A guide to Bayesian model checking for ecologists

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- Abstract. Checking that models adequately represent data is an essential component
- of applied statistical inference. Ecologists increasingly use hierarchical Bayesian statistical
- models in their research. The appeal of this modeling paradigm is undeniable, as
- 4 researchers can build and fit models that embody complex ecological processes while
- 5 simultaneously controlling for potential biases arising from sampling artifacts. However,
- 6 ecologists tend to be less focused on checking model assumptions and assessing potential
- ⁷ lack-of-fit when applying Bayesian methods than when they applying more traditional
- 8 modes of inference such as maximum likelihood. There are also multiple ways of assessing
- goodness-of-fit for Bayesian models, each of which has strengths and weaknesses. For
- instance, in ecological applications, the posterior predictive p-value is probably the most
- widely used approach for assessing lack of fit in Bayesian models. Such p-values are
- relatively easy to compute, but they are well known to be conservative, producing p-values

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biased toward 0.5. Alternatively, lesser known approaches to model checking, such as prior predictive checks, cross-validation probability integral transforms, and pivot discrepancy measures may produce more accurate characterizations of goodness-of-fit but are not as 15 well known to ecologists. In addition, a suite of visual and targeted diagnostics can be used 16 to examine violations of different model assumptions and lack-of-fit at different levels of the 17 modeling hierarchy, and to check for residual temporal or spatial autocorrelation. In this review, we synthesize existing literature in order to guide ecologists to the many available 19 options for Bayesian model checking. We illustrate methods and procedures with several ecological case studies, including i) explaining variation in simulated spatio-temporal count 21 data, and (ii) modeling survival and residence times of fur seal mothers on a rookery, and 22 (iii) using N-mixture models to estimate abundance and detection probability of sea otters from an aircraft. We find that commonly used procedures based on posterior predictive p-values have high power to detect extreme model inadequacy, but low power to detect more subtle cases of lack of fit. Tests based on cross-validation and pivot discrepancy measures (including the "sampled predictive p-value") appear to be much better suited to 27 this task and to have better overall statistical performance. We conclude that model checking is an essential component of scientific discovery and learning that should accompany most Bayesian analyses presented in the literature. Bayesian p-value, count data, qoodness-of-fit diagnostic check, hierarchical model, 31 model checking, N-mixture model, pivot discrepancy, posterior predictive check, probability

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interval transform, sampled predictive p-value, spatio-temporal model

Introduction

Ecologists increasingly use Bayesian methods to analyze complex hierarchical models for natural systems (Hobbs and Hooten 2015). There are clear advantages of adopting a Bayesian mode of inference, as one can entertain models that were previously intractable using common modes of statistical inference (e.g., maximum likelihood). Ecologists use Bayesian inference to fit rich classes of models to their datasets, allowing them to separate measurement error from process error, and to model features such as temporal or spatial autocorrelation, individual level random effects, and hidden states (Link et al. 2002, Clark 41 and Bjørnstad 2004, Cressie et al. 2009). Applying Bayesian calculus also results in posterior probability distributions for parameters of interest; used together with posterior model probabilities, these can provide the basis for mathematically coherent decision and risk analyses (Link and Barker 2006, Berger 2013, Williams and Hooten 2016). Ultimately, the reliability of inference from a fitted model (Bayesian or otherwise) 46 depends on how well the model approximates reality. There are multiple ways of assessing a model's performance in representing the system being studied. A first step is often to examine diagnostics that compare observed data to model output to pinpoint if and where any systematic differences occur. This process, which we term model checking, is a critical part of statistical inference, as it helps diagnose assumption violations and illuminate places where a model might be amended to more faithfully represent gathered data. Following this step, one might proceed to compare the performance of alternative models embodying different hypotheses using any number of model comparison or out-of-sample predictive performance metrics (see Hooten and Hobbs 2015, for a review) to gauge the support for alternative hypotheses or optimize predictive ability (Fig. 1). Note that

scientific inference can still proceed if models do not fit the data well, but conclusions need to be tempered; one approach in such situations is to estimate a variance inflation factor to adjust precision levels downward (e.g., Cox and Snell 1989, McCullagh and Nelder 1989). Non-Bayesian statistical software often include a suite of goodness-of-fit diagnostics 60 that examine different types of lack-of-fit (Table 2). For instance, when fitting generalized linear (McCullagh and Nelder 1989) or additive (Wood 2006) models in the R programming environment (R Development Core Team 2015), one can easily access diagnostics such as quantile-quantile, residual, and leverage plots. These diagnostics allow one to assess the reasonability of the assumed probability model, to examine whether there is evidence of heteroskedasticity, and to pinpoint outliers. Likewise, in capture-recapture analysis, there are established procedures for assessing overall fit as well as departures from specific model assumptions which are codified in user-friendly software such as U-CARE (Choquet et al. 2009). Results of such goodness-of-fit tests are routinely reported when publishing analyses in the ecological literature. The implicit requirement that one conduct model checking exercises is not often 71 adhered to when reporting results of Bayesian analyses in the ecological literature. For instance, a search of recent volumes of Ecology indicated that only 25% of articles employing Bayesian analysis on real datasets reported any model checking or goodness-of-fit testing (Fig. 2). There are several reasons why Bayesian model checking is uncommon. First, it likely has to do with momentum; the lack of precedent in ecological literature may lead some authors looking for templates on how to publish Bayesian analyses to conclude that model checking is unnecessary. Second, when researchers seek to publish new statistical methods, applications may be presented more as proof-of-concept exhibits than as definitive analyses that can stand up to scrutiny on their own. In such

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studies, topics like goodness-of-fit and model checking are often reserved for future
   research, presumably in journals with less impact. Third, all of the articles we examined
   did a commendable job in reporting convergence diagnostics to support their contention
   that Markov chains from MCMC output had reached their stationary distribution. Perhaps
   there is a mistaken belief among authors and reviewers that convergence to a stationary
   distribution, combined with a lack of prior sensitivity, implies that a model fits the data?
   Finally, it may just be that those publishing Bayesian analyses in ecological literature ". . .
   like artists, have the bad habit of falling in love with their models" (to borrow a quote
   attributed to G.E.P. Box and referenced by Link and Barker (2010) with regard to model
   checking). However, models can be poor at returning our affection; indeed this monograph
   can be viewed as a partial atonement for unrequited love.
       If we accept the premise that Bayesian models in ecology should be routinely checked
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   for compatibility with data, a logical next question is how best to conduct such checks.
   Unfortunately, there is no single best answer. Most texts in ecology (e.g., King et al. 2009,
   Link and Barker 2010, Kéry and Schaub 2012) focus on posterior predictive checks, as
   pioneered by Guttman (1967), Rubin (1981, 1984), and Gelman et al. (1996) (among
   others). These procedures are also the main focus of popular Bayesian analysis texts (e.g.,
   Cressie and Wikle 2011, Gelman et al. 2014) and are based on the intuitive notion that
   data simulated from the posterior distribution should be similar to the data one is
   analyzing. However, "Bayesian p-values" generated from these tests tend to be
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   conservative (biased toward 0.5) because the data are used twice (once to fit the model and
   once to test the model; Bayarri and Berger 2000, Robins et al. 2000). Depending on the
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   data, the conservatism of Bayesian p-values can be considerable (Zhang 2014) and can be
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   accompanied by low power to detect lack-of-fit (Yuan and Johnson 2012, Zhang 2014). By
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sampled posterior p-values, cross-validated probability integral transforms, and pivot 106 discrepancy measures) may produce more accurate characterizations of model fit. 107 In this monograph, we have collated relevant statistical literature with the goal of 108 providing ecologists with a practical guide to Bayesian model checking. We start by 109 defining a consistent notation that we use throughout the paper. Next, we work to compile 110 a bestiary of Bayesian model checking procedures, providing positives and negatives 111 associated with each approach. We illustrate Bayesian model checking using several 112 simulation studies (including spatial regression and N-mixture models), as well as a case 113 study involving capture-recapture sampling of adult female fur seals (Callorhinus ursinus) 114 on a rookery in Alaska, U.S.A., and estimating abundance and distribution of sea otters in Glacier Bay, Alaska, U.S.A. We conclude with several recommendations on how model 116 checking results should be presented in the ecological literature. 117

contrast, other approaches less familiar to ecologists (such as prior predictive checks,

Background and notation

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Before describing specific model checking procedures, we first establish common notation.

Bayesian inference seeks to describe the posterior distribution, $[\boldsymbol{\theta}|\mathbf{y}]$, of model parameters, $\boldsymbol{\theta}$, given data, \mathbf{y} . Throughout the paper, we use bold lowercase symbols to denote vectors.

Matrices are represented with bold, uppercase symbols, while roman (unbolded) characters are used for scalars. The bracket notation '[...]' denotes a probability distribution or mass function, and a bracket with a vertical bar '[' denotes that it is a conditional probability distribution.

