sprof internal

July 28, 2013

Encoding UTF-8

Type Package

2 sprof-package

sprof	-package	sprof: Analysis of R profiles	
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Description

Profiling, timing and optimization utilitites

Details

Package: sprof Type: Package Version: 0.0-5 Date: 2013-07-23

License: GPL-2 | GPL-3

 $\frac{readRprof()}{reads} \ a \ profile \ file \ from \ \frac{Rprof()}{reads} \ or \ other \ profilers \ and \ returns \ a \ composite \ structure \ of \ class \ sprof.$ The basic components of sprof are (conceptually) four data frames

info general information and summaries

nodes node specific information

stacks node specific information. Stacks are random snapshots from the program execution, possibly including side in profiles collected records of a profile, encoded as references to stacks

To create a profile on the fly, use sampleRprof.

To import profile information written by Rprof or other profilers, use readRprof.

For sprof, the usual access functions are supported.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

```
## Not run:
data(sprof01lm)
```

adjacency 3

```
str(sprof01lm)
plot(sprof01lm)
## End(Not run)
```

adjacency

sprof to adjacency matix

Description

convert node information from a sprof structure to adjacancy matrix.

Usage

```
adjacency(sprof,\,keep.names = TRUE,\,rmzero = TRUE)
```

Arguments

sprof a sprof structure.

keep.names boolean. Copy node names as row- and column names.

rmzero boolean. Remove nodes with no edges.

Value

a correspondency matrix

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

References

See the vignette of package sprof.

```
data(sprof01lm)
adjacency(sprof01lm)
```

barplot_s

|--|

Description

Creates a sorted bar plot with vertical or horizontal bars.

Usage

```
barplot_s(height,
sort_by, decreasing = TRUE,
lowtrim, hightrim, trimlegend = TRUE,
col, coli, colfun,
main, ...)
```

Arguments

height	either a vector or matrix of values describing the bars which make up the plot. See barplot.
sort_by	a variable to sort by. Defaults to height.
decreasing	boolean. Sorting direction.
lowtrim	A optional lower trim value. Observations with sort by values up to lowtrim are discarded.
hightrim	A optional upper trim value. Observations with $\operatorname{sort_by}$ values from lowtrim are discarded.
trimlegend	Boolean. Show a legend about trimmed values.
col	a vector of colors for the bars or bar components. By default, grey is used if height is a vector, and a gamma-corrected grey palette if height is a matrix.
coli	An index into the col table, based on original sorting.
colfun	A function to generate a col palette. grey is rescaled to 1n.
main	overall title for the plot

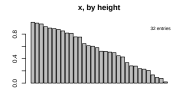
Details

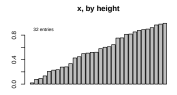
...

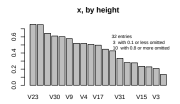
To come. Plots are from this collection:

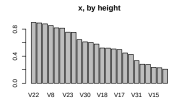
Passed to barplot

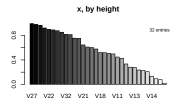
barplot_s 5

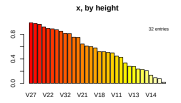












Value

Invisible: A data frame with components

x height

perm the permutation applied coli the colour index applied

col optional: the colours selected

Note

Part of this could go to the R base function barplot.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

See Also

barplot

Examples

 $x < \text{-} \, \operatorname{runif}(100)$

 $barplot_s(x)$

x <- rnorm(100)

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```
barplot\_s(x, colfun=heat.colors, lowtrim=-1) \\ rm(x)
```

list.as.matrix

Convert list to matrix

Description

Convert list to matrix. List entries go to matrix columns, filled for equal length.

Usage

```
list.as.matrix(x, filler = NA)
```

Arguments

x a list of numeric vectors.

filler a value to be used as a filler

Value

A matrix with the values from x, filled to matrix shape.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

Examples

```
\begin{array}{l} x <- \operatorname{list}(x1{=}c(1{,}2{,}3){,}x2{=}3,\; x3{=}4{:}8) \\ \operatorname{list.as.matrix}(x) \\ \operatorname{list.as.matrix}(x, \operatorname{filler}{=}0) \end{array}
```

nodepackage

Find a package that may contain a node

Description

getAnywhere() is used to look up x, and the package or namespace information is used to give a source package.

Usage

