Title: Integrated species distribution model improves spatial uncertainty of eDNA of Edomae-fish in Tokyo Bay

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Abstract

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1 Introduction

Understanding of spatial distribution of species and underling its mechanism is a major goal n ecology. Field surveys using environmental DNA (eDNA) are widely used for detecting invasive or rare species and hotspot of biodiversity because the surveys of eDNA are easy to detect occurrence of target species, non-invasiveness, and high cost effectiveness rather than previous direct sampling method (Rees et al. 2014; Thomsen & Willerslev 2015) However, uncertainty of the occurrence of eDNA is a issue in eDNA studies because various factors, such as biological and environmental features, intricately affect the shedding (i.e., production of eDNA from organisms) and degradation (i.e., decay of eDNA from a system) of eDNA (Fig. 1). Therefore, it is necessary to consider the spatial uncertainty included in eDNA to infer the spatial distribution of species. After the suggestion of the importance to understand about origin, state, transport, and 12 fate of eDNA ("ecology of eDNA") by Barnes & Turner (2016), the studies which used laboratory experiments and numerical hydrological models have been increased to overcome the uncertainty of eDNA. For example, the relationship between production rate 15 of eDNA and biological factors (e.g., species and body mass) or environmental factors (e.g., 16 temperature), and between decomposition of eDNA and the sate of eDNA (e.g., length of 17 eDNA) or environmental factors were measured. In numerical hydrological models, spatial 18 distribution of eDNA was simulated in aquatic area and obtain the production, transport and 19 decay of eDNA (e.g., Fukaya et al. 2020). However, keeps for all focal species in laboratory 20 are not realistic and low cost-efficiency when multi species studies such as biodiversity

monitoring and fisheries management. Moreover, biologists monitoring species are not often

familiar with hydrological models and the simulation of eDNA seems to be hard for them.

Integrated species distribution models (IDMs) are now common spatial model to 24 predict spatial pattern of species (Issac et al. 2020). The models combine the different type of data with strengths and weaknesses in a single model. For example, the model can use 26 both scientific survey data, which is high quality but less abundant due to restriction of cost, and citizen data, which is widely collected and abundant but may be low quality due to not 28 using consistent field methods in a model. Combining both types of data can capitalize on 29 the strengths of each data and perform better prediction than models when using single data (Pacifici et al. 2017; Miller et al. 2019). Hence, if we develop a IDM that uses not only the occurrence data of eDNA but also the second data which contains the information that species exists at a location, as well as environmental factors as covariance in the model, then the inference from the model should represent the reduction of the spatial uncertainty of eDNA and improvement (Fig. 1). 35

In this paper, to infer spatial distribution of species from the occurrence data of eDNA,
we first make a spatial distribution model which considers spatial uncertainty of eDNA by
using an integrated spatial distribution model. We then applied the model to both eDNA data
and catch weight data for four fish in Tokyo Bay, Japan, and predicted the spatial
distribution of these fish. We finally compared the differences in (i) the predicted spatial
distribution of eDNA among fish and (ii) the relationship between the occurrence of eDNA
and environmental factors among fish to discuss the effects of the biological factors on the
spatial uncertainty of eDNA.

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2 Materials and Methods

46 2.1 A general model to estimate species distribution from eDNA

Integrated spatial distribution model that account for explicitly spatial autocorrelation in occurrence were built by Pacifici et al. (2017), which shows three approaches to predict the spatial distribution of species: the joint likelihood (shared), correlation, and covariate 49 methods. The joint likelihood method uses multiple data types to simultaneously estimate a 50 shared set of parameters with constraining that the likelihoods of shared set of parameters to be equal across. The correlation method connects multiple data types indirectly through a shared covariance matrix that captures similar patterns present in each data sources. The covariate method incorporates information from a added dataset via a fixed effect. Although each methods estimate the spatial distribution of species using multiple data 55 sets, we need to select method depending on the data features for analysis because there are 56 strengths and weaknesses (Pacifici et al. 2017; Miller et al. 2018). The joint likelihood 57

sets, we need to select method depending on the data features for analysis because there are
strengths and weaknesses (Pacifici et al. 2017; Miller et al. 2018). The joint likelihood
method may be problematic when the second data is of poorly quality compared to
correlation and covariate methods because each data can directly inform the latent
occurrence state (probabilities?) and the weight given to estimate the parameters is naturally
determined by their relative size and quality. Thus, it is not the best method when our
second data is low quality while it is the best method when our second data is high quality
(vise versa). The correlation method is added robustness to the joint likelihood because the
second data indirectly inform the occurrence state. Thus, it is the best method when our
second data is low quality while it is inferior to the joint likelihood method when both data
are deemed reliable. The covariate method does not make full use of the information in the

67 second data because the second data as a constructed covariate in the mean occurrence state.

In addition, this method can reduce the computational cost because there are fewer
parameters to estimate and the number of data locations can be reduced. Thus, it is the best
method when the second data is low quality and/or there is computational limitation while it
may not the best method when the information of the second data is needed.

When predicting the spatial distribution of species from eDNA using integrated species distribution model, the information that a species exists is needed as second data to consider spatial uncertainties of eDNA due to complex factors (Fig. 1). Hence, the second data is preferred to high quality as possible.

