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Comparison of EM-algorithm and MLE using Cholesky decomposition

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Abstract

The intent of this work is to compare The EM algorithm to a MLE approach in the case of multivariate normal mixture models using the Cholesky decomposition. The EM algorithm is widely used in statistics and is proven to converge, however in pathological cases convergence slows down considerably.

methods(not done) results(not done)

CONTENTS

${\bf Contents}$

	roduction to normal mixture models	1
1.1	Definitions	1
1.2	The EM-Algorithm in Sketch	2
1.3	Choice of Notation	3
1.4	Models of Covariance Matrices	4
1.5	Problems of the EM-algorithm	6
1.6	Alternative Option	7
The	e norMmix Package	9
2.1	Introduction to the Package	9
	2.1.1 norMmixMLE	10
2.2	On The Development of norMmix	12
2.3	Demonstration	13
Cor	nparing Algorithms	15
3.1	Time Analysis	16
3.2	Behaviour in n	17
3.3	Behaviour in p	17
3.4	Diffixult Mixtures	18
3.5	Nonnormal Mixtures	19
Dis	cussion	27
Bib	liography	28
R C	ode	31
A.1	llnorMmix	31
Fur	ther Plots	37
R 1	Chapter 3	37
	1.2 1.3 1.4 1.5 1.6 The 2.1 2.2 2.3 Con 3.1 3.2 3.3 3.4 3.5 Disc Bib R C A.1 A.2 Fur	1.2 The EM-Algorithm in Sketch 1.3 Choice of Notation 1.4 Models of Covariance Matrices 1.5 Problems of the EM-algorithm 1.6 Alternative Option The norMmix Package 2.1 Introduction to the Package 2.1.1 norMmixMLE 2.2 On The Development of norMmix 2.3 Demonstration Comparing Algorithms 3.1 Time Analysis 3.2 Behaviour in n 3.3 Behaviour in p 3.4 Diffixult Mixtures 3.5 Nonnormal Mixtures Discussion Bibliography R Code A.1 11norMmix A.2 Example Simulation Script Further Plots

List of Figures

1.1	Parameters of MW.nm9
1.2	True and Estimated density
1.3	20 EM steps
1.4	200 EM steps
1.5	Log-likelihood Plotted against Iteration Count for the Example in 1.5
2.1	Demonstration of the MW Objects
	Correct Mixture (left) and Fitted overlayed in orange (right)
3.1	Example of Comparison Plot
	Log-log Plot of System Time against Parameter Length

LIST OF TABLES

T	• ,	C		1 1	
	ist	α t	' l '9	h	ΔC
	1100	\ //	10		

1.1	Table of Parameters of the Covariance Matrices	Ę
1.2	Full Table of Parameters	(
9 1	Translation Table: Mathematical Notation to R Code	(

1 Chapter 1

2 Introduction to normal mixture models

Definitions 1.1

- A good and thorough introductory book is the work of McLachlan and Peel (2000) and the
- reader is encouraged to study it to learn in depth about normal mixtures and clustering.
- We will here give a short overview of normal mixtures to fix notation and nomenclature.
- The motivating idea behind mixture models is, that in real world examples a sample
- might be suspected to arise from more than one population or be more simply modelled
- by several overlayed distributions. The example of this, that is generally considered to
- be the first of this kind, is the one by Karl Pearson, who fitted two normal distributions 11
- with different means and variances. In his book, Pearson and Henrici (1896) Section 4.d.; 12
- page 266, Pearson analyzed measurements of forehead to body length of crabs sampled 13
- from the bay of Naples. His mixture model-based approach suggested, that the crabs 14
- were evolving into two new subspecies. This is a historically important example, because 15
- it presents statistical evidence of evolution in process. Mixture models have been used
- since, but research took off after the availability of computing power made computational research possible 18
- While the theory of mixture models holds for a much broader class of distributions, we 19
- restrict ourselves here to the case of normal distributions, because this restriction fits
- more comfortably into the scope of this work and because normal distributions allow for 21
- a parsimonious parametrization, that is of interest to study. 22
- This parametrization is the $LDL\top$ decomposition, which allows a very simple parametriza-
- tion and a straightforward connection between degrees of freedom and necessarily gener-24
- ated numerical values. This will be explained further in section 1.4.
- But before we delve deeper into the topic of this research, we first define the concept of a
- normal mixture model:
- $\Sigma \in \mathbb{R}^{p \times p}$ be symmetric positive definite and $\phi(-; \mu, \Sigma)$ be the normal

distribution with mean μ and covariance matrix Σ with density function:

$$\phi(\boldsymbol{x}; \boldsymbol{\mu}, \boldsymbol{\Sigma}) = \frac{\exp(-\frac{1}{2}(\boldsymbol{x} - \boldsymbol{\mu})\boldsymbol{\Sigma}^{-\frac{1}{2}}(\boldsymbol{x} - \boldsymbol{\mu})^{\top})}{\sqrt{(2\pi)^k \det \boldsymbol{\Sigma}}}$$
(1.1.0.1)

for $\boldsymbol{x} \in \mathbb{R}^p$. Since we are studying mixture models, we will need several overlapping of normal distributions, of differing means and covariance. Therefore, we choose notation allowing us to refer to the components in shorthand. Let us assume we have $K \in \mathbb{N}$ normal distributions with means and covariance $\boldsymbol{\mu}_k, \boldsymbol{\Sigma}_k, \quad k \in \{1, \dots, K\}$, then we fix:

$$\phi_k(\boldsymbol{x}) := \phi(\boldsymbol{x}; \boldsymbol{\mu_k}, \boldsymbol{\Sigma_k}) \tag{1.1.0.2}$$

And going forward, we will refer to components by the subscript k.

Definition 1.1.0.1. Suppose we have a random sample Y_1, \ldots, Y_n , where Y_i is a pdimensional random vector with probability density function $Y_i \sim f(y_i)$ on \mathbb{R}^p .

We assume that the density $f(y_i)$ of Y_i can be written in the form:

$$f(\mathbf{y}_i) = \sum_{k=1}^{K} \pi_k \phi_k(\mathbf{y}_i)$$
(1.1.0.3)

The ϕ_k are normal distributions and are called the mixture components with parameters μ_k and Σ_k as described above (1.1.0.2). The π_k are called the component densities of the mixture and are constrained by the rules $\pi_k > 0$ and $\Sigma_k \pi_k = 1$.

For 'large' datasets there are more parsimonious parametrizations, that reduce computation time. These, for example, assume that all components have the same covariance, or have certain restrictions placed on them. We will give a detailed description of the models assumed in this thesis in section 1.4.

⁴⁵ 1.2 The EM-Algorithm in Sketch

With this definition, we immediately face the problem of how to fit these mixture components to given data. A popular algorithm to solve this problem is the Expectation-

48 Maximization algorithm, abbreviated as EM-algorithm.

We give here a sketch of the EM-algorithm in the case of all normal mixture components.

50 This roughly follows the content in McLachlan and Peel (2000). For a more thorough

treatment of the matter see chapter 3.

Suppose we have a p-dimensional dataset of n samples x_1, \ldots, x_n , onto which we would

like to fit a K component normal mixture with mixture components $\phi_k, k \in 1, \ldots, n$.

For the EM-algorithm further parameters are introduced. These are denoted $\tau_i(y_i)$ and

they represent the posterior probabilities that observation i is a member of component j.

The EM-algorithm is a two step, iterative process consisting of an 'e'-step and an 'm'-step.

