Exam-2

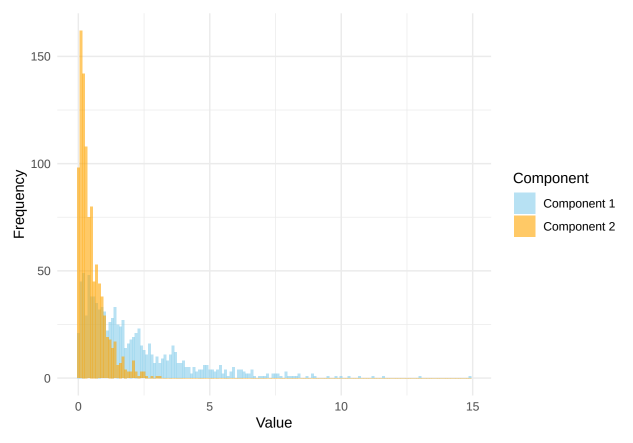
2024-11-16

Part I 1.a) What occurs in the E-step: Updates the probability that each data ponits belong to a specific cluster 1.b) What occurs in the M-Step: Updates the estimates of the parameters - both mixing coefficients and rate parameters. It leverages the probabilities calculated from E-step to get the parameter estimates by maximizing the likelihood.

1.c) Clusters can be overlapping due to the true rate parameters are close to each other. (soft clustering?)

Part II

```
library(ggplot2)
set.seed(1234)
# sample
n samples <- 2000
true_rates <- c(0.5, 2.0)
true_proportions \leftarrow c(0.5, 0.5)
# Sample group assignments (1 or 2) based on mixing proportions
component_labels <- sample(1:2, n_samples, replace = TRUE, prob = true_proportions)</pre>
# Stage 2: Generate data from the corresponding exponential
# distribution based on the group assignment
data <- numeric(n_samples) # Initialize the data vector</pre>
# Generate data for group 1 (rate = 0.5)
data[component_labels == 1] <- rexp(sum(component_labels == 1), rate = true_rates[1])</pre>
# Generate data for group 2 (rate = 2.0)
data[component_labels == 2] <- rexp(sum(component_labels == 2), rate = true_rates[2])</pre>
# Create a data frame for plotting
plot_data <- data.frame(</pre>
Value = data, # Data points
Component = factor(component_labels, labels = c("Component 1", "Component 2")) # Component labels
# Plot the data using ggplot2
ggplot(plot_data, aes(x = Value, fill = Component)) +
geom_histogram(binwidth = 0.1, position = "identity", alpha = 0.6) +
scale_fill_manual(values = c("skyblue", "orange")) + # Set colors for the components
labs(
x = "Value",
y = "Frequency",
fill = "Component"
) +
theme_minimal() +
theme(plot.title = element_blank()) # Remove the title
```



```
exponentialMixture <- function(data, K, max iter = 1000, tol = 1e-5) {
n <- length(data)</pre>
pi <- rep(1/K, K) # mixing proportions
lambda <- runif(K, 0.1, 1) # rate parameters</pre>
log_likelihoods <- numeric(max_iter) # store log-likelihood values</pre>
for (iter in 1:max_iter) { # E-step: numerator of the gammas
  gamma <- matrix(NA, nrow = n, ncol = K)
  for (k in 1:K) {
    gamma[, k] <- pi[k] * dexp(data, rate = lambda[k])</pre>
  row_sums <- rowSums(gamma) # denominator of the gammas</pre>
  gamma <- gamma / row_sums # normalize probabilities</pre>
  pi_old <- pi # M-step: Update mixing proportions and rate parameters</pre>
  lambda_old <- lambda</pre>
  pi <- colMeans(gamma) # update mixing proportions</pre>
  for (k in 1:K) {
    lambda[k] <- sum(gamma[, k]) / sum(gamma[, k] * data) # update rate parameters
  log_likelihoods[iter] <- sum(log(row_sums)) # calculate log-likelihood</pre>
  if (max(abs(pi - pi_old)) < tol && max(abs(lambda - lambda_old)) < tol) {
    log_likelihoods <- log_likelihoods[1:iter] # trim to the number of iterations</pre>
    cat("Convergence reached at iteration", iter,
    "with log-likelihood:", log_likelihoods[iter], "\n")
    break
    }
```

```
return(list(pi = pi, lambda = lambda, log_likelihood = log_likelihoods))
}

2.a) Cite: Exam Review Slide 26
set.seed(1234)
library(knitr)
# Fit the mixture model
result <- exponentialMixture(data, K = 2)

## Convergence reached at iteration 229 with log-likelihood: -2390.612
# Create a data frame to display the results in a table
results_table <- data.frame(
"Component" = 1:2,
"Estimated Mixing Proportion" = round(result$pi, 4),
"Estimated Rate Parameter (lambda)" = round(result$lambda, 4)
)
# move results to a table
kable(results_table, caption = "Estimated Parameters for the Mixture Model", format = "pipe")</pre>
```

