

Spatial models of carbon, nitrogen and sulphur stable isotope distributions (isoscapes) across a shelf sea: An INLA approach

Katie St. John Glew¹  | Laura J. Graham²  | Rona A. R. McGill³ |
Clive N. Trueman¹ 

¹Ocean and Earth Science, University of Southampton, Southampton, UK

²Geography and Environment, University of Southampton, Southampton, UK

³NERC LSMSF, Scottish Universities Environmental Research Centre, Glasgow, UK

Correspondence

Katie St. John Glew
Email: katie.stjohn@oton.ac.uk

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Abstract

1. Spatial models of variation in the isotopic composition of structural nutrients across habitats (isoscapes) offer information on physical, biogeochemical and anthropogenic processes occurring across space, and provide a tool for retrospective assignment of animals or animal products to their foraging area or geographic origin. The isotopic differences among reference samples used to construct isoscapes may vary spatially and according to non-spatial terms (e.g. sampling date, or among individual or species effects). Partitioning variance between spatially dependent and spatially independent terms is a critical but overlooked aspect of isoscape creation with important consequences for the design of studies collecting reference data for isoscape creation and the accuracy and precision of isoscape models.
2. We introduce the use of integrated nested Laplace approximation (INLA) to construct isoscape models. Integrated nested Laplace approximation provides a computationally efficient framework to construct spatial models of isotopic variability explicitly addressing additional variation introduced by including multiple reference species (or other recognized sources of variance).
3. We present carbon, nitrogen and sulphur isoscape models extending over c. 1 million km² of the UK shelf seas. Models were built using seven different species of jellyfish as spatial reference data and a suite of environmental correlates. Compared to alternative isoscape prediction methods, INLA-spatial isotope models show high spatial precision and reduced variance. We briefly discuss the likely biogeochemical explanations for the observed spatial isotope distributions. We show for the first time that sulphur isotopes display systematic spatial variation across open marine shelf seas and may therefore be a useful additional tool for marine spatial ecology.
4. The INLA technique provides a promising tool for generating isoscape models and associated uncertainty surfaces where reference data are accompanied by multiple, quantifiable sources of uncertainty.

KEYWORDS

Bayesian spatial modelling, carbon, isoscape, jellyfish, nitrogen, stable isotopes, sulphur, UK shelf seas

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1 | INTRODUCTION

The isotopic ratio of elements such as hydrogen, oxygen, carbon, nitrogen and sulphur varies systematically across the natural environment. Modelling these spatial differences through mechanistic or statistical models (isoscares [West, Bowen, Dawson, & Tu, 2010]) offers insight into the biogeochemical processes leading to spatial variation in isotopic expressions, and provides a tool for the retrospective assignment of animals or animal products back to their origin or foraging area (Hobson, Wassenaar, & Taylor, 1999), with broad uses in animal and human migration and tracking (Ehleringer et al., 2008; Hobson, 1999; Hobson, Barnett-Johnson, & Cerling, 2010), trophic ecology (Jennings & van der Molen, 2015; Olson et al., 2010) and traceability within consumer goods supply chains (Chesson, Valenzuela, O'Grady, Cerling, & Ehleringer, 2010; Kelly, Heaton, & Hoogewerff, 2005). Isoscares have been used extensively in terrestrial ecological and forensic applications, particularly isoscares describing spatial variations in hydrogen and oxygen isotopes of precipitation (Bowen, 2010). Spatial variation in isotopic compositions in marine systems has also been explored (Cherel & Hobson, 2007; Schell, Saupé, & Haubenstock, 1989). However, relatively few continuous surface isoscares have been published in marine compared to terrestrial systems, probably due to the difficulty in obtaining sufficient reference samples over appropriate spatial and temporal scales.

To construct a continuous surface isoscare model, isotopic compositions of reference materials or organisms are typically projected across space using either spatial interpolation methods (Trueman, MacKenzie, & St John Glew, 2017; Vander Zanden et al., 2015), by statistical inference based on correspondence between measured data and environmental correlates (Bowen & Wilkinson, 2002; Courtiol & Rousset, 2017; Jennings & Warr, 2003) or a combination of both (MacKenzie, Longmore, Preece, Lucas, & Trueman, 2014; Wunder, 2010). Ideally, the isotopic composition of reference samples should vary only according to spatially dependent effects. However, additional isotopic variance among reference samples is commonly introduced through processes such as collection of samples over different time-scales or differences in ecology or physiology among individuals or species. As the spatial scale of a study area increases, the ease of collecting uniform reference samples generally decreases, especially where isoscare reference data are compiled from opportunistically collected samples. Accounting for and quantifying spatially dependent and spatially independent variance is a key component of isoscare model creation. In simple interpolation models, the variance associated with the prediction of expected isotopic compositions at any point in space increases with distance from discrete sampling points; therefore, irregular spacing of reference samples produces spatial gradients in isoscare uncertainty, which may bias interpretations. When environmental correlates are introduced, estimating variance becomes more complex due to error associated with the relationship between measured data and environmental correlates, which is itself spatially varying, but rarely quantified across space (Bowen & Revenaugh, 2003; Courtiol &

Rousset, 2017). To date, many isoscare models either assume spatially invariant uncertainty in the relationship between measured data and environmental correlates (Jennings & Warr, 2003), infer spatial variance by interpolating residuals from regression models (MacKenzie et al., 2014) or draw on resampling methods to estimate spatially varying uncertainty (Wunder & Norris, 2008). Courtiol and Rousset (2017) introduced a frequentist mixed modelling approach that enables spatially explicit variance surfaces to be calculated by including location as a random effect but at the cost of slow computational processing. Here, we introduce an alternative approach to isoscare generation based on integrated nested Laplace approximations (INLAs). We aim to address the common issue of limited sample availability by modelling isoscares using multiple species and explicitly addressing spatial isotopic variation due to mixed sample sources. Many commonly used isoscare prediction methods are unable to incorporate multiple sample sources while quantifying associated spatial variance and including boundary effects (Table 1). We explore the use of recently developed Bayesian hierarchical modelling techniques using INLA. We firstly produce isoscare models for a restricted region, the North Sea, using a single jellyfish species as a spatial reference dataset, and compare the assignment accuracy and precision associated with INLA-produced and alternative North Sea isoscare models (Trueman et al., 2017). Secondly, we predict isoscares for carbon, nitrogen and sulphur across the wider UK shelf sea area using multiple reference jellyfish species.

The North Sea and wider UK shelf seas host some of the most globally productive fisheries, regionally significant oil, gas and renewable energy resources and infrastructure and intensive shipping activity. The UK shelf region has received extensive detailed investigation into spatial isotopic variability, with carbon and nitrogen isoscare models previously produced using purpose collected baseline samples, rather than commonly adopted opportunistic sampling. Barnes, Jennings, and Barry (2009b) and Jennings and Warr (2003) and Jennings and van der Molen (2015) used queen scallops *Aequipecten opercularis* from known catch locations as reference samples, coupled with environmental variables; however, variance surfaces were only calculated by Jennings and van der Molen (2015). High resolution, in situ sample-based isoscares have been modelled for the North Sea using lion's mane jellyfish *Cyanea capillata* as reference organisms through ordinary kriging of evenly spaced samples (Trueman et al., 2017) and with additional environmental variables (MacKenzie et al., 2014). Spatially explicit variance surfaces were calculated in both examples and initial assignments of invertebrate, fish and seabird samples have proven successful (St John Glew et al., 2018; Trueman et al., 2017). However, this approach is constrained by the availability and distribution of a single reference species across the region of interest, limiting marine isoscare modelling capabilities across larger spatial scales, as no single jellyfish species is distributed across the entire range of the UK shelf seas. In addition, barrier effects (e.g. uneven coastlines) are particularly important in basin scale marine isoscare predictions, yet many existing modelling techniques do not enable easy incorporation of coastlines and boundaries (Table 1).

