

# Package ‘autoReg’

October 12, 2022

**Type** Package

**Title** Automatic Linear and Logistic Regression and Survival Analysis

**Version** 0.2.6

**URL** <https://github.com/cardiomoon/autoReg>,  
<https://cardiomoon.github.io/autoReg/>

**BugReports** <https://github.com/cardiomoon/autoReg/issues>

**Description** Make summary tables for descriptive statistics and select explanatory variables automatically in various regression models. Support linear models, generalized linear models and cox-proportional hazard models. Generate publication-ready tables summarizing result of regression analysis and plots. The tables and plots can be exported in “HTML”, “pdf(LaTex)”, “docx(MS Word)” and “pptx(MS Powerpoint)” documents.

**License** GPL-3

**LazyData** true

**Encoding** UTF-8

**Imports** moonBook(>= 0.3.0), nortest, dplyr, crayon, stringr, tidyr,  
purrr, survival, mice, officer, flextable, rlang, patchwork,  
ggplot2, boot, broom, tidycmprsk, scales, maxstat, pammtools

**Suggests** knitr, finalfit, lme4, TH.data, rmarkdown, survminer, asaur,  
cmprsk, PairedData

**RoxygenNote** 7.1.2

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**NeedsCompilation** no

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---

addFitSummary	<i>Add model summary to an object of class gaze</i>
---------------	---

---

**Description**

Add model summary to an object of class gaze

**Usage**

```
addFitSummary(df, fit, statsname = "")
```

**Arguments**

df                    An object of class "gaze" or "autoReg"  
 fit                   An object of class "glm" or "lm" or "crr"  
 statsname           character Name of statistics

**Value**

addFitSummary returns an object of `gaze` or `autoReg` - the same class as `df`

**Examples**

```
require(survival)
require(dplyr)
data(cancer, package="survival")
fit=coxph(Surv(time, status)~rx+age+sex+nodes+obstruct+perfor, data=colon)
df=autoReg(fit, uni=FALSE)
final=fit2final(fit)
df %>% addFitSummary(final, statsname="HR (final)") %>% myft()
```

---

addLabelData

*Add labels to data*

---

**Description**

Add labels to data

**Usage**

```
addLabelData(data)
```

**Arguments**

data                A data.frame

**Value**

A data.frame

**Examples**

```
addLabelData(data.frame(ph.ecog=0:3, sex=c(1, 2, 2, 2), age=c(20, 30, 40, 70)))
```

---

adjustedPlot	<i>Draw an expected plot</i>
--------------	------------------------------

---

### Description

Draw an expected plot

### Usage

```
adjustedPlot(
  fit,
  xnames = NULL,
  pred.values = list(),
  newdata = NULL,
  maxy.lev = 5,
  median = TRUE,
  facet = NULL,
  se = FALSE,
  mark.time = FALSE,
  show.median = FALSE,
  type = "ggplot",
  ...
)
```

### Arguments

fit	An object of class "coxph" or "survreg"
xnames	Character Names of explanatory variable to plot
pred.values	A list A list of predictor values
newdata	A data.frame or NULL
maxy.lev	Integer Maximum unique length of a numeric variable to be treated as categorical variables
median	Logical
facet	Character Name of facet variable
se	logical Whether or not show se
mark.time	logical Whether or not mark time
show.median	logical
type	Character plot type
...	further arguments to be passed to plot.survfit

### Value

A ggplot or no return value(called for side effects)

## Examples

```

library(survival)
fit=coxph(Surv(time,status)~rx+logWBC,data=anderson)
adjustedPlot(fit)
adjustedPlot(fit,xnames="rx",se=TRUE,type="plot")
adjustedPlot(fit,xnames="rx",se=TRUE)
## Not run:
anderson$WBCgroup=ifelse(anderson$logWBC<=2.73,0,1)
anderson$WBCgroup=factor(anderson$WBCgroup,labels=c("low","high"))
anderson$rx=factor(anderson$rx,labels=c("treatment","control"))
fit=coxph(Surv(time,status)~rx,data=anderson)
adjustedPlot(fit,xnames=c("rx"),show.median=TRUE)
fit=coxph(Surv(time,status)~rx*WBCgroup,data=anderson)
adjustedPlot(fit,xnames=c("rx","WBCgroup"),show.median=TRUE)
adjustedPlot(fit,xnames=c("rx","WBCgroup"),facet="WBCgroup",show.median=TRUE)
data(cancer,package="survival")
fit=coxph(Surv(time,status)~rx+strata(sex)+age+differ,data =colon)
adjustedPlot(fit,xnames=c("sex"))
adjustedPlot(fit,xnames=c("sex"),pred.values=list(age=58,differ=3))
adjustedPlot(fit,xnames=c("sex","rx"),facet="sex")
adjustedPlot(fit,xnames=c("rx","sex","differ"),facet=c("sex","rx"),se=TRUE)
fit <- coxph(Surv(start, stop, event) ~ rx + number + size+ cluster(id), data = bladder2)
adjustedPlot(fit,xnames=c("rx","number","size"),facet=c("rx","size"),maxy.lev=8)

## End(Not run)

```

---

adjustedPlot.survreg    *Draw predicted survival curve with an object survreg*

---

## Description

Draw predicted survival curve with an object survreg

## Usage

```

adjustedPlot.survreg(
  x,
  xnames = NULL,
  pred.values = list(),
  maxy.lev = 5,
  median = TRUE,
  newdata = NULL,
  addCox = FALSE
)

```

## Arguments

x                    An object of class survreg

xnames	Character Names of explanatory variable to plot
pred.values	A list A list of predictor values
maxy.lev	Integer Maximum unique length of a numeric variable to be treated as categorical variables
median	Logical
newdata	A data.frame or NULL
addCox	logical Whether or not add KM

**Value**

No return value, called for side effects

**Examples**

```
library(survival)
x=survreg(Surv(time, status) ~ rx, data=anderson,dist="exponential")
adjustedPlot(x)
adjustedPlot(x,addCox=TRUE)
## Not run:
x=survreg(Surv(time, status) ~ sex, data=lung,dist="weibull")
adjustedPlot(x,addCox=TRUE)
x=survreg(Surv(time, status) ~ rx, data=anderson,dist="exponential")
adjustedPlot(x)
x=survreg(Surv(time, status) ~ ph.ecog + age + sex, data=lung, dist="weibull")
adjustedPlot(x)
adjustedPlot(x,addCox=TRUE)
adjustedPlot(x,pred.values=list(age=c(20,40,60,80),sex=2,ph.ecog=3),addCox=TRUE)
newdata=data.frame(ph.ecog=0:3,sex=c(1,2,2,2),age=c(20,40,60,80))
adjustedPlot(x,newdata=newdata,addCox=TRUE)

## End(Not run)
```

---

adjustedPlot2

*Draw a survfitted plot*


---

**Description**

Draw a survfitted plot

**Usage**

```
adjustedPlot2(fit, se = FALSE, mark.time = FALSE)
```

**Arguments**

fit	An object of class coxph or survfit
se	logical Whether or not show se
mark.time	logical Whether or not mark time

**Value**

a ggplot

**Examples**

```
library(survival)
fit=coxph(Surv(time,status)~rx+logWBC,data=anderson)
plot(survfit(fit),conf.int=TRUE)
adjustedPlot2(fit,se=TRUE)
```

---

anderson

*Remission survival times of 42 leukemia patients*

---

**Description**

A dataset containing survival time of 42 leukemia patients

**Usage**

anderson

**Format**

A data.frame with 42 rows and 5 variables

**time** survival time in weeks

**status** censoring status 1=failure 0=censored

**sex** sex 0=Female 1=Male

**logWBC** log white blood cell count

**rx** treatment status 1=control 0=treatment

**Source**

David G. Kleinbaum and Mitchel Klein. Survival Analysis. A Self-Learning Text(3rd ed)(Springer,2012)  
ISBN: 978-1441966452



---

anderson1

*Remission survival times of 42 leukemia patients*

---

### Description

A dataset containing survival time of 42 leukemia patients This data is the same data with anderson, but sex and rx variable are factors not numeric

### Usage

anderson1

### Format

A data.frame with 42 rows and 5 variables

**time** survival time in weeks

**status** censoring status 1=failure 0=censored

**sex** sex "Female" or "Male"

**logWBC** log white blood cell count

**rx** treatment status "treatment" or "control"

### Source

David G. Kleinbaum and Mitchel Klein. Survival Analysis. A Self-Learning Text(3rd ed)(Springer,2012)  
ISBN: 978-1441966452

---

anderson2

*Remission survival times of 31 leukemia patients*

---

### Description

This data is subdata of anderson with medium( $2.3 < \log WBC \leq 2.96$ ) and high WBC count( $\log WBC > 2.96$ )

