

# Package ‘recipes’

March 18, 2020

**Title** Preprocessing Tools to Create Design Matrices

**Version** 0.1.10

**Description** An extensible framework to create and preprocess design matrices. Recipes consist of one or more data manipulation and analysis “steps”. Statistical parameters for the steps can be estimated from an initial data set and then applied to other data sets. The resulting design matrices can then be used as inputs into statistical or machine learning models.

**URL** <https://github.com/tidymodels/recipes>,  
<https://tidymodels.github.io/recipes/>

**BugReports** <https://github.com/tidymodels/recipes/issues>

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add_step	<i>Add a New Operation to the Current Recipe</i>
----------	--

---

### Description

add\_step adds a step to the last location in the recipe. add\_check does the same for checks.

### Usage

```
add_step(rec, object)
```

```
add_check(rec, object)
```

### Arguments

rec	A <a href="#">recipe()</a> .
object	A step or check object.

### Value

A updated [recipe\(\)](#) with the new operation in the last slot.

---

bake	<i>Apply a Trained Data Recipe</i>
------	------------------------------------

---

### Description

For a recipe with at least one preprocessing operations that has been trained by [prep.recipe\(\)](#), apply the computations to new data.

### Usage

```
bake(object, ...)
```

```
## S3 method for class 'recipe'
bake(object, new_data = NULL, ..., composition = "tibble")
```

## Arguments

object	A trained object such as a <code>recipe()</code> with at least one preprocessing operation.
...	One or more selector functions to choose which variables will be returned by the function. See <code>selections()</code> for more details. If no selectors are given, the default is to use <code>everything()</code> .
new_data	A data frame or tibble for whom the preprocessing will be applied.
composition	Either "tibble", "matrix", "data.frame", or "dgCMatrix" for the format of the processed data set. Note that all computations during the baking process are done in a non-sparse format. Also, note that this argument should be called <b>after</b> any selectors and the selectors should only resolve to numeric columns (otherwise an error is thrown).

## Details

`bake()` takes a trained recipe and applies the operations to a data set to create a design matrix.

If the original data used to train the data are to be processed, time can be saved by using the `retain = TRUE` option of `prep()` to avoid duplicating the same operations. With this option set, `juice()` can be used instead of `bake` with `new_data` equal to the training set.

Also, any steps with `skip = TRUE` will not be applied to the data when `bake` is invoked. `juice()` will always have all of the steps applied.

## Value

A tibble, matrix, or sparse matrix that may have different columns than the original columns in `new_data`.

## Author(s)

Max Kuhn

## See Also

`recipe()`, `juice()`, `prep()`

---

check\_class

*Check Variable Class*

---

## Description

`check_class` creates a *specification* of a recipe check that will check if a variable is of a designated class.

**Usage**

```

check_class(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  class_nm = NULL,
  allow_additional = FALSE,
  skip = FALSE,
  class_list = NULL,
  id = rand_id("class")
)

## S3 method for class 'check_class'
tidy(x, ...)

```

**Arguments**

recipe	A recipe object. The check will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the check. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this check since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
class_nm	A character vector that will be used in <code>inherits</code> to check the class. If NULL the classes will be learned in prep. Can contain more than one class.
allow_additional	If TRUE a variable is allowed to have additional classes to the one(s) that are checked.
skip	A logical. Should the check be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations.
class_list	A named list of column classes. This is NULL until computed by <code>prep.recipe()</code> .
id	A character string that is unique to this step to identify it.
x	A <code>check_class</code> object.

**Details**

This function can check the classes of the variables in two ways. When the `class` argument is provided it will check if all the variables specified are of the given class. If this argument is NULL, the check will learn the classes of each of the specified variables in prep. Both ways will break bake if the variables are not of the requested class. If a variable has multiple classes in prep, all the classes are checked. Please note that in prep the argument `strings_as_factors` defaults to TRUE.

If the train set contains character variables the check will be break bake when strings\_as\_factors is TRUE.

### Value

An updated version of recipe with the new check added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms (the selectors or variables selected) and value (the type).

### See Also

[recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

### Examples

```
library(dplyr)
library(modeldata)
data(okc)

# Learn the classes on the train set
train <- okc[1:1000, ]
test <- okc[1001:2000, ]
recipe(train, age ~ .) %>%
  check_class(everything()) %>%
  prep(train, strings_as_factors = FALSE) %>%
  bake(test)

# Manual specification
recipe(train, age ~ .) %>%
  check_class(age, class_nm = "integer") %>%
  check_class(diet, location, class_nm = "character") %>%
  check_class(date, class_nm = "Date") %>%
  prep(train, strings_as_factors = FALSE) %>%
  bake(test)

# By default only the classes that are specified
# are allowed.
x_df <- tibble(time = c(Sys.time() - 60, Sys.time()))
x_df$time %>% class()
## Not run:
recipe(x_df) %>%
  check_class(time, class_nm = "POSIXt") %>%
  prep(x_df) %>%
  bake_(x_df)

## End(Not run)

# Use allow_additional = TRUE if you are fine with it
recipe(x_df) %>%
  check_class(time, class_nm = "POSIXt", allow_additional = TRUE) %>%
  prep(x_df) %>%
  bake(x_df)
```

---

 check\_cols

*Check if all Columns are Present*


---

### Description

check\_cols creates a *specification* of a recipe step that will check if all the columns of the training frame are present in the new data.

### Usage

```
check_cols(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  skip = FALSE,
  id = rand_id("cols")
)

## S3 method for class 'check_cols'
tidy(x, ...)
```

### Arguments

recipe	A recipe object. The check will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are checked in the check See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this check since no new variables are created.
trained	A logical for whether the selectors in ... have been resolved by <a href="#">prep()</a> .
skip	A logical. Should the check be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations.
id	A character string that is unique to this step to identify it.
x	A check_cols object.

### Details

This check will break the bake function if any of the specified columns is not present in the data. If the check passes, nothing is changed to the data.



## Examples

```
library(modeldata)
data(biomass)

biomass_rec <- recipe(HHV ~ ., data = biomass) %>%
  step_rm(sample, dataset) %>%
  check_cols(contains("gen")) %>%
  step_center(all_predictors())

## Not run:
bake(biomass_rec, biomass[, c("carbon", "HHV")])

## End(Not run)
```

---

check\_missing

*Check for Missing Values*

---

## Description

check\_missing creates a *specification* of a recipe operation that will check if variables contain missing values.

## Usage

```
check_missing(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  columns = NULL,
  skip = FALSE,
  id = rand_id("missing")
)

## S3 method for class 'check_missing'
tidy(x, ...)
```

## Arguments

recipe	A recipe object. The check will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are checked in the check See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this check since no new variables are created.
trained	A logical for whether the selectors in ... have been resolved by <a href="#">prep()</a> .

columns	A character string of variable names that will be populated (eventually) by the terms argument.
skip	A logical. Should the check be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations.
id	A character string that is unique to this step to identify it.
x	A <code>check_missing</code> object.

### Details

This check will break the bake function if any of the checked columns does contain NA values. If the check passes, nothing is changed to the data.

### Value

An updated version of recipe with the new check added to the sequence of existing operations (if any). For the tidy method, a tibble with columns terms (the selectors or variables selected).

### Examples

```
library(modeldata)
data(credit_data)
is.na(credit_data) %>% colSums()

# If the test passes, `new_data` is returned unaltered
recipe(credit_data) %>%
  check_missing(Age, Expenses) %>%
  prep() %>%
  bake(credit_data)

# If your training set doesn't pass, prep() will stop with an error

## Not run:
recipe(credit_data) %>%
  check_missing(Income) %>%
  prep()

## End(Not run)

# If `new_data` contain missing values, the check will stop bake()

train_data <- credit_data %>% dplyr::filter(Income > 150)
test_data <- credit_data %>% dplyr::filter(Income <= 150 | is.na(Income))

rp <- recipe(train_data) %>%
  check_missing(Income) %>%
  prep()

bake(rp, train_data)
```

```
## Not run:
bake(rp, test_data)

## End(Not run)
```

---

check\_new\_values      *Check for New Values*

---

## Description

check\_new\_values creates a *specification* of a recipe operation that will check if variables contain new values.

## Usage

```
check_new_values(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  columns = NULL,
  ignore_NA = TRUE,
  values = NULL,
  skip = FALSE,
  id = rand_id("new_values")
)
```

## Arguments

recipe	A recipe object. The check will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are checked in the check. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this check since no new variables are created.
trained	A logical for whether the selectors in ... have been resolved by <a href="#">prep()</a> .
columns	A character string of variable names that will be populated (eventually) by the terms argument.
ignore_NA	A logical that indicates if we should consider missing values as value or not. Defaults to TRUE.
values	A named list with the allowed values. This is NULL until computed by <a href="#">prep.recipe()</a> .
skip	A logical. Should the check be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations.
id	A character string that is unique to this step to identify it.

## Details

This check will break the bake function if any of the checked columns does contain values it did not contain when prep was called on the recipe. If the check passes, nothing is changed to the data.

## Value

An updated version of recipe with the new check added to the sequence of existing operations (if any). For the tidy method, a tibble with columns terms (the selectors or variables selected).

## Examples

```
library(modeldata)
data(credit_data)

# If the test passes, `new_data` is returned unaltered
recipe(credit_data) %>%
  check_new_values(Home) %>%
  prep() %>%
  bake(new_data = credit_data)

# If `new_data` contains values not in `x` at the `prep()` function,
# the `bake()` function will break.
## Not run:
recipe(credit_data %>% dplyr::filter(Home != "rent")) %>%
  check_new_values(Home) %>%
  prep() %>%
  bake(new_data = credit_data)

## End(Not run)

# By default missing values are ignored, so this passes.
recipe(credit_data %>% dplyr::filter(!is.na(Home))) %>%
  check_new_values(Home) %>%
  prep() %>%
  bake(credit_data)

# Use `ignore_NA = FALSE` if you consider missing values as a value,
# that should not occur when not observed in the train set.
## Not run:
recipe(credit_data %>% dplyr::filter(!is.na(Home))) %>%
  check_new_values(Home, ignore_NA = FALSE) %>%
  prep() %>%
  bake(credit_data)

## End(Not run)
```

**Description**

check\_range creates a *specification* of a recipe check that will check if the range of a numeric variable changed in the new data.

**Usage**

```
check_range(
  recipe,
  ...,
  role = NA,
  skip = FALSE,
  trained = FALSE,
  slack_prop = 0.05,
  warn = FALSE,
  lower = NULL,
  upper = NULL,
  id = rand_id("range_check_")
)

## S3 method for class 'check_range'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The check will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the check. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this check since no new variables are created.
skip	A logical. Should the check be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
slack_prop	The allowed slack as a proportion of the range of the variable in the train set.
warn	If TRUE the check will throw a warning instead of an error when failing.
lower	A named numeric vector of minimum values in the train set. This is NULL until computed by <a href="#">prep.recipe()</a> .
upper	A named numeric vector of maximum values in the train set. This is NULL until computed by <a href="#">prep.recipe()</a> .
id	A character string that is unique to this step to identify it.
x	A check_range object.

## Details

The amount of slack that is allowed is determined by the `slack_prop`. This is a numeric of length one or two. If of length one, the same proportion will be used at both ends of the train set range. If of length two, its first value is used to compute the allowed slack at the lower end, the second to compute the allowed slack at the upper end.

## Value

An updated version of `recipe` with the new check added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `value` (the means).

## See Also

[recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

## Examples

```
slack_df <- data_frame(x = 0:100)
slack_new_data <- data_frame(x = -10:110)

# this will fail the check both ends
## Not run:
recipe(slack_df) %>%
  check_range(x) %>%
  prep() %>%
  bake(slack_new_data)

## End(Not run)

# this will fail the check only at the upper end
## Not run:
recipe(slack_df) %>%
  check_range(x, slack_prop = c(0.1, 0.05)) %>%
  prep() %>%
  bake(slack_new_data)

## End(Not run)

# give a warning instead of an error
## Not run:
recipe(slack_df) %>%
  check_range(x, warn = TRUE) %>%
  prep() %>%
  bake(slack_new_data)

## End(Not run)
```

---

detect_step	<i>Detect if a particular step or check is used in a recipe</i>
-------------	---

---

**Description**

Detect if a particular step or check is used in a recipe

**Usage**

```
detect_step(recipe, name)
```

**Arguments**

recipe	A recipe to check.
name	Character name of a step or check, omitted the prefix. That is, to check if step_intercept is present, use name = intercept.

**Value**

Logical indicating if recipes contains given step.

**Examples**

```
rec <- recipe(Species ~ ., data = iris) %>%
  step_intercept()

detect_step(rec, "step_intercept")
```

---

discretize	<i>Discretize Numeric Variables</i>
------------	-------------------------------------

---

**Description**

discretize converts a numeric vector into a factor with bins having approximately the same number of data points (based on a training set).

**Usage**

```
discretize(x, ...)

## Default S3 method:
discretize(x, ...)

## S3 method for class 'numeric'
discretize(
  x,
```

```

cuts = 4,
labels = NULL,
prefix = "bin",
keep_na = TRUE,
infs = TRUE,
min_unique = 10,
...
)

## S3 method for class 'discretize'
predict(object, new_data, ...)

```

### Arguments

<code>x</code>	A numeric vector
<code>...</code>	Options to pass to <code>stats::quantile()</code> that should not include <code>x</code> or <code>probs</code> .
<code>cuts</code>	An integer defining how many cuts to make of the data.
<code>labels</code>	A character vector defining the factor levels that will be in the new factor (from smallest to largest). This should have length <code>cuts+1</code> and should not include a level for missing (see <code>keep_na</code> below).
<code>prefix</code>	A single parameter value to be used as a prefix for the factor levels (e.g. <code>bin1</code> , <code>bin2</code> , ...). If the string is not a valid R name, it is coerced to one.
<code>keep_na</code>	A logical for whether a factor level should be created to identify missing values in <code>x</code> .
<code>infs</code>	A logical indicating whether the smallest and largest cut point should be infinite.
<code>min_unique</code>	An integer defining a sample size line of dignity for the binning. If (the number of unique values)/(cuts+1) is less than <code>min_unique</code> , no discretization takes place.
<code>object</code>	An object of class <code>discretize</code> .
<code>new_data</code>	A new numeric object to be binned.

### Details

`discretize` estimates the cut points from `x` using percentiles. For example, if `cuts = 3`, the function estimates the quartiles of `x` and uses these as the cut points. If `cuts = 2`, the bins are defined as being above or below the median of `x`.

The `predict` method can then be used to turn numeric vectors into factor vectors.

If `keep_na = TRUE`, a suffix of `"_missing"` is used as a factor level (see the examples below).

If `infs = FALSE` and a new value is greater than the largest value of `x`, a missing value will result.

### Value

`discretize` returns an object of class `discretize` and `predict.discretize` returns a factor vector.



**Examples**

```

library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

median(biomass_tr$carbon)
discretize(biomass_tr$carbon, cuts = 2)
discretize(biomass_tr$carbon, cuts = 2, infs = FALSE)
discretize(biomass_tr$carbon, cuts = 2, infs = FALSE, keep_na = FALSE)
discretize(biomass_tr$carbon, cuts = 2, prefix = "maybe a bad idea to bin")

carbon_binned <- discretize(biomass_tr$carbon)
table(predict(carbon_binned, biomass_tr$carbon))

carbon_no_infs <- discretize(biomass_tr$carbon, infs = FALSE)
predict(carbon_no_infs, c(50, 100))

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)
rec <- rec %>% step_discretize(carbon, hydrogen)
rec <- prep(rec, biomass_tr)
binned_te <- bake(rec, biomass_te)
table(binned_te$carbon)

```

---

formula.recipe

*Create a Formula from a Prepared Recipe*


---

**Description**

In case a model formula is required, the formula method can be used on a recipe to show what predictors and outcome(s) could be used.

**Usage**

```

## S3 method for class 'recipe'
formula(x, ...)

```

**Arguments**

x                    A recipe object where all steps have been prepared.  
...                   Note currently used.

**Value**

A formula.

## Examples

```
formula(recipe(Species + Sepal.Length ~ ., data = iris))

iris_rec <- recipe(Species ~ ., data = iris) %>%
  step_center(all_numeric()) %>%
  prep(training = iris)
formula(iris_rec)
```

---

fully_trained	<i>Check to see if a recipe is trained/prepared</i>
---------------	---

---

## Description

Check to see if a recipe is trained/prepared

## Usage

```
fully_trained(x)
```

## Arguments

x                    A recipe

## Value

A logical which is true if all of the recipe steps have been run through prep. If no steps have been added to the recipe, TRUE is returned.

## Examples

```
rec <- recipe(Species ~ ., data = iris) %>%
  step_center(all_numeric())

rec %>% fully_trained

rec %>% prep(training = iris) %>% fully_trained
```

---

has_role	<i>Role Selection</i>
----------	-----------------------

---

### Description

has\_role(), all\_predictors(), and all\_outcomes() can be used to select variables in a formula that have certain roles. Similarly, has\_type(), all\_numeric(), and all\_nominal() are used to select columns based on their data type.

See ?selections for more details.

current\_info() is an internal function.

All of these functions have limited utility outside of column selection in step functions.

### Usage

```
has_role(match = "predictor")
```

```
all_predictors()
```

```
all_outcomes()
```

```
has_type(match = "numeric")
```

```
all_numeric()
```

```
all_nominal()
```

```
current_info()
```

### Arguments

match            A single character string for the query. Exact matching is used (i.e. regular expressions won't work).

### Value

Selector functions return an integer vector.

current\_info() returns an environment with objects vars and data.

### Examples

```
library(modeldata)
data(biomass)

rec <- recipe(biomass) %>%
  update_role(
    carbon, hydrogen, oxygen, nitrogen, sulfur,
    new_role = "predictor"
```

```

) %>%
  update_role(HHV, new_role = "outcome") %>%
  update_role(sample, new_role = "id variable") %>%
  update_role(dataset, new_role = "splitting indicator")

recipe_info <- summary(rec)
recipe_info

# Centering on all predictors except carbon
rec %>%
  step_center(all_predictors(), -carbon) %>%
  prep(training = biomass) %>%
  juice()

```

juice

*Extract Finalized Training Set***Description**

As steps are estimated by prep, these operations are applied to the training set. Rather than running bake to duplicate this processing, this function will return variables from the processed training set.

**Usage**

```
juice(object, ..., composition = "tibble")
```

**Arguments**

object	A recipe object that has been prepared with the option retain = TRUE.
...	One or more selector functions to choose which variables will be returned by the function. See <a href="#">selections()</a> for more details. If no selectors are given, the default is to use <a href="#">everything()</a> .
composition	Either "tibble", "matrix", "data.frame", or "dgCMatrix" for the format of the processed data set. Note that all computations during the baking process are done in a non-sparse format. Also, note that this argument should be called <b>after</b> any selectors and the selectors should only resolve to numeric columns (otherwise an error is thrown).

**Details**

When preparing a recipe, if the training data set is retained using retain = TRUE, there is no need to bake the recipe to get the preprocessed training set.

juice will return the results of a recipes where *all steps* have been applied to the data, irrespective of the value of the step's skip argument.

**See Also**

[recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```

library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

sp_signed <- rec %>%
  step_normalize(all_predictors()) %>%
  step_spatialsign(all_predictors())

sp_signed_trained <- prep(sp_signed, training = biomass_tr)

tr_values <- bake(sp_signed_trained, new_data = biomass_tr, all_predictors())
og_values <- juice(sp_signed_trained, all_predictors())

all.equal(tr_values, og_values)

```

names0

*Naming Tools***Description**

names0 creates a series of num names with a common prefix. The names are numbered with leading zeros (e.g. prefix01-prefix10 instead of prefix1-prefix10). dummy\_names can be used for renaming unordered and ordered dummy variables (in [step\\_dummy\(\)](#)).

**Usage**

```

names0(num, prefix = "x")

dummy_names(var, lvl, ordinal = FALSE, sep = "_")

```

**Arguments**

num	A single integer for how many elements are created.
prefix	A character string that will start each name.
var	A single string for the original factor name.
lvl	A character vectors of the factor levels (in order). When used with <a href="#">step_dummy()</a> , lvl would be the suffixes that result <i>after</i> <code>model.matrix</code> is called (see the example below).
ordinal	A logical; was the original factor ordered?
sep	A single character value for the separator between the names and levels.

**Value**

`names0` returns a character string of length `num` and `dummy_names` generates a character vector the same length as `lvl`,

**Examples**

```
names0(9, "x")
names0(10, "x")

example <- data.frame(y = ordered(letters[1:5]),
                     z = factor(LETTERS[1:5]))

dummy_names("z", levels(example$z)[-1])

after_mm <- colnames(model.matrix(~y, data = example))[-1]
after_mm
levels(example$y)

dummy_names("y", substring(after_mm, 2), ordinal = TRUE)
```

---

 prep

*Train a Data Recipe*


---

**Description**

For a recipe with at least one preprocessing operation, estimate the required parameters from a training set that can be later applied to other data sets.

**Usage**

```
prep(x, ...)

## S3 method for class 'recipe'
prep(
  x,
  training = NULL,
  fresh = FALSE,
  verbose = FALSE,
  retain = TRUE,
  strings_as_factors = TRUE,
  ...
)
```

**Arguments**

`x` an object  
 ... further arguments passed to or from other methods (not currently used).

training	A data frame or tibble that will be used to estimate parameters for preprocessing.
fresh	A logical indicating whether already trained operation should be re-trained. If TRUE, you should pass in a data set to the argument training.
verbose	A logical that controls whether progress is reported as operations are executed.
retain	A logical: should the <i>preprocessed</i> training set be saved into the template slot of the recipe after training? This is a good idea if you want to add more steps later but want to avoid re-training the existing steps. Also, it is advisable to use retain = TRUE if any steps use the option skip = FALSE. <b>Note</b> that this can make the final recipe size large. When verbose = TRUE, a message is written with the approximate object size in memory but may be an underestimate since it does not take environments into account.
strings_as_factors	A logical: should character columns be converted to factors? This affects the preprocessed training set (when retain = TRUE) as well as the results of bake.recipe.

### Details

Given a data set, this function estimates the required quantities and statistics required by any operations.

`prep()` returns an updated recipe with the estimates.

Note that missing data handling is handled in the steps; there is no global `na.rm` option at the recipe-level or in `prep()`.

Also, if a recipe has been trained using `prep()` and then steps are added, `prep()` will only update the new operations. If `fresh = TRUE`, all of the operations will be (re)estimated.

As the steps are executed, the training set is updated. For example, if the first step is to center the data and the second is to scale the data, the step for scaling is given the centered data.

### Value

A recipe whose step objects have been updated with the required quantities (e.g. parameter estimates, model objects, etc). Also, the `term_info` object is likely to be modified as the operations are executed.

### Author(s)

Max Kuhn

---



*Wrapper function for preparing recipes within resampling*


---

### Description

When working with the **rsample** package, a simple recipe must be *prepared* using the `prep` function first. When using recipes with **rsample** it is helpful to have a function that can prepare a recipe across a series of `split` objects that are produced in this package. `prepper` is a wrapper function around `prep` that can be used to do this. See the vignette on "Recipes and `rsample`" for an example.

**Usage**

```
prepper(split_obj, recipe, ...)
```

**Arguments**

split_obj	An rplit object
recipe	An untrained recipe object.
...	Arguments to pass to prep such as verbose or retain.

**Details**

prepper() sets the underlying prep() argument fresh to TRUE.

---

print.recipe	<i>Print a Recipe</i>
--------------	-----------------------

---

**Description**

Print a Recipe

**Usage**

```
## S3 method for class 'recipe'
print(x, form_width = 30, ...)
```

**Arguments**

x	A recipe object
form_width	The number of characters used to print the variables or terms in a formula
...	further arguments passed to or from other methods (not currently used).

