

Package ‘triageR’

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Title Automated Machine Learning and AI Agent Tools for Clinical Prediction Modelling

Version 0.1.0

Description Provides a streamlined workflow for building, validating, and reporting clinical prediction models. Combines standard machine learning tools with an optional AI agent that recommends appropriate statistical methods, runs sensitivity analyses, and flags common pitfalls. Includes automated generation of reports aligned with TRIPOD+AI reporting guidance (Collins et al. (2024 <[doi:10.1136/bmj-2023-078378](https://doi.org/10.1136/bmj-2023-078378)>)) for reproducible, guideline-aligned research.

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URL <https://github.com/DevWebWacky/triageR>

BugReports <https://github.com/DevWebWacky/triageR/issues>

Suggests knitr, missForest, mlbench, ranger, rmarkdown, spelling, testthat (>= 3.0.0), xgboost

Config/testthat/edition 3

Imports DALEX, dplyr, ellmer, ggplot2, mice, naniar, parsnip, pROC, quarto, recipes, tibble, tidyr, workflows, yardstick

VignetteBuilder knitr

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tr_agent_review	<i>Review a clinical modelling pipeline for common pitfalls</i>
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Description

Runs a series of rule-based checks for common clinical-ML pitfalls (class imbalance, low events-per-variable, possible leakage, near-zero variance predictors), then optionally asks an LLM to summarize the findings in plain language.

Usage

```
tr_agent_review(data, model, use_agent = TRUE)
```

Arguments

data	The data frame used to fit the model.
model	A fitted triageR_model object from tr_fit().
use_agent	Logical. If TRUE (default), also generates an AI-written plain-language summary of the findings via ellmer.

Value

A triageR_review object (list) containing a tibble of flags and, if use_agent = TRUE, an AI-generated summary.

Examples

```
## Not run:
tr_agent_review(data, model)

## End(Not run)
```

tr_check_missing	<i>Check missing data in a clinical dataset</i>
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Description

Produces a summary table of missingness per column, plus a visual plot showing the pattern of missing data across the dataset.

Usage

```
tr_check_missing(data)
```

Arguments

data A data frame, typically the output of `tr_load_clinical()`.

Value

Invisibly returns a summary tibble (columns, `n_missing`, `pct_missing`). Also prints a summary to console and displays a missingness plot as a side effect.

Examples

```
df <- data.frame(a = c(1, NA, 3), b = c(4, 5, NA))
tr_check_missing(df)
```

tr_explain	<i>Explain a clinical prediction model</i>
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Description

Generates variable importance or prediction explanations for a fitted `trriageR_model`, using the DALEX framework under the hood.

Usage

```
tr_explain(
  model,
  method = c("permutation", "shap"),
  newdata = NULL,
  observation = 1
)
```

Arguments

model	A fitted triageR_model object from tr_fit().
method	Character. One of "permutation" (global variable importance) or "shap" (SHAP values for a single prediction).
newdata	Optional data frame to explain predictions on. If NULL, uses the training data.
observation	Integer. Row index of the observation to explain, only used when method = "shap". Defaults to 1.

Value

A triageR_explanation object (list) containing the explanation result and a plot.

Examples

```
## Not run:
tr_explain(model, method = "permutation")
tr_explain(model, method = "shap", observation = 3)

## End(Not run)
```

tr_fit

Fit a clinical prediction model

Description

Fits a binary classification model using the parsnip/workflows framework. The user must specify the model engine explicitly.

Usage

```
tr_fit(
  data,
  outcome,
  engine = c("logistic_reg", "random_forest", "boost_tree"),
  predictors = NULL
)
```

Arguments

data	A data frame containing predictors and the outcome column.
outcome	Character. Name of the binary outcome column (must be a factor or coercible to one, with the event of interest as the second level).
engine	Character. Model engine to use. One of "logistic_reg", "random_forest", or "boost_tree".
predictors	Character vector of predictor column names. If NULL (default), all columns except outcome are used.

Value

A fitted triageR_model object — a list containing the fitted workflow, the engine used, and the outcome/predictor names.

Examples

```
set.seed(1)
df <- data.frame(
  age = round(rnorm(50, 55, 12)),
  sex = sample(c("M", "F"), 50, replace = TRUE),
  disease = sample(c(0, 1), 50, replace = TRUE)
)
model <- tr_fit(df, outcome = "disease", engine = "logistic_reg")
```

tr_impute

*Impute missing values in a clinical dataset***Description**

Fills in missing values using either multiple imputation (mice) or a random-forest based approach (missForest). The method must be chosen explicitly, this function does not guess for you.

Usage

```
tr_impute(data, method = c("mice", "missForest"), m = 5, seed = 123)
```

Arguments

data	A data frame with missing values, typically the output of tr_load_clinical().
method	Character. Either "mice" (default) or "missForest".
m	Integer. Number of multiple imputations to run if method = "mice". Defaults to 5. Ignored for "missForest".
seed	Integer. Random seed for reproducibility. Defaults to 123.

Value

A completed data frame with missing values filled in. If method = "mice", the first completed dataset is returned, and the full mids object is attached as an attribute ("mice_object") in case the user wants to inspect all imputations.

