Abstract

Determinantal Point Processes (DPPs) are a widely used probabilistic model for negatively correlated sets. DPPs have been successfully employed in Machine Learning applications to select a diverse, yet representative subset of data. In these applications, the parameters of the DPP need to be fitted to match the data; typically, we seek a set of parameters that maximize the likelihood of the data. The algorithms used for this task to date either optimize over a limited family of DPPs, or use local improvement heuristics that do not provide theoretical guarantees of optimality.

It is natural to ask if there exist efficient algorithms for finding a maximum likelihood DPP model for a given data set. In seminal work on DPPs in Machine Learning, Kulesza conjectured in his PhD Thesis (2012) that the problem is NP-complete. The lack of a formal proof prompted Brunel, Moitra, Rigollet and Urschel (2017a) to conjecture that, in opposition to Kulesza’s conjecture, there exists a polynomial-time algorithm for computing a maximum-likelihood DPP. They also presented some preliminary evidence supporting their conjecture.

In this work we prove Kulesza’s conjecture. In fact, we prove the following stronger hardness of approximation result: even computing a $1 - \frac{1}{\text{poly} \log N}$-approximation to the maximum log-likelihood of a DPP on a ground set of $N$ elements is NP-complete. At the same time, we also obtain the first polynomial-time algorithm that achieves a nontrivial worst-case approximation to the optimal log-likelihood: the approximation factor is $\frac{1}{(1+o(1)) \log m}$ unconditionally (for data sets that consist of $m$ subsets), and can be improved to $1 - \frac{1+o(1)}{\log N}$ if all $N$ elements appear in a $O(1/N)$-fraction of the subsets.

In terms of techniques, we reduce approximating the maximum log-likelihood of DPPs on a data set to solving a gap instance of a “vector coloring” problem on a hypergraph. Such a hypergraph is built on a bounded-degree graph construction of Bogdanov, Obata and Trevisan (2002), and is further enhanced by the strong expanders of Alon and Capalbo (2007) to serve our purposes.

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1. Introduction

Determinantal Point Processes (DPPs) are a family of probability distributions on sets that feature repulsion among elements in the ground set. Roughly speaking, a DPP is a distribution over all \(2^N\) subsets of \(\{1, \ldots, N\}\) defined by a positive semidefinite (PSD) \(N \times N\) matrix \(K\) (called a marginal kernel or correlation kernel) whose eigenvalues all lie in \([0, 1]\), such that, for any \(S \subseteq \{1, \ldots, N\}\), random subsets \(X\) drawn according to the distribution satisfy \(\Pr[S \subseteq X] = \det(K_S)\), where \(K_S\) is the principal submatrix of \(K\) indexed by \(S\).

DPPs were first proposed in quantum statistical physics to model fermion systems in thermal equilibrium (Macchi, 1975), but they also arise naturally in diverse fields such as random matrix theory, probability theory, random spanning trees and non-intersecting paths (Dyson, 1962; Burton and Pemantle, 1993; Rudnick and Sarnak, 1996; Soshnikov, 2000). After the seminal work of Kulesza and Taskar (2012), DPPs have attracted a flurry of attention from the machine learning community due to their computational tractability and excellent capability at producing diverse but representative subsets, and subsequently fast algorithms have been developed for sampling from DPPs (Hough et al., 2006; Kulesza and Taskar, 2010; Rebeschini and Karbasi, 2015; Li et al., 2016b,a; Anari et al., 2016; Dereziński et al., 2019; Launay et al., 2020). Furthermore, DPPs have since found a vast variety of applications throughout machine learning and data analysis, including text and image search, segmentation and summarization (Lin and Bilmes, 2012; Kulesza and Taskar, 2012; Zou and Adams, 2012; Gillenwater et al., 2012b,a; Yao et al., 2016; Kulesza and Taskar, 2011b; Affandi et al., 2014; Lee et al., 2016; Affandi et al., 2013b; Chao et al., 2015; Affandi et al., 2013a), signal processing (Xu and Ou, 2016; Krause et al., 2008; Guestrin et al., 2005), clustering (Zou and Adams, 2012; Kang, 2013; Shah and Ghahramani, 2013), recommendation systems (Zhou et al., 2010), revenue maximization (Dughmi et al., 2009), multi-agent reinforcement learning (Osogami and Raymond, 2019; Yang et al., 2020), modeling neural spikes (Snoek et al., 2013), sketching for linear regression (Dereziński and Warmuth, 2018; Dereziński et al., 2020), low-rank approximation (Guruswami and Sinop, 2012), and likely many more.

Maximum likelihood estimation. Many of these applications require inferring a set of parameters for a DPP capturing a given data set. As a DPP specifies a probability distribution, hence in contrast to supervised learning problems, the quality of a DPP cannot be estimated by the “error rate” of the model’s predictions. The standard approach to estimate a DPP from data is to find parameters that maximize the likelihood of the given data set being produced by a sample from the DPP (Kulesza and Taskar, 2012), i.e., the probability density of the observed data in the joint distribution. When the samples are identically and independently chosen from the DPP, the likelihood is the product of the probability densities the DPP assigns to the sampled subsets. The goal of the maximum likelihood estimator (MLE) method is to find a kernel matrix that maximizes the likelihood of the data set. Brunel et al. (2017b) showed that the maximum likelihood estimate indeed converges to the true kernel at a polynomial rate. In general, maximizing the likelihood of a DPP gives rise to a non-convex optimization problem, and has been approached with heuristics such as expectation maximization (Gillenwater et al., 2014), fixed point algorithms (Mariet and Sra, 2015), and MCMC (Affandi et al., 2014). In the continuous case, the problem has been studied under strong parametric assumptions (Lavancier et al., 2015), or smoothness assumptions (Baraud, 2013).
1.1. Our results

In spite of the wide applications of DPPs and the central role of the learning step, no efficient algorithms are known to find a maximum likelihood DPP. Instead, as mentioned above, two families of algorithms are known: one seeks to learn an optimal DPP within certain parameterized families of DPP structures (Kulesza and Taskar, 2012; Affandi et al., 2014; Gartrell et al., 2016; Mariet and Sra, 2016; Gartrell et al., 2017; Urschel et al., 2017; Dupuy and Bach, 2018), while the other invokes heuristics to maximize the likelihood in an unconstrained search over the parameter space (Kulesza and Taskar, 2011a; Gillenwater et al., 2014; Affandi et al., 2014; Mariet and Sra, 2015). Neither of these approaches provides any guarantees for how close the likelihood of the obtained parameters are to the maximum over all DPPs.

