An Adjoint-based Optimization Method Using the Solution of Gray-box Conservation Laws

by

Han Chen

Submitted to the Department of Aeronautics and Astronautics in partial fulfillment of the requirements for the degree of

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Abstract

Many design applications can be formulated as optimization constrained by conservation laws. Such optimization can be efficiently solved by the adjoint method, which computes the gradient of the objective to the design variables. Traditionally, the adjoint method has not been able to be implemented in "gray-box" conservation law simulations. In gray-box simulations, the analytical and numerical form of the conservation law is unknown, but the full solution of relevant flow quantities is available. Optimization constrained by gray-box simulations can be challenging for high-dimensional design because the adjoint method is not directly applicable.

We consider the case where the flux function is unknown in the gray-box conservation law. The twin model method is presented to estimated the gradient by inferring the flux function from the space-time solution. The method enables the estimation of the gradient by solving the adjoint equation associated with the inferred conservation law. Building upon previous research, a Gaussian process optimization (GPO) framework, the twin-model GPO, is presented that admits the estimated gradient. The effectiveness of the proposed optimization method is compared to a conventional GPO method where the gradient is unavailable. The performance of the conventional method is found to deteriorate as the optimization dimensionality increases. Adopting the twin-model GPO results in improved objective function given a limited number of gray-box simulations.

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Contents

1	Bac	kgroui	nd	15
	1.1	Motiv	ation	15
	1.2	Proble	em Formulation	21
	1.3	Litera	ture Review	22
		1.3.1	Review of Optimization Methods	22
		1.3.2	The Adjoint Method	26
		1.3.3	Adaptive Basis Construction	31
		1.3.4	Model Calibration	34
		1.3.5	Bayesian Optimization	36
	1.4	Notat	ions	40
	1.5	Thesis	s Objectives	41
	1.6	Outlin	ne	42
2	Esti	imate	the Gradient by Using the Space-time Solution	43
	2.1	Appro	oach	43
		2.1.1	Parameterization	47
		2.1.2	Elements for Adaptive Basis Construction	52
		2.1.3	Algorithm	57
		2.1.4	Minimizing the Truncation Error	59
	2.2	Nume	rical Results	62
		2.2.1	Buckley-Leverett Equation	63
		2.2.2	Navier-Stokes Flow	64
		2.2.3	Polymer Injection in Petroleum Reservoir	67

	2.3	Chapter Summary	74
3	Lev	eraging the Twin Model for Bayesian Optimization	7 5
	3.1	Approach	75
		3.1.1 Modeling the Objective and Gradient by Gaussian Processes .	75
		3.1.2 Algorithm	79
		3.1.3 Convergence Properties Using True Hyper Parameters	81
	3.2	Numerical Results	83
		3.2.1 Buckley-Leverett Equation	83
		3.2.2 Navier-Stokes Flow	86
		3.2.3 Polymer Injection in Petroleum Reservoir	86
	3.3	Chapter Summary	92
4	Con	nclusions	93
	4.1	Thesis Summary	93
	4.2	Contributions	95
	4.3	Future Work	96
\mathbf{A}	Pro	of of Theorems	97
	A.1	Theorem 1	97
	A.2	Theorem 2	99
	A.3	Theorem 3	102

List of Figures

1-1	The computational graph for (1.16) . The yellow nodes indicate the	
	input variables, the blue node indicates the output variable, and the	
	white nodes indicate the intermediate variables. The arrows indicate	
	elementary operations. The begining and end nodes of each arrow	
	indicate the independent and dependent variables for each operation.	29
1-2	Computational graphs for the PDE simulation and objective evaluation.	31
2-1	Domain of dependence: $u(t,x)$ depends on u at an earlier time within	
	a domain of dependence. The two planes in this figure indicates the	
	spatial solution at two adjacent timesteps. The domain of dependence	
	can be much smaller than Ω , the entire spatial domain	45
2-2	An illustration of B_u defined in Theorem 1. The blue line is u_0 and	
	the green dashed line is $\frac{du_0}{dx}$. B_u is the set of u_0 where the derivative	
	$\frac{du_0}{dx}$ has an absolute value larger than γ	46
2-3	An example of mother wavelet, the Meyer wavelet	48
2-4	Red line: the integral (2.7) of the Meyer wavelet. Black line: the	
	logistic sigmoid function	49
2-5	An ad hoc set of bases	50
2-6	Space time solutions	51
2-7	The objective function ξ evaluated by either the gray-box model or the	
	trained twin model	51

2-8	The first row shows the three different initial conditions used to generate		
	the gray-box space-time solution. The second row compares the trained		
	\tilde{F} (blue) and the Buckley-Leverett F (red). The third row compares		
	the trained $\frac{d\tilde{F}}{du}$ (blue) and the Buckley-Leverett $\frac{dF}{du}$ (red). The green		
	background highlights the domain of u where the gray-box space-time		
	solution exists.	52	
2-9	An illustration of the tuple representation and the corresponding univariate)	
	sigmoid	54	
2-10	Neighborhood for univariate bases. (a) shows the neighborhood (blue)		
	of a single basis (red). (b) shows the neighborhood (blue) of several		
	bases (red). The left column represents the basis on the $(j, \frac{\eta}{2^j})$ plane,		
	and the right column shows the actual basis $\phi_{j,\eta}$	56	
2-11	The discretized gray-box solution is shuffled into 3 sets, each indicated		
	by a color	56	
2-12	The basis dictionary for the three solutions in Figure 2-8	63	
2-13	The errors of estimated gradients for the three solutions	64	
2-14	The return bend geometry and the mesh for the simulation	65	
2-15	Left column: an example gray-box solution for a given geometry. Right		
	column: the solution mismatch after training a twin model	67	
2-16	The gray-box state equation (right column) and the trained state equation		
	(left column). The gray-box model uses either the ideal gas equation		
	(first row) or the Reclich-Kwong equation (second row). The convex		
	hull of the gray-box solution is shown by the dashed red line	68	
2-17	A comparison of the estimated gradient and the true gradient	69	
2-18	Water flooding in petroleum reservoir engineering (courtesy from PetroWik	ai).	70
2-19	The geometry of the petroleum reservoir	71	
2-20	The isosurfaces of $S_w = 0.25$ and $S_w = 0.7$ at $t = 30$ days	72	
2-21	The gradient of ξ with respect to rates at the two injectors. The lines		
	indicate the gradients estimated by the twin model, while the stars		
	indicate the true gradient evaluated by finite difference	73	

3-1	The flowchart of Algorithm 3	80
3-2	Optimized results for the Buckley-Leverett equation	84
3-3	A comparison of the optimized $u(t=1,x)$ after 20 gray-box simulations. The red line is obtained by the vanilla Bayesian optimization and the green line by the twin-model Bayesian optimization. The cyan dashed line indicates the $u(t=1,x)$ obtained by setting the source term to zero	85
3-4	The current best objective at each iterate. The red line is obtained	00
0 1	by the vanilla Bayesian optimization and the green line by the twin-model Bayesian optimization. The black horizontal line indicates the true optimal	85
3-5	The left plot shows the initial guess of control points (blue dots), the initial guess of the geometry (blue line), the optimized control points (red dots), and the optimized geometry (red line). The purple squares indicate the bound constraints for each control point. The right plot shows the pressure along the interior and the exterior boundaries for	
	the initial (blue) and the optimized (red) geometry	87
3-6	The current best objective at each iterate for the ideal gas and the Redlich-Kwong gas. The green lines are obtained by the twin-model Bayesian optimization. The red lines are obtained by the vanilla Bayesian optimization. The black horizontal lines indicate the true optimal	87
3-7	The cumulative and per-iterate wall clock time, in minutes	88
3-8	The permeability of the reservoir, in 100 milli Darcy. The 5 injectors are indicated by the black dots, and the producer is indicated by the	
	green dot	89
3-9	The current best objective evaluation against the number of iterates.	90
3-10	$\xi(t)$ for the initial and the optimized injection rates	90
3_11	The optimized time-dependent injection rates	91

3-12	The current best objective evaluation using the backtracking-Armijo	
	gradient descent method, where the gradient is provided by the twin	
	model	91
A-1	The state-space trajectories of the gray-box model and the twin model.	
	${\cal M}$ measures the difference of the twin model trajectory (blue) with the	
	gray-box trajectory (red). $\mathcal T$ measures the difference of the twin model	
	trajectory with restarts (green) and the gray-box trajectory (red)	100

List of Tables

2.1	The integrated errors of the estimated gradients for the three solutions.	64
2.2	The error of the gradient estimation, in percentage	67
2.3	The error of estimated gradient at day 2, 16, 30, and 44, in percentage.	73

Chapter 1

Background

1.1 Motivation

A conservation law states that a particular property of a physical system does not appear or vanish as the system evolves over time, such as the conservation of mass, momentum, and energy. Mathematically, a conservation law can be expressed locally as a continuity equation (1.1),

$$\frac{\partial u}{\partial t} + \nabla \cdot F = q, \qquad (1.1)$$

where u is the conserved physical quantity, t is time, F is the flux of u, and q is the source for u. Many equations fundamental to the physical world, such as the Navier-Stokes equation, the Maxwell equation, and the porous medium transport equation, can be described by (1.1).

Optimization constrained by conservation laws is present in many engineering applications. For example, in gas turbines, the rotor blades can operate at a temperature close to 2000K [9]. To prevent material failure due to overheating, channels can be drilled inside the rotor blades to circulate coolant air whose dynamics are governed by the Navier-Stokes equation [6]. The pressure used to drive the coolant flow is provided by the compressor, resulting in a penalty on the turbine's thermo-dynamic efficiency

[7]. Engineers are thereby interested in optimizing the coolant channel geometry in order to suppress the pressure loss. In this optimization problem, the control variables are the parameters that describe the channel geometry. The dimensionality of the optimization is the number of control variables, i.e. the control's degree of freedom. Another example is the field control of petroleum reservoir. In petroleum reservoir, the fluid flow of various phases and chemical components is dictated by porous medium transport equations [4]. The flow can be passively and actively controlled by a variety of techniques [1], such as the wellbore pressure control, the polymer injection, and the steam heating, where the reservoir is controlled by the pressure at each wells, by the the injection rate of polymer, and by the temperature of the steam [5]. These controls can be parameterized in time for each well, polymer injector, and steam injector. The dimensionality of the optimization is the total number of these control variables. Driven by economic interests, petroleum producers are devoted to optimizing the controls for enhanced recovery and reduced cost.

Such optimization is being revolutionized by the numerical simulation and optimization algorithms. On one hand, conservation law simulation can provide an evaluation of a candidate control that is cheaper, faster, and more scalable than conducting physical experiments. On the other hand, advanced optimization algorithms can guide the control towards the optimal with reduced number of simulation [34, 35, 36, 42, 46, 47, 48, 61]. However, optimization based on conservation law simulation can still be overwhelmingly costly. The cost is two-folded: Firstly, each simulation for a given control may run for hours or days even on a high-end computer. This is mainly because of the high-fidelity physical models, the complex numerical schemes, and the large scale space-time discretization employed in the simulation. Secondly, optimization algorithms generally take many iterations of simulation on various controls. The number of iterations required to achieve near-optimality usually increases with the control's degree of freedom [52]. The two costs are multiplicative. The multiplicative effect compromises the impact of computational efforts among field engineers.

Fortunately, the cost due to iteration can be alleviated by adopting gradient-based optimization algorithms [52]. A gradient-based algorithm requires significantly less iterations than a derivative-free algorithm for problems with many control variables. Gradient-based algorithms require the gradient of the optimization objective to the control variables, which is efficiently computable through the adjoint method [10]. The adjoint method propagates the gradient from the objective backward to the control variables through the path of time integration [10] or through the chain of numerical operations [17]. To keep track of the back propagation, the simulator source code needs to be available. In real-world industrial simulators, adjoint is scarcely implemented because most source codes are proprietary and/or legacy. For example, PSim, a reservoir simulator developed and owned by ConocoPhillips, is a multimillion-line Fortran-77 code that traces its birth back to the 1980's. Implementing adjoint directly into the source code is unpreferable because it can take tremendous amount of brain hours. Besides, the source code and its physical models are only accessible and modifiable by the computational team inside the company. For the sake of gradient computation, *PSim* has been superceded by adjoint-enabled simulators, but it is difficult to be replaced due to its legacy use and cost concerns. proprietary and legacy nature of many industrial simulators hinders the prevalence of the adjoint method and gradient-based algorithms in many real-world problems with high-dimensional control.

Despite their proprietary and legacy nature, most simulators for unsteady conservation laws are able to provide the discretized space-time solution of relevant flow quantities. For example, *PSim* provides the space-time solution of pressure, saturation, and concentration for multi-phase flow. Similarly, most steady state simulators are able to provide the spatial solution. the discussion will focus on the unsteady case, since a steady state simulator can be viewed as a special case of the unsteady one where the solution remains the same over many time steps.

I argue that the adjoint gradient computation may be enabled by leveraging

the space-time solution. The discretized space-time solution provides invaluable information about the conservation law hardwired in the simulator. For illustration, consider a code which simulates

$$\frac{\partial u}{\partial t} + \frac{\partial F(u)}{\partial x} = c, \quad x \in [0, 1], \quad t \in [0, 1]$$
(1.2)

with proper intial and boundary conditions and F being differentiable. c indicates the control that acts as a source for u. If the expression of F(u) in the simulator is not accessible by the user, adjoint can not be implemented directly. However, F may be partially inferred from a discretized space-time solution of u for a given c. To see this, let the discretized solution be $\mathbf{u} \equiv \{u(t_i, x_j)\}_{i=1,\dots,M,\ j=1,\dots,N}$, where $0 \le t_1 < t_2 < \dots < t_M \le 1$ and $0 \le x_1 < x_2 < \dots < x_N \le 1$ indicate the time and space discretization. Given \mathbf{u} , the $\frac{\partial u}{\partial t}$ and $\frac{\partial u}{\partial x}$ can be sampled by finite difference. Because (1.2) can be written as

$$\frac{\partial u}{\partial t} + \frac{dF}{du}\frac{\partial u}{\partial x} = c, \quad x \in [0, 1], \quad t \in [0, 1]$$
(1.3)

away from the shock wave, the samples of $\frac{\partial u}{\partial t}$ and $\frac{\partial u}{\partial x}$ can be plugged into (1.3) to obtain samples of $\frac{dF}{du}$. The reasoning remains intact at the shock wave, where $\frac{dF}{du}$ in (1.3) is replaced by the finite difference form $\frac{\Delta F}{\Delta u}$ according to the Rankine-Hugoniot condition. Based upon the sampled $\frac{dF}{du}$ and $\frac{\Delta F}{\Delta u}$, the unknown flux function F can be approximated up to a constant for values of u that appeared in the solution, by using indefinite integral. Let \tilde{F} be the approximation for F. An alternative conservation law can be proposed

$$\frac{\partial \tilde{u}}{\partial t} + \frac{\partial \tilde{F}(\tilde{u})}{\partial \tilde{u}} = c, \quad x \in [0, 1], \quad t \in [0, 1], \tag{1.4}$$

that approximates the true but unknown conservation law (1.2), where \tilde{u} is the solution associated with \tilde{F} , in the following sense: If \tilde{F} and F are off by a constant, the solutions of (1.2) and (1.4) to any initial value problem will be the same. In addition, the solutions to any adjoint equation, with any objective function, will be

the same. As a result, the gradient computed by the adjoint equation of (1.4) will be the true gradient, and therefore can drive the optimization constrained by (1.2). A simulator for the approximated conservation law is named **twin model**, since it behaves as an adjoint-enabled twin of the original simulator. If a conservation law has a system of equations and/or has a greater-than-one spatial dimension, the above simple method to recover the flux function from a solution will no longer work. Nonetheless, I expect a lot of information for the flux function can be extracted from the solution. Given some additional information of the conservation law, one may be able to recover the unknown aspects of the flux function. The details of this topic are discussed in Chapter 2.

My thesis focuses on a class of simulators that I call **gray-box**. A simulator is defined to be gray-box if the following two conditions are met:

- 1. the adjoint is unavailable, and is impractical to implement into the source code.
- 2. the full space-time solution of relevant flow quantities is available.

Many industrial simulators, such as *PSim*, satisfy both conditions. In contrast, a simulator is named **open-box** if condition 1 is violated. For example, *OpenFOAM* [53] is an open-source fluid simulator where adjoint can be implemented directly into its source code, so it is open-box by definition. Open-box simulators enjoy the benifit of efficient gradient computation brought by adjoint, thereby are not within the research scope of my thesis. If condition 1 is met but 2 is violated, a simulator is named **black-box**. For example, *Aspen* [54], an industrial chemical reactor simulator, provides neither the adjoint nor the full space-time solution. Black-box simulators are simply calculators for the objective function. Due to the lack of space-time solution, adjoint can not be enabled using the twin model. Gray-box simulators are ubiquitous in many engineering applications. Examples are ... for computational fluid dynamics, and ECLIPSE (Schlumberger), PSim (ConocoPhillips), and MORES (Shell) for petroleum reservoir simulations. My thesis will only investigate gray-box simulators.

My thesis aims at reducing the number of expensive iterations in the optimization constrained by gray-box simulators. Motivated by the adjoint gradient computation, a mathematical procedure will be developed to estimate the adjoint gradient by leveraging the full space-time solution. In addition, my thesis will investigate how the estimated gradient can faciliate a suitable optimization algorithm to reduce the number of iterations. Finally, the iteration reduction achieved by my approach will be assessed, especially for problems with many control parameters.

Instead of discussing gray-box simulators in general, my thesis only focuses on simulators with (partially) unknown flux function, while their boundary condition, initial condition, and the source term are known. ... a more precise definition of partially known ... This assumption is valid for some applications, such as simulating a petroleum reservoir with polymer injection. The flow in such reservoir is governed by multi-phase multi-component porous medium transport equations [4]. The initial condition is usually given at the equilibrium state, the boundary is usually described by a no-flux condition, and the source term can be modeled as controls with given flow rate or wellbore pressure. Usually the flux function is given by the Darcy's law. The Darcy's law involves physical models like the permeability and the viscosity². The mechanism that modifies the rock permeability and flow viscosity can be unavailable. Thereby the flux is partially unknown. The specific form of PDE considered in my thesis is given in Section 1.2. It is a future work to extend my research to more general gray-box settings where the initial condition, boundary condition, source term, and the flux are jointly unknown.

¹The permeability quantifies the easiness of liquids to pass through the rock.

²The viscosity quantifies the internal friction of the liquid flow.

1.2 Problem Formulation

Consider the optimization problem

$$c^* = \underset{c_{\min} \le c \le c_{\max}}{\operatorname{argmax}} \xi(\boldsymbol{u}, c)$$

$$\xi(\boldsymbol{u}, c) = \sum_{i=1}^{M} \sum_{j=1}^{N} w_{ij} f(\boldsymbol{u}_{ij}, c) \approx \int_{0}^{T} \int_{\Omega} f(u, c) d\boldsymbol{x} dt$$

$$(1.5)$$

where u is the discretized space-time solution of a gray-box conservation law simulator. The spatial coordinate is $x \in \Omega$ and the time is $t \in [0, T]$. $i = 1, \dots, M$ and $j = 1, \dots, N$ indicate the indices for the time and space discretization. w_{ij} 's are given quadrature weights for the space-time integration. $c \in \mathbb{R}^d$ indicates the control variable. c_{\min} and c_{\max} are elementwise bound constraints. The gray-box simulator solves the partial differential equation (PDE)

$$\frac{\partial u}{\partial t} + \nabla \cdot (DF(u)) = q(u, c), \qquad (1.6)$$

which is a system of k equation. The initial and boundary conditions are known. D is a known differential operator that may depend on u, and F is an unknown function that depends on u. q is a known source term that depends on u and c. Notice (1.6) degenerates to (1.1) when D equals 1. The simulator does not have the adjoint capability, and it is infeasible to implement the adjoint method into its source code. But the full space-time solution u is provided. The steady-state conservation law is a special case of the unsteady one, so it will not be discussed separately.

My thesis focuses on reducing the number of gray-box simulations in the optimization, especially for problems where d, the dimensionality of the control variable, is large. I assume that the computational cost is dominated by the repeated gray-box simulation, while the cost of optimization algorithm is relatively small. Chapter 2 develops a mathematical procedure, called the twin model method, that enables adjoint gradient computation by leveraging the full space-time solution. Based upon previous research

[55, 56, 60, 61, 63, 65, 66], Chapter 3 develops an optimization algorithm that takes advantage of the estimated gradient to achieve iteration reduction. The utility of the estimated gradient for optimization is analyzed both numerically and theoretically.

1.3 Literature Review

Given the background, I review the literature on derivative-free optimization and gradient-based optimization. In addition, I review the adjoint method since it is an essential ingredient for Chapter 2. Finally, I review model calibration and Bayesian optimization methods, which serve as the framework for the developments in Chapter 3.

1.3.1 Review of Optimization Methods

Optimization methods can be categorized into derivative-free and gradient-based methods [35], depending on whether the gradient information is used. In the sequel, I review the two types of methods.

Derivative-free Optimization

Derivative-free optimization (DFO) requires only the availability of objective function values but no gradient information [35], thus is useful when the gradient is unavailable, unreliable, or too expensive to obtain. Such methods are suitable for problems constrained by black-box simulators.

Depending on whether a local or global optimum is desired, DFO methods can be categorized into local methods and global methods [35]. Local methods seek a local optimum which is also the global optimum for convex problems. An important local method is the trust-region method [39]. Trust-region method introduces a surrogate model that is cheap to evaluate and presumably accurate within a trust region: an adaptive neighborhood around the current iterate [39]. At each iteration, the

surrogate is optimized in a domain bounded by the trust region to generate candidte steps for additional objective evaluations [39]. The surrogates can be constructed either by interpolating the objective evaluations [40, 43], or by running a low-fidelity simulation [41, 49]. Convergence to the objective function's optimum is guaranteed by ensuring that the surrogate have the same value and gradient as the objective function when the size of the trust region shrinks to zero [42, 43].

