# The Estimation of Distributions and the **Minimum Relative Entropy Principle**

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#### Abstract

Estimation of Distribution Algorithms (EDA) have been proposed as an extension of genetic algorithms. In this paper the relation of EDA to algorithms developed in statistics, artificial intelligence, and statistical physics is explained. The major design issues are discussed within a general interdisciplinary framework. It is shown that maximum entropy approximations play a crucial role. All proposed algorithms try to minimize the Kullback-Leibler divergence KLD between the unknown distribution  $p(\mathbf{x})$  and a class  $q(\mathbf{x})$  of approximations. The Kullback-Leibler divergence is not symmetric. Approximations which suppose that the function to be optimized is additively decomposed (ADF) minimize  $\tilde{K}LD(q|p)$ , the methods which learn the approximate model from data minimize KLD(p|q). This minimization is identical to maximizing the *log*likelihood. In the paper three classes of algorithms are discussed. FDA uses the ADF to compute an approximate factorization of the unknown distribution. The factors are marginal distributions, whose values are computed from samples. The Bethe-Kikuchi approach developed in statistical physics uses bi-variate or higher order marginals. The values of the marginals are computed from a difficult minimization problem. The third class learns the factorization from the data. We analyze our learning algorithm *LFDA* in detail. It is shown that learning is faced with two problems: first, to detect the important dependencies between the variables, and second, to create an acyclic Bayesian network of bounded clique size.

## **Keywords**

Estimation of distributions, Boltzmann distribution, factorization of distributions, maximum entropy principle, minimum relative entropy, minimum log-likelihood ratio, Bayesian information criterion, Bethe approximation.

# 1 Introduction

The Estimation of Distribution (EDA) family of population based search algorithms was introduced by Mühlenbein and Paaß (1996) as an an extension of genetic algorithms.<sup>1</sup> The following observations lead to this proposal. First, genetic algorithm have difficulties to optimize deceptive and non-separable functions, and second, the search distributions implicitly generated by recombination and crossover can be extended to include the correlation of the variables in samples of high fitness values.

<sup>&</sup>lt;sup>1</sup>In (Mühlenbein and Paaß, 1996) they have been named conditional distribution algorithms.

EDA uses probability distributions derived from the function to be optimized to generate search points instead of crossover and mutation as done by genetic algorithms. The other parts of the algorithms are identical. In both cases a population of points is used and points with good fitness are selected either to estimate a search distribution or to be used for crossover and mutation.

In (Mühlenbein and Paaß, 1996) the distribution has been estimated by computationally intensive Monte Carlo methods. The distribution was restricted to tree-like structures. It has been shown by Mühlenbein et al. (1999) that simpler and more effective methods exist which use a general factorization of the distribution.

The family of EDA algorithms can be understood and further developed without the background of genetic algorithms. The problem to estimate empirical distributions has been investigated independently in several scientific disciplines. In this paper we will show how results in statistics, belief networks and statistical physics can be used to understand and further develop EDA. In fact, an interdisciplinary research effort is well under way which cross-fertilizes the different disciplines.

Unfortunately each discipline uses a different language, has a slightly different application, and has developed different algorithms. In EDA we have to sample from a distribution, in belief networks one computes a single marginal distribution  $p(\mathbf{y}|\mathbf{z})$  for new evidence  $\mathbf{z}$ , and statistical physicists want to compute the free energy of a Boltzmann distribution. Thus the algorithms developed for belief networks concentrate on computing a single marginal distribution, whereas for EDA we want to sample  $p(\mathbf{x})$  in areas of high fitness values, i.e. we are interested in a sampling method which generates points with a high value of  $p(\mathbf{x})$ . All disciplines are interested in developing fast algorithms to compute marginal distributions. The foundation of the theory is the same for all disciplines. It is based on graphical models and their decomposition. We hope that the readers are interested to accompany us on our journey through the different disciplines. We will leave out a discussion of the approaches in probabilistic logic to simplify the presentation.

Today two major branches of EDA can be distinguished. In the first branch the factorization of the distribution is computed from the structure of the function to be optimized, in the second one the structure is computed from the correlations of the data. The second branch has been derived from the theory of belief networks (Jordan, 1999). For large real life applications often a hybrid between these two approaches is most successful (Mühlenbein and Mahnig, 2002a).

The paper is intended as a short introduction to the theory of EDA. We will only consider discrete variables. It is not intended as a survey of ongoing research. Here an excellent overview is already available (Larrañaga and Lozano, 2002).

The outline of the paper is as follows. In section 2 the basic steps to derive the Factorized Distribution Algorithm *FDA* are recapitulated. A factorization theorem will be discussed which uses the structure of the function to be optimized to factor the distribution. In section 2.2 the junction tree algorithm is described which computes an exact factorization by decomposing graphical models. Unfortunately many important problems do not allow an exact factorization useful for numerical computations. In section 3 the estimation problem is generalized. Here the concept of *maximum entropy* distributions is explained.

In section 4 the methods developed in statistical physics are described. In this approach the marginals are not computed from data, but from the known expression of the function. In section 5 the learning of models from samples of high fitness values is described. Then we compare the different approaches presented using a simple

example. In section 7 our learning algorithm *LFDA* is analyzed and its behavior and performance compared to *FDA* is investigated.

# 2 Factorization of the Search Distribution

EDA has been derived from a search distribution point of view. We just recapitulate the major steps published in (Mühlenbein et al., 1999; Mühlenbein and Mahnig, 2000; Mühlenbein and Mahnig, 2002a). We will use in the paper the following notation. Capital letters denote variables, lower cases instances of variables. If the distinction between variables and instances is not necessary, we will use lower case letters. Vectors are denoted by  $\mathbf{x}$ , a single variable by  $x_i$ .

Let a function  $f : \mathbf{X} \to \mathbb{R}_{\geq 0}$  be given. We consider the optimization problem

$$\mathbf{x}_{opt} = \operatorname*{argmax} f(\mathbf{x}) \tag{1}$$

A good candidate for optimization using a search distribution is the Boltzmann distribution.

**Definition 1.** For  $\beta \ge 0$  define the Boltzmann distribution<sup>2</sup> of a function  $f(\mathbf{x})$  as

$$p_{\beta}(\mathbf{x}) := \frac{e^{\beta f(\mathbf{x})}}{\sum_{\mathbf{y}} e^{\beta f(\mathbf{y})}} =: \frac{e^{\beta f(\mathbf{x})}}{Z_f(\beta)}$$
(2)

where  $Z_f(\beta)$  is the partition function. To simplify the notation  $\beta$  and/or f might be omitted.

The Boltzmann distribution concentrates with increasing  $\beta$  around the global optima of the function. Obviously, the distribution converges for  $\beta \to \infty$  to a distribution where only the optima have a probability greater than 0 (Mühlenbein and Mahnig, 2002b). Therefore, if it were possible to sample efficiently from this distribution for arbitrary  $\beta$ , optimization would be an easy task. But the computation of the partition function needs an exponential effort for a problem of *n* variables. We have therefore proposed an algorithm which incrementally computes the Boltzmann distribution from empirical data using Boltzmann selection.

**Definition 2.** Given a distribution p and a selection parameter  $\Delta\beta$ , Boltzmann selection calculates the distribution for selecting points according to

$$p^{s}(\mathbf{x}) = \frac{p(\mathbf{x})e^{\Delta\beta f(\mathbf{x})}}{\sum_{u} p(\mathbf{y})e^{\Delta\beta f(\mathbf{y})}}$$
(3)

The following theorem is easy to prove.

**Theorem 3.** If  $p_{\beta}(\mathbf{x})$  is a Boltzmann distribution, then  $p^{s}(\mathbf{x})$  is a Boltzmann distribution with inverse temperature  $\beta(t+1) = \beta(t) + \Delta\beta(t)$ .

Algorithm 1 describes BEDA, the Boltzmann Estimated Distribution Algorithm.

*BEDA* is a conceptional algorithm, because the calculation of the distribution requires a sum over exponentially many terms. In the next section we transform *BEDA* into a practical numerical algorithm.

<sup>&</sup>lt;sup>2</sup>The Boltzmann distribution is usually defined as  $e^{-\frac{E(\mathbf{x})}{T}}/Z$ . The term E(x) is called the energy and  $T = 1/\beta$  the temperature. We use the inverse temperature  $\beta$  instead of the temperature.

Algorithm 1: BEDA – Boltzmann Estimated Distribution Algorithm

 $\begin{array}{ll} 1 & t \equiv 1. & \mbox{Generate } N \mbox{ points according to the uniform distribution} \\ p(\mathbf{x},0) \mbox{ with } \beta(0) = 0. \end{array}$   $\begin{array}{ll} 2 & \mbox{do} \left\{ \\ 3 & \mbox{With a given } \Delta\beta(t) > 0, \mbox{let} \\ & p^s(\mathbf{x},t) = \frac{p(\mathbf{x},t)e^{\Delta\beta(t)f(\mathbf{x})}}{\sum_{\mathbf{y}} p(\mathbf{y},t)e^{\Delta\beta(t)f(\mathbf{y})}}. \end{array}$   $\begin{array}{ll} 4 & \mbox{Generate } N \mbox{ new points according to the distribution } p(\mathbf{x},t+1) = \\ & p^s(\mathbf{x},t). \end{array}$   $\begin{array}{ll} 5 & t \equiv t + 1. \\ 6 & \end{array} \right\} \mbox{until (stopping criterion reached)} \end{array}$ 

## 2.1 Factorization of the distribution

In this section an efficient numerical algorithm is derived if the fitness function is additively decomposed.

