Exercise 07

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1 Probabilistic Machine Learning

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1.1 Exercise Sheet No. 7 — Online Gaussian Process Regression

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```
[]: # For measuring wallclock time
from time import time

# Numpy, JAX
import jax
import jax.numpy as jnp
import numpy as np
jax.config.update("jax_enable_x64", True) # use double-precision numbers
```

```
[]: # Plotting
from matplotlib import pyplot as plt
from tueplots import bundles

import matplotlib as mpl

%config InlineBackend.figure_formats = ["svg"]
%matplotlib inline
plt.rcParams.update(bundles.beamer_moml())
plt.rcParams.update({'figure.dpi': 200})
```

```
[]: # Import `gaussians.py`
import gaussians
```

1.1.1 0. Overview

In this exercise sheet, we are going to implement Gaussian Process (GP) regression in an *online* setting. That means, data arrives sequentially over time and the GP regression model is incrementally updated to incorporate that new information.

In the lecture and previous exercise sheets, you have already seen and worked with the classes Gaussian, GaussianProcess, and ConditionalGaussianProcess implemented in gaussians.py. This functionality is also very straightforward to use in the online setting, but a naive approach will turn out to be prohibitively inefficient. We are going to investigate the underlying problem in more detail and then develop and implement a more efficient strategy by using some tools from linear algebra (like the Schur complement and a block-wise Cholesy factorization).

1.1.2 1. Training and Test Data

Since we want to simulate that the training data is provided *incrementally*, it is represented by two lists X_list (containing 6 subsets of 100 data points each) and Y_list (containing the corresponding regression targets). As a first step, we load the data and visualize these subsets.

```
[]: # Load data
data = np.load("data.npz")

# Extract from data
X_list = data["X_list"]
Y_list = data["Y_list"]
sigma = data["sigma"]
X_test = data["X_test"]
[]: # Check shape of sections
print("Training data")
```

```
[]: # Check shape of sections
print("Training data")
for idx, (Xi, Yi) in enumerate(zip(X_list, Y_list)):
    print(f"Subset index {idx}: X.shape = {Xi.shape}, Y.shape = {Yi.shape}")

print("\nTest data")
print("X_test.shape = ", X_test.shape)
```

```
print("\nTest data")
print("X_test.shape = ", X_test.shape)

Training data
Subset index 0: X.shape = (100, 1), Y.shape = (100,)
Subset index 1: X.shape = (100, 1), Y.shape = (100,)
Subset index 2: X.shape = (100, 1), Y.shape = (100,)
Subset index 3: X.shape = (100, 1), Y.shape = (100,)
Subset index 4: X.shape = (100, 1), Y.shape = (100,)
Subset index 5: X.shape = (100, 1), Y.shape = (100,)
Test data
X_test.shape = (100, 1)

[]: def setup_plot(ax, title=None):
    ax.set_xlabel("$x$")
```

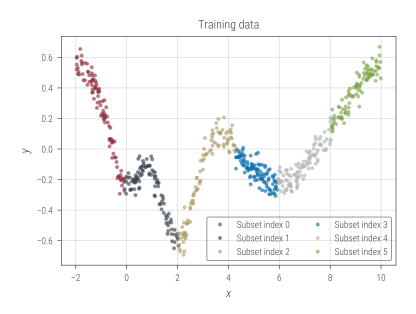
```
ax.set_ylabel("$y$")
ax.grid()
if title is not None:
    ax.set_title(title)

# Visualization of training data
fig, ax = plt.subplots(1, 1, figsize=(4, 3))

for idx, (Xi, Yi) in enumerate(zip(X_list, Y_list)):
    ax.plot(Xi, Yi, "o", ms=2, alpha=0.5, label=f"Subset index {idx}")
print(Xi.shape)

setup_plot(ax, "Training data")
ax.legend(ncol=2)
plt.show()
```

(100, 1)



1.1.3 2. Prior GP

Here, we set up the prior GP with a constant mean at zero and rational quadratic kernel as covariance function.

```
[]: import functools

def constant_mean(x, c=0.0):
```

```
[]: # Define mean, kernel and prior GP
mean = functools.partial(constant_mean, c=0.0)
kernel = functools.partial(RQ_kernel, ell=1.0, alpha=1.0, theta=0.5)
prior = gaussians.GaussianProcess(mean, kernel)
```

1.1.4 3. Approach 1

We can use the functionality implemented in gaussians.py to perform inference in an incremental way: In the first step, we condition the prior on the first subset (X_list[0], y_list[0]). From this, we obtain an instance of a ConditionalGaussianProcess that we can condition on the next subset (X_list[1], y_list[1]) etc.

