Scaling Integral Projection Models for Analyzing Size Demography

Alan E. Gelfand, Souparno Ghosh and James S. Clark

Abstract. Historically, matrix projection models (MPMs) have been employed to study population dynamics with regard to size, age or structure. To work with continuous traits, in the past decade, integral projection models (IPMs) have been proposed. Following the path for MPMs, currently, IPMs are handled first with a fitting stage, then with a projection stage. Model fitting has, so far, been done only with individual-level transition data. These data are used in the fitting stage to estimate the demographic functions (survival, growth, fecundity) that comprise the kernel of the IPM specification. The estimated kernel is then iterated from an initial trait distribution to obtain what is interpreted as steady state population behavior. Such projection results in inference that does not align with observed temporal distributions. This might be expected; a model for population level projection should be fitted with population level transitions.

Ghosh, Gelfand and Clark [*J. Agric. Biol. Environ. Stat.* **17** (2012) 641–699] offer a remedy by viewing the observed size distribution at a given time as a point pattern over a bounded interval, driven by an operating intensity. They propose a three-stage hierarchical model. At the deepest level, demography is driven by an unknown deterministic IPM. The operating intensities are allowed to vary around this deterministic specification. Further uncertainty arises in the realization of the point pattern given the operating intensities. Such dynamic modeling, optimized by fitting data observed over time, is better suited to projection.

Here, we address scaling of population IPM modeling, with the objective of moving from projection at plot level to projection at the scale of the eastern U.S. Such scaling is needed to capture climate effects, which operate at a broader geographic scale, and therefore anticipated demographic response to climate change at larger scales. We work with the Forest Inventory Analysis (FIA) data set, the only data set currently available to enable us to attempt such scaling. Unfortunately, this data set has more than 80% missingness; less than 20% of the 43,396 plots are inventoried each year. We provide a hierarchical modeling approach which still enables us to implement the desired scaling at annual resolution. We illustrate our methodology with a simulation as well as with an analysis for two tree species, one generalist, one specialist.

Key words and phrases: Hierarchical model, log Gaussian Cox process, Markov chain Monte Carlo, missing data.

Alan E. Gelfand is Professor, Department of Statistical Science, Duke University, Durham, North Carolina 27708, USA (e-mail: alan@stat.duke.edu). Souparno Ghosh is Assistant Professor, Department of Mathematics and Statistics, Texas Tech University, Lubbock, Texas 79409,

USA (e-mail: souparno.ghosh@ttu.edu). James S. Clark is Professor, Nicholas School of the Environment, Duke University, Durham, North Carolina 27708, USA (e-mail: jimclark@duke.edu).

1. INTRODUCTION

Population dynamics is a field with a long history in ecology and biology. Demography summarizes traits classified as stages. Matrix population models (MPMs), which assume stages are discrete classes, are usually used to describe changing structure (see, e.g., Keyfitz and Caswell, 2005, and references therein). Though changes in trait operate at the individual level, analysis of changing structure requires a translation of individual level data to the population level. In the past decade, the integral projection model (IPM) (Easterling, Ellner and Dixon, 2000; Ellner and Rees 2006, 2007; Rees and Ellner, 2009) has been offered as an alternative to matrix projection models when investigating continuous traits, for example, size, age, mass, leaf length. These models are built from demographic functions, parametric models for demographic processes specified in the form of vital rates like growth, maturation, survival, birth and fecundity; these rates are incorporated into a stationary redistribution kernel. For an estimated demographic model, projection refers to iterative projection of this kernel to steady state in order to attempt to answer questions regarding what would happen. So far, these models have only been fitted with individual-level transition data, that is, these data are used to estimate the demographic functions that comprise the kernel of the IPM specification. Then, projection proceeds through iteration, given the estimated kernel.

In recent work, Ghosh, Gelfand and Clark (2012) argue that such an approach introduces an inherent mismatch in scales. Working with tree diameters as the trait of interest, an individual level model describes the (conditional) transition of an individual of diameter xat time t to diameter y at time t + 1. On the other hand, an IPM essentially takes the distribution of diameters of individuals at time t to the distribution of diameters of individuals at time t + 1. We have a version of the familiar ecological fallacy (Wakefield, 2009). Moreover, in our application, we do not have any individual level transition data to attempt to scale up. For a given species, we only have the collection of diameters in a given plot, in a given year.

Ghosh, Gelfand and Clark (2012) offer a remedy by viewing the observed diameter distribution at a given time as a point pattern over a bounded interval, driven by an operating intensity. They propose a three-stage hierarchical model. At the deepest level, demography is driven by an unknown deterministic IPM. The operating intensities vary around this deterministic specification. Further uncertainty arises in the realization of the point pattern given the operating intensities. Ghosh, Gelfand and Clark (2012) argue that such dynamic modeling, optimized by fitting data observed over time, will better reveal how intensities, hence population structure, change over time. With individuallevel IPM model fitting, there is no mechanism to align projected trait distributions with trait distributions observed over time; consequential drift relative to the observed data can occur.

The contribution of this paper is to address scaling for population level IPMs with the objective of moving from plot level scale to larger scales, for example, the scale of the eastern U.S. Such scaling is intended to try to identify climate effects, which operate at a broader geographic scale, on demography. In turn, such scaling could allow assessment of changes in trait distributions and abundance, in response to climate change at larger scales. The threat of climate change is typically evaluated in terms of changes in distribution and abundance at regional scales (e.g., Guisan and Rahbeck, 2011). Our point pattern approach is attractive for scaling since we can cumulate intensities to explain aggregated point patterns. Individual level models cannot offer the desired scaling. Aggregation has to be done in climate space since the IPM kernels have arguments over trait space, not in geographic space, but introduce climate as covariates.

A further, critical contribution is an approach to handle severe sparsity in data collection. This emerges as a key feature with the USDA Forest Service's Forest Inventory and Analysis (FIA) data, which motivates our scaling objective. The FIA data is one of the few available data sets to investigate such scaling. Unfortunately, this database has enormous missingness. Of the roughly 44,000 plots, less than 20% are inventoried each year; more than 80% of plot level data over the time period 2000-2010 is missing. During this period a plot will have been inventoried two, possibly three times. Moreover, we have even more sparse sampling prior to 2004. Our study region includes the 31 eastern US states with climate conditions varying from hot and moist near the Gulf of Mexico to cold and dry near the Great Lakes. We focus on two illustrative species: Acer rubrum (ACRU) is a generalist and occupies a wide range of climate conditions. Liriodendron tulipifera (LITU) is restricted to the hot moist climate of the eastern and southeastern United States. Only in the recent work of Ghosh, Gelfand and Clark (2012) have IPM models been considered at the population scale. We are unaware of any applications at large geographic

scales or in the absence of data for consecutive time periods.

