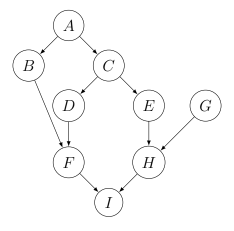
COMP0085 Summative Assignment

Jan 4, 2023

Question 1

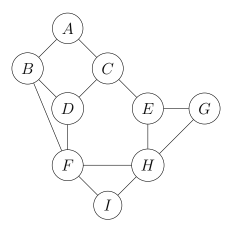
(a)

The directed acyclic graph:

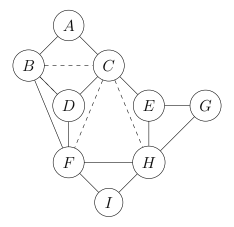


(b)

The moralised graph:

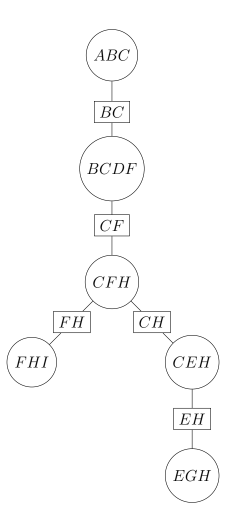


An effective triangulation:

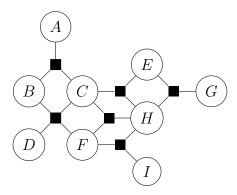


where the dashed lines are edges added to triangulate the moralised graph.

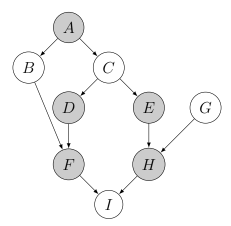
The resulting junction tree:



where the circular nodes are cliques and the square nodes are separators/factors. The junction tree redrawn as a factor graph:



(c)



The set $\{A, D, E, F, H\}$ is a non-unique smallest set of molecules such that if the concentrations of the species within the set are known, the concentrations of the others $\{B, C, G, I\}$ would all be independent (conditioned on the measured ones).

(d)

Using our factor analysis model, we can describe the biochemical pathway as:

$$\delta[\mathbf{x}] = \Lambda \mathbf{z} + \epsilon$$

where $\delta[\mathbf{x}]$ are the concentration perturbations, $\epsilon \sim \mathcal{N}(0, \Psi)$, and the latent factors $z \sim \mathcal{N}(0, I)$. From the graph structure, we know that:

Having observations for $\delta[B]$, $\delta[D]$, $\delta[E]$ and $\delta[G]$:

We can see that these simplify to the equations:

$$\delta[B] = \Lambda_{BA} z_A + \epsilon_B$$

$$\delta[D] = \Lambda_{DC} z_C + \epsilon_D$$

$$\delta[E] = \Lambda_{EC} z_C + \epsilon_E$$

$$\delta[G] = \epsilon_G$$

Thus, we see that the only latent variables present are z_A and z_C , so would expect to recover the factors of A and C, the two parent nodes of the observations.

(e)

Question 2

(a)

We want the posterior mean and covariance over a and b. Defining a weight vector \mathbf{w} :

$$\mathbf{w} = \begin{bmatrix} a \\ b \end{bmatrix}$$

Our distribution for w:

$$P(\mathbf{w}) = \mathcal{N}\left(\begin{bmatrix} \mu_a \\ \mu_b \end{bmatrix}, \begin{bmatrix} \sigma_a^2 & 0 \\ 0 & \sigma_b^2 \end{bmatrix}\right) = \mathcal{N}(\mu_\mathbf{w}, \Sigma_\mathbf{w})$$

Moreover, for our data $\mathcal{D} = \{X, Y\}$:

$$P(\mathcal{D}|\mathbf{w}) = \mathcal{N}\left(\mathbf{Y} - \mathbf{w}^T \mathbf{X}, \sigma^2 \mathbf{I}\right)$$

where
$$\mathbf{X} = \begin{bmatrix} t_1 & t_2 \cdots t_N \\ 1 & 1 \cdots 1 \end{bmatrix} \in \mathbb{R}^{2 \times N}$$
 and $\mathbf{Y} \in \mathbb{R}^{1 \times N}$.

Knowing:

$$P(\mathbf{w}|\mathcal{D}) \propto P(\mathcal{D}|\mathbf{w})P(\mathbf{w})$$

we can substitute the above distributions:

$$P(\mathbf{w}|\mathcal{D}) \propto \exp\left(\frac{-1}{2\sigma^2} \left(\mathbf{Y} - \mathbf{w}^T \mathbf{X}\right) \left(\mathbf{Y} - \mathbf{w}^T \mathbf{X}\right)^T\right) \exp\left(\frac{-1}{2} \left(\mathbf{w} - \mu_{\mathbf{w}}\right)^T \Sigma_{\mathbf{w}}^{-1} \left(\mathbf{w} - \mu_{\mathbf{w}}\right)\right)$$

expanding:

$$\log P(\mathbf{w}|\mathcal{D}) \propto \frac{-1}{2} \left(\frac{\mathbf{Y}\mathbf{Y}^T}{\sigma^2} - 2\mathbf{w}^T \frac{\mathbf{X}\mathbf{Y}^T}{\sigma^2} + \mathbf{w}^T \frac{\mathbf{X}\mathbf{X}^T}{\sigma^2} \mathbf{w} + \mathbf{w}^T \boldsymbol{\Sigma}_{\mathbf{w}}^{-1} \mathbf{w} - 2\mathbf{w}^T \boldsymbol{\Sigma}_{\mathbf{w}}^{-1} \boldsymbol{\mu}_{\mathbf{w}} + \boldsymbol{\mu}_{\mathbf{w}}^T \boldsymbol{\Sigma}_{\mathbf{w}}^{-1} \boldsymbol{\mu}_{\mathbf{w}} \right)$$

collecting w terms:

$$\log P(\mathbf{w}|\mathcal{D}) \propto \frac{-1}{2} \left(\mathbf{w}^T \left(\frac{\mathbf{X} \mathbf{X}^T}{\sigma^2} + \Sigma_{\mathbf{w}}^{-1} \right) \mathbf{w} - 2 \mathbf{w}^T \left(\frac{\mathbf{X} \mathbf{Y}^T}{\sigma^2} + \Sigma_{\mathbf{w}}^{-1} \mu_{\mathbf{w}} \right) \right)$$

Knowing that the posterior $P(\mathbf{w}|\mathcal{D})$ will be Gaussian with mean $\bar{\mu}_w$ and covariance $\bar{\Sigma}_w$, we can see that expanding the exponent component would have the form:

$$(\mathbf{w} - \bar{\mu}_w)^T \bar{\Sigma}_w^{-1} (\mathbf{w} - \bar{\mu}_w) = \mathbf{w}^T \bar{\Sigma}_w^{-1} \mathbf{w} - 2\mathbf{w}^T \bar{\Sigma}_w^{-1} \bar{\mu}_w + \bar{\mu}_w^T \bar{\Sigma}_w^{-1} \bar{\mu}_w$$

Thus we can identify the posterior covariance:

$$\bar{\Sigma}_w = \left(\frac{\mathbf{X}\mathbf{X}^T}{\sigma^2} + \Sigma_{\mathbf{w}}^{-1}\right)^{-1}$$

and the posterior mean:

$$\bar{\mu}_w = \bar{\Sigma}_w \left(\frac{\mathbf{X} \mathbf{Y}^T}{\sigma^2} + \Sigma_{\mathbf{w}}^{-1} \mu_{\mathbf{w}} \right)$$

Computing the posterior mean and covariance over a and b given by the CO_2 data:

		value
arameters	а	1.828457
	-	1.020101
	b	334.203782

Figure 1: The Posterior Mean

Figure 2: The Posterior Covariance

(b)

Plotting the residuals:

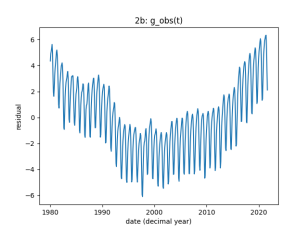


Figure 3: $g_{obs}(t)$

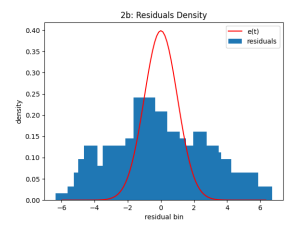


Figure 4: Density Estimation of Residuals vs $e(t) \sim \mathcal{N}(0,1)$

We can see that the residuals do not perfectly conform to our prior over $e(t) \sim \mathcal{N}(0,1)$. The density estimation shows that a mean of zero is a reasonable prior belief however the data does not seem to exhibit unit variance. Also we know it's not iid because timeseries.

(c & d)

We are considering the kernel:

$$k(s,t) = \theta^2 \left(\exp\left(-\frac{2\sin^2(\pi(s-t)/\tau)}{\sigma^2}\right) + \phi^2 \exp\left(-\frac{(s-t)^2}{2\eta^2}\right) \right) + \zeta^2 \delta_{s=t}$$

We can make qualitative observations this kernel by visualising the covariance (gram) matrix:



Figure 5: Covariance Matrix

We can observe a striped pattern which indicate higher covariance at regular intervals. This can be attributed to the sinusoidal term in the kernel and encourages sinusoidal functions. Additionally, we can see that covariance values also decay as they are further away from the diagonal. This can be attributed to the exponential term in the kernel, encouraging points closer in time to be more correlated and vice versa. From our CO_2 data, we would want a class of functions which exhibit both of these behaviours as the data looks sinusoidal (seasonal with respect to each year) and correlations locally.

We can also visualise some samples from a Gaussian Process with the same covariance matrix and zero mean. This verifies our observations about the covariance matrix.

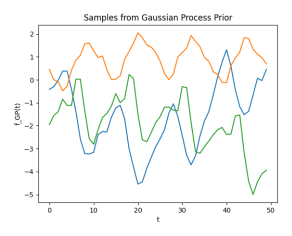


Figure 6: Samples from a zero mean GP with the provided covariance kernel

More specifically, we can see how changing each hyper-parameter will affect the characteristics of the function.

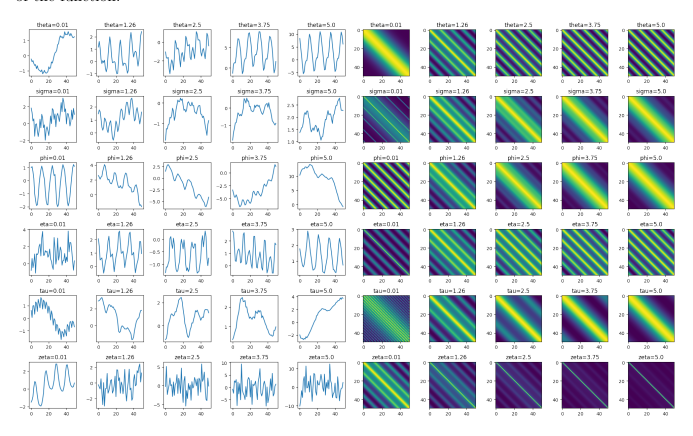


Figure 7: Samples for different parameters

Figure 8: Covariances for different parameters

- θ : As θ increases, we see more pronounced periodic behavior in the sample function. The covariance matrix shows how increasing θ visually reveals the striped periodic component. This is expected because it is the parameter that adjusts the weight of the periodic component.
- σ : As σ increases, we see reduced periodic behaviour in the sample function. The covariance matrix shows how increasing σ will increase covariance values in the off-diagonals. This is expected because it adjusts the lengthscale of the periodic portion of the kernel, which ends up dominating the function.
- ϕ : As ϕ increases, we see the ratio of the amplitude of the periodicity component of the sample function reduces compared to the baseline. The covariance matrix shows how increasing ϕ will start to increase the non-periodic component. This is expected because it adjusts the weight of the non-periodic portion of the kernel, thus the periodic component remains the same (i.e.same amplitude) but the large baseline shifts from increasing ϕ ends up dominating the function visually.
- η : As η increases we see smoother sample functions. This is expected because the η increases the lengthscale of the non-periodic component, allowing for smoother functions. This causes the off-diagonals of the gram matrix to increase, however the periodic component is still maintained because η doesn't affect the relative weight of the two components.

- τ : As τ increases, the period of the periodic function increases. We can see this reflected in the stripes in the gram matrix getting further apart. This makes sense because we are adjusting the period in the sinusoid function of the periodic term with τ .
- ζ : As ζ increases, the function becomes less smooth. This is because the ζ parameter adjusts the weight of the $\delta_{s=t}$ parameter. This places stronger emphasis on the independence of each timestep, which can be seen with the reduction of relative magnitude of off-diagonals in the gram matrix. However, this is simply masking the periodic and squared-exponential terms as we can see with the increased magnitude of the functions as ζ increases.

(e)

Suitable values for hyper-parameters can be chosen through a combination of visual inspection and prior knowledge. For example, it is a reasonable assumption that the CO_2 concentration levels have a strong yearly seasonality behaviour due to the cyclic changes in temperature, humidity, etc. Thus we can choose $\tau = 1$ to ensure functions with a period of one year to reflect this knowledge. It can be difficult to quantitatively choose values for the other parameters as they can relate to the uncertainty exhibited in the data (i.e.the smoothness of the function). One approach is to maximise:

$$\log P(\mathbf{Y}|\mathbf{X}) = -\frac{1}{2}\mathbf{Y}^{T}(\mathbf{K} + \sigma^{2}\mathbf{I})^{-1}\mathbf{Y} - \frac{1}{2}\log|\mathbf{K} + \sigma^{2}\mathbf{I}| - \frac{n}{2}\log(2\pi)$$

the log-likelihood of the posterior distribution with respect to the given data where \mathbf{K} is the gram matrix for the kernel (equation 2.30 from http://gaussianprocess.org/gpml/chapters/RW2.pdf). We can define a loss function as the negative log-likelihood and employ gradient-based algorithms to find optimal parameters.

Comparing the hyperparameters corresponding to before and after training side by side:

	value
parameter	
eta (kernel)	5.0
phi (kernel)	10.0
sigma	1.0
sigma (kernel)	5.0
tau (kernel)	1.0
heta (kernel)	5.0
zeta (kernel)	2.0

Figure 9: Untrained hyperparameters

Figure 10: Trained Hyperparmaeters

We can analyse some of the changes in these parameters after training to gain some insights. We can see that τ remains the same as we would expect given the yearly seasonality we have prior knowledge of. On the other hand, the value for ζ is significantly reduced signifying that $\delta_{s=t}$ is not a very good kernel for representing the data as datapoints at different timesteps do exhibit correlations.

(f)

Extrapolating the CO_2 concentration levels:

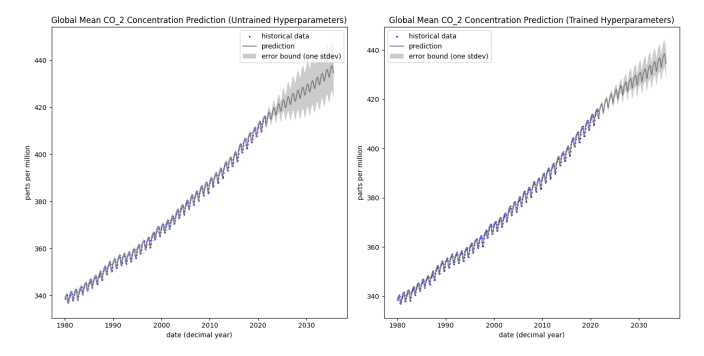


Figure 11: Untrained extrapolation

Figure 12: Trained extrapolation

We can see that the extrapolation shows a continued increase in CO_2 in the future. This follows our expectations given that the levels has been steadily increasing in the past. Moreover, the concentration continues to exhibit yearly seasonality (for the trained extrapolation) as we would expect. We can see that the conclusions can be quite sensitive to kernel hyperparameters when comparing the values from before and after training. Prior to training, the extrapolated prediction is not representative of the given data, with pretty much no seasonal behaviour and very large uncertainty. After training, we can see that the prediction is much more reasonable, and qualitatively the uncertainty bounds seem to exhibit the historical variability in the data.

(g)

This procedure is not fully Bayesian because despite using a posterior estimate of our linear regression terms, we only use a point estimate when making prediction. For a fully Bayesian approach, we should also incorporate the uncertainty of the linear regression parameters into our extrapolation/uncertainty bounds. For our procedure, we only include the uncertainty of g(t) however it can be observed in the plots that the trend is not perfectly linear so this should be reflected in the uncertainty of our extrapolation. Another approach could be to add a linear kernel to our combined kernel function and model f(t) directly with our kernel, removing the linear regression component in our procedure. Thus our kernel extrapolation would incorporate the uncertainty of all components of our signal.

