Finite Mixture Modeling via REBMIX

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Abstract- The REBMIX algorithm for fitting finite mixture models implemented in R package rebmix is presented. It provides functions for random univariate and multivariate finite mixture generation, the number of components, component weights and component parameter estimation, bootstrapping and the plotting of finite mixtures. It requires preprocessing of observations, information criterion and conditionally independent normal, lognormal, Weibull, gamma, binomial, Poisson or Dirac component densities. The algorithm optimizes the component parameters, mixing weights and number of components successively based on boundary conditions, such as the maximum number of components and number of bins or nearest neighbours. The algorithm is robust, time efficient and can be used either to assess an initial set of unknown parameters and number of components, e.g., for the EM algorithm, or as a standalone algorithm providing a good compromise between parametric and nonparametric methods of finite mixture estimation. Univariate and multivariate datasets are analysed for validation purposes.

Keywords- Continuous Variable; Discrete Variable; Mixture Estimation; R Package

I. INTRODUCTION

Finite mixture models are used increasingly to model the distributions of a wide variety of random phenomena. For multivariate data of a continuous nature, attention is paid to the use of multivariate normal components because of their computational convenience [1, 2, 3]. However, in fatigue and reliability analysis, lognormal and Weibull distributions are preferred due to their flexibility and applicability to continuous positive random variables only [4, 5, 6, 7].

The finite mixture models have seen a real boost in popularity over the last two decades due to a tremendous increase in available computing power. These models can be applied to datasets, where observations originate from categories whose affiliations are not known, or it can provide approximations for multimodal distributions [8]. Some of the latest models can be found in [9, 10, 11, 12, 13, 14, 15, 16, 17].

The REBMIX algorithm addresses the problem of finite mixture estimation numerically. It originates in [18] and is based on the assumption that the problem of finite mixture estimation can be broken into multiple problems of parameter estimation for basic parametric family types. Here the governing equations preexist and originate in the maximum likelihood. REBMIX is a sequential algorithm that avoids the following drawbacks of the simultaneous EM algorithm [19, 20]:

- The EM algorithm converges to a local maximum of the likelihood function very quickly.
- There are often several other promising local optimal solutions in the vicinity of the solutions obtained from methods that provide good initial guesses of the solution.
- Model selection criterion usually assumes that the global optimal solution of the log-likelihood function can be obtained. However, achieving this is computationally intractable.
- Some regions in the search space do not contain any promising solutions. The promising and non-promising regions co-exist, and it often becomes challenging to avoid wasting computational resources to search in non-promising regions.

Over time REBMIX has evolved [21, 22, 23, 24, 25]. This paper extends it by adding gamma, binomial, Poisson and Dirac parametric families. The reader should expect the derivation of a numerical algorithm, which combines basic statistical techniques in a closed unit and not a concise statistical derivation of the governing equations. The latter does not seem to be feasible.

REBMIX has been compared to the recognizable FlexMix [26] algorithm in [7]. The comparison shows that REBMIX is a robust, time efficient tool that can be used either to assess an initial set of unknown parameters and the number of components for other algorithms [27, 28] or as a standalone procedure that acts as a good compromise between the parametric and nonparametric methods of finite mixture estimation. The rebmix implementation of REBMIX [29] extends the set of algorithms available for random univariate and multivariate finite mixture generation, number of components, component weights and component parameter estimation, bootstrapping and plotting of the finite mixtures in the R language and environment for statistical computing [30].
II. ALGORITHM

Let \( y_1, \ldots, y_n \) be an observed \( d \) dimensional dataset of size \( n \) of continuous or discrete vector observations \( y_j \). Each observation is assumed to follow the predictive mixture density

\[
f(y|c, w, \Theta) = \sum_{l=1}^{c} w_l f(y|\theta_l)
\]

with conditionally independent component densities

\[
f(y|\theta_l) = \prod_{i=1}^{d} f(y_i|\theta_{il})
\]

indexed by the parameter vector \( \theta_l = (\theta_{1l}, \ldots, \theta_{dl})^\top \). The components can currently belong to either normal

\[
f(y_i|\theta_{il}) = \frac{1}{\sqrt{2\pi\sigma_{il}}} \exp \left\{ -\frac{1}{2} \frac{(y_i - \mu_{il})^2}{\sigma_{il}^2} \right\},
\]

lognormal

\[
f(y_i|\theta_{il}) = \frac{1}{\sqrt{2\pi\sigma_{il}y_i}} \exp \left\{ -\frac{1}{2} \frac{(\log(y_i) - \mu_{il})^2}{\sigma_{il}^2} \right\},
\]

Weibull

\[
f(y_i|\theta_{il}) = \frac{\beta_{il}}{\theta_{il}} \left( \frac{y_i}{\theta_{il}} \right)^{\beta_{il}-1} \exp \left\{ -\left( \frac{y_i}{\theta_{il}} \right)^{\beta_{il}} \right\},
\]

gamma

\[
f(y_i|\theta_{il}) = \frac{1}{\Gamma(\beta_{il})y_i} \left( \frac{y_i}{\theta_{il}} \right)^{\beta_{il}} \exp \left\{ -\frac{y_i}{\theta_{il}} \right\},
\]

binomial

\[
f(y_i|\theta_{il}) = \binom{y_i}{\theta_{il}} p_{il}^{y_i} (1 - p_{il})^{\theta_{il} - y_i},
\]

Poisson

\[
f(y_i|\theta_{il}) = e^{-\theta_{il}} \frac{\theta_{il}^{y_i}}{y_i!}
\]

or Dirac

\[
f(y_i|\theta_{il}) = \begin{cases} 1 & y_i = \theta_{il} \\ 0 & \text{otherwise} \end{cases}
\]

parametric family types. The object of the analysis is to obtain the number of components \( c \), component weights \( w_l \) summing to 1 and component parameters \( \theta_l \). The REBMIX algorithm is an iterative numerical procedure which does not rely on rigorous statistical analysis. Instead it makes good use of the suppositions:

- It is always possible to assign empirical densities to an arbitrary dataset.
- Based on the empirical densities, a global mode position can be identified.
- Once the global mode position and its empirical density are known, rough component parameters of the predictive component density can be estimated.
- Based on the rough component parameters, the dataset can be clustered successively into the classes linked to the predictive component densities and the residue.
- The number of components \( c \) equals the number of classes.
- Component parameters and the component weights can be enhanced for all classes.
- The unassigned observations can be distributed between the existing components by the Bayes decision rule and the parameters of the finite mixture can be fine-tuned.

