Package 'DistributionOptimization'

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Type Package

Title Distribution Optimization

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Description Fits Gaussian Mixtures by applying evolution. As fitness function a mixture of the chi square test for distributions and a novel measure for approximating the common area under curves between multiple Gaussians is used. The package presents an alternative to the commonly used Likelihood Maximization as is used in Expectation Maximization. The algorithm and applications of this package are published under: Lerch, F., Ultsch, A., Lotsch, J. (2020) <doi:10.1038/s41598-020-57432-w>. The evolution is based on the 'GA' package: Scrucca, L. (2013) <doi:10.18637/jss.v053.i04> while the Gaussian Mixture Logic stems from 'AdaptGauss': Ultsch, A, et al. (2015) <doi:10.3390/ijms161025897>.

Imports ggplot2, GA, AdaptGauss, graphics, stats, utils, pracma

Suggests parallelDist

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DistributionOptimization-package Distribution Optimization

Description

Distribution Optimization fits gaussian mixture models on to one dimensional samples by minimizing the Chi Squared Error by evolutional optimization. It is an alternative to likelihood maximizers like expection maximization. Through the included "Overlapping" Methods, single gaussians can be forced to be separated, achieving various significant models to choose from. The evolutionary part is done through the "GA" Package. The Gaussian Mixture Logic is based on the "AdaptGauss" Package.

Author(s)

Florian Lerch, Jorn Lotsch, Alfred Ultsch

References

Luca Scrucca (2013). GA: A Package for Genetic Algorithms in R. Journal of Statistical Software, 53(4), 1-37. URL http://www.jstatsoft.org/v53/i04/

BinProb4Mixtures Bin Probabilities

Description

Calculates the probability of bins/intervals within the dataspace defined by given breaks between them.

Usage

```
BinProb4Mixtures(Means, SDs, Weights, Breaks, IsLogDistribution = rep(F,
length(Means)), LimitsAreFinite = T)
```

Arguments

Means	Means of the GMM Components		
SDs	Standard Deviations of the GMM Components		
Weights	Weights of the GMM Components		
Breaks	Breaks Defining c-1 or c+1 bins (depending on LimitsAreFinite)		
IsLogDistribution			
	If True, the GMM is interpreted as a logarithmic		
LimitsAreFinite			
	If True, there are $c+1$ Bins, where the first and last bin are of inifinite size		

If True, there are c+1 Bins, where the first and last bin are of inifinite size

Value

Probabalities of either c-1 or c+1 bins/intervals (depending on LimitsAreFinite)

Author(s)

Florian Lerch

Examples

```
Data = c(rnorm(50,1,2), rnorm(50,3,4))
NoBins = 20
breaks = seq(min(Data),max(Data), length.out=length(NoBins)+1)
BinProb4Mixtures(c(1,3), c(2,4), c(0.5,0.5), breaks)
```

DistributionOptimization

Distribution Fitting

Description

Fits a Gaussian Mixture Model onto a Dataset by minimizing a fitting error through evolutionary optimization. Every individual encodes one GMM. Details over the evolutionary process itself can be taken from the 'GA' package. ga

Usage

```
DistributionOptimization(Data, Modes, Monitor = 1,
SelectionMethod = "UnbiasedTournament",
MutationMethod = "Uniform+Focused",
CrossoverMethod = "WholeArithmetic", PopulationSize = Modes * 3 * 25,
MutationRate = 0.7, Elitism = 0.05, CrossoverRate = 0.2,
Iter = Modes * 3 * 200, OverlapTolerance = NULL,
IsLogDistribution = rep(F, Modes), ErrorMethod = "chisquare",
NoBins = NULL, Seed = NULL, ConcurrentInit = F, ParetoRad = NULL)
```

Arguments

Data	Data to be modelled		
Modes	Number of expected Modes		
Monitor	0:no monitoring, 1: status messages, 2: status messages and plots, 3: status messages, plots and calculated error-measures		
SelectionMethod			
	1: LinearRank selection 4: UnbiasedTournament 5: FitnessProportional selec- tion		

