**Spatial modeling of Audubon Christmas Bird Counts reveals fine-scale patterns and drivers of relative-abundance trends**

Timothy D. Meehan1†, Nicole L. Michel2, and Håvard Rue3

1 *National Audubon Society, Boulder, Colorado, USA*

2 *National Audubon Society, Portland, Oregon, USA*

3 *King Abdulla University of Science and Technology, Thuwal, Saudi Arabia*

† **E-mail:** tmeehan@audubon.org

**Abstract.** Bird counts by community volunteers provide valuable information about the conservation needs of many bird species. The statistical modeling techniques commonly used to analyze these counts provide robust, long-term population trend estimates from heterogeneous community science data at regional, national, and continental scales. Here we present a new modeling approach that increases the spatial resolution of trend estimates and reduces the computational burden of trend estimation, each by an order of magnitude. We demonstrate the approach with data for the American Robin (*Turdus migratorius*) from Audubon Christmas Bird Counts conducted between 1966 and 2017. We show that aggregate regional trend estimates from the proposed method aligned well with those from the current standard method, and that spatial variation in trends were associated with winter temperatures and human population densities as predicted by ecological energetics. This technique can provide reasonable large-scale trend estimates for users interested in general patterns, while also providing higher-resolution estimates for examining correlates of abundance trends at finer spatial scales, which is a prerequisite for tailoring management plans to local conditions.

**Key words:** Audubon Christmas Bird Count; Bayesian hierarchical model; conditional autoregressive model; North American Breeding Bird Survey; population trends; range shifts; spatially varying coefficients model.

**Introduction**

Volunteers with the Audubon Christmas Bird Count (CBC) have been counting wintering birds across North America every year for the last 118 years (Dunn et al. 2005, Soykan et al. 2016). Population trends derived from CBC data, along with those derived from other large-scale monitoring programs like the North American Breeding Bird Survey (BBS, Robbins et al. 1989, Sauer et al. 2017), provide valuable information for understanding the conservation needs of North American bird species (Dickinson et al. 2010, Hochachka et al. 2012). For example, CBC and BBS trends are used to evaluate population change rates and population half-lives for many of the 448 bird species included in the Partners in Flight Landbird Conservation Plan (Rosenberg et al. 2016, 2017). CBC and BBS trends are also used by BirdLife International (http://datazone.birdlife.org/species/search) to make status recommendations to the International Union for Conservation of Nature, creators of the Red List of Threatened Species (https://www.iucnredlist.org/), and by the North American Bird Conservation Initiative (NABCI) to produce their State of the Birds conservation vulnerability assessment for all 1,154 native North American bird species (NABCI 2016). CBC trends are especially useful for species who are otherwise not monitored in their remote northern breeding locations, particularly those that breed in the poorly-surveyed boreal forest or Arctic tundra (Niven et al. 2004).

The current, standard approach for generating trends from CBC data (Link et al. 2006, Soykan et al. 2016) was derived from methods originally developed for BBS data (Link and Sauer 2002, Sauer and Link 2011). The general approach is to assign counts in Canada and the US to one of up to 169 spatial strata, which are intersections of US states or Canadian provinces, and Bird Conservation Regions (BCR, Sauer et al. 2003). Then, treating each stratum as independent, a non-linear function is used to correct for the effect of observer effort on counts, while simultaneous effects are estimated for the impact of count circle, year, and stratum by year (Link et al. 2006, Soykan et al. 2016). These parameter estimates are used to derive a relative abundance index per stratum and year, and those annual indices are used to compute annual percent change per stratum across defined time periods (Link and Sauer 2002, Sauer and Link 2011).

The standard CBC analysis provides robust long-term trend estimates from heterogeneous community science data across large spatial scales. By pooling count circles per stratum, this approach deals with the issue of count circles haphazardly becoming active or inactive over the time series (Sauer and Link 2011, Soykan et al. 2016). Additionally, pooling produces a sufficiently large sample of counts to generate a reasonably robust count-effort correction function (Link and Sauer 1999), which is critical given the wide variation in count effort among count circles (Bock and Root 1981, Dunn et al. 2005). This approach produces a relative abundance index per year and stratum, which can be used to explore variation around long-term log-linear trends, and can be summed across larger hierarchically-nested strata, such as states, provinces, BCRs, or Landscape Conservation Cooperative (LCC) regions, and used to estimate change in relative abundance at larger spatial scales. Producing annual abundance indices also permits summarizing abundance change between any desired pair of time points. Finally, the simplicity of the standard model enables a flexible and robust computational process, suitable for analysis of hundreds of species that vary enormously in their ubiquity, abundance, and population dynamics.

While the current approach produces trends that are useful for understanding population status of birds at regional to continental scales, the approach has some limitations. As implemented, it is a computationally intensive process, especially for wide-ranging species. This is due to the use of Markov chain Monte Carlo (MCMC) to estimate model parameters for relative abundance. From an ecological perspective, the coarse resolution of standard population trend analyses limits their ability to provide inference about local variation, processes, and drivers. While trends can be scaled up to larger spatial units, they cannot be scaled down to smaller ones. The analytical stratum is the finest level of resolution, which limits the extent to which variation in trends can be attributed to fine-scale processes such as change in local land cover or weather patterns (Thogmartin et al. 2004, Bled et al. 2013), and limits opportunities for tailoring conservation plans to local conditions (Ethier and Nudds 2015, Ethier et al. 2017). Abundance, distribution, and population trends of birds are affected by land cover, climate, topography, and other environmental conditions and processes at a variety of spatial scales ranging from local (e.g., home range or smaller) to landscape (Thogmartin et al. 2004, 2006; Zhang et al. 2012). Summarizing and evaluating correlates of trends at the scale of traditional strata may miss important local variation in abundance, distribution, and the environmental conditions affecting them.

Moreover, the current approach does not take full account or advantage of spatial relationships among counts. Modeling this structure would facilitate borrowing information across spatial boundaries, allowing more robust trend estimates in places where data are sparse (Waller and Gotway 2004, Banerjee et al. 2014, Blangiardo and Cameletti 2015) and enabling trend estimation at finer spatial scales (Thogmartin et al. 2004, Bled et al. 2013). Accounting for the spatial dependence across count sites also reduces the amount of spatial autocorrelation in model residuals, which leads to more reliable inference about trend estimates (Dormann et al. 2007, Beale et al. 2010).

