```
In [82]: # Initialize task.
         import jax.numpy as jnp
         import numpy as np
         import matplotlib.pyplot as plt
         import seaborn as snb
         import matplotlib.colors as colors
         from scipy.stats import norm
         from scipy.optimize import minimize
         from jax import hessian, random
         snb.set_theme(font_scale=1.25)
         # Load Data
         data = jnp.load('./data_exercise5b.npz')
         X = data['day']
         y = np.log(data['bike_count'])
         #Standardize data
         ym, ys = jnp.mean(y), jnp.std(y)
         y = (y - ym) / ys
```

# Part 1: Fully Bayesian inference for Gaussian process regression

## Task 1.1 Choose a value for v such that the prior probability of observing a lengthscale larger than 100 is approximately 1%

First, we identify that our lengthscale parameter follows af half normal distribution

$$\ell \sim \mathcal{N}_+(0,v)$$

We know the condition we need to fulfill is

$$P(\ell > 100) \approx 0.01$$

We can express our distribution  $\ell=|Z\sqrt{v}|$  with Z being a standard normal variable, since  $\ell$  is a half normal. Since the standard normal distribution is symmetric we view it as a two-tailed probability.

$$P(\ell > 100) = P(|Z\sqrt{v}| > 100) \approx 0.01$$
  $P(|Z| > \frac{100}{\sqrt{v}}) \approx 0.01$  (Two-tailed)  $2 \cdot P(\frac{Z > 100}{\sqrt{v}}) \approx 0.01$   $P(\frac{Z > 100}{\sqrt{v}}) \approx 0.005$   $P(\frac{Z > 100}{\sqrt{v}}) \approx 0.005$   $P(\frac{Z > 100}{\sqrt{v}}) \approx 0.005$ 

So when  $v \approx 1507$  the probability of the lengthscale parameter being above 100 will be roughly 1%

### Task 1.2 Determine the marginalized distribution $p(y, \sigma, k, \ell)$

The joint distribution is as follows

$$p(\boldsymbol{y}, f, \sigma, \kappa, \ell) = p(\boldsymbol{y} \mid \boldsymbol{f}, \sigma^2) p(f \mid \kappa, \ell) p(\kappa) p(\ell) p(\sigma)$$
  
=  $\mathcal{N}(\boldsymbol{y} \mid \boldsymbol{f}, \sigma^2 \boldsymbol{I}) \mathcal{N}(\boldsymbol{f} \mid \boldsymbol{0}, \boldsymbol{K}) \mathcal{N}_{+}(\kappa \mid 0, 1) \mathcal{N}_{+}(\ell \mid 0, v) \mathcal{N}_{+}(\sigma \mid 0, 1).$ 

Denoting  $\mathcal{N}_+(\kappa \mid 0, 1)\mathcal{N}_+(\ell \mid 0, v)\mathcal{N}_+(\sigma \mid 0, 1) = p(\kappa, \ell, \sigma)$  we get the following for marginalising out f from the joint distribution:

$$p(oldsymbol{y}, \sigma, \kappa, \ell) = \int p(oldsymbol{y}, f, \sigma, \kappa, \ell) \, \mathrm{d}f$$

$$= p(\kappa, \ell, \sigma) \underbrace{\int \mathcal{N}\left(oldsymbol{y} \mid oldsymbol{f}, \sigma^2 oldsymbol{I}
ight) \mathcal{N}(oldsymbol{f} \mid oldsymbol{0}, oldsymbol{K}) \, \mathrm{d}f}_{ ext{Marginal likelihood = linear Gaussian system}}$$

$$= p(\kappa, \ell, \sigma) \mathcal{N}(oldsymbol{y} \mid oldsymbol{0}, oldsymbol{K} + \sigma^2 oldsymbol{I})$$

