

Spatially Varying Coefficient Models with R-INLA

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Note: See <https://inlabru-org.github.io/inlabru/articles/svc.html> for an updated version of this vignette that avoids manual handling of mesh indices, mapping matrices, and data stacks, by using the `inlabru` interface instead of plain `INLA`.

Introduction

Spatially varying coefficient models (SVCs, Gelfand et al. 2003) are often used to model data when relationships between dependent and independent variables are not uniform across space, a common situation when exploring phenomena across large spatial extents (Finley 2011). Meehan et al. (2019) described an SVC model to evaluate continent-scaled variation in bird abundance trends. The SVC model used in that analysis employed discrete aerial units (100 km grid cells), with spatial structure described by neighborhood matrices and spatial relationships described by an intrinsic conditional autoregressive model (Besag 1974). The online supplement for the paper included code for building the model using the `R-INLA` package (Rue et al. 2009) for the `R` statistical programming language (R Core Team 2021). Both the manuscript and code can be accessed at <https://github.com/tmeeha/inlaSVCBC>.

In this vignette, we will describe how to build an SVC model similar to that described in Meehan et al. (2019), but within a continuous-space framework. This model will be computed using the stochastic partial differential equation (SPDE) approach of Lindgren et al. (2011, 2022), implemented in the `R-INLA` package for `R`. The SPDE approach employs a computationally efficient approximation of a Gaussian random field with parameters directly comparable to those of a Matérn covariance function. The benefits of a continuous-space versus a discrete-space SVC include the potential for finer resolution estimation and prediction, a better understanding of the range of spatial correlation, and a reduction in boundary effects associated with discrete-space analyses.

We build the model using a subset of the data described in Meehan et al. (2019). Specifically, we use counts of American Robin (*Turdus migratorius*) from south central North America collected between 1987 and 2016 during the Audubon Christmas Bird Count (CBC). The overall goal of the analysis is to produce spatially explicit estimates of annual relative abundance as well as long-term relative abundance trends for robins that account for spatial and temporal variation in count effort.

Model

The model used to analyze these data assumes that counts come from a negative binomial distribution with an expected count and dispersion parameter. The expected count has a log-linear predictor:

$$\log(\lambda_{st}) = \kappa_s + \alpha_s + \epsilon_s \log[\text{Effort}_{st}] + \tau_s \text{Year}_{st}$$

where the natural log of expected count, $\log(\lambda_{st})$, at site s during year t , is modeled with a zero-centered, normally distributed intercept per site, κ_s , a spatially varying intercept, α_s , a spatially varying effect of the log of count effort in hours, ϵ_s , and a spatially varying linear effect of year, τ_s . The spatially structured effects are modeled as Gaussian random fields with Matérn covariance functions with range and variance parameters.

Model parameters κ_s , α_s , ϵ_s and τ_s are analogous to those in Meehan et al. (2019). For example, κ_s is included to account for site-level differences in counts, possibly due to habitat availability or observer experience. α_s

can be interpreted as an effort-corrected abundance index at year zero. ϵ_s is the exponent for a power-law effort-correction function. And τ_s is the long-term temporal trend at a given site.

Environment

To get started with data analysis, we set up the environment, by loading some packages, and setting some options.

```
# libraries
library(maps)
library(ggplot2)
library(sf)
library(terra)
library(tidyterra) # raster plotting
library(tidyr)
library(scales)
library(inlabru)
library(INLA)
library(dplyr)
# Note: the 'splancs' package also needs to be installed,
# but doesn't need to be loaded

# set option
select <- dplyr::select
options(scipen = 99999)
options(max.print = 99999)
options(stringsAsFactors = FALSE)
```

Next we define a coordinate reference system (CRS) for spatial analysis and create a base map for later use. The CRS uses the USA Contiguous Albers Equal-Area Conic projection, and is identified by the EPSG code 6703. We modify the CRS slightly to have units of kilometers, so that distances between widespread count sites are not especially large numbers (Krainski et al. 2018).

```
# define a crs
epsg6703km <- paste(
  "+proj=aea +lat_0=23 +lon_0=-96 +lat_1=29.5",
  "+lat_2=45.5 +x_0=0 +y_0=0 +datum=NAD83",
  "+units=km +no_defs"
)

# make a base map
states <- maps::map("state", plot = FALSE, fill = TRUE) %>%
  sf::st_as_sf() %>%
  filter(ID %in% c(
    "texas", "oklahoma", "kansas", "missouri",
    "arkansas", "louisiana"
  )) %>%
  sf::st_make_valid() %>%
  sf::st_transform(epsg6703km) %>%
  sf::st_make_valid()
```

