# Minimizing the Kolmogorov distance between a random variable and an approximation of it with a fixed support size

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#### **Abstract**

- We present an algorithm that takes a discrete random variable X and a number m and computes a random variable whose support (set of possible outcomes) is of
- size at most m and whose Kolmogorov distance from X is minimal.

#### 4 1 Introduction

- 5 Many different approaches to approximation of probability distributions are studied in the litera-
- 6 ture [12, 15, 16]. The approaches vary in the types random variables considered, how they are rep-
- 7 resented, and in the criteria used for evaluation of the quality of the approximations. This paper is
- 8 on approximating discrete distributions represented as explicit probability mass functions with ones
- 9 that are simpler to store and to manipulate. This is needed, for example, when a discrete distribution
- o is given as a large data-set, obtained, e.g., by sampling, and we want to represent it approximately
- with a small table.
- 12 The main contribution of this paper is an efficient algorithm for computing the best possible ap-
- 13 proximation of a given random variable with a random variable whose complexity is not above a
- prescribed threshold, where the measures of the quality of the approximation and the complexity of
- the random variable are as specified in the following two paragraphs.
- We measure the quality of an approximation by the distance between the original variable and the
- 17 approximate one. Specifically, we use the Kolmogorov distance which is commonly used for com-
- $^{18}$  paring random variables in statistical practice and literature. Given two random variables X and
- 19 X' whose cumulative distribution functions (cdfs) are  $F_X$  and  $F_{X'}$ , respectively, the Kolmogorov
- distance between X and X' is  $d_K(X,X')=\sup_t |F_X(t)-F_{X'}(t)|$  (see, e.g., [9]). We say taht X'
- is a good approximation of X if  $d_K(X, X')$  is small.
- 22 The complexity of a random variable is measured by the size of its support, the number of values
- that it can take,  $|\operatorname{support}(X)| = |\{x \colon Pr(X = x) \neq 0\}|$ . When distributions are maintained as
- explicit tables, as done in many implementations of statistical software, the size of the support of
- a variable is proportional to the amount of memory needed to store it and to the complexity of the

- computations around it. In summary, the exact notion of optimality of the approximation targeted in this paper is:
- Definition 1. A random variable X' is an optimal m-approximation of a random variable X if  $|\operatorname{support}(X')| \leq m$  and there is no random variable X'' such that  $|\operatorname{support}(X'')| \leq m$  and
- 30  $d_K(X, X'') < d_K(X, X')$ .
- The main contribution of the paper is an efficient algorithm that takes X and m as parameters and constructs an optimal m-approximation of X.
- The rest of the paper is organized as follows. In Section 2 we describe how our work relates to other algorithms and problems studied in the literature. In Section 3 we detail the proposed algorithm, analyze its properties, and prove the main theorem. In Section 4 we demonstrate how the proposed approach performs on the problem of estimating the probability of hitting deadlines is plans and compare it to alternatives approximation approaches from the literature. We also demonstrate the performance of our approximation algorithm on some randomly generated random variables. The
- paper is concluded with a discussion in Section 5.

#### 40 **2 Related Work**

- The most relevant work related to this paper is the papers by Cohen at. al. [5, 4]. These papers study approximations of random variables in the context of estimating deadlines. In this context, X' is defined to be a good approximation of X if  $F_{X'}(t) > F_X(t)$  for any t and  $\sup_t F_{X'}(t) - F_X(t)$ 43 is small. This is not a distance because it is not symmetric. The motivation given by Cohen at. al. 44 for using this type of approximation is for cases where overestimation of the probability of missing 45 a deadline is acceptable but underestimation is not. In Section 4, we consider the same examples examined by Cohen at. al. and show how the algorithm proposed in this paper performs relative to 47 the algorithms proposed there when both over- and under- estimations are allowed. As expected, the 48 Kolmogorov distance between the approximation and the original random variable is smaller by a 49 factor of one half, on average, when using the algorithm proposed here.
- Another relevant prior work is the theory of Sparse Approximation (aka Sparse Representation) that deals with sparse solutions for systems of linear equations, as follows.