The idea is thus to assign the component densities one after another to the empirical mixture density. At the same time the observations belonging to the component densities are extracted from the dataset. By increasing the number of components, the number of observations in the dataset decreases. When the dataset attains a lower limit, it is assumed that the unassigned observations can be distributed between the existing components. Sections A. to G. give the theoretical backgrounds for the algorithm, while Section H. lists and explains its flow.
A. Preprocessing of Observations

The algorithm requires the preprocessing of observations. Using a histogram, the dataset is counted into a finite number of nonoverlapping, equally sized and regularly distributed bins. Assuming that the mean of a bin \( y_j = (\bar{y}_{i1}, \ldots, \bar{y}_{id})^\top \) is given by

\[
\bar{y}_{ij} = \bar{y}_{i0} + \text{‘An arbitrary integer’} \times h_{ij}, \quad i = 1, \ldots, d, \tag{3}
\]

the fraction of observations \( k_j \) for \( j = 1, \ldots, v \) falling into the volume \( V_j \) is counted out, where \( \bar{y}_{i0} \) stands for an arbitrary origin and \( v \) depicts the number of bins. Similarly, if the Parzen window is employed, the fraction of observations falling into \( V_j \) centered on observation \( y_j \) is obtained. In both cases, the volume is taken to be a hypersquare with sides of length \( h_{ij} \). This yields \( V_j = \prod_{i=1}^{d} h_{ij} \). Moreover, for both approaches bin widths \( h_{ij} = h_i \) and volumes \( V_j = V \) are kept constant. If the \( k \)-nearest neighbour is used, the fraction of observations falling into a normalized hypersphere \( V_j = \pi^{d/2} R_j^d / \Gamma[1 + d/2] \) of radius \( R_j \) centered on observation \( y_j \) contains a constant number \( k_j = k \) of observations. The bin widths for the histogram and Parzen window and continuous parametric families

\[
h_i = \frac{y_{i\text{max}} - y_{i\text{min}}}{v}
\]

depend on the minimum \( y_{i\text{min}} = \min y_{ij} \) and maximum \( y_{i\text{max}} = \max y_{ij} \) observations. For the histogram and continuous parametric families, the origin is preset to

\[
\bar{y}_{i0} = y_{i\text{min}} + \frac{h_i}{2}.
\]

Discrete parametric families require \( h_i = 1 \) and \( \bar{y}_{i0} = y_{i\text{min}} \). The \( k \)-th \(-\) 1 nearest neighbour \( y_j \) is searched around \( y_j \) based on the normalized Euclidean distance

\[
R_j = \sqrt{\sum_{i=1}^{d} \left( \frac{y_{ij} - y_{ij}}{y_{i\text{max}} - y_{i\text{min}}} \right)^2} \quad \text{for } j \neq j \\text{ and } h_{ij} = 2R_j(y_{i\text{max}} - y_{i\text{min}}).
\]

If \( N \geq k \) nearest neighbours coincide, then the normalized Euclidean distance \( R_j \) to the first nearest non-coincident neighbour \( y_j \) is multiplied by \((k/(N + 1))^{1/d}\). Infinite empirical density estimations are thus prevented.

B. Global Mode Detection

The argument \( m \) at which the empirical density \( f_{ij} \) attains its maximum

\[
m = \arg \max_j f_{ij} \tag{4}
\]
determines the global mode. If observations are binned into the histogram, then

\[
f_{ij} = \frac{k_{ij}}{n_l V_j}, \quad j = 1, \ldots, v, \tag{5}
\]

where frequencies \( k_{ij} \) are initially set to \( k_j \) and the number of observations in class \( l \) is

\[
n_l = \sum_{j=1}^{v} k_{ij}.
\]

If the Parzen window or \( k \)-nearest neighbour is applied,

\[
f_{ij} = \frac{k_{ij}}{n_l V_j}, \quad j = 1, \ldots, n. \tag{6}
\]

Frequencies \( k_{ij} \) are initially set to 1, \( n_l = \sum_{j=1}^{n} k_{ij} \) and the component weight \( w_l = n_l / n \). Moreover, the \( l \)-th component conditional empirical density at the global mode for the histogram

\[
f_{i_l|l,m} = \frac{k_{lm}}{\sum_{j=1}^{w} g_{ij} = g_{ilm}} \frac{1}{k_{ij|l,m} h_{im}} = \frac{k_{lm}}{k_{ij|l,m} h_{im}}, \tag{7}
\]

is required, where index \( i = 1, \ldots, i - 1, i + 1, \ldots, d \). If \( d = 1 \), \( k_{i_l|l,m} = n_l \) and \( f_{i_l|l,m} = f_{lm} \). For the Parzen window and \( k \)-nearest neighbour

\[
f_{i_l|l,m} = \frac{k_{lm}}{\sum_{j=1}^{n} |y_{ij} - y_{im}| \leq h_{im}} \frac{k_{lm}}{k_{ij|l,m} h_{im}} = \frac{k_{lm}}{k_{ij|l,m} h_{im}}. \tag{8}
\]
C. Clustering of Observations

The clustering of observations is an iterative procedure which identifies those observations belonging to the \( l \)th component. The deviations between \( k_{ij} \) and the predictive component frequencies for the histogram are given by

\[
e_{ij} = k_{ij} - nt f(\hat{y}_j | \theta_1)V_j.
\]  

(9)

For the Parzen window and \( k \)-nearest neighbour

\[
e_{ij} = k_{ij} - nt f(\hat{y}_j | \theta_1)V_j/k_j.
\]  

(10)

To identify those observations which deviate the most, relative positive deviations \( r_j \) where

\[
\epsilon = e_{ij}/k_{ij}
\]

and component weight estimation until

\[
l = \text{min} \{ e_{ij} \}
\]

and the predictive component frequencies for the histogram are given by

\[
l_j = \prod_{i=1}^{d} f(y_i = \hat{y}_{im} | \theta_d) = f_{im} = \prod_{i=1}^{d} e f_{i|i,lm},
\]  

(14)

from where the required restraints

\[
f(y_i = \hat{y}_{im} | \theta_d) = e f_{i|i,lm} = f_{i|i,lm} \text{max}, i = 1, \ldots, d
\]  

(15)

can be derived. In addition, from known \( f_{lm} \) and \( f_{i|i,lm} \), it follows that

\[
\epsilon = \min \left\{ 1, \left( \frac{f_{lm}}{\prod_{i=1}^{d} f_{i|i,lm}} \right)^{1/2} \right\}.
\]  