The Python code for Bayesian Linear Regression:

```
from dataclasses import dataclass
3
     import numpy as np
 6
     @dataclass
      class LinearRegressionParameters:
           Parameters for linear regression
10
11
12
           mean: np.ndarray # weight vector (1, number of features)
13
14
           covariance: np.ndarray # covariance matrix on mean (number of features, number of features)
           @property
16
17
           def precision(self) -> np.ndarray:
    return np.linalg.inv(self.covariance)
18
19
           \begin{array}{lll} \textbf{def} & \texttt{predict} \, (\, \texttt{self} \, \, , \, \, \, \texttt{x: np.ndarray} \, ) \, \, -\!\!\!> \, \texttt{np.ndarray} \, : \end{array}
20
21
22
                Linear regression prediction.
                :param x: design matrix (number of features, number of data points) :return: predicted response matrix (1, number of data points)
23
24
25
26
                return self.mean.T @ x
27
28
29
30
     {\bf class\ Theta:} \\ {\bf linear-regression-parameters:\ LinearRegressionParameters}
31
32
33
34
           @property
35
           def variance (self) -> float:
36
                return self.sigma**2
38
          def precision(self) -> float:
return 1 / self.variance
40
41
42
     def compute_linear_regression_posterior(
44
           x: np.ndarray,
45
           y: np.ndarray,
           prior_linear_regression_parameters: LinearRegressionParameters,
47
            esiduals_precision: float
     ) -> LinearRegressionParameters:
48
49
           Compute the parameters of the posterior distribution on the linear regression weights
51
           : param \ x: \ design \ matrix \ (number \ of \ features \ , \ number \ of \ data \ points)
           :param y: response matrix (1, number of data points)
:param prior_linear_regression_parameters: parameters for the prior distribution on the linear regression
           :param residuals-precision: the precision of the residuals of the linear regression :return: parameters for the posterior distribution on the linear regression weights
57
58
           posterior_covariance = np.linalg.inv(
59
                residuals\_precision * x @ x.T + prior\_linear\_regression\_parameters.precision
           posterior_mean = posterior_covariance @ (
61
                residuals_precision * x @ y.T
+ prior_linear_regression_parameters.precision
@ prior_linear_regression_parameters.mean
62
65
66
           return LinearRegressionParameters (
                mean=posterior_mean, covariance=posterior_covariance
```

src/models/bayesian_linear_regression.py

The Python code for kernels:

```
from abc import ABC, abstractmethod
     from dataclasses import dataclass
3
     import jax.numpy as jnp
     from jax import vmap
 6
     @dataclass
     class KernelParameters (ABC):
10
          An abstract dataclass containing the parameters for a kernel.
11
12
13
14
     class Kernel (ABC):
16
17
          An abstract kernel.
19
20
          Parameters: KernelParameters = None
21
22
          @abstractmethod
23
          def _kernel(
24
                self , parameters: KernelParameters , x: jnp.ndarray , y: jnp.ndarray
          ) -> jnp.ndarray:
25
26
                   Kernel evaluation between a single feature x and a single feature y.
27
28
29
                    parameters: parameters dataclass for the kernel
30
                    x: ndarray of shape (number_of_dimensions,)
y: ndarray of shape (number_of_dimensions,)
               The kernel evaluation. (1, 1)
34
35
36
                raise NotImplementedError
38
          self, parameters: KernelParameters, x: jnp.ndarray, y: jnp.ndarray = None
) -> jnp.ndarray:
          def kernel (
39
40
41
               """ Kernel evaluation for an arbitrary number of x features and y features. Compute k(x, x) if y is None. This method requires the parameters dataclass and is better suited for parameter optimisation.
42
44
45
                     parameters: parameters dataclass for the kernel
                    x: ndarray of shape (number_of_x_features, number_of_dimensions)
y: ndarray of shape (number_of_y_features, number_of_dimensions)
47
48
49
               A gram matrix k(x, y), if y is None then k(x,x). (number_of_x_features, number_of_y_features)
50
51
                # compute k(x, x) if y is None
               if y is None:
y = x
56
               # add dimension when x is 1D, assume the vector is a single feature
               x = jnp.atleast_2d(x)
58
               y = jnp.atleast_2d(y)
59
60
               61
62
63
                return vmap(
lambda x_i: vmap(
65
66
               )(x) (y),
                          lambda y_i: self._kernel(parameters, x_i, y_i),
67
          def __call__(
    self, x: jnp.ndarray, y: jnp.ndarray = None, **parameter_args
) -> jnp.ndarray:
70
71
72
73
74
75
76
               > jnp.ndarray:
""" Kernel evaluation for an arbitrary number of x features and y features.
This method is more user-friendly without the need for a parameter data class.
It wraps the kernel computation with the initial step of constructing the parameter data class from the provided parameter arguments.
77
78
79
                    x: ndarray of shape (number_of_x_features, number_of_dimensions)
y: ndarray of shape (number_of_y_features, number_of_dimensions)
**parameter_args: parameter arguments for the kernel
80
81
83
               A gram matrix k(x, y), if y is None then k(x,x). (number_of_x_features, number_of_y_features).
84
                parameters = self.Parameters(**parameter_args)
86
                return self.kernel(parameters, x, y)
          def diagonal (
89
90
                self,
91
                x: jnp.ndarray,
                y: jnp.ndarray = None,
92
                 **parameter_args ,
          ) -> jnp.ndarray:
```

```
""" Kernel evaluation of only the diagonal terms of the gram matrix.
 96
 97
                        x: ndarray of shape (number_of_x_features, number_of_dimensions)
y: ndarray of shape (number_of_y_features, number_of_dimensions)
**parameter_args: parameter arguments for the kernel
 98
 aa
101
                        A diagonal of gram matrix k(x, y), if y is None then trace(k(x, x)). (number_of_x_features, number_of_y_features)
103
                   \# compute k(x, x) if y is None if y is None:
106
107
108
                        y = x
109
110
                   \# add dimension when x is 1D, assume the vector is a single feature
                  x = jnp.atleast_2d(x)

y = jnp.atleast_2d(y)
113
114
                   assert (
                   x.shape[1] == y.shape[1]
), f"Dimension Mismatch: {x.shape[1]=} != {y.shape[1]=}"
116
117
                   assert (
                  x.shape[0] = y.shape[0]), f"Must have same number of features for diagonal: \{x.shape[0]=\}! = \{y.shape[0]=\}"
120
                   return vmap(
                        lambda x_i, y_i: self._kernel(
parameters=self.Parameters(**parameter_args),
123
124
                               y=y_i ,
126
127
                   )(x, y)
128
             def trace(
130
                   \verb|self|, x: jnp.ndarray|, y: jnp.ndarray| = None, **parameter\_args|
             ) \rightarrow jnp.ndarray: "" Trace of the gram matrix, calculated by summation of the diagonal matrix.
134
                   Args:
                        x: ndarray of shape (number_of_x_features, number_of_dimensions)
y: ndarray of shape (number_of_y_features, number_of_dimensions)
**parameter_args: parameter arguments for the kernel
135
136
                   The trace of the gram matrix k(x, y).
140
141
                   \begin{array}{ll} parameters = self. Parameters (**parameter\_args) \\ \hline return \ jnp.trace (self.kernel (parameters, \ x, \ y)) \end{array}
142
144
145
       class CombinedKernelParameters (KernelParameters):
147
148
149
             Parameters for the Combined Kernel:
151
             log_theta: float
             log_sigma: float log_phi: float
154
             log_eta: float
log_tau: float
156
             log_zeta: float
158
             @property
             def theta(self) -> float:
    return jnp.exp(self.log_theta)
160
161
162
163
             @property
             def sigma(self) -> float:
164
                  return jnp.exp(self.log_sigma)
165
166
             @property
167
            def phi(self) -> float:
    return jnp.exp(self.log_phi)
168
169
             @property
172
173
             def eta(self) -> float:
                  return jnp.exp(self.log_eta)
174
\frac{175}{176}
            @property
def tau(self) -> float:
                  return jnp.exp(self.log_tau)
178
179
             @property
             def zeta(self) -> float:
    return jnp.exp(self.log_zeta)
180
181
182
             @theta.setter
def theta(self, value: float) -> None:
    self.log_theta = jnp.log(value)
183
184
185
186
187
             @sigma.setter
             def sigma(self, value: float) -> None:
189
                   self.log_sigma = jnp.log(value)
190
```

```
191
          @phi.setter
          def phi(self, value: float) -> None:
    self.log_phi = jnp.log(value)
192
193
194
195
          def eta(self , value: float) -> None:
    self.log_eta = jnp.log(value)
196
197
198
          @tau.setter
199
          def tau(self, value: float) -> None:
200
201
               self.log\_tau = jnp.log(value)
202
203
          def zeta(self, value: float) -> None:
    self.log_zeta = jnp.log(value)
204
205
206
207
      class CombinedKernel(Kernel):
208
209
210
          The kernel defined as:  k(x, y) = \frac{2 \sin^2 2 (\exp(-(2 \sin^2 2(\pi - y)/\tan 2)) + \pi^2 * \exp(-(x - y)^2/(2 * \cot^2 2)) }{2 + 2 \cot^2 2 * \det(x - y)} 
211
212
213
          Parameters = CombinedKernelParameters
215
216
          def _kernel(
               self,
218
219
               parameters: CombinedKernelParameters,
          x: jnp.ndarray,
y: jnp.ndarray,
) -> jnp.ndarray:
""" Kernel evaluation between a single feature x and a single feature y.
220
221
223
224
226
                   parameters: parameters dataclass for the Gaussian kernel
227
                   x: ndarray of shape (1,)
y: ndarray of shape (1,)
229
230
               Returns:
               The kernel evaluation.
231
232
233
               return jnp.dot(
234
                   jnp.ones(1),
236
                         (parameters.theta**2)
                         * (
* (
237
238
                                  240
241
243
                             )
244
                        246
249
               )
```

src/models/kernels.py

The Python code for Gaussian Process Regression:

```
from dataclasses import dataclass
from typing import Any, Dict, Tuple
 3
      import jax.numpy as jnp
      import optax
      from jax import grad
from optax import GradientTransformation
10
      from src.models.kernels import Kernel
11
13
14
      @\,d\,a\,t\,a\,c\,l\,a\,s\,s
      class Gaussian Process Parameters:
           Parameters for a Gaussian Process: log_sigma: logarithm of the noise parameter
16
17
                 kernel: parameters for the chosen kernel
19
20
           log_sigma: float
kernel: Dict[str, Any]
21
22
23
24
           def variance(self) -> float:
    return self.sigma**2
25
26
27
28
           @property
29
           def sigma(self) -> float:
30
                 return jnp.exp(self.log_sigma)
           def sigma(self, value: float) -> None:
    self.log_sigma = jnp.log(value)
36
      class Gaussian Process:
38
           A Gaussian measure defined with a kernel, better known as a Gaussian Process.
39
40
41
           Parameters = GaussianProcessParameters
42
           def __init__(self, kernel: Kernel, x: jnp.ndarray, y: jnp.ndarray) -> None:
    """ Initialising requires a kernel and data to condition the distribution.
44
45
47
48
                       kernel: kernel for the Gaussian Process
49
                       x: design matrix (number_of_features, number_of_dimensions)
                 y: response vector (number_of_features, )
50
51
                 self.number_of_train_points = x.shape[0]
                 self.x = x
self.y = y
                  self.kernel = kernel
56
           \textcolor{red}{\textbf{def}} \hspace{0.2cm} \texttt{\_compute\_kxx\_shifted\_cholesky\_decomposition} \hspace{0.1cm} (
           self, parameters
) -> Tuple[jnp.ndarray, bool]:
58
59
60
                 Cholesky decomposition of (kxx + (1/ ^2)*I)
61
62
63
                       parameters: parameters dataclass for the Gaussian Process
                       cholesky_decomposition_kxx_shifted: the cholesky decomposition (number_of_features,
67
             number_of_features)
                 lower\_flag: ^{'}flag \ indicating \ whether \ the \ factor \ is \ in \ the \ lower \ or \ upper \ triangle \ """
68
                 kxx = self.kernel(self.x, **parameters.kernel)
                 kxx_shifted = kxx + parameters.variance * jnp.eye(self.number_of_train_points)
kxx_shifted_cholesky_decomposition , lower_flag = jax.scipy.linalg.cho_factor(
    a=kxx_shifted , lower=True
71
72
73
74
75
                 return kxx_shifted_cholesky_decomposition, lower_flag
76
77
78
           def posterior_distribution (
           self, x: jnp.ndarray, **parameter_args
) -> Tuple[jnp.ndarray, jnp.ndarray]:
    """Compute the posterior distribution for test points x.
    Reference: http://gaussianprocess.org/gpml/chapters/RW2.pdf
79
80
82
83
                       x: test points (number_of_features, number_of_dimensions)
85
                       **parameter\_args:\ parameter\ arguments\ for\ the\ Gaussian\ Process
86
                      mean: the distribution mean (number_of_features, )
covariance: the distribution covariance (number_of_features, number_of_features)
88
89
QΩ
                 parameters = self.Parameters(**parameter.args)
kxy = self.kernel(self.x, x, **parameters.kernel)
kyy = self.kernel(x, **parameters.kernel)
91
```

```
kxx_shifted_cholesky_decomposition,
              lower_flag ,
) = self._compute_kxx_shifted_cholesky_decomposition(parameters)
96
97
98
99
              mean = (
100
                  kxy.T
                  @ jax.scipy.linalg.cho_solve(
                       c_and_lower=(kxx_shifted_cholesky_decomposition, lower_flag), b=self.y
104
              ).reshape(
105
                   -1.
              107
108
109
              return mean, covariance
         112
113
114
                   **parameter_args: parameter arguments for the Gaussian Process
118
              Returns:
119
              The negative log likelihood.
122
              parameters = self.Parameters(**parameter_args)
124
                   kxx_shifted_cholesky_decomposition,
125
                   lower_flag
126
              ) = self._compute_kxx_shifted_cholesky_decomposition(parameters)
              negative_log_likelihood = -(
                  -0.5
* (
130
                        self.y.T
                       @ jax.scipy.linalg.cho_solve(
    c_and_lower=(kxx_shifted_cholesky_decomposition, lower_flag),
132
                            b=self.y,
136
                  / jnp.trace(jnp.log(kxx_shifted_cholesky_decomposition))
- (self.number_of_train_points / 2) * jnp.log(2 * jnp.pi)
138
139
140
              return negative_log_likelihood
141
         def -compute-gradient(self, **parameter-args) -> Dict[str, Any]:
    """ Calculate the gradient of the posterior negative log likelihood with respect to the parameters.
143
144
              Args:
146
                   **parameter_args: parameter arguments for the Gaussian Process
147
              A dictionary of the gradients for each parameter argument.
149
150
              gradients = grad(
              lambda params: self.posterior_negative_log_likelihood(**params))(parameter_args)
153
154
              return gradients
157
              optimizer: GradientTransformation,
              number_of_training_iterations: int,
160
              **parameter_args
         ) -> GaussianProcessParameters:
161
               ""Train the parameters for a Gaussian Process by optimising the negative log likelihood.
163
164
                  optimizer: jax optimizer object number_of_training_iterations: number_of_training_iterations: number of iterations to perform the optimizer
165
                   **parameter_args: parameter arguments for the Gaussian Process
167
168
              Returns:
              A parameters dataclass containing the optimised parameters. ""
171
172
              opt_state = optimizer.init(parameter_args)
for _ in range(number_of_training_iterations):
\frac{174}{175}
                   gradients = self._compute_gradient(**parameter_args)
updates, opt_state = optimizer.update(gradients, opt_state)
                   parameter_args = optax.apply_updates(parameter_args, updates)
              return self.Parameters(**parameter_args)
```

src/models/gaussian_process_regression.py

The rest of the Python code for question 2:

```
from dataclasses import asdict, fields
 3
      import dataframe_image as dfi
      import jax
      import jax.numpy as jnp
import matplotlib.pyplot as plt
      import numpy as np
      import optax
import pandas as pd
import scipy
10
11
      from src.models.bayesian_linear_regression import (
13
14
            {\tt Linear Regression Parameters} \ ,
            compute_linear_regression_posterior,
16
17
      from src.models.gaussian_process_regression import (
            Gaussian Process,
Gaussian Process Parameters,
19
20
21
22
      from src.models.kernels import CombinedKernel, CombinedKernelParameters
23
24
25
      jax.config.update("jax_enable_x64", True)
26
      def construct_design_matrix(t: np.ndarray):
27
28
            return np. stack ((t, np. ones(t. shape)), axis=1).T
29
30
      def a(
31
           t: np.ndarray,
32
            y: np.ndarray,
            sigma: float
34
            prior_linear_regression_parameters: LinearRegressionParameters,
35
            save_path: str,
      ) -> LinearRegressionParameters
36
37
           x = construct_design_matrix(t)
38
            prior_theta = Theta(
                 linear_regression_parameters=prior_linear_regression_parameters,
39
40
                  sigma=sigma,
41
            ,
posterior_linear_regression_parameters = compute_linear_regression_posterior(
42
                 х,
44
45
                  prior_linear_regression_parameters ,
                  residuals_precision=prior_theta.precision,
47
            df_mean = pd.DataFrame(
48
49
                  posterior_linear_regression_parameters.mean, columns=["value"]
50
            df-mean.index = ["a", "b"]
df-mean = pd.concat([df-mean], keys=["parameters"])
dfi.export(df-mean, save-path + "-mean.png")
51
55
            df_covariance = pd.DataFrame(
                  posterior_linear_regression_parameters.covariance, columns=["a", "b"]
56
            df_covariance.index = ["a", "b"]
df_covariance = pd.concat([df_covariance], keys=["parameters"])
df_covariance = pd.concat([df_covariance.T], keys=["parameters"])
dfi.export(df_covariance, save-path + "-covariance.png")
return posterior_linear_regression_parameters
58
59
60
61
62
63
65
      def b(
66
            t_year: np.ndarray,
67
           t: np.ndarray,
y: np.ndarray,
            j. np.ndaray,
linear_regression_parameters: LinearRegressionParameters,
error_mean: float,
error_variance: float,
70
71
72
73
74
75
76
            save\_path ,
      ) -> None:
                 construct_design_matrix(t)
            residuals = y - linear\_regression\_parameters.predict(x) \\ plt.plot(t\_year.reshape(-1), residuals.reshape(-1))
            plt.ylabel("date (decimal year)")
plt.ylabel("residual")
plt.title("2b: g_obs(t)")
plt.savefig(save_path + "-residuals-timeseries")
77
78
79
80
81
            plt.close()
            \begin{array}{lll} count \;,\; bins \;=\; np.\, histogram \, (\, residuals \;,\; bins = 100,\; density = True) \\ plt.\, bar \, (\, bins \, [1:] \;,\; count \;,\; label = "\, residuals "\,) \end{array}
83
84
                  bins[1:].
86
                 scipy.stats.norm.pdf(bins[1:], loc=error_mean, scale=error_variance), color="red", label="e(t)",
87
89
90
            plt.xlabel("residual bin")
plt.ylabel("density")
plt.title("2b: Residuals Density")
91
92
            plt.legend()
```

```
plt.savefig(save_path + "-residuals-density-estimation")
97
 98
 aa
      def c(
           kernel: CombinedKernel.
101
            kernel-parameters: CombinedKernelParameters,
            log_theta_range: np.ndarray,
           t: np.ndarray,
number_of_samples: int,
103
           save_path: str,
      ) -> None:
gram = kernel(t, **asdict(kernel_parameters))
106
107
           plt.imshow(gram)
plt.xlabel("t")
plt.ylabel("t")
108
109
110
           plt.title("Gram Matrix (Prior)")
plt.savefig(save_path + "-gram-matrix")
113
           plt.close()
114
            for _ in range(number_of_samples):
116
                 plt.plot(
                      np.random.multivariate_normal(
117
                           jnp.zeros(gram.shape[0]), gram, size=1
                      ). reshape(-1)
120
           plt.xlabel("t")
plt.ylabel("f_GP(t)")
plt.title("Samples from Gaussian Process Prior")
plt.savefig(save_path + "-samples")
123
124
            plt.close()
126
127
            fig_samples, ax_samples = plt.subplots(
                 len(fields(kernel_parameters.__class__)),
len(log_theta_range),
128
129
                 figsize=(
130
                      len(log_theta_range) * 2,
len(fields(kernel_parameters.__class__)) * 2,
                 frameon=False,
134
           for i, field in enumerate(fields(kernel-parameters.--class--)):
    default_value = getattr(kernel-parameters, field.name)
    for j, log_value in enumerate(log_theta_range):
136
138
                      setattr(kernel-parameters, field.name, log_value)
gram = kernel(t, **asdict(kernel-parameters))
ax_samples[i][j].plot(
140
141
                           np.random.multivariate_normal(
142
                                jnp.zeros(gram.shape[0]), gram, size=1
143
                           ). reshape(-1),
144
145
                      ax_samples[i][j].set_title(
                           f"{field.name.strip('log_')}={np.round(np.exp(log_value), 2)}"
147
148
149
                 setattr (kernel_parameters, field.name, default_value)
           plt.tight_layout()
plt.savefig(save_path + f"-parameter-samples", bbox_inches="tight")
151
            plt.close(fig_samples)
154
            fig_gram , ax_gram = plt.subplots(
                 len(fields(kernel_parameters.__class__)),
len(log_theta_range),
156
                 figsize=(
158
                      len(log-theta-range) * 2,
                      len (fields (kernel_parameters.__class__)) * 2,
161
                 frameon=False
           163
164
165
166
167
168
169
172
                 setattr(kernel_parameters, field.name, default_value)
           plt.tight_layout()
plt.savefig(save_path + f"-parameter-grams", bbox_inches="tight")
            plt.close(fig_gram)
176
      def f(
    t_train: np.ndarray,
178
179
180
            y_train: np.ndarray,
           t_test: np.ndarray,
min_year: float,
181
182
           prior_linear_regression_parameters: LinearRegressionParameters, linear_regression_sigma: float, kernel: CombinedKernel,
183
184
           {\tt gaussian\_process\_parameters}: \ {\tt GaussianProcessParameters} \ , \\ {\tt learning\_rate}: \ {\tt float} \ , \\
186
187
           number_of_iterations: int,
189
           save_path: str,
190
      ) -> None:
```

```
191
          # Train Bayesian Linear Regression
192
           x_train = construct_design_matrix(t_train)
          prior_theta = Theta(
    linear_regression_parameters=prior_linear_regression_parameters ,
193
194
195
               sigma=linear_regression_sigma,
196
197
           ,
posterior_linear_regression_parameters = compute_linear_regression_posterior(
198
199
               v_train
200
               prior_linear_regression_parameters ,
201
               residuals_precision=prior_theta.precision,
202
203
204
          residuals = y\_train - posterior\_linear\_regression\_parameters.predict(x\_train)\\ gaussian\_process = GaussianProcess(
205
206
               kernel, t_train.reshape(-1, 1), residuals.reshape(-1)
207
208
209
          # Prediction
           x_test = construct_design_matrix(t_test)
210
211
          linear_prediction = posterior_linear_regression_parameters.predict(x_test).reshape(
212
213
          \label{eq:continuous_prediction} \begin{subarray}{ll} mean\_prediction , covariance\_prediction = gaussian\_process\_posterior\_distribution ( \\ t\_test\_reshape(-1, 1) , **asdict(gaussian\_process\_parameters) \end{subarray}
214
216
218
          # Plot
          plt figure (figsize = (7, 7))
219
          plt. lighte (light2e = (1, plt. scatter ( t_train + min_year
220
222
               y_{train.reshape(-1)},
223
                s=2.
               color="blue"
224
               label="historical data",
226
           plt.plot(
               t_test + min_year,
               {\tt linear\_prediction} \ + \ {\tt mean\_prediction} \ ,
230
               color="gray",
label="prediction",
231
233
           plt.fill_between(
234
               t_test + min_year,
               linear_prediction
236
               + mean_prediction
                 1 * jnp.sqrt(jnp.diagonal(covariance-prediction)),
237
               linear_prediction
238
               + mean_prediction
240
               + 1 * jnp.sqrt(jnp.diagonal(covariance_prediction)),
               facecolor = (0.8, 0.8, 0.8), label="error bound (one stdev)",
241
243
          plt.xlabel("date (decimal year)")
plt.ylabel("parts per million")
plt.title("Global Mean CO_2 Concentration Prediction (Untrained Hyperparameters)")
244
246
           plt.legend()
248
           plt.tight_layout()
           plt.savefig(save_path + "-extrapolation-untrained", bbox_inches="tight")
249
250
           plt.close()
251
252
          df_parameters = pd.DataFrame(
               [
254
                         x.strip("log_") + " (kernel)",
255
                         np.exp(gaussian_process_parameters.kernel[x]),
257
258
                    for x in gaussian_process_parameters.kernel.keys()
259
               | [ "sigma", float(gaussian_process_parameters.sigma)]],
columns=["parameter", "value"],
260
261
262
          263
264
265
          # Train Gaussian Process Regression (Hyperparameter Tune)
266
          optimizer = optax.adam(learning_rate)
268
          {\tt gaussian\_process\_parameters} \ = \ {\tt gaussian\_process.train} \ (
269
               {\tt optimizer}\ ,\ {\tt number\_of\_iterations}\ ,\ ** {\tt asdict}\ ({\tt gaussian\_process\_parameters}\ )
270
271
           df_parameters = pd.DataFrame(
272
273
                        x.strip("log_") + " (kernel)",
np.exp(gaussian_process_parameters.kernel[x]),
274
276
277
                    for x in gaussian_process_parameters.kernel.keys()
279
                  \hbox{\tt [["sigma", float(gaussian\_process\_parameters.sigma)]],}\\
               columns = ["parameter", "value"
280
281
          282
283
285
          # Prediction
286
          x_test = construct_design_matrix(t_test)
```

```
287
            linear\_prediction \ = \ posterior\_linear\_regression\_parameters.predict(x\_test).reshape(
288
289
            290
291
292
293
294
            # Plot
            295
296
297
298
299
                 color="blue",
label="historical data",
300
301
302
            plt.plot(
    t_test + min_year,
    linear_prediction + mean_prediction,
    color="gray",
    label="prediction",
303
304
305
306
307
308
            plt.fill_between (
309
                 t_test + min_year,
linear_prediction
310
311
312
                 + mean_prediction
313
                    1 * jnp.sqrt(jnp.diagonal(covariance_prediction)),
                 linear_prediction
314
315
                 + mean_prediction
+ 1 * jnp.sqrt(jnp.diagonal(covariance_prediction)),
316
                 \begin{array}{lll} \text{facecolor} = (0.8 \,,\, 0.8 \,,\, 0.8) \,, \\ \text{label} = \text{"error bound (one stdev)"} \,, \end{array}
317
318
319
            plt.xlabel("date (decimal year)")
plt.ylabel("parts per million")
plt.title("Global Mean CO_2 Concentration Prediction (Trained Hyperparameters)")
plt.legend()
plt.tight.layout()
320
321
322
323
            plt.savefig(save_path + "-extrapolation-trained", bbox_inches="tight") plt.close()
325
326
```

