UNIVERSITY OF CALIFORNIA Los Angeles

Studies in Hyperparameter Tuning, Design Selection and Optimization

A dissertation submitted in partial satisfaction of the requirements for the degree Doctor of Philosophy in Statistics

by

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ABSTRACT OF THE DISSERTATION

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This dissertation explores advanced optimization techniques, focusing on hyperparameter tuning, design strategy selection, and novel optimization methods. First, we investigate a modified Differential Evolution (DE) algorithm for generating uniform projection designs, emphasizing the importance of hyperparameter configuration. We analyze the surface structure of these hyperparameters and provide guidelines for optimizing settings under various conditions. Next, we examine the role of initial design choices in prediction and sequential optimization using Efficient Global Optimization (EGO), demonstrating that uniform projection designs outperform traditional strategies such as maximin distance designs, particularly in high-dimensional spaces. Finally, we introduce a Kriging-based sequential region shrinking method that integrates EGO to efficiently reduce the search space by targeting promising data points. Comparative results show that this method not only requires fewer computational resources than conventional tuning techniques like grid and random search, but also outperforms other Bayesian optimization methods such as TREGO. These findings offer significant contributions to the optimization field, enhancing both theoretical understanding and practical applications.

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To my parents, Joseph and Jane, — and my siblings, Tabitha, Naomi, Jasper, and Lilian,— who among so many other things ensured that I gave my all to accomplish this task.

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CHAPTER 1

Introduction

1.1 Introduction to Optimization Algorithms

Optimization algorithms play a pivotal role in solving complex problems where analytical solutions are either unavailable or computationally infeasible. Among these, gradient-free algorithms are particularly valuable for functions that are non-differentiable, discontinuous, or noisy. Two prominent approaches in this category are metaheuristic algorithms and Bayesian optimization, each offering unique strengths and applications.

Metaheuristic algorithms are inspired by natural processes such as evolution, swarm behavior, or physical phenomena. Popular examples include Differential Evolution, Genetic Algorithms, Particle Swarm Optimization and Simulated Annealing. These algorithms excel in exploring vast and complex solution spaces, making them highly effective for global optimization. Their stochastic nature and population-based search strategies allow them to escape local optima and adapt to diverse problem landscapes. On the other hand, Bayesian optimization takes a probabilistic approach, using surrogate models (such as Gaussian processes) to approximate the objective function. By sequentially selecting points that balance exploration and exploitation, Bayesian optimization achieves efficiency in scenarios where function evaluations are expensive.

Both approaches are integral to my research, offering robust, gradient-free methods to address the challenges of experimental design construction and data-driven model optimization.

1.2 Metaheuristic Algorithms

Metaheuristic algorithms are high-level problem-solving frameworks designed to find nearoptimal solutions to complex optimization problems. These algorithms are particularly useful
for problems where traditional optimization methods may struggle due to factors like highdimensional search spaces, non-linearity, discontinuity, or multimodality. The key characteristics of metaheuristic algorithms include:

- 1. **General-Purpose**: Applicable to a wide range of problems without requiring problem-specific knowledge.
- 2. **Approximate Solutions**: Focus on finding good-enough solutions in a reasonable time rather than guaranteed optimal solutions.
- 3. **Stochastic Nature**: Often incorporate randomization to explore the search space and avoid local optima.
- 4. **Iterative Process**: Evolve candidate solutions over several iterations using a combination of exploration (searching broadly) and exploitation (refining promising areas).

1.2.1 Evolutionary Algorithms

Evolutionary algorithms are a family of optimization techniques inspired by the principles of natural evolution, such as selection, mutation, recombination or crossover, and survival of the fittest. These algorithms operate on a population of candidate solutions, iteratively improving them based on a defined fitness function. Common examples include Genetic Algorithms (GA), Differential Evolution (DE), and Evolution Strategies (ES), each with unique mechanisms for generating and selecting solutions.

Genetic Algorithm (GA)

Genetic Algorithms (GAs) are search heuristics inspired by the principles of natural selection and genetics, first introduced by Holland (1975). In GAs, a population of candidate solutions, called chromosomes, evolves over iterations, or generations, through processes like selection, crossover, and mutation. The algorithm aims to find an optimal or near-optimal solution to a given problem by mimicking the survival-of-the-fittest principle in biological systems (Mitchell 1998).

The GA process begins with the initialization of a population, typically generated randomly within the solution space. Each chromosome in the population is evaluated using a fitness function, which quantifies the quality of the solution it represents. Based on fitness scores, a selection mechanism such as roulette wheel or tournament selection is used to choose parent chromosomes for reproduction. The better the fitness, the more likely a chromosome is selected, promoting better solutions (Kawachi and Ando 1992).

Reproduction involves crossover, where segments of parent chromosomes are combined to produce offspring. Common crossover methods include single-point, two-point, and uniform crossover. This recombination introduces new solution candidates into the population. Additionally, mutation is applied to offspring with a small probability to alter random genes, maintaining diversity and preventing premature convergence to local optima (De Jong 1975).

The GA process iterates through selection, crossover, mutation, and fitness evaluation until a termination criterion is met, such as a maximum number of generations or convergence to a solution. GAs have been successfully applied to various optimization problems, including scheduling, engineering design, and machine learning hyperparameter tuning. Their adaptability and ability to handle complex, non-linear problems have made them a popular choice in optimization research (Whitley 1994).

However, GAs face challenges like computational cost due to their population-based nature and sensitivity to parameter settings, such as mutation rate, crossover probability, and population size. Hybrid approaches combining GAs with other optimization methods, like

local search or simulated annealing, have been developed to address these issues, enhancing convergence and efficiency (Michalewicz 2013).

Differential Evolution (DE)

Differential Evolution (DE) is a population-based optimization algorithm introduced by Storn and Price (1997), specifically designed for continuous optimization problems. Unlike GAs, DE focuses on vector differences to guide its search, making it particularly effective for high-dimensional and non-linear optimization tasks. DE is simple yet powerful, with few parameters to tune (Price, Storn, and Lampinen 2006).

The DE process starts with initializing a population of candidate solutions as vectors within the solution space. Each vector is evaluated using an objective function to determine its fitness. The algorithm then generates trial vectors for each population member through mutation, crossover, and selection. Mutation in DE involves creating a donor vector by adding the scaled difference between two randomly chosen population vectors to a third vector, a strategy unique to DE (Mezura-Montes, Velázquez-Reyes, and Coello Coello 2006).

Crossover combines the donor vector with the target vector to produce a trial vector. This process can be uniform or binomial, depending on whether crossover is controlled by a fixed or probabilistic scheme. The trial vector is evaluated, and if it outperforms the target vector in terms of fitness, it replaces the target in the population for the next generation. This selection mechanism ensures steady improvement in the population's quality over iterations (Das and Suganthan 2010).

DE's strength lies in its robustness and ability to balance exploration and exploitation through its mutation and selection strategies. Its simple structure and limited parameter requirements, primarily the mutation factor and crossover rate, make it easy to implement and tune. DE has been applied to diverse areas, including function optimization, neural network training, and control system design (Miettinen 1999).

Despite its effectiveness, DE may face challenges in multimodal optimization problems,

where it can become trapped in local optima. Variants of DE, such as adaptive DE and self-adaptive DE, have been developed to address these issues by dynamically adjusting parameters during the search process. Hybrid approaches integrating DE with other techniques, like gradient-based methods, have also shown promise in improving convergence and tackling complex optimization problems (Qin, Huang, and Suganthan 2008).

Evolution Strategies (ES)

Evolution Strategies (ES) are a prominent subclass of evolutionary algorithms tailored for optimizing real-valued continuous functions. Developed in the 1960s by Ingo Rechenberg and Hans-Paul Schwefel, these algorithms are inspired by natural evolution, focusing on adaptation and mutation while eschewing crossover mechanisms typical in other evolutionary approaches like Genetic Algorithms (Schwefel 1977; Vent 1975). ES are population-based, relying on iterative cycles of mutation, selection, and reproduction to refine candidate solutions. Their robustness, simplicity, and adaptability make them particularly effective for solving high-dimensional, noisy, or non-linear optimization problems (Beyer and Schwefel 2002).

The standard ES workflow begins with a population of candidate solutions, each represented as a vector of real-valued parameters. Mutation, the core variation operator, introduces randomness by perturbing these parameters using Gaussian noise. Selection mechanisms, such as $(\mu + \lambda)$ or (μ, λ) , then choose the fittest individuals to form the next generation. While $(\mu + \lambda)$ strategies retain both parents and offspring for survival, promoting stability, (μ, λ) strategies consider only offspring, encouraging greater exploration of the search space (Schwefel 1993). These mechanisms make ES versatile in balancing exploration and exploitation.

A significant extension of ES is the Directed Variation, introduced by Zhou and Li (2003), which enhances the mutation process by guiding it with directional information derived from the problem landscape (Zhou and Li 2003). Unlike standard random isotropic mutations,

directed variation biases mutation steps toward promising regions of the search space. Zhou and Li proposed frameworks for incorporating directional probabilities or fitness gradients into the mutation process, improving convergence rates and solution quality in complex optimization problems. This refinement allows ES to adapt dynamically to the fitness landscape, avoiding premature convergence while maintaining effective exploration.

The self-adaptation mechanism in ES, developed in earlier works (Schwefel 1981), complements directed variation by evolving strategy parameters, such as mutation step sizes, alongside candidate solutions. This approach enables the algorithm to adjust its search behavior based on the landscape's characteristics. When combined with directed variation, self-adaptation can achieve even better convergence properties by synergizing global exploration with locally informed refinement (Beyer and Arnold 2001). Directed variation thus enriches the evolutionary process by incorporating both dynamic parameter adjustment and problem-specific guidance.

Directed variation is particularly advantageous in applications involving constrained or multi-modal optimization. Zhou and Li's work highlights its efficacy in improving optimization outcomes for tasks with complex, irregular landscapes (Zhou and Li 2003). By biasing mutations toward beneficial directions, ES with directed variation can escape local optima and converge efficiently to global optima. This capability has been demonstrated in fields such as engineering design, robotics, and neural network training.

The theoretical foundation of directed variation draws on insights from gradient-based optimization while retaining the flexibility of stochastic methods. Zhou and Li's approach blends deterministic directional cues with stochastic exploration, bridging the gap between classical optimization techniques and evolutionary computation. This hybrid methodology underscores the broader adaptability of ES, allowing it to tackle a diverse range of optimization challenges (Beyer and Schwefel 2002; Zhou and Li 2003).

Another notable advancement is Covariance Matrix Adaptation (CMA-ES), introduced by Hansen and Ostermeier (2001). CMA-ES dynamically adjusts the covariance matrix of the mutation distribution to align with the topology of the fitness function. This technique enables ES to search more effectively along relevant dimensions of the landscape, particularly in ill-conditioned or anisotropic problems.

In summary, ES, enriched with innovations like directed variation, self-adaptation, and CMA-ES, offer a robust and versatile framework for optimization. Together, these advancements have solidified ES as a cornerstone of evolutionary computation, enabling it to excel in solving real-world problems with complex, high-dimensional, and noisy landscapes.

1.2.2 Swarm intelligence (SI)

Swarm intelligence algorithms are optimization techniques inspired by the collective behavior of decentralized, self-organized systems in nature, such as flocks of birds, schools of fish, or ant colonies. These algorithms use the interaction of simple agents to solve complex problems without central control. Swarm intelligence models excel in parallelism, adaptability, and robustness, making them well-suited for solving optimization problems across diverse domains, including engineering, scheduling, and machine learning (Kennedy and Eberhart 1995).

Particle Swarm Optimization (PSO)

Introduced by Kennedy and Eberhart (1995), PSO is inspired by the flocking behavior of birds and schooling of fish. In PSO, each individual in the swarm, called a particle, represents a candidate solution. The particles explore the solution space by updating their positions based on their own best-known position, the best-known position of their neighbors, and the overall global best position. These updates are guided by two key parameters: cognitive and social components, which balance personal learning and group influence.

The mathematical foundation of PSO lies in velocity and position updates. A particle's velocity is influenced by three components: inertia, which maintains momentum; the cognitive term, which pulls the particle toward its own best position; and the social term, which

attracts the particle toward the global best position. By iteratively updating positions and velocities, PSO achieves convergence to near-optimal solutions. Its simplicity and efficiency have made it popular for a wide range of problems, from neural network training to resource allocation (Clerc and Kennedy 2002).

One of the primary advantages of PSO is its ease of implementation, as it requires fewer hyperparameters compared to other optimization algorithms. However, it faces challenges such as premature convergence, especially in high-dimensional or multimodal optimization problems. Variants of PSO, such as constriction coefficient PSO and adaptive PSO, have been developed to address these limitations and improve exploration and exploitation (Shi and Eberhart 1998).

Ant Colony Optimization (ACO)

Ant Colony Optimization (ACO), another prominent SI algorithm, was introduced by Dorigo (1996) and is inspired by the foraging behavior of ants. Ants use pheromone trails to communicate and collectively find optimal paths between their nest and a food source. In ACO, artificial ants construct solutions by traversing a graph representation of the problem, depositing pheromones on paths to indicate their quality.

The algorithm operates iteratively, where ants build solutions based on probabilistic rules influenced by pheromone levels and heuristic information, such as edge lengths or costs. After constructing solutions, pheromones are updated: stronger paths receive more pheromone reinforcement, while weaker paths evaporate over time. This dynamic update mechanism balances exploration of new paths and exploitation of promising ones, allowing ACO to identify optimal or near-optimal solutions (Dorigo 2007).

ACO has been successfully applied to combinatorial optimization problems, such as the Traveling Salesman Problem, vehicle routing, and scheduling. Its strengths lie in its adaptability and ability to integrate heuristic knowledge. However, ACO's performance depends on parameters like evaporation rate, pheromone influence, and heuristic weighting, which

require careful tuning to avoid premature convergence or excessive exploration (Blum 2005).

Despite their differences, both PSO and ACO share a reliance on decentralized control and emergent behavior to solve problems. While PSO excels in continuous optimization due to its particle dynamics, ACO is better suited for discrete optimization tasks because of its graph-based representation. Hybrid approaches combining PSO and ACO have been developed to leverage the strengths of both algorithms, particularly for problems with mixed discrete and continuous variables (Talbi 2009).

Swarm intelligence algorithms, including PSO and ACO, are inherently parallelizable, making them attractive for large-scale optimization problems. Their ability to balance exploration and exploitation is a key factor in their success. Recent research has focused on adapting these algorithms to dynamic and multi-objective optimization problems, where solution landscapes change over time or involve conflicting objectives (Coello, Pulido, and Lechuga 2004).

One challenge in swarm intelligence algorithms is their computational complexity, especially for high-dimensional problems or large populations. To address this, researchers have developed variants like distributed swarm intelligence and hybrid metaheuristics. These approaches aim to reduce computational cost while maintaining solution quality, enabling the application of swarm intelligence in fields like robotics, logistics, and bioinformatics.

Moreover, advances in hardware, such as GPUs and parallel computing platforms, have further enhanced the scalability of swarm intelligence algorithms. These developments allow for real-time applications in domains such as autonomous navigation, sensor networks, and real-time scheduling, where traditional optimization methods may fall short (Yang and Karamanoglu 2013).

In conclusion, swarm intelligence algorithms like PSO and ACO have revolutionized optimization by drawing inspiration from nature. Their versatility, adaptability, and robustness have enabled solutions to a wide range of complex problems. With ongoing advancements in hybridization, parameter tuning, and computational power, swarm intelligence continues

to be a cornerstone of modern optimization research.

1.2.3 Simulated Annealing (SA)

Simulated Annealing (SA) is a probabilistic optimization technique inspired by the annealing process in metallurgy, where materials are heated and slowly cooled to alter their physical properties. Proposed by Kirkpatrick, Gelatt Jr, and Vecchi (1983), SA solves complex optimization problems by mimicking this physical process, exploring a solution space to find near-optimal or global optima. Unlike traditional optimization methods, it allows temporary acceptance of worse solutions to escape local minima, making it especially effective for problems with rugged landscapes.

The algorithm operates by iteratively modifying a candidate solution and evaluating its quality using a predefined objective function. A key feature of SA is its acceptance criterion, which is governed by a probabilistic function dependent on temperature and the difference in solution quality. This function, often based on the Metropolis criterion, reduces the likelihood of accepting worse solutions as the temperature decreases. This simulated "cooling" process, if managed properly, ensures convergence to an optimal solution while maintaining exploration of the solution space early on (Aarts and Korst 1989).

Temperature scheduling plays a critical role in SA's effectiveness. Commonly, exponential decay is used, where the temperature decreases geometrically over iterations. However, alternative cooling schedules like linear or logarithmic decay can also be employed, each with unique trade-offs in convergence speed and solution quality. The choice of cooling schedule and initial temperature significantly affects performance, as improper tuning may lead to suboptimal solutions or slow convergence.

SA has been successfully applied to a wide range of problems, including combinatorial optimization tasks like the Traveling Salesman Problem (TSP), job scheduling, and graph partitioning. For instance, Johnson et al. (1991) demonstrated SA's efficiency in solving the TSP by leveraging its capability to explore and exploit the problem's complex search space.

In continuous optimization, modifications to the algorithm have enabled its use in problems like parameter estimation and engineering design.

While SA has many strengths, it also faces challenges, such as the need for careful parameter tuning and the potential for slow convergence. Hybrid approaches combining SA with other optimization techniques, such as genetic algorithms or gradient-based methods, have been proposed to address these issues. Such hybrid algorithms often leverage SA's exploration capabilities while benefiting from the exploitation strengths of complementary methods (Yang 2010).

Theoretical work on SA has shown that it can asymptotically converge to the global optimum under certain conditions. Specifically, if the cooling schedule is sufficiently slow, the algorithm can theoretically explore all possible solutions and guarantee convergence. However, these conditions are often impractical due to computational constraints, necessitating heuristic adjustments in real-world applications (Hajek 1988).

In conclusion, simulated annealing remains a versatile and robust optimization technique suitable for a wide array of applications. Its success lies in its balance between exploration and exploitation, as well as its capacity to escape local minima. Advances in hybrid approaches and theoretical insights continue to expand its scope and effectiveness, making it a critical tool in modern optimization practices.

1.3 Bayesian Optimization

Beyond the metaheuristic optimization methods lies the notion of Bayesian Optimization. Bayesian optimization has become a widely adopted approach for black-box optimization problems, particularly for its sequential nature in handling noisy and non-convex functions commonly encountered in real-world scenarios. The framework often relies on Efficient Global Optimization (EGO), a sequential strategy that iteratively searches for optimal solutions by balancing exploration of the search space and exploitation of known high-performing

regions (Jones, Schonlau, and Welch 1998). At each step, Bayesian optimization evaluates new points based on their potential to improve the objective function while accounting for the uncertainty of the model predictions. This iterative, or sequential, nature allows Bayesian optimization to progressively refine its estimates, focusing on promising regions and thereby enhancing efficiency.

In Bayesian optimization, a Gaussian Process (GP) is typically used as the surrogate model, offering a flexible and powerful approach for modeling complex functions (Rasmussen and Williams 2006). GPs are particularly valuable because they predict the objective function's value at unexplored points and provide an estimate of prediction uncertainty, which can guide decisions on where to sample next. The choice of acquisition function, also known as the utility function, is central to Bayesian optimization, as it defines the criteria for selecting the next sample point. Common acquisition functions include Probability of Improvement (PI), Expected Improvement (EI), Upper Confidence Bound (UCB), and Lower Confidence Bound (LCB), each designed to guide the search process differently depending on the optimization strategy.

The Probability of Improvement (PI) function, for instance, favors points likely to outperform the current best-known solution, focusing on regions with high potential for incremental gains. Expected Improvement (EI), on the other hand, seeks points that offer the highest potential for improvement by integrating both the predicted mean and the uncertainty of the GP model, balancing exploration and exploitation more effectively (Mockus 1994). The Upper Confidence Bound (UCB) and Lower Confidence Bound (LCB) functions apply a degree of confidence to the GP predictions, where UCB emphasizes exploration by targeting areas with high uncertainty, while LCB may be used to address risk-averse or cost-sensitive problems by minimizing potential loss (Srinivas et al. 2009).

These acquisition functions collectively enable Bayesian optimization to adapt dynamically based on the task requirements. Each function aligns with different optimization objectives, allowing Bayesian optimization to tailor its approach to complex landscapes and

focus computational resources on the most promising areas of the search space. This adaptability, coupled with the sequential nature of EGO, makes Bayesian optimization a powerful method for tackling diverse and challenging optimization problems.

1.4 Leveraging Differential Evolution and Bayesian Optimization

As previously discussed, Differential Evolution (DE) is a versatile metaheuristic algorithm primarily used for optimizing continuous functions. Its simplicity and efficiency make it appealing across a wide range of applications, including experimental design. Notably, its strong global search capabilities, robust performance, and flexibility across diverse problem domains (Storn and Price 1997), combined with its high convergence speed for specific challenges (Babu and Jehan 2003) and inherent parallelizability (Kukkonen and Lampinen 2006), have established it as a highly favorable choice for design construction.

However, using Differential Evolution for discrete data tasks, such as experimental design construction, presents challenges. Applying DE in such cases requires modifications to its foundational structure to effectively manage discrete variables and discrete search space. Recent work by Stokes, Wong, and Xu (2024)introduced a modified DE algorithm specifically tailored for these discrete tasks, highlighting the need for adaptations to extend DE's functionality beyond continuous domains. Despite these advancements, the algorithm's performance remains highly sensitive to several hyperparameters, which play a critical role in influencing outcomes.

This sensitivity to hyperparameters emphasizes the importance of understanding and exploring the hyperparameter landscape. Developing optimized settings across different problem setups is essential for achieving reliable and efficient performance. Tailoring hyperparameters for specific tasks or datasets can greatly improve outcomes, but doing so requires a structured approach to hyperparameter tuning. As such, there is a need for strategies that systematically optimize hyperparameter configurations to align with varied and complex problem requirements.

In this dissertation, we employ the DE algorithm and a novel localized region shrinkage Bayesian optimization in the construction of the uniform projection designs. Three studies are done and the structure of these studies is laid as follows.