(16)

where the upper limit of \( \epsilon \) is set to 1. For Rayleigh, Poisson or binomial distributions with known \( \theta_d \) it is assumed

\[
\frac{\partial f(y_i = \hat{y}_{im} | \theta_d)}{\partial y_i} = 0, i = 1, \ldots, d.
\]  

(17)
The rough component parameters for single parameter distributions are thus gained from restraints (15) or (17). The restraint chosen should ensure that the predictive component density at \( \hat{y}_{im} \) attains its maximum. For two parameter normal, lognormal, Weibull or gamma distributions, the Lagrange multiplier

\[
\Lambda(\Theta_{il}, \lambda_{il}) = -\int_{-\infty}^{+\infty} f(y_i|\Theta_{il}) \log(f(y_i|\Theta_{il})) dy_i + \lambda_{il} \log(f(y_i = \hat{y}_{im}|\Theta_{il})/f_{\hat{y}_{im}}) \tag{18}
\]

provides a strategy for entropy maximization subject to the logarithm of (15). The rough component parameters for two parameter distributions are then a solution of

\[
\nabla_{\Theta_{il}, \lambda_{il}} \Lambda(\Theta_{il}, \lambda_{il}) = 0, \quad i = 1, \ldots, d. \tag{19}
\]

Constrained entropy (18) maximization enables rough Weibull and gamma parameter estimation for shape parameter \( \beta_{il} > 0 \) and not only for \( \beta_{il} > 1 \) as in [24, 25]. Rough normal component parameters are given by

\[
\mu_{il} = \hat{y}_{im} \text{ and } \sigma_{il} = \frac{1}{\sqrt{2\pi f_{\hat{y}_{im}}}}. \tag{20}
\]

Similarly, rough lognormal

\[
f(\lambda_{il}) = \frac{\lambda_{il} - 1}{\lambda_{il}} + \log(\lambda_{il}(\lambda_{il} - 1)) + 2 \log(\sqrt{2\pi f_{\hat{y}_{im}}}) = 0, \quad \mu_{il} = \lambda_{il} - 1 + \log(\hat{y}_{im}) \text{ and } \sigma_{il} = \sqrt{\lambda_{il}(\lambda_{il} - 1)}, \tag{21}
\]

Weibull

\[
f(\alpha_{il}) = \frac{\alpha_{il} - 1}{\alpha_{il}} e^{-\frac{\alpha_{il} - 1}{\alpha_{il}}} - f_{\hat{y}_{im}}\hat{y}_{im} = 0, \quad \lambda_{il} = \frac{\alpha_{il}}{\beta_{il}}, \beta_{il} = \alpha_{il} + \gamma + \log\left(\frac{\alpha_{il} - 1}{\alpha_{il}}\right), \quad \theta_{il} = \hat{y}_{im} \left(\frac{\alpha_{il}}{\alpha_{il} - 1}\right)^{\frac{1}{\alpha_{il}}} \quad \text{and } \beta_{il} > 0, \tag{22}
\]

gamma

\[
f(\alpha_{il}) = \frac{1}{2} \log(\beta_{il}) + \beta_{il} \left( \log\left(\frac{\alpha_{il} - 1}{\alpha_{il}}\right) + \frac{1}{\alpha_{il}} \right) - \log(\sqrt{2\pi f_{\hat{y}_{im}}}) = 0, \quad \beta_{il} = \frac{\gamma(1 + \alpha_{il})}{\gamma - 1 - \alpha_{il} \log\left(\frac{\alpha_{il} - 1}{\alpha_{il}}\right)}, \quad \lambda_{il} = \frac{\alpha_{il}}{\beta_{il}}, \quad \theta_{il} = \frac{\hat{y}_{im} \lambda_{il}}{\alpha_{il} - 1} \quad \text{and } \beta_{il} > 0, \tag{23}
\]

binomial

\[
p_{il} = \begin{cases} 1 - f_{\hat{y}_{im}}^{1/\alpha_{il}} & \hat{y}_{im} = 0 \\ f_{\hat{y}_{im}}^{1/\alpha_{il}} & \hat{y}_{im} = \theta_{il} \\ \hat{y}_{im}/\theta_{il} & \text{otherwise,} \end{cases} \tag{24}
\]

rough Poisson

\[
\theta_{il} = \begin{cases} -\log(f_{\hat{y}_{im}}) & \hat{y}_{im} = 0 \\ \hat{y}_{im} & \text{otherwise} \end{cases} \tag{25}
\]

and rough Dirac

\[
\theta_{il} = \hat{y}_{im} \tag{26}
\]

component parameters are derived, where \( \gamma \) is the Euler-Mascheroni constant. When deriving (23) \( \Gamma[\beta_{il}] \) is approximated by the Stirling’s formula and digamma function by \( \psi(\beta_{il}) = \log(\beta_{il}) - \gamma/\beta_{il} \), the rough binomial parameter \( \theta_{il} = \theta_{i} \) is fixed and equals the number of categories minus one.

Such rigid restraints result in poor component parameter estimation if the modes of several component densities coincide. The loose restraints introduced in [24] improve component parameter estimation and offer further evolution opportunities. The rigid restraints become loose if \( f_{\hat{y}_{im}} \) in equations (20) to (26) is replaced by \( f_{\hat{y}_{im}} \), where

\[
0 \leq f_{\hat{y}_{im}} \leq f_{\hat{y}_{im}}. \tag{27}
\]

Instead of minimizing the maximum relative positive deviation [24], the simpler root finding of the total of relative deviations is used here to attain the optimal \( f_{\hat{y}_{im}} \). For the histogram total of relative deviations

\[
D_{\hat{y}_{im}} = 1 - \sum_{j=1}^{v} f(y_{il} = \hat{y}_{ij}) h_{ij} \tag{28}
\]
equals the fraction of observations falling into regions on the $y_i$ axis with zero empirical probability. If $D_{i,l,m}$ is close to zero, e.g., 0.002, then observations not contributing significantly to the $l$th component should not affect the loose component parameter estimation. This yields

$$\sum_{j=1}^{v} f(\hat{y}_{ij}|\Theta_{il})h_{ij} = 0.998$$

Equation (29) can be solved for optimal $f_{i,l,m}$ by the bisection root finding method. If the root does not exist, then $f_{i,l,m} = f_{i,l,m}^{\text{max}}$. For the Parzen window and $k$-nearest neighbour the root of

$$\sum_{j=1}^{n} f(y_{ij}|\Theta_{il})h_{ij}/k_j = 0.998$$

is searched for optimal $f_{i,l,m}$. The Dirac parameter $\theta_{il}$ of (26) does not require $f_{i,l,m}$ optimization.

