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adasyn  Adaptive Synthetic Algorithm

Description

Generates synthetic positive instances using ADASYN algorithm.

Usage

adasyn(df, var, k = 5, over_ratio = 1)

Arguments

df data.frame or tibble. Must have 1 factor variable and remaining numeric variables.

var Character, name of variable containing factor variable.

k An integer. Number of nearest neighbor that are used to generate the new examples of the minority class.

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.
Details

All columns used in this function must be numeric with no missing data.

Value

A data.frame or tibble, depending on type of `df`.

References


See Also

`step_adasyn()` for step function of this method
Other Direct Implementations: `bsmote()`, `nearmiss()`, `smotenc()`, `smote()`, `tomek()`

Examples

circle_numeric <- circle_example[, c("x", "y", "class")]
res <- adasyn(circle_numeric, var = "class")
res <- adasyn(circle_numeric, var = "class", k = 10)
res <- adasyn(circle_numeric, var = "class", over_ratio = 0.8)

bsmote

`borderline SMOTE Algorithm`

Description

BSMOTE generates generate new examples of the minority class using nearest neighbors of these cases in the border region between classes.

Usage

bsmote(df, var, k = 5, over_ratio = 1, all_neighbors = FALSE)

Arguments

df data.frame or tibble. Must have 1 factor variable and remaining numeric variables.

var Character, name of variable containing factor variable.

k An integer. Number of nearest neighbor that are used to generate the new examples of the minority class.
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.

all_neighbors

Type of two borderline-SMOTE method. Defaults to FALSE. See details.

Details

This method works the same way as `smote()`, except that instead of generating points around every point of the minority class each point is first being classified into the boxes "danger" and "not". For each point the k nearest neighbors is calculated. If all the neighbors comes from a different class it is labeled noise and put in to the "not" box. If more then half of the neighbors comes from a different class it is labeled "danger."

If `all_neighbors = FALSE` then points will be generated between nearest neighbors in its own class. If `all_neighbors = TRUE` then points will be generated between any nearest neighbors. See examples for visualization.

The parameter `neighbors` controls the way the new examples are created. For each currently existing minority class example X new examples will be created (this is controlled by the parameter `over_ratio` as mentioned above). These examples will be generated by using the information from the `neighbors` nearest neighbor of each example of the minority class. The parameter `neighbors` controls how many of these neighbor are used.

All columns used in this step must be numeric with no missing data.

Value

A data.frame or tibble, depending on type of `df`.

References


See Also

`step_bsmote()` for step function of this method

Other Direct Implementations: `adasyn()`, `nearmiss()`, `smotenc()`, `smote()`, `tomek()`

Examples

circle_numeric <- circle_example[, c("x", "y", "class")]

res <- bsmote(circle_numeric, var = "class")

res <- bsmote(circle_numeric, var = "class", k = 10)

res <- bsmote(circle_numeric, var = "class", over_ratio = 0.8)
```r
res <- bsmote(circle_numeric, var = "class", all_neighbors = TRUE)
```

---

**circle_example**  
*Synthetic Dataset With a Circle*

**Description**  
A random dataset with two classes one of which is inside a circle. Used for examples to show how the different methods handles borders.

**Usage**  
circle_example

**Format**  
A data frame with 200 rows and 4 variables:
- **x**  Numeric.
- **y**  Numeric.
- **class**  Factor, values "Circle" and "Rest".
- **id**  character, ID variable.

---

**nearmiss**  
*Remove Points Near Other Classes*

**Description**  
Generates synthetic positive instances using nearmiss algorithm.

**Usage**  
nearmiss(df, var, k = 5, under_ratio = 1)

**Arguments**
- **df**  data.frame or tibble. Must have 1 factor variable and remaining numeric variables.
- **var**  Character, name of variable containing factor variable.
- **k**  An integer. Number of nearest neighbor that are used to generate the new examples of the minority class.
- **under_ratio**  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.
smote

Details

All columns used in this function must be numeric with no missing data.

Value

A data.frame or tibble, depending on type of df.