Given a matrix  $D \in \mathbb{R}^{n \times p}$  and a vector  $x \in \mathbb{R}^n$ , the most studied sparse representation problem is finding the sparsest possible representation  $\alpha \in \mathbb{R}^p$  satisfying  $x = D\alpha$ :

$$\min_{\alpha \in \mathbb{R}^p} \|\alpha\|_0 \text{ subject to } x = D\alpha.$$

where  $\|\alpha\|_0 = |\{i: \alpha_i \neq 0, i = 1, ..., p\}|$  is the  $\ell_0$  pseudo-norm, counting the number of non-zero coordinates of  $\alpha$ . This problem is known to be NP-Hard with a reduction to NP-complete subset selection problems.

In these terms, using also the  $\ell_{\infty}$  norm that represents the maximal coordinate and the  $\ell_1$  norm that represents the sum of the coordinates, our problem can be phrased as:

$$\min_{\alpha \in [0,\infty)^p} \|x - D\alpha\|_{\infty} \text{ subject to } \|\alpha\|_0 = m \text{ and } \|\alpha\|_1 = 1.$$

where D is the all-ones triangular matrix (the entry at row i and column j is one if  $i \leq j$  and zero otherwise), x is related to X such that the ith coordinate of x is  $F_X(x_i)$  where  $\operatorname{support}(X) = \{x_1 < x_2 < \dots < x_n\}$  and  $\alpha$  is related to X' such that the ith coordinate of  $\alpha$  is  $f_{X'}(x_i)$ . The functions  $F_X$  and  $f_{X'}$  represent, respectively, the cumulative distribution function of X and the

mass distribution function of X'. This, of course, means that the coordinates of x are assumed to be positive and monotonically increasing and that the last coordinate of x is assumed to be one. We demonstrate an application for this specific sparse representation problem and show that it can be solve in  $O(n^2m)$  time and  $O(m^2)$  memory.

The presented work is also related to the research on binning in statistical inference. Consider, for example, the problem of credit scoring [21] that deals with separating good applicants from bad applicants where the Kolmogorov–Smirnov statistic KS is a standard measure. The KS comparison is often preceded by a procedure called binning where a large table is translated to a smaller one by collecting nearby values together. There are many methods for binning [11, 17, 2, 19]. In this context, our algorithm can be consider as a new binning strategy that provides optimality guarantees with respect to the Kolmogorov distance that none of the existing binning technique that we are aware of provides.

The present study is also related to the work of Pavlikov and Uryasev [15], where a procedure for 72 producing a random variable X' that optimally approximates a random variable X is presented. 73 74 Their approximation scheme, achieved using linear programming, is designed for a different notion of distance (called CVaR). The new contribution of the present work in this context is that our 75 method is direct, not using linear programming, thus allowing tighter analysis of time and memory 76 complexity. Also, our method is designed for optimizing the Kolmogorov distance that is more 77 prevalent in applications. For comparison, in Section 4 we briefly discuss the performance of linear 78 programming approach similar to the one proposed in [15] for the Kolmogorov distance and compare 79 it to the algorithm proposed in this paper. 80

# 81 3 An Algorithm for Optimal Approximation

In the scope of this section, let X be a given random variable with a finite support of size n, and let  $0 < m \le n$  be a given complexity bound. The section evolves by developing notations and by collecting facts towards an algorithm for finding an optimal m-approximation of X.

The first useful fact is that it is enough to limit our search to approximations X's such that support $(X') \subseteq \operatorname{support}(X)$ :

Lemma 2. For every random variable X'' there is a random variable X' such that  $\operatorname{support}(X') \subseteq \operatorname{support}(X)$  and  $d_K(X,X') \le d_K(X,X'')$ .

Proof. Let  $\{x_1,\ldots,x_n\}= \operatorname{support}(X)$ , and let  $x_0=-\infty,x_{n+1}=\infty$ . Consider the random variable X' whose probability mass function is  $f_{X'}(x_i)=P(x_{i-1}< X''\leq x_i)$  for  $i=1,\ldots,n-1$ ,  $f_{X'}(x_n)=P(x_n-1< X''< x_{n+1})$ , and  $F_{X'}(x)=0$  if  $x\notin\operatorname{support}(X)$ . Since X' "accumulates" the non-zero probabilities of X'' to the support of X, we have that  $f_{X'}$  is a probability mass function and therefore X' is well defined.