**Definition 4.** Let  $s_1, \ldots, s_m$  be index sets,  $s_i \subseteq \{1, \ldots, n\}$ . Let  $f_i$  be functions depending only on the variables  $x_j$  with  $j \in s_i$ . Then

$$f(\mathbf{x}) = \sum_{i=1}^{m} f_i(\mathbf{x}_{s_i})$$
(4)

is an additive decomposition of the fitness function (ADF).

**Definition 5.** Let an ADF be given. Then the graph  $G_{ADF}^3$  is defined as follows: The vertices represent the variables of the ADF. Two vertices are connected by an arc iff the corresponding variables are contained in a common sub-function.

Given an ADF we want to estimate the Boltzmann distribution (2) using a product of marginals. We need the following sets:

**Definition 6.** Given  $s_1, \ldots, s_m$ , we define for  $i = 1, \ldots, m$  the sets  $d_i$ ,  $b_i$  and  $c_i$ :

$$d_{i} := \bigcup_{j=1}^{i} s_{j}, \qquad b_{i} := s_{i} \setminus d_{i-1}, \qquad c_{i} := s_{i} \cap d_{i-1}$$
(5)

We demand  $d_m = \{1, ..., n\}$  and set  $d_0 = \emptyset$ . In the theory of decomposable graphs,  $d_i$  are called histories,  $b_i$  residuals and  $c_i$  separators (Lauritzen, 1996).

The next definition is stated a bit informally.

**Definition 7.** A set of marginal distributions  $\tilde{q}(\mathbf{x}_{b_i}, \mathbf{x}_{c_i})$  is called consistent if the marginal distributions fulfill the laws of probability, e.g.

$$\sum_{\mathbf{x}_{b_i}, \mathbf{x}_{c_i}} \tilde{q}(\mathbf{x}_{b_i}, \mathbf{x}_{c_i}) = 1$$
(6)

$$\sum_{\mathbf{x}_{b_i}} \tilde{q}(\mathbf{x}_{b_i}, \mathbf{x}_{c_i}) = \tilde{q}(\mathbf{x}_{c_i})$$
(7)

<sup>&</sup>lt;sup>3</sup>Xiang et al. (1997) call it a decomposable Markov graph.

**Proposition 8.** Let a consistent set of marginal distributions  $\tilde{q}(\mathbf{x}_{b_i}, \mathbf{x}_{c_i})$  be given. If  $b_i \neq \emptyset$  then \_\_\_\_\_\_m

$$q_{\beta}(\mathbf{x}) = \prod_{i=1}^{m} \tilde{q}_{\beta}(\mathbf{x}_{b_i} | \mathbf{x}_{c_i})$$
(8)

defines a valid distribution ( $\sum q_{\beta}(\mathbf{x}) = 1$ ). Furthermore

$$q_{\beta}(\mathbf{x}_{b_i}|\mathbf{x}_{c_i}) = \tilde{q}_{\beta}(\mathbf{x}_{b_i}|\mathbf{x}_{c_i}), \quad i = 1, \dots m$$
(9)

whereas in general

$$q_{\beta}(\mathbf{x}_{b_i}, \mathbf{x}_{c_i}) \neq \tilde{q}_{\beta}(\mathbf{x}_{b_i}, \mathbf{x}_{c_i}), \quad i = 1, \dots m$$
(10)

The proof follows from the definition of marginal probabilities. The proof of equation (9) is somewhat technical, but straightforward. The inequality (10) is often overlooked. We need an additional constraint in order that also the marginal distributions become equal. This has been proven by Mühlenbein et al. (1999).

**Theorem 9 (Factorization Theorem).** Let  $f(\mathbf{x}) = \sum_{i=1}^{m} f_{s_i}(\mathbf{x})$  be an additive decomposition. If

$$\forall i = 1, \dots, m; \quad b_i \neq \emptyset \tag{11}$$

$$\forall i \ge 2 \ \exists j < i \ \text{such that} \ c_i \subseteq s_j \tag{12}$$

then

$$p_{\beta}(\mathbf{x}) = \prod_{i=1}^{m} p_{\beta}(\mathbf{x}_{b_i} | \mathbf{x}_{c_i}) = \frac{\prod_{i=1}^{m} p_{\beta}(\mathbf{x}_{b_i}, \mathbf{x}_{c_i})}{\prod_{i=2}^{m} p_{\beta}(\mathbf{x}_{c_i})}$$
(13)

There exists always a factorization fulfilling the assumptions of the factorization theorem. We just mention

$$p(\mathbf{x}) = p(x_1)p(x_2|x_1)p(x_3|x_1, x_2)\cdots p(x_n|x_1, \dots, x_{n-1})$$
(14)

But this factorization uses marginal distributions of size O(n), thus the computation is exponential in n. Therefore we are looking for factorizations where the size of the marginals is bounded, independent of n.

**Definition 10.** The constraint defined by equation (12) is called the running intersection property (*RIP*). The factorization is polynomially bounded (*PBF*) if the size of the sets  $\{b_i, c_i\}$  is bounded by a constant independent of n.

The connection of the factorization theorem to research done in non-sequential dynamic programming is not well known. If a factorization with *RIP* is possible, we can compute the maximum directly. In fact, the following maximization theorem has been proven earlier than the factorization theorem by Bertelé and Brioschi (1972).

**Theorem 11 (Maximization Theorem).** Let the assumptions of the factorization theorem be fulfilled. If the factorization is polynomially bounded, then  $\max p_{\beta}(\mathbf{x})$  and  $\operatorname{argmax} p_{\beta}(\mathbf{x})$  can be computed recursively in polynomial time.

A proof of the theorem using graphical models can be found in Jordan (1999). The basic idea is that maximization can be done in the same manner as marginalization. We have

$$\max_{a,b,c} f_1(a,b) f_2(b,c) = \max_{a,b} f_1(a,b) \max_c f_2(b,c)$$

For later use we just remark that any *FDA* factorization can be transformed into an acyclic Bayesian network (acBN) (see equation (14)).

$$q_{\beta}(\mathbf{x}) = \prod_{i=1}^{n} q_{\beta}(x_i | \pi_i)$$
(15)

where  $\pi_i$  are called the parents of  $x_i$ .

We next describe a well-known algorithm to obtain a factorization with marginals of small size and fulfilling the *RIP* given an arbitrary graph.

# 2.2 Computing a factorization by junction trees

The algorithm is defined for any graphical models G, an example is  $G_{ADF}$ . In order to find the separators  $c_i$  the method computes cliques and generates a junction tree J. A junction tree is an undirected tree the nodes of which are clusters of variables. The clusters satisfy the *junction property*: For any two clusters a and b and any cluster h on the unique path between a and b in the junction tree

$$a \cap b \subseteq h$$
 (16)

The edges between the clusters are labeled with the intersection of the adjacent clusters; we call these labels *separating sets* or *separators*.

**Remark:** The junction property is equivalent to the running intersection property (12).

A junction tree is constructed from the graphical model by the following steps:

- **Triangulating the graph** *G*: A graph is triangulated if it contains no chord-less circle with more than three vertices. An algorithm for adding the necessary edges is described in (Huang and Darwiche, 1996).
- **Finding the cliques:** A clique C in a graph is a maximal totally connected subgraph. That means that in C every node is connected to every other node in C, and there is no clique C' which contains C.
- **Generating the clusters:** For each clique generate a cluster containing its variables. This cluster will become a node of the junction tree *J*.
- **Building the junction tree:** Find pairs of clusters with maximal intersection and connect them. Label the edge with the separating set. Repeat this until the tree is complete.

This results in a tree which fulfills the junction property. There is plenty of literature available about this method, e. g. (Lauritzen, 1996; Huang and Darwiche, 1996; Jensen and Jensen, 1994).

A simple example to demonstrate this method is a circular graph G. It can be triangulated by connecting one node with all other nodes. The resulting junction tree is shown for 8 nodes in figure 1. The distribution can be factored into the cliques given by the clusters of the junction tree.

$$p(\mathbf{x}) = p(x_1, x_2, x_8) \prod_{i=3}^{7} p(x_i | x_{i-1}, x_8)$$
(17)

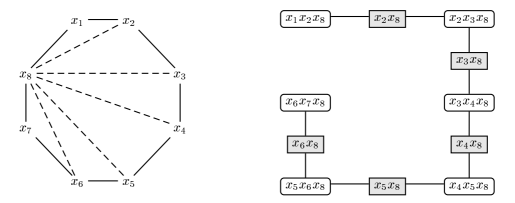


Figure 1: Graph model with triangulation and junction tree for a 1-D bi-variate circle. The left figure shows the graph G; the dashed lines are inserted for the triangulation. The cliques of the triangulated graph are the clusters of the junction tree J (right figure, white boxes). The separators are the shaded boxes.

