

Recursive Surrogate-Modeling for Stochastic Search

**Bachelor's Thesis
of**

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Karlsruhe, den 26. März 2021

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Zusammenfassung

Roboter werden in den nächsten Jahrzehnten zunehmend unseren Alltag durchdringen. Sie werden ein breites Spektrum an Aufgaben wie Altenpflege, Such- und Rettungsaufgaben und allgemeine Assistenz im täglichen Leben übernehmen können. Derzeit sind die meisten Roboter darauf angewiesen von einem erfahrenen menschlichen Bediener programmiert zu werden. Damit Roboter Teil des täglichen Lebens werden können, müssen sie sich in verändernden Umgebungen zurechtfinden und sich an neue Situationen anpassen. Daher werden selbstlernende Roboter zunehmend an Bedeutung gewinnen. Lernende Roboter ist bereits ein aktives Forschungsfeld und wird weiter wachsen. Es besteht eine enge Beziehung zwischen Reinforcement Learning und Robotik, wobei sich beide Forschungsgebiete gegenseitig ergänzen. Eine wichtige Herausforderung ist es dateneffiziente Methoden für selbstlernende Roboter zu entwickeln, da Stichproben aus realen Interaktionen sehr kostspielig sind. Policy-Suchmethoden, ein Teilgebiet des Reinforcement Learning, können eingesetzt werden um verschiedene Aufgaben durch Versuch und Irrtum zu erlernen.

In dieser Arbeit versuchen wir, die Dateneffizienz eines Policy-Suchalgorithmus zu verbessern. Dafür benutzen wir rekursive Schätzmethoden wie den Kalman Filter, der aus einer Bayes'schen Perspektive eingeführt wird. Wir implementieren verschiedene Versionen von Filteralgorithmen und vergleichen sie mit bisherigen Methoden und testen unsere Algorithmen für Optimierungs-Testfunktionen und einfache planare Greifaufgaben.

Abstract

Robots will increasingly permeate our daily lives over the next few decades. Fulfilling a wide range of tasks like elder care, search and rescue and general assistance in daily life. Today most robots rely on being taught and programmed by a skilled human operator. For robots to become part of daily life they need to cope with changing environments and adjust to new situations, therefore problem of autonomously learning robotics will become more important. Currently robot learning is already a very active field of research. There is a close relationship between Reinforcement learning and robotics, where both areas of research complement each other. A big challenge for robot learning is being sample efficiency, because real world samples are costly to obtain. Policy search methods, a sub-field of reinforcement learning, can be used to learn different task only through trial and error.

In this thesis we try to improve the sample efficiency of a policy search algorithm. This is done using classical recursive estimation techniques like the Kalman filter, introduced from a Bayesian perspective. We implement different versions of Filtering algorithms and compare them with previous methods and benchmark them on optimization test function and simple planar reaching tasks.

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Chapter 1.

Introduction

1.1. Motivation

Robots are already used extensively in industry to form production chains, where they perform the same task over and over. These robots are being programmed and fine tuned by a human engineer which requires experience and expertise.

Recently there has been a new development of robots becoming part of our daily life's, for example in the form of vacuum cleaner and lawn mowers . In the future areas like care giving and everyday assistance and household robots may become more established Schaal (2007). This poses a dramatic shift from the prior uses of robots in industry, where they mainly worked in isolated and predefined contexts. Especially countries like Japan, facing the problem of an aging population, put increasing effort to make robots viable in these domains.

In daily life the robots are confronted with different challenges, like adapting to different like lightening conditions and objectives being moved around. The robots which are currently already on the market like vacuum cleaners and lawn mowers have either a “one size fits all” approach or need a special setup in software or hardware. To solve this issue machine learning and especially reinforcement learning will be key technologies to enable robots to adjust to dynamic and stochastic environments.

Robotics and reinforcement learning complement each other with robotics providing a great, and reinforcement learning providing the framework for formulating solutions, the relationship may be similar to the one of math and physics Kober et al. (2013).

Compared to other domains like board games Go, or video games Atari Robotics poses special challenges for reinforcement algorithms. Making it necessary to explore methods specialized to the inherent challenges of Robotics, which include:

- high dimensional state and action space
- obtaining real world samples
- goal specification
- under-modeling and model uncertainty

In general Robots have potential to transform our society, by bringing robots from the factories into our homes there will be many possibilities for improvement and challenges likewise.

- TODO: include pictures of robots solving tasks with RL

1.2. Contribution

Policy search algorithms have shown promise as an alternative to value function-based reinforcement learning, especially for learning motor skills in robotics Deisenroth et al. (2013). The MORE algorithm as a policy search method based on information theoretic updates is introduced in Abdolmaleki et al. (2015). The key idea of MORE is to learn a surrogate Model of the objective function to efficiently computing the updates to the policy in closed form. One of the contributions of this thesis is to explore recursive estimation for learning the surrogate model of the objective function to increase sample efficiency. Besides that we aim to improve the runtime of the algorithm with this approach. We focus mainly on classical methods of parameter estimation and filtering like the Recursive Least Squares and the Kalman Filter, considering more advanced methods is part of future work. We benchmark the our version of the MORE algorithm on test functions and simple planar reaching tasks and compare them to previous results.

1.3. Structure of Thesis

The remainder of this thesis is structured as follows:

Chapter 2: In the fundamentals chapter we lay the foundation for the thesis, first introducing the basics of Reinforcement Learning and then focusing on the specifics of applying RL to robotic tasks. Then We focus on policy search as one method for solving the robot learning problem. Next we introduce the original MORE algorithm and finally we look at Bayesian filtering.

Chapter 3: Review of the related work in the field.

Chapter 4: In this chapter we motivate the idea of using recursive estimation for learning the surrogate model and then review methods used for data preprocessing. Finally we develop our approach of connecting the MORE algorithm with recursive parameter estimation.

Chapter 5: In the evaluation chapter we conduct experiments with the algorithms on several tests function and some simple reaching tasks for a planar robot. We compare our algorithm with original MORE and other stochastic search algorithms.

Chapter 6: We conclude the thesis with a summary of the achieved results and an outlook on future work.

Chapter 2.

Fundamentals

This chapter introduces basic concepts used throughout this thesis. First basics of reinforcement learning and the problem of robot learning. Next we discuss policy search, a sub-field of reinforcement learning, as one method to solve the robot learning problem. The Kullback-Leibler divergence (KL), as an important information theoretic distance metric between probability distributions. Having discussed the underlying basics we can then introduce the MORE Algorithm. Finally we will look at Filtering from a Bayesian Estimation viewpoint and review the Kalman Filter for parameter estimation.

2.1. Reinforcement Learning

Reinforcement Learning is a subfield of Machine Learning concerned with agents learning to interact with their environment. This is done through exploration and trial-and-error, trying to discover cause and effect relationship between actions. Compared to supervised learning and unsupervised learning it more closely resembles the way humans learn.

