# Package: dcurves (via r-universe)

August 23, 2024

Title Decision Curve Analysis for Model Evaluation

Version 0.5.0

Description Diagnostic and prognostic models are typically evaluated with measures of accuracy that do not address clinical consequences. Decision-analytic techniques allow assessment of clinical outcomes, but often require collection of additional information may be cumbersome to apply to models that yield a continuous result. Decision curve analysis is a method for evaluating and comparing prediction models that incorporates clinical consequences, requires only the data set on which the models are tested, and can be applied to models that have either continuous or dichotomous results. See the following references for details on the methods: Vickers (2006) <doi:10.1177/0272989X06295361>, Vickers (2008) <doi:10.1186/1472-6947-8-53>, and Pfeiffer (2020) <doi:10.1002/bimj.201800240>.

```
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```

URL https://github.com/ddsjoberg/dcurves,

https://www.danieldsjoberg.com/dcurves/

BugReports https://github.com/ddsjoberg/dcurves/issues

**Depends** R (>= 3.5)

- **Imports** broom (>= 0.7.10), dplyr (>= 1.0.5), ggplot2 (>= 3.3.3), glue (>= 1.4.2), purrr (>= 0.3.4), rlang (>= 0.4.10), scales (>= 1.1.1), survival, tibble (>= 3.1.0)
- **Suggests** broom.helpers (>= 1.15.0), covr (>= 3.5.1), gtsummary (>= 2.0.0), knitr (>= 1.32), rmarkdown (>= 2.7), spelling (>= 2.2), testthat (>= 3.0.2), tidyr (>= 1.1.3)

VignetteBuilder knitr

ByteCompile true

Config/testthat/edition 3

Config/testthat/parallel true

**Encoding** UTF-8

Language en-US

LazyData true

RoxygenNote 7.2.3

NeedsCompilation no

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**Repository** CRAN

Date/Publication 2024-07-23 23:20:01 UTC

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as\_tibble.dca Convert DCA Object to tibble

#### Description

Convert DCA Object to tibble

## Usage

## S3 method for class 'dca'
as\_tibble(x, ...)

#### Arguments

х	dca object created with dca()
	not used

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dca

#### Value

a tibble

#### Author(s)

Daniel D Sjoberg

#### See Also

dca(), net\_intervention\_avoided(), standardized\_net\_benefit(), plot.dca()

#### Examples

```
dca(cancer ~ cancerpredmarker, data = df_binary) %>%
    as_tibble()
```

dca

Perform Decision Curve Analysis

#### Description

Diagnostic and prognostic models are typically evaluated with measures of accuracy that do not address clinical consequences. Decision-analytic techniques allow assessment of clinical outcomes but often require collection of additional information may be cumbersome to apply to models that yield a continuous result. Decision curve analysis is a method for evaluating and comparing prediction models that incorporates clinical consequences, requires only the data set on which the models are tested, and can be applied to models that have either continuous or dichotomous results. The dca function performs decision curve analysis for binary outcomes. Review the DCA Vignette for a detailed walk-through of various applications. Also, see www.decisioncurveanalysis.org for more information.

#### Usage

```
dca(
   formula,
   data,
   thresholds = seq(0, 0.99, by = 0.01),
   label = NULL,
   harm = NULL,
   as_probability = character(),
   time = NULL,
   prevalence = NULL
)
```

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formula	a formula with the outcome on the LHS and a sum of markers/covariates to test on the RHS
data	a data frame containing the variables in formula=.
thresholds	vector of threshold probabilities between 0 and 1. Default is $seq(0, 0.99)$ , by = 0.01). Thresholds at zero are replaced with 10e-10.
label	named list of variable labels, e.g. list(age = "Age, years")
harm	named list of harms associated with a test. Default is NULL
as_probability	character vector including names of variables that will be converted to a probability. Details below.
time	if outcome is survival, time= specifies the time the assessment is made
prevalence	When NULL, the prevalence is estimated from data=. If the data passed is a case-control set, the population prevalence may be set with this argument.

#### Value

List including net benefit of each variable

#### as\_probability argument

While the as\_probability= argument can be used to convert a marker to the probability scale, use the argument only when the consequences are fully understood. For example, when the outcome is binary, logistic regression is used to convert the marker to a probability. The logistic regression model assumes linearity on the log-odds scale and can induce miscalibration when this assumption is not true. Miscalibration in a model will adversely affect performance on decision curve analysis. Similarly, when the outcome is time-to-event, Cox Proportional Hazards regression is used to convert the marker to a probability. The Cox model also has a linearity assumption and additionally assumes proportional hazards over the follow-up period. When these assumptions are violated, important miscalibration may occur.

Instead of using the as\_probability= argument, it is suggested to perform the regression modeling outside of the dca() function utilizing methods, such as non-linear modeling, as appropriate.

#### Author(s)

Daniel D Sjoberg

#### See Also

net\_intervention\_avoided(), standardized\_net\_benefit(), plot.dca(), as\_tibble.dca()

## Examples

#### df\_binary

