

Biosciences_Project_RMarkdown_Code_Version1

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2024-01-23

```
#Import the libraries and the Dataset
```

```
#import the used library and the csv Data File
```

```
library(dplyr)
```

```
##
```

```
## Attaching package: 'dplyr'
```

```
## The following objects are masked from 'package:stats':
```

```
##
```

```
##      filter, lag
```

```
## The following objects are masked from 'package:base':
```

```
##
```

```
##      intersect, setdiff, setequal, union
```

```
library(lubridate)
```

```
##
```

```
## Attaching package: 'lubridate'
```

```
## The following objects are masked from 'package:base':
```

```
##
```

```
##      date, intersect, setdiff, union
```

```
library(ggplot2)
```

```
library(tidyr)
```

```
library(scales)
```

```
library(MASS)
```

```
## Warning: package 'MASS' was built under R version 4.2.3
```

```
##
```

```
## Attaching package: 'MASS'
```

```
## The following object is masked from 'package:dplyr':
```

```
##
```

```
##      select
```

```
data <- read.csv ("Biosciences_Publications_Dataset_2024.csv")
head(data)
```

```
##
## 1          ArdA proteins from different mobile genetic elements can bind to the EcoKI Type
## 2          Bacteria repelling poly(methylmethacrylate-co-dimethylacry
## 3          Exosomes secreted by nematode parasites transfer small RNAs to mamma
## 4          Parasite-Derived MicroRNAs in Host Serum As
## 5 Characterisation of Translation Elongation Factor eEF1B Subunit Expression in Mammalian Cells and '
## 6          The translation elongation factor eEF1A1 couples transcription to
##          DOI ResGrp Type
## 1 https://doi.org/10.1016/j.bbapap.2013.12.008 Blakely InfD
## 2 https://doi.org/10.1039/C4TB01129E Blakely InfD
## 3 https://www.nature.com/articles/ncomms6488 Buck InfD
## 4 https://doi.org/10.1371/journal.pntd.0002701 Buck InfD
## 5 10.1371/journal.pone.0114117 Abbott NCD
## 6 10.7554/eLife.03164 Abbott NCD
##          Institution
## 1 Edinburgh Infectious Diseases Organisation
## 2 Edinburgh Infectious Diseases Organisation
## 3 Edinburgh Infectious Diseases Organisation
## 4 Edinburgh Infectious Diseases Organisation
## 5 IGC (Centre for Genomic and Experimental Medicine)
## 6 IGC (Centre for Genomic and Experimental Medicine)
##          Journal
## 1 Biochimica et Biophysica Acta (BBA) - Proteins and Proteomics (ScienceDirect)
## 2 Journal of Materials Chemistry B (The Royal Society of Chemistry)
## 3 Nature Communications (Nature)
## 4 PLOS Neglected Tropical Diseases (Public Library of Science PLOS)
## 5 PLoS ONE
## 6 eLife
## Year AnalysisPgrm CodeArchived DAS CorresAuthor Preprints Complete Reuse
## 1 2014 1 NA NA 0 0 2 3
## 2 2014 1 NA NA 0 0 2 2
## 3 2014 1 0 NA 1 0 2 4
## 4 2014 1 1 NA 1 0 4 4
## 5 2014 1 NA 1 0 0 1 1
## 6 2014 1 NA NA 0 0 1 1
## Access Licence Image Genomics Rescored
## 1 2 4 0 0 0
## 2 2 4 0 NA 0
## 3 4 4 NA 1 0
## 4 4 4 NA 1 0
## 5 1 1 0 NA 0
## 6 1 1 0 NA 0
```

#1 - Descriptive Statistics

#How many papers in each type

```
type_frequency <- table(data$Type)
print(type_frequency) #these results were used in the table s1 of the publication
```

```
##
```

```
## InfD  NCD
##    82  111
```

#frequency and percentage of paper in each year

```
total_counts_by_type <- data %>%
  group_by(Type) %>%
  summarise(Total = n(), .groups = "drop")

# Calculate frequency and percentage for each Type and Year
summarized_data <- data %>%
  group_by(Year, Type) %>%
  summarise(count = n(), .groups = "drop") %>%
  left_join(total_counts_by_type, by = "Type") %>%
  mutate(Percentage = (count / Total) * 100)

print(summarized_data) #these results were used in the table s1 of the publication
```

```
## # A tibble: 18 x 5
##   Year Type  count Total Percentage
##   <int> <chr> <int> <int>      <dbl>
## 1  2014 InfD     13    82      15.9
## 2  2014 NCD      7   111      6.31
## 3  2015 InfD      6    82      7.32
## 4  2015 NCD      9   111      8.11
## 5  2016 InfD      6    82      7.32
## 6  2016 NCD      7   111      6.31
## 7  2017 InfD      9    82     11.0
## 8  2017 NCD     17   111     15.3
## 9  2018 InfD     10    82     12.2
## 10 2018 NCD      8   111      7.21
## 11 2019 InfD     12    82     14.6
## 12 2019 NCD     15   111     13.5
## 13 2020 InfD     11    82     13.4
## 14 2020 NCD     11   111      9.91
## 15 2021 InfD      7    82      8.54
## 16 2021 NCD     18   111     16.2
## 17 2022 InfD      8    82      9.76
## 18 2022 NCD     19   111     17.1
```

Calculate the total number of papers for each year

```
yearly_totals <- data %>%
  group_by(Year) %>%
  summarise(Total = n(), .groups = "drop")

# Calculate the grand total of all papers
grand_total <- sum(yearly_totals$Total)

# Add a column for the percentage of each year's total relative to the grand total
```

```
yearly_totals <- yearly_totals %>%
  mutate(Percentage = (Total / grand_total) * 100)

# Print the yearly totals and percentages
print(yearly_totals)
```

```
## # A tibble: 9 x 3
##   Year Total Percentage
##   <int> <int>     <dbl>
## 1  2014     20      10.4
## 2  2015     15       7.77
## 3  2016     13       6.74
## 4  2017     26      13.5
## 5  2018     18       9.33
## 6  2019     27      14.0
## 7  2020     22      11.4
## 8  2021     25      13.0
## 9  2022     27      14.0
```

#to categorize papers based on the year of publication compared to FAIR principles implementation 2016 (overall)

```
data <- data %>%
  mutate(Period = ifelse(Year <= 2016, "Before or in 2016", "After 2016"))

# Calculate the total number of papers for each period
period_totals <- data %>%
  group_by(Period) %>%
  summarise(Total = n(), .groups = "drop")

# Calculate the grand total of all papers
grand_total <- sum(period_totals$Total)

# Add a column for the percentage of each period's total relative to the grand total
period_totals <- period_totals %>%
  mutate(Percentage = (Total / grand_total) * 100)

# Print the period totals and percentages
print(period_totals)
```

```
## # A tibble: 2 x 3
##   Period          Total Percentage
##   <chr>          <int>     <dbl>
## 1 After 2016        145       75.1
## 2 Before or in 2016   48       24.9
```

#to categorize papers based on the year of publication compared to FAIR principles implementation 2016 (depend on type)

```
#Frequency and Percentage of paper before and After FAIR principles (2016)
total_counts_by_type <- data %>%
  group_by(Type) %>%
```

```

summarise(Total = n(), .groups = "drop")

# Categorize, calculate frequency and percentage for each Type based on year
summarized_data <- data %>%
  mutate(Category = ifelse(Year <= 2016, "On/Before 2016", "After 2020")) %>%
  group_by(Category, Type) %>%
  summarise(count = n(), .groups = "drop") %>%
  left_join(total_counts_by_type, by = "Type") %>%
  mutate(Percentage = (count / Total) * 100)

print(summarized_data) #these results were used in the table s1 of the publication

```

```

## # A tibble: 4 x 5
##   Category      Type count Total Percentage
##   <chr>         <chr> <int> <int>      <dbl>
## 1 After 2020    InfD     57    82      69.5
## 2 After 2020    NCD     88   111      79.3
## 3 On/Before 2016 InfD     25    82      30.5
## 4 On/Before 2016 NCD     23   111      20.7

```

#to categorise papers based on the year of publication compared to COVID 19 - 2020 (overall)

```

data <- data %>%
  mutate(Period = ifelse(Year <= 2020, "Before or in 2020", "After 2020"))

# Calculate the total number of papers for each period
period_totals <- data %>%
  group_by(Period) %>%
  summarise(Total = n(), .groups = "drop")

# Calculate the grand total of all papers
grand_total <- sum(period_totals$Total)

# Add a column for the percentage of each period's total relative to the grand total
period_totals <- period_totals %>%
  mutate(Percentage = (Total / grand_total) * 100)

# Print the period totals and percentages
print(period_totals) #these results were used in the table s1 of the publication

```

```

## # A tibble: 2 x 3
##   Period      Total Percentage
##   <chr>         <int>      <dbl>
## 1 After 2020      52      26.9
## 2 Before or in 2020 141      73.1

```

#to categorise papers based on the year of publication compared to COVID 19 - 2020 (NCD vs InfD)

```

total_counts_by_type <- data %>%
  group_by(Type) %>%
  summarise(Total = n(), .groups = "drop")

```

```
# Categorize, calculate frequency and percentage for each Type based on year
summarized_data <- data %>%
  mutate(Category = ifelse(Year <= 2020, "On/Before 2020", "After 2020")) %>%
  group_by(Category, Type) %>%
  summarise(count = n(), .groups = "drop") %>%
  left_join(total_counts_by_type, by = "Type") %>%
  mutate(Percentage = (count / Total) * 100)

print(summarized_data) #these results were used in the table s1 of the publication
```

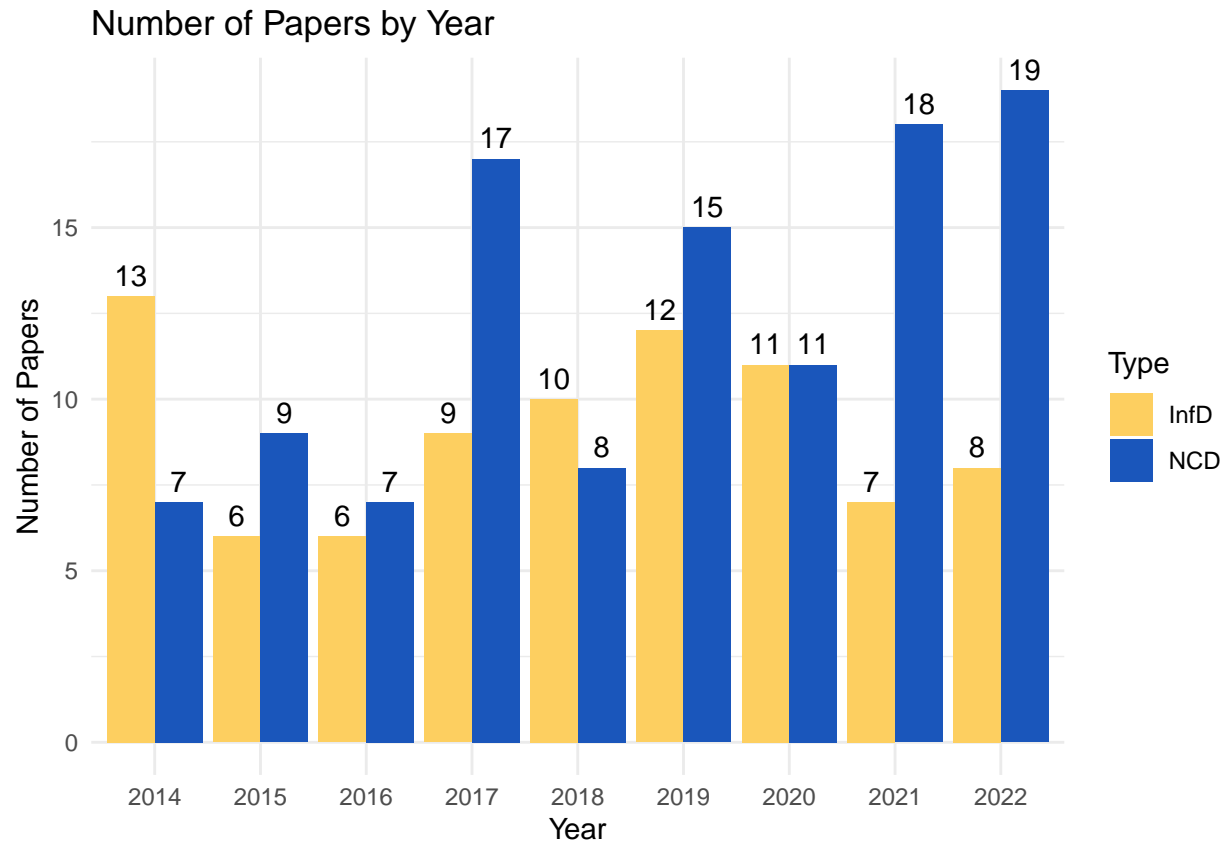
```
## # A tibble: 4 x 5
##   Category      Type count Total Percentage
##   <chr>         <chr> <int> <int>      <dbl>
## 1 After 2020    InfD     15     82       18.3
## 2 After 2020    NCD      37    111       33.3
## 3 On/Before 2020 InfD     67     82       81.7
## 4 On/Before 2020 NCD      74    111       66.7
```

#Plot the distribution of the included papers over the years

```
aggregated_data <- data %>%
  group_by(Year, Type) %>%
  summarise(count = n(), .groups = "drop")

# Create the clustered bar chart with numbers on top of the bars and specified colors
year <- ggplot(aggregated_data, aes(x = as.factor(Year), y = count, fill = Type)) +
  geom_bar(stat = "identity", position = "dodge") +
  geom_text(aes(label = count), position = position_dodge(width = 0.9), vjust = -0.5) +
  scale_fill_manual(values = c("#FDCF60", "#1A56BB")) +
  labs(x = "Year", y = "Number of Papers", title = "Number of Papers by Year") +
  theme_minimal()

print(year)
```



```
ggsave("year.png", year, width = 10, height = 8, dpi = 300, bg="white")
```

Frequency and Percentage of the papers that share data (Data Sharing= Data Completeness score >1) Overall

```
data <- data %>%
  mutate(Period = ifelse(Complete > 1, "Completeness = 1", "Completeness > 1"))

# Calculate the total number of papers for each period
period_totals <- data %>%
  group_by(Period) %>%
  summarise(Total = n(), .groups = "drop")

# Calculate the grand total of all papers
grand_total <- sum(period_totals$Total)

# Add a column for the percentage of each period's total relative to the grand total
period_totals <- period_totals %>%
  mutate(Percentage = (Total / grand_total) * 100)

# Print the period totals and percentages
print(period_totals) #these results were used in the table s1 of the publication
```

```
## # A tibble: 2 x 3
##   Period      Total Percentage
##   <chr>      <int>      <dbl>
## 1 Completeness = 1    151      78.2
## 2 Completeness > 1     42      21.8
```

Frequency and Percentage of the papers that share data (Data Sharing= Data Completeness score >1) by Type

```
total_counts_by_type <- data %>%
  group_by(Type) %>%
  summarise(Total = n(), .groups = "drop")

# Categorize based on Completeness, calculate frequency and percentage for each Type
summarized_data <- data %>%
  filter(Complete == 1 | Complete > 1) %>%
  mutate(Category = ifelse(Complete == 1, "Completeness = 1", "Completeness > 1")) %>%
  group_by(Category, Type) %>%
  summarise(count = n(), .groups = "drop") %>%
  left_join(total_counts_by_type, by = "Type") %>%
  mutate(Percentage = (count / Total) * 100)

print(summarized_data) #these results were used in the table s1 of the publication
```

```
## # A tibble: 4 x 5
##   Category      Type  count Total Percentage
##   <chr>      <chr> <int> <int>      <dbl>
## 1 Completeness = 1 InfD     19     82      23.2
## 2 Completeness = 1 NCD     23    111      20.7
## 3 Completeness > 1 InfD     63     82      76.8
## 4 Completeness > 1 NCD     88    111      79.3
```

#Chi square test to assess if there is any difference in data share between the two types (InfD vs NCD)

```
# Create a contingency table
table_completeness <- table(data$Type, data$Complete > 1)

# Perform the Chi-square test
result <- chisq.test(table_completeness)

#Print the chi-square results
print(result)
```

```
##
## Pearson's Chi-squared test with Yates' continuity correction
##
## data:  table_completeness
## X-squared = 0.053503, df = 1, p-value = 0.8171
```



```

# Get the expected values
expected_values <- result$expected

# Print the expected values
print(expected_values) ##in order to check the assumption of chi square test: Each cell in the contingency table has an expected count of 5 or more.

##
##          FALSE      TRUE
## InfD 17.84456 64.15544
## NCD  24.15544 86.84456

```

```

#The assumptions of that test were met:

#### Sample Size: Each cell in the contingency table has an expected count of 5 or more.
#### Independence: The observations are independent of each other. This means that the selection of one observation does not affect the selection of another.
#### Random Sampling: The data is a random sample.
#### Categorical Data: Both variables should be categorical (either nominal or ordinal).

```

#Frequency and percentage of preprints

```

# Calculate frequency and percentage for each Preprints category, across all types
preprints_summary <- data %>%
  group_by(Preprints) %>%
  summarise(count = n(), .groups = "drop")

# Calculate the overall total
overall_total <- sum(preprints_summary$count)

# Add a column for the percentage of each Preprints category relative to the overall total
preprints_summary <- preprints_summary %>%
  mutate(percentage = (count / overall_total) * 100)

# Print the summary
print(preprints_summary)

```

```

## # A tibble: 2 x 3
##   Preprints count percentage
##   <int> <int>     <dbl>
## 1     0  149     77.2
## 2     1   44     22.8

```

```

# Frequency and Percentage of the papers with preprints
data %>%
  group_by(Type, Preprints) %>%
  summarise(count = n(), .groups = "drop") %>%
  group_by(Type) %>%
  mutate(total = sum(count),
         percentage = (count/total)*100) %>%
  ungroup() %>%
  arrange(Type, Preprints)

```

```
## # A tibble: 4 x 5
##   Type Preprints count total percentage
##   <chr>      <int> <int> <int>      <dbl>
## 1 InfD         0     65     82       79.3
## 2 InfD         1     17     82       20.7
## 3 NCD          0     84    111       75.7
## 4 NCD          1     27    111       24.3
```

```
#these results were used in the table s1 of the publication
```

```
#Chi square test to assess if there is any difference in preprint between the two types (InfD, NCD)
```

```
# Create a contingency table
table_preprint <- table(data$Type, data$Preprints)
```

```
# Perform the Chi-square test
result <- chisq.test(table_preprint)
```

```
#Print the chi-square results
print(result)
```

```
##
## Pearson's Chi-squared test with Yates' continuity correction
##
## data: table_preprint
## X-squared = 0.17184, df = 1, p-value = 0.6785
```

```
# Get the expected values
expected_values <- result$expected
```

```
# Print the expected values
print(expected_values) ##in order to check the assumption of chi square test: Each cell in the contingency table has an expected count of 5 or more.
```

```
##
##           0           1
## InfD 63.3057 18.6943
## NCD  85.6943 25.3057
```

```
#The assumptions of that test were met:
```

```
#### Sample Size: Each cell in the contingency table has an expected count of 5 or more.
#### Independence: The observations are independent of each other. This means that the selection of one observation does not affect the selection of another.
#### Random Sampling: The data is a random sample.
#### Categorical Data: Both variables should be categorical (either nominal or ordinal).
```

Calculate frequency and percentage for DAS

```

DAS_summary <- data %>%
  group_by(DAS) %>%
  summarise(count = n(), .groups = "drop")

# Calculate the overall total
overall_total <- sum(DAS_summary$count)

# Add a column for the percentage of each DAS category relative to the overall total
DAS_summary <- DAS_summary %>%
  mutate(percentage = (count / overall_total) * 100)

print(DAS_summary)

```

```

## # A tibble: 3 x 3
##   DAS count percentage
##   <int> <int>     <dbl>
## 1     0    27     14.0
## 2     1    75     38.9
## 3    NA    91     47.2

```

```

data$DAS <- as.factor(data$DAS)

# Frequency and Percentage of the papers with DAS in the publications
data %>%
  group_by(Type, DAS) %>%
  summarise(count = n(), .groups = "drop") %>%
  group_by(Type) %>%
  mutate(total = sum(count),
         percentage = (count/total)*100) %>%
  ungroup() %>%
  arrange(Type, DAS)

```

```

## # A tibble: 6 x 5
##   Type DAS   count total percentage
##   <chr> <fct> <int> <int>     <dbl>
## 1 InfD  0       8    82      9.76
## 2 InfD  1      35    82     42.7
## 3 InfD <NA>    39    82     47.6
## 4 NCD   0      19   111     17.1
## 5 NCD   1      40   111     36.0
## 6 NCD <NA>    52   111     46.8

```

#these results were used in the table s1 of the publication

#Chi square test to assess if there is any difference in DAS between the two types (InfD, NCD)

```

# Create a contingency table
table_das <- table(data$Type, data$DAS)

# Perform the Chi-square test
result <- chisq.test(table_das)

```

```
#Print the chi-square results
print(result)
```

```
##
## Pearson's Chi-squared test with Yates' continuity correction
##
## data: table_das
## X-squared = 1.7161, df = 1, p-value = 0.1902
```

```
# Get the expected values
expected_values <- result$expected
```

```
# Print the expected values
print(expected_values) ##in order to check the assumption of chi square test: Each cell in the contingency table has an expected count of 5 or more.
```

```
##
##           0           1
## InfD 11.38235 31.61765
## NCD  15.61765 43.38235
```

```
#The assumptions of that test were met:
```

```
#### Sample Size: Each cell in the contingency table has an expected count of 5 or more.
#### Independence: The observations are independent of each other. This means that the selection of one observation does not affect the selection of another.
#### Random Sampling: The data is a random sample.
#### Categorical Data: Both variables should be categorical (either nominal or ordinal).
```

Calculate frequency and percentage for Code Sharing

```
CodeArchived_summary <- data %>%
  group_by(CodeArchived) %>%
  summarise(count = n(), .groups = "drop")

# Calculate the overall total
overall_total <- sum(CodeArchived_summary$count)

# Add a column for the percentage of each DAS category relative to the overall total
CodeArchived_summary <- CodeArchived_summary %>%
  mutate(percentage = (count / overall_total) * 100)

print(CodeArchived_summary)
```

```
## # A tibble: 3 x 3
##   CodeArchived count percentage
##       <int> <int>      <dbl>
## 1         0    21      10.9
## 2         1    48      24.9
## 3        NA   124      64.2
```

```

# Frequency and Percentage of the papers that shared the code used
data %>%
  group_by(Type, CodeArchived) %>%
  summarise(count = n(), .groups = "drop") %>%
  group_by(Type) %>%
  mutate(total = sum(count),
         percentage = (count/total)*100) %>%
  ungroup() %>%

  arrange(Type, CodeArchived)

```

```

## # A tibble: 6 x 5
##   Type CodeArchived count total percentage
##   <chr>         <int> <int> <int>      <dbl>
## 1 InfD           0      8     82       9.76
## 2 InfD           1     14     82      17.1
## 3 InfD          NA     60     82      73.2
## 4 NCD            0     13    111      11.7
## 5 NCD            1     34    111      30.6
## 6 NCD           NA     64    111      57.7

```

#these results were used in the table s1 of the publication

#Chi square test to assess if there is any difference in Code Sharing between the two types (InfD, NCD)

```

# Create a contingency table
table_code <- table(data$Type, data$CodeArchived)

# Perform the Chi-square test
result <- chisq.test(table_code)

#Print the chi-square results
print(result)

```

```

##
## Pearson's Chi-squared test with Yates' continuity correction
##
## data:  table_code
## X-squared = 0.20392, df = 1, p-value = 0.6516

```

```

# Get the expected values
expected_values <- result$expected

```

```

# Print the expected values
print(expected_values) ##in order to check the assumption of chi square test: Each cell in the contingency

```

```

##
##           0           1
##   InfD  6.695652 15.30435
##   NCD   14.304348 32.69565

```

```
#The assumptions of that test were met:
```

```
#### Sample Size: Each cell in the contingency table has an expected count of 5 or more.
```

```
#### Independence: The observations are independent of each other. This means that the selection of one
```

```
### Random Sampling: The data is a random sample.
```

```
### Categorical Data: Both variables should be categorical (either nominal or ordinal).
```

```
#these results were used in the table s1 of the publication
```

```
#Plot the 4 scoring criteria for all the papers
```

```
# Convert all relevant columns to factors first to avoid the error
```

```
data <- data %>%
```

```
  mutate(across(c(Complete, Reuse, Access, Licence), as.character), .groups = "drop") # Converts to character
```

```
# Reshape data to long format
```

```
long_data <- data %>%
```

```
  pivot_longer(cols = c(Complete, Reuse, Access, Licence), names_to = "Category", values_to = "Value")
```

```
  mutate(Value = as.character(Value)) # Ensure that Value is a character
```

```
# Rename and reorder categories
```

```
long_data$Category <- recode(long_data$Category,  
                             "Complete" = "Completeness",  
                             "Reuse" = "Reusability",  
                             "Access" = "Accessibility",  
                             "Licence" = "Licence")
```

```
long_data$Category <- factor(long_data$Category, levels = c("Completeness", "Reusability", "Accessibility", "Licence"))
```

```
long_data <- long_data %>%
```

```
  mutate(Value = factor(Value, levels = c("4", "3", "2", "1")))
```

```
# Calculate counts and percentages
```

```
long_data <- long_data %>%
```

```
  group_by(Category, Value) %>%
```

```
  summarise(Count = n(), .groups = "drop") %>%
```

```
  ungroup() %>%
```

```
  group_by(Category) %>%
```

```
  mutate(Total = sum(Count),
```

```
         Percentage = Count / Total * 100) %>%
```

```
  ungroup() %>%
```

```
  arrange(Category, desc(Value)) %>%
```

```
  group_by(Category) %>%
```

```
  mutate(Label = ifelse(Percentage > 5, paste0(round(Percentage, 1), "%"), ""),
```

```
         CumPercentage = cumsum(Percentage) - (0.5 * Percentage)) %>%
```

```
  ungroup()
```

```
# Plot
```

```
Figure_1 <- ggplot(long_data, aes(x = Category, y = Percentage, fill = Value)) +
```

```
  geom_bar(stat = "identity", position = "stack") +
```

```
  scale_fill_manual(values = c("#D0A136", "#36D0A1", "#50689B", "#E5696F")) +
```

```
  labs(y = "Percentage", fill = "Score") +
```

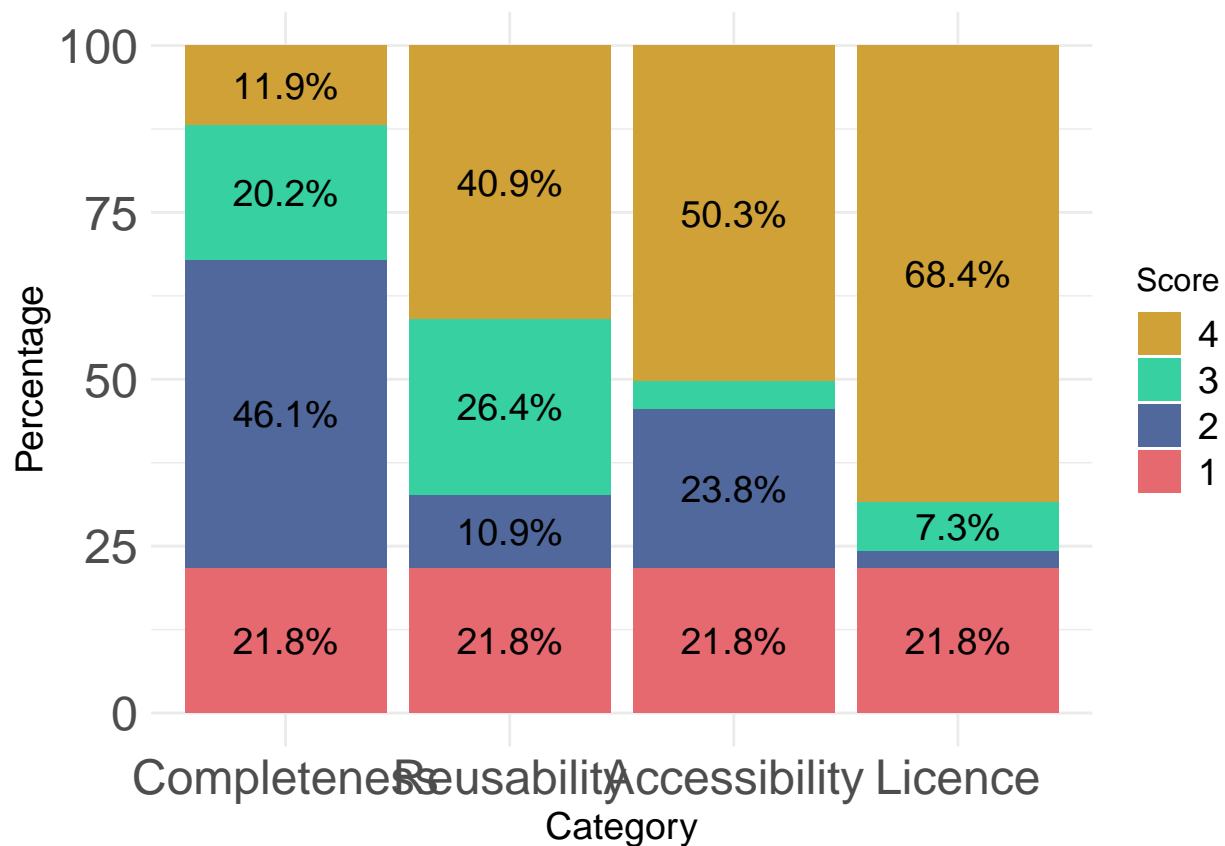
```
  geom_text(aes(label = Label, y = CumPercentage), size = 5) +
```

```

theme_minimal() +
  theme(
    axis.title = element_text(size = 14), # Increase axis titles
    axis.text = element_text(size = 18), # Increase axis text
    legend.title = element_text(size = 12), # Increase legend title
    legend.text = element_text(size = 14) # Increase legend text
  ) +
  ylim(0, 100)

# Show the plot
print(Figure_1) #these results were used in the figure 1 of the publication

```



```

# Save the plot
ggsave("Figure_1.png", Figure_1, width = 10, height = 8, dpi = 300, bg="white")

```

#Cluster bar chart fro the completeness criteria by the Type of the study

```

# Summarize data
summarized_data <- data %>%
  group_by(Type, Complete) %>%
  summarise(Count = n(), .groups = "drop") %>%
  left_join(data %>% group_by(Type) %>% summarise(Total = n(), .groups = "drop"), by = "Type") %>%
  mutate(Percentage = Count/Total * 100)

summarized_data <- summarized_data %>%

```

```

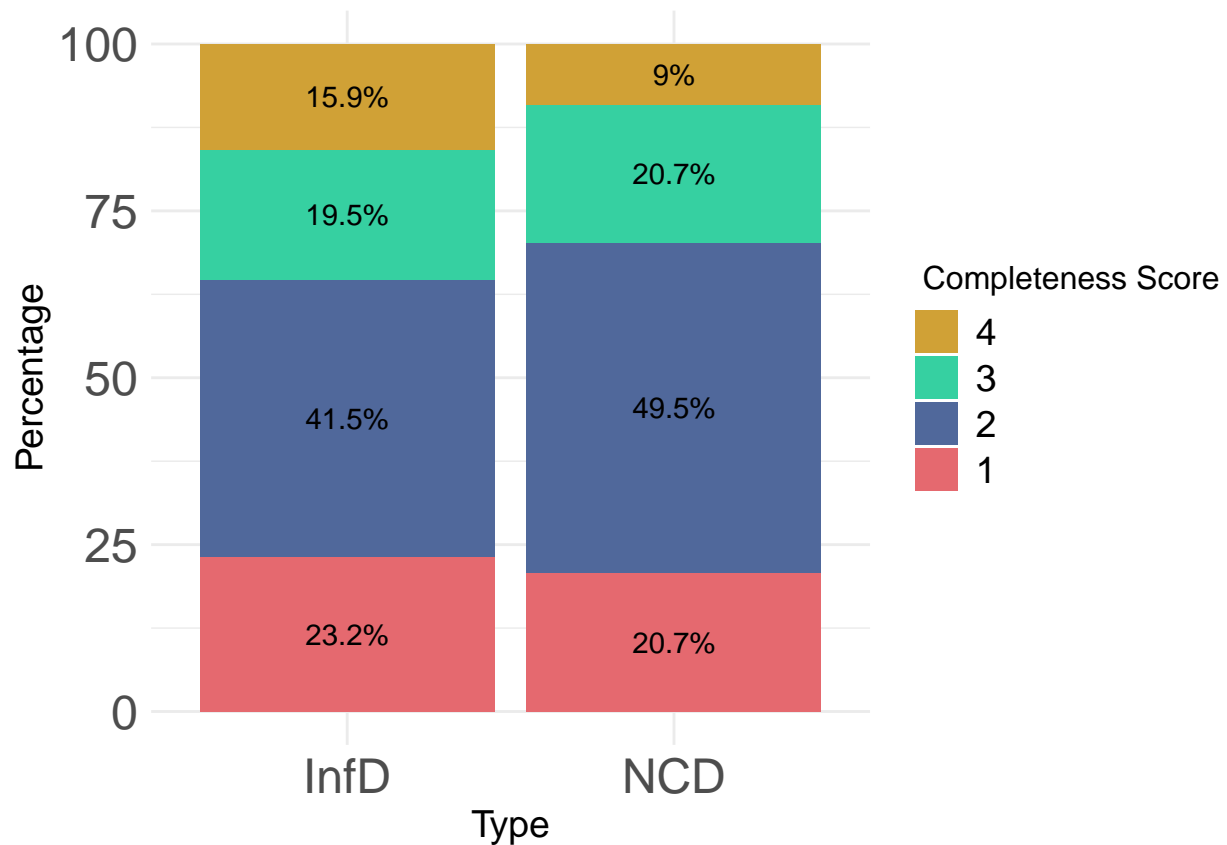
mutate(Complete = factor(Complete, levels = c("4", "3", "2", "1")))

# For cumulative percentages
summarized_data <- summarized_data %>%
  arrange(Type, desc(Complete)) %>%
  group_by(Type) %>%
  mutate(CumPercentage = cumsum(Percentage))

# Create the stacked bar chart for Completeness with colors and annotations
Figure_s1_1 <- ggplot(summarized_data, aes(x = Type, y = Percentage, fill = as.factor(Complete))) +
  geom_bar(stat = "identity", position = "stack") +
  scale_fill_manual(values = c("#D0A136", "#36D0A1", "#50689B", "#E5696F")) +
  labs(x = "Type", y = "Percentage", fill = "Completeness Score") +
  theme_minimal() +
  geom_text(aes(label = ifelse(Percentage > 5, paste0(round(Percentage, 1), "%"), ""),
    y = ifelse(Complete == 1, Percentage / 2, CumPercentage - (0.5 * Percentage))),
    color = "Black", size = 4) +
  theme(
    axis.title = element_text(size = 14), # Increase axis titles
    axis.text = element_text(size = 18), # Increase axis text
    legend.title = element_text(size = 12), # Increase legend title
    legend.text = element_text(size = 14) # Increase legend text
  ) +
  ylim(0, 100)

print(Figure_s1_1)

```

```
ggsave("Figure_s1_1.png", Figure_s1_1, width = 10, height = 8, dpi = 300)
```

