CobBO: Coordinate Backoff Bayesian Optimization with Two-Stage Kernels

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Abstract

Bayesian optimization is a popular method for optimizing expensive black-box functions. Yet it oftentimes struggles in high dimensions where the computation could be prohibitively expensive and a sufficient estimation of the global landscape requires more observations. We introduce Coordinate backoff Bayesian optimization (CobBO) with two-stage kernels to alleviate this problem. In each iteration, a promising subset of coordinates is selected in the first stage, as past observed points in the full space are projected to the selected subspace adopting a simple kernel that sacrifices the approximation accuracy for computational efficiency. Then in the second stage of the same iteration a more sophisticated kernel is applied for estimating the landscape in the selected low dimensional subspace where the computational cost becomes affordable. Effectively, this second stage kernel refines the approximation of the global landscape estimated by the first stage kernel through a sequence of observations in the local subspace. This refinement lasts until a stopping rule is met determining when to back off from a certain subspace and switch to another coordinate subset. This decoupling significantly reduces the computational burden in high dimensions, while the two-stage kernels of the Gaussian process regressions fully leverage the observations in the whole space rather than only relying on observations in each coordinate subspace. Extensive evaluations show that CobBO finds solutions comparable to or better than other state-of-the-art methods for dimensions ranging from tens to hundreds, while reducing the trial complexity and computational costs.

1 Introduction

Bayesian optimization (BO) has emerged as an effective zero-order paradigm for optimizing expensive black-box functions. The entire sequence of iterations rely only on the function values of the already queried points without information on their derivatives. Though highly competitive in low dimensions (e.g., the dimension $D \leq 20$ [15]), Bayesian optimization based on Gaussian Process (GP) regression has obstacles that impede its effectiveness, especially in high dimensions.

**Approximation accuracy**: GP regression assumes a class of random functions in a probability space as surrogates that iteratively yield posterior distributions by conditioning on the queried points. When suggesting new query points, for complex functions with numerous local optima and saddle points due to local fluctuations, always exactly using the values on the queried points as the conditional events may mismatch the function’s local landscape by overemphasizing the approximation accuracy of the global landscape.

**Curse of dimensionality**: As a sample efficient method, Bayesian optimization often suffers from high dimensions. Fitting the GP model (estimating the parameters, e.g., length_scales [14]), computing the Gaussian process posterior and optimizing the acquisition function in high dimensions all
To alleviate these issues, we design coordinate backoff Bayesian optimization (CobBO) with two-stage kernels, by challenging a seemingly natural intuition stating that it is always better for Bayesian optimization to have a more accurate approximation of the objective function at all times. We demonstrate that this is not necessarily true, by showing that smoothing out local fluctuations and using the estimated function values instead of the true observations to serve as the conditional events in selected subspaces can not only significantly reduce the computation time due to the curse of dimensionality but also help in capturing the large-scale properties of the objective function $f(x)$.

Specifically, CobBO introduces the two-stage kernels with a stopping rule. The first stage of each iteration adopts a simple kernel that sacrifices the approximation accuracy of $f(x)$ for computational efficiency. For example, by using a universal radial basis function (RBF) approximation without learnable parameters [50], CobBO can eliminate the model fitting time in the full space. It captures a smooth approximation $\hat{f}(x)$ of the global landscape by interpolating the values of queried points projected to selected promising subspaces. These projected points serve as the conditional events for GP regression. In a selected coordinate subspace, the second stage of the same iteration applies a sophisticated kernel that can tolerate high computational cost in low dimensions. For example, CobBO uses the Automatic Relevance Determination (ARD) Matérn 5/2 kernel [40]. It refines the approximation of the local landscape less accurately using the RBF kernel, and then explores selected subspaces $\Omega_t$ more accurately using the Matern kernel. This eventually better exploits the promising subspaces.

![Figure 1: Minimize the fluctuated Rastrigin function on $[-5, 10]^D$ with 20 initial samples. [Left] Computation times for training the GP regression model and maximizing the acquisition function at each iteration. CobBO significantly reduces the execution time compared with a vanilla BO, e.g. $\times 13$ faster in this case. [Right] The average error between the GP predictions before making queries and the true function values at the queried points (solid curves, the higher the better) and the best observed function value (dashed curves, the lower the better) at iteration $t$. CobBO captures the global landscape less accurately using the RBF kernel, and then explores selected subspaces $\Omega_t$ more accurately using the Matern kernel. This eventually better exploits the promising subspaces.](image)
This method can be viewed as a variant of block coordinate ascent tailored to Bayesian optimization by applying backoff stopping rules for switching coordinate blocks. While similar work exists [43, 48], CobBO differs by introducing the two-stage kernels and addressing the following three issues:

1. Selecting a block of coordinates for ascending requires determining the block size as well as the coordinates therein. CobBO selects the coordinate subsets by a multiplicative weights update method [2] to the preference probability associated with each coordinate. Thus, it samples more promising subspaces with higher probabilities.

