Approximate Inference in Continuous Determinantal Point Processes

Raja Hafiz Affandi¹, Emily B. Fox², and Ben Taskar²

¹University of Pennsylvania, rajara@wharton.upenn.edu
²University of Washington, {ebfox@stat, taskar@cs}.washington.edu

Abstract

Determinantal point processes (DPPs) are random point processes well-suited for modeling repulsion. In machine learning, the focus of DPP-based models has been on diverse subset selection from a discrete and finite base set. This discrete setting admits an efficient sampling algorithm based on the eigendecomposition of the defining kernel matrix. Recently, there has been growing interest in using DPPs defined on continuous spaces. While the discrete-DPP sampler extends formally to the continuous case, computationally, the steps required are not tractable in general. In this paper, we present two efficient DPP sampling schemes that apply to a wide range of kernel functions: one based on low rank approximations via Nyström and random Fourier feature techniques and another based on Gibbs sampling. We demonstrate the utility of continuous DPPs in repulsive mixture modeling and synthesizing human poses spanning activity spaces.

1 Introduction

Samples from a determinantal point process (DPP) [15] are sets of points that tend to be spread out. More specifically, given $\Omega \subseteq \mathbb{R}^d$ and a positive semidefinite kernel function $L : \Omega \times \Omega \mapsto \mathbb{R}$, the probability density of a point configuration $A \subset \Omega$ under a DPP with kernel $L$ is given by

$$P_L(A) \propto \det(L_A),$$

where $L_A$ is the $|A| \times |A|$ matrix with entries $L(x, y)$ for each $x, y \in A$. The tendency for repulsion is captured by the determinant since it depends on the volume spanned by the selected points in the associated Hilbert space of $L$. Intuitively, points similar according to $L$ or points that are nearly linearly dependent are less likely to be selected.

Building on the foundational work in [5] for the case where $\Omega$ is discrete and finite, DPPs have been used in machine learning as a model for subset selection in which diverse sets are preferred [2, 3, 9, 12, 13]. These methods build on the tractability of sampling based on the algorithm of Hough et al. [10], which relies on the eigendecomposition of the kernel matrix to recursively sample points based on their projections onto the subspace spanned by the selected eigenvectors.

Repulsive point processes, like hard core processes [7, 16], many based on thinned Poisson processes and Gibbs/Markov distributions, have a long history in the spatial statistics community, where considering continuous $\Omega$ is key. Many naturally occurring phenomena exhibit diversity—trees tend to grow in the least occupied space [17], ant hill locations are over-dispersed relative to uniform placement [1] and the spatial distribution of nerve fibers is indicative of neuropathy, with hard-core processes providing a critical tool [25]. Repulsive processes on continuous spaces have garnered interest in machine learning as well, especially relating to generative mixture modeling [18, 29].

The computationally attractive properties of DPPs make them appealing to consider in these applications. On the surface, it seems that the eigendecomposition and projection algorithm of [10] for discrete DPPs would naturally extend to the continuous case. While this is true in a formal sense as $L$
becomes an operator instead of a matrix, the key steps such as the eigendecomposition of the kernel and projection of points on subspaces spanned by eigenfunctions are computationally infeasible except in a few very limited cases where approximations can be made [14]. The absence of a tractable DPP sampling algorithm for general kernels in continuous spaces has hindered progress in developing DPP-based models for repulsion.

In this paper, we propose an efficient algorithm to sample from DPPs in continuous spaces using low-rank approximations of the kernel function. We investigate two such schemes: Nyström and random Fourier features. Our approach utilizes a dual representation of the DPP, a technique that has proven useful in the discrete Ω setting as well [11]. For k-DPPs, which only place positive probability on sets of cardinality k [13], we also devise a Gibbs sampler that iteratively samples points in the k-set conditioned on all k − 1 other points. The derivation relies on representing the conditional DPPs using the Schur complement of the kernel. Our methods allow us to handle a broad range of typical kernels and continuous subspaces, provided certain simple integrals of the kernel function can be computed efficiently. Decomposing our kernel into quality and similarity terms as in [13], this includes, but is not limited to, all cases where the (i) spectral density of the quality and (ii) characteristic function of the similarity kernel can be computed efficiently. Our methods scale well with dimension, in particular with complexity growing linearly in d.

In Sec. 2 we review sampling algorithms for discrete DPPs and the challenges associated with sampling from continuous DPPs. We then propose continuous DPP sampling algorithms based on low-rank kernel approximations in Sec. 3 and Gibbs sampling in Sec. 4. An empirical analysis of the two schemes is provided in Sec. 5. Finally, we apply our methods to repulsive mixture modeling and human pose synthesis in Sec. 6 and 7.

2 Sampling from a DPP

When Ω is discrete with cardinality N, an efficient algorithm for sampling from a DPP is given in [10]. The algorithm, which is detailed in the supplement, uses an eigendecomposition of the kernel matrix \( L = \sum_{n=1}^{N} \lambda_n v_n v_n^T \) and recursively samples points \( x_i \) as follows, resulting in a set \( A \sim \text{DPP}(L) \) with \( A = \{x_i\} \):

Phase 1 Select eigenvector \( v_n \) with probability \( \frac{\lambda_n}{\lambda_n + 1} \). Let \( V \) be the selected eigenvectors (\( k = |V| \)).

Phase 2 For \( i = 1, \ldots, k \), sample points \( x_i \in \Omega \) sequentially with probability based on the projection of \( x_i \) onto the subspace spanned by \( V \). Once \( x_i \) is sampled, update \( V \) by excluding the subspace spanned by the projection of \( x_i \) onto \( V \).

When Ω is discrete, both steps are straightforward since the first phase involves eigendecomposing a kernel matrix and the second phase involves sampling from discrete probability distributions based on inner products between points and eigenvectors. Extending this algorithm to a continuous space was considered by [14], but for a very limited set of kernels \( L \) and spaces Ω. For general \( L \) and Ω, we face difficulties in both phases. Extending Phase 1 to a continuous space requires knowledge of the eigendecomposition of the kernel function. When Ω is a compact rectangle in \( \mathbb{R}^d \), [14] suggest approximating the eigendecomposition using an orthonormal Fourier basis.

Even if we are able to obtain the eigendecomposition of the kernel function (either directly or via approximations as considered in [14] and Sec. 3), we still need to implement Phase 2 of the sampling algorithm. Whereas the discrete case only requires sampling from a discrete probability function, here we have to sample from a probability density. When Ω is compact, [14] suggest using a rejection sampler with a uniform proposal on Ω. The authors note that the acceptance rate of this rejection sampler decreases with the number of points sampled, making the method inefficient in sampling large sets from a DPP. In most other cases, implementing Phase 2 even via rejection sampling is infeasible since the target density is in general non-standard with unknown normalization. Furthermore, a generic proposal distribution can yield extremely low acceptance rates.

In summary, current algorithms can sample approximately from a continuous DPP only for translation-invariant kernels defined on a compact space. In Sec. 3 we propose a sampling algorithm that allows us to sample approximately from DPPs for a wide range of kernels \( L \) and spaces Ω.

3 Sampling from a low-rank continuous DPP

Again considering Ω discrete with cardinality N, the sampling algorithm of Sec. 2 has complexity dominated by the eigendecomposition, \( O(N^3) \). If the kernel matrix \( L \) is low-rank, i.e. \( L = B^\top B \),...
with $B$ a $D \times N$ matrix and $D \ll N$.\cite{NS} showed that the complexity of sampling can be reduced to $O(ND^2 + D^3)$. The basic idea is to exploit the fact that $L$ and the dual kernel matrix $C = BB^\top$, which is $D \times D$, share the same nonzero eigenvalues, and for each eigenvector $v_k$ of $L$, $Bv_k$ is the corresponding eigenvector of $C$. See the supplement for algorithmic details.

While the dependence on $N$ in the dual is sharply reduced, in continuous spaces, $N$ is infinite. In order to extend the algorithm, we must find efficient ways to compute $C$ for Phase 1 and manipulate eigenfunctions implicitly for the projections in Phase 2. Generally, consider sampling from a DPP on a continuous space $\Omega$ with kernel $L(x, y) = \sum_{n=1}^{\infty} \lambda_n \phi_n(x)\overline{\phi_n(y)}$, where $\lambda_n$ and $\phi_n(x)$ are eigenvalues and eigenfunctions, and $\overline{\phi_n(y)}$ is the complex conjugate of $\phi_n(y)$. Assume that we can approximate $L$ by a low-dimensional (generally complex-valued) mapping, $B(x) : \Omega \to \mathbb{C}^D$:

$$\tilde{L}(x, y) = B(x)^*B(y), \text{ where } B(x) = [B_1(x), \ldots, B_D(x)]^\top. \quad (2)$$

Here, $A^*$ denotes complex conjugate transpose of $A$. We consider two efficient low-rank approximation schemes in Sec. 3.1 and 3.2. Using such a low-rank representation, we propose an analog of the dual sampling algorithm for continuous spaces, described in Algorithm 1. A similar algorithm provides samples from a $k$-DPP, which only gives positive probability to sets of a fixed cardinality $k$. The only change required is to the for-loop in Phase 1 to select exactly $k$ eigenvectors using an efficient $O(Dk)$ recursion. See the supplement for details.

