Information-theoretic Inducing Point Placement for High-throughput Bayesian Optimisation

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Abstract

Sparse Gaussian Processes are a key component of high-throughput Bayesian optimisation (BO) loops — an increasingly common setting where evaluation budgets are large and highly parallelised. By using representative subsets of the available data to build approximate posteriors, sparse models dramatically reduce the computational costs of surrogate modelling by relying on a small set of pseudo-observations, the so-called inducing points, in lieu of the full data set. However, current approaches to design inducing points are not appropriate within BO loops as they seek to reduce global uncertainty in the objective function. Thus, the high-fidelity modelling of promising and data-dense regions required for precise optimisation is sacrificed and computational resources are instead wasted on modelling areas of the space already known to be sub-optimal. Inspired by entropy-based BO methods, we propose a novel inducing point design that uses a principled information-theoretic criterion to select inducing points. By choosing inducing points to maximally reduce both global uncertainty and uncertainty in the maximum value of the objective function, we build surrogate models able to support high-precision high-throughput BO.

1. Introduction

Countless design tasks in science, industry and machine learning can be formulated as high-throughput optimisation problems, as characterised by access to substantial evaluation budgets and an ability to make large batches of evaluations in parallel. Prominent examples include high-throughput screening within drug discovery, DNA sequencing, and experimental design pipelines, where automation allows researchers to efficiently oversee thousands of scientific experiments, field tests and simulations. In addition, these pipelines employ an ever increasing degree of parallelisation, through cheap access to high-fidelity sensor arrays and cloud compute resources. However, such design tasks have sufficiently large and multimodal search spaces that, even under large optimisation budgets, only a small proportion of candidate solutions can ever be evaluated, and often only approximately or under significant levels of observation noise. Consequently, most existing optimisation routines are unsuitable, as brute-force methods require too many evaluations and assume exact evaluations.

Bayesian Optimisation (BO, see Shahriari et al., 2016, for a review) has surfaced as the de facto approach for solving black-box optimisation tasks under restricted evaluation...
budgets, with numerous successful applications across the empirical sciences and industry. However, vanilla BO relies on a Gaussian process (GP, Rasmussen, 2004), which incurs a significant computational overhead for each individual optimisation step, and this cost becomes increasingly unwieldy as data volumes increase, making it unsuitable for the high-throughput tasks motivated above. Sparse GPs (Titsias, 2009) dramatically reduce the computational cost of GPs, allowing BO to scale efficiently with data volumes (Vakili et al., 2021). In a nutshell, sparse GPs replace the full set of observations by a smaller representative set of pseudo-observations referred to as inducing points. The choice of the inducing point locations has a critical influence on the behaviour of the model, in particular as it encodes its local expressivity. However, existing approaches for inducing point placement focus purely on regression tasks, i.e. the global accuracy of models and so sacrifice the high-fidelity modelling of promising regions that is required to guide precise optimisation, leading potentially to poor optimisation performance (Figure 1).

We propose the first inducing point selection strategy designed specifically for BO. Our subset selection method uses a principled information-theoretic criterion to explicitly trade off the modelling resources spent explaining global variations in the objective functions with those spent providing accurate modelling around potential optima (as demonstrated in Figure 1). Using synthetic benchmark problems, we demonstrate that our new inducing point selection method supports high-precision BO while existing selection strategies do not.

2. Preliminaries

2.1 Bayesian Optimisation with Sparse Gaussian Processes

GP models are a popular choice for Bayesian modelling, as they combine flexibility with reliable uncertainty estimates. Moreover, for regression with a Gaussian likelihood, the exact posterior can be computed in closed form. However, computing this posterior has computational complexity $O(N^3)$ and requires $O(N^2)$ memory.

To mitigate this, sparse approaches (Titsias, 2009; Hensman et al., 2013) have been developed which have complexity $O(M^2N)$ and $O(MN)$ memory cost, where we use $M \ll N$.
inducing points, which are typically selected to provide an effective representation of the full dataset. In this work, we focus on the stochastic variational Gaussian process (SVGP) approach introduced in Hensman et al. (2013), which allows for training with minibatches.

Finally, Vakili et al. (2021) have recently shown that the decoupled sampling approach based on Matheron’s rule introduced in Wilson et al. (2020) can be used with SVGP models for BO with large batch Thompson sampling, resulting in drastic efficiency gains over traditional full GP-based BO without significant impact on regret performance.

