
Supplementary Material for Large Linear Multi-output Gaussian Process Learning for Time Series

Anonymous Author(s)

Affiliation

Address

email

1 Implementation Details

LLGP was implemented in Python 3 from the Anaconda, which offered an Intel MKL-linked scipy [1]. The code made heavy use of other packages, namely climin [2], GPy [3], and paramz [4]. Code and benchmarks are available at <anonymous repository>.

Application of our approach to all replication studies was carried out on a large server in a multi-programming environment: CentOS 6.5 with 80 Intel(R) Xeon(R) CPU E7-4870 @ 2.40GHz. The representation evaluation benchmarks were done at once on a cluster of machines running CentOS 5.2-5.9 with Intel(R) Xeon(R) CPU E5430 @ 2.66GHz, where these jobs ran on a single thread per CPU.

2 Representation evaluation

We evaluated the performance of the different kernel representations over various rank and parameterization settings. In particular, we have the following parameters: n total sample size across all outputs, D number of outputs, Q number of kernels k_q , R average added rank, ϵ mean noise, and $ktype$ kernel type. Kernel type is one of `mat`, `periodic`, `rbf`, `mix` corresponding to Matérn-3/2, standard periodic¹, and radial basis functions. `mix` refers to a mix of all three kernels.

For each setting, we randomly sample entries for each A_q , κ_q , and ϵ and the inverse length scale γ for each kernel. Then, we investigate the average gradient construction accuracy and speed of LLGP for different settings of Q , ϵ , and $ktype$. Each kernel's inverse length scales and periods were selected by sampling uniformly in log space from 1 to 10 with Q samples. Next, we construct a random LMC kernel by sampling entries of each A_q from a standard normal distribution truncated to the unit interval, κ_q from an inverse gamma with unit shape and scale, and independent noise ϵ for every output from an inverse gamma with unit scale and mean ϵ . Inputs and outputs were independently generated from $\text{Unif}[0, 1]$ for benchmarking.

As expected from their asymptotic runtime, SUM, BT, and SLFM representations are complimentary in MVM speed for different configurations of D , R , Q —this results in sparse inversion computation that consistently outperforms Cholesky decomposition in runtime (Tab. 1). For inverting systems, all computations were carried out until the residual ℓ_2 norm was at most 10^{-4} .

We next evaluated the accuracy of the gradients for $N = 10$ trace samples. Fixing $R = 3$, $n = 5000$, $D = 10$, we quantified the accuracy LLGP's $\nabla \mathcal{L}$ by comparing against the exact Cholesky approach. The relative error in the gradients is generally low for smooth kernels that induce diagonally dominant covariance matrices (Fig. 1). Kernels such as the single Matérn or periodic kernel with noise on the order of 10^{-4} lead to less accurate gradients, owing to poor MINRES convergence in the

¹We define the periodic kernel as $k(r) = \exp\left(\frac{-\gamma}{2} \sin^2 \frac{\pi r}{T}\right)$.

Table 1: The runtime in seconds for solving $K\mathbf{x} = \mathbf{y}$ for a random kernel K constructed as in Sec. 2 using MINRES for each of the kernel representations. For comparison, the CHOL representation is wallclock time to compute the Cholesky decomposition of the matrix, which must be constructed, and use this decomposition to invert the system. We averaged over five runs. In every run, we use $n = 5000$ simulated data points, mix kernels, and $\epsilon = 0.1$.

| D | R | Q | CHOLSKY | SUM | BT | SLFM |
|-----|-----|-----|---------|-------------|-------------|-------------|
| 2 | 2 | 10 | 41.00 | 32.24 | 6.53 | 42.33 |
| 10 | 1 | 10 | 37.38 | 31.01 | 20.81 | 9.34 |
| 10 | 10 | 1 | 10.74 | 0.41 | 2.40 | 0.89 |

33 inversions (Fig. 2. We showed in the main paper that the stochastic gradients suffice for optimization
34 in practical examples.

