Package 'sambia'

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Type Package

Title A Collection of Techniques Correcting for Sample Selection Bias

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Description A collection of various techniques correcting statistical models for sample selection bias is provided. In particular, the resampling-based methods ``stochastic inverse-probability oversampling" and ``parametric inverse-probability bagging" are placed at the disposal which generate synthetic observations for correcting classifiers for biased samples resulting from stratified random sampling. For further information, see the article Krauten-bacher, Theis, and Fuchs (2017) <doi:10.1155/2017/7847531>. The methods may be used for further purposes where weighting and generation of new observations is needed.

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R topics documented:

costing	 	 	 	 	2
genSample	 	 	 	 4	ŧ.
IPbag	 	 	 	 6	5
ipOversampling	 	 	 	 8	3
rejSamp	 	 	 	 9)
smoteMod	 	 	 	 10)
smoteNew	 	 	 	 	2
synthIPbag	 	 	 	 13	3

16

Index

costing

Description

This method trains classifiers by correcting them for sample selection bias via Costing, a method proposed in Zadrozny et al. (2003). This method is similar to sambia's IP bagging function in terms of resampling from the learning data and aggregation of the learned algorithms. It differs in the implementation of resampling. Observations from the original data enters a resampled data set only once at most.

Usage

```
costing(..., learner, list.train.learner, list.predict.learner, n.bs,
mod = FALSE)
```

Arguments

	see the parameter rejSamp() of package sambia.		
learner	a character indicating which classifier is used to train. Note: set learner='rangerTree' if random forest should be applied as in Krautenbacher et al. (2017), i.e. the correction step is incorporated in the inherent random forest resampling procedure.		
list.train.lear	mer		
	a list of parameters specific to the classifier that will be trained. Note that the parameter 'data' need not to be provided in this list since the training data which the model will learn on is already attained by new sampled data produced by the method rejSamp().		
list.predict.learner			
	a list of parameters specifiying how to predict new data given the trained model. (This trained model is uniquely determined by parameters 'learner' and 'list.train.learner'		
n.bs	number of bootstramp samples.		
mod	If mod = TRUE: strategy for always having (at least) two outcome classes in subsets.		

Author(s)

Norbert Krautenbacher, Kevin Strauss, Maximilian Mandl, Christiane Fuchs

References

Zadrozny, B., Langford, J., & Abe, N. (2003, November). Cost-sensitive learning by cost-proportionate example weighting. In Data Mining, 2003. ICDM 2003. Third IEEE International Conference on (pp. 435-442). IEEE.

Krautenbacher, N., Theis, F. J., & Fuchs, C. (2017). Correcting Classifiers for Sample Selection Bias in Two-Phase Case-Control Studies. Computational and mathematical methods in medicine, 2017.

costing

```
## simulate data for a population
require(pROC)
set.seed(1342334)
N = 100000
x1 <- rnorm(N, mean=0, sd=1)</pre>
x^{2} <- rt(N, df=25)
x3 <- x1 + rnorm(N, mean=0, sd=.6)
x4 <- x2 + rnorm(N, mean=0, sd=1.3)
x5 <- rbinom(N, 1, prob=.6)
x6 <- rnorm(N, 0, sd = 1) # noise not known as variable
x7 <- x1*x5 # interaction
x <- cbind(x1, x2, x3, x4, x5, x6, x7)
## stratum variable (covariate)
xs <- c(rep(1,0.1*N), rep(0,(1-0.1)*N))</pre>
## effects
beta <- c(-1, 0.2, 0.4, 0.4, 0.5, 0.5, 0.6)
beta0 <- -2
## generate binary outcome
linpred.slopes <- log(0.5)*xs + c(x %*% beta)</pre>
eta <- beta0 + linpred.slopes</pre>
p <- 1/(1+exp(-eta)) \# this is the probability P(Y=1|X), we want the binary outcome however:
y<-rbinom(n=N, size=1, prob=p) #</pre>
population <- data.frame(y,xs,x)</pre>
#### draw "given" data set for training
sel.prob <- rep(1,N)</pre>
sel.prob[population$xs == 1] <- 9</pre>
sel.prob[population$y == 1] <- 8</pre>
sel.prob[population$y == 1 & population$xs == 1] <- 150</pre>
ind <- sample(1:N, 200, prob = sel.prob)</pre>
data = population[ind, ]
## calculate weights from original numbers for xs and y
w.matrix <- table(population$y, population$xs)/table(data$y, data$xs)</pre>
w <- rep(NA, nrow(data))</pre>
w[data$y==0 & data$xs ==0] <- w.matrix[1,1]</pre>
w[data$y==1 & data$xs ==0] <- w.matrix[2,1]</pre>
w[data$y==0 & data$xs ==1] <- w.matrix[1,2]</pre>
w[data$y==1 & data$xs ==1] <- w.matrix[2,2]</pre>
### draw a test data set
newdata = population[sample(1:N, size=200 ), ]
n.bs = 20
```

```
pred_nb <- sambia::costing(data = data, weights = w,
    learner='naiveBayes', list.train.learner = list(formula=formula(y~.)),
    list.predict.learner = list(newdata=newdata, type="raw"),n.bs = n.bs, mod=TRUE)
roc(newdata$y, pred_nb, direction="<")</pre>
```

```
genSample
```

