## Assignment4\_9628\_jzeiders

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## 1 John Zeiders (jzeiders) - Assignment 4

```
[]: import numpy as np
from numpy.linalg import inv, det
from scipy.stats import multivariate_normal
import pandas as pd
```

## 2 Part 1

```
[]: def Estep(data, prob, mean, Sigma):
         E-step: Calculate responsibilities
         Parameters:
         data: ndarray, shape (n, p)
             The input data matrix
         prob: ndarray, shape (G,)
             Mixing proportions
         mean: ndarray, shape (p, G)
             Mean vectors for each component
         Sigma: ndarray, shape (p, p)
             Shared covariance matrix
         Returns:
         responsibilities: ndarray, shape (n, G)
             Matrix of posterior probabilities P(Z_i=k/x_i)
         n = data.shape[0]
         G = prob.shape[0]
         # Initialize matrix to store densities
         densities = np.zeros((n, G))
         # Compute gaussian densities for each component
         for k in range(G):
```

```
densities[:, k] = multivariate_normal.pdf(data, mean=mean[:, k],__
 ⇔cov=Sigma)
    # Multiply by mixing proportions
    weighted_densities = densities * prob[np.newaxis, :]
    # Normalize to get responsibilities
    responsibilities = weighted_densities / np.sum(weighted_densities, axis=1)[:
 →, np.newaxis]
    return responsibilities
def Mstep(data, responsibilities):
    11 11 11
    M-step: Update parameters
    Parameters:
    data: ndarray, shape (n, p)
        The input data matrix
    responsibilities: ndarray, shape (n, G)
        Matrix of posterior probabilities
    Returns:
    _____
    prob: ndarray, shape (G,)
        Updated mixing proportions
    mean: ndarray, shape (p, G)
        Updated mean vectors
    Sigma: ndarray, shape (p, p)
        Updated shared covariance matrix
    n, p = data.shape
    G = responsibilities.shape[1]
    # Update mixing proportions
    prob = np.mean(responsibilities, axis=0)
    # Update means
    mean = np.zeros((p, G))
    for k in range(G):
        mean[:, k] = np.sum(responsibilities[:, k:k+1] * data, axis=0) / np.
 ⇔sum(responsibilities[:, k])
    # Update shared covariance matrix
    Sigma = np.zeros((p, p))
    for k in range(G):
```

```
diff = data - mean[:, k] # Shape: (n, p)
        weighted_diff = responsibilities[:, k].reshape(n, 1) * diff # Shape:
 \hookrightarrow (n, p)
        Sigma += weighted_diff.T @ diff # Shape: (p, p)
    Sigma /= n
    return prob, mean, Sigma
def loglik(data, prob, mean, Sigma):
    Compute log-likelihood
    Parameters:
    _____
    data: ndarray, shape (n, p)
        The input data matrix
    prob: ndarray, shape (G,)
       Mixing proportions
    mean: ndarray, shape (p, G)
       Mean vectors
    Sigma: ndarray, shape (p, p)
        Shared covariance matrix
    Returns:
    _____
    ll: float
       Log-likelihood value
   n = data.shape[0]
    G = prob.shape[0]
    # Initialize array for component densities
    densities = np.zeros((n, G))
    # Compute densities for each component
    for k in range(G):
        densities[:, k] = prob[k] * multivariate_normal.pdf(data, mean=mean[:, u
 →k], cov=Sigma)
    # Sum over components and take log
    11 = np.sum(np.log(np.sum(densities, axis=1)))
    return 11
def myEM(data, G, initial_params, itmax):
    Main EM algorithm function
```

```
Parameters:
_____
data: ndarray, shape (n, p)