**E. Enhanced Component Parameter Estimation**

Maximum likelihood is employed to obtain enhanced component parameters. For the histogram, enhanced normal component parameters are given by

$$\mu_{il} = \frac{1}{n_l} \sum_{j=1}^{v} k_{ij}\hat{y}_{ij} \text{ and } \sigma_{il}^2 = \frac{1}{n_l} \sum_{j=1}^{v} k_{ij}\hat{y}_{ij}^2 - \mu_{il}^2.$$  

Likewise, enhanced lognormal

$$\mu_{il} = \frac{1}{n_l} \sum_{j=1}^{v} k_{ij} \log(\hat{y}_{ij}) \text{ and } \sigma_{il}^2 = \frac{1}{n_l} \sum_{j=1}^{v} k_{ij} \log(\hat{y}_{ij})^2 - \mu_{il}^2.$$  

Weibull

$$\theta_{il} = \frac{1}{n_l} \sum_{j=1}^{v} k_{ij}\hat{y}_{ij}^{\beta_{il}} \text{ and } f(\beta_{il}) = \frac{1}{\beta_{il}} + \frac{1}{n_l} \sum_{j=1}^{v} k_{ij} \log(\hat{y}_{ij}) - \frac{\sum_{j=1}^{v} k_{ij}\hat{y}_{ij}^{\beta_{il}} \log(\hat{y}_{ij})}{\sum_{j=1}^{v} k_{ij}\hat{y}_{ij}^{\beta_{il}}} = 0,$$

gamma

$$\theta_{il} = \frac{1}{\beta_{il}n_l} \sum_{j=1}^{v} k_{ij}\hat{y}_{ij} \text{ and } f(\beta_{il}) = \frac{1}{n_l} \sum_{j=1}^{v} k_{ij} \log(\hat{y}_{ij}) - \log(\theta_{il}) - \frac{\Gamma'[\beta_{il}]}{\Gamma[\beta_{il}]} = 0,$$

binomial

$$p_{il} = \frac{1}{n_l\theta_{il}} \sum_{j=1}^{v} k_{ij}\hat{y}_{ij},$$

Poisson

$$\theta_{il} = \frac{1}{n_l} \sum_{j=1}^{v} k_{ij}\hat{y}_{ij}$$

and Dirac component parameters

$$\theta_{il} = \hat{y}_{ilm}$$

are estimated. Index $v$ is replaced by $n$ for the Parzen window or $k$-nearest neighbour.

**F. First and Second Moment Calculation**

The first and second moment of the normal

$$m_{il} = \mu_{il} \text{ and } V_{il} = \sigma_{il}^2 + \mu_{il}^2,$$

lognormal

$$m_{il} = e^{\mu_{il} + \frac{\sigma_{il}^2}{2}} \text{ and } V_{il} = e^{2\mu_{il} + 2\sigma_{il}^2},$$

Weibull

$$m_{il} = \theta_{il} \Gamma \left[1 + \frac{1}{\beta_{il}}\right] \text{ and } V_{il} = \theta_{il}^2 \Gamma \left[1 + \frac{2}{\beta_{il}}\right],$$

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gamma

\[ m_{il} = \theta_{il}\beta_{il} \quad \text{and} \quad V_{il} = \theta_{il}^2\beta_{il}(1 + \beta_{il}) \]  

(41)

and the first moment of binomial

\[ m_{il} = \theta_{il}p_{il}, \]  

(42)

Poisson

\[ m_{il} = \theta_{il} \]  

(43)

and Dirac

\[ m_{il} = \theta_{il} \]  

(44)

distributions are calculated to enable the classification of the unassigned observations.

**G. Bayes Classification of the Unassigned Observations**

With an increase in the number of components, the number \( n_l \) of unassigned observations decreases

\[ \sum_{k=1}^{l-1} w_k + \frac{n_l}{n} = 1. \]  

(45)

The upper limit \( l - 1 \) equals the assigned number of components \( c \) and \( l \) the index of an additional component that has to be checked whether or not it should be assigned to the finite mixture. The total of positive relative deviations \( D \) is then given by

\[ D = \sum_{k=1}^{l-1} w_k D_k + \frac{n_l}{n} D_l, \]

where \( n_l/n \) and \( D_l \) stand for the weight of the unassigned observations and the total of positive relative deviations of the unassigned observations, respectively. As \( e_{lp} \) in (11) for any unassigned component equals \( n_l \), \( D_l \) equals 1. This yields

\[ D = \sum_{k=1}^{l-1} w_k \frac{e_{lp}}{n_k} + \frac{n_l}{n}. \]  

(46)

From equations (12) and (46) it follows that

\[ D \leq D_{\text{min}}(l - 1) + \frac{n_l}{n}. \]  

(47)

It is reasonable to stop assigning new components when the weight of the unassigned observations is less than or equal to

\[ \frac{n_l}{n} \leq D_{\text{min}}(l - 1). \]  

(48)

Unassigned observations \( k_{lj} \) are then assumed to belong to the existing classes and do not form new ones. The classification of unassigned observations is accomplished by the Bayes decision rule [31]

\[ l = \arg \max_l w_l f(y_j|\Theta_l) \]

\[ w_l = w_l + \frac{k_{lj}}{n}, \quad m_{il} = m_{il} + \frac{k_{lj}(y_{ij} - m_{il})}{nw_l} \quad \text{and} \quad V_{il} = V_{il} + \frac{k_{lj}(y_{ij}^2 - V_{il})}{nw_l}, \]

(49)

where \( k_{lj} \) is added to the \( l \)th class and the component weight and both moments are recalculated [32]. Once all \( v \) bin means or all \( n \) observations are processed, the predictive mixture parameters are gained by inverting equations (38) to (44), inclusive.

**H. Algorithm Flow**

REBMIX is listed in Fig. 1. It requires fourteen arguments and, depending on the parametric families, five or six of them are mandatory whilst the rest are optional. It consists of four main loops: the inner \( 9 \rightarrow 37 \), the middle \( 6 \rightarrow 41 \), the outer \( 4 \rightarrow 47 \) and the outmost loop \( 1 \rightarrow 48 \). The numbers are line indices.