Generate synthetic observations using inverse-probability weights

Description

This method corrects a given data set for sample selection bias by generating new observations via Stochastic inverse-probability oversampling or parametric inverse-probability sampling using inverse-probability weights and information on covariance structure of the given strata (Krautenbacher et al, 2017).

Usage

```
genSample(data, strata.variables = NULL, stratum = NULL, weights = rep(1,
nrow(data)), distr = "mvnorm", type = c("parIP", "stochIP"))
```

Arguments

data	a data frame containing the observations rowwise, along with their correspond- ing categorical strata feature.
strata.variable	25
	a character vector of the names determined by the categorical stratum features.
stratum	a numerical vector of the length of the number of rows of the data specifying the stratum ID. Either 'strata.variables' or 'stratum' has to be provided. This vector will not be included as a column in the resulting data set.
weights	a numerical vector whose length must coincide with the number of the rows of data. The i-th value contains the inverse-probability e.g. determines how often the i-th observation of data shall be replicated.
distr	character object that describes the distribution
type	character which decides which method is used to correct a given data set for sample selection bias. Stochastic Inverse-Probability oversampling is applied if type = 'stochIP' or Parametric Inverse-Probability Bagging if type = 'parIP'.

Value

\$data data frame containing synthetic data which is corrected for sample selection bias by generating new observations via Stochastic inverse-probability oversampling or parametric inverseprobability oversampling.

\$orig.data original data frame which shall to corrected

\$stratum vector containing the stratum of each observation

genSample

\$method a character indicating which method was used. If method = 'stochIP' then Stochastic Inverse-Probability oversampling was used, if method = 'parIP' the Parametric Inverse-Probability sampling was used.

\$strata.tbl a data frame containing all variables and their feature occurences

\$N number of rows in data

\$n number of rows in original data

Author(s)