#these results were used in the Figure S1 of the publication

##Cluster bar chart fro the Reuse criteria by the Type of the study

```
summarized_data_reuse <- data %>%
  group_by(Type, Reuse) %>%
  summarise(Count = n(), .groups = "drop") %>%
  left_join(data %>% group_by(Type) %>% summarise(Total = n(), .groups = "drop"), by = "Type") %>%
  mutate(Percentage = Count/Total * 100)

summarized_data_reuse <- summarized_data_reuse %>%
  mutate(Reuse = factor(Reuse, levels = c("4", "3", "2", "1")))

summarized_data_reuse <- summarized_data_reuse %>%
  arrange(Type, desc(Reuse)) %>%
  group_by(Type) %>%
  mutate(CumPercentage = cumsum(Percentage))
print(summarized_data_reuse)
```

```
## # A tibble: 8 x 6
## # Groups:   Type [2]
##   Type  Reuse Count Total Percentage CumPercentage
```

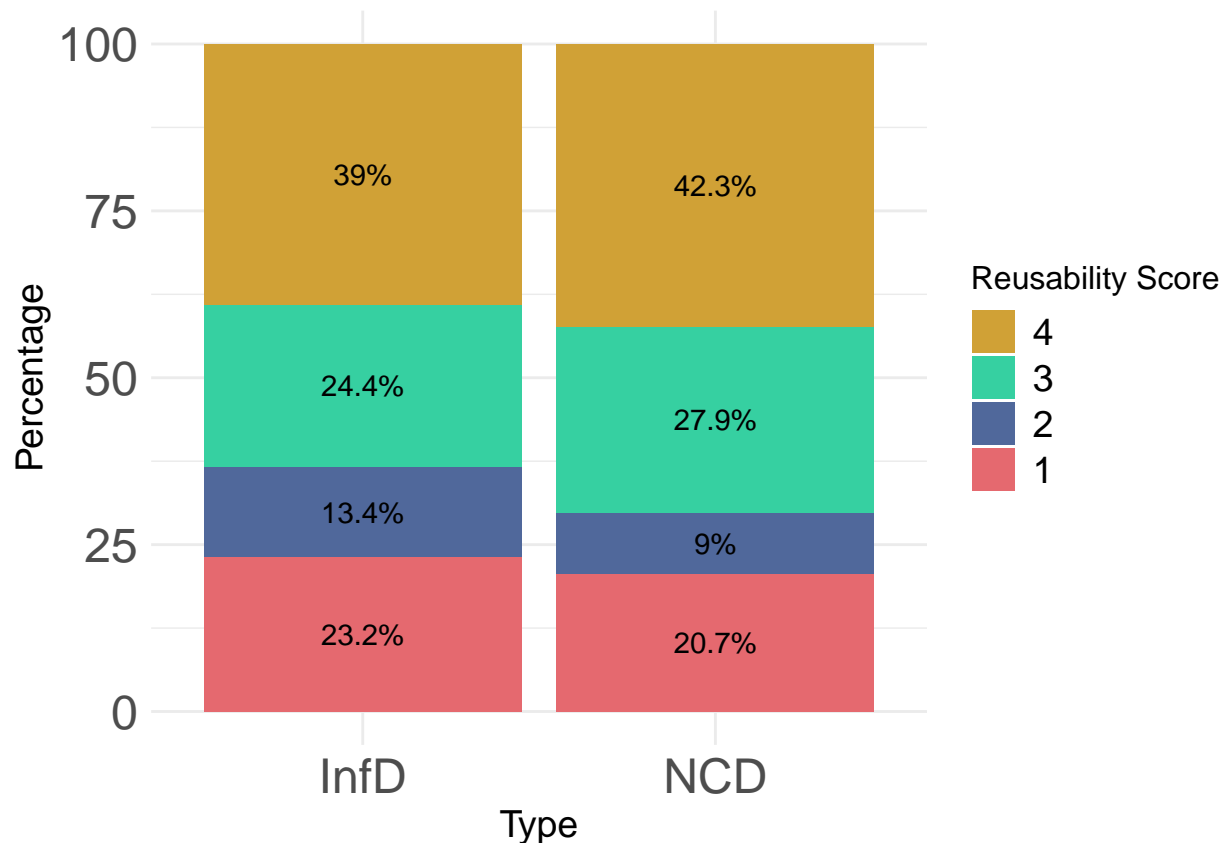
	<chr>	<fct>	<int>	<int>	<dbl>	<dbl>
## 1	InfD	1	19	82	23.2	23.2
## 2	InfD	2	11	82	13.4	36.6
## 3	InfD	3	20	82	24.4	61.0
## 4	InfD	4	32	82	39.0	100
## 5	NCD	1	23	111	20.7	20.7
## 6	NCD	2	10	111	9.01	29.7
## 7	NCD	3	31	111	27.9	57.7
## 8	NCD	4	47	111	42.3	100

```
colors <- c( "#D0A136", "#36D0A1", "#50689B", "#E5696F")
```

```
# Visualization
```

```
Figure_s1_2 <- ggplot(summarized_data_reuse, aes(x = Type, y = Percentage, fill = as.factor(Reuse))) +
  geom_bar(stat = "identity", position = "stack") +
  scale_fill_manual(values = colors) +
  labs(x = "Type", y = "Percentage", fill = "Reusability Score") +
  theme_minimal() +
  geom_text(aes(label = paste0(round(Percentage, 1), "%"),
    y = CumPercentage - (0.5 * Percentage)),
    color = "Black", size = 4) +
  theme(
    axis.title = element_text(size = 14), # Increase axis titles
    axis.text = element_text(size = 18), # Increase axis text
    legend.title = element_text(size = 12), # Increase legend title
    legend.text = element_text(size = 14) # Increase legend text
  ) +
  ylim(0, 100)

print(Figure_s1_2)
```



```
ggsave("Figure_s1_2.png", Figure_s1_2, width = 10, height = 8, dpi = 300)
```

#these results were used in the Figure S1 of the publication

#bar chart for the Accessibility criteria by the Type of the study

```
summarized_data_Access <- data %>%
  group_by(Type, Access) %>%
  summarise(Count = n(), .groups = "drop") %>%
  left_join(data %>% group_by(Type) %>% summarise(Total = n(), .groups = "drop"), by = "Type") %>%
  mutate(Percentage = Count/Total * 100)

summarized_data_Access <- summarized_data_Access %>%
  mutate(Access = factor(Access, levels = c("4", "3", "2", "1")))

summarized_data_Access <- summarized_data_Access %>%
  arrange(Type, desc(Access)) %>%
  group_by(Type) %>%
  mutate(CumPercentage = cumsum(Percentage))
print(summarized_data_Access)
```

```
## # A tibble: 8 x 6
## # Groups:   Type [2]
##   Type Access Count Total Percentage CumPercentage
##   <chr> <fct> <int> <int>     <dbl>         <dbl>
```

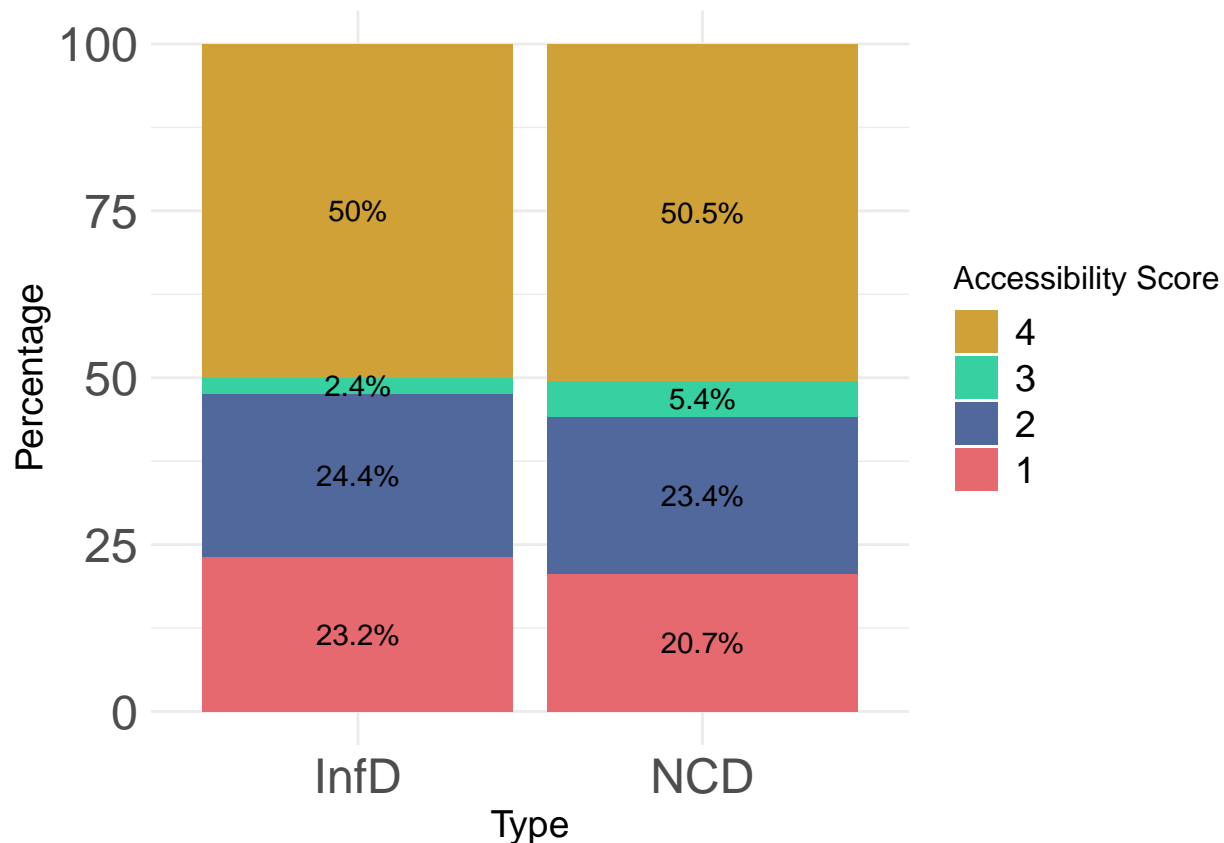
## 1	InfD	1	19	82	23.2	23.2
## 2	InfD	2	20	82	24.4	47.6
## 3	InfD	3	2	82	2.44	50
## 4	InfD	4	41	82	50	100
## 5	NCD	1	23	111	20.7	20.7
## 6	NCD	2	26	111	23.4	44.1
## 7	NCD	3	6	111	5.41	49.5
## 8	NCD	4	56	111	50.5	100

```
colors <- c("#D0A136", "#36D0A1", "#50689B", "#E5696F")
```

```
# Visualization
```

```
Figure_s1_3 <- ggplot(summarized_data_Access, aes(x = Type, y = Percentage, fill = as.factor(Access))) +
  geom_bar(stat = "identity", position = "stack") +
  scale_fill_manual(values = colors) +
  labs(x = "Type", y = "Percentage", fill = "Accessibility Score") +
  theme_minimal() +
  geom_text(aes(label = paste0(round(Percentage, 1), "%"),
    y = CumPercentage - (0.5 * Percentage)),
    color = "Black", size = 4) +
  theme(
    axis.title = element_text(size = 14), # Increase axis titles
    axis.text = element_text(size = 18), # Increase axis text
    legend.title = element_text(size = 12), # Increase legend title
    legend.text = element_text(size = 14) # Increase legend text
  ) +
  ylim(0, 100)

print(Figure_s1_3)
```



```
ggsave("Figure_s1_3.png", Figure_s1_3, width = 10, height = 8, dpi = 300)
```

#these results were used in the Figure S1 of the publication

#bar chart for the Licence criteria by the Type of the study

```
summarized_data_Licence <- data %>%
  group_by(Type, Licence) %>%
  summarise(Count = n(), .groups = "drop") %>%
  left_join(data %>% group_by(Type) %>% summarise(Total = n(), .groups = "drop"), by = "Type") %>%
  mutate(Percentage = Count/Total * 100)

summarized_data_Licence <- summarized_data_Licence %>%
  mutate(Licence = factor(Licence, levels = c("4", "3", "2", "1")))

summarized_data_Licence <- summarized_data_Licence %>%
  arrange(Type, desc(Licence)) %>%
  group_by(Type) %>%
  mutate(CumPercentage = cumsum(Percentage))
print(summarized_data_Licence)
```

```
## # A tibble: 8 x 6
## # Groups:   Type [2]
##   Type Licence Count Total Percentage CumPercentage
##   <chr> <fct>   <int> <int>      <dbl>         <dbl>
```

## 1	InfD	1	19	82	23.2	23.2
## 2	InfD	2	1	82	1.22	24.4
## 3	InfD	3	6	82	7.32	31.7
## 4	InfD	4	56	82	68.3	100
## 5	NCD	1	23	111	20.7	20.7
## 6	NCD	2	4	111	3.60	24.3
## 7	NCD	3	8	111	7.21	31.5
## 8	NCD	4	76	111	68.5	100

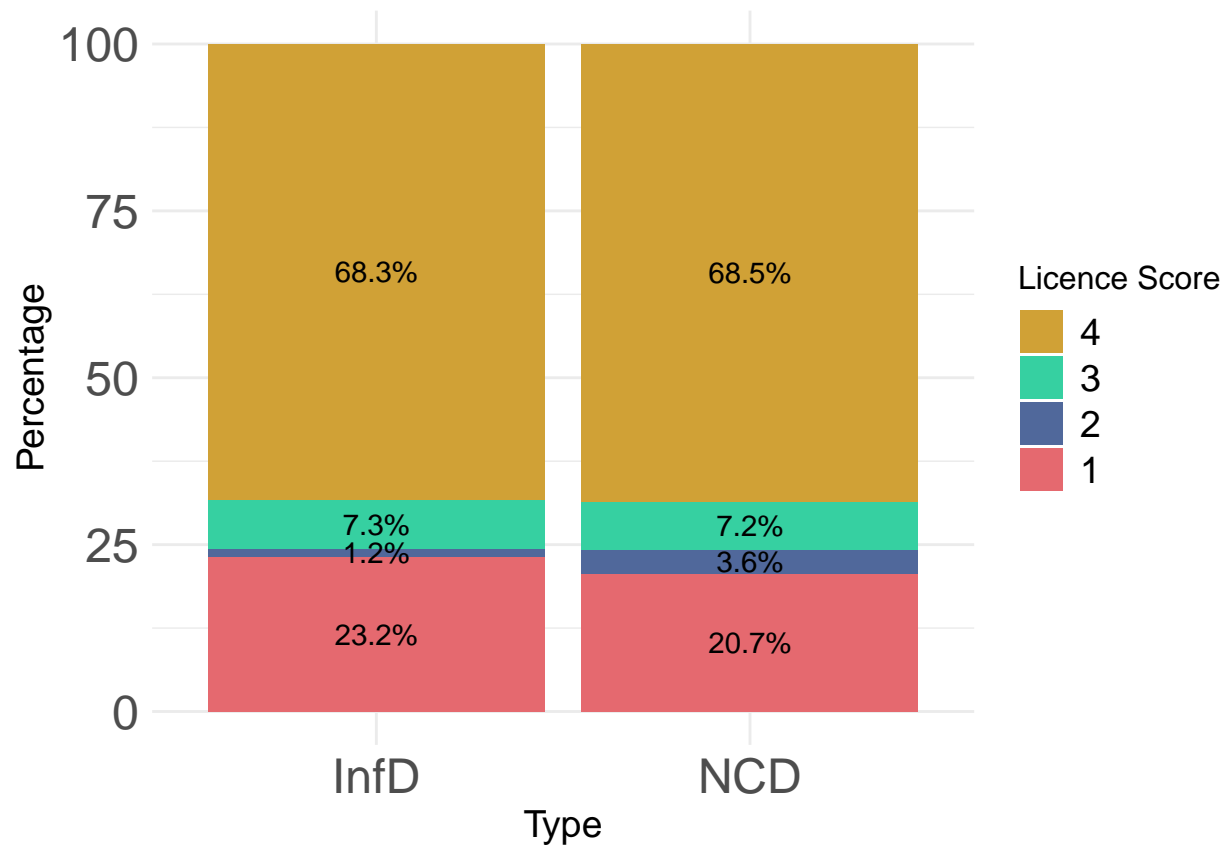
```
colors <- c("#D0A136", "#36D0A1", "#50689B", "#E5696F")
```

```
# Visualization
```

```
Figure_s1_4 <- ggplot(summarized_data_Licence, aes(x = Type, y = Percentage, fill = as.factor(Licence)))
```

```
  geom_bar(stat = "identity", position = "stack") +
  scale_fill_manual(values = colors) +
  labs(x = "Type", y = "Percentage", fill = "Licence Score") +
  theme_minimal() +
  geom_text(aes(label = paste0(round(Percentage, 1), "%"),
    y = CumPercentage - (0.5 * Percentage)),
    color = "Black", size = 4) +
  theme(
    axis.title = element_text(size = 14), # Increase axis titles
    axis.text = element_text(size = 18), # Increase axis text
    legend.title = element_text(size = 12), # Increase legend title
    legend.text = element_text(size = 14) # Increase legend text
  ) +
  ylim(0, 100)
```

```
print(Figure_s1_4)
```



```
ggsave("Figure_s1_4.png", Figure_s1_4, width = 10, height = 8, dpi = 300)
```

```
#these results were used in the Figure S1 of the publication
```

```
library(patchwork)
```

```
## Warning: package 'patchwork' was built under R version 4.2.2
```

```
##
```

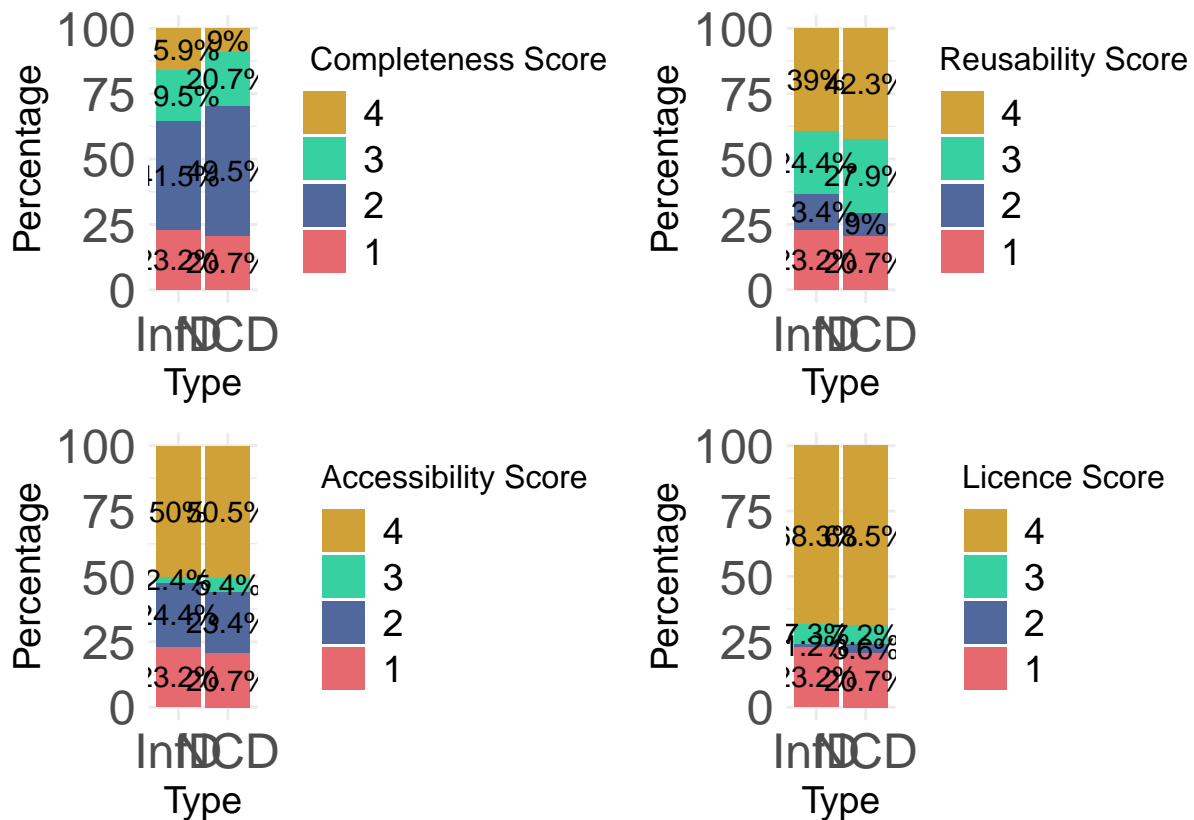
```
## Attaching package: 'patchwork'
```

```
## The following object is masked from 'package:MASS':
```

```
##
```

```
## area
```

```
Figure_s1 <- Figure_s1_1 + Figure_s1_2 + Figure_s1_3 + Figure_s1_4 +
  plot_layout(
    ncol = 2, heights = c(10, 10), widths = c(10, 10)
  )
print(Figure_s1)
```



```
ggsave("Figure_s1.png", Figure_s1_4, width = 15, height = 10, units = "in")
```

```
#these results were used in the Figure S1 of the publication
```

```
# Calculate the frequency and percentage for each 'Complete' score within each 'Year'
long_data <- data %>%
  count(Year, Complete) %>%
  group_by(Year) %>%
  mutate(Percentage = n / sum(n) * 100) %>%
  mutate(Label = ifelse(Percentage > 5, paste0(round(Percentage, 1), "%"), "")) %>%
  mutate(Cumulative_Percentage = cumsum(Percentage) - (0.5 * Percentage)) %>%
  ungroup()
```

```
long_data <- long_data %>%
  mutate(Complete = factor(Complete, levels = c("4", "3", "2", "1")))
```

```
# Custom colors
```

```
colors <- c("#D0A136", "#36D0A1", "#50689B", "#E5696F")
```

```
# Creating the stacked bar chart with percentage labels
```

```
Figure_2_1 <- ggplot(long_data, aes(x = as.factor(Year), y = n, fill = as.factor(Complete))) +
  geom_bar(stat = "identity", position = "fill") + # Scale the bar heights to proportions
  scale_y_continuous(labels = percent_format()) + # Convert the y-axis to percentage
  geom_text(
    aes(label = Label, y = Percentage),
    size = 4,
```

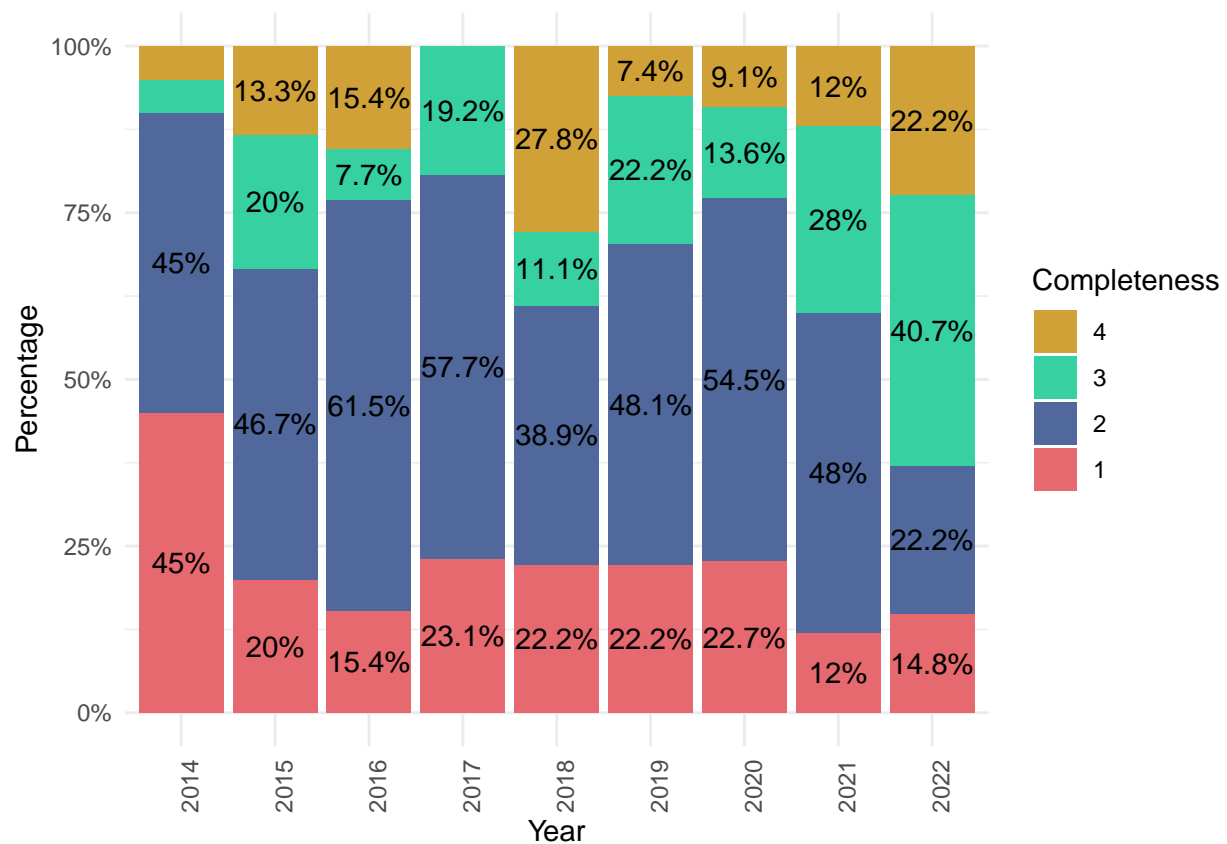


```

    color = "black",
    position = position_fill(vjust = 0.5)
) +
scale_fill_manual(values = colors) +
labs(x = "Year",
     y = "Percentage",
     fill = "Completeness") +
theme(
  axis.title = element_text(size = 14), # Increase axis titles
  axis.text = element_text(size = 20), # Increase axis text
  legend.title = element_text(size = 18), # Increase legend title
  legend.text = element_text(size = 20) # Increase legend text
) +
theme_minimal() +
scale_x_discrete(name = "Year", labels = unique(long_data$Year)) +
theme(axis.text.x = element_text(angle = 90, hjust = 1)) # Adjust text angle for readability

# Print the plot
print(Figure_2_1)

```



```

ggsave("Figure_2_1.png", Figure_2_1, width = 15, height = 10, units = "in", bg = "white")

#these results were used in the Figure 2 of the publication

```

```

# Calculate the frequency and percentage for each 'Reuse' score within each 'Year'
long_data <- data %>%
  count(Year, Reuse) %>%
  group_by(Year) %>%
  mutate(Percentage = n / sum(n) * 100) %>%
  mutate(Label = ifelse(Percentage > 5, paste0(round(Percentage, 1), "%"), "")) %>%
  mutate(Cumulative_Percentage = cumsum(Percentage) - (0.5 * Percentage)) %>%
  ungroup()

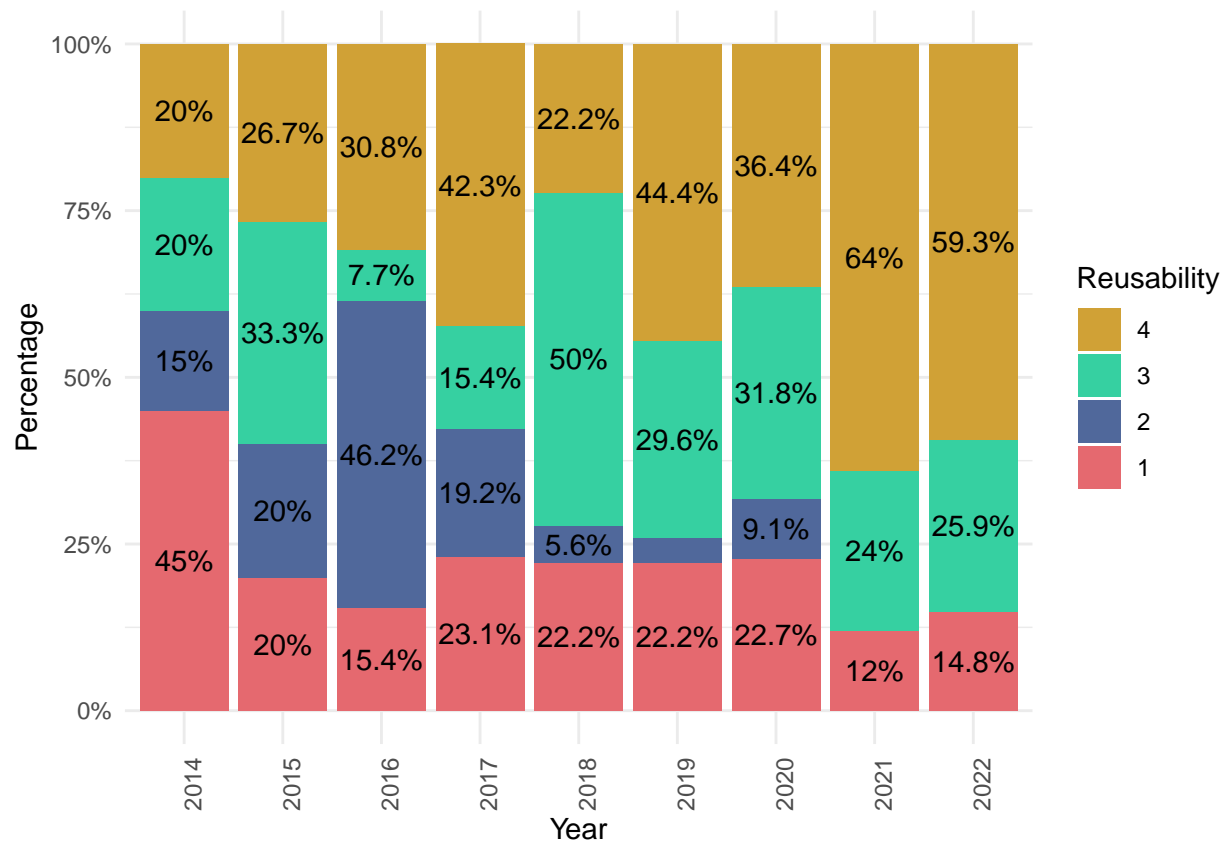
long_data <- long_data %>%
  mutate(Reuse = factor(Reuse, levels = c("4", "3", "2", "1")))

# Custom colors
colors <- c("#D0A136", "#36D0A1", "#50689B", "#E5696F")

# Creating the stacked bar chart with percentage labels
Figure_2_2 <- ggplot(long_data, aes(x = as.factor(Year), y = n, fill = as.factor(Reuse))) +
  geom_bar(stat = "identity", position = "fill") + # Scale the bar heights to proportions
  scale_y_continuous(labels = percent_format()) + # Convert the y-axis to percentage
  geom_text(
    aes(label = Label, y = Percentage),
    size = 4,
    color = "black",
    position = position_fill(vjust = 0.5)
  ) +
  scale_fill_manual(values = colors) +
  labs(x = "Year",
       y = "Percentage",
       fill = "Reusability") +
  theme(
    axis.title = element_text(size = 14), # Increase axis titles
    axis.text = element_text(size = 20), # Increase axis text
    legend.title = element_text(size = 18), # Increase legend title
    legend.text = element_text(size = 20) # Increase legend text
  ) +
  theme_minimal() +
  scale_x_discrete(name = "Year", labels = unique(long_data$Year)) +
  theme(axis.text.x = element_text(angle = 90, hjust = 1)) # Adjust text angle for readability

# Print the plot
print(Figure_2_2)

```



```
ggsave("Figure_2_2.png", Figure_2_2, width = 15, height = 10, units = "in", bg = "white")
```

#these results were used in the Figure 2 of the publication

```
# Calculate the frequency and percentage for each 'Access' score within each 'Year'
long_data <- data %>%
  count(Year, Access) %>%
  group_by(Year) %>%
  mutate(Percentage = n / sum(n) * 100) %>%
  mutate(Label = ifelse(Percentage > 5, paste0(round(Percentage, 1), "%"), "")) %>%
  mutate(Cumulative_Percentage = cumsum(Percentage) - (0.5 * Percentage)) %>%
  ungroup()
```

```
long_data <- long_data %>%
  mutate(Access = factor(Access, levels = c("4", "3", "2", "1")))
```

Custom colors

```
colors <- c("#D0A136", "#36D0A1", "#50689B", "#E5696F")
```

Creating the stacked bar chart with percentage labels

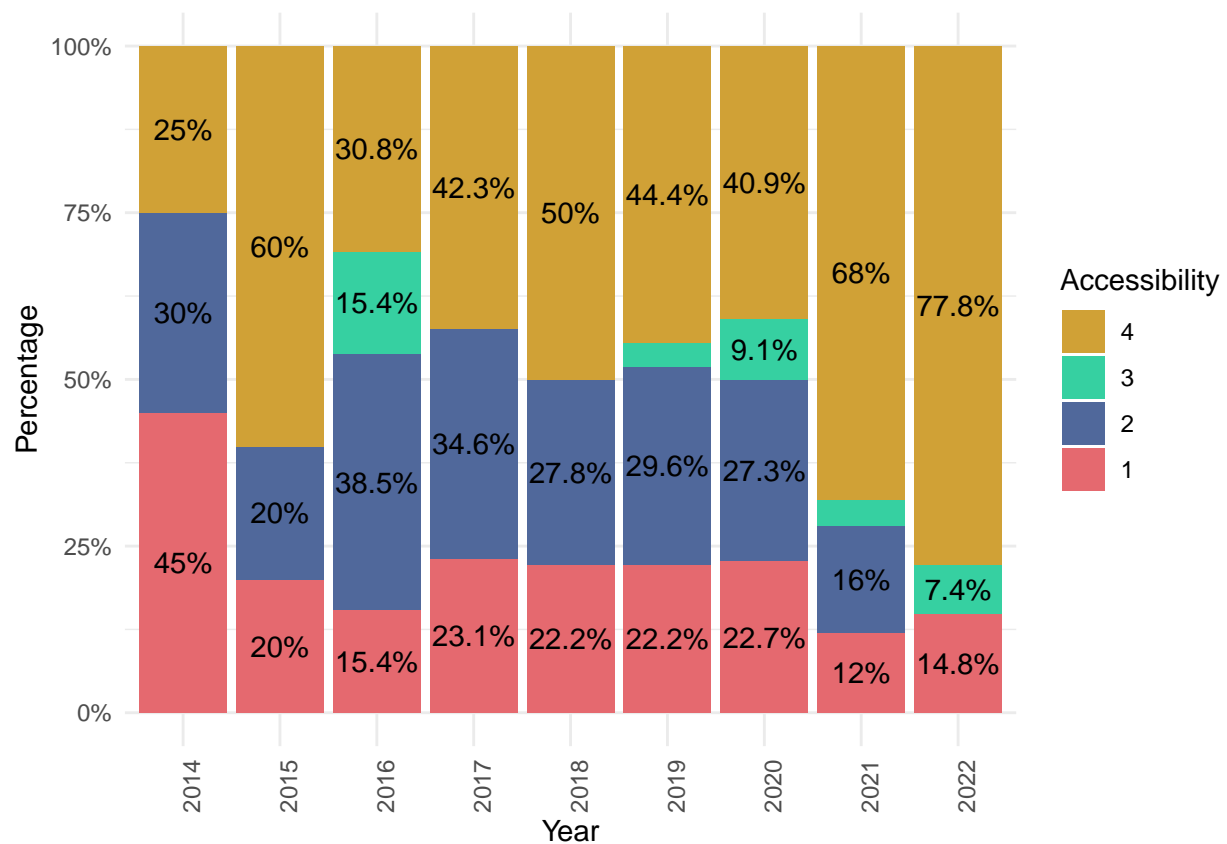
```
Figure_2_3 <- ggplot(long_data, aes(x = as.factor(Year), y = n, fill = as.factor(Access))) +
  geom_bar(stat = "identity", position = "fill") + # Scale the bar heights to proportions
  scale_y_continuous(labels = percent_format()) + # Convert the y-axis to percentage
  geom_text(
    aes(label = Label, y = Percentage),
    size = 4,
```

```

    color = "black",
    position = position_fill(vjust = 0.5)
) +
scale_fill_manual(values = colors) +
labs(x = "Year",
     y = "Percentage",
     fill = "Accessibility") +
theme(
  axis.title = element_text(size = 14), # Increase axis titles
  axis.text = element_text(size = 20), # Increase axis text
  legend.title = element_text(size = 18), # Increase legend title
  legend.text = element_text(size = 20) # Increase legend text
) +
  theme_minimal() +
scale_x_discrete(name = "Year", labels = unique(long_data$Year)) +
theme(axis.text.x = element_text(angle = 90, hjust = 1)) # Adjust text angle for readability

# Print the plot
print(Figure_2_3)