```
nodepackage(x)
```

Arguments

 \mathbf{x}

a character string or name, or a vector.

nodescloud 7

Details

There is no indication whether the information is from a namespace or from a package information.

No indication is given if multiple hits are encountered.

The information is based on the run time environment of this function. This may be different from the environment the object is taken from.

See the help information for getAnywhere() for more warnings.

Value

a character string or a vector of strings with the package names.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

See Also

```
getAnywhere()
```

Examples

```
nodepackage("getAnywhere")
```

nodescloud

Nodescloud of nodes from profile data

Description

Show the nodes from a profile, with class encoded as colour and frequency encoded as size.

Usage

```
nodescloud(sprof, min.freq = 3, icol, col)
```

Arguments

sprof A data structure as returned by readRprof.

min.freq Minimum frequency to be included.

icol An index vector to colour palette. Defaults to sprof\$nodes\$icol.

col A colour palette.

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Details

Note: these figures may be ourdated. Please run the examples.

Plots are from this collection:



Value

Used for the side effect of showing the plots.

Note

See the vignette of package sprof.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

```
## Not run:
data(sprof01lm)
nodescloud(sprof01lm)
## End(Not run)
```

plot.sprof 9

plot.sprof

plot for profiles

Description

plot a plot for the output of class scode.

Usage

```
## S3 method for class 'sprof' plot(x, ...)
```

Arguments

x A data structure as returned by readRprof.

... further arguments passed to or from other methods.

Value

subject to change

Note

See the vignette for in-context explanations.

Displays of the graph structure are given in the vignette.

References

```
http://sintro.r-forge.r-project.org/
```

See Also

```
summaryRprof
plot_profiles
plot_nodes plot_stacks
```

```
{\rm data(sprof01lm)} {\rm plot.sprof(sprof01lm)}
```

plot_nodes

plot nodes

Plot profiling information on node level.

Description

Various plots of a profile.

Usage

```
\begin{array}{l} plot\_nodes(x,\,which=c(1L,\,2L,\,3L,\,4L),\,col=NULL,\\ ask=prod(par("mfcol")) < length(which) \&\& \,dev.interactive(),\\ src=NULL,\,mincount=5,\,horiz=FALSE,\,\ldots) \end{array}
```

Arguments

x preferably a sprof object. Other data structures may be extended

which Selector of plots to show.

col Colour table

ask boolean. Ask for a new page?

src String to be used as source identifier.

mincount minimum total frequency count for node to be shown in barcharts.

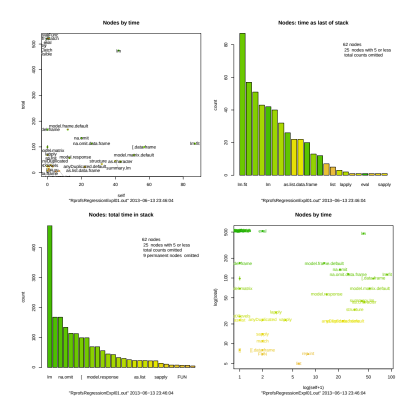
horiz draw horizontal bar plots.

... passed.

Details

Plots are from this collection:

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Value

To come.

Note

See the vignette of package sprof.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

References

See the vignette of package sprof.

See Also

See Also as plot.sprof, ~~~

```
\begin{array}{l} data(sprof01lm) \\ oldpar <- \ par(mfrow=c(2,2)) \\ plot\_nodes(sprof01lm) \\ par(oldpar) \end{array}
```

plot_profiles

plot	profile
piot	ргоше

Plot profiling information on profile level.

Description

Various plots of a profile.

Usage

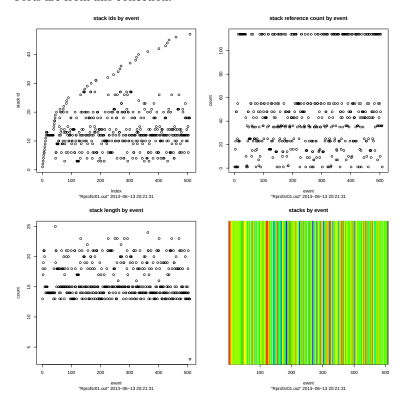
```
\begin{array}{l} plot\_profiles(x,\,which=c(1L,\,2L,\,3L,\,4L),\,col,\\ ask=prod(par("mfcol"))< length(which)\,\,\&\&\,\,dev.interactive(),\\ src=NULL,\,...) \end{array}
```

Arguments

X	preferably a sprof object. Other data structures may be extended
which	Selector of plots to show.
col	Colour table
ask	boolean. Ask for a new page?
src	String to be used as source identifier.
	passed.

Details

Plots are from this collection:



plot_stacks 13

Note

See the vignette of package sprof.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

References

See the vignette of package sprof.

See Also