#### Task 1.3: Implement a Metropolis sampler using the proposal distribution

```
In [83]: def metropolis(log_target, num_params, tau, num_iter, theta_init=None, seed=0):
             Runs a Metropolis-Hastings sampler.
             Arguments:
                 log_target: function evaluating the log target, expecting a vector (size
                 num_params: number of parameters
                           vector of proposal standard deviations (one per parameter)
                 num iter: number of iterations
                 theta init: initial parameter vector (or None)
                         random seed
                 seed:
             Returns:
                 thetas: jnp.array of shape (num_iter+1, num_params)
                 accepts: list of acceptance flags
             key = random.PRNGKey(seed)
             if theta init is None:
                 theta_init = jnp.zeros((num_params))
             thetas = [theta_init]
             accepts = []
             log_p_theta = log_target(theta_init)
             for k in range(num iter):
                 key, key_proposal, key_accept = random.split(key, num=3)
                 theta_cur = thetas[-1]
                 # Elementwise proposal using tau for each parameter.
                 theta star = theta cur + tau * random.normal(key proposal, shape=(num pa
                 log_p_theta_star = log_target(theta_star)
                 log r = log p theta star - log p theta
                 A = \min(1, jnp.exp(log_r))
                 if random.uniform(key_accept) < A:</pre>
```

```
theta_next = theta_star
    log_p_theta = log_p_theta_star
    accepts.append(1)
else:
    theta_next = theta_cur
    accepts.append(0)
    thetas.append(theta_next)

print("Acceptance ratio: %3.2f" % jnp.mean(jnp.array(accepts)))
thetas = jnp.stack(thetas)
assert thetas.shape == (
    num_iter + 1,
    num_params,
), f"Expected shape {(num_iter+1, num_params)}, got {thetas.shape}."
return thetas, accepts
```

```
In [ ]: def metropolis_multiple_chains(
            log_target, num_params, num_chains, tau, num_iter, theta_init, seeds, warm_u
        ):
            """Runs multiple Metropolis-Hastings chains. The i'th chain should be initia
            Arguments:
                                    function for evaluating the log joint distribution
                log_target:
                                     number of parameters of the joint distribution (inte
                num_params:
                                     number of MCMC chains
                num_chains:
                tau:
                                     proposal standard deviation (jnp.array with shape (n
                num_iter:
                                    number of iterations for each chain (integer)
                                    array of initial values (jnp.array with shape (num_c
                theta_init:
                                     seed for each chain (jnp.array with shape (num_chain
                seeds:
                                    number of warm up samples to be discarded
                warm_up:
            returns:
                thetas
                                     jnp.array of samples from each chain after warmup (s
                                    jnp.array of acceptances rate for each chain (shapes
                accept_rates
            0.00
            # verify dimension of initial parameters
            assert theta_init.shape == (
                num chains,
                num params,
            ), "theta init seems to have the wrong dimensions. Plaese check your code."
            # prepare arrays for storing samples
            thetas = []
            accept_rates = []
            # run sampler for each chain
            for idx_chain in range(num_chains):
                print(f"Running chain {idx_chain}. ", end="")
                thetas temp, accepts temp = metropolis(
                    log_target,
                    num params,
                    tau,
                    num iter,
                    theta_init=theta_init[idx_chain],
                    seed=seeds[idx_chain],
                thetas.append(thetas temp)
                accept_rates.append(jnp.array(accepts_temp))
```

```
thetas = jnp.stack(thetas, axis=0)
accept_rates = jnp.stack(accept_rates, axis=0)

# discard warm-up samples
thetas = thetas[:, warm_up:, :]

# verify dimensions and return
assert thetas.shape == (
    num_chains,
    num_iter + 1 - warm_up,
    num_params,
), f"The expected shape of chains is ({num_chains}, {num_iter+1-warm_up}, {num_sert len(accept_rates) == num_chains
return thetas, accept_rates
```

```
In [ ]: # Log half-normal prior.
        def log_halfnormal(x, scale=1.0):
            return jnp.log(2 / scale) + norm.logpdf(x, 0, scale)
        def squared_exponential(tau, kappa, lengthscale):
            return kappa**2 * jnp.exp(-0.5 * tau**2 / (lengthscale**2))
        class StationaryIsotropicKernel(object):
            def __init__(self, kernel_fun, kappa=1.0, lengthscale=1.0):
                the argument kernel_fun must be a function of three arguments kernel_fun
                squared_exponential = lambda tau, kappa, lengthscale: kappa**2*np.exp(-0
                self.kernel_fun = kernel_fun
                self.kappa = kappa
                self.lengthscale = lengthscale
            def contruct_kernel(self, X1, X2, kappa=None, lengthscale=None, jitter=1e-8)
                """compute and returns the NxM kernel matrix between the two sets of inp
                arguments:
                    X1
                                    -- NxD matrix
                    X2
                                    -- MxD matrix
                    kappa
                                   -- magnitude (positive scalar)
                    lengthscale
                                    -- characteristic lengthscale (positive scalar)
                    jitter
                                    -- non-negative scalar
                returns
                                    -- NxM matrix
                # extract dimensions
                N, M = X1.shape[0], X2.shape[0]
                # prep hyperparameters
                kappa = self.kappa if kappa is None else kappa
                lengthscale = self.lengthscale if lengthscale is None else lengthscale
                # compute all the pairwise distances efficiently
                dists = jnp.sqrt(
                    jnp.sum((jnp.expand_dims(X1, 1) - jnp.expand_dims(X2, 0)) ** 2, axis
```

```
# squared exponential covariance function
        K = self.kernel_fun(dists, kappa, lengthscale)
        # add jitter to diagonal for numerical stability
        if len(X1) == len(X2) and jnp.allclose(X1, X2):
            K = K + jitter * jnp.identity(len(X1))
        assert K.shape == (
           Ν,
           Μ,
        ), f"The shape of K appears wrong. Expected shape ({N}, {M}), but the ac
        return K
# Log joint function: returns -inf if any parameter ≤ 0.
def log_marginal_likelihood(theta, X, y, v=1507.0):
    kappa, ell, sigma = theta
   n = y.shape[0]
   if (kappa <= 0) or (ell <= 0) or (sigma <= 0):</pre>
        return -jnp.inf
    # Priors.
   lp_kappa = log_halfnormal(kappa, scale=1.0)
   lp_ell = log_halfnormal(ell, scale=jnp.sqrt(v))
   lp_sigma = log_halfnormal(sigma, scale=1.0)
    log_prior = lp_kappa + lp_ell + lp_sigma
    # Kernel matrix computation.
   kernel = StationaryIsotropicKernel(squared_exponential, kappa, ell)
    K = kernel.contruct kernel(X, X)
   C = K + sigma**2 * jnp.eye(n)
   # Compute the Cholesky decomposition.
    L = jnp.linalg.cholesky(C)
   v vec = jnp.linalg.solve(L, y)
    # Compute Log marginal likelihood.
    logdet_term = jnp.sum(jnp.log(jnp.diag(L)))
    quad term = 0.5 * jnp.sum(v vec**2)
    const_term = -0.5 * n * jnp.log(2 * jnp.pi)
    log_likelihood = const_term - logdet_term - quad_term
    return log_prior + log_likelihood
def log target(theta):
    """Log target function for the Metropolis-Hastings sampler.
    Arguments:
       theta:
                           jnp.array of parameters (jnp.array with shape (num_pa
    Returns:
                          log target distribution (real number)
       log_target:
    # Compute the log target distribution.
    log_mar_likelihood = log_marginal_likelihood(theta, X, y)
    log_target = log_mar_likelihood
    return log_target
```

```
In [98]:
        # mcmc settings
         num chains = 4
         num_iter = 10000
         tau = jnp.array([1.0, 100.0, 0.1])
         num_params = 3
         warm_up = 0
         seeds = jnp.arange(num chains)
         # generate initial values from uniform distribution
         key = random.PRNGKey(1)
         theta_init = random.uniform(
             key, shape=(num_chains, num_params), minval=0.1, maxval=2.0
         )
         # sample
         chains, accepts = metropolis_multiple_chains(
             log_target,
             num params,
             num_chains,
             tau,
             num_iter,
             theta_init=theta_init,
             seeds=seeds,
             warm_up=warm_up,
         # report estimated mean and variance
         print(f"\nEstimated mean:\t\t{jnp.mean(chains.ravel()):+3.2f}")
         print(f"Estimated variance:\t{jnp.var(chains.ravel()):+3.2f}")
        Running chain 0. Acceptance ratio: 0.02
        Running chain 1. Acceptance ratio: 0.02
        Running chain 2. Acceptance ratio: 0.02
        Running chain 3. Acceptance ratio: 0.02
        Estimated mean:
                                +18.20
        Estimated variance:
                                +644.37
```