In the e-step the expectation of component membership is updated.

$$\tau_j(\boldsymbol{y}_i; \boldsymbol{\Psi}) = \phi_j(\boldsymbol{y}_i) / \sum_{k=1}^K \phi_k(\boldsymbol{y}_i)$$
 (1.2.0.1)

and in the m-step given the component membership information we update the component means and covariances by weighted versions of the usual estimators.

$$\boldsymbol{\mu}_j = \sum_{i=1}^n \tau_j(\boldsymbol{y}_i) \boldsymbol{y}_i / \sum_{j=1}^n \tau_j(\boldsymbol{y}_i)$$
 (1.2.0.2)

$$\Sigma_j = \sum_{i=1}^n \tau_j(\boldsymbol{y}_i)(\boldsymbol{y}_i - \boldsymbol{\mu}_j)(\boldsymbol{y}_i - \boldsymbol{\mu}_j)^\top / \sum_{i=1}^n \tau_j(\boldsymbol{y}_i)$$
 (1.2.0.3)

There remains to be stated how to start the algorithm. Since both steps of the algorithm depend on data from the other, the EM-algorithm needs some form of initialization step.

Most popular implementations use some form of pre clustering and use the EM-algorithm as subsequent tools to fit the data. The R-package mclust for example uses hierarchical agglomerative clustering Scrucca, Fop, Murphy, and Raftery (2016).

₆ 1.3 Choice of Notation

The classification of models in this paper relies heavily on the work of Celeux and Govaert (1995), however, out of necessity for clarity, we break with their notation. So as to not confuse the reader we describe here in depth the differences in notation between Celeux and Govaert (1995) and ours.

The basis of classification in Celeux and Govaert (1995) is the decomposition of a symmetric matrix into an orthogonal and a diagonal component. A symmetric positive definite matrix Σ can be decomposed as follows

$$\Sigma = \lambda \mathbf{D} \mathbf{A} \mathbf{D}^{\top} \tag{1.3.0.1}$$

with D an orthogonal matrix and A a diagonal matrix and $\lambda = \sqrt[p]{\det(\Sigma)}$ the p-th root of the determinant of Σ .

This decomposition has an appealing geometric interpretation, with D as the orientation of the distribution, A the shape, and λ the volume. The problem of notation comes from standard conventions in linear algebra, where the letters A and D are usually occupied by arbitrary and diagonal matrices respectively. Furthermore, we intend to apply a variant of the Cholesky decomposition to Σ , the αLDL^{\top} decomposition. This obviously raises some conflicts in notation.

Therefore we, from here on, when referring to the decomposition as described by Celeux and Govaert (1995), will use the following modification of notation:

$$D \longmapsto Q$$
 (1.3.0.2)

$$A \longmapsto \Lambda$$
 (1.3.0.3)

$$\lambda \longmapsto \alpha$$
 (1.3.0.4)

$$\Sigma = \lambda \mathbf{D} \mathbf{A} \mathbf{D}^{\top} = \alpha \mathbf{Q} \mathbf{\Lambda} \mathbf{Q}^{\top}$$
 (1.3.0.5)

These were chosen according to general conventions of linear algebra. Q is usually chosen for orthonormal matrices; Λ is often a choice for diagonal matrices of eigenvectors and α was somewhat arbitrarily chosen.

92

7 1.4 Models of Covariance Matrices

As mentioned above, there are ways to constrain the covariance matrices for computational reasons. There are istances where the resulting loss of information is seen as acceptable, for example if, through consideration of the data, that a simplified model is acceptable.

Another is if the sheer size of the data makes application of full generality impossible.

- We restrict the complexity of the decomposition components
- We restrict the variability of the mixture components

Let us look at the first case. We take the decomposition of a covariance matrix as $\Sigma = \alpha Q \Lambda Q^{\top}$. Of these, we can simplify the structure of Q and Λ , by replacing them with the identity. If we set Q = Id, we lose the freedom of orientation and if we set $\Lambda = \text{Id}$ we restrict ourselves to spherical distributions.

of course, we cannot restrict λ while letting q free, since

$$\mathbf{Q}\Lambda\mathbf{Q}^{\top} = \mathbf{Q}\mathrm{Id}\mathbf{Q}^{\top} = \mathrm{Id} \tag{1.4.0.1}$$

The second restriction simply means we hold the decomposition fixed throughout all covariance matrices. There is however an issue with the cholesky decomposition. For 10 out of 14 cases as defined by Celeux and Govaert (1995), there exists a canonical translation of decompositions. The 6 diagonal cases need no translation; the eigen and cholesky decomposition are equal to identity. For the non-diagonal cases note that for a given sym. pos. def. matrix Σ we have decompositions:

$$\Sigma = \alpha \mathbf{Q} \Lambda \mathbf{Q}^{\top} \quad \Sigma = \alpha \mathbf{L} \mathbf{D} \mathbf{L}^{\top}$$
 (1.4.0.2)

Since in both cases the enclosing matrices Q and L have determinant 1 the determinant 105 of Σ falls entirely on α . therefore α , in these particular decompositions, is equal for both. 106 Celeux and Govaert (1995) vary σ by either varying or holding fixed the volume (α/α_k) , 107 shape (Λ/Λ_k) and orientation (Q/Q_k) . These 3 times 2 cases would yield the 8 out of 14 108 cases of non-diagonal cases. However there is no canonical transform for either variable 109 orientation and fixed shape or fixed orientation and variable shape. The reason for this is 110 that in the LDL^{\top} decomposition the lower diagonal matrix L holds some of the shape of 111 the matrix, which in the eigendecomposition is in the Λ matrix. In fact, L is orthogonal 112 if and only if $L = \mathrm{Id}_{n \times n}$. Therefore we can only decompose matrices where either both or neither shape and orientation vary. See table 1.1.

While we could in theory construct the cases $\boldsymbol{L}\boldsymbol{D}_k\boldsymbol{L}^{\top}$ and $\boldsymbol{L}_k\boldsymbol{D}\boldsymbol{L}_k^{\top}$, however they do not correspond to the desired geometric intent behind the differentiation of models and are therefore not included.

Model	Σ_k C&G	volume	shape	orientation	parameters	$oldsymbol{TDT}_{\perp}$	parameters	count
EII	$oldsymbol{a}$	ednal	ednal	1	σ	as in $C\&G$		П
VIII	$lpha_k m{I}$	var.	equal	1	α_k			K
EEI	$\alpha \mathbf{A}$	equal	equal	coord. axes	$lpha, \lambda_i$			1 + (p-1)
VEI	$lpha_k oldsymbol{\Lambda}$	var.	equal	coord. axes	$lpha_k, \lambda_i$			K + (p-1)
EVI	$lpha oldsymbol{\Lambda}_k$	equal	var.	coord. axes	$\alpha,\lambda_{i,k}$			1+K(p-1)
VVI	$lpha_k oldsymbol{\Lambda}_k$	var.	var.	coord. axes	$\alpha_k, \lambda_{i,k}$			K + K(p-1)
EEE	$lpha oldsymbol{Q} oldsymbol{\Lambda} oldsymbol{Q} oldsymbol{Q}^{ op}$	equal	equal	edual	$\alpha,\lambda_i,q_{i,j}$	$lpha m{LDL}^{ op}$	$\lambda, d_i, l_{i,j}$	$1 + (p-1) + \frac{p(p-1)}{2}$
EVE	$lpha oldsymbol{Q} oldsymbol{\Lambda}_k oldsymbol{Q}^ op$	equal	var.	ednal	$\alpha, \lambda_{i,k}, q_{i,j}$	doesn't exist		$1 + K(p-1) + \frac{p(p-1)}{2}$
VEE	$\alpha_k \boldsymbol{Q} \boldsymbol{\Lambda} \boldsymbol{Q}^\top$	var.	equal	ednal	$lpha_k, \lambda_i, q_{i,j}$	$\alpha_k \boldsymbol{L} \boldsymbol{D} \boldsymbol{L}^\top$	$\lambda_k, d_i, l_{i,j}$	$K + (p-1) + \frac{p(p-1)}{2}$
VVE	$lpha_k oldsymbol{Q} \mathbf{\Lambda}_k oldsymbol{Q}^ op$	var.	var.	ednal	$\alpha_k, \lambda_{i,k}, q_{i,j}$			$K + K(p-1) + \frac{p(p-1)}{2}$
EEV	$lpha oldsymbol{Q}_k oldsymbol{\Lambda} oldsymbol{Q}_k^ op$	equal	equal	var.	$\alpha,\lambda_i,q_{i,j,k}$	don't exist		$1 + (p-1) + K \frac{p(p-1)}{2}$
VEV	$lpha_k oldsymbol{Q}_k oldsymbol{\Lambda} oldsymbol{Q}_k^ op$	var.	equal	var.	$\alpha_k, \lambda_i, q_{i,j,k}$			$K + (p-1) + K \frac{p(p-1)}{2}$
EVV	$lpha oldsymbol{Q}_k oldsymbol{\Lambda}_k oldsymbol{Q}_k^ op$	equal	var.	var.	$lpha,\lambda_i,q_{i,j,k}$	$lpha oldsymbol{L}_k oldsymbol{D}_k oldsymbol{L}_k^ op$	$\lambda, d_{i,k}, l_{i,j,k} \ j > i$	$\lambda, d_{j,k}, l_{i,j,k} \ j > i $ $1 + K(p-1) + K^{\frac{p(p-1)}{2}}$
VVV	$lpha_k oldsymbol{Q}_k oldsymbol{\Lambda}_k oldsymbol{Q}_k^ op$	var.	var.	var.	$\alpha_k, \lambda_i, q_{i,j,k}$	$lpha_koldsymbol{L}_koldsymbol{D}_koldsymbol{L}_k^ op$	$\lambda_k, d_{i,k}, l_{i,j,k} \ j > i$	$\lambda_k, d_{i,k}, l_{i,j,k} \ j > i K + K(p-1) + K^{\frac{p(p-1)}{2}}$

