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(Article begins on next page)

A New Estimator of the Discovery Probability

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Summary. Species sampling problems have a long history in ecological and biological studies and a number of issues, including the evaluation of species richness, the design of sampling experiments, and the estimation of rare species variety, are to be addressed. Such inferential problems have recently emerged also in genomic applications, however, exhibiting some peculiar features that make them more challenging: specifically, one has to deal with very large populations (genomic libraries) containing a huge number of distinct species (genes) and only a small portion of the library has been sampled (sequenced). These aspects motivate the Bayesian nonparametric approach we undertake, since it allows to achieve the degree of flexibility typically needed in this framework. Based on an observed sample of size n, focus will be on prediction of a key aspect of the outcome from an additional sample of size m, namely, the so-called discovery probability. In particular, conditionally on an observed basic sample of size n, we derive a novel estimator of the probability of detecting, at the (n + m + 1)th observation, species that have been observed with any given frequency in the enlarged sample of size n + m. Such an estimator admits a closed-form expression that can be exactly evaluated. The result we obtain allows us to quantify both the rate at which rare species are detected and the achieved sample coverage of abundant species, as m increases. Natural applications are represented by the estimation of the probability of discovering rare genes within genomic libraries and the results are illustrated by means of two expressed sequence tags datasets.

KEY WORDS: Bayesian nonparametrics; Gibbs-type priors; Rare species discovery; Species sampling models; Two-parameter Poisson–Dirichlet process.

1. Introduction

Species sampling problems have a long history in ecological and biological studies. Suppose data are recorded from a population whose statistical units belong to different species. Therefore, data will consist of species labels and their corresponding frequencies. Based on a sample, various interesting predictive issues concerning the composition of the population arise. Such problems have regained popularity in recent years due to their frequent appearance in genomic applications, which are characterized by very large populations (genomic libraries) containing a huge number of distinct species (genes) and only a small portion of the library has been sampled (sequenced). Recently, a Bayesian nonparametric method, particularly suited to such genomic contexts, has been proposed and implemented in Lijoi, Mena, and Prünster (2007a, 2007b). This approach is based on the randomization of the unknown proportions p_i of the individuals belonging to the species iin the whole population, for $i \geq 1$. It is further assumed that the recorded data X_1, \ldots, X_n are part of an exchangeable sequence $(X_n)_{n\geq 1}$. In this case, it is well-known that by de Finetti's representation theorem, $(X_n)_{n\geq 1}$ can be characterized by a hierarchical model, namely, the X_n 's as a random sample from some distribution P and a prior Π on P, that is,

$$X_i | \tilde{P} \stackrel{\text{iid}}{\sim} \tilde{P},$$

 $\tilde{P} \sim \Pi.$ (1)

Moreover, \tilde{P} is a discrete random probability measure $\tilde{P} = \sum_{i \geq 1} w_i \, \delta_{Y_i}$, with $\sum_{i \geq 1} w_i = 1$ almost surely, belonging to the class of Gibbs-type priors (Gnedin and Pitman, 2006). Note that δ_c is used to denote the unit mass at point c, and the Y_i 's are independent and identically distributed from some distribution P_0 . Henceforth, we shall adhere to common practice and assume P_0 nonatomic thus implying that any two Y_i and Y_i , with $i \neq j$, are different with probability 1. On the basis of this fact. one can equivalently use the Y_i 's or the positive integers $\{1, 2, \ldots\}$ to label different species in the population whose random proportions are modeled through the w_i 's. It is worth lingering on this point a little further and, given its relevance, clarify it. To this end, introduce an auxiliary integer-valued sequence $(\xi_n)_{n\geq 1}$ such that $\mathbb{P}[\xi_n=j\mid P]=w_j$, for any n and j, and note that (1) corresponds to assuming that $X_i = Y_{\xi_i}$. Hence, the X_n 's can be interpreted as the observed species labels since, due to the nonatomic nature of P_0 , any two data points X_i and X_j , for $i \neq j$, differ if and only if ξ_i and ξ_j do. Moreover, one has that $\mathbb{P}[\xi_i = \xi_j] > 0$, for any $i \neq j$, and this entails that the *i*th and the *j*th observation may reveal the same species with positive probability. Accordingly, the *n* observations X_1, \ldots, X_n forming a basic sample may feature ties. If $K_n = j$ is the number of different species (labels) detected in the basic sample with respective frequencies n_1, \ldots, n_j , an additional potential and unobserved sample of size $m, X_{n+1}, \dots, X_{n+m}$, is considered.

Focus will be on prediction of a key aspect of the outcome of the additional sample, namely, the so-called discovery probability, which coincides with the probability of sampling certain types of genes in further sequencing. The discovery probability represents a natural tool for a quantitative assessment of qualitative concepts such as species richness, rare species variety, sample coverage, and optimal design of sampling experiments. A first partial result in this direction is obtained in Lijoi et al. (2007a), where an estimator for the probability that the (n+m+1)th observation leads to discover a new species without actually observing the additional m-size sample is provided.