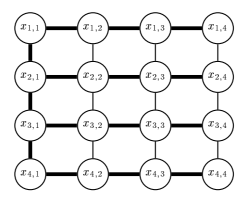


Figure 2: Graph model for a 2-D grid. The thick lines give a possible spanning tree

However, the junction tree contains non-local marginal distributions of order three.<sup>4</sup> One can show that there exists no exact factorization of a 1-D circle using bi-variate marginals only. For EDA this poses no problem in principle, because the marginals can be computed from the selected sample. But for larger marginals more samples are needed for a reliable estimate. If the graph contains many loops, the junction tree might be difficult to compute or might contain marginals with an exponential number of variables. We will investigate this problem for a 2-D grid.

# 2.3 The 2-D grid

Let there be a 2-D grid of variables  $x_{i,j}$ , i, j = 1, ..., n. Let the fitness function be composed of the sub-functions of pairs of neighboring variables,  $x_{i,j}$ ,  $x_{i+1,j}$  and  $x_{i,j}$ ,  $x_{i,j+1}$ . The goal is to compute a factorized distribution which is a good approximation to the true distribution.

An exact factorization can be found with a junction tree. The difficulty of the computation lies in the triangulation of the graphical model. One valid triangulation uses

<sup>&</sup>lt;sup>4</sup>Local marginals are defined in a 1-D neighborhood like  $p(x_{i-1}, x_i, x_{i+1})$ 



Figure 3: A  $3 \times 3$  grid and its factorization using (19).

the rows of the grid. Each variable is connected with all variables in the same row and the neighboring rows. This adds O(n) edges to the graph. The cliques in the junction tree consist of pairs of neighboring rows and have size 2n. Thus the exact factorization is not polynomially bounded.

Therefore it is advisable to look for approximations. A very straightforward approximation is to leave out some of the marginals and build a spanning tree of the grid. This could be the vertical edges in the first column and all the horizontal edges, forming a big "E" (see thick lines in figure 2).

Given this subset of the edges and disregarding the rest, we can define the following distribution:

$$q(\mathbf{x}) = p(x_{1,1}, x_{2,1}) \prod_{i=2}^{n-1} p(x_{i+1,1} | x_{i,1}) \prod_{i=1}^{n} \prod_{j=1}^{n-1} p(x_{i,j+1} | x_{i,j})$$
(18)

This is a valid probability distribution insofar as it sums up to 1 and complies with the regarded marginals. But obviously the choice of some marginals, while abandoning the rest, retains the stain of arbitrariness. Another possibility which regards all the given marginals, consists of combining blocks of four variables  $(x_{i,j}, x_{i+1,j}, x_{i,j+1}, x_{i+1,j+1})$ . The complete distribution can then be built up by:

$$q(\mathbf{x}) = p(x_{1,1}, x_{2,1}, x_{1,2}, x_{2,2}) \prod_{i=2}^{n-1} p(x_{i+1,1}, x_{i+1,2} | x_{i,1}, x_{i,2})$$
$$\prod_{j=2}^{n-1} p(x_{1,j+1}, x_{2,j+1} | x_{1,j}, x_{2,j}) \prod_{i=2}^{n-1} \prod_{j=2}^{n-1} p(x_{i+1,j+1} | x_{i,j}, x_{i+1,j}, x_{i,j+1})$$
(19)

However, the factorization (19) violates the running intersection property (12). It reproduces the given marginals only in the first tetra-variate row and column, but not in the areas where the running intersection property is violated. For simplicity we assume a  $3 \times 3$  grid. For abbreviation, we enumerate the variables with 1 through 9 (figure 3). We can calculate the marginal distribution  $q(x_6, x_9)$  as follows:

$$q(x_{6}, x_{9}) = \sum_{x_{5}, x_{8}} p(x_{9}|x_{5}, x_{6}, x_{8}) \sum_{x_{4}} p(x_{8}|x_{4}, x_{5}) \sum_{x_{2}} p(x_{6}|x_{2}, x_{5}) p(x_{2}, x_{4}, x_{5})$$

$$\approx \sum_{x_{5}, x_{8}} p(x_{9}|x_{5}, x_{6}, x_{8}) \sum_{x_{4}} p(x_{8}|x_{4}, x_{5}) p(x_{4}, x_{5}, x_{6})$$

$$\approx \sum_{x_{5}, x_{8}} p(x_{9}|x_{5}, x_{6}, x_{8}) p(x_{5}, x_{6}, x_{8})$$
(20)
(21)

$$\sum_{x_5, x_8} F(x_5|x_5, x_6, x_8)F(x_5, x_6, x_8)$$

$$= p(x_6, x_9)$$

The approximations (20) and (21) would be correct if

$$p(x_6|x_2, x_5) = p(x_6|x_2, x_4, x_5)$$
  
$$p(x_8|x_4, x_5) = p(x_8|x_4, x_5, x_6)$$

However, these equations cannot be deduced from the graphical model. The first approximation, for instance, is only true if  $x_6$  and  $x_4$  are conditionally independent given  $(x_2, x_5)$ . But this is not the case. There exists a path between  $x_4$  and  $x_6$  which does not have  $x_2$  or  $x_5$  as a node.

The optimal decomposition of a grid has already been investigated for non-serial dynamic programming by Martelli and Montari (1972). We next describe our factorized distribution algorithm *FDA* which also works with approximate factorizations.

## 2.4 The Factorized Distribution Algorithm FDA

If the factorization violates the assumption of the factorization theorem, then non-serial dynamic programming does not work. But an algorithm which estimates the marginals from samples might still find the optimum. One only has to compute a good approximate factorization given the graph  $G_{ADF}$ . We first describe our algorithm *FDA*.

# Algorithm 2: FDA – Factorized Distribution Algorithm

- Calculate b<sub>i</sub> and c<sub>i</sub> by the Sub-function Merger Algorithm.
   t ← 1. Generate an initial population with N individuals from the uniform distribution.
   do {
   Select M ≤ N individuals using Boltzmann selection<sup>a</sup> (see Def. 2).
   Estimate the conditional probabilities p(x<sub>bi</sub> | x<sub>ci</sub>, t) from the selected points.
  - 6 Generate new points according to  $p(\mathbf{x}, t+1) = \prod_{i=1}^{m} p(\mathbf{x}_{b_i} | \mathbf{x}_{c_i}, t)$ .
  - 7  $t \Leftarrow t+1.$
  - *8* } **until** (stopping criterion reached)

$$s_1 = \{1, 2\}, s_2 = \{2, 3\}, s_3 = \{1, 3\}$$

<sup>&</sup>lt;sup>a</sup>The algorithm works with any selection method

We next describe the sub-function merger algorithm which computes the *FDA* factorization. Let us first discuss the assumption  $b_i \neq \emptyset$  of the factorization theorem. This assumption is violated already for the loop

All possible sequences end in  $b_3 = \emptyset$  because the variables of the sub-function left are already contained in the two previous sets. One possibility to solve this problem is to choose only a subset of the  $s_i$  and disregard the others; in our example, we can use the factorization  $q(\mathbf{x}) = p(x_1, x_2)p(x_3|x_2)$  using  $s_1$  and  $s_2$ . An exact factorization is  $p(\mathbf{x}) = p(x_1, x_2)p(x_3|x_2, x_1)$ . This factorization will be generated if the two subfunctions  $s_2$  and  $s_3$  are merged. This observation leads to the idea to compute approximate factorizations by merging of sub-functions<sup>5</sup>.

#### **Algorithm 3: Sub-function Merger**

1  $\mathcal{S} \Leftarrow \{s_1, \ldots, s_m\}$  $2 \quad j \Leftarrow 1$ 3 while  $\tilde{d}_j \neq \{1, ..., n\}$  do { Chose an  $s_i \in S$  to be added 4  $\mathcal{S} \Leftarrow \mathcal{S} \setminus \{s_i\}$ 5 Let the indices of the new variables in  $s_i$  be  $b_i = \{k_1, \ldots, k_l\}$ 6 for  $\lambda = 1$  to l do { 7  $\delta_{\lambda} \leftarrow \{k \in \tilde{d}_{j-1} | (x_k, x_{k_{\lambda}}) \in G_{ADF} \}$ 8 9 } 10 for  $\lambda = 1$  to l do { **if** exists  $\lambda' \neq \lambda$  with  $\delta_{\lambda} \subseteq \delta_{\lambda'}$  and  $k_{\lambda'}$  not marked superfluous 11  $\delta_{\lambda'} \Leftarrow \delta_{\lambda'} \cup \{k_{\lambda}\}$ 12 Mark  $k_{\lambda}$  superfluous 13 } 14 for  $\lambda = 1$  to l do { 15 **if** not  $k_{\lambda}$  superfluous 16  $\tilde{s}_i \Leftarrow \delta_\lambda \cup \{k_1, \ldots, k_\lambda\}$ 17  $j \Leftarrow j + 1$ 18 } 19 20 Ĵ

A good merging heuristic tries to minimize the number of mergers but simultaneously to use all dependencies in  $G_{ADF}$ . Thus the heuristic generates graphs with  $b_i \neq \emptyset$  which violate the RIP only in a few regions.

Algorithm 3 describes our heuristic. The idea of the sub-function merger algorithm is that each new variable is included in a set together with the previous variables on which it depends. However, if another variable depends on a superset of variables, the two sets are merged. The algorithm calculates  $\tilde{c}_j$ ,  $\tilde{b}_j$  and  $\tilde{d}_j$  analogous to (5).

