- A simulation-based approach for quantifying
- and partitioning uncertainty to improve
 - forecasts of dynamic processes
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10 Abstract

- 11 Making informed decisions in the face of rapid environmental change requires forecasts from
- models of ecological processes. However, forecasts from ecological models are often
- associated with high degrees of uncertainty, making it difficult for such forecasts to inform
- decision-making processes. To make progress toward the goal of reliable and informative
- ecological forecasts, we need to know from where forecast uncertainty arises. Such knowledge
- can guide investment in future research that will most improve forecast skill. There is a rich
- 17 history of analytical expressions that partition the variance of future dynamics, but these
- 18 expressions suffer from necessary assumptions such as linear dynamics and small-variance

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approximations that exclude interactions. Similarly, the earth systems modeling community
has developed methods for paritioning uncertainty of model projections, but these operate at
a different modeling grain than most of ecology. Building on these approaches, we develop a
simulation-based approach for quantifying and partitioning forecast uncertainty from
Bayesian state-space models that overcomes the limitations of previous analytical approaches.
Our approach is similar to an Analysis of Variance, where the total variance of a forecast is
paritioned among its constituent parts, namely initial conditions uncertainty, parameter
uncertainty, driver uncertainty, process error, and their interactions. We apply the approach
to near-term forecasts of the Yellowstone bison population and measles in China,
demonstrating the broad utility of our approach. We also provide functions written in the
statistical programming language R, which will allow others using Bayesian state-space
models to employ our approach in their own research.

Keywords: Bayesian state-space model, forecast, Markov chain Monte Carlo, measles, prediction, population model, uncertainty, Yellowstone bison

Introduction

Ecology is entering an era of prediction. This new era is made possible by continuous data streams (e.g., the National Ecological Observatory Network, citizen science data, remote sensing), increased statistical literacy among ecologists, and the need to provide actionable information for decision makers (Dietze et al. 2018). Thus, we are on the verge of answering Clark et al.'s (2001) challenge to make forecasting a core goal of ecology. Ecologists are in an excellent position to meet this forecasting challenge because we have spent decades gaining understanding of the processes that regulate populations, communities, and ecosystems. But we lack a systematic understanding of the current limits to ecological forecasts. As a result, we do not know how to allocate research effort to improve our forecasts. Making poor forecasts is inevitable as ecology matures into a more predictive science.

The key is to learn from our failures so that forecasts become more accurate over time. The

success of meteorological forecasting tells us that basic research on the causes of forecast

uncertainty is an essential component of this learning process (Bauer et al. 2015). Therefore,

we need a systematic and robust way to quantify and partition forecast uncertainty into its

constituent parts (Dietze 2017).

Various approaches have been used to characterize and partition forecast uncertainty (Sobol' 1993, Cariboni et al. 2007). For example, consider a dynamic model designed to predict some state y in the future (y_{t+1}) based on the current state (y_t) , an environmental driver(s) (x), parameters (θ) , and process error (ϵ) . We can then write a general form of the model as:

$$y_{t+1} = f(y_t, x_t | \theta) + \epsilon_{t+1}, \tag{1}$$

which states that y at time t + 1 is a function of y and x at time t conditional on the model parameters (θ) plus process error (ϵ) . Ignoring covariance among factors and assuming linear dynamics, Dietze (2017), following Sobol' (1993) and Cariboni et al. (2007), suggests that forecast variance $(Var[y_{t+1}])$ is approximately:

$$Var[y_{t+1}] \approx \underbrace{\left(\frac{\delta f}{\delta y}\right)^2}_{\text{stability IC uncert.}} \underbrace{Var[y_t]}_{\text{IC uncert.}} + \underbrace{\left(\frac{\delta f}{\delta x}\right)^2}_{\text{driver sens. driver uncert.}} \underbrace{Var[x_t]}_{\text{param sens. param. uncert.}} \underbrace{Var[\theta]}_{\text{param sens. param. uncert.}} + \underbrace{Var[\epsilon_{t+1}]}_{\text{process error}}, \tag{2}$$

