

Spatial occupancy models for data collected on stream networks

Olivier Gimenez^{1*}

2024-07-25

¹ CEFE, Univ Montpellier, CNRS, EPHE, IRD, Montpellier, France

* Corresponding author: olivier.gimenez@cefe.cnrs.fr

To monitor streams and rivers biodiversity, we need to quantify species distribution. To do so, occupancy models allow distinguishing the non-detection of a species from its actual absence. Occupancy models can account for spatial autocorrelation, but are not suited for streams and rivers because of their spatial structure in networks. Here I propose spatial occupancy models for data collected on stream and river networks. I present the statistical developments of the model, then I illustrate the approach on a semi-aquatic mammal. Overall, spatial stream network occupancy models provide a formal approach to assess biodiversity in streams and rivers.

Keywords: Bayesian statistics, Spatial stream network models, Occupancy models, Spatial autocorrelation, Wildlife monitoring

18 Introduction

19 Streams and rivers provide essential habitats for numerous species of animals and plants ([Reid](#)
20 [et al. 2019](#)). The ecological health of these freshwater ecosystems is paramount not only for the
21 biodiversity they harbor but also for the ecosystem services they provide, which are
22 indispensable to both wildlife and human populations ([Vári et al. 2021](#)). However, human
23 activities are altering the natural conditions of streams, rivers, and their associated riparian
24 habitats, jeopardizing the persistence of these ecosystems ([Albert et al. 2020](#)).

25 In this context, species distribution models (SDMs) are essential tools in understanding and
26 preserving biodiversity ([Elith and Leathwick 2009](#)). SDMs predict the distribution of species,
27 helping scientists and conservationists in identifying critical habitats and biodiversity hotspots.
28 Additionally, SDMs inform strategies aimed at mitigating the impacts of climate and land-use
29 changes, managing invasive species, and enhancing habitat connectivity in freshwater
30 ecosystems ([Domisch et al. 2015](#)).

31 Species Distribution Models (SDMs) are influenced by two main issues: imperfect detection and
32 spatial autocorrelation ([Guélat and Kéry 2018](#)). First, imperfect detection occurs when a species
33 present in a given area may not be detected during surveys due to factors such as observer
34 experience, species behavior, and environmental conditions. If imperfect detection is ignored, it
35 can lead to biased estimates of species distribution and flawed inferences about the relationship
36 between species presence and environmental factors (e.g., [Lahoz-Monfort et al. 2014](#)). This can
37 misinform conservation strategies and habitat management decisions. To address this issue,
38 occupancy models are SDMs that rely on repeated visits of spatial sampling units for inferring
39 distribution ([MacKenzie et al. 2017](#)). These models have been widely used in freshwater
40 ecosystems for various taxa ([Preece et al. 2021](#), [Charbonnel et al. 2022](#), [Wedderburn et al. 2022](#)).

41 Second, SDMs rely on the assumption of independent residuals. This assumption may be
42 violated if nearby sampling sites tend to have similar probabilities of species presence, leading

to biased estimates of species distribution and potentially inflating the effects of environmental factors (Dormann et al. 2007). Several extensions of occupancy models have been proposed to account for spatial autocorrelation, building on the spatial statistics literature, e.g. conditional autoregressive models (Johnson et al. 2013) and geoaddivitive models (Rushing et al. 2019). However, these models rely on the Euclidean distance between the spatial sampling units, which does not acknowledge the spatial structure in networks of streams and rivers.

Here I propose spatial occupancy models that account for spatial autocorrelation based on stream flow and connectivity (Peterson et al. 2013). I build on the linear mixed modelling approach developed by Ver Hoef and Peterson (2010) and Peterson and Hoef (2010), which integrates various distance-based spatial correlation structures (both Euclidean and non-Euclidean) within a single model. I plug-in this variance component approach into occupancy models using a Bayesian approach. A similar approach was recently undertaken by Lu et al. (2024) for count data to estimate abundance. Below I outline the statistical developments of this model and demonstrate its application to a semi-aquatic mammal in French streams and rivers.