The posterior distribution is often written as

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$$[\boldsymbol{\theta}|\mathbf{y}] = \frac{[\mathbf{y}|\boldsymbol{\theta}][\boldsymbol{\theta}]}{[\mathbf{y}]}, \tag{1}$$

likelihood), $[\theta]$ denotes the joint prior distribution for parameters, and [y] is the marginal 128 distribution of the data. In Bayesian computation, the denominator [y] is frequently ignored because it is a fixed constant that does not affect inference (although it is needed 130 when computing Bayes factors for model comparison and averaging; Link and Barker 131 2006). The exact mechanics of Bayesian inference are well reviewed elsewhere (e.g., King 132 et al. 2009, Link and Barker 2010, Hobbs and Hooten 2015), and we do not attempt to 133 provide a detailed description here. For the remainder of this treatment, we assume that the reader has familiarity with the basics of Bayesian inference, including Markov chain 135 Monte Carlo (MCMC) as a versatile tool for sampling from $|\theta|y|$. 136 In describing different model checking procedures, we often refer to data simulated 137 under an assumed model. We use \mathbf{y}_i^{rep} to denote a single, simulated dataset under the 138 model that is being checked. In some situations, we may indicate that the dataset was simulated using a specific parameter vector, $\boldsymbol{\theta}_i$; in this case, denote the simulated dataset 140 as $\mathbf{y}_i^{rep}|\boldsymbol{\theta}_i$. We use the notation $T(\mathbf{y},\boldsymbol{\theta})$ to denote a discrepancy function that is dependent 141 upon data and possibly the parameters θ . For instance, we might compare the discrepancy $T(\mathbf{y}, \boldsymbol{\theta})$ calculated with observed data to a distribution obtained by applying $T(\mathbf{y}^{rep}, \boldsymbol{\theta})$ to 143 multiple replicated data sets. Examples of candidate discrepancy functions are provided in Table 4.

where $[\mathbf{y}|\boldsymbol{\theta}]$ is the assumed probability model for the data, given parameters (i.e., the

Model Checking Procedures

Our goal in this section is to review relevant Bayesian model checking procedures for typical models in ecology, with the requirement that such procedures be accessible to 148 statistics-savvy ecologists. As such, we omit several approaches that have good statistical 149 properties but have been criticized (e.g., Johnson 2007b, Zhang 2014) as too 150 computationally intensive, conceptually difficult, or problem-specific to be of relevant use in 151 common applications. For instance, we omit consideration of double sampling methods that 152 may increase the computational burden of a Bayesian analysis by an order of magnitude 153 (Johnson 2007b), including "partial posterior" and "conditional predictive" p-values (see 154 e.g., Bayarri and Berger 1999, Robins et al. 2000, Bayarri and Castellanos 2007).

Prior predictive checks

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Box (1980) argued that the hypothetico-deductive process of scientific learning can be
embodied through successive rounds of model formulation and testing. According to his
view, models are built to represent current theory and an investigator's knowledge of the
system under study; data are then collected to evaluate how well the existing theory (i.e.,
model) matches up with reality. If necessary, the model under consideration can be
amended, and the process repeats itself.

From a Bayesian standpoint, such successive rounds of estimation and criticism can be
embodied through posterior inference and model checking, respectively (Box 1980). If one
views a model, complete with all its set of assumptions and prior beliefs, as a working
model of reality, then data simulated under a model should look similar to data gathered in
the real world. This notion can be formalized through a prior predictive check, where

replicate data \mathbf{y}^{rep} are simulated via

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$$\boldsymbol{\theta}^{rep} \sim [\boldsymbol{\theta}]$$
 (2)
$$\mathbf{y}^{rep} \sim [\mathbf{y}|\boldsymbol{\theta}^{rep}]$$

and then compared to observed data y via a discrepancy function (Appendix A, Alg. 1). When the prior distribution(s) $[\theta]$ are proper statistical distributions, p-values from 170 prior predictive checks are uniformly distributed under the null model and have properly 171 stated frequentist properties. The main problem with this approach is that the models 172 being considered need to have considerable historical investment and proper prior 173 distributions informed by expert opinion or data from previous studies. In our experience, 174 when Bayesian inference is employed in ecological applications, this is not often the case. Still, prior predictive checks may be useful for hierarchical models that serve as an 176 embodiment of current theory about a study system (e.g., population or ecosystem 177 dynamics models). Alternatively, a subset of data (test data) can be withheld when fitting 178 a model, and the posterior distribution $[\theta|y]$ can be substituted for $[\theta]$ in Eq. 2. If used in 179 this manner, prior predictive checks can be viewed as a form of cross validation, a subject we shall examine in a later subsection (see *Cross-validation tests*). 181 Prior predictive checks appear to have found little use in applied Bayesian analysis 182 (but see Dev et al. 1998), at least in the original form proposed by Box (1980). However, 183 they are important as historical precursor to modern day approaches to Bayesian model 184 checking. Further, several researchers have recently used discrepancy measures calculated on prior predictive data sets to help calibrate posterior predictive (e.g., Hiort et al. 2006) 186 or joint pivot discrepancy (Johnson 2007a) p-values so that they have a uniform null

distribution. These calibration exercises are not conceptually difficult, but do have a high computational burden (Yuan and Johnson 2012). The properties (e.g., type I error probabilities, power) of p-values produced with these methods also depend critically on the similarity of the real world data-generating process with the prior distributions used for calibration (Zhang 2014).

Posterior predictive checks

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Posterior predictive checks are the dominant form of Bayesian model checking advanced in 194 statistical texts read by ecologists (e.g., King et al. 2009, Link and Barker 2010, Kéry and 195 Schaub 2012, Gelman et al. 2014). Although sample size was small (n=25), our survey of recent Ecology volumes indicated that posterior predictive checks are also the dominant 197 form of Bayesian model checking being reported in ecological literature (if any checking is 198 reported at all; Fig. 2). Posterior predictive checks are based on the intuition that data 199 simulated under a fitted model should be comparable to the real world data the model was 200 fitted to. If observed data differ from simulated data in a systematic fashion (e.g., excess zeros, increased skew, lower kurtosis), it is good indication that model assumptions are not 202 being met. 203 Posterior predictive checks can be used to look at differences between observed and 204 simulated data graphically, or can be used to calculate "Bayesian p-values" (Appendix A, 205 Alg. 2). Bayesian p-values necessarily involve application of a discrepancy function, $T(\mathbf{y}, \boldsymbol{\theta})$, for comparing observed and simulated data. There are several omnibus 207 discrepancy measures that can be employed to examine overall lack-of-fit, and targeted 208

discrepancy measures can be used to look for specific data features that systematically

differ between simulated and observed data (Table 4).

Posterior predictive checks are straightforward to implement. Unfortunately, Bayesian 211 p-values based on these checks tend to be conservative in the sense that the distribution of 212 p-values calculated under a null model (i.e., when the data generating model and 213 estimation model are the same) tends to be dome shaped instead of the uniform 214 distribution expected of frequentist p-values (Robins et al. 2000). This feature arises 215 because data are used twice: once to approximate the posterior distribution and to simulate the reference distribution for the discrepancy measure, and a second time to 217 calculate the tail probability (Bayarri and Berger 2000). As such, the power of posterior 218 predictive Bayesian p-values to detect significant differences in the discrepancy measure is 219 low. Evidently, the degree of conservatism can vary across data, models, and discrepancy 220 functions, making it difficult to interpret or compare Bayesian p-values across models. In a simulation study with two different model types, Zhang (2014) found that posterior 222 predictive p-values almost never rejected a model, even when the model used to fit the data 223 differed considerably from the model used to generate it. 224

Another possible criticism of posterior predictive checks is that they rely solely on
properties of simulated and observed data. Given that a lack of fit is observed, it may be
difficult to diagnose where misspecification is occurring within the modeling hierarchy (e.g.,
poorly specified priors, errant mean structure, underdispersed error distribution). Further,
a poorly specified mean structure may still result in reasonable fit of the model if the
model is made sufficiently flexible through inclusion of random effects.

These cautions do not imply that posterior predictive checks are completely devoid of value. Indeed, given that tests are conservative, small (e.g., < 0.05) or very large (e.g., > 0.95) p-values strongly suggest lack-of-fit. Further, graphical displays (see *Graphical techniques*) and targeted discrepancies (Table 4) may help pinpoint common assumption

violations (e.g., lack of independence, zero inflation, overdispersion). However, it is often
less clear how to interpret p-values and discrepancies that indicate no (or little) lack-of-fit.
P-values close to 0.15 or 0.25 are especially problematic. In these cases, it seems necessary
to conduct simulation-based exercises to determine the range of p-values that should be
regarded as extreme, and to possibly calibrate the observed p-value with those obtained in
simulation exercises (e.g., Dey et al. 1998, Hjort et al. 2006).

Some practical suggestions may help to reduce the degree of conservatism of posterior 241 predictive p-values. Lunn et al. (2013) suggest that the level of conservatism depends on 242 the discrepancy function used; discrepancy functions that are solely a function of simulated 243 and observed data (e.g., proportion of zeros, distribution of quantiles) may be less 244 conservative than those that also depend on model parameters (e.g., summed Pearson residuals). Similarly, Marshall and Spiegelhalter (2003) suggest reducing the impact of the 246 double use of data by iteratively resimulating random effects when generating posterior predictions for each data point, a procedure they term a "mixed predictive check" (also 248 called "ghosting"). For an example of this latter approach, see Spatial models for count 240 data.