Previous work by Thogmartin et al. (2004), Bled et al. (2013), and Smith et al. (2015), among others, offered spatially explicit variations of the standard trend analysis approach for community science data. These works were focused on analysis of BBS data, but their approaches are easily applied to analysis of CBC data. Instead of using the standard strata described above (Smith et al. 2015), Thogmartin et al. (2004) assigned count sites to irregular polygons, created by tessellation of BBS route locations. Bled et al. (2013) assigned routes to cells on a regular grid, with one-degree latitude and longitude spacing. All three studies utilized spatially structured random intercepts for relative abundance per polygon, grid cell, or stratum. Thogmartin et al. (2004) included an effect of year per polygon, but that effect did not incorporate spatial structure. Bled et al. (2013) and Smith et al. (2015) estimated relative abundances per year, and then trends were generated as derived parameters, as done in the standard analysis.

Here, we present a new approach for calculating temporal trends in relative abundance, one that takes advantage of the considerable spatial structure in CBC data. This approach borrows components from previous ones, incorporates new components that prioritize robust trend estimation at finer spatial scales, and employs a simplified and computationally efficient workflow. Similar to Bled et al. (2013), we assigned CBC count sites to cells on a uniform grid covering North America. In contrast to previous work, effort and year effects were modeled as random slopes with spatial structure, following a spatially varying coefficient (SVC) approach (Gelfand et al. 2003, Finley 2011, Congdon 2014). Finally, unlike prior studies using MCMC, we used integrated nested Laplace approximation (INLA) to estimate Bayesian posteriors for model parameters (Rue et al. 2009, Martins et al. 2013, Lindgren and Rue 2015, Blangiardo and Cameletti 2015, Rue et al. 2017, Bakka et al. 2018, Krainski et al. 2018), which led to a dramatic decrease in computing time. Our objective in developing this technique was to estimate long-term population trends at an appropriate spatial scale for evaluating fine-scale population processes and ecological drivers to inform conservation. The four goals of this report were to (*i*) describe an SVC approach for calculating trends in CBC data, (*ii*) employ the approach using data for the American Robin, (*iii*) compare trend results derived from the SVC approach to aggregate results derived from standard methods, and (*iv*) demonstrate use of fine-scaled trend results through a simple post-hoc analysis exploring correlations between precomputed SVC trends and potential energetic drivers related to climate and winter food resources.

**Materials and Methods**

***Christmas Bird Count***

We demonstrate use of the SVC approach using data from the CBC, a volunteer bird monitoring program begun in 1900. Since its inception, the program has grown tremendously and now includes more than 2,400 count sites and involves more than 70,000 volunteers (Soykan et al. 2016). The program extends across North and South America, but the majority of counts have been conducted in the US and Canada. The program has produced counts extending 118 years, but standard trend analyses are conducted using counts conducted after 1965, as protocols and count efforts have been most consistent since then. An individual CBC occurs over a 24-hour period, once per year, during late December or early January. During a count, volunteers record birds seen or heard within a circular study area with a 24.1 km diameter. The area covered and the amount of time spent counting birds in a given circle varies considerably across years and count locations, so annual effort metrics are recorded and used to standardize bird counts. More details about the CBC can be found in Bock and Root (1981) and Dunn et al. (2005), who describe the program and discuss the potential uses and limitations of CBC data.

***Statistical model***

We modeled CBC count data, , for grid cell *i* encompassing count circle *k* during year *t*, as a random variable from a negative binomial distribution. Expected values for counts per grid cell, , were assumed to be a function of spatially structured grid-cell, count-effort, and year effects, plus unstructured variation among count circles. The linear predictor for took the form:

. (Eq. 1)

Parameters were modelled as cell-specific random intercepts with an intrinsic conditional autoregressive (CAR) structure (Besag et al. 1991). With this structure, values came from a normal distribution, with a conditional mean related to the average of adjacent cells, and with conditional variance proportional to the variance across adjacent cells and inversely proportional to the number of adjacent cells. Spatial structure was incorporated into to allow for information about relative abundance to be shared across neighboring cells, and to reduce the spatial autocorrelation among model residuals that occurs when this spatial structure is ignored (Dormann et al. 2007, Finley 2011).

Parameters were modeled as spatially structured, cell-specific, random slope coefficients for the local effects of effort . These spatially varying coefficients (Gelfand et al. 2003, Banerjee et al. 2014, Congdon 2014) were also modelled with a CAR structure (Besag et al. 1991). Slopes were drawn from a normal distribution with a conditional mean related to the average of adjacent cells, and with conditional variance proportional to the variance across adjacent cells and inversely proportional to the number of adjacent cells. Spatial structure was incorporated into to allow for information about the effort effect to be shared across neighboring cells, and to accommodate a potential lack of stationarity (Finley 2011) in the effort effect. Effort was represented by , the number of party hours expended during a count, where a party hour was the count effort of one party of unspecified size for one hour. Pairing log-transformed counts with log-transformed effort in the linear predictor yielded a power function for effort correction, a flexible mathematical form that accommodated a decreasing, linear, or increasing impact of effort on expected counts (Butcher and McCulloch 1990, Link and Sauer 1999).

Parameters were modeled as spatially structured, cell-specific, random slope coefficients for the year effect. Spatially varying coefficients (Gelfand et al. 2003, Banerjee et al. 2014, Congdon 2014) were also modeled with CAR structure (Besag et al. 1991), where values came from a normal distribution, with conditional means and variances as described above. Spatial structure was incorporated into to allow for information about the year effect to be shared across neighboring cells, and to allow the year effect to vary across the cells in the study region. Year, represented by *T*, was transformed before analysis such that max(*T*) = 0, and each preceding year took an increasingly-negative integer value. Given the scaling of effort and year variables, could be interpreted as a cell-specific expected count given one party hour of effort during the final year in the time series.

