lab01

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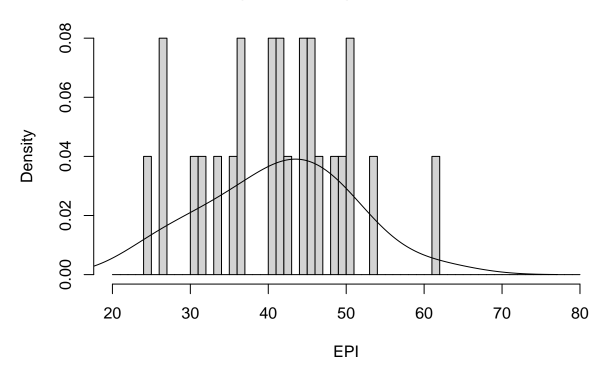
Load the results dataset

})

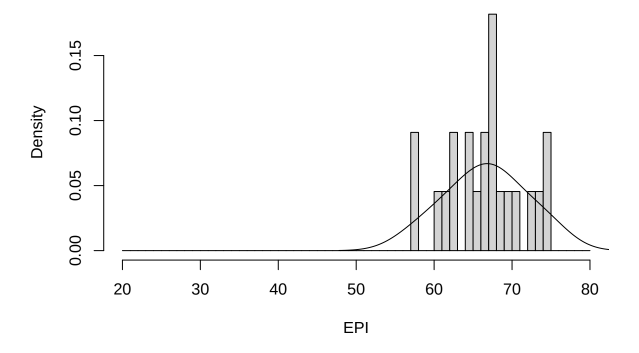
```
# read data
epi <- read.csv("../data/epi2024results_DA_F24_lab03.csv")</pre>
selected_regions <- epi %>%
 distinct(region) %>%
  slice_tail(n = 2) %>%
  pull(region)
epi %>%
  filter(region %in% selected_regions) %>%
  group_by(region) %>%
  group_walk(~ {
    region_name <- .y$region</pre>
        EPI_data <- .x$EPI</pre>
    # Create a histogram
    hist(EPI_data, breaks = seq(20, 80, 1.0), prob = TRUE,
         main = paste("Histogram for Region:", region_name),
         xlab = "EPI", ylab = "Density")
    # Add the density line
```

lines(density(EPI_data, na.rm = TRUE, bw = 'SJ'))