The format of the paper is as follows. In Section 2 we review IPMs and their properties as well as how we introduce uncertainty into the deterministic IPM specification. We also clarify the associated model fitting challenge, offering an approximation. To expedite flow and clarify the contribution here, explicit details of the model specification, as developed in Ghosh, Gelfand and Clark (2012), are deferred to the Appendix. In Section 3 we describe the FIA data set, as well as the climate data. In Section 4 we develop the scaling model ideas, first with a full data set, then for a very sparse data set. In Section 5 we provide a simulation investigation to demonstrate the loss of information due to the severe missingness as well as an analysis of the FIA data set. We offer some concluding remarks in Section 6.

2. INTEGRAL PROJECTION MODELS

In this section we briefly review the MPM, then turn to the IPM and its properties. We discuss how to introduce uncertainty into the IPM specification and conclude with a short discussion of how we fit these models. Again, MPMs and IPMs are techniques of choice for ecological demography. These models are specified with two indices, one for time, the other for trait level, for example, size, age, stage. There can be continuity or discreteness in time as well as continuity or discreteness in the trait space. With discrete time and categorical trait space, we have a MPM; with discrete time and continuous trait space, an IPM. As Sections 2.1 and 2.2 reveal, MPMs and IPMs are not associated with a specified region. There is no spatial index in these models. This reveals that scaling cannot be done over geographic space. A different approach is needed, which we propose in Section 4.

2.1 Matrix Projection Models

MPMs specify population structure dynamically. The state of the population at time t, as a vector of binned cell counts, $\mathbf{n}(t)$, is multiplied by a population projection matrix, \mathbf{A} , to yield the state of the population at time t + 1,

(1)
$$\mathbf{n}(t+1) = \mathbf{A}\mathbf{n}(t).$$

If the projection matrix is assumed to be time-invariant, a linear system of difference equations results to describe the evolution of the population. When one allows the projection matrix to vary because of external factors independent of the state of the population, a more general, time-varying difference equation version is obtained. If the projection matrix depends on the current state of population itself, denoted by $\mathbf{n}(t+1) = \mathbf{A}_n \mathbf{n}(t)$, we obtain a nonlinear model termed a density-dependent MPM. Tuljapurkar and Caswell (1997) and Caswell (2001) discuss the features of all these MPMs in detail. Caswell (2008) examines change in response of nonlinear matrix population models to changes in its parameters.

In (1), the A_{ij} give the average per-capita contribution from individuals in category *j* at time *t* to category *i* at time t + 1, either by survival, growth or reproduction. Typically, **A** is written as $\mathbf{A} = \mathbf{T} + \mathbf{F}$ with **T** describing transition (survival and growth) and F describing reproduction (fecundity). The stationary behavior of this matrix projection equation is obtained in terms of the eigenvalues (Λ_i) and eigenvectors (\mathbf{w}_i) of **A**. The long-term (*ergodic*) behavior of $\mathbf{n}(t)$ is determined by the dominant eigenvalue, $max(\Lambda_i)$, and associated right eigenvector (see the book of Caswell, 2001, for further details). Further eigenanalysis of the projection matrix yields a set of population statistics, viz., population growth rate, damping ratio, reproductive value and so on Caswell (2001). When the model is density dependent, the resulting behavior of the matrix equation cannot be written in terms of eigenvalues and eigenvectors (Caswell, 2001, page 504). In fact, equilibrium behavior need not exist.

2.2 The IPM and its Properties

For continuous traits and, in particular, for diameters, the MPM classes/stages are ordinal with definition being somewhat arbitrary. In this regard, Easterling, Ellner and Dixon (2000) and Ellner and Rees (2006) note that the IPM is proposed to remove the categorization required under the MPM approach. Here, we briefly review the behavior of an IPM as a deterministic specification. Working with intensities, $\gamma_t(\cdot)$, subscripted by time, we replace the MPM with

(2)
$$\gamma_{t+1}(y) = \int_{L}^{U} K(y; x) \gamma_t(x) \, dx.$$

The kernel K(y, x) is the IPM analog of the projection matrix **A** in a MPM. *L* and *U* are the lower and upper limits for the values of the trait.¹ Note that $\gamma_t(x)$ is an

¹Formally, *finite* point patterns are associated with bounded domains to ensure that the integral of the intensity over the domain is finite. In one dimension this means confining the support for γ to a bounded interval. Adopting this restriction in our setting implies that *L* can be 0, but we take $U < \infty$. This is a mild practical constraint.

intensity at time *t* implicitly associated with some region which we will refer to as a *plot*. So, if we integrate $\gamma_t(x)$ over an interval of diameters, we obtain the expected number of individuals in the plot with diameters in that interval, at time *t*. Therefore, $\gamma_{t,\cdot} = \int_L^U \gamma_t(x) dx$ is the expected number of individuals (population size) for the plot, at time *t*. Integrating (2) over *y* from *L* to *U* yields $\gamma_{t+1,\cdot} = \int_L^U K(\cdot, x)\gamma_t(x) dx$, where $K(\cdot, x) = \int_L^U K(y; x) dy; \gamma_{t+1,\cdot}$ can be compared with $\gamma_{t,\cdot}$. To give a population level interpretation to (2), it may be easiest to think in terms of intensity elements. That is, $\gamma_{t+1}(y) dy = \int_L^U K(y; x) dy \gamma_t(x) dy$ is the expected number of individuals in diameter interval (y, y + dy) at time t + 1 from all individuals in diameter interval (x, x + dx) at time *t*.

The eigenvalue theory for the IPM is directly connected to that for the MPM by viewing K(y; x) as a linear operator, that is, $Kh \equiv \int_{L}^{U} K(y; x)h(x) dx$. If Λ is the largest eigenvalue associated with K and w(x) is the associated right eigenfunction, $\int_{L}^{U} K(y; x) \cdot w(x) dx = \Lambda w(y)$, showing that, at steady state, Λ is the growth rate and w(x) (normalized) is the steady state diameter distribution.² As a result, $K^{t}w = \Lambda^{t}w$, but for arbitrary initial diameter distribution $\gamma_{0}(x)$, the projection $K^{t}\gamma_{0}$ need not be close to the projection $\Lambda^{t}\gamma_{0}$.

Specification of *K* introduces a survival and growth term as well as a fecundity or recruitment term. In practice, a time-independent *K* is not plausible; we employ a K_t that is dependent on levels of suitable environmental variables, say, \mathbf{Z}_t , during year *t*, as well as density dependent, that is, a function of $\gamma_{t,..}$ Details on how we introduce these features into K_t are supplied in the Appendix. The foregoing scaling challenge is apart from the form of $K_t(y; x)$ so, in the sequel, we treat K_t generically.

Sometimes normalization is introduced into the IPM. For instance, we might replace K(y; x) in (2) with, say, $K(y; x)/K(\cdot; x)$, a normalized version. However, this removes the interpretation of $\gamma_t(x)$ as an intensity since it imposes $\gamma_{t,\cdot}$ constant over *t*. Normalizing $\gamma_t(x)$ to the density $\tilde{\gamma}_t(x) = \gamma_t(x)/\gamma_{t,\cdot}$ is also unattractive since it now normalizes the resulting $\gamma_{t+1}(y)$ by $\gamma_{t,\cdot}$ rather than by $\gamma_{t+1,\cdot}$.

