Analyzing Stochastic Computer Models: A Review with Opportunities

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Abstract. In modern science, computer models are often used to understand complex phenomena and a thriving statistical community has grown around analyzing them. This review aims to bring a spotlight to the growing prevalence of stochastic computer models—providing a catalogue of statistical methods for practitioners, an introductory view for statisticians (whether familiar with deterministic computer models or not), and an emphasis on open questions of relevance to practitioners and statisticians. Gaussian process surrogate models take center stage in this review, and these, along with several extensions needed for stochastic settings, are explained. The basic issues of designing a stochastic computer experiment and calibrating a stochastic computer model are prominent in the discussion. Instructive examples, with data and code, are used to describe the implementation of, and results from, various methods.

Key words and phrases: Computer model, Gaussian process, uncertainty quantification, emulator, computer experiment, agent based model, surrogates, calibration.

1. INTRODUCTION

Computer models, also known as simulators, are in use everywhere. These are programs which describe and approximate a process of interest. The code typically takes a set of inputs and produces some output. Stochastic simulators, unlike deterministic ones, can produce a different output even with the same inputs due to the presence of random elements.¹ Such computer models are in wide use. For example, agent-based models (ABMs) deal with large populations of individuals, where specific actions taken at each time-step are to complex for deterministic modeling. ABMs are prevalent (Johnson, 2010, Johnson and Briggs, 2011, Ramsey and Efford, 2010, Smieszek

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¹This terminology can have different meanings and connotations in different fields. In weather modeling, a stochastic simulator might refer to a random weather generator (Richardson, 1981, Peleg et al., 2017). In this work, we use the term to refer to any code that includes (pseudo-) random elements in generating output.

et al., 2011, Grimm et al., 2006) and used to explore complex phenomena in sociology, transportation, ecology, epidemiology and other fields.

The following is a basic model of a stochastic simulator experiment. If the code is run at a (vector) input x producing a (scalar) output y(x), this could be represented as

(1.1)
$$y(x) = M(x) + v, \quad v \sim N(0, \sigma_v^2(x)),$$

where M(x) is the expected value, E[y(x)], of the output and v is independent variability representing the randomness of the simulator. Its variance, σ_v^2 , can depend on x, but constant variance is also possible. For deterministic simulators, $\sigma_v^2 = 0$.

Randomness in stochastic simulators invariably requires many simulations, which limits the complexity (including the size of the input dimension) that can be effectively analyzed. The prospect of replicated runs in stochastic simulators introduces a trade-off between replication and exploration, a challenging design issue. Analysis is also harder when the noise, v, has nonconstant variance. This article examines these basic issues, identifies accessible and effective methods, and points to unresolved questions that should be addressed.

Equation (1.1) is often used to model physical experiments, where an observation y(x) is the truth, M(x), plus measurement error (and, possibly, intrinsic variability as well), or for an observational study, where M(x) is fit to the observations with residual variance. Because they are structurally the same, physical experiments can be analyzed with the same methods used for stochastic simulators (Gao, Sacks and Welch, 1996). However, the contexts and goals are often different, leading to different problem formulations and different interpretations of results.

The choice of method, with its assumptions and limitations, is crucial for any experiment. A desire for simplicity, and availability of software, would encourage the use of a standard statistical regression model (e.g., linear regression) for M with a constant σ_v^2 . This approach can be effective under some circumstances, especially when the space, X, of possible inputs is small, which begs the question of how reliable it can be as a general prescription. Complex systems modeled by a simulator rarely allow for such simplification. The methods described in this review allow the simulated data to guide the choice of model under general conditions with little or no simplification.

Statistics (Sacks et al., 1989, Kennedy and O'Hagan, 2001) and Applied Mathematics (Sullivan, 2015) play prominent roles in the design and analysis of deterministic computer experiments. Unsurprisingly, some methods developed for deterministic simulators have modifications that can be used in the stochastic context. Alternatives, driven by the stochasticity, are necessary in many contexts. These structural differences will be noted in the narrative below.

1.1 Goals

We have three primary goals; all related to the crossdisciplinary nature of this topic.

One goal is to bring effective statistical methods to the attention of subject scientists and enable a deeper understanding of stochastic simulators in use. The descriptions below of statistical tools used (or cited) try to avoid being bogged down in mathematical intricacies. Some details of individual methods are included to help in understanding the strengths and weaknesses of the methods. Application of a number of methods is exemplified on testbed cases (Section 2), and available software for methods are identified where possible.

A second goal is to familiarize statisticians with an area of major importance that is crucial to the formation of evidence-based policy. Statisticians are sorely needed in the study and application of agent-based models (ABMs) and stochastic simulators in general. Researchers familiar with deterministic simulation techniques will see immediate opportunities, but statistical expertise of all kinds is essential to advance the study of stochastic simulators.

The analysis of stochastic simulators is a developing field with many unsolved problems. Challenges are often driven by the scale of the problems and a range of issues whose resolution requires close cooperation between statisticians, subject scientists and computer scientists. A third goal of this paper is to spur that process.

The review is structured as follows: Section 2 briefly outlines the simulators used as examples throughout. Section 3 describes the models that form the basis for the analyses. Section 4 is devoted to the fundamental question of what simulator runs to make. Section 5 addresses a common objective of simulation experiments: calibration. Section 6 discusses other models and objectives that are important, but are more on the "boundaries" of this review and are therefore less detailed. Finally, Section 7 summarizes conclusions and poses unanswered questions. The references here do not cover the entire body of work on stochastic simulators but, together with this overview, should provide adequate coverage of the problems discussed.

2. EXAMPLE SIMULATORS

Three stochastic simulators will be discussed throughout this review to aid understanding. Two are deliberately simplified and used to exhibit key features of the methods presented. In some cases, simpler strategies could be equally effective because the complexity of the models has been greatly reduced; the demonstration purpose is the one that is relevant in the discussion and reported computations.

The third is a model which we use to anchor and motivate methods. The specific model in question is an epidemiological model developed in response to the Ebola epidemic of 2014. For the Ebola model, a synthetic population representing the individuals in Liberia (population ~ 4.5 million) and their activity schedules, inducing a time-varying contact network of individuals and locations, was developed (Mortveit et al., 2015), and paired with an agent-based model (Bisset et al., 2009). Together, this ABM models a contagion spreading from one individual to another in Liberia. Since the parameter for contagion, transmissibility, only controls the probability of infection for a given interaction, this model is stochastic. The model is updated daily, with the progress of the disease determined by the activity schedule, contact details, and other epidemiological characteristics. This model is complex, with high dimensional outputs, multiple unknown inputs and nonnormality all present. The analysis performed by Fadikar et al. (2018) tackles all of these using ideas discussed within this article (see Sections 3.3.1, 3.4 and 5).

2.1 Fish Capture–Recapture

The first simplified stochastic simulator we consider mimics the movements and schooling behavior of fish in a mark-recapture application. Mark and recapture involves capturing a sample of the population, marking and releasing them and following up by capturing another sample and counting how many are marked—the "recaptured." The number recaptured allows estimation of the population size (Begon et al., 1979).

This process can be modeled by initializing a population of fish at random locations in a 2-d, rectangular lake with boundary conditions. The fish begin moving and schooling according to simple, agent-based rules. After an initial period of time, 100 fish are marked as they pass through a "net" in the lake. After a second period of time, 100 fish are captured using the same net and the number of "recaptured" are recorded. This agent-based model is a modified version of the flocking model developed in NetLogo (Wilensky, 1999). The collective behavior that emerges in the flocking model is the result of providing each individual agent with the same set of simple rules (Reynolds, 1987). The flocking model is modified to include the mark-recapture dynamics described above. Given an observed count of recaptured fish, this model can be used to estimate the total size of the fish population (see Section 5.3). The only input considered is the number of fish in the total population and the output is the number of recaptured fish. Other inputs for this model control the individual movement rules of the fish; for simplicity, these are ignored here and set to default values. The supplementary code and compiled Rmarkdown documents, corresponding to our analysis of this simulator can be found at https://github.com/jhuang672/fish. Running the simulator afresh will require the installation of NetLogo from https://ccl.northwestern.edu/netlogo/.

2.2 Ocean Circulation

The second simplified example is a stochastic simulator that models the concentration of oxygen in a thin water layer (around 2000 m deep) in the South Atlantic ocean (McKeague et al., 2005, Herbei and Berliner, 2014). The physical model is described via an advection-diffusion equation (equation (4) of McKeague et al., 2005), that is, a nonlinear partial differential equation (PDE) describing the dynamics of oxygen concentration in terms of the water velocities and diffusion coefficients. For a given set of inputs, the solution of the advection-diffusion equation is not available in closed form. However, using theoretical results (Feynman, 1948, Kac, 1949), the solution can be closely approximated through an associated random process (Herbei and Berliner, 2014). For a specific location within the domain, random paths of the process are generated, producing noisy outcomes that approximate the solution to the PDE at that location. This example is simplified by taking the oxygen concentration output to only depend on four inputs: two unknown diffusion constants (K_x and K_y) and the two location variables (latitude and longitude). All other inputs are held fixed at nominal values. Such stochastic approximations are numerous in physical sciences, either due to computational limitations, a lack of complete understanding of the underlying system or because the system under study is itself believed to be random. The supplementary code and compiled Rmarkdown documents, corresponding to our analysis of this simulator can be found at https://github.com/Demiperimetre/Ocean.

3. STATISTICAL MODELS

An experiment involving running a simulator and producing data whose output is described by equation (1.1) can have a multitude of goals. A principal objective, and the one we focus on here, is using the simulated data to predict values of the simulator, M(x) + v, and the uncertainties of these predictions, at untried xs in a context where getting new runs of the simulator is expensive. When M is believed to be "simple" (e.g., a polynomial function of the coordinates of x) there are many standard "classical" techniques that can be used to approximate M. For example, linear regression models and generalised linear models have been used by Andrianakis et al. (2017) and Marrel et al. (2012). Complex problems such as those in Section 2 are less easily managed: specifying a functional form for complex M requires sufficient prior knowledge or a huge abundance of data, both of which are often lacking. This article focuses on methods which can flexibly represent M.

