# **Bayesian Optimization**

Why & How?

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Sargent Centre Summer School on Bayesian Optimization, 2024

For copies of slides & code, see <a href="https://github.com/joelpaulson/Sargent\_Centre\_BO\_Summer\_School\_2024">https://github.com/joelpaulson/Sargent\_Centre\_BO\_Summer\_School\_2024</a>

#### **Outline**

- Introduction to Bayesian optimization
  - White-box vs. black-box, prevalence of expensive functions, bird's eye view
- Improvement-based acquisition functions
  - Expected improvement, knowledge gradient
- Information-theoretic acquisition functions
  - Predictive, max-value, and joint entropy search
- Constrained Bayesian optimization
  - Expected improvement with constraints, exact penalty methods
- Practical considerations
  - Optimizing the acquisition function, kernel adaptation, acquisition scheduling

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# What is an Optimization Problem?

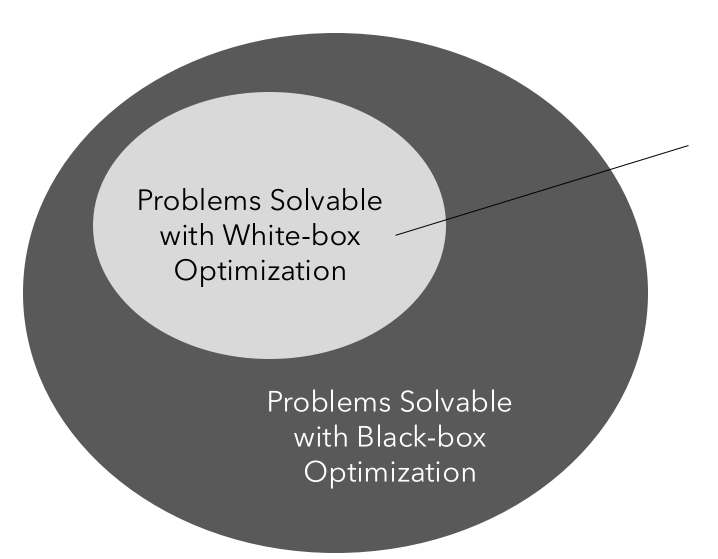
decision (design) 
$$\underbrace{ \ \ }_{x} \ \ f(x) \Big\} - \underbrace{ \ \ }_{constraints}$$
 objective function 
$$s.t. \ \ x \in \Omega \Big\} - \underbrace{ \ \ }_{(feasible region)}$$

- Optimization problems are **pervasive** in every application domain
  - differentiate problems based on characteristics  $\rightarrow$  determine what solver to use
- There are a huge number of available optimization algorithms; difficult to a priori know the best one but we can eliminate some options

# **How to Classify Optimization Algorithms?**

- A simple way to "partition" the algorithms into two major buckets are "white-box" and "black-box" (i.e., not white box)
- White-box means that we need an "equation-oriented model" of the system so that the mathematical structure of f(x) and  $\Omega$  satisfy certain important assumptions
  - The exact assumptions depend on the method, but they will typically require the functions to be differentiable and/or easy to build relaxations of them
- Any method that only requires evaluations of f(x) and  $x \in \Omega$  at specific points can then be classified as "black box"

# **How to Classify Optimization Algorithms?**



Since white-box algorithms make stronger assumptions, they can only be used to tackle a subset of problems when compared to black-box algorithms

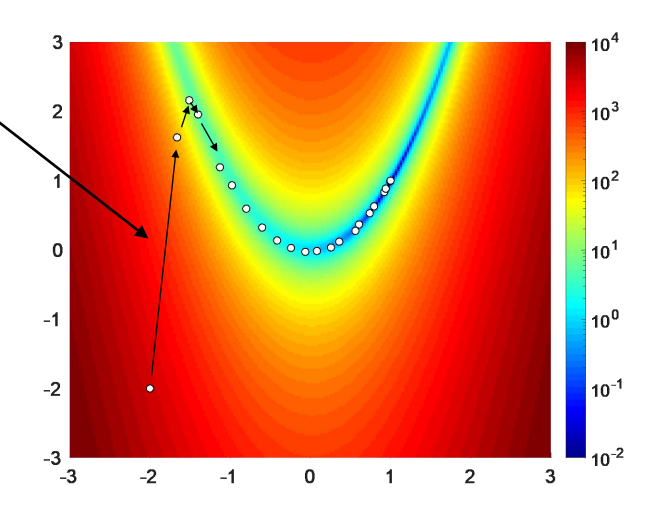
→ The main value of black-box methods are their generality (not necessarily efficient)

# **Example of White-Box Optimization: Newton's Method**

Use derivatives to take step toward reducing objective, i.e.,

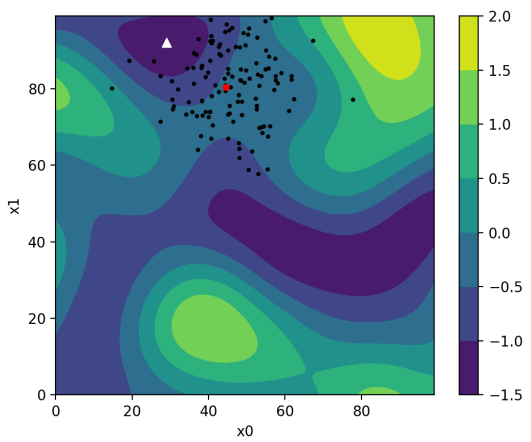
$$x_{k+1} = x_k - \alpha_k (\nabla^2 f(x_k))^{-1} \nabla f(x_k)$$

This type of algorithm is "local" (requires initial guess) & requires ability to compute derivatives (expensive when the structure of the function is unknown)



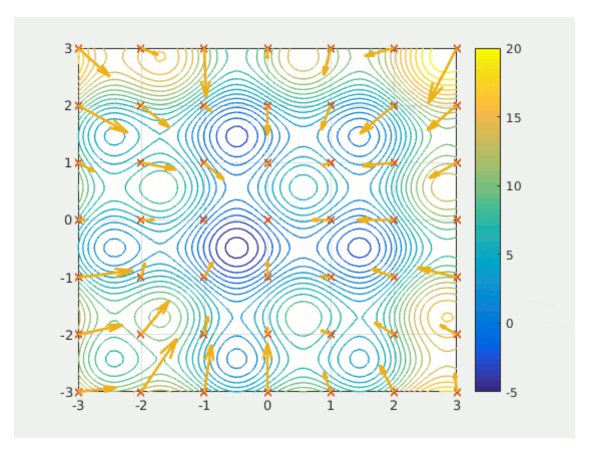
# **Examples of Black-Box (Derivative-Free) Optimization**

Covariance Matrix Adaptive Evolutionary Strategy (CMA-ES)



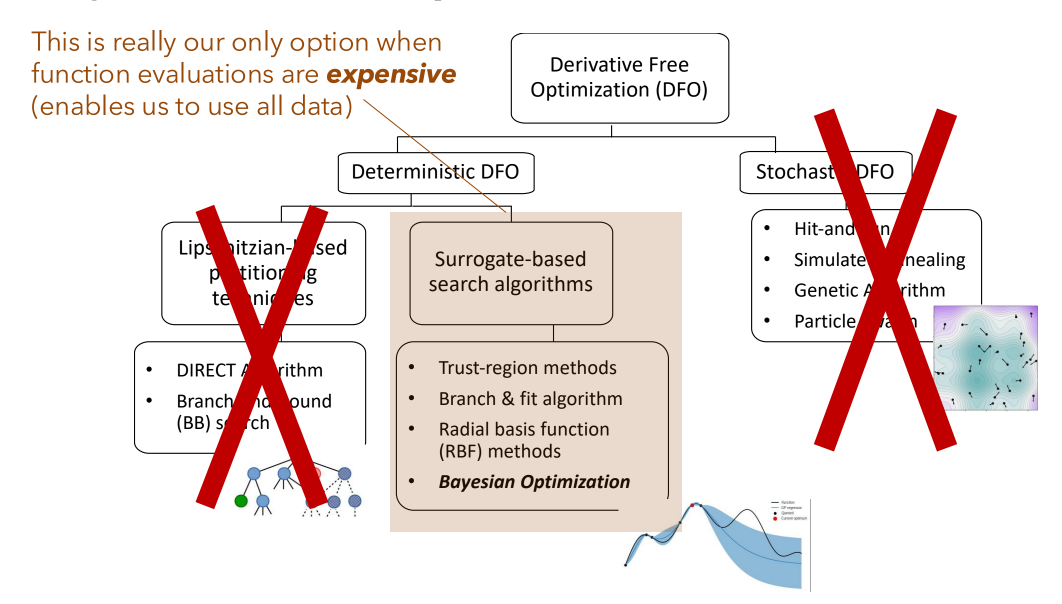
https://thurinj.github.io/CMA-ES.html

Particle Swarm Optimization (PSO)

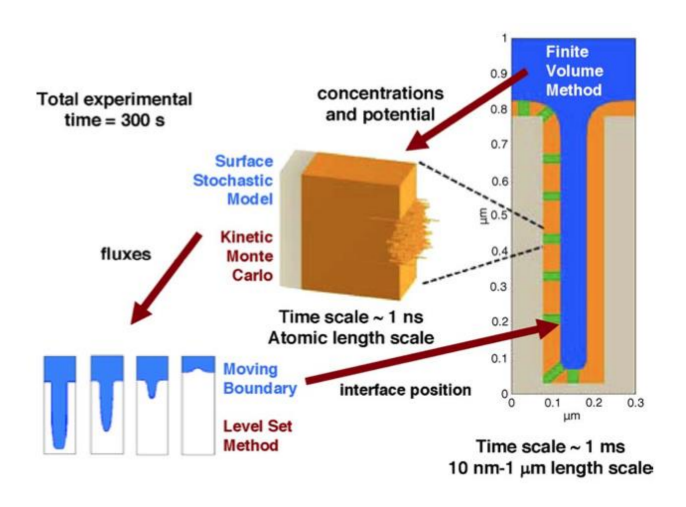


https://en.wikipedia.org/wiki/Particle\_swarm\_optimization

## Many derivative-free optimization methods, which to choose?



#### Optimizing multi-scale simulation models



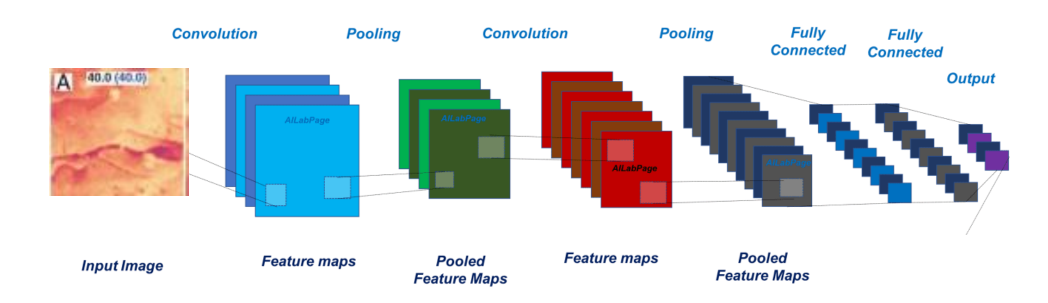
#### Objective:

Minimize surface roughness

#### Design variables:

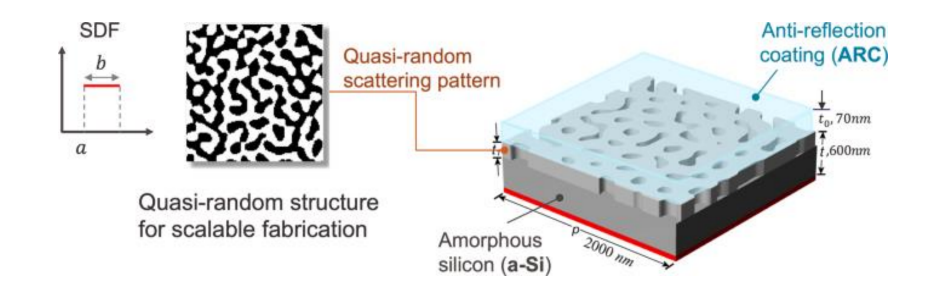
Chemical additive concentrations & reaction temperature

#### Automated machine learning



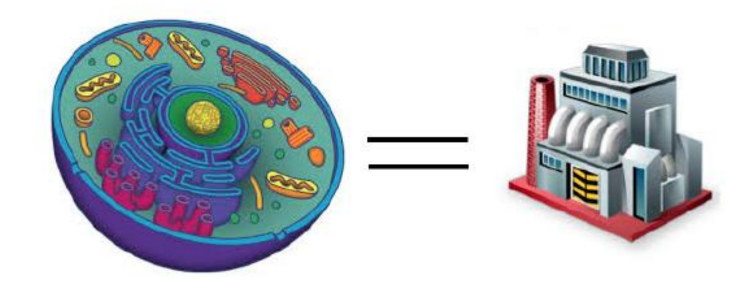
- Objective: Maximize classification accuracy for image-based chemical sensor
- Design variables: Number of layers, number of nodes per layer, learning rates, regularization penalties, activation functions, etc.

Material and drug discovery



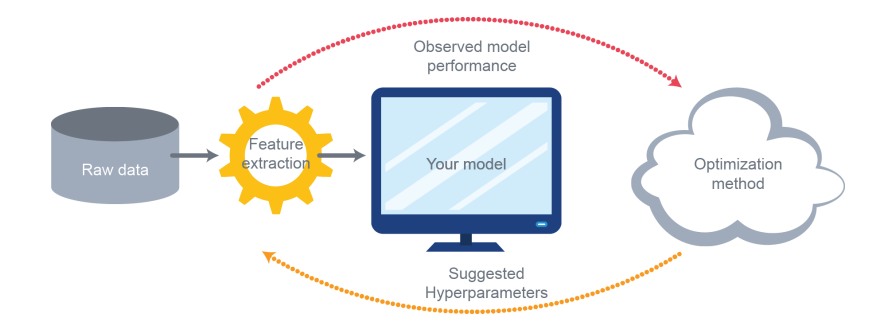
- Objective: Maximize light adsorption in quasi-random solar cell
- Design variables: Type of amorphous silicon (a-Si), light trapping pattern for fabrication,
   & overall thickness

Design of experiments: Gene optimization



- **Objective:** Maximize efficiency of the cell factory to make product (e.g., proteins)
- **Design variables:** Gene sequence (e.g., ATTGGTUGA...) & culture conditions (e.g., pH)

Tuning hyperparameters in optimization codes



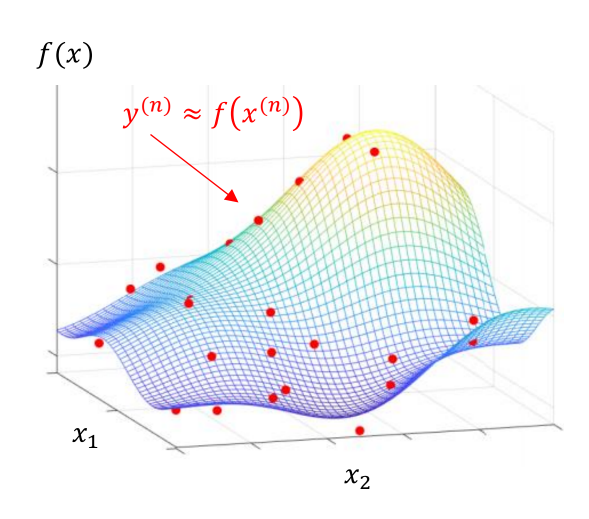
- Objective: Minimize solution time for family of scheduling/planning problems
- Design variables: Algorithmic parameters in solver (e.g., CPLEX has 76 design parameters)

#### Many other problems:

- Robotics, aerospace, control, reinforcement learning
- Tuning websites with A/B testing
- Calibrating expensive simulators to experimental data
- etc....

#### **Standard Goal in Bayesian Optimization:**

Optimize functions  $f: \mathbb{R}^d \to \mathbb{R}$  that are:

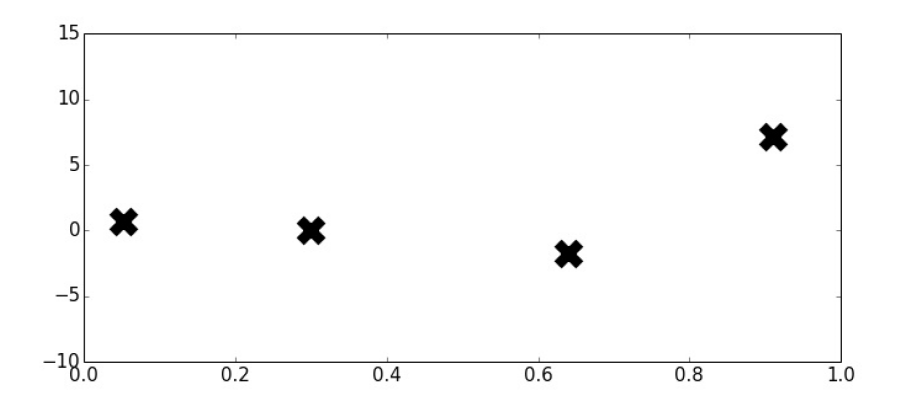


\*We will deal with black-box constraints later

- $f(\cdot)$  is explicitly <u>unknown</u> & <u>non-convex</u>
  - lacks known special structure, e.g., convexity
- $f(\cdot)$  is <u>derivative-free</u>
  - cannot simply get gradients
- $f(\cdot)$  is expensive to evaluate
  - # of evaluations is severely limited
- $f(\cdot)$ 's evaluations may be <u>noisy</u>
  - noise independent & ~normally distributed,
     but unknown variance

## Illustrative example to build some intuition

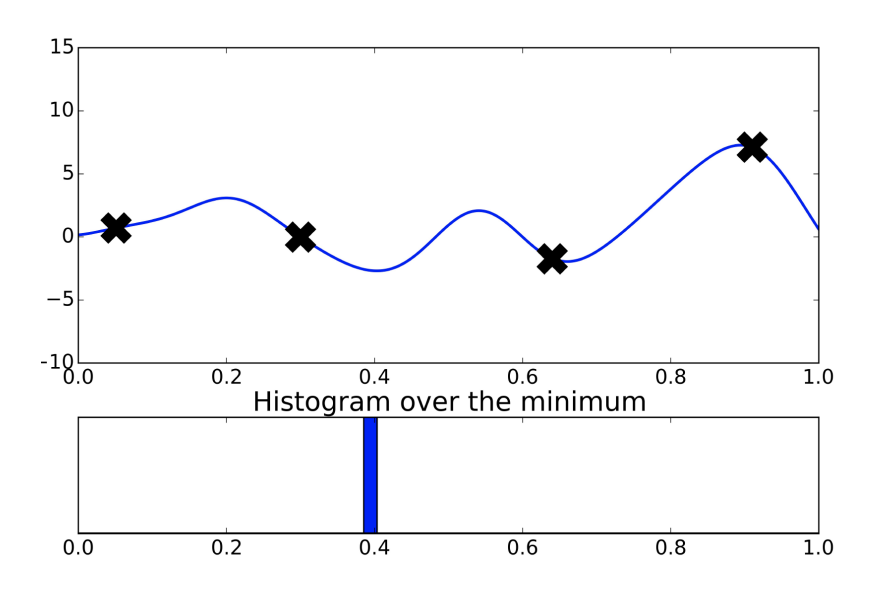
We have four function evaluations



- Where is the minimum of the function  $f(\cdot)$ ?
- Where should we take our next evaluation?

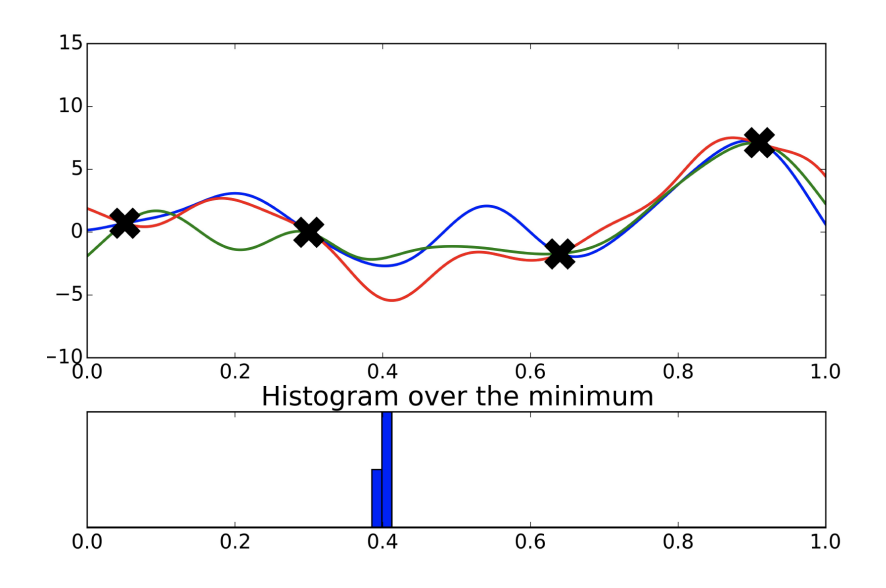
#### Intuitive solution, fit a surrogate model

One curve; which one should we select?