The final term in the model, , was an exchangeable random effect that accounted for variation in relative abundance among circles, possibly due to differences in habitat conditions or observer experience (Soykan et al. 2016). Note that the model did not include a normally-distributed, observation-level random effect to deal with overdispersed Poisson counts, i.e., and , as is done for the standard approach (Sauer and Link 2011, Soykan et al. 2016). Rather, we used a negative binomial count distribution for *y,* i.e., and (Linden and Mantyniemi 2011). These two approaches are expected to yield similar outcomes. However, as implemented in R-INLA, the latter approach returns a single dispersion estimate while foregoing estimation of individual observation effects, which reduces computing time and the size of posterior samples.

***Case study data***

We developed a case study, using data for the American Robin (*Turdus migratorius*) from CBCs conducted across the continental US and Canada between 1966 and 2017, to demonstrate the SVC modeling approach, compare results with those from the standard approach, and illustrate a straightforward analysis of fine-scaled trends. Before modeling the CBC data, extreme outliers (> 3 SD from the mean, after log transformation) in counts and effort were removed. After filtering, there were 36,650,191 American Robin observations from 78,140 counts from 3,195 count circles over 52 years available for modeling.

Locations of the 3,195 unique count circles were mapped using the North American Albers Equal Area Conic projection (EPSG 102008, https://epsg.io/102008) and assigned to 880 cells on a grid divided along 100 km increments in latitude and longitude (Fig. 1A). Grid cells formed a continuous lattice within a non-convex polygon created using circle locations. The number of count circles per grid cell varied from 0 to 20, and averaged 2.43 (Fig. 1A). The number of neighbors for a given grid cell ranged from 1 to 8, and averaged 7.48. Note that grid cells with zero counts were retained during model estimation to preserve the spatial relationships between counts. However, before analyzing resulting trend estimates, cells with no observed counts were removed from the dataset, as we were not interested in interpolating trends for grid cells without CBC sites.

***Model computation***

The SVC model described above was analyzed within a Bayesian framework using the R-INLA package (version 18.10.09, Martins et al. 2013, Lindgren and Rue 2015, Blangiardo and Cameletti 2015, Rue et al. 2017, Bakka et al. 2018, Krainski et al. 2018) for R statistical computing software (version 3.3.1, R Core Team 2016). For parameters , , and , with CAR structure, precision matrices were scaled such that the geometric mean of marginal variances was equal to one (Sørbye and Rue 2014, Riebler et al. 2016, Freni-Sterrantino et al. 2018), and priors for precision parameters were penalized complexity (PC) priors, with parameter values and (Simpson et al. 2017). Precision for the zero-centered, exchangeable, random circle effect, , was also assigned a PC prior with parameter values and (Simpson et al. 2017). The overdispersion term for the negative binomial count distribution, , was assigned a PC prior with parameter value *l* = 7. Very simply, as implemented here, PC priors were weakly informative priors with an innate tendency to shrink structured and unstructured random effects toward zero in the absence of a strong signal. Readers are referred to Simpson et al. (2017) for the details of, and rationale behind, PC priors, as well as the default structures and parameter values of PC priors used in the R-INLA package.

Along with parameter estimates, R-INLA produced values to evaluate individual model fit (Czado et al. 2009) and compare different models to one another (Gneiting and Raftery 2007, Link et al. 2017): cross-validation probability integral transform (PIT, Dawid 1984) and conditional predictive ordinate (CPO, Pettit 1990). For this application, we were not comparing multiple models. However, we extracted PIT values and visually inspected their histogram, as an approximate uniform distribution is expected for a model that fits the data reasonably well (Czado et al. 2009, Held et al. 2010).

Following model analysis, posterior medians and symmetric 95% credible intervals were computed per cell for , , and . Credible interval widths, representing estimate uncertainty, were computed by subtracting the lower credible limit from the upper credible limit per cell. Posterior summaries were then mapped to visualize spatial variation in abundance indices, effort effects, and relative-abundance trends.

***Trend comparisons***

It is common, following CBC and BBS analyses, to aggregate trend information to larger scales that might be of interest to resource managers designing and implementing policies across states, provinces, BCRs, LCCs, or nations (Sauer et al. 2003, Sauer and Link 2011, Soykan et al. 2016). After analysis of the SVC model for the American Robin, we aggregated 100-km results to the BCR level to compare them to those produced using standard CBC analysis methods (i.e., the implementation of hierarchical models in JAGS; Soykan et al. 2016). BCR-level trends were precomputed and are available for download from the National Audubon Society website (Meehan et al. 2018). SVC trends were aggregated for each BCR by averaging trends for all equal-area grid cells where the cell centroid fell within the BCR. We evaluated the uncertainty around SVC trend estimates by comparing credible interval widths for cells within a BCR to those calculated for a BCR using the standard approach. We note that full posterior distributions for aggregate SVC trend estimates could have been computed by averaging random draws from posteriors for cells within a BCR, or by using R-INLA functions for creating linear combinations (Blangiardo and Cameletti 2015). Further, those averages could have been weighted by the amount of area where the cells and BCR overlap (Bled et al. 2013). We did not employ these methods here because our main focus was on fine scale trends and only qualitative comparison with standard results.

***Correlates of American Robin trends***

One of the greatest potential benefits of the SVC method is the ability to produce fine-scale population trends, enabling investigation of both fine- and coarse-scale ecological processes and population trend drivers. To highlight this potential, we conducted a simple demonstration analysis of fine-scaled American Robin trends. Specifically, we explored correlations between precomputed SVC abundance trends and three covariates: latitude, historic minimum winter temperature, and historic human population density. We conducted this correlation analysis post-hoc on precomputed trends, rather than relating covariates to the original count data, because we believe that analysis of precomputed trends, while possibly less elegant, is a simpler process and thus more likely to be adopted by the larger scientific community.

