A NEW MODELLING FRAMEWORK FOR ROOST COUNT DATA

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> Roost counts, where individuals of a species are counted whilst arriving or departing from their roost site, are an important monitoring tool for several species around the world. However, the raw count data are an underestimate of the size of the monitored population at any one time because of individuals temporarily not using the roost (temporary emigration, TE) and because the probability of detection of individuals, even when using the roost, is typically much lower than one (observation error). In this paper, we develop a novel modelling framework for estimating population size, from roost count data, while accounting for both TE and observation error. Our framework builds on the popular class of N-mixture models but extends them in a number of ways. Specifically, we introduce two model classes for TE, a parametric, which relies on temporal models, and a non-parametric, which relies on Dirichlet process mixture models. Both model classes give rise to interesting ecological interpretations of the TE pattern while being parsimonious in terms of the number of parameters required to model the pattern. When accounting for observation error, we use mixed-effects models and implement an efficient Bayesian variable selection algorithm for identifying important predictors for the probability of detection. We demonstrate our new modelling framework using an extensive simulation study, which highlights the importance of using mixed-effects models for the probability of detection and illustrates the performance of the model when estimating population size and underlying TE patterns. We also assess the ability of the corresponding variable selection algorithm to identify important predictors under different scenarios for observation error and its corresponding model. When fitted to two motivating data sets of parrots, our results provide new insights into how each species uses the roost throughout the year, on changes in population size between and within years, and on important predictors for observation error.

1. Introduction. The loss of Earth's biological diversity negatively impacts ecosystem services that are vital for human health and prosperity (Cardinale et al., 2012). This global issue is recognised by International agreements and policy frameworks including the Convention on Biological Diversity (CBD) and the United Nations Sustainable Development Goals (SDGs), which call upon all United Nations Member States to take urgent action to restore and protect habitats and to halt further biodiversity loss (sdgs.un.org).

With an increasing number of species suffering population declines (Thomas, 2013; Almond, Grooten and Peterson, 2020), it is paramount to develop innovative monitoring methods in order to characterise population dynamics, understand how environmental changes

Keywords and phrases: N-mixture model, temporary emigration, Bayesian variable selection, Dirichlet process, temporal models, population size.

affect populations, identify species that require protection, and develop or appraise management practices, policies and guidelines (Jetz et al., 2019). However, some highly mobile species such as parrots (Dénes, Tella and Beissinger, 2018) and bats (Kunz, 1982) can be challenging to monitor because they are not individually identifiable and they often feed and nest in low densities among inaccessible habitats such as forest canopies (Dénes, Tella and Beissinger, 2018). Consequently, one of the only opportunities to survey such species at scale is at communal roosts where large numbers of individuals may regularly come together and interact socially for reasons including predator avoidance, cooperative breeding, information exchange, informing foraging strategy and meeting thermoregulatory demands (Kunz, 1982; Beauchamp, 1999; Salinas-Melgoza, Salinas-Melgoza and Wright, 2013; Seixas and Mourao, 2018). During these surveys, individuals are observed and counted as they arrive or depart from their roost, which is a more cost-effective method than others, such as capturemark-recapture, which can be impractical for such species (Kunz, 1982).

In practice, the specific methodology includes multiple simultaneous counts obtained by one or more observers positioned at one or more vantage points. In addition, due to the challenging nature of performing roost counts and the costs and challenges of identifying and accessing a roost, only a single roost is often monitored for a given species, with counts typically collected under different environmental conditions (Cougill and Marsden, 2004; Berg and Angel, 2006; Matuzak and Brightsmith, 2007). These roost survey counts cannot serve as an index of population size due to individuals exhibiting TE, and hence becoming temporarily unavailable for detection, and due to observation error, with the probability of detecting individuals that are available for detection typically being much lower than one. Therefore, statistical modelling needs to be employed for inferring population size and TE patterns from roost count data. This is the aim of this paper, as we describe below.

Count data for closed populations that do not exhibit TE are often analyzed using standard N-mixture models (Royle, 2004), which can estimate population size using spatiallyreplicated counts over time by accounting for observation error. The time-for-space substitution N-mixture model (Kéry and Royle, 2015) uses temporally replicated counts without spatial replication, giving temporal estimates of population size and enabling estimation of a single population trend, but also does not account for TE. However, Chandler, Royle and King (2011) showed that failure to account for TE can result in positively biased estimates of population size.

Roost survey sampling usually takes place under Pollock's robust design (Pollock, 1982), with several short secondary periods, eg days, across various primary periods, eg months. The population size is then assumed constant across secondary periods within the same primary period (closed population) but can change between primary periods (open population) due to births, deaths, immigration, or permanent emigration. In this case, Chandler, Royle and King (2011) extended the standard N-mixture models to account for TE. This model has two processes: an ecological process for the latent number of individuals present and available for detection, and an observation process, for the available individuals detected. The proportions of individuals in the population in any given primary period that are available for detection on each secondary period are either assumed constant for the duration of the study period (Chandler, Royle and King, 2011) or are estimated separately of each other, requiring one parameter to be estimated for each primary period (Kéry and Royle, 2020). However, the first option may be too restrictive and the latter is parameter-greedy, and does not allow for an intuitive ecological interpretation of the results. Finally, existing models do not provide information on TE cyclical patterns, where certain primary periods of each year correspond to certain levels of TE. Identifying and inferring these cyclical patterns can give new insights into the behaviors of the species, such as breeding patterns and seasonal availability of foods.