1. しかし、eDNA は直接的なモニタリングに比べて簡易的であるためより広い範囲で取得されている可能性が高く、eDNAのデータと同様の空間範囲で調査データのように質の高いデータを取得することは難しいかもしれない。その一方で、eDNAの空間的な不確実性を考慮するためには、種がいた証拠である2番目のデータの情報をeDNAのデータにしっかりと伝える必要がある。これらを考えると、integrated spetial distribution modelを用いたeDNAからの空間分布の推定には、以下のような correlation method が適切である:

$$p_{e}(s_{i}) = \alpha_{e} + \sum_{k} f_{e,k}(x_{e,k}(s_{i})) + w\theta(s_{i}) + u_{e}(s_{i})$$

$$p_{a}(s_{i}) = \alpha_{a} + \sum_{k} f_{a,k}(x_{a,k}(s_{i})) + \theta(s_{i}) + u_{a}(s_{i})$$
(1)

where α and $x_k(s_i)$ are the intercept and the covariates at sites i for occurrence probabilities at sites i of the added data (p_a) and eDNA data (p_e) , respectively. $u(s_i)$ is spatial error that is specific for each data following multivariate normal distributions $MVN(0, \mathbf{R})$, where the variance—covariance matrix \mathbf{R} is a Matérn correlation function. θ which is shared between two equations is the common spatial pattern between the two data, which cannot explain by each terms of the equations. That is, θ can be interpreted as "true" spatial distribution of species.

2.2 An application to a eDNA and catch data in Tokyo Bay

91 **2.2.1 eDNA data**

92 Field surveys

Field surveys were conducted by prefectural experimental station in Chiba, following the consistent sampling design at 14 sites in Tokyo Bay from April to December in 2018 (Fig. 1). In each sites, seawater and environmental data were simultaneously collected. For eDNA 95 analysis, two litter of bottom seawater was collected using a Niskin water sampler, and then 96 it was separated for two 1L samples for replicate. Each samples filtered glass fiber membrane GF/F (0.7 μm pore size; Cytiva, Sheffield, UK) onboard and then the filters were 98 frozen on a block of dry ice. These frozen filters were stored at -30° in the laboratory until eDNA extraction. To lower the levels of cross-contamination, equipments for eDNA 100 sampling were changed new one or washed in each sites. During sampling the bottom seawater, seawater temperature, salinity, pH, and dissolved oxygen (DO) at the same depth of seawater sampling for eDNA were measured by CTD $(\cancel{x} - \cancel{n} -)$. 103

104 Laboratory experiments

In laboratory, eDNA extraction, eDNA amplification, and eDNA sequence were conducted.

Total eDNA was extracted from the frozen filters using a DNeasy Blood and Tissue Kit

(Qiagen, Hilden, Germany) following Yamamoto et al. 2019. Mitochondorial 12S rRNA

gene was amplified using MiFish universal primers referring to Miya et al. 2015 with slight modification. The details was shown in Hongo et al. (受理されてないようだったら書くしかない). eDNA sequence were

2.2.2 Catch statistics

A part of catch statistics of small-scale bottom trawl fisheries recorded by several representative boats of Chiba Prefecture were provided by Chiba Prefecture. This data included date, geographic location, efforts (number of tows), gear, and catch weight (kg) in each fish. Almost of all gear was beam trawl although dredge net also used. The species which also detected by eDNA was *Conger myriaster* (マアナゴ), *Kareius bicoloratus* (イシガレイ), *Lateolabrax japonicus* (スズキ), and *Konosirus punctatus* (コノシロ). Thus, we estimated the spatial distribution of these four species using the eDNA-IDM. マコガレイ, カマス類, クロダイ, イシモチ類も解析できる??

2.2.3 Estimation of spatial distribution

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To estimate the spatial distribution of four focal species using eDNA and catch data by considering with spatial uncertainties of eDNA, we fitted the model (equation 1) to the presence/absence of eDNA and of catch collected in Tokyo Bay as follows:

logit
$$p_e(s_i) = \alpha_e + \sum_k f_k(x_k(s_i)) + w\theta(s_i) + u_e(s_i)$$

logit
$$p_c(s_i) = \alpha_c + \beta_i + \theta(s_i) + u_c(s_i)$$

where α is the intercept, and $x_k(s_i)$ is the covariates at sites i for occurrence probabilities of eDNA at sites i. In the study, seawater temperature, salinity, pH, and DO were used as

covariates which effect on the occurrence of eDNA (i.e., k=4). $u(s_i)$ is spatial error that is specific for each data following multivariate normal distributions $MVN(0,\mathbf{R})$, where the variance—covariance matrix \mathbf{R} is a Matérn correlation function. θ which is shared between eDNA and catch is the common spatial pattern between the two data, which cannot explain by each terms of the equations. That is, θ can be interpreted as the spatial distribution of species we want to know. 共変量の非線形性について書く Parameters in this model was estimated by Integrated Nested Laplace Approximation using using the R-INLA package (Lindgren, 2012) in R 3.6.1 (R Development Core Team, 2019).

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138 Authorship

YK conceived of the research idea. YH, YU, HM, MI, KA, and AK conducted field sampling. YH performed the laboratory experiments. YK, HO, and SN designed statistical analyses. YK wrote programs and performed the analyses. YK wrote the manuscript with input from all co-authors' comments.

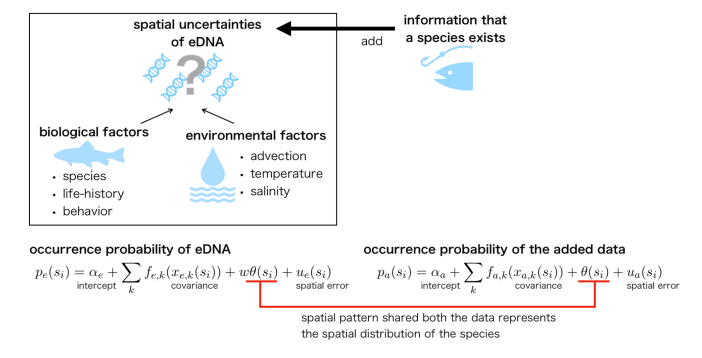


Fig. 1: Conceptual diagram of this study.