Distributed, partially collapsed MCMC for Bayesian nonparametrics

Avinava Dubey*  Michael M. Zhang*  Eric P. Xing  Sinead A. Williamson
Google Research  Princeton University  Carnegie Mellon Univ.  Univ. of Texas Austin

Abstract

Bayesian nonparametric (BNP) models provide elegant methods for discovering underlying latent features within a data set, but inference in such models can be slow. We exploit the fact that completely random measures, which commonly-used models like the Dirichlet process and the beta-Bernoulli process can be expressed using, are decomposable into independent sub-measures. We use this decomposition to partition the latent measure into a finite measure containing only instantiated components, and an infinite measure containing all other components. We then select different inference algorithms for the two components: uncollapsed samplers mix well on the finite measure, while collapsed samplers mix well on the infinite, sparsely occupied tail. The resulting hybrid algorithm can be applied to a wide class of models, and can be easily distributed to allow scalable inference without sacrificing asymptotic convergence guarantees.

1 INTRODUCTION

Bayesian nonparametric (BNP) models are a flexible class of models whose complexity adapts to the data under consideration. BNP models place priors on infinite-dimensional objects, such as partitions with infinitely many blocks; matrices with infinitely many columns; or discrete measures with infinitely many atoms. A finite set of observations is assumed to be generated from a finite—but random—subset of these components, allowing flexibility in the underlying dimensionality and providing the ability to incorporate previously unseen properties as our dataset grows.

While the flexibility of these models is a good fit for large, complex data sets, distributing existing inference algorithms across multiple machines is challenging. If we explicitly represent subsets of the underlying infinite-dimensional object—for example, using a slice sampler—we can face high memory requirements and slow convergence. Conversely, if we integrate out the infinite-dimensional object, we run into problems due to induced global dependencies.

Moreover, a key goal of distributed algorithms is to minimize communication between agents. This can be achieved by breaking the algorithm into independent sub-algorithms, which can be run independently on different agents. In practice, we usually cannot split an MCMC sampler on a Bayesian hierarchical model into entirely independent sub-algorithms since there are typically some global dependencies implied by the hierarchy. Instead, we make use of conditional independencies to temporarily partition our algorithm.

Contributions: In this paper, we propose a distributable sampler for models derived from completely random measures, which unifies exact parallel inference for a wide class of Bayesian nonparametric priors, including the popularly used Dirichlet process [Ferguson 1973] and the beta-Bernoulli process [Griffiths and Ghahramani 2011]. After reviewing the appropriate background material, we first introduce general recipes for (non-distributed) partially collapsed samplers appropriate for a wide range of BNP models, focusing on the beta-Bernoulli process and the Dirichlet process as exemplars. We then demonstrate that these methods can be easily extended to a distributed setting. Finally we provide experimental results for our hybrid and distributed sampler on DP and BB inference.

2 RELATED WORK

Completely random measures (CRMs, Kingman 1967) are random measures that assign independent masses to disjoint subsets of a space. For example, the gamma process assigns a gamma-distributed mass to each sub-
Inference in such models tend to fall into three categories: uncollapsed samplers that alternate between sampling the latent measure and the assignments (Isswaran and Zarepour, 2002; Paisley and Carin, 2009; Zhou et al., 2009; Walker, 2007; Teh et al., 2007); collapsed samplers where the latent measure is integrated out (Isswaran and James, 2001; Neal, 2000; Ghahramani and Griffiths, 2005); and optimization-based methods that work with approximating distributions where the parameters are assumed to have a mean-field distribution (Blei and Jordan, 2006; Doshi-Velez et al., 2009a).

Collapsed methods often mix slowly due to the dependency between assignments, while blocked updates mean uncollapsed methods typically have good mixing properties at convergence (Isswaran and James, 2011). Uncollapsed methods are often slow to incorporate new components, since they typically rely on sampling unoccupied components from the prior. In high dimensions, such components are unlikely to be close to the data. Conversely, collapsed methods can make use of the data when introducing new points, which tends to lead to faster convergence (Neal, 2000).

Other methods incorporate both uncollapsed and collapsed sampling, resulting in a “hybrid”, partially collapsed sampler, although such approaches have been restricted to specific models. Doshi-Velez and Ghahramani (2009) introduced a linear time accelerated Gibbs sampler for conjugate IBPs that effectively marginalized over the latent factors, while more recently Yerebakan and Dundar (2017) developed a sampler by partially marginalizing latent random measure for DPs. These methods can be seen as special cases of our hybrid framework (Section 3), but do not generalize to the distributed setting.