2. A coordinate subspace requires a sufficient number of query points acting as the conditional events for the GP regression. CobBO leverages all observations in the whole space by interpolating the values of queried points projected to selected promising subspaces, rather than simply starting from scratch in each subspace.

3. Querying a certain subspace, under some trial budget, comes at the expense of exploring other coordinate blocks. Yet prematurely shifting to different subspaces does not fully exploit the full potential of a given subspace. Hence determining the number of consecutive function queries within a subspace makes a trade-off between exploration and exploitation. CobBO uses a stopping rule in each subspace to switch the selected coordinates. When consecutively querying data points in the same subspace, CobBO does not need to conduct the first-stage function approximation in the full space, which is far more efficient.

Through comprehensive evaluations, CobBO demonstrates appealing performance for dimensions ranging from tens to hundreds. It obtains comparable or better solutions with fewer queries, in comparison with the state-of-the-art methods, for most of the problems tested in Section 4.2.

2 Related work

Certain assumptions are often imposed on the latent structure in high dimensions. Typical assumptions include low dimensional structures and additive structures. Their advantages manifest on problems with a low dimension or a low effective dimension. However, these assumptions do not necessarily hold for non-separable functions with no redundant dimensions.

Low dimensional structure: The black-box function \( f \) is assumed to have a low effective dimension [30, 58], e.g., \( f(x) = g(\Phi x) \) with some function \( g(\cdot) \) and a matrix \( \Phi \) of \( d \times D, d << D \). A number of different methods have been developed, including random embedding [66, 11, 63, 36, 44, 70, 5, 32], low-rank matrix recovery [11, 58], and learning subspaces by derivative information [11, 13]. In contrast to existing work on subspace selections, e.g., Hashing-enhanced Subspace BO (HeSBO) [44], Mahalanobis kernel for linear embeddings [33], DROPOUT [35] and LineBO [29] (which receives a special treatment in Appendix F), CobBO efficiently leverages all the observations in the whole space using the two-stage kernels and the stopping rule in each subspace for consecutive observations, rather than only relying on limited observations in each coordinate subspace. It exploits subspace structure from a perspective of block coordinate ascent, independent of the dimensions, different from some algorithms that are more suitable for low dimensions, e.g., BADS [1].

Additive structure: A decomposition assumption is often made by \( f(x) = \sum_{i=1}^{k} f^{(i)}(x_i) \), with \( x_i \) defined over low-dimensional components. In this case, the effective dimensionality of the model is the largest dimension among all additive groups [45], which is usually small. The Gaussian process is structured as an additive model [17, 28], e.g., projected-additive functions [36], ensemble Bayesian optimization (EBO) [61], latent additive structural kernel learning (HDBBO) [65] and group additive models [28, 36]. However, learning the unknown structure incurs a considerable computational cost [44], and is not applicable for non-separable functions, for which CobBO can still be applied.

Trust regions and space partitions: Trust region BO has been proven effective for high-dimensional problems. A typical pattern is to alternate between global and local search regions. In the local trust regions, many efficient methods have been applied, e.g., local Gaussian models (TurBO [14]), adaptive search on a mesh grid (BADS [1]) or quasi-Newton local optimization (BLOSSOM [41]). TurBO [14] uses Thompson sampling to allocate samples across multiple regions. A related method is to use space partitions, e.g., LA-MCTS[60] on a Monte Carlo tree search algorithm to learn efficient partitions. CobBO differs by selecting low dimensional subspaces. It can also incorporate trust regions in the first-stage global approximation, as shown in the Appendix.
3 Algorithm

Without loss of generality, suppose that the goal is to solve a maximization problem $x^* = \arg\max_{x \in \Omega} f(x)$ for a black-box function $f : \Omega \rightarrow \mathbb{R}$. The domain is normalized $\Omega = [0, 1]^D$ with the coordinates indexed by $I = \{1, 2, \cdots, D\}$.

For a sequence of points $H_t = \{(x_i, y_i = f(x_i))\}_{i=1}^t$ with $t$ indexing the most recent iteration, we observe $\mathcal{H}_t = \{(x_i, y_i = f(x_i))\}_{i=1}^t$. A random subset $C_t \subseteq I$ of the coordinates is selected, forming a subspace $\Omega_t \subseteq \Omega$ at iteration $t$. As a variant of coordinate ascent, the subspace $\Omega_t$ contains a pivot point $V_t$, which presumably is the maximum point $x_t^M = \arg\max_{x \in \mathcal{X}} f(x)$ with $M_t = f(x_t^M)$.