Task 1.4: Plot the trace for each parameter and report the convergence diagnostics

```
In [108...
          def compute_Rhat(chains):
              Compute the Gelman-Rubin Rhat diagnostic for each parameter.
              Expects chains to be a jnp.array of shape (num chains, num samples, num para
              Returns a jnp.array of shape (num_params,).
              num_chains, num_samples, num_params = chains.shape
              chain_means = jnp.mean(chains, axis=1) # shape: (num_chains, num_params)
              overall_mean = jnp.mean(chains, axis=(0, 1)) # shape: (num_params,)
              B = (
                  num samples
                  / (num_chains - 1)
                  * jnp.sum((chain_means - overall_mean) ** 2, axis=0)
              ) # between-chain variance.
              # Use ddof=1 for unbiased sample variance
              W = jnp.mean(jnp.var(chains, axis=1, ddof=1), axis=0) # within-chain varian
              var_plus = ((num_samples - 1) / num_samples) * W + (1 / num_samples) * B
              Rhat = jnp.sqrt(var_plus / W)
              return Rhat
```

```
def compute_effective_sample_size(chains):
    Compute a basic effective sample size (S_eff) approximation for each paramet
    This implementation uses the lag autocorrelations of the merged chains.
    Returns a jnp.array of shape (num_params,).
    num_chains, num_samples, num_params = chains.shape
   total_samples = int(num_chains * num_samples)
    S_eff = jnp.zeros(num_params)
   chains np = np.array(chains) # switch to numpy for autocorrelation computat
    for p in range(num_params):
        # merge chains for parameter p
        x = chains_np[:, :, p].reshape(-1)
        # compute autocorrelation for increasing lag until the correlation becom
        ac sum = 0.0
        for lag in range(1, num_samples):
            # compute autocorrelation at lag 'lag'
            corr = np.corrcoef(x[:-lag], x[lag:])[0, 1]
           if corr < 0:</pre>
                break
            ac sum += corr
        S_eff = S_eff.at[p].set(total_samples / (1 + 2 * ac_sum))
    return S_eff
warm_up = 1000
# Compute number of warm-up samples discarded (per chain)
total warm up = num chains * warm up
print(f"Number of warm-up samples discarded: {total_warm_up}")
# Compute estimated overall mean and variance, pooling all samples
all_samples = chains.reshape(-1, chains.shape[-1]) # shape: (total_samples, num
# Compute convergence diagnostics.
Rhat = compute Rhat(chains)
total_samples = num_chains * (chains.shape[1]) # after warm-up
S eff = compute effective sample size(chains)
# Print results
for p in range(all_samples.shape[1]):
   print(f"\nParameter {p+1}:")
   print(f" Effective sample size: {S eff[p]:.0f}")
   print(f" ^R:
                                      {Rhat[p]:3.2f}")
# Plot trace for each parameter.
num params = chains.shape[-1]
fig, axes = plt.subplots(num_params, 1, figsize=(10, 3 * num_params))
if num params == 1:
    axes = [axes] # ensure axes is iterable
for p in range(num params):
    for chain in range(chains.shape[0]):
        axes[p].plot(chains[chain, warm_up:, p], label=f"Chain {chain+1}")
   axes[p].set_xlabel("Iteration")
    axes[p].set_ylabel(f"Parameter {p+1}")
    axes[p].set_title(f"Trace plot for Parameter {p+1}")
    axes[p].legend()
```

```
plt.tight_layout()
  plt.show()
Number of warm-up samples discarded: 4000
Parameter 1:
  Effective sample size:
                                 304
   ^R:
                                 1.00
Parameter 2:
  Effective sample size:
                                 414
   ^R:
                                 1.00
Parameter 3:
  Effective sample size:
                                 440
   ^R:
                                 1.00
                                          Trace plot for Parameter 1
     3
                Chain 1
  Parameter 1
                Chain 2
                Chain 3
     2
                Chain 4
           0
                             2000
                                                4000
                                                                   6000
                                                                                       8000
                                                   Iteration
                                          Trace plot for Parameter 2
   100
                                                                                              Chain 1
Parameter 2
                                                                                              Chain 2
    75
                                                                                              Chain 3
                                                                                              Chain 4
    50
    25
           0
                             2000
                                                4000
                                                                   6000
                                                                                       8000
                                                   Iteration
                                          Trace plot for Parameter 3
  0.65
                                                                                              Chain 1
Parameter 3
                                                                                              Chain 2
  0.60
                                                                                              Chain 3
                                                                                              Chain 4
  0.55
  0.50
           0
                                                4000
                                                                   6000
                             2000
                                                                                       8000
                                                   Iteration
```

Task 1.5: Estimate and report the posterior mean for each hyperparameter. Report the MCSE for each estimate.

```
# Compute estimated overall mean and variance, pooling all samples
all_samples = chains.reshape(-1, chains.shape[-1]) # shape: (total_samples, num
estimated_mean = jnp.mean(all_samples, axis=0)
estimated_var = jnp.var(all_samples, axis=0)
estimated_std = jnp.sqrt(estimated_var)
MC_error = estimated_std / jnp.sqrt(S_eff)

for p in range(all_samples.shape[1]):
```

```
print(f"\nParameter {p+1}:")
     print(f" Estimated mean:
                                       {estimated_mean[p]:+3.2f}")
     print(f" MCSE:
                                       {MC_error[p]:+3.2f}")
Parameter 1:
                        +1.15
 Estimated mean:
 MCSE:
                         +0.02
Parameter 2:
 Estimated mean:
                        +52.89
 MCSE:
                         +0.56
Parameter 3:
 Estimated mean:
                         +0.57
                         +0.00
```

