Sequential design for computer experiments with a flexible Bayesian additive model

Hugh CHIPMAN*, Pritam RANJAN and Weiwei WANG

Department of Mathematics and Statistics, Acadia University, 12 University Avenue, Wolfville, NS, Canada
B4P 2R6

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Abstract: In computer experiments, a mathematical model implemented on a computer is used to represent complex physical phenomena. These models, known as computer simulators, enable experimental study of a virtual representation of the complex phenomena. Simulators can be thought of as complex functions that take many inputs and provide an output. Often these simulators are themselves expensive to compute, and may be approximated by “surrogate models” such as statistical regression models. In this paper we consider a new kind of surrogate model, a Bayesian ensemble of trees (Chipman, George, & McCulloch, 2010), with the specific goal of learning enough about the simulator so that a particular feature of the simulator can be estimated. We focus on identifying the simulator’s global minimum. Utilizing the Bayesian version of the expected improvement criterion (Jones, Schonlau, & Welch, 1998), we show that this ensemble is particularly effective when the simulator is ill-behaved, exhibiting nonstationarity or abrupt changes in the response. A number of illustrations of the approach are given, including a tidal power application.

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Résument: Lors d’expérimentation par ordinateur, un modèle mathématique implanté dans un ordinateur est utilisé pour représenter un phénomène physique complexe. Ces modèles, connus sous le nom de simulateur par ordinateur, permettent l’étude expérimentale d’une représentation virtuelle de phénomènes complexes. Nous pouvons considérer ces simulateurs comme étant des fonctions complexes qui demandent plusieurs intrants et qui fournissent un extrant. Souvent ces modèles sont eux-mêmes coûteux en terme de temps de calcul et ils peuvent être approximés par des modèles de substitution tels que les modèles de régression. Dans cet article, nous considérons un nouveau type de modèle de substitution, un ensemble bayésien d’arbres (Chipman et al., 2010) dont le but spécifique est d’en apprendre assez sur une caractéristique particulière du simulateur pour l’estimer. Notre attention portera particulièrement sur l’identification du minimum global du simulateur. En utilisant une version bayésienne du critère d’amélioration espérée (Jones et al., 1998), nous montrons que cet ensemble est particulièrement efficace lorsque le simulateur ne se comporte pas bien, par exemple s’il présente de la non-stationnarité ou des changements abrupts. Notre approche est illustrée à l’aide de nombreux exemples, incluant une application sur l’énergie marémotrice. La revue canadienne de statistique 40: 663–678; 2012 © 2012 Société statistique du Canada

1. INTRODUCTION

Many phenomena, such as tidal flow, nuclear reactions, climate behaviour, and universe expansion are sufficiently large and complex that direct scientific experiments are either impossible or impractical, due to time and cost constraints. As a result, mathematical models are often used to build a realistic representation of these phenomena, enabling experimentation. For example, with a model of tidal flow, researchers can study the effect of placing underwater electrical turbines,

* Author to whom correspondence may be addressed.
E-mail: hugh.chipman@acadiau.ca

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using the mathematical model to “observe” how tidal flow would change, and predict how much electrical power might be generated.

These mathematical models, often known as computer simulators, typically take a number of inputs and when they are run, generate an output. Running computer simulators can be time-consuming when output values are desired for a large number of different input values. In such situations, a second level of approximation, in which a “surrogate model” is used to approximate the input–output relationship of the simulator, is often used. These surrogate models are flexible regression models, taking an input vector \( x \) and predicting an output \( y \).

As with most statistical models, a surrogate model is estimated with training data, consisting of \( n \) observations \( (x_1, y_1), \ldots, (x_n, y_n) \). Unlike many real-world phenomena, a computer simulator is often deterministic. That is, every time the simulator is run with input \( x \), the same numeric value of output \( y \) will result. Surrogate models that capture such behaviour, interpolating exactly the training data, are popular tools in such computer experiments.

In this paper we consider computer experiments with the very specific goal of optimization. We wish to find the values of input \( x \) which minimize (or maximize) the output \( y \). This sort of approach is taken in Jones, Schonlau, & Welch (1998), where the training data are collected sequentially, allowing refinement of the surrogate model and increasingly accurate prediction of the location of the global minimizer.

By far the most popular statistical surrogate model is the Gaussian process (GP) model (for an overview, see Santner, Williams, & Notz, 2003 and Rasmussen & Williams, 2006). The GP assumes a multivariate normal distribution for a vector of responses at corresponding input locations. The covariance of the responses is taken to depend on the distances between the input locations. Closely related to spatial models (e.g., kriging), the GP model allows interpolation of a deterministic function, and can predict quite effectively in the input space.

The GP model makes rather strong assumptions, however. The assumption that covariance depends only on the distance between points and not their location leads to a stationary model, in which the smoothness of the response function is the same over the entire input space. Limitations such as this have led to extensions. For example, Gramacy & Lee (2008) develop a treed Gaussian process (TGP) model, in which a tree recursively partitions the input space into rectangular regions, and within each region, a stationary GP model is fit.