Several inference algorithms allow computation to be distributed across multiple machines—although again, such algorithms are specific to a single model. The approximate collapsed algorithm of Smyth et al. (2009) is only developed for Dirichlet process-based models, and lacks asymptotic convergence guarantees. Distributed split-merge methods have been developed for Dirichlet process-based models, but not extended to more general nonparametric models (Chang and Fisher, 2013, 2014). Partition-based algorithms based on properties of CRMs have been developed for Dirichlet process- and Pitman-Yor process-based models (Williamson et al., 2013; Dubey et al., 2014), but it is unclear how to extend to other model families. A low-communication, distributed-memory slice sampler has been developed for the Dirichlet process, but since it is based on an uncollapsed method it will tend to perform poorly in high dimensions (Ge et al., 2015). Doshi-Velez et al. (2009b) developed an approximate distributed inference algorithm for the Indian buffet process which is superficially similar to our distributed beta-Bernoulli sampler. However, their approach allows all processors to add new features, which will lead to overestimating the number of features. We contrast

**Table 1: Comparison of various parallel and distributed inference algorithm proposed for BNPs. Other CRMs include gamma-Poisson process, beta-negative binomial process etc.**

<table>
<thead>
<tr>
<th>Methods</th>
<th>Data Size</th>
<th>Exact</th>
<th>Parallel</th>
<th>Distributed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Smyth et al. 2009</td>
<td>1M</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Doshi-Velez and Ghahramani 2009</td>
<td>.01M</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Doshi-Velez et al. 2009b</td>
<td>.1M</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Williamson et al. 2013</td>
<td>1M</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Chang and Fisher 2013</td>
<td>0.3M</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Dubey et al. 2014</td>
<td>10M</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Ge et al. 2015</td>
<td>0.1M</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Yerebakan and Dundar 2017</td>
<td>0.05M</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
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</tbody>
</table>

This paper

1M ✓ ✓ ✓ ✓ ✓ ✓ ✓
3 HYBRID INFERENCE FOR CRM-BASED MODELS

By definition, completely random measures can be decomposed into independent random measures. If the CRM has been transformed in some manner we can often still decompose the resulting random measure into independent random measures – for example, a normalized random measure can be decomposed as a mixture of normalized random measures. Such representations allow us to decompose our inference algorithms, and use different inference techniques on the constituent measures.

As discussed in Section 2, collapsed and uncollapsed methods both have advantages and disadvantages. Loosely, collapsed methods are good at adding new components and exploring the tail of the distribution, while uncollapsed methods offer better mixing in established clusters and easy parallelization. We make use of the decomposition properties of CRMs to partition our model into two components: One containing (finitely many) components currently associated with multiple observations, and one containing the infinite tail of components.

3.1 Models Constructed Directly From CRMs

Consider a generic hierarchical model,

\[
M := \sum_{k=1}^{\infty} \mu_k \delta_{\theta_k} \sim \text{CRM}(\nu(d\mu)H(d\theta))
\]

\[Z_{i,k} \sim f(\mu_k), \quad X_i = \sum_{k=1}^{\infty} g(Z_{i,k}, \theta_k) + e_i \tag{1}\]

where \(\nu\) is a measure on \(\mathbb{R}_+\); \(H\) is a measure on the space of parameters \(\Theta\); \(g(\cdot, \cdot)\) is some deterministic transformation such that \(g(0, \theta) = 0\) for all \(\theta \in \Theta\); \(e_i\) is a noise term; and \(f(\cdot)\) is a likelihood that forms a conjugate pair, then the posterior distribution \(P(M^*|Z_i)\) can be decomposed into two CRMs, each with known distribution. If \(K\) is the number of features for which \(\sum_i Z_{i,k} > 0\), the first, \(M_{\leq K}^* = \sum_{k=1}^{K} \mu_k \delta_{\theta_k}\), is a finite measure with fixed-location atoms at locations \(\theta_k : \sum_i Z_{i,k} > 0\). The distribution over the corresponding weights is proportional to \(\nu(d\mu) \prod_{i=1}^{n} f(Z_{i,k}|\mu)\). The second, with infinitely many random-location atoms, has Lévy measure \(\nu(d\mu)H(d\theta)(f(0|\mu))^n\).