Indeed, Kulesza (2011a; 2012) conjectured over a decade ago that the problem of finding a set of parameters is NP-hard, but indicated that he was unable to formally establish a reduction: his thesis includes a sketch of a reduction from EXACT-3-COVER to a related problem\(^1\) with numerical evidence suggesting its correctness but without a formal proof. The subsequent literature adopted this belief, despite the lack of a solid theoretical foundation.

In this work, we resolve this question by proving Kulesza’s conjecture: computing maximum likelihood DPP kernels is indeed NP-hard. In fact, we establish a stronger, gapped hardness result:

**Theorem 1 (Informal Statement of the Main Theorem)** There is a ground set of size \(N\) such that it is NP-hard to \((1 - O((\log^k N))^{-\alpha})\)-approximate the maximum DPP log-likelihood value of a training set.

**Remark 2** Some comments on our (somewhat confusing) convention of approximation factors are in place. Since log likelihood functions are always negative real numbers and it is a bit awkward to work with optimizing negative quantities, we therefore add a minus sign at the front of our definition of log likelihood functions. Consequently, we minimize log likelihood functions instead of maximizing them. On the other hand, as our hard learning instances are reduced from MAX-3SAT and 3-COLORING, to be consistent with hardness results in the literature on these problems, we use \(\alpha\)-approximation (where \(0 < \alpha < 1\), for maximization problems) in the statements of our hardness and algorithmic results. Note that here “\(\alpha\)-approximation” actually means that the log likelihood function (in our definition and ought to be minimized) output by an algorithm is at most \(\frac{1}{\alpha}\) time the optimal log likelihood function.

Therefore, the difficulty of learning a DPP is not tied to any particular representation of the marginal kernel, as in fact estimating the maximum likelihood value itself is NP-hard. Note, however, that many problems in learning are hard merely due to the difficulty of finding a specific representation (Pitt and Valiant, 1988), which is not the case for our problem.

The NP-hardness of maximum likelihood learning naturally raises the question of whether any nontrivial approximation is possible. We show that such an approximation is possible: we present a simple, polynomial-time algorithm obtaining a \(\frac{1}{(1+o(1))\log m}\)-approximation for a data set with \(m\) subsets.

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\(^1\) Technically, the reduction proposed by Kulesza targets a variant of the maximum-likelihood DPP learning problem in which the instance specifies a set of positive-semidefinite matrices along with the data, and the objective is to find a DPP kernel in the cone of the given matrices that maximizes the likelihood.
Theorem 3 (Informal Statement of the Approximation Algorithm) There is a polynomial-time approximation algorithm achieving the following: on an input data set consisting of \( m \) subsets over a ground set of size \( N \), it returns a kernel that \( \frac{1}{(1+o(1)) \log m} \)-approximates the maximum log likelihood. Moreover, if every element in the ground set appears in at most \( O(\frac{1}{N}) \)-fraction of the subsets, the kernel achieves a \( (1 - \frac{1+o(1)}{\log N}) \)-approximation to the maximum log likelihood.

We stress that in contrast to the prior work on learning DPP kernels with guarantees (Urschel et al., 2017), our algorithm does not rely on the data being produced by a DPP to have a “cycle basis” of short cycles or large nonzero entries. We obtain an approximation to the optimal likelihood for any data set. Although a \( \frac{1}{(1+o(1)) \log m} \)-approximation is weak, when every element appears in relatively few subsets (which is common in practice), our algorithm is much better: the actual approximation factor is \( 1 - \frac{1}{\log(m/a_{\text{max}})} \), where \( a_{\text{max}}/m \) is the fraction of the data subsets containing the most frequent element in the ground set. Hence, if all elements appear in at most a \( \sim \frac{1}{N} \)-fraction of the subsets, we obtain a \( (1 - \frac{1+o(1)}{\log N}) \)-approximation to the log likelihood. Although we don’t expect our algorithm to obtain substantially better likelihood than various heuristics employed in practice, it may nevertheless serve as a benchmark to estimate how close to optimal these solutions are. Moreover, the family of instances constructed in our reduction indeed satisfies this condition; therefore, to improve the hardness of approximation bound beyond \( 1 - \frac{1+o(1)}{\log N} \), the hard instance of data set must have some elements appearing in \( \omega(\frac{1}{N}) \)-fraction of the subsets.

1.2. Our approach and techniques

We show that it is NP-hard to approximate the optimal DPP likelihood function by reducing from a coloring problem, rather than from EXACT-3-COVER, which was Kulesza’s (2012) initial approach.

We begin with some intuition leading to a notion of vector coloring that we use in the reduction. As any marginal kernel \( K \in \mathbb{R}^{N \times N} \) is positive semidefinite, it can be factored as \( K = Q^\top Q \), where \( Q \in \mathbb{R}^{k \times N} \), \( Q^\top \) stands for the transpose of \( M \), and \( k \) is called the dimension of the kernel. If we denote the column vectors of \( Q \) by \( q_1, \ldots, q_N \), each \( q_i \in \mathbb{R}^k \), then one can think of these \( q_i \)'s as providing an embedding of the elements in \( \{1, \ldots, N\} \) into the space \( \mathbb{R}^k \), and the embedding vectors capture similarities among elements. Specifically, the preference of DPPs for diverse subsets, roughly speaking, stems from the following fact: if a subset \( S \) includes elements that are similar, the submatrix \( K_S \) would contain columns that are nearly co-linear embedding vectors, and hence its determinant (and correspondingly, the probability that \( S \) is the random subset generated by the marginal kernel \( K \)) is close to zero.\(^2\)

Consider, for simplicity, a training set that consists of a collection of subsets of \( \{1, \ldots, N\} \), each of size \( r \) where \( r \) is a constant. What can we say about a maximum-likelihood DPP kernel for such a data set? Ideally, the embedding vectors should encode an “\( r \)-vector-coloring” of the elements in the following sense. Each of the \( r \) colors is represented by a unit vector (after normalizations) in an orthonormal basis; to maximize the likelihood function, every subset \( S = \{i_1, \ldots, i_r\} \) that appears in the training set corresponds to a “rainbow coloring” of the embedding vectors \( \{q_{i_1}, \ldots, q_{i_r}\} \) (“rainbow coloring” means that the \( r \) embedding vectors are all colored differently), while for the