In this paper we consider a more radical departure from the GP model. We build upon a Bayesian ensemble of trees model (Chipman, George, & McCulloch, 2010, here onwards referred to as CGM), which combines the outputs from a large number of regression trees via a summation. This gives a flexible model capable of capturing nonstationarity and complex relationships, and if present, additive structure. It is also able to emulate both deterministic and stochastic (noisy) simulators. By utilizing Markov chain Monte Carlo (MCMC) for estimation, full inference for the model is available. This in turn enables computation of merit-based criteria, for instance, an expected improvement (EI) criterion (Jones, Schonlau, & Welch 1998), that can guide the sequential design of experiments for identification of a global minimum (our focus in this article) or other features of interest.

Yu et al. (2010) develop an adaptive sampling algorithm for predictive modelling in a traditional regression setting. Like our approach, Yu et al. (2010) use CGM’s ensemble as the predictive model. Yu et al. (forthcoming) consider a similar approach for computer experiments using a variety of predictive models. In both papers, the goal is accurate estimation of an entire response surface. In contrast, we focus on the discovery of a particular feature of a computer simulator output, such as the global minimum.

The remainder of the paper is organized as follows. In Section 2, we introduce Bayesian additive regression trees (BART), the ensemble that will be used as the surrogate model. After discussing adaptation of prior distributions for BART to computer experiments, we briefly outline
how TGP may be used as another surrogate model. The two models are illustrated with a tidal power generation example. In Section 3, the EI criterion of Jones, Schonlau, & Welch (1998) is reviewed and adapted to the BART model. A number of illustrations are given in Section 4, with test functions playing the role of computer models, and up to four-dimensional inputs considered. Section 5 concludes with a discussion of future directions for research.

2. STATISTICAL SURROGATE MODELS

In this section, we present a brief review of the BART model (see CGM for more details). We also discuss the TGP model of Gramacy & Lee (2008), which will be compared to BART in Section 4.

2.1. The BART Model

Assume the simulator takes a $d$-dimensional input vector $x = (x_1, \ldots, x_d)$ and has real-valued output $y(x)$. The BART model represents the output as a sum of $m$ adaptively chosen functions and an independent normal error,

$$y(x) = \sum_{j=1}^{m} g(x; T_j, M_j) + \epsilon = h(x) + \epsilon, \quad \epsilon \sim N(0, \sigma^2).$$

(1)

The function $g(x; T, M)$ produces an output when provided with input vector $x$ and parameters $T$ and $M$. It is the shorthand notation for a “regression tree” model. The predictions for a particular value of $x$ are generated by following the sequence of decision rules in tree $T$ until arriving at a terminal node $b$, at which point an associated scalar prediction $\mu_b$ is returned. Decision rules in $T$ branch on a single element of the vector $x$, say $x_1$, yielding rules such as $x_1 < 2$. Different rules in each tree may use different variables. For a tree $T$ with $B$ terminal nodes (i.e., partitioning the input space into $B$ rectangular regions), let $M = (\mu_1, \ldots, \mu_B)$ denote the collection of terminal node predictions.

Thus, viewed as a function of $x$, tree model $g$ produces a piecewise-constant output. By combining together an “ensemble” of $m$ such tree models in (1), a flexible modelling framework is created. For instance, if each individual $T_j$ uses partitions on only a single input variable, then the BART model becomes an additive model. BART is however capable of incorporating higher-dimensional interactions, by adaptively choosing the structure and individual rules of the $T_j$’s. Furthermore, many individual trees ($T_j$) may place split points in the same area, allowing the predicted function to change rapidly nearby, effectively capturing nonstationary behaviour such as abrupt changes in the response.

Viewing this as a statistical model, we have parameters $\Theta = (T_1, \ldots, T_m, M_1, \ldots, M_m, \sigma)$. CGM take quite large values of $m$ (say, 50–200), and estimate (1) in a Bayesian framework using MCMC. With a sufficiently large number of terms in the ensemble, the interpretation of individual trees becomes irrelevant, and the most useful interpretation of BART is to view it as a way of placing a prior on $E(Y|X)$, that is a prior on functions. Of course, all regression models, including GPs, place a prior on functions. What is interesting is the sort of functions that receive prior mass.

The BART model places a prior on functions that differs from GPs in several important ways. First, as mentioned above, the BART model is biased towards additive and low-dimensional functions (i.e., $g(x; T_j, M_j)$ in (1) with small trees $T_j$, that are functions of one or a few variables). Second, the BART model does not assume continuity of the response, thus making it appropriate when there are abrupt changes or nonstationarity in the response. In contrast, the explicit specification of a spatial correlation structure in a GP model implies continuity and, for many common correlation functions, stationarity (i.e., a constant amount of “wiggle”) of the response.
The Bayesian framework equips the BART model with a full suite of inferential tools. Most importantly, uncertainty in predicting $h(x)$ due to all parameters is available. As will be demonstrated in Section 3, this is a key ingredient in sequential design.