Naturally, detection probability, and hence observation error (with the two terms used interchangeably in this paper), is expected to vary between sampling occasions as a response

to changes in environmental and weather conditions or effort. This variation can be captured within a logistic regression model accounting for the effect of covariates, such as time of sampling and weather conditions at the time of surveying (see for example Kéry and Royle, 2020; Neubauer et al., 2022). All of the existing modelling approaches can account for the effect of covariates (referred to as variables or predictors in the literature and in this paper) on detection probability through fixed effects models for a given variable set. However, it is unlikely that these fixed effects will capture all of the variation in detection, as other, unobserved or unobservable effects, such as the behaviour of the surveyed species, can have a substantial impact on observation error. As we demonstrate with our simulation study, using fixed-effects models can lead to substantial bias in the estimation of population size when the model for observation error is misspecified, that is when important variables for observation error are omitted, which is likely to be the case in reality. Additionally, the potential set of variables to be considered as predictors for observation error can be large, and hence corresponding tools are required to identify the subset of important variables in the model.

Motivated by two roost count survey data sets of parrots, in this paper we develop a novel modeling framework that can be used to estimate time-varying population size at a site, while accounting for TE and observation error. We extend the TE N-mixture model developed by Chandler, Royle and King (2011) by proposing two model classes: a parametric approach, which employs different temporal models that account for temporal auto-correlation of different order, and a non-parametric approach based on the Dirichlet process (DP) prior (Ferguson, 1973) that allows us to cluster the primary periods according to roost use by the surveyed individuals, and leads to interesting ecological insights about the behavior of the population.

To account for variation in observation error, in addition to that captured by a fixed-effects model, we introduce a mixed-effects logistic regression model on the detection probability. Additionally, we implement a recent efficient Bayesian variable selection (BVS) algorithm, the Bayesian Group Lasso Spike and Slab (BGLSS) (Xu and Ghosh, 2015; Liquet et al., 2017), to perform variable selection for the probability of detection in this mixed-effects model framework.

We implement our novel modelling framework in a Bayesian setting using Markov Chain Monte Carlo (MCMC) methods via R package NIMBLE (de Valpine et al., 2017) version 0.13.0 with the code freely available on https://github.com/Fabian-Ketwaroo/A-novel-modelling-framework-for-roost-count-data.

We present an extensive simulation study that assesses the performance of the proposed models in estimating population size and TE patterns under different scenarios, such as when the model for observation error is misspecified. For the first time in N-mixture models and related literature, we highlight the risks of using misspecified fixed-effects models for observation error and demonstrate how the risks are mitigated by instead using mixed-effects models, as we propose in this paper. We also demonstrate the performance of our proposed variable selection approach in identifying important predictors for observation error in our novel mixed-effects modelling framework under these scenarios.

Finally, we apply our new modelling framework to two case studies, considering roost count data on Ecuadorian Amazon parrots *Amazona lilacina* and on Orange-winged Amazon parrots *Amazona amazonica*. We use cross-validation to select the most appropriate model for the TE pattern in each case and obtain interesting ecological results on temporal population sizes, TE trends, and cyclical patterns and identify important predictors affecting observation error.

The paper is organized as follows. In Section 2 we define our new modelling framework, including background on the methods on which it builds. Simulation results are presented in Section 3 and the results for the two case studies are presented in Section 4. Section 5 concludes the paper and provides ideas for potential future directions.

2. Models. Sampling follows Pollock's robust design (Pollock, 1982) with T open primary periods (e.g. months) and J closed secondary periods (e.g. days within a month). Often, studies can have Y additional top-level primary periods, e.g. Y years, with T primary periods, e.g. months, and J secondary periods, e.g. days within them. The data are summarised in counts $C_{j,t,y}$ of individuals detected on secondary occasion j, primary period t, within top-level primary period y.

We assume there is an overall super-population of M individuals that can visit the roost at least once during the survey period. These M individuals can contribute to the Y superpopulation sizes (κ_y , y = 1, ..., Y), indicating the number of individuals that can visit the roost at least once in each top-level primary period and denote the probability that an individual from the super-population has used the roost at least once in top-level primary period y by δ_y . Conditional on κ_y , we denote the number of individuals using the roost in primary period t within top-level primary period y by $N_{t,y} \sim Bin(\kappa_y, \theta_{t,y})$ (temporal population size), with $\theta_{t,y}$ referred to as the availability parameters (meaning that these individuals are available for detection in that primary period). Finally, individuals that use the roost in primary period t within top-level primary period y are detected on secondary occasion j with probability $p_{j,t,y}$. The hierarchical representation of the model is given in equation (1), while a graphical representation of the model is given in Fig 1.

$$M \sim \text{Poisson}(\lambda)$$

$$\kappa_y \sim \text{Binomial}(M, \delta_y)$$

$$N_{t,y} \sim \text{Binomial}(\kappa_y, \theta_{t,y})$$
(1)
$$C_{j,t,y} \sim \text{Binomial}(N_{t,y}, p_{j,t,y})$$

The κ_y variables allow us to study the availability pattern within each top-level primary period, conditional on the corresponding population size, and hence identify changes in availability patterns across top-level primary periods, without these changes being confounded to changes in population size. When there are no top-level primary periods, this model can be simplified by dropping the κ_y level, i.e. setting $\kappa_y = M \ \forall y$, and the y subscript in all subsequent levels.

The main novelty of our proposed framework lies in the way in which we model detection probability, as described in Section 2.1, and the availability parameters, as described in Section 2.2.

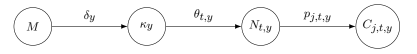


Fig 1: Graphical model representation

2.1. Detection probability. The model of equation (1) is a function of the detection probability on secondary occasion j, primary period t, top-level primary period y, $p_{j,t,y}$. This probability cannot be freely varying, as that introduces more parameters than we can estimate into the model. Instead, it can be assumed as constant for all j, t, y or, more realistically, as a function of variables (covariates), which can vary between secondary and/or primary periods, within a logistic regression framework, as for example in Kéry and Royle (2020). However, it is likely that, in practice, such models are misspecified, and that the variables considered are only a subset of the variables that affect detection probability in the field. In such cases, as we demonstrate in our simulation study in Section 3, the estimation of population size can be substantially biased, and for that reason we propose the use of a mixed effects model:

(2)
$$\operatorname{logit}(p_{j,t,y}) = \eta_{j,t,y} = \mu + \sum_{g=1}^{G} X_{j,t,y,g} \beta_g + \epsilon_{j,t,y}$$

where g = 1, ..., G are continuous/categorical variables, such that variable g requires C_g coefficients to model its effect, so that if g is a continuous variable, $C_g = 1$, and if g is a categorical variable, C_g is its number of levels (excluding baseline). Finally, β_g is the $(C_g \times 1)$ vector corresponding to the logistic regression coefficients for variable g, $X_{j,t,y,g}$ is the vector of length C_g containing variable g on occasion t, y, j, and $\epsilon_{j,t,y} \sim \text{Normal}(0, \sigma_{\epsilon}^2)$ are corresponding independent random effects.