where each additive term follows a pattern of *sensitivity* times *variance* and "IC uncert."
refers to "Initial Conditions uncertainty." The variance attributable to any particular factor
is a function of how sensitive the model is to the factor and the variance of that factor. For
example, the atmosphere is a chaotic system, meaning its dynamics are internally unstable

and sensitive to initial conditions uncertainty. This is why billions of dollars are spent each year to measure meterological variables – meterologists learned that the key to reducing forecast error $(Var[y_{t+1}])$ was to reduce the uncertainty of initial conditions $(Var[y_t])$. In contrast, ecologists are attempting to make actionable forecasts with little knowledge of which term in Eq. 2 dominates forecast error. Knowing which term dominates forecast error in different ecological settings will advance our fundamental understanding of the natural world and immediately impact practical efforts to monitor, model, and predict ecological dynamics.

While having an analytical expression such as Eq. 2 is satisfying, arriving at the expression involves strict assumptions. First, Eq. 2 only holds when the underlying dynamics are linear, which may not be the case for many populations and models. Second, Eq. 2 only includes additive effects of each factor because the Taylor series decomposition requires small-variance approximations that eliminate interactions. But, interactions among the factors are probably common. For example, in a simple simulation of an AR(1) process, we show that initial conditions uncertainty and parameter error interact to generate the full spread of forecast variance (Figure 1). Analyzing only the main effects of each source of uncertainty would lead to the false conclusion that initial conditions uncertainty is not important. Progress in quantifying and partioning forecast uncertainty therefore requires a more flexible approach than that provided by Eq. 2.

We have four objectives: First, we introduce a canonical equation for error
propagation and review approaches for quantifying sources of forecast uncertainty from other
fields. Second, we describe a general, simulation-based method for quantifying and
partitioning forecast uncertainty from Bayesian state-space models. Third, we apply our
method to near-term forecasts of the Yellowstone bison population and measlse cases in
China. These two applications demonstrate the generality of our approach. Fourth, through
our applications we introduce functions written in the statistical programming language R

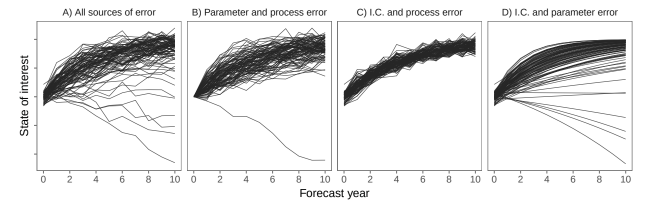


Figure 1: Example of forecast uncertainty with different sources of error set to zero. Each line represents one realization, out of 200, from an order-one autoregressive model (AR(1) process). Contrary to the analytical expression (Eq. 2), initial conditions (I.C.) uncertainty and parameter uncertainty clearly interact. The spread of lines in (A) is not wholly because of initial conditions uncertainty or parameter uncertainty (compare panels B and C). It is their combined influence that causes the spread of realizations in (A and D). At least in this example, process error (set to zero in D) does appear to be independent, but we used a small value of process error to highlight other interactions. Source code: generate_forecast_fxns.R.

- 88 (R Core Team 2016), which can be used to implement our method for many Bayesian
- 89 state-space models.

90 A Brief History of Quantifying and Partitioning

51 Forecast Uncertainty

92 Error propagation

- 93 Methods for quantifying and partitioning forecast uncertainty all share common roots in
- 94 statistical error propagation. Error propagation is concerned with translating the effects of
- variable (or parameter) uncertainty into the uncertainty of a function based those variables.
- That is, for the output q of some function $q = f(x_1, x_2, \dots, x_n)$, we are interested in the
- variance of q given the variance of input variables (\mathbf{x}) . It is this generic model formulation
- that leads to the canonical expression of error propagation

$$\sigma_q^2 = \left(\frac{\delta q}{\delta x_1} \sigma_{x_1}\right)^2 + \left(\frac{\delta q}{\delta x_2} \sigma_{x_2}\right)^2 + \dots + \left(\frac{\delta q}{\delta x_n} \sigma_{x_n}\right)^2 \tag{3}$$

$$= \sum_{i=1}^{n} \left(\frac{\delta q}{\delta x_i} \sigma_{x_i} \right)^2. \tag{4}$$

This expression states that the variance of the output of function q (σ_q^2) is equal to the sum of squares of the input variables weighted by the sensitivity of the output variable to each input variable, quantified as the partial derivative. Add more here on interactions, which require knowing the correlations among x's.