2.2. Prior Specification of BART for Computer Experiments

CGM demonstrate effective performance of BART in a wide range of simulated and real-data examples. In all examples, there was considerable noise, that is $\sigma \gg 0$. Often in computer experiments, the simulator is deterministic ($\sigma = 0$) or has small errors ($\sigma \approx 0$). This property underlies some important changes to CGM’s default choice of prior distributions. This section briefly outlines those choices, with an emphasis on modifications for surrogate modelling of the deterministic simulator output.

We follow the same general formulation of prior distributions as in CGM, with different parameter choices. CGM simplify prior specification by assuming that, a priori, (i) $T_1, \ldots, T_m$ are i.i.d., (ii) all elements of $M_1, \ldots, M_m$ (i.e., the $\mu_{i,b}$, node $b$ of tree $i$) are i.i.d. given all $T$’s, and $\sigma$ is independent of all $T$’s and $M$’s.

A special prior for a generic output $\mu$ of a tree is used. In a computer experiment, with $\sigma \approx 0$, we believe $h(x) \approx y(x)$, that is, the overall output is quite close to the observed response. Thus we choose a larger prior variance for individual tree output $\mu_{b,i}$ than the CGM default. This allows the outputs more flexibility, giving $h(x) \approx y(x)$ at the training points. This is accomplished as follows: After scaling data values of $Y$ to the interval $(-0.5, 0.5)$, CGM recommend a default prior $\mu \sim N(0, \sigma^2_\mu) = N(0, 1/(4k^2m))$ with $k = 2$. Since the prediction $h(x)$ in (1) is a sum of $m$ distinct $\mu$’s, the default prior means that $h(x) \sim N(0, 1/(4k^2))$. There is thus a prior probability of approximately $95\%$ that $h(x)$ falls within $0 \pm 2\sigma_{\mu} = 0 \pm 1/k$, or $(-0.5, 0.5)$ for $k = 2$. We recommend relaxing this prior to $k = 1$. Of note is the fact that CGM’s default prior specification applies considerable shrinkage to individual output $\mu_{i,b}$. Choosing a smaller value of $k$ (e.g., $k = 1$) increases the prior variance of output $h(x)$, applying less shrinkage (or smoothness) of the response.

The assumption that $\sigma \approx 0$ requires modification to the prior on $\sigma$. Essentially, we place most of the prior mass near 0. This is accomplished with the same inverted-chi-squared prior for $\sigma^2$ as in CGM, using their recommended value of 3 degrees of freedom, and anchoring the 90th percentile of the $\sigma$ prior at $0.20 \times s\bar{d}(Y)$, where $s\bar{d}(Y)$ is the sample standard deviation of the training $y$ values. The only difference from the CGM default is the value used to anchor the 90th percentile (i.e., $0.20 \times s\bar{d}(Y)$). This prior specification allows for some noise in the response values. This strategy facilitates MCMC mixing for BART, and can also be considered as having a similar motivation to Gramacy & Lee (2012) who advocate the inclusion of a residual error when approximating deterministic functions with GPs, for numeric stability and predictive accuracy.

Although not part of the prior specification, several other operating parameters of BART are chosen as follows: The number of trees in the ensemble is chosen as $m = 100$. Individual trees are allowed to split on a fine grid of 1,000 cutpoints along each axis. The MCMC algorithm uses 6,000 iterations, discarding the first 2,000 (burn in) and keeping every 20th thereafter, for a sample of 200 posterior draws. Larger posterior samples might be desirable, but with the quick mixing behaviour of BART observed by CGM, a sample of this size will be sufficient for the sequential design step covered in Section 3.

The remaining elements of the prior specification involve the tree $T$. We use the defaults from CGM, with a prior on tree $T$ that puts a probability mass of 0.05, 0.55, 0.28, 0.09, and 0.03 on trees with 1, 2, 3, 4, and $\geq 5$ terminal nodes. While favouring small trees, this prior does not rule out the possibility of very large trees.

2.3. Treed Gaussian Processes and Gaussian Process Models

Later in the paper we shall make comparisons between BART and another tree-based model, the TGP model of Gramacy & Lee (2008). We briefly review the model and prior specification here.
The TGP model has form similar to (1), but with \( m = 1 \), that is, a single tree. The replacement of the ensemble by a single tree is offset by an enrichment of the terminal node model. Instead of a constant mean in each terminal node, Gramacy and Lee assume that conditional on the tree structure, the response \( y(x) \) follows a GP model with \( \text{var}(y(x)) = \sigma^2_h + \sigma^2 \) and \( \text{cov}(y(x), y(x')) = \sigma^2_h K(x - x') \), where \( K(x - x') \) is the spatial correlation structure, and \( \sigma^2_h \) and \( \sigma^2 \) are the process and noise variance respectively. By allowing a distinct GP model for each terminal node, TGP can accommodate nonstationarity in the response.

