

# Package ‘mikropml’

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**Title** User-Friendly R Package for Supervised Machine Learning Pipelines

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**Description** An interface to build machine learning models for classification and regression problems. 'mikropml' implements the ML pipeline described by Topçuoğlu et al. (2020) [doi:10.1128/mBio.00434-20](https://doi.org/10.1128/mBio.00434-20) with reasonable default options for data preprocessing, hyperparameter tuning, cross-validation, testing, model evaluation, and interpretation steps. See the website <https://www.schlosslab.org/mikropml/> for more information, documentation, and examples.

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**URL** <https://www.schlosslab.org/mikropml/>,  
<https://github.com/SchlossLab/mikropml>

**BugReports** <https://github.com/SchlossLab/mikropml/issues>

**Depends** R (>= 4.1.0)

**Imports** caret, dplyr, e1071, glmnet, kernlab, MLmetrics, randomForest, rlang, rpart, stats, utils, xgboost

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---

*calc\_baseline\_precision*  
*Calculate the fraction of positives, i.e. baseline precision for a PRC curve*

---

**Description**

Calculate the fraction of positives, i.e. baseline precision for a PRC curve

**Usage**

```
calc_baseline_precision(dataset, outcome_colname = NULL, pos_outcome = NULL)
```

**Arguments**

dataset	Dataframe with an outcome variable and other columns as features.
outcome_colname	Column name as a string of the outcome variable (default NULL; the first column will be chosen automatically).
pos_outcome	the positive outcome from outcome_colname, e.g. "cancer" for the otu_mini_bin dataset.

**Value**

the baseline precision based on the fraction of positives

**Author(s)**

Kelly Sovacool, <sovacool@umich.edu>

**Examples**

```
# calculate the baseline precision
data.frame(y = c("a", "b", "a", "b")) %>%
  calc_baseline_precision("y", "a")

calc_baseline_precision(otu_mini_bin,
```

```

outcome_colname = "dx",
pos_outcome = "cancer"
)

# if you're not sure which outcome was used as the 'positive' outcome during
# model training, you can access it from the trained model and pass it along:
calc_baseline_precision(otu_mini_bin,
  outcome_colname = "dx",
  pos_outcome = otu_mini_bin_results_glmnet$trained_model$levels[1]
)

```

---

calc\_model\_sensspec    *Calculate and summarize performance for ROC and PRC plots*

---

### Description

Use these functions to calculate cumulative sensitivity, specificity, recall, etc. on single models, concatenate the results together from multiple models, and compute mean ROC and PRC. You can then plot mean ROC and PRC curves to visualize the results. **Note:** These functions assume a binary outcome.

### Usage

```

calc_model_sensspec(trained_model, test_data, outcome_colname = NULL)

calc_mean_roc(sensspec_dat)

calc_mean_prc(sensspec_dat)

```

### Arguments

trained_model	Trained model from <code>caret::train()</code> .
test_data	Held out test data: dataframe of outcome and features.
outcome_colname	Column name as a string of the outcome variable (default NULL; the first column will be chosen automatically).
sensspec_dat	data frame created by concatenating results of <code>calc_model_sensspec()</code> for multiple models.

### Value

data frame with summarized performance

**Functions**

- `calc_model_sensspec()`: Get sensitivity, specificity, and precision for a model.
- `calc_mean_roc()`: Calculate mean sensitivity over specificity for multiple models
- `calc_mean_prc()`: Calculate mean precision over recall for multiple models

**Author(s)**

Courtney Armour

Kelly Sovacool, <sovacool@umich.edu>

**Examples**

```
## Not run:
library(dplyr)
# get cumulative performance for a single model
sensspec_1 <- calc_model_sensspec(
  otu_mini_bin_results_glmnet$trained_model,
  otu_mini_bin_results_glmnet$test_data,
  "dx"
)
head(sensspec_1)

# get performance for multiple models
get_sensspec_seed <- function(seed) {
  ml_result <- run_ml(otu_mini_bin, "glmnet", seed = seed)
  sensspec <- calc_model_sensspec(
    ml_result$trained_model,
    ml_result$test_data,
    "dx"
  ) %>%
    mutate(seed = seed)
  return(sensspec)
}
sensspec_dat <- purrr::map_dfr(seq(100, 102), get_sensspec_seed)

# calculate mean sensitivity over specificity
roc_dat <- calc_mean_roc(sensspec_dat)
head(roc_dat)

# calculate mean precision over recall
prc_dat <- calc_mean_prc(sensspec_dat)
head(prc_dat)

# plot ROC & PRC
roc_dat %>% plot_mean_roc()
baseline_prec <- calc_baseline_precision(otu_mini_bin, "dx", "cancer")
prc_dat %>%
  plot_mean_prc(baseline_precision = baseline_prec)

## End(Not run)
```

---

calc\_perf\_metrics      *Get performance metrics for test data*

---

### Description

Get performance metrics for test data

### Usage

```
calc_perf_metrics(  
  test_data,  
  trained_model,  
  outcome_colname,  
  perf_metric_function,  
  class_probs  
)
```

### Arguments

`test_data`      Held out test data: dataframe of outcome and features.

`trained_model`    Trained model from `caret::train()`.

`outcome_colname`    Column name as a string of the outcome variable (default NULL; the first column will be chosen automatically).

`perf_metric_function`    Function to calculate the performance metric to be used for cross-validation and test performance. Some functions are provided by caret (see `caret::defaultSummary()`). Defaults: binary classification = `twoClassSummary`, multi-class classification = `multiClassSummary`, regression = `defaultSummary`.

`class_probs`      Whether to use class probabilities (TRUE for categorical outcomes, FALSE for numeric outcomes).

### Value

Dataframe of performance metrics.

### Author(s)

Zena Lapp, <zenalapp@umich.edu>

### Examples

```
## Not run:  
results <- run_ml(otu_small, "glmnet", kfold = 2, cv_times = 2)  
calc_perf_metrics(results$test_data,  
  results$trained_model,  
  "dx",
```

```
    multiClassSummary,  
    class_probs = TRUE  
  )  
  
  ## End(Not run)
```

---

combine\_hp\_performance

*Combine hyperparameter performance metrics for multiple train/test splits*

---

## Description

Combine hyperparameter performance metrics for multiple train/test splits generated by, for instance, [looping in R](#) or using a [snakemake workflow](#) on a high-performance computer.

## Usage

```
combine_hp_performance(trained_model_lst)
```

## Arguments

```
trained_model_lst  
  List of trained models.
```

## Value

Named list:

- `dat`: Dataframe of performance metric for each group of hyperparameters
- `params`: Hyperparameters tuned.
- `Metric`: Performance metric used.