First see by construction that  $|F_X(x_i) - F_{X'}(x_i)| = |F_X(x_i) - F_{X''}(x_i)|$  for every  $1 \le i \le n-1$ . For i = n we have  $|F_X(x_n) - F_{X'}(x_n)| = |1-1| = 0$ . Finally see that  $|F_X(x) - F_{X'}(x)| = |F_X(x_i) - F_{X'}(x_i)|$  for every  $0 \le i < n+1$  and  $x_i < x < x_{i+1}$ . Therefore we have that  $|F_X(X_i) - F_{X'}(X_i)| \le |F_X(X_i) - F_{X'}(X_i)| \le |F_X(X_i) - F_{X''}(X_i)| \le$ 

Next, note that every random variable X'' with support of size at most m that is contained in support (X) can be described by first setting the (at most m) elements of the support of X''; then for every such option, determine X'' by setting probability values for the elements in the chosen support of X', and setting 0 for rest of the elements.

- Denote the set of random variables with support  $S \subseteq \operatorname{support}(X)$  by  $\mathbb{X}_S$ . In Step 1 below, we find a random variable in  $\mathbb{X}_S$  that minimizes the Kolmogorov distance from X, and denote this distance by  $\varepsilon(X,S)$ . Next, in Step 2, that we will describe later, we will show how to efficiently find S that minimizes  $\varepsilon(X,S)$  among all the sets that satisfy  $S \subset \operatorname{support}(X)$  and  $|S| \leq m$ . Then the minimized random variable  $\mathbb{X}_S$  from the minimal S, is the m-optimal approximation to X.
- 107 Step 1: Finding an X' in  $X_S$  that minimizes  $d_K(X, X')$
- We first fix a set  $S\subseteq \operatorname{support}(X)$  of size at most m, and among all the random variables in
- 109  $X_S$  find one with a minimal distance from X. Denote the elements of S in increasing order by
- 110  $S = \{x_1 < \cdots < x_m\}$  and let  $x_0 = -\infty$ , and  $x_{m+1} = \infty$ . For every  $1 < i \le m$  let  $\hat{x}_i$  be the
- maximal element of support (X) that is smaller than  $x_i$ .
- Next, as the elements of S are also elements of support (X), we can define the following weight
- 113 function:
- Definition 3. For  $0 \le i \le m$  let

$$w(x_i, x_{i+1}) = \begin{cases} P(x_i < X < x_{i+1}) & \text{if } i = 0 \text{ or } i = m; \\ P(x_i < X < x_{i+1})/2 & \text{otherwise.} \end{cases}$$

- Note that  $x_i = -\infty$  for i = 0 and  $x_i = \infty$  for i = m + 1. Also note that  $P(x_i < X < x_{i+1}) =$
- 116  $F_X(\hat{x}_{i+1}) F_X(x_i)$ , a fact that we will use throughout this section.
- 117 **Definition 4.** Let  $\varepsilon(X,S) = \max_{i=0,...,m} w(x_i,x_{i+1})$ .
- We first show that  $\varepsilon(X,S)$  is a lower bound. That is, every random variable in  $\mathbb{X}_S$  has a distance at
- least  $\varepsilon(X,S)$ . Then, we present a random variable  $X' \in \mathbb{X}_S$  with distance  $\varepsilon(X,S)$ . It then follows
- that such X' is an optimal m-approximation random variable among all random variables in  $\mathbb{X}_S$ .
- The intuition behind choosing these specific weights and  $\varepsilon(X,S)$  being a lower bound is as follows.
- Since for every  $X' \in \mathbb{X}_S$  the probability values of X' for the elements not in S are set to 0, we have
- that  $F_{X'}(\hat{x}_{i+1}) = F_{X'}(x_i)$ . Therefore the distance between X' and X at points  $x_i$  and  $\hat{x}_{i+1}$  that we
- have to take into additional account is increased by  $F_X(\hat{x}_{i+1}) F_X(x_i) = P(x_i < X < x_{i+1})$ .
- Formally we have the following.
- Proposition 5. If  $X' \in \mathbb{X}_S$  then  $d_K(X, X') \geq \varepsilon(X, S)$ .
- 127 Proof. By definition, for every  $0 \le i \le m$ ,  $d_K(X,X') \ge \max\{|F_X(\hat{x}_{i+1})|$
- 128  $F_{X'}(\hat{x}_{i+1})|, |F_X(x_i) F_{X'}(x_i)|$ . Note that  $F_{X'}(\hat{x}_{i+1}) = F_{X'}(x_i)$  since the probability values
- for all the elements not in S are set to 0.
- 130 If i=0, that is  $x_i=-\infty$ , we have that  $F_X(x_i)=F_{X'}(x_i)=F_{X'}(\hat{x}_{i+1})=0$  and therefore
- 131  $d_K(X, X') \ge |F_X(\hat{x}_{i+1})| = |F_X(\hat{x}_{i+1}) F_X(x_i)| = P(x_i < X < x_{i+1}) = w(x_i, x_{i+1}).$
- 132 If i=m, that is  $x_{i+1}=\infty$ , we have that  $F_X(\hat{x}_{i+1})=F_{X'}(\hat{x}_{i+1})=F_{X'}(x_i)=1$ . and therefore
- 133  $d_K(X, X') \ge |1 F_X(\hat{x}_i)| = |F_X(\hat{x}_{i+1}) F_X(x_i)| = P(x_i < X < x_{i+1}) = w(x_i, x_{i+1}).$
- Otherwise for every  $1 \le i < m$ , we use the fact that  $max\{|a|,|b|\} \ge |a-b|/2$  for every  $a,b \in A$
- 135  $\mathbb{R}$ , to have  $d_K(X,X') \geq 1/2|F_X(\hat{x}_{i+1}) F_X(x_i) + F_{X'}(x_i) F_{X'}(\hat{x}_{i+1})|$ . So  $d_K(X,X') \geq 1/2|F_X(\hat{x}_{i+1}) F_X(x_i)|$
- 136  $1/2|F_X(\hat{x}_{i+1}) F_X(x_i)| = P(x_1 < X < x_2)/2 == w(x_i, x_{i+1}).$
- Therefore since  $d_K(X,X') \geq w(x_i,x_{i+1})$  for every  $0 \leq i \leq m$ , by definition of  $\varepsilon(X,S)$  proof
- 138 follows. □