**Value**

The original object (invisibly)

**Author(s)**

Max Kuhn



---

recipe	<i>Create a Recipe for Preprocessing Data</i>
--------	---

---

### Description

A recipe is a description of what steps should be applied to a data set in order to get it ready for data analysis.

### Usage

```
recipe(x, ...)

## Default S3 method:
recipe(x, ...)

## S3 method for class 'data.frame'
recipe(x, formula = NULL, ..., vars = NULL, roles = NULL)

## S3 method for class 'formula'
recipe(formula, data, ...)

## S3 method for class 'matrix'
recipe(x, ...)
```

### Arguments

x, data	A data frame or tibble of the <i>template</i> data set (see below).
...	Further arguments passed to or from other methods (not currently used).
formula	A model formula. No in-line functions should be used here (e.g. $\log(x)$ , $x:y$ , etc.) and minus signs are not allowed. These types of transformations should be enacted using step functions in this package. Dots are allowed as are simple multivariate outcome terms (i.e. no need for <code>cbind</code> ; see Examples).
vars	A character string of column names corresponding to variables that will be used in any context (see below)
roles	A character string (the same length of vars) that describes a single role that the variable will take. This value could be anything but common roles are "outcome", "predictor", "case_weight", or "ID"

### Details

Recipes are alternative methods for creating design matrices and for preprocessing data.

Variables in recipes can have any type of *role* in subsequent analyses such as: outcome, predictor, case weights, stratification variables, etc.

recipe objects can be created in several ways. If the analysis only contains outcomes and predictors, the simplest way to create one is to use a simple formula (e.g.  $y \sim x_1 + x_2$ ) that does not contain inline functions such as  $\log(x_3)$ . An example is given below.

Alternatively, a recipe object can be created by first specifying which variables in a data set should be used and then sequentially defining their roles (see the last example).

There are two different types of operations that can be sequentially added to a recipe. **Steps** can include common operations like logging a variable, creating dummy variables or interactions and so on. More computationally complex actions such as dimension reduction or imputation can also be specified. **Checks** are operations that conduct specific tests of the data. When the test is satisfied, the data are returned without issue or modification. Otherwise, any error is thrown.

Once a recipe has been defined, the `prep()` function can be used to estimate quantities required for the operations using a data set (a.k.a. the training data). `prep()` returns another recipe.

To apply the recipe to a data set, the `bake()` function is used in the same manner as `predict` would be for models. This applies the steps to any data set.

Note that the data passed to `recipe` need not be the complete data that will be used to train the steps (by `prep()`). The recipe only needs to know the names and types of data that will be used. For large data sets, `head` could be used to pass the recipe a smaller data set to save time and memory.

## Value

An object of class `recipe` with sub-objects:

<code>var_info</code>	A tibble containing information about the original data set columns
<code>term_info</code>	A tibble that contains the current set of terms in the data set. This initially defaults to the same data contained in <code>var_info</code> .
<code>steps</code>	A list of step or check objects that define the sequence of preprocessing operations that will be applied to data. The default value is <code>NULL</code>
<code>template</code>	A tibble of the data. This is initialized to be the same as the data given in the data argument but can be different after the recipe is trained.

## Author(s)

Max Kuhn

## Examples

```
#####
# simple example:
library(modeldata)
data(biomass)

# split data
biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

# When only predictors and outcomes, a simplified formula can be used.
rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

# Now add preprocessing steps to the recipe.
```

```

sp_signed <- rec %>%
  step_normalize(all_predictors()) %>%
  step_spatialsign(all_predictors())
sp_signed

# now estimate required parameters
sp_signed_trained <- prep(sp_signed, training = biomass_tr)
sp_signed_trained

# apply the preprocessing to a data set
test_set_values <- bake(sp_signed_trained, new_data = biomass_te)

# or use pipes for the entire workflow:
rec <- biomass_tr %>%
  recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur) %>%
  step_normalize(all_predictors()) %>%
  step_spatialsign(all_predictors())

#####
# multivariate example

# no need for `cbind(carbon, hydrogen)` for left-hand side
multi_y <- recipe(carbon + hydrogen ~ oxygen + nitrogen + sulfur,
                  data = biomass_tr)
multi_y <- multi_y %>%
  step_center(all_outcomes()) %>%
  step_scale(all_predictors())

multi_y_trained <- prep(multi_y, training = biomass_tr)

results <- bake(multi_y_trained, biomass_te)

#####
# Creating a recipe manually with different roles

rec <- recipe(biomass_tr) %>%
  update_role(carbon, hydrogen, oxygen, nitrogen, sulfur,
              new_role = "predictor") %>%
  update_role(HHV, new_role = "outcome") %>%
  update_role(sample, new_role = "id variable") %>%
  update_role(dataset, new_role = "splitting indicator")
rec

```

## Description

The recipes package can be used to create design matrices for modeling and to conduct preprocessing of variables. It is meant to be a more extensive framework than R's formula method. Some differences between simple formula methods and recipes are that

1. Variables can have arbitrary roles in the analysis beyond predictors and outcomes.
2. A recipe consists of one or more steps that define actions on the variables.
3. Recipes can be defined sequentially using pipes as well as being modifiable and extensible.

### Basic Functions

The three main functions are `recipe()`, `prep()`, and `bake()`.

`recipe()` defines the operations on the data and the associated roles. Once the preprocessing steps are defined, any parameters are estimated using `prep()`. Once the data are ready for transformation, the `bake()` function applies the operations.

### Step Functions

These functions are used to add new actions to the recipe and have the naming convention "step\_action". For example, `step_center()` centers the data to have a zero mean and `step_dummy()` is used to create dummy variables.

---

roles

*Manually Alter Roles*

---

### Description

`update_role()` alters an existing role in the recipe or assigns an initial role to variables that do not yet have a declared role.

`add_role()` adds an *additional* role to variables that already have a role in the recipe. It does not overwrite old roles, as a single variable can have multiple roles.

`remove_role()` eliminates a single existing role in the recipe.

### Usage

```
add_role(recipe, ..., new_role = "predictor", new_type = NULL)
```

```
update_role(recipe, ..., new_role = "predictor", old_role = NULL)
```

```
remove_role(recipe, ..., old_role)
```

### Arguments

<code>recipe</code>	An existing <code>recipe()</code> .
<code>...</code>	One or more selector functions to choose which variables are being assigned a role. See <code>selections()</code> for more details.
<code>new_role</code>	A character string for a single role.
<code>new_type</code>	A character string for specific type that the variable should be identified as. If left as <code>NULL</code> , the type is automatically identified as the <i>first</i> type you see for that variable in <code>summary(recipe)</code> .

`old_role` A character string for the specific role to update for the variables selected by `update_role()` accepts a NULL as long as the variables have only a single role.

### Details

`update_role()` should be used when a variable doesn't currently have a role in the recipe, or to replace an `old_role` with a `new_role`. `add_role()` only adds additional roles to variables that already have roles and will throw an error when the current role is missing (i.e. NA).

When using `add_role()`, if a variable is selected that already has the `new_role`, a warning is emitted and that variable is skipped so no duplicate roles are added.

Adding or updating roles is a useful way to group certain variables that don't fall in the standard "predictor" bucket. You can perform a step on all of the variables that have a custom role with the selector `has_role()`.

### Value

An updated recipe object.

### Examples

```
library(recipes)
library(modeldata)
data(biomass)

# Using the formula method, roles are created for any outcomes and predictors:
recipe(HHV ~ ., data = biomass) %>%
  summary()

# However `sample` and `dataset` aren't predictors. Since they already have
# roles, `update_role()` can be used to make changes:
recipe(HHV ~ ., data = biomass) %>%
  update_role(sample, new_role = "id variable") %>%
  update_role(dataset, new_role = "splitting variable") %>%
  summary()

# `update_role()` cannot set a role to NA, use `remove_role()` for that
## Not run:
recipe(HHV ~ ., data = biomass) %>%
  update_role(sample, new_role = NA_character_)

## End(Not run)

# -----

# Variables can have more than one role. `add_role()` can be used
# if the column already has at least one role:
recipe(HHV ~ ., data = biomass) %>%
  add_role(carbon, sulfur, new_role = "something") %>%
  summary()
```

```

# `update_role()` has an argument called `old_role` that is required to
# unambiguously update a role when the column currently has multiple roles.
recipe(HHV ~ ., data = biomass) %>%
  add_role(carbon, new_role = "something") %>%
  update_role(carbon, new_role = "something else", old_role = "something") %>%
  summary()

# `carbon` has two roles at the end, so the last `update_roles()` fails since
# `old_role` was not given.
## Not run:
recipe(HHV ~ ., data = biomass) %>%
  add_role(carbon, sulfur, new_role = "something") %>%
  update_role(carbon, new_role = "something else")

## End(Not run)

# -----

# To remove a role, `remove_role()` can be used to remove a single role.
recipe(HHV ~ ., data = biomass) %>%
  add_role(carbon, new_role = "something") %>%
  remove_role(carbon, old_role = "something") %>%
  summary()

# To remove all roles, call `remove_role()` multiple times to reset to `NA`
recipe(HHV ~ ., data = biomass) %>%
  add_role(carbon, new_role = "something") %>%
  remove_role(carbon, old_role = "something") %>%
  remove_role(carbon, old_role = "predictor") %>%
  summary()

# -----

# If the formula method is not used, all columns have a missing role:
recipe(biomass) %>%
  summary()

```

---

 selections

*Methods for Select Variables in Step Functions*


---

## Description

When selecting variables or model terms in step functions, dplyr-like tools are used. The *selector* functions can choose variables based on their name, current role, data type, or any combination of these. The selectors are passed as any other argument to the step. If the variables are explicitly stated in the step function, this might be similar to:

```

recipe( ~ ., data = USArrests) %>%
  step_pca(Murder, Assault, UrbanPop, Rape, num = 3)

```

The first four arguments indicate which variables should be used in the PCA while the last argument is a specific argument to `step_pca()`.

Note that:

1. The selector arguments should not contain functions beyond those supported (see below).
2. These arguments are not evaluated until the prep function for the step is executed.
3. The dplyr-like syntax allows for negative signs to exclude variables (e.g. `-Murder`) and the set of selectors will be processed in order.
4. A leading exclusion in these arguments (e.g. `-Murder`) has the effect of adding all variables to the list except the excluded variable(s).

Also, select helpers from the `tidyselect` package can also be used: `tidyselect::starts_with()`, `tidyselect::ends_with()`, `tidyselect::contains()`, `tidyselect::matches()`, `tidyselect::num_range()`, `tidyselect::everything()`, and `tidyselect::one_of()`. For example:

```
recipe(Species ~ ., data = iris) %>%
  step_center(starts_with("Sepal"), -contains("Width"))
```

would only select `Sepal.Length`

**Inline** functions that specify computations, such as `log(x)`, should not be used in selectors and will produce an error. A list of allowed selector functions is below.

Columns of the design matrix that may not exist when the step is coded can also be selected. For example, when using `step_pca`, the number of columns created by feature extraction may not be known when subsequent steps are defined. In this case, using `matches("^PC")` will select all of the columns whose names start with "PC" *once those columns are created*.

There are sets of functions that can be used to select variables based on their role or type: `has_role()` and `has_type()`. For convenience, there are also functions that are more specific: `all_numeric()`, `all_nominal()`, `all_predictors()`, and `all_outcomes()`. These can be used in conjunction with the previous functions described for selecting variables using their names:

```
data(biomass)
recipe(HHV ~ ., data = biomass) %>%
  step_center(all_numeric(), -all_outcomes())
```

This results in all the numeric predictors: carbon, hydrogen, oxygen, nitrogen, and sulfur.

If a role for a variable has not been defined, it will never be selected using role-specific selectors.

Selectors can be used in `step_interact()` in similar ways but must be embedded in a model formula (as opposed to a sequence of selectors). For example, the interaction specification could be `~ starts_with("Species"):Sepal.Width`. This can be useful if `Species` was converted to dummy variables previously using `step_dummy()`.

The complete list of allowable functions in steps:

- **By name:** `tidyselect::starts_with()`, `tidyselect::ends_with()`, `tidyselect::contains()`, `tidyselect::matches()`, `tidyselect::num_range()`, and `tidyselect::everything()`
- **By role:** `has_role()`, `all_predictors()`, and `all_outcomes()`
- **By type:** `has_type()`, `all_numeric()`, and `all_nominal()`

---

step_arrange	<i>Sort rows using dplyr</i>
--------------	------------------------------

---

### Description

step\_arrange creates a *specification* of a recipe step that will sort rows using `dplyr::arrange()`.

### Usage

```
step_arrange(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  inputs = NULL,
  skip = FALSE,
  id = rand_id("arrange")
)

## S3 method for class 'step_arrange'
tidy(x, ...)
```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	Comma separated list of unquoted variable names. Use <code>desc()</code> to sort a variable in descending order. See <code>[dplyr::arrange()]</code> for more details. For the <code>tidy</code> method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
inputs	Quosure of values given by ...
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_arrange object

### Details

When an object in the user's global environment is referenced in the expression defining the new variable(s), it is a good idea to use quasiquotation (e.g. `!!!`) to embed the value of the object in the expression (to be portable between sessions). See the examples.



**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms which contains the sorting variable(s) or expression(s). The expressions are text representations and are not parsable.

**Examples**

```

rec <- recipe( ~ ., data = iris) %>%
  step_arrange(desc(Sepal.Length), 1/Petal.Length)

prepped <- prep(rec, training = iris %>% slice(1:75))
tidy(prepped, number = 1)

library(dplyr)

dplyr_train <-
  iris %>%
  as_tibble() %>%
  slice(1:75) %>%
  dplyr::arrange(desc(Sepal.Length), 1/Petal.Length)

rec_train <- juice(prepped)
all.equal(dplyr_train, rec_train)

dplyr_test <-
  iris %>%
  as_tibble() %>%
  slice(76:150) %>%
  dplyr::arrange(desc(Sepal.Length), 1/Petal.Length)
rec_test <- bake(prepped, iris %>% slice(76:150))
all.equal(dplyr_test, rec_test)

# When you have variables/expressions, you can create a
# list of symbols with `rlang::syms()` and splice them in
# the call with `!!!`. See https://tidyeval.tidyverse.org

sort_vars <- c("Sepal.Length", "Petal.Length")

qq_rec <-
  recipe( ~ ., data = iris) %>%
  # Embed the `values` object in the call using !!!
  step_arrange(!!!syms(sort_vars)) %>%
  prep(training = iris)

tidy(qq_rec, number = 1)

```

## Description

step\_bagimpute creates a *specification* of a recipe step that will create bagged tree models to impute missing data.

## Usage

```
step_bagimpute(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  impute_with = imp_vars(all_predictors()),
  trees = 25,
  models = NULL,
  options = list(keepX = FALSE),
  seed_val = sample.int(10^4, 1),
  skip = FALSE,
  id = rand_id("bagimpute")
)

imp_vars(...)

## S3 method for class 'step_bagimpute'
tidy(x, ...)
```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose variables. For step_bagimpute, this indicates the variables to be imputed. When used with imp_vars, the dots indicates which variables are used to predict the missing data in each variable. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
impute_with	A call to imp_vars to specify which variables are used to impute the variables that can include specific variable names separated by commas or different selectors (see <a href="#">selections()</a> ). If a column is included in both lists to be imputed and to be an imputation predictor, it will be removed from the latter and not used to impute itself.
trees	An integer for the number bagged trees to use in each model.
models	The <code>ipred::ipredbag()</code> objects are stored here once this bagged trees have been trained by <code>prep.recipe()</code> .
options	A list of options to <code>ipred::ipredbag()</code> . Defaults are set for the arguments nbagg and keepX but others can be passed in. <b>Note</b> that the arguments X and y should not be passed here.

seed_val	A integer used to create reproducible models. The same seed is used across all imputation models.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_bagimpute</code> object.

### Details

For each variables requiring imputation, a bagged tree is created where the outcome is the variable of interest and the predictors are any other variables listed in the `impute_with` formula. One advantage to the bagged tree is that it can accept predictors that have missing values themselves. This imputation method can be used when the variable of interest (and predictors) are numeric or categorical. Imputed categorical variables will remain categorical. Also, integers will be imputed to integer too.

Note that if a variable that is to be imputed is also in `impute_with`, this variable will be ignored.

It is possible that missing values will still occur after imputation if a large majority (or all) of the imputing variables are also missing.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `model` (the bagged tree object).

### References

Kuhn, M. and Johnson, K. (2013). *Applied Predictive Modeling*. Springer Verlag.

### Examples

```
library(modeldata)
data("credit_data")

## missing data per column
vapply(credit_data, function(x) mean(is.na(x)), c(num = 0))

set.seed(342)
in_training <- sample(1:nrow(credit_data), 2000)

credit_tr <- credit_data[ in_training, ]
credit_te <- credit_data[-in_training, ]
missing_examples <- c(14, 394, 565)

rec <- recipe(Price ~ ., data = credit_tr)
## Not run:
```

```

impute_rec <- rec %>%
  step_bagimpute(Status, Home, Marital, Job, Income, Assets, Debt)

imp_models <- prep(impute_rec, training = credit_tr)

imputed_te <- bake(imp_models, new_data = credit_te, everything())

credit_te[missing_examples,]
imputed_te[missing_examples, names(credit_te)]

tidy(impute_rec, number = 1)
tidy(imp_models, number = 1)

## Specifying which variables to impute with

impute_rec <- rec %>%
  step_bagimpute(Status, Home, Marital, Job, Income, Assets, Debt,
                 impute_with = imp_vars(Time, Age, Expenses),
                 # for quick execution, nbagg lowered
                 options = list(nbagg = 5, keepX = FALSE))

imp_models <- prep(impute_rec, training = credit_tr)

imputed_te <- bake(imp_models, new_data = credit_te, everything())

credit_te[missing_examples,]
imputed_te[missing_examples, names(credit_te)]

tidy(impute_rec, number = 1)
tidy(imp_models, number = 1)

## End(Not run)

```

---

step\_bin2factor

---

*Create a Factors from A Dummy Variable*


---

### Description

step\_bin2factor creates a *specification* of a recipe step that will create a two-level factor from a single dummy variable.

### Usage

```

step_bin2factor(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  levels = c("yes", "no"),
  ref_first = TRUE,

```

```

    columns = NULL,
    skip = FALSE,
    id = rand_id("bin2factor")
  )

  ## S3 method for class 'step_bin2factor'
  tidy(x, ...)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	Selector functions that choose which variables will be converted. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
levels	A length 2 character string that indicate the factor levels for the 1's (in the first position) and the zeros (second)
ref_first	Logical. Should the first level, which replaces 1's, be the factor reference level?
columns	A vector with the selected variable names. This is NULL until computed by <a href="#">prep.recipe()</a> .
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_bin2factor</code> object.

### Details

This operation may be useful for situations where a binary piece of information may need to be represented as categorical instead of numeric. For example, naive Bayes models would do better to have factor predictors so that the binomial distribution is modeled in stead of a Gaussian probability density of numeric binary data. Note that the numeric data is only verified to be numeric (and does not count levels).

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns terms (the columns that will be affected).

### Examples

```

library(modeldata)
data(covers)

```

```

rec <- recipe(~ description, covers) %>%
  step_regex(description, pattern = "(rock|stony)", result = "rocks") %>%
  step_regex(description, pattern = "(rock|stony)", result = "more_rocks") %>%
  step_bin2factor(rocks)

tidy(rec, number = 3)

rec <- prep(rec, training = covers)
results <- bake(rec, new_data = covers)

table(results$rocks, results$more_rocks)

tidy(rec, number = 3)

```

---

step\_BoxCox

*Box-Cox Transformation for Non-Negative Data*


---

## Description

step\_BoxCox creates a *specification* of a recipe step that will transform data using a simple Box-Cox transformation.

## Usage

```

step_BoxCox(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  lambdas = NULL,
  limits = c(-5, 5),
  num_unique = 5,
  skip = FALSE,
  id = rand_id("BoxCox")
)

## S3 method for class 'step_BoxCox'
tidy(x, ...)

```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.

trained	A logical to indicate if the quantities for preprocessing have been estimated.
lambdas	A numeric vector of transformation values. This is NULL until computed by <code>prep.recipe()</code> .
limits	A length 2 numeric vector defining the range to compute the transformation parameter lambda.
num_unique	An integer where data that have less possible values will not be evaluate for a transformation.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_BoxCox</code> object.

### Details

The Box-Cox transformation, which requires a strictly positive variable, can be used to rescale a variable to be more similar to a normal distribution. In this package, the partial log-likelihood function is directly optimized within a reasonable set of transformation values (which can be changed by the user).

This transformation is typically done on the outcome variable using the residuals for a statistical model (such as ordinary least squares). Here, a simple null model (intercept only) is used to apply the transformation to the *predictor* variables individually. This can have the effect of making the variable distributions more symmetric.

If the transformation parameters are estimated to be very closed to the bounds, or if the optimization fails, a value of NA is used and no transformation is applied.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `value` (the lambda estimate).

### References

Sakia, R. M. (1992). The Box-Cox transformation technique: A review. *The Statistician*, 169-178..

### See Also

[step\\_YeoJohnson\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

### Examples

```
rec <- recipe(~ ., data = as.data.frame(state.x77))
```

```

bc_trans <- step_BoxCox(rec, all_numeric())

bc_estimates <- prep(bc_trans, training = as.data.frame(state.x77))

bc_data <- bake(bc_estimates, as.data.frame(state.x77))

plot(density(state.x77[, "Illiteracy"]), main = "before")
plot(density(bc_data$Illiteracy), main = "after")

tidy(bc_trans, number = 1)
tidy(bc_estimates, number = 1)

```

---

step\_bs

*B-Spline Basis Functions*


---

## Description

step\_bs creates a *specification* of a recipe step that will create new columns that are basis expansions of variables using B-splines.

## Usage

```

step_bs(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  deg_free = NULL,
  degree = 3,
  objects = NULL,
  options = list(),
  skip = FALSE,
  id = rand_id("bs")
)

## S3 method for class 'step_bs'
tidy(x, ...)

```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.



role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new columns created from the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
deg_free	The degrees of freedom.
degree	Degree of polynomial spline (integer).
objects	A list of <code>splines::bs()</code> objects created once the step has been trained.
options	A list of options for <code>splines::bs()</code> which should not include <code>x</code> , <code>degree</code> , or <code>df</code> .
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_bs</code> object.

### Details

`step_bs` can new features from a single variable that enable fitting routines to model this variable in a nonlinear manner. The extent of the possible nonlinearity is determined by the `df`, `degree`, or `knot` arguments of `splines::bs()`. The original variables are removed from the data and new columns are added. The naming convention for the new variables is `varname_bs_1` and so on.

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` which is the columns that will be affected and `holiday`.

### See Also

[step\\_poly\(\)](#) [recipe\(\)](#) [step\\_ns\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

### Examples

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

with_splines <- rec %>%
  step_bs(carbon, hydrogen)
with_splines <- prep(with_splines, training = biomass_tr)
```

```
expanded <- bake(with_splines, biomass_te)
expanded
```

---

step_center	<i>Centering numeric data</i>
-------------	-------------------------------

---

## Description

step\_center creates a *specification* of a recipe step that will normalize numeric data to have a mean of zero.

## Usage

```
step_center(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  means = NULL,
  na_rm = TRUE,
  skip = FALSE,
  id = rand_id("center")
)

## S3 method for class 'step_center'
tidy(x, ...)
```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
means	A named numeric vector of means. This is NULL until computed by <a href="#">prep.recipe()</a> .
na_rm	A logical value indicating whether NA values should be removed during computations.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_center object.

**Details**

Centering data means that the average of a variable is subtracted from the data. `step_center` estimates the variable means from the data used in the `training` argument of `prep.recipe`. `bake.recipe` then applies the centering to new data sets using these means.