2.3 Introducing Uncertainty

As specified, the foregoing IPM is deterministic, raising the question of where and how to insert uncertainty. Within the Bayesian framework, a natural choice is to make the parameters random. However, a broader concern involves uncertainty associated with the form of K itself. Insisting that the IPM model is correct (even with "best" parameter estimates) is too restrictive. Rather, we view the outcome of the IPM as a sequence of intensities, $\gamma_t(y)$. Then, the operating intensity, $\lambda_t(y)$ (i.e., the intensity that drives the observed point pattern a time t), is assumed to vary around $\gamma_t(y)$. It is easier and more direct to specify uncertainty through the λ 's than through the K's; the latter will prove computationally infeasible. With regard to the intensities, we write $\lambda_t(x) = \gamma_t(x)e^{\varepsilon_t(x)}$, where ε_t is a stationary Gaussian process over [L, U] with covariance function $\sigma_{\varepsilon}^2 \rho(\cdot, \phi)$ and mean 0.³ We have a log Gaussian Cox process (Møller and Waagepetersen, 2004).

To allow for time-varying redistribution kernels, at least two approaches emerge. The first assumes that a vector of parameters is randomly chosen at each time point so that K_t takes the form $K(y, x; \theta(t))$. This strategy is employed (with individual-level data) in, for example, Rees and Ellner (2009), where, under parametric modeling for $K(y, x; \theta)$, the posterior for θ provides draws for $\theta(t)$. These draws may be interpreted as providing temporal random effects rather than parameter uncertainty. A more general regression approach is to assume that K is specified as a fixed parametric function but involving time-varying climate covariates and density dependence. Again, specific choices that we employ are supplied in the Appendix. In any event, we note that propagation of intensities through the K_t 's will not yield explicit forms (even for stationary K's). In fact, starting from time 0, at time t we will have a t-dimensional integration for γ_t .

Last, we do not apply the $K_t(y, x)$ to $\lambda_t(x)$. Rather, we allow the IPM to provide dynamics in a deterministic fashion for the γ_t 's, again viewing the λ_t 's driving the point patterns as varying around their respective γ_t 's. This specification suggests a pseudo-IPM approximation, as discussed in Section 2.4. As a result, the λ_t 's are conditionally independent given the { γ_t }'s. At

²The Perron–Frobenius theory tells us that, at this Λ , $w(x) \ge 0$ $\forall x$.

³It might be more natural to set the mean equal to $-\sigma_{\varepsilon}^2/2$ so that $E(\varepsilon_t(x)) = 1$. However, mean 0 only implies a scaling of λ relative to γ .

the highest level, we assume the point patterns, the \mathbf{x}_t 's, are conditionally independent given their { λ_t }'s with a nonhomogeneous Poisson process likelihood given by

(3)
$$[\mathbf{x}_t | \lambda_t] \propto \left[\exp\left(-\int_L^U \lambda_t(x) \, dx \right) \prod_{i=1}^{n_t} \lambda_t(x_{ti}) \right]$$

Hence, across *t*, the observed diameters are conditionally independent given $\lambda_t(x)$, but are marginally dependent due to the log Gaussian Cox process model for $\lambda_t(x)$.

Modeling is initiated with γ_0 , a kernel intensity estimate (Diggle, 2003). Hence, the full posterior is proportional to

$$\prod_{t=1}^{T} [\mathbf{x}_{t} | \lambda_{t}(x), x \in [L, U]]$$

$$(4) \qquad \cdot \prod_{t=2}^{T} [\lambda_{t}(x), x \in [L, U] | \gamma_{t}(x), x \in [L, U], \sigma^{2}, \phi]$$

$$\cdot [\{\gamma_{t}(\boldsymbol{\theta}, \gamma_{0}), t = 1, \dots, T\}] [\sigma^{2}] [\phi] [\boldsymbol{\theta}].$$

In (4), the bracketed term involving $\{\gamma_t\}$ is a degenerate distribution. It is employed here to denote the deterministic functional specification for the γ_t 's given the IPM and θ .

2.4 Model Fitting

We handle the stochastic integral in (3) by discretization, as described in the Appendix. However, the model described in (4), using (2), is computationally demanding to fit. The challenge arises because (2), with $K_t(y; x)$ as in (10) in the Appendix, does not have a closed form solution; we need to resort to numerical integration to create the sequence of $\{\gamma_t(x)\}$. Moreover, the dimension of the numerical integrations increases as the number of time epochs increases and consequently will result in an explosion of summations over time. An MCMC scheme will be computationally prohibitive because we will have to perform these integrations iteration by iteration. Following Ghosh, Gelfand and Clark (2012), we propose an approximate "pseudo" IPM approach, using adjacent pairs of years, that allows us to handle general K_t .

As above (14) in the Appendix, we consider x_j^* to be the center of the grid cell *j*. Then the pseudo-IPM update is given by

(5)
$$\gamma_{t+1}(x_j^*) = \sum_l K_t(x_j^*|x_l^*; \mathbf{z}_t, \boldsymbol{\theta}, \gamma_{t,\cdot}) \hat{\gamma}_t(x_l^*),$$

where $\hat{\gamma}_t(x)$ is an empirical estimate of the intensity corresponding to the point pattern observed at



FIG. 1. Graphical model driving the dynamics in the "pseudo" IPM.

time point t evaluated at the grid centers x_l^* . Under this updating scheme, for each t, we replace the tdimensional integral required to get $\gamma_t(x)$ in (2) by a one-dimensional integral. In (4), to obtain $[\{\gamma_t\}; \boldsymbol{\theta}]$ would require computing the γ_t deterministically and sequentially for a given $\boldsymbol{\theta}$, that is, $\prod_{1}^{T} [\gamma_t | \boldsymbol{\theta}, \gamma_{t-1}]$. Using (5), this now becomes $\prod_{1}^{T} [\gamma_t | \boldsymbol{\theta}, \mathbf{x}_{t-1}]$, where \mathbf{x}_{t-1} yields $\hat{\gamma}_{t-1}(x)$. The graphical model shown in Figure 1 captures the pseudo-IPM approximation. By analogy, pseudo-likelihood approximations in the literature often work with pairs (in our case, years) of observations, often with good asymptotic properties, though we cannot make such claims here. However, the fact that our approximation from time t to time t + 1 omits the uncertainty provided by λ_t suggests that we will underestimate uncertainty.

3. DATA TYPES AND THE FIA DATA SET

In order to clarify the proposed scaling approach, we first describe the motivating FIA data set as well as the associated climate data. With MPMs, demographic data are customarily available at the individual level over time. However, the case where the data is in the form of a time series of population vectors is also discussed (see Caswell, 2001). With data of the latter type, we observe a sequence of population vectors $\mathbf{n}(t_1), \mathbf{n}(t_2), \ldots$ without distinguishing the individuals. Dennis et al. (1995, 1997) use nonlinear multivariate time-series methodology to obtain the maximum likelihood estimates of the model parameters in this setting. With IPMs, such data would consist of a time series of point patterns for the trait distribution, for example, diameter, over the study region.