Import data

Next we import some bird count data from the GitHub repository associated with Meehan et al. (2019), and turn the data set into spatially referenced points. We use a subset of the data (30 years, 6 US states) for this

analysis to reduce computing time (~ 1 min). Note that site selection and zero filling, important components of trend analyses, have already been conducted and this is the resulting data set.

```
# get data
## download_dat <- read.csv(paste0(
##     "https://raw.githubusercontent.com/tmeeha/inlaSVCBC",
##     "/master/code/modeling_data.csv"))
download_dat <- read.csv("data/modeling_data.csv.gz")

# select subset
count_dat <- download_dat %>%
  select(
    circle, bcr, state, year, std_yr, count, log_hrs,
    lon, lat, obs
  ) %>%
  mutate(year = year + 1899) %>%
  filter(
    state %in% c(
      "TEXAS", "OKLAHOMA", "KANSAS", "MISSOURI",
      "ARKANSAS", "LOUISIANA"
    ),
    year >= 1987
  )
```

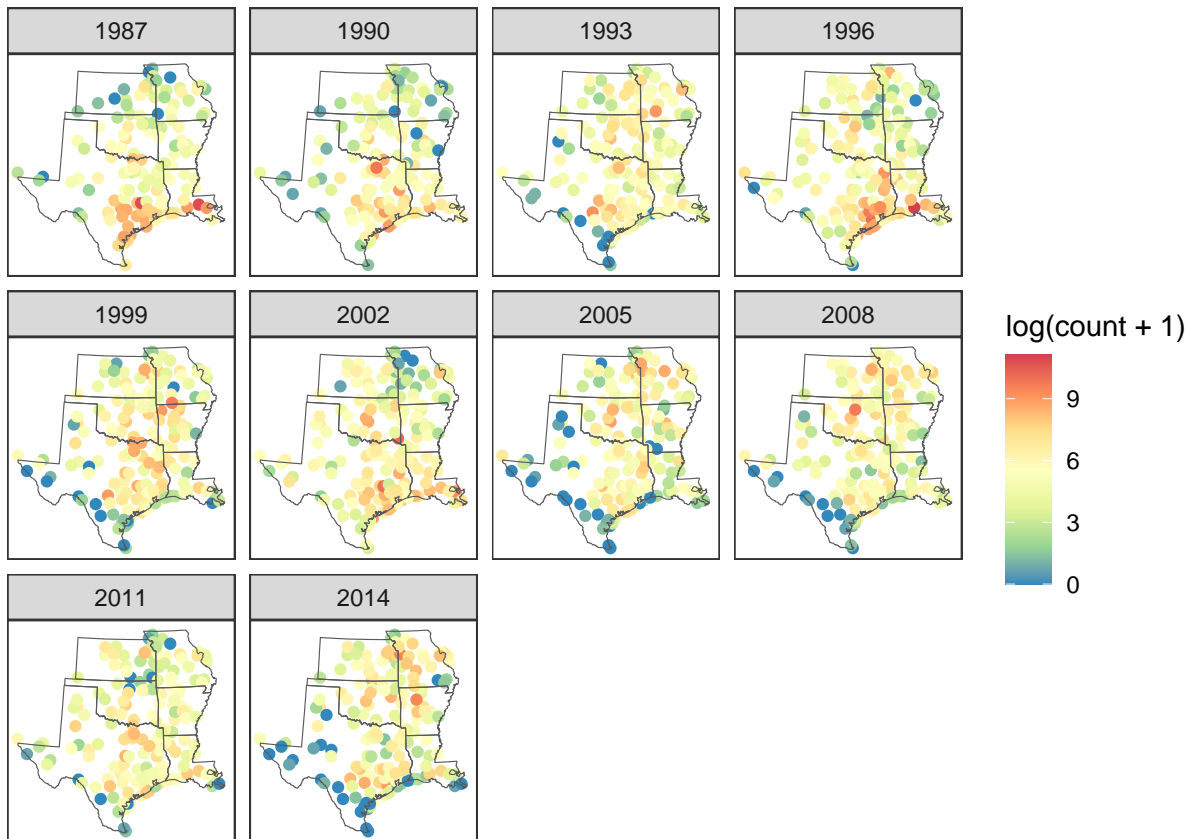
After loading the data, we filter out observation sites with less than 20 years of data, add index variables to uniquely index site and year information, and transform the coordinates to the `epsg6703km` CRS:

```
count_dat <- count_dat %>%
  mutate(site_idx = as.numeric(factor(paste(circle, lon, lat)))) %>%
  group_by(site_idx) %>%
  mutate(n_years = n()) %>%
  filter(n_years >= 20) %>%
  ungroup() %>%
  mutate(
    std_yr = year - max(year),
    obs = seq_len(nrow(.)),
    site_idx = as.numeric(factor(paste(circle, lon, lat))),
    year_idx = as.numeric(factor(year)),
    site_year_idx = as.numeric(factor(paste(circle, lon, lat, year)))
  ) %>%
  st_as_sf(coords = c("lon", "lat"), crs = 4326, remove = FALSE) %>%
  st_transform(epsg6703km) %>%
  mutate(
    easting = st_coordinates(.)[, 1],
    northing = st_coordinates(.)[, 2]
  ) %>%
  arrange(circle, year)
```