Table 1.1: Table of Parameters of the Covariance Matrices

126

Σ model	μ, π	Σ	reduced	$\mathcal{O}()$
EII	K-1+pK	1	Kp + K	Kp+K
VII	K-1+pK	K	Kp + 2K - 1	Kp + K
EEI	K-1+pK	1 + (p - 1)	Kp + p + K - 1	Kp + p + K
VEI	K-1+pK	K + (p - 1)	Kp + p + 2K - 2	Kp + p + K
EVI	K-1+pK	1 + K(p-1)	2Kp	Kp
VVI	K-1+pK	K + K(p-1)	2Kp+K-1	Kp
EEE	K-1+pK	$1 + (p-1) + \frac{p(p-1)}{2}$	$\frac{(p+2)(p-1)}{2} + Kp + K$	$p^2 + Kp + K$
VEE		$K + (p-1) + \frac{p(p-1)}{2}$	$\frac{(p+2)(p-1)}{2} + Kp + 2K - 2$	$p^2 + Kp + K$
EVV	K-1+pK	$1 + K(p-1) + K^{\frac{p(p-1)}{2}}$	$K^{\frac{(p+2)(p-1)}{2}} + Kp + K$	$Kp^2 + Kp + K$
VVV	K-1+pK	$K + K(p-1) + K^{\frac{p(p-1)}{2}}$	$K^{\frac{(p+2)(p-1)}{2}} + Kp + 2K - 1$	$Kp^2 + Kp + K$

Table 1.2: Full Table of Parameters

There is an attractive advantage in using the LDL^{\top} decomposition. Since both the LDL^{\top} and eigendecomposition derive from the same covariance matrix, the necessary parameters are the same in cardinality. In the case of the Q and L matrices, there need to be $\frac{p(p-1)}{2}$ parameters to be determined to uniquely define these matrices. In the case of the L matrix these are straightforward the entries of the lower diagonal matrix, whereas Q needs a nontrivial amount of work to determine a minimal generating set of parameters, which makes computation of the decomposition as in Celeux and Govaert (1995) a lot more difficult. Therefore the LDL^{\top} decomposition was chosen for the purpose of this thesis.

1.5 Problems of the EM-algorithm

The EM-algorithm has stalling problems especially close to a local optimum. In their seminal work, Dempster, Laird, and Rubin (1977), have proven that the EM-algorithm converges under mild regularity conditions. However, convergence does not guarantee fast convergence. In fact, a lot of the work, that has gone into the research around the EM-algorithm has been concerned with speeding up convergence, see McLachlan and Peel (2000)[section 2.17]. The concern here is that a slowing in convergence might be mistaken for actual convergence.

This phenomenon is not infrequent and in difficult mixtures quite visible. To illustrate let us look at a particular mixture taken from Marron and Wand (1992) and the nor1mix package from CRAN. nor1mix is a package designed and developed for educational purposes to teach about univariate normal mixtures. It is also the spiritual predecessor of this thesis' R code.

The mixture is a trimodal mixture of uneven weight, as shown in figure 1.2. While not the most difficult mixture studied by Marron and Wand (1992), it is certainly not trivial either. In the figures below, 1.5 and 1.5, we demonstrate, that even after 200 iterations of the EM-algorithm the convergence is poor. In this instance, the initialization is done using R 's CLARA implementation from the cluster package.

then an illustration of MW examples of pathological cases

We can see, that the EM-algorithm seems to converge to an intermediary solution, where the smaller middle solution is weighted lower, until it manages to correct back and find

Figure 1.1: Parameters of MW.nm9

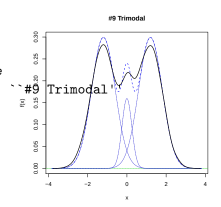


Figure 1.2: True and Estimated density

the correct components.

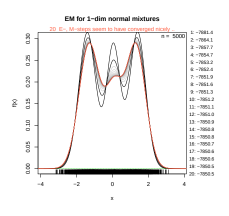


Figure 1.3: 20 EM steps

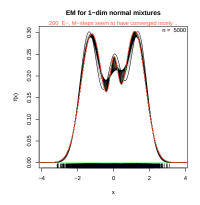


Figure 1.4: 200 EM steps

- We see how change in log-likelihood seems to stagnate. However, this does not stay that way. If we let EM run a bit further we see, the log-likelihood hits a flatspot, after which convergence accelerates again.
- In fact, it seems that the previous solution is a saddle point in the likelihood function, where EM has chronic problems continuing improvements.
- give 2D demonstration.

4 1.6 Alternative Option

- In conclusion, the EM-algorithm has very appealing advantages. However, as we have shown, there are chronic problems in convergence rates. The aim of this thesis is to test if some improvement could be achieved by a different method.
- The plan is reasonably straightforward:
- i.) Initialize using CLARA.
- ii.) Perform one m-step, to transform CLARA's results into the form of a normal mixture.

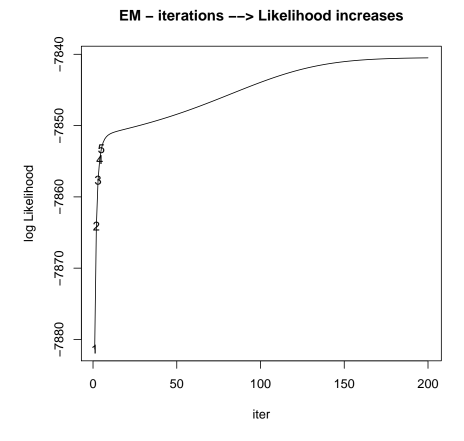


Figure 1.5: Log-likelihood Plotted against Iteration Count for the Example in 1.5

- iii.) Apply a general optimizer, using the mixture's log-likelihood function.
- what do we hope from this? better convergence proof of concept i.e. not complete failure
- raise questions about implementation, clara fctn optim params
- the subsequent chapter is devoted to answering this question by documenting the devel-
- 165 opment of norMmix

$\mathbf{c}_{\scriptscriptstyle{66}}$ Chapter $\mathbf{2}$

The norMmix Package

2.1 Introduction to the Package

For this thesis, an R package was developed that implements the algorithm that fits multivariate normal mixtures to given data. ¹ There is a lot of unused code still in the package. These were at one point implemented used and discarded. They are still included for demonstration. The norMmix package is constructed around the norMmix object that codifies a normal Multivariate mixture model, and the llnorMmix() function, that calculates the log-likelihood given a model and data.