Based on Lemma 2, we partition \(M^*\) into a finite CRM \(M_{\leq J}^* = \sum_{k=1}^{J} \mu_k \delta_{\theta_k}\) for some \(J \leq K\), that contains all, or a subset of, the fixed-location atoms; and an infinite CRM \(M_{> K}^*\) that contains the remaining atoms. We use an uncollapsed sampler to sample \(M_{> K}^*|\{Z_{i,k}\}_{k \leq K}\), and then sample \(\{Z_{i,k}\}_{k \notin \mathbb{K}}|X, \{M_{> K}^*\}_{\mathbb{K}}\). Then we use a collapsed sampler to sample the allocations \(Z_{i,k} : k > J\). The size \(J\) should be changed periodically to make sure \(J \leq K\) to avoid explicitly instantiating atoms that are not associated with data. In our experiments, we set \(J = K\) at the beginning of each iteration.

Example 1: Beta-Bernoulli Process. As a specific example, consider the beta-Bernoulli process. Let \(B := \sum_{k=1}^{\infty} \mu_k \delta_{\theta_k} \sim \text{Beta}(\alpha, c, H)\) be a homogeneous beta process [Thibaux and Jordan 2007], and let \(Z_{i,k} \sim \text{Bernoulli}(\mu_k)\). The posterior is given by

\[B^*|Z \sim \text{BetaP}(\frac{\alpha \sum_{k=1}^{K} \mu_k}{c + n}, \frac{\alpha H + \sum_{k=1}^{K} \mu_k \delta_{\theta_k}}{c + \sum_{k=1}^{K} \mu_k})\]

where \(m_k = \sum_{i=1}^{n} Z_{i,k}\). In this case the following lemma helps us in decomposing the posterior distribution:

\[\text{Lemma 2} \quad \text{[Thibaux and Jordan 2007]. If } K \text{ is the number of features where } \sum_i Z_{i,k} > 0 \text{ and } J \leq K, \text{ we can decompose the posterior distribution of beta-} \]

Algorithm 1 Hybrid Beta-Bernoulli Sampler

1: \textbf{while} not converged \textbf{do}
2: \hspace{1em} Select \(J\)
3: \hspace{1em} Sample \(\mu_k \sim \text{Beta}(m_k, n - m_k + c), \forall k \leq J\)
4: \hspace{1em} Sample \(\theta_k \sim p(\theta_k|H, Z, X), \forall k \leq J\)
5: \hspace{1em} for \(i = 1, \ldots, N\) \textbf{do}
6: \hspace{2em} Sample \(\{Z_{i,k}\}_{k=1}^{J}\) according to Eq. 6
7: \hspace{2em} Sample \(\{Z_{i,k}\}_{k=1}^{J}\) according to Eq. 7
8: \hspace{2em} Metropolis-Hastings sample \(Z_{i,k'}\) for \(k' \notin \{k : Z_{i,k} = 1, \sum_j Z_{j,k} = 0\}\).
9: \hspace{1em} end for
10: \textbf{end while}
Bernoulli process as $B^* = B_{<J}^* + B_{>J}^*$ where

$$B_{<J}^* \sim \text{Beta}(\sum_{k=1}^{J} m_k, c + n, \frac{\sum_{k=1}^{J} \mu_k \delta_{k}}{\sum_{k=1}^{J} m_k})$$

$$B_{>J}^* \sim \text{Beta}(\frac{c}{c + n}, c + n, \frac{\alpha + \sum_{k>J} m_k \delta_{k}}{\alpha + \sum_{k>J} m_k})$$

We note that the atom sizes of $B_{<J}^*$ are $\beta(m_k, n - m_k + c)$ random variables. This allows us to split the beta-Bernoulli process into two independent feature selection mechanisms: one with a finite number of currently instantiated features, and one with an unbounded number of features.

The likelihood of a given data point, given the latent variables and other data points (written as $P(X_i|\cdot)$ for space reasons), can be written as

$$p(X_i|\cdot) = \int p(X_i|Z_i, \theta_1, \ldots, \theta_J, \ldots, \theta_K)$$

$$p(\theta_{J+1}, \ldots, \theta_K|Z_{<i}, X_{<i}) d\theta_{J+1} \cdots d\theta_K.$$  

When working with conjugate likelihoods, we can typically evaluate the integral term in Equation 3 analytically (see Ghahramani and Griffiths (2008) for the case of Gaussian prior and linear Gaussian likelihood for $J = 0$). If this is not possible, we can sample $\theta_k$ for $J \geq k \geq K$ as auxiliary variables (Doshi-Velez and Ghahramani 2009).