## Author(s)

Zena Lapp, <zenalapp@umich.edu>

## Examples

```
## Not run:  
results <- lapply(seq(100, 102), function(seed) {  
  run_ml(otu_small, "glmnet", seed = seed, cv_times = 2, kfold = 2)  
})  
models <- lapply(results, function(x) x$trained_model)  
combine_hp_performance(models)  
  
## End(Not run)
```

---

compare_models	<i>Perform permutation tests to compare the performance metric across all pairs of a group variable.</i>
----------------	--

---

### Description

A wrapper for `permute_p_value()`.

### Usage

```
compare_models(merged_data, metric, group_name, nperm = 10000)
```

### Arguments

<code>merged_data</code>	the concatenated performance data from <code>run_ml</code>
<code>metric</code>	metric to compare, must be numeric
<code>group_name</code>	column with group variables to compare
<code>nperm</code>	number of permutations, default=10000

### Value

a table of p-values for all pairs of group variable

### Author(s)

Courtney R Armour, <armourc@umich.edu>

### Examples

```
df <- dplyr::tibble(  
  model = c("rf", "rf", "glmnet", "glmnet", "svmRadial", "svmRadial"),  
  AUC = c(.2, 0.3, 0.8, 0.9, 0.85, 0.95)  
)  
set.seed(123)  
compare_models(df, "AUC", "model", nperm = 10)
```



---

define_cv	<i>Define cross-validation scheme and training parameters</i>
-----------	---

---

**Description**

Define cross-validation scheme and training parameters

**Usage**

```
define_cv(
  train_data,
  outcome_colname,
  hyperparams_list,
  perf_metric_function,
  class_probs,
  kfold = 5,
  cv_times = 100,
  groups = NULL,
  group_partitions = NULL
)
```

**Arguments**

train_data	Dataframe for training model.
outcome_colname	Column name as a string of the outcome variable (default NULL; the first column will be chosen automatically).
hyperparams_list	Named list of lists of hyperparameters.
perf_metric_function	Function to calculate the performance metric to be used for cross-validation and test performance. Some functions are provided by caret (see <a href="#">caret::defaultSummary()</a> ). Defaults: binary classification = twoClassSummary, multi-class classification = multiClassSummary, regression = defaultSummary.
class_probs	Whether to use class probabilities (TRUE for categorical outcomes, FALSE for numeric outcomes).
kfold	Fold number for k-fold cross-validation (default: 5).
cv_times	Number of cross-validation partitions to create (default: 100).
groups	Vector of groups to keep together when splitting the data into train and test sets. If the number of groups in the training set is larger than kfold, the groups will also be kept together for cross-validation. Length matches the number of rows in the dataset (default: NULL).
group_partitions	Specify how to assign groups to the training and testing partitions (default: NULL). If groups specifies that some samples belong to group "A" and some

belong to group "B", then setting `group_partitions = list(train = c("A", "B"), test = c("B"))` will result in all samples from group "A" being placed in the training set, some samples from "B" also in the training set, and the remaining samples from "B" in the testing set. The partition sizes will be as close to `training_frac` as possible. If the number of groups in the training set is larger than `kfold`, the groups will also be kept together for cross-validation.

### Value

Caret object for `trainControl` that controls cross-validation

### Author(s)

Begüm Topçuoğlu, <topcuoglu.begum@gmail.com>

Kelly Sovacool, <sovacool@umich.edu>

### Examples

```
training_inds <- get_partition_indices(otu_small %>% dplyr::pull("dx"),
  training_frac = 0.8,
  groups = NULL
)
train_data <- otu_small[training_inds, ]
test_data <- otu_small[-training_inds, ]
cv <- define_cv(train_data,
  outcome_colname = "dx",
  hyperparams_list = get_hyperparams_list(otu_small, "glmnet"),
  perf_metric_function = caret::multiClassSummary,
  class_probs = TRUE,
  kfold = 5
)
```

---

`get_caret_processed_df`

*Get preprocessed dataframe for continuous variables*

---

### Description

Get preprocessed dataframe for continuous variables

### Usage

```
get_caret_processed_df(features, method)
```

### Arguments

<code>features</code>	Dataframe of features for machine learning
<code>method</code>	Methods to preprocess the data, described in <code>caret::preProcess()</code> (default: <code>c("center", "scale")</code> ), use <code>NULL</code> for no normalization).

**Value**

Named list:

- processed: Dataframe of processed features.
- removed: Names of any features removed during preprocessing.

**Author(s)**

Zena Lapp, <zenalapp@umich.edu>

**Examples**

```
get_caret_processed_df(mikropml::otu_small[, 2:ncol(otu_small)], c("center", "scale"))
```

---

```
get_feature_importance
```

*Get feature importance using the permutation method*

---

**Description**

Calculates feature importance using a trained model and test data. Requires the `future.apply` package.

**Usage**

```
get_feature_importance(  
  trained_model,  
  test_data,  
  outcome_colname,  
  perf_metric_function,  
  perf_metric_name,  
  class_probs,  
  method,  
  seed = NA,  
  corr_thresh = 1,  
  groups = NULL,  
  nperms = 100,  
  corr_method = "spearman"  
)
```

**Arguments**

<code>trained_model</code>	Trained model from <code>caret::train()</code> .
<code>test_data</code>	Held out test data: dataframe of outcome and features.
<code>outcome_colname</code>	Column name as a string of the outcome variable (default NULL; the first column will be chosen automatically).

perf_metric_function	Function to calculate the performance metric to be used for cross-validation and test performance. Some functions are provided by caret (see <a href="#">caret::defaultSummary()</a> ). Defaults: binary classification = twoClassSummary, multi-class classification = multiClassSummary, regression = defaultSummary.
perf_metric_name	The column name from the output of the function provided to perf_metric_function that is to be used as the performance metric. Defaults: binary classification = "ROC", multi-class classification = "logLoss", regression = "RMSE".
class_probs	Whether to use class probabilities (TRUE for categorical outcomes, FALSE for numeric outcomes).
method	ML method. Options: c("glmnet", "rf", "rpart2", "svmRadial", "xgbTree"). <ul style="list-style-type: none"> <li>• glmnet: linear, logistic, or multiclass regression</li> <li>• rf: random forest</li> <li>• rpart2: decision tree</li> <li>• svmRadial: support vector machine</li> <li>• xgbTree: xgboost</li> </ul>
seed	Random seed (default: NA). Your results will only be reproducible if you set a seed.
corr_thresh	For feature importance, group correlations above or equal to corr_thresh (range 0 to 1; default: 1).
groups	Vector of feature names to group together during permutation. Each element should be a string with feature names separated by a pipe character ( ). If this is NULL (default), correlated features will be grouped together based on corr_thresh.
nperms	number of permutations to perform (default: 100).
corr_method	correlation method. options or the same as those supported by stats::cor: spearman, pearson, kendall. (default: spearman)

## Details

For permutation tests, the p-value is the number of permutation statistics that are greater than the test statistic, divided by the number of permutations. In our case, the permutation statistic is the model performance (e.g. AUROC) after randomizing the order of observations for one feature, and the test statistic is the actual performance on the test data. By default we perform 100 permutations per feature; increasing this will increase the precision of estimating the null distribution, but also increases runtime. The p-value represents the probability of obtaining the actual performance in the event that the null hypothesis is true, where the null hypothesis is that the feature is not important for model performance.