\(^2\) Recall that, since \( K \) and hence \( K_S \) are Gram matrices, \( \det(K_S) \) is equal to the square of the volume of the parallelepiped spanned by the embedding vectors of elements in \( S \).
r-subsets that do not appear in the training set, we would like as many as possible of them to contain some repeated color.\(^3\)

Thus, it is natural to attempt a reduction from \textsc{Graph r-Coloring} to Maximum Likelihood Learning of DPP. However, if we view each edge as a 2-subset, then we fail to get a hard problem to begin with, since graph 2-coloring is easy. We overcome this by “lifting” each edge to a size-3 subset (or equivalently, we transform a graph into a 3-uniform hypergraph, and view all its hyperedges as size-3 subsets in the data set) so that we can still work with \textsc{3-Colorability}. On an input graph \(G = (V, E)\), our goal would be to show the following: if \(G\) is 3-colorable, then there is a DPP kernel whose likelihood is large (completeness); if \(G\) is not 3-colorable, then the likelihood of every DPP kernel is small (soundness).

As the column vectors of DPP kernels are in Euclidean spaces instead of discrete ones, the continuous variant of coloring that works for us is the notion of \textit{vector coloring} of graphs. A \textit{vector coloring} of a graph \(G = (V, E)\) is a mapping from vertices in \(V\) to points (vectors) in some low-dimensional metric space \(M\), such that the presence or absence of edges between any pair of vertices prescribes the value of the inner product between the two corresponding vectors. (See Section 1.3.)

Our problem differs from this one in two important ways: first, we do not care too much about the minimum dimension of the Euclidean space in which a vector representation exists; second, which is more subtle and will be elaborated below, we need a “gapped” reduction.

Secondly, instead of a “decision” hardness result on vector coloring of hypergraphs (e.g. it is \textit{NP}-hard to decide if there is a \(3\)-dimensional orthogonal representation for the elements in the set that satisfies certain orthogonality conditions), we rather require a “\textit{gapped}” reduction, obtaining something like “starting with a \textit{Yes} instance, we end up with a set of embedding vectors so that the average volume of the \(3\)-dimensional parallelepipeds spanned by these embedding vectors of hyperedges is large; starting with a \textit{No} instance, then every possible embedding scheme will make the average volume of those parallelepipeds small”. Namely, in an “averaging” sense, we need the resulting hypergraph transformed from a \textit{No} instance to be “far from” \(3\)-vector colorable. Accordingly, the \textit{No} instance of \textsc{3-Colorability} should have the property that, even after removing a small fraction of the edges, it is \textit{still not} \(3\)-colorable. On the other hand, the strongest known hardness results (Khanna et al., 2000; Guruswami and Khanna, 2004; Dinur et al., 2009; Brakensiek and Guruswami, 2016) on coloring \(3\)-colorable graphs are based on dense graphs; the requirement on \textit{No} instance mentioned above, when applied to dense graphs, would make the problem fall into the regime of property testing, which is unfortunately known to be computationally easy (Goldreich et al., 1998).

We circumvent this obstacle by adapting a \textit{sparse graph} construction of Bogdanov, Obata and Trevisan (2002) (referred to as \textit{BOT} graph henceforth). Based on the hardness of approximating \textsc{Max-3SAT}, \textit{BOT} graph was used in Bogdanov et al. (2002) to prove query lower bound for testing

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3. Implicitly, we would also like to have \(k = r\) so that no non-degenerate parallelepiped of dimension higher than \(r\) exists, as the number of size-\((r + 1)\) or larger subsets dominates those of size-\(r\) subsets; see Conjecture 12 below.
3-COLORABILITY in the bounded-degree model. We fix some minor mistakes in the construction and analysis of Bogdanov et al. (2002), further enhance the robustness of BOT graph with the strong expander construction of Alon and Capalbo (2007). These modifications allow us to show that, for some absolute constant $\delta$, we can decode a 3-coloring of the vertices which satisfies at least $(1 - \delta)$-fraction of the edges in the original BOT graph, as long as the DPP log likelihood of a training set constructed from the edges of the BOT graph were close enough to the maximum value of a 3-colorable graph. An overview of the reduction sequence is illustrated in Figure 1.

Algorithmic results. For the upper bound, we obtain an approximation algorithm by using some of the properties required for the analysis of our reduction. The algorithm itself is very simple: given a data set $X_1, X_2, \ldots, X_m$, output the DPP marginal kernel given by the $N \times N$ diagonal matrix $K$ such that $K_{ii} = |\{j : X_j \ni i\}|/m$ for all $i$ in the ground set. In other words, the diagonal entries of the marginal kernel are just the empirical probabilities of elements in the data set. Hadamard’s inequality gives a lower bound on the optimal likelihood that is similar to the likelihood of the diagonal kernel; if the elements all appear in at most a $a_{\text{max}}/m$-fraction of the subsets, the ratio $\frac{\log \text{likelihood output by the algorithm}}{\log \text{optimal log likelihood}}$ is at most $1 + \frac{\log((1-a_{\text{max}}/m)^{1-a_{\text{max}}/m})}{\log((a_{\text{max}}/m)^{a_{\text{max}}/m})} \approx 1 + \frac{1}{\log N}$ when $a_{\text{max}}/m$ is $O(1/N)$. For an unconditional upper bound, observe that elements in all $m$ sets should occur with probability 1 in the maximum likelihood DPP (and thus may be disregarded without loss of generality), hence we may plug $a_{\text{max}} = m - 1$ into the aforementioned bound and obtain a $(1 + o(1)) \log m$ upper bound on the ratio.

1.3. Related work

Learning DPPs. As mentioned earlier, Urschel et al. (2017) in particular proposed an algorithm for recovering a DPP’s kernel up to similarity, which is efficient when (i) the graph defined by interpreting the kernel as a weighted adjacency matrix has a “cycle basis” of cycles of bounded length and (ii) the nonzero entries are not too small. Furthermore, they gave a lower bound on the sample complexity of estimating the DPP kernel, showing that it indeed depends similarly on these quantities. Thus, when there is enough data to permit exact recovery of the kernel, this algorithm will perform well, but otherwise there is no guarantee that the algorithm produces a kernel for a DPP with likelihood close to maximum.