#### Task 1.6: Estimate a 95% posterior credibility interval for each hyperparameter.

```
In [ ]: # Compute 95% credibility intervals for each hyperparameter.
        cred_intervals = np.percentile(np.array(all_samples), [2.5, 97.5], axis=0)
        for p in range(cred_intervals.shape[1]):
            print(
                f"Parameter {p+1} 95% Credibility Interval: [{cred_intervals[0, p]:.2f},
       Parameter 1 95% Credibility Interval: [0.60, 2.11]
       Parameter 2 95% Credibility Interval: [30.05, 74.45]
       Parameter 3 95% Credibility Interval: [0.53, 0.61]
In [ ]: data = jnp.load('Gustav/data_assignment3.npz')
        x_data, y_data = data['x'], data['t']
In [ ]: def design_matrix(x):
            return np.column_stack((x,np.ones(len(x))))
In [ ]: # Normal distribution
        def log_npdf(x, m, v):
            return -0.5 * (x - m) ** 2 / (v) - 0.5 * jnp.log(2 * jnp.pi * v)
        def npdf(x, m, v):
            return jnp.exp(log_npdf(x, m, v))
        # Half-normal distribution
        def log_half_npdf(x, m, v):
            return jnp.log(2) - 0.5 * (x - m) ** 2 / (v) - 0.5 * jnp.log(2 * jnp.pi * v)
        def half_npdf(x, m, v):
            return jnp.exp(log_half_npdf(x, m, v))
        # Logistic function
        def sigmoid(z):
            return 1 / (1 + jnp.exp(-z))
In [ ]: def plot summary(
            ax,
            Χ,
```

```
interval=95,
   num_samples=0,
   sample_color="k",
   sample_alpha=0.4,
   interval alpha=0.25,
   color="r",
   legend=True,
   title="",
   plot_mean=True,
   plot_median=False,
   label="",
   seed=0,
):
   b = 0.5 * (100 - interval)
   lower = jnp.percentile(s, b, axis=0).T
   upper = jnp.percentile(s, 100 - b, axis=0).T
   if plot median:
       median = jnp.percentile(s, 50, axis=0).T
       lab = "Median"
       if label:
           lab += " " + label
       ax.plot(x.ravel(), median, label=lab, color=color, linewidth=4)
   if plot_mean:
       mean = jnp.mean(s, axis=0).T
       lab = "Mean"
       if label:
           lab += " " + label
       ax.plot(x.ravel(), mean, "--", label=lab, color=color, linewidth=4)
   ax.fill_between(
       x.ravel(),
       lower.ravel(),
       upper.ravel(),
       color=color,
       alpha=interval_alpha,
       label=f"{interval}% Interval",
   )
   if num_samples > 0:
        np.random.seed(seed)
        idx_samples = np.random.choice(s.shape[0], size=num_samples, replace=Fal
        ax.plot(x, s[idx_samples, :].T, color=sample_color, alpha=sample_alpha)
   if legend:
       ax.legend(loc="best")
        ax.set_title(title, fontweight="bold")
```

# Part 2: Regression modelling using mixture of experts

Mixture of experts (MoE) model for regression

We consider two different linear models  $\boldsymbol{y}_n = \boldsymbol{w}_0^T \boldsymbol{x}_n + e_n$  and  $\boldsymbol{y}_n = \boldsymbol{w}_1^T \boldsymbol{x}_n + e_n$ , with weights  $\boldsymbol{w}_0$  and  $\boldsymbol{w}_1$  in respect to two different regions of  $\boldsymbol{x}_n$ , controlled by the latent variable  $z_n \in \{0,1\}$  which we let be Bernoulli distributed given the data  $\boldsymbol{x}_n$ 

$$oldsymbol{z}_n \mid oldsymbol{v} \sim \mathrm{Ber}ig(\sigmaig(oldsymbol{v}^Toldsymbol{x}_nig)ig),$$

where (v) are parameters and  $x_n = [x_n, 1]$ .

We then construct the conditional likelyhood as Gaussians for both data regions determined by the latent variable such that

$$p\left(y_n \mid x_n, z_n, oldsymbol{w}_0, oldsymbol{w}_1, \sigma_0^2, \sigma_1^2
ight) = egin{cases} \mathcal{N}\left(y_n \mid oldsymbol{w}_1^T oldsymbol{x}_n, \sigma_1^2
ight) & ext{if} \quad z_n = 1, \ \mathcal{N}\left(y_n \mid oldsymbol{w}_{z_n}^T oldsymbol{x}_n, \sigma_0^2
ight) & ext{if} \quad z_n = 0, \end{cases} = \mathcal{N}\left(y_n \mid oldsymbol{w}_{z_n}^T oldsymbol{x}_n, \sigma_0^2
ight)$$

with noise variance  $\sigma_{z_n}^2$ . Adding the following generic priors, we write up the joint distribution

$$egin{aligned} au, \sigma_0, \sigma_1 &\sim \mathcal{N}_+(0,1), \ w_0, w_1, oldsymbol{v} &\sim \mathcal{N}\left(oldsymbol{0}, au^T oldsymbol{I}
ight), \ y_n \mid z_n &\sim \mathcal{N}\left(oldsymbol{w}_{z_n}^T oldsymbol{x}_n, \sigma_{z_n}^2
ight), \end{aligned}$$

joint distribution:

$$egin{aligned} p\left(oldsymbol{y}, oldsymbol{w}_1, oldsymbol{w}_0, oldsymbol{v}, oldsymbol{z}, au, \sigma_0, \sigma_1 \mid oldsymbol{x} 
ight) = & \left[ \prod_{n=1}^N \mathcal{N}\left(oldsymbol{y}_n \mid oldsymbol{w}_{z_n}^T oldsymbol{x}_n, \sigma_{z_n}^2 
ight) \operatorname{Ber}ig(oldsymbol{z}_n \mid oldsymbol{\sigma}\left(oldsymbol{v}^T oldsymbol{x}_n 
ight) ig) 
ight] \mathcal{N}\left(oldsymbol{w}_0 \mid oldsymbol{o}, au^2 oldsymbol{I} 
ight) \mathcal{N}_+\left( au \mid 0, 1
ight) \mathcal{N}_+\left( au \mid 0, 1
ight) \mathcal{N}_+\left(\sigma_0 \mid oldsymbol{\sigma}_0 \mid oldsymbol{\sigma}_n \mid olds$$

### Task 2.1: Marginalize out each $z_n$ from to joint model.

Using the sum rule we marginalize out  $z_n$  from the joint distribution. Remembering that  $z_n$  is discrete, we have to sum and not integrate over  $z_n$ .

