Project 3

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Network Structure Analysis

Since the graph is NOT strongly connected, we have to find the diameter and radius using the largest strongly connected sub component.

The dataset also does not have duplicate sender -> receiver pairs, so we will analyze the dataset as an unweighted graph.

```
# Read the text file into a dataframe
df <- read.table("email-Eu-core.txt", header = FALSE, sep = " ")</pre>
colnames(df) <- c("From", "To")</pre>
# The dataset does not hav
# df_weighted <- aggregate(weight ~ From + To, data = transform(df, weight = 1), FUN = sum)
g <- graph_from_data_frame(df, directed = TRUE)</pre>
# q <- graph from data frame(df weighted, directed = TRUE)
num_nodes = vcount(g)
num_edges = ecount(g)
# Check strong connectivity
components <- components(g, mode="strong")</pre>
# Extract the vertices belonging to the largest strongly connected component
largest_comp_vertices <- V(g)[components$membership == which.max(components$csize)]</pre>
g_largest <- induced_subgraph(g, largest_comp_vertices)</pre>
# Calculate diameter and radius on the largest SCC
diameter largest <- diameter(g largest, directed = TRUE)</pre>
radius_largest <- radius(g_largest, mode = "out")</pre>
output <- paste(</pre>
  "Graph Summary",
  paste("Number of nodes:", num_nodes),
  paste("Number of edges:", num_edges),
  paste("Density:", round(edge_density(g), 6)),
  paste(""),
  "Connectivity:",
```

```
paste("Number of strongly connected components:", components$no),
  paste("Size of largest strongly connected component:", max(components$csize)),
  paste("fraction of elements belonging to the largest strong subcomponent: ", max(components$csize)/nu
  "The graph is NOT strongly connected",
  paste(""),
  paste("Largest diameter:", diameter_largest),
  paste("Largest radius:", radius_largest),
  sep = "\n"
)
cat(output)
## Graph Summary
## Number of nodes: 1005
## Number of edges: 25571
## Density: 0.025342
## Connectivity:
## Number of strongly connected components: 203
## Size of largest strongly connected component: 803
## fraction of elements belonging to the largest strong subcomponent: 0.799004975124378
## The graph is NOT strongly connected
## Largest diameter: 6
## Largest radius: 3
Cluster coefficient computed through the ClustF function found in the DirectedClustering package:
adj_matrix <- as_adjacency_matrix(g, sparse = FALSE)</pre>
# Compute clustering coefficients (cycle type)
clustering_coeff <- ClustF(adj_matrix, type = "cycle")</pre>
## Warning in ClustF(adj_matrix, type = "cycle"): Loops have been removed
# Print the global cycle clustering coefficient
print(clustering_coeff$GlobalcycleCC)
## [1] 0.2934512
print(paste("Global out-clustering coefficient:", clustering_coeff$GlobaloutCC))
## [1] "Global out-clustering coefficient: 0.288824631612028"
```

Degree Distribution Analysis

```
# Calculate in-degree, out-degree, and total degree
in_degree <- degree(g, mode = "in")
out_degree <- degree(g, mode = "out")
total_degree <- degree(g, mode = "all")

# Set up a 1x3 plotting layout
par(mfrow = c(1, 3))</pre>
```

```
# Plot histogram for in-degree
hist(in_degree, breaks = 50, main = "In-Degree Distribution",
     xlab = "In-Degree", ylab = "Frequency", col = "skyblue", border = "black")
# Plot histogram for out-degree
hist(out_degree, breaks = 50, main = "Out-Degree Distribution",
     xlab = "Out-Degree", ylab = "Frequency", col = "salmon", border = "black")
# Plot histogram for total degree
hist(total_degree, breaks = 50, main = "Total Degree Distribution",
     xlab = "Total Degree", ylab = "Frequency", col = "lightgreen", border = "black")
          In-Degree Distribution
                                             Out-Degree Distribution
                                                                               Total Degree Distribution
                                     350
   250
                                     300
                                                                        250
   200
                                     250
                                                                        200
                                     200
  150
                                                                        150
                                     150
   100
                                                                        100
                                     9
   20
                                                                        20
                                     20
                100
                      150
                            200
                                               100 150 200 250 300
                                                                               100
                                                                                                500
     Ω
           50
                                           50
                                                                           0
                                                                                   200
                                                                                       300
                                                                                            400
               In-Degree
                                                  Out-Degree
                                                                                    Total Degree
# Reset plotting layout
par(mfrow = c(1, 1))
# Summary statistics
cat("In-Degree Summary:\n")
## In-Degree Summary:
print(summary(in_degree))
##
      Min. 1st Qu. Median
                                 Mean 3rd Qu.
                                                   Max.
                      18.00
##
      0.00
               5.00
                                25.44
                                         36.00 212.00
cat("\nOut-Degree Summary:\n")
##
## Out-Degree Summary:
print(summary(out_degree))
      Min. 1st Qu. Median
                                 Mean 3rd Qu.
                                                   Max.
##
##
      0.00
               2.00
                       15.00
                                25.44
                                         35.00 334.00
cat("\nTotal Degree Summary:\n")
```