References


See Also

step_nearmiss() for step function of this method
Other Direct Implementations: adasyn(), bsmote(), smotenc(), smote(), tomek()

Examples

circle_numeric <- circle_example[, c("x", "y", "class")]
res <- nearmiss(circle_numeric, var = "class")
res <- nearmiss(circle_numeric, var = "class", k = 10)
res <- nearmiss(circle_numeric, var = "class", under_ratio = 1.5)

---

smote

SMOTE Algorithm

Description

SMOTE generates new examples of the minority class using nearest neighbors of these cases.

Usage

smote(df, var, k = 5, over_ratio = 1)

Arguments

df data.frame or tibble. Must have 1 factor variable and remaining numeric variables.

var Character, name of variable containing factor variable.

k An integer. Number of nearest neighbor that are used to generate the new examples of the minority class.
SMOTENC Algorithm

**Description**

SMOTENC generates new examples of the minority class using nearest neighbors of these cases, and can handle categorical variables.

**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.

**Details**

The parameter `neighbors` controls the way the new examples are created. For each currently existing minority class example X new examples will be created (this is controlled by the parameter `over_ratio` as mentioned above). These examples will be generated by using the information from the neighbors nearest neighbor of each example of the minority class. The parameter `neighbors` controls how many of these neighbor are used. All columns used in this function must be numeric with no missing data.

**Value**

A data.frame or tibble, depending on type of `df`.

**References**


**See Also**

- `step_smote()` for step function of this method
- Other Direct Implementations: `adasyn()`, `bsmote()`, `nearmiss()`, `smotenc()`, `tomek()`

**Examples**

```r
circle_numeric <- circle_example[, c("x", "y", "class")]
res <- smote(circle_numeric, var = "class")
res <- smote(circle_numeric, var = "class", k = 10)
res <- smote(circle_numeric, var = "class", over_ratio = 0.8)
```
Usage

smotenc(df, var, k = 5, over_ratio = 1)

Arguments

df  data.frame or tibble. Must have 1 factor variable and remaining numeric variables.
var  Character, name of variable containing factor variable.
k  An integer. Number of nearest neighbor that are used to generate the new examples of the minority class.
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.

Details

The parameter `neighbors` controls the way the new examples are created. For each currently existing minority class example X new examples will be created (this is controlled by the parameter `over_ratio` as mentioned above). These examples will be generated by using the information from the `neighbors` nearest neighbor of each example of the minority class. The parameter `neighbors` controls how many of these neighbor are used. Columns can be numeric and categorical with no missing data.

Value

A data.frame or tibble, depending on type of `df`.

References


See Also

`step_smotenc()` for step function of this method
Other Direct Implementations: `adasyn()`, `bsmote()`, `nearmiss()`, `smote()`, `tomek()`

Examples

circle_numeric <- circle_example[, c("x", "y", "class")]
res <- smotenc(circle_numeric, var = "class")
res <- smotenc(circle_numeric, var = "class", k = 10)
res <- smotenc(circle_numeric, var = "class", over_ratio = 0.8)
## Description

`step_adasyn` creates a specification of a recipe step that generates synthetic positive instances using ADASYN algorithm.

## Usage

```r
step_adasyn(
  recipe,
  ..., 
  role = NA, 
  trained = FALSE, 
  column = NULL, 
  over_ratio = 1, 
  neighbors = 5, 
  skip = TRUE, 
  seed = sample.int(10^5, 1), 
  id = rand_id("adasyn")
)
```

## 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 `selections()` for more details. The selection should result in a single factor variable. 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.
- **column**: A character string of the variable name that will be populated (eventually) by the ... selectors.
- **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.
- **neighbors**: An integer. Number of nearest neighbor that are used to generate the new examples of the minority class.
- **skip**: A logical. Should the step be skipped when the recipe is baked by `bake()`? While all operations are baked when `prep()` 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.
seed  
An integer that will be used as the seed when applied.

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

Details

All columns in the data are sampled and returned by `juice()` and `bake()`.
All columns used in this step must be numeric with no missing data.
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.

Tidying

When you `tidy()` this step, a tibble with columns `terms` (the selectors or variables selected) will be returned.

Case weights

The underlying operation does not allow for case weights.