### Usage

anderson2

**Format**

A data.frame with 31 rows and 6 variables

**time** survival time in weeks

**status** censoring status 1=failure 0=censored

**sex** sex 0=Female 1=Male

**logWBC** log white blood cell count

**rx** treatment status 1=control 0=treatment

**WBCCAT** WBC count group 1=medium 2=high

**Details**

A dataset containing survival time of 31 leukemia patients

**Source**

David G. Kleinbaum and Mitchel Klein. Survival Analysis. A Self-Learning Text(3rd ed)(Springer,2012)  
ISBN: 978-1441966452

---

as\_printable

*Convert data.frame to printable form*

---

**Description**

Calculate character length and apply all data

**Usage**

```
as_printable(  
  data,  
  align.first = "left",  
  align.chr = "right",  
  align.num = "right"  
)
```

**Arguments**

data	A data.frame
align.first	character Alignment of first variable
align.chr	character Alignment of character variable
align.num	character Alignment of numeric variable

**Value**

A data.frame

**Examples**

```
as_printable(mtcars)
as_printable(iris)
```

---

autoReg	<i>Perform univariable and multivariable regression and stepwise backward regression automatically</i>
---------	--

---

**Description**

Perform univariable and multivariable regression and stepwise backward regression automatically

**Usage**

```
autoReg(x, ...)
```

## S3 method for class 'lm'  
autoReg(x, ...)

## S3 method for class 'glm'  
autoReg(x, ...)

## S3 method for class 'coxph'  
autoReg(x, ...)

## S3 method for class 'survreg'  
autoReg(x, ...)

**Arguments**

x	An object of class lm, glm or coxph
...	Further arguments

**Value**

autoReg returns an object of class "autoReg" which inherits from the class "data.frame" with at least the following attributes:

**attr(\*,"yvars)** character. name of dependent variable

**attr(\*,"model")** name of model. One of "lm", "glm" or "coxph"

**Methods (by class)**

- lm: S3 method for a class lm
- glm: S3 method for a class glm
- coxph: S3 method for a class coxph
- survreg: S3 method for a class survreg

**Examples**

```

data(cancer, package="survival")
fit=glm(status~rx+sex+age+obstruct+nodes, data=colon, family="binomial")
autoReg(fit)
autoReg(fit, uni=FALSE, final=TRUE)
autoReg(fit, uni=FALSE, imputed=TRUE)
fit=lm(mpg~wt*hp+am+I(wt^2), data=mtcars)
autoReg(fit, final=TRUE)
autoReg(fit, imputed=TRUE)

```

---

autoRegCox	<i>perform automatic regression for a class of coxph</i>
------------	--

---

**Description**

perform automatic regression for a class of coxph

**Usage**

```

autoRegCox(
  x,
  threshold = 0.2,
  uni = FALSE,
  multi = TRUE,
  final = FALSE,
  imputed = FALSE,
  keepstats = FALSE,
  ...
)

```

**Arguments**

x	An object of class coxph
threshold	numeric
uni	logical whether or not perform univariable regression
multi	logical whether or not perform multivariable regression
final	logical whether or not perform stepwise backward elimination
imputed	logical whether or not perform multiple imputation
keepstats	logical whether or not keep statistic
...	Further arguments to be passed to gaze()

**Value**

autoRegCox returns an object of class "autoReg" which inherits from the class "data.frame" with at least the following attributes:

**attr(\*,"yvars)** character. name of dependent variable

**attr(\*,"model")** name of model. One of "lm", "glm" or "coxph"

**Examples**

```

require(survival)
require(dplyr)
data(cancer)
fit=coxph(Surv(time,status==2)~log(bili)+age+cluster(edema),data=pubc)
autoReg(fit)
fit=coxph(Surv(time,status)~rx+age+sex+nodes+obstruct+perfor,data=colon)
autoReg(fit)
autoReg(fit,uni=TRUE,threshold=1)
autoReg(fit,uni=TRUE,final=TRUE) %>% myft()
data(colon_s,package="finalfit")
fit=coxph(Surv(time,status)~age.factor+sex.factor+obstruct.factor+perfor.factor,data=colon_s)
autoReg(fit,uni=TRUE,threshold=1)
autoReg(fit,uni=TRUE,imputed=TRUE)

```

---

autoRegsurvreg	<i>perform automatic regression for a class of survreg</i>
----------------	--

---

**Description**

perform automatic regression for a class of survreg

**Usage**

```

autoRegsurvreg(
  x,
  threshold = 0.2,
  uni = FALSE,
  multi = TRUE,
  final = FALSE,
  imputed = FALSE,
  keepstats = FALSE,
  mode = 1,
  ...
)

```

**Arguments**

x	An object of class survreg
threshold	numeric
uni	logical whether or not perform univariable regression
multi	logical whether or not perform multivariable regression
final	logical whether or not perform stepwise backward elimination
imputed	logical whether or not perform multiple imputation
keepstats	logical whether or not keep statistic
mode	integer
...	Further arguments to be passed to gaze()

**Value**

autoRegSurvreg returns an object of class "autoReg" which inherits from the class "data.frame" with at least the following attributes:

**attr(\*,"yvars)** character. name of dependent variable

**attr(\*,"model")** name of model. One of "lm","glm","coxph" or "survreg"

**Examples**

```
require(survival)
require(dplyr)
data(cancer)
fit=survreg(Surv(time,status)~rx+age+sex+nodes+obstruct+perfor,data=colon)
autoReg(fit)
autoReg(fit,uni=TRUE,threshold=1)
autoReg(fit,uni=TRUE,final=TRUE)
autoReg(fit,uni=TRUE,final=TRUE) %>% myft()
## Not run:
autoReg(fit,mode=2)
autoReg(fit,uni=TRUE,threshold=1,,mode=2)
autoReg(fit,uni=TRUE,final=TRUE,mode=2)
autoReg(fit,uni=TRUE,final=TRUE,mode=2) %>% myft()
autoReg(fit,final=TRUE,imputed=TRUE) %>% myft()
autoReg(fit,final=TRUE,imputed=TRUE,mode=2) %>% myft()

## End(Not run)
```

---

autoReg_sub	<i>Perform univariable and multivariable regression and stepwise backward regression automatically</i>
-------------	--

---

**Description**

Perform univariable and multivariable regression and stepwise backward regression automatically

**Usage**

```
autoReg_sub(
  fit,
  threshold = 0.2,
  uni = FALSE,
  multi = TRUE,
  final = FALSE,
  imputed = FALSE,
  keepstats = FALSE,
  showstats = TRUE,
  ...
)
```

**Arguments**

fit	An object of class lm or glm
threshold	numeric
uni	logical whether or not perform univariate regression
multi	logical whether or not perform multivariate regression
final	logical whether or not perform stepwise backward elimination
imputed	logical whether or not include imputed model
keepstats	logical whether or not keep statistics
showstats	logical whether or not show descriptive statistics
...	Further arguments to be passed to imputedReg()

**Value**

An object of class "autoReg" which inherits from the class "data.frame" with at least the following attributes:

**attr(\*,"yvars)** character. name of dependent variable

**attr(\*,"model")** name of model. One of "lm","glm" or "coxph"

---

beNumeric

*Whether a string vector can be converted to numeric*


---

**Description**

Whether a string vector can be converted to numeric

**Usage**

```
beNumeric(x)
```

**Arguments**

x                    A string vector

**Value**

A logical vector

---

bootPredict	<i>Bootstrap simulation for model prediction</i>
-------------	--

---

**Description**

Generate model predictions against a specified set of explanatory levels with bootstrapped confidence intervals.

**Usage**

```
bootPredict(fit, newdata, R = 100, type = "response", ...)
```

**Arguments**

fit	An object of class lm or glm
newdata	A data.frame
R	Number of simulations. Note default R=100 is very low.
type	he type of prediction required, see predict.glm. The default for glm models is on the scale of the response variable. Thus for a binomial model the default predictions are predicted probabilities.
...	Further arguments to be passed to boot::boot

**Value**

An object of class "data.frame"

**Examples**

```
data(GBSG2,package="TH.data")
fit=glm(cens~horTh+pnodes,data=GBSG2,family="binomial")
newdata=expand.grid(horTh=factor(c(1,2),labels=c("no","yes")),pnodes=1:51)
bootPredict(fit,newdata)
library(survival)
fit=coxph(Surv(time,cens)~age+horTh+progrec+pnodes,data=GBSG2)
```

---

countGroups	<i>Count groups</i>
-------------	---------------------

---

**Description**

Count groups

**Usage**

```
countGroups(data, yvars)
```



**Arguments**

data	A data.frame
yvars	variable names

**Value**

An object of class "tibble"

**Examples**

```
library(moonBook)
countGroups(acs, "sex")
countGroups(acs, c("sex", "Dx"))
```

---

 coxzphplot

*Graphical Test of Proportional Hazards*


---

**Description**

This is a ggplot version of plot.cox.zph. Displays a graph of the scaled Schoenfeld residuals, along with a smooth curve.