**Value**

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `value` (the means).

**See Also**

[recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

center_trans <- rec %>%
  step_center(carbon, contains("gen"), -hydrogen)

center_obj <- prep(center_trans, training = biomass_tr)

transformed_te <- bake(center_obj, biomass_te)

biomass_te[1:10, names(transformed_te)]
transformed_te

tidy(center_trans, number = 1)
tidy(center_obj, number = 1)
```

---

step\_classdist

*Distances to Class Centroids*

---

**Description**

`step_classdist` creates a *specification* of a recipe step that will convert numeric data into Mahalanobis distance measurements to the data centroid. This is done for each value of a categorical class variable.

**Usage**

```

step_classdist(
  recipe,
  ...,
  class,
  role = "predictor",
  trained = FALSE,
  mean_func = mean,
  cov_func = cov,
  pool = FALSE,
  log = TRUE,
  objects = NULL,
  prefix = "classdist_",
  skip = FALSE,
  id = rand_id("classdist")
)

## S3 method for class 'step_classdist'
tidy(x, ...)

```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
class	A single character string that specifies a single categorical variable to be used as the class.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that resulting distances will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
mean_func	A function to compute the center of the distribution.
cov_func	A function that computes the covariance matrix
pool	A logical: should the covariance matrix be computed by pooling the data for all of the classes?
log	A logical: should the distances be transformed by the natural log function?
objects	Statistics are stored here once this step has been trained by <a href="#">prep.recipe()</a> .
prefix	A character string that defines the naming convention for new distance columns. Defaults to "classdist_". See Details below.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations

id	A character string that is unique to this step to identify it.
x	A step_classdist object.

### Details

step\_classdist will create a new column for every unique value of the class variable. The resulting variables will not replace the original values and by default have the prefix classdist\_. The naming format can be changed using the prefix argument.

Note that, by default, the default covariance function requires that each class should have at least as many rows as variables listed in the terms argument. If pool = TRUE, there must be at least as many data points are variables overall.

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms (the selectors or variables selected), value (the centroid of the class), and class.

### Examples

```
# in case of missing data...
mean2 <- function(x) mean(x, na.rm = TRUE)

# define naming convention
rec <- recipe(Species ~ ., data = iris) %>%
  step_classdist(all_predictors(), class = "Species",
                 pool = FALSE, mean_func = mean2, prefix = "centroid_")

# default naming
rec <- recipe(Species ~ ., data = iris) %>%
  step_classdist(all_predictors(), class = "Species",
                 pool = FALSE, mean_func = mean2)

rec_dists <- prep(rec, training = iris)

dists_to_species <- bake(rec_dists, new_data = iris, everything())
## on log scale:
dist_cols <- grep("classdist", names(dists_to_species), value = TRUE)
dists_to_species[, c("Species", dist_cols)]

tidy(rec, number = 1)
tidy(rec_dists, number = 1)
```

## Description

step\_corr creates a *specification* of a recipe step that will potentially remove variables that have large absolute correlations with other variables.

## Usage

```
step_corr(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  threshold = 0.9,
  use = "pairwise.complete.obs",
  method = "pearson",
  removals = NULL,
  skip = FALSE,
  id = rand_id("corr")
)

## S3 method for class 'step_corr'
tidy(x, ...)
```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
threshold	A value for the threshold of absolute correlation values. The step will try to remove the minimum number of columns so that all the resulting absolute correlations are less than this value.
use	A character string for the use argument to the <a href="#">stats::cor()</a> function.
method	A character string for the method argument to the <a href="#">stats::cor()</a> function.
removals	A character string that contains the names of columns that should be removed. These values are not determined until <a href="#">prep.recipe()</a> is called.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_corr object.

**Details**

This step attempts to remove variables to keep the largest absolute correlation between the variables less than threshold.

When a column has a single unique value, that column will be excluded from the correlation analysis. Also, if the data set has sporadic missing values (and an inappropriate value of use is chosen), some columns will also be excluded from the filter.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms which is the columns that will be removed.

**Author(s)**

Original R code for filtering algorithm by Dong Li, modified by Max Kuhn. Contributions by Reynald Lescarbeau (for original in caret package). Max Kuhn for the step function.

**See Also**

[step\\_nzv\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```
library(modeldata)
data(biomass)

set.seed(3535)
biomass$duplicate <- biomass$carbon + rnorm(nrow(biomass))

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen +
              sulfur + duplicate,
              data = biomass_tr)

corr_filter <- rec %>%
  step_corr(all_predictors(), threshold = .5)

filter_obj <- prep(corr_filter, training = biomass_tr)

filtered_te <- bake(filter_obj, biomass_te)
round(abs(cor(biomass_tr[, c(3:7, 9)])), 2)
round(abs(cor(filtered_te)), 2)

tidy(corr_filter, number = 1)
tidy(filter_obj, number = 1)
```

step\_count

*Create Counts of Patterns using Regular Expressions***Description**

step\_count creates a *specification* of a recipe step that will create a variable that counts instances of a regular expression pattern in text.

**Usage**

```
step_count(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  pattern = ".",
  normalize = FALSE,
  options = list(),
  result = make.names(pattern),
  input = NULL,
  skip = FALSE,
  id = rand_id("count")
)

## S3 method for class 'step_count'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	A single selector functions to choose which variable will be searched for the pattern. The selector should resolve into a single variable. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	For a variable created by this step, what analysis role should they be assigned?. By default, the function assumes that the new dummy variable column created by the original variable will be used as a predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
pattern	A character string containing a regular expression (or character string for fixed = TRUE) to be matched in the given character vector. Coerced by as.character to a character string if possible.
normalize	A logical; should the integer counts be divided by the total number of characters in the string?.
options	A list of options to <a href="#">gregexpr()</a> that should not include x or pattern.



result	A single character value for the name of the new variable. It should be a valid column name.
input	A single character value for the name of the variable being searched. This is NULL until computed by <code>prep.recipe()</code> .
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_count</code> object.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `result` (the new column name).

### Examples

```
library(modeldata)
data(covers)

rec <- recipe(~ description, covers) %>%
  step_count(description, pattern = "(rock|stony)", result = "rocks") %>%
  step_count(description, pattern = "famil", normalize = TRUE)

rec2 <- prep(rec, training = covers)
rec2

count_values <- bake(rec2, new_data = covers)
count_values

tidy(rec, number = 1)
tidy(rec2, number = 1)
```

---

step\_date

*Date Feature Generator*

---

### Description

`step_date` creates a *specification* of a recipe step that will convert date data into one or more factor or numeric variables.

**Usage**

```

step_date(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  features = c("dow", "month", "year"),
  abbr = TRUE,
  label = TRUE,
  ordinal = FALSE,
  columns = NULL,
  skip = FALSE,
  id = rand_id("date")
)

## S3 method for class 'step_date'
tidy(x, ...)

```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be used to create the new variables. The selected variables should have class Date or POSIXct. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new variable columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
features	A character string that includes at least one of the following values: month, dow (day of week), doy (day of year), week, month, decimal (decimal date, e.g. 2002.197), quarter, semester, year.
abbr	A logical. Only available for features month or dow. FALSE will display the day of the week as an ordered factor of character strings, such as "Sunday". TRUE will display an abbreviated version of the label, such as "Sun". abbr is disregarded if label = FALSE.
label	A logical. Only available for features month or dow. TRUE will display the day of the week as an ordered factor of character strings, such as "Sunday." FALSE will display the day of the week as a number.
ordinal	A logical: should factors be ordered? Only available for features month or dow.
columns	A character string of variables that will be used as inputs. This field is a placeholder and will be populated once <a href="#">prep.recipe()</a> is used.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations

may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using `skip = TRUE` as it may affect the computations for subsequent operations

`id` A character string that is unique to this step to identify it.  
`x` A `step_date` object.

### Details

Unlike other steps, `step_date` does *not* remove the original date variables. `step_rm()` can be used for this purpose.

### Value

For `step_date`, an updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns `terms` (the selectors or variables selected), `value` (the feature names), and `ordinal` (a logical).

### See Also

[step\\_holiday\(\)](#) [step\\_rm\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

### Examples

```
library(lubridate)

examples <- data.frame(Dan = ymd("2002-03-04") + days(1:10),
                      Stefan = ymd("2006-01-13") + days(1:10))
date_rec <- recipe(~ Dan + Stefan, examples) %>%
  step_date(all_predictors())

tidy(date_rec, number = 1)

date_rec <- prep(date_rec, training = examples)

date_values <- bake(date_rec, new_data = examples)
date_values

tidy(date_rec, number = 1)
```

---

step\_depth

*Data Depths*

---

### Description

`step_depth` creates a *specification* of a recipe step that will convert numeric data into measurement of *data depth*. This is done for each value of a categorical class variable.

**Usage**

```

step_depth(
  recipe,
  ...,
  class,
  role = "predictor",
  trained = FALSE,
  metric = "halfspace",
  options = list(),
  data = NULL,
  prefix = "depth_",
  skip = FALSE,
  id = rand_id("depth")
)

## S3 method for class 'step_depth'
tidy(x, ...)

```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be used to create the new features. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
class	A single character string that specifies a single categorical variable to be used as the class.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that resulting depth estimates will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
metric	A character string specifying the depth metric. Possible values are "potential", "halfspace", "Mahalanobis", "simplicialVolume", "spatial", and "zonoid".
options	A list of options to pass to the underlying depth functions. See <a href="#">ddalpha::depth.halfspace()</a> , <a href="#">ddalpha::depth.Mahalanobis()</a> , <a href="#">ddalpha::depth.potential()</a> , <a href="#">ddalpha::depth.projection()</a> , <a href="#">ddalpha::depth.simplicial()</a> , <a href="#">ddalpha::depth.simplicialVolume()</a> , <a href="#">ddalpha::depth.spatial</a> and <a href="#">ddalpha::depth.zonoid()</a> .
data	The training data are stored here once after <a href="#">prep.recipe()</a> is executed.
prefix	A character string that defines the naming convention for new depth columns. Defaults to "depth_". See Details below.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations

id	A character string that is unique to this step to identify it.
x	A step_depth object.

### Details

Data depth metrics attempt to measure how close data a data point is to the center of its distribution. There are a number of methods for calculating death but a simple example is the inverse of the distance of a data point to the centroid of the distribution. Generally, small values indicate that a data point not close to the centroid. `step_depth` can compute a class-specific depth for a new data point based on the proximity of the new value to the training set distribution.

This step requires the **ddalpha** package. If not installed, the step will stop with a note about installing the package.

Note that the entire training set is saved to compute future depth values. The saved data have been trained (i.e. prepared) and baked (i.e. processed) up to the point before the location that `step_depth` occupies in the recipe. Also, the data requirements for the different step methods may vary. For example, using `metric = "Mahalanobis"` requires that each class should have at least as many rows as variables listed in the `terms` argument.

The function will create a new column for every unique value of the `class` variable. The resulting variables will not replace the original values and by default have the prefix `depth_`. The naming format can be changed using the `prefix` argument.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `class`.

### Examples

```
# halfspace depth is the default
rec <- recipe(Species ~ ., data = iris) %>%
  step_depth(all_predictors(), class = "Species")

# use zonoid metric instead
# also, define naming convention for new columns
rec <- recipe(Species ~ ., data = iris) %>%
  step_depth(all_predictors(), class = "Species",
             metric = "zonoid", prefix = "zonoid_")

rec_dists <- prep(rec, training = iris)

dists_to_species <- bake(rec_dists, new_data = iris)
dists_to_species

tidy(rec, number = 1)
tidy(rec_dists, number = 1)
```

---

step_discretize	<i>Discretize Numeric Variables</i>
-----------------	-------------------------------------

---

## Description

step\_discretize creates a *specification* of a recipe step that will convert numeric data into a factor with bins having approximately the same number of data points (based on a training set).

## Usage

```
step_discretize(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  num_breaks = 4,
  min_unique = 10,
  objects = NULL,
  options = list(),
  skip = FALSE,
  id = rand_id("discretize")
)

## S3 method for class 'step_discretize'
tidy(x, ...)
```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	For step_discretize, the dots specify one or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
num_breaks	An integer defining how many cuts to make of the data.
min_unique	An integer defining a sample size line of dignity for the binning. If (the number of unique values)/(cuts+1) is less than min_unique, no discretization takes place.
objects	The <a href="#">discretize()</a> objects are stored here once the recipe has been trained by <a href="#">prep.recipe()</a> .
options	A list of options to <a href="#">discretize()</a> . A default is set for the argument x. Note that using the options prefix and labels when more than one variable is being transformed might be problematic as all variables inherit those values.

skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_discretize</code> object

**Value**

`step_discretize` returns an updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `value` (the breaks).

---

step_downsample	<i>Down-Sample a Data Set Based on a Factor Variable</i>
-----------------	--

---

**Description**

`step_downsample` creates a *specification* of a recipe step that will remove rows of a data set to make the occurrence of levels in a specific factor level equal.

**Usage**

```
step_downsample(
  recipe,
  ...,
  under_ratio = 1,
  ratio = NA,
  role = NA,
  trained = FALSE,
  column = NULL,
  target = NA,
  skip = TRUE,
  seed = sample.int(10^5, 1),
  id = rand_id("downsample")
)

## S3 method for class 'step_downsample'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
--------	--

...	One or more selector functions to choose which variable is used to sample the data. See <code>selections()</code> for more details. The selection should result in <i>single factor variable</i> . For the <code>tidy</code> method, these are not currently used.
<code>under_ratio</code>	A numeric value for the ratio of the minority-to-majority frequencies. The default value (1) means that all other levels are sampled down to have the same frequency as the least occurring level. A value of 2 would mean that the majority levels will have (at most) (approximately) twice as many rows than the minority level.
<code>ratio</code>	Deprecated argument; same as <code>under_ratio</code>
<code>role</code>	Not used by this step since no new variables are created.
<code>trained</code>	A logical to indicate if the quantities for preprocessing have been estimated.
<code>column</code>	A character string of the variable name that will be populated (eventually) by the ... selectors.
<code>target</code>	An integer that will be used to subsample. This should not be set by the user and will be populated by <code>prep</code> .
<code>skip</code>	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
<code>seed</code>	An integer that will be used as the seed when downsampling.
<code>id</code>	A character string that is unique to this step to identify it.
<code>x</code>	A <code>step_downsample</code> object.

### Details

Down-sampling is intended to be performed on the *training* set alone. For this reason, the default is `skip = TRUE`. It is advisable to use `prep(recipe, retain = TRUE)` when preparing the recipe; in this way `juice()` can be used to obtain the down-sampled version of the data.

If there are missing values in the factor variable that is used to define the sampling, missing data are selected at random in the same way that the other factor levels are sampled. Missing values are not used to determine the amount of data in the minority level

For any data with factor levels occurring with the same frequency as the minority level, all data will be retained.

All columns in the data are sampled and returned by `juice()` and `bake()`.

Keep in mind that the location of down-sampling in the step may have effects. For example, if centering and scaling, it is not clear whether those operations should be conducted *before* or *after* rows are removed.

When used in modeling, users should strongly consider using the option `skip = TRUE` so that the extra sampling is *not* conducted outside of the training set.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` which is the variable used to sample.



**Examples**

```

library(modeldata)
data(okc)

sort(table(okc$diet, useNA = "always"))

ds_rec <- recipe(~ ., data = okc) %>%
  step_downsample(diet) %>%
  prep(training = okc)

table(juice(ds_rec)$diet, useNA = "always")

# since `skip` defaults to TRUE, baking the step has no effect
baked_okc <- bake(ds_rec, new_data = okc)
table(baked_okc$diet, useNA = "always")

```

step\_dummy

*Dummy Variables Creation***Description**

step\_dummy creates a *specification* of a recipe step that will convert nominal data (e.g. character or factors) into one or more numeric binary model terms for the levels of the original data.

**Usage**

```

step_dummy(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  one_hot = FALSE,
  preserve = FALSE,
  naming = dummy_names,
  levels = NULL,
  skip = FALSE,
  id = rand_id("dummy")
)

## S3 method for class 'step_dummy'
tidy(x, ...)

```

**Arguments**

recipe            A recipe object. The step will be added to the sequence of operations for this recipe.

...	One or more selector functions to choose which <i>factor</i> variables will be used to create the dummy variables. See <a href="#">selections()</a> for more details. The selected variables must be factors. For the <code>tidy</code> method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the binary dummy variable columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
one_hot	A logical. For C levels, should C dummy variables be created rather than C-1?
preserve	A single logical; should the selected column(s) be retained (in addition to the new dummy variables).
naming	A function that defines the naming convention for new dummy columns. See Details below.
levels	A list that contains the information needed to create dummy variables for each variable contained in <code>terms</code> . This is NULL until the step is trained by <a href="#">prep.recipe()</a> .
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_dummy</code> object.

## Details

`step_dummy` will create a set of binary dummy variables from a factor variable. For example, if an unordered factor column in the data set has levels of "red", "green", "blue", the dummy variable `bake` will create two additional columns of 0/1 data for two of those three values (and remove the original column). For ordered factors, polynomial contrasts are used to encode the numeric values.

By default, the excluded dummy variable (i.e. the reference cell) will correspond to the first level of the unordered factor being converted.

The function allows for non-standard naming of the resulting variables. For an unordered factor named `x`, with levels "a" and "b", the default naming convention would be to create a new variable called `x_b`. Note that if the factor levels are not valid variable names (e.g. "some text with spaces"), it will be changed by `base::make.names()` to be valid (see the example below). The naming format can be changed using the `naming` argument and the function `dummy_names()` is the default. This function will also change the names of ordinal dummy variables. Instead of values such as ".L", ".Q", or "^4", ordinal dummy variables are given simple integer suffixes such as "\_1", "\_2", etc.

To change the type of contrast being used, change the global contrast option via `options`.

When the factor being converted has a missing value, all of the corresponding dummy variables are also missing.

When data to be processed contains novel levels (i.e., not contained in the training set), a missing value is assigned to the results. See [step\\_other\(\)](#) for an alternative.

If no columns are selected (perhaps due to an earlier `step_zv()`), the `bake()` and `juice()` functions will return the data as-is (e.g. with no dummy variables).

The [package vignette for dummy variables](#) and interactions has more information.

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms (the selectors or original variables selected) and columns (the list of corresponding binary columns).

### See Also

[step\\_factor2string\(\)](#), [step\\_string2factor\(\)](#), [dummy\\_names\(\)](#), [step\\_regex\(\)](#), [step\\_count\(\)](#), [step\\_ordinalscore\(\)](#), [step\\_unorder\(\)](#), [step\\_other\(\)](#) [step\\_novel\(\)](#)

### Examples

```
library(modeldata)
data(okc)
okc <- okc[complete.cases(okc),]

rec <- recipe(~ diet + age + height, data = okc)

dummies <- rec %>% step_dummy(diet)
dummies <- prep(dummies, training = okc)

dummy_data <- bake(dummies, new_data = okc)

unique(okc$diet)
grep("^diet", names(dummy_data), value = TRUE)

# Obtain the full set of dummy variables using `one_hot` option
rec %>%
  step_dummy(diet, one_hot = TRUE) %>%
  prep(training = okc) %>%
  juice(starts_with("diet")) %>%
  names() %>%
  length()

length(unique(okc$diet))

# Without one_hot
length(grep("^diet", names(dummy_data), value = TRUE))

tidy(dummies, number = 1)
```

---

step\_factor2string      *Convert Factors to Strings*

---

### Description

step\_factor2string will convert one or more factor vectors to strings.

**Usage**

```

step_factor2string(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  columns = FALSE,
  skip = FALSE,
  id = rand_id("factor2string")
)

## S3 method for class 'step_factor2string'
tidy(x, ...)

```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will converted to strings See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variables that will be converted. This is NULL until computed by <a href="#">prep.recipe()</a> .
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_factor2string object.

**Details**

prep has an option `strings_as_factors` that defaults to TRUE. If this step is used with the default option, the string(s) produced by this step will be converted to factors after all of the steps have been prepped.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms (the columns that will be affected).

**See Also**

[step\\_string2factor\(\)](#) [step\\_dummy\(\)](#)

**Examples**

```
library(modeldata)
data(okc)

rec <- recipe(~ diet + location, data = okc)

rec <- rec %>%
  step_string2factor(diet)

factor_test <- rec %>%
  prep(training = okc,
        strings_as_factors = FALSE) %>%
  juice
# diet is a
class(factor_test$diet)

rec <- rec %>%
  step_factor2string(diet)

string_test <- rec %>%
  prep(training = okc,
        strings_as_factors = FALSE) %>%
  juice
# diet is a
class(string_test$diet)

tidy(rec, number = 1)
```

---

step\_filter

*Filter rows using dplyr*

---

**Description**

step\_filter creates a *specification* of a recipe step that will remove rows using `dplyr::filter()`.

**Usage**

```
step_filter(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  inputs = NULL,
  skip = FALSE,
  id = rand_id("filter")
)

## S3 method for class 'step_filter'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	Logical predicates defined in terms of the variables in the data. Multiple conditions are combined with &. Only rows where the condition evaluates to TRUE are kept. See <code>dplyr::filter()</code> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
inputs	Quosure of values given by ...
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_filter</code> object

**Details**

When an object in the user's global environment is referenced in the expression defining the new variable(s), it is a good idea to use quasiquotation (e.g. `!!`) to embed the value of the object in the expression (to be portable between sessions). See the examples.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms which contains the conditional statements. These expressions are text representations and are not parsable.

**Examples**

```
rec <- recipe( ~ ., data = iris) %>%
  step_filter(Sepal.Length > 4.5, Species == "setosa")

prepped <- prep(rec, training = iris %>% slice(1:75))

library(dplyr)

dplyr_train <-
  iris %>%
  as_tibble() %>%
  slice(1:75) %>%
  dplyr::filter(Sepal.Length > 4.5, Species == "setosa")

rec_train <- juice(prepped)
all.equal(dplyr_train, rec_train)
```

```

dplyr_test <-
  iris %>%
  as_tibble() %>%
  slice(76:150) %>%
  dplyr::filter(Sepal.Length > 4.5, Species != "setosa")
rec_test <- bake(prepped, iris %>% slice(76:150))
all.equal(dplyr_test, rec_test)

values <- c("versicolor", "virginica")

qq_rec <-
  recipe(~ ., data = iris) %>%
  # Embed the `values` object in the call using !!
  step_filter(Sepal.Length > 4.5, Species %in% !!values)

tidy(qq_rec, number = 1)

```

---

step\_geodist

*Distance between two locations*


---

## Description

step\_geodist creates a a *specification* of a recipe step that will calculate the distance between points on a map to a reference location.

## Usage

```

step_geodist(
  recipe,
  lat = NULL,
  lon = NULL,
  role = "predictor",
  trained = FALSE,
  ref_lat = NULL,
  ref_lon = NULL,
  log = FALSE,
  name = "geo_dist",
  columns = NULL,
  skip = FALSE,
  id = rand_id("geodist")
)

## S3 method for class 'step_geodist'
tidy(x, ...)

```

## Arguments

recipe            A recipe object. The step will be added to the sequence of operations for this recipe.

lon, lat	Selector functions to choose which variables are affected by the step. See <code>selections()</code> for more details.
role	or model term created by this step, what analysis role should be assigned?. By default, the function assumes that resulting distance will be used as a predictor in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
ref_lon, ref_lat	Single numeric values for the location of the reference point.
log	A logical: should the distance be transformed by the natural log function?
name	A single character value to use for the new predictor column. If a column exists with this name, an error is issued.
columns	A character string of variable names that will be populated (eventually) by the <code>terms</code> argument.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_geodist</code> object.
...	One or more selector functions to choose which variables are affected by the step. See <code>selections()</code> for more details. For the <code>tidy</code> method, these are not currently used.

## Details

`step_geodist` will create a

## Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns echoing the values of `lat`, `lon`, `ref_lat`, `ref_lon`, `name`, and `id`.

## Examples

```
library(modeldata)
data(Smithsonian)

# How close are the museums to Union Station?
near_station <- recipe(~ ., data = Smithsonian) %>%
  update_role(name, new_role = "location") %>%
  step_geodist(lat = latitude, lon = longitude, log = FALSE,
              ref_lat = 38.8986312, ref_lon = -77.0062457) %>%
  prep(training = Smithsonian)
```



```
juice(near_station) %>%
  arrange(geo_dist)

tidy(near_station, number = 1)
```

---

```
step_holiday      Holiday Feature Generator
```

---

## Description

step\_holiday creates a *specification* of a recipe step that will convert date data into one or more binary indicator variables for common holidays.

## Usage

```
step_holiday(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  holidays = c("LaborDay", "NewYearsDay", "ChristmasDay"),
  columns = NULL,
  skip = FALSE,
  id = rand_id("holiday")
)

## S3 method for class 'step_holiday'
tidy(x, ...)
```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used to create the new variables. The selected variables should have class Date or POSIXct. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new variable columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
holidays	A character string that includes at least one holiday supported by the timeDate package. See <a href="#">timeDate::listHolidays()</a> for a complete list.
columns	A character string of variables that will be used as inputs. This field is a placeholder and will be populated once <a href="#">prep.recipe()</a> is used.

skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_holiday</code> object.

### Details

Unlike other steps, `step_holiday` does *not* remove the original date variables. `step_rm()` can be used for this purpose.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns `terms` which is the columns that will be affected and `holiday`.

### See Also

[step\\_date\(\)](#) [step\\_rm\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#) [timeDate::listHolidays\(\)](#)

### Examples

```
library(lubridate)

examples <- data.frame(someday = ymd("2000-12-20") + days(0:40))
holiday_rec <- recipe(~ someday, examples) %>%
  step_holiday(all_predictors())

holiday_rec <- prep(holiday_rec, training = examples)
holiday_values <- bake(holiday_rec, new_data = examples)
holiday_values
```

---

step\_hyperbolic

*Hyperbolic Transformations*

---

### Description

`step_hyperbolic` creates a *specification* of a recipe step that will transform data using a hyperbolic function.

**Usage**

```
step_hyperbolic(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  func = "sin",
  inverse = TRUE,
  columns = NULL,
  skip = FALSE,
  id = rand_id("hyperbolic")
)

## S3 method for class 'step_hyperbolic'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
func	A character value for the function. Valid values are "sin", "cos", or "tan".
inverse	A logical: should the inverse function be used?
columns	A character string of variable names that will be populated (eventually) by the terms argument.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_hyperbolic object.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms (the columns that will be affected), inverse, and func.