Total Degree Summary:

```
print(summary(total_degree))
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 1.00 8.00 32.00 50.89 71.00 546.00
```

Community Detection

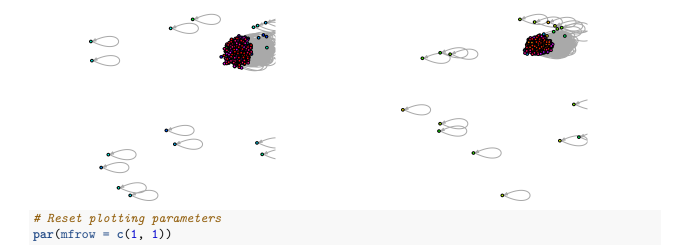
I chose to utilize the Edge Betweenness algorithm and the Walktrap algorithm for community detection. I used these two algorithms because they both work for unweighted, directed graphs and because of how differently they determine communities.

```
eb_communities <- cluster_edge_betweenness(g)</pre>
eb_membership <- membership(eb_communities)</pre>
eb_modularity <- modularity(eb_communities)</pre>
eb_sizes <- sizes(eb_communities)</pre>
# Walktrap Community Detection
wt_communities <- cluster_walktrap(g)</pre>
wt_membership <- membership(wt_communities)</pre>
wt_modularity <- modularity(wt_communities)</pre>
wt_sizes <- sizes(wt_communities)</pre>
# Print summary statistics for both algorithms
cat("Edge Betweenness Community Detection:\n")
## Edge Betweenness Community Detection:
cat("Number of communities:", length(unique(eb_membership)), "\n")
## Number of communities: 710
cat("Modularity:", round(eb_modularity, 4), "\n")
## Modularity: 0.0425
cat("Community sizes:\n")
## Community sizes:
print(sort(table(eb_membership), decreasing = TRUE)[1:10])
## eb_membership
     1
         3 146 60 91 97 103 29 56 96
## 233 17
                 5
                      4
                                      3
cat("\n")
cat("Walktrap Community Detection:\n")
## Walktrap Community Detection:
cat("Number of communities:", length(unique(wt membership)), "\n")
## Number of communities: 136
cat("Modularity:", round(wt modularity, 4), "\n")
## Modularity: 0.3673
```

```
cat("Community sizes:\n")
## Community sizes:
print(sort(table(wt_membership), decreasing = TRUE)[1:10])
## wt_membership
## 4 1 5 9 12 7 8 11 6
                                         3
## 352 109 91 85 85 84 30 22 12
# Visualize the communities
# Function to plot communities with layout optimization
plot_communities <- function(graph, membership, title) {</pre>
  # Create color palette for communities
  num_communities <- length(unique(membership))</pre>
  colors <- rainbow(num_communities)</pre>
  # Set vertex colors based on community membership
  V(graph)$color <- colors[membership]</pre>
  # Calculate layout (using Fruchterman-Reingold algorithm)
  layout <- layout_with_fr(graph)</pre>
  # Plot the graph
  plot(graph,
       layout = layout,
       vertex.size = 3,
      vertex.label = NA,
       edge.arrow.size = 0.2,
       main = title)
}
# Set up plotting area for side-by-side comparison
par(mfrow = c(1, 2))
# Plot both community structures
plot_communities(g, eb_membership, "Edge Betweenness Communities")
plot_communities(g, wt_membership, "Walktrap Communities")
```

Edge Betweenness Communitie

Walktrap Communities



Triad Census Analysis

For the walktrap algorithm, these results indicate decent community detection. For the triads, over half are in the same community together.

However, for the edge betweenness algorithm, the low percentage of about 14% indicates that the algorithm may be breaking up communities into too small of groups. The large difference between both algorithms suggests that walktrap may be better suited than edge betweenness for detection.

Also unrelated note, but the 'triad_census' function here is fantastic. I tried to manually compute all the different triads using my own naive R code and my computer was not able to do that even after 8 hours of computation. I looked into random sampling every N triads per subcomponent of the graph because of these compute restraints, but the following approach was already very efficient and effective, so I abandoned that the N sampling strategy.

```
analyze_triads_communities <- function(g, communities) {
    # Get triad census
    triad_counts <- triad_census(g)

# Get community memberships
    comm_mem <- membership(communities)

# Function to check if three nodes form a triangle
    is_triangle <- function(nodes) {
        subg <- induced_subgraph(g, nodes)
        return(ecount(subg) >= 2) # At least 2 edges for directed graph
    }

# Get triangles using cliques
# For directed graphs, we look at the underlying undirected graph for triangles
    g_undir <- as.undirected(g, mode="collapse")
    triangles <- cliques(g_undir, min=3, max=3)

# Count triangles in same community</pre>
```

```
same_comm_triangles <- 0</pre>
 total_triangles <- length(triangles)</pre>
 for(triangle in triangles) {
   if(length(unique(comm_mem[triangle])) == 1) {
      same_comm_triangles <- same_comm_triangles + 1</pre>
 }
 # Create results
 results <- list(
   triad_census = triad_counts,
   triangle_stats = list(
     total_triangles = total_triangles,
     same_comm_triangles = same_comm_triangles,
     percent_same_comm = if(total_triangles > 0)
                          (same_comm_triangles/total_triangles) * 100
   )
 )
 return(results)
}
# Analyze both community detection methods
wt_triad_analysis <- analyze_triads_communities(g, wt_communities)</pre>
## Warning: `as.undirected()` was deprecated in igraph 2.1.0.
## i Please use `as_undirected()` instead.
## This warning is displayed once every 8 hours.
## Call `lifecycle::last_lifecycle_warnings()` to see where this warning was
## generated.
eb_triad_analysis <- analyze_triads_communities(g, eb_communities)</pre>
# Print results
cat("Triad Census Analysis Results:\n")
## Triad Census Analysis Results:
cat("----\n")
## -----
cat("Total triads by type:\n")
## Total triads by type:
print(wt_triad_analysis$triad_census)
   [1] 153640073
                   6345756
                             7716387
                                         81896
                                                   38347
                                                             58745
                                                                      145903
## [8]
                      5639
                                 419
          262008
                                        279934
                                                    6984
                                                             11123
                                                                        7455
## [15]
           39656
                     34185
cat("\nTriangle Statistics:\n")
```