CobBO may set $V_t$ different from $x_t^M$ to escape local optima. Then, BO is conducted within $\Omega_t$ while fixing all the other coordinates $C_t = I \setminus C_t$, i.e., the complement of $C_t$.

**Algorithm 1: CobBO($f$, $\tau$, $T$)**

1. $\mathcal{H}_t \leftarrow$ sample $\tau$ initial points and evaluate their values
2. $V_t, M_t \leftarrow$ Find the tuple with the maximal objective value in $\mathcal{H}_t$
3. $q_t \leftarrow 0$ Initialize the number of consecutive failed queries
4. $\pi_t \leftarrow$ Initialize a uniform preference distribution on the coordinates
5. **for** $t \leftarrow \tau$ to $T$ **do**
6.   **if** switch $\Omega_{t-1}$ by the backoff stopping rule (Section 3.2) **then**
7.     $C_t \leftarrow$ Sample a promising coordinate block according to $\pi_t$ (Section 3.1)
8.     $\Omega_t \leftarrow$ Take the subspace of $\Omega_t$ over the coordinate block $C_t$, such that $V_t \in \Omega_t$
9.   **else**
10.    $\mathcal{X}_t \leftarrow P_{\Omega_t}(\mathcal{X}_t)$ [Project $\mathcal{X}_t$ onto $\Omega_t$ to obtain a set of virtual points (Eq. 1)]
11.   $\mathcal{H}_t \leftarrow R(\mathcal{X}_t, \mathcal{H}_t)$ [Smooth function values on $\mathcal{X}_t$ by interpolation using $\mathcal{H}_t$]
12.   $p[f_{\Omega_t}(x) | \mathcal{H}_t] \leftarrow$ Compute the posterior distribution of the Gaussian process in $\Omega_t$
13.   **conditional on** $\mathcal{H}_t$
14.    $x_{t+1} \leftarrow \arg\max_{x \in \Omega_t} Q_{f \sim p(f | \mathcal{H}_t)}(x | \mathcal{H}_t)$ [Suggest the next query in $\Omega_t$ (Section 3)]
15.    $y_{t+1} \leftarrow$ Evaluate the black-box function $y_{t+1} = f(x_{t+1})$
16.   **if** $y_{t+1} > M_t$ **then**
17.     $V_{t+1} \leftarrow x_{t+1}, M_{t+1} \leftarrow y_{t+1}, q_{t+1} \leftarrow 0$
18.   **else**
19.     $V_{t+1} \leftarrow V_t, M_{t+1} \leftarrow M_t, q_{t+1} \leftarrow q_t + 1$
20.    $\pi_{t+1} \leftarrow$ Update $\pi_t$ by a multiplicative weights update method (Eq. 2)
21.   $\mathcal{H}_{t+1} \leftarrow \mathcal{H}_t \bigcup \{(x_{t+1}, y_{t+1})\}, \mathcal{X}_{t+1} \leftarrow \mathcal{X}_t \bigcup \{x_{t+1}\}$
22. **end**

For BO in $\Omega_t$, we use Gaussian processes as the random surrogates $\hat{f} = \hat{f}_{\Omega_t}(x)$ to describe the Bayesian statistics of $f(x)$ for $x \in \Omega_t$. At each iteration, the next query point is generated by solving

$$x_{t+1} = \arg\max_{x \in \Omega_t, V_t \in \Omega_t} Q_{f \sim p(f | \mathcal{H}_t)}(x | \mathcal{H}_t),$$

where the acquisition function $Q(x | \mathcal{H}_t)$ incorporates the posterior distribution of the Gaussian processes $p(\hat{f} | \mathcal{H}_t)$. Typical acquisition functions include the expected improvement (EI) [42, 27], the upper confidence bound (UCB) [3, 54, 55], the entropy search [24, 25, 64], and the knowledge gradient [16, 53, 69].

Instead of directly computing the posterior distribution $p(\hat{f} | \mathcal{H}_t)$, we replace the conditional events $\mathcal{H}_t$ by

$$\hat{\mathcal{H}}_t := R(P_{\Omega_t}(\mathcal{X}_t), \mathcal{H}_t) = \{(\hat{x}_i, \hat{y}_i)\}_{i=1}^t$$

with an interpolation function $R(\cdot, \cdot)$ and a projection function $P_{\Omega_t}(\cdot)$,

$$P_{\Omega_t}(x)^{(j)} = \begin{cases} x^{(j)} & \text{if } j \in C_t \\ V_t^{(j)} & \text{if } j \notin C_t \end{cases}$$ (1)
at coordinate $j$. It simply keeps the values of $x$ whose corresponding coordinates are in $C_t$ and replaces the rest by the corresponding values of $V_t$, as illustrated in Fig. 2.