**Algorithm 1 Dual sampler for a low-rank continuous DPP**

| Input: $L(x, y) = B(x)^*B(y)$, a rank-$D$ DPP kernel |
| Phase 1 |
| Compute $C = \int_{\Omega} B(x)B(x)^* dx$ |
| Compute eigendecomp. $C = \sum_{k=1}^{D} \lambda_k v_k v_k^\top$ |
| for $k = 1, \ldots, D$ do |
| $J \leftarrow J \cup \{k\}$ with probability $\frac{\lambda_k}{\lambda_k + 1}$ |
| $V \leftarrow \left\{ \frac{v_k}{\sqrt{\lambda_k C v_k}} \right\}_{k \in J}$ |

| Phase 2 |
| $X \leftarrow \emptyset$ |
| while $|V| > 0$ do |
| Sample $\hat{x}$ from $f(x) = \frac{1}{N} \sum_{v \in V} |v^* B(x)|^2$ |
| $X \leftarrow X \cup \{\hat{x}\}$ |
| Let $v_0$ be a vector in $V$ such that $v_0^* B(\hat{x}) \neq 0$ |
| Update $V \leftarrow \{v - \frac{v^* B(\hat{x})}{v_0^* B(\hat{x})} v_0 \mid v \in V - \{v_0\}\}$ |
| Orthonormalize $V$ w.r.t. $(v_1, v_2) = v_1^* C v_2$ |

| Output: $X$ |

In this dual view, we still have the same two-phase structure, and must address two key challenges:

**Phase 1** Assuming a low-rank kernel function decomposition as in Eq. (2), we need to able to compute the dual kernel matrix, given by an integral:

$$C = \int_{\Omega} B(x)B(x)^* dx. \quad (3)$$

**Phase 2** In general, sampling directly from the density $f(x)$ is difficult; instead, we can compute the cumulative distribution function (CDF) and sample $\hat{x}$ using the inverse CDF method\cite{Chiu}:

$$F(\hat{x} = (\hat{x}_1, \ldots, \hat{x}_d)) = \prod_{l=1}^{d} \int_{-\infty}^{\hat{x}_l} f(x) 1_{\{x_l \in \Omega\}} dx_l. \quad (4)$$

 Assuming (i) the kernel function $\hat{L}$ is finite-rank and (ii) the terms $C$ and $f(x)$ are computable, Algorithm\cite{NS} provides exact samples from a DPP with kernel $\hat{L}$. In what follows, approximations only arise from approximating general kernels $L$ with low-rank kernels $\hat{L}$. If given a finite-rank kernel $L$ to begin with, the sampling procedure is exact.

One could imagine approximating $L$ as in Eq. (2) by simply truncating the eigendecomposition (either directly or using numerical approximations). However, this simple approximation for known decompositions does not necessarily yield a tractable sampler, because the products of eigenfunctions required in Eq. (3) might not be efficiently integrable. For our approximation algorithm to work, not only do we need methods that approximate the kernel function well, but also that enable us to solve Eq. (3) and (4) directly for many different kernel functions. We consider two such approaches that enable an efficient sampler for a wide range of kernels: Nyström and random Fourier features.
3.1 Sampling from RFF-approximated DPP

Random Fourier features (RFF) \cite{le2018random} is an approach for approximating shift-invariant kernels, \( k(x, y) = k(x - y) \), using randomly selected frequencies. The frequencies are sampled independently from the Fourier transform of the kernel function, \( \omega_j \sim \mathcal{F}(k(x - y)) \), and letting:

\[
\hat{k}(x - y) = \frac{1}{D} \sum_{j=1}^{D} \exp\{i\omega_j^\top (x - y)\}, \quad x, y \in \Omega. \tag{5}
\]

To apply RFFs, we factor \( L \) into a quality function \( q \) and similarity kernel \( k \) (i.e., \( q(x) = \sqrt{L(x, x)} \)):

\[
L(x, y) = q(x)k(x, y)q(y), \quad x, y \in \Omega \text{ where } k(x, x) = 1. \tag{6}
\]

The RFF approximation can be applied to cases where the similarity function has a known characteristic function, e.g., Gaussian, Laplacian and Cauchy. Using Eq. (5), we can approximate the similarity kernel function to obtain a low-rank kernel and dual matrix:

\[
\tilde{L}_{RFF}(x, y) = \frac{1}{D} \sum_{j=1}^{D} q(x) \exp\{i\omega_j^\top (x - y)\} q(y), \quad C_{RFF}^{jk} = \frac{1}{D} \int_{\Omega} q^2(x) \exp\{i(\omega_j - \omega_k)^\top x\} dx.
\]

The CDF of the sampling distribution \( f(x) \) in Algorithm 1 is given by:

\[
F_{RFF}(\tilde{x}) = \frac{1}{|V|} \sum_{v \in V} \sum_{j=1}^{D} v_j \prod_{l=1}^{d} \int_{-\infty}^{\tilde{x}_l} q^2(x) \exp\{i(\omega_j - \omega_k)^\top x\} dx, \tag{7}
\]

where \( v_j \) denotes the \( j \)-th element of vector \( v \). Note that equations \( C_{RFF} \) and \( F_{RFF} \) can be computed for many different combinations of \( \Omega \) and \( q(x) \). In fact, this method works for any combination of (i) translation-invariant similarity kernel \( k \) with known characteristic function and (ii) quality function \( q \) with known spectral density. The resulting kernel \( L \) need not be translation invariant. In the supplement, we illustrate this method by considering a common and important example where \( \Omega = \mathbb{R}^d \), \( q(x) \) is Gaussian, and \( k(x, y) \) is any kernel with known Fourier transform.

3.2 Sampling from a Nyström-approximated DPP

Another approach to kernel approximation is the Nyström method \cite{williams2001using}. In particular, given \( z_1, \ldots, z_D \) landmarks sampled from \( \Omega \), we can approximate the kernel function and dual matrix as,

\[
\tilde{L}_{Nys}(x, y) = \sum_{j=1}^{D} \sum_{k=1}^{D} W_{jk} L(x, z_j) L(z_k, y), \quad C_{Nys}^{jk} = \sum_{n=1}^{D} \sum_{m=1}^{D} W_{jn}W_{mk} \int_{\Omega} L(z_n, x)L(x, z_m)dx,
\]

where \( W_{jk} = L(z_j, z_k)^{-1/2} \). Denoting \( w_j(v) = \sum_{n=1}^{D} W_{jn}v_n \), the CDF of \( f(x) \) in Alg. 1 is:

\[
F_{Nys}(\tilde{x}) = \frac{1}{|V|} \sum_{v \in V} \sum_{j=1}^{D} w_j(v)w_k(v) \prod_{l=1}^{d} \int_{-\infty}^{\tilde{x}_l} L(x, z_j)L(z_k, x)dx. \tag{8}
\]

As with the RFF case, we consider a decomposition \( L(x, y) = q(x)k(x, y)q(y) \). Here, there are no translation-invariant requirements, even for the similarity kernel \( k \). In the supplement, we provide the important example where \( \Omega = \mathbb{R}^d \) and both \( q(x) \) and \( k(x, y) \) are Gaussians and also when \( k(x, y) \) is polynomial, a case that cannot be handled by RFF since it is not translationally invariant.

4 Gibbs sampling

For \( k \)-DPPs, we can consider a Gibbs sampling scheme. In the supplement, we derive that the full conditional for the inclusion of point \( x_k \) given the inclusion of the \( k-1 \) other points is a 1-DPP with a modified kernel, which we know how to sample from. Let the kernel function be represented as before: \( L(x, y) = q(x)k(x, y)q(y) \). Denoting \( J^k = \{ x_j \}_{j \neq k} \) and \( M^{k} = L_{J^k}^{-1} \) the full conditional can be simplified using Schur’s determinantal equality \cite{gittovsky2015efficient}:

\[
p(x_k|\{ x_j \}_{j \neq k}) \propto L(x_k, x_k) - \sum_{i,j \neq k} M_{ij}^{k} L(x_i, x_k)L(x_j, x_k). \tag{9}
\]
Variational Distance

Eigenvalues

We also focus on sampling from $k$-DPPs for which the size of the set $X$ is always $k$. Details are in the supplement.

