

# Package ‘SurvivalTests’

July 21, 2025

**Type** Package

**Title** Survival Tests for One-Way Layout

**Version** 1.0

**Date** 2024-01-16

**Depends** R (>= 3.2.0), weibullness

**Imports** stats, ggplot2

**Suggests** survival, arules

**Author** Osman Dag [aut, cre],  
Malwane Ananda [aut],  
Sam Weerahandi [aut]

**Maintainer** Osman Dag <osman.dag@outlook.com>

**Description** Performs survival analysis for one-way layout. The package includes the generalized test for survival ANOVA (Tsui and Weerahandi (1989) <doi:10.2307/2289949> and (Weerahandi, 2004; ISBN:978-0471470175)). It also performs pairwise comparisons and graphical approaches. Moreover, it assesses the weibullness of data in each group via test. The package computes mean and confidence interval under Weibull distribution.

**License** GPL (>= 2)

**NeedsCompilation** no

**Repository** CRAN

**Date/Publication** 2024-01-16 17:50:05 UTC

## Contents

plot.Sdescribe . . . . .	2
SANOVA . . . . .	3
Sdescribe . . . . .	4
Spaircomp . . . . .	5
weibull.test . . . . .	6

<b>Index</b>	<b>8</b>
--------------	----------

---

`plot.Sdescribe`*Error Bars with Means and Confidence Limits*

---

**Description**

`plot.Sdescribe` produce error bars with means and confidence limits of the given grouped values.

**Usage**

```
## S3 method for class 'Sdescribe'  
plot(x, ylim = NULL, xlab = NULL, ylab = NULL, title = NULL, width = NULL, ...)
```

**Arguments**

<code>x</code>	a <code>Sdescribe</code> object.
<code>ylim</code>	a limit for y axes.
<code>xlab</code>	a label for the x axis.
<code>ylab</code>	a label for the y axis.
<code>title</code>	a main title for the plot.
<code>width</code>	the little lines at the tops and bottoms of the error bars (defaults to 0.80).
<code>...</code>	additional arguments.

**Value**

No return value, called for side effects

**Author(s)**

Osman Dag

**Examples**

```
library(survival)  
lung$status <- ifelse(lung$status == 2, 1, 0)  
lung$age <- arules::discretize(lung$age, breaks = 3, labels = c("Low", "Medium", "High"))  
  
library(SurvivalTests)  
  
out <- Sdescribe(time~age+status, lung)  
  
plot(out, ylim = c(0,500))  
plot(out)
```

---

 SANOVA

*Generalized Test for Survival ANOVA*


---

**Description**

SANOVA performs generalized test for survival ANOVA.

**Usage**

```
SANOVA(formula, data, nM = 5000, seed = 123, alpha = 0.05, na.rm = TRUE, verbose = TRUE)
```

**Arguments**

formula	a formula of the form lhs ~ rhs where lhs gives the survival times and rhs the corresponding groups + the status variable (1: non-censored, 0: censored).
data	a data frame containing the variables in formula.
nM	a number of bootstrap samples.
seed	a seed number for the reproducibility of results. Default is set to 123.
alpha	the level of significance to assess the statistical difference. Default is set to alpha = 0.05.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
verbose	a logical for printing output to R console.

**Details**

SANOVA performs parametric survival ANOVA when the underlying data are distributed as Weibull or Gumbel. SANOVA tests are based on generalized p-value approach (cf. Tsui and Weerahandi (1989) and Weerahandi (2004)) extended for situations of affine invariance.

**Value**

A list with class "survtests" containing the following components:

p.value	the p-value of the test.
alpha	the level of significance to assess the statistical difference.
method	the character string "Generalized Test for Survival ANOVA".
data	a data frame containing the variables in which NA values (if exist) are removed.
formula	a formula of the form lhs ~ rhs where lhs gives the sample values and rhs the corresponding groups + the status variable (1: non-censored, 0: censored).
seed	a seed number for the reproducibility of results.

**Author(s)**

Sam Weerahandi, Malwane Ananda, Osman Dag

## References

Tsui K. and Weerahandi, S. (1989). Generalized P-values in Significance Testing of Hypotheses in the Presence of Nuisance Parameters. *Journal of the American Statistical Association*, **84**, 602-607.

