Determinantal Point Process Mixtures Via Spectral Density Approach

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Abstract. We consider mixture models where location parameters are a priori encouraged to be well separated. We explore a class of determinantal point process (DPP) mixture models, which provide the desired notion of separation or repulsion. Instead of using the rather restrictive case where analytical results are partially available, we adopt a spectral representation from which approximations to the DPP density functions can be readily computed. For the sake of concreteness the presentation focuses on a power exponential spectral density, but the proposed approach is in fact quite general. We later extend our model to incorporate covariate information in the likelihood and also in the assignment to mixture components, yielding a trade-off between repulsiveness of locations in the mixtures and attraction among subjects with similar covariates. We develop full Bayesian inference, and explore model properties and posterior behavior using several simulation scenarios and data illustrations. Supplementary materials for this article are available online (Bianchini et al., 2019).

Keywords: density estimation, nonparametric regression, repulsive mixtures, reversible jumps.

1 Introduction

1.1 Mixture models

Mixture models are an extremely popular class of models, that have been successfully used in many applications. For a review, see, e.g. Frühwirth-Schnatter (2006). Such models are typically stated as

\[ y_i \mid k, \theta, \pi \sim \text{iid} \sum_{j=1}^{k} \pi_j f(y_i \mid \theta_j), \quad i = 1, \ldots, n, \]  

where \( \pi = (\pi_1, \ldots, \pi_k) \) are constrained to be nonnegative and sum up to 1, \( \theta = (\theta_1, \ldots, \theta_k) \), and \( 1 \leq k \leq \infty \), with \( k = \infty \) corresponding to a nonparametric model. A common prior assumption is that \( \pi \sim \text{Dirichlet}(\delta_1, \ldots, \delta_k) \) and that the components of \( \theta \) are drawn i.i.d. from some suitable prior \( p_0 \). However, the weights \( \pi \) may be constructed differently, e.g. using a stick-breaking representation (finite or infinite), which poses a well-known connection with more general models, including nonparametric ones. See, e.g., Ishwaran and James (2001) and Miller and Harrison (2017). A popular class

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of Bayesian nonparametric models is the Dirichlet process mixture (DPM) model, introduced in Ferguson (1983) and Lo (1984). It is well-known that this class of mixtures usually overestimates the number of clusters, mainly because of “the rich gets richer” property of the Dirichlet process. By this we mean that both prior and posterior distributions are concentrated on a relatively large number of clusters, but a few are very large while the rest have very small sizes. Mixture models may even be inconsistent; see Rousseau and Mengersen (2011), where concerns about over-fitted mixtures are illustrated, and Miller and Harrison (2013), for inconsistency features of DPMs.

Despite their success, models like (1) tend to use excessively many mixture components. Xu et al. (2016) point out that this is due to the fact that the component-specific parameters are a priori i.i.d., and therefore, free to move. This motivated Petralia et al. (2012), Fúquene et al. (2016) and Quinlan et al. (2017) to explicitly define joint distributions for \( \theta \) having the property of repulsion among its components, i.e. \( p(\theta_1, \ldots, \theta_k) \) puts higher mass on configurations with well separated components. See also Quinlan et al. (2018) for an application to density estimation. For a different approach, via sparsity in the prior, see Malsiner-Walli et al. (2016).

Xu et al. (2016) explored a similar way to accomplish separation of mixture components, by means of a Determinantal Point Process (DPP) acting on the parameter space. DPPs have recently received increased attention in the statistical literature (Lavancier et al., 2015). DPPs are point processes having a product density function expressed as the determinant of a certain matrix constructed using a covariance function, evaluated at the pairwise distances among points, in such a way that higher mass is assigned to configurations of well-separated points. We give details below. DPPs have been used to make inference mostly on spatial data. Bardenet and Titsias (2015) and Affandi et al. (2014) applied DPPs to model spatial patterns of nerve fibers in diabetic patients, a basic motivation being that such fibers become more clustered as diabetes progresses. The latter discussed also applications to image search, showing how such processes could be used to study human perception of diversity in different image categories. Similarly, Kulesza and Taskar (2012) show how DPPs can be applied to various problems that are relevant to the machine learning community, such as finding diverse sets of high-quality search results, building informative summaries by selecting diverse sentences from documents, modeling non-overlapping human poses in images or videos, and automatically building timelines of important news stories. More recently, Shirota and Gelfand (2017) have described an approximate Bayesian computation method to fit DPPs to spatial point pattern data. Historically, the first paper where DPPs were adopted as a prior for statistical inference in mixture models is Affandi et al. (2013). The statistical literature also includes a number of papers illustrating theoretical properties for estimators of DPPs from a non-Bayesian viewpoint; see, for instance, Biscio and Lavancier (2016, 2017) and Bardenet and Titsias (2015).

1.2 Main contributions of this work

We discuss full Bayesian inference for a class of mixture densities where the locations follow stationary DPPs. Our first contribution is the introduction of an approach that generalizes and extends the model studied in Xu et al. (2016) who base their analysis
on a special, finite state space case of DPPs called L-ensembles, and with a particular closed-form expression for the determinant, in terms of a Gaussian covariance function. Instead, we resort to the spectral representation of the covariance function defining the determinant as the joint distribution of component-specific parameters. Our methods can thus be used with any such valid spectral representation, as described by Lavancier et al. (2015), which implies great generality of the proposal, that goes well beyond the Gaussian covariance function with a squared exponential kernel case of Xu et al. (2016). The extensions considered here are stated in the context of both uni- and multi-dimensional responses, and with different spectral densities. For brevity of presentation, we defer details of computational implementation to Section 1 of the on-line Supplementary Material file. For the sake of concreteness, our illustrations focus on the case of power exponential spectral representation; see examples with different spectral densities in Section 2 of the on-line Supplementary Material. This particular specification allows for flexible repulsion patterns, and we discuss how to set up different types of prior behavior, shedding light on the practical use of our approach in that particular scenario. Although we limit ourselves to the case of isotropic DPPs, inhomogeneous DPPs can be obtained by transforming or thinning a stationary process (Lavancier et al., 2015). A crucial point in our models and algorithms is the DPP density expression with respect to the unit rate Poisson process. This density is only defined for DPPs restricted to compact subsets $S$ of the state space, and if it exists, it explicitly depends on $S$. A sufficient condition for the existence of the density is that all the eigenvalues of (Mercer’s Theorem representation of) covariance function, restricted to $S$, are smaller than 1. We follow the spectral approach and assume that the covariance function defining the DPP has a spectral representation. One basic motivation for our choice is that conditions for the existence of a density become easier to check. Another motivation is the flexibility in assuming different degrees of repulsion in the DPP by way of the spectral density. We review here the basic theory on DPPs, making an effort to be as clear and concise as possible in the presentation of our subsequent models. We discuss applications in the context of synthetic and real data applications.

A second contribution of this work is the extension of the proposed spectral DPP model to incorporate covariate information in the likelihood and also in the assignment to mixture components. In particular, subjects with similar covariates are a priori more likely to co-cluster, just as in mixtures of experts models (see, e.g., McLachlan and Peel, 2005), where weights are defined as normalized exponential functions. From a computational viewpoint, a third contribution of our work is the generalization of the reversible jump (RJ) MCMC posterior simulation scheme proposed by Xu et al. (2016) to the general spectral approach and also to the covariate-dependent extensions we consider. We discuss two RJ MCMC versions, one for each of uni- and multi-variate responses. In all cases the algorithms require computing the DPP density with respect to the unit rate Poisson process. We explain how to carry out the calculations, and discuss the required restriction of the process to compact subsets. When extending the model to incorporate covariate information in both likelihood and prior assignment to mixture components, we discuss how to modify the RJ MCMC algorithms; see Section 3 of the on-line Supplementary Material.