Applying $P_{\Omega_t}(\cdot)$ on $\mathcal{X}_t$ and discarding duplicates generate a new set of distinct virtual points $\hat{\mathcal{X}}_t = \{\hat{x}_1, \hat{x}_2, \hat{x}_3, \cdots, \hat{x}_t\}$, $\hat{x}_i \in \Omega_t \forall 1 \leq i \leq t$. The function values at $\hat{x}_i \in \hat{\mathcal{X}}_t$ are interpolated as $\hat{y}_i = R(\hat{x}_i, \mathcal{H}_t)$ using the standard radial basis function [6, 7] and the observed points in $\mathcal{H}_t$. It not only significantly reduces the GP regression time due to the efficiency of RBF [6] and the acquisition function optimization in low dimensions [11], but also eventually improves the model accuracy using the more sophisticated kernel applied on $\Omega_t$.

Note that only a fraction of the points in $\hat{\mathcal{X}}_t \cap \mathcal{X}_t$ directly observe the exact function values. The function values on the rest ones in $\hat{\mathcal{X}}_t \setminus \mathcal{X}_t$ are estimated by interpolation, which captures the landscape of $f(x)$ by smoothing out the local fluctuations. To control the trade-off between the inaccurate estimations and the exact observations in $\Omega_t$, we design a stopping rule that optimizes the number of consistent queries in $\Omega_t$. The more consistent queries conducted in a given subspace, the more accurate observations could be obtained, albeit at the expense of a smaller remaining budget for exploring other regions.

The key features of CobBO are listed in Algorithm 1, with more details in the following sections. Several auxiliary components are utilized and presented in Appendix C to deal with a larger variety of problems and corner cases.

### 3.1 Block coordinate ascent and subspace selection

For Bayesian optimization, consider an infeasible assumption that each iteration can exactly maximize the function $f(x)$ in $\Omega_t$. This is not possible for one iteration but only if one can consistently query in $\Omega_t$, since the points converge to the maximum, e.g., under the expected improvement acquisition function with fixed priors [59] and the convergence rate can be characterized for smooth functions in the reproducing kernel Hilbert space [8]. However, even with this infeasible assumption, it is known that coordinate ascent with fixed blocks can cause stagnation at a non-critical point, e.g., for non-differentiable [67] or non-convex functions [49]. This motivates us to select a subspace with a variable-size coordinate block $C_t$ for each query. A good coordinate block can help the iterations to escape the trapped non-critical points. For example, one condition can be based on the result in [21] that assumes $f(x)$ to be differentiable and strictly quasi-convex over a collection of blocks. In practice, we do not restrict ourselves to these assumptions.

We induce a preference distribution $\pi_t$ over the coordinate set $I$, and sample a variable-size coordinate block $C_t$ accordingly. This distribution is updated at iteration $t$ through a multiplicative weights update method [2]. Specifically, the values of $\pi_t$ at coordinates in $C_t$ starts off uniform and increase in face of an improvement or decrease otherwise according to different multiplicative ratios $\alpha > 1$ and $\beta > 1$, respectively,

$$w_{t,j} = w_{t-1,j} \begin{cases} \alpha & \text{if } j \in C_t \text{ and } y_t > M_{t-1} \\ \frac{1}{\beta} & \text{if } j \in C_t \text{ and } y_t \leq M_{t-1} \\ 1 & \text{if } j \notin C_t \end{cases} ; \quad w_{0,j} = \frac{1}{D} ; \quad \pi_{t,j} = \frac{w_{t,j}}{\sum_{j=1}^{D} w_{t,j}} \quad (2)$$

This update characterizes how likely a coordinate block can generate a promising search subspace.

The multiplicative ratio $\alpha$ is chosen to be relatively large, e.g., $\alpha = 2.0$, and $\beta$ relatively small, e.g., $\beta = 1.1$, since the queries that improve the best observations $y_t > M_{t-1}$ happen more rarely than the opposite $y_t \leq M_{t-1}$.

How to dynamically select the size $|C_t|$? It is known that Bayesian optimization works well for low dimensions [15]. Thus, we specify an upper bound for the dimension of the subspace (e.g., $|C_t| \leq 30$). In principle, $|C_t|$ can be any random number in a finite set of possible block sizes $C$. This is different from the method that partitions the coordinates into fixed blocks and selects one according to, e.g., cyclic order [68], random sampling or Gauss-Southwell [46].
3.2 Backoff stopping rule for consistent queries

Applying BO on $\Omega_t$ requires a strategy to determine the number of consecutive queries for making a sufficient progress. This strategy is based on previous observations, thus forming a stopping rule. In principle, there are two different scenarios, exemplifying exploration and exploitation, respectively. Persistently querying a given subspace refrains from opportunistically exploring other coordinate combinations. Abruptly shifting to different subspaces does not fully exploit the potential of a given subspace.