Sampled posterior p-values

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Posterior predictive checks involve cyclically drawing parameter values from the posterior distribution (i.e., $\boldsymbol{\theta}_i \sim [\boldsymbol{\theta}|\mathbf{y}]$) and then generating a replicate dataset for each i, $\mathbf{y}_i^{rep} \sim [\mathbf{y}|\boldsymbol{\theta}_i]$, to compute the reference distribution for a discrepancy test statistic (Gelman et al. 2004, ; Appendix A, Alg. 2). Alternatively, one can simulate a single parameter vector from the posterior, $\tilde{\boldsymbol{\theta}} \sim [\boldsymbol{\theta}|\mathbf{y}]$, and then generate replicate datasets conditional on this parameter vector alone (i.e., $\mathbf{y}_i^{rep} \sim [\mathbf{y}|\tilde{\boldsymbol{\theta}}]$), otherwise calculating the p-value in the

same manner. This choice may seem strange because the resulting p-value can vary depending upon the posterior sample, $\hat{\theta}$, but a variety of theoretical arguments (e.g., 250 Johnson 2004; 2007a, Yuan and Johnson 2012, Gosselin 2011) and several simulation 260 studies (e.g., Gosselin 2011, Zhang 2014) suggest that it may be a preferable choice, both 261 in terms of Type I error control and power to detect lack-of-fit. In fact, sampled posterior 262 p-values are guaranteed to at least have an asymptotic uniform distribution under the null (i.e., when the model fit to the data is the "true" model Gosselin 2011). Sampled posterior 264 p-values can also be calculated using pivotal discrepancy measures, reducing computational 265 burden (i.e., eliminating the requirement that replicate datasets be generated). We 266 describe an example of this approach in Spatial models for count data. 267

Pivotal discrepancy measures (PDMs)

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In addition to overstated power to detect model lack-of-fit, posterior predictive p-values are limited to examining systematic differences between observed data and data simulated under a hypothesized model. As such, there is little ability to examine lack-of-fit at higher levels of modeling hierarchy. One approach to conducting goodness-of-fit at multiple levels of the model is to use discrepancy functions based on pivotal quantities (Johnson 2004, Yuan and Johnson 2012). Pivotal quantities are random variables that can be functions of data, parameters, or both, that have known probability distributions that are independent of parameters (see e.g., Casella and Berger 1990, section 9.2.2). For instance, if

$$y \sim \mathcal{N}(\mu, \sigma^2)$$

then $z = \frac{y-\mu}{\sigma}$ has a standard $f = \mathcal{N}(0,1)$ distribution. Thus, z is a pivotal quantity in that it has a known distribution independent of μ or σ .

This suggests a potential strategy for assessing goodness-of-fit; for instance, in a
Bayesian regression model

$$\mathbf{y} \sim \mathcal{N}(\mathbf{X}\boldsymbol{\beta}, \sigma^2 \mathbf{I}),$$
 (3)

where **X** represents a design matrix, $\boldsymbol{\beta}$ is a vector of regression coefficients, and **I** is an identity matrix, we might keep track of

$$z_{ij} = \frac{y_i - \mathbf{x}_i' \boldsymbol{\beta}_j}{\sigma_j} \tag{4}$$

for each of $j \in 1, 2, ..., n$ samples from the posterior distribution (i.e., drawing each (β_j, σ_j) pair from $[\boldsymbol{\theta}|\mathbf{y}]$). Systematic departures of z_{ij} from the theoretical $\mathcal{N}(0,1)$ distribution can 284 point to model misspecification. Although we have focused on the data model in Eq. 3, 285 note that the same approach could be used at higher levels of the modeling hierarchy. The advantage of using PDMs is that the reference distribution is known and does not 287 necessarily involve simulation of replicated datasets, \mathbf{y}^{rep} . However, in practice, there are 288 several difficulties with using pivotal quantities as discrepancy measures in Bayesian model 289 checking. First, as with the sampled predictive p-value, p-values using PDMs are only 290 guaranteed to be uniform under the null if calculated with respect to a single posterior 291 parameter draw, $\tilde{\boldsymbol{\theta}} \sim [\boldsymbol{\theta}|\mathbf{y}]$. The joint distribution of PDMs calculated across $i \in 1, 2, \dots, n$ 292 samples from the posterior distribution are not independent because they depend on the 293 same observed data, y (Johnson 2004). As with the Bayesian p-value calculated using a

posterior predictive check, this latter problem can result in p-values that are conservative.

Yuan and Johnson (2012) suggest comparing histograms of a pivotal discrepancy function $T(\mathbf{y}, \boldsymbol{\theta}_i)$ to its theoretical distribution, f, to diagnose obvious examples of model

misspecification. If an omnibus Bayesian p-value is desired, a test can be implemented by

appealing to limiting distributions of order statistics (Johnson 2004), but these tests are

conservative and have low power to detect lack of fit.

A second problem is that, to apply these techniques, one must first define a pivotal 301 quantity and ascertain its reference distribution. Normality assessment is relatively 302 straightforward using standardized residuals (e.g., Eq. 4), but pivotal quantities are not 303 necessarily available for other distributions (e.g., Poisson). However, Yuan and Johnson 304 (2012), building upon work of Johnson (2004) proposed an algorithm based on cumulative distribution functions (CDFs) that can apply to any distribution, and at any level of a 306 hierarchical model (Appendix A, Alg. 3). For continuous distributions, this algorithm 307 works by defining a quantity $w_{ij} = g(y_{ij}, \boldsymbol{\theta})$ (this can simply be $w_{ij} = y_{ij}$) with a known 308 CDF, F. Then, according to the probability integral transformation, $F(\mathbf{w})$ should be 300 uniformly distributed if the modeled distribution function is appropriate. Similarly, for discrete distributions, we can apply a randomization scheme (Smith 1985, Yuan and 311 Johnson 2012) to transform discrete variables into continuously distributed uniform 312 variates. For example, when y_{ij} has integer valued support, we can define

$$w_{ij} = F(y_{ij} - 1|\boldsymbol{\theta}) + u_{ij}f(y_{ij}|\boldsymbol{\theta}),$$

where u_{ij} is a continuously uniform random deviate on (0,1) and F() and f() are the cumulative mass and probability mass functions associated with $[\mathbf{y}|\boldsymbol{\theta}]$, respectively. In this

case, w_{ij} will be uniformly and continuously distributed on (0,1) if the assumed distribution is reasonable; deviation from uniformity can point to model misspecification. 317 We have written the PDM algorithm in terms of the data distribution $[\mathbf{y}|\boldsymbol{\theta}]$ (Appendix 318 A), but the algorithm can be applied (without loss of generality) to any level of a 319 hierarchical model. Further, the algorithm can be applied separately to different categories 320 of mean response (e.g., low, medium, or high levels of predicted responses). These advantages are extremely appealing in that one can more thoroughly test distributional 322 assumptions and look for places where lack-of-fit may be occurring, something that can be 323 difficult to do with posterior predictive checks. We apply this algorithm in Examples and 324 provide R code for applying this approach to generic MCMC data in the R package 325 HierarchicalGOF accompanying this paper (see Software for more information).

Cross-validation tests

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Cross-validation consists of leaving out one or more data points, running an analysis, and 328 seeing how model predictions match up with actual observations. This process is often repeated sequentially for different partitions of the data. It is most often used to examine 330 the relative predictive performance of different models (i.e., for model selection; see e.g. 331 Arlot and Celisse 2010). However, it also possible to use cross-validation techniques to 332 examine model fit and diagnose outlier behavior. The major advantage of conducting tests 333 in this fashion is that there is no duplicate use of data (as with posterior predictive tests or those based on joint PDMs). The major disadvantage is that it can be computationally 335 challenging for complicated hierarchical models. 336

One approach to checking models using cross-validation is the cross-validated

probability integral transform (PIT) test, which has long been exploited to examine the

adequacy of probabilistic forecasts (e.g., Dawid 1984, Früiwirth-Schnatter 1996, Gneiting et al. 2007, Czado et al. 2009). These tests work by simulating data at a set of times or 340 locations, and computing the CDF of the predictions evaluated at the realized data (where realized data are not used to fit the model). This can be accomplished in a sequential 342 fashion for time series data, or by withholding data (as with leave-one-out 343 cross-validation). In either case, divergence from a Uniform(0,1) distribution is indicative of a model deficiency. In particular, a U-shape suggests an underdispersed model, a dome 345 shape suggests an overdispersed model, and skew (i.e., mean not centered at 0.5) suggests bias. Congdon (2014) provides an algorithm for computing PIT diagnostic histograms for 347 both continuous and discrete data in Bayesian applications (see Appendix A, Alg. 4). 348 Cross-validation can also be useful for diagnosing outliers in spatial modeling applications. For instance, Stern and Cressie (2000) and Marshall and Spiegelhalter (2003) 350 use it to identify regions that have inconsistent behavior relative to the model. Such 351 outliers can either indicate that the model does not sufficiently explain variation in 352 responses, that there are legitimate "hot spots" worthy of additional investigation 353 (Marshall and Spiegelhalter 2003), or both. For certain types of data sets and models it is possible to approximate leave-one-out 355 cross validation tests with a single sample from the posterior distribution. For instance, in 356 random effects models, importance weighting and resampling can be used to approximate 357 the leave-one-out distribution (Stern and Cressie 2000, ?). Similarly, Marshall and 358 Spiegelhalter (2007) use a procedure known as "ghosting" to resimulate random effects and thereby approximate the leave-one-out distribution. When applicable, such approaches can 360 lead to well stated frequentist properties (i.e., a uniform distribution of p-values under the null; ?).