In line 1 the outmost loop begins. It runs over an ordered set of numbers \( K \) of bins \( v \) for the histogram and the Parzen window or numbers of nearest neighbours \( k \) for the \( k \)-nearest neighbour. If absolute distance between the number at which IC attains its optimum and the closest two numbers in \( K \) is greater than 1, set \( K \) is extended by the golden section search method. In line 2 the

\[^1\text{Mandatory argument.}\]
Require: Dataset, Preprocessing, cmmax, Criterion, Variables, pdf, Theta1, Theta2, K, y0, ymin, ymax, ar and Restraints.

Ensure: Dataset contains datasets, Preprocessing is one of "histogram", "Parzen window" or "k-nearest neighbour", cmmax ∈ N, Criterion is one of "AIC", "AIC3", "AIC4", "AICc", "BIC", "CAIC", "HQc", "MDL2", "MDL5", "AWE", "CLC", "ICl", "FC", "ICL-BIC", "D" or "SSE", Variables are "continuous" or "discrete", pdf is one of "normal", "lognormal", "Weibull", "gamma", "binomial", "Poisson" or "Dirac". Theta1 may contain initial binomial parameters, Theta2 is inactive, K ∈ N, y0 may contain origins, ymin and ymax may contain minimum and maximum observations, 0 < ar ≤ 1 and Restraints are "loose" or "rigid".

1: for all k do
2:   Preprocessing of observations
3:   \( I_1 \leftarrow 1, D_{\text{min}} \leftarrow 0.25, k_{ij} \leftarrow k_j \) for \( j = 1 \) to \( v \)
4:   while \( l < l_{\text{max}} \) do
5:     \( l \leftarrow 1, r \leftarrow n, n_l \leftarrow n \)
6:     while \( n_l/n > D_{\text{min}}(l - 1) \) do
7:       Global mode detection
8:       \( I_2 \leftarrow 1, n_l \leftarrow n_l/n, r_j \leftarrow 0 \) for \( j = 1 \) to \( v \)
9:       while \( l_2 < l_{\text{max}} \) do
10:      Rough component parameter estimation
11:     for \( j = 1 \) to \( v \) do
12:       \( e_{ij}^{(l)} \leftarrow 0, e_{in}^{(l)} \leftarrow 0, e_{\text{imax}}^{(l)} \leftarrow 0 \)
13:         \( e_{ij} \leftarrow 0, e_{ij} \leftarrow 0 \)
14:         if \( k_{ij} > 0 \) or \( r_j > 0 \) then
15:           \( e_{ij} \leftarrow k_{ij} - n_l/(y_j(\Theta)) V_j \)
16:         if \( e_{ij} > 0 \) then
17:           \( e_{ij} \leftarrow e_{ij}/k_{ij} \)
18:           \( e_{l_{\text{imax}}} \leftarrow \max\{e_{l_{\text{max}}}, e_{ij}\} \)
19:         end if
20:       end if
21:     end for
22:   end while
23:   \( D_l \leftarrow e_{\text{ip}}/n_l, e_{\text{imax}}^{(l)} \leftarrow e_{l_{\text{imax}}}(1 - \alpha) \)
24: if \( D_l > D_{\text{min}}/w_l \) then
25:   for all \( j \) such that \( 1 \leq j \leq v \) and \( e_{ij} > e_{l_{\text{imax}}} \) do
26:     \( k_{ij} \leftarrow k_{ij} - e_{ij}, r_j \leftarrow r_j + e_{ij}, n_l \leftarrow n_l - e_{ij} \)
27:   end for
28: if \( e_{\text{ip}} > e_{\text{ip}}/D_l - n_l, f \leftarrow e_{\text{ip}}/e_{in} \) if \( e_{in} > e_{\text{ip}} \) otherwise \( f \leftarrow 1 \)
29: if \( f > 1 \) then \( e_{ij} \leftarrow f e_{ij}, k_{ij} \leftarrow k_{ij} - e_{ij}, r_j \leftarrow r_j + e_{ij}, n_l \leftarrow n_l - e_{ij} \)
30: end for
31: \( w_l \leftarrow n_l/n \)
32: end if
33: end for
34: Enhanced component parameter estimation, break
35: end if
36: \( I_2 \leftarrow I_2 + 1 \)
37: end while
38: First and second moment calculation
39: \( c \leftarrow l, r \leftarrow r - n_l, l \leftarrow l + 1, n_l \leftarrow r, k_{ij} \leftarrow r_j \) for \( j = 1 \) to \( v \)
40: Stop \( \leftrightarrow c \geq v \) or \( c \geq \text{cmax}, \text{break} \) if \( \text{Stop} = \text{true} \)
41: end while
42: Bayes classification of the unassigned observations, log likelihood \( \log L \), information criterion IC and total of positive relative deviations \( D \) calculation
43: if IC < IC_{opt} then
44: \( \log L \rightarrow \log L_{\text{opt}}, IC \rightarrow IC_{opt}, c \rightarrow c_{opt}, w \rightarrow w_{\text{opt}}, \Theta \rightarrow \Theta_{\text{opt}} \)
45: end if
46: break if \( \text{Stop} = \text{true} \), \( D_{\text{min}} \leftarrow cD_{\text{min}}/(c + 1), I_1 \leftarrow I_1 + 1 \)
47: end while
48: end for

Fig. 1 REBMIX algorithm
observations are preprocessed as described in Section A.. In line 3, counter \( I_1 \), constant \( D_{\text{min}} \), and frequencies \( k_{ij} \) are initiated. Next, the outer loop begins. Line 5 assumes the mixture consists of one component, the number of observations \( r \) to separate is set to \( n \) and \( n_1 \) to \( n \). If inequality (48) in line 6 is not fulfilled, the middle loop begins.

In lines 7 and 8, the global mode argument \( m \) is detected as explained in Section B., counter \( I_2 \) is initiated, component weight \( w_1 \) is calculated and frequencies \( r_j \) are set to zero. If \( I_2 \leq I_{\text{max}} \), the inner loop begins, otherwise in line 38 the first and the second moments are calculated (see Section F.). In line 39 the number of components \( c \) is set to \( l \), the number of observations \( r \) is decreased by \( n_1 \), \( l \) is incremented, the number of the unassigned observations \( r \) joins \( n_1 \), and the frequencies of the unassigned observations \( r_j \) are moved to \( k_{ij} \). If the number of components \( c \) in line 40 is greater or equal than \( v \) or \( c_{\text{max}} \), Stop is set to true and the middle loop ends.