This formulation of the likelihood, combined with the partitioning of the Bernoulli process described in Equation 2, gives us the hybrid sampler, which we summarize in Algorithm 2. For each data point $X_i$, we sample $Z_{i,k}$ in a three step manner. For $k \leq J$,

$$P(Z_{i,k} = z|\cdot) \propto$$

$$\left\{ \begin{array}{ll}
\mu_k p(X_i|Z_{i,k} = 1, \cdot) & z = 1 \\
(1 - \mu_k) p(X_i|Z_{i,k} = 0, \cdot) & z = 0
\end{array} \right.$$

where $p(X_i|Z_{i,k} = 1, \cdot)$ is given by Equation 3 with $Z_{i,k}$ set to 1. For $k > J$ and $m_k > 0$, we have

$$P(Z_{i,k} = z|Z_{<i}, X_i) \propto$$

$$\left\{ \begin{array}{ll}
m_k p(X_i|Z_{i,k} = 1, \cdot) & z = 1 \\
(n - m_k) p(X_i|Z_{i,k} = 0, \cdot) & z = 0
\end{array} \right.$$

where $p(X_i|Z_{i,k} = 1, \cdot)$ is given by Equation 3 with $Z_{i,k}$ set to 0. Finally, we propose adding Poisson($\alpha/n$) new features, accepting using a Metropolis-Hastings step. Once we have sampled the $Z_{i,k}$, for every instantiated feature $k \leq J$, we sample $\mu_k \sim \text{Beta}(m_k, n - m_k + c)$ and its corresponding parameters $\theta_k \sim p(\theta_k|H, Z, X)$.

We note that similar algorithms can be easily derived for other nonparametric latent feature models such as those based on the infinite gamma-Poisson process (Thibaux and Jordan 2007) and the beta-negative Binomial process (Zhou et al. 2012; Broderick et al. 2018).

### 3.2 Models Based on Transformations of Random Measures

While applying transformations to CRMs means the posterior is no longer directly decomposable, in some cases we can still apply the above general ideas.

**Example 2: Dirichlet Process.** As noted in Section 2, the Dirichlet process with concentration parameter $\alpha$ and base measure $H$ can be constructed by normalizing a gamma process with base measure $\alpha H$. If the Dirichlet process is used as the prior in a mixture model (DPMM), the posterior distribution conditioned on the cluster allocations $Z_1, \ldots, Z_n$, having $K$ unique clusters is again a Dirichlet process:

$$D^*|Z_1, \ldots, Z_n \sim \text{DP}(\alpha + n, \frac{\alpha H + \sum_{k<J} m_k \delta_k}{n + \alpha})$$

where $m_k = \sum_i \delta(Z_i, k)$ and $K$ is the number of clusters with $m_k > 0$. In this case also the following lemma helps us in decomposing the posterior.

**Lemma 3.** Assuming $J$, and $n = \sum_{k<J} m_k$, we can decompose the posterior of $D^*$ as

$$D^*|Z_1, \ldots, Z_n = B^*D_{<J}^* + (1 - B^*)D_{>J}^*$$

where

$$D_{<J}^*|Z_1, \ldots, Z_n \sim \text{DP}(\frac{\sum_{k<J} m_k \delta_k}{n}, \frac{\sum_{k<J} m_k \delta_k}{n + \alpha})$$

$$D_{>J}^*|Z_1, \ldots, Z_n \sim \text{DP}(\alpha + n - \hat{n}, \frac{\alpha H + \sum_{k>J} m_k \delta_k}{\alpha + n - \hat{n}})$$

$$B^* \sim \text{Beta}(\hat{n}, n - \hat{n} + \alpha)$$

**Proof.** This is a direct extension of the fact that the Dirichlet process has Dirichlet-distributed marginals (Ferguson 1973). See Chapter 3 of Ghosh and Ramamoorthi (2003) for a detailed analysis.