    The input data matrix
G: int
   Number of components
initial_params: dict
    Dictionary containing initial parameters:
    'prob': mixing proportions
    'mean': mean vectors
    'Sigma': covariance matrix
itmax: int
    Maximum number of iterations
Returns:
_____
prob: ndarray, shape (G,)
   Final mixing proportions
mean: ndarray, shape (p, G)
   Final mean vectors
Sigma: ndarray, shape (p, p)
   Final shared covariance matrix
ll: float
   Final log-likelihood value
# Initialize parameters
prob = initial_params['prob']
mean = initial_params['mean']
Sigma = initial_params['Sigma']
for iteration in range(itmax):
    # E-step
   responsibilities = Estep(data, prob, mean, Sigma)
    # M-step
   prob, mean, Sigma = Mstep(data, responsibilities)
    # Optionally, compute log-likelihood for monitoring
   11 = loglik(data, prob, mean, Sigma)
   print(f"Iteration {iteration+1}, Log-Likelihood: {11:.3f}")
# Compute final log-likelihood
11 = loglik(data, prob, mean, Sigma)
return prob, mean, Sigma, 11
```

```
[]: def test_G2(data):
        """Test EM algorithm with G=2 components"""
        n, p = data.shape
         # Set initial parameters as specified
        p1 = 10/n
        prob = np.array([p1, 1-p1])
        mean = np.zeros((p, 2))
        mean[:, 0] = np.mean(data[:10], axis=0) # mean of first 10 samples
        mean[:, 1] = np.mean(data[10:], axis=0) # mean of remaining samples
        # Calculate initial Sigma
        diff1 = data[:10] - mean[:, 0]
        diff2 = data[10:] - mean[:, 1]
        Sigma = (diff1.T @ diff1 + diff2.T @ diff2) / n
        initial_params = {
            'prob': prob,
             'mean': mean,
             'Sigma': Sigma
        }
        # Run EM algorithm
        final_prob, final_mean, final_Sigma, final_ll = myEM(data, 2,_
      ⇒initial params, 20)
        return final_prob, final_mean, final_Sigma, final_ll
    def test_G3(data):
         """Test EM algorithm with G=3 components"""
        n, p = data.shape
        # Set initial parameters
        p1, p2 = 10/n, 20/n
        prob = np.array([p1, p2, 1-p1-p2])
        mean = np.zeros((p, 3))
        mean[:, 0] = np.mean(data[:10], axis=0) # mean of first 10 samples
        mean[:, 1] = np.mean(data[10:30], axis=0) # mean of next 20 samples
        mean[:, 2] = np.mean(data[30:], axis=0) # mean of remaining samples
        # Calculate initial Sigma
        diff1 = data[:10] - mean[:, 0]
        diff2 = data[10:30] - mean[:, 1]
        diff3 = data[30:] - mean[:, 2]
        Sigma = (diff1.T @ diff1 + diff2.T @ diff2 + diff3.T @ diff3) / n
```

```
initial_params = {
             'prob': prob,
             'mean': mean,
             'Sigma': Sigma
         }
         # Run EM algorithm
         final_prob, final_mean, final_Sigma, final_ll = myEM(data, 3,_

initial_params, 20)

         return final_prob, final_mean, final_Sigma, final_ll
[]: | # Load faithful dataset
     file_path = 'https://liangfgithub.github.io/Data/faithful.dat' # Update this_{\sqcup}
      ⇒path as necessary
     data = np.loadtxt(file_path, skiprows=1, dtype=float, usecols=(1,2))
     print("Data Shape:", data.shape)
     print("First 5 Data Points:\n", data[:5])
    Data Shape: (272, 2)
    First 5 Data Points:
     [[ 3.6
             79. 1
     Γ 1.8
             54.

√ 3.333 74.

                   1
     [ 2.283 62.
                   1
     [ 4.533 85.
