

Seminar for Statistics

Department of Mathematics						
Bachelor Thesis	Winter 2019					
Dachelor Thesis	Willier 2019					

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Comparison of EM-algorithm and MLE using Cholesky decomposition

Submission Date: placeholder

Advisor: placeholder

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)

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Chapter 1

Introduction to normal mixture models

4 1.1 Definitions

- 5 A good and thorough introductory book is the work of McLachlan and Peel (2000) and
- 6 the reader is encouraged to study it to learn in depth about normal mixtures and their
- 7 clustering. We will here give a short overwiev of normal mixtures to fix notation and
- 8 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. The original example
- of this, by Karl Pearson, who fitted two normal distributions with different means and
- variances. In his paper Pearson analyzed measurements of forehead to bodylength of
- crabs sampled from the bay of Naples. His mixture model-based approach suggested, that
- a the crabs were evolving into two new subspecies.
- here make clear we restrict to multivariate normal case.
- Let $\mu \in \mathbb{R}^p$, $\Sigma \in \mathbb{R}^{p \times p}$ symmetric positive definite and $\phi(-; \mu, \Sigma)$ be the normal distri-
- bution with mean μ and covariance matrix Σ .
- Y_1, \dots, Y_n
- Definition 1.1.0.1. Suppose we have a random sample Y_1, \ldots, Y_n with probability density
- 19 function $Y_j \sim f(y_j)$ on \mathbb{R}^p We assume that the density $f(y_j)$ of Y_j can be written in the
- 20 form

$$f(y_j) = \sum_{k=1}^{K} \pi_k \phi_k(y_k; \mu, \Sigma)$$

- The π_k are called the component densities of the mixture and the ϕ_k mixture components.
- 22 here small note about restricted cases. give ref to 1.4??

1.2 The EM-algorithm in scetch

- 24 With this definition we immediately face the problem of how to fit these mixture com-
- ponents to given data. A popular algorithm to solve this problem is the Expectation-
- Maximization algorithm, abbreviated as EM-algorithm.

37

We give here a scetch of the EM-algorithm in the case of all normal mixture components, since it is the scope of this thesis and simplifies it considerably.

Suppose we have a p-dimensional dataset of n samples x_1, \ldots, x_n , onto which we would like to fit K normal distributions ϕ_k , $k \in 1, \ldots, n$. We introduce a further explaining variable \mathbf{Z} in $\mathrm{Mat}^{n \times k}$, with entries in [0,1] which represent the expectation that observation i belongs to component k.

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.

$$\tau_i(y_j; \Psi) = \phi_i(y_j; \mu_i, \Sigma_i) / \sum_{k=1}^K \phi_k(y_j; \mu_k, \Sigma_k)$$

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

$$\mu_i = \sum_{j=1}^n \tau_{ij} y_j / \sum_{j=1}^n \tau_{ij}$$

 $\Sigma_{i} = \sum_{j=1}^{n} \tau_{ij} (y_{j} - \mu_{i}) (y_{j} - \mu_{i})^{\top} / \sum_{j=1}^{n} \tau_{ij}$

38 here note about initialization methods.

$_{ ext{\tiny 39}}$ 1.3 choice of notation

The classification of models in this paper relies heavily on the work of Celeux and Grovaert, 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 and ours.

The basis of classification in Celeux & Grovaert 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

$$\Sigma = \lambda \mathbf{D} \mathbf{A} \mathbf{D}^{\mathsf{T}}$$

with \boldsymbol{D} an orthogonal matrix and \boldsymbol{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 arbytrary 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 reffering to the decomposition as described by cng, will use the following modification of notation:

$$\begin{aligned} \boldsymbol{D} &\longmapsto \boldsymbol{Q} \\ \boldsymbol{A} &\longmapsto \boldsymbol{\Lambda} \\ \boldsymbol{\lambda} &\longmapsto \boldsymbol{\alpha} \\ \boldsymbol{\Sigma} &= \boldsymbol{\lambda} \boldsymbol{D} \boldsymbol{A} \boldsymbol{D}^\top = \boldsymbol{\alpha} \boldsymbol{Q} \boldsymbol{\Lambda} \boldsymbol{Q}^\top \end{aligned}$$

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 eigen vectors and α was somewhat arbitrarily chosen.