In the table 2.1 the notation used in code is listed along with a translation to the previously used mathematical notation. Additionally, functions with ambiguous names are listed here.

177 The package contains the following functionality:

The package relies on optim from the stats package for general optimization. we use the standard method implemented in optim which is BFGS, which is a quasi-Newton method (also known as a variable metric algorithm) as described in Broyden (1970) among others.

The workflow when using the package is as follows. The function rnorMmix can be used to generate data from a norMmix object. The MW objects provide ready made examples and

¹The package was written with R version 3.6.1 (2019-07-05) last updated on 2019-10-22.

In Notation	In Code
π_i	w, weights
Σ	Sigma
μ	mu
K	k
dimension	p, dim, dims
components	cl, components
Σ model	model
cluster's CLARA	clara
mclust's hierarchical clustering	mclVVV
mclust's Mclust fuction	mclust

Table 2.1: Translation Table: Mathematical Notation to R Code

- norMmix norMmix() is the 'init-method' for norMmix objects. There exist is.norMmix rnorMmix and dnorMmix functions.
- **parametrization** The main functions that handle reparametrization of models from and to LDL^{\top} decomposition are nMm2par and par2nMm, which are inverse to each other.
- MLE The function norMmixMLE marries the main components of this package. It initializes a model and parametrizes it for use with optim
- model choice Using norMmixMLE, the function fitnMm allows fitting of multiple models and components. Functions analyzing the output of this are also provided, e.g. BIC and print methods.
- misc There are also various methods of generics, like logLik, print, BIC, AIC and nobs as well as various print methods.
- example objects Following the paper of Marron and Wand (1992) various example objects are provided and used for study. They follow the naming convention: MW + dimension + number. For example MW213 for the 13th model of dimension 2.
- simulations A good portion of the package is designed with the study of simulations in mind. Therefore there are functions provided to study large collections of evaluated data. e.g compplot

objects of study and the norMmix function can be used to define normal mixtures from scratch. Of course, other data sets can be used for analysis. The following functions rely, however, on the matrix data structure. So dataframes must be converted beforehand and non numerical data is not accepted.

Given data, the functions that accept it for analysis are mainly norMmixMLE and fitnMm.
The former performs model fit on data, and the latter performs model selection, by calling norMmixMLE for specified k and model vectors.

190 **2.1.1** norMmixMLE

The core of norMmixMLE is the application of optim in conjunction with llnorMmix as function to be optimized. llnorMmix can be accessed directly, however, it needs a transposed dataset. As stated in section 1.6 the MLE implicitly performs initialization. There are two options for this initialization step. One is the CLARA clustering algoritm, with non-standard arguments. The standard arguments are somewhat historic in origin and were, at the time, chosen because of hardware limitations. The newer function, due to this thesis' advisor Martin Mächler, was designed to be a 'sensible' alternative, but should be subject to further scrutiny. It is reproduced here.

It is dependent on the size and dimension of the dataset, as well as the demanded number of clusters. The alternative to CLARA is mclust's hierarchical agglomerative clustering, which follows the work of Fraley (1998). The intention behind using mclust's initialization function is to directly compare how much difference the initialization process makes.

The initialization stage does not yield a normal mixture. This requires a way to transform a clustering into a mixture. The method chosen in this package is to use an m-step from the EM-algorithm. Unlike the EM-algorithm, clustering algorithms like CLARA produce binary cluster membership results, whereas the component membership of EM is determined as a probability value between 0 and 1. This is resolved by interpreting the results as probability values which are either 0 or 1. These are then used as the τ_j as described in section 1.2. This m-step is also taken from the mclust package for reasons better explained in section 2.2. It has the advantage of being able to generate a mixture object with the correct covariance model.

This mixture object is still in human readable form and not the necessary parameter vector demanded by optim. So an application of the function nMm2par is carried out, resulting in a starting value for optim.

Due to the nature of the package the returned results are more than abundant. Not only is the fitted model returned but also everything produced by optim and the entire dataset.

Here are listed the stucture the returned values:

```
data(fSMI.12, package="norMmix")
>
      str(fSMI.12$nMm[3,3][[1]], max=2)
List of 6
 $ norMmix:List of 6
            : num [1:20, 1:3] 15.9 30.7 36.2 21.8 753 ...
  ..$ Sigma : num [1:20, 1:20, 1:3] 0.358 0 0 0 0 ...
  ..$ weight: num [1:3] 0.219 0.419 0.362
  ..$ k
            : int 3
  ..$ dim
            : int 20
  ..$ model : chr "EEI"
  ..- attr(*, "name")= chr "model = EEI , clusters = 3"
  ..- attr(*, "class")= chr "norMmix"
 $ optr
          :List of 5
                 : num [1:82] 0.264 0.118 15.903 30.67 36.155 ...
  ..$ par
  ..$ value
                 : num 7370
                 : Named int [1:2] 232 88
  ..$ counts
  ... - attr(*, "names")= chr [1:2] "function" "gradient"
  ..$ convergence: int 0
  ..$ message
                 : NULL
 $ npar
          : int 82
 $ n
          : int 141
 $ x
          : num [1:141, 1:20] 16.1 15.7 15.7 16.1 16.6 ...
  ..- attr(*, "dimnames")=List of 2
          : num 1.72
 - attr(*, "class")= chr "norMmixMLE"
```

Besides mclust the package also relies on a number of other packages for various tasks.
Listed in no particular order: cluster, MASS, mvtnorm, mclust, mixtools and sfsmisc.

since mclust is one of the more popular packages implementing the EM algo, we employ a lot of functions from mclust, to keep things around EM as similar as possible.

222 also relies on mixtools package for random generating function rnorMmix using rmvnorm.

23 2.2 On The Development of norMmix

```
about Cholesky decomp as ldlt. has advantages: fast, parametrically parsimonious, can
224
   easily compute loglikelihood
   One dead-end was the parametrization of the weights of a mixture using the logit func-
226
   tion.
227
    > logit <- function(e) {</pre>
          stopifnot(is.numeric(e) ,all(e >= 0), all.equal(sum(e),1))
          qlogis(e[-1L])
    + }
    > logitinv <- function(e) {</pre>
          if (length(e)==0) {return(c(1))}
          stopifnot(is.numeric(e))
          e<- plogis(e)
          sp. <- sum(e)
          w \leftarrow c((1-sp.), e)
    + }
   This uses the logistical function logis to transform to reduce the number of weights
   from K to K-1. Much like clr1, given a list of weights logit will transform them
   and logiting will correctly reverse the transformation. However, unlike clr1, it will not
   transform an arbitrary list of length K-1 into a valid weight parameter. For example:
    > w <- runif(7); ret <- logitinv(w)
    > ret
    [1] -3.3177621
                      0.6679835 0.7244333
                                              0.6466517 0.5028572 0.5493769
                                                                                    0.6778167
         0.5486428
   The issue here is that the last line of logitiny, which is necessary to sum to one, but
    results in a negative value in ret[1] which is not a valid weight. The underlying issue is
233
    that not every tuple in \mathbb{R}^{K-1} is a result of logit.
    The option to use logit is still an argument to norMmixMLE by specifying trafo="logit",
235
    but it shouldn't be used.
236
   Another issue during development cropped up during fitting of high dimensional data. We
237
   studied the dataset SMI.12 from the package copula:
    > data(SMI.12, package="copula")
    > str(SMI.12)
    num [1:141, 1:20] 16.1 15.7 15.7 16.1 16.6 ...
     - attr(*, "dimnames")=List of 2
      ..$ : chr [1:141] "2011-09-09" "2011-09-12" "2011-09-13" "2011-09-14" ...
      ..$ : chr [1:20] "ABBN" "ATLN" "ADEN" "CSGN" ...
   A consequence of high dimensions is that matrix multiplication is no longer very stable.
   As a result, the covariance matrices produced by our own implementation of the EM-
```

As a result, the covariance matrices produced by our own implementation of the EMalgorithms m-step (mstep.nMm) were not positive definite. In the case of SMI.12, several
covariance matrices are degenerate, which results in cancellation error with near-zero entries. We attempted to correct this with the function forcePositive, which simply tries

2.3 Demonstration 13

> plot(MW215)

Figure 2.1: Demonstration of the MW Objects

to set D in LDL^{\top} greater than zero. This didn't resolve the issue, since a non-negligible part of the numerical error was in the L matrix and the resultant covariance matrix was still not positive definite.