Methods

Occupancy models

To address imperfect detection, I use occupancy models to estimate the true species distribution (MacKenzie et al. 2017). In these models, monitoring occurs across S spatial sampling units, or sites. If detection was perfect, the state z_i of a site i would be a Bernoulli random variable, taking value 1 with occupancy probability ψ_i if the site was occupied, and 0 otherwise with probability $1 - \psi$. However, because the ecological process of state occupancy z_i is only partially observable (since the species might be present but undetected), we must also account

for the observation process, which is also modeled as a Bernoulli random variable. When the species is detected at site i , i.e. $y_i = 1$, with detection probability p , it confirms that the site is occupied. Conversely, if the species is not detected, i.e. $y_i = 0$, with probability $1 - p$, we cannot determine whether the site is occupied or not. Both parameters, ψ and p , can be modeled as functions of explanatory spatial variables, in the spirit of generalized linear models and logistic regressions for example. To estimate occupancy and detection probabilities separately, data should be collected from at least two independent visits to each site within a short period, ensuring that sites remain in the same state.

Spatial autocorrelation for stream networks

How is spatial autocorrelation accounted for in occupancy models? The usual way is to write the probabilities of occupancy $\psi = (\psi_1, \dots, \psi_S)$ on some scale, say the logit scale, as a function of explanatory variables gathered in a matrix \mathbf{X} with corresponding regression parameters β that need to be estimated. To account for spatial autocorrelation, a random effect ϵ is added to the model, which captures the spatial dependencies among sites (Guélat and Kéry 2018):

$$\text{logit}(\psi) = \mathbf{X}\beta + \epsilon.$$

The random effect ϵ can be structured using methods such as conditional autoregressive models and their extensions (Johnson et al. 2013), or ge additive models (Rushing et al. 2019). However, these approaches typically rely on Euclidean distance to assess proximity among sites, which may not fully capture the complex spatial dependencies present in streams and rivers. Specifically, we are interested in flow connectivity and the topology of streams and rivers (Peterson et al. 2013). Following Ver Hoef and Peterson (2010) and Peterson and Hoef (2010), I define two sites as flow-connected if water flows from an upstream site to a downstream site, and as flow-unconnected if they share a common confluence downstream but

do not directly share flow. Then I parameterize occupancy by rewriting the random effect as a mixture of four components as follows:

$$\text{logit}(\psi) = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\tau}_{tu} + \boldsymbol{\tau}_{td} + \boldsymbol{\tau}_{eu} + \boldsymbol{\epsilon}$$

where $\boldsymbol{\tau}_{tu}$ is a random effect with spatial covariance between flow-connected sites that can occur in the same direction of the river flow (tail-up), which is suitable for organisms that move passively, such as mussels), $\boldsymbol{\tau}_{td}$ is a random effect with spatial covariance between flow-connected and flow-unconnected sites, which can occur with or against the direction of flow (tail-down), applicable to organisms that move actively, such as semi-aquatic mammals, $\boldsymbol{\tau}_{eu}$ is a random effect with a spatial covariance independent of the network topology, influenced by factors like air temperature or precipitation, and $\boldsymbol{\epsilon}$ is a random effect with variance, often referred to as the nugget, which accounts for additional variability. How to build these covariance components is described in details elsewhere (e.g., [Ver Hoef et al. 2019](#)), and I provide an example in the next section.

Case study

To illustrate the new approach, I investigated the impact of human disturbance on the occupancy of European otter (*Lutra lutra*) in France. The otter, a semi-aquatic mammal, faced near extinction in the 20th century in France due to extensive hunting for its fur. With hunting bans and protection efforts, the species is now recolonizing the country, and the ecological question is assessing its current distribution. Data on otter detection and non-detection were collected in 2003-2005 in the Midi-Pyrénées region (see panel a in Fig. 1). Observers searched for signs of otter presence at a small river catchment scale, which was used as the spatial sampling unit. These data were analyzed by Couturier et al. (2023), who found that human density and the proportion of cultivated area influenced occupancy. In this study, I focus on a