We strongly recommend providing multiple cores to speed up computation time. See [our vignette on parallel processing](#) for more details.

## Value

Data frame with performance metrics for when each feature (or group of correlated features; names) is permuted (perf\_metric), differences between the actual test performance metric on and the

permuted performance metric (`perf_metric_diff`; test minus permuted performance), and the p-value (pvalue: the probability of obtaining the actual performance value under the null hypothesis). Features with a larger `perf_metric_diff` are more important. The performance metric name (`perf_metric_name`) and seed (`seed`) are also returned.

### Author(s)

Begüm Topçuoğlu, <topcuoglu.begum@gmail.com>

Zena Lapp, <zenalapp@umich.edu>

Kelly Sovacool, <sovacool@umich.edu>

### Examples

```
## Not run:
# If you called `run_ml()` with `feature_importance = FALSE` (the default),
# you can use `get_feature_importance()` later as long as you have the
# trained model and test data.
results <- run_ml(otu_small, "glmnet", kfold = 2, cv_times = 2)
names(results$trained_model$trainingData)[1] <- "dx"
feat_imp <- get_feature_importance(results$trained_model,
  results$trained_model$trainingData,
  results$test_data,
  "dx",
  multiClassSummary,
  "AUC",
  class_probs = TRUE,
  method = "glmnet"
)

# We strongly recommend providing multiple cores to speed up computation time.
# Do this before calling `get_feature_importance()`.
doFuture::registerDoFuture()
future::plan(future::multicore, workers = 2)

# Optionally, you can group features together with a custom grouping
feat_imp <- get_feature_importance(results$trained_model,
  results$trained_model$trainingData,
  results$test_data,
  "dx",
  multiClassSummary,
  "AUC",
  class_probs = TRUE,
  method = "glmnet",
  groups = c(
    "Otu00007", "Otu00008", "Otu00009", "Otu00011", "Otu00012",
    "Otu00015", "Otu00016", "Otu00018", "Otu00019", "Otu00020", "Otu00022",
    "Otu00023", "Otu00025", "Otu00028", "Otu00029", "Otu00030", "Otu00035",
    "Otu00036", "Otu00037", "Otu00038", "Otu00039", "Otu00040", "Otu00047",
    "Otu00050", "Otu00052", "Otu00054", "Otu00055", "Otu00056", "Otu00060",
    "Otu00003|Otu00002|Otu00005|Otu00024|Otu00032|Otu00041|Otu00053",
    "Otu00014|Otu00021|Otu00017|Otu00031|Otu00057",
    "Otu00013|Otu00006", "Otu00026|Otu00001|Otu00034|Otu00048",
```

```

    "Otu00033|Otu00010",
    "Otu00042|Otu00004", "Otu00043|Otu00027|Otu00049", "Otu00051|Otu00045",
    "Otu00058|Otu00044", "Otu00059|Otu00046"
  )
)

# the function can show a progress bar if you have the `progressr` package installed.
## optionally, specify the progress bar format:
progressr::handlers(progressr::handler_progress(
  format = ":message :bar :percent | elapsed: :elapsed | eta: :eta",
  clear = FALSE,
  show_after = 0
))
## tell progressr to always report progress
progressr::handlers(global = TRUE)
## run the function and watch the live progress updates
feat_imp <- get_feature_importance(results$trained_model,
  results$trained_model$trainingData,
  results$test_data,
  "dx",
  multiClassSummary,
  "AUC",
  class_probs = TRUE,
  method = "glmnet"
)

# You can specify any correlation method supported by `stats::cor`:
feat_imp <- get_feature_importance(results$trained_model,
  results$trained_model$trainingData,
  results$test_data,
  "dx",
  multiClassSummary,
  "AUC",
  class_probs = TRUE,
  method = "glmnet",
  corr_method = "pearson"
)

## End(Not run)

```

---

get\_hp\_performance      *Get hyperparameter performance metrics*

---

## Description

Get hyperparameter performance metrics

## Usage

```
get_hp_performance(trained_model)
```

**Arguments**

trained\_model    trained model (e.g. from run\_ml())

**Value**

Named list:

- dat: Dataframe of performance metric for each group of hyperparameters.
- params: Hyperparameters tuned.
- metric: Performance metric used.

**Author(s)**

Zena Lapp, <zenalapp@umich.edu>

Kelly Sovacool <sovacool@umich.edu>

**Examples**

```
get_hp_performance(otu_mini_bin_results_glmnet$trained_model)
```

---

get\_hyperparams\_list    *Set hyperparameters based on ML method and dataset characteristics*

---

**Description**

For more details see the vignette on [hyperparameter tuning](#).

**Usage**

```
get_hyperparams_list(dataset, method)
```

**Arguments**

dataset	Dataframe with an outcome variable and other columns as features.
method	ML method. Options: c("glmnet", "rf", "rpart2", "svmRadial", "xgbTree"). <ul style="list-style-type: none"><li>• glmnet: linear, logistic, or multiclass regression</li><li>• rf: random forest</li><li>• rpart2: decision tree</li><li>• svmRadial: support vector machine</li><li>• xgbTree: xgboost</li></ul>

**Value**

Named list of hyperparameters.

**Author(s)**

Kelly Sovacool, <sovacool@umich.edu>

**Examples**

```
get_hyperparams_list(otu_mini_bin, "rf")
get_hyperparams_list(otu_small, "rf")
get_hyperparams_list(otu_mini_bin, "rpart2")
get_hyperparams_list(otu_small, "rpart2")
```

---

get\_outcome\_type      *Get outcome type.*

---

**Description**

If the outcome is numeric, the type is continuous. Otherwise, the outcome type is binary if there are only two outcomes or multiclass if there are more than two outcomes.

**Usage**

```
get_outcome_type(outcomes_vec)
```

**Arguments**

outcomes\_vec      Vector of outcomes.

**Value**

Outcome type (continuous, binary, or multiclass).