Although not our primary target, a sub-product of the approach is the estimation of a partition in the sample. Specifically, we consider the partition minimizing the posterior
expectation of Binder’s loss function (Binder, 1978) under equal misclassification costs, a common choice in the applied Bayesian nonparametric literature (Lau and Green, 2007). Nevertheless, we emphasize one conceptual advantage of the separation induced by the prior assumption, namely, the reduction in the estimated number of clusters a posteriori compared to the usual mixture models without this feature. The fact that our prior generates components from a repulsive process, automatically penalizes redundancy, thus inducing fewer clusters than if they were allowed to be independently selected. Reducing the effective number of clusters a posteriori helps scaling up our model – better than alternatives with no separation – when the sample size grows. We discuss this particular point in our data illustrations.

We also provide comparison with alternative models, via predictive goodness-of-fit tools. We specifically consider some “natural competitors”, namely DPMs and the repulsive mixtures such as those introduced in Petralia et al. (2012) and Quinlan et al. (2017), as well as finite mixture models, as implemented in the package \texttt{Mclust} (Fraley et al., 2012). We find that our model is a competent alternative to those previously listed.

1.3 Outline

The rest of this article is organized as follows. Section 2 presents notation and theoretical background necessary to understand how DPP mixture models are constructed; we also sketch alternative repulsive models already introduced in the literature, later to be used as a comparison in the context of specific illustrative examples. Section 3 illustrates the covariate-dependent extension. Here we build on regular mixture models, incorporating covariate dependence in the mixture weights and optionally in the likelihood, which still allows for repulsion among components after correcting for the regression effect. Section 4 presents results from a simulation study and for unidimensional and bidimensional reference datasets, and an application to Biopic data is discussed in Section 5. We conclude in Section 6 with final comments and discussion. The on-line Supplementary Material contains a description of the two RJ MCMC algorithms, additional illustrative examples based on the well-known Galaxy dataset, supplemental figures, and additional application to Air Quality Index data.

2 Using DPPs to induce repulsion

We review here the basic theory on DPPs to the extent required to explain our mixture model. We use the same notation as in Lavancier et al. (2015), where further details on this theory may be found.

2.1 Basic theory on DPPs

Let $B \subseteq \mathbb{R}^d$; we mainly consider the cases $B = \mathbb{R}^d$ and $B = S$, a compact subset in $\mathbb{R}^d$. Let $X$ be a simple locally finite spatial point process defined on $B$, i.e. the number of points of the process in any bounded region is a finite random variable, and
there is at most one point at any location. See Daley and Vere-Jones (2003; 2007) for a general presentation on point processes. The class of DPPs we consider is defined in terms of their moments, expressed by their product density functions \( \rho^{(n)} : B^n \to [0, +\infty) \), \( n = 1, 2, \ldots \). Intuitively, for any pairwise distinct points \( x_1, \ldots, x_n \in B \), \( \rho^{(n)}(x_1, \ldots, x_n)dx_1 \cdots dx_n \) is the probability that \( X \) has a point in an infinitesimal small region around \( x_i \) of volume \( dx_i \), for each \( i = 1, \ldots, n \). More formally, \( X \) has \( n \)-th order product density function \( \rho^{(n)} : B^n \to [0, +\infty) \) if this function is locally integrable (i.e. \( \int_S |\rho^{(n)}(x)|dx < +\infty \) for any compact \( S \)) and, for any Borel-measurable function \( h : B^n \to [0, +\infty) \),

\[
E\left(\sum_{x_1, \ldots, x_n \in X} h(x_1, \ldots, x_n)\right) = \int_{B^n} \rho^{(n)}(x_1, \ldots, x_n)h(x_1, \ldots, x_n)dx_1 \cdots dx_n,
\]

where the \( \neq \) sign over the summation means that \( x_1, \ldots, x_n \) are pairwise distinct. See also Møller and Waagepetersen (2007).

Let \( C : B \times B \to \mathbb{R} \) denote a covariance function. A simple locally finite spatial point process \( X \) on \( B \) is called a determinantal point process with kernel \( C \) if its product density functions are

\[
\rho^{(n)}(x_1, \ldots, x_n) = \text{det}(C)(x_1, \ldots, x_n), \quad (x_1, \ldots, x_n) \in B^n,
\]

for \( n = 1, 2, \ldots \), where \( [C](x_1, \ldots, x_n) \) is the \( n \times n \) matrix with entries \( C(x_i, x_j) \). We write \( X \sim \text{DPP}_B(C) \); when \( B = \mathbb{R}^d \) we write \( X \sim \text{DPP}(C) \). Note that, if \( A \) is a Borel subset of \( B \), then the restriction \( X_A := X \cap A \) of \( X \) to \( A \) is a DPP with kernel given by the restriction of \( C \) to \( A \times A \).

By Theorem 1 in Lavancier et al. (2015), first proved by Macchi (1975), such DPP’s exist under the two following conditions:

- \( C \) is a continuous covariance function, hence, by Mercer’s Theorem, for any compact subset \( S \),

\[
C(x, y) = \sum_{k=1}^{+\infty} \lambda_k^S \phi_k(x)\phi_k(y), \quad (x, y) \in S \times S,
\]

where \( \{\lambda_k^S\} \) and \( \{\phi_k(x)\} \) are the eigenvalues and eigenfunctions in Mercer’s Theorem representation of \( C \) restricted to \( S \times S \), respectively;

- \( \lambda_k^S \leq 1 \) for all compact \( S \) in \( \mathbb{R}^d \) and all \( k \).

Note that, since \( \rho^{(n)}(x_1, \ldots, x_n) \) defined above has the interpretation of the density of a point \( (x_1, \ldots, x_n) \), and the determinant of the matrix \( [C](x_1, \ldots, x_n) \) is the hypervolume of the parallelepiped spanned by the columns of this matrix, the density grows as the columns become more separated. Continuity of the covariance function \( C \) implies that \( \rho^{(n)}(x_1, \ldots, x_n) \) converges to 0 if \( x_i \to x_j \) for some \( i \neq j \). Moreover, since the determinant of \( [C](x_1, \ldots, x_n) \) is less than or equal to the product of its diagonal
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When $\lambda$ and $\rho$ define the covariance function to be isotropic, is the Fourier transform of $\phi_k(x)\phi_k(y)$, $x, y \in S$. (4)

When $n = 0$ the density (as well as the determinant) is defined to be equal to 0. See Möller and Waagepetersen (2007) for a thorough definition of absolute continuity of a spatial process with respect to the unit rate Poisson process. However, note that from the first part of (2) we have $\mathbb{P}(N(S) = 0) = \prod_{k=1}^{+\infty} (1 - \lambda_k^S)$; this probability could be positive due to the assumption $\lambda_k^S < 1$ for all $k = 1, 2, \ldots$.

From now on we restrict our attention to stationary DPP’s, that is, when $C(x, y) = C_0(x - y)$, where $C_0 \in L^2(\mathbb{R}^d)$ is such that its spectral density $\varphi$ exists, i.e.

$$C_0(x) = \int_{\mathbb{R}^d} \varphi(y) \cos(2\pi x \cdot y) dy, \quad x \in \mathbb{R}^d$$

and $x \cdot y$ is the scalar product in $\mathbb{R}^d$. If $\varphi \in L^1(\mathbb{R}^d)$ and $0 \leq \varphi \leq 1$, then the DPP$(C)$ process exists. Summing up, the distribution of a stationary DPP can be assigned by its spectral density; see Corollary 1 in Lavancier et al. (2015). Notice that $C_0$, which defines the covariance function to be isotropic, is the Fourier transform of $\varphi$, so that they are in one-to-one correspondence. Therefore, we can indistinctly identify one by the other, which explains why we can define the DPP by $\varphi$.