We expected that, in the Northern Hemisphere, latitude would be positively related to SVC trends, as several landbird species have demonstrably shifted their ranges towards the Earth’s poles (Thomas and Lennon 1999, La Sorte and Thompson 2007, Huang et al. 2017). A northward range shift would logically lead to negative trends in the southern portion of their range and positive trends in the northern portion of their range. One putative mechanism for northward range shifts in wintering birds is the increase in minimum winter temperatures across the continent over the past 50 years (Zuckerberg et al. 2011, Princé and Zuckerberg 2015), which makes the energetics of wintering in the north more favorable (Root 1988, Meehan et al. 2004). With this mechanism in mind, we expected that, for a given latitude, residual trends would be negatively related to historic minimum winter temperature, as birds began wintering in historically colder locations further from coasts and higher in elevation. Finally, we tested the hypothesis that winter bird distribution and abundance is also governed by resource availability (Dunning and Brown 1982, Meehan et al. 2004). We expected that, after accounting for latitude and minimum winter temperature, SVC trends would be positively related to historic human population density (Robb et al. 2008, Zuckerberg et al. 2011), as American Robins are well known to prosper in human-dominated landscapes, replete with mown grass and fruiting ornamentals (Beissinger and Osborne 1982, Marzluff 1997).

For this analysis, values were related to the latitude of the centroid for a given 100 km grid cell. A single, historical-average minimum December and January temperature was calculated per cell between 1965 and 2017, using the CRU TS v4.01 dataset (Harris et al. 2014). A single, historic-average human population size between 1970 and 2015 was calculated per cell using Global Population Count Grid Time Series Estimates (CIESIN 2017) for 1970 and the GPWv4 dataset (CIESIN 2016) for 2015. Cell-specific values were modeled with a latent Gaussian model, where , , and *Li*, *Ti*, and *Pi* represented cell-specific latitude, historic-average minimum winter temperature, and historic-average human population size variables, respectively. Note the transformation of *Pi*, which was highly right-skewed. The term represented cell-specific random intercepts with CAR structure (Besag et al. 1991), where the precision matrix was scaled such that the geometric mean of marginal variances was equal to one (Sørbye and Rue 2014, Riebler et al. 2016, Freni-Sterrantino et al. 2018). was added to the model to account for spatial structure in the model residuals (Dormann et al. 2007, Beale et al. 2010). *β*0, *β*1, *β*2, and *β*3 were given normal priors with mean of zero and precision equal to 0.001. The prior for precision associated with the CAR term was a PC prior (Simpson et al. 2017) with parameter values and .

**Results**

On average, 4.28 robins were counted per party hour in 2017, but that number varied by several orders of magnitude across the species range, from 0.01 to 73.50. A map of posterior median values illustrated that the species was most abundant in the central part of their range, and was least abundant along the northern and southern margins of their range (Fig. 1B). Calculation of the SVC model using R-INLA took approximately 10 minutes to complete. Inspection of the PIT histogram indicated satisfactory model fit.

Posterior median values for , the power-law exponent for the relationship between effort and counts, varied from 0.28 to 1.44, with a median value of 0.81. The 95% credible intervals for values indicated that 80% of estimates were not significantly different from 1, while all were significantly greater than 0. Estimates not significantly different from 1 indicated a positive linear relationship between effort and counts. Values significantly greater than 0 and less than 1 also indicated a positive relationship between effort and counts, but one with diminishing returns for additional count effort. A map of posterior median values highlighted the spatial structure in the effort effect (Fig. 1C). Locations with posterior medians well below 1 were frequently locations with relatively low abundance indices (Fig. 1B), suggesting that the majority of American Robins in a count circle could be counted with relatively low effort. Locations with posterior medians closer to 1 were frequently locations with relatively high abundance indices (Fig. 1B), suggesting an endless supply of American Robins for CBC volunteers to count. The two parameters, and , were correlated across space, with a rank correlation coefficient of 0.26.

Posterior median values for , the temporal trend from 1966 through 2017, when transformed to annual percent change, varied from -11.80% to 13.63% per year, with a median value of 2.63% per year (Fig. 1D). The 95% credible intervals for values indicated that 8% of estimates were significantly lower than 0, while 44% were significantly greater than 0 (Fig. 1E). A map of posterior median values showed that relative abundance trends had strong spatial structure. Credible intervals for values (Fig. 1F) were used to illustrate where trends were significantly negative or positive, showing that relative abundance during winter has generally decreased in the southern and coastal parts of their range and increased in the northern and interior parts of their range. The parameters and were significantly correlated across space, with a rank correlation coefficient of -0.16, indicating that the strongest trends were occurring at the margins of the geographic range where relative abundance was lowest.

The posterior median estimate for , the dispersion parameter, was , considerably larger than 1, highlighting overdispersion in American Robin counts relative to a Poisson distribution. Credible intervals for precision estimates for the random effects showed that all were important for explaining variation in the count data. When precision values were converted to a standard deviation scale, the random effects were ranked (cell-specific intercept, SD = 1.58), (circle-specific intercept, 1.05), (cell-specific effort effect, 0.25), and (cell-specific year effect, 0.04), in terms of the amount of variation explained.

Figure 2 shows the median of posterior median SVC trends across cells per BCR (Fig. 2A), along with the posterior median trend for each BCR from the standard analysis (Fig. 2B). Side-by-side visual comparison of these maps showed that aggregate trends were similar, regardless of method. The SVC approach gave a median trend of 2.11% across all BCRs, while the posterior median trend for the standard approach was 1.95% across all BCRs. Within BCRs, the trend direction was consistent across the two methods in 28 of 32 BCRs. The rank correlation between BCR trends generated by the two methods was 0.88. Regarding differences, trends derived from the SVC approach changed more smoothly across the continent, as would be expected using a spatial statistical model. Also, the range of posterior median SVC trends (-4.58% to 9.16%) was slightly less than that for standard trends (-7.96% to 14.26%), especially near geographic range boundaries, as would be expected given the sharing of information across space.

Figure 3 compares the credible interval widths for SVC trends per grid cell (Fig. 3A) with those from the standard approach for aggregate BCR estimates (Fig. 3B). When compared to the standard approach, some SVC grid cells within a BCR, ones in information-rich neighborhoods (Fig.1A), had SVC trend estimates with remarkably narrow confidence intervals (Fig. 3C, SVC minimum). Other grid cells, ones in information-poor neighborhoods (Fig.1A), had trend estimates with relatively broad confidence intervals (Fig. 3C, SVC maximum). On average, however, precision of estimates per BCR were similar, regardless of method, if not slightly higher using the SVC approach (Fig. 3C, SVC median).