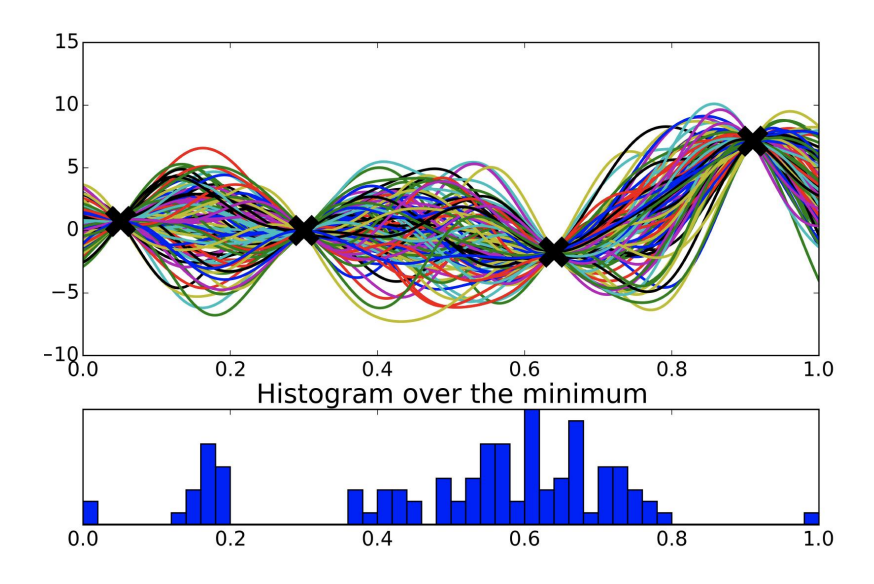
#### Intuitive solution, fit a surrogate model

Three curves



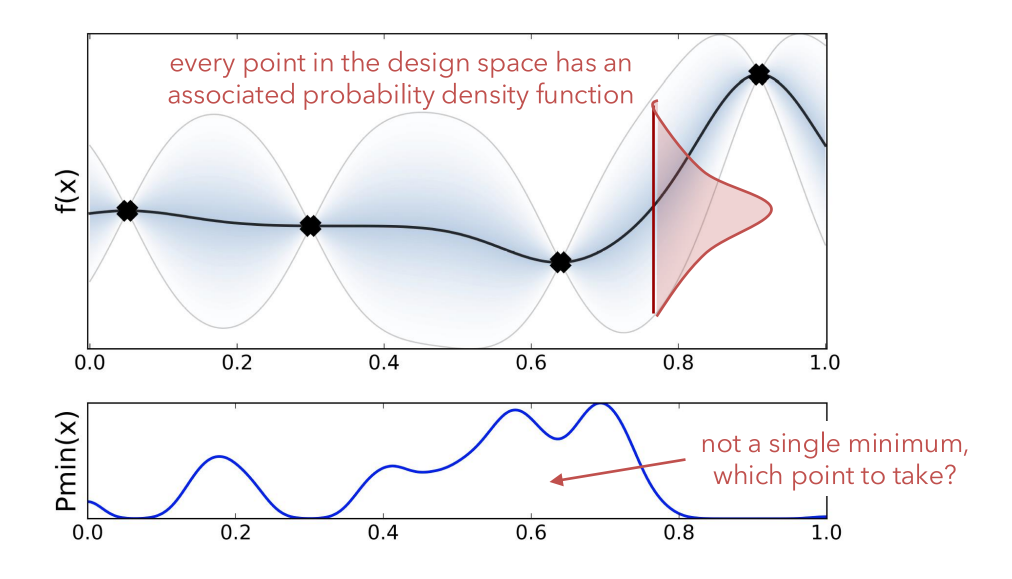
#### Intuitive solution, fit a surrogate model

One hundred curves



# Intuitive solution, fit a surrogate model Infinite curves

(Need the help of information theory to properly define models + metrics)



# Bird's-eye View of Bayesian Optimization

Last section of this presentation

while {budget not exhausted}

See "prior" GP slides

Fit a Bayesian machine learning model (usually Gaussian process regression) to observations {x, f(x)}

Find x that maximizes acquisition(x, posterior)

Sample x & then observe f(x)

Next few sections of *this* presentation

end

More Information

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# Bird's-eye View of Bayesian Optimization

while {budget not exhausted}

Fit a Bayesian machine learning model (usually Gaussian process regression) to observations {x, f(x)}

Find x that maximizes acquisition(x, posterior)

Sample x & then observe f(x)

end

Assume our goal is to minimize f(x) [same idea as maximize, just replace with -f(x)]

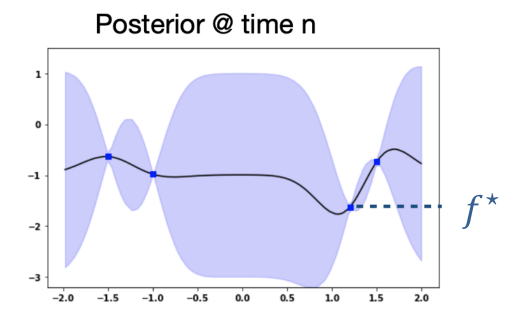
# How to Define an Acquisition Function $\alpha_n$ ?

- When properly selected, the value of  $\alpha_n(x)$  at any  $x \in \Omega$  should be a good measure of the (expected) benefit of querying f at that point in future
  - Must depend on the posterior distribution of  $f|y_{1:n}$
- This implies we should like to preferentially sample at the point that produces the highest possible value of the acquisition function:

$$x_{n+1} \in \operatorname{argmax}_{x \in \Omega} \alpha_n(x)$$

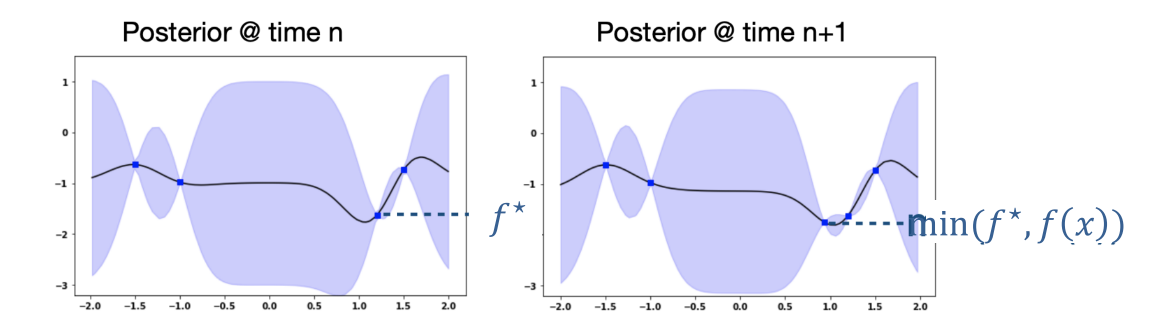
• It is expected for this problem to be much cheaper to solve since, unlike f, we have some equation-based form for  $\alpha_n$ 

[Mockus 1989; Jones, Schonlau, and Welch 1998]



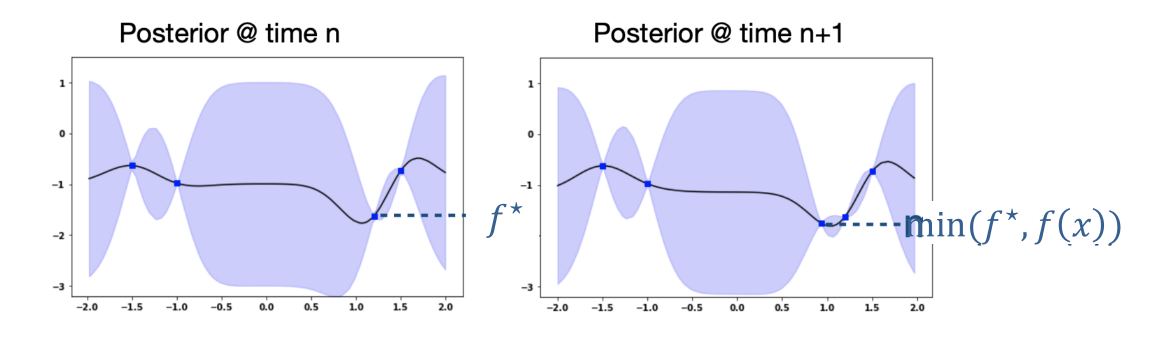
• Loss if we stop now:  $f^*$ 

[Mockus 1989; Jones, Schonlau, and Welch 1998]



- Loss if we stop now:  $f^*$
- Loss if we stop after sampling at f(x): min $(f^*, f(x))$

[Mockus 1989; Jones, Schonlau, and Welch 1998]



- Loss if we stop now: f\*
- Loss if we stop after sampling at f(x): min $(f^*, f(x))$
- Expected reduction in loss due to sampling:  $\mathbb{E}_n[f^* \min(f^*, f(x))]$

[Mockus 1989; Jones, Schonlau, and Welch 1998]

$$\operatorname{EI}_{n}(x) = \mathbb{E}_{n} \{ f^{*} - \min(f^{*}, f(x)) \}$$

$$= \mathbb{E}_{n} \{ \max\{ f^{*} - f(x), 0 \} \}$$

$$= \mathbb{E}_{Z} \{ \max\{ f^{*} - \mu_{n}(x) - \sigma_{n}(x)Z, 0 \} \}$$

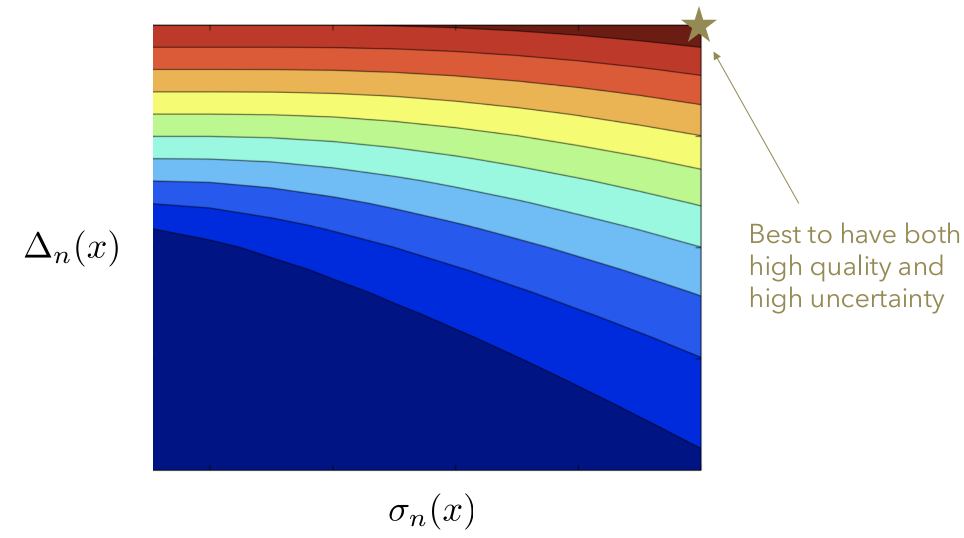
Integral can be carried out analytically using integration by parts