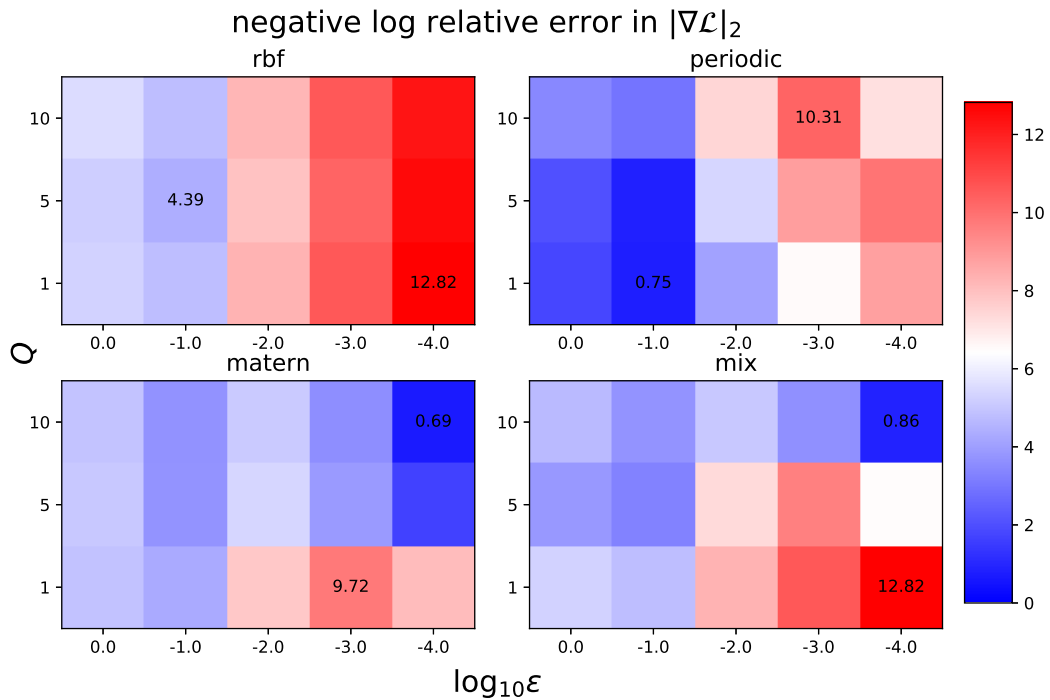


Figure 1: Negative logarithm of the relative error from the LLGP gradient construction to the exact log likelihood gradient. In each case, higher is better, and extremal values are noted. For each data point, the average was taken over five runs. Reducing average noise, corresponding to increases in $-\log_{10}\epsilon$, and increases in Q , generally make gradient reconstruction more difficult by making K more ill-conditioned, reducing the accuracy of MINRES.

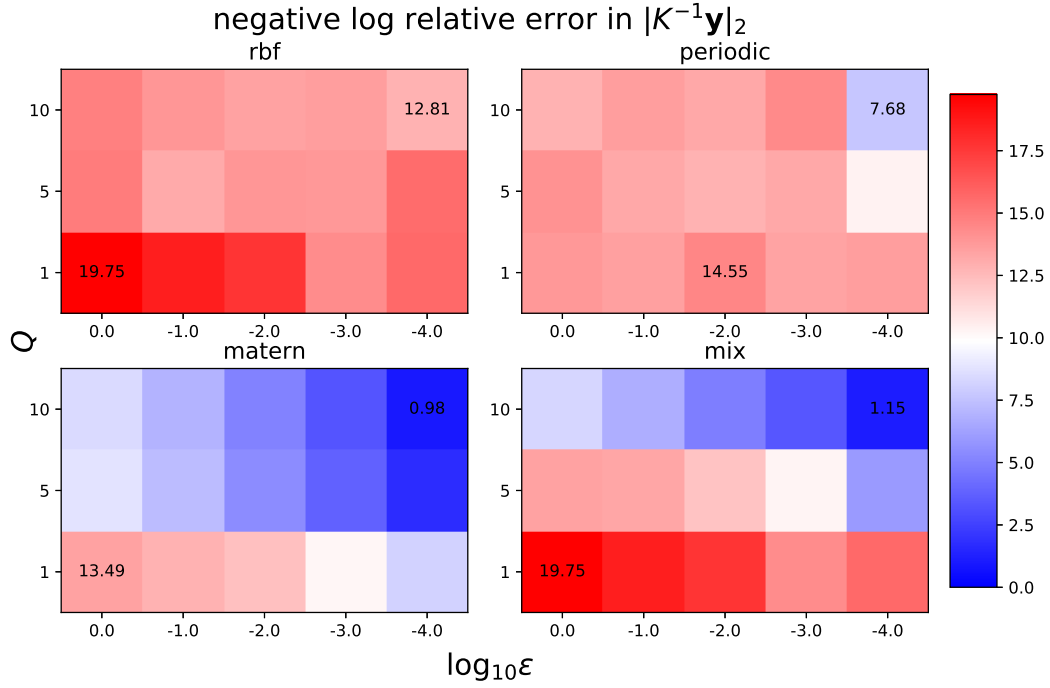


Figure 2: Negative logarithm of the relative error in $K^{-1}\mathbf{y}$, using MINRES compared to the Cholesky solution. Higher is better, and extremal values are noted. An average was taken over five runs.

References

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