More specifically, an important inferential goal in ecological and biological studies is the evaluation of the probability that further sampling reveals: (i) new species; (ii) species that appear only once in the observed sample, the so-called unique species; and (iii) rare species, where by rare species one refers to species whose frequency is below a certain threshold τ . In ecology, for instance, conservation of biodiversity is a fundamental theme and it can be formalized in terms of the proportion of species whose frequency is greater than a specified threshold of abundance. Indeed, any form of management on a sustained basis requires a certain proportion of sufficiently abundant species, the so-called breeding stock. See Magurran (2003), and references therein, for a comprehensive and stimulating survey on measurement of biodiversity, conservation of populations, commonness, and rarity of species. On the other hand, in genomics, rare genes are a fundamental issue in several problems, the most evident being that they are often associated with deleterious diseases. See, e.g., Laird and Lange (2010).

In this article, we provide an estimator for detecting a species of any given frequency at the (n+m+1)th observation, therefore addressing the issues (ii) and (iii) above and improving remarkably, both in terms of theory and applications potential, on the results in Lijoi et al. (2007a) concerning problem (i) only. Specifically, based on a sample of size n, we will provide, for any frequency $k=0,\ldots,n+m$ and additional unobserved sample size $m\geq 0$, an explicit estimator for the probability that the (n+m+1)th observation coincides with a species whose frequency, within the sample of size n+m, is exactly k. In the sequel, we refer to such a probability as m-step k-discovery probability or, in short, [m:k]-discovery. Expressed in terms of the unknown species proportions in the whole population, p_i for $i\geq 1$, the determination of the [m:k]-discovery corresponds to estimating

$$U_{n+m}(k) := \sum_{i \ge 1} p_i 1_{\{k\}}(N_{i,n+m}), \tag{2}$$

where $N_{i,n+m}$ is the frequency with which the *i*th species is recorded in the enlarged sample of size n+m and clearly $U_{n+m}(0)$ represents the proportion of yet unobserved species or, equivalently, the probability of observing a new species.

For m = 0, such a problem has already been tackled in the literature. In particular, Good (1953) proposed a popular estimator for (2) of the form

$$\check{U}_{n+0}(k) = (k+1)\frac{l_{k+1}}{n},\tag{3}$$

where l_r is the number of species in the sample with frequency r. Such an estimator is sometimes referred to as $Turing\ estimator$. In Mao and Lindsay (2002), an interesting new moment-based derivation of (3) is provided. Note that $1-\check{U}_{n+0}(0)$ estimates the $sample\ coverage\$ that is the proportion of distinct species present in the observed sample. On the other hand, Good and Toulmin (1956) and Mao (2004) faced the general case of $m \geq 0$, however, with the restriction k=0, which corresponds only to the discovery of a new species. The resulting estimator is known as $Good-Toulmin\ estimator$. Further references addressing related issues are Chao (1981); Starr (1979); Clayton and Frees (1987); Guindani and Müller (2010); Sepúlveda et al. (2010); Barger and Bunge (2010)).

Our contribution therefore fills in an important gap since it allows to deal with the cases of both the additional sample size m and the frequency k to be any desired integer. As a byproduct, also the corresponding coverage estimators are obtained. Moreover, the coherent probabilistic structure of Bayesian nonparametric modeling allows to avoid some possible drawbacks related to the estimator (3), which will be detailed in the sequel by comparing the two estimators.

The structure of the article is as follows. In Section 2, we describe the Bayesian nonparametric model and provide the main result. A completely explicit expression of the new estimator is then provided in the two-parameter Poisson–Dirichlet process case. Section 3 contains an illustration dealing with genomic expressed sequence tag (EST) datasets. The Matlab code for computing the proposed estimators and proofs of our results are available in a Web Appendix as Supplementary Material.

2. Estimation of the m-Step k-Discovery

As outlined in Section 1, the main aim of the present contribution is the estimation of the [m:k]-discovery. The deduced Bayesian estimator is based on a nonparametric model for the unknown distribution of species labels in (1) and, more importantly, can be exactly evaluated. The latter seems to be an attractive feature since it makes our approach readily implementable by practitioners. Let us first provide a quick introduction of the nonparametric process prior we resort to, along with some known results. We will then state the main result.

2.1 Preliminaries

It is apparent, from (1), that the prior Π can be defined as a distribution on the space of all probability measures for the observables $(X_n)_{n\geq 1}$. We now consider the case where the support of Π does not degenerate on a finite-dimensional space so that the model is nonparametric, and Π is concentrated on the set of all discrete probability distributions for the X_i 's. Such a specification is consistent with the nature of our data, since the process of observation leads to detecting multiple records for the same species. In other terms, if we let Π select discrete probability measures with probability 1, then $\mathbb{P}[X_i = X_j] > 0$ for $i \neq j$ which is what we aim at. In this setup, a sample X_1, \ldots, X_n is partitioned into K_n distinct values, i.e., species, $X_1^*, \ldots, X_{K_n}^*$ with vector of respective frequencies $\mathbf{N}_n = (N_{1,n}, \ldots, N_{K_n,n})$ being such that $\sum_{i=1}^{K_n} N_{i,n} = n$. In the sequel, we consider a broad class of discrete random probability measures defined as $\tilde{P} = \sum_{i \geq 1} w_i \, \delta_{Y_i}$, where the

nonnegative weights w_i 's sum up to 1, i.e., $\sum_i w_i = 1$ (almost surely), and are independent from the locations Y_i 's. The Y_i 's, in turn, are i.i.d. from some nonatomic distribution P_0 . It is further supposed that \tilde{P} is such that the probability distribution of the partition of a sample into K_n distinct values with respective frequencies N_n is of the form