```
See Also as plot.sprof, ~~~
```

Examples

```
data(sprof01lm)
oldpar <- par(mfrow=c(2,2))
plot_profiles(sprof01lm)
par(oldpar)
```

 $plot_stacks$

Plot profiling information on stack level.

Description

Various plots of a profile.

Usage

```
\begin{array}{l} plot\_stacks(x,\,which=c(1L,\,2L),\\ ask=prod(par("mfcol")) < length(which) \,\&\&\,\,dev.interactive(),\\ src=NULL,\,mincount=5,\,horiz=FALSE,\,\ldots) \end{array}
```

Arguments

x preferably a sprof object. Other data structures may be extended
--

which Selector of plots to show.
ask boolean. Ask for a new page?

src String to be used as source identifier.

mincount minimum total frequency count for stack to be shown in barcharts.

horiz draw horizontal bar plots.

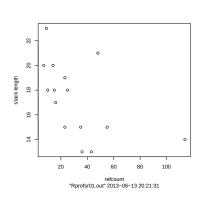
... passed.

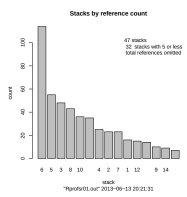
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Details

Note: these figures may be outdated. Please run the examples.

Plots are from this collection:





Value

To come.

Note

See the vignette of package sprof.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

References

See the vignette of package sprof.

See Also

See Also as plot.sprof, ~~~

```
data(sprof01lm)
oldpar <- par(mfrow=c(2,2))
plot_stacks(sprof01lm)
par(oldpar)
```

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print.sprof

print for profiles

Description

Print a print for the output of class scode.

Usage

```
## S3 method for class 'sprof' print(x, ...)
```

Arguments

x A data structure as returned by readRprof.

... further arguments passed to or from other methods.

Value

None.

References

```
http://sintro.r-forge.r-project.org/
```

See Also

```
{\bf summary} {\bf Rprof\ plot.sprof}
```

Examples

```
data(sprof01lm)
print(sprof01lm)
```

print_profiles

Print profile information

Description

Print profile information.

Usage

```
print\_profiles(x)
```

Arguments

Х

a sprof data structure.

profiles_matrix

Value

none

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

Examples

```
\frac{\mathrm{data}(\mathrm{sprof01lm})}{\mathrm{print\_profiles}(\mathrm{sprof01lm})}
```

profiles_matrix

Extract a node incidence matrix from profile information.

Description

Extract a node incidence matrix from profile information.

Usage

```
profiles_matrix(x)
```

Arguments

Х

an sprof data structure.

Value

an incidence matrix, NA filled.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

```
data(sprof01lm)
smat <-profiles_matrix(sprof01lm)
image(smat)
```

readRprof 17

readRprof	Read Rprof Output files and Stack Logs	

Description

Read a log of stack entries, such as the output of the Rprof function, and generate a more accessible representation.

Usage

```
\label{eq:readRprof} \begin{split} & \operatorname{readRprof}(\text{filename} = "Rprof.out", \ \operatorname{chunksize} = 5000, \\ & \operatorname{interval} = 0.02, \\ & \operatorname{head} = \operatorname{c}("\operatorname{auto"}, \, "\operatorname{none"}, \, "\operatorname{Rprofmem"}), \\ & \operatorname{id} = \operatorname{NULL}) \end{split}
```

Arguments

filename Name of a file produced by Rprof(). chunksize Number of lines to read at a time.

interval Real: time interval between samples, in s.

head c("auto", "none", "Rprofmem") to interpret control information as provided

by Rprof or Rprofmem. See details.

id An optional identification string. Defaults to filename and date.

Details

This function reads a log file of stacks, one stack snapshot per line, stack entries separated by space.

As profiling output file could be very large, it is read in blocks of chunksize lines. Increasing chunksize will make the function run faster if sufficient memory is available.

The input format is controlled by the head argument. Format "auto" tries to detect control lines as interspersed by Rprof. These lines are not included in the output.

Value

This data structure is subject to change.

Temporarily: A list with components

firstline A verbatim copy of the first line of the input file. Typically this contains timing

or formatting information.

nodes A vector of node names. This may include stray entries from interperspersed

lines.

stacks A vector of unique stacks found in input, stored as verbatim copies.

stacksrenc A list of unique stacks in top down order (top first), encoded as vectors or refer-

ences to stacks.

data A vector encoding the data file as referenes to stacks.

[&]quot;none" igores all control information and includes these lines as strange stacks.

[&]quot;Rprofmem" isolates headers as provided by Rprofmem. new page entries are encoded as malloc requests with length 0.

18 readRprof

timesRLE Vector of sampling intervals, in miliseconds. Run-length encoded.

freq A frequency table summarizing data.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>, based on the code of summaryRprof

References

