Package 'survex'

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Title Explainable Machine Learning in Survival Analysis

Version 1.2.0

Description Survival analysis models are commonly used in medicine and other areas. Many of them are too complex to be interpreted by human. Exploration and explanation is needed, but standard methods do not give a broad enough picture. 'survex' provides easy-to-apply methods for explaining survival models, both complex black-boxes and simpler statistical models. They include methods specific to survival analysis such as SurvSHAP(t) introduced in Krzyzin-ski et al., (2023)

<doi:10.1016/j.knosys.2022.110234>, SurvLIME described in Kovalev et al., (2020) <doi:10.1016/j.knosys.2020.106164> as well as

extensions of existing ones described in Biecek et al., (2021) <doi:10.1201/9780429027192>.

License GPL (>= 3)

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brier_score

Calculate Brier score

Description

A function for calculating the Brier score for a survival model.

Usage

```
brier_score(y_true = NULL, risk = NULL, surv = NULL, times = NULL)
```

```
loss_brier_score(y_true = NULL, risk = NULL, surv = NULL, times = NULL)
```

Arguments

y_true	a survival::Surv object containing the times and statuses of observations for which the metric will be evaluated
risk	ignored, left for compatibility with other metrics
surv	a matrix containing the predicted survival functions for the considered observa- tions, each row represents a single observation, whereas each column one time point
times	a vector of time points at which the survival function was evaluated

Details

Brier score is used to evaluate the performance of a survival model, based on the squared distance between the predicted survival function and the actual event time, weighted to account for censored observations.

Value

numeric from 0 to 1, lower scores are better (Brier score of 0.25 represents a model which returns always returns 0.5 as the predicted survival function)

References

- [1] Brier, Glenn W. "Verification of forecasts expressed in terms of probability." Monthly Weather Review 78.1 (1950): 1-3.
- [2] Graf, Erika, et al. "Assessment and comparison of prognostic classification schemes for survival data." Statistics in Medicine 18.17-18 (1999): 2529-2545.

See Also

cd_auc()

Examples

```
library(survival)
library(survex)

cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
cph_exp <- explain(cph)

y <- cph_exp$y
times <- cph_exp$times
surv <- cph_exp$times
surv <- cph_exp$predict_survival_function(cph, cph_exp$data, times)
brier_score(y, surv = surv, times = times)
loss_brier_score(y, surv = surv, times = times)</pre>
```

cd_auc

Calculate Cumulative/Dynamic AUC

Description

This function calculates the Cumulative/Dynamic AUC metric for a survival model. It is done using the estimator proposed proposed by Uno et al. [1], and Hung and Chang [2].

Usage

cd_auc(y_true = NULL, risk = NULL, surv = NULL, times = NULL)

Arguments

y_true	a survival::Surv object containing the times and statuses of observations for which the metric will be evaluated
risk	ignored, left for compatibility with other metrics
surv	a matrix containing the predicted survival functions for the considered observa- tions, each row represents a single observation, whereas each column one time point
times	a vector of time points at which the survival function was evaluated

Details

C/D AUC is an extension of the AUC metric known from classification models. Its values represent the model's performance at specific time points. It can be integrated over the considered time range.

Value

a numeric vector of length equal to the length of the times vector, each value (from the range from 0 to 1) represents the AUC metric at a specific time point, with higher values indicating better performance.

References

- [1] Uno, Hajime, et al. "Evaluating prediction rules for t-year survivors with censored regression models." Journal of the American Statistical Association 102.478 (2007): 527-537.
- [2] Hung, Hung, and Chin-Tsang Chiang. "Optimal composite markers for time dependent receiver operating characteristic curves with censored survival data." Scandinavian Journal of Statistics 37.4 (2010): 664-679.

See Also

```
loss_one_minus_cd_auc() integrated_cd_auc() brier_score()
```

Examples

```
library(survival)
library(survex)
cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
cph_exp <- explain(cph)
y <- cph_exp$y
times <- cph_exp$times
surv <- cph_exp$times
surv <- cph_exp$predict_survival_function(cph, cph_exp$data, times)
cd_auc(y, surv = surv, times = times)</pre>
```

Description

Helper function to transform between CHF and survival function

Usage

```
cumulative_hazard_to_survival(hazard_functions)
```

Arguments

hazard_functions

matrix or vector, with each row representing a cumulative hazard function

Value

A matrix or vector transformed to the form of a survival function.

Examples

library(survex)

```
vec <- c(1, 2, 3, 4, 5)
matr <- matrix(c(1, 2, 3, 2, 4, 6), ncol = 3)
cumulative_hazard_to_survival(vec)
cumulative_hazard_to_survival(matr)</pre>
```

c_index

Compute the Harrell's Concordance index

Description

A function to compute the Harrell's concordance index of a survival model.

Usage

```
c_index(y_true = NULL, risk = NULL, surv = NULL, times = NULL)
```

Arguments

y_true	a survival::Surv object containing the times and statuses of observations for which the metric will be evaluated
risk	a numeric vector of risk scores corresponding to each observation
surv	ignored, left for compatibility with other metrics
times	ignored, left for compatibility with other metrics

Value

numeric from 0 to 1, higher values indicate better performance

References

• [1] Harrell, F.E., Jr., et al. "Regression modelling strategies for improved prognostic prediction." Statistics in Medicine 3.2 (1984): 143-152.

See Also

loss_one_minus_c_index()

explain_survival

Examples

```
library(survival)
library(survex)

rotterdam <- survival::rotterdam
rotterdam$year <- NULL
cox_rotterdam_rec <- coxph(Surv(rtime, recur) ~ .,
    data = rotterdam,
    model = TRUE, x = TRUE, y = TRUE
)
coxph_explainer <- explain(cox_rotterdam_rec)</pre>
```

risk <- coxph_explainer\$predict_function(coxph_explainer\$model, coxph_explainer\$data)
c_index(y_true = coxph_explainer\$y, risk = risk)</pre>

explain_survival A model-agnostic explainer for survival models

Description

Black-box models have vastly different structures. explain_survival() returns an explainer object that can be further processed for creating prediction explanations and their visualizations. This function is used to manually create explainers for models not covered by the survex package. For selected models the extraction of information can be done automatically. To do this, you can call the explain() function for survival models from mlr3proba, censored, randomForestSRC, ranger, survival packages and any other model with pec::predictSurvProb() method.

Usage

```
explain_survival(
 model,
 data = NULL,
 y = NULL,
  predict_function = NULL,
 predict_function_target_column = NULL,
  residual_function = NULL,
 weights = NULL,
  ...,
  label = NULL,
  verbose = TRUE,
  colorize = !isTRUE(getOption("knitr.in.progress")),
 model_info = NULL,
  type = NULL,
  times = NULL,
  times_generation = "survival_quantiles",
```

```
predict_survival_function = NULL,
 predict_cumulative_hazard_function = NULL
)
explain(
 model,
 data = NULL,
 y = NULL,
  predict_function = NULL,
 predict_function_target_column = NULL,
  residual_function = NULL,
 weights = NULL,
  ...,
  label = NULL,
  verbose = TRUE,
  colorize = !isTRUE(getOption("knitr.in.progress")),
 model_info = NULL,
  type = NULL
)
## Default S3 method:
explain(
 model,
 data = NULL,
 y = NULL,
  predict_function = NULL,
 predict_function_target_column = NULL,
  residual_function = NULL,
 weights = NULL,
  ...,
  label = NULL,
  verbose = TRUE,
  colorize = !isTRUE(getOption("knitr.in.progress")),
  model_info = NULL,
  type = NULL
)
```

Arguments

model	object - a survival model to be explained
data	data.frame - data which will be used to calculate the explanations. If not pro- vided, then it will be extracted from the model if possible. It should not contain the target columns. NOTE: If the target variable is present in the data some functionality breaks.
У	survival::Surv object containing event/censoring times and statuses corresponding to data
predict_function	

function taking 2 arguments - model and newdata and returning a single number

	for each observation - risk score. Observations with higher score are more likely to observe the event sooner.	
predict_function	on_target_column	
[····	unused, left for compatibility with DALEX	
residual_funct:	ion	
	unused, left for compatibility with DALEX	
weights	unused, left for compatibility with DALEX	
	additional arguments, passed to DALEX::explain()	
label	character - the name of the model. Used to differentiate on visualizations with multiple explainers. By default it's extracted from the 'class' attribute of the model if possible.	
verbose	logical, if TRUE (default) then diagnostic messages will be printed	
colorize	logical, if TRUE (default) then WARNINGS, ERRORS and NOTES are col- orized. Will work only in the R console. By default it is FALSE while knitting and TRUE otherwise.	
model_info	a named list (package, version, type) containing information about model. If NULL, survex will seek for information on its own.	
type	type of a model, by default "survival"	
times	numeric, a vector of times at which the survival function and cumulative hazard function should be evaluated for calculations	
times_generation		
	either "survival_quantiles", "uniform" or "quantiles". Sets the way of generating the vector of times based on times provided in the y parameter. If "survival_quantiles" the vector contains unique time points out of 50 uniformly distributed survival quantiles based on the Kaplan-Meier estimator, and additional time point being the median survival time (if possible); if "uniform" the vector contains 50 equally spaced time points between the minimum and maximum observed times; if "quantiles" the vector contains unique time points out of 50 time points between 0th and 98th percentiles of observed times. Ignored if times is not NULL.	
<pre>predict_survival_function</pre>		
predict_cumula	function taking 3 arguments model, newdata and times, and returning a matrix whose each row is a survival function evaluated at times for one observation from newdata tive_hazard_function function taking 3 arguments model, newdata and times, and returning a matrix	
	function taking 3 arguments model, newdata and times, and returning a matrix whose each row is a cumulative hazard function evaluated at times for one observation from newdata	

Value

It is a list containing the following elements:

- model the explained model.
- data the dataset used for training.
- y response for observations from data.