Global methods seek the global optimum. Example methods include the branchand-bound search [44], evolution methods [45], and Bayesian methods [60, 62, 82]. The branch-and-bound search sequentially partitions the entire control space into a tree structure, and determines lower and upper bounds for the optimum [44]. Partitions that are inferior are eliminated in the course of the search [44]. The bounds are usually obtained through the assumption of the Lipschitz continuity or statistical bounds for the objective function [44]. Evolution methods maintain a population of candidate controls, which adapts and mutates in a way that resembles natural phenomenons such as the natural selection [46, 48] and the swarm intelligence [47]. Bayesian methods model the objective function as a random member function from a stochastic process. At each iteration, the statistics of the stochastic process are calculated and the posterior, a probability measure, of the objective is updated using Bayesian metrics [60, 61]. The posterior is used to pick the next candidate step that best balances the exploration of unsampled regions and the exploitation around the sampled optimum [62, 71, 58]. Details of Bayesian optimization methods are discussed in Section 1.3.5.

Because many real-world problems are non-convex, global methods are usually preferred to local methods if the global optimum is desired [35]. Besides, DFO methods usually require a large number of function evaluations to converge, especially when the dimension of control is large [35]. This issue can be alleviated by incorporating the gradient information [55, 63, 78, 79]. The details are discussed in the next subsection.

Gradient-based Optimization

Gradient-based optimization (GBO) requires the availability of the gradient values [52, 72]. A gradient value, if exists, provides the optimal infinitesimal change of control variables at each iterate, thus is useful in searching for a better control. Similar to DFO, GBO can also be categorized into local methods and global methods [52]. Examples of local GBO methods include the gradient descent methods [73, 93, the conjugate gradient methods [74, 75], and the quasi-Newton methods [34, 36. The gradient descent methods and the conjugate gradient methods choose the search step in the direction of either the gradient [73, 93] or a conjugate gradient [74, 75]. Quasi-Newton methods, such as the Broyden-Fletcher-Goldfarb-Shannon (BFGS) method [34], approximate the Hessian matrix using a series of gradient values. The approximated Hessian allows a local quadratic approximation to the objective function which determines the search direction and stepsize by the Newton's method [34]. In addition, some local DFO methods can be enhanced to use gradient information [50, 51]. For instance, in trust-region methods, the construction of local surrogates can incorporate gradient values if available [50, 51]. The usage of gradient usually improves the surrogate's accuracy thus enhances the quality of the search step, thereby reducing the required number of iterations [50, 51].

Global GBO methods search for the global optimum using gradient values [52, 72]. Many global GBO methods can trace their development to corresponding DFO methods [76, 77, 78, 79, 63]. For example, the stochastic gradient-based global optimization method (StoGo) [76, 77] works by partitioning the control space and bounding the optimum in the same way as the branch-and-bound method [44]. But the search in each partition is performed by gradient-based algorithms such as BFGS [34]. Similarly, some gradient-based evolution methods, such as the gradient-based particle swarm method [78] and the gradient-based cuckoo search method [79], can be viewed as gradient variations of corresponding derivative-free counterparts [47, 48].

For example, the gradient-based particle swarm method combines particle swarm algorithm with the stochastic gradient descent method [78]. The movement of each particle is dictated not only by the function evaluations of all particles, but also by its local gradient [78].

My thesis is particularly interested in the gradient-based Bayesian optimization method [64]. In this method, the posterior of the objective function assimilates both the gradient and function values in a CoKriging framework [55, 64]. The details of my treatment is discussed in Section 1.3.5 and Chapter 3. I expect that the inclusion of gradient values results in more accurate posterior mean and reduced posterior uncertainty, which in turn reduces the number of iterations required to achieve near-optimality. The effect of iteration reduction is analyzed numerically in Chapter 3.

A property of the Bayesian method is that the search step can be determined using all available objective and gradient values [60, 71]. In addition, the search step is optimal under a Bayesian metric [60, 71]. The advantage of such properties can be justified when the objective and gradient evaluations are dominantly more expensive than the overhead of optimization algorithm [60]. Besides, my thesis proves that the Baysian optimization method is convergent even if the gradient values are estimated inexactly, which is discussed in Section 3.1.3. The conclusion of Section 3.1.3 is: Under some assumptions of the objective and the inexact gradient, a Bayesian optimization algorithm can find the optimum regardless of the accuracy of the gradient estimation.

To achieve a desired objective value, GBO methods generally require much less iterations than DFO methods for problems with many control variables [52, 72]. GBO methods can be efficiently applied to optimization constrained by open-box simulators, because the gradient is efficiently computable by the adjoint method [10, 52], which is introduced in the next subsection. My thesis extends GBO to optimization constrained by gray-box simulation by estimating the gradient using the full space-time solution.

1.3.2 The Adjoint Method

Consider a differentiable objective function constrained by a conservation law PDE (1.6). Let the objective function be $\xi(u,c)$, $c \in \mathbb{R}^d$, and let the PDE (1.6) be abstracted as $\mathcal{F}(u,c) = 0$. \mathcal{F} is a parameterized differential operator, together with boundary conditions and/or initial conditions, that uniquely defines a u for each c. The gradient $\frac{d\xi}{dc}$ can be estimated trivially by finite difference. The ith component of the gradient is given by

$$\left(\frac{d\xi}{dc}\right)_{i} \approx \frac{1}{\delta} \left(\xi(u + \Delta u_{i}, c + \delta e_{i}) - \xi(u, c)\right), \tag{1.7}$$

where

$$\mathcal{F}(u,c) = 0$$
, $\mathcal{F}(u + \Delta u_i, c + \delta e_i) = 0$. (1.8)

 e_i indicates the *i*th unit Cartesian basis vector in \mathbb{R}^d , and $\delta > 0$ indicates a small perturbation. Because (1.8) needs to be solved for every δe_i , so that the corresponding Δu_i can be used in (1.7), d+1 PDE simulations are required to evaluate the gradient. As explained in Section 1.3.1, d can be large in many control optimization problems. Therefore, it can be costly to evaluate the gradient by finite difference.

In contrast, the adjoint method evaluates the gradient using only one PDE simulation plus one adjoint simulation [10]. To see this, linearize $\mathcal{F}(u,c) = 0$ into a variational form

$$\delta \mathcal{F} = \frac{\partial \mathcal{F}}{\partial u} \delta u + \frac{\partial \mathcal{F}}{\partial c} \delta c = 0, \qquad (1.9)$$

which gives

$$\frac{du}{dc} = -\left(\frac{\partial \mathcal{F}}{\partial u}\right)^{-1} \frac{\partial \mathcal{F}}{\partial c} \tag{1.10}$$

Using (1.10), $\frac{d\xi}{dc}$ can be expressed by

$$\frac{d\xi}{dc} = \frac{\partial \xi}{\partial u} \frac{du}{dc} + \frac{\partial \xi}{\partial c}
= -\frac{\partial \xi}{\partial u} \left(\frac{\partial \mathcal{F}}{\partial u}\right)^{-1} \frac{\partial \mathcal{F}}{\partial c} + \frac{\partial \xi}{\partial c},
= -\lambda^{T} \frac{\partial \mathcal{F}}{\partial c} + \frac{\partial \xi}{\partial c} \tag{1.11}$$

where λ , the adjoint state, is given by the adjoint equation

$$\left(\frac{\partial \mathcal{F}}{\partial u}\right)^T \lambda = \left(\frac{\partial \xi}{\partial u}\right)^T \tag{1.12}$$

Therefore, the gradient can be evaluated by (1.11) using one simulation of $\mathcal{F}(u,c) = 0$ and one simulation of (1.12) that solves for λ .

Adjoint methods can be categorized into continuous adjoint and discrete adjoint methods, depending on whether the linearization or the discretization is excuted first [14]. The above procedure, (1.9) thru. (1.12), is the continuous adjoint, where \mathcal{F} is a differential operator. The continuous adjoint method linearizes the continuous PDE $\mathcal{F}(u,c) = 0$ first, then discretizes the adjoint equation (1.12) [10]. In (1.12), $\left(\frac{\partial \mathcal{F}}{\partial u}\right)^T$ can be derived as another differential operator. With proper boundary and/or initial conditions, it uniquely determines the adjoint solution λ . See [18] for a detailed derivation of the continuous adjoint equation.

The discrete adjoint method [16] discretizes $\mathcal{F}(u, c) = 0$ first. After the discretization, u and c become vectors \mathbf{u} and \mathbf{c} . \mathbf{u} is defined implicitly by the system $\mathcal{F}_d(\mathbf{u}, \mathbf{c}) = 0$, where \mathcal{F}_d indicates the discretized difference operator. Using the same derivation as (1.9) thru. (1.12), the discrete adjoint equation can be obtained

$$\left(\frac{\partial \mathcal{F}_d}{\partial \boldsymbol{u}}\right)^T \boldsymbol{\lambda} = \left(\frac{\partial \boldsymbol{\xi}}{\partial \boldsymbol{u}}\right)^T, \tag{1.13}$$

which is a linear system of equations. $\left(\frac{\partial \mathcal{F}_d}{\partial u}\right)^T$ is derived as another difference operator.

With proper discretized boundary/initial conditions, it uniquely determines the discrete adjoint vector λ , which subsequently determines the gradient

$$\frac{d\xi}{d\boldsymbol{c}} = -\boldsymbol{\lambda}^T \frac{\partial \mathcal{F}_d}{\partial \boldsymbol{c}} + \frac{\partial \xi}{\partial \boldsymbol{c}}.$$
 (1.14)

See Chapter 1 of [19] for a detailed derivation of the discrete adjoint.

The discrete adjoint method can be implemented by automatic differentiation (AD) [17]. AD exploits the fact that a PDE simulation, no matter how complicated, executes a sequence of elementary arithmetic operations (e.g. addition, multiplication) and elementary functions (e.g. exp, sin) [17]. For example, consider the function

$$\xi = f(c_1, c_2) = c_1 c_2 + \sin(c_1). \tag{1.15}$$

The function can be broken down into a series of elementary arithmetic operations and elementary functions.

$$w_{1} = c_{1}$$
 $w_{2} = c_{2}$
 $w_{3} = w_{1}w_{2}$
 $w_{4} = \sin(w_{1})$
 $\xi = w_{3} + w_{4}$.
$$(1.16)$$

(1.16) can be represented by a computational graph in Figure 1-1. In the graph, the gradient of the output with respect to the input variables can be computed using the chain rule [17]. Let \bar{z} denote the gradient of ξ with respect to z, for any independent or intermediate variable z in (1.16). To compute the derivatives $\bar{c}_1 = \frac{\partial \xi}{\partial c_1}$ and $\bar{c}_2 = \frac{\partial \xi}{\partial c_2}$,

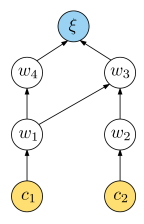


Figure 1-1: The computational graph for (1.16). The yellow nodes indicate the input variables, the blue node indicates the output variable, and the white nodes indicate the intermediate variables. The arrows indicate elementary operations. The beginning and end nodes of each arrow indicate the independent and dependent variables for each operation.

one can propogate the derivatives backward in the computational graph as follows

$$\bar{w}_{4} = 1$$

$$\bar{w}_{3} = 1$$

$$\bar{w}_{2} = \bar{w}_{3} \frac{\partial w_{3}}{\partial w_{2}} = 1 \cdot w_{1}$$

$$\bar{w}_{1} = \bar{w}_{4} \frac{\partial w_{4}}{\partial w_{1}} + \bar{w}_{3} \frac{\partial w_{3}}{\partial w_{1}} = 1 \cdot \cos(w_{1}) + 1 \cdot w_{2}$$

$$\bar{c}_{2} = \bar{w}_{2} = c_{1}$$

$$\bar{c}_{1} = \bar{w}_{1} = \cos(c_{1}) + c_{2}$$
(1.17)

The derivatives in (1.17) are straightforward to compute. This is because every forward operation in (1.16) is elementary, and their derivatives can be hardwired in AD softwares. Notice each arrow in Figure 1-1 is traversed for one and only one time in the backward propagation (1.17). Therefore, the backward gradient computation has a similar cost as the forward output computation, regardless of the number of input variables. See [17] for a thorough review of AD.

Because a PDE simulation can be viewed as performing a sequence of elementary

operations, AD can be used to evaluate the discrete adjoint. Consider a discretized PDE simulation that at each timestep solves

$$\mathcal{F}_{t+1} = \mathcal{F}(\mathbf{x}_t, \mathbf{x}_{t+1}, \mathbf{c}_{t+1}) = 0, \qquad (1.18)$$

for $t = 0, \dots, T - 1$, where \mathbf{x}_t and \mathbf{c}_t are the state and control variables at the tth timestep. AD can be used to compute the gradient of an objective function

$$\xi = \xi(\boldsymbol{x}_0, \cdots, \boldsymbol{x}_T; \boldsymbol{c}_1, \cdots \boldsymbol{c}_T)$$

to the control variables. To see this, consider the evaluation of (1.18) using an AD software. The gradients $\frac{\partial \mathcal{F}_{t+1}}{\partial \boldsymbol{x}_t}$, $\frac{\partial \mathcal{F}_{t+1}}{\partial \boldsymbol{x}_{t+1}}$, and $\frac{\partial \mathcal{F}_{t+1}}{\partial \boldsymbol{c}_{t+1}}$, for $t = 0, \dots, T-1$, are automatically computable. Therefore, one can obtain

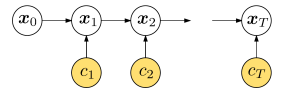
$$\frac{\partial \boldsymbol{x}_{t+1}}{\partial \boldsymbol{x}_{t}} = -\left(\frac{\partial \mathcal{F}_{t+1}}{\partial \boldsymbol{x}_{t+1}}\right)^{-1} \left(\frac{\partial \mathcal{F}_{t+1}}{\partial \boldsymbol{x}_{t}}\right)
\frac{\partial \boldsymbol{x}_{t+1}}{\partial \boldsymbol{c}_{t+1}} = -\left(\frac{\partial \mathcal{F}_{t+1}}{\partial \boldsymbol{x}_{t+1}}\right)^{-1} \left(\frac{\partial \mathcal{F}_{t+1}}{\partial \boldsymbol{c}_{t+1}}\right).$$
(1.19)

Therefore a computational graph, Figure 1-2a, can be constructed using the chain rule. The graph enables the evaluation of all $\frac{\partial x_t}{\partial c_{t-i}}$, for $t=1,\cdots,T$ and $i=0,\cdots,t-1$. Given the solutions x_t 's and the controls c_t 's, the evaluation of ξ is nothing but overlaying the graph by an additional layer of computations, shown in Figure 1-2b. Because $\frac{\partial \xi}{\partial x_t}$'s and $\frac{\partial \xi}{\partial c_t}$'s can be obtained by AD, the gradient

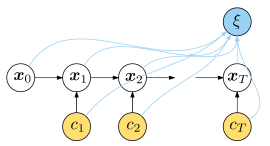
$$\frac{d\xi}{d\mathbf{c}_t} = \frac{\partial \xi}{\partial \mathbf{c}_t} + \frac{\partial \xi}{\partial \mathbf{x}_t} \frac{\partial \mathbf{x}_t}{\partial \mathbf{c}_t} + \frac{\partial \xi}{\partial \mathbf{x}_{t+1}} \frac{\partial \mathbf{x}_{t+1}}{\partial \mathbf{c}_t} + \dots + \frac{\partial \xi}{\partial \mathbf{x}_T} \frac{\partial \mathbf{x}_T}{\partial \mathbf{c}_t}$$
(1.20)

can be computed, for all $t = 1, \dots, T$.

The adjoint method has seen wide applications in optimization problems constrained by conservation law simulations, such as in airfoil design [11, 12, 13], adaptive mesh refinement [19], injection policy optimization in petroleum reservoirs [2], history matching in reservoir geophysics [14], and optimal well placement in reservoir management



(a) The computational graph for (1.18), which is constructed by (1.19). The yellow nodes indicate the input variables.



(b) The computational graph for evaluating the objective function ξ . The blue node indicates the output variable.

Figure 1-2: Computational graphs for the PDE simulation and objective evaluation.

[15]. Besides, there are many free AD softwares available for various languages, such as ADOL-C (C, C++) [20], Adiff (Matlab) [21], and Theano (Python) [22]. Unfortunately, the adjoint method is not directly applicable to gray-box simulations, as explained in Section 1.1. To break this limitation, Chapter 2 develops the twin model method that enables the adjoint gradient computation for gray-box simulations.

1.3.3 Adaptive Basis Construction

The unknown function F in (1.6) can be approximated by a linear combination of basis function. An over-complete or incomplete set of bases can negatively affects the approximation due to overfitting or underfitting. Therefore, adaptive basis construction is needed.

Square-integrable functions can be represented by the parameterization ³ [94]

$$\tilde{F}(\cdot) = \sum_{i \in \mathbb{N}} \alpha_i \phi_i(\cdot) , \qquad (1.21)$$

where ϕ_i 's are the basis functions, α_i 's are the coefficients, and i indices the basis. For example, a bivariate function can be represented by monomials (Weierstrass approximation theorem [94])

$$1, u_1, u_1^2, u_2, u_1u_2, u_1^2u_2, u_2^2, u_1u_2^2, u_1^2u_2^2, \cdots$$

on any real interval [a, b].

Let \mathcal{A} be a non-empty finite subset of \mathbb{N} , \tilde{F} can be approximated using a subset of the bases,

$$\tilde{F}(\cdot) \approx \sum_{i \in \mathcal{A}} \alpha_i \phi_i(\cdot) ,$$
 (1.22)

where $\{\phi_i\}_{i\in\mathcal{A}}$ is called a basis dictionary [25]. For example, in polynomial approximations, the basis dictionary can consists of the basis whose total polynomial degree does not exceed $p \in \mathbb{N}$ [23]. Given a basis dictionary, the coefficients can be determined by projection [23], such that the error between the approximation and the true function on a set of evaluation points is minimized. My thesis parameterizes the twin-model flux \tilde{F} and optimizes the coefficients, so the twin model serves as a proxy of the gray-box model. Details are discussed in Section 2.1.1.

$$\langle f, g \rangle = \int_A \overline{f(x)} g(x) dx,$$

where

- \bullet f and g are square integrable functions,
- $\overline{f(x)}$ is the complex conjugate of f(x), and
- A is the set over which the integral is defined.

It can be shown that square integrable functions is complete under the metric induced by the inner product, thus is a Hilbert space [94]. Any member function can be represented by a finite or countably infinite number of basis functions of the Hilbert space [94].

³By definition, the square integrable functions form a pre-Hilbert space with inner product given by

The cardinality of \mathcal{A} can increase drastically as the number of variables increases, and as the basis complexity (e.g. the total degree p of the polynomial basis) increases [23]. For example, if bound max degree of each variable, polynomial, exponential, if total degree, if Smolyak quadrature rule,

Fortunately, in many applications, many bases are insignificant so their coefficients can set to zero without affecting the approximation performance [25]. There are several methods that can determine the insignificant bases which result in a sparse representation, such as the Lasso regularization [37] and the matching pursuit [25]. Lasso regularization adds a penalty $\lambda \sum_{i \in \mathcal{A}} |\alpha_i|$ to the approximation error, where $\lambda > 0$ is a tunable parameter [37]. Therefore it balances the approximation error and the number of non-zero coefficients [37]. Matching pursuit adopts a greedy, stepwise approach to construct a basis [25]. It either selects a significant basis one-at-a-time (forward selection) [26], or prunes an insignificant basis one-at-a-time (backward pruning) [27]. Matching pursuit will be used in my thesis because it requires no tunable parameters.

In conventional matching pursuit, the basis dictionary needs to be pre-determined, with the belief that the dictionary is a superset of the significant basis [29]. This can be problematic because the significant ones are unknown a prior. To address this issue, adaptive methods have been devised [28, 29, 30]. Although different in details, such methods share the same heuristics: Starting from a trivial basis (e.g. 1), a dictionary is built up progressively by iterating over a forward step and a backward step [28, 29, 30]. The forward step searches over a candidate set of bases, and appends the significant ones to the dictionary [28, 29, 30]. The backward step searches over the current dictionary, and removes the insignificant ones from the dictionary [28, 29, 30]. The iteration stops when no alternation is made to the dictionary or a targeted approximation performance is met [28, 29, 30]. Such heuristics are adopted in my thesis to build up a sparse representation of \tilde{F} . Details are discussed in Section 2.1.2.

1.3.4 Model Calibration

Model is not reality. A computer simulation may not reproduce experimental results exactly due to several sources of uncertainties, including parameter uncertainty, model inadequacy, code uncertainty, and observation error [66]. Parameter uncertainty refers to uncertain parameters of a model [66]. Consider an example in the polymerinjected petroleum reservoir. The parameters in the permeability models and viscosity models, measured in laboratory conditions, may not equal the true parameters in reservoir conditions. Parameter uncertainty can be eliminated if the true parameters are known and specified in a simulation [66]. Model inadequacy relates to the fact that no model is perfect [66]. The model output differs from experimental results due to underlying missing physics and numerical approximations, even if the model's parameter uncertainty is eliminated [66]. For example, in the same petroleum reservoir simulation, model inadequacy would exist if the compressibility is neglected, or if the flux is approximated by numerical schemes like upwinding. Code uncertainty refers to the unknown output of a computer simulation given any particular control variables and model parameters until the code is run [66]. In theory, code uncertainty disappears if the model is simulated densely over its entire input domain [66]. In practice, however, the simulation output can only be interpolated on a limited number of simulated results since the code can be expensive to run. Observation error is the uncertainty associated with experimental measurements [66]. This error represents the variability in the experimental response if the experiment were repeated multiple times under the exact same settings [66].