# **Closed-form Expression Expected Improvement**

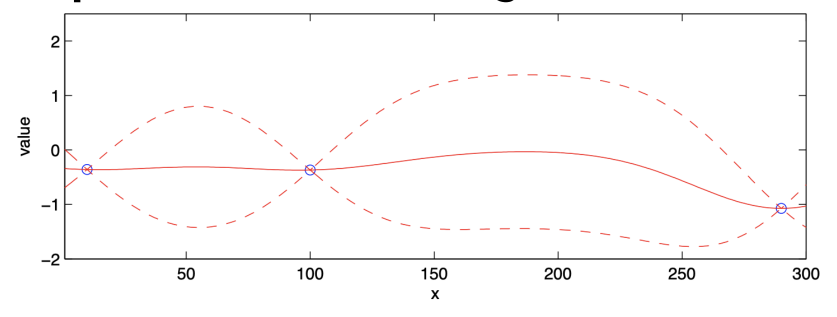
Standard normal cumulative distribution function (CDF) Standard normal probability density function (PDF)  $EI_n(x) = \Delta_n(x) \Phi\left(\frac{\Delta_n(x)}{\sigma_n(x)}\right) + \sigma_n(x) \phi\left(\frac{\Delta_n(x)}{\sigma_n(x)}\right)$ 

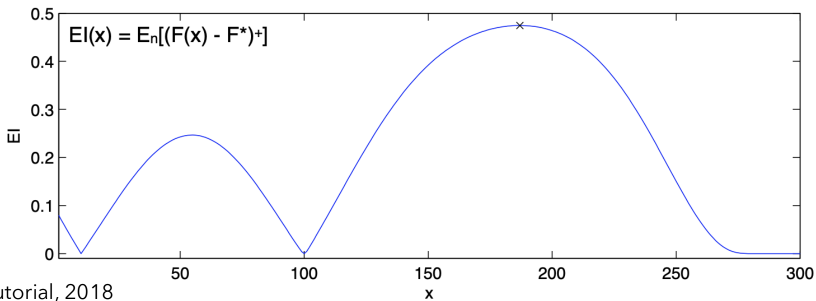
Where  $\Delta_n(x) = f_n^{\star} - \mu_n(x)$  is expected quality

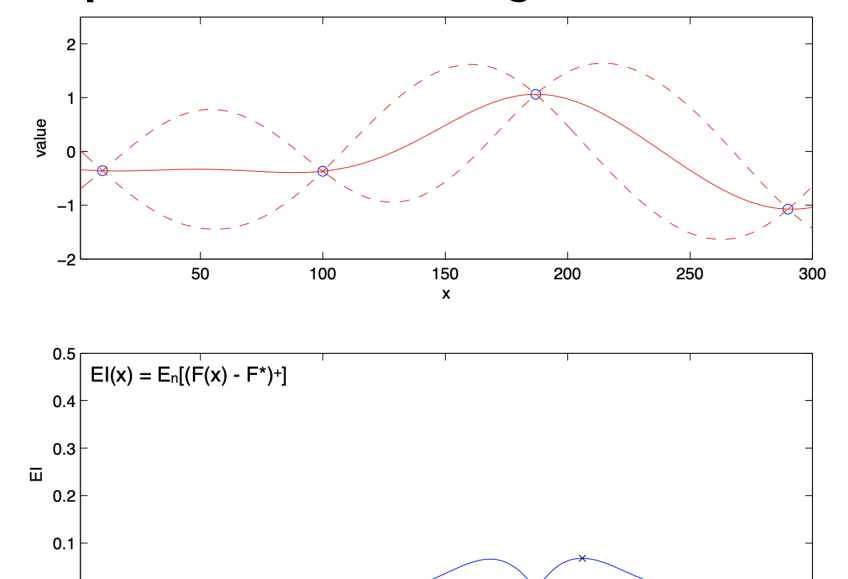
# El Tradeoffs Exploitation ( $\Delta_n(x)$ ) vs. Exploration ( $\sigma_n(x)$ )



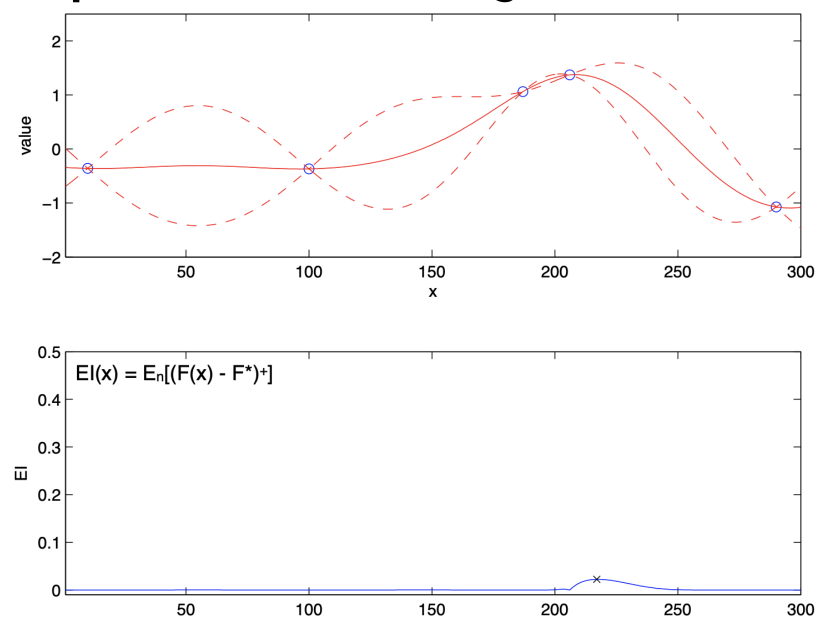
[Jones et al., 1998; Frazier, 2018]



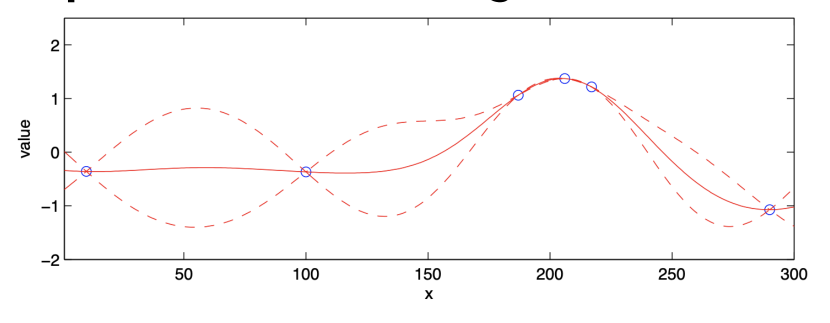


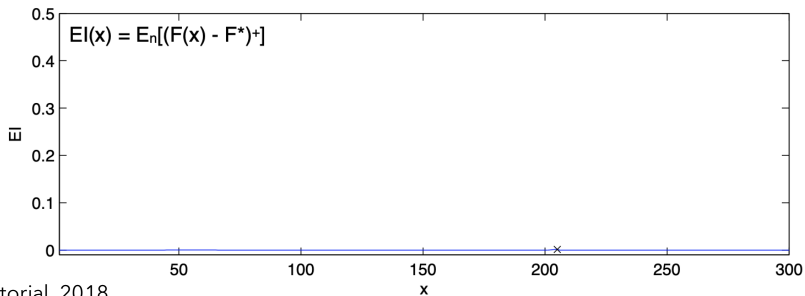


Χ



Χ





# **Thought Experiment**

- What should expected improvement (EI) reduce to when the variance of the prediction is zero everywhere?
  - We no longer have uncertainty, so we no longer need to sequentially search (simply find the minimum of the mean function)
  - We can think of traditional optimization as placing a GP prior with perfectly known mean function  $\mu_0(x) = f(x)$  (and zero variance/covariance), then running EI one step to find the "true" minimum
  - El in some sense generalizes traditional "white-box" optimization to the unknown "black-box" setting → attempts to be information-optimal

## Is El Optimal in any Sense?

- Yes, it turns out that EI is Bayes-optimal under some assumptions:
  - -There is no noise in the observations of the objective function
  - -We are only willing to select previously evaluated point as final solution
  - -We are risk neutral (i.e., we value a random outcome according to its expected value, hence  $\mathbb{E}[Reduction\ in\ Loss])$
  - -This is our last evaluation

Why is this assumption needed?

# In general, we must solve a sequential decision-making problem

- The loss that we calculated previously is only a function of the next sample that we take; however, in general, we have a budget of N remaining samples  $\{x_1, x_2, ..., x_N\}$
- Furthermore, every sample that we take yields more data, such that we have more information to make our next decision
- We can formulate this as a stochastic optimal control problem where our <u>state is current data</u>, <u>action is next sample</u>, and <u>immediate reward is reduction in loss</u>

# Best (finite-budget) sampling strategy is policy that optimizes the value function (total loss reduction)

- Policy:  $\pi = \{\pi_1, \pi_2, \dots, \pi_N\}$ ,  $x_k = \pi_k(\mathcal{D}_{k-1})$
- Value function:  $V_{\pi}(\mathcal{D}_0) = \mathbb{E}\left[\sum_{k=1}^N r(\mathcal{D}_{k-1}, \mathcal{D}_k)\right]$ ,  $r(\cdot)$  is loss reduction
- Optimal policy:  $V^{\star}(\mathcal{D}_0) = V_{\pi^{\star}}(\mathcal{D}_0) = \max_{\pi \in \Pi} V_{\pi}(\mathcal{D}_0)$
- Solution expressed using dynamic programing:

$$V_k(\mathcal{D}) = \max_{x \in \Omega} \mathbb{E}_{\mathcal{D}^+} \left[ r(\mathcal{D}, \mathcal{D}^+) + V_{k-1}(\mathcal{D}^+) \right], \quad \forall k = 1, \dots, N$$

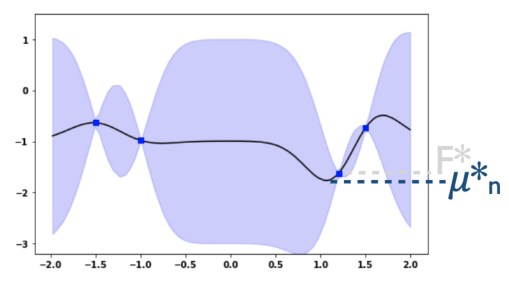
## Let's Drop Two of These Assumptions

- Yes, it turns out that EI is Bayes-optimal under some assumptions:
  - -There is no noise in the observations of the objective function
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  - -We are risk neutral (i.e., we value a random outcome according to its expected value, hence  $\mathbb{E}[Reduction\ in\ Loss])$
  - -This is our last evaluation

Yields Knowledge Gradient (KG) acquisition function, what should be loss?