**Usage**

```
coxzphplot(x, resid = TRUE, se = TRUE, var = NULL, hr = FALSE, add.lm = FALSE)
```

**Arguments**

x	result of the cox.zph function.
resid	a logical value, if TRUE the residuals are included on the plot, as well as the smooth fit.
se	a logical value, if TRUE, confidence bands at two standard errors will be added.
var	The set of variables for which plots are desired. It can be integer or variable names
hr	logical If true, plot for hazard ratio, If false, plot for coefficients
add.lm	logical If true, add linear regression line

**Value**

A faceted ggplot

**Examples**

```
library(survival)
vfit <- coxph(Surv(time,status) ~ trt + factor(celltype) + karno + age, data=veteran, x=TRUE)
x <- cox.zph(vfit)
coxzphplot(x)
coxzphplot(x, var="karno", add.lm=TRUE)
```

---

crr2stats	<i>Extract statistics from an object of class crr</i>
-----------	---

---

**Description**

Extract statistics from an object of class crr

**Usage**

```
crr2stats(x, digits = 2)
```

**Arguments**

x	an object of class crr
digits	integer indication the position of decimal place

**Value**

An object of class "data.frame"

**Examples**

```
data(melanoma, package="boot")
melanoma$status_crr=ifelse(melanoma$status==1,1,ifelse(melanoma$status==2,0,2))
x=crrFormula(time+status_crr~age+sex+thickness+ulcer,data=melanoma)
crr2stats(x)
```

---

crrFormula	<i>Competing Risk Regression with Formula</i>
------------	---

---

**Description**

Competing Risk Regression with Formula

**Usage**

```
crrFormula(x, data, ...)
```

**Arguments**

x	formula time+status~explanatory variables
data	data a data.frame
...	Further arguments to be passed to <a href="#">crr</a>

**Value**

An object of class "tidycrr" which is described in [crr](#)

**Examples**

```
data(melanoma, package="boot")
melanoma$status_crr=ifelse(melanoma$status==1,1,ifelse(melanoma$status==2,0,2))
crrFormula(time+status_crr~age+sex+thickness+ulcer,data=melanoma)
```

---

descNum	<i>Make description for numeric summary</i>
---------	---

---

**Description**

Make description for numeric summary

**Usage**

```
descNum(method = 1, p = NULL)
```

**Arguments**

method	numeric
p	A numeric or NULL

**Value**

A character vector of length 1

---

df2flexible	<i>Convert data.frame to flexible</i>
-------------	---------------------------------------

---

**Description**

Convert data.frame to flexible

**Usage**

```
df2flectable(
  df,
  vanilla = FALSE,
  fontname = NULL,
  fontsize = 12,
  add.rownames = FALSE,
  even_header = "transparent",
  odd_header = "#5B7778",
  even_body = "#EFEFEF",
  odd_body = "transparent",
  vlines = TRUE,
  colorheader = FALSE,
  digits = 2,
  digitp = 3,
  align_header = "center",
  align_body = "right",
  align_rownames = "left",
  NA2space = TRUE,
  pcol = NULL,
  ...
)
```

**Arguments**

df	A data.frame
vanilla	A Logical
fontname	Font name
fontsize	font size
add.rownames	logical. Whether or not include rownames
even_header	background color of even_header
odd_header	background color of even_header
even_body	background color of even_body
odd_body	background color of even_body
vlines	Logical. Whether or not draw vertical lines
colorheader	Logical. Whether or not use color in header
digits	integer indicating the number of decimal places
digitp	integer indicating the number of decimal places of p values
align_header	alignment of header. Expected value is one of 'left', 'right', 'center', 'justify'.
align_body	alignment of body. Expected value is one of 'left', 'right', 'center', 'justify'.
align_rownames	alignment of rownames. Expected value is one of 'left', 'right', 'center', 'justify'.
NA2space	A logical. If true, convert NA value to space

pcol            An integer indicating p value. If specified, convert value less than 0.01 to "< 0.001" in given column.

...            further arguments to be passed to [flextable](#)

**Value**

An object of class "flextable" which is described in [flextable](#)

---

drawline	<i>draw line character</i>
----------	----------------------------

---

**Description**

draw line character

**Usage**

```
drawline(n)
```

**Arguments**

n            Numeric

**Value**

No return value, called for side effects

**Examples**

```
drawline(10)
```

---

expectedPlot	<i>Draw an adjusted Plot for a numeric predictor</i>
--------------	--

---

**Description**

Select cutpoint for a numeric predictor with `maxstat.test()` and draw survival plot with this cutpoint

**Usage**

```

expectedPlot(
  fit,
  xname = NULL,
  no = 2,
  maxy.lev = 5,
  median = TRUE,
  mark.time = FALSE,
  se = FALSE,
  type = "ggplot",
  ...
)

```

**Arguments**

<code>fit</code>	An object of class "coxph"
<code>xname</code>	Character Name of explanatory variable to plot
<code>no</code>	integer Number of groups to be made
<code>maxy.lev</code>	Integer Maximum unique length of a numeric variable to be treated as categorical variables
<code>median</code>	Logical
<code>mark.time</code>	logical Whether or not mark time
<code>se</code>	logical Whether or not show se
<code>type</code>	Character "plot" or "ggplot"
<code>...</code>	further arguments to be passed to <code>plot.survfit</code>

**Value**

No return value, called for side effects

**Examples**

```

library(survival)
data(cancer, package="survival")
fit=coxph(Surv(time,status)~age+sex,data =colon)
expectedPlot(fit,xname="age")
fit=coxph(Surv(time,status)~rx+logWBC,data=anderson)
expectedPlot(fit,xname="logWBC",no=3)

```

---

filldown	<i>filldown vector with lead value</i>
----------	--

---

**Description**

filldown vector with lead value

**Usage**

```
filldown(x, what = c("", NA))
```

**Arguments**

x	a vector
what	Values to be filled

**Value**

A vector with the same class as x

**Examples**

```
x=rep(1:5,each=3)
x=removeDup(x,NA)
filldown(x)
```

---

find1stDup	<i>Find first duplicated position</i>
------------	---------------------------------------

---

**Description**

Find first duplicated position

**Usage**

```
find1stDup(x)
```

**Arguments**

x	a vector
---	----------

**Value**

A logical vector

**Examples**

```
x=rep(1:5,each=3)
which(find1stDup(x))
```

---

findDup	<i>Find duplicated term</i>
---------	-----------------------------

---

**Description**

Find duplicated term

**Usage**

```
findDup(x)
```

**Arguments**

x	A vector
---	----------

**Value**

A logical vector

**Examples**

```
x=rep(1:5, each=3)
findDup(x)
x=c(6,x)
findDup(x)
which(!findDup(x))
```

---

fit2final	<i>Make final model using stepwise backward elimination</i>
-----------	---

---

**Description**

Make final model using stepwise backward elimination

**Usage**

```
fit2final(fit, threshold = 0.2)
```

**Arguments**

fit	An object of class "coxph"
threshold	Numeric

**Value**

An object of class "coxph" which is described in [coxph](#)



**Examples**

```
require(survival)
data(cancer)
fit=coxph(Surv(time,status)~age+sex+obstruct+perfor,data=colon)
final=fit2final(fit)
fit2summary(final)
```

---

fit2lik	<i>extract likelihood information with a coxph object</i>
---------	---

---

**Description**

extract likelihood information with a coxph object

**Usage**

```
fit2lik(x)
```

**Arguments**

x                    An object of class "coxph" or "survreg"

**Value**

A string

**Examples**

```
library(survival)
fit=coxph(Surv(time,status) ~rx,data=anderson)
fit2lik(fit)
```

---

fit2list	<i>Make a list of univariable model with multivariable regression model</i>
----------	---

---

**Description**

Make a list of univariable model with multivariable regression model

**Usage**

```
fit2list(fit)
```

**Arguments**

fit                    An object of class "lm" or "glm"

**Value**

An object of class "fitlist" which is a list of objects of class "lm" or "glm"

**Examples**

```
library(survival)
data(cancer)
fit=glm(status~rx+sex+age+obstruct+nodes,data=colon,family="binomial")
fit2list(fit)
fit=lm(mpg~wt*hp+am,data=mtcars)
fit2list(fit)
```

---

fit2model

*Restore fit model data containing AsIs expressions*

---

**Description**

Restore fit model data containing AsIs expressions

**Usage**

```
fit2model(fit)
```

**Arguments**

`fit` An object of class "lm", "glm" or "coxph"

**Value**

An object of class "data.frame"

**Examples**

```
require(survival)
pbc$status2=ifelse(pbc$status==2,1,0)
fit=coxph(Surv(time,status2)~age+log(bili),data=pbc)
fit2model(fit)
```

---

fit2multi	<i>Make multivariable regression model by selecting univariable models with p.value below threshold</i>
-----------	---

---

**Description**

Make multivariable regression model by selecting univariable models with p.value below threshold

**Usage**

```
fit2multi(fit, threshold = 0.2)
```

**Arguments**

fit	An object of class "coxph"
threshold	Numeric

**Value**

An object of class "coxph"

**Examples**

```
require(survival)
data(cancer)
fit=coxph(Surv(time,status)~age+sex+obstruct+perfor,data=colon)
fit2multi(fit)
```