                   11
[]: # Test G=2
     print("\nTesting with G=2:")
     prob2, mean2, Sigma2, 112 = test_G2(data)
     print("\nprob")
     print(prob2)
     print("\nmean")
     print(mean2)
     print("\nSigma")
     print(Sigma2)
     print("\nloglik")
     print(112)
    Testing with G=2:
    Iteration 1, Log-Likelihood: -1289.766
    Iteration 2, Log-Likelihood: -1289.762
    Iteration 3, Log-Likelihood: -1289.758
    Iteration 4, Log-Likelihood: -1289.753
    Iteration 5, Log-Likelihood: -1289.747
```

```
Iteration 6, Log-Likelihood: -1289.741
    Iteration 7, Log-Likelihood: -1289.734
    Iteration 8, Log-Likelihood: -1289.726
    Iteration 9, Log-Likelihood: -1289.718
    Iteration 10, Log-Likelihood: -1289.708
    Iteration 11, Log-Likelihood: -1289.698
    Iteration 12, Log-Likelihood: -1289.686
    Iteration 13, Log-Likelihood: -1289.674
    Iteration 14, Log-Likelihood: -1289.661
    Iteration 15, Log-Likelihood: -1289.647
    Iteration 16, Log-Likelihood: -1289.632
    Iteration 17, Log-Likelihood: -1289.617
    Iteration 18, Log-Likelihood: -1289.601
    Iteration 19, Log-Likelihood: -1289.585
    Iteration 20, Log-Likelihood: -1289.569
    prob
    [0.04297883 0.95702117]
    mean
    [76.79789154 70.63205853]]
    Sigma
    [[ 1.29793612 13.92433626]
     [ 13.92433626 182.58009247]]
    loglik
    -1289.5693549424104
[]: # Test G=3
    print("\nTesting with G=3:")
    prob3, mean3, Sigma3, 113 = test_G3(data)
    print("\nprob")
    print(prob3)
    print("\nmean")
    print(mean3)
    print("\nSigma")
    print(Sigma3)
    print("\nloglik")
    print(113)
    Testing with G=3:
    Iteration 1, Log-Likelihood: -1289.750
    Iteration 2, Log-Likelihood: -1289.744
    Iteration 3, Log-Likelihood: -1289.738
```

```
Iteration 4, Log-Likelihood: -1289.730
    Iteration 5, Log-Likelihood: -1289.722
    Iteration 6, Log-Likelihood: -1289.712
    Iteration 7, Log-Likelihood: -1289.701
    Iteration 8, Log-Likelihood: -1289.688
    Iteration 9, Log-Likelihood: -1289.674
    Iteration 10, Log-Likelihood: -1289.658
    Iteration 11, Log-Likelihood: -1289.640
    Iteration 12, Log-Likelihood: -1289.619
    Iteration 13, Log-Likelihood: -1289.596
    Iteration 14, Log-Likelihood: -1289.571
    Iteration 15, Log-Likelihood: -1289.542
    Iteration 16, Log-Likelihood: -1289.511
    Iteration 17, Log-Likelihood: -1289.476
    Iteration 18, Log-Likelihood: -1289.438
    Iteration 19, Log-Likelihood: -1289.396
    Iteration 20, Log-Likelihood: -1289.351
    prob
    [0.04363422 0.07718656 0.87917922]
    mean
    [[ 3.51006918  2.81616674  3.54564083]
     [77.10563811 63.35752634 71.25084801]]
    Sigma
    [[ 1.26015772 13.51153756]
     [ 13.51153756 177.96419105]]
    loglik
    -1289.350958862739
       Part 2
    3
[]: import numpy as np
     from numpy.linalg import inv, det
     from scipy.stats import multivariate_normal
     from io import StringIO
     import unittest
     from numpy.testing import assert_array_almost_equal, assert_array_less
     import pandas as pd
[]: def forward_pass(data, w, A, B):
         Compute forward probabilities alpha(t,j)
```

Args:

```
data: T-by-1 observation sequence (1D array)
        w: Initial state distribution (mz,)
        A: Transition matrix (mz-by-mz)
        B: Emission matrix (mz-by-mx)
    Returns:
        alpha: Forward probabilities (T-by-mz)
    11 11 11
    T = len(data)
    mz = len(w)
    alpha = np.zeros((T, mz))
    # Initialize first time step
    alpha[0, :] = w * B[:, data[0]]
    # Forward recursion
    for t in range(1, T):
        alpha[t, :] = np.dot(alpha[t - 1, :], A) * B[:, data[t]].T
    return alpha
def backward_pass(data, A, B):
    Compute backward probabilities beta(t, j)
    Args:
        data: T-by-1 observation sequence (1D array)
        A: Transition matrix (mz-by-mz)
        B: Emission matrix (mz-by-mx)
    Returns:
        beta: Backward probabilities (T-by-mz)