The inner loop is divided into three sections. In line 10 the component parameters are roughly estimated (see Section D.). In the second section \( 11 \rightarrow 23 \), deviations \( e_{lj} \), the total of positive deviations \( e_{lp} \), the total of negative deviations \( e_{ln} \), the total of positive relative deviations \( D_1 \) and maximum relative deviation \( e_{\text{max}} \) are calculated. The number of iterations depends on the acceleration rate \( a \). In the third section \( 24 \rightarrow 35 \), the maximum and negative deviations are transferred between frequencies \( k_{lj} \) and \( r_j \). This way deviations \( e_{lj} \) are reduced gradually. The negative value of \( e_{lj} \) can never be higher than \( r_j \). If this is not true, deviation \( e_{lj} \) is corrected as listed in line 19. When the condition in line 24 is not fulfilled, the enhanced component parameter estimation is carried out (see Section E.) and the inner loop ends.

Enhanced component parameter estimation may fail. In this instance, the component parameters are reset to the state just before the failure occurred. In line 42 the unassigned observations are classified by the Bayes decision rule as depicted in Fig. 1. When the condition in line 24 is not fulfilled, the enhanced component parameter estimation is carried out (see Section E.) and the inner loop ends.

The binomial, Poisson and Dirac mixtures require \( M = cd + c - 1 \). The log likelihood function for the binned observations is given by

\[
\log L(c, w, \Theta) = \sum_{j=1}^{v} k_j \log f(\bar{y}_j | c, w, \Theta).
\]  

Otherwise

\[
\log L(c, w, \Theta) = \sum_{j=1}^{n} \log f(y_j | c, w, \Theta).
\]  

Finally, the total of positive relative deviations for the histogram

\[
D = \sum_{j=1}^{n} \left\langle \frac{k_j}{n} - f(\bar{y}_j | c, w, \Theta)V_j \right\rangle,
\]  

Parzen window or \( k \)-nearest neighbour

\[
D = \frac{1}{n} \sum_{j=1}^{n} \left\langle \frac{1}{k_j} - \frac{f(y_j | c, w, \Theta)V_j}{k_j} \right\rangle
\]  

is calculated, where \( \langle x \rangle = x \) if \( x > 0 \) and \( \langle x \rangle = 0 \) if \( x \leq 0 \). This way the optimum IC_{opt} corresponding to the optimal number of components \( c_{\text{opt}} \), weights \( w_{\text{opt}} \) and parameters \( \Theta_{\text{opt}} \) is reached. The outer loop ends if the Stop criterion in line 46 is true. Otherwise \( D_{\text{min}} \) is decreased in such a way that the total of positive relative deviations

\[
eD_{\text{old}} = (c + 1)D_{\text{new}}^\text{old}
\]

for the \( c \) and \( c + 1 \) components is preserved. If index \( v \) in Fig. 1 is replaced by \( n \) and line 15 is replaced by (10) the algorithm, presented for the histogram, can also be used with the Parzen window and \( k \)-nearest neighbour.

### III. APPLICATION

A comparison between the EM algorithm and the REBMIX algorithm for time efficiency and accuracy has already been studied in [22, 23]. Therefore, the latest results [7] obtained by comparing the EM algorithm implemented in FlexMix [26] to REBMIX are only summarized here. Two cases have been investigated. In case 1, a dataset of size \( n = 10000 \) generated from the three component multivariate Weibull-normal mixture density has been considered. The FlexMix algorithm attained
an AIC = 223736 with three components in 149.42s, whereas the REBMIX algorithm resulted in an AIC = 223999 with four components in 1.53s. In case 2, the Weibull-normal mixture density has been applied to a real dataset, where the number of components was not known. The FlexMix algorithm has shown some convergence problems and resulted in five components and an AIC = 562113 in 1801.52s, whereas the REBMIX algorithm resulted in nine components and an AIC = 559637 in 7.09s. The latest study has confirmed our previous experience with the EM algorithm. The REBMIX algorithm is time efficient and robust. The main advantage of the EM algorithm is its accuracy if convergence problems are avoided by a carefully selecting the initial set of unknown parameters. Since the efficiency and accuracy of the REBMIX algorithm have already been dealt with in [22, 23, 7], the rest of this section stresses other capabilities of the REBMIX algorithm by studying univariate and multivariate datasets.

### A. Univariate Normal Datasets

As the family of finite normal mixtures is very flexible, it was used by [34] to represent a wide variety of density shapes in their analytical study of the mean integrated squared error of the kernel density estimator. To demonstrate that the family of normal mixtures is very broad, 15 carefully chosen examples were given. The first five represent different types of problems that can arise for unimodal densities. The rest of the densities are multimodal. Densities 6 to 9 are mildly multimodal and might be estimated fairly well with a dataset of a moderate size. The remaining densities are strongly multimodal and are very hard to recover in full with moderate dataset size. However, they are still worth studying because the issue of just how much can be recovered is important.

To find out the extent to which `rebmix` has improved, datasets of sizes 100, 1000 and 10000 are generated for these 15 finite normal mixtures under identical conditions as in [24]. For each dataset size and each density from 1 to 15, 100 datasets are generated. See `help("RNGMIX")`, `help("REBMIX")` and `help("kseq")` in `rebmix` for details.

The `REBMIX` function is applied to the three dataset sizes and the 15 finite normal mixtures. The preprocessing is set to histogram, the maximum number of components to 20 and the information criterion to BIC. The number of bins ranges from Sturges [35] to Log10 corresponding to the Log10 rule or to the RootN rule. One or two values appear in Tables 1 and 2. Where two values are shown, they are separated by a slash sign. If the calculations are performed for $K$ ranging from Sturges:Log10, then the values in the tables are on the left side of the slash sign. If $K$ ranges from Sturges:RootN, then the values are on the right side of the slash sign. To speed up computation, $K$ is set to `kseq(from = Sturges, to = RootN, f = 0.05)` for datasets of size 10000. If there is no slash, $K = $Sturges:Log10. The origin $y_0 = 0.0$, minimum $y_{min}$ and maximum $y_{max}$ observations are set to the outmost values of 100 datasets attained by the `RNGMIX` function.