For individual level data, perhaps the most direct modeling strategy would be through a dynamic model with individual level random effects as in, for example, Clark et al. (2010). The state-space framework provides inference on individual variation in terms of population parameters, while being anchored directly by observations at the same scale (or with specifications that translate data to process scales, e.g., seed traps to trees). Afterward, desired population-level summaries can be created.

However, at large geographic scales, tracking of individuals is not feasible. For instance, we cannot hope to track individuals in thousands of forests on an annual basis over a span of decades. Collecting marginal point patterns at the scale of plots, without transition information on individuals, is more realistic. Even so, annual censusing of plots may not be. Hence, we need to be able to fit IPMs with data of the this type.

3.1 The FIA Data

The USDA Forest Services Forest Inventory and Analysis (FIA) program is the primary source for information about the extent, condition, status and trends of forest resources in the United States (Smith et al., 2009). FIA applies a nationally consistent sampling protocol using a quasi-systematic design covering all ownerships across the United States, resulting in a national sample intensity of one plot per 2428 hectare (Bechtold and Patterson, 2005) where plots are 54 m^2 . Aerial photography and/or classified satellite imagery is used to stratify the population (i.e., increase the precision of population estimates) and to establish permanent inventory plots in forest land uses. Forested land is defined as areas at least 10% covered by tree species, at least 0.4 ha in diameter, and at least 36.6 m wide. FIA inventory plots that are established in forested conditions consist of four 7.2 m fixed radius subplots spaced 36.6 m apart in a triangular arrangement with one subplot in the center (Bechtold and Patterson, 2005). All trees (standing live and dead) with a diameter at breast height (dbh) of at least 12.7 cm are inventoried on forested subplots. Within each subplot, a 2.07 m radius microplot offset 3.66 m from subplot center is established where only live trees with a dbh between 2.5 and 12.7 cm are inventoried. Within each microplot, all live tree seedlings are tallied according to species.

The program includes the measurement of a fixed proportion of the plots in each state, in each year, known as annual inventory. The legislative mandate requires measurement of 20% of the plots in each state, each year (FIA factsheet series, available online, http://www.fia.fs.fed.us/library/fact-sheets/datacollections/Sampling and Plot Design.pdf). In this analysis, the FIA data we employ were extracted from the most recent annual inventories (2000 to 2010) in 31 eastern states for a total of 43,396 inventory plots from FIADB version 4.0 on March 16, 2010 (available online http://fia.fs.fed.us/). The sampling is sparse in the initial years. Collection increases starts in 2003 and reaches its current level in 2006. The FIA plots sampled in year 2001 along with those sampled in 2006 are shown in Figure 2. A display of the set of plots sampled in 2007 would look almost the same as in Figure 2(b). Nonetheless, there would be no overlap between the two sets of plots!

3.2 The Climate Data

The climate data in this study was extracted from the 800m resolution Parameter-elevation Regressions on Independent Slopes Model (PRISM) data set (available online http://www.prism.oregonstate.edu/). Recognized as the highest quality spatial climate data sets, PRISM is a sophisticated interpolation that uses meteorological station data to produce continuous, digital grid estimates of climatic parameters, with consideration of location, elevation, coastal proximity, topographic facet orientation, vertical atmospheric layer, topographic position and orographic effectiveness of the terrain (Daly et al., 2008). In each FIA-measured plot, we used the climate data from the previous year to create the climatic covariates. We extracted the annual average precipitation (in mm) and the mean winter temperature (in °C), the average of January, February and March maximum and minimum monthly values.

Since climatological covariates operate over a broad geographical area, they may not explain the variation in, for example, diameter distribution of a species at the plot level (Canham and Thomas, 2010). Such variation will likely depend more on micro-scale covariates like soil moisture, nutrient availability and so on, which are not available to us. As a result, an approach to enable climate to provide explanation of demography is through suitable scaling. The scaling model described in the next section offers a viable way to study diameter distribution with this objective. The key idea is to partition climate space into bins.

4. A SCALING MODEL

We first present the extension of the model in (2) to enable the desired scaling in the context of a full data



FIG. 2. Map of the sampled FIA plots in (a) 2001 and (b) 2006.

set, that is, a data set supplying annual diameter distributions (point patterns) for every FIA plot. Then we present the model we employ to deal with the extreme sparseness. A convenient way to appreciate the scaling challenge is to envision a rectangular array where the rows represent the plots, the columns denote the years, and, in a given cell, we have a point pattern of diameters. In a full data set, we have an observed point pattern in every cell. Imagining this for the FIA data collection over ten years would entail more than 400,000 point patterns. With the actual FIA data collection, we have more than 80% of the cells empty and for no plot do we have point patterns in consecutive years.

4.1 The Full Data Scaling Model

Again, we note that the IPM is indexed by time and by trait value but there is no spatial index. In our context, there is redistribution over trait space but no redistribution over geographic space; the model specified in (2) operates at the plot level. With multiple plots, perhaps spatially-referenced, in principle, we could fit this model plot by plot. While such an analysis might be useful in some contexts, biogeographic studies require joint modeling across plots. In other words, it would be very difficult to develop a synthesis that would tell a big picture demography story from such an analysis and, as noted above, it would be difficult to capture climate effects. Moreover, it is evidently not scaling the data to larger geographic regions. In theory, with spatiallyreferenced plots, we might imagine introducing plotlevel spatial random effects into a joint model across plots. However, with the FIA data, plots are not contiguous; they are sparse across the eastern U.S. so such spatial analysis is not appropriate.⁴ Rather, as noted earlier, an attractive feature of working with point patterns and associated intensities in a Cox process setting is the convenience of aggregating intensities to explain aggregated point patterns. First, we consider how we might do this with a full data set.

Recall that each plot is subjected to a sequence of annual climate variables across the years of data collection. Suppose we partition the climate space into a collection of climate bins, indexed by l = 1, 2, ..., M. In the present setting, we have two climate covariates and we are partitioning the upper right quadrant of \mathbb{R}^2 . (Below, we say more about the choice of partition.) Suppose we label each climate bin by a suitable centroid (defined below). Then, we have a set of M climates which are labeled as \mathbf{z}_l^* . Suppose, in year t, we assign label $L_{j,t} = l$ to plot j if it received climate falling in bin l. Thus, in year t, all of these plots will have their diameter distributions operated on by the same redistribution kernel, that is, K as in (10) in the Appendix with $\mathbf{z}_t = \mathbf{z}_l^*$, apart from plot-specific density dependence. In particular, suppose $S_{t,l} = \{j : L_{j,t} = l\}$

⁴An alternative might be to introduce i.i.d. plot level random effects, but this will substantially increase the dimension of the model and such a model be difficult to fit, especially in our very sparse sampling setting.

and there are $n_{l,l}$ plots receiving climate \mathbf{z}_l^* in year t. Then, $\sum_{j \in S_{t,l}} \gamma_{j,t}(x)$ is the cumulated intensity subjected to K, with $\mathbf{z}_t = \mathbf{z}_l^*$ in year t. The only modification is that, if density dependence appears in K, then the population size for density dependence would be $\sum_{j \in S_{t,l}} \gamma_{j,t,\cdot}$. We acknowledge that density dependence operates at the plot scale, not at the scale of aggregation to climate bins. However, at larger scale, it is still arguable that an increase in aggregated abundance at year t will place a scaled increase in resource pressure on the species for year t + 1. Moreover, if we wish, we can compare models with and without aggregated density dependence.