103 Dynamic theoretical models

Lotka-Volterra style models and variance of t+1 (draws from error propagation and sensitivity analysis).

106 Weather forecasting

107 Chaos and initial conditions.

Earth system models

109 Carbon and uncertainty from models and scenarios.

110 A Simulation-Based Approach for Partitioning

111 Forecast Uncertainty

121

128

Analytical expressions of forecast uncertainty must rely on simplifying assumptions. Two 112 important assumptions are (1) that different sources of uncertainty do not interact and (2) 113 that the system of equations is linear. These analytical expressions are important for guiding 114 our intuition, but these strict assumptions limit our ability to partition forecast uncertainty 115 in practice. Thus, we present a simulation approach that is entirely model-based and 116 requires no assumptions, other than those embedded in the model itself. We are building on 117 the ideas put forth by Dietze (2017), who suggested a simulation approach for quantifying 118 the terms in Eq. 2. Here we generalize the approach to fully partition forecast uncertainty 119 among the main effects of different sources and their interactions. 120

As a starting point, consider the Bayesian state-space model

Data Model:
$$y_t \sim \left[y_t \mid z_t, \sigma_o^2 \right], \qquad t = 1, \dots, T,$$

Process Models: $z_t \sim \left[z_t \mid \mu_t, \sigma_p^2 \right],$

$$\mu_t = g\left(z_{t-1}, \mathbf{x}_t', \mathbf{\theta} \right), \qquad t = 2, \dots, T,$$

Parameter Models: $\mathbf{\phi} \sim \left[\mathbf{\theta}, \sigma_p^2, \sigma_o^2, z_{t-1} \right],$ (5)

where y_t is the observed state at time t, z_t is the latent state at time t, μ_t is the determinstic prediction of z at time t from the process model g, which is a function of z at time t-1, a vector of covariates (\mathbf{x}) at time t, and a set of unknown parameters, $\mathbf{\theta}$ (Berliner 1996). σ_0^2 is observation error and σ_p^2 is process error. The notation $[a \mid b, c]$ reads, "the probability of agiven b and c" (Gelfand and Smith 1990), and $\mathbf{\phi}$ refers to the prior probability distributions for all unknown parameters and the initial conditions for the latent state, $z_{t=1}$.

For our purposes, we are interested in the probability distributions of the true state z

at future points in time, conditional on previous observations (\mathbf{y}). This is referred to as the forecast distribution or the predictive process distribution (Hobbs and Hooten 2015 pp. 131 199–200), which, for one time step ahead of the final observation (T + 1), is defined as

$$[z_{T+1}|y_1,\ldots,y_T] = \int \int \ldots \int \left[z_{T+1}|z_T,\mathbf{x}_T,\mathbf{\theta},\sigma_{\mathbf{p}}^2\right] \times \left[z_1,\ldots,z_{T+1},\mathbf{\theta},\sigma_{\mathbf{p}}^2|y_1,\ldots,y_T\right] d\mathbf{\theta} d\sigma_{\mathbf{p}}^2 dz_1\ldots dz_T.$$
(6)