$$\mathbb{P}[K_n = j, \mathbf{N}_n = (n_1, \dots, n_j)] = V_{n,j} \prod_{i=1}^{j} (1 - \sigma)_{n_i - 1}, \quad (4)$$

where $(a)_n = a(a+1) \cdots (a+n-1)$ stands for the *n*th ascending factorial of a, with $(a)_0 \equiv 1$, σ is some parameter in (0,1), and the nonnegative weights $\{V_{n,j}: n \geq 1, 1 \leq j \leq n\}$ satisfy the following forward recursion $V_{n,j} = V_{n+1,j+1} + (n - 1)$ $j\sigma$) $V_{n+1,j}$. Such random probability measures and their corresponding distribution Π , which represents our prior distribution, are said of Gibbs-type and were introduced by Gnedin and Pitman (2006). The parameter σ that characterizes Π influences the partition structure of the data: indeed, as noted in Lijoi, Mena, and Prünster (2007c), it regulates a reinforcement mechanism that determines the concentration of the observations in the different groups forming the partition. In summary, large values of σ tend to favor partitions with a larger number of clusters, i.e., of different observed species: most of the clusters (species) display very low frequencies, whereas just a few clusters have very large abundances. Hence, a large value of σ reflects a prior opinion according to which a large number of species will be detected though only a few of them with high frequencies, which is a common situation in genomic datasets like the ones we are going to consider in Section 3.

In various applications, it is useful to describe the probability distribution of the random partition induced by Π in terms of the vector $\mathcal{L}^{(n)} := (\mathcal{L}_{1,n}, \ldots, \mathcal{L}_{n,n})$, where $\mathcal{L}_{i,n}$ stands for the number of distinct species in the basic sample, of size n, with frequency i. Hence, (4) can be rewritten as

$$\mathbb{P}[\mathcal{L}^{(n)} = (l_1, \dots, l_n)] = V_{n,j} \ n! \prod_{i=1}^n \left\{ \frac{(1-\sigma)_{i-1}}{i!} \right\}^{l_i} \frac{1}{l_i!}, (5)$$

for any vector of nonnegative integers (l_1, \ldots, l_n) such that $l_i \geq 0$ for any $i, \sum_{i=1}^n i l_i = n$ and $\sum_{i=1}^n l_i = j$. The probability distribution in (5) is termed as Gibbs-type sampling formula and it includes as a special case the well-known Ewens sampling formula (Ewens, 1972) and Ewens-Pitman sampling formula (Pitman, 1995).

Gibbs-type priors lead to predictive distributions, given a sample of size n (X_1, \ldots, X_n) , featuring $K_n = j$ distinct species X_1^*, \ldots, X_j^* with frequencies n_1, \ldots, n_j , having the following simple structural form, which can be deduced from (4),

$$\mathbb{P}[X_{n+1} \in A \mid X_1, \dots, X_n] = \frac{V_{n+1,j+1}}{V_{n,j}} P_0(A) + \frac{V_{n+1,j}}{V_{n,j}} \sum_{i=1}^{j} (n_i - \sigma) \delta_{X_i^*}(A).$$
(6)

Various noteworthy priors fall within the class of Gibbstype priors, according as to the specific form of the weights $V_{n,j}$. For example, the Dirichlet process (Ferguson, 1973) with parameter measure θP_0 is recovered when $V_{n,j} =$ $\theta^j/(\theta)_n$, whereas the two-parameter Poisson–Dirichlet process (Pitman, 1995) is obtained with $V_{n,j} = \prod_{i=1}^{j-1} (\theta+i\sigma)/(\theta+1)_{n-1}$. Also, the normalized inverse Gaussian process (Lijoi, Mena, and Prünster, 2005) belongs to this class, with the $V_{n,j}$'s expressed in terms of a linear combination of incomplete gamma functions. Another instance of Gibbs-type prior is the normalized generalized gamma process that has been used in Lijoi et al. (2007c) for hierarchical mixture modeling and, more recently, in Kolossiatis, Griffin, and Steel (2011) for modeling overdispersion in count data.

In Lijoi et al. (2007a), one can find a few results concerning Gibbs-type priors in view of their application to species sampling problems. For example, an estimator of the sample coverage can be easily deduced from the predictive distributions (6) so that

$$1 - \widehat{U}_n(0) := \widehat{C}_n = \frac{V_{n+1,j}}{V_{n,j}} (n - j\sigma).$$
 (7)

Moreover, they determine an estimator for the [m:0]-discovery given by

$$\widehat{U}_{n+m}(0) = \sum_{k=0}^{m} \frac{V_{n+m+1,j+k+1}}{V_{n,j}} \frac{\mathcal{C}(m,k;\sigma,-n+j\sigma)}{\sigma^{k}}$$
(8)

with $C(m,k;\sigma,-n+j\sigma)=(k!)^{-1}\sum_{r=0}^k (-1)^r {k\choose r}$ $(n-\sigma(r+j))_m$ being the noncentral generalized factorial coefficient; see Charalambides (2005) for details and properties. The corresponding estimator of the sample coverage, given an observed sample of size n and an additional unobserved sample of size m, \widehat{C}_{n+m} , is then obtained as $1-\widehat{U}_{n+m}(0)$. The estimators (8), and consequently, \widehat{C}_{n+m} can then be specialized to various particular cases by plugging in the corresponding $V_{n,j}$'s.