**Author(s)**

Zena Lapp, <zenalapp@umich.edu>

**Examples**

```
get_outcome_type(c(1, 2, 1))
get_outcome_type(c("a", "b", "b"))
get_outcome_type(c("a", "b", "c"))
```



---

get\_partition\_indices *Select indices to partition the data into training & testing sets.*

---

### Description

Use this function to get the row indices for the training set.

### Usage

```
get_partition_indices(  
  outcomes,  
  training_frac = 0.8,  
  groups = NULL,  
  group_partitions = NULL  
)
```

### Arguments

outcomes	vector of outcomes
training_frac	Fraction of data for training set (default: 0.8). Rows from the dataset will be randomly selected for the training set, and all remaining rows will be used in the testing set. Alternatively, if you provide a vector of integers, these will be used as the row indices for the training set. All remaining rows will be used in the testing set.
groups	Vector of groups to keep together when splitting the data into train and test sets. If the number of groups in the training set is larger than kfold, the groups will also be kept together for cross-validation. Length matches the number of rows in the dataset (default: NULL).
group_partitions	Specify how to assign groups to the training and testing partitions (default: NULL). If groups specifies that some samples belong to group "A" and some belong to group "B", then setting group_partitions = list(train = c("A", "B"), test = c("B")) will result in all samples from group "A" being placed in the training set, some samples from "B" also in the training set, and the remaining samples from "B" in the testing set. The partition sizes will be as close to training_frac as possible. If the number of groups in the training set is larger than kfold, the groups will also be kept together for cross-validation.

### Details

If groups is NULL, uses [createDataPartition](#). Otherwise, uses `create_grouped_data_partition()`.

Set the seed prior to calling this function if you would like your data partitions to be reproducible (recommended).

### Value

Vector of row indices for the training set.

**Author(s)**

Kelly Sovacool, sovacool@umich.edu

**Examples**

```
training_inds <- get_partition_indices(otu_mini_bin$dx)
train_data <- otu_mini_bin[training_inds, ]
test_data <- otu_mini_bin[-training_inds, ]
```

---

get\_performance\_tbl    *Get model performance metrics as a one-row tibble*

---

**Description**

Get model performance metrics as a one-row tibble

**Usage**

```
get_performance_tbl(  
  trained_model,  
  test_data,  
  outcome_colname,  
  perf_metric_function,  
  perf_metric_name,  
  class_probs,  
  method,  
  seed = NA  
)
```

**Arguments**

**trained\_model**    Trained model from `caret::train()`.

**test\_data**        Held out test data: dataframe of outcome and features.

**outcome\_colname**    Column name as a string of the outcome variable (default NULL; the first column will be chosen automatically).

**perf\_metric\_function**    Function to calculate the performance metric to be used for cross-validation and test performance. Some functions are provided by caret (see `caret::defaultSummary()`). Defaults: binary classification = `twoClassSummary`, multi-class classification = `multiClassSummary`, regression = `defaultSummary`.

**perf\_metric\_name**    The column name from the output of the function provided to `perf_metric_function` that is to be used as the performance metric. Defaults: binary classification = "ROC", multi-class classification = "logLoss", regression = "RMSE".

class_probs	Whether to use class probabilities (TRUE for categorical outcomes, FALSE for numeric outcomes).
method	ML method. Options: c("glmnet", "rf", "rpart2", "svmRadial", "xgbTree"). <ul style="list-style-type: none"> <li>• glmnet: linear, logistic, or multiclass regression</li> <li>• rf: random forest</li> <li>• rpart2: decision tree</li> <li>• svmRadial: support vector machine</li> <li>• xgbTree: xgboost</li> </ul>
seed	Random seed (default: NA). Your results will only be reproducible if you set a seed.

**Value**

A one-row tibble with a column for the cross-validation performance, columns for each of the performance metrics for the test data, plus the method, and seed.

**Author(s)**

Kelly Sovacool, <sovacool@umich.edu>

Zena Lapp, <zenalapp@umich.edu>

**Examples**

```
## Not run:
results <- run_ml(otu_small, "glmnet", kfold = 2, cv_times = 2)
names(results$trained_model$trainingData)[1] <- "dx"
get_performance_tbl(results$trained_model, results$test_data,
  "dx",
  multiClassSummary, "AUC",
  class_probs = TRUE,
  method = "glmnet"
)

## End(Not run)
```

---

get\_perf\_metric\_fn      *Get default performance metric function*

---

**Description**

Get default performance metric function

**Usage**

```
get_perf_metric_fn(outcome_type)
```

**Arguments**

outcome\_type    Type of outcome (one of: "continuous","binary","multiclass").

**Value**

Performance metric function.

**Author(s)**

Zena Lapp, <zenalapp@umich.edu>

**Examples**

```
get_perf_metric_fn("continuous")
get_perf_metric_fn("binary")
get_perf_metric_fn("multiclass")
```

---

get\_perf\_metric\_name    *Get default performance metric name*

---

**Description**

Get default performance metric name for cross-validation.

**Usage**

```
get_perf_metric_name(outcome_type)
```

**Arguments**

outcome\_type    Type of outcome (one of: "continuous","binary","multiclass").

**Value**

Performance metric name.

**Author(s)**

Zena Lapp, <zenalapp@umich.edu>

**Examples**

```
get_perf_metric_name("continuous")
get_perf_metric_name("binary")
get_perf_metric_name("multiclass")
```

---

get_tuning_grid	<i>Generate the tuning grid for tuning hyperparameters</i>
-----------------	--

---

**Description**

Generate the tuning grid for tuning hyperparameters

**Usage**

```
get_tuning_grid(hyperparams_list, method)
```

**Arguments**

`hyperparams_list`  
Named list of lists of hyperparameters.

`method`  
ML method. Options: `c("glmnet", "rf", "rpart2", "svmRadial", "xgbTree")`.

- `glmnet`: linear, logistic, or multiclass regression
- `rf`: random forest
- `rpart2`: decision tree
- `svmRadial`: support vector machine
- `xgbTree`: xgboost

**Value**

The tuning grid.