**See Also**

[step\\_logit\(\)](#) [step\\_invlogit\(\)](#) [step\\_log\(\)](#) [step\\_sqrt\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

## Examples

```
set.seed(313)
examples <- matrix(rnorm(40), ncol = 2)
examples <- as.data.frame(examples)

rec <- recipe(~ V1 + V2, data = examples)

cos_trans <- rec %>%
  step_hyperbolic(all_predictors(),
                 func = "cos", inverse = FALSE)

cos_obj <- prep(cos_trans, training = examples)

transformed_te <- bake(cos_obj, examples)
plot(examples$V1, transformed_te$V1)

tidy(cos_trans, number = 1)
tidy(cos_obj, number = 1)
```

---

step\_ica

*ICA Signal Extraction*

---

## Description

step\_ica creates a *specification* of a recipe step that will convert numeric data into one or more independent components.

## Usage

```
step_ica(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  num_comp = 5,
  options = list(),
  res = NULL,
  prefix = "IC",
  skip = FALSE,
  id = rand_id("ica")
)

## S3 method for class 'step_ica'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used to compute the components. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new independent component columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
num_comp	The number of ICA components to retain as new predictors. If <code>num_comp</code> is greater than the number of columns or the number of possible components, a smaller value will be used.
options	A list of options to <code>fastICA::fastICA()</code> . No defaults are set here. <b>Note</b> that the arguments <code>X</code> and <code>n.comp</code> should not be passed here.
res	The <code>fastICA::fastICA()</code> object is stored here once this preprocessing step has been trained by <code>prep.recipe()</code> .
prefix	A character string that will be the prefix to the resulting new variables. See notes below.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_ica</code> object.

**Details**

Independent component analysis (ICA) is a transformation of a group of variables that produces a new set of artificial features or components. ICA assumes that the variables are mixtures of a set of distinct, non-Gaussian signals and attempts to transform the data to isolate these signals. Like PCA, the components are statistically independent from one another. This means that they can be used to combat large inter-variables correlations in a data set. Also like PCA, it is advisable to center and scale the variables prior to running ICA.

This package produces components using the "FastICA" methodology (see reference below). This step requires the **dimRed** and **fastICA** packages. If not installed, the step will stop with a note about installing these packages.

The argument `num_comp` controls the number of components that will be retained (the original variables that are used to derive the components are removed from the data). The new components will have names that begin with `prefix` and a sequence of numbers. The variable names are padded with zeros. For example, if `num_comp < 10`, their names will be `IC1 - IC9`. If `num_comp = 101`, the names would be `IC001 - IC101`.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms (the selectors or variables selected), value (the loading), and component.

**References**

Hyvarinen, A., and Oja, E. (2000). Independent component analysis: algorithms and applications. *Neural Networks*, 13(4-5), 411-430.

**See Also**

[step\\_pca\(\)](#) [step\\_kpca\(\)](#) [step\\_isomap\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```
# from fastICA::fastICA
set.seed(131)
S <- matrix(runif(400), 200, 2)
A <- matrix(c(1, 1, -1, 3), 2, 2, byrow = TRUE)
X <- as.data.frame(S %%% A)

tr <- X[1:100, ]
te <- X[101:200, ]

rec <- recipe( ~ ., data = tr)

ica_trans <- step_center(rec, V1, V2)
ica_trans <- step_scale(ica_trans, V1, V2)
ica_trans <- step_ica(ica_trans, V1, V2, num_comp = 2)

if (require(dimRed) & require(fastICA)) {
  ica_estimates <- prep(ica_trans, training = tr)
  ica_data <- bake(ica_estimates, te)

  plot(te$V1, te$V2)
  plot(ica_data$IC1, ica_data$IC2)

  tidy(ica_trans, number = 3)
  tidy(ica_estimates, number = 3)
}
```

---

step\_integer

---

*Convert values to predefined integers*


---

**Description**

step\_integer creates a *specification* of a recipe step that will convert new data into a set of integers based on the original data values.

**Usage**

```
step_integer(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  strict = FALSE,
  zero_based = FALSE,
  key = NULL,
  skip = FALSE,
  id = rand_id("integer")
)

## S3 method for class 'step_integer'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used to create the integer variables. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
strict	A logical for whether the values should be returned as integers (as opposed to double).
zero_based	A logical for whether the integers should start at zero and new values be appended as the largest integer.
key	A list that contains the information needed to create integer variables for each variable contained in terms. This is NULL until the step is trained by <a href="#">prep.recipe()</a> .
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_integer object.

**Details**

step\_integer will determine the unique values of each variable from the training set (excluding missing values), order them, and then assign integers to each value. When baked, each data point

is translated to its corresponding integer or a value of zero for yet unseen data (although see the `zero_based` argument above). Missing values propagate.

Factor inputs are ordered by their levels. All others are ordered by sort.

Despite the name, the new values are returned as numeric unless `strict = TRUE`, which will coerce the results to integers.

## Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns `terms` (the selectors or variables selected) and `value` is a *list column* with the conversion key.

## See Also

[step\\_factor2string\(\)](#), [step\\_string2factor\(\)](#), [step\\_regex\(\)](#), [step\\_count\(\)](#), [step\\_ordinalscore\(\)](#), [step\\_unorder\(\)](#), [step\\_other\(\)](#), [step\\_novel\(\)](#), [step\\_dummy\(\)](#)

## Examples

```
library(modeldata)
data(okc)

okc$location <- factor(okc$location)

okc_tr <- okc[1:100, ]
okc_tr$age[1] <- NA

okc_te <- okc[101:105, ]
okc_te$age[1] <- NA
okc_te$diet[1] <- "fast food"
okc_te$diet[2] <- NA

rec <- recipe(Class ~ ., data = okc_tr) %>%
  step_integer(all_predictors()) %>%
  prep(training = okc_tr)

bake(rec, okc_te, all_predictors())
tidy(rec, number = 1)
```

---

step\_interact

*Create Interaction Variables*

---

## Description

`step_interact` creates a *specification* of a recipe step that will create new columns that are interaction terms between two or more variables.



**Usage**

```
step_interact(
  recipe,
  terms,
  role = "predictor",
  trained = FALSE,
  objects = NULL,
  sep = "_x_",
  skip = FALSE,
  id = rand_id("interact")
)

## S3 method for class 'step_interact'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
terms	A traditional R formula that contains interaction terms. This can include . and selectors.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new columns created from the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
objects	A list of terms objects for each individual interaction.
sep	A character value used to delineate variables in an interaction (e.g. var1_x_var2 instead of the more traditional var1:var2).
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_interact</code> object
...	One or more selector functions to choose which variables are affected by the step. See <code>selections()</code> for more details. For the <code>tidy</code> method, these are not currently used.

**Details**

`step_interact` can create interactions between variables. It is primarily intended for **numeric data**; categorical variables should probably be converted to dummy variables using `step_dummy()` prior to being used for interactions.

Unlike other step functions, the `terms` argument should be a traditional R model formula but should contain no inline functions (e.g. `log`). For example, for predictors A, B, and C, a formula such as

$\sim A:B:C$  can be used to make a three way interaction between the variables. If the formula contains terms other than interactions (e.g.  $(A+B+C)^3$ ) only the interaction terms are retained for the design matrix.

The separator between the variables defaults to `"_x_"` so that the three way interaction shown previously would generate a column named `A_x_B_x_C`. This can be changed using the `sep` argument.

When dummy variables are created and are used in interactions, selectors can help specify the interactions succinctly. For example, suppose a factor column `X` gets converted to dummy variables `x_2`, `x_3`, ..., `x_6` using `step_dummy()`. If you wanted an interaction with numeric column `z`, you could create a set of specific interaction effects (e.g. `x_2:z + x_3:z` and so on) or you could use `starts_with("z_"):z`. When `prep()` evaluates this step, `starts_with("z_")` resolves to `(x_2 + x_3 + x_4 + x_5 + x_6):z` so that the formula is now `(x_2 + x_3 + x_4 + x_5 + x_6):z` and all two-way interactions are created.

## Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns `terms` which is the interaction effects.

## Examples

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

int_mod_1 <- rec %>%
  step_interact(terms = ~ carbon:hydrogen)

int_mod_2 <- rec %>%
  step_interact(terms = ~ (matches("gen$") + sulfur)^2)

int_mod_1 <- prep(int_mod_1, training = biomass_tr)
int_mod_2 <- prep(int_mod_2, training = biomass_tr)

dat_1 <- bake(int_mod_1, biomass_te)
dat_2 <- bake(int_mod_2, biomass_te)

names(dat_1)
names(dat_2)

tidy(int_mod_1, number = 1)
tidy(int_mod_2, number = 1)
```

---

step_intercept	Add intercept (or constant) column
----------------	------------------------------------

---

### Description

`step_intercept` creates a *specification* of a recipe step that will add an intercept or constant term in the first column of a data matrix. `step_intercept` has defaults to *predictor* role so that it is by default called in the bake step. Be careful to avoid unintentional transformations when calling steps with `all_predictors`.

### Usage

```
step_intercept(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  name = "intercept",
  value = 1,
  skip = FALSE,
  id = rand_id("intercept")
)
```

### Arguments

<code>recipe</code>	A recipe object. The step will be added to the sequence of operations for this recipe.
<code>...</code>	Argument ignored; included for consistency with other step specification functions.
<code>role</code>	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new columns created from the original variables will be used as predictors in a model.
<code>trained</code>	A logical to indicate if the quantities for preprocessing have been estimated. Again included for consistency.
<code>name</code>	Character name for new added column
<code>value</code>	A numeric constant to fill the intercept column. Defaults to 1.
<code>skip</code>	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
<code>id</code>	A character string that is unique to this step to identify it.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any).

**See Also**

[recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)
rec_trans <- recipe(HHV ~ ., data = biomass_tr[, -(1:2)]) %>%
  step_intercept(value = 2) %>%
  step_scale(carbon)

rec_obj <- prep(rec_trans, training = biomass_tr)

with_intercept <- bake(rec_obj, biomass_te)
with_intercept
```

---

step\_inverse

*Inverse Transformation*

---

**Description**

step\_inverse creates a *specification* of a recipe step that will inverse transform the data.

**Usage**

```
step_inverse(
  recipe,
  ...,
  role = NA,
  offset = 0,
  trained = FALSE,
  columns = NULL,
  skip = FALSE,
  id = rand_id("inverse")
)

## S3 method for class 'step_inverse'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
offset	An optional value to add to the data prior to logging (to avoid 1/0).
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variable names that will be populated (eventually) by the terms argument.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_inverse object.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms which is the columns that will be affected.

**See Also**

[step\\_log\(\)](#) [step\\_sqrt\(\)](#) [step\\_hyperbolic\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```
set.seed(313)
examples <- matrix(runif(40), ncol = 2)
examples <- data.frame(examples)

rec <- recipe(~ X1 + X2, data = examples)

inverse_trans <- rec %>%
  step_inverse(all_predictors())

inverse_obj <- prep(inverse_trans, training = examples)

transformed_te <- bake(inverse_obj, examples)
plot(examples$X1, transformed_te$X1)

tidy(inverse_trans, number = 1)
tidy(inverse_obj, number = 1)
```

---

step_invlogit	<i>Inverse Logit Transformation</i>
---------------	-------------------------------------

---

### Description

step\_invlogit creates a *specification* of a recipe step that will transform the data from real values to be between zero and one.

### Usage

```
step_invlogit(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  columns = NULL,
  skip = FALSE,
  id = rand_id("invlogit")
)

## S3 method for class 'step_invlogit'
tidy(x, ...)
```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variable names that will be populated (eventually) by the terms argument.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_invlogit object.

### Details

The inverse logit transformation takes values on the real line and translates them to be between zero and one using the function  $f(x) = 1/(1+\exp(-x))$ .

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any).  
For the tidy method, a tibble with columns terms which is the columns that will be affected.

**See Also**

[step\\_logit\(\)](#) [step\\_log\(\)](#) [step\\_sqrt\(\)](#) [step\\_hyperbolic\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

ilogit_trans <- rec %>%
  step_center(carbon, hydrogen) %>%
  step_scale(carbon, hydrogen) %>%
  step_invlogit(carbon, hydrogen)

ilogit_obj <- prep(ilogit_trans, training = biomass_tr)

transformed_te <- bake(ilogit_obj, biomass_te)
plot(biomass_te$carbon, transformed_te$carbon)
```

---

step\_isomap

*Isomap Embedding*


---

**Description**

step\_isomap creates a *specification* of a recipe step that will convert numeric data into one or more new dimensions.

**Usage**

```
step_isomap(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  num_terms = 5,
  neighbors = 50,
  options = list(.mute = c("message", "output")),
  res = NULL,
```

```

  prefix = "Isomap",
  skip = FALSE,
  id = rand_id("isomap")
)

## S3 method for class 'step_isomap'
tidy(x, ...)

```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used to compute the dimensions. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new dimension columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
num_terms	The number of isomap dimensions to retain as new predictors. If num_terms is greater than the number of columns or the number of possible dimensions, a smaller value will be used.
neighbors	The number of neighbors.
options	A list of options to <a href="#">dimRed::Isomap()</a> .
res	The <a href="#">dimRed::Isomap()</a> object is stored here once this preprocessing step has been trained by <a href="#">prep.recipe()</a> .
prefix	A character string that will be the prefix to the resulting new variables. See notes below.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_isomap object

## Details

Isomap is a form of multidimensional scaling (MDS). MDS methods try to find a reduced set of dimensions such that the geometric distances between the original data points are preserved. This version of MDS uses nearest neighbors in the data as a method for increasing the fidelity of the new dimensions to the original data values.

This step requires the **dimRed**, **RSpectra**, **igraph**, and **RANN** packages. If not installed, the step will stop with a note about installing these packages.



It is advisable to center and scale the variables prior to running Isomap (`step_center` and `step_scale` can be used for this purpose).

The argument `num_terms` controls the number of components that will be retained (the original variables that are used to derive the components are removed from the data). The new components will have names that begin with `prefix` and a sequence of numbers. The variable names are padded with zeros. For example, if `num_terms < 10`, their names will be `Isomap1` - `Isomap9`. If `num_terms = 101`, the names would be `Isomap001` - `Isomap101`.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected).

### References

De Silva, V., and Tenenbaum, J. B. (2003). Global versus local methods in nonlinear dimensionality reduction. *Advances in Neural Information Processing Systems*. 721-728.

**dimRed**, a framework for dimensionality reduction, <https://github.com/gdkrmr>

### See Also

[step\\_pca\(\)](#) [step\\_kpca\(\)](#) [step\\_ica\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

### Examples

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

im_trans <- rec %>%
  step_YeoJohnson(all_predictors()) %>%
  step_normalize(all_predictors()) %>%
  step_isomap(all_predictors(), neighbors = 100, num_terms = 2)

if (require(dimRed) & require(RSpectra)) {
  im_estimates <- prep(im_trans, training = biomass_tr)

  im_te <- bake(im_estimates, biomass_te)

  rng <- extendrange(c(im_te$Isomap1, im_te$Isomap2))
  plot(im_te$Isomap1, im_te$Isomap2,
       xlim = rng, ylim = rng)

  tidy(im_trans, number = 3)
  tidy(im_estimates, number = 3)
```

```
}
```

---

```
step_knnimpute
```

```
Imputation via K-Nearest Neighbors
```

---

## Description

step\_knnimpute creates a *specification* of a recipe step that will impute missing data using nearest neighbors.

## Usage

```
step_knnimpute(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  neighbors = 5,
  impute_with = imp_vars(all_predictors()),
  options = list(nthread = 1, eps = 1e-08),
  ref_data = NULL,
  columns = NULL,
  skip = FALSE,
  id = rand_id("knnimpute")
)

## S3 method for class 'step_knnimpute'
tidy(x, ...)
```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose variables. For step_knnimpute, this indicates the variables to be imputed. When used with imp_vars, the dots indicates which variables are used to predict the missing data in each variable. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
neighbors	The number of neighbors.
impute_with	A call to imp_vars to specify which variables are used to impute the variables that can include specific variable names separated by commas or different selectors (see <a href="#">selections()</a> ). If a column is included in both lists to be imputed and to be an imputation predictor, it will be removed from the latter and not used to impute itself.

options	A named list of options to pass to <code>gower::gower_topn()</code> . Available options are currently <code>nthread</code> and <code>eps</code> .
ref_data	A tibble of data that will reflect the data preprocessing done up to the point of this imputation step. This is NULL until the step is trained by <code>prep.recipe()</code> .
columns	The column names that will be imputed and used for imputation. This is NULL until the step is trained by <code>prep.recipe()</code> .
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_knnimpute</code> object.

### Details

The step uses the training set to impute any other data sets. The only distance function available is Gower's distance which can be used for mixtures of nominal and numeric data.

Once the nearest neighbors are determined, the mode is used for predictor nominal variables and the mean is used for numeric data. Note that, if the underlying data are integer, the mean will be converted to an integer too.

Note that if a variable that is to be imputed is also in `impute_with`, this variable will be ignored.

It is possible that missing values will still occur after imputation if a large majority (or all) of the imputing variables are also missing.

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns `terms` (the selectors or variables for imputation), `predictors` (those variables used to impute), and `neighbors`.

### References

Gower, C. (1971) "A general coefficient of similarity and some of its properties," *Biometrics*, 857-871.

### Examples

```
library(recipes)
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training", ]
biomass_te <- biomass[biomass$dataset == "Testing", ]
biomass_te_whole <- biomass_te

# induce some missing data at random
set.seed(9039)
```

```

carb_missing <- sample(1:nrow(biomass_te), 3)
nitro_missing <- sample(1:nrow(biomass_te), 3)

biomass_te$carbon[carb_missing] <- NA
biomass_te$nitrogen[nitro_missing] <- NA

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

ratio_recipe <- rec %>%
  step_knnimpute(all_predictors(), neighbors = 3)
ratio_recipe2 <- prep(ratio_recipe, training = biomass_tr)
imputed <- bake(ratio_recipe2, biomass_te)

# how well did it work?
summary(biomass_te_whole$carbon)
cbind(before = biomass_te_whole$carbon[carb_missing],
      after = imputed$carbon[carb_missing])

summary(biomass_te_whole$nitrogen)
cbind(before = biomass_te_whole$nitrogen[nitro_missing],
      after = imputed$nitrogen[nitro_missing])

tidy(ratio_recipe, number = 1)
tidy(ratio_recipe2, number = 1)

```

---

step\_kpca

*Kernel PCA Signal Extraction*


---

## Description

step\_kpca a *specification* of a recipe step that will convert numeric data into one or more principal components using a kernel basis expansion.

## Usage

```

step_kpca(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  num_comp = 5,
  res = NULL,
  options = list(kernel = "rbfdot", kpar = list(sigma = 0.2)),
  prefix = "kPC",
  skip = FALSE,
  id = rand_id("kpca")
)

```

```
## S3 method for class 'step_kpca'
tidy(x, ...)
```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used to compute the components. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned? By default, the function assumes that the new principal component columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
num_comp	The number of PCA components to retain as new predictors. If <code>num_comp</code> is greater than the number of columns or the number of possible components, a smaller value will be used.
res	An S4 <code>kernlab::kpca()</code> object is stored here once this preprocessing step has been trained by <code>prep.recipe()</code> .
options	A list of options to <code>kernlab::kpca()</code> . Defaults are set for the arguments <code>kernel</code> and <code>kpar</code> but others can be passed in. <b>Note</b> that the arguments <code>x</code> and <code>features</code> should not be passed here (or at all).
prefix	A character string that will be the prefix to the resulting new variables. See notes below.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_kpca</code> object

### Details

Kernel principal component analysis (kPCA) is an extension a PCA analysis that conducts the calculations in a broader dimensionality defined by a kernel function. For example, if a quadratic kernel function were used, each variable would be represented by its original values as well as its square. This nonlinear mapping is used during the PCA analysis and can potentially help find better representations of the original data.

This step requires the **dimRed** and **kernlab** packages. If not installed, the step will stop with a note about installing these packages.

As with ordinary PCA, it is important to standardized the variables prior to running PCA (`step_center` and `step_scale` can be used for this purpose).

When performing kPCA, the kernel function (and any important kernel parameters) must be chosen. The **kernlab** package is used and the reference below discusses the types of kernels available and

their parameter(s). These specifications can be made in the `kernel` and `kpar` slots of the options argument to `step_kpca`.

The argument `num_comp` controls the number of components that will be retained (the original variables that are used to derive the components are removed from the data). The new components will have names that begin with `prefix` and a sequence of numbers. The variable names are padded with zeros. For example, if `num_comp < 10`, their names will be `kPC1 - kPC9`. If `num_comp = 101`, the names would be `kPC001 - kPC101`.

## Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected).

## References

Scholkopf, B., Smola, A., and Muller, K. (1997). Kernel principal component analysis. *Lecture Notes in Computer Science*, 1327, 583-588.