A rough view of changes in robin relative abundance, which does not account for variation in count effort, can be seen by plotting raw counts per site and year.

```
# map it
ggplot() +
  geom_sf(
    data = count_dat %>% filter(year_idx %in% seq(1, 30, 3)),
    aes(col = log(count + 1))
  )
```

```
) +
geom_sf(data = states, fill = NA) +
coord_sf(datum = NA) +
facet_wrap(~year) +
scale_color_distiller(palette = "Spectral") +
theme_bw()
```



Make spatial data

Next we use the count data to make a map of distinct count sites and save the coordinates of the sites, unique and across all years, for later spatial modeling and plotting.

```
# make a set of distinct study sites for mapping
site_map <- count_dat %>%
  select(circle, easting, northing) %>%
  distinct() %>%
  select(circle, easting, northing)

# save coordinates as matrices for later
unique_coords <- count_dat %>%
  select(circle, easting, northing) %>%
  distinct() %>%
  select(easting, northing) %>%
  st_drop_geometry() %>%
  as.matrix()
all_coords <- as.matrix(st_coordinates(count_dat))
```

SPDE components

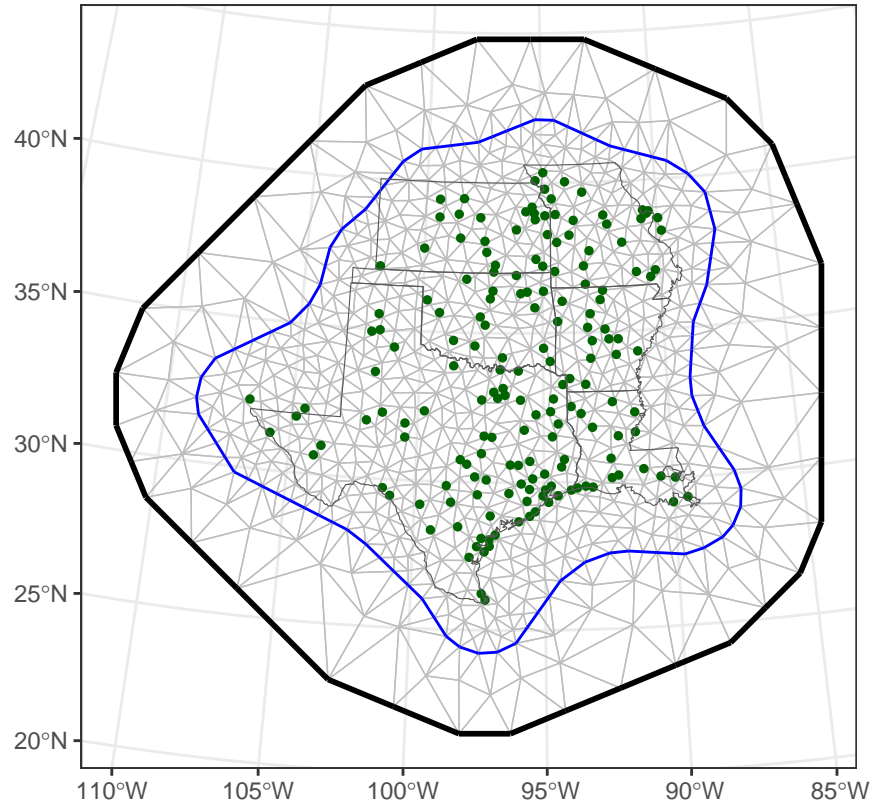
Computing a continuous-space model with R-INLA using the SPDE approach requires construction of four distinct sets of data and model objects (Blangiardo and Cameletti 2015, Krainski et al. 2018). First, we create a **modeling mesh**, which is used to provide a piecewise linear representation of the continuous spatial surface, based on a triangulation of the modeled region. Here, the same mesh will get reused for each of the spatial and spatio-temporal terms in the model. Second, we construct an **SPDE model object** that specifies properties of the spatial model. Again, we will use the same SPDE object for each of the spatial terms in the model. Third, we create **named index sets** that identify data locations and mesh nodes for use in model construction, computation, and output extraction. Note that a separate set of indices needs to be constructed for each spatial term in the model. Fourth, we construct **projector matrices** or **A matrices** that create formulaic mappings between the observed data locations and the mesh nodes. Again, note that a separate set of projector matrices needs to be constructed for each spatial term in the model. The four corresponding functions for these steps are: `fmesher::fm_mesh_2d()`, `inla.spde2.pcmatern()`, `inla.spde.make.index()`, and `inla.spde.make.A()`.