To explicitly evaluate (3) over $S = \left[ -\frac{1}{2}, \frac{1}{2} \right]^d$, we approximate $\tilde{C}$ as suggested in Lavancier et al. (2015). In other words, we approximate the density of $X$ on $S$ by

$$f_{\text{app}}(\{x_1, \ldots, x_n\}) = e^{[S] - D_{\text{app}} \text{det}[\tilde{C}_{\text{app}}]}(x_1, \ldots, x_n),$$ (5)
where \( \{x_1, \ldots, x_n\} \subset S \) and
\[
\tilde{C}_{\text{app}}(x, y) = \tilde{C}_{\text{app}, 0}(x - y) = \sum_{k \in \mathbb{Z}^d} \left[ \frac{\varphi(k)}{1 - \varphi(k)} \right] \cos(2\pi k \cdot (x - y)), \quad x, y \in S, \quad (6)
\]
\[
D_{\text{app}} = \sum_{k \in \mathbb{Z}^d} \log \left( 1 + \frac{\varphi(k)}{1 - \varphi(k)} \right). \quad (7)
\]

To understand why the approximation \( C(x, y) \approx C_{\text{app}, 0}(x - y) \) for \( x - y \in S \) follows, as well as the corresponding approximation for the tilted versions of these functions, we observe that the exact Fourier expansion of \( C_0(x - y) \) in \( S \) is as in (6) with the real part of \( \int_S C_0(y)e^{-2\pi ik \cdot y}dy \) instead of \( \varphi(k) \); assuming \( C_0 \) is such that \( C_0(t) \approx 0 \) for \( t \notin S \), then
\[
\Re\left( \int_S C_0(y)e^{-2\pi ik \cdot y}dy \right) \approx \varphi(k) := \Re\left( \int_{\mathbb{R}^d} C_0(y)e^{-2\pi ik \cdot y}dy \right).
\]
See also Lavancier et al. (2015), Section 4.1. See Figure 1 in the Supplementary Material, where we display an example of the function \( \varphi(k) \). Indeed, all the quantities in formula (2) can be obtained in this way.

When \( S = R \) is a rectangle in \( \mathbb{R}^d \), we can always find an affine transformation \( T \) such that \( T(R) = S = \left[ \begin{array}{cccc} -1 & 1 \\ 1 & 2 \end{array} \right] \). Define \( Y = T(X) \). If \( f_{Y_{\text{app}}} \) is the approximate density of \( Y \) as in (5), we can then approximate the density of \( X_R \) by
\[
f_{\text{app}}(\{x_1, \ldots, x_n\}) = |R|^{-n} \exp(-|S|f_{Y_{\text{app}}}(T(\{x_1, \ldots, x_n\}))), \quad (8)
\]
for \( \{x_1, \ldots, x_n\} \subset R \). In practice, the summation over \( \mathbb{Z}^d \) in (6) above is truncated to \( \mathbb{Z}_N^d \), where \( \mathbb{Z}_N := \{-N, -N + 1, \ldots, 0, \ldots, N - 1, N\} \) (see Section 4.3 in Lavancier et al., 2015).

As mentioned in the Introduction, one particular example of spectral density that we found useful is
\[
\varphi(x; \rho, \nu) = s^d \exp \left\{ - \left( \frac{s}{\sqrt{\pi}} \right)^\nu \left( \frac{\Gamma(\frac{\nu}{2} + 1)}{\Gamma(\frac{\nu}{2} + 1)} \right)^{\nu d} \rho^{\nu d} ||x||^\nu \right\}, \quad \rho, \nu > 0, \quad (9)
\]
for fixed \( s \in (0, 1) \) (e.g. \( s = \frac{1}{2} \)) and \( ||x|| \) is the Euclidean norm of \( x \in \mathbb{R}^d \). This function is the spectral density of a power exponential spectral model. See (3.7) in Lavancier et al. (2015) setting \( \alpha = s \alpha_{\max}(\rho, \nu) \). In this case, we write \( X \sim \text{PES} - \text{DPP}(\rho, \nu) \). The corresponding spatial process is isotropic. When \( \nu = 2 \), the spectral density is
\[
\varphi(x; \rho, \nu) = s^d \exp \left\{ - \frac{s^2 \rho^{2d}}{\sqrt{\pi}} ||x||^2 \right\}, \quad \rho > 0,
\]
corresponding to the Gaussian spectral density. We discuss more specifically the choice of (9) later in Section 4.
2.2 The mixture model with repulsive means

To deal with limitations of model (1) or DPMs, we consider repulsive mixtures. Our aim is to estimate a random partition of the available subjects, and we want to do so using “few” groups. By repulsion we mean that cluster locations are a priori encouraged to be well separated, thus inducing fewer clusters than if they were allowed to be independently selected. We start from parametric densities \( f(\cdot; \theta) \), which we take to be Gaussian, and assume that the collection of location parameters follows a DPP. We specify a hierarchical model that achieves the goals previously described. Concretely, we propose:

\[
\begin{align*}
    y_i \mid s_i = k, \{\mu_k\}, \{\sigma_k^2\}, K & \sim \text{N} \left( y_i; \mu_k, \sigma_k^2 \right) \text{ for each } i \\
    X = \{\mu_1, \mu_2, \ldots, \mu_K, K\} & \sim \text{PES-DPP}(\rho, \nu) \\
    (\rho, \nu) & \sim \pi \\
    p(s_i = k) = w_k, \ k = 1, \ldots, K \text{ for each } i \\
    w_1, \ldots, w_K \mid K & \sim \text{Dirichlet}(\delta, \delta, \ldots, \delta) \\
    \sigma_k^{-2} \mid K & \sim \text{Gamma}(a_0, b_0),
\end{align*}
\]

where the PES-DPP(\( \rho, \nu \)) assumption (11) is regarded as a default choice that could be replaced by any other valid DPP alternative. We stress the explicit role of \( K \) in formula (12), where \( K \) represents the (random) total number of points generated by the determinantal point process prior. By the approximation procedure described in Section 2.1, \( K \) represents the total number \( N(S) \) of points in the compact set \( S \), assumed to be large enough to contain all required latent cluster means (points); see (2). Recall also that in practice all we need is to work on the cube \([-1/2, 1/2]^d\] and then use a linear transformation of this cube onto a cube large enough to cover all required centers, as just described. The choice of \( \pi \) in (12) will be discussed below in Section 4. We note that, as stated, the prior model may assign a positive probability to the case \( K = 0 \). This case of course makes no sense from the viewpoint of the model described above. Nevertheless, we adopt the working convention of redefining the prior to condition on \( K \geq 1 \), i.e., truncating the DPP to having at least one point. In practice, the posterior simulation scheme later described simply ignores the case \( K = 0 \), which produces the desired result. Note also that we have assumed prior independence among blocks of parameters not involving the locations \( \mu_k \).

Model (10)-(15) is a DPP mixture model along the lines proposed in Xu et al. (2016). Indeed, we both use DPPs as priors for location points in the mixture of parametric densities. However, the specific DPP priors are different, as they restrict to a particular case of DPPs (L-ensembles) that require a finite state space, and choose a Gaussian covariance function for which eigenvalues and eigenfunctions in the corresponding Mercer’s Theorem representation are analytically available. We adopt instead the spectral approach for assigning the prior (11), which implies great generality of the proposal. Similar to Xu et al. (2016), we carry out posterior simulation using a reversible jump step as part of the Gibbs sampler. However, when updating the location points \( \mu_1, \ldots, \mu_K \)
we refer to formulas (5)-(8). Xu et al. (2016) take advantage of the analytical expressions that we do not have for our case, and that are also unavailable in other possible specific choices of the spectral density.

As a general comment, we underline that the numerical evaluation of the DPP density, involving the computation of the determinant of a $K \times K$ matrix, is not particularly expensive, even in case of a large dataset; in this case, the repulsion property will favor a moderate number $K$ of clusters. See Section 4.4, where we describe applications of this model to datasets, using the posterior simulation algorithms described in Section 1 in the Supplementary material. In our experience, the proposed model scales well compared to mixtures with independent components.