There are a range of factors that need to be taken into account before choosing a statistical model (hereon referred to as a *surrogate model*, as it acts as a surrogate for the computer model). In addition to methodological assumptions, it is important to consider the "context," that is, the conditions of the particular problem being studied. Some important contexts include:

- The space of inputs is usually a hyper-rectangle: each coordinate of an input *x* is constrained by upper and lower bounds. Section 2.2 simplifies issues by taking a rectangular input space even though the Atlantic Ocean is not rectangular.
- The output *y* in equation (1.1) is scalar, but multiple output, such as time series, is also common.
- Some inputs may be categorical rather than numerical.
- The probability distribution of the variability, *v*, is often taken to be normal, which is sometimes invalid (as with the Ebola model).

Stretching back to Sacks et al. (1989) and Currin et al. (1991), a vast literature, mostly on deterministic simulators, has found that Gaussian Processes (GPs) produce flexible, effective surrogates for M. This approach, and modifications needed to address the variability v, can be effective for stochastic simulation (Kleijnen, 2009, Kleijnen, 2017) and will be apparent below. A thorough, intuitive, explanation (for deterministic computer models) can be found in O'Hagan (2006). More technical descriptions of GPs from a statistical perspective can be found in Santner, Williams and Notz (2018) and Gramacy (2020); for a machine learning perspective, see Rasmussen and Williams (2006). In brief, the use of GPs allows computer model runs to play the key role in selecting a surrogate and assessments of its uncertainty in prediction. Deep learning methods, such as neural networks, are also in wide use. These methods can struggle to produce accurate uncertainties estimates (important for simulation experiments), but there is active research directed towards this end (Neal, 1996, Graves, 2011, Welling and Teh, 2011, Papamakarios et al., 2019, Gal and Ghahramani, 2016, Lakshminarayanan, Pritzel and Blundell, 2017).

3.1 Gaussian Process Surrogates

Suppose that the input space X is a hyper-rectangle in d-dimensions; the output y(x) is univariate (scalar); and the variability is normally distributed. Additionally, assume:

A1 The variability, v, has constant variance σ_v^2

A2 The mean $M(x) = \mu + Z(x)$

A3 μ is constant

A4 $Z(\cdot)$ is a Gaussian process on X with mean 0 and covariance function K, deconstructed as a product of a variance σ_Z^2 and a correlation function C.

The technical definition of a GP (Assumption A4) is: for *any* finite N and collection of inputs $X_N =$

 $(x_1, \ldots, x_N), Z_N = (Z(x_1), \ldots, Z(x_N))^{\top}$ is a multivariate normal random vector with mean **0** and $N \times N$ covariance matrix K_N , whose entries are $K(x_i, x_j)$. It follows that the simulator output, $Y_N = (y(x_1), \ldots, y(x_N))^{\top}$, is also multivariate normal but with mean $\mu \mathbf{1}$ and covariance matrix $K_N + \sigma_v^2 I_N$, where I_N is the identity $N \times N$ matrix and **1** is the *N*-vector of 1s.

A simpler interpretation is that these assumptions describe a *prior* distribution for all possible functions for the mean M. Different choices for the GP allow for different classes of possible M; the power of a GP is that these classes can be big enough to allow for all reasonable possibilities. After specifying μ , K, and σ_v^2 , a Bayesian analysis can then be carried out, resulting in a *posterior* distribution for all the functions that can still represent Mafter accounting for the observed simulator runs.

Another interpretation of M and Assumptions A2 and A4 is to think of M as a random function, with μ being a regression function (as in linear regression), and the GP for Z modeling local divergences from μ . Both formulations have the same mathematical structure but with slightly different interpretations.

The predictive distribution for any new run, $y(x_{\text{new}})$, given the observed simulator data $\{X_N, Y_N\}$ is also normal, and has a known analytical form. The mean $\mu_N(x_{\text{new}})$ and variance $\sigma_N^2(x_{\text{new}})$ of predictions are

(3.1)

$$\begin{aligned} & \mu_N(x_{\text{new}}) \\ &= \mu + k_N(x_{\text{new}})^\top (K_N + \sigma_v^2 I_N)^{-1} (Y_N - \mu \mathbf{1}), \\ & \sigma_N^2(x_{\text{new}}) \\ (3.2) &= \sigma_v^2 + \sigma_Z^2 \\ & - k_N(x_{\text{new}})^\top (K_N + \sigma_v^2 I_N)^{-1} k_N(x_{\text{new}}), \end{aligned}$$

with $k_N(x_{\text{new}})$ denoting the *N*-vector ($K(x_{\text{new}}, x_1), \ldots, K(x_{\text{new}}, x_N)$)^{\top} of covariances between the desired prediction and observed data. Once the correlation function *C* is specified, the parameters (μ, σ_Z^2 , and σ_v^2) can be estimated from the data. *C* is often specified to also contain parameters, θ , which can be estimated, thereby tailoring *C* to observations. One example for *C* is the squared-exponential correlation function (also known as the Gaussian kernel):

(3.3)
$$C(x,w) = \exp\left\{-\sum_{j=1}^{d} \frac{(x_j - w_j)^2}{\theta_j}\right\}.$$

This correlation function is suited for approximating very smooth, infinitely differentiable, functions. Alternative correlation functions exist and are used; one commonly used alternative is the Matérn 5/2 correlation function



FIG. 1. Fish example: Predictive means and 95% uncertainty intervals for the fish model using homGP and hetGP. Data are 400 simulator runs, consisting of 20 replicates at 20 different population sizes (a maximin Latin hypercube scaled to [150, 4000], rounded down to nearest integers). The left panels use homGP—the solid red line is the median of the predictive distribution and the dashed red-lines form the 95% uncertainty intervals. The right panels use hetGP. The upper panels include the data used to fit the surrogates; the lower panels omit the data but include the "true" values in black.

(Stein, 2012), which is appropriate for approximating less-smooth functions (only 2 derivatives).²

With a choice of *C* and Assumptions A1–A4, the likelihood of the observed output is available and maximum likelihood estimates (MLEs) $\hat{\mu}$, $\hat{\sigma}_v^2$, $\hat{\sigma}_Z^2$ and $\hat{\theta}$ can be calculated. Henceforth, $\mu_N(x_{\text{new}})$ and $\sigma_N^2(x_{\text{new}})$ will be used to denote the mean and variance of the predictive distribution even when the parameters in equations (3.1) and (3.2) are estimated. The predictive probability distribution for the computer model output $y(x_{\text{new}})$ is then

(3.4)
$$y(x_{\text{new}}) \sim N(\mu_N(x_{\text{new}}), \sigma_N^2(x_{\text{new}})).$$

A complete assessment of uncertainty is lost by plugging in estimated parameters without accounting for their uncertainty. Accordingly, the predictive variance, $\sigma_N^2(x_{\text{new}})$, obtained this way is called the *plug-in* (or *nominal*) predictive variance. The alternative of a full Bayesian analy-

$$\prod_{i=1}^{d} \left(1 + \frac{\sqrt{5}|x_j - w_j|}{\theta_j} + \frac{5|x_j - w_j|^2}{3\theta_j^2}\right) \exp\left(-\frac{\sqrt{5}|x_j - w_j|}{\theta_j}\right)$$

sis to estimate the parameters can be computationally impractical in many circumstances, though not impossible (intermediate schemes and approximations have proven to be useful, e.g., Spiller et al., 2014).

For the correlation function in equation (3.3), and for others such as the Matérn 5/2, the correlation between Z(x) and Z(w) depends only on x - w, the difference between the two vectors of inputs. That is, Z is assumed to be a stationary GP (and, consequently, so is y). For functions exhibiting markedly different behavior in one region of input space than in another part, stationarity is problematic. This issue is tackled and discussed in Gramacy and Lee (2008), Ba and Joseph (2012), Kersaudy et al. (2015) and Chen et al. (2016), among others, and Section 6.1 discusses one solution.

Despite the fairly complex mathematical expressions above, Gaussian processes are easily accessible thanks to numerous available packages (e.g., DiceKriging in R (Roustant, Ginsbourger and Deville, 2018), the hetGP R package (Binois and Gramacy, 2018) mentioned later, and the GaussianProcessRegressor function from scikitlearn in Python (Pedregosa et al., 2011)). In general, a GP is a flexible method for estimating the mean M(x) of the simulator output, even with a lack of prior knowledge. This is illustrated in the top panels of Figures 1 and 2, but

²The Matérn 5/2 can be written as

and is the product of d one-dimensional covariance functions. Further discussion of the features of different kernels can be found in Chapter 4.2 of Rasmussen and Williams (2006), Chapter 2.2 of Santner, Williams and Notz (2018) or Chapter 5.3 of Gramacy (2020).



FIG. 2. Ocean example: Predictive mean and standard deviation surfaces for the ocean model output using homGP and hetGP. Data are 1000 simulator runs consisting of 20 replicates at 50 different input locations from (a maximin Latin hypercube of size 50 in 2 dimensions). The top row provides predictive means, μ_N , and the bottom gives standard deviations, σ_N , of the predictive distribution of oxygen concentration. The left column uses homGP, the right uses hetGP.

we first introduce a vital modeling twist to cope with a common feature of stochastic computer simulations.

3.2 Heteroscedastic GP Surrogates

The constant variance Assumption (A1) simplifies the construction of a statistical model because only one intrinsic variance parameter σ_v^2 needs to be estimated. When $\sigma_v^2(x)$ is believed to vary over the input space more must be done. Boukouvalas, Cornford and Stehlík (2014) model $\sigma_v^2(x)$ as $\exp(h(x))$ for simple functions h (e.g., polynomials), a simple extension to assuming just one variance parameter σ_v^2 (the exponential transform ensures positivity of the variance). Like analogous approaches to predicting the mean (briefly discussed in Section 3.1), it is not clear what to use for h, and its simplicity may not meet the complexities found in many applications.

GPs are used for σ_v^2 by several authors (Goldberg, Williams and Bishop, 1997, Kersting et al., 2007, Boukouvalas and Cornford, 2009, Ankenman, Nelson and Staum, 2010, Binois, Gramacy and Ludkovski, 2018). The difficulty is that doing so directly depends on knowing the value of $\sigma_v^2(x)$ at the inputs X_n , but these values are not observed. If there are enough replicated simulation runs, r_i , at the inputs x_i , then the sample variances at the x_i s can be used to estimate the $\sigma_v^2(x)$ at the inputs X_n . Equations (3.1) and (3.2) can then be used to predict $\sigma_v^2(x_{new})$ (working with the logarithm of the sample variances and then exponentiating the results to avoid negative variance predictions). But this approach, called *stochastic kriging* (SK, Ankenman, Nelson and Staum, 2010), is limited by the need for many replicates at each input and the possible inefficiency of treating the variance and mean processes separately.