```



```

ggsave("Figure_2_3.png", Figure_2_3, width = 15, height = 10, units = "in", bg = "white")

#these results were used in the Figure 2 of the publication

```

```

# Calculate the frequency and percentage for each 'Licence' score within each 'Year'
long_data <- data %>%
  count(Year, Licence) %>%
  group_by(Year) %>%
  mutate(Percentage = n / sum(n) * 100) %>%
  mutate(Label = ifelse(Percentage > 5, paste0(round(Percentage, 1), "%"), "")) %>%
  mutate(Cumulative_Percentage = cumsum(Percentage) - (0.5 * Percentage)) %>%
  ungroup()

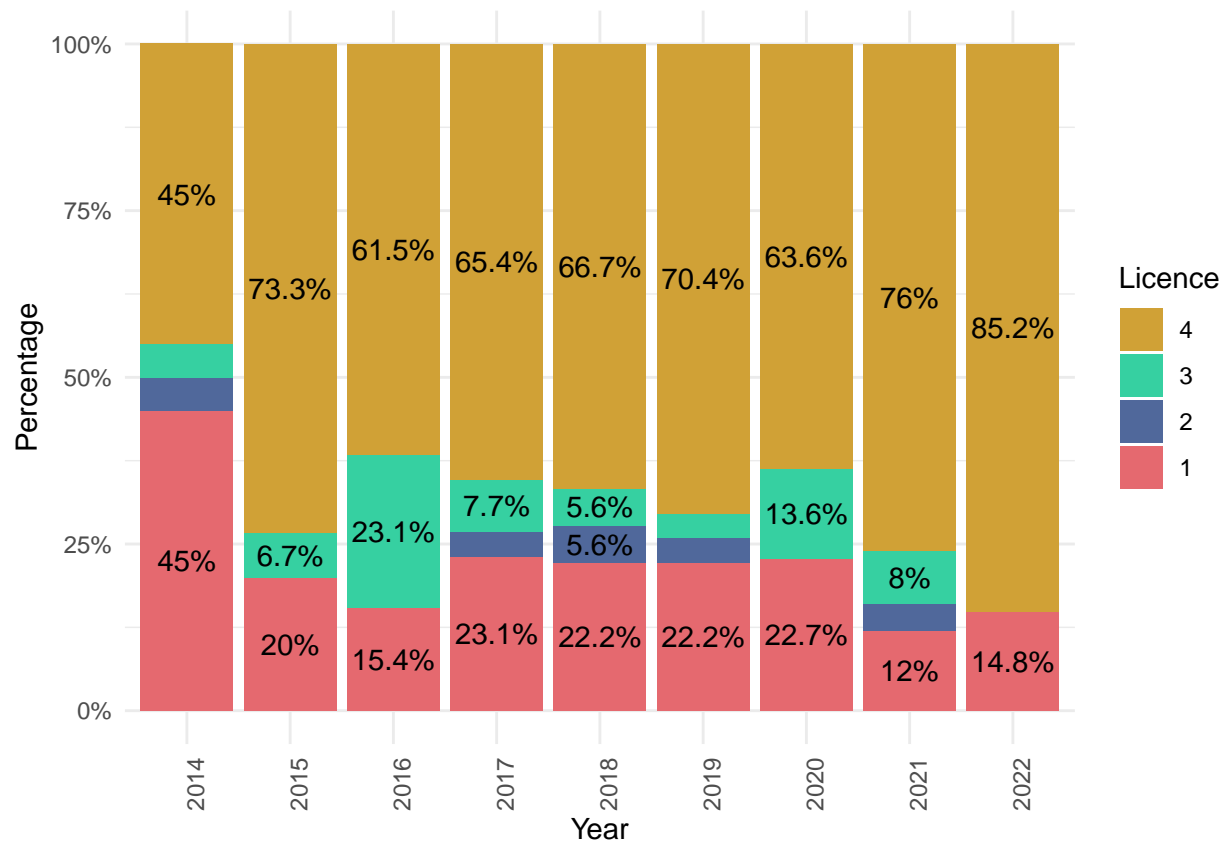
long_data <- long_data %>%
  mutate(Licence = factor(Licence, levels = c("4", "3", "2", "1")))

# Custom colors
colors <- c("#D0A136", "#36D0A1", "#50689B", "#E5696F")

# Creating the stacked bar chart with percentage labels
Figure_2_4 <- ggplot(long_data, aes(x = as.factor(Year), y = n, fill = as.factor(Licence))) +
  geom_bar(stat = "identity", position = "fill") + # Scale the bar heights to proportions
  scale_y_continuous(labels = percent_format()) + # Convert the y-axis to percentage
  geom_text(
    aes(label = Label, y = Percentage),
    size = 4,
    color = "black",
    position = position_fill(vjust = 0.5)
  ) +
  scale_fill_manual(values = colors) +
  labs(x = "Year",
       y = "Percentage",
       fill = "Licence") +
  theme(
    axis.title = element_text(size = 14), # Increase axis titles
    axis.text = element_text(size = 20), # Increase axis text
    legend.title = element_text(size = 18), # Increase legend title
    legend.text = element_text(size = 20) # Increase legend text
  ) +
  theme_minimal() +
  scale_x_discrete(name = "Year", labels = unique(long_data$Year)) +
  theme(axis.text.x = element_text(angle = 90, hjust = 1)) # Adjust text angle for readability

# Print the plot
print(Figure_2_4)

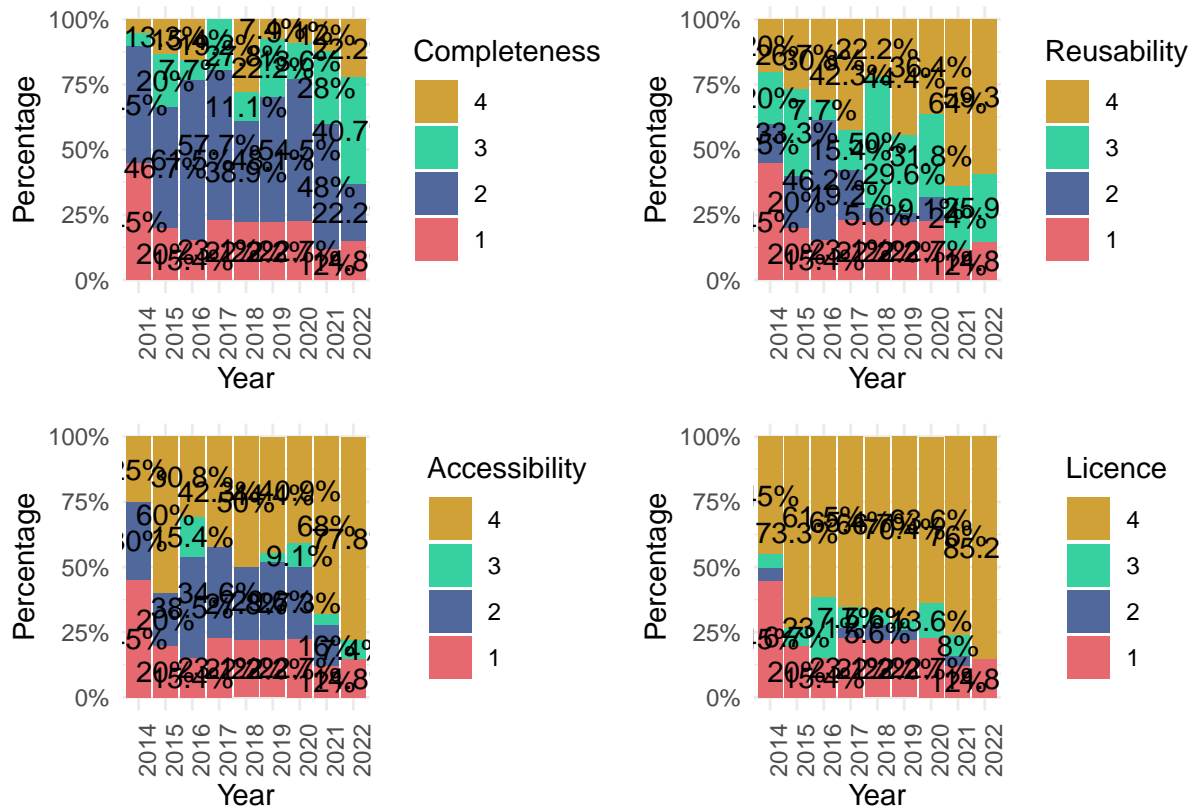
```



```
ggsave("Figure_2_4.png", Figure_2_4, width = 15, height = 10, units = "in", bg = "white")
```

#these results were used in the Figure 2 of the publication

```
Figure_2 <- Figure_2_1 + Figure_2_2 + Figure_2_3 + Figure_2_4 +
  plot_layout(
    ncol = 2, heights = c(10, 10), widths = c(10, 10)
  )
print(Figure_2)
```



```
ggsave("Figure_2.png", Figure_2, width = 15, height = 10, units = "in")
```

```
#these results were used in the Figure 2 of the publication
```

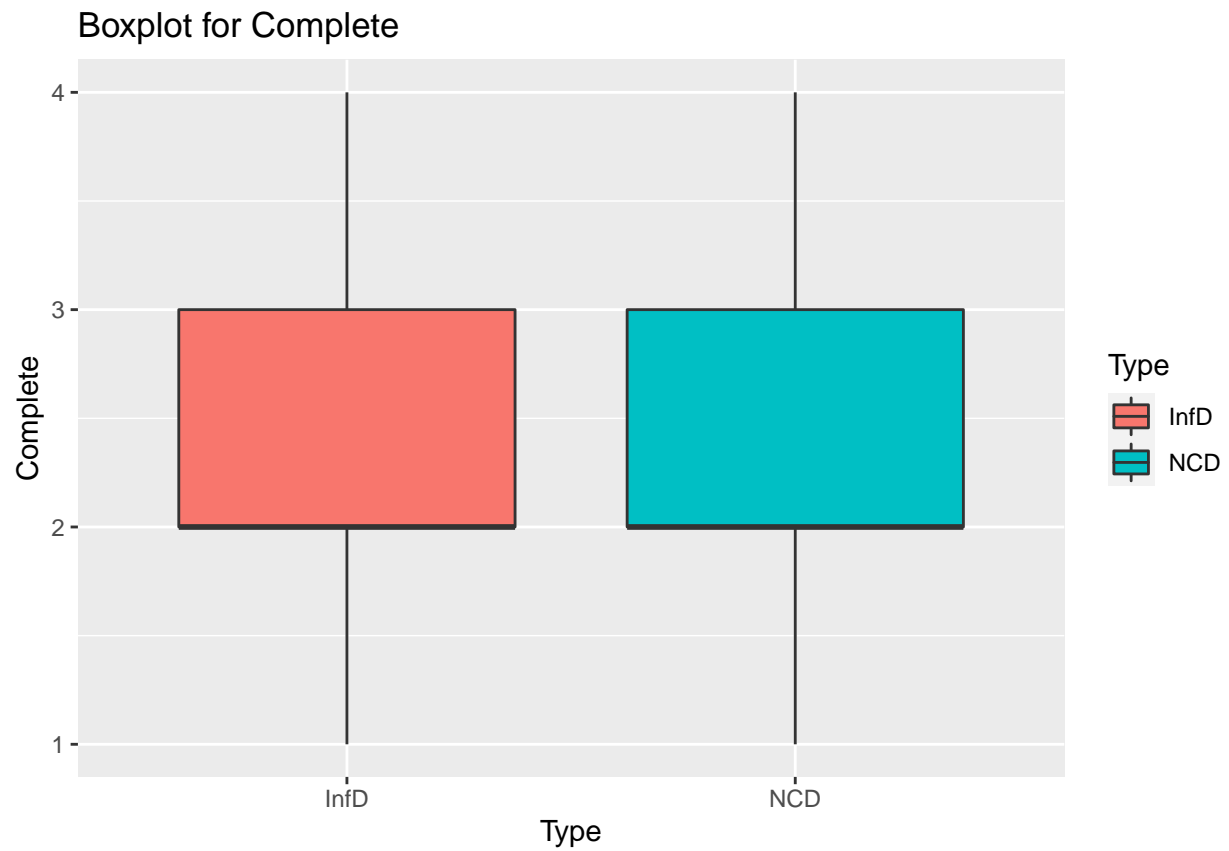
```
#study the significant difference between two type of the reseach for each scoring criteria
```

```
# Convert scoring criteria to numeric (if not already)
data <- data %>%
  mutate(across(c(Complete, Reuse, Access, Licence), ~as.numeric(as.character(.))))

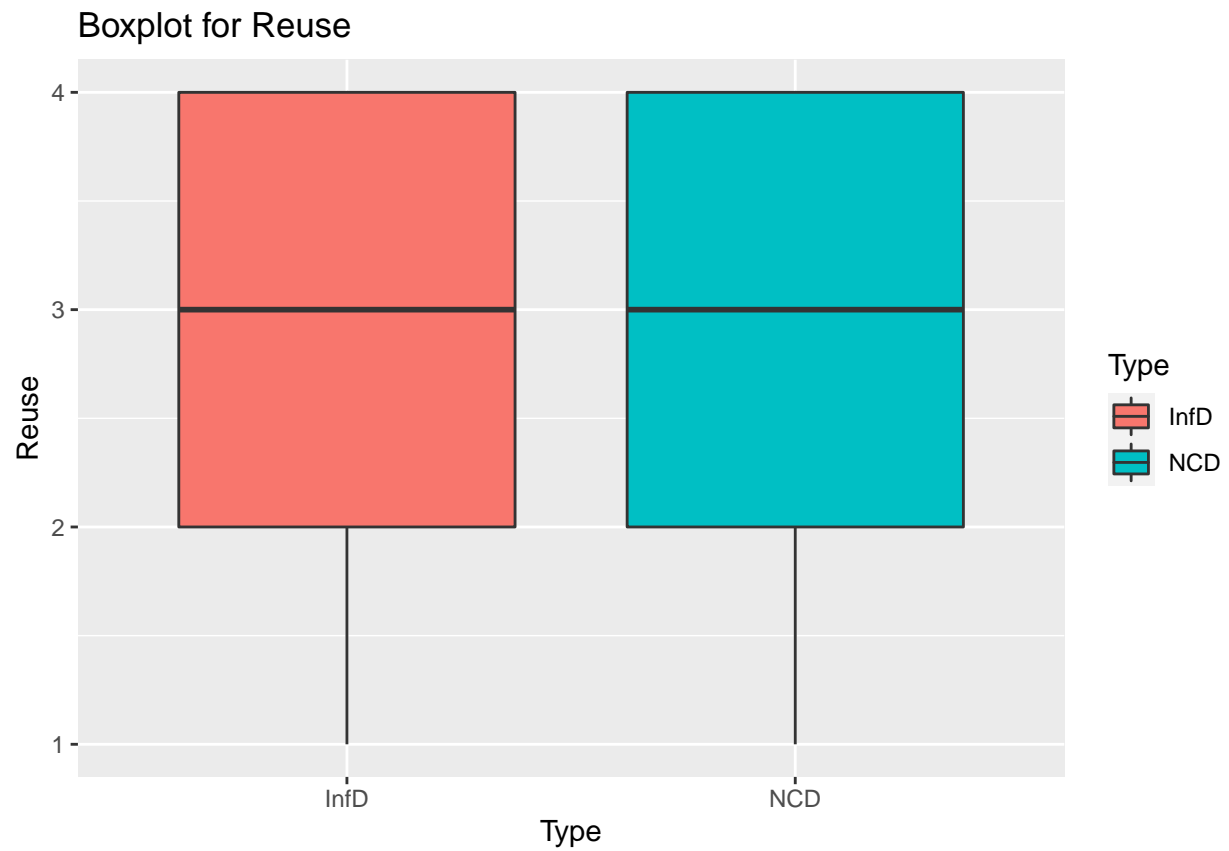
# Assumption checks for each criterion
for (criterion in c("Complete", "Reuse", "Access", "Licence")) {
  # Create and display a box plot to check the distribution shape and spread
  print(ggplot(data, aes_string(x = "Type", y = criterion, fill = "Type")) +
    geom_boxplot() +
    ggtitle(paste("Boxplot for", criterion)))

  # Print the median for a basic comparison
  cat(paste("Median of", criterion, "by Type:\n"))
  print(aggregate(. ~ Type, data[c("Type", criterion)], median))

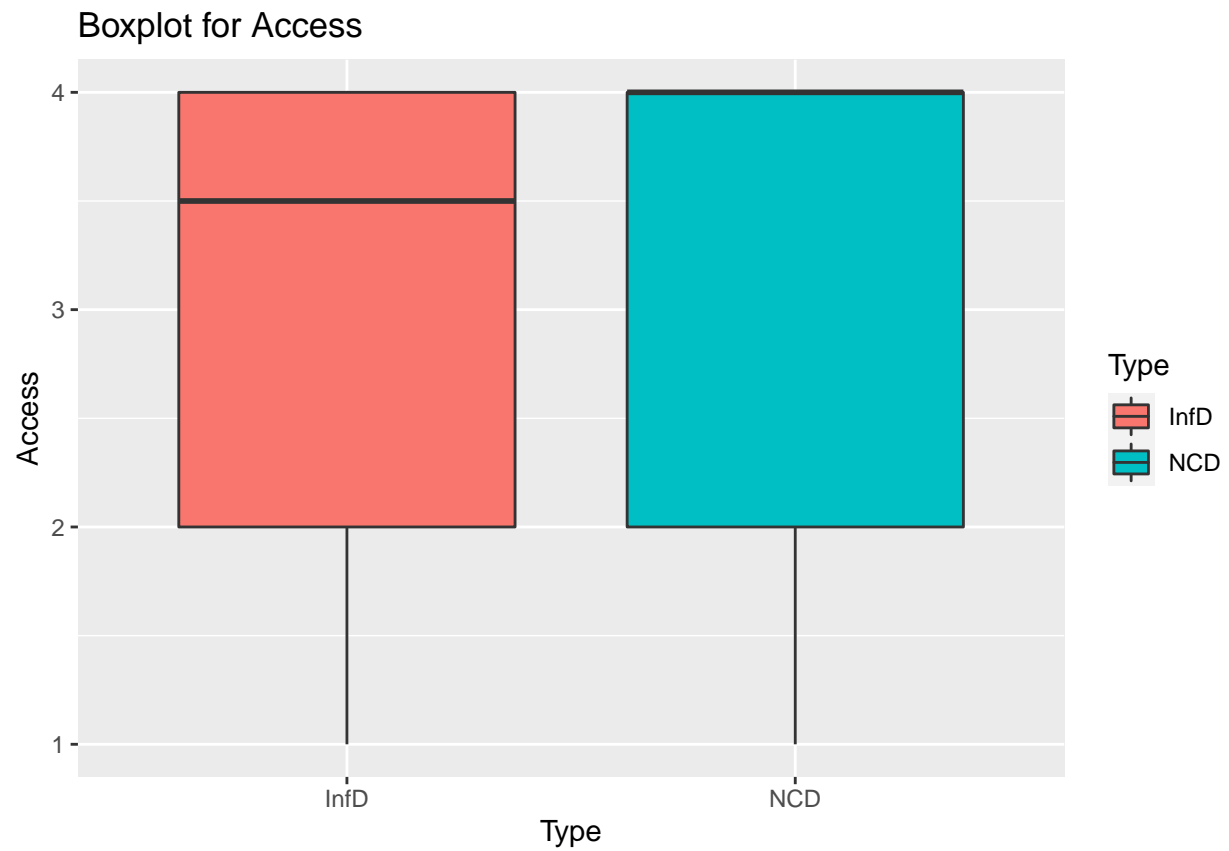
  cat("\n")
}
```



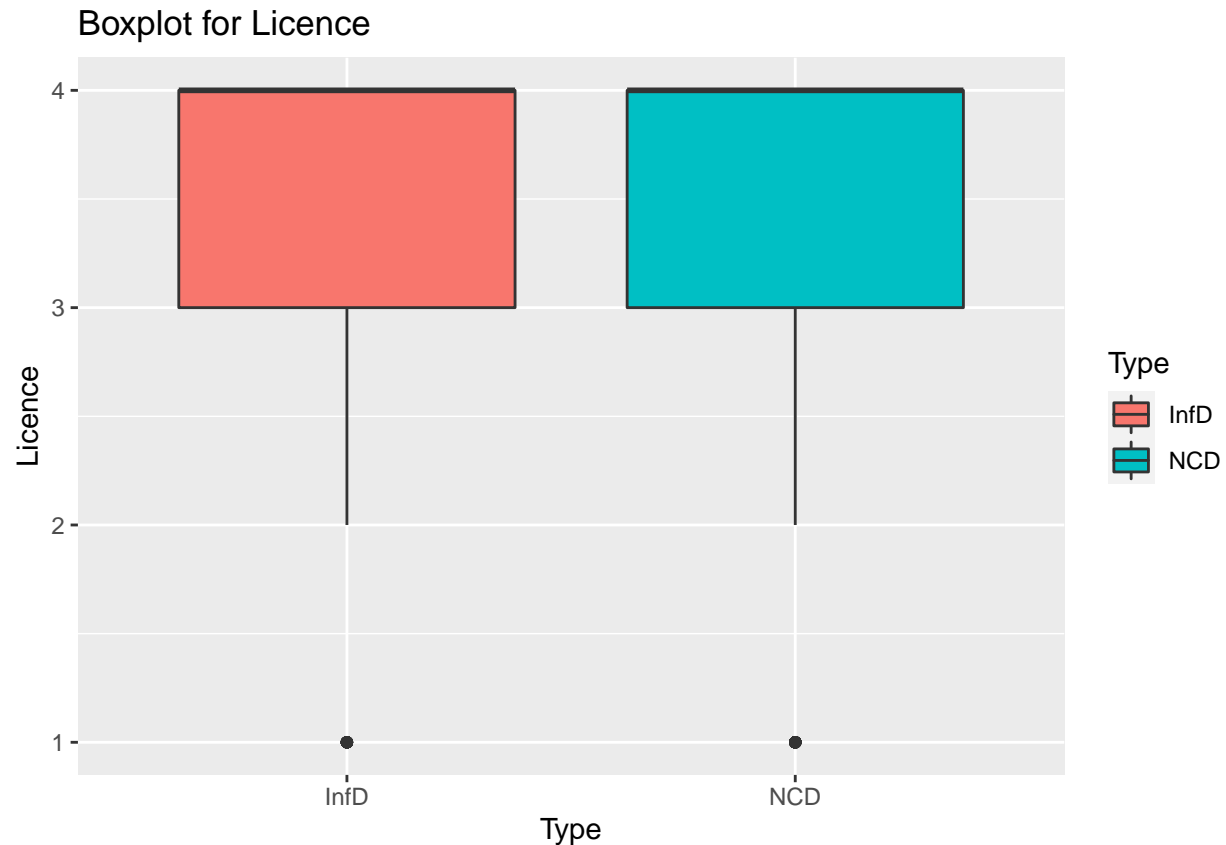
```
## Median of Complete by Type:
##   Type Complete
## 1 InfD         2
## 2 NCD          2
```

```
## Median of Reuse by Type:
##   Type Reuse
## 1 InfD     3
## 2 NCD     3
```



```
## Median of Access by Type:
##   Type Access
## 1 InfD   3.5
## 2 NCD   4.0
```



```
## Median of Licence by Type:
```

```
##   Type Licence
```

```
## 1 InfD      4
```

```
## 2 NCD       4
```

```
# Function to perform Mann-Whitney U test
perform_mann_whitney <- function(data, criterion) {
  test_result <- wilcox.test(
    reformulate("Type", response = criterion),
    data = data,
    exact = FALSE
  )
  list(criterion = criterion, p.value = test_result$p.value)
}

# Perform the test for each criterion
mw_results <- lapply(c("Complete", "Reuse", "Access", "Licence"), function(criterion) {
  perform_mann_whitney(data, criterion)
})

# Convert the list of results to a dataframe
mw_results_df <- do.call(rbind, mw_results)

print(mw_results_df)
```

```
##      criterion  p.value
```

```
## [1,] "Complete" 0.5901688
## [2,] "Reuse"    0.4855373
## [3,] "Access"   0.7664346
## [4,] "Licence"  0.9162592
```

```
#The assumptions of the test were met:
### Ordinal Data Check: The scoring criteria are ordinal variables
### Similar Distribution Shapes: For each criterion, it creates a box plot to visually inspect the dist.
### Independence of Observations:the variables are independence observations.

#these results were used in the First result Paragraph of the publication
```

```
#Creat variables for FAIR principles in 2016 and COVID-19 in 2020
```

```
data <- data %>%
  mutate(Period2020 = ifelse(Year <= 2020, "Before 2020", "After 2020")) %>%
  mutate(Period2016 = ifelse(Year <= 2016, "Before 2016", "After 2016"))#
```

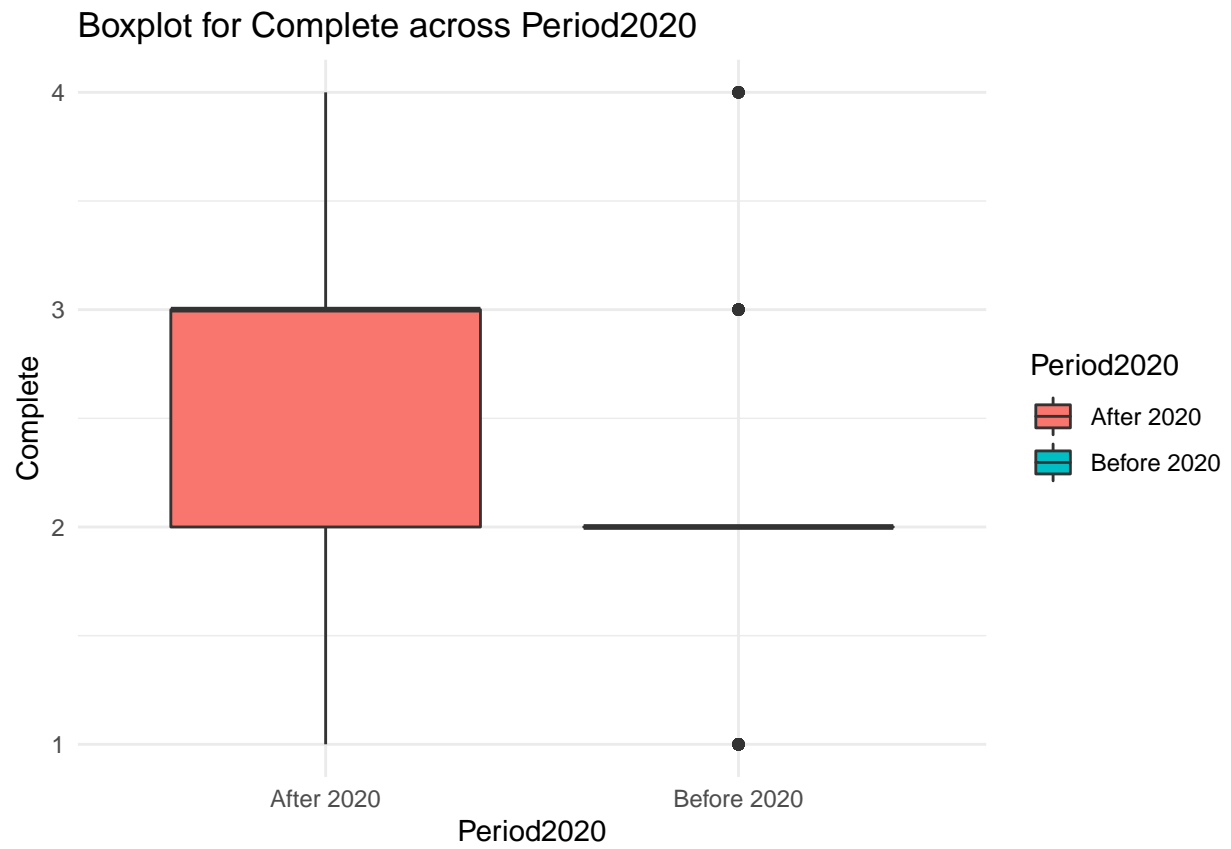
```
#Study the significant difference before and afterCovid-19 2020 using Median and Mann Whitney U test
```

```
# Ensure the scoring criteria are numeric
data <- data %>%
  mutate(across(c(Complete, Reuse, Access, Licence), ~as.numeric(as.character(.))))

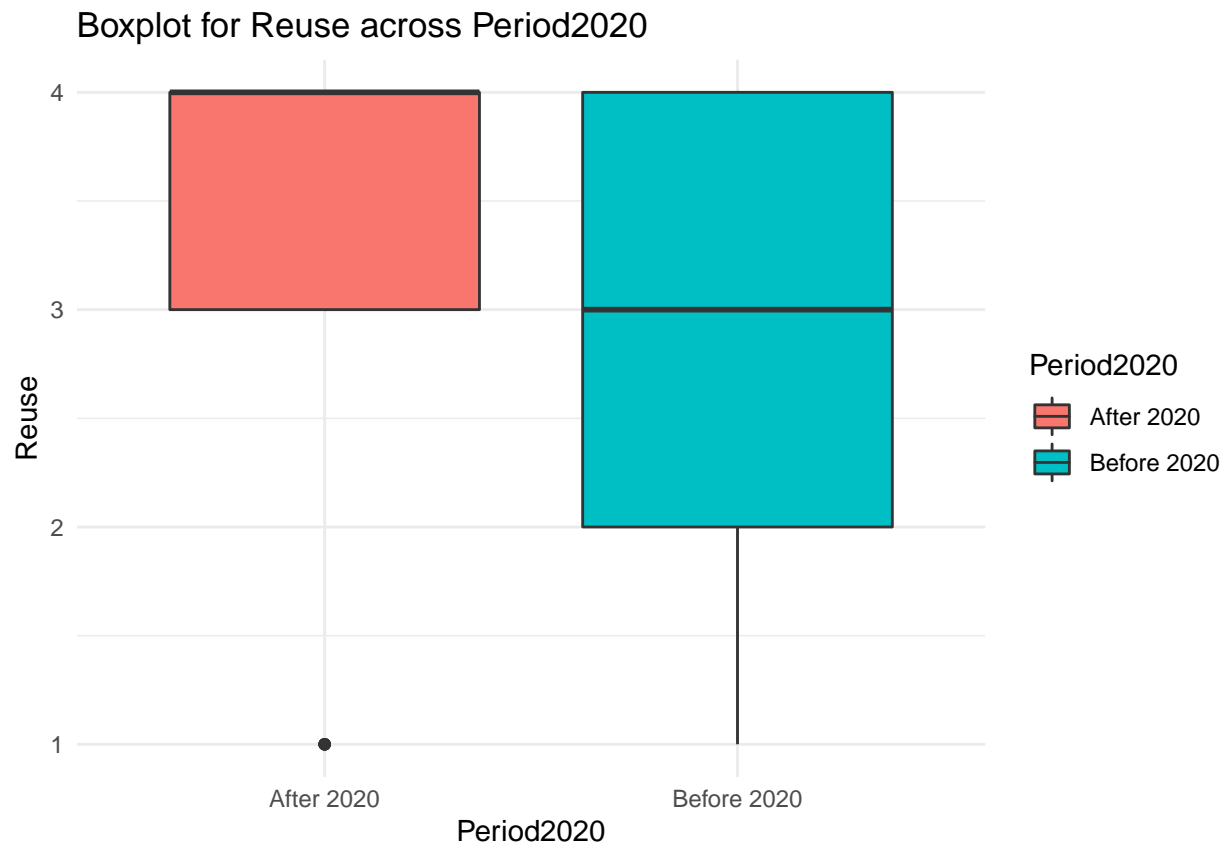
# Function to generate box plots for distribution checks
generate_boxplot <- function(data, score_var, period_var) {
  ggplot(data, aes_string(x = period_var, y = score_var, fill = period_var)) +
    geom_boxplot() +
    labs(title = paste("Boxplot for", score_var, "across", period_var),
         x = period_var,
         y = score_var) +
    theme_minimal()
}

# Generate and display box plots for each score with Period2020
lapply(c("Complete", "Reuse", "Access", "Licence"), function(score_var) {
  generate_boxplot(data, score_var, "Period2020")
})
```