If we do the above for each l, then in year t, every plot will have been assigned to a unique climate bin and we can fit the IPM across the M bins for year t. Then, if we do this for each year, we have jointly fitted the IPM across all plots for all years.

Finally, we assign as the "centroid" to bin *l* the average of all of the climates for all of the plots across all of the time points that fell into bin *l*. That is, we keep the partitions the same from year to year and we keep the labels constant across time as well. As for the creation of the partitions, we overlay a bounding rectangle on the observed climates for all plots and all years in the study. We then partition the temperature axis (mean winter temperature) as well as the precipitation axis (average annual precipitation) to create a rectangular grid. Some bins will be empty in one or more years; they are not considered. In fact, if, for a given species, there were no occurrences across all plots in a climate bin in a given year, then the bin is not considered for that year. We have no point pattern to drive the pseudo-IPM for the species in that year.

Again, we emphasize that we are scaling in climate space, not in geographic space; we are aggregating plots receiving essentially a common climate in a given year regardless of where they are in physical space. But, this raises the question of what projection means under such scaling? In climate space, we fit data across $M \times T$ cells but it makes no sense to project a climate bin in time. If we aggregated plots in geographic space, we would not necessarily find common climate for all plots in each year. So, we cannot do projection for such spatially aggregated plots? The conclusion is that we should think of projection at the plot level or for an aggregated collection of plots receiving the same sequence of climate variables over time. Projection under such aggregation may be adequate to address large scale response to broad, coarse spatial resolution climate scenarios. In any event, these limitations are not a criticism of our approach to scaling. Rather, they clarify what projection can entail. Moreover, as we shall see in the next subsection, our approach offers the only way to implement demographic modeling for the FIA data with its inherent sparseness.

To conclude here, we consider inference summaries under our modeling. First, we can present posterior summaries of the model parameters, that is, all of the parameters in the K_t 's. Next, we can develop posterior predictive intensities to compare with empirical intensities for different (l, t) combinations. Also, we can provide comparison of observed and predicted population sizes under various (l, t) combinations. Last, at the plot level, we can implement projection under the model and compare predicted intensities and population sizes with their observed counterparts.

4.2 Accommodating the Sparseness in the FIA Data

With regard to the discussion of the previous subsection, now, for every species, our plot by year array has more then 80% cells with no observed point pattern. As above, for plot j at time t, if $\mathbf{z}_{j,t}$ is the associated covariate, we will assign label $L_{j,t} = l$ if $\mathbf{z}_{j,t}$ falls in bin l. So, if $L_{j,t} = l$, the redistribution kernel operating at time t is $K_t(y, x; \mathbf{z}_l, \boldsymbol{\theta})$.

We formalize our modeling at the plot level, that is, for plot *j* in year *t*, if $L_{j,t} = l$,

(6)
$$\gamma_{j,t+1}(y) = \int K(y,x;\mathbf{z}_l,\boldsymbol{\theta})\gamma_{j,t}(x) dx.$$

With the foregoing notation, for year *t*, we have $S_{t,l} =$ $\{j: L_{j,t} = l\}, l = 1, 2, ..., M$, with $n_{t,l}$, the number of plots in $S_{t,l}$. The number of plots in $\bigcup_l S_{t,l}$ is n_t , the number of FIA plots in year t. (There is year to year variation.) Next, let $I_{j,t} = 1, 0$ if plot j is measured (observed) or not in year t. That is, if $I_{j,t} = 1$, we observe a point pattern, $\mathbf{x}_{l,t}$. Then, let $S_{t,l,1} =$ $\{j: L_{j,t} = l, I_{j,t} = 1\}, S_{t,l,0} = \{j: L_{j,t} = l, I_{j,t} = l\}$ 0}. Evidently, $S_{t,l,1} \cup S_{t,l,0} = S_{t,l}$. Similarly, define $R_{t,l,1} = \{j : L_{j,t} = l, I_{j,t+1} = 1\}, R_{t,l,0} = \{j : L_{j,t} = l\}$ $l, I_{j,t+1} = 0$. So, $R_{t,l,1} \cup R_{t,l,0} = S_{t,l}$. The idea here is, for plots that experienced \mathbf{z}_l in year t, we wish to capture the set which was observed at the start of the year $(S_{t,l,1})$ and the set which was observed at the *end* of the year $(R_{t,l,1})$. Again, $S_{t,l,1}$ and $R_{t,l,1}$ are disjoint, providing the crux of the missing data challenge. Let $n_{t,l,1}$ and $m_{t,l,1}$ denote the number of plots in $S_{t,l,1}$ and $R_{t,l,1}$, respectively.

From (6), we have the following conceptual IPM scaling:

(7)
$$\sum_{j \in S_{t,l}} \gamma_{j,t+1}(y) = \int_D K(y,x;\mathbf{z}_l,\boldsymbol{\theta}) \sum_{j \in S_{t,l}} \gamma_{j,t}(x) \, dx.$$

Again, we only see a subset of the plots on the left-hand side of (7) and also only a subset of the plots on the right-hand side. However, dividing both sides by $n_{t,l}$, we have the "per plot" (or average) intensity,

(8)
$$\bar{\gamma}_{l,t+1}(y) = \int_D K(y,x;\mathbf{z}_l,\boldsymbol{\theta})\bar{\gamma}_{l,t}(x)\,dx$$

with the obvious definition for the $\bar{\gamma}$'s.

But, this leads to the natural approximations: $\tilde{\gamma}_{l,t+1}(y) \approx \bar{\gamma}_{l,t+1}(y)$ and $\tilde{\gamma}_{l,t}(x) \approx \bar{\gamma}_{l,t}(x)$, where $\tilde{\gamma}_{l,t+1}(y) = \sum_{j \in R_{t,l,1}} \gamma_{j,t+1}(y)/m_{t,l,1}$ and $\tilde{\gamma}_{l,t}(x) = \sum_{j \in S_{t,l,1}} \gamma_{j,t}(x)/n_{t,l,1}$.