The inclusion of the random effect terms allows for any variability in detection probability that is not captured by the variables considered by the fixed effects to be absorbed by the random effect variance, which, as we demonstrate using simulation, leads to reliable inference on population size, even when the detection probability model is misspecified. However, an overparameterised fixed effects model can lead to increased uncertainty around variable effects and population size, and therefore, we suggest the use of a Bayesian variable selection algorithm, and specifically of the Bayesian Group Lasso Spike-and-Slab (BGLSS) algorithm (Xu and Ghosh, 2015), for identifying important predictor variables for p. The BGLSS places a prior on each group of coefficients, where a group can consist of coefficients introduced to model the effect of a categorical variable and can number a single coefficient in the case of continuous variables. This prior is given in equation (3) below, and more details are provided in the Supplementary material.

$$\beta_{g} | \tau_{g}^{2} \sim (1 - \gamma_{g}) \delta_{0}(\beta_{g}) + \gamma_{g} N(0, \tau_{g}^{2} I_{C_{g}})$$

$$\tau_{g}^{2} \sim \text{Gamma}\left(\frac{C_{g} + 1}{2}, \frac{\psi^{2}}{2}\right)$$

$$\gamma_{g} \sim \text{Bernoulli}(\phi_{g})$$

$$\psi \sim \text{Gamma}(a, b)$$
(3)

where γ_g is a binary variable that indicates whether variable g is included (1) in the model or not (0), $\delta_0(\beta_g)$ denotes a point mass at $0 \in \mathbb{R}^{C_g}$, I_{C_g} is the identity matrix $(C_g \times C_g)$, ψ is the shrinkage parameter, and ϕ_g is the prior inclusion probability, which can be fixed to 0.5 or can be assigned a uniform or Beta prior distribution.

The BGLSS accommodates group-level variable selection by using a spike and slab prior (Mitchell and Beauchamp, 1988), with coefficients exactly zero for excluded variables, and the Bayesian group lasso (BGL) (Casella et al., 2010) for included variables, enforcing the L_1 penalization (Tibshirani, 1996), giving more parsimonious models. This Bayesian formulation can reduce the computational cost by proposing a prior on ψ rather than testing several values and choosing the best value by cross-validation. In addition, the BGLS produces reliable standard errors of coefficients without any extra cost in comparison to the frequentist group lasso (Yuan and Lin, 2006).

2.2. Availability parameters. We propose two model classes for modelling the availability parameters, a nonparametric approach and a parametric approach, both of which are described below. We define $\theta_{\ell} = \theta_{t,y}$, with $\ell = t + T(y-1)$ for $\ell = 1, \ldots, T \cdot Y$ to model correlation in the availability parameters for the whole time series, across primary periods. When

there are no top-level primary periods, Y = 1 and $\theta_{\ell} = \theta_t$ for $\ell = 1, ..., T$. Table 1 provides the terminology used hereafter for each model considered for the availability parameters.

TABLE 1Models proposed for availability parameters.

Notation	Model
DP	Dirichlet process (DP) mixture model
RW1	Random walk of order 1
RW2	Random walk of order 2
Cor	Across level correlation model
AR1	Auto-regressive model of order 1

2.2.1. *Nonparametric approach.* We model availability non-parametrically via a Beta Dirichlet process (DP) mixture model (Kottas, 2006). This formulation expresses the distribution of availability parameters as a mixture model, and provides a flexible and robust specification of the corresponding density, by describing it as a mixture model with an unknown number of components, with primary periods clustered according to their corresponding availability parameters, eg low, medium, and high. This is ecologically relevant as it enables the study of TE trends and hence roost use patterns throughout the season(s).

The Beta DP mixture model can be represented using the Chinese restaurant process (CRP) algorithm, which relies on the inferred cluster allocation variables, z_{ℓ} , $\ell = 1, ..., T \cdot Y$, indicating the cluster to which primary period ℓ has been allocated. The CRP is used to represent the sequential way in which cases, i.e. periods in our case, are allocated to clusters, with the number of clusters being infinite a priori, but finite in practice and inferred as part of the process. The corresponding model for the availability parameters is given in equation (4).

$$\begin{split} \theta_{\ell} | \tilde{\gamma}, \tilde{\psi}, z_{\ell} &\sim \operatorname{Beta}(\tilde{\gamma}_{z_{\ell}}, \tilde{\psi}_{z_{\ell}}), \quad \ell = 1, \dots, (T \cdot Y) \\ z_{\ell} &\sim \operatorname{CRP}(\alpha), \quad \alpha \sim \operatorname{Gamma}(\zeta, \tau) \\ \tilde{\gamma}_{k} &\sim \operatorname{Gamma}(\mu, \nu), \quad \tilde{\psi}_{k} \sim \operatorname{Gamma}(\vartheta, \omega), \quad k = 1, \dots, K. \end{split}$$

(4)

where $\zeta, \tau, \mu, \nu, \vartheta, \omega \in \mathbb{R}$ and $K \leq (T \cdot Y)$.

2.2.2. *Parametric approach.* Alternatively, availability can be modelled parametrically using temporal models, specifically random walk models and auto-regressive models. These temporal models share information across primary periods by accounting for temporal auto-correlation, which is meaningful ecologically as, as also mentioned above, the availability pattern is expected to be smooth and allows for borrowing strength in cases where the data are sparse.