In general, sampling directly from this full conditional is difficult. However, for a wide range of kernel functions, including those which can be handled by the Nyström approximation in Sec. 3.2, the CDF can be computed analytically and $x_k$ can be sampled using the inverse CDF method:

$$F(x_i | \{x_j \}_{j \neq k}) = \frac{\int_{-\infty}^{x_i} L(x_i, x_l) - \sum_{i,j \neq k} M_{ij}^k L(x_i, x_j) L(x_j, x_l) 1_{\{x_l \in \Omega\}} \, dx_l}{\int_{\Omega} L(x, x) - \sum_{i,j \neq k} M_{ij}^k L(x_i, x) L(x_j, x) \, dx}$$  \hspace{1cm} (10)

In the supplement, we illustrate this method by considering the case where $\Omega = \mathbb{R}^d$ and $q(x)$ and $k(x, y)$ are Gaussians. We use this same Schur complement scheme for sampling from the full conditionals in the mixture model application of Sec. 6. A key advantage of this scheme for several types of kernels is that the complexity of sampling scales linearly with the number of dimensions $d$ making it suitable in handling high-dimensional spaces.

As with any Gibbs sampling scheme, the mixing rate is dependent on the correlations between variables. In cases where the kernel introduces low repulsion we expect the Gibbs sampler to mix well, while in a high repulsion setting the sampler can mix slowly due to the strong dependencies between points and fact that we are only doing one-point-at-a-time moves. We explore the dependence of convergence on repulsion strength in the supplementary materials. Regardless, this sampler provides a nice tool in the $k$-DPP setting. Asymptotically, theory suggests that we get exact (though correlated) samples from the $k$-DPP. To extend this approach to standard DPPs, we can first sample $k$ (this assumes knowledge of the eigenvalues of $L$) and then apply the above method to get a sample. This is fairly inefficient if many samples are needed. A more involved but potentially efficient approach is to consider a birth-death sampling scheme where the size of the set can grow/shrink by 1 at every step.

5 Empirical analysis

To evaluate the performance of the RFF and Nyström approximations, we compute the total variational distance $||P_L - P_{\hat{L}}|| = \frac{1}{2} \sum_{X} |P_L(X) - P_{\hat{L}}(X)|$, where $P_L(X)$ denotes the probability of set $X$ under a DPP with kernel $L$, as given by Eq. (1). We restrict our analysis to the case where the quality function and similarity kernel are Gaussians with isotropic covariances $\Gamma = \text{diag}(\sigma^2, \ldots, \sigma^2)$, respectively. We also focus on sampling from $k$-DPPs for which the size of the set $X$ is always $k$.

Fig. 1 displays estimates of the total variational distance for the RFF and Nyström approximations when $\rho^2 = 1$, varying $\sigma^2$ (the repulsion strength) and the dimension $d$. Note that the RFF method performs slightly worse as $\sigma^2$ increases and is rather invariant to $d$ while the Nyström method performs much better for increasing $\sigma^2$ but worse for increasing $d$.

While this phenomenon seems perplexing at first, a study of the eigenvalues of the Gaussian kernel across dimensions sheds light on the rationale (see Fig. 1). Note that for fixed $\sigma^2$ and $\rho^2$, the decay of eigenvalues is slower in higher dimensions. It has been previously demonstrated that the Nyström method performs favorably in kernel learning tasks compared to RFF in cases where there is a large eigengap in the kernel matrix [28]. The plot of the eigenvalues seems to indicate the same phenomenon here. Furthermore, this result is consistent with the comparison of RFF to Nyström in approximating DPPs in the discrete $\Omega$ case provided in [3].

This behavior can also be explained by looking at the theory behind these two approximations. For the RFF, while the kernel approximation is guaranteed to be an unbiased estimate of the true kernel element-wise, the variance is fairly high [19]. In our case, we note that the RFF estimates of minors are biased because of non-linearity in matrix entries, overestimating probabilities for point

![Figure 1: Estimates of total variational distance for Nyström and RFF approximation methods to a DPP with Gaussian quality and similarity with covariances $\Gamma = \text{diag}(\rho^2, \ldots, \rho^2)$ and $\Sigma = \text{diag}(\sigma^2, \ldots, \sigma^2)$, respectively. (a)-(c) For dimensions $d=1, 5$ and $10$, each plot considers $\rho^2 = 1$ and varies $\sigma^2$. (d) Eigenvalues for the Gaussian kernels with $\sigma^2 = \rho^2 = 1$ and varying dimension $d$.](image-url)
configurations that are more spread out, which leads to samples that are overly-dispersed. For the Nyström method, on the other hand, the quality of the approximation depends on how well the landmarks cover Ω. In our experiments the landmarks are sampled i.i.d. from q(x). When either the similarity bandwidth σ² is small or the dimension d is high, the effective distance between points increases, thereby decreasing the accuracy of the approximation. Theoretical bounds for the Nyström DPP approximation in the case when Ω is finite are provided in [3]. We believe the same result holds for continuous Ω by extending the eigenvalues and spectral norm of the kernel matrix to operator eigenvalues and operator norms, respectively.

In summary, for moderate values of σ² it is generally good to use the Nyström approximation for low-dimensional settings and RFF for high-dimensional settings.

6 Repulsive priors for mixture models

Mixture models are used in a wide range of applications from clustering to density estimation. A common issue with such models, especially in density estimation tasks, is the introduction of redundant, overlapping components that increase the complexity and reduce interpretability of the resulting model. This phenomenon is especially prominent when the number of samples is small. In a Bayesian setting, a common fix to this problem is to consider a sparse Dirichlet prior on the mixture weights, which penalizes the addition of non-zero-weight components. However, such approaches run the risk of inaccuracies in the parameter estimates [13]. Instead, [13] show that sampling the location parameters using repulsive priors leads to better separated clusters while maintaining the accuracy of the density estimate. They propose a class of repulsive priors that rely on explicitly defining a distance metric and the manner in which small distances are penalized. The resulting posterior computations can be fairly complex.

The theoretical properties of DPPs make them an appealing choice as a repulsive prior. In fact, [29] considered using DPPs as repulsive priors in latent variable models. However, in the absence of a feasible continuous DPP sampling algorithm, their method was restricted to performing MAP inference. Here we propose a fully generative probabilistic mixture model using a DPP prior for the location parameters, with a K-component model using a K-DPP.

In the common case of mixtures of Gaussians (MoG), our posterior computations can be performed using Gibbs sampling with nearly the same simplicity of the standard case where the location parameters μ_k are assumed to be i.i.d.. In particular, with the exception of updating the location parameters {μ_1, ..., μ_K}, our sampling steps are identical to standard MoG Gibbs updates in the uncollapsed setting. For the location parameters, instead of sampling each μ_k independently from its conditional posterior, our full conditional depends upon the other locations μ_i,k as well. Details are in the supplement, where we show that this full conditional has an interpretation as a single draw from a tilted 1-DPP. As such, we can employ the Gibbs sampling scheme of Sec. 4.

We assess the clustering and density estimation performance of the DPP-based model on both synthetic and real datasets. In each case, we run 10,000 Gibbs iterations, discard 5,000 as burn-in and thin the chain by 10. Hyperparameter settings are in the supplement. We randomly permute the labels in each iteration to ensure balanced label switching. Draws are post-processed following the algorithm of [23] to address the label switching issue.

Synthetic data To assess the role of the prior in a density estimation task, we generated a small sample of 100 observations from a mixture of two Gaussians. We consider two cases, the first with well-separated components and the second with poorly-separated components. We compare a mixture model with locations sampled i.i.d. (IID) to our DPP repulsive prior (DPP). In both cases, we set an upper bound of six mixture components. In Fig. 2, we see that both IID and DPP provide very similar density estimates. However, IID uses many large-mass components to describe the density. As a measure of simplicity of the resulting density description, we compute the average entropy of the posterior mixture membership distribution, which is a reasonable metric given the similarity of the overall densities. Lower entropy indicates a more concise representation in an information-theoretic sense. We also assess the accuracy of the density estimate by computing both (i) Hamming distance error relative to true cluster labels and (ii) held-out log-likelihood on 100 observations. The results are summarized in Table I. We see that DPP results in (i) significantly lower entropy, (ii) lower overall clustering error, and (iii) statistically indistinguishable held-out log-likelihood. These results signify that we have a sparser representation with well-separated (interpretable) clusters while maintaining the accuracy of the density estimate.
We consider another possible application of continuous-space sampling. In many applications of inverse reinforcement learning or inverse optimal control, the learner is presented with control trajectories executed by an expert and tries to estimate a reward function that would approximately reproduce such policies [1]. In order to estimate the reward function, the learner needs to compare the rewards of a large set of trajectories (or all, if possible), which becomes intractable in high-dimensional spaces with complex non-linear dynamics. A typical approximation is to use a set of perturbed expert trajectories as a comparison set, where a good set of trajectories should cover as large a part of the space as possible.