So, for each t and l, we work with the approximate IPM relationship,

(9)
$$\tilde{\gamma}_{l,t+1}(y) = \int_D K(y,x;\mathbf{z}_l,\boldsymbol{\theta})\tilde{\gamma}_{l,t}(x)\,dx.$$

To use this relationship, we have to do two things:

(i) create an empirical estimate of $\tilde{\gamma}_{l,t}(x)$ with the observed $\mathbf{x}_{j,t}$ for $j \in S_{l,t,1}$ to use on the right-hand side. We do this by creating an empirical intensity based upon all of the plots in $S_{t,l,1}$ and then scaling the intensity by $n_{t,l,1}$.

(ii) use the observed $\mathbf{x}_{j,t+1}$ for $j \in R_{t,l,1}$ to inform about the left-hand side. That is, the observed $\mathbf{x}_{j,t+1}$ are linked to $m_{t,l,1}\lambda_{l,t+1}(y)$, as above in expression (9), in the likelihood and $\lambda_{l,t+1}(y)$ is linked to $\tilde{\gamma}_{l,t+1}(y)$, up to log GP error, as in Section 2.3. To do this, we introduce a "per plot" log GP error for each climate bin in each year.

A further complication arises due to lack of information about new recruits [again, see (10)–(12) and related discussion in the Appendix]. In the absence of consecutive years of data, we cannot distinguish if an individual observed at time t + 1 in bin l is actually a new recruit (crossing the boundary from seedling to trees) or was in an unsampled plot in that bin at time t. Due to this ambiguity, δ_1 in (12) cannot be estimated reliably. Hence, the effect of density dependence on recruitment rate cannot be ascertained at this level of sparsity. As a result, we set $\delta_1 = 0$ and thereby assume a time-invariant Δ . As with a full data set but even more so, some blocks will be empty in one or more years; they are not considered. Again, if, for a given species, there were no occurrences across all plots in a bin in a given year, then we have no point pattern to drive the pseudo-IPM for that bin for that year. For that species, the bin is not considered for that year. Inference summaries will parallel those we proposed above to create with a full data set.

5. A SIMULATION AND AN FIA DATA ANALYSIS

In Section 5.1 we provide a simulation to show how well our approach works with a full data set and also to reveal the effect of the loss of information as we go to 50% and also 80% missingness. Then, in Section 5.2, we turn to the FIA data, to look at two species, recognizing the inference challenges imposed by the sparsity.

5.1 A Simulation Example

To illustrate the performance of the model under various level of missingness, we performed the following simulation study. We envisioned a covariate with four levels assigned as z = 0, 1, 2, 3 and to each covariate level we assigned 100 FIA plots from year 2005. Thus, the covariate level can be viewed as a climate, defining four "climate bins" with, initially, 100 plots in each bin. We used the empirical intensity associated with each plot (from the 2005 FIA data), say, $\hat{\gamma}_{lj,0}(x^*), l = 1, 2, 3, 4; j = 1, 2, \dots, 100$, as the initial condition, for x^* 's as described in Section 2.4. We plug this initial intensity in the pseudo-IPM (5) with covariate information inserted in the redistribution kernel (10) to obtain $\gamma_1(\cdot)$ (for each plot) at the following time point. With regard to K, we used the forms in the Appendix, fixing $Q_0 = 1$, $\delta_1 = 0$, and $\mu = 0$, with the remaining parameter values set as in Table 1. We seek to infer about these remaining parameters, including the regression coefficient β for climate, as well as to project 10 years forward. To mimic the FIA data set, we randomize the plots to changing climate at each time point following the illustrative transition matrix shown below:

Location at $t + 1$				
Location at t	0	1	2	3
0	0.7	0.2	0.07	0.03
1	0.2	0.7	0.03	0.07
2	0.07	0.03	0.7	0.2
3	0.03	0.07	0.2	0.7

TABLE 1Posterior summaries of model parameters under various levels ofmissingness for the simulated data. Posterior mean and 95%equal tail credible intervals provided

Parameters	True value	Level of missingness	Posterior summary
Q_1	0.01	0%	0.0171 (0.0058, 0.0292)
σ	0.25		0.2456 (0.1054, 0.3923)
δ_0	0.30		0.3467 (0.2161, 0.4860)
η	0.10		0.1253 (0.0548, 0.1953)
β	0.01		0.0091 (0.0034, 0.0148)
Q_1	0.01	50%	0.0174 (0.0016, 0.0275)
σ	0.25		0.2695 (0.0596, 0.4849)
δ_0	0.30		0.3281 (0.1181, 0.5643)
η	0.10		0.1242 (0.0530, 0.1962)
β	0.01		0.0146 (-0.0002, 0.0292)
Q_1	0.01	80%	0.0293 (0.0125, 0.0582)
σ	0.25		0.5702 (0.2756, 0.8323)
δ_0	0.30		0.3536 (0.1783, 0.5315)
η	0.10		0.0824 (0.0089, 0.2201)
β	0.01		0.0691 (0.0220, 0.1166)

In this fashion, for each plot, we generate a sequence of $\gamma_t(x^*)$ for 10 time points and then an associated point pattern. We then fit the scaling model (9) using the first nine time points of the complete data set. The average empirical intensity at each covariate bin at t = 1 [$\tilde{\gamma}_{l,1}(x^*)$] is used in fitting this full data model. The posterior summary of the parameters (Table 1) suggests that, when there is no missingness, the scaling model can recover the parameters of the plot level model.

In order to investigate our ability to recover the parameters under a moderate amount of missingness, we randomly remove 50 plots from each covariate bin at each of the first nine time points and fit the scaling model (9) using the remaining available 200 plots at each time point in the training set. The posterior summary in Table 1 shows that we can still recover certain parameters, although, as expected, the uncertainty associated with these estimates is higher than that obtained for the complete data set. Turning to extreme missingness, as in the FIA data, we randomly remove 80 plots from each covariate bin at every time point in the training set and then fit the scaling model to the remaining plots. The posterior summary in Table 1 shows that the intervals are now even longer and not well centered. In summary, our modeling approach is viable but, with very high levels of missingness, our ability to learn about the process will be limited.

Next, we illustrate how missingness leads to increased uncertainty in projection. Let Θ^{comp} and Θ^{ext}

be posterior samples of all the parameters obtained from fitting the scaling model on the complete data set and the data set with 80% missing plots, respectively. Using Θ_b^{comp} , the *b*th posterior sample of Θ^{comp} and $\tilde{\gamma}_{l,1}(x^*)$, we generate $\tilde{\gamma}_{l,2}(x^*; \Theta_b^{\text{comp}}), \ldots, \tilde{\gamma}_{l,10}(x^*, \Theta_b^{\text{comp}}), l = 1, 2, 3, 4; b = 1, 2, \ldots, B$. Similarly, using Θ_b^{ext} , we generate $\tilde{\gamma}_{l,1}(x^*; \Theta_b^{\text{ext}}), \ldots, \tilde{\gamma}_{l,10}(x^*, \Theta_b^{\text{ext}})$. Figure 3 shows the true $\tilde{\gamma}_{l,10}(x^*)$ along with the pointwise 95% CI obtained for $\tilde{\gamma}_{l,10}(x^*, \Theta_b^{\text{comp}})$ and $\tilde{\gamma}_{l,10}(x^*, \Theta_b^{\text{ext}})$ for l = 1, 2, 3, 4. In all cases we are able to contain the true ten year projections under the extreme missingness but, as expected, the uncertainties associated with the projection using Θ^{ext} are substantially higher than those obtained for Θ^{comp} .