To simplify the sum, we will denote all Gaussians that are not dependent on  $z_n$  as  $p(\neg z)$  and sum over all  $2^N$  combinations of  $z_n$ ,  $z = \{z_n\}_{n=1}^N$ . Since each observation is independent of each other, we can move the sum inside the product

$$\left(\sum_{oldsymbol{z}} = \sum_{z_1} \sum_{z_2} \dots \sum_{z_N} = \prod_{n=1}^N \sum_{z_n} \right)$$
.

$$egin{aligned} \sum_{oldsymbol{z}} p\left(oldsymbol{y}, oldsymbol{w}_1, oldsymbol{w}_0, oldsymbol{v}, oldsymbol{z}, au_0, \sigma_1 \mid oldsymbol{x} 
ight) &= p\left(
eg oldsymbol{z}
ight) \sum_{oldsymbol{z}} \left[ \prod_{n=1}^N \mathcal{N}\left(y_n \mid oldsymbol{w}_{z_n}^T oldsymbol{x}_n, \sigma_{z_n}^2
ight) oldsymbol{\sigma}\left(oldsymbol{v}^T oldsymbol{x}_n
ight)^{z_n} (1) \ &= p\left(
eg oldsymbol{z}
ight) \prod_{n=1}^N \left[ \mathcal{N}\left(y_n \mid oldsymbol{w}_1^T oldsymbol{x}_n, \sigma_1^2
ight) oldsymbol{\sigma}\left(oldsymbol{v}^T oldsymbol{x}_n
ight) + \mathcal{N}\left(y_n \mid oldsymbol{w}_1^T oldsymbol{x}_n, \sigma_1^2
ight) oldsymbol{\sigma}\left(oldsymbol{v}^T oldsymbol{x}_n
ight) + \mathcal{N}\left(y_n \mid oldsymbol{w}_1^T oldsymbol{x}_n, \sigma_1^2
ight) oldsymbol{\sigma}\left(oldsymbol{v}^T oldsymbol{x}_n
ight) + \mathcal{N}\left(y_n \mid oldsymbol{w}_1^T oldsymbol{w}_1, \sigma_1^2
ight) oldsymbol{\sigma}\left(oldsymbol{v}^T oldsymbol{x}_n
ight) + \mathcal{N}\left(y_n \mid oldsymbol{w}_1, oldsymbol{v}^T oldsymbol{w}_n
ight) oldsymbol{\sigma}\left(oldsymbol{v}^T oldsymbol{x}_n
ight) + \mathcal{N}\left(y_n \mid oldsymbol{w}_1, oldsymbol{w}_1, oldsymbol{w}_1, oldsymbol{v}^T oldsymbol{w}_n
ight) + \mathcal{N}\left(y_n \mid oldsymbol{w}_1, oldsymbol{w}_1, oldsymbol{w}_2, oldsymbol{w}_1, oldsymbol{w}_2, oldsymbol{w}_2, oldsymbol{w}_1, oldsymbol{w}_2, oldsymbol{w}_$$

We see that the likelihood for each observation has become a mixture of gaussians after the marginalization of  $z_n$ .

### Task 2.2: Python function to evaluate the marginalized log joint distribution

We will separate the terms from above, such that we have:

$$\log p\left(\boldsymbol{y}, \boldsymbol{w}_0, \boldsymbol{w}_1, \boldsymbol{v}, \tau, \sigma_0, \sigma_1 | \boldsymbol{x}\right) = \log p(\neg \boldsymbol{z}) + \log p\left(\boldsymbol{y} \mid \boldsymbol{x}, \boldsymbol{w}_0, \boldsymbol{w}_1, \boldsymbol{v}, \sigma_0, \sigma_1\right)$$

$$\log p(
eg oldsymbol{z}) = -3\log(2\pi) - 6\log au - rac{oldsymbol{w_0}^Toldsymbol{w_0} + oldsymbol{w_1}^Toldsymbol{w_0} + oldsymbol{v}^Toldsymbol{v}}{2 au^2} + rac{3}{2}\log au$$

$$\log p\left(\boldsymbol{y} \mid \boldsymbol{x}, \boldsymbol{w}_{0}, \boldsymbol{w}_{1}, \boldsymbol{v}, \sigma_{0}, \sigma_{1}\right) = \sum_{n=1}^{N} \log \left[\mathcal{N}\left(y_{n} \mid \boldsymbol{w}_{1}^{T} \boldsymbol{x}_{n}, \sigma_{1}^{2}\right) \sigma\left(\boldsymbol{v}^{T} \boldsymbol{x}_{n}\right) + \mathcal{N}\left(y_{n} \mid \boldsymbol{w}_{0}^{T} \boldsymbol{x}_{n}\right)\right]$$

