- A Bayesian Model of the DNA Barcode Gap
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- Running Title: Bayesian inference for DNA barcode gap estimation

8 Abstract

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A simple statistical model of the DNA barcode gap is outlined. Specifically, accuracy of recently introduced nonparametric metrics, inspired by coalescent theory, to characterize the extent of proportional overlap/separation in maximum and minimum pairwise genetic distances within and among species, respectively, is explored in both frequentist and Bayesian contexts. The empirical cumulative distribution function (ECDF) is utilized to estimate probabilities associated with positively skewed extreme tail distribution quantiles bounded on the closed unit interval [0, 1] based on a straightforward binomial distance overlap count. Using R and Stan, the proposed maximum likelihood estimators and Bayesian model are demonstrated on cytochrome b (CYTB) gene sequences from two Agabus diving beetle species exhibiting limits in the extent of representative taxonomic sampling. Large-sample theory and MCMC simulations show much uncertainty in parameter estimates, particularly when specimen sample sizes for target species are small. Findings highlight the promise of the Bayesian approach using a conjugate beta prior for reliable posterior uncertainty estimation when available data are sparse. Obtained results can shed light on foundational and applied research questions concerning DNA-based specimen identification and species delineation for studies in evolutionary biology and ecology, as well as biodiversity conservation and management, of wide-ranging taxa.

Keywords: Bayesian/frequentist inference, DNA barcoding, intraspecific genetic distance, interspecific genetic distance, specimen identification, species discovery

₁₉ 1 Introduction

The routine use of DNA sequences (particularly mitochondrial DNA (mtDNA)) to support broad evolutionary hypotheses and questions concerning demographic processes, like gene flow and speciation, that have produced a distinctive and measurable pattern of genetic polymorphism in diverse and spatially-distributed taxonomic lineages such as birds, fishes,

insects, and arachnids, among other extensively studied groups, took flight in the late 1980s (Avise et al., 1987). The application of genomic data to applied fields like biodiversity forensics, conservation, and management for the molecular identification of unknown specimen samples came later (e.q., Forensically Informative Nucleotide Sequencing (FINS); 37 Bartlett and Davidson (1992)). Since its inception over 20 years ago, DNA barcoding (Hebert et al., 2003a,b) built significantly on earlier work and has emerged as a robust method of specimen identification and species discovery across myriad multicellular Eukaryotes which have been sequenced at easily obtained short, standardized gene regions like the cytochrome c oxidase subunit I (5'-COI) mitochondrial locus for animals. However, the success of the single-locus approach, particularly for regulatory and forensic applications, depends crucially on two important factors: (1) the availability of high-quality specimen records found in public reference sequence databases such as the Barcode of Life Data Systems (BOLD; http://www.barcodinglife.org) (Ratnasingham and Hebert, 2007) and GenBank 46 (https://www.ncbi.nlm.nih.gov/genbank/), and (2) the establishment of a DNA barcode gap 47 — the notion that the maximum genetic distance observed within species is much smaller than the minimum degree of marker variation found among species (Meyer and Paulay, 2005; Meier et al., 2008). Early work has demonstrated that the presence of a DNA barcode gap hinges strongly on extant levels of species haplotype diversity gauged from comprehensive specimen sampling at wide geographic and ecological scales (Bergsten et al., 2012; Candek 52 and Kuntner, 2015). Despite this, many taxonomic groups lack adequate separation in their 53 pairwise intraspecific and interspecific genetic distances due to varying rates of evolution in both genes and taxa (Pentinsaari et al., 2016). Furthermore, it has been well-demonstrated that the presence of a DNA barcode gap becomes less certain with increasing spatial scale of sampling since interspecific distances increase, while intraspecific distances shrink as more closely-related species are sampled (Phillips et al., 2022). This can pose problems in cases of rare species or monotypic taxa, for instance (Ahrens et al., 2016) and compromise rapid matching of unknown samples to expertly-validated references, leading to cases of false positives (taxon oversplitting) and false negatives (excessive lumping of taxa) as a result of incomplete lineage sorting, interspecies hybridization, genome introgression, species synonymy, cryptic species diversity, and misidentifications (Hubert and Hanner, 2015; Phillips et al., 2022).

Recent work has argued that DNA barcoding, in its current form, is lacking in statistical 65 rigor, as most studies rely strongly on heuristic distance-based measures to infer taxonomic identity. Of these studies, few report measures of uncertainty, such as standard errors (SEs) and confidence intervals (CIs), around estimates of intraspecific and interspecific variation, calling into question the existence of a true species' DNA barcode gap (Candek and Kuntner, 2015; Phillips et al., 2022). To support this notion, novel nonparametric locus- and species-specific metrics based on the multispecies coalescent (MSC) were recently outlined by Phillips et al. (2024). Unlike previously proposed MSC algorithmic approaches (of which there are too many to exhaustively list here), which generally assume a strict molecular clock 73 and a simplified model of DNA sequence evolution across closely-related taxa from which an estimated species phylogeny may be constructed (e.g., with or without use of a guide tree) (e.g., Rannala and Yang (2003, 2017); Yang and Rannala (2010, 2014, 2017)), Phillips et al.'s (2024) approach is tree-free and does not require judicious parameter setting. The statistics have been shown to hold strong promise for reliable DNA barcode gap assessment when applied to predatory Agabus (Coleoptera: Dytiscidae) diving beetles (Phillips et al., 79 2024). Despite their ease of sampling and well-established taxonomy, this group possesses few morphologically-distinct taxonomic characters that readily facilitate their assignment to the 81 species level (Bergsten et al., 2012). Further, the proposed metrics indicate that sister species pairs from this taxon are often difficult to distinguish on the basis of their DNA barcode 83 sequences (Phillips et al., 2024). Using sequence data from three mitochondrial cytochrome markers (5'-COI, 3'-COI, and cytochrome b (CYTB)) obtained from BOLD and GenBank, results highlight that DNA barcoding has been a one-sided argument. Phillips et al.'s (2024) findings point to the need to balance both the sufficient collection of specimens, as well as the

