Computation-Aware Gaussian Process Inference

Jonathan Wenger

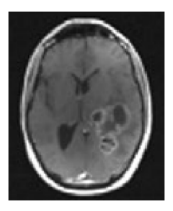








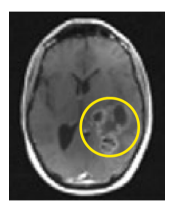
Accurate Reconstruction





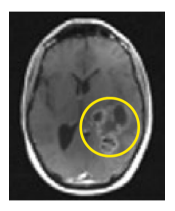


Accurate Reconstruction



Motivation

Accurate Reconstruction





(Radmanesh et al., 2022)

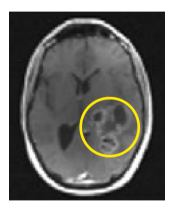
Subsampled Reconstruction (100x)



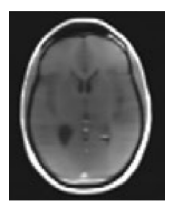


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Accurate Reconstruction



Learned Reconstruction (100x)

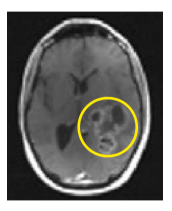


Importance of Uncertainty Quantification

Crucial information to benefit from the 100x acceleration is missing!

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Accurate Reconstruction



Learned Reconstruction (100x)



Uncertainty quantification is essential to make critical decisions.

Gaussian Process Regression

Supervised learning of an unknown function $f : \mathbb{R}^d \to \mathbb{R}$ with uncertainty quantification.

earning an unknown function from data.

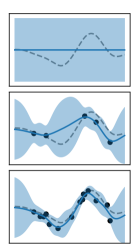


Goal: Supervised learning from n data points (X, y)

Prior: Gaussian process $f \sim \mathcal{GP}(\mu, k)$

Likelihood: Observations $\mathbf{y} = f(\mathbf{X}) + \boldsymbol{\varepsilon} \sim \mathcal{N}(f(\mathbf{X}), \sigma^2 \mathbf{I})$

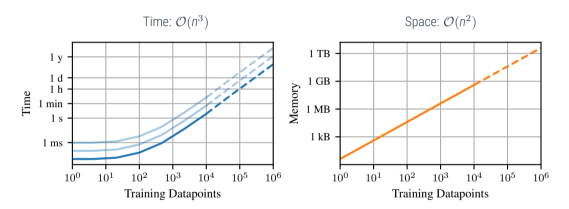
Posterior:
$$f \mid \mathbf{X}, \mathbf{y} \sim \mathcal{GP}(\mu_*, k_*)$$
 with
 $\mu_*(\cdot) = \mu(\cdot) + k(\cdot, \mathbf{X})\hat{\mathbf{K}}^{-1}(\mathbf{y} - \mu(\mathbf{X}))$
 $k_*(\cdot, \cdot) = k(\cdot, \cdot) - k(\cdot, \mathbf{X})\hat{\mathbf{K}}^{-1}k(\mathbf{X}, \cdot)$
where $\hat{\mathbf{K}} = \mathbf{K} + \sigma^2 \mathbf{I} \in \mathbb{R}^{n \times n}$.



Computational Cost of Gaussian Processes



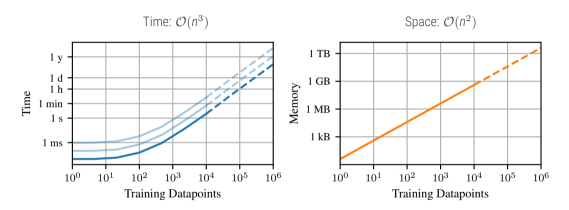
Uncertainty quantification can be expensive.



Computational Cost of Gaussian Processes



Uncertainty quantification can be expensive.

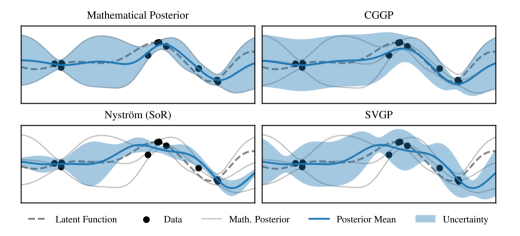


We need to approximate the posterior.

Approximate Gaussian Process Inference

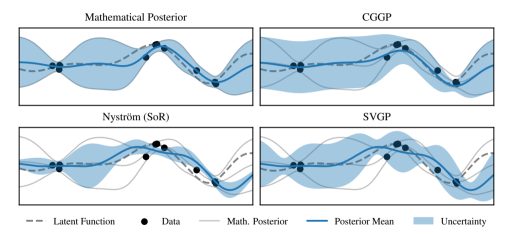
Impact of approximations on uncertainty quantification and decision-making.





Approximate Gaussian Process Inference

Impact of approximations on uncertainty quantification and decision-making.



Approximations introduce error, which impacts downstream decisions.

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Question 1:

How can we perform Gaussian process inference at scale?



Question 1:

How can we perform Gaussian process inference at scale?

Question 2:

How can we quantify the inevitable approximation error?

Q1: Gaussian Process Inference at Scale?

Efficiently approximating the posterior of a Gaussian process.

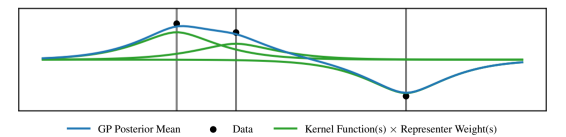


The posterior mean is a linear combination of kernel functions centered at datapoints.

$$f \mid \mathbf{X}, \mathbf{y} \sim \mathcal{GP}(\mu_*, k_*)$$

$$\mu_*(\cdot) = \mu(\cdot) + k(\cdot, \mathbf{X}) \frac{\hat{\mathbf{K}}^{-1}(\mathbf{y} - \mu(\mathbf{X}))}{\text{representer weights } \mathbf{v}_*} = \mu(\cdot) + \sum_{j=1}^n k(\cdot, \mathbf{X}_j)(\mathbf{v}_*)_j$$

 \mathbf{n}



Interlude: Method of Conjugate Gradients

Efficiently solving linear systems with positive definite system matrix via matrix-vector multiplies.

Goal: Approximately solve linear system Ax = b, where A symmetric positive definite.