src/solutions/q2.py

Question 3

(a)

The free energy is can be calculated as:

$$\mathcal{F}(q, \theta) = \langle \log P(\mathbf{x}, \mathbf{s} | \theta) \rangle_{q(\mathbf{s})} + H[Q(\mathbf{s})]$$

Knowing,

$$\log P(\mathbf{x}, \mathbf{s}|\theta) = \log P(\mathbf{x}|\mathbf{s}, \theta) + \log P(\mathbf{s}|\theta)$$

we can write:

$$\mathcal{F}(Q, \theta) = \langle \log P(\mathbf{x}|\mathbf{s}, \theta) \rangle_{q(\mathbf{s})} + \langle \log P(\mathbf{s}|\theta) \rangle_{q(\mathbf{s})} + H[q(\mathbf{s})]$$

Moreover, our mean field approximation:

$$q(\mathbf{s}) = \prod_{i=1}^{K} q_i(s_i)$$

where $q_i(s_i) = \lambda_i^{s_i} (1 - \lambda_i)^{(1-s_i)}$.

To compute the first term:

$$P(\mathbf{x}|\mathbf{s}, \theta) = \mathcal{N}\left(\sum_{i=1}^{K} s_i \mu_i, \sigma^2 \mathbf{I}\right)$$

substituting the appropriate terms:

$$P(\mathbf{x}|\mathbf{s}, \theta) = 2\pi^{-\frac{d}{2}} |\sigma^2 \mathbf{I}|^{-\frac{1}{2}} \exp\left(-\frac{1}{2} \left(\mathbf{x} - \sum_{i=1}^K s_i \mu_i\right)^T \frac{1}{\sigma^2} \mathbf{I} \left(\mathbf{x} - \sum_{i=1}^K s_i \mu_i\right)\right)$$

with d being the number of dimensions.

Taking the logarithm:

$$\log P(\mathbf{x}|\mathbf{s}, \theta) = -\frac{d}{2}\log(2\pi\sigma^2) - \frac{1}{2\sigma^2} \left(\mathbf{x}^T\mathbf{x} - 2\mathbf{x}^T\sum_{i=1}^K s_i\mu_i + \sum_{i=1}^K\sum_{j=1}^K s_is_j\mu_i^T\mu_j\right)$$

The expectation distributed to the relevant terms:

$$\langle \log P(\mathbf{x}|\mathbf{s}, \theta) \rangle_{q(\mathbf{s})} = -\frac{d}{2} \log(2\pi\sigma^2) - \frac{1}{2\sigma^2} \left(\mathbf{x}^T \mathbf{x} - 2\mathbf{x}^T \sum_{i=1}^K \langle s_i \rangle_{q_i(s_i)} \mu_i + \sum_{i=1}^K \sum_{j=1}^K \langle s_i s_j \rangle_{q_i(s_i)q_j(s_j)} \mu_i^T \mu_j \right)$$

Evaluating the expectations:

$$\langle \log P(\mathbf{x}|\mathbf{s}, \theta) \rangle_{q(\mathbf{s})} = -\frac{d}{2} \log(2\pi\sigma^2) - \frac{1}{2\sigma^2} \left(\mathbf{x}^T \mathbf{x} - 2\mathbf{x}^T \sum_{i=1}^K \lambda_i \mu_i + \sum_{i=1}^K \sum_{j=1, j \neq i}^K \lambda_i \lambda_j \mu_i^T \mu_j + \sum_{i=1}^K \lambda_i \mu_i^T \mu_i \right)$$

where $\langle s_i s_i \rangle_{q_i(s_i)} = \langle s_i \rangle_{q_i(s_i)}$ because $s_i \in \{0, 1\}$.

To compute the second term:

$$P(\mathbf{s}|\theta) = \prod_{i=1}^{K} \pi_i^{s_i} (1 - \pi_i)^{(1-s_i)}$$

Taking the logarithm:

$$\log P(\mathbf{s}|\theta) = \sum_{i=1}^{K} s_i \log \pi_i + (1 - s_i) \log(1 - \pi_i)$$

The expectation distributed to the relevant terms:

$$\langle \log P(\mathbf{s}|\theta) \rangle_{q(\mathbf{s})} = \sum_{i=1}^{K} \langle s_i \rangle_{q_i(s_i)} \log \pi_i + (1 - \langle s_i \rangle_{q_i(s_i)}) \log(1 - \pi_i)$$

Evaluating the expectations:

$$\langle \log P(\mathbf{s}|\theta) \rangle_{q(\mathbf{s})} = \sum_{i=1}^{K} \lambda_i \log \pi_i + (1 - \lambda_i) \log(1 - \pi_i)$$

To compute the third term, we use the mean field factorisation:

$$H\left[q(\mathbf{s})\right] = \sum_{i=1}^{K} H\left[q_i(s_i)\right]$$

Thus,

$$H[q(\mathbf{s})] = -\sum_{i=1}^{K} \sum_{s_i \in \{0,1\}} q_i(s_i) \log q_i(s_i)$$

Substituting the appropriate values:

$$H[q(\mathbf{s})] = -\sum_{i=1}^{K} \lambda_i \log \lambda_i + (1 - \lambda_i) \log(1 - \lambda_i)$$

Combining, we have our free energy expression:

$$\mathcal{F}(q,\theta) = \frac{\frac{-d}{2}\log(2\pi\sigma^{2}) - \frac{1}{2\sigma^{2}}\left(\mathbf{x}^{T}\mathbf{x} - 2\mathbf{x}^{T}\sum_{i=1}^{K}\lambda_{i}\mu_{i} + \sum_{i=1}^{K}\sum_{j=1,j\neq i}^{K}\lambda_{i}\lambda_{j}\mu_{i}^{T}\mu_{j} + \sum_{i=1}^{K}\lambda_{i}\mu_{i}^{T}\mu_{i}\right) + \sum_{i=1}^{K}\lambda_{i}\log\pi_{i} + (1-\lambda_{i})\log(1-\pi_{i}) - \sum_{i=1}^{K}\lambda_{i}\log\lambda_{i} + (1-\lambda_{i})\log(1-\lambda_{i})$$

To derive the partial update for $q_i(s_i)$ we take the variational derivative of the Lagrangian, enforcing the normalisation of q_i :

$$\frac{\partial}{\partial q_i} \left(\mathcal{F}(q, \theta) + \lambda^{LG} \int q_i - 1 \right) = \langle \log P(\mathbf{x}, \mathbf{s} | \theta) \rangle_{\prod_{j \neq i} q_j(s_j)} - \log q_i(s_i) - 1 + \lambda^{LG}$$

where λ^{LG} is the Lagrange multiplier.

Setting this to zero we can solve for the λ_i that maximises the free energy:

$$\log q_i(s_i) = \langle \log P(\mathbf{x}, \mathbf{s} | \theta) \rangle_{\prod_{j \neq i} q_j(s_j)} - 1 + \lambda^{LG}$$

Similar to our free energy derivation:

$$\langle \log P(\mathbf{x}|\mathbf{s}, \theta) \rangle_{\prod_{j \neq i} q_j(s_j)} \propto -\frac{1}{2\sigma^2} \left(\mathbf{x}^T \mathbf{x} - 2\mathbf{x}^T \sum_{k=1}^K \langle s_k \rangle_{\prod_{j \neq i} q_j(s_j)} \mu_i + \sum_{k=1}^K \sum_{j=1}^K \langle s_k s_j \rangle_{\prod_{j \neq i} q_j(s_j)} \right)$$

and

$$\langle \log P(\mathbf{s}|\theta) \rangle_{\prod_{j \neq i} q_j(s_j)} = \sum_{k=1}^K \langle s_k \rangle_{\prod_{j \neq i} q_j(s_j)} \log \pi_k + (1 - \langle s_k \rangle_{\prod_{j \neq i} q_j(s_j)}) \log(1 - \pi_k)$$

We can write:

$$\log q_i(s_i) \propto \log P(\mathbf{x}|\mathbf{s}, \theta)_{\prod_{j \neq i} q_j(s_j)} + \langle \log P(\mathbf{s}|\theta) \rangle_{\prod_{j \neq i} q_j(s_j)}$$

Substituting the relevant terms:

$$\log q_i(s_i) \propto -\frac{1}{2\sigma^2} \left(-2s_i \mathbf{x}^T \mu_i + s_i s_i \mu_i^T \mu_i + 2 \sum_{j=1, j \neq i}^K s_i \lambda_j \mu_i^T \mu_j \right) + s_i \log \pi_i + (1 - s_i) \log(1 - \pi_i)$$

Knowing $\log q_i(s_i) = s_i \log \lambda_i + (1 - s_i) \log(1 - \lambda_i)$:

$$\log q_i(s_i) \propto s_i \log \frac{\lambda_i}{1 - \lambda_i}$$

Thus,

$$s_i \log \frac{\lambda_i}{1 - \lambda_i} \propto -\frac{1}{2\sigma^2} \left(-2s_i \mathbf{x}^T \mu_i + s_i s_i \mu_i^T \mu_i + 2 \sum_{j=1, j \neq i}^K s_i \lambda_j \mu_i^T \mu_j \right) + s_i \log \frac{\pi_i}{1 - \pi_i}$$

Also, because $s_i \in \{0, 1\}$ we know that $s_i^2 = s_i$:

$$s_i \log \frac{\lambda_i}{1 - \lambda_i} \propto -\frac{1}{2\sigma^2} \left(-2s_i \mathbf{x}^T \mu_i + s_i \mu_i^T \mu_i + 2 \sum_{j=1, j \neq i}^K s_i \lambda_j \mu_i^T \mu_j \right) + s_i \log \frac{\pi_i}{1 - \pi_i}$$

Because we have only kept terms with s_i , this is an equality:

$$s_i \log \frac{\lambda_i}{1 - \lambda_i} = \frac{s_i \mu_i^T}{2\sigma^2} \left(2\mathbf{x} - \mu_i - 2\sum_{j=1, j \neq i}^K \lambda_j \mu_j \right) + s_i \log \frac{\pi_i}{1 - \pi_i}$$

Solving for λ_i :

$$\lambda_i = \frac{1}{1 + \exp\left[-\left(\frac{\mu_i^T}{\sigma^2} \left(\mathbf{x} - \frac{\mu_i}{2} - \sum_{j=1, j \neq i}^K \lambda_j \mu_j\right) + \log\frac{\pi_i}{1 - \pi_i}\right)\right]}$$

we have our partial update.

(b)

The provided derivations for the M step of the mean parameter μ :

$$\mu = \left(\left\langle \mathbf{s} \mathbf{s}^T \right\rangle_{q(\mathbf{s})} \right)^{-1} \left\langle \mathbf{s} \right\rangle_{q(\mathbf{s})} \mathbf{x}$$

where $\mu \in \mathbb{R}^{K \times D}$, $\mathbf{s} \in \mathbb{R}^{K \times N}$, and $\mathbf{x} \in \mathbb{R}^{N \times D}$.

This mimics the least squares solution:

$$\hat{\beta} = (\mathbf{X}\mathbf{X}^T)\mathbf{X}\mathbf{Y}$$

for the linear regression problem $\mathbf{Y} = \mathbf{X}^T \boldsymbol{\beta}$ where $\boldsymbol{\beta}$ corresponds to the mean parameters $\boldsymbol{\mu}$, the design matrix \mathbf{X} corresponds to the input \mathbf{s} and the response Y corresponds to the image pixels denoted \mathbf{x} . This makes sense because our resulting images \mathbf{x} are modeled as linear combinations of features $\boldsymbol{\mu}$, weighted by \mathbf{s} .

(c)

The computational complexity of the implemented M step function can be broken down for each parameter:

 μ : - The inversion ESS⁻¹ where ESS $\in \mathbb{R}^{K \times K}$ is $\mathcal{O}(K^3)$

- The dot product $ESS^{-1}ES^T$ where $ESS^{-1} \in \mathbb{R}^{K \times K}$ and $ES \in \mathbb{R}^{N \times K}$ is $\mathcal{O}(K^2N)$

- The dot product $(ESS^{-1}ES^T)\mathbf{x}$ where $(ESS^{-1}ES^T) \in \mathbb{R}^{K \times N}$ and $\mathbf{x} \in \mathbb{R}^{N \times D}$ is $\mathcal{O}(KND)$

 σ : - The dot product $(\mathbf{x}^T\mathbf{x})$ where $\mathbf{x} \in \mathbb{R}^{N \times D}$ is $\mathcal{O}(D^2N)$

- The dot product $\mu^T \mu$ where $\mu \in \mathbb{R}^{D \times K}$ is $\mathcal{O}(K^2 D)$

– The dot product $(\mu^T \mu)$ ESS where $\mu^T \mu \in \mathbb{R}^{K \times K}$ and ESS $\in \mathbb{R}^{K \times K}$ is $\mathcal{O}(K^3)$

 π : — The mean operation for ES $\in \mathbb{R}^{N \times K}$ along the first dimension is $\mathcal{O}(NK)$

Thus, the computational complexity of the M step is $\mathcal{O}(K^3 + K^2N + KND + D^2N + K^2D)$ where we do not assume that any of N, K, or D is large compared to the others.