50 1.4 Models of Covariance Matrices

make clear that the models can not be translated one to one to ldlt model There is howerver an issue with the Cholesky decomposition. For 10 out of 14 cases as defined by Celeux & Grovaert, 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:

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

Since in both cases the bracketing matrices Q and L have determinant 1 the determinant of Σ falls entirely on α . Therefore α , in these particular decompositions, is equal for both. Celeux & Grovaert vary Σ by either varying or holding fixed the volume (α/α_k) , shape (Λ/Λ_k) and orientation (Q/Q_k) . These 3 times 2 cases would yield the 8 out of 14 cases of non-diagonal cases. However there is no canonical transform for either variable orientation and fixed shape or fixed orientation and variable shape. The reason for this is that in the LDL^{\top} decomposition the lower diagonal matrix L holds some of the shape of the matrix, which in the Eigen decomposition is in the Λ matrix. In fact, L is orthogonal 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 $\boldsymbol{L}_k\boldsymbol{D}\boldsymbol{L}^{\top}$, however they do not correspond to the desired geometric intent behind the differentiation of models and are therefore not included.

make nice table(maybe sideways to account for parameter list)

count													$1 + pK + K \frac{p(p-1)}{2}$	$K + pK + K\frac{p(p-1)}{2}$
parameters													$\lambda, d_{i,k}, l_{i,j,k} \ j > i \qquad 1 + pK + K \frac{p(p-1)}{2}$	$\lambda_k, d_{i,k}, l_{i,j,k} \ j > i K + pK + K \frac{p(p-1)}{2}$
TDT	same as $C\&G$						$lpha m{T}m{D}m{L}^{ op}$	doesn't exist	$\alpha_k \boldsymbol{LDL}^\top$		don't exist		$lpha oldsymbol{L}_k oldsymbol{D}_k oldsymbol{L}_k^ op$	$lpha_koldsymbol{L}_koldsymbol{D}_koldsymbol{L}_k^ op$
count	1	K	1+p	K + p	1 + pK	K + pK	$1 + p + p^2$	$1 + pK + p^2$	$K + p + p^2$	$K + pK + p^2$	$1 + p + Kp^2$	$K + p + Kp^2$	$1 + pK + Kp^2$	$\alpha_k, \lambda_i, q_{i,j,k} K + pK + Kp^2$
parameters	α	α_k	$lpha, \lambda_i$	$lpha_k, \lambda_i$	$lpha, \lambda_{i,k}$	$lpha_k, \lambda_{i,k}$	$\alpha, \lambda_i, q_{i,j}$	$lpha,\lambda_{i,k},q_{i,j}$	$\alpha_k, \lambda_i, q_{i,j}$	$lpha_k, \lambda_{i,k}, q_{i,j}$	$lpha, \lambda_i, q_{i,j,k}$	$lpha_k, \lambda_i, q_{i,j,k}$	$lpha, \lambda_i, q_{i,j,k}$	$lpha_k, \lambda_i, q_{i,j,k}$
orientation	ı	ı	coordinate axes	coordinate axes	coordinate axes	coordinate axes	equal	ednal	equal	ednal	variable	variable	variable	variable
$_{ m shape}$	equal	ednal	ednal	ednal	variable	variable	ednal	variable	ednal	variable	ednal	equal	variable	variable
volume	equal	variable	equal	variable	equal	variable	equal	equal	variable	variable	equal	variable	equal	variable
$\mathbf{\Sigma}_k$ C&G	αI	$lpha_k m{I}$	$\Delta \Delta$	$lpha_k \mathbf{A}$	$lpha {f \Lambda}_k$	$lpha_k \mathbf{\Lambda}_k$	$lpha oldsymbol{Q} oldsymbol{V} oldsymbol{Q}_{\perp}$	$lpha oldsymbol{Q} oldsymbol{\Lambda}_k oldsymbol{Q}^{ op}$	$lpha_k oldsymbol{Q} oldsymbol{\Lambda} oldsymbol{Q}^{ op}$	$lpha_k oldsymbol{Q} oldsymbol{\Lambda}_k oldsymbol{Q}^{ op}$	$lpha oldsymbol{Q}_k oldsymbol{\Lambda} oldsymbol{Q}_k^ op$	$lpha_k oldsymbol{Q}_k oldsymbol{\Lambda} oldsymbol{Q}_k^ op$	$lpha oldsymbol{Q}_k oldsymbol{\Lambda}_k oldsymbol{Q}_k^ op$	$lpha_k oldsymbol{Q}_k oldsymbol{\Lambda}_k oldsymbol{Q}_k^ op$
Model	EII	VII	EEI	VEI	EVI	VVI	EEE	EVE	VEE	AVE	EEV	VEV	EVV	VVV

$_{ iny 50}$ 1.5 problems of EM

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]. In common software implementations, The concern here is that a slowing in convergence might be mistaken for actual convergence.