Idea: Rephrase as quadratic optimization problem and optimize. Let

$$f(\mathbf{x}) = \frac{1}{2}\mathbf{x}^{\mathsf{T}}\mathbf{A}\mathbf{x} - \mathbf{b}^{\mathsf{T}}\mathbf{x}$$

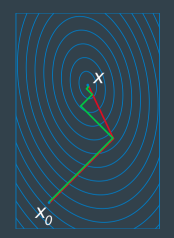
then $\nabla f(\mathbf{x}) = \mathbf{0} \iff A\mathbf{x} = \mathbf{b} \iff r(\mathbf{x}) \coloneqq \mathbf{b} - A\mathbf{x} = \mathbf{0}$.

Question: How should we optimize?

Gradient descent: Follow $d_i = r(x_i) = -\nabla f(x_i)$ s.t. $\langle d_i, d_j \rangle = 0$.

2 Conjugate direction method: Follow d_i s. t. $\langle d_i^{\mathsf{T}} d_j \rangle_A = d_i^{\mathsf{T}} A d_j = 0$ for $i \neq j$. \Rightarrow convergence in at most *n* steps.

Conjugate gradient method: First step $d_0 = r(x_0)$.



Oleg Alexandrov, commons.wikimedia.org/w/index.php?curid=2267598





Approximating Representer Weights

Iterative linear solvers can approximate the representer weights.

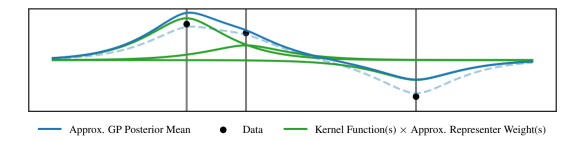


(Gardner et al., 2018; Charlier et al., 2021)

$$\mu_*(\cdot) = \mu(\cdot) + k(\cdot, \mathbf{X}) \frac{\hat{\mathbf{K}}^{-1}(\mathbf{y} - \mu(\mathbf{X}))}{r_{\text{encesenter weights } \mathbf{y}}} \approx \mu(\cdot) + k(\cdot, \mathbf{X}) \mathbf{v}_i$$

representer weights **v***

Observation: Can use iterative linear solvers (e.g. CG) to approximate the representer weights $v_* \approx v_i$.



Approximating Representer Weights

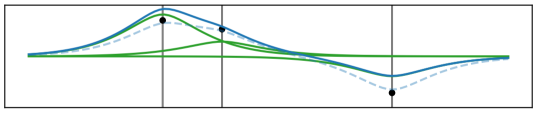
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Approx. GP Posterior Mean
 Data
 Kernel Function(s) × Approx. Representer Weight(s)

Benefit: Time complexity $\mathcal{O}(n^2)$ and space complexity $\mathcal{O}(nd)$.

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Approximating Representer Weights

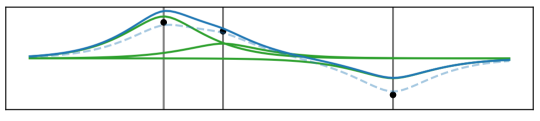
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Approx. GP Posterior Mean
 Data
 Kernel Function(s) × Approx. Representer Weight(s)

Question: Can we quantify the impact of this approximation on the posterior?

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Q2: Can We Quantify Approximation Error?

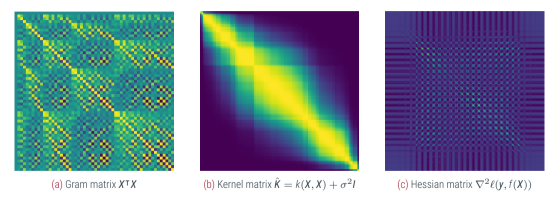
Probabilistic error quantification at prediction time using probabilistic linear solvers.

Probabilistic Linear Solvers for Machine Learning

Leveraging structure and quantifying approximation error.



Problem: Solve linear system(s) $Ax_* = b$ for $x_* \in \mathbb{R}^n$.

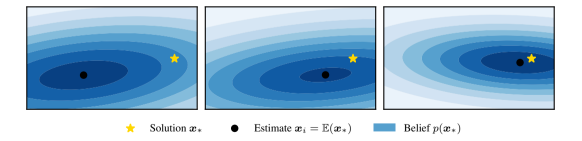


Linear systems in ML are large-scale, have model-induced structure and are often solved repeatedly.

Interpreting solving linear systems numerically as statistical inference.

Core Insights of Probabilistic Numerics

> The solution to any numerical problem is fundamentally uncertain.

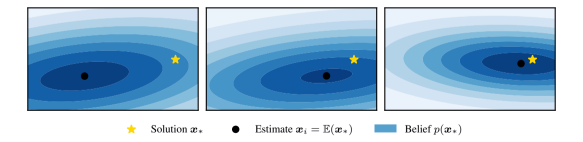


Interpreting solving linear systems numerically as statistical inference.

(Hennig, 2015; Cockayne et al., 2019; Wenger et al., 2020)

Core Insights of Probabilistic Numerics

- ► The solution to any numerical problem is fundamentally **uncertain**.
- Numerical algorithms are learning agents, which actively collect data and make predictions.



Estimating representer weights with a probabilistic linear solver.

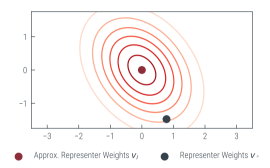


(Wenger et al., 2022a

Goal: Solve
$$\hat{K}v_* = y - \mu$$
 approximately.