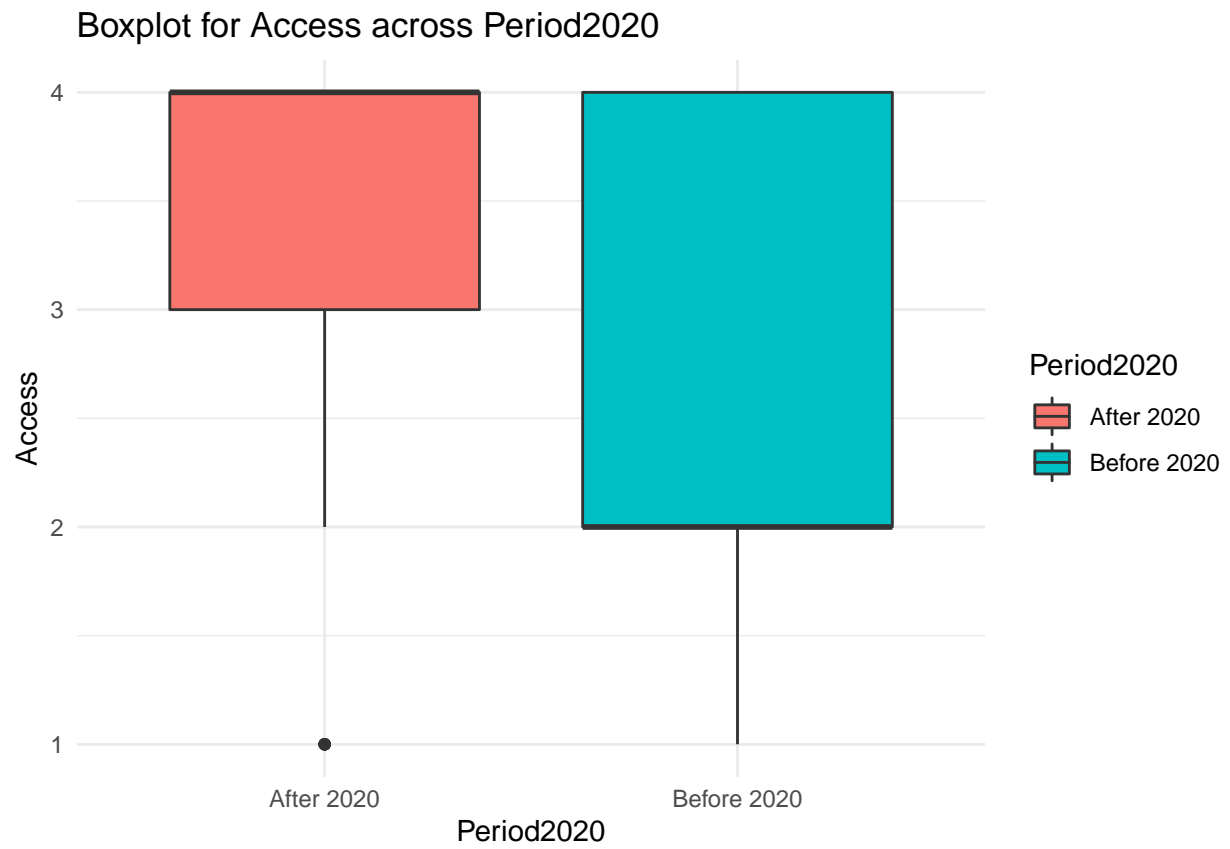
```
## [[1]]
```



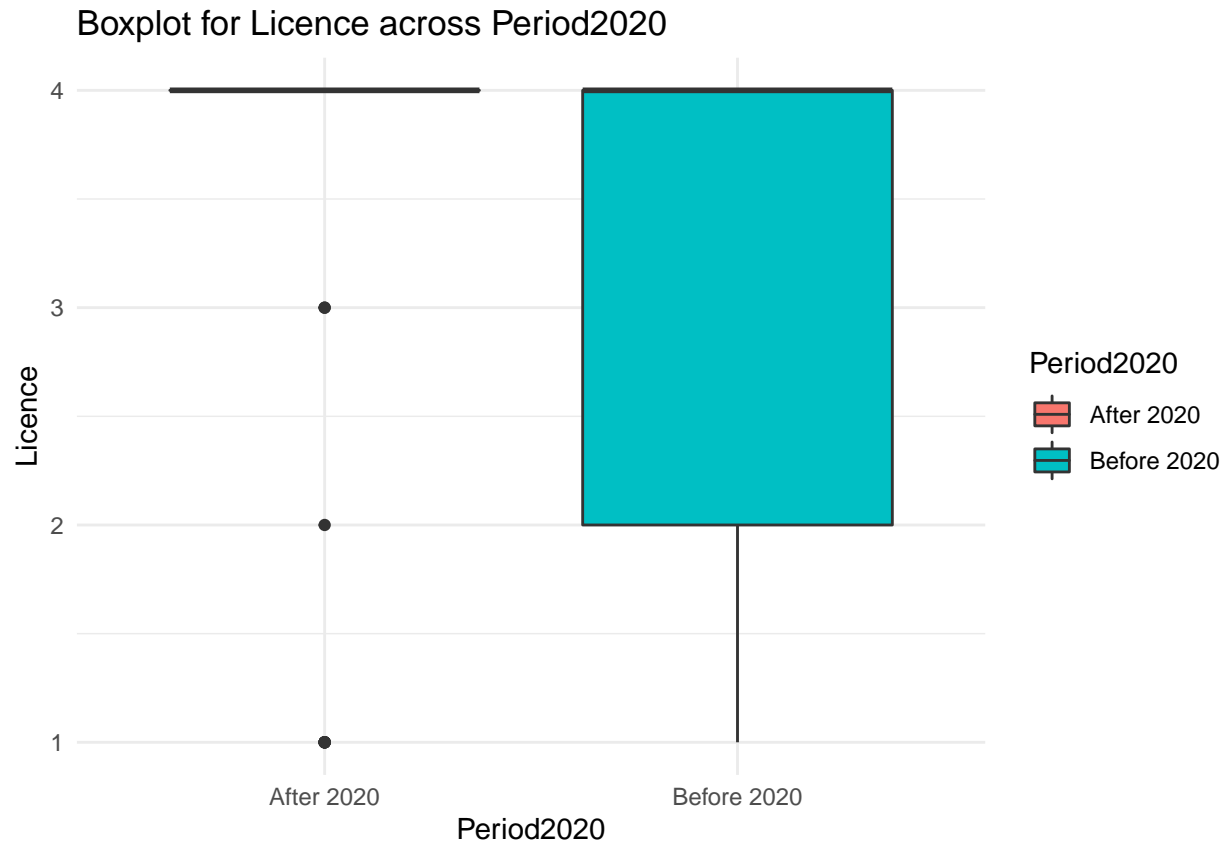
```
##  
## [[2]]
```



```
##  
## [[3]]
```



```
##  
## [[4]]
```



```
# Function to perform the Mann-Whitney test and calculate medians
perform_analysis <- function(data, score_var, period_var) {
  test_result <- wilcox.test(reformulate(period_var, score_var), data = data, exact = FALSE)

  medians <- data %>%
    group_by(!sym(period_var)) %>%
    summarise(median = median(!sym(score_var), na.rm = TRUE), .groups = 'drop')

  list(median = medians, p.value = test_result$p.value)
}
```

```
# Analysis for each score for 2020
results_complete_2020 <- perform_analysis(data, "Complete", "Period2020")
results_reuse_2020 <- perform_analysis(data, "Reuse", "Period2020")
results_access_2020 <- perform_analysis(data, "Access", "Period2020")
results_licence_2020 <- perform_analysis(data, "Licence", "Period2020")
```

```
# Print results
print(results_complete_2020)
```

```
## $median
## # A tibble: 2 x 2
##   Period2020 median
##   <chr>         <dbl>
## 1 After 2020     3
## 2 Before 2020    2
```



```
##
## $p.value
## [1] 0.00147346
```

```
print(results_reuse_2020)
```

```
## $median
## # A tibble: 2 x 2
##   Period2020 median
##   <chr>         <dbl>
## 1 After 2020      4
## 2 Before 2020     3
##
## $p.value
## [1] 0.0002688369
```

```
print(results_access_2020)
```

```
## $median
## # A tibble: 2 x 2
##   Period2020 median
##   <chr>         <dbl>
## 1 After 2020      4
## 2 Before 2020     2
##
## $p.value
## [1] 0.0002703917
```

```
print(results_licence_2020)
```

```
## $median
## # A tibble: 2 x 2
##   Period2020 median
##   <chr>         <dbl>
## 1 After 2020      4
## 2 Before 2020     4
##
## $p.value
## [1] 0.02901313
```

```
#The assumptions of the test were met:
```

```
### Ordinal Data Check: The scoring criteria are ordinal variables
```

```
### Similar Distribution Shapes: For each criterion, it creates a box plot to visually inspect the dist.
```

```
### Independence of Observations:the variables are independence observations.
```

```
#these results were used in the sixth result Paragraph of the publication
```

```
#Study the significant difference before and afterCovid-19 2020 using Median and Mann Whitney U test for
the NCD Papers
```

```

data_NCD <- data %>%
  filter(Type == "NCD")

perform_analysis <- function(data_NCD, score_var, period_var) {
  test_result <- wilcox.test(reformulate(period_var, score_var), data = data_NCD, exact = FALSE)

  medians <- data_NCD %>%
    group_by(!sym(period_var)) %>%
    summarise(median = median(!sym(score_var), na.rm = TRUE), .groups = 'drop')

  list(median = medians, p.value = test_result$p.value)
}

# Analysis for each score for 2020
results_complete_2020_NCD <- perform_analysis(data_NCD, "Complete", "Period2020")
results_reuse_2020_NCD <- perform_analysis(data_NCD, "Reuse", "Period2020")
results_access_2020_NCD <- perform_analysis(data_NCD, "Access", "Period2020")
results_licence_2020_NCD <- perform_analysis(data_NCD, "Licence", "Period2020")

# Print results
results_complete_2020_NCD

```

```

## $median
## # A tibble: 2 x 2
##   Period2020 median
##   <chr>         <dbl>
## 1 After 2020      2
## 2 Before 2020     2
##
## $p.value
## [1] 0.004012932

```

```
results_reuse_2020_NCD
```

```

## $median
## # A tibble: 2 x 2
##   Period2020 median
##   <chr>         <dbl>
## 1 After 2020      4
## 2 Before 2020     3
##
## $p.value
## [1] 0.001191915

```

```
results_access_2020_NCD
```

```

## $median
## # A tibble: 2 x 2
##   Period2020 median
##   <chr>         <dbl>
## 1 After 2020      4

```

```
## 2 Before 2020      2
##
## $p.value
## [1] 0.0007444155
```

```
results_licence_2020_NCD
```

```
## $median
## # A tibble: 2 x 2
##   Period2020 median
##   <chr>      <dbl>
## 1 After 2020      4
## 2 Before 2020      4
##
## $p.value
## [1] 0.0153904
```

#these results were used in the sixth result Paragraph of the publication

#Study the significant difference before and afterCovid-19 2020 using Median and Mann Whitney U test for the InfD Papers

```
data_InfD <- data %>%
  filter(Type == "InfD")

perform_analysis <- function(data_NCD, score_var, period_var) {
  test_result <- wilcox.test(reformulate(period_var, score_var), data = data_InfD, exact = FALSE)

  medians <- data_InfD %>%
    group_by(!sym(period_var)) %>%
    summarise(median = median(!sym(score_var), na.rm = TRUE), .groups = 'drop')

  list(median = medians, p.value = test_result$p.value)
}

# Analysis for each score for 2020
results_complete_2020_InfD <- perform_analysis(data_InfD, "Complete", "Period2020")
results_reuse_2020_InfD <- perform_analysis(data_InfD, "Reuse", "Period2020")
results_access_2020_InfD <- perform_analysis(data_InfD, "Access", "Period2020")
results_licence_2020_InfD <- perform_analysis(data_InfD, "Licence", "Period2020")

# Print results
results_complete_2020_InfD

## $median
## # A tibble: 2 x 2
##   Period2020 median
##   <chr>      <dbl>
## 1 After 2020      3
## 2 Before 2020      2
##
## $p.value
## [1] 0.07239354
```

```
results_reuse_2020_InfD
```

```
## $median
## # A tibble: 2 x 2
##   Period2020 median
##   <chr>         <dbl>
## 1 After 2020      4
## 2 Before 2020     3
##
## $p.value
## [1] 0.1105719
```

```
results_access_2020_InfD
```

```
## $median
## # A tibble: 2 x 2
##   Period2020 median
##   <chr>         <dbl>
## 1 After 2020      4
## 2 Before 2020     2
##
## $p.value
## [1] 0.1227605
```

```
results_licence_2020_InfD
```

```
## $median
## # A tibble: 2 x 2
##   Period2020 median
##   <chr>         <dbl>
## 1 After 2020      4
## 2 Before 2020     4
##
## $p.value
## [1] 0.6975206
```

```
#these results were used in the sixth result Paragraph of the publication
```

```
#Study the significant difference before and after Fair 2016 using Median and Mann Whitney U for all the papers
```

```
data <- data %>%
  mutate(Period2016 = ifelse(Year <= 2016, "Before 2016", "After 2016"))

# Function to perform the Mann-Whitney test, calculate medians, and generate box plots
perform_analysis <- function(data, score_var, period_var) {
  test_result <- wilcox.test(reformulate(period_var, score_var), data = data, exact = FALSE)

  medians <- data %>%
    group_by(!!sym(period_var)) %>%
    summarise(median = median(!!sym(score_var), na.rm = TRUE), .groups = 'drop')
```

```

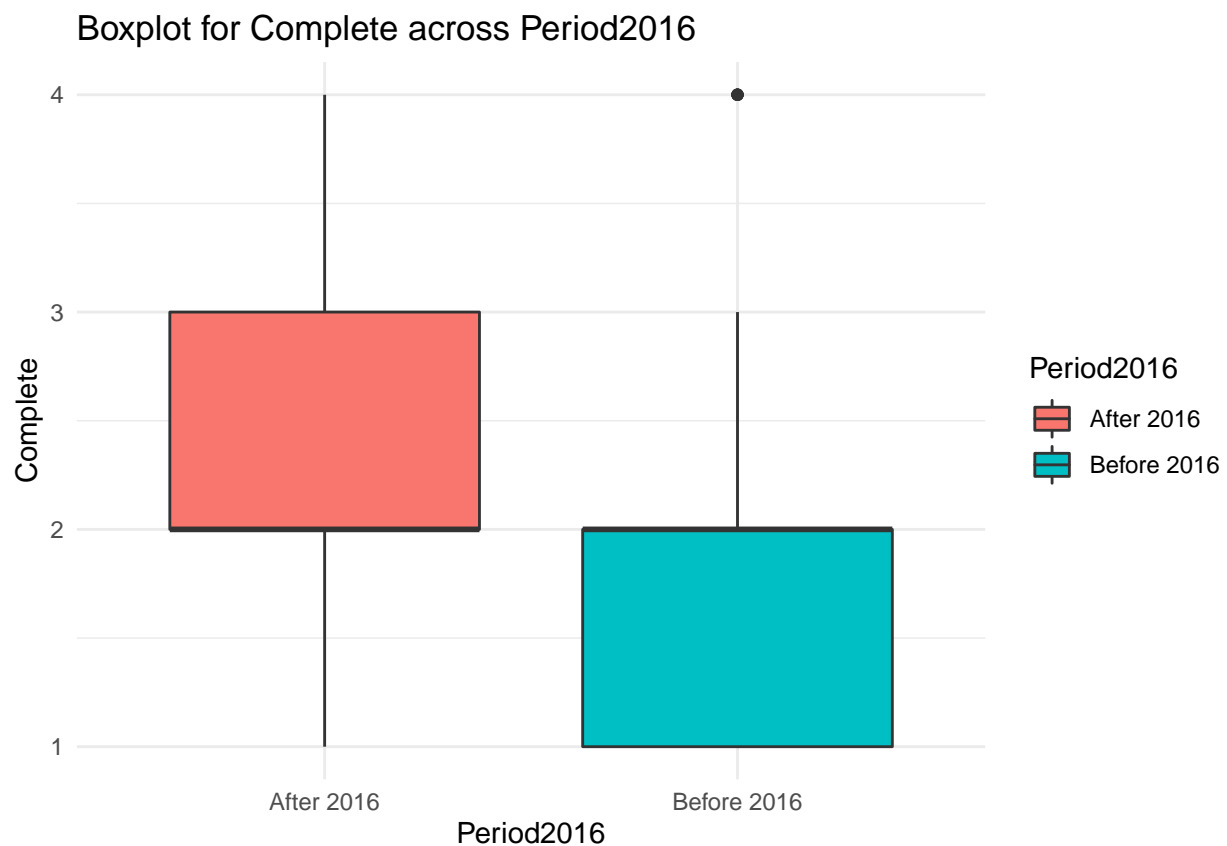
# Generate a box plot
box_plot <- ggplot(data, aes_string(x = period_var, y = score_var, fill = period_var)) +
  geom_boxplot() +
  labs(title = paste("Boxplot for", score_var, "across", period_var),
       x = period_var,
       y = score_var) +
  theme_minimal()

# Print the box plot
print(box_plot)

list(median = medians, p.value = test_result$p.value, plot = box_plot)
}

# Analysis for each score for 2016
results_complete_2016 <- perform_analysis(data, "Complete", "Period2016")

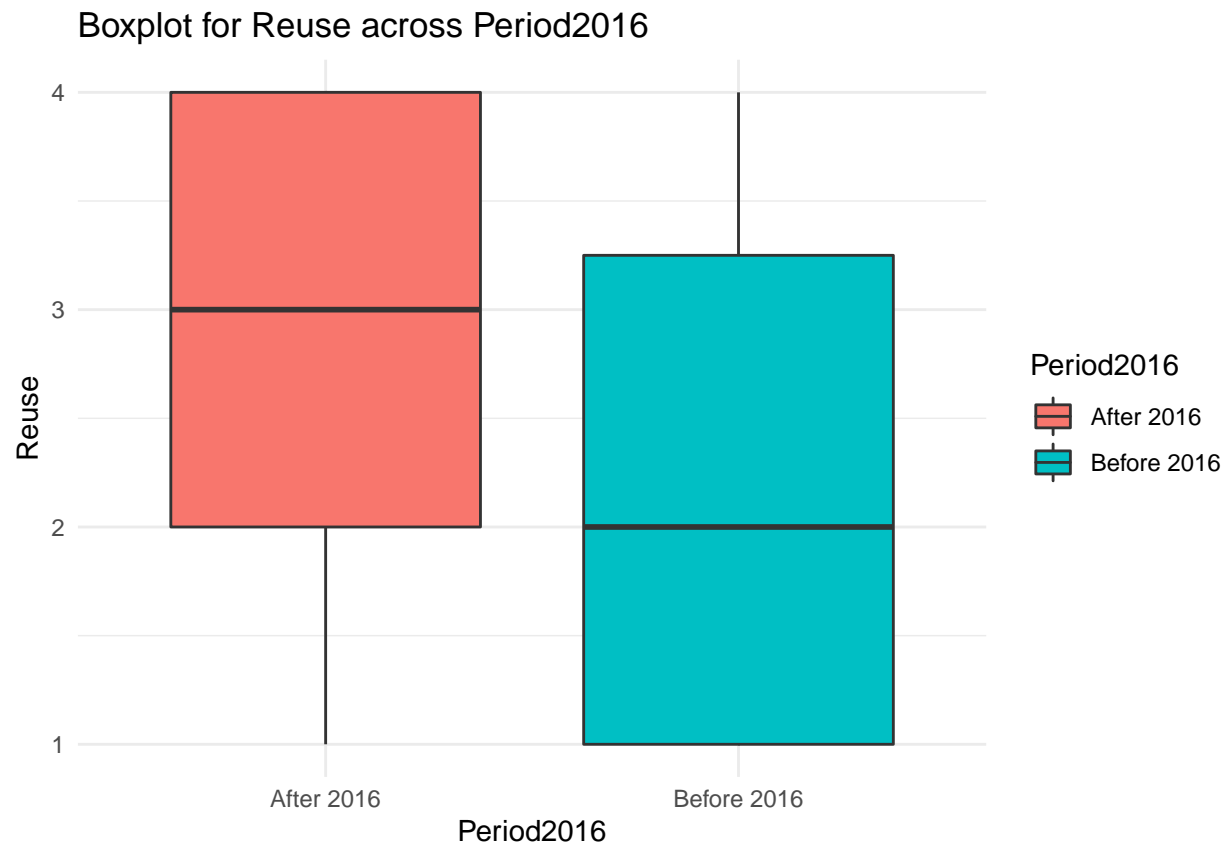
```



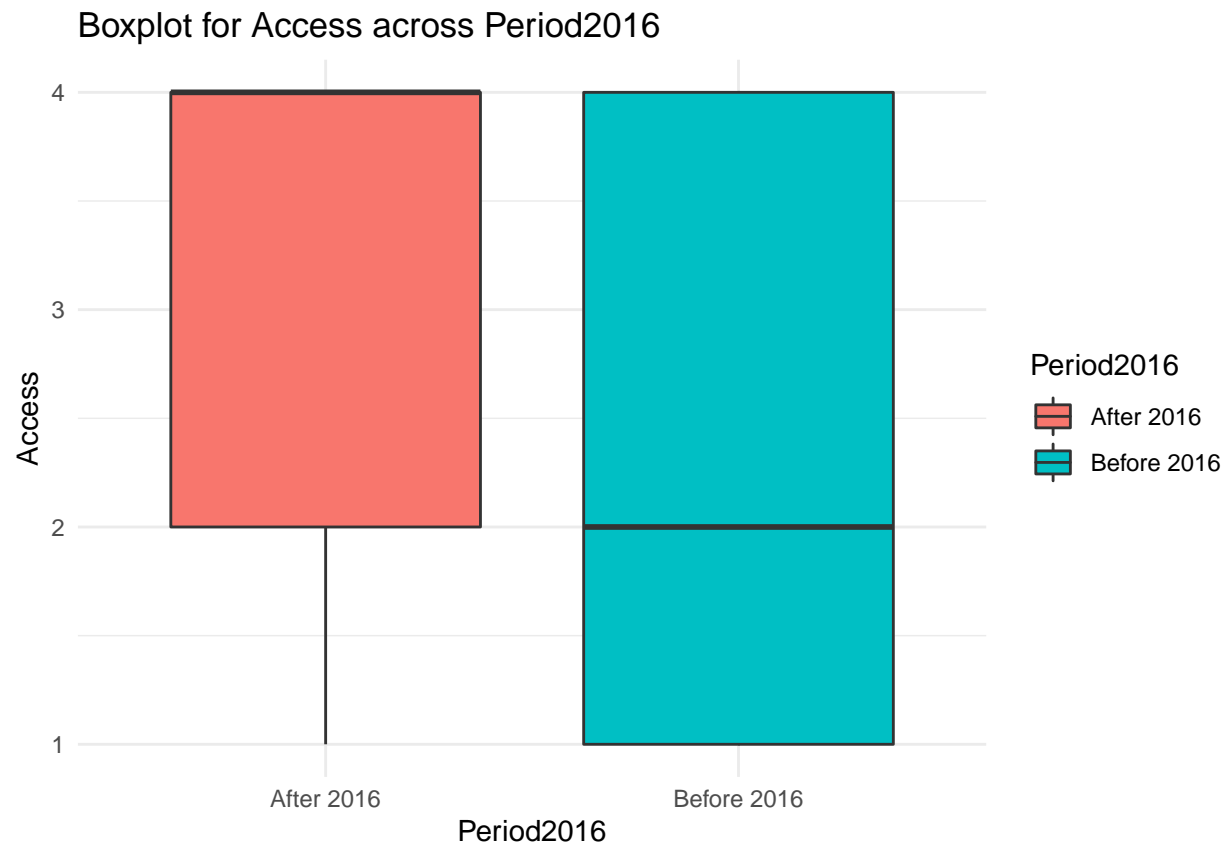
```

results_reuse_2016 <- perform_analysis(data, "Reuse", "Period2016")

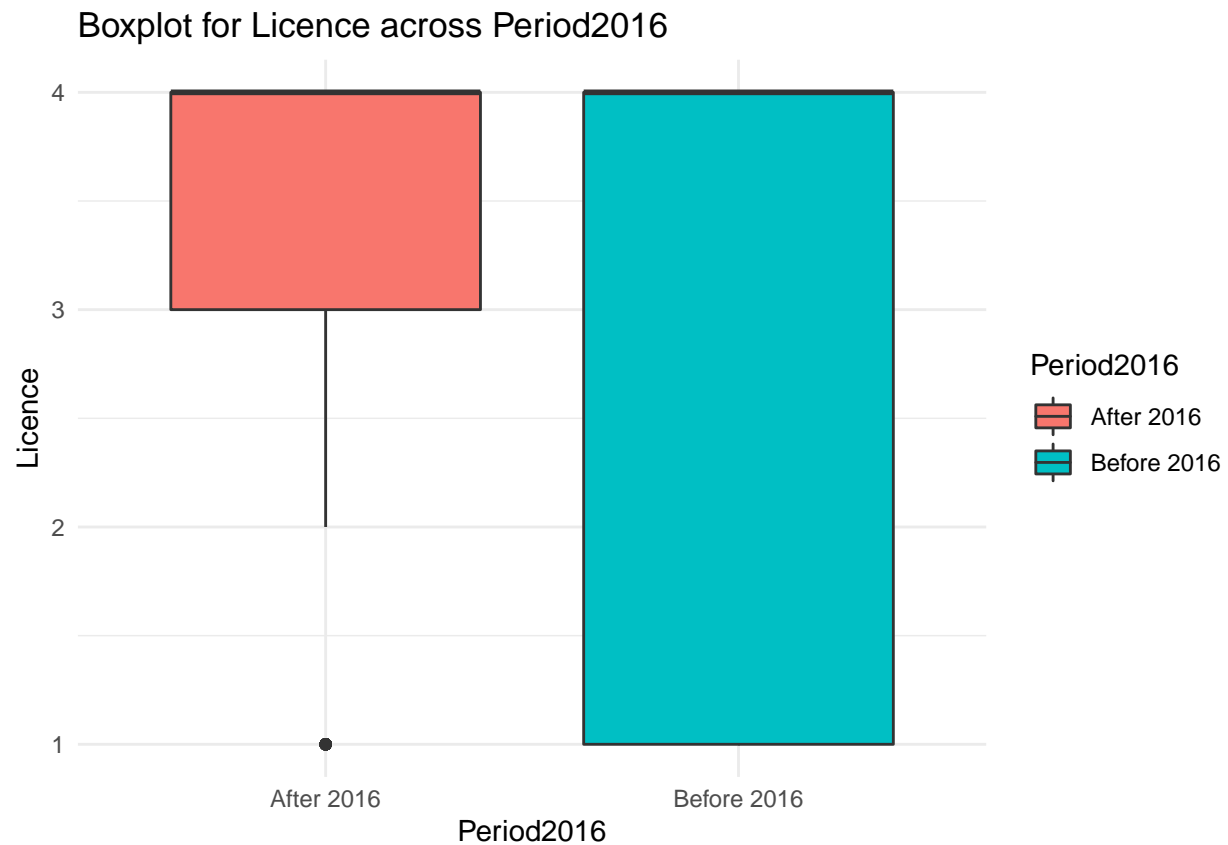
```



```
results_access_2016 <- perform_analysis(data, "Access", "Period2016")
```

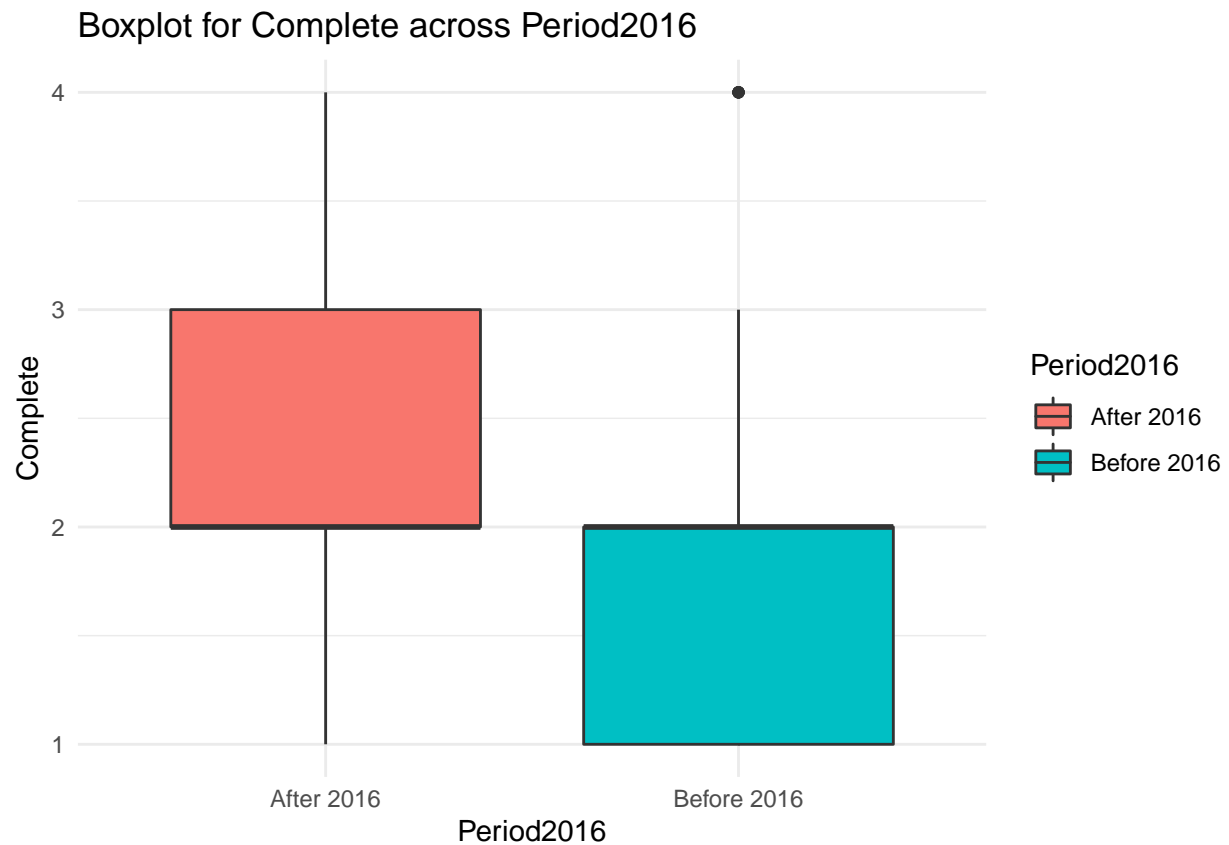


```
results_licence_2016 <- perform_analysis(data, "Licence", "Period2016")
```



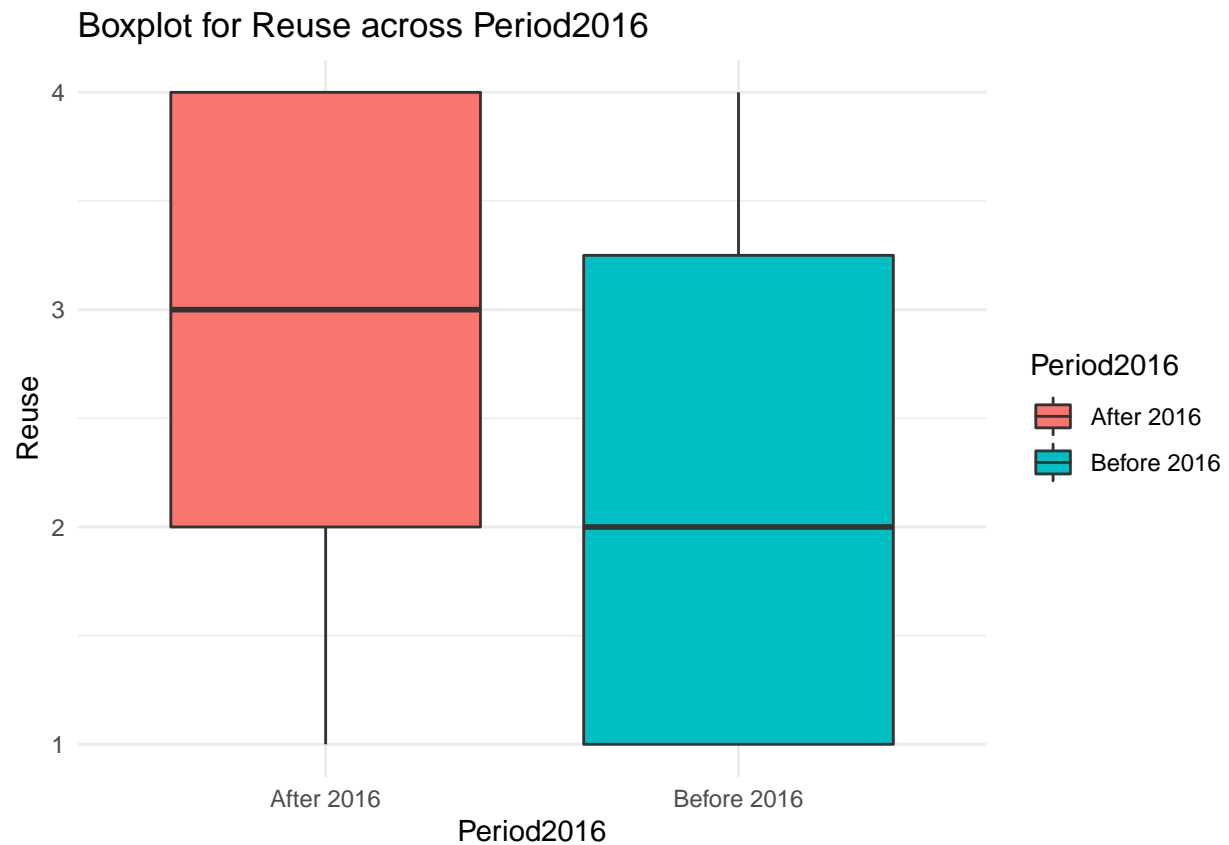
```
# Print results
results_complete_2016
```

```
## $median
## # A tibble: 2 x 2
##   Period2016 median
##   <chr>      <dbl>
## 1 After 2016      2
## 2 Before 2016      2
##
## $p.value
## [1] 0.05617637
##
## $plot
```

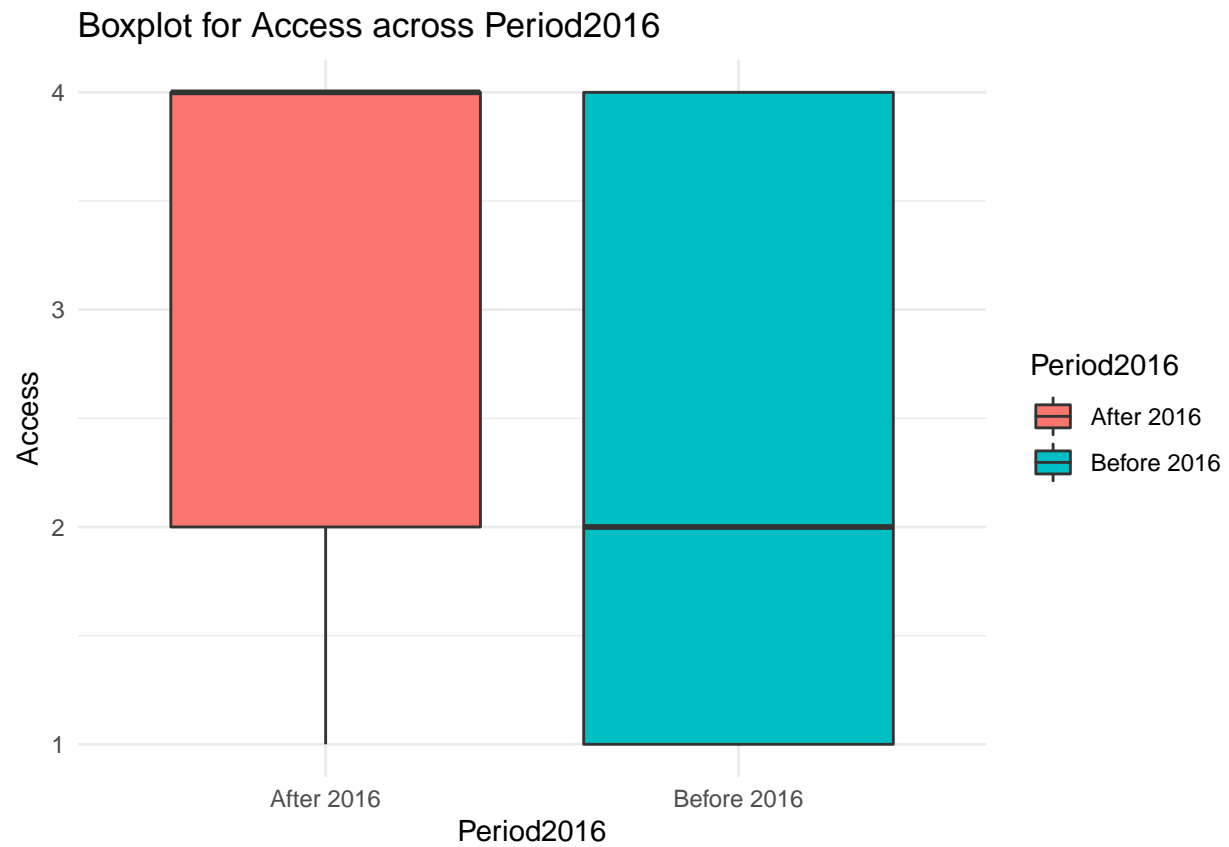
```
results_reuse_2016
```

```
## $median
## # A tibble: 2 x 2
##   Period2016 median
##   <chr>      <dbl>
## 1 After 2016      3
## 2 Before 2016     2
##
## $p.value
## [1] 0.002078694
##
## $plot
```