In a companion paper, Brunel et al. (2017a) also studied the rate of estimation obtained by the maximum likelihood kernel. Again, they determined classes of DPPs for which it is efficient (or not). Moreover, they identified an exponential number of saddle points, and conjectured that these are the only critical points; they further suggested that a proof of this conjecture might lead to an efficient algorithm for computing a maximum likelihood kernel. But, they did not actually provide algorithms for computing the likelihood or the kernel itself.
Starting with the pioneering work of Kulesza and Taskar (2011a), various empirical learning algorithms have been proposed for learning discrete DPPs, such as Bayesian methods (Affandi et al., 2014), expectation-maximization (EM) algorithms (Gillenwater et al., 2014), fixed-point iteration (Mariet and Sra, 2015), learning non-symmetric DPPs (Gartrell et al., 2019), learning with negative sampling (Mariet et al., 2019), and minimizing Wasserstein distance (Anquetil et al., 2020). However, none of these algorithms has theoretical guarantees. Efficient learning algorithms have also been designed for restricted classes of DPPs (Mariet and Sra, 2016; Gartrell et al., 2017; Dupuy and Bach, 2018; Osogami et al., 2018). A related problem, namely testing DPPs, recently has been investigated by (Gatmiry et al., 2020).

We note that in contrast to the problem of learning the DPP kernel from a data set as considered here, the problem of computing the mode (“MAP estimate”) of a DPP given by its kernel has long been known to be NP-complete (Ko et al., 1995; Civril and Magdon-Ismail, 2009). The inapproximability for this problem was recently strengthened substantially by (Ohsaka, 2021).

**Vector coloring problems.** The notion of *orthogonal representation* (in which there is an edge between two vertices if and only if the two corresponding representation vectors are orthogonal) was introduced by Lovász (1979), and was used in the definition of the famous Lovász’s $\vartheta$ function, which has been employed to bound quantities such as Shannon capacity, the clique numbers or the chromatic numbers of graphs. More generally, a *geometric representation* of a graph is a mapping from vertices in $V$ to points in a metric space $M$, such that two vertices are connected by an edge if and only if the distance between the two corresponding points satisfies certain condition. For example, orthogonal representation is a special case of the *unit-distance* graph where (in the framework of geometric representation) the underlying metric space is the unit sphere (with distance 1 replaced by sphere distance $\pi/2$). Geometric representation of graphs is a well-studied subject, revealing many surprising connections between parameters (e.g. dimension) of geometric representations and properties of the original graph, such as chromatic number, connectivity, excluded subgraphs, tree width, planarity, etc; see e.g. (Lovász, 1979; Lovász et al., 1989; Parsons and Pisanski, 1989; Karger et al., 1998; Lovász and Vesztergombi, 1999; Lovász et al., 2000; Haynes et al., 2010) and the recent textbook (Lovász, 2019).

**Matrix completion problem.** Geometric representations of graphs are intimted connected to another class of problems, *matrix completion problems*. For instance, the celebrated result of Peeters (1996), showing NP-hardness of deciding whether a 3-dimensional orthogonal representation over a finite field exists for a graph, was obtained through reducing *3-COLORABILITY* to a low-rank matrix completion problem. Matrix completion studies under what conditions a partially specified matrix can be completed into one which belongs to a certain class of matrices, such as low-rank matrices, semidefinite matrices, Euclidean distance matrices, etc. See e.g. Laurent (2009) for an overview of the important results in this area. Interestingly, a recent work of Hardt et al. (2014), which proved the hardness of low-rank matrix completion problem under the incoherence assumption (a commonly used assumption for many matrix completion results), was also based on gapped versions of computationally hard problems on graphs such as the $r$-*COLORING* problem and the $(r, \epsilon)$-*INDEPENDENT-SET* problem.5

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4. Some authors, for example Lovász (1979), define orthogonal representation by mandating the vectors of two non-adjacent vertices to be orthogonal.

5. In this problem, one is given an undirected graph that is promised to be $r$-colorable and is asked to find an independent set of size $\epsilon n$, where $\epsilon < 1/r$ and $n$ is the number of the vertices in the graph.
2. Maximum likelihood learning of DPP and our main hardness result

2.1. Preliminaries

Unless stated otherwise, all logarithms in this paper are to the base $e$ (i.e. natural logarithms). For positive integer $n$, we write $[n]$ to denote the set $\{1, 2, \ldots, n\}$. For an $n$-dimensional real vector $x$, we use $\|x\|_2 = \sqrt{x_1^2 + \cdots + x_n^2}$ to denote the $\ell_2$ or Euclidean norm of $x$.

**Matrix analysis.** Let $A$ be an $m \times n$ matrix. The $(i, j)^{th}$ entry of $A$ will be denoted by $A_{i,j}$. All matrices in this paper are over real numbers $\mathbb{R}$; therefore the Hermitian adjoint of $A$, $A^H$ is the same as $A^\top$, the transpose of $A$. By the spectral theorem, the eigenvalues of a real, symmetric matrix $M \in \mathbb{R}^{n \times n}$ are all real numbers, and will be denoted $\lambda_1(M) \geq \lambda_2(M) \geq \cdots \geq \lambda_n(M)$. A real, symmetric matrix $M$ is called positive semidefinite (PSD) if all its eigenvalues are non-negative (i.e., $\lambda_n(M) \geq 0$). Well-known equivalent characterizations of PSD matrices include $x^\top M x \geq 0$ for all $x \in \mathbb{R}^n$, and the existence of a matrix $Q \in \mathbb{R}^{k \times n}$ for some $k > 0$ such that $M = Q^\top Q$.