- Next we show a random variable  $X' \in \mathbb{X}_S$  with a distance of  $\varepsilon(X,S)$  from X. Thus X' is an
- optimal m-approximation among the set  $X_S$ . We define X' as follows:
- **Definition 6.** Let  $f_{X'}(x_i) = w(x_{i-1}, x_i) + w(x_i, x_{i+1}) + f_X(x_i)$  for i = 1, ..., m and  $f_{X'}(x) = 0$
- 142 for  $x \notin S$ .
- We first show that X' is a properly defined random variable:
- Lemma 7.  $f_{X'}$  is a probability mass function.
- 145 *Proof.* From definition  $f_{X'}(x_i) \geq 0$  for every i. To see that  $\sum_i f_{X'}(x_i) = 1$ , we have
- 146  $\sum_{i} f_{X'}(x_i) = \sum_{i} (w(x_{i-1}, x_i) + w(x_i, x_{i+1}) + f_X(x_i)) = \sum_{x_i \in S} f_X(x_i)) + w(x_0, x_1) + w(x_0, x_$
- 147  $\sum_{0 < i < m} 2w(x_i, x_{i+1}) + w(x_m, x_{m+1}) = \sum_{x_i \in S} P(X = x_i) + P(x_0 < X < X_1) + P(x_i)$
- 148  $\sum_{0 < i < m} P(x_i < X < X_{i+1}) + P(x_m < X < X_{m+1}) = 1$  since this sum is the entire cpt of
- 149 X.
- Note that X' can be constructed in linear time to the size of the cdf of X. Intuitively the setting of
- 151 X' allows to take an "advantage" of distance from X at the elements of support (X'), to avoid the
- overall increased distance of X from X' at the elements that are not at support(X) and in which
- $f_{X'}$  is set to 0. Formally we have the following.
- Lemma 8. Let  $x \in \text{support}(X)$  and  $0 \le i \le m$  be such that  $x_i \le x \le x_{i+1}$  then  $-w(x_i, x_{i+1}) \le x_i$
- 155  $F_X(x) F_{X'}(x) \le w(x_i, x_{i+1}).$
- 156 *Proof.* We prove by induction on  $0 \le i < m$ .
- First see that  $F_{X'}(j) = 0$  for every  $x_0 < j < x_1$  and therefore  $F_X(j) F_{X'}(j) = F_X(j) 0 \le 0$
- 158  $F_X(\hat{x}_1) = F_X(\hat{x}_1) F_X(x_0) = w(x_0, x_1)$ . For  $j = x_1$  we have  $F_X(x_1) F_{X'}(x_1) = F_X(\hat{x}_1) + F_X(\hat{x}_1) = F_X(\hat{x$
- 159  $f_X(x_1) (w(x_0, x_1) + w(x_1, x_2) + f_X(x_1) = w(x_0, x_1) + f_X(x_1) (w(x_0, x_1) + w(x_1, x_2) + f_X(x_1) +$
- 160  $f_X(x_1) = -w(x_1, x_2).$
- Next assume that  $F_X(\hat{x}_i) F_{X'}(\hat{x}_i) = w(x_{i-1}, x_i)$ . Then  $F_X(x_i) F_{X'}(x_i) = F_X(\hat{x}_i) + f_X(x_i) F_{X'}(x_i)$
- 162  $(w(x_{i-1},x_i)+w(x_i,x_{i+1})+f_X(x_i))=w(x_{i-1},x_i)+f_X(x_i)-(w(x_{i-1},x_i)+w(x_i,x_{i+1})+f_X(x_i))$
- 163  $f_X(x_i) = -w(x_i, x_{i+1}).$
- As before we have that for all  $x_i < j < x_{i+1}$ , we have  $F_X(j) F_{X'}(j) = F_X(j) F_{X'}(\hat{x}_{i+1}) \le$
- 165  $F_X(\hat{x}_{i+1}) F_{X'}(\hat{x}_{i+1})$ . Then  $F_X(\hat{x}_{i+1}) F_{X'}(\hat{x}_{i+1}) = (F_X(x_i) + P(x_i < x < x_{i+1})) F_{X'}(\hat{x}_{i+1})$
- 166  $F_{X'}(x_i) = -w(x_i, x_{i+1}) + 2w(x_i, x_{i+1}) = w(x_i, x_{i+1}).$
- Finally for  $x_m \leq j \leq x_{m+1}$  we have that  $F_{X'}(x_m) = 1$  therefore  $F_X(x_m) F_{X'}(x_m) = (1 1)$
- 169  $x_{m+1}$  we have  $F_X(j) F_{X'}(j) < (1 P(x_m < X < x_{m+1})) 1 < -P(x_m < X < x_{m+1})) = 0$
- $-w(x_m,x_{m+1})$  as required.
- From Lemma 8, by the definition of  $\varepsilon(X,S)$ , we then have:
- 172 Corollary 9.  $d_K(X, X') = \varepsilon(X, S)$ .
- Step 2: Finding a set S that minimizes  $\varepsilon(X,S)$
- Finding the m-optimal random variable with a distance  $\varepsilon(X,S)$  among the set  $\mathbb{X}_S$ , we next proceed
- to find an S that minimizes  $\varepsilon(X,S)$ . To obtain that we use a dynamic programming approach,
- motivated by a method described in [3]. Intuitively we construct a directed graph with a source and
- a target, in which every (acyclic) source-target path of size  $\leq m$  corresponds to a possible support