Since the TGP model was developed for the emulation of computer simulators, we need less modification of the prior settings. In all our experiments we used default parameters in the R implementation (the \texttt{tgp} package), except for \( \text{BTE} = c(2,000,6,000,20), \text{nug.p} = c(1,10,1,10^5) \). The BTE parameter indicates that the predictive samples are saved every E (20) MCMC rounds starting at round B (2,000), and stopping at T (6,000). That is, the TGP predictions were also based on 200 posterior draws, the same as for BART. The \text{nug.p} parameter specifies a mixture of exponential priors (with expected values \( 10^{-1} \) and \( 10^{-5} \)) on the “nugget” (\( \omega \)), where \( \sigma^2 = \sigma^2_h \cdot \omega \).

GP models are also popular for sequential optimization. Our comparisons also include GPs, using the implementation from the \texttt{tgp} package in R. All settings for the GP model are as above for TGP, except of course without the trees.

2.4. Small Illustration: 1D Tidal Power

To give a taste of these two models, we briefly describe a tidal power modelling application. A full description of the application is given in Wang (2010) and Ranjan et. al. (2011). The basic problem is to predict the amount of kinetic energy (which in turn will be used for producing electricity) that could be extracted from tidal flow through the Minas Passage in Nova Scotia, Canada. This 7-km wide passage has some of the highest tides in the world. According to Karsten et al. (2008), an individual turbine can generate up to 1 MW of power, and approximately 2.5 GW of power can be harnessed from the tidal kinetic energy by placing large collections of turbines in the Minas Passage. Karsten et al. suggest that a “fence” of turbines across the passage is optimal for extracting the maximum possible energy at the minimum cost.

Wang considered the emulation of a simplified tidal flow model (simulator), in which fences of turbines are placed in the Minas Passage. The only parameter to control in this simulator is the location of the fence along the passage. That is, our “\( x \)”, when scaled to the interval \((0, 1)\), represents the fence position along the passage. Variation in the shape of the shoreline leads to a considerable difference in the flow at different locations. A second very significant influence on the flow is the placement of other turbine fences. In the area surrounding a previously placed turbine fence, the flow can be expected to be much lower.

A tidal model can provide output of the power (MW) as a function of the fence location. Thus we have a one-dimensional optimization problem. The true power function (multiplied by \(-1\) so our goal is minimization) is displayed in each of the four panels of Figure 1. Note that in this simplified model the power function can be efficiently evaluated at many \( x \), making the study of emulators more of an academic exercise. For this particular power function, the effect of a pre-existing turbine fence at roughly \( x = 0.83 \) is evident, as the power dips (or negative power spikes) at this location.

The power function is the same in all panels. Each panel displays the same training set, and the corresponding predictions (posterior mean at a given \( x \)) for BART and TGP. The power function displays considerable nonstationarity (smooth, gradual variation for \( 0 \leq x \leq 0.65 \) and rapid fluctuations for \( 0.65 \leq x \leq 1 \)). BART deals more consistently with this nonstationarity, closely tracking the observed values. TGP is more likely to declare that this variation is noise, and the resultant prediction is attenuated, relative to the true function. Although TGP is able to

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Figure 1: Consistency comparison in fitting surrogate model using BART and TGP for the negative power function in the tidal power example. (a) Realization 1; (b) Realization 2; (c) Realization 3; (d) Realization 4. [Colour figure can be viewed in the online issue, which is available at wileyonlinelibrary.com]

partition the space into different regions and fit a GP model in each area, in this case the TGP model is unable to consistently capture nonstationarity.

3. SEQUENTIAL DESIGN WITH BART AND EXPECTED IMPROVEMENT

As mentioned in the introduction, we consider a specific kind of sequential design, in which the goal is to find the optimum of a computer simulator. In the remainder of the paper, we shall assume that the goal is minimization, since it is trivial to switch between minimization and maximization.

We follow the approach of Jones, Schonlau, & Welch (1998). Let $f_{min} = \min\{y_1, \ldots, y_{n_0}\}$ be the current best (i.e., smallest) function value among the $n_0$ points sampled thus far. Jones, Schonlau, & Welch (1998) define the improvement at a point $x$ as $I(x) = \max\{f_{min} - y(x), 0\}$, which is positive if the (unobserved) response $y(x)$ at location $x$ is less than the current best function value, and 0 otherwise. They follow the maximum likelihood approach and take the expectation of this improvement function with respect to $y(x) \sim N(\hat{y}(x), s^2(x))$, where $\hat{y}(x)$ and $s^2(x)$ are the BLUP and the mean square error of response $y$ at $x$. The GP assumption leads to a
closed-form expression,

\[
EI(x) = E[I(x)] = (f_{\min} - \hat{y}) \Phi \left( \frac{f_{\min} - \hat{y}}{s} \right) + s \phi \left( \frac{f_{\min} - \hat{y}}{s} \right),
\]

where \( \Phi \) and \( \phi \) are the standard normal CDF and density. The first term in (2) captures “local search” that seeks to improve the estimate of the minimum near a currently identified minimum. The second term captures “global search” which places points in regions where there is sufficient uncertainty that the minimum response could be nearby.