A useful variational characterization of the eigenvalues of real, symmetric matrices is the Courant-Fischer theorem, which states that for every $1 \leq k \leq n$ (when a set of vectors whose indices are outside the range $[n]$, then the set is understood to be empty) we have

$$\lambda_k(A) = \min_{x_1, \ldots, x_{k-1} \in \mathbb{R}^n} \max_{y \neq 0, y \perp x_1, \ldots, x_{k-1}} \frac{y^\top A y}{y^\top y},$$

and

$$\lambda_k(A) = \max_{x_{k+1}, \ldots, x_n \in \mathbb{R}^n} \min_{y \neq 0, y \perp x_{k+1}, \ldots, x_n} \frac{y^\top A y}{y^\top y}.$$

The singular values of a matrix $A \in \mathbb{R}^{m \times n}$ are defined as the (positive) square roots of the eigenvalues of $A^H A = A^\top A$ (a real, symmetric $n \times n$ matrix). Namely, $\sigma_i(A) = \sqrt{\lambda_i(A^H A)}$, $i = 1, \ldots, n$. We also arrange the singular values of a matrix $A$ in decreasing order, that is $\sigma_1(A) \geq \sigma_2(A) \geq \cdots \geq \sigma_n(A)$. The Frobenius norm of $A$, denoted $\|A\|_F$, is defined to be $\|A\|_F = \sqrt{\sum_{i=1}^m \sum_{j=1}^n |A_{i,j}|^2}$. It is well-known that $\|A\|_F^2 = \sigma_1^2(A) + \cdots + \sigma_n^2(A)$. Finally, the spectral norm of a square $n \times n$ matrix $A$ is defined as the square root of the maximum eigenvalue of $A^H A$, i.e.,

$$\|A\|_2 = \sqrt{\lambda_1(A^H A)} = \max_{x \neq 0} \frac{\|Ax\|_2}{\|x\|_2} = \sigma_1(A).$$

**Discrete determinantal point processes.** A discrete determinantal point process (DPP) $P$ over a finite set $\mathcal{X}$ is a probability measure over the set of all subsets of the ground set $\mathcal{X}$. The distribution of $P$ is specified by a marginal kernel $K \in \mathbb{R}^{\mathcal{X} \times \mathcal{X}}$, which is a positive semidefinite matrix with eigenvalues in $[0, 1]$, in the following manner: if $Y \subseteq \mathcal{X}$ is a random subset drawn according to $P$, then its probability mass function $P_K$ is defined such that, for every $S \subseteq \mathcal{X}$,

$$\Pr_{Y \sim P_K}[S \subseteq Y] = \det(K_S).$$

Here $K_S$ is the principal submatrix of $K$ indexed by $S \subseteq \mathcal{X}$.
If it is the case that all eigenvalues of $K$ are in $[0,1)$, then $\mathcal{P}$ is called an $L$-ensemble, whose kernel can be defined to be the positive definite\textsuperscript{6} matrix $L = K(I - K)^{-1}$. In this case, the corresponding probability mass function, denoted $\mathcal{P}_L$, can be shown to be

$$Pr_{Y \sim \mathcal{P}_L}[Y = S] = \frac{\det(L_S)}{\det(I + L)}$$

for every $S \subseteq \mathcal{X}$. Hence, $Pr_{Y \sim \mathcal{P}_L}[Y = \emptyset] = \det(I - K)$, and consequently a DPP is an $L$-ensemble if and only if the random variable $Y = \emptyset$ with non-zero probability.

### 2.2. Maximum Likelihood Learning of DPPs

We define the problem of Maximum Likelihood Learning of DPPs as follows. A learning algorithm receives a training data set $\{X_t\}_{t=1}^{T'}$ (viewed as a multiset), typically drawn independently and identically from a distribution $D$ over the subsets of a ground set $\mathcal{X}$. The goal of the learning algorithm is to find a DPP kernel $K$ based on the training set that minimizes\textsuperscript{7} the following maximum log likelihood estimator

$$\ell(K) = -\frac{1}{T'} \log \prod_{t=1}^{T'} Pr_{Y \sim \mathcal{P}_K}[Y = X_t] = -\frac{1}{T'} \sum_{t=1}^{T'} \log Pr_{Y \sim \mathcal{P}_K}[Y = X_t],$$

where $\{X_1, \ldots, X_{T'}\}$ is the set of distinct elements in the multiset $\{X_t\}_{t=1}^{T'}$. When the training data set $\{X_t\}_{t=1}^{T'}$ is clear from context, we simply denote the value of the maximum log likelihood of an optimal DPP kernel by $\ell^*$.  

One common way to establish the hardness of maximum likelihood learning problems is to show that even computing the maximum value of the log likelihood $\ell^*$ is hard (if one could efficiently find an optimal DPP kernel $K$, then since evaluations of determinants can be performed in polynomial time, clearly the corresponding log likelihood $\ell^*$ would be efficiently computable as well). That is also the approach we take in this paper. In fact, the lower bound actually proved is much stronger: we show that it is NP-hard even to compute a $1 - O(1/\log N)$-approximation of $\ell^*$ for a ground set of size $N$.

**Theorem 4 (Main)** There are infinitely many positive even integers $N$ such that the following holds.\textsuperscript{8} Let $\mathcal{X} = \{1, 2, \ldots, N\}$. There is a training data set $\{X_t\}_{t=1}^{N/2}$ of size $N/2$, where $X_i \subseteq \mathcal{X}$ for each $i$, such that it is NP-hard to $\left(1 - O\left(\frac{1}{\log N}\right)\right)$-approximate the maximum log likelihood value that a DPP kernel can achieve on the training set.

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\textsuperscript{6} A real, symmetric matrix is called positive definite (PD) if all its eigenvalues are positive.

\textsuperscript{7} We add a negative sign at the front to make the estimator to be always positive, and thus changing the maximization problem into a minimization one. To be consistent, we still call the quantity as a “maximum” likelihood estimator; see Remark 2.

\textsuperscript{8} As we will see later that $N/2$ is the number of edges in a specially constructed graph. We then construct a 3-uniform hypergraph based on this graph, and add a new node to the hypergraph for each edge in the graph. The ground set consists of these newly added nodes together with the set of vertices in the original graph, hence the cardinality of the ground set is at most $N$. 