Residual tests

Lunn et al. (2013) suggest several informal tests based on distributions of Pearson and deviance residuals. These tests are necessarily informal in Bayesian applications, as residuals all depend on $\boldsymbol{\theta}$ and are thus not truly independent as required in unbiased application of goodness-of-fit tests. Nevertheless, several rules of thumb can be used to screen residuals for obvious assumption violations. For example, standardized Pearson residuals for continuous data,

$$r_i = \frac{y_i - E(y_i|\boldsymbol{\theta})}{\sqrt{\operatorname{Var}(y_i|\boldsymbol{\theta})}},$$

should generally take on values between -2.0 and 2.0. Values very far out of this range
represent outliers. Similarly, for the Poisson and binomial distributions, an approximate
rule of thumb is that the mean saturated deviance should approximately equal sample size
for a well fitting model (Lunn et al. 2013).

For time series, spatial, and spatio-temporal models, failure to account for autocorrelation can result in bias and overstated precision (Lichstein et al. 2002). For this 375 reason, it is important to look for evidence of residual spatio-temporal autocorrelation in 376 analyses where data have a spatio-temporal index. There are a variety of metrics to 377 quantify autocorrelation, depending upon the ecological question and types of data 378 available (e.g., Perry et al. 2002). For Bayesian regression models, one versatile approach is to compute a posterior density associated with a statistic such as Moran's I (Moran 1950) 380 or Getis-Ord G* (Getis and Ord 1992) on residuals. For example, calculating Moran's I for 381 each posterior sample j relative to posterior residuals $\mathbf{Y} - \mathrm{E}(\mathbf{Y}|\boldsymbol{\theta}_j)$, a histogram of I_j 382 values can be constructed; substantial overlap with zero suggests little evidence of residual

spatial autocorrelation. As calculation of Moran's I is dependent upon a a pre-specified
distance weighting scheme, investigators might simulate a posterior sample of Moran's I at
several different choices of weights or neighborhoods to evaluate residual spatial
autocorrelation at different scales.

Just build a bigger model! Tradeoffs between fit and prediction

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important.

One way to ensure a model fits the data is simply to build a model high complexity. To take an extreme example, one could simply start with a saturated model (one where there 390 is a separate parameter for each datum) so that the model fits the data perfectly. No one 391 would actually do this in practice; science proceeds be establishing generalities, and there is no generality implicit in such a model. Further, there is no way to predict future outcomes 393 with such a model. Indeed, models with high complexity can fit the data well, but may 394 have poorer predictive ability than a model of lower complexity (Burnham and Anderson 395 2002, Hooten and Hobbs 2015). 396 When unsure of the desirable level of complexity or number of predictive covariates to 397 include in a model, one approach is to fit a number of different models and to average 398 among the models according to some criterion (see, e.g., Green 1995, Hoeting et al. 1999, Link and Barker 2006). Still, unless one conducts model checking exercises, there is no 400 assurance that any of the models fit the data. Further, there are costs to using this 401 approach, especially in Bayesian applications where considerable effort is needed to implement an appropriate algorithm. In such cases, it may make more sense to iterate on a 403 single model (Ver Hoef and Boveng 2015), and thus, model checking becomes even more

Graphical techniques

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Many of the tests described previously require discrepancy functions, and it may be 407 difficult to formulate such functions for different types of lack-of-fit (e.g., Table 2). Many scientists are visual learners, and displaying model checking information graphically can 409 lead to more rapid intuition about where models fit or do not fit the data. Alternative 410 plots can be made for each type of model checking procedure (e.g., posterior predictive checks, sampled predictive checks, or even PDMs). For instance, Gelman et al. (2014) 412 argues that residual and binned residual plots can be instructive for revealing patterns of 413 model misspecification. In spatial problems, maps of residuals can be helpful in detecting whether lack-of-fit is spatially clustered. The types of plots that are possible are many and 415 varied, so it is difficult to provide a comprehensive list in this space. However, we illustrate 416 several types of diagnostic plots in the following examples. 417

Computing

We conduct all subsequent analyses using a combination of R (R Development Core Team
2015) and JAGS (Plummer 2003). We used R to simulate data and to conduct model
testing procedures; JAGS was used to conduct MCMC inference and produce posterior
predictions. We developed an R package, HierarchicalGOF, that contains all of our code.
This package is publicly available at
https://github.com/pconn/HierarchicalGOF/releases, and will be published to a
permanent repository following manuscript acceptance. The code is predominantly
model-specific; however, we hope it can be used as a template for ecologists conducting
their own model checking exercises.

SIMULATION STUDIES

We conducted several simulation studies to illustrate application of alternative model

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checking procedures when trying to detect departures from model assumptions. Simulation 430 is extremely useful for illustrating concepts as truth is known, and we can examine the 431 large-scale properties of model testing procedures, including the important case when the same model is used to fit the data as is used to generate them. For example, we can 433 describe the empirical null distribution of Bayesian p-values, which must be uniformly distributed on (0,1) to provide an unbiased test. 435 Simulation has been previously used to assess the properties of alternative Bayesian 436 model checking procedures (e.g., Gosselin 2011, Yuan and Johnson 2012, Zhang 2014), but the problems studied have often been simplistic relative to the types of Bayesian models 438 used in real world ecological applications. In this section, we study the properties of different model checking methods when applied to simulated data sets more typical for 440 ecological inference. First, we examine spatial regression models applied to simulated count data. In this case, we are interested in detecting residual spatial autocorrelation and overdispersion relative the Poisson distribution. In our second example, we investigate N-mixture models.

Spatial models for count data

We examined alternative model checking procedures for spatially explicit regression models applied to simulated count data. Such models are often used to describe variation in animal or plant abundance over space and time, and can be used to map abundance distributions or examine trends in abundance (e.g., Sauer and Link 2011, Conn et al.

- relative to the commonly chosen Poisson distribution. In ecological data, several sources of overdispersion are often present, including a greater number of zero counts than expected under the Poisson (zero inflation; Agarwal et al. 2002), and heavier tails than predicted by the Poisson (Potts and Elith 2006, Ver Hoef and Boveng 2007). Another important question is whether there is residual spatial autocorrelation that needs to be taken into account for proper inference (Legendre 1993, Lichstein et al. 2002).
- In this simulation study, we generate count data under a Poisson distribution where
 the true mean response is a function of a hypothetical covariate, spatially autocorrelated
 error, and additional Gaussian noise. Data simulated in this manner can be viewed as
 arising from a spatially autocorrelated Poisson-normal mixture, and can be expected to be
 overdispersed relative to the Poisson, in much the same way that a negative binomial
 distribution (a Poisson-gamma mixture) is. We then examine the effectiveness of
 alternative model checking procedures for diagnosing incorrect model specification, such as
 when spatial independence is assumed. We also study properties of model checking
 procedures when the correct estimation model is specified.
- For a total of 1000 simulation replicates, this study consisted of the following steps:
- 1. Locate n=200 points at random in a square study area \mathcal{A}_1 , where $\mathcal{A}_1 \subset \mathcal{A}_2 \subset \mathbb{R}^2$, and \mathcal{A}_1 and \mathcal{A}_2 are subsets of \mathbb{R}^2 . Call the set of n=200 points \mathcal{S} .
- 2. Generate a hypothetical, spatially autocorrelated covariate \mathbf{x} using a Matérn cluster process on \mathcal{A}_2 (see Appendix B).
- 3. Generate expected abundance for all $s \in \mathcal{S}$ as $\boldsymbol{\mu} = \exp(\mathbf{X}\boldsymbol{\beta} + \boldsymbol{\eta} + \boldsymbol{\epsilon})$, where \mathbf{X} is a two-column design matrix specifying a linear effect of \mathbf{x} , $\boldsymbol{\eta}$ are spatially

- autocorrelated random effects, and ϵ are iid Gaussian errors.
- 4. Simulate count data, $y_i|\mu_i \sim \text{Poisson}(\mu_i)$, at each of the $i \in \{1, 2, \dots, 200\}$ points.
- 5. Fit a sequence of three models to each data set according to the following naming convention:
 - Pois0: Poisson model with no overdispersion

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$$Y_i \sim \text{Poisson}(\exp(\mathbf{x}_i'\boldsymbol{\beta}))$$

• PoisMix: A Poisson-normal mixture with iid error

$$Y_i \sim \text{Poisson}(\exp(\nu_i))$$

$$\nu_i \sim \text{Normal}(\mathbf{x}_i'\boldsymbol{\beta}, \tau_{\epsilon}^{-1})$$

• PoisMixSp: The data-generating model, consisting of a Poisson-normal mixture
with iid and spatially autocorrelated errors induced by a predictive process (cf.

Banerjee et al. 2008):

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$$Y_i \sim \operatorname{Poisson}(\exp(\nu_i))$$

$$\nu_i \sim \operatorname{Normal}(\mathbf{x}_i'\boldsymbol{\beta} + \eta_i, \tau_{\epsilon}^{-1})$$

$$\eta_i = \mathbf{w}_i'\tilde{\boldsymbol{\eta}}$$

$$\tilde{\boldsymbol{\eta}} \sim \mathcal{N}(\mathbf{0}, \boldsymbol{\Sigma})$$

- 6. Finally, a number of model checking procedures were employed on each simulated dataset.
- A depiction of the data generating algorithm (i.e., steps 1-4) is provided in Fig. 3;
 mathematical details of this procedure, together with a description of Bayesian analysis
- methods used in step 5 are provided in Appendix B. As it is the main focus of the paper,
- we next describe model checking procedures (step 6) in greater detail.