Histogram for Region: Asia-Pacific

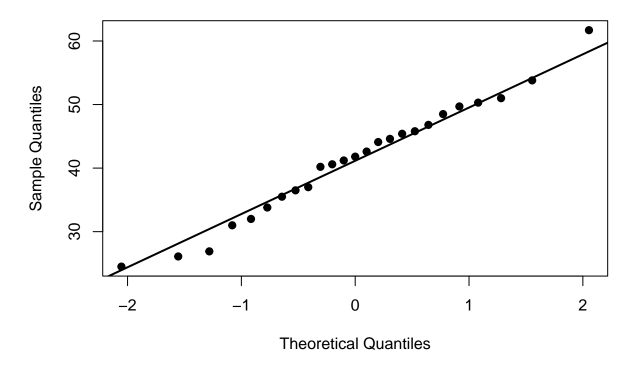


Histogram for Region: Global West

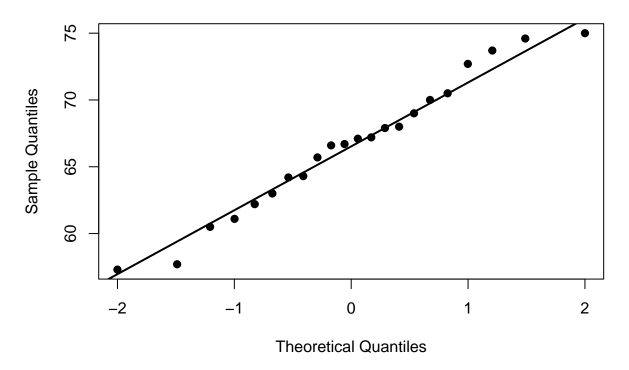


```
epi %>%
  filter(region %in% selected_regions) %>%
  group_by(region) %>%
 group_walk(~ {
    region_name <- .y$region</pre>
    EPI_data <- .x$EPI</pre>
    # Calculate the mean and standard deviation of the data
    mean_EPI <- mean(EPI_data, na.rm = TRUE)</pre>
    sd_EPI <- sd(EPI_data, na.rm = TRUE)</pre>
    # Create the Q-Q plot against a normal distribution
    qqnorm(EPI_data,
           main = paste("Q-Q Plot for Region:", region_name),
           xlab = "Theoretical Quantiles", ylab = "Sample Quantiles",
           pch = 19)
    \# Add a Q-Q line for a normal distribution
    qqline(EPI_data, distribution = qnorm, lwd = 2)
 })
```

Q-Q Plot for Region: Asia-Pacific



Q-Q Plot for Region: Global West

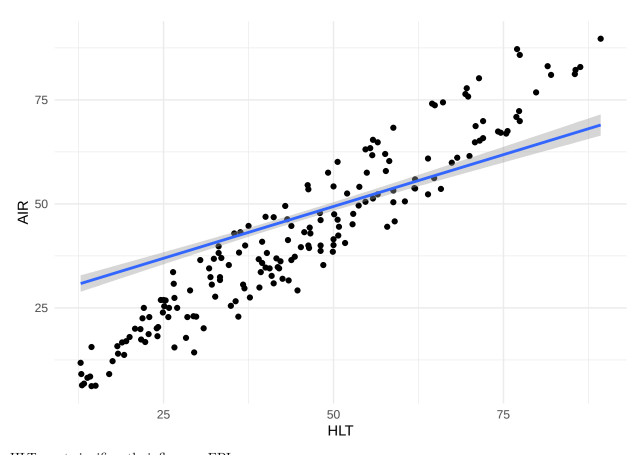


```
linear_model <- lm(EPI ~ WWT + WWR + HLT + AIR + HPE, data = epi)
summary(linear_model)</pre>
```

```
##
## Call:
## lm(formula = EPI ~ WWT + WWR + HLT + AIR + HPE, data = epi)
##
## Residuals:
##
        Min
                  1Q
                                     ЗQ
                                             Max
                       Median
## -15.5124 -4.4814
                       0.6935
                                4.2839
                                        18.2293
##
##
  Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
                           1.41326
                                    17.438
## (Intercept) 24.64374
                                            < 2e-16 ***
                0.04772
                           0.02855
                                     1.672
                                             0.0964 .
## WWT
## WWR
                0.02863
                           0.03665
                                     0.781
                                              0.4358
                                     5.304 3.42e-07 ***
## HLT
                0.75468
                           0.14230
## AIR
               -0.36305
                           0.16481
                                    -2.203
                                              0.0289 *
## HPE
                0.01975
                           0.04835
                                     0.409
                                              0.6834
##
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## Residual standard error: 6.048 on 174 degrees of freedom
## Multiple R-squared: 0.7335, Adjusted R-squared: 0.7259
## F-statistic: 95.79 on 5 and 174 DF, p-value: < 2.2e-16
```

```
ggplot(epi, aes(x = HLT, y = AIR)) +
geom_point() +
stat_smooth(method = "lm", aes(y = EPI)) +
theme_minimal()
```

'geom_smooth()' using formula = 'y ~ x'



