# Hierarchical Generalized Additive Models: an introduction with mgcv

#### ABSTRACT

This is just placeholder text until we write a proper abstract

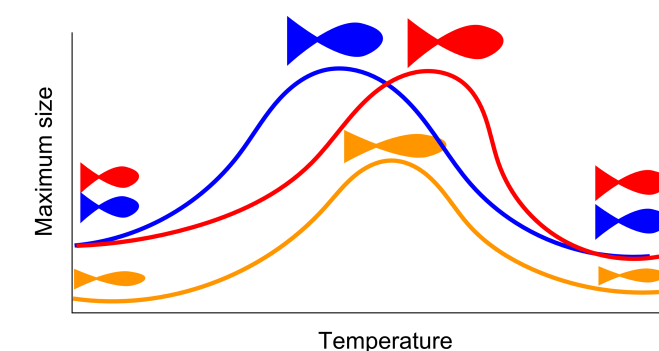
#### 5 I: INTRODUCTION

As ecology has progressed as a quantitative discipline and the questions ecologists ask have become more complicated, the statistical techniques ecologists use have increased in their flexibility to model complex relationships. Two of the more popular and powerful techniques now in use are generalized additive models (GAMs; Simon N Wood 2006) for modelling flexible regression functions, and generalized linear mixed models ("hierarchical generalized linear models" HGLMs or simply "hierarchical models"; Bolker et al. 2009, A. Gelman et al. (2013)) for modelling between-group variability in regression relationships.

At first glance, GAMs and HGLMs are very different tools. GAMs are used to estimate smooth functional relationships between predictor variables and the response, assuming that the phenomena under investigation is not linear (in the GLM sense) but is a smooth function of the predictor variables. Example of such relationships would be the vertical distribution of abundance of a population as a function of depth (Stanley, Pedersen, and Snelgrove 2016) or the swimming speed of snakes as function of temperature (Vickers, Aubret, and Coulon 2017). HGLMs, on the other hand, are used to estimate linear relationships between predictor variables and response, but impose a structure where predictors are organized into groups (often referred to as "blocks") and the relationships between predictor and response may differ between those groups. Either or both slope and intercept may be subject to grouping. A typical example of HGLM use might be to include site-specific effects in a model of counts, or to model individual level heterogeneity in a study with repeated observations of multiple individuals.

Both GAMs and HGLMs can be used to fit potentially highly variable models by "pooling" parameter estimates towards one another. The connection between the two methods is quite deep and a GAMs may be interpreted (and fitted) as HGLMs and vice-versa. Given this connection, the obvious extension to the standard GAM framework is to allow the smooth functional relationship between predictor and response to vary between different grouping levels, but in such a way that the different functions are in some sense pooled toward each other. We often want to know both how the functional relationship between varies between groups, and if there is a strong relationship on average across groups. We will refer to this type of model as a hierarchical GAM, or HGAM.

There are many potential uses for HGAMs. For example, to estimate how the maximum size different fish species reach varies along a common temperature gradient (figure 1). Each species will typically have its own response function, but since the species overlap in range, they should have similar responses over at least some of the temperature gradient; figure 1 shows all three species reach their largest maximum sizes in the center of the temperature gradient. Estimating a seperate function for each species throws away a lot of shared information and could result in highly noisy function estimates if there were only a few data points for each species. Estimating a single average relationship could result in an average function that did not predict any specific group well. In our example, using a single global temperature-size relationship would miss the three species distinct temperature optima, and that the orange species is significantly smaller at



**Figure 1.** Hypothetical example of functional variability between different group levels. Each line indicates how the maximum possible body size for different species of fish in a community might vary as a function of average water temperature. While the orange species shows lower maximum size at all temperatures, and the red and blue species differ in what temperature they can acheive the maximum possible size, all three curves are similarly smooth, and peak close to one another, relative to the entire range of tested temperatures.

- all temperatures than the other two (figure 1). We prefer a hierarchical model that fit a single global temperature-size curve plus species-specific curves that were penalized to be close to the mean function.
- The capability to fit HGAMs already exists in the popular mgcv package for the R statistical programming language. There are many different options available respresenting different model assumptions with corresponding trade-offs. This paper will cover the different approaches to group-level smoothing, the options for each one and why a user might choose it, and demonstrate the different approaches across a range of case studies.
- This paper is divided into six sections. Part II is a brief (and friendly) review of how generalized additive models work and their relation to hierarchical models. In part III, we discuss different ways of modelling hierarchical generalized additive models, what assumptions each model makes about how information is shared between groups, and different ways of specifying these models in mgcv. In part IV, we discuss some of the tools available for plotting model output and assessing model goodness of fit. In part V, we discuss some of the computational and statistical issues involved in fitting HGAMs in mgcv. Finally, in part VI, we work through a few examples of analyses using this approach, to demonstrate the modelling process and how hierarchical GAMs can be incorporated into the quantitative ecologist's toolbox.

#### II: AND INTRODUCTION TO GENERALIZED ADDITIVE MODELS

One of the most common model formulations in statistics is the generalized linear model (McCullagh and Nelder 1989) — models that relate some response (y) to linear combinations of

explanatory variables. We may allow allow the response to be distributed according to some exponential family distribution (e.g., letting the response be a trial, a count or a strictly positive number – binomial, Poisson or Gamma distributions, respectively). The generalized additive modelling (GAM) framework (Hastie and Tibshirani 1990; Ruppert, Wand, and Carroll 2003; Simon N Wood 2006) allows the relationships between the explanatory variables (henceforth covariates) and the response to be described by smooth terms (usually *splines* (Boor 1978), but potentially other structures). In general we are then talking about models of the form:

$$\mathbb{E}(y) = g^{-1} \left( \beta_0 + \sum_{j=1}^J f_j(x_j) \right),\,$$

where y is the response (with an appropriate distribution and link function g),  $f_j$  is a smooth function of the covariate  $x_j$ ,  $\beta_0$  is an intercept term and  $g^{-1}$  is the inverse link function. Here there are J smooths and each is a function of only one covariate, though it is possible to construct smooths of multiple variables.

Each smooth  $f_j$ s is represented by a sum of simpler basis functions  $(b_k)$  multiplied by corresponding coefficients  $(\beta_k)$ , which need to be estimated to be estimated:

$$f_j(x_j) = \sum_{k=1}^K \beta_k b_k(x_j),$$

The size of K of each smooth will determine the flexibility of the resulting term (referred to as "basis size", "basis complexity" or "basis richness"). Though it seems like the basis can be overly complex ("how big should I make K?") and lead to overfitting, we need not worry about this as we use a penalty to ensure that the functions complexity is appropriate; hence the basis only need to be "large enough" and we let the penalty deal with excess wigglyness.

The penalty for a term is usually based on derivatives of that term — as the derivatives give the wigglyness of the function and hence its flexibility. We trade-off the fit of the model against the wigglyness penalty to obtain a model that both fits the data well but does not overfit. To control this trade-off we estimate a *smoothing parameter*. Figure 2 shows optimal smoothing (where the smoothing parameter is estimated to give a parsimonious model) in the first plot; the second plot shows what happens when the smoothing parameter is set to zero, so the penalty has no effect (interpolation); the right plot shows when the smoothing parameter is set to a very large value, giving a straight line. Smooths of this kind are often referred to as a *basis-penalty smoothers*.

#### say something about knots!

#### say something about penalty matrices and also smoothing parameters

The number of basis functions, K, limits the maximum basis complexity for a given smooth term. To measure the wigglyness of a given term, we use the *effective degrees of freedom* (EDF) which, at a maximum is the number of coefficients to be estimated in the model, minus any constraints. The EDF can take non-integer values and a larger value indicates a more wiggly term. See Simon N Wood (2006) Section 4.4 for further details.

There are many possible basis functions that can be used to model the  $b_k$ s. Here we'll use thin plate regression splines, which have the appealing property that knot choice is somewhat automatic (the best approximation to including knots at each data point is used; Simon N. Wood (2003)).

#### DLM:: put the cubic splines back in here

TPRS are also defined for any number of predictors, so multivariate smoothers can be constructed easily. The basis is *isotropic* so smoothing is treated the same in all directions. So if one had, a bivariate smooth of temperature and time, a one degree change in temperature would equate to

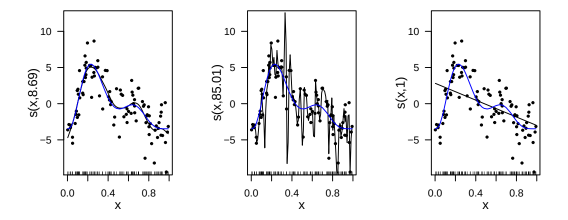


Figure 2. Examples of how different choices of the smoothing parameter effect the resulting function. Data (points) were generated from the blue function and noise added to them. In the left plot the smoothing parameter was estimated to give a good fit to the data, in the middle plot the smoothing parameter was set to zero, so the penalty has no effect and the function interpolates the data, the right plot shows when the smoothing parameter is set to a very large value, so the penalty removes all terms that have any wigglyness, giving a straight line. Numbers in the y axis labels show the estimated degrees of freedom for the term.

a one second change in time, which is an odd assumption to make. In the more general case where units are not alike, we can use *tensor products* to combine two or more univariate smooths into a more complex basis. Each component can be made up from a different basis, playing to their particular strengths.

In the linear modelling literature we can specify a single interaction between terms (in R, a:b) or a "full interaction", which includes the marginal terms (a\*b in R, which is equivalent to a + b + a:b). There are parallels for smooths too, allowing us to separate-out the main effect terms from the interactions (in R te(a, b) specifies the tensor product which is equivalent to ti(a) + ti(b) + ti(a, b)). The ability to separate out the interactions and main effects will become very useful in the next section, once we start looking at group-level smooths. For an example of a tensor product interaction, see figure 3, which illustrates marginal smooths and a tensor product interaction of depth and salinity predicting mackerel egg counts from a marine survey.