Posterior predictive p-values

- For each dataset and estimation model, we several posterior predictive p-values with different discrepancy measures. These included χ^2 , Freeman-Tukey, and deviance-based omnibus p-values, as well as directed p-values examining tail probabilities (Table 4). Tail probabilities were examined by comparing the 95% quantile of simulated and estimated data.
- For the Pois0 model, calculation of posterior predictive p-values was straightforward; posterior predictions (\mathbf{y}^{rep}) were simply simulated from a Poisson distribution, with an expectation that depends on posterior samples of $[\boldsymbol{\beta}|\mathbf{y}]$. For the other two models (i.e.,

PoisMix and PoisMixSp), it was less obvious how best to calculate posterior predictions.

For instance, we identified at least three ways to simulate replicated data, \mathbf{y}^{rep} for

PoisMixSp (Fig. 4). Initial explorations suggested similar performance of predictions

generated via the schematics in Figs. 4A-B, but the approach in Fig. 4B was used in

reported results. We also examined the relative performance of a "mixed predictive check"

(Marshall and Spiegelhalter 2007, ; Fig. 4C) for the PoisMixSp model.

To calculate some of the omnibus discrepancy checks (Table 4), one must also specify a method for calculating the expectation, $E(y_i|\boldsymbol{\theta})$. As with posterior predictions, this calculation depends on what one admits to being a parameter (e.g., are the latent $\boldsymbol{\nu}$ variables part of the parameter set, $\boldsymbol{\theta}$?). We opted to start with the lowest level parameters possible. For instance, for PoisMix we calculate the expectation relative to the parameter set $\boldsymbol{\theta} \equiv \{\boldsymbol{\beta}, \tau_{\epsilon}\}$; as such $E(y_i|\boldsymbol{\theta}) = \exp(\mathbf{x}_i\boldsymbol{\beta} + 0.5\tau_{\epsilon}^{-1})$. For PoisMixSp, we compute the expectation relative to $\boldsymbol{\theta} \equiv \{\boldsymbol{\beta}, \tau_{\epsilon}, \tau_{\eta}\}$, so that $E(y_i|\boldsymbol{\theta}) = \exp(\mathbf{x}_i\boldsymbol{\beta} + 0.5(\tau_{\epsilon}^{-1} + \tau_{\eta}^{-1}))$.

510 Pivotal discrepancy measures

We used Alg. 3 (Appendix A) to conduct PDM tests on each simulated data set and model type. For all models, we assessed fit of the Poisson stage; for the PoisMix and PoisMixSp models, we also applied PDM tests on the Gaussian stage (see e.g., Fig. 5). These tests produce a collection of p-values for each fitted model; one for each posterior sample of parameters (i.e., one for each MCMC iteration). We used the median p-value from this collection to summarize overall PDM goodness-of-fit.

517 Sampled predictive p-values

In addition to the median p-value from applying PDM tests, we also sampled a single PDM p-value at random from each MCMC run. This p-value was used as the sampled predictive p-value for each fitted model.

521 K-fold cross-validation

We used a cross-validation procedure to estimate an omnibus p-value for the PoisMix model, but did not attempt to apply it to the PoisO or PoisMixSp models owing to high computational cost. To improve computational efficiency, we modified Alg. 4 (Appendix A) to use k-fold cross-validation instead of leave-one-out cross-validation. For each simulated dataset, we partitioned data into k = 40 "folds" of m = 5 observations each. We then fit the PoisMix model to each unique combination of 39 of these groups, systematically leaving out a single fold for testing (each observation was left out of the analysis exactly once). We then calculated an empirical CDF value for each omitted observation i as

$$u_i = n^{-1} \sum_{i=1}^n I(y_{ij}^{rep} < y_i) + 0.5I(y_{ij}^{rep} = y_i).$$

Here, $I(y_{ij}^{rep} < y_i)$ is a binary indicator function taking on the value 1.0 if the posterior prediction of observation i at MCMC sample j (y_{ij}^{rep}) is less than the observed data at i.

The binary indicator function $I(y_{ij}^{rep} = y_i)$ takes on the value 1.0 if $y_{ij}^{rep} = y_i$.

According to PIT theory, the u_i values should be uniformly distributed on (0,1) if the model being tested does a reasonable job of predicting the data. For each simulated dataset, we used a χ^2 test (with ten equally space bins) to test for uniformity; the associated p-value was used as an omnibus cross-validation p-value.

Posterior Moran's I for spatial autocorrelation

To test for residual spatial autocorrelation, we calculated a posterior distribution for the Moran's I statistic on residuals for each model fitted to simulated data. For each of $j \in 1, 2, ..., n$ samples from the posterior distribution (e.g., for each MCMC sample), Moran's I was calculated using the residuals $\mathbf{y} - E(\mathbf{y}|\theta_j)$. For Pois0, we set $E(\mathbf{y}|\theta_j) = \exp(\mathbf{X}\boldsymbol{\beta})$; for PoisMix and PoisMixSp, we set $E(\mathbf{y}|\theta_j) = \exp(\boldsymbol{\nu})$.

543 Spatial regression simulation results

Posterior predictive p-values were extremely conservative, with p-values highly clustered near 0.5 under the null case where the data generating model and estimation model were 545 the same (Fig. 6). By contrast, an unbiased test should generate an approximately uniform 546 distribution of p-values under the null. Tests using the median p-value associated with 547 PDMs were also conservative, as were mixed predictive checks and those calculated relative 548 to posterior Moran's I statistics. At least in this example, there did not appear to be much reason to go to the extra effort of computing a mixed predictive check, as they actually 550 appeared slightly more conservative than their posterior predictive counterparts. Posterior 551 predictive checks that depended on parameters in the discrepancy function (e.g., χ^2 , 552 deviance based discrepancies) appeared to be slightly more conservative than those that 553 depended solely on observed and simulated data properties (e.g., the 'tail' discrepancy comparing upper quantiles). In fact, the only p-values that appeared to have good nominal 555 properties were sampled predictive p-values and cross-validation p-values. We did not 556 explicitly quantify null properties of cross-validation p-values, but these should be uniform 557 under the null because the data used to fit and test the model are truly independent in this 558 case.

For the Pois0 model, the mean directed posterior predictive p-value examining tail
probabilities was 0.09 over all simulated data sets; the means of all other p-values
(posterior predictive and otherwise) were < 0.01 for the Pois0 model. As such, all model
checking procedures had high power to appropriately detect the inadequacy of the basic
Poisson model.

For the PoisMix model, only the cross-validation test, the Moran I test, and tests based on PDMs of the Gaussian portion of the model had any power to detect model inadequacy (Fig. 6). Of these, the sampled predictive p-value had higher power than the p-value based on the median PDM. The remaining model checking approaches (notably including those based on posterior predictive checks) had no power to detect model inadequacy (Fig. 6).

EXAMPLES

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The need for closure: N-mixture models

N-mixture models are a class of hierarchical models that use count data collected from repeated visits to multiple sites to estimate abundance in the presence of an unknown 573 detection probability (Royle 2004). That is, counts $y_{i,j}$ are collected during sampling visits 574 $j=1,\ldots,J$, at sites $i=1,\ldots,n$, and are assumed to be independent binomial random 575 variables, conditional on constant abundance N_i and detection probability p; 576 $y_{i,j} \sim \text{Binomial}(N_i, p)$. Additionally, N_i is assumed to be an independent random variable with probability mass function $[N_i|\theta]$ (e.g., Poisson, Negative Binomial, Conway-Maxwell 578 Poisson). The assumption of constant abundance $N_{i,j} = N_i \, \forall j$ is critical for accurate 579 estimates of N_i and p. In practice, this assumption implies that a population at site i is 580 closed with respect to births, deaths, immigration, and emigration, for all replicate

temporal surveys at the site. Violation of this assumption can lead to non-identifiability of the N and p parameters, or worse, posterior distributions that converge, but result in N_i 583 being biased high and p being biased low (Kéry and Royle 2015, Apendix C). Additionally, the expected abundance, λ , will be biased high, and is not necessarily biologically 585 interpretable (e.g., it does not necessarily provide an estimate of the total number of 586 individuals ever associated with a site, c.f. superpopulation; Kéry and Royle 2015). Assessing the closure assumption of N-mixture models can be challenging because 588 scientifically plausible alternative models in which N_i (or λ_i) is non-identifiable or does not even exist, lead to data that are practically indistinguishable from data generated under an 590 N-mixture model (Barker et al. In Review). In practice, the appropriateness of the closure 591 assumption has typically been determined by judgment of the investigators, who assess whether time between replicate surveys is short relative to the dynamics of the system, and 593 whether individual movement is small, compared to the size of sample plots (e.g., Efford and Dawson 2012; but see Dail and Madsen 2011, for a frequentist test of this assumption 595 using a model selection approach). As an alternative, we consider the utility of Bayesian 596 model checking to assess the closure assumption for N-mixture models. We first consider a brief simulated example where truth is known. We then examine real data consisting of 598 counts of sea otters from aerial photographs taken in Glacier Bay National Park, 599 southeastern Alaska. For additional model checking examples for other violations of 600 assumptions of the N-mixture model, including: zero-inflation, extra-Poisson dispersion, 601 extra-binomial dispersion, unmodeled site covariates, and unmodeled detection covariates, see Kéry and Royle (2015, section 6.8). 603

