

Package ‘ganGenerativeData’

December 12, 2024

Type Package

Title Generate Generative Data for a Data Source

Version 2.1.4

Date 2024-12-13

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Description Generative Adversarial Networks are applied to generate generative data for a data source. A generative model consisting of a generator and a discriminator network is trained. During iterative training the distribution of generated data is converging to that of the data source. Direct applications of generative data are the created functions for data evaluation, missing data completion and data classification. A software service for accelerated training of generative models on graphics processing units is available. Reference: Goodfellow et al. (2014) <[doi:10.48550/arXiv.1406.2661](https://doi.org/10.48550/arXiv.1406.2661)>.

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Imports Rcpp (>= 1.0.3), tensorflow (>= 2.0.0), httr (>= 1.4.7)

LinkingTo Rcpp

RoxygenNote 7.2.3

SystemRequirements TensorFlow (<https://www.tensorflow.org>)

NeedsCompilation yes

Encoding UTF-8

Repository CRAN

Date/Publication 2024-12-12 13:20:01 UTC

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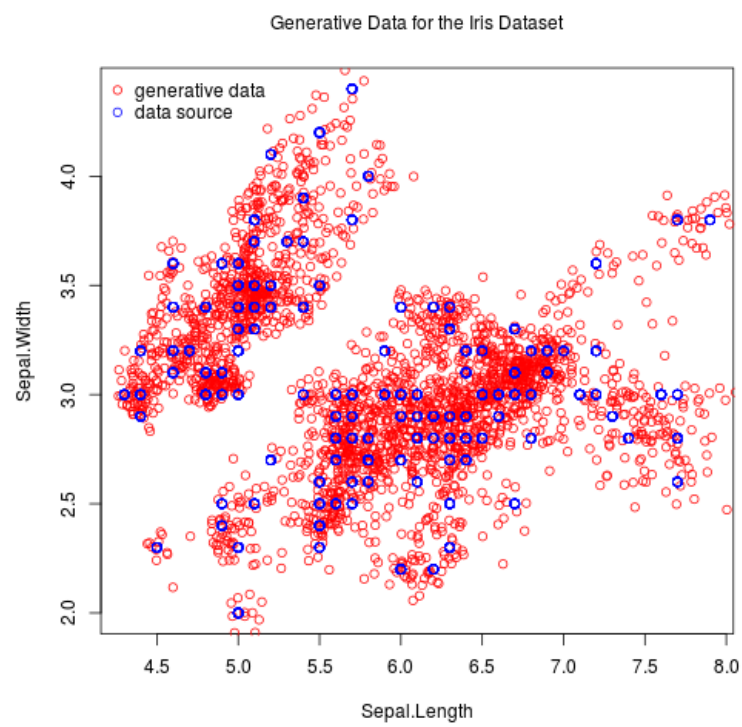
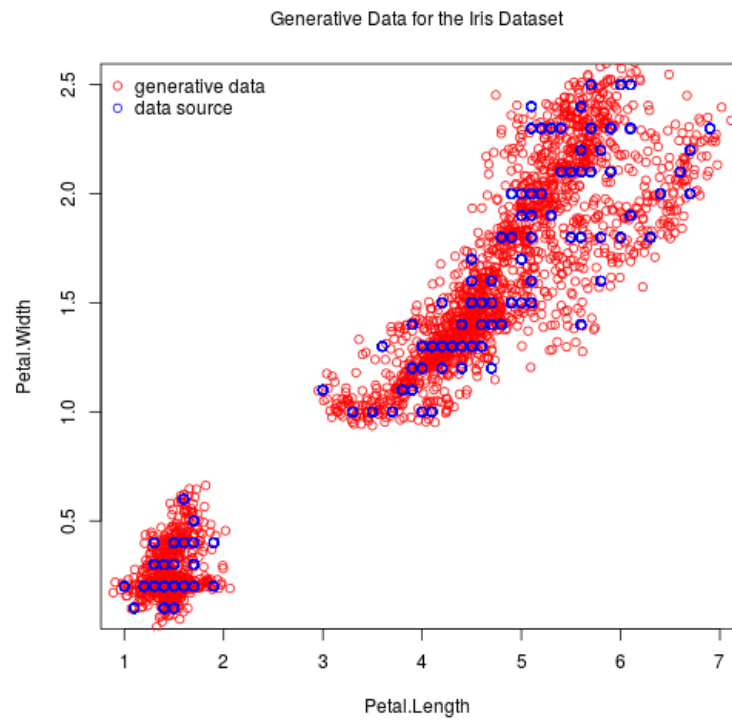
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Index**41****ganGenerativeData-package***Generate generative data for a data source***Description**

Generative Adversarial Networks are applied to generate generative data for a data source. A generative model consisting of a generator and a discriminator network is trained. During iterative training the distribution of generated data is converging to that of the data source.

Generated data can be written to a file in training and after finished training in a separate generation step. First method accumulates generative data using a dynamic model, second method generates generative data using a static model.

Inserted images show two-dimensional projections of generative data for the iris dataset:



Method of data evaluation:

Let $M = (R^D, d)$ be the metric space with Euclidean distance d on R^D . Let X be a random vector consisting of D random variables and $(x_i)_{i=1}^n, x_i \in R^D$ a sequence of occurrences of X .