extensive sampling of species: DNA barcode libraries are biased toward the latter (Phillips et al., 2022). The coalescent (Kingman, 1982a,b) encompasses a backwards continuous-time stochastic Markov process of allelic sampling within natural, neutrally-evolving, species populations towards the most recent common ancestor (MRCA). The estimators from Phillips 91 et al. (2024) represent a clear improvement over simple, yet arbitrary, distance heuristics such as the 2% rule noted by Hebert et al. (2003a) and the 10× rule (Hebert et al., 2004) that form the basis of single-locus species delimation tools like Automatic Barcode Gap Discovery (ABGD) (Puillandre et al., 2011), Assemble Species by Automatic Partitioning (ASAP) (Puillandre et al., 2021), and the Barcode Index Number (BIN) framework (Ratnasingham and Hebert, 2013). The 2% rule asserts that DNA sequences differing by at least 2% at 97 sequenced genomic regions should be expected to originate from different biological species, whereas the $10\times$ rule suggests that sequences displaying 10 times more genetic variation among species than within taxa is evidence for a distinct evolutionary origin. However, 100 the lack of adoption of an explicit, universally agreed upon, species concept that readily 101 governs lineage formation and proliferation necessary to establish rigorous taxon definitions 102 for successful delimitation of hypothesized and heuristic evolutionary units using these 103 well-known criteria, is missing (Rannala, 2015; Pante et al., 2015; Wells et al., 2022). In 104 addition, the reliance on visualization approaches, such as frequency histograms, dotplots, 105 and quadrant plots to expose DNA barcoding's limitations, has also been criticized 106 (Collins and Cruickshank, 2013; Phillips et al., 2022). Up until the work of Phillips et al. 107 (2024), the majority of studies (e.g., Young et al. (2021)) have treated the DNA barcode 108 gap as a binary response. However, given poor sampling depth for most taxa, a Yes/No 109 dichotomy is inherently flawed because it can falsely imply a DNA barcode gap is present for 110 a taxon of interest when in fact no such separation in distances exists. The proposed statistics 111 quantify the extent of asymmetric directionality of proportional distance distribution overlap/separation for species within well-sampled taxonomic genera based on a 113 straightforward distance count, in a similar vein to established measures of statistical

similarity such as the Kullback-Leibler (KL) divergence (Kullback and Leibler, 1951) and other related statistics of f-divergence. The metrics can be employed in a variety of ways, 116 including to validate performance of marker genes for specimen identification to the species 117 level (as in Phillips et al. (2024)), as well as to assess whether computed values are consistent 118 with population genetic-level parameters like effective population size (N_e) , mutation rates 119 (μ) and divergence times (τ) for species under study in a statistical phylogeographic setting 120 (Knowles and Maddison, 2002; Mather et al., 2019). Early on, DNA barcoding was presumed 121 to only work for reciprocally monophyletic groups and thus concerned itself with terminal 122 branches of generated phylogenies rather than more basal lineages occurring deeper in 123 hypothesized species trees (Mutanen et al., 2016). Furthermore, the occurrence of short 124 branches within resolved phylogenies increases the probability of deep coalescence, clouding species delimitations. As DNA barcoding is a single-locus approach, it is problematic for 126 evolutionarily young taxa, wherein incomplete lineage sorting within gene genealogies is 127 a common phenomenon. The most promising way forward in this regard seems to be 128 through the use of software such as BPP (Bayesian Phylogenetics and Phylogeography), 129 which permits efficient full Bayesian simulations under various MSC models (e.q., MSC-I 130 (MSC with introgression) or MSC-M (MSC with migration), among others) using MCMC 131 for tree parameter estimation (using the A00 option, for instance) (Flouri et al., 2018), or 132 PHRAPL (Phylogeographic Inference using Approximate Likelihoods) (Jackson et al., 2017), 133 which employs tractable phylogenetic likelihood calculations. 134

While introduction of the metrics is a step in the right direction, what appears to be missing is a rigorous statistical treatment of the DNA barcode gap. This includes an unbiased way to compute the statistical accuracy of Phillips et al.'s (2024) estimators arising through problems inherent in frequentist maximum likelihood estimation for probability distributions having bounded positive support on the closed unit interval [0, 1]. To this end, here, a Bayesian model of the DNA barcode gap coalescent is introduced to rectify such issues. The model allows accurate estimation of posterior means, posterior standard deviations (SDs),

posterior quantiles, and credible intervals (CrIs) for the metrics given datasets of intraspecific and interspecific distances for species of interest.

$_{\scriptscriptstyle{144}}$ 2 Methods

2.1 DNA Barcode Gap Metrics

The novel nonparametric maximum likelihood estimators (MLEs) of proportional overlap/separation between intraspecific and interspecific distance distributions for a given species (x) to aid assessment of the DNA barcode gap are as follows:

$$p_x = \frac{\#\{d_{ij} \ge a\}}{\#\{d_{ij}\}} \tag{1}$$

$$q_x = \frac{\#\{d_{XY} \le b\}}{\#\{d_{XY}\}} \tag{2}$$

$$p_x' = \frac{\#\{d_{ij} \ge a'\}}{\#\{d_{ij}\}} \tag{3}$$

$$q_x' = \frac{\#\{d_{XY}' \le b\}}{\#\{d_{XY}'\}} \tag{4}$$

where d_{ij} are distances within species, d_{XY} are distances among species for an entire genus of concern, and d'_{XY} are combined interspecific distances for a target species and its closest neighbouring species. The notation # reflects a count. Quantities a, a', and b correspond to $\min(d_{XY})$, $\min(d'_{XY})$, and $\max(d_{ij})$, the minimum interspecific distance, the minimum combined interspecific distance, and the maximum intraspecific distance, respectively (Figure 1).