Modeling mesh

There are various things to consider when constructing a mesh (Lindgren and Rue 2015, Blangiardo and Cameletti 2015, Krainski et al. 2018, Bakka et al. 2018). In constructing one, we balance a trade-off between capturing fine-scaled features of the Gaussian random field and computing times. Here, we create a non-convex hull around the count sites, and then build a triangular mesh by specifying minimum and maximum edge lengths within and slightly outside the hull.

```
# make a hull and mesh for spatial model
hull <- fm_nonconvex_hull_inla(
  count_dat,
  convex = 200,
  concave = 350
)
mesh <- fm_mesh_2d(
  boundary = hull, max.edge = c(100, 600), # km inside and outside
  cutoff = 50, offset = c(100, 300),
  crs = fm_crs(count_dat)
) # cutoff is min edge

# plot it
ggplot() +
  gg(data = mesh) +
  geom_sf(data = site_map, col = "darkgreen", size = 1) +
  geom_sf(data = states, fill = NA) +
  theme_bw() +
  labs(x = "", y = "")
```



SPDE model object

Next we create an SPDE object to define the model smoothness, with prior distributions for the variance and range parameters, and the mesh. Here we assume a Gaussian random field characterized with a Matérn covariance function with penalized complexity priors (Simpson et al. 2017) for the practical range (distance where spatial correlation approaches 0.1) and variation explained by the function (Fuglstad et al. 2019). The prior for the spatial range is set such that the probability of a range exceeding 500 km is 0.5. The prior for the variance explained by the spatial effect is set such that the probability of a standard deviation exceeding 1 is 0.5 (Krainski et al. 2018). If one wants to constrain this kind of spatial effect to integrate to zero, `constr=TRUE` should be added at this stage.

```
# make spde
spde <- inla.spde2.pcmatern(
  mesh = mesh,
  prior.range = c(500, 0.5),
  prior.sigma = c(1, 0.5)
)
```

Index sets

Next we create index sets for the spatial terms in the model. Note that the `tag` names applied to the index sets in this step will be used in the model formula statement below. The index vectors for the spatial terms are essentially vectors of integers going from 1 to the number of nodes on the mesh.

```
# make index sets
alpha_idx <- inla.spde.make.index(name = "alpha", n.spde = mesh$n)
eps_idx <- inla.spde.make.index(name = "eps", n.spde = mesh$n)
```

```
tau_idx <- inla.spde.make.index(name = "tau", n.spde = mesh$n)
```

Projector matrices

Next we create projector matrices for the spatial terms in the model. Note that this is the step where SVCs are specified. In our model, ϵ_s is a spatially varying effect of the log of count effort. The SVC effort effect is specified using the `weights` argument to `inla.spde.make.A()` function. Similarly, τ_s is a spatially varying effect of year, so standardized year (with $1987 = 0$) is also specified in the `weights` argument. α_s is also an SVC, but it is a spatially varying intercept. For intercepts it is not necessary to specify a constant weight of 1 when making a projector matrix. The only model component ignored here is κ_s , because it is not spatially structured and not modeled on the mesh. Note that the current use of the term ‘weights’ is different from that often encountered when defining mixed effect models in R. Here it is used to define covariate values, as opposed to importance values in other contexts.

```
# make projector matrices
A_alpha <- inla.spde.make.A(mesh = mesh, loc = count_dat)
A_eps <- inla.spde.make.A(
  mesh = mesh, loc = count_dat,
  weights = count_dat$log_hrs
) # note weights argument
A_tau <- inla.spde.make.A(
  mesh = mesh, loc = count_dat,
  weights = count_dat$std_yr
) # note weights argument
```

Data stack for model fitting

With the SPDE method, there are several data structures that need to be bundled and used for analysis of the spatial model. There is the observed response at the count sites, the covariate values at the count sites, and the indices and mapping of those data to the modeling mesh. All of this information gets bundled into a data stack for model analysis using the `inla.stack()` function. This function is a very flexible and can be difficult to understand at first. See Blandgiardo and Cameletti (2015) and Gomez-Rubio (2020) for practical information on how to use it.