subsample of this dataset, covering $S = 56$ sites in the Lot, Aveyron and Cantal counties, which were visited 3 times (see panel b in Fig. 1). I used human population density as a proxy for human disturbance, calculated as the number of inhabitants per km² within a 200-m buffer around each stream (see panel c in Fig. 1). Additionally, I considered the proportion of cultivated areas as an explanatory variable. Detection was considered as constant. Regarding spatial autocorrelation, since otters can move both downstream and upstream, I used a tail-down model

$$\text{logit}(\psi) = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\tau}_{td}$$

with a exponential covariance:

$$\text{Cov}(\tau_i, \tau_j) = \begin{cases} \sigma^2 \exp(-h/\theta), & \text{if sites } i \text{ and } j \text{ are flow-connected} \\ \sigma^2 \exp(-(a+b)/\theta), & \text{if sites } i \text{ and } j \text{ are flow-unconnected} \end{cases}$$

where I dropped the td notation in τ_{td} for clarity, h is the stream distance between sites i and j , b denotes the longer of the distances to the common downstream junction, a denotes the shorter of the two distances, σ^2 is a variance parameter usually referred to as the partial sill and θ is a range parameter (see Appendix A in [Peterson and Hoef 2010](#)).

Implementation

For all analyses, I used the statistical language R ([R Core Team 2023](#)). I used the openSTARS ([Kattwinkel et al. 2020](#)) and SSN ([Hoef et al. 2014](#)) packages to build and characterize the network and calculate hydrological distances. I fitted models within the Bayesian framework, specifying weakly informative priors ([Northrup and Gerber 2018](#)), implementing a marginalized likelihood ([Clark and Altwegg 2019](#)), and using the rstan ([Stan Development Team 2023](#)) package. I ran two chains for a total of 15,000 iterations with a burn-in of 5,000

iterations. For simplicity, I summarized posterior distributions with posterior mean and 95% credible intervals. I assessed model convergence using R-hat values (< 1.05), effective sample size (> 100), and visual inspection of the trace plots. I provide additional information in the code available at <https://github.com/oliviergimenez/spatial-stream-network-occupancy-model>.

Results and discussion

Here, I provide the parameter estimates from the new model accommodating spatial autocorrelation, unless otherwise specified. Detection probability was less than one, estimated at 0.71 (0.59, 0.80). The proportion of cultivated area had no effect, with a slope estimated at 0.60 (-0.67, 1.96). Population density also had no effect on occupancy probability, with a slope estimated at -0.96 (-2.24, 0.17). However, I did find a negative effect when spatial autocorrelation was ignored, with a slope estimated at -1.10 (-1.99, -0.34). This latter result aligns with a previous analysis of a more extensive dataset (Couturier et al. 2023) that also ignored spatial autocorrelation, in areas highly connected to otter-occupied catchments.

As anticipated, the effect size of human density increased when spatial autocorrelation was ignored. The most likely explanation for this is a bias due to an omitted variable. There is spatial autocorrelation in human density (see panel c in Fig. 1), which inflates its effect size; this bias is controlled for when spatial autocorrelation is included in the model. There may be spatial variation in occupancy probabilities attributable to another variable that needs to be accounted for.

Two short-term perspectives arise from this work. From a methodological perspective, the new approach could be extended to multi-season occupancy models, enabling the modeling of colonization probability as a function of distance to habitat features that may impede species movement. This would facilitate the quantification of landscape connectivity in freshwater

ecosystems (Kervellec et al. 2023). Such development requires moving to spatio-temporal models for stream and river data, which have recently become available (Santos-Fernandez et al. 2022). From an ecological perspective, the new approach presents significant potential for the analysis of environmental DNA (eDNA). The eDNA methodology offers substantial promise for the non-invasive monitoring of biodiversity in freshwater ecosystems (Carraro et al. 2020). While spatial stream network models have been employed to analyze eDNA data (Winkowski et al. 2024), these models have overlooked the issue of imperfect detection. Previous studies have recognized occupancy models as effective tools for eDNA data analysis (Burian et al. 2021), with some considering spatial autocorrelation (Chen and Ficetola 2019), however they have yet to integrate spatial stream networks. The new approach addresses this gap by incorporating both imperfect detection and spatial stream networks.

