Package 'sanon'

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Title Stratified Analysis with Nonparametric Covariable Adjustment
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Depends R (>= 3.5)
Description There are several functions to implement the method for analysis in a randomized clinical trial with strata with following key features. A stratified Mann-Whitney estimator addresses the comparison between two randomized groups for a strictly ordinal response variable. The multivariate vector of such stratified Mann-Whitney estimators for multivariate re-

cal trial with strata with following key features. A stratified Mann-Whitney estimator addresses the comparison between two randomized groups for a strictly ordinal response variable. The multivariate vector of such stratified Mann-Whitney estimators for multivariate response variables can be considered for one or more response variables such as in repeated measurements and these can have missing completely at random (MCAR) data. Non-parametric covariance adjustment is also considered with the minimal assumption of randomization. The p-value for hypothesis test and confidence interval are provided.

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R topics documented:

Type Package

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Description

A Package for Implementation of the method in Kawaguchi, Koch, and Wang (2011)

Author(s)

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References

Kawaguchi A., Koch, G. G. (2015). sanon: An R Package for Stratified Analysis with Nonparametric Covariable Adjustment. Journal of Statistical Software, 67(9), 1-37. doi:10.18637/jss.v067.i09

Kawaguchi A., Koch, G. G., Wang, X. (2011). Stratified Multivariate Mann-Whitney Estimators for the Comparison of Two Treatments with Randomization Based Covariance Adjustment. Statistics in Biopharmaceutical Research, Vol. 3, No. 2, 217-231.

See Also

sanon

catecovar

Identify Categorical Covariables

Description

This is a special function used in the context of sanon. It identifies categorical covariables when they appear on the right hand side of a formula.

Usage

```
catecovar(x, ref = NULL)
```

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Arguments

x variable name

ref character for the reference group for the categorical covariable.

Details

In the sanon, the categorical covariable is converted into a dummy variable. The reference group is specified in the ref argument.

coef.sanon

Extract Model Coefficients

Description

coef is a generic function which extracts model coefficients from objects returned by modeling functions, coefficients is an alias for it.

Usage

```
## S3 method for class 'sanon'
coef(object, ...)
```

Arguments

object an object of class "sanon", usually, a result of a call to sanon further arguments passed to or from other methods.

Details

All object classes which are returned by model fitting functions should provide a coef method or use the default one.

Value

Coefficients extracted from the model object object.

Examples

```
##### Example 3.1 Randomized Clinical Trial of Chronic Pain #####
data(cpain)
out1 = sanon(response ~ grp(treat, ref="placebo") + strt(center) + strt(diagnosis), data=cpain)
coef(out1)
coefficients(out1)

##### Example 3.2 Randomized Clinical Trial of Respiratory Disorder ####
data(resp)
P = rbind(rep(0, 4), diag(4), rep(0, 4))
out23 = sanon(cbind(baseline, visit1, visit2, visit3, visit4) ~ grp(treatment, ref="P")
```

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```
+ strt(center) + strt(sex) + covar(age), data=resp, P=P)
# each four visits
coef(out23)
coefficients(out23)
```

confint.sanon

Confidence Intervals for Model Parameters

Description

Computes confidence intervals for one or more parameters in a fitted model.

Usage

```
## S3 method for class 'sanon'
confint(object, parm = NULL, level = 0.95, ...)
## S3 method for class 'confint.sanon'
print(x, ...)
```

Arguments

object, x an object of class "sanon", usually, a result of a call to sanon

parm a specification of which parameters are to be given confidence intervals, either

a vector of numbers or a vector of names. If missing, all parameters are consid-

ered.

level the confidence level required.

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

Details

Confidence intervals for adjusted parameters in the weighted least squares are computed based on an asymptotic normal.

Value

ci A matrix (or vector) with columns giving Mann-Whiteney estimates and their

lower and upper confidence limits for each parameter with estimates. The interval will be labelled as Lower for (1 - level)/2 limit and Upper for 1 - (1 - level)/2

limit (by default 0.025 and 0.975).

level Confidence level

advarnames Adjust variable names in the weighted least squares method

contrast 5

Examples