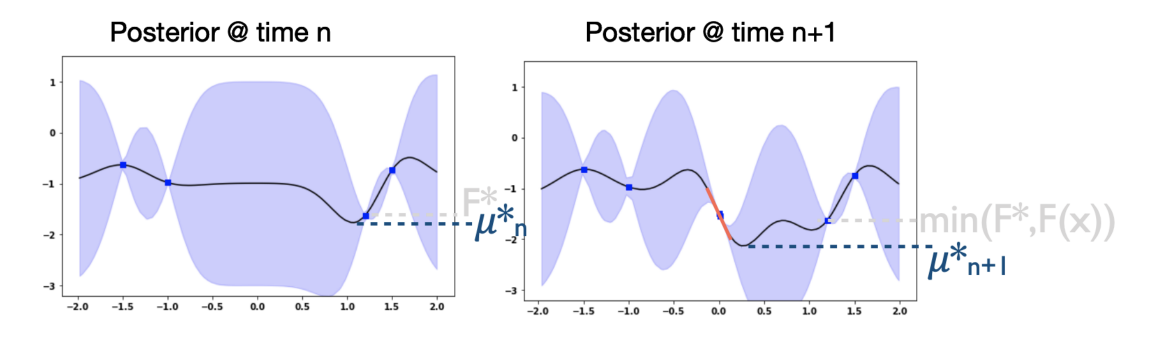
# **Knowledge Gradient (KG) Acquisition Function**

#### Posterior @ time n



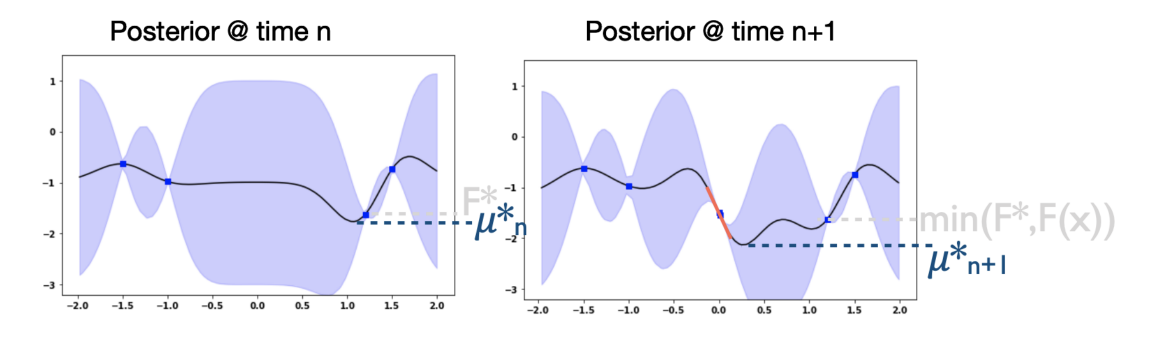
• Loss if we stop now:  $\mu_n^{\star} = \min_{x \in \Omega} \mu_n(x)$ 

## **Knowledge Gradient (KG) Acquisition Function**



- Loss if we stop now:  $\mu_n^{\star} = \min_{x \in \Omega} \mu_n(x)$
- Loss if we stop after sampling f(x):  $\mu_{n+1}^{\star} = \min_{x \in \Omega} \mu_{n+1}(x)$

## **Knowledge Gradient (KG) Acquisition Function**



- Loss if we stop now:  $\mu_n^{\star} = \min_{x \in \Omega} \mu_n(x)$
- Loss if we stop after sampling f(x):  $\mu_{n+1}^{\star} = \min_{x \in \Omega} \mu_{n+1}(x)$
- Expected reduction in loss due to sampling:  $\mathbb{E}_n[\mu_n^{\star} \mu_{n+1}^{\star} \mid \text{sample } x]$

### Do you see any challenges with KG?

- The main disadvantage of KG is that we lose the analytic formula that we were able to derive for EI
  - Makes is harder to maximize than EI, as we now have a two-stage optimization
- Are there alternative strategies for handling observation noise?
  - Yes, many other strategies exist including:
    - Modifications to El to mitigate impact of noise
    - Alternative acquisition functions

# Use a "plug-in" value for the incumbent

• Computing EI with noise is challenging because we no longer know the incumbent  $f^*$ ; a simple way to address this is to replace with an alternative value such as the GP estimate of the best function value

$$\hat{f}^{\star} = \min_{x \in \Omega} \mu_n^{\star}$$

- The main issue with this approach is that it tends to underestimate the improvement potential in regions of  $\Omega$  with large uncertainty
  - Because mean smooths out noise, potentially ignores regions where there could be significant improvement due to high variance in  $f(x) \rightarrow$  over-exploitation

## **Noisy Expected Improvement (NEI)**

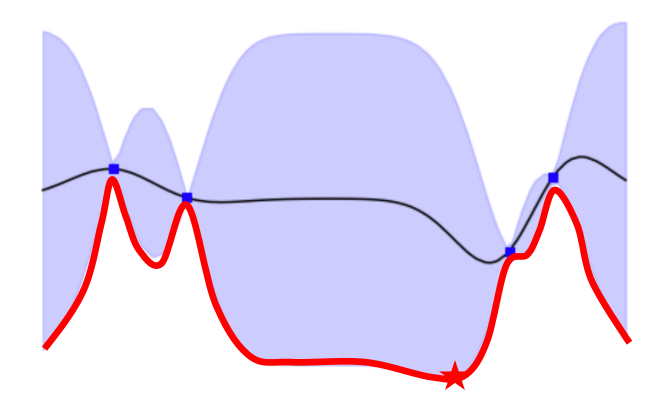
The noisy expected improvement function

$$NEI(x|\mathcal{D}) = \mathbb{E}_{\mathbf{f}^n}[EI(x|\mathbf{f}^n)|\mathcal{D}] = \int_{\mathbf{f}^n} EI(x|\mathbf{f}^n)p(\mathbf{f}^n|\mathcal{D}) d\mathbf{f}^n$$

Where  $\mathrm{EI}(x|\mathbf{f}^n)$  denotes the standard El expression assuming noiseless observations of the function  $\mathbf{f}^n$  and  $p(\mathbf{f}^n|\mathcal{D}) \sim N(\boldsymbol{\mu}_f^n, \Sigma_f^n)$  is the GP posterior prediction of the function values given noisy data  $\mathcal{D}$ 

• By incorporating noise into the decision-making process, NEI promotes a more exploratory behavior, especially in early iterations when the model is still uncertain about the objective function landscape.

#### **Alternative Acquisition: Lower Confidence Bound (LCB)**



• Simple idea: Just directly minimize a lower bound on the function

$$\min_{x \in \Omega} \mu_n(x) - \sqrt{\beta_{n+1}} \sigma_n(x)$$

## We can establish rigorous bounds on "regret" for LCB

- Lower confidence bound:  $l_n(x) = \mu_{n-1}(x) \sqrt{\beta_n}\sigma_{n-1}(x)$
- Upper confidence bound:  $u_n(x) = \mu_{n-1}(x) + \sqrt{\beta_n}\sigma_{n-1}(x)$
- Assume that true function satisfies  $f(x) \in [l_n(x), u_n(x)]$ (can prove this holds with high probability for sufficiently large  $\beta_n$ )
- Performance measure: Regret  $r_n$  defined as distance to optimal solution:

$$r_n = f(x_n) - f(x^*)$$

## We can establish rigorous bounds on "regret" for LCB

• The following sequence of inequalities hold:

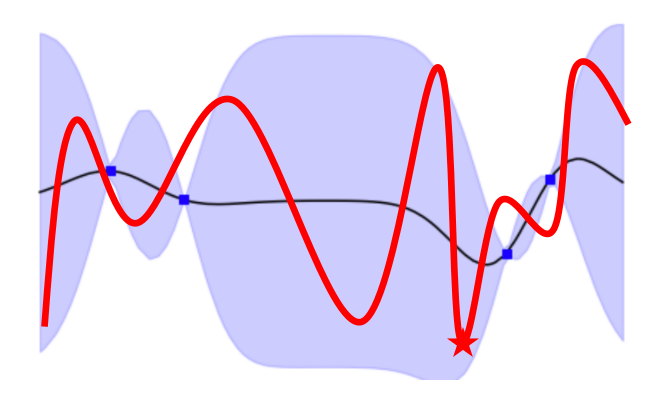
$$r_n = f(x_n) - \min_{x \in \Omega} f(x)$$
 [Definition of regret] 
$$\leq u_n(x_n) - \min_{x \in \Omega} f(x)$$
 [Property of upper bound] 
$$\leq u_n(x_n) - \min_{x \in \Omega} l_n(x)$$
 [Property of lower bound] 
$$= u_n(x_n) - l_n(x_n)$$
 [Definition of our sample choice  $x_n = \operatorname{argmin}_x l_n(x)$ ] 
$$= 2\sqrt{\beta_n} \sigma_{n-1}(x_n)$$
 [Difference between bounds given by standard deviation]

## Be careful with LCB in practice...

• Although theoretical values for  $\{\beta_n\}$  exist, they are often too conservative to be very useful in practice  $\rightarrow$  not great short-term performance

• In published experiments, it is common to set  $\sqrt{\beta_n} \approx 2$  as a heuristic, which can result in reasonable performance in certain problems; however, performance can be quite sensitive to the choice of  $\beta_n$  for some problems

# **Alternative Acquisition: Thompson Sampling (TS)**



• Minimize random sample of the GP, i.e.,  $f^{(n)} \sim \mathcal{GP}(\mu_n(x), \sigma_n^2(x))$   $\min_{x \in \Omega} f^{(n)}(x)$ 

# Efficient generation of differentiable TS from GP posteriors

- Interesting recent work showing how to get an efficient, differentiable TS
  - I highly recommend the following paper that goes in-depth on GP sampling:
    - [Wilson et al., "Efficiently sampling functions from GP posteriors", ICML, 2020]
- Core argument is one can use Matheron's rule to break down posterior sample into a "prior" and "update" step that has the following form for GPs

$$(f \mid \mathbf{y})(\cdot) \approx \sum_{i=1}^{l} w_i \phi_i(\cdot) + \sum_{j=1}^{n} v_j k(\cdot, \mathbf{x}_j), \qquad \mathbf{v} = \left(\mathbf{K}_{n,n} + \sigma_n^2 \mathbf{I}\right)^{-1} (\mathbf{y} - \mathbf{\Phi} \mathbf{w} - \mathbf{\varepsilon})$$
random noise realizations, weight-space prior function-space update 
$$\mathbf{\varepsilon} \sim N(\mathbf{0}, \sigma_n^2 \mathbf{I})$$

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## **A Quick Primer on Information Theory**