The first study delves into the construction of space-filling designs focusing mainly on Uniform Projection Designs (UPDs) by investigating the surface structure of DE's hyper-parameters and their respective contributions. UPDs, introduced by Sun, Wang, and Xu (2019), are a special class of space-filling designs that ensure an even distribution of sample points across all lower-dimensional projections of a high-dimensional design space. By analyzing the impact of hyperparameters on the performance of the modified DE algorithm, the study aims to derive optimal settings for generating efficient UPDs using a second-order model. Through comparisons of various experimental designs and surrogate models, the research provides crucial guidelines for enhancing DE's performance. The insights gained from this study are intended to equip practitioners with effective strategies for optimizing hyperparameter configurations in practical applications.

Another critical aspect of optimization algorithms is the initial design choices, which can significantly impact both prediction accuracy and the efficiency of sequential optimization processes. The second study utilizes gaussian process and various initial experimental designs to model test functions determining the prediction power of the initial design. In addition, the effect of the initial design on the optimization performance via active learning is evaluated. This study highlights how initial design strategies greatly influence prediction prowess and early optimization results, with their effects diminishing as iterations proceed toward the global optimum. In particular, the research demonstrates that distance-based designs, like maximin distance designs, struggle in high-dimensional settings, whereas Uniform Projection Designs consistently perform well across varying dimensionalities.

In response to the challenges associated with traditional optimization approaches, the third study introduces a novel Kriging-based sequential region shrinking method, which effectively incorporates the EGO algorithm. This innovative method aims to progressively reduce the region of interest by focusing on the most promising data points during each iteration. The efficiency of this approach is validated through its application to various well-known physical test functions, where it significantly reduces the computational resources required compared to traditional hyperparameter tuning techniques like grid and random search. Furthermore, it demonstrates advantages over other Bayesian optimization strategies, such as TREGO, highlighting its practicality in resource-constrained environments.

Together, these studies provide a comprehensive exploration of advanced optimization techniques, ranging from hyperparameter tuning in DE to effective design strategy selection, culminating in novel optimization methods. The findings contribute valuable insights to both theoretical research and practical applications, showcasing the evolving landscape of optimization strategies. By bridging the gaps in existing methodologies, this work aims to enhance the efficacy of optimization processes across diverse fields, ultimately leading to more robust and efficient solutions for complex challenges.

CHAPTER 2

Tuning Differential Evolution Algorithm for Constructing Uniform Projection Designs

2.1 Introduction

Experimental design construction is a fundamental aspect of research and data-driven inquiry, aimed at organizing experimental runs to extract maximum information while minimizing resource use. By strategically selecting input combinations, well-constructed designs ensure that researchers can identify key factors, estimate model parameters, and predict responses accurately (Montgomery 2017). The primary goal is to balance efficiency and comprehensiveness, whether in exploring high-dimensional spaces, optimizing processes, or assessing system robustness. Central to this process is the notion of space-filling, where design points are distributed to capture variations across the entire experimental domain, providing a solid foundation for modeling and inference (Santner et al. 2003).

Different design strategies address diverse experimental objectives and constraints. For instance, factorial and fractional factorial designs are widely used to study the main effects and interactions of factors systematically, while response surface designs, such as central composite and Box-Behnken designs, support optimization and curvature estimation (Myers, Montgomery, and Anderson-Cook 2016). In other cases, non-traditional approaches like space-filling designs (e.g., Latin hypercube or maximin designs) and discrepancy-based designs (e.g., maxpro or uniform designs) are preferred for high-dimensional and computational experiments (Joseph 2016). Each method offers unique strengths, and the choice depends on

the experimental goals, computational resources, and the nature of the underlying system being studied. Through thoughtful design construction, researchers can ensure that their experiments are not only scientifically rigorous but also cost-effective and impactful.

In the quest for robust experimental designs, Uniform Projection Designs (UPDs) have emerged as a powerful tool for ensuring uniformity across all low-dimensional projections of the design space (Sun, Wang, and Xu 2019). UPDs, introduced by Sun, Wang, and Xu (2019), are specialized space-filling designs characterized by robust performance across various design criteria and impressive space-filling properties in multiple dimensions. These designs are particularly valuable in high-dimensional settings where relationships between subsets of factors often carry critical information. However, existing algorithms for generating UPDs remain underexplored, highlighting an important area for further research. To construct UPDs, we leverage the use of Differential Evolution (DE)

Recently, Stokes, Wong, and Xu (2024) proposed a DE-based approach for constructing order-of-addition designs, showcasing its efficiency compared to other metaheuristic algorithms, such as Simulated Annealing, Threshold Accepting, Genetic Algorithms, and Particle Swarm Optimization. Inspired by their findings, we adapt and extend their modified DE algorithm for the construction of UPDs.

The performance of the DE algorithm is significantly influenced by its hyperparameters, which dictate the learning process of the optimization (Price, Storn, and Lampinen 2006). An inappropriate setting of these hyperparameters can lead to suboptimal performance of the DE algorithm. While the DE algorithm proposed by Stokes, Wong, and Xu (2024) encompasses several hyperparameters, their effects remain largely unexamined. Therefore, we aim to conduct a comprehensive study of the hyperparameters to elucidate their impacts on the algorithm's performance, providing insights that could enhance its effectiveness.

The challenge of determining the optimal hyperparameter settings for any learning process has been widely studied. Two primary frameworks dominate this landscape: the model-based framework and the model-free framework. Model-based hyperparameter optimization

focuses on tuning hyperparameters by approximating the true learning algorithm, while model-free methods approach the optimization problem without parametric assumptions. Relevant literature on model-based hyperparameter optimization includes works by Falkner, Klein, and Hutter (2018), Hutter, Hoos, and Leyton-Brown (2011), Li et al. (2018), Lujan-Moreno et al. (2018), Mockus, Tiesis, and Zilinskas (1978), Snoek, Larochelle, and Adams (2012), Wu, Chen, and Liu (2020), and Zoph and Le (2016). In contrast, model-free frameworks encompass techniques such as manual search, grid search, random search, genetic algorithms, and orthogonal array tuning methods (Liashchynskyi and Liashchynskyi 2019).

Our approach leverages various types of designs and models to investigate the effectiveness of DE in constructing UPDs. Specifically, we aim to address three objectives: (i) identifying useful design types in understanding the surface structure extended by the DE hyperparameters, (ii) determining the most effective models, and (iii) developing an efficient algorithm for the construction of UPDs. We employ different types of models to evaluate the performance of different designs, enabling us to visualize the response surface of the DE algorithm's hyperparameters. This framework outlines the data generation, modeling, and analysis procedures. The insights gained from our analysis provide a general guideline for optimal hyperparameter settings necessary for generating superior uniform projection designs. This approach is quite different from the Lujan-Moreno et al. (2018) method which used a 2^k factorial design with the response surface method (RSM) and ridge regression for screening to select the important factors in the data. While Lujan-Moreno et al. (2018) focuses on factor screening and selection with the traditional RSM approach, we emphasize on the comparisons of different types of designs and surrogate models in approximating the surface structure of the DE algorithm.

2.2 Differential Evolution Algorithm

Originating from the pioneering work of Storn and Price (1997), Differential Evolution (DE) has emerged as a powerful heuristic optimization algorithm, drawing inspiration from the

mechanisms of biological evolution. To simulate survival-of-the-fittest dynamics, DE treats each candidate or agent as a chromosome made up of several genes and implements mutation and crossover procedures that allow beneficial genes to persist into future generations (Storn and Price 1997). DE operates on the principle of population-based search, where a set of candidate solutions evolves over successive generations towards optimal or near-optimal solutions. At its core, DE employs mutation, crossover, and selection operators to iteratively improve the quality of solutions. The algorithm's efficacy stems from its robustness, simplicity, and ability to handle non-linear, non-convex, and noisy optimization landscapes.

Without loss of generality, we assume that we want to minimize a real-valued objective function ϕ over an m-dimension space Ω . It has five steps

- 1. Genetic Representation: Let π_1, \ldots, π_N be the initial population, where each agent $\pi_i = (\pi_{i1}, \ldots, \pi_{im})$ is randomly chosen from Ω .
- 2. **Mutation**: Mutation expands the search space of the current population. For each i = 1, ..., N, mutation produces a potential donor ν_i in Ω by adding the weighted difference of two agents to a third, all randomly chosen and distinct from the target (π_i) , that is,

$$\nu_i = \pi_a + w(\pi_b - \pi_c) \tag{2.1}$$

where $a \neq b \neq c$ are randomly chosen three distinct numbers from $1, \ldots, N$, and they are all different from i.

3. Crossover: Crossover blends the current generation of agents with the population of potential donors in order to form candidates for the next generation known as trial agents. For each i = 1, ..., N, one of the m variables of ν_i is randomly selected to directly enter the trial agent u_i . In this way, one variable is forced to change so that each u_i will certainly differ from its original target π_i . Next, with probability pCR, more variables are taken from u_i and placed in the trial agent. Whichever variables do not take their value from the donor inherit their original value from π_i . Assuming j_0 is a

random number from $1, \ldots, m$, this process can be written as follows: for $j = 1, \ldots, m$,

$$u_{ij} = \begin{cases} \nu_{ij} & \text{with probability pCR or if } j = j_0, \\ \pi_{ij} & \text{otherwise} \end{cases}$$

4. **Selection**: Selection creates the next generation of agents by comparing each target to its respective trial agent. The trial agent is adopted if it leads to an improvement and is discarded otherwise. For minimization problems, this process is given by,

$$\pi_i = \begin{cases} u_i & \text{if } \phi(u_i) < \phi(\pi_i) \\ \pi_i & \text{otherwise} \end{cases}$$

5. **Repeat**: Repeat steps 2 through 4 over many generations until a specified stopping condition is satisfied.

Though quite simplistic, its ability to balance exploration and exploitation makes it ideal for solving non-linear and multimodal problems.

Since experimental designs lie on a discrete and constrained space, we leverage the modified DE by Stokes, Wong, and Xu (2024). This is to ensure that the resulting mutated design is feasible. The proposed method modifies the mutation step by using the swap mutation (Michalewicz 2013), one of the structural mutation operators in genetic algorithm. In this operator, two positions in a solution are randomly selected, and their values are exchanged. This maintains the feasibility of the solution by preserving its permutation structure while introducing variation to explore the search space. The swap mutation is computationally efficient and effective at diversifying the population, reducing the risk of premature convergence. The mutation step is controlled by mutation rate pMut which is the probability of swapping two elements. In addition borrowing from PSO, they induced the influence of the global best solution with a probability pGBest. This yielded an algorithm which contained the following hyperparameters:

• NP - The size of the population (N).

- itermax the maximum number of iterations/generations used.
- pCR Probability of crossover.
- pMut Probability of mutation.
- pGBest Probability of using the global best for mutation.
- pSelf Probability of using the current agent for mutation.

They proposed three different choices of the initial agent to be mutated to obtain the proposal agent. This lead to three different variants which they referred to as DE1 which uses the global best, DE2 which uses the current agent and DE3 which uses a random agent.

Regarding the hypermarameters, the first two, NP and itermax, determine the budget size, whereas the other four hyperparameters affect the evolution process. The hyperparameters, pGBest and pSelf, determine the probability of using the global best and the current agent in the mutation process, respectively. There is a constraint between these two hyperparameters, that is, pGBest + pSelf \leq 1. The question that arises is how these hyperparameters interact with each other. Also whether we can do better than the proposed fixed probabilities to obtain the better settings for the DE algorithm for design generation.

In this study, we shall consider values between [10, 100] for NP, [500, 1500] for itermax, and [0.05, 0.95] for pCR, pMut and pGBest. We fix pSelf = (1 - pGBest)/2 so that there is an equal chance for selecting a current agent and a random agent if the global best is not used.

2.3 Designs for Hyperparameter Settings

Various designs can be used to set the DE hyperparameters before the optimization process. As the functions optimized by DE are often complex with many local minima, one has to carefully choose the initial point for the optimization process. These initial points are

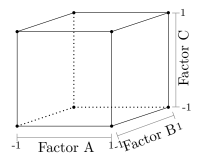


Figure 2.1: Geometric illustration of a 2^3 full factorial design

determined using any of the methods discussed below. In each subsection below one method is described, and its benefits and drawbacks are discussed.

Full factorial designs (FFD)

Factorial designs are a research method for studying the effects of multiple independent variables on a response variable, formalized by Sir Ronald A. Fisher in the early 20th century (Fisher 1935). These designs typically involve defining factors at two or three levels, forming a grid of all possible combinations, resulting in 2^m or 3^m observations for m factors. Figure 2.1 shows a 2^3 full factorial design.

Full factorial designs sample points at the corners of a hypercube, ensuring uniform distribution across the design space. They allow for the analysis of main effects and interactions (Montgomery 2017), but can be complex and require larger sample sizes for adequate power (Tabachnick and Fidell 2019).

To mitigate the need for larger samples, fractional factorial designs were introduced by David John Finney (Finney 1945). These designs use a fraction of runs based on the sparsity-of-effects principle, focusing on main effects and lower-order interactions. They are expressed as 2^{m-p} or 3^{m-p} , depending on the factors set as products of others. Selecting defining relations for fractional designs is essential, with criteria such as maximum resolution and minimum aberration guiding this process (Wu and Hamada 2011).

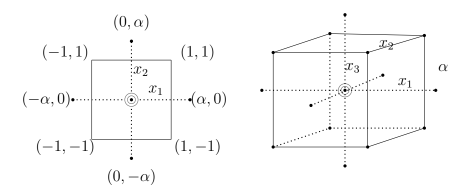


Figure 2.2: CCD for m = 2 and m = 3

Central composite designs (CCDs)

Introduced by Box and Wilson (1951) as an extension of factorial designs, they were developed as a way to efficiently fit quadratic response surfaces and identify optimal process settings in industrial experiments. CCDs are full or fractional factorial designs that are augmented with two additional sets of sampling points described as "center" and "axial or star" points (Box and Wilson 1951). The center point is defined by all factors being set at their center level. The CCD uses 2m axial points, each of which is defined by all but one factor being at their center level and the level of the remaining factor is denoted by α , which is generally chosen to be between 1 and \sqrt{m} (Montgomery 2017). The basic concepts of the CCD for m = 2 and m = 3 are depicted in Figure 2.2, where the bold dots are the design points.

Orthogonal array composite designs (OACD)

Introduced by Xu, Jaynes, and Ding (2014), an OACD is a class of composite designs based on a two-level factorial design and a three-level orthogonal array (OA). An OA of n runs, m columns, s levels, and strength t, denoted by OA (n, s^m, t) , is an $n \times m$ matrix in which all s^t level-combinations appear equally often in every $n \times t$ submatrix (Wu and Hamada 2011). For example, a 2^m factorial design can be viewed as OA $(n, 2^m, t)$ with $n = 2^m$ and t = m.

Similarly, a three-level OA can be written as OA $(n, 3^m, t)$. Thus, an OACD is a composite design which consists of a two-level factorial design as its factorial points, a three-level OA as its additional points, plus any number of center points (Luna et al. 2022).

Space filling designs

While the previously discussed designs utilize sampling points that are at the boundaries of the design space, space filling designs generate samples that are dispersed throughout the multidimensional design space and not just at the boundary of the design space. These designs are important in sampling a surface as they could capture important regions thereby minimizing the bias between the true structure of the surface and the estimated surface from the sampled points (Gardner et al. 2006; Giunta, Wojtkiewicz, and Eldred 2003). They are of various types depending on the approach used to construct them, e.g., sampling-based – Latin Hypercube designs, distance-based – maximin designs, and distribution – based uniform designs (Burton, Xu, and Yi 2022).

Latin hypercube designs (LHDs) – Based on McKay (1992)'s Latin hypercube sampling, it divides the range of each factor into bins of equal size, where n also corresponds to the number of samples to be generated resulting in a total of n^m combinations where m is the number of factors being considered. The n samples are then randomly generated such that for all one-dimensional projections, there will be only one sample in each bin.

Maximin distance designs – Introduced by Johnson, Moore, and Ylvisaker (1990), this design aims at spreading out the design points in the design space by maximizing the minimum distance between any two design points. It thus tends to place a large proportion of points at the corners and on the boundaries of the design space. Mathematically, this can be formulated as follows. Suppose we want to construct an n-run design in m factors. Let the design region be the unit hypercube \mathcal{X} and let the design be $D = \{x_1, \dots, x_n\}$, where each design point x_i is in $\mathcal{X} = [0, 1]^m$. The maximin design

optimizes the function below:

$$\max_{D} \min_{i \neq j} d\left(\boldsymbol{x}_{i}, \boldsymbol{x}_{j}\right),$$

where $d(\mathbf{x}_i, \mathbf{x}_j)$ is the distance between the points \mathbf{x}_i and \mathbf{x}_j .

Maximin Latin hypercube designs – Unlike Latin hypercube designs, maximin distance designs do not have good projection properties for each factor. Morris and Mitchell (1995) proposed to overcome this problem by searching for the maximin distance design within the class of Latin hypercube designs. They also proposed to use the following criterion to achieve maximin distance:

$$\min_{D} \left\{ \sum_{i=1}^{n-1} \sum_{j=i+1}^{n} \frac{1}{d^{p}(\boldsymbol{x}_{i}, \boldsymbol{x}_{j})} \right\}^{1/p}$$
(2.2)

where p > 0 is chosen large enough, say p = 15.

Maximum projection (MaxPro) designs – Although maximin Latin hypercube designs ensure good space-filling in m dimensions and uniform projections in each dimension, their projection properties in two to m-1 dimensions are not known. By the effect sparsity principle (Wu and Hamada 2011), only a few factors are expected to be important. To curb this, Joseph, Gul, and Ba (2015) proposed a different criterion:

$$\min_{D} \psi(D) = \left\{ \frac{2}{n(n-1)} \sum_{i=1}^{n-1} \sum_{j=i+1}^{n} \frac{1}{\prod_{l=1}^{m} (x_{il} - x_{jl})^{2}} \right\}^{1/m}.$$
 (2.3)

and showed that the design that minimizes $\psi(D)$ tends to maximize its projection capability in all subspaces of factors, and thus named these designs as maximum projection designs.

2.4 Modeling

We consider three different frameworks to model the data: (a) Linear Model, (b) Kriging Model and (c) Heterogeneous Gaussian Process (HetGP).

Linear Model (lm)

For m quantitative factors, denoted by x_1, \ldots, x_m , a second-order linear model is defined as

$$y = \beta_0 + \sum_{i=1}^{m} \beta_i x_i + \sum_{i=1}^{m} \beta_{ii} x_i^2 + \sum_{i < j} \beta_{ij} x_i x_j + \epsilon$$
 (2.4)

where $\beta_0, \beta_i, \beta_{ii}$, and β_{ij} are the intercept, linear, quadratic, and bilinear (or interaction) terms, respectively, and ϵ is the error term. This model is simple and provides a straightforward way to model and understand relationships between the response variable and the factors. The main effects are easy to to interpret.

Kriging Model (km)

Proposed by South African geostatistician Krige (1951), Kriging is one of the methods used to interpolate intermediate values, whereby these intermediate values are modeled using Gaussian Process (GP) which is governed by prior co-variances. It provides a probabilistic prediction of the output variable, as well as an estimate of the uncertainty of the prediction (Chevalier, Picheny, and Ginsbourger 2014). The kriging predictors interpolating the observations are assumed to be noise-free (Roustant, Ginsbourger, and Deville 2012). Intermediate interpolated values obtained by Kriging are the best linear unbiased predictors.

The Kriging model consists of two parts: a trend and a GP. The trend part is often modeled as a regression on some fixed basis functions. In the specific case where the basis functions reduce to a constant function, it is referred to as ordinary Kriging (Roustant, Ginsbourger, and Deville 2012). The general form is as given below

$$Y(\mathbf{x}) = \sum_{i=1}^{k} \beta_i f_i(\mathbf{x}) + Z(\mathbf{x}), \qquad (2.5)$$

where f_1, \ldots, f_k are k basis functions, β_1, \ldots, β_k are corresponding regression coefficients, and $Z(\boldsymbol{x})$ is a stationary GP with zero mean and covariance function ψ . The covariance

function ψ completely defines the behavior of the Gaussian Process Z(x). It is defined as

$$\psi\left(\boldsymbol{x}_{i}, \boldsymbol{x}_{j}\right) = \operatorname{Cov}\left(Z\left(\boldsymbol{x}_{i}\right), Z\left(\boldsymbol{x}_{j}\right)\right) = \sigma^{2} \prod_{l=1}^{m} K\left(h_{l}; \theta_{l}\right), \tag{2.6}$$

where σ^2 is the scale parameter called the process variance, $h_l = |x_{i,l} - x_{j,l}|$, $x_{i,l}$ and $x_{j,l}$ are the lth elements of the i^{th} run \boldsymbol{x}_i and the j^{th} run \boldsymbol{x}_j , and $K(h;\theta)$ is the correlation function. The parameters θ_l chosen for the correlation function $K(h_l;\theta_l)$ must be positive. Otherwise the correlation function will not be feasible. These parameters are chosen to be physically interpretable in the same unit as the corresponding variables. They are often referred to as the *characteristic length-scales* by Rasmussen and Williams (2006). Popular correlation functions include Gaussian, Matérn, and power-exponential family correlation functions. The Matérn function with parameter $\nu = 5/2$ is often chosen as the default when fitting kriging models. It is defined as:

$$K(h;\theta) = \left(1 + \frac{\sqrt{5}h}{\theta} + \frac{5h^2}{3\theta^2}\right) \exp\left(-\frac{\sqrt{5}h}{\theta}\right). \tag{2.7}$$

The unknown parameters can be estimated via MLE or cross validation. In R, the km function in the DiceKriging package was used to fit the kriging model.

Heteroskedastic Gaussian Process (HetGP)

HetGP follows the simplifying assumption in the computer experiments literature in using a mean zero GP, which shifts all of the modeling effort to the covariance structure (Binois, Gramacy, and Ludkovski 2018). Observation model is given by

$$y_i = y(\mathbf{x}_i) = f(\mathbf{x}_i) + \varepsilon_i, \quad \varepsilon_i \sim \mathcal{N}(0, r(\mathbf{x}_i)),$$
 (2.8)

where $f(\mathbf{x}_i)$ is a GP with covariance or kernel $k(\cdot, \cdot)$ and $r(\mathbf{x}_i)$ is the variance of ϵ_i which depends on \mathbf{x}_i . The kernel $k(\cdot, \cdot)$ is positive definite, with parameterized families such as the Gaussian or Matérn being typical choices. If $r(\mathbf{x}_i) = \tau^2$ is a constant, then the process is homoskedastic. In matrix notation, the modeling framework just described is equivalent to

writing

$$Y \sim \mathcal{N}\left(\mathbf{0}, \mathbf{K}_n + \mathbf{\Sigma}_n\right)$$

where \mathbf{K}_n is the $n \times n$ matrix with (i, j) coordinate $k(\mathbf{x}_i, \mathbf{x}_j)$, and $\mathbf{\Sigma}_n = \mathrm{Diag}(r(\mathbf{x}_1), \dots, r(\mathbf{x}_n))$ is the variance matrix of the vector of independent noise ε_i .