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2.3. Proof of the Main Theorem: an outline

\textbf{MAX-3SAT with bounded occurrence.} Our starting point is the hardness of MAX-3SAT, in which given a Boolean formula \( \phi \) in 3-CNF form, the goal is to output the maximum number of clauses of \( \phi \) that can be satisfied by any truth assignment of the variables. A classical hardness result is Håstad’s 3-bit PCP theorem (Håstad, 2001), which states that it is NP-hard to \((7/8 + \epsilon)\)-approximate MAX-3SAT for any constant \( \epsilon > 0 \). However, for our purposes, we need the formula \( \phi \) to have bounded occurrences of any variable. Let MAX-3SAT\((k)\) denote a subclass of MAX-3SAT, in which the instances satisfy that every variable occurs in at most \( k \) clauses. Håstad (2000) showed that it is NP-hard to \(7/8 + 1/(\log k)^c\)-approximate MAX-3SAT\((k)\) where \( c \) is some absolute constant. Therefore,

\textbf{Lemma 5 (Håstad (2000))} There is a constant integer \( k \) and constant \( \epsilon > 0 \) (depending only on \( k \)) such that it is NP-hard to \((1 - \epsilon)\)-approximate MAX-3SAT\((k)\).

That is, for infinitely many integers \( n \), there are two families of instances \( \phi_Y \) and \( \phi_N \) in MAX-3SAT\((k)\) of size \( n \) each with the following property: \( \phi_Y \) is satisfiable; every truth assignment can satisfy at most an \( 1 - \epsilon \) fraction of the clauses in \( \phi_N \); and it is NP-hard to distinguish between the two cases.

\textbf{3-COLORING for bounded degree graphs.} Next, we adapt a gap-preserving reduction of Bogdanov, Obata and Trevisan (2002), which was originally used to prove an \( \Omega(n) \) query lower bound for testing 3-Colorability in bounded-degree graphs under the property testing model. On input an instance \( \phi \) of MAX-3SAT\((k)\), the reduction outputs a degree-bounded graph \( G_\phi \) (BOT graph) which satisfies the following: if \( \phi \) is satisfiable then \( G_\phi \) is 3-colorable; and if every truth assignment can satisfy at most \( 1 - \epsilon \) fraction of the clauses in \( \phi \) then every 3-coloring of the vertices of \( G_\phi \) can make at most \( 1 - \epsilon' \) fraction of the edges in \( G_\phi \) non-monochromatic. Here \( \epsilon' \) is a constant depending only on \( \epsilon \) and \( k \).

\textbf{Lemma 6 (Bogdanov et al. (2002))} There are absolute constants \( d \) and \( \epsilon' > 0 \) such that the following holds. For infinitely many integers \( n \), there are two degree-d bounded graphs \( G_{\phi,Y} \) and \( G_{\phi,N} \) of size \( n \) such that: \( G_{\phi,Y} \) is 3-colorable; no \( 1 - \epsilon' \) fraction of the edges of \( G_{\phi,N} \) is 3-colorable; and yet it is NP-hard to distinguish between the two cases.

\textbf{Very strong expanders.} The main idea of the reduction of Bogdanov et al. (2002) is to make \( k \) copies of \textsc{True}, \textsc{False} and \textsc{Dummy} for each variable and its negation, and use an expander to connect these copies together to ensure truth value consistency among different copies. Any constant-degree expander with reasonable vertex-expansion suffices: on one hand, the resulting graph \( G_\phi \) is of bounded degree; on the other hand, by the expansion property, deleting a small fraction of the edges in \( G_\phi \) will still leave the graph with a large connected component, using which, one can — from the coloring of \( G_\phi \) — decode a satisfying assignment that satisfies most of the clauses.

However, for the purpose of proving hardness of DPP Maximum Likelihood Learning, we need the expander to have some additional properties, which are encapsulated in the following definition.

\footnote{We may also use the NP-hardness results of Berman et al. (2003) for 3-SAT instances in which every variable appears exactly 4 times, or assuming \( \text{RP} \neq \text{NP} \), use the hardness result of Trevisan (2001) with better parameters.}

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Definition 7 (Very strong expanders (Alon and Capalbo, 2007)) A graph $G = (V, E)$ is called a $d$-regular very strong expander on $n$ vertices if the average degree in any subgraph of $G$ on at most $n/10$ vertices is at most $d/6$, and the average degree in any subgraph of $G$ on at most $n/2$ vertices is at most $2d/3$.

The nice properties of very strong expanders that we require are summarized in the following theorem from Alon and Capalbo (2007).

Theorem 8 (Alon and Capalbo (2007)) Let $G = (V, E)$ be a very strong $d$-regular expander on $n$ vertices. If we delete an arbitrary subset of $m' \leq nd/150$ edges from $G$ and denote the resulting graph by $G'$, then $G'$ contains a subgraph $H$ on at least $n - 15m'/d$ vertices and the diameter of $H$ is $O(\log n)$.

The known explicit constructions of Ramanujan graphs (Lubotzky et al., 1988; Margulis, 1988) yield families of $d$-regular strong expanders on $n$ vertices for infinitely many $n$’s.

3-uniform hypergraph. To obtain the training data set, we transform a BOT graph $G_\phi = (V, E)$ into a 3-uniform hypergraph $H_\phi = (V', E')$ as follows. The vertex set $V'(H_\phi)$ is $V(G) \cup E(G)$, and for notational convenience, we will simply label the “graph-vertex” vertices by $a_v$ for every $v \in V(G)$, and label the “graph-edge” vertices in $V'(H_\phi)$ by $a_{(u,v)}$ for every edge $(u,v) \in E(G)$. Then the set of hyper-edges is $E'(H_\phi) = \{(a_u, a_v, a_{(u,v)}) : (u,v) \in E(G)\}$. It follows that $H_\phi$ is a 3-uniform hypergraph with $N = |V'(H_\phi)| = n + m$ and $|E'(H_\phi)| = m$, where $n$ and $m$ denote the number of vertices and edges of the BOT graph $G_\phi$, respectively.

Now, what happens if we use the set of hyper-edges of $H_\phi$ as the training data set and feed it to a DPP Maximum Likelihood Learning algorithm? Our first step in understanding the optimal DPP kernel is to establish a connection between DPP kernel of learning BOT hypergraphs and a problem called “vector coloring”.