- residuals calculated residuals.
- predict_function function that may be used for model predictions, shall return a single numerical value for each observation.
- residual_function function that returns residuals, shall return a single numerical value for each observation.
- class class/classes of a model.
- label label of explainer.
- model_info named list containing basic information about model, like package, version of package and type.
- times a vector of times, that are used for evaluation of survival function and cumulative hazard function by default
- predict_survival_function function that is used for model predictions in the form of survival function
- predict_cumulative_hazard_function function that is used for model predictions in the form of cumulative hazard function

```
library(survival)
library(survex)
cph <- survival::coxph(survival::Surv(time, status) ~ .,</pre>
    data = veteran,
    model = TRUE, x = TRUE
)
cph_exp <- explain(cph)</pre>
rsf_ranger <- ranger::ranger(survival::Surv(time, status) ~ .,</pre>
    data = veteran,
    respect.unordered.factors = TRUE, num.trees = 100, mtry = 3, max.depth = 5
)
rsf_ranger_exp <- explain(rsf_ranger,</pre>
    data = veteran[, -c(3, 4)],
    y = Surv(veteran$time, veteran$status)
)
rsf_src <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)</pre>
rsf_src_exp <- explain(rsf_src)</pre>
library(censored, quietly = TRUE)
bt <- parsnip::boost_tree() %>%
    parsnip::set_engine("mboost") %>%
    parsnip::set_mode("censored regression") %>%
    generics::fit(survival::Surv(time, status) ~ ., data = veteran)
bt_exp <- explain(bt, data = veteran[, -c(3, 4)], y = Surv(veteran$time, veteran$status))</pre>
###### explain_survival() ######
```

```
cph <- coxph(Surv(time, status) ~ ., data = veteran)
veteran_data <- veteran[, -c(3, 4)]
veteran_y <- Surv(veteran$time, veteran$status)
risk_pred <- function(model, newdata) predict(model, newdata, type = "risk")
surv_pred <- function(model, newdata, times) pec::predictSurvProb(model, newdata, times)
chf_pred <- function(model, newdata, times) -log(surv_pred(model, newdata, times))
manual_cph_explainer <- explain_survival(
    model = cph,
    data = veteran_data,
    y = veteran_y,
    predict_function = risk_pred,
    predict_survival_function = surv_pred,
    predict_cumulative_hazard_function = chf_pred,
    label = "manual coxph"
)</pre>
```

extract_predict_survshap

Extract Local SurvSHAP(t) from Global SurvSHAP(t)

Description

Helper function to extract local SurvSHAP(t) explanation from global one. Can be can be useful for creating SurvSHAP(t) plots for single observations.

Usage

extract_predict_survshap(aggregated_survshap, index)

Arguments

aggregated_su	rvshap
	an object of class aggregated_surv_shap containing the computed global SHAP values
index	a numeric value, position of an observation to be extracted in the result of global explanation

Value

An object of classes c("predict_parts_survival", "surv_shap"). It is a list with the element result containing the results of the explanation.

Examples

```
veteran <- survival::veteran</pre>
rsf_ranger <- ranger::ranger(</pre>
    survival::Surv(time, status) ~ .,
    data = veteran,
    respect.unordered.factors = TRUE,
    num.trees = 100,
    mtry = 3,
    max.depth = 5
)
rsf_ranger_exp <- explain(</pre>
    rsf_ranger,
    data = veteran[, -c(3, 4)],
    y = survival::Surv(veteran$time, veteran$status),
    verbose = FALSE
)
ranger_global_survshap <- model_survshap(</pre>
    explainer = rsf_ranger_exp,
    new_observation = veteran[
        c(1:4, 17:20, 110:113, 126:129),
        !colnames(veteran) %in% c("time", "status")
    ]
)
local_survshap_1 <- extract_predict_survshap(ranger_global_survshap, index = 1)</pre>
plot(local_survshap_1)
```

integrated_brier_score

Calculate integrated Brier score

Description

This function calculates the integrated Brier score metric for a survival model.

Usage

```
integrated_brier_score(y_true = NULL, risk = NULL, surv = NULL, times = NULL)
loss_integrated_brier_score(
  y_true = NULL,
  risk = NULL,
  surv = NULL,
  times = NULL
)
```

Arguments

y_true	a survival::Surv object containing the times and statuses of observations for which the metric will be evaluated
risk	ignored, left for compatibility with other metrics
surv	a matrix containing the predicted survival functions for the considered observa- tions, each row represents a single observation, whereas each column one time point
times	a vector of time points at which the survival function was evaluated

Details

It is useful to see how a model performs as a whole, not at specific time points, for example for easier comparison. This function allows for calculating the integral of Brier score metric numerically using the trapezoid method.

Value

numeric from 0 to 1, lower values indicate better performance

References

- [1] Brier, Glenn W. "Verification of forecasts expressed in terms of probability." Monthly Weather Review 78.1 (1950): 1-3.
- [2] Graf, Erika, et al. "Assessment and comparison of prognostic classification schemes for survival data." Statistics in Medicine 18.17-18 (1999): 2529-2545.

See Also

brier_score() integrated_cd_auc() loss_one_minus_integrated_cd_auc()

```
library(survival)
library(survex)

cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
cph_exp <- explain(cph)

y <- cph_exp$y
times <- cph_exp$y
times surv <- cph_exp$times
surv <- cph_exp$predict_survival_function(cph, cph_exp$data, times)

# calculating directly
integrated_brier_score(y, surv = surv, times = times)</pre>
```

integrated_cd_auc Calculate integrated C/D AUC

Description

This function calculates the integrated Cumulative/Dynamic AUC metric for a survival model.

Usage

integrated_cd_auc(y_true = NULL, risk = NULL, surv = NULL, times = NULL)

Arguments

y_true	a survival::Surv object containing the times and statuses of observations for which the metric will be evaluated
risk	ignored, left for compatibility with other metrics
surv	a matrix containing the predicted survival functions for the considered observa- tions, each row represents a single observation, whereas each column one time point
times	a vector of time points at which the survival function was evaluated

Details

It is useful to see how a model performs as a whole, not at specific time points, for example for easier comparison. This function allows for calculating the integral of the C/D AUC metric numerically using the trapezoid method.

Value

numeric from 0 to 1, higher values indicate better performance

#' @section References:

- [1] Uno, Hajime, et al. "Evaluating prediction rules for t-year survivors with censored regression models." Journal of the American Statistical Association 102.478 (2007): 527-537.
- [2] Hung, Hung, and Chin-Tsang Chiang. "Optimal composite markers for time-dependent receiver operating characteristic curves with censored survival data." Scandinavian Journal of Statistics 37.4 (2010): 664-679.

See Also

```
cd_auc() loss_one_minus_cd_auc()
```

loss_adapt_mlr3proba

Examples

```
library(survival)
library(survex)
cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
cph_exp <- explain(cph)
y <- cph_exp$y
times <- cph_exp$y
times <- cph_exp$times
surv <- cph_exp$predict_survival_function(cph, cph_exp$data, times)
integrated_cd_auc(y, surv = surv, times = times)</pre>
```

loss_adapt_mlr3proba Adapt mlr3proba measures for use with survex

Description

This function allows for usage of standardized measures from the mlr3proba package with survex.

Usage

```
loss_adapt_mlr3proba(measure, reverse = FALSE, ...)
```

Arguments

measure	• a MeasureSurv object from the mlr3proba package, the object to adapt
reverse	• boolean, FALSE by default, whether the metric should be reversed in order to be treated as loss (for permutational variable importance we need func- tions with lower values indicating better performance). If TRUE, the new metric value will be (1 - metric_value)
	other parameters, currently ignored

Value

a function with standardized parameters ($y_true, risk, surv, times$) that can be used to calculate loss

```
if(FALSE){
    measure <- msr("surv.calib_beta")
    mlr_measure <- loss_adapt_mlr3proba(measure)
}</pre>
```

loss_integrate

Description

This function allows for creating a function for calculation of integrated metrics based on a time dependent metric. A possibility to cut off the data at certain quantiles is implemented, as well as weighting the integrated metric by max time and marginal survival function [1]

Usage

```
loss_integrate(loss_function, ..., normalization = NULL, max_quantile = 1)
```

Arguments

loss_function	• A time dependent loss function taking arguments (y_true, risk, surv, times)
	• other parameters, currently ignored
normalization	 either NULL, "t_max" or "survival". Decides what kind of weighting should be applied to the integrated metric. If "t_max", then the integral is calcu- lated using dw(t) where w(t) = t/t_max. If "survival", then the integral is calculated using dw(t) where w(t) = (1 - S(t))/(1 - S(t_max)) and S(t) denotes the estimated marginal survival function. If NULL (default), the integral is calculated using dt.
<pre>max_quantile</pre>	• a number from the interval (0,1]. The integral will be calculated only up to the time value of quantile(max_quantile) of the observed event/censoring times in y_true.

Value

a function that can be used to calculate metrics (with parameters y_true, risk, surv, and times)

References

• [1] Graf, Erika, et al. "Assessment and comparison of prognostic classification schemes for survival data." Statistics in Medicine 18.17-18 (1999): 2529-2545.

loss_one_minus_cd_auc Calculate Cumulative/Dynamic AUC loss

Description

This function subtracts the C/D AUC metric from one to obtain a loss function whose lower values indicate better model performance (useful for permutational feature importance)

Usage

loss_one_minus_cd_auc(y_true = NULL, risk = NULL, surv = NULL, times = NULL)

Arguments

y_true	a survival::Surv object containing the times and statuses of observations for which the metric will be evaluated
risk	ignored, left for compatibility with other metrics
surv	a matrix containing the predicted survival functions for the considered observa- tions, each row represents a single observation, whereas each column one time point
times	a vector of time points at which the survival function was evaluated

Value

a numeric vector of length equal to the length of the times vector, each value (from the range from 0 to 1) represents 1 - AUC metric at a specific time point, with lower values indicating better performance.

#' @section References:

- [1] Uno, Hajime, et al. "Evaluating prediction rules for t-year survivors with censored regression models." Journal of the American Statistical Association 102.478 (2007): 527-537.
- [2] Hung, Hung, and Chin-Tsang Chiang. "Optimal composite markers for time-dependent receiver operating characteristic curves with censored survival data." Scandinavian Journal of Statistics 37.4 (2010): 664-679.