Given the kernel function $k(\cdot, \cdot)$ and data $\boldsymbol{y} = (y_1, \dots, y_n)^\top$, multivariate normal (MVN) conditional identities provide a predictive distribution at site $\boldsymbol{x} : Y(\boldsymbol{x}) \mid \boldsymbol{y}$, which is Gaussian with parameters

$$\mu(\boldsymbol{x}) = \mathbb{E}(Y(\boldsymbol{x}) \mid \boldsymbol{y}) = \mathbf{k}(\boldsymbol{x})^{\top} (\mathbf{K}_n + \boldsymbol{\Sigma}_n)^{-1} \boldsymbol{y},$$

$$\sigma^2(\boldsymbol{x}) = \mathbb{V} \operatorname{ar}(Y(\boldsymbol{x}) \mid \boldsymbol{y}) = k(\boldsymbol{x}, \boldsymbol{x}) + r(\boldsymbol{x}) - \mathbf{k}(\boldsymbol{x})^{\top} (\mathbf{K}_n + \boldsymbol{\Sigma}_n)^{-1} \mathbf{k}(\boldsymbol{x}),$$

where $\mathbf{k}(\boldsymbol{x}) = (k(\boldsymbol{x}, \boldsymbol{x}_1), \dots, k(\boldsymbol{x}, \boldsymbol{x}_n))^{\top}$. In R, the mleHetGP function in hetGP package is used to fit this model with the default Gaussian kernel.

2.5 The Data Generation Process

We aim to obtain optimal DE hyperparameter settings that can be used to generate UPDs.

The Objective Function: Uniform Projection Design Criterion

Proposed by Sun, Wang, and Xu (2019), the uniform projection criterion solely focuses on two-dimensional projections. This is due to two factor interactions being more important than three-factor or higher-order interactions. The motivating idea was that although designs with low discrepancy have good uniformity in the full-dimensional space, they can have bad projections in lower dimensional spaces, which is undesirable when only a few factors are active. Thus designs with better projection properties are preferred. Sun, Wang, and Xu (2019) argued that the uniform projection designs scatter points uniformly in all dimensions and have good space-filling properties in terms of distance, uniformity and orthogonality.

The uniform projection design criterion is defined using the centered L_2 -discrepancy. For an $n \times m$ design $D = (x_{ik})$ with s levels from $\{0, 1, \ldots, s-1\}$, its (squared) centered L_2 -discrepancy is defined as

$$CD(D) = \frac{1}{n^2} \sum_{i=1}^{n} \sum_{j=1}^{n} \prod_{k=1}^{m} \left(1 + \frac{1}{2} |z_{ik}| + \frac{1}{2} |z_{jk}| - \frac{1}{2} |z_{ik} - z_{jk}| \right)$$
$$- \frac{2}{n} \sum_{i=1}^{n} \prod_{k=1}^{m} \left(1 + \frac{1}{2} |z_{ik}| - \frac{1}{2} |z_{ik}|^2 \right) + \left(\frac{13}{12} \right)^m,$$

where $z_{ik} = \left(2x_{ik} - s + 1\right)/(2s)$. Then the uniform projection criterion is to minimize

$$\phi(D) = \frac{2}{m(m-1)} \sum_{|u|=2} \text{CD}(D_u), \qquad (2.9)$$

where u is a subset of $\{1, 2, ..., m\}$, |u| denotes the cardinality of u and D_u is the projected design of D onto dimensions indexed by the elements of u. The $\phi(D)$ is the average centered L_2 -discrepancy values of all two-dimensional projections of D.

We implemented the DE and uniform projection criterion in the package UniPro. The following code generates an $n \times m$ UPD with s levels

where NP, itermax, pMut, pCR and pGBest are DE hyperparameters described in Section 2, and seed is an optional seed for random number generators that ensures reproducibility.

As the task of design generation is quite complex, only 3 design sizes are considered. A UPD of size 30×3 is considered as a small and easy task, 50×5 as a medium task and 70×7 as a large and difficult task. We only consider the construction of designs with s = n so that the resulting UPD is an LHD.

Training and testing data

Designs discussed in Section 2.3 are used to determine the parameter settings for the DE algorithm hyperparameters. Specifically, we construct five designs: a CCD with 43 runs (ccd3_43), an OACD with 50 runs (oacd3_50), 50-run random LHD, 50-run maximin LHD, and 50-run maxpro LHD. All designs have five factors, one for each hyperparameter. Each

run corresponds to a setting of the five hyperparameters. The CCD and OACD have 3 levels while the rest have 50 levels. The levels are linearly interpolated within the minimum and maximum factor values for each hyperparameter. Generating the test data from the 3^5 factorial, 243-run random LHD, and the 4^5 factorial seems good enough. This is because the random LHD enjoys the maximum space-filling property in all one dimensions, while the 3^5 and 4^5 factorial designs cover the entire 5-dimensional input space in a uniform fashion.

Given the target design size $(n \times m)$ and a setting of the five hyperparameters, we run the UniPro function to generate an $n \times m$ UPD and the resulting $\phi(D)$ value defined in (2.9). This is recorded as the response value for that particular hyperparameter setting and target design size. For each setting, the DE algorithm is replicated ten times yielding ten replicates for the response. These are then aggregated to obtain the mean and the standard deviation of the response. Thus we obtain five training datasets for each target size.

The same procedure is taken to generate the testing dataset with the exception that the designs used for the hyperparameter setting combinations being a 3⁵ full factorial design, a random LHD with 243 runs, a combination of these two, and a 4⁵ full factorial design.

Density plots of the response for the testing and training data are presented in Figure 2.3 for the target size 50×5 . All of the distributions are skewed to the right. For a 50×5 UPD all the training designs lead to similar minimum $\phi(.)$ values, around 0.17, whereas different types of designs lead to different maximum $\phi(.)$ values. Indeed, all space filling designs have maximum $\phi(.)$ values around 0.28, while the factorial designs and the hybrid design have a maximum $\phi(.)$ values around 0.34. The narrower range of the $\phi(.)$ values suggests that the space filling designs do not explore the entire space of hyperparameters.

Model evaluation

For each training dataset, we fit the three models descibed in Section 2.4 and test on the four testing datasets. Designs are evaluated by considering their ability to collect informative data for building a statistical model that specifies the relationship between the response and

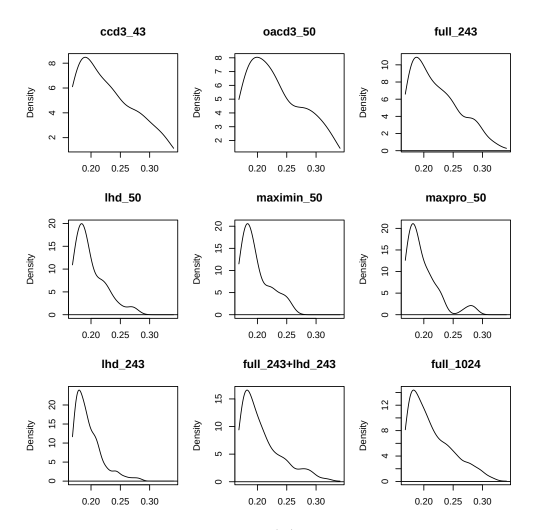


Figure 2.3: Density plots of the $\phi(D)$ values with target size 50×5

Table 2.1: Comparison of designs and model evaluations with target size 30×3

	(a) Testing on the 3 ⁵ FFD						(b) Testing on the 243 LHD						
	correlation			RMSE			correlation			RMSE			
Design	lm	km	hetGP	lm	km	hetGP	lm	km	hetGP	lm	km	hetGP	
ccd3_43	0.88	0.88	0.88	1.51	1.59	1.62	0.65	0.03	0.60	1.62	3.75	1.51	
oacd3_50	0.88	0.94	0.93	1.62	1.25	1.32	0.68	0.63	0.61	1.94	2.30	2.22	
lhd_50	0.41	0.36	0.34	3.19	3.30	3.27	0.28	0.20	0.19	1.08	1.14	1.14	
maximin_50	0.71	0.73	0.74	2.86	3.19	3.03	0.64	0.63	0.64	0.66	0.66	0.65	
maxpro_50	0.71	0.62	0.66	2.35	2.97	2.72	0.69	0.71	0.74	0.74	0.69	0.61	
	(c) Testing on the 3 ⁵ FFD+243 LHD						(d) Testing on the 4 ⁵ FFD						
	correlation			RMSE			С	orrela	tion	RMSE			
Design	lm	km	hetGP	lm	km	hetGP	lm	km	hetGP	lm	km	hetGP	
ccd3_43	0.84	0.60	0.83	1.57	2.88	1.57	0.84	0.66	0.85	1.55	2.83	1.50	
oacd3_50	0.82	0.83	0.83	1.79	1.85	1.83	0.85	0.87	0.87	1.68	1.68	1.68	
lhd_50	0.42	0.35	0.36	2.38	2.47	2.45	0.45	0.39	0.37	2.47	2.57	2.56	
maximin_50	0.69	0.63	0.68	2.08	2.31	2.19	0.74	0.75	0.76	2.16	2.38	2.27	
maxpro_50	0.74	0.60	0.68	1.75	2.15	1.98	0.74	0.67	0.71	1.80	2.22	2.03	

the hyperparameters, which is measured by the test root mean squared error (RMSE). The correlation (ρ) between the response and the predicted together with the RMSE are reported.

2.6 Results and Analysis

Table 2.1(a)(b) and Figure 2.4(a)(b) present comparison of designs and model evaluations with target size 30×3 for testing the two 243-run data sets. One striking observation is that the performance of the training data set depends on the nature of the testing data set. The

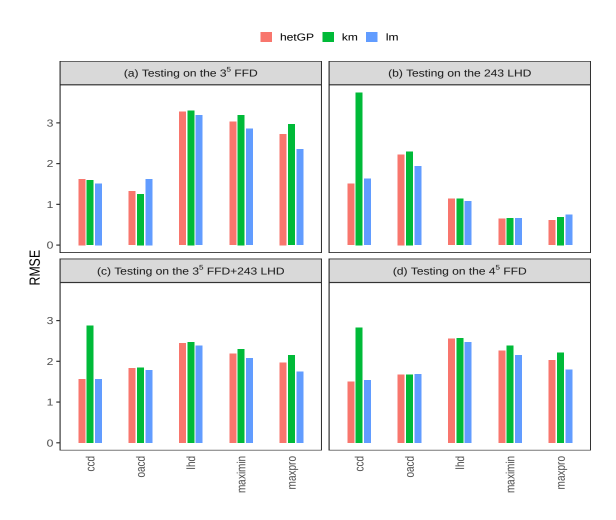


Figure 2.4: Comparison of RMSE with target size 30×3

composite designs, CCD and OACD, seem to be better when tested on the 3⁵ FFD while the space filling designs (random LHD, maximin LHD, and maxpro LHD) did better when tested on the 243-run random LHD. As this does not give a general idea as to which designs might perform better in general, we invoke the combined data with 486 (runs 3⁵ FFD and 243-run LHD) and the 4⁵ FFD as the testing dataset. Here we see that the composite designs perform better than the space filling designs; see Table 2.1(c)(d) and Figure 2.4(c)(d). The 50-run random LHD performed the worst in terms of correlation regardless of the testing data. The correlation is strikingly low whereas the RMSE is high. This might be due to randomness, but it does show the weakness of the random LHD.

One bizarre observation from Table 2.1(b) is the correlation of 0.03 when using the CCD as the training design and testing it on the 243-run random LHD. This value is strikingly lower than any other values given in Table 2.1. No apparent reason could be deduced as to why this is so. Multiple replications indicated that this is not an error. From all the results, we can deduce the robustness of OACD over CCD. This gives a reason to use OACD for the hyperparameter initialization.

With regards to the models, there seems to be no striking observation to be made as to whether one fitting method performs better than the other two, with exception for one 30×3 case when the kriging model fitting to the CCD training data had a much higher RMSE value than the other cases.

The three models have quite different assumptions. The linear model assumes a polynomial trend and independent random errors with homoskedastic variance. The Kriging model assumes a stationary covariance structure, that is, the covariance between two points in the DE hyperparameter surface structure depends only on the distance or spatial lag between those points, and not on their specific locations within the domain. The HetGP model assumes a heteroskedastic variance-covariance structure. For the DE algorithm, the homoskedastic and stationary assumptions are questionable. Due to this, the HetGP model is preferred to the linear model and the kriging model. However, the linear model is easy

to interpret and fits as well as the HetGP model, and from the results, there is no striking difference between the two. We use the linear model to determine factor importance and optimal hyperparameter settings.

Tables 2.3-2.4 and Figures 2.9-2.10 in the Appendix show results when the target design sizes are 50×5 and 70×7 , respectively. Looking at the results, apart from the random LHD with 50 runs, previously stated observations are upheld.

A natural question is why composite designs perform better than the space filling designs. We perceive that the hyperparameters at the boundaries lead to some extreme cases in this experiment and the composite designs do capture this phenomena while the space filling designs do not. Figure 2.5 presents the histograms of the distances from design points to the design center for all the designs, where each column is rescaled to [-1,1] and the euclidean distance from each point to the center of the design is calculated. This gives an insight as to why the composite designs, OACD and CCD, tend to perform better than the space filling designs. This is because the composite designs tend to capture information lying at the boundaries compared to the space filling designs which tend to capture the information lying at the center of the design. This is confirmed by the notion that three of the hyperparameters tend to be optimized around their highest level as discussed in the next section.

2.7 Factor Importance and Optimal Settings

The results obtain call for a deeper look into the model and how each factor is involved in the surface approximation. This enables us to have a better picture of the surface generated by the DE hyperparameters.

Table 2.2 shows the estimated coefficients of the second-order models based on the four different training datasets: CCD, OACD, maximin LHD, and maxpro LHD, for the target size 30×3 . Looking at the various models above, the model obtained using the maxpro LHD as the training data is the worst performing model. It has the lowest adjusted R^2 of

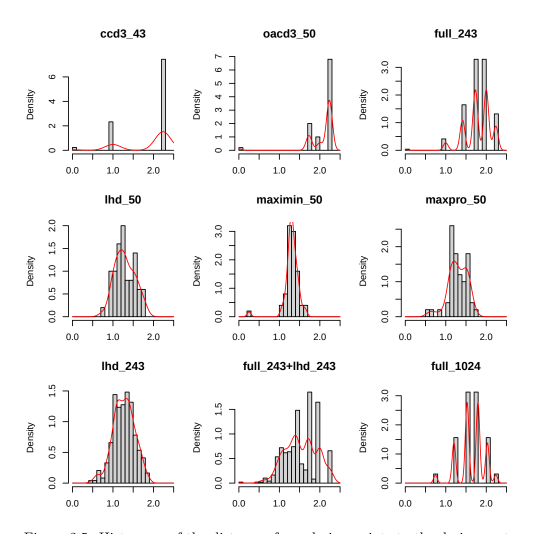


Figure 2.5: Histogram of the distances from design points to the design center

				_
	ccd3_43	oacd3_50	maximin_50	maxpro_50
(Intercept)	0.3815***	0.3876***	0.3878***	0.3814***
NP	-0.0134***	-0.0143^{***}	-0.0042^{***}	-0.0055^*
pMut	0.0020	0.0028	0.0043***	0.0045^{*}
pGBest	-0.0204***	-0.0224***	-0.0047^{***}	-0.0082^{***}
pCR	-0.0022	-0.0030	0.0004	0.0007
itermax	-0.0146***	-0.0152^{***}	-0.0046***	-0.0021
itermax_q	0.0090	-0.0081	0.0011	0.0057
NP_q	0.0119	0.0080	0.0022	-0.0004
pCR_q	0.0071	0.0074	-0.0008	0.0049
$pGBest_q$	0.0083	0.0192*	0.0035	0.0159***
pMut_q	0.0150	0.0174*	0.0083***	0.0104*
NP:pMut	0.0058	0.0064*	-0.0002	-0.0037
NP:pGBest	-0.0041	-0.0038	0.0017	0.0006
NP:pCR	0.0002	-0.0007	-0.0006	-0.0002
NP:itermax	0.0049	0.0033	0.0023	0.0052
pMut:pGBest	-0.0080^*	-0.0093***	-0.0089^{***}	-0.0139**
pMut:pCR	0.0203***	0.0197***	0.0042**	0.0018
pMut:itermax	0.0032	0.0025	-0.0061^{***}	0.0004
pGBest:pCR	0.0034	0.0036	0.0000	0.0051
pGBest:itermax	0.0057	0.0068**	0.0076***	0.0058
pCR:itermax	0.0037	0.0021	0.0018	0.0040
R^2	0.9034	0.9235	0.8970	0.7297
$Adj. R^2$	0.8157	0.8708	0.8260	0.5433
Num. obs.	43	50	50	50

^{***}p < 0.001; **p < 0.01; *p < 0.05

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Table 2.2: Statistical models

only 0.54. This model does not capture important main effects. For example, it indicates that the number of iterations (itermax) for optimization is not significant. Yet this is well known to be important. On the other hand, the model obtained by using OACD as the training dataset performs the best. It has an adjusted R^2 of 0.87 and captures important main effects and interactions. In addition, CCD might be a little worse than OACD because of the fewer number of points (43) used for training compared to the other models which used 50 points. The model from the CCD does not identify any of the quadratic effects to be significant while the other models do.

Looking at the corresponding p-values from the models obtained, it is evident that all five hyperparameters are important. The main effects of three hyperparameters (NP, itermax and pGBest) are very significant, whereas the several interactions involving one of the other two hyperparameters (pMut and pCR) are also very significant. The probability of using the global best (pGBest) is quite important because its main effect is highly significant in all the models in Table 2.2. It can also be inferred that the maximum number of iterations (itermax) and the population size (NP) are important. The linear models indicate that to minimize the objective function, it is ideal to use a larger pGBest, increase the population size (NP) and increase the number of iterations (itermax). The population size and the number of iterations could be constrained by the available budget.

With regards to interactions, all significant interaction terms involve either pMut or pGBest. The interaction of pMut and pGBest is negative. On the other hand, the interaction between pMut and pCR is positive. As this is a minimization problem, this positive interaction indicate that the two variables need to be either at their low levels or one has to be low while the other high. Both should not be at their high levels.

To have a better understanding of the interactions, we examine some interaction plots from the full 4⁵ factorial, shown in Figure 2.6. Here, the number of population (NP), number of iterations (itermax) and the probability of using global best (pGBest) are held at their highest levels, i.e., 100, 1500 and 0.95, respectively. These results are consistent

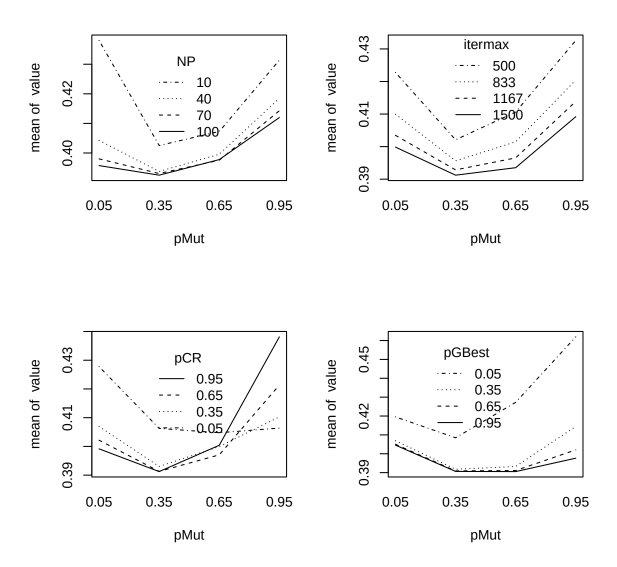


Figure 2.6: Interaction plots involving pMut based on the 4^5 FFD and target size 30×3 .

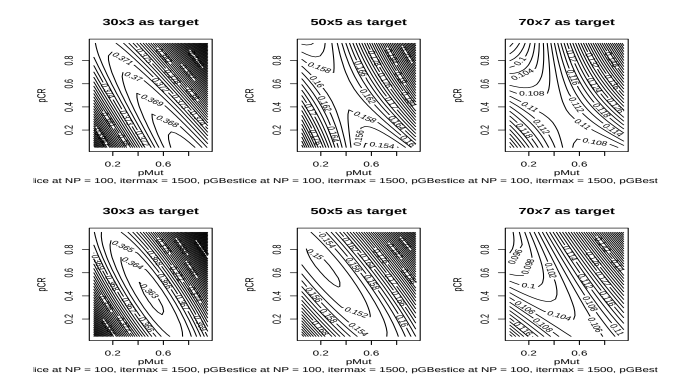


Figure 2.7: Contour plots of pMut and pCR while fixing other hyperparameters at high levels. Top row uses CCD as the training data; bottom row uses OACD as training data.

with the second order models in Table 2.2, where NP, itermax and pGBest are all negative and significant. Thus they should be set at the highest level to minimize the response values. These results are consistent for other target design sizes. For pMut and pCR, their interaction is more complicated as it is positive and thus a further analysis is called for to determine the preferred levels to set these parameters. We use contour plots to visualize the two parameters when the other three are held at their highest level.

Figure 2.7 shows the contour plots of pMut and pCR while fixing NP=100, itermax=1500 and pGBest=0.95. From the contour plot based on CCD and OACD training data set, and target size 30×3 , we note that the response value could be minimized by taking values over the off diagonal. Either a smaller pMut and larger pCR or a larger pMut and a smaller pCR. This is also the case for target sizes 50×5 and 70×7 . Searching the optimal setting is achieved by varing pMut from $0.05, 0.15, \ldots, 0.95$ with fixed parameters pCR = 1 - pMut.