Model calibration addresses the problem of adjusting the model parameters and quantifying the uncertainties [66]. An important approach for model calibration, proposed by Kennedy and O'Hagan, adopts a Bayesian modeling for the observation, the code output, and the uncertainties [66, 67]. Represent the observation by z and the code output by y. y not only depends on the input control variables c but also on the model parameters θ . The relationship between the observation and the code

output is modeled by

$$z_i = \rho y(c_i, \theta) + \epsilon(c_i) + \delta_i, \qquad (1.23)$$

where z_i is the *i*th observation, ρ is an unknown regression parameter, $\epsilon(\cdot)$ is the model inadequacy that depends on c, and δ_i is the observation error for the *i*th observation [66]. δ_i 's are assumed to be i.i.d. Gaussian distributed, which models the idiosyncratic observation error for individual experiments [66]. $\epsilon(\cdot)$ accounts for the model inadequacy, which is assumed to be an unknown member of a Gaussian process [66]. The Gaussian process model expresses a belief in the smoothness of underlying function, and offers the flexibility in adjusting the smoothness by tuning hyperparameters like the correlation length [66]. To incorporate the code uncertainty, $y(\cdot, \cdot)$ is modeled by a joint Gaussian process that distributes on c and θ [66]. Usually, $y(\cdot, \cdot)$ is assumed to be independent of $\epsilon(\cdot)$ [66]. I refer to the reference [68] for a justification of the independence assumption. The parameter uncertainty is accounted for by assuming a distribution on θ [66]. Denote the hyperparameters in these Gaussian processes by ϕ . A joint posterior distribution

$$p(\theta, \rho, \phi | \mathcal{D}) \propto p(\theta, \rho, \phi) p(\mathcal{D} | \theta, \rho, \phi)$$
 (1.24)

can be obtained, given the prior $p(\theta, \rho, \phi)$ and a set of observations $\mathcal{D} = (c_i, z_i)_{i=1}^N$. Thereby, the model is calibrated by the posterior distribution of θ , ρ , and δ , as well as the posterior Gaussian processes of y and ϵ .

The model calibration method is applicable not only when z_i 's are observed experimentally, but also when z_i 's are simulated by another computer model [65]. A goal in my thesis is to estimate the true gradient by the twin model's adjoint gradient, as well as to quantify the uncertainty in such estimation. The goal can be achieved by model calibration, where z and y in (1.23) indicate the true gradient and the estimated gradient. My thesis calibrates the twin model by using the space-time solution of deterministic gray-box simulations. Deterministic simulations have no observation error; therefore this error is not considered in my thesis. Besides, instead

of jointly calibrating the parameters and uncertainties using a Bayesian approach, the model parameters are calibrated by the twin model method developed in Chapter 2. The twin model method provides a point estimate of the model parameters. Once the point estimate is determined, the model inadequacy and the code uncertainty are calibrated in the next step while fixing the model parameters as known constants. It is a future work to extend the developments to joint calibration.

1.3.5 Bayesian Optimization

Similar to other kinds of optimization, Bayesian optimization aims at finding the maximum of a function $\xi(\cdot)$ in a bounded set $\mathcal{C} \subset \mathbb{R}^d$ [60, 61, 71]. However, Bayesian optimization distinguishes from other methods by maintaining a probabilistic model for ξ [60, 61, 71]. The probabilistic model is exploited to make decisions about where to invest the next function evaluation in \mathcal{C} [60, 61, 71]. In addition, it uses all information of available evaluations, not just local evaluations, to direct the search step [60, 61, 71].

Consider the case when the objective function evaluation is available. Bayesian optimization begins by assuming that the objective function is sampled from a stochastic process [60, 61, 71]. A stochastic process is a function

$$f: \mathcal{C} \times \Omega \to \mathbb{R}$$

 $(c, \omega) \to f(c, \omega)$, (1.25)

where for any $c \in \mathcal{C}$, $f(c,\cdot)$ is a random variable defined on the probability space $(\Omega, \Sigma, \mathbb{P})$. The objective function ξ is assumed to be a sample function from the stochastic process $\xi(\cdot) = f(\cdot, \omega^*)$ with an unknown $\omega^* \in \Omega$. My thesis will use the notations $\xi(\cdot)$, $f(\cdot, \omega)$, and $f(\cdot, \omega^*)$ interchangeably when the context is clear.

Stationary Gaussian process is a stochastic process that is used ubiquitously in

Bayesian optimization [80]. For any given ω and any finite set of N points $\{c_i \in \mathcal{C}\}_{i=1}^N$, a stationary Gaussian process $f(\cdot,\cdot)$ has the property that $\{f(c_i,\cdot)\}_{i=1}^N$ are multivariate Gaussian distributed; in addition, the distribution remains unchanged if c_i 's are all added by the same constant in \mathcal{C} . The Gaussian process is solely determined by its mean function m(c) and its covariance function K(c,c') [80]

$$m(c) = \mathbb{E}_{\omega} [f(c, \omega)]$$

$$K(c, c') = \mathbb{E}_{\omega} [(f(c, \omega) - m(c))(f(c', \omega) - m(c'))],$$
(1.26)

for any $c, c' \in \mathcal{C}$, which is denoted by $f \sim \mathcal{N}(m, K)$. Conditioned on a set of samples $\{\xi(c_1), \dots, \xi(c_N)\}$, the posterior is also a Gaussian process with the mean and covariance [80]

$$\tilde{m}(c) = m(c) + K(c, \underline{c}_n) K(\underline{c}_n, \underline{c}_n)^{-1} \left(\xi(\underline{c}_n) - m(\underline{c}_n) \right),$$

$$\tilde{K}(c, c') = K(c, c') - K(c, \underline{c}_n) K(\underline{c}_n, \underline{c}_n)^{-1} K(\underline{c}_n, c')$$

$$(1.27)$$

where $\underline{c}_n = (c_1, \dots, c_N)$, $\xi(\underline{c}_n) = (\xi(c_1), \dots, \xi(c_N))^T$, $m(\underline{c}_n) = (m(c_1), \dots, m(c_N))^T$, $K(c, \underline{c}_n) = K(\underline{c}_n, c)^T = (K(c, c_1), \dots, K(c, c_N))$, and

$$K(\underline{c}_n, \underline{c}_n) = \begin{pmatrix} K(c_1, c_1) & \cdots & K(c_1, c_N) \\ \vdots & \ddots & \vdots \\ K(c_N, c_1) & \cdots & K(c_N, c_N) \end{pmatrix}.$$

Without prior knowledge about the underlying function, $m(\cdot)$ is usually modeled as a constant independent of c [80]. In many cases, the covariance are assumed isotropic, indicating that K(c, c') only depends on the L_2 norm ||c - c'|| [80]. There are many choices for K, such as the exponential kernel, the squared exponential kernel, and the Matérn kernels, each embeds different degrees of smoothness (differentiability) for the underlying function. For a survey of various covariance functions, I refer to

the Chapter 4 in [80]. Among such choices, the Matérn 5/2 kernel [81]

$$K(c,c') = \sigma^2 \left(1 + \frac{\sqrt{5}\|c - c'\|}{L} + \frac{5\|c - c'\|^2}{3L^2} \right) \exp\left(-\frac{\sqrt{5}\|c - c'\|}{L} \right), \quad (1.28)$$

has been recommended because it results in functions that are twice differentiable, an assumption made by, e.g. quasi-Newton methods, but without further smoothness [60]. My thesis will focus on using the Matérn 5/2 kernel. Notice the parameters L and σ , known as the hyperparametes, are yet to be determined. They can be determined by the posterior maximum likelihood estimation (MLE) or by a fully-Bayesian approach [60, 71]. I refer to the reference [60] for the details and a comparison of these treatments. My thesis will focus on MLE due to its simpler numerical implementation.

Based on the posterior and the current best evaluation $c_{\text{best}} = \operatorname{argmax}_{c \in c_n} \xi(c)$, Bayesian optimization introduces an acquisition function, $a : \mathcal{C} \to \mathbb{R}^+$, that evaluates the expected utility of investing the next sample at $c \in \mathcal{C}$ [58, 60, 61, 71, 82]. The location of the next sample is determined by an optimization $c_{N+1} = \operatorname{argmax}_{c \in \mathcal{C}} a(c)$ [58, 60, 61, 71, 82]. In most cases, a greedy acquisition function is used, which evaluates the one-step-lookahead utility [58, 60, 61, 71, 82]. There are several choices for the acquisition function, such as

• the probability of improvement (PI) [82],

$$a_{\rm PI}(c) = \Phi(\gamma(c)), \qquad (1.29)$$

• the expected improvement (EI) [61, 62],

$$a_{\rm EI}(c) = \sigma(c) \left(\gamma(c) \Phi(\gamma(c)) + \mathcal{N}(\gamma(c)) \right), \tag{1.30}$$

• and the upper confidence bound (UCB) [58],

$$a_{\text{UCB}}(c) = \mu(c) + \kappa \sigma(c), \qquad (1.31)$$

with a tunable parameter $\kappa > 0$,

where μ, σ are the posterior mean and variance, $\gamma(c) = \sigma^{-1}(c) (\mu(c) - \xi(c_{\text{best}}))$, and Φ, \mathcal{N} indicate the cumulative and density functions for the standard normal distribution. My thesis will focus on the EI acquisition function, as it behaves better than the PI, and requires no extra tunable parameters [60]. Because (1.30) has a closed-form gradient, the acquisition function can be maximized by a global GBO method, e.g. StoGo [77], to obtain its global maximum.

Although my thesis only focuses on bound constraints as shown in (1.5), Bayesian optimization can accommodate more general inequality and equality constraints [88]. The constraints can be enforced by modifying the objective, such as the penalty method [83], the augmented Lagrangian method [84], and the barrier function method [85]. They can also be enforced by modifying the acquisition function, such as the recently developed expected improvement with constraints (EIC) method [86], and the integrated expected conditional improvement (IECI) method [87]. See Chapter 2 of [88] for a detailed review of constrained Bayesian optimization.

In addition to function evaluations $\xi(\underline{c}_n)$, Bayesian optimization admits gradient information [55, 63]. In Chapter 3, I investigate the scenario where the gradient evaluations are inexact, possibly corrupted by model inadequacy and observation error [66]. The Bayesian optimization method developed in my thesis allows both the exact function evaluation and the inexact gradient evaluation. Details of this topic will be discussed in Section 3.1.

1.4 Notations

The general notations are declared here.

- $t \in [0, T]$: the time,
- $\{t_i\}_{i=1}^M$: the time discretization,
- $x \in \Omega$: the space,
- $\{x_j\}_{j=1}^N$: the space discretization,
- u: the space-time solution of gray-box conservation law,
- \tilde{u} : the space-time solution of twin-model conservation law,
- u: the discretized space-time solution of gray-box simulator,
- $\tilde{\boldsymbol{u}}$: the discretized space-time solution of twin-model simulator,
- k: 1) the number of equations of the conservation law; or 2) the number of folds in cross validation.
- D: a differential operator,
- F: the unknown function of the gray-box model,
- \tilde{F} : the inferred F,
- q: the source term,
- c: the control variables,
- $\underline{c}_n = (c_1, \dots, c_n)$: a sequence of n control variables,
- w: the quadrature weights in the numerical space-time integration,
- ξ : the objective function,
- c_{\min} , c_{\max} : bound constraints,

- $\xi_{\tilde{\nabla}}$: the estimated gradient of ξ with respect to c,
- d: the number of control variables,
- $\mathcal{C} \subset \mathbb{R}^d$: the control space,
- K, G: the covariance functions,
- a: the acquisition function,
- \mathcal{M} : the solution mismatch,
- $\overline{\mathcal{M}}$: the mean solution mismatch in cross validation,
- ϕ : the basis functions for \tilde{F} ,
- α : the coefficients for ϕ ,
- A: the index set for a basis dictionary,
- T: twin model,
- τ : residual,
- τ : discretized residual,
- \mathcal{T} : integrated truncation error.

1.5 Thesis Objectives

Based the motiviation and literature review, we find a need to enable adjoint gradient computation for gray-box conservation law simulations. We also need to exploit the estimated gradient to optimize more efficiently, especially for problems with many control variables. To summarize, the objectives of my thesis are

to develop an adjoint approach that estimates the gradient of objective functions
constrained by gray-box conservation law simulations with unknown flux functions,
by leveraging the space-time solution;

- 2. to assess the utility of the estimated gradient in a suitable gradient-based optimization method; and
- 3. to demonstrate the effectiveness of the developed procedure in several numerical examples, given a limited computational budget.

1.6 Outline

In Chapter 2, a solution mismatch metric is presented. The metric is used for the training of twin models. In addition, an adaptive basis construction scheme is developed. By summarizing the developments, a twin model algorithm is presented. Furthermore, a truncation error metric and a pre-train step is developed in order to reduce the computational cost. Finally, the twin model algorithm is demonstrated in several numerical examples. In Chapter 3, the twin-model gradient is modeled stochastically by the Gaussian process. Based on the Gaussian process model, a Bayesian optimization algorithm is devised that leverages the twin-model gradient. Its convergence properties are studied. Finally, the twin-model Bayesian optimization algorithm is demonstrated in several numerical examples. Chapter 3 summarizes the thesis and my contributions, and proposes several interesting future works.

Chapter 2

Estimate the Gradient by Using the Space-time Solution

This chapter develops a method to estimate the gradient by using the space-time solution of gray-box conservation law simulations. An example, equation (1.1), has been given in Section 1.1 to illustrate why such estimation is possible when the conservation law involves only one equation and one dimensional space. In this example, the derivative of the flux can be first interpolated by using the discretized gray-box solution; then the adjoint method is applied to the inferred conservation law (1.4) to estimate the gradient. This chapter develops a more general procedure suitable for systems of equations and for problems with a spatial dimension greater than one.

2.1 Approach

Consider a gray-box simulator that solves the PDE (1.6), a system of k equations, for u(t,x) with $t \in [0,T]$ and $x \in \Omega$. The PDE has an unknown flux F, but known source term q, and known initial and boundary conditions. Let its discretized space-time solution be u. My thesis introduces an open-box simulator solving another PDE,

namely the twin model,

$$\frac{\partial \tilde{u}}{\partial t} + \nabla \cdot \left(D\tilde{F}(\tilde{u}) \right) = q(\tilde{u}, c) , \qquad (2.1)$$

which is a system of k equations with the same source term and the same initial and boundary conditions. Equation (2.1) differs from (1.6) in its flux. For simplicity, let the open-box simulator use the same space-time discretization, and let its discretized solution be $\tilde{\boldsymbol{u}}$. Define the solution mismatch

$$\mathcal{M}(\tilde{F}) = \sum_{i=1}^{M} \sum_{j=1}^{N} w_{ij} (\tilde{\boldsymbol{u}}_{ij} - \boldsymbol{u}_{ij})^{2}, \qquad (2.2)$$

where w_{ij} 's are the quadrature weights for the space-time integration. \mathcal{M} approximates the space-time integration of the continous solutions' mismatch,

 $\mathcal{M} \approx \int_0^T \int_{\Omega} (\tilde{u}(t,x) - u(t,x))^2 dx dt$. Given a set \mathcal{S}_F consisted of all possible guesses for F, I propose to infer a flux \tilde{F} such that the mismatch between \boldsymbol{u} and $\tilde{\boldsymbol{u}}$ is minimized, i.e.

$$\tilde{F}^* = \operatorname*{argmin}_{\tilde{F} \in \mathcal{S}_F} \mathcal{M} \,, \tag{2.3}$$

The choice for S_F will be discussed later in Section 2.1.1 and 2.1.2. Because the twin model is open-box, (2.3) can be solved by gradient-based methods. Once \mathcal{M} is minimized, the adjoint method can be applied to the twin model to estimate the gradient.

The key to inferring the flux is to leverage the gray-box space-time solution. Its inferrability can be loosely explained by the following reasonings. Firstly, the conserved quantity u in (1.6) depends on u in a previous time only inside a domain of dependence, illustrated in Figure 2-1. Similarly, in discretized PDE simulation, the solution at any gridpoint only depends on a numerical domain of dependence. Besides, in a PDE simulation, a one-step time marching at any gridpoint can be viewed as a mapping whose input only involves the numerical domain of dependence. Because the number of state variables in the numerical domain of dependence can be small,

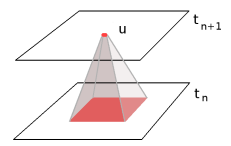


Figure 2-1: Domain of dependence: u(t,x) depends on u at an earlier time within a domain of dependence. The two planes in this figure indicates the spatial solution at two adjacent timesteps. The domain of dependence can be much smaller than Ω , the entire spatial domain.

inference of the mapping is potentially feasible. Secondly, the space-time solution at every space-time gridpoint can be viewed as a sample for this mapping. Because the scale of space-time discretization in a conservation law simulation is usually large, a large number of samples are available for the inference, thus making the inference potentially accurate.

The inferrability is difficult to prove. However, it can be partially justified by the following theorem for PDEs with k=1 and one dimensional space.

Theorem 1. Consider two PDEs

$$\frac{\partial u}{\partial t} + \frac{\partial F(u)}{\partial x} = 0$$
, and (2.4)

$$\frac{\partial \tilde{u}}{\partial t} + \frac{\partial \tilde{F}(\tilde{u})}{\partial x} = 0, \qquad (2.5)$$

with the same initial condition $u(0,x) = u_0(x)$, and $x \in \mathbb{R}$. u_0 is bounded, differentiable, Lipschitz continuous with constant L_u , and has a finite support. F and \tilde{F} are both twice-differentiable and Lipschitz continuous with constant L_F . Let

$$B_u \equiv \left\{ u \left| u = u_0(x) \text{ that satisfies } \left| \frac{du_0}{dx} \right| \ge \gamma > 0, \text{ for all } x \in \mathbb{R} \right. \right\} \subseteq \mathbb{R}.$$

be a non-empty and measurable set. We have:

For any $\epsilon > 0$, there exist $\delta > 0$ and T > 0 such that

• if $|\tilde{u}(t,x) - u(t,x)| < \delta$ for any $x \in \mathbb{R}$ and $t \in [0,T]$, then $\left| \frac{d\tilde{F}}{du} - \frac{dF}{du} \right| < \epsilon$ for any $u \in B_u$.

The proof is given in Appendix A.1. An illustration of B_u is given in Figure 2-2. Several observations can be made from Theorem 1. Firstly, if the solutions of (2.4) and (2.5) match closely, then the derivatives of their flux functions must match closely in B_u . Secondly, the conclusion can only be drawn for $u \in B_u$ where the initial condition has coverage and has large enough variation. For more general problems involving 1) systems of equations, 2) higher spatial dimensions, and 3) discretization, the inferrability is difficult to show theoretically. Instead, it will be demonstrated numerically.

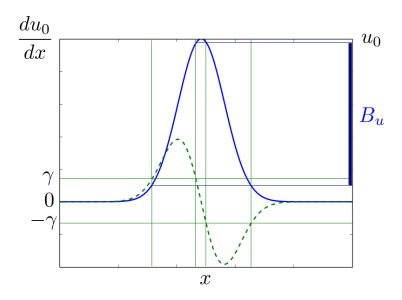


Figure 2-2: An illustration of B_u defined in Theorem 1. The blue line is u_0 and the green dashed line is $\frac{du_0}{dx}$. B_u is the set of u_0 where the derivative $\frac{du_0}{dx}$ has an absolute value larger than γ .

The remainder of this chapter is organized as follows. Section 2.1.1 discusses the choices of S_F occurred in (2.3). A suitable parameterization for \tilde{F} will be chosen that takes into account the observations from Theorem 1. Section 2.1.2 develops a procedure that adaptively refines the parameterization. The procedure adopts a

greedy search of the best refinement at every iterate and refrains from overfitting. Section 2.1.3 summarizes the algorithm for training the twin model. Section 2.1.4 presents a numerical shortcut for (2.3) that is more computationally efficient. Section 2.2 demonstrates the algorithm in several numerical examples. Finally, Section 2.3 summarizes the chapter.

2.1.1 Parameterization

Functions can be parameterized by a linear combination of basis functions. Firstly, consider the case when \tilde{F} is univariate. There are many types of basis functions to parameterize a univariate function, such as polynomial basis, Fourier basis, and wavelet basis. Based on the observations from Theorem 1, \tilde{F} and F are expected to match only on a domain of u where the gray-box space-time solution exists and has large variation. Besides, \tilde{F} may match F better on a domain where the gray-box discretized solution u are more densely sampled. Therefore, an ideal parameterization should admit local refinements so \tilde{F} can match F better at some domain; similarly, it should allow local dropouts when bases are redundant at some domain. This section presents a choice of the parameterization for \tilde{F} that allows such local refinements, while a procedure for basis refinement is given in the next section.

A parameterization that allows local refinements and local dropouts is the wavelet. Wavelet is the basis developed for multi-resolution analysis (MRA) [91]. MRA is an increasing sequence of closed function spaces $\{V_j\}_{j\in\mathbb{Z}}$,

$$\cdots \subset V_{-1} \subset V_0 \subset V_1 \subset \cdots$$

which can approximate functions with increasing resolutions as j increases. For

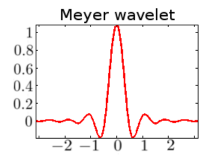


Figure 2-3: An example of mother wavelet, the Meyer wavelet.

univariate MRA, V_j 's satisfy the following properties known as self-similarity:

$$f(u) \in V_j \Leftrightarrow f(2u) \in V_{j+1}, \ j \in \mathbb{Z}$$

 $f(u) \in V_j \Leftrightarrow f(u - \frac{\eta}{2^j}) \in V_j, \ j \in \mathbb{Z}, \ \eta \in \mathbb{Z}$

The wavelet bases for V_j are given by

$$\hat{\phi}_{j,\eta}(u) = 2^{j/2} \hat{\phi}(2^j u - \eta), \quad \eta \in \mathbb{Z}$$
(2.6)

where $\hat{\phi}$ is called the mother wavelet satisfying $\hat{\phi}(u) \to 0$ for $u \to -\infty$ and ∞ . An example mother wavelet, the Meyer wavelet, is shown in Figure 2-3.