 Random walk models, which enable estimation of non-linear temporal trends retaining the smoothing-varying feature that is present in observed time series data. As highlighted in Fahrmeir and Lang (2001), random walk models can be rewritten in an undirected symmetric form, as a one-dimensional version of the spatial intrinsic conditional autoregressive (ICAR) model (Besag, 1974). Generally, random walk models can be defined as a set of conditional probability distributions under the ICAR models as

(5)
$$\theta_{\ell}|\theta_{-\ell}, \sigma^2, W^{\mathrm{RW}} \sim N\left[\frac{\sum_{n=1}^{T \cdot Y} w_{\ell n} \theta_n}{w_{\ell +}}, \frac{\sigma^2}{w_{\ell +}}\right], \ \ell = 1, \dots, T \cdot Y.$$

where W^{RW} represents the temporal weights matrix with entry $\omega_{\ell n}$ in the ℓ th row and the *n*th column, $w_{\ell+}$ is the sum of the elements in the ℓ th row, σ^2 is the ICAR variance and $\sigma^2/\omega_{\ell+}$ is the conditional variance.

Consequently, random walk models possess the same set of properties as the ICAR model. That is, positive auto-correlation is assumed via a chosen W that imposes a neighbourhood structure on time points in the study period and determines the amount of information borrowed from other time points. This shared information across temporal neighbours results in temporally smooth time trends, with estimation of θ_{ℓ} borrowing information from past time points eg. $(\ell - 1, \ell - 2)$ but also from future time points eg. $(\ell + 1, \ell + 2)$, provided that these time points are within the study period. In addition, as the conditional variance increases, θ_{ℓ} can deviate more from its neighbours, producing a temporal pattern that is less smooth but more flexible. This model representation allows us to infer the variance of the ICAR model (σ^2) and $\theta_{\ell} \forall \ell$.

- Random walk of order 1 (RW1) can be defined as an ICAR model with binary weights, W^{RW1} , such that the entry $\omega_{\ell,n} = 1$ if points ℓ, n are neighbours and 0 otherwise. In the RW1 model, each ℓ has 2 neighbours $\ell 1, \ell + 1$, except the first and the last, which only have one neighbour, adjacent to the right and left respectively. The binary temporal weights matrix, W^{RW1} , assumes that equal strength of information is borrowed from adjacent neighbours.
- Random walk of order 2 (RW2). Similarly, the RW2 model can be defined as an ICAR model but with a general weights matrix (W^{RW2}). The elements in W^{RW2} are derived from the conditional distributions of each θ_{ℓ} conditioned on all other parameters in θ and the variance σ^2 (conditional distributions listed in the Supplementary material). The elements are the coefficients in the numerator of the conditional mean for θ_{ℓ} . As can be seen in equation (5), the conditional variance depends on the number of neighbours, hence, the RW2 model generally produces smoother temporal trends than the RW1 model as it borrows information from more time points. In addition, using a general weights matrix instead of a binary weights matrix specifies the strength of the information borrowed, with more information borrowed from close neighbours.
- Across level correlation (Cor) model. We extend the RW1 model to allow a time point to borrow information from other specific time points, in addition to $\ell - 1, \ell + 1$ time points given time points are within the study period. For instance, this allows a specific month in a year to be correlated to months directly before and after that month, but also the same month across years. This model is defined similarly to the RW1 model with a binary weights matrix (W^{Cor}) such that the entry $\omega_{\ell,n} = 1$ if points ℓ, n are neighbours and 0 otherwise, where neighbours in this case are the adjacent time points, but also time points that are c time periods apart, where c = 12 in the case of monthly patterns across years. Therefore, the first time point are neighbour with ($\ell + 1, \ell + qc$) time points, the last time point with ($\ell - 1, \ell - qc$) neighbours and others with ($\ell - 1, \ell + 1, \ell \pm qc$) for $q = 1, \ldots, ((T \cdot Y)/c) - 1$, provided time points are within the study period.

2. Auto-regressive models. An auto-regressive model of order 1 (AR1) on the set of timespecific parameters can be defined as

(6)
$$\theta_{\ell} = \rho \theta_{\ell-1} + \epsilon_{\ell}, \ \ell = 2, \dots, T \cdot Y,$$
$$\theta_1 \sim \mathcal{N}(0, \sigma_1^2(1 - \rho^2))$$

where ρ is the temporal correlation coefficient ($|\rho| < 1$) and $\epsilon_{\ell} \sim N(0, \sigma^2)$ are iid noise effect terms. The RW1 model is a subset of the AR1 model when $\rho = 1$. As such, the AR1 is a more flexible model as it accommodates both positive and negative temporal autocorrelation. However, if positive auto-correlation is present, the RW1 model is preferable as one fewer parameter needs to be estimated.

2.3. Inference. We fit models in a Bayesian framework using MCMC methods via R package NIMBLE (de Valpine et al., 2017) version 0.13.0 with all code freely available on https://github.com/Fabian-Ketwaroo/A-novel-modelling-framework-for-roost-count-data. Specifically, for variables assigned an ICAR model, we follow NIMBLE's recommendation and update these variables without the zero constraints and then centering (Paciorek, 2009). We implement the Beta mixture DP model by using the collapsed sampler (Neal, 2000) provided in NIMBLE. We use methods developed by Wade and Ghahramani (2018) to summarise DP cluster results. We employ median thresholding in variable selection (Barbieri and Berger, 2004), that is, $Pr(\gamma_q = 1|y) > 0.5$, $g = 1 \dots, G$ to identify significant variables.