Examples

```
df <- data.frame(
  a = c(5, 7, 3, 9, 2, 8, 6, 4, 5, 7),
  b = c(1, 3, 2, 4, 5, 3, 2, NA, 1, 3)
)
completed <- tr_impute(df, method = "mice", m = 2)
```

tr_load_clinical	<i>Load and standardize a clinical dataset</i>
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Description

Reads a flat clinical dataset (CSV or data frame) and returns it as a standardized triageR object with basic structure checks. This is the entry point for most triageR workflows.

Usage

```
tr_load_clinical(data, id_col = "id")
```

Arguments

data	A file path to a CSV, or an existing data frame.
id_col	Character. Name of the column identifying unique patients. Defaults to "id".

Value

A tibble of class triageR_data, with basic metadata attached.

Examples

```
df <- data.frame(
  id = 1:5,
  age = c(45, 62, 38, 71, 55),
  sex = c("F", "M", "F", "M", "F")
)
tr_load_clinical(df, id_col = "id")
```

tr_recommend_method	<i>Recommend an appropriate statistical or ML method (AI agent)</i>
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Description

Uses an LLM to inspect the structure of a clinical dataset and suggest an appropriate statistical or machine learning approach, with reasoning. This is an assistive tool, not a replacement for expert judgment.

Usage

```
tr_recommend_method(data, outcome, context = NULL)
```

Arguments

data	A data frame, typically the output of <code>tr_load_clinical()</code> .
outcome	Character. Name of the outcome column of interest.
context	Optional character string giving extra clinical context (e.g. "predicting 30-day readmission in heart failure patients").

Value

Invisibly returns the raw text recommendation (character string). Also prints the recommendation to console.

Examples

```
## Not run:
tr_recommend_method(data, outcome = "disease",
  context = "predicting diabetes onset in adults")

## End(Not run)
```

tr_sensitivity

*Run an automated sensitivity analysis battery***Description**

Re-fits a clinical prediction model under several robustness checks, complete-case vs imputed data, outlier exclusion, and subgroup consistency, and compares performance metrics across them.

Usage

```
tr_sensitivity(data, model, subgroup_col = NULL, outlier_sd = 3)
```

Arguments

data	The original (pre-imputation) data frame, containing the same predictors and outcome used in model.
model	A fitted <code>trriageR_model</code> object from <code>tr_fit()</code> .
subgroup_col	Optional character. Name of a categorical column (e.g. "sex") to check subgroup consistency across.
outlier_sd	Numeric. Number of standard deviations beyond which a numeric predictor value is considered an outlier and excluded in the outlier-robustness check. Defaults to 3.

Value

A `trriageR_sensitivity` object (list) with a comparison tibble of metrics across all sensitivity scenarios.

Examples

```
## Not run:
tr_sensitivity(data, model, subgroup_col = "sex")

## End(Not run)
```

tr_tripod_report	<i>Generate a TRIPOD+AI-aligned clinical model report</i>
------------------	---

Description

Renders a reproducible report summarizing model fit, validation, sensitivity analysis, and pipeline review, aligned with TRIPOD+AI reporting guidance. This is a drafting aid, not a certified compliance tool.

Usage

```
tr_tripod_report(
  model,
  validation = NULL,
  review = NULL,
  sensitivity = NULL,
  recommendation = NULL,
  output_file = file.path(tempdir(), "trriageR_report"),
  format = c("html", "docx")
)
```

Arguments

model	A fitted triageR_model object from tr_fit().
validation	Optional triageR_validation object from tr_validate().
review	Optional triageR_review object from tr_agent_review().
sensitivity	Optional triageR_sensitivity object from tr_sensitivity().
recommendation	Optional character string from tr_recommend_method().
output_file	Character. File path (without extension) to save the report to. Defaults to a file in tempdir(). Set explicitly (e.g. file.path("my_folder", "report")) to save elsewhere.
format	Character. Either "html" (default) or "docx".

Value

Invisibly returns the path to the rendered report file.

Examples

```
## Not run:
tr_tripod_report(model, validation = val, review = rev,
  output_file = file.path(tempdir(), "report"))

## End(Not run)
```

tr_validate

*Validate a clinical prediction model***Description**

Computes discrimination (AUC, sensitivity, specificity) and calibration metrics for a fitted `trriageR_model`. Can validate on new (external/holdout) data, or fall back to the training data with a clear warning.

Usage

```
tr_validate(model, newdata = NULL, threshold = 0.5)
```

Arguments

<code>model</code>	A fitted <code>trriageR_model</code> object from <code>tr_fit()</code> .
<code>newdata</code>	Optional data frame to validate on. If <code>NULL</code> (default), validation runs on the original training data, with a warning.
<code>threshold</code>	Numeric. Probability threshold for classifying the positive class. Defaults to 0.5.

Value

A `trriageR_validation` object (list) containing a metrics tibble and the underlying predictions, invisibly printed as a summary.

Examples

```
set.seed(1)
df <- data.frame(
  age = round(rnorm(50, 55, 12)),
  sex = sample(c("M", "F"), 50, replace = TRUE),
  disease = sample(c(0, 1), 50, replace = TRUE)
)
model <- tr_fit(df, outcome = "disease", engine = "logistic_reg")
tr_validate(model, newdata = df)
```

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