2.2 Main Results

As anticipated, our main goal is the estimation of $U_{n+m}(k)$, for any value of m and k, conditional on an observed sample of size n featuring j distinct species. We first state a simple result yielding a nonparametric Bayes estimator of $U_{n+0}(k)$ for any integer k. Such an estimation involves the one-step predictive distribution since it amounts to evaluating

$$\mathbb{P}[X_{n+1} \in \mathcal{G}_{k,n} \mid X_1, \dots, X_n], \tag{9}$$

where $\mathcal{G}_{k,n} = \{X_i^* : N_{i,n} = k\}$ is the set of all distinct species that have appeared k times in the sample X_1, \ldots, X_n . For k = 0, it is apparent that (9) coincides with the probability that the (n + 1)th observation reveals a new species and is readily available from the predictive distribution (6), i.e., $V_{n+1,j+1}/V_{n,j}$. See also (7). Hence, we can focus on $k \geq 1$ and provide an explicit expression of the predictive probability (9).

Theorem 1. If $(X_n)_{n\geq 1}$ is an exchangeable sequence with Gibbs-type prior Π in (1)

$$\widehat{U}_{n+0}(k) = \frac{V_{n+1,j}(k-\sigma)}{V_{n,j}} l_k,$$
(10)

where $l_k \geq 0$ is the number of distinct species with frequency k observed within the sample (X_1, \ldots, X_n) .

It is worth noting that (10) implies that $(K_n, \mathcal{L}_{k,n})$ is sufficient for estimating the [0:k]-discovery $U_{n+0}(k)$.

Remark 1. The estimator $\widehat{U}_{n+0}(k)$ represents a Bayesian nonparametric counterpart to the popular Turing estimator $U_{n+0}(k)$ recalled in (3) and used in a frequentist approach. The most notable difference between the two estimators can be traced back to the different frequency count used. Indeed, $\check{U}_{n+0}(k)$ in (3) depends only on the frequency l_{k+1} of species that appeared k+1 times in the basic sample. This seems, in our opinion, counterintuitive since l_k should be the main ingredient for the estimation of the discovery probability of species appearing k times. And the Bayesian estimator $U_{n+0}(k)$ in (10) perfectly adheres to such an intuition. Note that if there are no species appearing k+1 times (i.e., $l_{k+1} = 0$) in the original sample of size n, then one would estimate as zero the probability of detecting a species with frequency k at the (n+1)th observation; and this regardless of how many species with frequency k one has observed in the basic sample. This is not the case for the Bayesian nonparametric estimator (10).

In many applications, one is interested, rather than in identifying the [0:k]-discovery, in evaluating the chances of observing species that can be considered as rare, namely, the sum of those not yet observed and of those having frequency below a certain threshold τ . In this case, an estimator of $\sum_{k=0}^{\tau} U_{n+0}(k)$ is

$$\widehat{U}_{n+0,\tau} = \sum_{k=0}^{\tau} \widehat{U}_{n+0}(k),$$

and it depends on the vector of summary statistics $(K_n, \mathcal{L}_{1,n}, \dots, \mathcal{L}_{\tau,n})$.

Remark 2. It is worth pointing out that the Bayesian nonparametric estimator of the sample coverage (7) can also be recovered from (10). Indeed, by definition, one has $\widehat{C}_n = 1 - \widehat{U}_{n+0}(0) = \sum_{k=1}^n \widehat{U}_{n+0}(k)$ and the expression in (7) can be determined by resorting to Theorem 1 and by recalling that $\sum_{k=1}^n k l_k = n$ and $\sum_{k=1}^n l_k = j$. Another related quantity of interest can be determined from Theorem 1, namely, an estimated abundant species coverage, where by abundant we mean species that are not rare. This is given by the proportion of species with frequency larger than τ represented in the sample, i.e., $\widehat{C}_n^{abund} = 1 - \sum_{k=0}^{\tau} \widehat{U}_{n+0}(k)$.

Let us now consider the case where $m \ge 1$: we are then going to evaluate the conditional probability, given the sample X_1, \ldots, X_n , that the (n+m+1)th observation displays a species observed k times within $X_1, \ldots, X_n, X_{n+1}, \ldots, X_{n+m}$. This implies that we wish to estimate

$$\mathbb{P}[X_{n+m+1} \in \mathcal{G}_{k,n+m} | X_1, \dots, X_n, X_{n+1}, \dots, X_{n+m}]. \tag{11}$$

Assuming a squared loss function, we estimate (11) by evaluating the expectation with respect to the conditional distribution of the unobserved portion of the sample, X_{n+1}, \ldots, X_{n+m} , given the sample that has been actually observed, X_1, \ldots, X_n , i.e., the m-step ahead predictive distribution. In other terms,

the desired estimator $\widehat{U}_{n+m}(k)$ is obtained by minimizing

$$\mathbb{E}\bigg\{\left.(\mathbb{P}[X_{n+m+1}\in\mathcal{G}_{k,n+m}\mid X_1,\ldots,X_n,X_{n+1},\ldots,X_{n+m}]-U)^2\mid \\ X_1,\ldots,X_n\right\}$$

with respect to U and coincides with $\mathbb{P}[X_{n+m+1} \in \mathcal{G}_{k,n+m} | X_1,\ldots,X_n]$. For k=0, the [m:0]-discovery coincides with the probability of discovering a new species at the (n+m+1)th step given the basic sample of size n, which has been derived in Lijoi et al. (2007a) and is recalled in (7). The case of k > 1 is dealt with by the following result.

