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#### Abstract

This tutorial introduces Gaussian Process Regression as a method to explore (learn) and exploit (maximize) unknown functions. Gaussian Process Regression is a popular non-parametric Bayesian approach to sequential function learning problems. This tutorial aims to provide an accessible introduction to these techniques. We will introduce and define Gaussian Processes as a distribution over functions used for Bayesian inference and demonstrate different applications thereof. Examples will focus on pure exploration within optimal design problems, bandit-like exploration-exploitation scenarios, and on scenarios with additional constraints, for example safe explorations, where the goal is to never sample below a certain threshold. Software pointers will be provided.

Keywords: Gaussian Process, Exploration-Exploitation, Bandit Problems

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## 1. Introduction

Many applied research problems require us to explore (learn) or exploit (optimize) an unknown function that maps inputs to outputs  $f: x \to y$ (Mockus, 2010). No matter if we try to find a function that describes participants' behavior (Cavagnaro, Aranovich, McClure, Pitt, and Myung, 2014), estimate parameters of psychological models (Wetzels, Vandekerckhove, Tuerlinckx, and Wagenmakers, 2010), try to sequentially optimize stimuli in an experiment (Myung and Pitt, 2009), or model how participants themselves learn to interact with their environment (Meder and Nelson, 2012). Often times the underlying function might be unknown, hard to evaluate analytically, or other requirements such as design costs might complicate the process of information acquisition. In these situations, Gaussian Process regression can serve as a useful multi-purpose tool towards learning and optimization (Rasmussen, 2006). Gaussian Process regression is a non-parametric Bayesian approach (Gershman and Blei, 2012), applied to regression problems. It can capture many different relations between inputs and outputs by utilizing potentially infinite number of parameters and letting the data decide directly upon the level of complexity through the means of Bayesian inference (Williams, 1998). 19 This tutorial will introduce Gaussian Process Exploration-Exploitation 20 as an approach towards learning and optimization of unknown functions. It is intended for a general readership and contains many practical examples and high level explanations. It consists of 5 main parts: The first part will brush over the mathematical underpinnings of Gaussian Process regression. The second part will show how Gaussian Processes can be used in problems of optimal experimental design, when the goal is pure exploration,

that is to learn a function as well as possible. The third part will describe
how Bayesian Optimization (exploitation) with Gaussian Processes works.
In the fourth part, we will talk about ways of utilizing Gaussian Process
exploration-exploitation methods in situations with additional requirements
and show one example where the goal is to avoid areas that are below a certain threshold. We will conclude by sketching out the possibility of Gaussian
Processes as a low level description of cognitive processes.

## 2. Problem Statement

Imagine you have a function  $f: x \to y$  that maps an input x to an output y. For both an *exploration* as well as an *exploration-exploitation* scenario the goal is to make stepwise queries as wisely as possible. Querying a function means that you determine a point  $x^*$  that you want to observe next and then see an output of the function  $y = f(x^*) + \epsilon$  with some noise  $\epsilon$  at that point. Normally it is assumed that this noise is independent and identically distributed, following a Gaussian distribution  $\epsilon \sim \mathcal{N}(0, \sigma^2)$ .

## 2.1. Exploration

The goal within exploration is to chose  $x^*$  in a way that allows you to learn an unknown function as well as possible. This is sometimes referred to as an optimal experimental design problem (Goos and Jones, 2011). More formally, the task is to find an informative set of points  $\mathcal{A} \subset \mathcal{D}$ , that is a sequential query  $\mathcal{A}$  out of the design set  $\mathcal{D}$ , chosen to maximize information gain over all potential steps. Generally, the information gain is measured as the difference in entropy between the state before a sample point was queried and afterwards as shown in Equation 1.

$$I(y_A; f) = H(y_A) - H(y_A|f) \tag{1}$$

In the discrete case, the entropy H can be calculated as shown in Equation 2.1.

$$H(X) = \sum_{i}^{n} p(x_i) \log p(x_i)$$

Finding the absolute information gain maximizer is generally NP-hard (Ko, Lee, and Queyranne, 1995), which means that one can never know the best strategy of how to pick all of the observations over time to certainly reach the maximum information gain at the end as this requires an exponentially increasing problem space to be evaluated. Optimal design problems are ubiquitous in psychology. No matter if we have to find out how participants integrate information (Borji and Itti, 2013) or if we have to find stimuli in order to distinguish between different models (Cavagnaro, Myung, Pitt, and Kujala, 2010), often times we have to try and learn an underlying function that we do not know a priori and do so both adaptively and efficiently. As we will see later, Gaussian Process