set of the same size, and the weights along that path correspond to the weight as defined in Definition 3. Thus the problem of finding the S that minimizes  $\varepsilon(X,S)$  is reduced to the problem of finding the source-target path  $\vec{p}$  of size  $\leq m$  in that graph such that the maximal weight of an edge in  $\vec{p}$  is minimal among all other such maximal edges in all other paths of size  $\leq m$ .

In details, we initially constructs a directed weighted graph G = (V, E, s, t, w), where V =182  $v_0, \cdots v_{n+1}$  correspond to the support of X with the additional  $v_0 = s$  as the source vertex and 183  $v_{n+1} = t$  as the target vertex. We set an edge  $(v_i, v_j)$  for every i < j and a weight that is defined 184 as follows. If 1 < i < j < n then  $w(v_i, v_j) = P(x_i < X < x_{i+1})/2$ . For i = 0 or j = n+1 we 185 set  $w(v_i, v_j) = P(x_i < X < x_{i+1})$ . Thus the weight of every edge (i, j) corresponds to Definition 186 3 for a case in which  $x_i, x_i$  where consecutive elements in an optional set S of size  $\leq m$ . Note that 187 there is a 1-1 correspondence between a subset S of support (X) of size m, and an s-t path  $\vec{p}_S$  in G 188 where every  $x_i \in S$  corresponds to the vertex  $v_i \in \vec{p}_S$ . On that sense note that the maximal weight 189 of an edge on  $\vec{p}_S$  is identical to  $\varepsilon(X,S)$ . We denote this maximal weight of an edge by  $w(\vec{p}_S)$ , and 190 denote the set of all acyclic s-to-t paths in G with at most m edges is called  $paths_m(G, s, t)$ . Thus, 191 the problem of finding the set S with the minimal  $\varepsilon(X,S)$  is trivially reduced to the problem of find-192 ing a path  $\vec{p} \in paths_m(G, s, t)$  such that  $w(\vec{p})$  is minimal among all  $\{w(\vec{p}')|\vec{p'} \in paths_m(G, s, t)\}$ . 193 This problem can be solved by the standard Bellman-Ford algorithm in m iterations (e.g. [10, 18]) 194 denoted by BellmanFord(G, m). We thus have the following from the standard Bellman-Ford 195 196