Theorem 2. If $(X_n)_{n\geq 1}$ is an exchangeable sequence with Gibbs-type prior Π in (1), a Bayesian nonparametric estimator of the [m:k]-discovery coincides with

$$\widehat{U}_{n+m}(k) = \sum_{i=1}^{k} l_i (i - \sigma)_{k+1-i} \binom{m}{k-i} Q_{m,k}^{(n,j)}(i, 0, i - \sigma) + \sigma (1 - \sigma)_k \binom{m}{k} Q_{m,k}^{(n,j)}(0, 1, 0),$$
(12)

where

$$egin{aligned} Q_{m,l}^{(n,j)}(lpha,eta,\gamma) := \sum_{k=eta}^{m-l+lpha} rac{V_{n+m+1,j+k}}{V_{n,j}} \ & imes rac{\mathcal{C}(m-l+lpha,k-eta;\sigma,-n+j\sigma+\gamma)}{\sigma^k} \end{aligned}$$

and $l_i \geq 0$ is the number of species observed with frequency i in the sample (X_1, \ldots, X_n) .

The analytic form of the estimator in (12) implies that the vector of the number of species and frequency counts $(K_n, \mathcal{L}_{1,n}, \ldots, \mathcal{L}_{k,n})$ is sufficient for estimating the [m:k]-discovery. It is worth remarking that the novel estimator in Theorem 2 does not have any counterpart both in the frequentist and in the Bayesian frameworks. As recalled in the introduction, frequentist estimators for m-step k-discovery exist only when either m = 0 and $k \geq 0$ or $m \geq 1$ and k = 0: the former corresponds to the Turing estimator displayed in (3), whereas the latter corresponds to the so-called Good-Toulmin estimator (Good and Toulmin, 1956), which, however, becomes unstable for $m \geq 2n$ (see, e.g., Lijoi et al. (2007b) for an illustration).

Obviously, from (12), one can also deduce estimators for cumulative [m:k]-discoveries, which allow to evaluate the probability of observing a rare species at the (n+m+1)th draw from the population. Hence, for an abundance threshold τ , one just needs to evaluate

$$\widehat{U}_{n+m,\tau} = \sum_{k=0}^{\tau} \widehat{U}_{n+m}(k).$$
 (13)

This implies that $(K_n, \mathcal{L}_{1,n}, \dots, \mathcal{L}_{\tau,n})$ is sufficient for estimating the [m:k]-discovery probability for species with frequencies less than or equal to τ , conditionally on the basic sample. An important application of the estimator in (13) is related to sample size determination. Indeed, one is often willing to

determine the size of the additional sample m as the smallest integer m^* for which $\widehat{U}_{n+m^*,\tau}$ is not smaller than a certain threshold κ . See, e.g., Christen and Nakamura (2003). The rationale is that the smaller the probability of sampling rare species, the less informative is the sampling procedure since, in this case, it mainly consists of reobserving species with large abundances. Hence, a further enlargement of the sample size beyond m^* would not yield enough relevant information compared the cost of further sampling. For instance, in applications, one often sets such a threshold to $\kappa \in \{0.1, 0.2, 0.5\}$.

The result in Theorem 2 allows to obtain a decomposition of the sample coverage estimator, given an observed sample of size n and an additional unobserved sample of size m, \widehat{C}_{n+m} . Indeed, one has that

$$\widehat{C}_{n+m} = \sum_{k=1}^{n+m} \widehat{U}_{n+m}(k), \tag{14}$$

which provides an alternative derivation of \widehat{C}_{n+m} w.r.t. the derivation based on (8). Similarly to the case m=0, based on Theorem 2, one can deduce an estimator for abundant species coverage after sampling n+m species, which is given by the proportion of species with frequency larger than τ represented in both the observed basic and unobserved additional sample, i.e.,

$$\widehat{C}_{n+m}^{abund} = 1 - \sum_{k=0}^{\tau} \widehat{U}_{n+m}(k).$$

2.3 The Two-Parameter Poisson-Dirichlet Process

As already mentioned, a prominent special case of Gibbs type prior is the two-parameter Poisson–Dirichlet process. The system of predictive distributions it induces for the data in (1) can be deduced from (6) by recalling the simple form of its weights $V_{n,j}$'s, namely, $V_{n,j} = \prod_{i=1}^{j-1} (\theta + i\sigma)/(\theta + 1)_{n-1}$, which leads to

$$\mathbb{P}[X_{n+1} \in A \mid X_1, \dots, X_n] = \frac{\theta + k\sigma}{\theta + n} P_0(A) + \frac{1}{\theta + n} \sum_{i=1}^{j} (n_i - \sigma) \, \delta_{X_i^*}(A)$$
(15)

for any set A. The vector of parameters is such that $\sigma \in (0,1)$ and $\theta > -\sigma$ and we shall, henceforth, adopt the concise notation $PD(\sigma, \theta; P_0)$ for denoting such a prior. Its specification is particularly appealing: besides being analytically tractable for addressing various inferential issues in a Bayesian setting, it also represents a very flexible model, as detailed, e.g., in Lijoi et al. (2007a). For these reasons, it has become an increasingly popular prior in several applications such as, e.g., mixture models (Ishwaran and James, 2001), linguistics (Teh, 2006), species sampling models (Navarrete, Quintana, and Müller, 2008), information retrieval in document modeling (Teh and Jordan, 2010), and survival analysis (Jara et al., 2010). And while its use can be considered as routine in Bayesian nonparametrics modeling, its effective implementation has become easily accessible also to practitioners due to the availability of an efficient R package that embodies various MCMC algorithms with the two-parameter Poisson–Dirichlet process. See Jara et al. (2011) for details.