```
##### Example 3.1 Randomized Clinical Trial of Chronic Pain #####
data(cpain)
out1 = sanon(response ~ grp(treat, ref="placebo") + strt(center) + strt(diagnosis), data=cpain)
confint(out1)

##### Example 3.2 Randomized Clinical Trial of Respiratory Disorder ####
data(resp)
P = rbind(rep(0, 4), diag(4), rep(0, 4))
out23 = sanon(cbind(baseline, visit1, visit2, visit3, visit4) ~ grp(treatment, ref="P")
+ strt(center) + strt(sex) + covar(age), data=resp, P=P)
# each four visits
confint(out23)
```

contrast

Contrast for Model Parameters

Description

Inference by contrast of parameters in a fitted model.

Usage

```
contrast(
  object,
  C = diag(length(object$b)),
  confint = FALSE,
  level = 0.95,
   ...
)

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

Arguments

object, x an object of class "sanon", usually, a result of a call to sanon.

C contrast matrix. The number of column should be same as the length of b in outputs of sanon.

confint logical value for whether the confidence interval is computed (only if C has one row).

level the confidence level required (only if C has one row).

further arguments passed to or from other methods.

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Details

This function provide the inference based on contrast after applying the function sanon. The contrast matrix C should be defined by the user. If the number of row of C = 1, the confidence interval for the estimator is produced.

Value

С	contrast matrix
Cb	contrast estimates
VCb	variance and covariance matrix of Cb
se	standard error of Cb
level	confidence level
UL	upper confidence limit (only if the number of row of C = 1, otherwise NULL)
LL	lower confidence limit (only if the number of row of $C = 1$, otherwise NULL)
Q	test statistic
df	degree of freedom
p	p-value

Examples

```
##### Example 3.2 Randomized Clinical Trial of Respiratory Disorder #####
data(resp)
P = rbind(rep(0, 4), diag(4), rep(0, 4))
out23 = sanon(cbind(baseline, visit1, visit2, visit3, visit4) ~ grp(treatment, ref="P")
+ strt(center) + strt(sex) + covar(age), data=resp, P=P)

# Homogeneity of the xi_k across the four visits
contrast(out23, C=cbind(diag(3), rep(-1, 3)))

# Comparison between treatments for the average of the xi_k across the 4 visits
contrast(out23, C=matrix(rep(1, 4)/4, ncol=4))
```

covar

Identify Covariables

Description

This is a special function used in the context of sanon. It identifies covariables when they appear on the right hand side of a formula.

Usage

```
covar(x)
```

Arguments

x variable name

cpain 7

cpain

Chronic Pain Data

Description

The data are from a multicenter randomized clinical trial to compare test and control treatments for the management of chronic pain, and they have had previous consideration in Stokes et al. (2000, chap. 13).

Usage

```
data(cpain)
```

Format

A data frame with 193 observations and 4 variables

Details

treat a factor with levels active and placebo for treatment

response a factor with five levels poor, fair, moderate, good and excel for pain status after treatment for 4 weeks

center a factor with two levels I and II for two centers

diagnosis a factor with four levels A, B, C, and D for diagnoses

References

Stokes, M. E., Davis, C. S., and Koch, G. G. (2000), Categorical Data Analysis using the SAS System, Cary: SAS Publishing.

grp

Identify Group Variables

Description

This is a special function used in the context of sanon. It identifies group variables when they appear on the right hand side of a formula.

Usage

```
grp(x, ref = NULL)
```

Arguments

x variable name

ref character for the reference group for treatment group.

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heartburn

Relief of heartburn Data

Description

The data are from two period cross-over design clinical trial for relief of heartburn, and listings of the data appear in Koch, Gitomer, Skalland, and Stokes (1983). The variables are as follows:

Usage

data(heartburn)

Format

A data frame with 60 observations and 9 variables.

Details

center a factor vector for two centers

sequence a factor with levels AP and PA for sequence groups

age a numeric vector for age

sex a factor for sex with levels female and male

freq a numeric vector for weekly frequency of condition from previous medical history

MD1 a numeric vector for time to relief from first dose during period 1

MD2 a numeric vector for time to relief from first dose during period 2

res1 a factor vector for relief status for period 1 (R = relief from first dose within 15 min, NF = no relief from first dose within 15 min)

ref2 a factor vector for relief status for period 2 with same categories as res1

References

Koch G, Gitomer S, Skalland L, Stokes M (1983). "Some non-parametric and categorical data analyses for a change-over design study and discussion of apparent carry-over effects." Statistics in Medicine, 2(3), 397–412.