This sub-function merger algorithm might still compute too large cliques. Therefore a cut parameter k is needed which bounds the clique size. If the size of a clique becomes larger than k our implementation will randomly leave out arcs from  $G_{ADF}$ .

Our presentation of the sub-function merger algorithm has been very short. The interested reader is referred to Bertelé and Brioschi (1972) for an in depth discussion of different fusion and folding heuristics. In the area of Bayesian networks, the problem

<sup>&</sup>lt;sup>5</sup> (Bertelé and Brioschi, 1972) have called it fusion.

	n = 30, N =	= 50		n = 60, N = 70				
Gen	Max prob	Opt prob	Gen		Opt prob			
1	$5.6 \cdot 10^{-6}$	$1.4 \cdot 10^{-7}$	1		$1.6 \cdot 10^{-16}$			
2	$4.0 \cdot 10^{-4}$	$3.5 \cdot 10^{-6}$	2	$3.5 \cdot 10^{-10}$	$9.5 \cdot 10^{-13}$			
3	$2.8\cdot 10^{-3}$	$7.7 \cdot 10^{-5}$	3	$3.0 \cdot 10^{-8}$	$2.7 \cdot 10^{-10}$			
4	$5.9\cdot10^{-2}$	$4.7 \cdot 10^{-4}$	4	$2.0 \cdot 10^{-6}$	$1.2 \cdot 10^{-7}$			
5	$9.7\cdot10^{-2}$	$1.9 \cdot 10^{-2}$	5	$9.3 \cdot 10^{-5}$	$4.1 \cdot 10^{-5}$			
6	$1.8 \cdot 10^{-1}$	$1.2 \cdot 10^{-1}$	6	$1.2 \cdot 10^{-3}$	$9.5 \cdot 10^{-4}$			
7	$4.0 \cdot 10^{-1}$	$4.0 \cdot 10^{-1}$	7	$1.1 \cdot 10^{-2}$	$1.1 \cdot 10^{-2}$			

Table 1: Runs of *FDA* with truncation selection ( $\tau = 0.3$ ) on a separable function (deceptive-3). For each generation, the probabilities of the most probable configuration and of the optimum are shown.

has been investigated by Almond (1995).

If the conditions of the factorization theorem are fulfilled, the convergence proof of *BEDA* is valid for *FDA*, too. Since *FDA* uses finite samples of points to estimate the conditional probabilities, convergence to the optimum will depend on the size of the sample. For small sample sizes the convergence rate is higher if a number of steps with low selection is used instead of just one step using strong selection. Thus this method is numerically more efficient than to use a very large sample size and strong selection.

Table 1 gives some numerical results. For n = 30 the probability to generate the maximum increases from about  $10^{-7}$  to  $10^{-1}$ , and from  $10^{-16}$  to  $10^{-2}$  for n = 60. Note that in the first generations the maximum of the estimated probability is not achieved by the maximum of the function.

*FDA* has experimentally proven to be very successful on a number of functions where standard genetic algorithms fail to find the global optimum. In (Mühlenbein and Mahnig, 1999) the scaling behavior for various test functions has been studied. For recent surveys the reader is referred to (Mühlenbein and Mahnig, 2002a, 2003).

## **3** The maximum entropy principle

*FDA* uses marginals for a factorization  $q(\mathbf{x})$  which estimates the unknown distribution. This problem can be formulated more general.

#### Problem

Given a set of consistent marginal distributions  $p(\mathbf{x}_{s_i})$  from an unknown distribution compute a distribution which satisfies the marginals.

If only a small number of marginals is given the problem is under-specified. Consequently, for incomplete specifications the missing information must be added by some automatic completion procedure. This is achieved by the *maximum entropy principle*. Let us recall

Definition 12. The entropy (Cover and Thomas, 1989) of a distribution is defined by

$$H(p) = -\sum_{x} p(\mathbf{x}) \ln(p(\mathbf{x}))$$
(22)

**Maximum entropy principle (MaxEnt):** Find the maximum entropy distribution  $q(\mathbf{x})$  which satisfies the given marginals.

The maximum entropy principle formulates the *principle of indifference*. If no constraints are specified, the uniform random distribution is assumed. MaxEnt has a long history in physics and probabilistic logic. The interested reader is referred to (Jaynes, 1957, 1978). MaxEnt is especially attractive because there exists a constructive way to obtain the solution. The following important theorem holds:

**Theorem 13.** If the given marginals are consistent then there exists a unique distribution  $q(\mathbf{x})$  of maximum entropy which satisfies the marginals. The distribution can be obtained by Iterative Proportional Fitting (IPF).

IPF iteratively computes a distribution  $q_{\tau}(\mathbf{x})$  from the given marginals  $p_k(\mathbf{x}_k)$ ,  $k = 1, \ldots, K$ , where  $\mathbf{x}_k$  is a sub-vector of  $\mathbf{x}$  and  $\tau = 0, 1, 2, \ldots$  is the iteration index. Let n be the dimension of  $\mathbf{x}$  and  $d_k$  be the dimension of  $\mathbf{x}_k$ .  $q_{\tau=0}$  is the uniform distribution. The update formula is

$$\forall \mathbf{x} \quad q_{\tau+1}(\mathbf{x}) = q_{\tau}(\mathbf{x}) \frac{p_k(\mathbf{x}_k)}{\sum\limits_{\mathbf{y} \in \{0,1\}^{n-d_k}} q_{\tau}(\mathbf{x}_k, \mathbf{y})}$$
(23)

with  $k = ((\tau - 1) \mod K) + 1$ .

The proof that IPF converges against the maximum entropy solution was first tried by Kullback (1968), but was faulty. The correct proof in a most general measure-theoretic framework was given in (Csiszár, 1975). Since the distribution q, which has to be stored and updated in every time step, has exponential size, this implementation takes exponential time and space.

We next connect the solution given by the factorization theorem with the MaxEnt solution.

**Theorem 14.** Let consistent marginal distributions  $p(\mathbf{x}_{s_i})$  be given. Let the assumptions of the factorization theorem be fulfilled. Then the factorization

$$p_{\beta}(\mathbf{x}) = \prod_{i=1}^{m} p_{\beta}(\mathbf{x}_{b_i} | \mathbf{x}_{c_i})$$

is the MaxEnt solution.

*Proof.* Use IPF to compute the MaxEnt solution. Apply the junction tree algorithm. Using these cliques one can show that IPF converges in just one sweep.  $\Box$ 

**Remark:** This theorem has important implications. It shows that all factorizations computed by the the junction tree algorithm generate the same distribution, namely the unique MaxEnt solution.

For many problems IPF cannot be performed in polynomial time. But IPF can be easily used locally to estimate higher order marginals from lower marginals computed from data. Thus it might be advantageous to compute only marginals of low order from the data, but use a factorization containing higher order marginals. The higher order marginals can be computed by *IPF*. A confirmation of this result can be found in (Ochoa et al., 2003). In this paper the structure is learned from the data. The structure is restricted to singly connected poly-trees. The poly-tree is constructed by bi-variates only. For sampling the junction tree is used. The higher order marginals are computed from the bi-variates using IPF. This algorithm performs by far better than computing the higher order marginals directly from the data.

Optimization problems which have a polynomially bounded factorization fulfilling *RIP* can be solved in polynomial time. But this is a *sufficient condition*, not a *necessary*  *condition*. Many problems do not admit a PBF fulfilling *RIP*, but an approximate factorization might still lead to the optimum. Our results obtained so far can be formulated in a conjecture.

**Conjecture:** In the class of ADF's with non-polynomially bounded factorization there exist instances which can only be solved in exponential time. But the number of instances which can be solved polynomially seems to be very large.

**Example:** 

$$f(\mathbf{x}) = \prod_{i=1}^{n} x_i$$
$$f(\mathbf{x}) = \prod_{i=1}^{n} x_i + \sum_{i=1}^{n} x_i$$

Both problems do not admit a polynomially bounded exact factorization. But whereas the first problem can only be solved in exponential time, the second problem can be solved by a simple univariate approximation:

$$p(\mathbf{x}) = \prod_{i=1}^{n} p(x_i)$$

Let us summarize the results: For all sets of consistent marginal distributions, there exists a unique MaxEnt distribution. If the graph corresponding to the marginal distributions admits a junction tree with polynomially bounded clique sizes, then the MaxEnt solution can be computed in polynomial time. If the graph does not admit a bounded junction tree, then the known algorithm (IPF) to compute the MaxEnt solution is exponential in the number of vertices. Nevertheless, approximate factorizations might still lead to the optimum.

The last observation points into the direction to proceed – one should aim to compute a good approximate factorization. Here two approaches are possible.

- If the *ADF* is explicitly given, compute a factorization which leads to a good approximation of the true distribution. This method is used by *FDA*.
- If only data points are given, compute a good factorization from the selected set of points ("learning a model").

The first approach has already been discussed. Before we describe the second approach, we discuss the methods pursued in statistical physics. They belong to the first class, but the marginals are not computed from sampling.

