# Package 'parsnip'

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```
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add\_rowindex

Add a column of row numbers to a data frame

#### Description

Add a column of row numbers to a data frame

# Usage

add\_rowindex(x)

## Arguments

Х

A data frame

#### Value

The same data frame with a column of 1-based integers named .row.

## Examples

mtcars %>% add\_rowindex()

boost\_tree

General Interface for Boosted Trees

# Description

boost\_tree() is a way to generate a *specification* of a model before fitting and allows the model to be created using different packages in R or via Spark. The main arguments for the model are:

- mtry: The number of predictors that will be randomly sampled at each split when creating the tree models.
- trees: The number of trees contained in the ensemble.
- min\_n: The minimum number of data points in a node that are required for the node to be split further.
- tree\_depth: The maximum depth of the tree (i.e. number of splits).
- learn\_rate: The rate at which the boosting algorithm adapts from iteration-to-iteration.
- loss\_reduction: The reduction in the loss function required to split further.
- sample\_size: The amount of data exposed to the fitting routine.

These arguments are converted to their specific names at the time that the model is fit. Other options and argument can be set using the set\_engine() function. If left to their defaults here (NULL), the values are taken from the underlying model functions. If parameters need to be modified, update() can be used in lieu of recreating the object from scratch.

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#### Usage

```
boost_tree(
  mode = "unknown",
  mtry = NULL,
  trees = NULL,
  min_n = NULL,
  tree_depth = NULL,
  learn_rate = NULL,
  loss_reduction = NULL,
  sample\_size = NULL
)
## S3 method for class 'boost_tree'
update(
  object,
  parameters = NULL,
  mtry = NULL,
  trees = NULL,
  min_n = NULL,
  tree_depth = NULL,
  learn_rate = NULL,
  loss_reduction = NULL,
  sample_size = NULL,
  fresh = FALSE,
)
```

# Arguments

	A • 1 1 .	· · · · · · · · · · · · · · · · · · ·	c 1 1	D 11 1 C 111
mode	A single character	string for the type	of model.	Possible values for this

model are "unknown", "regression", or "classification".

mtry A number for the number (or proportion) of predictors that will be ran-

domly sampled at each split when creating the tree models (xgboost only).

trees An integer for the number of trees contained in the ensemble.

min\_n An integer for the minimum number of data points in a node that are

required for the node to be split further.

tree\_depth An integer for the maximum depth of the tree (i.e. number of splits)

(xgboost only).

learn\_rate A number for the rate at which the boosting algorithm adapts from

iteration-to-iteration (xgboost only).

loss\_reduction A number for the reduction in the loss function required to split further

(xgboost only).

sample\_size A number for the number (or proportion) of data that is exposed to the

fitting routine. For xgboost, the sampling is done at at each iteration

while C5.0 samples once during training.

object A boosted tree model specification.

parameters A 1-row tibble or named list with main parameters to update. If the indi-

vidual arguments are used, these will supersede the values in parameters.

Also, using engine arguments in this object will result in an error.

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fresh	A logical for whether the arguments should be modified in-place of or
	replaced wholesale.
	Not used for update().

### Details

The data given to the function are not saved and are only used to determine the *mode* of the model. For boost\_tree(), the possible modes are "regression" and "classification".

The model can be created using the fit() function using the following engines:

```
R: "xgboost" (the default), "C5.0"Spark: "spark"
```

### Value

An updated model specification.

# **Engine Details**

The standardized parameter names in parsnip can be mapped to their original names in each engine:

parsnip	xgboost	C5.0	spark
${\it tree\_depth}$	$\max_{-depth}$	NA	$\max_{-depth}$
trees	nrounds	trials	$\max_{i}$ iter
$learn\_rate$	eta	NA	$step\_size$
$\operatorname{mtry}$	$colsample\_bytree$	NA	$feature\_subset\_strategy$
$\min_{-n}$	$\min_{\text{child\_weight}}$	minCases	$min\_instances\_per\_node$
loss_reduction	gamma	NA	NA
$sample\_size$	subsample	sample	$subsampling\_rate$
$\min_{i}$ info_gain	NA	NA	$loss\_reduction$

Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are:

```
{\bf xgboost} classification
```

```
parsnip::xgb_train(x = missing_arg(), y = missing_arg(), nthread = 1,
    verbose = 0)

xgboost regression

parsnip::xgb_train(x = missing_arg(), y = missing_arg(), nthread = 1,
    verbose = 0)

C5.0 classification

parsnip::C5.0_train(x = missing_arg(), y = missing_arg(), weights = missing_arg())

spark classification

sparklyr::ml_gradient_boosted_trees(x = missing_arg(), formula = missing_arg(),
    type = "classification", seed = sample.int(10^5, 1))
```

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## spark regression

```
sparklyr::ml_gradient_boosted_trees(x = missing_arg(), formula = missing_arg(),
    type = "regression", seed = sample.int(10^5, 1))
```

#### Note

For models created using the spark engine, there are several differences to consider. First, only the formula interface to via fit() is available; using fit\_xy() will generate an error. Second, the predictions will always be in a spark table format. The names will be the same as documented but without the dots. Third, there is no equivalent to factor columns in spark tables so class predictions are returned as character columns. Fourth, to retain the model object for a new R session (via save()), the model\$fit element of the parsnip object should be serialized via ml\_save(object\$fit) and separately saved to disk. In a new session, the object can be reloaded and reattached to the parsnip object.

#### See Also

```
[fit(), set_engine()
```

## Examples

```
boost_tree(mode = "classification", trees = 20)
# Parameters can be represented by a placeholder:
boost_tree(mode = "regression", mtry = varying())
model <- boost_tree(mtry = 10, min_n = 3)
model
update(model, mtry = 1)
update(model, mtry = 1, fresh = TRUE)

param_values <- tibble::tibble(mtry = 10, tree_depth = 5)
model %>% update(param_values)
model %>% update(param_values, mtry = 3)

param_values$verbose <- 0
# Fails due to engine argument
# model %>% update(param_values)
```

control\_parsnip

Control the fit function

### Description

Options can be passed to the fit() function that control the output and computations

# Usage

```
control_parsnip(verbosity = 1L, catch = FALSE)
fit_control(verbosity = 1L, catch = FALSE)
```

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#### Arguments

verbosity An integer where a value of zero indicates that no messages or output

should be shown when packages are loaded or when the model is fit. A value of 1 means that package loading is quiet but model fits can produce output to the screen (depending on if they contain their own verbose-type argument). A value of 2 or more indicates that any output should

be seen.

catch A logical where a value of TRUE will evaluate the model inside of try(, silent

= TRUE). If the model fails, an object is still returned (without an error)

that inherits the class "try-error".

#### **Details**

fit\_control() is deprecated in favor of control\_parsnip().

#### Value

An S3 object with class "fit\_control" that is a named list with the results of the function call

decision\_tree

General Interface for Decision Tree Models

#### Description

decision\_tree() is a way to generate a *specification* of a model before fitting and allows the model to be created using different packages in R or via Spark. The main arguments for the model are:

- cost\_complexity: The cost/complexity parameter (a.k.a. Cp) used by CART models (rpart only).
- tree\_depth: The *maximum* depth of a tree (rpart and spark only).
- min\_n: The minimum number of data points in a node that are required for the node to be split further.

These arguments are converted to their specific names at the time that the model is fit. Other options and argument can be set using <code>set\_engine()</code>. If left to their defaults here (NULL), the values are taken from the underlying model functions. If parameters need to be modified, <code>update()</code> can be used in lieu of recreating the object from scratch.