    11 11 11
    T = len(data)
    mz = A.shape[0]
    beta = np.zeros((T, mz))
    # Initialize last time step
    beta[-1, :] = 1
    # Backward recursion
    for t in range(T - 2, -1, -1):
        beta[t, :] = np.dot(A, (B[:, data[t + 1]] * beta[t + 1, :]))
    return beta
def BW_onestep(data, w, A, B):
```

```
One step of the Baum-Welch algorithm (E-step + M-step)
  Arqs:
       data: T-by-1 observation sequence (1D array)
      w: Initial state distribution (mz,)
      A: Current transition matrix (mz-by-mz)
      B: Current emission matrix (mz-by-mx)
  Returns:
      A new: Updated transition matrix
      B_new: Updated emission matrix
  T = len(data)
  mz = A.shape[0]
  mx = B.shape[1]
  # E-step: Compute forward and backward probabilities
  alpha = forward_pass(data, w, A, B)
  beta = backward_pass(data, A, B)
  # Compute xi(t,i,j) = P(z_t=i, z_{t+1}=j | x_{1}:T)
  xi = np.zeros((T - 1, mz, mz))
  for t in range(T - 1):
       denominator = np.dot(np.dot(alpha[t, :], A) * B[:, data[t + 1]], beta[t_{\square}]
+ 1, :])
      numerator = alpha[t, :, np.newaxis] * A * B[:, data[t + 1]] * beta[t +__
\hookrightarrow 1, :]
      xi[t, :, :] = numerator / denominator
  # Compute state probabilities (gamma)
  gammas_j = np.zeros((T, mz))
  gammas_j[:-1] = np.sum(xi, axis=2)
  # Fix for last time step - should use alpha and beta
  gammas_j[-1] = (alpha[-1] * beta[-1]) / np.sum(alpha[-1] * beta[-1])
  # M-step: Update parameters
  # Update A
  A_new = np.zeros_like(A)
  for i in range(mz):
      for j in range(mz):
           numerator = np.sum(xi[:, i, j])
           denominator = np.sum(xi[:, i, :])
           A_new[i, j] = numerator / denominator
  # Update B (vectorized version)
  B_new = np.zeros_like(B)
```

```
for 1 in range(mx):
        mask = (data == 1)
        B_new[:, 1] = np.sum(gammas_j[mask], axis=0) / (np.sum(gammas_j,_
 \Rightarrowaxis=0) + 1e-300)
    # Verify row stochasticity
    assert np.allclose(np.sum(A_new, axis=1), 1, rtol=1e-5), "A matrix not row_
 ⇔stochastic"
    assert np.allclose(np.sum(B_new, axis=1), 1, rtol=1e-5), "B matrix not row_
 \hookrightarrowstochastic"
    return A_new, B_new
def myBW(data, initial_params, itmax):
    Main Baum-Welch algorithm with specified initial parameters
    Arqs:
        data: T-by-1 observation sequence (1D array)
        initial_params: Dictionary containing initial parameters:
            'w': Initial state distribution (mz,)
            'A': Initial transition matrix (mz-by-mz)
            'B': Initial emission matrix (mz-by-mx)
        itmax: Maximum number of iterations
    Returns:
        A final: Final transition matrix
        B_final: Final emission matrix
        ll_final: Final log likelihood
    n n n
    # Ensure data is a 1D array
    data = np.asarray(data).flatten()
    # Extract initial parameters
    w = initial_params["w"]
    A = initial_params["A"]
    B = initial_params["B"]
    for iteration in range(itmax):
        # Compute log likelihood
        alpha = forward_pass(data, w, A, B)
        current_ll = np.log(np.sum(alpha[-1]))
        # Update parameters
        A, B = BW_onestep(data, w, A, B)
        # Debugging: Print row sums to ensure they sum to 1
```

```
if (iteration + 1) % 10 == 0 or iteration == 0:
            print(f"\nIteration {iteration + 1}")
            print("Row sums of A:", A.sum(axis=1))
            print("Row sums of B:", B.sum(axis=1))
            print(f"Log-Likelihood: {current_ll:.3f}")
   return A, B, current_ll
def myViterbi(data, w, A, B):
    Compute the most likely state sequence using the Viterbi algorithm
   Args:
        data: T-by-1 observation sequence (1D array)
        w: Initial state distribution (mz,)
        A: Transition matrix (mz-by-mz)
        B: Emission matrix (mz-by-mx)
   Returns:
       path: Most likely state sequence
       max_prob: Probability of the most likely path
    .....