---

fit2newdata	<i>Make a new data of mean value or most frequent value</i>
-------------	---

---

**Description**

Make a new data of mean value or most frequent value

**Usage**

```
fit2newdata(
  fit,
  xnames = NULL,
  pred.values = list(),
  maxy.lev = 5,
  median = TRUE,
  digits = 1
)
```

**Arguments**

fit	An object of class "coxph"
xnames	character Names of explanatory variable to plot
pred.values	A list A list of predictor values
maxy.lev	Integer Maximum unique length of a numeric variable to be treated as categorical variables
median	logical If TRUE, select median value for numerical variable. Otherwise select most frequent value
digits	integer indicating the number of decimal places

**Value**

A data.frame

**Examples**

```
require(survival)
data(cancer, package="survival")
fit=coxph(Surv(time, status)~rx+sex+age, data=colon)
fit=coxph(Surv(time, status)~rx+age+strata(sex), data=colon)
fit=survreg(Surv(time, status) ~ ph.ecog + age + sex, data=lung, dist="weibull")
fit2newdata(fit)
fit2newdata(fit, pred.values=list(sex=0, age=58))
fit2newdata(fit, pred.values=list(age=c(20, 40, 60, 80), sex=2, ph.ecog=3))
```

---

fit2stats

*Summarize statistics with a model*


---

**Description**

Summarize statistics with a model

**Usage**

```
fit2stats(fit, method = "default", digits = 2, mode = 1)
```

**Arguments**

fit	An object of class lm or glm or coxph or survreg
method	character choices are one of the c("likelihood", "wald")
digits	integer indicating the number of decimal places
mode	integer

**Value**

An object of class "data.frame"

## Examples

```
library(survival)
data(cancer)
fit=glm(status~rx+sex+age+obstruct+nodes,data=colon,family="binomial")
fit2stats(fit)
fit=lm(mpg~wt*hp+am,data=mtcars)
fit2stats(fit)
fit=survreg(Surv(time,status)~rx+sex+age+obstruct+nodes,data=colon)
fit2stats(fit)
```

---

fit2summary

*Summarize statistics with a model or model list*

---

## Description

Summarize statistics with a model or model list

## Usage

```
fit2summary(fit, mode = 1, ...)
```

## Arguments

fit	An object of class "lm" or "glm" or "fitlist" which is a result of <a href="#">fit2list</a>
mode	integer
...	Further argument to be passed to fit2stats

## Value

An object of class "data.frame"

## Examples

```
library(survival)
data(cancer)
fit=glm(status~rx+sex+age+obstruct+nodes,data=colon,family="binomial")
fit2summary(fit)
fitlist=fit2list(fit)
fit2summary(fitlist)
fit=survreg(Surv(time,status)~rx+sex+age+obstruct+nodes,data=colon)
fit2summary(fit)
```

---

`gaze`*Produce table for descriptive statistics*

---

### Description

Produce table for descriptive statistics by groups for several variables easily. Depending on the nature of these variables, different descriptive statistical methods were used (t-test, ANOVA, Kruskal-Wallis, chi-squared, Fisher's,...)

### Usage

```
gaze(x, ...)  
  
## S3 method for class 'formula'  
gaze(x, ...)  
  
## S3 method for class 'data.frame'  
gaze(x, ...)  
  
## S3 method for class 'coxph'  
gaze(x, ...)  
  
## S3 method for class 'survreg'  
gaze(x, ...)  
  
## S3 method for class 'glm'  
gaze(x, ...)  
  
## S3 method for class 'lm'  
gaze(x, ...)  
  
## S3 method for class 'tidycrr'  
gaze(x, ...)
```

### Arguments

<code>x</code>	An R object, formula or data.frame
<code>...</code>	arguments to be passed to gaze.data.frame or gaze.formula

### Value

An object of class "gaze" which inherits from the class "data.frame" with at least the following attributes:

**attr(\*,"yvars)** character. name of dependent variable

**Methods (by class)**

- formula: S3 method for formula
- data.frame: default S3 method
- coxph: default S3 method
- survreg: default S3 method
- glm: default S3 method
- lm: default S3 method
- tidycrr: default S3 method

**Examples**

```

library(moonBook)
library(dplyr)
gaze(acs)
gaze(~age+sex,data=acs)
gaze(sex~.,data=acs,digits=1,method=1,show.p=TRUE) %>% myft()

gaze(sex~age+Dx,data=acs)
gaze(EF~.,data=acs) %>% myft()
gaze(sex+Dx~.,data=acs,show.p=TRUE) %>% myft()
gaze(sex+Dx~.,data=acs)
gaze(Dx+sex~cardiogenicShock,data=acs,show.p=TRUE) %>% myft()
gaze(Dx+sex+HBP~cardiogenicShock,data=acs,show.p=TRUE)
gaze(~mpg+cyl,data=mtcars)
gaze(~.,data=mtcars)
gaze(cyl~.,data=mtcars,show.p=TRUE)
gaze(hp~.,data=mtcars)
gaze(cyl+am~.,data=mtcars)

library(survival)
x=coxph(Surv(time,status) ~rx,data=anderson1)
gaze(x)
x=coxph(Surv(time,status) ~rx*logWBC,data=anderson1)
gaze(x)
library(survival)
x=survreg(Surv(time, status) ~ rx, data=anderson,dist="exponential")
gaze(x)
x=survreg(Surv(time, status) ~ ph.ecog + age + sex, lung)
gaze(x)
data(cancer,package="survival")
fit=glm(status~rx+sex+age+obstruct+nodes,data=colon,family="binomial")
gaze(fit)
fit=lm(mpg~wt*hp+am+I(wt^2),data=mtcars)
gaze(fit)
data(melanoma,package="boot")
melanoma$status_crr=ifelse(melanoma$status==1,1,ifelse(melanoma$status==2,0,2))
fit=crrFormula(time+status_crr~age+sex+thickness+ulcer,data=melanoma)
gaze(fit)

```

---

gaze.formula_sub	<i>Produce table for descriptive statistics</i>
------------------	---

---

### Description

Produce table for descriptive statistics by groups for several variables easily. Depending on the nature of these variables, different descriptive statistical methods were used (t-test, ANOVA, Kruskal-Wallis, chi-squared, Fisher's, ...)

### Usage

```
## S3 method for class 'formula_sub'
gaze(x, data, missing = FALSE, ...)
```

### Arguments

x	An object of class "formula". Left side of ~ must contain the name of one grouping variable or two grouping variables in an additive way (e.g. sex+group~), and the right side of ~ must have variables in an additive way.
data	A data.frame
missing	logical If true, missing value analysis performed
...	Further arguments to be passed to gaze()

### Value

An object of class "gaze" which inherits from the class "data.frame" with at least the following attributes:

**attr(\*,"yvars)** character. name of dependent variable

---

gazeCat	<i>Summary function for categorical variable</i>
---------	--

---

### Description

Summary function for categorical variable



**Usage**

```
gazeCat(
  data,
  x,
  y = NULL,
  max.ylev = 5,
  digits = 1,
  show.total = FALSE,
  show.n = FALSE,
  show.missing = FALSE,
  show.stats = TRUE,
  origData = NULL,
  show.p = TRUE,
  method = 1,
  catMethod = 2,
  maxCatLevel = 20,
  ...
)
```

**Arguments**

<code>data</code>	A data frame
<code>x</code>	Name of a categorical variable
<code>y</code>	Name of a variable, either continuous or categorical
<code>max.ylev</code>	<code>max.ylev</code> An integer indicating the maximum number of levels of grouping variable ('y'). If a column have unique values less than <code>max.ylev</code> it is treated as a categorical variable. Default value is 5.
<code>digits</code>	Numeric
<code>show.total</code>	logical. Whether or not show total column
<code>show.n</code>	logical. Whether or not show N column
<code>show.missing</code>	logical. Whether or not show missing column
<code>show.stats</code>	logical. Whether or not show stats column
<code>origData</code>	A data.frame containing original data
<code>show.p</code>	logical. Whether or not show p column
<code>method</code>	<code>method</code> An integer indicating methods for continuous variables. Possible values in methods are 1 forces analysis as normal-distributed 2 forces analysis as continuous non-normal 3 performs a Shapiro-Wilk test or <code>nortest::ad.test</code> to decide between normal or non-normal Default value is 1.
<code>catMethod</code>	An integer indicating methods for categorical variables. Possible values in methods are <ul style="list-style-type: none"> <li><b>0</b> Perform <code>chisq.test</code> first. If warning present, perform fisher test</li> <li><b>1</b> Perform <code>chisq.test</code> without continuity correction</li> <li><b>2</b> Perform <code>chisq.test</code> with continuity correction</li> <li><b>3</b> perform <code>fisher.test</code></li> </ul>

	<b>4</b>	perform prop.trend test Default value is 2.
maxCatLevel		An integer indicating the maximum number of unique levels of categorical variable. If a column have unique values more than maxCatLevel, categorical summarization will not be performed.
...		Further arguments

