Package ‘nonnest2’

March 14, 2020

Title    Tests of Non-Nested Models
Version  0.5-3
Date     2020-03-14
Description
    Testing non-nested models via theory supplied by Vuong (1989) <DOI:10.2307/1912557>. Includes tests of model distinguishability and of model fit that can be applied to both nested and non-nested models. Also includes functionality to obtain confidence intervals associated with AIC and BIC. This material is partially based on work supported by the National Science Foundation under Grant Number SES-1061334.

Depends  R (>= 3.0.0)
Imports  CompQuadForm, mvtnorm, lavaan (>= 0.6-3), sandwich
License  GPL-2 | GPL-3
LazyData yes
Suggests mirt (>= 1.26.3), AER, MASS, faraway, mlogit, ordinal, pscl, knitr, rmarkdown, testthat

VignetteBuilder knitr
RoxygenNote 7.0.2

NeedsCompilation no
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Repository CRAN
Date/Publication 2020-03-14 22:20:02 UTC

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Description

Calculate confidence intervals of AIC and BIC for non-nested models.

Usage

\[ \text{icci}(\text{object1}, \text{object2}, \text{conf.level} = 0.95, \text{ll1} = \text{llcont}, \text{ll2} = \text{llcont}) \]

Arguments

- **object1**: a model object
- **object2**: a model object
- **conf.level**: confidence level of the interval
- **ll1**: an optional function for computing log-likelihood contributions of object1
- **ll2**: an optional function for computing log-likelihood contributions of object2

Details

Functionality is currently available for models of classes `lm`, `glm`, `glm.nb`, `clm`, `hurdle`, `zeroinfl`, `mlogit`, `nls`, `polr`, `rlm`, and `lavaan`.

Users should take care to ensure that the two models have the same dependent variable (or, for lavaan objects, identical modeled variables), with observations ordered identically within each model object. Assuming the same data matrix is used to fit each model, observation ordering should generally be identical. There are currently no checks for this, however.

Note: if models are nested or if the "variance test" from `vuongtest()` indicates models are indistinguishable, then the intervals returned from `icci()` will be incorrect.

Value

an object of class `icci` containing test results.

Author(s)

Ed Merkle and Dongjun You

References


Examples

## Not run:
## Count regression comparisons
require(MASS)
house1 <- glm(Freq ~ Infl + Type + Cont, family=poisson, data=housing)
house2 <- glm(Freq ~ Infl + Sat, family=poisson, data=housing)

## CI for BIC
icci(house2, house1)

## Further comparisons to hurdle, zero-inflated models
require(pscl)
bio1 <- glm(art ~ fem + mar + phd + ment, family=poisson, data=bioChemists)
bio2 <- hurdle(art ~ fem + mar + phd + ment, data=bioChemists)
bio3 <- zeroinfl(art ~ fem + mar + phd + ment, data=bioChemists)
icci(bio2, bio1)
icci(bio3, bio1)
icci(bio3, bio2)

## Latent variable model comparisons
require(lavaan)
HS.model <- "visual =~ x1 + x2 + x3
textual =~ x4 + x5 + x6
speed =~ x7 + x8 + x9"
fit1 <- cfa(HS.model, data=HolzingerSwineford1939, meanstructure=TRUE)
fit2 <- cfa(HS.model, data=HolzingerSwineford1939, group="school")
icci(fit1, fit2)

## End(Not run)

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llcont

Individual Log-Likelihoods

Description

Obtain log-likelihood values associated with individual observations, evaluated at the ML estimates.

Usage

llcont(x, ...)

Arguments

x a model object
... arguments passed to specific methods
Details

This is a S3 generic function. Currently, the method is defined for lm, glm, glm.nb, clm, hurdle, zeroinfl, mlogit, nls, polr, rlm, lavaan, vglm, and mirt objects.

Value

An object of class numeric containing individuals’ contributions to the log-likelihood. The sum of these contributions equals the model log-likelihood.