Within the present framework, Lijoi et al. (2007b) and Favaro et al. (2009) provided closed-form expressions for $PD(\sigma,\theta;P_0)$ -estimators of: (i) the expected number of new species $K_m^{(n)}$ that will be observed from a further sampling of size m and (ii) the [m:0]-discovery as described in (8). However, as mentioned before, it is important to extend the analysis to rare species, whose quantification is often a major goal not only in EST analysis but also in different genomic applications such as cap analysis gene expression (CAGE) (see, e.g., Valen (2009)). Based on the general results of Section 2.2, here, we provide completely explicit estimators of the [m:k]-discovery in this noteworthy particular case.

Proposition 3: If $(X_n)_{n\geq 1}$ is an exchangeable sequence with Π in (1) being a Poisson-Dirichlet process with parameters (σ,θ) , then

$$\widehat{U}_{n+0}(k) = \frac{k-\sigma}{\theta+n} l_k, \tag{16}$$

where $l_k \geq 0$ is the number of distinct species with frequency k observed within the sample (X_1, \ldots, X_n) . Moreover,

$$\widehat{U}_{n+m}(k) = \sum_{i=1}^{k} l_i (i-\sigma)_{k+1-i} \binom{m}{k-i} \frac{(\theta+n-i+\sigma)_{m-k+i}}{(\theta+n)_{m+1}} + \binom{m}{k} \frac{(1-\sigma)_k}{(\theta+n)_{m+1}} \left\{ (\theta+j\sigma)(\theta+n+\sigma)_{m-k} - \prod_{i=j}^{j+m-k} (\theta+i\sigma) \right\}.$$

$$(17)$$

From the previous result, one then immediately deduces the cumulative discovery estimator $\widehat{U}_{n+m,\tau}$. Finally, for completeness, we also consider the special case of the Dirichlet process with parameter $\theta>0$ and baseline probability measure P_0 , which coincides, in distribution, with the PD $(0,\theta;P_0)$ process. Therefore, from (16)–(17), it is straightforward to see that the corresponding estimators reduce to, respectively, $\widehat{U}_{n+0}(k)=k\,l_k/(\theta+n)$ and

$$\begin{split} \widehat{U}_{n+m}(k) &= k! \sum_{i=1}^{k} \frac{l_i}{(i-1)!} \binom{m}{k-i} \frac{(\theta+n-i)_{m-k+i}}{(\theta+n)_{m+1}} \\ &+ \binom{m}{k} \frac{k!}{(\theta+n)_{m+1}} \left\{ \theta (\theta+n)_{m-k} - \theta^{m-k+1} \right\}. \end{split}$$

It is interesting to note that in the Dirichlet case, $\widehat{U}_{n+0}(k+1) = (\theta+n)\check{U}_{n+0}(k)/n$, where \check{U}_{n+0} is the Turing estimator in (3).

3. An Application to Genomic Data

The genomic datasets we analyze consist of ESTs samples, which play an important role in the identification, discovery, and characterization of organisms as they provide an attractive and efficient alternative to full genome sequencing. ESTs are single-read sequences of complementary DNA (cDNA) fragments obtained by sequencing randomly selected cDNA clones from a cDNA library. Since a cDNA library consists of millions of cDNA clones, only a small fraction is usually sequenced because of cost constraints. This is a natural

Table 1
ESTs from two Naegleria gruberi libraries. Reported data include: frequency counts l_k , for different values of k, total number of distinct genes j, and sample size n. Source: Susko and Roger (2004).

1	2	3	4	5	6	7	8	9	10	11
346	57	19	12	9	5	4	2	4	5	4
491	72	30	9	13	5	3	1	2	0	1
12	13	14	15	16	17	18	27	55	j	n
1	0	0							473	959
0	1	3	0	0	0	0	0	0	631	969
	491 12 1	346 57 491 72 12 13 1 0	346 57 19 491 72 30 12 13 14 1 0 0	346 57 19 12 491 72 30 9 12 13 14 15 1 0 0 0	346 57 19 12 9 491 72 30 9 13 12 13 14 15 16 1 0 0 0 1	346 57 19 12 9 5 491 72 30 9 13 5 12 13 14 15 16 17 1 0 0 0 1 1	346 57 19 12 9 5 4 491 72 30 9 13 5 3 12 13 14 15 16 17 18 1 0 0 0 1 1 1 1	346 57 19 12 9 5 4 2 491 72 30 9 13 5 3 1 12 13 14 15 16 17 18 27 1 0 0 0 1 1 1 1	346 57 19 12 9 5 4 2 4 491 72 30 9 13 5 3 1 2 12 13 14 15 16 17 18 27 55 1 0 0 0 1 1 1 1 1	346 57 19 12 9 5 4 2 4 5 491 72 30 9 13 5 3 1 2 0 12 13 14 15 16 17 18 27 55 j 1 0 0 0 1 1 1 1 1 473

setting in which prediction and, in particular, the estimation of discovery probabilities are of great relevance.