604 Simulation

We examine the most common form of N-mixture model for ecological data,

$$y_{i,j} \sim \text{Binomial}(N_i, p_i),$$

$$N_i \sim \text{Poisson}(\lambda_i),$$

$$\log(\lambda_i) = \mathbf{x}_i' \boldsymbol{\beta},$$

$$\log(t_i) = \mathbf{w}_i' \boldsymbol{\alpha},$$
(5)

where p_i and the expected abundance λ_i depend on covariates \mathbf{w}_i and \mathbf{x}_i , respectively. We used equation (5) to simulate data, with one additional step to induce violation of the closure assumption. We examined a series of eight cases where the closure assumption was increasingly violated by letting

$$N_{i,j} \sim \text{Discrete Uniform}(N_{i,j-1}(1-c), N_{i,j-1}(1+c)),$$
 (6)

for j = 2, ..., J, and $c = \{0, 0.05, 0.10, 0.15, 0.20, 0.25, 0.30, 0.35\}$, where c can be 610 interpreted as the maximum proportion of the population that could move in or out of a site between j-1 and j. For all values of c, we arbitrarily set $\beta = (4,1)'$, and set 612 $\boldsymbol{\alpha} = (1, -1)', i = 1, \dots, n = 300, j = 1, \dots, J = 5$. The covariate matrices **X** and **W** each 613 had dimensions 300×2 , where the first column was all ones, and the second column was 614 generated by sampling from a Bernoulli distribution with probability $0.5 \ \forall i$. We then fit 615 Eq. 5 to the generated data using a Markov Chain Monte Carlo Algorithm (MCMC) 616 written in R. Using the fitted model, we assessed the effectiveness of Bayesian p-values, 617 and sampled predictive p-values for diagnosing the closure assumption. When c=0, the 618

model used to generate the data was the same as the model used to fit the data, and our model checking procedures should indicate no lack of model fit. In all other cases, the 620 closure assumption was violated, with the degree of violation proportional to the value of c. Annotated R code, results, and figures from the simulation are provided in Appendix 3. 622 When the closure assumption was met (c=0), the estimated posterior Results distributions recovered true parameter values well, which was expected (Table 1, Appendix 624 C). The posterior predictive p-value was 0.48, and the sampled predictive p-value was 0.27, 625 suggesting no lack of model fit from either model checking proceedure (Table 1). 626 When the closure assumption was violated (i.e., c > 0), MCMC chains appeared to 627 converge to stationary posterior distributions (Appendix C), and convergence was often supported by Gelman-Rubin diagnostics (Table 1). However, abundance was always 629 overestimated when the closure assumption was violated, and the true abundance value used to simulate the data was always outside estimated 95% credible intervals (Table 1). 631 The posterior predictive p-values did not suggest lack of model fit when c < 0.10, and 632 suggested lack of model fit otherwise (Table 1). The sampled predictive p-value correctly identified violation in the closure assumption (assuming a type I error rate of 0.05) for all 634 values of c, for this simulation (Table 1). The effective sample sizes of the MCMC chains 635 were small due to the autocorrelation between abundance and detection probability in the 636 N-mixture model (Table 1). Mean abundance estimates erroneously increased, with 637 increased violation in the closure assumption, and confidence intervals failed to cover the true abundance value by allowing just 5\% of the population to move in or out of a site 639

between surveys.

\overline{c}	p-value	sppv	Abundance (truth=50,989)	95% CRI	GR	ESS
0.00	0.48	0.27	51,200	(49,295, 53,481)	1.00	3,420
0.05	0.40	1.00	60,047	(56,605, 63,868)	1.00	3,260
0.10	0.00	1.00	81,299	(75,223, 89,601)	1.01	3,194
0.15	0.00	1.00	97,066	(89,149, 104,360)	1.13	3,199
0.20	0.00	0.02	117,624	(108,825, 127,007)	1.03	3,184
0.25	0.00	0.01	119,397	(110,477, 125,992)	1.06	3,206
0.30	0.00	0.00	133,797	(124,194, 141,117)	1.10	3,195
0.35	0.00	0.00	139,951	(133,351, 147,086)	1.00	3,213

TABLE 1. Results of one simulation for examining the effect of the closure assumption on model fit. The notation c represents the maximum proportion of the population that could move in or out of a site between j-1 and j, p-value is the posterior predictive p-value using a χ -squared goodness-of-fit statistic, sppv is the sampled predictive p-value using the sum of variance test statistic, Abundance is the mean of the marginal posterior distribution for total abundance at the 300 sites, the 95% CRI are the 95% credible intervals, GR is the multi-variate Gelman-Rubin convergence diagnostic, and ESS is the effective sample size of 1,000,000 MCMC iterations.

Estimating sea otter detection probability from aerial photographs

Williams et al. (In Press) describe a framework for using aerial photograph data to fit N-mixture models, where photographs are taken such that a subset of images overlap in 643 space. The subset of overlapping images provide temporal replication of counts of individuals at spatial locations that can be used to estimate p in the N-mixture modeling 645 framework. To assess the utility of their approach, Williams et al. (In Press) conducted an aerial survey in Glacier Bay National Park, southeastern Alaska, in which they identified groups of sea otters at the surface of the ocean, flew over the groups of sea otters multiple 648 times, and captured an image of the group of sea otters for each flight over the group. In their study, a primary observer operated the camera, and a secondary observer watched the 650 groups of sea otters to ensure the closure assumption of N-mixture models was met. That 651 is, whether sea otters dispersed out of, or into, the footprint of the photograph among temporal replicates. Of the 21 groups of sea otters that were photographed multiple times, 653 20 groups did not appear to violate the closure assumption based on the secondary

observer's observations. At one site, sea otters began moving for an unknown reason. For analysis, Williams et al. (In Press) omitted the one site that violated the closure 656 assumption, based on the secondary observer's observations. Here, we use Bayesian model 657 checking as a formal method for assessing the closure assumption of two data sets that are 658 used to fit the N-mixture model. The first data set is the complete set of 21 observations 659 initially collected for Williams et al. (In Press), including the site where the secondary observer noted a violation in assumption. The second data set is the data provided by 661 Williams et al. (In Press), Table 1, which omits the problematic site. The full data set is 662 provided in the R package HierarchicalGOF. As in our N-mixture model simulation study 663 above, we used Bayesan p-values and sampled posterior predictive values to check our 664 model. We used each data set to fit the model

$$y_{i,j} \sim \text{Binomial}(N_i, p),$$

$$N_i \sim \text{Poisson}(\lambda_i),$$

$$\lambda_i \sim \text{Gamma}(0.001, 0.001),$$

$$p \sim \text{Beta}(1, 1),$$
(7)

using an MCMC algorithm written in R (Appendix C). The Bayesian p-value for the full
data set (21 sites) was 0.048 and the sampled posterior predictive value was 0.059,
suggesting potential lack of model fit. The Bayesian p-value for the restricted data set used
in Williams et al. (In Press) was 0.5630 and the sampled posterior predictive value was
0.823, suggesting no lack of model fit. These results confirm the results of the secondary
observer who noted a violation of closure while in the field. Thus, model checking
proceedures can provide a formal method for examining the closure assumption of

N-mixture models for our example, and corroborates the auxillary information collected by the secondary observer.

Fur seal capture-recapture

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DISCUSSION

Previous articles in the ecological literature that use Bayesian analysis have tended to focus on prior sensitivity, convergence diagnostics and sometimes model comparison (e.g., DIC or cross-validation) - not as much focus on GOF.

GOF on most general model, then model selection/comparison/averaging (Burnham and Anderson 2002).

Standard regression analysis software In capture-recapture analysis and other areas of ecological statistics, there has been considerable focus on developing procedures to assess goodness-of-fit (e.g., Choquet et al. 2009)

Mean structure vs. dispersion - not always obvious where misspecification occurs.

The purpose of this paper was to provide a general overview of common approaches to
Bayesian model checking and the strengths and weaknesses of each. We do not, however,
claim to be entirely comprehensive. For instance, we limited our discussion to approaches
that are relatively straightforward to implement. There are a variety of options to
producing p-values with good statistical properties provided one has the time and technical
acumen to implement them (e.g., "partial posterior" and "conditional predictive" p-values;
see Bayarri and Berger 1999, Robins et al. 2000, Bayarri and Castellanos 2007). Another
possibility is to use data sets simulated from the prior predictive distribution to study the
realized Bayesian p-values under the null model and calibrate p-values accordingly (Hjort

et al. 2006). For example, if p-values exhibit a dome shaped pattern (c.f., dashed lines in Fig. 6), we might adjust p-value cut off values to be the 5th and 95th quantiles of the realized values. Other approaches to model checking may be useful in more specialized areas of ecology. For instance, Shipley (2009) introduced directional-separation tests for assessing the path structure of directed, acyclic graphs. These tests can be useful for assessing the graph structure of ecological networks.