 HLT most significantly influences EPI.

```
sub_saharan_africa_epi <- epi %>%
  filter(region == "Sub-Saharan Africa")
linear_model <- lm(EPI ~ WWT + WWR + HLT + AIR + HPE, data = sub_saharan_africa_epi)
summary(linear_model)</pre>
```

```
##
## Call:
## Im(formula = EPI ~ WWT + WWR + HLT + AIR + HPE, data = sub_saharan_africa_epi)
##
## Residuals:
## Min    1Q Median   3Q Max
## -8.4780 -2.7365 -0.2571  2.1588  10.6573
##
## Coefficients:
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) 38.16511  2.56907  14.856  <2e-16 ***</pre>
```

```
## WWT
               0.07774
                          0.04585
                                    1.695
                                            0.0978 .
               0.20308
                          0.09536
                                    2.130
                                            0.0394 *
## WWR
## HLT
              -0.14346
                          0.46958
                                   -0.306
                                            0.7616
## AIR
               0.07154
                          0.49606
                                    0.144
                                            0.8861
## HPE
               0.01275
                          0.09763
                                    0.131
                                            0.8968
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 4.273 on 40 degrees of freedom
## Multiple R-squared: 0.4241, Adjusted R-squared: 0.3521
## F-statistic: 5.891 on 5 and 40 DF, p-value: 0.0003606
```

The full model is a better fit. It captures more variance, as shown by and Adjusted R-Squared of 0.7259 versus 0.3521. It has higher residuals, but that can be explained by the variance of EPI being higher. It is also worth noting that the significance of predictors varies, implying there are different relationships between the predictors and EPI in different regions.

```
epi.norm <- epi %>%
  select(EPI, WWT, WWR, HLT, AIR)
epi.norm <- as.data.frame(scale(epi.norm))</pre>
epi.norm$region <- epi$region
# Define the regions
regions_a <- c("Latin America & Caribbean", "Asia-Pacific", "Eastern Europe")
regions_b <- c("Sub-Saharan Africa", "Global West", "Greater Middle East")
epi.norm.a <- epi.norm %>%
  filter(region %in% regions_a)
epi.norm.b <- epi.norm %>%
  filter(region %in% regions_b)
# Create a random sample of 80% of the data
set.seed(123) # Set seed for reproducibility
sample_index.a <- sample(seq_len(nrow(epi.norm.a)), size = 0.8 * nrow(epi.norm.a))</pre>
sample_index.b <- sample(seq_len(nrow(epi.norm.b)), size = 0.8 * nrow(epi.norm.b))</pre>
train_a <- epi.norm.a[sample_index.a, ]</pre>
test_a <- epi.norm.a[-sample_index.a, ]</pre>
train_b <- epi.norm.b[sample_index.b, ]</pre>
test_b <- epi.norm.b[-sample_index.b, ]</pre>
KNNpred.a \leftarrow knn(train = train_a[1:5], test = test_a[1:5], cl = train_a$region)
KNNpred.b \leftarrow knn(train = train_b[1:5], test = test_b[1:5], cl = train_b$region)
contingency_a <- table(Predicted = KNNpred.a, Actual = test_a$region)</pre>
contingency_b <- table(Predicted = KNNpred.b, Actual = test_b$region)</pre>
contingency_a
```

```
##
                               Actual
## Predicted
                                Asia-Pacific Eastern Europe
##
     Asia-Pacific
##
     Eastern Europe
                                            1
                                                            1
     Latin America & Caribbean
                                            2
                                                            2
##
##
                               Actual
```

```
## Predicted
                                Latin America & Caribbean
##
     Asia-Pacific
                                                          1
     Eastern Europe
##
                                                          0
    Latin America & Caribbean
                                                          6
##
contingency b
##
                         Actual
## Predicted
                          Global West Greater Middle East Sub-Saharan Africa
##
    Global West
                                    2
##
     Greater Middle East
                                    0
                                                         5
                                                                              0
     Sub-Saharan Africa
                                    0
##
                                                          1
                                                                              9
accuracy_a <- sum(diag(contingency_a)) / sum(contingency_a)</pre>
accuracy_b <- sum(diag(contingency_b)) / sum(contingency_b)</pre>
print(paste("Accuracy a:", round(accuracy_a * 100, 2)))
## [1] "Accuracy a: 56.25"
print(paste("Accuracy b:", round(accuracy_b * 100, 2)))
## [1] "Accuracy b: 94.12"
```