TABLE 1 Comparison of isoscape prediction methods and their ability to incorporate multiple species sources, environmental data, and boundary effects while explicitly quantifying spatial variance

Method	Able to incorporate multiple species?	Able to incorporate environmental data?	Able to take into account boundary effects?	Able to quantify spatial variance due to species random effect?	Fast processing time?
Simple interpolation/kriging of sample data	X	X	X	X	Y
Linear regression models of sample data (with species as a random effect) and additional environmental variables, followed by interpolation	Y	Y	X	X	Y
General additive models of sample data (with species as a random effect) and additional environmental variables, followed by interpolation	Y	Y	X	X	Y
Ocean Data View: Data Interpolating Variation Analysis	X	X	Y	X	Y
Mixed modelling with spatial dependency term.	Y	Y	X	Y	X
Bayesian hierarchical spatial modelling (integrated nested Laplace approximation)	Y	Y	Y	Y	Y

2 | MATERIALS AND METHODS

2.1 | Data collection and stable isotope analysis

To construct isoscape models of UK shelf seas, we collected 627 jellyfish samples of seven different species (Barrel *Rhizostoma pulmo*, Blue *Cyanea lamarckii*, Compass *Chrysaora hysoscella*, Crystal *Aequorea victoria*, Lion's Mane *Cyanea capillata*, Mauve stinger *Pelagia noctiluca* and Moon *Aurelia aurita*) from 308 stations across the UK shelf between August 2015 and December 2016 (Figure 1). Samples were collected on board the RV Cefas Endeavour (Cefas), MRVs Scotia (Marine Scotland), Thalassa (Ifremer) and RV Celtic Explorer (Marine Institute) during annual fisheries surveys. We opportunistically collected further samples from small commercial fisheries, research and private vessels. Jellyfish were collected, identified, weighed and measured on-board before thorough washing with salt water and immediately freezing to -20°C . In the laboratory, samples were thawed, washed repeatedly (Mackenzie, Trueman, Lucas, & Bortoluzzi, 2017) and a section of bell tissue (mesoglea) removed and refrozen prior to freeze-drying for 24 hr, subsampling and submission for isotopic analysis.

We analysed all 627 samples across the UK shelf sea for $\delta^{13}\text{C}$, $\delta^{15}\text{N}$ and $\delta^{34}\text{S}$ at the Life Sciences Mass Spectrometry Facility (LSMSF), East Kilbride, United Kingdom, in autumn 2017. A subset of North Sea lion's mane jellyfish samples (57) from 51 stations was also analysed for $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ at Elementex laboratories, Cornwall, United Kingdom, in autumn 2015. Accuracy and precision were monitored through laboratory internal standards (LSMSF: MSAG, M2 and SAAG2) and an in-house comparison standard (ARCOS glutamic acid) nested within samples. We compared North Sea samples analysed in both laboratories for consistency and no significant

differences were observed between results ($\delta^{13}\text{C}$: $t = -0.38$, $p > 0.05$, $\delta^{15}\text{N}$: $t = 0.31$, $p > 0.05$).

Jellyfish bell tissue $\delta^{13}\text{C}$ values showed a significant negative linear relationship with C:N ratios ($p < 0.005$, slope = -2.22 , adjusted $R^2 = 0.06$). To correct for lipid-related variance in $\delta^{13}\text{C}$ values, we applied an algebraic correction (Kiljunen et al. (2006)). Lipid-corrected carbon and nitrogen isotopic data from Queen scallops of known location were taken from Jennings and Warr (2003) and Barnes et al. (2009b), for scallops collected between 25 July and 29 September 2001 and from Barnes, Jennings, and Barry (2009a) for scallops collected in 2010.

We estimated within-species variation in jellyfish stable isotope compositions by averaging (mean) the among-individual standard deviation of the same species occurring at the same sampling location. We calculated among-species average isotopic differences by calculating the mean isotopic difference between species at the same location and then averaging across all locations.

2.2 | Environmental data

We downloaded chlorophyll (Chl-a) and night-time short-wave length sea surface temperature (SST) level three (instrument calibrated) monthly average data from the MODISA satellite (NASA Goddard Space Flight Center, 2014) between March and September in 2015 and 2016 over the spatial range of the UK shelf seas at 4-km resolution (Figure 2). We downloaded bottom temperature (BT), surface salinity and mixed layer depth (MLD) daily mean data from the Forecasting Ocean Assimilation Model 7 km Atlantic Margin model (FOAM AMM7) at a 0.11 by 0.07 degree resolution across the UK shelf seas between March and September in 2015 and 2016 (CMEMS, 2017). We calculated 2-year median spring-summer raster surfaces for each variable

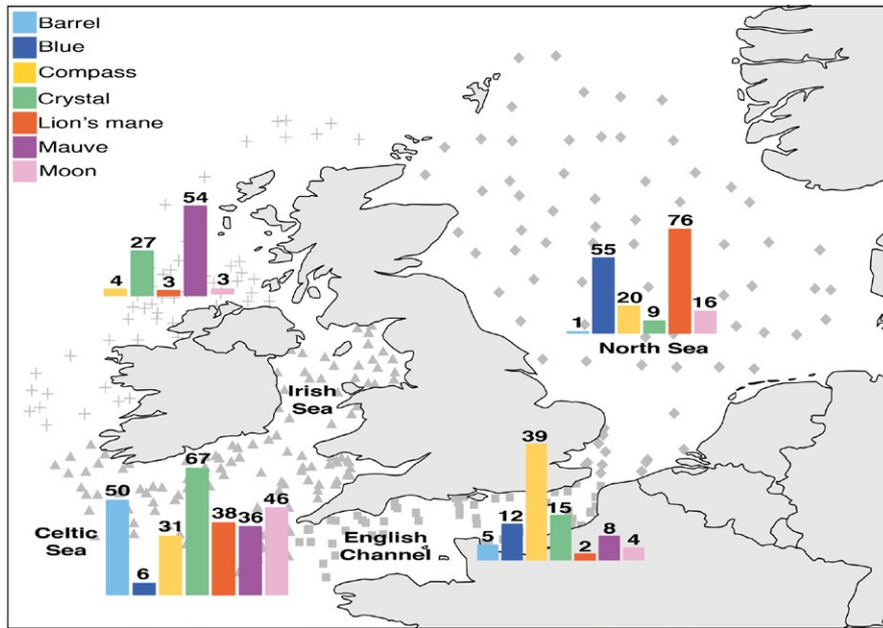


FIGURE 1 Jellyfish sampling locations around the UK shelf seas and a summary of the number of each individuals of each species collected within the North Sea (diamond), English Channel (square), Celtic and Irish Seas (triangle) and off West Scotland and Ireland (cross)

(Figure 2) and a temperature difference (Tdiff) surface by subtracting bottom temperature from SST raster surfaces. We acquired water column depths from NOAA bathymetry database at 1° resolution (Figure 2). Raster surfaces were resampled to $0.1 \times 0.1^\circ$ resolution over the coordinates $(-13, 8, 48, 62)$. We extracted covariate values at jellyfish sampling points and scaled the data by subtracting the variable mean from each value and dividing by the variable standard deviation (Figure 2).