Table 1: For IID and DPP on synthetic datasets: mean (stdev) for mixture membership entropy, cluster assignment error rate and held-out log-likelihood of 100 observations under the posterior mean density estimate.

| DATASET          | ENTROPY  | CLUSTERING ERROR | HELDOUT LOG-LIKE |
|------------------|----------|------------------|------------------|
|                  | IID      | DPP              | IID              | DPP              | IID            | DPP          |
| Well-separated   | 1.11 (0.3) | 0.88 (0.2)        | 0.19 (0.1)       | 0.19 (0.1)       | -169 (6)      | -171 (8)|
| Poorly-separated | 1.46 (0.2) | 0.92 (0.3)        | 0.47 (0.1)       | 0.39 (0.1)       | -211 (10)     | -207 (9) |

Real data We also tested our DPP model on three real density estimation tasks considered in [20]: 82 measurements of velocity of galaxies diverging from our own (galaxy), acidity measurement of 155 lakes in Wisconsin (acidity), and the distribution of enzymatic activity in the blood of 245 individuals (enzyme). We once again judge the complexity of the density estimates using the posterior mixture membership entropy as a proxy. To assess the accuracy of the density estimates, we performed 5-fold cross validation to estimate the predictive held-out log-likelihood. As with the synthetic data, we find that DPP visually results in better separated clusters (Fig. 2). The DPP entropy measure is also significantly lower for data that are not well separated (acidity and galaxy) while the differences in predictive log-likelihood estimates are not statistically significant (Table 2).

Finally, we consider a classification task based on the iris dataset: 150 observations from three iris species with four length measurements. For this dataset, there has been significant debate on the optimal number of clusters. While there are three species in the data, it is known that two have very low separation. Based on loss minimization, [24][26] concluded that the optimal number of clusters was two. Table 2 compares the classification error using DPP and IID when we assume for evaluation the real data has three or two classes (by collapsing two low-separation classes), but consider a model with a maximum of six components. While both methods perform similarly for three classes, DPP has significantly lower classification error under the assumption of two classes, since DPP places large posterior mass on only two mixture components. This result hints at the possibility of using the DPP mixture model as a model selection method.

7 Generating diverse sample perturbations

We consider another possible application of continuous-space sampling. In many applications of inverse reinforcement learning or inverse optimal control, the learner is presented with control trajectories executed by an expert and tries to estimate a reward function that would approximately reproduce such policies [1]. In order to estimate the reward function, the learner needs to compare the rewards of a large set of trajectories (or all, if possible), which becomes intractable in high-dimensional spaces with complex non-linear dynamics. A typical approximation is to use a set of perturbed expert trajectories as a comparison set, where a good set of trajectories should cover as large a part of the space as possible.
Table 2: For IID and DPP, mean (stdev) of (left) mixture membership entropy and held-out log-likelihood for three density estimation tasks and (right) classification error under 2 vs. 2 of true classes for the iris data.

| DATA    | ENTROPY   | HELDOUT LL. |   | DATA    | CLASS ERROR |
|---------|-----------|--------------|---|---------|-------------|
|         | IID DPP   | IID DPP      |   |         | IID DPP      |
| Galaxy  | 0.89 (0.2)| 0.74 (0.2)   | -20(2) | Iris (3 cls) | 0.43 (0.02) |
| Acidity | 1.32 (0.1)| 0.98 (0.1)   | -49 (2) | Iris (2 cls) | 0.23 (0.03) |
| Enzyme  | 1.01 (0.1)| 0.96 (0.1)   | -55(2) |            | 0.15 (0.03) |

Figure 3: Left: Diverse set of human poses relative to an original pose by sampling from an RFF (top) and Nyström (bottom) approximations with kernel based on MoCap of the activity dance. Right: Fraction of data having a DPP/i.i.d. sample within an $\epsilon$ neighborhood.

We propose using DPPs to sample a large-coverage set of trajectories, in particular focusing on a human motion application where we assume a set of motion capture (MoCap) training data taken from the CMU database. Here, our dimension $d$ is 62, corresponding to a set of joint angle measurements. For a given activity, such as dancing, we aim to select a reference pose and synthesize a set of diverse, perturbed poses. To achieve this, we build a kernel with Gaussian quality and similarity using covariances estimated from the training data associated with the activity. The Gaussian quality is centered about the selected reference pose and we synthesize new poses by sampling from our continuous DPP using the low-rank approximation scheme. In Fig. 3, we show an example of such DPP-synthesized poses. For the activity dance, to quantitatively assess our performance in covering the activity space, we compute a coverage rate metric based on a random sample of 50 poses from a DPP. For each training MoCap frame, we compute whether the frame has a neighbor in the DPP sample within an $\epsilon$ neighborhood. We compare our coverage to that of i.i.d. sampling from a multivariate Gaussian chosen to have variance matching our DPP sample. Despite favoring the i.i.d. case by inflating the variance to match the diverse DPP sample, the DPP poses still provide better average coverage over 100 runs. See Fig. 3 (right) for an assessment of the coverage metric. A visualization of the samples is in the supplement. Note that the i.i.d. case requires on average $\epsilon = 253$ to cover all data whereas the DPP only requires $\epsilon = 82$. By $\epsilon = 40$, we cover over 90% of the data on average. Capturing the rare poses is extremely challenging with i.i.d. sampling, but the diversity encouraged by the DPP overcomes this issue.

8 Conclusion

Motivated by the recent successes of DPP-based subset modeling in finite-set applications and the growing interest in repulsive processes on continuous spaces, we considered methods by which continuous-DPP sampling can be straightforwardly and efficiently approximated for a wide range of kernels. Our low-rank approach harnessed approximations provided by Nyström and random Fourier feature methods and then utilized a continuous dual DPP representation. The resulting approximate sampler garners the same efficiencies that led to the success of the DPP in the discrete case. One can use this method as a proposal distribution and correct for the approximations via Metropolis-Hastings, for example. For $k$-DPPs, we devised an exact Gibbs sampler that utilized the Schur complement representation. Finally, we demonstrated that continuous-DPP sampling is useful both for repulsive mixture modeling (which utilizes the Gibbs sampling scheme) and in synthesizing diverse human poses (which we demonstrated with the low-rank approximation method). As we saw in the MoCap example, we can handle high-dimensional spaces $d$, with our computations scaling just linearly with $d$. We believe this work opens up opportunities to use DPPs as parts of many models.

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Supplementary Material: Approximate Inference in Continuous Determinantal Processes

Abstract

We provide further details for the NIPS 2013 submission “Approximate Inference in Continuous Determinantal Processes”. First, we elaborate upon the existing DPP samplers for the discrete and finite $\Omega$ case. We then provide a list of standard cases when our (approximate) DPP sampling scheme can be performed. We derive the low-rank approximation and Gibbs sampling schemes for a few standard cases along with the details of empirical analysis of the low-rank approximations. For our mixture of Gaussian example application, we detail the model specification and Gibbs sampler and contrast with a standard (non-repulsive) mixture model. Finally, we provide additional details on the settings used in our experiments and present some additional figures of results.

A DPP, k-DPP, and dual DPP sampling

For $\Omega$ discrete and finite with cardinality $N$, we provide the algorithms for sampling from DPPs, $k$-DPPs, and DPPs via the dual representation in Algorithms 2, 3, 4. In the $k$-DPP sampler, $e_i$ denotes the $i$th elementary symmetric polynomial. For $\Omega$ continuous, we provide the continuous $k$-DPP dual sampler in Algorithm 5. Note that the only difference relative to the DPP dual sampler is in the for loop of Phase 1. The revision exactly parallels the story for the discrete $\Omega$ case.