We see that we can vectorize the Gaussian mixture terms wrt. the number of data points for the logaritmic transformation which was not possible for the joint distribution.

```
In [ ]: def log_joint(y, x, theta):
            Evaluates the log joint distribution
            Parameters:
                    : array of N observations (shape: (N,))
                    : design matrix for N observations (shape: (N,2)); each row is [x_n]
              theta: vector of parameters of shape (9,), where
                       theta[0:2] -> w0 (slope and intercept)
                       theta[2:4] -> w1 (slope and intercept)
                       theta[4:6] -> v (slope and intercept)
                       theta[6] -> tau
                       theta[7]
                                   -> sigma0
                       theta[8] -> sigma1
            Returns:
              log joint density evaluated at theta.
            # Unpack parameters
            w0 = theta[0:2]
            w1 = theta[2:4]
            v = theta[4:6]
            tau = theta[6]
            sigma0 = theta[7]
            sigma1 = theta[8]
            # ----- Prior term (log p(~z)) ------
            # \log p(\sim z) = -3\log(2pi) - 6\log(tau) - (w0^T w0 + w1^T w1 + v^T v)/(2*tau^2)
                     + 1.5 log(2) - 1.5 log(pi) - (tau^2+sigma0^2+sigma1^2)/2
            log_p_neg_z = (
                -3 * jnp.log(2 * jnp.pi)
                - 6 * jnp.log(tau)
                -(jnp.dot(w0, w0) + jnp.dot(w1, w1) + jnp.dot(v, v)) / (2 * tau**2)
                + 1.5 * jnp.log(2)
```

```
- 1.5 * jnp.log(jnp.pi)
    - (tau**2 + sigma0**2 + sigma1**2) / 2
)
# ----- Likelihood term -----
# Compute predictions:
# For each observation x_n (a row vector of shape (2,)), we compute:
# m1 = w1^T x_n, m0 = w0^T x_n, and v_{dot} = v^T x_n
m1 = jnp.sum(x * w1, axis=1)
m0 = jnp.sum(x * w0, axis=1)
v_{dot} = jnp.sum(x * v, axis=1)
# For each observation, form the likelihood contribution (a mixture) as:
\# term1 = N(y_n \mid m1, sigma1^2) * sigmoid(v_dot)
# term2 = N(y_n \mid m0, sigma0^2) * sigmoid(-v_dot)
term1 = jnp.exp(log_npdf(y, m1, sigma1**2)) * sigmoid(v_dot)
term2 = jnp.exp(log_npdf(y, m0, sigma0**2)) * sigmoid(-v_dot)
# Sum of both (for each observation) and then take the log and sum over all
log_likelihood = jnp.sum(jnp.log(term1 + term2))
return log_p_neg_z + log_likelihood
```

Task 2.3: Metropolis-Hasting sampler to infer all parameters

Below is the code for the Metropolis-Hasting sampler, 'MetH'. Since the we have a mixed Gaussian as target distribution, we use unimodal Gaussians as proposal distributions for the parameters.

```
In [ ]: def MetH(log_target, num_params, tau, num_iter, theta_init=None, seed=0):
            """Runs a Metropolis-Hastings sampler
            Arguments:
            log target:
                                function for evaluating the log target distribution, i.e
                                number of parameters of the joint distribution (integer)
            num params:
                               standard deviation of the Gaussian proposal distribution
            tau:
            num iter:
                              number of iterations (integer)
            theta_init:
                                vector of initial parameters (np.array with shape (num_p
            seed:
                                seed (integer)
            returns
            thetas
                                np.array with MCMC samples (np.array with shape (num ite
            # set initial key
            key = random.PRNGKey(seed)
            if theta init is None:
                theta_init = jnp.zeros((num_params))
            # prepare lists
            thetas = [theta_init]
            accepts = []
            log_p_theta = log_target(theta_init)
            for k in range(num_iter):
```

```
# update keys: key_proposal for sampling proposal distribution and key_a
    key, key_proposal, key_accept = random.split(key, num=3)
    # get the last value for theta and generate new proposal candidate
    theta cur = thetas[-1]
    theta_star = theta_cur + tau * random.normal(key_proposal, shape=(num_pa
    # evaluate the log density for the candidate sample
    log_p_theta_star = log_target(theta_star)
    # compute acceptance probability
    log_r = log_p_theta_star - log_p_theta
    A = min(1, jnp.exp(log_r))
    # accept new candidate with probability A
    if random.uniform(key_accept) < A:</pre>
        theta_next = theta_star
        log_p_theta = log_p_theta_star
        accepts.append(1)
    else:
        theta_next = theta_cur
        accepts.append(0)
    thetas.append(theta_next)
print("Acceptance ratio: %3.2f" % jnp.mean(jnp.array(accepts)))
# return as np.array
thetas = jnp.stack(thetas)
# check dimensions and return
assert thetas.shape == (
    num_iter + 1,
    num params,
), f"The shape of thetas was expected to be ({num_iter+1}, {num_params}), bu
return thetas
```

Looking at the data and the derived log joint distribution, we expect  $z_n=1$  to be the first part with a positive slope and  $z_n=0$  to be the second part with a negative slope. Because of this convention, we expect the slope parameter of  $\boldsymbol{v}$  to be negative. We also notice the variance for the data points in region  $z_n=0$  is much larger than in the in region  $z_n=1$ . With good initial guesses for the parameters, we might have a good chance of being close to convergence (for relatively few iterations), when picking a low proposal variance to get a high acceptance ratio.

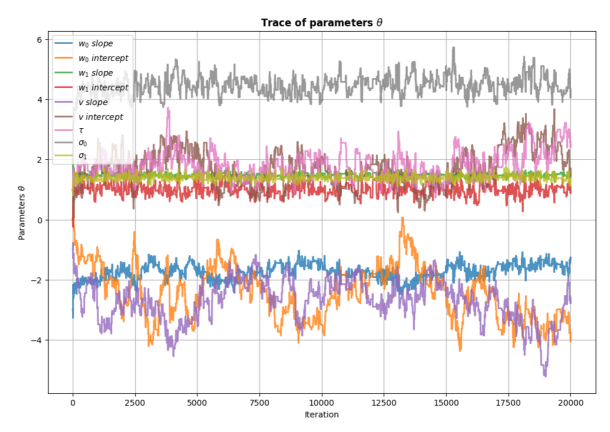
Hence, we guess for the parameters to be near:

$$heta=\left[oldsymbol{w}_0=inom{-3}{0},\;oldsymbol{w}_1=inom{2}{0},\;oldsymbol{v}=inom{-1}{1},\; au=1,\;\sigma_0=2,\;\sigma_1=1
ight]$$