See Also

cd_auc()

```
library(survival)
library(survex)
```

```
cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
cph_exp <- explain(cph)</pre>
```

```
y <- cph_exp$y
times <- cph_exp$times
surv <- cph_exp$predict_survival_function(cph, cph_exp$data, times)</pre>
```

```
loss_one_minus_cd_auc(y, surv = surv, times = times)
```

loss_one_minus_c_index

Calculate the Concordance index loss

Description

This function subtracts the C-index metric from one to obtain a loss function whose lower values indicate better model performance (useful for permutational feature importance)

Usage

```
loss_one_minus_c_index(y_true = NULL, risk = NULL, surv = NULL, times = NULL)
```

Arguments

y_true	a survival::Surv object containing the times and statuses of observations for which the metric will be evaluated
risk	a numeric vector of risk scores corresponding to each observation
surv	ignored, left for compatibility with other metrics
times	ignored, left for compatibility with other metrics

Value

numeric from 0 to 1, lower values indicate better performance

References

• [1] Harrell, F.E., Jr., et al. "Regression modelling strategies for improved prognostic prediction." Statistics in Medicine 3.2 (1984): 143-152.

See Also

c_index()

```
library(survival)
library(survex)
rotterdam <- survival::rotterdam
rotterdam$year <- NULL
cox_rotterdam_rec <- coxph(Surv(rtime, recur) ~ .,
    data = rotterdam,
    model = TRUE, x = TRUE, y = TRUE
)
coxph_explainer <- explain(cox_rotterdam_rec)</pre>
```

```
risk <- coxph_explainer$predict_function(coxph_explainer$model, coxph_explainer$data)
loss_one_minus_c_index(y_true = coxph_explainer$y, risk = risk)</pre>
```

loss_one_minus_integrated_cd_auc

Calculate integrated C/D AUC loss

Description

This function subtracts integrated the C/D AUC metric from one to obtain a loss function whose lower values indicate better model performance (useful for permutational feature importance)

Usage

```
loss_one_minus_integrated_cd_auc(
  y_true = NULL,
  risk = NULL,
  surv = NULL,
  times = NULL
)
```

Arguments

y_true	a survival::Surv object containing the times and statuses of observations for which the metric will be evaluated
risk	ignored, left for compatibility with other metrics
surv	a matrix containing the predicted survival functions for the considered observa- tions, each row represents a single observation, whereas each column one time point
times	a vector of time points at which the survival function was evaluated

Value

numeric from 0 to 1, lower values indicate better performance

#' @section References:

- [1] Uno, Hajime, et al. "Evaluating prediction rules for t-year survivors with censored regression models." Journal of the American Statistical Association 102.478 (2007): 527-537.
- [2] Hung, Hung, and Chin-Tsang Chiang. "Optimal composite markers for time-dependent receiver operating characteristic curves with censored survival data." Scandinavian Journal of Statistics 37.4 (2010): 664-679.

See Also

```
integrated_cd_auc() cd_auc() loss_one_minus_cd_auc()
```

Examples

```
library(survival)
library(survex)

cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
cph_exp <- explain(cph)

y <- cph_exp$y
times <- cph_exp$times
surv <- cph_exp$predict_survival_function(cph, cph_exp$data, times)

# calculating directly
loss_one_minus_integrated_cd_auc(y, surv = surv, times = times)</pre>
```

model_diagnostics Dataset Level Model Diagnostics

Description

This function calculates martingale and deviance residuals.

Usage

```
model_diagnostics(explainer)
```

```
## S3 method for class 'surv_explainer'
model_diagnostics(explainer)
```

Arguments

explainer an explainer object - model preprocessed by the explain() function

Value

An object of class c("model_diagnostics_survival"). It's a list with the explanations in the result element.

Examples

```
library(survival)
library(survex)
cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
rsf_ranger <- ranger::ranger(Surv(time, status) ~ .,
    data = veteran,
    respect.unordered.factors = TRUE,</pre>
```

model_parts

```
num.trees = 100,
mtry = 3,
max.depth = 5
)
cph_exp <- explain(cph)
rsf_ranger_exp <- explain(rsf_ranger,
    data = veteran[, -c(3, 4)],
    y = Surv(veteran$time, veteran$status)
)
cph_residuals <- model_diagnostics(cph_exp)
rsf_residuals <- model_diagnostics(rsf_ranger_exp)
head(cph_residuals$result)
plot(cph_residuals, rsf_residuals, xvariable = "age")
plot(cph_residuals, rsf_residuals, plot_type = "Cox-Snell")
```

model_parts

Dataset Level Variable Importance for Survival Models

Description

This function calculates variable importance as a change in the loss function after the variable values permutations.

Usage

```
model_parts(explainer, ...)
## S3 method for class 'surv_explainer'
model_parts(
    explainer,
    loss_function = survex::loss_brier_score,
    ...,
    type = "difference",
    output_type = "survival",
    N = 1000
)
```

Arguments

explainer	an explainer object - model preprocessed by the explain() function
	$Arguments\ passed\ on\ to\ surv_feature_importance,\ surv_integrated_feature_importance$
	B numeric, number of permutations to be calculated

	variables a character vector, names of variables to be included in the calcula- tion
	variable_groups a list of character vectors of names of explanatory variables. For each vector, a single variable-importance measure is computed for the joint effect of the variables which names are provided in the vector. By default, variable_groups = NULL, in which case variable-importance mea- sures are computed separately for all variables indicated in the variables argument
	label label of the model, if provides overrides x\$label
loss_function	a function that will be used to assess variable importance, by default loss_brier_score for survival models. The function can be supplied manually but has to have these named parameters (y_true, risk, surv, times), where y_true represents the survival::Surv object with observed times and statuses, risk is the risk score calculated by the model, and surv is the survival function for each observation evaluated at times.
type	a character vector, if "raw" the results are losses after the permutation, if "ratio" the results are in the form loss/loss_full_model and if "difference" the results are of the form loss - loss_full_model. Defaults to "difference".
output_type	either "survival" or "risk" the type of survival model output that should be used for explanations. If "survival" the explanations are based on the survival function. Otherwise the scalar risk predictions are used by the DALEX::model_profile function.
Ν	number of observations that should be sampled for calculation of variable impor- tance. If NULL then variable importance will be calculated on the whole dataset.

Details

Note: This function can be run within progressr::with_progress() to display a progress bar, as the execution can take long, especially on large datasets.

Value

An object of class c("model_parts_survival", "surv_feature_importance"). It's a list with the explanations in the result element.

```
library(survival)
library(survex)

cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
rsf_ranger <- ranger::ranger(Surv(time, status) ~ .,
    data = veteran,
    respect.unordered.factors = TRUE,
    num.trees = 100,
    mtry = 3,
    max.depth = 5
)</pre>
```

```
cph_exp <- explain(cph)

rsf_ranger_exp <- explain(rsf_ranger,
   data = veteran[, -c(3, 4)],
   y = Surv(veteran$time, veteran$status)
)

cph_model_parts_brier <- model_parts(cph_exp)
print(head(cph_model_parts_brier$result))
plot(cph_model_parts_brier)

rsf_ranger_model_parts <- model_parts(rsf_ranger_exp)
print(head(rsf_ranger_model_parts$result))
plot(cph_model_parts_brier, rsf_ranger_model_parts)</pre>
```

model_performance Dataset Level Performance Measures

Description

This function calculates metrics for survival models. The metrics calculated are C/D AUC, Brier score, and their integrated versions, as well as concordance index. It also can calculate ROC curves for specific selected time points.

Usage

```
model_performance(explainer, ...)
## S3 method for class 'surv_explainer'
model_performance(
    explainer,
    ...,
    type = "metrics",
    metrics = c(`C-index` = c_index, `Integrated C/D AUC` = integrated_cd_auc,
    `Brier score` = brier_score, `Integrated Brier score` = integrated_brier_score,
    `C/D AUC` = cd_auc),
    times = NULL
)
```

Arguments

explainer	an explainer object - model preprocessed by the explain() function
	other parameters, currently ignored
type	character, either "metrics" or "roc". If "metrics" then performance metrics
	are calculated, if "roc" ROC curves for selected time points are calculated.

metrics	a named vector containing the metrics to be calculated. The values should be standardized loss functions. The functions can be supplied manually but has to have these named parameters (y_true, risk, surv, times), where y_true represents the survival::Surv object with observed times and statuses, risk is the risk score calculated by the model, and surv is the survival function for each observation evaluated at times.
times	a numeric vector of times. If type == "metrics" then the survival function is evaluated at these times, if type == "roc" then the ROC curves are calculated at these times.

Value

An object of class "model_performance_survival". It's a list of metric values calculated for the model. It contains:

- Harrell's concordance index [1]
- Brier score [2, 3]
- C/D AUC using the estimator proposed by Uno et. al [4]
- integral of the Brier score
- integral of the C/D AUC

References

- [1] Harrell, F.E., Jr., et al. "Regression modelling strategies for improved prognostic prediction." Statistics in Medicine 3.2 (1984): 143-152.
- [2] Brier, Glenn W. "Verification of forecasts expressed in terms of probability." Monthly Weather Review 78.1 (1950): 1-3.
- [3] Graf, Erika, et al. "Assessment and comparison of prognostic classification schemes for survival data." Statistics in Medicine 18.17-18 (1999): 2529-2545.
- [4] Uno, Hajime, et al. "Evaluating prediction rules for t-year survivors with censored regression models." Journal of the American Statistical Association 102.478 (2007): 527-537.

```
library(survival)
library(survex)

cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
rsf_ranger <- ranger::ranger(Surv(time, status) ~ .,
    data = veteran,
    respect.unordered.factors = TRUE,
    num.trees = 100,
    mtry = 3,
    max.depth = 5
)</pre>
```

```
rsf_src <- randomForestSRC::rfsrc(Surv(time, status) ~ .,</pre>
    data = veteran
)
cph_exp <- explain(cph)</pre>
rsf_ranger_exp <- explain(rsf_ranger,</pre>
    data = veteran[, -c(3, 4)],
    y = Surv(veteran$time, veteran$status)
)
rsf_src_exp <- explain(rsf_src)</pre>
cph_model_performance <- model_performance(cph_exp)</pre>
rsf_ranger_model_performance <- model_performance(rsf_ranger_exp)</pre>
rsf_src_model_performance <- model_performance(rsf_src_exp)</pre>
print(cph_model_performance)
plot(rsf_ranger_model_performance, cph_model_performance,
    rsf_src_model_performance,
    metrics_type = "scalar"
)
plot(rsf_ranger_model_performance, cph_model_performance, rsf_src_model_performance)
cph_model_performance_roc <- model_performance(cph_exp, type = "roc", times = c(100, 250, 500))</pre>
plot(cph_model_performance_roc)
```

<pre>model_profile</pre>	Dataset Level Variable Profile as Partial Dependence Explanations for
	Survival Models

Description

This function calculates explanations on a dataset level that help explore model response as a function of selected variables. The explanations are calculated as an extension of Partial Dependence Profiles with the inclusion of the time dimension.