Acknowledgments

I would like to warmly thank Jay Ver Hoef and Edgar Santos Fernández for useful discussions on spatial stream network models. I also thank Maëlis Kervellec for sharing her code to wrangle the otter data.

Ethics and Integrity statements

Data availability statement

Data and code are available at

<https://github.com/oliviergimenez/spatial-stream-network-occupancy-model>.

Funding statement

This research is a product of the DISCAR group funded by the French Foundation for Research on Biodiversity (FRB) through its synthesis center CESAB.

Conflict of interest disclosure

The author has no conflicts of interest to declare.

References

- Albert, J. S., G. Destouni, S. M. Duke-Sylvester, A. E. Magurran, T. Oberdorff, R. E. Reis, K. O. Winemiller, and W. J. Ripple. 2020. Scientists' warning to humanity on the freshwater biodiversity crisis. *Ambio* 50:85–94.
- Burian, A., Q. Mauvisseau, M. Bulling, S. Domisch, S. Qian, and M. Sweet. 2021. Improving the reliability of eDNA data interpretation. *Molecular Ecology Resources* 21:1422–1433.
- Carraro, L., E. Mächler, R. Wüthrich, and F. Altermatt. 2020. Environmental DNA allows upscaling spatial patterns of biodiversity in freshwater ecosystems. *Nature Communications* 11:3585.
- Charbonnel, A., F. Blanc, P. Laffaille, M. Némot, and L. Buisson. 2022. Combining spatial dependence occupancy models and conservation gap analyses to promote species conservation: A case study with a threatened semi-aquatic mammal. *Biological Conservation* 270:109567.
- Chen, W., and G. F. Ficetola. 2019. Conditionally autoregressive models improve occupancy analyses of autocorrelated data: An example with environmental DNA. *Molecular Ecology Resources* 19:163–175.
- Clark, A. E., and R. Altwegg. 2019. Efficient Bayesian analysis of occupancy models with logit

link functions. *Ecology and Evolution* 9:756–768.

Couturier, T., J. Steinmetz, P. Defos du Rau, D. Marc, E. Trichet, R. Gomes, and A. Besnard. 2023.

Intensive agriculture as the main limiting factor of the otter's return in southwest france.

Biological Conservation 279:109927.

Domisch, S., S. Jähnig, J. Simaika, M. Kuemmerlen, and S. Stoll. 2015. Application of species

distribution models in stream ecosystems: The challenges of spatial and temporal scale,

environmental predictors and species occurrence data. *Fundamental and Applied*

Limnology 186:45–61.

Dormann, F. C., J. M. McPherson, M. B. Araújo, R. Bivand, J. Bolliger, G. Carl, R. G. Davies, A.

Hirzel, W. Jetz, W. Daniel Kissling, I. Kühn, R. Ohlemüller, P. R. Peres-Neto, B. Reineking, B.

Schröder, F. M. Schurr, and R. Wilson. 2007. Methods to account for spatial autocorrelation

in the analysis of species distributional data: A review. *Ecography* 30:609–628.

Elith, J., and J. R. Leathwick. 2009. Species distribution models: Ecological explanation and

prediction across space and time. *Annual Review of Ecology, Evolution, and Systematics*

40:677–697.

Guélat, J., and M. Kéry. 2018. Effects of spatial autocorrelation and imperfect detection on

species distribution models. *Methods in Ecology and Evolution* 9:1614–1625.

Hoef, J. M. V., E. E. Peterson, D. Clifford, and R. Shah. 2014. SSN: An R package for spatial

statistical modeling on stream networks. *Journal of Statistical Software* 56:1–45.

Johnson, D. S., P. B. Conn, M. B. Hooten, J. C. Ray, and B. A. Pond. 2013. Spatial occupancy

models for large data sets. *Ecology* 94:801–808.

Kattwinkel, M., E. Szöcs, E. Peterson, and R. Schäfer. 2020. Preparing GIS data for analysis of

stream monitoring data: The R package openSTARS. *Plos One* 15:e0239237.