The k nearest neighbor density estimation function is defined by

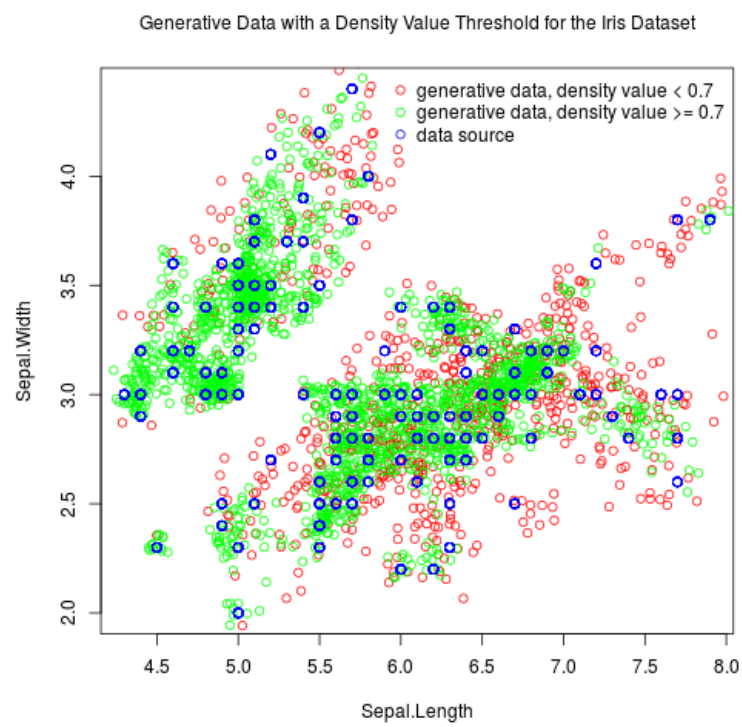
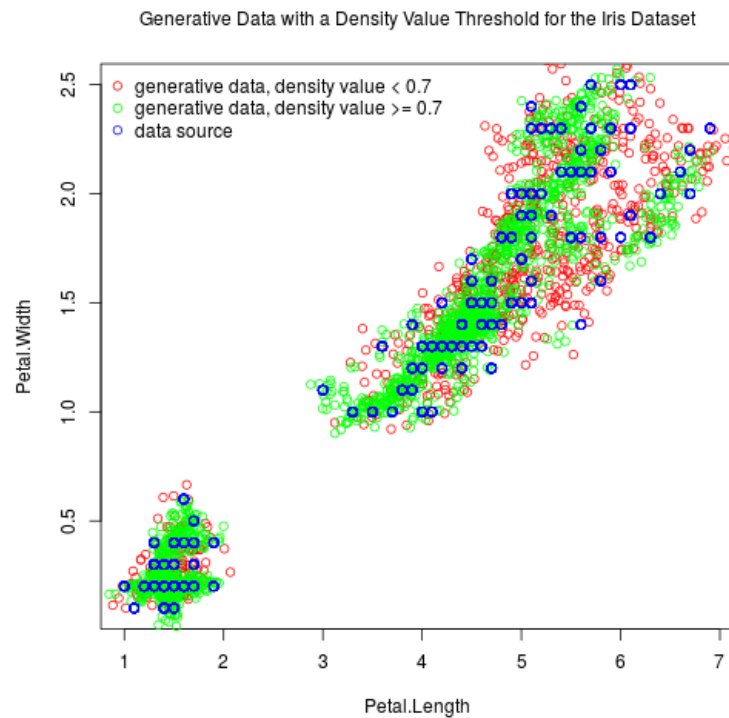
$$de(x) = \frac{k}{nd^D(x_{i_k}, x)V_D}, x \in R^D$$

where k is the number of nearest neighbors, n is the number of occurrences, x_{i_k} is the k -th nearest neighbor to x , $d(x_{i_k}, x)$ is the Euclidean distance, $V_D = \frac{\pi^{D/2}}{\Gamma(\frac{D}{2}+1)}$ is the volume of a unit sphere and $\Gamma(x)$ is the Gamma function.

We calculate density values by logarithmizing and normalizing $de(x)$. Next we set the density value of $x \in R^D$ into relation to density values of a sequence of occurrences of X by computing the inverse quantile. The received percent value is then used for evaluation of x .

In practice function `gdCalulcateDensityValues()` is called to calculate density values for generated generative data and for $x \in R^D$ function `gdCalculateDensityValue()` and `gdDensityValueInverseQuantile()` is called to receive a percent value for evaluation of x .

For illustration inserted images show generative data for a density value threshold for the iris dataset:



Method of missing data completion:

Without loss of generality we consider the case of two random variables X, Y . Let $g_{XY}(x, y)$ be the continuous probability density function of joint distribution X, Y and $G = ((x_i, y_i))_{i=1}^n$ a sequence of occurrences of X, Y .

In order to complete missing data for a given $X = x_0$ we consider the conditional probability density function $g_{Y|X=x_0}(y)$.

We compute the mean of $g_{XY}(x, y)$ over interval $[x_0 - \epsilon, x_0 + \epsilon]$ and then the limit by applying the rule of de l'Hospital

$$\lim_{\epsilon \rightarrow 0} \frac{1}{2\epsilon} \int_{x_0 - \epsilon}^{x_0 + \epsilon} g_{XY}(x, y) dx = g_{XY}(x_0, y)$$

By deviding a normalization term instead of 2ϵ we get

$$\lim_{\epsilon \rightarrow 0} \frac{1}{\int_{x_0 - \epsilon}^{x_0 + \epsilon} \int_{-\infty}^{+\infty} g_{XY}(x, y) dy dx} \int_{x_0 - \epsilon}^{x_0 + \epsilon} g_{XY}(x, y) dx = \frac{\int_{-\infty}^{+\infty} g_{XY}(x_0, y) dy}{\int_{-\infty}^{+\infty} g_{XY}(x_0, y) dy} = g_{Y|X=x_0}(y)$$

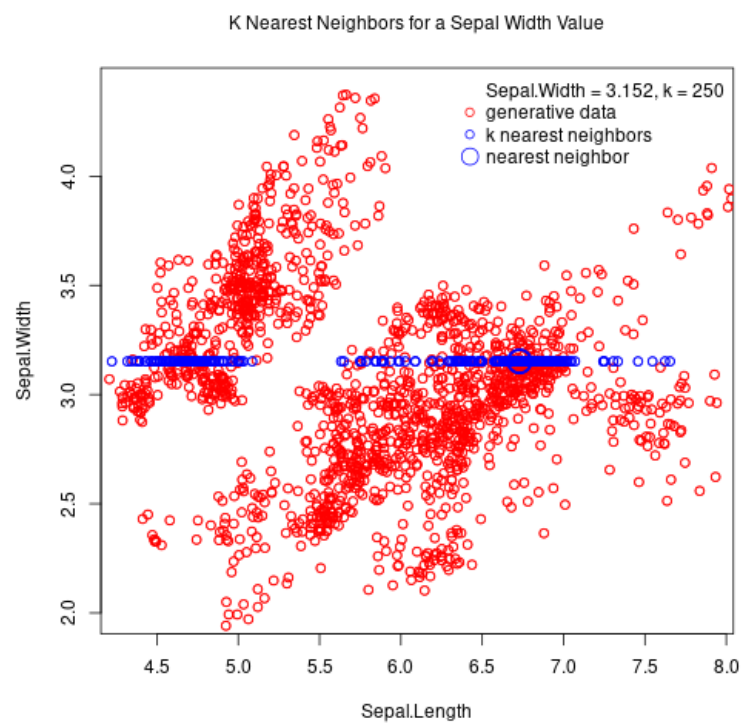
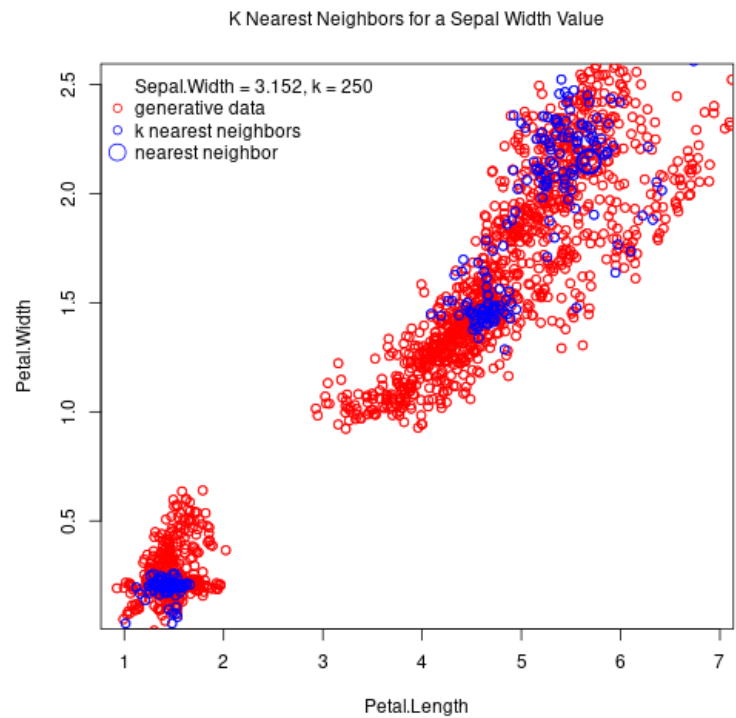
So the integral of $g_{XY}(x, y)$ over interval $[x_0 - \epsilon, x_0 + \epsilon]$ divided by a normalization term converges to $g_{Y|X=x_0}(y)$ for $\epsilon \rightarrow 0$ and the subsequence of occurrences of G in the interval is given by $G_{x_0, \epsilon} = ((x_{i_j}, y_{i_j}))$ with $1 \leq i_j \leq n, x_{i_j} \in [x_0 - \epsilon, x_0 + \epsilon]$.