**Author(s)**

Begüm Topçuoğlu, <topcuoglu.begum@gmail.com>

Kelly Sovacool, <sovacool@umich.edu>

**Examples**

```
ml_method <- "glmnet"  
hparams_list <- get_hyperparams_list(otu_small, ml_method)  
get_tuning_grid(hparams_list, ml_method)
```

---

`group_correlated_features`*Group correlated features*

---

**Description**

Group correlated features

**Usage**

```
group_correlated_features(  
  features,  
  corr_thresh = 1,  
  group_neg_corr = TRUE,  
  corr_method = "spearman"  
)
```

**Arguments**

<code>features</code>	a dataframe with each column as a feature for ML
<code>corr_thresh</code>	For feature importance, group correlations above or equal to <code>corr_thresh</code> (range 0 to 1; default: 1).
<code>group_neg_corr</code>	Whether to group negatively correlated features together (e.g. <code>c(0,1)</code> and <code>c(1,0)</code> ).
<code>corr_method</code>	correlation method. options or the same as those supported by <code>stats::cor</code> : <code>spearman</code> , <code>pearson</code> , <code>kendall</code> . (default: <code>spearman</code> )

**Value**

vector where each element is a group of correlated features separated by pipes (|)

**Author(s)**

Kelly Sovacool, <sovacool@umich.edu>

**Examples**

```
features <- data.frame(  
  a = 1:3, b = 2:4, c = c(1, 0, 1),  
  d = (5:7), e = c(5, 1, 4), f = c(-1, 0, -1)  
)  
group_correlated_features(features)
```

---

mikropml	<i>mikropml: User-Friendly R Package for Robust Machine Learning Pipelines</i>
----------	--

---

## Description

mikropml implements supervised machine learning pipelines using regression, support vector machines, decision trees, random forest, or gradient-boosted trees. The main functions are `preprocess_data()` to process your data prior to running machine learning, and `run_ml()` to run machine learning.

## Authors

- Begüm D. Topçuoğlu ([ORCID](#))
- Zena Lapp ([ORCID](#))
- Kelly L. Sovacool ([ORCID](#))
- Evan Snitkin ([ORCID](#))
- Jenna Wiens ([ORCID](#))
- Patrick D. Schloss ([ORCID](#))

## See vignettes

- [Introduction](#)
- [Preprocessing data](#)
- [Hyperparameter tuning](#)
- [Parallel processing](#)
- [The mikropml paper](#)

---

otu_data_preproc	<i>Mini OTU abundance dataset - preprocessed</i>
------------------	--

---

## Description

This is the result of running `preprocess_data("otu_mini_bin")`

## Usage

```
otu_data_preproc
```

## Format

An object of class `list` of length 3.

---

otu_mini_bin	<i>Mini OTU abundance dataset</i>
--------------	-----------------------------------

---

**Description**

A dataset containing relative abundances of OTUs for human stool samples with a binary outcome, dx. This is a subset of otu\_small.

**Usage**

```
otu_mini_bin
```

**Format**

A data frame The dx column is the diagnosis: healthy or cancerous (colorectal). All other columns are OTU relative abundances.

---

otu_mini_bin_results_glmnet	<i>Results from running the pipeline with L2 logistic regression on otu_mini_bin with feature importance and grouping</i>
-----------------------------	---

---

**Description**

Results from running the pipeline with L2 logistic regression on otu\_mini\_bin with feature importance and grouping

**Usage**

```
otu_mini_bin_results_glmnet
```

**Format**

An object of class list of length 4.



---

`otu_mini_bin_results_rf`

*Results from running the pipeline with random forest on  
otu\_mini\_bin*

---

**Description**

Results from running the pipeline with random forest on otu\_mini\_bin

**Usage**

```
otu_mini_bin_results_rf
```

**Format**

An object of class list of length 4.

---

`otu_mini_bin_results_rpart2`

*Results from running the pipeline with rpart2 on otu\_mini\_bin*

---

**Description**

Results from running the pipeline with rpart2 on otu\_mini\_bin

**Usage**

```
otu_mini_bin_results_rpart2
```

**Format**

An object of class list of length 4.

---

otu\_mini\_bin\_results\_svmRadial

*Results from running the pipeline with svmRadial on otu\_mini\_bin*

---

**Description**

Results from running the pipeline with svmRadial on otu\_mini\_bin

**Usage**

otu\_mini\_bin\_results\_svmRadial

**Format**

An object of class list of length 4.

---

otu\_mini\_bin\_results\_xgbTree

*Results from running the pipeline with xgbTree on otu\_mini\_bin*

---

**Description**

Results from running the pipeline with xgbTree on otu\_mini\_bin

**Usage**

otu\_mini\_bin\_results\_xgbTree

**Format**

An object of class list of length 4.

---

otu\_mini\_cont\_results\_glmnet

*Results from running the pipeline with glmnet on otu\_mini\_bin with Otu00001 as the outcome*

---

**Description**

Results from running the pipeline with glmnet on otu\_mini\_bin with Otu00001 as the outcome

**Usage**

otu\_mini\_cont\_results\_glmnet

**Format**

An object of class list of length 4.

---

otu\_mini\_cont\_results\_nocv

*Results from running the pipeline with glmnet on otu\_mini\_bin with Otu00001 as the outcome column, using a custom train control scheme that does not perform cross-validation*

---

**Description**

Results from running the pipeline with glmnet on otu\_mini\_bin with Otu00001 as the outcome column, using a custom train control scheme that does not perform cross-validation

**Usage**

otu\_mini\_cont\_results\_nocv

**Format**

An object of class list of length 4.

---

otu\_mini\_cv                      *Cross validation on train\_data\_mini with grouped features.*

---

**Description**

Cross validation on train\_data\_mini with grouped features.

**Usage**

```
otu_mini_cv
```

**Format**

An object of class list of length 27.

---

otu\_mini\_multi                      *Mini OTU abundance dataset with 3 categorical variables*

---

**Description**

A dataset containing relative abundances of OTUs for human stool samples

**Usage**

```
otu_mini_multi
```

**Format**

A data frame The dx column is the colorectal cancer diagnosis: adenoma, carcinoma, normal. All other columns are OTU relative abundances.

---

otu\_mini\_multi\_group                      *Groups for otu\_mini\_multi*

---

**Description**

Groups for otu\_mini\_multi

**Usage**

```
otu_mini_multi_group
```

**Format**

An object of class character of length 490.

---

`otu_mini_multi_results_glmnet`

*Results from running the pipeline with glmnet on otu\_mini\_multi for multiclass outcomes*

---

**Description**

Results from running the pipeline with glmnet on otu\_mini\_multi for multiclass outcomes

**Usage**

```
otu_mini_multi_results_glmnet
```

**Format**

An object of class list of length 4.

---

`otu_small`

*Small OTU abundance dataset*

---

**Description**

A dataset containing relative abundances of 60 OTUs for 60 human stool samples. This is a subset of the data provided in `extdata/otu_large.csv`, which was used in [Topçuoğlu \*et al.\* 2020](#).

**Usage**

```
otu_small
```

**Format**

A data frame with 60 rows and 61 variables. The `dx` column is the diagnosis: healthy or cancerous (colorectal). All other columns are OTU relative abundances.

