

Package ‘SuperCurveGUI’

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Title 'SuperCurve' GUI Package

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Description Graphical User Interface for RPPA analysis using the 'SuperCurve' package.

Depends R (>= 2.15)

Imports methods, tcltk, SuperCurve (>= 1.5), tclish (>= 1.0)

SystemRequirements Tcl/Tk

NeedsCompilation no

URL <http://supercurve.r-forge.r-project.org/#SuperCurveGUI>

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LazyLoad yes

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R topics documented:

SuperCurveGUI-package	2
supercurveGUI	2

Index	4
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SuperCurveGUI-package *GUI for reverse phase protein lysate array analysis*

Description

A package for analyzing reverse phase protein lysate arrays (RPPA).

Details

Package: SuperCurveGUI
Type: Package
Version: 2.1.5
Date: 2015-07-27
License: Artistic-2.0

For a complete list of functions, use `library(help="SuperCurveGUI")`.
For a high-level summary of the changes for each revision, use
`file.show(system.file("NEWS", package="SuperCurveGUI"))`.

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supercurveGUI *Graphical User Interface for the SuperCurve package*

Description

Graphical User Interface for the SuperCurve package.

Usage

```
supercurveGUI()  
scui()
```

Details

The function `supercurveGUI` launches a Graphical User Interface for the **SuperCurve** package. The GUI uses Tk widgets (via the R-Tcl/Tk interface by Peter Dalgaard) in order to provide a simple interface to the SuperCurve functions for analyzing reverse phase protein lysate arrays.

The function `scui` is an alias for the aforementioned function.

If set, the environment variable `SC_DIR` will be used as the initial location for the directory containing quantification files.

Resources

The Tk options database is loaded from user-specific defaults files, such as `$.Xdefaults`, and resource databases loaded into the X server. The package will attempt to load its resources upon startup, following the X11R5 method of merging app-default files from multiple sources. The standard X11 paths are searched first, followed by paths specified by the environment variables `XFILESEARCHPATH`, `XAPPLRESDIR`, `XUSERFILESEARCHPATH`, respectively. Unlike X11, **ALL** matching files will be loaded, not just the first.

One way to make use of the options database is to create a file named `supercurveGUI.ad` in your home directory, specifying any desired resources. Set your `XUSERFILESEARCHPATH` environment variable within R with the command:

```
Sys.setenv(XUSERFILESEARCHPATH=file.path(path.expand("~"), "%N.ad"))
```

For example, to cause **SuperCurve** to convert each quantification file as it's read from a single subgrid physical layout into its actual logical one by default, add the following resource value to the app-defaults file mentioned above:

```
*software: superslide
```

Author(s)

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Examples

```
## Not run:
supercurveGUI()
# -or-
scui()

## End(Not run)
```

Index

*Topic **IO**

supercurveGUI, [2](#)

*Topic **package**

SuperCurveGUI-package, [2](#)

scui (supercurveGUI), [2](#)

supercurveGUI, [2](#)

SuperCurveGUI-package, [2](#)