The model in Eq. 5 can be fit using Markov chain Monte Carlo (MCMC) algorithms, 132 which makes calculating the forecast distribution a relatively simple task. To obtain $[z_{T+1}|y_1,\ldots,y_T]$, we can change the indexing of t in Eq. 5 to $t=2,\ldots,T+1$ and then 134 sample $z_{T+1}^{(k)}$ from $\left[z_{T+1}|g(z_T^{(k)},\mathbf{x}_{T+1}^{(j(k))},\boldsymbol{\theta}^{(k)}),\sigma_{\mathbf{p}}^{2(k)}\right]$ given the current values for $\boldsymbol{\theta}^{(k)}$ and $z_T^{(k)}$ 135 on each k = 1, ..., K iteration of the MCMC algorithm (Hobbs and Hooten 2015, Williams 136 et al. 2018). Note that we index the external covariate vector \mathbf{x} using j and k, where j(k) is 137 realization j of the covariate vector \mathbf{x} associated with MCMC sample k. In some cases, the 138 external covariate will have only one value, in which case all K MCMC samples will share 139 the same x. In other cases, their may be a distribution of external covariate values 140 associated with uncertainty from the forecast of \mathbf{x} , resulting in n values for each x_{T+1} . When 141 n < K, which we anticipate it often will be, then x can be sampled with replacement and a 142 value can be assigned to each MCMC sample. Making forecasts further into the future than 143 T+1 requires extending T+1 to $T+2,\ldots,T+q$ and iteratively sampling 144 $\left[z_{T+q}|g(z_{T+q-1}^{(k)},\mathbf{x}_{T+q}^{(j(k))},\boldsymbol{\theta}^{(k)}),\sigma_{\mathbf{p}}^{2(k)}\right]$ (Hobbs and Hooten 2015).

The forecast distribution (Eq. 6) has all of the quantities that contribute to forecast uncertainty by incorporating their uncertainty explicitly across the K MCMC iterations. For example, initial conditions uncertainty is incorporated because $z_{T+1}^{(k)}$ is a function of $z_T^{(k)}$, resulting in a total of K point forecasts for z that comprise the posterior distribution of z at each time t. If we wish to ignore initial conditions uncertainty (I.C. uncertainty), we can make all K point forecasts starting from the mean of z_T

Table 1: Sampling equations for generating forecast distributions at times T+q (where T is the time of the last observation) across $k=1,\ldots,K$ MCMC samples with only certain sources of uncertainty present.

Source of Uncertainty	Notation	Sampling Equation	
I.C. Uncertainty	$V^{(I)} = V^{(I,\overline{PA},\overline{D},\overline{PS})}$	$\left[z_{T+q} \mid g(z_{T+q-1}^{(k)}, \mathbf{x}_T^{(*)}, \mathbf{\theta}^{(*)}), 0\right]$	
Parameter Uncertainty	$V^{(PA)} = V^{(\overline{I}, PA, \overline{D}, \overline{PS})}$	$ [z_{T+q} \mid g(z_{T+q-1}^{(k)}, \mathbf{x}_T^{(*)}, \boldsymbol{\theta}^{(k)}), 0], $ $ [z_{T+q} \mid g(z_T^{(*)}, \mathbf{x}_T^{(*)}, \boldsymbol{\theta}^{(k)}), 0], $	q > 1 $q = 1$
Driver Uncertainty	$V^{(D)} = V^{(\overline{I}, \overline{PA}, D, \overline{PS})}$	$ [z_{T+q} \mid g(z_{T+q-1}^{(k)}, \mathbf{x}_T^{(j(k))}, \boldsymbol{\theta}^{(*)}), 0], $ $ [z_{T+q} \mid g(z_T^{(*)}, \mathbf{x}_T^{(j(k))}, \boldsymbol{\theta}^{(*)}), 0], $	q > 1 $q = 1$
Process Uncertainty	$V^{(PS)} = V^{(\overline{I}, \overline{PA}, \overline{D}, PS)}$	$ [z_{T+q} \mid g(z_{T+q-1}^{(k)}, \mathbf{x}_{T}^{(*)}, \boldsymbol{\theta}^{(*)}), \sigma_{\mathbf{p}}^{2(k)}], $ $ [z_{T+q} \mid g(z_{T}^{(*)}, \mathbf{x}_{T}^{(*)}, \boldsymbol{\theta}^{(*)}), \sigma_{\mathbf{p}}^{2(k)}], $	q > 1 $q = 1$

Note: The notation $V^{(A,\overline{B},\overline{C})}$ means that uncertainty from A enters the forecast from its posterior distribution (as approximated from MCMC samples), while sources B and C are set to their means (as calculated across the MCMC samples). In the main text we leave out the averaged terms (those with overbars) to reduce clutter. Thus, $V^{(A)} = V^{(A,\overline{B},\overline{C})}$.