For each setting of pMut and pCR, the DE algorithm is run 100 times to construct 100 designs while fixing NP = 100, itermax = 1500 and pGBest = 0.95. The average response values are then calculated across these runs, allowing us to identify the optimal settings of pMut and pCR for each target size. This approach results in the following optimal settings of (pMut, pCR): (0.95, 0.05), (0.25, 0.75), and (0.15, 0.85) for target size 30×3 , 50×5 and 70×7 respectively.

The optimal settings are compared with two DE variants, DE1 and DE4, described by Stokes, Wong, and Xu (2024). DE1 uses only the global best solution (pGBest = 1), while DE4 is a hybrid approach with pGBest = 0.5, where the global best, the current agent, and a random agent are selected with probabilities of 0.5, 0.25, and 0.25, respectively, for each column independently. Both DE1 and DE4 utilize fixed values of pMut = 0.1 and pCR = 0.5.

For the comparison, the optimal settings for pMut and pCR, determined for each target size, are used while maintaining pGBest = 0.95. In all cases, NP = 100 and itermax = 1500 are set, and the DE algorithm is executed 100 times to construct 100 designs for each method. Figure 2.8 presents boxplots of the $\phi(\cdot)$ values of these designs. The results demonstrate that DE with optimal settings (DEoptim) produces significantly better uniform projection designs compared to the two variants provided by Stokes, Wong, and Xu (2024), confirming the effectiveness of hyperparameter tuning.

2.8 Conclusion

This paper compared small designs in exploring the response surface of a DE algorithm hyperparameters. Five numerical hyperparameters were considered: population size, the number of maximum iterations, probability of crossover, probability of mutation, and probability of using the global best for mutation. Various composite designs and space-filling designs for selecting combinations of these hyperparameters were also examined. The performance of a design was evaluated via building a second order model, a kriging model and

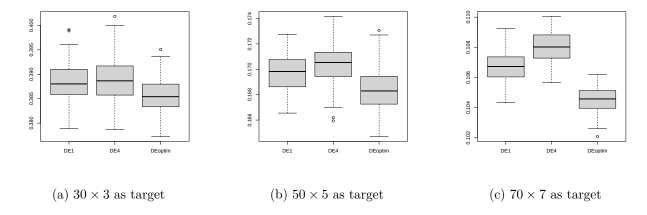


Figure 2.8: Performance of the DE algorithms under three settlings: DE1, DE4 and DEoptim (optimal settings)

a heterogeneous GP model. The performance was measured in terms of testing RMSEs and correlation. The comparison was made based on data simulated using the uniform projection criterion. Under the settings considered, the comparison demonstrates that OACD and CCD are the better choices over space-filling designs for exploring the response surface of the DE algorithm hyperparameters. In addition, the second-order model is simple and works just as well as the Kriging model and the heterogeneous GP model in this situation. The importance of tuning the DE algorithm is demonstrated and a simple strategy on determining optimal hyperparameter settings for constructing UPDs with different target sizes is provided.

While the primary goal of optimizing hyperparameters is to find an optimal hyperparameter combination that maximizes the overall performance of a learning algorithm, the paper additionally examines the impact of different design configurations on the effectiveness of hyperparameter tuning. The insights gained are subsequently used to select the best hyperparameters, which are then applied by the DE algorithm to construct UPDs.

Appendix: Additional tables and figures for target sizes 50×5 and 70×7

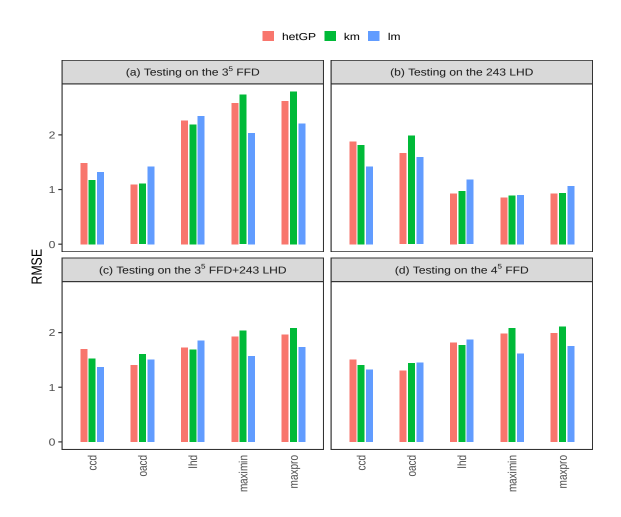


Figure 2.9: Comparison of RMSE with target size 50×5

Table 2.3: Comparison of designs and model evaluations with target size 50×5

	(a) Testing on the 3 ⁵ FFD						(b) Testing on the 243 LHD						
	С	orrela	tion	RMSE			С	orrela	tion	RMSE			
Design	lm	km	hetGP	lm	km	hetGP	lm	km	hetGP	lm	km	hetGP	
ccd3_43	0.94	0.96	0.93	1.32	1.17	1.49	0.83	0.82	0.80	1.42	1.81	1.88	
oacd3_50	0.94	0.96	0.96	1.42	1.11	1.09	0.83	0.84	0.86	1.59	1.98	1.66	
lhd_50	0.83	0.85	0.83	2.35	2.19	2.26	0.89	0.92	0.93	1.18	0.97	0.92	
maximin_50	0.87	0.84	0.81	2.03	2.73	2.58	0.92	0.92	0.93	0.90	0.89	0.85	
maxpro_50	0.85	0.81	0.82	2.21	2.79	2.62	0.90	0.92	0.92	1.06	0.94	0.93	
	(c) 7	(c) Testing on the 3 ⁵ FFD+243 LHD						(d) Testing on the 4 ⁵ FFD					
	correlation			RMSE			correlation			RMSE			
Design	lm	km	hetGP	lm	km	hetGP	lm	km	hetGP	lm	km	hetGP	
ccd3_43	0.92	0.92	0.89	1.37	1.53	1.70	0.93	0.93	0.92	1.32	1.40	1.50	
oacd3_50	0.91	0.92	0.93	1.51	1.61	1.41	0.92	0.94	0.94	1.45	1.44	1.30	
lhd_50	0.86	0.88	0.88	1.86	1.69	1.73	0.87	0.87	0.87	1.87	1.77	1.82	
maximin_50	0.90	0.85	0.85	1.57	2.03	1.92	0.90	0.87	0.86	1.61	2.08	1.98	
maxpro_50	0.88	0.83	0.85	1.73	2.08	1.97	0.88	0.85	0.86	1.75	2.11	1.99	

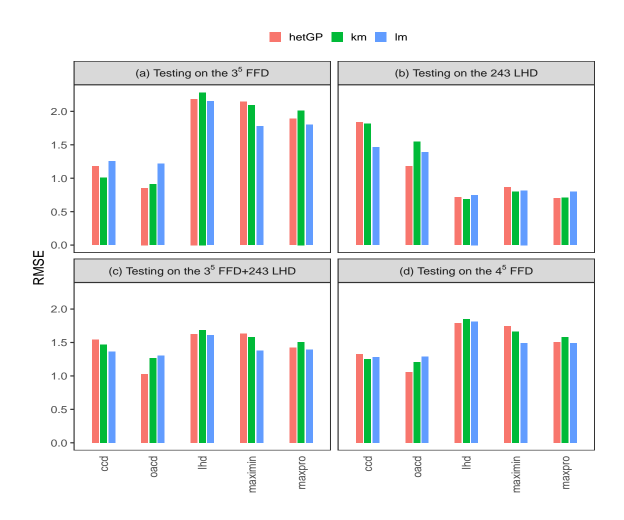


Figure 2.10: Comparison of RMSE with target size 70×7

Table 2.4: Comparison of designs and model evaluations with target size 70×7

	(a) Testing on the 3 ⁵ FFD						(b) Testing on the 243 LHD						
	C	orrela	tion	RMSE			c	orrela	tion	RMSE			
Design	lm	km	hetGP	lm	km	hetGP	lm	km	hetGP	lm	km	hetGP	
ccd3_43	0.93	0.96	0.94	1.25	1.01	1.18	0.81	0.84	0.83	1.46	1.82	1.84	
oacd3_50	0.94	0.97	0.97	1.22	0.91	0.86	0.83	0.88	0.92	1.39	1.54	1.18	
lhd_50	0.83	0.84	0.82	2.15	2.28	2.18	0.94	0.95	0.95	0.75	0.69	0.72	
maximin_50	0.87	0.87	0.81	1.77	2.09	2.14	0.94	0.94	0.93	0.82	0.80	0.86	
maxpro_50	0.88	0.88	0.87	1.80	2.01	1.89	0.94	0.95	0.95	0.80	0.71	0.70	
	(c) T	(c) Testing on the 3 ⁵ FFD+243 LHD						(d) Testing on the 4 ⁵ FFD					
	correlation			RMSE			correlation			RMSE			
Design	lm	km	hetGP	lm	km	hetGP	lm	km	hetGP	lm	km	hetGP	
ccd3_43	0.90	0.91	0.90	1.36	1.47	1.54	0.92	0.94	0.93	1.28	1.26	1.33	
oacd3_50	0.91	0.93	0.95	1.30	1.27	1.03	0.92	0.94	0.95	1.29	1.21	1.06	
lhd_50	0.87	0.87	0.87	1.61	1.69	1.62	0.86	0.87	0.86	1.81	1.85	1.79	
maximin_50	0.90	0.88	0.86	1.38	1.58	1.63	0.89	0.89	0.86	1.49	1.67	1.75	
maxpro_50	0.90	0.89	0.90	1.39	1.51	1.43	0.90	0.90	0.90	1.49	1.58	1.51	

CHAPTER 3

Evaluating Space-Filling Designs for Prediction and Sequential Optimization

3.1 Introduction

In optimization and experimental design, selecting the right strategy is key to achieving precise and efficient results, particularly in methods like Gaussian processes (GP) and Efficient Global Optimization (EGO). These techniques rely heavily on initial design strategies to guide optimization and improve predictive accuracy. Although the Latin hypercube design (LHD) (McKay, Beckman, and Conover 1979) is a commonly used space-filling strategy for computer experiments, research suggests that alternative designs may outperform it in certain cases, especially for screening and prediction tasks (Welch et al. 1992). Studies have explored various design strategies, finding that even subtle differences in design can significantly affect performance (Chen et al. 2016).

The choice of a design is particularly important in complex systems, including fields like engineering and machine learning. Prior research has highlighted how design strategies can impact the performance of Gaussian processes, emphasizing the importance of selecting appropriate methods for high-dimensional problems (Harari and Steinberg 2014).

In a sequential optimization such as EGO, as optimization progresses, the GP model is updated with additional sample points. As the process converges, the influence of the initial design strategy is reduced. Nevertheless, the initial design strategy remains crucial in determining the speed and accuracy of convergence. Choosing the appropriate design

strategy for a given problem is essential for enhancing both predictive performance and optimization efficiency.

The objective of this paper is to evaluate which space-filling designs offer greater efficiency and robustness for both prediction and optimization. Various types of space-filling designs have been proposed in literature. They include LHDs, maximin distance designs (Johnson, Moore, and Ylvisaker 1990; Morris and Mitchell 1995), orthogonal array-based designs (Tang 1993; Xiao and Xu 2018), uniform designs (Fang et al. 2000), maximum projection designs (Joseph, Gul, and Ba 2015), and uniform projection designs (Sun, Wang, and Xu 2019). Traditionally, comparisons have been focused on LHDs and maximin distance LHDs. With new software developments, we can now compare the performance of other types of space-filling designs, including maximum projection designs, uniform designs, and uniform projection designs.

Recent studies, such as those by Shi, Chiu, and Xu (2023), have evaluated space-filling designs in deep neural network predictions, but our research extends this by focusing on the performance of space filling designs in optimization as well. We assess both predictive and minimization capabilities using deterministic test functions, with GP models serving as surrogates and the expected improvement method guiding optimization.

This study evaluates the effectiveness of various space-filling design strategies, particularly in high-dimensional prediction and optimization tasks. We find that uniform designs and uniform projection designs, especially those with 16 levels or more, demonstrate consistent robustness and they often outperform other types of space-filling designs. In contrast, distance-based designs like maximin distance designs underperform in higher dimensions. These findings emphasize the importance of selecting effective design strategies for high-dimensional prediction and optimization.

The paper is organized as follows. Section 2 provides background on the GP surrogate model and active learning techniques. Section 3 reviews various types of space-filling designs, followed by Section 4, which describes the test functions used. Section 5 presents experi-

mental results, and Section 6 concludes with key findings and recommendations for future optimization challenges.

3.2 Background

3.2.1 The surrogate model

Gaussian Processes (GPs) are well-regarded for their flexibility and accuracy in regression tasks, providing a probabilistic framework that accounts for uncertainties in predictions (Snelson 2008). The kriging model, proposed by South African geostatistician Krige (1951), is taken to be one of the surrogate models used in modelling the data. Kriging is one of the methods used to interpolate intermediate values, whereby these intermediate values are modeled using GP which is governed by prior co-variances. It provides a probabilistic prediction of the output variable, as well as an estimate of the uncertainty of the prediction (Chevalier, Picheny, and Ginsbourger 2014). The kriging predictors interpolating the observations are assumed to be noise-free (Roustant, Ginsbourger, and Deville 2012). Intermediate interpolated values obtained by kriging are the best linear unbiased predictors.

The kriging model is

$$Y(\boldsymbol{x}) = \mu(\boldsymbol{x}) + Z(\boldsymbol{x}),$$

where $\mu(\mathbf{x})$ is a trend function and $Z(\mathbf{x})$ is a stationary GP with zero mean. The ordinary kriging assumes that the trend is a constant, i.e., $\mu(\mathbf{x}) = \mu$, while the universal kriging assume that the trend is a linear combination of some basis functions.

The stationary GP $Z(\mathbf{x})$ is determined by its covariance structure $Cov(\mathbf{x}_i, \mathbf{x}_j) = \sigma^2 r(\mathbf{x}_i, \mathbf{x}_j)$ where σ^2 is the common variance and $r(\mathbf{x}_i, \mathbf{x}_j)$ is the correlation between \mathbf{x}_i and \mathbf{x}_j . It is often assumed that $r(\mathbf{x}_i, \mathbf{x}_j)$ is a decreasing function of some distance. Various correlation functions can be used in the fitting of the kriging model. Here, we make use of the Matérn 5/2 correlation function. We fit an ordinary kriging with the constant trend. Chen et al. (2016) showed that a regression model more complex than a constant mean either has lit-

tle impact on prediction accuracy or is an impediment and that the choice of correlation function has modest effect, but there is little to separate two common choices, the power exponential and the Matérn, if the latter is optimized with respect to its smoothness.

Given n design points $\{x_1, \ldots, x_n\}$ and the response $\mathbf{y} = (y_1, \ldots, y_n)$, let $\mathbf{R} = (r(\mathbf{x}_i, \mathbf{x}_j))$ be the $n \times n$ correlation matrix. In this setup, for the ordinary kriging, the μ and σ^2 are estimated as follows:

$$\hat{\mu} = \frac{\mathbf{1}^{\top} \mathbf{R}^{-1} \mathbf{y}}{\mathbf{1}^{\top} \mathbf{R}^{-1} \mathbf{1}}, \qquad \hat{\sigma}^2 = \frac{1}{n} (\mathbf{y} - \hat{\mu} \mathbf{1})^{\top} \mathbf{R}^{-1} (\mathbf{y} - \hat{\mu} \mathbf{1})$$
(3.1)

For any point \boldsymbol{x} , let $\boldsymbol{r}(\boldsymbol{x}) = (r(\boldsymbol{x}, \boldsymbol{x}_1), \dots, r(\boldsymbol{x}, \boldsymbol{x}_n))^{\top}$. Then the best linear unbiased predictor (BLUP) of $Y(\boldsymbol{x})$ is

$$\hat{y}(\boldsymbol{x}) = \hat{\mu} + \boldsymbol{r}(\boldsymbol{x})^{\top} \boldsymbol{R}^{-1} (\boldsymbol{y} - \hat{\mu} \boldsymbol{1}), \tag{3.2}$$

where $\hat{\mu}$ is given in (3.1). The variance of the BLUP is

$$s^{2}(\boldsymbol{x}) = \sigma^{2} \left[1 - \boldsymbol{r}(\boldsymbol{x})^{\top} \boldsymbol{R}^{-1} \boldsymbol{r}(\boldsymbol{x}) + \frac{(1 - \mathbf{1}^{\top} \boldsymbol{R}^{-1} \boldsymbol{r}(\boldsymbol{x}))^{2}}{\mathbf{1}^{\top} \boldsymbol{R}^{-1} \mathbf{1}} \right]$$
(3.3)

and σ^2 can be estimated from the data as given in (3.1).

3.2.2 The expected improvement (EI)

The expected improvement is a measure of how promising a particular set of inputs is in terms of improving the objective function value. To quantify this measure, the objective function needs to be evaluated at various points, yet these evaluations are quite costly. Often when the evaluation of the objective function is costly, the need of specific strategies to optimize these functions arises. In most of these evaluations, the non-availability of derivatives prevents the use of gradient based techniques. Similarly, the use of meta-heuristics (e.g., genetic algorithm) is compromised due to severely limited evaluation budgets (Roustant, Ginsbourger, and Deville 2012). In order to curb these limitations, Jones, Schonlau, and Welch (1998) proposed the use of kriging model as the surrogate model to estimate the

expected improvement of selecting a new set of inputs over the current best solution. When carrying out minimization for example, the improvement at a new point x is

$$I(\boldsymbol{x}) = \max(y_{min} - Y(\boldsymbol{x}), 0),$$

where $y_{min} = \min(y_1, \dots, y_n)$ is the existing minimum value. The new point will bring a positive improvement if $Y(\boldsymbol{x})$ is less than y_{min} , and an improvement of 0 otherwise. The expected improvement (EI) is simply the expectation of $I(\boldsymbol{x})$, that is,

$$E[I(\boldsymbol{x})] = E[\max(y_{min} - Y(\boldsymbol{x}), 0)].$$

Under the ordinary kriging, $Y(\mathbf{x})$ follows a normal distribution with mean $\hat{y}(\mathbf{x})$ and variace $s^2(\mathbf{x})$ given in (3.2) and (3.3), respectively. Then one can express EI in a closed form (Jones, Schonlau, and Welch 1998):

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

where Φ and ϕ are the cumulative and probability density function of the standard normal distribution, respectively. The EI criterion has important properties for sequential exploration: It is null at the already visited sites, and non-negative everywhere else with a magnitude that is increasing with s(x) and decreasing with $\hat{y}(x)$ (Jones, Schonlau, and Welch 1998). This guides the selection of the next set of inputs to evaluate. It encourages the exploration of regions with high uncertainty (large predicted variances) and exploitation of regions with potentially high rewards (small predicted means). The mean function and variance function of the Gaussian process are typically smooth as they are constructed based on a combination of smooth kernel functions and observed data. The cumulative distribution function $\Phi(z)$ and the probability density function $\phi(z)$ of the standard normal distribution, are also smooth functions. These functions are well-defined and infinitely differentiable for all real values of z. Therefore, combining these smooth components in the closed form EI expression results in a smooth function overall. This smoothness property of the closed form EI enables the use of gradient-based optimization methods, in particular, the L-BFGS (Lim-

ited memory Broyden-Fletcher-Goldfarb-Shanno) optimization algorithm, to search for the point of maximum expected improvement efficiently within a specified constrained domain.

3.2.3 The Efficient Global Optimization (EGO) Algorithm

This optimization method sequentially builds upon the EI criterion. EGO enhances this modeling process by intelligently selecting the most informative data points for evaluation. Rather than passively using a fixed dataset, EGO iteratively queries the objective function, focusing on areas that maximize information gain. This targeted approach reduces the number of expensive evaluations needed, accelerating the learning process. EGO (Jones, Schonlau, and Welch 1998) leverages the surrogate GP model to find the global optimum of the objective function. Using EI as the acquisition function, EGO balances the trade-off between exploring new regions of the design space and exploiting known promising areas. This results in a more efficient optimization process that converges on optimal solutions with fewer evaluations.

Algorithm 1 summarizes the procedure. Starting with an initial design X (typically, a Latin hypercube design), EGO sequentially visits a current global maximizer of EI and updates the metamodel at each iteration, including hyperparameters re-estimation (Roustant, Ginsbourger, and Deville 2012). This is done until convergence or until the budgets are exhausted.

Algorithm 1 EGO algorithm

```
Require: X, f = function to be minimized, n_{new}=number of points to add Evaluate f at the design points X; y = f(X) Build a kriging model based on X and y for i in 1 to n_{new} do

Find x^* \leftarrow \arg \max_x \mathrm{E}[I(x)]

Evaluate y^* \leftarrow f(x^*)

Update X and y with the new point x^* and response y^*

Update the kriging model end for Return X, y
```

The EGO algorithm has been shown to be effective and has been adopted in many computer experiments and are nowadays considered as reference global optimization methods in dimension $m \leq 10$ in cases where the number of objective function evaluations is drastically limited (Jones 2001).

3.3 Space-Filling Designs

We briefly review various types of space-filling designs.

Latin Hypercube Design (LHD): Based on McKay (1992)'s Latin hypercube sampling, it divides the range of each factor into bins of equal size, where n also corresponds to the number of samples to be generated resulting in a total of n^m combinations where m is the number of factors being considered. The n samples are then randomly generated such that for all one-dimensional projections, there will be only one sample in each bin. In this paper, the random Latin hypercube sampling was used to generate the random LHD with levels $0, \dots, n-1$. In R, this was accomplished using the 'lhs' package. The following command generates an LHD with n runs and m factors.

> floor(lhs::randomLHS(n, m) * n)

Maximin Distance Designs: Introduced by Johnson, Moore, and Ylvisaker (1990), this design aims at spreading out the design points in the design space by maximizing the minimum distance between any two design points. It thus tends to place a large proportion of points at the corners and on the boundaries of the design space. Mathematically, this can be formulated as follows. Suppose we want to construct an n-run design in m factors, say $D = \{x_1 \dots, x_n\}$, where x_i represents the ith run. The maximin distance design optimizes the function below:

$$\max_{D} \min_{i < j} d\left(\boldsymbol{x}_{i}, \boldsymbol{x}_{j}\right), \tag{3.4}$$

where $d(\mathbf{x}_i, \mathbf{x}_j)$ is the distance between the points \mathbf{x}_i and \mathbf{x}_j . Here we use the Euclidean distance, i.e., $d(\mathbf{x}_i, \mathbf{x}_j) = \sqrt{\sum_{l=1}^{m} (x_{il} - x_{jl})^2}$.