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TABLES

FIGURE CAPTIONS

867

FIGURE 1. A decision diagram describing the steps we suggest ecologists adopt when 868 reporting the results of Bayesian analyses in the literature, particularly when results will be used for conservation and management or to inform ecological theory. The first step is to 870 formulate reasonable ecological models, ensuring that the model(s) and associated software 871 is free of errors and that convergence to the posterior distribution can be achieved (using Markov chain Monte Carlo, for instance). Following this step, models should be checked 873 against observed data to diagnose possible model misspecification (the subject of this 874 article). Assuming no obvious inadequacies, various model comparison or averaging 875 techniques can be used to compare the predictive performance of alternative models that 876 embody different ecological hypotheses. Finally, we suggest conducting robustness analyses (prior sensitivity analyses, simulation analyses where model assumptions are violated) to 878 gauge the importance of implicit parametric assumptions on ecological inference. 879 FIGURE 2. Type of model checking procedures used in n=31 articles published in the 880 journal Ecology during 2014 and 2015. Articles were found via a Web of Science for articles 881 including the topic "Bayesian" (search conducted 10/1/2015). Six articles were determined to be non-applicable (N/A) because they either (1) were simulation studies, or (2) used 883 approximate Bayesian computation, which is conceptually different than traditional 884 Bayesian inference (see e.g. Beaumont et al. 2002). Of the remaining 25, 20 did not report 885 any model checking procedures. Five articles reported specific model checking procedures, 886 which included a combination of Bayesian cross-validation (Cross.val.), frequentist software (Non-Bayes), posterior predictive p-values (Pp.pval), and posterior predictive 888 graphical checks (Pp,qc). Some articles also investigated prior sensitivity which can be 880 regarded as a form of model checking, but we do not report prior sensitivity checks here.

FIGURE 3. A depiction of how simulated count data are generated. First, a spatially 891 autocorrelated covariate is generated using a Matérn cluster process (A) over a region A_2 . 892 Second, a spatially autocorrelated random effect is simulated according to a predictive process formulation (B), where the parent process occurs at a knot level (C; open circles). 894 The covariate and spatial random effect values combine on the log scale to generate 895 expected abundance (C). Sampling locations (C; small points) are randomly placed over a subregion, A_1 of the study area, where A_1 is defined by the inner box of knot values. 897 Finally, counts are simulated according to a Poisson distribution (D). Note that counts are simulated in $A_1 \subset A_2$ to eliminate possible edge effects. 899 FIGURE 4. Three possible ways of simulating replicate data to calculate posterior 900 predictive p-values for the spatial regression simulation study. Solid boxes indicate 901 parameters or latent variables that occur in the directed graph for observed counts, while 902 dashed boxes indicate posterior predictions. In (A), replicate data (y_i^{rep}) for a given observation i depend only upon the latent variable ν_i , posterior samples of which are 904 available directly from MCMC sampling. In (B), replicate values of ν_i are simulated (ν_i^{rep}) 905 prior to generating posterior predictions. In (C), an example of a "mixed predictive check," spatially autocorrelated random effects are also resimulated (η_i^{rep}) , conditional on the 907 values of random effects at other sites, η_{-i} , and parameters describing spatial 908 autocorrelation (i.e., τ_{η} and ϕ). 909 FIGURE 5. Example computation of a χ^2 discrepancy test using a CDF pivot for a 910 single posterior sample of a Normal-Poisson mixture model (without spatial autocorrelation) fit to simulated count data. In this case, the test focuses on the fit of the 912 the latent variable ν to a Gaussian distribution with mean given by the linear predictor 913 (i.e., $X\beta$) and precision τ as specified in the PoisMix model. The test we employed

partitions the linear predictor based on 20%, 40%, 60%, and 80% quantiles (solid lines), and assesses whether the Gaussian CDF in these ranges is uniformly distributed within five 916 bins. If modeling assumptions are met, there should be a roughly equal number of 917 observations in each bin. For the data presented here, there appears to underpredictions at 918 low and high values of the linear predictor. 910 FIGURE 6. Histogram bin heights showing the relative frequency of 1000 p-values as obtained in the spatial regression simulation study (histograms have 10 bins). The dashed 921 line represents the case where the simulation and estimation model were the same 922 (PoisMixSp). An unbiased test should have a roughly uniform distribution in this case, 923 whereas concave distributions indicate that the test is conservative. A greater frequency of 924 low p-values (e.g., < 0.1) under PoisMix (solid lines) indicate a higher power of rejecting the PoisMix model, a model that incorrectly omits the possibility of residual spatial 926 autocorrelation. The following types of p-values were calculated: k-fold cross-validation 927 ('Cross.val'; PoisMix model only), a mixed predictive p-value using the Freeman-Tukey 928 discrepancy ('Mixed.FT'; PoisMixSp model only), posterior Moran's I ('Moran'), median 920 pivot discrepancy on the Gaussian ('Pivot.Gauss') and Poisson ('Pivot.Pois') parts of the

autocorrelation. The following types of p-values were calculated: k-fold cross-validation ('Cross.val'; PoisMix model only), a mixed predictive p-value using the Freeman-Tukey discrepancy ('Mixed.FT'; PoisMixSp model only), posterior Moran's I ('Moran'), median pivot discrepancy on the Gaussian ('Pivot.Gauss') and Poisson ('Pivot.Pois') parts of the model, a posterior predictive p-value with a χ^2 discrepancy function ('PP.ChiSq'), posterior predictive p-values using a deviance-based discrepancy calculated relative to the Poisson ('PP.Dev.Pois') and Gaussian ('PP.Dev.Gauss') portions of the likelihood, a posterior predictive p-value calculated with the Freeman-Tukey discrepancy ('PP.FT'), a posterior predictive p-value using a 95th quantile discrepancy ('PP.Tail'), and sampled predictive p-values relative the Gaussian ('Sampled.Gauss') and Poisson ('Sampled.Pois') parts of the model.

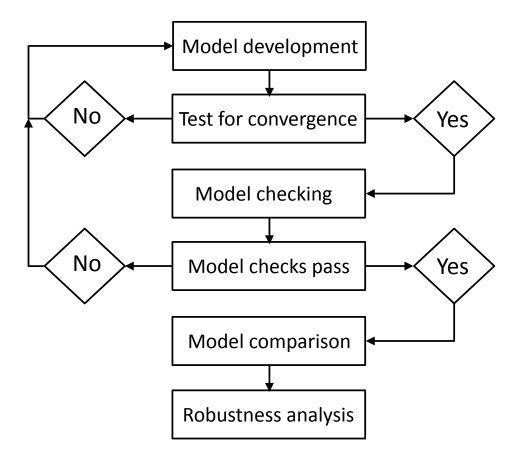


Fig 1

Table 2. Types and causes of lack-of-fit in statistical models $\,$

Concept	Description
Dependent	Many statistical models assume independent response variables. Lack of
responses	independence can have multiple causes, including behavioral coupling
	and unmodeled explanatory variables, with the latter often inducing
	residual spatial or temporal autocorrelation. The usual result is inflated
	sample size, underestimated variance, and overdispersion relative to the
	assumed model.
Over dispersion	Although dependent responses can certainly induce it, the term overdis-
	persion is more a symptom of lack-of-fit, namely that the statistical model
	is incapable of reproducing the amount of variation seen in a data set.
	Three common types of overdispersion in ecological data are (i) unmod-
	eled heterogeneity, (ii) zero inflation in count data (more zero obser-
	vations are obtained than expected under canonical models such as the
	Poisson), and (iii) heavy tails (more extreme observations than predicted
	under the assumed model). The latter is often a result of kurtosis mis-
Higher moments	specification (see <i>Higher moments</i> below).
moments	Overdispersion refers to a misspecification (underestimate) of variance, which is defined as a second moment when studied in terms of moment
	generating functions. However, higher moments may also be misspecified.
	For instance, <i>skewness</i> refers to the third moment and depicts the amount
	of asymmetry of an assumed probability density about its mean; kurtosis
	refers to the fourth moment and to the tail behavior of the distribution.
Outliers	Outliers consist of observations that are surprisingly different than those
	predicted by a statistical model. They can arise because of measurement
	error, or because of model misspecification (particularly with regard to
	kurtosis). Outliers can often have undue influence on the results of an
	analysis (i.e., high leverage), and it may be advantageous to choose mod-
	els that are robust to the presence of outliers.
Nonidentical	Statistical models often assume that responses are identically distributed
distribution	(i.e., have the same underlying probability distribution). However, this
	need not be the case. For instance, <i>Heteroskedasticity</i> refers to the case in
	which variance increases as a function of the magnitude of the response.
Over-	A model is overparameterized whenever two or more combinations of
parameterization	parameters give the same, optimal solution given the data and assumed
	model. If overparameterization is a function of the model only (i.e., could
	not be resolved by collection of more data), a particular parameter set
	is said to be <i>non-identifiable</i> . If it is overparameterized because data
	are too sparse to discriminate between alternative solutions, a particu-
	lar parameter set is said to be <i>non-estimable</i> . Overparameterization can
	be studied analytically or (perhaps more commonly) through numerical
	techniques such as singular value decomposition. It can be difficult to
	diagnose in Bayesian applications because it typically results in a multi-
	modal posterior distribution, and it can be difficult to discern whether
	all the modes have been reached.

Table 3. Discrepancy functions and pivotal quantities useful for hierarchical model checking.