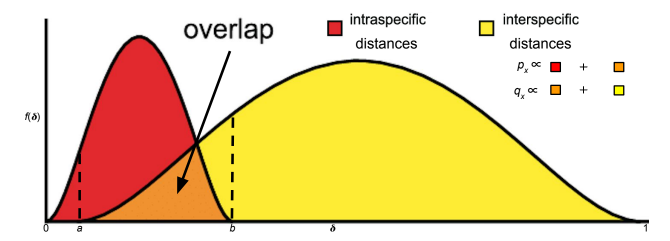


Figure 1: Modified depiction from Meyer and Paulay (2005) and Phillips et al. (2024) of the overlap/separation of intraspecific and interspecific distances (δ) for calculation of the DNA barcode gap metrics (p_x and q_x) for a hypothetical species x. The minimum interspecific distance is denoted by a and the maximum intraspecific distance is indicated by b. The quantity $f(\delta)$ is akin to a kernel density estimate of the probability density function of distances. A similar visualization can be displayed for p'_x and q'_x within the interval [a', b].

Hence, Equations (1)-(4) are simply empirical partial means of distances falling at and below, or at and exceeding, given distribution thresholds. Notice further that a/a', and b are also 156 the first and nth order statistics, $X_{(1)}$ and $X_{(n)}$, respectively, with a/a' < b, which have been 157 pointed out by Phillips et al. (2022) as important for developing a mathematical theory to 158 test the existence of the DNA barcode gap. Equations (1)-(4) can also be expressed in terms 159 of empirical cumulative distribution functions (ECDFs) (see next section). Distances form a 160 continuous distribution and are easily computed from a model of DNA sequence evolution, 161 such as uncorrected or corrected p-distances (Jukes and Cantor, 1969; Kimura, 1980) using, 162 for example, the dist.dna() function available in the ape R package (Paradis et al., 2004); 163 however, values are not independent and identically distributed (IID) because estimated 164 standard errors (SEs) will depend on both the number of species sampled with the genus 165 under study, as well as the number of specimens sampled within a target species. In Phillips 166 et al. (2024), To tease this out, Phillips et al. (2024) suggests plotting estimator values against 167 their estimated SEs, along with a simple random downsampling scheme. In the case of two 168 species comprising a focal genus, one well sampled and the other poorly sampled, values 169

of the metrics close to zero obtained from Equations (1)-(4) for the sufficiently sampled species will likely possess larger SEs following downsizing to match the number of poorly 171 sampled specimens (Phillips et al., 2024). The approach of Phillips et al. (2024) differs 172 markedly from the traditional definition of the DNA barcoding gap laid out by Meyer and 173 Paulay (2005) and Meier et al. (2008) in that the proposed metrics incorporate interspecific 174 distances which *include* the target species of interest. Furthermore, if a focal species is found to have multiple nearest neighbours, then the species possessing the smallest average distance 176 is used. These schemes more accurately account for species' coalescence processes inferred 177 from contemporaneous samples of DNA sequences leading to instances of barcode sequence 178 sharing, such as interspecific hybridization/introgression events (Phillips et al., 2024). Within 179 equations (3) and (4), the degree of distance distribution overlap between a target taxon and 180 its nearest neighbouring species, gauged from magnitudes of p'_x and q'_x , is directly proportional 181 to the amount of time in which the two lineages diverged from the MRCA (Phillips et al., 182 2024). Thus, the quantities can be used as a criterion to assess the failure of DNA barcoding 183 in recently radiated taxonomic groups, among other plausible biological explanations. Note, 184 distances are constrained to the interval [0, 1], whereas the metrics are defined only on 185 the interval [a/a', b]. Values of the estimators obtained from equations (1)-(4) close to or 186 equal to zero give evidence for separation between intraspecific and interspecific distance 187 distributions; that is, values suggest the presence of a DNA barcode gap for a target species. 188 Conversely, values near or equal to one give evidence for distribution overlap; that is, values 189 likely indicate the absence of a DNA barcode gap. 190

191 **2.2** The Model

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Before delving into the derivation of the proposed DNA barcode gap metrics, review of some fundamental statistical theory is necessary.

For a given random variable X, its cumulative distribution function (CDF) is defined by

$$F_X(t) = \mathbb{P}(X \le t) = 1 - \mathbb{P}(X > t). \tag{5}$$

Rearranging Equation (5) gives

$$\mathbb{P}(X > t) = 1 - F_X(t) = 1 - \mathbb{P}(X \le t), \tag{6}$$

196 from which it follows that

$$\mathbb{P}(X \ge t) = 1 - F_X(t) + \mathbb{P}(X = t). \tag{7}$$

Equations (1)-(4) can thus be expressed in terms of ECDFs as follows, since the true underlying CDFs, $F(\cdot)$, are unknown *a priori*, and therefore must be estimated using available data:

$$p_x = \mathbb{P}(d_{ij} \ge a)$$

$$= 1 - \hat{F}_{d_{ij}}(a) + \mathbb{P}(d_{ij} = a)$$

$$= \hat{F}_{d_{ij}}(b) - \hat{F}_{d_{ij}}(a) + \mathbb{P}(d_{ij} = a)$$
(8)

$$q_x = \mathbb{P}(d_{XY} \le b)$$

$$= \hat{F}_{d_{XY}}(b) \tag{9}$$

$$p'_{x} = \mathbb{P}(d_{ij} \ge a')$$

$$= 1 - \hat{F}_{d_{ij}}(a') + \mathbb{P}(d_{ij} = a')$$

$$= \hat{F}_{d_{ij}}(b) - \hat{F}_{d_{ij}}(a') + \mathbb{P}(d_{ij} = a')$$
(10)