**Value**

An object of class "data.frame" or "tibble"

**Examples**

```
require(moonBook)
gazeCat(acs, "Dx")
gazeCat(acs, "Dx", "smoking")
gazeCat(acs, "sex", "Dx", show.p=TRUE)
gazeCat(acs, "Dx", "sex", show.p=TRUE)
gazeCat(acs, "Dx", "EF")
gazeCat(acs, "sex", "EF", method=2)
gazeCat(mtcars, "cyl", "hp")
```

---

gazeCont

*Summary function for continuous variable*


---

**Description**

Summary function for continuous variable

**Usage**

```
gazeCont(
  data,
  x,
  y = NULL,
  max.ylev = 5,
  digits = 1,
  show.total = FALSE,
  show.n = FALSE,
  show.missing = FALSE,
  show.stats = TRUE,
  show.p = TRUE,
  method = 1,
  origData,
  ...
)
```

**Arguments**

<code>data</code>	A data.frame
<code>x</code>	A name of variable
<code>y</code>	A name of variable, either continuous or categorical
<code>max.ylev</code>	<code>max.ylev</code> An integer indicating the maximum number of levels of grouping variable ('y'). If a column have unique values less than <code>max.ylev</code> it is treated as a categorical variable. Default value is 5.
<code>digits</code>	integer indicating the number of decimal places
<code>show.total</code>	logical. Whether or not show total column
<code>show.n</code>	logical. Whether or not show N column
<code>show.missing</code>	logical. Whether or not show missing column
<code>show.stats</code>	logical. Whether or not show stats column
<code>show.p</code>	logical. Whether or not show p column
<code>method</code>	<code>method</code> An integer indicating methods for continuous variables. Possible values in methods are 1 forces analysis as normal-distributed 2 forces analysis as continuous non-normal 3 performs a Shapiro-Wilk test or <code>nortest::ad.test</code> to decide between normal or non-normal Default value is 1.
<code>origData</code>	A data.frame containing original data
<code>...</code>	Further arguments

**Value**

An object of class "data.frame" or "tibble"

**Examples**

```
gazeCont(mtcars, "hp")
gazeCont(mtcars, "hp", "mpg")
require(moonBook)
gazeCont(acs, "log(age)")
gazeCont(acs, "age", method=2)
gazeCont(acs, "age", "EF", method=2)
gazeCont(acs, "age", "Dx", method=1)
gazeCont(acs, "age", "Dx", show.p=TRUE, method=3)
```

---

gaze\_sub

*Summary function for categorical/continuous variable*


---

**Description**

Summary function for categorical/continuous variable

**Usage**

```
gaze_sub(data, xname, y = NULL, max.ylev = 5, ...)
```

**Arguments**

data	A data.frame
xname	A name of categorical/continuous vector
y	A name of vector, either continuous or categorical
max.ylev	max.ylev An integer indicating the maximum number of levels of grouping variable ('y'). If a column have unique values less than max.ylev it is treated as a categorical variable. Default value is 5.
...	Further arguments to be passed to gazeCont() or gazeCat()

**Value**

An object of class "data.frame" or "tibble"

**Examples**

```
require(moonBook)
gaze_sub(acs, "age")
gaze_sub(acs, "log(age)")
gaze_sub(acs, "I(age^2)")
gaze_sub(acs, "sex")
gaze_sub(acs, "age", "EF")
gaze_sub(acs, "sex", "EF")
gaze_sub(acs, "age", "Dx")
gaze_sub(acs, "sex", "Dx")
gaze_sub(iris, "Species", "Sepal.Length")
```

---

getInteraction	<i>Get interaction data from data</i>
----------------	---------------------------------------

---

**Description**

Get interaction data from data

**Usage**

```
getInteraction(name, data)
```

**Arguments**

name	a string with interaction term
data	a data.frame

**Value**

An object of class "data.frame"

**Examples**

```
data(acs, package="moonBook")
getInteraction("TC:Dx:sex", data=acs)
```

---

getN

*Get number of data specified by 'name' and 'desc'*


---

**Description**

Get number of data specified by 'name' and 'desc'

**Usage**

```
getN(name, desc, data)
```

**Arguments**

name	a string with interaction term
desc	character
data	a data.frame

**Value**

A numeric vector

**Examples**

```
data(acs, package="moonBook")
df=getInteraction("TC:Dx:sex", data=acs)
getN(name=df$name, desc=df$desc, data=acs)
```

---

getSigVars

*Get explanatory variables of a model with significance level below the threshold*


---

**Description**

Get explanatory variables of a model with significance level below the threshold

**Usage**

```
getSigVars(fit, threshold = 0.2, final = TRUE)
```

**Arguments**

<code>fit</code>	An object of class <code>lm</code> or <code>glm</code>
<code>threshold</code>	Numeric
<code>final</code>	logical if true, perform stepwise regression using <code>step()</code>

**Value**

A list containing the following components:

**sigVars** names of explanatory variables which have significant levels below the threshold in univariable model

**finalVars** names of explanatory variables included in final model as a result of `step`

**Examples**

```
library(survival)
data(cancer, package="survival")
fit=glm(status~rx+sex+age+obstruct+nodes, data=colon, family="binomial")
getSigVars(fit)
fit=lm(mpg~hp*wt+am, data=mtcars)
getSigVars(fit)
```

---

ggcmprsk

---

*Draw Cumulative Incidence Curves for Competing Risks*


---

**Description**

Draw Cumulative Incidence Curves for Competing Risks

**Usage**

```
ggcmprsk(x, data, id = NULL, se = FALSE, strata = NULL, facet = NULL, ...)
```

**Arguments**

<code>x</code>	A formula as <code>time+status~1</code>
<code>data</code>	A <code>data.frame</code>
<code>id</code>	character vector label for status
<code>se</code>	logical whether or not show confidence interval
<code>strata</code>	character vector label for strata
<code>facet</code>	numeric if facet is not <code>NULL</code> , draw plot with selected facets
<code>...</code>	Further arguments to be passed to <code>tidycmprsk::cuminc</code>

**Value**

An object of class `"ggplot"`

**Examples**

```

data(melanoma, package="boot")
melanoma$status1 = ifelse(melanoma$status==1, 1, ifelse(melanoma$status==2, 0, 2))
ggcmprsk(time/365+status1~1, data=melanoma)
ggcmprsk(time/365+status1~1, data=melanoma, id=c("alive", "melanoma", "other"), se=TRUE)
ggcmprsk(time/365+status1~sex, data=melanoma)
ggcmprsk(time/365+status1~sex, data=melanoma, facet=1)
ggcmprsk(time/365+status1~sex, data=melanoma,
id=c("alive", "melanoma", "other"), strata=c("female", "male"))
ggcmprsk(time/365+status1~sex, data=melanoma,
id=c("alive", "melanoma", "other"), strata=c("female", "male"), facet=1)

```

ggcmprsk2

*Compare cumulative incidence to th Kaplan-Meier estimate***Description**

Compare cumulative incidence to th Kaplan-Meier estimate

**Usage**

```

ggcmprsk2(
  x,
  data,
  id = c("disease", "other"),
  se = FALSE,
  xpos = c(2, 2),
  ypos = c(0.25, 0.7),
  ylabs = NULL,
  xlab = NULL,
  label = NULL,
  plot = TRUE
)

```

**Arguments**

x	A formula as time+status~1
data	A data.frame
id	Character vector of length 2
se	logical whether or not show confidence interval
xpos	numeric x-axis position of label
ypos	numeric y-axis position of label
ylabs	string vector of length 2. y axis labels
xlab	A character. The x-axis label
label	string vector of length 2. Label names
plot	logical Whether or not print plot

**Value**

A list containing the following components:

**df** A long-form data.frame consist of time, est, upper,lower, id, method

**df3** A data.frame for label consist of x, y, label, id

**p** A ggplot object

**Examples**

```
require(dplyr)
data(prostateSurvival,package="asaur")
prostateHighRisk <- prostateSurvival %>%
  filter(grade=="poor" & stage=="T2",ageGroup=="80+")
ggcmprsk2(survTime/12+status~1,data=prostateHighRisk,
  id=c("prostate cancer","other causes"))
```

---

highlight2

*Highlight a data.frame*


---

**Description**

Highlight a data.frame

**Usage**

```
highlight2(x, i = NULL, j = NULL, style = NULL, include.colname = FALSE)
```

**Arguments**

x	A data.frame
i	numeric rows to highlight
j	numeric columns to highlight
style	A style function or NULL
include.colname	logical Whether or not include colname

**Value**

a data.frame



**Examples**

```

head(mtcars) %>% highlight2(i=3) %>% printdf()
library(crayon)
head(mtcars) %>% highlight2(i=2) %>% highlight2(j=3,style=blue$bold) %>% printdf()
fit=lm(mpg~wt*hp,data=mtcars)
gaze(fit)
gaze(fit) %>% highlight2(j=4,include.colname=TRUE)
gaze(fit) %>% highlight2(i=2,j=4) %>% highlight2(i=2,j=2:3,style=blue$bold)
gaze(fit) %>% highlight2(i=2) %>% highlight2(j=3,style=blue$bold)