Connecting DPP kernels with vector colorings. Since $K$ is a positive semidefinite matrix, we can write $K$ as $K = Q^T Q$ for some matrix $Q$. Let $q_1, \ldots, q_N \in \mathbb{R}^k$ be the columns of $Q$. We can further decompose these vectors as $q_i = \|q_i\|_2 \chi_i$, where $\chi_i \in \mathbb{R}^k$ is a unit vector. The quantity $\|q_i\|_2$ is a measure of the “importance” of item $i$, and $\chi_i$ is a normalized vector which encodes diversity features of item $i$. Now the entries of the marginal kernel satisfy $K_{ij} = \|q_i\|_2^2 \chi_i^T \chi_j \|q_j\|_2^2$, where $\chi_i^T \chi_j \in [-1,1]$ is a signed measure of the similarity between item $i$ and item $j$. In particular, the diagonal entries satisfy that $K_{ii} = \|q_i\|_2^2$ for every $i \in [N]$.

We prove the following theorem, which allows us to somewhat decouple $\|q_i\|_2$ and $\chi_i$ for each item $i$, and identify (from the training set) the value of the “importance” (i.e. value $\|q_i\|_2$) for each item.

Theorem 9 Let $K$ be a marginal kernel with likelihood $\ell(K)$. Then there exists a marginal kernel $K'$ with $\ell(K') \leq \ell(K)$ such that the diagonal of $K'$ (indexed by vertices and edges of $G_\phi$) satisfies

$$
K'_{ii} = \begin{cases} 
\deg_{G_\phi}(u) / m & \text{for } i = u \in V(G_\phi); \\
\frac{1}{m} & \text{for } i = (u,v) \in E(G_\phi).
\end{cases}
$$

10. In order to not overload the statement in Theorem 4 with multiple parameters, we may add isolated vertices to graph $G_\phi$ so that $n = m$ and hence the ground set size is $N$ and the sample size is $m = N/2$.

11. It is no coincidence that our simple algorithm, using this “first-moment” information from the training set in a similar manner, constructs its DPP that achieves nontrivial worst-case approximation to the optimal log likelihood.
Thus, it remains to determine the diversity features that maximize the likelihood. To this end, we use a variant of 3-colorability in which the “colors” are generalized to vectors in \( \mathbb{R}^k \) and the “coloring constraint” for an edge \((i, j)\) is satisfied if the vectors \( \chi_i \) and \( \chi_j \) assigned to the vertices \( i \) and \( j \) are orthogonal.

More formally, let \( S^{k-1} \) be the unit sphere in \( k \)-dimensional Euclidean space; that is, \( S^{k-1} = \{ x \in \mathbb{R}^k : \| x \|_2 = 1 \} \). Given a graph \( G = (V, E) \), we define a vector \( k \)-coloring of \( G \) to be a function \( \chi : V(G) \rightarrow S^{k-1} \). We say that a vector \( k \)-coloring \( \chi \) is orthogonal if, for every edge \((u, v) \in E(G)\), we have \( \chi_u^\top \chi_v = 0 \). We define the error of a vector \( k \)-coloring \( \chi \) of \( G \) to be

\[
\text{err}_\chi(G) := \frac{1}{|E(G)|} \sum_{(u,v) \in E(G)} |\chi_u^\top \chi_v|^2
\]

so that a vector \( k \)-coloring \( \chi \) of \( G \) is orthogonal if and only if \( \text{err}_\chi(G) = 0 \). Since \( \chi_u \) and \( \chi_v \) are unit vectors, \( |\chi_u^\top \chi_v| = |\cos \theta_{uv}| \) for all \((u, v) \in E(G)\), where \( \theta_{uv} \) is the angle between \( \chi_u \) and \( \chi_v \).

Now, by combining Theorem 9 with the fact that any 3-coloring of \( G \) naturally induces a vector 3-coloring of \( G \), it is not hard to prove the following “completeness” theorem.

**Theorem 10 (Completeness theorem)** Let \( G_\phi \) be a BOT graph, and let \( n = |V(G_\phi)| \) be the number of vertices and \( m = |E(G_\phi)| \) be the number of edges of \( G_\phi \), respectively. If \( \phi \) is satisfiable, then there exists a rank-3 DPP marginal kernel \( K \) such that

\[
\ell(K) = \ell^* = 3 \log m - \frac{1}{m} \sum_{(u,v) \in E(G_\phi)} \left( \log(\deg_{G_\phi}(u)) + \log(\deg_{G_\phi}(v)) \right).
\]

**Projecting DPP kernels to \( \mathbb{R}^3 \).** Intuitively, the maximum likelihood marginal kernel has dimension 3 so that zero probability measure will be assigned for subsets of size at least 4. We were unable to prove this, but we nevertheless manage to show that the loss in making such an assumption is not too great:

**Theorem 11** Let \( G_\phi \) be a BOT graph with maximum degree at most \( k \). There is a constant \( C_k \) depending only on \( k \) such that the following holds. Let \( K \) be an optimal marginal kernel with likelihood \( \ell(K) \leq \ell^* + \delta \) for some \( 0 < \delta \leq 1/(128k)^2 \), then there exists a marginal kernel \( K' \) of dimension 3 such that \( \ell(K') \leq \ell^* + C_k \delta^{1/4} \).

We conjecture that an even stronger statement is in fact true.

**Conjecture 12 (Cardinality-rank conjecture)** If the cardinality of every subset in a training set is at most \( k \geq 1 \), then every optimal maximum likelihood marginal kernel for the training set has dimension at most \( k \).

This conjecture may be of independent interest outside the realm of maximum likelihood learning of DPPs.

**Decoding truth assignments from vector colorings.** Because of Theorem 11, from now on we assume that the dimension of \( Q \) is 3. Therefore, for each \((a_u, a_v, a_{(u,v)}) \in E(H_\phi), \)

\[
\Pr_{Y \sim P_K}[Y = \{a_u, a_v, a_{(u,v)}\}] = \det(K_{\{a_u, a_v, a_{(u,v)}\}}),
\]

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where $K = Q^TQ$ is the marginal kernel of DPP $P$. To maximize the likelihood, we want to maximize the product of determinants of the above form. Since each $a_{(u,v)}$ occurs in only one example, we can assume that $\chi_{a_{(u,v)}}$ is always taken to be orthogonal to $\chi_{a_u}$ and $\chi_{a_v}$. Thus, the likelihood contribution from $(a_u, a_v, a_{(u,v)})$ is maximized when $\chi_{a_u}^\top \chi_{a_v} = 0$; or equivalently, when $\chi_{a_u}$ and $\chi_{a_v}$ are orthogonal. We formally prove this correspondence between the “orthogonality” of the associated vector 3-coloring of $G_\phi$ and the likelihood of the corresponding DPP with marginal kernel $K = Q^TQ$, where the column of $Q$ corresponding to vertex $u$ is $\|q_u\|_2 \chi_{a_u}$. Moreover, we can even decode the truth-assignment of the Boolean formula $\phi$ if the vector 3-coloring of $G_\phi$ is very close to satisfying all edges of $G_\phi$.