Maximin LHD: Introduced by Morris and Mitchell (1995) these designs combine the principles of Latin Hypercube Sampling (LHS) and the maximin distance criterion to optimize the spread of sample points across the design space. This hybrid approach ensures that points are uniformly distributed in each dimension, while also maximizing the minimum distance between any two points in the design. For computational purpose, Morris and Mitchell (1995) also reformatted the maximin criterion as a minimization problem given as:

$$\min_{D} \phi_{p}(D) = \left\{ \sum_{i=1}^{n-1} \sum_{j=i+1}^{n} \frac{1}{d^{p}(\boldsymbol{x}_{i}, \boldsymbol{x}_{j})} \right\}^{1/p}$$
(3.5)

where p > 0 is chosen large enough so that the resulting design achieves maximin distance. We use the SLHD package to generate maximin LHDs because they are better than those generated from the lhs package using the function maximinLHS. The following command generates a maximin LHD with n runs and m factors.

> SLHD::maximinSLHD(t = 1, n, m)\$Design - 1

Maximum Projection Design (Maxpro Design): Although maximin LHDs ensure good space-filling in m dimensions and uniform projections in each dimension, their projection properties in two to m-1 dimensions are not known (Joseph, Gul, and Ba 2015).

By the effect sparsity principle (Wu and Hamada 2011), only a few factors are expected to be important. To curb this, Joseph, Gul, and Ba (2015) proposed the maximum projection (MaxPro) criterion:

$$\min_{D} \psi(D) = \left\{ \frac{2}{n(n-1)} \sum_{i=1}^{n-1} \sum_{j=i+1}^{n} \frac{1}{\prod_{l=1}^{m} (x_{il} - x_{jl})^{2}} \right\}^{1/m}.$$
 (3.6)

They argued that the design that minimizes $\psi(D)$ tends to maximize its projection capability in all sub spaces of factors, and thus named these designs as maximum projection designs. We use the MaxPro package to generate a MaxPro LHD with n runs and m factors.

> MaxPro::MaxProLHD(n, m)\$Design * n - 0.5

Uniform Design (UD): These designs seek to scatter points uniformly on the domain. This is often achieved by minimizing the centered L_2 -discrepancy of the design (Fang et al. 2000). For an $n \times m$ design $D = (x_{ik})$ with s levels, denoted by $0, \dots, s-1$, its (squared) centered L_2 -discrepancy is defined as

$$CD(D) = \frac{1}{n^2} \sum_{i=1}^{n} \sum_{j=1}^{n} \prod_{k=1}^{m} \left(1 + \frac{1}{2} |z_{ik}| + \frac{1}{2} |z_{jk}| - \frac{1}{2} |z_{ik} - z_{jk}| \right) - \frac{2}{n} \sum_{i=1}^{n} \prod_{k=1}^{m} \left(1 + \frac{1}{2} |z_{ik}| - \frac{1}{2} |z_{ik}|^2 \right) + \left(\frac{13}{12} \right)^m,$$
(3.7)

where $z_{ik} = (2x_{ik} - s + 1)/(2s)$. We use the R package UniDOE to generate UDs. The following command generates a UD with n runs, m factors, and s levels.

> UniDOE::GenUD(n, m, s)\$final_design - 1

The UniDOE package uses the threshold accepting algorithm to generate UDs, which often takes an excessive time in comparison with the construction of maximin and MaxPro LHDs. Thus we only used UDs for prediction, not optimization.

The UniDOE package was removed from the CRAN (Comprehensive R Archive Network) repository in 2021 as it was no longer maintained and problems were not corrected on time. Though one can still be able to install it from github.

Uniform Projection Design (UPD): Proposed by Sun, Wang, and Xu (2019), this design solely focuses on two-dimensional projections. This is due to two factor interactions being more important than three-factor or higher-order interactions. The motivating idea is that although designs with low discrepancy have good uniformity in the full-dimensional space, they can have bad projections in lower dimensional spaces, which is undesirable when only a few factors are active. Thus it is preferable to have a design with better projection properties. The uniform projection criterion is defined using the centered L_2 -discrepancy as follows:

$$\phi(D) = \frac{2}{m(m-1)} \sum_{|u|=2} \text{CD}(D_u), \qquad (3.8)$$

where u is a subset of $\{1, 2, ..., m\}$, |u| denotes the cardinality of u and D_u is the projected design of D onto dimensions indexed by the elements of u. The $\phi(D)$ is the average centered L_2 -discrepancy values of all two-dimensional projections of D. The UPD scatters points uniformly in all dimensions and have good space-filling properties in terms of distance, uniformity and orthogonality (Sun, Wang, and Xu 2019). We use the UniPro package and the Differential Evolution algorithm to generate UPDs. The following command generates a UPD with n runs, m factors, and s levels.

Note that the SLHD and MaxPro packages can only generate LHDs while the UniDOE and UniPro packages can generate balanced multi-level UDs and UPDs as long as the number of runs is a multiple of the number of levels.

3.4 Test Functions

To test the efficiency of various space-filling designs, two methods were analyzed. First the efficiency of the designs was analysed via its predictability efficiency. Here the objective was to minimize the prediction root mean square error (RMSE) to ensure accurate and reliable

predictions. The second method was to look at designs through the problem of minimization convergence. Under these two methods, various test functions were taken into consideration.

3.4.1 Prediction Functions

Currin: This is a simple two dimensional polynomial function evaluated on the square $[0, 1]^2$ used for illustrating methods of modeling computer experiment output (Currin et al. 1991). It has the form:

$$f(\mathbf{x}) = 4.90 + 21.15x_1 - 2.17x_2 - 15.88x_1^2 - 1.38x_2^2 - 5.26x_1x_2.$$

Circuit: This is a six-dimensional function that models an output transformerless pushpull circuit (Surjanovic and Bingham 2013) with the response being the midpoint voltage. It takes the following form:

$$f(\boldsymbol{x}) = \frac{(V_{b1} + 0.74) \beta (R_{c2} + 9)}{\beta (R_{c2} + 9) + R_f} + \frac{11.35 R_f}{\beta (R_{c2} + 9) + R_f} + \frac{0.74 R_f \beta (R_{c2} + 9)}{(\beta (R_{c2} + 9) + R_f) R_{c1}},$$
where $V_{b1} = \frac{12 R_{b2}}{R_{b1} + R_{b2}}$.

The variables are of different domains: $R_{b1} \in [50, 150]$, $R_{b2} \in [25, 70]$, $R_f \in [0.5, 3]$, $R_{c1} \in [1.2, 2.5]$, $R_{c2} \in [0.25, 1.2]$ and $\beta \in [50, 300]$.

Piston: This is a seven-dimensional function that models the circular motion of a piston within a cylinder. It involves a chain of nonlinear functions with the response f(x) being the cycle time (Surjanovic and Bingham 2013). It takes the following form:

$$f(\mathbf{x}) = 2\pi \sqrt{\frac{M}{k + S^2 \frac{P_0 V_0}{T_0} \frac{T_a}{V^2}}},$$

$$V = \frac{S}{2k} \left(\sqrt{A^2 + 4k \frac{P_0 V_0}{T_0} T_a} - A \right) \text{ and } A = P_0 S + 19.62M - \frac{kV_0}{S},$$

where $M \in [30, 60]$ is piston weight (kg), $S \in [0.005, 0.020]$ is piston surface area (m²), $V_0 \in [0.002, 0.010]$ is initial gas volume (m³), $k \in [1000, 5000]$ is spring coefficient (N/m), $P_0 \in [90000, 110000]$ is atmospheric pressure (N/m²), $T_a \in [290, 296]$ is ambient temperature (K) and $T_0 \in [340, 360]$ is filling gas temperature (K).

Borehole: This is a 8-dimensional function that models water flow through a borehole with the response being water rate flow in m^3/yr (Surjanovic and Bingham 2013). It takes the following form:

$$f(\boldsymbol{x}) = \frac{2\pi T_u \left(H_u - H_l \right)}{\ln \left(r/r_w \right) \left(1 + \frac{2LT_u}{\ln \left(r/r_w \right) r_w^2 K_w} + \frac{T_u}{T_l} \right)}.$$

The variable names and their corresponding domains are given as follows:

Domain	Variable Name
$r_w \in [0.05, 0.15]$	radius of borehole (m)
$r \in [100, 50000]$	radius of influence (m)
$T_u \in [63070, 115600]$	transmissivity of upper aquifer (m ² /yr)
$H_u \in [990, 1110]$	potentiometric head of upper aquifer (m)
$T_{I} \in [63.1, 116]$	transmissivity of lower aquifer (m ² /yr)
$H_{\rm I} \in [700, 820]$	potentiometric head of lower aquifer (m)
$L \in [1120, 1680]$	length of borehole (m)
$K_{\rm w} \in [9855, 12045]$	hydraulic conductivity of borehole (m/yr)

G-function: This is an *m*-dimensional function designed by Ilya M. Sobol to study the sensitivity of complex mathematical models and is particularly useful for benchmarking and validating global sensitivity analysis methods. It has the form:

$$f(\mathbf{x}) = \prod_{i=1}^{m} \frac{|4x_i - 2| + a_i}{1 + a_i},$$

where $\mathbf{x} = (x_1, x_2, \dots, x_m)$ is a vector of input variables, each typically in the range [0, 1] and $a_i = i/2-1$, for all $i = 1, \dots, m$, are coefficients that determine the sensitivity of the function to the *i*th input variable. Larger values of a_i indicate less sensitivity to the corresponding input.

Wing weight: This is a function that models a light aircraft wing with the response being the wing's weight.

$$f(\boldsymbol{x}) = 0.036 S_w^{0.758} W_{fw}^{0.0035} \left(\frac{A}{\cos^2(\Lambda)}\right)^{0.6} q^{0.006} \lambda^{0.04} \left(\frac{100t_c}{\cos(\Lambda)}\right)^{-0.3} (N_z W_{dg})^{0.49} + S_w W_p,$$

where $S_W \in [150, 200]$ is wing area (ft²), $W_{fw} \in [220, 300]$ is weight of fuel in the wing (lb), $A \in [6, 10]$ is aspect ratio, $\Lambda \in [-10, 10]$ is quarter-chord sweep (degrees), $q \in [16, 45]$ is dynamic pressure at cruise (lb/ft²), $\lambda \in [0.5, 1]$ is taper ratio, $t_c \in [0.08, 0.18]$ is aerofoil thickness to chord ratio, $N_Z \in [2.5, 6]$ is ultimate load factor, $W_{dg} \in [1700, 2500]$ is flight design gross weight (lb) and $W_p \in [0.025, 0.08]$ is paint weight (lb/ft²).

Oakley & O'hagan (2004): This is a 15-dimensional function that take the following form:

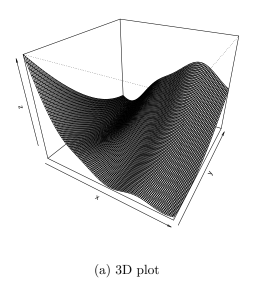
$$f(\boldsymbol{x}) = \mathbf{a}_1^T \boldsymbol{x} + \mathbf{a}_2^T \sin(\boldsymbol{x}) + \mathbf{a}_3^T \cos(\boldsymbol{x}) + \boldsymbol{x}^T \mathbf{M} \boldsymbol{x},$$

where the **a**-coefficients are chosen so that 5 of the input variables contribute significantly to the output variance, 5 have a much smaller effect, and the remaining 5 have almost no effect on the output variance. Values of the coefficient vectors $\mathbf{a}_1, \mathbf{a}_2$ and \mathbf{a}_3 , and the matrix \mathbf{M} , are available at Surjanovic and Bingham (2013).

Rosenbrock: The Rosenbrock function is a classic *m*-dimensional optimization problem, also known as the Valley or banana function. The global optimum lays inside a long, narrow, parabolic shaped flat valley (Molga and Smutnicki 2005). To find the valley is trivial, however convergence to the global optimum is difficult and hence this problem has been frequently used to test the performance of optimization algorithms (Picheny, Wagner, and Ginsbourger 2013). For comparison, the modified 4d version presented by Picheny, Wagner, and Ginsbourger (2013) was used. This modified version has the following form:

$$f(\mathbf{x}) = \frac{1}{3.755 \times 10^5} \left[\sum_{i=1}^{3} \left(100 \left(\bar{x}_{i+1} - \bar{x}_i^2 \right)^2 + (1 - \bar{x}_i)^2 \right) - 3.827 \times 10^5 \right], \quad (3.9)$$

where $\bar{x}_i = 15x_i - 5$ for all i = 1, ..., 4. The test region is usually restricted to $0 \le x_i \le 1$ for i = 1, ..., 4.



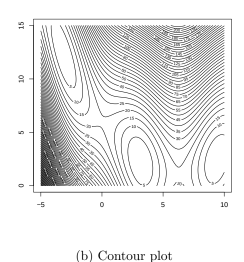


Figure 3.1: Branin Function

3.4.2 Minimization Functions

Branin function: This is a well-known 2-dimensional function with multiple local minima and global minimum introduced by Dixon and Szegö (1978). It is often used to test optimization algorithms due to its complexity and multi-modal nature. The Branin function has the form

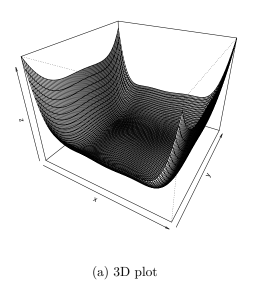
$$f(\mathbf{x}) = f(x_1, x_2) = a(x_2 - bx_1^2 + cx_1 - r)^2 + s(1 - t)\cos(x_1) + s$$

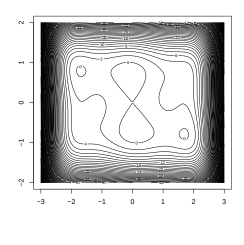
where the typical parameter values are $a=1,b=5.1/(4\pi^2)$, $c=5/\pi,r=6,s=10$ and $t=1/(8\pi)$. The function is usually evaluated over the square $x_1 \in [-5,10], x_2 \in [0,15]$. Within this domain the function has 3 global minima: $\boldsymbol{x}^*=(-\pi,12.275), (\pi,2.275)$ and (9.42478,2.475), with the minimum function value being $f(\boldsymbol{x}^*)=0.397887$. The contour plot of the function is as shown in Figure 3.1.

Camel Six-Hump function: This is a 2-dimensional function with 6 local minima and 2 global minima evaluated on the rectangle $x_1 \in [-3, 3]$ and $x_2 \in [-2, 2]$. It has the form:

$$f(\mathbf{x}) = \left(4 - 2.1x_1^2 + \frac{x_1^4}{3}\right)x_1^2 + x_1x_2 + \left(-4 + 4x_2^2\right)x_2^2.$$

The global minima is $f(x^*) = -1.0316$ which occurs at $x^* = (0.0898, -0.7126)$ and (-0.0898, 0.7126).





(b) Contour plot

Figure 3.2: Camel Six-Hump Function

Goldstein-Price function: This is a 2-dimensional function with several local minima evaluated on the rectangle $x_i \in [-2, 2]$ for i = 1, 2. It has the form:

$$f(\mathbf{x}) = \left[1 + (x_1 + x_2 + 1)^2 \left(19 - 14x_1 + 3x_1^2 - 14x_2 + 6x_1x_2 + 3x_2^2\right)\right] \times \left[30 + (2x_1 - 3x_2)^2 \left(18 - 32x_1 + 12x_1^2 + 48x_2 - 36x_1x_2 + 27x_2^2\right)\right].$$
(3.10)

The global minima is $f(\boldsymbol{x}) = 3$ at $\boldsymbol{x} = (0, -1)^{\top}$.

Ackley function: The Ackley function is a widely used function for testing optimization algorithms (Adorio and Diliman 2005; Molga and Smutnicki 2005). In its two-dimensional form, it is characterized by a nearly flat outer region, and a large hole at the center. The function poses a risk for optimization algorithms, particularly hill climbing algorithms, as they often get trapped in one of its many local minima (Surjanovic and Bingham 2013). The Ackley function has the following form:

$$f(\boldsymbol{x}) = -a \exp\left(-b\sqrt{\frac{1}{m}\sum_{i=1}^{m}x_i^2}\right) - \exp\left(\frac{1}{m}\sum_{i=1}^{m}\cos\left(cx_i\right)\right) + a + \exp(1)$$

with the recommended values for a, b and c being a = 20, b = 0.2 and $c = 2\pi$. It is usually evaluated on the hypercube $x_i \in [-32.768, 32.768]$ for all $i = 1, \dots, m$ with a global minimum of $f(\mathbf{x}) = 0$ at $x_i = 0$ for $i = 1, \dots, m$.

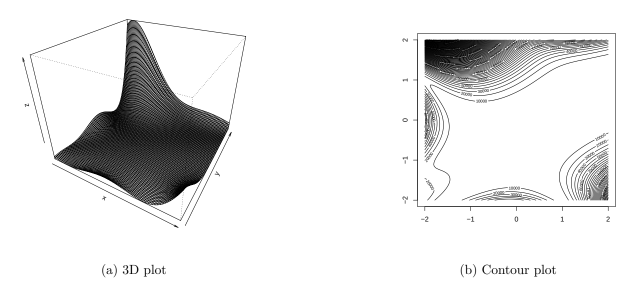


Figure 3.3: Goldstein-Price Function

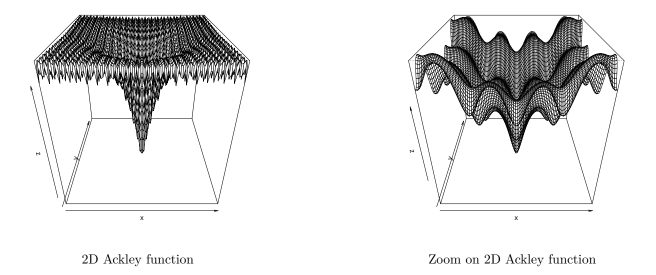
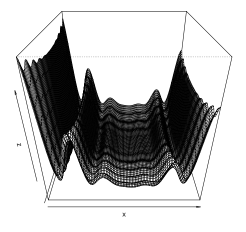


Figure 3.4: Ackley function



(a) 2D levy Function

Figure 3.5: Levy function

Levy function: This is a m-dimensional function often used as a test function for optimization. It is usually evaluated on the hypercube $x_i \in [-10, 10] \ \forall i = 1, ..., m$ and has the global minimum of $f(\boldsymbol{x}^*) = 0$ at $\boldsymbol{x}^* = (1, ..., 1)^\top$. It is defined as

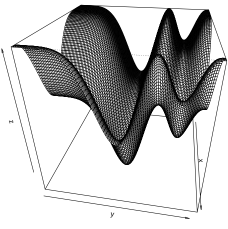
$$f(\boldsymbol{x}) = \sin^2(\pi w_1) + \sum_{i=1}^{m-1} (w_i - 1)^2 \left[1 + 10 \sin^2(\pi w_i + 1) \right] + (w_m - 1)^2 \left[1 + \sin^2(2\pi w_m) \right],$$

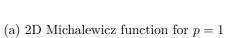
where $w_i = 1 + (x_i - 1)/4$ for i = 1, ..., m.

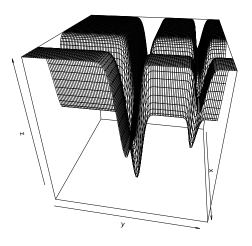
Michalewicz Function: This is a multimodal test function with m! local optima for m dimensions. It is characterized by its steep valleys and ridges. The "steepness" of the valley or edges is defined by the parameter p (Molga and Smutnicki 2005). For larger p it is quite difficult to obtain the optimum as the function values for points in the space outside the narrow peaks give very little to no information on the location of the global optimum. At the same time an increase in dimensionality increases the difficulty due to the increased number of local minima. The function is defined as:

$$f(\boldsymbol{x}) = -\sum_{i=1}^{m} \sin(x_i) \left[\sin\left(\frac{ix_i^2}{\pi}\right) \right]^{2p}$$

with p usually set at p=10 and the domain restricted to $x_i \in [0,\pi]$ for $i=1,\cdots,m$. The







(b) 2D Michalewicz function for p=10

Figure 3.6: Michalewicz function

global minimum is approximated to be $f(\mathbf{x}) = -1.8013$ for m = 2, $f(\mathbf{x}) = -4.6877$ for m = 5 and $f(\mathbf{x}) = -9.6602$ for m = 10.

3.5 Numerical Experiments

This section presents the numerical results from applying Gaussian Process (GP) to the prediction functions and Efficient Global Optimization (EGO) to the minimization functions. In order to determine the effectiveness of the different initial designs, both prediction and optimization functions were taken into consideration.

We first generate different types of space-filling designs using various packages as described in Section 3.3 and examine their performance and relationship under different criteria. For each type of design, we ran the corresponding algorithm to construct 80×8 LHDs 200 times. Table 3.1 shows the mean and median of the various design criteria for each design type. The design criteria are the maximin LHD criterion (3.5) with p = 15, the MaxPro criterion (3.6), the centered L_2 -discrepancy (3.7), the UPD criterion (3.8), and the absolute average correlation criterion (3.11) defined below:

$$\rho_{ave} = \frac{2}{m(m-1)} \sum_{k=1}^{m} \sum_{l=k+1}^{m} \frac{\left| \sum_{i=1}^{n} (x_{ik} - \bar{x}_{\cdot k}) (x_{il} - \bar{x}_{\cdot l}) \right|}{\sqrt{\sum_{i=1}^{n} (x_{ik} - \bar{x}_{\cdot k})^{2} \sum_{i=1}^{n} (x_{il} - \bar{x}_{\cdot l})^{2}}}.$$
 (3.11)

For all criteria, smaller values represent better designs.