References


See Also

`adasyn()` for direct implementation
Other Steps for over-sampling: `step_bsmote()`, `step_rose()`, `step_smotenc()`, `step_smote()`, `step_upsample()`

Examples

```
library(recipes)
library(modeldata)
data(hpc_data)

hpc_data0 <- hpc_data %>%
  select(-protocol, -day)

orig <- count(hpc_data0, class, name = "orig")
orig

up_rec <- recipe(class ~ ., data = hpc_data0) %>%
  # Bring the minority levels up to about 1000 each
# 1000/2211 is approx 0.4523
step_adasyn(class, over_ratio = 0.4523) %>%
  prep()

training <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "training")
training

# Since `skip` defaults to TRUE, baking the step has no effect
baked <- up_rec %>%
  bake(new_data = hpc_data0) %>%
  count(class, name = "baked")
baked

# Note that if the original data contained more rows than the
# target n (= ratio * majority_n), the data are left alone:
orig %>%
  left_join(training, by = "class") %>%
  left_join(baked, by = "class")

library(ggplot2)

ggplot(circle_example, aes(x, y, color = class)) +
  geom_point() +
  labs(title = "Without ADASYN")

recipe(class ~ x + y, data = circle_example) %>%
  step_adasyn(class) %>%
  prep() %>%
  bake(new_data = NULL) %>%
  ggplot(aes(x, y, color = class)) +
  geom_point() +
  labs(title = "With ADASYN")

---

**step_bsmote**

**Apply borderline-SMOTE Algorithm**

**Description**

*step_bsmote* creates a *specification* of a recipe step that generate new examples of the minority class using nearest neighbors of these cases in the border region between classes.

**Usage**

```r
step_bsmote(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
)```
step_bsmote

column = NULL,
over_ratio = 1,
neighbors = 5,
all_neighbors = FALSE,
skip = TRUE,
seed = sample.int(10^5, 1),
id = rand_id("bsmote")
)

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 selections() for more details. The selection should result in single factor variable. 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.
column  A character string of the variable name that will be populated (eventually) by the ... selectors.
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.
neighbors  An integer. Number of nearest neighbor that are used to generate the new examples of the minority class.
all_neighbors Type of two borderline-SMOTE method. Defaults to FALSE. See details.
skip  A logical. Should the step be skipped when the recipe is baked by bake()? While all operations are baked when prep() 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.
seed  An integer that will be used as the seed when smote-ing.
id  A character string that is unique to this step to identify it.

Details

This methods works the same way as step_smote(), expect that instead of generating points around every point of the minority class each point is first being classified into the boxes "danger" and "not". For each point the k nearest neighbors is calculated. If all the neighbors comes from a different class it is labeled noise and put in to the "not" box. If more then half of the neighbors comes from a different class it is labeled "danger.

If all_neighbors = FALSE then points will be generated between nearest neighbors in its own class. If all_neighbors = TRUE then points will be generated between any nearest neighbors. See examples for visualization.
The parameter `neighbors` controls the way the new examples are created. For each currently existing minority class example X new examples will be created (this is controlled by the parameter `over_ratio` as mentioned above). These examples will be generated by using the information from the neighbors nearest neighbor of each example of the minority class. The parameter `neighbors` controls how many of these neighbor are used.

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

All columns used in this step must be numeric with no missing data.

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.

**Tidying**

When you `tidy()` this step, a tibble with columns `terms` (the selectors or variables selected) will be returned.

**Case weights**

The underlying operation does not allow for case weights.