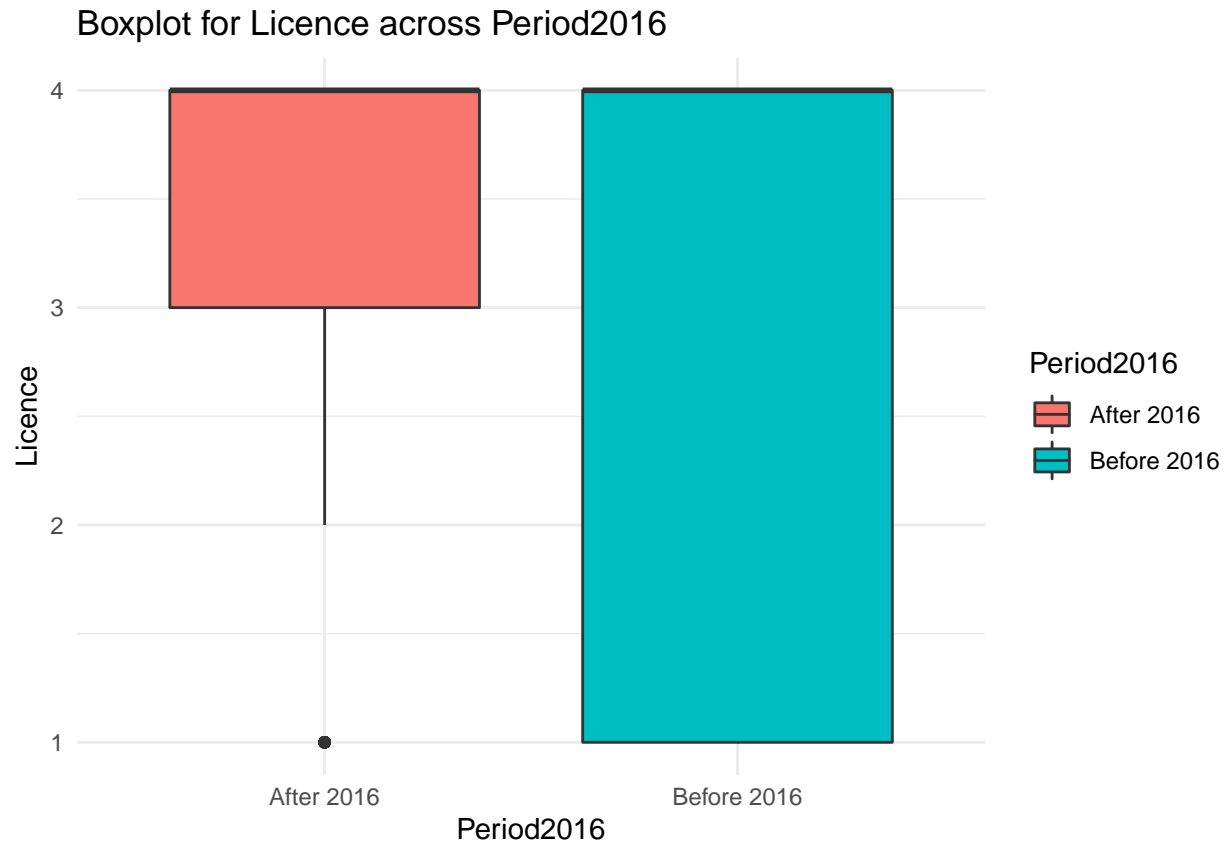
```
results_access_2016
```

```
## $median
## # A tibble: 2 x 2
##   Period2016 median
##   <chr>      <dbl>
## 1 After 2016      4
## 2 Before 2016     2
##
## $p.value
## [1] 0.03765665
##
## $plot
```



```
results_licence_2016
```

```
## $median
## # A tibble: 2 x 2
##   Period2016 median
##   <chr>         <dbl>
## 1 After 2016     4
## 2 Before 2016    4
##
## $p.value
## [1] 0.08882603
##
## $plot
```



#these results were used in the sixth result Paragraph of the publication

#Study the significant difference before and after Fair 2016 using Median and Mann Whitney U test for the NCD Papers

```
perform_analysis <- function(data_NCD, score_var, period_var) {
  test_result <- wilcox.test(reformulate(period_var, score_var), data = data_NCD, exact = FALSE)

  medians <- data_NCD %>%
    group_by(!sym(period_var)) %>%
    summarise(median = median(!sym(score_var), na.rm = TRUE), .groups = 'drop')

  list(median = medians, p.value = test_result$p.value)
}

# Analysis for each score for 2016
results_complete_2016_NCD <- perform_analysis(data_NCD, "Complete", "Period2016")
results_reuse_2016_NCD <- perform_analysis(data_NCD, "Reuse", "Period2016")
results_access_2016_NCD <- perform_analysis(data_NCD, "Access", "Period2016")
results_licence_2016_NCD <- perform_analysis(data_NCD, "Licence", "Period2016")

# Print results
results_complete_2016_NCD
```

```
## $median
## # A tibble: 2 x 2
```

```
##   Period2016  median
##   <chr>         <dbl>
## 1 After 2016      2
## 2 Before 2016     2
##
## $p.value
## [1] 0.09162029
```

```
results_reuse_2016_NCD
```

```
## $median
## # A tibble: 2 x 2
##   Period2016  median
##   <chr>         <dbl>
## 1 After 2016      3
## 2 Before 2016     3
##
## $p.value
## [1] 0.03941278
```

```
results_access_2016_NCD
```

```
## $median
## # A tibble: 2 x 2
##   Period2016  median
##   <chr>         <dbl>
## 1 After 2016      4
## 2 Before 2016     2
##
## $p.value
## [1] 0.1542596
```

```
results_licence_2016_NCD
```

```
## $median
## # A tibble: 2 x 2
##   Period2016  median
##   <chr>         <dbl>
## 1 After 2016      4
## 2 Before 2016     4
##
## $p.value
## [1] 0.15619
```

```
#these results were used in the sixth result Paragraph of the publication
```

```
#Study the significant difference before and after Fair 2016 using Median and Mann Whitney U test for the InfD Papers
```

```
# Analysis for each score for 2016
results_complete_2016_InfD <- perform_analysis(data_InfD, "Complete", "Period2016")
results_reuse_2016_InfD <- perform_analysis(data_InfD, "Reuse", "Period2016")
results_access_2016_InfD <- perform_analysis(data_InfD, "Access", "Period2016")
results_licence_2016_InfD <- perform_analysis(data_InfD, "Licence", "Period2016")
```

```
# Print results
```

```
results_complete_2016_InfD
```

```
## $median
## # A tibble: 2 x 2
##   Period2016 median
##   <chr>         <dbl>
## 1 After 2016      2
## 2 Before 2016    2
##
## $p.value
## [1] 0.2801454
```

```
results_reuse_2016_InfD
```

```
## $median
## # A tibble: 2 x 2
##   Period2016 median
##   <chr>         <dbl>
## 1 After 2016      3
## 2 Before 2016    2
##
## $p.value
## [1] 0.02973854
```

```
results_access_2016_InfD
```

```
## $median
## # A tibble: 2 x 2
##   Period2016 median
##   <chr>         <dbl>
## 1 After 2016      4
## 2 Before 2016    2
##
## $p.value
## [1] 0.147289
```

```
results_licence_2016_InfD
```

```
## $median
## # A tibble: 2 x 2
##   Period2016 median
##   <chr>         <dbl>
## 1 After 2016      4
## 2 Before 2016    4
```

```
##
## $p.value
## [1] 0.3305261
```

```
#these results were used in the sixth result Paragraph of the publication
```

```
#plot the distribution of DAS for each completeness score
```

```
# Transform the DAS variable
data <- data %>%
  mutate(NewDAS = case_when(
    is.na(DAS) ~ "Not Presented",
    DAS == 1 ~ "Shared",
    DAS == 0 ~ "Not Shared",
    TRUE ~ as.character(DAS))) # This line is just a fallback to handle unexpected values

class(data$NewDAS)
```

```
## [1] "character"
```

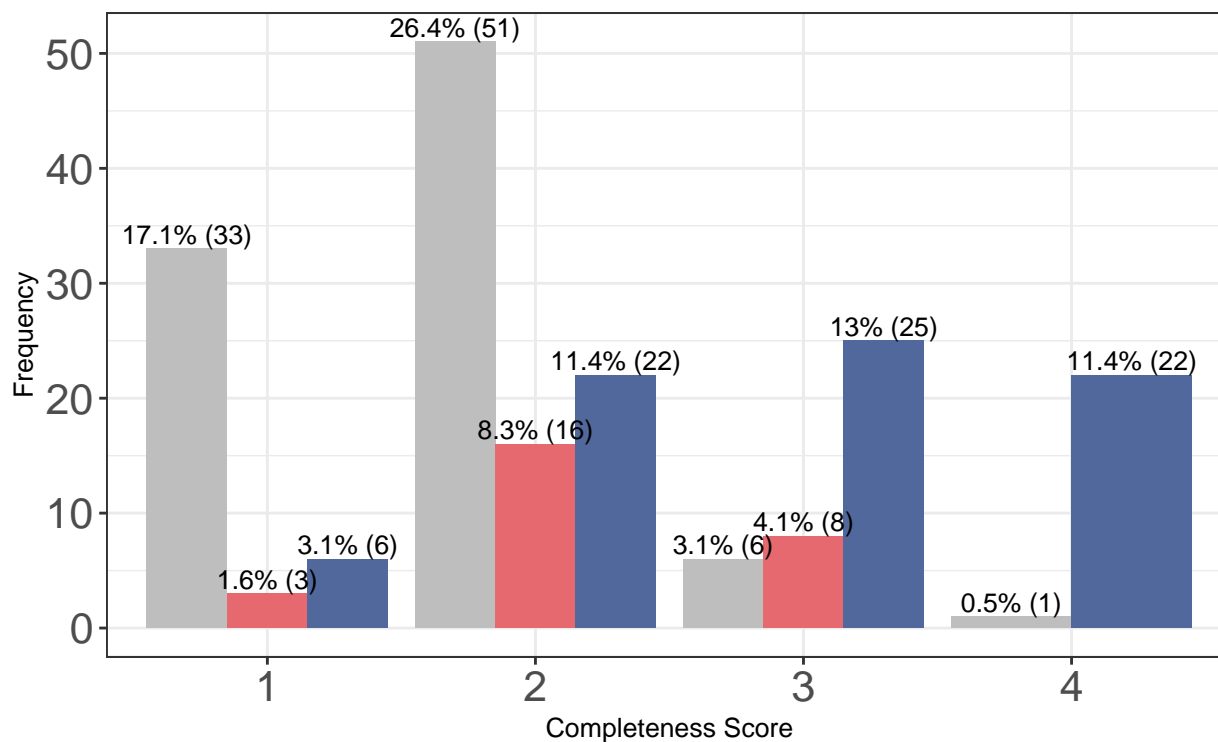
```
# Convert the new DAS variable to a factor for plotting
data$NewDAS <- factor(data$NewDAS, levels = c("Not Presented", "Not Shared", "Shared"))

# Calculate frequency and percentage for each combination of 'Complete' score and 'NewDAS' status
das_frequency <- data %>%
  group_by(Complete, NewDAS) %>%
  summarise(Frequency = n(), .groups = 'drop') %>%
  mutate(Percentage = (Frequency / sum(Frequency)) * 100)

# Define specific colors for the new DAS values
colors <- c("Not Presented" = "gray", "Not Shared" = "#E5696F", "Shared" = "#50689B")

# Create the plot
Figure_4 <- ggplot(das_frequency, aes(x = as.factor(Complete), y = Frequency, fill = NewDAS)) +
  geom_bar(stat = "identity", position = "dodge") +
  geom_text(aes(label = paste0(round(Percentage, 1), "% (", Frequency, ")")),
            position = position_dodge(width = 0.9), vjust = -0.25, size = 3.5) +
  scale_fill_manual(values = colors, name = "DAS Status") +
  theme_bw() + # Use theme_bw for a white background
  theme(
    axis.title = element_text(size = 10), # Increase axis titles
    axis.text = element_text(size = 16), # Increase axis text
    legend.title = element_text(size = 14), # Increase legend title
    legend.text = element_text(size = 16) # Increase legend text
  ) +
  labs(x = "Completeness Score", y = "Frequency") +
  theme(legend.position = "bottom")

# Print and save the plot
print(Figure_4)
```



DAS Status Not Presented Not Shared Shared

```
ggsave("Figure_4.png", Figure_4, width = 8, height = 6, bg = "white")
```

#these results were used in the Figure 4 of the publication

```
# Calculate the total count per year
yearly_totals <- data %>%
  group_by(Year) %>%
  summarise(Total = n(), .groups = 'drop')
```

```
# Join the totals back to the original data and calculate proportions
proportion_data <- data %>%
  left_join(yearly_totals, by = "Year") %>%
  group_by(Year, NewDAS) %>%
  summarise(Count = n(), .groups = 'drop') %>%
  mutate(Proportion = Count / n())
```

```
# Get all unique years for the x-axis
all_years <- sort(unique(proportion_data$Year))
```

```
colors <- c("Not Presented" = "#828282", "Not Shared" = "#E5696F", "Shared" = "#50689B")
```

```
# Plot with every year on the x-axis
Figure_S2 <- ggplot(proportion_data, aes(x = Year, y = Proportion, color = NewDAS, group = NewDAS)) +
  geom_line() +
  geom_point() +
  geom_vline(xintercept = 2016, linetype = "dashed", color = "black") +
```

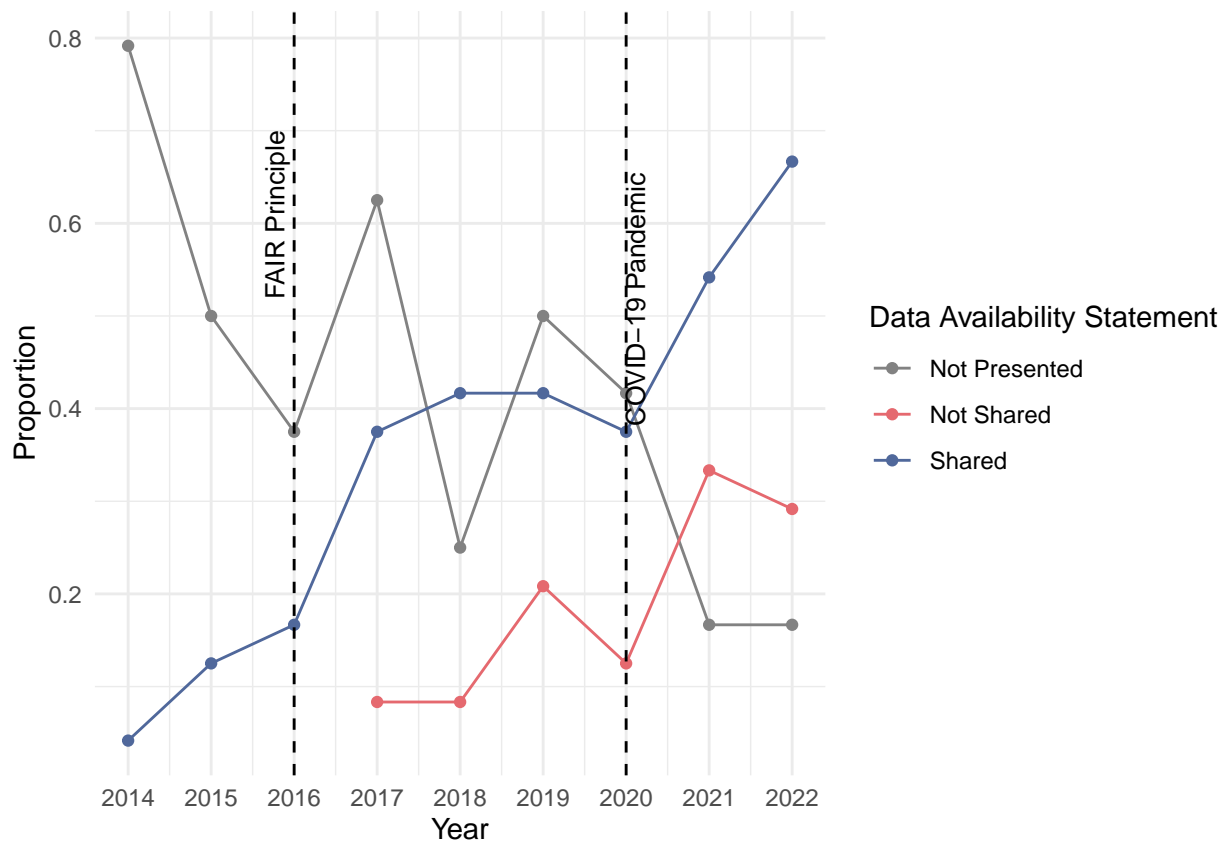


```

geom_vline(xintercept = 2020, linetype = "dashed", color = "black") +
annotate("text", x = 2016, y = max(proportion_data$Proportion), label = "FAIR Principle", angle = 90,
annotate("text", x = 2020, y = max(proportion_data$Proportion), label = "COVID-19 Pandemic", angle = 90) +
scale_x_continuous(breaks = all_years) +
labs(x = "Year", y = "Proportion", color = "Data Availability Statement") +
theme(
  axis.title = element_text(size = 10), # Increase axis titles
  axis.text = element_text(size = 16), # Increase axis text
  legend.title = element_text(size = 14), # Increase legend title
  legend.text = element_text(size = 16) # Increase legend text
) +
theme_minimal() +
scale_color_manual(values = colors)

print(Figure_S2)

```



```

ggsave("Figure_S2.png", Figure_S2, width = 8, height = 6, bg = "white")

```

#these results were used in the Figure S2 of the publication

```

#Convert the Preprint numeric values to factor levels 'Yes' and 'No'
data$Preprints <- factor(ifelse(data$Preprints == 1, "Yes", "No"))

```

```

# Calculate the total count per year and proportion for Preprint
proportion_data <- data %>%

```

```

group_by(Year) %>%
count(Preprints) %>%
mutate(Total = sum(n),
       Proportion = n / Total)

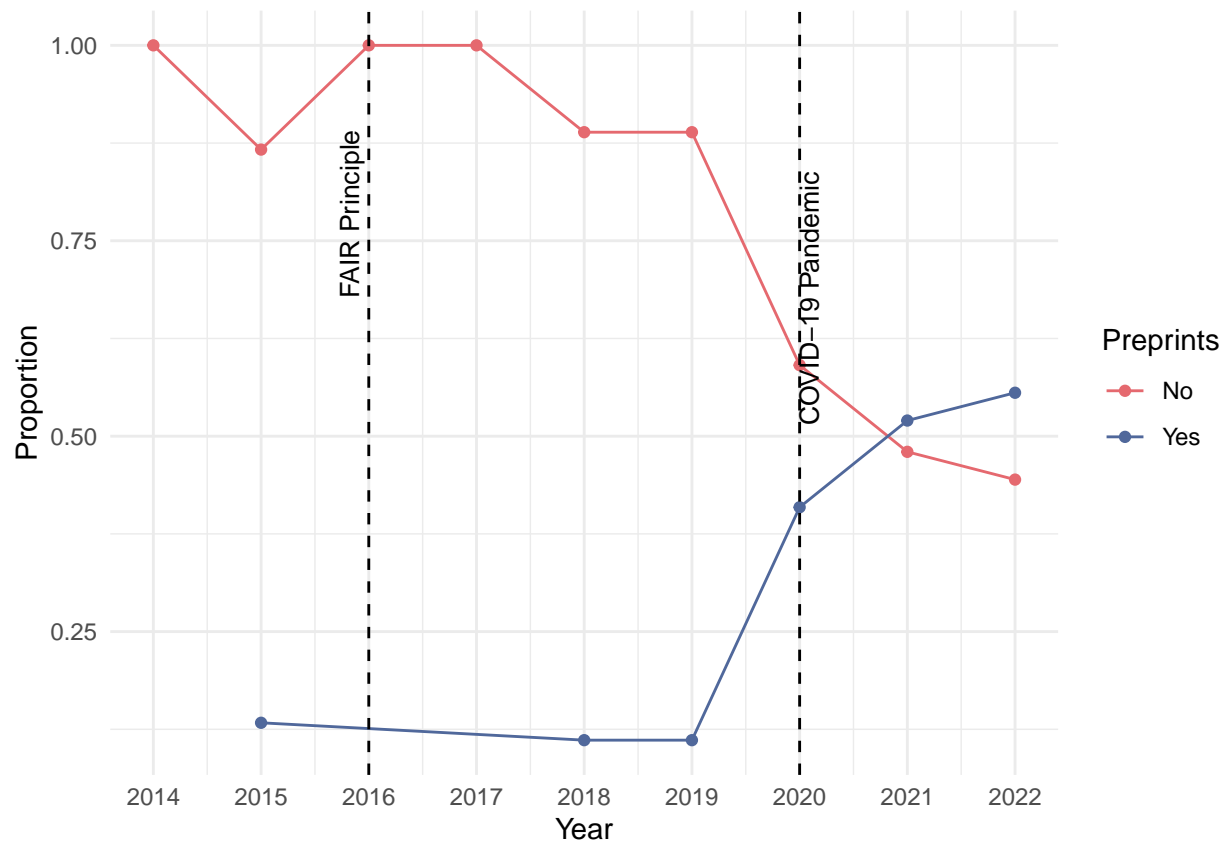
# Get all unique years for the x-axis
all_years <- sort(unique(proportion_data$Year))

# Define colors for 'Yes' and 'No'
colors <- c("No" = "#E5696F", "Yes" = "#50689B")

# Plot with every year on the x-axis for Preprints
Figure_S3 <- ggplot(proportion_data, aes(x = Year, y = Proportion, color = Preprints, group = Preprints)) +
  geom_line() +
  geom_point() +
  geom_vline(xintercept = 2016, linetype = "dashed", color = "black") +
  geom_vline(xintercept = 2020, linetype = "dashed", color = "black") +
  annotate("text", x = 2016, y = max(proportion_data$Proportion), label = "FAIR Principle", angle = 90,
  ) +
  annotate("text", x = 2020, y = max(proportion_data$Proportion), label = "COVID-19 Pandemic", angle = 90,
  ) +
  scale_x_continuous(breaks = all_years) +
  labs(x = "Year", y = "Proportion", color = "Preprints") +
  theme(
    axis.title = element_text(size = 10), # Increase axis titles
    axis.text = element_text(size = 16), # Increase axis text
    legend.title = element_text(size = 14), # Increase legend title
    legend.text = element_text(size = 16) # Increase legend text
  ) +
  theme_minimal() +
  scale_color_manual(values = colors)

print(Figure_S3)

```



```
# Save the plot for Preprint
ggsave("Figure_S3.png", Figure_S3, width = 8, height = 6, bg = "white")

#these results were used in the Figure S3 of the publication
```

#study if FAIR implementation and Covid19 have an effect on DAS and Preprint # 1- Fair implementataion

```
total_by_period <- data %>%
  group_by(Period2016) %>%
  summarise(Total = n(), .groups = "drop")

# Join this total with the DAS and Preprints summaries and calculate the percentage
Preprints_summary <- data %>%
  group_by(Period2016, Preprints) %>%
  summarise(Frequency = n(), .groups = "drop") %>%
  left_join(total_by_period, by = "Period2016") %>%
  mutate(Percentage = (Frequency / Total) * 100)

# Print the Preprints summary table
print(Preprints_summary)
```

```
## # A tibble: 4 x 5
##   Period2016 Preprints Frequency Total Percentage
##   <chr>      <fct>      <int> <int>     <dbl>
## 1 After 2016 No          103   145     71.0
```

```
## 2 After 2016 Yes          42   145    29.0
## 3 Before 2016 No         46    48    95.8
## 4 Before 2016 Yes        2    48    4.17
```

```
Preprints_table <- table(data$Preprints, data$Period2016)
```

```
# Perform the Chi-square test
```

```
result <- chisq.test(Preprints_table)
```

```
#Print the chi-square results
```

```
print(result)
```

```
##
```

```
## Pearson's Chi-squared test with Yates' continuity correction
```

```
##
```

```
## data: Preprints_table
```

```
## X-squared = 11.231, df = 1, p-value = 0.0008044
```

```
# Get the expected values
```

```
expected_values <- result$expected
```

```
# Print the expected values
```

```
print(expected_values) ##in order to check the assumption of chi square test: Each cell in the contingency table has an expected count of 5 or more.
```

```
##
```

```
##      After 2016 Before 2016
```

```
## No   111.94301  37.05699
```

```
## Yes   33.05699  10.94301
```

```
#The assumptions of that test were met:
```

```
#### Sample Size: Each cell in the contingency table has an expected count of 5 or more.
```

```
#### Independence: The observations are independent of each other. This means that the selection of one observation does not affect the selection of another.
```

```
### Random Sampling: The data is a random sample.
```

```
### Categorical Data: Both variables should be categorical (either nominal or ordinal).
```

```
#these results were used in the seventh result paragraph and Table S3/S4 of the publication
```

```
total_by_period <- data %>%
```

```
  group_by(Period2016) %>%
```

```
  summarise(Total = n(), .groups = "drop")
```

```
# Join this total with the NewDAS and Preprints summaries and calculate the percentage
```

```
das_summary <- data %>%
```

```
  group_by(Period2016, NewDAS) %>%
```

```
  summarise(Frequency = n(), .groups = "drop") %>%
```

```
  left_join(total_by_period, by = "Period2016") %>%
```

```
  mutate(Percentage = (Frequency / Total) * 100)
```

```
# Print the DAS summary table
```

```
print(das_summary)
```

```
## # A tibble: 5 x 5
##   Period2016 NewDAS      Frequency Total Percentage
##   <chr>      <fct>      <int> <int>      <dbl>
## 1 After 2016 Not Presented      51   145      35.2
## 2 After 2016 Not Shared       27   145      18.6
## 3 After 2016 Shared          67   145      46.2
## 4 Before 2016 Not Presented    40    48      83.3
## 5 Before 2016 Shared          8    48      16.7
```

```
das_table <- table(data$NewDAS, data$Period2016)
```

```
# Perform the Chi-square test
result <- chisq.test(das_table)
```

```
#Print the chi-square results
print(result)
```

```
##
## Pearson's Chi-squared test
##
## data:  das_table
## X-squared = 34.776, df = 2, p-value = 2.809e-08
```

```
# Get the expected values
expected_values <- result$expected
```

```
# Print the expected values
print(expected_values) ##in order to check the assumption of chi square test: Each cell in the contingency table has an expected count of 5 or more.
```

```
##
##           After 2016 Before 2016
## Not Presented  68.36788  22.632124
## Not Shared    20.28497   6.715026
## Shared        56.34715  18.652850
```

```
#The assumptions of that test were met:
```

```
#### Sample Size: Each cell in the contingency table has an expected count of 5 or more.
#### Independence: The observations are independent of each other. This means that the selection of one observation does not affect the selection of another.
### Random Sampling: The data is a random sample.
### Categorical Data: Both variables should be categorical (either nominal or ordinal).
```

```
#these results were used in the seventh result paragraph and Table S3/S4 of the publication
```

```
#2- Covid 19 # the impact of COVID-19 on Preprints
```

```
total_by_period <- data %>%
  group_by(Period2020)%>%
  summarise(Total = n(), .groups = "drop")
```

```
# Join this total with the DAS and Preprints summaries and calculate the percentage
Preprints_summary <- data %>%
```

```

group_by(Period2020, Preprints) %>%
summarise(Frequency = n(), .groups = "drop") %>%
left_join(total_by_period, by = "Period2020") %>%
mutate(Percentage = (Frequency / Total) * 100)

# Print the Preprints summary table
print(Preprints_summary)

```

```

## # A tibble: 4 x 5
##   Period2020 Preprints Frequency Total Percentage
##   <chr>      <fct>      <int> <int>      <dbl>
## 1 After 2020 No          24     52        46.2
## 2 After 2020 Yes         28     52        53.8
## 3 Before 2020 No        125    141        88.7
## 4 Before 2020 Yes         16    141        11.3

```

```
Preprints_table <- table(data$Preprints, data$Period2020)
```

```

# Perform the Chi-square test
result <- chisq.test(Preprints_table)

#Print the chi-square results
print(result)

```

```

##
## Pearson's Chi-squared test with Yates' continuity correction
##
## data: Preprints_table
## X-squared = 36.607, df = 1, p-value = 1.445e-09

```

```

# Get the expected values
expected_values <- result$expected

```

```

# Print the expected values
print(expected_values) ##in order to check the assumption of chi square test: Each cell in the contingency table has an expected count of 5 or more.

```

```

##
##      After 2020 Before 2020
## No    40.14508  108.85492
## Yes   11.85492   32.14508

```

```
#The assumptions of that test were met:
```

```

#### Sample Size: Each cell in the contingency table has an expected count of 5 or more.
#### Independence: The observations are independent of each other. This means that the selection of one observation does not affect the selection of another.
#### Random Sampling: The data is a random sample.
#### Categorical Data: Both variables should be categorical (either nominal or ordinal)

```

```
#these results were used in the seventh result paragraph of the publication
```

the impact of COVID-19 on DAS

```
total_by_period <- data %>%
  group_by(Period2020) %>%
  summarise(Total = n(), .groups = "drop")

# Join this total with the NewDAS and Preprints summaries and calculate the percentage
das_summary <- data %>%
  group_by(Period2020, NewDAS)%>%
  summarise(Frequency = n(), .groups = "drop") %>%
  left_join(total_by_period, by = "Period2020") %>%
  mutate(Percentage = (Frequency / Total) * 100)

# Print the DAS summary table
print(das_summary)
```

```
## # A tibble: 6 x 5
##   Period2020 NewDAS      Frequency Total Percentage
##   <chr>      <fct>      <int> <int>    <dbl>
## 1 After 2020 Not Presented      8    52    15.4
## 2 After 2020 Not Shared     15    52    28.8
## 3 After 2020 Shared       29    52    55.8
## 4 Before 2020 Not Presented  83   141    58.9
## 5 Before 2020 Not Shared    12   141     8.51
## 6 Before 2020 Shared      46   141    32.6
```

```
das_table <- table(data$NewDAS, data$Period2016)
```

```
# Perform the Chi-square test
result <- chisq.test(das_table)
```

```
#Print the chi-square results
print(result)
```

```
##
## Pearson's Chi-squared test
##
## data:  das_table
## X-squared = 34.776, df = 2, p-value = 2.809e-08
```

```
# Get the expected values
expected_values <- result$expected
```

```
# Print the expected values
print(expected_values) ##in order to check the assumption of chi square test: Each cell in the contingency
```

```
##
##           After 2016 Before 2016
## Not Presented  68.36788  22.632124
## Not Shared    20.28497   6.715026
## Shared        56.34715  18.652850
```

```
#The assumptions of that test were met:
```

```
#### Sample Size: Each cell in the contingency table has an expected count of 5 or more.
```

```
#### Independence: The observations are independent of each other. This means that the selection of one
```

```
#### Random Sampling: The data is a random sample.
```

```
#### Categorical Data: Both variables should be categorical (either nominal or ordinal)
```

```
#these results were used in the seventh result paragraph of the publication
```

```
#2 - Ordinal regression models (Scoring Criteria and year)
```

```
library(ordinal) # ordinal logistic regression: cumulative link mixed models, clm function is in this p
```

```
## Warning: package 'ordinal' was built under R version 4.2.3
```

```
##
```

```
## Attaching package: 'ordinal'
```

```
## The following object is masked from 'package:dplyr':
```

```
##
```

```
## slice
```

```
library(VGAM) # more ordinal regression
```

```
## Warning: package 'VGAM' was built under R version 4.2.3
```

```
## Loading required package: stats4
```

```
## Loading required package: splines
```

```
##
```

```
## Attaching package: 'VGAM'
```

```
## The following objects are masked from 'package:ordinal':
```

```
##
```

```
## dgumbel, dlgamma, pgumbel, plgamma, qgumbel, rgumbel, wine
```

```
library(dplyr) # Data manipulation
```

```
library(chisq.posthoc.test) # If needed
```

```
## Warning: package 'chisq.posthoc.test' was built under R version 4.2.3
```

```
library(gmodels) # For SPSS style chi-sq/ contingency tables
```

```
## Warning: package 'gmodels' was built under R version 4.2.3
```



```
library(emmeans)
```

```
## Warning: package 'emmeans' was built under R version 4.2.3
```

```
library(sure)
```

```
## Warning: package 'sure' was built under R version 4.2.3
```

```
data$Complete <- factor(data$Complete, ordered = TRUE)
data$Reuse <- factor(data$Reuse, ordered = TRUE)
data$Access <- factor(data$Access, ordered = TRUE)
data$Licence <- factor(data$Licence, ordered = TRUE)

levels(data$Complete)
```

```
## [1] "1" "2" "3" "4"
```

```
levels(data$Reuse)
```

```
## [1] "1" "2" "3" "4"
```

```
levels(data$Access)
```

```
## [1] "1" "2" "3" "4"
```

```
levels(data$Licence)
```

```
## [1] "1" "2" "3" "4"
```

```
#Ordinal regression model for the Completeness by year
```

```
m1a <- clm(Complete ~ Year, data = data)
```

```
## Warning in x$code == 0L || action == "silent": 'length(x) = 2 > 1' in coercion
## to 'logical(1)'
```

```
## Warning: (2) Model is nearly unidentifiable: very large eigenvalue
## - Rescale variables?
## In addition: Absolute and relative convergence criteria were met
```

```
summary_m1a <- summary(m1a)
```

```
# Extract coefficients and standard errors
coefs <- summary(m1a)$coefficients
OR <- exp(coefs[,1])
lower_ci <- exp(coefs[,1] - 1.96 * coefs[,2])
upper_ci <- exp(coefs[,1] + 1.96 * coefs[,2])
```

```
# Calculate z-values and p-values
z_values <- coefs[,1] / coefs[,2]
p_values <- 2 * (1 - pnorm(abs(z_values)))

# Combine everything into a data frame for easy viewing
results <- data.frame(OR = OR, LowerCI = lower_ci, UpperCI = upper_ci, p_value = p_values)
print(results)
```

```
##              OR      LowerCI      UpperCI      p_value
## 1|2  7.098397e+156  3.817148e+64  1.320023e+249  0.0008626939
## 2|3  5.984716e+157  2.942469e+65  1.217237e+250  0.0008077966
## 3|4  2.197013e+158  1.041525e+66  4.634421e+250  0.0007751331
## Year  1.196741e+00  1.077146e+00  1.329615e+00  0.0008275139
```

```
#Check the Assumptions
nominal_test(m1a) # This is a test of the proportional odds assumption, odds are proportional in this c
```

```
## Tests of nominal effects
##
## formula: Complete ~ Year
##      Df  logLik    AIC LRT Pr(>Chi)
## <none>   -238.52 485.05
## Year
```

```
scale_test(m1a) # scale test checks for equal variance/ scale across year, assumptions are not violated
```

```
## Warning: (-2) Model failed to converge: degenerate Hessian with 1 negative eigenvalues
## In addition: iteration limit reached
```

```
## Tests of scale effects
##
## formula: Complete ~ Year
##      Df  logLik    AIC LRT Pr(>Chi)
## <none>   -238.52 485.05
## Year
```

