

Package ‘h5vcData’

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Type Package

Title Example data for the h5vc package

Version 1.114.0

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Description This package contains the data used in the vignettes and examples of the 'h5vc' package

License GPL (>= 3)

Suggests h5vc

biocViews CancerData

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R topics documented:

h5vcData-package 1

Index 3

h5vcData-package	<i>Example data for the h5vc package</i>
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Description

This package contains the example data needed for the vignettes and examples of the h5vc package.

Details

Package: h5vcData
Type: Package
Version: 1.0.0
Date: 2013-10-16
License: |GPL (>= 3)

This package contains the following files in inst/extdata:

example.tally.hfs5: The example HDF5 tally file

NRAS.AML.bam: BAM file containig reads spanning the NRAS locus from an AML sample

NRAS.AML.bam.bai: BAM file index for NRAS.AML.bam

NRAS.Control.bam: BAM file containig reads spanning the NRAS locus from the matched control sample

NRAS.Control.bam.bai: BAM file index for NRAS.Control.bam

Pt*bam: BAM file containing reads spannign DNMT3A locus of cancer or control samples from a total of 6 pairs Pt*bam.bam: Corresponding index files for the set of bam files overlapping the DNMT3A locus

This package contains the following data objects in data:

variantCalls is the data.frame containing a set of example variant calls on the example tally file

Author(s)

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See Also

[h5vc](#)

Examples

```
tallyFile <- system.file("extdata", "example.tally.hfs5", package = "h5vcData")
caseBamFile <- system.file("extdata", "NRAS.AML.bam", package = "h5vcData")
controlBamFile <- system.file("extdata", "NRAS.Control.bam", package = "h5vcData")
data( "example.variants", package = "h5vcData" )
head(variantCalls)
```

Index

*Topic **package**

h5vcData-package, [1](#)

h5vc, [2](#)

h5vcData (h5vcData-package), [1](#)

h5vcData-package, [1](#)

variantCalls (h5vcData-package), [1](#)