# 4 Computing approximate factorizations in statistical physics

We discuss the method using an important example, the 2-D Ising model. Each cell of the grid, called spin s, is in one of two states, +1 or -1. The cell is influenced by the four neighbors only. It is important to note (at least for a computer scientist) that Ising did *not specify any dynamics*. Instead Ising assumed that the system behaves according to a stationary distribution which is given by the *Boltzmann* distribution

$$p_{\beta}(\mathbf{s}) = \frac{1}{\tilde{Z}_{\beta}} e^{\beta \sum_{(ij)} \bar{J}_{ij} s_i s_j + \sum_i \bar{J}_i s_i}$$

 $\tilde{J}_{ij}$  are the coupling constants. (i, j) denotes a neighboring pair. For all non-neighbors, we can set  $\tilde{J}_{ij} = 0$ . In particular, without loss of generality, we set  $\tilde{J}_{ii} = 0$ , because this adds only a constant, which cancels out with  $\tilde{Z}_{\beta}$ .

Thus we again encounter the problem to *approximate a Boltzmann distribution*. Using  $s_i = 2x_i - 1$  we can change the variables to  $x_i \in \{0, 1\}$ . We obtain

$$p_{\beta}(\mathbf{x}) = \frac{1}{Z_{\beta}} e^{\beta \sum_{(ij)} J_{ij} x_i x_j + \sum_i J_i x_i}$$
(24)

$$J_{ij} = 4\tilde{J}_{ij} \tag{25}$$

$$J_i = 2\tilde{J}_i - 2\sum_j \left(\tilde{J}_{ij} + \tilde{J}_{ji}\right)$$
(26)

and a different partition function  $Z_{\beta}$ .

In statistical physics the maximum entropy principle is extended because additional information is available. Instead of minimizing the distance to the uniform random distribution (this is another formulation of MaxEnt), the distance to the Boltzmann distribution is minimized. As distance measure the *Kullback-Leibler divergence* is used.

Definition 15. The Kullback-Leibler divergence (KLD) between two distributions is defined by

$$KLD(q||p) = \sum_{\mathbf{x}} q(\mathbf{x}) \ln \frac{q(\mathbf{x})}{p(\mathbf{x})}$$
(27)

Note that KLD is not symmetric! Thus we have two choices.

**Minimum relative entropy principle (MinRel)** Given a set of consistent marginal distributions, find a distribution q with these marginals which minimizes KLD(q||p) to the target distribution  $p(\mathbf{x})$ .

**Remark:** If  $p(\mathbf{x})$  is the uniform random distribution, then MinRel is identical to MaxEnt. This justifies the above definition. But from a mathematical point of view, it is also possible to minimize the complementary divergence KLD(p||q) instead. Cover and Thomas (1989) (p. 18) call KLD(p||q) the expected logarithm of the *likelihood ratio*. It is a measure of the inefficiency of assuming q when the true distribution is p. It is connected to the description length. If we knew p we could construct a code with average description length H(p). If, instead, we used the code for distribution q, we would need H(p) + KLD(p||q) bits on the average to describe the random variable. Thus the following principle is also justified:

**Minimum expected log-likelihood ratio principle (MinLike)** Given a set of consistent marginal distributions, find a distribution q with these marginals which minimizes KLD(p||q) to the target distribution  $p(\mathbf{x})$ .

If p is the uniform random distribution, then MinLike minimizes  $\sum_{x} \ln q(\mathbf{x})$ . This is not the entropy of  $q(\mathbf{x})$ . The MinLike principle will be later used for learning Bayesian networks.

In physics MinRel is used. Let us now introduce some physical terms. This is not necessary, but it will help the reader to understand the statistical physics papers better. **Definition 16**. *The average energy U and Gibbs free energy G are defined by* 

$$U(q) = -\sum_{n} q(\mathbf{x}) f(\mathbf{x})$$
(28)

$$G(q) = U(q) - H(q)$$
(29)

We can set  $\beta = 1$ . Then we obtain

$$KLD(q||p) = U(q) - H(q) + \ln Z.$$
 (30)

KLD = 0 will be achieved for  $q(\mathbf{x}) = p(\mathbf{x})$ . This gives the minimal value of  $G(q) = -\ln Z$ . This leads to the idea to minimize G(q) for a class of distributions q. This means we have to compute U(q) and H(q). The computation of U(q) is easy because the following theorem holds.

**Theorem 17.** (Mühlenbein and Mahnig, 2002a) : Let  $I = \{1, ..., n\}$ . Let  $s \subseteq I$  be a multiindex. Then every binary function can be written as

$$f(\mathbf{x}) = \sum_{s \subseteq I} a_s \mathbf{x}_s \tag{31}$$

Furthermore, the average of f with respect to a distribution  $q(\mathbf{x})$  is given by

$$E_q(f) = \sum_{\mathbf{x}} q(\mathbf{x}) f(\mathbf{x}) = \sum_{s \subseteq I} a_s q_s$$
(32)

where  $q_s$  is the marginal distribution  $q_s := q(X_i = 1 | i \in s)$ 

The above theorem can easily be applied to the Ising problem, and more general to ADF's defined on grids. Thus U(q) can efficiently be computed, but the computation of H(q) is more difficult. We will discuss two approximations , the first one uses univariate marginals and the second one bi-variate marginals.

## 4.1 The mean-field approximation

Let us assume that a product distribution is given.

$$q(\mathbf{x}) = \prod_{i=1}^{n} q(x_i)$$
(33)

Then we can compute its entropy

$$H(q) = -\sum_{x} \prod_{i=1}^{n} q(x_i) \sum_{j=1}^{n} \ln q(x_j)$$
  
=  $-\sum_{x_1} q(x_1) \ln q(x_1) - \sum_{x_2,\dots,x_n} \prod_{i=2} q(x_i) \sum_{j=2} \ln q(x_j)$   
=  $-\sum_{i=1}^{n} \sum_{x_i} q(x_i) \ln q(x_i)$ 

We can now try to find a local minimum by setting the derivative of KLD equal to zero, using the uni-variates as variables. We abbreviate  $q_i = q(x_i = 1)$ 

**Theorem 18.** The mean-field approximation minimizes the Kullback-Leibler divergence to the Boltzmann distribution. The local minima of the divergence are given by the nonlinear equation

$$q_i^* = \frac{1}{1 + e^{\frac{\partial U}{\partial q_i}}} \tag{34}$$

*Proof.* From (30) we obtain

$$\frac{\partial KLD}{\partial q_i} = \ln \frac{q_i}{1 - q_i} + \frac{\partial U}{\partial q_i} = 0$$
(35)

The solution gives (34).

In (35) the derivative of the average energy U(q) enters. From theorem 17 we obtain for  $\beta=1$ 

$$U(q) = -\sum_{(ij)} J_{ij} q_i q_j - \sum_i J_i q_i$$
(36)

Taking the derivative gives

$$q_i^* = \frac{1}{1 + e^{-\sum_{j \neq i} (J_{ij} + J_{ji})q_j - J_i}}$$
(37)

This equation can be solved by iteration.

**Remark:** In the mean-field approximation the univariate marginals are considered to be variables. The marginals are computed from (37). G(q) is an upper bound of  $-\ln Z$ . The uni-variate approximation *UMDA* (Mühlenbein and Mahnig, 2001) computes the marginals from the samples. It has to be investigated if the additional computational effort needed for the mean-field approach pays off.

We next extend the mean-field approach to higher order marginals.

# 4.2 Bethe-Kikuchi approximation and the FDA factorization

An obvious extension of the mean-field approach is the use of higher order marginals. This has been done by Bethe (1935) using bi-variate marginals and Kikuchi (1951) for higher order marginals. The interested reader is referred to Yedidia et al. (2001) for the original statistical physics approach. A state-of the art report has recently been written by the same authors (Yedidia et al., 2004).

The approximations of Bethe-Kikuchi may contain loops if  $G_{ADF}$  contains loops. Therefore these approximations are not sufficient to sample from a distribution as is needed for our optimization problem. Therefore we decided to use our *FDA* factorization as an approximation instead. This factorization does not contain cycles. For the *FDA* factorization  $q(\mathbf{x}) = \prod_{i=1}^{m} \tilde{q}(\mathbf{x}_{b_i} | \mathbf{x}_{c_i})$  we have

$$H(q) = -\sum_{i=1}^{n} \sum_{\mathbf{x}_{b_i}, \mathbf{x}_{c_i}} q(\mathbf{x}_{b_i}, \mathbf{x}_{c_i}) \ln \tilde{q}(\mathbf{x}_{b_i} | \mathbf{x}_{c_i})$$
(38)

The proof is based on marginalization and left to the reader. But from equation (9) we know that  $q(\mathbf{x}_{b_i}, \mathbf{x}_{c_i}) \neq \tilde{q}(\mathbf{x}_{b_i}, \mathbf{x}_{c_i})$  if the RIP is violated. Therefore we approximate

$$H(q) \approx -\sum_{i=1}^{n} \sum_{\mathbf{x}_{b_i}, \mathbf{x}_{c_i}} \tilde{q}(\mathbf{x}_{c_i}) \tilde{q}(\mathbf{x}_{b_i} | \mathbf{x}_{c_i}) \ln \tilde{q}(\mathbf{x}_{b_i} | \mathbf{x}_{c_i})$$
(39)

U(q) can be computed using (32). We approximate the true marginals as before by setting  $q(\mathbf{x}_{b_i}, \mathbf{x}_{c_i}) \approx \tilde{q}(\mathbf{x}_{b_i}, \mathbf{x}_{c_i})$ . Having approximations of U(q) and H(q) we can minimize G(q) as before. For the minimization the marginals  $\tilde{q}(x_i, x_j)$  are considered to be variables. The minimization is very difficult, because the variables are not independent, but the marginals have to be consistent. The minimization becomes easier if the number of variables can be reduced substantially, e.g. the unknown Boltzmann distribution is homogenous.