There are a variety of approaches for evaluating (2) or similar expressions. All involve some form of integration over the predictive distribution for the unobserved response at a new input \( x \). In this paper, we take a fully Bayesian approach, calculating the EI by taking the expectation of \( I(x) \) with respect to the posterior for all parameters, using the MCMC samples. This accounts for both predictive uncertainty and parameter uncertainty. We employ a similar approach for TGP and GP. Although we do not use (2), the ideas of local and global search are still useful concepts to bear in mind when considering how EI chooses runs in a sequential design.

The details of the computation of EI are as follows: For BART, we calculate EI directly from the MCMC output. For every posterior draw \( \Theta^{(i)} = (T_1^{(i)}, \ldots, T_m^{(i)}, m_1^{(i)}, \ldots, m_m^{(i)}, \sigma^{(i)}) \), it is straightforward to get the posterior realization of \( y(x) \) and hence \( I(x) \). Taking a sample average of \( I(x) \) values over the \( N \) MCMC draws yields a MCMC approximation to the EI

\[
EI(x) = \frac{1}{N} \sum_{i=1}^{N} I(x; \Theta^{(i)}) = \frac{1}{N} \sum_{i=1}^{N} \max \{ f_{\min} - y(x; \Theta^{(i)}), 0 \},
\]

where \( y(x; \Theta^{(i)}) = h(x; \Theta^{(i)}) \). This expression for EI incorporates uncertainty in all BART parameters. We use the same MCMC-based strategy for calculating EI with TGP and GP models.

In order to select a new design point, EI must be maximized over all possible \( x \). The current implementation of BART simultaneously runs MCMC and generates posterior draws for \( h(x) \) at a specified set of \( x \) points. Thus, we use space-filling designs (random latin hypercube designs) to generate a large candidate set, evaluate (3) at all points in the candidate set, and select the point with the largest value of EI as the next input at which to obtain simulator output \( y(x) \). To make comparisons with TGP, we employ the same strategy.

As will be illustrated in the higher-dimensional examples in the next section, this “candidate set” strategy for locating the maximizer of EI becomes computationally challenging as the dimensionality of the input space grows.

The sequential design algorithm can thus be summarized as follows:

1. Obtain an initial design \( X_{n_0} \) with \( n_0 \) points, and evaluate the simulator at these points, yielding corresponding simulator outputs \( Y_{n_0} \).
2. Set iteration \( i = 0 \)
3. Select a candidate set \( X^{\text{cand}}_{n_0+i} \) for prediction and evaluation of EI.
4. Fit the model using \( X_{n_0+i}, Y_{n_0+i} \).
5. Calculate EI for each point \( x \in X^{\text{cand}}_{n_0+i} \), identifying the maximizer \( x^* \) of EI.
6. Evaluate the simulator at \( x^* \), augment \( X_{n_0+i} \) and \( Y_{n_0+i} \) with \( x^* \) and \( y(x^*) \), and set \( i = i + 1 \).
7. Unless a stopping condition is met, go to Step 3.

The initial design in Step 1 is a maximin LHD, with \( n_0 - 2 \) runs. The two other runs are \( x = (0, 0, \ldots, 0) \) and \( x = (1, 1, \ldots, 1) \), which are chosen to aid BART in assessing uncertainty near the boundaries of the input space. In Step 3, a new set of candidate points is generated with a
random LHD at every iteration of the algorithm. In Step 4, the “model” is BART, TGP, or GP. In Step 5 EI is calculated using MCMC integration. The “stopping condition” in Step 7 could either be a convergence criterion or tied to a budget. In the remainder of the article, we shall assume a fixed budget of a total of \( n = n_{\text{new}} + n_0 \) evaluations of the simulator.

4. EXPERIMENTAL RESULTS

In this section, we demonstrate the performance of our proposed approach using a few simulated examples and the tidal power application discussed in Section 2.4. First a simple one-dimensional example is used to illustrate the sequential design approach with BART as the emulator. Then, we demonstrate the proposed approach on more complex simulators with one or more inputs. In all the examples presented here, our objective is to find the global minimum.

The average performance of BART in the sequential design framework is compared with TGP, a popular tree-based surrogate for emulating nonstationary computer simulators. We also make comparisons with a standard GP model. The performance of the sequential techniques is also compared with non-sequential “one-shot” space-filling designs. Under the one-shot design setup, the simulator outputs based on an \((n_0 + k)\)-point random maximin LHD was used to estimate the global minimum, where \(0 \leq k \leq n_{\text{new}}\). Although a series of different LHDs of increasing size is not a viable sequential design strategy, it does provide a basis for comparison. All results presented in this section are based on 100 realizations (i.e., 100 different starting designs based on random maximin LHDs, each followed by sequential design generation). For convenience, the EI-based sequential design approach using BART with (3) is referred to as BART-EI, and TGP-EI (GP-EI) refers to the sequential procedure if TGP (GP, respectively) is used instead of BART.