5.2 Data Analysis for Two Species

We illustrate the scaling model on two species *Acer rubrum* (ACRU) and *Liriodendron tulipifera* (LITU). ACRU has a broad geographic distribution, its range extending from the Gulf Coast of the eastern United States to Canada. It can thrive on mesic (moderate moisture) to xeric (dry) sites. Compared with ACRU, LITU is much less widespread. Its range does not extend as far north, nor does it occupy xeric sites.

Due to the extreme sparsity of sampling prior to 2005, the scaling model is fitted on data from 2005 through 2010. We use the data from 2005 to provide the initial intensities. We partition the available information (from the PRISM data set) on pairs of mean winter temperature and average annual precipitation for the period under study in a 5×5 equi-spaced grid and use the "centroid" (see Section 5.1) of each bin as z_l . Figures 4(a) and 4(b) show climate bins for ACRU and LITU, respectively, along with the logarithm of the total number of plots observed in each bin, for each species during the period of study. We see that, for ACRU and, even more so, for LITU, there are climate bins in which the species were not observed. Also, for convenience in display, the bins are numbered from 1 to 25 as indicated.

For display purposes, we choose four climate bins that have at least 100 individuals in each of the years under study. Figure 5a shows the summary of the estimated per-plot intensities for ACRU for these grids for 2006. The corresponding observed empirical perplot intensity is overlaid. Figure 5b shows the same for year 2009. Figures 6a and 6b are similar to Figures 5a and 5b but for the species LITU. Two remarks emerge, neither unexpected in view of the severe sparsity. We evidently do better in some climate bins than others and we have a very large amount of uncertainty.



FIG. 3. Plot of true simulated $\tilde{\gamma}_{l,10}(x^*)$ (solid) for four covariate bins (see text for details). Overlaid are the pointwise 95% CI of the projected $\gamma_{l,10}(x^*)$ under the complete data set (dashed) and those under the sparse data set with 80% missingness per time point (dotted). The covariate levels are noted atop the figures. Note: the posterior medians are not displayed in Figure 3. Those figures are available upon request.

To further assess the goodness of fit, we plot the posterior summary of the per-plot estimated population size for nonempty climate bins (indexed as in Figure 4) in Figures 7a and 7b. Overlaid are the observed perplot population size in the corresponding climate grid. Generally, our prediction is successful but, again, our uncertainty is very high.

The posterior summaries of the parameters for both species are shown in Table 2. We assumed the upper bound for survival probability for both ACRU and LITU to be 0.9 and the estimates of Q_1 suggest a stronger density dependence for ACRU as compared to LITU. The recruitment rate (Δ) for ACRU is higher than that for LITU, explaining the relatively higher abundance of the former compared to the latter. The climate covariates do not seem to significantly impact the evolution of population size for either species. We attribute this, again, to the severe sparsity which leads to high uncertainty in these parameters, manifested by very wide credible intervals. Projection would proceed



FIG. 4. The climate bins along with the logarithm of the number of plots in which (a) ACRU were observed during the period of study (from 2005 to 2010) in each bin, and (b) LITU were observed during the period of study (from 2005 to 2010) in each bin. Also, the bins are indexed from 1 to 25.



FIG. 5a. Posterior mean (solid) and pointwise 95% CI (dashed) for the estimated per-plot intensity for ACRU for 2006. The observed empirical per-plot intensity is overlaid (dash-dotted). The temperature and precipitation corresponding to the grid centroid are noted atop.



FIG. 5b. Posterior mean (solid) and pointwise 95% CI (dashed) for the estimated per-plot intensity for ACRU for 2009. The observed empirical per-plot intensity is overlaid (dash-dotted). The temperature and precipitation corresponding to the grid centroid are noted atop.

at the plot level, assuming we know climate in the intervening, unobserved years.

6. A BRIEF SUMMARY AND FUTURE WORK

Only in recent work of Ghosh, Gelfand and Clark (2012) have IPM models been considered at the population scale and never have IPM models been considered at large regional scales or in the absence of data for consecutive time periods. Here we have presented a modeling approach to enable this in the context of an important demographic data set, the FIA data which samples plots, not individuals, roughly every five years, for the entire eastern half of the United States. After specifying our population level IPM, we have shown how to scale this IPM from plots to large regions and then we have shown how to approximately fit this scaled specification in the presence of the more

than 80% absence in the IPM data. We have illustrated the analysis for two species in the FIA data set.

Future work will see us investigating additional species in the FIA database (there are roughly 100 and many are not prevalent) to compare IPMs. We will also explore the possibility of building a joint IPM specification to allow for dependence between species. Such dependence could obviously affect both population size and diameter distribution. Scaling such joint models from plots to large geographic regions will add further challenge.

APPENDIX

The redistribution kernel K is specified as a parametric form. We assume K to be comprised of growth and recruitment with climate scaling and density de-



FIG. 6a. Posterior mean (solid) and pointwise 95% CI (dashed) for the estimated per-plot intensity for LITU for 2006. The observed empirical per-plot intensity is overlaid (dash-dotted). The temperature and precipitation corresponding to the grid centroid are noted atop.

pendence in the form

(10)
$$K_{t}(y, x; \mathbf{z}_{t}, \boldsymbol{\theta}, \gamma_{t, \cdot}) = (G_{t}(y, x; \mathbf{z}_{t}, \boldsymbol{\theta}, \gamma_{t, \cdot}) + R_{t}(y; \mathbf{z}_{t}, \boldsymbol{\theta}, \gamma_{t, \cdot}))e^{\mathbf{z}_{t}^{T}\boldsymbol{\beta}}.$$

In (10), the exponential term implies multiplicative scaling of climate effects regardless of x. It is introduced illustratively and to facilitate model identifiability and fitting; thus, the z_t 's are removed from the G and R terms. We have considered other forms for K. For instance, climate might drive growth, that is, be introduced in f while population size or growth could be introduced to drive survival. The flexibility in specifying K is attractive, but the more complex K is, the weaker the identifiability, the greater the sensitivity to prior specification, the more difficult the model fitting. The suggestion is that simple forms for the vital rates below, which determine K, may be more sensible.