The first argument to the function, shown below, is the `tag` name for the stack. In this case we are including observed data for model estimation, so we name the stack ‘obs’. The next argument, `data`, specifies the response variable. Here we have a single response variable called ‘count’. We tell the stacking function that observed counts reside in the ‘count_dat’ data set. The next argument, `effects`, is used to specify the model predictors. These are defined using the observed data for non-spatial predictors, and using the named index sets for the spatial predictors. The last argument, `A`, asks for a list of projector matrices or A matrices for the predictor variables. We did not need to make a projector for κ_s , because it is not defined as a spatial parameter. Nevertheless, we add a value of 1 because it is the value that will be multiplied by the effect during model computations and serves as a way for the stacking function to match the number of A matrices with the number of effects – if there are 4 effects, then there must be four entries for A. Note that if there were more than one non-spatial predictor in the model, whether a grouped random intercept or a global fixed effect, then each one would be given its own value of 1 during this step. And note that the order of the A matrices must match the order of the effects.

```
# stack observed data
stack_fit <- inla.stack(
  tag = "obs",
  data = list(count = as.vector(count_dat$count)), # response from data frame
  effects = list(data.frame(
    intercept = 1,
    kappa = count_dat$site_idx
```

```

), # predictors from data frame
alpha = alpha_idx, # or index sets if spatial
eps = eps_idx,
tau = tau_idx
),
A = list(
  1, # a value of 1 is given for non-spatial terms
  A_alpha,
  A_eps,
  A_tau
)
)

```

Model formula

The last required input for the analysis is the model formula, which includes information on the prior for explained variation for the unstructured random intercept. We define the prior for κ_s as a penalized complexity prior (Simpson et al. 2017), set such that the probability of the standard deviation associated with the random effect exceeding 1 is 0.01.

```

# iid prior
pc_prec <- list(prior = "pcprec", param = c(1, 0.1))

```

Notice that if one wants to constrain a spatial spde effect to integrate to zero, it should be added `constr=TRUE` in the SPDE model definition rather than in the `f()` terms. As we want to constrain κ_s and it is a non-spatial term we can use `constr=TRUE` in its corresponding `f()` below, which imposes a sum-to-zero constraint.

The model described above is translated to R-INLA modeling syntax as:

```

# formula
svc_form <- count ~ -1 +
  f(kappa, model = "iid", constr = TRUE, hyper = list(prec = pc_prec)) +
  f(alpha, model = spde) +
  f(eps, model = spde) +
  f(tau, model = spde)

```

Here, we define the response as `count`, remove the automatic global intercept with a `-1`, and then specify the other terms in the model with `f()` statements. The first `f()` statement defines κ_s , the site effect, as a normally distributed (`model="iid"`), zero-centered (`constr=TRUE`), deviation from α_s . The name ‘kappa’ given in the `f()` argument is associated with the effect ‘kappa’ defined in the estimation stack. The second `f()` statement defines α_s as a spatially varying intercept with spatial structure described by the SPDE object called ‘spde’. The third `f()` statement defines ϵ_s as an SVC for the effect of count effort, with spatial structure also described by the SPDE object. The weights for this spatially structured random slope were specified during construction of the projector A matrix, `A_eps`. The fourth `f()` statement defines τ_s as an SVC for the year effect, with spatial structure described in the SPDE object. Again, the weights for this spatially structured random slope were specified during construction of the projector A matrix, `A_tau`. Again, note that the current use of the term ‘weights’ is different from that often encountered when defining mixed effect models in R. Here it is used to define covariate values, as opposed to importance values in other contexts.

Run model

We estimate the model with a call to `inla()`. First we set the option to use the (new) experimental way to do internal computations, see Van Niekerk et. al. (2022), for the sake of computing speed and better numerics. In the call to `inla()`, we give the model formula, specify the negative binomial distribution for the counts, define the estimation data, and describe the location of the projector A matrix. Note that we are not generating predictions (`compute=F`). Nevertheless, the `control.predictor` argument still needs to be

included so the software knows how to find the A matrices. Then we ask `inla()` to compute WAIC and CPO to evaluate model fit, and to save the information necessary for posterior sampling (`config=T`). For computing speed, we choose to use the adaptive integration strategy and Empirical Bayes estimation. The `inla()` run for this model takes about 1 minutes on a standard laptop computer. Another option is to use the Variational Bayes approximation as detailed in Van Niekerk and Rue (2021) and Van Niekerk et. al. (2022).

```
res <- inla(svc_form,
  family = "nbinomial",
  data = inla.stack.data(stack_fit), # stack the stack
  control.predictor = list(
    A = inla.stack.A(stack_fit),
    compute = FALSE
  ), # must define A
  control.compute = list(waic = TRUE, cpo = TRUE, config = TRUE),
  control.inla = list(strategy = "adaptive", int.strategy = "eb"),
  verbose = FALSE
)
```

