

Estimation and Comparison of Summary Statistics for the Receiver Operating Characteristic Curve (ROC)

Description

Estimate and compare ROC summary statistics between two markers. Choices for summary statistics are: ROC(f), the True positive rate corresponding to False positive rate f; $\text{ROC}^{-1}(t)$, the False positive rate corresponding to True positive rate t; AUC, the area under the ROC curve; and pAUC(f), the partial area under the ROC curve from 0 to f. Algorithms use the percentile value formulation of the ROC curve. When percentile values are calculated empirically, the estimates are the standard non-parametric estimations of ROC summary indices. Optional covariate adjustment can be achieved.

Usage

```
comproc(dataset = NULL, d, markers, auc = FALSE, pauc = NULL, roc = NULL, rocinv = NULL,
        pvcmeth = "empirical", tiecorr = FALSE, adjcov = NULL, adjmodel = "stratified",
        nsamp = 1000, nobstrap = FALSE, noccsamp = FALSE, nostsamp = FALSE, cluster = NULL,
        resfile = NULL, replace = FALSE, level = 95)
```

Arguments

- dataset** optional character string specifying the name of the dataset to be used for analysis.
- d** character string specifying the name of the 0/1 outcome vector.
- markers** vector of character strings specifying the names of the test markers/variables.
- auc** logical. If TRUE, compare markers with respect to the area under the curve (AUC). This is the default if no summary statistics are specified.
- pauc** specify FPR, f, such that the markers are compared with respect to the partial area under the curve (pAUC) for false positive range $\text{FPR} < f$. The argument must be between 0 and 1. A tie correction is included in the percentile value (PV) calculation if this option is included among the specified summary statistics options and the `pvcmeth="empirical"`.
- roc** specify FPR, f, such that the markers are compared with respect to the ROC at the specified $\text{FPR} = f$. The argument must be between 0 and 1.
- rocinv** specify TPR, t, such that the markers are compared with respect to the inverse ROC, $\text{ROC}^{-1}(t)$, at the specified $\text{TPR} = t$. The argument must be between 0 and 1.
- pvcmeth** character string specifying PV calculation method as "empirical" (default) or "normal". "empirical" uses the empirical distribution of the test measure among controls ($D=0$) as the reference distribution for the calculation of case PVs. The PV for the case measure y_i is the proportion of control measures that smaller than y_i . "normal" models the test measure among controls with a normal distribution. The PV for the case measure y_i is the standard normal cumulative distribution function of $(y_i - \text{mean})/\text{sd}$, where the mean and the standard deviation (sd) are calculated by using the control sample.
- tiecorr** logical. If FALSE (default), no correction for ties. If TRUE, it indicates that a correction for ties between case and control values is included in the empirical PV calculation. The correction is relevant only in calculating summary indices, such as the area under the ROC curve. The tie-corrected PV for a case with the marker value y_i is the proportion of control values $Y_{\text{Db}} < y_i$ plus one half the proportion of control values $Y_{\text{Db}} = y_i$, where Y_{Db} denotes controls. By default, the PV calculation includes only the first term, i.e. the proportion of control values $Y_{\text{Db}} < y_i$. This option applies only to the empirical PV calculation method.
- adjcov** character string vector specifying covariates to adjust for.
- adjmodel** character string specifying how the covariate adjustment is to be done: "stratified" (default), "linear", "oprobit" (ordered probit), or "ologit" (ordered logit). If "stratified", PVs are calculated separately for each stratum defined by `adjcov`. This is the default if `adjmodel` is not specified and `adjcov` is. Each case-containing stratum must include at least two controls. Strata that do not include cases are excluded from calculations. "linear" fits a linear regression of the marker distribution on the adjustment covariates among controls. Standardized residuals based on this fitted linear model are used in place of the marker values for cases and controls. "oprobit" calculates PVs based on the fit of an ordered probit regression model of the marker on the adjustment covariates among controls. "ologit" calculates PVs based on the fit of an ordered logit regression model of the marker on

the adjustment covariates among controls. "oprobit" and "ologit" assume that `markers` consists of ordinal-valued marker variables.

`nsamp` number of bootstrap samples to be drawn for estimating sampling variability of summary measures; default is `nsamp=1000`.

`nobstrap` logical. If TRUE, omit bootstrap sampling and estimation of standard errors and CIs. If `nsamp` is specified, `nobstrap` will override it. Default is FALSE.

`noccsamp` logical. If TRUE, bootstrap samples are drawn from the combined sample (cohort sampling) rather than sampling separately from cases and controls (case-control sampling); default is FALSE (case-control sampling).

`nostsamp` logical. If TRUE (default), bootstrap samples are drawn without respect to covariate strata. By default, samples are drawn from within covariate strata when stratified covariate adjustment is requested via the `adjcov` and `adjmodel` options.

`cluster` character string specifying variables that identify bootstrap resampling clusters.

`resfile` character string specifying the filename to save bootstrap results for the included statistics in. The .txt file is called [filename].txt if a single marker is specified or [filename]#.txt for the #th marker if more than 1 marker is included in `markers`.

`replace` logical. If TRUE, overwrite existing specified bootstrap results file if it already exists; default is FALSE.