## 2.2. Exploration-Exploitation

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The goal within exploration-exploitation tasks is to both learn and op-64 timize a function. Here, it is essentially the intention to learn a function quickly so that we then can find the maximum of that function and keep on exploiting it; exploiting means entering the input that produces the highest 67 output. This is normally measured by regret, the distance between what your current guess of the maximum has produced and what the best argument would have produced. If you are sequentially producing an estimate of

exploration algorithms provide us with an excellent tool for such tasks.

what you currently think might be the maximum  $x^*$ , then it is possible to measure regret R by the difference to what the actual best argument  $x_{\text{max}}$ would have produced.

$$R = \sum_{i=1}^{T} r_i$$
$$= \sum_{i=1}^{T} f(x_{\text{max}}) - f(x^*)$$

Problems where we have to optimize an unknown function are very common; be it to estimate parameters of a complex model (Snoek, Larochelle, and Adams, 2012) or finding the stimulus that produces the maximal response in an experiment (Daunizeau, Preuschoff, Friston, and Stephan, 2011), many problems require us to explore and exploit functions. Sometimes this approach is also called Bayesian Optimization (Brochu, Cora, and De Freitas, 2010).

## 3. Gaussian Processes-a distribution over functions

- 82 3.1. Motivation
- If our goal is to learn or optimize an unknown function, then we need the following two ingredients to be successful.
- 1. A model for f, that is a model that learns what f looks like.
- 2. A method to select observations, given the problem statement.
- In this section we want to find a good model for f before we can then use it to either explore or exploit the function.

3.2. Weight space view

Let us first start by considering the text book-approach to learn functions, which is linear Bayesian regression, before then transitioning over to Gaussian Process regression. Remember that it is the task to either learn or optimize an unknown function. Therefore, regressing observed input points to observed output points by standard regression seems to be an apparent thing to do. Once we are able to capture the function—so the intuition we can easily quantify our uncertainty about it or try to find the point that produces the highest expected output. In classic linear regression, we assume normally distributed noise,  $\epsilon \sim \mathcal{N}(0, \Sigma)$ , and our goal is to predict a value y based on observations  $\mathbf{x}$  using weights  $\mathbf{w}$ .

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$$f(\mathbf{x}|\mathbf{w}) = \mathbf{x}^{\top}\mathbf{w}$$
$$y = f + \epsilon$$

Here,  $\mathbf{x}$  are our observations and  $\mathbf{w}$  is a vector containing the weights (sometimes notated as  $\beta$ ). If we assume a Gaussian prior over the parameters  $p(\mathbf{w}) = \mathcal{N}(0, \Sigma)$  and the likelihood  $p(\mathbf{y}|\mathbf{X}, \mathbf{w}) = \mathcal{N}(\mathbf{X}^{\top}\mathbf{w}, \sigma^{2}\mathbf{I})$ , then -by applying standard Bayesian inference—we get the posterior

$$p(\mathbf{w}|\mathbf{y}, \mathbf{X}) \propto p(\mathbf{y}|\mathbf{X}, \mathbf{w})p(\mathbf{w})$$
$$= \mathcal{N}(\frac{1}{\sigma^2}\mathbf{A}^{-1}\mathbf{X}\mathbf{y}, \mathbf{A}^{-1})$$

with  $\mathbf{A} = \mathbf{\Sigma}^{-1} + \sigma^{-2} \mathbf{X} \mathbf{X}^{\top}$ . In order to make predictions at a new test points  $\mathbf{x}_{\star}$ , we have to average over all posterior predictions

$$p(f_{\star}|\mathbf{x}_{\star}, \mathbf{X}, \mathbf{y}) = \int p(f_{\star}|\mathbf{x}_{\star}, \mathbf{w}) p(\mathbf{w}|\mathbf{X}, \mathbf{y}) d\mathbf{w}$$
$$= \mathcal{N}(\sigma^{-2}\mathbf{x}_{\star}^{\top} \mathbf{A}^{-1} \mathbf{X} \mathbf{y}, \mathbf{x}_{\star}^{\top} \mathbf{A}^{-1} \mathbf{x}_{\star})$$