Those limitations can be removed by considering the intrinsic variances at the inputs, $(\sigma_v^2(x_1), \ldots, \sigma_v^2(x_n))$, as unknown parameters (a.k.a., latent variables) to be estimated in the same manner as all the other unknown parameters. Goldberg, Williams and Bishop (1997) do so in a fully Bayesian, but computationally taxing, way. Reducing these costs forms the essence of approaches by Kersting et al. (2007) and Boukouvalas and Cornford (2009). A recent variant, proposed in Binois, Gramacy and Ludkovski (2018), along with accessible software hetGP (Binois and Gramacy, 2018), tackles the compu-

tational hazards and is the method described and used in this review.

The technical details addressing the computational barriers of a heteroscedastic GP (hetGP) have three elements. One, hetGP models the log variances as the *mean* output of a GP on latent (hidden) variables. The second uses Woodbury matrix identities (Harville, 1997) to reduce computations from treating all N observations to computations involving only the n unique inputs, a reduction of computational complexity from $O(N^3)$ to $O(n^3)$ (especially relevant when there are many replicates). The third element uses MLE to learn all parameters.

While full details are provided by Binois, Gramacy and Ludkovski (2018), some specifics of the description above are worth noting. With $\lambda(x) = \sigma_v^2(x) / \sigma_Z^2$ and $\Lambda_n = (\lambda(x_1), \dots, \lambda(x_n))$ for the *n* distinct inputs, $\log \Lambda_n$ is taken to be the predictive mean of a GP on latent (hidden) variables, $\Delta_n = (\delta_1, \dots, \delta_n)$. For ease of exposition, assume the GP has 0-mean (a constant mean is actually the default setting in hetGP) and take the covariance matrix for Δ_n to be $\sigma_g^2(C_g + gR^{-1})$ where g > 0, $R = \text{diag}(r_1, \dots, r_n)$, and C_g is a correlation matrix (obtained using a correlation function with parameters θ_g). Then $\log \Lambda_n = C_g (C_g + g R^{-1})^{-1} \Delta_n$, with Δ_n to be learnt. This latent Δ_n approach facilitates smooth estimates of Λ_n and provides a fixed functional form for $\lambda(x)$, but does not incorporate the resulting uncertainty. Given Λ_n , the Woodbury identities (Harville, 1997) reduce the likelihood of Y_N , the output at *all* inputs including replicates, to depend only on quantities of size n. Maximum likelihood estimates for the unknown parameters can then be computed at a cost of $O(n^3)$, as can derivatives further facilitating optimization for maximizing likelihood.

As a side note, heteroscedastic measurement error is sometimes present in spatial statistics models (which are often related to surrogate models); however, we know of no such models, which allow for the full modeling and predictions of the intrinsic variance process in the same way as a hetGP. For example, the model in Nguyen, Cressie and Braverman (2017) allows for non-constant measurement error at different sites, but it does not estimate these measurement errors jointly with the other model parameters, nor does it allow for the prediction of the measurement errors at new unseen sites. This is mostly because there is little interest in predicting the measurement error process in spatial statistics (the "true" underlying signal is the objective), whereas with stochastic simulators the intrinsic variability can be of direct modeling interest.

Fish example. We apply both an ordinary homoscedastic GP (homGP) and a hetGP surrogate to the fish example from Section 2.1. The simulation budget is constrained to 400 runs and focuses on the relationship between the total number, x, of fish in a population and the number, y(x), of fish recaptured in the second round of capture. The total population is an integer between 150 and 4000. The simulator is run 20 times at each of 20 unique x locations in [150, 4000], chosen via a maximin Latin hypercube design (see Section 4). The number of fish counted cannot be less than zero, but the normality assumption would allow negative fish counts, so we square root the simulated output before performing our analysis, squaring the resulting predictions to return to the original scale afterwards. In addition, we estimate the "truth" by generating another data set: replicating 500 times at each of the same 20 sites.

Applying a homGP surrogate with squared exponential correlation function produces the results in the upper left panel of Figure 1; the upper right panel shows the results of hetGP. The predicted intervals for the fish model are obtained in the transformed (square-root) space, and squared to get back to the original space.³ The lower panels are plots with the "true" 2.5%, 50% and 97.5% quantiles superimposed.

The key conclusion is that both homGP and hetGP capture the nonlinear trend (though a bit off in the region near 800). The presence of nonconstant intrinsic variability is clear from the truth plot, with the region near 800 showing higher variability than elsewhere. The hetGP surrogate does not fully capture the nonconstant variability, but it does improve on homGP. Full resolution is largely a matter of simulation budget though alternative designs may further improve hetGP. Our supplementary material includes improved results using the sequential design scheme of Section 4.3. The takeaway message here is that the trend is readily captured by both homGP and hetGP and the presence of heteroscedastic variability favours the use of hetGP (perhaps with added simulations or improved designs).

Ocean example. For the ocean model (Section 2.2), we take each simulation "run" to be the average of 6 simulation runs. The true simulator is known to be nonnormal and this adjustment makes the example more Gaussian. For now, we fix the two diffusion coefficients at $K_x = 700$ and $K_y = 200$, leaving the two spatial coordinates as the only varying inputs. Using 1000 simulations (50 sites each replicated 20 times), we obtain, for surrogates homGP and hetGP, the predictive mean surface and the predictive standard deviation surface (which includes both the uncertainty around the predictive mean and the intrinsic variance estimates σ_v^2). These surfaces are plotted in Figure 2, with the left column for homGP and the right column for hetGP.

³If a large portion of the predictive distribution was negative in the transformed space, these un-transformed intervals would be invalid, but this does not appear to be a problem in our example. Monotonic transforms exist to avoid this problem (Johnson et al., 2018), or sampling could be performed to obtain intervals in the un-transformed space. Predictions in the transformed space are also provided in the supplementary material (Baker et al., 2022).

The mean surfaces for both surrogates are similar. The predictive standard deviation for homGP (bottom-left) is relatively constant across the input area (clearly affected by the constraint that the intrinsic variance σ_v^2 is constant). The standard deviation surface for hetGP is markedly different, which could be evidence that the intrinsic variance is non-constant.

To compare these two predictions, we obtain the "truth" using replicate runs (up to 100,000) of the simulator at 500 sites (chosen via a LHD, Section 4), averaging the replicates at each site to get the true mean and the square deviations from the mean to get the true variance). These are plotted in the Appendix (Section 7) and the supplementary material, and they confirm the presence of non-constant intrinsic variance. Moreover, the standard deviation plot for hetGP exhibits a structure similar to the truth plot, leading to the conclusion that hetGP is the better surrogate for this problem. However, this conclusion comes with a caution: repeating this experiment multiple times reveals a great deal of variability in the standard deviation plot, due to variability in the design and the simulations (discussed further in Section 4.3).

Overall, reliable predictions of the mean are achieved, but the uncertainties are less certain. This is similar to the fish example, and improving the uncertainties would require more simulation. The results point to the superiority of hetGP to homGP, which is confirmed via a numerical comparison in Section 4.3, where a sequential design is also examined and compared.

3.3 Nonnormal Variability

In many applications, assuming the variability v is normally distributed is inappropriate. With the fish example, the number of fish cannot be less than 0. In the Ebola example, even with the inputs x fixed, repeated simulations can lead to two distinct groups of possible infection counts (bimodality). In other simulators, there may be a greater tendency for extreme values (fatter tails) in the distribution of v. With these possibilities, normality can be a strong assumption to be used with caution.

Transformation of the data is a time-honored device that sometimes induces "enough" normality in the data to permit the use of Gaussian-based methodology (as in Section 3.2 for the fish model). For example, Henderson et al. (2009) uses the logit transformation $(\log y/(1 - y))$ in analyzing the proportion of deletions in mitochondrial DNA. Plumlee and Tuo (2014) take a different route by focusing on the quantiles of the output distribution— normality is not needed. Both of these approaches have the appeal of leading to relatively simple modifications of the methods in Sections 3.1 and 3.2.

There are also other, more complex, methods that generally lack the same ease of implementation. For example, Moutoussamy, Nanty and Pauwels (2015) attempt to model the underlying probability density function itself, rather than the output y. Xie and Chen (2017) devise a Student *t*-process that is not much different than the GP process while at the same time allowing heavier tails in the distribution of the data.⁴

3.3.1 *Quantile kriging*. Quantile Kriging (QK) is a popular tool for the emulation of stochastic computer models (Rannou et al., 2002, Plumlee and Tuo, 2014, Zhang and Xie, 2017, Fadikar et al., 2018). These approaches are a natural extension of spatial kriging formulations (Zhang, Cressie and Craigmile, 2008, Zhou, Chang and Fuentes, 2012, Opitz et al., 2018) used in environmental applications.

The QK method directly models specific quantiles of interest, such as the median and the lower/upper 95% quantiles at each input. Minimal assumptions are required. $Q_q(x)$, the qth quantile of the simulator output at input x, is modeled with a GP. Given values $Q_q(x_i)$ at inputs x_1, \ldots, x_n , the quantile, $Q_q(x_{new})$ for x_{new} , can be predicted using equations (3.1) and (3.2). This framework allows the distribution of the variability v to take on almost any shape. Although a true generative process for the output y is lost, its distribution can still be described.

To implement QK, values of the targeted quantiles at the inputs are needed. Just as in Section 3.2, where sample variance estimates are used, sample quantiles can be used here given enough replicates r_i at each x_i . The GPs used to predict new quantile values, $Q_q(x_{new})$, should also include a noise term σ_q^2 to acknowledge that the sample quantiles are estimates. Assuming the variability of the sample quantiles is normally distributed may also be invalid, but this is a level further removed from the quantity of interest, y, and can be acceptable in practice.

As a useful modification, the quantile q can be included as an additional input to the GP model. The quantile $Q_q(x)$ can be reformulated as Q(x,q), increasing the dimensionality of the inputs from d to d + 1. This strategy allows for the prediction of Q(x,q) for any desired quantile q, not just those that were empirically estimated and is used by Fadikar et al. (2018) for the Ebola model.