```
http://sintro.r-forge.r-project.org/
```

See Also

```
summaryRprof
summaryRprof
flatProfile in library(proftools).
{parse_rprof} in library(profr).
The chapter on "Tidying and profiling R code" in "Writing R Extensions" (see the 'doc/manual' subdirectory of the R source tree).
Rprof is a sampling profiler.
tracemem traces copying of an object via the C function duplicate.
Rprofmem is a non-sampling memory-use profiler.
```

Examples

```
\#\# Not run:
## Rprof() is not available on all platforms
profinterval <- 0.001
simruns < -100
n < -10000
x < -runif(n)
y0 < -2 + 3 * x
sinknull <- textConnection(NULL, "w"); sink(sinknull)</pre>
Rprof(tmp <- tempfile(), interval = profinterval)
for (i in 1:simruns) \{y \leftarrow y0 + rnorm(n); xxx \leftarrow summary(lm(y^x))\}
Rprof()
Rprof\_out < - readRprof(tmp)
unlink(tmp)
sink(); close(sinknull)
str(Rprof out)
## End(Not run)
```

http://developer.r-project.org/memory-profiling.html

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rrle

Recursive run length encoding.

Description

Encode a matrix as run-length, top down. Encoding respects previous runs, e.g line 2 encodes rns in each run of line 1.

Usage

```
rrle(x)
```

Arguments

 \mathbf{X}

a matrix.

Value

list of run length encoded lines

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

See Also

```
rrleb, ~~~
```

```
 \begin{array}{l} x <- \ matrix(c(\\ 1,1,1,2,2,\\ 3,3,4,4,4,\\ 5,5,6,6,7,\\ 8,9,9,0,0\\ ),nrow=4,\ ncol\ =5,\ byrow=TRUE)\\ xrrle <-\ rrle(x)\\ xrrle \\ t(sapply(xrrle,\ inverse.rle)) \end{array}
```

20 rrleb

rrleb

Recursive run length encoding bottom up.

Description

Encode a matrix as run-length, bottom up. Encoding respects previous runs, e.g line n-1 encodes rns in each run of line n.

May be removed.

Usage

```
rrleb(x)
```

Arguments

х

a matrix.

Value

list of run length encoded lines

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

See Also

```
rle, ~~~
```

```
 \begin{array}{l} x <- \ matrix(c(\\ 3,3,4,4,4,\\ 5,5,6,6,7,\\ 8,9,9,0,0,\\ 1,1,1,2,2 \\ \\ ),nrow=4,\ ncol\ =5,\ byrow=TRUE)\\ xrrleb <- \ rrleb(x)\\ xrrleb \\ t(sapply(xrrleb,\ inverse.rle)) \\ \end{array}
```

sampleRprof 21

sampleRprof

Get a sample profile

Description

Get a sample profile and return it as a sprof data structure.

Usage

```
sampleRprof(expr, runs = NULL, gcFirst = TRUE, interval = 0.001, ...)
```

Arguments

expr an expression to be profiled.

runs nr of runs to profile. gcFirst boolean. Profile GC.

interval Real: time interval between samples, in s. ... additional parameters, passed to Rprof

Value

A list of type sprof

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

References

```
http://sintro.r-forge.r-project.org/
```

Examples

```
res\_lm <- sampleRprof(for \ (i \ in \ 1:1000) \ yy <- \ lm(runif(1000) \ \~rnorm(1000)), \ runs = 100)
```

shownodes

Show node information from a profile

Description

Plot node information from a profile in various plots.

Usage

