

Package ‘flowWorkspaceData’

April 4, 2023

Type Package

Title A data package containing two flowJo, one diva xml workspace and the associated fcs files as well as three GatingSets for testing the flowWorkspace, openCyto and CytoML packages.

Version 3.10.0

Date 2011-03-31

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Description The necessary external data to run the flowWorkspace and openCyto vignette is found in this package.

biocViews ExperimentData, FlowCytometryData

License GPL-2

LazyLoad yes

git_url <https://git.bioconductor.org/packages/flowWorkspaceData>

git_branch RELEASE_3_16

git_last_commit 36f5e80

git_last_commit_date 2022-11-01

Date/Publication 2023-04-04

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flowWorkspaceData-package

A flowJo XML Workspace and Associated Data for testing the flow-Workspace package.

Description

This package contains fcs data files and xml workspaces for testing the flowWorkspace and openCyto packages and building the their vignettes. These can be located in the inst/extdata directory of the flowWorkspaceData package. The sample files and workspaces contain data from whole blood. The primary purpose of this data set is to test the import code and export code in the flowWorkspace package and running automated gating pipeline in the openCyto package.

Details

Package:	flowWorkspaceData
Type:	Package
Version:	1.1.0
Date:	2011-03-31
License:	GPL 2.0
LazyLoad:	yes

flowWorkspaceDataInfo() provides information about the package.

Author(s)

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See Also

[flowWorkspace](#) [openCyto](#)

Examples

```
flowWorkspaceDataInfo()
```

flowWorkspaceDataInfo *Print some information about this package and the data.*

Description

Contains the data necessary to build the flowWorkspace and openCyto vignettes.

Usage

```
flowWorkspaceDataInfo()
```

Author(s)

Greg Finak

See Also

[flowWorkspace-package](#) [openCyto](#)

Examples

```
flowWorkspaceDataInfo();
```

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