Estimating invasive rodent abundance using removal data and hierarchical models

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Invasive rodents pose significant ecological, economic, and public health challenges. Robust

methods are needed for estimating population abundance to guide effective management.

⁹ Traditional methods such as capture-recapture are often impractical for invasive species due to

ethical and logistical constraints. Here, I showcase the application of hierarchical multinomial

11 N-mixture models for estimating the abundance of invasive rodents using removal data. First, I

performed a simulation study which demonstrated minimal bias in abundance estimates across

a range of sampling scenarios. Second, I analyzed removal data for two invasive rodent species:

coypus (Myocastor coypus) in France and muskrats (Ondatra zibethicus) in the Netherlands. Using

hierarchical multinomial N-mixture models, I examined the effects of temperature on abundance

6 while accounting for imperfect and time-varying capture probabilities. I also showed how to

accommodate spatial variability using random effects, and quantified uncertainty in parameter

estimates. Overall, I hope to demonstrate the flexibility and utility of hierarchical models in

invasive species management. I provide reproducible code and data to encourage broader

20 adoption of multinomial N-mixture models.

21 Keywords: Invasive species, Multinomial N-mixture, Population size, Statistical ecology

2 Introduction

Invasive species are a significant global issue, with wide-ranging impacts on ecosystems, economies, and public health (Pyšek et al. 2020, Roy et al. 2024). Among these, the financial, epidemiological, social, and ecological costs associated with invasive rodents are substantial, as 25 they damage infrastructures, degrade agricultural systems, and act as reservoirs for zoonotic diseases (Diagne et al. 2023). Effective management of invasive species requires the estimation of population abundance for guiding control efforts and evaluating the success of eradication or regulation programs (Williams et al. 2002, Thompson et al. 2021). However, the challenge in estimating animal 30 abundance is that individuals are not always observed even when present due to imperfect 31 detection (Borchers et al. 2002, Seber and Schofield 2023). Ignoring imperfect detection leads to biased estimates of population abundance (Kéry and Schmidt 2008). To account for imperfect detection, capture-recapture methods are usually used to correct observed counts (McCrea and Morgan 2015). Yet, for invasive species, capture-recapture is often impractical, as ethical and management concerns typically prevent the release of captured animals. 36 An alternative approach involves the use of removal methods (Rodriguez de Rivera and McCrea 37 2021) in which individuals are captured and permanently removed from the study area during successive sampling occasions. This process leads to a decrease in the expected number of captures by a consistent proportion over time (rather than by a fixed amount decline), which informs on the total abundance as the initial population determines how quickly the number of individuals available for capture diminishes. While standard removal methods are well-established (Moran 1951, Zippin 1956, 1958, Ro-43 driguez de Rivera and McCrea 2021), recent advances in population ecology remain underutilized in the context of invasive species. Hierarchical models, in particular, have gained traction (Royle and Dorazio 2008, Kéry and Royle 2015) due to their ability to: (i) explicitly separate

biological processes of interest (e.g., population dynamics) from observation processes (e.g., imperfect detection), thus enabling more accurate modeling; (ii) incorporate environmental, spatial, or temporal covariates at multiple levels, allowing exploration of how various factors influence ecological processes; and (iii) share information across groups by modeling parameters 50 hierarchically with random effects, which improves estimates for groups with fewer data. 51 In this paper, I showcase the application of a hierarchical formulation of removal models, the multinomial N-mixture model (Dorazio et al. 2005), to estimate the abundance of rodents in Europe. In this study, I focus on the coypu (Myocastor coypus) in France and the muskrat (Ondatra 54 zibethicus) in the Netherlands. Both species are semi-aquatic rodents introduced to Europe in 55 the early 20th century following escapes or releases from fur farms. The coypu, native to South America, has formed widespread invasive populations in France (Bonnet et al. 2023), where 57 it causes significant damage to infrastructure and crops. Additionally, it serves as a healthy carrier of leptospirosis, a zoonotic disease with potentially serious consequences. Similarly, the muskrat, native to North America, has established extensive populations in the Netherlands. 60 By burrowing into riverbanks, dykes, and dams, muskrats compromise the integrity of these 61 structures, posing a threat to public safety (Loon et al. 2017). Using removal data, I demonstrate the application of the multinomial N-mixture model to 63 estimate the abundance of rodent populations. First, I conduct a simulation study to evaluate the model's performance under varying numbers of sampling sites and sampling occasions. Second, I present a case study on a coypu population in France to illustrate the hierarchical structure 66 of the multinomial N-mixture model, demonstrating how covariates can be incorporated to 67 account for variations in abundance and capture probabilities. Third, I use a case study on muskrats in the Netherlands to demonstrate the integration of random effects within the model. 69 To facilitate reproducibility, I provide the accompanying code and data, aiming to promote the broader adoption of removal models in the study of biological invasions.