## 2.3 Competitor repulsive models

We briefly introduce the class of parsimonious mixture models in Quinlan et al. (2017), to be used as a competitor model for our applications. Quinlan et al. (2017) exploit the idea of repulsion, i.e. when any two mixture components are encouraged to be well separated, as we do. For the sake of comparison, we introduce their model for unidimensional data: similarly to our case, they consider a mixture of $K$ Gaussian components, but assume a fixed value $k$ for $K$ in (10) and (13)-(15). The prior for the location parameters $\mu_1, \ldots, \mu_k$ is called repulsive distribution, and denoted by $NRep_k(\mu, \Sigma, \tau)$, where $\mu \in \mathbb{R}, \Sigma, \tau > 0$; see (3.4)-(3.6) in Quinlan et al. (2017). This prior is characterized by a repulsion potential that assumes the following expression:

$$\phi_1(r; \tau) = -\log \left(1 - e^{-\frac{1}{2\tau}r^2}\right) 1_{(0, +\infty)}(r), \quad \tau > 0;$$

Petralia et al. (2012) use a similar model, where the repulsion potential is

$$\phi_2(r; \tau) = \frac{\tau}{r^2} 1_{(0, +\infty)}(r), \quad \tau > 0.$$

Potential $\phi_2$ introduces a stronger repulsion than $\phi_1$, in the sense that in Petralia et al. (2012), locations are encouraged to be further apart than in Quinlan et al. (2017). Note also that, by nature of the point process, our approach does not require an upper bound on the allowed number of mixture components (similar to DPM models), contrary to the approach in Quinlan et al. (2017) and Petralia et al. (2012).

All models under comparison for a specific application will be matched in such a way that they have the same prior expected number $K$ of components in the mixtures.

## 3 Generalization to covariate-dependent models

The methods discussed in Section 2 were devised for density estimation-like problems. We now extend the previous modeling to the case where $p$-dimensional covariates $z_1, \ldots, z_n$ are recorded as well. We do so by allowing the mixture weights to depend on such covariates. In this case, there is a trade-off between repulsiveness of locations in
the mixtures and attraction among subjects with similar covariates. We also entertain
the case where covariate dependence is added to the likelihood part of the model. Our
modeling choice here is akin to mixtures of experts models (see, e.g., McLachlan and
Peel, 2005), i.e., the weights are defined by means of normalized exponential function.

Building on the model from Section 2.2, we assume the same likelihood (10) and the
DPP prior for $X = \{\mu_1, \mu_2, \ldots, \mu_K, K\}$ in (11)-(12), but change (13) and (14) to

$$p(s_i = k) = w_k(z_i) = \frac{\exp(\beta_k^T z_i)}{\sum_{l=1}^K \exp(\beta_l^T z_i)}, \quad k = 1, \ldots, K$$

(16)

$$\beta_2, \ldots, \beta_K \mid K \overset{\text{iid}}{\sim} \mathcal{N}_p(\beta_0, \Sigma_0), \quad \beta_1 = 0,$$

(17)

where the $\beta_1 = 0$ assumption is to ensure identifiability. To complete the model, we
assume (15) as the conditional marginal for $\sigma_k^2$; the prior for $(\rho, \nu)$ in (12) is later
specified. Here $\beta_0 \in \mathbb{R}^p$, and to choose $\Sigma_0$, we use a g-prior approach, namely
$\Sigma_0 = \phi \times (Z^T Z)^{-1}$, where $\phi$ is fixed, typically of the same order of magnitude of
the sample size (see Zellner, 1986). Here $Z$ denotes the design matrix.

Assuming now (10) on top of (16)-(17) rules out the case of a likelihood explicitly
dependent on covariates, which instead would generally achieve a better fit than other-
wise. Of course, there are many ways in which such dependence may be added. For the
sake of concreteness, we assume here a Gaussian regression likelihood, where only the
intercept parameters arise from the DPP prior. More precisely, we assume

$$y_i \mid s_i = k, z_i, \{\mu_k\}, \{\sigma_k^2\}, K \overset{\text{iid}}{\sim} \mathcal{N}(y_i; \mu_k + z_i^T \gamma_k, \sigma_k^2)$$

(18)

for all $i = 1, \ldots, n$, and

$$\gamma_i \mid K, \sigma_i^2 \overset{\text{iid}}{\sim} \mathcal{N}(\gamma_0, \sigma_i^2 \Lambda_0) \quad \text{and} \quad \sigma_i^{-2} \overset{\text{iid}}{\sim} \text{Gamma}(a_0, b_0),$$

(19)

where the $\gamma_k$'s are $p$-dimensional regression coefficients, $\gamma_0 \in \mathbb{R}^p$ and $\Lambda_0$ is a covariance
matrix. The prior for $\{s_i\}$ and $\beta_j$'s is given in (16)-(17) as in the previous model. Note
that (18) implies that only the intercept term is distributed according to the repulsive
prior. Thus, we allow the response mean to be corrected by a linear combination of the
covariates with cluster-specific coefficients, with the repulsion acting only on the residual
of this regression. The result is a more flexible model than the repulsive mixture (10)-
(15). Observe that there is no need to assume the same covariate vector in (18) and
(16), but we do so for illustration purposes only.

The Gibbs sampler algorithm employed to carry out posterior inference for this
model is detailed in Section 3 of the Supplementary material. However, it is worth noting
that the reversible jump step related to updating the number of mixture components
$K$ and the update of the coefficients $\{\beta_2, \beta_3, \ldots, \beta_K\}$ are complicated by the presence
of the covariates. For the $\beta$ coefficients, we resort to a Metropolis-Hastings step, with a
multivariate Gaussian proposal centered in the current value. For $K$, we employ an ad
hoc Reversible Jump move.
4 Simulated data and reference datasets

Before illustrating the application of our models to specific datasets, we discuss some general choices that apply to all examples.

4.1 Spectral density and other general choices

As discussed earlier, our methods are quite general in the sense that they can be used with any valid spectral density. Practical implementation, however, requires choosing one such function, and the power exponential function in (9) is a convenient and flexible choice. In this case, note that \( \varphi(x; \rho, \nu) < 1 \) when \( 0 < s < 1 \) for any \( x \in \mathbb{R}^d, \rho, \nu > 0 \), so that \( X_S \) has a density as described in (3). Figure 2 in the Supplementary material shows a plot of the power exponential spectral density (9) for different values of parameters \( \rho, \nu \). Note that \( \nu \) controls the shape of \( \varphi(x; \rho, \nu) \), which ranges from a slowly decreasing function of \( x \) when \( \nu \) is small, to an indicator function when \( \nu \) is large. On the other hand \( \rho \) plays the role that resembles a location parameter, with higher values retarding the decay speed of \( \varphi(x; \rho, \nu) \). As discussed in Section 2.1, knowledge about the spectral density is all that is needed for the approximations to work. Moreover, even if the analytic expression of \( C(x, y), (x, y) \in \mathbb{R}^d \times \mathbb{R}^d \) is known, as in, e.g. (9), we still need to compute the eigenvalues and eigenfunctions (of Mercer’s Theorem representation) of \( C \) restricted to \( S \times S \) for any compact \( S \), and this may be analytically impossible. A potential disadvantage derived from this is that parameter interpretation in the spectral domain becomes unclear. We reckon this is a problem that arises for any particular parametrized version of \( \varphi \) one may choose.

Prior elicitation for \( (\rho, \nu) \) is complicated due to the reasons stated above. Therefore, an extensive sensitivity analysis regarding \( \pi(\rho, \nu) \) was carried out for the datasets considered below. See Sections 4.2 and 4.3. We point out that an initial prior independence assumption \( \pi(\rho, \nu) = \pi(\rho) \pi(\nu) \) produced bad mixing of the chain. In particular, when \( \rho \) is small with respect to \( \nu \), the spectral function \( \varphi(\cdot) \) has a very narrow support, concentrated near the origin, forcing the covariance function \( \tilde{C}_{app}(x, y) \) in (6) to become nearly constant for \( x, y \in S \) and thus producing nearly singular matrices. We next investigated the case \( \pi(\rho, \nu) = \pi(\rho | \nu) \times \pi_\nu(\nu) \), where

\[
\rho | \nu \overset{d}{=} M(s, \varepsilon, \nu) + \rho_0, \quad \rho_0 \sim \text{Gamma}(a_\rho, b_\rho).
\]

Here, \( M(s, \varepsilon, \nu) \) is a constant that is the minimum value of \( \rho \) such that \( \varphi(2) > \varepsilon \) (here \( \varphi(2) \) is a reference value chosen to avoid a small support), and \( \varepsilon \) is a threshold value, assumed to be small (0.05, for instance). From Figure 2 in the Supplementary Material, it is clear that \( \varphi(\cdot; \rho, \nu) \) goes to 0 too fast when \( \nu \) is small relative to \( \rho \). It follows that

\[
M(s, \varepsilon, \nu) = \frac{2s\Gamma(1/\nu + 1)\pi^{1/2}}{\Gamma(3/2) \left( \log \left( \frac{s}{\varepsilon} \right) \right)^{1/\nu}}.
\]