Alternative QK-based approaches are also under development. For example; a promising variant of QK called Asymmetric Kriging (AK, Zhang and Xie, 2017) does not require sample quantiles by leveraging quantile regression techniques (Koenker and Bassett, 1978).

Fish example. For the fish simulator, QK is implemented with the same simulated data set as before (possible because many replicates were obtained). The sample 5%, 27.5%, 50%, 72.5% and 95% quantiles at each of the

⁴The hetGP package also implements a Student-*t* variant (Wang, Shi and Lee, 2017, Shah, Wilson and Ghahramani, 2014, Chung et al., 2019).

20 population sizes form the observed data, and the modification using the quantile q as an added input dimension is adopted. Figure 1 presents the predicted Q(x,q)mean for 5 different quantiles along with the data (the left plot) and compares the "true" values with predictions at the 5%, 50% and 95% quantiles (the right plot).

The center purple curve in Figure 3 is the predicted median. The outer red lines are the predicted 5% and 95% quantiles; the inner blue curves are the predicted 25% and 75% quantiles. The nonmonotone "wavy" lines for the 5% and 95% quantiles reflect the natural variability of extreme quantiles based on only 20 observations. Without an abundance of replicates, accurately capturing extreme quantiles is difficult, a drawback of QK. The other quantiles presented display more regularity.

The results from QK do not differ much from those in Figure 1 where the square-root transformation was sufficient. With more complex problems, such as the Ebola model, the method is suitable while other approaches may be less so. In any case, QK can be a good robust choice, given adequate data for estimating quantiles.

3.4 Multiple Outputs

The discussion so far has assumed that the simulator outputs a single scalar quantity of interest. For multivariate output, a more comprehensive model would be ideally used.

In the geostatistical literature—typically focused on 2d spatial applications—multivariate methods incorporating cross-correlation between outputs are often called cokriging (Cressie, 1993). Adapting these to multivariate simulators must consider higher dimensional inputs, computational tractability and output smoothness. Specific features of the simulator (e.g., hierarchies, dynamics, no missing outputs) have allowed some such adaptations (Kennedy and O'Hagan, 2000, Conti and O'Hagan, 2010, Fricker, Oakley and Urban, 2013, Paulo, García-Donato and Palomo, 2012).

Bespoke, problem-specific formulations for time-series and other outputs have also been entertained (Farah et al., 2014, Sun et al., 2019).

Though less than ideal, more general methods can also be employed with good effect. If there are a small number of outputs, treating each independently, with its own surrogate model, often suffices. This method can be effective, despite ignoring any correlation between the different outputs (and thus wasting information). For example, Spiller et al. (2014) deploy independent surrogates at each of a multitude of sites in a region to good effect.

Alternatively, by treating the index, t, of the T outputs as an additional *input* dimension (changing the dimension of the input space from d to d + 1) a GP surrogate on d + 1 input dimensions can be formed (Bayarri et al., 2009). This method allows correlation structures between the different outputs to be modeled. This is similar to the QK modification where quantile levels are treated as an added input (Section 3.3.1). A drawback of this technique is that, if T is very large, computational issues will arise because the GP must be trained on NT data points rather than just N. Intrinsic variability prevents simplifications of the sort used in Bernardo et al. (1992) for deterministic simulators in this setting.

A different approach reduces the the size of T to a smaller K_0 by representing the multivariate output through the use of basis functions, $\psi_k(t)$:

(3.5)
$$y(x,t) = \sum_{k=1}^{K_0} w_k(x)\psi_k(t) + \delta(x,t).$$

Coefficients $w_k(x_i)$, $k = 1, ..., K_0$ are determined by the data, and $\delta(x, t)$ is the residual error between the basis function representation and the data y. If $K_0 = T$, then $\delta = 0$. Typically, K_0 is taken to be much less than T, but large enough so that the error, δ , is sufficiently small. Each $w_k(x)$ can then be independently modeled with a surrogate and predictions for y(x, t) are obtained from equation (3.5), ignoring δ .

Different choices for the bases can be appropriate in different settings. For example, Bayarri et al. (2007a) use wavelets for the ψ_k s in a deterministic setting where *t* is time. A common choice of basis functions are principal components: the ψ_k s are the eigenvectors of the matrix $Y_N^{\top}Y_N$, the first K_0 corresponding to the first K_0 eigenvalues in decreasing order. Often, the first few (five or less) principal components are enough to capture sufficient information about the full (*T*) data set. Coefficients $w_k(x_i)$ are then equal to $\sum_{t=1}^{T} y(x_i, t)\psi_k(t)$. More information about principal components can be found in Jolliffe (2011) and software for obtaining ψ_k and w_k is prevalent.

Further discussion about using principal components to model high-dimensional simulator output can be found in Higdon et al. (2008). Principal components are also utilized in Fadikar et al. (2018) to model the time-series output of the stochastic Ebola simulator. While principal components are a common default, there is concern that key features of the data set may be left within the discarded $\delta(x, t)$ preventing reliable prediction. Salter et al. (2019) document these concerns with regards to calibration and suggest an alternative.

For problems with functional outputs, with potentially missing data and/or irregularly spaced data (such as irregularly spaced timesteps or spatial locations), a functional decomposition can also be useful. For example, Ma et al. (2021) use functional principal component analysis to model satellite observation simulations.

Overall, multiple output analysis lies on the frontier of research, especially for stochastic simulations (e.g., capturing dependence in the variability between different outputs is one future research direction). The techniques discussed are a start.



FIG. 3. Fish example: Same setup as Figure 1 but with a QK surrogate. Mean predictions of 5 quantiles (5%, 25%, 50%, 75% and 95%) are provided on the left along with data, and mean predictions of 3 quantiles (5%, 50% and 95%) are provided on the right along with the "true" values.

4. EXPERIMENTAL DESIGN

For an experiment, the design (the choice of x values) and analysis (the assessment of the output y(x)) are, in principle, closely connected. Other considerations can also enter. For physical experiments, controlling for external influences or nuisance factors by blocking and randomization is often a vital part of the design. External influences are absent in computer experiments and so controlling for nuisance factors is usually irrelevant. However, many minor parameters are often fixed which could instead be randomized over, with a consequent addition to intrinsic error.

Extensive study of deterministic computer experiments has lead to the recommendation of readily computed "space-filling" designs where no large region of the input space is missed. Multiple methods exist for obtaining space-filling designs, the most popular being Latin hypercube designs (LHDs, McKay, Beckman and Conover, 1979).⁵ LHDs have proved adequate, especially when joined with an additional criterion, such as the maximin criterion, where one also maximizes the minimum distance between points in the design.⁶ Sobol sequence designs (Sobol, 1967) provide an alternative where the xsare generated sequentially making it easy to retain the space-filling character when additional simulations are made at a later stage.⁷ Pronzato and Müller (2012) have a lengthy discussion of these and other space-filling methods, some pertinent to nonrectangular input spaces.

For stochastic simulators, the picture is less clear. The presence of intrinsic variability raises the complication of replication, which is not present in deterministic experiments. With the same inputs, a stochastic simulator can be run multiple times (replicated), providing different output values each time due to the intrinsic randomness. Replicates obviously have an effect on the estimation of the intrinsic variance, σ_v^2 , and, therefore, on prediction (see Section 3.2), and so the number and location of replicates are important. A simple approach is to use a space-filling design to establish the sites $X_n = (x_1, \ldots, x_n)$ of the experiment and then add replicates at each site. Determining the number, r_i , of replicates at each site x_i and how to budget between replicates and sites is not well understood. In fact, there is limited theoretical evidence of the need for replicates altogether (unless the surrogate explicitly needs them), although there is numerical evidence and wide belief that replicates can be advantageous (at least in appropriate contexts). For example, Wang and Haaland (2019) produce designs by minimizing bounds on the integrated mean squared prediction error (IMSPE).⁸ Their numerical results show no need for replicates unless $\sigma_v^2(x)$ is large compared to σ_Z^2 (the uncertainty around the mean, *M*).

The presence of intrinsic variability suggests there is value in multistage designs where stage 1 is used to get information about $\sigma_v^2(x)$ and later stages exploit this information to allocate replicates and select new inputs. Questions arise as to how inputs should be selected for stage 1, and also how to properly leverage the results from stage 1 to select new inputs and replicates. These two factors, replication and multiple stages, are central to developing effective design strategies.

⁵A Latin hypercube design is one where: in each dimension the input space is divided into intervals and each interval is constrained to contain exactly one data point.

⁶Such maximin LHDs are produced, for example, by the maximinSLHD function of the R package SLHD (Ba, 2019), or the lefthand side function from the Python package pyDOE (Lee, 2015).

⁷In R, the sobol function in the R package randtoolbox (Chalabi et al., 2019) can be used to generate Sobol sequences.

⁸With the predictive variance, $\sigma_N^2(x)$, the IMSPE, of a design *D* is equal to $\int_{x \in X} \sigma_N^2(x) dx$.

4.1 Single-Stage Design

A common approach in single-stage studies is to use space-filling designs for inputs, say n in number, and rreplicates at each input, sometimes with no repeats, that is, r = 1. Predictions follow as described in Section 3 depending on the particular prediction model selected. Choices have to be made about the total number of runs and the number of replicates at each input site (N = nr). Often, N is a question of budget, but there is little insight into how r should be chosen except when meeting a specific surrogate model requirement, as in SK (Section 3.2).

For their single-stage study, Marrel et al. (2012) use a standard LHD with no repeats to compare the performance of different statistical models. On the other hand, Plumlee and Tuo (2014) use a LHD with varying numbers of replicates r_i at each x_i . In their case, the number of replicates *must* be large, because the QK method (Section 3.3.1) depends on computing quantiles of the output $y(x_i)$ at each input site of the design.

4.2 Two-Stage Design

The case for a two-stage design is largely to enable estimation of σ_v^2 at stage 1 and use it for the second stage. Ankenman, Nelson and Staum (2010) provide one solution in the context of SK. A first-stage design chooses the x_i s via an LHD of size n_1 with a common number, r, of replicates at each of the inputs, resulting in a total number of $N_1 = n_1 r$ runs at stage 1. The first-stage analysis uses the r replicates at each input to estimate $\sigma_v^2(x_i)$ using the sample variances. As outlined in Section 3.2, a GP (working with $\log s^2(x_i)$) is then used to produce "plugin" estimates of $\sigma_v^2(x)$ for all x. A different GP uses these variance estimates to build a predictor for the mean output M.