Methods

3 Multinomial N-mixture model

Think of a dice with six sides. The dice has a 1 in 6 chance of landing on face 1, the same for face 2, and so on. If I roll the dice 30 times, I expect to get face 1 five times, face 2 five times, and so on, on average. In this experiment, y_1 , the vector made of the number of 1s, y_2 , the number of 2s, ..., and y_6 , the number of 6s, follows a multinomial distribution with parameters the 77 number of rolls (30) and probabilities (1/6, 1/6, ..., 1/6). Now think of a removal campaign conducted over 3 months. We record the number of rodents y_1 captured in month 1, y_2 in month 2, y_3 in month 3, and let y_4 represent the number of rodents never captured. Let p be the probability of capturing a rodent in a given month. The probability of capturing a rodent in the first month is $\pi_1 = p$. The probability of capturing 82 a rodent in the second month is $\pi_2 = (1-p)p$ the probability of not capturing it in the first month (1-p) multiplied by the probability of capturing it in the second month p. The probability of capturing a rodent in the third month is $\pi_3 = (1 - p)(1 - p)p$, the probability of not capturing it in the first and second months, (1-p)(1-p), multiplied by the probability of capturing it in the third month, p. Finally, the probability of never being captured is π_4 87 1 - (p + (1-p)p + (1-p)(1-p)p) the complement of the probability of being captured in the 88 first, second, or third month. If we assume that N represents the abundance, then we have that the vector of counts (y_1, y_2, y_3, y_4) follows a multinomial distribution with parameters N and probabilities $(\pi_1, \pi_2, \pi_3, \pi_4)$. In general, we assume that N follows a Poisson distribution with parameter the expected number of rodents denoted λ . And there you have it, the multinomial N-mixture model for a removal experiment, which is similar to throwing a dice N times and 93 the $\pi's$ give the probabilities that I get a given face of that dice. In general, we monitor rodents 94 in several populations or sites i = 1, ..., S and we need to estimate local abundance N_i . To do so, Dorazio et al. (2005) extended multinomial N-mixture models to account for spatial

- variation in abundance and/or capture, and showed that abundance estimates had similar or
- better precision than those obtained from analyzing removal data for each population or site
- 99 separately.
- Parameters N, p, and λ are unknown and need to be estimated. In a frequentist framework,
- marginalization is performed by summing over all possible values of N (Dorazio et al. 2005). In
- a Bayesian framework, all these parameters are estimated directly, which simplifies the process
- (Royle and Dorazio 2006). Both parameters, λ and p, can be modeled as functions of explanatory
- spatial and temporal variables, in the spirit of generalized linear models, and Poisson or logistic
- regressions for example. I warmly recommend chapter 7 in Kéry and Royle (2015) for a detailed
- description of the multinomial mixture model.

of Simulation study

I conducted a simulation study to evaluate the model's performance by examining parameter bias under varying numbers of sampling sites and sampling occasions. I simulated removal data over 1, 5, 10 and 50 sites using a Poisson distribution with expected number of animals λ between 10 and 100 (20 values) for the ecological process. I simulated the observation process with a capture probability p varying between 0.3 and 0.9 (20 values) across 3, 5 and 10 occasions per site. In total, I considered 4800 scenarios. I fitted the multinomial N-mixture model to the simulated data within the frequentist framework using function multinomPois() in the R package unmarked (Kellner et al. 2023), and I repeated this procedure 500 times. Eventually, I calculated the relative bias for each parameter.