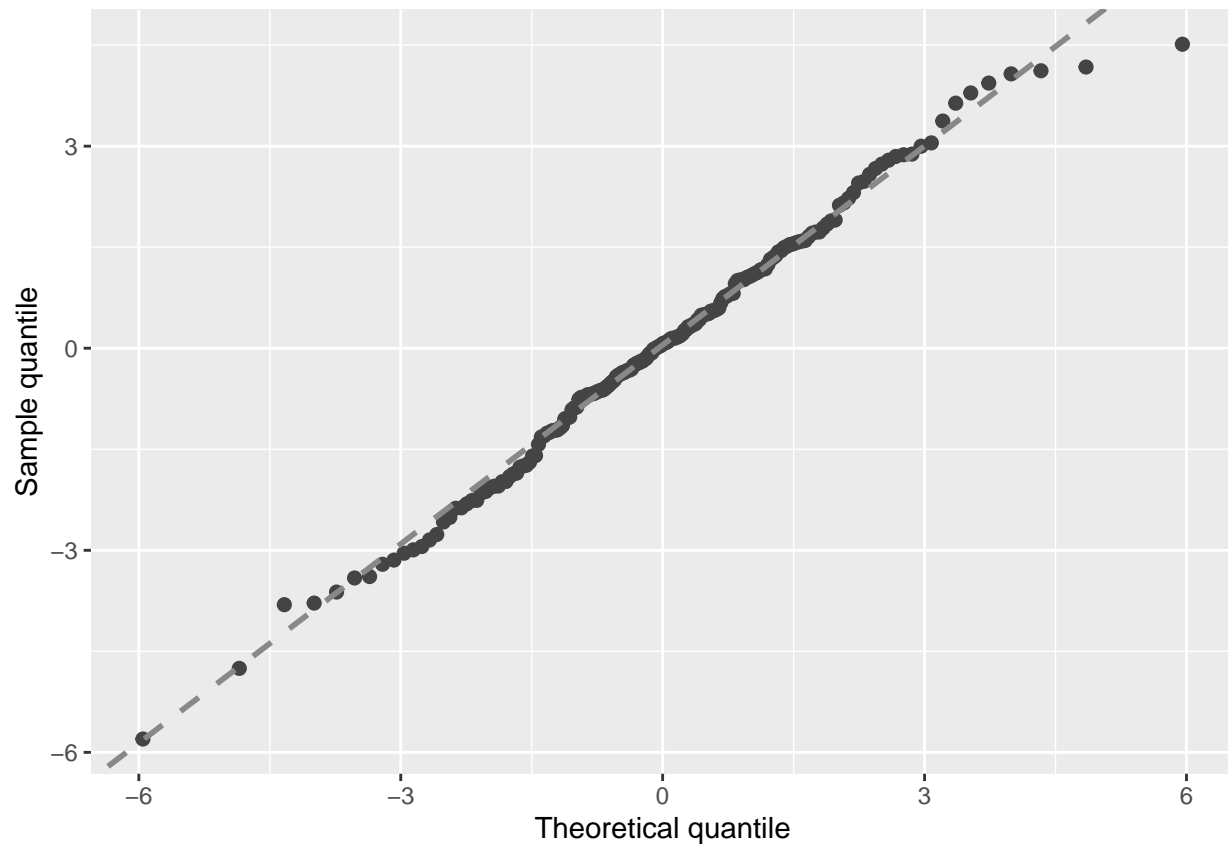
```
convergence(m1a) # This is another way to assess the model
```

```
## nobs logLik  niter max.grad cond.H  logLik.Error
## 193  -238.52 5(0)  9.97e-08 8.1e+12 <1e-10
##
##      Estimate  Std.Err  Gradient      Error Cor.Dec Sig.Dig
## 1|2  361.1631 108.39704  4.74e-10 -3.95e-09      8      11
## 2|3  363.2951 108.44275 -4.15e-10 -3.97e-09      8      11
## 3|4  364.5955 108.46135 -1.03e-11 -3.97e-09      8      11
## Year   0.1796   0.05372 -9.97e-08 -1.96e-12     11     11
##
## Eigen values of Hessian:
## 2.299e+08 7.161e+01 2.950e+01 2.836e-05
##
```

```
## Convergence message from clm:
## (2) Model is nearly unidentifiable: very large eigenvalue
## - Rescale variables?
## (3) Model is nearly unidentifiable: large eigenvalue ratio
## - Rescale variables?
## In addition: Absolute and relative convergence criteria were met
```

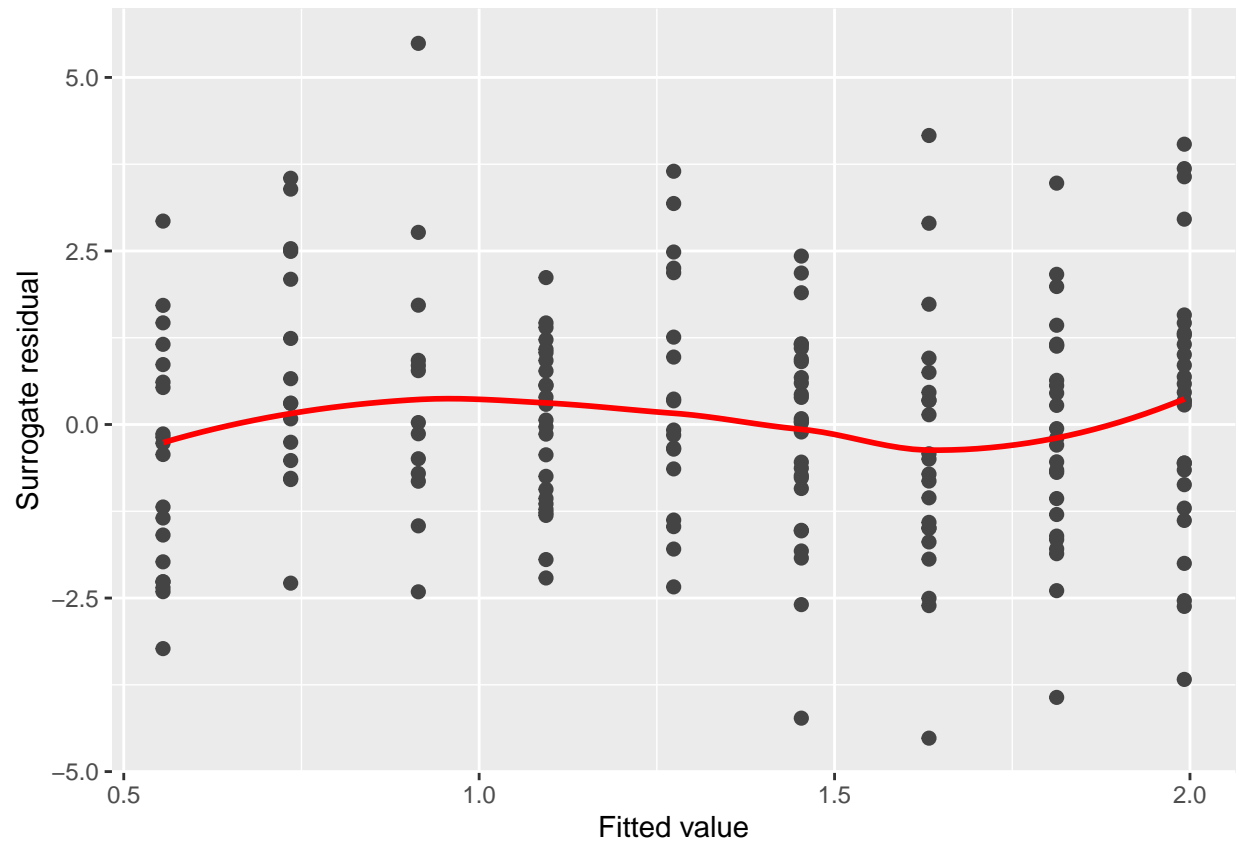
```
##### Graphically validate proportional odds using the sure package #####
```

```
autoplot.clm(m1a, what = c("qq")) # most of the points fall along the line, so no violation of linearity
```



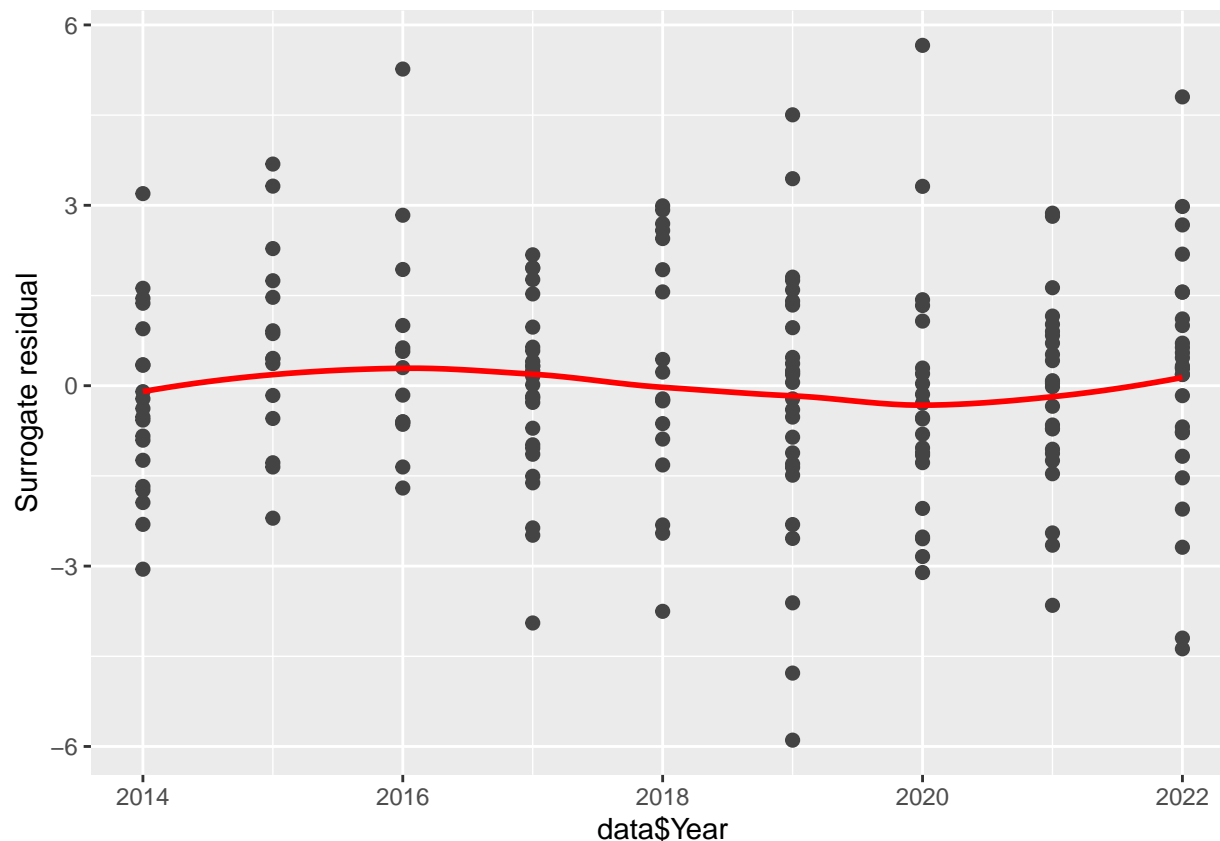
```
autoplot.clm(m1a, what = c("fitted")) # the residuals do not show a clear pattern or trend
```

```
## 'geom_smooth()' using method = 'loess' and formula 'y ~ x'
```



```
autoplot.clm(m1a, what = c("covariate"), x = data$Year) # scale test indicated no variance issues. This
```

```
## 'geom_smooth()' using method = 'loess' and formula 'y ~ x'
```



N.B. - Interpreting residual plots is largely subjective!

###plotting the model

Predict probabilities for each level of 'Complete'

```
new_data <- data.frame(Year = sort(unique(data$Year)))
```

```
pred_probs <- predict(m1a, newdata = new_data, type = "prob")
```

Add the 'Year' column to the predicted probabilities dataframe

```
pred_probs_df <- cbind(new_data, as.data.frame(pred_probs))
```

Convert the predicted probabilities to a long format for ggplot

```
library(reshape2)
```

```
## Warning: package 'reshape2' was built under R version 4.2.3
```

```
##
```

```
## Attaching package: 'reshape2'
```

```
## The following object is masked from 'package:tidyr':
```

```
##
```

```
## smiths
```

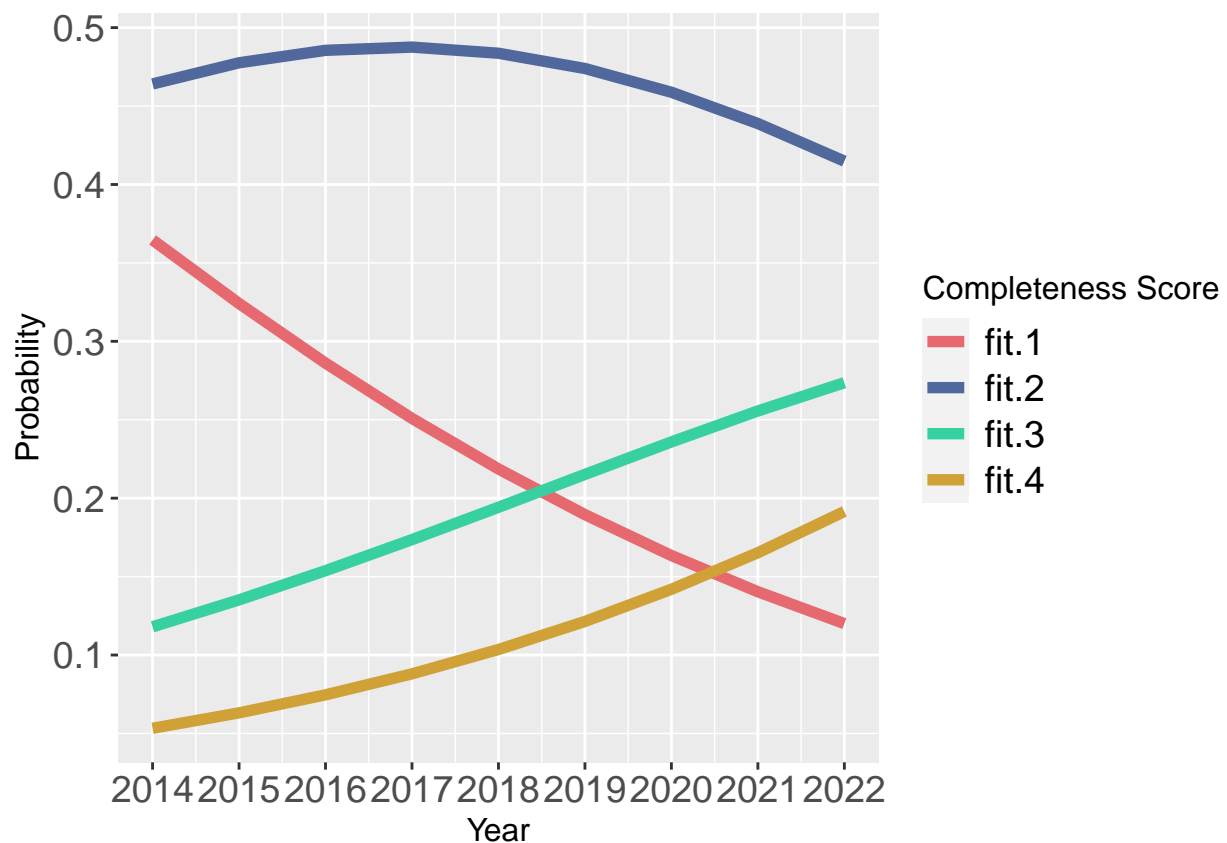
```
pred_probs_long <- melt(pred_probs_df, id.vars = 'Year', variable.name = 'CompleteLevel', value.name =
```

```

# Set the colors
my_colors <- c("#E5696F", "#50689B", "#36D0A1", "#D0A136")

# Modify ggplot command
Figure_3_1 <- ggplot(pred_probs_long, aes(x = Year, y = PredictedProbability, group = CompleteLevel, color = CompleteLevel)) +
  geom_line(size = 2) + # Set size of the lines to make them thicker
  scale_color_manual(values = my_colors) +
  scale_x_continuous(breaks = unique(pred_probs_long$Year)) + # Show all years on x-axis
  labs(x = "Year", y = "Probability", color = "Completeness Score") +
  theme(
    axis.title = element_text(size = 12), # Increase axis titles
    axis.text = element_text(size = 14), # Increase axis text
    legend.title = element_text(size = 12), # Increase legend title
    legend.text = element_text(size = 14) # Increase legend text
  )
print(Figure_3_1)

```



```

ggsave("Figure_3_1.png", Figure_3_1, width = 15, height = 10, units = "in", bg = "white")

#these results were used in the Forth result paragraph and Figure 3 of the publication

```

#Ordinal regression model for the Reusibility by year

```
m1a <- clm(Reuse ~ Year, data = data)
```

```
## Warning in x$code == 0L || action == "silent": 'length(x) = 2 > 1' in coercion
## to 'logical(1)'
```

```
## Warning: (2) Model is nearly unidentifiable: very large eigenvalue
## - Rescale variables?
## In addition: Absolute and relative convergence criteria were met
```

```
summary_m1a <- summary(m1a)
```

```
# Extract coefficients and standard errors
```

```
coefs <- summary(m1a)$coefficients
OR <- exp(coefs[,1])
lower_ci <- exp(coefs[,1] - 1.96 * coefs[,2])
upper_ci <- exp(coefs[,1] + 1.96 * coefs[,2])
```

```
# Calculate z-values and p-values
```

```
z_values <- coefs[,1] / coefs[,2]
p_values <- 2 * (1 - pnorm(abs(z_values)))
```

```
# Combine everything into a data frame for easy viewing
```

```
results <- data.frame(OR = OR, LowerCI = lower_ci, UpperCI = upper_ci, p_value = p_values)
print(results)
```

```
##              OR      LowerCI      UpperCI      p_value
## 1|2  2.349088e+189  9.057314e+96  6.092548e+281  5.910538e-05
## 2|3  4.254302e+189  1.602643e+97  1.129328e+282  5.785682e-05
## 3|4  1.399145e+190  4.955124e+97  3.950672e+282  5.549741e-05
## Year  1.241997e+00  1.117686e+00  1.380133e+00  5.629614e-05
```

```
#Check the Assumptions
```

```
nominal_test(m1a) # This is a test of the proportional odds assumption, odds are proportional in this c
```

```
## Tests of nominal effects
```

```
##
```

```
## formula: Reuse ~ Year
```

```
##      Df  logLik    AIC LRT Pr(>Chi)
```

```
## <none>    -240.66 489.32
```

```
## Year
```

```
scale_test(m1a) # scale test checks for equal variance/ scale across year, assumptions are not violated
```

```
## Warning: (-1) Model failed to converge with max|grad| = 451.581 (tol = 1e-06)
```

```
## In addition: iteration limit reached
```

```
## Tests of scale effects
```

```
##
```

```
## formula: Reuse ~ Year
```

```
##      Df  logLik    AIC LRT Pr(>Chi)
```

```
## <none>    -240.66 489.32
```

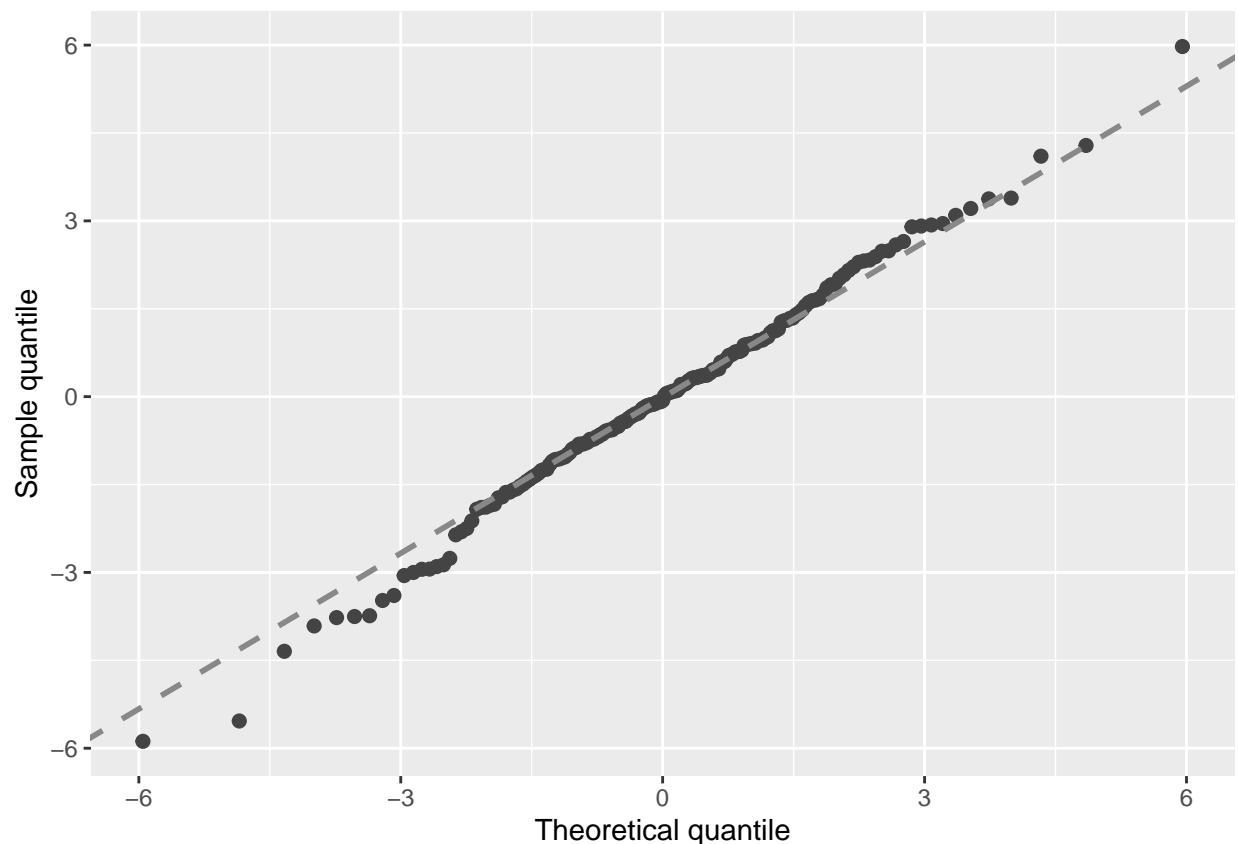
```
## Year
```

```
convergence(m1a) # This is another way to assess the model
```

```
## nobs logLik niter max.grad cond.H logLik.Error
## 193 -240.66 7(0) 1.79e-09 8.2e+12 <1e-10
##
##      Estimate   Std.Err   Gradient   Error Cor.Dec Sig.Dig
## 1|2  436.0426 108.56677  2.06e-12 -2.72e-11    10    13
## 2|3  436.6365 108.57863 -2.78e-12 -2.73e-11    10    13
## 3|4  437.8270 108.61013  1.60e-12 -2.73e-11    10    13
## Year   0.2167   0.05381 -1.79e-09 -1.35e-14    13    13
##
## Eigen values of Hessian:
## 2.310e+08 1.550e+02 5.908e+01 2.827e-05
##
## Convergence message from clm:
## (2) Model is nearly unidentifiable: very large eigenvalue
## - Rescale variables?
## (3) Model is nearly unidentifiable: large eigenvalue ratio
## - Rescale variables?
## In addition: Absolute and relative convergence criteria were met
```

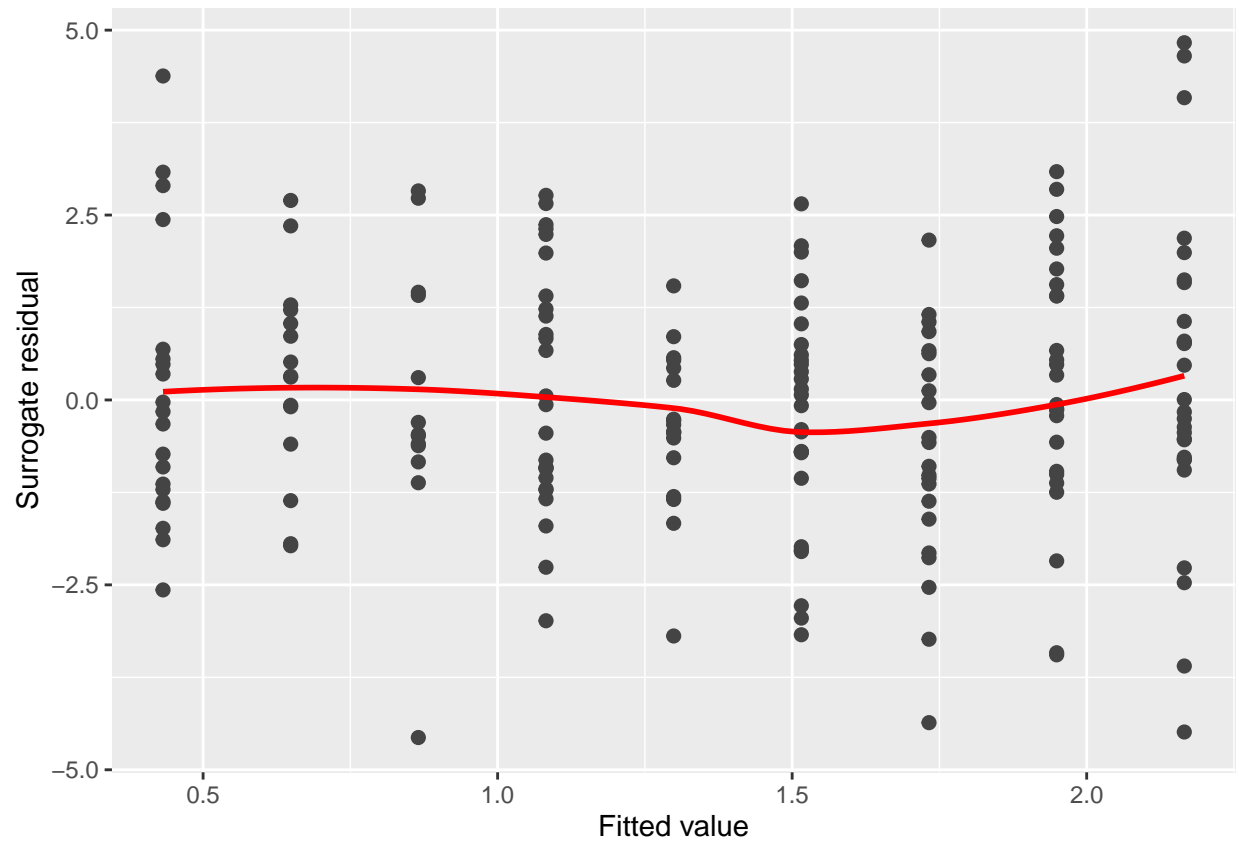
```
##### Graphically validate proportional odds using the sure package #####
```

```
autoplot.clm(m1a, what = c("qq")) # most of the points fall along the line, so no violation of linearity
```



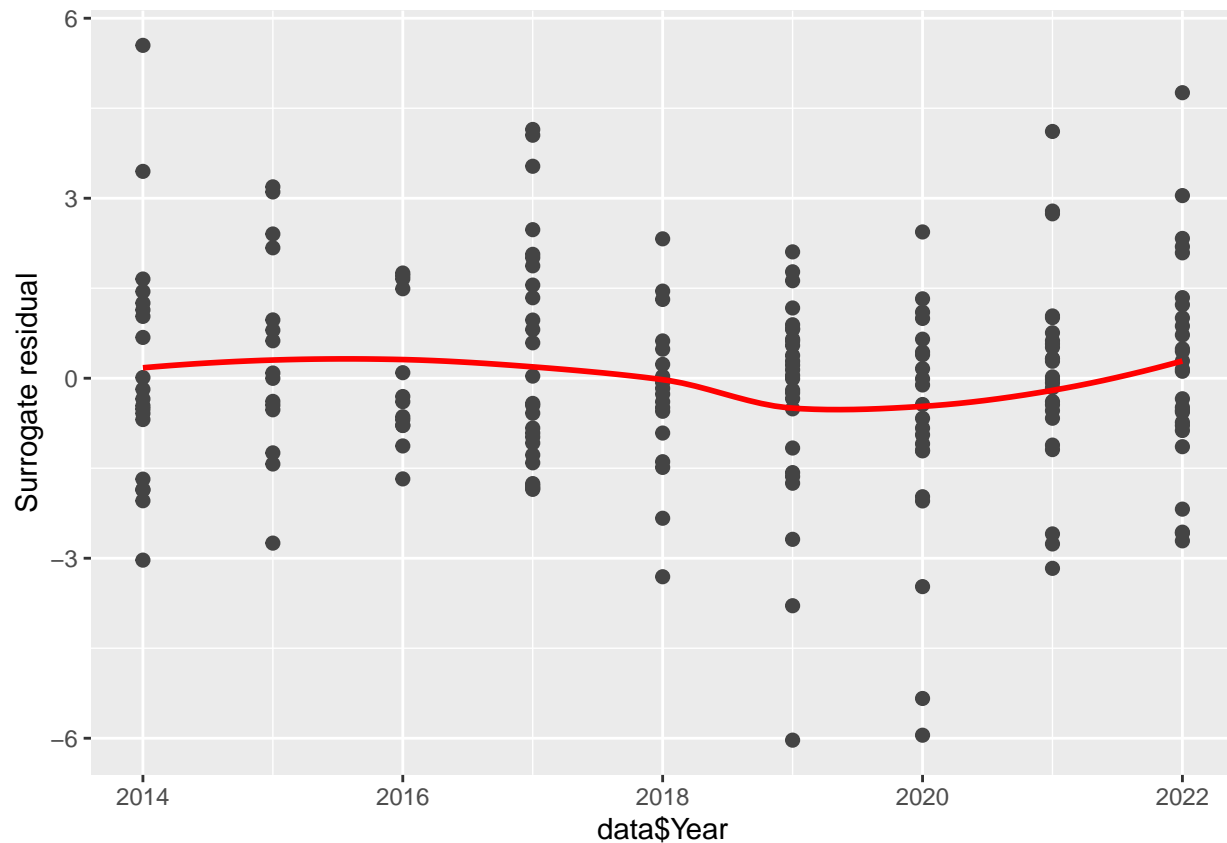

```
autoplot.clm(m1a, what = c("fitted")) # the residuals do not show a clear pattern or trend
```

```
## 'geom_smooth()' using method = 'loess' and formula 'y ~ x'
```



```
autoplot.clm(m1a, what = c("covariate"), x = data$Year) # scale test indicated no variance issues. This
```

```
## 'geom_smooth()' using method = 'loess' and formula 'y ~ x'
```



N.B. - Interpreting residual plots is largely subjective!

###plotting the model

Predict probabilities for each level of 'Complete'

```
new_data <- data.frame(Year = sort(unique(data$Year)))
```

```
pred_probs <- predict(m1a, newdata = new_data, type = "prob")
```

Add the 'Year' column to the predicted probabilities dataframe

```
pred_probs_df <- cbind(new_data, as.data.frame(pred_probs))
```

Convert the predicted probabilities to a long format for ggplot

```
library(reshape2)
```

```
pred_probs_long <- melt(pred_probs_df, id.vars = 'Year', variable.name = 'Reuselevel', value.name = 'Pr
```

Set the colors

```
my_colors <- c("#E5696F", "#50689B", "#36D0A1", "#D0A136")
```

Modify ggplot command

```
Figure_3_2 <- ggplot(pred_probs_long, aes(x = Year, y = PredictedProbability, group = Reuselevel, color
```

```
  geom_line(size = 2) + # Set size of the lines to make them thicker
```

```
  scale_color_manual(values = my_colors) + # Use colors
```

```
  scale_x_continuous(breaks = unique(pred_probs_long$Year)) + # Show all years on x-axis
```

```
  labs(x = "Year", y = "Probability", color = "Reusability Score") +
```

```
  theme(
```

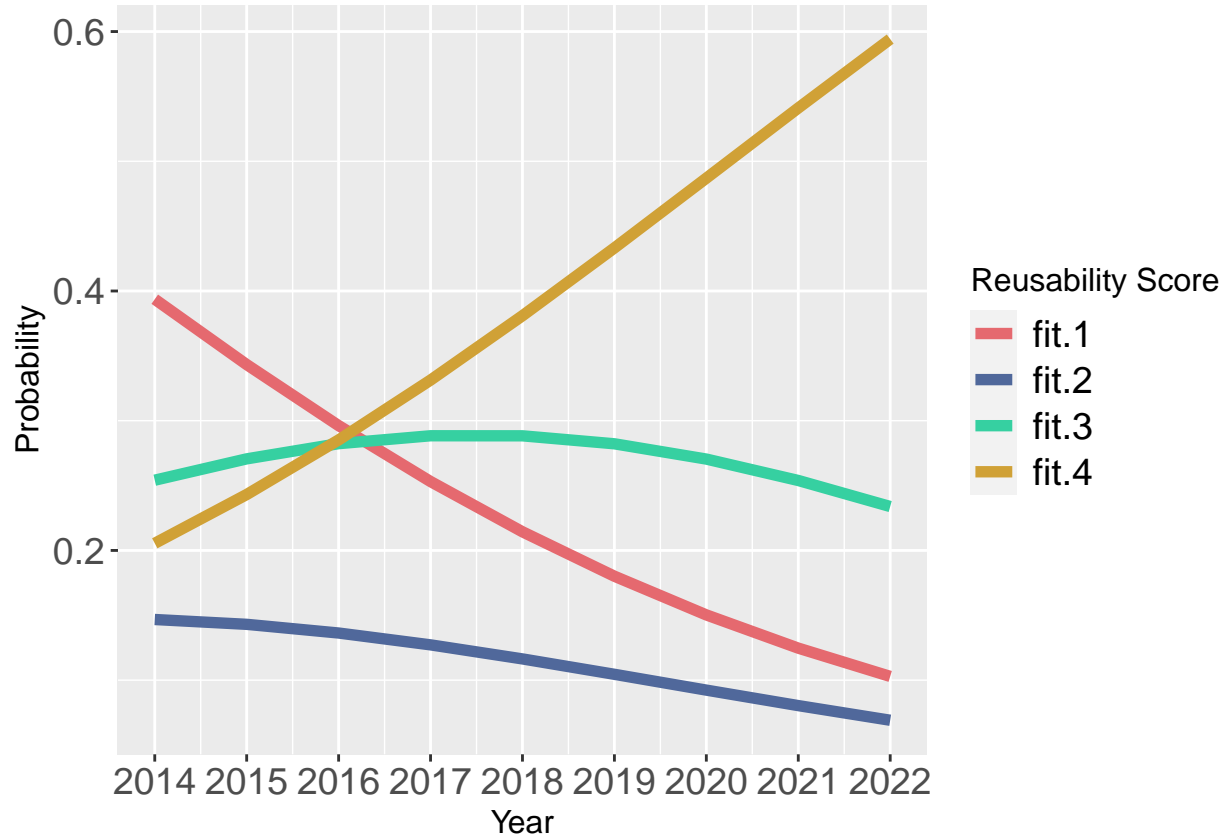
```
    axis.title = element_text(size = 12), # Increase axis titles
```

```
    axis.text = element_text(size = 14), # Increase axis text
```

```

    legend.title = element_text(size = 12), # Increase legend title
    legend.text = element_text(size = 14) # Increase legend text
  )
print(Figure_3_2)

```



```

ggsave("Figure_3_2.png", Figure_3_2, width = 15, height = 10, units = "in", bg = "white")

```

#these results were used in the Forth result paragraph and Figure 3 of the publication

```

m1a <- clm(Access ~ Year, data = data)

```

```

## Warning in x$code == 0L || action == "silent": 'length(x) = 2 > 1' in coercion
## to 'logical(1)'

```

```

## Warning: (2) Model is nearly unidentifiable: very large eigenvalue
## - Rescale variables?
## In addition: Absolute and relative convergence criteria were met

```

```

summary_m1a <- summary(m1a)

```

```

# Extract coefficients and standard errors
coefs <- summary_m1a$coefficients[, 1] # Coefficients

```

```

se_coefs <- summary_m1a$coefficients[, 2] # Standard errors

# Extract coefficients and standard errors
coefs <- summary(m1a)$coefficients
OR <- exp(coefs[,1])
lower_ci <- exp(coefs[,1] - 1.96 * coefs[,2])
upper_ci <- exp(coefs[,1] + 1.96 * coefs[,2])

# Calculate z-values and p-values
z_values <- coefs[,1] / coefs[,2]
p_values <- 2 * (1 - pnorm(abs(z_values)))

# Combine everything into a data frame for easy viewing
results <- data.frame(OR = OR, LowerCI = lower_ci, UpperCI = upper_ci, p_value = p_values)
print(results)

```

```

##              OR      LowerCI      UpperCI      p_value
## 1|2  3.320521e+169  6.180429e+74  1.783996e+264  0.0004524204
## 2|3  1.065638e+170  1.898440e+75  5.981670e+264  0.0004360872
## 3|4  1.273957e+170  2.252216e+75  7.206093e+264  0.0004336612
## Year  1.214171e+00  1.089767e+00  1.352777e+00  0.0004336952

```

#Check the Assumptions

`nominal_test(m1a)` *# This is a test of the proportional odds assumption, odds are proportional in this c*

```

## Tests of nominal effects
##
## formula: Access ~ Year
##      Df  logLik    AIC LRT Pr(>Chi)
## <none>   -215.84 439.68
## Year