Corollary 10. BellmanFord(G,m) returns a path  $\vec{p} \in paths_m(G,s,t)$  such that  $w(\vec{p})$  is minimal among all  $\{w(\vec{p'})|\vec{p'} \in paths_m(G,s,t)\}$ .

### 199 Step 3: Constructing the overall algorithm

We combine Step 1 and Step 2 in the following algorithm called KolmogorovApprox (Algorithm 1) that follows naturally from the two steps. Given X and  $\operatorname{support}(X)$  we add  $x_0, x_{n+1}$  and construct the graph (line 2) as in Step 2. Then we execute the Bellman-Ford algorithm on G for m iterations to obtained a path  $\vec{p} = (v_0, \dots, v_{m+1})$  (line 2) as described in Corollary 10. Finally we use Definition 6 to construct X' from the weights of  $\vec{p}$  (lines 4-5).

## **Algorithm 1:** KolmogorovApprox(X, m)

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1 support(X) \cup \{x_0, x_{n+1}\}

2 Construct G = (V, E, s, t, w) according to Step 2

3 \vec{p} = (v_0, \dots, v_{m+1}) = Belmman - Ford(G, m)

4 for 0 < i < m+1 do

5 \bigcup_i f_{X'}(x_i) = w(v_{i-1}, v_i) + w(v_i, v_{i+1}) + f_X(x_i)

6 return X'
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#### **Theorem 11.** KolmogorovApprox returns an m-optimal-approximation of X.

Proof. From Corollary 10 and the construction of G we get that the path  $\vec{p}$  obtained in line 4 of KolmogorovApprox describes a set S of support of size at most m for which  $\varepsilon(S,X)$  is minimal. Then from Definition 6 and Corollary 9 we construct X' in lines 4-5 of KolmogorovApprox such that  $d_K(X,X')=\varepsilon(X,S)$ . Therefore X' is an m-approximation among all random variables with support contained in support(X). Finally from Lemma 2 we have that X' is m-approximation among all random variables os support of size at most m, thus X' is an m-optimal-approximation of X.

213 Finally we analyze the complexity of KolmogorovApprox as follows.

Theorem 12. The KolmogorovApprox(X, m) algorithm runs in time  $O(mn^2)$ , using  $O(n^2)$  mem-215 ory where  $n = |\operatorname{support}(X)|$ .

*Proof.* Constructing the graph G as described in Step 2 takes  $O(n^2)$ . The number of edges in G 216 is  $O(|E|) = O(n^2)$ , for every edge the weight is at most the sum of all probabilities between the 217 source node s and the target node t, which can be calculated efficiently by aggregating the weights 218 of already calculated edges. The construction is also the only stage that requires memory allocation, 219 specifically  $O(|E| + |V|) = O(n^2)$ . Next using the Bellman-Ford algorithm on G for m iterations 220 takes  $O(m(|E|+|V|)) \approx O(mn^2)$ . [[DF: cite Corman or some algorithms book]]. Finally deriving 221 the new random variable X' from the computed path  $\vec{p}$  takes O(m) time: For every node  $x_i$  in  $\vec{p}$ 222 (at most m nodes), use the already calculated weights to find the probability mass function  $f_{X'}(x_i)$ . 223 To conclude, the time complexity of KolmogorovApprox(X, m) is  $O(n^2 + mn^2 + m) = O(mn^2)$ 224 and memory complexity is  $O(E + V) = O(n^2)$ . 225