Even though this approach is commonly chosen to model functions, the 107 way it is set up above only allows to make linear predictions. However, 108 only few relations in the real world are actually linear. Therefore, what 109 we need is a way to model non-linear dependencies as well. One possible 110 adjustment is to use a projection of the inputs  $\mathbf{x}$  onto a feature space by using 111 a function  $\phi(\mathbf{x})$ . One common projection is to use polynomials, resulting 112 into polynomial regression. Take a cubic regression as an example, which 113 assumes a function  $f(x) = \beta_0 + \beta_1 x + \beta_2 x^2 + \beta_3 x^3 + \epsilon$ . Deriving the posterior 114 for this model is similar to the linear regression described before, only that 115 all **X** are replaced by the projection  $\Phi = \phi(\mathbf{X})$ . However, the problem then becomes to find appropriate projections for our input variables as infinitely 117 many projections might be possible and we have to choose one a priori (or 118 by model comparison between finite sets of parametric forms). Especially if 119 the problem is to explore and exploit a completely unknown function, this 120 approach will not be beneficial as we would not know which projections we 121 should try out (we do not know the parametric form a priori). Fortunately, 122 there exists another approach to this problem which treats functions as 123 distributions and models this distribution directly. This approach is called 124 Gaussian Process regression and shifts the view away from a weight space 125 perspective to a function space view.

#### 3.3. Function space view

Another approach to regression problems is –instead of modeling distributions over weights– to focus on distributions over functions. A common
way to describe a distribution over functions is a Gaussian Process. A
Gaussian process is defined as a collection of random variables, any finite
number of which have a jointly Gaussian distribution. Therefore, a function
is distributed as

$$f(\mathbf{x}) \sim \mathcal{GP}(m(\mathbf{x}), k(\mathbf{x}, \mathbf{x}'))$$

Here,  $m(\mathbf{x})$  is a mean function modeling the expected output of the function and  $k(\mathbf{x}, \mathbf{x}')$  is a kernel function modeling the covariance between different points.

$$m(\mathbf{x}) = \mathbb{E}[f(\mathbf{x})]$$
$$k(x, x') = \mathbb{E}\left[ (f(\mathbf{x}) - m(x))(f(\mathbf{x}') - m(x')) \right]$$

As k(x, x') models the covariance between two different points, we have 137 to choose a function that will produce sensible estimates. The function k is 138 commonly called the kernel of the Gaussian Process (Jäkel, Schölkopf, and 139 Wichmann, 2007). The choice of an appropriate kernel is normally based on 140 assumptions such as smoothness and likely patterns to be expected in the data. If the data is not expected to be periodic, then a sensible assumption 142 is normally that the correlation between two points decays according to a 143 power function in dependency of the distance between the two points and that the covariance is symmetric, that is that only the distance between two 145 points matters, but not the direction. This just means that closer points

are expected to be more similar than points which are further away from each other in all possible directions. One very popular choice of a kernel fulfilling those requirement is the so-called squared exponential (sometimes also called Gaussian or Radial Basis Function) kernel.

$$k(x, x') = \sigma^2 \exp\left(-\frac{(x - x')^2}{2\lambda^2}\right)$$

The squared exponential is an expressive way to model smooth functions and the hyper parameters  $\lambda$  (called the length-scale) and  $\sigma^2$  (the noise constant) are normally optimized by using the marginal likelihood.

This implies the aforementioned distribution over functions as we can easily generate samples for new input points at location  $X_{\star}$ .

$$\mathbf{f}_{\star} \sim \mathcal{N}(0, K(X_{\star}, X_{\star}))$$

Given observations  $\mathcal{D} = \{\mathbf{X}, \mathbf{y}\}$  with a noise level  $\sigma$ , we can draw new predictions from our function  $\mathbf{f}_{\star}$  for inputs  $X_{\star}$  as described below.

$$\begin{bmatrix} \mathbf{y} \\ \mathbf{f}_{\star} \end{bmatrix} \sim \mathcal{N} \left( \mathbf{0}, \begin{bmatrix} K(X, X) + \sigma^2 I & K(X, X_{\star}) \\ K(X_{\star}, X) & K(X_{\star}, X_{\star}) \end{bmatrix} \right)$$

This means that we treat a function as a vector of infinite size. However, as we always only have to make predictions for finitely many points, we can simply draw outputs for these points by using a multivariate normal distribution with a covariance matrix generated by our kernel. Calculating the expectation of the Gaussian Process at the new points then is straight forward.

$$\mathbf{f}_{\star}|X,\mathbf{y},X_{\star} \sim \mathcal{N}(\overline{\mathbf{f}}_{\star},\mathrm{cov}(\mathbf{f}_{\star}))$$
 where

This means that predictions for new points are generated based on the expected mean value and covariance function of the posterior Gaussian Process.

$$\mathbb{E}[\mathbf{f}_{\star}|X,\mathbf{y},X_{\star}] = K(X_{\star},X)[K(X,X) + \sigma^{2}I]^{-1}\mathbf{y}$$
$$\operatorname{cov}(\mathbf{f}_{\star}) = K(X_{\star},X_{\star}) - K(X_{\star},X)[K(X,X) + \sigma^{2}I]^{-1}K(X,X_{\star})$$

Figure 1 shows an example of samples from a squared exponential Gaussian Process prior and the updated mean functions after some points have
been observed.