Karatzoglou, K., Smola, A., Hornik, K., and Zeileis, A. (2004). kernlab - An S4 package for kernel methods in R. *Journal of Statistical Software*, 11(1), 1-20.

## See Also

[step\\_pca\(\)](#) [step\\_ica\(\)](#) [step\\_isomap\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

## Examples

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

kpca_trans <- rec %>%
  step_YeoJohnson(all_predictors()) %>%
  step_normalize(all_predictors()) %>%
  step_kpca(all_predictors())

if (require(dimRed) & require(kernlab)) {
  kpca_estimates <- prep(kpca_trans, training = biomass_tr)

  kpca_te <- bake(kpca_estimates, biomass_te)

  rng <- extendrange(c(kpca_te$kPC1, kpca_te$kPC2))
  plot(kpca_te$kPC1, kpca_te$kPC2,
       xlim = rng, ylim = rng)

  tidy(kpca_trans, number = 3)
  tidy(kpca_estimates, number = 3)
```

```
}

```

---

step\_kpca\_poly

*Polynomial Kernel PCA Signal Extraction*


---

## Description

step\_kpca\_poly a *specification* of a recipe step that will convert numeric data into one or more principal components using a polynomial kernel basis expansion.

## Usage

```
step_kpca_poly(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  num_comp = 5,
  res = NULL,
  degree = 2,
  scale_factor = 1,
  offset = 1,
  prefix = "kPC",
  skip = FALSE,
  id = rand_id("kpca_poly")
)

## S3 method for class 'step_kpca_poly'
tidy(x, ...)
```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used to compute the components. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned? By default, the function assumes that the new principal component columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
num_comp	The number of PCA components to retain as new predictors. If num_comp is greater than the number of columns or the number of possible components, a smaller value will be used.
res	An S4 <a href="#">kernlab::kpca()</a> object is stored here once this preprocessing step has been trained by <a href="#">prep.recipe()</a> .

degree, scale_factor, offset	Numeric values for the polynomial kernel function.
prefix	A character string that will be the prefix to the resulting new variables. See notes below.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_kpca_poly</code> object

## Details

Kernel principal component analysis (kPCA) is an extension a PCA analysis that conducts the calculations in a broader dimensionality defined by a kernel function. For example, if a quadratic kernel function were used, each variable would be represented by its original values as well as its square. This nonlinear mapping is used during the PCA analysis and can potentially help find better representations of the original data.

This step requires the **dimRed** and **kernlab** packages. If not installed, the step will stop with a note about installing these packages.

As with ordinary PCA, it is important to standardized the variables prior to running PCA (`step_center` and `step_scale` can be used for this purpose).

The argument `num_comp` controls the number of components that will be retained (the original variables that are used to derive the components are removed from the data). The new components will have names that begin with `prefix` and a sequence of numbers. The variable names are padded with zeros. For example, if `num_comp < 10`, their names will be `kPC1 - kPC9`. If `num_comp = 101`, the names would be `kPC001 - kPC101`.

## Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected).

## References

Scholkopf, B., Smola, A., and Muller, K. (1997). Kernel principal component analysis. *Lecture Notes in Computer Science*, 1327, 583-588.

Karatzoglou, K., Smola, A., Hornik, K., and Zeileis, A. (2004). kernlab - An S4 package for kernel methods in R. *Journal of Statistical Software*, 11(1), 1-20.

## See Also

[step\\_pca\(\)](#) [step\\_ica\(\)](#) [step\\_isomap\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)



**Examples**

```

library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

kpca_trans <- rec %>%
  step_YeoJohnson(all_predictors()) %>%
  step_normalize(all_predictors()) %>%
  step_kpca_poly(all_predictors())

if (require(dimRed) & require(kernlab)) {
  kpca_estimates <- prep(kpca_trans, training = biomass_tr)

  kpca_te <- bake(kpca_estimates, biomass_te)

  rng <- extendrange(c(kpca_te$kPC1, kpca_te$kPC2))
  plot(kpca_te$kPC1, kpca_te$kPC2,
       xlim = rng, ylim = rng)

  tidy(kpca_trans, number = 3)
  tidy(kpca_estimates, number = 3)
}

```

---

step\_kpca\_rbf

*Radial Basis Function Kernel PCA Signal Extraction*


---

**Description**

step\_kpca\_rbf a *specification* of a recipe step that will convert numeric data into one or more principal components using a radial basis function kernel basis expansion.

**Usage**

```

step_kpca_rbf(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  num_comp = 5,
  res = NULL,
  sigma = 0.2,
  prefix = "kPC",
  skip = FALSE,
  id = rand_id("kpca_rbf")
)

```

```
)

## S3 method for class 'step_kpca_rbf'
tidy(x, ...)
```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used to compute the components. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned? By default, the function assumes that the new principal component columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
num_comp	The number of PCA components to retain as new predictors. If <code>num_comp</code> is greater than the number of columns or the number of possible components, a smaller value will be used.
res	An S4 <code>kernlab::kpca()</code> object is stored here once this preprocessing step has been trained by <code>prep.recipe()</code> .
sigma	A numeric value for the radial basis function parameter.
prefix	A character string that will be the prefix to the resulting new variables. See notes below.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_kpca_rbf</code> object

### Details

Kernel principal component analysis (kPCA) is an extension a PCA analysis that conducts the calculations in a broader dimensionality defined by a kernel function. For example, if a quadratic kernel function were used, each variable would be represented by its original values as well as its square. This nonlinear mapping is used during the PCA analysis and can potentially help find better representations of the original data.

This step requires the **dimRed** and **kernlab** packages. If not installed, the step will stop with a note about installing these packages.

As with ordinary PCA, it is important to standardized the variables prior to running PCA (`step_center` and `step_scale` can be used for this purpose).

The argument `num_comp` controls the number of components that will be retained (the original variables that are used to derive the components are removed from the data). The new components

will have names that begin with `prefix` and a sequence of numbers. The variable names are padded with zeros. For example, if `num_comp < 10`, their names will be `kPC1 - kPC9`. If `num_comp = 101`, the names would be `kPC001 - kPC101`.

## Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected).

## References

Scholkopf, B., Smola, A., and Muller, K. (1997). Kernel principal component analysis. *Lecture Notes in Computer Science*, 1327, 583-588.

Karatzoglou, K., Smola, A., Hornik, K., and Zeileis, A. (2004). kernlab - An S4 package for kernel methods in R. *Journal of Statistical Software*, 11(1), 1-20.

## See Also

[step\\_pca\(\)](#) [step\\_ica\(\)](#) [step\\_isomap\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

## Examples

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

kpca_trans <- rec %>%
  step_YeoJohnson(all_predictors()) %>%
  step_normalize(all_predictors()) %>%
  step_kpca_rbf(all_predictors())

if (require(dimRed) & require(kernlab)) {
  kpca_estimates <- prep(kpca_trans, training = biomass_tr)

  kpca_te <- bake(kpca_estimates, biomass_te)

  rng <- extendrange(c(kpca_te$kPC1, kpca_te$kPC2))
  plot(kpca_te$kPC1, kpca_te$kPC2,
       xlim = rng, ylim = rng)

  tidy(kpca_trans, number = 3)
  tidy(kpca_estimates, number = 3)
}
```

---

step_lag	<i>Create a lagged predictor</i>
----------	----------------------------------

---

### Description

step\_lag creates a *specification* of a recipe step that will add new columns of lagged data. Lagged data will by default include NA values where the lag was induced. These can be removed with [step\\_naomit\(\)](#), or you may specify an alternative filler value with the default argument.

### Usage

```
step_lag(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  lag = 1,
  prefix = "lag_",
  default = NA,
  columns = NULL,
  skip = FALSE,
  id = rand_id("lag")
)
```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details.
role	Defaults to "predictor"
trained	A logical to indicate if the quantities for preprocessing have been estimated.
lag	A vector of positive integers. Each specified column will be lagged for each value in the vector.
prefix	A prefix for generated column names, default to "lag_".
default	Passed to <code>dplyr::lag</code> , determines what fills empty rows left by lagging (defaults to NA).
columns	A character string of variable names that will be populated (eventually) by the terms argument.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.

**Details**

The step assumes that the data are already *in the proper sequential order* for lagging.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any).

**See Also**

[recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#) [step\\_naomit\(\)](#)

**Examples**

```
n <- 10
start <- as.Date('1999/01/01')
end <- as.Date('1999/01/10')

df <- data.frame(x = runif(n),
                 index = 1:n,
                 day = seq(start, end, by = "day"))

recipe(~ ., data = df) %>%
  step_lag(index, day, lag = 2:3) %>%
  prep(df) %>%
  bake(df)
```

---

step\_lincomb

*Linear Combination Filter*


---

**Description**

step\_lincomb creates a *specification* of a recipe step that will potentially remove numeric variables that have linear combinations between them.

**Usage**

```
step_lincomb(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  max_steps = 5,
  removals = NULL,
  skip = FALSE,
  id = rand_id("lincomp")
)

## S3 method for class 'step_lincomb'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
max_steps	A value.
removals	A character string that contains the names of columns that should be removed. These values are not determined until <a href="#">prep.recipe()</a> is called.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_lincomb</code> object.

**Details**

This step finds exact linear combinations between two or more variables and recommends which column(s) should be removed to resolve the issue. This algorithm may need to be applied multiple times (as defined by `max_steps`).

**Value**

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns `terms` which is the columns that will be removed.

**Author(s)**

Max Kuhn, Kirk Mettler, and Jed Wing

**See Also**

[step\\_nzv\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```
library(modeldata)
data(biomass)

biomass$new_1 <- with(biomass,
  .1*carbon - .2*hydrogen + .6*sulfur)
biomass$new_2 <- with(biomass,
  .5*carbon - .2*oxygen + .6*nitrogen)
```

```

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen +
              sulfur + new_1 + new_2,
              data = biomass_tr)

lincomb_filter <- rec %>%
  step_lincomb(all_predictors())

lincomb_filter_trained <- prep(lincomb_filter, training = biomass_tr)
lincomb_filter_trained

tidy(lincomb_filter, number = 1)
tidy(lincomb_filter_trained, number = 1)

```

---

step\_log

*Logarithmic Transformation*


---

### Description

step\_log creates a *specification* of a recipe step that will log transform data.

### Usage

```

step_log(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  base = exp(1),
  offset = 0,
  columns = NULL,
  skip = FALSE,
  signed = FALSE,
  id = rand_id("log")
)

## S3 method for class 'step_log'
tidy(x, ...)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.

role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
base	A numeric value for the base.
offset	An optional value to add to the data prior to logging (to avoid $\log(0)$ ).
columns	A character string of variable names that will be populated (eventually) by the terms argument.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
signed	A logical indicating whether to take the signed log. This is $\text{sign}(x) * \text{abs}(x)$ when $\text{abs}(x) \geq 1$ or 0 if $\text{abs}(x) < 1$ . If <code>TRUE</code> the <code>offset</code> argument will be ignored.
id	A character string that is unique to this step to identify it.
x	A <code>step_log</code> object.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the columns that will be affected) and `base`.

### See Also

[step\\_logit\(\)](#) [step\\_invlogit\(\)](#) [step\\_hyperbolic\(\)](#) [step\\_sqrt\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

### Examples

```
set.seed(313)
examples <- matrix(exp(rnorm(40)), ncol = 2)
examples <- as.data.frame(examples)

rec <- recipe(~ V1 + V2, data = examples)

log_trans <- rec %>%
  step_log(all_predictors())

log_obj <- prep(log_trans, training = examples)

transformed_te <- bake(log_obj, examples)
plot(examples$V1, transformed_te$V1)

tidy(log_trans, number = 1)
tidy(log_obj, number = 1)

# using the signed argument with negative values
examples2 <- matrix(rnorm(40, sd = 5), ncol = 2)
```



```

examples2 <- as.data.frame(examples2)

recipe(~ V1 + V2, data = examples2) %>%
  step_log(all_predictors()) %>%
  prep(training = examples2) %>%
  bake(examples2)

recipe(~ V1 + V2, data = examples2) %>%
  step_log(all_predictors(), signed = TRUE) %>%
  prep(training = examples2) %>%
  bake(examples2)

```

---

step\_logit

*Logit Transformation*


---

## Description

step\_logit creates a *specification* of a recipe step that will logit transform the data.

## Usage

```

step_logit(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  columns = NULL,
  skip = FALSE,
  id = rand_id("logit")
)

## S3 method for class 'step_logit'
tidy(x, ...)

```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variable names that will be populated (eventually) by the terms argument.

skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_logit</code> object.

### Details

The logit transformation takes values between zero and one and translates them to be on the real line using the function  $f(p) = \log(p/(1-p))$ .

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` which is the columns that will be affected.

### See Also

[step\\_invlogit\(\)](#) [step\\_log\(\)](#) [step\\_sqrt\(\)](#) [step\\_hyperbolic\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

### Examples

```
set.seed(313)
examples <- matrix(runif(40), ncol = 2)
examples <- data.frame(examples)

rec <- recipe(~ X1 + X2, data = examples)

logit_trans <- rec %>%
  step_logit(all_predictors())

logit_obj <- prep(logit_trans, training = examples)

transformed_te <- bake(logit_obj, examples)
plot(examples$X1, transformed_te$X1)

tidy(logit_trans, number = 1)
tidy(logit_obj, number = 1)
```

---

step\_lowerimpute

*Impute Numeric Data Below the Threshold of Measurement*

---

### Description

`step_lowerimpute` creates a *specification* of a recipe step designed for cases where the non-negative numeric data cannot be measured below a known value. In these cases, one method for imputing the data is to substitute the truncated value by a random uniform number between zero and the truncation point.

**Usage**

```
step_lowerimpute(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  threshold = NULL,
  skip = FALSE,
  id = rand_id("lowerimpute")
)

## S3 method for class 'step_lowerimpute'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
threshold	A named numeric vector of lower bounds This is NULL until computed by <a href="#">prep.recipe()</a> .
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_lowerimpute</code> object.

**Details**

`step_lowerimpute` estimates the variable minimums from the data used in the `training` argument of `prep.recipe`. `bake.recipe` then simulates a value for any data at the minimum with a random uniform value between zero and the minimum.

**Value**

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `value` for the estimated threshold.

## Examples

```
library(recipes)
library(modeldata)
data(biomass)

## Truncate some values to emulate what a lower limit of
## the measurement system might look like

biomass$carbon <- ifelse(biomass$carbon > 40, biomass$carbon, 40)
biomass$hydrogen <- ifelse(biomass$hydrogen > 5, biomass$carbon, 5)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

impute_rec <- rec %>%
  step_lowerimpute(carbon, hydrogen)

tidy(impute_rec, number = 1)

impute_rec <- prep(impute_rec, training = biomass_tr)

tidy(impute_rec, number = 1)

transformed_te <- bake(impute_rec, biomass_te)

plot(transformed_te$carbon, biomass_te$carbon,
      ylab = "pre-imputation", xlab = "imputed")
```

---

step\_meanimpute

*Impute Numeric Data Using the Mean*

---

## Description

step\_meanimpute creates a *specification* of a recipe step that will substitute missing values of numeric variables by the training set mean of those variables.

## Usage

```
step_meanimpute(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  means = NULL,
  trim = 0,
  skip = FALSE,
```

```

  id = rand_id("meanimpute")
)

## S3 method for class 'step_meanimpute'
tidy(x, ...)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
means	A named numeric vector of means. This is NULL until computed by <a href="#">prep.recipe()</a> . Note that, if the original data are integers, the mean will be converted to an integer to maintain the same a data type.
trim	The fraction (0 to 0.5) of observations to be trimmed from each end of the variables before the mean is computed. Values of trim outside that range are taken as the nearest endpoint.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_meanimpute</code> object.

### Details

`step_meanimpute` estimates the variable means from the data used in the training argument of `prep.recipe`. `bake.recipe` then applies the new values to new data sets using these averages.

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `model` (the mean value).

### Examples

```

library(modeldata)
data("credit_data")

## missing data per column
vapply(credit_data, function(x) mean(is.na(x)), c(num = 0))

```

```

set.seed(342)
in_training <- sample(1:nrow(credit_data), 2000)

credit_tr <- credit_data[ in_training, ]
credit_te <- credit_data[-in_training, ]
missing_examples <- c(14, 394, 565)

rec <- recipe(Price ~ ., data = credit_tr)

impute_rec <- rec %>%
  step_meanimpute(Income, Assets, Debt)

imp_models <- prep(impute_rec, training = credit_tr)

imputed_te <- bake(imp_models, new_data = credit_te, everything())

credit_te[missing_examples,]
imputed_te[missing_examples, names(credit_te)]

tidy(impute_rec, number = 1)
tidy(imp_models, number = 1)

```

---

step_medianimpute	<i>Impute Numeric Data Using the Median</i>
-------------------	---

---

## Description

step\_medianimpute creates a *specification* of a recipe step that will substitute missing values of numeric variables by the training set median of those variables.

## Usage

```

step_medianimpute(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  medians = NULL,
  skip = FALSE,
  id = rand_id("medianimpute")
)

## S3 method for class 'step_medianimpute'
tidy(x, ...)

```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <code>selections()</code> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
medians	A named numeric vector of medians. This is NULL until computed by <code>prep.recipe()</code> . Note that, if the original data are integers, the median will be converted to an integer to maintain the same a data type.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_medianimpute</code> object.

**Details**

`step_medianimpute` estimates the variable medians from the data used in the training argument of `prep.recipe`. `bake.recipe` then applies the new values to new data sets using these medians

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns `terms` (the selectors or variables selected) and `model` (the median value).

**Examples**

```
library(modeldata)
data("credit_data")

## missing data per column
vapply(credit_data, function(x) mean(is.na(x)), c(num = 0))

set.seed(342)
in_training <- sample(1:nrow(credit_data), 2000)

credit_tr <- credit_data[ in_training, ]
credit_te <- credit_data[-in_training, ]
missing_examples <- c(14, 394, 565)

rec <- recipe(Price ~ ., data = credit_tr)

impute_rec <- rec %>%
```

```

step_medianimpute(Income, Assets, Debt)

imp_models <- prep(impute_rec, training = credit_tr)

imputed_te <- bake(imp_models, new_data = credit_te, everything())

credit_te[missing_examples,]
imputed_te[missing_examples, names(credit_te)]

tidy(impute_rec, number = 1)
tidy(imp_models, number = 1)

```

---

step\_modeimpute

*Impute Nominal Data Using the Most Common Value*


---

### Description

step\_modeimpute creates a *specification* of a recipe step that will substitute missing values of nominal variables by the training set mode of those variables.

### Usage

```

step_modeimpute(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  modes = NULL,
  skip = FALSE,
  id = rand_id("modeimpute")
)

## S3 method for class 'step_modeimpute'
tidy(x, ...)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
modes	A named character vector of modes. This is NULL until computed by <a href="#">prep.recipe()</a> .



skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_modeimpute</code> object.

## Details

`step_modeimpute` estimates the variable modes from the data used in the training argument of `prep.recipe`. `bake.recipe` then applies the new values to new data sets using these values. If the training set data has more than one mode, one is selected at random.

## Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns `terms` (the selectors or variables selected) and `model` (the mode value).

## Examples

```
library(modeldata)
data("credit_data")

## missing data per column
vapply(credit_data, function(x) mean(is.na(x)), c(num = 0))

set.seed(342)
in_training <- sample(1:nrow(credit_data), 2000)

credit_tr <- credit_data[ in_training, ]
credit_te <- credit_data[-in_training, ]
missing_examples <- c(14, 394, 565)

rec <- recipe(Price ~ ., data = credit_tr)

impute_rec <- rec %>%
  step_modeimpute(Status, Home, Marital)

imp_models <- prep(impute_rec, training = credit_tr)

imputed_te <- bake(imp_models, new_data = credit_te, everything())

table(credit_te$Home, imputed_te$Home, useNA = "always")

tidy(impute_rec, number = 1)
tidy(imp_models, number = 1)
```

---

step_mutate	<i>Add new variables using mutate</i>
-------------	---------------------------------------

---

### Description

step\_mutate creates a *specification* of a recipe step that will add variables using `dplyr::mutate()`.

### Usage

```
step_mutate(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  inputs = NULL,
  skip = FALSE,
  id = rand_id("mutate")
)

## S3 method for class 'step_mutate'
tidy(x, ...)

## S3 method for class 'step_mutate_at'
tidy(x, ...)
```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	Name-value pairs of expressions. See <code>dplyr::mutate()</code> . If the argument is not named, the expression is converted to a column name.
role	For model terms created by this step, what analysis role should they be assigned? By default, the function assumes that the new dimension columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
inputs	Quosure(s) of ...
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_mutate object

## Details

When an object in the user's global environment is referenced in the expression defining the new variable(s), it is a good idea to use quasiquotation (e.g. `!!`) to embed the value of the object in the expression (to be portable between sessions). See the examples.

## Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with column values which contains the mutate expressions as character strings (and are not parsable).

## Examples

```
rec <-
  recipe( ~ ., data = iris) %>%
  step_mutate(
    dbl_width = Sepal.Width * 2,
    half_length = Sepal.Length / 2
  )

prepped <- prep(rec, training = iris %>% slice(1:75))

library(dplyr)

dplyr_train <-
  iris %>%
  as_tibble() %>%
  slice(1:75) %>%
  mutate(
    dbl_width = Sepal.Width * 2,
    half_length = Sepal.Length / 2
  )

rec_train <- juice(prepped)
all.equal(dplyr_train, rec_train)

dplyr_test <-
  iris %>%
  as_tibble() %>%
  slice(76:150) %>%
  mutate(
    dbl_width = Sepal.Width * 2,
    half_length = Sepal.Length / 2
  )

rec_test <- bake(prepped, iris %>% slice(76:150))
all.equal(dplyr_test, rec_test)

# Embedding objects:
const <- 1.414

qq_rec <-
  recipe( ~ ., data = iris) %>%
```

```

step_mutate(
  bad_approach = Sepal.Width * const,
  best_approach = Sepal.Width * !!const
) %>%
prep(training = iris)

juice(qq_rec, contains("appro")) %>% slice(1:4)

# The difference:
tidy(qq_rec, number = 1)

```

---

<code>step_mutate_at</code>	<i>Mutate multiple columns</i>
-----------------------------	--------------------------------

---

### Description

`step_mutate_at` creates a *specification* of a recipe step that will modify the selected variables using a common function.

### Usage

```

step_mutate_at(
  recipe,
  ...,
  fn,
  role = "predictor",
  trained = FALSE,
  inputs = NULL,
  skip = FALSE,
  id = rand_id("mutate_at")
)

```

### Arguments

<code>recipe</code>	A recipe object. The step will be added to the sequence of operations for this recipe.
<code>...</code>	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
<code>fn</code>	A function <code>fun</code> , a quosure style lambda <code>'~ fun(.)'</code> or a list of either form. (see <a href="#">dplyr::mutate_at()</a> ). <b>Note that this argument must be named.</b>
<code>role</code>	For model terms created by this step, what analysis role should they be assigned? By default, the function assumes that the new dimension columns created by the original variables will be used as predictors in a model.
<code>trained</code>	A logical to indicate if the quantities for preprocessing have been estimated.
<code>inputs</code>	A vector of column names populated by <code>prep()</code> .

skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.

**Value**

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` which contains the columns being transformed.

**Examples**

```
library(dplyr)
recipe(~ ., data = iris) %>%
  step_mutate_at(contains("Length"), fn = ~ 1/.) %>%
  prep() %>%
  juice() %>%
  slice(1:10)

recipe(~ ., data = iris) %>%
  # leads to more columns being created.
  step_mutate_at(contains("Length"), fn = list(log = log, sqrt = sqrt)) %>%
  prep() %>%
  juice() %>%
  slice(1:10)
```

---

step\_naomit

*Remove observations with missing values*


---

**Description**

`step_naomit` creates a *specification* of a recipe step that will add remove observations (rows of data) if they contain NA or NaN values.