```

---

imputedReg	<i>Make a multiple imputed model</i>
------------	--------------------------------------

---

**Description**

Make a multiple imputed model

**Usage**

```
imputedReg(fit, data = NULL, m = 20, seed = 1234, digits = 2, mode = 1, ...)
```

**Arguments**

<code>fit</code>	An object of class <code>lm</code> , <code>glm</code> , <code>coxph</code> or <code>survreg</code>
<code>data</code>	a <code>data.frame</code>
<code>m</code>	Number of multiple imputations. The default is <code>m=20</code> .
<code>seed</code>	An integer that is used as argument by the <code>set.seed()</code> for offsetting the random number generator.
<code>digits</code>	Integer indicating the number of decimal place
<code>mode</code>	integer indicating summary mode of class <code>survreg</code>
<code>...</code>	Further argument to be passed to <code>mice</code>

**Value**

An object of class "imputedReg" which inherits from the class "data.frame"

**Examples**

```

data(cancer,package="survival")
fit=glm(status~rx+sex+age+obstruct+nodes,data=colon,family="binomial")
imputedReg(fit)

library(survival)
fit=coxph(Surv(time,status)~rx+age+sex+nodes+obstruct+perfor,data=colon)
imputedReg(fit)
fit=survreg(Surv(time,status)~rx+age+sex+nodes+obstruct+perfor,data=colon)
imputedReg(fit)
imputedReg(fit,mode=2)

```

---

is.mynumeric	<i>Decide whether a vector can be treated as a numeric variable</i>
--------------	---

---

**Description**

Decide whether a vector can be treated as a numeric variable

**Usage**

```
is.mynumeric(x, maxy.lev = 5)
```

**Arguments**

x	A vector
maxy.lev	An integer indicating the maximum number of unique values of a numeric variable be treated as a categorical variable

**Value**

A logical value

**Examples**

```
x=1:5  
is.mynumeric(x)  
x=1:13  
is.mynumeric(x)
```

---

loglogplot	<i>Draw log-log plot</i>
------------	--------------------------

---

**Description**

Draw log-log plot

**Usage**

```
loglogplot(  
  fit,  
  xnames = NULL,  
  main = NULL,  
  labels = NULL,  
  no = 3,  
  add.loess = FALSE,  
  add.lm = TRUE,  
  type = "l",
```

```

    se = TRUE,
    what = "surv",
    legend.position = NULL,
    ...
  )

```

### Arguments

<code>fit</code>	An object of class "coxph" or "survfit"
<code>xnames</code>	character Names of explanatory variable to plot
<code>main</code>	String Title of plot
<code>labels</code>	String vector Used as legend in legend
<code>no</code>	Numeric The number of groups to be converted
<code>add.loess</code>	logical If true, add loess regression line
<code>add.lm</code>	logical If true, add linear regression line
<code>type</code>	character "l" or "p"
<code>se</code>	logical If true, add se
<code>what</code>	character One of c("surv","survOdds","failureOdds")
<code>legend.position</code>	legend position. One of c("left","top","bottom","right") or numeric vector of length 2.
<code>...</code>	Furhter arguments to be passed to plot()

### Value

A ggplot or no return value, called for side effects

### Examples

```

require(survival)
data(cancer,package="survival")
fit=coxph(Surv(time,status)~x,data=leukemia)
loglogplot(fit)
fit=survfit(Surv(time,status)~1,data=anderson)
loglogplot(fit)
fit=survfit(Surv(time,status)~sex,data=anderson)
loglogplot(fit)
fit=survfit(Surv(time,status)~logWBC,data=anderson)
loglogplot(fit)
fit=survfit(Surv(time,status)~logWBC+rx,data=anderson)
loglogplot(fit,no=2)
fit=survfit(Surv(time,status)~rx,data=anderson)
loglogplot(fit,type="p")
fit=survfit(Surv(time,status)~WBCCAT,data=anderson2)
loglogplot(fit,type="p",what="survOdds")
loglogplot(fit,type="p",what="failureOdds")

```

`maxnchar`*Return maximum character number except NA*

---

**Description**

Return maximum character number except NA

**Usage**

```
maxnchar(x)
```

**Arguments**

`x` a vector

**Value**

A numeric vector of length 1

**Examples**

```
x=c(1,2,"sadf",NA)
maxnchar(x)
data(acs,package="moonBook")
lapply(acs,maxnchar)
```

---

`modelPlot`*Draw coefficients/odds ratio/hazard ratio plot*

---

**Description**

Draw coefficients/odds ratio/hazard ratio plot

**Usage**

```
modelPlot(
  fit,
  widths = NULL,
  change.pointsize = TRUE,
  show.OR = TRUE,
  show.ref = TRUE,
  bw = TRUE,
  legend.position = "top",
  ...
)
```

**Arguments**

<code>fit</code>	An object of class <code>glm</code>
<code>widths</code>	Numeric vector
<code>change.pointsize</code>	logical Whether or not change point size
<code>show.OR</code>	logical Whether or not show odds ratio
<code>show.ref</code>	logical Whether or not show reference
<code>bw</code>	logical If true, use grey scale
<code>legend.position</code>	legend position default value is 'top'
<code>...</code>	Further arguments to be passed to <code>autoReg()</code>

**Value**

`modelPlot` returns an object of class "modelPlot" An object of class `modelPlot` is a list containing at least of the following components:

**tab1** The first table containing names

**tab2** The 2nd table containing levels

**tab3** The 3rd table containing coefficients or odds ratio or hazards ratio

**p** A `ggplot`

**widths** the widths of the tables and the `ggplot`

**Examples**

```
fit=lm(mpg~wt*hp+am,data=mtcars)
modelPlot(fit,widths=c(1,0,2,3))
modelPlot(fit,uni=TRUE,threshold=1,widths=c(1,0,2,3))
fit=lm(Sepal.Width~Sepal.Length*Species,data=iris)
modelPlot(fit)
modelPlot(fit,uni=TRUE,change.pointsize=FALSE)

data(cancer,package="survival")
fit=glm(status~rx+age+sex+nodes+obstruct+perfor,data=colon,family="binomial")
modelPlot(fit)
modelPlot(fit,uni=TRUE,multi=TRUE,threshold=1)
modelPlot(fit,multi=TRUE,imputed=TRUE,change.pointsize=FALSE)
data(colon_s,package="finalfit")
fit=glm(mort_5yr~age.factor+sex.factor+obstruct.factor+perfor.factor,data=colon_s,family="binomial")
modelPlot(fit)
modelPlot(fit,uni=TRUE,multi=TRUE,threshold=1)
modelPlot(fit,uni=TRUE,multi=TRUE)
modelPlot(fit,uni=TRUE,multi=TRUE,threshold=1,show.ref=FALSE)
library(survival)
fit=coxph(Surv(time,status)~rx+age+sex+obstruct+perfor,data=colon)
modelPlot(fit)
modelPlot(fit,uni=TRUE,threshold=1)
modelPlot(fit,multi=FALSE,final=TRUE,threshold=1)
```

```
fit=coxph(Surv(time,status)~age.factor+sex.factor+obstruct.factor+perfor.factor,data=colon_s)
modelPlot(fit)
modelPlot(fit,uni=TRUE,threshold=1)
modelPlot(fit,uni=TRUE,threshold=1,show.ref=FALSE)
modelPlot(fit,imputed=TRUE)
```

---

modelsSummary	<i>Makes table summarizing list of models</i>
---------------	---

---

### Description

Makes table summarizing list of models

### Usage

```
modelsSummary(fitlist, show.lik = FALSE)
```

### Arguments

fitlist	A list of objects of class "coxph"
show.lik	logical Whether or not show likelihood test results

### Value

No return value, called for side effects

### Examples

```
library(survival)
fit1=coxph(Surv(time,status) ~rx,data=anderson)
fit2=coxph(Surv(time,status) ~rx+logWBC,data=anderson)
fit3=coxph(Surv(time,status) ~rx*logWBC,data=anderson)
fitlist=list(fit1,fit2,fit3)
modelsSummary(fitlist)
```

---

modelsSummaryTable	<i>Makes flextable summarizing list of models</i>
--------------------	---

---

### Description

Makes flextable summarizing list of models

### Usage

```
modelsSummaryTable(fitlist, labels = NULL, show.lik = FALSE)
```

**Arguments**

fitlist	A list of objects of class "coxph"
labels	character labels of models
show.lik	logical Whether or not show likelihood test results

**Value**

A flextable

**Examples**

```
library(survival)
fit1=coxph(Surv(time,status) ~rx,data=anderson)
fit2=coxph(Surv(time,status) ~rx+logWBC,data=anderson)
fit3=coxph(Surv(time,status) ~rx*logWBC,data=anderson)
fitlist=list(fit1,fit2,fit3)
modelsSummaryTable(fitlist)
```