Author(s)

Ed Merkle, Dongjun You, and Lennart Schneider

Examples

```r
## Fit gamma glm, check that sum of llcont() equals
## the model loglikelihood:
clotting <- data.frame(u = c(5,10,15,20,30,40,60,80,100),
  lot1 = c(118,58,42,35,27,25,21,19,18),
  lot2 = c(69,35,26,21,18,16,13,12,12))
gam1 <- glm(lot1 ~ log(u), data = clotting, family = Gamma)
sum(llcont(gam1))
logLik(gam1)
```

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**vuongtest**

**Vuong Tests for Model Comparison**

Description

Test pairs of models using Vuong’s (1989) <DOI:10.2307/1912557> theory. This includes a test of model distinguishability and a test of model fit.

Usage

```r
vuongtest(
  object1,
  object2, 
  nested = FALSE, 
  adj = "none", 
  ll1 = llcont, 
  ll2 = llcont, 
  score1 = NULL, 
  score2 = NULL, 
  vc1 = vcov, 
  vc2 = vcov 
)
```
Arguments

- `object1`: a model object
- `object2`: a model object
- `nested`: if `TRUE`, models are assumed to be nested
- `adj`: Should an adjusted test statistic be calculated? Defaults to “none”, with possible adjustments being “aic” and “bic”
- `ll1`: an optional function for computing log-likelihood contributions of `object1`
- `ll2`: an optional function for computing log-likelihood contributions of `object2`
- `score1`: an optional function for computing scores of `object1`
- `score2`: an optional function for computing scores of `object2`
- `vc1`: an optional function for computing the asymptotic covariance matrix of the parameters of `object1`
- `vc2`: an optional function for computing the asymptotic covariance matrix of the parameters of `object2`

Details

For non-nested models, the test of distinguishability indicates whether or not the models can possibly be distinguished on the basis of the observed data. The LRT then indicates whether or not one model fits better than another.

For nested models (`nested=TRUE`), both tests serve as robust alternatives to the classical likelihood ratio tests. In this case, the `adj` argument is ignored.

Users should take care to ensure that the two models have the same dependent variable (or, for `lavaan` objects, identical modeled variables), with observations ordered identically within each model object. Assuming the same data matrix is used to fit each model, observation ordering should generally be identical. There are currently no checks for this, however.

Value

an object of class `vuongtest` containing test results.

Author(s)

Ed Merkle and Dongjun You

References


Examples

## Not run:
## Count regression comparisons
require(MASS)
house1 <- glm(Freq ~ Infl + Type + Cont, family=poisson, data=housing)
house2 <- glm(Freq ~ Infl + Sat, family=poisson, data=housing)
house3 <- glm(Freq ~ Infl, family=poisson, data=housing)
## house3 is nested within house1 and house2
anova(house3, house1, test="Chisq")
anova(house3, house2, test="Chisq")

## house 2 is not nested in house1, so this test is invalid
anova(house2, house1, test="Chisq")

## Use vuongtest() instead
vuongtest(house2, house1)

## Application to models with different distributional assumptions
require(pscl)
bio1 <- glm(art ~ fem + mar + phd + ment, family=poisson, data=bioChemists)
bio2 <- hurdle(art ~ fem + mar + phd + ment, data=bioChemists)
bio3 <- zeroinfl(art ~ fem + mar + phd + ment, data=bioChemists)
vuongtest(bio2, bio1)
vuongtest(bio3, bio1)
vuongtest(bio1, bio2)
vuongtest(bio1, bio3)
vuongtest(bio3, bio2)

## Application to latent variable models
require(lavaan)
HS.model <- 'visual =~ x1 + x2 + x3
  textual =~ x4 + x5 + x6
  speed =~ x7 + x8 + x9'
fit1 <- cfa(HS.model, data=HolzingerSwineford1939)
fit2 <- cfa(HS.model, data=HolzingerSwineford1939, group="school")
vuongtest(fit1, fit2)

## Supplying custom vcov function
require(lme4)
require(merDeriv)
fm1 <- lmer(Reaction ~ Days + (Days | Subject), sleepstudy, REML=FALSE)
fm2 <- lmer(Reaction ~ Days + (Days || Subject), sleepstudy, REML=FALSE)
vcl <- function(obj) vcov(obj, full=TRUE)
vuongtest(fm1, fm2, vc1=vcl, vc2=vcl, nested=TRUE)

## End(Not run)
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