Implementations of BART-EI, TGP-EI, and GP-EI follow the algorithm outlined at the end of Section 3. For all examples presented here, the BART runs follow the prior parameters and operational settings described in Section 2.2, whereas the TGP parameter settings are discussed in 2.3. The GP settings are the same as for TGP, but with the constraint that there must be a single node, giving the same GP over the entire input space. Both TGP and GP also use MCMC integration to evaluate EI.

For all examples, the following details apply. We start by fitting a surrogate to the simulator data on an \(n_0\)-point random maximin Latin hypercube design (LHD). These designs are generated using the R function `maximinLHS` (Carnell, 2009) which takes random starting points and hence the output designs are random. Jones, Schonlau, & Welch (1998) and Loeppky, Sacks, & Welch (2009) suggest using approximately \(n_0 = 10d\) points as a reasonable rule-of-thumb for an initial design. However, the optimal choice of \(n_0\) varies with the complexity of the computer simulator. The remaining \(n_{\text{new}} = n - n_0\) points are sequentially added one at a time by optimizing the EI criterion and refitting the surrogate (BART, TGP, or GP). The size of the candidate sets \(X_{\text{cand}}\) for the evaluation of EI at unsampled input locations varies with the complexity of the simulators.

We use the running best estimate of the global minimum (i.e., the smallest \(y\) value observed thus far) as our performance measure. An alternative considered, but not reported here, is the distance between the input locations of the true and the estimated global minimum. This may not be useful if the simulator has multiple global minima, which is the case in Example 3.

Example 1. Suppose the simulator outputs are generated using the simple one-dimensional test function

\[
y(x) = \frac{\sin(10\pi x)}{2x} + (x - 1)^4, \quad x \in [0.5, 2.5].
\]

(4)

The inputs are scaled to [0, 1] for the implementation. Gramacy & Lee (2012) used this example to demonstrate the benefits of including a nugget in a GP model. Figure 2 illustrates the sequential BART-EI methodology for \(n_0 = 10\) and \(n_{\text{new}} = 15\) runs.
Although BART does not accurately capture all high-frequency oscillations of the function, it appears to effectively guide sequential design for the minimum, utilizing a combination of local and global search. Figure 2a shows that the first follow-up trial is in the vicinity of the current best estimate of the global minimum (i.e., supporting the local search). Figure 2b shows that the next four follow-up points support exploration and the points are away from the running global minimum. Most additional points are near the running global minimum (see panel c), although some points such as 11, 12, 14, and 15 are placed in regions of high uncertainty (i.e., global search). The emulator constructed using BART exhibits “football shaped” uncertainty bounds as typically shown by GP models for deterministic simulators. The variance of the predicted response at the training points is not zero, however, due to the inclusion of a noise term in the BART model.

To compare GP, TGP, and BART, we repeated the experiment 100×. Following the $n_0 = 10d$ rule of thumb, we started with $n_0 = 10$ points and added $n_{\text{new}} = 40$ runs sequentially one
at a time. We used 1,000-point random LHDs as candidate sets \( X_{\text{cand}} \) at which EI was evaluated to choose the next run. Figure 3 presents the performance comparison between BART-EI, TGP-EI, GP-EI and the one-shot design method. The performance is summarized using the mean and median (over 100 realizations) of the running best estimates of the global minimum for each method.

As expected, the sequential design approaches outperform the one-shot design scheme. BART-EI slightly outperforms TGP-EI and is clearly superior to GP-EI. The mean value of the running best estimate is typically larger than the median because the best estimate is bounded below by the global minimum of \( y(x) \). Thus, the horizontal lines just below \(-0.85\) indicate that the global minimum has been found in over half of the 100 replicates. The more gradual approach of the mean lines indicates that in some cases, the global optimum has not yet been found.

**Example 2.** We revisit the tidal power example, where our objective is to maximize the power function (or equivalently, minimize the negative power function). Recall from Figure 1 that the negative power function is very spiky near the global minimum and there is another spike with the function value close to the global minimum. This makes the optimization problem tricky especially if the design points are not densely sampled in both the spikes. Thus, we started the sequential algorithms with a slightly larger initial design, \( n_0 = 15 \), and added \( n_{\text{new}} = 45 \) follow-up points one at a time. The EI criterion was calculated on a 1,000-point random LHD. Figure 4 summarizes the mean and median (over 100 realizations) of the running best estimates of the global minimum.
As in Example 1, the sequential design approaches outperform the naive one-shot approach. However, the performance comparison of BART-EI and TGP-EI shows interesting results. The mean curve for BART-EI is converging at a slower rate than TGP-EI or GP-EI, but in terms of median performance, BART dominates. It turns out that BART-EI can sometimes get stuck in the wrong spike if there are not enough design points in the regions of interest (see Figure 1).