**References**


**See Also**

`bsmote()` for direct implementation

Other Steps for over-sampling: `step_adasyn()`, `step_rose()`, `step_smotenc()`, `step_smote()`, `step_upsample()`

**Examples**

```r
library(recipes)
library(modeldata)
data(hpc_data)

hpc_data0 <- hpc_data %>%
  select(-protocol, -day)

orig <- count(hpc_data0, class, name = "orig")
orig

up_rec <- recipe(class ~ ., data = hpc_data0) %>%
```
# Bring the minority levels up to about 1000 each
# 1000/2211 is approx 0.4523
step_bsmote(class, over_ratio = 0.4523) %>%
  prep()

training <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "training")

# Since `skip` defaults to TRUE, baking the step has no effect
baked <- up_rec %>%
  bake(new_data = hpc_data0) %>%
  count(class, name = "baked")

# Note that if the original data contained more rows than the
# target n (= ratio * majority_n), the data are left alone:
orig %>%
  left_join(training, by = "class") %>%
  left_join(baked, by = "class")

library(ggplot2)

ggplot(circle_example, aes(x, y, color = class)) +
  geom_point() +
  labs(title = "Without SMOTE")

recipe(class ~ x + y, data = circle_example) %>%
  step_bsmote(class, all_neighbors = FALSE) %>%
  prep() %>%
  bake(new_data = NULL) %>%
  ggplot(aes(x, y, color = class)) +
  geom_point() +
  labs(title = "With borderline-SMOTE, all_neighbors = FALSE")

recipe(class ~ x + y, data = circle_example) %>%
  step_bsmote(class, all_neighbors = TRUE) %>%
  prep() %>%
  bake(new_data = NULL) %>%
  ggplot(aes(x, y, color = class)) +
  geom_point() +
  labs(title = "With borderline-SMOTE, all_neighbors = TRUE")

### `step_downsample` - Down-Sample a Data Set Based on a Factor Variable

**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

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

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 `selections()` for more details. The selection should result in single factor variable. For the tidy method, these are not currently used.
- **under_ratio**: 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.
- **ratio**: Deprecated argument; same as `under_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 `bake()`? While all operations are baked when `prep()` 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.
- **seed**: An integer that will be used as the seed when downsampling.
- **id**: A character string that is unique to this step to identify it.

Details

Down-sampling is intended to be performed on the **training** set alone. For this reason, the default is `skip = TRUE`. 
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.

**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.

**Tidying**

When you `tidy()` this step, a tibble with columns `terms` (the selectors or variables selected) will be returned.

**Case weights**

This step performs an unsupervised operation that can utilize case weights. To use them, see the documentation in `recipes::case_weights` and the examples on tidymodels.org.

**See Also**

Other Steps for under-sampling: `step_nearmiss()`, `step_tomek()`

**Examples**

```r
library(recipes)
library(modeldata)
data(hpc_data)

hpc_data0 <- hpc_data %>%
  select(-protocol, -day)

orig <- count(hpc_data0, class, name = "orig")
orig

up_rec <- recipe(class ~ ., data = hpc_data0) %>%
  step_downsample(class, under_ratio = 3.862) %>%
  prep()

training <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "training")
```
step_nearmiss

training

# Since 'skip' defaults to TRUE, baking the step has no effect
baked <- up_rec %>%
  bake(new_data = hpc_data0) %>%
  count(class, name = "baked")

# Note that if the original data contained more rows than the
# target n (= ratio * majority_n), the data are left alone:
orig %>%
  left_join(training, by = "class") %>%
  left_join(baked, by = "class")

library(ggplot2)

ggplot(circle_example, aes(x, y, color = class)) +
  geom_point() +
  labs(title = "Without downsample")

recipe(class ~ x + y, data = circle_example) %>%
  step_downsample(class) %>%
  prep() %>%
  bake(new_data = NULL) %>%
  ggplot(aes(x, y, color = class)) +
  geom_point() +
  labs(title = "With downsample")

---

**step_nearmiss**  
*Remove Points Near Other Classes*

**Description**

*step_nearmiss* creates a *specification* of a recipe step that removes majority class instances by undersampling points in the majority class based on their distance to other points in the same class.

**Usage**

```r
step_nearmiss(
  recipe,
  ..., 
  role = NA,
  trained = FALSE,
  column = NULL,
  under_ratio = 1,
  neighbors = 5,
  skip = TRUE,
  seed = sample.int(10^5, 1),
  id = rand_id("nearmiss")
)
```
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 `selections()` for more details. The selection should result in a single factor variable. 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.
- **column**: A character string of the variable name that will be populated (eventually) by the ... selectors.
- **under_ratio**: 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.
- **neighbors**: An integer. Number of nearest neighbor that are used to generate the new examples of the minority class.
- **skip**: A logical. Should the step be skipped when the recipe is baked by `bake()`? While all operations are baked when `prep()` 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.
- **seed**: An integer that will be used as the seed when applied.
- **id**: A character string that is unique to this step to identify it.