Because the gray-box model is unchanged by adding a constant to F, the derivative of \tilde{F} , instead of \tilde{F} itself, should be approximated. Thereby, the bases for \tilde{F} shall be chosen as the integrals of the wavelets, i.e.

$$\phi_{j,\eta}(u) = \int_{-\infty}^{u} \hat{\phi}_{j,\eta}(u')du'. \qquad (2.7)$$

 $\phi_{j,\eta}$'s are sigmoid functions which satisfy

$$\phi_{j,\eta}(u) = \begin{cases} 0, u \to -\infty \\ 1, u \to \infty \end{cases}$$
(2.8)

It's easy to show $\phi_{j,\eta}$ also satisfies self-similarity, and can be represented by a mother

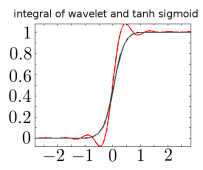


Figure 2-4: Red line: the integral (2.7) of the Meyer wavelet. Black line: the logistic sigmoid function.

sigmoid ϕ ,

$$\phi_{j,\eta}(u) = \phi(2^j u - \eta) \,, \quad j \in \mathbb{Z} \,, \, \eta \in \mathbb{Z}$$
(2.9)

There are many types of sigmoid functions. My thesis will use the logistic sigmoid function as the mother sigmoid,

$$\phi(u) = \frac{1}{1 + e^{-u}} \,. \tag{2.10}$$

If \tilde{F} is univariate, the logistic sigmoids $\phi_{j,\eta}$'s are used as the bases. If \tilde{F} is multivariate, the basis can be formed by the tensor product of univariate sigmoids,

$$\phi_{j,\eta}(u_1,\dots,u_k) = \phi_{j_1,\eta_1}(u_1)\dots\phi_{j_k,\eta_k}(u_k),$$
 (2.11)

where $\boldsymbol{j}=(j_1,\cdots,j_k)\in\mathbb{Z}^k,\,\boldsymbol{\eta}=(\eta_1,\cdots,\eta_k)\in\mathbb{Z}^k.$ To sum up,

$$\tilde{F}(\cdot) = \sum_{j,\eta} \alpha_{j,\eta} \phi_{j,\eta}(\cdot), \qquad (2.12)$$

where α 's are the coefficients of the bases. A systematic procedure for choosing a set of $\{j, \eta\}$ is presented in Section 2.1.2.

For illustration, consider a numerical example where the gray-box model solves

the 1-D Buckley-Leverett equation [3]

$$\frac{\partial u}{\partial t} + \frac{\partial}{\partial x} \left(\underbrace{\frac{u^2}{1 + 2(1 - u)^2}}_{F} \right) = c, \qquad (2.13)$$

with the initial condition $u(0,x) = u_0(x)$ and the periodic boundary condition u(t,0) = u(t,1). Let $0 \le u_0(x) \le 1$ for all $x \in [0,1]$. This is because the Buckley-Leverett equation models the two-phase porous media flow where u stands for the saturation of a phase, and the saturation is always positive and no larger than one. $c \in \mathbb{R}$ is a constant-valued control. F is assumed unknown and is inferred by a twin model. The twin model solves

$$\frac{\partial \tilde{u}}{\partial t} + \frac{\partial}{\partial x} \tilde{F}(\tilde{u}) = c, \qquad (2.14)$$

with the same c and the same initial and boundary conditions. \tilde{F} is parameterized by (2.12) where j's and η 's are chosen ad hoc. Figure 2-5 gives an example of the bases used in this section. To ensure the well-posedness of (2.3), an ad hoc L_1 regularization on α is used in minimizing \mathcal{M} .

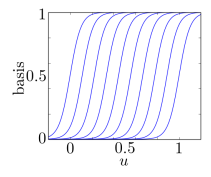


Figure 2-5: An ad hoc set of bases

Figure 2-6a shows the discretized space-time solution of (2.13) for $x \in [0, 1]$, $t \in [0, 1]$ and c = 0. The solution is used to train a twin model according to (2.3). The discretized space-time solution of the trained twin model is shown in Figure 2-6b.

Once a twin model is trained, its adjoint can be used for gradient estimation.

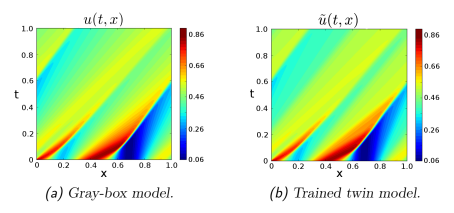


Figure 2-6: Space time solutions.

Consider an objective function

$$\xi(c) \equiv \int_{x=0}^{1} \left(u(1, x; c) - \frac{1}{2} \right)^{2} dx.$$
 (2.15)

Its gradient $\frac{d\xi}{dc}$ can be estimated by the trained twin model. Figure 2-7 shows the objective function, evaluated using the gray-box model and the trained twin model. It is observed that the gradients of ξ match closely at c=0, i.e. the control where the twin model is trained.

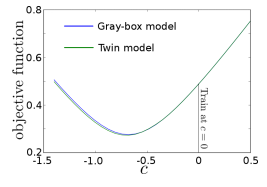


Figure 2-7: The objective function ξ evaluated by either the gray-box model or the trained twin model.

In addition to the gradient estimation, the inferred \tilde{F} is examined. If different solutions are used in the training, it is expected that the trained twin model will also be different. Figure 2-8 shows the training results for three different initial conditions.

Some observations can be made: 1) As expected, \tilde{F} can differ from F by a constant without affecting \mathcal{M} ; 2) $\frac{d\tilde{F}}{du}$ matches $\frac{dF}{du}$ only in a domain of u where the solution exists (indicated by the green area); 3) Sometimes the bases seem redundant thus can be safely dropped out. The issue seems particularly important in the third column, where most bases are suppressed; 4) Sometime the bases seem too coarse thus may be refined in order to reduce the minimal \mathcal{M} . The issue seems particularly important in the first column, where $\frac{d\tilde{F}}{du}$ exhibits a wavy deviation from $\frac{dF}{du}$. Addressing these issues systematically is crucial to the rigorous development of the twin model method.

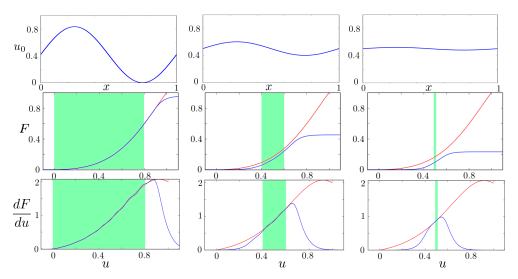


Figure 2-8: The first row shows the three different initial conditions used to generate the gray-box space-time solution. The second row compares the trained \tilde{F} (blue) and the Buckley-Leverett F (red). The third row compares the trained $\frac{d\tilde{F}}{du}$ (blue) and the Buckley-Leverett $\frac{dF}{du}$ (red). The green background highlights the domain of u where the gray-box space-time solution exists.

2.1.2 Elements for Adaptive Basis Construction

This section develops several key elements that lead to the adaptive basis construction for twin models. The heuristics for the adaptive basis construction, discussed in Section 1.3.3, are applied to build up a basis dictionary consisted of only the significant candidates. The section is organized as follows: Firstly, a formulation is provided to

efficiently assess the significance of each candidate basis; Secondly, the neighborhood of a sigmoid basis is defined; Thirdly, a metric is devised that determines when to add or remove a candidate basis. The three elements are then employed to build the twin model algorithm in Section 2.1.3.

Given a basis dictionary $\phi_{\mathcal{A}} = \{\phi_i\}_{i \in \mathcal{A}}$, define the "minimal mismatch"

$$\mathcal{M}^*(\mathcal{A}) = \min_{\alpha_{\mathcal{A}} \in \mathbb{R}^{|\mathcal{A}|}} \mathcal{M}\left(\sum_{i \in \mathcal{A}} \alpha_i \phi_i\right), \qquad (2.16)$$

to be the minimal solution mismatch (2.2) if \tilde{F} were parameterized by $\phi_{\mathcal{A}}$. \mathcal{A} is a set containing $\{j, \eta\}$'s. $\alpha_{\mathcal{A}} = \{\alpha_i\}_{i \in \mathcal{A}}$ is the coefficient for $\phi_{\mathcal{A}}$. Let $\alpha_{\mathcal{A}}^* = \{\alpha_i^*\}_{i \in \mathcal{A}}$ be the optimal coefficients, and let $\tilde{F}_{\mathcal{A}}^* = \sum_{i \in \mathcal{A}} \alpha_i^* \phi_i$. Consider appending $\phi_{\mathcal{A}}$ by an additional basis ϕ_l , and let $\phi_{\mathcal{A}'} = \{\phi_{\mathcal{A}}, \phi_l\}$, $\mathcal{A}' = \{\mathcal{A}, l\}$. The minimal mismatch for the appended basis dictionary $\phi_{\mathcal{A}'}$ is

$$\mathcal{M}^*(\mathcal{A}') = \min_{\alpha_{\mathcal{A}'} \in \mathbb{R}^{|\mathcal{A}|+1}} \mathcal{M}\left(\sum_{i \in \mathcal{A}'} \alpha_i \phi_i\right) , \qquad (2.17)$$

Clearly $\mathcal{M}^*(\mathcal{A}') \leq \mathcal{M}^*(\mathcal{A})$. Define the "mismatch improvement" to be

$$\Delta \mathcal{M}^* (\mathcal{A}, l) = \mathcal{M}^* (\mathcal{A}) - \mathcal{M}^* (\mathcal{A}')$$
 (2.18)

Approximate (2.18) by Taylor expansion, we get

$$\Delta \mathcal{M}^* (\mathcal{A}, l) \approx -\left(\int_{u \in \mathbb{R}^k} \frac{d\mathcal{M}}{d\tilde{F}} \Big|_{\tilde{F}_{A}^*} \phi_l \, du \right) \alpha_l, \qquad (2.19)$$

where $\frac{d\mathcal{M}}{d\tilde{F}}$ is the derivative of $\mathcal{M}(\tilde{F})$ with respect to \tilde{F} , evaluated on $\tilde{F} = \tilde{F}_{\mathcal{A}}^*$. For a twin model consisted of a system of k equations, \tilde{F} is a function of $u \in \mathbb{R}^k$, thus $\frac{d\mathcal{M}}{d\tilde{F}}$ is also a function of $u \in \mathbb{R}^k$. As discussed in the previous sections, $\frac{d\mathcal{M}}{d\tilde{F}}$ is non-zero only in a domain where there is solution. Thus (2.19) can be integrated by quadrature

only over a bounded domain. The absolute value of the coefficient for α_l ,

$$s_l(\mathcal{A}) \equiv \left| \int_{u \in \mathbb{R}^k} \frac{d\mathcal{M}}{d\tilde{F}} \right|_{\tilde{F}_{\mathcal{A}}^*} \phi_l \, \mathrm{d}u \right|,$$
 (2.20)

estimates the significance of the basis ϕ_l [31]. If there are multiple candidate bases, (2.20) can be used to rank their significance.

In the sequel, a compact representation of the sigmoid bases is introduced. The univariate basis function, $\phi_{j,\eta}$ in (2.9), is represented by a tuple $(j, \frac{\eta}{2^j})$, where j can be viewed as the "resolution", and $\frac{\eta}{2^j}$ is the center of the basis. Similarly, the k-variate basis function, $\phi_{j,\eta}$ in (2.11), is represented by a tuple $(j, \frac{\eta}{2^j}) = (\{j_1, \dots, j_k\}, \{\frac{\eta_1}{2^{j_1}}, \dots, \frac{\eta_k}{2^{j_k}}\})$. Thus, a sigmoid can be represented by a point in a 2k-dimensional space. The representation is illustrated in Figure 2-9a thru. 2-9d for the univariate case.

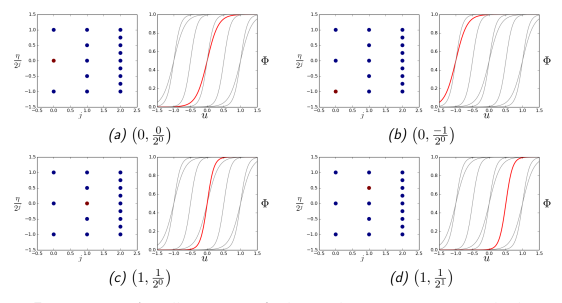


Figure 2-9: An illustration of the tuple representation and the corresponding univariate sigmoid.

Using this representation, define the "neighborhood" of a univariate sigmoid $(j, \frac{\eta}{2^j})$ to be

$$\mathcal{N}\left[\left(j, \frac{\eta}{2^{j}}\right)\right] = \left\{\left(j+1, \frac{\eta}{2^{j}}\right), \left(j, \frac{\eta \pm 1}{2^{j}}\right)\right\}. \tag{2.21}$$

The neighborhood contains 1) a basis $(j+1,\frac{\eta}{2^j})$ with an increment of resolution; and 2) two basis $(j,\frac{\eta\pm 1}{2^j})$ with the same resolution but a marginal shift of center. For illustration, the neighborhood of $(0,\frac{0}{2^0})$ is shown in Figure 2-10a. Similarly, define the neighborhood of a multivariate sigmoid to be

$$\mathcal{N}\left[\left(j,\frac{\eta}{2^{j}}\right)\right] = \mathcal{N}\left[\left(\left\{j_{1},\cdots,j_{k}\right\},\left\{\frac{\eta_{1}}{2^{j_{1}}},\cdots,\frac{\eta_{k}}{2^{j_{k}}}\right\}\right)\right] \\
= \left\{\left(\left\{j_{1}+1,\cdots,j_{k}\right\},\left\{\frac{\eta_{1}}{2^{j_{1}+1}},\cdots,\frac{\eta_{k}}{2^{j_{k}}}\right\}\right)\cdots,\left(\left\{j_{1},\cdots,j_{k}+1\right\},\left\{\frac{\eta_{1}}{2^{j_{1}}},\cdots,\frac{\eta_{k}}{2^{j_{k}+1}}\right\}\right), \\
\left(\left\{j_{1},\cdots,j_{k}\right\},\left\{\frac{\eta_{1}\pm1}{2^{j_{1}}},\cdots,\frac{\eta_{k}}{2^{j_{k}}}\right\}\right)\cdots,\left(\left\{j_{1},\cdots,j_{k}\right\},\left\{\frac{\eta_{1}}{2^{j_{1}}},\cdots,\frac{\eta_{k}\pm1}{2^{j_{k}}}\right\}\right)\right\}, \\
(2.22)$$

which consists of k bases with incremental resolution, and 2k bases with center shifts. It is easy to see that a basis $(\boldsymbol{j}_0, \frac{\eta_0}{2^{j_0}})$ can be connected to any basis $(\boldsymbol{j}, \frac{\eta}{2^{j}})$ with $\boldsymbol{j} \geq \boldsymbol{j}_0$ through a chain of neighborhoods. In addition, define the neighborhood of multiple sigmoid functions to be the union of each individual's neighborhood, as illustrated by Figure 2-10b.

$$\mathcal{N}\left[(\boldsymbol{j}_1, \frac{\boldsymbol{\eta}_1}{2^{\boldsymbol{j}_1}}), \cdots, (\boldsymbol{j}_n, \frac{\boldsymbol{\eta}_n}{2^{\boldsymbol{j}_n}})\right] = \mathcal{N}\left[(\boldsymbol{j}_1, \frac{\boldsymbol{\eta}_1}{2^{\boldsymbol{j}_1}})\right] \bigcup \cdots \bigcup \mathcal{N}\left[(\boldsymbol{j}_n, \frac{\boldsymbol{\eta}_n}{2^{\boldsymbol{j}_n}})\right]. \tag{2.23}$$

Although the mismatch improvement, $\Delta \mathcal{M}^*(\mathcal{A}, l)$, is always non-negative, it is inadvisable to cram the basis dictionary with too many bases, otherwise a twin model can be overfitted. Therefore, a criterion is required to determine if a candidate basis shall be added to or removed from the basis dictionary. This can be achieved by cross validation, in particular, k-fold cross validation [32]. Given a basis dictionary, the k-fold cross validation proceeds in the following three steps: Firstly, the gray-box solution u is shuffled randomly into k disjoint sets $\{u_1, u_2, \dots, u_k\}$. An illustration for k = 2 is shown in Figure 2-11.

Secondly, k twin models who share the same basis dictionary are trained so their space-time solutions match all but one sets, shown in (2.24). T_i indicates the ith twin

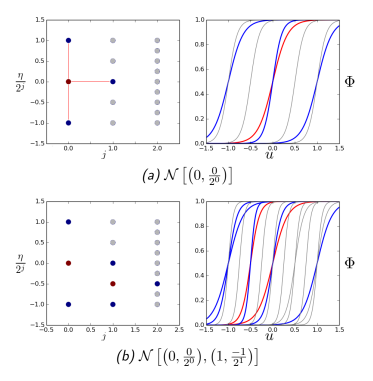


Figure 2-10: Neighborhood for univariate bases. (a) shows the neighborhood (blue) of a single basis (red). (b) shows the neighborhood (blue) of several bases (red). The left column represents the basis on the $(j, \frac{\eta}{2^j})$ plane, and the right column shows the actual basis $\phi_{j,\eta}$.

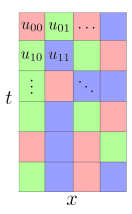


Figure 2-11: The discretized gray-box solution is shuffled into 3 sets, each indicated by a color.

model.

$$T_1 = \texttt{TrainTwinModel}(\boldsymbol{u}_2, \boldsymbol{u}_3, \cdots, \boldsymbol{u}_k)$$

$$T_2 = \texttt{TrainTwinModel}(\boldsymbol{u}_1, \boldsymbol{u}_3, \cdots, \boldsymbol{u}_k)$$

$$\cdots$$

$$T_k = \texttt{TrainTwinModel}(\boldsymbol{u}_1, \boldsymbol{u}_2, \cdots, \boldsymbol{u}_{k-1})$$

$$(2.24)$$

Thirdly, each trained twin model is validated on the remaining set. In particular, the solution mismatch for the validation set is computed, as shown in (2.25).

$$\mathcal{M}_1 = exttt{MismatchValidation} \left(T_1, oldsymbol{u}_1
ight) \ \mathcal{M}_2 = exttt{MismatchValidation} \left(T_2, oldsymbol{u}_2
ight) \ \ldots \ \mathcal{M}_k = exttt{MismatchValidation} \left(T_k, oldsymbol{u}_k
ight) \ \end{cases}$$

The mean value of validation errors,

$$\overline{\mathcal{M}} = \frac{1}{k} \left(\mathcal{M}_1 + \mathcal{M}_2 + \dots + \mathcal{M}_k \right) \tag{2.26}$$

measures the performance of the basis dictionary. A basis shall be added to or removed from the dictionary only if such action reduces $\overline{\mathcal{M}}$. In practice, cross validation proliferates the computational cost. Therefore, a small k is preferrable if cost is a concern. All the numerical examples in the thesis use k=2.

2.1.3 Algorithm

Based upon the developments in the previous sections, a twin model algorithm with adaptive basis construction is devised.

```
Input: Initial basis dictionary \phi_{\mathcal{A}}, coefficients \alpha_{\mathcal{A}} = \mathbf{0}, Validation error \overline{\mathcal{M}}_0 = \infty,
           Gray-box solution u.
   1: Minimize solution mismatch \alpha_{\mathcal{A}} \leftarrow \operatorname{argmin}_{\alpha} \mathcal{M} \left( \sum_{i \in \mathcal{A}} \alpha_i \phi_i \right)
   2: loop
                             Find \phi_l \in \mathcal{N}(\phi_A) \setminus \phi_A with the maximal s_l(A)
   3:
                             \mathcal{A} \leftarrow \mathcal{A} \bigcup \{l\}, \, \phi_{\mathcal{A}} \leftarrow \phi_{\mathcal{A}} \bigcup \{\phi_{l}\}, \, \alpha_{l} = 0, \, \alpha_{\mathcal{A}} \leftarrow \{\alpha_{\mathcal{A}}, \alpha_{l}\}
                             Compute \overline{\mathcal{M}} by k-fold cross validation.
   4:
                             if \overline{\mathcal{M}} < \overline{\mathcal{M}}_0 then
   5:
                                                \overline{\mathcal{M}}_0 \leftarrow \overline{\mathcal{M}}
   6:
                                                \alpha_{\mathcal{A}} \leftarrow \operatorname{argmin}_{\alpha} \mathcal{M} \left( \sum_{i \in \mathcal{A}} \alpha_i \phi_i \right)
                             else
   7:
                                               \mathcal{A} \leftarrow \mathcal{A} \setminus \{l\}, \ \phi_{\mathcal{A}} \leftarrow \phi_{\mathcal{A}} \setminus \{\phi_{l}\}, \ \alpha_{\mathcal{A}} \leftarrow \alpha_{\mathcal{A}} \setminus \{\alpha_{l}\} \ \mathbf{break}
   8:
                             end if
   9:
                             Find \phi_{l'} \in \phi_{\mathcal{A}} with the least s_{l'}(\mathcal{A})
10:
                             if l' \neq l then
11:
                                                \mathcal{A} \leftarrow \mathcal{A} \setminus \{l'\}, \ \phi_{\mathcal{A}} \leftarrow \phi_{\mathcal{A}} \setminus \{\phi_{l'}\}, \ \alpha_{\mathcal{A}} \leftarrow \alpha_{\mathcal{A}} \setminus \{\alpha_{l'}\}
12:
                                                Compute \overline{\mathcal{M}} by k-fold cross validation.
13:
                                                if \overline{\mathcal{M}} < \overline{\mathcal{M}}_0 then
14:
                                                                   \overline{\mathcal{M}}_0 \leftarrow \overline{\mathcal{M}}
15:
                                                                   \alpha_{\mathcal{A}} \leftarrow \operatorname{argmin}_{\alpha} \mathcal{M} \left( \sum_{i \in \mathcal{A}} \alpha_i \phi_i \right)
                                                else
16:
                                                                   \mathcal{A} \leftarrow \mathcal{A} \bigcup \{l'\}, \ \phi_{\mathcal{A}} \leftarrow \phi_{\mathcal{A}} \bigcup \{\phi_{l'}\}, \ \alpha_{\mathcal{A}} \leftarrow \alpha_{\mathcal{A}} \bigcup \{\alpha_{l'}\}
17:
                                                end if
18:
19:
                             end if
20: end loop
Output: \mathcal{A}, \phi_{\mathcal{A}}, \alpha_{\mathcal{A}}.
```

Algorithm 1 adopts the heuristics of the forward-backward iteration discussed in Section 1.3.3. The algorithm starts from training a twin model using a simple basis dictionary. Usually the starting dictionary contains one basis for each dimension

Algorithm 1: Training twin model with adaptive basis construction.

with very low resolution. Details of the choice are given in Section 2.2 along with numerical examples. The main part of the algorithm iterates over a forward step (line 3-9) and a backward step (line 10-19). The forward step firstly finds the most promising candidate in the neighborhood of the current dictionary for addition, according to (2.20). If the addition indeed reduces the cross validation error, the candidate is appended to the dictionary; otherwise it is rejected. If the basis is appended, the coefficients are updated by minimizing the solution mismatch, which can be implemented by the Broyden-Fletcher-Goldfarb-Shannon (BFGS) algorithm. The backward step finds the most promising candidate in the current dictionary for deletion. If the deletion reduces the cross validation error, the candidate is removed from the dictionary. If the basis is deleted, the coefficients are updated by BFGS again. The iteration exits when the most promising addition no longer reduces the validation error. In the end, the algorithm provides the basis dictionary and its coefficients as the output.