As Sutton and Barto (2018) puts it, the term reinforcement learning relates to a class of problems, solution methods and the field that studies these problems and solutions. Some famous examples include playing games like Go Silver et al. (2016) and Atari games Mnih et al. (2013) Generally RL is applicable to a large range of problems. Whereas in supervised learning the best action is presented to the system, the agent in a reinforcement learning setting receives an occasional reward (or punishment). To gain information about the rewards the agent needs to explore previously unused actions. Dare to try new things or keep performing safe well-known actions, this is the *exploration-exploitation tradeoff*. The agent should exploit actions he knows that give decent reward, but he first has to try different things to learn about

these actions, and then he has to progressively focus in on them. The reward signal is only given occasionally, hence the amount of information the agent receives is minimal compared to supervised and unsupervised learning approaches.

We model the agent and its environment as a state $s \in S$. The agent may perform action $a \in A$ which can be either discrete or continuous. For every action step the agent receives a Reward R , which is a scalar value. The overarching goal is to find a mapping from states to actions, called policy π , that picks actions in a way that the reward is maximized. We can distinguish an *episodic* setting from an on-going task. In the episodic setting the task is restarted after the end of an episode and the goal is to maximize the total reward per episode. In on-going tasks the goal is to simply achieve high average reward over the whole life-time or one can use a formulation with a discounted return (weighting the future and past differently).

The classical approach to formalizing problems in RL is through Markov Decision Processes (MDPs). MDPs are a mathematical framework for decision making in deterministic and stochastic environments. MDPs focus on only three aspects - sensation, action and goal, which are central for reinforcement learning problems. MDPs satisfy the Markov property (cite), which state that “the future is independent of the past given the present”. In our case this means the next state s' and the reward only depend on the previous state s and action a Sutton et al. (1992). A MDP can be formally defined as a tuple (S, A, P, r) :

- a set of states $s \in S$ that describe the environment.
- a set of actions $a \in A$ that can be performed by the agent in the environment
- a transition function $P(s_{t+1}|s_t, a_t)$ that gives the probability of a new state s_{t+1} after an action a_t has been taken in state s_t
- a reward function $r(s_t, a_t)$ that specifies the immediate reward after taking action a_t in state s_t

The Markov property can be expressed as

$$P(s_{t+1}|s_t, a_t, s_{t-1}, a_{t-1} \dots) = P(s_{t+1}|s_t, a_t)$$

This recapitulates the notion of state - a state is a sufficient statistic for predicting the future, rendering previous observations irrelevant. In robotics we may only find some approximate notion of state.

Generally the goal is to find an optimal policy π^* , a mapping from states to actions that maximizing the expected return J . Optimal behavior can be modeled in different ways, resulting in different definitions for expected return.

Finite-horizon model, for horizon H meaning the next H timesteps:

$$J = E \left\{ \sum_{t=0}^H R_t \right\}$$

Discounted reward, discounting future rewards by a discount factor γ (with $0 \leq \gamma < 1$):

$$J = E \left\{ \sum_{t=0}^{\infty} \gamma^t R_t \right\}$$

Reinforcement learning can also be seen as general case of optimal control as in Sutton et al. (1992), whereas optimal control assumes perfect knowledge, RL uses approximations and data-driven techniques.

- write about Value learning, and shortly name traditional methods (multiarmedbandit, DP, Temporal difference learning)

2.2. Robot Learning

The sub-field where reinforcement learning and machine learning intersect with robotics is called *Robot Learning*. It aims to bridge the gap between programmed robots, with fine tuned controllers and fully autonomous robots. As proposed in Deisenroth et al. (2013), robot control can be modeled as a reinforcement learning problem.

The state space \mathbf{x} in robotic tasks is high dimensional and made up of the internal state of the robot (e.g., joint position, body position/ orientation) and external state (e.g. object locations, lighting). The true state is not observable and also not noise free. The Robot chooses its next motor control u according to a policy π . This policy π may be deterministic $a = \pi(s)$ or stochastic $a \sim \pi(s, a) = P(a|s)$. The motor commands u alter the state according to the probabilistic transition function $p(\mathbf{x}_{t+1} | x_t, u_t)$. This transition function is not known, in model-based policy search this function is learned from data and used to improve the policy. Collectively the states and actions of the robot form a *trajectory* $\tau = (x_0, u_0, x_1, u_1, \dots)$ which is also called a *rollout* or a *path*. There has to be a numeric scoring system assessing the quality of the robots trajectory and returning a reward signal $R(\tau)$. For *episodic learning tasks* the task ends after a given number T of time steps. Then the accumulated reward $R(\tau)$ for a trajectory is given by

$$R(\tau) = r_T(x_T) + \sum_{t=0}^{T-1} r_t(x_t, u_t)$$

where r_t is an instantaneous reward function (e.g. a punishment for energy consumed) and r_T the final reward, when performing a reaching task this may take the form of a quadratic punishment term for deviation from the goal posture.

If we consider an infinite-horizon for an on-going task we get

$$R(\tau) = \sum_{t=0}^{\infty} \gamma^t r(x_t, u_t)$$

where $\gamma \in [0, 1)$ is a discount factor that discounts rewards further in the future.

Many tasks in robotics can thus be formulated as choosing a optimal control policy π^* that maximized the expected accumulated reward

$$J_{\pi} = \mathbb{E}[R(\tau)|\pi] = \int R(\tau) p_{\pi}(\tau) d\tau$$

where $R(\tau)$ defines the objectives of the task, and $p_{\pi}(\tau)$ is the distribution over trajectories τ .

Formulating robotic tasks in this way allows us to use methods from reinforcement learning.

2.2.1. Challenges

Reinforcement learning in general is a difficult problem, it can be harder to make RL work compared to supervised or unsupervised learning. One reason for this is that the reward signal may only be given occasionally and even then it may be unclear which of the agents actions were responsible for a certain reward signal.

Robotics is different compared to other fields where Reinforcement Learning is used extensively. First of all the states and actions of the robots in the real world are inherently continuous requiring us to deal with the resolution. In addition the state space can have a high dimensionality and working with real world systems on real hardware is costly and makes manual interventions necessary. Robots require algorithms to run in real-time and working with real sensors further introduces discrepancy between sensing and execution.

Most traditional methods from RL like TD-learning Sutton and Barto (2018) have been unfit for these particular requirements of robotic tasks. Stressing the importance of robotics as a testing ground for RL that demands new developments and innovative research. We will now discuss some problems encountered when applying RL to robotics, this treatment does only focus on a certain set of challenges and is not exhaustive.

Curse of Dimensionality

The term ‘‘Curse of Dimensionality’’ was coined by Bellman (1957) when he explored optimal control in higher dimensions and encountered an exponential explosion of the states and actions. For example, for the evaluation of our algorithm we run a simulation of a simple

planar reaching task with a 5 link robot arm using DMPs for policy representation and get a 25 state dimension. Especially modern anthropomorphic robots tend to have many degrees of freedom.