### Usage

```
decision_tree(
  mode = "unknown",
  cost_complexity = NULL,
  tree_depth = NULL,
  min_n = NULL
)

## S3 method for class 'decision_tree'
update(
```

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```
object,
parameters = NULL,
cost_complexity = NULL,
tree_depth = NULL,
min_n = NULL,
fresh = FALSE,
...
)
```

## **Arguments**

mode A single character string for the type of model. Possible values for this

model are "unknown", "regression", or "classification".

cost\_complexity

A positive number for the the cost/complexity parameter (a.k.a. Cp) used

by CART models (rpart only).

tree\_depth An integer for maximum depth of the tree.

min\_n An integer for the minimum number of data points in a node that are

required for the node to be split further.

object A random forest model specification.

parameters A 1-row tibble or named list with main parameters to update. If the indi-

 $vidual \ arguments \ are \ used, \ these \ will \ supersede \ the \ values \ in \ {\tt parameters}.$ 

Also, using engine arguments in this object will result in an error.

fresh A logical for whether the arguments should be modified in-place of or

replaced wholesale.

.. Not used for update().

## **Details**

The model can be created using the fit() function using the following engines:

 $\bullet$  R: "rpart" (the default) or "C5.0" (classification only)

• Spark: "spark"

Note that, for rpart models, but cost\_complexity and tree\_depth can be both be specified but the package will give precedence to cost\_complexity. Also, tree\_depth values greater than 30 rpart will give nonsense results on 32-bit machines.

# **Engine Details**

The standardized parameter names in parsnip can be mapped to their original names in each engine:

parsnip	rpart	C5.0	spark
${\it tree\_depth}$	$\max$ depth	NA	$\max_{-depth}$
$\min_{-n}$	minsplit	$\min Cases$	$min\_instances\_per\_node$
$cost\_complexity$	ср	NA	NA

Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are::

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```
rpart classification

rpart::rpart(formula = missing_arg(), data = missing_arg(), weights = missing_arg())

rpart regression

rpart::rpart(formula = missing_arg(), data = missing_arg(), weights = missing_arg())

C5.0 classification

parsnip::C5.0_train(x = missing_arg(), y = missing_arg(), weights = missing_arg(), trials = 1)

spark classification

sparklyr::ml_decision_tree_classifier(x = missing_arg(), formula = missing_arg(), seed = sample.int(10^5, 1))

spark regression

sparklyr::ml_decision_tree_classifier(x = missing_arg(), formula = missing_arg(), seed = sample.int(10^5, 1))
```

#### Note

For models created using the spark engine, there are several differences to consider. First, only the formula interface to via fit() is available; using fit\_xy() will generate an error. Second, the predictions will always be in a spark table format. The names will be the same as documented but without the dots. Third, there is no equivalent to factor columns in spark tables so class predictions are returned as character columns. Fourth, to retain the model object for a new R session (via save()), the model\$fit element of the parsnip object should be serialized via ml\_save(object\$fit) and separately saved to disk. In a new session, the object can be reloaded and reattached to the parsnip object.

## See Also

[fit()

# Examples

```
decision_tree(mode = "classification", tree_depth = 5)
# Parameters can be represented by a placeholder:
decision_tree(mode = "regression", cost_complexity = varying())
model <- decision_tree(cost_complexity = 10, min_n = 3)
model
update(model, cost_complexity = 1)
update(model, cost_complexity = 1, fresh = TRUE)</pre>
```

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descriptors

Data Set Characteristics Available when Fitting Models

# Description

When using the fit() functions there are some variables that will be available for use in arguments. For example, if the user would like to choose an argument value based on the current number of rows in a data set, the .obs() function can be used. See Details below.

# Usage

- .cols()
- .preds()
- .obs()
- .lvls()
- .facts()
- .x()
- .y()
- .dat()

## Details

Existing functions:

- .obs(): The current number of rows in the data set.
- .preds(): The number of columns in the data set that are associated with the predictors prior to dummy variable creation.
- .cols(): The number of predictor columns available after dummy variables are created (if any).
- $\bullet$  .facts (): The number of factor predictors in the dat set.
- .lvls(): If the outcome is a factor, this is a table with the counts for each level (and NA otherwise).
- $\bullet\,$  .x(): The predictors returned in the format given. Either a data frame or a matrix.
- .y(): The known outcomes returned in the format given. Either a vector, matrix, or data frame.
- .dat(): A data frame containing all of the predictors and the outcomes. If fit\_xy() was used, the outcomes are attached as the column, ..y.

For example, if you use the model formula  ${\sf Sepal.Width}$   $\tilde{\ }$  . with the iris data, the values would be

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```
.preds() =
                       (the 4 columns in `iris`)
                        (3 numeric columns + 2 from Species dummy variables)
.cols() =
            5
.obs()
       = 150
.1vls() = NA
                       (no factor outcome)
.facts() =
                        (the Species predictor)
        = <vector>
                        (Sepal.Width as a vector)
.y()
        = <data.frame> (The other 4 columns as a data frame)
.x()
        = <data.frame> (The full data set)
.dat()
```

If the formula  $\mathsf{Species}\ \widetilde{\ }$  . where used:

To use these in a model fit, pass them to a model specification. The evaluation is delayed until the time when the model is run via fit() (and the variables listed above are available). For example:

```
library(modeldata)
data("lending_club")

rand_forest(mode = "classification", mtry = .cols() - 2)
```

When no descriptors are found, the computation of the descriptor values is not executed.

fit.model\_spec

Fit a Model Specification to a Dataset

# Description

fit() and fit\_xy() take a model specification, translate the required code by substituting arguments, and execute the model fit routine.

### Usage

```
## S3 method for class 'model_spec'
fit(object, formula, data, control = control_parsnip(), ...)
## S3 method for class 'model_spec'
fit_xy(object, x, y, control = control_parsnip(), ...)
```

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## Arguments

object	An object of class model_spec that has a chosen engine (via set_engine()).
formula	An object of class "formula" (or one that can be coerced to that class): a symbolic description of the model to be fitted.
data	Optional, depending on the interface (see Details below). A data frame containing all relevant variables (e.g. outcome(s), predictors, case weights, etc). Note: when needed, a <i>named argument</i> should be used.
control	A named list with elements verbosity and catch. See control_parsnip().
• • •	Not currently used; values passed here will be ignored. Other options required to fit the model should be passed using set_engine().
x	A matrix or data frame of predictors.
у	A vector, matrix or data frame of outcome data.

#### **Details**

fit() and fit\_xy() substitute the current arguments in the model specification into the computational engine's code, checks them for validity, then fits the model using the data and the engine-specific code. Different model functions have different interfaces (e.g. formula or x/y) and these functions translate between the interface used when fit() or fit\_xy() were invoked and the one required by the underlying model.

When possible, these functions attempt to avoid making copies of the data. For example, if the underlying model uses a formula and fit() is invoked, the original data are references when the model is fit. However, if the underlying model uses something else, such as x/y, the formula is evaluated and the data are converted to the required format. In this case, any calls in the resulting model objects reference the temporary objects used to fit the model.

If the model engine has not been set, the model's default engine will be used (as discussed on each model page). If the verbosity option of control\_parsnip() is greater than zero, a warning will be produced.

## Value

A model\_fit object that contains several elements:

- lvl: If the outcome is a factor, this contains the factor levels at the time of model fitting.
- spec: The model specification object (object in the call to fit)
- fit: when the model is executed without error, this is the model object. Otherwise, it is a try-error object with the error message.
- preproc: any objects needed to convert between a formula and non-formula interface (such as the terms object)

The return value will also have a class related to the fitted model (e.g. "\_glm") before the base class of "model\_fit".

# See Also

```
set_engine(), control_parsnip(), model_spec, model_fit
```

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#### Examples

```
# Although `glm()` only has a formula interface, different
# methods for specifying the model can be used
library(dplyr)
library(modeldata)
data("lending_club")
lr_mod <- logistic_reg()</pre>
using_formula <-
  1r_mod %>%
  set_engine("glm") %>%
  fit(Class ~ funded_amnt + int_rate, data = lending_club)
using_xy <-
  1r_mod %>%
   set_engine("glm") %>%
  fit_xy(x = lending_club[, c("funded_amnt", "int_rate")],
         y = lending_club$Class)
using_formula
using_xy
```

linear\_reg

General Interface for Linear Regression Models

### Description

linear\_reg() is a way to generate a *specification* of a model before fitting and allows the model to be created using different packages in R, Stan, keras, or via Spark. The main arguments for the model are:

- penalty: The total amount of regularization in the model. Note that this must be zero for some engines.
- mixture: The proportion of L1 regularization in the model. Note that this will be ignored for some engines.

These arguments are converted to their specific names at the time that the model is fit. Other options and argument can be set using set\_engine(). If left to their defaults here (NULL), the values are taken from the underlying model functions. If parameters need to be modified, update() can be used in lieu of recreating the object from scratch.

# Usage

```
linear_reg(mode = "regression", penalty = NULL, mixture = NULL)
## S3 method for class 'linear_reg'
update(
  object,
  parameters = NULL,
  penalty = NULL,
  mixture = NULL,
```

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```
fresh = FALSE,
...
)
```

## Arguments

mode A single character string for the type of model. The only possible value

for this model is "regression".

penalty A non-negative number representing the total amount of regularization

(glmnet, keras, and spark only). For keras models, this corresponds to purely L2 regularization (aka weight decay) while the other models can be a combination of L1 and L2 (depending on the value of mixture).

mixture A number between zero and one (inclusive) that represents the proportion

of regularization that is used for the L2 penalty (i.e. weight decay, or ridge

regression) versus L1 (the lasso) (glmnet and spark only).

object A linear regression model specification.

parameters A 1-row tibble or named list with main parameters to update. If the indi-

vidual arguments are used, these will supersede the values in parameters.