**Theorem 13 (Soundness theorem)** Let $\ell^*$ be the optimal log likelihood as in Theorem 10. Then there exists a constant $C > 0$ which depends only on $k$ and $\epsilon'$ as those defined in Theorem 6, such that the following holds. If there is a DPP marginal kernel $K$ of rank 3, which satisfies $\ell(K) \leq \ell^* + \frac{C}{\log^k n}$ where $n = |V(G_\phi)|$ is the number of vertices in the BOT graph, then there is a truth assignment that satisfies at least $(1 - \epsilon)$ fraction of the clauses in $\phi$, where $\epsilon$ is the constant defined in Theorem 5.

To see why Theorem 13 implies our main theorem, Theorem 4, consider that we start the reduction with two families of instances $\phi_Y$ and $\phi_N$ in MAX-3SAT($k$) which are NP-hard to distinguish. Then we construct their corresponding BOT hypergraphs, $H_{\phi_Y}$ and $H_{\phi_N}$, and use the edge sets of these two hypergraphs as training sets of size $m$ for a DPP maximum likelihood learning algorithm. The log likelihood estimator of $\phi_Y$ is $\ell^* = \Theta(\log N)$ by Theorem 10. What is the log likelihood estimator of $\phi_N$? Well, by Theorem 11, if the marginal kernel of $G_{\phi_N}$ has log likelihood $\ell(K) \leq \ell^* + \delta$ for some small enough $\delta > 0$, then there exists a marginal kernel $K'$ of dimension 3 such that $\ell(K') \leq \ell^* + C_k \delta^{1/4}$. By Theorem 13, we must have $\delta = \Omega(\frac{1}{\log^k N})$ for $\phi_N$. That is, the log likelihood estimator of $\phi_N$ is $\ell^* + \Omega(\frac{1}{\log^k N})$. Now if there were an polynomial-time algorithm $A$ which approximates the log likelihood estimator within a factor of $1 - 1/\Omega(\log^k N)$, then $A$ would be able to tell apart $\phi_Y$ from $\phi_N$ — thus solving an NP-complete problem — simply by approximating the maximum log likelihood estimators on training data sets from $H_{\phi_Y}$ and $H_{\phi_N}$, respectively.

**3. Discussion and open problems**

In this work, we establish that it is NP-hard to obtain a $1 - O(\frac{1}{\log^k N})$-approximation to the maximum log likelihood of DPPs. We also demonstrate a simple polynomial-time algorithm that achieves $\frac{1}{(1+\phi(1)) \log m}$-approximation. One immediate open problem is to close this large gap. A natural and plausible approach is to prove the cardinality-rank conjecture or at least to improve the bound in Theorem 11. Note that our hardness result does not rule out efficient learning with some constant factor of approximation: it is still possible that there is a polynomial-time algorithm that obtains a DPP kernel with a 99%-approximation to the maximum log likelihood. As observed earlier, we cannot preclude constant-factor approximations by a better analysis of the constructed hard instance: the approximation algorithm shows that our hardness result is tight up to a polynomial factor for the type of subset collections employed in the proof. Therefore any stronger hardness proof would require constructing a collection of subsets in which some element appears in a non-trivial fraction of the subsets.
Our investigation just takes a first stab at understanding the computational landscape of learning DPPs. In particular, our knowledge for the complexity of learning DPPs when the data set is indeed generated by an unknown DPP is still very limited: Can one design efficient algorithms for such a task? Can the DPP kernel be learned with arbitrary accuracy? And if not, what is the best approximation factor can such an algorithm achieve? Note that the data here is no longer a worst-case data set, but only sampled from a worst-case DPP. The underlying model is thus a semi-random one, and it seems challenging to extend NP-completeness hardness to such settings; some kind of average-case hardness is likely to be the best one can hope for. This is essentially the approach that Brunel et al. (2017a) had envisioned when examining the optimization landscape of the likelihood function for DPP kernels. The convergence of the empirical log-likelihood function to the true log-likelihood function only holds with high probability, and so in particular doesn’t carry over to the kinds of worst-case data sets produced by our reduction. Thus, their conjectured property may still hold, and may be a route to an efficient algorithm in this setting. On the other hand, “realizability” is probably a too strong assumption for practical purposes; DPPs are generally used to model processes featuring negative association, and it often seems implausible that the data actually follows a DPP distribution. Therefore, finding more appropriate assumptions is yet to be explored, and an algorithm for a realizable setting would be a natural first step along this direction.

As the other side of the coin, it is entirely conceivable that such efficient algorithms may not exist at all. One may view our main result as proving the hardness of “agnostic-learning” DPPs, while here the task would be proving hardness of “PAC-learning” DPPs. Presumably this is more difficult, as PAC-learning is in general easier than agnostic learning, it is thus harder to obtain lower bounds in the former setting. In particular, the usual approach of proving PAC-learning lower bounds involves uniform distributions over some prescribed collections of subsets. Such distributions are within the scope of PAC-learning model as it allows arbitrary distributions. By contrast, DPPs are normally unable to generate the uniform distribution over an arbitrary collection of subsets. Indeed, we believe that the data set we construct would be atypical for all DPPs. This is why contrary to the usual representation-specific hardness theorems in PAC-learning, we believe that an average-case hardness assumption will be necessary here.

Indeed, this discussion also highlights that maximum likelihood estimation is a kind of representation-specific (or “proper”) learning. One more potential route to learning DPPs is thus to use a more expressive representation that may possibly achieve better likelihood than the best DPP for a given data set, thwarting our reduction. Probabilistic Generating Circuits (Zhang et al., 2021), for example, can efficiently represent DPPs, and therefore may be an appropriate representation to use for this purpose.

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