Statistical Analysis of Nodal Network Structures

2025-03-22

Dataset Information

Let's read in two datasets containing graph theory measures of patients who were sleeping while undergoing EEG. The target variable is Dreams, a binary (0,1) value that indicates if a patient reported a conscious experience. The data is messy, let's read it in, inspect it, and do some pre-processing. We will need a non-parametric test to compare the two groups, this PDF outlines the original supplement, AND the assumptions that led to the test statistics on the box/violin plot.

```
suppressPackageStartupMessages({
  library(readr)
  library(dplyr)
  library(tidyr)
})
## Warning: package 'tidyr' was built under R version 4.4.3
df1 <- read_csv("C:/Users/User/Desktop/group1.csv", show_col_types = FALSE)</pre>
df2 <- read_csv("C:/Users/User/Desktop/group2.csv", show_col_types = FALSE)</pre>
#head(df1, 3);
head(df2, 3)
## # A tibble: 3 x 10
     Subject Band threshold wGlobEff wTrans mean wBetw bTrans mean bBetw Dreams
##
     <chr>>
                                                                                <dbl>
##
              <chr>
                         <dbl>
                                   <dbl>
                                          <dbl>
                                                      <dbl>
                                                             <dbl>
                                                                         <dbl>
## 1 sub_2012 delta
                         0.202
                                   0.125 0.105
                                                       58.3 0.439
                                                                          151.
                                                                                     0
## 2 sub_2010 beta
                                   0.164 0.125
                                                                                     0
                         0.181
                                                      132.
                                                             0.309
                                                                          201.
## 3 sub_2010 alpha
                         0.294
                                 NA
                                          0.151
                                                       47.8 0.302
                                                                          233.
                                                                                     0
## # i 1 more variable: degree <dbl>
process_combined_data <- function(df1, df2) {</pre>
  # Combine both data frames in a row-wise manner
  combined <- rbind(df1, df2)</pre>
  # Number of rows before dropping NA values
  n before <- nrow(combined)</pre>
  # Remove rows with any NA values
  combined_clean <- tidyr::drop_na(combined)</pre>
  # Number of rows after dropping NA values
  n_after <- nrow(combined_clean)</pre>
  # print the total number of rows dropped
  cat("Dropped", n_before - n_after, "rows containing NA values.\n")
  # Convert the Dreams column: 0 -> "No Dreams", 1 -> "Dreaming"
  combined_clean$Dreams <- factor(</pre>
```

```
combined_clean$Dreams,
levels = c(0, 1),
labels = c("No Dreams", "Dreaming")
)

# Create a new column 'obs' to track observation number
combined_clean$obs <- seq_len(nrow(combined_clean))

# Return the cleaned, merged dataset
return(combined_clean)
}
processed_data <- process_combined_data(df1, df2)</pre>
```

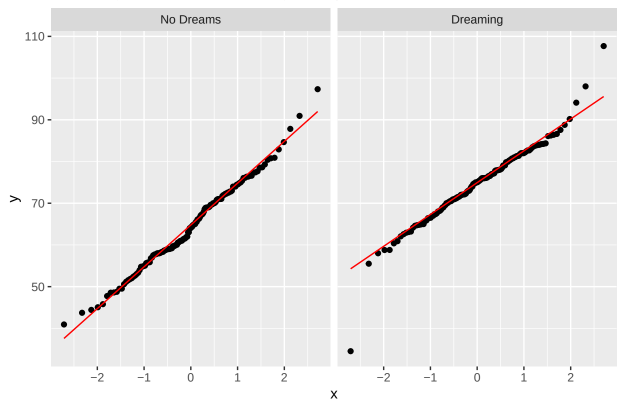
Dropped 103 rows containing NA values.

Statistical Assumption Checking

Normality Tests

```
# Shapiro-Wilk test for normality by group
shapiro_results <- processed_data %>%
 group_by(Dreams) %>%
 summarise(
   statistic = shapiro.test(degree)$statistic,
   p.value = shapiro.test(degree)$p.value
print(shapiro_results)
## # A tibble: 2 x 3
   Dreams statistic p.value
##
   <fct> <dbl>
                           <dbl>
## 1 No Dreams 0.990 0.375
                0.959 0.000253
## 2 Dreaming
# QQ Plots for further normality tests
suppressPackageStartupMessages(library(ggplot2))
ggplot(processed_data, aes(sample = degree)) +
 geom_qq() +
 geom_qq_line(color = "red") +
 facet_wrap(~Dreams) +
 ggtitle("Normality Check: Q-Q Plots by Group")
```

Normality Check: Q-Q Plots by Group



Shapiro-Wilk test showed significant evidence against the null hypothesis of normality, the powerful non-normality is evidenced by p-values ($\mathbf{p}=0.00025$) in the dreaming group, the non-dreaming group appears normal with ($\mathbf{p}=0.3752400104$). This test is further supported by the Dreaming group's qq-plot showing heavier tails or skewness compared to a normal distribution via the upward curve in the high x-range.