We represent the terms in our model as basis functions, which end up as additional columns in our design matrix and parameter vector, and penalties, which penalize the likelihood and stop our model from being too wiggly. Taking a pragmatic Bayesian approach to the problem, the penalty is really a prior on how we think the model should act. In which case the penalty matrix itself is a prior precision matrix (inverse variance) for the term. With that in mind, we can think about random effects as "smooths" in our model, albeit ones with ridge penalies (Kimeldorf and Wahba 1970; Simon N. Wood 2017). For instance, to include a random effect modelling between group variation in intercepts there will be one basis function for each level of the grouping variable, that takes a value of 1 for any observation in that group and 0 for any observation not in the group. The penalty matrix for these terms is a  $n_g$  by  $n_g$  identity matrix, where  $n_g$  is the number of groups. This means that each group-level coefficient will be penalized in proportion to its squared deviation from zero. This is equivilent to how random effects are estimated in standard mixed effect models. The penalty term here is proportionate to the inverse of the variance of the fixed effect estimated by standard hierarchical model solvers (Verbyla et al. 1999 does this contradict what's above??).

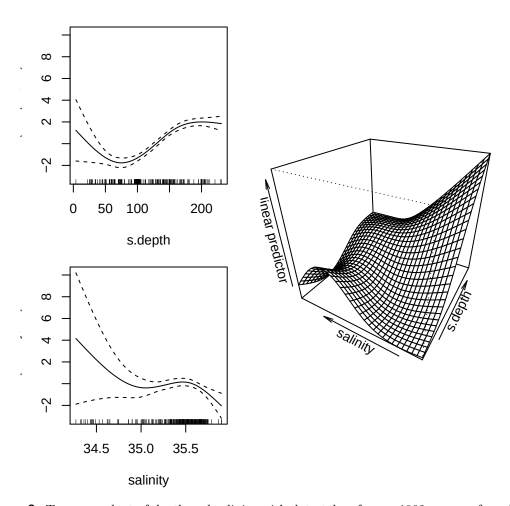


Figure 3. Tensor product of depth and salinity with data taken from a 1992 survey of mackerel eggs. The two left plots show the marginal smooths of each term in the model (ti(s.depth) above and ti(salinity) below), the right plot shows the interaction effect (ti(s.depth, salinity)). Data are from Simon N Wood (2006).

This connection between random effects and basis function smooths extends beyond the varyingintercept case. Any basis-function representation of a smooth function can be transformed so 133 that it can be represented as a combination of a random effect with an associated variance, and 134 possibly one or more fixed effects, corresponding to functions in the null space of the original 135 136 basis-function (see below). While this is beyond the scope of this paper, see Verbyla et al. (1999) or Simon N. Wood, Scheipl, and Faraway (2013) for a more detailed discussion on the connections 137 between these approaches. 138

#### Smoothing penalties vs. shrinkage penalties

does this go above??

#### EJP: I think this makes sense here

Penalties can have two effects on how well a model fits: they can penalize how wiggly a given term is (smoothing) and they can penalize the absolute size of the function (shrinkage). The penalty can only effect the components of the smooth that have derivatives (the range space), not the other parts (the nullspace). For 1-dimensional thin plate regression splines, this means 145 that there is a linear term left in the model, even when the penalty is in full force (as  $\lambda \to \infty$ ), as shown in figure 4 **BLAH**. It is often useful to be able to remove nullspace functions as well, 147 to be able to shrink them to zero if they do not contribute significantly to a given model fit. This can be done either by tweaking the penalty matrix so that it both smooths and shrinks 149 as the single penalty term increases, or by adding a new penalty term that just penalizes the 150 null space for the model. Figure 4 shows an example of what the basis functions (Fig. 4A), 151 and smoothing penalties and shrinkage penalties (Fig. 4B) look like for a 6-basis function cubic 152 spline and for a 6-basis function thin-plate spline. The random effects smoother we discussed 153 earlier is an example of a pure shrinkage penalty; it penalizes all deviations away from zero, no 154 matter the pattern of those deviations. This will come into play in section III, when we use 155 random effects smoothers as one of the components in fitting HGAMs. 156

#### DLM: say something about cs basis here

EDIT this figure 158

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Should this figure include the intercept basis function as well for each? Right now I've excluded as mgcv drops it automatically, but it is part of the basis, and comes into play with tensor products...

Do we actually need the cubic spline here? It would simplify the presentation to just show the thin plate spline, and it gets the idea across.

#### Comparison to hierarchical linear models

Generalized linear mixed effect models (GLMMs; also referred to as hiearchical generalized linear models, multilevel models etc; e.g., Bolker et al. 2009; Andrew Gelman 2006) are an extension 166 of regression modelling that allow the modeller to include structure in the data – the structure is usually of the form of a nesting of the observations. For example individuals are nested within sample sites, sites are nested within forests and forests within states. The depth of the nesting is limited by the fitting procedure and number of parameters to estimate.

HGLMs are a highly flexible way to think about groupings in the data, the groupings used in 171 the models often refer to the spatial or temporal scale of the data (McMahon and Diez 2007) though can be based on any useful grouping.

We would like to be able to think about the groupings in our data in a simple way, even when the covariates in our model are related to the response in a non-linear way. The next section investigates the extension of the smoothers we showed above to the case where each observation is in a group, with a group-level smooth.

\*\* this last section should say something like: these are both latent gaussian thingos, the model structure is the difference – we want to be able to heirarchically structure our smoothers\*\*

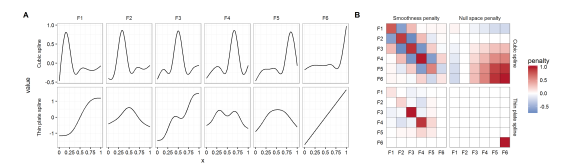


Figure 4. a) Examples of the basis functions associated with a six basis function thin plate spline (top) and a cubic spline (bottom) calculated for x data spread evenly between x=0 and x=1. Each line represents a single basis function. To generate a given smooth function, each basis function would be multiplied by its own coefficient. b) The smoothing and shinkage penalty matrices for the thin plate and cubic smoothers shown on left. Red entries indicate positive values and blue indicate negative values. For example, for the thin plate spline, functions f3 and f4 would have the greatest proportionate effect on the total penalty (as they have the largest values on the diagonal), whereas function f6 would not contribute to the penalty at all (all the values in the 6th row and column of the penalty matrix are zero). This means function f6 is in the null space of this basis, and would be treated as completely smooth.

#### III: WHAT ARE HIERARCHICAL GAMS?

#### What do we mean by hierarchical smooths?

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The smoothers in section II allowed us to model flexible relationships between our response and predictor variables. In this section, we will describe how to model model inter-group variability using smooth curves and how to fit these models in mgcv. Model structure is key in this framework, so we start with three choices:

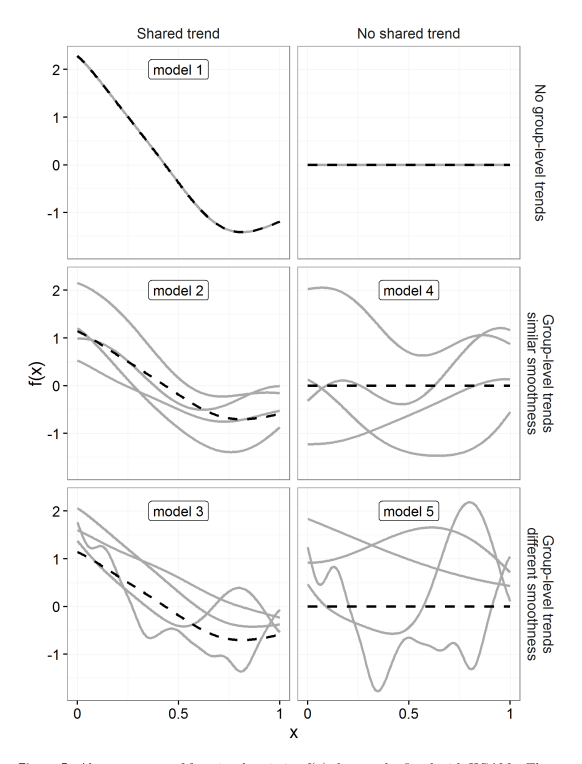
- 1. Should each group have its own smooth, or will a global smooth term suffice?
- 2. Do all of the group-specific curves have the same smoothness, or should each group have its own smoothing parameter?
- 3. Will the smooths for each group have a similar shape to one another a shared average curve<sup>1</sup>?

These three choices result in five possible models (figure 5), beyond the null model of "no relation between response and predictor(s)".:

- 1. A single common smooth for all observations.
- 2. A single common smooth plus group-level smooths that have the same wigglyness.
  - 3. A single common smooth plus group-level smooths with differing wigglyness.
- 4. Group-specific smooths without an average trend, but with all smooths having the same wigglyness.
  - 5. Group-specific smooths with different wigglyness.

It is important to note that "similar wiggliness" and "similar shape" are distinct ideas; two functions can have very similar wiggliness but very different shapes. Wiggliness simply measures

<sup>&</sup>lt;sup>1</sup>For this paper, we consider two functions to have similar shape if the average squared distance between the functions is small (assuming the functions have been scaled to have a mean value of zero across their ranges). This definition is somewhat restricted; for instance, a cyclic function would not be considered to have the same shape as a phase-shifted version of that function, nor would two normal distributions with the same mean but different standard deviations. The benefit of this definition of shape, however, is that it is straightforward to translate into quadratic penalties as we have been using.



**Figure 5.** Alternate types of functional variation f(x) that can be fitted with HGAMs. The dashed line indicates the average function value for all groups, and each solid line indicates the functional value at a given predictor value for an individual group level.

how quickly a function changes across its range, and it is easy to construct two functions that differ in shape but have the same wiggliness. For example, a logistic curve might have the same squared total second derivative between -1 and +1 as a sine curve, but they have very different shapes. Figure 5, model 4 illustrates this case. Similarly, two curves could have very similar overall shape, but differ in their wiggliness. For instance, if one function was equal to the second function plus a high-frequency osscilation. Figure 5 model 3 illustrates this.