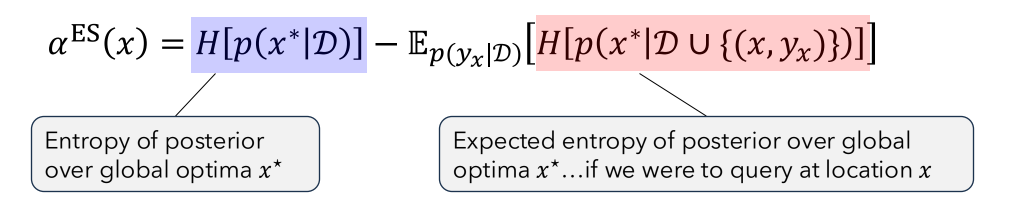
- The concept of information entropy was introduced by Claude Shannon in his 1948 paper "A Mathematical Theory of Communication"
- In essence, for a discrete random variable X that takes values in  $\Omega$ , we can compute its entropy as follows

$$H[X] = \mathbb{E}[-\log p(X)] = -\sum_{x \in \Omega} p(x) \log p(x)$$

- The entropy H[X] quantifies the average level of uncertainty or information associated with the variable's potential outcomes
  - Generalizes to continuous & multivariate distributions > differential entropy

# Can we use the notion of entropy to define our acquisition function in Bayesian optimization?

- Yes!
  - Similar to the idea of the improvement-based acquisitions, we can now attempt to decrease the entropy = increase our information by querying new points
- Expected information gain (EIG) expected decrease in entropy over posterior for global optima  $x^*$  if we were to query f at point x is:



#### Entropy search acquisition is challenging to compute!

[Predictive entropy search (PES) to the rescue]

- Two difficulties
  - Need to compute  $p(x^*|\mathcal{D} \cup \{(x,y_x)\})$  for many different values
  - Neither entropy expression can be computed analytically
- Luckily, we can use a nice trick to simplify computation
  - Notice that  $\alpha^{ES}(x)$  equivalent to mutual information (MI) between  $x^*|\mathcal{D}$  and  $y_x|\mathcal{D}$
  - MI is symmetric such that we can reverse the order and obtain:

$$\alpha^{\text{PES}}(x) = H[p(y_x|\mathcal{D})] - \mathbb{E}_{p(x^*|\mathcal{D})}[H[p(y_x|\mathcal{D}, x^*)]]$$

We can derive analytic expression using GP

Generate MC samples before optimization by finding optima of TS

Generate approximate samples using expectation propagation

## Entropy search concept easily generalizes to other statistics

• In general, can write out MI for some statistic of unknown function S(f):

$$\alpha^{S}(x) = MI(y_{x}; S(f)|x, \mathcal{D})$$

$$= H[p(y_{x}|\mathcal{D})] - \mathbb{E}_{p(S(f)|\mathcal{D})}[p(y_{x}|x, \mathcal{D}, S(f))]$$

- Predictive entropy search (PES) sets  $S(f) = x^*$ 
  - Hernández-Lobato et al., NeurIPS, 2014
- Max-value entropy search (MES) sets  $S(f) = y^* = \max_{x \in \Omega} f(x)$ 
  - Wang and Jegelka, ICML, 2017
- Joint entropy search (JES) sets  $S(f) = (x^*, y^*)$ 
  - Hvarfner, Hutter, and Nardi, NeurIPS, 2022

#### Visual Illustration of PES, MES, and JES

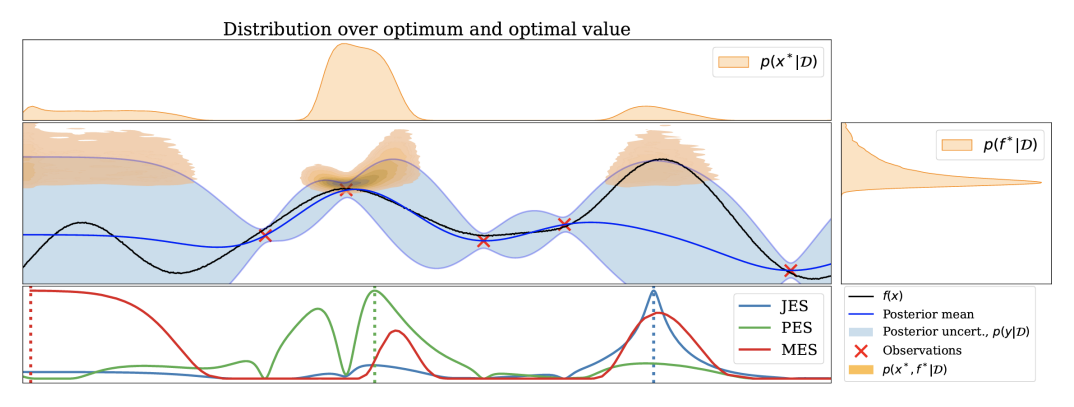
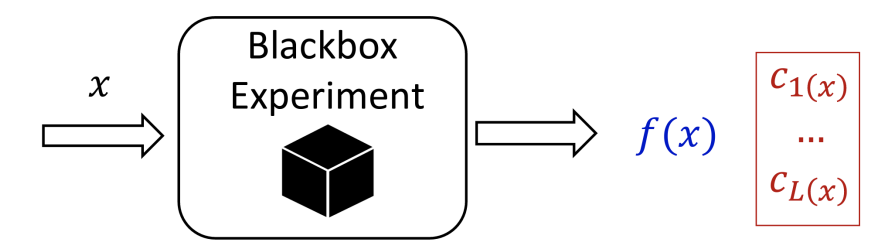


Figure 1: The densities considered by ES/PES (top), MES (right) and JES (center) on a one-dimensional toy example. The multimodal density  $p(x^*, f^*)$  is reduced to a heavy-tailed density over  $f^*$  for the density used by MES (right), which does not capture the multi-modality of the density over the optimum. The density over  $x^*$  used by PES (top) does not capture the apparent exploration/exploitation trade-off that exists between the modes. The acquisition functions and their next point selections are highlighted with dashed lines (bottom).

#### **Outline**

- Introduction to Bayesian optimization
  - White-box vs. black-box, prevalence of expensive functions, bird's eye view
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- Practical considerations
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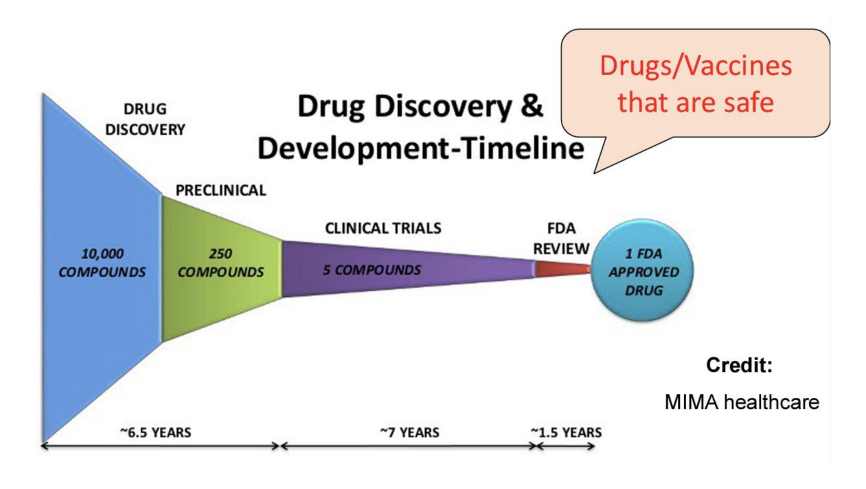
#### What about expensive black-box constraints?



Objective and constraints evaluation of design *x* 

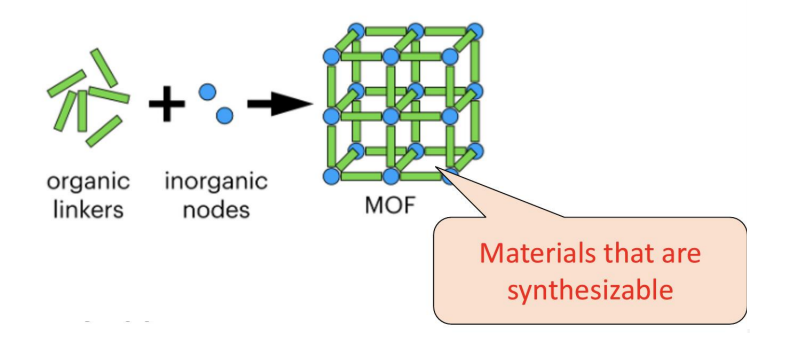
- **Goal:** find the constrained optima that minimizes the objective subject to satisfying constraints across the entire design space  $\Omega$ 
  - Evaluations of objective and constraint could be coupled or uncoupled

### **Example #1: Drug/Vaccine Design**



Accelerate the discovery of safe and promising designs

#### **Example #2: Design of Nanoporous Materials**



- Sustainability applications:
  - Storing gases (e.g., hydrogen powered cars)
  - Separating gases (e.g.,  $CO_2$  from flue gas in power plant)
  - Detecting gases (e.g., pollutants in outdoor spaces)

## **BO with Black-Box Constraints: Key Challenges**

- Modeling challenge: How to model the black-box constraints?
  - GP models can be straightforwardly trained for each constraint
  - Need to be a bit careful when the objective and/or constraints are correlated or there are a large number of constraints > efficient multi-output GPs
- Reasoning challenge: How to select the next input location using simultaneous information from all objective and constraint models?
  - Especially in the case that we have no feasible inputs that satisfy constraints from the previous experiments (also must be careful with observation noise!)

## **Expected Improvement with Constraints (EIC)**

Feasibility indicator function = 1 if  $c(x) \le 0$  and 0 otherwise Improvement over our best incumbent value  $f_n^* = \min_{i=1,...,n} f(x_i)$  s.t.  $c(x_i) \le 0$ 

$$EIC(x) = \mathbb{E}_n \{ \mathbf{1}_{\{c(x) \le 0\}}(x) \max\{0, f_n^* - f(x) \} \}$$
$$= \mathbb{E}_n \{ \mathbf{1}_{\{c(x) \le 0\}}(x) \} \mathbb{E}_n \{ \max\{0, f_n^* - f(x) \} \}$$

Conditional independence of objective and constraints\*

$$= \Pr_n\{c(x) \le 0\} \operatorname{EI}_n(x)$$

$$\Phi\left(-\frac{\mu_n^c(x)}{\sigma_n^c(x)}\right)$$

Standard expected improvement value that has analytic solution (shown on previous slide)

Probability of feasibility; analytic solution available for GP model using normal CDF

## **Expected Improvement with Constraints (EIC)**

• If a best feasible incumbent  $f^*$  exists, we can use previous expression

$$\mathrm{EIC}(x; f^{\star}, \mathcal{D}) = \mathrm{EI}(x; f^{\star}, \mathcal{D}) \prod_{i=1}^{L} \mathbb{P}(c_i(x) \leq 0 | \mathcal{D})$$