Prior:

 $\mathbf{v}_* \sim \mathcal{N}(\mathbf{v}_0, \mathbf{\Sigma}_0)$



Estimating representer weights with a probabilis<u>tic linear solver.</u>



(Wenger et al., 2022a)

Goal: Solve
$$\hat{K}v_* = y - \mu$$
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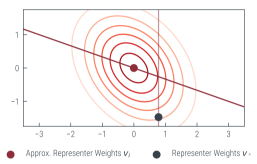
Prior: $v_* \sim \Lambda$

$$m{v}_* \sim \mathcal{N}(m{v}_0, m{\Sigma}_0)$$

Likelihood: Observe representer weights via arbitrarily chosen actions $s_i \in \mathbb{R}^n$:

$$\begin{aligned} \alpha_i &\coloneqq \mathbf{s}_i^{\mathsf{T}} \mathbf{r}_{i-1} = \mathbf{s}_i^{\mathsf{T}} ((\mathbf{y} - \boldsymbol{\mu}) - \hat{\mathbf{K}} \mathbf{v}_{i-1}) \\ &= \mathbf{s}_i^{\mathsf{T}} \hat{\mathbf{K}} (\mathbf{v}_* - \mathbf{v}_{i-1}) \end{aligned}$$

 $p(\alpha_i \mid \mathbf{v}_*) = \lim_{\varepsilon \to 0} \mathcal{N}(\alpha_i; 0, \varepsilon)$



Estimating representer weights with a probabilistic linear solver.



(Wenger et al., 2022a)

3

Goal: Solve
$$\hat{K}v_* = y - \mu$$
 approximately.

 $\mathbf{v}_{*} \sim \mathcal{N}(\mathbf{v}_{0}, \mathbf{\Sigma}_{0})$ Prior:

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Posterior: Affine Gaussian inference!



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-2

-1

-3

Estimating representer weights with a probabilistic linear solver.



3

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Goal:Solve
$$\hat{K} \mathbf{v}_* = \mathbf{y} - \boldsymbol{\mu}$$
 approximately.Prior: $\mathbf{v}_* \sim \mathcal{N}(\mathbf{v}_0, \boldsymbol{\Sigma}_0)$ Likelihood:Observe representer weights via arbitrarily chosen actions $\mathbf{s}_i \in \mathbb{R}^n$: $\alpha_i := \mathbf{s}_i^\mathsf{T} \mathbf{r}_{i-1} = \mathbf{s}_i^\mathsf{T} ((\mathbf{y} - \boldsymbol{\mu}) - \hat{K} \mathbf{v}_{i-1})$ $= \mathbf{s}_i^\mathsf{T} \hat{K}(\mathbf{v}_* - \mathbf{v}_{i-1})$ $p(\alpha_i \mid \mathbf{v}_*) = \lim_{\varepsilon \to 0} \mathcal{N}(\alpha_i; 0, \varepsilon)$

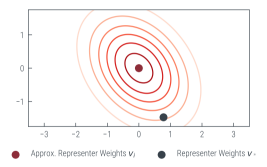
Posterior:
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, where
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 $\mathbf{\Sigma}_i = \mathbf{\Sigma}_{i-1} - \mathbf{\Sigma}_{i-1}\hat{\mathbf{K}}\mathbf{s}_i(\mathbf{s}_i^\mathsf{T}\hat{\mathbf{K}}\mathbf{\Sigma}_{i-1}\hat{\mathbf{K}}\mathbf{s}_i)^{-1}\mathbf{s}_i^\mathsf{T}\hat{\mathbf{K}}\mathbf{\Sigma}_{i-1}$

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The Gaussian process prior makes assumptions about the representer weights.



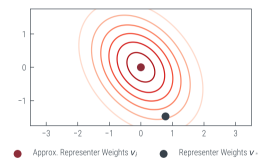
Question: How to choose the linear solver prior?



The Gaussian process prior makes assumptions about the representer weights.

Question: How to choose the linear solver prior?

Remember:
$$\mathbf{y} = f(\mathbf{X}) + \boldsymbol{\varepsilon}$$
, where $\boldsymbol{\varepsilon} \sim \mathcal{N}(\mathbf{0}, \sigma^2 \mathbf{I})$.
 $\Rightarrow \mathbf{y} - \boldsymbol{\mu} \sim \mathcal{N}(\mathbf{0}, \mathbf{k}(\mathbf{X}, \mathbf{X}) + \sigma^2 \mathbf{I}) = \mathcal{N}(\mathbf{0}, \hat{\mathbf{K}})$



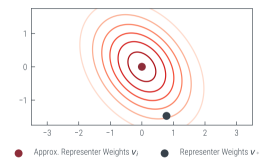


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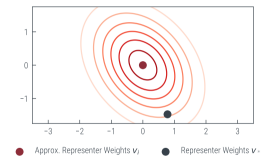
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Setting $\mathbf{v}_0 = 0$ and $\mathbf{\Sigma}_0 = \hat{\mathbf{K}}^{-1}$, we have

$$\begin{aligned} \mathbf{v}_i &= \mathbf{S}_i (\mathbf{S}_i^{\mathsf{T}} \hat{\mathbf{K}} \mathbf{S}_i)^{-1} \mathbf{S}_i^{\mathsf{T}} (\mathbf{y} - \boldsymbol{\mu}) = \mathbf{C}_i (\mathbf{y} - \boldsymbol{\mu}) \\ \mathbf{\Sigma}_i &= \mathbf{\Sigma}_0 - \mathbf{S}_i (\mathbf{S}_i^{\mathsf{T}} \hat{\mathbf{K}} \mathbf{S}_i)^{-1} \mathbf{S}_i^{\mathsf{T}} = \mathbf{\Sigma}_0 - \mathbf{C}_i \end{aligned}$$

where S_i is the matrix of actions s_1, \ldots, s_i .





The Gaussian process prior makes assumptions about the representer weights.

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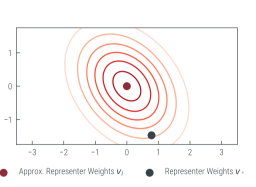
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IterGP: Computation-Aware Gaussian Process Inference

Quantifying uncertainty arising from observing finite data and performing a finite amount of computation.



(Wenger et al., 2022a)

Goal: Approximate the Gaussian process posterior $f \mid \mathbf{y} \sim \mathcal{GP}(\mu_*, k_*)$, where

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Idea: Propagate uncertainty about representer weights to posterior.

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1 Pathwise form of posterior: $(f \mid \mathbf{y})(\cdot) \stackrel{d}{=} f(\cdot) + k(\cdot, \mathbf{X})\hat{\mathbf{K}}^{-1}(\mathbf{y} - \boldsymbol{\mu})$

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2 Reparametrize posterior: $(f \mid \mathbf{v}_*)(\cdot) \stackrel{d}{=} f(\cdot) + k(\cdot, \mathbf{X})\mathbf{v}_*$

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(Wenger et al., 2022a)

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Idea: Propagate uncertainty about representer weights to posterior.