We will discuss the trade-offs between different models and guidelines about when each of these models is appropriate in section IV. The remainder of this section will focus on how to specify each of these five models using mgcv.

#### 210 Coding hierarchical GAMs in R

# EJP: Going with canned and simulated data for the examples rather than real as it's a bit less messy

Each of these models can be coded straightforwardly in mgcv. To help illustrate this throughout the section when describing how to set these models up, we will refer to the response variable as y, continuous predictor variables as x (or  $x_1$  and  $x_2$ , in the case multiple predictors), and fac to designate the discrete grouping factor whose variation we are interested in understanding.

We will also use two example datasets to demonstrate how to code these models (see the appendix for code to generate these examples):

A. The CO2 dataset, available in R in the datasets package. This data is from an experimental study by Potvin, Lechowicz, and Tardif (1990) of CO<sub>2</sub> uptake in grasses under varying concentrations of CO<sub>2</sub>, measuring how concentration-uptake functions varied between plants from two locations (Mississippi and Quebec) and two temperature treatments (chilled and warm). A total of 12 plants were measured, and uptake measured at 7 concentration levels for each plant (figure 6a). Here we will focus on how to use these techniques to estimate inter-plant variation in functional responses.

B. A hypothetical study of what bird movement might look like along a migration corridor, sampled throughout the year. We have simulated this data for this paper (see supplemental code XX). This dataset consists of records of numbers of observed locations of 100 tagged individuals each from six species of bird, at ten locations along a latitudinal gradient, with one observation taken every four weeks. Not every bird was observed at each time point, so counts vary randomly between location and week. The data set (bird\_move) consists of the variables count, latitude, week and species (figure 6b). This example will allow us to demonstrate how to fit these models with interactions and with non-normal (count) data.

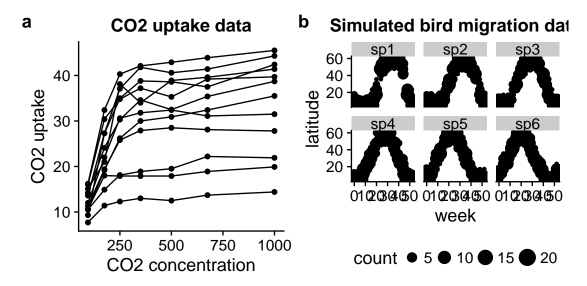
It is important to note that the grouping variable should be coded in R as an unordered factor – a character will raise an error and numeric will lead to a completely different model specification.
Whether the factor is ordered or not will not matter for most of the smoothers we use here.
However, for models 3&5 order will matter (see below for further details).

Throughout the examples we use Restricted Maximum Likelihood (REML) to estimate model coefficients and smoothing parameters. We strongly recommend using either REML or marginal likelihood (ML) when fitting GAMs for the reasons outlined in (Simon N. Wood 2011).

In each case some data processing and manipulation has been done to obtain the graphics and results below. We recommend readers take a look at the source RMarkdown [CITE] document for this paper to get the full code.

#### A single common smooth for all observations (Model 1)

We start with the simplest model we can in our framework and include many details here to ensure that readers are comfortable with the terminology and R functions we are going to use later.



**Figure 6.** Example data sets used throughout section III. a) Grass CO2 uptake versus CO2 concentration for 12 individual plants (black lines). b) Simulated data set of bird migration, with point size corresponding to weekly counts of 6 species along a latituidinal gradient (zeros excluded for clarity).

For our C02 data set, we will model  $\log_e(\text{uptake})$  as a function of two smooths: a thin plate regression spline of log concentration, and a random effect for species to model species-specific intercepts.<sup>2</sup> Mathematically:

$$\log_e(\mathtt{uptake}_i) = f(\log_e(\mathtt{conc}_i)) + \zeta_{\mathtt{Plant\_uo}} + \epsilon_i$$

where  $\zeta_{\text{Plant\_uo}}$  is the random effect for plant and  $\epsilon_i$  is a Gaussian error term. We assume that  $\log_e(\text{uptake}_i)$  is normally distributed.

DLM: not sure if this note is necessary...

DLM: need to justify why we use log concentration not just concentration? (could just cite?)

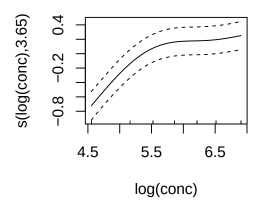
I think it's useful here as it shows one approach to deal with multipliciative functional variation. I've simplified it to remove the note about log-transforming concentration though. The reasons for that should be obvious

In R we can write our model as:

This is the typical GAM setup, with a single smooth term for each variable. Specifying the model is similar to specifying a glm in R, with the addition of s() terms to include one-dimensional or isotropic multidimensional smooths. The first argument to s() is the terms to be smoothed, the type of smooth to be used for the term is specified by the bs=... argument, and the number of basis functions is specified by k=....

<sup>&</sup>lt;sup>2</sup>Note that we're actually modelling ln(uptake); this can be a useful approach when dealing with estimating multiple functional relationships as it means that functions that differ from each other by a multiplicative constant (so  $f_1(x) = \alpha \cdot f_2(x)$ ) will differ by an additive constant when log-transformed (which can be estimated by simple random effects):  $ln(f_1(x)) = ln(\alpha) + ln(f_2(x))$ .

## s(Plant\_uo,10.83)



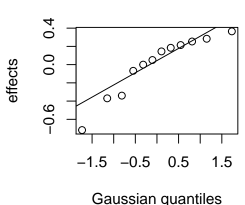


Figure 7. mgcv plotting output for model 1 applied to the CO2 dataset.

Figure 7 illustrates mgcv's default plotting out for CO2\_mod1: the left panel shows the estimated global functional relationship, and the right shows a quantile-quantile plot of the estimates effects vs Guassian quantiles, which can be used to check our model.

#### DLM: add more to plot description!

Looking at the effects by term is useful but we are often interested in fitted values or predictions our models. This can be useful to construct plots (like those in Figure ??). The next block of code shows how you could plot this to illustrate inter-plant variation in the functional response, plotting untransformed uptake and concentration to make the figure easier to interpret.

"'{r co2\_mod1\_ggplot, fig.width=6, fig.height=3, fig.cap="\label{fig:co2\_mod1\_predict}" # setup prediction data CO2\_mod1\_pred <- with(CO2, expand.grid(conc=seq(min(conc), max(conc), length=100), Plant\_uo=levels(Plant\_uo))) # make the prediction, add this and a column of standard errors to the prediction # data.frame. Predictions are on the log scale. CO2\_mod1\_pred <- cbind(CO2\_mod1\_pred, predict(CO2\_mod1, CO2\_mod1\_pred, se.fit=TRUE))

#### MAKE THE PLOT

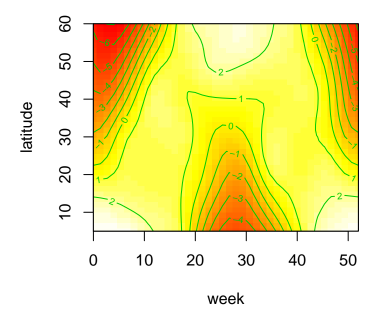
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ggplot(data=CO2, aes(x=conc, y=uptake, group=Plant\_uo)) + facet\_wrap(~Plant\_uo) + geom\_point() + geom\_line(aes(y=exp(fit)), data=CO2\_mod1\_pred) + geom\_ribbon(aes(ymin=exp(fit - 2se.fit), ymax=exp(fit + 2se.fit), x=conc), data=CO2\_mod1\_pred, alpha=0.3, inherit.aes=FALSE) "'

We can include interactions in an s() term via isotropic smooths such as thin plate regression splines or we can use the tensor product (te()) function, if we don't believe the composite terms are isotropic. In this case bs and k can be specified as a single value (in which case each marginal smooth has the same basis or complexity) or as a vector of basis types or complexities. For example, y~te(x1,x2, k=c(10,5), bs=c("tp","cs")), would specify a non-isotropic smooth of x1 and x2, with the marginal basis for x1 being a thin plate regression spline with 10 basis functions, and the smooth of x2 being a cubic regression spline with a penalty on the null space.

For our bird example, we want to look at the interaction between location and time, so for this we setup the model as:

### te(week,latitude,44.85)



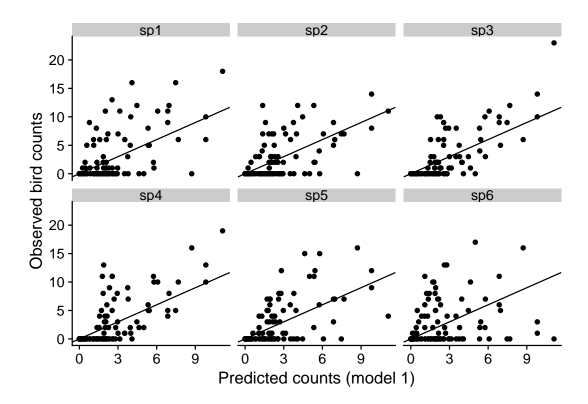
**Figure 8.** The default plot for this GAM illustrates the average log-abundance of all bird species at each latitude for each week, with yellow colours indicating more individuals and red colours fewer.

```
count_i = exp(f(week_i, latitude_i))
```

where we assume that  $\mathtt{count}_i \sim \mathtt{Poisson}$ . For the smooth term, f, we employ a tensor product of latitude and week, using a thin plate regression spline for the marginal latitude effects, and a cyclic cubic spline for the marginal week effect to account for the cyclic nature of weekly effects (we expect week 1 and week 52 to have very similar values), both splines had basis complexity (k) of 10. We will also assume the counts of individuals at each location in each week follow a Poisson distribution, and we will ignore species-specific variability.

Figure 8 shows birds starting at low latitudes in the winter then migrating to high latitudes from the 10th to 20th week, staying there for 15-20 weeks, then migrating back. However, the plot also indicates a large amount of variability in the timing of migration. The source of this variability is apparent when looking at the specifics of migration timing of each species (figure 6b).