Usage

```
model_profile(
    explainer,
    variables = NULL,
    N = 100,
    ...,
    groups = NULL,
    k = NULL,
    type = "partial",
```

```
center = FALSE,
 output_type = "survival"
)
## S3 method for class 'surv_explainer'
model_profile(
 explainer,
 variables = NULL,
 N = 100,
  . . . ,
 categorical_variables = NULL,
  grid_points = 51,
 variable_splits_type = "uniform",
  groups = NULL,
  k = NULL,
  center = FALSE,
  type = "partial",
 output_type = "survival"
)
```

Arguments

explainer	an explainer object - model preprocessed by the explain() function
variables	character, a vector of names of variables to be explained
Ν	number of observations used for the calculation of aggregated profiles. By de- fault 100. If NULL all observations are used.
	<pre>other parameters passed to DALEX::model_profile if output_type == "risk", otherwise ignored</pre>
groups	<pre>if output_type == "risk" a variable name that will be used for grouping. By default NULL, so no groups are calculated. If output_type == "survival" then ignored</pre>
k	<pre>passed to DALEX::model_profile if output_type == "risk", otherwise ig- nored</pre>
type	<pre>the type of variable profile, "partial" for Partial Dependence, "accumulated" for Accumulated Local Effects, or "conditional" (available only for output_type == "risk")</pre>
center	logical, should profiles be centered around the average prediction
output_type	either "survival", "chf" or "risk" the type of survival model output that should be considered for explanations. If "survival" the explanations are based on the survival function. If "chf" the explanations are based on the cu- mulative hazard function. Otherwise the scalar risk predictions are used by the DALEX::predict_profile function.
categorical_var	iables
	character, a vector of names of additional variables which should be treated as categorical (factors are automatically treated as categorical variables). If it contains variable names not present in the variables argument, they will be added at the end.

grid_points maximum number of points for profile calculations. Note that the final number of points may be lower than grid_points. Will be passed to internal function. By default 51.

variable_splits_type

character, decides how variable grids should be calculated. Use "quantiles" for percentiles or "uniform" (default) to get uniform grid of points.

Value

An object of class model_profile_survival. It is a list with the element result containing the results of the calculation.

Examples

```
library(survival)
library(survex)
cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
rsf_src <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)</pre>
cph_exp <- explain(cph)</pre>
rsf_src_exp <- explain(rsf_src)</pre>
cph_model_profile <- model_profile(cph_exp,</pre>
    output_type = "survival",
    variables = c("age")
)
head(cph_model_profile$result)
plot(cph_model_profile)
rsf_model_profile <- model_profile(rsf_src_exp,</pre>
    output_type = "survival",
    variables = c("age", "celltype"),
    type = "accumulated"
)
head(rsf_model_profile$result)
plot(rsf_model_profile, variables = c("age", "celltype"), numerical_plot_type = "contours")
```

model_profile_2d Dataset Level 2-Dimensional Variable Profile for Survival Models

Description

This function calculates explanations on a dataset level that help explore model response as a function of selected pairs of variables. The explanations are calculated as an extension of Partial Dependence Profiles or Accumulated Local Effects with the inclusion of the time dimension.

Usage

```
model_profile_2d(
  explainer,
  variables = NULL,
 N = 100,
  categorical_variables = NULL,
  grid_points = 25,
  center = FALSE,
  variable_splits_type = "uniform",
  type = "partial",
  output_type = "survival"
)
## S3 method for class 'surv_explainer'
model_profile_2d(
  explainer,
  variables = NULL,
 N = 100,
  categorical_variables = NULL,
  grid_points = 25,
  center = FALSE,
  variable_splits_type = "uniform",
  type = "partial",
  output_type = "survival"
)
```

Arguments

explainer	an explainer object - model preprocessed by the explain() function	
variables	list of character vectors of length 2, names of pairs of variables to be explained	
Ν	number of observations used for the calculation of aggregated profiles. By default 100. If NULL all observations are used.	
categorical_variables		
	character, a vector of names of additional variables which should be treated as categorical (factors are automatically treated as categorical variables). If it contains variable names not present in the variables argument, they will be added at the end.	
grid_points	maximum number of points for profile calculations. Note that the final number of points may be lower than grid_points. Will be passed to internal function. By default 25.	
center	logical, should profiles be centered around the average prediction	

variable_splits_type	
	character, decides how variable grids should be calculated. Use "quantiles" for quantiles or "uniform" (default) to get uniform grid of points. Used only if type = "partial".
type	the type of variable profile, "partial" for Partial Dependence or "accumulated" for Accumulated Local Effects
output_type	either "survival", "chf" or "risk" the type of survival model output that should be considered for explanations. If "survival" the explanations are based on the survival function. If "chf" the explanations are based on the cu- mulative hazard function. Otherwise the scalar risk predictions are used by the DALEX::predict_profile function.

Value

An object of class model_profile_2d_survival. It is a list with the element result containing the results of the calculation.

Examples

```
library(survival)
library(survex)

cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
cph_exp <- explain(cph)

cph_model_profile_2d <- model_profile_2d(cph_exp,
    variables = list(c("age", "celltype"))
)
head(cph_model_profile_2d$result)
plot(cph_model_profile_2d]

cph_model_profile_2d_ale <- model_profile_2d(cph_exp,
    variables = list(c("age", "karno")),
    type = "accumulated"
)
head(cph_model_profile_2d_ale$result)
plot(cph_model_profile_2d_ale$result)
plot(cph_model_profile_2d_ale$result)</pre>
```

model_survshap Global SHAP Values

Description

This function computes global SHAP values.

Usage

```
model_survshap(explainer, ...)
## S3 method for class 'surv_explainer'
model_survshap(
    explainer,
    new_observation = NULL,
    y_true = NULL,
    N = NULL,
    calculation_method = "kernelshap",
    aggregation_method = "integral",
    output_type = "survival",
    ...
)
```

Arguments

explainer	an explainer object - model preprocessed by the explain() function	
	additional parameters, passed to internal functions	
new_observatio	n	
	new observations for which predictions need to be explained	
y_true	a two element numeric vector or matrix of one row and two columns, the first el- ement being the true observed time and the second the status of the observation, used for plotting	
Ν	a positive integer, number of observations used as the background data	
calculation_method		
	a character, either "kernelshap" for use of kernelshap library (providing faster Kernel SHAP with refinements), "exact_kernel" for exact Kernel SHAP estimation, or "treeshap" for use of treeshap library (efficient implementation to compute SHAP values for tree-based models).	
aggregation_method		
	a character, either "integral", "integral_absolute", "mean_absolute", "max_absolute", or "sum_of_squares"	
output_type	a character, either "survival" or "chf". Determines which type of prediction should be used for explanations.	

Details

If specifying y_true, also new_observation must be specified. Using the argument new_observation, global SHAP values are computed for the provided data. Otherwise, global SHAP values are computed for the data, the explainer was trained with.

Value

An object of class aggregated_surv_shap containing the computed global SHAP values.

Examples

```
veteran <- survival::veteran</pre>
rsf_ranger <- ranger::ranger(</pre>
    survival::Surv(time, status) ~ .,
    data = veteran,
    respect.unordered.factors = TRUE,
    num.trees = 100,
    mtry = 3,
    max.depth = 5
)
rsf_ranger_exp <- explain(</pre>
    rsf_ranger,
    data = veteran[, -c(3, 4)],
    y = survival::Surv(veteran$time, veteran$status),
    verbose = FALSE
)
ranger_global_survshap <- model_survshap(</pre>
    explainer = rsf_ranger_exp,
    new_observation = veteran[
        c(1:4, 17:20, 110:113, 126:129),
        !colnames(veteran) %in% c("time", "status")
    ],
    y_true = survival::Surv(
        veteran$time[c(1:4, 17:20, 110:113, 126:129)],
        veteran$status[c(1:4, 17:20, 110:113, 126:129)]
    ),
    aggregation_method = "integral",
    calculation_method = "kernelshap",
)
plot(ranger_global_survshap)
plot(ranger_global_survshap, geom = "beeswarm")
plot(ranger_global_survshap, geom = "profile", color_variable = "karno")
```

plot.aggregated_surv_shap Plot Aggregated SurvSHAP(t) Explanations for Survival Models

Description

This functions plots objects of class aggregated_surv_shap - aggregated time-dependent explanations of survival models created using the model_survshap() function.

Usage

S3 method for class 'aggregated_surv_shap'

```
plot(
    x,
    geom = "importance",
    ...,
    title = "default",
    subtitle = "default",
    max_vars = 7,
    colors = NULL
)
```

Arguments

an object of class aggregated_surv_shap to be plotted
character, one of "importance", "beeswarm", "profile" or "curves". Type of chart to be plotted; "importance" shows the importance of variables over time and aggregated, "beeswarm" shows the distribution of SurvSHAP(t) values for variables and observations, "profile" shows the dependence of SurvSHAP(t) values on variable values, "curves" shows all SurvSHAP(t) curves for selected variable colored by its value or with functional boxplot if boxplot = TRUE.
additional parameters passed to internal functions
character, title of the plot
character, subtitle of the plot, 'default' automatically generates "created for the XXX model ($n = YYY$)", where XXX is the explainer label and YYY is the number of observations used for calculations
maximum number of variables to be plotted (least important variables are ignored), by default 7
character vector containing the colors to be used for plotting variables (containing either hex codes "#FF69B4", or names "blue")

Value

An object of the class ggplot.