Also, using engine arguments in this object will result in an error.

fresh A logical for whether the arguments should be modified in-place of or

replaced wholesale.

... Not used for update().

#### Details

The data given to the function are not saved and are only used to determine the *mode* of the model. For linear\_reg(), the mode will always be "regression".

The model can be created using the fit() function using the following engines:

• R: "lm" (the default) or "glmnet"

Stan: "stan"Spark: "spark"keras: "keras"

## **Engine Details**

The standardized parameter names in parsnip can be mapped to their original names in each engine:

parsnip	${f glmnet}$	spark	keras
penalty	lambda	$reg\_param$	penalty
mixture	alpha	elastic_net_param	NA

Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are:

## lm

```
stats::lm(formula = missing_arg(), data = missing_arg(), weights = missing_arg())
```

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#### glmnet

```
glmnet::glmnet(x = missing_arg(), y = missing_arg(), weights = missing_arg(),
    family = "gaussian")

stan

rstanarm::stan_glm(formula = missing_arg(), data = missing_arg(),
    weights = missing_arg(), family = stats::gaussian, refresh = 0)

(note that the refresh default prevents logging of the estimation process. Change this value in set_engine() will show the logs)

spark

sparklyr::ml_linear_regression(x = missing_arg(), formula = missing_arg(),
    weight_col = missing_arg())
```

parsnip::keras\_mlp(x = missing\_arg(), y = missing\_arg(), hidden\_units = 1, act = "linear")

For glmnet models, the full regularization path is always fit regardless of the value given to penalty. Also, there is the option to pass multiple values (or no values) to the penalty argument. When using the predict() method in these cases, the return value depends on the value of penalty. When using predict(), only a single value of the penalty can be used. When predicting on multiple penalties, the multi\_predict() function can be used. It returns a tibble with a list column called .pred that contains a tibble with all of the penalty results.

For prediction, the stan engine can compute posterior intervals analogous to confidence and prediction intervals. In these instances, the units are the original outcome and when std\_error = TRUE, the standard deviation of the posterior distribution (or posterior predictive distribution as appropriate) is returned.

### Note

For models created using the spark engine, there are several differences to consider. First, only the formula interface to via fit() is available; using fit\_xy() will generate an error. Second, the predictions will always be in a spark table format. The names will be the same as documented but without the dots. Third, there is no equivalent to factor columns in spark tables so class predictions are returned as character columns. Fourth, to retain the model object for a new R session (via save()), the model\$fit element of the parsnip object should be serialized via ml\_save(object\$fit) and separately saved to disk. In a new session, the object can be reloaded and reattached to the parsnip object.

# See Also

```
fit(), set_engine()
```

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#### Examples

```
linear_reg()
# Parameters can be represented by a placeholder:
linear_reg(penalty = varying())
model <- linear_reg(penalty = 10, mixture = 0.1)
model
update(model, penalty = 1)
update(model, penalty = 1, fresh = TRUE)</pre>
```

 $logistic\_reg$ 

General Interface for Logistic Regression Models

## Description

logistic\_reg() is a way to generate a *specification* of a model before fitting and allows the model to be created using different packages in R, Stan, keras, or via Spark. The main arguments for the model are:

- penalty: The total amount of regularization in the model. Note that this must be zero for some engines.
- mixture: The proportion of L1 regularization in the model. Note that this will be ignored for some engines.

These arguments are converted to their specific names at the time that the model is fit. Other options and argument can be set using set\_engine(). If left to their defaults here (NULL), the values are taken from the underlying model functions. If parameters need to be modified, update() can be used in lieu of recreating the object from scratch.

# Usage

```
logistic_reg(mode = "classification", penalty = NULL, mixture = NULL)
## S3 method for class 'logistic_reg'
update(
   object,
   parameters = NULL,
   penalty = NULL,
   mixture = NULL,
   fresh = FALSE,
   ...
)
```

## Arguments

mode

A single character string for the type of model. The only possible value for this model is "classification".

penalty

A non-negative number representing the total amount of regularization (glmnet, keras, and spark only). For keras models, this corresponds to purely L2 regularization (aka weight decay) while the other models can be a combination of L1 and L2 (depending on the value of mixture).

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mixture A number between zero and one (inclusive) that represents the proportion

of regularization that is used for the L2 penalty (i.e. weight decay, or ridge  $\,$ 

regression) versus L1 (the lasso) (glmnet and spark only).

object A logistic regression model specification.

parameters A 1-row tibble or named list with main parameters to update. If the indi-

vidual arguments are used, these will supersede the values in parameters.

Also, using engine arguments in this object will result in an error.

fresh A logical for whether the arguments should be modified in-place of or

replaced wholesale.

... Not used for update().

#### Details

For logistic\_reg(), the mode will always be "classification".

The model can be created using the fit() function using the following engines:

ullet R: "glm" (the default) or "glmnet"

• Stan: "stan"

• Spark: "spark"

• keras: "keras"

## **Engine Details**

The standardized parameter names in parsnip can be mapped to their original names in each engine:

parsnip	$\mathbf{glmnet}$	spark	keras
penalty	lambda	$reg\_param$	penalty
mixture	alpha	elastic_net_param	NA

Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are:

#### glm

```
stats::glm(formula = missing_arg(), data = missing_arg(), weights = missing_arg(),
    family = stats::binomial)
```

# ${f glmnet}$

```
glmnet::glmnet(x = missing_arg(), y = missing_arg(), weights = missing_arg(),
    family = "binomial")
```

#### stan

```
rstanarm::stan_glm(formula = missing_arg(), data = missing_arg(),
    weights = missing_arg(), family = stats::binomial, refresh = 0)
```

(note that the refresh default prevents logging of the estimation process. Change this value in set\_engine() will show the logs)

## spark

18 logistic\_reg

```
sparklyr::ml_logistic_regression(x = missing_arg(), formula = missing_arg(),
    weight_col = missing_arg(), family = "binomial")
```

keras

```
parsnip::keras_mlp(x = missing_arg(), y = missing_arg(), hidden_units = 1,
    act = "linear")
```

For glmnet models, the full regularization path is always fit regardless of the value given to penalty. Also, there is the option to pass multiple values (or no values) to the penalty argument. When using the predict() method in these cases, the return value depends on the value of penalty. When using predict(), only a single value of the penalty can be used. When predicting on multiple penalties, the multi\_predict() function can be used. It returns a tibble with a list column called .pred that contains a tibble with all of the penalty results.

For prediction, the stan engine can compute posterior intervals analogous to confidence and prediction intervals. In these instances, the units are the original outcome and when std\_error = TRUE, the standard deviation of the posterior distribution (or posterior predictive distribution as appropriate) is returned. For glm, the standard error is in logit units while the intervals are in probability units.

#### Note

For models created using the spark engine, there are several differences to consider. First, only the formula interface to via fit() is available; using fit\_xy() will generate an error. Second, the predictions will always be in a spark table format. The names will be the same as documented but without the dots. Third, there is no equivalent to factor columns in spark tables so class predictions are returned as character columns. Fourth, to retain the model object for a new R session (via save()), the model\$fit element of the parsnip object should be serialized via ml\_save(object\$fit) and separately saved to disk. In a new session, the object can be reloaded and reattached to the parsnip object.