Model summaries

Once computation is complete, we look at the initial results to see how things went. First we check the posterior means for the hyperparameters of the model, mainly the variance components and the spatial ranges of the spatially structured parameters.

```
# view results
res$summary.hyperpar[-1, c(1, 2)]
```

```
##              mean      sd
## Precision for kappa    2.1524  0.4018
## Range for alpha      1030.9565 335.5792
## Stdev for alpha       2.0924  0.4800
## Range for eps        5544.8429 3679.2972
## Stdev for eps         0.4443  0.1953
## Range for tau        760.3879 274.5707
## Stdev for tau         0.0634  0.0127
```

Next we examine some summaries of the random effect estimates, starting with $\exp(\alpha_s)$, which is effort-corrected relative abundance at year = 0 (1987), given 1 hour of count effort (i.e., $\log[1]=0$).

```
summary(exp(res$summary.random$alp$"0.5quant")) # exp(alpha) posterior median
```

```
##      Min. 1st Qu.  Median    Mean 3rd Qu.    Max.
## 0.0341  1.0981  3.3320  6.9666  9.1488 56.1732
```

Note that to avoid issues due to $E[\exp(x)|y] \neq \exp[E(x|y)]$, we use the posterior median instead of the posterior mean.

The summary for ϵ_s shows variation in the exponent for the effort correction function across space.

```
summary(res$summary.random$eps$mean) # epsilon
```

```
##      Min. 1st Qu.  Median    Mean 3rd Qu.    Max.
## 0.732  0.879  0.986  0.977  1.051  1.270
```

The τ_s summary shows how long-term, log-linear trends of robin relative abundance have varied across space, from annual decreases of around 10% to annual increases of around 10%.

```
summary((exp(res$summary.random$tau$"0.5quant") - 1) * 100) # (exp(tau)-1)*100
```

```
##      Min. 1st Qu.  Median      Mean 3rd Qu.      Max.
## -13.06   -5.10   -1.51   -1.04    2.87   11.09
```

SVC maps

Next we create maps of α_s , ϵ_s , and τ_s to inspect the spatial structure of these parameter estimates. We start by creating a 25-km mapping grid, and then projecting this mapping grid to the modeling mesh.

```
# get easting and northing limits
bbox <- fm_bbox(hull[[1]])
grd_dims <- round(c(x = diff(bbox[[1]]), y = diff(bbox[[2]])) / 25)

# make mesh projector to get model summaries from the mesh to the mapping grid
mesh_proj <- fm_evaluator(
  mesh,
  xlim = bbox[[1]], ylim = bbox[[2]], dims = grd_dims
)
```

Then we populate the mapping grids with parameter estimates (posterior median and range95), turn them into a raster stack, and mask the raster stack to the study area.

```
# pull data
kappa <- data.frame(
  median = exp(res$summary.random$kappa$"0.5quant"),
  range95 = exp(res$summary.random$kappa$"0.975quant") -
    exp(res$summary.random$kappa$"0.025quant")
)
alph <- data.frame(
  median = exp(res$summary.random$alpha$"0.5quant"),
  range95 = exp(res$summary.random$alpha$"0.975quant") -
    exp(res$summary.random$alpha$"0.025quant")
)
epsi <- data.frame(
  median = res$summary.random$eps$"0.5quant",
  range95 = (res$summary.random$eps$"0.975quant" -
    res$summary.random$eps$"0.025quant")
)
taus <- data.frame(
  median = (exp(res$summary.random$tau$"0.5quant") - 1) * 100,
  range95 = (exp(res$summary.random$tau$"0.975quant") -
    exp(res$summary.random$tau$"0.025quant")) * 100
)

# loop to get estimates on a mapping grid
pred_grids <- lapply(
  list(alpha = alph, epsilon = epsi, tau = taus),
  function(x) as.matrix(fm_evaluate(mesh_proj, x))
)

# make a terra raster stack with the posterior median and range95
out_stk <- rast()
for (j in 1:3) {
  mean_j <- cbind(expand.grid(x = mesh_proj$x, y = mesh_proj$y),
    Z = c(matrix(pred_grids[[j]][, 1], grd_dims[1]))
  )
  mean_j <- rast(mean_j, crs = epsg6703km)
```

```

range95_j <- cbind(expand.grid(X = mesh_proj$x, Y = mesh_proj$y),
  Z = c(matrix(pred_grids[[j]][, 2], grd_dims[1]))
)
range95_j <- rast(range95_j, crs = epsg6703km)
out_j <- c(mean_j, range95_j)
terra::add(out_stk) <- out_j
}
names(out_stk) <- c(
  "alpha_median", "alpha_range95", "epsilon_median",
  "epsilon_range95", "tau_median", "tau_range95"
)
out_stk <- terra::mask(
  out_stk,
  terra::vect(sf::st_union(states)),
  updatevalue = NA,
  touches = FALSE
)