Plot options

plot.aggregated_surv_shap(geom = "importance"):

- rug character, one of "all", "events", "censors", "none" or NULL. Which times to mark on the x axis in geom_rug().
- rug_colors character vector containing two colors (containing either hex codes "#FF69B4", or names "blue"). The first color (red by default) will be used to mark event times, whereas the second (grey by default) will be used to mark censor times.
- xlab_left, ylab_right axis labels for left and right plots (due to different aggregation possibilities)

plot.aggregated_surv_shap(geom = "beeswarm"):

• no additional parameters

plot.aggregated_surv_shap(geom = "profile"):

- variable variable for which the profile is to be plotted, by default first from result data
- color_variable variable used to denote the color, by default equal to variable

plot.aggregated_surv_shap(geom = "curves"):

- variable variable for which SurvSHAP(t) curves are to be plotted, by default first from result data
- boxplot whether to plot functional boxplot with marked outliers or all curves colored by variable value
- coef length of the functional boxplot's whiskers as multiple of IQR, by default 1.5

```
veteran <- survival::veteran</pre>
rsf_ranger <- ranger::ranger(</pre>
    survival::Surv(time, status) ~ .,
   data = veteran,
    respect.unordered.factors = TRUE,
    num.trees = 100,
   mtry = 3,
   max.depth = 5
)
rsf_ranger_exp <- explain(</pre>
    rsf_ranger,
    data = veteran[, -c(3, 4)],
    y = survival::Surv(veteran$time, veteran$status),
    verbose = FALSE
)
ranger_global_survshap <- model_survshap(</pre>
    explainer = rsf_ranger_exp,
    new_observation = veteran[
        c(1:4, 17:20, 110:113, 126:129),
        !colnames(veteran) %in% c("time", "status")
   ],
    y_true = survival::Surv(
        veteran$time[c(1:4, 17:20, 110:113, 126:129)],
        veteran$status[c(1:4, 17:20, 110:113, 126:129)]
   ),
    aggregation_method = "integral",
    calculation_method = "kernelshap",
)
plot(ranger_global_survshap)
plot(ranger_global_survshap, geom = "beeswarm")
plot(ranger_global_survshap, geom = "profile"
     variable = "age", color_variable = "karno")
plot(ranger_global_survshap, geom = "curves",
     variable = "age")
plot(ranger_global_survshap, geom = "curves",
     variable = "age", boxplot = TRUE)
```

Description

This function plots objects of class "model_diagnostics_survival" created using the model_diagnostics() function.

Usage

```
## S3 method for class 'model_diagnostics_survival'
plot(
    x,
    ...,
    plot_type = "deviance",
    xvariable = "index",
    smooth = as.logical(xvariable != "index"),
    facet_ncol = NULL,
    title = "Model diagnostics",
    subtitle = "default",
    colors = NULL
)
```

Arguments

х	an object of class model_diagnostics_survival to be plotted
	$additional \ objects \ of \ class \ model_diagnostics_survival \ to \ be \ plotted \ together$
plot_type	character, either "deviance", "martingale" or "Cox-Snell". Selects the type of plot to be prepared. If "deviance" or "martingale then deviance/martingale residuals are plotted against xvariable. If "Cox-Snell" then diagnostic plot of Cox-Snell residuals is prepared, which is CHF estimated based on Cox-Snell residuals against theoretical cumulative hazard trajectory of the Exp(1) – diagonal line.
xvariable	character, name of the variable to be plotted on x-axis (can be name of the variable to be drawn on the x-axis (can be any column from the x\$result: explanatory variable, time, other residuals). By default "index" which gives the order of observations.
smooth	<pre>logical, shall the smooth line be added. Only used when plot_type = "deviance" or plot_type = "martingale".</pre>
facet_ncol	number of columns for arranging subplots
title	character, title of the plot

subtitle	character, subtitle of the plot, "default" automatically generates "created for XXX, YYY models", where XXX and YYY are the explainer labels
colors	character vector containing the colors to be used for plotting (containing either hex codes "#FF69B4", or names "blue").

Value

An object of the class ggplot.

plot.model_parts_survival

Plot Model Parts for Survival Models

Description

This function is a wrapper for plotting model_parts objects created for survival models using the model_parts() function.

Usage

S3 method for class 'model_parts_survival'
plot(x, ...)

Arguments

х	an object of class "model_parts_survival" to be plotted
	additional parameters passed to the $\verb"plot.surv_feature_importance"$ function

Value

An object of the class ggplot.

Plot options

- title character, title of the plot
- subtitle character, subtitle of the plot, if NULL automatically generated as "created for XXX, YYY models", where XXX and YYY are explainer labels
- max_vars maximum number of variables to be plotted (least important variables are ignored)
- colors character vector containing the colors to be used for plotting variables (containing either hex codes "#FF69B4", or names "blue")
- rug character, one of "all", "events", "censors", "none" or NULL. Which times to mark on the x axis in geom_rug().
- rug_colors character vector containing two colors (containing either hex codes "#FF69B4", or names "blue"). The first color (red by default) will be used to mark event times, whereas the second (grey by default) will be used to mark censor times.

See Also

Other functions for plotting 'model_parts_survival' objects: plot.surv_feature_importance()

Examples

```
library(survival)
library(survex)
model <- coxph(Surv(time, status) ~ ., data = veteran, x = TRUE, model = TRUE, y = TRUE)
explainer <- explain(model)
mp <- model_parts(explainer)
plot(mp)</pre>
```

Description

This function is a wrapper for plotting model_performance objects created for survival models using the model_performance() function.

Usage

```
## S3 method for class 'model_performance_survival'
plot(x, ...)
```

Arguments

х	an object of class "model_performance_survival" to be plotted
	additional parameters passed to the plot.surv_model_performance or plot.surv_model_performance function

Value

An object of the class ggplot.

Plot options

plot.surv_model_performance:

- x an object of class "surv_model_performance" to be plotted
- ... additional objects of class "surv_model_performance" to be plotted together
- metrics character, names of metrics to be plotted (subset of C/D AUC", "Brier score" for metrics_type %in% c("time_dependent", "functional") or subset of "C-index", "Integrated Brier score", "Integrated C/D AUC" for metrics_type == "scalar"), by default (NULL) all metrics of a given type are plotted
- metrics_type character, either one of c("time_dependent", "functional") for functional metrics or "scalar" for scalar metrics
- title character, title of the plot
- subtitle character, subtitle of the plot, 'default' automatically generates "created for XXX, YYY models", where XXX and YYY are the explainer labels
- facet_ncol number of columns for arranging subplots
- colors character vector containing the colors to be used for plotting variables (containing either hex codes "#FF69B4", or names "blue")
- rug character, one of "all", "events", "censors", "none" or NULL. Which times to mark on the x axis in geom_rug().
- rug_colors character vector containing two colors (containing either hex codes "#FF69B4", or names "blue"). The first color (red by default) will be used to mark event times, whereas the second (grey by default) will be used to mark censor times.

plot.surv_model_performance_rocs:

- x an object of class "surv_model_performance_rocs" to be plotted
- ... additional objects of class "surv_model_performance_rocs" to be plotted together
- title character, title of the plot
- subtitle character, subtitle of the plot, 'default' automatically generates "created for XXX, YYY models", where XXX and YYY are the explainer labels
- colors character vector containing the colors to be used for plotting variables (containing either hex codes "#FF69B4", or names "blue")
- facet_ncol number of columns for arranging subplots

See Also

Other functions for plotting 'model_performance_survival' objects: plot.surv_model_performance_rocs(), plot.surv_model_performance()

```
library(survival)
library(survex)
model <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)
exp <- explain(model)
m_perf <- model_performance(exp)
plot(m_perf, metrics_type = "functional")
m_perf_roc <- model_performance(exp, type = "roc", times = c(100, 300))
plot(m_perf_roc)</pre>
```

plot.model_profile_2d_survival

Plot 2-Dimensional Model Profile for Survival Models

Description

This function plots objects of class "model_profile_2d_survival" created using the model_profile_2d() function.

Usage

```
## S3 method for class 'model_profile_2d_survival'
plot(
    x,
    ...,
    variables = NULL,
    times = NULL,
    marginalize_over_time = FALSE,
    facet_ncol = NULL,
    title = "default",
    subtitle = "default",
    colors = NULL
)
```

Arguments

х	an object of class model_profile_2d_survival to be plotted	
	additional objects of class model_profile_2d_survival to be plotted together	
variables	list of character vectors of length 2, names of pairs of variables to be plotted	
times	numeric vector, times for which the profile should be plotted, the times must be present in the 'times' field of the explainer. If NULL (default) then the median survival time (if available) or the median time from the explainer object is used.	
marginalize_over_time		
	logical, if TRUE then the profile is calculated for all times and then averaged over time, if FALSE (default) then the profile is calculated for each time separately	
facet_ncol	number of columns for arranging subplots	
title	character, title of the plot. 'default' automatically generates either "2D partial dependence survival profiles" or "2D accumulated local effects survival profiles" depending on the explanation type.	
subtitle	character, subtitle of the plot, 'default' automatically generates "created for the XXX model", where XXX is the explainer labels, if marginalize_over_time = FALSE, time is also added to the subtitle	
colors	character vector containing the colors to be used for plotting variables (contain- ing either hex codes "#FF69B4", or names "blue")	

Value

A collection of ggplot objects arranged with the patchwork package.

Examples

```
library(survival)
library(survex)
cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
cph_exp <- explain(cph)</pre>
cph_model_profile_2d <- model_profile_2d(cph_exp,</pre>
    variables = list(
        c("age", "celltype"),
        c("age", "karno")
    )
)
head(cph_model_profile_2d$result)
plot(cph_model_profile_2d, variables = list(c("age", "celltype")), times = cph_exp$times[20])
cph_model_profile_2d_ale <- model_profile_2d(cph_exp,</pre>
    variables = list(c("age", "karno")),
    type = "accumulated"
)
head(cph_model_profile_2d_ale$result)
plot(cph_model_profile_2d_ale, times = cph_exp$times[c(10, 20)], marginalize_over_time = TRUE)
```

Description

This function plots objects of class "model_profile_survival" created using the model_profile() function.