$$q'_{x} = \mathbb{P}(d'_{XY} \le b)$$

$$= \hat{F}_{d'_{XY}}(b) \tag{11}$$

From this, it can be seen that $\hat{F}_{d_{ij}}(b) = 1$ in Equations (8) and (10). Given n 200 increasing-ordered data points, the (discrete) ECDF, $\hat{F}_n(t) = \frac{1}{n} \sum_{i=1}^n \mathbb{1}_{[x_i \leq t]}$, comprises a step 201 function having jump discontinuities of size $\frac{1}{n}$ at each sample observation (x_i) , excluding ties 202 (or steps of weight $\frac{i}{n}$ with duplicate observations), where $\mathbb{1}(x)$ is the indicator function. Note, 203 $\mathbb{P}(X=t) \neq 0$. Equations (8)-(11) clearly demonstrate the asymmetric directionality of the 204 proposed metrics. Furthermore, calculation of the DNA barcode gap estimators is convenient 205 as they implicitly account for total distribution area (including overlap). 206 A major criticism of large sample (frequentist) theory is that it relies on asymptotic 207 properties of the MLE (whose population parameter is assumed to be a fixed but unknown 208 quantity), such as estimator normality and consistency as the sample size approaches infinity. 209 This problem is especially pronounced in the case of binomial proportions (Newcombe, 1998). 210 The estimated Wald SE of the sample proportion, is given by 211 $\widehat{SE[\hat{p}]} = \sqrt{\frac{\hat{p}(1-\hat{p})}{n}}$, where $\hat{p} = \frac{Y}{n}$ is the MLE, Y is the total number of successes $(Y = \sum_{i=1}^{n} y_i)$

and n is the total number of trials (i.e., sample size). However, the above formula for the standard error is problematic for several reasons. First, it is a Normal approximation which 214 makes use of the central limit theorem (CLT); thus, large sample sizes are required for reliable 215 estimation. When few observations are available, SEs will be large and inaccurate, leading 216 to low statistical power to detect a true DNA barcode gap when one actually exists. Further, 217 resulting interval estimates could span values less than zero or greater than one, or have zero width, which is practically meaningless. Second, when proportions are exactly equal to zero 219 or one, resulting SEs will be exactly zero, rendering $\widehat{SE[\hat{p}]}$ given above completely useless. 220 In the context of the proposed DNA barcode gap metrics, values obtained at the boundaries 221 of their support are often encountered. Therefore, reliable calculation of SEs is not feasible. 222 Given the importance of sufficient sampling of species genetic diversity for DNA barcoding 223 initiatives, a different statistical estimation approach is necessary. 224

Bayesian inference offers a natural path forward in this regard since it allows for 225 straightforward specification of prior beliefs concerning unknown model parameters and 226 permits the seamless propagation of uncertainty, when data are lacking and sample sizes 227 are small, through integration with the likelihood function associated with true generating 228 processes. The posterior distribution $(\pi(\theta|Y))$ is given by Bayes' theorem up to a 229 proportionality $\pi(\theta|Y) \propto \pi(Y|\theta)\pi(\theta)$, where θ are unobserved parameters, Y are known 230 data, $\pi(Y|\theta)$ is the likelihood, and $\pi(\theta)$ is the prior. As a consequence, because parameters 231 are treated as random variables, Bayesian models are much more flexible and generally more 232 easily interpretable compared to frequentist approaches. Under the Bayesian paradigm, entire 233 posterior distributions, along with their summaries (e.g., CrIs) are outputted, rather than just 234 long run behaviour reflected in sampling distributions, p-values, and CIs as in the frequentist 235 case, thus allowing direct probability statements to be made. 236

Essentially, from a statistical perspective, the goal herein is to nonparametrically estimate probabilities corresponding to extreme tail quantiles for positive highly skewed distributions on the unit interval (or any closed subinterval thereof). Here, it is sought to numerically

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approximate the extent of proportional overlap/separation of intraspecific and interspecific distance distributions within the subinterval [a/a', b]. This is a challenging computational problem within the current study as detailed in subsequent sections. The usual approach 242 employs kernel density estimation (KDE), along with numerical or Monte Carlo integration 243 of complicated probability distribution functions (PDFs), and invocation of extreme value 244 theory (EVT); however, this requires careful selection of the bandwidth parameter, among other considerations. This becomes problematic when fitting finite mixture models where nonidentifiability is rampant. For DNA barcode gap estimation, this would correspond 247 to a two-component mixture (one for intraspecific distance comparisons, and the other for 248 interspecific comparisons), with one or more curve intersection points between components, 249 and the presence of zero distance inflation. This makes parameter estimation difficult using 250 methods like the Expectation-Maximization (EM) algorithm (Dempster et al., 1977) as 251 the algorithm may become stuck in suboptimal regions of the parameter search space and 252 prematurely converge to local optima. Here, for simplicity, a different route is taken to 253 avoid these obstacles. Counts, y, of overlapping distances (as expressed in the numerator of 254 Equations (1)-(4)) are treated as binomially distributed with expectation $\mathbb{E}[Y] = k\theta$, where 255 $k = \{N, C\}$ are total count vectors of intraspecific and combined interspecific distances, 256 respectively, for a target species along with its nearest neighbour species, and k = M is a 257 total count vector for all interspecific species comparisons. This follows from the fact that 258 the ECDF is binomially distributed. The quantity thus being estimated is the parameter 259 vector $\underline{\theta} = \{p_x, q_x, p'_x, q'_x\}.$ 260 The metrics encompassing $\underline{\theta}$ are presumed to follow a Beta(α , β) distribution, with real 261 shape parameters α and β , which is a natural choice of prior on probabilities. The beta 262 distribution has a prior mean of $\mathbb{E}[\theta] = \frac{\alpha}{\alpha + \beta}$ and a prior variance equal to $\mathbb{V}[\theta] = \frac{\alpha\beta}{(\alpha + \beta)^2(\alpha + \beta + 1)}$. In the case where $\alpha = \beta$, all generated Beta(α, β) distributions will possess the same prior