3. Simulation study. In this section, we present an extensive simulation study to explore a number of different cases, listed in Table 2. For each case, we perform 50 simulation runs and we set T = 36, J = 8, assuming no top-level primary periods with $\lambda = 100$ and consider high and low detection levels, $p \approx (0.6, 0.3)$, with p as a function of covariates (variables). The coefficients for fixed effects are set as: $\beta = (\beta_1 = 1.25, \beta_2 = 0.2, \beta_3 = 2, \beta_4 = 0, \beta_5 = -0.6, \beta_6 = 0.5, \beta_7 = -1, \beta_8 = 0)$ with the first five corresponding to continuous variables, x_1, \ldots, x_5 , and last three to categorical variables, x_6 and x_7 , with two and three levels, respectively. Continuous variables were generated from a standard normal distribution and categorical variables were from a multinomial distribution with equal probabilities. To obtain the desired level of average detection, as stated above, the intercepts, β_0 , were set to (0.75, -1.5), for high and low detection probability, respectively. To introduce misspecification in the model for detection, variables x_1 and x_7 were not included in the model in each of the two cases described in Table 2. When the DP model was used to generate the data, we specified two clusters of equal size (18) from Beta(10,10) and Beta(10,1) respectively. When the RW1 model was used to generate data, we set $\sigma = 1$.

The following prior distributions were used in all cases: $\lambda \sim \text{Gamma}(0.01, 0.01), \psi \sim \text{Gamma}(0.001, 0.001), \phi_g = 0.5, \beta_1 \sim \text{Normal}(0, 2), \sigma \sim \text{Uniform}(0, 15), \alpha \sim \text{Gamma}(1, 1), \tilde{\gamma}_k \sim \text{Gamma}(2, 0.1), \tilde{\psi}_k \sim \text{Gamma}(2, 0.1)$. The MCMC settings in terms of the number of iterations, burn-in, and thinning in each case are reported in the Supplementary material.

We use median relative bias and median 95% credible interval (CI) coverage to summarise the estimation of population size and covariate effects. We also use median misclassification rate for summarising the DP mixture clustering and the BGLSS performance. The detailed results of the simulation study for each case are presented in the Supplementary material and the key findings are summarised in Table 3 and discussed below.

MODELLING ROOST COUNT DATA

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Simulation settings.

Case	Description
1	Comparing estimation of population size under different models for the availability parameters
	when the correct model for these parameters is fitted to the data and we do not perform variable
	selection and
	a) the model for detection probability is correctly specified.
	b) the model for detection probability is misspecified (fixed vs mixed effects models).
2	Assessing the performance of BGLSS in variable selection under the RW1 model for the availability
	parameters when
	a) the model for detection probability is correctly specified.
	b) the model for detection probability is misspecified (mixed effects model).

TABLE 3

Median relative bias and median coverage of population size and covariate coefficients for each simulation scenario and setting for detection probability, as described in Table 2.

CS: correctly specified; MS: misspecified; FE: fixed effects; ME: mixed effects.

Case	Model for θ	Model for p	Average p	Parameters	RB	Coverage
1. a)						
				Coefficients	0.008	94
	DP	CS - FE	0.6	Population size	-0.001	100
	DI	00 11	0.3	Coefficients	0.002	98
			0.0	Population size	-0.002	98
			0.6	Coefficients	0.003	96
	RW1	CS - FE		Population size	-0.001	98
	KW I	COPIL	0.3	Coefficients	-0.003	96
			0.5	Population size	-0.005	98
1. b)						
			0.6	Coefficients	-0.787	2
	DP	MS - FE	0.0	Population size	8.928	0
	DI	MS - FE —		Coefficients	-0.590	4
			0.3	Population size	4.417	0
			0.6	Coefficients	-0.745	4
	RW1	MS - FE	0.0	Population size	6.066	2
	K W I	M3 - FE	0.3	Coefficients	-0.502	12
			0.5	Population size	3.347	4
			0.6	Coefficients	0.036	94
	DP	MS - ME	0.0	Population size	-0.005	98
	DF	M3 - ME	0.3	Coefficients	0.011	90
			0.5	Population size	-0.019	90
			0.6	Coefficients	0.046	98
	RW1	MS - ME	0.0	Population size	-0.004	100
		1013 - 101E	0.3	Coefficients	0.043	96
			0.0	Population size	0.013	92
2. a)						
			0.6	Coefficients	0.001	96
	RW1	CS - FE	0.0	Population size	-0.001	98
	KW I	C3 - LE	0.3	Fixed effects	-0.006	96
			0.5	Population size	-0.001	98
2. b)				-		
			0.6	Coefficients		
	DW/1			Population size	-0.005	100
	RW1	MS - ME	0.2	Coefficients		
			0.3	Population size	0.031	90
L						

3.0.1. *Case 1.* When the model for detection probability is correctly specified (a), both the DP and the RW1 models perform well in terms of inference, with low median relative bias and high coverage for covariate coefficients and population size. The DP mixture model has a low misclassification rate, on average equal to 0.055 for both levels of detection. In addition, the standard deviation of the RW1 model (σ) is also estimated well with low relative bias (0.011, -0.035) and high coverage (0.98, 1) at high and low levels of detection respectively. Consequently, this scenario shows that both models for the availability parameters perform well in terms of inference when the model for detection probability is correctly specified.

However, when the model for detection probability is misspecified (b) and a fixed effects detection model is used, the estimation of population size is considerably positively biased with very poor coverage in all cases. Similarly, covariate coefficients are estimated with high bias and low coverage and the DP mixture model performs poorly, with a misclassification rate on average equal to (0.111, 0.444) for high and low detection probability, respectively. However, using a mixed effects model for detection probability corrects for the misspecification and produces population size and covariate coefficient estimates with negligible bias and high coverage. The DP mixture model also performs better, with a misclassification rate on average equal to (0.055, 0.111) for high and low detection probability, respectively.

3.0.2. *Case 2.* Similarly, when the model for detection probability is correctly specified (a), BGLSS performs well in identifying both significant (strong and weak) and non-significant effects with mean misclassification rates of 0 across both levels of detection. As such, population size and covariate coefficients are estimated well in all cases.