As predicted, trends were positively related to degrees north latitude, with trends being negative at low latitudes and positive at high latitudes (Fig. 4A), suggesting a poleward shift in winter range for the American Robin. After accounting for latitude, there was a clear negative relationship between historic average minimum temperature and SVC trends (Fig. 4B), indicating that robins were becoming more abundant in historically colder areas at higher elevations and further from coasts. After accounting for latitude and historic minimum winter temperature, SVC trends were positively related to the number of people historically residing in a grid cell, which we use as a proxy for resource availability for this anthropophilic species. Posterior median standardized coefficients from this model indicated that latitude (2.22) was the strongest predictor of trends, followed by historic minimum temperature (-0.89) and historic human population size (0.35). We used posterior medians for *β*0, *β*1, *β*2, and *β*3 to compute a predicted tau value, and found that the rank correlation coefficient for the association between input tau values and predicted tau values was 0.62.

**Discussion**

This analysis demonstrated use of Bayesian SVC models to estimate fine-scaled, long-term relative abundance trends along with relevant environmental drivers from community science data collected across North America. The inclusion of spatially structured random slope terms allowed for robust estimation of effort effects and relative abundance trends at spatial resolutions much finer than the standard analysis approach. Fine scaled trend estimates for the American Robin, when aggregated to the BCR level, were very similar to those produced by the standard analysis approach. The use of INLA for model analysis resulted in dramatically reduced computing time compared to the standard analysis approach. When we explored correlates of precomputed American Robin SVC trends, we found that trends were related to latitude, historic winter temperature, and historic human population size as expected from the literature on factors governing the distribution and abundance of wintering birds.

To put resolution gains into context, consider that a CBC circle has a radius of approximately 12 km and an area of 452 km2. A 100 km grid cell, covering 10,000 km2, is approximate 22 times larger than a CBC circle. In comparison, the average analytical stratum has an area of 104,378 km2, approximately 231 times the area of a CBC circle. Thus, the SVC approach brought an order of magnitude increase in spatial resolution when compared to the standard approach. This increased resolution facilitated finer-scaled investigations into the drivers of winter bird trends (Thogmartin et al. 2004, Bled et al. 2013, Smith et al. 2015). Understanding the fine-scale drivers of trends creates opportunities to tailor conservation plans to local conditions (Ethier and Nudds 2015, Ethier et al. 2017).

Estimating trends at relatively high resolution was made possible by adopting spatial-statistical techniques designed to borrow information across neighboring regions (Thogmartin et al. 2004, Bled et al. 2013, Smith et al. 2015). Employing spatial techniques also had implications for uncertainty in trend estimates. In the standard analysis, the uncertainty in a trend estimate depended upon the variation in trends across the circles within a stratum and the number of circles in a stratum. In the SVC analysis, uncertainty depended upon those same two factors, but also depended upon those characteristics in the neighborhood of a grid cell. The consequences of this difference are demonstrated in Figure 3. In regions with many CBC circles (e.g., Piedmont BCR), SVC methods produced trend estimates with relatively low uncertainty (minimum credible interval width of 1.48%) compared to the standard method (3.40%), due to the density of information. Similar to Bled et al. (2013), we found that precision of SVC estimates also tended to be relatively high in regions at the edge of a species range where there were few counts (e.g., Boreal Softwood Shield BCR, maximum interval width of 12.02%) when compared to the standard approach (19.14%), due to borrowing of information across neighboring cells that crossed regional boundaries. In other parts of the continent with fewer, more isolated CBC circles (e.g., Southern Rockies Colorado Plateau BCR), the SVC methods produced trend estimates with relatively high uncertainty (minimum interval width of 3.47%) compared to the standard method (2.80%). It is not entirely clear if the small intervals of the standard approach are justified in this context. If the relatively few and far-between circles that fall within those large BCRs can be considered representative samples of that larger area, then estimates with high precision are reasonable, and certainly preferred. If it cannot be assumed that those circles are representative of the larger area, then estimating trends for smaller areas, in neighborhoods with more information, and basing uncertainty estimates on the amount of local information, seems more appropriate. Critical evaluation of this representative-sample assumption is particularly important when analyzing data from the CBC, because count-site selection is not based on sampling design principles, and count locations tend to be biased toward areas of high human population density and areas of high bird density and diversity (Drennan 1981, Dunn et al. 2005).

On a typical laptop computer, SVC model analysis using R-INLA took roughly 10 minutes for full Bayesian results. The standard approach, which employs MCMC, took approximately 10 hours for full Bayesian results on the same hardware. Had spatial statistical models been analyzed using MCMC, processing times would have been much longer. The difference in computing time was due to R-INLA producing highly accurate approximations of Bayesian posteriors, orders of magnitude faster than MCMC (Rue et al. 2009, 2017). The obvious benefit of shorter processing times is that, for a given set of computing resources, more time periods, more distinct model forms (e.g., Link and Sauer 2016), or more species can be evaluated. Even small differences in computing time add up when analyzing counts from tens of years, for hundreds of species, across thousands of count sites.

There were, as there usually are, tradeoffs for rapid model analysis. Specifically, R-INLA is an analysis option whenever a statistical model can be expressed as a latent Gaussian model (Rue et al 2009, Blangiardo and Cameletti 2013, Rue et al. 2017). This option was possible for the model used in this analysis. However, this would not have been possible had we chosen to use the effort-correction function developed by Link and Sauer (1999, 2006) and used in the standard analysis (Soykan et al. 2016). Here, we used a single-parameter, power-law function for effort correction because it could fit positive, negative, linear, increasing, and decreasing relationships (Butcher and McCulloch 1990) and was easily incorporated into a latent Gaussian model. In contrast, the effort-correction function used for the standard approach is a two-parameter nonlinear function, which is more flexible and so will better-fit relationships that come to a rapid asymptote. Ideally, we would have tools for rapid analysis of spatial statistical models that incorporate the standard effort-correction function. In this choose-two situation, we erred towards rapid analysis of a spatial model with the simpler effort-correction function, because it allowed for more robust, if occasionally slightly biased (Link and Sauer 1999), estimates of the effort effect in regions where information was sparse. Robust estimates of effort effects are particularly critical when generating trends from CBC data, as count effort varies widely across time and space (Bock and Root 1981, Butcher et al. 1990, Dunn et al. 2005).