If we look at how to generate predictions for single points, we get the

following equation of the expectation for new points.

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 $\mathbf{f}_{\star}(\mathbf{x}) = \sum_{i=1}^{n} \alpha_{i} k(\mathbf{x}_{i}, \mathbf{x}_{\star})$ 

with  $\alpha = (K(X,X) + \sigma^2 I)^{-1} \mathbf{y}$ . What this equation tells us is that a 172 Gaussian Process can be written as a sum of basis functions. This means 173 that a potentially infinitely parametrized model boils down to a finite sum 174 when making predictions. This sum only depends on the chosen kernel and 175 the data observed thus far (Kac and Siegert, 1947). 176 This is also why Gaussian Process regression is referred to as non-177 parametric. It is not the case that this regression approach has no param-178 eters. Instead, it has potentially infinitely many parameters (parametrized 179 by the chosen kernel), but only manifests itself by a finite sum when making

## Samples from a Gaussian Process

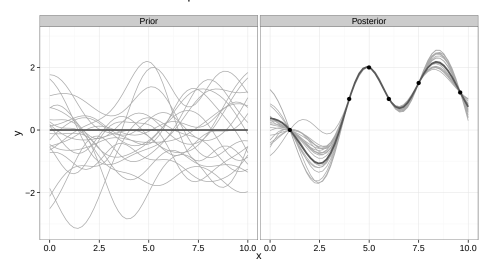


Figure 1: Example of samples from a Gaussian Process prior and posterior. Grey lines inidcate samples from the GP. Black dots mark empirical observations. The dark grey line marks the current mean of the GP.

predictions. Therefore, Gaussian Process regression is a powerful tool to capture many stationary functions. This in turn can be easily applied to contexts where the task is to explore or exploit these functions sequentially.

## 3.4. General setup

Having found a good model to learn functions, that is our first ingredient for successful sequential function learning, we now have to find a way to smartly explore or exploit the function we are learning over time. Within the Gaussian Process approach both pure *exploration* and *exploitation* can be viewed as two sides of the same coin. They both take a Gaussian Process to model an underlying unknown function <sup>1</sup> and then estimate the utility
of all available queries (candidate points from which to sample next) by
using what is called an acquisition function. An acquisition function can be
seen as a measurement of usefulness (or utility) that candidate points under
consideration promise to produce. The approach then goes on to choose as
the next point the one that currently produces the highest utility. The way
the general set up works is shown in Algorithm 1.

# Algorithm 1 General $\mathcal{GP}$ optimization Algorithm

Require: Input space  $\mathcal{D} = \{X, \mathbf{y}\}; \mathcal{GP}\text{-prior } \mu_0 = 0, \sigma_0, k$ 

for t = 1, 2, ... do

Choose  $x_t^* = \operatorname{argmax} f_{\text{Acquisition}}(\mathbf{x})$ 

Sample  $y_t = f(x_t^*) + \epsilon_t$ 

**Perform** Bayesian update for  $\mu_t$  and  $\sigma_t$ 

end for

This algorithm starts out with a Gaussian Process distribution over functions, then assesses the usefulness of the available samples by utilizing the acquisition function and selects the point that currently maximizes this function. Afterwards, the new output at the chosen sample point is observed, the Gaussian Process is updated, and the process starts anew.

<sup>&</sup>lt;sup>1</sup>Sometimes the Gaussian Process here is referred to as a surrogate model (Gramacy and Lee, 2008).