When the model for detection probability is misspecified (b) and a mixed effects detection model is employed, BGLSS has, as expected, lower power to identify weak effects ($\beta_2 =$ 0.2) with average misclassification rate (0.38, 0.4) at high and low detection probability, respectively, but still high power to identify strong effects with average misclassification rate 0 at both levels of detection. In addition, the power to identify non-significant variables also declines, with a mean misclassification rate (0.1, 0.06) at high and low detection levels respectively. However, importantly, inference on population size is unaffected in all cases when mixed effects models for detection probability are employed.

4. Case studies.

4.1. *Ecuadorian Amazon parrots.* We consider roost count data collected as part of an ongoing conservation project for the Ecuadorian Amazon parrot (*Amazona lilacina*) in Ecuador (Biddle et al., 2020, 2021a,b). Counts were obtained from a single site close to the El Salado Mangrove Reserve, where parrots roost overnight, for 36 consecutive months between 2016 and 2019. Each year, surveys took place between November and October, with surveys taking place on three to five days within each month, and two surveys being performed each day, AM and PM. We assume that the population is closed within each month, but open between months.

We model the data using the model defined in equation (1), fitting all models listed in Table 1 and using k-fold cross-validation to select the most appropriate model for the availability parameters. In each case, we consider a mixed effects model for detection probability, and perform variable selection via BGLSS, considering the following variables: median temperature, average relative humidity, visibility, average wind speed, rain/drizzle, storm/thunder (taken from the Simon Bolivar weather station approximately 14km from the roost site (https://www.tutiempo.net/clima/01-1999/ws-842030.html), time of sampling (AM/PM), and weather recorded by the observer at the roost site (clear, cloud, rain, sunshine). The prior distributions were set as described in the simulation study.

MODELLING ROOST COUNT DATA

k-fold cross-validation was performed by splitting the data into monthly subsets (k = 36) and using root mean square error (RMSE) to evaluate the predictive accuracy of the models considered when leaving one month out at a time. Cor was selected as the model with the lowest RMSE, as seen in Table 4. Cor, RW1, and DP are the top three models, having similar RMSE values. Notably, all these models considered produced similar estimates of population size, BVS results, and model fit. Consequently, we display the results obtained from the Cor model in the paper, while the results obtained from the other models are presented in the Supplementary material, with the exception of the DP model clustering results, which are shown in Table 5 and discussed as they provide us with new insights about the use of the roost throughout and across years.

TABLE 4								
Ecuadorian Amazon parrots case study. Cross-validation results.								
Model DP RW1 RW2 Cor AR1								

Model	DP	RW1	RW2	Cor	AR1
RMSE	41.079	40.586	41.188	39.590	41.999

Fig 2a shows posterior summaries of the month-specific population sizes, N_1, \ldots, N_{36} , obtained from the Cor model. The pattern suggests two peaks in the year, January/February/March and then June/July/August. The first peak, which is more consistent across years, could represent chicks fledging and returning to the roost with the adults, while the second peak, which varies more between years, could represent social gathering before the breeding season, giving opportunities for time to create breeding pairs and highlighting the importance of these communal roosts for the formation of new breeding pairs.

We assessed the fit of models using posterior predictive goodness of fit. For that, we define *monthly rate* to be the sum of the counts obtained in a month divided by the number of surveys in that particular month. Using MCMC samples, we simulated counts, and hence rates, from our models and compared these to the observed rates. Fig 2b displays that the Cor model fits the data well as it produces similar monthly rates to the observed rates, with the true values falling within the 95% posterior credible interval of simulated values and with no consistent pattern of bias observed.

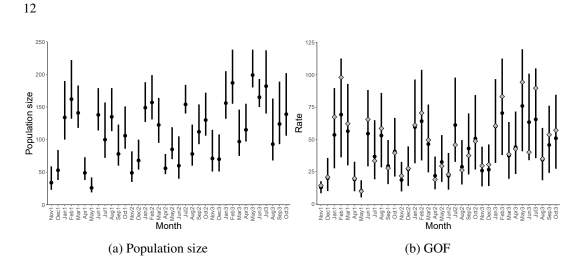


Fig 2: Ecuadorian Amazon parrots case study. (a) The black dots represent the posterior mean population size for each month and the thick bands represent the corresponding 95% posterior credible interval. (b) The diamonds are the observed monthly rates and the thick bands represent the 95% intervals of simulated monthly rates. In both cases, the x-axis represents the months in each year with months ending in 1, 2, and 3 denoting months in the 1st, 2nd, and 3rd year, respectively.

The results of the Cor model are consistent with the clustering output of the DP model (Table 5), where two clusters of equal size (18) have been identified for each year. These correspond to months with low (L) and months with high (H) availability probabilities, with the clustering pattern fairly consistent across years and agreeing with the general trend identified by the Cor model. Locating and observing individual nests for this species can be difficult, and hence this clustering pattern of the overall roosting population provides supportive evidence to reports of seasonal breeding behaviour. The first peak corresponds with months when chicks fledge from nests (January / February / March) and so is likely to represent population recruitment, whilst the second peak in October occurs just before breeding pairs start to nest together in the dry forest and could represent an increase in attendance at the social roost to form or strengthen pair bonds. Due to the fluctuating nature of this particular roost site, accounting for detection probability allows us to identify robust patterns for ecological interpretation that would not be visible clearly in the raw data, helping conservation managers to determine breeding phenology more broadly so that efforts can be more focused on finding nest cavities and documenting breeding success at the right time of year. In other Amazon parrot species roost attendance is also linked with food availability (i.e. in times of food scarcity, roost attendance is greater to allow information sharing) so it is also possible that fluctuating food availability in this seasonal climate may drive high/low distinction.

 TABLE 5

 Ecuadorian Amazon parrots case study. Cluster allocations from the DP model.