- Pathwise form of posterior: $(f \mid \mathbf{y})(\cdot) \stackrel{d}{=} f(\cdot) + k(\cdot, \mathbf{X})\hat{\mathbf{K}}^{-1}(\mathbf{y} \boldsymbol{\mu})$
- **2** Reparametrize posterior: $(f | \mathbf{v}_*)(\cdot) \stackrel{d}{=} f(\cdot) + k(\cdot, \mathbf{X})\mathbf{v}_*$
- 3 Marginalize representer weights belief: $p(f(\cdot)) = \int p(f(\cdot) | \mathbf{v}_*) p(\mathbf{v}_*) d\mathbf{v}_* = \mathcal{GP}(f; \mu_i, k_i)$,

$$\mu_{i}(\cdot) = \mu(\cdot) + k(\cdot, \mathbf{X})\mathbf{v}_{i}$$

$$k_{i}(\cdot, \cdot) = \underbrace{k(\cdot, \cdot) - k(\cdot, \mathbf{X})\hat{\mathbf{K}}^{-1}k(\mathbf{X}, \cdot)}_{\text{mathematical uncertainty}} + \underbrace{k(\cdot, \mathbf{X})\boldsymbol{\Sigma}_{i}k(\mathbf{X}, \cdot)}_{\text{computational uncertainty}} = \underbrace{k(\cdot, \cdot) - k(\cdot, \mathbf{X})\mathbf{C}_{i}k(\mathbf{X}, \cdot)}_{\text{combined uncertainty}}$$

Probabilistic Quantification of Approximation Error

The covariance can be interpreted as a squared error.

Combined Uncertainty

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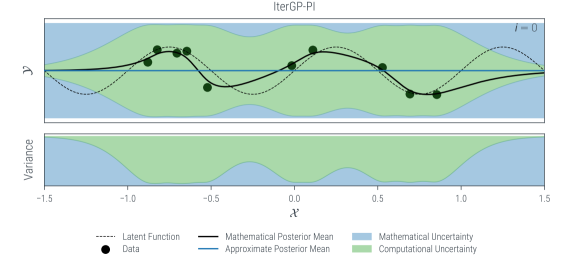
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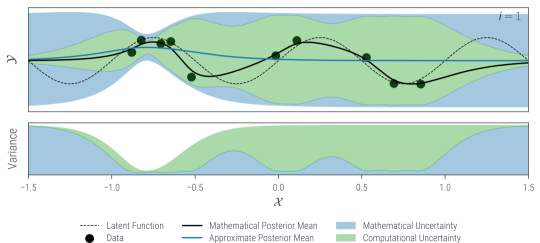
$$\overset{\text{comp}}{\underset{i}{\text{computational uncertainty}}} = \underbrace{k(\mathbf{X}, \mathbf{X}) \boldsymbol{\Sigma}_{i}k(\mathbf{X}, \mathbf{X})}_{\text{computational uncertainty}} = \underbrace{\mathbb{E}\left((v_{*} - v_{i})(v_{*} - v_{i})^{\mathsf{T}}\right)}_{\mathbb{E}\left((\mu_{*}(\mathbf{X}) - \boldsymbol{\mu}_{i}(\mathbf{X}))^{2}\right)}$$

Interpreting computational and combined uncertainty as error quantification.





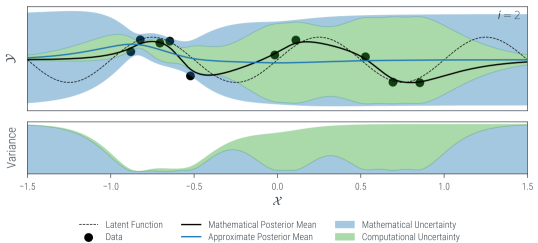
Interpreting computational and combined uncertainty as error quantification.





IterGP-PI

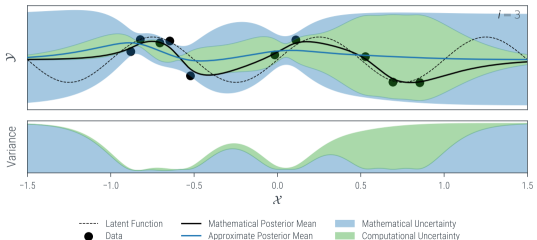
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lterGP-PI

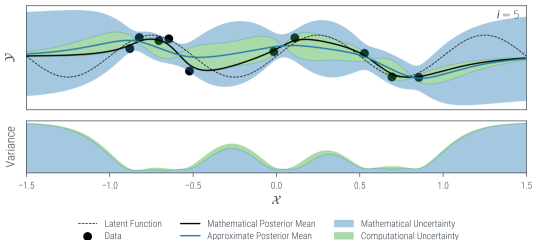
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IterGP-PI



Interpreting computational and combined uncertainty as error quantification.



IterGP-PI



Theoretical Analysis

Uncertainty as a bound on the relative predictive error.



Theorem (Relative Error Bound)

$$\sup_{g \in \mathcal{H}_{k^{\sigma}} : \|g\|_{\mathcal{H}_{k^{\sigma}}} \leq 1} \underbrace{g(\mathbf{x}) - \mu_*^g(\mathbf{x})}_{\text{error of math. post. mean } \odot} \sup_{g \in \mathcal{H}_{k^{\sigma}}} \frac{|g(\mathbf{x}) - \mu_*^g(\mathbf{x})|}{\|g\|_{\mathcal{H}_{k^{\sigma}}}} = \sqrt{k_*(\mathbf{x}, \mathbf{x}) + \sigma^2} \tag{1}$$

Theoretical Analysis

The combined uncertainty is a tight worst case bound on the relative error to the latent function



(Wenger et al., 2022a)

Theorem (Relative Error Bound) error of approximate posterior mean -+ $\sup_{g \in \mathcal{H}_{k^{\sigma}} : \|g\|_{\mathcal{H}_{k^{\sigma}}} \leq 1} \underbrace{g(\mathbf{X}) - \mu_{*}^{g}(\mathbf{X})}_{\text{error of math. post. mean } \bullet} + \underbrace{\mu_{*}^{g}(\mathbf{X}) - \mu_{i}^{g}(\mathbf{X})}_{\text{computational error } \bullet} = \sqrt{k_{i}(\mathbf{X}, \mathbf{X}) + \sigma^{2}}$ (1 Variance Latent Function Mathematical Posterior Mean Mathematical Uncertainty Data Approximate Posterior Mean Computational Uncertainty Combined Uncertainty

Computation-Aware Gaussian Process Inference - Jonathan Wenger - July 19, 2023



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What About:

How does IterGP relate to other numerical (approximation) methods, e.g. Cholesky, CGGP, SVGP?



