MoTBFs: An R Package for Learning Hybrid Bayesian Networks Using Mixtures of Truncated Basis Functions

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Abstract This paper introduces **MoTBFs**, an R package for manipulating mixtures of truncated basis functions. This class of functions allows the representation of joint probability distributions involving discrete and continuous variables simultaneously, and includes mixtures of truncated exponentials and mixtures of polynomials as special cases. The package implements functions for learning the parameters of univariate, multivariate, and conditional distributions, and provides support for parameter learning in Bayesian networks with both discrete and continuous variables. Probabilistic inference using forward sampling is also implemented. Part of the functionality of the **MoTBFs** package relies on the **bnlearn** package, which includes functions for learning the structure of a Bayesian network from a data set. Leveraging this functionality, the **MoTBFs** package supports learning of MoTBF-based Bayesian networks over hybrid domains. We give a brief introduction to the methodological context and algorithms implemented in the package. An extensive illustrative example is used to describe the package, its functionality, and its usage.

Introduction

Mixtures of truncated basis functions (MoTBFs) (Langseth et al., 2012a) have been proposed as a general framework for handling hybrid Bayesian networks, i.e., Bayesian networks where discrete and continuous variables coexist. As special cases, the framework includes the so-called mixtures of truncated exponentials (MTEs) (Moral et al., 2001) and mixtures of polynomials (MoPs) (Shenoy and West, 2011; López-Cruz et al., 2012).

One of the advantages of MoTBFs is that they allow for hybrid Bayesian networks with no structural restrictions on the relations between the continuous and discrete variables; this is in contrast to conditional Gaussian (CG) models (Lauritzen, 1992), where discrete variables are not allowed to have continuous parents. Restricting arc directions is problematic, for instance in situations where the network structure is given a causal interpretation; the fact that MoTBFs have no such restriction make them suitable for this kind of analysis. Furthermore, MoTBFs are closed under addition, multiplication, and integration, which facilitates the use of exact probabilistic inference methods like the Shenoy-Shafer architecture (Shenoy and Shafer, 1990) or the variable elimination algorithm (Zhang and Poole, 1996).

Methods for learning MoTBFs from data have previously been studied and cover algorithms for learning both marginal (Langseth et al., 2012b,a) and conditional MoTBF densities from data (Langseth et al., 2009, 2014; Pérez-Bernabé et al., 2015). To address situations where data availability is limited, Pérez-Bernabé et al. (2016) proposed a methodology for integrating prior knowledge when learning univariate and conditional MoTBFs from data. The underlying idea of the integration is to represent the prior knowledge as an MoTBF density that is later combined with the density learned from data, thus resulting in a new MoTBF density. Together, the learning algorithms for MoTBF-based Bayesian networks facilitate learning in data-rich domains as well as domains where limited quantitative data is counterbalanced by qualitative domain knowledge.

The aim of the MoTBFs package is to provide a free and accessible implementation of algorithms for learning MoTBFs from data. The package implements state-of-the-art learning algorithms for univariate, conditional, and joint MoTBF densities. By extension, functionality is also provided for learning MoTBF-based Bayesian networks by leveraging functionality from the bnlearn package (Scutari, 2010). Hybrid Bayesian networks supported by bnlearn are restricted to CG models, which implies that conditional distributions of discrete variables given continuous ones are not permitted. Other packages like deal (Bøttcher and Dethlefsen, 2003) and pcalg (Kalisch et al., 2012) are also restricted to CG models. By adopting the MoTBF framework, the MoTBFs package sidesteps this restriction on the possible network structures. Furthermore, the MoTBFs package also provides methods for integrating prior domain knowledge in the learning process, thus also supporting data sparse domains.

Another package that includes hybrid Bayesian networks functionality is **HydeNet** (Dalton and Nutter, 2019). This package is able to handle conditional distributions beyond the Gaussian model class, but discrete variables conditional on continuous ones are modeled using generalized linear models, and inference is only possible using Markov Chain Monte Carlo through an interface to

JAGS (Plummer, 2003). Unlike **MoTBFs** and **bnlearn**, **HydeNet** does not provide functionality for learning the network structure. **HydeNet** is especially appropriate for modeling decision problems, since it implements influence diagrams, which are extensions of Bayesian networks that include decision and utility nodes, similarly to decision trees.

The package **abn** (Kratzer et al., 2019) deals with so-called *additive Bayesian networks*, which are Bayesian networks where each node holds a generalized linear model, and the effect of the parents of each node is additive in terms of the exponential family expression of the conditional distribution. Unlike **HydeNet**, **abn** is able to learn the network structure from data, and adopts a fully Bayesian approach, thus allowing the specification of prior distributions on the parameters in a natural way. The main difference with respect to package **MoTBFs** is that MoTBF distributions do not belong to the exponential family, and in that sense both packages are complementary.

Mixtures of truncated basis functions

The MoTBF framework (Langseth et al., 2012a) is based on the abstract notion of real-valued *basis functions*, which includes both polynomial and exponential functions as special cases.

More formally, let **X** be a mixed n-dimensional random vector. Let $\mathbf{Y} = (Y_1, \dots, Y_d)$ and $\mathbf{Z} = (Z_1, \dots, Z_c)$ be the discrete and continuous parts of **X**, respectively, with c + d = n. Let $\Psi = \{\psi_i(\cdot)\}_{i=0}^{\infty}$ with $\psi_i : \mathbb{R} \to \mathbb{R}$ define a collection of real basis functions. We say that a function $\hat{f} : \Omega_{\mathbf{X}} \mapsto \mathbb{R}_0^+$ is a mixture of truncated basis functions potential to level k wrt. Ψ if one of the following two conditions holds:

• *f* can be written as

$$f(\mathbf{x}) = f(\mathbf{y}, \mathbf{z}) = \sum_{i=0}^{k} \prod_{j=1}^{c} a_{i,\mathbf{y}}^{(j)} \psi_i \left(z_j \right), \tag{1}$$

where $a_{i,\mathbf{v}}^{(j)}$ are real numbers.