The agent in RL generally needs to collect data throughout the entire state-space to ensure global optimization, for robotics dealing with three dimensional space makes this very difficult. To alleviate this problem a widespread idea is to use expert demonstration to get a good initialization for the agents policy, this eliminates the need to search the entire search space, instead the agent can focus on locally optimizing the initialized policy. This concept of imitation learning focuses on the problem of “learning from demonstrations” which plays an important role for robotics Osa et al. (2018). Imitation learning will enable domain experts to teach motions and skills without special knowledge about robotics, which will be crucial when robots start making their way from factories into everyday life.

Curse of Real-world Samples

Robot hardware is expensive and suffers from wear and tear, making costly maintenance necessary. Hence safe methods for real robots should avoid big jumps in policy updates, as such sudden changes may result in unpredictable movements and consequently damage the robot. This problem is commonly referred to as *safe exploration*. Whereas in traditional reinforcement learning safe exploration does not receive much attention, it has become a key issue for robot learning Schneider (1997). Different external factors like temperature may change the robot dynamics and additional uncertainty from the sensors makes it difficult to reproduce and compare results.

Most task also require a “human in the loop” who either supervises the robot or resets the setup after one episode. Even if this can be avoided the learning speed simply cannot be sped up. For a single robot the training time has a natural limit and in total only relatively few executions can be completed. One method for gathering more data is “collective robot learning” described in Kehoe et al. (2015). The idea is for multiple robots to share their data on trajectories, policies and outcomes. Currently this seems only viable for large corporations with significant capital. Thus in robot learning the constraint of using only a small number of trials is given more weight than limiting memory consumption or computational complexity. Thus sample efficient algorithms are essential. Generally the state represented by sensors slightly lags behind the real state due to processing and communication delays. This is in stark contrast to most other reinforcement learning algorithms, which assume actions to take effect instantaneously.

Curse of Goal Specification

Defining a good reward function in robot reinforcement learning is difficult and often needs a lot of domain knowledge and expertise. There are certain trade-offs to keep in mind, for

example performing a powerful swing for a hitting task may yield high reward but may damage or shorten the life-time the robot. Reinforcement learning algorithms also tend to solve tasks in unintended ways exploiting the reward function in some unforeseen way.

Inverse reinforcement learning Russell (1998) provides an alternative to specifying the reward function manually. The goal of inverse reinforcement learning is to recover the unknown reward function from the expert's trajectories.

Still learning algorithms are rarely employed on robots for real daily usage. Also most algorithms are over fitted to a particular robot architecture and do not generalize to other robots easily. A large number of different methods exist, but in general there is no clear recipe for robot learning. Showing that robot learning still has a lot open problems, and will continue to grow as a research field. Even minor flaws or errors in the employed method can completely prevent success of learning. Nonetheless appropriately chosen algorithms and rewards functions already achieve promising results.

2.2.2. Policy search

Many traditional methods in RL try to estimate the expected long-term reward of a policy for each state \mathbf{x} and time step t , this leads to formulation of the value function $V_t^\pi(\mathbf{x})$. With the value function we can assess the quality of executing action \mathbf{u} in state \mathbf{x} . This assessment is used to directly compute the policy by action selection or to update the policy π . As value function methods struggle with the challenges in reinforcement learning, the main approach for robotics has become policy search. Policy search methods opposed to value-based methods use parameterized policies π_θ and search directly in the parameter space Θ . This allows using RL with high-dimensional continuous action spaces encountered in robotics by reducing the search space of possible policies. Policy search further allows the usage of predefined task-appropriate policy representations like Dynamic Movement Primitives Schaal et al. (2005), as well as easily integrating imitation learning for policy initialization.

Generally we can divide policy search into model-free and model-based and differentiate whether stochastic or deterministic trajectories are used. Model-free policy search uses trajectories from the robot directly for updating the policy. Model-based methods use the data from the robot to learn a model of the robot. This model is then used to generate trajectories that are used for policy updates.

- include figure for policy search taxonomy (recreated from Deisenroth et al. (2013)) also include examples for each type (REINFORCE, REPS, PILCO,...)

The most important concept is computing the policy updates. Both model-free and model-based policy search use policy gradients (PG), expectation-maximization (EM)-based updates, or information-theoretic insights (Inf.Th.).

In this thesis we will focus on model free policy search methods where the trajectories are generated by “sampling” from the robot. Due to their simplicity and ease of use of these algorithms are used more frequently than model-based policy search methods. Specifically we will focus on stochastic search algorithms, which are general black-box optimizers. They are used in a wide range of fields like operations research, machine learning and also policy search. Since these algorithms do not use any knowledge about the objective function it is straightforward to apply them to policy search in the episode-based formulation.

Using stochastic search algorithms we keep an upper-level policy $\pi_{\omega}(\theta)$ which selects the parameters of the actual control policy $\pi_{\theta}(\mathbf{u}|\mathbf{x})$ of the robot. With this instead of directly finding the parameters θ of the lower-level policy we want to find the parameter vector ω which defines a distribution over θ . We can then use this search distribution to directly explore the parameter space.

2.3. Kullback-Leibler (KL) Divergence

Many algorithms in policy search rely on the Kullback-Leibler Divergence, also known as the relative entropy, for controlling the difference between the the old and updated policy. Working with real robots additionally requires to perform safe exploration, big exploration steps, may result in damaging the hardware. Specifically, it measures the Shannon entropy of one distribution relative to the other.

$$KL(p||q) = \int p(\theta) \log \frac{p(\theta)}{q(\theta)} d\theta$$

Note that in general the relative entropy is not symmetric under interchange of the distributions p and q . In general $KL(p||q) \neq KL(q||p)$, therefore in a mathematical sense it is not strictly a distance.

The KL-bound has been used successfully for many algorithms (cite REPS, Natural gradient) It is widely used in Machine Learning algorithms. (cite..)

2.4. MORE Algorithm

The MORE algorithm is a stochastic search algorithm. It can be used for model-free policy search. The key idea is using information-theoretic policy updates by bounding the relative entropy (Kullback Leibler divergence) between two subsequent policies. We approximate the objective function with a quadratic surrogate model hence we can satisfy the KL-bound in closed form.