---

my.chisq.test2	<i>Statistical test for categorical variables Statistical test for categorical variables</i>
----------------	--

---

**Description**

Statistical test for categorical variables Statistical test for categorical variables

**Usage**

```
my.chisq.test2(x, y, catMethod = 2, all = FALSE)
```

**Arguments**

x	a vector
y	a vector
catMethod	An integer indicating methods for categorical variables. Possible values in methods are <ul style="list-style-type: none"> <li><b>0</b> Perform chisq.test first. If warning present, perform fisher test</li> <li><b>1</b> Perform chisq.test without continuity correction</li> <li><b>2</b> Perform chisq.test with continuity correction</li> <li><b>3</b> perform fisher.test</li> <li><b>4</b> perform prop.trend test</li> </ul> Default value is 2.
all	A logical

**Value**

A numeric vector of length 1

**Examples**

```
library(moonBook)
x=acs$sex
y=acs$Dx
my.chisq.test2(x,y)
```

---

my.t.test2

*Statistical test for continuous variables*

---

**Description**

Statistical test for continuous variables

**Usage**

```
my.t.test2(y, x, method = 1, all = FALSE)
```

**Arguments**

y	a categorical vector
x	a numeric vector
method	method An integer indicating methods for continuous variables. Possible values in methods are 1 forces analysis as normal-distributed 2 forces analysis as continuous non-normal 3 performs a Shapiro-Wilk test or nortest::ad.test to decide between normal or non-normal Default value is 1.
all	A logical

**Value**

A numeric vector of length 1

**Examples**

```
library(moonBook)
y=acs$sex
x=acs$height
my.t.test2(y,x)
```



---

mycphSimple

*Fit Simple Proportional Hazards Regression Model*


---

**Description**

Fit Simple Proportional Hazards Regression Model

**Usage**

```
mycphSimple(fit, threshold = 0.2, digits = 2)
```

**Arguments**

fit	An object of class coxph
threshold	numeric p-value threshold to enter multiple model
digits	integer indicating the position decimal place

**Value**

An object of class "data.frame"

**Examples**

```
require(survival)
data(cancer)
fit=coxph(Surv(time,status)~age+sex+obstruct+perfor,data=colon)
mycphSimple(fit)
```

---

myformat

*Convert data.frame to printable format*


---

**Description**

Convert data.frame to printable format

**Usage**

```
myformat(x, showid = FALSE, digits = 3)
```

**Arguments**

x	A data.frame
showid	logical if TRUE, show id
digits	Integer indicating the number of decimal places

**Value**

A data.frame

**Examples**

```
fit=lm(mpg~wt*hp,data=mtcars)
gaze(fit) %>% myformat()
```

---

myft

*Convert data.frame into flextable*

---

**Description**

Convert data.frame into flextable

**Usage**

```
myft(x, vanilla = TRUE, fontsize = 10, digits, showid = FALSE, ...)
```

**Arguments**

x	A data.frame
vanilla	logical
fontsize	Numeric
digits	integer indicating the position of decimal place
showid	logical if TRUE, show id
...	Further arguments to be passed to df2flextable()

**Value**

An object of class `flextable`

**Examples**

```
data(acs,package="moonBook")
library(dplyr)
gaze(acs) %>% myft()
gaze(sex~.,acs) %>% myft()
fit=lm(mpg~hp*wt,data=mtcars)
gaze(fit) %>% myft()
library(survival)
fit=coxph(Surv(time,status) ~rx,data=anderson1)
gaze(fit) %>% myft()

gaze(sex+Dx~.,data=acs,show.p=TRUE,show.total=TRUE,show.n=TRUE,shiw.missing=TRUE) %>% myft()
gaze(Dx+sex~cardiogenicShock,data=acs,show.p=TRUE) %>% myft()
gaze(Dx+sex+HBP~cardiogenicShock,data=acs,show.p=TRUE) %>% myft()
```

---

mysurvregSimple	<i>Fit Simple AFT Model</i>
-----------------	-----------------------------

---

**Description**

Fit Simple AFT Model

**Usage**

```
mysurvregSimple(fit, threshold = 0.2, digits = 2, mode = 1)
```

**Arguments**

fit	An object of class survreg
threshold	numeric p-value threshold to enter multiple model
digits	integer indicating the position decimal place
mode	integer

**Value**

An object of class "data.frame"

**Examples**

```
require(survival)
data(cancer)
fit=survreg(Surv(time,status)~rx+age+strata(sex)+obstruct+perfor,data=colon)
mysurvregSimple(fit)
```

---

num2factor	<i>Convert a numeric column in a data.frame to a factor</i>
------------	---

---

**Description**

Convert a numeric column in a data.frame to a factor

**Usage**

```
num2factor(data, call, name, no = 3)
```

**Arguments**

data	A data.frame
call	a function call
name	character Name of numeric column
no	numeric

**Value**

A data.frame

**Examples**

```
num2factor(anderson,name="logWBC")
library(survival)
fit=coxph(Surv(time,status)~logWBC+rx,data=anderson)
num2factor(anderson,call=fit$call,name="logWBC",no=2)
```

---

num2stat

*Summarize numeric vector to statistical summary*

---

**Description**

Summarize numeric vector to statistical summary

**Usage**

```
num2stat(x, digits = 1, method = 1, p = NULL)
```

**Arguments**

x	A numeric vector
digits	integer indicating the number of decimal places
method	An integer indicating methods for continuous variables. Possible values in methods are 1 forces analysis as normal-distributed 2 forces analysis as continuous non-normal 3 performs a Shapiro-Wilk test or nortest::ad.test to decide between normal or non-normal Default value is 1.
p	A numeric

**Value**

A character vector of length 1

**Examples**

```
library(moonBook)
num2stat(acs$age)
num2stat(acs$age,method=2)
```

---

OEplot *Draw an Observed vs Expected plot*

---

**Description**

Draw an Observed vs Expected plot

**Usage**

```
OEplot(fit, xnames = NULL, no = 3, maxy.lev = 5, median = TRUE)
```

**Arguments**

fit	An object of class "coxph"
xnames	Character Names of explanatory variable to plot
no	integer Number of groups to be made
maxy.lev	Integer Maximum unique length of a numeric variable to be treated as categorical variables
median	logical

**Value**

No return value, called for side effects

**Examples**

```
library(survival)
data(cancer, package="survival")
fit=coxph(Surv(time, status)~rx+age+sex, data=colon)
OEplot(fit)
OEplot(fit, xnames="sex")
## Not run:
fit=coxph(Surv(time, status)~age, data=colon)
OEplot(fit)
fit=coxph(Surv(time, status)~logWBC, data=anderson)
OEplot(fit)

## End(Not run)
```

---

p2character2	<i>Change p value to string</i>
--------------	---------------------------------

---

**Description**

Change p value to string

**Usage**

```
p2character2(x, digits = 3, add.p = TRUE)
```

**Arguments**

x	a numeric
digits	integer indicating decimal place
add.p	logical

**Value**

A character vector

---

print.autoReg	<i>S3 method print for an object of class autoReg</i>
---------------	---

---

**Description**

S3 method print for an object of class autoReg

**Usage**

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

**Arguments**

x	An object of class autoReg
...	Further arguments

**Value**

No return value, called for side effects

**Examples**

```
data(cancer, package="survival")
fit=glm(status~rx+sex+age+obstruct+nodes, data=colon, family="binomial")
autoReg(fit)
```

---

print.gaze	<i>S3 method print for an object of class gaze</i>
------------	--

---

**Description**

S3 method print for an object of class gaze

**Usage**

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

**Arguments**

x	An object of class gaze
...	Further arguments

**Value**

No return value, called for side effects

**Examples**

```
data(acs, package="moonBook")  
x=gaze(acs, show.n=TRUE, show.missing=TRUE)  
gaze(sex~, acs, show.p=TRUE, show.n=TRUE, show.missing=TRUE, show.total=TRUE)  
  
gaze(Dx+sex~, acs, show.p=TRUE)  
gaze(sex+Dx+HBP~, acs, show.p=TRUE)
```

---

print.modelPlot	<i>S3 method for an class modelPlot</i>
-----------------	---

---

**Description**

S3 method for an class modelPlot

**Usage**

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

**Arguments**

x	An object of class modelPlot
...	Further arguments to be passed to plot()

printdf *Print function for data.frame*

---

**Description**

Print function for data.frame

**Usage**

```
printdf(x)
```

**Arguments**

x                    A data.frame

**Value**

No return value, called for side effects

**Examples**

```
x=mtcars[1:5,1:5]
printdf(x)
```

---

removeDup *Remove duplicated term*

---

**Description**

Remove duplicated term

**Usage**

```
removeDup(x, replacement = "")
```

**Arguments**

x                    A vector  
replacement        A character to be replaced or NA