We can summarize the major points of the Bethe-Kikuchi approximation:

The energy function is explicitly given. The marginals used for the factorization are not computed from a sample, but are determined by computing a local minimum of the Kullback-Leibler divergence to the Boltzmann distribution. For the 2-D grid G(q) can be computed only approximatively, therefore the goodness of the approximation cannot be assessed theoretically.

To our knowledge there exists no computer implementation of the full Bethe-Kikuchi method. Yedidia et al. (2004) have recently shown that loopy belief propagation converges to local minima of the approximation of G(q), if it converges at all. We will implement our proposal in the near future.

We next turn to an approach which uses only samples of good fitness values. This approach has first been investigated in artificial intelligence. It is called learning of the model (Jordan, 1999).

# 5 Learning a Bayesian network from data

This section will be very brief, compared to the difficulty of the subject. An excellent indepth discussion can be found in (Larrañaga and Lozano, 2002). We will just motivate some of the major design decisions. Let  $p(\mathbf{x})$  be the true distribution. The learning algorithm uses acyclic Bayesian networks (acBN) as models.

$$q(\mathbf{x}) = \prod_{i=1}^{n} \hat{p}(x_i | \pi_i)$$
(40)

 $\Pi_i$  are called the parents of  $X_i$ ,  $\hat{p}(x_i|\pi_i)$  is a numerical approximation of the true conditional marginal  $p(x_i|\pi_i)$ . If the running intersection property is fulfilled, the Bayesian network is *singly connected*. If the number of the parents  $|\Pi_i|$  is bounded by a constant independent from *n*, we say the Bayesian network is polynomially bounded (PBB).

Both the MaxEnt and the MinRel principle assume that a *fixed* set of marginal distributions is given. But if the data is provided by a sample, we can choose which marginal distributions should be used in order to obtain a Bayesian network which reproduces the data accurately. This is called *model selection*.

Therefore we have to deal with the problem *how to choose the appropriate model*. This problem can be solved in the following way. Let Q be the set of all possible distributions q defined by the Bayesian networks considered. (Because of the efficiency the number of parents is bound usually by a small number.

We first compute the average of  $\ln q$  over the true distribution p

$$E(\ln q) = \sum_{\mathbf{x}} p(\mathbf{x}) \ln q(\mathbf{x})$$
(41)

We have

$$E(\ln q) = -H(p) - KLD(p||q)$$
(42)

Therefore the minimization of KLD(p||q) in Q is equivalent to maximization of  $E(\ln q)$ . The next proposition shows that  $E(\ln q)$  can be computed exactly, in contrast to G(q) which is used for the MinRel method.

**Proposition 19.** For the distribution  $q(x) = \prod_{i=1}^{n} \hat{p}(x_i | \pi_i)$  we have

$$E(\ln q) = \sum_{i=1}^{n} \sum_{x_i, \pi_i} p(x_i, \pi_i) \ln \hat{p}(x_i | \pi_i)$$
(43)

Proof.

$$\sum_{\mathbf{x}} p(\mathbf{x}) \ln q(\mathbf{x}) = \sum_{\mathbf{x}} p(\mathbf{x}) \sum_{i=1}^{n} \ln \hat{p}(x_i | \pi_i)$$
$$= \sum_{i=1}^{n} \sum_{x_i, \pi_i} p(x_i, \pi_i) \ln \hat{p}(x_i | \pi_i)$$

Equation (43) can be approximated using a finite sample by setting  $p(x_i, \pi_i) \approx \hat{p}(x_i, \pi_i)$ . We introduce the following notation. Let N denote the size of the sample. Let  $N_{ijk}$  denote the number of instances with  $x_i = k$  and  $\pi_i = j$ , where the states of  $\Pi_i$  are numbered  $1 \le j \le 2^{|\Pi_i|}$ . Let  $N_{ij} = \sum_k N_{ijk}$ . We can now approximate

$$\hat{p}(x_i = k | \pi_i = j) = \frac{N_{ijk}}{N_{ij}}$$
(44)

$$p(x_i = k, \pi_i = j) = \lim_{N \to \infty} \frac{N_{ijk}}{N}$$
(45)

$$E(\ln q) = \lim_{N \to \infty} \left( E_N(\ln q) = \sum_{i=1}^n \sum_{j=1}^{2^{|\Pi_i|}} \sum_{k=1}^2 \frac{N_{ijk}}{N} \ln \frac{N_{ijk}}{N_{ij}} \right)$$
(46)

Thus we have arrived at the following principle

**Finite sample MaxLike principle (FinMaxLike)** *Maximize in the class of Bayesian networks Q* 

$$max_{q\in Q}E_N(\ln q) = \sum_{i=1}^n \sum_{j=1}^{2^{|\Pi_i|}} \sum_{k=1}^2 \frac{N_{ijk}}{N} \ln \frac{N_{ijk}}{N_{ij}}$$
(47)

FinMaxLike can also be derived from the maximum log-likelihood principle. **Proposition 20.** Let l(G|D) be the log-likelihood of the data given graph *G*. Then

$$l(G|D) = N \cdot E(\ln q) \tag{48}$$

*Proof.* Let  $D = {\mathbf{x}_1, \dots, \mathbf{x}_N}$ . Then the likelihood of D is given by

$$L(G|D) = \prod_{l=1}^{N} q(\mathbf{x}_{l}|G) = \prod_{l=1}^{N} \prod_{i=1}^{n} q(x_{li}|\pi_{li})$$
(49)

Therefore we obtain

$$l(G|D) = \ln L(G|D) = \sum_{l=1}^{N} \sum_{i=1}^{n} \ln q(x_{li}|\pi_{li})$$
$$= \sum_{i=1}^{n} \sum_{l=1}^{N} \ln q(x_{li}|\pi_{li})$$
$$\approx \sum_{i=1}^{n} \sum_{j=1}^{2^{|\Pi_i|}} \sum_{k=1}^{2} N_{ijk} \ln \frac{N_{ijk}}{N_{ij}}$$

For given N maximizing  $E(\ln q)$  gives the same result as maximizing l(G|D). But this principle is not yet sufficient for practical computations. FinMaxLike does not prefer exact models of small complexity (small number of variables) to exact models of large complexity. This can be seen as follows. If we approximate  $\hat{p}(x_i, \pi_i) \approx p(x_i, \pi_i)$ , then we obtain

$$E(\ln q) = \sum_{i=1}^{n} \left( \sum_{x_i, \pi_i} p(x_i, \pi_i) \ln \frac{p(x_i, \pi_i)}{p(x_i)p(\pi_i)} + \sum_{x_i} p(x_i) \ln p(x_i) \right)$$
(50)

The first term is called mutual information (Cover and Thomas, 1989).

$$I(X;Y) = \sum_{x,y} p(x,y) \ln \frac{p(x,y)}{p(x)p(y)}$$
(51)

If *X* and *Y* are independent, we have I(X; Y) = 0. Using the mutual information we obtain

$$E(\ln q) = \sum_{i=1}^{n} (I(X_i; \Pi_i) - H(q(x_i)))$$
(52)

 $E(\ln q)$  remains unchanged if a vertex between two independent variables is inserted. In order to solve this over-fitting problem, we need a criterion which *maximizes the log-likelihood* and *minimizes the complexity*. Such a criterion can be derived using Bayesian principles or a concept called the *minimum description length* MDL. The interested reader is referred to (Jordan, 1999). One of the most popular criterion has been derived by Schwarz (1978). It has been called the *Bayesian Information Criterion* BIC.

**Definition 21.** Let V be the degrees of freedom of the marginal distributions defined by the graph G. Then the weighted BIC measure is defined by

$$BIC_{\alpha} = N \cdot E(\ln q) - \alpha \ln N \cdot V$$

It has been shown that  $-BIC = -N \cdot E(\ln q) + \alpha \ln(N) \cdot V$  is asymptotically equivalent to the minimum description length. Schwarz (1978) computed  $\alpha = 0.5$  as the best weighting factor for  $N \to \infty$ .

The *BIC* criterion can be used to construct a Bayesian network. Basically there are two approaches. The first one starts with an *empty* network and adds arcs between *dependent* variables, the second one starts with the *fully connected* network and *deletes* arcs between *independent* variables. In most implementations the first approach is used together with a simple greedy heuristic for adding a single arc at each step. The reader is referred to (Larrañaga and Lozano, 2002; Mühlenbein and Mahnig, 1999). The quality of both approaches depends on a *good estimate of the mutual information*, the heuristic to *construct the network*, and the *weighting factor*.

These topics will be shortly investigated next. A detailed report will be available soon.