## 4. Exploration and Optimal Design

203 4.1. Acquisition function

The goal in an optimal design setting is to learn an unknown func-204 tion as well and quickly as possible. As said before, this is the same as 205 the attempt to maximize information gain over all trials. For a Gaussian, 206 the entropy  $H(\mathcal{N}(\mu, \Sigma)) = \frac{1}{2} \log |2\pi e\Sigma|$ , which means that in our setting 207  $I(\mathbf{y};f) = \frac{1}{2}\log|I + \sigma^{-2}K|$ , where K = [k(x,x')]. Even though finding the 208 overall information gain maximizer is NP-hard, it can be approximated by an 209 efficient greedy algorithm based on Gaussian Processes. If  $F(A) = I(\mathbf{y}_A; f)$ , 210 then this algorithm picks  $x_t = \operatorname{argmax} F(A_{t-1} \cup \{\mathbf{x}\})$ . This in turn can be 211 shown to be equivalent to the following acquisition function

$$f_{\text{Acquisition}}(\mathbf{x}_t) = \operatorname{argmax} \sigma_{t-1}(\mathbf{x})$$
 (2)

What this algorithm does, is to start out with a Gaussian Process prior, and to sequentially sample from the points that currently show the highest uncertainty in a greedy fashion. Even though this algorithm sounds naïve at first, it can actually be shown to converge to at least a constant fraction of the maximum information gainer (Krause, Singh, and Guestrin, 2008).

$$F(A_T) \ge \left(1 - \frac{1}{e}\right) \max F(A)$$
 (3)

This is based on a property of the acquisition function called *submodularity* (Krause and Golovin, 2012). Intuitively, submodularity here means that information never hurts (it is always helpful to observe more points), but also that the usefulness of newly acquired points decreases the more points have been sampled before. This diminishing returns property is crucial to

show that the greedy algorithm can be successful over all (see Appendix).

A simple example of the Gaussian Process uncertainty reduction sampler is
shown in Figure 2 below. We have sampled a function from a Gaussian Process prior (a squared exponential) and let the algorithm select observations
by picking as the next obvservation the one that currently has the highest
standard deviation attached.

Figure 2: GP-uncertainty reduction example. The dark grey line marks the current mean of the GP. The dashed line shows the upper part of the standard deviation. The light grey lines are samples from the GP.

Optimal design

# Step 1 Step 2 (X)J tndtnO O

10.0 0.0 Input 2.5

5.0

7.5

10.0

# 224 4.2. Example: Learning unknown functions

5.0

7.5

0.0

225

226

227

228

2.5

In order to demonstrate how Gaussian Process based exploration works, let us simulate how the algorithm learns unknown functions and compare it to other algorithms. If we assume that we have to learn an unknown function as quickly as possible and that the function f only takes a one-dimensional input x = [0, 0.1, 0.2, ..., 10] and to which it maps an output y, then we can

set up different functions that the models can learn actively over time. As
the GP is considered to learn many different functions well, we will test the
following different functions: a linear, quadratic, cubic, logarithmic, sine,
and a non-stationary function (see Appendix for details).

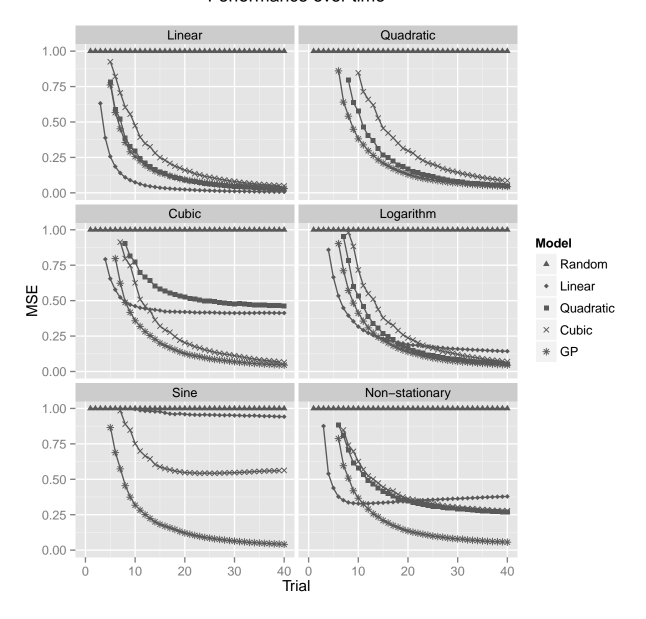
We have deliberately chosen different parametric forms to assess the 234 learning methods. The models we have used to learn the different function 235 are a linear, a quadratic, a cubic, and a Gaussian Process (with a squared 236 exponential kernel) regression. Each model was set up to learn the under-237 lying function by picking as the next observation the one that currently has 238 the highest uncertainty (standard deviation of the predicted mean). We let 239 each model run 100 times for each underlying function and averaged the 240 mean squared error over the whole discretized input space for each step. 241 Results are shown in Figure 3. 242

It can be seen that the Gaussian Process model learns all functions both efficiently and well. Only in the cases in which the used learning function is indeed the same as the learning function (for example, using a linear function to learn an underlying linear function), does another model learn faster than the Gaussian Process.