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**Figure 9.** Observed counts by species versus predicted counts from bird\_mod1 (1-1 line added as reference). If our model fitted well we would expect that all species should show similiar dispersions around the 1-1 line. Instead we see that variance around the predicted is much higher for species 1 and 6.

All six species in figure 6b) show relatively precise migration patterns, but they differ in the timing of when they leave their winter grounds and the time they spend at their summer grounds. Averaging over all of this variation results in a relatively imprecise (diffuse) average estimate of migration timing (figure 8, 9), and viewing species-specific plots of observed versus predicted values (figure 9), it is appearent that the model fits some of the species better than others. This model could potentially be improved by adding inter-group variation in migration timing. The rest of this section will focus on how to model this type of variation.

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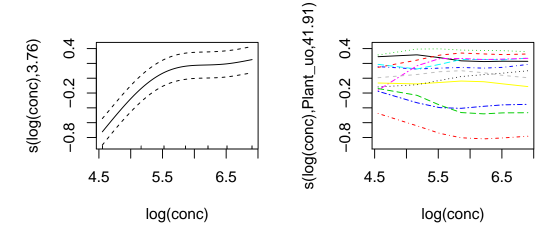
```
bird_move$mod1 = predict(bird_mod1, type="response")

ggplot(bird_move, aes(x=mod1, y= count))+
  facet_wrap(~species)+
  geom_point()+
  geom_abline()+
  labs(x="Predicted counts (model 1)", y= "Observed bird counts")+
  cowplot::theme_cowplot()
```

#### A single common smooth plus group-level smooths that have the same wigglyness (Model 2)

Model 2 is a close analogue to a GLMM with varying slopes: all groups have similar functional responses, but allows for inter-group variation in responses. This approach works by allowing each grouping level to have its own functional response, but penalizing functions that are too far from the average.

This can be coded in mgcv by explicitly specifying one term for the global smooth (as in model 1 above) then adding a second smooth term specifying the group level smooth terms, using



**Figure 10.** Global function (left) and group-specific deviations from the global function (right) for CO2\_mod2

a penalty term that tends to draw these group-level smooths to zero. For one-dimensional smooths, mgcv provides an explicit basis type to do this, the factor smooth or "fs" basis (see ?smooth.construct.fs.smooth.spec for detailed notes). This smoother creates a copy of each set of basis functions for each level of the grouping variable, but only estimates one set of smoothing parameters for all groups. The penalty is also set up so each component of its null space is given its own penalty (so that all components of the smooth are penalized towards zero)<sup>3</sup>. As there can be issues of co-linearity between the global smooth term and the group-specific terms (see section V for more details), it is generally necessary to use a smoother with a more restricted null space than the global smooth; for thin plate splines this can be done by setting m=2 for the global smooth and m=1 for the group smooth (Baayen et al. 2016) [also cite Wieling paper here]. e.g.: y~s(x,bs="tp",m=2)+s(x,fac,bs="fs",m=1,xt=list(bs="tp")).

We modify our previous  $CO_2$  model as follows:

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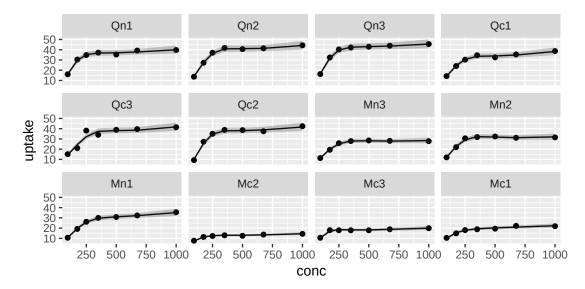
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$$\log_e(\mathtt{uptake}_i) = f(\log_e(\mathtt{conc}_i)) + f_{\mathtt{Plant\_uo}_i}(\log_e(\mathtt{conc}_i)) + \epsilon_i$$

where  $f_{\texttt{Plant\_uo}_i}(\log_e(\texttt{conc}_i))$  is the smooth of concentration for the given plant. In R we then have:

Figure 10 shows the fitted smoothers for CO2\_mod2. The plots of group-specific smooths indicate that plants differ not only in average log-uptake (which would correspond to each plant having a

<sup>&</sup>lt;sup>3</sup>As part of the penalty construction, each group will also have its own intercept (part of the penalized null space), so there is no need to add a separate term for group specific intercepts as we did in model 1.



**Figure 11.** Predicted uptake values (lines) verus oberved uptake for each plant, based on CO2 model 2.

straight line at different levels for the group-level smooth), but differ slightly in the shape of their functional responses. Figure 11 shows how the global and group-specific smooths combine to predict uptake rates for individual plants:

The "fs"-based approach mentioned above does not work for higher-dimensional tensor product smooths (if one is willing to use thin plate regression splines for the multivariate smooth then one can use "fs"). Instead, the group-specific term can be specified with a tensor product of the continuous smooths and a random effect for the grouping parameter. This term will again have a separate set of basis functions for each group, one penalty for the smooth term, and a second penalty drawing all basis functions toward zero<sup>4</sup>. e.g.: y~te(x1,x2,bs="tp",m=2)+te(x1,x2,fac,bs=c("tp","tp","re"),m=1). We illustrate this approach below on the bird migration data.

Model 2 is able to effectively capture the observed patterns of interspecific variation in migration behaviour (figure 12a), shows a much tigher fit between observed and predicted values, as well as less evidence of over-dispersion in some species compared to model 1 (figure 12b).

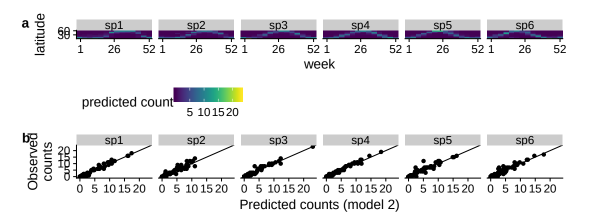
#### A single common smooth plus group-level smooths with differing wigglyness (Model 3)

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This model class is very similar to model 2, but we now allow each group-specific smooth to have its own smoothing parameter and hence it's own level of wigglyness. This increases the computational cost of the model, and means that the only information shared between groups is through the global smoothing term. This is useful if different groups differ substantially in how variable they are.

Fitting a seperate smooth term (with its own penalties) can be done in mgcv by using the by=fac argument in the s() function. Therefore, we can code this model as: y~s(x,bs="tp") + s(x, by=fac, m=1, bs= "ts") + s(fac, bs="re"). Note two major differences from how model 2

<sup>&</sup>lt;sup>4</sup>Note that this differs from the "fs" penalty, which assigned one penalty per null space term.



**Figure 12.** a) Predicted migration paths for each species based on bird\_mod2, with lighter colors corresponding to higher predicted counts. b) Observed counts versus predictions from bird\_mod2.

was specified: 1., we explicitly include a random effect for the intercept (the bs="re" term), as group-specific intercepts are not incorporated into these smooth terms automatically (as would be the case with bs="fs" or a tensor product random effect); 2., we explicitly use a basis with a fully penalized null space for the group-level smooth (bs="ts", for "thin plate with shrinkage"), as this method does not automatically penalize the nullspace, so there is potential for co-linearity issues between unpenalized components of the global and group-level smoothers.

Our CO2 model is then modified as follows:

Figure 13 shows a subsample of the group-specific smooths from this model, to prevent crowding.

It is appearent from this that some groups (e.g. Qc1) have very similar shapes to the global smooth (differing only in intercept), others do differ from the global trend, with higher uptake at low concentrations and lower uptake at higher concentrations (e.g. Mc1, Qn1), or the reverse pattern (e.g. Mn1).

Using model 3 with higher-dimensional data is also straightforward; by=fac terms work as well in tensor-product smooths as they do with isotrophic smooths. We can see this with our bird model:

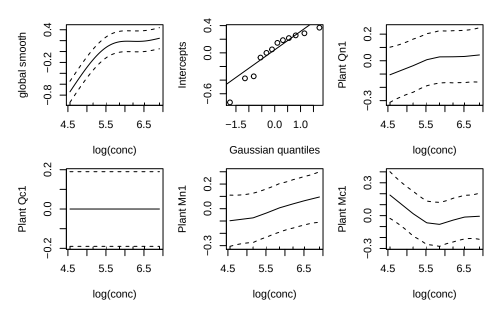
The fitted model for bird\_mod3 is visually indistinguishable from bird\_mod2 (figure 12) so we do not illustrate it here.

#### Models without global smooth terms (models 4 and 5)

We can modify the above models to exclude the global term (which is generally faster; see section V). When we don't model the global term, we are allowing each factor to be different, though there may be some similarities in the shape of the functions.

#### Model 4:

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**Figure 13.** Functional relationships for the CO2 data estimated for model 3. Top left: the global smooth; Top middle: species-specific random effect intercepts. The remaining plots are a selected subset of the plant-specific smoothers, indicating how the functional response of that plant differs from the global smooth.

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Model 4 (shared smooths) is simply model 2 without the global smooth term: y~s(x,fac,bs="fs") or y~te(x1,x2,fac,bs=c("tp","tp","re"). This model assumes all groups have the same smoothness, but that the individual shapes of the smooth terms are not related. (Plots are very similar to model 2.)
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#### Model 5:

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Model 5 is simply model 3 without the first term: y~s(x,by=fac) or y~te(x1,x2, by=fac). (Plots are very similar to model 3.)

Where group-level smooths are coded using the by=fac argument in the s() function,; if the factor is unordered, mgcv will set up a model with one smooth for each grouping level. If the factor is ordered, mgcv will not set the basis functions for the first grouping level to zero. In model 3 (with an ungrouped smooth included) the ungrouped smooth will then correspond to the first grouping level, rather than the average functional response, and the group-specific smooths will correspond to deviations from the first group. In model 5, using an ordered factor will result in the first group not having a smooth term associated with it at all.