We focus on EST data obtained from Naegleria gruberi cDNA libraries. Naegleria gruberi is a widespread free-living soil and freshwater amoeboflagellate widely studied in the biological literature. The two datasets we consider are sequenced from two cDNA libraries prepared from cells grown under different culture conditions, aerobic and anaerobic, and have been previously analyzed in Susko and Roger (2004), where a full account of their preparation is detailed. The data, which will constitute our basic samples, can be effectively summarized in terms of their frequency counts l_k , for k = 1, ..., nand are reported in Table 1 below. One observes that the frequency counts l_k are large for small values of k and, for initial values of k, l_k decreases monotonically with a few isolated points of increase for larger values of k. These are features common to most EST datasets and to samples drawn from genomic libraries in general. Moreover, the Naegleria Aerobic dataset exhibits several genes with high frequency and, in particular, two genes appearing 27 and 55 times, respectively. On the other hand, the Naegleria Anaerobic dataset yields $l_k = 0$ for any k > 14.

The nonparametric prior in (1) we adopt for analyzing these datasets is given by the two-parameter Poisson-Dirichlet process, concisely discussed in Section 2.3. We will focus on quantifying the rare species variety and provide estimates of both the [m:k]-discovery and of the cumulative discovery $U_{n+m,\tau}$ for different values of the additional sample size m, frequency k, and threshold τ . It is worth noting that such an analysis, still within the two-parameter Poisson-Dirichlet framework, has been carried out in Lijoi et al. (2007b): there the analysis is focused on overall species variety and a comparison of the heterogeneity of the two cDNA libraries, which in this context is typically quantified in terms of their redundancy (the counterpart of species variety), is carried out. According to such a prior specification, one is left to eliciting the actual values of σ and θ that will be used in the inferential process. A natural way to fix these parameters is through an empirical Bayes argument. In particular, $(\widehat{\sigma}, \theta)$ are fixed so to maximize the distribution of $\mathcal{L}^{(n)}$ in correspondence to the observed frequency counts (l_1, \ldots, l_n) , i.e., $(\widehat{\sigma}, \widehat{\theta})$ result from

Table 2
Bayesian nonparametric and Turing estimates of the [0:k]-discoveries (for $k=0,\ldots,4$) and 0e cumulative [0:k]-discoveries (for $\tau=3,4,5$) for the two EST datasets.

	Naegleri	a aerobic	Naegleria anaerobic		
	BNP	Turing	BNP	Turing	
$\overline{U_{n+0}(0)}$	0.3613	0.3608	0.5086	0.5067	
$U_{n+0}(1)$	0.1136	0.1189	0.1485	0.1486	
$U_{n+0}(2)$	0.0754	0.0594	0.0858	0.0929	
$U_{n+0}(3)$	0.0440	0.0501	0.0624	0.0372	
$U_{n+0}(4)$	0.0397	0.0469	0.0267	0.0671	
$U_{n+0,3}$	0.5943	0.5892	0.8053	0.7854	
$U_{n+0,4}$	0.6341	0.6361	0.8320	0.8524	
$U_{n+0,5}$	0.6728	0.6674	0.8822	0.8834	

$$\arg\max_{(\sigma,\theta)} \mathbb{P}\left[\mathcal{L}^{(n)} = (l_1, \dots, l_n)\right] = \arg\max_{(\sigma,\theta)} \times \frac{\prod_{i=1}^{j-1} (\theta + i\sigma)}{(\theta + 1)_{n-1}} n! \times \prod_{i=1}^{n} \left\{\frac{(1-\sigma)_{i-1}}{i!}\right\}^{l_i}_{l_i!}.$$

$$(18)$$

For the two considered libraries, we then obtain $(\widehat{\sigma}, \widehat{\theta}) = (0.67, 46.3)$ in the aerobic case and $(\widehat{\sigma}, \widehat{\theta}) = (0.66, 155.5)$ in the anaerobic case. Alternatively, one could specify a prior for (σ, θ) and devise a suitable Markov Chain Monte Carlo scheme for sampling from their posteriors as in Lijoi et al. (2008). However, with such relatively large basic samples, the posteriors are highly peaked and the results essentially coincide.

In Table 2, we show the estimates of the [0:k]-discoveries and of the cumulative [0:k]-discoveries for different values of k and of τ resulting from the Bayesian nonparametric and the Turing estimators. The estimates from the two methods are basically the same for k=0 and k=1, which correspond to the probabilities of sequencing a new gene and a unique gene, respectively. Larger discrepancies are detectable as kgets bigger. This is also reflected in the estimation of the cumulative discoveries. With reference to the discussion in Remark 1, it is worth noting that the Naegleria anaerobic basic sample displays two species with frequency 9, meaning that $l_9 = 2$ and no species with frequency equal to 10, i.e., $l_{10} = 0$. In such a case, the Turing estimator (3) that depends on $l_{k+1} = l_{10}$ is 0: one would then conclude that the probability that the (n+1)th observation coincides with one of the two species appearing nine times in the basic sample is zero. Moreover, if we were to estimate the [0:10] discovery, the frequentist estimator would lead to a positive value, since $l_{11} > 0$, a conclusion that seems again counterintuitive since there are no species with frequency equal to 10 in the basic sample. This puzzling feature is due to the fact that the estimator of the [0:k]-discovery in (3) depends on l_{k+1} . In contrast, the Bayesian nonparametric estimator (10) is positive and actually equal to 0.014833 for the [0:9]-discovery and 0 for the [0:10]-discovery.