# See Also

fit()

# Examples

```
logistic_reg()
# Parameters can be represented by a placeholder:
logistic_reg(penalty = varying())
model <- logistic_reg(penalty = 10, mixture = 0.1)
model
update(model, penalty = 1)
update(model, penalty = 1, fresh = TRUE)</pre>
```

mars 19

mars

General Interface for MARS

## Description

mars() is a way to generate a *specification* of a model before fitting and allows the model to be created using R. The main arguments for the model are:

- num\_terms: The number of features that will be retained in the final model.
- prod\_degree: The highest possible degree of interaction between features. A value of 1 indicates and additive model while a value of 2 allows, but does not guarantee, two-way interactions between features.
- prune\_method: The type of pruning. Possible values are listed in ?earth.

These arguments are converted to their specific names at the time that the model is fit. Other options and argument can be set using <code>set\_engine()</code>. If left to their defaults here (NULL), the values are taken from the underlying model functions. If parameters need to be modified, <code>update()</code> can be used in lieu of recreating the object from scratch.

## Usage

```
mars(
   mode = "unknown",
   num_terms = NULL,
   prod_degree = NULL,
   prune_method = NULL
)

## S3 method for class 'mars'
update(
   object,
   parameters = NULL,
   num_terms = NULL,
   prod_degree = NULL,
   prune_method = NULL,
   fresh = FALSE,
   ...
)
```

### Arguments

A single character string for the type of model. Possible values for this

 ${\it model are "unknown", "regression", or "classification"}.$ 

num\_terms The number of features that will be retained in the final model, including

the intercept.

prod\_degree The highest possible interaction degree.

prune\_method The pruning method.

object A MARS model specification.

20 mars

parameters	A 1-row tibble or named list with <i>main</i> parameters to update. If the individual arguments are used, these will supersede the values in parameters. Also, using engine arguments in this object will result in an error.
fresh	A logical for whether the arguments should be modified in-place of or replaced wholesale.
	Not used for update().

#### **Details**

The model can be created using the fit() function using the following engines:

• R: "earth" (the default)

## **Engine Details**

The standardized parameter names in parsnip can be mapped to their original names in each engine:

parsnip	$\mathbf{earth}$
$num\_terms$	nprune
$\operatorname{prod\_degree}$	degree
$prune\_method$	pmethod

Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are:

earth classification

```
earth::earth(x = missing_arg(), y = missing_arg(), weights = missing_arg(),
    glm = list(family = stats::binomial), keepxy = TRUE)

earth regression

earth::earth(x = missing_arg(), y = missing_arg(), weights = missing_arg(),
    keepxy = TRUE)
```

Note that, when the model is fit, the **earth** package only has its namespace loaded. However, if multi\_predict is used, the package is attached.

# See Also

```
fit()
```

## Examples

```
mars(mode = "regression", num_terms = 5)
model <- mars(num_terms = 10, prune_method = "none")
model
update(model, num_terms = 1)
update(model, num_terms = 1, fresh = TRUE)</pre>
```

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mlp

General Interface for Single Layer Neural Network

## Description

mlp(), for multilayer perceptron, is a way to generate a *specification* of a model before fitting and allows the model to be created using different packages in R or via keras The main arguments for the model are:

- hidden\_units: The number of units in the hidden layer (default: 5).
- penalty: The amount of L2 regularization (aka weight decay, default is zero).
- dropout: The proportion of parameters randomly dropped out of the model (keras only, default is zero).
- epochs: The number of training iterations (default: 20).
- activation: The type of function that connects the hidden layer and the input variables (keras only, default is softmax).

If parameters need to be modified, this function can be used in lieu of recreating the object from scratch.

# Usage

```
mlp(
  mode = "unknown",
  hidden_units = NULL,
  penalty = NULL,
  dropout = NULL,
  epochs = NULL,
  activation = NULL
## S3 method for class 'mlp'
update(
  object,
  parameters = NULL,
  hidden_units = NULL,
  penalty = NULL,
  dropout = NULL,
  epochs = NULL,
  activation = NULL,
  fresh = FALSE,
)
```

# Arguments

mode A single character string for the type of model. Possible values for this

model are "unknown", "regression", or "classification".

hidden\_units An integer for the number of units in the hidden model.

penalty A non-negative numeric value for the amount of weight decay.

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dropout A number between 0 (inclusive) and 1 denoting the proportion of model

parameters randomly set to zero during model training.

epochs An integer for the number of training iterations.

activation A single character strong denoting the type of relationship between the

original predictors and the hidden unit layer. The activation function between the hidden and output layers is automatically set to either "linear" or "softmax" depending on the type of outcome. Possible values are:

"linear", "softmax", "relu", and "elu"

object A random forest model specification.

parameters A 1-row tibble or named list with main parameters to update. If the indi-

 $vidual \ arguments \ are \ used, \ these \ will \ supersede \ the \ values \ in \ {\tt parameters}.$ 

Also, using engine arguments in this object will result in an error.

fresh A logical for whether the arguments should be modified in-place of or

replaced wholesale.

... Not used for update().

### Details

These arguments are converted to their specific names at the time that the model is fit. Other options and argument can be set using set\_engine(). If left to their defaults here (see above), the values are taken from the underlying model functions. One exception is hidden\_units when nnet::nnet is used; that function's size argument has no default so a value of 5 units will be used. Also, unless otherwise specified, the linout argument to nnet::nnet() will be set to TRUE when a regression model is created. If parameters need to be modified, update() can be used in lieu of recreating the object from scratch.

The model can be created using the fit() function using the following engines:

• R: "nnet" (the default)

• keras: "keras"

An error is thrown if both penalty and dropout are specified for keras models.

## **Engine Details**

The standardized parameter names in parsnip can be mapped to their original names in each engine:

parsnip	keras	$\mathbf{nnet}$
hidden_units	$hidden\_units$	size
penalty	penalty	decay
dropout	dropout	NA
epochs	epochs	$\max$ it
activation	activation	NA

Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are:

keras classification

```
parsnip::keras_mlp(x = missing_arg(), y = missing_arg())
```

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```
keras regression
   parsnip::keras_mlp(x = missing_arg(), y = missing_arg())
    nnet classification
    nnet::nnet(formula = missing_arg(), data = missing_arg(), weights = missing_arg(),
        size = 5, trace = FALSE, linout = FALSE)
    nnet regression
   nnet::nnet(formula = missing_arg(), data = missing_arg(), weights = missing_arg(),
        size = 5, trace = FALSE, linout = TRUE)
See Also
    fit()
Examples
    mlp(mode = "classification", penalty = 0.01)
    # Parameters can be represented by a placeholder:
   mlp(mode = "regression", hidden_units = varying())
   model <- mlp(hidden_units = 10, dropout = 0.30)</pre>
   model
   update(model, hidden_units = 2)
   update(model, hidden_units = 2, fresh = TRUE)
```

 $model_fit$ 

Model Fit Object Information

## Description

An object with class "model\_fit" is a container for information about a model that has been fit to the data.

## **Details**

The main elements of the object are:

- lvl: A vector of factor levels when the outcome is a factor. This is NULL when the outcome is not a factor vector.
- spec: A model\_spec object.
- fit: The object produced by the fitting function.
- preproc: This contains any data-specific information required to process new a sample point for prediction. For example, if the underlying model function requires arguments x and y and the user passed a formula to fit, the preproc object would contain items such as the terms object and so on. When no information is required, this is NA.

As discussed in the documentation for <code>model\_spec</code>, the original arguments to the specification are saved as quosures. These are evaluated for the <code>model\_fit</code> object prior to fitting. If the resulting model object prints its call, any user-defined options are shown in the call preceded by a tilde (see the example below). This is a result of the use of quosures in the specification.

This class and structure is the basis for how **parsnip** stores model objects after to seeing the data and applying a model.

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#### Examples