Usage

```
## S3 method for class 'model_profile_survival'
plot(
    x,
    ...,
    geom = "time",
    variables = NULL,
    variable_type = NULL,
    facet_ncol = NULL,
```

```
numerical_plot_type = "lines",
times = NULL,
marginalize_over_time = FALSE,
plot_type = NULL,
title = "default",
subtitle = "default",
colors = NULL,
rug = "all",
rug_colors = c("#dd0000", "#222222")
)
```

Arguments

х	an object of class model_profile_survival to be plotted
	additional objects of class model_profile_survival to be plotted together. Only available for geom = "time".
geom	character, either "time" or "variable". Selects the type of plot to be prepared. If "time" then the x-axis represents survival times, and variable is denoted by colors, if "variable" then the x-axis represents the variable values, and y-axis represents the predictions at selected time points.
variables	character, names of the variables to be plotted. When geom = "variable" it needs to be a name of a single variable, when geom = "time" it can be a vector of variable names. If NULL (default) then first variable (for geom = "variable") or all variables (for geom = "time") are plotted.
variable_type	character, either "numerical", "categorical" or NULL (default), select only one type of variable for plotting, or leave NULL for all. Only used when geom = "time".
facet_ncol	number of columns for arranging subplots. Only used when geom = "time".
<pre>numerical_plot_</pre>	
	character, either "lines", or "contours" selects the type of numerical variable plots. Only used when geom = "time".
times	numeric vector, times for which the profile should be plotted, the times must be present in the 'times' field of the explainer. If NULL (default) then the median survival time (if available) or the median time from the explainer object is used. Only used when geom = "variable" and marginalize_over_time = FALSE.
marginalize_ove	r_time
	logical, if TRUE then the profile is calculated for all times and then averaged over time, if FALSE (default) then the profile is calculated for each time separately. Only used when geom = "variable".
plot_type	character, one of "pdp", "ice", "pdp+ice", or NULL (default). If NULL then the type of plot is chosen automatically based on the number of variables to be plotted. Only used when geom = "variable".
title	character, title of the plot
subtitle	character, subtitle of the plot, "default" automatically generates "created for XXX, YYY models", where XXX and YYY are the explainer labels

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colors	character vector containing the colors to be used for plotting variables (contain- ing either hex codes "#FF69B4", or names "blue").
rug	character, one of "all", "events", "censors", "none" or NULL. Which times to mark on the x axis in geom_rug(). Only used when geom = "time".
rug_colors	character vector containing two colors (containing either hex codes "#FF69B4", or names "blue"). The first color (red by default) will be used to mark event times, whereas the second (grey by default) will be used to mark censor times.

Value

A collection of ggplot objects arranged with the patchwork package.

Examples

```
library(survival)
library(survex)

model <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)
exp <- explain(model)

m_prof <- model_profile(exp, categorical_variables = "trt")
plot(m_prof)
plot(m_prof, numerical_plot_type = "contours")
plot(m_prof, variables = c("trt", "age"), facet_ncol = 1)
plot(m_prof, geom = "variable", variables = "karno", plot_type = "pdp+ice")
plot(m_prof, geom = "variable", times = exp$times[c(5, 10)],
    variables = "trt", plot_type = "pdp+ice")</pre>
```

plot.predict_parts_survival

Plot Predict Parts for Survival Models

Description

This function plots objects of class "predict_parts_survival" - local explanations for survival models created using the predict_parts() function.

Usage

```
## S3 method for class 'predict_parts_survival'
plot(x, ...)
```

Arguments

х	an object of class "predict_parts_survival" to be plotted
	additional parameters passed to the ${\tt plot.surv_shap}$ or ${\tt plot.surv_lime}$ functions

Value

An object of the class ggplot.

Plot options

plot.surv_shap:

- x an object of class "surv_shap" to be plotted
- ... additional objects of class surv_shap to be plotted together
- title character, title of the plot
- subtitle character, subtitle of the plot, 'default' automatically generates "created for XXX, YYY models", where XXX and YYY are the explainer labels
- max_vars maximum number of variables to be plotted (least important variables are ignored)
- colors character vector containing the colors to be used for plotting variables (containing either hex codes "#FF69B4", or names "blue")
- rug character, one of "all", "events", "censors", "none" or NULL. Which times to mark on the x axis in geom_rug().
- rug_colors character vector containing two colors (containing either hex codes "#FF69B4", or names "blue"). The first color (red by default) will be used to mark event times, whereas the second (grey by default) will be used to mark censor times.

plot.surv_lime:

- x an object of class "surv_lime" to be plotted
- type character, either "coefficients" or "local_importance", selects the type of plot
- show_survival_function logical, if the survival function of the explanations should be plotted next to the barplot
- ... other parameters currently ignored
- title character, title of the plot
- subtitle character, subtitle of the plot, 'default' automatically generates "created for XXX, YYY models", where XXX and YYY are the explainer labels
- max_vars maximum number of variables to be plotted (least important variables are ignored)
- colors character vector containing the colors to be used for plotting variables (containing either hex codes "#FF69B4", or names "blue")

See Also

Other functions for plotting 'predict_parts_survival' objects: plot.surv_lime(), plot.surv_shap()

Examples

```
library(survival)
library(survex)
model <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)
exp <- explain(model)
p_parts_shap <- predict_parts(exp, veteran[1, -c(3, 4)], type = "survshap")
plot(p_parts_shap)
p_parts_lime <- predict_parts(exp, veteran[1, -c(3, 4)], type = "survlime")
plot(p_parts_lime)</pre>
```

Description

This function plots objects of class "predict_profile_survival" created using the predict_profile() function.

Usage

```
## S3 method for class 'predict_profile_survival'
plot(
  х,
  ...,
  geom = "time"
  variables = NULL,
  variable_type = NULL,
  facet_ncol = NULL,
  numerical_plot_type = "lines",
  times = NULL,
  marginalize_over_time = FALSE,
  title = "default",
  subtitle = "default",
  colors = NULL,
  rug = "all",
  rug_colors = c("#dd0000", "#222222")
)
```

Arguments

х	an object of class predict_profile_survival to be plotted
	additional objects of class "predict_profile_survival" to be plotted together. Only available for geom = "time".
geom	character, either "time" or "variable". Selects the type of plot to be prepared. If "time" then the x-axis represents survival times, and variable is denoted by colors, if "variable" then the x-axis represents the variable values, and y-axis represents the predictions at selected time points.
variables	character, names of the variables to be plotted. When geom = "variable" it needs to be a name of a single variable, when geom = "time" it can be a vector of variable names. If NULL (default) then first variable (for geom = "variable") or all variables (for geom = "time") are plotted.
variable_type	character, either "numerical", "categorical" or NULL (default), select only one type of variable for plotting, or leave NULL for all. Only used when geom = "time".
facet_ncol	number of columns for arranging subplots. Only used when geom = "time".
numerical_plot	
	character, either "lines", or "contours" selects the type of numerical variable plots. Only used when geom = "time".
times	numeric vector, times for which the profile should be plotted, the times must be present in the 'times' field of the explainer. If NULL (default) then the median survival time (if available) or the median time from the explainer object is used. Only used when geom = "variable" and marginalize_over_time = FALSE.
marginalize_ov	
	logical, if TRUE then the profile is calculated for all times and then averaged over time, if FALSE (default) then the profile is calculated for each time separately. Only used when geom = "variable".
title	character, title of the plot
subtitle	character, subtitle of the plot, 'default' automatically generates "created for XXX, YYY models", where XXX and YYY are the explainer labels
colors	character vector containing the colors to be used for plotting variables (contain- ing either hex codes "#FF69B4", or names "blue")
rug	character, one of "all", "events", "censors", "none" or NULL. Which times to mark on the x axis in geom_rug().
rug_colors	character vector containing two colors (containing either hex codes "#FF69B4", or names "blue"). The first color (red by default) will be used to mark event times, whereas the second (grey by default) will be used to mark censor times.

Value

A collection of ggplot objects arranged with the patchwork package.

plot.surv_feature_importance

Examples

```
library(survival)
library(survex)

model <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)
exp <- explain(model)

p_profile <- predict_profile(exp, veteran[1, -c(3, 4)])

plot(p_profile)

p_profile_with_cat <- predict_profile(
    exp,
    veteran[1, -c(3, 4)],
    categorical_variables = c("trt", "prior")
)

plot(p_profile_with_cat)</pre>
```

plot.surv_feature_importance

Plot Permutational Feature Importance for Survival Models

Description

This function plots feature importance objects created for survival models using the model_parts() function with a time-dependent metric, that is loss_one_minus_cd_auc() or loss_brier_score().

Usage

```
## S3 method for class 'surv_feature_importance'
plot(
    x,
    ...,
    title = "Time-dependent feature importance",
    subtitle = "default",
    max_vars = 7,
    colors = NULL,
    rug = "all",
    rug_colors = c("#dd0000", "#222222")
)
```

Arguments

х	an object of class "surv_feature_importance" to be plotted
•••	additional objects of class "surv_feature_importance" to be plotted together
title	character, title of the plot
subtitle	character, subtitle of the plot, 'default' automatically generates "created for XXX, YYY models", where XXX and YYY are the explainer labels
max_vars	maximum number of variables to be plotted (least important variables are ignored)
colors	character vector containing the colors to be used for plotting variables (containing either hex codes "#FF69B4", or names "blue")
rug	character, one of "all", "events", "censors", "none" or NULL. Which times to mark on the x axis in geom_rug().
rug_colors	character vector containing two colors (containing either hex codes "#FF69B4", or names "blue"). The first color (red by default) will be used to mark event times, whereas the second (grey by default) will be used to mark censor times.

Value

An object of the class ggplot.