Table 1: Estimated Parameters for the Mixture Model

Component	${\bf Estimated. Mixing. Proportion}$	Estimated. Rate. Parameter lambda.
1	0.5135	0.4998
2	0.4865	1.9721

The true rates for cluster 1 and cluster 2 are 0.5 and 2.0 respectively, while the mixing proportions are 0.5 and 0.5. The estimates closely resemble the true rates (0.4998 for cluster 1 and 1.9721 for cluster 2; both with error smaller than 0.02) and the mixing proportion (0.5135 for cluster 1 and 0.4865 for cluster; both with error less than 0.02).

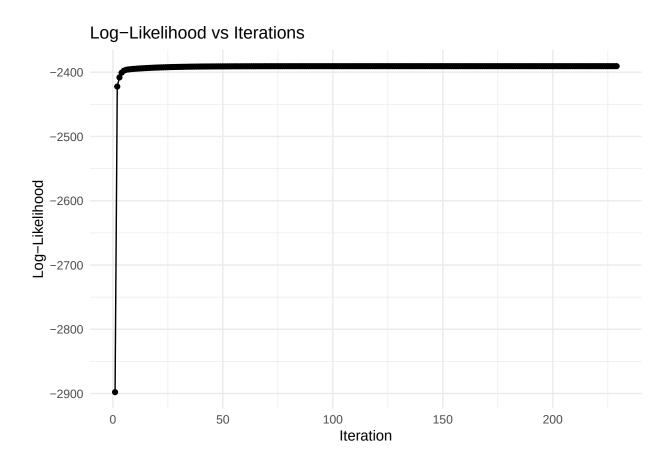
2.b) The parameter estimates do appear close to the true estimates. Cluster 1's true rate is 0.5 and its estimated rate is 0.4998, while cluster 2's true rate is 2.0 and its estimated rate is 1.9721. The true mixing proportions are 0.5 and the estimated proportion are 0.5135 and 0.4865. All the errors are less than 0.002.

2.c)

```
# Create a data frame for plotting
log_likelihoods <- result[["log_likelihood"]]

log_likelihood_data <- data.frame(
    Iteration = 1:length(log_likelihoods),
    LogLikelihood = log_likelihoods
)

# Generate the plot
ggplot(log_likelihood_data, aes(x = Iteration, y = LogLikelihood)) +
    geom_line() +
    geom_point() +
    labs(
        title = "Log-Likelihood vs Iterations",
        x = "Iteration",
        y = "Log-Likelihood"
) +
    theme_minimal()</pre>
```



The log-likelihood becomes stable after roughly 50 iterations, suggesting that the EM algorithm has converged to the optimal parameters (local maximum).

2.d) We typically monitor convergence of the algorithm using the log-likelihood.(Exam Review Slide 12) It is also evident by the code as we calculated the log likelihood (log_likelihoods[iter] <- sum(log(row_sums))) and didn't include the latent variable.

```
2.e)
```

```
n = 5
# an empty data frame to store results
results <- data.frame()</pre>
for (run in 1:n) {
  #set.seed(1234)
  # Fit the mixture model
  result <- exponentialMixture(data, K = 2)
  results <- rbind(</pre>
    results,
    data.frame(
      Run = run,
      Component = 1:2, # Components 1 and 2
      Mixing_Proportion = result$pi,
      Rate_Parameter = result$lambda
    )
  )
```

```
## Convergence reached at iteration 203 with log-likelihood: -2390.612
## Convergence reached at iteration 198 with log-likelihood: -2390.612
## Convergence reached at iteration 237 with log-likelihood: -2390.612
## Convergence reached at iteration 203 with log-likelihood: -2390.612
## Convergence reached at iteration 202 with log-likelihood: -2390.612
## Convergence reached at iteration 202 with log-likelihood: -2390.612
kable(results, caption = "Estimates from Five Runs of the EM Algorithm", format = "pipe")
```