Name	Definition	Comments	
A. Omnibus discrepa	ancy functions		
χ^2	$T(\mathbf{y}, \boldsymbol{\theta}) = \sum_{i} \frac{(y_i - E(y_i \boldsymbol{\theta}))^2}{E(y_i \boldsymbol{\theta})}$	Often used for count data; suggested by Gelman et al. (2014) (among others).	
Deviance (D)	$T(\mathbf{y}, \boldsymbol{\theta}) = -2\log[\mathbf{y} \boldsymbol{\theta}]$	used by King et al. (2009)	
Likelihood ratio statistic	$T(\mathbf{y}, \boldsymbol{\theta}) = 2\sum_{i} y_{i} \log(\frac{y_{i}}{E(y_{i} \boldsymbol{\theta})})$	used by Lunn et al. (2013)	
Freeman-Tukey Statistic	$T(\mathbf{y}, \boldsymbol{\theta}) = \sum_{i} (\sqrt{y_i} - \sqrt{\mathbf{E}(y_i \boldsymbol{\theta})})^2$	Less sensitive to small expected values than χ^2 ; suggested by Kéry and Royle (2016) for count data.	
B. Targeted discrepa	ancy functions		
Proportion of zeros	$T(\mathbf{y}) = \sum_{i} I(y_i = 0)$	Zero inflation check for count data	
Kurtosis checks	$T(\mathbf{y}) = y_{p\%}$	Using the $p\%$ quantile can be useful for checking for proper tail behavior.	
C. Pivotal quantities	5		
$Y \sim \text{Exponential}(\lambda)$	$\lambda \bar{Y} \sim \operatorname{Gamma}(n, n)$	Note n is sample size	
$Y \sim \mathcal{N}(\mu, \sigma^2)$ (Gaussian)	$\frac{Y-\mu}{\sigma} \sim \mathcal{N}(0,1)$	For mean μ and standard deviation σ	
$Y \sim \text{Weibull}(\alpha, \beta)$	$\beta Y^{\alpha} \sim \text{Exponential}(1)$		
Y from any distribution	$Z = \frac{\bar{Y} - \mu}{\sigma / \sqrt{n}} \xrightarrow{\mathcal{D}} \mathcal{N}(0, 1)$	For large sample size (n) , Z converges in distribution to a standard normal (Slutsky's theorem) and Z is tarmed on "computationly"	
		termed an "asymptotically pivotal quantity."	

TABLE 4. A summary of Bayesian model checking approaches. For each method, we describe whether each method allows for (1) computation of an overall p-value ("p-value?"), (2) whether the method tends to be conservative (i.e., has overstated power to detect goodness-of-fit; "conservative?"), (3) whether all levels of the modeling hierarchy can be evaluated ("all levels?"), (4) whether out-of-sample data are used to assess lack-of-fit ("out-of-sample?"), and (5) computing cost ("cost").

Method	p-value?	conservative?	all levels?	out-of-sample?	cost?
Pivotal discrepancy	Yes	Yes	Yes	No	medium
Posterior predictive check	Yes	Yes	No	No	low
Prior predictive check	Yes	No	Yes	No	low
Predictive PIT tests	No	No	No	Yes	very high
Sampled predictive p-value	Yes	No	Maybe	No	low
Graphical	No	Maybe	Yes	No	low-medium

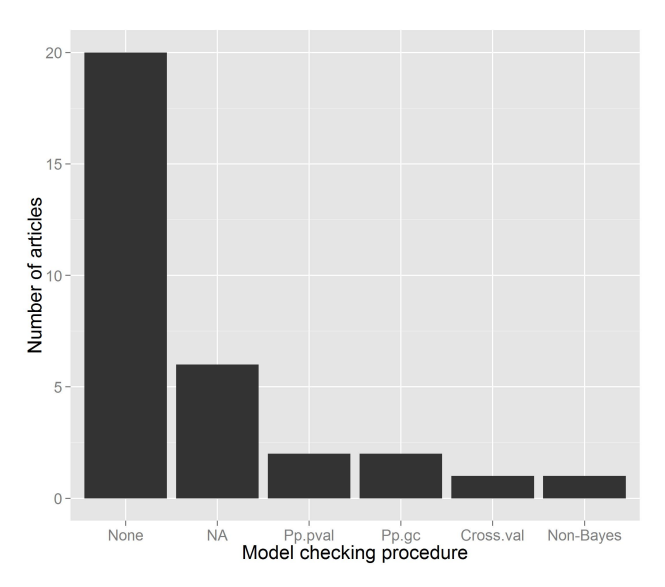


Fig 2

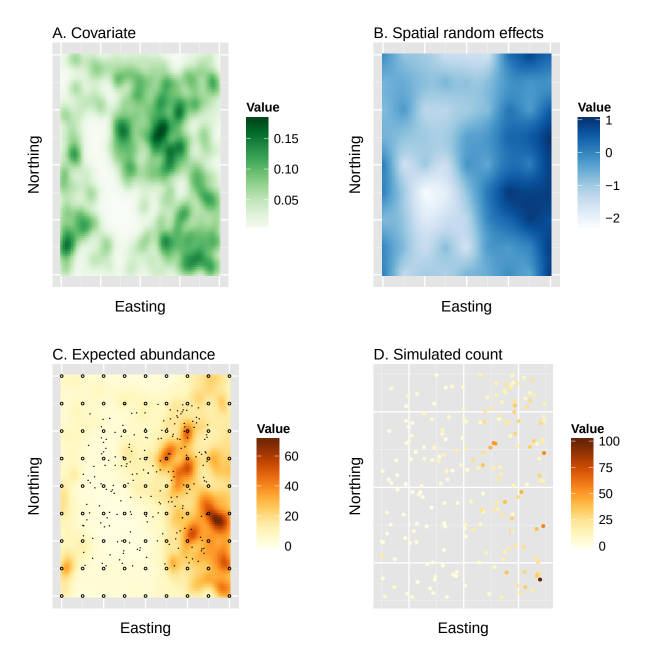


Fig 3

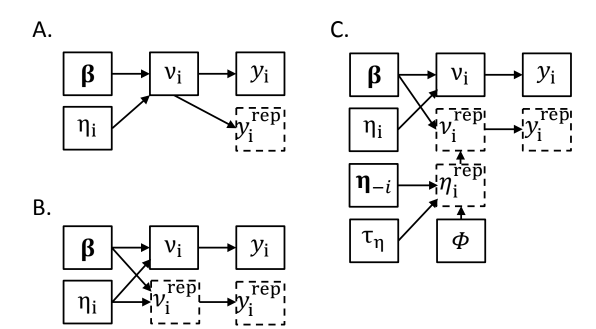


Fig 4

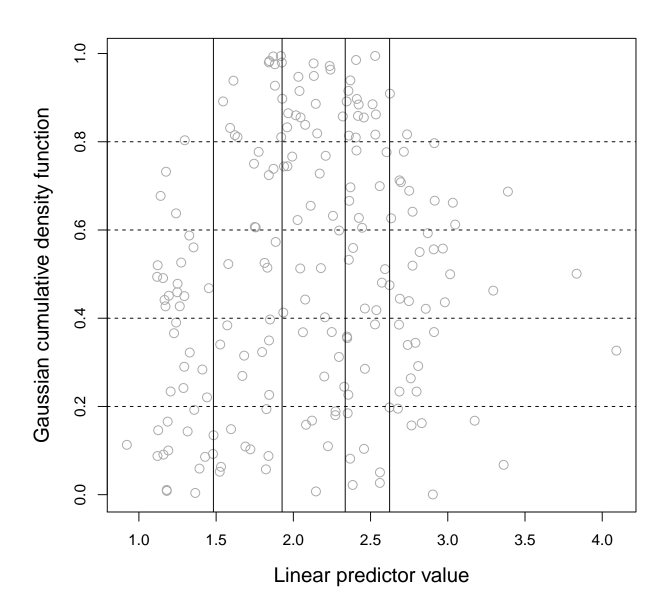


Fig 5

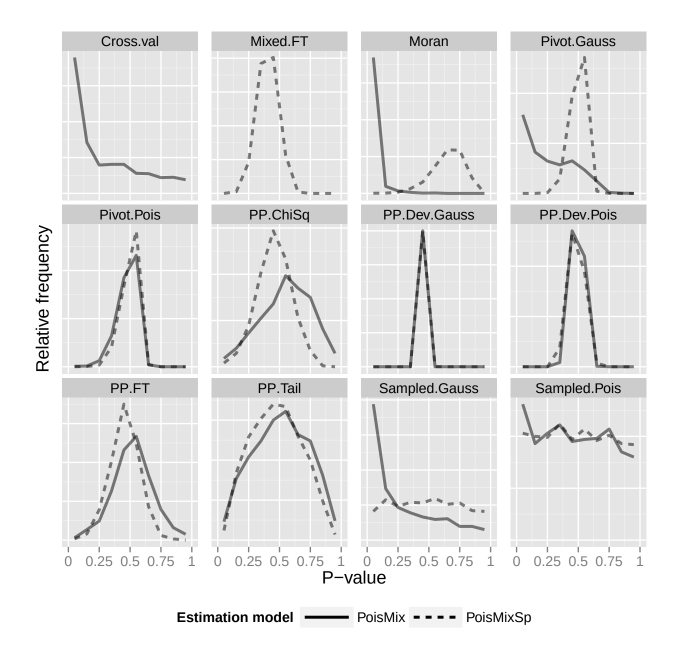


Fig 6