Algorithm 2 DPP-Sample(L)

| Input: kernel matrix $L$ of rank $D$ | PHASE 1 |
|-------------------------------------|---------|
| $\{(v_n, \lambda_n)\}_{n=1}^D$ ← eigendecomposition of $L$ | $J ← \emptyset$ |
| for $n = 1, \ldots, D$ do | $J ← J \cup \{n\}$ with prob. $\frac{\lambda_n}{\sum_{v \in V} (v^\top e_i)^2}$ |
| $V ← \{v_n\}_{n \in J}$ | Output: $Y$ |

Algorithm 3 $k$-DPP-Sample(L)

| Input: kernel matrix $L$ of rank $D$, size $k$ | PHASE 1 |
|-----------------------------------------------|---------|
| $\{(v_n, \lambda_n)\}_{n=1}^D$ ← eigendecomposition of $L$ | $J ← \emptyset$ |
| for $n = D, \ldots, 1$ do | $J ← J \cup \{n\}$ |
| if $u ∼ U[0, 1] < \frac{\lambda_n e_i^{n-1}}{\sum_{v \in V} (v^\top e_i)^2}$ then | $k ← k - 1$ |
| $J ← J \cup \{n\}$ | if $k = 0$ then |
| $k ← k - 1$ | break |
| $V ← \{v_n\}_{n \in J}$ | Output: $Y$ |

Algorithm 4 Dual DPP-Sample(L)

| Input: kernel matrix $L$ of rank $D$, size $k$ | PHASE 1 |
|-----------------------------------------------|---------|
| $\{(v_n, \lambda_n)\}_{n=1}^D$ ← eigendecomposition of $L$ | $J ← \emptyset$ |
| for $n = D, \ldots, 1$ do | $J ← J \cup \{n\}$ |
| if $u ∼ U[0, 1] < \frac{\lambda_n e_i^{n-1}}{\sum_{v \in V} (v^\top e_i)^2}$ then | $k ← k - 1$ |
| $J ← J \cup \{n\}$ | if $k = 0$ then |
| $k ← k - 1$ | break |
| $V ← \{v_n\}_{n \in J}$ | Output: $Y$ |
Algorithm 4 Dual-DPP-Sample(B)

Input: $B \in \mathbb{C}^{D \times N}$ such that $L = B^*B$.

PHASE 1
$C \leftarrow BB^*$
$\{(\mathbf{v}_n, \lambda_n)\}_{n=1}^D \leftarrow$ eigendecompositon of $C$
$J \leftarrow \emptyset$
for $n = 1, \ldots, D$
do
$J \leftarrow J \cup \{n\}$ with prob. $\frac{\lambda_n}{\sum_{k \in J} \lambda_k}$
$\tilde{V} \leftarrow \left\{ \frac{\mathbf{v}_n}{\sqrt{\sum_{n \in J} \lambda_n}} \right\}_{n \in J}$

PHASE 2
$Y \leftarrow \emptyset$
while $|V| > 0$
do
Select $i$ from $\Omega$ with $\Pr(i) = \frac{1}{|V|} \sum_{n \in V} \langle \hat{B}^*B_i \rangle^2$
$Y \leftarrow Y \cup \{i\}$
Let $\hat{v}_0$ be a vector in $V$ with $B_i^*\hat{v}_0 \neq 0$
Update $\tilde{V} \leftarrow \left\{ \hat{v} - \frac{\hat{v}^* B_i \hat{v}_0}{\langle \hat{v}, B_i^* \hat{v} \rangle} \mid \hat{v} \in \tilde{V} \right\}$
Orthonormalize $\tilde{V}$ w.r.t. $\langle \hat{v}_1, \hat{v}_2 \rangle = \hat{v}_1^*C\hat{v}_2$
Output: $Y$

Algorithm 5 Dual sampler for a low-rank continuous $k$-DPP

Input: $\tilde{L}(x, y) = B(x)^*B(y)$, a rank-$D$ DPP kernel

PHASE 1
Compute $C = \int_{\Omega} B(x)B(x)^*dx$
$\{(\mathbf{v}_n, \lambda_n)\}_{n=1}^D \leftarrow$ eigendecompositon of $C$
for $n = D, \ldots, 1$
do
if $u \sim U[0, 1] < \frac{\lambda_n}{\sum_{k \in J} \lambda_k}$ then
$J \leftarrow J \cup \{n\}$
$k \leftarrow k - 1$
if $k = 0$ then
break
$V \leftarrow \left\{ \frac{\mathbf{v}_n}{\sqrt{\sum_{n \in J} \lambda_n}} \right\}_{n \in J}$

PHASE 2
$X \leftarrow \emptyset$
while $|V| > 0$
do
Sample $\mathbf{x}$ from density $f(\mathbf{x}) = \frac{1}{|V|} \sum_{v \in V} |v^*B(\mathbf{x})|^2$
$X \leftarrow X \cup \{\mathbf{x}\}$
Let $\mathbf{v}_0$ be a vector in $V$ such that $\mathbf{v}_0^*B(\mathbf{x}) \neq 0$
Update $V \leftarrow \left\{ v - \frac{v^*B(\mathbf{x})}{\mathbf{v}_0^*B(\mathbf{x})} \mathbf{v}_0 \mid v \in V \right\}$
Orthonormalize $V$ w.r.t. $\langle \mathbf{v}_1, \mathbf{v}_2 \rangle = \mathbf{v}_1^*C\mathbf{v}_2$
Output: $X$

B Derivation of the Gibbs sampling scheme

For a $k$-DPP, the probability of choosing a specific $k$ point configuration is given by
$$p(\{x_j\}_{j=1}^k) \propto \det(L(\{x_j\}_{j=1}^k)).$$

Denoting $J^k = \{x_j\}_{j \neq k}$ and $M^k = L_{-1}^{-1}_{J^k}$, the Schur’s determinantal identity formula yields
$$\det(L(\{x_j\}_{j=1}^k)) = \det(L_{J^k}) \left( L(x_k, x_k) - \sum_{i, j \neq k} M^k_{ij} L(x_i, x_k) L(x_j, x_k) \right).$$

Conditioning on the inclusion of the other $k - 1$ points, and suppressing constants not dependent on $x_k$ we can now write the conditional distribution as
$$p(x_k | \{x_j\}_{j \neq k}) \propto L(x_k, x_k) - \sum_{i, j \neq k} M^k_{ij} L(x_i, x_k) L(x_j, x_k),$$

Normalizing and integrating this density yields a full conditional CDF given by
$$F(x_k | \{x_j\}_{j \neq k}) = \frac{\int_{-\infty}^{x_k} L(x_k, x_l) - \sum_{i, j \neq k} M^k_{ij} L(x_i, x_l) L(x_j, x_k) 1_{\{x_i \in \Omega\}} dx_l}{\int_{\Omega} L(x, x_k) - \sum_{i, j \neq k} M^k_{ij} L(x_i, x_k) L(x_j, x_k) dx_k}. $$

C Overview of analytically tractable kernel types under RFF or Nyström

Sampling from a DPP with kernel $L$ using Algorithm 1 of the main paper requires that (i) we can compute a low-rank decomposition $\hat{L}$ of $L$ and (ii) the terms $C$ and $f(x)$ are computable. In the main paper, we consider a decomposition of $L(x, y) = q(x)k(x, y)g(y)$ where $q(x)$ is a quality function and $k(x, y)$ a similarity kernel. We then use either
random Fourier features (RFF) or the Nyström method to approximate $L$ with $\tilde{L}$. In general, we can consider RFF approximations whenever the spectral density of $q(x)$ and characteristic function of $k(x, y)$ are known. For Nyström, the statement is not quite as clear. Instead, we provide a list of standard choices and their associated feasibilities for DPP sampling in Table 3. The list is by no means exhaustive, but is simply to provide some insight. We also elaborate upon some standard kernels in the following sections.

Table 3: Examination of the feasibility of DPP sampling using Nyström and RFF approximations for a few standard examples of quality functions $q$ and similarity kernels $k$.

| $q(x)$           | $k(x, y)$       | Method          |
|------------------|-----------------|-----------------|
| Gaussian, Laplacian | Gaussian, Laplacian | Nyström ✓     |
|                  | Cauchy          | RFF ✓                   |
|                  |                 | Gibbs ✓                |
| Gaussian, Laplacian | Cauchy          | Nyström ?                |
|                  |                 | RFF ?                  |
|                  |                 | Gibbs ?                |
| Cauchy           | Gaussian, Laplacian | Nyström ?                |
|                  |                 | RFF ?                  |
|                  |                 | Gibbs ?                |
| Cauchy           | Cauchy          | Nyström ?                |
|                  |                 | RFF ?                  |
|                  |                 | Gibbs ?                |
| Gaussian, Laplacian | Linear, Polynomial | Nyström ✓     |
|                  |                 | RFF X                  |
|                  |                 | Gibbs ✓                |

Example: Sampling from RFF-approximated DPP with Gaussian quality

Assuming $q(x) = \exp\left\{-\frac{1}{2}(x - a)^\top\Gamma^{-1}(x - a)\right\}$ and $k(x, y) = k(x - y)$ is given by a translation-invariant kernel with known characteristic function. We start by sampling $\omega_1, \ldots, \omega_D \sim \mathcal{F}(k(x - y))$. Note, for example, that the Fourier transform of a Gaussian kernel is a Gaussian while that of the Laplacian is Cauchy and vice versa. The approximated kernel is given by

$$\tilde{L}_{RFF} = q(x) \left[\frac{1}{D} \sum_{j=1}^{D} \exp i \omega_j^\top(x - y)\right] q(y) \quad \text{where} \quad q(x) = \exp\left\{-\frac{1}{2}(x - a)^\top\Gamma^{-1}(x - a)\right\}. \quad (15)$$