Norbert Krautenbacher, Kevin Strauss, Maximilian Mandl, Christiane Fuchs

References

Krautenbacher, N., Theis, F. J., & Fuchs, C. (2017). Correcting Classifiers for Sample Selection Bias in Two-Phase Case-Control Studies. Computational and mathematical methods in medicine, 2017.

```
## simulate data for a population
require(pROC)
set.seed(1342334)
N = 100000
x1 <- rnorm(N, mean=0, sd=1)</pre>
x^{2} < -rt(N, df=25)
x3 <- x1 + rnorm(N, mean=0, sd=.6)
x4 <- x2 + rnorm(N, mean=0, sd=1.3)
x5 <- rbinom(N, 1, prob=.6)</pre>
x6 <- rnorm(N, 0, sd = 1) # noise not known as variable
x7 <- x1*x5 # interaction
x <- cbind(x1, x2, x3, x4, x5, x6, x7)
## stratum variable (covariate)
xs <- c(rep(1,0.1*N), rep(0,(1-0.1)*N))</pre>
## effects
beta <- c(-1, 0.2, 0.4, 0.4, 0.5, 0.5, 0.6)
beta0 <- -2
## generate binary outcome
linpred.slopes <- log(0.5)*xs + c(x %*% beta)</pre>
eta <- beta0 + linpred.slopes</pre>
p <- 1/(1+exp(-eta)) # this is the probability P(Y=1|X), we want the binary outcome however:
y<-rbinom(n=N, size=1, prob=p) #</pre>
population <- data.frame(y,xs,x)</pre>
#### draw "given" data set
sel.prob <- rep(1,N)</pre>
```

```
sel.prob[population$xs == 1] <- 9</pre>
sel.prob[population$y == 1] <- 8</pre>
sel.prob[population$y == 1 & population$xs == 1] <- 150</pre>
ind <- sample(1:N, 200, prob = sel.prob)</pre>
data = population[ind, ]
## calculate weights from original numbers for xs and y
w.matrix <- table(population$y, population$xs)/table(data$y, data$xs)</pre>
w <- rep(NA, nrow(data))</pre>
w[data$y==0 & data$xs ==0] <- w.matrix[1,1]</pre>
w[data$y==1 & data$xs ==0] <- w.matrix[2,1]</pre>
w[data$y==0 & data$xs ==1] <- w.matrix[1,2]</pre>
w[data$y==1 & data$xs ==1] <- w.matrix[2,2]</pre>
## parametric IP bootstrap sample
sample1 <- sambia::genSample(data=data, strata.variables = c('y', 'xs'),</pre>
                            weights = w, type='parIP')
## stochastic IP oversampling; treating 'y' and 'xs' as usual input variable
## but using strata info unambiguously defined by the weights w
sample2 <- sambia::genSample(data=data,</pre>
                              weights = w, type='stochIP', stratum= round(w))
```

IPbag

Predicting outcomes using non-parametric Inverse-Probability bagging

Description

This method trains classifiers by correcting them for sample selection bias via non-parametric inverse-probability bagging. This method fits classifiers from different resampled data whose observations are increased per stratum to correct for the bias in the original sample. The so attained ensemble of predictors is aggregated by averaging.

Usage

IPbag(..., learner, list.train.learner, list.predict.learner, n.bs)

Arguments

	see the parameter ipOversampling() of package sambia.	
learner	a character indicating which classifier is used to train. Note: set learner='rangerTree' if random forest should be applied as in Krautenbacher et al. (2017), i.e. the correction step is incorporated in the inherent random forest resampling procedure.	
list.train.learner		
	a list of parameters specific to the classifier that will be trained. Note that the parameter 'data' need not to be provided in this list since the training data which the model will learn on is already attained by new sampled data produced by the method genSample().	

IPbag

list.predict.le	arner
	a list of parameters specifiying how to predict new data given the learned model. (This learned model is uniquely determined by parameters 'learner' and 'list.train.learner').
n.bs	number of bootstramp samples.

Author(s)

Norbert Krautenbacher, Kevin Strauss, Maximilian Mandl, Christiane Fuchs

References

Krautenbacher, N., Theis, F. J., & Fuchs, C. (2017). Correcting Classifiers for Sample Selection Bias in Two-Phase Case-Control Studies. Computational and mathematical methods in medicine, 2017.

```
## simulate data for a population
require(pROC)
set.seed(1342334)
N = 100000
x1 <- rnorm(N, mean=0, sd=1)</pre>
x2 <- rt(N, df=25)
x3 <- x1 + rnorm(N, mean=0, sd=.6)
x4 <- x2 + rnorm(N, mean=0, sd=1.3)
x5 <- rbinom(N, 1, prob=.6)</pre>
x6 <- rnorm(N, 0, sd = 1) # noise not known as variable
x7 <- x1*x5 # interaction
x <- cbind(x1, x2, x3, x4, x5, x6, x7)
## stratum variable (covariate)
xs <- c(rep(1,0.1*N), rep(0,(1-0.1)*N))</pre>
## effects
beta <- c(-1, 0.2, 0.4, 0.4, 0.5, 0.5, 0.6)
beta0 <- -2
## generate binary outcome
linpred.slopes <- log(0.5)*xs + c(x \%) beta)
eta <- beta0 + linpred.slopes</pre>
p <- 1/(1+exp(-eta)) # this is the probability P(Y=1|X), we want the binary outcome however:
y<-rbinom(n=N, size=1, prob=p) #</pre>
population <- data.frame(y,xs,x)</pre>
#### draw "given" data set for training
sel.prob <- rep(1,N)</pre>
sel.prob[population$xs == 1] <- 9</pre>
sel.prob[population$y == 1] <- 8</pre>
sel.prob[population$y == 1 & population$xs == 1] <- 150</pre>
```

```
ind <- sample(1:N, 200, prob = sel.prob)</pre>
data = population[ind, ]
## calculate weights from original numbers for xs and y
w.matrix <- table(population$y, population$xs)/table(data$y, data$xs)</pre>
w <- rep(NA, nrow(data))</pre>
w[data$y==0 & data$xs ==0] <- w.matrix[1,1]</pre>
w[data$y==1 & data$xs ==0] <- w.matrix[2,1]</pre>
w[data$y==0 & data$xs ==1] <- w.matrix[1,2]</pre>
w[data$y==1 & data$xs ==1] <- w.matrix[2,2]</pre>
### draw a test data set
newdata = population[sample(1:N, size=200 ), ]
n.bs = 5
pred_nb <- sambia::IPbag(data = data, weights = w,</pre>
            learner='naiveBayes', list.train.learner = list(formula=formula(y~.)),
            list.predict.learner = list(newdata=newdata, type="raw"),
            n.bs = n.bs)
roc(newdata$y, pred_nb, direction="<")</pre>
```