2.3 | Model formation

We describe a Bayesian hierarchical spatial modelling framework for the shelf seas, predicting $\delta^{13}\text{C}$, $\delta^{15}\text{N}$ and $\delta^{34}\text{S}$ values using INLA via the *R-INLA* package (<http://www.r-inla.org>) (Rue, Martino, & Chopin, 2009). This approach differs from the frequentist mixed modelling approach introduced by Courtiol and Rousset (2017) by adopting a Bayesian framework enabling uncertainty to be more easily interpretable, allowing the inclusion of boundary effects and solving the spatial dependency term in an alternative and faster way.

When modelling across a spatial range, ordinary linear regression ignores spatial dependency between sampling locations. Through the latent Gaussian field with Matérn correlation, *R-INLA* provides a means to explicitly incorporate spatial dependency:

$$y_{(si)} \sim N(x_{(si)}, \sigma^2), x_{(si)} = \text{covariates}_{(si)} + u_{(si)},$$

where $y_{(si)}$ are the response values at all sampling locations which are assumed to be normally distributed with mean $x_{(si)}$ and variance σ^2 . $u_{(si)}$ is the spatial dependency random effect. *R-INLA* includes a stochastic partial differential equation (SPDE) approach that allows fast modelling of Gaussian Random Fields (GRFs) similar to kriging approaches, but is better adapted to handling data with complex spatial structures (Lindgren, Rue, & Lindström, 2011). The SPDE approach

enables the covariance matrix of the Gaussian field to be approximated as a Gaussian Markov Random Field (GMRF) using a Matérn covariance structure and Delaunay triangulation to create prediction locations in the form of a mesh (Figure 3). Observations are treated as initial vertices; then, further vertices are added with the aim of reducing the number of triangles required, but including all observations in denser sampled areas.

To compare between isoscape models created through INLA and previously described kriging approaches (Trueman et al., 2017), we developed an INLA North Sea model using reference data from lion's mane jellyfish. For the UK shelf sea model, we included observations of all seven jellyfish species and species identity was a random effect. All individual jellyfish data were included in the model, including in locations where multiple individuals of the same or different species were sampled at the same location. Put simply, we modelled each isotope value as a function of a set of covariates X_i where species and the underlying spatial structure were included as random effects. Models were specified as:

$$Y_i \sim \text{Intercept} + \beta_i X_i + f(U_i) + f(W_i) + \varepsilon_i,$$

$$U_i \sim N(0, \sigma_{\text{species}}^2),$$

$$W_i \sim N(0, \Omega),$$

$$\varepsilon_i \sim N(0, \sigma^2),$$

where Y_i is the isotope value ($\delta^{13}\text{C}$, $\delta^{15}\text{N}$ or $\delta^{34}\text{S}$) at location i , X_i is a vector containing the environmental covariates as linear fixed effects, β_i is a vector of parameters to be estimated, U_i is the species random effect with assumed Gaussian distribution, W_i represents the smooth spatial effect, linking each observation with a spatial location, with the elements of Ω estimated using

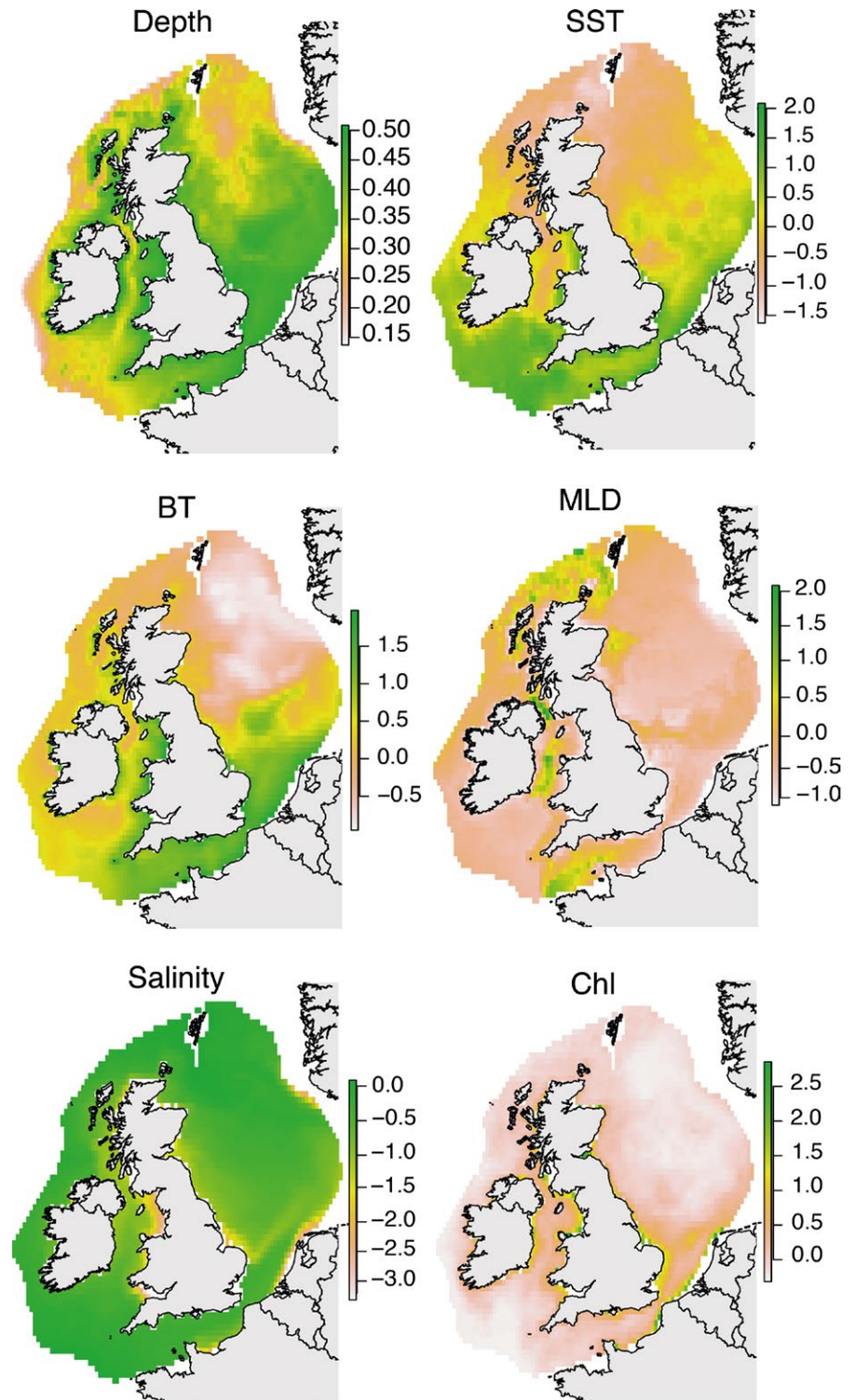


FIGURE 2 Scaled environmental covariate raster surfaces (depth, sea surface temperature (SST), bottom temperature (BT), mixed layer depth (MLD), salinity and chlorophyll (Chl))

the Matérn correlation, and ϵ_i contains the independently distributed residuals. Full global models including all environmental covariates, specifying no interaction terms and first-order interaction terms were tested (Table 3), as we had no prior expectations of covariate or interaction significance. Model selection and inclusion or exclusion of interactions were based on deviance information criterion (DIC) and model fit (Pearson correlation between predicted and observed values). Within the North

Sea, we also compared models with and without a spatial effect. When similar DIC values were observed (within 2), the simpler model was selected (Burnham & Anderson, 2003). We only compared spatial models within the shelf seas to capture and model isotopic spatial variation driven both by the larger spatial extent of the UK shelf study area, and additional variance introduced by multispecies reference samples. We used non-informative default priors for each model.