We note that the SVC approach described here differed from the standard approach in that it was optimized, specifically, for computing long-term, log-linear trends in relative abundance at fine spatial scales. The emphasis on long-term, log-linear trends was motivated by requests from resource managers, who desire simple summary statistics that reflect overall population status for many species (Rosenberg et al. 2016, 2017). The emphasis on fine spatial resolution was motivated by requests from researchers wishing to conduct research at relatively fine spatial scales and by Audubon Christmas Bird Count volunteers desiring to learn how bird numbers have changed over the years in their local area. Given these two emphases, we did not incorporate additional model terms necessary for creating annual abundance indices. These indices are critical for those who wish to look beyond single, long-term trends, at detailed time series that give more information about the nature of abundance changes (Sauer and Link 2002, 2011). Creating these annual indices is done by adding an additional random effect per cell and year, and combining these effects with and . Adding this effect to the SVC model is easily done in R-INLA. This effect could be specified as exchangeable, or have spatial or temporal structure. For this dataset, preliminary trials showed that adding an exchangeable effect to the model increased computing time to approximately 1 hour. We did not explore this model variant in depth because producing annual abundance indices was not a primary goal of this effort.

When we applied the SVC approach to data from the American Robin, we learned that aggregate trends resulting from the SVC and standard approaches were similar in direction and magnitude. Precision at aggregate levels was generally similar, if not a bit lower with the SVC approach, due to different assumptions about how precision should, or should not, be related to the spatial distribution of counts. Spatial variation in SVC trends for the American Robin were also consistent with expectations from the literature on factors governing winter bird distribution and abundance (Root 1988, Meehan et al. 2004, La Sorte and Thompson 2007, Zuckerberg et al. 2011). Specifically, trends were negative at low latitudes, positive at high latitudes and, for a given latitude, were more positive in historically cold regions and in areas with higher human population density. While it is likely that latitudinal effects would have been found using aggregate trends from the standard analysis (Fig. 2), it is less likely that residual winter temperature and human population density effects would have been detected. This is because winter temperatures and human population sizes both tend to vary at spatial scales considerably finer than analytical strata, states, provinces or BCRs. Future studies could explore relationships between SVC trends and other potential variables that can be summarized at comparable spatial scales, such as forest management practices, agricultural intensification, and other factors related to land use and land cover.

Dunn et al. (2005) suggested that analyses of CBC data by the broader scientific community would increase in number and quality as preprocessed, effort-corrected trends were made available to the public, as they have been for the BBS results (Sauer et al. 2003). Work by Soykan et al. (2016) marks the initial outcome of an effort by Audubon to make this happen, and standard trend analysis results have recently become publically available via the internet for over 500 species (https://www.audubon.org/conservation/where-have-all-birds-gone). With further fine tuning, it is possible that SVC trends could also become publically available for researchers interested in conducting finer-scaled analyses of preprocessed, effort-corrected trends and their environmental drivers.

**Acknowledgments**

We thank the thousands of volunteers who participate in the Audubon Christmas Bird Count, as well as the two anonymous reviewers who helped improve this manuscript.

**Literature Cited**

Bakka, H., H. Rue, G. A. Fuglstad, A. Riebler, D. Bolin, J. Illian, E. T. Krainski, D. P. Simpson, and F. K. Lindgren. 2018. Spatial modeling with R-INLA: a review. WIREs Computational Statistics e1443.

Banerjee, S., B. P. Carlin, and A. E. Gelfand. 2014. Hierarchical modeling and analysis for spatial data. CRC Press, Boca Raton, Florida, USA.

Beale, C. M., J. J. Lennon, J. M. Yearsley, M. J. Brewer, and D. A. Elston. 2010. Regression analysis of spatial data. Ecology Letters 13:246–264.

Beissinger, S. R., and D. R. Osborne. 1982. Effects of urbanization on avian community organization. Condor 84:75–83.

Besag, J., J. York, and A. Molliè. 1991. Bayesian image restoration, with two applications in spatial statistics. Annals of the Institute of Statistical Mathematics 43:1–59.

Blangiardo, M., and M. Cameletti. 2015. Spatial and spatio-temporal Bayesian models with R-INLA. John Wiley and Sons, New York, New York, USA.

Bled, F., J. R. Sauer, K. Pardieck, P. Doherty, and J. A. Royle. 2013. Modeling trends from North American Breeding Bird Survey data: a spatially explicit approach. PLoS ONE 8:e81867.

Bock, C. E., and T. L. Root. 1981. The Christmas Bird Count and avian ecology. Studies in Avian Biology 6:17–23.

Butcher, G. S., M. R. Fuller, L. S. McAllister, and P. H. Geissler. 1990. An evaluation of the Christmas Bird Count for monitoring population trends of selected species. Wildlife Society Bulletin 18:129–134.

Butcher, G. S., and C. E. McCulloch. 1990. The influence of observer effort on the number of individual birds recorded on Christmas Bird Counts. Pages 120–129 *in* J. R. Sauer and S. Droege, editors. Survey designs and statistical methods for the estimation of avian population trends. Biological Report 90. US Fish and Wildlife Service, Washington, DC, USA.

CIESIN [Center for International Earth Science Information Network]. 2016. Gridded population of the world, version 4, GPWv4, population count. <http://dx.doi.org/10.7927/H4BG2KXS>

CIESIN [Center for International Earth Science Information Network]. 2017. Global population count grid time series estimates, version 1, 1970-2000. <https://doi.org/10.7927/H4CC0XNV>

Congdon, P. 2014. Applied Bayesian modelling. John Wiley and Sons, New York, New York, USA.

Czado, C., T. Gneiting, and L. Held. 2009. Predictive model assessment for count data. Biometrics 65:1254–1261.

Dawid, A. P. 1984. Statistical theory: the prequential approach. Journal of the Royal Statistical Society, Series A 147:278–292.

Dickinson, J. L., B. Zuckerberg, and D. N. Bonter. 2010. Citizen science as an ecological research tool: challenges and benefits. Annual Review of Ecology, Evolution, and Systematics 41:149–172.