We eventually resolved this issue by abandoning our own implementation and using the functions from the Mclust package. Not only were these numerically stable they were also able to differentiate between models, whereas ours would assume VVV for every fit.

250 testing of mytnorm as proof that ldlt is in fact faster parametrization

set.seed(2019); x <- rnorMmix(500, MW215)</pre>

mention, that there may be faster ways to apply backsolve. quote knuth about premature optimization?

not possible to sensibly compare normal mixtures except maybe a strange sorting algorithm using Mahalanobis distance or Kullback-Leibler distance or similar (Hellinger), but not numerically sensible to integrate over potentially high-dimensional spaces.

$_{256}$ 2.3 Demonstration

>

To end this chapter, here a small demonstration of the capabilities of norMmix. First a small plot to show an MW mixture.

It is a trimodal mixture along the diagonal.

```
system.time(mleResult <- norMmixMLE(x, 3, "VEE"))</pre>
initial value 2206.907425
iter 10 value 2147.633703
iter 20 value 2125.658743
final value 2125.658364
converged
   user
         system elapsed
  0.256
          0.008
                  0.265
      mleResult
object of class 'norMmixMLE'
norMmix object:
multivariate normal mixture model with the following attributes:
                       model = VEE , components = 3
name:
dimension:
components:
weight of components 0.365 0.325 0.31
returned from optim:
function gradient
      75
               22
```

log-likelihood: -2125.658

```
        nobs
        npar
        nobs/npar

        500
        13
        38.46154
```

Here are the results of a run of norMmixMLE and below the graphical display of the results.

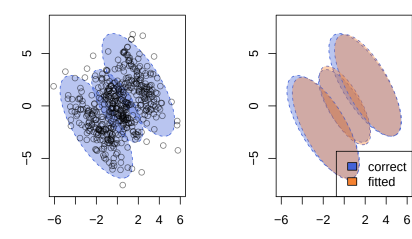


Figure 2.2: Correct Mixture (left) and Fitted overlayed in orange (right)

Chapter 3

⁶² Comparing Algorithms

With the norMmix package explained, we can turn to comparing it to existing methods.

As previously stated, the implementation representing the EM-algorithm is the mclust
package. It will be used with very little deviation from out-of-the-box, safe for restriction
of the covariance models. This is done, so we can compare like with like. The specific
command that performs the EM-algorithm is:

> #mclust::Mclust(x, G=cl, modelNames=mo)\$BIC

Where cl is a vector of integers of however many components we are trying to fit and mo are the model names:

The \$BIC element of the results is taken as the main tool for model selection, as it is advertised in the package authors paper Scrucca et al. (2016).

There is however a small but crucial change applied to these results. The mclust package authors have flipped the definition of the BIC to mean:

$$2ln(\hat{L}) - ln(n)\theta$$

274 instead of the more common

$$ln(n)\theta - 2ln(\hat{L})$$

Where n is the number of observations, θ is the cardinality of the parameter vector and \hat{L} is the estimated log-likelihood.

277 So even if not explicitly mentioned, we use the negative of the values returned by mclust.

Another thing that should be stated before all else is the difference in initialization between mclust's pre-clustering and CLARA. CLARA is dependent on random number generators (RNG). As such, unless a fixed seed is chosen, every iteration of CLARA will return a different result. Unlike mclust, which will, for given data, always return the same results.

The effect on the following findings is that results will spread out for data obtained from CLARA results.

First, we illustrate the structure of the graphical results we will be presenting hereafter.
The basic shape of the plots will be the BIC value plotted against the number of components. This is in line with mclust's manner of visualizing data, however since our method

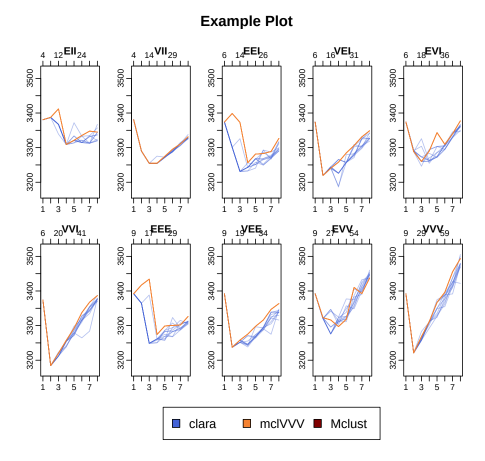


Figure 3.1: Example of Comparison Plot

is to some extent RNG dependent, we are forced to display multiple runs of the algorithm on the same graph. Therefore we split the plot according to covariance model, putting 10 models in 10 graphs in a plot. Here an example:

As can be seen from the formula of the BIC value, lower is better. When selecting a model based on BIC, we take the model and component with the lowest value to be the best fitting model. Although this may not necessarily the 'correct' model, that is, the model from which the data arises.

There are many ways in which this type of model selection might miss the correct model, for example by 'gluing together' multiple components into one, or covering the dataset in a 'patchwork' of smaller components, to name a few.

297 We will discuss them as they arise in the following analysis of simulations

here explain simulations conducted, A.2 here explain the various sections: time, n, p, difficult, nonnormal

3.0 3.1 Time Analysis

here how much time they take, in p,k and n give approximate O(x) value

3.2 Behaviour in n 17

```
library(norMmix, lib.loc="~/ethz/BA/norMmix.Rcheck/")
      # change this dir to whereever the simulations are saved
>
      mainsav <- normalizePath("~/ethz/BA/Rscripts/")</pre>
      savdir <- file.path(mainsav, "2time")</pre>
      filelist <- list.files(savdir, pattern=".rds")</pre>
>
>
      filelist <- grep("mcl.rds", filelist, invert=TRUE, value=TRUE)</pre>
>
      ## need to split these better
>
      f <- lapply(file.path(savdir,filelist), function(j) readRDS(j)$fit)</pre>
      times <- unlist(lapply(f, function(j) extracttimes(j)[,,1]))</pre>
      dims <- unlist(lapply(f, function(j) attr(extracttimes(j), "p")))</pre>
      size <- unlist(lapply(f, function(j) attr(extracttimes(j), "n")))</pre>
>
>
      ddims <- rep(dims, each=80)</pre>
      ssize <- rep(size, each=80)
>
      pars <- unlist(lapply(f, npar))</pre>
      r <- lm(log(times) ~ log(pars) + log(ddims) + log(ssize))
      summary(r)
Call:
lm(formula = log(times) ~ log(pars) + log(ddims) + log(ssize))
Residuals:
             1Q Median
                              3Q
    Min
                                      Max
-3.4428 -0.2986 0.0671 0.4579 2.0936
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
                         0.10598 -91.91 <2e-16 ***
(Intercept) -9.74133
log(pars)
             2.75983
                         0.01181 233.75
                                            <2e-16 ***
log(ddims)
            -2.06063
                         0.02483 -82.99
                                            <2e-16 ***
log(ssize)
             0.61301
                         0.01446
                                  42.38
                                            <2e-16 ***
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
Residual standard error: 0.6946 on 7196 degrees of freedom
Multiple R-squared: 0.8887,
                                     Adjusted R-squared: 0.8887
F-statistic: 1.916e+04 on 3 and 7196 DF, p-value: < 2.2e-16
can see that time is almost one to one proportional to parameter length.
```

$_{\scriptscriptstyle 03}$ 3.2 Behaviour in $_{\scriptscriptstyle 1}$

here show as expected narrower scattering as n increases

305 [h]