A candidate for missing $Y = y_0$ for a given $X = x_0$ is then y_k of a uniform randomly selected (x_k, y_k) from $G_{x_0, \epsilon}$.

This is applied for generated generative data as follows:

In function gdComplete() the nearest neighbor for a given incomplete data record is searched with respect to present values and then utilized to complete the incomplete data record. Also function gdKNearestNeighbors() can be called. A nearest neighbor uniform randomly selected from returned k nearest neighbors can then be used for data record completion.

For illustration inserted images show k nearest neighbors for a sepal width value:



Accelerated training of generative models:

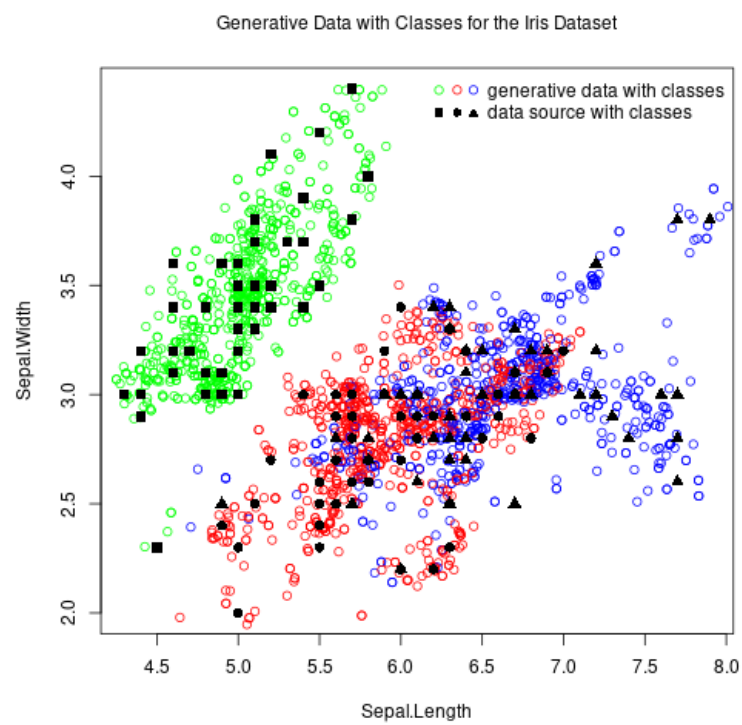
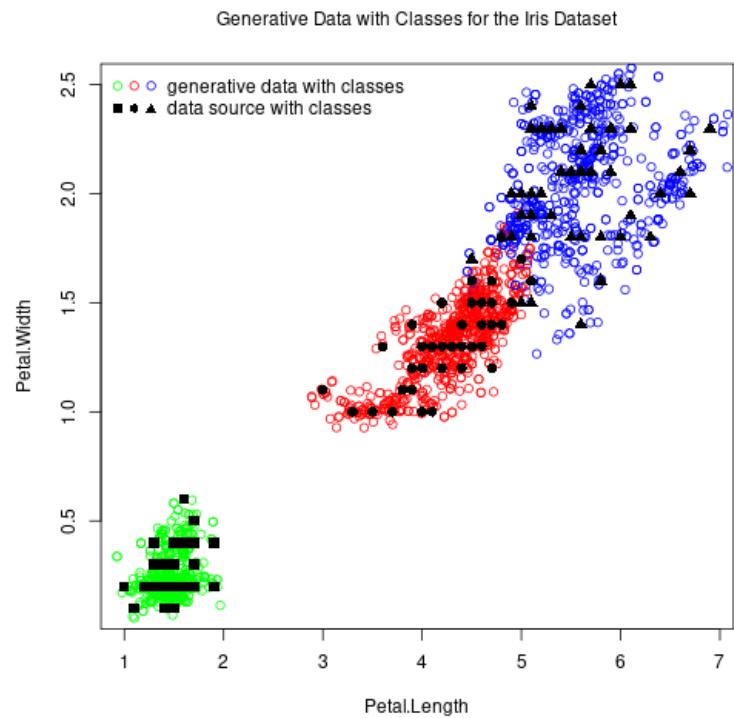
Training of generative models can be accelerated by using graphics processing units for computation. For this kind of training a software service is available. For access details contact the author of the package and see in the documentation in section Examples how training can be executed. The software service also supports training of generative models for mixed numerical and categorical variables.

Method of data classification:

With support of training of generative models for mixed numerical and categorical variables data classification can be done for a given labeled dataset and unlabeled data as follows:

1. Considering class labels as a categorical variable train a generative model for joint distribution of data records and class labels.
2. Apply method of missing data completion to determine class labels for unlabeled data records.

For illustration inserted images show generative data for the iris dataset with data records consisting of numerical variables sepal length, sepal width, petal length, petal width and assigned class labels:



Method of generative model evaluation:

A specific method for generative model evaluation compares the distribution of density values for generated generative data with that of the data source as follows:

1. Create histograms representing distributions of density values for generative data and data source
2. Compute the Kullback-Leibler divergence for created generative data and data source histogram to measure the distribution divergence.