CobBO designs a heuristic stopping rule in compromise. It takes the above two scenarios into joint consideration, by considering not only the number of consecutive queries that fail to improve the objective function but also other factors including the improved difference $M_t - M_{t-1}$, the point distance $||x_t - x_{t-1}||$, the query budget $T$ and the problem dimension $D$. On the one hand, switching to another subspace $\Omega_{t+1} (\neq \Omega_t)$ prematurely without fully exploiting $\Omega_t$ incurs an additional approximation error associated with the interpolation of observations in $\Omega_t$ projected to $\Omega_{t+1}$. On the other hand, it is also possible to over-exploit a subspace, spending high query budget on marginal improvements around local optima. In order to mitigate this, even when a query leads to an improvement, other factors are considered for sampling a new subspace.

3.3 Theoretical Analysis

One can view our block coordinate selection approach in section 3.1 as a combinatorial mixture of experts problem [10], where each coordinate is a single expert and the forecaster aims at choosing the best combination of experts in each step. Under this view, we bound the regret of our selection method with respect to the policy of selecting the best (unknown) block of coordinates at each step.

Assume that there is a fixed optimal choice $\mathcal{I}^*$ for the block of coordinates to pick at all steps. This block is characterized by improving the objective function for the largest number of times among all the possible coordinate blocks when performing Bayesian optimization over the corresponding subspaces. The following particular design of losses expresses this cause:

$$
\ell_{t,i} = \begin{cases} 
-\log(\hat{\alpha}) & \text{if } i \in C_t \text{ and } y_t > M_{t-1} \\
\log(\hat{\beta}) & \text{if } i \in C_t \text{ and } y_t \leq M_{t-1} \\
0 & \text{if } i \notin C_t 
\end{cases} \quad \hat{\alpha}, \hat{\beta} > 1
$$

(3)

as all the coordinates participating in the selected block incur the same loss that effectively rewards these coordinates for improving the objective and penalizes these for failing to improve the objective. All other coordinates that are not selected receive a zero loss and remain untouched.

Note that $\hat{\alpha}$ and $\hat{\beta}$ express the extent of reward and penalty, e.g., for $\hat{\alpha} = \hat{\beta} = e$ we have losses of $\ell_{t,i} \in \{-1, 1, 0\}$. Yet, $\hat{\alpha}$ is better chosen to be larger than $\hat{\beta}$, since the frequency of improving the objective is expected to be smaller.

The loss received by the forecaster is to reflect the same motivation. This is done by averaging the losses of the individual coordinates in the selected block, so that the size of the block does not matter explicitly, i.e., a bigger block should not incur more loss just due to its size but only due to its performance. Such that for each coordinate block $\mathcal{I}_t \subset \mathcal{I} = \{1, \cdots, D\}$ selected at time step $t$, the loss incurred by the forecaster is $L_{t,\mathcal{I}_t} = \frac{1}{|\mathcal{I}_t|} \sum_{i \in \mathcal{I}_t} \ell_{t,i}$. This is also the common loss incurred by all the coordinates participating in that block.

In each step we have the following multiplicative update rule of the weights associated with each coordinate (setting $\alpha = \hat{\alpha}^\eta$ and $\beta = \hat{\beta}^\eta$ yields the update rule in Eq. 2):

$$
w_{t,i} = w_{t-1,i} \cdot e^{-\eta \ell_{t,i}} = w_{t-1,i} \cdot \begin{cases} 
\hat{\alpha}^\eta & \text{if } i \in C_t \text{ and } y_t > M_{t-1} \\
1/\hat{\beta}^\eta & \text{if } i \in C_t \text{ and } y_t \leq M_{t-1} \\
1 & \text{if } i \notin C_t 
\end{cases}
$$

(4)

The probability $\tilde{\pi}_{t,\mathcal{I}_t}$ of selecting a certain coordinate block $\mathcal{I}_t$ is induced by $\pi_t$ as specified next. Thus the expected cumulative loss of the forecaster is:

$$
L_T = \sum_{t=1}^T \sum_{c \in \mathcal{C}} \sum_{t \in \mathcal{S}_c} \tilde{\pi}_{t,\mathcal{I}_t} \cdot \frac{1}{|\mathcal{I}_t|} \sum_{i \in \mathcal{I}_t} \ell_{t,i}
$$
Assume the best coordinate block is \( \mathcal{I}^* \), then the corresponding cumulative loss is:

\[
L^*_T = \sum_{t=1}^{T} L_t \mathbb{1}_{\mathcal{I}^*} = \sum_{t=1}^{T} \frac{1}{|\mathcal{I}^*|} \sum_{i \in \mathcal{I}^*} \ell_{t,i}
\]

We hence aim at bounding the regret \( \text{Regret}_T = L_T - L^*_T \).