The elements of the dual matrix $C_{RFF}$ are then given by

$$C_{jk}^{RFF} = \frac{1}{D} \int_{\mathbb{R}^d} \exp\{- (x - a)^\top \Gamma^{-1} (x - a) + i (\omega_j - \omega_k)^\top x\} dx. \quad (16)$$

Letting $R \Delta R^\top$ be the spectral decomposition of $\Gamma^{-1}$ with $\Delta = \text{diag}(\frac{1}{\sigma_1^2}, \ldots, \frac{1}{\sigma_D^2})$, $\tilde{\omega}_j = R^\top \omega_j$, $\tilde{a} = R^\top a$ and $y = R^\top x$, one can straightforwardly derive:

$$C_{jk}^{RFF} = \frac{1}{D} \prod_{l=1}^{d} \left[ \sqrt{\pi \sigma_l^2} \exp \left\{ - \frac{\delta_l^2 (\tilde{\omega}_j - \tilde{\omega}_k)^2}{4} \right\} + i \tilde{a}_l (\tilde{\omega}_j - \tilde{\omega}_k) \right]. \quad (17)$$

Likewise,

$$F_{RFF}(y) = \frac{1}{D |V|} \sum_{\omega \in V} \sum_{j=1}^{D} \sum_{k=1}^{D} v^{(j)} v^{(k)} \prod_{l=1}^{d} g(\tilde{\omega}_j, \tilde{\omega}_k, \tilde{a}_l, \delta_l, y_l), \quad (18)$$

where

$$g(\tilde{\omega}_j, \tilde{\omega}_k, \tilde{a}_l, \delta_l, y_l) = \frac{1}{2} \sqrt{\pi \sigma_l^2} \exp \left\{ - \frac{\delta_l^2 (\tilde{\omega}_j - \tilde{\omega}_k)^2}{4} \right\} + i \tilde{a}_l \left( \tilde{\omega}_j - \tilde{\omega}_k \right) \left( 1 - \text{erf} \left( \frac{i \sqrt{\sigma_l^2} (\tilde{\omega}_j - \tilde{\omega}_k)}{2 \sqrt{\delta_l^2}} - \frac{y_l - \tilde{a}_l}{2 \sqrt{\delta_l^2}} \right) \right).$$

Once samples $y$ are obtained, we transform back into our original coordinate system by letting $x = Ry$. 

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Example: Sampling from Nyström-approximated DPP with Gaussian quality and similarity

Assuming \( q(x) = \exp \left\{ -\frac{1}{2}(x - a)^\top \Gamma^{-1}(x - a) \right\} \) and \( k(x, y) = \exp \left\{ -\frac{1}{2}(x - y)^\top \Sigma^{-1}(x - y) \right\} \), the approximated kernel is given by

\[
\tilde{K}_{Ny}(x, y) = \sum_{j=1}^{D} \sum_{k=1}^{D} W_{jk}^2 q(x)q(z_{jk}) \exp \left\{ -\frac{1}{2}(x - z_j)^\top \Sigma^{-1}(x - z_j) - \frac{1}{2}(y - z_k)^\top \Sigma^{-1}(y - z_k) \right\} q(z_k)q(y),
\]

(19)

Let \( \Sigma^{-1} = QAQ^\top \) with \( A = \text{diag}(\frac{1}{\sigma_1^2}, \ldots, \frac{1}{\sigma_n^2}) \), \( \Gamma^{-1} = R\Delta R^\top \) with \( \Delta = \text{diag}(\frac{1}{\sigma_1^2}, \ldots, \frac{1}{\sigma_n^2}) \) and \( (\Sigma^{-1} + \Gamma^{-1}) = T\Theta T^\top \) with \( \Theta = \text{diag}(\frac{1}{\sigma_1^2}, \ldots, \frac{1}{\sigma_n^2}) \). Furthermore, let \( \tilde{z}_j = T^\top(\Gamma^{-1} + \Sigma^{-1})\Sigma^{-1}z_j \), \( \tilde{a} = T^\top(\Gamma^{-1} + \Sigma^{-1})\Gamma^{-1}a \) and \( y = T^\top x \). Then, the elements of the dual matrix \( C_{Ny}^{Nys} \) are then given by

\[
C_{Ny,k}^{Nys} = \sum_{m=1}^{D} \sum_{n=1}^{D} W_{jm}W_{nk}A_{mn} \prod_{l=1}^{d} \sqrt{\pi\theta_l^2},
\]

(20)

where

\[
A_{mn} = \exp \left\{ -\frac{1}{2}(z_m - a)^\top \Gamma^{-1}(z_m - a) - \frac{1}{2}(z_m - a)^\top \Gamma^{-1}(z_m - a) - \frac{1}{2}z_m^\top \Sigma^{-1}z_m - \frac{1}{2}z_n^\top \Sigma^{-1}z_n 
+ (\Gamma^{-1}a + \Sigma^{-1}(z_m + z_n)/2)^\top (\Sigma^{-1} + \Gamma^{-1})^{-1}(\Gamma^{-1}a + \Sigma^{-1}(z_m + z_n)/2) - a^\top \Gamma^{-1}a \right\}.
\]

Finally, the CDF of \( f(y) \) is given by

\[
F_{Ny}(y) = \frac{1}{|V|} \sum_{v \in V} \sum_{j,k=1}^{D} w_j(v)w_k(v)A_{jk} \prod_{l=1}^{d} \frac{\sqrt{\pi}\theta_l^2}{2} \left[ 1 - \text{erf} \left( \frac{2\tilde{a}_l + \tilde{z}_j + \tilde{z}_k - 2y_l}{2\sqrt{\theta_l^2}} \right) \right].
\]

(21)

Once samples \( y \) are obtained, we transform back to our original coordinate system by letting \( x = Ty \).

Example: Sampling from Nyström-approximated DPP with Gaussian quality and polynomial similarity

For simplicity of exposition, we consider a linear similarity kernel and \( d = 1 \), although the result can straightforwardly be extended to higher order polynomials and dimensions \( d \). Assuming \( q(x) = \exp \left\{ -\frac{x^2}{2T^2} \right\} \) and \( k(x, y) = xy \), the approximated kernel is given by

\[
\tilde{K}_{Ny}(x, y) = \sum_{j=1}^{D} \sum_{k=1}^{D} W_{jk}^2 \exp \left\{ -\frac{(x^2 + z_j^2 + z_k^2 + y^2)}{2T^2} \right\} (xz_j)(yz_k).
\]

(22)

The elements of the dual matrix \( C_{Ny}^{Nys} \) are then given by

\[
C_{Ny,k}^{Nys} = \sum_{m=1}^{D} \sum_{n=1}^{D} W_{jm}W_{nk} \frac{z_mz_n}{2} \exp \left\{ -\frac{z_m^2 + z_n^2}{2T^2} \right\} \sqrt{\pi\rho^3}.
\]

(23)

The CDF is given by

\[
F_{Ny}(y) = \frac{1}{|V|} \sum_{v \in V} \sum_{j,k=1}^{D} w_j(v)w_k(v) \frac{z_jz_k}{2} \exp \left\{ -\frac{z_j^2 + z_k^2}{2T^2} \right\} \left[ \sqrt{\pi\rho^3} \left\{ \text{erf} \left( \frac{y}{\sqrt{T}} \right) + 1 \right\} - 2ye^{-\frac{y^2}{2T^2}} \right].
\]

(24)

Example: Gibbs sampling with Gaussian quality and similarity

For generic kernels \( L(x, y) = q(x)k(x, y)q(y) \), we recall that the CDF of \( x_k \) given \( \{x_j\}_{j \neq k} \) for a \( k \)-DPP is given by

\[
F(\hat{x}_k|\{x_j\}_{j \neq k}) = \frac{\int_{\mathbb{R}^d} q(x_k)^2 (1 - \sum_{i,j \neq k} M_{ij}q(x_i)q(x_j)k(x_k, x_i)k(x_j, x_k)) 1_{\{x_k \in \Omega\}} dx_k}{\int_{\Omega} q(x)^2 (1 - \sum_{i,j \neq k} M_{ij}q(x_i)q(x_j)k(x, x_i)k(x, x_j)) dx}.
\]

(25)
Assuming \( q(x) = \exp \left\{ -\frac{1}{2}(x-a)^\top \Gamma^{-1}(x-a) \right\} \) and \( k(x, y) = \exp \left\{ -\frac{1}{2}(x-y)^\top \Sigma^{-1}(x-y) \right\} \), the integrals above can be solved to yield

\[
F(\tilde{x}_k|\{x_j\}_{j \neq k}) = \prod_{i=1}^d \left[ \frac{\sqrt{\pi \delta_i^2}}{2} \left[ 1 - \text{erf} \left( \frac{2\tilde{a}_i - 2x_{ik}}{2\sqrt{\delta_i^2}} \right) \right] - \sum_{i,j \neq k} M_{ij} a_{ij} \sqrt{\pi \theta_i^2} \left[ 1 - \text{erf} \left( \frac{2\tilde{a}_i + \tilde{z}_i + \tilde{z}_{ij} - 2x_{ik}}{2\sqrt{\theta_i^2}} \right) \right] \right] \prod_{i=1}^d \left[ \frac{\sqrt{\pi \delta_i^2}}{2} - \sum_{i,j \neq k} W_{ij} a_{ij} \sqrt{\pi \theta_i^2} \right],
\]

(26)

where \( \tilde{a}, \tilde{z}, \delta_i, A_{ij}, \) and \( \theta_i \) are as given in the previous examples.