			Months										
[Year	Nov	Dec	Jan	Feb	Mar	Apr	May	Jun	Jul	Aug	Sep	Oct
ſ	1	L	L	Н	Н	Н	L	L	Н	L	Н	L	Н
	2	L	L	Н	Н	Н	L	L	L	H	L	Н	Н
	3	L	L	Н	Н	L	L	Н	H	Н	L	L	Н

Baseline detection probability is fairly low (posterior mean = 0.358 with (0.261, 0.445) 95% posterior credible interval). Rain, storm, and time of sampling are identified as important predictors for observation error with posterior inclusion probabilities (PIP): 0.519, 0.651, and 0.721 respectively, but all with 95% posterior credible intervals covering 0 (Table 6). Rain, storm, and surveying in PM instead of AM have an estimated positive effect on the probability of detection. The presence of rain and storm can force parrots to fly lower down in the sky and land close to the observation point to gain shelter, increasing the probability of detection. Higher detection probability in PM than in AM is possibly due to the character of final destination: in the PM parrots are flying to one communal roost while in the AM parrots fly in multiple directions based on food dispersal and nest location, making it more difficult to detect them.

Coefficient	Mean	SD	95% PCI
Intercept	-0.586	0.209	(-1.040, -0.219)
Median Temperature	0.014	0.047	(-0.043, 0.162)
Humidity	0.003	0.033	(-0.065, 0.099)
Visibility	-0.000	0.023	(-0.053, 0.059)
Wind Speed	-0.025	0.062	(-0.224, 0.031)
Rain	0.053	0.110	(-0.036, 0.377)
Storm	0.224	0.334	(-0.031, 1.090)
Time-PM	0.128	0.140	(-0.001, 0.420)
Weather-Cloud	-0.013	0.044	(-0.152, 0.035)
Weather-Rain	0.000	0.039	(-0.092, 0.087)
Weather-Sunshine	0.000	0.039	(-0.088, 0.094)

 TABLE 6

 Ecuadorian Amazon parrots case study. Posterior summaries of coefficients for the detection probability model.

4.2. Orange-winged Amazon parrots. We next consider roost count data from Orangewinged Amazon parrots (Amazona amazonica) in Brazil. Counts were collected from a single site at an island near Belém, Pará between September 2004 and September 2005, with 96 surveys conducted (54 in the afternoon and 42 in the morning) across 50 weeks. More details can be found in De Moura, Vielliard and Da Silva (2010). We assume that the population is closed within each week, but open between weeks. Therefore, in this case, the primary periods correspond to weeks, and there are no top-level primary periods. Detection probability is modelled as a function of the following categorical covariates: Cloud (cloudy, partially cloudy, no cloud), wind (strong wind, medium wind, low wind), rain (yes, no) and time of sampling (AM or PM).

k-fold cross-validation, performed by leaving one week out at the time (k = 50), selected RW2 as the best model as seen in Table 7. We note that the Cor model is not an option in this case as the data are collected in a single year, so we cannot model correlation between weeks across different years. All models considered produced similar estimates of temporal population size, with a similar model fit. We display the results produced from the RW2 model in the main body of the paper, with the results obtained from the other models in the Supplementary material.

 TABLE 7

 Orange-winged Amazon parrots case study. Cross-validation results.

Model	DP	RW1	RW2	AR1
RMSE	2273.242	819.262	797.985	2638.177

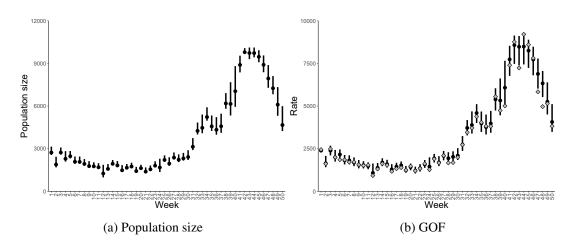


Fig 3: Orange-winged Amazon parrots case study. (a) The black dots represent the posterior mean population size each week and the thick bands represent the corresponding 95% posterior credible interval. (b) The diamonds are the observed weekly rates and the thick bands represent the 95% intervals of simulated weekly rates.

Fig 3a shows the posterior summaries of the temporal population size estimates obtained for each week using the RW2 model. The primary factor influencing the fluctuation in population size at the roosting site is the breeding season (De Moura, Vielliard and Da Silva, 2010). Consequently, the period of low population size (weeks 1-31) is possibly when paired individuals leave the roost in search of a nest, where they breed, nest, and rear young until the nestlings can fly. This long period of low population size may be due to the asynchronous reproduction of Orange-winged Amazons. The period of high population size (weeks 41-48) corresponds to the return of pairs with young, while the period of medium population size (weeks 32-40 and 49-50) corresponds to the time when individuals start returning with young (weeks 32-40) and when individuals start to disperse (weeks 49-50). Finally, like the Ecuadorian Amazon parrots, we use posterior goodness of fit to assess model fit, defining weekly rate to be the sum of counts obtained in a week divided by the number of surveys in that particular week. Fig 3b suggests that the RW2 model fits the data well as it produced similar weekly rates to the observed rates for the majority of the weeks.

Baseline detection probability is estimated as high (posterior mean = 0.875 with (0.807, 0.926) 95% posterior credible interval), possibly because in this case parrots were counted from a boat by a minimum of three teams of two observers, each team oriented in a different direction. Predictors cloud, rain and time are the only ones with PIP>0.5, but only marginally so (0.501, 0.504, 0.503, respectively), and their coefficients are estimated close to 0. In this case, rain and surveying PM instead of AM decrease the probability of detection (Table 8), and we discuss this result and compare it to that obtained for the Ecuadorian Amazon parrots in Section 5.

MODELLING ROOST COUNT DATA

Coefficient	Mean	SD	95% PCI
Intercept	1.945	0.282	(0.807, 0.926)
Partially cloudy	-0.003	0.031	(-0.067, 0.015)
Cloudy	-0.006	0.0348	(-0.109, 0.007)
Low wind	0.000	0.016	(-0.009, 0.029)
Strong wind	-0.001	0.018	(-0.020, 0.013)
Rain-Yes	-0.012	0.071	(-0.204, 0.009)
Time-PM	-0.001	0.025	(-0.034, 0.024)

 TABLE 8

 Orange-winged Amazon parrots case study. Posterior summaries of coefficients for detection probability.