The true numbers of components $c$ from which the datasets are generated for the 15 densities, predictive mean numbers of components $\mu_c$ and the corresponding standard deviations $\sigma_c$ are shown in Table 1 and are compared to Table 2 in [24]. In testing the null hypothesis, that the predictive mean number of components $\mu_c$ with the corresponding standard deviation $\sigma_c$ equals the

| Density          | $c$ | $\mu_c$ | $\sigma_c$ | $c$ | $\mu_c$ | $\sigma_c$ | $c$ | $\mu_c$ | $\sigma_c$ |
|------------------|-----|---------|------------|-----|---------|------------|-----|---------|------------|
| Gaussian         | 1   | 1.0     | 0.00       | 1   | 1.0     | 0.00       | 1   | 1.0     | 0.24       |
| Skewed unimodal  | 3   | 1.2     | 0.36       | 2   | 2.1     | 0.29       | 4.4/2.4 | 1.53/0.84 |
| Strongly skewed  | 8   | 2.5     | 0.56       | 7.3 | 1.09    | 16.2       | 1.74 |
| Kurtotic unimodal| 2   | 2.0     | 0.10       | 2.1 | 0.22    | 2.0        | 2.0  |
| Outlier          | 2   | 2.2     | 0.55       | 2.5 | 0.50    | 2.2        | 0.49 |
| Bimodal          | 2   | 1.6     | 0.50       | 2.0 | 0.00    | 2.2        | 0.66 |
| Separated bimodal| 2   | 2.0     | 0.10       | 2.0 | 0.20    | 2.4/2.0    | 0.69/0.00 |
| Skewed bimodal   | 2   | 1.5     | 0.50       | 2.2 | 0.36    | 4.0        | 1.23 |
| Trinodal         | 3   | 1.8     | 0.42       | 2.4/2.5 | 0.48/0.50 | 4.2/3.0    | 1.36/0.20 |
| Claw             | 6   | 1.1     | 0.27       | 3.4/5.8 | 1.64/1.55 | 9.5        | 1.33 |
| Double claw      | 9   | 1.6     | 0.50       | 2.0 | 0.10    | 2.1        | 0.62 |
| Asymmetric claw  | 6   | 1.1     | 0.27       | 4.3 | 1.94    | 10.1       | 2.28 |
| Asymmetric double claw | 8   | 1.7     | 0.45       | 2.0 | 0.17    | 5.0        | 3.08 |
| Smooth comb      | 6   | 3.2     | 0.55       | 5.1 | 0.96    | 10.0       | 2.15 |
| Discrete comb    | 6   | 3.9     | 0.32       | 5.0 | 0.83    | 11.0       | 1.32 |

...
true number of components $c$, one sample $t$-test

$$t = \frac{\mu_c - c}{\sigma_c / \sqrt{n_c}}$$ \hspace{1cm} (56)

is used. Let statistics $t$ calculated from Table 2 \([24]\) be $t^{\text{old}}$ and let those from Table 1 be $t^{\text{new}}$. If $t^{\text{new}} < t^{\text{old}}$, an improvement denoted by + in Table 2 is observed. If $t^{\text{new}} = t^{\text{old}}$, notation 0 means unchanged. If $t^{\text{new}} > t^{\text{old}}$, a deterioration denoted by – is observed.

Table 2 shows that for $n = 100$ an improvement of 20% is achieved. For $n = 1000$ an improvement of 27% is achieved. However, for $n = 10000$ a worsening by 7% is observed. This means that for the three dataset sizes a 13% change of the old state is observed. If $\text{RootN}$ is chosen for maximum $K$, no improvement is observed for $n = 100$ as the value of $K$ for $n = 100$ for both $\text{Log10}$ and $\text{RootN}$ rules is identical. For $n = 1000$ and 10000 an improvement of 53% and 20% is observed, respectively. This means that for the three dataset sizes the old state is improved by 31% on average.

The gaussian density can be recovered by the REBMIX algorithm. When the unimodal density is skewed, $\mu$, tends towards $c = 3$ for higher $n$. The strongly skewed density can not be recovered satisfactorily even for $n = 10000$. These kind of densities point out one of the limitations of the REBMIX algorithm which will be difficult to suppress. The kurtotic unimodal, outlier, bimodal and separated bimodal densities can also be recovered however the skewed bimodal density is slightly overfitted for $n = 10000$. The trimodal density can be better recovered for higher $n$. The claw density consists of one broad density with $w_1 = 1/2$ and five superimposed narrow densities with $w_2 = \ldots = w_6 = 1/10$. It can be recovered for $n = 1000$, however overfitting occurs for high $n$. This suggests further improvements of the loose restraints are required. The double claw density where three narrow densities with component weights of $7/300$ can be recovered completely for the highest $n$. The asymmetric double claw density can be better recovered for higher $n$ values. Further improvements of the loose restraints could also lead to improvements in this area. The smooth comb and discrete comb densities can be modelled if $n$ is high enough although overfitting occurs. The optimal number of bins should preferably be searched within the broadest limits of $K$. The rebmix.univariate.normal.R script file that generates the results is available in the rebmix package demo subdirectory. To run it enter demo("rebmix.univariate.normal.R","rebmix") and press return.

| Density               | $n = 100$ | $n = 1000$ | $n = 10000$ |
|-----------------------|-----------|------------|-------------|
| 1 Gaussian            | 0         | 0          | 0           |
| 2 Skewed unimodal     | –         | +          | –/+         |
| 3 Strongly skewed      | +         | +          | –           |
| 4 Kurtotic unimodal   | +         | –          | 0           |
| 5 Outlier             | –         | –          | +           |
| 6 Bimodal             | +         | 0          | +           |
| 7 Separated bimodal   | 0         | 0          | –/0         |
| 8 Skewed bimodal      | +         | +          | +           |
| 9 Trimodal            | –         | –/+        | –/0         |
| 10 Claw               | +         | –/+        | –           |
| 11 Double claw        | +         | +          | +           |
| 12 Asymmetric claw    | +         | +          | +           |
| 13 Asymmetric double claw | –     | +          | +           |
| 14 Smooth comb        | +         | +          | –           |
| 15 Discrete comb      | –         | +          | –           |

|               | 20% | 27%/53% | -7%/20% |

B.Galaxy Dataset

The dataset analysed in \([36]\) contains the measurements of the velocities of 82 galaxies diverging away from our own galaxy. The multimodality of the velocities may indicate the presence of super clusters of galaxies surrounded by large voids, each mode representing a cluster moving away at its own speed \([36, \text{gives more background}]\). Richardson and Green \([37]\) concluded from their approach that the number of components ranged from 5 to 7, while \([38]\) provided the support for six components. Stephens \([39]\) reported that three components were optimal for the mixture of normal and four for the mixture of $t$ distributions.
The \textsc{rebmix} function in \texttt{rebmix} is entered for normal, lognormal, Weibull and gamma parametric families. The maximum number of components $c_{\text{max}}$ is set to 10. The influence of the Akaike [33] information criterion AIC and the Bayesian [40] information criterion BIC for the histogram and Parzen window preprocessing on predictive number of components $c$ is studied. The optimal number of bins is searched within the broadest limits for $K = 7:20$. The minimum information criterion and