MutationMethod	1: UniformRandom mutation 2: NonuniformRandom mutation 4: Focused mu-		
	tation, alternative random mutation around solution 5: GaussMutationCust 6:		
	TwoPhaseMutation - mutation is uniform random during the first half of itera-		
	tions, and than focuses around current solution		
CrossoverMethod			
	1: single point crossover 2: whole arithmetic crossover 3: local arithmetic crossover 4: blend crossover 5: GaussCrossover - exchange complete gaussian components 6: MultiPointCrossover - Random amount of information between mixtures get exchanged		
PopulationSize	Size of the population		
MutationRate	amount (01) of population that gets mutated		
Elitism	amount of best individuals that will survive generation unchanged		
CrossoverRate	amount of individuals that will be used for crossover		
Iter	number of iterations of this algorithm		
OverlapTolerance			
	$ratio \ between \ Chi-Square \ and \ Overlap Error \ (only \ if \ Fitness Method = 4 \ (Chi2Value With Overlap))$		
IsLogDistribution			
	which gauss components should be considered as log gaussian		
ErrorMethod	"pde": fitting is measured by pareto density estimation. "chisquare": fitting is measured by Chi-Square test		
NoBins	Number of Bins that will be used for evaluating fitting		
Seed	Random Seed for reproducible results		
ConcurrentInit	If true, before initialization a number of short optimizations are done to find a good starting point for evolution		
ParetoRad	Pareto Radius for Pareto Density Estimation and its plots		

Value

The GA object containing the evolutionary training and a description of the final GMM consisting of means, sdevs and weights.

Author(s)

Florian Lerch Jorn Lotsch Alfred Ultsch

Examples

```
## Not run:
DistributionOptimization(c(rnorm(200),rnorm(200,3), 2))
```

End(Not run)

MixedDistributionError

MixedDistributionError

Description

Calculates a fitting error as well as the overlapping measure for the mixtures. Combines them with ratio rho in favor of Overlapping.

Usage

```
MixedDistributionError(Means, SDs, Weights, Data, rho = 0.5,
    breaks = NULL, Kernels = NULL, ErrorMethod = "chisquare")
```

Arguments

Means of the GMM Components
Standard Deviations of the GMM Components
Weights of the GMM Components
Empirical Data based on which the GMM is build
Ratio of OverlappingError vs Fitting Error
vector containing the breaks between bins
positions at which density is to be compared
"pdeerror": fitting error is measured through Pareto Density Estimation. "chisquare": fitting error is measured through the Chi Square fitting error.

Value

Mixed Error

Author(s)

Florian Lerch

Examples

```
Data = c(rnorm(50,1,2), rnorm(50,3,4))
MixedDistributionError(c(1,3), c(2,4), c(0.5,0.5), Data = Data)
```

OverlapErrorByDensity OverlapErrorByDensity

Description

Similarity in GMM by Density

Usage

```
OverlapErrorByDensity(Means, SDs, Weights, Data = NULL, Kernels = NULL)
```

Arguments

Means	Means of the GMM Components
SDs	Standard Deviations of the GMM Components
Weights	Weights of the GMM Components
Data	Dataset that the GMM should be compared with
Kernels	if length(Kernels) = 1: amount of kernels if length(Kernels) > 1: kernels in dataspace at which the GMM Components will be compared with each other

Details

Calculates the similarity (overlap) between multiple modes in Gaussian Mixture Models. Kernels at equally distanced positions are used, if not explicitly given.

Value

List: OverlapError Error for estimating the maximal Overlap of AUC of PDFs of each pair of GMM Components Kernels Kernels that were used for comparing the GMM Components

Author(s)

Florian Lerch

Examples

```
Data = c(rnorm(50,1,2), rnorm(50,3,4))
V<-OverlapErrorByDensity(c(1,3), c(2,4), c(0.5,0.5), Data = Data)
AdaptGauss::PlotMixtures(Data, c(1,3), c(2,4), SingleGausses = TRUE)
print(V$OverlapError)</pre>
```

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