For stage 2, n_2 additional unique input locations are chosen so that the combined set of design locations, $X_n = (x_1, ..., x_n)$, remains space-filling. The IMSPE is then calculated using the GP model constructed in stage 1. Minimizing the IMSPE with respect to the number of replicates $R_n = (r_1, ..., r_n)$ provides the optimal number of replicates for the chosen X_n . Details are in Ankenman, Nelson and Staum (2010). One difficulty is that the optimal R_n might produce an r_i for a first-stage site that is smaller than the *r* already obtain in stage 1. Some modification to the design is then necessary.

In this setting, a Sobol sequence could also be used to obtain a design that is space filling at both stage 1 and stage 2. This is not what is done in Ankenman, Nelson and Staum (2010), but a Sobol sequence is easier to implement and likely to yield similar results. Choosing the unique inputs X_n for stage 2 by optimizing the IMSPE could also be done, but this adds to the computational burden. Suitable recommendations for the values of n_1 , n_2 , N and the replicates at each distinct input are lacking (in

Ankenman, Nelson and Staum (2010) the recommendations are ad hoc) and, as for one-stage experiments, open for study. A third-stage design (or indeed, any multistage design) can be constructed by repeating stage 2 in this process.

4.3 Sequential Design

In some cases, it can be feasible to carry out a sequential process where, after the first stage, simulation runs are chosen one-at-a-time. After each run, the relevant quantities can be updated before determining the next run. This can address the issue of learning about σ_v^2 and choosing new runs, without prespecifying the proportion of replicates. An advantage of a sequential design is the possibility of stopping when a criterion is met before a budget constraint is reached. An additional advantage is the increased likelihood of performing useful simulator runs, replicates or otherwise. For some objectives, such as optimization (Section 6.3), a sequential design is usually essential. For global prediction, Binois et al. (2019) present one sequential design, implemented in the previously mentioned hetGP package.

The strategy in Binois et al. (2019) begins at stage 1 with a space-filling design D_1 of n_1 inputs and an allocation of runs $(r(x_1), \ldots, r(x_{n_1}))$. Using a GP for M and a latent GP prior on σ_v^2 (as in Section 3.2), estimating the parameters by MLE leads to a calculable estimate of IMSPE (D_1) . A new point z is considered, either as a new unique input x_{n_1+1} or as a replicate of an existing input in D_1 . Selection z is added to the design D_1 if z minimizes IMSPE $(D_1 + z)$, yielding a new design D_2 . This myopic rule can be iterated, and each time a new point is added, the surrogate is updated. The process stops when a criterion is met or the computational budget exhausted.

Computational viability is strained by the updating required after each run. On the other hand, the computational burden is less than it could be—by being "greedy" it only seeks the optimal data point for the very next simulator run, ignoring runs that may be better in the long run.

This is not the only sequential design scheme available for global prediction problems. For example, the treegenerating processes used in TGP and BART (see Section 6.1) can lead to specialized sequential design strategies. Details are available in Gramacy and Lee (2009) and Chipman, George and McCulloch (2010).

Blurring the lines between multistage and sequential designs, it can sometimes be practical to run additional simulations in batches (e.g., as in making efficient use of a multicore supercomputer). In such circumstances, a "batch design" would be desirable. These have been developed for deterministic experiments (Loeppky, Moore and Williams, 2010, Duan et al., 2017, Erickson et al., 2018), but not yet explicitly extended to stochastic cases.



FIG. 4. Ocean example: Predictions for the ocean model output using seqhetGP. Left plot is the mean, μ_N , of the predictive distribution; right plot the standard deviation, σ_N . Design sites with their replicates are superimposed on the right-hand plot.

When fully sequential methods are feasible, the seqhetGP strategy sketched above is valuable. There are several aspects worth considering with sequential designs:

- The extensive use of a surrogate in the construction of the design may require scrutiny by diagnostics that assess the quality of the surrogate.
- The first stage of a sequential strategy should avoid a poor (e.g., too small) initial design lest a poor starting surrogate leads to poor choices thereafter.
- The utility of a sequential design depends on the relative cost of implementation compared to a simulator run. For challenging problems, simulator runs are likely to be costly enough to make sequential design attractive.
- There may be modifications to a sequential design that reduce computational load without paying a significant cost in accuracy. For example, by reestimating parameters periodically rather than after each step.

Ocean example. For the ocean model, we implement the sequential design scheme from Binois et al. (2019)using an initial design of 50 sites (chosen by a maximin LHD in 2-d), giving each site 5 runs. The remaining 750 data points are then assigned via the sequential scheme. The resulting mean and standard deviation surfaces are in Figure 4. For the standard deviation surface, the design sites are superimposed along with the number of runs taken at each site.

The mean surface in the left panel is slightly different than for the nonsequential analyses (Figure 2, top row). The standard deviation surfaces look very different. For the design itself, new inputs are heavily replicated in regions where the standard deviation is larger, and less so in regions where it is smaller. Additionally, the sequential design includes more unique sites than the fixed design, and more points on the boundaries of the input space.

Using the "truth" established in Section 3.2, we can compare the performances of the three methods. As dis-

cussed previously, the visual presence of heteroscedasticity is a reason to avoid homGP. Visually distinguishing between the performances of the hetGP and seqhetGP surrogates is more difficult: the means appear similar, and while some patterns in the true standard deviation appear to be captured by hetGP, imperfections are visible and the magnitude is not always correct. With the seqhetGP standard deviation, a lot of nuance seems lost. To properly compare the different methods, a numerical comparison can be more valuable.

Two useful numerical measures are root mean squared error, RMSE (the square-root of the average squared difference between the surrogate's prediction of the mean and the "true" mean) and Score (the proper scoring rule from equation (27) in Gneiting and Raftery, 2007). RMSE measures the accuracy of the mean predictions and Score is an overall measure testing the accuracy of the combined mean and variance predictions. With a test set of inputs x_1, \ldots, x_p and simulator outputs y_1, \ldots, y_p , surrogate predictive means $\mu_N(x_1), \ldots, \mu_N(x_p)$ and variances $\sigma_N^2(x_1), \ldots, \sigma_N^2(x_p)$, Score is

(4.1)
$$\frac{1}{p} \sum_{i=1}^{p} \left(-\left(\frac{y_i - \mu_N(x_i)}{\sqrt{\sigma_N^2(x_i)}}\right)^2 - \log(\sigma_N^2(x_i)) \right).$$

Smaller RMSE is better while for Score, larger is better.

For the three methods, the RMSE for homGP, hetGP and seqhetGP are respectively 2.056, 1.985 and 1.567; and the Scores are respectively -3.999, -3.880 and -3.834. The RMSE results reveals that seqhetGP is best at predicting the mean, which was not obvious from the plots. The Scores for hetGP and seqhetGP are close but both noticeably better than homGP, affirming the presence of heteroscedasticity.

The randomness in stochastic simulators, as well as variability in design (there are many possible maximin LHDs), can induce a large degree of variability in specific results (such as those just cited). It is therefore difficult to rely on a single result for making general comparisons. As such, the above experiment is repeated 100 times and the resulting 100 RMSEs and Scores are summarized in boxplots in Figure 5.

The boxplots confirm what was found with the single data set: heteroscedasticity is present and seqhetGP is preferred. Visual inspection of many of the standard deviation plots for the repeated experiments (as discussed in Section 3.2 and found in the supplement) reveals considerable variation and departure from the true standard deviation. We find the variance (the intrinsic variance and the GP uncertainty for the mean) can be hard to get right without an abundance of data, and the difficulty is compounded by the use of plug-in estimates whose uncertainty is not accounted for.

4.4 Designing for Statistical Model Parameter Estimation

Sections 4.2 and 4.3 construct designs that rely on a surrogate model based on stage 1 data in order to choose subsequent data points. The quality of the designs depends on the accuracy of the surrogate which, in turn, depends on the accuracy of its parameters. An alternate approach to those used in Sections 4.2 and 4.3, is to build a design with the express purpose of better estimating these parameters.

Boukouvalas, Cornford and Stehlík (2014) address this idea, using a simple parametric function for the variance $(\sigma_v^2(x) = \exp(h(x)))$, where *h* is a simple function (e.g., a polynomial). They propose designs that maximize a criterion previously used for deterministic simulators by Abt and Welch (1998): the logarithm of the determinant of the Fisher information matrix, $\log |I|$. Numerical results suggest this method gives improvements in estimating the parameters, but overall global prediction is no better, and sometimes worse, than using a space-filling design. When prediction is of prime importance, the question arises about how to make use of such designs for stage 1 in a multistage or sequential setting, where its impact on obtaining better parameter estimates might be felt. For example, see Zhang, Cole and Gramacy (2021).

5. CALIBRATION

Calibration is needed when there are inputs to the simulator that are neither known nor measurable, which is a common condition in practice. Transmissibility in the Ebola simulator and the diffusion coefficients in the ocean model are examples of such inputs. In order to infer (indirectly) values for these inputs and produce predictions, added information in the form of field data (experimental or otherwise) are necessary. Inclusion of field data and calibration parameters, labeled u_C , leads to the observation model:

(5.1)
$$y_F(x) = y_S(x, u_C) + \delta_{\text{MD}}(x) + \epsilon,$$

where $y_F(x)$ are real-world field observations at controllable (or measurable) inputs x, y_S is the simulator with additional unknown, nonmeasurable, inputs u_C , ϵ is measurement error for the observations $y_F(x)$ (with variance σ_{ϵ}^2) and $\delta_{\text{MD}}(x)$ is an important term that accounts for the simulator not being a perfect representation of reality. y_F "observes" reality with error ϵ ; reality = $y_S + \delta_{\text{MD}}$.

Multiple competing methodologies and even philosophies exist for calibration. Several solutions to the calibration problem are outlined below. Despite the centrality of calibration in computer experiments, comprehensive comparisons are lacking.

5.1 Kennedy–O'Hagan Calibration (KOH)

The formulation in equation (5.1) was made by Kennedy and O'Hagan (2001) for deterministic simulators and is the basis for much of the calibration and related prediction work since. The strategy pursued by Kennedy and O'Hagan (2001), as implemented in Bayarri et al. (2007b), obtains a surrogate for y_S and models $\delta_{MD}(x)$ with a GP (although other choices are possible). After replacing y_S with the surrogate, posterior distributions for all unknowns can be obtained via a Bayesian analysis. In practice, the surrogate model is fit only using the simulator data, ignoring possible influences from the field data. Details and discussion of this modular approach can be found in Bayarri et al. (2007b) and Liu, Bayarri and Berger (2009).