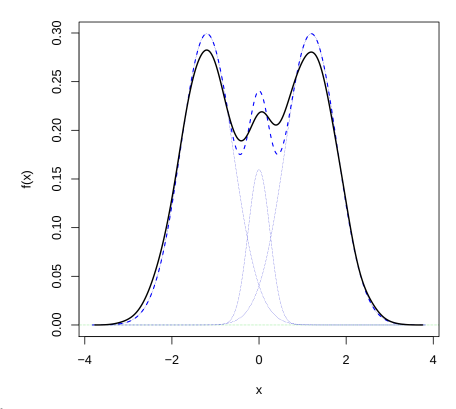
This phenomenon is not infrequent and in difficult mixtures quite visible. To illustrate let us look at a particular mixture taken from Marron and Wand (1992) and the nor1mix package from CRAN.

[2,] 1.2 0.60 0.45

[3,] 0.0 0.25 0.10

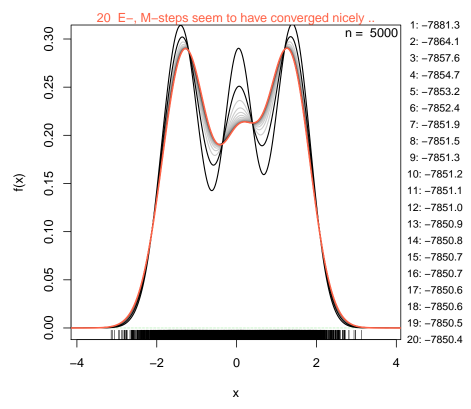
91 show an example using nor1mix

#9 Trimodal



93 then an illustration of MW examples of pathological cases

EM for 1-dim normal mixtures



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- yay, got figure to print. solution was use of fig=TRUE, instead of various mutations like figure=true.
- here we see how change in loglik seems to stagnate. However, this does not stay that way, if we let EM run a bit further.
- to conclude example show part of mixest that shows it takes 1200 iterations to converge
- In fact, it seems that the previous solution is a saddle point in the likelihood function, where EM has chronic problems continuing improvements.
- should include animations?? like mix_est_1d.R line 249+24 lines
- maybe show Marr Wand's examples of 'difficult' mixtures
- give conclusion recapping the just demonstrated, and lead in for next chapter

EM for 1-dim normal mixtures

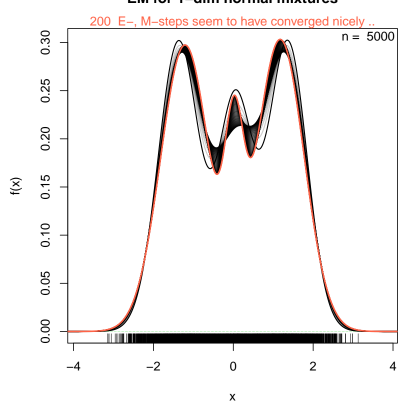


Figure 1.1: 200 EM steps

Chapter 2

$^{_{ imes}}$ ${ m The}$ norMmix ${ m Package}$

- explain, that this package was written purposefully for this paper.
- The norMmix package is constructed around the norMmix object, that codifies a normal
- Multivariate mixture model, and the llnorMmix() function.
- 110 quickly list contents of norMmix object
- relies on optim() generic optimizer. maximizes llnormix by varying model parameters.
- 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.
- 114 Conceptually, at the start of a fitting algo, e.g. EM we need to initialize a mixture object.
- thereafter the paths diverge. at the heart of norMmix's functionality lie the functions:
- 116 llnorMmix and nMm2par which are in turn employed by norMmixMLE to funnel a mixture
- object into optim and give optim a function to optimize.
- also relies on mixtools package for random generating function rnorMmix using rmvnorm.

1.1 concept of package

- (this Section maybe one chapter earlier)
- about Cholesky decomp as ldlt. has advantages: fast, parametrically parsimonious, can easily compute loglikelihood
- maybe reread section in McLachlan about accelerating EM algo
- 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.
- So caomparison of algos done through throwing difficult mixtures and non-mixtures at it
- and hoping that norMmix finds better solutions than EM. So the criteria for "better fit"
- are 1. better log-likelihood 2. correct model, where EM fails.

2.2 finer details of norMmix package

Chapter 3

32 Comparing Algorithms

3.1 General Setup

- display abilities of norMmix on its own. can find correct models
- 135 Mention, that mclust doesn't depend on seed(double check) and therefore norMmix has
- advantage of 'confidence intervals'. We can run 50 simulations and see if there might be
- more sensible clusters.
- maybe apply to MW[0-9] objects?
- 139 not sure
- as in Raftery 2002, Benaglia 2009, Roeder 1997, maybe compare to MISE of various forms.
- They all did and see it as adequate method for comparing accuracy of algorithm.
- also wanted is accuracy of model selection. generate from model and then compare fitted
- to original. either by acc-model==fit-model and acc-k==fit-k or acc-ll fit-ll.

$_{144}$ 3.2 Findings

- Chapter 4
- Discussion

14 Discussion

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