$_{ imes}$ 3.3 Behaviour in p

here show how norMmix is consistently competitive with mclust

```
> plot(times~pars, log="xy", yaxt="n", xaxt="n")
> sfsmisc::eaxis(1)
> sfsmisc::eaxis(2)
```

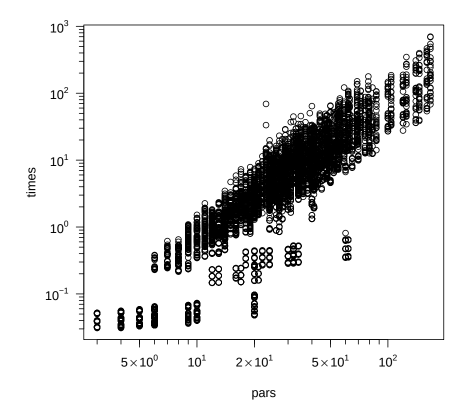


Figure 3.2: Log-log Plot of System Time against Parameter Length

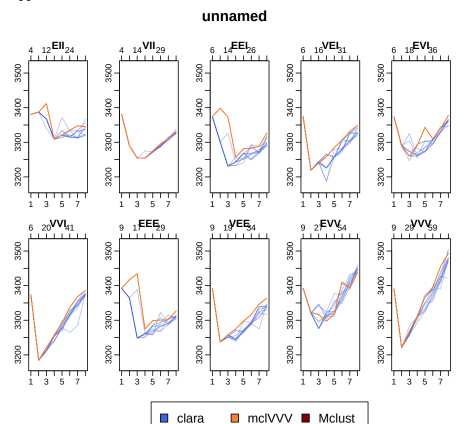
3.4 Diffixult Mixtures

309 here show behaviour in difficult cases

```
> savdir <- file.path(mainsav, "2init")
```

- > filenames <- list.files(savdir, pattern=".rds")</pre>
- > MW214fn <- grep("MW214", filenames, value="TRUE")
- > mclustfiles <- grep("mcl.rds", MW214fn, value=TRUE)
- > MW214fn <- grep("mcl.rds", MW214fn, value="TRUE", invert=TRUE)
- > claraMW <- grep("clara", MW214fn, value=TRUE)</pre>
- > mclMW <- grep("mclVVV", MW214fn, value=TRUE)
- > clarabic <- massbic(claraMW, savdir)</pre>
- > mclbic <- massbic(mclMW, savdir)</pre>
- > mclustbic <- readRDS(file.path(savdir,mclustfiles[1]))</pre>
- here some examples of fitted mixtures
- We can see, that, subtracting the obvious hiccups of the small erroneous components,
- 312 norMmix has correctly found the 'intended' distribution. This is remarkable, given the

> compplot(s05mw34bic, m0534)



small sample size and difficulty of distribution

3.5 Nonnormal Mixtures

 $_{\rm 315}$ here 2smi and 2var, maybe others as well.

316 here 2smi:

- > savdir <- file.path(mainsav, "2smi")</pre>
- > filenames <- list.files(savdir, pattern=".rds")</pre>
- > fnclara <- grep("clara_seed", filenames, value=TRUE)</pre>
- > fnmclVV <- grep("mclVVV_see", filenames, value=TRUE)</pre>
- > fnmclus <- grep("__mcl.rds", filenames, value=TRUE)</pre>

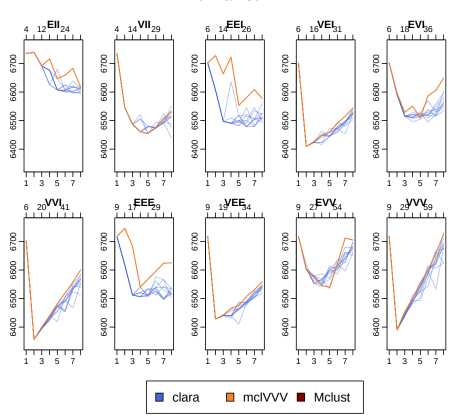
While not very spectacular, the graphs show that even at large parameter counts our algorithm closes in on the same values as mclust. At these dimensions it is difficult to compare if these are actually equal, or even similar fits, but going by BIC values, it is at the very least equally viable as a working model.

To illustrate, here are the parameter sizes for this simulation:

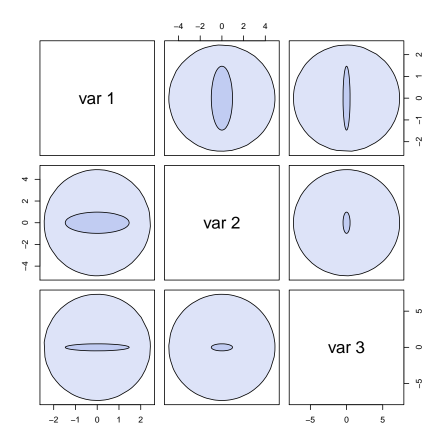
```
EII VII EEI VEI EVI VVI EEE VEE EVV VVV
1 21 21 40 40 40 40 230 230 230 230
```

> compplot(s10mw34bic, m1034)

unnamed



> plot(MW34)



```
      2
      42
      43
      61
      62
      80
      81
      251
      252
      460
      461

      3
      63
      65
      82
      84
      120
      122
      272
      274
      690
      692

      4
      84
      87
      103
      106
      160
      163
      293
      296
      920
      923

      5
      105
      109
      124
      128
      200
      204
      314
      318
      1150
      1154

      6
      126
      131
      145
      150
      240
      245
      335
      340
      1380
      1385

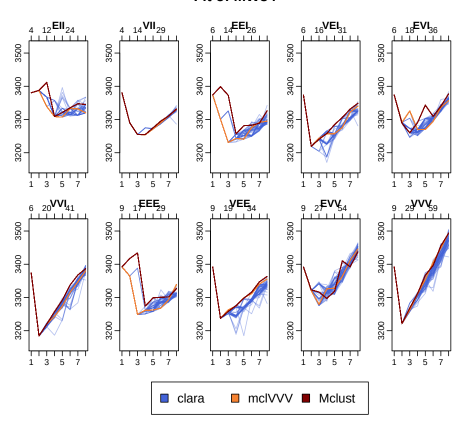
      7
      147
      153
      166
      172
      280
      286
      356
      362
      1610
      1616

      8
      168
      175
      187
      194
      320
      327
      377
      384
      1840
      1847
```

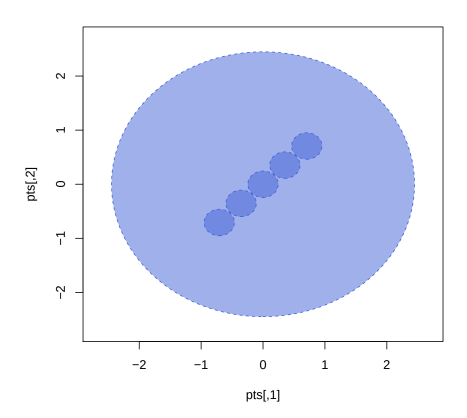
>

compplot(clarabic, mclbic, mclustbic, main="Fit of MW34")

Fit of MW34

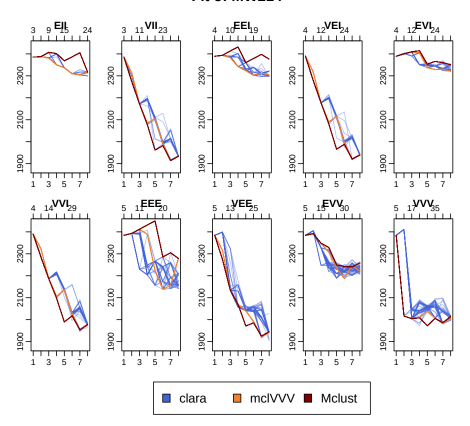


> plot(MW214)