Weerahandi, S. (2004). *Generalized Inference in Repeated Measures: Exact Methods in MANOVA and Mixed Models*, Wiley.

## Examples

```
library(survival)
lung$status <- ifelse(lung$status == 2, 1, 0)
lung$age <- arules::discretize(lung$age, breaks = 3, labels = c("Low", "Medium", "High"))

library(SurvivalTests)

SANOVA(time~age+status, lung, alpha = 0.05)
```

---

Sdescribe

*Descriptive Statistics*

---

## Description

Sdescribe produces number of observations (n), number of events (nE), and the mean survival with confidence limits for the mean.

## Usage

```
Sdescribe(formula, data, level = 0.95, nM = 5000, na.rm = TRUE,
  verbose = TRUE)
```

## Arguments

formula	a formula of the form lhs ~ rhs where lhs gives the survival times and rhs the corresponding groups + the status variable (1: non-censored, 0: censored).
data	a data frame containing the variables in formula.
level	the confidence level.
nM	a number of bootstrap samples.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
verbose	a logical for printing output to R console.

## Value

Returns a data.frame of output with class "Sdescribe".

**Author(s)**

Malwane Ananda, Osman Dag

**Examples**

```
library(survival)
lung$status <- ifelse(lung$status == 2, 1, 0)
lung$age <- arules::discretize(lung$age, breaks = 3, labels = c("Low", "Medium", "High"))

library(SurvivalTests)

Sdescribe(time~age+status, lung)
```

---

Spaircomp

*Pairwise Comparisons of Survival Tests for One-Way Layout*

---

**Description**

Spaircomp is a generic function for pairwise comparisons of survival tests in one-way layout by adjusting p-values.

**Usage**

```
## S3 method for class 'survtests'
Spaircomp(x, adjust.method = c("bonferroni", "holm", "hochberg", "hommel", "BH",
  "BY", "fdr", "none"), verbose = TRUE, ...)
```

**Arguments**

x	a survtests object.
adjust.method	Method for adjusting p values (see <a href="#">p.adjust</a> ). Default is set to "bonferroni".
verbose	a logical for printing output to R console.
...	Additional arguments affecting multiple comparisons of groups in one-way independent designs.

**Value**

Returns a data.frame of output.

**Author(s)**

Osman Dag

**Examples**

```

library(survival)
lung$status <- ifelse(lung$status == 2, 1, 0)
lung$age <- arules::discretize(lung$age, breaks = 3, labels = c("Low", "Medium", "High"))

library(SurvivalTests)

out = SANOVA(time~age+status, lung, alpha = 0.05)

Spaircomp(out, adjust.method = "bonferroni")

# to illustrate pairwise comparison, alpha is set to 0.5
out2 = SANOVA(time~age+status, lung, alpha = 0.5)

Spaircomp(out2, adjust.method = "bonferroni")

Spaircomp(out2, adjust.method = "hochberg")

Spaircomp(out2, adjust.method = "hommel")

Spaircomp(out2, adjust.method = "holm")

```

---

weibull.test

*Weibullness Test*


---

**Description**

weibull.test performs weibullness test from weibull plot in each group.

**Usage**

```
weibull.test(formula, data, alpha = 0.05, na.rm = TRUE, verbose = TRUE)
```

**Arguments**

formula	a formula of the form lhs ~ rhs where lhs gives the survival times and rhs the corresponding groups.
data	a tibble or data frame containing the variables in formula.
alpha	the level of significance to assess normality. Default is set to alpha = 0.05.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
verbose	a logical for printing output to R console.

**Value**

A data frame gives the test results for the weibullness of groups.

**Author(s)**

Osman Dag

**See Also**

[wp.test](#)

**Examples**

```
library(survival)
lung$status <- ifelse(lung$status == 2, 1, 0)
lung$age <- arules::discretize(lung$age, breaks = 3, labels = c("Low", "Medium", "High"))

library(SurvivalTests)

weibull.test(time~age, lung)
```

# Index

## \* functions

plot.Sdescribe, 2

SANOVA, 3

Sdescribe, 4

Spaircomp, 5

weibull.test, 6

p.adjust, 5

plot.Sdescribe, 2

SANOVA, 3

Sdescribe, 4

Spaircomp, 5

weibull.test, 6

wp.test, 7