In practice the Kullback-Leibler divergence is calculated by calling functions `gdDensityValueInverseQuantile()` and `dsDensityValueInverseQuantile()` as follows:

$$D_{KL}(h_{gd}||h_{ds}) = \sum_{i=1}^n h_{gd,i} \log\left(\frac{h_{gd,i}}{h_{ds,i}}\right)$$

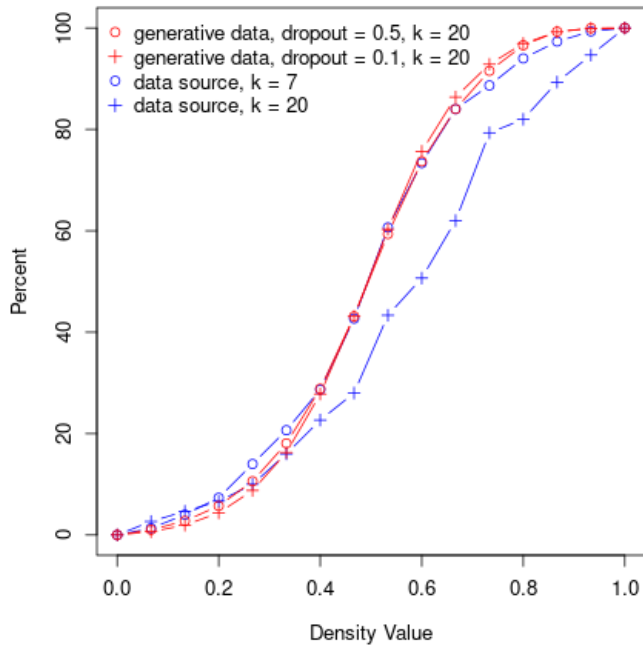
$$h_{gd,i} = (gdDensityValueInverseQuantile(i/n) - gdDensityValueInverseQuantile((i-1)/n)) / 100$$

$$h_{ds,i} = (dsDensityValueInverseQuantile(i/n) - dsDensityValueInverseQuantile((i-1)/n)) / 100$$

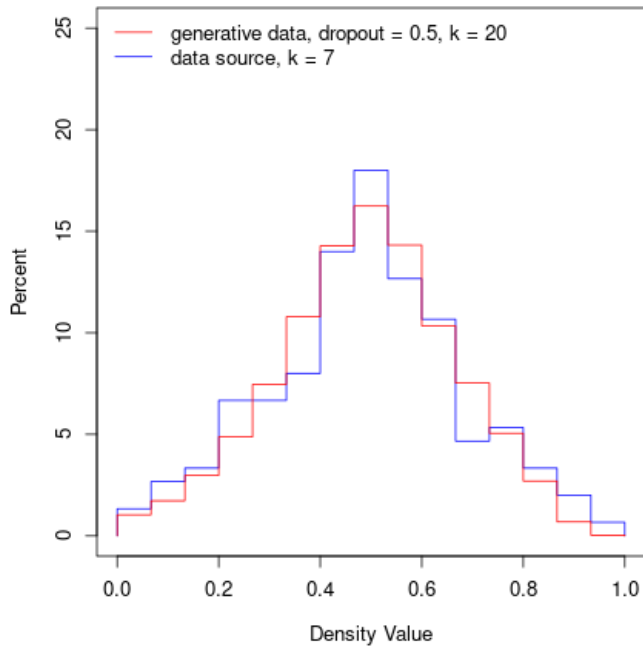
where $h_{gd,i}$ and $h_{ds,i}$ are percent values of density values for generative data and data source lying in interval $((i-1)/n, i/n]$, $1 \leq i \leq n$.

For illustration inserted images show inverse quantiles and histograms for the iris dataset. For the small size iris data source density values are calculated for different k nearest neighbor values.

Density Value Inverse Quantiles for the Iris Dataset



Density Value Histograms for the Iris Dataset



Details

The API includes functions for topics "definition of data source" and "generation of generative data". Main function of first topic is `dsCreateWithDataFrame()` which creates a data source with passed data frame. Main functions of second topic are `gdTrain()` which trains a generative model for a data source and `gdGenerate()` which uses a trained generative model to generate generative data. Additionally a software service for accelerated training of generative models is available.

1. Definition of data source

`dsCreateWithDataFrame()` Create a data source with passed data frame.

`dsActivateColumns()` Activate columns in a data source in order to include them in training of generative models. By default columns are active.

`dsDeactivateColumns()` Deactivate columns in a data source in order to exclude them from training of generative models. Note that the training function in the package supports only columns of type R-class numeric, R-type double. All columns of other type have to be deactivated. The training function in the software service for accelerated training of generative models supports columns of any type.

`dsGetActiveColumnNames()` Get names of active columns of a data source.

`dsGetInactiveColumnNames()` Get names of inactive columns of a data source.

`dsWrite()` Write created data source including settings of active columns to a file in binary format. This file will be used as input in functions of topic "generation of generative data".

`dsRead()` Read a data source from a file that was written with `dsWrite()`.

`dsGetNumberOfRows()` Get number of rows in a data source.

`dsgetRow()` Get a row in a data source.

`dsCalculateDensityValues()` Read a data souce from a file, calculate density values and write the data source with density values to original file.

`dsDensityValueInverseQuantile()` Calculate inverse density value quantile for a density value.

2. Training of generative model and generation of generative data

`gdTrainParameters()` Specify parameters for training of generative model.

`gdTrain()` Read a data source from a file, train a generative model that generates generative data for the data source in iterative training steps, write trained generative model and generated data in

training steps to a file in binary format..

`gdGenerateParameters()` Specify parameters for generation of generative data.

`gdGenerate()` Read a generative model and a data source from a file, generate generative data for the data source and write generated data to a file in binary format.

`gdCalculateDensityValues()` Read generative data from a file, calculate density values and write generative data with density values to original file.

`gdRead()` Read generative data and data source from specified files.

`gdPlotParameters()` Specify plot parameters for generative data.

`gdPlotDataSourceParameters()` Specify plot parameters for data source.

`gdPlotProjection()` Create an image file containing two-dimensional projections of generative data and data source.

`gdGetNumberOfRows()` Get number of rows in generative data.

`gdGetRow()` Get a row in generative data.

`gdCalculateDensityValue()` Calculate density value for a data record.

`gdDensityValueQuantile()` Calculate density value quantile for a percent value.

`gdDensityValueInverseQuantile()` Calculate inverse density value quantile for a density value.

`gdKNearestNeighbors()` Search for k nearest neighbors in generative data.

`gdComplete()` Complete incomplete data record.

`gdWriteSubset()` Write subset of generative data.

3. Software service for accelerated training of generative models

`gdServiceTrain()` Send a request to software service to train a generative model.