- TODO: include 3D figure of more algorithm

2.4.1. MORE Framework

The MORE algorithm maintains a search distribution (policy) over the parameter space of the objective function. We can use the method of constraint optimization to formulate an optimization problem for obtaining a new search distribution that maximizes the expected objective value while upper-bounding the KL-divergence and lower-bounding the entropy of the distribution.

$$\max_{\pi} \int \pi(\theta) \mathbf{R}_{\theta} d\theta, \quad (2.1)$$

$$\text{s.t. } \text{KL}(\pi(\theta) || q(\theta)) \leq \varepsilon, \quad (2.2)$$

$$H(\pi) \geq \beta, \quad (2.3)$$

$$1 = \int \pi(\theta) d\theta \quad (2.4)$$

Forming the Lagrangian we get

$$\mathcal{L}(\pi, \eta, \omega) = \int \pi(\theta) \mathcal{R}_{\theta} d\theta + \eta \left(\varepsilon - \int \pi(\theta) \log \frac{\pi(\theta)}{q(\theta)} d\theta \right) - \omega \left(\beta + \int \pi(\theta) \log(\pi(\theta)) d\theta \right) \quad (2.5)$$

The solution is

$$\pi(\theta) \propto q(\theta)^{\eta/(\eta+\omega)} \exp \left(\frac{\mathcal{R}_{\theta}}{\eta + \omega} \right) \quad (2.6)$$

it depends on the quadratic and linear term of the surrogate model and the Lagrangian multipliers. These can be obtained by minimizing the dual problem:

$$g(\eta, \omega) = \eta \varepsilon - \omega \beta + (\eta - \omega) \log \left(\int q(\theta)^{\frac{\eta}{\eta+\omega}} \exp \left(\frac{\mathcal{R}_{\theta}}{\eta + \omega} \right) d\theta \right) \quad (2.7)$$

We get the closed form solution for the equation (see full derivation in Appendix).

With these equations we can formulate an iterative algorithm for updating the policy.

Entropy constraint

The bound β is defined such that relative difference between the entropy of the policy $H(\pi)$ and a minimum exploration policy $H(\pi_0)$ is decreased for a certain percentage:

$$H(\pi) - H(\pi_0) \geq \gamma(H(q) - H(\pi_0)) \rightarrow \beta = \gamma(H(q) - H(\pi_0) + H(\pi_0))$$

2.4.2. Surrogate Model

The key idea of the MORE algorithm is using a surrogate model of the objective function for satisfying the bound set by the optimization problem. It has proven to be sufficient to use a quadratic model, since the exponent of Gaussian distribution is also quadratic, it is not possible to exploit the information of a more complex surrogate model. This concept of using a surrogate model has been used in numerical optimization (cite from book numerical optimization)

The original approach of MORE is based on weighted Bayesian linear model with dimensionality reduction.

2.4.3. Analytic Solution for Dual-Function and Policy

Assuming we are given a quadratic surrogate model

$$R_\theta \approx \theta^T R \theta + \theta^T r + r_0$$

we can now solve the dual function in closed form.

$$g(\eta, \omega) = \eta \varepsilon - \beta \omega + \frac{1}{2} (\mathbf{f}^T \mathbf{F} \mathbf{f} - \eta \mathbf{b}^T \mathbf{Q}^{-1} \mathbf{b} - \eta \log |2\pi \mathbf{Q}| + (\eta + \omega) \log |2\pi(\eta + \omega) \mathbf{F}|)$$

Assuming π is Gaussian we get the solution, and can update the mean and covariance matrix of the search distribution.

$$\pi_{t+1} = \mathcal{N}(\mu_{t+1}, \Sigma_{t+1}) \quad (2.8)$$

$$\mu_{t+1} = (\eta \Sigma_t^{-1} \mu_t + r) / (\eta + \omega) \quad (2.9)$$

$$\Sigma_{t+1} = (\eta \Sigma_t^{-1} + R) / (\eta + \omega) \quad (2.10)$$

With these equations we can iteratively update our search distribution. It is important to note that MORE can be used as a stochastic search algorithm, a “black box solver” because

only the samples and objective function value is needed to update the search distribution. In Robotics it can be used to learn motor tasks by using Dynamic Movement Primitives (DMPs) as a policy representation.

2.5. Bayesian Filtering

TODO: rewrite, because currently this part to closely resembles Särkkä (2013).

Optimal filtering is concerned with estimating the state of a time-varying system which is only partially observed through noisy measurements. *Bayesian Filtering* refers to the Bayesian way of formulating optimal filtering.

In general the term “Bayesian” refers to inference methods that represent “degrees of certainty” using probability theory, and basically use **Bayes’ rule** to update the degree of certainty given data.

Filtering methods are important in robotics for handling the uncertainty in the environment. Sensing and modeling the state of the robot itself. Methods like the Kalman Filter are extensively used since we are dealing with noisy sensors.

Also the data is very limited and when the robots needs to make decisions it can only work with small amounts of data, therefore therefore Bayesian methods are a great fit and are widely used (cite..)

In general we solve linear Gaussian and non-linear/non-Gaussian state space models with the Bayesian filtering equations.

- optimal recursive estimators first presented for linear systems (simplicity)
- most natural optimality criterion is the least squares optimality, minimum mean squared error
- statistical inversion problem: estimate hidden states from observed measurements in Bayesian sense we want to compute joint posterior distribution. Straightforward application of Bayes rule
- Bayesian viewpoint sees probabilities as degree of belief.
- Bayes rule can be used to update our information about a quantity
- Bayesian probability building blocks

2.5.1. Building Blocks of Bayesian models

In general all Bayesian models have some basic components to them.

prior distribution

Encodes the information on parameter θ before seeing any observations. When we are uncertain about our prior information we can choose a high variance of the distribution or use a non-informative prior (which imposes the minimal amount of structure on the data).

$$p(\theta) = \text{information on parameter } \theta \text{ before seeing any observations}$$

measurement model

Models the relationship between true parameters and the measurements.

$$p(y|\theta) = \text{distribution of observation } y \text{ given the parameters } \theta$$

posterior distribution

The conditional distribution of the parameters given the observations. It represents the updated belief about the parameters after obtaining the measurements. It can be computed by using Bayes' rule.

$$p(\theta|y) = \frac{p(y|\theta)p(\theta)}{p(y)} \propto p(y|\theta)p(\theta)$$

The normalization constant can be computed by integrating θ out.

$$p(y) = \int p(y|\theta)p(\theta)d\theta$$

-TODO: create graphic (2d or 3d) for merging prior with observation to get posterior (show with multivariate Gaussian distribution)

2.5.2. Gaussian probability function

A random variable $x \in \mathbb{R}^n$ has a Gaussian distribution with mean $m \in \mathbb{R}^n$ and covariance $P \in \mathbb{R}^{n \times n}$ if its probability density has the form

$$N(x|m, P) = \frac{1}{(2\pi)^{n/2} |P|^{1/2}} \exp \left(-\frac{1}{2} (x-m)^T P^{-1} (x-m) \right)$$

where $|P|$ is the determinant of the matrix P .

2.5.3. Least squares

To grasp the Bayesian Filtering viewpoint we will now look at a simple regression problem from a Bayesian perspective and derive the least squares solution.

We consider a simple linear regression problem,

$$y_k = \theta_1 + \theta_2 x_k + \varepsilon_k$$

note that the response y_k is only dependent on one regressor x_k .