# 4 A case study and experimental results

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The case study examined in our experiments is the problem of task trees with deadlines [5, 4]. 227 Hierarchical planning is a well-established field in AI [6, 7, 8], and is still relevant nowadays [1, 228 20]. A hierarchical plan is a method for representing problems of automated planning in which 229 the dependency among tasks can be given in the form of networks, here we focus on hierarchical 230 plans represented by task trees. The leaves in a task tree are primitive actions (or tasks), and the 231 internal nodes are either sequence or parallel actions. The plans we deal with are of stochastic 232 nature, and the task duration is described as probability distributions in the leaf nodes. We assume 233 that the distributions are independent but not necessarily identically distributed and that the random 234 variables are discrete and have a finite support. 235

A sequence node denotes a series of tasks that should be performed consecutively, whereas a parallel 236 node denotes a set of tasks that begin at the same time. A valid plan is one that is fulfilled before 237 some given deadline, i.e., its makespan is less than or equal to the deadline. The objective in this 238 context is to compute the probability that a given plan is valid, or more formally computing P(X <239 T), where X is a random variable representing the makespan of the plan and T is the deadline. The 240 problem of finding the probability that a task tree satisfies a deadline is known to be NP-hard. In 241 fact, even the problem of summing a set of random variables is NP-hard [13]. This is an example of 242 an explicitly given random variable that we need to estimate deadline meeting probabilities for. 243

The first experiment we focus on is the problem of task trees with deadlines, and consider three types of task trees. The first type includes logistic problems of transporting packages by trucks and 245 airplanes (from IPC2 http://ipc.icaps-conference.org/). Hierarchical plans of those logistic problems 246 were generated by the JSHOP2 planner [14], one parallel node with all descendant task nodes being 247 in sequence. The second type consists of task trees used as execution plans for the ROBIL team 248 entry in the DARPA robotics challenge (DRC simulation phase), and the third type is of linear plans 249 (sequential task trees). The primitive tasks in all the trees are modeled as discrete random variables 250 with support of size M obtained by discretization of uniform distributions over various intervals. 251 The number of tasks in a tree is denoted by N. 252

We implemented the approximation algorithm for solving the deadline problem with four different methods of approximation. The first two are for achieving a one-sided Kolmogorov approximation – the OptTrim [4] and the Trim [5] operators, and the third is a simple sampling scheme. We used those methods as a comparison to the Kolmogorov approximation with the suggested KolmogorovApprox algorithm. The parameter m of  $\operatorname{OptTrim}$  and  $\operatorname{KolmogorovApprox}$  corresponds to the inverse of  $\varepsilon$  given to the  $\operatorname{Trim}$  operator. Note that in order to obtain some error  $\varepsilon$ , one must take into consideration the size of the task tree N, therefore,  $m/N=1/(\varepsilon\cdot N)$ . We ran also an exact computation as a reference to the approximated one in order to calculate the error. The experiments conducted with the following operators and their parameters:  $\operatorname{KolmogorovApprox}$  operator with  $m=10\cdot N$ , the  $\operatorname{OptTrim}$  operator with  $m=10\cdot N$ , the  $\operatorname{Trim}$  as operator with  $\varepsilon=0.1/N$ , and two simple simulations, with a different samples number  $s=10^4$  and  $s=10^6$ .

Task Tree	M	KolmogorovApprox	OptTrim	Trim	Sampling	
		m/N=10	m/N=10	$\varepsilon \cdot N = 0.1$	$s=10^4$	$s=10^6$
Logistics $(N = 34)$	2	0	0	0.0019	0.007	0.0009
	4	0.0024	0.0046	0.0068	0.0057	0.0005
Logistics (N=45)	2	0.0002	0.0005	0.002	0.015	0.001
	4	0	0.003	0.004	0.008	0.0006
DRC-Drive (N=47)	2	0.0014	0.004	0.009	0.0072	0.0009
	4	0.001	0.008	0.019	0.0075	0.0011
Sequential (N=10)	2	0.0093	0.015	0.024	0.0063	0.0008
	4	0.008	0.024	0.04	0.008	0.0016

Table 1: Comparison of estimated errors with respect to the reference exact computation on various task trees.

Table 1 shows the results of the case study experiment. The quality of the solutions provided by using the KolmogorovApprox operator are better than those provided by the Trim and OptTrim operators, following the optimality guarantees, but is interesting to see that the quality gaps happen in practice in each of the examined task trees. However, in some of the task trees the sampling method produced better results than the approximation algorithm with KolmogorovApprox. Nevertheless, the approximation algorithm comes with an inherent advantage of providing an exact quality guarantees, as opposed to the probabilistic guarantees provided by sampling.