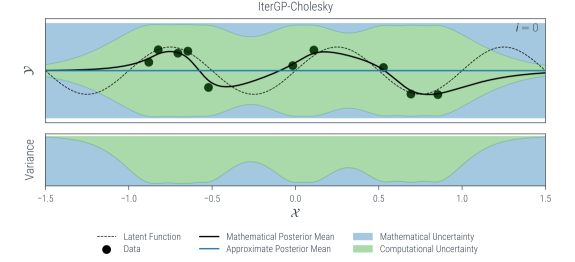
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What About:

- How does IterGP relate to other numerical (approximation) methods, e.g. Cholesky, CGGP, SVGP?
- ► Is quadratic time $O(n^2)$ the limit? Can we approximate more cheaply?

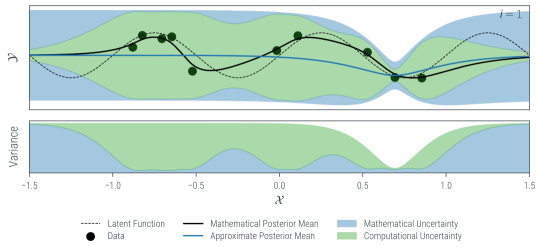
IterGP with unit vector actions recovers vanilla GP inference.





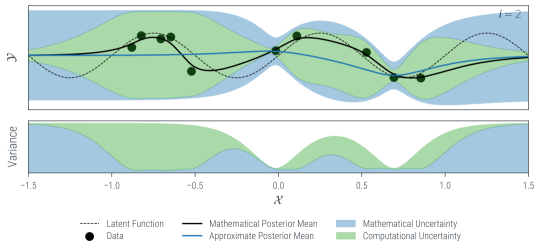
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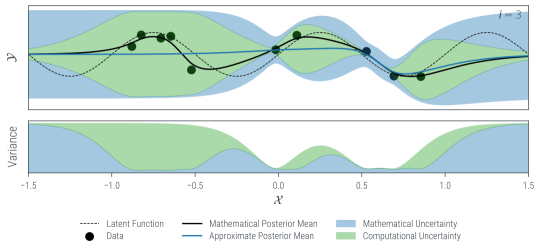
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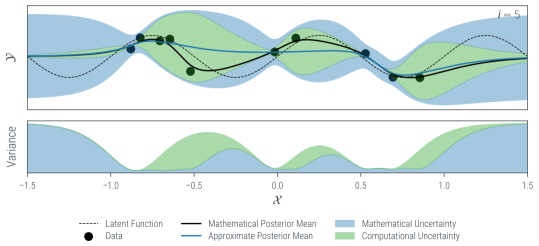
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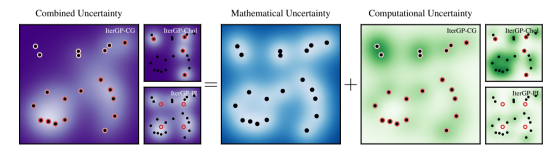


Policy Choice and Connection to Other Approximations



IterGP extends the most commonly used GP approximations to include computational uncertainty, with at most quadratic cost.

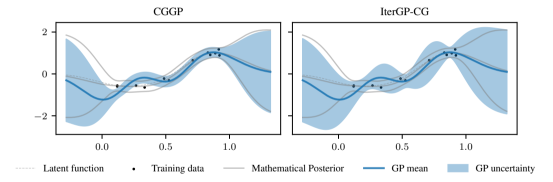
Method	Actions s i	Classic Analog
IterGP-Cholesky IterGP-EVD IterGP-CG IterGP-PseudoInput		(Partial) Cholesky / subset of data (Partial) Eigenvalue decomposition (Preconditioned) CG ≈ SVGP



CGGP versus IterGP-CG

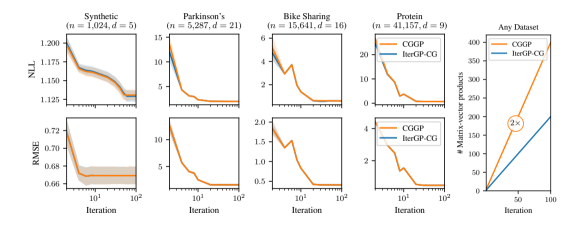
IterGP reduces the necessary computations for CG-based GP inference.





CGGP versus IterGP-CG

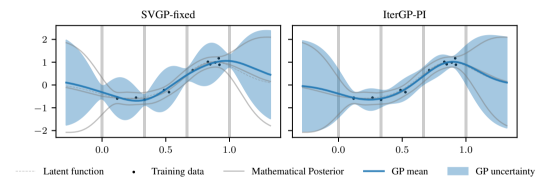
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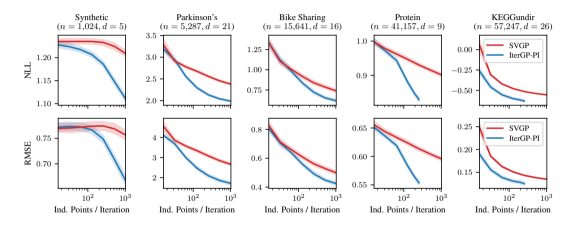


Quantifying computational uncertainty improves generalization of inducing point methods like SVGP (Titsias, 2009; Hensman et al., 2013).