Writing this in probabilistic terms we get:

$$p(y_k|\theta) = N(y_k|H_k\theta, \sigma^2) \tag{2.11}$$

$$p(\theta) = N(\theta|m_0, P_0) \tag{2.12}$$

Here $H_k = (1 x_k)$ and $N(\cdot)$ is the Gaussian probability density function. The batch solution can then be easily obtained by application of Bayes' rule

$$\begin{aligned} p(\theta|y_{1:T}) &\propto p(\theta) \prod_{k=1}^T p(y_k|\theta) \\ &= N(\theta|m_0, P_0) \prod_{k=1}^T N(y_k|H_k\theta, \sigma^2) \end{aligned}$$

Because the prior and likelihood are Gaussian, the *posterior distribution* will also be Gaussian

$$p(\theta|y_{1:T}) = N(\theta|m_t, P_t)$$

Mean and covariance obtained by completing the quadratic form in the exponent: - TODO: Do calculation in Appendix, (use canonical form)

$$\mathbf{m}_T = \left[\mathbf{P}_0^{-1} + \frac{1}{\sigma^2} \mathbf{H}^T \mathbf{H} \right]^{-1} \left[\frac{1}{\sigma^2} \mathbf{H}^T \mathbf{y} + \mathbf{P}_0^{-1} \mathbf{m}_0 \right] \quad (2.13)$$

$$\mathbf{P}_T = \left[\mathbf{P}_0^{-1} + \frac{1}{\sigma^2} \mathbf{H}^T \mathbf{H} \right]^{-1} \quad (2.14)$$

This gives us the batch least squares solution.

- TODO talk more about how to why this is ordinary least squares?

2.5.4. Recursive least squares

With the Bayesian formulation we can now introduce a recursive solution to the regression problem, resulting in a recursive least squares algorithm.

For this we have to assume the next step only depends on the previous state, (markovian assumption). The main idea is using the previous posterior as the prior in the estimation step. The measurements are assumed to be conditionally independent.

To get the *recursive Bayesian solution* we proceed as follows: In the beginning of estimation all the information about the parameter θ are contained in the prior distribution $p(\theta)$. The measurements are obtained on at a time, to obtain the next posterior distribution we use the measurement and the information from the previous posterior distribution as the prior.

$$\begin{aligned} p(\theta|y_1) &= \frac{1}{Z_1} p(y_1|\theta) p(\theta) \\ p(\theta|y_{1:2}) &= \frac{1}{Z_2} p(y_2|\theta) p(\theta|y_1) \\ &\vdots \\ p(\theta|y_{1:T}) &= \frac{1}{Z_T} p(y_T|\theta) p(\theta|y_{1:T-1}) \end{aligned}$$

Normalization term: $Z_k = \int p(\theta) p(y_k|\theta) d\theta$

Using this approach to solve our simple linear regression model we get

$$p(\theta|y_{1:k}) \propto p(y_k|\theta) p(\theta|y_{1:k-1}) \quad (2.15)$$

$$\propto \mathbf{N}(\theta|\mathbf{m}_k, \mathbf{P}_k) \quad (2.16)$$

from which we get:

$$\mathbf{m}_k = \left[\mathbf{P}_{k-1}^{-1} + \frac{1}{\sigma^2} \mathbf{H}_k^T \mathbf{H}_k \right]^{-1} \left[\frac{1}{\sigma^2} \mathbf{H}_k^T \mathbf{y}_k + \mathbf{P}_{k-1}^{-1} \mathbf{m}_{k-1} \right] \quad (2.17)$$

$$\mathbf{P}_k = \left[\mathbf{P}_{k-1}^{-1} + \frac{1}{\sigma^2} \mathbf{H}_k^T \mathbf{H}_k \right]^{-1} \quad (2.18)$$

TODO: do calculation with matrix inversion lemma (in appendix?)

By using the *matrix inversion lemma* the covariance calculation can be written as

$$P_k = P_{k-1} - P_{k-1} H_k^T [H_k P_{k-1} H_k^T + \sigma^2]^{-1} H_k P_{k-1}$$

By introducing temporary variables S_k and \mathbf{K}_k the calculation of the mean and covariance can be written in the form

$$\begin{aligned} S_k &= \mathbf{H}_k \mathbf{P}_{k-1} \mathbf{H}_k^T + \sigma^2 \\ \mathbf{K}_k &= \mathbf{P}_{k-1} \mathbf{H}_k^T S_k^{-1} \\ \mathbf{m}_k &= \mathbf{m}_{k-1} + \mathbf{K}_k [y_k - \mathbf{H}_k \mathbf{m}_{k-1}] \\ \mathbf{P}_k &= \mathbf{P}_{k-1} - \mathbf{K}_k S_k \mathbf{K}_k^T \end{aligned}$$

Here \mathbf{K} is the Kalman gain that controls the weight we give to new measurements. This is actually a special case of The Kalman filter update equations. Since here the parameters θ are assumed to stay constant there is no stochastic dynamics model used for the prediction step.

For estimating the surrogate model for the MORE algorithm we have to deal with dynamic parameters instead. In the next chapter we will therefore introduce a drift model for the parameters.

Drift model

We assume the parameter θ performs a *Gaussian Random Walk* after each time step.

$$p(y_k | \theta) = N(y_k | \mathbf{H}_k \theta_k, \sigma^2) \quad (2.19)$$

$$p(\theta_k | \theta_{k-1}) = N(\theta_k | \theta_{k-1}, \mathbf{Q}) \quad (2.20)$$

$$p(\theta_0) = N(\theta_0 | \mathbf{m}_0, \mathbf{P}_0) \quad (2.21)$$

Where \mathbf{Q} is the covariance of the random walk. Starting with the distribution

$$p(\theta_{k-1} | y_{1:k-1}) = N(\theta_{k-1} | \mathbf{m}_{k-1}, \mathbf{P}_{k-1})$$

the joint distribution of θ_k and θ_{k-1} is

$$p(\theta_k, \theta_{k-1} | y_{1:k-1}) = p(\theta_k | \theta_{k-1}) p(\theta_{k-1} | y_{1:k-1})$$

The distribution of θ_k given the measurement history up to time step $k-1$ can be calculated by integrating over θ_{k-1} :

$$p(\theta_k | y_{1:k-1}) = \int p(\theta_k | \theta_{k-1}) p(\theta_{k-1} | y_{1:k-1}) d\theta_{k-1}$$

This relationship is sometimes called *Chapman-Kolmogorov equation*. The result of the marginalization is Gaussian

$$p(\theta_k | y_{1:k-1}) = \mathcal{N}(\theta_k | m_k^-, P_k^-)$$

with

$$m_k^- = m_{k-1} \quad (2.22)$$

$$P_k^- = P_{k-1} + Q \quad (2.23)$$

Now we simply replace P_{k-1} with P_k^- in our update equations:

$$\begin{aligned} S_k &= \mathbf{H}_k \mathbf{P}_k^- \mathbf{H}_k^T + \sigma^2 \\ \mathbf{K}_k &= \mathbf{P}_k^- \mathbf{H}_k^T S_k^{-1} \\ \mathbf{m}_k &= \mathbf{m}_k^- + \mathbf{K}_k [y_k - \mathbf{H}_k \mathbf{m}_k^-] \\ \mathbf{P}_k &= \mathbf{P}_k^- - \mathbf{K}_k S_k \mathbf{K}_k^T \end{aligned}$$

These equations are a special case of the Kalman Filter, only doing a adding noise to the parameters in the prediction step and using an identity matrix as the state transition model. There is no obvious state transition model for the general case, maybe some method based on an momentum approach on the differences in subsequent estimated parameters. Otherwise this could be engineered to a certain objective function, but this is part of future work.