117 Case studies

In this section, I analyzed removal data from two rodent species: coypus in France and muskrats in the Netherlands. With these case studies, I aimed at illustrating specific features of hierarchical multinomial N-mixture models. For both species, I explored the potential effect of temperature on abundance (e.g., Gosling 1981, Simpson and Boutin 1993). A comprehensive analysis of the ecological factors influencing population dynamics was beyond the scope of this work and will be addressed in future studies.

124 Coypus in France

Removal data on coypus were collected from annual control operations conducted since 2015 in 125 several cities within the Hérault department, located in the Occitanie region of southern France. 126 These operations are carried out year-round, with the exception of July and August. Coypus are 127 trapped using cages by a network of volunteers coordinated by the Syndicat Mixte du Bassin de 128 l'Or and the Fédération Départementale des Chasseurs de l'Hérault. For this study, I focus on 129 data from 2022, specifically from sampling occasions in February, March, and April. The data, covering S = 6 cities, are summarized in Table 1. I fitted a model where the expected number of 1.31 coypus was modeled as a function of temperature, while the capture probability was allowed 132 to vary by month.

	Removed in	Removed in	Removed in	Averaged
City	February	March	April	temperature
Candillargues	18	12	38	9.5
Lansargues	15	17	75	8.8
Mauguio	20	9	6	9.2
Saint-Nazaire-de-Pézan	169	41	15	9.3
Saint-Just	85	61	77	9.2
Valergues	0	1	3	9.4

Table 1: Number of invasive coypus removed monthly and the average 3-month temperature across several cities in the Hérault department, France, in 2022.

Muskrats in the Netherlands

Removal data on muskrats in the Netherlands were collected by professional trappers. The data were registered in atlas blocks (5 x 5 km) per periods of four weeks. For this study, I focus on data from 2014, specifically from sampling occasions in January, February, and March. The data were made available through the LIFE MICA project (Cartuyvels et al. 2024) and can be freely downloaded from https://www.gbif.org/dataset/7d75109d-a6cb-4e90-89d0-79d08577c580. The data, covering S = 342 cities, are presented in Figure 1. I fitted the same model as for the coypus data, except that I added a site random effect on abundance to accommodate the spatial variation that was not explained by temperature.

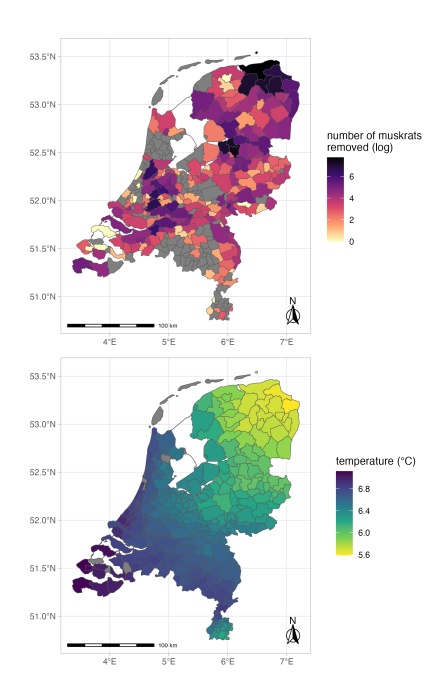


Figure 1: Total number of invasive muskrats removed in January, February and March (top panel), and the average 3-month temperature (bottom panel) across the Netherlands in 2014.