• There is a partition $\Omega^1_{X'}, \dots, \Omega^m_{X}$ of Ω_X for which the domain of the continuous variables, Ω_Z , is divided into hyper-cubes and such that f is defined as

$$f(\mathbf{x}) = f_{\ell}(\mathbf{x}) \quad \text{if } \mathbf{x} \in \Omega^{\ell}_{\mathbf{X}},$$

where each f_{ℓ} , $\ell = 1, ..., m$ can be written in the form of Equation 1.

Typically, a univariate MoTBF for a variable X does not rely on a partitioning of Ω_X .

To see the relationship between MoTBFs and MoPs (Shenoy and West, 2011), we can instantiate the basis functions as polynomials (i.e., $\psi_i(x) = x^i$, for i = 0, ..., k) in which case the MoTBF model reduces to an MoP model. For example, with polynomial basis functions, a univariate MoTBF of level k for a variable X is given by

$$f(x) = \sum_{i=0}^{k} \theta_i \, \psi_i(x) \,.$$

Similarly, by having exponential basis functions the MoTBF model implements an MTE model (Moral et al., 2001). For ease of exposition, we shall in the remainder of this paper assume polynomial basis functions unless explicitly stated otherwise.

An MoTBF potential is a *density* if $\sum_{\mathbf{y} \in \Omega_{\mathbf{Y}}} \int_{\Omega_{\mathbf{z}}} f(\mathbf{y}, \mathbf{z}) d\mathbf{z} = 1$. Similarly, we say that an MoTBF $f(\mathbf{y}, \mathbf{z})$ is a *conditional density* for $\mathbf{Z}' \subseteq \mathbf{Z}$ and $\mathbf{Y}' \subseteq \mathbf{Y}$ given $(\mathbf{Z} \setminus \mathbf{Z}')$ and $(\mathbf{Y} \setminus \mathbf{Y}')$ if

$$\sum_{\mathbf{y}' \in \Omega_{\mathbf{x}'}} \int_{\Omega_{\mathbf{z}'}} f(\mathbf{y}', \mathbf{y}'', \mathbf{z}', \mathbf{z}') d\mathbf{z}' = 1,$$

for all $\mathbf{z}'' \in \Omega_{\mathbf{Z} \setminus \mathbf{Z}'}$ and $\mathbf{y}'' \in \Omega_{\mathbf{Y} \setminus \mathbf{Y}'}$. Following Langseth et al. (2012a) we furthermore assume that the influence that a set of continuous parent variables \mathbf{Z} have on their child variable X is encoded only through the partitioning of $\Omega_{\mathbf{Z}}$ into hyper-cubes, and not directly in the functional form of $f(x|\mathbf{z})$ inside the hyper-cube $\Omega_{\mathbf{Z}}^{j}$. That is, for a partitioning $\mathcal{P} = \{\Omega_{\mathbf{Z}}^{1}, \dots, \Omega_{\mathbf{Z}}^{m}\}$ of $\Omega_{\mathbf{Z}}$, the conditional MoTBF is defined for $\mathbf{z} \in \Omega_{\mathbf{Z}}^{j}$, $1 \leq j \leq m$, as

$$f_k^{(j)}(x|\mathbf{z} \in \Omega_{\mathbf{Z}}^j) = \sum_{i=0}^k \theta_i^{(j)} \, \psi_i^{(j)}(x).$$
 (2)

In the remainder of this paper we shall assume that a conditional MoTBF density includes only a single 'head' variable, i.e., $|\mathbf{Z}' \cup \mathbf{Y}'| = 1$.

Learning univariate MoTBFs from data

Langseth et al. (2014) present a method for learning univariate MoTBF distributions from data. This method is also implemented in the **MoTBFs** package and is briefly described here together with its extension to both conditional and joint distributions. The estimation procedure relies on the empirical cumulative distribution function (CDF) as a representation of the data, which, for a sample $D = \{x_1, \ldots, x_N\}$, is defined as

$$G_N(x) = \frac{1}{N} \sum_{\ell=1}^{N} \mathbf{1}\{x_{\ell} \le x\}, \quad x \in \mathbb{R},$$
 (3)

where $\mathbf{1}\{\cdot\}$ is the indicator function.

The algorithm developed by Langseth et al. (2014) fits a potential, whose derivative is an MoTBF, to the empirical CDF using least squares. As an example, if we use polynomials as basis functions, $\Psi = \{1, x, x^2, x^3, \ldots\}$, the parameters of the CDF, denoted as c_0, \ldots, c_k , are estimated by solving the optimization problem

$$\begin{aligned}
& \underset{c_0, \dots, c_k}{\text{minimize}} \sum_{\ell=1}^{N} \left(G_N(x_{\ell}) - \sum_{i=0}^{k} c_i x_{\ell}^i \right)^2 \\
& \text{subject to } \sum_{i=1}^{k} i c_i x^{i-1} \ge 0 \quad \forall x \in \Omega, \\
& \sum_{i=0}^{k} c_i \alpha^i = 0 \text{ and } \sum_{i=0}^{k} c_i \beta^i = 1,
\end{aligned} \tag{4}$$

where the constraints ensure that the obtained parameter estimates define a valid CDF, with α and β being the minimum and maximum values of the data sample, respectively. More precisely, the first constraint guarantees that the corresponding density is non-negative and the last two restrictions ensure that it integrates to 1. The latter is equivalent to stating that the CDF should be equal to 0 at the minimum and equal to 1 at the maximum. Note that the estimated function is not actually a density, but a CDF instead. An MoTBF density can be obtained by simply taking the derivative of the CDF.

Note that the optimization program above is convex, and can be efficiently solved in theory. However, the infinite number of constraints introduced by imposing that $\frac{\mathrm{d}F(x)}{\mathrm{d}x} \geq 0$ for all $x \in \Omega_X$ complicates the implementation on a computer. In practice, we only check that the constraint is fulfilled for a limited set of points spread across Ω_X .

In learning scenarios involving a large amount of data (i.e., when N is large), solving the program can be time consuming. In such cases we define a *grid* on Ω_X , that is selected so that the number of observations is the same between each pair of consecutive grid-points. The grid-points are used to evaluate the objective function instead of the sample points.