2.2 Inducing Point Selection Through the Lens of Experimental Design

Common approaches for selecting inducing points include taking a random subset of the data, computing centroids obtained by running a k-means algorithm on the data, or directly optimising their locations by maximising the ELBO. Recent theoretical results on sparse GP methods (Burt et al., 2019; Vakili et al., 2021) assume that the set of \( M \) inducing points \( Z \) are sampled from the available data according to an \( M \)-Determinantal Point Process (\( M \)-DPP, Kulesza and Taskar, 2012). For regression tasks, this is a meaningful design criterion, as the repulsion properties of DPPs ensure that inducing points are spread across the whole search space with high probability. More precisely, an \( M \)-DPP inducing point selection criterion chooses the set \( Z \) for our \( M \) inducing points with probability proportional to the determinant of the \( M \times M \) Gram matrix \( K_Z = [k(z_i, z_j)]_{(z_i, z_j) \in Z \times Z} \), i.e.,

\[
P(Z = Z) \propto |K_Z|.
\]  

(1)

To help motivate our BO-specific inducing point selection routine, it is worth stressing a link between the DPP formulation above and experimental design. In particular, the arguments of the celebrated paper of Srinivas et al. (2009) can be modified to show that the MAP \( M \)-DPP objective of Eq. 1 is equivalent to choosing the \( M \) inducing points \( Z \) providing the largest gain in information about the unknown function \( f \),

\[
Z = \arg\max_{Z \subseteq D_n:|Z|=M} \text{MI}(y_Z, f),
\]  

(2)

where the mutual information \( \text{MI}(y_Z, f) = H(y_Z) - E[H(y_Z|f)] \) quantifies the reduction in the differential entropy \( H \) of \( f \) provided by revealing the evaluations \( y_Z \).

Maximising Eq. 1 corresponds to the maximum a posteriori (MAP) estimation of a DPP, which is known to be an NP-hard problem (Ko et al., 1995), making the use of Eq. 1 impractical. However, it is possible to obtain a high-performing approximation of the MAP estimate using a greedy algorithm which we refer to as conditional variance reduction (CVR). In practice, this algorithm can be seen to greedily build a set of inducing points by maximising the posterior predictive variance of a noise-free and zero-mean Gaussian process model \( f \sim \mathcal{GP}(0, k) \) conditioned on previously selected observations. Through incremental updates of Cholesky decompositions, this algorithm returns \( M \) points from a set of \( N \) candidates with only \( O(M^2N) \) complexity, i.e. the same cost as a single fit of the resulting sparse GP model (see Hennig and Garnett, 2016; Chen et al., 2018; Burt et al., 2019, for detailed description and discussion).
3. Inducing Point Selection for Bayesian Optimisation Surrogate Models

Our primary considerations to build a BO-specific routine are a) to allow the focusing of modelling resources into promising areas of the space whilst b) maintaining a sufficiently accurate global model. Accurate modelling in promising areas are necessary to allow the precise identification of the optimum, while a level of global accuracy is necessary to prevent the re-investigation of areas already identified as sub-optimal.

3.1 Reducing Uncertainty in $f^*$

We propose choosing a set of inducing points that provide an optimal trade-off between providing information about the whole objective function and providing information about the function’s maximum value. This idea is motivated by the empirical success of max-value entropy search acquisition functions (Wang and Jegelka, 2017; Takeno et al., 2019; Moss et al., 2020), where query points in BO loops are chosen to reduce our current uncertainty in the function’s maximal value $f^* = \max f(x)$.

This entropy-based criterion for inducing point placement, henceforth referred to as ENT-DPP (for reasons that will become obvious below), seeks inducing points $Z$ that maximise the following trade-off:

$$C(Z) = \alpha \times IG(y_Z; f^*) + (1 - \alpha) \times IG(y_Z; f),$$

(3)

where $IG(y_Z, f^*) = H(f^*) - H(f^* | y_Z)$ quantifies the reduction in uncertainty provided by these inducing points about the objective function’s maximal value $f^*$, and $IG(y_Z; f)$ is an in Eq. 2. The trade-off parameter $\alpha \in [0, 1]$ controls the balance of resources spent modelling global variations and modelling variations around potential maxima. Note that setting $\alpha = 1$ returns the criterion of Eq. 2. In our experiments we consider an equal trade-off (ie. $\alpha = 0.5$); however, future work will investigate the practical and theoretical effects of other choices.

As demonstrated in Figure 2, we now have an intuitive inducing point selection strategy well-suited to the demands of BO; however, our criterion is only practically useful if its inducing points can be identified with at most $O(M^2 N)$ cost.

3.2 Efficient (Approximate) Maximisation of the ENT-DPP Criterion

The lack of closed-form expression for the distribution of a Gaussian process’s maximum value $f^*$ renders the calculation of the ENT-DPP criterion (3) challenging. Fortunately, there exists a rich literature of methods for approximately optimising similar information-theoretic quantities (Hennig and Schuler, 2012; Hernández-Lobato et al., 2014; Moss et al., 2021).