Table 2: Estimates from Five Runs of the EM Algorithm

Run	Component	Mixing_Proportion	Rate_Parameter
1	1	0.5135368	0.4998434
1	2	0.4864632	1.9721059
2	1	0.4864629	1.9721072
2	2	0.5135371	0.4998436
3	1	0.5135381	0.4998440
3	2	0.4864619	1.9721106
4	1	0.4864611	1.9721136
4	2	0.5135389	0.4998445
5	1	0.4864616	1.9721118
5	2	0.5135384	0.4998442

The cluster labeling switches sometimes (Run 1 cluster 1 is the true cluster 1 with rate of 0.5, but Run 2 Cluster 2 is the true cluster 1). The estimated cluster rates and proportion vary slightly across runs. For examples, Run 2 Cluster 1 and Run 4 Cluster 1 both correspond to true cluster 1 but have estimated proportion of 0.4864629 and 0.4864611 (error less than 0.00001).

2.f) I found the most difficult when trying to code the latent variable. The exam review and the tutorials were particularly helpful for clarifying questions. Have tutorials to follow and check answers with could make learning on material easier in the future.

Part III.

```
# Simulation parameters
set.seed(123) # For reproducibility
n_samples <- 1000 # Number of samples
true_rates <- c(0.5, 1.5) # True rate parameters for two components
true_proportions <- c(0.6, 0.4) # True mixing proportions
# Generate synthetic data
data <- c(rexp(n_samples * true_proportions[1], rate = true_rates[1]),
rexp(n_samples * true_proportions[2], rate = true_rates[2]))
3.a)
n = 5
# an empty data frame to store results
results2 <- data.frame()

for (run in 1:n) {
    #set.seed(1234)
# Fit the mixture model</pre>
```

```
result2 <- exponentialMixture(data, K = 2)</pre>
  results2 <- rbind(
   results2,
    data.frame(
      Run = run,
      Component = 1:2, # Components 1 and 2
      Mixing_Proportion = result2$pi,
      Rate Parameter = result2$lambda
   )
  )
}
## Convergence reached at iteration 720 with log-likelihood: -1383.459
## Convergence reached at iteration 802 with log-likelihood: -1383.459
## Convergence reached at iteration 748 with log-likelihood: -1383.459
## Convergence reached at iteration 718 with log-likelihood: -1383.459
## Convergence reached at iteration 734 with log-likelihood: -1383.459
```

Table 3: Estimates from Five Runs of the EM Algorithm

kable(results2, caption = "Estimates from Five Runs of the EM Algorithm", format = "pipe")

Run	Component	Mixing_Proportion	Rate_Parameter
1	1	0.2326631	1.9503575
1	2	0.7673369	0.5624366
2	1	0.7673356	0.5624362
2	2	0.2326644	1.9503492
3	1	0.7673365	0.5624365
3	2	0.2326635	1.9503552
4	1	0.7673368	0.5624366
4	2	0.2326632	1.9503574
5	1	0.2326639	1.9503526
5	2	0.7673361	0.5624364

- 3.b) The cluster labeling switches sometimes (Run 1 cluster 1 is the true cluster 1 with rate of 0.5, but Run 2 Cluster 2 is the true cluster 1). The estimated parameters vary between runs (e.g. Run 3 Cluster 1 is 0.5624365 and Run 4 Cluster 1 is 0.5624366) and vary significantly from the true estimates. Est.proportions are around (0.7673,0.2327) contrasted with the true proportions (0.6,0.4), while the est.rates are around (0.5624,1.9503) contrasted with the true rates (0.5,1.5).
- 3.c) The root problem of switching label is because the model inherently considered the component labels being interchangeable. The root problem of significant variance between est.parameters and the true parameters can be a result that the EM algorithm converges to a local maximum, potentially due to the choice of initialization values.
- 3.d) K-means clustering can help with the root problem. We can group the data in clusters and estimate the mixing proportion using proportion of points in each cluster and rate using the inverse of the mean of data points in the cluster (since this is an exponential mixture model with a mean of 1/rate).