#### V: MODELLING ISSUES

Which of the five models should you choose for a given data set? There are two major trade-offs to take into account. The first is the bias-variance trade-off: more complex models can account for more fluctuations in the data, but also tend to give more variable predictions, and can overfit. The second tradeoff is model complexity versus computer time: more complex models can include more potential sources of variation and give more information about a given data set, but will generally take more time and computational resources to fit and debug. We will discuss both of these trade-offs in this section.

#### Bias-variance tradeoffs

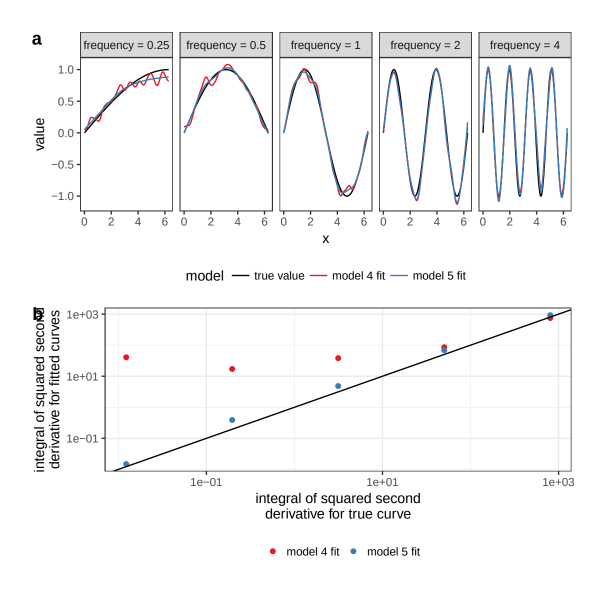
The bias-variance tradeoff is a fundamental concept in classical statistical analysis. When trying to estimate any value (in the cases we are focusing on, a smooth functional relationship between predictors and data), bias measures how on average an estimate is from the true value of the thing we are trying to estimate, and the variance of an estimator corresponds to how much that estimator would fluctuate if applied to multiple different samples taken from the same population. These two properties tend to be traded off when fitting models; for instance, rather than estimating a population mean from data, we could simply use a fixed value regardless of the observed data. This estimate would have no variance (as it is always the same) but would have high bias unless the true population mean happened to equal zero. The core insight into why penalization is useful is that the penalty term slightly increases the bias but can substantially decrease the variance of an estimator, relative to its unpenalized version (Efron and Morris 1977).

In GAMs and HGLMs, the bias-variance tradeoff is managed by the penalty terms (random effect variances in HGLM terminology). Larger penalties correspond to lower variance, as the estimated function is unable to wiggle a great deal, but also correspond to higher bias unless the true function is close to the null space for a given smoother (e.g. a straight line for thin-plate splines with 2nd derivative penalties, or zero for a standard random effect). The computational machinery used by mgcv to fit smooth terms is designed to find penalty terms that best trade off bias for variance to find a smooth that can effectively to predict new data.

The bias—variance tradeoff comes into play with HGAMs when choosing whether to fit separate penalties for each group level or assign a common penalty for all group levels (i.e. deciding between models 2 & 3 or models 4 & 5). If the functional relationships we are trying to estimate for different group levels actually vary in how wiggly they are, setting the penalty for all group-level smooths equal (models 2&4) will either lead to overly variable estimates for the least variable group levels, overly smoothed (biased) estimates for the most wiggly terms, or a mixture of these two, depending on how the fitting criteria used (ML, REML, or GCV) determines where the optimal smoothing parameter should be set.

We developed a simple numerical experiment to determine whether mgcv fitting criteria tend to set estimated smoothness penalties high or low in the presence of among-group variability in smoothness. We simulated data from five different groups, with all groups having the same levels of the covariate x, ranging from 0 to  $2\pi$ . For each group, the true function relating x to y was a sine wave, but the frequency varied from 0.25 (equal to half a cycle across the range of x) to 4 (corresponding to 4 full cycles across the range). We added normally distributed error to all y-values, with a standard deviation of 0.2. We then fit both model 4 (where all curves were assumed to be equally smooth) and model 5 (with varying smoothness) to the entire data set, using REML criteria to estimate penalties. For this example (Fig. 14a), requiring equal smoothness for all group levels resulted mgcv underestimating the penalty for the lowest frequency (most smooth) terms, but accurately estimating the true smoothness of the highest frequency terms as measured by the squared second derivative of the smooth fit versus that of the true function (Fig. 14b).

<sup>&</sup>lt;sup>5</sup>While this example may seem contrived, this is exactly what happens when we assume a given fixed effect is equal to zero (and thus exclude it from a model).



**Figure 14.** plotting example

This implies that assuming equal smoothness will tend to lead to underestimating the true smoothness of low-variability terms, and thus leading to more variable estimates of these terms. If this is a potential issue, we recommend fitting both models and using the model evalution techniques discussed in Section IV to determine if there is evidence for among-group variability in smoothness. For instance, the AIC for model 4 fit to this data is -180, whereas it is -211, implying a substantial improvement in fit by allowing smoothness to vary. However, it may be the case that there is too few data points per group to estimate seperate smoothness levels, in which case model 2 or model 4 may still be the better option even in the face of varying smoothness.

The ideal case would be to assume that among group penalties follow their own distribution (estimated from the data), to allow variation in smoothness while still getting the benefit of pooling information on smoothness between groups. However, this is currently not implemented in mgcv (and would be difficult to set up via mgcv's method of structuring penalties). It is possible to set up this type of varying penalty model in flexible Bayesian modelling software such as Stan or INLA (see below for a discussion of these tools), but how to set up this type of model has not been well studied, and is beyond the scope of this paper.

It may seem like there is also a bias-variance tradeoff between choosing to use a single global smoother (model 1) or a global smoother plus group-level terms (model 2-3), as in model 1, all the data is used to estimate a single smooth term, and thus should have lower variance than models 2-3, but higher bias for any given group in the presence of inter-group functional variability. However, in practice, this tradeoff will already be handled by mgcv via estimating penalties; if there are no average differences between functional responses, mgcv will penalize the group-specific functions toward zero, and thus toward the global model. The choice between using model 1 versus models 2-3 should generally be driven by computational costs; model 1 is typically much faster to fit than models 2-3, even in the absence of among-group differences, so if there is no need to estimate inter-group variability, model 1 will typically be more efficient.

A similar issue exists when choosing between models 2/3 and 4/5; if all group levels have very different functional shapes, the global term will get penalized toward zero in models 2/3, so they will reduce to models 4/5. Again, the choice to include a global term or not should be made based on scientific considerations (is the global term of interest to estimate) and computational considerations (which we will discuss next).

#### Complexity - computation tradeoffs

GAMs and GLMMs have substantially increased the range of flexible models available to the average researcher, and the HGAM models we discussed in section III extend on this broad base. However, the more flexible a model is, the larger an effective parameter space any fitting software has to search to find parameters that can predict the observed data. While numerical algorithms for solving complex models are always improving, it can still be surprisingly easy to use up massive computational resources trying to fit a model to even relatively small datasets. While we typically want to choose a model based on model fit (see above and section IV) and our goals for what the model will be used for, computing resources can often act as an effective upper limit on possible model complexity. Fitting an HGAM means adding extra computational complexity on top of either a GAM model with only global terms or a GLMM without smooth terms. For a given data set (with a fixed number n data points) and assuming a fixed family and link function, the time it takes to compute a given HGAM will depend, roughly, on four factors: the number of basis functions to be estimated, the number of smooth penalties to be estimated, whether the model needs to estimate both a global smooth and groupwise smooths, and the algorithm used to estimate parameters and fitting criteria used.

The most straightforward factor that will affect the amount of computational resources is the number of parameters in the model. Adding group-level smooths (moving from model 1 to 2-5) means that there will be more regression parameters to estimate, since each grouping level needs a separate coefficient for each basis function in the smooth. For a dataset with g different groups and n data points, fitting a model will just a global smooth,  $y\sim s(x,k=k)$  will require only k coefficients, and takes  $\mathcal{O}(nk^2)$  operations<sup>6</sup> to evaluate, but fitting the same data using a group-level smooth (model 4,  $y\sim s(x,fac,bs="fs",k=k)$ ) will require  $\mathcal{O}(nk^2g^2)$  operations to evaluate; in effect, adding a group-level smooth will increase computational time by an order of the number of groups squared<sup>7</sup>. The effect of this is visible in the examples we fit in section III when comparing the number of coefficients and relative time it takes to compute model 1 versus the other models (Table 1). One way to deal with this issue would be to reduce the number of basis functions (k) used when fitting group-level smooths when the number of groups is large; in effect, this would increase the flexibility of the model to accommodate inter-group differences, while reducing its ability to model variance within any given group. It can also make sense to use more computationally efficient basis functions when fitting large data sets, such as p-splines or

<sup>&</sup>lt;sup>6</sup>To understand the effects of these terms, we will use "big-O" notation; when we say a given computation is of order  $\mathcal{O}(n\log n)$ , it means that, for that computation, as n gets large, the amount of time the computation will take will grown proportionally to  $n\log n$ , so more quickly than linearly with n, but not as fast as n squared.

<sup>&</sup>lt;sup>7</sup>Including a global smooth (models 2-3) or not (models 4-5) will not generally substantially affect the number of coefficients needed to estimate (compare the number of coefficients in Table 1, model 2 vs. model 4, or model 3 versus model 5). Adding a global term will only add at most k extra terms, and it actually ends up being less that that, as mgcv drops basis functions from co-linear smooths to ensure that the model matrix is full rank.

**Table 1.** Relative computational time and model complexity for different HGAM formulations of the two example data sets from section III. All times are scaled relative to the length of time model 1 takes to fit to that data set. The # of coefficients measures the total number of model parameters (including intercepts). The # of smooths is the total number of unique penalty values estimated by the model.

		# of terms	
model	relative time	coefficients	penalties
A. CO2 data			
1	1	17	2
2	5	65	3
3	14	65	14
4	4	61	3
5	9	61	13
B. bird movement data			
1	1	90	2
2	110	540	5
3	140	624	14
4	100	541	3
5	66	535	12

 $_{503}$  cubic splines, rather than thin-plate splines, as thin-plate splines can take a substantial amount of overhead to compute the actual basis functions to use [CITE].