## 6 A comparison of the approaches

In this section we will discuss the three different methods presented in this paper -MaxEnt, MinRel, and MinLike. We restrict the discussion to problems where an additive decomposed function (ADF) is given.

FDA uses a factorization defined by equation (13). If the factorization fulfills the running intersection property it follows from the factorization theorem that the factorization gives the true distribution. But in general, the factorization will not fulfill the

*RIP.*  $G_{\text{FDA}}$  can easily be transformed into an acyclic Bayesian network. This class is used by *LFDA*. Therefore *FDA* and *LFDA* use the same class of graphical models, only the computation of the graph is different.

The approximations originally proposed by Bethe-Kikuchi contain  $G_{ADF}$ . But they compute only the marginals, not the whole distribution. Let us discuss the methods using a simple example.

Example:

$$f(\mathbf{x}) = J_{12}x_1x_2 + J_{23}x_2x_3 + J_{34}x_3x_4 + J_{14}x_1x_4 + \sum J_ix_i$$
$$p(\mathbf{x}) = \frac{e^{f(\mathbf{x})}}{Z}$$

 $G_{ADF}$  is a loop. The following approximations can be used

- 1.  $q(x) = \tilde{q}(x_1)\tilde{q}(x_2|x_1)\tilde{q}(x_3|x_2)\tilde{q}(x_4|x_3)$ - It defines an acBN with *RIP*, but it does not contain *G*<sub>ADF</sub> (the arc between  $x_4$  and  $x_1$  is missing).
- 2. q(x) = q̃(x<sub>1</sub>)q̃(x<sub>2</sub>|x<sub>1</sub>)q̃(x<sub>3</sub>|x<sub>2</sub>)q̃(x<sub>4</sub>|x<sub>1</sub>, x<sub>3</sub>)
   It defines an acBN *without RIP*. It contains G<sub>ADF</sub>. This factorization is computed by *FDA* using merging of sub-functions. The last factor is a tri-variate marginal.
- 3.  $q(x) = \tilde{q}(x_1)\tilde{q}(x_2|x_1)\tilde{q}(x_3|x_1, x_2)\tilde{q}(x_4|x_3, x_1)$ - It defines an acBN with *RIP* which contains  $G_{ADF}$ . It uses two tri-variate marginals. This graph will be obtained by the junction tree algorithm. The approximation is exact.
- 4. Original Bethe approach: Compute marginals (q̃(xi, xi+1), i = 1, ... 4) by minimizing G(q) = U(q) H(q)
  The marginals define a graph with a loop. It needs an iterative method to sample from the marginals.
- 5. Given all bi-variate marginals of the unknown distribution p(x1, x2), ... p(x1, x4), compute the unique MaxEnt distribution qME(x)
  We have shown that qME(x) = p(x), but the computation of qME(x) is exponential.

From a theoretical standpoint the junction tree factorization is the best. But for 2-D grids this method leads to large clique sizes (*O* (size of the grid)). We have proposed to use the *FDA* factorization. For 2-D grids the sub-function merge algorithm computes tri-variate marginals in the interior.

**Remark:** In this paper we have discussed a specific version of the MaxEnt principle, namely a consistent set of marginals is given as constraints. The original MaxEnt has been introduced with other classes of constraints, mainly with moments of given functions. If the averages of the sub-functions are taken as the constraints then the MaxEnt solution is an exponential distribution, in our case the Boltzmann distribution of the ADF (Jaynes, 1978; Cover and Thomas, 1989)!

The comparison of *LFDA* and *FDA* needs a more detailed investigation. This is done next.

# 7 How to test EDA algorithms

In our opinion most researchers so far have concentrated on the benchmark method to show the power of EDA algorithms. A popular benchmark or a difficult function

Table 2: Number of edges added by *LFDA* for a uniform random data set (average over ten runs).

$\alpha$	n	N	# edges	n	N	# edges	n	N	# edges
1.00	25	200	0.3	50	400	0.4	100	800	0.8
0.75	25	200	1.5	50	400	3.4	100	800	6.5
0.50	25	200	7.1	50	400	17.2	100	800	45.8
0.25	25	200	38.4	50	400	89.4	100	800	197.6
0.10	25	200	113.1	50	400	254.7	100	800	536.5
0.50	25	10000	0.5	50	10000	4.3	100	10000	10.9

is chosen and the success of the optimization algorithm is shown. The success rate is usually the percentage of runs computing the optimum. The internal behavior of the algorithm (e.g. which Bayesian network it has constructed etc.) is not investigated. Therefore a generalization of the results to other problems is difficult.

We propose to test EDA algorithms in carefully selected steps instead – starting from theoretically understood problems to more complex ones. Learning the structure of the Bayesian network from data is a difficult task. Many factors contribute to the success or failure of the learning method. The Bayesian network community has investigated this problem intensively. We just mention (Xiang et al., 1997) as a starting point. The test procedure is obvious. A Bayesian network is used to generate the data, then a Bayesian network is learned from this data. It is measured how close the learned Bayesian network is to the network generating the data. The results show that in general large data sets are needed to learn a network which is close to the given one. We first test *LFDA* on problems where all variables are independent, i.e  $G_{\rm FDA}$  contains no arcs.

## 7.1 The penalty weight $\alpha$

Schwarz (1978) has computed an optimal penalty factor  $\alpha = 0.5$  under very restrictive assumptions. (One of the assumptions is  $N \to \infty$ .) Since we want to use fairly small population sizes, we investigate the influence of  $\alpha$  on the computed network in the neighborhood of  $\alpha = 0.5$ . In a first test we generate uniform random data. In this case the exact network has no edges at all. Table 2 shows empirical results. How is an optimal  $\alpha$  defined? It is obvious that no edges will be generated for a large  $\alpha$ . For very small  $\alpha$  many edges will be generated. Thus we are looking for a value of  $\alpha$  at the transition between these two regimes. Informally speaking, we look for  $\alpha_{tr}$  with  $\#edges \leq 5$  for  $\alpha > \alpha_{tr}$ ,  $4 \leq \#edges \leq 10$  for  $\alpha = \alpha_{tr}$  and #edges > 10 for  $\alpha < \alpha_{tr}$ .

The results of table 2 suggest that a value of  $\alpha_{tr} = 0.75$  fulfills the requirements for reasonably large population sizes. The last row shows the results for a very large population. In this case  $\alpha_{tr} = 0.5$  might be indeed the best value.

We have also analyzed the weighting factor in optimization tasks. In many applications the success of *LFDA* can dramatically be increased by a suitable weighting factor. But in general truncation selection with  $\tau = 0.2$  and  $\alpha = 0.5$  is a good starting point.

## 7.2 The connection between FDA and LFDA

At first it seems that an internal testing of the learned network of *LFDA* is impossible, because the structure of the true network seems to be unknown. But the theory helps.

*FDA* can be seen as the infinite sample size limit of any plausible learning method, e.g *LFDA*. The relation between *FDA* and *LFDA* is difficult to formulate precisely. The following conjecture is a first try.

**Conjecture:** Let the empirical distribution  $\hat{p}(\mathbf{x})$  be generated by selection from an ADF. Then for  $N \to \infty$  the mutual information is the largest between those variables which are contained in a common sub-function. This means that the graph obtained by connecting the variables with highest mutual information contains  $G_{ADF}$ .

Thus for  $N \to \infty$  the learning algorithm *LFDA* has to solve the same problem as *FDA*: Given the graph  $G_{ADF}$  compute an acyclic Bayesian network. *LFDA* has an advantage, because it can use the mutual information to leave out less important arcs. For some practical problems with a sparsely connected  $G_{ADF}$  the graph  $G_{FDA}$  contains  $G_{ADF}$ . Thus it does not leave out arcs. The same is true for 2-D grids and *LFDA*.

**Observation:** If the ADF is defined on a 2-D grid, then  $G_{LFDA}$  for  $N \to \infty$  contains  $G_{ADF}$ .

The above observation depends on the specific learning algorithm. *LFDA*, for instance, uses a greedy heuristic which adds a single arc at each step. This method has limitations in constructing the correct network for some artificial cases, as is shown in (Xiang et al., 1997).

We will discuss the results for three typical benchmark functions. The first function is separable of order 5 and deceptive.

$$F_{\text{Dec5}}(x) = \sum_{i=1}^{m} f_{\text{dec5}}(x_{5i-4}, \dots, x_{5i}) \qquad (l = 1, 2, 3)$$

with

$$f_{\text{dec5}}(x_1, x_2, x_3, x_4, x_5) = \begin{cases} 0.9 \iff \sum x_i = 0\\ 0.8 \iff \sum x_i = 1\\ 0.7 \iff \sum x_i = 2\\ 0.6 \iff \sum x_i = 3\\ 0.0 \iff \sum x_i = 4\\ 1.0 \iff \sum x_i = 5 \end{cases}$$
(53)

The second function is non-separable. It consists of m overlapping blocks of size 3 (n = 2m + 1). We have a different function for the last block.

$$F_{\text{IsoPeak}}(\mathbf{x}) := \sum_{i=1}^{m-1} f(x_{2i-1}, x_{2i}, x_{2i+1}) + g(x_{2m-1}, x_{2m}, x_{2m+1})$$
(54)

$$f(x, y, z) = \begin{cases} m & \iff (x, y, z) = (0, 0, 0) \\ m - 1 & \iff (x, y, z) = (1, 1, 1) \\ 0 & \text{otherwise} \end{cases}$$
(55)

$$g(x, y, z) = \begin{cases} m & \iff (x, y, z) = (1, 1, 1) \\ 0 & \text{otherwise} \end{cases}$$
(56)

The third class of functions ( $F_{3-SAT}$ ) are 3-SAT problems with 20 and 50 variables.