We used the best model for each isotope to predict isotopic compositions across the whole spatial domain using continuous raster surfaces of scaled environmental variables as predictors. Response variables were estimated at all mesh vertices, which were then linearly interpolated within each triangle into a finer regular grid ($0.2 \times 0.2^\circ$) via Bayesian kriging. To avoid extrapolating beyond the environmental covariates range, we masked grid cells where predictor covariate values fell largely outside the range of values observed at jellyfish sampling locations. This was particularly important when incorporating depth as a covariate. All jellyfish samples were collected on the UK continental shelf; therefore, the prediction area was also limited to the shallower shelf sea range, in order to prevent skewed isotope predictions in deeper unsampled regions. Mean and variance predictions were obtained for each grid cell and isotope and mapped to produce isoscapes and model variance surfaces representing expected isotopic compositions for jellyfish (or a similar pelagic generalist consumer) accounting for the species random effect.

2.4 | Comparing INLA and kriging isoscape models for single species isoscapes

To assess the differences between the traditional ordinary kriging and the INLA isoscape prediction approaches, we compared North Sea carbon and nitrogen isotope prediction and variance surfaces (Figure 4). INLA prediction surfaces were subtracted from the Trueman et al. (2017) North Sea isoscapes and results are displayed as difference surfaces. Following the methods described in Trueman et al. (2017), we assigned scallops from known locations to their most likely origin within the North Sea (Figure 5) based on similarity between measured isotopic compositions and isoscape predictions using multivariate normal probability distributions. Assignments were made by estimating the likelihood that each raster cell of the carbon and nitrogen isoscapes represented the origin of each individual. We displayed assignment accuracy and precision results as per Trueman et al. (2017) using odds ratios to set probability threshold values to differentiate between cells of likely and unlikely origin. Assignment precision

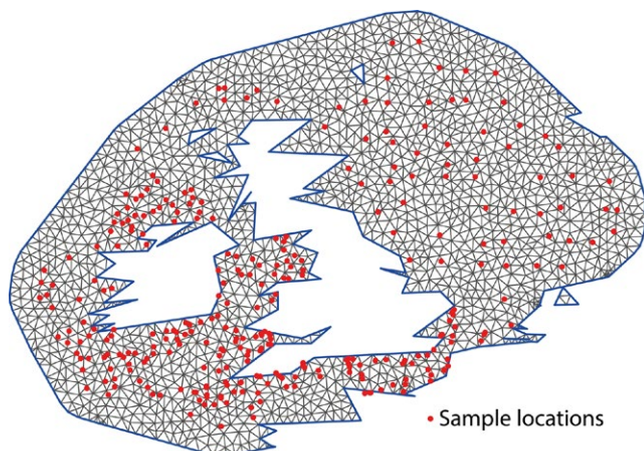


FIGURE 3 Delaunay triangulation mesh designs for the UK shelf sea model. Sampling locations are indicated in red

was defined by the odds ratio threshold and represents the proportion of the surface area with probability values above this set threshold, and assignment accuracy is defined as the proportion of individuals where the true location falls within the assigned area (Trueman et al., 2017; Vander Zanden et al., 2015). We compared accuracy and precision of known-origin scallops assignments sampled in 2001 and 2010 between INLA-predicted North Sea isoscapes and ordinary kriging isoscapes produced by Trueman et al. (2017). We performed all analyses using R 3.4.2 (R Core Development Team, 2016).

3 | RESULTS

3.1 | Within- and between-species variability

The variations in average stable isotope ratios within species sampled at the same locations were relatively consistent across species in both carbon (0.37–0.62‰) and sulphur (0.41–0.63‰) apart from crystal jellyfish where among-individual variation was higher for both $\delta^{13}\text{C}$ (1.08‰) and $\delta^{34}\text{S}$ (0.75‰) (Table 2). Within-species differences in nitrogen were more variable ranging from 0.44‰ in blue jellyfish to 1.65‰ in crystal jellyfish (Table 2). Among-species differences ranged considerably between species and isotope (Table 2). $\delta^{34}\text{S}$ differences were relatively constrained with differences ranging from 0.01 to 1.45‰ whereas $\delta^{13}\text{C}$ differences ranged from 0.03‰ (between mauve stingers and compass jellyfish) to 3‰ (between barrel and compass jellyfish). Among-species differences in $\delta^{15}\text{N}$ varied over the largest range, from 0.02‰ between lion's mane and blue jellyfish up to 7.1‰ between barrel and mauve jellyfish.

3.2 | The North Sea isoscape models

The best-fit carbon and nitrogen isoscape models for the North Sea were non-spatial models excluding interaction terms (Table 3):

$$y \sim 1 + X + \text{SST} + \text{BT} + \text{Tdiff} + \text{Chl} + \text{MLD} + \text{Sal} + \text{Depth}.$$

In both cases, interaction and spatial dependency terms did not improve model fit; therefore, the simplest models were selected (Table 3).

Broad spatial patterns in $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ ranges (Figure 4a,c, respectively) are consistent with previous studies (Jennings & van der Molen, 2015; Trueman et al., 2017), indicating that isoscape surfaces predicted within an INLA modelling framework are comparable with more traditional ordinary kriging approaches. Associated variance surface values for carbon and nitrogen (Figure 4b,d) are considerably lower than those calculated using ordinary kriging approaches (Trueman et al., 2017), with both carbon and nitrogen variance values predominately below 1‰.

3.3 | Method comparison and scallop assignment

Both the INLA approach and the Trueman et al. (2017) ordinary kriging approach predicted similar carbon and nitrogen isoscape

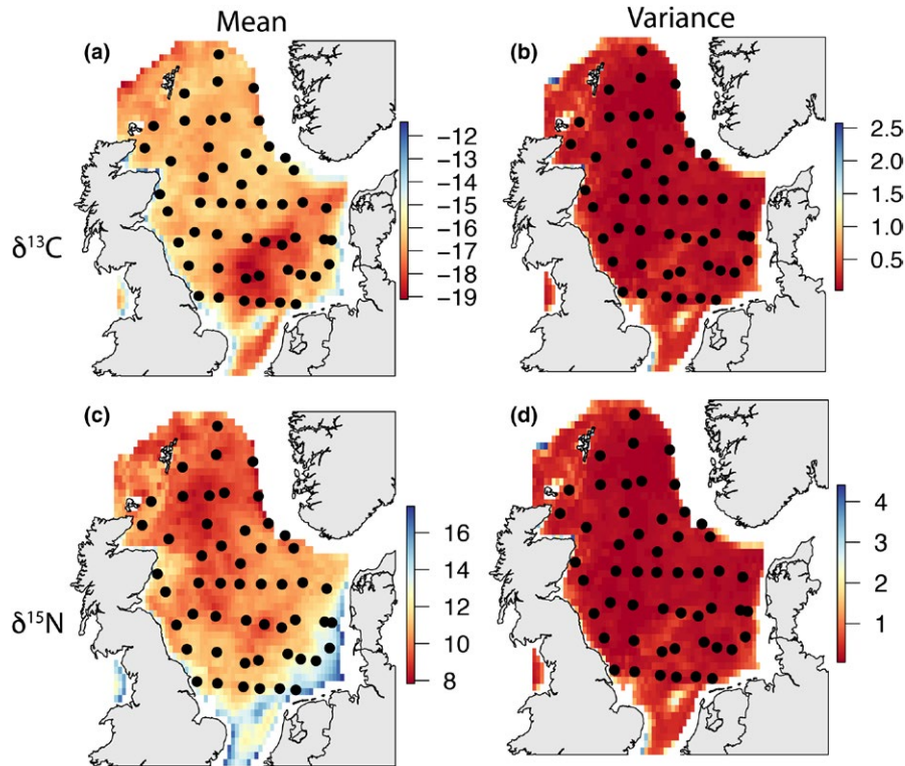


FIGURE 4 North Sea carbon (a) and nitrogen (c) isoscape models and associated variance of the posterior predicted distribution surfaces (b, d). Values based on *Cyanea capillata* sampled in August 2015. Filled circles represent sampling locations