Kervellec, M., T. Couturier, S. Bauduin, D. Chenesseau, P. D. du Rau, N. Drouet-Hoguet, C.

Duchamp, J. Steinmetz, J.-M. Vandiel, and O. Gimenez. 2023. Bringing circuit theory into

spatial occupancy models to assess landscape connectivity. *bioRxiv*.

220 Lahoz-Monfort, J. J., G. Guillera-Arroita, and B. A. Wintle. 2014. Imperfect detection impacts
 221 the performance of species distribution models. *Global Ecology and Biogeography*
 222 23:504–515.

223 Lu, X., Y. Kanno, G. P. Valentine, J. M. Rash, and M. B. Hooten. 2024. Using multi-scale spatial
 224 models of dendritic ecosystems to infer abundance of a stream salmonid. *Journal of*
 225 *Applied Ecology* 61:1703–1715.

226 MacKenzie, D. I., J. D. Nichols, J. A. Royle, K. H. Pollock, L. L. Bailey, and J. E. Hines. 2017.
 227 *Occupancy Estimation and Modeling: Inferring Patterns and Dynamics of Species*
 228 *Occurrence*. Elsevier.

229 Northrup, J., and B. Gerber. 2018. A comment on priors for Bayesian occupancy models. *Plos*
 230 *One* 13:e0192819.

231 Peterson, E. E., and J. M. V. Hoef. 2010. A mixed-model moving-average approach to
 232 geostatistical modeling in stream networks. *Ecology* 91:644–651.

233 Peterson, E. E., J. M. Ver Hoef, D. J. Isaak, J. A. Falke, M.-J. Fortin, C. E. Jordan, K. McNyset, P.
 234 Monestiez, A. S. Ruesch, A. Sengupta, N. Som, E. A. Steel, D. M. Theobald, C. E. Torgersen,
 235 and S. J. Wenger. 2013. Modelling dendritic ecological networks in space: An integrated
 236 network perspective. *Ecology Letters* 16:707–719.

237 Preece, E. P., M. Bryan, S. M. Mapes, C. Wademan, and R. Dorazio. 2021. Monitoring for
 238 freshwater mussel presence in rivers using environmental DNA. *Environmental DNA*
 239 3:591–604.

240 R Core Team. 2023. [R: A language and environment for statistical computing](#). R Foundation for
 241 Statistical Computing, Vienna, Austria.

242 Reid, A. J., A. K. Carlson, I. F. Creed, E. J. Eliason, P. A. Gell, P. T. J. Johnson, K. A. Kidd, T. J.
 243 MacCormack, J. D. Olden, S. J. Ormerod, J. P. Smol, W. W. Taylor, K. Tockner, J. C. Vermaire,
 244 D. Dudgeon, and S. J. Cooke. 2019. Emerging threats and persistent conservation challenges
 245 for freshwater biodiversity. *Biological Reviews* 94:849–873.

Rushing, C., J. Andrew Royle, D. Ziolkowski Jr, and K. Pardieck. 2019. Modeling spatially and temporally complex range dynamics when detection is imperfect. *Scientific Reports* 9.

Santos-Fernandez, E., J. M. Ver Hoef, E. E. Peterson, J. McGree, D. J. Isaak, and K. Mengersen. 2022. Bayesian spatio-temporal models for stream networks. *Computational Statistics & Data Analysis* 170:107446.

Stan Development Team. 2023. [RStan: The R interface to Stan](#).

Vári, Á., S. Podschun, T. Eros, T. Hein, B. Pataki, C. Ioja, C. Adamescu, A. Gerhardt, T. Gruber, A. Dedić, M. Ciric, B. Gavrilović, and A. Báldi. 2021. Freshwater systems and ecosystem services: Challenges and chances for cross-fertilization of disciplines. *Ambio* 51:135–151.

Ver Hoef, J., and E. Peterson. 2010. A moving average approach for spatial statistical models of stream networks. *Journal of the American Statistical Association* 105:6–18.

Ver Hoef, J., E. Peterson, and D. Isaak. 2019. Spatial statistical models for stream networks. Pages 421–441 in G. A. E., F. M., H. J. A., and L. S. R., editors. *Handbook of environmental and ecological statistics*. Chapman; Hall/CRC.