    # Ensure data is 1D array
   data = np.asarray(data).flatten()
   T = len(data)
                        # Length of sequence
   mz = len(w)
                         # Number of hidden states
    # Initialize tables
   V = np.zeros((mz, T)) # Viterbi table
   bp = np.zeros((mz, T), dtype=int) # Backpointer table
    # Initialize first column of Viterbi table
   V[:, 0] = np.log(w) + np.log(B[:, data[0]])
    # Forward pass: Fill tables
   for t in range(1, T):
       for j in range(mz):
            # Calculate probabilities for all possible previous states
            probs = V[:, t-1] + np.log(A[:, j]) + np.log(B[j, data[t]])
            # Find maximum probability and its index
            V[j, t] = np.max(probs)
            bp[j, t] = np.argmax(probs)
    # Backward pass: Retrieve the most likely path
   path = np.zeros(T, dtype=int)
    # Find the most likely final state
```

```
path[-1] = np.argmax(V[:, -1])
max_prob = V[path[-1], -1]

# Backtrack through the sequence
for t in range(T-2, -1, -1):
    path[t] = bp[path[t+1], t+1]

return path, max_prob
```

```
[]: # Load HMM data
     data url = 'https://liangfgithub.github.io/Data/coding4 part2 data.txt'
     →Update if necessary
     data = pd.read_table(data_url, sep="\s+", header=None).values.flatten() - 1
     print("HMM Data Shape:", data.shape)
     print("First 5 Observations:\n", data[:5])
     # Load true state sequence (for comparison)
     Z_url = 'https://liangfgithub.github.io/Data/Coding4_part2_Z.txt' # Update ifu
      \rightarrownecessary
     Z true = pd.read_table(Z url, sep="\s+", header=None).values.flatten() - 1 #__
     →Assuming states are 1-indexed
     Z_true = Z_true[~np.isnan(Z_true)]
     print("True State Sequence Shape:", Z true.shape)
     print("First 5 True States:\n", Z_true[:5])
     # Initialize parameters
     mz = 2 # number of hidden states
     mx = 3 # number of observation symbols
     w_{initial} = np.array([0.5, 0.5])
     A_{initial} = np.array([[0.5, 0.5], [0.5, 0.5]])
     B_{initial} = np.array([[1/9, 3/9, 5/9], [1/6, 2/6, 3/6]])
     initial_params = {
         'w': w_initial,
         'A': A_initial,
         'B': B_initial
     }
     # Run Baum-Welch algorithm
     itmax = 100
     A_final, B_final, ll_final = myBW(data, initial_params, itmax)
     print("\nFinal Transition Matrix A:")
     print(A_final)
     print("\nFinal Emission Matrix B:")
     print(B_final)
```

```
print("\nFinal Log-Likelihood:")
print(ll_final)
HMM Data Shape: (200,)
First 5 Observations:
 [1 2 2 2 2]
True State Sequence Shape: (200,)
First 5 True States:
 [0. 0. 0. 0. 0.]
Iteration 1
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -221.373
Iteration 10
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -202.314
Iteration 20
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -202.313
Iteration 30
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -202.313
Iteration 40
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -202.312
Iteration 50
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -202.312
Iteration 60
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -202.311
Iteration 70
Row sums of A: [1. 1.]
```

Row sums of B: [1. 1.]

```
Log-Likelihood: -202.310
   Iteration 80
   Row sums of A: [1. 1.]
   Row sums of B: [1. 1.]
   Log-Likelihood: -202.309
   Iteration 90
   Row sums of A: [1. 1.]
   Row sums of B: [1. 1.]