- If  $f^*$  does not exist, EIC is not well-defined. This issue is often addressed by maximizing the probability of constraint satisfaction  $\prod_{i=1}^L \mathbb{P}(c_i(x) \leq 0 | \mathcal{D})$ 
  - This is not ideal behavior, as it ignores learning about f in these iterations
- Suffers from a pathology when dealing with equality constraints  $c_i(x) = 0$ 
  - Must convert to two-sided inequality  $\rightarrow \mathbb{P}(c_i(x) \leq 0 | \mathcal{D}) \cdot \mathbb{P}(c_i(x) \geq 0 | \mathcal{D})$ , which can be extremely small especially when no feasible solutions known

## **Exact Penalty Bayesian Optimization (EPBO)\***

- Idea: Take an LCB (optimistic) perspective of the problem by relaxing the objective and constraints with high probability
  - Can build upon theory for deterministic constrained optimization where certain types of penalty functions are known to preserve exact solutions
  - Extend standard LCB theory to prove convergence to constrained global optima

#### Advantages:

- Does not depend on a potentially undefined incumbent  $f^*$
- Naturally handles both equality and inequality constraints
- Theoretical convergence guarantees, even in the presence of noise

#### **EPBO Acquisition Function: Penalized Lower Confidence Bound**

Convergence holds for sufficiently large  $\rho$ , can set  $\rho = \infty$  in practice (hard constraints)

$$\min_{x,\epsilon} \ \mu_{f,t}(x) - \beta_{t+1}^{1/2} \sigma_{f,t}(x) + \rho \|\epsilon\|_1,$$

s.t. 
$$|\mu_{h,t}(x)| - \beta_{t+1}^{1/2} \sigma_{h,t}(x) \leq \epsilon_h,$$
  
 $\mu_{g,t}(x) - \beta_{t+1}^{1/2} \sigma_{g,t}(x) \leq \epsilon_g,$   
 $\epsilon = [\epsilon_h^\top, \epsilon_g^\top]^\top \geq 0,$   
 $x \in \Omega,$ 

No penalty whenever we can find a mean prediction within a confidence band set by the predicted standard deviation

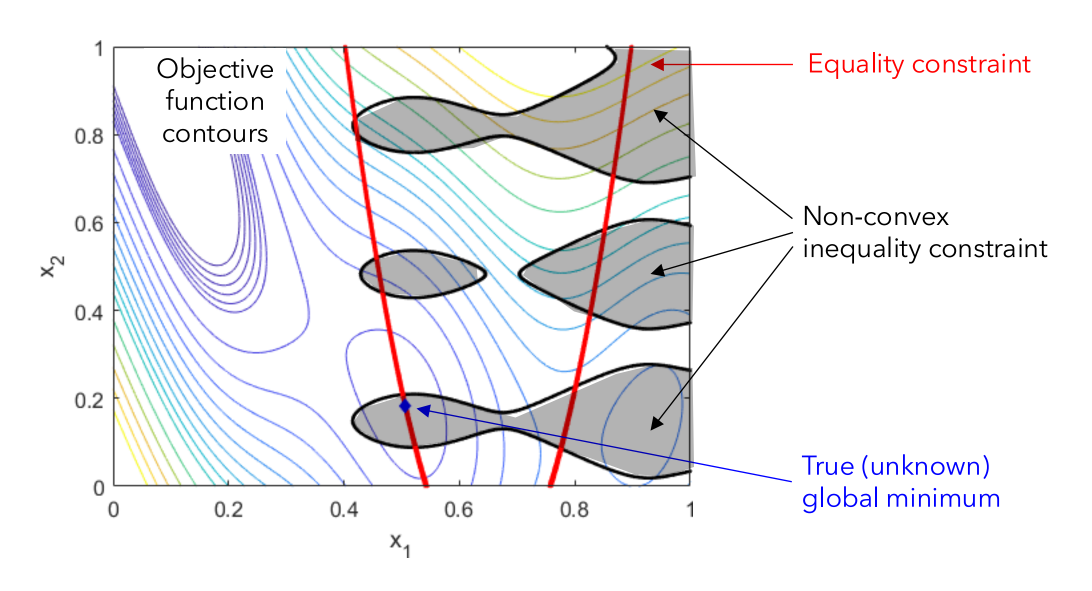
$$\mu_{h,t}(x) \in \sqrt{\beta_{t+1}}[-\sigma_{h,t}(x), \sigma_{h,t}(x)]$$

We can interpret inequality constraint representation as a <u>high probability</u> <u>relaxation</u> of the true feasible region

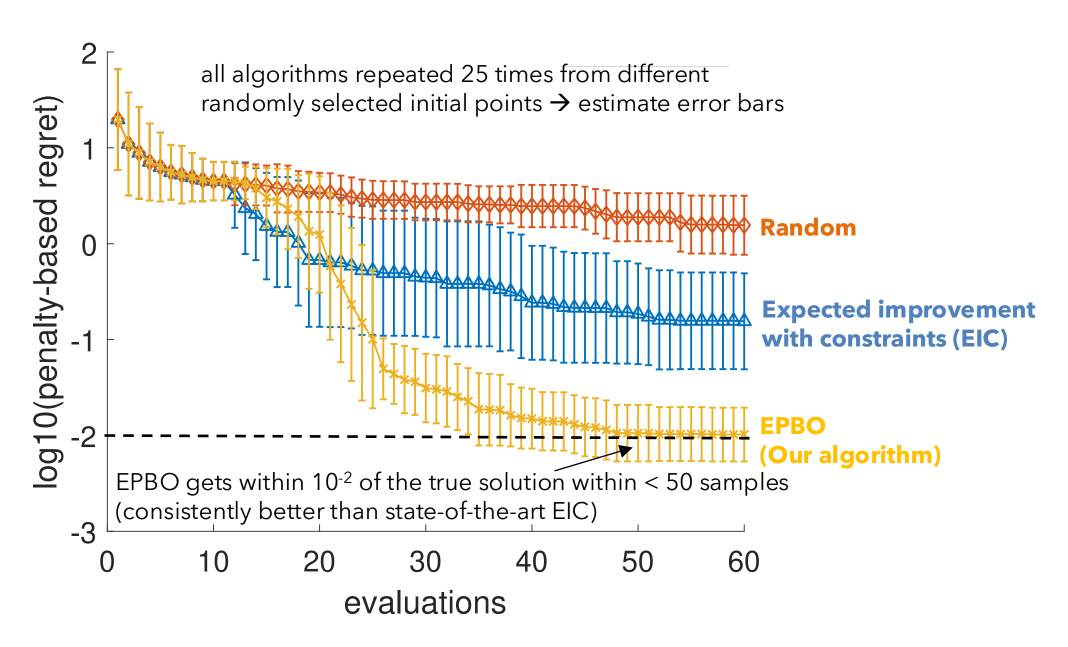
$${x: g(x) \le 0} \subseteq {x: \mu_{g,t}(x) \le \beta_{t+1}^{1/2} \sigma_{g,t}(x)}$$

#### **Example Problem with Equality and Inequality Constraints:**

**Modified Branin Function** 



#### **EPBO versus EIC: Results on Modified Branin Function**



## Can we use information theory for the constrained setting?

 Yes, PES with constraints (PESC) showed good performance on problems related to meta-optimization of machine learning and sampling methods

Now this is *constrained* optimizer

$$\alpha(\mathbf{x}) = \mathbf{H}\left[\mathbf{x}_{\star}|\mathcal{D}\right] - \mathbb{E}_{\mathbf{y}}\left\{\mathbf{H}\left[\mathbf{x}_{\star}|\mathcal{D}\cup(\mathbf{x},\mathbf{y})\right]\right\}$$

Vector of all evaluations

- Advantage: Nicely handles the <u>decoupled evaluation</u> case wherein  $\alpha(\cdot)$  becomes a sum of individual acquisition functions (one per function)
  - For example, imagine you are designing a cookie recipe and want to find the lowest calorie cookie possible that meets a constraint on the taste (>95% of test subjects like it)
  - Much easier for us to estimate calories (objective) than the taste (constraint) function, and these two functions can be evaluated at different inputs
- Disadvantage: Main disadvantage is its difficultly to implement, as it involves several
  approximations that can become numerically unstable

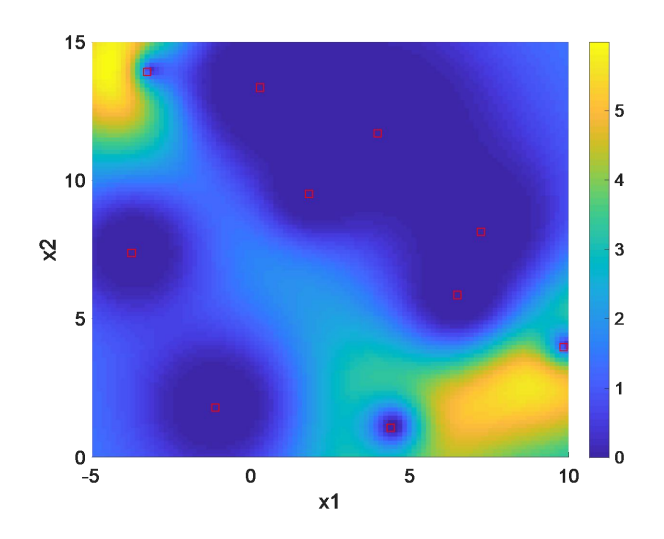
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### Two Major Tasks at Each Iteration in BO

- 1. Train hyperparameters of Gaussian process model(s)
  - The more accurate estimates we achieve for kernel + hyperparameters, the better the decision we can make for the next sample
  - In practice, we re-optimize the hyperparameters at every iteration (given the most recent data) by maximizing the log-likelihood function
  - A trick to reduce cost is to "warm start" the initial guess for the hyperparameters, which works well once they have roughly "stabilized"
- 2. Maximize the acquisition function,  $x_{n+1} \in \operatorname{argmax}_{x \in \Omega} \alpha_n(x)$ 
  - Many methods exist, no consensus in literature (use your favorite method)
  - Complexity of GP model + operators in acquisition function both important

- We expect  $\alpha_n(x)$  to be non-convex but typically differentiable
- Easily apply local optimization methods (e.g., L-BFGS, IPOPT)
- Important to perform some type of multi-start procedure to globalize
  - How effective is this method?