The algorithm requires to train multiple twin models at each iteration. For k=2, 6 twin models are trained if both the forward and the backward step are acceptive. In practice, the trained coefficients at the last iterate usually provide good initial guess for the next iterate. Nonetheless, the algorithm can be costly if the dictionary turns out to have a high cardinality which results in a large number of iterates before the dictionary construction completes. Therefore, a numerical shortcut is provided in Section 2.1.4 that significantly reduces the cost.

2.1.4 Minimizing the Truncation Error

In the previous sections, a twin model is trained to minimize the solution mismatch. The training can be expensive. Because the minimization of the solution mismatch, coupled with the adaptive basis construction, can require a large number of solution mismatch evaluations, and each evaluation involves one twin model simulation. To reduce the computational cost, a "pre-training" step is proposed where an "integrated

truncation error" is minimized. A pre-trained twin model is then "fine tuned" to minimize the solution mismatch. The applicability of the pre-training is studied; in particular, I study under what condition can the solution mismatch be bounded by the integrated truncation error. Finally, a stochastic gradient descent approach is adopted that efficiently minimizes the integrated truncation error.

Define

$$\tau = \frac{\partial u}{\partial t} + \nabla \cdot \left(D\tilde{F}(u) \right) - q(u, c) , \qquad (2.27)$$

which is the residual if the gray-box PDE's solution is plugged in the twin-model PDE (2.1). Let its discretization be τ . For simplicity, assume the gray-box simulator and its twin model use the same space-time discretization. τ can be obtained by plugging the discretized gray-box solution in the twin model simulator. Define the integrated truncation error to be

$$\mathcal{T}(\tilde{F}) = \sum_{i=1}^{M} \sum_{j=1}^{N} w_{ij} \tau_{ij}^{2}, \qquad (2.28)$$

where w_{ij} are the same quadrature weights as in (2.2). i, j are the indices for time and space discretization as in the previous sections. I propose to pre-train a twin model using Algorithm 1 with \mathcal{M} replaced by \mathcal{T} . In other words, in the pre-training step, the coefficients are determined by

$$\alpha_{\mathcal{A}} \leftarrow \underset{\alpha}{\operatorname{argmin}} \mathcal{T} \left(\sum_{i \in \mathcal{A}} \alpha_i \phi_i \right) .$$
 (2.29)

Besides, the estimator for the significance of a candidate basis, $s_l(\mathcal{A})$, is replaced by

$$s_l^t(\mathcal{A}) \equiv \left| \int_{u \in \mathbb{R}^k} \frac{d\mathcal{T}}{d\tilde{F}} \right|_{\tilde{F}_{\mathcal{A}}^*} \phi_l \, \mathrm{d}u \right| .$$
 (2.30)

Finally, the validation error, $\overline{\mathcal{M}}$, is replaced by

$$\overline{\mathcal{T}} = \frac{1}{k} \left(\mathcal{T}_1 + \mathcal{T}_2 + \dots + \mathcal{T}_k \right) , \qquad (2.31)$$

where

$$\mathcal{T}_i = \text{IntegratedTruncationError}(T_i, \boldsymbol{u}_i).$$
 (2.32)

for $i = 1, \dots, k$. Using the pre-trained basis dictionary $\phi_{\mathcal{A}}^t$, the twin model is then fine tuned by minimizing the solution mismatch, where $\alpha_{\mathcal{A}}^t$ is used as the initial guess and is adjusted according to (2.3). For a simulation with implicit schemes, the residual and the integrated truncation error are cheaper to evaluate than the solution mismatch, thereby the benefit of the pre-train.

However, \mathcal{M} may not be bounded by \mathcal{T} . A sufficient condition under which the bound exists is provided by Theorem 2.

Theorem 2. Consider a twin model simulator whose one-step time marching is

$$\mathcal{G}_i: \mathbb{R}^N \mapsto \mathbb{R}^N, \ \tilde{\boldsymbol{u}}_{i\cdot} \to \tilde{\boldsymbol{u}}_{i+1\cdot} = \mathcal{G}_i \tilde{\boldsymbol{u}}_{i\cdot}, \quad i = 1, \cdots, M-1.$$
 (2.33)

Assume the quadrature weights are time-independent, i.e. $w_{ij} = w_j$ for all i, j. If \mathcal{G}_i satisfies

$$\|\mathcal{G}_i a - \mathcal{G}_i b\|_W^2 \le \beta \|a - b\|_W^2, \qquad (2.34)$$

with $\beta < 1$, for any $a, b \in \mathbb{R}^N$ and for all i, then

$$\mathcal{M} \le \frac{1}{1-\beta} \mathcal{T}, \tag{2.35}$$

where

$$||v||_W^2 \equiv v^T \begin{pmatrix} w_1 & & \\ & \ddots & \\ & & w_N \end{pmatrix} v \tag{2.36}$$

for any $v \in \mathbb{R}^N$.

The proof is given in Appendix A.2. If the twin model is a contractive dynamical system [33], as given by (2.34), then the solution mismatch can be bounded by

the integrated truncation error. In contrast, the bound may not exist for non-contractive dynamical systems, for example for systems that exhibit bifurcation. It is a future work to further investigate the applicability of the pre-training theoretically, in particular, to investigate the necessary and sufficient condition for the bound. My thesis will explore the usefulness of the pre-training by several numerical test cases.

Because the residual τ can be evaluated explicitly given the gray-box solution, the evaluation can be decoupled for different space-time grid points $\{i, j\}$. By viewing the truncation error at each $\{i, j\}$ as a stochastic sample, (2.29) can be solved by stochastic gradient descent, Algorithm 2.

```
Input: \alpha = \alpha_0

1: for (i, j) = (1, 1) to (M, N) do

2: if not converged then

3: \alpha \leftarrow \alpha - \lambda w_{ij} \frac{\partial}{\partial \alpha} \tau_{ij}

4: else

5: break

6: end if

7: end for

Output: \alpha
```

Algorithm 2: Minimizing the integrated truncation error by stochastic gradient descent.

 $\lambda > 0$ is a tunable step size. Usually λ is tuned by experience to increase convergence speed while avoiding divergence. In practice, it is beneficial to compute the gradient against more than one grid points (called a "mini-batch") at each iterate. This is because the code can take advantage of vectorization libraries rather than computing the residual at each grid point separately.

2.2 Numerical Results

This section demonstrates the twin model on the estimation of the gradients for several numerical examples.

2.2.1 Buckley-Leverett Equation

Section 2.1.1 has applied a sigmoid parameterization to the gray-box model governed by the Buckley-Leverett equation (2.13). In this section, the same problem is studied but using the adaptive basis construction developed in Section 2.1.3 and 2.1.4. The initial dictionary, ϕ_A , is selected to contain a single basis $(0, \frac{0}{2^0})$. Clearly the choice is not unique. As long as the initial basis has a low resolution and is centered around $[u_{\min}, u_{\max}]$, Algorithm 1 shall build the dictionary adaptively.

Figure 2-12 shows the selected bases for the three solutions in Figure 2-8, respectively, by using the pre-train step. As $[u_{\min}, u_{\max}]$ shrinks, the dictionary's cardinality reduces and the resolution increases.

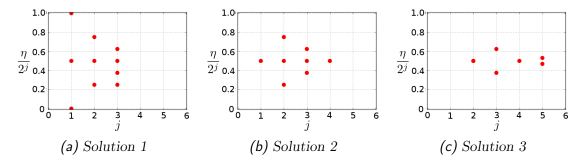


Figure 2-12: The basis dictionary for the three solutions in Figure 2-8.

Consider a time-space-dependent control c = c(t, x) in (2.13) and (2.14). The gradient of ξ , (2.15), is estimated using the trained twin model. The estimated gradients are compared with the true adjoint gradients of the gray-box model, and the errors are shown in Figure 2-13.

The adaptive basis construction improves the accuracy of the gradient estimation. Table 2.1 shows the integrated gradient error ¹ by using either the ad hoc bases in Figure 2-5 or by using the bases constructed adaptively.

¹The gradient error is integrated using the same quadrature rule as in (2.2).



Figure 2-13: The errors of estimated gradients for the three solutions.

	Solution 1	Solution 2	Solution 3
Ad hoc basis	2.5×10^{-3}	6.6×10^{-4}	7.3×10^{-5}
Adaptive basis	4.2×10^{-6}	1.5×10^{-6}	8.9×10^{-7}

Table 2.1: The integrated errors of the estimated gradients for the three solutions.

2.2.2 Navier-Stokes Flow

Consider a compressible internal flow in a 2-D return bend channel driven by the pressure difference between the inlet and the outlet. The return bend is bounded by no-slip walls. The inlet static pressure and the outlet pressure are fixed. The geometry of the return bend is given in Figure 2-14. The inner and outer boundaries of the bending section are each generated by 6 control points using quadratic B-spline.

The flow is governed by Navier-Stokes equations. Let ρ , u, v, E, and p denote the density, Cartesian velocity components, total energy, and pressure. The steady-state Navier-Stokes equation is

$$\frac{\partial}{\partial x} \begin{pmatrix} \rho u \\ \rho u^2 + p - \sigma_{xx} \\ \rho uv - \sigma_{xy} \\ u(E\rho + p) - \sigma_{xx}u - \sigma_{xy}v \end{pmatrix} + \frac{\partial}{\partial y} \begin{pmatrix} \rho v \\ \rho uv - \sigma_{xy} \\ \rho v^2 + p - \sigma_{yy} \\ v(E\rho + p) - \sigma_{xy}u - \sigma_{yy}v \end{pmatrix} = \mathbf{0}, \quad (2.37)$$

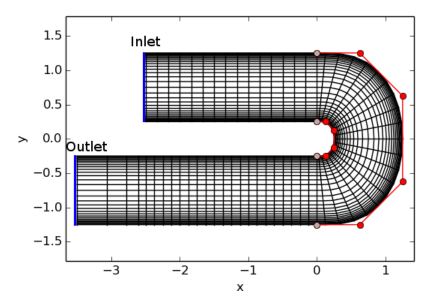


Figure 2-14: The return bend geometry and the mesh for the simulation.

where

$$\sigma_{xx} = \mu \left(2 \frac{\partial u}{\partial x} - \frac{2}{3} \left(\frac{\partial u}{\partial x} + \frac{\partial v}{\partial y} \right) \right)$$

$$\sigma_{yy} = \mu \left(2 \frac{\partial v}{\partial y} - \frac{2}{3} \left(\frac{\partial u}{\partial x} + \frac{\partial v}{\partial y} \right) \right).$$

$$\sigma_{xy} = \mu \left(\frac{\partial u}{\partial y} + \frac{\partial v}{\partial x} \right)$$

$$(2.38)$$

The Navier-Stokes equation requires an additional equation, the state equation, for closure. The state equation has the form

$$p = p(U, \rho), \qquad (2.39)$$

where U denotes the internal energy per unit volume,

$$U = \rho \left(E - \frac{1}{2} (u^2 + v^2) \right) . \tag{2.40}$$

Many models have been developed for the state equation, such as the ideal gas equation, the van der Waals equation, and the Redlich-Kwong equation [92]. In the sequel, the true state equation in the gray-box simulator is assumed unknown and

will be inferred from the gray-box solution. Let ρ_{∞} be the steady state density, $\mathbf{u}_{\infty} = (u_{\infty}, v_{\infty})$ be the steady state Cartesian velocity, and E_{∞} be the steady state energy density. The steady state mass flux is

$$\xi = -\int_{\text{outlet}} \rho_{\infty} u_{\infty} \big|_{\text{outlet}} dy = \int_{\text{inlet}} \rho_{\infty} u_{\infty} \big|_{\text{inlet}} dy$$
 (2.41)

The goal is to estimate the gradient of ξ to the red control points' coordinates.

Two state equations are tested: the ideal gas equation and the Redlich-Kwong equation, given by

$$p_{ig} = (\gamma - 1)U$$

$$p_{rk} = \frac{(\gamma - 1)U}{1 - b_{rk}\rho} - \frac{a_{rk}\rho^{5/2}}{((\gamma - 1)U)^{1/2}(1 + b_{rk}\rho)}$$
(2.42)

where $a_{rk} = 10^7$ and $b_{rk} = 0.1$.

The solution mismatch, (2.2), is given by

$$\mathcal{M} = w_{\rho} \int_{\Omega} \left| \tilde{\rho}_{\infty} - \rho_{\infty} \right|^{2} d\boldsymbol{x} + w_{u} \int_{\Omega} \left| \tilde{u}_{\infty} - u_{\infty} \right|^{2} d\boldsymbol{x} + w_{v} \int_{\Omega} \left| \tilde{v}_{\infty} - v_{\infty} \right|^{2} d\boldsymbol{x} + w_{E} \int_{\Omega} \left| \tilde{E}_{\infty} - E_{\infty} \right|^{2} d\boldsymbol{x} ,$$

where w_{ρ} , w_{u} , w_{v} , and w_{E} are non-dimensionalization constants. Figure 2-15 shows the gray-box solution and the solution mismatch after training the twin model ². Figure 2-16 compares the true state equation and the corresponding trained state equation, where the convex hull of $(U_{\infty}, \rho_{\infty})$, the internal energy and the density of the gray-box solution, is shown by the dashed red line. Because the state equation is expected to be inferrable only inside the domain of the gray-box solution, large deviation is expected outside the convex hull.

²Both the solution and the mismatch are normalized.

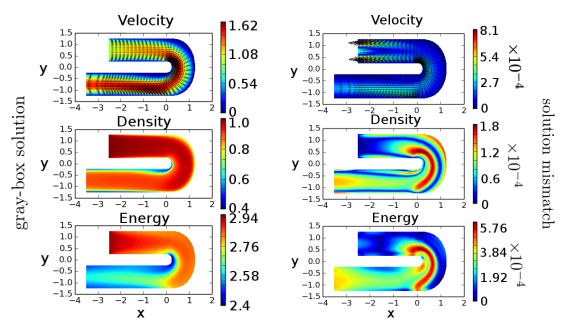


Figure 2-15: Left column: an example gray-box solution for a given geometry. Right column: the solution mismatch after training a twin model.

The trained twin model enables the adjoint gradient estimation. Figure 2-17 shows the estimated gradient of ξ with respect to the control points coordinates. It also compares the estimated gradient with the true gradient. The two gradients are indistinguishable, and the error is given in Table 2.2.

Gas	Interior control points			Exterior control points				
Ideal	0.13	0.04	0.05	0.32	0.16	0.15	0.07	0.02
Redlich-Kwong	0.32	0.03	0.07	0.50	0.40	0.12	0.06	0.05

Table 2.2: The error of the gradient estimation, in percentage.

2.2.3 Polymer Injection in Petroleum Reservoir

Water flooding is a technique to enhance the secondary recovery in petroleum reservoirs, as illustrated in Figure 2-18. Injecting pure water can be cost-inefficient due to low water viscosity and high water cut. Therefore, water-solvent polymer can be utilized

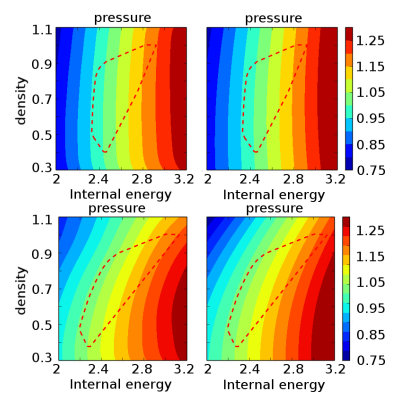
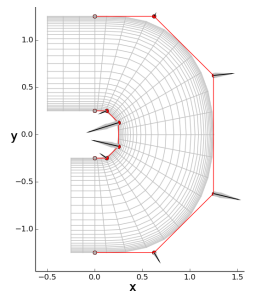
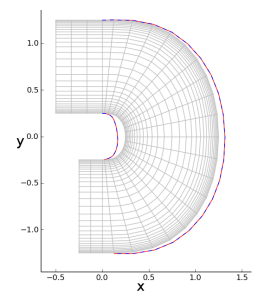


Figure 2-16: The gray-box state equation (right column) and the trained state equation (left column). The gray-box model uses either the ideal gas equation (first row) or the Reclich-Kwong equation (second row). The convex hull of the gray-box solution is shown by the dashed red line.





(a) The gradient of ξ to the control points for the Redlich-Kwong gas. The wide gray arrow is the gradient evaluated by the gray-box model, while the thin black arrow is the gradient evaluated by the twin model using finite difference.

(b) The boundary perturbed according to the gradient. The blue dashed line is computed by finite difference of the graybox model, while the red dashed line is computed by the twin model's gradient.

Figure 2-17: A comparison of the estimated gradient and the true gradient.

to increase the water-phase viscosity and to reduce the residual oil.



Figure 2-18: Water flooding in petroleum reservoir engineering (courtesy from PetroWiki).

Consider a reservoir governed by the two-phase porous media flow equations

$$\frac{\partial}{\partial t} (\rho_{\alpha} \phi S_{\alpha}) + \nabla \cdot (\rho_{\alpha} \vec{v}_{\alpha}) = 0, \quad \alpha \in \{w, o\},
\frac{\partial}{\partial t} (\rho_{w} \phi S_{w} c) + \nabla \cdot (c \rho \vec{v}_{wp}) = 0$$
(2.43)

for $x \in \Omega$ and $t \in [0,T]$, where the phase velocities are given by the Darcy's law

$$\vec{v}_{\alpha} = -M_{\alpha}k_{r\alpha}\mathbf{K} \cdot (\nabla p - \rho_{w}g\nabla z), \quad \alpha \in \{w, o\}$$

$$\vec{v}_{wp} = -M_{wp}k_{rw}\mathbf{K} \cdot (\nabla p - \rho_{w}g\nabla z)$$

$$(2.44)$$

w, o indicate the water and oil phases. ρ is the phase density. ϕ is the porosity. S is the phase saturation where $S_w + S_o = 1$. c is the polymer concentration in the water phase. v_w , v_o , v_{wp} are the componentwise velocities of water, oil, and polymer. K is the permeability tensor. k_r is the relative permeability. p is the pressure. z is the depth. g is the gravity constant. The mobility factors, M_o , M_w , M_{wp} , model the modification of the componentwise mobility due to the presence of polymer. In the sequel, the models for the mobility factors are unknown. The only knowledge about the mobility factors is that they depend on S_w , p, and c.

PSim, the simulator aforementioned in Section 1.1, is used as the gray-box simulator,

which uses the IMPES time marching, i.e. implicit in pressure and explicit in saturation, as well as the upwind scheme. Its solution, S_w , c, and p can be used to train the twin model. The twin model uses fully implicit time marching and the upwind scheme. The solution mismatch is defined by

$$\mathcal{M} = w_{S_w} \int_0^T \int_{\Omega} |S_w - \tilde{S}_w|^2 d\boldsymbol{x} dt + w_c \int_0^T \int_{\Omega} |c - \tilde{c}|^2 d\boldsymbol{x} dt + w_p \int_0^T \int_{\Omega} |p - \tilde{p}|^2 d\boldsymbol{x} dt,$$
(2.45)

where w_{S_w} , w_c , and w_p are non-dimensionalization constants.

Consider a reservoir setup shown in Figure 2-19, which is a 3D block with two injectors and one producer. The permeability is 100 milli Darcy, and the porosity is 0.3. A constant injection rate of $10^6 \text{ft}^3/\text{day}$ is used at both the injectors. The reservoir is simulated for $t \in [0, 50] \text{day}$. The solution of S_w is illustrated in Figure 2-20 for the untrained twin model, the gray-box model, and the trained twin model, respectively. After the training, the twin-model solution matches the gray-box solution closely.



Figure 2-19: The geometry of the petroleum reservoir.