---

permute_p_value	<i>Calculated a permuted p-value comparing two models</i>
-----------------	---

---

### Description

Calculated a permuted p-value comparing two models

### Usage

```
permute_p_value(  
  merged_data,  
  metric,  
  group_name,  
  group_1,  
  group_2,  
  nperm = 10000  
)
```

### Arguments

merged_data	the concatenated performance data from run_ml
metric	metric to compare, must be numeric
group_name	column with group variables to compare
group_1	name of one group to compare
group_2	name of other group to compare
nperm	number of permutations, default=10000

### Value

numeric p-value comparing two models

### Author(s)

Begüm Topçuoğlu, <topcuoglu.begum@gmail.com>  
Courtney R Armour, <armourc@umich.edu>

### Examples

```
df <- dplyr::tibble(  
  model = c("rf", "rf", "glmnet", "glmnet", "svmRadial", "svmRadial"),  
  AUC = c(.2, 0.3, 0.8, 0.9, 0.85, 0.95)  
)  
set.seed(123)  
permute_p_value(df, "AUC", "model", "rf", "glmnet", nperm = 100)
```

---

plot\_hp\_performance *Plot hyperparameter performance metrics*

---

**Description**

Plot hyperparameter performance metrics

**Usage**

```
plot_hp_performance(dat, param_col, metric_col)
```

**Arguments**

dat	dataframe of hyperparameters and performance metric (e.g. from <code>get_hp_performance()</code> or <code>combine_hp_performance()</code> )
param_col	hyperparameter to be plotted. must be a column in dat.
metric_col	performance metric. must be a column in dat.

**Value**

ggplot of hyperparameter performance.

**Author(s)**

Zena Lapp, <zenalapp@umich.edu>

Kelly Sovacool <sovacool@umich.edu>

**Examples**

```
# plot for a single `run_ml()` call
hp_metrics <- get_hp_performance(otu_mini_bin_results_glmnet$trained_model)
hp_metrics
plot_hp_performance(hp_metrics$dat, lambda, AUC)
## Not run:
# plot for multiple `run_ml()` calls
results <- lapply(seq(100, 102), function(seed) {
  run_ml(otu_small, "glmnet", seed = seed)
})
models <- lapply(results, function(x) x$trained_model)
hp_metrics <- combine_hp_performance(models)
plot_hp_performance(hp_metrics$dat, lambda, AUC)

## End(Not run)
```

---

plot\_mean\_roc                      *Plot ROC and PRC curves*

---

### Description

Plot ROC and PRC curves

### Usage

```
plot_mean_roc(dat, ribbon_fill = "#C6DBEF", line_color = "#08306B")

plot_mean_prc(
  dat,
  baseline_precision = NULL,
  ribbon_fill = "#C7E9C0",
  line_color = "#00441B"
)
```

### Arguments

dat                      sensitivity, specificity, and precision data calculated by calc\_mean\_roc()  
ribbon\_fill              ribbon fill color (default: "#D9D9D9")  
line\_color              line color (default: "#000000")  
baseline\_precision              baseline precision from calc\_baseline\_precision()

### Functions

- plot\_mean\_roc(): Plot mean sensitivity over specificity
- plot\_mean\_prc(): Plot mean precision over recall

### Author(s)

Courtney Armour  
Kelly Sovacool <sovacool@umich.edu>

### Examples

```
## Not run:
library(dplyr)
# get performance for multiple models
get_sensspec_seed <- function(seed) {
  ml_result <- run_ml(otu_mini_bin, "glmnet", seed = seed)
  sensspec <- calc_model_sensspec(
    ml_result$trained_model,
    ml_result$test_data,
    "dx"
  )
}
```



```
) %>%
  mutate(seed = seed)
return(sensspec)
}
sensspec_dat <- purrr::map_dfr(seq(100, 102), get_sensspec_seed)

# plot ROC & PRC
sensspec_dat %>%
  calc_mean_roc() %>%
  plot_mean_roc()
baseline_prec <- calc_baseline_precision(otu_mini_bin, "dx", "cancer")
sensspec_dat %>%
  calc_mean_prc() %>%
  plot_mean_prc(baseline_precision = baseline_prec)

## End(Not run)
```

---

plot\_model\_performance

*Plot performance metrics for multiple ML runs with different parameters*

---

### Description

ggplot2 is required to use this function.

### Usage

```
plot_model_performance(performance_df)
```

### Arguments

performance\_df dataframe of performance results from multiple calls to run\_ml()

### Value

A ggplot2 plot of performance.

### Author(s)

Begüm Topçuoğlu, <topcuoglu.begum@gmail.com>

Kelly Sovacool, <sovacool@umich.edu>

**Examples**

```

## Not run:
# call `run_ml()` multiple times with different seeds
results_lst <- lapply(seq(100, 104), function(seed) {
  run_ml(otu_small, "glmnet", seed = seed)
})
# extract and combine the performance results
perf_df <- lapply(results_lst, function(result) {
  result[["performance"]]
}) %>%
  dplyr::bind_rows()
# plot the performance results
p <- plot_model_performance(perf_df)

# call `run_ml()` with different ML methods
param_grid <- expand_grid(
  seeds = seq(100, 104),
  methods = c("glmnet", "rf")
)
results_mtx <- mapply(
  function(seed, method) {
    run_ml(otu_mini_bin, method, seed = seed, kfold = 2)
  },
  param_grid$seeds, param_grid$methods
)
# extract and combine the performance results
perf_df2 <- dplyr::bind_rows(results_mtx["performance", ])
# plot the performance results
p <- plot_model_performance(perf_df2)

# you can continue adding layers to customize the plot
p +
  theme_classic() +
  scale_color_brewer(palette = "Dark2") +
  coord_flip()

## End(Not run)

```

---

preprocess\_data

*Preprocess data prior to running machine learning*


---

**Description**

Function to preprocess your data for input into `run_ml()`.

**Usage**

```
preprocess_data(
```

```

dataset,
outcome_colname,
method = c("center", "scale"),
remove_var = "nzv",
collapse_corr_feats = TRUE,
to_numeric = TRUE,
group_neg_corr = TRUE,
prefilter_threshold = 1
)

```

### Arguments

dataset	Dataframe with an outcome variable and other columns as features.
outcome_colname	Column name as a string of the outcome variable (default NULL; the first column will be chosen automatically).
method	Methods to preprocess the data, described in <a href="#">caret::preProcess()</a> (default: c("center", "scale"), use NULL for no normalization).
remove_var	Whether to remove variables with near-zero variance ('nzv'; default), zero variance ('zv'), or none (NULL).
collapse_corr_feats	Whether to keep only one of perfectly correlated features.
to_numeric	Whether to change features to numeric where possible.
group_neg_corr	Whether to group negatively correlated features together (e.g. c(0,1) and c(1,0)).
prefilter_threshold	Remove features which only have non-zero & non-NA values N rows or fewer (default: 1). Set this to -1 to keep all columns at this step. This step will also be skipped if to_numeric is set to FALSE.