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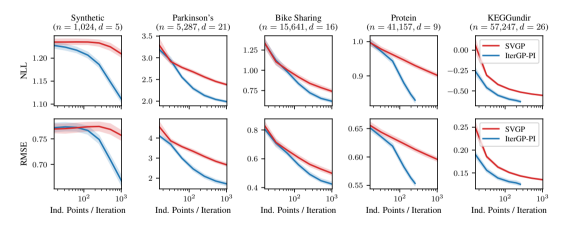


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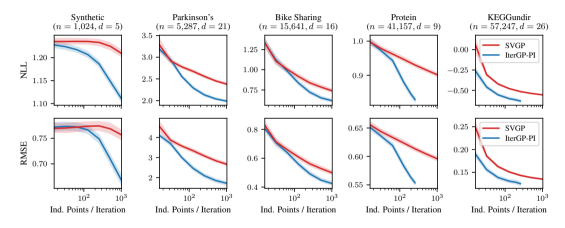


What about optimizing inducing point locations?

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SVGP versus IterGP-PI

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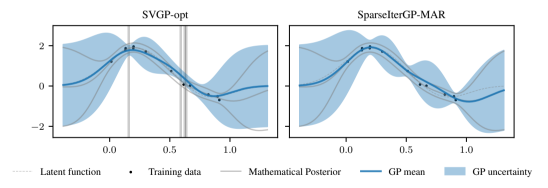


What about **computational cost**? SVGP: $\mathcal{O}(nm^2)$ versus IterGP-PI: $\mathcal{O}(n^2m)$.

UNIVERSITA TÜBINGEN Linear-time computation-aware GP inference with IterGP.



Policy: Unit vector actions $s_i = e_j$ which select points greedily as $j = \arg \max r_{i-1} \implies \mathcal{O}(nm^2)$.

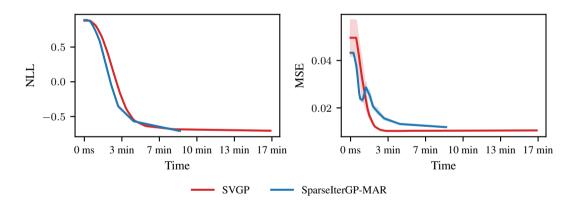


SVGP versus IterGP-MAR: Large-Scale Problem

Linear-time computation-aware GP inference with IterGP on a problem with $n \approx 10^5$ datapoints.



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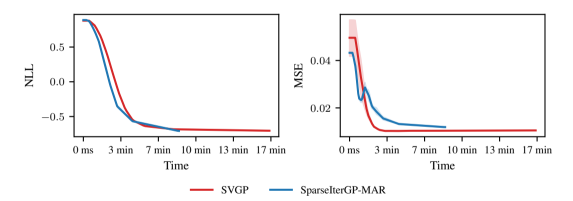


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Scalable GP approximation without inadvertently comprimising uncertainty quantification.

Bonus: Getting Philosophical

Blurring the lines between data and computation.

Working with Infinite Data

For IterGP it does not matter how large the dataset is, or whether we have it stored on our machine.

Theorem (Online GP Approximation with IterGP)

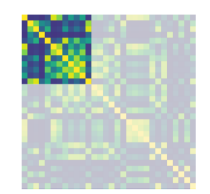
Let $n, n' \in \mathbb{N}$ and consider training data sets $X \in \mathbb{R}^{n \times d}, y \in \mathbb{R}^{n}$ and $X' \in \mathbb{R}^{n' \times d}, y' \in \mathbb{R}^{n'}$. Consider two sequences of actions $(\mathbf{s}_{i})_{i=1}^{n} \in \mathbb{R}^{n}$ and $(\tilde{\mathbf{s}}_{i})_{i=1}^{n+n'} \in \mathbb{R}^{n+n'}$ such that

$$ilde{\mathbf{s}}_i = egin{pmatrix} \mathbf{s}_i \ \mathbf{0} \end{pmatrix}$$

(2)

Then the posterior returned by IterGP for the dataset (X, y) using actions s_i is identical to the posterior returned by IterGP for the extended dataset using actions \tilde{s}_i :

ITERGP
$$(\mu, k, \mathbf{X}, \mathbf{y}, (\mathbf{s}_i)_i) = I$$
TERGP $\left(\mu, k, \begin{pmatrix} \mathbf{X} \\ \mathbf{X}' \end{pmatrix}, \begin{pmatrix} \mathbf{y} \\ \mathbf{y}' \end{pmatrix}, (\tilde{\mathbf{s}}_i)_i \right).$





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An alternative view of IterGP as a better model for the way we do inference instead of an approximation.

Observation: Only once we perform computation on data, does it enter our prediction.





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What if we modelled this situation with a Gaussian process?

 $\begin{aligned} & f \sim \mathcal{GP}(\mu, k) \\ & \tilde{\mathbf{y}} \mid f(\mathbf{X}) \sim \mathcal{N}\big(\mathbf{S}_i^\mathsf{T} f(\mathbf{X}), \sigma^2 \mathbf{S}_i^\mathsf{T} \mathbf{S}_i\big) \\ & f \mid \mathbf{X}, \tilde{\mathbf{y}} \sim \mathcal{GP}(\mu_i, k_i) \end{aligned}$



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IterGP's combined posterior is equivalent to exact GP regression for linearly projected data.



Takeaways

► Large-scale models are often as much about the approximation as they are about the data.



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- Policy design for downstream tasks and decision making problems.
 - Active learning
 - Bayesian optimization
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Extension to non-Gaussian likelihoods.



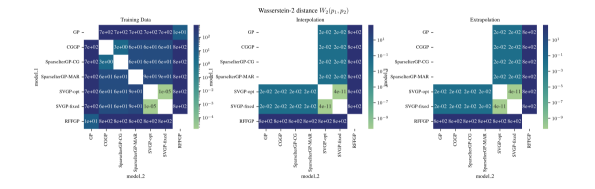
Comparison of GP Approximations

Gaussian Process Classification

Large-scale Model Selection

Comparison of GP Approximations: Wasserstein-2 Distance

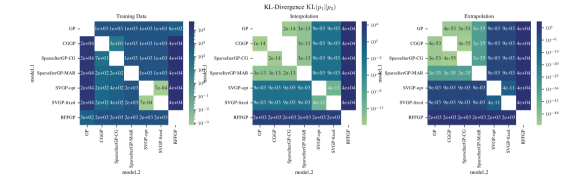
Comparison of different GP approximations at the training data, for interpolation and extrapolation.