Alg	n	$F_{\text{Dec5}}$	$F_{IsoPeak}$	Alg	F <sub>Dec5</sub>	$F_{\text{IsoPeak}}$
FDA	25	400	250	LFDA	*3500	900
FDA	50	600	700	LFDA	*18000	*5000
FDA	100	800	500			
FDA	200	1200	† <b>3500</b>			

Table 3: The minimal population size to find the optimum with 95%.

Table 3 gives the population size for which the optimum is found with 95% in 20 runs. The selection threshold for truncation selection was set to  $\tau = 0.3$ , except \* ( $\tau = 0.1$ ), and † ( $\tau = 0.7$ ). For missing values, the required population size is too large.

The population size needed for finding the optimum is much larger for *LFDA* than for *FDA*. The results of *LFDA* for the 'easy' separable function  $F_{Dec5}$  is especially disappointing. The 3-SAT problem with 20 variables problem was very easy to solve. Both *FDA* and *LFDA* need a population size of 250. But for n = 50 the 3-SAT problems turn out to be very difficult to solve, both for *FDA* and *LFDA*. The reason is that the graphs  $G_{ADF}$  are densely connected. For these graphs the computing of good factorizations with bounded clique size is very difficult or even impossible.

The factorization of the separable function  $F_{Dec5}$  is obvious. For  $F_{IsoPeak}$  the following factorization is computed by *FDA* 

$$q(\mathbf{x}) = \hat{p}(x_1, x_2, x_3)\hat{p}(x_4, x_5 | x_3) \cdots \hat{p}(x_{n-1}, x_n | x_{n-2})$$
(57)

Table 4: Characterization of the Bayesian network computed by *LFDA* in the first generations. In column **arcs comp.** the number in brackets denotes the total number of arcs, the first number gives the number of arcs contained in  $G_{ADF}$ . **Mutual information** gives the number of arcs in  $G_{ADF}$  which are contained in the set of arcs having largest mutual information (number in brackets). \* denotes that the *LFDA* run did not find the optimum. Maximum number of parents Q = 8.  $F_{3-SAT}$  with n = 50 was run with very strong selection ( $\tau = 0.03$ ).

F	n	arcs	Ν	arcs comp.	mutual information
$F_{\text{Dec5}}$	50	100	*1000	3(28),5(38),4(34)	3(50),7(50),6(50)
			*5000	2(12),2(20),3(23)	3(50),6(50),6(50)
			*10000	0(2),4(14),6(20)	4(50),5(50),12(50)
			*30000	28(68),63(95),79(105)	20(50),37(50),68(70)
$F_{IsoPeak}$	24	36	500	16(23),19(23),22(28)	21(28), 23(28), 24(28)
			5000	34(39),36(39)	36(40), 36(40)
	48	72	*2000	44(47),54(59),13(18)	60(70),62(70), 35(70)
			5000	58(63),66(73),69(74)	69(80),73(80),75(80)
	74	112	5000	77(85),90(97),98(105)	103(115),109(115),112(115)
$F_{3-SAT}$	20	141	100	55(65),30(479	90(120),90(120)
			500	63(81),65(82),57(70)	90(120),94(120),95(120)
			5000	101(108)	115(120)
$\alpha = 0.25$	50	491	*30000	131(138),131(141),114(135)	175(200),158(200),135(200)
$\alpha = 0.10$			30000	183(288),195(254),180(225)	180(200),184(200),158(200)

In table 4 we investigate the structure learned by *LFDA*. Here we see the reason why *LFDA* needs a huge population size to find the optimum of  $F_{\text{Dec5}}$ . If the popula-

tion is small, the network computed by *LFDA* contains a few arcs only. A closer look shows that this result is not a problem of the learning procedure, but of the computation of the mutual information. Only for N = 30000 a reasonable number of the correct dependencies are contained in the set of the variables with largest mutual information. The reason can easily be given. If selection is done, then it seems that the sub-function is almost linear.

The results for  $F_{IsoPeak}$  are very good. The learned network is almost identical to the network  $G_{FDA}$  computed by the sub-function merger algorithm if one observes that

$$\hat{p}(x_{i-1}, x_i | x_{i-2}) = \hat{p}(x_{i-1} | x_{i-2}) \hat{p}(x_i | x_{i-1}, x_{i-2})$$

The performance for  $F_{3-SAT}$  with 20 variables is surprisingly good. A closer look at the network generated by *LFDA* shows that it contains only 1/3 of the arcs of  $G_{ADF}$ . The situation changes dramatically for  $F_{3-SAT}$  with 50 variables. The optimum is found only with very strong selection ( $\tau = 0.03$ ), a large population size (N = 30000), and a small weight factor ( $\alpha = 0.1$ ). A look into table 4 shows the reason for the bad performance. *LFDA* computes a network which leaves out too many arcs of  $G_{ADF}$ , despite that the mutual information gives correct information about the dependencies. Even for  $\alpha = 0.1$ the learned network  $G_{LFDA}$  contains only about 180 edges from the 491 edges in  $G_{ADF}$ . The same problem occurs with *FDA*.

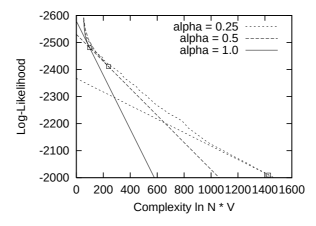


Figure 4: Trajectories of three runs of the *LFDA* learning algorithm in the *BIC* space. Maximum number of parents Q = 8, N = 600,  $\alpha$  varies. The squares mark the endpoints of the runs. The tangents with gradient  $\alpha$  are also shown.

In figure 4 we analyze the dynamic behavior of our greedy algorithm computing the network. The search starts with complexity of almost 0 and a small log-likelihood. The log-likelihood is increased until the increase is equal to the increase in complexity. We summarize the results of this section.

The learning algorithms face two problems, first to identify the dependent variables, and second to compute an acBN which has a bound on the number of parents, but does not leave out important dependencies.

In this paper we have only discussed a learning algorithm which starts with an empty network. In the near future we will investigate learning algorithms which start with the fully connected network and *delete* the arcs between *independent* variables. It seems that the detection of independent variables is more reliable for small population sizes than the test for dependence. Unfortunately the construction of the Bayesian network is much more complicated. Just deleting arcs might lead to a very difficult graphical model, which is no Bayesian network at all.

# 8 Conclusion and Outlook

The efficient estimation and sampling of distributions is a common problem in several scientific disciplines. Unfortunately each discipline uses a different language to formulate its algorithms. We have identified two principles used for the approximation – minimum relative entropy and minimum expected log-likelihood ratio. Both principles are closely related. If p is the distribution to be estimated, then MinRel minimizes the Kullback-Leibler divergence KLD(q||p) whereas MinLike minimizes KLD(p||q).

We have shown that the basic theory is the same for all algorithms. This theory deals with the decomposition of graphical models and the computation of approximate factorizations. If the unknown distribution allows an exact factorization, then both methods lead to KLD = 0, thus they compute the exact distribution.

We have discussed two EDA algorithms in detail. *FDA* computes a factorization from the graph representing the structure. If the corresponding graphical model does not fulfill the assumptions of the factorization theorem the exact distribution is only approximated. Promising factorizations can be obtained by merging of sub-functions.

*LFDA* learns the structure from the data. We have discussed some of the problems of this method. A crucial point for the success seems to be the correct detection of the important dependencies. Statistical physics uses the most complex approach, selected marginal distributions are computed by minimizing the distance to the Boltzmann distribution. The marginals generate a dependence graph with loops. Therefore they are not sufficient to define a distribution, any factorization  $q(\mathbf{x})$  using these marginals has to be normalized ( $\sum_{\mathbf{x}} q(\mathbf{x}) = 1$ ). But this summation is exponential. We have proposed an extension of the original approach which circumvents the normalization by using the marginals of a *FDA* factorization.

An important improvement of *FDA* and *LFDA* could not be discussed, this is the use of a local hill-climber. This topic is discussed for large bi-partitioning problems in Mühlenbein and Mahnig (2002a). We have now implemented a local hill-climber proposed by Lin and Kernighan (1973), which can be used for many combinatorial optimization problems. It increases the performance of *FDA* and *LFDA* on 3-SAT problems substantially. In fact, problems up to size 250 pose no difficulties. The reason is that the L-K local optima have a structure which suits *LFDA*.

The goal of this paper is to inspire researchers to implement some of the methods presented and make a detailed comparison between the different methods. Especially interesting would be an implementation of the full Kikuchi method, which has been implemented only for special *ADF* 's so far. Whereas the theory of EDA algorithms is very convincing, we all have to work hard to design numerically efficient EDA algorithms. Efficiency can only be achieved by having a close look at the different developments in the neighboring disciplines mentioned.

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