Dormann, C. F., J. M. McPherson, M. B. Araujo, R. Bivand, J. Bolliger, G. Carl, R. G. Davies, A. Hirzel, W. Jetz, W. D. Kissling, I. Kuhn, R. Ohlemüller, P. R. Peres‐Neto, B. Reineking, B. Schröder, F. M. Schurr, R. Wilson. 2007. Methods to account for spatial autocorrelation in the analysis of species distributional data: a review. Ecography 30:609–628.

Drennan, S. R. 1981. The Christmas Bird Count: an overlooked and underused sample. Studies in Avian Biology 6:24–29.

Dunn, E. H., C. M. Francis, P. J. Blancher, S. R. Drennan, M. A. Howe, D. Lepage, C. S. Robbins, K. V. Rosenberg, J. R. Sauer, and K. G. Smith. 2005. Enhancing the scientific value of the Christmas Bird Count. Auk 122:338–346.

Dunning, J. B., and J. H. Brown. 1982. Summer rainfall and winter sparrow densities: a test of the food limitation hypothesis. Auk 99:123–129.

Ethier, D. M., N. Koper, and T. D. Nudds. 2017. Spatiotemporal variation in mechanisms driving regional-scale population dynamics of a threatened grassland bird. Ecology and Evolution 7:4152–4162.

Ethier, D. M., and T. D. Nudds. 2015. Scalar considerations in population trend estimates: implications for recovery strategy planning for species of conservation concern. Condor 117:545–559.

Finley, A. O. 2011. Comparing spatially varying coefficients models for analysis of ecological data with non-stationary and anisotropic residual dependence. Methods in Ecology and Evolution 2:143–154.

Freni-Sterrantino, A., M. Ventrucci, and H. Rue. 2018. A note on intrinsic conditional autoregressive models for disconnected graphs. Spatial and Spatio-Temporal Epidemiology 26:25–34.

Gelfand, A. E., H. J. Kim, C. F. Sirmans, and S. Banerjee. 2003. Spatial modeling with spatially varying coefficient processes. Journal of the American Statistical Association 98:387–396.

Gneiting, T., and A. E. Raftery. 2007. Strictly proper scoring rules, prediction, and estimation. Journal of the American Statistical Association 102:359–378.

Harris, I., P. D. Jones, T. J. Osborn, and D. H. Lister. 2014. Updated high-resolution grids of monthly climatic observations: the CRU TS3.10 dataset. International Journal of Climatology 34:623–642.

Held, L., B. Schrödle, and H. Rue. 2010. Posterior and cross-validatory predictive checks: a comparison of MCMC and INLA. Pages 91-110 *in* T. Kneib and G. Tutz, editors. Statistical modelling and regression structures. Springer-Verlag, Berlin, Germany.

Hochachka, W. M., D. Fink, R. A. Hutchinson, D. Sheldon, Wong, Weng-Keen, and S. Kelling. 2012. Data-intensive science applied to broad-scale citizen science. Trends in Ecology and Evolution 27:130–137.

Huang, Q., J. R. Sauer, and R. O. Dubayah. 2017. Multidirectional abundance shifts among North American birds and the relative influence of multifaceted climate factors. Global Change Biology 23:3610–3622.

Krainski, E. T., V. Gómez-Rubio, H. Bakka, A. Lenzi, D. Castro-Camilo, D. P. Simpson, F. K. Lindgren, and H. Rue. 2018. Advanced spatial modeling with stochastic partial differential equations using R and INLA. CRC Press, Boca Raton, Florida, USA.

La Sorte, F. A., and F. R. Thompson. 2007. Poleward shifts in winter ranges of North American birds. Ecology 88:1803–1812.

Linden, A., and S. Mantyniemi. 2011. Using the negative binomial distribution to model overdispersion in ecological count data. Ecology 92:1414–1421.

Lindgren, F., and H. Rue. 2015. Bayesian spatial modelling with R-INLA. Journal of Statistical Software 63:19.

Link, W. A., and J. R. Sauer. 1999. Controlling for varying effort in count surveys: an analysis of Christmas Bird Count data. Journal of Agricultural, Biological, and Environmental Statistics 4:116–125.

Link, W. A., and J. R. Sauer. 2002. A hierarchical analysis of population change with application to cerulean warblers. Ecology 83:2832–2840.

Link, W. A., and J. R. Sauer. 2016. Bayesian cross-validation for model evaluation and selection, with application to the North American Breeding Bird Survey. Ecology 97:1746–1758.

Link, W. A., J. R. Sauer, and D. K. Niven. 2006. A hierarchical model for regional analysis of population change using Christmas Bird Count data, with application to the American black duck. Condor 108:13–24.

Link, W. A., J. R. Sauer, and D. K. Niven. 2017. Model selection for the North American Breeding Bird Survey: a comparison of methods. Condor 119:546–556.

Martins, T. G., D. P. Simpson, F. K. Lindgren, and H. Rue. 2013. Bayesian computing with INLA: new features. Computational Statistics and Data Analysis 67:68–83.

Marzluff, J. M. 1997. Effects of urbanization and recreation on songbirds. Pages 89–102 *in* W. M. Block and D. M. Finch, editors. Songbird ecology in southwestern ponderosa pine forests: a literature review. RMRS GTR-292. USDA Forest Service, Rocky Mountain Research Station, Fort Collins, Colorado, USA.

Meehan, T. D., W. Jetz, and J. H. Brown. 2004. Energetic determinants of abundance in winter landbird communities. Ecology Letters 7:532–537.

Meehan, T. D., G. S. LeBaron, K. Dale, N. L. Michel, G. M. Verutes, and G. M. Langham. 2018. Abundance trends of birds wintering in the USA and Canada, from Audubon Christmas Bird Counts, 1966-2017, version 2.1. National Audubon Society, New York, New York, USA. Accessed 1 May 2018, from https://www.audubon.org/conservation/where-have-all-birds-gone

Niven, D. K., J. R. Sauer, G. S. Butcher, and W. A. Link. 2004. Christmas Bird Count provides insights into population change in land birds that breed in the boreal forest. American Birds 58:10–20.

NABCI [North American Bird Conservation Initiative]. 2016. The state of North America’s birds, 2016. Environment and Climate Change Canada, Ottawa, Ontario, Canada.

Pettit, L. I. 1990. The conditional predictive ordinate for the normal distribution. Journal of the Royal Statistical Society, Series B 52:175–184.