$$z_T^{(*)} = E(z_T|y_1, \dots, y_T) \approx \frac{\sum_{k=1}^K z_T^{(k)}}{K},$$
 (7)

which we call $z_T^{(*)}$ (as opposed to $z_T^{(k)}$), while retaining the uncertainty for all other parameters. Our iterative sampling to obtain the forecast distribution then becomes a conditional statement,

$$z_{T+q} \sim \begin{cases} \left[z_{T+q} | g(z_{T+q-1}^{(k)}, \mathbf{x}_{T}^{(j(k))}, \boldsymbol{\theta}^{(k)}), \sigma_{\mathbf{p}}^{2(k)} \right], & q > 1 \\ \left[z_{T+q} | g(z_{T}^{(*)}, \mathbf{x}_{T}^{(j(k))}, \boldsymbol{\theta}^{(k)}), \sigma_{\mathbf{p}}^{2(k)} \right], & q = 1. \end{cases}$$
(8)

We can extend the basic idea of setting subsets of parameters and states to their posterior means (or medians, depending on the distribution) to make partitioned forecasts where only prescribed sources of uncertainty contribute to forecast uncertainty (Table 1). It is important to note that although our discussion has centered on obtaining forecast distributions within the MCMC algorithm used to fit the model, it is only feasible to do this for the full forecast distribution (Eq. 6). In all other cases, where states or parameters must be averaged over the K MCMC iterations, forecast simulations must be done post hoc using saved MCMC samples (Box 1). In other words, estimating z_1, z_2, \ldots, z_T is done within the MCMC fitting algorithm, while estimating $z_{T+1}, z_{T+2}, \ldots, z_{T+q}$ is done after fitting the model, but with the full MCMC output.

With these basics in mind, we now develop our approach for partitioning and quantifying uncertainty, which is similar to an Analysis of Variance (ANOVA). Let the variance of the forecast distribution at time T+q be $V_{T+q}^{(X)}$, where X=F (full forecast distribution), I (initial conditions uncertainty only), PA (parameter uncertainty only), D (driver uncertainty), or PS (process uncertainty only) (Table 1). $V_{T+q}^{(I)}$, $V_{T+q}^{(PA)}$, $V_{T+q}^{(D)}$, and $V_{T+q}^{(PS)}$ are the main effects of each factor on the forecast distribution such that

$$V_{T+q}^{(F)} = V_{T+q}^{(I)} + V_{T+q}^{(PA)} + V_{T+q}^{(D)} + V_{T+q}^{(PS)}$$

$$+ \epsilon_{T+q}^{(I,PA)} + \epsilon_{T+q}^{(I,D)} + \epsilon_{T+q}^{(I,PS)} + \epsilon_{T+q}^{(PA,PS)} + \epsilon_{T+q}^{(PA,D)} + \epsilon_{T+q}^{(D,PS)}$$

$$+ \epsilon_{T+q}^{(I,PA,D)} + \epsilon_{T+q}^{(I,PA,PS)} + \epsilon_{T+q}^{(I,D,PS)} + \epsilon_{T+q}^{(PA,D,PS)}$$

$$+ \epsilon_{T+q}^{(I,PA,D,PS)},$$

$$(9)$$

where the notation $\epsilon_{T+q}^{(X,Y)}$ represents the remaining interactive effect of X and Y on $V_{T+q}^{(F)}$ after accounting for their main effects. For example, if the full forecast variance is a function of only initial conditions uncertainty I and parameter uncertainty PA, then

$$V_{T+q}^{(F)} = V_{T+q}^{(I)} + V_{T+q}^{(PA)} + \epsilon_{T+q}^{(I,PA)}, \tag{10}$$

which rearranges to

$$\epsilon_{T+q}^{(I,PA)} = V_{T+q}^{(F)} - \left[V_{T+q}^{(I)} + V_{T+q}^{(PA)} \right]. \tag{11}$$

We show an example of applying Eqs. 10-11 in a hypothetical situation where forecast uncertainty is determined by initial conditions uncertainty and parameter uncertainty alone in Figure 2.