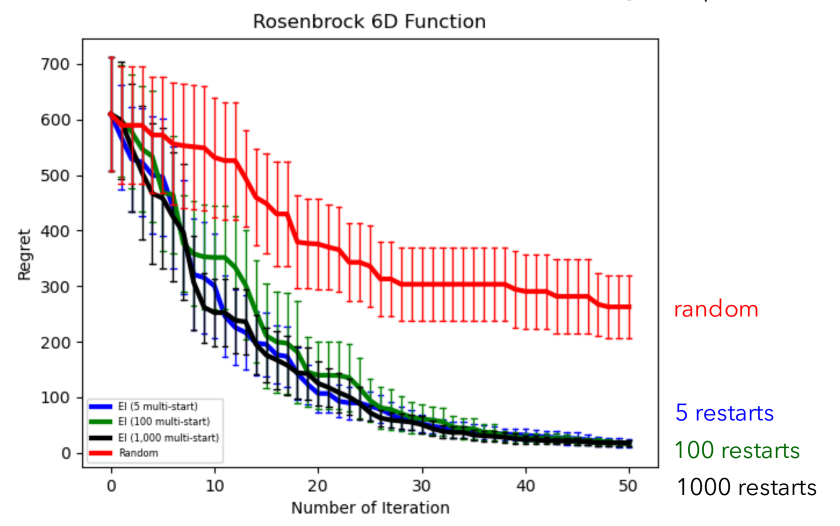
#### **How to Measure Performance?**

 Most BO papers use simple regret, which is the minimum over the recommended point after a finite number of iterations

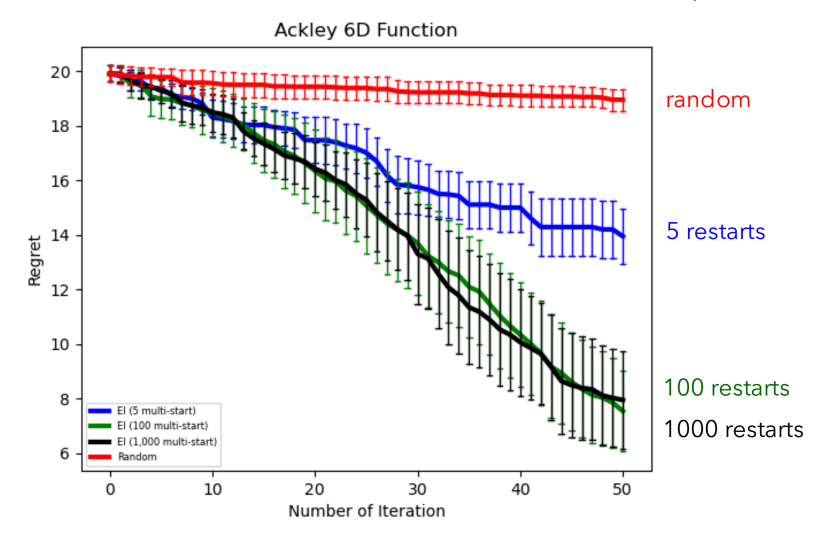
$$SimpleRegret_n = \min_{i \in \{1, ..., n\}} r_i = \min_{i \in \{1, ..., n\}} \{f(x_i) - \underline{f(x^*)}\}$$
Use best known solution when true solution unknown

- Since the regret sequence depends on the initial data, it is very common to do 50-100 replicates of the entire procedure (with initial data randomly generated) to assess average performance
  - We cannot conclude that every run performs not obvious what the distribution of performance is either, so often do not try to estimate

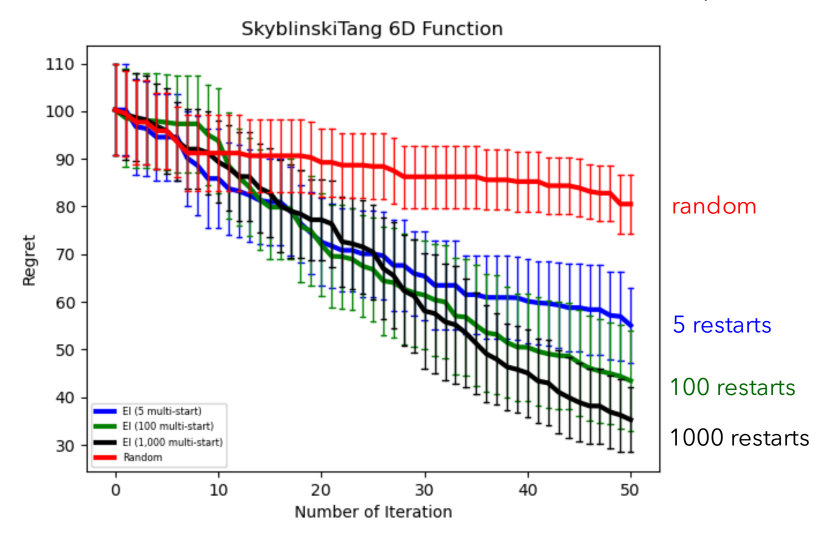
[100 replicates over initial data]



[100 replicates over initial data]



[100 replicates over initial data]



### Why Not Global Optimization?

Short Answer: Existing Methods Struggle

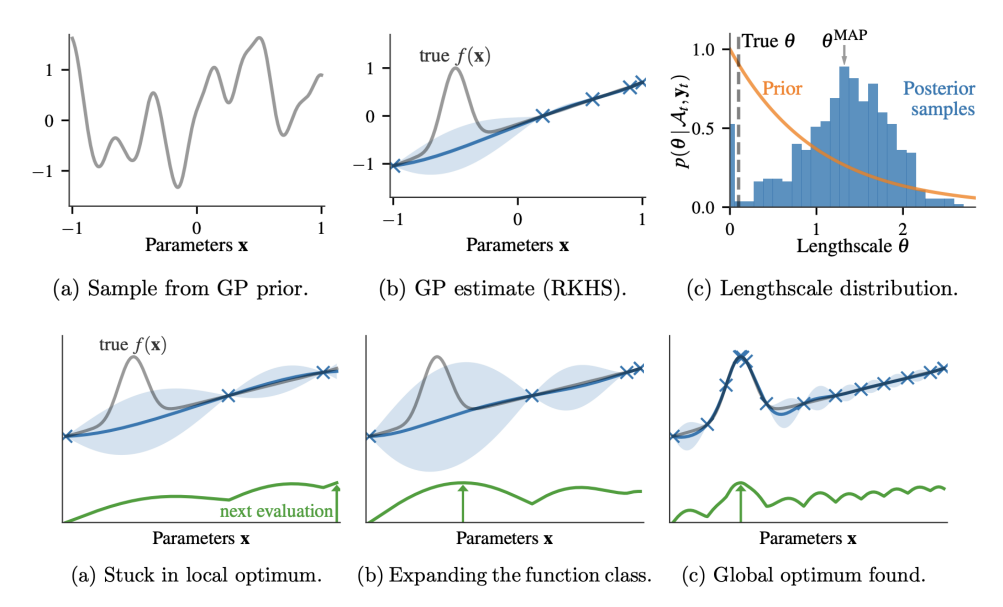
- McCormick relaxations end up being very weak for the posterior mean and covariance functions since they are the sum over several terms that can have alternating sign
- Would be great if we could (cheaply) construct underestimators that are not too weak → active research area in my group
- Since multi-start is trivially parallelizable (run local solves in parallel), it seems that is best approach to use for now
  - First-order methods, like Adam, also can take advantage of GPU acceleration, so end up having fast wall-times in practice

# Some Adaptive Modifications to (Potentially) Improve Performance

### **Dynamic Kernel Selection**

- Normally, we pick a single kernel and keep it fixed at every iteration
- There has been recent work suggesting that some performance gains can be obtained by training multiple GP models (with different kernels) and using some criteria to dynamically select the one to use at each iteration
- Heuristic in nature, so more work needed to find best ways to systematically select between GP models → usually based on some random select process to induce more exploration in the BO process

#### **Adaptation of Kernel Hyperparameters**



# **Dynamic Scheduling of Acquisition Functions**

- General practice in the BO literature has been for a practitioner to pick their favorite acquisition and use it for the entire optimization process
- Relatively recent work has discussed the value of adopting an adaptive sampling that chooses different acquisition functions at different iterations instead of attempting to pick "the best one"
- Different sampling strategies exist. Simple one is to assume m acquisition functions with weights  $\{w_i\}_{i=1}^m$ . The probability of sampling  $i^{\text{th}}$  acquisition is  $w_i/\sum_{i=1}^n w_i$  and, if the acquisition takes a successful move, then the weight is updated

#### **Exploitative Guard Against Model Misspecification**

[especially for information-theoretic approaches]

- In the paper that introduced the JES acquisition function, noted informationtheoretic acquisitions aimed at reducing uncertainty in location of optimum
- Thus, information-theoretic acquisitions are less likely to query the perceived optimum than other approaches like EI, which can greatly impact their performance in the case of a misspecified surrogate model
- To remedy this, they propose to use a  $\gamma$ -exploit approach that implies, with probability  $\gamma$ , the algorithm will query the point that maximizes the posterior mean to confirm its belief on the optimum location
  - If the model is misspecified, these exploitative steps help the algorithm to reconsider its beliefs rather than continuing to act based on faulty ones  $\rightarrow$  suggest  $\gamma = 0.1$

Extra material (if time permits)\*

Knowledge Gradient Maximization [More Advanced Optimization]

#### Let's recall the Knowledge Gradient (KG) acquisition function

• Earlier in this slide deck, we saw that the KG acquisition function is:

$$KG_{n}(x) = \mathbb{E}_{n} \{ \mu_{n}^{\star} - \mu_{n+1}^{\star} | x_{n+1} = x \}$$

$$= \mathbb{E}_{n} \left\{ \mu_{n}^{\star} - \min_{x' \in \Omega} \mu_{n+1}(x') | x_{n+1} = x \right\}$$

• Our goal is to maximize this function, so can ignore constant and convert the max to a min (due to the -1)

# Maximization of KG is a two-stage stochastic program

$$\min_{x \in \Omega} \mathbb{E}_n \left\{ \min_{x' \in \Omega} \mu_{n+1}(x') | x_{n+1} = x \right\}$$

How can we express the expectation in terms of things we can compute?

$$\mu_{n+1}(x') = \mu_n(x') + \tilde{\sigma}_n(x', x_{n+1})Z, \qquad Z \sim \mathcal{N}(0, 1)$$

$$\tilde{\sigma}_n(x, x') = \frac{k_n(x, x')}{\sqrt{k_n(x', x') + \sigma^2}}$$

# Maximization of KG is a two-stage stochastic program

$$\min_{x_{n+1}\in\Omega} \mathbb{E}_Z \left\{ \min_{x'\in\Omega} \{ \mu_n(x') + \tilde{\sigma}_n(x', x_{n+1})Z \right\}$$

- Two main approaches for solving this problem:
- 1. Stochastic gradient descent (SGD) (+ envelope theorem)
- 2. Sample average approximation (SAA)

this has become more popular recently

# Sample average approximation for KG acquisition

$$\min_{x_{n+1} \in \Omega} \frac{1}{N} \sum_{i=1}^{N} \left\{ \min_{x^{(i)} \in \Omega} \{ \mu_n(x^{(i)}) + \tilde{\sigma}_n(x^{(i)}, x_{n+1}) Z_i \right\}$$



The point that minimizes the next mean function depends on our sample selection

$$\min_{\substack{x_{n+1},x^{(1)},\dots,x^{(N)}\in\Omega\\}}\frac{1}{N}\sum_{i=1}^N\{\mu_n(x^{(i)})+\tilde{\sigma}_n(x^{(i)},x_{n+1})Z_i\}$$
 "here-and-now" "wait-and-see"