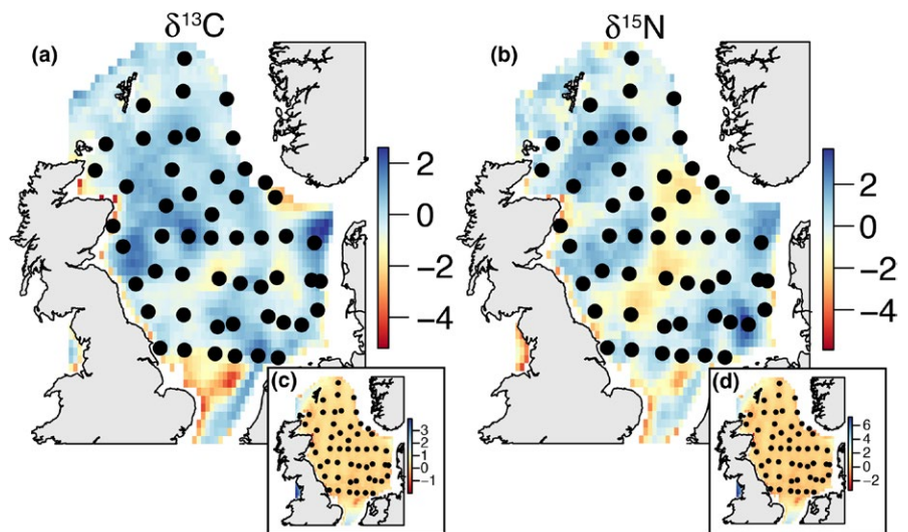


FIGURE 5 Difference between integrated nested Laplace approximation (INLA) predicted and Trueman et al. (2017) kriging predicted carbon (a) and nitrogen (b) isoscape surfaces (kriging–INLA), and the respective difference between the variance surfaces (c, d)

surfaces for the North Sea, with isotopic differences in each grid cell primarily falling between ± 1 – 1.5% (Figure 5). The regions where isotopic difference is seen to be greater (2–4%) are the areas where no jellyfish samples occur, such as the southern North Sea. Larger differences in nitrogen isotope predictions are also seen across the central North Sea (Figure 5b), demonstrating the strong depth influence on INLA-predicted isotope value. Minimal differences were observed in the carbon and nitrogen variance surfaces (Figure 5c,d) in the range where samples were collected.

Accuracy and precision of known catch location scallop assignments to the original Trueman et al. (2017) North Sea isoscapes and the INLA-modelled isoscapes were compared over a range of odds ratio threshold values (Figure 6). Assignment accuracy to the new INLA-modelled isoscapes was better than random at all precision values (Figure 6). Assignment accuracy for scallops sampled in 2001 and 2010 was over 90% when assigning to areas representing on average over 40% of the total North Sea isoscape area. When precision was increased to an area representing 20% of the isoscape, assignment accuracy was greater than 70% for both scallop

	Barrel	Blue	Compass	Crystal	Lion's mane	Mauve	Moon
Carbon (‰)							
Barrel	0.37	NA	-3.0	-2.16	-2.73	-0.93	-2.34
Blue	0	0.62	1.14	-0.40	0.26	2.22	2.27
Compass	5	11	0.59	0.74	0.63	-0.03	0.63
Crystal	8	5	16	1.08	1.05	0.71	0.06
Lion's Mane	4	24	9	10	0.60	0.84	0.39
Mauve	1	1	7	30	1	0.48	-0.71
Moon	5	9	11	21	8	14	0.46
Nitrogen (‰)							
Barrel	0.79	NA	-2.08	-1.85	-1.37	7.10	-0.60
Blue		0.44	0.49	-1.47	0.02	1.63	0.78
Compass			0.52	-3.55	-0.30	-0.19	-3.68
Crystal				1.65	1.34	5.22	2.54
Lion's Mane					0.62	2.97	0.32
Mauve						0.95	-5.25
Moon							0.63
Sulphur (‰)							
Barrel	0.45	NA	0.13	0.37	0.83	1.45	0.88
Blue		0.62	-0.74	0.65	-0.1	-0.01	-0.49
Compass			0.41	0.15	0.53	-0.19	-0.39
Crystal				0.75	-0.62	0.50	-0.34
Lion's Mane					0.44	0.29	-0.47
Mauve						0.59	-0.03
Moon							0.63

TABLE 2 Within (red)- and between (row 1 - column 1)-species isotopic differences (black). Calculated at locations where multiple individuals of the same species or multiple species occur and averaged across locations. Within-species isotopic difference is the among-individual standard deviation of the same species occurring at the same sampling locations and averaged across all locations. Between-species isotopic difference is the difference between different species sampled at the same locations and averaged across all locations. Blue numbers indicate the number of locations where pairs of species were sampled

datasets. At higher precision values, assignment of the 2001 dataset to the INLA-modelled isoscapes appears more accurate than assignment to the kriging isoscapes by Trueman et al. (2017), whereas the opposite is observed with the 2010 scallop dataset. Overall assignment to the original kriging isoscapes is slightly more accurate than to the INLA-modelled isoscapes, but both methods are largely comparable.

3.4 | UK shelf sea isoscape models

Global models, including first-order interaction terms, were the best-fit for carbon, nitrogen and sulphur isoscapes;

$$\begin{aligned}
 y \sim & 1 + X + \text{SST} + \text{BT} + \text{Tdiff} + \text{Chl} + \text{MLD} + \text{Sal} + \text{Depth} \\
 & + \text{SST:BT} + \text{SST:Tdiff} + \text{SST:MLD} + \text{SST:Depth} + \text{SST:Chl} + \text{SST:Sal} \\
 & + \text{BT:Tdiff} + \text{BT:MLD} + \text{BT:Depth} + \text{BT:Chl} + \text{BT:Sal} + \text{Tdiff:MLD} \\
 & + \text{Tdiff:Depth} + \text{Tdiff:Chl} + \text{Tdiff:Sal} + \text{MLD:Depth} \\
 & + \text{MLD:Chl} + \text{MLD:Sal} + \text{Depth:Chl} + \text{Depth:Sal} + \text{Chl:Sal} \\
 & + f(\text{Species}) + f(\text{Spatial})
 \end{aligned}$$

Best-fit models for carbon and sulphur UK shelf sea isoscapes had moderate fit ($R = 0.47$, $p < 0.05$ and $R = 0.50$, $p < 0.05$ respectively) (Table 3). The best fitting nitrogen isoscape model had a stronger fit ($R = 0.80$, $p < 0.05$) (Table 3).