Homogeneity of Variance test

```
# Levene's Test to see how to set the eq.val argument in ggbetweenstats()
suppressPackageStartupMessages(library(car))
leveneTest(degree ~ Dreams, data = processed_data, center = median)

## Levene's Test for Homogeneity of Variance (center = median)

## Df F value Pr(>F)
## group 1 6.9179 0.008982 **

## 295

## ---

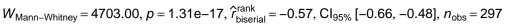
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

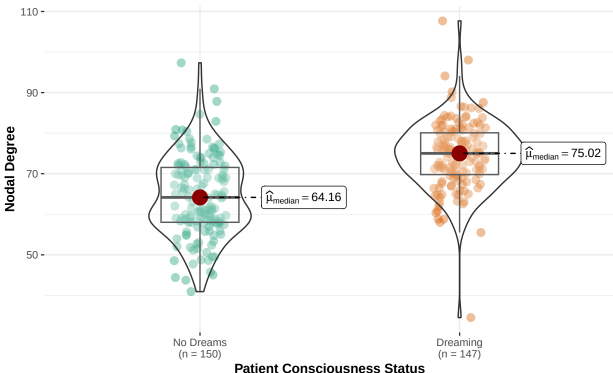
Levene's test rejects homogeneity of variance, and assumes heterogeneity of variances (p = 0.009)

Based on the prior statistical tests, we will use Mann-Whitney U test. The Mann-Whitney U test does not assume normality or equal variances and is robust to outliers and skewed distributions. This matches the data'a characteristics

```
suppressPackageStartupMessages(library(ggstatsplot))
ggbetweenstats(
  data = processed_data,
  x = Dreams,  # 'Dreams' is a factor that indicates group membership
  y = degree,  # 'degree' is the graph measure to compare the two groups
  type = "nonparametric", # Nonparametric test is selected
  bf.message = FALSE,  # set to TRUE for postier probabilities
  xlab = "Patient Consciousness Status",
  ylab = "Nodal Degree",
  title = "Comparison of Degree by Dreams Group",
  messages = FALSE
)
```

Comparison of Degree by Dreams Group





Patients who reported dreaming had significantly lower nodal degree (a network connectivity measure) compared to those who did not, with a clinically meaningful effect, as evidenced by a large effect size ($\mathbf{r} = -0.57$), non-overlapping confidence intervals (CI: [-0.66, -0.48]), and extreme statistical significance ($\mathbf{p} < 0.05$)

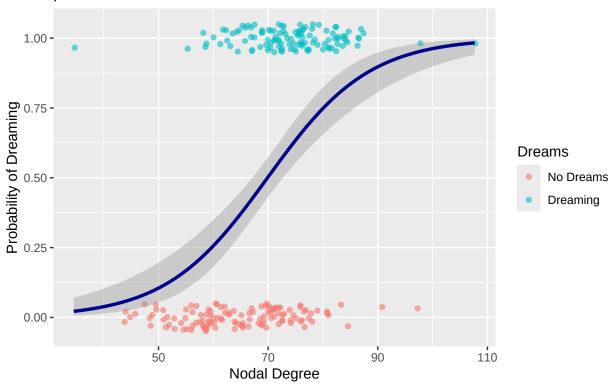
While statistical differences don't always translate to predictive power, I want to see how a model might do, let's try logistic regression.

```
suppressPackageStartupMessages({library(pROC)})
# 1) Split the data
set.seed(222) # I'll set a seed for hold-out reproducibility
train_index <- sample(seq_len(nrow(processed_data)), size = 0.8 * nrow(processed_data)) # 80% train
train_data <- processed_data[train_index, ]
test_data <- processed_data[-train_index, ] # 20% test</pre>
```

```
logit_model <- glm(</pre>
  Dreams ~ degree, # Dream is target, degree is our predictor
  data = train_data,
 family = binomial() # Sets glm() to a logistic regression
model_summary <- summary(logit_model)</pre>
# Lets visualize the logistic (sigmoid) curve and p-value
ggplot(train_data, aes(x = degree, y = as.numeric(Dreams == "Dreaming"))) +
  # Using geom_jitter to offset points to avoid clutter
  geom_jitter(aes(color = Dreams), height = 0.05, width = 0.5, alpha = 0.6) +
  geom_smooth(
   method
              = "glm",
    method.args = list(family = "binomial"),
           = TRUE,
= "darkblue",
   color
   linewidth = 1.2
  ) +
 labs(
           = "Nodal Degree",
   X
          = "Probability of Dreaming",
   title = "Logistic Regression: Nodal Degree vs Dreaming Probability (Train Set)",
    # Extract p-value from coefficient #2 (degree)
    subtitle = paste("p =", format.pval(model_summary$coefficients[2, 4], digits = 3))
  )
```

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

Logistic Regression: Nodal Degree vs Dreaming Probability (Train Set) p = 5.48e-10

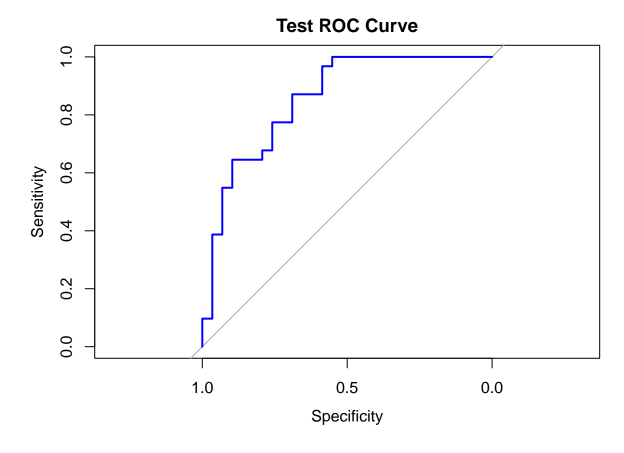


```
# Evaluation on test data
test_predictions <- predict(logit_model, newdata = test_data, type = "response")
test_roc <- roc(
    response = as.numeric(test_data$Dreams == "Dreaming"),
    predictor = test_predictions
)

## Setting levels: control = 0, case = 1

## Setting direction: controls < cases

plot(test_roc, main = "Test ROC Curve", col = "blue")</pre>
```



```
cat("Test AUC:", auc(test_roc), "\n")
```

Test AUC: 0.8542825

ROC of 0.85 indicates a classifier discriminating much better than a random-guessing model $\#(illustrated\ via\ diagonal\ line)$