```
[]: def get_posteriors():
    """Construct posteriors from prior by conditioning on subsets"""

posteriors = [prior]
    for posterior_idx, (Xi, Yi) in enumerate(zip(X_list, Y_list)):

    # Compute posterior
    posterior = posteriors[-1].condition(Yi, Xi, sigma)
    posteriors.append(posterior)
    posteriors.pop(0) # remove prior from list
    return posteriors

posteriors_A1 = get_posteriors()
```

No GPU/TPU found, falling back to CPU. (Set TF_CPP_MIN_LOG_LEVEL=0 and rerun for more info.)

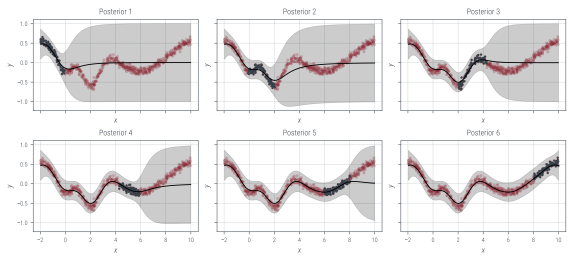
Let's visualize those posteriors.

```
[]: from math import ceil

def get_nrows_ncols(nplots):
    nrows = ceil((1/3) * nplots)
    ncols = ceil(nplots / nrows)
    return nrows, ncols

def plot_conditional_gps(list_of_conditional_gps):
```

```
"""Plot a list of conditional GPs in one figure."""
    # Concatenate training data
    X = jnp.concatenate(X_list, axis=0)
    Y = jnp.concatenate(Y_list, axis=0)
    # Define grid of subplots
    num_gps = len(list_of_conditional_gps)
    nrows, ncols = get_nrows_ncols(num_gps)
    fig, axs = plt.subplots(
        nrows,
        ncols,
        figsize=(ncols * 3, nrows * 2),
        sharex=True,
        sharey=True,
    )
    # Plot GPs
    for idx, gp in enumerate(list_of_conditional_gps):
        ax = axs.flatten()[idx]
        # Plot GP and data
        ax.plot(X, Y, ".", ms=5, alpha=0.2)
        ax.plot(gp.X, gp.y, ".", ms=3)
        gp.plot(ax, X_test, color="k", std_kwargs={"alpha": 0.2})
        setup_plot(ax, f"Posterior {idx + 1}")
    plt.show()
plot_conditional_gps(posteriors_A1)
```



Hm, that took quite some time. Let's investigate the runtime in more detail.

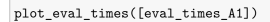
```
def measure_eval_times(nreps=3):
    """Construct posteriors and measure the time required for evaluating
    them on `X_test`"""

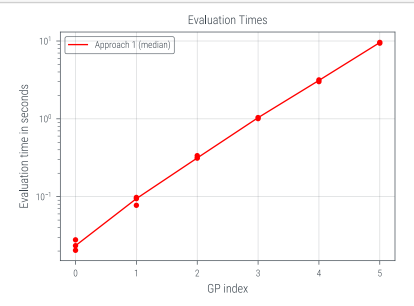
posteriors = [prior]
    eval_times = np.zeros((nreps, len(X_list)))
    for posterior_idx, (Xi, Yi) in enumerate(zip(X_list, Y_list)):
        for rep_idx in range(nreps):

        # Measure the time for evaluating the posterior on `X_test`
        start_time = time()
        posterior = posteriors[-1].condition(Yi, Xi, sigma)
        posterior(X_test)
        eval_times[rep_idx, posterior_idx] = time() - start_time
        posteriors.append(posterior)

return eval_times
eval_times()
```

```
[]: import matplotlib.colors as mcolors
     COLS = ["red", "green", "blue"]
     def plot_eval_times(list_of_eval_times):
         """Plot the measured evaluation times. Entry 0 in `list_of_eval_times`
         is assumed to belong to approach 1, entry 1 to approach 2 etc.
         fig, ax = plt.subplots(1, 1, figsize=(4, 3))
         for idx, eval_times in enumerate(list_of_eval_times):
             ax.plot(eval_times.T, "o", ms=3, c=COLS[idx])
             label = f"Approach {idx + 1} (median)"
             ax.plot(np.median(eval times, axis=0), "-", c=COLS[idx], label=label)
         # General plot settings
         ax.set_xlabel("GP index")
         ax.set_ylabel("Evaluation time in seconds")
         ax.grid()
         ax.legend()
         ax.set_yscale("log")
         ax.set_title(f"Evaluation Times")
         plt.show()
```