##

```
## Triangle Statistics:
cat("Walktrap Communities:\n")
## Walktrap Communities:
cat(" Total triangles:", wt_triad_analysis$triangle_stats$total_triangles, "\n")
##
     Total triangles: 105461
cat(" Triangles in same community:", wt_triad_analysis$triangle_stats$same_comm_triangles, "\n")
##
     Triangles in same community: 60034
cat(" Percentage in same community:",
    round(wt_triad_analysis$triangle_stats$percent_same_comm, 2), "%\n")
     Percentage in same community: 56.93 %
cat("\nEdge Betweenness Communities:\n")
##
## Edge Betweenness Communities:
cat(" Total triangles:", eb_triad_analysis$triangle_stats$total_triangles, "\n")
     Total triangles: 105461
cat(" Triangles in same community:", eb_triad_analysis$triangle_stats$same_comm_triangles, "\n")
     Triangles in same community: 14697
##
cat(" Percentage in same community:",
    round(eb_triad_analysis$triangle_stats$percent_same_comm, 2), "%\n")
##
     Percentage in same community: 13.94 %
Modularity Analysis
wt_mod_value <- modularity(g, wt_membership)</pre>
eb_mod_value <- modularity(g, eb_membership)</pre>
cat("Walktrap Modularity: ", wt_mod_value, "\n")
## Walktrap Modularity: 0.3821017
```

Evaluation

Based off this analysis, neither algorithm did very well at determining the correct communities. The edge betweenness algorithm did better than the walktrap algorithm in terms of NMI (0.444 vs 0.181), however both algorithms scored close to zero in the ARI score. In fact, the edge betweenness ARI score of (-0.002) indicates that the algorithm may actually be worse than random.

cat("Edge Betweenness Modularity: ", eb_mod_value, "\n")

Edge Betweenness Modularity: 0.04254974

```
gt_communities <- read.table("email-Eu-core-department-labels.txt", header=FALSE)
colnames(gt_communities) <- c("Node", "Community")</pre>
```

```
# Convert memberships to vectors ensuring they use the same node ordering
gt_membership <- gt_communities$Community</pre>
wt_membership_ordered <- membership(wt_communities)[order(as.numeric(V(g)))]</pre>
eb_membership_ordered <- membership(eb_communities)[order(as.numeric(V(g)))]
# Calculate NMI
nmi_wt <- NMI(gt_membership, wt_membership_ordered)</pre>
nmi_eb <- NMI(gt_membership, eb_membership_ordered)</pre>
# Calculate ARI
ari_wt <- ARI(gt_membership, wt_membership_ordered)</pre>
ari_eb <- ARI(gt_membership, eb_membership_ordered)</pre>
# Print results
cat("Comparison with Ground Truth:\n")
## Comparison with Ground Truth:
cat("Walktrap Algorithm:\n")
## Walktrap Algorithm:
cat(" NMI:", nmi_wt, "\n")
    NMI: 0.1806671
##
cat(" ARI:", ari_wt, "\n")
     ARI: 0.001814829
##
cat("\nEdge Betweenness Algorithm:\n")
##
## Edge Betweenness Algorithm:
cat(" NMI:", nmi_eb, "\n")
    NMI: 0.4440126
cat(" ARI:", ari_eb, "\n")
     ARI: -0.001998724
# Create visualization comparing community sizes
par(mfrow=c(1,3))
# Ground Truth distribution
barplot(table(gt_membership),
        main="Ground Truth Community Sizes",
        xlab="Community ID",
        ylab="Size")
# Walktrap distribution
barplot(table(wt_membership),
        main="Walktrap Community Sizes",
        xlab="Community ID",
        ylab="Size")
# Edge Betweenness distribution
```

```
barplot(table(eb_membership),
       main="Edge Betweenness Community Sizes",
        xlab="Community ID",
       ylab="Size")
```

Ground Truth Community Size

Walktrap Community Sizes Edge Betweenness Community S