This means that Gaussian Processes are especially useful in cases where
the underlying function is not known or can be ignored. For examle, one
could easily use Gaussian Processes to learn participants' utility function
over different experiments or simply use them to generate stimuli that are
most informative overall.

Figure 3: GP-uncertainty reduction example.

# Performance over time



## 5. Exploration-Exploitation and Bayesian Optimization

In an exploration-exploitation scenario the goal is to find the argument to a function that produces the maximum output as quickly as possible. One way to measure the quality of this search process is to quantify regret. Regret is the distance between the output of the currently chosen argument and the maximum output.

$$r(\mathbf{x}) = f(\mathbf{x}^*) - f(\mathbf{x}) \tag{4}$$

The goal then is to minimize regret over all available trials.

$$\min \sum_{t}^{T} r(x_t) = \max \sum_{t}^{T} f(x_t)$$
 (5)

Again, finding the global function maximizer is NP hard. That is finding the minimum sequence of queries that leads to the lowest regret overall is almost impossible. However, there is again a greedy trick one can apply in this scenario. This trick is based on redefining the function maximization problem as a bandit task. In a bandit tasks the goal is to maximize output by playing the right arm out of many available arms (it is named after the one armed-bandits that can be found in casinos). As the tasks is to maximize output of a function, the discretized input points to that functions are seen as arms, that are correlated in dependency of the underlying covariance kernel. Taking this perspective, one easy way to approach these kind of problems is the following acquisition function called Upper Confidence Band sampling.

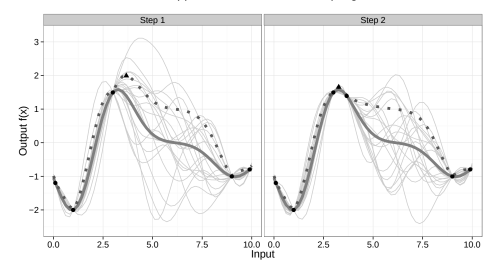
$$f_{\text{Acquisition}}(\mathbf{x}) = \mu_{t-1}(\mathbf{x}) + k\sigma_{t-1}(\mathbf{x})$$
 (6)

Upper Confidence Band sampling (UCB) plays the arm that currently shows the highest upper confidence interval. This strategy is sometimes called

naive optimism in the face of uncertainty. Intuitively, the upper confidence 256 band is determined by two factors, the current estimate of the mean at a 257 particular point (the higher the estimate, the higher the band) and the un-258 certainty attached to that estimate (the higher the uncertainty, the higher 259 the band). Therefore, the UCB algorithm trades off naturally between ex-260 pectations and uncertainties. An example of how the UCB works is shown 261 in Figure 4. Again, we have sampled a function from a Gaussian Process 262 prior and have applied the algorithm to this function. 263

Figure 4: GP-UCB example. The dark grey line marks the current mean of the GP. The dashed line marks the GP's upper confidence bound. The light grey lines are samples from the GP.

## Upper Confidence Band sampling



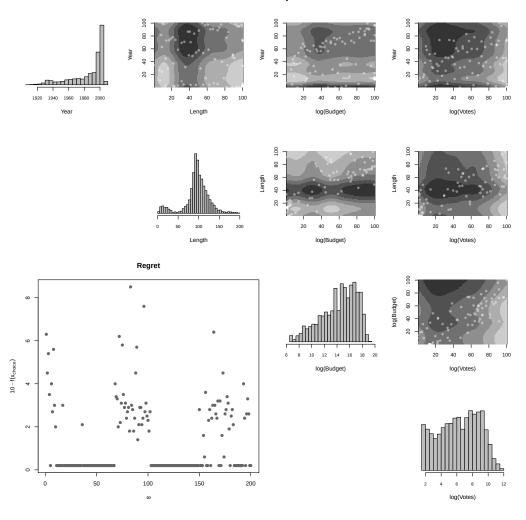
Even though the greedy UCB strategy is again naïve, it can be shown that its regret is sublinear, again based on the submodularity of the overall information gain (Srinivas, Krause, Kakade, and Seeger, 2009).