   Log-Likelihood: -202.308
   Iteration 100
   Row sums of A: [1. 1.]
   Row sums of B: [1. 1.]
   Log-Likelihood: -202.306
   Final Transition Matrix A:
   [[0.49793938 0.50206062]
    [0.44883431 0.55116569]]
   Final Emission Matrix B:
   [[0.22159897 0.20266127 0.57573976]
    [0.34175148 0.17866665 0.47958186]]
   Final Log-Likelihood:
   -202.3062728417872
[]: # Run Viterbi algorithm
    path, max_prob = myViterbi(data, w_initial, A_final, B_final)
    print("\nMost Likely State Sequence:")
    print(path)
    print("\nProbability of the Most Likely Path:")
    print(max_prob)
    # Compare with true states
    matches = (path == Z_true).sum()
    total = len(Z_true)
    accuracy = matches / total
    print(f"\nViterbi Path Accuracy: {accuracy * 100:.2f}%")
   Most Likely State Sequence:
```

```
[]: # Initialize B uniformly
     B uniform = np.full((mz, mx), 1/3)
     initial_params_uniform = {
         'w': w initial,
         'A': A_initial,
         'B': B_uniform
     }
     # Run Baum-Welch for 20 iterations
     print("\nRunning Baum-Welch with Uniform B for 20 Iterations:")
     A final_20, B_final_20, ll_final_20 = myBW(data, initial_params_uniform,
      ⇒itmax=20)
     print("\nTransition Matrix A after 20 Iterations:")
     print(A final 20)
     print("\nEmission Matrix B after 20 Iterations:")
     print(B final 20)
     print("\nLog-Likelihood after 20 Iterations:")
     print(ll_final_20)
     # Run Baum-Welch for 100 iterations
     print("\nRunning Baum-Welch with Uniform B for 100 Iterations:")
     A final_100, B final_100, ll_final_100 = myBW(data, initial_params uniform, ___
      ⇒itmax=100)
     print("\nTransition Matrix A after 100 Iterations:")
     print(A final 100)
     print("\nEmission Matrix B after 100 Iterations:")
     print(B_final_100)
     print("\nLog-Likelihood after 100 Iterations:")
     print(ll_final_100)
```

Running Baum-Welch with Uniform B for 20 Iterations:

```
Iteration 1
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -219.722
```

```
Iteration 10
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -202.315
Iteration 20
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -202.315
Transition Matrix A after 20 Iterations:
[[0.5 0.5]
[0.5 \ 0.5]]
Emission Matrix B after 20 Iterations:
[[0.285 0.19 0.525]
 [0.285 0.19 0.525]]
Log-Likelihood after 20 Iterations:
-202.31544020689537
Running Baum-Welch with Uniform B for 100 Iterations:
Iteration 1
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -219.722
Iteration 10
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -202.315
Iteration 20
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -202.315
Iteration 30
Row sums of A: [1. 1.]
Row sums of B: [1. 1.]
Log-Likelihood: -202.315
Iteration 40
Row sums of A: [1. 1.]
```

Row sums of B: [1. 1.] Log-Likelihood: -202.315

Iteration 50 Row sums of A: [1. 1.] Row sums of B: [1. 1.] Log-Likelihood: -202.315 Iteration 60 Row sums of A: [1. 1.] Row sums of B: [1. 1.] Log-Likelihood: -202.315 Iteration 70 Row sums of A: [1. 1.] Row sums of B: [1. 1.] Log-Likelihood: -202.315 Iteration 80 Row sums of A: [1. 1.] Row sums of B: [1. 1.] Log-Likelihood: -202.315 Iteration 90 Row sums of A: [1. 1.] Row sums of B: [1. 1.] Log-Likelihood: -202.315 Iteration 100 Row sums of A: [1. 1.] Row sums of B: [1. 1.] Log-Likelihood: -202.315 Transition Matrix A after 100 Iterations:  $[[0.5 \ 0.5]]$ [0.5 0.5]] Emission Matrix B after 100 Iterations: [[0.285 0.19 0.525]

Log-Likelihood after 100 Iterations: -202.31544020689537

[0.285 0.19 0.525]]