Let the objective function be the residual oil at $T=50\,\mathrm{day},$

$$\xi = \int_{\Omega} \rho_o(T) \phi S_o(T) \, \mathrm{d}\boldsymbol{x} \,. \tag{2.46}$$



Figure 2-20: The isosurfaces of $S_w=0.25$ and $S_w=0.7$ at t=30 days.

The gradient of ξ with respect to the time-dependent injection rate is computed. The gradient estimated by the twin model is shown in Figure 2-21, where the red and blue lines indicate the gradient for the two injectors. In comparison, the star markers show the true gradient at day 2, 16, 30, and 44, evaluated by finite difference. Clearly, a rate increase at the injector 1 leads to more residual oil reduction than the injector 2. This is because the injector 2 is closer to the producer, where a larger rate accelerates the water breakthrough that impedes further oil production. It is observed that the estimated gradient closely matches the true gradient, although the error slightly increases for smaller t, possibly because of the different numerical schemes used in the twin and gray-box models. The error is given in Table 2.3.

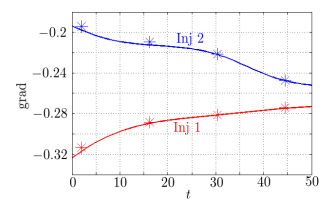


Figure 2-21: The gradient of ξ with respect to rates at the two injectors. The lines indicate the gradients estimated by the twin model, while the stars indicate the true gradient evaluated by finite difference.

Error	t = 0.04	t = 0.32	t = 0.6	t = 0.88
Inj 1	1.7	1.0	0.6	0.2
Inj 2	2.2	1.9	0.7	0.2

Table 2.3: The error of estimated gradient at day 2, 16, 30, and 44, in percentage.

2.3 Chapter Summary

This chapter develops a method for gradient estimation by using the space-time solution of gray-box conservation law simulations. In particular, an adjoint-enabled twin model is trained to minimize the solution mismatch metric. The inferrability of the twin model is studied theoretically for a simple PDE with only one equation and one dimensional space. To enable the training computationally, a sigmoid parameterization is presented. However, an ad hoc choice for the bases does not fully exploit the information contained in the gray-box solution. To address this issue, an adaptive basis construction procedure is presented. The adaptive procedure builds upon three key elements: the approximated basis significance, the basis neighborhood, and the cross validation. The algorithm for training the twin model is summarized. To alleviate the training cost, a pre-train step is suggested that minimizes the integrated truncation error instead of the solution mismatch.

The proposed twin model algorithm has a wide applicability, which is demonstrated on a variety of numerical examples. The first example is the Buckley-Leverett equation, whose flux function is inferred. The trained twin model accurately estimates the gradient of an objective to the source term. The second example is the steady-state Navier-Stokes equation in a return bend, whose state equation is inferred. The inferred state equation allows estimating the gradient of mass flux to the control surface geometry. The third example is the petroleum reservoir with polymer injection, where the mobility factors are inferred. The gradient of the residual oil to the injection rate is estimated. With the aid of the estimated gradient, the objective can be optimized more efficiently, which will be discussed in the next chapter.

Chapter 3

Leveraging the Twin Model for Bayesian Optimization

This chapter develops a Bayesian optimization framework to solve (1.5). The estimated gradient, provided by the twin model, is utilized to improve the optimization performance. The goal is to reduce the number of gray-box simulations required to achieve a desired objective evaluation, as well as to reduce the overall computational cost. The chapter is organized as follows. Section 3.1.1 develops the probabilistic ingredients for the Bayesian optimization. These ingredients are applied to build an algorithm in Section 3.1.2. Its convergence properties are investigated in Section 3.1.3. Finally, the algorithm is demonstrated in Section 3.2 through several numerical examples.

3.1 Approach

3.1.1 Modeling the Objective and Gradient by Gaussian Processes

Assume the gray-box simulator evaluates the objective function ξ accurately. The adjoint gradient estimated by the twin model is not exactly the true gradient for several reasons. For example, the discretized gray-box solution can be under-resolved, thus limiting the accuracy of the inference of F. In addition, the simulators for the

twin and gray-box models may use different numerical schemes, so the \tilde{F} that yields the minimal solution mismatch may not be exactly F. It is difficult to identify the various sources of errors and unrealistic to quantify all the errors separately. Instead, my thesis treats the gradient error as a whole as a model inadequacy. The model inadequacy is then calibrated using a Bayesian approach.

Given a gray-box solution, the trained twin model is deterministic. Assume the numerical schemes of the gray-box simulator to be deterministic too. Then the gradient error is deterministic. Besides, the gray-box solutions are generally correlated for different controls. Thus the twin models and their gradient estimation errors are also correlated. The deterministic and correlated error can be treated as model inadequacy as discussed in Section 1.3.4. Let $\nabla \xi$ be the true gradient, $\xi_{\tilde{\nabla}}$ be the estimated gradient ¹, and $\xi_{\tilde{\nabla}i}$ be its *i*th component. The relationship between $\nabla \xi$ and $\xi_{\tilde{\nabla}}$ is modeled by the model inadequacy formulation (1.23),

$$\xi_{\tilde{\nabla}i} = \nabla \xi_i + \epsilon_i \,, \tag{3.1}$$

for $i = 1, \dots, d$. (3.1) has the idiosyncratic noise term removed because $\nabla \xi$ and $\xi_{\tilde{\nabla}}$ are deterministic. It also sets the regression parameter, ρ in (1.23), to be 1 for computational simplicity.

Gaussian processes are adopted to model the terms in (3.1). In particular, I made the following assumptions.

- 1. ξ is a realization of a stationary Gaussian process with mean μ , and covariance kernel $K(\cdot, \cdot)$;
- 2. $\epsilon_1, \dots, \epsilon_d$ are realizations of zero-mean stationary Gaussian processes with covariances $G_1(\cdot, \cdot), \dots, G_d(\cdot, \cdot)$, respectively;

 $^{{}^1\}tilde{\xi}(c)$ is the objective evaluation provided by a twin model trained using the gray-box solution at c. $\xi_{\tilde{\nabla}}(c)$ is the adjoint gradient estimated by the twin model. If $\tilde{\xi}$ is differentiable, there is another gradient, $\nabla \tilde{\xi}$, which can be evaluated by finite difference. Usually $\nabla \tilde{\xi} \neq \xi_{\tilde{\nabla}}$. Notice $\xi_{\tilde{\nabla}}$ may not be a conservative vector field.

3. The gradient errors, ϵ_i 's, are independent with the objective,

$$\operatorname{cov}\left[\xi(c_1), \epsilon_i(c_2)\right] = 0, \tag{3.2}$$

for all $c_1, c_2 \in \mathbb{R}^d$, $i = 1, \dots, d$;

4. The components of the gradient error are pairwise independent,

$$cov \left[\epsilon_i(c_1), \epsilon_j(c_2)\right] = 0,$$

for all $c_1, c_2 \in \mathbb{R}^d$ and $i \neq j$;

5. The covariances are isotropic, i.e. $K(c_1, c_2)$, $G_1(c_1, c_2)$, \cdots $G_d(c_1, c_2)$ only depend on $||c_1 - c_2||_{L_2}$.

Suppose ξ and $\xi_{\tilde{\nabla}}$ have been evaluated on \underline{c}_n ². Based upon the assumptions above, the joint distribution of $\xi(c)$, $\xi(\underline{c}_n)$, and $\xi_{\tilde{\nabla}}(\underline{c}_n)$ is multivariate normal, and is given by

$$\begin{pmatrix} \xi(c) \\ \xi(\underline{c}_n) \\ \xi_{\tilde{\nabla}}(\underline{c}_n) \end{pmatrix} \sim \mathcal{N} \begin{pmatrix} \mu \\ \mu \\ 0 \end{pmatrix}, \begin{pmatrix} K(c,c) & \boldsymbol{v} & \boldsymbol{w} \\ \boldsymbol{v}^T & \boldsymbol{D} & \boldsymbol{H} \\ \boldsymbol{w}^T & \boldsymbol{H}^T & \boldsymbol{E} + \overline{\boldsymbol{G}} \end{pmatrix} \end{pmatrix}, \tag{3.3}$$

where

$$\mathbf{v} = (K(c, c_1), \cdots, K(c, c_N)), \qquad (3.4)$$

$$\boldsymbol{w} = (\nabla_{c_1} K(c, c_1), \cdots, \nabla_{c_N} K(c, c_N)), \qquad (3.5)$$

$$\mathbf{D} = \begin{pmatrix} K(c_1, c_1) & \cdots & K(c_1, c_N) \\ \vdots & \ddots & \vdots \\ K(c_N, c_1) & \cdots & K(c_N, c_N) \end{pmatrix}, \tag{3.6}$$

$$\boldsymbol{H} = \begin{pmatrix} \nabla_{c_1} K(c_1, c_1) & \cdots & \nabla_{c_N} K(c_1, c_N) \\ \vdots & \ddots & \vdots \\ \nabla_{c_1} K(c_N, c_1) & \cdots & \nabla_{c_N} K(c_N, c_N) \end{pmatrix},$$
(3.7)

²The notations are consistent with Section 1.3.5. The objective and estimated gradient evaluations are assumed to be collocated, which will be revealed in Section 3.1.2.

$$\boldsymbol{E} = \begin{pmatrix} \nabla_{c_1} \nabla_{c'_1} K(c_1, c'_1) & \cdots & \nabla_{c_1} \nabla_{c'_N} K(c_1, c'_N) \\ \vdots & \ddots & \vdots \\ \nabla_{c_1} \nabla_{c'_N} K(c_N, c'_1) & \cdots & \nabla_{c_N} \nabla_{c'_N} K(c_N, c'_N) \end{pmatrix},$$
(3.8)

$$\overline{\boldsymbol{G}} = \begin{pmatrix} \boldsymbol{G}(c_1, c_1) & \cdots & \boldsymbol{G}(c_1, c_N) \\ \vdots & \ddots & \vdots \\ \boldsymbol{G}(c_N, c_1) & \cdots & \boldsymbol{G}(c_N, c_N) \end{pmatrix},$$
(3.9)

$$G(c_i, c_j) = diag(G_1(c_i, c_j), \dots, G_d(c_i, c_j)), i, j = 1, \dots, d.$$
 (3.10)

The derivation of (3.7) and (3.8) can be found in [63].

The Matérn 5/2 kernel is used for K and G_i 's, in particular,

$$K(c_{1}, c_{2}) = \sigma_{\xi}^{2} \left(1 + \frac{\sqrt{5} \|c_{1} - c_{2}\|_{L_{2}}}{L_{\xi}} + \frac{5 \|c_{1} - c_{2}\|_{L_{2}}^{2}}{3L_{\xi}^{2}} \right) \exp\left(-\frac{\sqrt{5} \|c_{1} - c_{2}\|_{L_{2}}}{L_{\xi}} \right),$$

$$(3.11)$$

$$G_{i}(c_{1}, c_{2}) = \sigma_{G_{i}}^{2} \left(1 + \frac{\sqrt{5} \|c_{1} - c_{2}\|_{L_{2}}}{L_{G_{i}}} + \frac{5 \|c_{1} - c_{2}\|_{L_{2}}^{2}}{3L_{G_{i}}^{2}} \right) \exp\left(-\frac{\sqrt{5} \|c_{1} - c_{2}\|_{L_{2}}}{L_{G_{i}}} \right).$$

$$(3.12)$$

Let θ denote the hyper parameters L_{ξ} , σ_{ξ} , L_{G_i} 's, σ_{G_i} 's, and μ . θ can be estimated by log maximum likelihood. The likelihood of observing $\xi(\underline{c}_n)$ and $\xi_{\tilde{\nabla}}(\underline{c}_n)$ is given by

$$p\left(\xi(\underline{c}_{n}), \xi_{\tilde{\nabla}}(\underline{c}_{n})|\theta\right) = \int p\left(\xi(\underline{c}_{n}), \xi_{\tilde{\nabla}}(\underline{c}_{n}), \nabla\xi(\underline{c}_{n})|\theta\right) d\left(\nabla\xi(\underline{c}_{n})\right)$$

$$= \int p\left(\xi(\underline{c}_{n}), \nabla\xi(\underline{c}_{n})|\theta\right) p\left(\xi_{\tilde{\nabla}}(\underline{c}_{n})|\xi(\underline{c}_{n}), \nabla\xi(\underline{c}_{n});\theta\right) d\left(\nabla\xi(\underline{c}_{n})\right),$$
(3.13)

which is marginalized over $\nabla \xi(\underline{c}_n)$. Because

$$\xi(\underline{c}_n), \nabla \xi(\underline{c}_n) \Big| \theta \sim \mathcal{N} \left(\begin{pmatrix} \boldsymbol{\mu} \\ \mathbf{0} \end{pmatrix}, \begin{pmatrix} \boldsymbol{D} & \boldsymbol{H} \\ \boldsymbol{H}^T & \boldsymbol{E} \end{pmatrix} \right),$$
 (3.14)

and

$$\xi_{\tilde{\nabla}}(\underline{c}_n)|\xi(\underline{c}_n), \nabla \xi(\underline{c}_n); \theta \sim \mathcal{N}\left(\nabla \xi(\underline{c}_n), \overline{G}\right),$$
 (3.15)

the log marginal likelihood has the closed form

$$\log p(\xi(\underline{c}_{n}), \xi_{\tilde{\nabla}}(\underline{c}_{n}) | \theta)$$

$$= -\frac{1}{2} \begin{pmatrix} \xi(\underline{c}_{n}) - \boldsymbol{\mu} \\ \xi_{\tilde{\nabla}}(\underline{c}_{n}) \end{pmatrix}^{T} \begin{pmatrix} \boldsymbol{D} & \boldsymbol{H} \\ \boldsymbol{H}^{T} & \boldsymbol{E} + \overline{\boldsymbol{G}} \end{pmatrix}^{-1} \begin{pmatrix} \xi(\underline{c}_{n}) - \boldsymbol{\mu} \\ \xi_{\tilde{\nabla}}(\underline{c}_{n}) \end{pmatrix} - \frac{1}{2} \log \left(\det \begin{pmatrix} \boldsymbol{D} & \boldsymbol{H} \\ \boldsymbol{H}^{T} & \boldsymbol{E} + \overline{\boldsymbol{G}} \end{pmatrix} \right)$$

$$- \frac{N(d+1)}{2} \log(2\pi), \qquad (3.16)$$

which can be optimized efficiently using GBO methods such as StoGo, as discussed in Section 1.3.1. Given the joint distribution (3.3), the posterior of $\xi(c)$, for any $c \in \mathbb{R}^d$, can be obtained by (1.27).

Using the posterior, the acquisition function, $\rho(c)$, can be constructed and optimized to find the next evaluation point. My thesis will use the expected improvement acquisition function. See Section 1.3.5 for the details.

3.1.2 Algorithm

Based upon the review in Section 1.3.5 and the developments in Section 3.1.1, I present a Bayesian optimization algorithm with bound constraints $c_{\min} \leq c \leq c_{\max}$, Algorithm 3. The flowchart of the algorithm is sketched in Figure 3-1.

In line 3 of Algorithm 3, it is beneficial to reuse twin models, because the twin models that are trained on previous controls may be good initial guesses for the current training. If the gray-box solutions are similar, it is expected that the trained twin models may be similar too. A rigorous investigation of the topic is a future work. Instead, I suggest a tentative procedure to reuse trained twin models as follows: 1) When running Algorithm 3, store u(c) and the trained twin models at all iterates; 2) At each iterate, find a previously trained twin model whose solution is closest to the current solution according to (2.2); 3) Loop over the backward step in Algorithm 1 to prune all possible redundant bases of the twin model; and 4) Apply Algorithm 1

Input: Initial guess c. Current best control c_0^* . Current best objective ξ_0^* . Max iteration n_{max} . Expected improvement threshold EI_{\min} . $D_c = [], D_{\xi_{\tilde{\nabla}}} = [].$ 1: for i = 1 to n_{max} do Simulate the gray-box model on c, obtain $\xi(c)$ and $\boldsymbol{u}(c)$. 2: Train a twin model using $\boldsymbol{u}(c)$, obtain $\xi_{\tilde{\nabla}}(c)$. 3: $D_c = [D_c, c], D_{\xi} = [D_{\xi}, \xi(c)], D_{\xi_{\tilde{\Sigma}}} = [D_{\xi_{\tilde{\Sigma}}}, \xi_{\tilde{\Sigma}}(c)].$ 4: if $\xi(c) > \xi_0^*$ then 5: $c_0^* \leftarrow c$ 6: 7: end if Update hyper parameters by MLE. 8:
$$\begin{split} c \leftarrow \operatorname{argmax}_{c_{\min} \leq c \leq c_{\max}} \log(\rho_{\mathtt{EI}}(c)). \\ \mathbf{if} \ \rho_{\mathtt{EI}}(c) < \mathtt{EI}_{\min} \ \mathbf{then} \end{split}$$
9: 10: break 11: end if 12: 13: end for **Output:** $c_0^*, \, \xi_0^*$

Algorithm 3: Bayesian optimization with twin model.

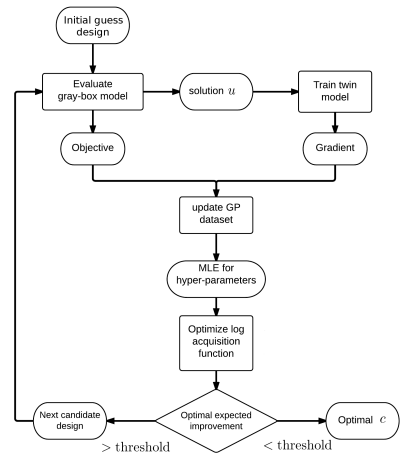


Figure 3-1: The flowchart of Algorithm 3.

to train the twin model as usual.

3.1.3 Convergence Properties Using True Hyper Parameters

This section investigates the convergence properties of Algorithm 3. For Bayesian optimization with only the objective evaluation, the convergence properties have been explored in the past. M. Locatelli [62] proved that Bayesian optimization with EI acquisition generates a dense search sequence for the 1-D optimization problem $c^* = \operatorname{argmax}_{c \in [0,1]} \xi(c)$, if ξ is a realization of the Wienner process. E. Vazquez [56] generalized the results by showing that the sequence is still dense for higher dimensional space and for more general classes of stochastic processes. Recently, A. Bull [57] showed that Bayesian optimization with EI has a convergence rate at $\mathcal{O}(n^{-\nu/d})$, where $\nu > 0$ is a constant parameter controlling the kernel smoothness. Similar results have been given for UCB acquisition. N. Srinivas [58] bounded the convergence rate from above at $\mathcal{O}(n^{-\frac{\nu}{2\nu+d(d+1)}})$ for UCB, and also established the relationship between the convergence rate and the information gain due to evaluating the objective. In this section, I analyze the convergence properties of Algorithm 3 when the true hyper parameters are used. My contribution is to extend the convergence analysis to incorporate estimated gradient evaluations. Under the assumptions in Section 3.1.1, it is proven that the search sequence is indeed dense. The conclusion implies that the algorithm is able to find the optimal as $n_{\text{max}} \to \infty$, regardless of the gradient estimation quality.

The true hyper parameters values are taken as known constants throughout the section. Without loss of generality, assume the objective function to be a realization of zero-mean stationary Gaussian process. The assumptions aforementioned in Section 3.1.1 are reiterated more formally as follows. ξ belongs to the reproducing kernel Hilbert space (RKHS) \mathcal{H}_K generated by a semi-positive definite kernel $K: \mathcal{C} \times \mathcal{C} \to [0, \infty)$. Let K be differentiable, then the gradients of all functions in \mathcal{H}_K form a reproducing kernel Hilbert space $\mathcal{H}_{K_{\nabla}}$ with the kernel $K_{\nabla}(c_1, c_2) \equiv \nabla_{c_1} \nabla_{c_2} K(c_1, c_2)$

for all $c_1, c_2 \in \mathcal{C}$ (theorem 1 in [55]). Besides, ϵ_i , for $i = 1, \dots, d$, belongs to the RKHS \mathcal{H}_G^i generated by a semi-positive definite kernel $G_i : \mathcal{C} \times \mathcal{C} \to [0, \infty)$. ϵ_i 's are pairwise independent. Denote the tensor product of the RKHSs by $\mathcal{H}_G \equiv \mathcal{H}_G^1 \otimes \cdots \otimes \mathcal{H}_G^d$.