**Usage**

```
step_naomit(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  columns = NULL,
  skip = FALSE,
  id = rand_id("naomit")
)

## S3 method for class 'step_naomit'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used to create the dummy variables. See <code>selections()</code> for more details. The selected variables must be factors.
role	Unused, include for consistency with other steps.
trained	A logical to indicate if the quantities for preprocessing have been estimated. Again included for consistency.
columns	A character string of variable names that will be populated (eventually) by the terms argument.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_naomit</code> object.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any).

**See Also**

`recipe()` `prep.recipe()` `bake.recipe()`

**Examples**

```
recipe(Ozone ~ ., data = airquality) %>%
  step_naomit(Solar.R) %>%
  prep(airquality, verbose = FALSE) %>%
  juice()
```

---

step\_nnmf

*NNMF Signal Extraction*


---

**Description**

`step_nnmf` creates a *specification* of a recipe step that will convert numeric data into one or more non-negative components.

**Usage**

```

step_nnmf(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  num_comp = 2,
  num_run = 30,
  options = list(),
  res = NULL,
  prefix = "NNMF",
  seed = sample.int(10^5, 1),
  skip = FALSE,
  id = rand_id("nnmf")
)

## S3 method for class 'step_nnmf'
tidy(x, ...)

```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used to compute the components. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new component columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
num_comp	The number of components to retain as new predictors. If <code>num_comp</code> is greater than the number of columns or the number of possible components, a smaller value will be used.
num_run	A positive integer for the number of computations runs used to obtain a consensus projection.
options	A list of options to <code>nmf()</code> in the NMF package by way of the <code>NNMF()</code> function in the <code>dimRed</code> package. <b>Note</b> that the arguments <code>data</code> and <code>ndim</code> should not be passed here.
res	The <code>NNMF()</code> object is stored here once this preprocessing step has been trained by <a href="#">prep.recipe()</a> .
prefix	A character string that will be the prefix to the resulting new variables. See notes below.
seed	An integer that will be used to set the seed in isolation when computing the factorization.

skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_nnmf</code> object.

## Details

Non-negative matrix factorization computes latent components that have non-negative values and take into account that the original data have non-negative values.

The argument `num_comp` controls the number of components that will be retained (the original variables that are used to derive the components are removed from the data). The new components will have names that begin with `prefix` and a sequence of numbers. The variable names are padded with zeros. For example, if `num < 10`, their names will be `NNMF1 - NNMF9`. If `num = 101`, the names would be `NNMF001 - NNMF101`.

## Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and the number of components.

## See Also

[step\\_pca\(\)](#), [step\\_ica\(\)](#), [step\\_kpca\(\)](#), [step\\_isomap\(\)](#), [recipe\(\)](#), [prep.recipe\(\)](#), [bake.recipe\(\)](#)

## Examples

```
library(modeldata)
data(biomass)

# rec <- recipe(HHV ~ ., data = biomass) %>%
#   update_role(sample, new_role = "id var") %>%
#   update_role(dataset, new_role = "split variable") %>%
#   step_nnmf(all_predictors(), num_comp = 2, seed = 473, num_run = 2) %>%
#   prep(training = biomass)
#
# juice(rec)
#
# library(ggplot2)
# ggplot(juice(rec), aes(x = NNMF2, y = NNMF1, col = HHV)) + geom_point()
```



---

step_normalize	<i>Center and scale numeric data</i>
----------------	--------------------------------------

---

### Description

step\_normalize creates a *specification* of a recipe step that will normalize numeric data to have a standard deviation of one and a mean of zero.

### Usage

```
step_normalize(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  means = NULL,
  sds = NULL,
  na_rm = TRUE,
  skip = FALSE,
  id = rand_id("normalize")
)

## S3 method for class 'step_normalize'
tidy(x, ...)
```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
means	A named numeric vector of means. This is NULL until computed by <a href="#">prep.recipe()</a> .
sds	A named numeric vector of standard deviations This is NULL until computed by <a href="#">prep.recipe()</a> .
na_rm	A logical value indicating whether NA values should be removed when computing the standard deviation and mean.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_normalize object.

**Details**

Centering data means that the average of a variable is subtracted from the data. Scaling data means that the standard deviation of a variable is divided out of the data. `step_normalize` estimates the variable standard deviations and means from the data used in the `training` argument of `prep.recipe`. `bake.recipe` then applies the scaling to new data sets using these estimates.

**Value**

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected), `value` (the standard deviations and means), and `statistic` for the type of value.

**Examples**

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

norm_trans <- rec %>%
  step_normalize(carbon, hydrogen)

norm_obj <- prep(norm_trans, training = biomass_tr)

transformed_te <- bake(norm_obj, biomass_te)

biomass_te[1:10, names(transformed_te)]
transformed_te
tidy(norm_trans, number = 1)
tidy(norm_obj, number = 1)
```

---

step\_novel

*Simple Value Assignments for Novel Factor Levels*


---

**Description**

`step_novel` creates a *specification* of a recipe step that will assign a previously unseen factor level to a new value.

**Usage**

```
step_novel(
  recipe,
  ...,
```

```

    role = NA,
    trained = FALSE,
    new_level = "new",
    objects = NULL,
    skip = FALSE,
    id = rand_id("novel")
  )

  ## S3 method for class 'step_novel'
  tidy(x, ...)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be affected by the step. These variables should be character or factor types. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
new_level	A single character value that will be assigned to new factor levels.
objects	A list of objects that contain the information on factor levels that will be determined by <a href="#">prep.recipe()</a> .
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_novel</code> object.

### Details

The selected variables are adjusted to have a new level (given by `new_level`) that is placed in the last position. During preparation there will be no data points associated with this new level since all of the data have been seen.

Note that if the original columns are character, they will be converted to factors by this step.

Missing values will remain missing.

If `new_level` is already in the data given to `prep`, an error is thrown.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the columns that will be affected) and `value` (the factor levels that is used for the new value)

**See Also**

[step\\_factor2string\(\)](#), [step\\_string2factor\(\)](#), [dummy\\_names\(\)](#), [step\\_regex\(\)](#), [step\\_count\(\)](#), [step\\_ordinalscore\(\)](#), [step\\_unorder\(\)](#), [step\\_other\(\)](#)

**Examples**

```
library(modeldata)
data(okc)

okc_tr <- okc[1:30000,]
okc_te <- okc[30001:30006,]
okc_te$diet[3] <- "cannibalism"
okc_te$diet[4] <- "vampirism"

rec <- recipe(~ diet + location, data = okc_tr)

rec <- rec %>%
  step_novel(diet, location)
rec <- prep(rec, training = okc_tr)

processed <- bake(rec, okc_te)
tibble(old = okc_te$diet, new = processed$diet)

tidy(rec, number = 1)
```

---

step\_ns

*Nature Spline Basis Functions*


---

**Description**

step\_ns creates a *specification* of a recipe step that will create new columns that are basis expansions of variables using natural splines.

**Usage**

```
step_ns(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  objects = NULL,
  deg_free = 2,
  options = list(),
  skip = FALSE,
  id = rand_id("ns")
)

## S3 method for class 'step_ns'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <code>selections()</code> for more details. For the tidy method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new columns created from the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
objects	A list of <code>splines::ns()</code> objects created once the step has been trained.
deg_free	The degrees of freedom.
options	A list of options for <code>splines::ns()</code> which should not include x or df.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_ns object.

**Details**

`step_ns` can new features from a single variable that enable fitting routines to model this variable in a nonlinear manner. The extent of the possible nonlinearity is determined by the `df` or `knot` arguments of `splines::ns()`. The original variables are removed from the data and new columns are added. The naming convention for the new variables is `varname_ns_1` and so on.

**Value**

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns `terms` which is the columns that will be affected and `holiday`.

**See Also**

[step\\_poly\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
```

```

      data = biomass_tr)

with_splines <- rec %>%
  step_ns(carbon, hydrogen)
with_splines <- prep(with_splines, training = biomass_tr)

expanded <- bake(with_splines, biomass_te)
expanded

```

---

step\_num2factor

*Convert Numbers to Factors*


---

### Description

step\_num2factor will convert one or more numeric vectors to factors (ordered or unordered). This can be useful when categories are encoded as integers.

### Usage

```

step_num2factor(
  recipe,
  ...,
  role = NA,
  transform = function(x) x,
  trained = FALSE,
  levels,
  ordered = FALSE,
  skip = FALSE,
  id = rand_id("num2factor")
)

## S3 method for class 'step_num2factor'
tidy(x, ...)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be converted to factors. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
transform	A function taking a single argument <code>x</code> that can be used to modify the numeric values prior to determining the levels (perhaps using <code>base::as.integer()</code> ). The output of a function should be an integer that corresponds to the value of levels that should be assigned. If not an integer, the value will be converted to an integer during <code>bake()</code> .

trained	A logical to indicate if the quantities for preprocessing have been estimated.
levels	A character vector of values that will be used as the levels. These are the numeric data converted to character and ordered. This is modified once <code>prep.recipe()</code> is executed.
ordered	A single logical value; should the factor(s) be ordered?
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_num2factor</code> object.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `ordered`.

**See Also**

[step\\_factor2string\(\)](#), [step\\_string2factor\(\)](#), [step\\_dummy\(\)](#)

**Examples**

```
library(dplyr)
library(modeldata)
data(attrition)

attrition %>%
  group_by(StockOptionLevel) %>%
  count()

amnt <- c("nothin", "meh", "some", "copious")

rec <-
  recipe(Attrition ~ StockOptionLevel, data = attrition) %>%
  step_num2factor(
    StockOptionLevel,
    transform = function(x) x + 1,
    levels = amnt
  )

encoded <- rec %>% prep() %>% juice()

table(encoded$StockOptionLevel, attrition$StockOptionLevel)

# an example for binning

binner <- function(x) {
```

```

x <- cut(x, breaks = 1000 * c(0, 5, 10, 20), include.lowest = TRUE)
# now return the group number
as.numeric(x)
}

inc <- c("low", "med", "high")

rec <-
  recipe(Attrition ~ MonthlyIncome, data = attrition) %>%
  step_num2factor(
    MonthlyIncome,
    transform = binner,
    levels = inc,
    ordered = TRUE
  ) %>%
  prep()

encoded <- juice(rec)

table(encoded$MonthlyIncome, binner(attrition$MonthlyIncome))

# What happens when a value is out of range?
ceo <- attrition %>% slice(1) %>% mutate(MonthlyIncome = 10^10)

bake(rec, ceo)

```

---

step\_nzv

*Near-Zero Variance Filter*


---

## Description

step\_nzv creates a *specification* of a recipe step that will potentially remove variables that are highly sparse and unbalanced.

## Usage

```

step_nzv(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  freq_cut = 95/5,
  unique_cut = 10,
  options = list(freq_cut = 95/5, unique_cut = 10),
  removals = NULL,
  skip = FALSE,
  id = rand_id("nzv")
)

```



```
## S3 method for class 'step_nzv'
tidy(x, ...)
```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will evaluated by the filtering. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
freq_cut, unique_cut	Numeric parameters for the filtering process. See the Details section below.
options	A list of options for the filter (see Details below).
removals	A character string that contains the names of columns that should be removed. These values are not determined until <a href="#">prep.recipe()</a> is called.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_nzv object.

### Details

This step diagnoses predictors that have one unique value (i.e. are zero variance predictors) or predictors that are have both of the following characteristics:

1. they have very few unique values relative to the number of samples and
2. the ratio of the frequency of the most common value to the frequency of the second most common value is large.

For example, an example of near zero variance predictor is one that, for 1000 samples, has two distinct values and 999 of them are a single value.

To be flagged, first the frequency of the most prevalent value over the second most frequent value (called the "frequency ratio") must be above `freq_cut`. Secondly, the "percent of unique values," the number of unique values divided by the total number of samples (times 100), must also be below `unique_cut`.

In the above example, the frequency ratio is 999 and the unique value percentage is 0.0001.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` which is the columns that will be removed.

**See Also**

[step\\_corr\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```
library(modeldata)
data(biomass)

biomass$sparse <- c(1, rep(0, nrow(biomass) - 1))

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen +
              nitrogen + sulfur + sparse,
              data = biomass_tr)

nzv_filter <- rec %>%
  step_nzv(all_predictors())

filter_obj <- prep(nzv_filter, training = biomass_tr)

filtered_te <- bake(filter_obj, biomass_te)
any(names(filtered_te) == "sparse")

tidy(nzv_filter, number = 1)
tidy(filter_obj, number = 1)
```

---

step\_ordinalscore

*Convert Ordinal Factors to Numeric Scores*

---

**Description**

step\_ordinalscore creates a *specification* of a recipe step that will convert ordinal factor variables into numeric scores.

**Usage**

```
step_ordinalscore(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  columns = NULL,
  convert = as.numeric,
  skip = FALSE,
  id = rand_id("ordinalscore")
)
```

```
## S3 method for class 'step_ordinalscore'
tidy(x, ...)
```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variables that will be converted. This is NULL until computed by <a href="#">prep.recipe()</a> .
convert	A function that takes an ordinal factor vector as an input and outputs a single numeric variable.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_ordinalscore</code> object.

### Details

Dummy variables from ordered factors with C levels will create polynomial basis functions with C-1 terms. As an alternative, this step can be used to translate the ordered levels into a single numeric vector of values that represent (subjective) scores. By default, the translation uses a linear scale (1, 2, 3, ... C) but custom score functions can also be used (see the example below).

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms (the columns that will be affected).

### Examples

```
fail_lvls <- c("meh", "annoying", "really_bad")

ord_data <-
  data.frame(item = c("paperclip", "twitter", "airbag"),
            fail_severity = factor(fail_lvls,
                                  levels = fail_lvls,
                                  ordered = TRUE))

model.matrix(~fail_severity, data = ord_data)
```

```

linear_values <- recipe(~ item + fail_severity, data = ord_data) %>%
  step_dummy(item) %>%
  step_ordinalscore(fail_severity)

linear_values <- prep(linear_values, training = ord_data)

juice(linear_values, everything())

custom <- function(x) {
  new_values <- c(1, 3, 7)
  new_values[as.numeric(x)]
}

nonlin_scores <- recipe(~ item + fail_severity, data = ord_data) %>%
  step_dummy(item) %>%
  step_ordinalscore(fail_severity, convert = custom)

tidy(nonlin_scores, number = 2)

nonlin_scores <- prep(nonlin_scores, training = ord_data)

juice(nonlin_scores, everything())

tidy(nonlin_scores, number = 2)

```

---

step\_other

*Collapse Some Categorical Levels*


---

### Description

step\_other creates a *specification* of a recipe step that will potentially pool infrequently occurring values into an "other" category.

### Usage

```

step_other(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  threshold = 0.05,
  other = "other",
  objects = NULL,
  skip = FALSE,
  id = rand_id("other")
)

## S3 method for class 'step_other'
tidy(x, ...)

```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will potentially be reduced. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
threshold	A numeric value between 0 and 1 or an integer greater or equal to one. If it's less than one then factor levels whose rate of occurrence in the training set are below threshold will be "othered". If it's greater or equal to one then it's treated as a frequency and factor levels that occur less than threshold times will be "othered".
other	A single character value for the "other" category.
objects	A list of objects that contain the information to pool infrequent levels that is determined by <a href="#">prep.recipe()</a> .
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_other</code> object.

**Details**

The overall proportion (or total counts) of the categories are computed. The "other" category is used in place of any categorical levels whose individual proportion (or frequency) in the training set is less than threshold.

If no pooling is done the data are unmodified (although character data may be changed to factors based on the value of `strings_as_factors` in [prep.recipe\(\)](#)). Otherwise, a factor is always returned with different factor levels.

If `threshold` is less than the largest category proportion, all levels except for the most frequent are collapsed to the other level.

If the retained categories include the value of `other`, an error is thrown. If `other` is in the list of discarded levels, no error occurs.

If no pooling is done, novel factor levels are converted to missing. If pooling is needed, they will be placed into the other category.

When data to be processed contains novel levels (i.e., not contained in the training set), the other category is assigned.

**Value**

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the columns that will be affected) and `retained` (the factor levels that were not pulled into "other")

**See Also**

[step\\_factor2string\(\)](#), [step\\_string2factor\(\)](#), [dummy\\_names\(\)](#), [step\\_regex\(\)](#), [step\\_count\(\)](#), [step\\_ordinalscore\(\)](#), [step\\_unorder\(\)](#), [step\\_novel\(\)](#)

**Examples**

```
library(modeldata)
data(okc)

set.seed(19)
in_train <- sample(1:nrow(okc), size = 30000)

okc_tr <- okc[ in_train, ]
okc_te <- okc[-in_train, ]

rec <- recipe(~ diet + location, data = okc_tr)

rec <- rec %>%
  step_other(diet, location, threshold = .1, other = "other values")
rec <- prep(rec, training = okc_tr)

collapsed <- bake(rec, okc_te)
table(okc_te$diet, collapsed$diet, useNA = "always")

tidy(rec, number = 1)

# novel levels are also "othered"
tahiti <- okc[1, ]
tahiti$location <- "a magical place"
bake(rec, tahiti)

# threshold as a frequency
rec <- recipe(~ diet + location, data = okc_tr)

rec <- rec %>%
  step_other(diet, location, threshold = 2000, other = "other values")
rec <- prep(rec, training = okc_tr)

tidy(rec, number = 1)
# compare it to
# okc_tr %>% count(diet, sort = TRUE) %>% top_n(4)
# okc_tr %>% count(location, sort = TRUE) %>% top_n(3)
```

---

step\_pca

*PCA Signal Extraction*


---

**Description**

`step_pca` creates a *specification* of a recipe step that will convert numeric data into one or more principal components.

**Usage**

```

step_pca(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  num_comp = 5,
  threshold = NA,
  options = list(),
  res = NULL,
  prefix = "PC",
  skip = FALSE,
  id = rand_id("pca")
)

## S3 method for class 'step_pca'
tidy(x, ...)

```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used to compute the components. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new principal component columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
num_comp	The number of PCA components to retain as new predictors. If <code>num_comp</code> is greater than the number of columns or the number of possible components, a smaller value will be used.
threshold	A fraction of the total variance that should be covered by the components. For example, <code>threshold = .75</code> means that <code>step_pca</code> should generate enough components to capture 75%. Note: using this argument will override and reset any value given to <code>num_comp</code> .
options	A list of options to the default method for <a href="#">stats::prcomp()</a> . Argument defaults are set to <code>retx = FALSE</code> , <code>center = FALSE</code> , <code>scale. = FALSE</code> , and <code>tol = NULL</code> . <b>Note</b> that the argument <code>x</code> should not be passed here (or at all).
res	The <a href="#">stats::prcomp.default()</a> object is stored here once this preprocessing step has been trained by <a href="#">prep.recipe()</a> .
prefix	A character string that will be the prefix to the resulting new variables. See notes below
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations

may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using `skip = TRUE` as it may affect the computations for subsequent operations

`id` A character string that is unique to this step to identify it.  
`x` A `step_pca` object.

### Details

Principal component analysis (PCA) is a transformation of a group of variables that produces a new set of artificial features or components. These components are designed to capture the maximum amount of information (i.e. variance) in the original variables. Also, the components are statistically independent from one another. This means that they can be used to combat large inter-variables correlations in a data set.

It is advisable to standardized the variables prior to running PCA. Here, each variable will be centered and scaled prior to the PCA calculation. This can be changed using the `options` argument or by using `step_center()` and `step_scale()`.

The argument `num_comp` controls the number of components that will be retained (the original variables that are used to derive the components are removed from the data). The new components will have names that begin with `prefix` and a sequence of numbers. The variable names are padded with zeros. For example, if `num_comp < 10`, their names will be PC1 - PC9. If `num_comp = 101`, the names would be PC001 - PC101.

Alternatively, `threshold` can be used to determine the number of components that are required to capture a specified fraction of the total variance in the variables.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected), `value` (the loading), and `component`.

### References

Jolliffe, I. T. (2010). *Principal Component Analysis*. Springer.

### See Also

[step\\_ica\(\)](#) [step\\_kpca\(\)](#) [step\\_isomap\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

### Examples

```
rec <- recipe(~ ., data = USArrests)
pca_trans <- rec %>%
  step_center(all_numeric()) %>%
  step_scale(all_numeric()) %>%
  step_pca(all_numeric(), num_comp = 3)
pca_estimates <- prep(pca_trans, training = USArrests)
pca_data <- bake(pca_estimates, USArrests)

rng <- extendrange(c(pca_data$PC1, pca_data$PC2))
```



```

plot(pca_data$PC1, pca_data$PC2,
     xlim = rng, ylim = rng)

with_thresh <- rec %>%
  step_center(all_numeric()) %>%
  step_scale(all_numeric()) %>%
  step_pca(all_numeric(), threshold = .99)
with_thresh <- prep(with_thresh, training = USArrests)
bake(with_thresh, USArrests)

tidy(pca_trans, number = 3)
tidy(pca_estimates, number = 3)

```

---

step\_pls

*Partial Least Squares Feature Extraction*


---

### Description

step\_pls creates a *specification* of a recipe step that will convert numeric data into one or more new dimensions.

### Usage

```

step_pls(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  num_comp = 2,
  outcome = NULL,
  options = NULL,
  res = NULL,
  prefix = "PLS",
  skip = FALSE,
  id = rand_id("pls")
)

## S3 method for class 'step_pls'
tidy(x, ...)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used to compute the dimensions. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.

role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new dimension columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
num_comp	The number of pls dimensions to retain as new predictors. If num_comp is greater than the number of columns or the number of possible dimensions, a smaller value will be used.
outcome	When a single outcome is available, character string or call to <code>dplyr::vars()</code> can be used to specify the variable. When there are multiple outcomes, <code>dplyr::vars()</code> must be used. This that can include specific variable names separated by commas or different selectors (see <code>selections()</code> ).
options	A list of options to <code>pls::plsr()</code> .
res	The <code>pls::plsr()</code> object is stored here once this preprocessing step has been trained by <code>prep.recipe()</code> .
prefix	A character string that will be the prefix to the resulting new variables. See notes below.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_pls</code> object

### Details

PLS is a supervised version of principal component analysis that requires one or more numeric outcomes to compute the new features. The data should be scaled (and perhaps centered) prior to running these calculations.

This step requires the **pls** package. If not installed, the step will stop with a note about installing the package.

The argument `num_comp` controls the number of components that will be retained (the original variables that are used to derive the components are removed from the data). The new components will have names that begin with `prefix` and a sequence of numbers. The variable names are padded with zeros. For example, if `num_comp < 10`, their names will be `PLS1 - PLS9`. If `num_comp = 101`, the names would be `PLS001 - PLS101`.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected).