See Also

Other functions for plotting 'model_parts_survival' objects: plot.model_parts_survival()

```
library(survival)
library(survex)
model <- coxph(Surv(time, status) ~ ., data = veteran, x = TRUE, model = TRUE, y = TRUE)
model_rf <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)
explainer <- explain(model)
explainer_rf <- explain(model_rf)
mp <- model_parts(explainer)
mp_rf <- model_parts(explainer_rf)
plot(mp, mp_rf)</pre>
```

plot.surv_lime

Description

This functions plots objects of class surv_lime - LIME explanations of survival models created using predict_parts(..., type="survlime") function.

Usage

```
## S3 method for class 'surv_lime'
plot(
    x,
    type = "local_importance",
    show_survival_function = TRUE,
    ...,
    title = "SurvLIME",
    subtitle = "default",
    max_vars = 7,
    colors = NULL
)
```

Arguments

х	an object of class "surv_lime" to be plotted
type	character, either "coefficients" or "local_importance" (default), selects the type of plot
show_survival_f	iunction
	logical, if the survival function of the explanations should be plotted next to the barplot
	other parameters currently ignored
title	character, title of the plot
subtitle	character, subtitle of the plot, 'default' automatically generates "created for XXX, YYY models", where XXX and YYY are the explainer labels
max_vars	maximum number of variables to be plotted (least important variables are ignored)
colors	character vector containing the colors to be used for plotting variables (containing either hex codes "#FF69B4", or names "blue")

Value

An object of the class ggplot.

See Also

Other functions for plotting 'predict_parts_survival' objects: plot.predict_parts_survival(), plot.surv_shap()

Examples

```
library(survival)
library(survex)
model <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)
exp <- explain(model)
p_parts_lime <- predict_parts(exp, veteran[1, -c(3, 4)], type = "survlime")
plot(p_parts_lime)</pre>
```

plot.surv_model_performance

Plot Model Performance Metrics for Survival Models

Description

This function plots objects of class "surv_model_performance" - visualization of metrics of different models created using the model_performance(..., type="metrics") function.

Usage

```
## S3 method for class 'surv_model_performance'
plot(
    x,
    ...,
    metrics = NULL,
    metrics_type = "time_dependent",
    title = "Model performance",
    subtitle = "default",
    facet_ncol = NULL,
    colors = NULL,
    rug = "all",
    rug_colors = c("#dd0000", "#222222")
)
```

Arguments

х	an object of class "surv_model_performance" to be plotted
	additional objects of class "surv_model_performance" to be plotted together

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metrics	<pre>character, names of metrics to be plotted (subset of C/D AUC", "Brier score" for metrics_type %in% c("time_dependent", "functional") or subset of "C- index","Integrated Brier score", "Integrated C/D AUC" for metrics_type == "scalar"), by default (NULL) all metrics of a given type are plotted</pre>
metrics_type	character, either one of c("time_dependent","functional") for functional metrics or "scalar" for scalar metrics
title	character, title of the plot
subtitle	character, subtitle of the plot, 'default' automatically generates "created for XXX, YYY models", where XXX and YYY are the explainer labels
facet_ncol	number of columns for arranging subplots
colors	character vector containing the colors to be used for plotting variables (containing either hex codes "#FF69B4", or names "blue")
rug	character, one of "all", "events", "censors", "none" or NULL. Which times to mark on the x axis in geom_rug().
rug_colors	character vector containing two colors (containing either hex codes "#FF69B4", or names "blue"). The first color (red by default) will be used to mark event times, whereas the second (grey by default) will be used to mark censor times.

Value

An object of the class ggplot.

See Also

Other functions for plotting 'model_performance_survival' objects: plot.model_performance_survival(), plot.surv_model_performance_rocs()

```
library(survival)
library(survex)
```

```
model <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)
exp <- explain(model)</pre>
```

```
m_perf <- model_performance(exp)
plot(m_perf)</pre>
```

plot.surv_model_performance_rocs

Plot ROC Curves for Survival Models

Description

This function plots objects of class "surv_model_performance_rocs" - ROC curves for specific time points for survival models created using the model_performance(..., type="roc").

Usage

```
## S3 method for class 'surv_model_performance_rocs'
plot(
    x,
    ...,
    title = "ROC curves for selected time points",
    subtitle = "default",
    auc = TRUE,
    colors = NULL,
    facet_ncol = NULL
)
```

Arguments

х	an object of class "surv_model_performance_rocs" to be plotted
	additional objects of class "surv_model_performance_rocs" to be plotted to- gether
title	character, title of the plot
subtitle	character, subtitle of the plot, 'default' automatically generates "created for XXX, YYY models", where XXX and YYY are the explainer labels
auc	boolean, whether the AUC values should be plotted
colors	character vector containing the colors to be used for plotting variables (contain- ing either hex codes "#FF69B4", or names "blue")
facet_ncol	number of columns for arranging subplots

Value

An object of the class ggplot.

See Also

Other functions for plotting 'model_performance_survival' objects: plot.model_performance_survival(), plot.surv_model_performance()

plot.surv_shap

Examples

```
library(survival)
library(survex)
model <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)
exp <- explain(model)
m_perf_roc <- model_performance(exp, type = "roc", times = c(100, 300))
plot(m_perf_roc)</pre>
```

plot.surv_shap Plot SurvSHAP(t) Explanations for Survival Models

Description

This functions plots objects of class surv_shap - time-dependent explanations of survival models created using the predict_parts(..., type="survshap") function.

Usage

```
## S3 method for class 'surv_shap'
plot(
    x,
    ...,
    title = "SurvSHAP(t)",
    subtitle = "default",
    max_vars = 7,
    colors = NULL,
    rug = "all",
    rug_colors = c("#dd0000", "#222222")
)
```

Arguments

х	an object of class surv_shap to be plotted
	additional objects of class surv_shap to be plotted together
title	character, title of the plot
subtitle	character, subtitle of the plot, 'default' automatically generates "created for XXX, YYY models", where XXX and YYY are the explainer labels
max_vars	maximum number of variables to be plotted (least important variables are ignored)
colors	character vector containing the colors to be used for plotting variables (contain- ing either hex codes "#FF69B4", or names "blue")

rug	character, one of "all", "events", "censors", "none" or NULL. Which times to mark on the x axis in geom_rug().
rug_colors	character vector containing two colors (containing either hex codes "#FF69B4", or names "blue"). The first color (red by default) will be used to mark event times, whereas the second (grey by default) will be used to mark censor times.

Value

An object of the class ggplot.

See Also

Other functions for plotting 'predict_parts_survival' objects: plot.predict_parts_survival(), plot.surv_lime()

Examples

```
library(survival)
library(survex)
model <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)
exp <- explain(model)
p_parts_shap <- predict_parts(exp, veteran[1, -c(3, 4)], type = "survshap")
plot(p_parts_shap)</pre>
```

predict.surv_explainer

Model Predictions for Survival Models

Description

This function allows for calculating model prediction in a unified way.

Usage

```
## S3 method for class 'surv_explainer'
predict(object, newdata = NULL, output_type = "survival", times = NULL, ...)
```

Arguments

object	an explainer object - model preprocessed by the explain() function
newdata	data used for the prediction
output_type	character, either "risk", "survival" or "chf" depending on the desired output

predict_parts

times	a numeric vector of times for the survival and cumulative hazard function predic-
	tions to be evaluated at. If "output_type == "risk" this argument is ignored,
	if left NULL then it is extracted from object\$times.
	other arguments, currently ignored

Value

A vector or matrix containing the prediction.

Examples

```
library(survival)
library(survex)
cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
rsf_ranger <- ranger::ranger(Surv(time, status) ~ .,</pre>
    data = veteran,
    respect.unordered.factors = TRUE,
    num.trees = 100,
    mtry = 3,
    max.depth = 5
)
cph_exp <- explain(cph)</pre>
rsf_ranger_exp <- explain(rsf_ranger,</pre>
    data = veteran[, -c(3, 4)],
    y = Surv(veteran$time, veteran$status)
)
predict(cph_exp, veteran[1, ], output_type = "survival")[, 1:10]
predict(cph_exp, veteran[1, ], output_type = "risk")
predict(rsf_ranger_exp, veteran[1, ], output_type = "chf")[, 1:10]
```

predict_parts

Instance Level Parts of Survival Model Predictions

Description

This function decomposes the model prediction into individual parts, which are attributions of particular variables. The explanations can be made via the SurvLIME and SurvSHAP(t) methods.

Usage

```
predict_parts(explainer, ...)
## S3 method for class 'surv_explainer'
predict_parts(
    explainer,
    new_observation,
    ...,
    N = NULL,
    type = "survshap",
    output_type = "survival",
    explanation_label = NULL
)
```

Arguments

explainer	an explainer object - model preprocessed by the explain() function
	other parameters which are passed to iBreakDown::break_down if output_type=="risk", or if output_type=="survival" to surv_shap() or surv_lime() functions depending on the selected type
new_observatio	n
	a new observation for which prediction need to be explained
Ν	the number of observations used for calculation of attributions. If NULL (de- fault) all explainer data will be used for SurvSHAP(t) and 100 neigbours for SurvLIME.
type	<pre>if output_type == "survival" must be either "survshap" or "survlime", otherwise refer to the DALEX::predict_parts</pre>
output_type	either "survival", "chf" or "risk" the type of survival model output that should be considered for explanations. If "survival" the explanations are based on the survival function. If "chf" the explanations are based on the cu- mulative hazard function. Otherwise the scalar risk predictions are used by the DALEX::predict_parts function.
explanation_label	
	a label that can overwrite explainer label (useful for multiple explanations for the same explainer/model)

Value

An object of class "predict_parts_survival" and additional classes depending on the type of explanations. It is a list with the element result containing the results of the calculation.