Example 3. Let \( x = (x_1, x_2) \in [0, 1]^2 \), and the simulator outputs be generated from a two-dimensional additive function given by (Rönkkönen et al., 2008)

\[
y(x) = \frac{1}{4} \sum_{i=1}^{2} \left[ \cos(4\pi w_i) + \alpha \cos(8\pi w_i) \right],
\]

where \( \alpha = 0.8 \) and \( w_i = \sum_{j=0}^{n_i} (n_i) P_{i,j}(1-x_i)^{n_i-j} x_i^j \) for \( i = 1, \ldots, d, n_1 = n_2 = 4 \) and \( P_1 = (0, 0.1, 0.2, 0.5, 1), P_2 = (0, 0.5, 0.8, 0.9, 1) \). The contour plot in Figure 5 shows that the test function has 16 global minima with \( y_{\text{min}} \approx -0.478 \) (shown in red diamonds).

Since the function has multiple global minima, one might suspect that the estimation of the global minimum would be relatively easier. But the local versus global search trade-off feature of the EI criterion forces the follow-up trials to jump from the neighbourhood of one global minimum to another. That is, instead of precisely estimating one global minimum, the EI criterion tries to minimize the prediction uncertainty near several global minima. Consequently, the sequential algorithms require a large number of follow-up trials to accurately estimate the global minimum.
Following the $n_0 = 10d$ rule, we started the implementation with $n_0 = 20$ points chosen using random maximin LHD and sequentially added $n_{\text{new}} = 40$ follow-up points using the EI criterion, which was evaluated at 5,000-point random LHD over $[0, 1]^2$. Figure 6 presents the mean and median (over 100 simulations) of the running best estimate of the global minimum.

It is clear from Figure 6 that BART-EI performs significantly better than TGP-EI and GP-EI. The fact that the spikes are aligned to the axes should help both BART and TGP. TGP should benefit more from the smoothness of the response function, while BART may be better suited for identifying the additive structure. The slower convergence of TGP-EI can perhaps be attributed to the inconsistent prediction of TGP.

In this example a larger candidate set $X^{\text{cand}}$ was used to evaluate EI and select the next run. Recall from Section 3 that the EI criterion is evaluated only at these candidate points, and thus their distribution and density play a role in estimating the global minimum. With a two-dimensional input space, we elected to use more candidate points.

**Example 4.** Suppose the computer simulator outputs are generated using the four-dimensional test function given by

$$y(x) = \sum_{i=1}^{4} -\sin(x_i) - 2 \exp \left( -30x_i^2 \right), \quad x_i \in [-2, 2],$$

where the input variables, $x = (x_1, x_2, x_3, x_4)$, are scaled to $[0, 1]^4$. This test function is based on a one-dimensional function from DiMatteo, Genovese, & Kass (2001), and has a unique global minimum with $y_{\text{min}} = -8$, but the global minimum is in a narrow spike. Figure 7 shows the one-dimensional function. The detection of this spike in the four-dimensional $[0, 1]^4$ space would
require at least a few design points in $[0.4, 0.6]^4$, that is in 0.16% of the total volume, otherwise the surrogate models can get misled by the overall shape (i.e., excluding the spike).

A space-filling LHD-based initial design would have to be very large to guarantee a few points in the spiky region $[0.4, 0.6]^4$. On the other hand, if we ignore this spike, the test function or simulator is relatively simple to emulate. We start the sequential procedure with only $n_0 = 30$ points (smaller than the recommended $n_0 = 10d$ rule) and leave the discovery of the spiky region to the follow-up runs. We sequentially added $n_{\text{new}} = 120$ follow-up points. To ensure the evaluation of the EI criterion in the spiky region, we considered dense candidate sets $X_{\text{cand}}$ with 20,000 and 50,000 points. Figure 8 summarizes the running global minimum.

BART-EI very clearly outperforms the other methods, with the median curves indicating that the global optimum is attained or nearly attained with an additional 50 points, in over 50% of the 100 replicates. Evidently, BART is much better suited for exploiting the additive structure in (6). There is a small but non-negligible benefit to evaluating the EI criterion for BART with a larger (50K) candidate set.

5. DISCUSSION

The various examples in Section 4 demonstrate that BART can be an effective engine for sequential design and optimization. In situations where a GP model or perhaps localized GP models are appropriate surrogates, optimization with BART is still competitive. When there are nonstationarities, abrupt changes or additive structure, we see that BART can deliver the optimum with fewer runs of the simulator.
Figure 7: True one-dimensional test function used to build the simulator in (6).