143 Implementation

For all analyses, I used the statistical language R (R Core Team 2024). I used the tidyverse (Wickham et al. 2019) suite of packages for data manipulation and visualization, sf (Pebesma and Bivand 2023) for dealing with spatial data and krigR (Kusch and Davy 2022) to get temperature data. For the two case studies, I fitted models within a Bayesian framework using Markov chain Monte Carlo (MCMC) algorithms. I used both the NIMBLE (de Valpine et al. 2017) and the ubms (Kellner et al. 2021) packages. The former offers high flexibility, enabling users to define custom likelihoods, though it requires manual coding, while the latter features simpler 150 syntax with pre-built multinomial N-mixture models, albeit limited to a Poisson distribution for abundance. I specified weakly informative priors for all parameters, specifically normal 152 distributions with mean 0 and standard deviation 1.5 for regression parameters, and a uniform 153 distribution for the standard deviation of the random effects. I ran two chains for a total of 200,000 iterations with a burn-in of 20,000 iterations. I summarized posterior distributions with 155 posterior mean and 95% credible intervals. I assessed model convergence using R-hat values (< 156 1.1), effective sample size (> 100), and visual inspection of the trace plots.

158 Results and discussion

The results of the simulation are presented in Figure 2. Overall, the analysis revealed minimal bias, with the exception of one site that showed a notable deviation.

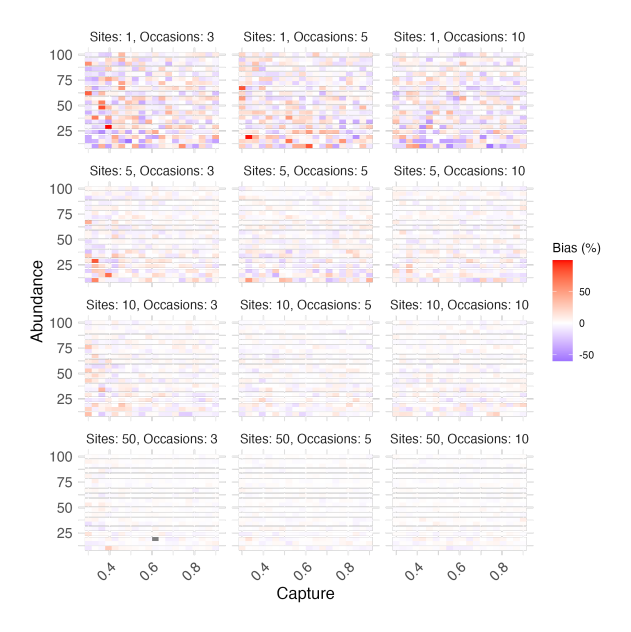


Figure 2: Relative bias in abundance estimates from a multinomial N-mixture model with constant parameters. Capture probabilities (X-axis) range from 0.3 to 0.9, while abundance (Y-axis) varies between 10 and 100 individuals. Scenarios consider 3, 5, and 10 capture occasions (columns) and 1, 5, 10, and 50 sites (rows). Results are based on 500 simulations.

Increasing the number of sites to 10 significantly reduced this bias, and no bias was observed with 50 sites, supporting the recommendation by (Dorazio et al. 2005) to analyze data jointly rather than separately. These findings align with previous simulation studies, which also reported minimal bias in removal models (e.g., Womack-Bulliner et al. 2019). To enhance

reproducibility, I provide the code for the simulation study in the Supplementary Material. This
resource can be adapted for various purposes, such as conducting custom simulation studies or
designing removal protocols and conducting power analyses.

In the coypus case study, temperature was found to have a negative effect on abundance, with a slope estimate of -0.14 (-0.22, -0.07). Capture probabilities were estimated at 0.43 (0.29, 0.49) in February, 0.35 (0.19, 0.44) in March and 0.84 (0.35, 1.00) in April. The posterior distributions of abundance across the different sites are presented in Figure 3.