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expectation, whereas the prior variance will shrink as both α and β increase. Such a scheme is

quite convenient since the beta distribution is conjugate to the binomial distribution. Thus,

the posterior distribution is also beta distributed, specifically, $\text{Beta}(\alpha+Y, \beta+n-Y)$, having expectation $\mathbb{E}[\theta|Y] = \frac{\alpha+Y}{\alpha+\beta+n}$ and variance $\mathbb{V}[\theta|Y] = \frac{(\alpha+Y)(\beta+n-Y)}{(\alpha+\beta+n)^2(\alpha+\beta+n+1)}$. In the context of 268 DNA barcoding, it is important that the DNA barcode gap metrics effectively differentiate 269 between extremes of no overlap/complete separation and complete overlap/no separation, 270 corresponding to values of the metrics equal to 0 and 1 (equivalent to total distance counts of 271 0 and n), respectively. These extremes yield a posterior expectation of $\mathbb{E}[\theta|Y=0]=\frac{\alpha}{\alpha+\beta+n}$ and a posterior variance of $\mathbb{V}[\theta|Y] = 0$ = $\frac{\alpha(\beta+n)}{(\alpha+\beta+n)^2(\alpha+\beta+n+1)}$ and $\mathbb{E}[\theta|Y = n] = \frac{\alpha+n}{\alpha+\beta+n}$ 273 and $\mathbb{V}[\theta|Y=n]=\frac{(\alpha+n)\beta}{(\alpha+\beta+n)^2(\alpha+\beta+n+1)}$. Note, the posterior variances are equivalent at these 274 thresholds for all $\alpha = \beta$. 275 Parameters were given an uninformative Beta(1, 1) prior, which is equivalent to a standard 276 uniform (Uniform(0, 1)) prior since it places equal probability on all parameter values within 277 its support. This distribution has an expected value of $\mu = \frac{1}{2}$ and a variance of $\sigma^2 = \frac{1}{12}$. 278 Further, the posterior is Beta(Y+1, n-Y+1), from which various moments such as the 279 expected value $\mathbb{E}[Y] = \frac{Y+1}{n+2}$ and variance $\mathbb{V}[Y] = \frac{(Y+1)(n-Y+1)}{(n+2)^2(n+3)}$, and other quantities, can be 280 easily calculated. Clearly, $\mathbb{E}[\theta|Y=0] = \frac{1}{n+2}$ and $\mathbb{V}[\theta|Y=0] = \frac{n+1}{(n+2)^2(n+3)}$, and 281 $\mathbb{E}[\theta|Y=n]=\frac{n+1}{n+2}$ and $\mathbb{V}[\theta|Y=n]=\frac{n+1}{(n+2)^2(n+3)}$. In general however, when possible, 282 it is always advisable to incorporate prior information, even if only weak, rather than 283 simply imposing complete ignorance in the form of a flat prior distribution. In the case 284 of unimodal distributions, the (estimated) posterior mean often possesses the property that 285 it readily decomposes into a convex linear combination, in the form of a weighted sum, of the 286 (estimated) prior mean and the MLE. That is $\hat{\mu}_{posterior} = w\hat{\mu}_{prior} + (1-w)\hat{\mu}_{MLE}$, where for the 287 beta distribution, $w = \frac{\alpha+\beta}{\alpha+\beta+n}$. Therefore, with sufficient data, $w \to 0$ as $n \to \infty$, regardless 288 of the values of α and β , and the choice of prior distribution becomes less important since 289 the posterior will be dominated by the likelihood. For the Beta(1, 1), $w = \frac{2}{2+n}$, with n = 2290 giving $w = \frac{1}{2}$; that is, the posterior is the arithmetic average of the prior and the likelihood. 291 The full Bayesian model for species x is thus given by

$$y_{\mathrm{lwr}} \sim \mathrm{Binomial}(N, p_{\mathrm{lwr}})$$

$$y_{\mathrm{upr}} \sim \mathrm{Binomial}(M, p_{\mathrm{upr}})$$

$$y_{\mathrm{lwr}}' \sim \mathrm{Binomial}(N, p_{\mathrm{lwr}}')$$

$$y_{\mathrm{upr}}' \sim \mathrm{Binomial}(C, p_{\mathrm{upr}}')$$

$$p_{\mathrm{lwr}}, p_{\mathrm{upr}}, p_{\mathrm{lwr}}', p_{\mathrm{upr}}' \sim \mathrm{Beta}(1, 1).$$

$$(12)$$

Equation (12) for distinction between MLEs and Bayesian posterior estimates. The above 294 statistical theory and derivations lay a good foundation for the remainder of this paper. 295 The proposed model is inherently vectorized to allow processing of multiple species 296 datasets simultaneously. Model fitting was achieved using the Stan probabilistic 297 programming language (Carpenter et al., 2017) framework for Hamiltonian Monte Carlo 298 (HMC) via the No-U-Turn Sampler (NUTS) sampling algorithm (Hoffman and Gelman, 299 2014) through the rstan R package (version 2.32.6) (Stan Development Team, 2023) in R 300 (version 4.4.1) (R Core Team, 2024). Four Markov chains were run for 2000 iterations each in 301 parallel across four cores with random parameter initializations. Within each chain, a total 302 of 1000 samples was discarded as warmup (i.e., burnin) to reduce dependence on starting 303 conditions and to ensure posterior samples are reflective of the equilibrium distribution. 304 Further, 1000 post-warmup draws were utilized per chain during the sampling phase. Because 305 HMC/NUTS results in dependent samples that are minimally autocorrelated, chain thinning 306 is not required. Each of these tuning parameters reflect default Markov Chain Monte Carlo 307 (MCMC) settings in Stan to control both bias and variance respectively in the resulting 308 draws. All analyses in the present work were carried out on a 2023 Apple MacBook Pro 309 with M2 chip and 16 GB RAM running macOS Ventura 13.2. A random seed was set to 310 ensure reproducibility of model results. Outputted estimates were rounded to three decimal 311

Note that p_x , q_x , $p_x^{'}$, and $q_x^{'}$ in Equations (1)-(4) are denoted p_{lwr} , p_{upr} , $p_{\text{lwr}}^{'}$, $q_{\text{upr}}^{'}$ within

places of precision. Posterior distributions were visualized as KDE plots using the ggplot2
R package (version 3.5.1) (Wickham, 2016) with the default Gaussian kernel and optimal
smoothness selection. To successfully run the Stan program, end users must have installed
an appropriate compiler (such as GCC or Clang) which is compatible with their operating
system, such as macOS.