The level k of the estimated MoP can be decided using different model selection techniques. For the results presented in this paper we have performed a greedy search, choosing the value for k maximizing the Bayesian information criterion (BIC) (Schwarz, 1978):

$$BIC(f, D) = \sum_{\ell=1}^{N} \log f(x_{\ell}) - \frac{k+2}{2} \log N.$$
 (5)

This choice is motivated by Langseth et al. (2014), who showed that the estimators based on Equation 4 are consistent in terms of the mean squared error for all $x \in \Omega_X$.

Learning conditional MoTBFs from data

In a conditional MoTBF density, the continuous parent variables \mathbf{Z} only influence the child variable X through the partitioning of the domain of \mathbf{Z} into hyper-cubes, and not directly in the functional form of $f(x|\mathbf{z})$ inside each hyper-cube (Langseth et al., 2012a). Thus, learning a conditional MoTBF basically consists in finding a partitioning of the domain of the parent variables and using the procedure above for estimating the density of the child variable for each of these partitions.

This procedure is formally described by Langseth et al. (2014), where the domain of the parent

variables, $\Omega_{\mathbf{Z}}$, is incrementally split as long as the BIC score improves. Equal frequency binning is used to determine candidate split points. After splitting a variable Z, the algorithm fits a univariate MoTBF density for each induced sub-partition $\Omega_{\mathbf{Z}}^1$ and $\Omega_{\mathbf{Z}}^2$. A candidate partition $\Omega_{\mathbf{Z}'}$ is only accepted if the BIC score is improved, i.e., if

$$BIC$$
-Gain $(\Omega'_{\mathbf{Z}}, Z) = BIC(f', D) - BIC(f, D) > 0$,

where f' is the conditional MoTBF potential defined over the candidate partition.

Learning joint MoTBFs from data

The procedure for learning joint densities is an extension of the program in Equation 4 to random vectors of arbitrary dimension (Pérez-Bernabé et al., 2015). In the multivariate case, the sample is a set of d-dimensional observations, $\mathcal{D} = \{\mathbf{x}_1, \dots, \mathbf{x}_N\}$, $\mathbf{x} \in \Omega_{\mathbf{X}} \subset \mathbb{R}^d$. We say that the event $\mathbf{x}_\ell \leq \mathbf{x}$ is true if and only if $\mathbf{x}_{\ell,i} \leq \mathbf{x}_i$ for each dimension $i = 1, \dots, d$. For notational convenience we use $\Omega_{\mathbf{X}}^- \in \mathbb{R}^d$ to denote the minimal point of $\Omega_{\mathbf{X}}$ (obtained by choosing the minimum of $\Omega_{\mathbf{X}}$ in each dimension), and let $\Omega_{\mathbf{X}}^+ \in \mathbb{R}^d$ be the corresponding maximal point. Then, the empirical CDF is defined as

$$G_N(\mathbf{x}) = \frac{1}{N} \sum_{\ell=1}^N \mathbf{1}\{\mathbf{x}_\ell \leq \mathbf{x}\}, \quad \mathbf{x} \in \Omega_{\mathbf{X}} \subset \mathbb{R}^d.$$

The goal is to find a representation of the empirical CDF of the form

$$F(\mathbf{x}) = \sum_{\ell_1=0}^{k} \dots \sum_{\ell_d=0}^{k} c_{\ell_1,\ell_2,\dots,\ell_d} \prod_{i=1}^{d} x_i^{\ell_i},$$

obtained by solving the optimization problem

minimize
$$\sum_{\ell=1}^{N} (G_N(\mathbf{x}_{\ell}) - F(\mathbf{x}_{\ell}))^2$$
subject to
$$\frac{\partial^d F(\mathbf{x})}{\partial x_1, \dots, \partial x_d} \ge 0 \quad \forall \mathbf{x} \in \Omega_{\mathbf{X}},$$

$$F(\Omega_{\mathbf{X}}^-) = 0 \text{ and } F(\Omega_{\mathbf{X}}^+) = 1.$$
(6)

The solution to this problem is the parameter-set that defines the joint CDF, and the density can be obtained by differentiation of the joint CDF. As in the univariate case, it is a quadratic optimization problem, that can be solved efficiently if the objective function is only evaluated on a set of grid-points.

Incorporating prior knowledge

There are real world situations, where the amount of available data is insufficient for accurate density estimation. The MoTBFs package includes implementations oriented to face these kinds of situations by allowing prior knowledge to be taken into account during density estimation. In Bayesian statistics (Bernardo and Smith, 2009), prior information is encoded as prior probability distributions over the parameters. As an example, consider the case of a random variable representing the body temperature of a patient in a hospital, and assume that the variable is normally distributed with mean μ and standard deviation σ . Prior knowledge could be provided in the form of a prior distribution on μ by establishing that $\mu \sim \mathcal{N}(37,0.1)$. However, in the case of MoTBF distributions, the parameters do not have a meaning in general. Therefore, there is no clear way in which a practitioner could provide prior information on any of the parameters, although some information could still be specified. For instance, in the body temperature example, the practitioner could choose not to give prior information on any single parameter, but instead provide a full distribution of the variable, reflecting his or her prior knowledge when no data is available. Such prior information could, e.g., include that the body temperature follows a normal distribution with mean 37 and standard deviation 0.5. This is the approach followed by Pérez-Bernabé et al. (2016), where prior knowledge is encoded as an MoTBF distribution over the random variable, which is then later combined with the MoTBF density learned

In order to represent such a prior distribution as an MoTBF, a sample is drawn from the prior – e.g., $\mathcal{N}(37,0.1)$ in the example above – and the sample is then used to learn an MoTBF using the methods previously described in this paper. Alternatively, one may also rely on direct translation schemes as described in, e.g., (Langseth et al., 2010; Cobb et al., 2006).

Given a prior MoTBF density f_{prior} over a variable X, Pérez-Bernabé et al. (2016) obtain a posterior density f_{post} by making a linear combination of the prior density and the density f_{data} learned from data, expressed as

$$f_{post} = w_p f_{prior} + w_d f_{data}$$
,

where w_p is the weight of f_{prior} and w_d is the weight of f_{data} , with $0 \le w_p \le 1$, $0 \le w_d \le 1$ and $w_p + w_d = 1$. The weights w_p and w_d reflect the way in which they explain the observed data. They are computed by measuring the difference between the log-likelihood produced by each one of the densities (f_{prior} and f_{data}) and the expected log-likelihood that would be produced by a randomly generated MoTBF. See (Pérez-Bernabé et al., 2016) for a detailed explanation of the procedure.