`gdServiceGetGenerativeData()` Get generated generative data from software service.

`gdServiceGetGenerativeModel()` Get trained generative model from software service.

gdServiceGetStatus() Get status of generated job from software service.

gdServiceDelete() Delete generated job from software service.

Author(s)

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References

Ian J. Goodfellow, Jean Pouget-Abadie, Mehdi Mirza, Bing Xu, David Warde-Farley, Sherjil Ozair, Aaron Courville, Yoshua Bengio (2014), "*Generative Adversarial Nets*", <arXiv:1406.2661v1>

Examples

```
# Environment used for execution of examples:

# Operating system: Ubuntu 22.04.1
# Compiler: g++ 11.3.0 (supports C++17 standard)
# R applications: R 4.1.2, RStudio 2022.02.2
# Installed packages: 'Rcpp' 1.0.10, 'tensorflow' 2.11.0,
# 'ganGenerativeData' 2.1.4

# Package 'tensorflow' provides an interface to machine learning framework
# TensorFlow. To complete the installation function install_tensorflow() has to
# be called.
## Not run:
library(tensorflow)
install_tensorflow()
## End(Not run)

# Generate generative data for the iris dataset

# Load library
library(ganGenerativeData)

# 1. Definition of data source for the iris dataset

# Create a data source with iris data frame.
dsCreateWithDataFrame(iris)

# Deactivate the column with name Species and index 5 in order to exclude it in
# trainng of generative model.
dsDeactivateColumns(c(5))

# Get the active column names: Sepal.Length, Sepal.Width, Petal.Length,
# Petal.Width.
dsGetActiveColumnNames()
```

```
# Write the data source including settings of active columns to file
# "ds.bin" in binary format.
## Not run:
dsWrite("ds.bin")
## End(Not run)

# 2. Training of generative model and generation of generative data for the iris
# data source

# Read data source from file "ds.bin", train a generative model in iterative
# training steps (used number of iterations in tests is in the range of 10000 to
# 50000), write trained generative model and generated data in training steps to
# files "gm.bin" and "gd.bin".
## Not run:
gdTrain("gm.bin", "gd.bin", "ds.bin", c(1, 2),
gdTrainParameters(numberOfTrainingIterations = 1000))
## End(Not run)

# Read generative data from file "gd.bin", calculate density values and
# write generative data with density values to original file.
## Not run:
gdCalculateDensityValues("gd.bin")
## End(Not run)

# Read generative data from file "gd.bin" and data source from "ds.bin". Read in
# data will be accessed in following function calls.
## Not run:
gdRead("gd.bin", "ds.bin")
## End(Not run)

# Create an image showing two-dimensional projections of generative data and
# data source for column indices 3, 4 and write it to file "gd34d.png".
## Not run:
gdPlotProjection("gd34d.png",
"Generative Data for the Iris Dataset",
c(3, 4),
gdPlotParameters(50),
gdPlotDataSourceParameters(100))
## End(Not run)

# Create an image showing two-dimensional projections of generative data and
# data source for column indices 3, 4 with density value threshold 0.71 and
# write it to file "gd34ddv.png".
## Not run:
gdPlotProjection("gd34ddv.png",
"Generative Data with a Density Value Threshold for the Iris Dataset",
c(3, 4),
gdPlotParameters(50, c(0.38, c("red", "green"))),
gdPlotDataSourceParameters(100))
## End(Not run)

# Get number of rows in generative data
## Not run:
```

```

gdGetNumberOfRows()
## End(Not run)

# Get row with index 1000 in generative data
## Not run:
gdGetRow(1000)
## End(Not run)

# Calculate density value for a data record
## Not run:
gdCalculateDensityValue(list(6.1, 2.6, 5.6, 1.4))
## End(Not run)

# Calculate density value quantile for 50 percent
## Not run:
gdDensityValueQuantile(50)
## End(Not run)

# Calculate inverse density value quantile for density value 0.5
## Not run:
gdDensityValueInverseQuantile(0.5)
## End(Not run)

# Search for k nearest neighbors for a data record
## Not run:
gdKNearestNeighbors(list(5.1, 3.5, 1.4, 0.2), 3)
## End(Not run)

# Complete incomplete data record containing an NA value
## Not run:
gdComplete(list(5.1, 3.5, 1.4, NA))
## End(Not run)

# Write subset containing 50 percent of randomly selected rows of
# generative data
## Not run:
gdRead("gd.bin")
gdWriteSubset("gds.bin", 50)
## End(Not run)

# 3. Usage of software service for accelerated training of a generative
# model

# Initialize variables for URL and access key.
## Not run:
url <- "http://xxx.xxx.xxx.xxx/gdService"
accessKey <- "xxxxxxxx-xxxx-xxxx-xxxx-xxxxxxxxxxxx"
## End(Not run)

# Send a request to software service to train a generative model for a data
# source. A job id will be returned.
## Not run:
trainParameters <- gdTrainParameters(numberOfTrainingIterations = 10000,

```

```
numberOfInitializationIterations = 2500)
jobId <- gdServiceTrain(url, accessKey, "gmService.bin", "gdService.bin", "ds.bin",
trainParameters)
## End(Not run)

# Get status of generated job from software service. When job is processed
# successfully status will be set to TRAINED.
## Not run:
gdServiceGetStatus(url, accessKey, jobId)
## End(Not run)

# Get generated generative data from software service for processed job
## Not run:
gdServiceGetGenerativeData(url, accessKey, jobId, "gdService.bin")
## End(Not run)

# Get trained generative model from software service for processed job
## Not run:
gdServiceGetGenerativeModel(url, accessKey, jobId, "gmService.bin")
## End(Not run)
```

dsActivateColumns *Activate columns*

Description

Activate columns in a data source in order to include them in training of generative models. By default columns are active.

Usage

```
dsActivateColumns(columnVector)
```

Arguments

columnVector Vector of column indices

Value

None

Examples

```
dsCreateWithDataFrame(iris)
dsGetActiveColumnNames()
dsDeactivateColumns(c(5))
dsGetActiveColumnNames()
dsActivateColumns(c(5))
dsGetActiveColumnNames()
```

dsCalculateDensityValues

Calculate density values for data source

Description

Read a data source from a file, calculate density values and write the data source with density values to original file. Calculated density values are used to evaluate a data source.

Usage

```
dsCalculateDensityValues(dataSourceFileName, nNearestNeighbors)
```

Arguments

dataSourceFileName	Name of data source file name
nNearestNeighbors	number of used nearest neighbors

Value

None

Examples

```
## Not run:  
dsCalculateDensityValues("ds.bin")  
## End(Not run)
```

dsCreateWithDataFrame *Create a data source with passed data frame*

Description

Create a data source with passed data frame.

