title**  
Copy Neutral Illumina Methylation Arrays

**Version**  
1.22.0

**Description**  
Provides a set of genomic copy neutral samples hybridized using Illumina Methylation arrays (450k and EPIC).

**Depends**  
R (>= 3.5.0)

**License**  
Artistic-2.0

**Encoding**  
UTF-8

**LazyData**  
true

**biocViews**

- ExperimentData
- Homo_sapiens_Data
- MicroarrayData
- TwoChannelData
- MethylationArrayData
- GEO

**Imports**

- ExperimentHub
- Rdpack (>= 0.8)

**Suggests**

- BiocStyle
- knitr
- rmarkdown
- minfi
- conumee
- minfiData

**RdMacros**

- Rdpack

**VignetteBuilder**

- knitr

**RoxygenNote**  
6.0.1

**NeedsCompilation**  
no

**git_url**  
https://git.bioconductor.org/packages/CopyNeutralIMA

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CopyNeutralIMA-package

Reference samples for analysis of IlluminaHumanMethylation arrays.

Description

The CopyNeutralIMA package provides a set of 51 samples for IlluminaHumanMethylation450k arrays and a set of 13 samples for IlluminaHumanMethylationEPIC arrays.

Details

The package includes two RGChannelSetExtended objects, one with 51 samples hybridized with the IlluminaHumanMethylation450k array and another one with 13 samples hybridized with the IlluminaHumanMethylationEPIC array.

For the 450k arrays, samples from GEO series GSE49618 (Ley et al. 2013), GSE61441 (Wei et al. 2015) and GSE106089 (Tomlinson et al. 2017) are provided.

For the EPIC arrays, samples from GEO series GSE86831/ GSE86833 #’ (Pidsley et al. 2016), GSE98990 (Zhou et al. 2017) and GSE100825 (Guastafierro et al. 2017) are also provided.

The provided samples consist of material from healthy patients with nominally no copy number aberrations.

Users may use this data package as controls for their copy number profiling analysis or for testing other tools.

Check the vignette on how to use these data as control samples for conumee.

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References


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**getCopyNeutralRGSet**  
*Accessor to the data objects.*

**Description**

`getCopyNeutralRGSet` simplifies the access to the data of the package in ExperimentHub. The allowed values matched of those in the array definition of the RGChannelSet objects from package ‘minfi’. If ‘ima’ is set to ‘IlluminaHumanMethylation450k’ it will return the object with index ‘EH1453’ in ExperimentHub; if set to ‘IlluminaHumanMethylationEPIC’ it will return the object with index ‘EH1454’.

**Usage**

```r
getCopyNeutralRGSet(ima = c("IlluminaHumanMethylation450k", 
"IlluminaHumanMethylationEPIC"))
```

**Arguments**

- **ima**

  A character string specifying for which array type to retrieve data. Valid values are ‘IlluminaHumanMethylation450k’ and ‘IlluminaHumanMethylationEPIC’.

**Value**

A `RGChannelSet-class` object

**Examples**

```r
rgset_450k <- getCopyNeutralRGSet("IlluminaHumanMethylation450k")
rgset_450k
```
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