The KOH approach emphasizes the need to address calibration and model discrepancy together. Confounding between u_C and $\delta_{MD}(x)$ inevitably occurs because there are multiple combinations of u_C and $\delta_{MD}(x)$ that result in the same observed field data. As such, u_C is nonidentifiable and its estimation is compromised, as is the discrepancy. Nonetheless, the resulting *predictions* for y and E(y) are sound, even if the individual estimates for u_C and $\delta_{MD}(x)$ are not. For details and further discussion, see Higdon et al. (2004), Bayarri et al. (2007b), Brynjarsdóttir and O'Hagan (2014) and Tuo and Jeff Wu (2016).

Multiple attempts to circumvent confounding have surfaced. Tuo and Wu (2015) alleviates the ambiguity in u_C by formally defining it as a least-squares quantity; Gu and Wang (2018) propose novel priors for the discrepancy that compromise between the Tuo and Wu (2015) strategy and KOH; and Plumlee (2017) introduces priors on the discrepancy that are orthogonal to the prior mean. In the stochastic simulator literature, Oakley and Youngman (2017) removes δ_{MD} but compensates by inflating the variability in the simulator output. Ignoring δ_{MD} altogether can sometimes be justified by strong evidence of the simulator being accurate, but such evidence is rare.

For stochastic problems, where reality is stochastic, the discrepancy term $\delta_{MD}(x)$ cannot be assumed deterministic. Discrepancy in stochastic settings is an open research



FIG. 5. Ocean example: Performance results of the three ocean model surrogate fits, repeated 100 times. The boxplots are for computed RMSE and Score, for each repetition, at the 500 test locations.

question, with little attention so far. The model for the discrepancy may need to be similar to the model for the simulator; for example, if modeling y_S calls for a hetGP with a Matérn 5/2 correlation function, then it is possible that a hetGP is needed for the discrepancy as well (perhaps with a smoother squared exponential correlation). A full Bayesian analysis in such circumstances may be prohibitively expensive and the above procedure might need to be modified. Sung, Barber and Walker (2019) use a hetGP for the discrepancy (but with a deterministic simulator), estimating parameters via maximum likelihood and following Tuo and Wu (2015) to avoid confounding.

Revisiting Ebola. The Ebola study (Fadikar et al., 2018) calibrates an ABM using the KOH framework. The simulator y_S has 5 unknown, unmeasured inputs u_C , and the output is the log of the cumulative number of infected individuals up to week 1 and every week thereafter up to 57 weeks. The field data y_F is a set of reported cumulative counts. For the statistical model, a QK strategy (Section 3.3.1) is followed by replicating each distinct simulation 100 times and then calculating the 5%, 27.5%, 50%, 72.5% and 95% quantiles at each time step. These quantile output trajectories are then reduced to a more manageable 5 dimensions using the principal component decomposition outlined in Section 3.4, which are then used to fit the QK model.

Underlying the approach is an assumption that the epidemic trajectories (actual and simulated) can be approximated by quantile trajectories (i.e., a realized epidemic that resembles the qth quantile at time 1 will also resemble the qth quantile at a later time). As such, the quantile q is included as an input parameter (see Section 3.3.1) to allow KOH calibration to learn about the (unknown) value of q, as well as the 5 calibration parameters, for the observed epidemic. The simulator and reality quantile trajectories are deterministic and unknown, so the discrepancy is also deterministic. Posterior distributions for unknown u_C , δ_{MD} , and q are then obtained and used to make predictions of the cumulative counts and other quantities.

In the main analysis, which restricts the field data to only the first 20 weeks, the estimate of model discrepancy is found to be almost zero. A subsequent analysis done using field data up to week 42 exposes some inaccuracy of the simulator (nonzero $\delta_{MD}(x)$). This reveals that, for this example, the simulator is flawed because it continues to predict infections even after the epidemic dies in reality (and so a discrepancy term is important). Extrapolation to later weeks using only the first 20 weeks of field data would be misguided because the simulator flaw would not be identified. This is a general issue for simulators: extrapolation can be tricky and a degree of faith is needed in the simulator.

Ocean example. The previous ocean analyses fixed the two diffusion coefficients. In reality, they are unknown and calibration is necessary. "Field" data are artificially created by averaging over 200 simulations at 150 different longitude–latitude coordinates, using the previously fixed values of the diffusion coefficients ($K_x = 700$ and $K_y = 200$). "True" values are obtained by adding a fake discrepancy, taken as a single realization from a GP with a squared-exponential correlation function, a variance of 1.64, and θ values of (1, 2) (equation (3.3)). To these, normally distributed pretend "observation errors" with a variance of 4 are added, two such observations at each site. In real problems, the field data would be observed and not generated like this. Note that field data for this problem

corresponds with the mean of the simulator, not individual draws from the simulator; a result of this simulator being a stochastic approximation.

With the diffusion coefficients now uncertain, the simulator has four inputs. A computer experiment is designed with runs at the 150 sites used for the field data and 500 unique selections of the calibration parameters K_x and K_{y} . This is done by combining copies of the 150 longitude and latitude sites with a size-500 maximin LHD for (K_x, K_y) , and then improving the combined design by maximizing the minimum distance between design points in the 4-dimensional space. We call this set of points $D_{\rm oc}$. The simulator experiment is carried out by taking 10 replicates at each point in D_{oc} . Two distinct surrogates (a homGP and a hetGP) are fit with this fixed design. In addition, a seqhetGP surrogate is constructed, with an initial design of only 4 replicates of D_{oc} and the remainder of the budget assigned following the strategy of Binois et al. (2019), described in Section 4.3.

For a KOH analysis done in modular fashion, the surrogates are fit only using the simulated data. Because reality here is represented by the expectation of the simulator (rather than the simulator output itself), y_S in equation (5.1) is replaced with $E(y_S)$. Similarly, because reality is deterministic, δ_{MD} is modeled as a standard GP. Of course, the simulated data are outputs from y_S , not from $E(y_S)$ —the surrogate is used to approximate the deterministic $E(y_S)$ (M). MCMC is then used to obtain posterior distributions for the remaining unknowns: the diffusion coefficients, K_x and K_y ; the variance and correlation parameters of the model discrepancy GP, $\sigma^2_{\rm MD}$ and $\theta_{\rm MD}$; and the observational error, σ_{ϵ}^2). The obtained posterior distributions for the key parameters are in Figure 6; their true values are $K_x = 700$, $K_y = 200$, $\sigma_{\text{MD}}^2 = 1.64$ and $\sigma_{\epsilon}^2 = 4$.

For all three surrogate models, the posterior distributions for K_x are fairly diffuse. The K_y posteriors are highly concentrated, but not quite around the true value. The three posteriors for observational error are quite similar but all point to estimates closer to 5 rather than the true 4. The posteriors for the discrepancy variance are also diffuse. These plots underline the dilemma of calibration: obtaining accurate values of calibration (and other) parameters in the presence of model discrepancy is difficult. Additionally, with noisy data, it is difficult to obtain precise estimates. However, KOH does yield useful posterior predictive distributions.

Table 1 compares predictions by KOH calibration with 3 other calibration approaches. The first one estimates K_x and K_y by ordinary least squares (OLS): (K_x, K_y) is chosen such that the sum of the squared residual difference between the mean surrogate prediction and the observed data is minimized. New observations are then predicted by running the surrogate with the parameters (K_x, K_y)

replaced by the OLS estimates (\hat{K}_x, \hat{K}_y) . The second approach follows a frequently adopted practice by guessing, or "judiciously selecting," specific values for K_x and K_y . Here, the choices $K_x = 600$ and $K_y = 400$ are made, and then predictions are made using the surrogate. Call this method SINGLE. The third method, NOCAL, generates predictions as if there were no field data and the distribution for (K_x, K_y) is taken as their prior distribution, independent uniform priors on [100, 1000]. In these alternative methods, the observational error variance is fixed at the true value, and 0-discrepancy is assumed (the former is overly generous and the latter is all too common in practice). For NOCAL, a distribution for the oxygen concentration is obtained by sampling values of K_x and K_y from their prior distribution and plugging them into the surrogate, while for KOH, by sampling from the posterior distributions of all unknowns.

Although the differences in RMSE are negligible, the Scores indicate that KOH performs the best. It is also possible that the accuracy of OLS, SINGLE and NO-CAL is overstated, because the observational error variance is taken as known while in KOH it is estimated. That the least squares estimates (\hat{K}_x, \hat{K}_y) are not always close to the true values is unsurprising given the presence of discrepancy, along with possible imperfections and high variability in the surrogate. For similar reasons, scant differences appear among the three surrogates.

The similarity of RMSEs is a consequence of large variability in the surrogate, the presence of discrepancy, the dominance of the longitude and latitude inputs and a weak effect from the calibration inputs. The first explains the magnitude of the RMSEs and the last explains why fairly inaccurate calibration inputs (in OLS and SINGLE) do not matter. Because KOH addresses discrepancy, its Score exceeds the others, showing that accounting for the discrepancy is necessary and cannot be wished away.

5.2 History Matching (HM)

History Matching (HM) is a common alternative to KOH calibration (Craig et al., 1997, Vernon, Goldstein and Bower, 2010, Boukouvalas et al., 2014, Andrianakis et al., 2017). HM searches for inputs where the simulator outputs closely match observed data, while recognizing the presence of the various uncertainties, including model discrepancy. The HM approach rules-out "implausible" inputs in a straightforward way, rather than attempting to find probable inputs. With an observation y_F , and initially assuming u_C makes up all the unspecified simulator inputs, u_C is deemed implausible if

(5.2)
$$\frac{|y_F - \mu_N(u_C)|}{\sqrt{\sigma_N^2(u_C) + \sigma_{\rm MD}^2 + \sigma_{\epsilon}^2)}} \ge 3,$$

where σ_N^2 , σ_{MD}^2 and σ_{ϵ}^2 are the variances of the surrogate, the model discrepancy and the observational error, respectively. In other words, an input is implausible if the difference between the observation and the simulator output



FIG. 6. Ocean example: KOH calibration results for the ocean model. The top row shows the posterior densities for the four parameters using homGP with the fixed design, the middle row uses hetGP with the fixed design and the bottom row uses the hetGP surrogate with the sequential design. The budget for all three is 5000 runs. True values are superimposed as red vertical lines.