```
# Keep the `x` matrix if the data are not too big.
spec_obj <-
    linear_reg() %>%
    set_engine("lm", x = ifelse(.obs() < 500, TRUE, FALSE))
spec_obj
fit_obj <- fit(spec_obj, mpg ~ ., data = mtcars)
fit_obj
nrow(fit_obj$fit$x)</pre>
```

model\_spec

Model Specification Information

# Description

An object with class "model\_spec" is a container for information about a model that will be fit.

# **Details**

The main elements of the object are:

- args: A vector of the main arguments for the model. The names of these arguments may be different form their counterparts n the underlying model function. For example, for a glmnet model, the argument name for the amount of the penalty is called "penalty" instead of "lambda" to make it more general and usable across different types of models (and to not be specific to a particular model function). The elements of args can varying(). If left to their defaults (NULL), the arguments will use the underlying model functions default value. As discussed below, the arguments in args are captured as quosures and are not immediately executed.
  - ...: Optional model-function-specific parameters. As with args, these will be quosures and can be varying().
  - mode: The type of model, such as "regression" or "classification". Other modes will be added once the package adds more functionality.
  - method: This is a slot that is filled in later by the model's constructor function. It
    generally contains lists of information that are used to create the fit and prediction
    code as well as required packages and similar data.
  - engine: This character string declares exactly what software will be used. It can be a package name or a technology type.

This class and structure is the basis for how **parsnip** stores model objects prior to seeing the data.

## **Argument Details**

An important detail to understand when creating model specifications is that they are intended to be functionally independent of the data. While it is true that some tuning parameters are *data dependent*, the model specification does not interact with the data at all.

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For example, most R functions immediately evaluate their arguments. For example, when calling mean(dat\_vec), the object dat\_vec is immediately evaluated inside of the function. parsnip model functions do not do this. For example, using

```
rand_forest(mtry = ncol(iris) - 1)
```

**does not** execute ncol(iris) -1 when creating the specification. This can be seen in the output:

```
> rand_forest(mtry = ncol(iris) - 1)
Random Forest Model Specification (unknown)
Main Arguments:
    mtry = ncol(iris) - 1
```

The model functions save the argument *expressions* and their associated environments (a.k.a. a quosure) to be evaluated later when either fit() or fit\_xy() are called with the actual data.

The consequence of this strategy is that any data required to get the parameter values must be available when the model is fit. The two main ways that this can fail is if:

- 1. The data have been modified between the creation of the model specification and when the model fit function is invoked.
- 2. If the model specification is saved and loaded into a new session where those same data objects do not exist.

The best way to avoid these issues is to not reference any data objects in the global environment but to use data descriptors such as .cols(). Another way of writing the previous specification is

```
rand_forest(mtry = .cols() - 1)
```

This is not dependent on any specific data object and is evaluated immediately before the model fitting process begins.

One less advantageous approach to solving this issue is to use quasiquotation. This would insert the actual R object into the model specification and might be the best idea when the data object is small. For example, using

```
rand_forest(mtry = ncol(!!iris) - 1)
```

would work (and be reproducible between sessions) but embeds the entire iris data set into the mtry expression:

```
> rand_forest(mtry = ncol(!!iris) - 1)
Random Forest Model Specification (unknown)

Main Arguments:
   mtry = ncol(structure(list(Sepal.Length = c(5.1, 4.9, 4.7, 4.6, 5, <snip>
```

However, if there were an object with the number of columns in it, this wouldn't be too bad:

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```
> mtry_val <- ncol(iris) - 1
> mtry_val
[1] 4
> rand_forest(mtry = !!mtry_val)
Random Forest Model Specification (unknown)
Main Arguments:
    mtry = 4
```

More information on quosures and quasiquotation can be found at <a href="https://tidyeval.tidyverse.org">https://tidyeval.tidyverse.org</a>.

 $multinom\_reg$ 

General Interface for Multinomial Regression Models

# Description

multinom\_reg() is a way to generate a *specification* of a model before fitting and allows the model to be created using different packages in R, keras, or Spark. The main arguments for the model are:

- penalty: The total amount of regularization in the model. Note that this must be zero for some engines.
- mixture: The proportion of L1 regularization in the model. Note that this will be ignored for some engines.

These arguments are converted to their specific names at the time that the model is fit. Other options and argument can be set using set\_engine(). If left to their defaults here (NULL), the values are taken from the underlying model functions. If parameters need to be modified, update() can be used in lieu of recreating the object from scratch.

## Usage

```
multinom_reg(mode = "classification", penalty = NULL, mixture = NULL)
## S3 method for class 'multinom_reg'
update(
  object,
  parameters = NULL,
  penalty = NULL,
  mixture = NULL,
  fresh = FALSE,
    ...
)
```

# Arguments

mode

A single character string for the type of model. The only possible value for this model is "classification".

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penalty	A non-negative number representing the total amount of regularization (glmnet, keras, and spark only). For keras models, this corresponds to purely L2 regularization (aka weight decay) while the other models can be a combination of L1 and L2 (depending on the value of mixture).
mixture	A number between zero and one (inclusive) that represents the proportion of regularization that is used for the L2 penalty (i.e. weight decay, or ridge regression) versus L1 (the lasso) (glmnet only).
object	A multinomial regression model specification.
parameters	A 1-row tibble or named list with <i>main</i> parameters to update. If the individual arguments are used, these will supersede the values in parameters. Also, using engine arguments in this object will result in an error.
fresh	A logical for whether the arguments should be modified in-place of or replaced wholesale.
	Not used for update().

### **Details**

For multinom\_reg(), the mode will always be "classification".

The model can be created using the fit() function using the following engines:

```
R: "glmnet" (the default), "nnet"Stan: "stan"keras: "keras"
```

# **Engine Details**

The standardized parameter names in parsnip can be mapped to their original names in each engine:

parsnip	$\mathbf{glmnet}$	spark	keras	$\mathbf{nnet}$
penalty	lambda	reg_param	penalty	decay
mixture	alpha	elastic_net_param	NA	NA

Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are:

# glmnet

```
glmnet::glmnet(x = missing_arg(), y = missing_arg(), weights = missing_arg(),
    family = "multinomial")

nnet

nnet::multinom(formula = missing_arg(), data = missing_arg(),
    weights = missing_arg(), trace = FALSE)

spark

sparklyr::ml_logistic_regression(x = missing_arg(), formula = missing_arg(),
    weight_col = missing_arg(), family = "multinomial")
```

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#### keras

```
parsnip::keras_mlp(x = missing_arg(), y = missing_arg(), hidden_units = 1,
    act = "linear")
```

For glmnet models, the full regularization path is always fit regardless of the value given to penalty. Also, there is the option to pass multiple values (or no values) to the penalty argument. When using the predict() method in these cases, the return value depends on the value of penalty. When using predict(), only a single value of the penalty can be used. When predicting on multiple penalties, the multi\_predict() function can be used. It returns a tibble with a list column called .pred that contains a tibble with all of the penalty results.

### Note

For models created using the spark engine, there are several differences to consider. First, only the formula interface to via fit() is available; using fit\_xy() will generate an error. Second, the predictions will always be in a spark table format. The names will be the same as documented but without the dots. Third, there is no equivalent to factor columns in spark tables so class predictions are returned as character columns. Fourth, to retain the model object for a new R session (via save()), the model\$fit element of the parsnip object should be serialized via ml\_save(object\$fit) and separately saved to disk. In a new session, the object can be reloaded and reattached to the parsnip object.

# See Also

fit()

# Examples

```
multinom_reg()
# Parameters can be represented by a placeholder:
multinom_reg(penalty = varying())
model <- multinom_reg(penalty = 10, mixture = 0.1)
model
update(model, penalty = 1)
update(model, penalty = 1, fresh = TRUE)</pre>
```

multi\_predict

Model predictions across many sub-models

## Description

For some models, predictions can be made on sub-models in the model object.

# Usage

```
multi_predict(object, ...)
## Default S3 method:
multi_predict(object, ...)
```

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```
## S3 method for class '`_xgb.Booster`'
multi_predict(object, new_data, type = NULL, trees = NULL, ...)