Probabilistic inference

In a Bayesian network, probabilistic inference is the task of computing the posterior distribution of any variable X given that some other variables Y have been observed to take some value y. Therefore, the goal of probabilistic inference is to compute the density f(x|y), where the value y is fixed.

An easy approach to estimate this density is by *forward sampling* (Henrion, 1988). The idea is to draw a sample of configurations of the variables in the Bayesian network by simulating each variable using its conditional distribution following a top-down order. Prior to sampling, all the densities in the network are restricted to the value $\mathbf{Y} = \mathbf{y}$ and when a variable is to be sampled, its conditional distribution is restricted to the values already obtained for its parents. Sampling is always possible since the variables are sampled following the topological ordering of the network. Once the sample has been obtained, $f(x|\mathbf{y})$ is estimated as a univariate MoTBF as we described before.

Package description and illustrative example

The MoTBFs package is designed using \$3 objects. The functions provided by the package implement the methods explained in the previous sections. The package implements functions for learning univariate, multidimensional, and conditional distributions, and provides support for parameter learning in hybrid Bayesian networks. In addition, it includes functions for incorporating prior knowledge when there is lack of data and for carrying out probabilistic inference. Moreover, two classes are incorporated in the package, "motbf" for defining univariate mixtures of truncated basis functions and "jointmotbf" for specifying multidimensional MoTBFs.

The functionality of the **MoTBFs** package is illustrated through an analysis carried out on a real world dataset. More precisely, we use the ecoli dataset (Lichman, 2013), which is provided along with the package. The dataset contains information about *Escherichia coli* and consists of n = 336 records, 8 input variables, and 1 output variable (the class). It is a bacterium of the genus *Escherichia* that is commonly found in the lower intestine of warm-blooded organisms. This dataset can be downloaded from http://archive.ics.uci.edu/ml/datasets/Ecoli.

To begin the analysis, the package and the data are loaded by

```
> install.packages("MoTBFs")
> library("MoTBFs")
> data("ecoli", package = "MoTBFs")
> str(ecoli)
'data.frame':
                  336 obs. of 9 variables:
$ Sequence.Name: chr "AAT_ECOLI" "ACEA_ECOLI" "ACEK_ECOLI" "ACKA_ECOLI" ...
$ mcg : num 0.49 0.07 0.56 0.59 0.23 0.67 0.29 0.21 0.2 0.42 ...
$ gvh
              : num 0.29 0.4 0.4 0.49 0.32 0.39 0.28 0.34 0.44 0.4 ...
              : chr "0.48" "0.48" "0.48" "0.48" ...
$ lip
              : chr "0.5" "0.5" "0.5" "0.5" ...
$ chg
              : num 0.56 0.54 0.49 0.52 0.55 0.36 0.44 0.51 0.46 0.56 ...
$ alm1
              : num 0.24 0.35 0.37 0.45 0.25 0.38 0.23 0.28 0.51 0.18 ...
              : num 0.35 0.44 0.46 0.36 0.35 0.46 0.34 0.39 0.57 0.3 ...
$ alm2
              : chr "cp" "cp" "cp" "cp" ...
$ class
```

The ecoli dataset is a data frame with 336 rows corresponding to proteins and 9 columns corresponding to variables. The dataset contains 4 discrete variables, stored as characters, and 5 continuous variables. The variables provide measurements of the cells used for predicting the localization site of proteins. The first variable, Sequence.Name, which is the accession number for the SWISS-PROT database, and the output variable class will not be used in this running example, and we will therefore remove them from the data frame. The discrete variables lip and chg are binary attributes, where

character numbers are used as states; "0.48" and "1", and "0.5" and "1", respectively. For validation purposes, the dataset is split into a training and a test set.

```
data <- ecoli[,-c(1,9)]
    set.seed(2)
    dataTT <- TrainingandTestData(data, percentage_test = 0.2)</pre>
>
    trainingData <- dataTT$Training</pre>
    testData <- dataTT$Test
```

The seed value determines the partitioning of the data into training and test, and is therefore key to reproducing the experiments. From now on, we will carry out all the analyses on the training data, leaving the test dataset for estimating the predictive capabilities of the learned models.

Our illustrative example basically consists of fitting MoTBF densities to a previously learned Bayesian network structure over the variables in the dataset. The structure can, for instance, be obtained, using the function hc() from the bnlearn package. This function returns a directed acyclic graph obtained from the dataset using a local search method. For the sake of simplicity, we have included the function LearningHC() in our package, which automatically converts into factors those columns that are non-numeric, before calling the function hc() in **bnlearn**. LearningHC() can also be used to discretize the dataset before calling hc(), but we are not using this functionality in the running example.

```
dag <- LearningHC(trainingData)</pre>
dag
```

Bayesian network learned via Score-based methods

```
[lip][alm1][mcg|lip:alm1][chg|lip][aac|alm1][gvh|mcg][alm2|gvh:lip:alm1]
nodes:
                                        8
arcs:
undirected arcs:
                                        0
directed arcs:
                                        8
average markov blanket size:
                                        3.14
                                        2.29
average neighbourhood size:
average branching factor:
                                        1.14
learning algorithm:
                                        Hill-Climbing
                                        BIC (cond. Gauss.)
score:
penalization coefficient:
                                        2.797356
tests used in the learning procedure:
optimized:
                                        TRUF
```

> plot(dag)

The network structure obtained is shown in Figure 1.

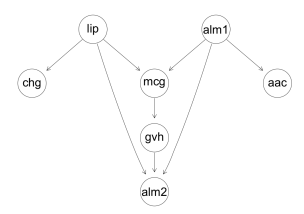


Figure 1: Directed acyclic graph learned from 80 percent of the ecoli dataset used as training data.

Before describing how to learn the MoTBF distributions associated with the network structure, we first present the basic functionality for learning different types of MoTBF representations, i.e., univariate, conditional, and joint MoTBF densities.