We considered two different choices for \( \pi_\nu \): a gamma distribution, which gave a bad chain mixing, and a discrete distribution on \( \mathcal{V}_2 = \{0.5, 1, 2, 3, 5, 10, 15, 20, 30, 50\} \) (or on
one of its subsets). In this case, the mixing of the chain was better, but the posterior for \( \nu \) did not discriminate among the values in the support. For this reason, in Sections 4.5 and 5 and in Section 5 of the Supplementary Material, we assume \( \nu = 2, s = 1/2 \) and

\[
\rho = \sqrt{\frac{\pi}{\log(\frac{1}{2\epsilon})}} + \rho_0, \quad \rho_0 \sim \text{Gamma}(a_\rho, b_\rho),
\]

These choices guarantee that a reasonable number of nonzero terms are present when evaluating \( \tilde{C}_{\text{app}}(x, y) \), avoiding either too few, and hence a near singularity when evaluating (8), or too many, which will cause excessive computational burden.

In what follows, every run of the Gibbs sampler (implemented in \( \text{R} \)) produced a final sample size of 5,000 or 10,000 iterations (unless otherwise specified), after a thinning of 10 and initial burn-in of 5,000 iterations. In all cases, convergence was checked using both visual inspection of the chains and standard diagnostics available in the \text{CODA} package.

### 4.2 Data illustration on reference datasets without covariates

We illustrate our model via two datasets without covariates with unidimensional (Galaxy data) and bidimensional (Air Quality data) observations, both publicly available in \( \text{R} \) (galaxy from the \text{DPpackage} and \text{airquality} in the base version). For the latter data set we removed 42 incomplete observations.

The popular dataset Galaxy contains \( n = 82 \) measured velocities of different galaxies from six well-separated conic sections of space. Values are expressed in Km/s, scaled by a factor of \( 10^{-3} \). We set the hyperparameters in this way: for the variance \( \sigma_k^2 \) of the components, \((a_0, b_0) = (3, 3)\) (such that the mean is 1.5 and the variance is \( 9/4 \)) and for the weights \( \{w_k\} \) the Dirichlet has parameter \((1, 1, \ldots, 1)\). The other hyperparameters are modified in the tests, as in Table 1, where we report summaries of interest, such as the prior and posterior mean and variance for the number of components \( K \). In addition, we also display the mean squared error (MSE) and the log-pseudo marginal likelihood (LPML) as indices of goodness of fit, defined as

\[
MSE = \sum_{i=1}^{n} (y_i - \hat{y}_i)^2 \quad \text{and} \quad LPML = \sum_{i=1}^{n} \log \left( f(y_i \mid \hat{y}_{(-i)}^i) \right),
\]

where \( \hat{y}_i \) is the posterior predictive mean and \( f(y_i \mid \hat{y}_{(-i)}^i) \) is the \( i \)-th conditional predictive ordinate, that is the predictive distribution obtained using the dataset without the \( i \)-th observation. Figure 1 shows density estimates and the estimated partition of the data, obtained as the partition that minimizes the posterior expectation of Binder’s loss function under equal misclassification costs (see Lau and Green, 2007). The points at the bottom of the plots represent observations, while colors refer to the corresponding cluster. See Figure 4 below for the posterior distribution of \( K \) for Test 4 and 6 in Table 1.

As a comparison, the same posterior quantities than in Table 1 were computed using the DPM, the Repulsive Gaussian Mixture Models (RGMM) by Quinlan et al. (2017), and also the proposal by Petralia et al. (2012). To make results comparable, we assumed the same prior information on hyperparameters common to all the mixture models. See Table 4. From these tables, it is clear that alternative repulsive models are
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Table 1: Prior specification for \((\rho, \nu)\) and \(K\) and posterior summaries for the Galaxy dataset; \((a_\rho, b_\rho)\) appear in (20); here \(\mathcal{V}_1\) is \(\{1, 2, 5, 10, 20\}\) and \(\mathcal{V}_2 = \{0.5, 1, 2, 3, 5, 10, 15, 20, 30, 50\}\).

| Test | \(\rho\) | \(\nu\) | \(E(K)\) | \(Var(K)\) | \(E(K|\text{data})\) | \(Var(K|\text{data})\) | MSE | LPML |
|------|----------|----------|----------|----------|-----------------|-----------------|-----|-----|
| 1    | 2        | 2        | 2        | 1.67     | 6.09            | 1.10            | 78.95 | -171.72 |
| 2    | 5        | 10       | 5.00     | 7.12     | 6.07            | 1.09            | 78.33 | -167.96 |
| 3    | \(a_\rho = 1, b_\rho = 1\) | 2 | 2.18 | 1.978 | 6.10 | 1.10 | 73.89 | -164.47 |
| 4    | \(a_\rho = 1, b_\rho = 1\) | 10 | 2.73 | 2.15 | 6.11 | 1.12 | 74.93 | -162.71 |
| 5    | \(a_\rho = 1, b_\rho = 1\) | \(\text{discr}(\mathcal{V}_1)\) | 2.47 | 2.21 | 6.06 | 1.08 | 74.02 | -172.54 |
| 6    | \(a_\rho = 1, b_\rho = 1\) | \(\text{discr}(\mathcal{V}_2)\) | 2.51 | 2.27 | 6.10 | 1.13 | 76.64 | -170.94 |

Figure 1: Density estimates and estimated partition for the Galaxy dataset under Test 4 in Table 1, including 90% credibility bands (light blue).

good competitors to ours, and that they generally achieve a better fit to the dataset. The tests showing the best indexes of goodness of fit are typically those overestimating the number of clusters. It is well-known that, in general, clustering in the context of DPMs and mixture models as those in Quinlan et al. (2017) and Petralia et al. (2012) is affected by the base measure. Our model, on the other hand, avoids the delicate choice of the base measure leading to more stable estimates of \(K\). And in fact, we found that our model has better goodness-of-fit indexes than the competitor repulsive mixtures in the case of a simulated dataset with 8 well-separated mixture components; see Section 4.4. This shows that our proposal is indeed a competitive alternative.

Finally, we recall that in Section 2 in the Supplementary material we report some further tests on the Galaxy dataset to show the influence of various choices of spectral density on the inference. We conclude that there is evidence that the approach is not sensitive with respect to the choice of spectral density.
We have considered one further application, this time using the same variables from the dataset Air Quality (ozone and solar radiation) as considered in Quinlan et al. (2017). Instead of (10), we assume that our likelihood is a bidimensional Gaussian, with bidimensional mean vectors distributed according to the PES $- DPP(\rho, \nu)$ prior as before, and with covariance matrices $\Sigma_k$ independent and identically distributed according to the inverse-Wishart distribution. See Section 1 in the Supplementary Material for changes in the Gibbs sampler with multidimensional data points, this time adapted from Dellaportas and Papageorgiou (2006). Table 2 reports summaries of interest for a few tests carried out, including the prior and posterior mean and variance for the number of components $K$, and the LPML. As usual in the context of other mixture models, we find that the inference depends on the chosen hyperparameters. If we compare with corresponding inference in Quinlan et al. (2017), we got lower estimates of $K$, and a better fit of the model to the data. The posterior predictive densities, not shown here, seem very similar to those in Quinlan et al. (2017), Fig. 9 (b).

| Test | $\rho$ | $\nu$ | $E(K)$ | $Var(K)$ | $E(K|data)$ | $Var(K|data)$ | LPML  |
|------|--------|-------|--------|----------|-------------|--------------|-------|
| 7    | 3      | 2     | 3      | 2.62     | 2.18        | 0.39         | -246.81|
| 8    | $\rho_0 \sim$ Gamma(1, 0.5) | 2     | 2.7    | 2.37     | 2.15        | 0.21         | -257.66|

Table 2: Prior specification for $(\rho, \nu)$ and $K$ and posterior summaries for the airquality dataset; $\rho_0$ appear in (20).