Let the stochastic dependence of ξ to be ω_{ξ} , and the stochastic dependence of ϵ_{i} to be ω_{ϵ}^{i} . Let $(\Omega_{\xi}, \Sigma_{\xi}, \mathbb{P}_{\xi})$ be the probability space for ω_{ξ} , and let $(\Omega_{\epsilon}^{i}, \Sigma_{\epsilon}^{i}, \mathbb{P}_{\epsilon}^{i})$ be the probability space for ω_{ϵ}^{i} . Then

$$\xi : \mathcal{C} \times \Omega_{\xi} \to \mathbb{R}$$

$$(c, \omega_{\xi}) \to \xi(c; \omega_{\xi}),$$
(3.17)

and

$$\epsilon : \mathcal{C} \times \Omega^i_{\epsilon} \to \mathbb{R}^d$$

$$(c, \omega^i_{\epsilon}) \to \epsilon(c; \omega^i_{\epsilon})$$
(3.18)

for $i=1,\cdots,d$. Let $\omega_{\epsilon}=(\omega_{\epsilon}^1,\cdots,\omega_{\epsilon}^d)$ and $\Omega_{\epsilon}=\Omega_{\epsilon}^1\otimes\cdots\otimes\Omega_{\epsilon}^d$. The true objective function is $\xi(c;\omega_{\xi}^*)$ for $\omega_{\xi}^*\in\Omega_{\xi}$, and the true estimated gradient error is $\epsilon(c;\omega_{\epsilon}^*)$ for $\omega_{\epsilon}^*\in\Omega_{\epsilon}$. In other words, $\xi(c;\omega_{\xi}^*)=\xi(c)$ and $\epsilon(c;\omega_{\epsilon}^*)=\epsilon(c)$ for all $c\in\mathcal{C}$. Conditioned on $\xi(\underline{c}_n)$ and $\xi_{\widetilde{\nabla}}(\underline{c}_n)$, Bayesian optimization generates the next search point deterministically. Given the initial control c_{init} , the search sequence can be seen as a mapping

$$\underline{C}(\omega_{\xi}, \omega_{\epsilon}) = (C_1(\omega_{\xi}, \omega_{\epsilon}), C_2(\omega_{\xi}, \omega_{\epsilon}), \cdots) , \qquad (3.19)$$

The search strategy \underline{C} generates a random search sequence C_1, C_2, \cdots in C, with the property that C_{n+1} is \mathcal{F}_n -measurable, where \mathcal{F}_n is the σ -algebra generated by $\xi(\underline{c}_n)$ and $\xi_{\tilde{\nabla}}(\underline{c}_n)$. At the n-th search step, the posterior mean and variance of $\xi(c)$ conditioned on $\xi(\underline{c}_n)$ and $\xi_{\tilde{\nabla}}(\underline{c}_n)$ are written as

$$\hat{\xi}_n(c;\underline{c}_n) = \mathbb{E}_{\omega_{\xi},\omega_{\epsilon}} \left[\xi(c,\omega_{\xi}) \middle| \underline{c}_n, \xi(\underline{c}_n), \xi_{\tilde{\nabla}}(\underline{c}_n) \right], \tag{3.20}$$

and

$$\sigma_n^2(c;\underline{c}_n) = \mathbb{E}_{\omega_{\xi},\omega_{\epsilon}} \left[\left(\xi(c) - \hat{\xi}_n(c) \right)^2 \left| \underline{c}_n, \xi(\underline{c}_n), \xi_{\tilde{\nabla}}(\underline{c}_n) \right| \right]. \tag{3.21}$$

Notice $\sigma_n^2(c;\underline{c}_n)$ only depends on \underline{c}_n , and is independent of $\xi(\underline{c}_n), \xi_{\tilde{\nabla}}(\underline{c}_n)$ because of the Gaussian process assumption.

The following theorem holds, which is proven in Appendix A.3.

Theorem 3. Let $\Phi(c) \equiv K(c,0)$ for all $c \in \mathcal{C}$, and let $\hat{\Phi}$ be its Fourier transform. If there exist $C \geq 0$ and $k \in \mathbb{N}^+$, such that $(1 + |\eta|^2)^k |\hat{\Phi}(\eta)| \geq C$ for all $\eta \in \mathbb{R}^d$, and if $n_{\max} \to \infty$ and $EI_{\min} = 0$, then \underline{c}_n is dense in \mathcal{C} for all $c_{init} \in \mathcal{C}$, all $\xi \in \mathcal{H}_K$ and all $\epsilon_i \in \mathcal{H}_G^i$, for $i = 1, \dots, d$.

In the limiting case of $n_{\rm max} \to \infty$ and ${\rm EI}_{\rm min} = 0$, the theorem implies that Algorithm 3 can find the maximum regardless of the accuracy of the gradient estimation if the true hyper parameters are known. It is a future work to extend the theory when the hyper parameters are estimated.

3.2 Numerical Results

This section demonstrates the optimization algorithm on several numerical examples.

3.2.1 Buckley-Leverett Equation

Consider the same problem setup as in Section 2.2.1. Represent the source term by 25 parameters

$$c(t,x) = \sum_{i=1}^{5} \sum_{j=1}^{5} c_{ij} B_{ij}(t,x)$$

$$B_{ij} = \exp\left(-\frac{(t-t_i)^2}{L_t^2}\right) \exp\left(-\frac{(x-x_j)^2}{L_x^2}\right),$$
(3.22)

where $L_t = L_x = 0.15$, and $(t_1, \dots, t_5) = (x_1, \dots, x_5) = \text{linspace(0,1,5)}$. Consider minimizing the objective

$$\xi(c) = \int_{x=0}^{1} \left| u(t=1,x) - \frac{1}{2} \right|^2 + \frac{1}{100} \sum_{ij} c_{ij}^2,$$
 (3.23)

with the bound constraints $-1 \le c_{ij} \le 1$ for $i, j = 1, \dots, 5$.

Figure 3-2a shows the optimized source term. Figure 3-2b shows the corresponding gray-box solution. Notice the solution at t=1 is close to $\frac{1}{2}$ due to the optimized source.

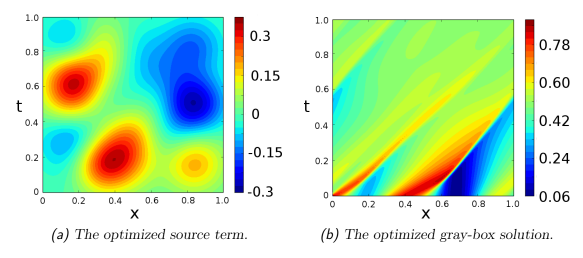


Figure 3-2: Optimized results for the Buckley-Leverett equation.

Constrained by a limited number of gray-box simulations, the optimized solution and objective are examined. Figure 3-3 compares the optimized u(t=1,x) obtained by either the twin-model Bayesian optimization and the vanilla Bayesian optimization³, after 20 gray-box simulations. Figure 3-4 shows the current best (minimal) objective at each iterate. The use of twin model accelerates the optimization, especially when the number of iterate is small.

³The vanilla Bayesian optimization uses only the objective evaluation.



Figure 3-3: A comparison of the optimized u(t = 1, x) after 20 gray-box simulations. The red line is obtained by the vanilla Bayesian optimization and the green line by the twin-model Bayesian optimization. The cyan dashed line indicates the u(t = 1, x) obtained by setting the source term to zero.

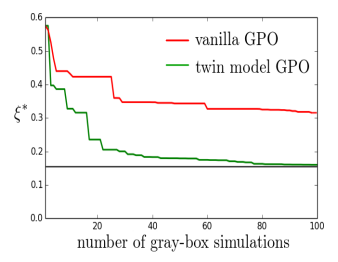


Figure 3-4: The current best objective at each iterate. The red line is obtained by the vanilla Bayesian optimization and the green line by the twin-model Bayesian optimization. The black horizontal line indicates the true optimal.

3.2.2 Navier-Stokes Flow

Consider the same Navier-Stokes flow as in Section 2.2.2. Let S(c) be the area of the return bend, which is a function of the control points' coordinates, c. Let S_0 be the area that corresponds to the initial guess of the control points. The objective function is the steady-state mass flux with a penalty term representing the difference of S and S_0 ,

$$\xi(c) = -\int_{\text{outlet}} \rho_{\infty} u_{\infty} \big|_{\text{outlet}} dy - \lambda (S - S_0)^2, \qquad (3.24)$$

where $\lambda > 0$. The goal is to maximize $\xi(c)$ in a bounded domain $c_{\min} \leq c \leq c_{\max}$. Because there are 4 adjustable control points at each boundary, the control is 16-dimensional. Figure 3-5 shows the initial and the optimized geometries as well as the bound constraints. It also shows the pressure profiles at the interior and the exterior boundaries along the streamwise direction. The optimized geometry reduces the adverse pressure gradient at the flow separation, thus decreases the drag and increases the mass flux.

Figure 3-6 shows the current best objective evaluation at each iterate. Twin model enables faster objective improvement than the vanilla Bayesian optimization. In particular, at the 8th iterate, twin-model Bayesian optimization already achieves near optimality. Figure 3-7 shows the wall clock time of the optimization against the number of iterates. Although twin model increases the per-iterate computational cost, the increased cost is offset by faster objective improvement. After around 80 minutes (8 twin-model optimization iterations), twin model achieves near optimality whereas the vanilla Bayesian optimization is still far from optimal.

3.2.3 Polymer Injection in Petroleum Reservoir

Consider a 2D horizontal reservoir governed by (2.43) and (2.44). The permeability is heterogeneous, and is shown in Figure 3-8. Five injectors are placed along the southern boundary, and one producer is placed in the northeastern corner. The

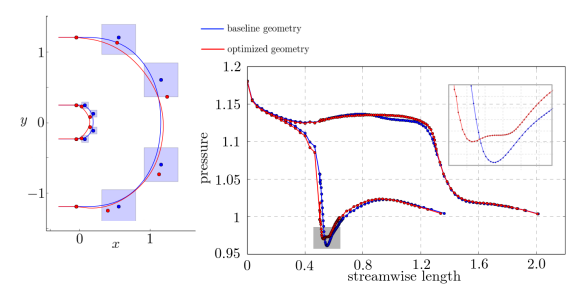


Figure 3-5: The left plot shows the initial guess of control points (blue dots), the initial guess of the geometry (blue line), the optimized control points (red dots), and the optimized geometry (red line). The purple squares indicate the bound constraints for each control point. The right plot shows the pressure along the interior and the exterior boundaries for the initial (blue) and the optimized (red) geometry.

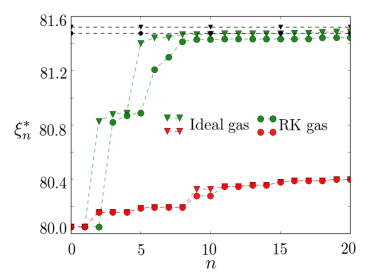


Figure 3-6: The current best objective at each iterate for the ideal gas and the Redlich-Kwong gas. The green lines are obtained by the twin-model Bayesian optimization. The red lines are obtained by the vanilla Bayesian optimization. The black horizontal lines indicate the true optimal.

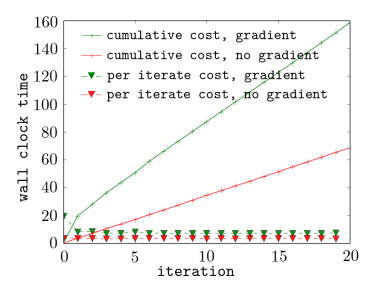


Figure 3-7: The cumulative and per-iterate wall clock time, in minutes.

reservoir is simulated for $t \in [0, T = 10]$ day.

Firstly, consider constant-in-time injection rates at the injectors. Define

$$\xi(t) = -\int_{\Omega} \rho_o(t)\phi S_o(t)d\mathbf{x} - \lambda t \sum_{i=1}^{5} I_{\text{inj}i}, \qquad (3.25)$$

which is the negative oil residual minus the water cost at time t. $\lambda = 0.4$ is the cost of water per unit volume. $I_{\mathtt{inj}i}$ is the injection rate at the ith injector. The goal is to maximize $\xi(T)$ with bound constraints on the injection rates $0 \leq I_{\mathtt{inj}i} \leq I_{\mathtt{max}}$. Since there are five injectors, the optimization is 5-dimensional.

Figure 3-9 shows the current best objective evaluation against the number of iterates. The black line indicates the true optimal⁴. The twin model Bayesian optimization achieves near-optimality faster than the vanilla Bayesian optimization. Figure 3-10 shows $\xi(t)$ for the initial and the optimized injection rates. The initial rates are set at $I_{\text{inj}i} = I_{\text{max}}$ for all injectors, which results in early water breakthrough and high water cut. Although the profit is high at smaller t, it deteriorates for larger

⁴The true optimal is obtained by simulated annealing after running 192 gray-box simulations.

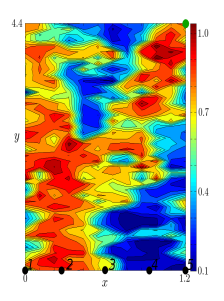


Figure 3-8: The permeability of the reservoir, in 100 milli Darcy. The 5 injectors are indicated by the black dots, and the producer is indicated by the green dot.

t due to the water being wasted.

Secondly, consider time-dependent injection rates. If [0, T] is discretized uniformly into 200 segments, each I_{inji} becomes a vector with a length of 200. Thus the optimization is one thousand dimensional. Clearly the Bayesian optimization algorithm developed in Section 3.1.2 is not long suitable, because the large dimensionality leads to a huge covariance matrix⁵. Instead, the twin model is tested on a simple gradient descent method, the backtracking-Armijo gradient descent method [93]. Clearly, such practice is theoretically flawed, because the gradient error can not be estimated or bounded, and the optimization may not converge at all. However, it is interesting to examine the practical utility of the twin model.

Figure 3-11 shows the optimized injection rates. The 1st and 5th injectors, at the

 $[\]overline{}^5$ As aforementioned, the covariance matrix for evaluating the posterior is N(d+1)-by-N(d+1). For example, after 100 iterates, the matrix becomes 10^5 -by- 10^5 . The optimization algorithm can dominate the computational cost instead of the conservation law simulation, which violates my assumptions in Chapter 1. Such scaling problem is suffered by most non-parametric methods.

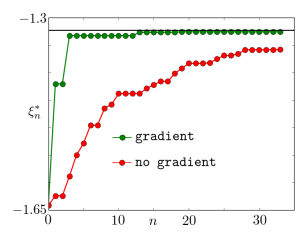


Figure 3-9: The current best objective evaluation against the number of iterates.

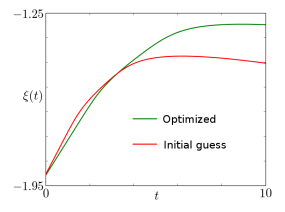


Figure 3-10: $\xi(t)$ for the initial and the optimized injection rates.

southeastern and southwestern corners, are turned on first. The rate at injector 5 is particularly large, possibly because the permeability is relatively low. Once water breaks through and a low-resistance water channel forms, the oil around the injector 5 will be harder to extract. Later, all injectors are turned on, and their rates gradually decrease when the water cut increases. Figure 3-12 shows the current best objective evaluation against the number of iterates. Using the time-dependent control, the objective evaluation gets more improvement than the constant-rate control.

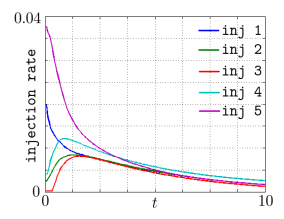


Figure 3-11: The optimized time-dependent injection rates.

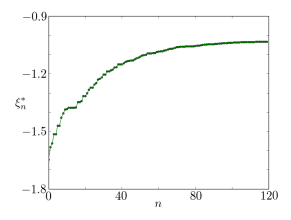


Figure 3-12: The current best objective evaluation using the backtracking-Armijo gradient descent method, where the gradient is provided by the twin model.

3.3 Chapter Summary

Based upon previous research, this chapter develops a Bayesian framework for the optimization problems constrained by gray-box conservation law simulations. Gaussian process models are presented for the objective function, the true gradient, the estimated gradient, and the gradient error. Using the Gaussian process models, the formulation of the joint and the posterior distributions are given, where the hyper parameters are estimated by maximum likelihood. The developments are summarized in a Bayesian optimization algorithm which leverages the twin-model gradient estimation. In addition, the convergence property of the algorithm is theoretically studied. The algorithm is guaranteed to find the optimal regardless of the gradient estimation accuracy, if the true hyper parameters are used. It is a future work to extend the theory to estimated hyper parameters.

The proposed optimization method is demonstrated on several numerical examples. The first example is the Buckley-Leverett equation whose flux is assumed unknown. The objective function is optimized by adjusting the source term represented by 25 control variables. The second example is a Navier-Stokes flow in a return bend, where the state equation is unknown. The mass flux with a penalty on the geometry is maximized by adjusting the flow boundaries which are controlled by 16 variables. The third example is a petroleum reservoir with polymer injections, where the mobility factors are unknown. The profit is maximized by adjusting the constant-time injection rates at five injectors. In all three examples, the twin-model optimization achieves near-optimality with less iterations than the vanilla Bayesian optimization. Finally, the time-dependent control is considered on the same petroleum reservoir example, which yields a 1000-dimensional problem. Conventionally, such high-dimensional optimization can be hard without the adjoint gradient. The twin-model gradient is tested to work well using a gradient descent approach.

Chapter 4

Conclusions

In this thesis, I addressed the optimization constrained by gray-box simulations. I enabled the adjoint gradient computation for gray-box simulations by leveraging the space-time solution. In addition, I utilized the gradient information in a Bayesian framework to faciliate a more efficient optimization. To conclude, this chapter summarizes the developments and highlights the contributions of this work. I close with suggestions for continuing work on this topic.

4.1 Thesis Summary

Optimization constrained by conservation law simulations are prevelant in many engineering applications. In many cases, the code of the simulator is proprietary, legacy, and lacks the adjoint capability. Chapter 1 categorizes such simulators as gray-box. The gray-box scenario limits the efficient application of gradient-based optimization methods. I motivates the need for the adjoint gradient, and explains the feasibility of estimating the adjoint gradient in the gray-box scenario. The key is to leverage the gray-box space-time solution, which contains information of the gray-box simulator but is usually abandoned by conventional optimization methods. To restrict the scope of my thesis, a class of problems is formulated where the flux functions are partially unknown.

To address this issue, an adjoint-enabled twin model is proposed to match the space-time solution. In Chapter 2, I develop a two-stage procedure to estimate the gradient. In the first stage, a twin model is trained to minimize the solution mismatch. In the second stage, the trained twin model computes an adjoint gradient which approximates the true gray-box gradient. For a simple conservation law with only one equation and one dimensional space, I demonstrate theoretically that the twin model can indeed infer the gray-box conservation law on a domain that has large solution variation. To implement the twin model numerically, the unknown part of the flux function is parameterized by a set of basis. I argue that the sigmoid bases are well suited for this problem since their gradients are local. The procedure is demonstrated on a Buckley-Leverett equation using an ad hoc set of sigmoids. Although the estimated gradient is accurate, several limitations are observed which lead to the developments of adaptive basis construction. Several tools are introduced for the adaptive basis construction, including a metric of the basis significance, the basis neighborhood, and the cross validation. The adaptive basis construction fully exploits the information contained in the gray-box solution, and avoids the problem of overfitting. Based upon these developments, a twin model algorithm is presented. To reduce the computational cost of the algorithm, a pre-train step is suggested that minimizes the integrated truncation error instead of the solution mismatch. The twin model algorithm is demonstrated on a variety of numerical examples, including a 1D convection equation with unknown flux function, a 2D steady-state Navier-Stokes flow with unknown state equation, and a 3D petroleum reservoir flow with unknown mobility factors. In all the three examples, the twin model algorithm provides accurate estimates of the true gradient, which represents a major contribution towards enabling the adjoint gradient computation for gray-box simulations.

Using the twin-model gradient, optimization can be done more efficiently. Chapter 3 incorporates the twin-model gradient into a Bayesian optimization framework, in which the objective function, the true gradient, the estimated gradient, and the gradient error are modeled by Gaussian processes. The model provide analytical

expressions for the posterior distributions and the acquisition function, while the hyper parameters are estimated by maximum likelihood. I present a Bayesian optimization algorithm that utilizes the twin-model gradient. In addition, I show that the algorithm is able to find the optimal regardless of the gradient estimation accuracy, if the true hyper parameters are used. The optimization algorithm is demonstrated on several problems similar to Chapter 2, including a Buckley-Leverett equation with source term controls, a Navier-Stokes flow in a return bend with boundary geometry controls, and a petroleum reservoir with polymer-water injection rate controls. In all the three examples, the twin-model optimization achieves near-optimality with less iterations than the vanilla Bayesian optimization. Finally, the twin model gradient is tested on a 1000-dimensional control problem, by employing a simple gradient descent approach. The gradient efficiently enables the optimization of the high-dimensional problem, which represents another major contribution of my thesis.

4.2 Contributions

The main contributions of this work are:

- 1. a twin model algorithm that enables the adjoint gradient computation for graybox conservation law simulations;
- an adaptive basis construction scheme that fully exploits the information of gray-box solutions and avoids overfitting;
- 3. a Gaussian process model of the twin-model gradient and a Bayesian optimization algorithm that employs the twin model; and
- 4. a theoretical and numerical demonstration of the algorithms in a variety of problems.

4.3 Future Work

There are several potential thrusts of further research: A useful extension is to investigate the inferrability of twin models for various conservation laws. In particular, Theorem 1 may be extended for problems with a system of equations and higher spatial dimension. Another interesting extension is to study the applicability of the pre-train step, especially to obtain a necessary and sufficient condition for bounding \mathcal{M} with \mathcal{T} . Finally, in the twin-model Bayesian optimization algorithm, it is of great practical value to reuse twin model more efficiently. My current approach uses the twin model with the closest solution as an initial guess, and re-trains the twin model at every iterate. In the future, an important research topic is on how to utilize all previously trained twin models. Another important topic is to employ "trust region" in the optimization: the same twin model can be used multiple times at different controls inside a trust region¹, thus reducing the training cost. Finally, it is interesting to generalize the formulation (1.6) to incorporate unknown source terms and boundary conditions.

¹In my thesis, the twin model is re-trained at each new control. Generally, gradient-based trust region methods require the gradient to satisfy a property called full-linearity. Unfortunately, this property is not guaranteed by the twin-model gradient. The lack of full-linearity is a key factor that refrains me from exploring the trust-region methods. See [42] and [43] for the details.

Appendix A

Proof of Theorems

A.1 Theorem 1

Proof:

We prove false the contradiction of the theorem, which reads:

For any $\delta > 0$ and T > 0, there exist $\epsilon > 0$, and F, \tilde{F} satisfying the conditions stated in theorem 1, such that $\|\tilde{u} - u\|_{\infty} < \delta$ and $\left\| \frac{d\tilde{F}}{du} - \frac{dF}{du} \right\|_{\infty} > \epsilon$ on B_u .

We show the following exception to the contradiction in order to prove it false. For any $\epsilon > 0$ and any F, \tilde{F} satisfying $\left\| \frac{d\tilde{F}}{du} - \frac{dF}{du} \right\|_{\infty} > \epsilon$ on B_u , we can find $\delta > 0$ and T > 0 such that $\|\tilde{u} - u\|_{\infty} > \delta$.