Details

This method retained the points form the majority classes which has the smallest mean distance to the k nearest points in the other classes.

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

All columns used in this step must be numeric with no missing data.

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.

Tidying

When you `tidy()` this step, a tibble with columns `terms` (the selectors or variables selected) will be returned.
Case weights

The underlying operation does not allow for case weights.

References


See Also

`nearmiss()` for direct implementation

Other Steps for under-sampling: `step_downsample()`, `step_tomek()`

Examples

```r
library(recipes)
library(modeldata)
data(hpc_data)

hpc_data0 <- hpc_data %>%
  select(-protocol, -day)

orig <- count(hpc_data0, class, name = "orig")
orig

up_rec <- recipe(class ~ ., data = hpc_data0) %>%
  # Bring the majority levels down to about 1000 each
  # 1000/259 is approx 3.862
  step_nearmiss(class, under_ratio = 3.862) %>%
  prep()

training <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "training")
training

# Since `skip` defaults to TRUE, baking the step has no effect
baked <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "baked")
baked

# Note that if the original data contained more rows than the
# target n (= ratio * majority_n), the data are left alone:
orig %>%
  left_join(training, by = "class") %>%
  left_join(baked, by = "class")

library(ggplot2)

ggplot(circle_example, aes(x, y, color = class)) +
```
```
geom_point() +
labs(title = "Without NEARMISS") +
xlim(c(1, 15)) +
ylim(c(1, 15))

recipe(class ~ x + y, data = circle_example) %>%
  step_nearmiss(class) %>%
  prep() %>%
  bake(new_data = NULL) %>%
  ggplot(aes(x, y, color = class)) +
  geom_point() +
  labs(title = "With NEARMISS") +
  xlim(c(1, 15)) +
  ylim(c(1, 15))
```

---

**step_rose**  
*Apply ROSE Algorithm*

**Description**

`step_rose` creates a *specification* of a recipe step that generates sample of synthetic data by enlarging the features space of minority and majority class example. Using `ROSE::ROSE()`.

**Usage**

```r
step_rose(
  recipe,
  ..., 
  role = NA,
  trained = FALSE,
  column = NULL,
  over_ratio = 1,
  minority_prop = 0.5,
  minority_smoothness = 1,
  majority_smoothness = 1,
  skip = TRUE,
  seed = sample.int(10^5, 1),
  id = rand_id("rose")
)
```

**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 `selections()` for more details. The selection should result in *single factor variable*. For the tidy method, these are not currently used.
- `role` Not used by this step since no new variables are created.
## step_rose

**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.

**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.

**minority_prop**  
A numeric. Determines the of over-sampling of the minority class. Defaults to 0.5.

**minority_smoothness**  
A numeric. Shrink factor to be multiplied by the smoothing parameters to estimate the conditional kernel density of the minority class. Defaults to 1.

**majority_smoothness**  
A numeric. Shrink factor to be multiplied by the smoothing parameters to estimate the conditional kernel density of the majority class. Defaults to 1.

**skip**  
A logical. Should the step be skipped when the recipe is baked by `bake()`? While all operations are baked when `prep()` 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.

**seed**  
An integer that will be used as the seed when rose-ing.

**id**  
A character string that is unique to this step to identify it.

### Details

The factor variable used to balance around must only have 2 levels.

The ROSE algorithm works by selecting an observation belonging to class k and generates new examples in its neighborhood is determined by some matrix $H_k$. Smaller values of these arguments have the effect of shrinking the entries of the corresponding smoothing matrix $H_k$. Shrinking would be a cautious choice if there is a concern that excessively large neighborhoods could lead to blur the boundaries between the regions of the feature space associated with each class.

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.

### Tidying

When you `tidy()` this step, a tibble with columns `terms` (the selectors or variables selected) will be returned.
Case weights

The underlying operation does not allow for case weights.