### 4.3 Tests on data from a mixture with 8 components

We simulated a dataset with $n = 100$ observations from a mixture of 8 components. Each component is the Gaussian density with mean $\theta_k$ and $\sigma_k^2 = \sigma^2 = 0.05$: the means $\{\theta_k\}$ are evenly spaced in the interval $(-10, 10)$. In the model (10)-(15), we set $a_0 = 2.0025$, $b_0 = 0.050125$ so that $E(\rho_0) = 0.05$ and $Var(\rho_0) = 1$; again, $s = 0.5$ and $\delta = 1$.

Table 3 reports hyperparameters values for different tests and posterior summaries of interest, as well as prior mean and variance of $K$. In particular, we show the posterior mean and variance for the number of components $K$ (with which we assess the effectiveness of the model for clustering), the mean squared error (MSE) and the log-pseudo marginal likelihood (LPML) (that helps quantifying the goodness-of-fit). In all cases we obtain a pretty satisfactory estimate of the exact number of components, which is 8: the posterior is concentrated around the true value with a very small variance. In particular, we got that $P(K = 8|data)$ is equal to 0.96 and 0.93 under Tests $S_2$ and $S_7$ in Table 3, respectively.

From the density estimation viewpoint, we have from Table 3 that both MSE and LPML are similar for all the tests, thus indicating insensitiveness with respect to the prior choice of parameters $\rho$ and $\nu$. However, preferable tests seem to be $S_2$ and $S_7$; see Figure 2, where density estimates and estimated partitions for these two cases are displayed. The posterior density of $\rho$ under Tests $S_2$ and $S_7$ is shown in Figure 3.
Table 3: Prior specification for $(\rho, \nu)$ and the corresponding mean and variance induced on $K$ (top). Hyperparameters $(a_\rho, b_\rho)$ appear in (20), while $\mathcal{V}_1 = \{1, 2, 5, 10, 20\}$ and $\mathcal{V}_2 = \mathcal{V}_1 \cup \{0.5, 3, 15, 30, 50\}$. Posterior summaries for the simulated dataset from a mixture with 8 components are in the bottom subtable.

Figure 2: Density estimate and estimated partition for the simulated dataset from the mixture of 8 components under Tests $S_2$ (left) and $S_7$ (right) in Table 3. The points at the bottom of the density estimate represent the data, and each color represents one of the eight estimated clusters.
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Figure 3: Posterior distribution of $\rho$ for the simulated dataset from the mixture of 8 components under Tests $S_2$ (left) and $S_7$ (right) in Table 3.

| Test | $\alpha$ | $\mathbb{E}(K)$ | $\mathbb{E}(K|data)$ | $\text{Var}(K|data)$ | MSE | LPML |
|------|----------|-----------------|----------------------|----------------------|-----|------|
| $D_1$ | Gamma(0.5, 1) | 7.7 | 7.271 | 1.594 | 36.708 | -149.258 |
| $D_2$ | 0.8 | 4.3 | 5.936 | 1.25 | 61.255 | -151.146 |
| $D_3$ | 0.45 | 3 | 4.371 | 1.142 | 139.659 | -169.978 |
| $D_4$ | Gamma(4, 2) | 2.9 | 6.166 | 1.549 | 62.703 | -151.797 |

Table 4: Prior specification for $\alpha$ and posterior summaries for the Galaxy dataset using the function DPdensity in DPpackage (top) and repulsive models (bottom).

As a comparison, the same posterior quantities than in Table 1 were computed; see Tables 4 and 5. The DPM was fitted via the function DPdensity available from DPpackage (Jara et al., 2011), while the code for the alternative repulsive models was gently provided by José Quinlan and Garritt Page.

4.4 Comparison to alternative models

We now consider fitting alternative models to the Galaxy and two simulated datasets, one from the mixture with 8 components introduced in the previous section, and the second consisting of 10,000 observations generated from a mixture of 20 components. We consider first the gold standard of Bayesian nonparametric models, the DPM, and
Table 5: Posterior summaries for the simulated dataset from the mixture of 8 components using the function DPdensity in DPpackage (top) and repulsive models (bottom).

| DPM      | Test | $\alpha$ | $E(K)$ | $E(K|data)$ | $Var(K|data)$ | MSE  | LPML  |
|----------|------|----------|--------|-------------|--------------|------|-------|
| DPM      | $D_5$ | 0.43     | 3      | 7.961       | 0.5          | 4.779 | -11.246 |
|          | $D_6$ | Gamma(4,2) | 8.17   | 8.665       | 0.910        | 4.248 | -10.116 |

| Repulsive models | Model | $E(K|data)$ | $Var(K|data)$ | MSE  | LPML  |
|------------------|-------|-------------|--------------|------|-------|
| Quinlan et al. (2017) | 10.73 | 1.407       | 3.121        | -4.754 |
| Petralia et al. (2012) | 8.51 | 0.357       | 4.152        | -4.022 |

Comparison of the tables above and Tables 1 and 3 show that the alternative repulsive models are good competitors to ours, and according to the dataset and hyperparameters specification, they may achieve a better (Galaxy) or worse (simulated data) fit to the data. The tests showing the best indexes of goodness of fit are typically those overestimating the number of clusters. It is well-known that, in general, clustering in the context of nonparametric mixture models as DPMs is strongly affected by the base measure (see, e.g. Miller and Harrison, 2017). The same disadvantage affects the mixture models in Quinlan et al. (2017) and Petralia et al. (2012). Our model, on the other hand, avoids the delicate choice of the base measure leading to more stable estimates of $K$.

As a further comparison, see also Figure 4 which displays the posterior distribution of $K$ under the DPM mixture and our models for the Galaxy dataset.

For the second simulated dataset, we considered applicability for a moderately large sample size, generating 10,000 observations from a 20-component mixture, 10 of them being Gaussian, and the rest being skew-normal distributions with positive and negative skewness. The true density is showed in Figure 5. To estimate the true number of clusters, we fitted different alternative models to this dataset: our model, the repulsive mixture models by Quinlan et al. (2017) and Petralia et al. (2012), and the finite mixture model implemented in the mclust R package via the function Mclust (Fraley et al., 2012) with a number of components between 10 and 25. The same prior information on hyperparameters common to all the Bayesian mixture models was assumed. The Mclust function returns the estimates of the number of components corresponding to the “best three” models, in this case 11, 17 and 18. See Table 6 for posterior summaries of the inference under the three repulsive models.
Figure 4: Posterior distribution of the number $K$ of components under Test 4 (black) and 6 (blue) in Table 1 and under the DPM model (red) as in Test D1 in Table 4.

Figure 5: Histogram, true density (red) and density estimate (black) of the large simulated dataset, including 90% credibility bands (light blue).

| Model              | $E(K|data)$ | $Var(K|data)$ | MSE    | LPML     |
|--------------------|-------------|---------------|--------|----------|
| $PES - DPP$        | 16.41       | 1.38          | 1356.43| -13239.54|
| Quinlan et al. (2017) | 14.13      | 0.146         | 1475.98| -13771.56|
| Petralia et al. (2012) | 20.81    | 0.564         | 1002.05| -12940.49|

Table 6: Posterior summaries for the large simulated dataset.

Though the run-time for this application is around 15 times longer than in the case of the Galaxy data, our algorithm reduces the effective number of clusters a posteriori, thus helping our model scaling up. Intuitively, the increase in the run-time is mostly due to the larger number of mixture components and the much larger sample size than in the case of other datasets illustrated here.
4.5 Simulated data with covariates

We consider the same simulated dataset as in Müller et al. (2011), Section 5.2; the simulation “truth” consists of 12 different distributions, corresponding to different covariate settings (see Figure 1 of that paper). Model (10)-(12), (15)-(17) was fitted to the dataset, assuming $\beta_0 = 0$, $\Sigma_0 = 400 \times (Z^TZ)^{-1}$, $a_\rho = 1$, $b_\rho = 1.2$, and $a_0, b_0$ such that the prior mean of $\sigma_k^2$ is 50 and variance is 300. Recall also that here we assume $\nu = 2$.