Additional parameters

There are additional parameters that are passed to internal functions

- for survlime
 - N a positive integer, number of observations generated in the neighbourhood

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- distance_metric character, name of the distance metric to be used, only "euclidean" is implemented
- kernel_width a numeric, parameter used for calculating weights, by default it's sqrt(ncol(data)*0.75)
- sampling_method character, name of the method of generating neighbourhood, only "gaussian" is implemented
- sample_around_instance logical, if the neighbourhood should be generated with the new observation as the center (default), or should the mean of the whole dataset be used as the center
- max_iter a numeric, maximal number of iteration for the optimization problem
- categorical_variables character vector, names of variables that should be treated as categories (factors are included by default)
- k a small positive number > 1, added to chf before taking log, so that weigths aren't negative
- for survshap
 - y_true a two element numeric vector or matrix of one row and two columns, the first element being the true observed time and the second the status of the observation, used for plotting
 - calculation_method a character, either "kernelshap" for use of kernelshap library (providing faster Kernel SHAP with refinements) or "exact_kernel" for exact Kernel SHAP estimation
 - aggregation_method a character, either "mean_absolute" or "integral", "max_absolute", "sum_of_squares"

References

- [1] Krzyziński, Mateusz, et al. "SurvSHAP(t): Time-dependent explanations of machine learning survival models." Knowledge-Based Systems 262 (2023): 110234
- [2] Kovalev, Maxim S., et al. "SurvLIME: A method for explaining machine learning survival models." Knowledge-Based Systems 203 (2020): 106164.

```
library(survival)
library(survex)

cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
cph_exp <- explain(cph)

cph_predict_parts_survshap <- predict_parts(cph_exp, new_observation = veteran[1, -c(3, 4)])
head(cph_predict_parts_survshap$result)
plot(cph_predict_parts_survshap)

cph_predict_parts_survlime <- predict_parts(
    cph_exp,
    new_observation = veteran[1, -c(3, 4)],
    type = "survlime"
)</pre>
```

```
head(cph_predict_parts_survlime$result)
plot(cph_predict_parts_survlime, type = "local_importance")
```

predict_profile Instance Level Profile as Ceteris Paribus for Survival Models

Description

This function calculates Ceteris Paribus Profiles for a specific observation with the possibility to take the time dimension into account.

Usage

```
predict_profile(
  explainer,
  new_observation,
  variables = NULL,
  categorical_variables = NULL,
  . . . .
  type = "ceteris_paribus",
  output_type = "survival",
  variable_splits_type = "uniform",
  center = FALSE
)
## S3 method for class 'surv_explainer'
predict_profile(
  explainer,
  new_observation,
  variables = NULL,
  categorical_variables = NULL,
  . . . ,
  type = "ceteris_paribus",
  output_type = "survival",
  variable_splits_type = "uniform",
  center = FALSE
)
```

Arguments

explaineran explainer object - model preprocessed by the explain() functionnew_observationa new observation for which the prediction need to be explainedvariablesa character vector containing names of variables to be explained

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categorical_variables a character vector of names of additional variables which should be treated as categorical (factors are automatically treated as categorical variables). If it contains variable names not present in the variables argument, they will be added at the end. additional parameters passed to DALEX::predict_profile if output_type =="risk" . . . character, only "ceteris_paribus" is implemented type either "survival", "chf" or "risk" the type of survival model output that output_type should be considered for explanations. If "survival" the explanations are based on the survival function. If "chf" the explanations are based on the cumulative hazard function. Otherwise the scalar risk predictions are used by the DALEX::predict_profile function. variable_splits_type character, decides how variable grids should be calculated. Use "quantiles" for percentiles or "uniform" (default) to get uniform grid of points. center logical, should profiles be centered around the average prediction

Value

An object of class c("predict_profile_survival", "surv_ceteris_paribus"). It is a list with the final result in the result element.

```
library(survival)
library(survex)

cph <- coxph(Surv(time, status) ~ ., data = veteran, model = TRUE, x = TRUE, y = TRUE)
rsf_src <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)

cph_exp <- explain(cph)
rsf_src_exp <- explain(rsf_src)

cph_predict_profile <- predict_profile(cph_exp, veteran[2, -c(3, 4)],
    variables = c("trt", "celltype", "karno", "age"),
    categorical_variables = "trt"
)
plot(cph_predict_profile, facet_ncol = 2)</pre>
```

```
rsf_predict_profile <- predict_profile(rsf_src_exp, veteran[5, -c(3, 4)], variables = "karno")
plot(cph_predict_profile, numerical_plot_type = "contours")</pre>
```

risk_from_chf

Description

Some models do not come with a ready to use risk prediction. This function allows for its generation based on the cumulative hazard function.

Usage

```
risk_from_chf(predict_cumulative_hazard_function, times)
```

Arguments

predict_cumulative_hazard_function	
	a function of three arguments (model, newdata, times) that allows for making cumulative hazard predictions.
times	a numeric vector of times at which the function should be evaluated.

Value

A function of two arguments (model, newdata) returning a vector of risks.

```
library(survex)
library(survival)

rsf_src <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)

chf_function <- transform_to_stepfunction(predict,
    type = "chf",
    prediction_element = "chf",
    times_element = "time.interest"
)
risk_function <- risk_from_chf(chf_function, unique(veteran$time))

explainer <- explain(rsf_src,
    predict_cumulative_hazard_function = chf_function,
    predict_function = risk_function</pre>
```

set_theme_survex Default Theme for survex plots

Description

Default Theme for survex plots

Usage

```
set_theme_survex(
   default_theme = "drwhy",
   default_theme_vertical = default_theme
)
```

theme_default_survex()

theme_vertical_default_survex()

Arguments

<pre>default_theme</pre>	object - string ("drwhy" or "ema") or an object of ggplot theme class.	Will be
	applied by default by survex to all horizontal plots	
default_theme_v	vertical	

object - string ("drwhy" or "ema") or an object of ggplot theme class. Will be applied by default by survex to all vertical plots

Value

list with current default themes

```
old <- set_theme_survex("ema")
library(survival)
library(survex)
model <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)
exp <- explain(model)
p_parts_lime <- predict_parts(exp, veteran[1, -c(3, 4)], type = "survlime")
old <- set_theme_survex("drwhy")
plot(p_parts_lime)
old <- set_theme_survex(ggplot2::theme_void(), ggplot2::theme_void())
plot(p_parts_lime)</pre>
```

survival_to_cumulative_hazard

Transform Survival to Cumulative Hazard

Description

Helper function to transform between survival function and CHF

Usage

```
survival_to_cumulative_hazard(survival_functions, epsilon = 0)
```

Arguments

survival_functions	
	matrix or vector, with each row representing a survival function
epsilon	a positive numeric number to add, so that the logarithm can be taken

Value

A matrix or vector transformed to the form of a cumulative hazard function.

Examples

library(survex)
vec <- c(1, 0.9, 0.8, 0.7, 0.6)
matr <- matrix(c(1, 0.9, 0.8, 1, 0.8, 0.6), ncol = 3)
survival_to_cumulative_hazard(vec)
survival_to_cumulative_hazard(matr)</pre>

surv_model_info Extract additional information from the model

Description

This generic function let user extract base information about model. The function returns a named list of class model_info that contain information about package of model, version and task type. For wrappers like mlr or parsnip both, package and wrapper information are stored

surv_model_info

Usage

```
surv_model_info(model, ...)
## S3 method for class 'coxph'
surv_model_info(model, ...)
## S3 method for class 'rfsrc'
surv_model_info(model, ...)
## S3 method for class 'ranger'
surv_model_info(model, ...)
## S3 method for class 'model_fit'
surv_model_info(model, ...)
## S3 method for class 'cph'
surv_model_info(model, ...)
## S3 method for class 'LearnerSurv'
surv_model_info(model, ...)
## S3 method for class 'sksurv'
surv_model_info(model, ...)
## S3 method for class 'flexsurvreg'
surv_model_info(model, ...)
## Default S3 method:
surv_model_info(model, ...)
```

Arguments

model	 model object
	• other arguments

Details

Currently supported packages are:

- class coxph Cox proportional hazards regression model created with survival package
- class model_fit models created with parsnip package
- class ranger random survival forest models created with ranger package
- class rfsrc random forest models created with randomForestSRC package

Value

A named list of class model_info

Examples

```
library(survival)
library(survex)
cph <- survival::coxph(survival::Surv(time, status) ~ .,
    data = veteran,
    model = TRUE, x = TRUE, y = TRUE
)
surv_model_info(cph)
library(ranger)
rsf_ranger <- ranger::ranger(survival::Surv(time, status) ~ .,
    data = veteran,
    num.trees = 50, mtry = 3, max.depth = 5
)
surv_model_info(rsf_ranger)</pre>
```

transform_to_stepfunction Transform Fixed Point Prediction into a Stepfunction

Description

Some models return the survival function or cumulative hazard function prediction at the times of events present in the training data set. This is a convenient utility to allow the prediction to be evaluated at any time.

Usage

```
transform_to_stepfunction(
    predict_function,
    eval_times = NULL,
    ...,
    type = NULL,
    prediction_element = NULL,
    times_element = NULL
)
```

Arguments

predict_function

a function making the prediction based on model and newdata arguments, the ... parameter is also passed to this function. It has to return either a numeric vector of the same length as eval_times, a matrix with this number of columns and the same number of rows as nrow(newdata). It can also return a list, with one of the elements containing such an object.

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eval_times	a numeric vector of times, at which the fixed predictions are made. This can be NULL, if predict_function returns a list which contains such a vector.
	other parameters passed to predict_function
type	the type of function to be returned, either "survival", "chf" or NULL this chooses the value of the step function before the first prediction time. If "survival" then it is 1, if "chf" then 0, otherwise, it is the value of the prediction for the first time in numerical order.
prediction_element	
	if predict_function returns a list with the matrix as one of its elements, this parameter should contain the name of this element
times_element	if predict_function returns a list with the matrix as one of its elements, this parameter should contain the name of this element

Value

The function returns a function with three arguments, (model, newdata, times), ready to supply it to an explainer.

```
library(survex)
library(survival)

rsf_src <- randomForestSRC::rfsrc(Surv(time, status) ~ ., data = veteran)

chf_function <- transform_to_stepfunction(predict,
    type = "chf",
    prediction_element = "chf",
    times_element = "time.interest"
)

explainer <- explain(rsf_src, predict_cumulative_hazard_function = chf_function)</pre>
```

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