```
[]: # Determine the slopes of the log evaluation time
  eval_times_A1_median = np.median(eval_times_A1, axis=0)
  slopes = eval_times_A1_median[1:] / eval_times_A1_median[:5]
  print("slopes = ", slopes)
```

slopes = [4.02063229 3.32992997 3.29469608 3.01359783 3.06525569]

Observations: The logarithmic plot shows that the runtime increases approximately exponentially reaching up to 10 seconds (on your machine, it may be different of course) for the last posterior. The runtime increases by a factor of around 3-4 for each additional subset of data.

Task: What's going on here? Why is the runtime increasing approximately exponentially?

Your answer: As the data size that the Conditional Gaussian conditioned on increases, the cost of operations (particularly matrix inversions) applied on the current (say N sized) matrix increases as well. Nearly exponential increase is due to matrix inversions (kernel size in Cholesky).

1.1.5 4. Approach 2

Here, we condition the prior on a growing data set. For this, we overwrite the condition method of the ConditionalGaussianProcess class.

```
[]: from gaussians import Gaussian, ConditionalGaussianProcess

def condition_on_prior(self, y_new, X_new, sigma):

# Check data
```

```
y_new = jnp.atleast_1d(y_new)
   X_new = jnp.atleast_2d(X_new)
    # Concatenate data
   X_both = jnp.concatenate([self.X, X_new], axis=0)
   y_both = jnp.concatenate([self.y, y_new], axis=0)
    # Concatenate Gaussians
   mu new = jnp.zeros like(y new)
   mu_both = jnp.concatenate([self.epsilon.mu, mu_new], axis=0)
   Sigma_new = sigma * jnp.eye(len(y_new))
   Sigma_both = jax.scipy.linalg.block_diag(self.epsilon.Sigma, Sigma_new)
   epsilon_both = Gaussian(mu=mu_both, Sigma=Sigma_both)
    # Build conditional GP from prior (!)
   return ConditionalGaussianProcess(self.prior, y=y_both, X=X_both,__
 ⇔epsilon=epsilon_both)
# Assign method to class
ConditionalGaussianProcess.condition = condition_on_prior
```

```
[]: # Evaluate approach 2
posteriors_A2 = get_posteriors()
eval_times_A2 = measure_eval_times()
```

Note: In theory, you can show that the posteriors from approaches 1 and 2 are identical, i.e. conditioning multiple times in a convoluted manner (approach 1) is equivalent to conditioning on the union of the subsets (approach 2). Let's check if we can confirm this claim empirically.

Task: Evaluate the posteriors on X_test and check if the resulting Gaussian distributions are identical (jnp.allclose might be useful here).

```
# TODO

# Check if the resulting Gaussian distributions are identical (`jnp.allclose`u might be useful here)

for posterior_A1, posterior_A2 in zip(posteriors_A1, posteriors_A2):
    same_mu = jnp.allclose(posterior_A1(X_test).mu, posterior_A2(X_test).mu)
    same_Sigma = jnp.allclose(posterior_A1(X_test).Sigma, posterior_A2(X_test).

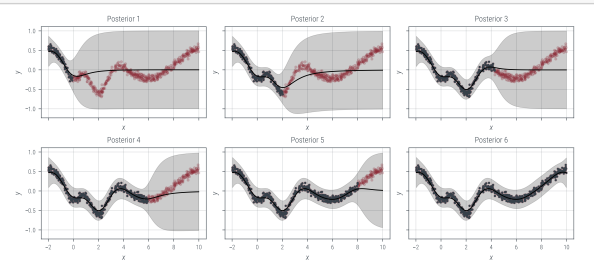
Sigma)

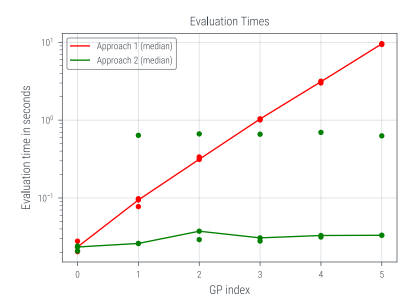
if not same_mu or not same_Sigma:
    print("The posteriors are not identical.")
    break