Wedderburn, S. D., N. S. Whiterod, and L. Vilizzi. 2022. Occupancy modelling confirms the first extirpation of a freshwater fish from one of the world’s largest river systems. *Aquatic Conservation: Marine and Freshwater Ecosystems* 32:258–268.

Winkowski, J. J., J. D. Olden, and S. Brown. 2024. Integrating spatial stream network models and environmental DNA to estimate current and future distributions of nonnative smallmouth bass. *Transactions of the American Fisheries Society* 153:180–199.

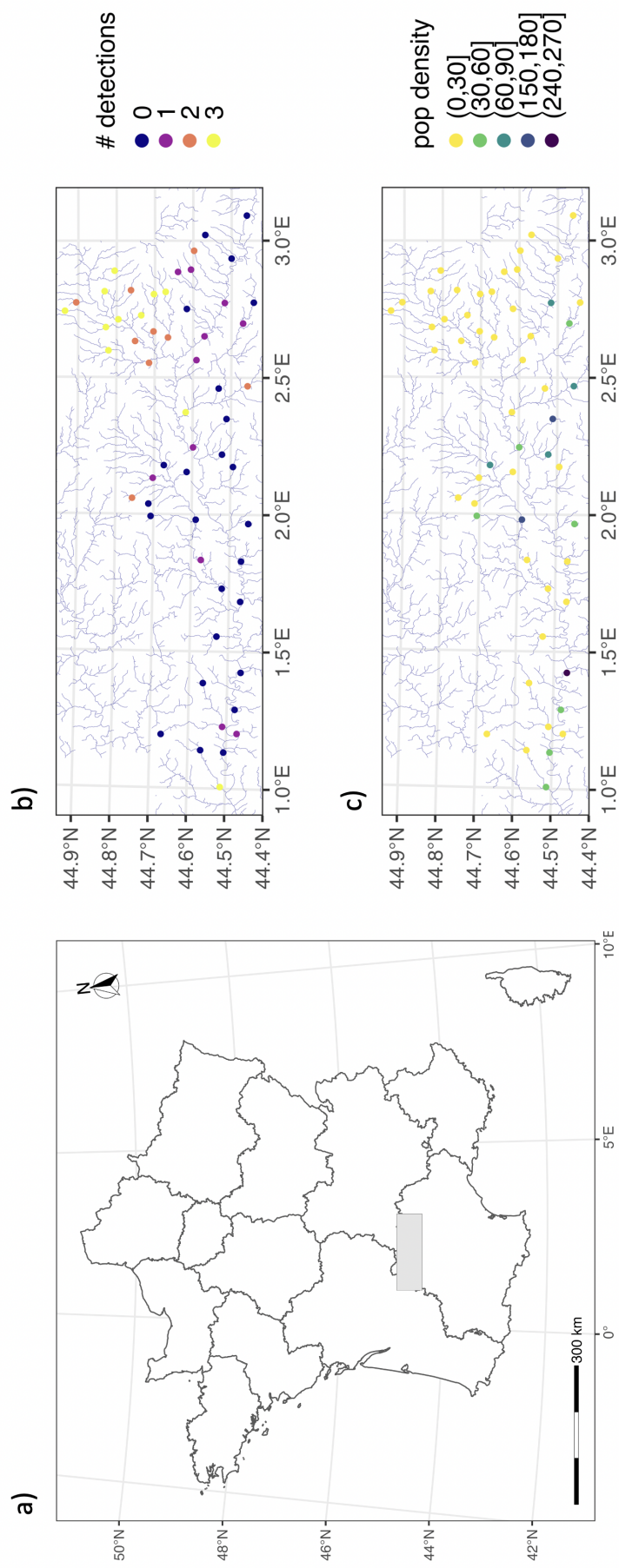


Figure 1: Information on the otter data. In panel a), the study area is given in a grey rectangle on a map of France. In panel b) the number of detections is given on a map of the study area. In panel c) the human population density is represented on a map of the study area, in number of inhabitants per km².