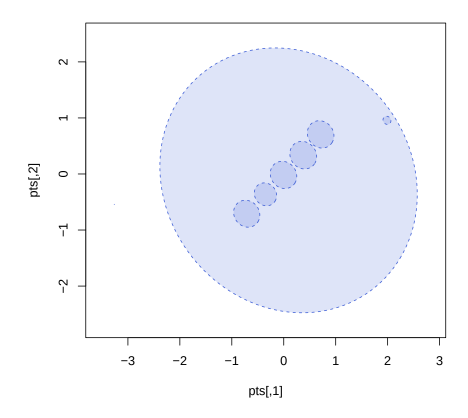


> compplot(clarabic, mclbic, mclustbic, main="Fit of MW214")

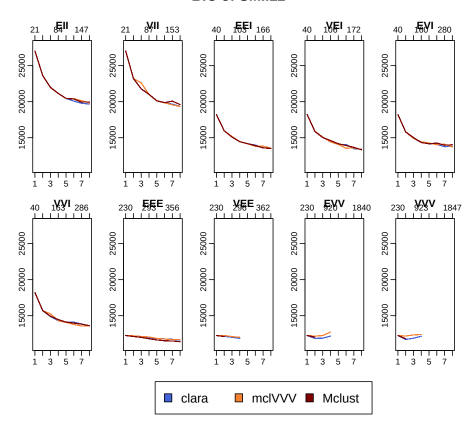
Fit of MW214



- > f <- readRDS(file.path(savdir, claraMW[28]))</pre>
- > ff <- f\$fit\$nMm[8,8][[1]]
- > plot(ff\$norMmix)
- > #points(ff\$x)



BIC of SMI.12



Chapter 4

₂₃ Discussion

- one shortcoming is time inefficiency. largely due to implementation. mclust has 16'000 lines of Fortran code, impossible in the scope of this thesis.
- proof of concept?? definitely possible to do model selection using a general optimizer.
- 327 strong points: 'randomness' of clara/optim allows 'confidence intervalls' for selected model
- $_{328}$ flexibility of approach: given an logLik fctn can do mixture fitting w/ arbitrary models
- further study might include: other presumed component distributions, 'high' dimensions
- failures of implementation: no lower boundary for variance, can lead to minuscule components not as bad as it used to be.

28 Discussion

32 Bibliography

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 New York: Springer. ISBN 0-387-95457-0.

364 Appendix A

R Code

$_{ extsf{366}}$ $\mathbf{A.1}$ llnorMmix

```
Here llnorMmix, since it is the central piece of the package, and 2time.R as an example of a simulation script.
```

```
#### the llnorMmix function, calculating log likelihood for a given
#### parameter vector
## Author: Nicolas Trutmann 2019-07-06
## Log-likelihood of parameter vector given data
# par:
         parameter vector
# tx:
         transposed sample matrix
         number of components
# model: assumed distribution model of normal mixture
# trafo: either centered log ratio or logit
llnorMmix <- function(par, tx, k,</pre>
                       trafo=c("clr1", "logit"),
                       model=c("EII","VII","EEI","VEI","EVI",
                                "VVI", "EEE", "VEE", "EVV", "VVV")
                       ) {
    stopifnot(is.matrix(tx),
               length(k \leftarrow as.integer(k)) == 1, k >= 1)
    p <- nrow(tx)
    x \leftarrow t(x) ## then only needed in
                                          (x-mu[,i])^2 i=1..k
    # 2. transform
    model <- match.arg(model)</pre>
    trafo <- match.arg(trafo)</pre>
    12pi <- log(2*pi)
    # 3. calc log-lik
    # get w
    w \leftarrow if (k==1) 1
         else switch(trafo,
                      "clr1" = clr1inv (par[1:(k-1)]),
                      "logit"= logitinv(par[1:(k-1)]),
```

32 R Code

```
stop("invalid 'trafo': ", trafo)
# start of relevant parameters:
f \leftarrow k + p*k \# weights -1 + means +1 => start of alpha
mu <- matrix(par[k:(f-1L)], p,k)</pre>
f1 <- f
             # end of alpha if uniform
f2 \leftarrow f+k-1L \# end of alpha if var
f1.1 <- f1 +1L # start of D. if alpha unif.
f2.1 <- f1 + k # start of D. if alpha variable
f11 <- f1 + p-1
                   # end of D. if D. uniform and alpha uniform
f12 <- f1 + (p-1)*k # end
                             D. if D.
                                         var
                                               and alpha unif.
f21 <- f2 + p-1
                  # end of D. if D. uniform and alpha variable
f22 \leftarrow f2 + (p-1)*k \# end of D. if D.
                                               and alpha var.
                                         var
f11.1 <- f11 +1L # start of L if alpha unif D unif
f21.1 <- f21 +1L # start of L if alpha var
f12.1 <- f12 +1L # start of L if alpha unif D var
f22.1 <- f22 +1L # start of L if alpha var
                                               D var
f111 \leftarrow f11 + p*(p-1)/2 \# end of L if alpha unif D unif
                p*(p-1)/2 # end of L if alpha var
f211 <- f21 +
f121 <- f12 + k*p*(p-1)/2 # end of L if alpha unif D var
f221 \leftarrow f22 + k*p*(p-1)/2 \# end of L if alpha var
# initialize f(tx_i) i=1..n vector of density values
invl <- 0
# calculate log-lik, see first case for explanation
switch(model,
"EII" = {
    alpha <- par[f]
    invalpha \leftarrow exp(-alpha) = 1/exp(alpha)
    for (i in 1:k) {
        rss <- colSums(invalpha*(tx-mu[,i])^2)
        # this is vector of length n=sample size
        # calculates (tx-mu)t * Sigma^-1 * (tx-mu) for diagonal
        # cases.
        invl \leftarrow invl+w[i]*exp(-0.5*(p*(alpha+l2pi)+rss))
        # adds likelihood of one component to invl
        # the formula in exp() is the log of likelihood
        # still of length n
    }
},
# hereafter differences are difference in dimension in alpha and D.
# alpha / alpha[i] and D. / D.[,i]
\"\!\" = {
    alpha <- par[f:f2]</pre>
    for (i in 1:k) {
        rss <- colSums((tx-mu[,i])^2/exp(alpha[i]))
        invl <- invl+w[i]*exp(-0.5*(p*(alpha[i]+l2pi)+rss))</pre>
    }
},
```

A.1 llnorMmix

```
"EEI" = {
    alpha <- par[f]
    D. <- par[f1.1:f11]</pre>
    D. \leftarrow c(-sum(D.), D.)
    D. \leftarrow D.-sum(D.)/p
    invD <- exp(alpha+D.)</pre>
    for (i in 1:k) {
        rss <- colSums((tx-mu[,i])^2/invD)
         invl <- invl+w[i]*exp(-0.5*(p*(alpha+l2pi)+rss))</pre>
    }
},
"VEI" = {
    alpha <- par[f:f2]
    D. <- par[f2.1:f21]</pre>
    D. <- c(-sum(D.), D.)
    D. \leftarrow D.-sum(D.)/p
    for (i in 1:k) {
        rss <- colSums((tx-mu[,i])^2/exp(alpha[i]+D.))
         invl <- invl+w[i]*exp(-0.5*(p*(alpha[i]+l2pi)+rss))</pre>
    }
},
"EVI" = {
    alpha <- par[f]
    D. <- matrix(par[f1.1:f12],p-1,k)</pre>
    D. \leftarrow apply(D.,2, function(j) c(-sum(j), j))
    D. <- apply(D.,2, function(j) j-sum(j)/p)</pre>
    for (i in 1:k) {
        rss <- colSums((tx-mu[,i])^2/exp(alpha+D.[,i]))
         invl \leftarrow invl+w[i]*exp(-0.5*(p*(alpha+l2pi)+rss))
    }
},
"VVI" = {
    alpha <- par[f:f2]
    D. <- matrix(par[f2.1:f22],p-1,k)</pre>
    D. \leftarrow apply(D.,2, function(j) c(-sum(j), j))
    D. \leftarrow apply(D.,2, function(j) j-sum(j)/p)
    for (i in 1:k) {
        rss <- colSums((tx-mu[,i])^2/exp(alpha[i]+D.[,i]))
         invl \leftarrow invl+w[i]*exp(-0.5*(p*(alpha[i]+l2pi)+rss))
    }
},
# here start the non-diagonal cases. main difference is the use
# of backsolve() to calculate tx^t Sigma^-1 tx, works as follows:
# assume Sigma = L D L^t, then Sigma^-1 = (L^t)^-1 D^-1 L^-1
# y = L^-1 tx => tx^t Sigma^-1 tx = y^t D^-1 y
# y = backsolve(L., tx)
"EEE" = {
    alpha <- par[f]
    D. <- par[f1.1:f11]</pre>
```