We illustrate the learning of a univariate MoTBF density by considering the continuous variable mcg.

```
> f1 <- univMoTBF(trainingData[,1], POTENTIAL_TYPE = "MTE", nparam = 13)
> f2 <- univMoTBF(trainingData[,1], POTENTIAL_TYPE = "MOP", nparam = 11)</pre>
```

The univMoTBF() function is used for learning univariate densities. The function is at the core of a collection of functions included in the package to learn densities of class "motbf" from data. Least squares optimization is used to minimize the mean squared error between the empirical cumulative distribution and the estimated MoTBF.

The function takes two mandatory arguments, data and POTENTIAL_TYPE, where the latter can either be "MOP" or "MTE" if polynomial or exponential basis functions should be used, respectively. univMoTBF() also accepts optional arguments: it is possible to specify the domain over which the model will be fitted, evalRange, the exact number of basis functions to be used, nparam, and the maximum number of parameters in the function, maxParam, which selects the best fit using the log-likelihood score. If nparam or maxParam are not given, then the Bayesian information criterion (BIC) (Schwarz, 1978) is used for scoring and function selection: it evaluates the two next functions and if the BIC value does not improve then the function with the best BIC score so far is returned.

An overview of the obtained results is shown via print() and summary().

```
R> print(f1)
[1] 31692.5765-19886.9389*exp(2*x)-3430.0930*exp(-2*x)+6520.8968*exp(4*x)
-91374.9603*exp(-4*x)-1269.7572*exp(6*x)+189285.9358*exp(-6*x)
+145.8260*exp(8*x)-186348.2886*exp(-8*x)-8.9962*exp(10*x)+94460.6774*exp(-10*x)
+0.2244*exp(12*x)-19787.1017*exp(-12*x)
> summarv(f2)
MoTBFs FOR UNIVARIATE DISTRIBUTIONS
Model:
0.0009+31.8820*x-1161.8247*x^2+16513.1506*x^3-121362.4662*x^4+529132.7157*x^5
-1434074.5145 \times x^6 + 2426253.6805 \times x^7 - 2482246.1917 \times x^8 + 1401057.0127 \times x^9 - 334304.5309 \times x^{10} + 1401057.0127 \times x
Class: motbf
Subclass: mop
Coefficients:
0.001 31.8821 -1161.825 16513.15 -121362.5 529132.7 -1434075 2426254
-2482246 1401057 -334304.5
Domain:
(0, 0.89)
Number of Iterations: 7
Processing Time: 0.002254009 secs
```

The object returned by univMoTBF() is a list containing several elements, including its mathematical expression and other hidden elements related to the learning task. The processing time is one of the values returned by this function and it can be extracted by \$Time. Although the learning process is always the same for a particular data sample, the processing time can vary inasmuch as it depends on the CPU.

```
> hist(trainingData[,1], prob = TRUE , main = "", xlab = "X")
> plot(f1, xlim = range(trainingData[,1]), col = "red", add = TRUE)
> plot(f2, xlim = range(trainingData[,1]), col = "blue", add = TRUE)
```

Figure 2 shows the model fits, provided by univMoTBF(), displayed using the generic method plot().

¹In order to save space and increase readability, we are only printing the 4 most significant digits in the examples in this paper.

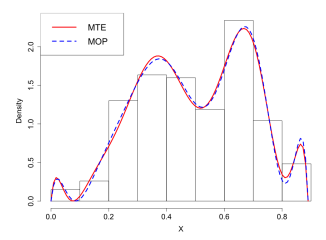


Figure 2: Univariate learning with blue dashed line for MOPs and red solid line for MTEs overlaying the histogram of the training data of the mcg variable.

To evaluate the predictive ability of the models we use the generic method as.function() developed for the "motbf" class to get the log-likelihood as well as BICMoTBF() to obtain the BIC score.

```
> sum(log(as.function(f1)(testData[,1])))
[1] 9.1945
> sum(log(as.function(f2)(testData[,1])))
[1] 8.7249
> BICMoTBF(f1,testData[,1])
[1] -20.2383
> BICMoTBF(f2,testData[,1])
[1] -16.5032
```

An alternative way to visually check the goodness of fit of the estimated models is to simulate a data sample from the learned functions and compare it with the training data. For doing this, we use the inverse transform method, a technique for generating random samples from a specific probability distribution based on evaluating the inverse of the CDF on a uniform random number, yielding a value for the random variable being sampled. This is done by function rMoTBF(). For the sake of reproducibility, we fix the seed for the random numbers to be used by the rMoTBF() function, which is set to 5 in this example. In the next code snippet, the previous function fitted with a polynomial basis, f2, will be used.

```
> set.seed(5)
> X <- rMoTBF(size = 400, fx = f2)
> ks.test(trainingData[,1], X)

Two-sample Kolmogorov-Smirnov test

data: trainingData[, 1] and X
D = 0.065167, p-value = 0.5018
alternative hypothesis: two-sided
```

In this example the two-sample Kolmogorov-Smirnov test is used. The p-value is notably above 0.05, so there is no evidence to reject the null hypothesis that both samples are drawn from the same population.

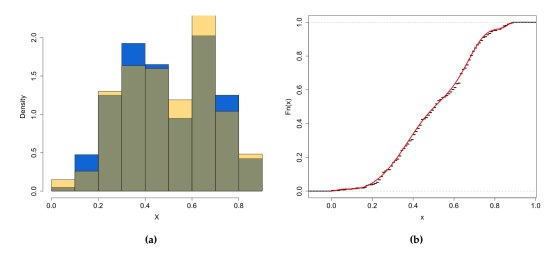


Figure 3: Histogram of variable mcg plotted over the histogram of the generated sample (a). Illustration of the inverse transform method (b), red solid line is the CDF of the generated sample, and black dashed line is the empirical CDF of the training data for variable mcg.

Figure 3 shows two plots comparing the training data of variable mcg and the sample simulated from the distribution learned using the same training data.