Usage

```
dsCreateWithDataFrame(dataFrame)
```

Arguments

dataFrame	Name of data frame
-----------	--------------------

Value

None

Examples

```
dsCreateWithDataFrame(iris)
```

```
dsDeactivateColumns      Deactivate columns
```

Description

Deactivate columns in a data source in order to exclude them from training of generative models. Note that the training function in the package supports only columns of type R-class numeric, R-type double. All columns of other type have to be deactivated. The training function in the software service for accelerated training of generative models supports columns of any type.

Usage

```
dsDeactivateColumns(columnVector)
```

Arguments

columnVector Vector of column indices

Value

None

Examples

```
dsCreateWithDataFrame(iris)
dsDeactivateColumns(c(5))
dsGetInactiveColumnNames()
```

dsDensityValueInverseQuantile
Calculate inverse density value quantile

Description

Calculate inverse density value quantile for a density value.

Usage

```
dsDensityValueInverseQuantile(densityValue)
```

Arguments

densityValue Normalized density value

Value

Percent value

Examples

```
## Not run:  

dsRead("ds.bin")  

dsDensityValueInverseQuantile(0.5)  

## End(Not run)
```

dsGetActiveColumnNames
Get active column names

Description

Get active column names of a data source

Usage

```
dsGetActiveColumnNames()
```

Value

Vector of names of active columns

Examples

```
dsCreateWithDataFrame(iris)  

dsDeactivateColumns(c(5))  

dsGetActiveColumnNames()
```

```
dsGetInactiveColumnNames  
    Get inactive column names
```

Description

Get inactive column names of a data source

Usage

```
dsGetInactiveColumnNames()
```

Value

Vector of names of inactive columns

Examples

```
dsCreateWithDataFrame(iris)  
dsDeactivateColumns(c(5))  
dsGetInactiveColumnNames()
```

```
dsGetNumberOfRows      Get number of rows
```

Description

Get number of rows in a data source

Usage

```
dsGetNumberOfRows()
```

Value

Number of rows

Examples

```
dsCreateWithDataFrame(iris)  
dsGetNumberOfRows()
```

dsGetRow	<i>Get a row in a data source</i>
----------	-----------------------------------

Description

Get a row in a data source for a row index.

Usage

```
dsGetRow(index)
```

Arguments

index	Index of row
-------	--------------

Value

List containing row in data source

Examples

```
dsCreateWithDataFrame(iris)  
dsGetRow(1)
```

dsRead	<i>Read a data source from file</i>
--------	-------------------------------------

Description

Read a data source from a file in binary format

Usage

```
dsRead(fileName)
```

Arguments

fileName	Name of data source file
----------	--------------------------

Value

None

Examples

```
## Not run:  
dsCreateWithDataFrame(iris)  
dsDeactivateColumns(c(5))  
dsWrite("ds.bin")  
dsRead("ds.bin")  
## End(Not run)
```

dsWrite

Write a data source to file

Description

Write a data source including settings of active columns to a file in binary format. This file will be used as input in functions for generation of generative data.

Usage

```
dsWrite(fileName)
```

Arguments

fileName	Name of data source file
----------	--------------------------

Value

None

Examples

```
## Not run:  
dsCreateWithDataFrame(iris)  
dsDeactivateColumns(c(5))  
dsWrite("ds.bin")  
## End(Not run)
```

gdCalculateDensityValue

Calculate density value for a data record

Description

Calculate density value for a data record. By default for the calculation a linear search is performed on generative data. When a search tree is used search is performed on a tree for generative data which is built once in the first function call.

Usage

```
gdCalculateDensityValue(dataRecord, useSearchTree = FALSE)
```

Arguments

dataRecord	List containing a data record
useSearchTree	Boolean value indicating if a search tree should be used.

Value

Normalized density value number

Examples

```
## Not run:
gdRead("gd.bin")
dv <- gdCalculateDensityValue(list(6.1, 2.6, 5.6, 1.4))
## End(Not run)
```

gdCalculateDensityValues

Calculate density values for generative data

Description

Read generative data from a file, calculate density values and write generative data with density values to original file. Calculated density values are used to classiy generative data. In function gdPlotParameters() density value thresholds with assigned colors can be passed to draw generative data for different density value ranges.

Usage

```
gdCalculateDensityValues(generativeDataFileName)
```

Arguments

generativeDataFileName
Name of generative data file name

Value

None

Examples

```
## Not run:  
gdCalculateDensityValues("gd.bin")  
## End(Not run)
```

gdComplete

Complete incomplete data record

Description

Search for first nearest neighbor in normalized generative data for incomplete data record containing NA values. Found row in generative data is then used to replace NA values in incomplete data record. This function calls gdKNearestNeighbors() with parameter k equal to 1.

Usage

```
gdComplete(dataRecord, useSearchTree = FALSE)
```

Arguments

dataRecord List containing incomplete data record
useSearchTree Boolean value indicating if a search tree should be used.

Value

List containing completed data record

Examples

```
## Not run:  
gdRead("gd.bin")  
gdComplete(list(5.1, 3.5, 1.4, NA))  
## End(Not run)
```

gdDensityValueInverseQuantile
Calculate inverse density value quantile

Description

Calculate inverse density value quantile for a density value.

Usage

```
gdDensityValueInverseQuantile(densityValue)
```

Arguments

densityValue Normalized density value

Value

Percent value

Examples

```
## Not run:  

gdRead("gd.bin")  

gdDensityValueInverseQuantile(0.5)  

## End(Not run)
```

gdDensityValueQuantile
Calculate density value quantile

Description

Calculate density value quantile for a percent value.

Usage

```
gdDensityValueQuantile(percent)
```

Arguments

percent Percent value

Value

Normalized density value quantile number

Examples

```
## Not run:  
gdRead("gd.bin")  
gdDensityValueQuantile(50)  
## End(Not run)
```

gdGenerate

Generate generative data for a data source

Description

Read a trained generative model for a data source, generate generative data and write generated data to a file in binary format.

Usage

```
gdGenerate(  
  generativeDataFileName,  
  generativeModelFileName,  
  generateParameters = gdGenerateParameters(numberOfSamples = 10000, dropout = 0.05)  
)
```

Arguments

```
generativeDataFileName  
  Name of generative data file  
generativeModelFileName  
  Name of generative model file  
generateParameters  
  Generation of generative data parameters, see function gdGenerateParameters().
```

Value

None

Examples

```
## Not run:  
generateParameters <- gdGenerateParameters(numberOfSamples = 10000)  
gdGenerate("gd.bin", "gm.bin", generateParameters)  
## End(Not run)
```

gdGenerateParameters *Specify parameters for generation of generative data*

Description

Specify parameters for generation of generative data. These parameters are passed to function gdGenerate().