Minimal residual isotopic variability between species remained with the chosen carbon and sulphur isoscape prediction models, indicating that the majority of species isotopic variability was able to be explained by the combination of covariates, and interactions between these covariates, included within the models (Figure 7a,c). Residual nitrogen isotopic variability has a larger range between species (c. 6‰), particularly between mauve stinger jellyfish depleted in ^{15}N and crystal jellyfish displaying relatively high $\delta^{15}\text{N}$ values (Figure 7b, Table 1).

Spatial distributions of $\delta^{13}\text{C}$ values within the North Sea are consistent with previous findings showing relatively low $\delta^{13}\text{C}$ values (-18 to -17‰) in the central North Sea and higher $\delta^{13}\text{C}$ values in the northern and southern North Sea (Figure 8a). Similar $\delta^{13}\text{C}$ values of between -17 and -16‰ are predicted within the western English Channel and into the Celtic and Irish Seas. Higher $\delta^{13}\text{C}$ values (-15.5 to -14.5‰) are predicted along the French and Belgian coasts of the English Channel and southern North Sea, off the southwest coasts of Cornwall and north of the Irish Sea (Figure 8a). Spatial distributions of $\delta^{15}\text{N}$ values are also consistent with previous North Sea predictions, with a strong isotopic gradient between the northern (8-10‰) and southern (11-13‰) North Sea (Figure 8c). Higher $\delta^{15}\text{N}$ values are also observed into the English Channel and within the Irish Sea, whereas lower $\delta^{15}\text{N}$ values are predicted around north and west Scotland and Ireland

TABLE 3 Model fit results for the carbon and nitrogen North Sea, and carbon, nitrogen, and sulphur UK shelf sea isoscape prediction models. Global models excluding and including first-order interaction terms were tested. Inclusion of the spatial term was also tested within the North Sea models. Model fit was tested using the deviance information criteria (DIC), and assessing the Pearson correlation between observed and fitted values. The t , R , 95% confidence intervals around R , and the degrees of freedom (df) are reported. The models displayed in red were the chosen models for isoscape predictions

Region	Isoscape	Model	DIC	t	R	95%	df
NS	Nitrogen	No interactions	207.5	8.3	0.75	0.60,0.84	55
NS	Nitrogen	First-order interactions	220.2	11.8	0.85	0.75,0.90	55
NS	Nitrogen	No interactions + spatial	207.2	8.3	0.75	0.60,0.84	55
NS	Nitrogen	First-order interactions + spatial	219.2	11.8	0.85	0.75,0.91	55
NS	Carbon	No interactions	176.7	5.5	0.59	0.39,0.74	55
NS	Carbon	First-order interactions	194.4	8.3	0.75	0.60,0.84	55
NS	Carbon	No interactions + spatial	176.5	5.5	0.60	0.40, 0.74	55
NS	Carbon	First-order interactions + spatial	193.4	8.3	0.75	0.60,0.84	55
UK Shelf	Nitrogen	No interactions + spatial + $f(\text{species})$	2,411.8	31.0	0.78	0.75,0.81	604
UK Shelf	Nitrogen	First-order interactions + spatial + $f(\text{species})$	2,398.6	32.9	0.80	0.77,0.82	604
UK Shelf	Carbon	No interactions + spatial + $f(\text{species})$	2,109.4	10.3	0.38	0.32,0.45	604
UK Shelf	Carbon	First-order interactions + spatial + $f(\text{species})$	2,090.4	13.0	0.47	0.40,0.53	604
UK Shelf	Sulphur	No interactions + spatial + $f(\text{species})$	1,469.8	11.9	0.43	0.36,0.50	604
UK Shelf	Sulphur	First-order interactions + spatial + $f(\text{species})$	1,458.1	14.2	0.50	0.44,0.56	604

(Figure 8c). The isotopic range in sulphur is relatively small across the shelf, with the majority of cells falling between values of 20.5 and 22.5‰, in comparison to the large variability observed in carbon and nitrogen isotope ratios (Figure 8e). The highest $\delta^{34}\text{S}$ values (>21.5‰) are observed in the northern North Sea, north Scotland and Ireland into the northwest Irish Sea (Figure 8e). A clear isotopic gradient is predicted between the northern and southern North Sea, with decreasing values into the southwest North Sea. Lowest $\delta^{34}\text{S}$ values are predicted off the southwest coast of the United Kingdom and into the Celtic Sea (Figure 8e).

Variance surfaces show broadly similar patterns for each isotope element, with low variance values (<2‰) across the majority of the shelf, and increased variance values observed within the eastern English Channel, eastern Irish Sea and in coastal regions (Figure 8b,d,f).

4 | DISCUSSION

This study has two main aims to introduce INLA as a powerful tool for creating isoscape models incorporating environmental correlates as predictors and where reference samples contain a source of variance that is not spatially dependent and to describe the spatial variation in $\delta^{13}\text{C}$, $\delta^{15}\text{N}$ and $\delta^{34}\text{S}$ values across the shelf seas of the British Isles. We have shown that INLA-generated isoscape models have

comparable accuracy and precision to simple kriging where reference samples are evenly distributed and common form (in our case the same species). We then extended the approach to draw isoscape models and uncertainty surfaces across a wide shelf sea area where collection of reference samples from a single species would be impossible.

4.1 | INLA as a tool for creating isoscape models

Creating isoscape models with associated uncertainty surfaces in regions where reference samples are either irregularly spaced and/or contain additional sources of isotopic variability is challenging (Courtillot & Rousset, 2017). Spatial modelling using the INLA approach addresses many common constraints. By using the INLA approach, we are able to incorporate environmental data and a species random effect into our isoscape prediction models. Although this can also be achieved by using a mixed effects model approach (Courtillot & Rousset, 2017), the INLA approach is unique in that it provides a computationally rapid technique to quantify the spatial variance due to the species random effect, which is essential for accurate measures of variance and subsequent isoscape assignments. In addition, INLA enables the incorporation of boundary effects, to model and predict around physical barriers, which is particularly useful in marine environments.

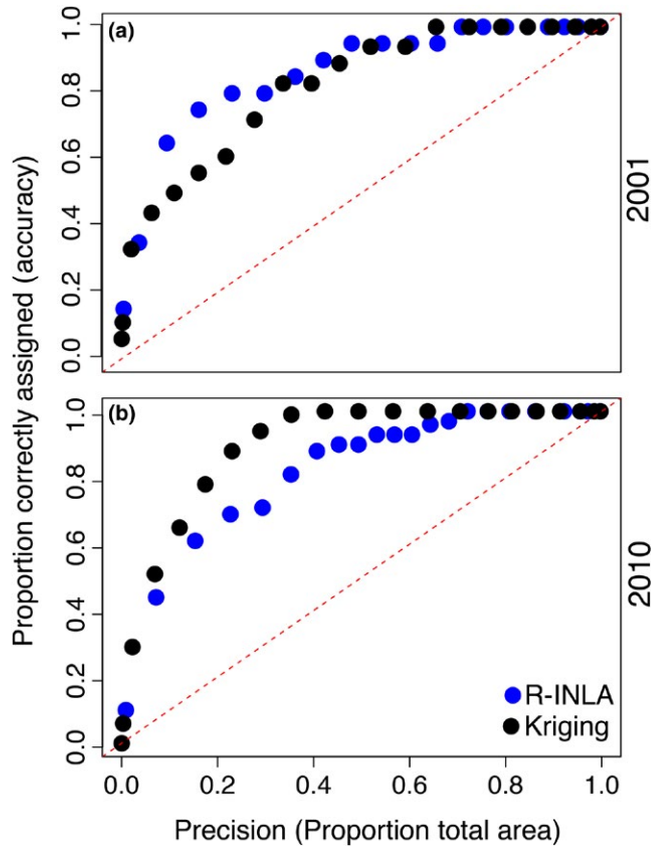


FIGURE 6 Accuracy (the proportion correctly assigned) and precision (proportion of the total surface area) of assignment to both the original North Sea kriging isoscape models (Trueman et al., 2017) shown in black, and the new integrated nested Laplace approximation (INLA) modelled North Sea isoscapes shown in blue for the 2001 and 2010 scallop datasets. The red line represents the accuracy and precision values if assignments were no better than random