if same_mu and same_Sigma:
    print("The posteriors are identical.")
```

The posteriors are identical.

[]: # Visualizations plot_conditional_gps(posteriors_A2) plot_eval_times([eval_times_A1, eval_times_A2])





Even though the slope is less in the plot of approach 2 in logarithmic scale, the (almost) linear trend in the logarithmic scale is present in the second approach too. Empirically, there is a little discrepancy yet the general trend is the same.

1.1.6 5. Approach 3

While approach 2 is less inefficient, it is still quite wasteful: For every new subset, the posterior GP is built entirely from scratch! This can be improved by exchanging information between the

posteriors: Simply put, a posterior that is conditioned on some data set should inform the next posterior what it has already learned about the data. Before we dive into that, we need to get the theory straight.

Our goal is to condition the GP on a data set (X,y) that is the union of (X_{old},y_{old}) and (X_{new},y_{new}) . Let's assume that we have conditioned a GP on (X_{old},y_{old}) and have already triggered the computation of its properties. So, in particular, the following quantities are already cached:

- 1. predictive_covariance: $k(X_{old}, X_{old}) + \sigma^2 I$
- 2. predictive_mean: $m(X_{old}) + \mu(X_{old})$
- 3. predictive_covariance_cho: the Cholesky factorization of predictive_covariance
- 4. representer_weights: $[k(X_{old}, X_{old}) + \sigma^2 I]^{-1}(y_{old} m(X_{old}) + \mu(X_{old}))$

Improved version: Now, the core idea is to use the quantities above to save computations when conditioning on (X, y).

5. predictive_covariance: This matrix now has the following block structure:

$$k(X,X) + \sigma^2 I = \frac{k(X_{old}, X_{old}) + \sigma^2 I}{k(X_{new}, X_{old})} \frac{k(X_{old}, X_{new})}{k(X_{new}, X_{new}) + \sigma^2 I} =: \frac{A \mid B}{C \mid D}.$$

If we have already conditioned on (X_{old}, X_{old}) , block A was already computed (see 1. above) and we only have to evaluate the remaining blocks B, C and D.

6. predictive_mean: The predictive mean also splits into two parts

$$m(X) + \mu(X) = \left(\frac{m(X_{old}) + \mu(X_{old})}{m(X_{new}) + \mu(X_{new})}\right) =: \left(\frac{a}{b}\right).$$

Note that a was already computed, see 2. above.

- 7. predictive_covariance_cho: In order to compute a Cholesky factorization of $k(X, X) + \sigma^2 I$, we can use that we already have the Cholesky factor $A^{1/2}$ of A, see 3. above. It should be clear from theory exercise 1 (b) on the current exercise sheet how to construct the Cholesky factor for $k(X, X) + \sigma^2 I$.
- 8. representer_weights: We already computed the representer weights $A^{-1} \cdot u$, with $u = y_{old} a$, see 4. above. Now, we have to solve a larger linear system

$$\begin{pmatrix} A & B \\ C & D \end{pmatrix} \cdot \begin{pmatrix} x \\ y \end{pmatrix} = \begin{pmatrix} u \\ v \end{pmatrix}$$

with $v = y_{new} - b$ and the unknowns x and y. After computing v, y and x are given by

$$y = Q^{-1}(v - CA^{-1}u) \tag{1}$$

$$x = A^{-1}u - A^{-1}By, (2)$$

where $Q=D-CA^{-1}B$ is the so-called Schur-complement. Note: All necessary quantities (like $Q^{1/2}$ and $A^{-1}u$) were already computed in previous steps.

Task: Implement an improved version of the condition method that constructs 5, 6, 7, and 8. Try to make this as efficient as possible by using the quantities 1, 2, 3, and 4. Make sure that the resulting posteriors are the same as the ones from approach 2 (e.g. by evaluating them on the same test data and comparing the Gaussians).

```
[]: from gaussians import Gaussian, ConditionalGaussianProcess
    def condition_on_prior_improved(self, y_new, X_new, sigma):
        # Check data
        y_new = jnp.atleast_1d(y_new)
        X_new = jnp.atleast_2d(X_new)
        # Concatenate data
        X_both = jnp.concatenate([self.X, X_new], axis=0)
        y_both = jnp.concatenate([self.y, y_new], axis=0)
        # Concatenate Gaussians
        mu_new = jnp.zeros_like(y_new)
        mu_both = jnp.concatenate([self.epsilon.mu, mu_new], axis=0)
        Sigma_new = sigma * jnp.eye(len(y_new))
        Sigma_both = jax.scipy.linalg.block_diag(self.epsilon.Sigma, Sigma new)
        epsilon_both = Gaussian(mu=mu_both, Sigma=Sigma_both)
        # Build conditional GP from prior (!)
        cond_gp = ConditionalGaussianProcess(self.prior, y=y_both, X=X_both,_
     ⇔epsilon=epsilon_both)
        # -----
        # 5. predictive_covariance
        # -----
        # TODO
        global A
        A = self.predictive_covariance
        B = self.prior.k(self.X[:, None, :], X_new[None, :, :])
        global D
        D = self.prior.k(X_new[:, None, :], X_new[None, :, :]) + Sigma_new
        def predictive_covariance_new(A, B, C, D):
           # Append A, B, C, D to a single square matrix of shape (n + m, n + m)_{\sqcup}
     \hookrightarrow and return
           return jnp.block([[A, B], [C, D]])
        cond_gp.predictive_covariance = predictive_covariance_new(A, B, C, D)
        # -----
        # 6. predictive_mean
        # -----
```