**Value**

A vector with the same class as x

**Examples**

```
x=rep(1:5,each=3)
removeDup(x)
```



---

residualNull	<i>Make a residual plot of NULL model</i>
--------------	---

---

**Description**

Make a residual plot of NULL model

**Usage**

```
residualNull(x, add.log = TRUE, type = "martingale")
```

**Arguments**

x	An object of class coxph
add.log	logical If true, log of predictor variables are added
type	character type of residuals

**Examples**

```
library(survival)
data(pharmacoSmoking, package="asaur")
pharmacoSmoking$priorAttemptsT=pharmacoSmoking$priorAttempts
pharmacoSmoking$priorAttemptsT[pharmacoSmoking$priorAttemptsT>20]=20
x=coxph(Surv(ttr, relapse)~age+priorAttemptsT+longestNoSmoke, data=pharmacoSmoking)
residualNull(x)
```

---

residualPlot	<i>Draw a residual plot with an object of class coxph</i>
--------------	---

---

**Description**

Draw a residual plot with an object of class coxph

**Usage**

```
residualPlot(
  fit,
  type = "martingale",
  vars = NULL,
  ncol = 2,
  show.point = TRUE,
  se = TRUE,
  topn = 5,
  labelsize = 4
)
```

**Arguments**

<code>fit</code>	An object of class <code>coxph</code> or <code>survreg</code>
<code>type</code>	character One of the <code>c("martingale", "deviance", "score", "schoenfeld", "dfbeta", "dfbetas", "scaledsch", "partial")</code> . Default value is "martingale".
<code>vars</code>	character Names of variables to plot. default value is <code>NULL</code>
<code>ncol</code>	numeric number of columns
<code>show.point</code>	logical Whether or not show point
<code>se</code>	logical Whether or not show se
<code>topn</code>	numeric number of data to be labelled
<code>labelsize</code>	numeric size of label

**Value**

A patchwork object

**Examples**

```

require(survival)
data(cancer)
fit=coxph(Surv(time, status==2)~log(bili)+age+cluster(edema), data=dbc)
residualPlot(fit)
residualPlot(fit, vars="age")
fit=coxph(Surv(time, status==2)~age, data=dbc)
residualPlot(fit)
residualPlot(fit, "partial")
fit=coxph(Surv(time, status)~rx+sex+logWBC, data=anderson)
residualPlot(fit, ncol=3)
## Not run:
data(pharmacoSmoking, package="asaur")
fit=coxph(Surv(ttr, relapse)~grp+employment+age, data=pharmacoSmoking)
residualPlot(fit)
residualPlot(fit, var="age")
residualPlot(fit, type="dfbeta")
residualPlot(fit, type="dfbeta", var="age")
residualPlot(fit, type="dfbeta", var="employment")
residualPlot(fit, type="dfbeta", var="employmentother")
pharmacoSmoking$ttr[pharmacoSmoking$ttr==0]=0.5
fit=survreg(Surv(ttr, relapse)~grp+age+employment, data=pharmacoSmoking, dist="weibull")
residualPlot(fit, type="response")
residualPlot(fit, type="deviance")
residualPlot(fit, type="dfbeta", vars="age")
fit=survreg(Surv(time, status)~ph.ecog+sex*age, data=lung, dist="weibull")
residualPlot(fit, "dfbeta")
residualPlot(fit, "deviance")

## End(Not run)

```

---

restoreData	<i>restore data with factor in column name</i>
-------------	--

---

**Description**

restore data with factor in column name

**Usage**

```
restoreData(data)
```

**Arguments**

data	An object of class "data.frame"
------	---------------------------------

**Value**

An object of class "data.frame"

---

restoreData2	<i>restore data with I() in column name</i>
--------------	---

---

**Description**

restore data with I() in column name

**Usage**

```
restoreData2(df)
```

**Arguments**

df	An object of class "data.frame"
----	---------------------------------

**Value**

An object of class "data.frame"

restoreData3            *restore data with operator in column name*

---

**Description**

restore data with operator in column name

**Usage**

```
restoreData3(df, changeLabel = FALSE)
```

**Arguments**

df                    An object of class "data.frame"  
changeLabel        logical

**Value**

An object of class "data.frame"

---

revOperator            *get opposite arithmetic operator*

---

**Description**

get opposite arithmetic operator

**Usage**

```
revOperator(operator)
```

**Arguments**

operator            A character

**Value**

A character

---

roundDf	<i>Convert numeric columns of data.frame to character</i>
---------	---

---

**Description**

Convert numeric columns of data.frame to character

**Usage**

```
roundDf(df, digits = 2)
```

**Arguments**

df	a data.frame
digits	integer indicating the number of decimal places

**Value**

An object of class "data.frame"

---

setLabel	<i>Add label to a vector</i>
----------	------------------------------

---

**Description**

Add label to a vector

**Usage**

```
setLabel(x, label = "")
```

**Arguments**

x	a vector
label	string

**Value**

a labelled vector

---

shorten	<i>Shorten an object of class gaze</i>
---------	--

---

**Description**

Shorten an object of class gaze

**Usage**

```
shorten(x, xname = NULL, ref = 1)
```

**Arguments**

x	an object of class gaze
xname	A variable name
ref	Numeric Th number to be used as reference

**Value**

An object of class "gaze" which is described in [gaze](#)

**Examples**

```
data(acs, package="moonBook")
x=gaze(sex~., data=acs)
shorten(x)
```

---

showEffect	<i>Show effects of covariates</i>
------------	-----------------------------------

---

**Description**

Show effects of covariates

**Usage**

```
showEffect(
  fit,
  x = NULL,
  color = NULL,
  facet = NULL,
  pred.values = list(),
  se = TRUE,
  logy = TRUE,
  collabel = label_both,
  rowlabel = label_both
)
```

**Arguments**

fit	An object of class survreg
x	character name of x-axis variable
color	character name of color variable
facet	character name of facet variable
pred.values	list list of values of predictor variables
se	logical whether or not show se
logy	logical Whether or not draw y-axis on log scale
collabel	labeller for column
rowlabel	labeller for row

**Value**

A ggplot

**Examples**

```
library(survival)
library(ggplot2)
fit=survreg(Surv(time,status)~ph.ecog+sex*age,data=lung,dist="weibull")
showEffect(fit)
fit=survreg(Surv(time,status)~rx+sex+age+obstruct+adhere,data=colon,dist="weibull")
showEffect(fit)
showEffect(fit,rowlabel=label_value)
fit=survreg(Surv(time,status)~ph.ecog+sex,data=lung,dist="weibull")
showEffect(fit)
fit=survreg(Surv(time,status)~ph.ecog+age,data=lung,dist="weibull")
showEffect(fit)
fit=survreg(Surv(time,status)~sex*age,data=lung,dist="weibull")
showEffect(fit)
fit=survreg(Surv(time,status)~age,data=lung,dist="weibull")
showEffect(fit)
```

---

strata2df

---

*Convert a character vector to a data.frame*


---

**Description**

Convert a character vector to a data.frame

**Usage**

```
strata2df(strata)
```

**Arguments**

strata	A character vector
--------	--------------------

**Value**

A data.frame

---

survfit2df	<i>Extract survival data from an object of class "survfit"</i>
------------	--

---

**Description**

Extract survival data from an object of class "survfit"

**Usage**

```
survfit2df(fit, labels = NULL)
```

**Arguments**

fit	An object of class "survfit"
labels	Character

**Value**

A data.frame

**Examples**

```
library(survival)
data(cancer, package="survival")
fit=survfit(coxph(Surv(time, status)~sex+age+strata(rx), data=colon))
survfit2df(fit)
## Not run:
fit=coxph(Surv(time, status)~sex+age+strata(rx), data=colon)
fit=survfit(as.formula(deparse(fit$terms)), data=fit2model(fit))
survfit2df(fit)
fit=survfit(Surv(time, status)~rx+sex+age, data=colon)
survfit2df(fit)
fit=survfit(Surv(time, status)~1, data=colon)
survfit2df(fit)

## End(Not run)
```



---

survreg2final	<i>Make final model using stepwise backward elimination</i>
---------------	---

---

**Description**

Make final model using stepwise backward elimination

**Usage**

```
survreg2final(fit, threshold = 0.2)
```

**Arguments**

fit	An object of class "survreg"
threshold	Numeric

**Value**

An object of class "survreg" which is described in [survreg](#)

**Examples**

```
require(survival)
data(cancer)
fit=survreg(Surv(time,status)~rx+age+sex+obstruct+perfor,data=colon)
survreg2final(fit)
```

---

survreg2multi	<i>Make multivariable regression model by selecting univariable models with p.value below threshold</i>
---------------	---

---

**Description**

Make multivariable regression model by selecting univariable models with p.value below threshold

**Usage**

```
survreg2multi(fit, threshold = 0.2)
```

**Arguments**

fit	An object of class "survreg"
threshold	Numeric

**Value**

An object of class "survreg"

**Examples**

```
require(survival)
data(cancer)
fit=survreg(Surv(time,status)~rx+age+sex+obstruct+perfor,data=colon)
survreg2multi(fit)
```

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