

iFlow

October 25, 2011

guiEnv

Accessor to the internal GUI environment

Description

All state information for the GUI is stored in an internal environment. This function can be used to access this environment.

Usage

```
## .guiEnv$list name such as Gate.list, workflowlist
```

Value

The internal gui environment.

Author(s)

Kyongryun Lee

Examples

```
## 1. Run iFlow
## Not run: iflow()
## 2. Load data
## Not run: data(GvHD)
## 3. Activate data by clicking a data name on Data panel
## 4. Create a gate
## 5. It is able to access created gate objects or workflow of the data on the command line
## The workflow is automatically created whenever the operations of the data are run.

## wf.list <- .guiEnv$workflow.list
## wf.list

## gt.list <- .guiEnv$Gate.list
## gt.list

## Not run: iFlow:::guiEnv$Gatelist
```

iFlow-package *iFlow*

Description

GUI based visualization for preprocessing of analyzing Flow Cytometry data.

Details

Package:

Type:

Version:

Date:

License:

LazyLoad:

The tutorial video can be downloaded from <http://bioconductor.fhcrc.org/docs/workflows/flowcytometry/tutorial.mpeg>

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References

<http://www.hindawi.com/journals/abi/2009/103839.html>

iflow *Main function for iFlow*

Description

Main function for iFlow. It calls up the main graphical user interface (GUI) of the iFlow package.

Usage

`iflow()`

Details

The GUI Pull-Down Menu consists of "File", "Data", "Graphics", "Gate", "ProbBin", and "Help" selections. The user can start with "File" pull-down menu to load data files of interest. The data file can be in either rda or FCS format. Please see the vignette or tutorial video for more detail (<http://bioconductor.fhcrc.org/docs/workflows/flowcytometry/tutorial.mpeg>).

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Examples

```
## on R console
iflow()
data(ITN)
```

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