## S3 method for class '`_C5.0`'
multi_predict(object, new_data, type = NULL, trees = NULL, ...)

## S3 method for class '`_elnet`'
multi_predict(object, new_data, type = NULL, penalty = NULL, ...)

## S3 method for class '`_lognet`'
multi_predict(object, new_data, type = NULL, penalty = NULL, ...)

## S3 method for class '`_earth`'
multi_predict(object, new_data, type = NULL, num_terms = NULL, ...)

## S3 method for class '`_multnet`'
multi_predict(object, new_data, type = NULL, penalty = NULL, ...)

## S3 method for class '`_train.kknn`'
multi_predict(object, new_data, type = NULL, neighbors = NULL, ...)
```

### Arguments

object	A model_fit object.
	Optional arguments to pass to predict.model_fit(type = "raw") such as type.
new_data	A rectangular data object, such as a data frame.
type	A single character value or NULL. Possible values are "numeric", "class", "prob", "conf_int", "pred_int", "quantile", or "raw". When NULL, predict() will choose an appropriate value based on the model's mode.
trees	An integer vector for the number of trees in the ensemble.
penalty	A numeric vector of penalty values.
num_terms	An integer vector for the number of MARS terms to retain.
neighbors	An integer vector for the number of nearest neighbors.

#### Value

A tibble with the same number of rows as the data being predicted. There is a list-column named .pred that contains tibbles with multiple rows per sub-model. Note that, within the tibbles, the column names follow the usual standard based on prediction type (i.e. .pred\_class for type = "class" and so on).

30 nearest\_neighbor

## Description

nearest\_neighbor() is a way to generate a *specification* of a model before fitting and allows the model to be created using different packages in R. The main arguments for the model are:

- neighbors: The number of neighbors considered at each prediction.
- weight\_func: The type of kernel function that weights the distances between samples.
- dist\_power: The parameter used when calculating the Minkowski distance. This corresponds to the Manhattan distance with dist\_power = 1 and the Euclidean distance with dist\_power = 2.

These arguments are converted to their specific names at the time that the model is fit. Other options and argument can be set using set\_engine(). If left to their defaults here (NULL), the values are taken from the underlying model functions. If parameters need to be modified, update() can be used in lieu of recreating the object from scratch.

## Usage

```
nearest_neighbor(
  mode = "unknown",
  neighbors = NULL,
  weight_func = NULL,
  dist_power = NULL)
```

# Arguments

mode	A single character string for the type of model. Possible values for this model are "unknown", "regression", or "classification".
neighbors	A single integer for the number of neighbors to consider (often called k). For <b>kknn</b> , a value of 5 is used if <b>neighbors</b> is not specified.
weight_func	A <i>single</i> character for the type of kernel function used to weight distances between samples. Valid choices are: "rectangular", "triangular", "epanechnikov", "biweight", "triweight", "cos", "inv", "gaussian", "rank", or "optimal".
$dist_{\mathtt{power}}$	A single number for the parameter used in calculating Minkowski distance.

#### **Details**

The model can be created using the fit() function using the following engines:

• R: "kknn" (the default)

# **Engine Details**

The standardized parameter names in parsnip can be mapped to their original names in each engine:

parsnip	kknn
neighbors	ks
$weight\_func$	kernel
$dist\_power$	distance

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Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are:

```
kknn (classification or regression)
```

```
kknn::train.kknn(formula = missing_arg(), data = missing_arg(),
    ks = 5)
```

#### Note

For kknn, the underlying modeling function used is a restricted version of train.kknn() and not kknn(). It is set up in this way so that parsnip can utilize the underlying predict.train.kknn method to predict on new data. This also means that a single value of that function's kernel argument (a.k.a weight\_func here) can be supplied

### See Also

```
fit()
```

### Examples

```
nearest_neighbor(neighbors = 11)
```

nullmodel

Fit a simple, non-informative model

# Description

Fit a single mean or largest class model. nullmodel() is the underlying computational function for the null\_model() specification.

## Usage

```
nullmodel(x, ...)
## Default S3 method:
nullmodel(x = NULL, y, ...)
## S3 method for class 'nullmodel'
print(x, ...)
## S3 method for class 'nullmodel'
predict(object, new_data = NULL, type = NULL, ...)
```

# Arguments

Х	An optional matrix or data frame of predictors. These values are not used in the model fit
	Optional arguments (not yet used)
у	A numeric vector (for regression) or factor (for classification) of outcomes
object	An object of class nullmodel
new_data	A matrix or data frame of predictors (only used to determine the number of predictions to return)
type	Either "raw" (for regression), "class" or "prob" (for classification)

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#### **Details**

nullmodel() emulates other model building functions, but returns the simplest model possible given a training set: a single mean for numeric outcomes and the most prevalent class for factor outcomes. When class probabilities are requested, the percentage of the training set samples with the most prevalent class is returned.

#### Value

The output of nullmodel() is a list of class nullmodel with elements

the function call

value the mean of y or the most prevalent class

levels when y is a factor, a vector of levels. NULL otherwise

pct when y is a factor, a data frame with a column for each class (NULL

otherwise). The column for the most prevalent class has the proportion of the training samples with that class (the other columns are zero).

n the number of elements in y

predict.nullmodel() returns a either a factor or numeric vector depending on the class of y. All predictions are always the same.

## Examples

null\_model

General Interface for null models

# Description

null\_model() is a way to generate a *specification* of a model before fitting and allows the model to be created using R. It doesn't have any main arguments.

#### Usage

```
null_model(mode = "classification")
```

## Arguments

mode

A single character string for the type of model. Possible values for this model are "unknown", "regression", or "classification".

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#### **Details**

The model can be created using the fit() function using the following engines:

• R: "parsnip"

# **Engine Details**

Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are:

```
parsnip classification
```

```
nullmodel(x = missing_arg(), y = missing_arg())
parsnip regression
nullmodel(x = missing_arg(), y = missing_arg())
```

#### See Also

fit()

#### Examples

```
null_model(mode = "regression")
```

predict.model\_fit

Model predictions

## Description

Apply a model to create different types of predictions. predict() can be used for all types of models and used the "type" argument for more specificity.

# Usage

```
## S3 method for class 'model_fit'
predict(object, new_data, type = NULL, opts = list(), ...)
## S3 method for class 'model_fit'
predict_raw(object, new_data, opts = list(), ...)
```

# Arguments

object An object of class model\_fit

new\_data A rectangular data object, such as a data frame.

type A single character value or NULL. Possible values are "numeric", "class",

"prob", "conf\_int", "pred\_int", "quantile", or "raw". When NULL, predict()

will choose an appropriate value based on the model's mode.

opts A list of optional arguments to the underlying predict function that will

be used when type = "raw". The list should not include options for the

model object or the new data being predicted.

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Arguments to the underlying model's prediction function cannot be passed here (see opts). There are some parsnip related options that can be passed, depending on the value of type. Possible arguments are:

- level: for types of "conf\_int" and "pred\_int" this is the parameter for the tail area of the intervals (e.g. confidence level for confidence intervals). Default value is 0.95.
- std\_error: add the standard error of fit or prediction for types of "conf\_int" and "pred\_int". Default value is FALSE.
- quantile: the quantile(s) for quantile regression (not implemented yet)
- time: the time(s) for hazard probability estimates (not implemented yet)

#### **Details**

If "type" is not supplied to predict(), then a choice is made (type = "numeric" for regression models and type = "class" for classification).

predict() is designed to provide a tidy result (see "Value" section below) in a tibble output format.

When using type = "conf\_int" and type = "pred\_int", the options level and std\_error can be used. The latter is a logical for an extra column of standard error values (if available).

#### Value

With the exception of type = "raw", the results of predict.model\_fit() will be a tibble as many rows in the output as there are rows in new\_data and the column names will be predictable.

For numeric results with a single outcome, the tibble will have a .pred column and .pred\_Yname for multivariate results.

For hard class predictions, the column is named .pred\_class and, when type = "prob", the columns are .pred\_classlevel.

type = "conf\_int" and type = "pred\_int" return tibbles with columns .pred\_lower and .pred\_upper with an attribute for the confidence level. In the case where intervals can be produces for class probabilities (or other non-scalar outputs), the columns will be named .pred\_lower\_classlevel and so on.

Quantile predictions return a tibble with a column .pred, which is a list-column. Each list element contains a tibble with columns .pred and .quantile (and perhaps other columns).

Using type = "raw" with predict.model\_fit() will return the unadulterated results of the prediction function.

In the case of Spark-based models, since table columns cannot contain dots, the same convention is used except 1) no dots appear in names and 2) vectors are never returned but type-specific prediction functions.

When the model fit failed and the error was captured, the predict() function will return the same structure as above but filled with missing values. This does not currently work for multivariate models.

#### Examples

library(dplyr)

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```
lm_model <-</pre>
  linear_reg() %>%
  set_engine("lm") %>%
  fit(mpg ~ ., data = mtcars %>% slice(11:32))
pred_cars <-</pre>
  mtcars %>%
  slice(1:10) %>%
  select(-mpg)
predict(lm_model, pred_cars)
predict(
  lm_model,
  pred_cars,
  type = "conf_int",
  level = 0.90
predict(
  lm_model,
  pred_cars,
  type = "raw".
  opts = list(type = "terms")
```

rand\_forest

General Interface for Random Forest Models

# Description

rand\_forest() is a way to generate a *specification* of a model before fitting and allows the model to be created using different packages in R or via Spark. The main arguments for the model are:

- mtry: The number of predictors that will be randomly sampled at each split when creating the tree models.
- trees: The number of trees contained in the ensemble.
- min\_n: The minimum number of data points in a node that are required for the node to be split further.