### See Also

[step\\_pca\(\)](#) [step\\_kpca\(\)](#) [step\\_ica\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```

library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

pls_rec <- recipe(HHV ~ ., data = biomass_tr) %>%
  step_rm(sample, dataset) %>%
  step_normalize(all_predictors()) %>%
  # If the outcome(s) need standardization, do it in separate
  # steps with skip = FALSE so that new data where the
  # outcome is missing can be processed.
  step_normalize(all_outcomes(), skip = TRUE) %>%
  step_pls(all_predictors(), outcome = "HHV")

pls_rec <- prep(pls_rec, training = biomass_tr)

pls_test_scores <- bake(pls_rec, new_data = biomass_te[, -8])

tidy(pls_rec, number = 4)

```

---

step\_poly

*Orthogonal Polynomial Basis Functions*


---

**Description**

step\_poly creates a *specification* of a recipe step that will create new columns that are basis expansions of variables using orthogonal polynomials.

**Usage**

```

step_poly(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  objects = NULL,
  degree = 2,
  options = list(),
  skip = FALSE,
  id = rand_id("poly")
)

## S3 method for class 'step_poly'
tidy(x, ...)

```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the new columns created from the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
objects	A list of <a href="#">stats::poly()</a> objects created once the step has been trained.
degree	The polynomial degree (an integer).
options	A list of options for <a href="#">stats::poly()</a> which should not include x, degree, or simple. Note that the option raw = TRUE will produce the regular polynomial values (not orthogonalized).
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_poly object.

**Details**

step\_poly can new features from a single variable that enable fitting routines to model this variable in a nonlinear manner. The extent of the possible nonlinearity is determined by the degree argument of [stats::poly\(\)](#). The original variables are removed from the data and new columns are added. The naming convention for the new variables is varname\_poly\_1 and so on.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms (the columns that will be affected) and degree.

**See Also**

[step\\_ns\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]
```

```
rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

quadratic <- rec %>%
  step_poly(carbon, hydrogen)
quadratic <- prep(quadratic, training = biomass_tr)

expanded <- bake(quadratic, biomass_te)
expanded

tidy(quadratic, number = 1)
```

---

step\_profile

Create a Profiling Version of a Data Set

---

## Description

`step_profile` creates a *specification* of a recipe step that will fix the levels of all variables but one and will create a sequence of values for the remaining variable. This step can be helpful when creating partial regression plots for additive models.

## Usage

```
step_profile(
  recipe,
  ...,
  profile = NULL,
  pct = 0.5,
  index = 1,
  grid = list(pctl = TRUE, len = 100),
  columns = NULL,
  role = NA,
  trained = FALSE,
  skip = FALSE,
  id = rand_id("profile")
)

## S3 method for class 'step_profile'
tidy(x, ...)
```

## Arguments

<code>recipe</code>	A recipe object. The step will be added to the sequence of operations for this recipe.
<code>...</code>	One or more selector functions to choose which variables will fixed to a single value. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.

profile	A call to <code>dplyr::vars()</code> to specify which variable will be profiled (see <code>selections()</code> ). If a column is included in both lists to be fixed and to be profiled, an error is thrown.
pct	A value between 0 and 1 that is the percentile to fix continuous variables. This is applied to all continuous variables captured by the selectors. For date variables, either the minimum, median, or maximum used based on their distance to pct.
index	The level that qualitative variables will be fixed. If the variables are character (not factors), this will be the index of the sorted unique values. This is applied to all qualitative variables captured by the selectors.
grid	A named list with elements <code>pct1</code> (a logical) and <code>len</code> (an integer). If <code>pct1 = TRUE</code> , then <code>len</code> denotes how many percentiles to use to create the profiling grid. This creates a grid between 0 and 1 and the profile is determined by the percentiles of the data. For example, if <code>pct1 = TRUE</code> and <code>len = 3</code> , the profile would contain the minimum, median, and maximum values. If <code>pct1 = FALSE</code> , it defines how many grid points between the minimum and maximum values should be created. This parameter is ignored for qualitative variables (since all of their possible levels are profiled). In the case of date variables, <code>pct1 = FALSE</code> will always be used since there is no quantile method for dates.
columns	A character string that contains the names of columns that should be fixed and their values. These values are not determined until <code>prep.recipe()</code> is called.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_profile</code> object.

### Details

This step is atypical in that, when baked, the `new_data` argument is ignored; the resulting data set is based on the fixed and profiled variable's information.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (which is the columns that will be affected), and `type` (fixed or profiled).

### Examples

```
library(modeldata)
data(okc)

# Setup a grid across date but keep the other values fixed
```

```

recipe(~ diet + height + date, data = okc) %>%
  step_profile(-date, profile = vars(date)) %>%
  prep(training = okc) %>%
  juice

#####

# An *additive* model; not for use when there are interactions or
# other functional relationships between predictors

lin_mod <- lm(mpg ~ poly(dis, 2) + cyl + hp, data = mtcars)

# Show the difference in the two grid creation methods

disp_pctl <- recipe(~ disp + cyl + hp, data = mtcars) %>%
  step_profile(-disp, profile = vars(dis)) %>%
  prep(training = mtcars)

disp_grid <- recipe(~ disp + cyl + hp, data = mtcars) %>%
  step_profile(
    -disp,
    profile = vars(dis),
    grid = list(pctl = FALSE, len = 100)
  ) %>%
  prep(training = mtcars)

grid_data <- juice(disp_grid)
grid_data <- grid_data %>%
  mutate(pred = predict(lin_mod, grid_data),
         method = "grid")

pctl_data <- juice(disp_pctl)
pctl_data <- pctl_data %>%
  mutate(pred = predict(lin_mod, pctl_data),
         method = "percentile")

plot_data <- bind_rows(grid_data, pctl_data)

library(ggplot2)

ggplot(plot_data, aes(x = disp, y = pred)) +
  geom_point(alpha = .5, cex = 1) +
  facet_wrap(~ method)

```

**Description**

step\_range creates a *specification* of a recipe step that will normalize numeric data to be within a pre-defined range of values.

**Usage**

```
step_range(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  min = 0,
  max = 1,
  ranges = NULL,
  skip = FALSE,
  id = rand_id("range")
)

## S3 method for class 'step_range'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be scaled. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
min	A single numeric value for the smallest value in the range.
max	A single numeric value for the largest value in the range.
ranges	A character vector of variables that will be normalized. Note that this is ignored until the values are determined by <a href="#">prep.recipe()</a> . Setting this value will be ineffective.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_range object.

**Details**

When a new data point is outside of the ranges seen in the training set, the new values are truncated at min or max.



**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms (the selectors or variables selected), min, and max.

**Examples**

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

ranged_trans <- rec %>%
  step_range(carbon, hydrogen)

ranged_obj <- prep(ranged_trans, training = biomass_tr)

transformed_te <- bake(ranged_obj, biomass_te)

biomass_te[1:10, names(transformed_te)]
transformed_te

tidy(ranged_trans, number = 1)
tidy(ranged_obj, number = 1)
```

---

step\_ratio

*Ratio Variable Creation*


---

**Description**

step\_ratio creates a a *specification* of a recipe step that will create one or more ratios out of numeric variables.

**Usage**

```
step_ratio(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  denom = denom_vars(),
  naming = function(numer, denom) make.names(paste(numer, denom, sep = "_o_")),
  columns = NULL,
  skip = FALSE,
```

```

  id = rand_id("ratio")
)

denom_vars(...)

## S3 method for class 'step_ratio'
tidy(x, ...)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used in the <i>numerator</i> of the ratio. When used with <code>denom_vars</code> , the dots indicates which variables are used in the <i>denominator</i> . See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	For terms created by this step, what analysis role should they be assigned?. By default, the function assumes that the newly created ratios created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
denom	A call to <code>denom_vars</code> to specify which variables are used in the denominator that can include specific variable names separated by commas or different selectors (see <a href="#">selections()</a> ). If a column is included in both lists to be numerator and denominator, it will be removed from the listing.
naming	A function that defines the naming convention for new ratio columns.
columns	The column names used in the ratios. This argument is not populated until <a href="#">prep.recipe()</a> is executed.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_ratio</code> object

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `denom`.

### Examples

```

library(recipes)
library(modeldata)
data(biomass)

biomass$total <- apply(biomass[, 3:7], 1, sum)

```

```

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen +
              sulfur + total,
              data = biomass_tr)

ratio_recipe <- rec %>%
  # all predictors over total
  step_ratio(all_predictors(), denom = denom_vars(total)) %>%
  # get rid of the original predictors
  step_rm(all_predictors(), -ends_with("total"))

ratio_recipe <- prep(ratio_recipe, training = biomass_tr)

ratio_data <- bake(ratio_recipe, biomass_te)
ratio_data

```

---

step\_regex

*Create Dummy Variables using Regular Expressions*


---

## Description

step\_regex creates a *specification* of a recipe step that will create a new dummy variable based on a regular expression.

## Usage

```

step_regex(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  pattern = ".",
  options = list(),
  result = make.names(pattern),
  input = NULL,
  skip = FALSE,
  id = rand_id("regex")
)

## S3 method for class 'step_regex'
tidy(x, ...)

```

## Arguments

recipe            A recipe object. The step will be added to the sequence of operations for this recipe.

...	A single selector functions to choose which variable will be searched for the pattern. The selector should resolve into a single variable. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	For a variable created by this step, what analysis role should they be assigned?. By default, the function assumes that the new dummy variable column created by the original variable will be used as a predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
pattern	A character string containing a regular expression (or character string for <code>fixed = TRUE</code> ) to be matched in the given character vector. Coerced by <code>as.character</code> to a character string if possible.
options	A list of options to <a href="#">grepl()</a> that should not include <code>x</code> or <code>pattern</code> .
result	A single character value for the name of the new variable. It should be a valid column name.
input	A single character value for the name of the variable being searched. This is <code>NULL</code> until computed by <a href="#">prep.recipe()</a> .
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_regex</code> object.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `result` (the new column name).

### Examples

```
library(modeldata)
data(covers)

rec <- recipe(~ description, covers) %>%
  step_regex(description, pattern = "(rock|stony)", result = "rocks") %>%
  step_regex(description, pattern = "ratake families")

rec2 <- prep(rec, training = covers)
rec2

with_dummies <- bake(rec2, new_data = covers)
with_dummies
tidy(rec, number = 1)
tidy(rec2, number = 1)
```

---

step_relevel	<i>Relevel factors to a desired level</i>
--------------	---

---

### Description

step\_relevel creates a *specification* of a recipe step that will reorder the provided factor columns so that the level specified by ref\_level is first. This is useful for contr.treatment contrasts which take the first level as the reference.

### Usage

```
step_relevel(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  ref_level,
  objects = NULL,
  skip = FALSE,
  id = rand_id("relevel")
)

## S3 method for class 'step_relevel'
tidy(x, ...)
```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be affected by the step. These variables should be character or factor types. See <a href="#">selections()</a> for more details.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
ref_level	A single character value that will be used to relevel the factor column(s) (if the level is present).
objects	A list of objects that contain the information on factor levels that will be determined by <a href="#">prep.recipe()</a> .
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_relevel object.

**Details**

The selected variables are releveled to a level (given by `ref_level`). Placing the `ref_level` in the first position.

Note that if the original columns are character, they will be converted to factors by this step.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any).

**Examples**

```
library(modeldata)
data(okc)
rec <- recipe(~ diet + location, data = okc) %>%
  step_unknown(diet, new_level = "UNKNOWN") %>%
  step_relevel(diet, ref_level = "UNKNOWN") %>%
  prep()

data <- bake(rec, okc)
levels(data$diet)
```

---

step\_relu

*Apply (Smoothed) Rectified Linear Transformation*


---

**Description**

`step_relu` creates a *specification* of a recipe step that will apply the rectified linear or softplus transformations to numeric data. The transformed data is added as new columns to the data matrix.

**Usage**

```
step_relu(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  shift = 0,
  reverse = FALSE,
  smooth = FALSE,
  prefix = "right_relu_",
  columns = NULL,
  skip = FALSE,
  id = rand_id("relu")
)

## S3 method for class 'step_relu'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details.
role	Defaults to "predictor".
trained	A logical to indicate if the quantities for preprocessing have been estimated.
shift	A numeric value dictating a translation to apply to the data.
reverse	A logical to indicate if the left hinge should be used as opposed to the right hinge.
smooth	A logical indicating if the softplus function, a smooth approximation to the rectified linear transformation, should be used.
prefix	A prefix for generated column names, default to "right_relu_" when right hinge transformation and "left_relu_" for reversed/left hinge transformations.
columns	A character string of variable names that will be populated (eventually) by the terms argument.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_relu object.

**Details**

The rectified linear transformation is calculated as

$$\max(0, x - c)$$

and is also known as the ReLu or right hinge function. If `reverse` is true, then the transformation is reflected about the y-axis, like so:

$$\max(0, c - x)$$

Setting the `smooth` option to true will instead calculate a smooth approximation to ReLu according to

$$\ln(1 + e^{(x - c)})$$

The `reverse` argument may also be applied to this transformation.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any).

## Connection to MARS

The rectified linear transformation is used in Multivariate Adaptive Regression Splines as a basis function to fit piecewise linear functions to data in a strategy similar to that employed in tree based models. The transformation is a popular choice as an activation function in many neural networks, which could then be seen as a stacked generalization of MARS when making use of ReLU activations. The hinge function also appears in the loss function of Support Vector Machines, where it penalizes residuals only if they are within a certain margin of the decision boundary.

## See Also

[recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

## Examples

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

transformed_te <- rec %>%
  step_relu(carbon, shift = 40) %>%
  prep(biomass_tr) %>%
  bake(biomass_te)

transformed_te
```

---

step\_rename

*Rename variables by name*

---

## Description

step\_rename creates a *specification* of a recipe step that will add variables using `dplyr::rename()`.

## Usage

```
step_rename(
  recipe,
  ...,
  role = "predictor",
  trained = FALSE,
  inputs = NULL,
  skip = FALSE,
  id = rand_id("rename")
```



```
)

## S3 method for class 'step_rename'
tidy(x, ...)

## S3 method for class 'step_rename_at'
tidy(x, ...)
```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more unquoted expressions separated by commas. See <code>dplyr::rename()</code> where the convention is <code>new_name = old_name</code> .
role	For model terms created by this step, what analysis role should they be assigned? By default, the function assumes that the new dimension columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
inputs	Quosure(s) of ...
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_rename</code> object

### Details

When an object in the user's global environment is referenced in the expression defining the new variable(s), it is a good idea to use quasiquotation (e.g. `!!`) to embed the value of the object in the expression (to be portable between sessions).

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns values which contains the rename expressions as character strings (and are not reparable).

### Examples

```
recipe(~ ., data = iris) %>%
  step_rename(Sepal_Width = Sepal.Width) %>%
  prep() %>%
  juice() %>%
  slice(1:5)

vars <- c(var1 = "cyl", var2 = "am")
```

```

car_rec <-
  recipe(~ ., data = mtcars) %>%
  step_rename(!!vars)

car_rec %>%
  prep() %>%
  juice()

car_rec %>%
  tidy(number = 1)

```

---

step_rename_at	<i>Rename multiple columns</i>
----------------	--------------------------------

---

### Description

step\_rename\_at creates a *specification* of a recipe step that will rename the selected variables using a common function.

### Usage

```

step_rename_at(
  recipe,
  ...,
  fn,
  role = "predictor",
  trained = FALSE,
  inputs = NULL,
  skip = FALSE,
  id = rand_id("rename_at")
)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
fn	A function fun, a quosure style lambda ‘~ fun(.)’ or a list of either form (but containing only a single function, see <a href="#">dplyr::rename_at()</a> ). <b>Note that this argument must be named.</b>
role	For model terms created by this step, what analysis role should they be assigned? By default, the function assumes that the new dimension columns created by the original variables will be used as predictors in a model.
trained	A logical to indicate if the quantities for preprocessing have been estimated.

inputs	A vector of column names populated by <code>prep()</code> .
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` which contains the columns being transformed.

### Examples

```
library(dplyr)
recipe(~ ., data = iris) %>%
  step_rename_at(everything(), fn = ~ gsub(".", "_", ., fixed = TRUE)) %>%
  prep() %>%
  juice() %>%
  slice(1:10)
```

---

step_rm	<i>General Variable Filter</i>
---------	--------------------------------

---

### Description

`step_rm` creates a *specification* of a recipe step that will remove variables based on their name, type, or role.

### Usage

```
step_rm(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  removals = NULL,
  skip = FALSE,
  id = rand_id("rm")
)

## S3 method for class 'step_rm'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be evaluated by the filtering bake. See <code>selections()</code> for more details. For the <code>tidy</code> method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
removals	A character string that contains the names of columns that should be removed. These values are not determined until <code>prep.recipe()</code> is called.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_rm</code> object.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` which is the columns that will be removed.

**Examples**

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

library(dplyr)
smaller_set <- rec %>%
  step_rm(contains("gen"))

smaller_set <- prep(smaller_set, training = biomass_tr)

filtered_te <- bake(smaller_set, biomass_te)
filtered_te

tidy(smaller_set, number = 1)
```

---

step_rollimpute	<i>Impute Numeric Data Using a Rolling Window Statistic</i>
-----------------	---

---

## Description

step\_rollimpute creates a *specification* of a recipe step that will substitute missing values of numeric variables by the a measure of location (e.g. median) within a moving window.

## Usage

```
step_rollimpute(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  columns = NULL,
  statistic = median,
  window = 5,
  skip = FALSE,
  id = rand_id("rollimpute")
)

## S3 method for class 'step_rollimpute'
tidy(x, ...)
```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. These columns should be non-integer numerics (i.e., double precision). For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A named numeric vector of columns. This is NULL until computed by <a href="#">prep.recipe()</a> .
statistic	A function with a single argument for the data to compute the imputed value. Only complete values will be passed to the function and it should return a double precision value.
window	The size of the window around a point to be imputed. Should be an odd integer greater than one. See Details below for a discussion of points at the ends of the series.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome

	variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
<code>id</code>	A character string that is unique to this step to identify it.
<code>x</code>	A <code>step_rollimpute</code> object.

### Details

On the tails, the window is shifted towards the ends. For example, for a 5-point window, the windows for the first four points are 1:5, 1:5, 1:5, and then 2:6.

When missing data are in the window, they are not passed to the function. If all of the data in the window are missing, a missing value is returned.

The statistics are calculated on the training set values *before* imputation. This means that if previous data within the window are missing, their imputed values are not included in the window data used for imputation. In other words, each imputation does not know anything about previous imputations in the series prior to the current point.

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `window` (the window size).

### Examples

```
library(lubridate)

set.seed(145)
example_data <-
  data.frame(
    day = ymd("2012-06-07") + days(1:12),
    x1 = round(runif(12), 2),
    x2 = round(runif(12), 2),
    x3 = round(runif(12), 2)
  )
example_data$x1[c(1, 5, 6)] <- NA
example_data$x2[c(1:4, 10)] <- NA

library(recipes)
seven_pt <- recipe(~ . , data = example_data) %>%
  update_role(day, new_role = "time_index") %>%
  step_rollimpute(all_predictors(), window = 7) %>%
  prep(training = example_data)

juice(seven_pt)
```

---

step_sample	<i>Sample rows using dplyr</i>
-------------	--------------------------------

---

## Description

step\_sample creates a *specification* of a recipe step that will sample rows using `dplyr::sample_n()` or `dplyr::sample_frac()`.

## Usage

```
step_sample(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  size = NULL,
  replace = FALSE,
  skip = FALSE,
  id = rand_id("sample")
)

## S3 method for class 'step_sample'
tidy(x, ...)
```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	Argument ignored; included for consistency with other step specification functions. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
size	An integer or fraction. If the value is within (0, 1), <code>dplyr::sample_frac()</code> is applied to the data. If an integer value of 1 or greater is used, <code>dplyr::sample_n()</code> is applied. The default of NULL uses <code>dplyr::sample_n()</code> with the size of the training set (or smaller for smaller new_data).
replace	Sample with or without replacement?
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_sample object

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns size, replace, and id.

**Examples**

```
# Uses `sample_n`
recipe( ~ ., data = iris) %>%
  step_sample(size = 1) %>%
  prep(training = iris) %>%
  juice() %>%
  nrow()

# Uses `sample_frac`
recipe( ~ ., data = iris) %>%
  step_sample(size = 0.9999) %>%
  prep(training = iris) %>%
  juice() %>%
  nrow()

# Uses `sample_n` and returns _at maximum_ 120 samples.
smaller_iris <-
  recipe( ~ ., data = iris) %>%
  step_sample() %>%
  prep(training = iris %>% slice(1:120))

juice(smaller_iris) %>% nrow()
bake(smaller_iris, iris %>% slice(121:150)) %>% nrow()
```

---

step\_scale

*Scaling Numeric Data*


---

**Description**

step\_scale creates a *specification* of a recipe step that will normalize numeric data to have a standard deviation of one.

**Usage**

```
step_scale(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  sds = NULL,
  factor = 1,
  na_rm = TRUE,
  skip = FALSE,
```



```

  id = rand_id("scale")
)

## S3 method for class 'step_scale'
tidy(x, ...)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <code>selections()</code> for more details. For the <code>tidy</code> method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
sds	A named numeric vector of standard deviations. This is NULL until computed by <code>prep.recipe()</code> .
factor	A numeric value of either 1 or 2 that scales the numeric inputs by one or two standard deviations. By dividing by two standard deviations, the coefficients attached to continuous predictors can be interpreted the same way as with binary inputs. Defaults to 1. More in reference below.
na_rm	A logical value indicating whether NA values should be removed when computing the standard deviation.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_scale</code> object.

### Details

Scaling data means that the standard deviation of a variable is divided out of the data. `step_scale` estimates the variable standard deviations from the data used in the `training` argument of `prep.recipe`. `bake.recipe` then applies the scaling to new data sets using these standard deviations.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `value` (the standard deviations).

### References

Gelman, A. (2007) "Scaling regression inputs by dividing by two standard deviations." Unpublished. Source: <http://www.stat.columbia.edu/~gelman/research/unpublished/standardizing.pdf>.

## Examples

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

scaled_trans <- rec %>%
  step_scale(carbon, hydrogen)

scaled_obj <- prep(scaled_trans, training = biomass_tr)

transformed_te <- bake(scaled_obj, biomass_te)

biomass_te[1:10, names(transformed_te)]
transformed_te
tidy(scaled_trans, number = 1)
tidy(scaled_obj, number = 1)
```

---

step\_shuffle

*Shuffle Variables*

---

## Description

step\_shuffle creates a *specification* of a recipe step that will randomly change the order of rows for selected variables.

## Usage

```
step_shuffle(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  columns = NULL,
  skip = FALSE,
  id = rand_id("shuffle")
)

## S3 method for class 'step_shuffle'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be permuted. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string that contains the names of columns that should be shuffled. These values are not determined until <a href="#">prep.recipe()</a> is called.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_shuffle</code> object.

**Value**

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns `terms` which is the columns that will be affected.

**Examples**

```
integers <- data.frame(A = 1:12, B = 13:24, C = 25:36)

library(dplyr)
rec <- recipe(~ A + B + C, data = integers) %>%
  step_shuffle(A, B)

rand_set <- prep(rec, training = integers)

set.seed(5377)
bake(rand_set, integers)

tidy(rec, number = 1)
tidy(rand_set, number = 1)
```

---

step\_slice

*Filter rows by position using dplyr*


---

**Description**

`step_slice` creates a *specification* of a recipe step that will filter rows using [dplyr::slice\(\)](#).

**Usage**

```
step_slice(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  inputs = NULL,
  skip = FALSE,
  id = rand_id("slice")
)

## S3 method for class 'step_slice'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	Integer row values. See <code>dplyr::slice()</code> for more details. For the <code>tidy</code> method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
inputs	Quosure of values given by ...
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_slice</code> object

**Details**

When an object in the user's global environment is referenced in the expression defining the new variable(s), it is a good idea to use quasiquotation (e.g. `!!`) to embed the value of the object in the expression (to be portable between sessions). See the examples.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` which contains the filter indices.

**Examples**

```
rec <- recipe( ~ ., data = iris) %>%
  step_slice(1:3)
```

```

prepped <- prep(rec, training = iris %>% slice(1:75))
tidy(prepped, number = 1)

library(dplyr)

dplyr_train <-
  iris %>%
  as_tibble() %>%
  slice(1:75) %>%
  slice(1:3)

rec_train <- juice(prepped)
all.equal(dplyr_train, rec_train)

dplyr_test <-
  iris %>%
  as_tibble() %>%
  slice(76:150) %>%
  slice(1:3)
rec_test <- bake(prepped, iris %>% slice(76:150))
all.equal(dplyr_test, rec_test)

# Embedding the integer expression (or vector) into the
# recipe:

keep_rows <- 1:6

qq_rec <-
  recipe( ~ ., data = iris) %>%
  # Embed `keep_rows` in the call using !!
  step_slice(!!keep_rows) %>%
  prep(training = iris)

tidy(qq_rec, number = 1)

```

---

step\_spatialsign      *Spatial Sign Preprocessing*

---

### Description

step\_spatialsign is a *specification* of a recipe step that will convert numeric data into a projection on to a unit sphere.