Convergence was assessed both visually and quantitatively as follows: (1) through 317 examining parameter traceplots, which depict the trajectory of accepted MCMC draws as 318 a function of the number of iterations, (2) through monitoring the Gelman-Rubin potential 319 scale reduction factor statistic (\hat{R}) (Gelman and Rubin, 1992; Vehtari et al., 2021), which 320 measures the concordance of within-chain versus between-chain variance, and (3) through 321 calculating the effective sample size (ESS) for each parameter, which quantifies the number 322 of independent samples generated Markov chains are equivalent to. Mixing of chains was 323 deemed sufficient when traceplots looked like "fuzzy caterpillars", $\hat{R} < 1.01$, and effective 324 sample sizes were reasonably large (Gelman et al., 2020). After sampling, a number of 325 summary quantities were reported, including posterior means, posterior SDs, and posterior 326 quantiles from which 95% CrIs could be computed to make probabilistic inferences concerning 327 true population parameters. To validate the overall correctness of the proposed statistical 328 model given by Equation (12), as a means of comparison, posterior predictive checks (PPCs) 329 were also employed to generate binomial random variates in the form of counts from the 330 posterior predictive distribution; that is $\gamma = \{Np_x, Mq_x, Np_x', Cq_x'\}$ to verify that the model 331 adequately captures relevant features of the observed data. The proposed Bayesian model 332 outlined here has a straightforward interpretation (**Table 1**). 333

334 3 Results and Discussion

The Agabus CYTB dataset analyzed by Phillips et al. (2024) is revisited herein.

Briefly, using the R package MACER (Young et al., 2021), DNA sequences were downloaded

Table 1: Interpretation of the DNA barcode gap estimators within $[a/a',\,b]$

Parameter	Explanation		
$p_x/p_{ m lwr}$	When p_{lwr} is close to 0 (1), it suggests that the probability of intraspecific (interspecific) distances being larger (smaller) than interspecific (intraspecific) distances is low (high) on average, while the probability of interspecific (intraspecific) distances being larger (smaller) than intraspecific (interspecific) distances is high (low) on average; that is, there is (no) evidence for a DNA barcode gap.		
$q_x/p_{ m upr}$	When $p_{\rm upr}$ is close to 0 (1), it suggests that the probability of interspecific (intraspecific) distances being larger (smaller) than intraspecific (interspecific) distances is high (low) on average, while the probability of intraspecific (interspecific) distances being larger (smaller) than interspecific (intraspecific) distances is low (high) on average; that is, there is (no) evidence for a DNA barcode gap.		
$p_x^{'}/p_{ m lwr}^{'}$	When $p'_{\rm lwr}$ is close to 0 (1), it suggests that the probability of intraspecific (combined interspecific distances for a target species and its nearest neighbour species) distances being larger than combined interspecific distances for a target species and its nearest neighbour species (intraspecific distances) is low (high) on average, while the probability of combined interspecific distances for a target species and its nearest neighbour species (intraspecific distances) being larger than intraspecific distances (combined interspecific distances for a target species and its nearest neighbour species) is high (low) on average; that is, there is (no) evidence for a DNA barcode gap.		
$q_x^{'}/p_{ m upr}^{'}$	When $p'_{\rm upr}$ is close to 0 (1), it suggests that the probability of combined interspecific distances for a target species and its earest neighbour species (intraspecific distances) being larger than attraspecific distances (combined interspecific distances for a target species and its nearest neighbour species) is high (low) on average, hile the probability of intraspecific distances (combined interspecific istances for a target species and its nearest neighbour species) being arger than combined interspecific distances for a target species and is nearest neighbour species (intraspecific distances) is low (high) on average; that is, there is (no) evidence for a DNA barcode gap.		

from GenBank and BOLD and processed to obtain a 343 bp FASTA alignment representing 46 unique haplotypes. Genetic distances were calculated using uncorrected p-distances. 338 Specifically, the proposed Bayesian model is demonstrated on the species A. bipustulatus 339 and A. nevadensis, since these taxa were the sole representatives for this locus, with the 340 most and the least specimen records, respectively (N = 701 and N = 2) across all three 341 assessed molecular markers. Further, A. bipustulatus comprised 46 total haplotypes, whereas A. nevadensis possessed two haplotypes. Note, DNA barcode gap estimation is only possible for species having at least two specimen records. This dataset is a prime illustrative example highlighting the issue of inadequate taxon sampling, which arises frequently in large-scale 345 phylogenetic and phylogeographic studies, in several respects. First, from a statistical 346 viewpoint, sample sizes reflect extremes in reliable parameter estimation. Second, from a 347 DNA barcoding perspective, Aqabus currently comprises about 200 extant species according 348 to the Global Biodiversity Information Facility (GBIF) (https://www.gbif.org); yet, due to 349 the level of convenience sampling inherent in taxonomic collection efforts for this genus, 350 adequate representation of species and genetic diversity is far from complete. 351

MCMC parameter traceplots showed rapid mixing of chains to the stationary distribution
(Supplementary Figure 1). Further, all \hat{R} and ESS values (not shown) were close to their
recommended cutoffs of one and thousands of samples, respectively, indicating chains are
both well-mixed and have converged to the posterior distribution.