```

`scale_test(m1a)` *# scale test checks for equal variance/ scale across year, assumptions are not violated*

```

## Warning: (-2) Model failed to converge: degenerate Hessian with 1 negative eigenvalues
## In addition: iteration limit reached

```

```

## Tests of scale effects
##
## formula: Access ~ Year
##      Df  logLik    AIC LRT Pr(>Chi)
## <none>   -215.84 439.68
## Year

```

`convergence(m1a)` *# This is another way to assess the model*

```

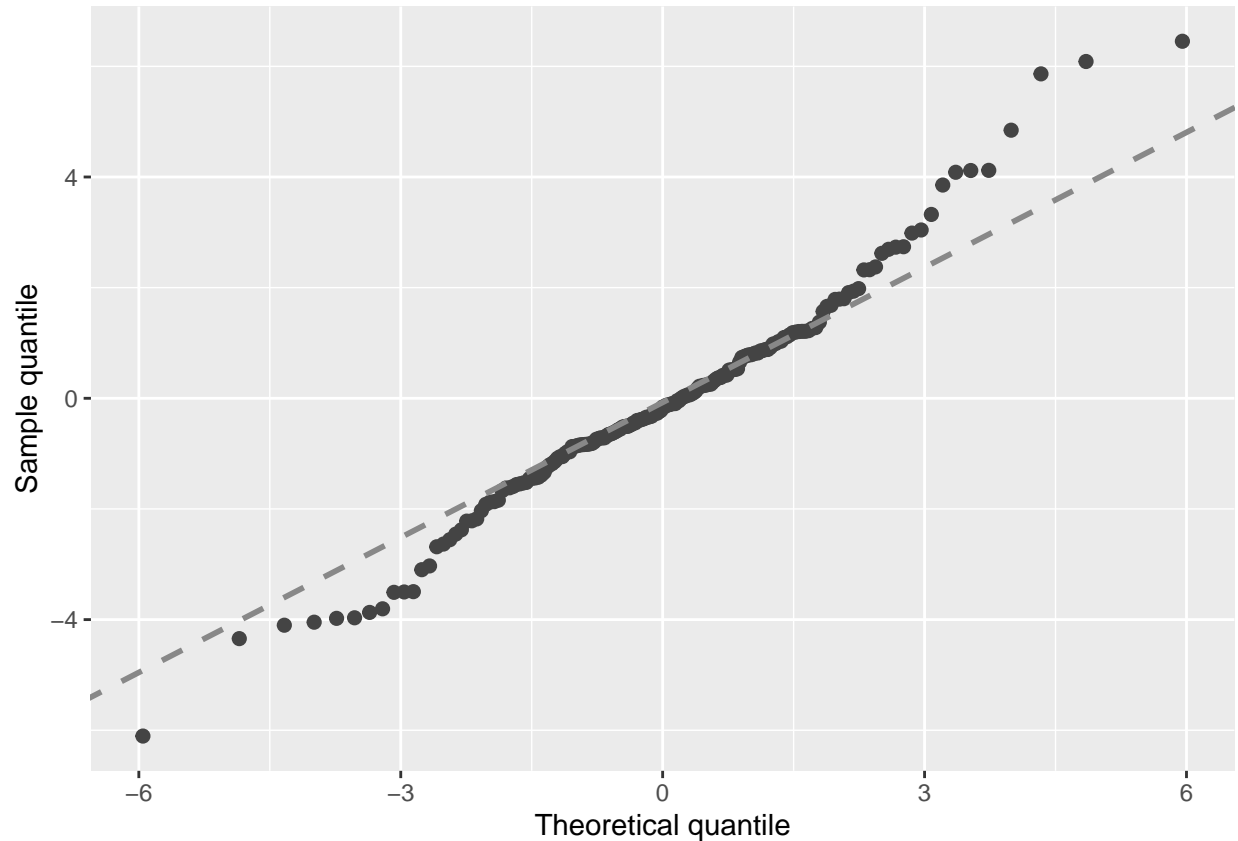
##  nobs logLik  niter max.grad cond.H  logLik.Error
##  193  -215.84 6(2)  4.89e-09 8.0e+12 <1e-10
##
##      Estimate  Std.Err  Gradient      Error Cor.Dec Sig.Dig

```

```
## 1|2 390.3370 111.28792 2.50e-13 -2.16e-10 9 12
## 2|3 391.5030 111.31027 -5.25e-12 -2.16e-10 9 12
## 3|4 391.6816 111.31418 7.41e-12 -2.16e-10 9 12
## Year 0.1941 0.05515 -4.89e-09 -1.07e-13 12 12
##
## Eigen values of Hessian:
## 2.159e+08 5.342e+02 6.066e+01 2.691e-05
##
## Convergence message from clm:
## (2) Model is nearly unidentifiable: very large eigenvalue
## - Rescale variables?
## (3) Model is nearly unidentifiable: large eigenvalue ratio
## - Rescale variables?
## In addition: Absolute and relative convergence criteria were met
```

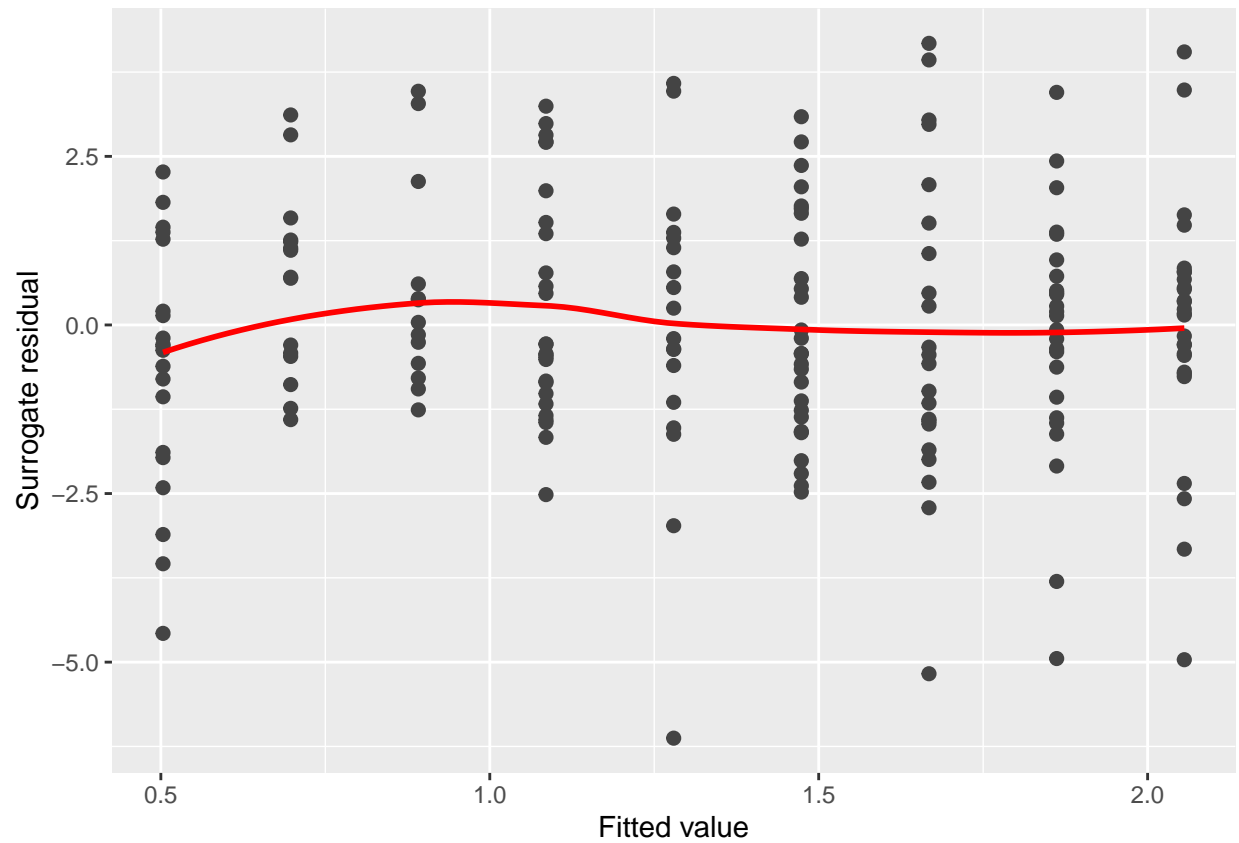
```
##### Graphically validate proportional odds using the sure package #####
```

```
autoplot.clm(m1a, what = c("qq")) # most of the points fall along the line, so no violation of linearity
```



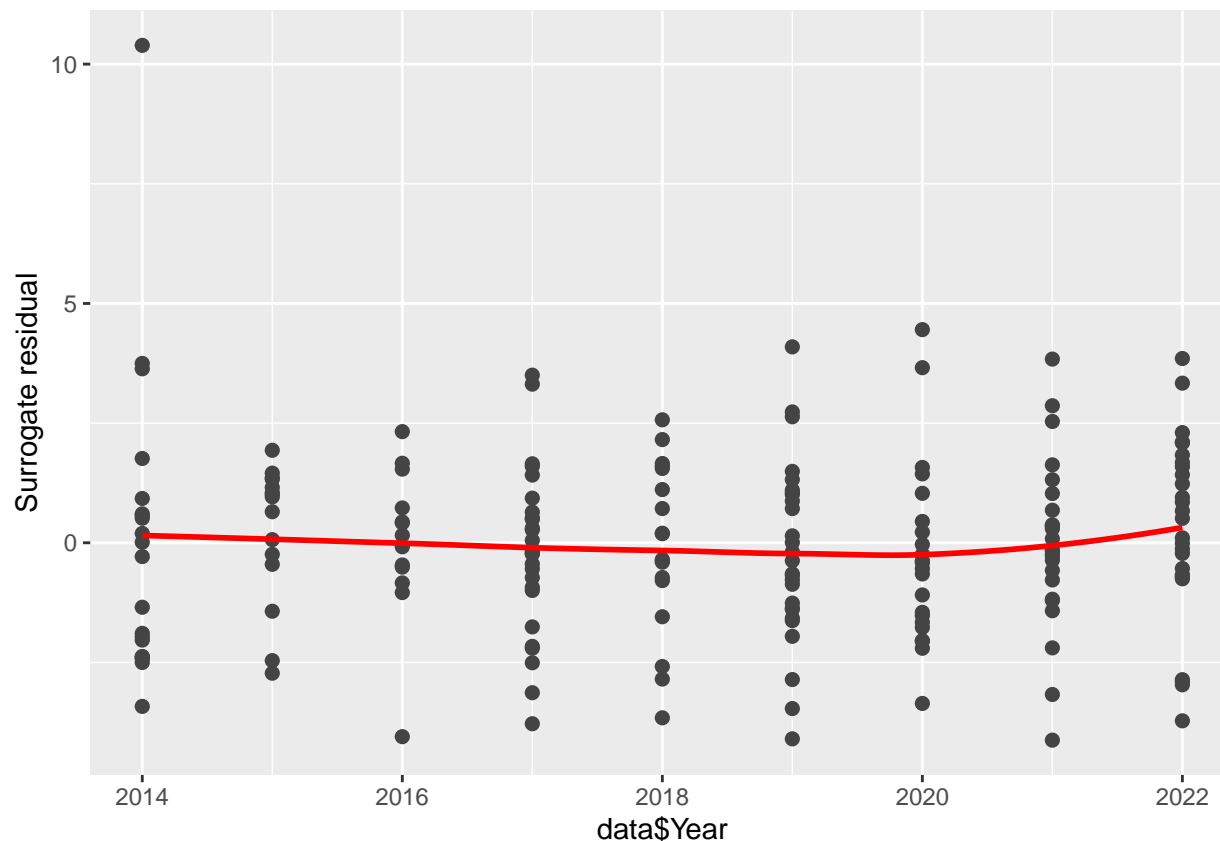
```
autoplot.clm(m1a, what = c("fitted")) # the residuals do not show a clear pattern or trend
```

```
## 'geom_smooth()' using method = 'loess' and formula 'y ~ x'
```



```
autoplot.clm(m1a, what = c("covariate"), x = data$Year) # scale test indicated no variance issues. This
```

```
## 'geom_smooth()' using method = 'loess' and formula 'y ~ x'
```



N.B. - Interpreting residual plots is largely subjective!

###plotting the model

Predict probabilities for each level of 'Complete'

```
new_data <- data.frame(Year = sort(unique(data$Year)))
```

```
pred_probs <- predict(m1a, newdata = new_data, type = "prob")
```

Add the 'Year' column to the predicted probabilities dataframe

```
pred_probs_df <- cbind(new_data, as.data.frame(pred_probs))
```

Convert the predicted probabilities to a long format for ggplot

```
library(reshape2)
```

```
pred_probs_long <- melt(pred_probs_df, id.vars = 'Year', variable.name = 'AccessLevel', value.name = 'P')
```

Set the colors

```
my_colors <- c("#E5696F", "#50689B", "#36D0A1", "#D0A136")
```

Modify ggplot command

```
Figure_3_3 <- ggplot(pred_probs_long, aes(x = Year, y = PredictedProbability, group = AccessLevel, color = AccessLevel))
```

```
  geom_line(size = 2) + # Set size of the lines to make them thicker
```

```
  scale_color_manual(values = my_colors) + # Use colors
```

```
  scale_x_continuous(breaks = unique(pred_probs_long$Year)) + # Show all years on x-axis
```

```
  labs(x = "Year", y = "Probability", color = "Accessibility Score") +
```

```
  theme(
```

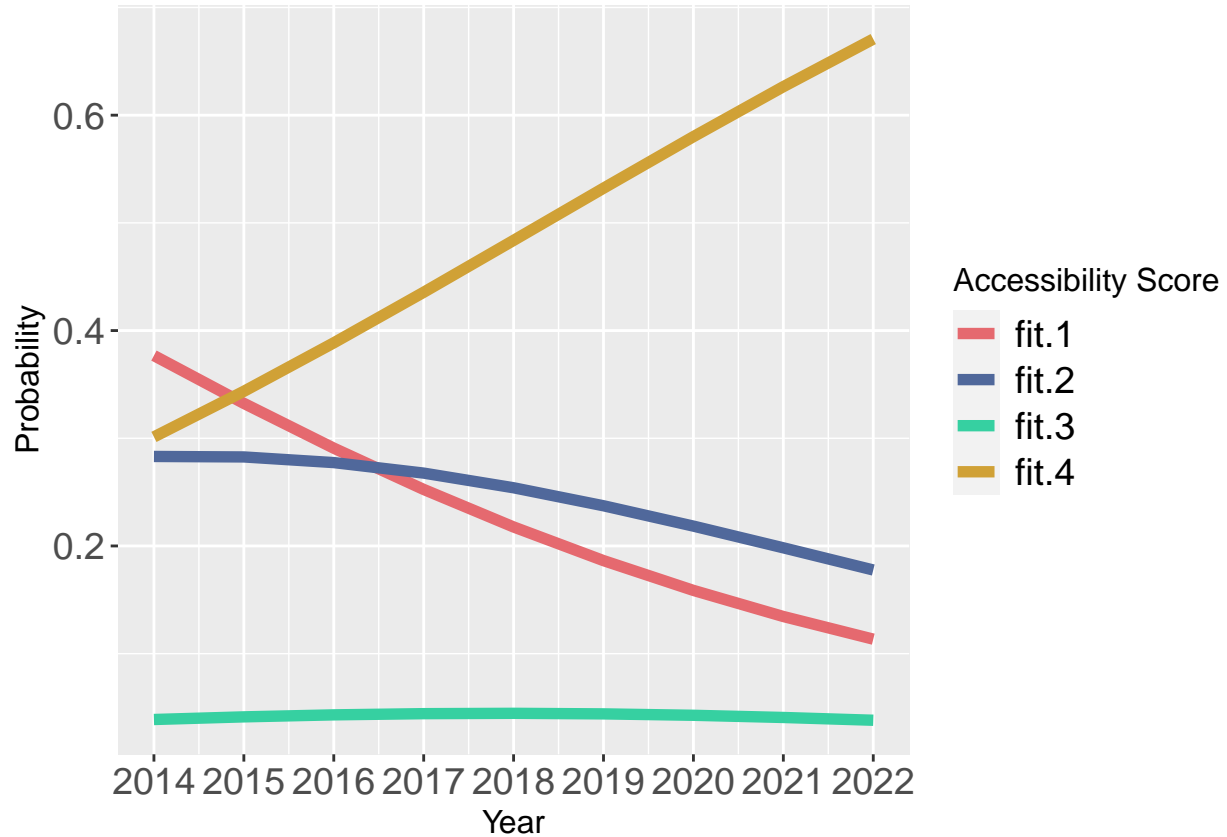
```
    axis.title = element_text(size = 12), # Increase axis titles
```

```
    axis.text = element_text(size = 14), # Increase axis text
```

```

    legend.title = element_text(size = 12), # Increase legend title
    legend.text = element_text(size = 14) # Increase legend text
  )
print(Figure_3_3)

```



```

ggsave("Figure_3_3.png", Figure_3_3, width = 15, height = 10, units = "in", bg = "white")

```

#these results were used in the Forth result paragraph and Figure 3 of the publication

```

m1a <- clm(Licence ~ Year, data = data)

```

```

## Warning in x$code == 0L || action == "silent": 'length(x) = 2 > 1' in coercion
## to 'logical(1)'

```

```

## Warning: (2) Model is nearly unidentifiable: very large eigenvalue
## - Rescale variables?
## In addition: Absolute and relative convergence criteria were met

```

```

summary_m1a <- summary(m1a)

```

```

# Extract coefficients and standard errors
coefs <- summary_m1a$coefficients[, 1] # Coefficients

```



```

se_coefs <- summary_m1a$coefficients[, 2] # Standard errors

# Extract coefficients and standard errors
coefs <- summary(m1a)$coefficients
OR <- exp(coefs[,1])
lower_ci <- exp(coefs[,1] - 1.96 * coefs[,2])
upper_ci <- exp(coefs[,1] + 1.96 * coefs[,2])

# Calculate z-values and p-values
z_values <- coefs[,1] / coefs[,2]
p_values <- 2 * (1 - pnorm(abs(z_values)))

# Combine everything into a data frame for easy viewing
results <- data.frame(OR = OR, LowerCI = lower_ci, UpperCI = upper_ci, p_value = p_values)
print(results)

```

```

##              OR      LowerCI      UpperCI      p_value
## 1|2  2.256630e+132  9.929195e+27  5.128692e+236  0.01292492
## 2|3  2.621908e+132  1.149732e+28  5.979135e+236  0.01288181
## 3|4  3.805940e+132  1.653113e+28  8.762366e+236  0.01277571
## Year  1.163746e+00  1.033102e+00  1.310911e+00  0.01255954

```

```

#Check the Assumptions
nominal_test(m1a) # This is a test of the proportional odds assumption, odds are proportional in this c

```

```

## Warning in x$code == OL || action == "silent": 'length(x) = 2 > 1' in coercion
## to 'logical(1)'

```

```

## Warning in !inherits(nfit, "try-error") && nfit$convergence$code >= 0:
## 'length(x) = 2 > 1' in coercion to 'logical(1)'

```

```

## Tests of nominal effects
##
## formula: Licence ~ Year
##      Df  logLik    AIC      LRT Pr(>Chi)
## <none>   -166.02 340.03
## Year     2 -165.98 343.97 0.062447   0.9693

```

```

scale_test(m1a) # scale test checks for equal variance/ scale across year, assumptions are not violated

```

```

## Warning: (-1) Model failed to converge with max|grad| = 41.6269 (tol = 1e-06)
## In addition: iteration limit reached

```

```

## Tests of scale effects
##
## formula: Licence ~ Year
##      Df  logLik    AIC  LRT Pr(>Chi)
## <none>   -166.02 340.03
## Year

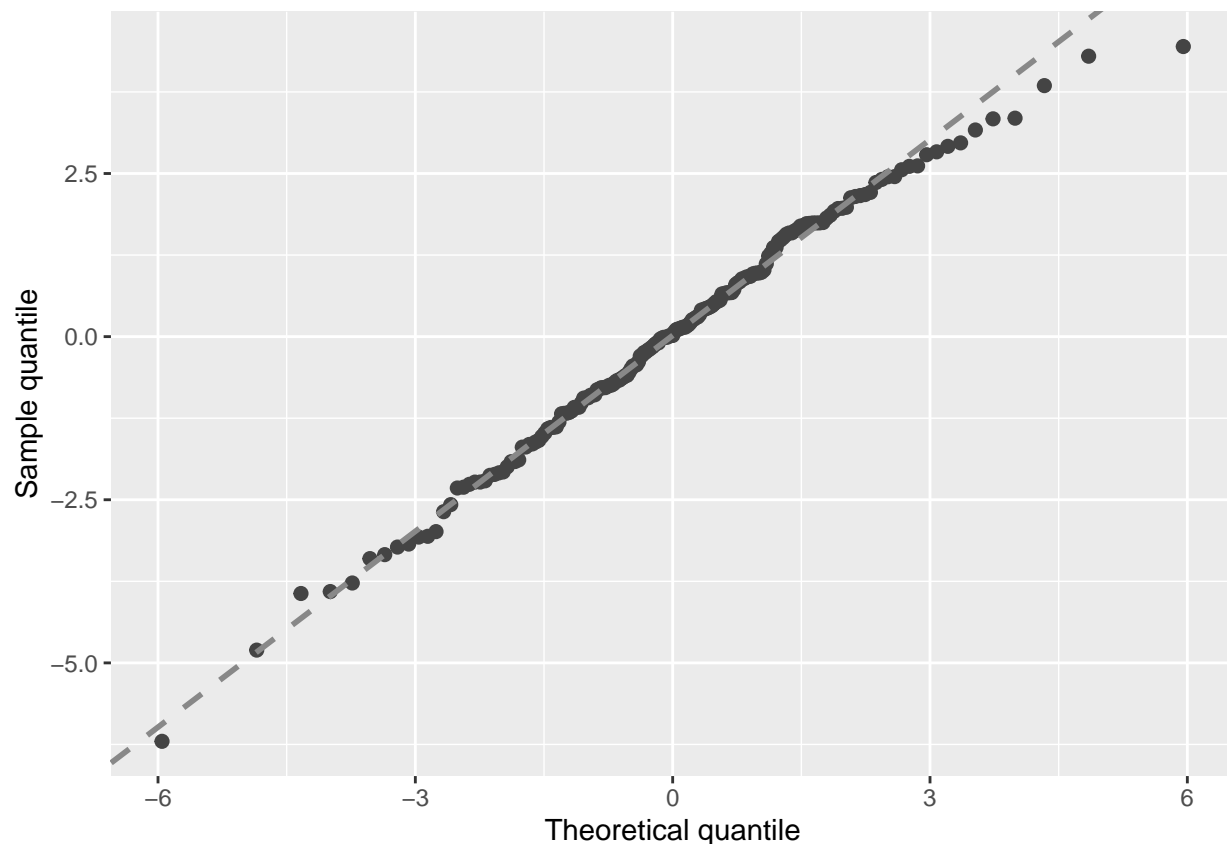
```

```
convergence(m1a) # This is another way to assess the model
```

```
## nobs logLik niter max.grad cond.H logLik.Error
## 193 -166.02 6(2) 1.68e-08 7.7e+12 <1e-10
##
##      Estimate   Std.Err   Gradient   Error Cor.Dec Sig.Dig
## 1|2  304.7551 122.59685  1.30e-09 -3.27e-10     9    12
## 2|3  304.9051 122.59858 -1.30e-09 -3.33e-10     9    12
## 3|4  305.2778 122.60345  7.54e-12 -3.32e-10     9    12
## Year   0.1516   0.06075 -1.68e-08 -1.64e-13    12    12
##
## Eigen values of Hessian:
## 1.709e+08 5.184e+02 1.427e+02 2.219e-05
##
## Convergence message from clm:
## (2) Model is nearly unidentifiable: very large eigenvalue
## - Rescale variables?
## (3) Model is nearly unidentifiable: large eigenvalue ratio
## - Rescale variables?
## In addition: Absolute and relative convergence criteria were met
```

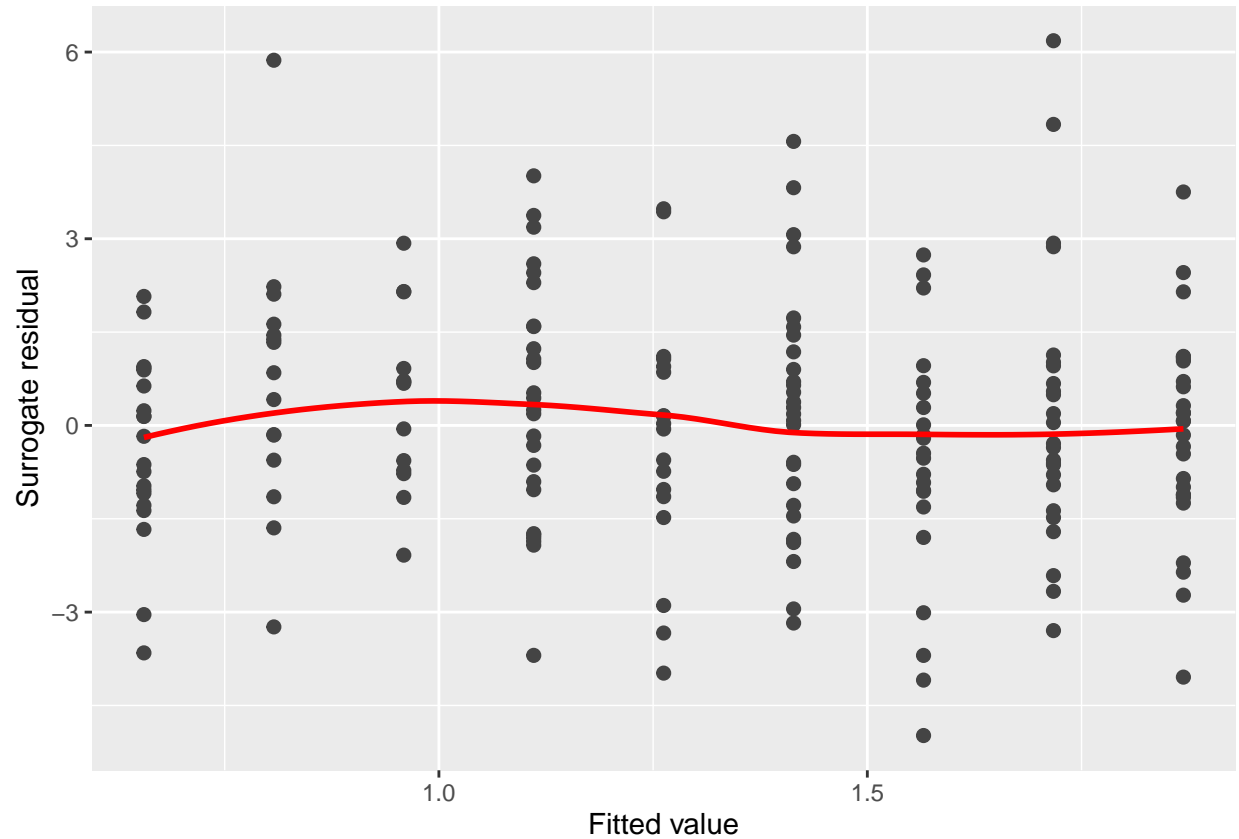
```
##### Graphically validate proportional odds using the sure package #####
```

```
autoplot.clm(m1a, what = c("qq")) # most of the points fall along the line, so no violation of linearity
```



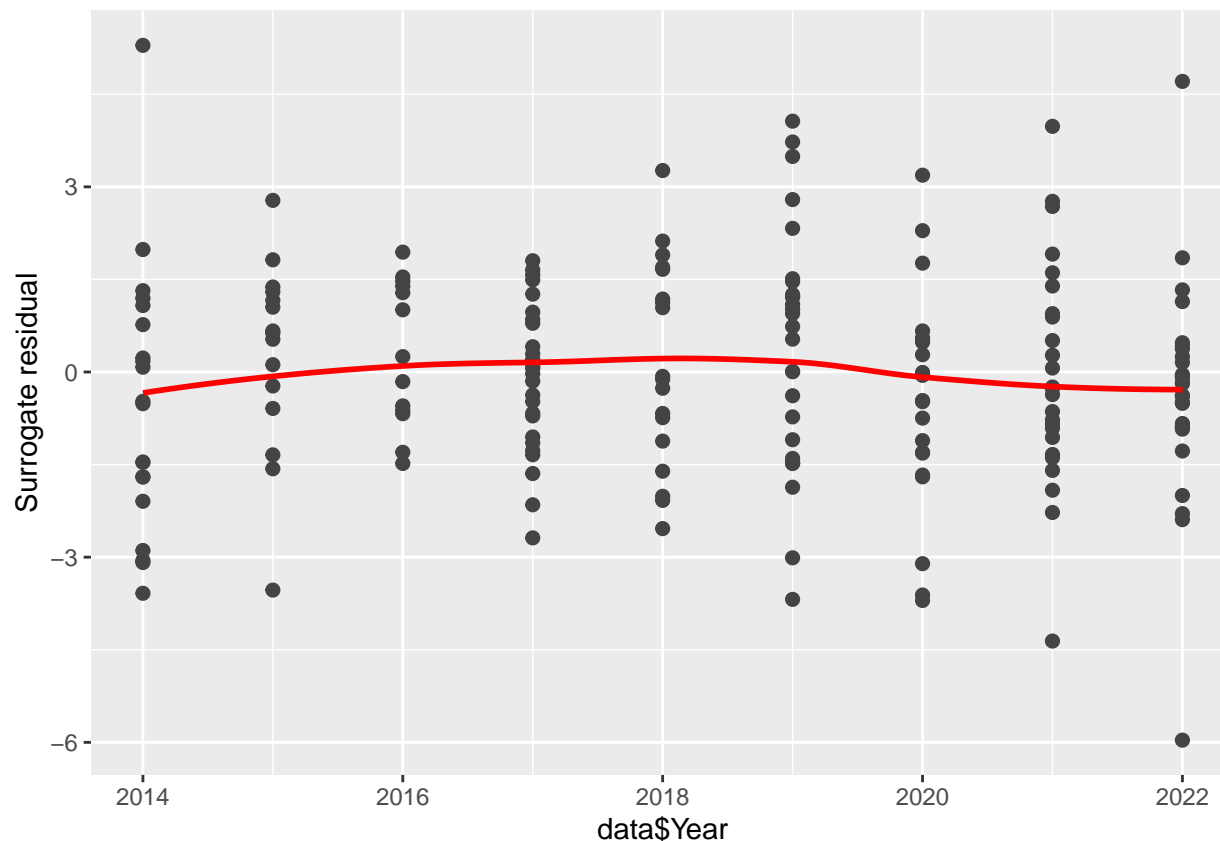
```
autoplot.clm(m1a, what = c("fitted")) # the residuals do not show a clear pattern or trend
```

```
## 'geom_smooth()' using method = 'loess' and formula 'y ~ x'
```



```
autoplot.clm(m1a, what = c("covariate"), x = data$Year) # scale test indicated no variance issues. This
```

```
## 'geom_smooth()' using method = 'loess' and formula 'y ~ x'
```



N.B. - Interpreting residual plots is largely subjective!