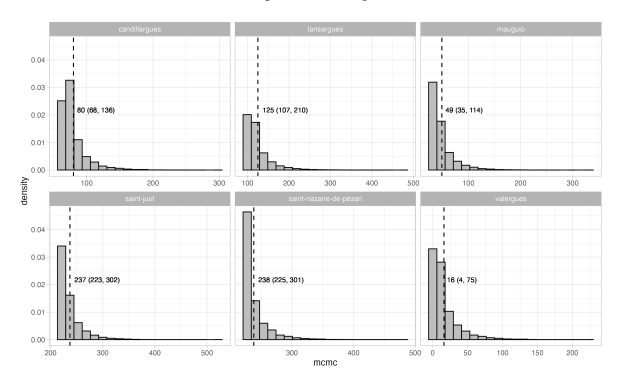


Figure 3: Posterior density plots for coypu abundance across several cities in the Hérault department, France, in 2022. The vertical shaded line indicates the posterior mean abundance, accompanied by its credible interval. See Table 1 for the raw data.

A key assumption of the multinomial N-mixture model is that abundance follows a Poisson distribution, which assumes the mean and variance are equal. However, this assumption does not appear valid based on the estimated abundance. Fortunately, this limitation can be addressed by relaxing the Poisson assumption. A straightforward approach is to use a negative

- binomial distribution to account for overdispersion. This adjustment can be implemented in
 both NIMBLE and unmarked but is not currently supported by ubms. I demonstrate how to fit such
 a model to the coypus data in the Supplementary Material. Interestingly, under the negative
 binomial model, the effect of temperature on abundance was no longer significant, with a slope
 estimate of -0.27 (-1.29, 0.60).
- In the muskrats case study, temperature was found to have a negative effect on abundance, with a slope estimate of -0.48 (-0.70, -0.26). The standard deviation of the site random effect was estimated at 1.62 (1.46, 1.79). Capture probabilities were estimated at 0.12 (0.05, 0.25) in January, 0.25 (0.15, 0.35) in February and 0.60 (0.53, 0.75) in March. Estimated abundance across sites is presented in Figure 4.

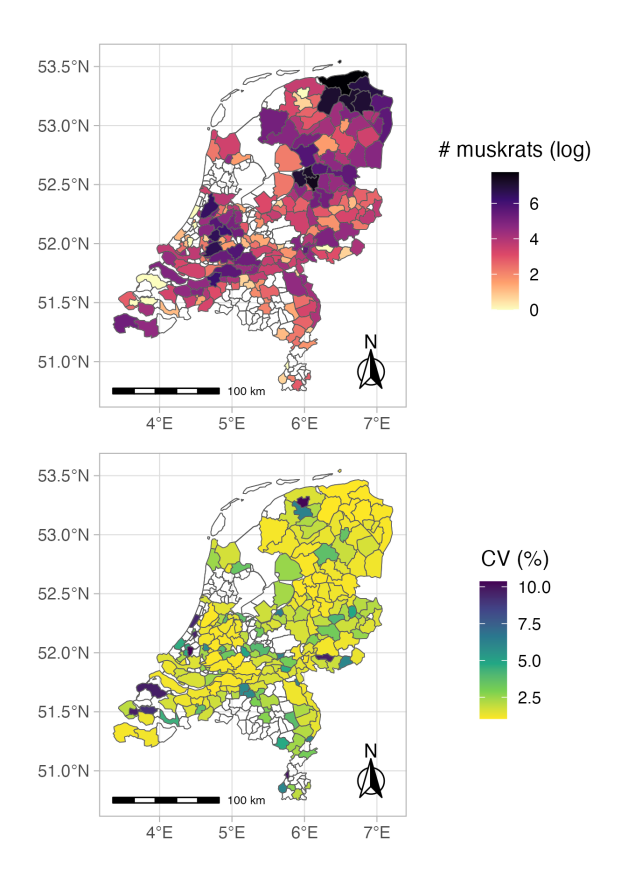


Figure 4: Posterior mean estimates of muskrat abundance in the Netherlands in 2022 (top panel) and the corresponding coefficient of variation (bottom panel). See Figure 1 for the raw data.