# 267 5.1. GP-UCB Example: Choosing movies

We use the imdb-movie data as an example for doing GP-UCB exploratio-268 exploitation. For this, we extracted 5141 movies from the data base, includ-269 ing the year they appeared, the budget that was used to make them, their 270 length, as well as how many people had evaluated the movie on the imdb. 271 The dependent variable was the actual imdb score and we set up a GP-UCB 272 with a squared exponential kernel, set  $\beta = 3$  and let the algorithm pick 200 273 movies sequentially. Results are shown in Figure 5. 274 It can be seen that the algorithm quickly starts picking movies that 275 produce low to zero regret. However, it still keeps exploring other movies 276 from time to time, just to alous come back to recommending movies with 277 really high scores. Additionally, it explores all of the given input areas for 278 quite some time, just to find good areas in the end.

Figure 5: GP-UCB example.

# **Gaussian Process Optimization**



## 6. Safe Exploration-Exploitation

Sometimes an experimentalist's goal might not only be to learn or max-281 imize a function, but additional requirements can also be important. One 282 such requirement, for example, can be to avoid certain outputs as much as 283 possible. One example for such a scenario is excitatory stimulation, espe-284 cially in a physiologically setting. Imagine a medical scenario, where the 285 task is to stimulate the spinal chord in such a way that certain movements 286 are achieved (Desautels, 2014). Here, it is important to stimulate the spinal 287 chord such that optimal recovery is achieved, but not too much as this might 288 lead to painful reactions within the patients. Again, Gaussian Process opti-289 mization methods can be used here to learn the underlying function of what 290 stimulations lead to what strength of reaction. However, an additional re-291 quirement now is to avoid these expectedly painful areas. It turns out that 292 there is a smart way to adapt the GP-UCB approach described above while 293 accommodating for this additional requirement. This is acchieved by an al-294 gorithm called SafeOpt (Sui, Gotovos, Burdick, and Krause, 2015). Leaving 295 the detailed technical explanation of this algorithm for the interested the 296 reader to look up, this algorithm basically works by trading-off two different 297 things. Firstly, it keeps a set of safe options it considers to be above the 298 given threshold and tries to expand this set as much as it can. Secondly, 299 it maintains a set of potential maximizers that, if used as an input, would 300 potentially achieve the highest output. It then choses as the next point, a 301 point within the intersection of these two sets that has the highest predictive 302 variance.

# 304 6.1. Example: Stimulus Optimisation

Imagine that you have to optimize a stimulus that can be described by 305 two dimensions  $X_1$  and  $X_2$ . This means you have to find the maximum 306 output y. However, you also want to avoid sampling values below 0 at all 307 costs. For this we have drawn a two dimensional function from a Gaussian 308 Process prior with a squared exponential kernel. This can be seen as a 309 similar to the case where one want to present stimuli to participants, but 310 make sure that participants never react with an intensity below a certain 311 threshold. Results are shown in Figure 6. 312 It can be seen that the SafeOpt algorithm finds the maximum of the

It can be seen that the SafeOpt algorithm finds the maximum of the function well, but also tries to expand the space of possible inputs more and more. At the same time, the algorithm does not sample from the white area (values below 0) for a single time. We therefore think that this algorithm could potentially applied in many optimal design settings that require additional constraints.

Figure 6: GP-SafeOpt example. White areas represent areas below 0. The black crosses show where the SafeOpt algorithm has sampled.

# **GP Safe Optimization** $\omega\,{=}\,10$ $\omega = 1$ $\overset{\mathsf{z}}{\mathsf{x}}$ **X**<sub>1</sub> **X**<sub>1</sub> $\omega = 50$ $\omega = 100$

## 9 7. Gaussian Processes and cognition

As we have seen, Gaussian Processes (in combination with smart ac-320 quisition functions) are a powerful tool to learn and maximize unknown 321 functions. However, they might also be applied in a different, even more 322 psychological context, that is as as a model for human cognition within 323 the tradition of Bayesian cognitive science (Chater and Oaksford, 2008). 324 Non-parametric Bayesian approaches have been used before to describe associative learning (Gershman and Niv, 2012) and categorization learning 326 (Kemp, Tenenbaum, Griffiths, Yamada, and Ueda, 2006), among others. 327 Recently, Lucas, Griffiths, Williams, and Kalish (2015) have proposed to 328 use Gaussian Processes as a rational model of human function learning for 329 passive learning tasks. Schulz, Tenenbaum, Reshef, Speekenbrink, and Ger-330 shman (2015c) used Gaussian Processes to assess participants judgments 331 about the predictability of functions in dependency of the smoothness of 332 the underlying kernel. Taking recent developments within the active learn-333 ing community into account, we expect Gaussian Processes to soon become 334 a fruitful descriptive model for human cognition in many different domains 335 where participants have to act actively, that is to select information sequen-336 tially. In a first attempt, we have found that participants' behavior can be 337 well described by Gaussian Process algorithms when the task is to learn and 338 maximize a function both in a static (Schulz, Konstantinidis, and Speeken-339 brink, 2015b) as well as in a dynamic environment (Schulz, Konstantinidis, 340 and & Speekenbrink, 2015a). However, given GP's expressiveness and the 341 mathematical guarantees that come with them, we expect them to be used more frequently as a model for human cognition in the near future and hope 343 that this tutorial can help people to apply them more often.