Bayesian posterior estimates were reported alongside frequentist MLEs, in addition to SEs, posterior SDs, 95% CIs and 95% CrIs (**Table 2**).

Table 2: Nonparametric frequentist and Bayesian estimates of distance distribution overlap/separation for the DNA barcode gap coalescent model parameters applied to A. bipustulatus (N = 701) and A. nevadensis (N = 2) for CYTB, including 95% CIs and CrIs. CrIs are based on 4000 posterior draws. All parameter estimates are reported to three decimal places of precision.

Species	Parameter	MLE (SE, 95% CI)	Bayes Est. (SD; 95% CrI)
A. bipustulatus	$p_x/p_{ m lwr}$	1.000 (0.000; 1.000-1.000)	1.000 (0.000; 1.000-1.000)
A. bipustulatus	$q_x/p_{ m upr}$	1.000 (0.000; 1.000-1.000)	$1.000 \ (0.000; \ 0.999-1.000)$
A. bipustulatus	$p_x^{'}/p_{ m lwr}^{'}$	1.000 (0.000; 1.000-1.000)	1.000 (0.000; 1.000-1.000)
$A.\ bipustulatus$	$q_x^{'}/p_{ m upr}^{'}$	1.000 (0.000; 1.000-1.000)	$1.000 \ (0.000; \ 0.999-1.000)$
$A.\ nevadensis$	$p_x/p_{ m lwr}$	1.000 (0.000; 1.000-1.000)	$0.835\ (0.144;\ 0.470 - 0.996)$
$A.\ nevadensis$	$q_x/p_{ m upr}$	$0.010 \ (0.002; \ 0.006 - 0.014)$	$0.010\ (0.002;\ 0.007 - 0.014)$
$A.\ nevadensis$	$p_x^{'}/p_{ m lwr}^{'}$	1.000 (0.000; 1.000-1.000)	$0.834\ (0.138;\ 0.481 - 0.994)$
$A.\ nevadensis$	$q_x^{'}/p_{ m upr}^{'}$	0.010 (0.070; -0.128-0.148)	$0.010\ (0.002;\ 0.007 \text{-} 0.014)$

CIs were calculated using the usual large sample $(1-\alpha)100\%$ -level interval estimate given by $\hat{p} \pm z_{1-\frac{\alpha}{2}} \sqrt{\frac{\hat{p}(1-\hat{p})}{n}}$, where $z_{1-\frac{\alpha}{2}} = 1.960$ for 95% confidence and α is the stated significance 359 level (here, 5%). Given a $(1-\alpha)100\%$ CI, with repeated sampling, on average $(1-\alpha)100\%$ 360 of constructed intervals will contain the true parameter of interest; on the other hand, any 361 given CI will either capture or exclude the true parameter with 100% certainty. This in 362 stark contrast to a CrI, where the true parameter is contained within said interval with 363 $(1-\alpha)100\%$ probability. Note, by default Stan computes equal-tailed (central) CrIs such 364 that there is equal area situated in the left and right tails of the posterior distribution. For 365 a 95% CrI, this corresponds to the 2.5th and 97.5th percent quantiles. However, constructed 366 intervals are usually only valid for symmetric or nearly symmetric distributions. Given the 367 bounded nature of the DNA barcode gap metrics, whose posterior distributions, as expected, 368 show considerable skewness, a different approach to reporting CrIs, such as Highest Posterior Density (HPD) intervals (Chen and Shao, 1999) or shortest probability intervals (SPIn) (Liu 370 et al., 2015) is warranted. As such asymmetric intervals generally attain greater statistical 371 efficiency (in the form of smaller Mean Squared Error (MSE) or variance) and higher coverage 372 probabilities than more standard interval estimates, careful in-depth comparison is left for 373 future work. 374

Findings based on nonparametric MLEs and Bayesian posterior means were quite

375

comparable with one another and show evidence of complete overlap in intraspecific, interspecific, and combined interspecific distances for A. bipustulatus in both the p/q and p'/q' directions since the metrics attain magnitudes very close to one (**Table 2**). As a result, this likely indicates that no DNA barcode gap is present for this species. Such findings are strongly reinforced by the very tight clustering of posterior draws (**Figure 2**) and associated interval estimates owing to the large number of specimens sampled for this species.

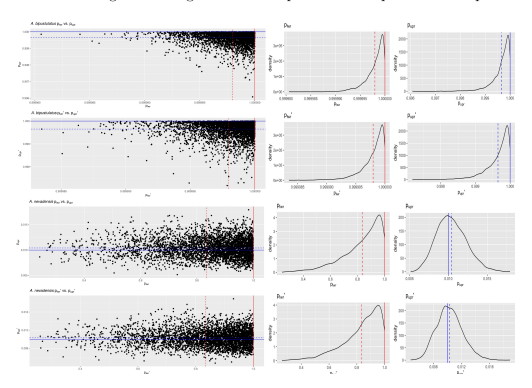


Figure 2: Scatterplots (black solid points) and distributions (black solid lines) depicting the DNA barcode gap metrics for A. bipustulatus (N = 701) and A. nevadensis (N = 2) across CYTB based on 4000 Bayesian posterior draws. MLEs and posterior means are displayed as coloured (red/blue) solid and dashed lines for the metrics, respectively.