As an initial step, inference for the complete dataset (1000 observations) was carried out, yielding a posterior of $K$, not reported here, mostly concentrated over the set $\{8, 9, \ldots, 16\}$, with a mode at 11. Figure 3 in the Supplementary Material shows posterior predictive distributions for the 12 different reference covariate values, along with 90% credibility intervals. These are in good accordance with the simulation truth (compare with Figure 1 in Müller et al., 2011).

To replicate the tests in Müller et al. (2011), a total of $M = 100$ datasets of size 200 were generated by randomly subsampling 200 out of the 1000 available observations. Computational burden over multiple repetitions was controlled by limiting the posterior sample sizes to 2,000. Table 7 displays the root MSE for estimating $\mathbb{E}(y \mid z_1, z_2, z_3)$ for each of the 12 covariate combinations defining the different clusters for our model and for the PPM, as in Table 1 of Müller et al. (2011). The computations also include evaluation of the root MSE and LPML for all the 100 datasets for estimating the data used to train the model, with $MSE_{\text{train}} = \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$, where $\hat{y}_i$ is the expected value of the estimated predictive distribution, and for a test dataset of 100 new data, $MSE_{\text{test}} = \sum_{i=1}^{n} (y_{i \text{test}} - \hat{y}_i)^2$. In addition, we report $LPML_{\text{train}}$, value of the Log Pseudo Marginal Likelihood for the training dataset. Table 8 shows the values compared to other competitor models, i.e the linear dependent Dirichlet process mixture (LDDP) defined in De Iorio et al. (2004), the product partition model (PPM) in Müller et al. (2011) and the linear dependent tailfree process model (LDTFP) in Jara and Hanson (2011). The best values are in bold: our model performs well according to the LPML, while the MSE suggests to use PPM or LDTFP. In general, our model is competitive with respect to other popular models in the literature. Moreover, in the LDDP case, we have that the average number of clusters is 20.6 with variance 2.266, thus indicating a less parsimonious model compared to ours.

In summary, our extensive simulations suggest that the proposed approach tends to require less mixture components than non-repulsive counterpart, while still providing a reasonably good fit to the data. Note that, with respect to other repulsive competitors, our results are comparable.

5 Biopic movies dataset

For this illustrative example we consider the Biopics data available in the R package fivethirtyeight (Ismay and Chunn, 2017). This dataset is based on the IMDB database, related to biographical films released from 1915 through 2014. An interest-
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Table 7: Root MSE for estimating \( \mathbb{E}(y \mid z_1, z_2, z_3) \) for 12 combinations of covariates \( (z_1, z_2, z_3) \) and \( PPM_x \) as competing model of reference (compare also the results in Table 1 of Müller et al., 2011).

| z1 | z2 | z3 | DPP | PPM_X |
|----|----|----|-----|-------|
| -1 | 0  | 0  | 6.1 | 7.9   |
| 0  | 0  | 0  | 6.7 | 3.9   |
| 1  | 0  | 0  | 7.2 | 2.8   |
| -1 | 1  | 0  | 6.5 | 5.4   |
| 0  | 1  | 0  | 6.5 | 4.6   |
| 1  | 1  | 0  | 6.8 | 4.0   |
| -1 | 0  | 1  | 6.8 | 6.1   |
| 0  | 0  | 1  | 6.1 | 4.2   |
| 1  | 0  | 1  | 5.7 | 4.5   |
| -1 | 1  | 1  | 5.9 | 9.5   |
| 0  | 1  | 1  | 6.6 | 8.3   |
| 1  | 1  | 1  | 5.8 | 6.2   |
| avg | | | **6.4** | **5.6** |

Table 8: Comparison with competitors for the simulated dataset with covariates: best values according to each index are in bold. DPPx denotes our model, while LDDP is the linear dependent Dirichlet process mixture, \( PPM_x \) is the product partition model with covariates, and LDTFP is the linear dependent tailfree process model.

|          | DPPx | LDDP | PPM_X | LDTFP |
|----------|------|------|-------|-------|
| Root MSE | 324.531 | 304.742 | **278.395** | 304.374 |
| Root MSE | 216.675 | 215.169 | 217.2459 | **212.761** |
| LPML     | **-871.8** | -902.2295 | -873.1671 | -901.465 |

We consider the logarithm of the gross earnings at US box office as a response variable, with the following covariates: (i) year of release of the movie (in a suitable scale, continuous); (ii) a binary variable that indicates whether the main character is a person of color; and (iii) a categorical variable that considers if the country of the movie is US, UK or other. After removing the missing data from the dataset, we were left with \( n = 437 \) observations and \( p = 4 \). We note that 76 biopics have a person of color as a subject and the frequencies of the category “origin” are \( (256, 79, 64) \) for US, UK and “other”, respectively; “other” means mixed productions (e.g. US and Canada, or US and UK). In what follows, the hyperparameters in model (18)-(19), (11)-(12), (16)-(17) are chosen as \( \beta_0 = 0, (a_0, b_0) = (1, 1) \). The prior mean and variance of \( K \) induced by these hyperparameters are 2.162 and 1.978, respectively. The scale hyperparameter \( \phi \) in the \( g \)-prior for \( \beta \) and \( (a_0, b_0) \) vary as determined in Table 9, where \( m \) and \( v \) denote the prior

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The explorative analysis of the data can be found in [https://fivethirtyeight.com/features/straight-outta-compton-is-the-rare-biopic-not-about-white-dudes/](https://fivethirtyeight.com/features/straight-outta-compton-is-the-rare-biopic-not-about-white-dudes/).
mean $b_0/(a_0 - 1)$ and variance $b_0^2/((a_0 - 1)^2(a_0 - 2))$, respectively, of the inverse gamma distribution for $\sigma_k^2$ as in (19). We also assume $\gamma_0$ equal to the vector of all 0’s, while $\Lambda_0$ is such that the marginal a priori variance of $\gamma_k$ is equal to diag(0.01, 0.1, 0.1, 0.1), in accordance to the variances of the corresponding frequentist estimators.

We have found that the posterior of $K$ is stable with respect to the choice of prior hyperparameters, i.e. it does not change significantly when changing the prior hyperparameters; on the other hand, our results show that by not including covariates in the likelihood, i.e. setting all $\gamma_k$’s are equal to 0, inference on $K$ is much more sensitive to the choice of $(a_0, b_0)$ (results not shown here).

Predictive inference was also considered, by evaluating the posterior predictive distribution at the following combinations of covariate values: (i) (mean value for covariate year, US, white); (ii) (mean value for covariate year, US, color); (iii) (mean value for covariate year, UK, white); (iv) (mean value for covariate year, UK, color); (v) (mean value for covariate year, “other”, white); and (vi) (mean value for covariate year, “other”, color). Corresponding plots are shown in Figure 6. These distributions appear to be quite different in the six cases: in particular, we can observe that in cases (i) and (ii), the posterior is shifted towards higher values. This is quite easy to interpret, since the measurements are given by the earnings in the US box offices; therefore, we expect that in general US movies will be more profitable in that market. The difference due to the race is, on the other hand, less evident. However, the predictive densities show slightly higher earnings for movies where the subject is a person of color, if the origin is “other” ((v) and (vi)). Movies from the UK, on the other hand, exhibit the opposite behavior ((iii) and (iv)).