(d)

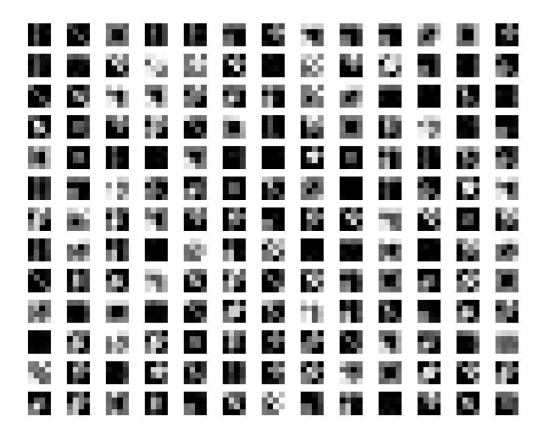


Figure 13: Images generated by randomly combined features with noise Examining the generated images, we can see eight features:

- (1) a cross
- (2) a border
- (3) a two by two square in the middle
- (4) a two by two square in the bottom left corner
- (5) a diagonal from top left to bottom right
- (6) a vertical line in the second column
- (7) a vertical line in the fourth column
- (8) a a horizontal line in the first row

Factor analysis assumes a model:

$$\mathbf{x} = \mathbf{W}\mathbf{s} + \epsilon$$

where $\epsilon \sim \mathcal{N}(\mu_{\epsilon}, \Sigma_{\epsilon})$ and $\mathbf{s} \sim \mathcal{N}(\mu_{\mathbf{s}}, \Sigma_{\mathbf{s}})$. Factor analysis would be inappropriate for this data because the our latent variables are binary (i.e. whether or not a feature is present) and not Gaussians. Moreover, the presence of each feature is independent of the presence of another which is not enforced in this model with a covariance matrix that might not be diagonal.

A mixture of Gaussians assumes as model:

$$\mathbf{x} = \sum_{k=1}^{K} \pi_k \mu_k + \epsilon$$

where $\epsilon \sim \mathcal{N}(0, \Sigma_{\epsilon})$. This also wouldn't be appropriate because each mixture component (feature) is assumed to have some covariance, whereas our mixtures are defined as binary vectors (a cross, a border, etc) and added together before adding some noise.

The independent component analysis assumes a model:

$$\mathbf{x} = \mathbf{W}\mathbf{s} + \epsilon$$

where $\epsilon \sim \mathcal{N}(0, \sigma^2 \mathbf{I})$ and $p(\mathbf{s}) = \prod_{k=1}^K p(s_k)$. This is appropriate for our data because we are linearly combining different features and then adding noise.

Thus, it would be expected that ICA does a good job modelling this data while factor analysis and mixture of Gaussians would not.

(e)

We can plot the free energy to make sure it increases each iteration:

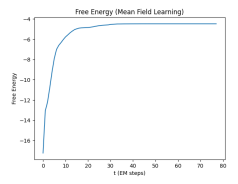


Figure 14: Free Energy

(f)

The initialised features:



Figure 15: Initial Latent Factors

The features learned by the algorithm:

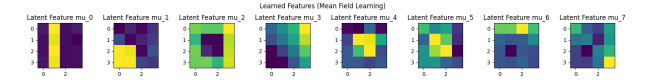


Figure 16: Learned Latent Factors

We can see that it has learned some of previously identified features, such as the vertical line in the second column, the two by two square in the bottom left corner, the border, and the a two by two square in the middle. The other features seem to be some linear combination of two or more features, such as μ_4 which looks like a combination of the cross and two by two square in the middle.

A possible way to improve our algorithm is reinitialising our algorithm a few times to find better potential convergence results (i.e. choose model with best free energy). Another way to improve the algorithm could be to increase the K, although it may learn some duplicate features, there is also a higher chance of capturing all the features. We can visualise this:

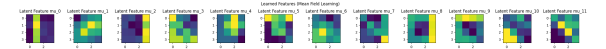


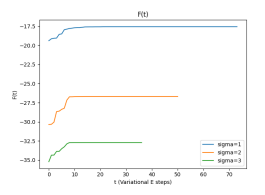
Figure 17: Increasing Number of Latent Factors

Here we can identify a few more features such as the vertical line in the fourth column the cross, and some of the diagonal feature in μ_7 .

When implementing the algorithm, the mean field parameters were initialised randomly, each independently from a uniform distribution. However π , σ , and μ by running the maximisation step using the randomly initialised mean field parameters. K was set to eight, after visually identifying eight features in part d.

(g)

Plotting the free energy at each partial expectation step of the variational approximation for different σ 's:



log(F(t)-F(t-1)

sigma=1
sigma=2
sigma=3

-10

-25

-30

10 20 30 40 50 60 70

Figure 18: Free energy vs σ

Figure 19: Free energy convergence vs σ

We know that our free energy is a upper bounded on the log likelihood:

$$\log P(\mathcal{X}|\theta) \ge \mathcal{F}(q,\theta)$$

In the variational expectation step, $\log P(\mathcal{X}|\theta)$ is fixed and we adjust our approximation q to reach this upper bound. We know that σ quantifies the noise of \mathbf{x} , thus a higher σ means a wider spread in our distribution $\log P(\mathcal{X}|\theta)$, meaning we are reducing our upper bound for $\mathcal{F}(q,\theta)$. As such, we can see in the plot for free energy above that when σ is increased, our free energy converges to a lower value, due to being bounded above by a lower log-likelihood. Moreover, by reducing the upper bound, we see in the plot of $\log(F(t) - F(t-1))$ that our free energy is able to converge faster. Because we have reduced the upper bound by increasing σ , our free energy can reach this upper bound faster.

The Python code for the binary latent factor model:

```
from typing import TYPE_CHECKING, Tuple
 3
     import numpy as np
     from demo_code.MStep import m_step from src.models.binary_latent_factor_approximations.abstract_binary_latent_factor_approximation import (
          AbstractBinaryLatentFactorApproximation,
           {\tt src.models.binary\_latent\_factor\_models.abstract\_binary\_latent\_factor\_model \ import \ (
10
          AbstractBinaryLatentFactorModel\;,
11
12
13
14
     class BinaryLatentFactorModel(AbstractBinaryLatentFactorModel):
          def __init__(
self,
15
16
17
              mu: np.ndarray,
               sigma: float
19
               pi: np.ndarray,
20
21
22
23
               :param mu: matrix of means (number_of_dimensions, number_of_latent_variables)
               :param sigma: Gaussian noise parameter :param pi: vector of priors (1, number_of_latent_variables)
24
25
26
27
               s\,e\,l\,f\,\,.\, \lrcorner m\,u\,\,=\,\,mu
28
               self._sigma = sigma
self._pi = pi
29
30
          @property
          def mu(self):
               return self._mu
34
35
          @mu.setter
          def mu(self, value):
    self._mu = value
36
38
39
          @property
          def sigma(self):
40
41
               return self._sigma
42
          @sigma.setter
          def sigma(self, value):
44
45
               self._sigma = value
          @property
def pi(self):
    return self._pi
47
48
49
50
51
          @pi.setter
          def pi(self, value):
    self._pi = value
          def variance(self) -> float:
    return self.sigma**2
56
58
59
          @staticmethod
60
          {\tt def} \ \ {\tt calculate\_maximisation\_parameters} \, (
               x: np.ndarray,
binary_latent_factor_approximation: AbstractBinaryLatentFactorApproximation,
61
62
          ) -> Tuple[np.ndarray, float, np.ndarray]:
               return m_step(
65
                    x=x,
66
                    es = binary\_latent\_factor\_approximation.expectation\_s \;,
67
                    ess=binary_latent_factor_approximation.expectation_ss ,
70
71
72
          def maximisation_step (
               x: np.ndarray,
73
74
75
76
               binary\_latent\_factor\_approximation: \ AbstractBinaryLatentFactorApproximation \ ,
              mu, sigma, pi = self.calculate_maximisation_parameters(
                    x, binary_latent_factor_approximation
               self.mu = mu
               self.sigma = sigma
80
               self.pi = pi
81
83
     def init_binary_latent_factor_model(
84
          x \colon \text{ np.ndarray} \ , \\ binary\_latent\_factor\_approximation : \ AbstractBinaryLatentFactorApproximation \ , \\
86
     ) -> BinaryLatentFactorModel:
87
          Initialise binary latent factor model by running a maximisation step with the parameters of the
          binary latent factor approximation
89
90
          : param \ x: \ data \ matrix \ (number\_of\_points \ , \ number\_of\_dimensions) \\ : param \ binary\_latent\_factor\_approximation: \ a \ binary\_latent\_factor\_approximation
91
92
          return: an initialised binary latent factor model
```

 $src/models/binary_latent_factor_models/binary_latent_factor_model.py$

The Python code for mean field learning:

```
from typing import List
 3
     import numpy as np
     from src.models.binary_latent_factor_approximations.abstract_binary_latent_factor_approximation import (
          AbstractBinaryLatentFactorApproximation,
     from src.models.binary_latent_factor_models.binary_latent_factor_model import (
          AbstractBinaryLatentFactorModel,
10
11
13
14
     {\bf class} \quad {\bf Mean Field Approximation (Abstract Binary Latent Factor Approximation):}
          def __init__(
               self, lambda_matrix: np.ndarray, max_steps: int, convergence_criterion: float
16
17
               self._lambda_matrix = lambda_matrix
               self.max_steps = max_steps
self.convergence_criterion = convergence_criterion
19
20
21
22
          def lambda_matrix(self) -> np.ndarray:
23
24
               lambda_matrix: parameters variational approximation (number_of_points, number_of_latent_variables)
25
26
               return self._lambda_matrix
27
28
          @lambda_matrix.setter
29
          def lambda_matrix(self, value):
30
               self._lambda_matrix = value
          {\tt def\ lambda\_matrix\_exclude(self\ ,\ exclude\_latent\_index:\ int)\ ->\ np.\,ndarray:}
               # (number_of_points, number_of_latent_variables-1)
return np.concatenate(
34
35
                         self.lambda_matrix[:, :exclude_latent_index],
self.lambda_matrix[:, exclude_latent_index + 1 :],
36
38
39
                     axis=1.
41
          def _partial_expectation_step(
42
               self,
44
               x: np.ndarray
               binary_latent_factor_model: AbstractBinaryLatentFactorModel,
45
               latent_factor: int,
          ) -> np.ndarray:
""" Partial Variational E step for factor i for all data points
47
48
49
               :param x: data matrix (number_of_points, number_of_dimensions)
:param binary_latent_factor_model: a binary_latent_factor_model
:param latent_factor: latent factor to compute partial update
:return: lambda_vector: new lambda parameters for the latent factor (number_of_points, 1)
"""
50
51
               lambda_matrix_excluded = self.lambda_matrix_exclude(latent_factor)
56
               mu\_excluded \ = \ binary\_latent\_factor\_model. \, mu\_exclude \, (\, latent\_factor \, )
58
               \verb|mu_latent = binary_latent_factor_model.mu[:, latent_factor]|
59
               # (number_of_points, 1)
partial_expectation_log_p_x_given_s_theta_proportion = (
61
                    \verb|binary_latent_factor_model.precision|
62
                         63
65
66
67
                    @ mu_latent # (number_of_dimensions, 1)
               )
70
71
72
73
74
75
76
               # (1, 1)
partial_expectation_log_p_s_given_theta_proportion = np.log(
    binary_latent_factor_model.pi[0, latent_factor]
    / (1 - binary_latent_factor_model.pi[0, latent_factor])
               partial_expectation_log_p_x_s_given_theta_proportion = (
    partial_expectation_log_p_x_given_s_theta_proportion
80
                    + partial_expectation_log_p_s_given_theta_proportion
81
83
84
               # (number_of_points, 1)
               lambda_vector = 1 / (
    1 + np.exp(-partial_expectation_log_p_x_s_given_theta_proportion)
86
               [lambda\_vector[lambda\_vector == 0] = 1e-10
               lambda\_vector[lambda\_vector == 1] = 1 - 1e-10
89
               return lambda_vector
90
91
          def variational_expectation_step (
          self , x: np.ndarray , binary_latent_factor_model: AbstractBinaryLatentFactorModel
) -> List[float]:
92
                   'Variational E step
```

```
96
97
                   :param binary_latent_factor_model: a binary_latent_factor_model
                   :param x: data matrix (number_of_points, number_of_dimensions)
 98
                  99
101
102
103
105
                                    self.compute_free_energy(x, binary_latent_factor_model)
106
107
108
                                \begin{array}{ll} \text{if} & \text{free\_energy} \, [\, -1] \,\, - \,\, \text{free\_energy} \, [\, -2] \,\, < = \,\, \text{self.convergence\_criterion} \, : \end{array} 
109
110
                          \text{if } \text{free\_energy} \left[-1\right] \ - \ \text{free\_energy} \left[-2\right] \ <= \ \text{self.convergence\_criterion} : 
\begin{array}{c} 111 \\ 112 \end{array}
                  break
return free_energy
113
114
       def init_mean_field_approximation (
116
117
       k: int, n: int, max_steps, convergence_criterion ) -> MeanFieldApproximation:
            return MeanFieldApproximation (
119
                  \label{eq:lambda_matrix} \begin{array}{l} \texttt{lambda_matrix} = \texttt{np.random.random} \left( \, \, \texttt{size} = (\texttt{n} \,, \  \, \texttt{k} \,) \, \right) \,, \end{array}
120
                   max_steps=max_steps,
                   convergence_criterion=convergence_criterion,
```

src/models/binary_latent_factor_approximations/mean_field_approximation.py

The Python code for expectation maximisation:

```
from __future__ import annotations
 3
     from typing import List, Tuple
     import numpy as np
     from src.models.binary_latent_factor_approximations.abstract_binary_latent_factor_approximation import (
          AbstractBinaryLatentFactorApproximation\;,\\
            src.models.binary_latent_factor_models.binary_latent_factor_model import (
11
          AbstractBinaryLatentFactorModel,
12
13
14
     def is_converge(
          free_energies: List[float],
current_lambda_matrix: np.ndarray,
previous_lambda_matrix: np.ndarray,
16
17
19
     ) -> bool:
20
21
          Check for convergence of free energy and lambda matrix
22
23
          :param free_energies: list of free energies
          :param current_lambda_matrix: current lambda matrix
:param previous_lambda_matrix: previous lambda matrix
24
25
          :return: boolean indicating convergence
26
27
           \begin{array}{lll} \textbf{return} & (abs(free\_energies[-1] - free\_energies[-2]) == 0) \ and \ np. \\ linalg.norm(current\_lambda\_matrix - previous\_lambda\_matrix \end{array} 
28
29
30
          ) == 0
     def learn_binary_factors(
34
          x: np.ndarray,
em_iterations: int,
35
          binary_latent_factor_model: AbstractBinaryLatentFactorModel, binary_latent_factor_approximation: AbstractBinaryLatentFactorApproximation,
36
     ) -> Tuple [
AbstractBinaryLatentFactorApproximation,
38
39
40
          AbstractBinaryLatentFactorModel,
41
          List [float],
42
     1:
44
          Expectation maximisation algorithm to learn binary factors.
45
          :param x: data matrix (number_of_points, number_of_dimensions)
          :param em_iterations: number of iterations to run EM
:param binary_latent_factor_model: a binary_latent_factor_model
:param binary_latent_factor_approximation: a binary_latent_factor_approximation
47
48
49
          :return: a Tuple containing the updated binary_latent_factor_model, updated binary_latent_factor_approximation,
50
                      and free energies during each step of EM
          free\_energies: List[float] = [
54
               binary\_latent\_factor\_approximation.compute\_free\_energy (
                    x, binary_latent_factor_model
57
58
          for _ in range (em_iterations):
59
               previous_lambda_matrix = np.copy(
                    \verb|binary_latent_factor_approximation.lambda_matrix|
               )
61
               # E step
               binary_latent_factor_approximation.variational_expectation_step(
65
                    binary_latent_factor_model=binary_latent_factor_model,
66
               )
68
               # M step
69
                binary_latent_factor_model.maximisation_step(
71
72
73
74
75
                     binary_latent_factor_approximation,
               free_energies.append(
76
77
78
                    binary_latent_factor_approximation.compute_free_energy(
                        x, binary_latent_factor_model
79
80
               if is_converge(
                     free_energies ,
82
                     binary_latent_factor_approximation.lambda_matrix,
                    previous_lambda_matrix ,
83
85
          {\bf return \ binary\_latent\_factor\_approximation \ , \ binary\_latent\_factor\_model \ , \ free\_energies \ }
```

src/expectation_maximisation.py

The rest of the Python code for question 3:

```
from typing import List
     import matplotlib.pyplot as plt
     import numpy as np
      from src.expectation_maximisation import is_converge, learn_binary_factors
      from src.models.binary_latent_factor_approximations.mean_field_approximation import (
          init_mean_field_approximation
10
            {\tt src.models.binary\_latent\_factor\_models.binary\_latent\_factor\_model \ \underline{import} \ \ (
11
          AbstractBinaryLatentFactorModel, init_binary_latent_factor_model,
13
14
16
17
     def e_and_f(
          x: np.ndarray,
k: int,
           em_iterations: int.
19
20
          e_maximum_steps: int
           e_convergence_criterion: float,
22
     save_path: str ,
) -> AbstractBinaryLatentFactorModel:
23
24
25
          n \; = \; x \, . \, s \, h \, a \, p \, e \; [ \, 0 \, ]
           {\tt mean\_field\_approximation} \ = \ {\tt init\_mean\_field\_approximation} \ (
26
                k, n, max_steps=e_maximum_steps, convergence_criterion=e_convergence_criterion
27
           binary_latent_factor_model = init_binary_latent_factor_model(
28
29
               x, mean_field_approximation
30
          fig , ax = plt.subplots(1, k, figsize=(k * 2, 2))
for i in range(k):
    ax[i].imshow(binary_latent_factor_model.mu[:, i].reshape(4, 4))
    ax[i].set_title(f"Latent Feature mu_{i}")
fig.suptitle("Initial Features (Mean Field Learning)")
plt_title("Initial Features (Mean Field Learning)")
34
35
36
           plt.tight_layout()
           plt.savefig(save_path + "-init-latent-factors", bbox_inches="tight")
38
           _, binary_latent_factor_model, free_energy = learn_binary_factors(
39
40
41
                em_iterations , binary_latent_factor_model ,
42
                \verb|binary_latent_factor_approximation=mean\_field_approximation|,
44
          45
47
48
49
          plt.tight_layout()
plt.savefig(save_path + "-latent-factors", bbox_inches="tight")
50
51
           plt.close()
           plt.title("Free Energy (Mean Field Learning)")
          plt.xlabel("t (EM steps)")
plt.ylabel("Free Energy")
plt.plot(free_energy)
56
           plt.savefig(save_path + "-free-energy", bbox_inches="tight")
58
59
           plt.close()
           return binary_latent_factor_model
61
62
     def g(
          x: np.ndarray, binary_latent_factor_model: AbstractBinaryLatentFactorModel,
65
66
           sigmas: List [float],
67
          k: int,
           em_iterations: int
          e_maximum_steps: int,
           e_convergence_criterion: float,
70
71
72
73
74
75
76
          save_path: str,
     ) -> None:
          n = x.shape[0]
free_energies = []
          for sigma in sigmas:
binary_latent_factor_model.sigma = sigma
                mean\_field\_approximation = init\_mean\_field\_approximation (
                     k,
80
81
                      convergence_criterion=e_convergence_criterion ,
                free_energy: List[float] = [
    mean_field_approximation.compute_free_energy(x, binary_latent_factor_model)
83
84
86
                      _ in range (em_iterations):
                      free_energy.pop(-1)
87
                     previous.lambda_matrix = np.copy(mean_field_approximation.lambda_matrix)
new_free_energy = mean_field_approximation.variational_expectation_step(
binary_latent_factor_model=binary_latent_factor_model,
89
90
91
92
                      free_energy.extend(new_free_energy)
```

```
\begin{array}{lll} & free\_energy \, [\, -1] \, - \, free\_energy \, [\, -2] \\ <= \, mean\_field\_approximation\_convergence\_criterion \end{array}
 97
                              ):
 98
                                       free\_energy.pop(-1)
 99
                               if is_converge (
101
                                      free_energy ,
                                      {\tt mean\_field\_approximation.lambda\_matrix}\;,
                                      previous_lambda_matrix,
103
                       {\tt free\_energies.append} \, (\, {\tt free\_energy} \, )
106
                for i, free_energy in enumerate(free_energies):
    plt.plot(
        free_energy,
108
109
110
\begin{array}{c} 111 \\ 112 \end{array}
                               label=f"sigma={sigmas[i]}",
                plt.title(f"F(t)")
plt.xlabel("t (Variational E steps)")
plt.ylabel("F(t)")
113
114
115
                plt.tight.layout()
plt.legend()
plt.savefig(save_path + f"-free-energy-sigma.png", bbox_inches="tight")
116
117
119
120
                for i, free_energy in enumerate(free_energies):
    diffs = np.log(np.diff(free_energy))
    plt.plot(
        diffs,
123
124
                               label = f" sigma = \{ sigmas [i] \}",
126
                plt.title(f"\log(F(t)-F(t-1)"))
plt.xlabel("t (Variational E steps)")
plt.ylabel("\log(F(t)-F(t-1)"))
plt.tight_layout()
plt.legend()
plt.savefig(save_path + f"-free-energy-diff-sigma.png", bbox_inches="tight")
127
128
130
131
                 plt.close()
```

src/solutions/q3.py

Question 4

(a)

We begin by writing the expression for x_d :

$$P(x_d|s, \mathbf{w}_d, \sigma^2) = \mathcal{N}\left(\mathbf{s}^T \mathbf{w}_d, \sigma^2\right)$$

where we know from the diagonal covariance of $P(\mathbf{x}|\mathbf{s}, \mu, \sigma^2)$ that each dimension is independent. Moreover, $\mathbf{w}_d \in \mathbb{R}^{K \times 1}$, which is the d^{th} row of $\mu \in \mathbb{R}^{D \times K}$

Thus, we can write the posterior:

$$\log P(\mathbf{x}, \mathbf{s}, \mu | \pi, \sigma^2, \alpha) = \log P(\mathbf{s} | \pi) + \sum_{d=1}^{D} \log P(x_d | s, \mathbf{w}_d, \sigma^2) + \log P(\mathbf{w}_d | \alpha)$$

where we introduce priors on each \mathbf{w}_k with $\alpha \in \mathbb{R}^{K \times 1}$.