References


See Also

Other Steps for over-sampling: step_adasyn(), step_bsmote(), step_smotenc(), step_smote(), step_upsample()

Examples

library(recipes)
library(modeldata)
data(hpc_data)

hpc_data0 <- hpc_data %>%
  mutate(class = factor(class == "VF", labels = c("not VF", "VF"))) %>%
  select(-protocol, -day)

orig <- count(hpc_data0, class, name = "orig")
orig

up_rec <- recipe(class ~ ., data = hpc_data0) %>%
  step_rose(class) %>%
  prep()

training <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "training")
training

# Since `skip` defaults to TRUE, baking the step has no effect
baked <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "baked")
baked

orig %>%
  left_join(training, by = "class") %>%
  left_join(baked, by = "class")

library(ggplot2)

ggplot(circle_example, aes(x, y, color = class)) +
  geom_point() +
step_smote

labs(title = "Without ROSE")

recipe(class ~ x + y, data = circle_example) %>%
  step_rose(class) %>%
  prep() %>%
  bake(new_data = NULL) %>%
  ggplot(aes(x, y, color = class)) +
  geom_point() +
  labs(title = "With ROSE")

---

**step_smote**  
*Apply SMOTE Algorithm*

**Description**

`step_smote` creates a specification of a recipe step that generate new examples of the minority class using nearest neighbors of these cases.

**Usage**

```r
step_smote(
  recipe, ...
  role = NA, trained = FALSE, column = NULL, over_ratio = 1, neighbors = 5, skip = TRUE, 
  seed = sample.int(10^5, 1), id = rand_id("smote")
)
```

**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 `selections()` for more details. The selection should result in single factor variable. 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.
- **column**: A character string of the variable name that will be populated (eventually) by the ... selectors.
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.

neighbors An integer. Number of nearest neighbor that are used to generate the new examples of the minority class.

skip A logical. Should the step be skipped when the recipe is baked by bake()? While all operations are baked when prep() 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.

seed An integer that will be used as the seed when smote-ing.

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

Details

The parameter neighbors controls the way the new examples are created. For each currently existing minority class example X new examples will be created (this is controlled by the parameter over_ratio as mentioned above). These examples will be generated by using the information from the neighbors nearest neighbor of each example of the minority class. The parameter neighbors controls how many of these neighbor are used.

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

All columns used in this step must be numeric with no missing data.

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.

Tidying

When you tidy() this step, a tibble with columns terms (the selectors or variables selected) will be returned.

Case weights

The underlying operation does not allow for case weights.

References

See Also

`smote()` for direct implementation

Other Steps for over-sampling: `step_adasyn()`, `step_bsmote()`, `step_rose()`, `step_smotenc()`, `step_upsample()`

Examples

```r
library(recipes)
library(modeldata)
data(hpc_data)

hpc_data0 <- hpc_data %>%
  select(-protocol, -day)

orig <- count(hpc_data0, class, name = "orig")
orig

up_rec <- recipe(class ~ ., data = hpc_data0) %>%
  step_smote(class, over_ratio = 0.4523) %>%
  prep()

training <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "training")
training

# Since `skip` defaults to TRUE, baking the step has no effect
baked <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "baked")
baked

# Note that if the original data contained more rows than the
# target n (= ratio * majority_n), the data are left alone:
orig %>%
  left_join(training, by = "class") %>%
  left_join(baked, by = "class")

library(ggplot2)

ggplot(circle_example, aes(x, y, color = class)) +
  geom_point() +
  labs(title = "Without SMOTE")

recipe(class ~ x + y, data = circle_example) %>%
  step_smote(class) %>%
  prep() %>%
  bake(new_data = NULL) %>%
  ggplot(aes(x, y, color = class)) +
  geom_point() +
```
step_smotenc

Description

step_smotenc creates a specification of a recipe step that generate new examples of the minority class using nearest neighbors of these cases. Gower’s distance is used to handle mixed data types. For categorical variables, the most common category along neighbors is chosen.