We can also manipulate the distributions with a collection of methods for class "motbf". Here is an example of the use of three of them, coef(), integralMoTBF(), and derivMoTBF().

```
> coef(f1)
[1] 3.1692e+04 -1.9886e+04 -3.4300e+03 6.5208e+03 -9.1374e+04
[6] -1.2697e+03 1.8928e+05 1.4582e+02 -1.8634e+05 -8.9962e+00
[11] 9.4460e+04 2.2449e-01 -1.9787e+04

> integralMoTBF(f2)
[1] 0.0009*x+15.9410*x^2-387.2749*x^3+4128.2876*x^4-24272.4932*x^5+88188.7859*x^6-204867.7877*x^7+303281.7100*x^8-275805.1324*x^9+140105.7012*x^10-30391.3209*x^11

> integralMoTBF(f2, min = min(trainingData[,1]), max = max(trainingData[,1]))
[1] 1

> derivMoTBF(f2)
[1] 31.8820-2323.6495*x+49539.4519*x^2-485449.8648*x^3+2645663.5790*x^4-8604447.0881*x^5+16983775.7655*x^6-19857969.5358*x^7+12609513.1160*x^8-3343045.3101*x^9
```

The learning process for multidimensional variables is similar to the previous one. The function parametersJointMoTBF() is used to solve the quadratic optimization problem. It returns Parameters, Range and Time, among other values. The function jointMoTBF() is used for obtaining the analytical expression, where the returned object is of class "jointmotbf". The expression is the only visible element, while the others can be retrieved using attributes(). In this example only two variables are used, mcg and alm1, in order to be able to plot the results.

> plot(P, type = "perspective", data = trainingData[,c(1,6)], orientation=c(60,20))

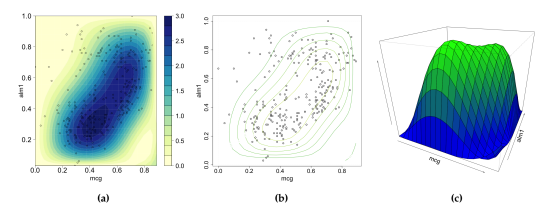


Figure 4: Filled contour (a), simple contour (b) and perspective (c) plots of the joint MoTBF f(Y, X) of variables mcg and alm1.

The plots in Figure 4 are generated using plot(), developed for objects of class "jointmothf". This function accepts optional arguments such as type, where one can choose between "perspective" and "contour", ranges, used to specify the plotting range, orientation, which indicates the orientation of the perspective graph, and filled for getting a filled contour plot.

The function print() can be used to obtain an expression of the learned joint density, while summary() yields a more thorough excerpt of the "jointmotbf" object.

> summary(P)

MoTBFs FOR MULTIVARIATE DISTRIBUTIONS

```
Model:
```

```
0.0355-0.2915*y+0.8351*y^2-0.9711*y^3+0.3939*y^4-2.9706*x+88.6144*x*y
-339.2388*x*y^2+451.7559*x*y^3-198.1810*x*y^4-9.2802*x^2+494.5320*x^2*y
-1392.3658*x^2*y^2+1175.9828*x^2*y^3-268.7925*x^2*y^4+35.3166*x^3
-1708.9509*x^3*y+5662.3268*x^3*y^2-5878.6313*x^3*y^3+1889.8292*x^3*y^4
-22.2566*x^4+1164.5531*x^4*y-4117.2185*x^4*y^2+4479.3097*x^4*y^3-1504.3348*x^4*y^4
```

Class: jointmotbf

Coefficients:

```
0.0355 -0.2915 0.8351 -0.9711 0.3939 -2.9706 88.6144 -339.2388 451.7559 -198.1811 -9.2802 494.5321 -1392.366 1175.983 -268.7926 35.3166 -1708.951 5662.327 -5878.631 1889.829 -22.2566 1164.553 -4117.219 4479.31 -1504.335
```

Domain x: (0, 0.89) Domain y: (0.03, 1)

Number of Iterations: 96

Processing Time: 1.144651 secs

As in the univariate case, the processing time, P\$Time, can vary depending on the CPU, but the learning outcome will always be the same for a specific data sample. The marginalJointMoTBF() function computes the marginals of joint densities. In this example we have two variables, so there are two marginal densities.

```
> marginalJointMoTBF(P, var = 1)
[1] 0.0031+1.6119*x+14.1692*x^2-23.7487*x^3+6.7540*x^4
> marginalJointMoTBF(P, var = 2)
[1] -0.2716+13.0463*y-32.4520*y^2+32.5558*y^3-12.8781*y^4
```

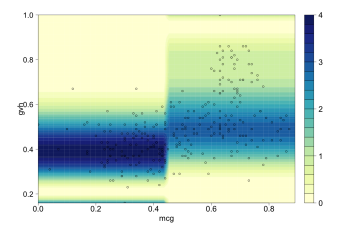


Figure 5: Conditional density of gvh given mcg.

The next step in our analysis is learning conditional densities, which is implemented by the function conditionalMethod(). Five of its arguments are compulsory: data, the dataset; nameParents, a character vector indicating the name of the parents; nameChild, a character string containing the name of the child; numIntervals, the maximum number of intervals for splitting the domain of the parent variables; POTENTIAL_TYPE, the type of basis function. Other arguments are optional, like maxParam, indicating the maximum number of parameters for each function, and s, the expert's relative confidence in any prior knowledge, and priorData if prior knowledge is incorporated in the analysis.

We will do the conditional analysis for only two variables in order to be able to make a 2-dimensional plot of the obtained results using plotConditional(). For example, taking into account the relationship found by the dag, we consider the child variable gvh with parent variable mcg.

Figure 5 shows the resulting conditional density (a MOP in this case) with the sample points overlaid. It can be noticed that the learning algorithm decides to split the domain of the parent into two intervals even though we have set the argument numIntervals to five. This is because the BIC score is not improved any further by splitting the domain into more than two intervals.

The last step is to learn the distributions tied to the Bayesian network learned previously. For doing this task, the MoTBFs_Learning() function of the **MoTBFs** package is used. The graph is a mandatory argument, that can be of class "bn", "graphNEL" or "network". Other mandatory arguments are the data, the maximum number of intervals for splitting the domain of the parents, and the type of basis function. The function also accepts additional arguments, but they are not listed here.