`level` specify confidence level for CIs as a percentage; default is `level=95`.

Details

`comproc` compares two continuous marker or test variables with respect to one or more ROC statistics: the AUC, the pAUC for $FPR < f$, the ROC at $FPR = f$, and the inverse ROC at $TPR = t$. `d` is the 0/1 outcome indicator variable.

Alternatively, a single marker variable can be specified, in which case the requested ROC statistics are returned without comparison statistics.

All ROC statistics are calculated by using PVs of the disease case measures relative to the corresponding marker distribution among controls (Pepe and Longton (2005), Huang and Pepe (in press)).

Optional covariate adjustment can be achieved either by stratification or with a linear regression approach (Janes and Pepe (2008); Janes and Pepe (2009)). Ordered regression covariate adjustment options are available if the `markers` measures are ordinal (Morris, Pepe, Barlow (in press)).

Bootstrap standard errors and confidence intervals (CIs) for the requested statistics and marker differences are calculated. Percentile CIs are displayed.

Wald test results for marker comparisons are based on the bootstrap standard errors for the difference between markers.

A companion program for the Stata software package is available. A detailed description of the methods and algorithms are provide in two articles in the Stata Journal which can be obtained upon request from Gary Longton (glongton@fhcrc.org). Corresponding articles for this program are forthcoming.

Value

List containing properties for requested summary statistics, where `stat` is one or more of `auc`, `pauc`, `roc` or `rocinv`. Returned list items include the following:

```
[stat]1      statistic estimate for first marker
[stat]2      statistic estimate for second marker
[stat]delta  estimate difference, [stat]2 - [stat]1
se_[stat]1   bootstrap standard-error estimate for first marker statistic
se_[stat]2   bootstrap standard-error estimate for second marker statistic
se_[stat]delta bootstrap standard-error estimate for the difference, [stat]2 - [stat]1
```

Author(s)

Aasthaa Bansal, University of Washington, Seattle, WA. abansal@u.washington.edu

Daryl Morris, University of Washington, Seattle, WA. daryl@u.washington.edu

Gary Longton, Fred Hutchinson Cancer Research Center, Seattle, WA. glongton@fhcrc.org

Margaret Pepe, Fred Hutchinson Cancer Research Center and University of Washington, Seattle, WA. mspepe@u.washington.edu

Holly Janes, Fred Hutchinson Cancer Research Center and University of Washington, Seattle, WA. hjanes@fhcrc.org

References

Dodd, L., Pepe, M.S. 2003. Partial AUC estimation and regression. *Biometrics* **59**,614–623.

Huang, Y., Pepe, M.S. 2009. Biomarker evaluation using the controls as a reference population. *Biostatistics* **2**,228–44.

Janes, H., Pepe, M.S. 2008. Adjusting for covariates in studies of diagnostic, screening, or prognostic markers: an old concept in a new setting. *American Journal of Epidemiology* **168**,89–97.

Janes, H., Pepe, M.S. 2009. Adjusting for covariate effects on classification accuracy using the covariate-adjusted ROC curve. *Biometrika* **96**,383–398.

Janes, H., Longton G, Pepe, M.S. 2009. Accommodating covariates in receiver operating characteristic analysis. *Stata Journal* **9**(1),17–39.

Morris, D.E., Pepe, M.S., Barlow, W.E. Contrasting Two Frameworks for ROC Analysis of Ordinal Ratings. *Medical Decision Making* (in press)

Pepe, M.S., Longton, G. 2005. Standardizing markers to evaluate and compare their performances. *Epidemiology* **16**(5),598-603.

Pepe MS, Longton G, Janes H. 2009. Estimation and comparison of receiver operating characteristic curves. *Stata Journal* **9**(1),1–16.

Pepe, M.S. 2003. *The Statistical Evaluation of Medical Tests for Classification and Prediction*. Oxford University Press.

See Also

[roccurve](#), [roclog](#).

Examples

```
nnhs2 <- read.csv("http://labs.fhcrc.org/pepe/book/data/nnhs2.csv",
  header = TRUE, sep = ",")

comproc(dataset="nnhs2", d="d", markers=c("y1","y2"))
comproc(dataset="nnhs2", d="d", markers=c("y1","y2"), pauc=0.10)
comproc(dataset="nnhs2", d="d", markers=c("y1","y2"), auc=TRUE, pauc=0.10, level=90)
comproc(dataset="nnhs2", d="d", markers=c("y1","y2"), auc=TRUE, pauc=0.20, roc=0.20,
  nsamp=5000)
comproc(dataset="nnhs2", d="d", markers=c("y1","y2"), pauc=0.20,
  pvcmeth="normal", resfile="rfile1", replace=TRUE)
comproc(dataset="nnhs2", d="d", markers=c("y1","y2"), pvcmeth="normal",
  noccsamp=TRUE, cluster="y1")
comproc(dataset="nnhs2", d="d", markers=c("y1","y2"),
  adjcov=c("currence","gender"))
comproc(dataset="nnhs2", d="d", markers=c("y1","y2"), adjcov=c("currence","gender"),
  adjmodel="linear")
comproc(dataset="nnhs2", d="d", markers=c("y1","y2"), adjcov="currence",
  adjmodel="linear", pvcmeth="normal")
```