5. Discussion. Roost count surveys are widely used and, for certain populations, are the only viable monitoring tool, as individuals may nest in elevated cavities in trees or cliffs that are difficult to find, reach, and capture (Dénes, Tella and Beissinger, 2018). In this paper, we have developed a new modelling framework for roost count survey data that accounts for observation error and TE, non-parametrically and parametrically to provide key estimates of population size, information on TE trends, and predictors of detection via variable selection. All of these estimates can serve as fundamental tools in adaptive wildlife monitoring, conservation, and management.

Moreover, we have performed an extensive simulation study to assess the performance of our novel modelling framework under different scenarios. When the model for detection probability is correctly specified, reliable estimates of population size and patterns of TE are obtained using both the nonparametric and parametric approaches introduced in the paper, even when the probability of detection is low. However, when the model for detection probability is misspecified, which is likely to be the case in practice, our results demonstrate the importance of using a mixed effect model for the probability of detection, so that the random effects part can absorb the lack of fit introduced by omitting important predictors for observation error. Failure to employ a mixed-effects model, in this case, gives rise to highly biased estimates of population size.

We applied our modelling framework to two case studies on parrots. We found substantially different sizes of population, detection probabilities, and variable effects on detection. The observation methods and roost site characteristics for each parrot species can explain in part these differences. Detection probability was much higher for the Orange-winged amazons, which were counted by a team of six people from a boat directly under the flight path between the mainland and an island roost, vastly reducing the chance of missing individuals. Detection however was lower for the Ecuadorian Amazon parrots, which were counted by two people from an observation tower on the mainland, where birds fly over and amongst buildings and human development to patches of scattered mangroves interspersed with aquaculture. Detection probability was higher for the Ecuadorian Amazon parrots when surveyed in the afternoon, whereas for the Orange-winged amazons, they were marginally more detectable during morning surveys. This again corresponds to the observation methods and direction of travel associated with the AM and PM surveys - with both cases showing higher detection probability when observers are at closest proximity to the roost i.e. on the observation tower when birds are departing from the mainland dry forest (Ecuadorian amazon parrots), on the boat when birds are departing from the mangrove roost (Orange-winged amazons). The effect of rain also differed, increasing the detection probability for the Ecuadorian Amazon parrots but decreasing it for the Orange-winged Amazon parrots. This can be attributed to the differing flight path birds have to make, with a 1km flight over a water body not possible in the rain (orange-winged Amazon parrots), thus lowering detection probability, whereas a shorter flight that can be taken lower down and in shorter stages by landing on

trees on the edge of the town close to the observation tower (Ecuadorian Amazon parrots) making birds more detectable.

Similarly, we identified differences in phenology between the two species, with the roost use pattern of Ecuadorian Amazon parrots being described by a two-mixture model, whereas that of Orange-winged Amazon parrots by a three-mixture model, when the DP approach is used to describe TE. This can be due to different levels of population and habitat fragmentation. There was a large difference in the population size between the two species, with the Ecuadorian Amazon parrots being just a few hundred birds, whilst the Orange-winged Amazon parrots population consists of over ten thousand birds. The Ecuadorian Amazon parrots have faced a 60 percent population decline at this roost site in the past two decades, in part attributed to habitat fragmentation, with the feeding, nesting, and roosting areas now occurring amongst a highly transformed landscape on the edges of a large city, vastly different to the relatively undisturbed roosting habitat of the Orange-winged Amazon parrots.

We have demonstrated our new modelling framework on parrot data, but bats and other species are also routinely monitored in the same way. The model can be readily fitted to such data and can be extended to account for data from multiple sites, when these are available, and to account for spatial correlation between sites. Spatial models such as the ICAR and the Besag, York and Mollié (BYM) model (Besag, York and Mollié, 1991) can be considered to account for spatial correlation.

Variable selection on detection probability via BGLSS performed well when the model is correctly specified or when misspecified and a mixed effect model is used for detection. BGLSS had lower power to identify weaker effects when using a mixed effect model for observation error. Additionally, BGLSS can only identify significant categorical covariates not significant levels of categorical variables. We also considered Bayesian Sparse Group selection (BSGS). BSGS developed by Chen et al. (2016) enables variable selection of both continuous and categorical variables. It has the advantage of identifying both significant categorical covariates and their relative levels. However, results shown in the Supplementary material suggest that BGLSS generally outperforms BSGS. Performance of other BVS methods such as the variable selection method of Griffin et al. (2020) can also be investigated in this scenario. Thus, future work can be focused on investigating/improving BVS methods when using a mixed-effect model.

The Beta DP mixture model in this framework enables our model to perform clustering of primary periods independently for top-level primary periods, and hence treats the observations as being from one long time series, with clusters, as a result, independent across top-level primary periods. An alternative would be to implement a hierarchical Dirichlet process (HDP) model (Teh et al., 2004), which allows clusters with the same locations but potentially different weights to be identified across top-level primary periods, providing a way to model dependence between top-level primary periods.

Another direction of future work is model selection. The proposed options for modelling the availability patterns define different, competing models (Table 1), for the TE pattern, each with its own advantages. We use the well-established approach of cross-validation to select between competing models. However, cross-validation can be computationally intensive as it requires fitting the model multiple times. Other model selection methods such as the Wantanable-Akaike information criterion (WAIC) (Watanabe and Opper, 2010) only require fitting the model once and can be easily computed using popular software, such as NIMBLE and STAN (Carpenter et al., 2017). Notably, WAIC computation relies on the independence assumption of data given the parameters. This assumption is often violated in temporal models where dependence among the data is a key modelling feature. Hence, future work can be focused on investigating/developing efficient model selection methods for temporally correlated data.

SUPPLEMENTARY MATERIAL

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