The idea is to construct such an exception by the method of lines [90]. Firstly, assume there is no shock wave for (2.4) and (2.5) for $t \in [0, T]$. Choose a segment in space, $[x_0 - \Delta, x_0]$ with $0 < \Delta < \frac{\epsilon}{L_F L_u}$, that satisfies

- $u_0(x) \in B_u$ for any $x \in [x_0 \Delta, x_0]$;
- $\left| \frac{d\tilde{F}}{du} (u_0(x_0)) \frac{dF}{du} (u_0(x_0)) \right| > \epsilon;$
- $x_0 \Delta + \frac{dF}{du} (u_0(x_0 \Delta))T = x_0 + \frac{d\tilde{F}}{du} (u_0)T \equiv x^*$.

Without loss of generality, we assume $\frac{dF}{du}>0$ and $\frac{d\tilde{F}}{du}>0$ for $\{u\big|u=u_0(x)\,,\ x\in$

 $[x_0 - \Delta, x_0]$. Using the method of lines, we have

$$u\left(T, x_0 - \Delta + \frac{dF}{du}(u_0(x_0 - \Delta))T\right) = u_0(x_0 - \Delta),$$

and

$$\tilde{u}\left(T, x_0 + \frac{d\tilde{F}}{du}(u_0(x_0))T\right) = u_0(x_0).$$

Therefore

$$|\tilde{u}(x^*,T) - u(x^*,T)| = |u_0(x_0) - u_0(x_0 - \Delta)| \ge \gamma \Delta \equiv \delta,$$

by using the definition of B_u .

Set
$$T = \frac{\Delta}{\left|\frac{d\tilde{F}}{du}\left(u_0(x_0-\Delta)\right) - \frac{dF}{du}\left(u_0(x_0)\right)\right|}$$
, we have
$$\left|\frac{d\tilde{F}}{du}\left(u_0(x_0-\Delta)\right) - \frac{dF}{du}\left(u_0(x_0)\right)\right|$$

$$= \left|\frac{dF}{du}\left(u_0(x_0)\right) - \frac{d\tilde{F}}{du}\left(u_0(x_0)\right) + \frac{dF}{du}\left(u_0(x_0-\Delta)\right) - \frac{dF}{du}\left(u_0(x_0)\right)\right|$$

$$= \left|\frac{dF}{du}\left(u_0(x_0)\right) - \frac{d\tilde{F}}{du}\left(u_0(x_0)\right) + \frac{\overline{d^2F}}{du^2}\left(u_0(x_0-\Delta) - u_0(x_0)\right)\right|$$

$$\geq \left|\frac{dF}{du}\left(u_0(x_0)\right) - \frac{d\tilde{F}}{du}\left(u_0(x_0)\right)\right| - L_uL_F\Delta$$

$$> \epsilon - L_uL_F\Delta \equiv \epsilon_F > 0$$

by using the mean value theorem. Therefore $T \leq \frac{\Delta}{\epsilon_F} < \infty$. So we find a $\delta = \gamma \Delta$ and a $T < \infty$ that provides an exception to the contradiction of the theorem.

Secondly, if there is shock wave within [0,T] for either (2.4) or (2.5), we let T^* be the time of the shock occurrence. Without loss of generality, assume the shock occurs for (2.4) first. The shock implies the intersection of two characteristic lines. Choose a $\Delta > 0$ such that $\left| \frac{dF}{du} (u_0(x)) - \frac{dF}{du} (u_0(x-\Delta)) \right| T^* = \Delta$. Using the mean

value theorem, we have

$$T^* = \frac{\Delta}{\frac{\overline{d^2 F}}{du^2} (u_0(x) - u_0(x - \Delta))} \ge \frac{1}{L_u L_F}$$

Thus, if we choose

$$T = \min \left\{ \frac{1}{L_u L_K}, \ \frac{\Delta}{\epsilon_{\Delta}} \right\} \ ,$$

no shock occurs in $t \in [0,T]$. Since the theorem is already proven for the no-shock scenario, the proof completes.

A.2 Theorem 2

Proof:

Let the one-step time marching of the gray-box simulator be

$$\mathcal{H}: \mathbb{R}^n \mapsto \mathbb{R}^n, \, \boldsymbol{u}_{i\cdot} \to \boldsymbol{u}_{i+1\cdot} = \mathcal{H}_i \boldsymbol{u}_{i\cdot}, \quad i = 1, \cdots, M-1,$$

The integrated truncation error can be written as

$$\mathcal{T}(\tilde{F}) = \sum_{i=1}^{M} \sum_{j=1}^{N} w_{j} (u_{i+1j} - (\mathcal{G}u_{i\cdot})_{j})^{2}$$

$$= \sum_{i=1}^{M} (u_{i+1} - \mathcal{G}u_{i\cdot})^{T} W(u_{i+1} - \mathcal{G}u_{i\cdot})$$

$$= \sum_{i=1}^{M} \|u_{i+1} - \mathcal{G}u_{i\cdot}\|_{W}^{2}$$

$$= \sum_{i=1}^{M} \|\mathcal{H}u_{i\cdot} - \mathcal{G}u_{i\cdot}\|_{W}^{2}$$

$$= \sum_{i=1}^{M} \|(\mathcal{H}^{i} - \mathcal{G}\mathcal{H}^{i-1}) u_{0\cdot}\|_{W}^{2}.$$

Similarly, the solution mismatch can be written as

$$\mathcal{M}(\tilde{F}) = \sum_{i=1}^{M} \left\| \left(\mathcal{H}^{i} - \mathcal{G}^{i} \right) u_{0}. \right\|_{W}^{2}$$

Fig A-1 gives an explanation of \mathcal{M} and \mathcal{T} by viewing the simulators as discrete-time dynamical systems.

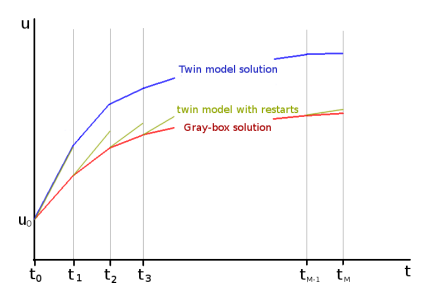


Figure A-1: The state-space trajectories of the gray-box model and the twin model. \mathcal{M} measures the difference of the twin model trajectory (blue) with the gray-box trajectory (red). \mathcal{T} measures the difference of the twin model trajectory with restarts (green) and the gray-box trajectory (red).

Using the equality

$$\mathcal{G}^i - \mathcal{H}^i = (\mathcal{G}^i - \mathcal{G}^{i-1}\mathcal{H}) + (\mathcal{G}^{i-1}\mathcal{H} - \mathcal{G}^{i-2}\mathcal{H}^2) + \dots + (\mathcal{G}\mathcal{H}^{i-1} - \mathcal{H}^i), \quad i \in \mathbb{N},$$

and triangular inequality, we have

$$\mathcal{M} \leq \left\{ \begin{aligned} \|(\mathcal{G}^{M-1}\mathcal{G} - \mathcal{G}^{M-1}\mathcal{H})u_{0}.\|_{W}^{2} + \|(\mathcal{G}^{M-2}\mathcal{G}\mathcal{H} - \mathcal{G}^{M-2}\mathcal{H}^{2})u_{0}.\|_{W}^{2} & + \dots + \|(\mathcal{G}\mathcal{H}^{M-1} - \mathcal{H}^{M})u_{0}.\|_{W}^{2} \\ & + \|(\mathcal{G}^{M-2}\mathcal{G} - \mathcal{G}^{M-2}\mathcal{H})u_{0}.\|_{W}^{2} & + \dots + \|(\mathcal{G}\mathcal{H}^{M-2} - \mathcal{H}^{M-1})u_{0}.\|_{W}^{2} \\ & \ddots & & \vdots \\ & & + \|(\mathcal{G} - \mathcal{H})u_{0}.\|_{W}^{2} \end{aligned} \right\}.$$

Therefore,

$$\mathcal{M}-\mathcal{T} \leq$$

$$\begin{cases}
\|(\mathcal{G}^{M-1}\mathcal{G} - \mathcal{G}^{M-1}\mathcal{H})u_{0}\|_{W}^{2} + \|(\mathcal{G}^{M-2}\mathcal{G}\mathcal{H} - \mathcal{G}^{M-2}\mathcal{H}^{2})u_{0}\|_{W}^{2} + \dots + \|(\mathcal{G}\mathcal{G}\mathcal{H}^{M-2} - \mathcal{G}\mathcal{H}^{M-1})u_{0}\|_{W}^{2} \\
+ \|(\mathcal{G}^{M-2}\mathcal{G} - \mathcal{G}^{M-2}\mathcal{H})u_{0}\|_{W}^{2} + \dots + \|(\mathcal{G}\mathcal{G}\mathcal{H}^{M-3} - \mathcal{G}\mathcal{H}^{M-2})u_{0}\|_{W}^{2} \\
\vdots \\
+ \|(\mathcal{G}\mathcal{G} - \mathcal{G}\mathcal{H})u_{0}\|_{W}^{2}
\end{cases}$$

Under the assumption

$$\left\|\mathcal{G}a - \mathcal{G}b\right\|_{W}^{2} \le \beta \|a - b\|_{W}^{2},$$

and its implication

$$\left\|\mathcal{G}^i a - \mathcal{G}^i b\right\|_W^2 \le \beta^i \|a - b\|_W^2, \quad i \in \mathbb{N},$$

we have

$$\mathcal{M}-\mathcal{T} \leq \left\{ \begin{array}{ccc} \beta^{M-1} \| (\mathcal{G} - \mathcal{H}) u_{0 \cdot} \|_{W}^{2} + \beta^{M-2} \| (\mathcal{G} \mathcal{H} - \mathcal{H}^{2}) u_{0 \cdot} \|_{W}^{2} & + \dots + \beta \| (\mathcal{G} \mathcal{H}^{M-2} - \mathcal{H}^{M-1}) u_{0 \cdot} \|_{W}^{2} \\ & + \beta^{M-2} \| (\mathcal{G} - \mathcal{H}) u_{0 \cdot} \|_{M-1}^{2} & + \dots + \beta \| (\mathcal{G} \mathcal{H}^{n-3} - \mathcal{H}^{n-2}) u_{0 \cdot} \|_{W}^{2} \\ & \ddots & \vdots \\ & + \beta \| (\mathcal{G} - \mathcal{H}) u_{0 \cdot} \|_{W}^{2} \end{array} \right\}.$$

Reorder the summation, we get

$$\mathcal{M}-\mathcal{T} \leq \left\{ \begin{array}{ccc} \beta^{M-1} \| (\mathcal{G} - \mathcal{H}) u_{0 \cdot} \|_{W}^{2} + \beta^{M-2} \| (\mathcal{G} - \mathcal{H}) u_{0 \cdot} \|_{W}^{2} & + \dots + \beta \| (\mathcal{G} - \mathcal{H}) u_{0 \cdot} \|_{W}^{2} \\ & + \beta^{M-2} \| (\mathcal{G} \mathcal{H} - \mathcal{H}^{2}) u_{0 \cdot} \|_{W}^{2} & + \dots + \beta \| (\mathcal{G} \mathcal{H} - \mathcal{H}^{2}) u_{0 \cdot} \|_{W}^{2} \\ & & \vdots \\ & + \beta \| (\mathcal{G} \mathcal{H}^{M-2} - \mathcal{H}^{M-1}) u_{0 \cdot} \|_{W}^{2} \end{array} \right\}.$$

Therefore,

$$\mathcal{M} - \mathcal{T} \le (\beta^{M-1} + \beta^{M-2} + \dots + \beta) \mathcal{T}$$

If β is strictly less than 1, then

$$\mathcal{M} \le \frac{1}{1-\beta}\mathcal{T}\,,$$

thus completes the proof.

A.3 Theorem 3

Proof:

Firstly, we have the following lemma (Chapter 1, Theorem 4.1, [59]).

lemma 1. Let K_1, K_2 be the reproducing kernels of functions on C with norms $\|\cdot\|_{\mathcal{H}_1}$ and $\|\cdot\|_{\mathcal{H}_2}$ respectively. Then $K = K_1 + K_2$ is the reproducing kernel of the space

$$\mathcal{H} = \mathcal{H}_1 \oplus \mathcal{H}_2 = \{ f = f_1 + f_2, f_1 \in \mathcal{H}_1, f_2 \in \mathcal{H}_2 \}$$

with norm $\|\cdot\|_{\mathcal{H}}$ defined by

$$\forall f \in \mathcal{H} \quad \|f\|_{\mathcal{H}}^2 = \min_{f = f_1 + f_2, \ f_1 \in \mathcal{H}_1, f_2 \in \mathcal{H}_2} \left(\|f_1\|_{\mathcal{H}_1^2} + \|f_2\|_{\mathcal{H}_2}^2 \right)$$

Using lemma 1, we prove the following Cauchy-Schwarz inequality, lemma 2.

$$\begin{split} & \left| \xi(c, \omega_{\xi}) - \hat{\xi}(c; \underline{c}_n) \right|^2 \leq \\ & \left(\left(1 + \frac{4d}{3} \right) \left\| \xi(c; \omega_{\xi}) \right\|_{\mathcal{H}_K} + \frac{4d}{3} \left\| \nabla_c \xi(c; \omega_{\xi}) \right\|_{\mathcal{H}_{K_{\nabla}}} + \frac{4}{3} \sum_{i=1}^d \left\| \epsilon_i(c; \omega_{\epsilon}^i) \right\|_{\mathcal{H}_G^i} \right) \sigma^2(c; \underline{c}_n) \end{split}$$

To prove lemma 2, we define a vector

$$u = (u_1, \cdots, u_d)^T \in \mathcal{U}$$
,

where $\mathcal{U} = [0, 1]^d$. Define an auxiliary function

$$\mathcal{Y}(c, u; \omega_{\xi}, \omega_{\epsilon}) = \left(1 - \sum_{i=1}^{d} u_{i}\right) \xi(c, \omega_{\xi}) + u^{T} \left[\nabla_{c} \xi(c, \omega_{\xi}) + \epsilon(c; \omega_{\epsilon})\right].$$

 u_1, \dots, u_d are functions from the Sobolev space $W^{1,2}$ defined on \mathcal{U} , equipped with the inner product

$$\langle \phi, \psi \rangle = \int_{\mathcal{U}} \phi \psi + (\nabla \phi)^T (\nabla \psi) \ du,$$

The Sobolev space is a RKHS with the kernel

$$K_u(\phi, \psi) = \frac{1}{2} \exp\left(-|\phi - \psi|\right)$$

on $\mathcal{U} = [0, 1]$. Given ω_{ξ} and ω_{ϵ} , $\mathcal{Y}(\cdot, \cdot; \omega_{\xi}, \omega_{\epsilon})$ can be viewed as a realization from a RKHS $\mathcal{H}_{\mathcal{Y}}$, defined on $\mathcal{C} \times \mathcal{U}$. Let the kernel function of $\mathcal{H}_{\mathcal{Y}}$ be

$$K_{\mathcal{Y}}: \mathcal{C} \times \mathcal{U}, \mathcal{C} \times \mathcal{U} \to \mathbb{R}$$

$$(c_1, u_1), (c_2, u_2) \to K_{\mathcal{Y}}((c_1, u_1), (c_2, u_2))$$

Notice

$$\mathcal{Y}(c, \mathbf{0}; \omega_{\xi}, \omega_{\epsilon}) = \xi(c, \omega_{\xi})$$

is the objective function, and

$$\left(\mathcal{Y}(c, e_1; \omega_{\xi}, \omega_{\epsilon}), \cdots, \mathcal{Y}(c, e_d; \omega_{\xi}, \omega_{\epsilon})\right) = \nabla_c \xi(c; \omega_{\xi}) + \epsilon(c; \omega_{\epsilon})$$

is the estimated gradient, where $e_i, i = 1, \dots, d$ indicates the *i*th unit Cartesian basis vector in \mathbb{R}^d . Conditioned on the samplings $\xi(\underline{c}_n)$ and $\xi_{\tilde{\nabla}}(\underline{c}_n)$, we can bound the error of the estimation of $\mathcal{Y}(c, \mathbf{0}; \omega_{\xi}, \omega_{\epsilon})$ by the Cauchy-Scharz inequality [59] in $\mathcal{H}_{\mathcal{Y}}$,

$$\left| \mathcal{Y}(c, \mathbf{0}; \omega_{\xi}, \omega_{\epsilon}) - \hat{\mathcal{Y}}(c, \mathbf{0}; \underline{c}_{n}) \right| = \left| \xi(c; \omega_{\xi}) - \hat{\xi}_{n}(c; \underline{c}_{n}) \right| \leq \sigma(c; \underline{c}_{n}) \|\mathcal{Y}\|_{\mathcal{H}_{\mathcal{Y}}}$$

Besides,

$$\|\mathcal{Y}\|_{\mathcal{H}_{\mathcal{Y}}} = \left\| \left(1 - \sum_{i=1}^{d} u_{i} \right) \xi(c; \omega_{\xi}) + u^{T} \left[\nabla_{c} \xi(c; \omega_{\xi}) + \epsilon(c; \omega_{\epsilon}) \right] \right\|_{\mathcal{H}_{\mathcal{Y}}}$$

$$\leq \|\xi(c; \omega_{\xi})\|_{\mathcal{H}_{K}} + \left(\sum_{i=d}^{d} \|u_{i}\|_{\mathcal{H}_{u}} \right) \|\xi(c; \omega_{\xi})\|_{\mathcal{H}_{K}} + \left(\sum_{i=d}^{d} \|u_{i}\|_{\mathcal{H}_{u}} \right) \|\nabla_{c} \xi(c; \omega_{\xi})\|_{\mathcal{H}_{K_{\nabla}}}$$

$$+ \sum_{i=1}^{d} \|u_{i} \epsilon_{i}(c; \omega_{\epsilon}^{i})\|_{\mathcal{H}_{u} \otimes \mathcal{H}_{G}^{i}}$$

$$= \|\xi(c, \omega)\|_{\mathcal{H}_{K}} + \frac{4d}{3} \|\xi(c, \omega)\|_{\mathcal{H}_{K}} + \frac{4d}{3} \|\nabla_{c} \xi(c; \omega_{\xi})\|_{\mathcal{H}_{K_{\nabla}}} + \frac{4}{3} \sum_{i=1}^{d} \|\epsilon_{i}(c; \omega_{\epsilon}^{i})\|_{\mathcal{H}_{G}^{i}},$$

where the inequality obtained by lemma 1. The proof for lemma 2 completes.

Using lemma 2, we prove

lemma 3. Let $(\underline{c}_n)_{n\geq 1}$ and $(\underline{a}_n)_{n\geq 1}$ be two sequences in C. Assume that the sequence (a_n) is convergent, and denote by a^* its limit. Then each of the following conditions implies the next one:

- 1. a^* is an adherent point of \underline{c}_n (there exists a subsequence in \underline{c}_n that converges to a^*),
- 2. $\sigma^2(a_n; \underline{c}_n) \to 0 \text{ when } n \to \infty$,
- 3. $\hat{\xi}(a_n;\underline{c}_n) \to \xi(a^*,\omega)$ when $n \to \infty$, for all $\xi \in \mathcal{H}_K$, $\epsilon \in \mathcal{H}_G$.

The proof of lemma 3 is the similar as the proposition 8 in [56], except that the Cauch-Schwarz inequality used in the paper is replaced by lemma 2. We do not repeat the proof but refer to [56] for the details.

Next, we show the three conditions are equivalent in lemma 3. Using the assumption: There exist $C \geq 0$ and $k \in \mathbb{N}^+$, such that $(1 + |\eta|^2)^k |\hat{\Phi}(\eta)| \geq C$ for all $\eta \in \mathbb{R}^d$, we have, for any $\xi \in \mathcal{H}_K$ and its Fourier transform $\hat{\xi}$,

$$\|\xi\|_{W^{k,2}} = \int (1+|\eta|^2)^k |\hat{\xi}|^2 d\eta \ge C \int |\hat{\Phi}(\eta)|^{-1} |\hat{\xi}(\eta)|^2 d\eta = C\sqrt{(2\pi)^d} \|\xi\|_{\mathcal{H}_K},$$

where $W^{k,2}$ is the Sobolev space whose weak derivatives up to order k have a finite L^2 norm [57]. Therefore, $W^{k,2} \subseteq \mathcal{H}_K$. The result can be extended to $\xi \in \mathcal{H}_K(\mathcal{C})$ defined on the domain $\mathcal{C} \in \mathbb{R}^d$, because $\mathcal{H}_K(\mathcal{C})$ embeds isometrically into $\mathcal{H}_K(\mathbb{R}^d)$ [69]. Besides, we have that C_c^{∞} is dense in $W^{k,2}$ (Chapter 2, Lemma 5.1 [70]), where C_c^{∞} is the C^{∞} functions with compact support on \mathcal{C} . As a consequence, $\mathcal{C}_c^{\infty} \subseteq \mathcal{H}_K$ [56]. If the condition 1 is false, then there exist a neighborhood U of a^* that does not intersect \underline{c}_n . There exist $\xi \in \mathcal{H}_K$ that is compactly supported in U, and $\epsilon = \mathbf{0}$, such that $\hat{\xi}(a^*;\underline{c}_n) = 0$ whereas $\xi(a^*) \neq 0$, which violates the condition 3. Therefore, the three conditions in lemma 3 are equivalent.

Finally, we have:

lemma 4. (E. Vazquez, Theorem 5 [56]) If the three conditions in lemma 3 are equivalent, $n_{\text{max}} \to \infty$, and $EI_{\text{min}} = 0$, then for all $c_{init} \in \mathcal{C}$ and all $\omega \in \mathcal{H}$, the sequence \underline{c}_n generated by the Bayesian optimization with expected improvement acquisition is dense in \mathcal{C} .

We do not repeat the proof. See [56] for the details. To sum up, under the conditions in Theorem 3, \underline{c}_n is dense in the search space.

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