**D Details of the empirical analysis**

To evaluate the performance of the RFF and Nyström approximations, we compute the total variational distance

\[
\|P_L - P_\infty\|_1 = \frac{1}{2} \sum_X |P_L(X) - P_\infty(X)|,
\]

(27)

where \( P_L(X) \) denotes the probability of set \( X \) under a DPP with kernel \( L \), as given by Eq. (1). One can show that the normalized density is \( P_L(X) = \frac{\det(L_X)}{\prod_{i=1}^d \lambda_i(x_i)} \), which requires the eigenvalues of the kernel \( L \). Thus, we restrict our analysis to the case where the quality function and similarity kernel are Gaussians with isotropic covariances \( \Gamma = \text{diag}(\rho^2, \ldots, \rho^2) \) and \( \Sigma = \text{diag}(\sigma^2, \ldots, \sigma^2) \), respectively, since the eigenvalues of the kernel is easily computable in this setting [8]. In this case, letting \( n = (n_1, \ldots, n_d) \) with \( n_j \in \mathbb{Z}_+ \), the eigenvalues (indexed by multi-index \( n \)) are given by:

\[
\lambda_n = \prod_{j=1}^d \sqrt{\frac{\pi \rho^2}{2 \epsilon_2 + (\frac{1}{c_1} + 1)}} n_j^{-1} \quad c_1 = (\beta^2 + 1) \quad c_2 = \frac{\rho^2}{\sigma^2}.
\]

(28)

where \( \beta = (1 + 2\rho^2)^{\frac{1}{4}} \). Since the eigenvalues are known in closed form, we can estimate the total variation distance by sampling sets \( X \) from the approximated DPP and calculating the absolute difference between \( P_L(X) \) and \( P_\infty(X) \).

**E Empirical analysis of Gibbs sampling**

To assess the mixing rate of the Gibbs sampling scheme, we run the Gibbs sampler to sample points from a 1-dimensional 15-DPP with uniform quality and Gaussian similarity kernels in the space \( \Omega = [-\frac{1}{2}, \frac{1}{2}] \). We perform this sampling under two values of repulsion parameter, \( \sigma^2 = 0.01 \) (high repulsion) and \( \sigma^2 = 0.001 \) (low repulsion). We run 100 Gibbs chains, each of length 3000, discard the first 1500 samples as burn-in and thin every 15 iterations which we call cycles. Each cycle represents a full resampling of the set, having cycled through the past 15 points. We compare the results to i.i.d. sampling of Nyström-approximated DPP as a baseline.

Figure 4(a)-(b) shows a visualization of the 15 points of the 15-DPPs. Figure 4(c)-(d) shows the plots of the Nyström-approximated DPP samples. As an ordered set, we see qualitatively that the locations of the points are highly correlated from cycle to cycle in the high repulsion Gibbs samples while less correlation is observed in the low-repulsion counterpart. In the Nyström approximated case, there are no correlations as the samples are generated i.i.d..

Quantitatively, we use two measures as a proxy to the mixing rate: the average movement of point from cycle to cycle and the effective sample size. The average movement, \( m \), is simply defined as the average difference in distance between points from one cycle to another averaged over the cycles:

\[
m = \frac{1}{T-1} \frac{1}{k} \sum_{t=1}^{T-1} \sum_{i=1}^k (x_{i,t+1} - x_{i,t})^2,
\]

(29)

where \( T \) is the length of the chain after burn-in and thinning, \( k \) is the number of points and \( x_{i,t} \) is the coordinate of point \( x_i \) at cycle \( t \). In our experiment, \( T \) and \( k \) are 100 and 15, respectively. When the Gibbs chain is mixing well, we expect the average movement to be high as this signals that the points are less correlated across cycles.

The effective sample size is a standard measure in assessing the mixing of a Gibbs chain. To compute this, we first compute the lag-\( s \) autocorrelation function of each point in the sampled sets. We then average the autocorrelation...
function at lag-$s$ across the $k$ points and denote this quantity $\bar{\rho}_s$. The effective sample size is then given by: $\alpha T$, where

$$\alpha = \frac{1}{1 + 2 \sum_{s=1}^{2^\delta + 1} \bar{\rho}_s},$$

(30)

where $\delta$ is the smallest positive integer satisfying $\bar{\rho}_{2^\delta} + \bar{\rho}_{2^\delta + 1} > 0$. In the case of i.i.d. samples, we expect $\alpha$ to be close to 1 while in cases where the mixing is bad, $\alpha$ will be much lower.

Table 4 shows the average values of $m$ and $\alpha$ for our Gibbs samples with i.i.d. Nyström-approximated DPP samples serving as a benchmark. We see that in the low repulsion setting, the Gibbs chain mixes well with values close to the benchmarks while for the Gibbs sampler in the high repulsion setting, the values of $m$ and $\alpha$ are much lower, indicating slow mixing.

|         | Gibbs High Repulsion | Gibbs Low Repulsion | Nyström High Repulsion | Nyström Low Repulsion |
|---------|----------------------|---------------------|------------------------|------------------------|
| $m$     | 0.08 (0.07,0.08)     | 0.1 (0.10,0.11)     | 0.11 (0.10,0.11)       | 0.11 (0.11,0.12)       |
| $\alpha$| 0.39 (0.31,0.45)     | 0.92 (0.80,1)       | 0.98 (0.82, 1)         | 0.98 (0.90, 1)         |

Table 4: The mean and 95% confidence interval for average movement, $m$ and the effective sample size coefficient, $\alpha$ for Gibbs samples and i.i.d. Nyström samples in high and low repulsion settings.
Algorithm 6 Mixture of Gaussians sampler

**Input:** Previous mixture weights $\pi$, emission parameters $\{\mu_k, \sigma_k^2\}^2$.

for $i = 1, \ldots, N$ do

Sample cluster indicators $z_i \mid y_i, \{\mu_k, \sigma_k^2\}, \pi_k \propto \frac{1}{c_n} \sum_{k=1}^K \pi_k N(y_i; \mu_k, \sigma_k^2) \delta(z_i, k)$

Sample mixture weights $\pi \mid \{z_i\}, \alpha \sim \text{Dir}(\alpha + N_1, \ldots, \alpha + N_K)$

for $k = 1, \ldots, K$ do

Sample scale parameters $\sigma_k^2 \mid \{y_i : z_i = k\}, \mu_k, a_\sigma, b_\sigma \sim \text{IG}(a_\sigma, b_\sigma), \ k = 1, \ldots, K$

Sample location parameters $\{\mu_1, \ldots, \mu_K\} \mid \{y_i\}, \{z_i\}, \{\sigma_k^2\} \sim F_{\text{post}}$

**Output:** New mixture weights $\pi$, emission parameters $\{\mu_k, \sigma_k^2\}$.

F  Gibbs sampling for repulsive mixtures of Gaussians

**Generative Model** We consider a Bayesian mixture of Gaussians with either an independent normal (IID) or $K$-DPP (DPP) prior on the location parameters. In both cases, the $K$-component model with $N$ observations is specified as:

$$
\pi \mid \alpha \sim \text{Dir}(\alpha, \ldots, \alpha) \\
\sigma_k^2 \mid a_\sigma, b_\sigma \sim \text{IG}(a_\sigma, b_\sigma), \ k = 1, \ldots, K \\
\{\mu_1, \ldots, \mu_K\} \sim F \\
z_i \mid \pi \sim \pi, \ i = 1, \ldots, N
$$

$$
y_i \mid \pi, \{\mu_k, \sigma_k^2\} \sim N(\mu_{z_i}, \sigma_{z_i}^2), \ i = 1, \ldots, N.
$$

(31)

Here, IG denotes the inverse gamma distribution and Dir a $K$-dimensional Dirichlet. For simplicity, we consider the univariate case here, though the multivariate case follows directly by considering an inverse Wishart prior in place of the inverse gamma and likewise modifying $F$ accordingly. Such a multivariate case is examined in the iris classification example in the main paper.