Usage

```
gdGenerateParameters(numberOfSamples = 1e+05, dropout = 0.05)
```

Arguments

numberOfSamples

Number of generated samples

dropout

Value in the range of 0 to 1. Specifies the rate of hidden units that are dropped. Dropout is a regularization method to prevent overfitting. See function gdTrainParameters().

Value

List of parameters for generation of generative data

Examples

```
## Not run:  
generateParameters <- gdGenerateParameters(numberOfSamples = 100000)  
## End(Not run)
```

gdGetNumberOfRows *Get number of rows*

Description

Get number of rows in generative data

Usage

```
gdGetNumberOfRows()
```

Value

Number of rows

Examples

```
## Not run:  
gdRead("gd.bin")  
gdGetNumberOfRows()  
## End(Not run)
```

gdGetRow

Get a row in generative data

Description

Get a row in generative data for a row index

Usage

```
gdGetRow(index)
```

Arguments

index	Index of row
-------	--------------

Value

List containing row in generative data

Examples

```
## Not run:  
gdRead("gd.bin")  
gdGetRow(1000)  
## End(Not run)
```

gdKNearestNeighbors

Search for k nearest neighbors

Description

Search for k nearest neighbors in normalized generative data for a data record. When the data record contains NA values only the non-NA values are considered in search. By default a linear search is performed. When a search tree is used search is performed on a tree which is built once in the first function call. Building a tree is also triggered when NA values in data records change in subsequent function calls.

Usage

```
gdKNearestNeighbors(dataRecord, k = 1L, useSearchTree = FALSE)
```

Arguments

- `dataRecord` List containing a data record
`k` Number of nearest neighbors
`useSearchTree` Boolean value indicating if a search tree should be used.

Value

A list of rows in denormalized generative data

Examples

```
## Not run:
gdRead("gd.bin")
gdKNearestNeighbors(list(5.1, 3.5, 1.4, 0.2), 3)
## End(Not run)
```

gdPlotDataSourceParameters

Specify plot parameters for data source

Description

Specify plot parameters for data source passed to function `gdPlotProjection()`.

Usage

```
gdPlotDataSourceParameters(percent = 100, color = "blue")
```

Arguments

- `percent` Percent of randomly selected rows in data source
`color` Colour for data points of data source

Value

List of plot parameters for data source

Examples

```
## Not run:
gdPlotDataSourceParameters(2500)
## End(Not run)
```

<code>gdPlotParameters</code>	<i>Specify plot parameters for generative data</i>
-------------------------------	--

Description

Specify plot parameters for generative data passed to function `gdPlotProjection()`. When density value thresholds with assigned colors are specified generative data is drawn for density value ranges in increasing order.

Usage

```
gdPlotParameters(
  percent = 10,
  densityValueThresholds = c(),
  densityValueColors = c("red")
)
```

Arguments

<code>percent</code>	Percent of randomly selected rows in generative data
<code>densityValueThresholds</code>	Vector of density value thresholds
<code>densityValueColors</code>	Vector of colors assigned to density value thresholds. The size must be the size of <code>densityValueThresholds</code> plus one.

Value

List of plot parameters for generative data

Examples

```
## Not run:
gdPlotParameters(50, c(0.75), c("red", "green"))
## End(Not run)
```

<code>gdPlotProjection</code>	<i>Create an image file for generative data and data source</i>
-------------------------------	---

Description

Create an image file containing two-dimensional projections of generative data and data source. Plot `pagd_2500_6.binrameters` for generative data and data source are passed by functions `gdPlotParameters()` and `gdPlotDataSourceParameters()`. Data points of data source are drawn above data points of generative data.

Usage

```
gdPlotProjection(
  imageName,
  title,
  columnIndices,
  generativeDataParameters = gdPlotParameters(percent = 10, densityValueThresholds = c(),
    densityValueColors = c("red")),
  dataSourceParameters = gdPlotDataSourceParameters(percent = 100, color = "blue")
)
```

Arguments

imageName Name of image file
title Title of image
columnIndices Vector of two column indices that are used for the two-dimensional projections.
Indices refer to indices of active columns of data source.
generativeDataParameters
Plot generative data parameters, see function `gdPlotParameters()`.
dataSourceParameters
Plot data source parameters, see function `gdPlotDataSourceParameters()`.

Value

None

Examples

```
## Not run:
gdRead("gd.bin", "ds.bin")
gdPlotProjection("gd12ddv.png",
  "Generative Data with a Density Value Threshold for the Iris Dataset", c(1, 2),
  gdPlotParameters(250000, c(0.71), c("red", "green")),
  gdPlotDataSourceParameters(2500))
gdPlotProjection("gd34ddv.png",
  "Generative Data with a Density Value Threshold for the Iris Dataset", c(3, 4),
  gdPlotParameters(250000, c(0.71), c("red", "green")),
  gdPlotDataSourceParameters(2500))
## End(Not run)
```

gdRead

Read generative data and data source

Description

Read generative data and data source from specified files. Read in generative data and data source are accessed in `gdPlot2dProjection()`, generative data is accessed in `gdGetRow()`, `gdCalculateDensityValue()` and `gdCalculateDensityValueQuantile()`.

Usage

```
gdRead(generativeDataFileName, dataSourceFileName = "")
```

Arguments

generativeDataFileName	Name of generative data file
dataSourceFileName	Name of data source file

Value

None

Examples

```
## Not run:  
gdRead("gd.bin", "ds.bin")  
## End(Not run)
```

gdServiceDelete	<i>Delete a generated job from software service for accelerated training of generative models</i>
-----------------	---

Description

Delete a generated job from software service. If the job is currently executed it will be stopped.

Usage

```
gdServiceDelete(url, accessKey, jobId)
```

Arguments

url	URL of software service for accelerated training of generative models
accessKey	Unique key for access to software service
jobId	Job id for generated job for training a generative model

Value

None

Examples

```
## Not run:  
url <- "http://xxx.xxx.xxx.xxx/gdService"  
accessKey <- "xxxxxxxx-xxxx-xxxx-xxxx-xxxxxxxxxxxx"  
gdServiceDelete(url, accessKey, 1)  
## End(Not run)
```

gdServiceGetGenerativeData

Get generative data from software service for accelerated training of generative models for processed job

Description

Download generated generative data from software service for processed job. The status of the job has to be TRAINED.

Usage

```
gdServiceGetGenerativeData(
  url,
  accessKey,
  jobId,
  generativeDataFileName = NULL
)
```

Arguments

url	URL of software service for accelerated training of generative models
accessKey	Unique key for access to software service
jobId	Job id for generated job for training a generative model
generativeDataFileName	Name of generative data file. If name is NULL or empty string name of passed name to function gdServiceTrain() will be used.