3.extra Credit)

```
#Cite: below from data-clean/06-more-clustering/code/kclust.R
exponentialMixture_kmeans <- function(data, K, max_iter = 1000, tol = 1e-5) {
    n <- length(data) # Initialize parameters using k-means
    kmeans_result <- kmeans(data, centers = K)</pre>
```

```
cluster_assignments <- kmeans_result$cluster</pre>
  # Mixing proportions
  pi <- table(cluster_assignments) / n</pre>
  # Rate parameters
  lambda <- sapply(1:K, function(k) {</pre>
    1 / mean(data[cluster assignments == k])
  })
  log_likelihoods <- numeric(max_iter) # store log-likelihood values</pre>
  for (iter in 1:max_iter) { # E-step: numerator of the gammas
    gamma <- matrix(NA, nrow = n, ncol = K)</pre>
    for (k in 1:K) {
      gamma[, k] <- pi[k] * dexp(data, rate = lambda[k])</pre>
    row_sums <- rowSums(gamma) # denominator of the gammas</pre>
    gamma <- gamma / row_sums # normalize probabilities</pre>
    pi\_old \leftarrow pi # M-step: Update mixing proportions and rate parameters
    lambda_old <- lambda</pre>
    pi <- colMeans(gamma) # update mixing proportions</pre>
    for (k in 1:K) {
      lambda[k] <- sum(gamma[, k]) / sum(gamma[, k] * data) # update rate parameters</pre>
    log_likelihoods[iter] <- sum(log(row_sums)) # calculate log-likelihood</pre>
    if (max(abs(pi - pi_old)) < tol && max(abs(lambda - lambda_old)) < tol) {</pre>
      log_likelihoods <- log_likelihoods[1:iter] # trim to the number of iterations</pre>
      cat("Convergence reached at iteration", iter,
      "with log-likelihood:", log_likelihoods[iter], "\n")
      break
    return(list(pi = pi, lambda = lambda, log_likelihood = log_likelihoods))
  }
n = 5
# an empty data frame to store results
results3 <- data.frame()
for (run in 1:n) {
  #set.seed(1234)
  # Fit the mixture model
  result3 <- exponentialMixture_kmeans(data, K = 2)
  results3 <- rbind(</pre>
    results3,
    data.frame(
      Run = run,
      Component = 1:2, # Components 1 and 2
      Mixing_Proportion = result3$pi,
      Rate_Parameter = result3$lambda
```

```
## Convergence reached at iteration 856 with log-likelihood: -1383.459
## Convergence reached at iteration 856 with log-likelihood: -1383.459
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## Convergence reached at iteration 856 with log-likelihood: -1383.459
kable(results3, caption = "Estimates from Five Runs of the EM Algorithm", format = "pipe")
```

Table 4: Estimates from Five Runs of the EM Algorithm

Run	Component	Mixing_Proportion	Rate_Parameter
1	1	0.2326635	1.9503554
1	2	0.7673365	0.5624365
2	1	0.2326635	1.9503554
2	2	0.7673365	0.5624365
3	1	0.2326635	1.9503554
3	2	0.7673365	0.5624365
4	1	0.2326635	1.9503554
4	2	0.7673365	0.5624365
5	1	0.2326635	1.9503554
5	2	0.7673365	0.5624365

K-means doesn't work as well in this case likely because k-means work best for cluster/spherical shape while exponential distribution's shape differs.

3.e) The first simulation taught me the foundation of mixture models and EM algorithm, including how to code it and how E-steps fit into M-steps and how to navigate the inherited randomness in the algorithm. The second simulation taught me the limitation of EM algorithm and potential solution to address the limitation of trapping in the local maximum. Moving forward, I will make sure to carefully choose reasonable initial values for the algorithm even when the truth is unknown and apply appropriate technique such as clustering to increase the accuracy and precision of EM algorithm.