### TABLE 3

| Parametric family | Preprocessing | Criterion | $c$ | $v$ | IC   | log $L$ |
|-------------------|---------------|-----------|-----|-----|------|--------|
| normal            | histogram     | AIC       | 6   | 20  | 424  | -195   |
|                   | histogram     | BIC       | 4   | 19  | 453  | -202   |
|                   | Parzen window | AIC       | 6   | 18  | 431  | -199   |
|                   | Parzen window | BIC       | 4   | 16  | 459  | -205   |
| lognormal         | histogram     | AIC       | 6   | 20  | 422  | -194   |
|                   | histogram     | BIC       | 3   | 19  | 442  | -203   |
|                   | Parzen window | AIC       | 6   | 19  | 436  | -201   |
|                   | Parzen window | BIC       | 3   | 8   | 461  | -213   |
| Weibull           | histogram     | AIC       | 5   | 15  | 428  | -200   |
|                   | histogram     | BIC       | 4   | 17  | 458  | -205   |
|                   | Parzen window | AIC       | 4   | 11  | 435  | -207   |
|                   | Parzen window | BIC       | 4   | 11  | 462  | -207   |
| gamma             | histogram     | AIC       | 6   | 20  | 423  | -194   |
|                   | histogram     | BIC       | 3   | 19  | 444  | -204   |
|                   | Parzen window | AIC       | 6   | 18  | 432  | -199   |
|                   | Parzen window | BIC       | 4   | 14  | 462  | -207   |

The maximum log likelihood in Table 3 are observed for the histogram preprocessing. The maximum log likelihood resulting in 6 components coincides with the lognormal and gamma parametric family types and the AIC. Most frequently six and four components appear. For this particular dataset the AIC is favorable giving 4 to 6 components. The \texttt{plot} method delivers a fitted

### TABLE 4

| $l$  | 1  | 2  | 3  | 4  | 5  | 6  |
|------|----|----|----|----|----|----|
| $w_l$| 0.456 | 0.154 | 0.243 | 0.0854 | 0.0372 | 0.0244 |
| $\mu_l$| 2.99 | 3.11 | 3.18 | 2.28 | 3.49 | 2.78 |
| $\sigma_l$| 0.041 | 0.034 | 0.0648 | 0.0511 | 0.029 | 0.0312 |

finite mixture, complete with legend, in Fig. 2. The corresponding predictive lognormal mixture parameters in Table 4 are given

Fig. 2 Empirical density and distribution function (circles) and predictive lognormal mixture density and distribution function (solid line) for galaxy dataset
by the \texttt{coef} method. For the details on specifying arguments for the \texttt{plot} and \texttt{coef} methods see \texttt{help("plot.REBMIX")}
and \texttt{help("coef.REBMIX")}, respectively. The \texttt{rebmix.galaxy.R} script file is made available via \texttt{demo("rebmix.
galaxy","rebmix")}.

\section*{C.Mixed Continuous-discrete Dataset}

A multivariate mixed continuous-discrete 3 component mixture is generated here by calling the \texttt{RNGMIX} function. The

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig3.png}
\caption{Empirical densities (large coloured circles), predictive multivariate marginal lognormal-Poisson-binomial-Weibull mixture densities (coloured lines and small circles), empirical densities (circles), predictive univariate marginal lognormal, Poisson, binomial and Weibull mixture densities and progress charts (solid line)}
\end{figure}

REBMIX function is called for the multivariate lognormal-Poisson-binomial-Weibull parametric family type. To plot the \texttt{rebmix} mixture in Fig. 3 the \texttt{plot} method is called. By calling the \texttt{boot.REBMIX} function \texttt{B} bootstrap datasets of length \(n\) are generated for the \(x\) object of class \texttt{REBMIX} at position \(pos\), where bootstrap \texttt{Bootstrap} can be one of the default "parametric" or "nonparametric" options. Arguments \texttt{replace} and \texttt{prob} affect the nonparametric bootstrap only, see \texttt{help("sample")} and [38] for details regarding replacement and weighted bootstrap.

The \texttt{mixedboot} object of class \texttt{boot.REBMIX} holds a data frame \(c = \{9, 8, 6, \ldots, 7, 5, 5, 5, 5\}\) containing numbers of components \(c\) for \(B = 100\) bootstrap datasets, standard error \(c.se = 1.54\), coefficient of variation \(c.cv = 0.259\), mode \(c.mode = 5\) and mode probability \(c.prob = 0.31\) of the numbers of components. Component weights \(w\), component parameters \(\theta_1.i\) and \(\theta_2.i\), standard errors \(w.se, \theta_1.i.se\) and \(\theta_2.i.se\) and coefficients of variation \(w.cv, \theta_1.i.cv\) and \(\theta_2.i.cv\) for those bootstrap datasets for which \(c = 5\) are also returned. See \texttt{help("boot.REBMIX")} in \texttt{rebmix} for details. The \texttt{demo("rebmix.mixed.continuous.discrete","rebmix")} makes the section reproducible.

\section*{IV. Conclusions}

The article presents the REBMIX algorithm and the \texttt{rebmix} package. The datasets are studied on the x64 architecture. By applying the \texttt{tikzDevice} package [41], plots with legends are obtained. The REBMIX algorithm can be used to assess an initial set of unknown parameters and number of components for, e.g., the EM algorithm or as a standalone procedure that is a good
compromise between the parametric and nonparametric methods of finite mixture estimation. Its major advantages are robustness and time efficiency. Its advantages are more apparent if mixtures are composed of a larger number of components. On the other hand, it requires the preprocessing of observations, which is a drawback, especially where smaller datasets are concerned. The restoration of overlapping components is not always adequate either. However, further improvements of the loose restraints seem to be feasible. The rebmix package can be broadened to other parametric family types, including those with conditionally dependent component densities, e.g., multivariate normal mixtures with full covariance matrices. Currently multivariate normal mixtures with diagonal covariance matrices are available. In time further improvements will see an extensibility of the REBMIX algorithm whereby users should have the ability to provide their own parametric family types for the rapid prototyping of new finite mixture models. The RCLSMIX function that enables class membership prediction is available in the rebmix package. See help("RCLSMIX") for details. The REBMIX is thus also intended to be used for pattern recognition.

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