These arguments are converted to their specific names at the time that the model is fit. Other options and argument can be set using set\_engine(). If left to their defaults here (NULL), the values are taken from the underlying model functions. If parameters need to be modified, update() can be used in lieu of recreating the object from scratch.

# Usage

```
rand_forest(mode = "unknown", mtry = NULL, trees = NULL, min_n = NULL)
## S3 method for class 'rand_forest'
update(
  object,
```

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```
parameters = NULL,
mtry = NULL,
trees = NULL,
min_n = NULL,
fresh = FALSE,
...
)
```

# Arguments

mode	A single character string for the type of model. Possible values for this model are "unknown", "regression", or "classification".
mtry	An integer for the number of predictors that will be randomly sampled at each split when creating the tree models.
trees	An integer for the number of trees contained in the ensemble.
min_n	An integer for the minimum number of data points in a node that are required for the node to be split further.
object	A random forest model specification.
parameters	A 1-row tibble or named list with <i>main</i> parameters to update. If the individual arguments are used, these will supersede the values in parameters. Also, using engine arguments in this object will result in an error.
fresh	A logical for whether the arguments should be modified in-place of or
	replaced wholesale.

# **Details**

The model can be created using the fit() function using the following engines:

```
\bullet~R\hbox{:}~"{\tt ranger"}~({\rm the~default})~{\rm or}~"{\tt randomForest"}
```

• Spark: "spark"

# **Engine Details**

The standardized parameter names in parsnip can be mapped to their original names in each engine:

parsnip	ranger	${f random Forest}$	spark
$\operatorname{mtry}$	$\operatorname{mtry}$	mtry	$feature\_subset\_strategy$
trees	num.trees	ntree	$num\_trees$
$\min_{-n}$	min.node.size	nodesize	min_instances_per_node

Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are::

ranger classification

```
ranger::ranger(formula = missing_arg(), data = missing_arg(),
    case.weights = missing_arg(), num.threads = 1, verbose = FALSE,
    seed = sample.int(10^5, 1), probability = TRUE)
```

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For ranger confidence intervals, the intervals are constructed using the form estimate  $+/-z * std_error$ . For classification probabilities, these values can fall outside of [0, 1] and will be coerced to be in this range.

#### Note

For models created using the spark engine, there are several differences to consider. First, only the formula interface to via fit() is available; using fit\_xy() will generate an error. Second, the predictions will always be in a spark table format. The names will be the same as documented but without the dots. Third, there is no equivalent to factor columns in spark tables so class predictions are returned as character columns. Fourth, to retain the model object for a new R session (via save), the model\$fit element of the parsnip object should be serialized via ml\_save(object\$fit) and separately saved to disk. In a new session, the object can be reloaded and reattached to the parsnip object.

## See Also

fit()

#### Examples

```
rand_forest(mode = "classification", trees = 2000)
# Parameters can be represented by a placeholder:
rand_forest(mode = "regression", mtry = varying())
model <- rand_forest(mtry = 10, min_n = 3)
model
update(model, mtry = 1)
update(model, mtry = 1, fresh = TRUE)</pre>
```

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set\_args

Change elements of a model specification

# Description

set\_args() can be used to modify the arguments of a model specification while set\_mode() is used to change the model's mode.

## Usage

```
set_args(object, ...)
set_mode(object, mode)
```

# Arguments

object A model specification.

... One or more named model arguments.

mode A character string for the model type (e.g. "classification" or "regres-

sion")

# **Details**

set\_args() will replace existing values of the arguments.

#### Value

An updated model object.

# Examples

```
rand_forest()
rand_forest() %>%
  set_args(mtry = 3, importance = TRUE) %>%
  set_mode("regression")
```

set\_engine

Declare a computational engine and specific arguments

# Description

set\_engine() is used to specify which package or system will be used to fit the model, along with any arguments specific to that software.

#### Usage

```
set_engine(object, engine, ...)
```

surv\_reg 39

## Arguments

object A model specification.

engine A character string for the software that should be used to fit the model.

This is highly dependent on the type of model (e.g. linear regression,

random forest, etc.).

... Any optional arguments associated with the chosen computational engine.

These are captured as quosures and can be varying().

#### Value

An updated model specification.

#### Examples

```
# First, set general arguments using the standardized names
mod <-
    logistic_reg(mixture = 1/3) %>%
    # now say how you want to fit the model and another other options
    set_engine("glmnet", nlambda = 10)
translate(mod, engine = "glmnet")
```

surv\_reg

General Interface for Parametric Survival Models

# Description

surv\_reg() is a way to generate a *specification* of a model before fitting and allows the model to be created using R. The main argument for the model is:

• dist: The probability distribution of the outcome.

This argument is converted to its specific names at the time that the model is fit. Other options and argument can be set using set\_engine(). If left to its default here (NULL), the value is taken from the underlying model functions.

If parameters need to be modified, this function can be used in lieu of recreating the object from scratch.

## Usage

```
surv_reg(mode = "regression", dist = NULL)
## S3 method for class 'surv_reg'
update(object, parameters = NULL, dist = NULL, fresh = FALSE, ...)
```

# Arguments

Mode A single character string for the type of model. The only possible value

for this model is "regression".

dist A character string for the outcome distribution. "weibull" is the default.

object A survival regression model specification.

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parameters A 1-row tibble or named list with *main* parameters to update. If the individual arguments are used, these will supersede the values in parameters.

Also, using engine arguments in this object will result in an error.

fresh A logical for whether the arguments should be modified in-place of or

replaced wholesale.

... Not used for update().

#### **Details**

The data given to the function are not saved and are only used to determine the *mode* of the model. For <code>surv\_reg()</code>,the mode will always be "regression".

Since survival models typically involve censoring (and require the use of survival::Surv() objects), the fit() function will require that the survival model be specified via the formula interface.

Also, for the flexsurv::flexsurvfit engine, the typical strata function cannot be used. To achieve the same effect, the extra parameter roles can be used (as described above).

For surv\_reg(), the mode will always be "regression".

The model can be created using the fit() function using the following engines:

• R: "flexsurv", "survival" (the default)

#### **Engine Details**

The standardized parameter names in parsnip can be mapped to their original names in each engine:

```
parsnip flexsurv survival dist dist dist
```

Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are:

#### flexsurv

```
flexsurv::flexsurvreg(formula = missing_arg(), data = missing_arg(),
    weights = missing_arg())
```

## survival

```
survival::survreg(formula = missing_arg(), data = missing_arg(),
    weights = missing_arg(), model = TRUE)
```

Note that model = TRUE is needed to produce quantile predictions when there is a stratification variable and can be overridden in other cases.

# References

Jackson, C. (2016). flexsurv: A Platform for Parametric Survival Modeling in R. *Journal of Statistical Software*, 70(8), 1 - 33.

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#### See Also

```
fit(), survival::Surv()
```

## Examples

```
surv_reg()
# Parameters can be represented by a placeholder:
surv_reg(dist = varying())

model <- surv_reg(dist = "weibull")
model
update(model, dist = "lnorm")</pre>
```

svm\_poly

General interface for polynomial support vector machines

# Description

svm\_poly() is a way to generate a *specification* of a model before fitting and allows the model to be created using different packages in R or via Spark. The main arguments for the model are:

- cost: The cost of predicting a sample within or on the wrong side of the margin.
- degree: The polynomial degree.
- scale\_factor: A scaling factor for the kernel.
- margin: The epsilon in the SVM insensitive loss function (regression only)

These arguments are converted to their specific names at the time that the model is fit. Other options and argument can be set using <code>set\_engine()</code>. If left to their defaults here (NULL), the values are taken from the underlying model functions. If parameters need to be modified, <code>update()</code> can be used in lieu of recreating the object from scratch.

# Usage