The difference between the models is in how the location parameters are specified. For the IID case, we simply have:

$$
\mu_k \mid \mu_0, \sigma_0^2 \sim N(\mu_0, \sigma_0^2)
$$

(32)

For the DPP case, we jointly sample:

$$
\{\mu_1, \ldots, \mu_K\} \mid L \sim K\text{-DPP}(L).
$$

(33)

We consider $L$ decomposed into Gaussian quality and similarity terms:

$$
L(\mu_m, \mu_n) = q(\mu_m)k(\mu_m, \mu_n)q(\mu_n)
$$

(34)

with

$$
k(\mu_m, \mu_n) = \exp\left\{-\frac{(\mu_m - \mu_n)^2}{\gamma_0^2}\right\}, \ q(\mu_m) = N(\mu_0, 2\sigma_0^2).
$$

(35)

**Gibbs sampling** For the uncollapsed setting, where mixture weights $\pi$ and emission parameters $\{\mu_k, \sigma_k^2\}$ are sampled, Algorithm 6 summarizes the Gibbs sampler for the finite mixture of Gaussians. We write the algorithm generically so that the overlap between IID and DPP is clear. In particular, the locations are sampled from $F_{\text{post}}$, which generically refers to the full conditional of the cluster means. For the IID case, we sample i.i.d. for each $k$ from

$$
\mu_k \mid \{y_i : z_i = k\}, \sigma_k^2, \mu_0, \sigma_0^2 \sim N\left(\hat{\mu}_k, \hat{\sigma}_k^2\right),
$$

(36)
where \( \hat{\mu}_k = \left( \frac{1}{\sigma^2} + \frac{N_k}{\sigma^2} \right)^{-1} \left( \frac{\mu_0}{\sigma^2} + \frac{1}{\sigma^2} \sum_{i: z_i = k} y_i \right) \) and \( \sigma_k^2 = \left( \frac{1}{\sigma^2} + \frac{N_k}{\sigma^2} \right)^{-1} \). Here, \( N_k = |\{y_i : z_i = k\}| \), i.e., the cardinality of the set of observations assigned to cluster \( k \).

For DPP, note that \( p(\{\mu_j\}_{j=1}^k \mid \{y_i\}, \{z_i\}, \{\mu_k, \sigma_k^2\}) \propto \det(L_{\mu_1,\ldots,\mu_k}) \prod_{i=1}^k \prod_{i: z_i = j} N(y_i; \mu_j, \sigma_j^2) \). Unfortunately, this posterior distribution is not a \( k \)-DPP. However, fixing the rest of \( k - 1 \) centroids, the full conditional of \( \mu_k \) is (dropping constant terms that do not depend on \( \mu_k \))

\[
p(\mu_k \mid \{y_i\}, \{z_i\}, \{\mu_j, \sigma_j^2\}_{j \neq k}, \sigma_k^2) \propto \det(L_{\mu_1,\ldots,\mu_k}) \prod_{i: z_i = k} N(y_i; \mu_k, \sigma_k^2). \tag{37}
\]

As before, we can use Schur’s determinantal equality \([22]\) to get

\[
det(L_{\mu_1,\ldots,\mu_k}) \propto L(\mu_k, \mu_k) - \sum_{i,j \neq k} M_{ij}^k L(\mu_i, \mu_k) L(\mu_j, \mu_k) \tag{38}
\]

\[
= q^2(\mu_k) \left( 1 - \sum_{i,j \neq k} M_{ij}^k q(\mu_i) k(\mu_i, \mu_k) k(\mu_j, \mu_k) q(\mu_j) \right). \tag{39}
\]

Combining the previous two equations, we get the full conditional

\[
p(\mu_k \mid \{y_i\}, \{z_i\}, \{\mu_j, \sigma_j^2\}_{j \neq k}, \sigma_k^2) \propto q^2(\mu_k) \left( 1 - \sum_{i,j \neq k} M_{ij}^k q(\mu_i) k(\mu_i, \mu_k) k(\mu_j, \mu_k) q(\mu_j) \right) \prod_{i: z_i = k} N(y_i; \mu_k, \sigma_k^2). \tag{40}
\]

The CDF of the distribution above can be computed easily, since it only involves exponential quadratic forms. The inverse CDF method can then be used to obtain a sample from the above distribution. Note once again that \( q^2(\mu_k) \prod_{i: z_i = k} N(y_i; \mu_k, \sigma_k^2) \) is defined to be exactly the same as the Gaussian distribution where \( \mu_k \) would have been sampled from in the IID case. Thus the equation above gives a nice intuition on the conditional density of \( \mu_k \) in the DPP setting: it is an exponentially tilted distribution in which \( q^2(\mu_k) \prod_{i: z_i = k} N(y_i; \mu_k, \sigma_k^2) \) is corrected by a factor that depends on the location of the other centroids. In the case where all of the other centroids are far away from the cluster center \( \mu_k \), the correction factor is close to one and we would recover the density for the IID case.

To get a sense of why the DPP leads to more diverse cluster centers than IID, consider the full conditional for \( \mu_k \) at some iteration \( m \) of our sampler, as visualized in Fig. 5. We have some data points currently assigned to cluster \( k \) via cluster indicators \( z_i = k \). The IID model assumes that \( \mu_k \) is independent of the other \( \mu_j \)’s whereas the DPP conditions on the other cluster centers leading to a conditional distribution for \( \mu_k \) that puts more mass on uncovered regions. In subsequent iterations, the data that had been assigned to cluster \( k \) but are not well covered by the sampled (and repulsed) \( \mu_k \) will instead be assigned to one of the existing cluster centers that have mass near that data item. Such an alternative cluster exists, and is why \( \mu_k \) was repulsed from that region, or will likely exist in future draws.

One attractive aspect of our DPP formulation is the fact that the sampling strategy maintains nearly the same simplicity as the standard IID sampler. This is in contrast, for example, to the repulsive mixture formulation of [18] which relied on slice sampling and draws from truncated normals, where the truncating region could only be computed in closed form for a restricted set of repulsive functions.

\section{Additional details on experiments}

\subsection{Hyperparameter settings}

For our mixture of Gaussian experiments, we used an inverse Wishart \( \text{IW}(\nu, \Psi) \) with \( \nu = d + 1 \) and \( \Psi = I \), which corresponds to \( a_\sigma = 2 \) and \( b_\sigma = 1 \) for the inverse Gamma in 1-dimensional. Here, we use an inverse Wishart specification such that \( \Sigma \sim \text{IW}(\nu, \Psi) \) has mean \( \mathbb{E}[\Sigma] = \frac{\Psi}{\nu - d - 1} \). The Dirichlet hyperparameters were set to \( \alpha = \frac{1}{2} \), just as in [18]. For the location hyperparameters, in the IID case we set \( \mu_0 = 0 \) and \( \sigma_0^2 = 1 \). In the DPP case, we use \( \mu_0 \) and \( \sigma_0^2 \) as in the IID case and set the repulsion parameter \( \rho_0^2 = 1 \).

For the MoCap experiment, we computed the covariance estimate from the training data, and set the similarity covariance parameter \( \Sigma \) equal to this estimate. We then take the quality covariance parameter to be \( \Gamma = \frac{1}{2} \Sigma \).
Figure 6: Comparison between the full conditional for $\mu_k$ using the IID and DPP models at a given iteration $m$ of the sampler.

Figure 7: (a)-(c) DPP (blue) and i.i.d. multivariate Gaussian (red) samples projected onto the top 4 principal components of the dance data.

G.2 Additional figures for MoCap experiments

In Fig. 7 we provide a visualization of poses sampled from the DPP relative to i.i.d. sampling of poses from a multivariate Gaussian. From these plots, we see how the sample of poses from the DPP covers a broader space, even when the covariance of the multivariate Gaussian is inflated to match that of the DPP. The reason for this broader coverage is the fact that the under the DPP, sampled poses repulse from regions already covered by other sampled poses.

Fig. 8 displays additional human poses that are drawn i.i.d. from a multivariate Gaussian, and compares to our DPP draws from both the RFF and Nyström approximations.
Figure 8: Synthesizing perturbed human poses relative to an original pose by sampling (1) i.i.d. from a multivariate Gaussian versus (2) drawing a set from an RFF- or Nyström- approximated DPP with kernel based on MoCap data from the activity category. The Gaussian covariance is likewise formed from the activity data.