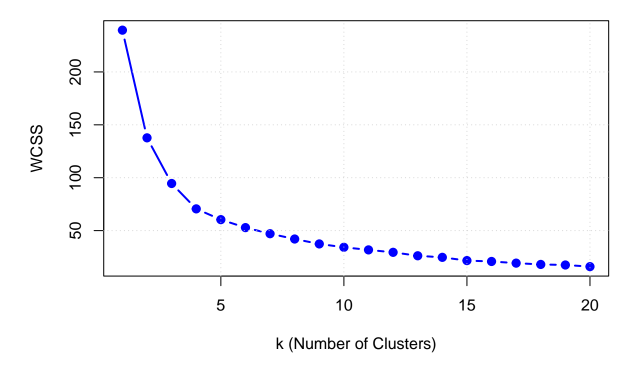
The accuracy for region b is significantly greater. This is likely due to greater differences in the chosen columns between the regions chosen in the group.

```
k fold kmeans <- function(data, k values = seq(1, 20), k folds = 10, seed = 123) {
  # Set the seed for reproducibility
  set.seed(seed)
  \# Create a k-fold cross-validation partition
  folds <- createFolds(data$region, k = k_folds, list = FALSE)</pre>
  # Initialize a vector to store WCSS for different k values
  wcss_values <- numeric(length(k_values))</pre>
  # Loop through different values of k
  for (i in seq_along(k_values)) {
    k <- k_values[i] # Get the current value of k</pre>
    fold_wcss <- numeric(k_folds) # Initialize vector to store WCSS for each fold
    # Loop through each fold
    for (fold in seq_len(k_folds)) {
      # Split the data into training and testing sets based on the fold
      train_data <- data[folds != fold, ]</pre>
      # Perform K-means clustering on the training data
      kmeans_model <- kmeans(train_data[, -ncol(train_data)], centers = k)</pre>
      # Calculate WCSS for this fold and store it
      fold_wcss[fold] <- sum(kmeans_model$withinss)</pre>
```

```
# Average WCSS across all folds for this k value
    wcss_values[i] <- mean(fold_wcss)</pre>
  }
  # Plot the WCSS vs. k values
  plot(k_values, wcss_values, type = "b", pch = 19, col = "blue", lwd = 2,
       xlab = "k (Number of Clusters)", ylab = "WCSS",
       main = "K-Means WCSS vs. k (K-Fold CV)")
  grid() # Add grid lines
epi.norm <- epi %>%
  select(EPI, WWT, WWR, HLT, AIR)
epi.norm <- as.data.frame(scale(epi.norm))</pre>
epi.norm$region <- epi$region</pre>
# Define the regions
regions_a <- c("Latin America & Caribbean", "Asia-Pacific", "Eastern Europe")
regions_b <- c("Sub-Saharan Africa", "Global West", "Greater Middle East")</pre>
epi.norm.a <- epi.norm %>%
  filter(region %in% regions_a)
epi.norm.b <- epi.norm %>%
  filter(region %in% regions_b)
```

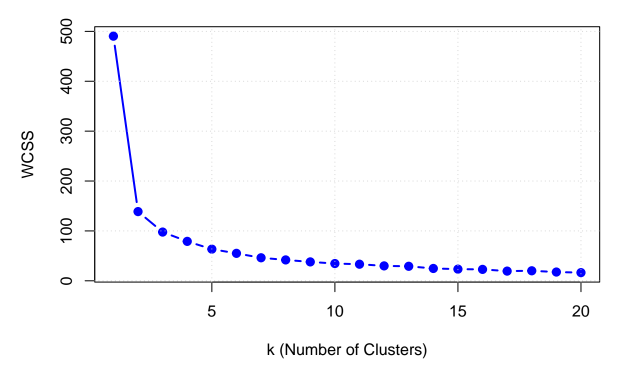
k_fold_kmeans(epi.norm.a)

K-Means WCSS vs. k (K-Fold CV)



k_fold_kmeans(epi.norm.b)

K-Means WCSS vs. k (K-Fold CV)



With group a we see tighter clusters, with a wcss of $\sim\!65$ at the elbow vs. $\sim\!100$ for b. This either indicates less variance in group a or better separation of clusters.