(using that input) is sufficiently large relative to those uncertainties. The number 3 comes from Pukelsheim (1994) who shows that at least 95% of any unimodal distribution is contained within three standard deviations. When there are multiple outputs or additional, controllable, inputs there are modifications to equation (5.2) (Vernon, Goldstein and Bower, 2010).

The process can be repeated in so-called "waves," using nonimplausible u_C found at one wave to generate simulation runs for the next wave, sequentially reducing the space where u_C could lie. With these waves, HM aims to avoid regions of inputs where u_C is unlikely to be and, in that regard, HM is a also calibration design scheme. At any given wave, it is possible for all values of u_C to be deemed implausible—the so-called terminal case (Salter et al., 2019)—usually implying that σ_{MD}^2 is set too low or that the simulator is not fit for purpose. Andrianakis et al. (2015) contain a thorough description of HM whilst applying it to a complex epidemiology model of HIV.

HM and KOH. With KOH, the estimation of u_C is confounded with discrepancy, but predictions and their uncertainties are available. However, implementing KOH in complex problems may be burdensome if not intractable. Speculatively, a hybrid strategy may be to use HM to reduce the input space, confirm the absence of the terminal case, and then apply KOH in the narrowed space to get predictions and uncertainties. Complex models, unlike the fish and ocean examples in this review, would be ones for which this approach would be most appealing. Such hybrid strategies are a topic for further exploration.

| TABLE | 1 |
|-------|---|
|-------|---|

Ocean example: Performance results of the three ocean model surrogates under KOH calibration. RMSE at the 500 test locations with the "true" values used for Figure 5; similarly for Score. (\hat{K}_x, \hat{K}_y) are least squares estimates for (K_x, K_y) , OLS presents the predictive results from least squares calibration, SINGLE the results from arbitrarily choosing (600, 400) for the diffusion coefficients, NOCAL the results from sampling the prior for (K_x, K_y) and KOH the results from performing KOH calibration

| | | (\hat{K}_x, \hat{K}_y) | OLS | SINGLE | NOCAL | КОН |
|-------|----------|--------------------------|-------|--------|-------|-------|
| RMSE | homGP | (824.9, 295.4) | 9.16 | 9.16 | 9.22 | 9.14 |
| | hetGP | (754.9, 295.8) | 9.16 | 9.16 | 9.20 | 9.15 |
| | seqhetGP | (496.3 276.0) | 9.15 | 9.15 | 9.22 | 9.16 |
| Score | homGP | | -2.50 | -2.66 | -2.59 | -2.32 |
| | hetGP | | -2.55 | -2.71 | -2.62 | -2.32 |
| | seqhetGP | | -2.55 | -2.69 | -2.61 | -2.30 |



FIG. 7. Fish example: ABC calibration directly (left) and via hetGP surrogate (right) for the fish model. The prior for the true population size is uniform on $\{200, \ldots, 4000\}$ and 25 recaptured fish were observed. The left plot shows 10,000 simulations, highlights the 52 in agreement with the observation, and shows the histogram of accepted simulations. The histogram in the right panel is for the 3811 accepted draws out of 1,000,000 from hetGP.

5.3 Approximate Bayesian Computation (ABC)

ABC methods offer alternatives which have been useful in moderately complex contexts (Rutter et al., 2019); but less so in more ambitious settings (McKinley et al., 2018).

ABC is a general method for producing samples from $\pi(u_C|Y_F)$, the posterior distribution of unknowns u_C , given data Y_F . ABC does this by generating samples for the unknowns $u_C^{(s)}$ and the output $z^{(s)}$ from $\pi(Y_F|u_C) \times$ $\pi(u_C)$, that is, from the likelihood of the data given the unknowns, multiplied by the prior probability of the unknowns. For computer models, generating samples from the likelihood is equivalent to running the simulator. Such samples are only accepted if $z^{(s)} = Y_F$. For continuous settings, where exact equality cannot occur, acceptance is instead made if $B(z^{(s)}, Y_F) < \tau$, where B is a measure of distance and τ a level of tolerance. An approximated posterior distribution is then given by the collection of accepted $u_C^{(s)}$ s. When there are multiple outputs (or there are other controllable inputs x, and so for any given $u_C^{(s)}$ there are effectively multiple outputs), Y_F and $z^{(s)}$ can be replaced with informative summary statistics. Finding a single statistic sufficient for all outputs is challenging, and a poorly chosen one can invalidate results.

The choice of the tolerance τ is also important. If τ is small, then it may take a very long time to generate a single sample which satisfies the inequality. If τ is not small, then the approximation to the posterior is less reliable. For calibration, τ can be interpreted as a bound on the observational error and model discrepancy, leading to a "correct" posterior rather than an approximation (Wilkinson, 2013). This is then similar to HM with the subjective choice of bounds.

ABC can be done without the use of a surrogate, but this will require many runs of the simulator itself. Otherwise, very few accepted u_C will be obtained, or an overly high value of τ will be required. In either case, accuracy is compromised. Such computational barriers are alleviated by the use of a surrogate.

Fish example. Here, we apply ABC to the fish simulator in order to estimate how many fish are in the total population. Suppose that 25 fish are recaptured in the second round. A straightforward method to determine the total population size is to simulate many times from the NetLogo fish model for many different values of the total fish population, and "accept" every simulation that leads to 25 fish being recaptured. This is exactly ABC, and is a fairly common practice with ABMs. Simulating 10,000 times, using a uniform prior on the integers between 200 and 4000 (so each such population size has prior probability 1/3801), yields the results in the left panel of Figure 7. This direct use of the simulator produces only 52 accepted samples, which is a very small number, and this is from 10,000 simulator runs. In comparison, the hetGP surrogate fit from only 400 simulator runs (from which 1,000,000 samples can be quickly drawn) yields 3811 accepted samples. This result, illustrated in the right panel of Figure 7, gives a less noisy histogram with the same overall shape. If the simulator is even marginally costly, then a surrogate is unquestionably valuable for ABC computations.

5.4 Related Calibration Techniques

The three calibration techniques above are the more popular techniques in the literature, but others also exist.

Bound-to-Bound (Frenklach et al., 2016) is akin to HM, where an error bound that sweeps up all uncertainties

is similarly defined and quadratic programming is then used to find feasible bounds for u_C . Bayesian melding (Poole and Raftery, 2000, Raftery, Givens and Zeh, 1995) is a technique related to Bayesian calibration, used to reconcile differences between elicited prior distributions on inputs and outputs of a simulator. It has been applied in ecology, epidemiology, urban modeling and pollution monitoring (Ševčíková, Raftery and Waddell, 2007, Alkema, Raftery and Clark, 2007, Radtke, Burk and Bolstad, 2002, Fuentes and Raftery, 2005).

6. OTHER METHODS AND OBJECTIVES

Here, we briefly outline some other surrogate modeling and downstream tasks.

6.1 Regression Trees

In some situations, the simulator mean M may have discontinuities or "regime changes," where a very different relationship between y and x exists in one part of input space compared to another part (i.e., nonstationarity). Regression trees (Breiman et al., 1984) form a class of methods that can be useful in these situations. They are also useful in contexts where some inputs are categorical rather than numerical. The problems are treated by dividing the input space into mutually exclusive regions within which independent surrogates (GPs or other regression methods) are fit.

Two approaches: the trees GP (TGP, Gramacy and Lee, 2008) and Bayesian Additive Regression Trees (BART, Chipman, George and McCulloch, 2010) have found wide application. Both use the data to automatically partition the input space, rely on Bayesian computation and have publicly available software: TGP in tgp on CRAN (Gramacy and Taddy, 2016, Gramacy, 2007); BART in several R packages, including BayesTree (Chipman and McCulloch, 2016) and BART (McCulloch et al., 2019).

Other approaches by Rullière et al. (2018), and via Voronoi tessellations instead of trees (e.g., Kim, Mallick and Holmes, 2005, Rushdi et al., 2017, Park and Apley, 2018), have received less attention. Pratola et al. (2020) extends BART to heteroscedastic σ_v^2 (HBART) by modeling *M* as a sum of Bayesian regression trees (as in BART) and the intrinsic variance $\sigma_v^2(x)$ as a product of Bayesian regression trees, in a joint approach similar to that in Section 3.2.

Calibration methods capitalizing on the KOH approach and using TGP are explored in Konomi et al. (2017). In each terminal node of the partition a GP with an independent constant intrinsic variance term is assumed for the computer model output. An independent GP is also deployed for the discrepancy term. Though σ_v^2 is constant at each terminal node the constants can vary across the terminal nodes so heteroscedasticity is automatically incorporated.

6.2 Qualitative Inputs

Categorical (qualitative) variables are often present in stochastic simulators, especially those that incorporate characteristics of human behavior. While regression trees are capable of dealing with categorical inputs (Broderick and Gramacy, 2011, Gramacy and Taddy, 2010), GPs may be more effective as surrogates for smooth simulator output.

Qian, Wu and Wu (2008), Zhou, Qian and Zhou (2011) and Chen, Wang and Yang (2013) describe ways to extend the kernels used for numerical inputs to incorporate qualitative variables. Painting with a broad brush, their approaches take the correlation between two outputs $y(x_i)$ and $y(x_j)$ as the product of two correlation functions: $C_c(w_i, w_j)$ dealing with the continuous inputs, w, and $C_q(z_i, z_j)$ for the qualitative variables, z. A simple way of building C_q takes

(6.1)
$$C_q(w_i, w_j) = \prod_{k=1}^K \tau_{k, w_{ik}, w_{jk}},$$

where *K* is the number of qualitative variables and $\tau_{k,w_{ik},w_{jk}}$ represents the correlation between w_{ik} and w_{jk} . One example of $\tau_{k,w_{ik},w_{jk}}$ is

(6.2)
$$\tau_{j,w_{ik},w_{jk}} = \exp\{-(\phi_{ik} + \phi_{jk})I[w_{ik} \neq w_{jk}]\},\$$

where *I* is the indicator function (= 1 if its argument is true, = 0 if false), and $\phi > 0$. The cited references provide other ways of modeling $\tau_{k,w_{ik},w_{jk}}$. Alternatives also exist; for example, Zhang et al. (2020) make use of latent variables for qualitative models.