**Theorem 1.** Sample from the combinatorial space of all possible coordinate blocks \( \mathcal{I}_t \in \bigcup_{c \in \mathcal{C}} \mathcal{S}_c \) with probability \( \hat{\pi}_t, \mathcal{I}_t = \prod_{i \in \mathcal{I}_t} w_{t,i}^{1/|\mathcal{I}_t|} / \sum_{c \in \mathcal{C}} \sum_{\mathcal{I} \in \mathcal{S}_c} \prod_{i \in \mathcal{I}} w_{t,i}^{1/|\mathcal{I}_t|} \). Then the update rule in Eq. 2 with \( \alpha = \tilde{\alpha} \), \( \beta = \tilde{\beta} \) and \( \eta = \log(\tilde{\alpha} \tilde{\beta})^{-1} \sqrt{T^{-1}|\mathcal{C}|D \log(D)} \) yields:

\[
\text{Regret}_t \leq O \left( (\log(\tilde{\alpha} \tilde{\beta}) \cdot \sqrt{T|\mathcal{C}|D \log(D)}) \right)
\]  

(5)

where \( w_{t,\mathcal{I}_t} = \prod_{i \in \mathcal{I}_t} w_{t,i}^{1/|\mathcal{I}_t|} \) is the geometric mean of weights in block \( \mathcal{I}_t \). The upper bound in Eq. 5 is tight, as the lower bound can be shown to be of \( \Omega(\sqrt{T \log(N)}) \) [23] where the number of experts is \( N = \sum_{c \in \mathcal{C}} \mathcal{S}_c \leq D^{|\mathcal{C}|} \) in our combinatorial setup, as typically \( |\mathcal{C}| \ll D \).

In practice, the direct sampling policy introduced in Theorem 1 involves high computational costs due to the exponential growth of combinations in \( D \). Thus CobBO suggests an alternative computationally efficient sampling policy with a linear growth in \( D \).

**Theorem 2.** Sample a block size \( c \in \mathcal{C} \) with probability \( p_c \) and \( c \) coordinates without replacement according to \( \pi_c \). Assume \( \mathcal{C} \supseteq \{1\} \), then the update rule in Eq. 2, with \( \alpha = \tilde{\alpha} \), \( \beta = \tilde{\beta} \) and \( \eta = \sqrt{\frac{\log(D)}{T \log(\tilde{\alpha} \tilde{\beta})^2 - \log(p_1)}} \geq 1 \) yields:

\[
\text{Regret}_t \leq O \left( \sqrt{(\log(\tilde{\alpha} \tilde{\beta})^2 - \log(p_1)) \cdot T \log(D)} \right)
\]  

(6)

where \( p_c > 0 \) for all \( c \in \mathcal{C} \) and \( \sum_{c \in \mathcal{C}} p_c = 1 \), e.g., uniformly set \( p_c = |\mathcal{C}|^{-1} \). The proof and detailed sampling policy are in Appendix A. The regret upper bound in Eq. 6 is tight, as the lower bound for an easier setup can be shown to be of \( \Omega(\sqrt{T \log(D)}) \) [23]. The implication on \( \eta \) is valid only for settings of a very high dimensionality and low query budget. In particular, CobBO is designed for this kind of problems.

**Remark:** Similar analysis and results follow when incorporating consistent queries from Section 3.2 and sampling a new coordinate block once every several steps. This is done by effectively performing less steps of aggregated temporal losses, as shown in Appendix A.

### 4 Numerical Experiments

This section presents detailed ablation studies of the key components presented in Section 3 and comparisons with other algorithms.

#### 4.1 Empirical analysis and ablation study

Ablation studies are designed to study the contributions of the key components in Algorithm 1 by experimenting with the Rastrigin function on \([-5, 10]^{50}\) with 20 initial points. The best performing run out of 5 experiments for each configuration is presented in Figure 3.