Usage

step_smotenc(
  recipe,
  ...,
  role = NA,
  trained = FALSE,
  column = NULL,
  over_ratio = 1,
  neighbors = 5,
  skip = TRUE,
  seed = sample.int(10^5, 1),
  id = rand_id("smotenc")
)

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 selections() for more details. The selection should result in single factor variable. 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.
- **column**: A character string of the variable name that will be populated (eventually) by the ... selectors.
- **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.
- **neighbors**: An integer. Number of nearest neighbors that are used to generate the new examples of the minority class.
step_smotenc

skip A logical. Should the step be skipped when the recipe is baked by `bake()`? While all operations are baked when `prep()` 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.

seed An integer that will be used as the seed when smote-ing.

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

Details

The parameter `neighbors` controls the way the new examples are created. For each currently existing minority class example X new examples will be created (this is controlled by the parameter `over_ratio` as mentioned above). These examples will be generated by using the information from the `neighbors` nearest neighbor of each example of the minority class. The parameter `neighbors` controls how many of these neighbor are used.

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

Columns can be numeric and categorical with no missing data.

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.

Tidying

When you `tidy()` this step, a tibble with columns `terms` (the selectors or variables selected) will be returned.

Case weights

The underlying operation does not allow for case weights.

References


See Also

`smotenc()` for direct implementation

Other Steps for over-sampling: `step_adasyn()`, `step_bsmote()`, `step_rose()`, `step_smote()`, `step_upsample()`
Examples

library(recipes)
library(modeldata)
data(hpc_data)

orig <- count(hpc_data, class, name = "orig")
orign

up_rec <- recipe(class ~ ., data = hpc_data) %>%
  step_impute_knn(all_predictors()) %>%
  # Bring the minority levels up to about 1000 each
  # 1000/2211 is approx 0.4523
  step_smotenc(class, over_ratio = 0.4523) %>%
  prep()

training <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "training")
training

# Since 'skip' defaults to TRUE, baking the step has no effect
baked <- up_rec %>%
  bake(new_data = hpc_data) %>%
  count(class, name = "baked")
baked

# Note that if the original data contained more rows than the
# target n (= ratio * majority_n), the data are left alone:
orig %>%
  left_join(training, by = "class") %>%
  left_join(baked, by = "class")

---

step_tomek

Remove Tomek's Links

Description

step_tomek creates a specification of a recipe step that removes majority class instances of tomek links.

Usage

step_tomek(
  recipe,
  ..., 
  role = NA,
  trained = FALSE,
  column = NULL,
  skip = TRUE,
seed = sample.int(10^5, 1),
id = rand_id("tomek")
)

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 selections() for more details. The selection should result in single factor variable. 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.

column  A character string of the variable name that will be populated (eventually) by the ... selectors.

skip  A logical. Should the step be skipped when the recipe is baked by bake()? While all operations are baked when prep() 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.

seed  An integer that will be used as the seed when applied.

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

Details

The factor variable used to balance around must only have 2 levels. All other variables must be numerics with no missing data.

A tomek link is defined as a pair of points from different classes and are each others nearest neighbors.

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.

Tidying

When you tidy() this step, a tibble with columns terms (the selectors or variables selected) will be returned.

Case weights

The underlying operation does not allow for case weights.
References


See Also
tomek() for direct implementation
Other Steps for under-sampling: step_downsample(), step_nearmiss()

Examples

```r
library(recipes)
library(modeldata)
data(hpc_data)

hpc_data0 <- hpc_data %>%
  select(-protocol, -day)

orig <- count(hpc_data0, class, name = "orig")
orig

up_rec <- recipe(class ~ ., data = hpc_data0) %>%
  step_tomek(class) %>%
  prep()

training <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "training")
training

# Since `skip` defaults to TRUE, baking the step has no effect
baked <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "baked")
baked

orig %>%
  left_join(training, by = "class") %>%
  left_join(baked, by = "class")

library(ggplot2)

ggplot(circle_example, aes(x, y, color = class)) +
  geom_point() +
  labs(title = "Without Tomek") +
  xlim(c(1, 15)) +
  ylim(c(1, 15))

recipe(class ~ x + y, data = circle_example) %>%
  step_tomek(class) %>%
  prep() %>%
  bake(new_data = NULL) %>%
  ggplot(aes(x, y, color = class)) +
  geom_point() +
  labs(title = "With Tomek") +
  xlim(c(1, 15)) +
  ylim(c(1, 15))
```

step_upsample

geom_point() +
labs(title = "With Tomek") +
xlim(c(1, 15)) +
ylim(c(1, 15))