We choose each prior to be:

$$P(\mathbf{w}_d|\alpha) = \mathcal{N}(0, \mathbf{A}^{-1})$$

where $\mathbf{A} = diag(\alpha)$, the precision matrix.

Combining, we have our expression:

$$\log P(\mathbf{x}, \mathbf{s}, \mu | \pi, \sigma^{2}, \alpha) =$$

$$+ \sum_{d=1}^{D} -\frac{-1}{2} \log(2\pi\sigma^{2}) - \frac{1}{2\sigma^{2}} \left(x_{d}^{2} - 2x_{d}\mathbf{s}^{T}\mathbf{w}_{d} + \mathbf{w}_{d}^{T}\mathbf{s}\mathbf{s}^{T}\mathbf{w}_{d} \right)$$

$$+ \sum_{k=1}^{K} s_{k} \log \pi_{k} + (1 - s_{k}) \log(1 - \pi_{k})$$

$$+ \sum_{d=1}^{D} -\frac{K}{2} \log(2\pi) + \frac{1}{2} \sum_{k=1}^{K} (\log \alpha_{k}) - \frac{1}{2} \mathbf{w}_{d}^{T}\mathbf{A}\mathbf{w}_{d}$$

For the Variational Bayes expectation step, we minimise $\mathbf{KL}[q_s(\mathbf{s}|\text{everything else})||P(\mathbf{s}|\text{everything else})]$ by setting:

$$q_s(\mathbf{s}) \propto \exp \left\langle \log P(\mathbf{x}, \mathbf{s}, \mu | \pi, \sigma^2, \alpha) \right\rangle_{q(\mu)}$$

Substituting the relevant terms:

$$q_s(\mathbf{s}) \propto \exp\left\{-\frac{1}{2\sigma^2}\left(-2\mathbf{x}^T\sum_{k=1}^K s_k \mu_k + \sum_{k=1}^K \sum_{k'=1, k' \neq k}^K s_k s_{k'} \mu_k^T \mu_{k'} + \sum_{k=1}^K s_k \mu_k^T \mu_k\right) + \sum_{k=1}^K s_k \log\frac{\pi_k}{1 - \pi_k}\right\}_{q(\mu)}$$

Given our factored approximation $q(\mathbf{s}) = \prod_{i=1}^{K} q_i(s_i)$, we can see that we can derive a similar partial update for $q_i(s_i)$ as in Question 3, by taking the variation derivative of the Lagrangian to enforce the normalisation of q_i :

$$\frac{\partial}{\partial q_i} \left(\exp \left\langle \log P(\mathbf{x}, \mathbf{s}, \mu | \pi, \sigma^2, \alpha) \right\rangle_{q(\mu)} + \lambda^{LG} \int q_i - 1) \right) \propto \exp \left\langle \log P(\mathbf{x}, \mathbf{s}, \mu | \pi, \sigma^2, \alpha) \right\rangle_{q(\mu) \prod_{j \neq i} q_j(s_j)} - \log q_i(s_i)$$

Setting this to zero we can solve for λ_i where $q_i(s_i) = \lambda_i^{s_i} (1 - \lambda_i)^{(1-s_i)}$:

$$\lambda_i = \frac{1}{1 + \exp\left[-\left(\frac{\langle \mu_i \rangle_{q_{\mu_i}}^T}{\sigma^2} \left(\mathbf{x} - \frac{\langle \mu_i \rangle_{q_{\mu_i}}}{2} - \sum_{j=1, j \neq i}^K \lambda_j \langle \mu_j \rangle_{q_{\mu_j}}\right) + \log \frac{\pi_i}{1 - \pi_i}\right)\right]}$$

we have our partial E step update.

For the maximisation step, we perform maximisation steps for the parameters σ and π in the same way as question 3. However, having defined a prior on μ (through **w**) so we will have to derive our expression for $\langle \mu_k \rangle_{q_{\mu_k}}$ the expectation of the posterior on μ_k . This involves deriving the posterior distribution of \mathbf{w}_d

$$q_{\mathbf{w}_d}(\mathbf{w}_d) \propto P(\mathbf{w}_d) \exp \left\langle \log P(\mathbf{x}, \mathbf{s}, \mu | \pi, \sigma^2, \alpha) \right\rangle_{q_{\mathbf{s}}(\mathbf{s})q_{\neg \mathbf{w}_d}(\mathbf{w}_d)}$$

Substituting the appropriate terms:

$$q_{\mathbf{w}_d}(\mathbf{w}_d) \propto \exp\left(-\frac{1}{2}\mathbf{w}_d^T \mathbf{A} \mathbf{w}_d\right) \exp\left\langle-\frac{1}{2\sigma^2} \left(-2x_d \mathbf{s}^T \mathbf{w}_d + \mathbf{w}_d^T \mathbf{s} \mathbf{s}^T \mathbf{w}_d\right)\right\rangle_{q_{\mathbf{s}}(\mathbf{s})q_{\neg \mathbf{w}_d}(\mathbf{w}_d)}$$

Simplifying:

$$q_{\mathbf{w}_d}(\mathbf{w}_d) \propto \exp\left(-\frac{1}{2}\left(\mathbf{w}_d^T\left(\mathbf{A} + \frac{\langle \mathbf{s}\mathbf{s}^T\rangle_{q_{\mathbf{s}}(\mathbf{s})}}{\sigma^2}\right)\mathbf{w}_d - 2\left(\frac{x_d\langle \mathbf{s}^T\rangle_{q_{\mathbf{s}}(\mathbf{s})}}{\sigma^2}\right)\mathbf{w}_d\right)\right)$$

We see that the posterior:

$$q_{\mathbf{w}_d}(\mathbf{w}_d) = \mathcal{N}\left(\mu_{\mathbf{w}_d}, \Sigma_{\mathbf{w}_d}\right)$$

where:

$$\Sigma_{\mathbf{w}_d} = \left(\frac{\langle \mathbf{s} \mathbf{s}^T \rangle_{q_{\mathbf{s}}(\mathbf{s})}}{\sigma^2} + \mathbf{A}\right)^{-1}$$

and

$$\mu_{\mathbf{w}_d} = \Sigma_{\mathbf{w}_d} \left(\frac{x_d \langle \mathbf{s}^T \rangle_{q_{\mathbf{s}}(\mathbf{s})}}{\sigma^2} \right)$$

Thus, $\langle \mu_k \rangle_{q_{\mu_k}} \in \mathbb{R}^{D \times 1}$ is simply the concatenation of the k^{th} elements of $\mu_{\mathbf{w}_d}$ for $d \in \{1, ..., D\}$ For ARD, we must also optimise α with a hyper-M step. We start by choose $Ga(\alpha_k|a,b)$, a Gamma prior on α_k , with a and b being hyperparameters. Thus, to optimise α we want to maximise the penalised objective:

$$\alpha = \arg \max_{\alpha} \left\langle \log P(\mathbf{x}, \mathbf{s}, \mu | \pi, \sigma^2, \alpha) \right\rangle_{q(\mathbf{w})} + \sum_{k=1}^{K} \log P(\alpha_k | a, b)$$

Substituting the appropriate terms, we have our penalised objective Q:

$$Q = \left\langle \sum_{d=1}^{D} \frac{1}{2} \sum_{k=1}^{K} (\log \alpha_k) - \frac{1}{2} \mathbf{w}_d^T \mathbf{A} \mathbf{w}_d \right\rangle_{q(\mathbf{w})} + \sum_{k=1}^{K} (a-1) \log \alpha_k - b\alpha_k$$

Simplifying:

$$Q = \frac{D}{2} \sum_{k=1}^{K} (\log \alpha_k) - \frac{1}{2} \sum_{d=1}^{D} \left(tr \left[\mathbf{A} \left\langle \mathbf{w}_d \mathbf{w}_d^T \right\rangle_{q(\mathbf{w}_d)} \right] \right) + \sum_{k=1}^{K} (a-1) \log \alpha_k - b\alpha_k$$

Setting $\frac{dQ}{d\alpha_k} = 0$ we get:

$$\frac{D}{2\alpha_k} - \frac{1}{2} \sum_{d=1}^{D} \left\langle (w_{d,k})^2 \right\rangle_{q(\mathbf{w}_d)} + \frac{a-1}{\alpha_k} - b = 0$$

where $w_{d,k}$ is the k^{th} element of \mathbf{w}_d . Knowing $\langle (w_{d,k})^2 \rangle_{q(\mathbf{w}_d)} = (\mu_{\mathbf{w}_{d,k}})^2 + \Sigma_{\mathbf{w}_{d,(k,k)}}$, we can solve for α_k :

$$\alpha_k = \frac{2a + D - 2}{2b + \sum_{d=1}^{D} \left((\mu_{\mathbf{w}_{d,k}})^2 + \sum_{\mathbf{w}_{d,(k,k)}} \right)}$$

we have our hyper-M steps for optimising α .

(b)

Running variational Bayes for different values of k, we can visualise the learned features μ_k and corresponding α_k^{-1} :

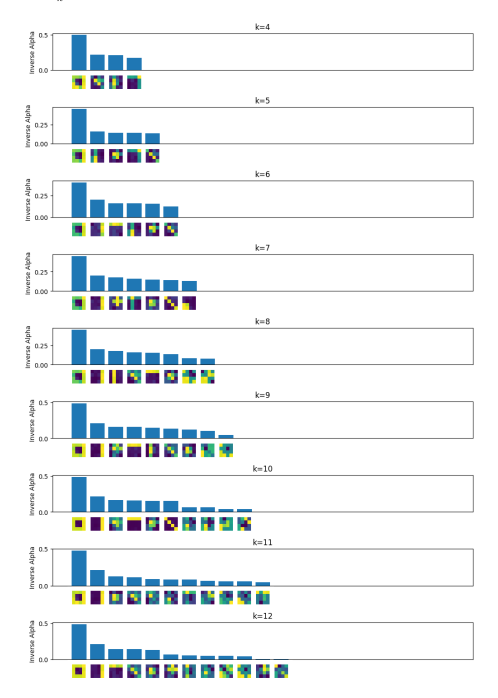


Figure 20: Learned Latent Factors vs Inverse Alpha

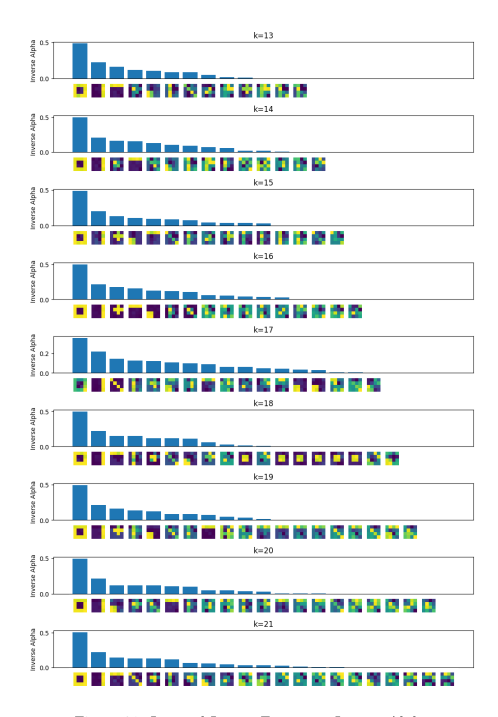


Figure 21: Learned Latent Factors vs Inverse Alpha

As we expect, when running the algorithm for higher K values, many of the features have $\alpha_k \to \infty$, depicted as α_k^{-1} for visual convenience. Moreover, visualising the learned features, we can see the clearest features often have the highest α_k^{-1} while the features deemed irrelevant are often noisy or duplicates.

Comparing the free energy plots of models trained on different K values:

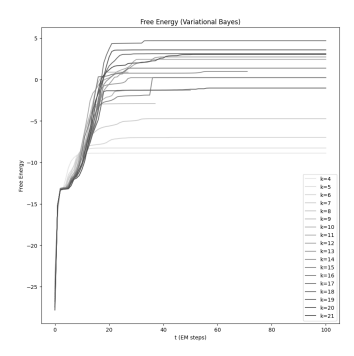


Figure 22: Free Energy for different values of k

We can see that initially, for k = 4 to k = 8, increasing k significantly increases the convergence value of the free energy. However, beyond k = 8 there is a no clear trend of k versus the free energy convergence value. We can see that this corresponds to the visualisation of α^{-1} where beyond k = 11, the effective number of features remains more or less the same. We know that there are only eight latent features, thus models with k > 8 should be learning duplicate or irrelevant features. As such, we wouldn't expect a model to be able to increase it's free energy significantly when provided with additional degrees of freedom by increasing the value of k beyond eight. We see that for models with k >> 8, there are typically ten or eleven features that might be deemed relevant (depending on how you threshold) and this is likely from slight overfitting, noise in the data, or duplicate features. Thus, the relationship between the free energy and the effective number of latent features for each model is as we would expect with ARD.

The Python code for Variational Bayes:

```
import numpy as np
 3
     from src.models.binary_latent_factor_approximations.abstract_binary_latent_factor_approximation import (
           AbstractBinaryLatentFactorApproximation,
     from src.models.binary_latent_factor_models.binary_latent_factor_model import (
           AbstractBinaryLatentFactorModel,
           BinaryLatentFactorModel,
10
12
     class Gaussian Prior:
13
14
           def_{n,n} = i n i t_{--} (self, a, b, d, k):
                Gaussian prior on mu matrix
16
17
                :param a: alpha parameter of Gamma Prior
                :param b: beta parameter of Gamma Prior
                :param d: number of dimensions
:param k: number of latent factors
"""
19
20
21
22
                self.a = a
23
                self.b = b
                self.mu = np.zeros((d, k))
self.alpha = np.ones((k,))
24
25
26
                self.w_covariance = np.zeros((k, k))
27
28
           def mu_k(self, k):
29
30
                Column vector of mu matrix, the latent feature vector
                : \mathtt{param} \ k \colon \ \mathtt{latent} \ \mathtt{factor} \ \mathtt{index}
                :return: column vector (number_of_dimensions, 1)
34
35
                \textcolor{return}{\mathtt{return}} \hspace{0.2cm} \mathtt{self.mu} [:, \hspace{0.2cm} k \hspace{0.2cm} : \hspace{0.2cm} k \hspace{0.2cm} + \hspace{0.2cm} 1]
36
           def w_d(self, d):
38
                Row vector of mu matrix, the weight vector for a particular dimension (pixel) of the data
39
40
                :param d: data dimension index
:return: row vector (1, number_of_latent_variables)
"""
41
42
44
                return self.mu[d : d + 1, :]
45
           def a_matrix(self) -> np.ndarray:
47
48
49
                Precision matrix for a weight vector w_d
                : return: \ matrix \ of \ shape \ ( \ number\_of\_latent\_variables \ , \ number\_of\_latent\_variables )
50
51
                return np.diag(self.alpha)
55
     {\bf class} \quad Variational Bayes Binary Latent Factor Model (Abstract Binary Latent Factor Model): \\
56
           \begin{array}{lll} \textbf{def} & \texttt{\_init\_\_(self} \;, \; mu \colon \; Gaussian Prior \;, \; \; variance \colon \; \textbf{float} \;, \; \; pi \colon \; np.\, ndarray) \colon \\ & \texttt{`````} \end{array}
57
                Variational Bayes implementation with prior on mu
58
59
                :param mu: Gaussian prior on latent features
                :param variance: Gaussian noise parameter
:param pi: vector of priors (1, number_of_latent_variables)
61
62
                \verb|self.gaussian_prior| = mu
65
                self._variance = variance
66
                self._pi = pi
67
           @property
           def variance(self) -> float:
70
71
72
73
74
75
76
                return self._variance
           @property
           def pi(self) -> np.ndarray:
    return self._pi
           @property
77
78
79
           def mu(self) -> np.ndarray:
    return self.gaussian.prior.mu
80
           def _update_w_d_covariance(
81
                binary\_latent\_factor\_approximation: \ AbstractBinaryLatentFactorApproximation\;,
83
84
                     (number_of_latent_variables, number_of_latent_variables)
                86
89
90
           def _update_w_d_mean(
91
                 \begin{array}{lll} \textbf{x}: & \texttt{np.ndarray}\,, & \# \; (\texttt{number\_of\_points}\,, & \texttt{number\_of\_dimensions}) \\ \textbf{binary\_latent\_factor\_approximation}: & \texttt{AbstractBinaryLatentFactorApproximation}\,, \\ \end{array} 
92
```

```
) -> None:
97
                 Update mean vector for w_d.
 98
                 :param x: data matrix (number_of_points, number_of_dimensions)
:param binary_latent_factor_approximation: a binary_latent_factor_approximation
:param d: index of data dimension to update
 99
101
                # (number_of_latent_variables, 1)
self.gaussian_prior.mu[d : d + 1, :] = (
    self.gaussian_prior.w_covariance
106
108
                      @ ( \# (number_of_latent_variables, number_of_latent_variables) self.precision
109
                            *\ binary\_latent\_factor\_approximation.expectation\_s.T\ \#\ (number\_of\_latent\_variables\ ,
             number_of_points)
                            @ x[:', d: d+1] # (number_of_points, 1)
112
                 ) . T
113
114
115
            def _hyper_maximisation_step(self) -> None:
116
                 Hyper M step updating alpha, which parameterize the covariance matrix of the Gaussian prior on mu
118
119
                 for k in range(self.k):
                       122
124
125
126
            def maximisation_step (
                 self , x: np.ndarray
                 binary\_latent\_factor\_approximation: \ AbstractBinaryLatentFactorApproximation\;,
130
            ) -> None:
131
                 \label{lem:maximisation} \begin{tabular}{lll} Maximisation step which runs the usual $M$-step followed by posterior updates to the distribution of mu as well as a hyper $M$-step updating the prior parameters on mu, the alpha vector :param x: data matrix (number_of_points, number_of_dimensions) \\ \end{tabular}
134
                 :param binary_latent_factor_approximation: a binary_latent_factor_approximation
136
                               pi \ = \ Binary Latent Factor Model. \ calculate\_maximis at ion\_parameters \, (
138
                       {\tt x, binary\_latent\_factor\_approximation}
139
140
                 self._variance = sigma**2
                 self._pi = pi
self._update_w_d_covariance(binary_latent_factor_approximation)
141
                 for d in range(self.d):
    self._update_w_d_mean(x, binary_latent_factor_approximation, d)
self._hyper_maximisation_step()
143
144
```