As expected, each type of design performs the best under the corresponding criterion used to generate the design. For example, maximin LHD have smaller $\phi_p(D)$ values and MaxPro designs have smaller $\psi(D)$ values. In addition, random LHDs are worse than all other types of space-filling designs. Regarding to the ρ_{ave} criterion, UPDs and uniform designs (ud) are considerably better than other types of designs. Figure 3.7 gives a visualization where UPDs are fairly comparable to the uniform designs. These two types of designs have the highest correlation. From Table 3.1 and Figure 3.7, we are capable to determine that uniform designs and UPDs are more robust than other types of designs under various design criteria. We have examined cases with different design sizes, and the conclusions are similar.

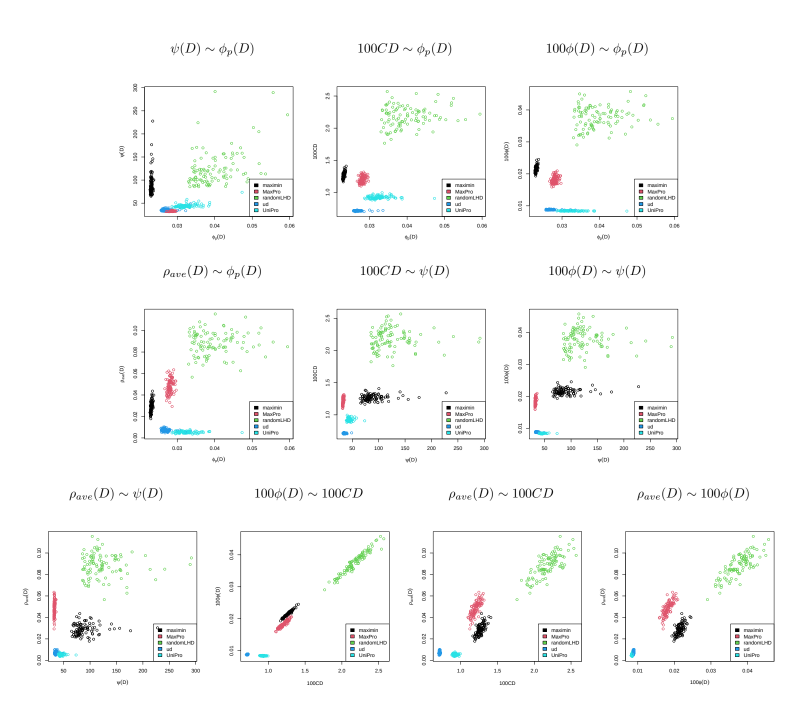


Figure 3.7: Comparison of 100 80×8 LHDs using various criteria

Table 3.1: Comparison of 100 80×8 designs using various criteria

design	summary	$\phi_p(D)$	$\psi(D)$	100 <i>CD</i>	$100\phi(D)$	$ ho_{ave}$
maximin	mean	0.0231	90.5940	1.2676	0.0215	0.0289
	median	0.0231	83.5325	1.2616	0.0214	0.0289
MaxPro	mean	0.0281	32.5730	1.2028	0.0182	0.0483
	median	0.0280	32.3859	1.1962	0.0181	0.0481
ud	mean	0.0269	35.0794	0.7129	0.0088	0.0072
	median	0.0266	34.6913	0.7131	0.0088	0.0071
UniPro	mean	0.0326	44.4252	0.9240	0.0084	0.0054
	median	0.0321	43.6042	0.9215	0.0084	0.0054
randomLHD	mean	0.0402	125.7954	2.1808	0.0377	0.0883
	median	0.0385	116.8441	2.1881	0.0379	0.0891

3.5.1 Prediction results

The test functions are evaluated on various space-filling LHDs of sizes 64×15 and 128×31 . In addition, UPDs with 8 and 16 levels are also constructed and used for model fitting and prediction with the purpose of evaluating whether the number of levels affects the performance. All designs are scaled to the function domain accordingly. Since the functions are of various dimensions, yet the designs used are of a pre-specified dimension, the unused dimensions are considered to be inert and serve as 'noise'. This is quite common in practice whereby some inert factors are involved in an experimental study. For example, the borehole function has 8 variables, thus we have 7 inert variables when the design has 15 columns. For each design type, an ordinary kriging model with a constant trend using the matérn correlation function with $\nu = 5/2$ is then fitted. To evaluate the performance of different types of designs, the normalized root mean square error is used. This is given by

Normalized RMSE =
$$\left[\frac{N^{-1} \sum_{i=1}^{N} \left\{\hat{y}\left(\boldsymbol{x}_{i}\right) - y\left(\boldsymbol{x}_{i}\right)\right\}^{2}}{N^{-1} \sum_{i=1}^{N} \left\{\bar{y}_{train} - y\left(\boldsymbol{x}_{i}\right)\right\}^{2}}\right]^{1/2},$$

where $\mathbf{x}_1, \dots, \mathbf{x}_N$ are the inputs of the test dataset, $y(\mathbf{x}_i)$ is the true response, $\hat{y}(\mathbf{x}_i)$ is the predicted value from the GP model, and \bar{y} is the mean of the responses in the training dataset. This criterion is related to R^2 in regression, but it measures performance for a new test dataset and smaller values are desirable (Chen et al. 2016). We used a random LHD with N = 10,000 runs as the test dataset.

In order to reduce the randomness effect, the process is replicated 200 times. The replication of the design generation is done via row and column permutations. For each design type, except for the random LHD, 100 designs are generated and the best design among these 100 generated designs determined by each design type criterion is chosen as the candidate design. Using row and column permutation of the best design for each design type, 100 designs are thus generated. This method of design generation is chosen as it is almost infeasible to generate uniform designs of size 64×15 and 128×31 200 times. In particular to generate a 10×3 uniform design with 10 levels using a machine equipped with a 13th Gen

Intel CoreTM i7-1360P 16-core CPU running at 2.2 GHz, it takes an average of 5.8 seconds while a machine with Intel® Xeon® Platinum 8160 CPU with 48 cores running at 2.10 GHz takes an average of 5.77 seconds. In order to use the row and column permutation, a fairly large enough design is to be used. For example to generate a 80×8 UD design with 80 levels requires ≈ 439 minutes (7.32 hours) just for one design in the first machine, while needing 432 minute (7.2 hours) in the second machine. This shows as to why permutation is the best way to do the design generation.

Figure 3.8 shows the normalized RMSEs for the various test functions using various 64×15 designs. The mean and median RMSEs are also obtained and are reported in Table 3.2. A general conclusion is that the use of uniform designs and uniform projection designs in prediction is efficient as they result in minimal normalized RMSEs for most of the test functions, compared to the other types of designs. This could be attributed to their robustness in nature. On the other hand, the random LHD is often the worst as expected with one exception. For the 9-dimensional G-function, the maximin, MaxPro and uniform LHDs perform worse than the random LHD whereas all three of the UPDs perform better than the random LHD. In particular the 8-level UPD (upd8q) performs the best in this case.

Among all space-filling designs, the UPD appears to be the most robust as it is never worse than the random LHD. Also it is better than the maximin and MaxPro designs with one exception for the Wingweight function, where the maximin design is better. In addition, the 16-level UPD (upd16q) performs better than the 8-level UPD except for the case of the G-function. At the same time 16-level UPD performs better than the 64-level UPD. This gives the notion of the importance of the number of levels. A fair enough number of levels in comparison to the number of runs is important in prediction. While in physical experiments, 3-5 levels are enough, in computer experiments, more levels are needed.

The conclusions reached are further cemented by observing the results obtained when using 128×31 design size for training and predicting on the test dataset with 10,000 runs. Here 8-level UPD is not used as it produces many outliers which would distort the visual

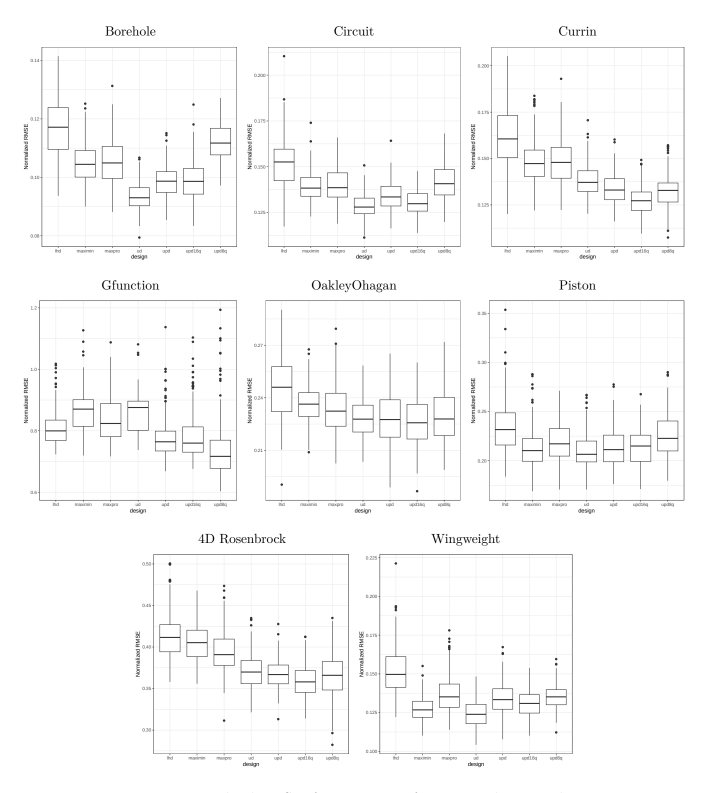


Figure 3.8: Normalized RMSEs for various test functions and 64×15 designs

Table 3.2: Means and medians of normalized RMSEs for 64×15 designs

fun		lhd	maximin	maxpro	ud	upd	upd16q	upd8q
Borehole	mean	0.1168	0.1048	0.1053	0.0936	0.0986	0.0990	0.1119
	median	0.1171	0.1044	0.1049	0.0930	0.0987	0.0986	0.1117
Circuit	mean	0.1519	0.1394	0.1397	0.1286	0.1337	0.1308	0.1417
	median	0.1526	0.1383	0.1386	0.1280	0.1335	0.1298	0.1408
Currin	mean	0.1622	0.1487	0.1480	0.1380	0.1333	0.1273	0.1326
	median	0.1606	0.1473	0.1479	0.1372	0.1330	0.1273	0.1328
Gfunction	mean	0.8092	0.8618	0.8328	0.8540	0.7795	0.7830	0.7423
	median	0.7996	0.8704	0.8237	0.8757	0.7642	0.7598	0.7167
OakleyOhagan	mean	0.2461	0.2358	0.2329	0.2284	0.2277	0.2258	0.2300
	median	0.2460	0.2364	0.2324	0.2278	0.2276	0.2257	0.2279
Piston	mean	0.2351	0.2127	0.2195	0.2101	0.2137	0.2146	0.2254
	median	0.2317	0.2103	0.2172	0.2065	0.2113	0.2149	0.2227
Rosenbrock	mean	0.4134	0.4051	0.3943	0.3708	0.3672	0.3583	0.3655
	median	0.4115	0.4053	0.3908	0.3699	0.3669	0.3580	0.3659
Wingweight	mean	0.1520	0.1275	0.1367	0.1244	0.1337	0.1310	0.1355
	median	0.1496	0.1268	0.1352	0.1239	0.1334	0.1309	0.1352

comparison of the results. The 16-level UPD still has the best performance in most of the test functions; see Table 3.3 and Figure 3.9.

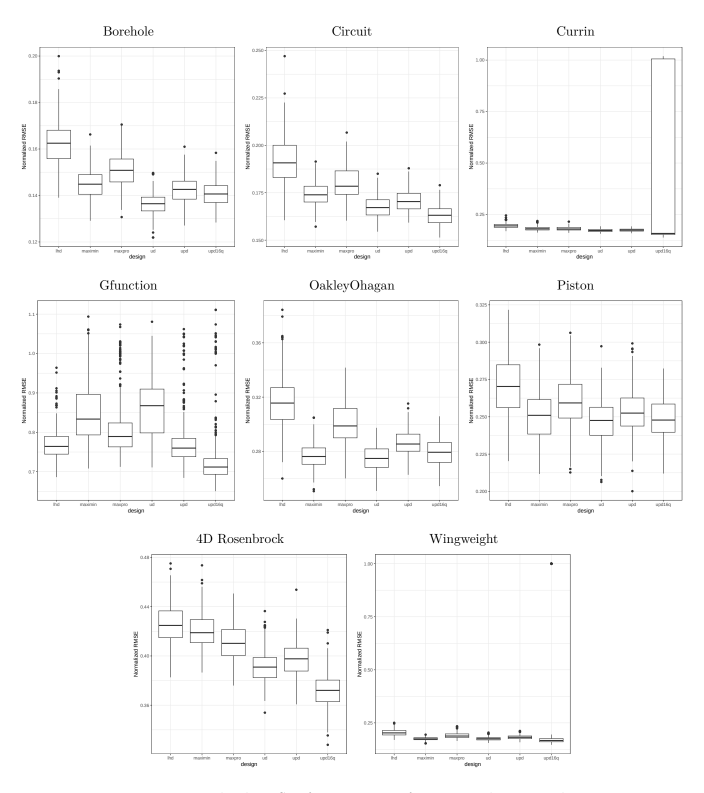


Figure 3.9: Normalized RMSEs for various test functions and 128×31 designs

Table 3.3: Means and medians of normalized RMSEs for 128×31 designs

fun		lhd	maximin	maxpro	ud	upd	upd16q
Borehole	mean	0.1624	0.1451	0.1511	0.1366	0.1425	0.1408
	median	0.1625	0.1449	0.1508	0.1365	0.1426	0.1406
Circuit	mean	0.1920	0.1742	0.1798	0.1676	0.1709	0.1630
	median	0.1908	0.1739	0.1784	0.1672	0.1704	0.1631
Currin	mean	0.1960	0.1818	0.1815	0.1730	0.1747	0.4376
	median	0.1949	0.1809	0.1799	0.1724	0.1744	0.1582
Gfunction	mean	0.7697	0.8550	0.8074	0.8682	0.7724	0.7297
	median	0.7642	0.8336	0.7894	0.8677	0.7599	0.7119
OakleyOhagan	mean	0.3166	0.2769	0.3006	0.2752	0.2865	0.2797
	median	0.3156	0.2763	0.2988	0.2748	0.2854	0.2794
Piston	mean	0.2699	0.2504	0.2601	0.2475	0.2535	0.2483
	median	0.2703	0.2510	0.2593	0.2475	0.2525	0.2478
Rosenbrock	mean	0.4257	0.4204	0.4105	0.3916	0.3975	0.3724
	median	0.4248	0.4189	0.4102	0.3909	0.3977	0.3720
Wingweight	mean	0.2037	0.1748	0.1900	0.1756	0.1821	0.3112
	median	0.2027	0.1745	0.1879	0.1752	0.1819	0.1669

From Figure 3.9 we see that the Currin and the wing weight functions produce a few large RMSE values, likely due to the failure of the optimization routine when fitting the

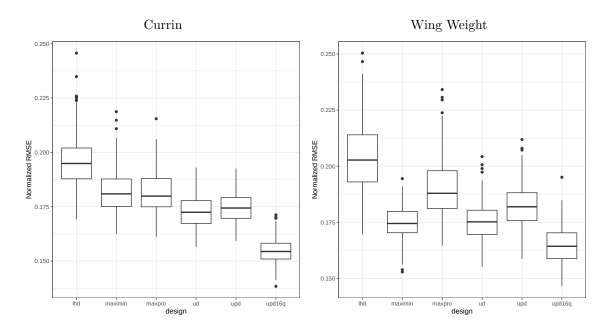


Figure 3.10: Normalized RMSEs for Currin and Wing Weight functions without the outliers

kriging model. These outliers distort the display of the boxplots even though the other RMSE values are generally concentrated below the RMSE value of the other functions. For a fair comparison, we got rid of these outliers and plotted the rest as shown in Figure 3.10.

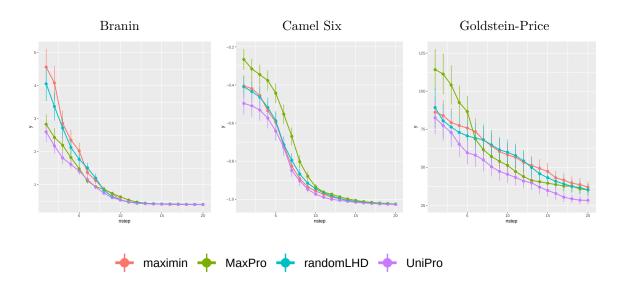


Figure 3.11: Minimization path for 2 dimensional test functions

3.5.2 Minimization results

We consider four types of space-filling designs for sequential minimization: random LHDs, maximin LHDs, MaxPro LHDs and UPDs. The test functions are evaluated at $n_0 = 10 \times m$ initial points, apart from the Branin function which is evaluated at $n_0 = 10$ initial points. These functions are then minimized using the EGO approach. We use the function fastEGO.nsteps in the R DiceOptim package to sequentially add a point at a time with n_1 steps. For each step, the cumulative minimum is recorded. This process is replicated 200 times.

Figure 3.11 shows the average minimal values at each step, together with the 2 standard error bar, of the resulting minimization path with $n_1 = 20$ added points for three 2-dimensional test functions. For the Branin function, MaxPro designs and UPDs yield smaller y-values at early stages than maximin and random LHDs. The difference gradually diminishes over the sequential optimization process. After 15 additional points, all the methods have the same y-value indicating that the function minimum has been achieved. The result is similar for the Camel Six function. For the Goldstein-Price function, the UPD maintains its advantage over others even after 20 steps.

The results presented above are not unique. They are determined by the computation of the first y-value from the initial design. The design that initially produces the least function value tends to generally converge faster to the function minimum, as the sequential points majorly depend on the expected improvement with influence from the design type. This indicates that the choice of the initial design for optimization is quite significant. A poor initial design will yield poor objective values at the initialization stage which will lead to slower convergence.

As the number of iterations (nsteps) increases, the influence of the initial design type on the optimization process diminishes. Initially, the choice of design strategy can play a significant role in guiding the search for the global optimum. However, as more sample points are iteratively added based on the acquisition function, the optimization process becomes increasingly driven by the updated GP model. This model incorporates all collected data points, thereby reducing the relative impact of the initial design. Consequently, the optimization converges towards the global optimum, and differences attributed to the initial design strategy become less pronounced.

Though this is the case, in high dimensional setup, or using complicated functions, the number of sequentially added points needed for convergence might be quite large. With a limited budget, the effect of initial design could be profound.

Figure 3.12 shows the minimization path for 3 other test functions with varying dimensions from 4 to 8. One striking observation is the decreasing efficiency of using maximin designs when the dimension increases. At lower dimensions, maximin designs fairly compete with the other designs but as the dimensions increase to 6 and 8, it is observed that the maximin designs do poorly. This is even after increasing the number of steps to 30 or more. We also note that MaxPro designs are worse than random LHDs and UPDs for the Ackley and Levy function. Overall, UPDs are robust and perform well under all situations, especially in high dimensions and complicated test functions.

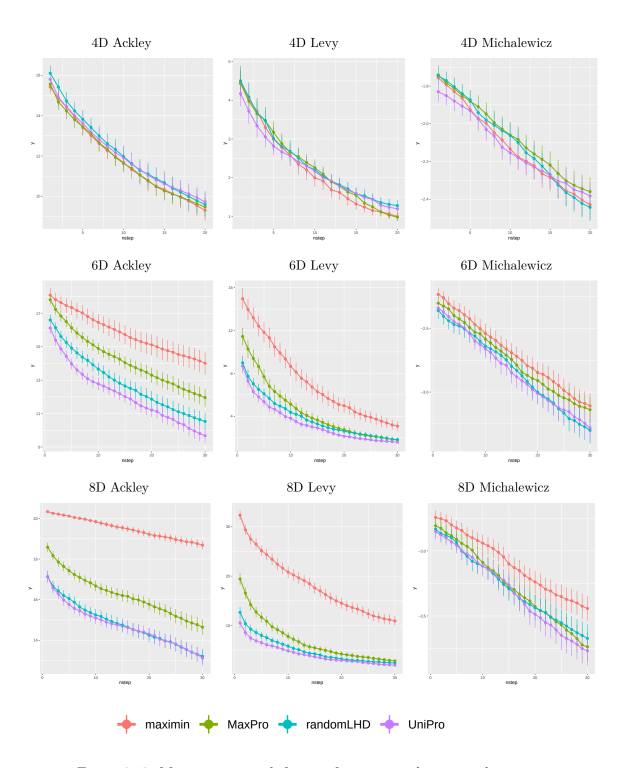


Figure 3.12: Minimization path for test functions with varying dimensions

3.6 Concluding Remarks

In conclusion, the numerical evaluation of Gaussian Process (GP) and Efficient Global Optimization (EGO) techniques reveal that the uniform design (UD) and uniform projection design (UPD) stand out for their robustness and efficiency. When compared across various design criteria, the UD and UPD consistently demonstrate superior performance. The extensive experimentation, including the generation of 100 designs and their permutations, highlighting the efficiency of the uniform designs in reducing normalized root mean square error in prediction tasks. Notably, the UPD with 16 levels proved to be particularly effective, underscoring the importance of having a sufficiently large number of levels in design, yet it is not necessary to use LHDs with the number of levels being equal to the number of runs when the latter is large.

The comparative analysis of different initial designs for EGO further confirmed the efficacy of UPDs. As evidenced by the results, UPDs converged more quickly to the function minimum compared to other design methods. Although the differences in optimization outcomes among design strategies became less pronounced with more iterations, the initial design choice had a significant impact at early stages, which could prolong to later stages when we encounter complicated optimization tasks in high dimensions. This observation underscores the value of selecting a good design strategy for enhancing the efficiency of the optimization process. In conclusion, UPDs are simple to construct, offering a feasible solution for large-scale, high-dimensional prediction and optimization tasks.