An important feature of multinomial N-mixture models is their ability to quantify uncertainty, which is often overlooked in spatial analyses. Here, I provide the coefficient of variation 187 to represent the uncertainty surrounding abundance estimates. This metric can help identify 188 specific areas where increased sampling effort might be beneficial to improve estimate precision. 189 Another key assumption underlying the proper use of multinomial N-mixture models is that of 190 population closure, which assumes no births, deaths, immigration, or emigration occur during 191 the trapping period. A straightforward approach to relax this assumption is to fit multiple years of data (i.e., stacking the data) into a standard multinomial N-mixture model. In this 193 approach, year-site combinations are treated as separate sites, and year is included as a site 194 covariate or random effect in the model. In the muskrats example, this method can be used to evaluate a temporal effect on the relationship between temperature and abundance. Assuming 196 an increase in temperature over time, one might hypothesize a decoupling or weakening of 197 the relationship between abundance and temperature. To test this hypothesis, I conducted an 198 additional analysis of the muskrat dataset spanning the 1987–2014 period, modeling the slope 199 of the temperature-abundance relationship as a linear function of time. The results revealed a 200 negative trend in the slope, estimated at -0.040 (-0.042, -0.039), providing evidence to support 201 the hypothesis of a temporal weakening of this relationship. 202 Several perspectives arise from this work. From a methodological standpoint, this study 203 highlights the suitability of hierarchical models for capturing dependencies in space and time, 204 which are common in ecological systems and removal experiments in particular. Two areas 205 stand out as particularly worth exploring. First, I focused in this paper exclusively on static 206 models under the assumption of population closure (see however previous section). When the 207 mechanisms underlying population dynamics—such as survival, recruitment, or dispersal—are 208 of interest, multinomial N-mixture models can be extended to open populations (Matechou et 209

al. 2016, Link et al. 2018, Zhou et al. 2019). These extended models can be implemented using

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unmarked or NIMBLE, though they are not yet available in ubms.

A second area of investigation concerns the spatial dimension of multinomial N-mixture models, 212 particularly the assumption of independence among removal sites (i.e., that removals at one 213 site do not influence those at another). One possible solution is to include site random effects, as demonstrated in the muskrat case study. To better address spatial autocorrelation, restricted 215 spatial regression (RSR) can also be employed (Johnson et al. 2013, Broms et al. 2014) tp 216 impose a structure where spatially adjacent sites are modeled to have correlated random effects, 217 effectively accounting for spatial autocorrelation. RSR models are advantageous because their 218 random effects are constructed to be uncorrelated with fixed covariates, avoiding potential 219 confounding issues, and they are computationally efficient. These models are easy to fit using ubms and can also be implemented in NIMBLE (Cook et al. 2022), although they are 221 not currently supported by unmarked. A promising extension would involve adapting the covariance structure in these models to account for stream networks, which is particularly relevant for semi-aquatic rodents (Gimenez 2024, Lu et al. 2024). 224 From an ecological perspective, the main contribution of this paper is to demonstrate the 225 estimation of abundance for rodent populations in Europe. The European Union Regulation No. 1143/2014 was established to mitigate the negative impacts of invasive alien species on 227 biodiversity. This regulation mandates measures to prevent the introduction of invasive alien 228 species and manage their established populations. In this study, I focused on two species listed as species of Union Concern under the regulation, which requires member states to implement 230 appropriate management actions. 231 In this context, I argue that estimates of abundance alone are of limited utility for supporting evidence-based management of invasive rodents. While it is straightforward to calculate 233 the number of coypus or muskrats remaining after removal campaigns (including associated 234

uncertainty), these estimates achieve their full potential only when integrated into a decision-

making framework. Such a framework enables the evaluation of the efficiency of eradication or control operations, facilitates the optimal allocation of limited resources, and incorporates uncertainty into the management of invasive species (Shea et al. 2002, Yackel Adams et al. 2024).

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244 Data availability statement

Data and code are available at https://github.com/oliviergimenez/counting-rodents.

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