src/models/binary_latent_factor_models/variational_bayes.py

The Python code for Automatic Relevance Determination:

```
from typing import List, Tuple
 \frac{3}{4}
     import numpy as np
     from src.expectation_maximisation import learn_binary_factors
     from src.models.binary_latent_factor_approximations.mean_field_approximation import (
          init_mean_field_approximation,
     from src.models.binary_latent_factor_models.binary_latent_factor_model import (
10
          {\tt BinaryLatentFactorModel}\ ,
11
12
     from src.models.binary_latent_factor_models.variational_bayes import (
13
14
          Gaussian Prior,
          VariationalBayesBinaryLatentFactorModel,
15
16
17
18
19
     def run_automatic_relevance_determination (
         x: np.ndarray,
a_parameter: int,
b_parameter: int,
20
21
22
          k: int,
23
          em_iterations: int,
24
25
          e_maximum_steps: int,
e_convergence_criterion: float,
     ) -> Tuple [VariationalBayesBinaryLatentFactorModel, List [float]]:
26
27
28
          Run automatic relevance determination with variational Bayes.
29
30
          : param \ x: \ data \ matrix \ (number\_of\_points \, , \ number\_of\_dimensions)
31
          :param a_parameter: alpha parameter for gamma prior :param b_parameter: beta parameter for gamma prior
33
34
          :param k: number of latent variables
          :param k: number of latent variables :param em_iterations: number of iterations to run EM :param e_maximum_steps: maximum number of iterations of partial expectation steps
35
          :param e_convergence_criterion: minimum required change in free energy for each partial expectation step :return: a Tuple containing the optimised VB model and a list of free energies during each EM step
36
38
39
          n = x.shape[0]
          mean_field_approximation = init_mean_field_approximation (
41
               {\tt k,\ n,\ max\_steps=e\_maximum\_steps},\ {\tt convergence\_criterion=e\_convergence\_criterion}
42
          (_, sigma, pi,) = BinaryLatentFactorModel.calculate_maximisation_parameters(
               x, mean_field_approximation
44
45
          mu = Gaussian Prior (
47
               a=a_parameter
48
               b=b_parameter,
49
               k=k,
               d=x \cdot shape[1],
50
51
52
          binary_latent_factor_model: VariationalBayesBinaryLatentFactorModel = (
53
54
               Variational Bayes Binary Latent Factor Model (\\
                   mu=mu,
55
                    variance=sigma**2,
56
                    рі=рі,
58
59
          , binary_latent_factor_model, free_energy = learn_binary_factors(
60
               em_iterations=em_iterations ,
binary_latent_factor_model=binary_latent_factor_model ,
61
               \verb|binary_latent_factor_approximation| = \verb|mean_field_approximation|,
          return binary_latent_factor_model, free_energy
```

src/automatic_relevance_determination.py

The rest of the Python code for question 4:

```
from typing import List
     import matplotlib.pyplot as plt
     import numpy as np from matplotlib.offsetbox import AnnotationBbox, OffsetImage
     from src.automatic_relevance_determination import run_automatic_relevance_determination
10
     def _offset_image(coord: int, path: str, ax: plt.axis):
12
13
14
          Add image to matplotlib axis.
          :param coord: coordinate on axis
          :param path: path to image :param ax: plot axis
16
17
          \begin{array}{ll} img = plt.imread(path) \\ im = OffsetImage(img, zoom=0.72) \end{array}
19
20
21
          im.image.axes = ax
22
23
          ab = AnnotationBbox(
24
                (\text{coord}, 0),

xybox = (0.0, -19.0),
25
26
27
                frameon=False,
28
                xvcoords="data
29
                boxcoords="offset points",
30
                pad=0,
          ax.add_artist(ab)
34
35
     def b(
          x: np.ndarray,
a_parameter: int,
b_parameter: int,
36
38
          ks: List[int],
39
          max_k: int,
40
41
           em_iterations: int .
42
          e_maximum_steps: int
           e_convergence_criterion: float,
     save_path: str ,
) -> None:
44
45
47
          binary_latent_factor_models = []
48
          free_energies = []
# automatic relevance determination
49
50
           for i, k in enumerate(ks):
51
                      binary_latent_factor_model,
                free_energy ,
) = run_automatic_relevance_determination(
56
                     a_parameter, b_parameter,
58
59
                     em_iterations ,
                     e_maximum_steps
61
                      e_convergence_criterion ,
62
                binary_latent_factor_models.append(binary_latent_factor_model)
                free_energies.append(free_energy)
65
          # store each feature as an image for later use
for i, k in enumerate(ks):
    sort_indices = np.argsort(binary_latent_factor_models[i].gaussian_prior.alpha)
    for j, idx in enumerate(sort_indices):
        fig = plt.figure(figsize = (0.3, 0.3))
        ax = plt.Axes(fig, [0.0, 0.0, 1.0, 1.0])
        ax.set_axis_off()
        fig add axes(ax)
67
70
71
72
73
74
75
76
                     fig.add.axes(ax)
ax.imshow(binary_latent_factor_models[i].mu[:, idx].reshape(4, 4))
fig.savefig(save_path + f"-latent-factor-{i}-{j}", bbox_inches="tight")
plt.close()
77
78
79
          \# bar plot of alphas fig , ax = plt.subplots(len(ks), 1, figsize=(12, 2 * len(ks)))
80
           plt.subplots_adjust(hspace=1)
for i, k in enumerate(ks):
81
                sort_indices = np.argsort(binary_latent_factor_models[i].gaussian_prior.alpha)
          83
84
86
89
90
91
                92
```

```
-offset_image(j, path, ax[i])
os.remove(path)
fig.savefig(save_path + f"-latent-factors-comparison", bbox_inches="tight")
plt.close()

# free energy plot
fig = plt.figure()
fig.set_figwidth(10)
fig.set_figheight(10)
shades = np.flip(np.linspace(0.3, 0.9, len(ks)))
for i, k in enumerate(ks):
    plt.plot(free_energies[i], label=f"{k=}", color=np.ones(3) * shades[i])

plt.title("Free Energy (Variational Bayes)")
plt.xlabel("t (EM steps)")
plt.ylabel("Free Energy")
plt.legend()
plt.savefig(save_path + "-free-energy", bbox_inches="tight")
plt.close()
```

src/solutions/q4.py

Question 5

(a)

The log-joint probability for a single observation-source pair:

$$\log p(\mathbf{s}, \mathbf{x}) = \log p(\mathbf{s}) + (\mathbf{x}|\mathbf{s})$$

Knowing $p(\mathbf{s}) = \prod_{i=1}^K p(s_i|\pi_i)$ and $p(\mathbf{x}|\mathbf{s}) = \mathcal{N}(\sum_{i=1}^K s_i \mu_i, \sigma^2 \mathbf{I})$:

$$\log p(\mathbf{s}, \mathbf{x}) \propto \frac{-1}{2} \left(\mathbf{x} - \sum_{i=1}^{K} s_i \mu_i \right)^T \frac{1}{\sigma^2} \mathbf{I} \left(\mathbf{x} - \sum_{i=1}^{K} s_i \mu_i \right) + \sum_{i=1}^{K} \left(s_i \log \pi_i + (1 - s_i) \log(1 - \pi_i) \right)$$

Expanding:

$$\log p(\mathbf{s}, \mathbf{x}) \propto \frac{-1}{2\sigma^2} \left(\mathbf{x}^T \mathbf{x} - 2\mathbf{x}^T \sum_{i=1}^K s_i \mu_i + \sum_{i=1}^K \sum_{j=1}^K s_i s_j \mu_i^T \mu_j \right) + \sum_{i=1}^K \left(s_i \log \pi_i + (1 - s_i) \log(1 - \pi_i) \right)$$

Collecting terms pertaining to s_i :

$$\log p(\mathbf{s}, \mathbf{x}) = \sum_{i=1}^{K} \left(\left(\frac{\mathbf{x}^T \mu_i}{\sigma^2} + \log \frac{\pi_i}{1 - \pi_i} \right) s_i \right) + \sum_{i=1}^{K} \sum_{j=1}^{K} \left(\frac{-\mu_i^T \mu_j}{2\sigma^2} s_i s_j \right) + C$$

where C are all other terms without s_i .

Knowing that $s_i^2 = s_i$:

$$\log p(\mathbf{s}, \mathbf{x}) = \sum_{i=1}^{K} \left(\left(\frac{\mathbf{x}^T \mu_i}{\sigma^2} + \log \frac{\pi_i}{1 - \pi_i} - \frac{\mu_i^T \mu_i}{2\sigma^2} \right) s_i \right) + \sum_{i=1}^{K} \sum_{j=1}^{i-1} \left(\frac{-\mu_i^T \mu_j}{\sigma^2} s_i s_j \right) + C$$

Thus:

$$\log p(\mathbf{s}, \mathbf{x}) = \sum_{i=1}^{K} \log f_i(s_i) + \sum_{i=1}^{K} \sum_{j=1}^{i-1} \log g_{ij}(s_i, s_j)$$

where the factors are defined:

$$\log f_i(s_i) = \left(\frac{\mathbf{x}^T \mu_i}{\sigma^2} + \log \frac{\pi_i}{1 - \pi_i} - \frac{\mu_i^T \mu_i}{2\sigma^2}\right) s_i$$

and

$$\log g_{ij}(s_i, s_j) = \frac{-\mu_i^T \mu_j}{\sigma^2} s_i s_j$$

as required.

The Boltzmann Machine can be defined:

$$P(\mathbf{s}|\mathbf{W}, \mathbf{b}) = \frac{1}{Z} \exp\left(\sum_{i=1}^{K} \sum_{j=1}^{i-1} W_{ij} s_i s_j - \sum_{i=1}^{K} b_i s_i\right)$$

where $s_i \in \{0, 1\}$, the same as our source variables.

From our factorisation, we can see that $p(\mathbf{s}, \mathbf{x})$ is a Boltzmann Machine with:

$$W_{ij} = \frac{-\mu_i^T \mu_j}{\sigma^2}$$

and

$$b_i = -\left(\frac{\mathbf{x}^T \mu_i}{\sigma^2} + \log \frac{\pi_i}{1 - \pi_i} - \frac{\mu_i^T \mu_i}{2\sigma^2}\right)$$

and

$$\log Z = -C$$

(b)

For $f_i(s_i)$, we will choose a Bernoulli approximation:

$$\tilde{f}_i(s_i) = \lambda_i^{s_i} + (1 - \lambda_i)^{1 - s_i}$$

Thus,

$$\log \tilde{f}_i(s_i) \propto \log \left(\frac{\lambda_i}{1-\lambda_i}\right) s_i$$

For $g_{ij}(s_i, s_j)$, we will choose a product of Bernoulli's approximation:

$$\tilde{g}_{ij}(s_i, s_j) = \tilde{g}_{ij, \neg s_j}(s_i) \tilde{g}_{ij, \neg s_i}(s_j)$$

where

$$\tilde{g}_{ij,\neg s_i}(s_i) = (\theta_{ji})^{s_i} + (1 - \theta_{ji})^{1 - s_i}$$

and

$$\tilde{g}_{ii,\neg s_i}(s_i) = (\theta_{ij})^{s_j} + (1 - \theta_{ij})^{1 - s_j}$$

Thus,

$$\log \tilde{g}_{ij}(s_i, s_j) \propto \log \left(\frac{\theta_{ji}}{1 - \theta_{ji}}\right) s_i + \log \left(\frac{\theta_{ij}}{1 - \theta_{ij}}\right) s_j$$

we can define $\xi_{ji} = \log\left(\frac{\theta_{ji}}{1-\theta_{ji}}\right)$ and $\xi_{ij} = \log\left(\frac{\theta_{ij}}{1-\theta_{ij}}\right)$:

$$\log \tilde{g}_{ij}(s_i, s_j) \propto \xi_{ji} s_i + \xi_{ij} s_j$$

To derive the a message passing scheme, we first define the incoming message to node i from the singleton factor:

$$\mathcal{M}_i(s_i) = \tilde{f}_i(s_i)$$

and the message incoming message to node i from node j:

$$\mathcal{M}_{j\to i}(s_i) = \sum_{s_1 \in \{0,1\}} \cdots \sum_{s_{i-1} \in \{0,1\}} \sum_{s_{i+1} \in \{0,1\}} \cdots \sum_{s_1 \in \{0,1\}} \tilde{f}_j(s_j) \tilde{g}_{ji}(s_j, s_i) \prod_{k \in ne(j), k \neq i}^{K} \mathcal{M}_{k\to j}(s_j)$$

where ne(j) are indices of neighbouring nodes of node j. Because $\tilde{g}_{ji}(s_j, s_i)$ is a product:

$$\mathcal{M}_{j\to i}(s_i) = \tilde{g}_{ji,\neg s_j}(s_i) \sum_{s_1 \in \{0,1\}} \cdots \sum_{s_{i-1} \in \{0,1\}} \sum_{s_{i+1} \in \{0,1\}} \cdots \sum_{s_1 \in \{0,1\}} \tilde{f}_j(s_j) \tilde{g}_{ji,\neg s_i}(s_j) \prod_{k \in ne(j), k \neq i}^K \mathcal{M}_{k\to j}(s_j)$$

Simplifying:

$$\mathcal{M}_{j\to i}(s_i) = \tilde{g}_{ji,\neg s_i}(s_i)$$

and,

$$\mathcal{M}_{j\to i}(s_i) \propto \exp\left(\xi_{ji}s_i\right)$$

Thus, the cavity distributions are:

$$q_{\neg \tilde{f}_i(s_i)}(s_i) = \prod_{j \in ne(i)}^K \mathcal{M}_{j \to i}(s_i)$$

and

$$q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) = \left(\mathcal{M}_i(s_i) \prod_{k \in ne(i), k \neq j}^K \mathcal{M}_{k \to i}(s_i)\right) \left(\mathcal{M}_j(s_j) \prod_{k \in ne(j), k \neq i}^K \mathcal{M}_{k \to j}(s_j)\right)$$

For $\tilde{f}_i(s_i)$, we do not need to make an approximation step. This is because we are minimising:

$$\tilde{f}_i(s_i) = \arg\min_{\tilde{f}_i(s_i)} \mathbf{KL} \left[f_i(s_i) q_{\neg \tilde{f}_i(s_i)}(s_i) \| \tilde{f}_i(s_i) q_{\neg \tilde{f}_i(s_i)}(s_i) \right]$$

We know that the factor $\log f_i(s_i)$ is a Bernoulli of the form $b_i s_i$. Because our approximation for this site is also Bernoulli, we can simply solve for λ_i in $\log \tilde{f}_i(s_i)$:

$$\log \tilde{f}_i(s_i) = \log f_i(s_i)$$

$$\log\left(\frac{\lambda_i}{1-\lambda_i}\right)s_i = b_i s_i$$

$$\lambda_i = \frac{1}{1 + \exp(-b_i)}$$

On the other hand, for $\tilde{g}_{ij}(s_i, s_j)$, we will approximate with:

$$\tilde{g}_{ij}(s_i, s_j) = \arg\min_{\tilde{g}_{ij}(s_i, s_j)} \mathbf{KL} \left[g_{ij}(s_i, s_j) q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \left\| \tilde{g}_{ij}(s_i, s_j) q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \right\| \right]$$

We can define natural parameters $\eta_{i,\neg s_j}$ and $\eta_{j,\neg s_i}$ for $q_{\neg \tilde{g}_{ij}(s_i,s_j)}(s_i,s_j)$ such that:

$$\mathcal{M}_i(s_i) \prod_{k \in ne(i), k \neq j}^K \mathcal{M}_{k \to i}(s_i) \propto \exp(\eta_{i, \neg s_j} s_i)$$

$$\mathcal{M}_{j}(s_{j}) \prod_{k \in ne(j), k \neq j}^{K} \mathcal{M}_{k \to j}(s_{j}) \propto \exp(\eta_{j, \neg s_{i}} s_{j})$$

Note that $\tilde{g}_{ij}(s_i, s_j)$ was chosen as the product of two Bernoulli distributions, updates to this site approximation involves updating the parameters ξ_{ij} and ξ_{ji} , for s_i and s_j respectively.