Chapter 3.

Related Work

3.1. Policy Search Algorithms

Generally model-free policy search algorithms have been used for their ease of use, at often being black box optimizers.

For model-free policy search the agent generates trajectories evaluates them and uses the reward and the trajectories to iteratively update the policy such that the expected reward is maximized.

There are different methods for computing the update of the policy, common strategies are based on gradient ascent Peters and Schaal (2006).

- natural gradient by NES (update direction of parameters like standard gradient but still in bound by KL-divergence)
- evolutionary strategies
- expectation-maximization algorithms

The MORE algorithm is part of information theoretic approaches to PS. Information theoretic approaches use the Kullback-Leibler Divergence in an optimization problem to bound the distance of the old policy to the new one. There are different techniques to solve the Integrals that arise when optimizing the problem. The REPS algorithm proposed in Peters et al. (2010)

uses a sample-based approximation for the KL-divergence. Using Taylor-expansions for the KL-divergence resulted in the natural evolutionary strategies (NES) Wierstra et al. (2014).

Some additions to the original MORE have been explored like formulating a contextual version CMORE Tangkaratt et al. (2017), enabling learning from high-dimensional context variables like camera images.

3.2. Surrogate modeling

Model-based methods for approximating the objective function with a local surrogate have been used in derivative free optimization Nocedal and Wright (2006). Local surrogate models have been used in trust region methods like Powell (2009). These methods maintain a point estimate and a trust region around this point instead of a search distribution, they update the point estimate by optimizing the surrogate and staying in the trust region.

Recursive estimation with methods like the Kalman Filter is extensively used in robotics, mostly to deal with noisy measurements. Probabilistic methods from a Bayesian especially with a focus on decision making is introduced in Thrun (2002).

Chapter 4.

Recursive Surrogate-Modeling for MORE

In this chapter we will first motivate the idea of using recursive filtering for estimating the surrogate model. Then we formulate the surrogate estimation as a regression problem and derive the recursive least squares approach for the drift model. Then we formulate our version of the MORE algorithm with recursive estimation of the surrogate model. Next we discuss various data preprocessing techniques we use or explored.

4.1. Motivation

We want to be sample efficiently, because of high cost of real world samples.

- do 3D example on simple sphere function

Previously estimating the surrogate model for the MORE algorithm has been solely based on samples and corresponding rewards. There was a sample pool used to be more sample efficient. By using an recursive filtering approach we want to utilize the information of previous surrogate models instead of using only samples and rewards for estimation.

Since the MORE algorithm uses the KL-bound to limit the distance from the previous search distribution the subsequent surrogate models are also locally correlated.

The surrogate model has the form

$$\mathbf{M}(\mathbf{x}) = -\frac{1}{2}\mathbf{x}^T \mathbf{A} \mathbf{x} + \mathbf{x}^T \mathbf{b} + c \quad (4.1)$$

With a quadratic term A , a spd(semi positive definite) matrix, a linear term b and a scalar c .

- TODO: include 2D contour line

The learned quadratic models are locally correlated and thus we can use recursive filtering algorithms to exploit the information of previous models at the current time step, this has the potential to result in using less samples and also reducing the overall runtime of the algorithm.

- do 3 figures: first mean and samples, then resulting surrogate, then updated mean

4.2. Surrogate Model Estimation

4.2.1. Regression problem

If we formulate this as a regression problem we get:

$$y_k = \mathbf{x}^T \mathbf{A} \mathbf{x} + \mathbf{x}^T \mathbf{b} + c + \varepsilon_k$$

So we want to estimate the surrogate model parameters $\theta = (\mathbf{A}, \mathbf{b}, c)$ from the samples \mathbf{x} and the rewards y . We also assume that the measurement noise ε_k is zero mean Gaussian $\varepsilon_k \sim N(0, \sigma^2)$ distributed.

In our algorithm we solve this problem by using the Recursive Least squares equations Section 2.5.4.

4.3. Data Preprocessing Techniques

TODO: move this whole section into evaluation/experiments?

To improve the performance of the algorithm we examined several data processing techniques. Here talk theoretically about the methods and in evaluation part show experiments about different techniques (for example show whitened model parameters)

4.3.1. Whitening

Whitening is common data preprocessing method in statistical analysis to transform a correlated random vector into an uncorrelated one Kessy et al. (2018).

We employ whitening for our algorithm, because uncorrelated random variables often greatly simplify multivariate data analysis (cite).

Whitening is a linear transformation that converts a d -dimensional random vector $\mathbf{x} = (x_1, \dots, x_d)^T$ with mean $E(\mathbf{x}) = \boldsymbol{\mu} = (\mu_1, \dots, \mu_d)^T$ and positive definite $d \times d$ covariance matrix $\text{var}(\mathbf{x}) = \boldsymbol{\Sigma}$ into a new random vector

$$\mathbf{z} = (z_1, \dots, z_d)^T = \mathbf{W}\mathbf{x} \quad (4.2)$$

of the same dimension d and with unit diagonal “white” covariance $\text{var}(\mathbf{z}) = \mathbf{I}$. The square $d \times d$ matrix \mathbf{W} is called the whitening matrix.

The whitening transformation defined in Equation Equation (4.2) requires the choice of a suitable whitening matrix W . Since $\text{var}(\mathbf{z}) = \mathbf{I}$ it follows that $\mathbf{W}\boldsymbol{\Sigma}\mathbf{W}^T = \mathbf{I}$ and thus $\mathbf{W}(\boldsymbol{\Sigma}\mathbf{W}^T) = \mathbf{W}$, which is fulfilled if \mathbf{W} satisfies the condition

$$\mathbf{W}^T \mathbf{W} = \boldsymbol{\Sigma}^{-1}$$

This constrain does not uniquely determine the whitening matrix \mathbf{W} , instead given $\boldsymbol{\Sigma}$ there are infinitely many possible matrices \mathbf{W} , because it allows for rotational freedom.

We used *Cholesky whitening* which is based on Cholesky factorization of the precision matrix $\mathbf{L}\mathbf{L}^T = \boldsymbol{\Sigma}^{-1}$, which leads to the whitening matrix:

$$\mathbf{W}^{\text{Chol}} = \mathbf{L}^T$$

.