In the example, the DAG was obtained using the **bnlearn** package and therefore it is an object of class "bn". As an example, we will use a maximum of 4 intervals and "MTE" potentials when learning the densities (i.e. exponential basis functions).

```
Parent: alm1
                            Range: 0.33 < alm1 < 1
                           Range = "0.48"
Parent: lip
[1] -5.4620+3.3347*exp(2*x)+3.6655*exp(-2*x)-0.4237*exp(4*x)-1.0852*exp(-4*x)
Parent: lip Range = "1"
[1] -11.4070+6.0014*exp(2*x)+4.4488*exp(-2*x)-0.7109*exp(4*x)+2.3944*exp(-4*x)
Potential(gvh)
Parent: mcg
                           Range: 0 < mcg < 0.51
[1] -97.6780+34.1822*exp(2*x)-282.2167*exp(-2*x)-2.7274*exp(4*x)+2028.8986*exp(-4*x)
-0.12096*exp(6*x)-3609.5088*exp(-6*x)+0.0175*exp(8*x)+2069.4440*exp(-8*x)
                           Range: 0.51 < mcg < 0.89
[1] 685.6507-360.2896 \times \exp(2 \times x) + 252.1477 \times \exp(-2 \times x) + 79.5432 \times \exp(4 \times x) - 3074.9108 \times \exp(-4 \times x)
-8.3530*\exp(6*x)+4377.0877*\exp(-6*x)+0.3410*\exp(8*x)-2004.8518*\exp(-8*x)
Potential(lip)
0.9630 0.0369
Potential(chg)
Parent: lip
                           Range = "0.48"
                           Range = "1"
Parent: lip
0.8181 0.1818
Potential(aac)
Parent: alm1
                            Range: 0.03 < alm1 < 1
[1] -3742.3665+2528.6429*exp(2*x)+1860.6477*exp(-2*x)-894.0268*exp(4*x)
+2121.1450 \times \exp(-4 \times x) + 175.4771 \times \exp(6 \times x) - 3473.1126 \times \exp(-6 \times x) - 18.0836 \times \exp(8 \times x)
+1679.6126 \times \exp(-8 \times x) + 0.7634 \times \exp(10 \times x) - 238.3277 \times \exp(-10 \times x)
Potential(alm1)
[1] 158.6127-95.2652*exp(2*x)-56.9318*exp(-2*x)+25.4631*exp(4*x)-97.0824*exp(-4*x)
-3.1624 \times \exp(6 \times x) + 52.8052 \times \exp(-6 \times x) + 0.1480 \times \exp(8 \times x) + 17.6287 \times \exp(-8 \times x)
Potential(alm2)
Parent: alm1
                           Range: 0.03 < alm1 < 0.33
Parent: gvh
                           Range: 0.16 < gvh < 1
Parent: lip
                          Range = "0.48"
[1] 193.4903-112.6569*exp(2*x)-52.9058*exp(-2*x)+28.5291*exp(4*x)-185.8167*exp(-4*x)
-3.3698 \times \exp(6 \times x) + 155.0492 \times \exp(-6 \times x) + 0.1517 \times \exp(8 \times x) - 21.3863 \times \exp(-8 \times x)
Parent: alm1 Range: 0.33 < alm1 < 0.45
Parent: gvh
                          Range: 0.16 < gvh < 1
Parent: lip
                          Range = "0.48"
[1] 395.0603-217.7061*exp(2*x)+16.0642*exp(-2*x)+51.0248*exp(4*x)-1009.4114*exp(-4*x)
-5.6246 \times \exp(6 \times x) + 1224.5254 \times \exp(-6 \times x) + 0.2387 \times \exp(8 \times x) - 454.1703 \times \exp(-8 \times x)
Parent: alm1
                           Range: 0.45 < alm1 < 0.71
Parent: gvh
                           Range: 0.16 < gvh < 1
                          Range = "0.48"
Parent: lip
[1] -3.0260+3.1097*exp(2*x)+6.9062*exp(-2*x)-0.8013*exp(4*x)-12.5762*exp(-4*x)
+0.0578*exp(6*x)+6.3306*exp(-6*x)
               Range = "1"
Parent: lip
[1] 1.9648-0.2660*exp(2*x)-0.2660*exp(-2*x)
Parent: alm1 Range: 0.71 < alm1 < 1
Parent: gvh Range: 0.16 < gvh < 1
Parent: lin Range = "0.48"
                         Range = "0.48"
Parent: lip
[1] -1.2697+0.6353*exp(2*x)+0.6353*exp(-2*x)
Parent: lip
                           Range = "1"
[1] 1.0101+0*exp(2*x)
```

The results are reported using the printBN() function. Notice how nodes in the DAG with only discrete parents contain as many functions as configurations of the parents, nodes that have continuous parents have at most 4 functions for each parent and nodes that have mixed parents contain as many functions as configurations of the discrete parents times the number of regions into which the domain

of the continuous parents is split. The BIC criterion is used to decide the number of splitting points of the domain of the continuous parent nodes and to choose the number of basis functions used. The function BiC.MoTBFBN() can be used to compute the log-likelihood and the BIC score of a dataset given the Bayesian network.

```
> bic <- BiC.MoTBFBN(bn, data = testData)
> attributes(bic)
$names
[1] "LogLikelihood" "BIC"
> bic$LogLikelihood
[1] 173.5496
> bic$BIC
[1] -51.40147
```

We will now exemplify the use of prior knowledge in the learning process. In order to illustrate the approach, we first select a small subset of the Ecoli dataset using TrainingandTestData(). In the next example the percentage of the test data is 99%, which means the training data is only 1% of the full dataset.

```
> set.seed(4)
> dataTT <- TrainingandTestData(data, percentage_test = 0.99)
> trainingData <- dataTT$Training
> testData <- dataTT$Test
> nrow(trainingData)
[1] 13
```

There are 13 entries in the training dataset. We are going to fit MoTBFs with and without prior information. To generate an artificial prior dataset the generateNormalPriorData() function can be used.