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resp

Respiratory Disorder Data

Description

The data are from a randomized clinical trial to compare a test treatment to placebo for a respiratory disorder, and listings of the data appear in Stokes et al. (2000, chap. 15, pp. 495-496) and Koch et al. (1990). The variables are as follows:

Usage

data(resp)

Format

A data frame with 111 observations and 9 variables.

Details

center a factor vector for two centers

treatment a factor with levels A and P for active and placebo treatments, respectively

sex a factor with levels F and M for female and male, respectively

age a numeric vector for age

- baseline a numeric vector for patient global ratings of symptom control according to 5 categories (4 = excellent, 3 = good, 2 = fair, 1 = poor, 0 = terrible) at baseline measurement
- visit1 a numeric vector for patient global ratings of symptom control at visit 1 with same categories as baseline
- visit2 a numeric vector for patient global ratings of symptom control at visit 2 with same categories as baseline
- visit3 a numeric vector for patient global ratings of symptom control at visit 3 with same categories as baseline
- visit4 a numeric vector for patient global ratings of symptom control at visit 4 with same categories as baseline

References

Stokes, M. E., Davis, C. S., and Koch, G. G. (2000), Categorical Data Analysis using the SAS System, Cary: SAS Publishing.

Koch, G. G., Carr, G. J., Amara, I. A., Stokes, M. E., and Uryniak, T. J. (1990), "Categorical Data Analysis," in Statistical Methodology in Pharmaceutical Sciences, ed. D. A. Berry, New York: Marcel Dekker, pp. 291-475.

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sanon

Non-Parametric Covariable Adjustment for Stratified Rank Measures of Association

Description

This is a function for computing a stratified multivariate Mann-Whitney estimator that addresses the comparison between two randomized groups for a strictly ordinal response variable. Response variables may have some missing completely at random (MCAR) values for some patients. Non-parametric covariable adjustment is considered through the difference estimates between mean covariable and the weighted least squares method. Although such estimators can be computed directly as weighted linear combinations of within-stratum Mann-Whitney estimators, consistent estimation of their covariance matrix is done using methods for multivariate U-statistics.

Usage

```
sanon(outcome, ...)
## S3 method for class 'formula'
sanon(formula, data = list(), ...)
## Default S3 method:
sanon(
  outcome,
  group,
  strt = NULL,
  covar = NULL,
  catecovar = NULL,
  ref = NULL,
  covref = NULL,
  P = NULL
  res.na.action = "default",
)
## S3 method for class 'sanon'
print(x, ...)
```

Arguments

outcome	outcome) variables
	further arguments passed to or from other methods.
formula	a formula object, with the response on the left of a \sim operator, and the terms on the right.
data	a data.frame in which to interpret the variables named in the formula.

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group	numeric vector of observations of length n for treatment group. The reference group can be specified in ref.
strt	numeric or factor vector of observations of length n, or a matrix with n rows for strata.
covar	numeric or factor vector of observations of length n, or a matrix with n rows for covariable.
catecovar	numeric or factor vector of observations of length n, or a matrix with n rows for categorical covariable.
ref	character for the reference group for treatment group in group.
covref	$character\ vector\ for\ the\ reference\ group\ for\ categorical\ covariables\ in\ {\tt catecovar}.$
P	a matrix for weighted least squares estimation.
res.na.action	character for setting NA actions. "default", "LOCF1", "LOCF2", "replace", and "remove" are available. default is "default". see the details.
X	an object of class "sanon", usually, a result of a call to sanon