### Usage

```

step_spatialsign(
  recipe,
  ...,
  role = "predictor",
  na_rm = TRUE,

```

```

    trained = FALSE,
    columns = NULL,
    skip = FALSE,
    id = rand_id("spatialsign")
  )

  ## S3 method for class 'step_spatialsign'
  tidy(x, ...)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be used for the normalization. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	For model terms created by this step, what analysis role should they be assigned?
na_rm	A logical: should missing data be removed from the norm computation?
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variable names that will be populated (eventually) by the terms argument.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_spatialsign object.

### Details

The spatial sign transformation projects the variables onto a unit sphere and is related to global contrast normalization. The spatial sign of a vector  $w$  is  $w/\text{norm}(w)$ .

The variables should be centered and scaled prior to the computations.

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms which is the columns that will be affected.

### References

Serneels, S., De Nolf, E., and Van Espen, P. (2006). Spatial sign preprocessing: a simple way to impart moderate robustness to multivariate estimators. *Journal of Chemical Information and Modeling*, 46(3), 1402-1409.

**Examples**

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

ss_trans <- rec %>%
  step_center(carbon, hydrogen) %>%
  step_scale(carbon, hydrogen) %>%
  step_spatialsign(carbon, hydrogen)

ss_obj <- prep(ss_trans, training = biomass_tr)

transformed_te <- bake(ss_obj, biomass_te)

plot(biomass_te$carbon, biomass_te$hydrogen)

plot(transformed_te$carbon, transformed_te$hydrogen)

tidy(ss_trans, number = 3)
tidy(ss_obj, number = 3)
```

---

step\_sqrt

*Square Root Transformation*

---

**Description**

step\_sqrt creates a *specification* of a recipe step that will square root transform the data.

**Usage**

```
step_sqrt(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  columns = NULL,
  skip = FALSE,
  id = rand_id("sqrt")
)

## S3 method for class 'step_sqrt'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will be transformed. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
columns	A character string of variable names that will be populated (eventually) by the terms argument.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_sqrt object.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms which is the columns that will be affected.

**See Also**

[step\\_logit\(\)](#) [step\\_invlogit\(\)](#) [step\\_log\(\)](#) [step\\_hyperbolic\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#)  
[bake.recipe\(\)](#)

**Examples**

```
set.seed(313)
examples <- matrix(rnorm(40)^2, ncol = 2)
examples <- as.data.frame(examples)

rec <- recipe(~ V1 + V2, data = examples)

sqrt_trans <- rec %>%
  step_sqrt(all_predictors())

sqrt_obj <- prep(sqrt_trans, training = examples)

transformed_te <- bake(sqrt_obj, examples)
plot(examples$V1, transformed_te$V1)

tidy(sqrt_trans, number = 1)
tidy(sqrt_obj, number = 1)
```



---

 step\_string2factor      *Convert Strings to Factors*


---

## Description

step\_string2factor will convert one or more character vectors to factors (ordered or unordered).

## Usage

```
step_string2factor(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  levels = NULL,
  ordered = FALSE,
  skip = FALSE,
  id = rand_id("string2factor")
)

## S3 method for class 'step_string2factor'
tidy(x, ...)
```

## Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables will converted to factors. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
levels	An options specification of the levels to be used for the new factor. If left NULL, the sorted unique values present when bake is called will be used.
ordered	A single logical value; should the factor(s) be ordered?
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_string2factor object.

**Details**

If `levels` is given, `step_string2factor` will convert all variables affected by this step to have the same levels.

Also, note that `prep` has an option `strings_as_factors` that defaults to `TRUE`. This should be changed so that raw character data will be applied to `step_string2factor`. However, this step can also take existing factors (but will leave them as-is).

**Value**

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `ordered`.

**See Also**

[step\\_factor2string\(\)](#) [step\\_dummy\(\)](#) [step\\_other\(\)](#) [step\\_novel\(\)](#)

**Examples**

```
library(modeldata)
data(okc)

rec <- recipe(~ diet + location, data = okc)

make_factor <- rec %>%
  step_string2factor(diet)
make_factor <- prep(make_factor,
                    training = okc,
                    strings_as_factors = FALSE)

# note that `diet` is a factor
juice(make_factor) %>% head
okc %>% head
tidy(make_factor, number = 1)
```

---

step\_unknown

*Assign missing categories to "unknown"*


---

**Description**

`step_unknown` creates a *specification* of a recipe step that will assign a missing value in a factor level to "unknown".

**Usage**

```
step_unknown(
  recipe,
  ...,
  role = NA,
```

```

    trained = FALSE,
    new_level = "unknown",
    objects = NULL,
    skip = FALSE,
    id = rand_id("unknown")
  )

  ## S3 method for class 'step_unknown'
  tidy(x, ...)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be affected by the step. These variables should be character or factor types. See <a href="#">selections()</a> for more details. For the <code>tidy</code> method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
new_level	A single character value that will be assigned to new factor levels.
objects	A list of objects that contain the information on factor levels that will be determined by <a href="#">prep.recipe()</a> .
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_unknown</code> object.

### Details

The selected variables are adjusted to have a new level (given by `new_level`) that is placed in the last position.

Note that if the original columns are character, they will be converted to factors by this step.

If `new_level` is already in the data given to `prep`, an error is thrown.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the columns that will be affected) and `value` (the factor levels that is used for the new value)

### See Also

[step\\_factor2string\(\)](#), [step\\_string2factor\(\)](#), [dummy\\_names\(\)](#), [step\\_regex\(\)](#), [step\\_count\(\)](#), [step\\_ordinalscore\(\)](#), [step\\_unorder\(\)](#), [step\\_other\(\)](#), [step\\_novel\(\)](#)

**Examples**

```
library(modeldata)
data(okc)

rec <-
  recipe(~ diet + location, data = okc) %>%
  step_unknown(diet, new_level = "unknown diet") %>%
  step_unknown(location, new_level = "unknown location") %>%
  prep()

table(juice(rec)$diet, okc$diet, useNA = "always") %>%
  as.data.frame() %>%
  dplyr::filter(Freq > 0)

tidy(rec, number = 1)
```

---

step\_unorder

---

*Convert Ordered Factors to Unordered Factors*


---

**Description**

step\_unorder creates a *specification* of a recipe step that will transform the data.

**Usage**

```
step_unorder(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  columns = NULL,
  skip = FALSE,
  id = rand_id("unorder")
)

## S3 method for class 'step_unorder'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.

columns	A character string of variable names that will be populated (eventually) by the terms argument.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_unorder</code> object.

### Details

The factors level order is preserved during the transformation.

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns terms (the columns that will be affected).

### See Also

[step\\_ordinalscore\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

### Examples

```
lmh <- c("Low", "Med", "High")

examples <- data.frame(X1 = factor(rep(letters[1:4], each = 3)),
                      X2 = ordered(rep(lmh, each = 4),
                                   levels = lmh))

rec <- recipe(~ X1 + X2, data = examples)

factor_trans <- rec %>%
  step_unorder(all_predictors())

factor_obj <- prep(factor_trans, training = examples)

transformed_te <- bake(factor_obj, examples)
table(transformed_te$X2, examples$X2)

tidy(factor_trans, number = 1)
tidy(factor_obj, number = 1)
```

step\_upsample

*Up-Sample a Data Set Based on a Factor Variable***Description**

step\_upsample creates a *specification* of a recipe step that will replicate rows of a data set to make the occurrence of levels in a specific factor level equal.

**Usage**

```
step_upsample(
  recipe,
  ...,
  over_ratio = 1,
  ratio = NA,
  role = NA,
  trained = FALSE,
  column = NULL,
  target = NA,
  skip = TRUE,
  seed = sample.int(10^5, 1),
  id = rand_id("upsample")
)

## S3 method for class 'step_upsample'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variable is used to sample the data. See <a href="#">selections()</a> for more details. The selection should result in <i>single factor variable</i> . For the tidy method, these are not currently used.
over_ratio	A numeric value for the ratio of the majority-to-minority frequencies. The default value (1) means that all other levels are sampled up to have the same frequency as the most occurring level. A value of 0.5 would mean that the minority levels will have (at most) (approximately) half as many rows than the majority level.
ratio	Deprecated argument; same as over_ratio.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
column	A character string of the variable name that will be populated (eventually) by the ... selectors.

target	An integer that will be used to subsample. This should not be set by the user and will be populated by prep.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
seed	An integer that will be used as the seed when upsampling.
id	A character string that is unique to this step to identify it.
x	A <code>step_upsample</code> object.

### Details

Up-sampling is intended to be performed on the *training* set alone. For this reason, the default is `skip = TRUE`. It is advisable to use `prep(recipe, retain = TRUE)` when preparing the recipe; in this way `juice()` can be used to obtain the up-sampled version of the data.

If there are missing values in the factor variable that is used to define the sampling, missing data are selected at random in the same way that the other factor levels are sampled. Missing values are not used to determine the amount of data in the majority level (see example below).

For any data with factor levels occurring with the same frequency as the majority level, all data will be retained.

All columns in the data are sampled and returned by `juice()` and `bake()`.

When used in modeling, users should strongly consider using the option `skip = TRUE` so that the extra sampling is *not* conducted outside of the training set.

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` which is the variable used to sample.

### Examples

```
library(modeldata)
data(okc)

orig <- table(okc$diet, useNA = "always")

sort(orig, decreasing = TRUE)

up_rec <- recipe(~ ., data = okc) %>%
  # Bring the minority levels up to about 200 each
  # 200/16562 is approx 0.0121
  step_upsample(diet, over_ratio = 0.0121) %>%
  prep(training = okc)

training <- table(juice(up_rec)$diet, useNA = "always")

# Since `skip` defaults to TRUE, baking the step has no effect
```

```

baked_okc <- bake(up_rec, new_data = okc)
baked <- table(baked_okc$diet, useNA = "always")

# Note that if the original data contained more rows than the
# target n (= ratio * majority_n), the data are left alone:
data.frame(
  level = names(orig),
  orig_freq = as.vector(orig),
  train_freq = as.vector(training),
  baked_freq = as.vector(baked)
)

```

---

step\_window

*Moving Window Functions*


---

### Description

step\_window creates a *specification* of a recipe step that will create new columns that are the results of functions that compute statistics across moving windows.

### Usage

```

step_window(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  size = 3,
  na_rm = TRUE,
  statistic = "mean",
  columns = NULL,
  names = NULL,
  skip = FALSE,
  id = rand_id("window")
)

## S3 method for class 'step_window'
tidy(x, ...)

```

### Arguments

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.



role	For model terms created by this step, what analysis role should they be assigned? If names is left to be NULL, the rolling statistics replace the original columns and the roles are left unchanged. If names is set, those new columns will have a role of NULL unless this argument has a value.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
size	An odd integer $\geq 3$ for the window size.
na_rm	A logical for whether missing values should be removed from the calculations within each window.
statistic	A character string for the type of statistic that should be calculated for each moving window. Possible values are: 'max', 'mean', 'median', 'min', 'prod', 'sd', 'sum', 'var'
columns	A character string that contains the names of columns that should be processed. These values are not determined until <code>prep.recipe()</code> is called.
names	An optional character string that is the same length of the number of terms selected by terms. If you are not sure what columns will be selected, use the summary function (see the example below). These will be the names of the new columns created by the step.
skip	A logical. Should the step be skipped when the recipe is baked by <code>bake.recipe()</code> ? While all operations are baked when <code>prep.recipe()</code> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A <code>step_window</code> object.

### Details

The calculations use a somewhat atypical method for handling the beginning and end parts of the rolling statistics. The process starts with the center justified window calculations and the beginning and ending parts of the rolling values are determined using the first and last rolling values, respectively. For example if a column `x` with 12 values is smoothed with a 5-point moving median, the first three smoothed values are estimated by `median(x[1:5])` and the fourth uses `median(x[2:6])`.

step will stop with a note about installing the package.

### Value

An updated version of `recipe` with the new step added to the sequence of existing steps (if any). For the `tidy` method, a tibble with columns `terms` (the selectors or variables selected) and `statistic` (the summary function name), and `size`.

### Examples

```
library(recipes)
library(dplyr)
library(rlang)
library(ggplot2, quietly = TRUE)
```

```

set.seed(5522)
sim_dat <- data.frame(x1 = (20:100) / 10)
n <- nrow(sim_dat)
sim_dat$y1 <- sin(sim_dat$x1) + rnorm(n, sd = 0.1)
sim_dat$y2 <- cos(sim_dat$x1) + rnorm(n, sd = 0.1)
sim_dat$x2 <- runif(n)
sim_dat$x3 <- rnorm(n)

rec <- recipe(y1 + y2 ~ x1 + x2 + x3, data = sim_dat) %>%
  step_window(starts_with("y"), size = 7, statistic = "median",
              names = paste0("med_7pt_", 1:2),
              role = "outcome") %>%
  step_window(starts_with("y"),
              names = paste0("mean_3pt_", 1:2),
              role = "outcome")
rec <- prep(rec, training = sim_dat)

# If you aren't sure how to set the names, see which variables are selected
# and the order that they are selected:
terms_select(info = summary(rec), terms = quos(starts_with("y")))

smoothed_dat <- bake(rec, sim_dat, everything())

ggplot(data = sim_dat, aes(x = x1, y = y1)) +
  geom_point() +
  geom_line(data = smoothed_dat, aes(y = med_7pt_1)) +
  geom_line(data = smoothed_dat, aes(y = mean_3pt_1), col = "red") +
  theme_bw()

tidy(rec, number = 1)
tidy(rec, number = 2)

# If you want to replace the selected variables with the rolling statistic
# don't set `names`
sim_dat$original <- sim_dat$y1
rec <- recipe(y1 + y2 + original ~ x1 + x2 + x3, data = sim_dat) %>%
  step_window(starts_with("y"))
rec <- prep(rec, training = sim_dat)
smoothed_dat <- bake(rec, sim_dat, everything())
ggplot(smoothed_dat, aes(x = original, y = y1)) +
  geom_point() +
  theme_bw()

```

---

step\_YeoJohnson

*Yeo-Johnson Transformation*


---

### Description

step\_YeoJohnson creates a *specification* of a recipe step that will transform data using a simple Yeo-Johnson transformation.

**Usage**

```
step_YeoJohnson(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  lambdas = NULL,
  limits = c(-5, 5),
  num_unique = 5,
  na_rm = TRUE,
  skip = FALSE,
  id = rand_id("YeoJohnson")
)

## S3 method for class 'step_YeoJohnson'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables are affected by the step. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
lambdas	A numeric vector of transformation values. This is NULL until computed by <a href="#">prep.recipe()</a> .
limits	A length 2 numeric vector defining the range to compute the transformation parameter lambda.
num_unique	An integer where data that have less possible values will not be evaluate for a transformation.
na_rm	A logical value indicating whether NA values should be removed during computations.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using <code>skip = TRUE</code> as it may affect the computations for subsequent operations
id	A character string that is unique to this step to identify it.
x	A step_YeoJohnson object.

**Details**

The Yeo-Johnson transformation is very similar to the Box-Cox but does not require the input variables to be strictly positive. In the package, the partial log-likelihood function is directly optimized within a reasonable set of transformation values (which can be changed by the user).

This transformation is typically done on the outcome variable using the residuals for a statistical model (such as ordinary least squares). Here, a simple null model (intercept only) is used to apply the transformation to the *predictor* variables individually. This can have the effect of making the variable distributions more symmetric.

If the transformation parameters are estimated to be very closed to the bounds, or if the optimization fails, a value of NA is used and no transformation is applied.

### Value

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms (the selectors or variables selected) and value (the lambda estimate).

### References

Yeo, I. K., and Johnson, R. A. (2000). A new family of power transformations to improve normality or symmetry. *Biometrika*.

### See Also

[step\\_BoxCox\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

### Examples

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
              data = biomass_tr)

yj_transform <- step_YeoJohnson(rec, all_numeric())

yj_estimates <- prep(yj_transform, training = biomass_tr)

yj_te <- bake(yj_estimates, biomass_te)

plot(density(biomass_te$sulfur), main = "before")
plot(density(yj_te$sulfur), main = "after")

tidy(yj_transform, number = 1)
tidy(yj_estimates, number = 1)
```

step\_zv

*Zero Variance Filter***Description**

step\_zv creates a *specification* of a recipe step that will remove variables that contain only a single value.

**Usage**

```
step_zv(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  removals = NULL,
  skip = FALSE,
  id = rand_id("zv")
)

## S3 method for class 'step_zv'
tidy(x, ...)
```

**Arguments**

recipe	A recipe object. The step will be added to the sequence of operations for this recipe.
...	One or more selector functions to choose which variables that will be evaluated by the filtering. See <a href="#">selections()</a> for more details. For the tidy method, these are not currently used.
role	Not used by this step since no new variables are created.
trained	A logical to indicate if the quantities for preprocessing have been estimated.
removals	A character string that contains the names of columns that should be removed. These values are not determined until <a href="#">prep.recipe()</a> is called.
skip	A logical. Should the step be skipped when the recipe is baked by <a href="#">bake.recipe()</a> ? While all operations are baked when <a href="#">prep.recipe()</a> is run, some operations may not be able to be conducted on new data (e.g. processing the outcome variable(s)). Care should be taken when using skip = TRUE as it may affect the computations for subsequent operations.
id	A character string that is unique to this step to identify it.
x	A step_zv object.

**Value**

An updated version of recipe with the new step added to the sequence of existing steps (if any). For the tidy method, a tibble with columns terms which is the columns that will be removed.

**See Also**

[step\\_nzv\(\)](#) [step\\_corr\(\)](#) [recipe\(\)](#) [prep.recipe\(\)](#) [bake.recipe\(\)](#)

**Examples**

```
library(modeldata)
data(biomass)

biomass$one_value <- 1

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

rec <- recipe(HHV ~ carbon + hydrogen + oxygen +
              nitrogen + sulfur + one_value,
              data = biomass_tr)

zv_filter <- rec %>%
  step_zv(all_predictors())

filter_obj <- prep(zv_filter, training = biomass_tr)

filtered_te <- bake(filter_obj, biomass_te)
any(names(filtered_te) == "one_value")

tidy(zv_filter, number = 1)
tidy(filter_obj, number = 1)
```

---

summary.recipe

*Summarize a Recipe*

---

**Description**

This function prints the current set of variables/features and some of their characteristics.

**Usage**

```
## S3 method for class 'recipe'
summary(object, original = FALSE, ...)
```

**Arguments**

object	A recipe object
original	A logical: show the current set of variables or the original set when the recipe was defined.
...	further arguments passed to or from other methods (not currently used).

**Details**

Note that, until the recipe has been trained, the current and original variables are the same.

It is possible for variables to have multiple roles by adding them with `add_role()`. If a variable has multiple roles, it will have more than one row in the summary tibble.

**Value**

A tibble with columns `variable`, `type`, `role`, and `source`.

**See Also**

[recipe\(\)](#) [prep.recipe\(\)](#)

**Examples**

```
rec <- recipe( ~ ., data = USArrests)
summary(rec)
rec <- step_pca(rec, all_numeric(), num = 3)
summary(rec) # still the same since not yet trained
rec <- prep(rec, training = USArrests)
summary(rec)
```

---

terms\_select

*Select Terms in a Step Function.*


---

**Description**

This function bakes the step function selectors and might be useful when creating custom steps.

**Usage**

```
terms_select(terms, info, empty_fun = abort_selection)
```

**Arguments**

terms	A list of formulas whose right-hand side contains quoted expressions. See <a href="#">rlang::quos()</a> for examples.
info	A tibble with columns <code>variable</code> , <code>type</code> , <code>role</code> , and <code>source</code> that represent the current state of the data. The function <a href="#">summary.recipe()</a> can be used to get this information from a recipe.
empty_fun	A function to execute when no terms are selected by the step. The default function throws an error with a message.

**Value**

A character string of column names or an error if there are no selectors or if no variables are selected.

**See Also**

`recipe()` `summary.recipe()` `prep.recipe()`

**Examples**

```
library(rlang)
library(modeldata)
data(okc)
rec <- recipe(~ ., data = okc)
info <- summary(rec)
terms_select(info = info, quos(all_predictors()))
```

---

tidy.recipe

*Tidy the Result of a Recipe*


---

**Description**

tidy will return a data frame that contains information regarding a recipe or operation within the recipe (when a tidy method for the operation exists).

**Usage**

```
## S3 method for class 'recipe'
tidy(x, number = NA, id = NA, ...)

## S3 method for class 'step'
tidy(x, ...)

## S3 method for class 'check'
tidy(x, ...)
```

**Arguments**

x	A recipe object (trained or otherwise).
number	An integer or NA. If missing and id is not provided, the return value is a list of the operations in the recipe. If a number is given, a tidy method is executed for that operation in the recipe (if it exists). number must not be provided if id is.
id	A character string or NA. If missing and number is not provided, the return value is a list of the operations in the recipe. If a character string is given, a tidy method is executed for that operation in the recipe (if it exists). id must not be provided if number is.
...	Not currently used.



**Value**

A tibble with columns that would vary depending on what tidy method is executed. When `number` and `id` are NA, a tibble with columns `number` (the operation iteration), `operation` (either "step" or "check"), `type` (the method, e.g. "nzv", "center"), a logical column called `trained` for whether the operation has been estimated using `prep`, a logical for `skip`, and a character column `id`.

**Examples**

```
library(modeldata)
data(okc)

okc_rec <- recipe(~ ., data = okc) %>%
  step_other(all_nominal(), threshold = 0.05, other = "another") %>%
  step_date(date, features = "dow") %>%
  step_center(all_numeric()) %>%
  step_dummy(all_nominal()) %>%
  check_cols(starts_with("date"), age, height)

tidy(okc_rec)

tidy(okc_rec, number = 2)
tidy(okc_rec, number = 3)

okc_rec_trained <- prep(okc_rec, training = okc)

tidy(okc_rec_trained)
tidy(okc_rec_trained, number = 3)
```

---

 update.step

*Update a recipe step*


---

**Description**

This step method for `update()` takes named arguments as `...` whose values will replace the elements of the same name in the actual step.

**Usage**

```
## S3 method for class 'step'
update(object, ...)
```

**Arguments**

<code>object</code>	A recipe step.
<code>...</code>	Key-value pairs where the keys match up with names of elements in the step, and the values are the new values to update the step with.

## Details

For a step to be updated, it must not already have been trained. Otherwise, conflicting information can arise between the data returned from `juice()` and the information in the step.

## Examples

```
library(modeldata)
data(biomass)

biomass_tr <- biomass[biomass$dataset == "Training",]
biomass_te <- biomass[biomass$dataset == "Testing",]

# Create a recipe using step_bs() with degree = 3
rec <- recipe(
  HHV ~ carbon + hydrogen + oxygen + nitrogen + sulfur,
  data = biomass_tr
) %>%
  step_bs(carbon, hydrogen, degree = 3)

# Update the step to use degree = 4
rec2 <- rec
rec2$steps[[1]] <- update(rec2$steps[[1]], degree = 4)

# Prep both recipes
rec_prepped <- prep(rec, training = biomass_tr)
rec2_prepped <- prep(rec2, training = biomass_tr)

# Juice both to see what changed
juice(rec_prepped)
juice(rec2_prepped)

# Cannot update a recipe step that has been trained!
## Not run:
update(rec_prepped$steps[[1]], degree = 4)

## End(Not run)
```

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