```

Finally, we plot the SVCs with the following code. We plot the posterior median and 95% uncertainty width (“range95”) for $\exp(\kappa_s)$, $\exp(\alpha_s)$, ϵ_s , and $100(\exp(\tau_s) - 1)$.

```

make_plot_field <- function(data_stk, scale_label) {
  ggplot(states) +
    geom_sf(fill = NA) +
    coord_sf(datum = NA) +
    geom_spatraster(data = data_stk) +
    labs(x = "", y = "") +
    scale_fill_distiller(
      scale_label,
      palette = "Spectral",
      na.value = "transparent"
    ) +
    theme_bw() +
    geom_sf(fill = NA)
}

make_plot_site <- function(data, scale_label) {
  ggplot(states) +
    geom_sf() +
    coord_sf(datum = NA) +
    geom_sf(data = data, size = 1, mapping = aes(colour = value)) +
    scale_colour_distiller(scale_label, palette = "Spectral") +
    labs(x = "", y = "") +
    theme_bw() +
    geom_sf(fill = NA)
}

# medians
# fields alpha_s, epsilon_s, tau_s
pa <- make_plot_field(
  data_stk = out_stk[["alpha_median"]],
  scale_label = "posterior\nmedian\nexp(alpha_s)"
)
pe <- make_plot_field(
  data_stk = out_stk[["epsilon_median"]],

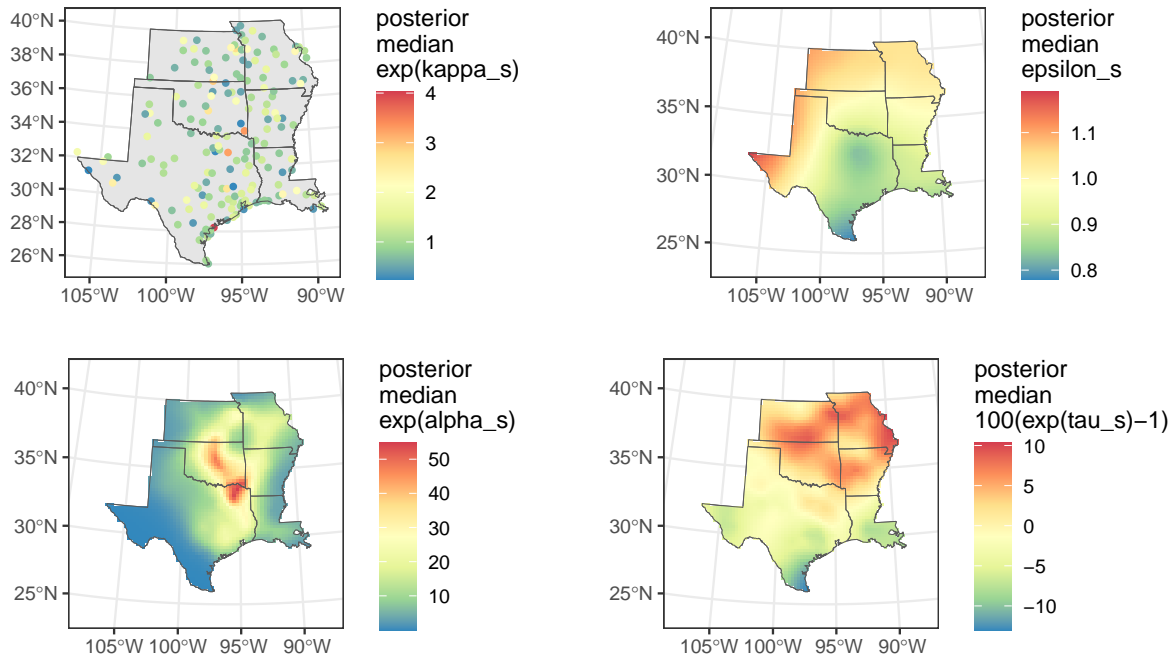
```

```

    scale_label = "posterior\nmedian\nepsilon_s"
  )
  pt <- make_plot_field(
    data_stk = out_stk[["tau_median"]],
    scale_label = "posterior\nmedian\n100(exp(tau_s)-1)"
  )
  # sites kappa_s
  ps <- make_plot_site(
    data = cbind(site_map, data.frame(value = kappa$median)),