Here, it is not only the assumption of not having to choose a parametric 346 shape a priori, but letting the data speak directly, that makes these models powerful as a psychological model. It is also the attached measure of un-348 certainty that comes for free when doing computations with Gaussian Pro-349 cesses, a characteristic more and more used in numerics (Hennig, Osborne, 350 and Girolami, 2015) and optimization problems (Hennig and Kiefel, 2013). 351 The resulting uncertainty of computation can then also be propagated be-352 tween different systems easily (Damianou and Lawrence, 2012). Using this 353 fact, one could build of a model that involves control, learning, and opti-354 mization and in which uncertainty is modeled over the whole system, sort 355

#### 8. Discussion

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of a Bayesian Cognitivism approach.

345

This tutorial has introduced Gaussian Process regression as a general 358 purpose inference engine to learn about (explore) and maximize (exploit) 359 unknown functions. Gaussian Processes have been introduced mathemati-360 cally and the greedy variance minimization exploration algorithm as well as 361 the upper confidence bound sampling method for exploration-exploitation 362 scenarios have been introduced. Within a simulated exploration experiment 363 we have shown how Gaussian Processes can be used to efficiently learn about unknown functions in an optimal design set-up and gradually presented in-365 formative stimuli. Within a parameter tuning example, we have shown how 366 GP-UCB can outperform other commonly used methods when the goal is to optimize hyper-parameters of a neural network. Additionally, we have 368 talked about introducing additional requirements to the used acquisition

function and have shown one example within a scenario, where the task was
to show the optimal excitatory stimulus while avoiding some areas of the input space. Finally, we have talked about utilizing Gaussian Process models
as actual models of cognition in the hope that they will be picked up and
used more frequently in the near future.

375

Of course a tutorial like this can never be fully comprehensive. Therefore, 376 let us briefly point out some things that we have not covered here. First of 377 all, using the Gaussian Process regression approach as we have parametrized 378 here is only one possible approach towards regression problems. In fact, it 379 can be shown that many standard Bayesian regression approaches can be 380 re-parametrized to be equivalent to Gaussian Process regression, given spe-381 cific assumptions about the kernel (Duvenaud, Lloyd, Grosse, Tenenbaum, 382 and Ghahramani, 2013). The two chosen utility functions here also are just 383 two options out of a pool of different acquisition functions. Another com-384 monly used acquisition function for the pure exploration case is the attempt to minimize the expected variance summed up over the whole input space 386 (Gramacy and Apley, 2014). Even though this method tends to sample less 387 from the boundary cases as compared to the algorithm introduced here, it 388 can be hard to compute, especially if the input space is large. There exist 389 many different acquisition functions in the exploration-exploitation context, 390 that are mostly discussed under the umbrella term Bayesian optimization 391 (de Freitas, Smola, and Zoghi, 2012). One other common acquisition func-392 tion that is frequently used here is the expected probability of improvement 393 (Mockus, 2012), which choses as the next point the one that promises to 394 have the highest likelihood of improving the currently expected maximum. Again, this method of assessing candidate points can be computationally expensive and mathematical guarantees are only given for the UCB algorithm.

Last not least, there are also many different acquisition functions with ad-

ditional constraints available. Important to mention hereby are Bayesian

400 black box optimization with additional constraints, the estimation of level

sets, as well as Bayesian Quadrature-based algorithms.

402

Here we have introduced Gaussian Process-based algorithms to explore and exploit unknown functions. We hope that this tutorial will help others to pick up these methods and that their usage will gradually become more common within psychology.

## 9. Software packages

Table 1 below contains further Gaussian Process software pointers.

Table 1: Gaussian Process Software Packages

Name	Algorithm	Language	Author
GPML	GP Toolbox	Matlab	Rasmussen & Williams
SFO	Submodular Optimization	Matlab	Krause
GPy	GP Toolbox	Python	Sheffield ML Group
gptk	GP Toolbox	R	Lawrence
$\operatorname{tgp}$	Tree GPs, GP regression	R	Gramacy & Taddy

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