34 R Code

D. <- c(-sum(D.), D.)

```
D. \leftarrow D.-sum(D./p)
    invD <- exp(alpha+D.)</pre>
    L. \leftarrow diag(1,p)
    L.[lower.tri(L., diag=FALSE)] <- par[f11.1:f111]</pre>
    for (i in 1:k) {
         rss <- colSums(backsolve(L.,(tx-mu[,i]), upper.tri=FALSE)^2/invD)
         invl <- invl+w[i]*exp(-0.5*(p*(alpha+12pi)+rss))</pre>
    }
},
"VEE" = {
    alpha <- par[f:f2]
    D. <- par[f2.1:f21]</pre>
    D. <- c(-sum(D.), D.)
    D. \leftarrow D.-sum(D./p)
    L. \leftarrow diag(1,p)
    L.[lower.tri(L., diag=FALSE)] <- par[f21.1:f211]
    for (i in 1:k) {
         rss <- colSums(backsolve(L., (tx-mu[,i]), upper.tri=FALSE)^2/exp(alpha[i]+D
         invl <- invl+w[i]*exp(-0.5*(p*(alpha[i]+l2pi)+rss))</pre>
},
"EVV" = {
    alpha <- par[f]
    D. <- matrix(par[f1.1:f12],p-1,k)</pre>
    D. \leftarrow apply(D.,2, function(j) c(-sum(j), j))
    D. \leftarrow apply(D.,2, function(j) j-sum(j)/p)
    L.temp <- matrix(par[f12.1:f121],p*(p-1)/2,k)
    for (i in 1:k) {
        L. <- diag(1,p)
         L.[lower.tri(L., diag=FALSE)] <- L.temp[,i]</pre>
         rss <- colSums(backsolve(L., (tx-mu[,i]), upper.tri=FALSE)^2/exp(alpha+D.[,
         invl \leftarrow invl+w[i]*exp(-0.5*(p*(alpha+l2pi)+rss))
    }
},
"VVV" = {
    alpha <- par[f:f2]</pre>
    D. <- matrix(par[f2.1:f22],p-1,k)</pre>
    D. \leftarrow apply(D.,2, function(j) c(-sum(j), j))
    D. <- apply(D.,2, function(j) j-sum(j)/p)</pre>
    invalpha <- exp(rep(alpha, each=p)+D.)
    L.temp <- matrix(par[f22.1:f221],p*(p-1)/2,k)
    L. \leftarrow diag(1,p)
    for (i in 1:k) {
         L.[lower.tri(L., diag=FALSE)] <- L.temp[,i]</pre>
         rss <- colSums(backsolve(L., (tx-mu[,i]), upper.tri=FALSE)^2/invalpha[,i])
         invl <- invl+w[i]*exp(-0.5*(p*(alpha[i]+l2pi)+rss))</pre>
    }
},
```

A.1 llnorMmix 35

```
## otherwise
    stop("invalid model:", model)
    ## return sum_{i=1}^n log( f(tx_i) ) :
    sum(log(invl))
sllnorMmix <- function(x, obj, trafo=c("clr1", "logit")) {</pre>
    stopifnot(is.character(model <- obj$model))</pre>
    trafo <- match.arg(trafo)</pre>
    llnorMmix(nMm2par(obj, model=model),
               tx = t(x), k = obj$k,
               model=model, trafo=trafo)
}
## log-likelihood function relying on mvtnorm function
        parameter vector as calculated by nMm2par
# par:
# x:
         matrix of samples
         number of cluster
# k:
# trafo: transformation of weights
# model: assumed model of the distribution
llmvtnorm <- function(par, x, k,</pre>
                       trafo=c("clr1", "logit"),
                       model=c("EII","VII","EEI","VEI","EVI",
                                "VVI", "EEE", "VEE", "EVV", "VVV")
               ) {
    stopifnot(is.matrix(x),
               length(k \leftarrow as.integer(k)) == 1, k >= 1)
    model <- match.arg(model)</pre>
    trafo <- match.arg(trafo)</pre>
    p \leftarrow ncol(x)
    nmm <- par2nMm(par, p, k, model=model, trafo=trafo)</pre>
    ## FIXME (speed!): dmvnorm(*, sigma= S) will do a chol(S) for each component
    ## ---- *instead* we already have LDL' and chol(S) = sqrt(D) L' !!
    ## another par2*() function should give L and D, or from that chol(Sagma), rather than S
    w <- nmm$w
    mu <- nmm$mu
    sig <- nmm$Sigma
    y <- 0
    for (i in 1:k) {
        y <- y + w[i]*mvtnorm::dmvnorm(x,mean=mu[,i],sigma=sig[,,i])</pre>
    sum(log(y))
}
```

36 R Code

369 A.2 Example Simulation Script

```
here e.g. 2init.R and write some remarks on it.
 ## Intent: analyse time as function of p,k,n
 nmmdir <- normalizePath("~/BachelorArbeit/norMmix.Rcheck/")</pre>
 savdir <- normalizePath("~/BachelorArbeit/Rscripts/2time")</pre>
 stopifnot(dir.exists(nmmdir), dir.exists(savdir))
 library(norMmix, lib.loc=nmmdir)
 library(mclust)
 ## at n=500,p=2 can do about 250xfitnMm(x,1:10) in 24h
 seeds <- 1:10
 sizes <- c(500, 1000, 2000)
nmm <- list(MW214, MW34, MW51)
 ## => about 100 cases
 # for naming purposes
 nmnames <- c("MW214", "MW34", "MW51")
 sizenames <- c("500", "1000", "2000")
 files <- vector(mode="character")</pre>
 for (nm in 1:3) {
     for (size in sizes) {
     set.seed(2019); x <- rnorMmix(size, nmm[[nm]])</pre>
         for (seed in seeds) {
             set.seed(2019+seed)
             r <- tryCatch(fitnMm(x, k=1:8,
                                    optREPORT=1e4, maxit=1e4),
                            error = identity)
             filename <- sprintf("%s_size=%0.4d_seed=%0.2d.rds",
                                   nmnames[nm], size, seed)
             files <- append(files, filename)</pre>
              cat("===> saving to file:", filename, "\n")
              saveRDS(list(fit=r), file=file.path(savdir, filename))
         }
     }
 }
 fillis <- list()</pre>
 for (i in seq_along(sizes)) {
     for (j in seq_along(nmnames)) {
         # for lack of AND matching, OR match everything else and invert
         ret <- grep(paste(sizenames[-i], nmnames[-j], sep="|"),</pre>
                      files, value=TRUE, invert=TRUE)
         fillis[[paste0(sizenames[i], nmnames[j])]] <- ret</pre>
     }
 }
 epfl(fillis, savdir)
```

371 Appendix B

Further Plots

373 here further plots:

B.1 Chapter 3

not nec best way to go about it, section differently.

dsfasdf

38 Further Plots

Declaration of Originality

The signed declaration of originality is a component of every semester paper, Bachelor's thesis, Master's thesis and any other degree paper undertaken during the course of studies, including the respective electronic versions.

Lecturers may also require a declaration of originality for other written papers compiled for

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