```
# TODO
a = self.predictive_mean
b = self.prior.m(X_new) + mu_new
def predictive_mean_new(a, b):
   # Append a, b and return
   return jnp.concatenate([a, b], axis=0)
cond_gp.predictive_mean = predictive_mean_new(a, b)
# 7. predictive_covariance_cho
global A_1_2
A_1_2, lower_A = self.predictive_covariance_cho
if not lower_A:
    # print('A is not lower triangular')
   A_1_2 = A_1_2.T
   lower_A = True
# assert jnp.allclose(A_1_2 @ A_1_2.T, A)
# if not jnp.allclose(A_1_2 @ A_1_2.T, A):
     A \ 1 \ 2 = A \ 1 \ 2.T
# elif not jnp.allclose(A_1_2.T @ A_1_2, A):
     A \ 1 \ 2 = A \ 1 \ 2.T
U_12 = jax.scipy.linalg.solve_triangular(A_1_2, B, lower=True)
L 21 = U 12.T
global Q
global Q_1_2
A_inverse_B = jax.scipy.linalg.cho_solve((A_1_2, lower_A), B)
Q = D - C @ A_inverse_B
Q_1_2, lower_Q = jax.scipy.linalg.cho_factor(Q, lower=True)
if not lower_Q:
    # print('Q is not lower triangular')
   Q_1_2 = Q_1_2.T
   lower_Q = True
# assert jnp.allclose(Q_1_2 @ Q_1_2.T, Q)
def predictive_covariance_cho_alt(cond_gp):
   return jax.scipy.linalg.cho_factor(cond_gp.predictive_covariance)
def predictive_covariance_cho_new(A_1_2, U_12, L_21, Q_1_2):
   \#U = jnp.block([[A_1_2.T, U_12], [jnp.zeros_like(L_21), Q_1_2.T]])
   L = jnp.block([[A_1_2, jnp.zeros_like(U_12)], [L_21, Q_1_2]])
   return (L, True)
```

```
#cond_qp.predictive_covariance_cho = predictive_covariance_cho_alt(cond_qp)
   cond_gp.predictive_covariance_cho = predictive_covariance_cho_new(A_1_2,__
 \cup U_12, L_21, Q_1_2)
   # -----
   # 8. representer weights
   # -----
   # TODO
   u = self.y - a
   v = y_new - b
   repr_w_old = self.representer_weights
   y_vec = jax.scipy.linalg.cho_solve((Q_1_2, lower_Q), (v-C @ repr_w_old))
   x_vec = repr_w_old - A_inverse_B @ y_vec
   def representer_weights_new(x_vec,y_vec):
       return jnp.concatenate([x_vec, y_vec], axis=0)
   def representer_weights_alt():
       return jax.scipy.linalg.cho_solve(cond_gp.predictive_covariance_cho,_
 →jnp.concatenate([u, v], axis=0))
   #cond_gp.representer_weights = representer_weights_alt()
   cond_gp.representer_weights = representer_weights new(x_vec,y_vec)
   return cond_gp
# Assign method to class
ConditionalGaussianProcess.condition = condition_on_prior_improved
# Evaluate approach 3
posteriors_A3 = get_posteriors()
eval_times_A3 = measure_eval_times()
# Make sure that the posteriors are identical to the ones from approach 2
# TODO
for posterior_A2, posterior_A3 in zip(posteriors_A2, posteriors_A3):
   same_mu = jnp.allclose(posterior_A2(X_test).mu, posterior_A3(X_test).mu)
   same_Sigma = jnp.allclose(posterior_A2(X_test).Sigma, posterior_A3(X_test).

Sigma)
   if not same mu or not same Sigma:
       print("The posteriors are not identical.")
       break
if same_mu and same_Sigma:
   print("The posteriors are identical.")
# Visualizations
plot_conditional_gps(posteriors_A3)
```

plot_eval_times([eval_times_A1, eval_times_A2, eval_times_A3])

The posteriors are identical.