The necessary terms for partitioning forecast variance can be obtained by calculating
the variance of the partitioned forecast distributions (equations in Table 1 and combinations
thereof). To take this one step further, and to reiterate the core idea, let V(F) be a function
of initial conditions uncertainty I, parameter uncertainty PA, and driver uncertainty D. We
can then write the equation for forecast uncertainty and the derived interaction effects as

There must be a way to write a concis form of these equations vai summations.

$$V_{T+q}^{(F)} = V_{T+q}^{(I)} + V_{T+q}^{(PA)} + V_{T+q}^{(D)} + \epsilon_{T+q}^{(I,PA)} + \epsilon_{T+q}^{(I,D)} + \epsilon_{T+q}^{(D,PA)} + \epsilon_{T+q}^{(I,PA,D)}, \quad \text{where} \quad (12)$$

$$\epsilon_{T+q}^{(I,PA)} = V_{T+q}^{(I+PA)} - \left[V_{T+q}^{(I)} + V_{T+q}^{(PA)} \right] \tag{13}$$

$$\epsilon_{T+q}^{(I,D)} = V_{T+q}^{(I+D)} - \left[V_{T+q}^{(I)} + V_{T+q}^{(D)} \right] \tag{14}$$

$$\epsilon_{T+q}^{(PA,D)} = V_{T+q}^{(PA+D)} - \left[V_{T+q}^{(PA)} + V_{T+q}^{(D)} \right] \tag{15}$$

$$\epsilon_{T+q}^{(I,PA,D)} = V_{T+q}^{(F)} - \left[V_{T+q}^{(I)} + V_{T+q}^{(PA)} + V_{T+q}^{(D)} \right] - \left[\epsilon_{T+q}^{(I,PA)} + \epsilon_{T+q}^{(I,D)} + \epsilon_{T+q}^{(PA,D)} \right], \tag{16}$$

where the notation $V_{T+q}^{(A+B)}$ is the forecast variance under scenario where both A and B are allowed to contribute to forecast uncertainty (i.e., a combination of the equations in Table 1). We present the equations for the full partition among I, PA, D, and PS in Appendix 1. In the next two sections, we apply our approach (Box 1) to quantify and partition the uncertainty of near-term forecasts of the Yellowstone bison population and measles cases in China.

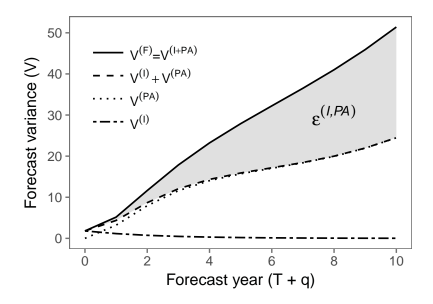


Figure 2: Graphical example of partitioning forecast uncertainty into main effects (V) and interaction effects (ϵ) . The grey shaded area shows the interaction effect $(\epsilon^{(I,PA)})$ that must be accounted for to fully partition forecast uncertainty between initial conditions uncertainty $(V^{(I)})$ and parameter uncertainty $(V^{(PA)})$. Source code: generate_forecast_fxns.R.

Box 1. Pseudocode for quantifying and partitioning forecast uncertainty from a Bayesian state-space model.

- 1. Fit a Bayesian state-space model (i.e., Eq. 5) with data y_1, \ldots, y_T and save the MCMC samples.
- 2. Forecast $z_{T+q}^{(k)}$ for all k = 1, ..., K MCMC samples to generate the full forecast distribution following Eq. 6 (this can be done within the MCMC algorithm or *post hoc* with saved MCMC samples).
- 3. Forecast $z_{T+q}^{(k)}$ for all k = 1, ..., K MCMC samples to generate the partitioned forecast distributions for each source of uncertainty, averaging quantities over the K MCMC samples as necessary (equations in Table 1 and combinations thereof).
- 4. For each forecast time q, calculate the variance of each forecast distribution from steps 2-3.
- 5. Partition forecast variance using Eq. X.

Box 1:

Application: Yellowstone Bison Population

Application: Measles in China

Discussion

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