We note that the posterior atom weights $(\pi_1, \ldots, \pi_J)$ for the finite component are distributed according to Dirichlet$(m_1, \ldots, m_J)$, and can easily be sampled as part of an uncollapsed sampler. Conditioned on $\{\pi_k, \theta_k : k \leq J\}$ and $B^*$ we can sample the cluster
allocation, $Z_i$ of point $X_i$ as

$$P(Z_i = k|\cdot) \propto \begin{cases} B^* \pi_k f(x_n; \theta_k) & k \leq J \\ \sum_{j=1}^{J} \frac{B^* m_j}{j} f_k(x_n) & J < k \leq K \\ \sum_{j=k}^{\infty} \frac{B^* m_j}{j} f_H(x_n) & k = K + 1 \end{cases}$$

where $f(X_i; \theta_k)$ is the likelihood for each mixing component: $f_k(X_i) = \int_{\Theta} f(X_i; \theta)p(\theta)[X_j : Z_j = k, j \neq i)]d\theta$ is the conditional probability of $x_i$ given other members of the $k$th cluster; and $f_H(x_i) = \int_{\Theta} f(x_i; \theta)H(d\theta)$. This procedure is summarized in Algorithm 2.

**Example 3: Pitman-Yor processes** The Pitman-Yor process [Perman et al., 1992; Pitman and Yor, 1997] is a distribution over probability measures, parameterized by $0 \leq \alpha < 1$ and $\alpha > -\sigma$, that is obtained from a $\sigma$-stable CRM via a change of measure and normalization. Provided $\alpha > 0$, it can also be represented as a Dirichlet process mixture of normalized $\sigma$-stable CRMs (Lemma 22, Pitman and Yor, 1997). This representation allows us to decompose the posterior distribution into a beta mixture of a finite-dimensional Pitman-Yor process and an infinite-dimensional Pitman-Yor process. We provide more details in the supplementary section A.

**Example 4: Hierarchical Dirichlet processes** We can decompose the hierarchical Dirichlet process (HDP, Teh et al., 2006) in a manner comparable to the Dirichlet process, allowing our hybrid sampler to be used on the HDP. For space reasons, we defer discussion to the supplementary section A.

### 4 DISTRIBUTED INFERENCE FOR CRM-BASED MODELS

The sampling algorithms in Section 2 can easily be adapted to a distributed setting, where data are partitioned across several $P$ machines and communication is limited. In this setting, we set $J = K$ (i.e. the number of currently instantiated features) after every communication step. We instantiate the finite measure ($M_{\leq J}$ in the case of CRMs, $D_{\leq J}$ for DPMMs), with globally shared atom sizes and locations, on all processors.

We then randomly select one out of $P$ processors by sampling $P^{*} \sim \text{Uniform}(1, \ldots, P)$. On the remaining $P - 1$ processors, we sample the allocations $Z_i$ using restricted Gibbs sampling (Neal, 2000), enforcing $Z_{i,k} = 0$ for $k > J$. When working with DPMMs, this means that we only need to calculate cluster probabilities that depend on the instantiated measure $D_{\leq J}$. In the CRM case, it means that we avoid the integral in Equation 8 and hence avoid any dependence on $Z_{i}$ or $X_{-i}$ when conditioning on $M_{\leq J}$. In both cases, this means that we do not need knowledge of the feature allocations on other processors, and can sample the $Z_i$ independent of each other.

On the remaining processor $P^{*}$, we sample the $Z_i$ using unrestricted Gibbs sampling. Let $P^{*}$ be the set of indices of data points on processor $P^{*}$. Since we know that $Z_{j,k} = 0$ for all $j \notin P^{*}$, then we can calculate the full sufficient statistics for features $k > J$ without knowledge of data or latent features on other processors. While $P(Z_i|\cdot)$ does depend on $\{Z_i, X_j : j \in P\}$, it is independent of $\{Z_j, X_j : j \notin P\}$ conditioned on $D_{\leq J}$ (or $M_{\leq J}$) plus the fact that $Z_{j,k} = 0$ for all $j \notin P^{*}$, so can be calculated without further information from the other processors.

At each global step, we gather the sufficient statistics from all instantiated clusters – from both the finite component $M_{\leq J}$ / $D_{\leq J}$ and the infinite component $M_{> J}$ / $D_{> J}$ – and sample parameters for those clusters. We then create a new partition, redefining $J$ as the current number of instantiated component parameters. In the case of the DPMM, we also resample $B \sim \text{Beta}(N, \alpha)$. We summarize the distributed algorithm for the special cases of the beta-Bernoulli process and the DPMM in Algorithms 3 and 4, and for PYMM in algorithm 6 in supplementary.