In order to better understand the quality gaps in practice between KolmogorovApprox, OptTrim, and Trim, we investigate their relative errors when applied on single random variables with support size n=100, and different support sizes of the resulting random variable approximation (m). In each instance of this experiment, a random variable is randomly generated by choosing the probabilities of each element in the support uniformly and then normalizing these probabilities so that they sum to 1.

Figure 1 present the error produced by the above methods. The depicted results are averages over several instances (50 instances) of random variables. The curves in the figure show the average error of  $\operatorname{OptTrim}$  and  $\operatorname{Trim}$  operators with comparison to the average error of the optimal approximation provided by  $\operatorname{KolmogorovApprox}$  as a function of m. According to the depicted results it is evident that increasing the support size of the approximation m reduces the error, as expected, in all three methods. However, errors produced by the  $\operatorname{KolmogorovApprox}$  are significantly smaller, a half of the error produced by  $\operatorname{OptTrim}$  and  $\operatorname{Trim}$ .

We also examined how our algorithm compares to linear programing as described and discussed, for example, in [15]. We ran an experiment to compare the run-time between the KolmogorovApprox algorithm with the run-time of a state-of-art implementation of linear programing. We used the "Minimize" function of Wolfram Mathematica and fed it with the equations  $\min_{\alpha \in \mathbb{R}^n} \|x - \alpha\|_{\infty}$  subject to  $\|\alpha\|_0 \le m$  and  $\|\alpha\|_1 = 1$ . The run-time comparison results were clear and persuasive, for a random variable with support size n = 10 and m = 5, the LP algorithm run-time was 850

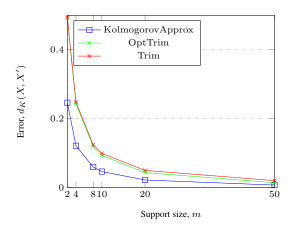


Figure 1: Error comparison between KolmogorovApprox,  $\operatorname{OptTrim}$ , and  $\operatorname{Trim}$ , on randomly generated random variables as function of m.

seconds, where the KolmogorovApprox algorithm run-time was less than a tenth of a second. For n=100 and m=5, the KolmogorovApprox algorithm run-time was 0.14 seconds and the LP algorithm took more than a day. Due to these timing results of the LP algorithm we did not proceed to examine it any further. Since it is not trivial to formally analyze the run-time of the LP algorithm, we conclude by the reported experiment that in this case the LP algorithm might not be as efficient as KolmogorovApprox algorithm whose complexity is proven to be polynomial in Theorem 12.

# 5 Discussion

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Compact representations of distributions is mentioned in the literature in various contexts for various applications. In this paper, we are interested in finding optimal approximation of a random variables under the Kolmogorov metric which we define as optimal m-approximation. In order to achieve this optimal approximation two steps were taken, find the support of the optimal random variable and then calculate the pmf of each and every value in that support to minimize the error. Proofs of existences, optimality and run-time were detailed in Section 3 and the main algorithm was presented, the KolmogorovApprox algorithm. Establishing the main contribution of this paper which is to present an optimal approximation scheme and to show it can be achieved in polynomial run-time. Furthermore, empirical evaluation was conducted on different domains and application to examine the algorithm performance in practice. We were interested in two aspects of performance - accuracy and run-time. Regarding to accuracy, as expected, the suggested KolmogorovApprox algorithm results much smaller error compared to the other methods, sometimes, in more then factor 0f 2. Regarding to run-time, KolmogorovApprox algorithm run-time is significantly much faster then LP approach. However, compared to other approximation methods accuracy vs. run-time is a trade off yet to be examined. Another interesting experiment that can be conducted in future work is to add the presented approach as one of the methods examined in [21] and compare it to the binning approaches.

As elaborated in the paper, our algorithm improves on the approach of Cohen, Shimony and Weiss [5] and [4] in that it finds an optimal two sided Kolmogorov approximation, and not just one sided. We consider this paper as a step in the examination of algorithms for optimal approximations of random variables. Beyond the Kolmogorov measure studied here we believe that similar approaches may apply also to total variation, the Wasserstein distance, and to other measures of approximations for other purposes.

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