In fact, the growth term $G_t(\cdot)$ is further decomposed as

$$G_t(y|x; \boldsymbol{\theta}, \gamma_{t,\cdot}) = q(x, \gamma_{t,\cdot}) f_t(y - x; \boldsymbol{\theta}).$$

Again, returning to density elements, we interpret $q(x, \gamma_{t,.}) f_t(y-x; \theta) \gamma_t(x) dx dy$ as the expected number of individuals in diameter interval (y, y + dy) at time t + 1 from survivors in diameter interval (x, x + dx) at time t. In particular, we assume f_t to be Gaussian density. Note that a translation-invariant f_t can be appropriate at the population level, though it would almost never be sensible at the individual level. In the sequel, again for convenience, we assume survival probability declines as a function of γ_t , due to resource limitation (but not as a function of diameter) and consider a logit form,

(11)
$$q(\gamma_{t,\cdot}) = \frac{Q_0 e^{-Q_1 \gamma_{t,\cdot}}}{1 + Q_0 e^{-Q_1 \gamma_{t,\cdot}}},$$



FIG. 6b. Posterior mean (solid) and pointwise 95% CI (dashed) for the estimated per-plot intensity for LITU for 2009. The observed empirical per-plot intensity is overlaid (dash-dotted). The temperature and precipitation corresponding to the bin centroid are noted atop.

where Q_0 and Q_1 (both > 0) are parameters that govern the rate of decay of the survival probability.

The recruitment term takes a form similar to the growth term,

$$R_t(y; \boldsymbol{\theta}, \gamma_{t,\cdot}) = \Delta(x, \gamma_{t,\cdot})g_t(y; \boldsymbol{\theta}).$$

With density elements, analogously, we interpret $\Delta(x, \gamma_{t,.})g_t(y; \theta)\gamma_t(x) dx dy$ as the expected number of recruited individuals in diameter interval (y, y + dy) at time t + 1 from individuals in diameter interval (x, x + dx) at time t. Usually, the terms on the right-hand side reflect flowering and seed production (see, e.g., Rees and Ellner, 2009). However, with trees, as in our FIA data set, seeds are not monitored. Hence, the recruitment simply describes the diameter intensity for new trees in year t + 1. Δ is the expected influx in year t + 1 and g_t is a diameter density on $\tilde{y} = y - L$ (since all new recruits to our point patterns are at least size L

in the year they arrive), which is assumed to be an exponential distribution translated to $[L, \infty)$. We assume influx declines with $\gamma_{t,.}$ due to reduced availability of resources and consider the form

(12)
$$\log \Delta(\gamma_{t,\cdot}) = \delta_0 - \delta_1 \gamma_{t,\cdot}$$

with δ_0 and δ_1 both nonnegative.

With the resulting kernel inserted into (10), integrating over *y*, we obtain

(13)
$$\gamma_{t+1,\cdot} = (q(\gamma_{t,\cdot}) + \Delta(\gamma_{t,\cdot}))e^{\mathbf{z}_t^T \boldsymbol{\beta}} \times \gamma_{t,\cdot},$$

which clarifies how the expected number of individuals changes from time t to time t + 1. Evidently, we can propagate (13) across t to learn about the behavior of population size over time.

Returning to (3), we approximate the stochastic integral with a Riemann sum. We divide the interval [L, U]into a fine grid consisting of *B* cells of equal length



FIG. 7a. Posterior mean (x) and 95% CI for the estimated per-plot abundance for ACRU for nonempty climate bins obtained for years 2007 through 2010. The observed per-plot abundance for the corresponding climate bins are overlaid (o). The bins are indexed on the x-axis following Figure 4.

with the centers given by x_j^* . We assume that the intensity is constant within each cell and that the centers, x_j^* , remain fixed across all of the time periods. The length and cell level intensity for cell *b* are denoted by *d* and $\lambda_t(b)$; b = 1, ..., B, respectively. Then the operational likelihood becomes

(14)
$$\prod_{t=1}^{T} \left[\exp\left(-\sum_{b=1}^{B} \lambda_t(b)d\right) \prod_{b=1}^{B} \left[\lambda_t(b)\right]^{n_{tb}} \right]$$

where n_{tb} is the number of points in cell b in year t.

As noted above, we assume $f_t(y - x; \mu_t, \sigma_t^2) = \phi(y - x; \mu_t, \sigma_t^2)$ and $g_t(y; \eta_t) = \eta_t e^{-\eta_t(y-L)}, y > L$. For the forms in (11) and (12), imposing priors

on $q(\gamma_{t,.})$ and $\Delta(\gamma_{t,.})$ requires specifying priors on Q_0, Q_1, δ_0 and δ_1 , respectively. We interpret $\frac{Q_0}{1+Q_0}$ as the survival probability when the population size tends to 0 and $\frac{\delta_0}{1+\delta_0}$ as the replacement rate when the population size tends to 0. We can roughly interpret Q_1 to be the global survival probability of the species and δ_1 to be the average rate of influx shown by that species.

 Q_0, Q_1, δ_0 and δ_1 are not well identified. In fact, from (13), the sum $q(\cdot) + \Delta(\cdot)$ is well identified but not its components. Estimation of $q(\gamma_{t,\cdot})$ and $\Delta(\gamma_{t,\cdot})$ requires using knowledge of the ecological processes driving the survival and influx for the population. According to the species, we assume a known upper



FIG. 7b. Posterior mean (x) and 95% CI for the estimated per-plot abundance for LITU for nonempty climate bins obtained for years 2007 through 2010. The observed per-plot abundance for the corresponding climate bins are overlaid (o). The bins are indexed on the x-axis following Figure 4.

 TABLE 2

 Posterior summary of model parameters for ACRU and for LITU.

 Posterior mean and 95% equal tail credible intervals provided

Parameters	Posterior summary for ACRU	Posterior summary for LITU
Q_1	0.08 (0.009, 0.19)	0.06 (0.0052, 0.099)
σ	0.38 (0.08, 0.49)	0.45 (0.28, 0.72)
δ	0.19 (0.002, 0.47)	0.16 (0.001, 0.47)
η	0.08 (0.018, 0.14)	0.07 (0.017, 0.15)
$\beta_{\text{intercept}}$	0.0746 (-5.16, 5.02)	1.96 (-5.90, 10.06)
β_{temp}	0.02 (-0.28, 0.30)	-0.16(-0.59, 0.24)
β_{precip}	0.0023 (-0.0019, 0.0059)	0.0014 (-0.0043, 0.0061)

bound of the survival and recruitment function which are achieved when $\gamma_{t,\cdot} = 0$. Solving these boundary conditions, we obtain the values of Q_0 and δ_0 and do not estimate them as part of model fitting. Q_1 and δ_1 , on the other hand, are estimated as a part of fitting using an additional constraint. Let $\rho_{t+1,t} = (N_{t+1} - N_t)/N_t$, t = 0, 1, ..., T - 1, where N_t is the total observed population size in year t. Then ρ_t denotes the relative change in population size in two consecutive years. To induce identifiability, we impose that $q(\gamma_{t,\cdot}) + \Delta(\gamma_{t,\cdot}) \in (1 - \max_t(\rho_{t+1,t}), 1 + \max_t(\rho_{t+1,t}))$. The priors on Q_1 and δ_1 are chosen such that this constraint is satisfied. The $\boldsymbol{\beta}$'s are well identified since they are regression coefficients associated with time varying covariates \mathbf{z}_t . Hence, we impose a vague Normal(0, 100) prior on each component of $\boldsymbol{\beta}$ independently.

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