#### 4.1 Warm-start Heuristics

In our distributed approach, only $1/P$ of the data points eligible to start a new cluster or feature, mean-
Algorithm 4 Distributed DPMM Sampler

1: procedure LOCAL({X_i, Z_i})
   ▷ Global variables J, P, \{\theta_k, \pi_k\}_{k=1}^D, B^*
2:   if P = P* then
3:     Sample Z_i according to (7)
4:   else
5:     P(Z_i = k) \propto \pi_k f(X_i; \theta_k)
6:   end if
7: end procedure

8: procedure GLOBAL({X_i, Z_i})
9:   Gather cluster counts m_k and parameter sufficient statistics \Psi_k from all processors.
10:  Sample B^* \sim \text{Beta}(n, \alpha)
11:  Let J be the number of instantiated clusters.
12:  Sample (\pi_1, \ldots, \pi_J) \sim \text{Dir}(m_1, \ldots, m_J)
13:  For k \in \{1, \ldots, J\}, sample \theta_k \sim \text{p}(\theta_k|\Psi_k, H).
14:  Sample P^* \sim \text{Uniform}(1, \ldots, P)
15: end procedure

5 EXPERIMENTAL EVALUATION

While our primary contribution is in the development of distributed algorithms, we first consider, in Section 5.1, the performance of the hybrid algorithms developed in Section 3 in a non-distributed setting. We show that this performance extends to the distributed setting, and offers impressive scaling, in Section 5.2.

5.1 Evaluating the Hybrid Sampler

We begin by considering the performance of the hybrid samplers introduced in Section 3 in a non-distributed setting. For this, we focus on the Dirichlet process, since there exist a number of collapsed and uncollapsed inference algorithms; we expect similar results under other models.

We compare the hybrid sampler of Algorithm 2 with a standard collapsed Gibbs sampler and an uncollapsed sampler based on Algorithm 8 of Neal (2000). Algorithm 8 collapses occupied clusters and instantiates a subset of unoccupied clusters; we modify this to instantiate the atoms associated with unoccupied clusters. Concretely, at each iteration, we sample weights for the K instantiated clusters plus U uninstantiated clusters as (\pi_1, \ldots, \pi_K, \pi_{K+1}, \ldots, \pi_{K+U}) \sim \text{Dir}(m_1, \ldots, m_K, \frac{J}{K+U}, \ldots, \frac{J}{K+U}) and sample locations for the uninstantiated clusters from the base measure H. Note that this method can be distributed easily.

Figure 1 shows convergence plots for the three algorithms. The data set is a D dimensional synthetic data set consisting of 100 observations of Gaussian mixtures with 2 true mixture components centered at 5 \times \{1\}^D and -5 \times \{1\}^D with an identity covariance matrix.

While the three algorithms perform comparably on low-dimensional data, as the dimension increases the performance of the uncollapsed sampler degrades much more than the collapsed sampler. This is because in high dimensions, it is unlikely that a proposed parameter will be near our data, so the associated likelihood of any given data point will be low. This is in contrast to the collapsed setting, where we integrate over all possible locations. While the hybrid method performs worse in high dimensions than the collapsed method, it outperforms the uncollapsed method.

The synthetic data in Figure 1 has fairly low-dimensional structure, so we do not see negative effects due to the poor mixing of the collapsed sampler. Next, we evaluate the algorithms on the CIFAR-100 dataset (Krizhevsky 2009). We used PCA to reduce the dimension of the data to between 8 and 64, and plot the test set log likelihood over time in Figure 2. Each marker represents a single iteration. We see that the uncollapsed sampler requires more iterations to con-
verge than the collapsed sampler; however since each iteration takes less time, in some cases the wall time to convergence is slower. The hybrid method has comparable iteration time to the collapsed, but, in general, converges faster. We see that, even without taking advantage of parallelization, the hybrid method is a compelling competitor to pure-collapsed and pure-uncollapsed algorithms.

5.2 Evaluating the Distributed Samplers

Here, we show that the distributed inference algorithms introduced in Section 4 allow inference in BNP models to be scaled to large datasets, without sacrificing accuracy. We focus on two cases: the beta-Bernoulli process (Algorithm 3) and the Dirichlet process (Algorithm 4).