On the other hand, the situation for A. nevadensis is more nuanced, as posterior values are further spread out (**Table 2** and **Figure 2**), suggesting less overall certainty in true parameter values given the low specimen sampling coverage for this taxon. Of note, the 95% CIs and 95% CrIs are quite wide for A. nevadensis, consistent with much uncertainty regarding the computed frequentist and Bayesian posterior means of the DNA barcode gap metrics. For instance, the Bayesian analysis for A. nevadensis suggests that the data are

consistent with both $p_{\rm lwr}$ and $p_{\rm lwr}^{'}$ ranging from approximately 0.250-1.000. Further, regarding the frequent ist analysis for the same species, the 95% CI for $q_x^{'}$ extends to negative values 389 at the left endpoint, due to the corresponding SE of 0.070 being too high as a result of 390 the extremely low sample size of n=2 individuals sampled (**Table 2**). Since the 95% CI 391 truncated at the lower endpoint includes the value of zero, the null hypothesis for the presence 392 of a DNA barcode gap cannot be rejected. Despite this, it is worth noting that truncation 393 is not standard statistical practice and will likely lead to an interval with less than 95% nominal coverage. In such cases, more appropriate confidence interval methods like the 395 Wilson score interval, the exact (Clopper-Pearson) interval, or the Agresti-Coull interval 396 should be employed (Newcombe, 1998; Agresti and Coull, 1998). KDEs for A. bipustulatus 397 are strongly left (negatively) skewed (**Figure 2**), whereas those for A. nevadensis exhibit 398 more symmetry, especially for p_{upr} and p'_{upr} (**Figure 2**). These differences are likely due to the 399 stark contrast in sample sizes for the two examined species. Nevertheless, simulated counts 400 of overlapping specimen records from the posterior predictive distribution (Supplementary 401 **Table 1**) were found to be very close to observed counts for both species, indicating that the 402 proposed model adequately captures underlying variation. Obtained results suggest that use 403 of the Beta(1, 1) prior may not be appropriate given a low number of collected individuals 404 for most taxa in DNA barcoding efforts. This suggests that further consideration of more 405 informative beta priors is worthwhile. 406

4 Conclusion

Herein, the accuracy of the DNA barcode gap was analyzed from a rigorous statistical lens to expedite both the curation and growth of reference sequence libraries, ensuring they are populated with high quality, statistically defensible specimen records fit for purpose to address standing questions in ecology, evolutionary biology, management, and conservation. To accomplish this, recently proposed, easy to calculate nonparametric MLEs were formally

derived using ECDFs and applied to assess the extent of overlap/separation of distance 413 distributions within and among two species of predatory water beetles in the genus Aqabus 414 sequenced at CYTB using a Bayesian binomial count model with conjugate beta priors. 415 Findings highlight a high level of parameter uncertainty for A. nevadensis, whereas posterior 416 estimates of the DNA barcode gap metrics for A. bipustulatus are much more certain. 417 Based on these results, it is imperative that specimen sampling be prioritized to better reflect actual species boundaries. More generally, apart from the metrics being employed to 419 better highlighting the importance of within-species genetic diversity versus between-species 420 divergence, it is expected that the approach developed herein will be of broad utility in applied 421 fields, such as DNA-based detection of seafood fraud within global supply chains, and in the 422 determination of species occupancy/detection probabilities at ecological sites of interest using 423 active and passive environmental DNA (eDNA) methods such as metabarcoding. 424

Since the DNA barcode gap metrics often attain values very close to zero (suggesting no 425 overlap and complete separation of distance distributions) and/or very near one (indicating no 426 separation and complete overlap), in addition to more intermediate values, a noninformative 427 $Beta(\frac{1}{2},\frac{1}{2})$ prior may be more appropriate over complete ignorance imposed by a Beta(1, 1)428 prior. The former distribution is U-shaped symmetric and places greater probability density 429 at the extremes of the distribution due to its heavier tails, while still allowing for variability 430 in parameter estimates within intermediate values along its domain. Note that this prior 431 is Jeffreys' prior density (Jeffreys, 1946), which is proportional to the square root of the 432 Fisher information $\mathcal{I}(\theta)$. That is $\pi(\theta) \propto \theta^{-\frac{1}{2}} (1-\theta)^{-\frac{1}{2}}$. Jeffreys' prior has several desirable 433 statistical properties as a prior: that it is inversely proportional to the standard deviation of 434 the binomial distribution, and most notably, that it is invariant to model reparameterization 435 (Gelman et al., 2014). However, this prior can lead to divergent transitions, among other 436 pathologies, imposed by complex geometry (i.e., curvature) in the posterior space since many 437 iterative stochastic MCMC sampling algorithms experience difficulties when exploring high 438 density distribution regions. Thus, remedies to resolve them, such as lowering the step size of

the HMC/NUTS sampler, should be attempted in future work, along with other approaches such as empirical Bayes estimation to approximate beta prior hyperparameters from observed 441 data through the MLE or other methods of parameter estimation, such as the method 442 of moments. Alternatively, hierarchical modelling could be employed to estimate separate 443 distribution model hyperparameters for each species and/or compute distinct estimates for 444 the directionality/comparison level of the DNA barcode gap metrics (i.e., lower vs. upper, non-prime vs. prime) separately within the genus under study. This would permit greater flexibility through incorporating more fine-grained structure seen in the data; however, low taxon sample sizes may preclude valid inferences to be reasonably ascertained due to the 448 large number additional parameters which would be introduced through the specification of 449 the hyperprior distributions. Methods outlined in Gelman et al. (2014), such as dealing with 450 non-exchangeability of observations and alternate model parameterizations like the logit, may 451 prove useful in this regard. Even though more work remains, it is clear that both frequentist 452 and Bayesian inference hold much promise for the future of molecular biodiversity science. 453

⁴⁵⁴ Supplementary Information

None declared.

456 Data Availability Statement

- Raw data, R, and Stan code can be accessed via Dryad at:
- http://datadryad.org/stash/share/
- 459 RZIfMixcEODe0RWP7eyXWQewSVbqEIA9UTrH3ZVKyn4.
- A GitHub repository can be found at:
- https://github.com/jphill01/Bayesian-DNA-Barcode-Gap-Coalescent.

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472 Conflict of Interest

None declared.

474 Author Contributions

JDP wrote the manuscript, wrote R and Stan code, as well as analyzed and interpreted all model results.

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