However, the results also revealed some limitations, particularly with distance-based designs like the maximin criterion in high-dimensional spaces. One reason as to why this might be the case is because of the use of Euclidean distance as the criterion to be optimized by the maximin design yet it is well documented that Euclidean distance is not a good metric in high dimensions. In high dimensions, the natural intuition which comes from our three-dimensional world fails and thus do not apply (Domingos 2012). Taking an example of Gaussian distribution, in high dimensions, most of the mass of a multivariate Gaussian

distribution is not near the mean, but in an increasingly distant "shell" around it; and most of the volume of a high dimensional orange is in the skin, and not the pulp. If a constant number of points is distributed uniformly in a high-dimensional hypercube, beyond some dimensionality most points are closer to a face of the hypercube than to their nearest neighbor. When a hypersphere is approximated by inscribing it in a hypercube, in high dimensions almost all the volume of the hypercube is outside the hypersphere (Domingos 2012). At the same time, in high dimensions, the ratio between the nearest and farthest points using Euclidean distance metric approaches 1, i.e., the points essentially become uniformly distant from each other (Aggarwal, Hinneburg, and Keim 2001). Thus, as all points are essentially uniformly distant from each other, the distinction is meaningless. This indicates as to why the Euclidean distance would therefore not be a good measure of distance in high dimensions. Since the spreading of points using the maximin design is done using the Euclidean distance, the design will experience the issues highlighted above making it inefficient as compared to the other methods. Perhaps this phenomenon also impacts the MaxPro designs, making the efficiency of the MaxPro design to decline as the dimensions increases as seen in Figure 3.12. Though this is not known.

As the UPD focusing on 2-dimensional uniformity, it is not highly impacted by the curse of dimensionality. Thereby retaining its high performance in comparison to the distance based designs with regards to minimizing high dimensional functions. The inefficacy of the maximin design in higher dimensions, due to the limitations of Euclidean distance metrics, suggests that such distance-based methods may not be well-suited for complex high-dimensional problems. Conversely, the UPD's use of the centered L_2 -discrepancy criterion allows it to mitigate the curse of dimensionality, maintaining its effectiveness across various dimensions. This highlights the need for careful consideration of design criteria and metrics in high-dimensional optimization tasks to ensure accurate and efficient results.

CHAPTER 4

Kriging Based Sequential Region Shrinkage with EGO for Hyperparameter Optimization

4.1 Introduction

Black-box optimization is a key challenge in many fields, including engineering, finance, and machine learning. It involves finding the optimal solution to a complex function that is either expensive or impossible to evaluate analytically. Formally, this can be expressed as:

$$oldsymbol{x}^* = \operatorname*{arg\,min}_{oldsymbol{x} \in \Omega} f(oldsymbol{x}),$$

where Ω represents the search space of \boldsymbol{x} . Optimizing $f(\boldsymbol{x})$ is a non-trivial task, typically requiring either derivative-based or derivative-free methods. Derivative-based methods depend on the calculation of the objective function's derivatives, making them suitable when $f(\boldsymbol{x})$ is smooth and differentiable, and its derivatives are easy to compute. However, in many real-world scenarios, the function is unknown or difficult to differentiate, making derivative-free methods preferable. These methods are especially useful in cases of black-box functions, that is, where the objective is not directly accessible.

Hyper-parameter Optimization (HPO) is a classic example of a black-box optimization problem, where the goal is to select the best set of hyper-parameters for a learning algorithm. These hyper-parameters are typically set before training and they control various aspects of the model. They include the learning rate, regularization, and the choice of the optimization algorithm. The model's performance can vary significantly based on the hyper-parameters chosen (Feurer and Hutter 2019), with results sometimes fluctuating drastically depending on

the architecture (Liu, Simonyan, and Yang 2018). Several methods are widely used for HPO, including grid search, random search (Bergstra and Bengio 2012), and Bayesian optimization (Pelikan, Goldberg, Cantú-Paz, et al. 1999).

Grid search systematically evaluates the model's performance for every combination of hyper-parameters within a predefined grid, choosing the configuration that yields the best result. Random search, on the other hand, samples hyper-parameters randomly from a predefined distribution and selects the best-performing configuration. Bayesian optimization constructs a probabilistic model of the objective function by combining prior knowledge with previously evaluated configurations. The posterior model is then optimized using an acquisition function, and the process is repeated until no further improvements can be made (Brochu, Cora, and De Freitas 2010).

While grid search and random search are simple and easy to implement, they become computationally expensive, especially in high-dimensional hyper-parameter spaces. Bayesian optimization is typically more efficient and effective for HPO (Snoek, Larochelle, and Adams 2012). One way to enhance the efficiency of Bayesian optimization is through the use of a surrogate model, which approximates the objective function and directs the search toward promising regions of the hyper-parameter space (Jones, Schonlau, and Welch 1998).

Bayesian optimization has gained popularity in solving black-box optimization problems due to its ability to handle noisy and non-convex functions, which are common in real-world scenarios. The probabilistic surrogate model used in Bayesian optimization is typically a Gaussian Process (GP), which is a flexible and a powerful tool for modeling complex functions (Rasmussen and Williams 2006). GPs can predict the value of the objective function at unexplored points while also providing an estimate of the uncertainty of these predictions.

In this paper, we propose a Kriging-based region shrinkage method that builds on Efficient Global Optimization (EGO) (Jones, Schonlau, and Welch 1998). The method sequentially refines the region of interest (ROI) based on a proportion of most informative data points, progressively shrinking the search space by reducing the size of the interval for each

hyper-parameter at each step. This approach allows us to focus more precisely on promising regions. By using EGO, the method balances exploration and exploitation within the domain.

The effectiveness of the proposed method is demonstrated using several well-known physical test functions from the Virtual Library of Simulation Experiments (Surjanovic and Bingham 2013). We compare our approach to existing optimization methods and show that the proposed derivative-free method achieves results on par with or exceeding expectations. Empirical results on DE hyperparameter optimization for constructing uniform projection designs show that our method requires fewer computational resources while performing comparably to, or better than, traditional hyper-parameter tuning methods such as grid search and random search. Although the theoretical guarantees are limited, the empirical results indicate strong potential for practical applications.

4.2 Background Theories

4.2.1 Related Work

In pursuit of minimizing loss, numerous optimization techniques based on the Bayesian approach have been developed. While some methods emphasize dimensionality reduction to enhance computational efficiency, the majority focus on reducing the size of the search space, commonly referred to as variable interval size reduction. The work presented here falls within this category.

Among the various strategies explored are the Controlled Gutmann-RBF (CG-RBF) method (Regis and Shoemaker 2007), the Trust Region Implementation in Kriging-based optimization with Expected Improvement (TRIKE) (Regis 2016), and the Trust-Region framework for Efficient Global Optimization (TREGO) (Diouane et al. 2023). Although many of these methods confine the search to a local region, the TREGO method alternates between local and global searches, returning to the global scale search after a successful local

search.

In TREGO, a local search is conducted when an iteration fails to achieve meaningful progress, meaning there is no significant improvement over the current best solution. Repeated failures progressively reduce the size of the local search space, while successful iterations trigger a global search. However, this process can lead to slow convergence.

To address this, we propose a modification: successful iterations will continue to focus on a reduced local search space, while unsuccessful iterations will initiate a global search. Here, a successful iteration is defined as one in which a point is found with a lower function value than any previously evaluated. We show that this proposed method leads to faster convergence compared to existing techniques, particularly TREGO.

4.2.2 The Efficient Global Optimization (EGO) Algorithm

The Efficient Global Optimization (EGO) algorithm builds upon the Expected Improvement (EI) criterion to sequentially explore the objective function. It begins with an initial design (typically a Latin hypercube) and iteratively visits the global maximizer of the EI criterion. At each step, the Kriging surrogate model is updated, including the re-estimation of hyperparameters, based on newly sampled points. This process continues until a convergence criterion is met or the budget of function evaluations is exhausted (Roustant, Ginsbourger, and Deville 2012). Algorithm 2 summarizes the procedure.

The EGO algorithm has proven to be highly effective and is frequently used in optimization problems where the number of objective function evaluations is severely limited. It is considered as one of the reference methods for global optimization in moderate-dimensional problems (typically $d \leq 10$) (Jones 2001).

Algorithm 2 EGO algorithm

Require: X, f = function to be minimized, n_{new} =number of points to add

- 1: Evaluate f at the design points X; y = f(X)
- 2: Build a kriging model based on X and y
- 3: **for** i in 1 to n_{new} **do**
- 4: Find $\boldsymbol{x}^* \leftarrow \arg \max_{\boldsymbol{x}} \mathbb{E}[I(\boldsymbol{x})]$
- 5: Evaluate $y^* \leftarrow f(\boldsymbol{x}^*)$
- 6: Update X and y with the new point x^* and response y^*
- 7: Update the kriging model
- 8: end for
- 9: Return $\boldsymbol{X}, \boldsymbol{y}$

4.2.3 The Trust Region EGO (TREGO)

Proposed by Diouane et al. (2023), Trust Region Efficient Global Optimization (TREGO) operates by searching within a dynamically adjusted trust region centered at the current best solution. The algorithm identifies new candidate points by optimizing within this region and evaluates the objective function. If the result is a success, the region expands, controlled by α . An iteration is deemed successful when there is a significant improvement on the result obtained from the current result by a margin. On the other hand, if poor, it contracts, controlled by β . After each evaluation, the surrogate model is updated with the new data, and the trust region is resized accordingly. This process balances global exploration and local refinement until stopping criteria are met. The stopping criteria mostly used is the maximum iterations. The default values used for α and β are 1/0.9 and 0.9 respectively. Algorithm 3 summarizes the procedure. A forcing function is used to determine whether a search is deemed successful or not. The forcing function $\rho(\sigma)$ is a positive continuous non-decreasing function such that $\rho(\sigma) \to 0$ when $\sigma \to 0$.

Algorithm 3 Simplified Trust-Region EGO (TREGO)

```
1: Input: Initial DoE X_0, function evaluations y_0, constants \alpha, \beta, d_{\min}, d_{\max}, initial step-
      size \sigma_0, initial best point \boldsymbol{x}_0^* \in \boldsymbol{X}_0
 2: Output: Best found point x_k^*
 3: Initialize: k = 0
 4: while stopping criterion is not met do
 5:
             Global Phase
                                                                                                                   /* Find global candidate */
             \boldsymbol{x}_{\text{global}} = \mathop{\mathrm{argmax}}_{\boldsymbol{x} \in \Omega} \, \mathbb{E} \left( \mathbf{I}(\boldsymbol{x}) \right)
 6:
            Update \boldsymbol{X}_{k+1} := \boldsymbol{X}_k \cup \boldsymbol{x}_k^{\text{global}} and y_{k+1} := \boldsymbol{y}_k \cup f(\boldsymbol{x}_k^{\text{global}})
 7:
            if f(\boldsymbol{x}_{\text{global}}) \leq f(\boldsymbol{x}_k^*) - \rho(\sigma_k): then
                                                                                                                    /* Successful global step */
 8:
                   Update best point: \boldsymbol{x}_{k+1}^* = x_{\text{global}}
 9:
                   Increase step size: \sigma_{k+1} = \alpha \sigma_k
10:
             else
11:
12:
                   Local Phase
                   Define trust region \Omega_k = \{ \boldsymbol{x} \in \Omega \mid d_{\min} \sigma_k \leq ||\boldsymbol{x} - \boldsymbol{x}_k^*|| \leq d_{\max} \sigma_k \}
13:
                                                                                                                     /* Find local candidate */
14:
                   \boldsymbol{x}_{\text{local}} = \operatorname{argmax} \mathbb{E}\left(\mathbf{I}(\boldsymbol{x})\right)
                   Update \boldsymbol{X}_{k+1} := \boldsymbol{X}_k \cup \boldsymbol{x}_k^{\mathrm{local}} and y_{k+1} := \boldsymbol{y}_k \cup f(\boldsymbol{x}_k^{\mathrm{local}})
15:
                   if f(\boldsymbol{x}_{local}) \leq f(\boldsymbol{x}_k^*) - \rho(\sigma_k): then
                                                                                                                      /* Successful local step */
16:
                         Update best point: \boldsymbol{x}_{k+1}^* = \boldsymbol{x}_{\text{local}}
17:
                         Increase step size: \sigma_{k+1} = \alpha \sigma_k
18:
                   else
19:
                         Keep current best point: \boldsymbol{x}_{k+1}^* = \boldsymbol{x}_k^*
20:
                         Reduce step size: \sigma_{k+1} = \beta \sigma_k
21:
                   end if
22:
             end if
23:
             Increment iteration counter: k = k + 1
24:
25: end while
               return Best found point x_k^*
```

4.3 The Proposed Algorithm

One key feature of the Efficient Global Optimization (EGO) algorithm is its ability to balance exploration and exploitation to efficiently search the solution space. By exploring broadly, EGO can uncover regions that are more likely to contain the global optimum, which is especially useful when dealing with complex and multimodal objective functions. However, this global search strategy can be computationally expensive, particularly when the search space is large or evaluating the objective function is time-consuming. In some cases, a localized search focusing on a smaller region may be more efficient.

To address these challenges, we propose a sequential region shrinkage method that alternates between a localized search and a global search. The global search evaluates whether further improvements can be made beyond the current local search region. This approach aims to strike a balance between efficient local searches and the broader exploration required for finding the global optimum. Algorithm 4 summarizes the procedure.

4.3.1 Region of Interest (ROI) Determination

In this method, a controlled parameter ρ determines the proportion of the top $100\rho\%$ of data points to use in calculating the new ROI. In the first iteration, the entire domain Ω serves as the initial ROI. EGO adds n_{new} points to the data set \boldsymbol{X} and \boldsymbol{y} , constituting a global search. Afterward, the top $100\rho\%$ of data points, based on a predefined ρ value, are selected. The ROI's boundaries are defined as the minimum and maximum of each dimension from these selected points. The ROI is then shifted to center it at the best-performing point, focusing the subsequent local search on this smaller region.

Figure 4.1 illustrates the ROI determination process.

In Figure 4.1(a), the contours represent the entire function domain Ω . We start with 10 points, selected using a maximin Latin hypercube design. These are represented as the black dots in Figure 4.1. Then 5 additional points are added using EGO. These additional points

Algorithm 4 Sequential Region Shrinkage Method

Require: X, f, Ω = domain, n_{new} , ρ

- 1: Evaluate f at the design points X; $y \leftarrow f(X)$
- 2: Build a Kriging model based on \boldsymbol{X} and \boldsymbol{y}
- 3: Set the region of interest (ROI) $\Omega' \leftarrow \Omega$ (initial domain)
- 4: while stopping criteria not met do
- 5: Run EGO within the domain Ω' to obtain the next n_{new} points
- 6: Update X and y with the new points
- 7: Update the Kriging model
- 8: Determine the new ROI $\Omega' \leftarrow \text{ROI}(\boldsymbol{X}, \boldsymbol{y}, \Omega, \rho)$
- 9: **if** small or no improvement (unsuccessful iteration) **then**
- 10: Restore the original domain: $\Omega' \leftarrow \Omega$
- 11: end if
- 12: end while
- 13: Return \boldsymbol{X} , \boldsymbol{y}

Algorithm 5 Determine Region of Interest (ROI)

Require: X, $\Omega = \text{domain}$, y, ρ .

- 1: $\tau \leftarrow \text{indices of the top } 100\rho\% \text{ of } \boldsymbol{y}$.
- 2: Select points in \boldsymbol{X} associated with τ .
- 3: Find the best point \boldsymbol{x}^* that minimizes the objective function
- 4: Determine the lower and upper bounds for points associated with τ ; set $L_j := \min_{i \in \tau} x_{ij}$ and $U_j := \max_{i \in \tau} x_{ij}$ for each $j = 1, \dots, d$
- 5: Set $\mathbf{D} := (\mathbf{U} \mathbf{L})/2$, where $\mathbf{U} = (U_1, \dots, U_d)$ and $\mathbf{L} = (L_1, \dots, L_d)$.
- 6: Set $\Omega' = \{ x^* D, x^* + D \} \cap \Omega$.
- 7: return Ω' .

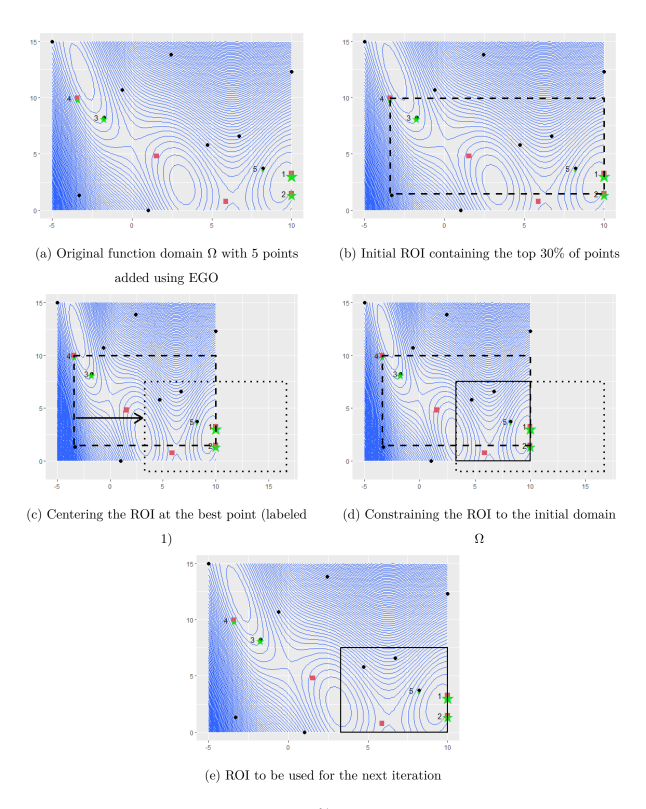


Figure 4.1: ROI determination using the top 30% of total points. The top points are labeled 1-5.

are indicated by the red squares. Taking $\rho = 0.3$, the best 30% of the 15 points, labeled 1-5 and indicated by green stars in decreasing order, are then used to define the new ROI, as shown in Figure 4.1(b). The ROI is centered around the best-performing point (labeled 1) in Figure 4.1(c). Since part of the ROI extends outside the original domain, it is constrained within Ω , resulting in the boxed region in Figure 4.1(d) and 4.1(e). This becomes the new ROI, Ω' , for the first iteration.

Next, a local search is performed within the newly defined ROI, and m points are added that maximize the Expected Improvement (EI). With each additional point, the Kriging model is updated to reflect the newly sampled points. If a point significantly improves upon the current minimum, the iteration is considered successful, and a new ROI is determined using the updated top $100\rho\%$ of data points. Otherwise, the method reverts to a global search.

This iterative process continues until a predefined stopping criterion is met, such as reaching a specified tolerance level or a maximum number of function evaluations. The stopping criterion can be adapted depending on whether the true optimum is known. If the optimum is known, we stop when the tolerance error is reached. Otherwise, we allow for a default of three global search iterations based on empirical tests, which showed that by the third global search, convergence typically occurs.

The region bounding strategy is designed to leverage the fact that a well-spaced design has a high likelihood of sampling points near the optimal region. By focusing the search in the most promising areas and selectively switching to global searches when necessary, the proposed method efficiently balances exploration and exploitation. As a result, it achieves the optimal solution with fewer function evaluations compared to standard EGO and TREGO algorithms.

4.3.2 Difference between RSO and TREGO

TREGO uses EGO to do optimization within the function domain. Once a point is obtained whose value is below the current function minimum by a given threshold, the iteration is considered to be successful. The next optimization is to run EGO in the entire function domain. In the case of an unsuccessful iteration, the region is shrunk by a factor of beta. The shrunk region is then centered in the current function minimum, where EGO is then run. The local optimization is only done after the global iteration is unsuccessful. This is different from the RSO since in the RSO, the region is determined by the top $100\rho\%$ of the data so far used. Also the RSO reverts to the global search after an unsuccessful iteration.

4.4 Numerical Experiments

To illustrate the efficiency of the method, various test functions of various dimensions from the Virtual Library of Simulation Experiments (Surjanovic and Bingham 2013) were taken into consideration. A maximin Latin hypercube design with n = 5d runs was selected as the initial design. The results were compared to those obtained by EGO and TREGO.

The test functions used and their settings are elaborated below:

1. Branin function (d=2)

$$f(x_1, x_2) = \left(x_2 - \frac{5}{4\pi^2}x_1^2 + \frac{5}{\pi}x_1 - 6\right)^2 + 10\left(1 - \frac{1}{8\pi}\right)\cos x_1 + 10$$

with $x_1 \in [-5, 10], x_2 \in [0, 15]$. The global minimum $f(\boldsymbol{x}^*) = 0.397887$ are located at $\boldsymbol{x}^* = (-\pi, 12.275), (\pi, 2.275)$ and (9.42478, 2.475).

2. Six-hump camel function (SixCamel) (d=2)

$$f(x_1, x_2) = 4x_1^2 - 2.1x_1^4 + x_1^6/3 + x_1x_2 - 4x_2^2 + 4x_2^4$$

with $-2 \le x_1 \le 2, -1 \le x_2 \le 1$. It has six extreme points, among them the global minimum are $\boldsymbol{x}^* = (0.0898, -0.7126), (-0.0898, 0.7126)$ and $f(\boldsymbol{x}^*) = -1.0316$.

3. Goldstein-Price function (d=2)

$$f(\mathbf{x}) = \left[1 + (x_1 + x_2 + 1)^2 \left(19 - 14x_1 + 3x_1^2 - 14x_2 + 6x_1x_2 + 3x_2^2\right)\right] \times \left[30 + (2x_1 - 3x_2)^2 \left(18 - 32x_1 + 12x_1^2 + 48x_2 - 36x_1x_2 + 27x_2^2\right)\right]$$

with $-2 \le x_1, x_2 \le 2$. The global minimum $f(\boldsymbol{x}^*) = 3$ is at point $\boldsymbol{x}^* = (0, -1)$ with several local minima around it.

4. Hartmann functions with d=3 (Hartmann3), d=4 (Hartmann4) and d=6 (Hartmann6)

$$f(\boldsymbol{x}) = -\sum_{i=1}^{4} \alpha_i \exp\left(-\sum_{j=1}^{d} A_{ij} (x_j - P_{ij})^2\right),$$

where $0 \le x_i \le 1, i = 1, 2, \dots, d$ and $\alpha = (1.0, 1.2, 3.0, 3.2)^T$. The elements of matrices $\boldsymbol{A}, \boldsymbol{P}$ and the global minimum are given in Table 4.1.

Functions with dimensions 6 or less were considered to be low dimensional functions. In the optimization, we used error between the function value evaluated at the best point and the known global optimum. We were interested in the number of function evaluations used by the algorithm to converge. This was replicated 20 times. The results were compared to the ones obtained when running EGO alone and when using TREGO. The results are reported in Table 4.2.