```
svm_poly(
 mode = "unknown",
 cost = NULL,
 degree = NULL,
  scale_factor = NULL,
 margin = NULL
## S3 method for class 'svm_poly'
update(
 object,
 parameters = NULL,
 cost = NULL,
 degree = NULL,
  scale_factor = NULL,
 margin = NULL,
  fresh = FALSE,
)
```

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## Arguments

Mode A single character string for the type of model. Possible values for this model are "unknown", "regression", or "classification".

A positive number for the cost of predicting a sample within or on the

wrong side of the margin

degree A positive number for polynomial degree.

scale\_factor A positive number for the polynomial scaling factor.

margin A positive number for the epsilon in the SVM insensitive loss function

(regression only)

object A polynomial SVM model specification.

parameters A 1-row tibble or named list with main parameters to update. If the indi-

vidual arguments are used, these will supersede the values in parameters.

Also, using engine arguments in this object will result in an error.

fresh A logical for whether the arguments should be modified in-place of or

replaced wholesale.

... Not used for update().

#### **Details**

The model can be created using the fit() function using the following engines:

• R: "kernlab" (the default)

# **Engine Details**

The standardized parameter names in parsnip can be mapped to their original names in each engine:

parsnip	kernlab
cost	$\mathbf{C}$
degree	degree
$scale\_factor$	scale
margin	epsilon

Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are::

kernlab classification

```
kernlab::ksvm(x = missing_arg(), y = missing_arg(), kernel = "polydot",
    prob.model = TRUE)
```

## kernlab regression

```
kernlab::ksvm(x = missing_arg(), y = missing_arg(), kernel = "polydot")
```

# See Also

fit()

svm\_rbf 43

#### Examples

```
svm_poly(mode = "classification", degree = 1.2)
# Parameters can be represented by a placeholder:
svm_poly(mode = "regression", cost = varying())
model <- svm_poly(cost = 10, scale_factor = 0.1)
model
update(model, cost = 1)
update(model, cost = 1, fresh = TRUE)</pre>
```

 $svm_rbf$ 

General interface for radial basis function support vector machines

# Description

svm\_rbf() is a way to generate a *specification* of a model before fitting and allows the model to be created using different packages in R or via Spark. The main arguments for the model are:

- cost: The cost of predicting a sample within or on the wrong side of the margin.
- rbf\_sigma: The precision parameter for the radial basis function.
- margin: The epsilon in the SVM insensitive loss function (regression only)

These arguments are converted to their specific names at the time that the model is fit. Other options and argument can be set using set\_engine(). If left to their defaults here (NULL), the values are taken from the underlying model functions. If parameters need to be modified, update() can be used in lieu of recreating the object from scratch.

#### Usage

```
svm_rbf(mode = "unknown", cost = NULL, rbf_sigma = NULL, margin = NULL)
## S3 method for class 'svm_rbf'
update(
  object,
  parameters = NULL,
  cost = NULL,
  rbf_sigma = NULL,
  margin = NULL,
  fresh = FALSE,
  ...
)
```

## Arguments

mode	A single character string for the type of model. Possible values for this model are "unknown", "regression", or "classification".
cost	A positive number for the cost of predicting a sample within or on the wrong side of the margin
rbf_sigma	A positive number for radial basis function.

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margin	A positive number for the epsilon in the SVM insensitive loss function (regression only)
object	A radial basis function SVM model specification.
parameters	A 1-row tibble or named list with <i>main</i> parameters to update. If the individual arguments are used, these will supersede the values in parameters. Also, using engine arguments in this object will result in an error.
fresh	A logical for whether the arguments should be modified in-place of or replaced wholesale.

... Not used for update().

#### **Details**

The model can be created using the fit() function using the following engines:

• R: "kernlab" (the default)

## **Engine Details**

The standardized parameter names in parsnip can be mapped to their original names in each engine:

parsnip	kernlab
$\cos t$	$\mathbf{C}$
$rbf\_sigma$	$_{ m sigma}$
margin	epsilon

Engines may have pre-set default arguments when executing the model fit call. For this type of model, the template of the fit calls are::

kernlab classification

```
kernlab::ksvm(x = missing_arg(), y = missing_arg(), kernel = "rbfdot",
    prob.model = TRUE)
kernlab regression
```

```
kernlab::ksvm(x = missing_arg(), y = missing_arg(), kernel = "rbfdot")
```

# See Also

fit()

# Examples

```
svm_rbf(mode = "classification", rbf_sigma = 0.2)
# Parameters can be represented by a placeholder:
svm_rbf(mode = "regression", cost = varying())
model <- svm_rbf(cost = 10, rbf_sigma = 0.1)
model
update(model, cost = 1)
update(model, cost = 1, fresh = TRUE)</pre>
```

tidy.model\_fit 45

tidy.model\_fit

Turn a parsnip model object into a tidy tibble

# Description

This method tidies the model in a parsnip model object, if it exists.

## Usage

```
tidy.model_fit(x, ...)
```

# Arguments

x An object to be converted into a tidy tibble::tibble().... Additional arguments to tidying method.

#### Value

a tibble

translate

Resolve a Model Specification for a Computational Engine

#### Description

translate() will translate a model specification into a code object that is specific to a particular engine (e.g. R package). It translates generic parameters to their counterparts.

# Usage

```
translate(x, ...)
## Default S3 method:
translate(x, engine = x$engine, ...)
```

# Arguments

x A model specification.... Not currently used.engine The computational engine for the model (see ?set\_engine).

#### **Details**

translate() produces a *template* call that lacks the specific argument values (such as data, etc). These are filled in once fit() is called with the specifics of the data for the model. The call may also include varying arguments if these are in the specification.

It does contain the resolved argument names that are specific to the model fitting function/engine.

This function can be useful when you need to understand how parsnip goes from a generic model specific to a model fitting function.

**Note**: this function is used internally and users should only use it to understand what the underlying syntax would be. It should not be used to modify the model specification.

#### Examples

```
lm_spec <- linear_reg(penalty = 0.01)

# `penalty` is tranlsated to `lambda`
translate(lm_spec, engine = "glmnet")

# `penalty` not applicable for this model.
translate(lm_spec, engine = "lm")

# `penalty` is tranlsated to `reg_param`
translate(lm_spec, engine = "spark")

# with a placeholder for an unknown argument value:
translate(linear_reg(mixture = varying()), engine = "glmnet")</pre>
```

varying

A placeholder function for argument values

# Description

varying() is used when a parameter will be specified at a later date.

# Usage

varying()

varying\_args.model\_spec

Determine varying arguments

# Description

varying\_args() takes a model specification or a recipe and returns a tibble of information on all possible varying arguments and whether or not they are actually varying.

#### Usage

```
## S3 method for class 'model_spec'
varying_args(object, full = TRUE, ...)
## S3 method for class 'recipe'
varying_args(object, full = TRUE, ...)
## S3 method for class 'step'
varying_args(object, full = TRUE, ...)
```

#### Arguments

 $object \hspace{1cm} A \hspace{0.1cm} \mathsf{model\_spec} \hspace{0.1cm} or \hspace{0.1cm} a \hspace{0.1cm} \mathsf{recipe}.$ 

full A single logical. Should all possible varying parameters be returned? If

FALSE, then only the parameters that are actually varying are returned.

... Not currently used.

#### **Details**

The id column is determined differently depending on whether a model\_spec or a recipe is used. For a model\_spec, the first class is used. For a recipe, the unique step id is used.

#### Value

A tibble with columns for the parameter name (name), whether it contains *any* varying value (varying), the id for the object (id), and the class that was used to call the method (type).

#### Examples

```
# List all possible varying args for the random forest spec
rand_forest() %>% varying_args()
# mtry is now recognized as varying
rand_forest(mtry = varying()) %>% varying_args()
# Even engine specific arguments can vary
rand_forest() %>%
  set_engine("ranger", sample.fraction = varying()) %>%
  varying_args()
# List only the arguments that actually vary
rand_forest() %>%
  set_engine("ranger", sample.fraction = varying()) %>%
  varying_args(full = FALSE)
rand_forest() %>%
  set_engine(
    "randomForest",
    strata = Class,
    sampsize = varying()
  ) %>%
  varying_args()
```

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