![Ablation study using Rastrigin on \([-5, 10]^{50}\) with 20 initial random samples](image-url)
**RBF interpolation**: RBF calculation is time efficient. Specifically, this is much beneficial in high dimensions. Figure 1 (left) shows the computation time of plain Bayesian optimization compared to CobBO’s. While the former applies GP regression using the Matérn kernel in the high dimensional space directly, the later applies RBF interpolation in the high dimensional space and GP regression with the Matérn kernel in the low dimensional subspace. This two-step composite kernel leads to a significant speed-up. Other time efficient alternatives are, e.g., the inverse distance weighting [26] and the simple approach of assigning the value of the observed nearest neighbour. Figure 3 (a) shows that RBF is the most favorable.

**Backoff stopping rule**: CobBO applies a stopping rule to query a variable number of points in subspace $\Omega_t$ (Section 3.2). To validate its effectiveness, we compare it with schemes that use a fixed budget of queries for $\Omega_t$. Figure 3 (b) shows that the stopping rule yields superior results.

**Coordinate blocks of a varying size**: CobBO selects a block of coordinates of a varying size $C_t$ (Section 3.1). Figure 3 (c) shows that a varying size is better than fixed.

**Preference probability over coordinates**: For demonstrating the effectiveness of coordinate selection (Section 3.1), we artificially let the function value only depend on the first 25 coordinates of its input and ignore the rest. It forms two separate sets of active and inactive coordinates, respectively. We expect CobBO to refrain from selecting inactive coordinates. Figure 4 shows the entropy of this preference probability $\pi_t$ over coordinates and the overall probability for picking active and inactive coordinate at each iteration. We see that the entropy decreases, as the preference distribution concentrates on the significant active coordinates.

![Figure 4: The preference probability focuses on active coordinates as the entropy decreases](image)

![Figure 5: Performance over low (left) medium (middle) and high (right) dimensional problems](image)

**4.2 Comparisons with other methods**

The default configuration for CobBO is specified in the supplementary materials. CobBO performs on par or outperforms a collection of state-of-the-art methods across the following experiments. Most of the experiments are conducted using the same settings as in TurBO [14], where it is compared with a comprehensive list of baselines, including BFGS, BOCK [47], BOHAMIANN, CMA-ES [22], BOBYQA, EBO [61], GP-TS, HeSBO [44], Nelder-Mead and random search. To avoid repetitions, we only show TuRBO and CMA-ES that achieve the best performance among this list, and additionally compare CobBO with BADS [1], REMBO [63], Differential Evolution (Diff-Evo) [56], Tree Parzen Estimator (TPE) [4] and Adaptive TPE (ATPE) [12].
4.2.1 Low dimensional tests

To evaluate CobBO on low dimensional problems, we use the lunar landing [38, 14] and robot pushing [62], by following the setup in [14]. Confidence intervals (95%) over 30 independent experiments for each problem are shown in Fig. 5.

Lunar landing (maximization): This controller learning problem (12 dimensions) is provided by the OpenAI gym [38] and evaluated in [14]. Each algorithm has 50 initial points and a budget of 1,500 trials. TuRBO is configured with 5 trust regions and a batch size of 50 as in [14]. Fig. 5 (upper left) shows that, among the 30 independent tests, CobBO quickly exceeds 300 along some good sample paths.

Robot pushing (maximization): This control problem (14 dimensions) is introduced in [62] and extensively tested in [14]. We follow the setting in [14], where TuRBO is configured with a batch size of 50 and 15 trust regions, each of which has 30 initial points. Each experiment has a budget of 10,000 evaluations. On average CobBO exceeds 10 within 5500 trials, as shown in Fig. 5 (lower left).

4.2.2 High dimensional tests

Since the duration of each experiment in this section is long, confidence intervals (95%) over repeated 10 independent experiments for each problem are presented.

Additive latent structure (minimization): As mentioned in Section 2, additive latent structures have been exploited in high dimensions. We construct an additive function of 56 dimensions, defined as $f_{56}(x) = \text{Ackley}(x_1) + \text{Levy}(x_2) + \text{Rastrigin}(x_3) + \text{Hartmann}(x_4) + \text{Rosenbrock}(x_5) + \text{Schwefel}(x_6)$, where the first three terms express the exact functions and domains described in Section 4.2.1, the Hartmann function on $[0, 1]^6$ and the Rosenbrock and Schwefel functions on $[-5, 10]^6$ and $[-500, 500]^6$, respectively.

We compare CobBO with TPE, BADS, CMA-ES and TuRBO, each with 100 initial points. Specifically, TuRBO is configured with 15 trust regions and a batch size 100. ATPE is excluded as it takes more than 24 hours per run to finish. The results are shown in Fig. 5 (upper middle), where CobBO quickly finds the best solution among the algorithms tested.