Details

sanon has two specifications for the input, variable and formula based. In the variable based input, one can specify R objects to outcome, group, and strata variables, and covariable. In the formula based input, the formula consists of variable names in a data.frame. The strata and group variables, and covariable are recognized by functions strt, grp, covar, and catecovar. outcome can be contained missing values, which should be coded by NA. Five options for the management of missing values can be specifed in the argument res.na.action; "default" = the method in Kawaguchi et al. (2011), "LOCF1" and "LOCF2" = last observation carried forward with respect to kernels of U-statistics and observed velues, repsectively, "replace" = missing values are managed as tied with all other values in the same stratum, and "remove" = the complete cases analaysis. For res.na.action = "LOCF1" or "LOCF2", the order in the outcome is considered as the time order in imputing. if the baseline measurement is missing, then the corresponding subject is removed. outcome can be also multiple (repeatly measured). If more than two strata are specified, these are taking a cross-classification. The group variable can be specifies its reference group in the argument ref in the sanon or in the function grp.

Value

N	Sample size
Nna	tne number of subjects with missing values
nhik	Sample size in each strata, group, and response
nik	Sample size in each group and response
хi	$(multivariate)\ Mann-Whitney\ estimate(s)\ that\ addresses\ the\ comparison\ between\ two\ randomized\ groups$
g	the difference estimates between mean covariable
f	a vector consisting of xi and g
Vf	estimated covariance matrix of f
b	fully adjustmented estimators for all covariables and the strata

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Vb covariance matrix of b
se standard error of b
Q test statistics for b
p - value for b

outnames outcome or response names

covarnames covariable names

advarnames variable names adjusting in the weighted least squares bnames variable names of adjusted in the weighted least squares

reslevels levels for response variables grouplevels levels for the group variable

strtout resulting (cross-classification) strata

strtlevels resulting (cross-classification) strata levels strtnames resulting (cross-classification) strata names

matP design matrix used in the weighted least squares

References

Kawaguchi A., Koch, G. G. (2015). sanon: An R Package for Stratified Analysis with Nonparametric Covariable Adjustment. Journal of Statistical Software, 67(9), 1-37. doi:10.18637/jss.v067.i09

Kawaguchi, A., Koch, G. G., Wang, X. (2011): Stratified Multivariate Mann-Whitney Estimators for the Comparison of Two Treatments with Randomization Based Covariance Adjustment. Statistics in Biopharmaceutical Research, Vol. 3, No. 2, 217-231.

Examples

```
##### Example 3.1 Randomized Clinical Trial of Chronic Pain #####
data(cpain)
out11 = sanon(response ~ grp(treat, ref="placebo") + strt(center) + strt(diagnosis), data=cpain)
summary(out11)
# R objects are also available
attach(cpain)
out12 = sanon(outcome=response, group=treat,
strt=cbind(center, diagnosis), ref="placebo")
out12
summary(out12)
##### Example 3.2 Randomized Clinical Trial of Respiratory Disorder #####
data(resp)
out21 = sanon(cbind(baseline, visit1, visit2, visit3, visit4)
~ grp(treatment, ref="P") + strt(center) + strt(sex) + covar(age), data=resp)
out21
summary(out21)
# the matrix P can be specified
```