### Value

Named list including:

- `dat_transformed`: Preprocessed data.
- `grp_feats`: If features were grouped together, a named list of the features corresponding to each group.
- `removed_feats`: Any features that were removed during preprocessing (e.g. because there was zero variance or near-zero variance for those features).

If the `progressr` package is installed, a progress bar with time elapsed and estimated time to completion can be displayed.

### More details

See the [preprocessing vignette](#) for more details.

Note that if any values in `outcome_colname` contain spaces, they will be converted to underscores for compatibility with `caret`.

**Author(s)**

Zena Lapp, <zenalapp@umich.edu>

Kelly Sovacool, <sovacool@umich.edu>

**Examples**

```
preprocess_data(mikropml::otu_small, "dx")

# the function can show a progress bar if you have the progressr package installed
## optionally, specify the progress bar format
progressr::handlers(progressr::handler_progress(
  format = ":message :bar :percent | elapsed: :elapsed | eta: :eta",
  clear = FALSE,
  show_after = 0
))
## tell progressor to always report progress
## Not run:
progressr::handlers(global = TRUE)
## run the function and watch the live progress updates
dat_preproc <- preprocess_data(mikropml::otu_small, "dx")

## End(Not run)
```

---

randomize\_feature\_order

*Randomize feature order to eliminate any position-dependent effects*

---

**Description**

Randomize feature order to eliminate any position-dependent effects

**Usage**

```
randomize_feature_order(dataset, outcome_colname)
```

**Arguments**

dataset	Dataframe with an outcome variable and other columns as features.
outcome_colname	Column name as a string of the outcome variable (default NULL; the first column will be chosen automatically).

**Value**

Dataset with feature order randomized.

**Author(s)**

Nick Lesniak, <nlesniak@umich.edu>  
Kelly Sovacool, <sovacool@umich.edu>

**Examples**

```
dat <- data.frame(  
  outcome = c("1", "2", "3"),  
  a = 4:6, b = 7:9, c = 10:12, d = 13:15  
)  
randomize_feature_order(dat, "outcome")
```

---

remove\_singleton\_columns

*Remove columns appearing in only threshold row(s) or fewer.*

---

**Description**

Removes columns which only have non-zero & non-NA values in threshold row(s) or fewer.

**Usage**

```
remove_singleton_columns(dat, threshold = 1)
```

**Arguments**

dat	dataframe
threshold	Number of rows. If a column only has non-zero & non-NA values in threshold row(s) or fewer, it will be removed.

**Value**

dataframe without singleton columns

**Author(s)**

Kelly Sovacool, <sovacool@umich.edu>  
Courtney Armour

**Examples**

```
remove_singleton_columns(data.frame(a = 1:3, b = c(0, 1, 0), c = 4:6))  
remove_singleton_columns(data.frame(a = 1:3, b = c(0, 1, 0), c = 4:6), threshold = 0)  
remove_singleton_columns(data.frame(a = 1:3, b = c(0, 1, NA), c = 4:6))  
remove_singleton_columns(data.frame(a = 1:3, b = c(1, 1, 1), c = 4:6))
```

---

replace_spaces	<i>Replace spaces in all elements of a character vector with underscores</i>
----------------	--

---

**Description**

Replace spaces in all elements of a character vector with underscores

**Usage**

```
replace_spaces(x, new_char = "_")
```

**Arguments**

x	a character vector
new_char	the character to replace spaces (default: _)

**Value**

character vector with all spaces replaced with new\_char

**Author(s)**

Kelly Sovacool, <sovacool@umich.edu>

**Examples**

```
dat <- data.frame(  
  dx = c("outcome 1", "outcome 2", "outcome 1"),  
  a = 1:3, b = c(5, 7, 1)  
)  
dat$dx <- replace_spaces(dat$dx)  
dat
```

---

run_ml	<i>Run the machine learning pipeline</i>
--------	--

---

**Description**

This function runs machine learning (ML), evaluates the best model, and optionally calculates feature importance using the framework outlined in Topçuoğlu *et al.* 2020 ([doi:10.1128/mBio.00434-20](https://doi.org/10.1128/mBio.00434-20)). Required inputs are a dataframe with an outcome variable and other columns as features, as well as the ML method. See `vignette('introduction')` for more details.

**Usage**

```
run_ml(
  dataset,
  method,
  outcome_colname = NULL,
  hyperparameters = NULL,
  find_feature_importance = FALSE,
  calculate_performance = TRUE,
  kfold = 5,
  cv_times = 100,
  cross_val = NULL,
  training_frac = 0.8,
  perf_metric_function = NULL,
  perf_metric_name = NULL,
  groups = NULL,
  group_partitions = NULL,
  corr_thresh = 1,
  seed = NA,
  ...
)
```

**Arguments**

dataset	Dataframe with an outcome variable and other columns as features.
method	ML method. Options: c("glmnet", "rf", "rpart2", "svmRadial", "xgbTree"). <ul style="list-style-type: none"> <li>• glmnet: linear, logistic, or multiclass regression</li> <li>• rf: random forest</li> <li>• rpart2: decision tree</li> <li>• svmRadial: support vector machine</li> <li>• xgbTree: xgboost</li> </ul>
outcome_colname	Column name as a string of the outcome variable (default NULL; the first column will be chosen automatically).
hyperparameters	Dataframe of hyperparameters (default NULL; sensible defaults will be chosen automatically).
find_feature_importance	Run permutation importance (default: FALSE). TRUE is recommended if you would like to identify features important for predicting your outcome, but it is resource-intensive.
calculate_performance	Whether to calculate performance metrics (default: TRUE). You might choose to skip this if you do not perform cross-validation during model training.
kfold	Fold number for k-fold cross-validation (default: 5).
cv_times	Number of cross-validation partitions to create (default: 100).