6.3 Optimization

A common experimental objective is to maximize an output of the simulator, that is, to find an input x_{max} that maximizes the output y(x). For minimization instead, replace y(x) with -y(x). Optimisation is usually a sequential process where successive xs are chosen to get closer and closer to the optimal x_{max} , which is a sequential design problem (see Section 4). With stochastic simulators, y(x) is random, and optima are less concretely defined because the output is different every time the simulator is run at the same x. As a consequence, interest usually lies in maximizing a nonrandom quantity of interest, such as the mean, M, or a quantity such as the qth quantile.

For deterministic simulators, Bayesian optimization (Mockus, Tiesis and Zilinskas, 1978, Jones, Schonlau and Welch, 1998) is a popular technique. An initial set of runs is used to build a GP surrogate and new runs are chosen by maximizing an "acquisition function" $\alpha(x)$. Iteratively choosing $x_{\text{new}} = \arg \max_x \alpha(x)$) provides a progressively improved estimate for the maximum. A widely used choice for $\alpha(x)$ is the *expected improvement* (EI):

(6.3)
$$\alpha_{\rm EI}(x) = E[\max(y(x) - y_{\rm max}, 0)].$$

Maximizing EI chooses the input x_{new} that maximizes the expected increase in the maximum value, y_{max} , of already observed runs. With y modeled by a GP,

(6.4)

$$\alpha_{\rm EI}(x) = \left(y_{\rm max} - \mu_N(x)\right) \Phi\left(\frac{\mu_N(x) - y_{\rm max}}{\sigma_N(x)}\right) + \sigma_N(x) \phi\left(\frac{y_{\rm max} - \mu_N(x)}{\sigma_N(x)}\right),$$

where $\mu_N(x)$ is the predictive mean of the GP, $\sigma_N(x)$ its standard deviation, ϕ is the standard normal density and Φ the standard normal distribution function.

Alternative acquisition functions have generated extensive work on Bayesian optimization in recent years, mostly in the machine learning literature. The probability of improvement (Kushner, 1964) is an early example, and the GP upper confidence bound (GP-UCB, Srinivas et al. (2012)) considers homoscedastic simulator error. A recent summary of Bayesian optimisation can be found in Frazier (2018).

For stochastic simulators, the EI procedure can be extended by replacing y_{max} with the maximum estimated mean of currently run inputs, $\mu_{max} = \max_{i \in \{1,...,N\}} \mu_N(x_i)$ (Vazquez et al., 2008). Alternatively, one can seek improvement over the maximum estimated mean of any possible input, $\max_x \mu_N(x)$ (Gramacy and Lee, 2011). In these cases, the $\sigma_N(x)$ term must exclude the $\sigma_v^2(x)$ term. Implementation of this method is provided in the hetGP package.

Alternative criteria for stochastic problems with constant intrinsic noise are discussed and compared in Picheny, Wagner and Ginsbourger (2013); with the above method is referred to as the "plugin" method. An R package for implementing several of these choices is available in DiceOptim (Picheny, Ginsbourger and Roustant, 2016, Picheny and Ginsbourger, 2014). Jalali, Van Nieuwenhuyse and Picheny (2017) also do a similar comparison for heteroscedastic noise.

The related goal of level set estimation to find regions where the output exceeds a threshold T can also be targeted with sequential criteria similar to EI. A simple criterion is maximum contour uncertainty (MCU), wherein new points are chosen by weighting the sum of a point's proximity to T with its uncertainty. The method is implemented in hetGP, and Lyu, Binois and Ludkovski (2018) provides some discussion.

Optimization using Gaussian processes, specifically in the presence of intrinsic variability that is potentially heteroscedastic (and potentially nonnormal) is an interesting research question and possibly deserving of its own review. Nonetheless, the references provided here should provide a good introduction.

6.4 Sensitivity Analysis

Determining and measuring the effect of inputs on the output is usually part of any simulator experiment. Doing so assists scientific understanding of the system and enables screening out potentially superfluous variables. This goal has many related names: sensitivity analysis, screening, variable selection, etc., but the overall objective is generally the same—summarize and measure the influence of each input.

For deterministic simulators, Sobol indices (Sobol, 1993) are widely used. Probabilistic distributions are assumed on the inputs of the simulator in order to represent their range of possible values. Then a functional Analysis of Variance (ANOVA) decomposition splits the variation of the simulator output into multiple components, each representing the individual contribution of an input variable x_i or combination of input variables. A Sobol index is then computed as the percentage of the total simulator output variation explained by a component. Key Sobol indices include main effects (the percentage of variation explained by the individual x_i s *alone*) and the variation explained by interactive additive effects with other inputs. Computing the components takes large numbers of runs but the use of surrogates make the calculations feasible (Schonlau and Welch, 2006, Marrel et al., 2009). An enveloping discussion of sensitivity analysis is provided by Oakley and O'Hagan (2004).

Two extensions of Sobol indices by Marrel et al. (2012) and Hart, Alexanderian and Gremaud (2017) for stochastic simulators yield the following expression for the stochastic simulator:

(6.5)
$$y(x) = y(x, \epsilon_{\text{seed}}),$$

where x is the set of controllable inputs. The input ϵ_{seed} is responsible for output stochasticity, standing in for intrinsic variability, and is sometimes called a seed variable. As with a deterministic simulator, a probabilistic distribution (typically uniform) is assumed to represent the range of variation in controllable inputs.

In Marrel et al. (2012), the total variation in the *mean* of the stochastic simulator is analyzed through a functional ANOVA decomposition and Sobol indices are computed based on the percentage of the total *simulator* variation each component explains. The variation explained by the seed variable ϵ_{seed} can also be computed, representing the total variation explained by the intrinsic variance. Additionally, a sensitivity analysis of the intrinsic variance $\sigma_N^2(x)$ can be conducted separately to gather information on which input variables most impact the heteroscedasticity.

Hart, Alexanderian and Gremaud (2017) assume the simulator can be run at different inputs x with the same seed ϵ_{seed} . Rather than building a joint surrogate for the mean and variance, as described in Section 3.2, they build

a separate surrogate for a number of seeds. For each seed, they obtain a realization of each Sobol index, and by aggregating the realizations, they obtain distributions for the indices.

The extensive literature on model selection may have counterparts that can be effective for stochastic simulators, but a fully satisfactory approach even for deterministic simulators remains somewhat elusive.

7. CONCLUDING REMARKS

There are several key messages to be drawn from this review, each pointing to open or new research questions:

Gaussian process surrogates. GPs are discussed extensively because they provide a flexible way of allowing the data to inform about the shape of the underlying process. Moreover, they can be effective predictors and quantifiers of uncertainty. Diagnosing shortcomings in a GP for stochastic simulators (available in deterministic settings (Bastos and O'Hagan, 2009)) is not yet well established.

As noted in Section 3, neural network (deep learning) methods are in active use and under study, some of which may, in combination with GPs, offer promising research directions (Schultz and Sokolov, 2018).

Additionally, it can be difficult to effectively capture nonnormal variability. Doing so with as few simulations as possible, while also properly quantifying the various uncertainties, is likely to be an important research direction for stochastic simulator analysis. The wider quantile regression literature is likely a good starting point.

Design. Stochastic simulators differ from deterministic ones because they require much larger sample sizes and permit the use of replicates, whose treatment is generally ad hoc. This leads to the questions raised in Section 4, forming a direction of important research. Design size rules of thumb, useful even if imperfect, exist for deterministic simulators (Loeppky, Sacks and Welch, 2009), but are lacking for stochastic simulators.

Calibration. Accounting for model discrepancy in calibration is critical but there is no obvious "one-size-fitsall" method. A broad empirical comparison is needed with guidance about which strategies are effective under which conditions. Assessing the effectiveness of different methods can be challenging (see McKinley et al., 2018, for one comparison between ABC and HM), but sorely needed.

Simulator complexity. For complex stochastic simulators, it may not be feasible to obtain enough runs. In some instances, the simulator can be replaced with a less complex one (e.g., Molina, Bayarri and Berger, 2005) that captures key features and permits adequate numbers of simulations. Another path, coupling stochastic simulators with deterministic simulators has been explored as a way to deal with low simulation budgets (Baker, Challenor and Eames, 2020). Multifidelity modeling, where multiple simulators of varying complexity are coupled together (Kennedy and O'Hagan, 2000, Kennedy, Henderson and Wilson, 2020) is a promising solution where possible.

In a similar vein, certain outputs may be less noisy than others, and the modeling of the less-noisy outputs can improve the modeling of the noisier ones. For example, Wang and Ng (2020) use the expectation of a simulator to improve the estimation of noisier quantiles. This is related to the wider variance reduction literature, which has a long history (Barton, Nakayama and Schruben, 2017). Variance reduction has been applied in a number of examples but its use in ABMs is not apparent, perhaps due to the profusion of stochastic elements in an ABM. Fixing the initial seed in a stochastic simulator has played a role in sensitivity analysis (see Section 6.4), but leveraging information about the nature of the intrinsic randomness for wider purposes is an open problem.

This review strives to raise awareness of existing tools and strategies for treating stochastic simulators and provide a starting point for practitioners interested in utilizing up-to-date statistical approaches. Despite the problems being pervasive and challenging, there is a shortage of statistical research in this field. The problems pose computational and technical questions, as well as theoretical and philosophical ones. Current solutions are often capable, but there is a lack of comprehensive comparison between different solutions, and a lack of testing regarding their generalizability to complex situations (such as very large data sets). The hope is that the review provokes statistical researchers to engage the open questions discussed, and for practitioners to make use of the tools which are available.

APPENDIX—OCEAN TRUTH

Throughout, reference to plots of the "truth" of the ocean model is made. These plots are presented in Figure 8, as well as in the supplementary material, for convenience.

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FIG. 8. Ocean example: The "true" mean and standard deviation for the ocean model, for 500 different sites.

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SUPPLEMENTARY MATERIAL

Supplement A to "Analyzing Stochastic Computer Models: A Review with Opportunities" (DOI: 10.1214/ 21-STS822SUPPA; .zip). Supplementary code for the fish example.

Supplement B to "Analyzing Stochastic Computer Models: A Review with Opportunities" (DOI: 10.1214/ 21-STS822SUPPB; .zip). Supplementary code for the ocean example.

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