If the cholesky whitening is unsuccessful we fall back to standardizing the random vector meaning $\text{var}(\mathbf{z}) = 1$ but the correlations are not removed.

- TODO: include example samples before and after whitening

4.3.2. Sample pool

The original MORE algorithm and other policy search algorithms (REPS) we use a sample pool.

Theoretically we would want to avoid using a sample pool and instead use the information from past samples in the form of the previously predicted model parameters. But with our algorithm using no sample pool led to unsatisfactory estimation results.

We increased the model noise for older samples, encoding the fact that older samples should have a higher covariance, meaning we are less certain about them. We implemented this by simply adding a constant amount to the covariance of the RLS equations for each time a sample is used. For higher dimensional task this is around 5-10 times. Still the use of a sample pool is theoretically unsound and problematic, instead a accurate state transition model could be introduced.

4.3.3. Normalization

Original MORE approach uses standard score normalization, doing it in a batch way at each iteration for all samples in the sample pool.

This is not fit for recursive estimation. - different types of reward functions difficulties: - rosenbrock (high range of values) - sharp spikes, in rewards (punishing term for reaching task)

- simply online calculation of mean and var for normalization is not a good fit (include plots and results of investigation)
- instead use moving average to calculate mean and var only of window (discard old data)

We can use a moving mean average to normalize the data for example the weighted moving average or exponential moving average

moving average - Write analysis about using different methods and showing that the mean is not what we want but the weighted average

4.4. MORE with Recursive Surrogate-Modeling

Algorithmus 1 : MORE Algorithm with Recursive Surrogate-Modeling

Input : Parameters ε and β , initial distribution,
 K number of iterations, N samples per iteration

```

for  $k = 1, \dots, K$  do
  for  $n = 1, \dots, N$  do
    Draw parameters  $\theta_n \sim \pi$ 
    Execute task with  $\theta_n$  and receive  $R(\theta_n)$ 
  end
  begin Learn quadratic model with Recursive Least Squares
    for  $n = 1, \dots, N$  do
      Whitening:  $\mathbf{W}\theta_n$ 
      Compute surrogate model parameters with RLS
    end
  end
  Minimize dual function  $g(\eta, \omega)$  using Eq.
  Update search distribution  $\pi$  using Equation (2.8)
end

```

- Add 3d Graphic of search distribution mean path

4.4.1. Higher model noise for older samples

Using the recursive least squares algorithm we had to maintain a sample pool, because otherwise the resulting estimation was not satisfactory. From a theoretic perspective this sloppy and ideally we would refrain from using the sample pool.

We introduced a simple counter for each sample, indicating how many times the sample has been used. Then for older samples (higher counter) we increased the model noise proportionally. We tried a linear relationship between the counter and the increased covariance for the model parameters.

$$new_cov = constant_cov_matrix + w * counter * \mathbf{I}$$

- TODO: try different weightings: exponential

This improved our results on the rosenbrock test function compared to the RLS version with constant model noise for all samples. In each MORE iteration first the oldest samples from the sample pool are used, meaning the recent samples with lowest model noise are processed last by RLS.

4.4.2. Moving back to prior

One technique we investigated is - to prior calculation (optimal for whitened space)

- set up equation

4.4.3. State Transition Model

Let us quickly state the full Kalman filter equations, the derivation can be found in the appendix (TODO ref)

Prediction step:

Update step:

Introducing a an accurate state transition model to incorporate the prediction step of the kalman filter would be ideal. We tried some simple momentum approaches based on the differences of subsequent surrogate model parameters, but those did not yield good results. Our Recursive least squares algorithm is a special case of these equations using an state transition model matrix $\mathbf{A} = \mathbf{I}$.

Chapter 5.

Evaluation

This chapter will introduce the setup for the experiments and the tasks.

5.1. Setup

All algorithms were implemented in python, the optimization algorithm for MORE is Low-storage BFGS from nlopt¹. The clusterwork2 (cw2) framework, developed at the ALR group, was used for running experiments. The experiments were conducted on a machine with two 8-core AMD Ryzen 2700X processores clocked at 2.6 GHz and 31GB of RAM.

- code is uploaded to gitlab/github (archive repo)

5.1.1. Hyperparameter search

We did a mixture of manual hyperparameter search and grid search with the cw2 framework. At a later stage we incorporated optuna for hyperparameter optimization.

¹<https://nlopt.readthedocs.io/en/latest/>

5.2. Experiments

We now evaluate the performance of our algorithms on various problems.

5.2.1. Test Functions for Optimization

Based on Molga and Smutnicki (2005).

Figure 5.1 shows the rosenbrock function, a uni-model optimization function. The global minimum $f(\mathbf{x}) = 0$. In the experiments the mean of the initial distribution has been chosen randomly.

$$f(x) = \sum_{i=1}^{n-1} [100(x_{i+1} - x_i^2)^2 + (1 - x_i)^2]$$

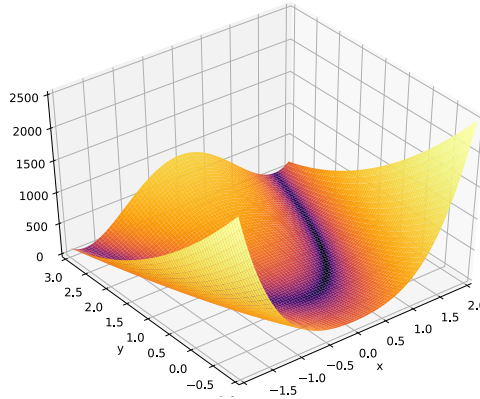


Figure 5.1.: rosenbrock function

In figures Figure 5.2 and Figure 5.3 we show the best results achieved on 5 dimensional and 15 dimensional rosenbrock. We observe that the our RLS approach uses the least amount of samples. For 5 dimensional rosenbrock the BLR approach is omitted due to considerably worse performance. One observation to make is that RLS converges only to around $1e-9$, while the LS approach manages to