The INLA-predicted North Sea $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ isoscapes (Figure 4) are broadly similar to isoscapes produced from ordinary kriging of identical lion's mane jellyfish data (Trueman et al., 2017) (Figure 5), with similar low variance estimates within the spatial confines of the reference sample (Figure 5c,d). Accordingly, the accuracy and precision by which scallops of known origin could be assigned back to origin were also comparable to that demonstrated by Trueman et al. (2017) (Figure 6). However, it must be noted that accuracy and precision results cannot be extrapolated outside the North Sea range and do not reflect wider shelf sea isoscape accuracy and precision. Given that the INLA approach draws on environmental correlates to predict isotopic compositions, one might expect simple kriging to produce more accurate isoscape models where reference sample collection is evenly spaced and dense compared to the spatial scale of isotopic gradients. The similarity in uncertainty between the two methods found here reflects the relatively strong statistical relationships between environmental correlates and reference isotope data. Where evenly spaced reference samples cannot be recovered across the entire region of interest, or where spatial variation in isotope values is expected to occur at smaller spatial scales than the spacing

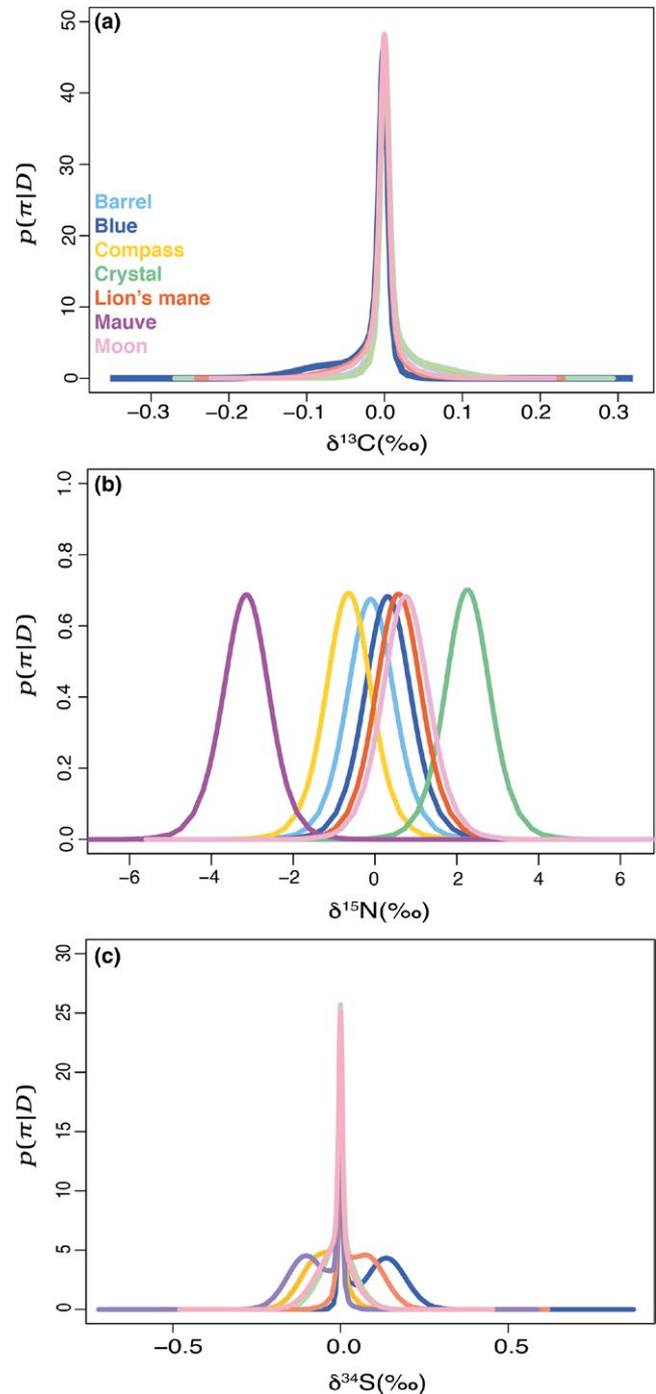


FIGURE 7 Marginal posterior distributions of the species random effect for the chosen carbon, nitrogen and sulphur isotope prediction models. π is the species-level deviation from the overall mean isotope value, and D is the data. Distributions represent the probability density of a given isotopic difference, given the data and represents species differences that remain after the models have been applied. Differences between species represent isotopic differences unable to be explained by environmental variables

between reference samples, isoscapes drawn from environmental predictors may produce more accurate and precise assignments.

One significant benefit of the spatial INLA approach is the ability to account for sources of isotopic variance in the reference data other