ipOversampling Plain replication of each observation by inverse-probability weights

Description

This method corrects for the sample selection bias by the plain replication of each observation in the sample according to its IP weight, i.e. in a stratified random sample one replicates an observation of stratum h by the factor w_h.

Usage

```
ipOversampling(data, weights, normalize = FALSE)
```

Arguments

data	a data frame containing the observations rowwise, along with their correspond- ing categorical strata feature(s).
weights	a numerical vector whose length must coincide with the number of the rows of data. The i-th value contains the inverse-probability e.g. determines how often the i-th observation of data shall be replicated.
normalize	If weight vector should be normalized, i.e. the smallest entry of the vector will be set to 1.

Details

If the numeric vector contains numbers which are not natural but real, they will be rounded.

rejSamp

Author(s)

Norbert Krautenbacher, Kevin Strauss, Maximilian Mandl, Christiane Fuchs

Examples

```
library(smotefamily)
library(sambia)
data.example <- sample_generator(100,ratio = 0.80)
result <- gsub('n','0',data.example[,'result'])
result <- gsub('p','1',result)
data.example[,'result'] <- as.numeric(result)
weights <- data.example[,'result']
weights <- ifelse(weights==1,1,4)
sample <- sambia::ipOversampling(data.example,weights)</pre>
```

rejSamp

Rejection Sampling is a method used in sambia's function 'costing' (*Krautenbacher et al, 2017*).

Description

Rejection Sampling is a method used in sambias costing function. It is sampling scheme that allows us to draw examples independently from a distribution X, given examples drawn independently from distribution Y.

Usage

rejSamp(data, weights)

Arguments

data	a data frame containing the observations rowwise, along with their correspond- ing categorical strata feature
weights	a numerical vector whose length must coincide with the number of the rows of data. The i-th value contains the inverse-probability e.g. determines how often the i-th observation of data shall be replicated.

Author(s)

Norbert Krautenbacher, Kevin Strauss, Maximilian Mandl, Christiane Fuchs

References

Krautenbacher, N., Theis, F. J., & Fuchs, C. (2017). Correcting Classifiers for Sample Selection Bias in Two-Phase Case-Control Studies. Computational and mathematical methods in medicine, 2017.

Examples

```
library(smotefamily)
library(sambia)
data.example <- sample_generator(100,ratio = 0.80)
result <- gsub('n','0',data.example[,'result'])
result <- gsub('p','1',result)
data.example[,'result'] <- as.numeric(result)
weights <- data.example[,'result']
weights <- ifelse(weights==1,1,4)
rej.sample <- sambia:::rejSamp(data=data.example, weights = weights)</pre>
```

smoteMod

smoteMod is a modified version of the 'synthetic minority oversampling technique to generate new data.