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Comparison of GP Approximations: KL-Divergence

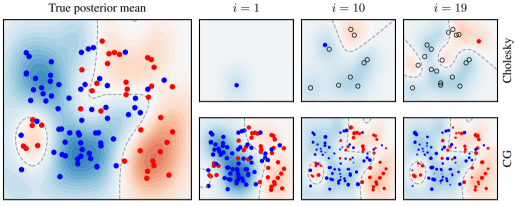
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Gaussian Process Classification

Extension to non-Gaussian likelihoods via Laplace Approximation.





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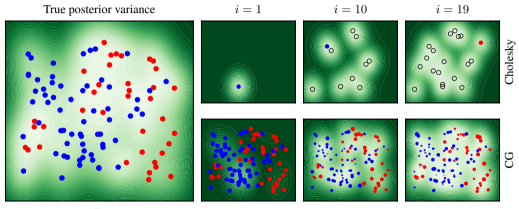
 x_2

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Gaussian Process Classification

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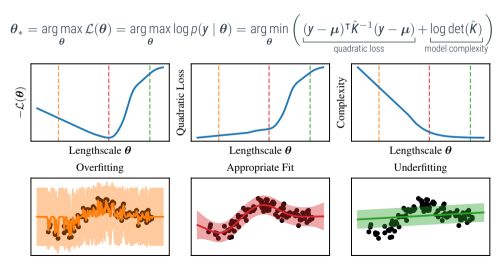


 x_1

Model Selection for Gaussian Processes



We can identify kernel hyperparameters by optimizing the log-marginal likelihood.



35

Large-scale GP Hyperparameter Optimization

A numerical linear algebra bottleneck.

Need to: Evaluate log-marginal likelihood and its derivative repeatedly.

- ► log-marginal likelihood $\mathcal{L}(\boldsymbol{\theta}) = -\frac{1}{2} \left(\boldsymbol{y}^{\mathsf{T}} \hat{\boldsymbol{K}}^{-1} \boldsymbol{y} + \log \det(\hat{\boldsymbol{K}}) + n \log(2\pi) \right)$
- $\blacktriangleright \quad \text{derivative } \tfrac{\partial}{\partial \theta} \mathcal{L}(\theta) = \tfrac{1}{2} y^{\mathsf{T}} \hat{K}^{-1} \tfrac{\partial \hat{K}}{\partial \theta} \hat{K}^{-1} y \tfrac{1}{2} \operatorname{tr}(\hat{K}^{-1} \tfrac{\partial \hat{K}}{\partial \theta})$

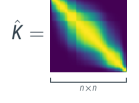
Challenge: Computationally costly operations with the kernel matrix.

- $\models \text{ linear solves } \mathbf{v} \mapsto \hat{\mathbf{K}}^{-1} \mathbf{v}$
- matrix traces log det $(\hat{K}) = \text{tr}(\log(\hat{K}))$ and $\text{tr}(\hat{K}^{-1} \frac{\partial \hat{K}}{\partial \theta_i})$

Linear solves and matrix traces can be computed solely via matrix-vector multiplication!

This is great because ...

- matrix-vector multiplies have complexity $\mathcal{O}(n^2)$.
- structured or sparse matrices are efficient to multiply with.
- the kernel matrix does not need to be stored in memory explicitly (Charlier et al., 2021).
- Computation Aware cash exploit parallelization, and modern hardware (GPUs).



lower time and space complexity



Preconditioning

How to encode and leverage structural prior knowledge about matrices.

Preconditioner

$$\hat{P} \approx \hat{K}$$

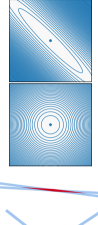
such that $\kappa(\hat{P}^{-1}\hat{K}) \ll \kappa(\hat{K})$ and \hat{P} is computationally tractable.

- Computing and storing \hat{P} is cheap.
- ► Linear solves $v \mapsto \hat{P}^{-1}v$ are efficient.
- Derived properties, such as the determinant or spectrum are known.

Asymptotic approx. error $g(\ell) \to 0$ of sequence of preconditioners $\hat{P}_{\ell} \to \hat{K}$:

 $\kappa(\hat{\boldsymbol{P}}_{\ell}^{-1}\hat{\boldsymbol{K}}) \leq (1 + \mathcal{O}(\boldsymbol{g}(\ell)) \|\hat{\boldsymbol{K}}\|_{\mathrm{F}})^2$

Known Use: Accelerate and stabilize linear solves via $\mathrm{CG} \Rightarrow \mathrm{bias}\ \mathrm{reduction}$

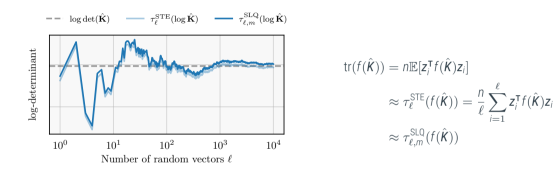






Stochastic Trace Estimation

Computing matrix traces tr $(f(\hat{K}))$ via matrix-vector multiplication.



Problems:

• Worst-case convergence in the number of random vectors is $\mathcal{O}(\ell^{-\frac{1}{2}})$

⇒ slows down training

Introduces stochasticity into hyperparameter optimization

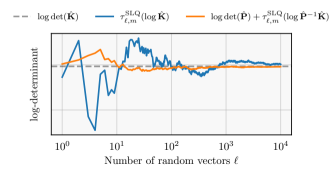
Preconditioned Log-Determinant Estimation

Variance-reduced stochastic trace estimation via preconditioning.

Idea: Decompose log-determinant into deterministic and stochastic approximation.

$$\log \det(\hat{K}) = \log \det(\hat{P}_{\ell}\hat{P}_{\ell}^{-1}\hat{K}) = \underbrace{\log \det(\hat{P}_{\ell})}_{\text{known}} + \underbrace{\operatorname{tr}(\log(\hat{K}) - \log(\hat{P}_{\ell}))}_{\approx \operatorname{stochastic trace estimate}}$$

The better the preconditioner, the smaller the stochastic approximation \Rightarrow variance reduction



- Backward pass analogously via automatic differentiation.
- If we compute a preconditioner for CG, we can simply reuse it at negligible overhead.
- ► If $\hat{P}_{\ell} \rightarrow \hat{K}$ at rate $g(\ell)$, then the STE only requires $\mathcal{O}(\ell^{-\frac{1}{2}}g(\ell))$ random vectors.