    scale_label = "posterior\nmedian\nexp(kappa_s)"
  )
  # range95
  # fields alpha_s, epsilon_s, tau_s
  pa_range95 <- make_plot_field(
    data_stk = out_stk[["alpha_range95"]],
    scale_label = "posterior\nrange95\nexp(alpha_s)"
  )
  pe_range95 <- make_plot_field(
    data_stk = out_stk[["epsilon_range95"]],
    scale_label = "posterior\nrange95\nepsilon_s"
  )
  pt_range95 <- make_plot_field(
    data_stk = out_stk[["tau_range95"]],
    scale_label = "posterior\nrange95\n100(exp(tau_s)-1)"
  )
  # sites kappa_s
  ps_range95 <- make_plot_site(
    data = cbind(site_map, data.frame(value = kappa$range95)),
    scale_label = "posterior\nrange95\nexp(kappa_s)"
  )

  # plot together
  multiplot(ps, pa, pe, pt, cols = 2)

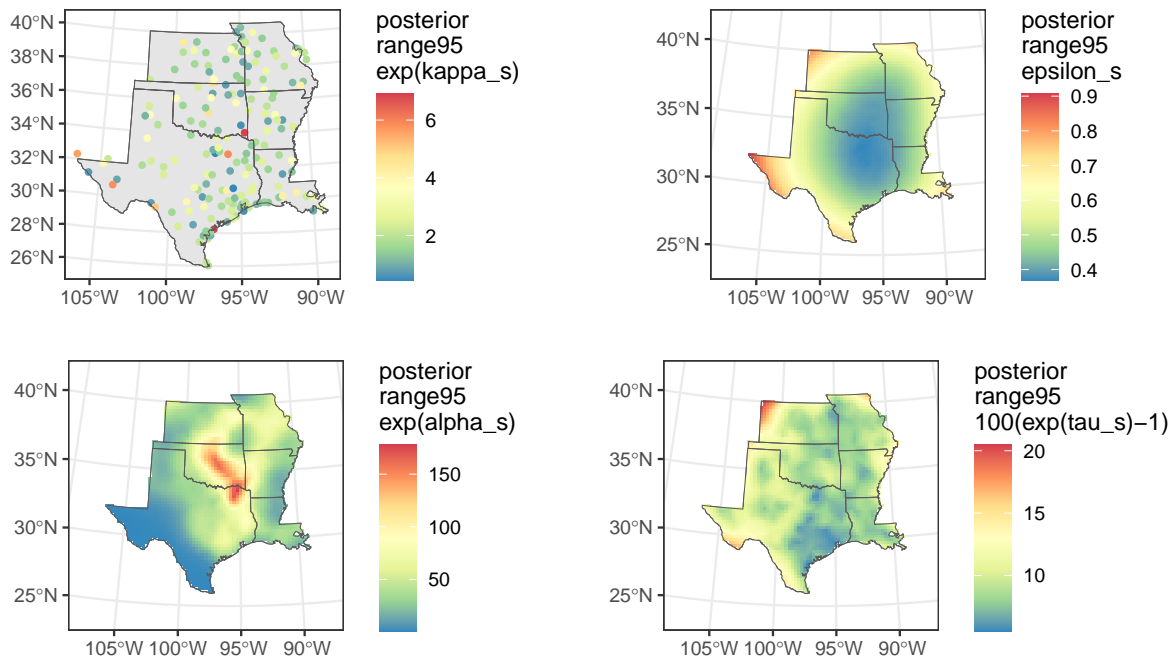
```



The map for the posterior mean of τ_s shows that robins have decreased in the southern part of the study area and increased in the northern part. This demonstrates how the wintering range of robins is shifting northward as winters become warmer due to climate change.

plot together

`multiplot(ps_range95, pa_range95, pe_range95, pt_range95, cols = 2)`



More information

More information on building spatial models using the SPDE approach in R-INLA can be found in Lindgren and Rue (2015), Blangiardo and Caimaletti (2015), Bakka et al. (2018), Krainski et al. (2018), and Moraga (2019).

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