Description

This method adapts SMOTE to the context of stratified random samples. Rather than enlarging only the minority class, smoteMod generates synthetic data for all strata with a weight bigger than 1. Note: this function has to apply SMOTE H-1 times: 1. subsample data by smallest stratum and a stratum to oversample 2. oversample with modified SMOTE function according to weight of the stratum 3. do this for the other H-2 to subsamples 4. build new data set with strata where H-1 strata contain synthetic data (stratum with smallest weight remains as is)

Usage

smoteMod(data.x, stratum, weights, data.y = NULL, K)

Arguments

data.x	A data frame or matrix of numeric-attributed dataset
stratum	a numerical vector of the same length as the number of the rows of data. De- pending on the number of strata variables and their number of exposures each such combination is assigned to a numeric class id. The i-th entry of stratum contains the class id (and therefore class belonging) of the i-th row (=observa- tion) of data.
weights	a numerical vector whose length must coincide with the number of the rows of data. The i-th value contains the inverse-probability e.g. determines how often the i-th observation of data shall be replicated.
data.y	A vector of a target class attribute corresponding to a dataset data.x.
К	The number of nearest neighbors during sampling process

Author(s)

Norbert Krautenbacher, Kevin Strauss, Maximilian Mandl, Christiane Fuchs

smoteMod

```
## simulate data for a population
require(pROC)
set.seed(1342334)
N = 100000
x1 <- rnorm(N, mean=0, sd=1)</pre>
x^{2} <- rt(N, df=25)
x3 <- x1 + rnorm(N, mean=0, sd=.6)
x4 <- x2 + rnorm(N, mean=0, sd=1.3)
x5 <- rbinom(N, 1, prob=.6)</pre>
x6 <- rnorm(N, 0, sd = 1) # noise not known as variable
x7 <- x1*x5 # interaction
x <- cbind(x1, x2, x3, x4, x5, x6, x7)
## stratum variable (covariate)
xs <- c(rep(1,0.1*N), rep(0,(1-0.1)*N))</pre>
## effects
beta <- c(-1, 0.2, 0.4, 0.4, 0.5, 0.5, 0.6)
beta0 <- -2
## generate binary outcome
linpred.slopes <- log(0.5)*xs + c(x %*% beta)</pre>
eta <- beta0 + linpred.slopes</pre>
p <- 1/(1+exp(-eta)) # this is the probability P(Y=1|X), we want the binary outcome however:
y<-rbinom(n=N, size=1, prob=p) #</pre>
population <- data.frame(y,xs,x)</pre>
#### draw "given" data set for training
sel.prob <- rep(1,N)</pre>
sel.prob[population$xs == 1] <- 9</pre>
sel.prob[population$y == 1] <- 8</pre>
sel.prob[population$y == 1 & population$xs == 1] <- 150</pre>
ind <- sample(1:N, 200, prob = sel.prob)</pre>
data = population[ind, ]
## calculate weights from original numbers for xs and y
w.matrix <- table(population$y, population$xs)/table(data$y, data$xs)</pre>
w <- rep(NA, nrow(data))</pre>
w[data$y==0 & data$xs ==0] <- w.matrix[1,1]</pre>
w[data$y==1 & data$xs ==0] <- w.matrix[2,1]</pre>
w[data$y==0 & data$xs ==1] <- w.matrix[1,2]</pre>
w[data$y==1 & data$xs ==1] <- w.matrix[2,2]</pre>
### draw a test data set
newdata = population[sample(1:N, size=200 ), ]
K = 5
```

```
genData = smoteMod(data.x = data[ , -which(colnames(data) %in% c('y', 'xs'))] ,
stratum = w, data.y = data$y, weights = w, K=K)
```

smoteNew

smoteNew is a necessary function that modifies the SMOTE algorithm.

Description

smoteNewis a necessary function that modifies the SMOTE algorithm in the following ways: (1) correct bug in original smotefamily::SMOTE() function and (2) lets the user specifiy which class to be oversampled.

Usage

smoteNew(data.x, data.y, K, dup_size = 0, class.to.oversample)

Arguments

data.x	A data frame or matrix of numeric-attributed dataset	
data.y	A vector of a target class attribute corresponding to a dataset X	
К	The number of nearest neighbors during sampling process	
dup_size	The number or vector representing the desired times of synthetic minority in- stances over the original number of majority instances	
class.to.oversample		
	Class to be oversampled	

Author(s)

Norbert Krautenbacher, Kevin Strauss, Maximilian Mandl, Christiane Fuchs

Examples

```
library(smotefamily)
library(sambia)
data.example = sample_generator(10000,ratio = 0.80)
genData = sambia:::smoteNew(data.example[,-3],data.example[,3],K = 5,class.to.oversample = 'p')
```

synthIPbag

Train a classifier via synthetic observations using inverse-probability weights

Description

This method trains classifiers by correcting them for sample selection bias via stochastic inverseprobability oversampling or parametric inverse-probability bagging (Krautenbacher et al 2017). Classifiers are trained from differently resampled data whose observations are weighted by inverseprobability weights per stratum to correct for the bias in the original sample. The so attained ensemble of predictors is aggregated by averaging.