Value

None

Examples

```
## Not run:
url <- "http://xxx.xxx.xxx.xxx/gdService"
accessKey <- "xxxxxxxx-xxxx-xxxx-xxxx-xxxxxxxxxxxx"
gdServiceGetGenerativeData(url, accessKey, 1, "gd.bin")
## End(Not run)
```

gdServiceGetGenerativeModel

Get generative model from software service for accelerated training of generative models for processed job

Description

Download trained generative model from software service for processed job. The status of the job has to be TRAINED.

Usage

```
gdServiceGetGenerativeModel(  
  url,  
  accessKey,  
  jobId,  
  generativeModelFileName = NULL  
)
```

Arguments

url	URL of software service for accelerated training of generative models
accessKey	Unique key for access to software service
jobId	Job id for generated job for training a generative model
generativeModelFileName	Name of generative model file. If name is NULL or empty string name of passed name to function gdServiceTrain() will be used.

Value

None

Examples

```
## Not run:  
url <- "http://xxx.xxx.xxx.xxx/gdService"  
accessKey <- "xxxxxxxx-xxxx-xxxx-xxxx-xxxxxxxxxxxx"  
gdServiceGetGenerativeModel(url, accessKey, 1, "gm.bin")  
## End(Not run)
```

gdServiceGetStatus	<i>Get status of generated job from software service for accelerated training of generative models</i>
--------------------	--

Description

Get status of generated job from software service. Defined status values are: CREATED, UPLOADED, TRAINING, TRAINED, DELETED, ERROR.

Usage

```
gdServiceGetStatus(url, accessKey, jobId)
```

Arguments

url	URL of software service for accelerated training of generative models
accessKey	Unique key for access to software service
jobId	Job id for generated job for training a generative model

Value

List containing status information

Examples

```
## Not run:
url <- "http://xxx.xxx.xxx.xxx/gdService"
accessKey <- "xxxxxxxx-xxxx-xxxx-xxxx-xxxxxxxxxxxx"
gdServiceGetStatus(url, accessKey, 1)
## End(Not run)
```

gdServiceTrain	<i>Send a request to software service for accelerated training of generative models to train a generative model for a data source</i>
----------------	---

Description

Send a request to software service to train a generative model. A data source file will be uploaded and a job for training will be generated. A job id for the generated job will be returned which has to be used in related requests. The job will be processed as soon as other waiting jobs are processed. When a name of an existing generative model file is passed the file will be also uploaded and the job will continue the training. See also functions gdTrain(), gdServiceGetGenerativeModel(), gdServiceGetGenerativeData(), gdServiceGetStatus(), gdServiceDelete().

Usage

```
gdServiceTrain(
  url,
  accessKey,
  generativeModelFileName,
  generativeDataFileName,
  dataSourceFileName,
  trainParameters = gdTrainParameters(numberOfTrainingIterations = 10000,
    numberOfInitializationIterations = 2500, numberOfHiddenLayerUnits = 1024,
    learningRate = 7e-05, dropout = 0.05)
)
```

Arguments

url	URL of software service for accelerated training of generative models
accessKey	Unique key for access to software service
generativeModelFileName	Name of generative model file
generativeDataFileName	Name of generative data file. If name is NULL or empty string generated data will not be written to a file.
dataSourceFileName	Name of data source file
trainParameters	Generative model training parameters, see function gdTrainParameters().

Value

Job Id number

Examples

```
## Not run:
url <- "http://xxx.xxx.xxx.xxx/gdService"
accessKey <- "xxxxxxxx-xxxx-xxxx-xxxx-xxxxxxxxxxxx"
trainParameters <- gdTrainParameters(numberOfTrainingIterations = 10000)
jobId <- gdServiceTrain(url, accessKey, "gm.bin", "gd.bin", "ds.bin", trainParameters)
## End(Not run)
```

gdTrain

Train a generative model for a data source

Description

Read a data source from a file, train a generative model that generates generative data for the data source in iterative training steps, write trained generative model and generated data in training steps to a file in binary format. When a higher number of iterations is used the distribution of generated data will get closer to that of the data source. When a name of an existing generative model file is passed training will be continued.

Usage

```
gdTrain(
  generativeModelFileName,
  generativeDataFileName,
  dataSourceFileName,
  columnIndices,
  trainParameters = gdTrainParameters(numberOfTrainingIterations = 10000,
    numberOfInitializationIterations = 1500, numberOfRowsHiddenLayerUnits = 1024,
    learningRate = 7e-05, dropout = 0.05)
)
```

Arguments

<code>generativeModelFileName</code>	Name of generative model file
<code>generativeDataFileName</code>	Name of generative data file. When name is NULL or empty string generated data will not be written to a file.
<code>dataSourceFileName</code>	Name of data source file
<code>columnIndices</code>	Vector of two column indices that are used to plot two-dimensional projections of normalized generated generative data and data source for a training step. Indices refer to indices of active columns of data source. Plotting can be disabled by passing NULL or an empty vector.
<code>trainParameters</code>	Generative model training parameters, see function <code>gdTrainParameters()</code> .

Value

None

Examples

```
## Not run:
trainParameters <- gdTrainParameters(numberOfTrainingIterations = 10000)
gdTrain("gm.bin", "gd.bin", "ds.bin", c(1, 2), trainParameters)
## End(Not run)
```

<code>gdTrainParameters</code>	<i>Specify parameters for training of generative model</i>
--------------------------------	--

Description

Specify parameters for training of neural networks used for generation of generative data. These parameters are passed to function `gdTrain()`.

Usage

```
gdTrainParameters(
  numberOfTrainingIterations = 10000,
  numberOfInitializationIterations = 1500,
  numberOfHiddenLayerUnits = 1024,
  learningRate = 7e-05,
  dropout = 0.1
)
```

Arguments

numberOfTrainingIterations	Number of training iterations
numberOfInitializationIterations	Number of initialization iterations
numberOfHiddenLayerUnits	Number of hidden layer units
learningRate	Learning rate for training of neural networks
dropout	Value in the range of 0 to 1. Specifies the rate of hidden units that are dropped. Dropout is a regularization method to prevent overfitting.

Value

List of parameters for training of generative model

Examples

```
## Not run:
generateParameters <- gdGenerateParameters(numberOfTrainingIterations = 10000)
## End(Not run)
```

gdWriteSubset

Write subset of generative data

Description

Write subset of randomly selected rows of generative data

Usage

```
gdWriteSubset(fileName, percent)
```

Arguments

fileName	Name of subset generative data file
percent	Percent of randomly selected rows

Value

None

Examples

```
## Not run:  
gdRead("gd.bin")  
gdWriteSubset("gds.bin", 50)  
## End(Not run)
```

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