```
# plot DCA curves with ggplot
plot(smooth = TRUE) +
# add ggplot formatting
ggplot2::labs(x = "Treatment Threshold Probability")
# calculate DCA with time to event endpoint
dca(Surv(ttcancer, cancer) ~ cancerpredmarker, data = df_surv, time = 1)
```

df\_binary

Simulated data with a binary outcome

#### Description

Simulated data with a binary outcome

#### Usage

df\_binary

#### Format

A data frame with 750 rows:

patientid Identification Number cancer Cancer Diagnosis: 0=No, 1=Yes dead Dead (1=yes; 0=no) risk\_group Patient Risk Group (Low, Intermediate, High) age Patient Age, years famhistory Family History of Cancer: 0=No, 1=Yes marker Marker cancerpredmarker Prob. of Cancer based on Age, Family History, and Marker

df\_case\_control Simulated data with a case-control outcome

# Description

Simulated data with a case-control outcome

#### Usage

df\_case\_control

#### Format

A data frame with 750 rows:

patientid Identification Number casecontrol Case-control Status: 1=Case, 0=Control risk\_group Patient Risk Group (Low, Intermediate, High) age Patient Age, years famhistory Family History of Cancer: 0=No, 1=Yes marker Marker cancerpredmarker Prob. of Cancer based on Age, Family History, and Marker

df\_surv

Simulated data with a survival outcome

## Description

Simulated data with a survival outcome

#### Usage

df\_surv

# Format

A data frame with 750 rows:

patientid Identification Number

cancer Cancer Diagnosis: 0=No, 1=Yes

cancer\_cr Cancer Diagnosis, competing event: "censor", "dead other causes", "diagnosed with cancer"

ttcancer Years to Cancer Dx/Censor

risk\_group Patient Risk Group (Low, Intermediate, High)

age Patient Age, years

famhistory Family History of Cancer: 0=No, 1=Yes

marker Marker

cancerpredmarker Prob. of Cancer based on Age, Family History, and Marker

net\_intervention\_avoided

Add Net Interventions Avoided

#### Description

Add the number of net interventions avoided to dca() object.

# Usage

```
net_intervention_avoided(x, nper = 1)
```

# Arguments

х	object of class 'dca' calculated with dca()
nper	Number to report net interventions per. Default is 1

# Value

'dca' object

# Author(s)

Daniel D Sjoberg

# See Also

dca(), standardized\_net\_benefit(), plot.dca(), as\_tibble.dca()

# Examples

```
dca(
  cancer ~ cancerpredmarker,
  data = df_binary
) %>%
  net_intervention_avoided()
dca(
  Surv(ttcancer, cancer) ~ cancerpredmarker,
  data = df_surv,
  time = 1
) %>%
  net_intervention_avoided(nper = 100)
```

plot.dca

# Description

Plot DCA Object with ggplot

# Usage