We report here the posterior cluster estimate for Test B in Table 9, that is, the partition that minimizes the posterior expectation of Binder’s loss function under equal misclassification costs (see Lau and Green, 2007). We note that here, as it is also a common situation with point estimates of partitions, the number of clusters can be different from the posterior mean of $K$, 4.45 in this case. Indeed, we found three groups, with sizes 10, 193, 234, respectively; see Figure 7 for the estimated clusters and boxplots of the response. As a comparison, it can be useful to report the total average values for the response, 15.36, and for the covariates: 7.89 (year), 0.18 (UK), 0.15 (“other”),

| Test | $\phi$ | $m$ | $v$ | $\mathbb{E}(K \mid \text{data})$ | sd($K \mid \text{data}$) | MSE | LPML |
|------|--------|-----|-----|---------------------------|-----------------|-----|------|
| A    | 50     | 5   | 1   | 4.49                      | 1.10             | 1126.32 | -960.89 |
| B    | 200    | 5   | 10  | 4.45                      | 1.19             | 983.55 | -954.55 |
| C    | 50     | 3   | $+\infty$ | 5.66                  | 1.27             | 501.22 | -918.74 |
| D    | 200    | 10  | 5   | 4.21                      | 1.33             | 1805.83 | -980.61 |
| E    | 100    | 2   | 1   | 5.31                      | 1.21             | 564.26 | -935.56 |
| F    | 200    | 2   | 10  | 5.51                      | 1.26             | 557.44 | -925.22 |

Table 9: Prior specification for $\beta_k$’s and $\sigma_k^2$’s parameters and posterior summaries for the Biopics dataset; $m$ and $v$ are prior mean and variance, respectively, of $\sigma_k^2$. Posterior mean and variance of the number $K$ of mixture components are in the fifth and sixth columns, respectively, while the last two columns report MSE and LPML, respectively.
Figure 6: Predictive distribution for cases (i) – (vi) under Test E in Table 9 for the Biopics dataset.

Figure 7: Cluster estimate (left) under our model (Test B in Table 9) for the Biopics dataset. Each color represents one of the three estimated clusters. Coordinate $y$ is the response, i.e. box-office earning, while coordinate $x$ is the covariate year of release. The boxplot of the response per group is in the right panel.

These 3 groups have a nice interpretation in terms of covariates: group 1 is the smallest, with a high average response (17.18), and it is characterized by a high percentage of movies from “other” countries, with a person of color as its subject. Group 2 corresponds also to a high average response (16.42), but the average values of UK, “other” and person of color are similar to the total averages (0.14, 0.09, 0.84, 0.83) (white).
respectively). The average response in group 3 is smaller (14.40) than the total sample mean, while the average values of UK, “other” and person of color are 0.22, 0.17, 0.84, respectively.

To assess effectiveness of the proposed model, we compare the results with the linear dependent Dirichlet process mixture model introduced in De Iorio et al. (2004) and implemented in the \texttt{LDDPdensity} function of \texttt{DPpackage} (Jara et al., 2011). Prior information has been fixed as follows: for Test G the mass parameter of the Dirichlet process $\alpha$ is set equal to 0.3 such that $E(K) = 2.87$ and $Var(K) = 1.81$, that approximately match the prior information we gave on the parameter $K$. Similarly, under Test H, $\alpha$ is distributed according to the Gamma($1/4, 1/2$), such that the prior mean on $K$ is 3.6 and variance 22.18. The normal baseline distribution is a multivariate Gaussian with mean vector 0 and a random covariance matrix which is given a non-informative prior and the inverse-gamma distribution for the variances of the mixture components has parameters such that mean and variance are equal to 5, 1, respectively similarly as in Table 9. Posterior summaries can be found in Table 10.

| Case | $E(K | data)$ | $sd(K | data)$ | MSE | LPML |
|------|--------------|---------------|-----|------|
| $G$  | 2.95         | 1.03          | 1282.49 | -937.51 |
| $H$  | 3.56         | 2.36          | 682.98  | -914.00  |

Table 10: Posterior summaries for the tests on the Biopics dataset under a linear dependent Dirichlet process mixture.

As a comparison between the estimated partitions under our model (Figure 7) and the LDDP mixture model, Figure 4 in the Supplementary Material displays the estimated partition obtained under the LDDP model under Test $G$, that has 3 groups with sizes \{300, 127, 10\}.

6 Conclusion

This work deals with mixture models where the prior has the property of repulsion across location parameters. Specifically, the discussion is centered on mixtures built on determinantal point processes (DPPs), that can be constructed using a general spectral representation. The methods work with any valid spectral density, but for the sake of concreteness, illustrations were discussed in the context of the power exponential case. Implementing the posterior inference for our models requires the use of numerical approximations of the density function of the DPP.

Our approach has focused on mixture modeling. One may use essentially the same models for clustering purposes. In such case, it is quite simple to modify the cluster-specific covariance matrices to have any particular form, such as enforcing elliptically- or circularly-shaped contours. We have here used standard assumptions such as unstructured covariance matrices as a default choice in our discussion, but again, such assumptions, or others, such as stick-breaking mixture weights, can be easily incorporated to the model definitions.
Repulsiveness can be measured in a number of ways, as discussed in the online supplementary material in Lavancier et al. (2015), Appendix J. These include Ripley’s K-function and the Pair Correlation Function. They both measure how dispersed the points appear to be and they are usually compared to the Poisson process, where for a given a number of points, they are randomly scattered. Some analytical results can be obtained, but in general, one must evaluate these functions numerically. The power exponential case includes the most repulsive type of stationary DPP when $\nu \to \infty$. When $d = 2$, this corresponds to a “jinc-like” covariance function, that is, the case in which the spectral density is of the form $\phi(x) = I_{\{\|x\| \leq \tau\}}$. We found out that in most applications a value of $\nu = 2$ is sufficient for reasonable inference results, but larger values of $\nu$ also provide stable enough inference.

Though we limit ourselves to the case of isotropic DPPs, inhomogeneous DPPs can be obtained by transforming or thinning a stationary process. However, we believe that this case is not very interesting, unless there is a strong reason to assume non-homogeneous locations a priori.

Our computational experiments and data illustrations show that the repulsion induced by the DPP priors indeed tends to eliminate the annoying case of many very small clusters that commonly arises when using models that do not constrain location/centering parameters. This happens with very small sacrifice of model fit compared to the usual mixture models. In this sense, since it is a parsimonious model with respect to the clustering structure, interpretation is facilitated because there are typically less clusters to interpret.

From a computational viewpoint, we found our model to scale reasonably well with respect to sample size, compared to similar alternatives. However, scaling with respect to data dimension is not a property we claim to have. In fact, this is a situation shared with mixture models either with a finite or infinite number of components.

Another advantage of our model over DPMs and natural competitors in the repulsive mixture models framework, such as those in Quinlan et al. (2017) and Petralia et al. (2012), is that we avoid the delicate choice of the base measure of the Dirichlet process, leading to more stable estimates on the number $K$ of components in the mixture. Moreover, with respect to competitor repulsive mixtures, and just in the case of DPM models, our approach does not require an upper bound on the allowed number of mixture components. This particular aspect can also be considered as a con, because the competitor repulsive mixtures may produce slightly better performance in terms of goodness of fit. On the other hand, since $K$ is finite and random, our algorithm requires a reversible jump implementation, which could be seen in practice as an extra computational price to pay when using the proposed model.

Another worth noting aspect of the model is the roles of subsets $S$ and $R$. They are both required to be compact so that the prior joint density of latent cluster centers is well defined. Subset $S$, in practice fixed as the unit hypercube, is specifically needed to guarantee that approximations (6) and (7) work well. In turn, the role of the rectangle $R$ is to cover the empirical data range, and the affine transformation $T$, which maps $R$ back onto $S$, is used to ensure that all calculations are carried in $S$. The size of $R$ has
little consequence in itself, beyond obvious changes in formula (8), because the prior
distribution of number of points is still determined by (2), i.e. by $S$ and $C_0$. And a very
large set $R$ would simply have large portions with very low posterior mass assigned,
corresponding to places with no observed data.

Summing up, through extensive computational experience in various settings, we
believe our model to be a valid and competitive alternative that, with little sacrifice in
model fit, may help achieving a more parsimonious representation of mixtures, compared
to other similar models.

**Supplementary Material**

Supplementary Material for “Determinantal Point Process Mixtures Via Spectral Den-
sity Approach” (DOI: 10.1214/19-BA1150SUPP; .pdf). Supplementary materials for this article are
available online. In particular, the document contains a description of the two Gibbs
samplers used for posterior inference under the DPP mixture model (Sections 1 and 3),
without and with covariate dependence, respectively. Moreover, a new test on the Galaxy
dataset when different spectral densities are considered (Section 2). Finally, supplemen-
tal figures that did not fit in the paper and additional analysis of an Air Quality Index
dataset are provided.

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