```
shownodes(sprof, col)
```

Arguments

sprof A data structure as returned by readRprof.

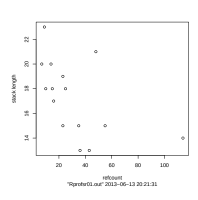
col A colour palette for the plots.

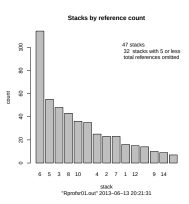
sprof01lm

Details

Note: these figures may be outdated. Please run the examples.

Plots are from this collection:





Value

Used for the side effect of showing the plots.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

Examples

```
## Not run:
data(sprof01lm)
shownodes(sprof01lm)
## End(Not run)
```

sprof01lm

sprof sample data

Description

An example data set for the functions in package sprof.

Usage

data(sprof01lm)

Format

The format is: A List of 4 \$ info :'data.frame': 1 obs. of 8 variables: \$ nodes :'data.frame': 62 obs. of 5 variables: \$ stacks :'data.frame': 50 obs. of 7 variables: \$ profiles:List of 4

References

See the vignette of package sprof.

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Examples

```
data(sprof01lm)
str(sprof01lm)
plot(sprof01lm)
```

stackstoadj

Stacks to adjacency matrix

Description

convert stack information to adjacency matrix

Usage

```
stackstoadj(xstacks, xfreq, maxnode)
```

Arguments

xstacks list of stack ids

xfreq vector of frequencies or weights

maxnode maximum of nodes (maybe higher then in stacks)

Value

the adjacency matrix

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

Examples

#

 str_prof

str for sprof objects

Description

```
str for sprof objects
```

Usage

```
str\_prof(x)
```

Arguments

 \mathbf{X}

an sprof object

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Author(s)

Günther Sawitzki @r-forge.r-project.org>

Examples

```
\frac{\mathrm{data}(\mathrm{sprof01lm})}{\mathrm{str\_prof}(\mathrm{sprof01lm})}
```

summary.sprof

Summary for profiles

Description

Print a summary for the output of class scode.

Usage

```
## S3 method for class 'sprof' summary(object, ...)
```

Arguments

object A data structure as returned by readRprof.

... further arguments passed to or from other methods.

Value

None.

References

```
http://sintro.r-forge.r-project.org/
```

See Also

```
{\bf summary} {\bf Rprof}
```

```
## Not run:  
## Rprof() is not available on all platforms  
profinterval <- 0.001  
simruns <- 100  

n <- 10000  
x <- runif(n)  
y0 <- 2+ 3 * x  

sinknull <- textConnection(NULL, "w"); sink(sinknull)  
Rprof(tmp <- tempfile(), interval = profinterval)  
for (i in 1:simruns) \{y <- y0 + rnorm(n); xxx <- summary(lm(y^x))\}  
Rprof()
```

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```
Rprof_out <- readProf(tmp)
unlink(tmp)
sink(); close(sinknull)
summary(Rprof_out)
## End(Not run)</pre>
```

 $summary_terminals$

Tabulate leaf nodes

Description

Tabulate leaf nodes

Usage

```
summary_terminals(x)
```

Arguments

x

an sprof data structure.

Value

A table of frequencies, bystack.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

Examples

```
\frac{\mathrm{data}(\mathrm{sprof}01\mathrm{lm})}{\mathrm{summary\_terminals}(\mathrm{sprof}01\mathrm{lm})}
```

updateRprof

Update statistics and tables in a sprof obejct

Description

Synchronize information from profiles and stack tables, and update statistics.

Usage

```
updateRprof(sprof)
```

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Arguments

sprof A data structure as returned by readRprof.

Value

An updated sprof data structure.

Note

See the vignette of package sprof.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

writeRprof

Write profile data

Description

Write a profile data file from a sprof data structure.

Usage

```
writeRprof(sprof, filename = "Rprof.Out")
```

Arguments

sprof a data structure from package sprof

filename The file to be used for exporting the profiling results.

Details

writeRprof only writes the stack entries for the profile.

This can be used to export information after preprocessing with sprof to some package designed for Rprof output.

Value

An invisble list with the profile entries, headers removed.

Note

See the vignette for in-context explanations.

Displays of the graph structure are given in the vignette.

Author(s)

Günther Sawitzki <gsawitzki@r-forge.r-project.org>

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References

http://sintro.r-forge.r-project.org/

```
## Not run:
data(sprof01lm)
writeRprof(sprof01lm)
## End(Not run)
```

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