Rover trajectory planning (maximization): This problem (60 dimensions) is introduced in [62]. The objective is to find a collision-avoiding trajectory of a sequence consisting of 30 positions in a 2-D plane. We compare CobBO with TuRBO, TPE and CMA-ES, each with a budget of 20,000 evaluations and 200 initial points. TuRBO is configured with 15 trust regions and a batch size of 100, as in [14]. ATPE, BADS and REMBO are excluded for this problem and the following ones, as they all take more than 24 hours per run. Fig. 5 (lower middle) shows that CobBO has a good performance.

The 200-dimensional Levy and Ackley functions (minimization): We minimize the Levy and Ackley functions over $[-5, 10]^{200}$ with 500 initial points. TuRBO is configured with 15 trust regions and a batch size of 100. These two problems are challenging and have no redundant dimensions. For Levy, in Fig. 5 (upper right), CobBO reaches 100.0 within 2,000 trials, while CMA-ES and TuRBO obtain 200.0 after 8,000 trials. TPE cannot find a comparable solution within 10,000 trials in this case. For Ackley, in Fig. 5 (lower right), CobBO reaches the best solution among all of the algorithms tested.

Regarding running times, for Ackley, CobBO runs for 12.8 CPU hours and TuRBO-1 run for more than 80 CPU hours or 9.6 GPU hours. Most other methods either cannot make any progress or find far worse solutions.

5 Conclusion

CobBO is a variant of coordinate ascent tailored for Bayesian optimization with a stopping rule to switch coordinate subspaces. The sampling policy of subspaces is proven to have tight regret bounds with respect to the best subspace in hindsight. Combining this projection on random subspaces with a two-stage kernels for function value interpolation and GP regression, we provide a practical Bayesian optimization method of affordable computational costs in high dimensions. Empirically, CobBO consistently finds comparable or better solutions with reduced trial complexity in comparison with the state-of-the-art methods across a variety of benchmarks.
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Broader Impact

As stated in [32], Bayesian optimization is a powerful optimization technique used in a wide range of industries and applications, such as robotics [37, 9, 52], internet tech companies [18, 34], designing novel molecules for pharmaceutics [19], material design for increasing efficiency of solar cells [71], and aerospace engineering [31]. All of these settings have high-dimensional optimization problems, and advances in BO will reflect on improved capabilities on these fields as well. We have fully open-sourced our code for CobBO using the MIT license to be available for researchers and practitioners in these fields, and many others. The ability to optimize a larger number of parameters than has previously been possible will bring further improvements to and further accelerate work in these areas.

Checklist

The checklist follows the references. Please read the checklist guidelines carefully for information on how to answer these questions. For each question, change the default [TODO] to [Yes], [No], or
You are strongly encouraged to include a justification to your answer, either by referencing the appropriate section of your paper or providing a brief inline description. For example:

- Did you include the license to the code and datasets? [Yes] See Section ??.
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1. For all authors...
   (a) Do the main claims made in the abstract and introduction accurately reflect the paper’s contributions and scope? [Yes] See Section 3.
   (b) Did you describe the limitations of your work? [No]
   (c) Did you discuss any potential negative societal impacts of your work? [Yes] See Broader Impact.
   (d) Have you read the ethics review guidelines and ensured that your paper conforms to them? [Yes]

2. If you are including theoretical results...
   (a) Did you state the full set of assumptions of all theoretical results? [Yes] See Section 3.3.
   (b) Did you include complete proofs of all theoretical results? [Yes] See the Appendix A.

3. If you ran experiments...
   (a) Did you include the code, data, and instructions needed to reproduce the main experimental results (either in the supplemental material or as a URL)? [Yes] In the supplemental material.
   (b) Did you specify all the training details (e.g., data splits, hyperparameters, how they were chosen)? [Yes] See Table 2 in the appendix, which contains the default hyperparameters.
   (c) Did you report error bars (e.g., with respect to the random seed after running experiments multiple times)? [Yes] See Fig.5 and Fig.8-10.
   (d) Did you include the total amount of compute and the type of resources used (e.g., type of GPUs, internal cluster, or cloud provider)? [Yes] See page 9, line 324.

4. If you are using existing assets (e.g., code, data, models) or curating/releasing new assets...
   (a) If your work uses existing assets, did you cite the creators? [N/A]
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   (e) Did you discuss whether the data you are using/curating contains personally identifiable information or offensive content? [N/A] It does not contain personal identifiable information or offensive content.

5. If you used crowdsourcing or conducted research with human subjects...
   (a) Did you include the full text of instructions given to participants and screenshots, if applicable? [N/A]
   (b) Did you describe any potential participant risks, with links to Institutional Review Board (IRB) approvals, if applicable? [N/A]
   (c) Did you include the estimated hourly wage paid to participants and the total amount spent on participant compensation? [N/A]