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```
P = rbind(rep(0, 4), diag(4), rep(0, 4))
out22 = sanon(cbind(baseline, visit1, visit2, visit3, visit4)
~ grp(treatment, ref="P") + strt(center) + strt(sex) + covar(age), data=resp, P=P)
out22
summary(out22)
```

sebor

Seborrheic Dermatitis Data

Description

The data are from a randomized clinical trial to compare a test treatment to placebo for a seborrheic dermatitis, and listings of the data appear in Ramaswamy, Koch, and Amara (1997). The variables are as follows:

Usage

data(sebor)

Format

A data frame with 167 observations and 8 variables.

Details

center a factor vector for eight centers

treat a factor with levels placebo and test for placebo and test treatments, resectively

score1 a numeric vector for patient global scores for the face according to 6 categories (0 = cleared, 1 = excellent improvement, 2 = moderate improvement, 3 = slight improvement, 4 = no change, 5 = exacerbation)

score2 a numeric vector for patient global scores for the scalp with same categories as score1

score3 a numeric vector for patient global scores for the chest with same categories as score1

severity1 a numeric vector for the baseline desease severity for the face according to 3 categories (1 = mild, 2 = moderate, 3 = severe)

severity2 a numeric vector for the baseline desease severity for the scalp with same categories as severity1

severity3 a numeric vector for the baseline desease severity for the chest with same categories as severity1

References

Ramaswamy R, Koch G, Amara I (1997). "Application of rank analysis of covariance methods to analysis of multiple anatomical regions with treatment for seborrheic dermatitis." Journal of Biopharmaceutical Statistics, 7(3), 403–416.

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skin

Skin Condition Data

Description

The data are from a randomized clinical trial to compare a test treatment to placebo for skin conditions, and listings of the data appear in Stanish, Gillings, Koch (1978a, b). The variables are as follows:

Usage

data(skin)

Format

A data frame with 172 observations and 6 variables.

Details

center a factor vector for two centers

treat a factor with levels A and P for active and placebo treatments, skinectively

stage a numeric vector for initial severity of the skin condition according to 3 categories (3 = fair, 4 = poor, 5 = exacerbation) at baseline measurement

res1 a numeric vector for extent of improvement at visit 1 according to 5 categories (1 = rapidly improving, 2 = slowly improving, 3 = stable, 4 = slowly worsening, 5 = rapidly worsening)

res2 a numeric vector for extent of improvement at visit 2 with same categories as res1

res3 a numeric vector for extent of improvement at visit 3 with same categories as res1

References

Stanish W, Gillings D, Koch G (1978a). "An application of multivariate ratio methods for the analysis of a longitudinal clinical trial with missing data." Biometrics, 34(2), pp. 305–317.

Stanish WM, Koch GG, Landis JR (1978b). "A computer program for multivariate ratio analysis (MISCAT)." Computer Programs in Biomedicine, 8(3-4), 197–207.

strt 15

strt

Identify Stratification Variables

Description

This is a special function used in the context of sanon. It identifies stratification variables when they appear on the right hand side of a formula.

Usage

```
strt(x)
```

Arguments

Х

variable name

summary.sanon

Summarizing Weighted Least Squares Fits

Description

summary method for class "sanon".

Usage

```
## S3 method for class 'sanon'
summary(object, ...)
## S3 method for class 'summary.sanon'
print(x, ...)
```

Arguments

```
object, x an object of class "sanon", usually, a result of a call to sanon further arguments passed to or from other methods.
```

Details

This function provide the p value for the hypothesis test of coefficient in the model of weighted least squares method. Note that the estimates in the output are for the $(xi_k - 0.5)$.

Value

coefficients a p x 4 matrix with columns for the estimated coefficient, its standard error,

chi-squared statistic and corresponding (two-sided) p-value.

advarnames adjust variable names in weighted least squares method

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Examples

```
##### Example 3.1 Randomized Clinical Trial of Chronic Pain #####
data(cpain)
sum1 = summary(sanon(response ~ grp(treat, ref="placebo") + strt(center) + strt(diagnosis)
, data=cpain))
sum1

##### Example 3.2 Randomized Clinical Trial of Respiratory Disorder ####
data(resp)
sum22 = summary(sanon(cbind(baseline, visit1, visit2, visit3, visit4)
~ grp(treatment, ref="P") + strt(center) + strt(sex) + covar(age), data=resp))
sum22
```

vcov sanon

Calculate Variance-Covariance Matrix for a Fitted Model Object

Description

Returns the variance-covariance matrix of the main parameters of a fitted model object.

Usage

```
## S3 method for class 'sanon'
vcov(object, ...)
```

Arguments

object an object of class "sanon", usually, a result of a call to sanon further arguments passed to or from other methods.

Details

This is a generic function.

Value

Coefficients extracted from the model object object.

Examples

```
##### Example 3.1 Randomized Clinical Trial of Chronic Pain #####
data(cpain)
out1 = sanon(response ~ grp(treat, ref="placebo") + strt(center) + strt(diagnosis), data=cpain)
vcov(out1)
##### Example 3.2 Randomized Clinical Trial of Respiratory Disorder #####
data(resp)
P = rbind(rep(0, 4), diag(4), rep(0, 4))
```

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```
out23 = sanon(cbind(baseline, visit1, visit2, visit3, visit4) ~ grp(treatment, ref="P")
    + strt(center) + strt(sex) + covar(age), data=resp, P=P)
# each four visits
vcov(out23)
```

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