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Adding additional smoothing parameters (moving from model 2 to model 3, or moving from model 4 to 5) is even more costly than increasing the number of coefficients to estimate, as estimating smoothing parameters is computationally intensive (Simon N. Wood 2011). This means that models 2 and 4 will generally be substantially faster than 3 and 5 when the number of groups is reasonably large, as models 3 and 5 fit a separate set of penalties for each group level. The effect of this is visible in comparing the time it takes to fit model 2 to model 3 (which has a smooth for each group) or models 4 and 5 for the example data (Table 1). Note that this will not hold for every model, though; for instance, model 5 takes less time to fit the bird movement data than model 4 does (Table 1B).

#### Alternative formulations: bam, gamm, and gamm4 (with a brief foray into Bayes)

When fitting models with large numbers of different group levels, it is often possible to speed up computation substantially by using one of the alternative fitting algorithms available through mgcv.

The first tool available, that requires the least changes to your code compared to the base gam 518 function, is the bam function. This function is designed to improve performance when fitting 519 data sets with large amounts of data. It uses two tools to do this. First, it saves on the amount 520 of memory needed to compute a given model by using a random subset of the data to calculate 521 the basis functions for the smoothers, then breaking the data up into blocks and updating model fit within each block (Simon N. Wood, Goude, and Shaw 2015). While this is primarily designed 523 to reduce the amount of memory needed to fit these models, it can also substantially reduce computation time. Second, the bam function, when fitting using its default "fREML" (for "Fast 525 REML") method, you can use the discrete when fitting the model. This option causes bam to 526 simplify each covariate to a set of discrete levels (instead of a continuous range), substantially 527 reducing the amount of computation needed. Setting "discrete = TRUE" lets bam estimate the number of bins to use for each covariate. It is also possible to manually specify the number of 529 bins by passing discrete a vector of values. See ?mgcv::bam for more details.

It also takes more computational overhead compared to gam to set a bam model up, so for small numbers of groups, it can actually be slower than gam (Figure 15), however, as the number

of groups increases, computational time for bam increases more slowly than for gam; in our simulation tests, when the number of groups is greater than 16, bam can be upward of an order of magnitude faster (Figure 15). Note that bam can be somewhat less computationally stable when estimating these models (i.e. less likely to converge) so it does typically make sense to still use gam for smaller data sets.

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The second option is to fit these models using one of two dedicated mixed effect model estimation packages, nlme and lme4. The mgcv package includes the function gamm that allows you to call nlme to solve a given GAM, automatically handling the transformation of smooth terms into random effects (and back into basis-function representations for plotting and other statistical analyses). To use 1me4, you will have to install the gamm4 package, and use the gamm4 function from this package. Using gamm or gamm4 to fit models rather than gam can substantially speed up computation when the number of groups is large, as both nlme and lme4 take advantage of the sparse structure of the random effects, where most basis functions will be zero for most groups (i.e. any group-specific basis function will only take a non-zero value for observations in that group level). As with bam, gamm and gamm4 are generally slower than gam for fitting HGAMs when the number of group levels is small (in our simulations, <8 group levels), however they do show substantial speed improvements even with a moderate number of groups, and were as fast as or faster to calculate than bam for all numbers of grouping levels we tested (Figure 15)<sup>8</sup>.

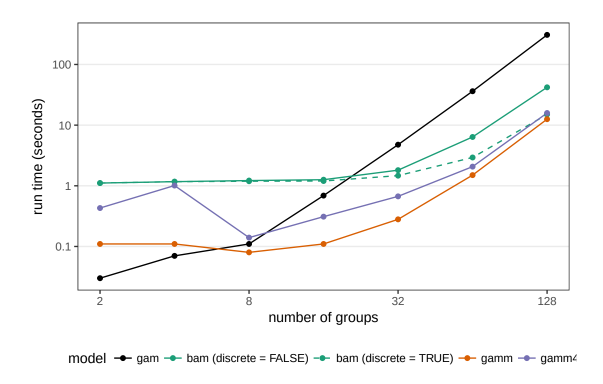
Setting up models 1-5 in bam uses the same code as we have previously covered; the only difference is that you use the bam instead of gam function, and have the additional option of discretizing your covariates. The advantage of this approach is that bam allows you to use almost all of the same families available to the gam function, and bam model output can be evaluated using the same functions (e.g. summary, AIC, plot, etc.) so it is simple to substitute for gam if you need to speed a model up.

Both gamm and gamm4 require at least a few changes to how you code models. First, there are a few limitations on how you are able to specify models 1-5 in both frameworks. Factor smooth (bs="fs") basis setups work in both gamm and gamm4. However, as the nlme package does not support crossed random effects, it is not possible to have two "fs" terms for the same grouping variable in gamm models (e.g. y~s(x1, grp,bs="fs"+s(x2, grp, bs="fs"). These type of crossed random effects are allowed in gamm4. The use of te and ti terms are not possible in gamm4 however, due to issues with how random effects are specified in the 1me4 package, making it impossible to code models where multiple penalties apply to a single basis function. Instead, for multidimensional group-level smooths, the alternate function t2 needs to be used to generate these terms, as it creates tensor products with only a single penalty for each basis function (see ?mgcv::t2 for details on these smoothers, and Simon N. Wood, Scheipl, and Faraway (2013) for the theoretical basis behind this type of tensor product). So for instance, model 2 for the bird movement data we discussed in section III would need to be coded as:

```
bird_mod4_gamm4 = gamm4(count ~ t2(week,latitude,species, bs= c("cc", "tp","re"),
                         k=c(10,10,6), m = 2),
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                    data= bird_move, family= poisson)
```

These packages also do not support the same range of families for the dependent variable; gamm only supports non-Gaussian families by using a fitting method called penalized quasi-likelihood (PQL) that is slower and not as numerically stable as the methods used in gam, bam, and gamm4. Non-Gaussian families are well supported by lme4 (and thus gamm4), but can only fit them using marginal likelihood (ML) rather than REML, so may tend to over-smooth relative to gam using REML estimation. Further, neither gamm nor gamm4 supports several of the extended families

<sup>&</sup>lt;sup>8</sup>It is also possible to speed up both gam and bam by using multiple processors in parallel, whereas this is not currently possible for gamm and gamm4. For large numbers of grouping levels, this should speed up computation as well, at the cost of using more memory. However, computation time will likely not decline linearly with the number of cores used, since not all model fitting sets are parallizable, and performance of the cores can vary. As parallel processing can be complicated and dependent on the type of computer you are using to configure properly, we do not go into how to use these methods here. The help file ?mgcv::mgcv.parallel explains how to use parallel computations for gam and bam in detail.



**Figure 15.** Elapsed time to estimate the same model using each of the four approaches. Each data set was generated with 20 observations per group using a unimodal global function and random group-specific functions consisting of an intercept, a quadratic term, and logistic trend for each group. Observation error was normally distributed. Models were fit using model 2:  $y\sim s(x,k=10,b=c'cp') + s(x,fac,k=10,b=c'fs',xt=list(b=c'cp'),m=1)$  All models were run on a single core.

available through mgcv, such as zero-inflated, negative binomial, or ordered categorical and multinomial distributions.

#### 581 Estimation issues when fitting both global and groupwise smooths

When fitting models with separate global and groupwise smooths (models 2 and 3), one issue to 582 be aware of is concurvity between the global smooth and groupwise terms. Concurvity measures how well one smooth term can be approximated by some combination of the other smooth terms 584 in the model (see ?mgcv::concurvity for details). For models 2 and 3, the global term is entirely 585 concurve with the groupwise smooths. This is because, in the absence of the global smooth term, 586 it would be possible to recreate that average effect by shifting all the groupwise smooths so they were centered around the global mean. In practical terms, this has the consequence of increasing 588 uncertainty around the global mean relative to a model with only a global smooth. In some 589 cases, it can result in the estimated global smooth being close to flat, even in simulated examples 590 with a known strong global effect. This concurvity issue may also increase the time it takes 591 to fit these models (for example, compare the time it takes to fit models 3 and 5 in Table 1). 592 That these models can still be estimated is because of the penalty terms; all of the methods we 593 have discussed for fitting model 2 ("fs" terms or random effect tensor products) automatically 594 create a penalty for the null space of the group-level terms, so that only the global term has 595 its own unpenalized null space, and both the REML and ML criteria work to balance penalties 596 between nested smooth terms (this is why nested random effects can be fitted). We have noted, 597 however, that mgcv still occasionally finds degenerate solutions with simulated data where the fitted global term ends up over-smoothed. 590

What we recommend to avoid this issue is to use a combination of smoother choice and setting model degrees of freedom so that the groupwise terms are either slightly less flexible or have a smaller null space. For instance, in the examples in section III, we used smoothers with an unpenalized null space (standard thin-plate splines) for the global smooth and ones with no null space for the groupwise terms<sup>9</sup>. When using thin-plate splines, it may also help to use splines with a lower order of derivative penalized in the groupwise smooths than the global smooths, as lower-order "tp" splines have fewer basis functions in the null space. For example, we used m=2 (penalizing squared second derivatives) for the global smooth, and m=1 (penalizing squared first derivatives) for groupwise smooths in models 2 and 3. Another option would be to use a lower number of basis functions (k) for groupwise relative to global terms, as this will reduce the maximum flexibility possible in the groupwise terms. We do caution that these are just rules of thumb. As of this writing, there is no published work looking what the effect of adding groupwise smooths has on the statistical properties of estimating a global smooth. In cases where an accurately estimated global smooth is essential, we recommend either fitting model 1, or using Markov Random Fields (Appendix A) and calculate the global smooth by averaging across grouping levels.