Table 3 Estimates of the [m:k]-discoveries (for $k=0,\ldots,4$) and of the cumulative [m:k]-discoveries (for $\tau=3,4,5$) for different sizes of the additional sample m.

			Naegleria Aerobi	\overline{c}		
	m = 250	m = 500	m = 750	m = 1000	m = 1250	m = 1500
$\overline{U_{n+m}\left(0\right)}$	0.3358	0.3162	0.3006	0.2877	0.2768	0.2673
$U_{n+m}\left(1\right)$	0.1066	0.1011	0.0965	0.0927	0.0894	0.0865
$U_{n+m}(2)$	0.0703	0.0664	0.0634	0.0609	0.0587	0.0569
$U_{n+m}\left(3\right)$	0.0475	0.0476	0.0467	0.0455	0.0443	0.0432
$U_{n+m}(4)$	0.0373	0.0370	0.0366	0.0361	0.0355	0.0348
$U_{n+m,3}$	0.5602	0.5313	0.5072	0.4867	0.4692	0.4539
$U_{n+m,4}$	0.5974	0.5683	0.5438	0.5228	0.5046	0.4887
$U_{n+m,5}$	0.6307	0.5996	0.5743	0.5528	0.5342	0.5178
			Naegleria Anaerol	bic		
$U_{n+m}\left(0\right)$	0.4751	0.4489	0.4275	0.4097	0.3945	0.3813
$U_{n+m}(1)$	0.1428	0.1377	0.1330	0.1289	0.1251	0.1218
$U_{n+m}(2)$	0.0849	0.0834	0.0817	0.0800	0.0783	0.0767
$U_{n+m}\left(3\right)$	0.0612	0.0602	0.0593	0.0584	0.0575	0.0565
$U_{n+m}(4)$	0.0388	0.0429	0.0443	0.0447	0.0446	0.0444
$U_{n+m,3}$	0.7639	0.7301	0.7015	0.6769	0.6554	0.6363
$U_{n+m,4}$	0.8027	0.7729	0.7458	0.7216	0.7000	0.6807
$U_{n+m,5}$	0.8384	0.8074	0.7809	0.7572	0.7360	0.7167

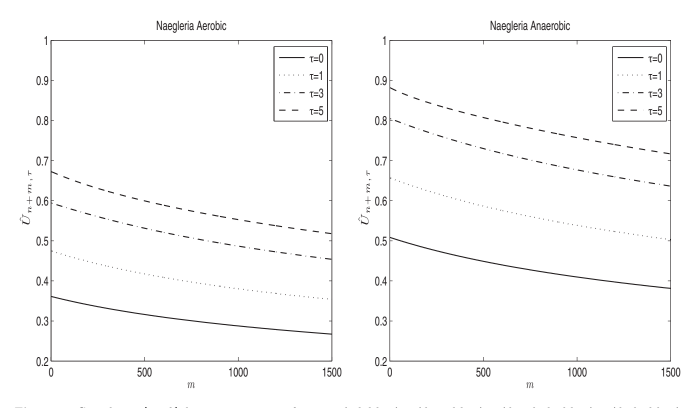


Figure 1. Cumulative [m:k]-discovery estimates for $\tau=0$ (solid line), 1 (dotted line), 3 (dot-dashed line), 5 (dashed line), and additional sample sizes $m=0,\ldots,1500$. Left and right panels refer, respectively, to Naegleria aerobic and anaerobic datasets.

In addition to one-step prediction estimates, we are also able to provide estimates of $U_{n+m}\left(k\right)$, and for the corresponding cumulative values, in the case where both m and k are strictly positive. These represent the main quantities of interest in such genomic experiments and as noted in Section 2,

no other estimators are available in the literature. The results are summarized in Table 3. It is worth remarking that the displayed numbers are exact, in the sense that (12) and (13) can be computed without the need to resorting to any approximation scheme.

The behavior of the cumulative [m:k]-discoveries, that provide evidence of the possibility of detecting rare genes as the additional sample size increases, is consistent with intuition: the larger the additional sample size m, the lower the probability that the (n+m+1)th observation identifies a rare gene, i.e., a gene that has never been observed before or that has been observed with frequency not greater than τ . Moreover, the larger is the considered frequency level k, the slower is the decay rate. The two datasets display quite different results, which become even more apparent from Figure 1. In particular, the *Naegleria Anaerobic* is a richer (or, equivalently, in biological terminology, less redundant) library in the sense that the probability of detecting rare species is significantly larger if compared with the Naegleria Aerobic library. This is consistent with the fact that the basic sample of the Naegleria Aerobic library exhibits several genes with high frequency, which, in fact, hints toward redundancy. As anticipated in Section 2.2, one can also address the issue of sample size determination. If we agree that $\tau = 3$ defines the upper bound for the frequency of species considered as rare, one can consider $m \mapsto \widehat{U}_{n+m,3}$ and identify the largest m that makes \widehat{U}_{n+m} not smaller than a certain threshold κ . For example, still referring to the Naegleria Aerobic dataset, if we were to choose $\kappa = 0.5$ then one should set a value of the additional sample size m = 833.

4. Supplementary Materials

A Web Appendix, which contains the Matlab code for computing the proposed estimators and the proofs of our results, is available with this article at the Biometrics website on Wiley Online Library.

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