Princé, K., and B. Zuckerberg. 2015. Climate change in our backyards: the reshuffling of North America’s winter bird communities. Global Change Biology 21:572–585.

Riebler, A., S. H. Sørbye, D. P. Simpson, and H. Rue. 2016. An intuitive Bayesian spatial model for disease mapping that accounts for scaling. Statistical Methods in Medical Research 25:1145–1165.

Robb, G. N., R. A. McDonald, D. E. Chamberlain, and S. Bearhop. 2008. Food for thought: supplementary feeding as a driver of ecological change in avian populations. Frontiers in Ecology and the Environment 6:476–484.

Robbins, C. S., J. R. Sauer, R. S. Greenberg, and S. Droege. 1989. Population declines in North American birds that migrate to the Neotropics. Proceedings of the National Academy of Sciences of the USA 86:7658–7662.

Root, T. 1988. Energy constraints on avian distributions and abundances. Ecology 69:330–339.

Rosenberg, K. V., P. J. Blancher, J. C. Stanton, and A. O. Panjabi. 2017. Use of North American Breeding Bird Survey data in avian conservation assessments. Condor 119:594–606.

Rosenberg, K. V., J. A. Kennedy, R. Dettmers, R. P. Ford, D. Reynolds, J. D. Alexander, C. J. Beardmore, P. J. Blancher, R. E. Bogart, and G. S. Butcher. 2016. Partners in Flight landbird conservation plan: 2016 revision for Canada and continental United States. Partners in Flight Science Committee.

Rue, H., S. Martino, and N. Chopin. 2009. Approximate Bayesian inference for latent Gaussian models using integrated nested Laplace approximations. Journal of the Royal Statistical Society, Series B 71:319–392.

Rue, H., A. Riebler, S. H. Sørbye, J. B. Illian, D. P. Simpson, and F. K. Lindgren. 2017. Bayesian computing with INLA: a review. Annual Review of Statistics and its Application 4:395–421.

Sauer, J. R., J. E. Fallon, and R. Johnson. 2003. Use of North American Breeding Bird Survey data to estimate population change for bird conservation regions. Journal of Wildlife Management 67:372–389.

Sauer, J. R., and W. A. Link. 2002. Using Christmas Bird Count data in analysis of population change. American Birds 102:10–14.

Sauer, J. R., and W. A. Link. 2011. Analysis of the North American Breeding Bird Survey using hierarchical models. Auk 128:87–98.

Sauer, J. R., K. L. Pardieck, D. J. Ziolkowski Jr, A. C. Smith, M.-A. R. Hudson, V. Rodriguez, H. Berlanga, D. K. Niven, and W. A. Link. 2017. The first 50 years of the North American Breeding Bird Survey. Condor 119:576–593.

Simpson, D., H. Rue, A. Riebler, T. G. Martins, and S. H. Sørbye. 2017. Penalising model component complexity: a principled, practical approach to constructing priors. Statistical Science 32:1–28.

Smith, A. C., M. A. R. Hudson, C. M. Downes, and C. M. Francis. 2015. Change points in the population trends of aerial-insectivorous birds in North America: synchronized in time across species and regions. PLoS ONE 10:e0130768.

Sørbye, S. H., and H. Rue. 2014. Scaling intrinsic Gaussian Markov random field priors in spatial modelling. Spatial Statistics 8:39–51.

Soykan, C. U., J. R. Sauer, J. G. Schuetz, G. S. LeBaron, K. Dale, and G. M. Langham. 2016. Population trends for North American winter birds based on hierarchical models. Ecosphere 7:e01351.

Thogmartin, W. E., J. R. Sauer, and M. G. Knutson. 2004. A hierarchical spatial model of avian abundance with application to cerulean warblers. Ecological Applications 14:1766–1779.

Thogmartin, W. E., M. G. Knutson, and J. R. Sauer. 2006. Predicting regional abundance of rare grassland birds with a hierarchical spatial count model. Condor 108:25–46.

Thomas, C. D., and J. J. Lennon. 1999. Birds extend their ranges northwards. Nature 399:213.

Waller, L. A., and C. A. Gotway. 2004. Applied spatial statistics for public health data. John Wiley and Sons, New York, New York, USA.

Zhang, J., W.D. Kissling, and F. He. 2013. Local forest structure, climate and human disturbance determine regional distribution of boreal bird species richness in Alberta, Canada. Journal of Biogeography 40:1131–1142.

Zuckerberg, B., D. N. Bonter, W. M. Hochachka, W. D. Koenig, A. T. DeGaetano, and J. L. Dickinson. 2011. Climatic constraints on wintering bird distributions are modified by urbanization and weather: wintering birds, weather, food, and climate. Journal of Animal Ecology 80:403–413.

**Figure Legends**

Fig. 1. Select inputs and outputs to the SVC model, including: (A) a map of the grid cell structure used in the American Robin analysis, along with the number of CBC circles included in each cell; (B) Alpha, representing the relative abundance index for 2017; (C) Epsilon, representing the effort-effect exponent; and (D, E, F) Tau, representing the percent change in relative abundance per year from 1966 through 2017. Note that values shown in E are significantly different from 0, based on (F) 95% credible intervals for Tau.

Fig. 2. Comparisons of aggregated relative abundance trends for American Robin produced using SVC (A, C) and standard methods (B, C). SVC trends (Tau, percent change per year from 1966 through 2017) were fine scaled trends averaged per Bird Conservation Region. The dark grey diagonal line in C represents equality.

Fig. 3. Comparisons of 95% credible interval widths for relative abundance trends for American Robin produced using SVC (A, C) and standard methods (B, C). Note that credible interval widths have the same units as SVC trends (Tau) and standard trends, which are percent change per year from 1966 through 2017.

Fig. 4. Model predictions and 95% credible intervals illustrating associations between American Robin SVC trends and (A) degrees north latitude, (B) mean minimum winter temperature between 1965 and 2017, and (C) mean human population size between 1971 and 2015. Note the non-linear transformation of mean human population size.

**Supporting Information**

All code and data needed to reproduce the SVC analysis is freely available at <https://github.com/tmeeha/inlaSVCBC>. A text file of the code is also available here as Data S1.