#### A brief foray into the land of Bayes

#### VI: EXAMPLES

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EJP: I think we should aim for 3-4 good examples here, highlighting different aspects of the model fitting problem. The example I'll be showing is a set of zooplankton community time series data, where multiple species were tracked throughout the year for a period of roughtly 20 years. This example can highlight both testing models 4/5 (for comparing different species' season cycles) and models 2/3 (for testing for differences between years for a single species). I think we need at least one example showing how to use these methods for multivariate regression (e.g. spatial analysis), and potentially an example showing how to these models work for

<sup>&</sup>lt;sup>9</sup>For model 2, the "fs" smoother, and tensor products of random effect ("re") and other smooth terms do not have a penalized null space by construction (they are full rank), as noted above. For model 3 groupwise terms, we used basis types that had a penalty added to the null space: bs="tp", "cs", or "ps" have this property.

non-normal data, and for including other covariates. In all examples, I think we should focus on how to fit each data set, visualize models, and compare different model fits.

629 In this final section, we will go through a few example analyses, to highlight how to use these 630 models in practice, and to illustrate how to fit, test, and visualize each model.

#### 631 Example 1: Inter- and intra-specific variation in zooplankton seasonal cycles over time

This first example will demonstrate how to use these models to fit community data, to show when using a global trend may or may not be justified, and to illustrate how to use these 633 models to fit seasonal time series. Here, we are using data from the Wisconsin Department 634 of Natural Resources collected by Richard Lathrop from a chain of lakes (Mendota, Menona, 635 Kegnonsa, and Waubesa) in Wisconsin, to study long-term patterns in the seasonal dynamics of 636 zooplankton. This data consists of roughly bi-weekly samples (during open-water conditions) of the zooplankton communities, taken from the deepest point of each lake via vertical tow collected 638 every year from 1976 to 1994 (the collection and processing of this data is fully described in Lathrop (2000)). We will use this data estimate variability in seasonality among species in the 640 community, and between lakes for the most abundant taxon in the sample (Daphnia mendotae). As we are focusing on seasonal cycles rather than average or maximum abudances, we have scaled all densities by log-transforming them then scaling by the within year species- and lake-specific mean and standard deviation (so all species in all lake-years will have a mean scaled density of zero and standard deviation of one).

This is what the data looks like:

zoo\_test = subset(zooplankton, year%2==1)

```
zooplankton = read.csv("../data/zooplankton_example.csv")
   str(zooplankton)
   ##
      'data.frame':
                        5848 obs. of 6 variables:
647
       $ day
                              10 10 10 10 10 10 10 10 12 12 ...
648
                               ##
       $ year
   ##
       $ lake
                        : Factor w/ 4 levels "Kegonsa", "Mendota", ...: 2 2 2 2 2 2 2 1 1
650
                        : Factor w/ 8 levels "Calanoida copepodites",..: 1 2 3 4 5 6 7 8 1 2 .
   ##
       $ taxon
651
                               5000 28000 3000 6000 1867000 ...
       $ density
                        : num
652
       $ density_scaled: num
                               -1.4173 -0.0714 -2.3837 0.1945 1.2613 ...
   ##
   levels(zooplankton$taxon)
   ## [1] "Calanoida copepodites"
                                       "Chydorus sphaericus"
   ## [3] "Cyclopoida copepodites"
                                       "Daphnia mendotae"
   ## [5] "Diacyclops thomasi"
                                       "Keratella cochlearis"
   ## [7] "Leptodiaptomus siciloides" "Mesocyclops edax"
   levels(zooplankton$lake)
   ## [1] "Kegonsa" "Mendota" "Menona"
                                         "Waubesa"
   We will split the data into testing and training sets, so we can evaluate how well our models fit
   out of sample. As there are multiple years of data here, we will use data from the even years to
   fit (train) models, and that from the odd years to test the fit:
   library(mgcv)
   zoo_train = subset(zooplankton, year\\\2==0)
   #the modulus (%%) finds the remainder after division by the right.
   #here we use it to find even numbers
```

Our first exercise here will be to demonstrate how to model community-level variability in seasonality, by regressing scaled density on day of year, with species-specific curves. As we are not interested here in average seasonal dynamics, we will focus on models  $4\&5^{10}$ . As this is seasonal data, we will use cyclic smoothers as the basis for seasonal dynamics.

zoo\_comm\_mod4 = gam(density\_scaled~s(day, taxon, bs="fs",k=10,xt=list(bs="cc")),

```
data=zoo_train,
           #we need to specify the start and end knots for day
           knots = list(day = c(1,365)),
           #We'll use ML as we are comparing models that differ in fixed effects
           method = "ML"
           )
summary(zoo_comm_mod4)
##
## Family: gaussian
## Link function: identity
##
## Formula:
## density_scaled ~ s(day, taxon, bs = "fs", k = 10, xt = list(bs = "cc"))
##
## Parametric coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) -0.02597
                           0.02621 - 0.991
##
## Approximate significance of smooth terms:
##
                  edf Ref.df
                                 F p-value
## s(day,taxon) 54.52
                          71 18.44 <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## R-sq.(adj) = 0.308
                         Deviance explained = 32.1%
## -ML = 3615.4 Scale est. = 0.64168
                                        n = 2947
# as all of the model features except the formula are the same in model 4arphi5,
# we just use the update function to refit the model with the new formula
zoo_comm_mod5 = update(zoo_comm_mod4,
                       formula = density_scaled~s(day, by=taxon, k=10,bs="cc"))
summary(zoo_comm_mod5)
##
## Family: gaussian
## Link function: identity
##
```

```
687
   ## Formula:
689
      density_scaled ~ s(day, by = taxon, k = 10, bs = "cc")
690
691
   ## Parametric coefficients:
692
   ##
                     Estimate Std. Error t value Pr(>|t|)
693
      (Intercept) -0.001563
                               0.014736 -0.106
                                                       0.916
694
   ##
695
   ## Approximate significance of smooth terms:
```

666

667

669

671

672

673

675

676

677

678

679

680

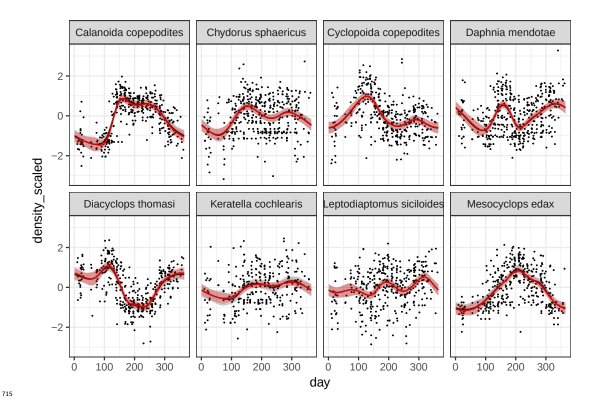
681 682

683

685

<sup>&</sup>lt;sup>10</sup>Here we are focusing on only the most common species in the data set. If we wanted to estimate the seasonal dynamics for rarer species, adding a global smooth term might be useful, so we could could borrow information from the more common species.

```
##
                                                edf Ref.df
                                                                F p-value
697
                                                         8 44.248 < 2e-16 ***
   ## s(day):taxonCalanoida copepodites
                                             6.968
                                                         8 8.933 2.47e-15 ***
   ## s(day):taxonChydorus sphaericus
                                             5.595
   ## s(day):taxonCyclopoida copepodites
                                             5.806
                                                         8 21.013 < 2e-16 ***
   ## s(day):taxonDaphnia mendotae
                                             6.941
                                                         8 15.977 < 2e-16 ***
                                             6.663
   ## s(day):taxonDiacyclops thomasi
                                                         8 38.303 < 2e-16 ***
   ## s(day):taxonKeratella cochlearis
                                             3.904
                                                         8 3.527 1.35e-06 ***
                                                         8 5.891 3.36e-09 ***
   ## s(day):taxonLeptodiaptomus siciloides 6.223
   ## s(day):taxonMesocyclops edax
                                             5.137
                                                        8 27.439 < 2e-16 ***
   ## ---
   ## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
707
   ## R-sq.(adj) =
                      0.31
                             Deviance explained = 32.1%
709
                                            n = 2947
   ## -ML = 3601.5 Scale est. = 0.63978
   We can see that both models have very similar fits, with an adusted R^2 of 0.308 for model
   4 and 0.31 for model 5. Model 5 has a somewhat lower AIC (AIC(zoo_comm_mod4) = 7115,
   AIC(zoo_comm_mod5) = 7104), implying a better overall fit. However, the two models show very
   similar fit to the data:
   library(ggplot2)
   library(dplyr)
   #Create synthetic data to use to compare predictions
   zoo_plot_data = expand.grid(day = 1:365, taxon = factor(levels(zoo_train$taxon)))
   zoo_mod4_fit = predict(zoo_comm_mod4, zoo_plot_data, se.fit = T)
   zoo_mod5_fit = predict(zoo_comm_mod5, zoo_plot_data, se.fit = T)
   zoo_plot_data$mod4_fit = as.numeric(zoo_mod4_fit$fit)
   zoo_plot_data$mod5_fit = as.numeric(zoo_mod5_fit$fit)
   zoo_plot_data$mod4_se = as.numeric(zoo_mod4_fit$se.fit)
   zoo_plot_data$mod5_se = as.numeric(zoo_mod5_fit$se.fit)
   zoo_plot = ggplot(zoo_plot_data, aes(x=day))+
     facet_wrap(~taxon, nrow = 2)+
     geom_point(data= zoo_train, aes(y=density_scaled),size=0.1)+
     geom_line(aes(y=mod4_fit))+
     geom_line(aes(y=mod5_fit),color="red")+
     geom_ribbon(aes(ymin = mod4_fit - 2*mod4_se,
                      ymax = mod4_fit + 2*mod4_se),
                  alpha=0.25)+
     geom_ribbon(aes(ymin = mod5_fit - 2*mod5_se,
                      ymax = mod5_fit + 2*mod5_se),
                  alpha=0.25, fill="red")+
     theme_bw()
   print(zoo_plot)
```