cross_val	a custom cross-validation scheme from <code>caret::trainControl()</code> (default: NULL, uses <code>kfold</code> cross validation repeated <code>cv_times</code> ). <code>kfold</code> and <code>cv_times</code> are ignored if the user provides a custom cross-validation scheme. See the <code>caret::trainControl()</code> docs for information on how to use it.
training_frac	Fraction of data for training set (default: 0.8). Rows from the dataset will be randomly selected for the training set, and all remaining rows will be used in the testing set. Alternatively, if you provide a vector of integers, these will be used as the row indices for the training set. All remaining rows will be used in the testing set.
perf_metric_function	Function to calculate the performance metric to be used for cross-validation and test performance. Some functions are provided by <code>caret</code> (see <code>caret::defaultSummary()</code> ). Defaults: binary classification = <code>twoClassSummary</code> , multi-class classification = <code>multiClassSummary</code> , regression = <code>defaultSummary</code> .
perf_metric_name	The column name from the output of the function provided to <code>perf_metric_function</code> that is to be used as the performance metric. Defaults: binary classification = "ROC", multi-class classification = "LogLoss", regression = "RMSE".
groups	Vector of groups to keep together when splitting the data into train and test sets. If the number of groups in the training set is larger than <code>kfold</code> , the groups will also be kept together for cross-validation. Length matches the number of rows in the dataset (default: NULL).
group_partitions	Specify how to assign groups to the training and testing partitions (default: NULL). If <code>groups</code> specifies that some samples belong to group "A" and some belong to group "B", then setting <code>group_partitions = list(train = c("A", "B"), test = c("B"))</code> will result in all samples from group "A" being placed in the training set, some samples from "B" also in the training set, and the remaining samples from "B" in the testing set. The partition sizes will be as close to <code>training_frac</code> as possible. If the number of groups in the training set is larger than <code>kfold</code> , the groups will also be kept together for cross-validation.
corr_thresh	For feature importance, group correlations above or equal to <code>corr_thresh</code> (range 0 to 1; default: 1).
seed	Random seed (default: NA). Your results will only be reproducible if you set a seed.
...	All additional arguments are passed on to <code>caret::train()</code> , such as case weights via the <code>weights</code> argument or <code>n</code> tree for rf models. See the <code>caret::train()</code> docs for more details.

## Value

Named list with results:

- `trained_model`: Output of `caret::train()`, including the best model.
- `test_data`: Part of the data that was used for testing.



- **performance**: Dataframe of performance metrics. The first column is the cross-validation performance metric, and the last two columns are the ML method used and the seed (if one was set), respectively. All other columns are performance metrics calculated on the test data. This contains only one row, so you can easily combine performance dataframes from multiple calls to `run_ml()` (see `vignette("parallel")`).
- **feature\_importance**: If feature importances were calculated, a dataframe where each row is a feature or correlated group. The columns are the performance metric of the permuted data, the difference between the true performance metric and the performance metric of the permuted data (true - permuted), the feature name, the ML method, the performance metric name, and the seed (if provided). For AUC and RMSE, the higher `perf_metric_diff` is, the more important that feature is for predicting the outcome. For log loss, the lower `perf_metric_diff` is, the more important that feature is for predicting the outcome.

### More details

For more details, please see [the vignettes](#).

### Author(s)

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Kelly Sovacool, <sovacool@umich.edu>

### Examples

```
## Not run:

# regression
run_ml(otu_small, "glmnet",
      seed = 2019
)

# random forest w/ feature importance
run_ml(otu_small, "rf",
      outcome_colname = "dx",
      find_feature_importance = TRUE
)

# custom cross validation & hyperparameters
run_ml(otu_mini_bin[, 2:11],
      "glmnet",
      outcome_colname = "Otu00001",
      seed = 2019,
      hyperparameters = list(lambda = c(1e-04), alpha = 0),
      cross_val = caret::trainControl(method = "none"),
      calculate_performance = FALSE
)

## End(Not run)
```

---

tidy_perf_data	<i>Tidy the performance dataframe</i>
----------------	---------------------------------------

---

**Description**

Used by `plot_model_performance()`.

**Usage**

```
tidy_perf_data(performance_df)
```

**Arguments**

`performance_df` dataframe of performance results from multiple calls to `run_ml()`

**Value**

Tidy dataframe with model performance metrics.

**Author(s)**

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**Examples**

```
## Not run:
# call `run_ml()` multiple times with different seeds
results_lst <- lapply(seq(100, 104), function(seed) {
  run_ml(otu_small, "glmnet", seed = seed)
})
# extract and combine the performance results
perf_df <- lapply(results_lst, function(result) {
  result[["performance"]]
}) %>%
  dplyr::bind_rows()
# make it pretty!
tidy_perf_data(perf_df)

## End(Not run)
```

---

train_model	<i>Train model using <code>caret::train()</code>.</i>
-------------	---

---

### Description

Train model using `caret::train()`.

### Usage

```
train_model(
  train_data,
  outcome_colname,
  method,
  cv,
  perf_metric_name,
  tune_grid,
  ...
)
```

### Arguments

train_data	Training data. Expected to be a subset of the full dataset.
outcome_colname	Column name as a string of the outcome variable (default NULL; the first column will be chosen automatically).
method	ML method. Options: <code>c("glmnet", "rf", "rpart2", "svmRadial", "xgbTree")</code> . <ul style="list-style-type: none"> <li>• <code>glmnet</code>: linear, logistic, or multiclass regression</li> <li>• <code>rf</code>: random forest</li> <li>• <code>rpart2</code>: decision tree</li> <li>• <code>svmRadial</code>: support vector machine</li> <li>• <code>xgbTree</code>: xgboost</li> </ul>
cv	Cross-validation caret scheme from <code>define_cv()</code> .
perf_metric_name	The column name from the output of the function provided to <code>perf_metric_function</code> that is to be used as the performance metric. Defaults: binary classification = "ROC", multi-class classification = "logLoss", regression = "RMSE".
tune_grid	Tuning grid from <code>get_tuning_grid().#'</code>
...	All additional arguments are passed on to <code>caret::train()</code> , such as case weights via the <code>weights</code> argument or <code>ntree</code> for <code>rf</code> models. See the <code>caret::train()</code> docs for more details.

### Value

Trained model from `caret::train()`.

**Author(s)**

Zena Lapp, <zenalapp@umich.edu>

**Examples**

```
## Not run:
training_data <- otu_mini_bin_results_glmnet$trained_model$trainingData %>%
  dplyr::rename(dx = .outcome)
method <- "rf"
hyperparameters <- get_hyperparams_list(otu_mini_bin, method)
cross_val <- define_cv(training_data,
  "dx",
  hyperparameters,
  perf_metric_function = caret::multiClassSummary,
  class_probs = TRUE,
  cv_times = 2
)
tune_grid <- get_tuning_grid(hyperparameters, method)

rf_model <- train_model(
  training_data,
  "dx",
  method,
  cross_val,
  "AUC",
  tune_grid,
  ntree = 1000
)
rf_model$results %>% dplyr::select(mtry, AUC, prAUC)

## End(Not run)
```

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