Convergence Rates for Kernel – Preconditioner Combinations



Fhe faster the preconditioner converges to the kernel matrix (i.e. $g(\ell) o 0)$ the fewer random vectors are needed.

If $\hat{P}_{\ell} \to \hat{K}$ at rate $g(\ell)$, then the STE only requires $\mathcal{O}(\ell^{-\frac{1}{2}}g(\ell))$ random vectors.

Kernel	d	Preconditioner	$g(\ell)$	Condition
any	\mathbb{N}	none	1	
any	\mathbb{N}	truncated SVD	$\ell^{-rac{1}{2}}$	
any	\mathbb{N}	random. SVD	$\ell^{-rac{1}{2}} + \mathcal{O}(\ell^{rac{1}{4}} \mathtt{S}^{-rac{1}{4}})$	w/ high prob. for <i>s</i> samples
any	\mathbb{N}	random. Nyström	$\ell^{-\frac{1}{2}} + \mathcal{O}(\ell^{\frac{1}{4}}S^{-\frac{1}{4}})$	w/ high prob. for <i>s</i> samples
any	\mathbb{N}	RFF	$\ell^{-\frac{1}{2}}$	w/ high prob.
RBF	1	partial Cholesky	$\exp(-c\ell)$	for some $c > 0$
RBF	\mathbb{N}	QFF	$\exp(-b\ell^{\frac{1}{d}})$	for some $b>0$ if $\ell^{rac{1}{d}}>2\gamma^{-2}$
Matérn (ν)	\mathbb{N}	partial Cholesky	$\ell^{-(\frac{2\nu}{d}+1)}$	$2 u\in\mathbb{N}$, maximin ordering Schaefer2021a
$Matérn(\nu)$	1	QFF	$\ell^{-(s(\nu)+1)}$	where $s(\nu) \in \mathbb{N}$
mod. Matérn (ν)	\mathbb{N}	QFF	$\ell^{-\frac{s(\nu)+1}{d}}$	where $s(u) \in \mathbb{N}$
additive	\mathbb{N}	any	$dg(\ell)$	all summands have rate $g(\ell)$
any	\mathbb{N}	any kernel approx.	$g(\ell)$	∃ uniform convergence bound

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Theoretical Guarantees

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Probabilistic error bounds for the estimates of the log-marginal likelihood and its derivative.

Theorem (Log-marginal likelihood)

[...] Then with probability $1 - \delta$, the error in the estimate η of the log-marginal likelihood $\mathcal L$ satisfies

$$|\eta - \mathcal{L}| \le \varepsilon_{\text{CG}} + \frac{1}{2}(\varepsilon_{\text{Lanczos}} + \varepsilon_{\text{STE}}) ||\log(\hat{K})||_{\text{F}},$$

where the individual errors are bounded by

$$\varepsilon_{CG}(\kappa, i) \le K_3 \left(\frac{\sqrt{\kappa}-1}{\sqrt{\kappa}+1}\right)^i$$
 (3)

$$\varepsilon_{\text{Lanczos}}(\kappa, i) \le K_1 \left(\frac{\sqrt{2\kappa+1}-1}{\sqrt{2\kappa+1}+1}\right)^{2i}$$
 (4)

$$\varepsilon_{\text{STE}}(\delta, \ell) \le C_1 \sqrt{\log(\delta^{-1})} \ell^{-\frac{1}{2}} g(\ell)$$
 (5)

Theorem (Derivative)

[...] Then with probability $1 - \delta$, the error in the estimate ϕ of the derivative of the log-marginal likelihood $\frac{\partial}{\partial \theta} \mathcal{L}$ satisfies

$$|\phi - \tfrac{\partial}{\partial \theta}\mathcal{L}| \leq \varepsilon_{\mathrm{CG}} + \tfrac{1}{2}(\varepsilon_{\mathrm{CG}'} + \varepsilon_{\mathrm{STE}}) \|\hat{K}^{-1} \tfrac{\partial \hat{K}}{\partial \theta}\|_{\mathrm{F}}$$

where the individual errors are bounded by

$$\varepsilon_{CG}(\kappa, i) \le K_4 \left(\frac{\sqrt{\kappa}-1}{\sqrt{\kappa}+1}\right)^i$$
 (6)

$$\varepsilon_{\mathbb{CG}'}(\kappa,i) \le K_2 \left(\frac{\sqrt{\kappa}-1}{\sqrt{\kappa}+1}\right)^i$$
 (7)

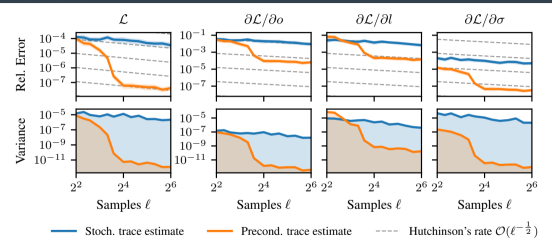
$$\varepsilon_{\text{STE}}(\delta,\ell) \le C_1 \sqrt{\log(\delta^{-1})} \ell^{-\frac{1}{2}} g(\ell)$$
(8)

We leverage preconditioning not only to reduce bias, but crucially also to reduce variance. Computation-Aware Gaussian Process Inference – Jonathan Wenger – July 19, 2023

Preconditioning Reduces Bias and Variance

Estimating the log-marginal likelihood and its derivatives <u>on synthetic data.</u>





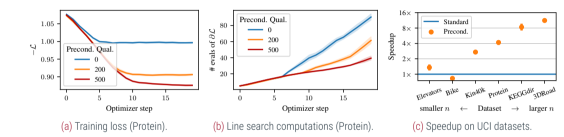
Experiment Details:

Randomly sampled synthetic data (n = 10,000, d = 1)

Computation-AvRBFGkesnelpwithsnoisecsCalerrathar=wellg0r=2July 19, 2023

Preconditioning Accelerates Hyperparameter Optimization

Gaussian process hyperparameter optimization on UCI data.



Experiment Details:

- UCI datasets (n = 12,449 to n = 326,155)
- Matérn $(\frac{3}{2})$ kernel with noise scale $\sigma^2 = 10^{-2}$
- Partial Cholesky preconditioner of size 500
- l = 50 random vectors

TUBINGE

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