We set the variance of the proposal distribution to 0.2 (called step\_size in code) and run the sampler for 20.000 iterations. We expect higher acceptance ratio with the low proposal variance.

```
In [ ]: # specify number of parameters in the target distribution
        num params = 9
        # target distribution = log joint distribution
        num_iterations = 20000
        # proposal variance for each parameter
        step_size = 0.2
        # theta = \{w0[0], w0[1], w1[0], w1[1], v[0], v[1], tau, sigm0, sigma1\}
        theta_lab = [r'$w_0\ slope$', r'$w_0\ intercept$', r'$w_1\ slope$', r'$w_1\ inte
                     r'$v\ slope$', r'$v\ intercept$', r'$\tau$', r'$\sigma_0$', r'$\sigm
        w0_{init} = [-3.0, 0.0]
        w1_{init} = [2.0, 0.0]
        v_{init} = [-1.0, 1.0]
        tau_init = [1.0]
        sigma0 init = [2.0]
        sigma1_init = [1.0]
        theta_init = jnp.array(w0_init + w1_init + v_init + tau_init + sigma0_init + sig
        p_target = lambda theta: log_joint(y_data, design_matrix(x_data), theta)
        # run sampler
        thetas = MetH(p_target, num_params, step_size, num_iterations, theta_init=theta_
        # plot resutls
        xs = np.linspace(-12, 12, 1000)
        fig, axes = plt.subplots(1, 1, figsize=(12, 8))
        for i in range(num_params):
            axes.plot(thetas[:,i], alpha=0.8, linewidth=2,label=theta_lab[i])
        axes.set_xlabel('Iteration')
        axes.set_ylabel('Parameters $\\theta$')
        axes.set_title('Trace of parameters $\\theta$', fontweight='bold')
        axes.legend()
        axes.grid()
        plt.show()
```

Acceptance ratio: 0.07

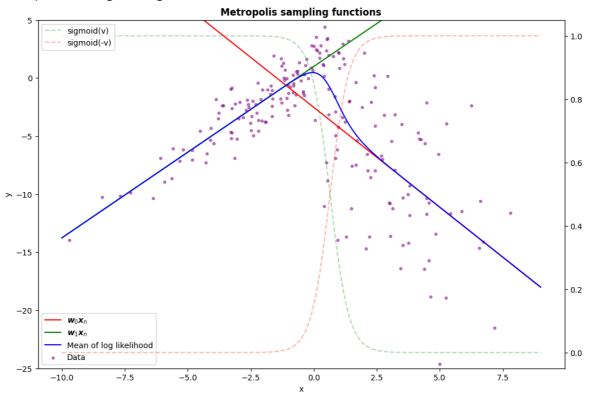


```
In [ ]: warm_up = int(0.2*num_iterations)
        theta_mean = np.mean(thetas[warm_up:], axis=0)
        w0 = jnp.array([theta_mean[0],theta_mean[1]])
        w1 = jnp.array([theta_mean[2],theta_mean[3]])
        v = jnp.array([theta_mean[4],theta_mean[5]])
        tau = theta_mean[6]
        sigma0 = theta_mean[7]
        sigma1 = theta_mean[8]
        x = np.linspace(-10, 9, 500)
        # design matrix
        X = design matrix(x)
        X1 = X@w1
        X0 = X@w0
        fig, axes = plt.subplots(1,1, figsize=(12, 8))
        #### Metro-Hastings samples mean:
        y = sigmoid(X@v)*X1 + sigmoid(-X@v)*X0
        ax2 = axes.twinx()
        ax2.plot(x,sigmoid(X@v),'--', color='green', alpha=0.3, label='sigmoid(v)')\\
        ax2.plot(x,sigmoid(-X@v),'--', color='red', alpha=0.3, label='sigmoid(-v)')
        ax2.legend(loc='upper left')
        axes.plot(x, X0, color='red', label=r'$\boldsymbol{w}_0 \boldsymbol{x}_n$')
        axes.plot(x, X1, color='green', label=r'\boldsymbol\{w\}_1 \boldsymbol\{x\}_n$')
        axes.plot(x, y, color='blue', label='Mean of log likelihood')
        # plot contour of log joint distribution and data
        axes.set_title('Metropolis sampling functions', fontweight='bold')
        axes.set_xlabel('x')
```

```
axes.set_ylabel('y')

#log_joint_vectorized = vmap(lambda xi, yi: log_joint(yi, xi, theta_mean))
axes.set_ylim(-25,5)
axes.scatter(x_data, y_data, c='purple', s=10, alpha=0.5, label='Data')
axes.legend(loc='lower left')
```

<matplotlib.legend.Legend at 0x1bfc5a74ce0>



Task 2.4: posterior mean and 95% credibility intervals for all parameters

```
In [ ]: # Assuming you already have these:
        # thetas, warm_up, theta_init
        theta_sample = thetas[warm_up:, :]
        # Round to 3 decimals
        lower = jnp.round(jnp.percentile(theta_sample, 2.5, axis=0), 3)
        upper = jnp.round(jnp.percentile(theta_sample, 97.5, axis=0), 3)
        mean = jnp.round(jnp.mean(theta_sample, axis=0), 3)
        # Labels with HTML
        labels = [
             'w<sub>0</sub>: slope', 'w<sub>0</sub>: intercept',
            'w<sub>1</sub>: slope', 'w<sub>1</sub>: intercept',
             'v slope', 'v intercept',
             'τ',
             '&\pmx03C3;<sub>0</sub>', # \sigma_0
             'σ<sub>1</sub>' # \sigma_1
        1
        # Format the intervals as strings: "[lower, upper]"
        low_up = [f''[\{1:.3f\}; \{u:.3f\}]'' for 1, u in zip(lower, upper)]
        # Create DataFrame
        cred int = pd.DataFrame({
```

```
'Parameter': labels,
   'Initial': theta_init,
   'Mean': mean,
   '95%-Credibility interval': low_up,
})

cred_int = cred_int.set_index('Parameter')

# Display with HTML rendering
display(HTML(cred_int.to_html(escape=False)))
```