```
## S3 method for class 'dca'
plot(
    x,
    type = NULL,
    smooth = FALSE,
    span = 0.2,
    style = c("color", "bw"),
    show_ggplot_code = FALSE,
    ...
)
```

# Arguments

x	dca object created with dca()
type	<pre>indicates type of plot to produce. Must be one of c("net_benefit", "net_intervention_avoided", "standardized_net_benefit"). The default is "net_benefit", unless the net intervention has been calculated when "net_intervention_avoided" is used, or if "standardized_net_benefit" has been calculated.</pre>
smooth	Logical indicator whether plot will be smooth with ggplot2::stat_smooth(). Default is FALSE
span	when smooth = TRUE, Controls the amount of smoothing for loess smoother. Smaller numbers produce wigglier lines, larger numbers produce smoother lines. Default is 0.2.
style	Must be one of c("color", "bw"). Default is "color", and "bw" will print a black and white figure
<pre>show_ggplot_cod</pre>	le
	Logical indicating whether to print ggplot2 code used to create figure. Default is FALSE. Set to TRUE to perform advanced figure customization
	not used

# Value

a ggplot2 object

#### Author(s)

Daniel D Sjoberg

#### See Also

```
dca(), net_intervention_avoided(), standardized_net_benefit(), as_tibble.dca()
```

#### Examples

```
p <-
dca(cancer ~ cancerpredmarker, data = df_binary) %>%
plot(smooth = TRUE, show_ggplot_code = TRUE)
p
# change the line colors
p + ggplot2::scale_color_manual(values = c('black', 'grey', 'purple'))
```

standardized\_net\_benefit

Add Standardized Net Benefit

# Description

Add the standardized net benefit to dca() object.

#### Usage

```
standardized_net_benefit(x)
```

# Arguments ×

object of class 'dca' calculated with dca()

#### Value

'dca' object

#### Author(s)

Daniel D Sjoberg

#### See Also

dca(), net\_intervention\_avoided(), plot.dca(), as\_tibble.dca()

#### Examples

```
dca(Surv(ttcancer, cancer) ~ cancerpredmarker, data = df_surv, time = 1) %>%
    standardized_net_benefit()
```

test\_consequences Test Consequences

## Description

Test Consequences

# Usage

```
test_consequences(
  formula,
  data,
  statistics = c("pos_rate", "neg_rate", "test_pos_rate", "test_neg_rate", "tp_rate",
    "fp_rate", "fn_rate", "tn_rate", "ppv", "npv", "sens", "spec", "lr_pos", "lr_neg"),
    thresholds = seq(0, 1, by = 0.25),
    label = NULL,
    time = NULL,
    prevalence = NULL
)
```

# Arguments

formula	a formula with the outcome on the LHS and a sum of markers/covariates to test on the RHS
data	a data frame containing the variables in formula=.
statistics	Character vector with statistics to return. See below for details
thresholds	vector of threshold probabilities between 0 and 1. Default is seq(0, 0.99, by = 0.01). Thresholds at zero are replaced with 10e-10.
label	named list of variable labels, e.g. list(age = "Age, years")
time	if outcome is survival, time= specifies the time the assessment is made
prevalence	When NULL, the prevalence is estimated from data=. If the data passed is a case-control set, the population prevalence may be set with this argument.

#### Value

a tibble with test consequences

#### statistics

The following diagnostic statistics are available to return.

Statistic	Abbreviation	Definition
Outcome Positive Rate	"pos_rate"	(a + c) / (a + b + c + d)
Outcome Negative Rate	"neg_rate"	(b + d) / (a + b + c + d)
Test Positive Rate	"test_pos_rate"	(a + b) / (a + b + c + d)

Test Negative Rate	"test_neg_rate"	(c + d) / (a + b + c + d)
True Positive Rate	"tp_rate"	a / (a + b + c + d)
False Positive Rate	"fp_rate"	b / (a + b + c + d)
False Negative Rate	"fn_rate"	c / (a + b + c + d)
True Negative Rate	"tn_rate"	d / (a + b + c + d)
Positive Predictive Value	"pp∨"	a / (a + b)
Negative Predictive Value	"np∨"	d / (c + d)
Sensitivity	"sens"	a / (a + c)
Specificity	"spec"	d / (b + d)
Positive Likelihood Ratio	"lr_pos"	sens / (1 - spec)
Negative Likelihood Ratio	"lr_neg"	(1 - sens) / spec

# Examples

test\_consequences(cancer ~ cancerpredmarker, data = df\_binary)

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