Usage

synthIPbag(..., learner, list.train.learner, list.predict.learner, n.bs)

Arguments

	see the parameter genSample() of package sambia.		
learner	a character indicating which classifier is used to train. Note: set learner='rangerTree' if random forest should be applied as in Krautenbacher et al. (2017), i.e. the correction step is incorporated in the inherent random forest resampling procedure.		
list.train.learner			
	a list of parameters specific to the classifier that will be trained. Note that the parameter 'data' need not to be provided in this list since the training data which the model will learn on is already attained by new sampled data produced by the method genSample().		
list.predict.learner			
	a list of parameters specifiying how to predict new data given the trained model.		
n.bs	number of bootstramp samples. This trained model is uniquely determined by parameters 'learner' and 'list.train.learner'.		

Author(s)

Norbert Krautenbacher, Kevin Strauss, Maximilian Mandl, Christiane Fuchs

References

Krautenbacher, N., Theis, F. J., & Fuchs, C. (2017). Correcting Classifiers for Sample Selection Bias in Two-Phase Case-Control Studies. Computational and mathematical methods in medicine, 2017.

Krautenbacher, N., Theis, F. J., & Fuchs, C. (2017). Correcting Classifiers for Sample Selection Bias in Two-Phase Case-Control Studies. Computational and mathematical methods in medicine, 2017.

Examples

```
## simulate data for a population
require(pROC)
set.seed(1342334)
N = 100000
x1 <- rnorm(N, mean=0, sd=1)</pre>
x^{2} <- rt(N, df=25)
x3 <- x1 + rnorm(N, mean=0, sd=.6)
x4 <- x2 + rnorm(N, mean=0, sd=1.3)
x5 <- rbinom(N, 1, prob=.6)</pre>
x6 <- rnorm(N, 0, sd = 1) # noise not known as variable
x7 <- x1*x5 # interaction
x <- cbind(x1, x2, x3, x4, x5, x6, x7)
## stratum variable (covariate)
xs <- c(rep(1,0.1*N), rep(0,(1-0.1)*N))</pre>
## effects
beta <- c(-1, 0.2, 0.4, 0.4, 0.5, 0.5, 0.6)
beta0 <- -2
## generate binary outcome
linpred.slopes <- log(0.5)*xs + c(x %*% beta)</pre>
eta <- beta0 + linpred.slopes</pre>
p <- 1/(1+exp(-eta)) # this is the probability P(Y=1|X), we want the binary outcome however:
y<-rbinom(n=N, size=1, prob=p) #</pre>
population <- data.frame(y,xs,x)</pre>
#### draw "given" data set for training
sel.prob <- rep(1,N)</pre>
sel.prob[population$xs == 1] <- 9</pre>
sel.prob[population$y == 1] <- 8</pre>
sel.prob[population$y == 1 & population$xs == 1] <- 150</pre>
ind <- sample(1:N, 200, prob = sel.prob)</pre>
data = population[ind, ]
## calculate weights from original numbers for xs and y
w.matrix <- table(population$y, population$xs)/table(data$y, data$xs)</pre>
w <- rep(NA, nrow(data))</pre>
w[data$y==0 & data$xs ==0] <- w.matrix[1,1]</pre>
w[data$y==1 & data$xs ==0] <- w.matrix[2,1]</pre>
w[data$y==0 & data$xs ==1] <- w.matrix[1,2]</pre>
w[data$y==1 & data$xs ==1] <- w.matrix[2,2]</pre>
### draw a test data set
newdata = population[sample(1:N, size=200 ), ]
n.bs = 10
```

Index

 $\operatorname{costing}, 2$

genSample,4

IPbag,6 ipOversampling,8

rejSamp,<mark>9</mark>

smoteMod, 10
smoteNew, 12
synthIPbag, 13