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 = deprecated(),
  role = NA,
  trained = FALSE,
  column = NULL,
  target = NA,
  skip = TRUE,
  seed = sample.int(10^5, 1),
  id = rand_id("upsample")
)

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 selections() for more details. The selection should result in single factor variable. 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.
<table>
<thead>
<tr>
<th>target</th>
<th>An integer that will be used to subsample. This should not be set by the user and will be populated by prep.</th>
</tr>
</thead>
<tbody>
<tr>
<td>skip</td>
<td>A logical. Should the step be skipped when the recipe is baked by <code>bake()</code>? While all operations are baked when <code>prep()</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.</td>
</tr>
<tr>
<td>seed</td>
<td>An integer that will be used as the seed when upsampling.</td>
</tr>
<tr>
<td>id</td>
<td>A character string that is unique to this step to identify it.</td>
</tr>
</tbody>
</table>

**Details**

Up-sampling is intended to be performed on the training set alone. For this reason, the default is `skip = TRUE`.

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()`.

**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.

**Tidying**

When you `tidy()` this step, a tibble with columns `terms` (the selectors or variables selected) will be returned.

**Case weights**

This step performs an unsupervised operation that can utilize case weights. To use them, see the documentation in `recipes::case_weights` and the examples on tidymodels.org.

**See Also**

Other Steps for over-sampling: `step_adasyn()`, `step_bsmote()`, `step_rose()`, `step_smotenc()`, `step_smote()`

**Examples**

```r
library(recipes)
library(modeldata)
data(hpc_data)

hpc_data0 <- hpc_data %>%
```
tidy.step_adasyn

select(-protocol, -day)

orig <- count(hpc.data0, class, name = "orig")
orig

up_rec <- recipe(class ~ ., data = hpc.data0) %>%
  # Bring the minority levels up to about 1000 each
  # 1000/2211 is approx 0.4523
  step_upsample(class, over_ratio = 0.4523) %>%
  prep()

training <- up_rec %>%
  bake(new_data = NULL) %>%
  count(class, name = "training")
training

# Since 'skip' defaults to TRUE, baking the step has no effect
baked <- up_rec %>%
  bake(new_data = hpc.data0) %>%
  count(class, name = "baked")
baked

# Note that if the original data contained more rows than the
# target n (= ratio * majority_n), the data are left alone:
orig %>%
  left_join(training, by = "class") %>%
  left_join(baked, by = "class")

library(ggplot2)

ggplot(circle.example, aes(x, y, color = class)) +
  geom_point() +
  labs(title = "Without upsample")

recipe(class ~ x + y, data = circle.example) %>%
  step_upsample(class) %>%
  prep() %>%
  bake(new_data = NULL) %>%
  ggplot(aes(x, y, color = class)) +
  geom_jitter(width = 0.1, height = 0.1) +
  labs(title = "With upsample (with jittering)")

---

**tidy.step_adasyn**  
* 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). See `recipes::tidy.recipe` for more information.
Usage

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

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

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

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

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

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

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

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

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

Arguments

- `x` A `step_upsample` object.
- `...` Not currently used.

---

tomek | Remove Tomek's links

Description

Removed observations that are part of tomek links.

Usage

tomek(df, var)
Arguments
df data.frame or tibble. Must have 1 factor variable and remaining numeric variables.
var Character, name of variable containing factor variable.

Details
All columns used in this function must be numeric with no missing data.

Value
A data.frame or tibble, depending on type of df.

References

See Also
step_tomek() for step function of this method
Other Direct Implementations: adasyn(), bsmote(), nearmiss(), smotenc(), smote()

Examples
circle_numeric <- circle_example[, c("x", "y", "class")]

res <- tomek(circle_numeric, var = "class")
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