### Initial Mean 95%-Credibility interval

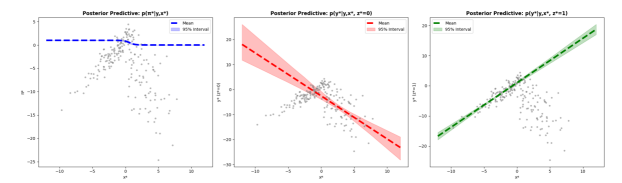
Parameter			
w <sub>0</sub> : slope	-3.0	-1.719	[-2.241 ; -1.291]
w <sub>0</sub> : intercept	0.0	-2.528	[-3.897 ; -0.686]
w <sub>1</sub> : slope	2.0	1.476	[1.356 ; 1.595]
w <sub>1</sub> : intercept	0.0	0.997	[0.631 ; 1.363]
v slope	-1.0	-2.682	[-4.247 ; -1.534]
v intercept	1.0	1.727	[0.648 ; 2.991]
τ	1.0	1.821	[1.051 ; 2.970]
$\sigma_0$	2.0	4.499	[3.926 ; 5.148]
$\sigma_1$	1.0	1.396	[1.208 ; 1.595]

### Task 2.5: Plot posterior predictive distribution for

```
p(\pi^* \mid \boldsymbol{y}, \boldsymbol{x}^*), \ p(y^* \mid \boldsymbol{y}, \boldsymbol{x}^*, z^* = 0) \ \text{and} \ p(y^* \mid \boldsymbol{y}, \boldsymbol{x}^*, z^* = 1)
```

```
In [ ]: # -----
        # Compute posterior predictive samples
        # -----
        # Create new x* values
        x_star = jnp.linspace(-12, 12, 1000)
        X_star = design_matrix(x_star) # design matrix (1000, 2)
        # Assume theta_sample is available from your MCMC sampler (shape: (M, 9))
        # Extract the relevant posterior samples:
          w0_samples: parameters for branch z=0, shape (M, 2)
          w1_samples: parameters for branch z=1, shape (M, 2)
          v_samples : parameters for mixing, shape (M, 2)
        w0_samples = theta_sample[:, 0:2]
        w1_samples = theta_sample[:, 2:4]
        v samples = theta sample[:, 4:6]
        # For each sample we compute the predictive functions:
        # \pi^* = sigmoid(X_star @ v)
          y*|z*=0 = X_star @ w0
        y*/z*=1 = X_star @ w1
        # Compute \pi^* samples: iterate over M samples to get an array of shape (M, len(x
        pi_star_samples = jnp.array(
          [sigmoid(X_star @ v_samples[i]) for i in range(v_samples.shape[0])]
```

```
# Compute y^* samples for branch z^*=0
y_star_z0_samples = jnp.array(
    [X_star @ w0_samples[i] for i in range(w0_samples.shape[0])]
# Compute y^* samples for branch z^*=1
y_star_z1_samples = jnp.array(
   [X_star @ w1_samples[i] for i in range(w1_samples.shape[0])]
# -----
# Plot predictions on top of the data
fig, axes = plt.subplots(1, 3, figsize=(20, 6))
# Plot posterior predictive for \pi^*
plot_summary(
   axes[0],
   x star,
   pi_star_samples,
   title="Posterior Predictive: p(\pi^*|y,x^*)",
   color="blue",
)
axes[0].set_xlabel("x*")
axes[0].set_ylabel("\pi*")
axes[0].scatter(
   np.array(x_data), np.array(y_data), color="grey", s=10, alpha=0.5, label="Da
# Plot posterior predictive for y^* with z^* = 0
plot_summary(
   axes[1],
   x_star,
   y_star_z0_samples,
   title="Posterior Predictive: p(y*|y,x*, z*=0)",
   color="red",
)
axes[1].set_xlabel("x*")
axes[1].set_ylabel("y* (z*=0)")
axes[1].scatter(
   np.array(x data), np.array(y data), color="grey", s=10, alpha=0.5, label="Da
# Plot posterior predictive for y^* with z^* = 1
plot_summary(
   axes[2],
   x_star,
   y star z1 samples,
   title="Posterior Predictive: p(y*|y,x*, z*=1)",
   color="green",
)
axes[2].set_xlabel("x*")
axes[2].set_ylabel("y* (z*=1)")
axes[2].scatter(
   np.array(x_data), np.array(y_data), color="grey", s=10, alpha=0.5, label="Da
)
plt.tight_layout()
plt.show()
```



Task 2.6: Plot posterior predictive distribution for  $p(y^* \mid y, x^*)$ 

```
In [ ]: # Create new x* values
        x_{star} = jnp.linspace(-12, 12, 1000)
        X_star = design_matrix(x_star) # design matrix (1000, 2)
        # Extract posterior samples from theta sample (shape: (M,9))
        w0_samples = theta_sample[:, 0:2]
        w1_samples = theta_sample[:, 2:4]
        v_samples = theta_sample[:, 4:6]
        # Compute predictive y* for each sample:
        \# y^* = sigmoid(X_star @ v) * (X_star @ w1) + (1 - sigmoid(X_star @ v)) * (X_star @ v)
        y_star_pred_samples = jnp.array([
            sigmoid(X_star @ v_samples[i]) * (X_star @ w1_samples[i]) +
            (1 - sigmoid(X_star @ v_samples[i])) * (X_star @ w0_samples[i])
            for i in range(theta_sample.shape[0])
        ])
        # Plot using plot_summary on top of the data
        fig, ax = plt.subplots(1, 1, figsize=(12, 8))
        plot_summary(ax, x_star, y_star_pred_samples, title="Posterior Predictive: p(y*|
        ax.set_xlabel("x*")
        ax.set_ylabel("y*")
        ax.scatter(np.array(x data), np.array(y data), color="grey", s=10, alpha=0.5, la
        plt.tight_layout()
        plt.show()
```