THe two curves are very close for all species, but the differences in smoothness that resulted in model 5 having an higher AIC than model 4 seem to be driven by the low seasonality of Keratella cochlearis and Leptodiaptomus siciloides relative to the other species. Still, both models show very similar fits to the training data, model 5 is only slightly better at predicting out of sample fits for K. cochlearis, and not at all better for L. siciloides:

```
# A tibble: 8 x 3
721
                                      mod4_cor mod5_cor
         taxon
722
         <fct>
                                         <dbl>
                                                   <dbl>
723
   ## 1 Calanoida copepodites
                                         0.690
                                                   0.700
724
   ## 2 Chydorus sphaericus
                                         0.360
                                                   0.350
725
   ## 3 Cyclopoida copepodites
                                         0.540
                                                   0.530
726
   ## 4 Daphnia mendotae
                                         0.340
                                                   0.350
   ## 5 Diacyclops thomasi
                                         0.810
                                                   0.810
728
   ## 6 Keratella cochlearis
                                         0.230
                                                   0.200
729
   ## 7 Leptodiaptomus siciloides
                                         0.330
                                                   0.330
730
   ## 8 Mesocyclops edax
                                         0.570
                                                   0.570
731
```

716

717

718

719

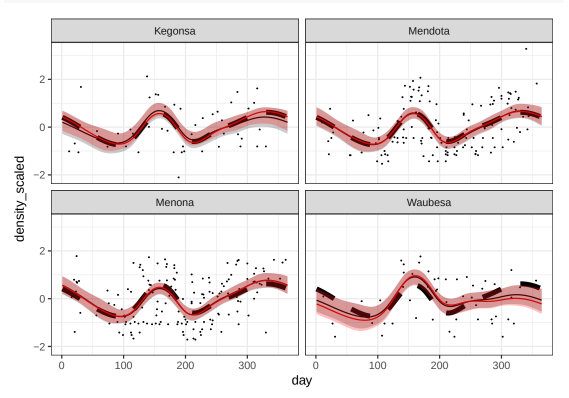
Now let's look at how to fit inter-lake variability in dynamics for just *Daphnia mendotae*. Here, we will compare models 1,2, and 3, to determine if a single global function is appropriate for

all four lakes, or if we can effectively model variation between lakes with a shared smooth or lake-specific smooths.

```
daphnia_train = subset(zoo_train,taxon=="Daphnia mendotae")
   daphnia_test = subset(zoo_test,taxon=="Daphnia mendotae")
   zoo_daph_mod1 = gam(density_scaled~s(day, bs="cc",k=10),
               data=daphnia_train,
               knots = list(day = c(1,365)),
               method = "ML"
   summary(zoo_daph_mod1)
736
   ## Family: gaussian
737
   ## Link function: identity
738
   ## Formula:
740
   ## density_scaled ~ s(day, bs = "cc", k = 10)
   ##
742
   ## Parametric coefficients:
                    Estimate Std. Error t value Pr(>|t|)
744
   ## (Intercept) 2.309e-16 4.245e-02
746
   ## Approximate significance of smooth terms:
               edf Ref.df
                               F p-value
748
   ## s(day) 6.824
                     8 13.96 <2e-16 ***
749
   ## ---
750
   ## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
751
752
   ## R-sq.(adj) = 0.217 Deviance explained = 23.1\%
   ## -ML = 519.41 Scale est. = 0.7262
   zoo_daph_mod2 = update(zoo_daph_mod1,
                           formula = density_scaled~s(day, bs="cc",k=10) +
                                                     s(day, lake, k=10, bs="fs",
                                                       xt=list(bs="cc")))
   summary(zoo_daph_mod2)
   ## Family: gaussian
   ## Link function: identity
757
   ##
758
   ## Formula:
759
   ## density_scaled \sim s(day, bs = "cc", k = 10) + s(day, lake, k = 10,
          bs = "fs", xt = list(bs = "cc"))
761
   ##
   ## Parametric coefficients:
763
   ##
                   Estimate Std. Error t value Pr(>|t|)
   ## (Intercept) 0.007772 0.043919
                                        0.177
765
   ##
   ## Approximate significance of smooth terms:
767
                     edf Ref.df
   ##
                                     F p-value
768
   ## s(day)
                  6.797 8 10.881 <2e-16 ***
769
                             35 0.432 0.0039 **
   ## s(day, lake) 5.348
```

```
## ---
771
                        0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
   ## Signif. codes:
772
773
   ## R-sq.(adj) = 0.246
                               Deviance explained = 26.9%
   ## -ML = 516.48 Scale est. = 0.69947
   zoo_daph_mod3 = update(zoo_daph_mod1,
                             formula = density_scaled~s(day, bs="cc",k=10) +
                                                        s(day,by=lake, k=10, bs="cc"))
   summary(zoo_daph_mod3)
776
   ## Family: gaussian
   ## Link function: identity
778
   ##
779
780
   ## density_scaled \sim s(day, bs = "cc", k = 10) + s(day, by = lake,
781
           k = 10, bs = "cc")
   ##
782
   ##
   ## Parametric coefficients:
784
   ##
785
                      Estimate Std. Error t value Pr(>|t|)
   ##
       (Intercept) -0.0007017 0.0416980 -0.017
786
   ##
   ## Approximate significance of smooth terms:
788
   ##
                                  edf Ref.df
                                                    F p-value
789
   ## s(day)
                                            8 13.985 < 2e-16 ***
                            6.8552361
790
   ## s(day):lakeKegonsa 0.0488015
                                            8
                                               0.006 0.34599
   ## s(day):lakeMendota 0.0005693
                                            8
                                               0.000 0.73813
   ## s(day):lakeMenona 0.9270931
                                               0.198 0.16129
   ## s(day):lakeWaubesa 2.2353408
                                            8 1.454 0.00116 **
   ## ---
795
                        0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
   ## Signif. codes:
796
797
   ## R-sq.(adj) = 0.246
                               Deviance explained = 26.5%
   ## -ML = 514.72 Scale est. = 0.69938
   The AIC values indicate that both model 2 and 3 are better fits than model 1, but models 2
800
   and 3 have similar fits to one another. There does not seem to be a large amount of inter-lake
   variability (all three models have similar adjusted R^2), and model 3 indicates that only Lake
   Waubesa deviates substantially from the overall dynamics. The plots for all three models (model
803
   1 as dashed line, model 2 in black and model 3 in red) show that Medota and Menona lakes are
804
   very close to the average and to one another for both models (which is unsurprising, as they are
   very closely connected by a short river) but both Kegons and Waubesa show evidence of a more
806
   pronouced spring bloom and lower winter abundances. While this is stronger in Lake Waubesa,
   model 2 (in black) shows that it is still detectable in Lake Kegonsa if we do not need to fit a
   separate penalty for each lake.
   library(ggplot2)
   library(dplyr)
   #Create synthetic data to use to compare predictions
   daph_plot_data = expand.grid(day = 1:365, lake = factor(levels(zoo_train$lake)))
   daph_mod1_fit = predict(zoo_daph_mod1, daph_plot_data, se.fit = T)
   daph_mod2_fit = predict(zoo_daph_mod2, daph_plot_data, se.fit = T)
```

```
daph_mod3_fit = predict(zoo_daph_mod3, daph_plot_data, se.fit = T)
daph_plot_data$mod1_fit = as.numeric(daph_mod1_fit$fit)
daph_plot_data$mod2_fit = as.numeric(daph_mod2_fit$fit)
daph_plot_data$mod3_fit = as.numeric(daph_mod3_fit$fit)
daph_plot_data$mod1_se = as.numeric(daph_mod1_fit$se.fit)
daph_plot_data$mod2_se = as.numeric(daph_mod2_fit$se.fit)
daph_plot_data$mod3_se = as.numeric(daph_mod3_fit$se.fit)
daph_plot = ggplot(daph_plot_data, aes(x=day))+
  facet_wrap(~lake, nrow = 2)+
  geom_point(data= daphnia_train, aes(y=density_scaled),size=0.1)+
  geom_line(aes(y=mod1_fit),linetype=2, size=2)+
  geom_line(aes(y=mod2_fit),color="black")+
  geom_line(aes(y=mod3_fit),color="red")+
  geom_ribbon(aes(ymin = mod2_fit - 2*mod2_se,
                  ymax = mod2_fit + 2*mod2_se),
              alpha=0.25)+
  geom_ribbon(aes(ymin = mod2_fit - 2*mod3_se,
                  ymax = mod2_fit + 2*mod2_se),
              alpha=0.25, fill="red")+
  theme_bw()
print(daph_plot)
```



In this case, model 2 is able to predict as good or better out of sample as model 1 or 3, indicating that jointly smoothing the lake together improved model prediction. However, None of the models did well in terms of predicting Lake Kegonsa dynamics out of sample (with a correlation

810

811

812

of only 0.11 between predicted and observed densities), indiciating that this model may be be missing substantial year-to-year variability in *D. mendotae* dynamics:

```
#Getting the out of sample predictions for both models:
   daphnia_test$mod1 = as.numeric(predict(zoo_daph_mod1,daphnia_test))
   daphnia_test$mod2 = as.numeric(predict(zoo_daph_mod2,daphnia_test))
   daphnia_test$mod3 = as.numeric(predict(zoo_daph_mod3,daphnia_test))
   # We'll look at the correlation between fitted and observed values for all species:
   daph_test_summary = daphnia_test %>%
     group_by(lake)%>%
     summarise(mod1_cor = round(cor(density_scaled, mod1),2),
                mod2_cor = round(cor(density_scaled, mod2),2),
                mod3_cor = round(cor(density_scaled, mod3),2))
   print(daph_test_summary)
   ## # A tibble: 4 x 4
816
   ##
        lake
                mod1_cor mod2_cor mod3_cor
817
         <fct>
                    <dbl>
                             <dbl>
                                      <dbl>
818
                   0.0800
                                     0.0900
   ## 1 Kegonsa
                             0.110
   ## 2 Mendota
                   0.420
                             0.430
                                     0.430
```

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0.390

0.270

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0.400

0.290

0.350

0.290

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