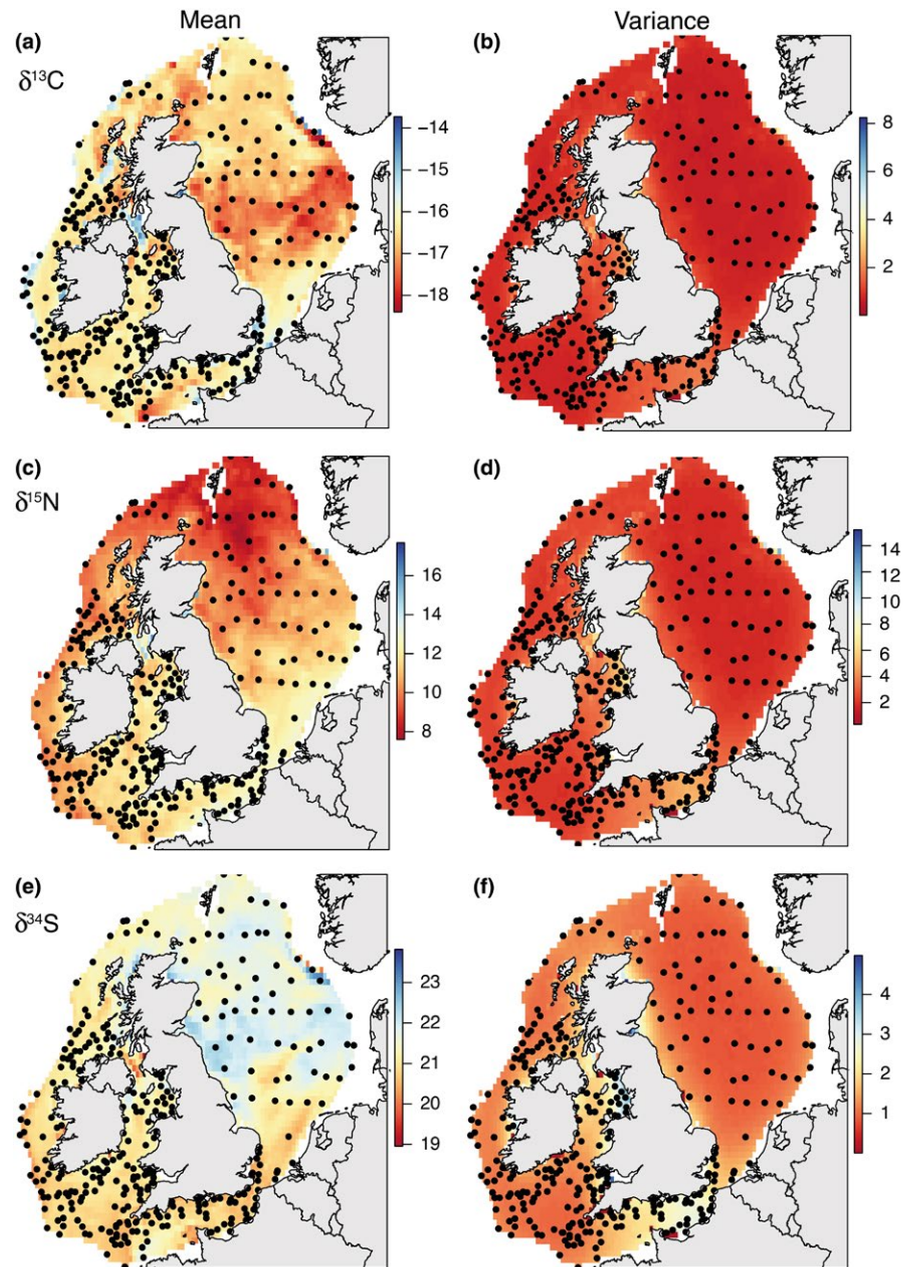


FIGURE 8 UK shelf sea carbon, nitrogen and sulphur isoscape models (a, c, e) and associated variance of the posterior predicted distribution, after species random effects have been accounted for (b, d, f). Values based on seven species of jellyfish (Barrel, Blue, Crystal, Compass, Lion's mane, Mauve and Moon) sampled between August 2015 and December 2016. Filled circles represent sampling locations

than spatially varying terms. In our case, INLA allowed us to identify and account for large, among species isotopic differences, ranging between 0.03–3.0‰ in $\delta^{13}\text{C}$, 0.02–7.1‰ in $\delta^{15}\text{N}$ and 0.01–1.45‰ in $\delta^{34}\text{S}$ (Table 2) into the spatial model. The 'species effects' are quantified as residual differences unaccounted for by the environmental predictors within the final models and displayed as marginal distributions in Figure 7. Both carbon and sulphur 'best-fit' models were able to explain all isotopic differences between species, whereas residual nitrogen isotopic differences were still observed. Mauve stinger and crystal jellyfish had markedly different $\delta^{15}\text{N}$ values with mauve stingers displaying consistently low and crystal jellyfish consistently high $\delta^{15}\text{N}$ values. Isotopic variation among different species is expected, likely due to different diets, habitat uses and metabolic processes. Deciphering the reasons behind these species isotopic differences is beyond the scope of this study, but we emphasize the

importance of treating gelatinous zooplankton as separate species in any isotopic study.

In this example, we use INLA to incorporate isotopic differences between species; however, the same concept applies whenever data with known, or assumed, differences must be combined. For example, in isoscape models where plankton or zooplankton are sampled and grouped (McMahon, Hamady, & Thorrold, 2013; Schell, Barnett, & Vinette, 1998); where data have been collected from multiple sources (Bataille et al., 2018); or where different sampling techniques have been adopted. The same approach could also be used to incorporate temporal variability in sample collection (Bowen & Revenaugh, 2003; Flockhart et al., 2013). While samples in the current study were collected over 2 years, sampling locations did not overlap across different times, so temporal effects could not be explicitly quantified.

4.2 | Isotopic variability across the UK shelf seas

Stratification and mixing extent are strong drivers of spatial isotopic variability, with front locations closely matching isotope ratio boundaries in carbon, nitrogen and sulphur (Miller & Christodoulou, 2014). In shallow well-mixed regions, (e.g. Irish Sea and southern North Sea), isotopically heavy nutrients become resuspended (MacKenzie et al., 2014; Miller & Christodoulou, 2014) (Figure 8), whereas deeper, seasonally stratified regions (e.g. northern North Sea and Celtic Sea), experience nutrient limitation and reduced fractionation (Goericke & Fry, 1994), resulting in higher $\delta^{13}\text{C}$ values (Figure 8).

Isotopic ratios are also strongly influenced by freshwater and terrestrial inputs. Freshwater has a lower $\delta^{34}\text{S}$ ratio compared to seawater (Fry, 2002), reducing $\delta^{34}\text{S}$ values in regions with high freshwater input (e.g. eastern Irish Sea, southern North Sea and English Channel and areas off West Scotland) (Painting et al., 2013) (Figure 8). Anthropogenic nutrient sources enter the marine environment through estuaries (Howarth, 1998) and influence productivity causing increased $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values in coastal and estuarine environments (e.g. southern North Sea, eastern English Channel, eastern Irish Sea) (Painting et al., 2013).

Production source also influences isotopic variability, with phytoplankton community structure differing between the northern and southern North Sea (Ford et al., 2016), the presence of cyanobacteria within the western English Channel (Rees, Gilbert, & Kelly-Gerrey, 2009) decreasing $\delta^{15}\text{N}$ but increasing $\delta^{13}\text{C}$ values due to nitrogen fixation (Levitan et al., 2007) and influence of microalgae increasing $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values around the East Anglian coast and into the southern North Sea (Bristow et al., 2013) (Figure 8). Variance surfaces are similar for each isoscape, with uniform variance across the majority of the UK shelf, but greater values found within more dynamic regions such as the eastern English Channel and eastern Irish Sea.

5 | CONCLUSION

The principle reason for adopting an INLA (or mixed model) approach to generate an isoscape is to account for variance in reference samples that is not explicitly spatial in origin. Where reference datasets can be assembled from the same species, collected at the same time and processed in the same way, simpler spatial modelling or kriging interpolation approaches may be favoured; however, in many cases, some extra non-spatially dependent variance terms are introduced because of the difficulty of obtaining uniform reference samples. In our example, jellyfish species have varying distributions across the spatial range, so to generate a single isoscape model required use of multiple species and therefore the introduction of random effect of species. The INLA approach is a promising method for accounting for additional non-spatially dependent isotopic variance within reference samples. Although our study focuses on marine carbon and nitrogen and the newly introduced sulphur isotopes, the same methods and benefits and limitations are applicable across all environments and isotope systems.

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AUTHORS' CONTRIBUTIONS

K.S.G. and C.T. conceived the project. K.S.G. collected the samples, carried out sample and data analysis and wrote the manuscript. K.S.G. and L.G. designed the statistical analysis. R.M. carried out stable isotope analysis and provided data interpretation expertise. All authors contributed to the manuscript.

DATA ACCESSIBILITY

All data and R code scripts are stored in the Git Hub repository found: <https://github.com/katiestjohnlew/UKShelfIsoscape> (<https://doi.org/10.5281/zenodo.2152638>).

ORCID

Katie St. John Glew  <https://orcid.org/0000-0002-0425-1494>

Laura J. Graham  <https://orcid.org/0000-0002-3611-7281>

Clive N. Trueman  <https://orcid.org/0000-0002-4995-736X>

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