From the results above, we note that the proposed method converges with fewer function evaluations than both EGO and TREGO in all but one instance. This efficiency not only conserves computational resources but also is particularly beneficial in resource-limited scenarios. The exploitation characteristics of our method contribute to this performance. Notably, while our method generally outperform EGO and TREGO in all instances, for the Hartmann3 function and tolerance of 10^{-4} , TREGO performs slightly better than the proposed method.

Additionally, Figure 4.2 shows the minimization path for different test functions using the 3 methods, along with the 2 standard error bars at each step of the minimization path.

Table 4.1: Parameters for the Hartmann functions

Functions	Parameters
Hartmann3 $x^* = (0.1146, 0.5556, 0.8525)$ $f(x^*) = -3.86278$	$\mathbf{A} = \begin{pmatrix} 3.0 & 10 & 30 \\ 0.1 & 10 & 35 \\ 3.0 & 10 & 30 \\ 0.1 & 10 & 35 \end{pmatrix}; \mathbf{P} = 10^{-4} \begin{pmatrix} 3689 & 1170 & 2673 \\ 4699 & 4387 & 7470 \\ 1091 & 8732 & 5547 \\ 381 & 5743 & 8828 \end{pmatrix}$
Hartmann4 $\mathbf{x}^* = (0.1873, 0.1906, \\ 0.5566, 0.2647)$ $f(\mathbf{x}^*) = -3.135474$	$ \mathbf{A} = \begin{pmatrix} 10.0 & 0.05 & 3.0 & 17.00 \\ 3.0 & 10.00 & 3.5 & 8.00 \\ 17.0 & 17.00 & 1.7 & 0.05 \\ 3.5 & 0.10 & 10.0 & 10.00 \\ 1.7 & 8.00 & 17.0 & 0.10 \\ 8.0 & 14.00 & 8.0 & 14.00 \end{pmatrix}; \mathbf{P} = 10^{-4} \begin{pmatrix} 1312 & 2329 & 2348 & 4047 \\ 1696 & 4135 & 1451 & 8828 \\ 5569 & 8307 & 3522 & 8732 \\ 124 & 3736 & 2883 & 5743 \\ 8283 & 1004 & 3047 & 1091 \\ 5886 & 9991 & 6650 & 381 \end{pmatrix} $
Hartmann6 $\boldsymbol{x}^* = (0.2017, 0.1500, 0.4769 \\ 0.2753, 0.3117, 0.6573) \\ f(\boldsymbol{x}^*) = -3.32237$	$\mathbf{A} = \begin{pmatrix} 10 & 3 & 17 & 3.5 & 1.7 & 8 \\ 0.05 & 10 & 17 & 0.1 & 8 & 14 \\ 3 & 3.5 & 1.7 & 10 & 17 & 8 \\ 17 & 8 & 0.05 & 10 & 0.1 & 14 \end{pmatrix}$ $\mathbf{P} = 10^{-4} \begin{pmatrix} 1312 & 1696 & 5569 & 124 & 8283 & 5886 \\ 2329 & 4135 & 8307 & 3736 & 1004 & 9991 \\ 2348 & 1451 & 3522 & 2883 & 3047 & 6650 \\ 4047 & 8828 & 8732 & 5743 & 1091 & 381 \end{pmatrix}$

Table 4.2: Number of function evaluations to achieve the desired tolerance level

		mean			median		
Function	tolerr	EGO	TREGO	RSO	EGO	TREGO	RSO
D '	10^{-3}	35.75	28.75	26.25	35	30	25
Branin	10^{-4}	42	37.25	32.5	40	40	30
S:C1	10^{-3}	43.50	41.00	33.75	45	40	35
SixCamel	10^{-4}	48.75	46.75	40.75	50	50	40
	10^{-3}	67	52.5	41	70	57.5	35
Goldstein-Price	10^{-4}	70	70	42.5	70	70	37.5
	10^{-3}	23.4	21.2	20.8	24	20	20
Hartmann3	10^{-4}	30.8	25.75	26.4	26	22	24
II	10^{-3}	46.0	44.0	40.0	44	42	36
Hartmann4	10^{-4}	63.6	55.2	53.6	46	44	40
II	10^{-3}	60.5	58.5	49.5	64	61	42.5
Hartmann6	10^{-4}	67.5	62	52	70	67.5	47.5

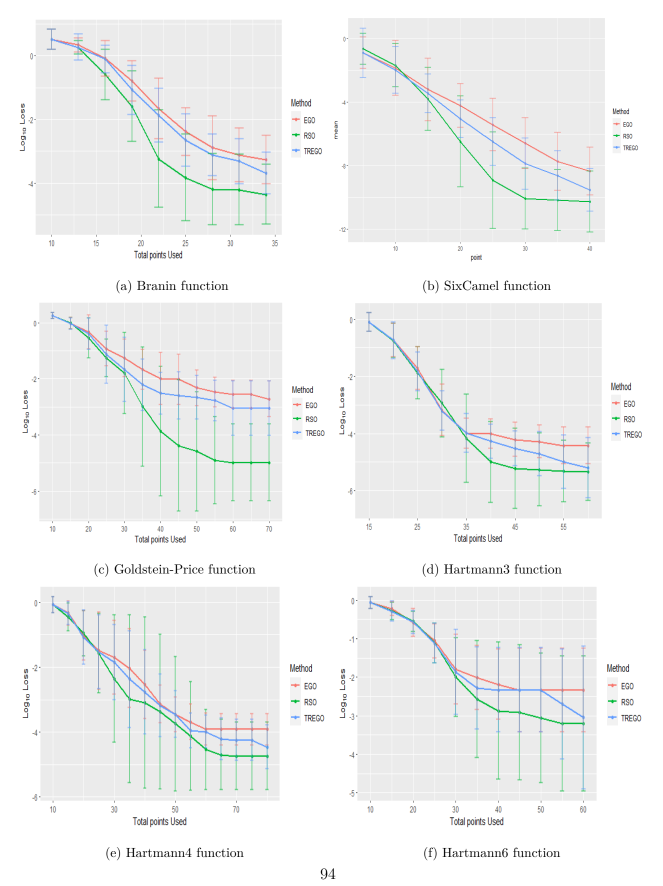


Figure 4.2: Comparison of 3 methods for different test functions

The log mean error analysis indicate that the proposed method rapidly decreases the error rate before stabilizing, signifying a faster convergence compared to the alternatives. In constrained resource settings, where only 15-25 new observations can be made, our method achieves a $\log 10$ error of at most -3 across all tested functions.

4.5 Application in Generating Uniform Projection Designs

We consider tuning the DE algorithm from UniPro package to generate uniform projection designs (UPDs). As the DE algorithm is stochastic, the generation is replicated 10 times and the mean of the 10 criterion values obtained. This is done to minimize variation. This is considered to be the objective function to be optimized.

As described in Chapter 2, the DE algorithm has five hyperparameters: NP, itermax, pCR, pMut and pGBest. We use EGO, TREGO and RSO to tune DE hyperparameters. Starting with a random Latin hypercube of size 5d = 25, we obtain the minimization path by sequentially adding 25 points. This is replicated 20 times. Figure 4.3 shows the minimization path and Figure 4.4 shows the boxplots of the final optimal objective values for three methods. The RSO method tends to generally provide smaller objective values than the EGO or TREGO method.

We also take a look at the distribution of the optimal hyperparameter settings from the 20 replicates in Figure 4.5 obtained from the RSO method, where NP and itermax are rescaled to [0, 1]. This determines as to whether the optimal region is the same point or we are converging to a local optimum. The 30×3 target design indicates that the process does not converge to the same value each time. On the other hand, from the 50×5 and 70×7 target designs, we see that the values for NP and itermax are concentrated at the upper vertex. Finally, from the 70×7 target design, we find out that the values for pCR and pGBest are also concentrated at the upper end. However, none of the target designs give information about where the probability of mutation (pMut) should lie. We examine the correlations to determine as to whether there exists a two way correlation among the

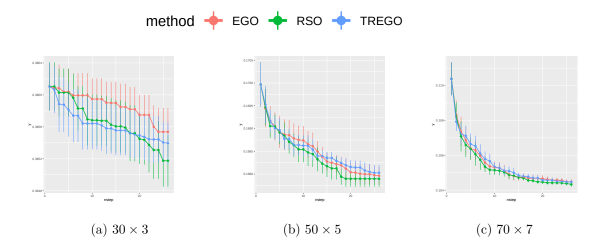


Figure 4.3: Minimization path for 3 methods in the construction of UPDs using DE

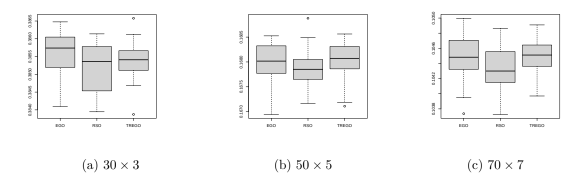


Figure 4.4: Comparison of optimal results from 3 methods

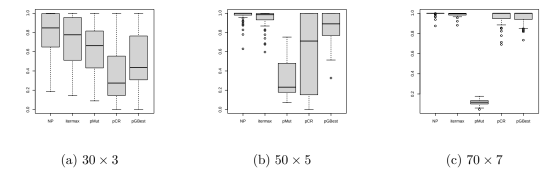


Figure 4.5: Distribution of optimal hyperparameter settings

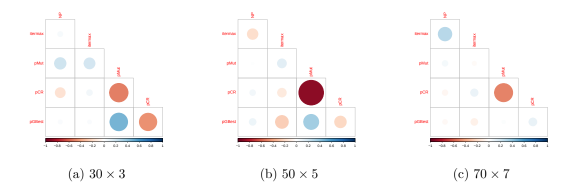


Figure 4.6: Correlation plot of the optimal hyperparameter settings

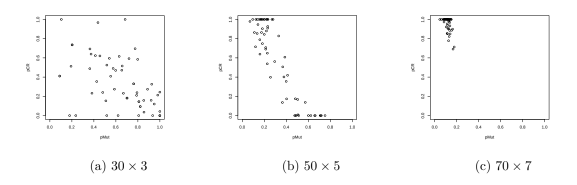


Figure 4.7: Scatter plots showing the relation between pMut and pCR

parameters.

From the correlation plots in Figure 4.6, we do notice that pMut is highly correlated to pCR. These two hyperparameters could be optimized simultaneously while holding the other 3 hyperparameters (NP, itermax, pGBest) at their highest level. The justification for this is due to the fact that NP and itermax could be thought of as the budget size. For the probability of using global best, this value should be high since we would want to be as close as possible to the global minimum. Figure 4.7 shows the scatter plots of pMut and pCR. These two hyperparameters are intriguing as they are highly negatively correlated.

Having set these three at their maximum level, an optimization is carried out on the two remaining parameters, pMut and pCR. The results obtained do not differ from the 5 hyperparameter optimization, though takes a bit longer since NP and itermax have been

Table 4.3: Optimal hyperparameter settings for each target design with 1024-run LHD

	NP	itermax	pMut	pCR	pGBest	$\phi(\cdot)$
30×3	95	917	0.63	0.19	0.72	0.3850
50×5	98	1485	0.40	0.38	0.88	0.1685
70×7	84	1408	0.11	0.92	0.86	0.1066

held at their highest level.

Finally, we compare the RSO method with other methods in generating UPDs. We consider the random search method, the grid search method, the DE1 and DE4 methods provided by Stokes, Wong, and Xu (2024), and the DE0ptim method based on the second order model in Chapter 2. The random search used a random Latin hypercube design with 1024 runs and the grid search method used a 4⁵ full factorial design. The optimal factor combinations obtained for the random search and grid search are given in Tables 4.3 and 4.4, respectively.

DE1 always uses the global best (i.e., pGBest=1), while DE4 is a hybrid version with pGBest=0.5, which randomly chooses the global best, the current agent and a random agent with probability 50%, 25%, and 25% for each column independently. Both DE1 and DE4 use NP = 100, itermax = 1500, pMut = 0.1 and pCR = 0.5. The DEoptim method from Chapter 2 uses NP = 100, itermax = 1500, pGBest = 0.95 and optimal settings of (pMut, pCR) as (0.95, 0.05), (0.25, 0.75), and (0.15, 0.85), respectively, for target size 30×3 , 50×5 and 70×7 . Table 4.5 gives the optimal hyperparameter settings obtained from the RSO method.

For a given target size and each method, we generate 100 UPDs using the respective optimal factor combination and compute their objective $\phi(\cdot)$ values. Figure 4.8 compares the objective values of the generated UPDs from the six methods.

Figure 4.8 indicates the advantage of using optimized methods as compared to random

Table 4.4: Optimal hyperparameter settings for each target design with 4^5 FFD

	NP	itermax	pMut	pCR	pGBest	$\phi(\cdot)$
30×3	100	1167	0.95	0.05	0.65	0.3845
50×5	100	1500	0.35	0.95	0.95	0.1684
70×7	100	1500	0.35	0.35	0.95	0.1055

Table 4.5: Optimal hyperparameter settings for each target design using RSO

	NP	itermax	pMut	pCR	pGBest
30×3	100	1452	0.95	0.05	0.74
50×5	100	1500	0.19	0.63	0.74
70×7	100	1498	0.13	0.95	0.95

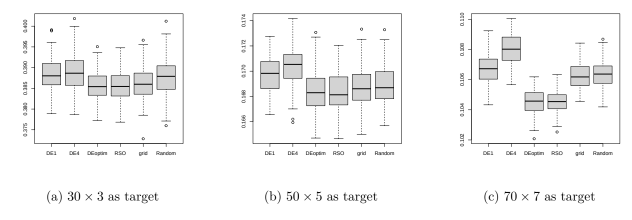


Figure 4.8: Comparison of six methods for constructing UPDs

search or grid search. Even using 1024-runs to determine the optimal settings, the grid and random search methods perform poorly as compared to RSO and DEoptim. On the other hand, DE1 and DE4 perform the worst as these methods use arbitrary settings selected by Stokes, Wong, and Xu (2024).

Figure 4.8 also indicates that using the RSO as compared to the default DE1 and DE4 is highly effective when the dimension of the target design increases. This is because with increase in dimension of target design, the DE1/DE4 algorithms are still far from the optimal solution, while the RSO carries out an additional minimization which moves closer to the optimal hyperparameter settings. In addition, RSO and DEoptim have similar performance despite the use of different settings for pCR, pMut and pGBest. One advantage of using RSO over DEoptim is the notion that for DEoptim, one has to determine which design is best, to carry out the optimization. In Chapter 2, it is shown that the 50-run OACD is the best. On the other hand, RSO uses a 25-run LHD with additional 25 points for optimization. No prior knowledge of the design is needed. Furthermore, determining the optimal settings in DEoptim is quite a task when there exists significant interactions between the factors, while for RSO, the method yields the optimal settings directly.

CHAPTER 5

Conclusion and Future Directions

The studies presented in this work advance the understanding and application of optimization methods and experimental design in addressing complex, high-dimensional challenges in design construction. Optimization algorithms, especially Differential Evolution (DE) and Bayesian optimization, have demonstrated their capability to tackle intricate real-world problems requiring efficient navigation of extensive search spaces. This dissertation has focused on three interlinked studies: optimizing DE hyperparameters for constructing Uniform Projection Designs (UPDs), evaluating initial design choices for prediction and optimization, and introducing a Kriging-based sequential shrinkage method for enhancing Bayesian optimization. Collectively, these studies provide essential insights into how the structure and strategy of optimization methods can be tailored to suit varied problem domains, resulting in improved accuracy, efficiency, and scalability.

This chapter synthesizes the findings of these studies, discussing their implications and highlighting directions for future work in advancing optimization techniques.

5.1 Key Findings Across the Studies

5.1.1 Optimizing Hyperparameters in Differential Evolution for UPD Generation

The first study explored the hyperparameter landscape of a modified DE algorithm applied to discrete tasks, such as generating Uniform Projection Designs (UPDs). DE is traditionally

effective for continuous optimization, but modifications to its structure are necessary to handle discrete variables. Inspired by recent work on DE adaptations, this study focused on five critical DE hyperparameters, including population size, maximum iterations, and probabilities related to mutation, crossover and use of the global best. Through extensive testing, OACD and CCD emerged as the most effective experimental designs for exploring DE's hyperparameter response surface, outperforming space-filling designs in terms of root mean squared error and correlation with the underlying objective.

This analysis, built upon second-order model as the surrogate model, revealed that second-order models offer a reliable yet simpler alternative to more complex surrogate approaches, such as Kriging and heteroskedastic Gaussian Processes (GPs). This is significant because identifying optimal hyperparameter settings is not only crucial for enhancing DE's performance but also for constructing high-quality UPDs, which have applications in a variety of fields that require uniformity across high-dimensional spaces. The study highlights the importance of structured hyperparameter tuning in optimizing DE's utility, establishing OACD and CCD as powerful tools for practitioners seeking to leverage DE in discrete optimization contexts.

5.1.2 Efficiency and Robustness of UPDs in High-Dimensional Prediction and Optimization Tasks

The second study assessed the effectiveness of various initial design strategies within the framework of prediction and optimization. By using Gaussian Process (GP) models as surrogate and focusing on active learning through selective sampling, this study highlighted the influence of initial design on prediction accuracy and optimization efficiency. In particular, Uniform Projection Designs (UPDs) consistently outperformed distance-based designs, such as maximin and MaxPro designs, especially in high-dimensional spaces. This advantage is attributed to the UPD's focus on two-dimensional uniformity via the centered L_2 -discrepancy criterion, which circumvents the inefficiencies associated with Euclidean distance metrics in

high-dimensional contexts and curse of dimensionality.

The performance of UPDs highlights the limitations of distance-based designs in high dimensions. Euclidean-based metrics, such as those used in maximin designs, tend to falter in high-dimensional settings where points become uniformly distant from one another. In contrast, UPDs maintain efficiency by avoiding reliance on high-dimensional distances, instead leveraging two-dimensional uniformity, which ensures more consistent prediction accuracy across varying dimensionalities. This study's findings underscore the importance of initial design choice in prediction and also in optimization via EGO processes, particularly when early-stage optimization has a pronounced impact on convergence and overall accuracy.

5.1.3 Introducing a Sequential Shrinking Approach to Bayesian Optimization for Resource-Efficient Tuning

The third study introduced a Kriging-based sequential region shrinking method, enhancing traditional Bayesian optimization techniques by focusing on progressively smaller, more promising regions in the search space. This novel approach incorporates aspects of EGO by iteratively narrowing the search region, allowing the optimization process to converge more efficiently. By targeting high-potential areas in each iteration, the approach significantly reduces the computational resources required compared to traditional grid and random search methods.

The sequential shrinking method not only performs effectively across various test functions but also demonstrates advantages over existing Bayesian strategies, such as TREGO. This is particularly important in resource-constrained scenarios, where reducing computational load without sacrificing optimization quality is paramount. The method's integration of Kriging with a focused region-based approach makes it a practical choice for applications in high-dimensional spaces, where traditional optimization strategies may become computationally prohibitive. As such, this study's findings contribute to a growing body of work emphasizing the role of adaptive, resource-efficient techniques in optimization.

5.2 Implications for Optimization Strategies and Practical Applications

This dissertation's findings emphasize the role of both design selection and adaptive optimization strategies in achieving high performance in complex, high-dimensional tasks. The successful use of UPDs as initial designs in EGO demonstrates that high-dimensional optimization can be more effectively approached by leveraging two-dimensional uniformity rather than high-dimensional Euclidean distances. This insight is essential for practitioners in fields where high-dimensional optimization is necessary, offering a robust alternative to conventional designs that suffer in high-dimensional spaces.

Moreover, the effectiveness of structured search techniques, particularly the novel Kriging-based sequential region shrinking method, highlights the potential of region-focused optimization in resource-constrained applications. By progressively shrinking the search region, this method achieves convergence with reduced computational requirements, which has practical implications for industries where efficiency and speed are critical. Similarly, the use of OACD and CCD in hyperparameter tuning underscores the continued relevance of classical factorial composite designs and the response surface methodologies, particularly when dealing with highly sensitive or discrete parameters in algorithms like DE.

For practitioners, these studies provide practical insights into choosing appropriate design and search techniques based on problem dimensionality, optimization objectives, and computational constraints. The findings also underscore the value of adaptive frameworks in both design and search, offering a balance between exploration and exploitation that improves optimization outcomes.

5.3 Limitations and Future Directions

Despite the strengths of the findings, this research also identifies limitations that warrant further investigation. First, while the current studies provide evidence for the effectiveness of UPDs in high-dimensional optimization, additional work is needed to explore the theoretical underpinnings of UPDs' performance. The impact of factor interactions in the DE algorithm remains an area for future exploration, particularly in developing more nuanced models that account for interaction effects without introducing excessive model complexity.

Additionally, the sequential shrinking approach, while efficient, could benefit from integration with TREGO or with other advanced machine learning models or ensemble methods, potentially enhancing its adaptability and performance in even more complex search spaces. For instance, exploring hybrid approaches that combine GP models with UPD-based designs or integrating deep learning models for dynamic surrogate modeling may further improve accuracy and computational efficiency.

Furthermore, the Kriging-based shrinking approach and DE hyperparameter optimization would benefit from more comprehensive evaluations across different domains. Future work might explore alternative acquisition functions that enhance adaptability in Bayesian optimization, especially for applications requiring specific trade-offs between exploration and exploitation. Extending these techniques to larger datasets or real-world applications would also validate their utility in practical settings, where high dimensionality and computational costs pose ongoing challenges.

Finally, this research although focuses exclusively on the construction of Uniform Projection Designs (UPDs), it is generalizable to the construction of various space filling designs. We considered the UPD as there is no established construction methods that currently exist. In future studies, however, we can apply the proposed procedures to construct other space filling designs such as MaxPro designs or uniform designs and then compare these with existing designs to assess the efficiency of the proposed methods.

5.4 Final Remarks

In conclusion, this work advances the field of high-dimensional optimization by evaluating and enhancing design and optimization strategies for hyperparameter tuning, experimental design generation, and adaptive region-focused optimization. The findings from the three studies provide a cohesive framework that balances theoretical insights with practical applicability, ultimately paving the way for more robust and efficient optimization techniques. This research not only enhances our understanding of the design and optimization landscape but also contributes valuable tools for tackling complex challenges across scientific, engineering, and industrial applications. The evolving methodologies and insights presented here have significant potential to guide future innovations, leading to more accurate, scalable, and resource-efficient solutions for a wide array of optimization problems.

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