We follow the ideas of Moss et al. (2021) and use common information-theoretic inequalities to replace our desired criterion (which we will be attempting to maximise) with a simpler lower bound. Specifically, we use the well-known fact that differential entropy reduces under conditioning, i.e. that $H(f^* | y_Z) \leq H(f^* | y_{z_i}) \forall i \in \{1, ..., M\}$ and write $H(f^* | y_Z) \leq \frac{1}{M} \sum_{i=1}^{M} H(f^* | y_{z_i})$. After simple mathematical manipulation, the resulting lower bound for the ENT-DPP criterion can be expressed in the following form:

$$C(Z) \geq \frac{1 - \alpha}{2} \log |L_Z|,$$

(4)
Figure 2: The 50 inducing points (red) chosen from 250 candidate evaluations (green) using three different inducing point selection strategies when seeking to minimise the 2d Logarithmic Goldstein-price function (background colour-map). We see that existing approaches that (a) use the centroids from a k-means clustering of the available data or (b) use the CVR strategy provide balanced coverage of the whole search space, whereas our ENT-DPP strategy (c) is able to focus resources modelling resources into promising (blue) areas.

for $L_Z = q_Z^T K_Z q_Z$, where $K_Z$ is the kernel Gram matrix and where $q_Z$ is the diagonal matrix containing the elements $q_z$ defined such that $(M(1 - \alpha)/\alpha) \log q_z = IG(y_z; f^*)$. Note that we cannot calculate $IG(y_z; f^*)$ exactly, so we follow Ru et al. (2018) and use a moment-matching approximation. By replacing $f^*$ with a Gaussian variable of mean $\mu$ and variance $\sigma^2$ (as extracted from a set of sampled $f^*$), we have $IG(y_z; f^*) \approx \frac{\gamma \phi(\gamma)}{2 \Phi(\gamma)} - \log \Phi(\gamma) \text{ for } \gamma = \frac{y_z - \mu}{\sigma}$ (see Wang and Jegelka (2017) for a derivation and strategies for sampling $f^*$).

Just like the criterion described in section 2.2, maximising the derived lower bound (4) corresponds to the MAP estimate of a DPP, however, with $K_Z$ replaced by $L_Z$. Hence, we can employ exactly the same greedy algorithm to efficiently build our set of inducing points.

4. Experimental Results

We now provide an empirical evaluation across high-throughput versions of some popular synthetic optimisation benchmarks using the open-source BO library trieste (Berkeley et al., 2022).

For clarity, all our experiments follow the same setup. We consider an SVGP model with either $M = 250$ or $500$ inducing points using either 1) our proposed ENT-DPP method, 2) the CVR of Burt et al. (2019) (see Section 2.2), 3) choosing the centroids of a K-means clustering of the data, and 4) choosing inducing points spread uniformly across the search space. When the total number of queried points is less than the desired number of inducing points (e.g. for the first 4 optimisation steps when $M = 500$), we use just the $N$ available training points as our inducing point locations. SVGP models are fit using an ADAM optimiser with learning rate $0.1$, using an early stopping criteria with a patience of 50 and a learning rate halving schedule with a patience of 10.

We allocate a total evaluation budget of $N = 5,000$ evaluations split across 50 BO steps in batches of 100 points, starting from an initial batch of random evaluations. Subsequent
batches are collected via the Decoupled Thompson sampling scheme presented in Vakili et al. (2021). We use 100 random Fourier features to build the samples and maximise each sample using an L-BFGS optimiser starting from the best of a random sample of 1,000 points. For additional context, BO using an exact GP model is included as a baseline, however, we can report only the first 10 optimisation steps after which it became prohibitively expensive (i.e. for \( N > 1,000 \)).

Figure 3 demonstrates experimental optimisation performance across the 4d Shekel, 5d Michalewicz and 4d Ackley function, where we have contaminated the evaluations of each with Gaussian noise of variance 0.1. Unsurprisingly, greater performance is achieved when using larger number of inducing points for all the considered methods, with ENT-DPP providing the lowest regret across all three tasks. Note that, ENT-DPP outperforms all other approaches on the Shekel function even when using just \( M = 250 \) inducing points. Interestingly, the SVGP-based methods are less prone to converging to one of the many local minima of the Ackley function than exact GPs. Future work will investigate exactly why this is the case.

5. Conclusions and Future Work

We have proposed ENT-DPP — the first BO-specific method for selecting the locations for the inducing points of sparse GPs. Although ENT-DPP’s approximate maximisation relies on potentially coarse approximations, our experimentation found this new inducing point allocation strategy to enable effective high-throughput BO. In future work we will apply ENT-DPP to real-world problems where sparse GPs are already being used, e.g. quantile optimisation (Torossian et al., 2020) or molecular search (Vakili et al., 2021). We will also investigate the applicability of ENT-DPP to other inducing point-based methods like Deep GPs (Damianou and Lawrence, 2013).
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