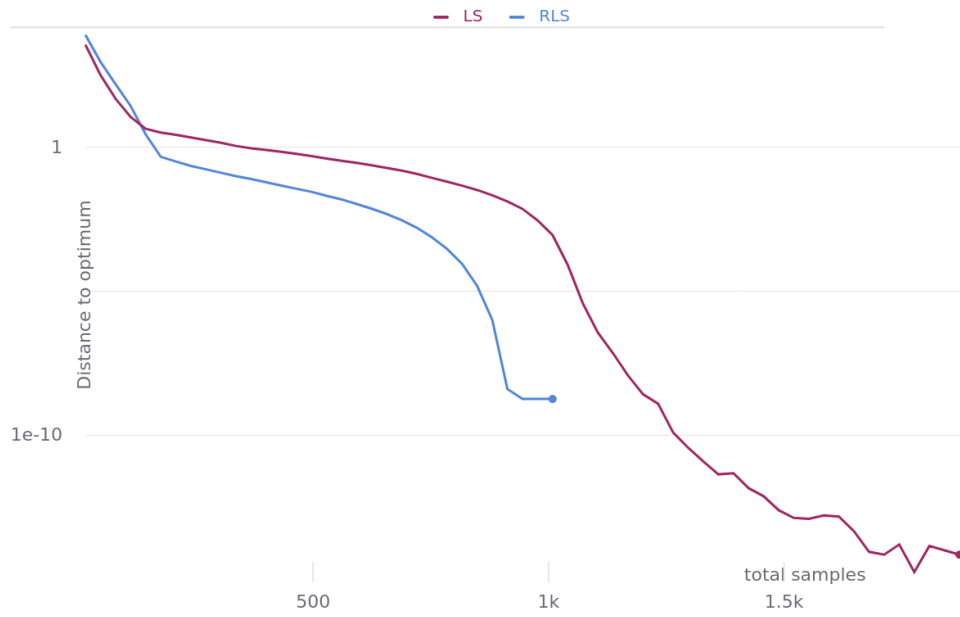


Figure 5.2.: This figure shows the median of (10 runs) for RLS and LS on 5 dimensional Rosenbrock

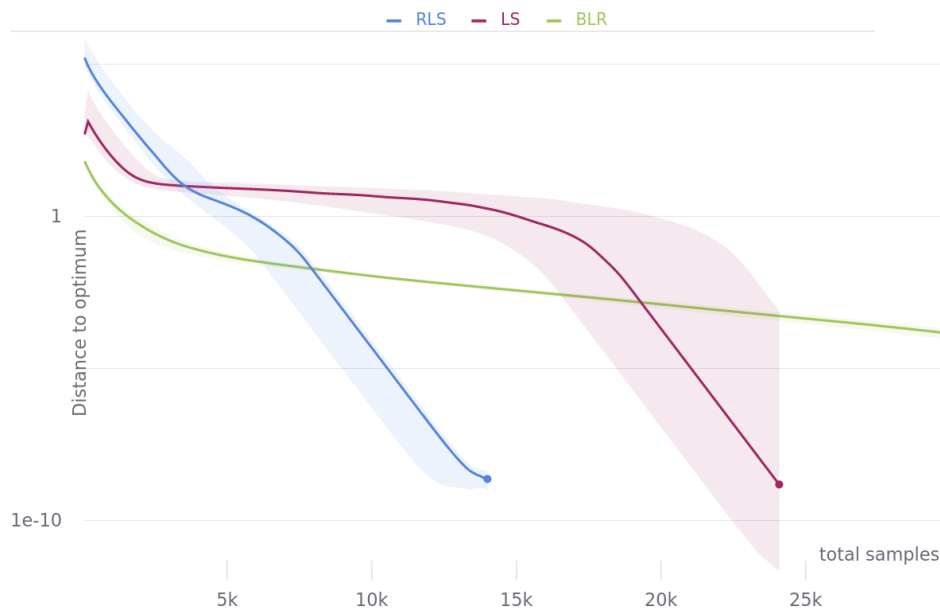


Figure 5.3.: This figure shows the median of (10 runs) for RLS, LS and BLR (original MORE) on 15 dimensional Rosenbrock

5.2.2. Planar Reaching Tasks

Dynamic Movement Primitives

- dynamic system-based motor primitives based on work [Ijspeert 2002, Schaal 2003,2007] improved imitation and reinforcement learning
- represent both discrete point-to-point movements as well as rhythmic motion with motor primitives
- first system canonical system (phase variable)
- we can build complex motor skills from basic primitives, its based on dynamical systems
- discrete movement primitives using only a single first order system
- key advantage the second system is linear in the shape parameters and can be efficiently learned
- initialized with imitation learning
- DMPs as policy representation

5.2.3. Via Point Reaching Task

We used a 5 link robot, similar to MORE setup. Therefore 25 dimensions of the problem, which should be at viapoint (1,1) at time step 100 and at viapoint (5,0) at timestep 200. In Figure 5.4 we see the task solved by the original MORE algorithm.

5.2.4. Hole Reaching Task

5 link robot, that has to reach into hole at [2,0] without collision with ground or walls.

5.2.5. Tests for Preprocessing Methods

- TODO: place this into appendix?

Tests on data preprocessing methods, different normalization techniques. Whitening very important, making it possible to track the model parameters more easily.

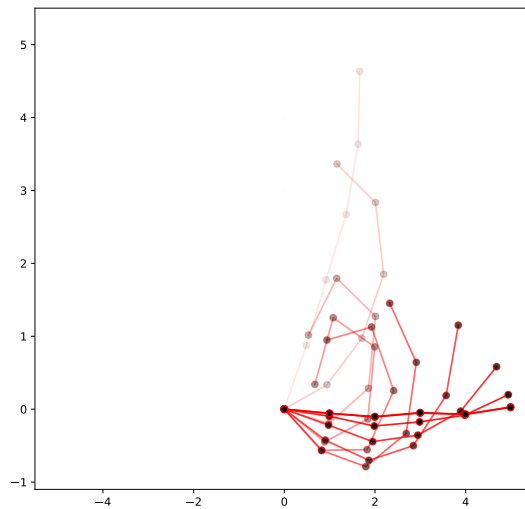


Figure 5.4.: This figure shows the movement resulting from the policy learned by BLR for the viapoint reaching task. The postures of the resulting motion are shown as overlay, where darker postures indicate a posture which is close in time to the hole.

- compare normalization techniques (moving average, batch normalization)
- best performance comparison (more iterations)
- best performance comparison of different methods (total samples)
- test influence of KL bound
- test influence of whitening, normalization?
- test influence of noise of drift model

5.3. Evaluation

- some theoretical problems: sample pool, model noise
- improve sample efficiency of rosenbrock and reaching task, better than original MORE and beating CMA-ES benchmark

Chapter 6.

Conclusion and Future Work

6.1. Conclusion

- managed to improve sample efficiency on simple test functions

We achieved some good results on the rosenbrock test function considerably improving the sample efficiency. Nonetheless for the planar reaching tasks we did not manage to achieve a significant improvement. Also our proposed algorithm did not work with multi modal and noisy objective functions.

- One main problem seems to be tuning the recursive estimation for one specific task.

The data seems to be difficult to Still

- inexperience of author with kalman filter (setting up matrices, choosing good state transition model)

6.2. Future Work

- remove sample pool
- usage of state transition model (kalman filter prediction step)

- make learning more stable for reaching tasks (safe exploration)
- learning the model noise: run original MORE approach and do backpropagation with pytorch over the weights

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Appendix A.

Appendix

A.1. MORE: Closed Form Solution

- add derivation of lagrangian function for MORE with clear steps

A.2. Kalman Filter Derivation