We can write:

$$\log \tilde{g}_{ij}(s_i, s_j) q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \propto \xi_{ji} s_i + \xi_{ij} s_j + \eta_{i, \neg s_j} s_i + \eta_{j, \neg s_i} s_j$$

Simplifying:

$$\log \tilde{g}_{ij}(s_i, s_j) q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \propto (\xi_{ji} + \eta_{i, \neg s_j}) s_i + (\xi_{ij} + \eta_{j, \neg s_i}) s_j$$

Thus, the first moments:

$$\mathbb{E}_{s_i} \left[\sum_{s_j \in \{0,1\}} \tilde{g}_{ij}(s_i, s_j) q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \right] = \frac{1}{1 + \exp\left(-\left(\xi_{ji} + \eta_{i, \neg s_j}\right)\right)}$$

and

$$\mathbb{E}_{s_j} \left[\sum_{s_i \in \{0,1\}} \tilde{g}_{ij}(s_i, s_j) q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \right] = \frac{1}{1 + \exp\left(-\left(\xi_{ij} + \eta_{j, \neg s_i}\right)\right)}$$

Moreover:

$$\log g_{ij}(s_i, s_j) q q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \propto W_{ij} s_i s_j + \eta_{i, \neg s_j} s_i + \eta_{j, \neg s_i} s_j$$

To derive the first moment for $g_{ij}(s_i, s_j)q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j)$ with respect to s_i , we first marginalise out s_j :

$$\sum_{s_j \in \{0,1\}} g_{ij}(s_i, s_j) q_{\neg \tilde{g}_{ij}(s_i, s_j)}(\mathbf{s}) \propto \exp\left(W_{ij} s_i + \eta_{i, \neg s_j} s_i + \eta_{j, \neg s_i}\right) + \exp\left(\eta_{i, \neg s_j} s_i\right)$$

Thus, the first moment:

$$\mathbb{E}_{s_i} \left[\sum_{s_j \in \{0,1\}} g_{ij}(s_i, s_j) q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \right] = \frac{\exp\left(W_{ij} + \eta_{i, \neg s_j} + \eta_{j, \neg s_i}\right) + \exp\left(\eta_{i, \neg s_j}\right)}{\left[\exp\left(W_{ij} + \eta_{i, \neg s_j} + \eta_{j, \neg s_i}\right) + \exp\left(\eta_{i, \neg s_j}\right)\right] + \left[\exp\left(\eta_{j, \neg s_i}\right) + 1\right]}$$

Simplifying:

$$\mathbb{E}_{s_i} \left[\sum_{s_j \in \{0,1\}} g_{ij}(s_i, s_j) q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \right] = \frac{\exp\left(\eta_{i, \neg s_j}\right) \left(\exp\left(W_{ij} + \eta_{j, \neg s_i}\right) + 1\right)}{\left[\exp\left(\eta_{i, \neg s_j}\right) \left(\exp\left(W_{ij} + \eta_{j, \neg s_i}\right) + 1\right)\right] + \left[\exp\left(\eta_{j, \neg s_i}\right) + 1\right]}$$

Similarly:

$$\mathbb{E}_{s_{j}}\left[\sum_{s_{i}\in\{0,1\}}g_{ij}(s_{i},s_{j})q_{\neg\tilde{g}_{ij}(s_{i},s_{j})}(s_{i},s_{j})\right] = \frac{\exp\left(\eta_{j,\neg s_{i}}\right)\left(\exp\left(W_{ij} + \eta_{i,\neg s_{j}}\right) + 1\right)}{\left[\exp\left(\eta_{j,\neg s_{i}}\right)\left(\exp\left(W_{ij} + \eta_{i,\neg s_{j}}\right) + 1\right)\right] + \left[\exp\left(\eta_{i,\neg s_{j}}\right) + 1\right]}$$

By setting:

$$\mathbb{E}_{s_i} \left[\sum_{s_j \in \{0,1\}} \tilde{g}_{ij}(s_i, s_j) q_{-\tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \right] = \mathbb{E}_{s_i} \left[\sum_{s_j \in \{0,1\}} g_{ij}(s_i, s_j) q_{-\tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \right]$$

and

$$\mathbb{E}_{s_j} \left[\sum_{s_i \in \{0,1\}} \tilde{g}_{ij}(s_i, s_j) q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \right] = \mathbb{E}_{s_j} \left[\sum_{s_i \in \{0,1\}} g_{ij}(s_i, s_j) q_{\neg \tilde{g}_{ij}(s_i, s_j)}(s_i, s_j) \right]$$

we can solve for the parameters of $\tilde{g}_{ij}(s_i, s_j)$ with moment matching:

$$\frac{1}{1 + \exp\left(-\left(\xi_{ji} + \eta_{i, \neg s_{j}}\right)\right)} = \frac{\exp\left(\eta_{i, \neg s_{j}}\right) \left(\exp\left(W_{ij} + \eta_{j, \neg s_{i}}\right) + 1\right)}{\left[\exp\left(\eta_{i, \neg s_{j}}\right) \left(\exp\left(W_{ij} + \eta_{j, \neg s_{i}}\right) + 1\right)\right] + \left[\exp\left(\eta_{j, \neg s_{i}}\right) + 1\right]}$$

Simplifying:

$$\exp(\eta_{j,\neg s_{i}}) + 1 = \exp\left(-\left(\xi_{ji} + \eta_{i,\neg s_{j}}\right)\right) \exp\left(\eta_{i,\neg s_{j}}\right) \left(\exp\left(W_{ij} + \eta_{j,\neg s_{i}}\right) + 1\right)$$
$$\frac{\exp(\eta_{j,\neg s_{i}}) + 1}{\exp(W_{ij} + \eta_{j,\neg s_{i}}) + 1} = \exp\left(-\xi_{ji}\right)$$

Our parameter update:

$$\xi_{ji} = \log \left(\frac{1 + \exp\left(W_{ij} + \eta_{j, \neg s_i}\right)}{1 + \exp\left(\eta_{j, \neg s_i}\right)} \right)$$

Similarly:

$$\xi_{ij} = \log \left(\frac{1 + \exp\left(W_{ij} + \eta_{i, \neg s_j}\right)}{1 + \exp\left(\eta_{i, \neg s_j}\right)} \right)$$

(c)

Using factored approximate messages, we see that:

$$\eta_{i, \neg s_j} = \log\left(\frac{\lambda_i}{1 - \lambda_i}\right) + \sum_{k \in ne(i), k \neq j}^K \log\left(\frac{\theta_{ki}}{1 - \theta_{ki}}\right)$$

Knowing $b_i = \log\left(\frac{\lambda_i}{1-\lambda_i}\right)$ and $\xi_{ki} = \log\left(\frac{\theta_{ki}}{1-\theta_{ki}}\right)$:

$$\eta_{i,\neg s_j} = b_i + \sum_{k \in ne(i), k \neq j}^K \xi_{ki}$$

and

$$\eta_{j,\neg s_i} = b_j + \sum_{k \in ne(j), k \neq i}^K \xi_{kj}$$

The summation of the natural parameters of the singleton factor for node i with the natural parameters of messages from all the neighbouring nodes.

This leads to a loopy BP algorithm because the nodes are fully connected (i.e. every node is the neighbour of all other nodes). Thus, we cannot simply move from one end of the graph to the other like BP for tree structured graphs.

(d)

Similar to question 3, we can use automatic relevance determination (ARD) as a hyperparameter method to select relevant features by placing a prior on μ_i . With a hyper-M step, certain features will have diverging precision, indicating that they are not relevant to the model output. Thus, the number of remaining features will be our selection for K.

Question 6

Implementing the EP/loopy-BP algorithm, we can compare the learned latent factors with those of the variational mean-field algorithm:

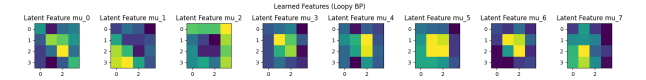


Figure 23: Learned Latent factors learned with EP/Loopy-BP

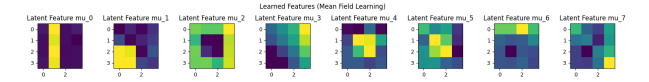
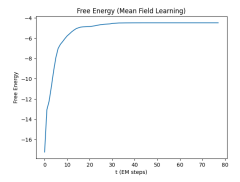


Figure 24: Learned Latent Factors with Mean Field Approximation

We can see that the mean field algorithm seems to learn better latent features. In particular there's are fewer duplicates, unlike loopy BP that has a few duplicates of the two by two square in the middle. Moreover, the learned features have less noise. For example μ_0 for the mean field algorithm looks almost like a binary image. We can understand the reason for this by comparing the free energies of the two algorithms:



-12 - -14 - -18 - -18 - -20 - 0 20 40 60 80 100 t (EM steps)

Figure 25: Mean Field Approximation

Figure 26: Loopy BP

We can observe that the free energy of the mean field algorithm converges while our loopy belief propagation is unable to converge to a free energy. Because loopy BP does not have convergence guarantees, this is one of the limitations of this approach.

The Python code for the Boltzmann machine:

```
import numpy as np
 3
     from src.models.binary_latent_factor_approximations.abstract_binary_latent_factor_approximation import (
           AbstractBinaryLatentFactorApproximation,
            src.models.binary_latent_factor_models.binary_latent_factor_model import (
           BinaryLatentFactorModel,
10
11
     class BoltzmannMachine (BinaryLatentFactorModel):
          def __init__(
self,
12
13
14
               mu: np.ndarray, sigma: float,
16
17
                pi: np.ndarray,
19
                Binary latent factor model as a Boltzmann Machine
20
21
                :param mu: matrix of means (number_of_dimensions, number_of_latent_variables)
                :param sigma: gaussian noise parameter:
:param pi: vector of priors (1, number_of_latent_variables)
22
23
24
25
                super().__init__(mu, sigma, pi)
26
27
28
          @property
          def w_matrix(self) -> np.ndarray:
29
30
                Weight matrix of the Boltzmann machine
                : return: \ matrix \ of \ weights \ (number\_of\_latent\_variables \,, \ number\_of\_latent\_variables)
34
                return -self.precision * (self.mu.T @ self.mu)
35
          def w_matrix_index(self, i, j) \rightarrow float:
36
38
                Weight matrix at a specific index
39
40
                :param i: row index
                :param j: column index
:return: weight value
41
42
44
                return -self.precision * (self.mu[:, i] @ self.mu[:, j])
45
          def b(self, x) \rightarrow np.ndarray:
47
48
                b term in the Boltzmann machine for all data points
49
                :param x: design matrix (number_of_points, number_of_dimensions)
:return: matrix of shape (number_of_points, number_of_latent_variables)
50
51
                return -(
                     self.precision * x @ self.mu
                     + self.log_pi_ratio - 0.5 * self.precision * np.multiply(self.mu, self.mu).sum(axis=0)
56
58
59
          def b_index(self, x, node_index) -> float:
60
61
                b term for a specific node in the Boltzmann machine for all data points
62
                :param x: design matrix (number_of_points, number_of_dimensions)
:param node_index: node index
:return: vector of shape (number_of_points, 1)
65
66
67
                     rn -(
self.precision * x @ self.mu[:, node_index]
+ (self.log_pi[0, node_index] - self.log_one_minus_pi[0, node_index])
- 0.5 * self.precision * self.mu[:, node_index] @ self.mu[:, node_index]
70
71
72
73
74
75
76
                ).reshape(
                      -1.
          @property
def log_pi_ratio(self) -> np.ndarray:
    return self.log_pi - self.log_one_minus_pi
80
     def init_boltzmann_machine(
          x \colon \text{ np.ndarray} \ , \\ binary\_latent\_factor\_approximation : \ AbstractBinaryLatentFactorApproximation \ , \\ \\
81
     ) -> BinaryLatentFactorModel:
83
84
           Initialise a boltzmann machine by running a maximisation step with the parameters of the
86
          binary latent factor approximation
          :param x: data matrix (number_of_points, number_of_dimensions)
:param binary_latent_factor_approximation: a binary_latent_factor_approximation
:return: an initialised Boltzmann machine model
"""
87
89
90
91
          mu, sigma, pi = BinaryLatentFactorModel.calculate_maximisation_parameters(
92
                x, binary_latent_factor_approximation
```

 $src/models/binary_latent_factor_models/boltzmann_machine.py$

The Python code for message passing:

```
from typing import List
  3
          import numpy as np
         from \  \  src.models.binary\_latent\_factor\_approximations.abstract\_binary\_latent\_factor\_approximation \  \  import \  \  (abstractBinary\_latentFactor\_approximation, abstractBinary\_latent\_factor\_approximation, abstractBinary\_approximation, abstractBin
          from src.models.binary_latent_factor_models.boltzmann_machine import BoltzmannMachine
10
          {\tt class} \ \ {\tt MessagePassingApproximation(AbstractBinaryLatentFactorApproximation):}
12
                   13
14
16
17
                                                 (number_of_points, number_of_latent_variables, number_of_latent_variables)
                   def __init__(self, eta_matrix: np.ndarray):
    self.eta_matrix = eta_matrix
19
20
22
                   @property
23
                   def lambda_matrix(self) -> np.ndarray:
24
                             Aggregate messages and compute parameter for Bernoulli distribution
25
26
                             :return:
27
                             \begin{array}{l} lambda\_matrix = 1 \ / \ (1 + np.exp(-self.xi.sum(axis=1))) \\ lambda\_matrix [ lambda\_matrix == 0] = 1e-10 \\ lambda\_matrix [ lambda\_matrix == 1] = 1 - 1e-10 \\ \end{array}
28
29
30
                             return lambda_matrix
                   @property
def xi(self) -> np.ndarray:
    return np.log(np.divide(self.eta_matrix, 1 - self.eta_matrix))
34
35
36
                   def aggregate_incoming_binary_factor_messages(
38
                             self, node_index: int, excluded_node_index: int
                   ) -> np.ndarray:
39
40
                            # (number_of_points,
41
                                    exclude message from excluded_node_index -> node_index
42
                                      np.sum(self.xi[:, :excluded_node_index, node_index], axis=1)
+ np.sum(self.xi[:, excluded_node_index + 1 :, node_index], axis=1)
44
45
                             ).reshape(
47
48
49
                   @staticmethod
                   def calculate_eta(xi: np.ndarray) -> np.ndarray:
    eta = 1 / (1 + np.exp(-xi))
    eta[eta == 0] = 1e-10
    eta[eta == 1] = 1 - 1e-10
    return eta
50
51
                   def variational_expectation_step(
    self, x: np.ndarray, binary_latent_factor_model: BoltzmannMachine)
    ->_List[float]:
56
58
59
                             Iteratively update singleton and binary factors
                             :param x: data matrix (number_of_points, number_of_dimensions)
:param binary_latent_factor_model: a binary_latent_factor_model
61
62
                             return: free energies after each update
65
                             free_energy = [self.compute_free_energy(x, binary_latent_factor_model)]
                             for i in range(self.k):
# singleton factor update
xi-new_ii = self.calculate_singleton_message_update(
66
67
                                                boltzmann\_machine=binary\_latent\_factor\_model\ ,
70
71
72
73
74
75
76
                                                x=x,
                                       /
self.eta_matrix[:, i, i] = self.calculate_eta(xi_new_ii)
free_energy.append(self.compute_free_energy(x, binary_latent_factor_model))
                                       for j in range(i):
                                                # binary factor update
xi_new_ij = self.calculate_binary_message_update(
boltzmann_machine=binary_latent_factor_model,
80
81
                                                           i=i ,
83
84
                                                )
self.eta_matrix[:, i, j] = self.calculate_eta(xi_new_ij)
xi_new_ji = self.calculate_binary_message_update(
86
                                                          boltzmann_machine=binary_latent_factor_model,
87
                                                          x=x,
89
                                                         j=i ,
90
                                                /self.eta_matrix[:, j, i] = self.calculate_eta(xi_new_ji)free_energy.append(
91
92
                                                          self.compute_free_energy(x, binary_latent_factor_model)
```

```
return free_energy
97
           {\tt def} \ \ {\tt calculate\_binary\_message\_update} \, (
 98
                 self,
 99
                    np.ndarray
                 boltzmann_machine: BoltzmannMachine,
101
                 i: int ,
j: int ,
           ) -> float:
                 Calculate new parameters for a binary factored message.
106
                 :param x: data matrix (number_of_points, number_of_dimensions)
                 :param boltzmann_machine: Boltzmann machine model :param i: starting node for the message :param j: ending node for the message
108
109
110
                 :return: new parameter from aggregating incoming messages
                 eta_i_not_j = boltzmann_machine.b_index(
    x=x, node_index=i
) + self.aggregate_incoming_binary_factor_messages(
113
114
116
                      node_index=i, excluded_node_index=j
117
                 119
120
           {\tt def} \ \ {\tt calculate\_singleton\_message\_update} \, (
123
                 x: np.ndarrav
124
                 boltzmann_machine: BoltzmannMachine,
           i: int,
) -> float:
126
127
                 Calculate the parameter update for the singleton message. Note that this does not require any approximation.
128
130
                 :param x: data matrix (number_of_points, number_of_dimensions)
:param boltzmann_machine: Boltzmann machine model
                 :param i: node to update
:return: new parameter
134
135
136
                 {\tt return boltzmann\_machine.b\_index(x=x, node\_index=i)}
      \label{eq:def_def} \begin{array}{ll} def & init\_message\_passing(k:\ int\ ,\ n:\ int\ ) \ -\!\!\!> \ MessagePassingApproximation: \end{array}
140
           Message passing initialisation
142
            :param k: number of latent variables
144
            :param n: number of data points
           :return: message passing
145
           eta_matrix = np.random.random(size=(n, k, k))
return MessagePassingApproximation(eta_matrix)
147
148
```

src/models/binary_latent_factor_approximations/message_passing_approximation.py

The rest of the Python code for question 6:

```
import matplotlib.pyplot as plt import numpy as np
 3
4
       from src.expectation_maximisation import learn_binary_factors
       {\bf from } \  \  {\bf src.models.binary\_latent\_factor\_approximations.message\_passing\_approximation \ \ \underline{\bf import} \ \ (
             \verb"init_message_passing",
      from src.models.binary_latent_factor_models.boltzmann_machine import ( init_boltzmann_machine ,
10
11
12
13
14
       def run(x: np.ndarray, k: int, em_iterations: int, save_path: str) -> None:
             n = x.shape[0]
message\_passing = init\_message\_passing(k, n)
15
16
17
             boltzmann\_machine = init\_boltzmann\_machine(x, message\_passing)
18
19
             # pre-training features plot
fig , ax = plt.subplots(1, k, figsize=(k * 2, 2))
for i in range(k):
20
             ax[i].imshow(boltzmann_machine.mu[:, i].reshape(4, 4))
ax[i].set_title(f"Latent Feature mu-{i}")
fig.suptitle("Initial Features (Loopy BP)")
plt_tight_layout()
21
22
23
24
25
             plt.savefig(save_path + "-init-latent-factors", bbox_inches="tight")
26
             plt.close()
27
28
29
             message_passing, boltzmann_machine, free_energy = learn_binary_factors(
30
31
                    em_iterations=em_iterations
                    binary_latent_factor_model=boltzmann_machine,
32
33
34
                    \verb|binary_latent_factor_approximation=message\_passing|,
35
            # post training features plot
fig , ax = plt.subplots(1, k, figsize=(k * 2, 2))
for i in range(k):
    ax[i].imshow(boltzmann_machine.mu[:, i].reshape(4, 4))
    ax[i].set_title(f"Latent Feature mu_{i})
fig.suptitle("Learned Features (Loopy BP)")
plt.tight_layout()
plt.savefig(save_path + "-latent-factors", bbox_inches="tight")
plt_close()
36
38
41
42
44
             plt.close()
45
            # free energy plot
plt.title("Free Energy (Loopy BP)")
plt.xlabel("t (EM steps)")
plt.ylabel("Free Energy")
47
48
49
             plt.plot(free_energy)
plt.savefig(save_path + "-free-energy", bbox_inches="tight")
50
51
             plt.close()
```