5.2.1 Beta-Bernoulli Process

We evaluate the beta-Bernoulli sampler on synthetic data based on the “Cambridge” dataset, used in the original IBP paper (Griffiths and Ghahramani, 2011), where each datapoint is the superposition of a randomly selected subset of four binary features of dimension 36, plus Gaussian noise with standard deviation 0.5. We model this data using a linear Gaussian likelihood, with \( Z \sim \text{Beta-Bern}(\alpha, 1) \), \( A_k \sim \text{Normal}(0, \sigma_A^2 \mathbf{1}) \), \( \mathbf{X_n} \sim \text{Normal} \left( \sum_k z_{nk} A_k, \sigma_X^2 \mathbf{1} \right) \).

We initialized to a single feature, and ran the hybrid sampler for 1,000 total observations with a synchronization step every 5 iterations, distributing over 1, 4, 8, 16, 32, 64 and 128 processors.

We first evaluate the hybrid algorithm under a “cold start”, where only one processor is allowed to introduce new features for the entire duration of the sampler. In the top left plot of Figure 3, we see that the cold start results in slow convergence of the test set log likelihood for large numbers of processors. We can see in the bottom left plot of Figure 3 that the number of features grows very slowly, as only \( 1/P \) processors are allowed to propose new features in the exact setting.

Next, we explore warm-start initialization, as described in Section 4.4. For the first one-eighth of the total number of MCMC iterations, all processors can propose new features; after this we revert to the standard algorithm. The top central plot of Figure 3 shows predictive log likelihood over time, and the bottom central plot shows number of features. We see that convergence is significantly improved relative to the cold-start experiments. Since we revert to the asymptotically correct sampler, the final number of features is generally close to the true number of features, 4.

Additionally, we see that convergence rate increases monotonically in the number of processors.

Next, we allowed all processors to propose new features for the entire duration (“always-hot”). This setting approximately replicates the behavior of the parallel IBP sampler of Doshi-Velez et al. (2009b). In the top right plot of Figure 3, we can see that all experiments roughly converge to the same test log likelihood. However, the number of features introduced (bottom right plot) is much greater than the warm start experiment, grows with the number of processors. Moreover, the difference in convergence rates between processors is not as dramatic as in the warm-start trials.

Next, we demonstrate the scalability of our distributed algorithm on a massive synthetic example, showing it can be used for large-scale latent feature models. We generate one million “Cambridge” synthetic data points, as described for the previous experiments, and distribute the data over 256 processors. This experiment represents the largest experiment ran for a beta-Bernoulli process algorithm (the next largest being 100,000 data points, in Doshi-Velez et al. 2009b). We limited the sampler to run for one day and completed 860 MCMC iterations. In Figure 4, we see in the test set log likelihood traceplot that we can converge to a local mode fairly quickly under a massive distributed

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1. Our code is available online at https://github.com/michaelzhang01/hybridBNP
2. See Figure 6 in the supplement for more details
3. Note that BNP models are not guaranteed to find the correct number of features in the posterior; see Miller and Harrison (2013).

Figure 2: Test log-likelihood with time on CIFAR-100 dataset as the dimensionality increases from 8 to 64.
Figure 3: **Top row left to right:** Test set log likelihood (y-axis) on synthetic data without warm-start initialization, with warm start, and with all processors on. The x-axis represents CPU wall time in seconds, on a log scale. **Bottom row left to right** number of features over iteration with cold-start, warm-start, and all processors introducing features. Y-axis represents number of instantiated features.

Figure 4: Test set log likelihood trace plot for a million observation “Cambridge” data set.

Figure 5: Comparison of CIFAR-100 test log-likelihood with time, with baseline methods (**left**) and with varying number of processor (**right**)

6 Conclusion

We have proposed a general inference framework for a wide variety of BNP models. We use the inherent decomposability of the underlying completely random measures to partition the latent random measures into a finite-dimensional component that represents the majority of the data, and an infinite-dimensional component that represents mostly uninstantiated tail. This allows us to take advantage of the inherent parallelizability of the uncollapsed sampler on the finite partition and the better performance of the collapsed sampler for proposing new components. Thus the proposed hybrid inference method can be easily distributed over multiple machines, providing provably correct inference for many BNP models. Experiments show that, for both the DP and the beta-Bernoulli process, our proposed distributed hybrid sampler converges faster than the comparison methods.
References