Figure 8: Comparison of the running best estimate of the global minimum using BART-EI and TGP-EI for the four-dimensional simulator (6), with \( n_0 = 30 \) and \( n_{\text{new}} = 120 \). [Colour figure can be viewed in the online issue, which is available at wileyonlinelibrary.com]
As in any sequential design procedure, the size and configuration of points in the initial design can impact the performance of the sequential design. Jones, Schonlau, & Welch (1998) and Loeppky, Sacks, & Welch (2009) suggest using approximately \( n_0 = 10d \) points as a reasonable rule-of-thumb for an initial design. However, the optimal choice of \( n_0 \) depends on the complexity of the simulator. Certainly if a very small number of runs is used in the initial design, the algorithm based on BART (or TGP) may take longer to find the optimum. On the other hand, one may not want use a very large initial design and put several points in the unimportant regions of the input space.

The reader may notice that the BART model is not continuous, but rather piecewise constant. However, by utilizing many trees in the sum (\( m = 100 \) in our examples) and also employing posterior averaging over these trees, the BART model builds a surrogate that can have many small steps, providing an effective approximation to a continuous function. In our experience, the BART model seems effective at deciding where to probe the simulator (i.e., carry out additional runs), even in situations where it may not provide as “nice” a fit as a GP model. Of course, if a smooth prediction is desired (and possibly a better extrapolation to the minimum, if there is smoothness nearby) one may train a GP or TGP model on the set of all simulator runs, once the sequential design procedure has terminated. If, on the other hand, there is a possibility of nonstationarity, abrupt changes or even discontinuities in the response function, then BART seems more likely to be an effective engine for sequential design and optimization.

As observed in Section 3, to evaluate EI (3), we must specify (or design) a candidate set of points at which to generate predictions from the BART model. Since this set can be large (e.g., 20,000–50,000 in the four-dimensional example), we used a random LHD. A maximin LHD (or other space-filling design) with thousands of points could be computationally challenging to identify.

Also related to the evaluation of EI, for the sake of comparison we used a “batch predict” approach to both BART and TGP. The current implementation of BART is not equipped with a “predict” function than allows predictions for new inputs. Instead the predictions are generated during the MCMC estimation of the model. To make comparisons with TGP, we thus used the same “batch predict” strategy. We are currently exploring alternative ways to evaluate EI and find the most promising points to add during the sequential design.

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BIBLIOGRAPHY

Carnell, R. (2009). lhs: Latin Hypercube Samples, R package version 0.5.
Chipman, H. A., George, E. I., & McCulloch, R. E. (2010). BART: Bayesian additive regression trees. *Annals of Applied Statistics*, 4, 266–298.
DiMatteo, I. Genovese, C. R., & Kass, R. E. (2001). Bayesian curve fitting with free-knot splines. *Biometrika*, 88, 1055–1071.
Gramacy, R. B. & Lee, H. K. H. (2008). Bayesian treed Gaussian process models with an application to computer modeling. *Journal of the American Statistical Association*, 103, 1119–1130.
Gramacy, R. B. & Lee, H. K. H. (2012). Cases for the Nugget in modelling computer experiments. *Statistics and Computing*, 22, 713–722.

DOI: 10.1002/cjs
Jones, D., Schonlau, M., & Welch, W. (1998). Efficient global optimization of expensive black-box functions. *Journal of Global Optimization*, 13, 455–492.

Karsten, R., McMillan, J., Lickley, M., & Haynes, R. (2008). Assessment of tidal current energy for the Minas Passage, Bay of Fundy. *Proceedings of the Institution of Mechanical Engineers, Part A: Journal of Power and Energy*, 222, 493–507.

Loeppky, J. L., Sacks, J., & Welch, W. J. (2009). Choosing the sample size of a computer experiment: A practical guide. *Technometrics*, 51(4), 366–376.

Ranjan, P., Haynes, R., & Karsten, R. (2011). A computationally stable approach to Gaussian process interpolation of deterministic computer simulation data. *Technometrics*, 53(4), 366–378.

Rasmussen, C. E. & Williams, C. K. I. (2006). *Gaussian Processes for Machine Learning*, The MIT Press, Cambridge, MA, USA.

Rönkkönen, J., Li, X., Kyrki, V., & Lampinen, J. A. (2008). Generator for multimodal test functions with multiple global optima. In *Proceedings of 7th International Conference on Simulated Evolution and Learning (SEAL’08)*, Melbourne, Australia, 239–248.

Santner, T. J., Williams, B. J., & Notz, W. I. (2003). *The Design and Analysis of Computer Experiments*, Springer-Verlag, New York.

Wang, W. (2010). Modelling Energy Output to Optimize Tidal Turbine Placement, B.Sc. Thesis, Acadia University, Canada.

Yu, Q., Li, B., Fang, Z., & Peng, L. (2010). An adaptive sampling scheme guided by BART-with an application to predict processor performance. *The Canadian Journal of Statistics*, 38(1), 136–152.

Yu, Q., Li, B., Fang, Z., & Peng, L. Model guided adaptive design and analysis in computer experiment, *Statistical Analysis and Data Mining, forthcoming*, http://onlinelibrary.wiley.com/doi/10.1002/sam.11156/abstract;jsessionid=6B182CE0F0B444148639BB087EDE5F36.d03t01?deniedAccessCustomisedMessage=&userIsAuthenticated=false.

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