src/solutions/q6.py

Appendix 1: abstract_binary_latent_factor_model.py

```
from __future__ import annotations
      from abc import ABC, abstractmethod
      from typing import TYPE_CHECKING
     import numpy as np
           from \  \  src.models.binary\_latent\_factor\_approximations.abstract\_binary\_latent\_factor\_approximation \  \  import \  \  (abstract\_binary\_latent\_factor\_approximation \  \  import \  \  )
10
                AbstractBinaryLatentFactorApproximation,
12
13
      {\tt class} \quad AbstractBinaryLatentFactorModel (ABC):
15
16
           @property
@abstractmethod
17
18
           def mu(self) \rightarrow np.ndarray:
19
                 matrix of means (number_of_dimensions, number_of_latent_variables)
20
21
                pass
23
24
           @\,property
           @abstractmethod
           def variance(self) -> float:
26
                gaussian noise parameter
29
30
31
           @property
           @abstractmethod
           def pi(self) -> np.ndarray:
34
35
                (1, number_of_latent_variables)
37
38
39
           @\,abstractmethod
40
           def maximisation_step (
41
                 self,
                 binary\_latent\_factor\_approximation: \ AbstractBinaryLatentFactorApproximation \ ,
43
44
           ) -> None:
46
           def mu_exclude(self, exclude_latent_index: int) -> np.ndarray:
    return np.concatenate( # (number_of_dimensions, number_of_latent_variables -1)
        (self.mu[:, :exclude_latent_index], self.mu[:, exclude_latent_index + 1 :]),
47
48
49
                      \dot{a} \times i s = 1,
51
52
           def log_pi(self) -> np.ndarray:
    return np.log(self.pi)
54
57
58
           def log_one_minus_pi(self) -> np.ndarray:
                return np.log(1 - self.pi)
60
           def precision(self) -> float:
    return 1 / self.variance
62
63
           @property
def d(self) -> int:
65
66
                return self.mu.shape[0]
68
69
           @property
70
71
           def k(self) -> int:
                return self.mu.shape[1]
```

src/models/binary_latent_factor_models/abstract_binary_latent_factor_model.py

Appendix 2: abstract_binary_latent_factor_approximation.py

```
from __future__ import annotations
      from abc import ABC, abstractmethod
      from typing import TYPE_CHECKING, List
      if TYPE_CHECKING:
           from src.models.binary_latent_factor_models.binary_latent_factor_model import (
                 AbstractBinaryLatentFactorModel,
      import numpy as np
12
13
14
             AbstractBinaryLatentFactorApproximation (ABC):\\
           @property
@abstractmethod
16
17
18
           \begin{array}{ll} d\,ef & lambda\_matrix\,(\,s\,elf\,) \,\, -\!\!> \,\, np\,.\,ndarray\,; \end{array}
19
                 lambda_matrix: parameters variational approximation (number_of_points, number_of_latent_variables)
20
21
                 pass
23
           @abstractmethod
24
           def variational_expectation_step(
26
                 x: np.ndarray
                 binary\_latent\_factor\_model: \ AbstractBinaryLatentFactorModel \, ,
29
                 pass
30
31
           def expectation_s(self):
    return self.lambda_matrix
34
35
           def expectation_ss(self):
37
38
                 ess = self.lambda_matrix.T @ self.lambda_matrix
np.fill_diagonal(ess, self.lambda_matrix.sum(axis=0))
40
41
           @property
           def log.lambda_matrix(self) -> np.ndarray:
    return np.log(self.lambda_matrix)
43
44
           \begin{array}{lll} \textbf{def} & \textbf{log\_one\_minus\_lambda\_matrix} \, (\, \, \textbf{self} \, ) \, \, \, \rightarrow \, \, \textbf{np.ndarray} \, ; \\ \end{array}
46
                 return np.log(1 - self.lambda_matrix)
48
49
           @property
           def n(self) \rightarrow int:
51
52
                 Number of data points
54
                 {\tt return self.lambda\_matrix.shape} \, [\, 0 \, ]
57
58
           def k(self) \rightarrow int:
                 Number of latent variables
60
                 {\tt return self.lambda\_matrix.shape} \, [\, 1\, ]
62
           def compute_free_energy(
                 x: np.ndarray,
binary_latent_factor_model: AbstractBinaryLatentFactorModel,
65
66
           ) -> float:
68
69
                 free energy associated with current EM parameters and data x
70
71
72
                 :param x: data matrix (number_of_points, number_of_dimensions)
:param binary_latent_factor_model: a binary_latent_factor_model
73
74
75
76
77
78
                 return: average free energy per data point:
                 expectation_log_p_x_s_given_theta = (
                       self._compute_expectation_log_p_x_s_given_theta(
    x, binary_latent_factor_model
79
                 approximation_model_entropy = self._compute_approximation_model_entropy()
82
                  \begin{array}{l} {\rm expectation\_log\_p\_x\_s\_given\_theta} \ + \ {\rm approximation\_model\_entropy} \\ ) \ / \ {\rm self.n} \end{array} 
85
           \textcolor{red}{\textbf{def}} \hspace{0.2cm} \texttt{\_compute\_expectation\_log\_p\_x\_s\_given\_theta} \hspace{0.1cm} (
88
                 binary_latent_factor_model: AbstractBinaryLatentFactorModel,
90
                 The first term of the free energy, the expectation of log P(X,S \mid \text{theta})
91
```

```
:param x: data matrix (number_of_points, number_of_dimensions) :param binary_latent_factor_model: a binary_latent_factor_model :return: the expectation of log P(X,S \mid theta) """
 95
 96
                 # (number_of_points, number_of_dimensions)
                 mu_lambda = self.lambda_matrix @ binary_latent_factor_model.mu.T
98
 99
100
                 # (number_of_latent_variables, number_of_latent_variables)
                 expectation_s_i_s_j_mu_i_mu_j = np.multiply(
self.lambda_matrix.T @ self.lambda_matrix
                       \verb|binary_latent_factor_model.mu.T @ binary_latent_factor_model.mu|,
104
                 106
107
108
                      0.5 \ * \ binary\_latent\_factor\_model.precision
110
                      np.sum(np.multiply(x, x))
- 2 * np.sum(np.multiply(x, mu_lambda))
+ np.sum(expectation_s_i_s_j_mu_i_mu_j)
113
114
                         np.trace(
                      expectation_s_i_s_j_mu_i_mu_j
) # remove incorrect E[s_i s_i] = lambda_i * lambda_i
+ np.sum( # add correct E[s_i s_i] = lambda_i
self.lambda_matrix
116
117
118
                            @ np.multiply(
120
                                  binary\_latent\_factor\_model.mu, \ binary\_latent\_factor\_model.mu
121
                 expectation_log_p_s_given_theta = np.sum(
124
125
                      np.multiply(
                            self.lambda_matrix,
binary_latent_factor_model.log_pi,
126
128
                      + np.multiply(
1 - self.lambda_matrix,
                            binary_latent_factor_model.log_one_minus_pi,
133
134
                 return expectation_log_p_x_given_s_theta + expectation_log_p_s_given_theta
135
           {\tt def\_compute\_approximation\_model\_entropy(self)} \; -\!\!\!> \; {\tt float}:
138
                 Compute the model entropy
139
                 :return: model entropy
140
142
                 {\tt return -np.sum}\,(
143
                      np.multiply(
                            self.lambda_matrix,
145
                            self.log\_lambda\_matrix,
146
                      + np. multiply (
                            1 - self.lambda_matrix,
self.log_one_minus_lambda_matrix,
148
149
```

src/models/binary_latent_factor_approximations/abstract_binary_latent_factor_approximation.py

Appendix 3: main.py

```
import os
from dataclasses import asdict
      import jax
      import jax numpy as jnp
import numpy as np
      import pandas as pd
      from src.constants import CO2_FILE_PATH, DEFAULT_SEED, OUTPUTS_FOLDER
      from src.generate_images import generate_images from src.models.bayesian_linear_regression import LinearRegressionParameters
      \begin{array}{lll} from & \verb|src.models.gaussian.process.regression| import| Gaussian Process Parameters \\ from & \verb|src.models.kernels| import| Combined Kernel,| Combined Kernel Parameters \\ \end{array}
13
      from src.solutions import q2, q3, q4, q6
16
      jax.config.update("jax_enable_x64", True)
17
18
      if __name__ == "__main
19
           np.random.seed (DEFAULT_SEED)
20
21
           if not os.path.exists(OUTPUTS.FOLDER):
    os.makedirs(OUTPUTS.FOLDER)
23
24
           if not os.path.exists(Q2_OUTPUT_FOLDER, "q2")
os.makedirs(Q2_OUTPUT_FOLDER):
26
            with open (CO2_FILE_PATH) as file
                  lines = [line.rstrip().split() for line in file]
29
30
           df_co2 = pd.DataFrame(
    np.array([line for line in lines if line[0] != "#"]).astype(float)
31
34
            column_names = lines [max([i for i, line in enumerate(lines) if line[0] == "#"])][1:]
           \begin{array}{ll} df\_co2.columns = column\_names \\ t = df\_co2.decimal.values [:] - np.min(df\_co2.decimal.values [:]) \\ y = df\_co2.average.values [:].reshape(1, -1) \end{array}
35
37
38
39
40
           mean \, = \, np.\,array\,(\,[\,0\;,\ 3\,6\,0\,]\,)\;.\,reshape\,(\,-1\,,\ 1\,)
41
           covariance = np.array(
                        [10**2, 0]
43
                        [0, 100**2],
44
46
47
            kernel = CombinedKernel()
48
            \label{eq:kernel-parameters} \begin{aligned} kernel\_parameters &= CombinedKernelParameters (\\ log\_theta=&jnp.log (1) \;, \end{aligned}
49
                  log_sigma=jnp.log(1),
50
51
52
                  \log_{-p} hi = jnp \cdot \log(1)
                  log_eta=jnp.log(1),
53
                  log_tau=jnp.log(1)
54
                  \log_{-}z \operatorname{et} a = \operatorname{jnp} \cdot \log (1 \operatorname{e} - 1),
56
57
58
            \verb|prior_linear_regression_parameters| = LinearRegressionParameters| (
                  mean=mean,
                  covariance=covariance,
60
            posterior_linear_regression_parameters = q2.a(
62
                 prior_linear_regression_parameters,
save_path=os.path.join(Q2_OUTPUT_FOLDER, "a"),
65
66
68
           q2.b(
69
                  t_year=df_co2.decimal.values[:],
70
71
                  t=t,
                  v=v.
                  linear_regression_parameters=posterior_linear_regression_parameters,
                  error_mean=0,
                  error_variance=1.
                  save_path=os.path.join(Q2_OUTPUT_FOLDER, "b"),
76
77
78
79
                  kernel=kernel
80
                  kernel_parameters=kernel_parameters,
                  log\_theta\_range=jnp.log(jnp.linspace(1e-2, 5, 5)),
82
                  t=t [:50]. reshape(-1, 1),
number_of_samples=3,
                  {\tt save\_path$=$os.path.join} \; ( {\tt Q2\_OUTPUT\_FOLDER}, \;\;"c") \; ,
85
            init\_kernel\_parameters = Combined Kernel Parameters (
                 log_theta=jnp.log(5),
log_sigma=jnp.log(5),
log_phi=jnp.log(10),
log_eta=jnp.log(5),
88
90
91
                  log_tau=jnp.log(1),
```

```
\log_z z e t a = j n p \cdot \log(2),
 94
            gaussian_process_parameters = GaussianProcessParameters(
 95
 96
                 kernel=asdict(init_kernel_parameters),
                 log_sigma=jnp.log(1),
 97
 98
 99
           /years_to_predict = 14
t_new = t[-1] + np.linspace(0, years_to_predict, years_to_predict * 12)
100
            t_{test} = np.concatenate((t, t_{new}))
           q2.f(
                 \dot{t}_{-}train=t,
                 y_train=y,
t_test=t_test,
104
105
106
                 \min_{y \in ar = np. min(df_{co2}.decimal.values[:])}
                 prior_linear_regression_parameters=prior_linear_regression_parameters,
107
108
                 linear_regression_sigma=sigma,
                 kernel=kernel,
110
                 gaussian_process_parameters=gaussian_process_parameters,
                 learning_rate=1e-2,
112
                 number_of_iterations=100.
                 save_path=os.path.join(Q2_OUTPUT_FOLDER, "f"),
113
114
115
116
           g3.OUTPUT.FOLDER = os.path.join(OUTPUTS.FOLDER, "q3")
if not os.path.exists(Q3.OUTPUT.FOLDER):
    os.makedirs(Q3.OUTPUT.FOLDER)
118
120
           number_of_images = 2000
           \begin{array}{ll} x = \text{generate\_images} (n=\text{number\_of\_images}) \\ k = 8 \end{array}
121
           {\tt em\_iterations} \, = \, 100
124
           e_maximum_steps = 50
            e_convergence_criterion = 0
126
127
            binary_latent_factor_model = q3.e_and_f(
128
                 k=k
130
                 em_iterations=em_iterations,
                 e_maximum_steps=e_maximum_steps,
e_convergence_criterion=e_convergence_criterion,
                 save_path=os.path.join(Q3_OUTPUT_FOLDER, "f"),
134
            _ = q3.e_and_f(
135
136
                  \begin{array}{l} x=x \,, \\ k=int \, (k \ * \ 1.5) \,, \\ em\_iterations=em\_iterations \,, \end{array} 
138
139
                 e\_maximum\_steps = e\_maximum\_steps \;,
                 e_convergence_criterion=e_convergence_criterion ,
save_path=os.path.join(Q3_OUTPUT_FOLDER, "f-larger-k"),
140
141
142
           q3.g(
143
                 \verb|binary_latent_factor_model| = \verb|binary_latent_factor_model| ,
145
146
                 sigmas = [1, 2, 3],
                 em_iterations=em_iterations
148
149
                 e_maximum_steps=e_maximum_steps,
150
                 e_convergence_criterion=e_convergence_criterion ,
                 \verb|save_path| = \verb|os.path.join| (Q3\_OUTPUT\_FOLDER, "g") \;,
152
153
           # Question 4
           Q4_OUTPUT_FOLDER = os.path.join(OUTPUTS_FOLDER, "q4")
            if not os.path.exists(Q4_OUTPUT_FOLDER):
    os.makedirs(Q4_OUTPUT_FOLDER)
156
           q4.b(
                 x=x,
                 a-parameter=1,
162
                 b_parameter = 0,
                 ks=np.arange(4, 22),
163
164
                 max_k=max_k,
                 em_iterations=em_iterations .
166
                 e_maximum_steps=e_maximum_steps,
                 e_convergence_criterion=e_convergence_criterion, save_path=os.path.join(Q4_OUTPUT_FOLDER, "b"),
167
168
170
171
            q4.b(
                 x=x.
                 a_parameter=1,
\frac{173}{174}
                 b_parameter = 0,
                 ks=np.arange(4, 13),
175
                 max_k=max_k,
\frac{176}{177}
                 em_iterations=em_iterations,
e_maximum_steps=e_maximum_steps,
                 e_convergence_criterion=e_convergence_criterion, save_path=os.path.join(Q4_OUTPUT_FOLDER, "b-1"),
178
179
180
181
            q4.b(
                 x=x.
182
183
                 a-parameter=1,
184
                 b_parameter=0
                 ks=np.arange(13, 22),
185
                 max_k=max_k,
187
                 em_iterations=em_iterations
188
                 e_maximum_steps=e_maximum_steps.
```

main.py

Appendix 4: constants.py

```
import os

DATAFOLDER = "data"

CO2_FILE_PATH = os.path.join(DATA_FOLDER, "co2.txt")
IMAGES_FILE_PATH = os.path.join(DATA_FOLDER, "images.jpg")

OUTPUTS_FOLDER = "outputs"

DEFAULT_SEED = 0

II

M1 = [0, 0, 1, 0, 0, 1, 1, 1, 0, 0, 1, 0, 0, 0, 0]

M2 = [0, 1, 0, 0, 0, 1, 0, 0, 0, 1, 0, 0, 0, 0, 0]

M3 = [1, 1, 1, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0]

M4 = [1, 0, 0, 0, 0, 1, 0, 0, 0, 1, 0, 0, 0, 1, 1, 0, 0]

M5 = [0, 0, 0, 0, 0, 0, 0, 1, 1, 0, 0, 1, 1, 1, 1]

M6 = [1, 1, 1, 1, 0, 0, 1, 1, 0, 0, 1, 1, 1, 1]

M7 = [0, 0, 0, 0, 0, 1, 1, 0, 0, 1, 1, 0, 0, 0, 0]

M8 = [0, 0, 0, 1, 0, 0, 0, 1, 0, 0, 0, 1, 0, 0, 0, 0]
```

src/constants.py

Appendix 5: generate_images.py

```
import numpy as np
      from src.constants import DEFAULT_SEED, M1, M2, M3, M4, M5, M6, M7, M8
5
6
7
8
9
10
11
12
13
14
15
16
       def generate_images(n: int = 400, seed: int = DEFAULT_SEED, sigma: float = 0.1):
            Image generation, adapted from provided demo code
            :param n: number of data points
:param seed: random seed
:param sigma: Gaussian noise
:return: images as a data matrix (number_of_points, number_of_dimensions)
"""
            d=16 # dimensionality of the data
            np.random.seed(seed)
17
18
19
            \# Define the basic shapes of the features number\_of\_features\,=\,8\, \# number of features
20
21
22
23
24
25
26
27
28
29
30
31
32
33
            rr [0] * M1,
rr [1] * M2,
rr [2] * M3,
rr [3] * M4,
rr [4] * M5,
rr [5] * M6,
rr [6] * M7,
rr [7] * M8,
                        rr [0]
rr [1]
rr [2]
rr [3]
rr [4]
rr [5]
rr [6]
rr [7]
34
35
36
                  np.random.rand(n, number_of_features) < 0.3
            ) # each feature occurs with prob 0.3 independently
            # Generate Data - The Data is stored in Y
\frac{40}{41}
                np.dot(s, mut) + np.random.randn(n, d) * sigma # some Gaussian noise is added
```

src/generate_images.py

Appendix 6: MStep.py

```
import numpy as np
  1
2
3
4
5
6
7
8
9
              def m_step(x, es, ess):
                         mu, sigma, pie = MStep(x, es, ess)
                                          x: shape (n, d) data matrix es: shape (n, k) E_{-q}[s] ess: shape (k, k) sum over data points of E_{-q}[ss'] (n, k, k) if E_{-q}[ss'] is provided, the sum over n is done for you.
10
11
12
13
14
15
16
                         Outputs:
                         mu: shape (d, k) matrix of means in p(y|\{s_i\},mu,sigma) sigma: shape (,) standard deviation in same pie: shape (1, k) vector of parameters specifying generative distribution for s """
17
18
 19
20
21
                         n, d = x.shape
if es.shape[0] != n:
    raise TypeError('es must have the same number of rows as x')
k = es.shape[1]
if ess.shape == (n, k, k):
    ess = np.sum(ess, axis=0)
if ess.shape != (k, k):
    raise TypeError('ess must be square and have the same number of columns as es')
23
24
25
26
27
28
29
                         \begin{array}{lll} mu = np. \det (np. \det (np. \operatorname{linalg.inv}(\operatorname{ess})\,,\, \operatorname{es.T})\,,\, x)\,.T \\ sigma = np. \operatorname{sqrt}\left((np. \operatorname{trace}(np. \det (x.T,\, x))\,+\, np. \operatorname{trace}(np. \det (\operatorname{np. \det}(\operatorname{mu.T},\, \operatorname{mu})\,,\, \operatorname{ess})\right) \\ & - 2 \,*\, np. \operatorname{trace}(\operatorname{np. \det}(\operatorname{np. \det}(\operatorname{es.T},\, x)\,,\, \operatorname{mu}))) \,\,/\, \, (n \,*\, d)) \\ pie = np. \operatorname{mean}(\operatorname{es}\,,\, \operatorname{axis} = 0,\, \operatorname{keepdims} = \operatorname{True}) \end{array}
32
                           return mu, sigma, pie
```

demo_code/MStep.py