Learning univariate and conditional distributions and Bayesian networks can be done using the functions learnMoTBFpriorInformation() and MoTBFs_Learning(). The arguments for these functions are the same as previously explained and, in addition, it is necessary to specify the expert confidence in the prior knowledge, s, and the prior dataset priorData. Argument s takes values on the interval [0, N], where N is the sample size, and is used to synchronize the support of the prior knowledge and the sample. We refer the reader to (Pérez-Bernabé et al., 2016) for the details. In this example we will use the aac variable from the data set, have s = 5 as confidence level, and set "MOP" as potential type.

```
> f <- learnMoTBFpriorInformation(priorData$aac, trainingData$aac,</pre>
        s = 5, POTENTIAL_TYPE = "MOP")
> attributes(f)
$names
[1] "coeffs"
                        "posteriorFunction" "priorFunction"
[4] "dataFunction"
                        "domain"
> print(f)
$coeffs
[1] 0.5509206 0.4490794
$posteriorFunction
[1] 0.2911+2.7626*x+4.3721*x^2-30.4218*x^3-0.8389*x^4+965.8444*x^5-4183.3860*x^6
+7735.8433*x^7-7321.9450*x^8+3498.9449*x^9-671.5104*x^10
$priorFunction
[1] 0.0673+1.7639*x+11.8532*x^2-55.2199*x^3-1.5228*x^4+1753.1465*x^5-7593.4470*x^6
+14041.6676*x^7-13290.3826*x^8+6351.0879*x^9-1218.8879*x^10
```

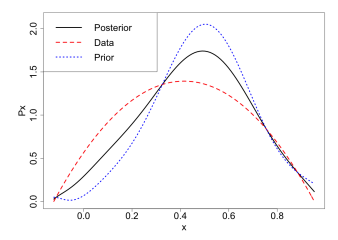


Figure 6: Univariate density estimation using prior knowledge. Solid black, dashed red and dotted blue lines represent the posterior, the data, and the prior function, respectively.

```
$dataFunction
[1] 0.5656+3.9878*x-4.8055*x^2

$domain
[1] -0.1232  0.9531

> sum(log(as.function(f$posteriorFunction)(testData$aac)))
[1] 134.566
> sum(log(as.function(f$dataFunction)(testData$aac)))
[1] 78.96405
```

The best model, taking into account the log-likelihood, is the MoTBF which uses the prior data, f\$posteriorFunction. The generic method plot() for "motbf" object is used for displaying the functions depicted in Figure 6.

```
> plot(f$posteriorFunction, xlim = f$domain, ylim = c(0,2.1))
> plot(f$dataFunction, xlim = f$domain, add = TRUE, col = 2)
> plot(f$priorFunction, xlim = f$domain, add = TRUE, col = 4)
```

The last step is to incorporate the prior knowledge in the full Bayesian network. For this analysis we are not going to print out the results, because the structure is similar to the previous Bayesian network representations. As an example, we will use numIntervals = 2, POTENTIAL_TYPE = "MOP", and s = 5.

Looking at the log-likelihood corresponding to the network with and without prior data, we can see that, in this example, incorporating prior knowledge is better when data is scarce.

After a Bayesian network has been constructed, the **MoTBFs** package can be used to obtain the conditional density of any variable in the network given that some other variables have been observed. The conditional distribution is obtained by forward sampling. As an example, consider a network estimated from the ecoli dataset:

```
> data("ecoli", package = "MoTBFs")
> data <- ecoli[,-c(1,9)]
> dag <- LearningHC(data)
> bn <- MoTBFs_Learning(dag, data = data, numIntervals = 4, POTENTIAL_TYPE = "MTE")</pre>
```

The observed values are specified using a data frame. In the example, we are assuming that we want to compute the conditional density of alm2 given that lip="0.48", alm1 = 0.55 and gvh = 1. This is achieved by using the function forward_sampling were we have chosen a sample size equal to 10 specified by parameter size = 10.

```
> obs <- data.frame(lip = "0.48", alm1 = 0.55, gvh = 1, stringsAsFactors=FALSE)
> node <- "alm2"
> set.seed(5)
> forward_sampling(bn, dag, target = node, evi = obs, size = 10, maxParam = 15)
Processing Time: 0.209545850753784secs
[1] -4.3738+2.4552*exp(2*x)+1.5054*exp(-2*x)-0.2392*exp(4*x)+2.0045*exp(-4*x)
$sample
mcg gvh lip chg
                    aac alm1
                                  alm2
1 0.7450156 1 0.48 0.5 0.3564026 0.55 0.53709571
2 0.2493266 1 0.48 0.5 0.4873396 0.55 0.55395502
3 0.4075408 1 0.48 0.5 0.5052861 0.55 0.74832025
4 0.3205169 1 0.48 0.5 0.4844040 0.55 0.82546475
5 0.4448844 1 0.48 0.5 0.4248584 0.55 0.07672959
6 0.5717975 1 0.48 0.5 0.4611889 0.55 0.35412707
7 0.7548104 1 0.48 0.5 0.5695499 0.55 0.68322911
8 0.5475842 1 0.48 0.5 0.7797320 0.55 0.45343460
9 0.3048408 1 0.48 0.5 0.5251502 0.55 0.73064241
```

The output consists of the posterior density and the sample from which the density parameters were estimated.

Conclusions

This paper has presented the R package **MoTBFs** for learning Mixtures of Truncated Basis Functions in hybrid Bayesian networks. It provides a free and accessible implementation of algorithms for learning the parameters of MoTBFs densities as well as MoTBF-based Bayesian networks relying on state-of-the-art learning algorithms.

The **MoTBFs** package is designed to provide the required implementation to tackle experimental data analysis with both discrete and continuous data. Not only does the package provide methods for learning distributions from data, it also includes a set of auxiliary functions to perform descriptive statistics as well as other basic operations like inference using forward sampling.

The **MoTBFs** package expands the functionality for handling hybrid Bayesian networks already provided by packages **bnlearn** and **HydeNet**, by implementing MoTBF distributions, resulting in unrestricted network structures, regardless of the discrete or continuous nature of the variables involved, and by providing methods for building models from data that are compatible with exact inference methods. The package **MoTBFs** is complementary to **abn** in the sense that the former is based on MoTBF densities, which do not belong to the exponential family.

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