###plotting the model

Predict probabilities for each level of 'Complete'

```
new_data <- data.frame(Year = sort(unique(data$Year)))
```

```
pred_probs <- predict(m1a, newdata = new_data, type = "prob")
```

Add the 'Year' column to the predicted probabilities dataframe

```
pred_probs_df <- cbind(new_data, as.data.frame(pred_probs))
```

Convert the predicted probabilities to a long format for ggplot

```
library(reshape2)
```

```
pred_probs_long <- melt(pred_probs_df, id.vars = 'Year', variable.name = 'Licencelevel', value.name = 'PredictedProbability')
```

Set the colors

```
my_colors <- c("#E5696F", "#50689B", "#36D0A1", "#D0A136")
```

Modify ggplot command

```
Figure_3_4 <- ggplot(pred_probs_long, aes(x = Year, y = PredictedProbability, group = Licencelevel, color = Licencelevel))
```

```
  geom_line(size = 2) + # Set size of the lines to make them thicker
```

```
  scale_color_manual(values = my_colors) + # Use colors
```

```
  scale_x_continuous(breaks = unique(pred_probs_long$Year)) + # Show all years on x-axis
```

```
  labs(x = "Year", y = "Probability", color = "Licence Score") +
```

```
  theme(
```

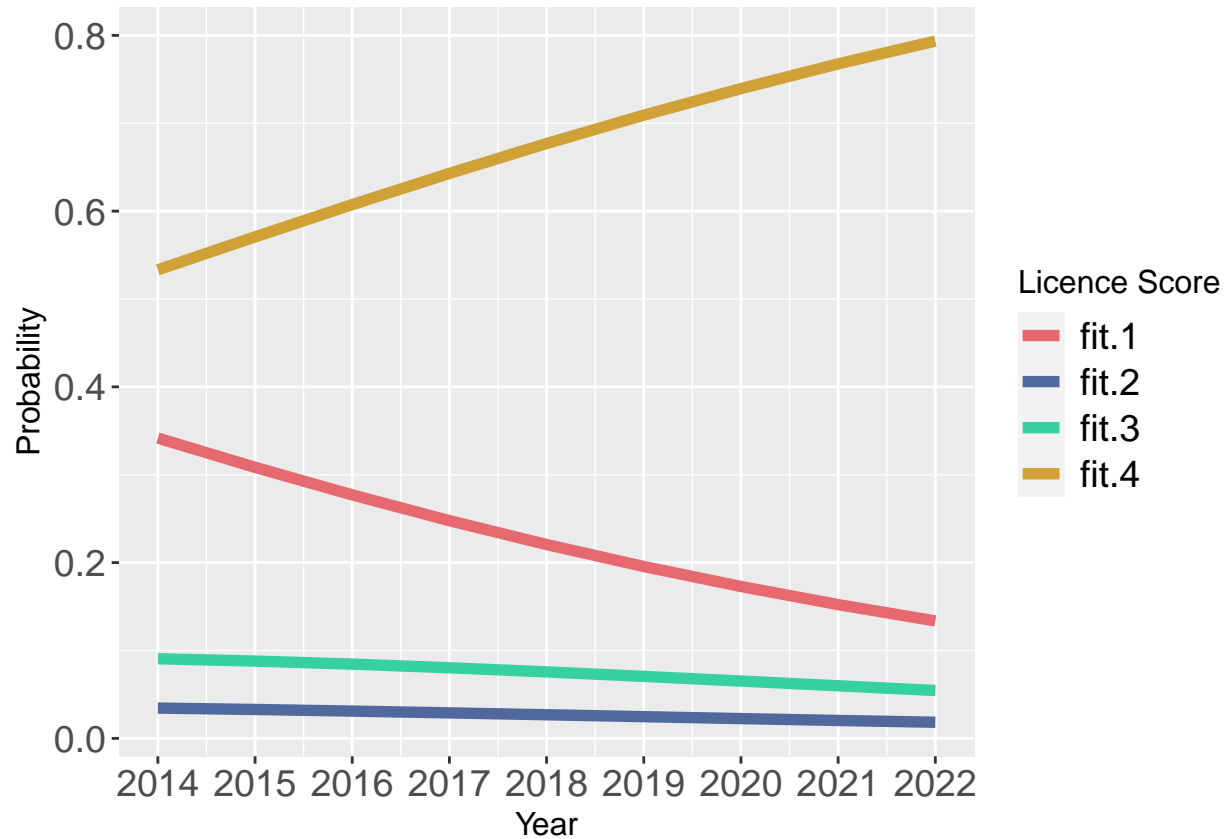
```
    axis.title = element_text(size = 12), # Increase axis titles
```

```
    axis.text = element_text(size = 14), # Increase axis text
```

```

    legend.title = element_text(size = 12), # Increase legend title
    legend.text = element_text(size = 14)  # Increase legend text
  )
  print(Figure_3_4)

```



```

ggsave("Figure_3_4.png", Figure_3_4, width = 15, height = 10, units = "in", bg = "white")

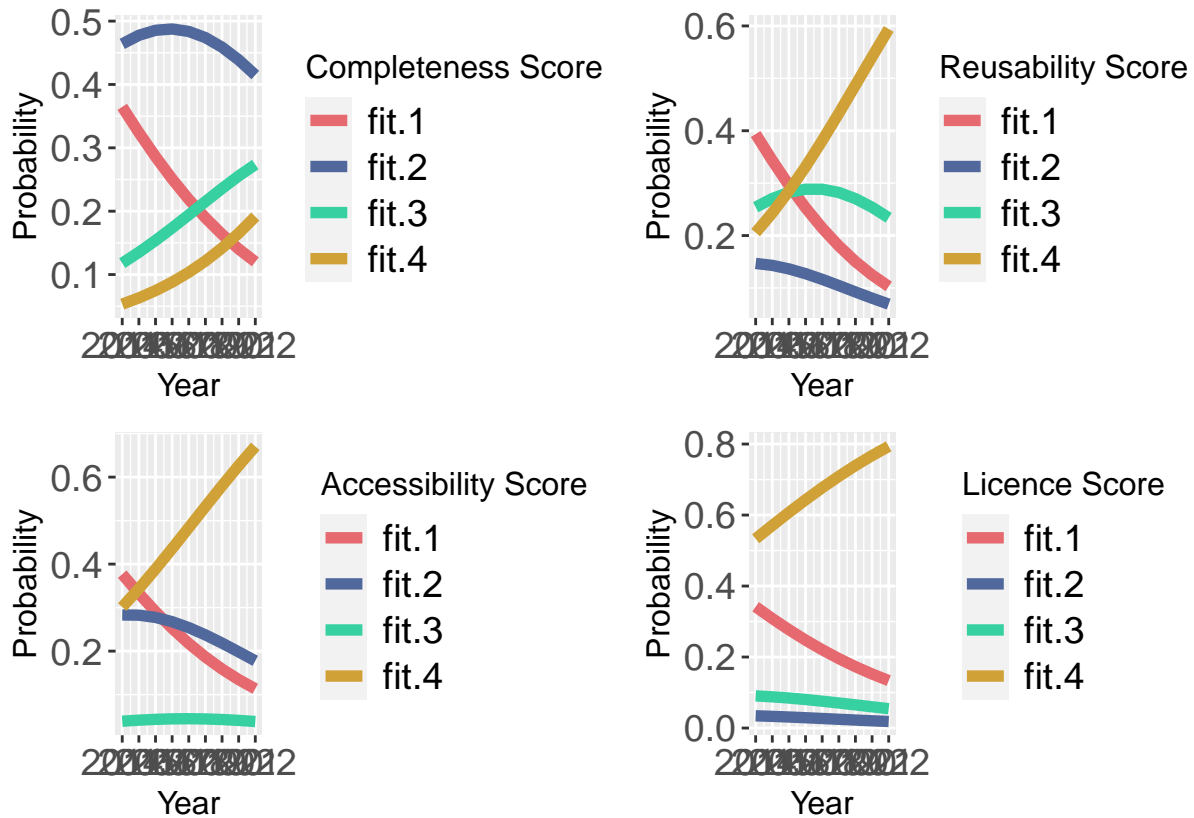
```

#these results were used in the Forth result paragraph and Figure 3 of the publication

```

Figure_3 <- Figure_3_1 + Figure_3_2 + Figure_3_3 + Figure_3_4 +
  plot_layout(
    ncol = 2, heights = c(10, 10), widths = c(10, 10)
  )
print(Figure_3)

```



```
ggsave("Figure_3.png", Figure_3, width = 15, height = 10, units = "in", bg="white")
```

#these results were used in the Figure 3 of the publication

####3

#Ordinal Regression models depending on the sharing projects

```
m1a <- clm(Complete ~ NewDAS + Preprints, data = data)
summary_m1a <- summary(m1a)

# Extract coefficients and standard errors
coefs <- summary_m1a$coefficients[, 1] # Coefficients
se_coefs <- summary_m1a$coefficients[, 2] # Standard errors

# Extract coefficients and standard errors
coefs <- summary(m1a)$coefficients
OR <- exp(coefs[,1])
lower_ci <- exp(coefs[,1] - 1.96 * coefs[,2])
upper_ci <- exp(coefs[,1] + 1.96 * coefs[,2])

# Calculate z-values and p-values
z_values <- coefs[,1] / coefs[,2]
p_values <- 2 * (1 - pnorm(abs(z_values)))

# Combine everything into a data frame for easy viewing
```

```
results <- data.frame(OR = OR, LowerCI = lower_ci, UpperCI = upper_ci, p_value = p_values)
print(results)
```

```
##              OR    LowerCI    UpperCI    p_value
## 1|2          0.6532438  0.432356   0.9869818 4.315240e-02
## 2|3          10.0537125  5.590398  18.0804911 1.287859e-14
## 3|4          52.1239514 24.658787 110.1800485 0.000000e+00
## NewDASNot Shared 3.2602625 1.392494   7.6332896 6.472506e-03
## NewDASShared    13.1606441 6.295668  27.5113854 7.357448e-12
## PreprintsYes     2.2372515 1.138340   4.3970105 1.949855e-02
```

#Check the Assumptions

`nominal_test(m1a)` *# This is a test of the proportional odds assumption, odds are proportional in this c*

```
## Tests of nominal effects
##
## formula: Complete ~ NewDAS + Preprints
##          Df logLik   AIC   LRT Pr(>Chi)
## <none>      -207.28 426.57
## NewDAS     4 -202.41 424.82 9.7501  0.04485 *
## Preprints  2 -207.14 430.28 0.2913  0.86448
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

`scale_test(m1a)` *# scale test checks for equal variance/ scale across year, assumptions are not violated*

```
## Tests of scale effects
##
## formula: Complete ~ NewDAS + Preprints
##          Df logLik   AIC   LRT Pr(>Chi)
## <none>      -207.28 426.57
## NewDAS     2 -203.91 423.83 6.7398  0.03439 *
## Preprints  1 -207.28 428.56 0.0031  0.95535
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

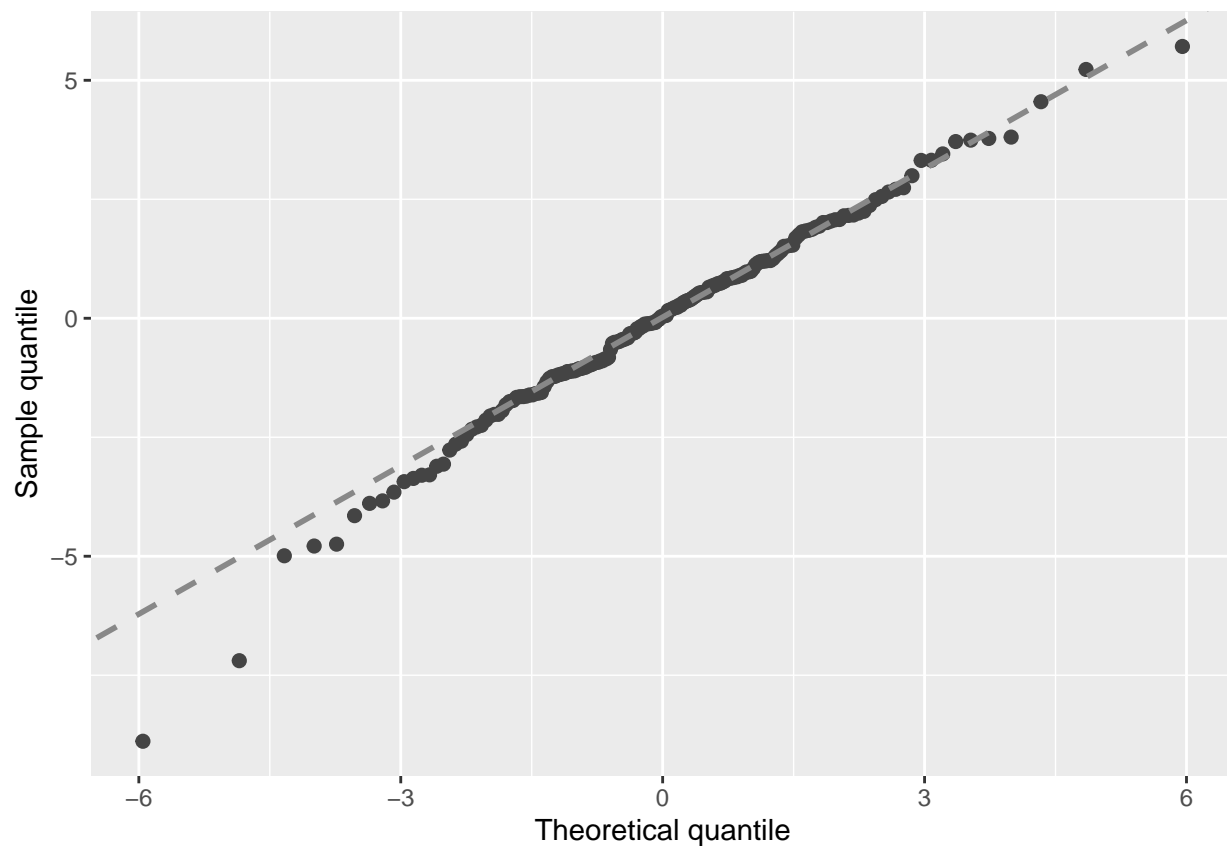
`convergence(m1a)` *# This is another way to assess the model*

```
## nobs logLik  niter max.grad cond.H  logLik.Error
## 193 -207.28 6(0)  3.61e-12 1.9e+01 <1e-10
##
##          Estimate Std.Err  Gradient    Error Cor.Dec Sig.Dig
## 1|2          -0.4258  0.2106  3.09e-12  1.17e-14     13     13
## 2|3           2.3079  0.2994 -3.61e-12 -2.81e-13     12     13
## 3|4           3.9536  0.3819 -3.23e-14 -2.81e-13     12     13
## NewDASNot Shared 1.1818  0.4340 -3.21e-13 -2.26e-13     12     13
## NewDASShared    2.5772  0.3762 -3.15e-13 -2.72e-13     12     13
## PreprintsYes     0.8052  0.3447  2.07e-14 -2.27e-14     13     13
##
## Eigen values of Hessian:
## 48.919 34.721 28.580  9.736  6.586  2.565
```

```
##
## Convergence message from clm:
## (0) successful convergence
## In addition: Absolute and relative convergence criteria were met
```

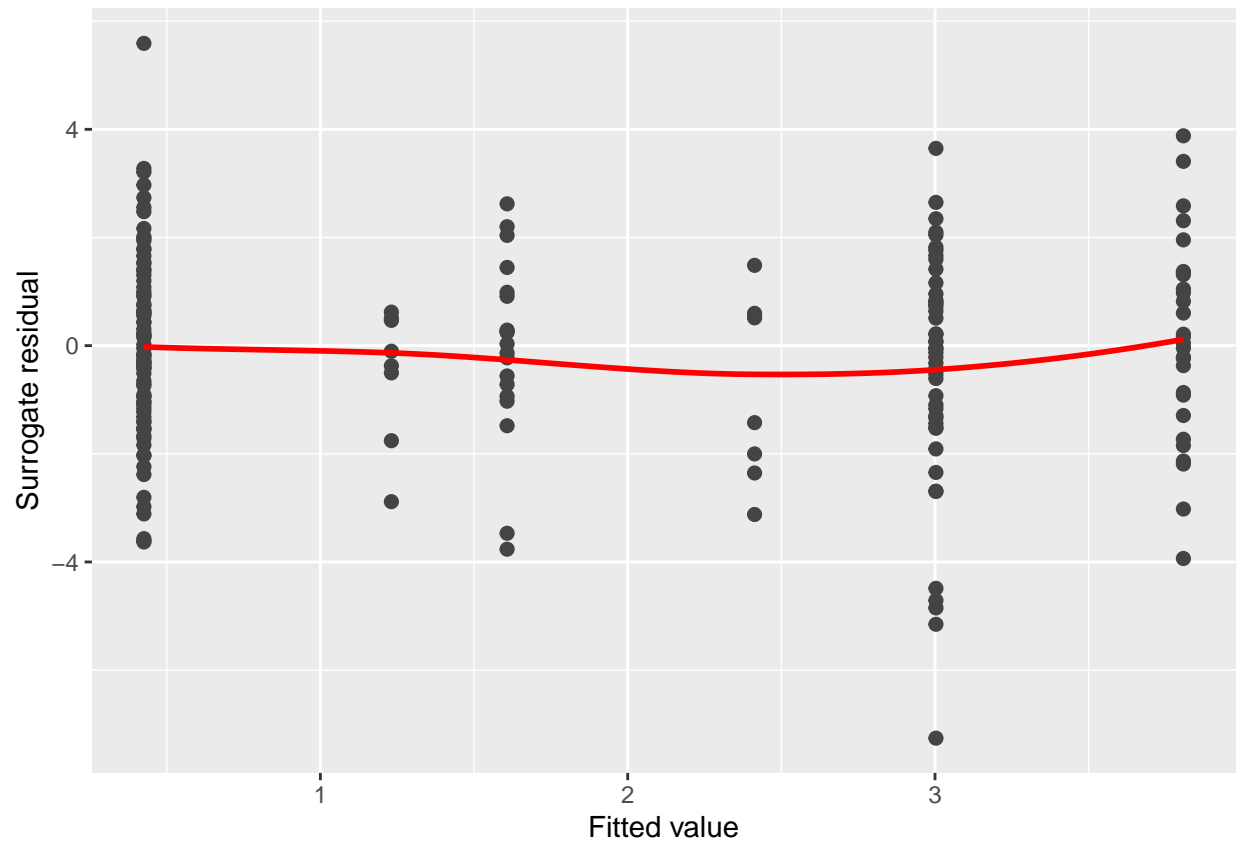
```
##### Graphically validate proportional odds using the sure package #####
```

```
autoplot.clm(m1a, what = c("qq")) # most of the points fall along the line, so no violation of linearity
```



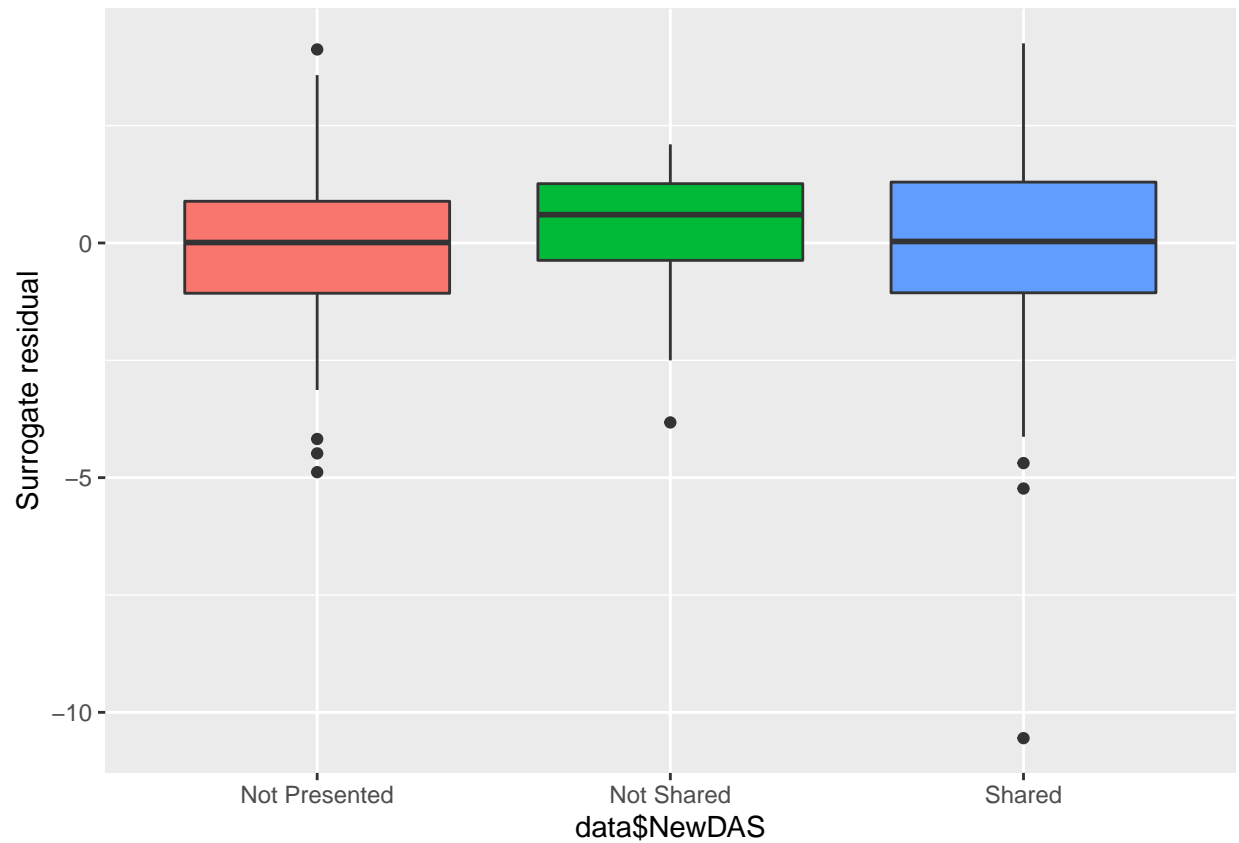
```
autoplot.clm(m1a, what = c("fitted")) # the residuals do not show a clear pattern or trend
```

```
## 'geom_smooth()' using method = 'loess' and formula 'y ~ x'
```

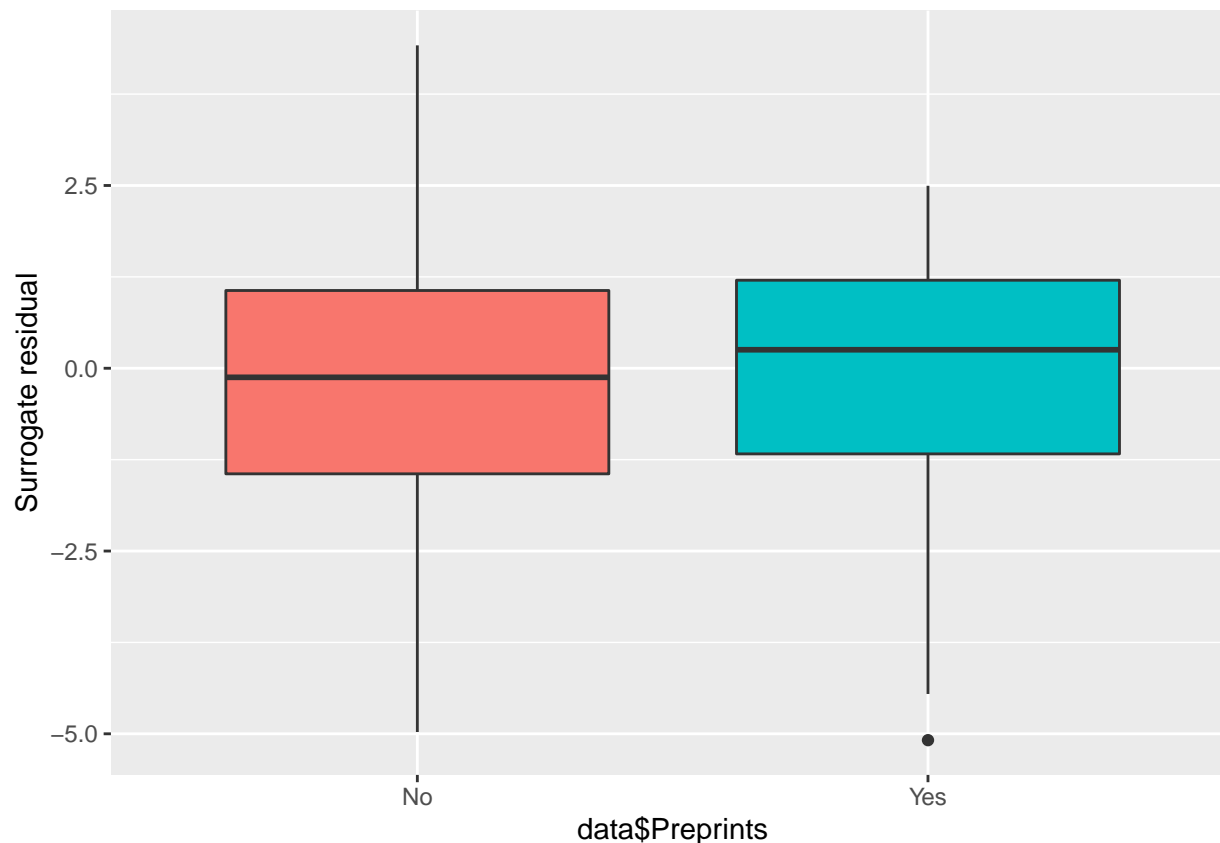
```
autoplot.clm(m1a, what = c("covariate"), x = data$NewDAS) # scale test indicated no variance issues. Th
```

```
## Warning: 'guides(<scale> = FALSE)' is deprecated. Please use 'guides(<scale> =  
## "none")' instead.
```



```
autoplot.clm(m1a, what = c("covariate"), x = data$Preprints)
```

```
## Warning: 'guides(<scale> = FALSE)' is deprecated. Please use 'guides(<scale> =  
## "none")' instead.
```



N.B. - Interpreting residual plots is largely subjective!

#these results were used in the Fifth result paragraph and Table 2 of the publication

```
m1a <- clm(Reuse ~ NewDAS + Preprints, data = data)
summary_m1a <- summary(m1a)

# Extract coefficients and standard errors
coefs <- summary_m1a$coefficients[, 1] # Coefficients
se_coefs <- summary_m1a$coefficients[, 2] # Standard errors

# Extract coefficients and standard errors
coefs <- summary(m1a)$coefficients
OR <- exp(coefs[,1])
lower_ci <- exp(coefs[,1] - 1.96 * coefs[,2])
upper_ci <- exp(coefs[,1] + 1.96 * coefs[,2])

# Calculate z-values and p-values
z_values <- coefs[,1] / coefs[,2]
p_values <- 2 * (1 - pnorm(abs(z_values)))

# Combine everything into a data frame for easy viewing
results <- data.frame(OR = OR, LowerCI = lower_ci, UpperCI = upper_ci, p_value = p_values)
print(results)
```

```
##           OR   LowerCI   UpperCI   p_value
## 1|2       0.5851276 0.3875115 0.8835203 1.080242e-02
## 2|3       1.1384211 0.7593092 1.7068181 5.303823e-01
## 3|4       4.4224283 2.7525267 7.1054250 7.981718e-10
## NewDASNot Shared 5.5189574 2.3571077 12.9221468 8.306053e-05
## NewDASShared    6.7456538 3.5058381 12.9794487 1.085746e-08
## PreprintsYes    1.0835906 0.5520896 2.1267718 8.154950e-01
```

#Check the Assumptions

```
nominal_test(m1a) # This is a test of the proportional odds assumption, odds are proportional in this c
```

```
## Tests of nominal effects
```

```
##
```

```
## formula: Reuse ~ NewDAS + Preprints
```

```
##           Df logLik   AIC   LRT Pr(>Chi)
```

```
## <none>      -226.59 465.19
```

```
## NewDAS      4 -223.98 467.96 5.2220 0.26527
```

```
## Preprints   2 -222.23 460.46 8.7302 0.01271 *
```

```
## ---
```

```
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
scale_test(m1a) # scale test checks for equal variance/ scale across year, assumptions are not violated
```

```
## Tests of scale effects
```

```
##
```

```
## formula: Reuse ~ NewDAS + Preprints
```

```
##           Df logLik   AIC   LRT Pr(>Chi)
```

```
## <none>      -226.59 465.19
```

```
## NewDAS      2 -226.39 468.78 0.41107 0.8142
```

```
## Preprints   1 -225.55 465.11 2.07927 0.1493
```

```
convergence(m1a) # This is another way to assess the model
```

```
## nobs logLik niter max.grad cond.H logLik.Error
```

```
## 193 -226.59 6(0) 1.80e-13 3.3e+01 <1e-10
```

```
##
```

```
##           Estimate Std.Err Gradient Error Cor.Dec Sig.Dig
```

```
## 1|2       -0.53593 0.2102 1.80e-13 2.64e-15 14 14
```

```
## 2|3       0.12964 0.2066 -1.56e-13 -5.47e-16 14 14
```

```
## 3|4       1.48669 0.2419 6.55e-15 -3.82e-16 15 16
```

```
## NewDASNot Shared 1.70819 0.4341 -9.99e-16 -5.94e-16 14 15
```

```
## NewDASShared    1.90890 0.3339 -7.88e-15 -1.01e-15 14 15
```

```
## PreprintsYes    0.08028 0.3440 2.89e-15 7.09e-16 14 13
```

```
##
```

```
## Eigen values of Hessian:
```

```
## 123.454 60.436 27.013 10.229 7.437 3.706
```

```
##
```

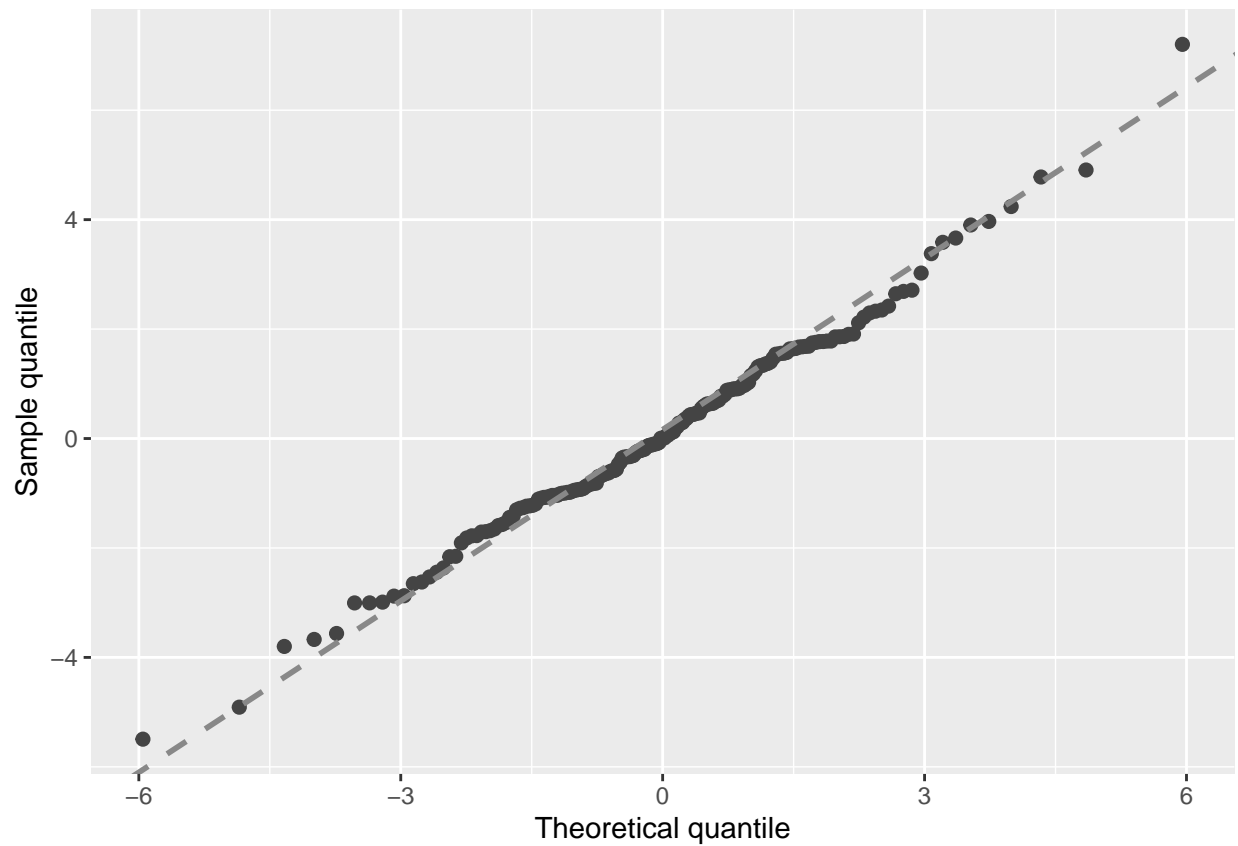
```
## Convergence message from clm:
```

```
## (0) successful convergence
```

```
## In addition: Absolute and relative convergence criteria were met
```

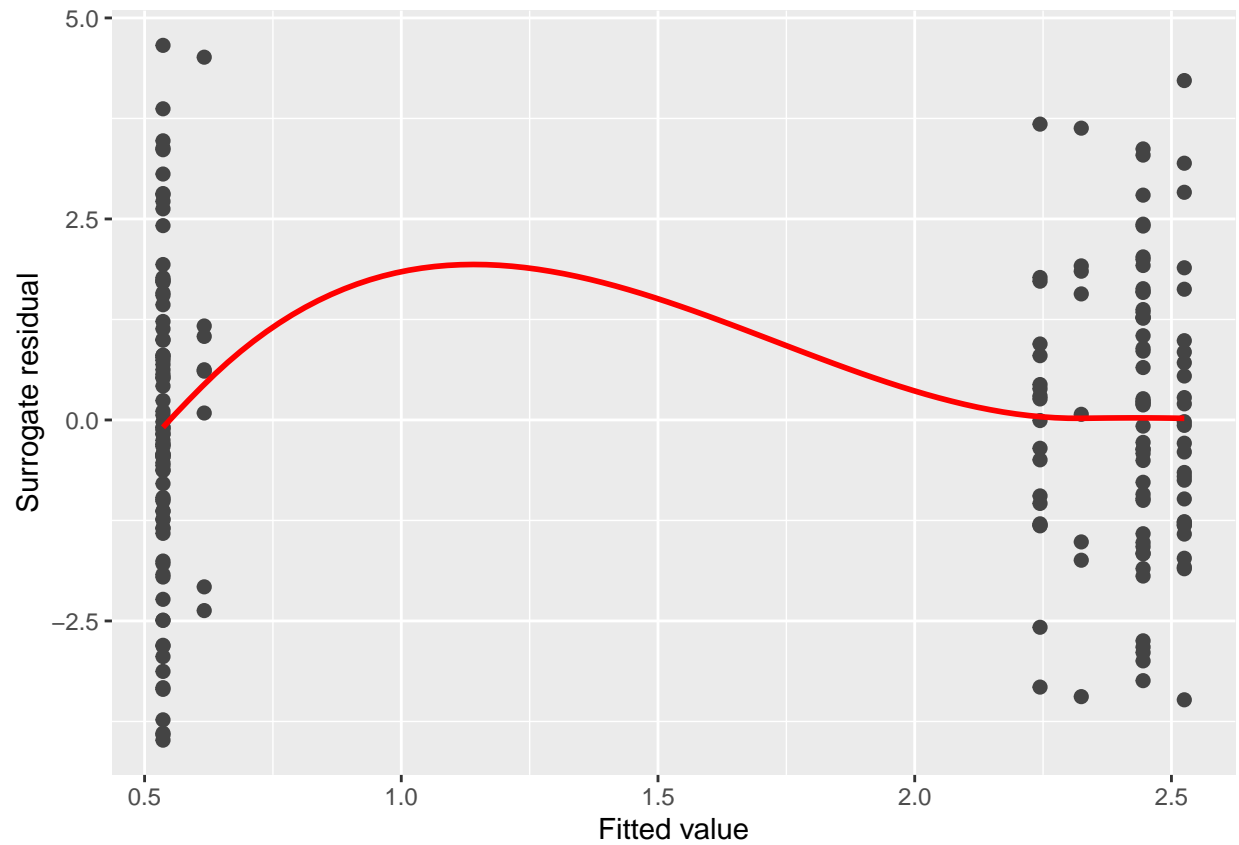
```
##### Graphically validate proportional odds using the sure package #####
```

```
autoplot.clm(m1a, what = c("qq")) # most of the points fall along the line, so no violation of linearity
```



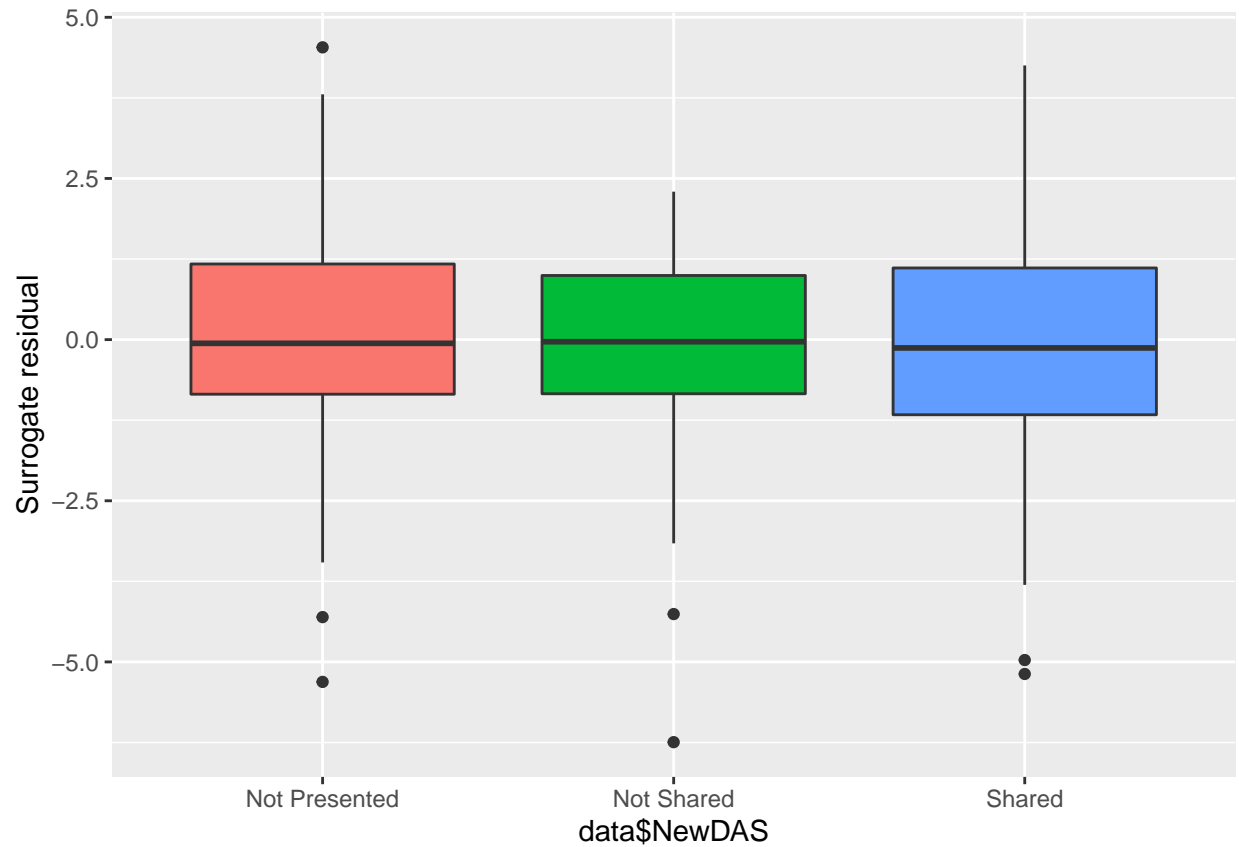
```
autoplot.clm(m1a, what = c("fitted")) # the residuals do not show a clear pattern or trend
```

```
## 'geom_smooth()' using method = 'loess' and formula 'y ~ x'
```



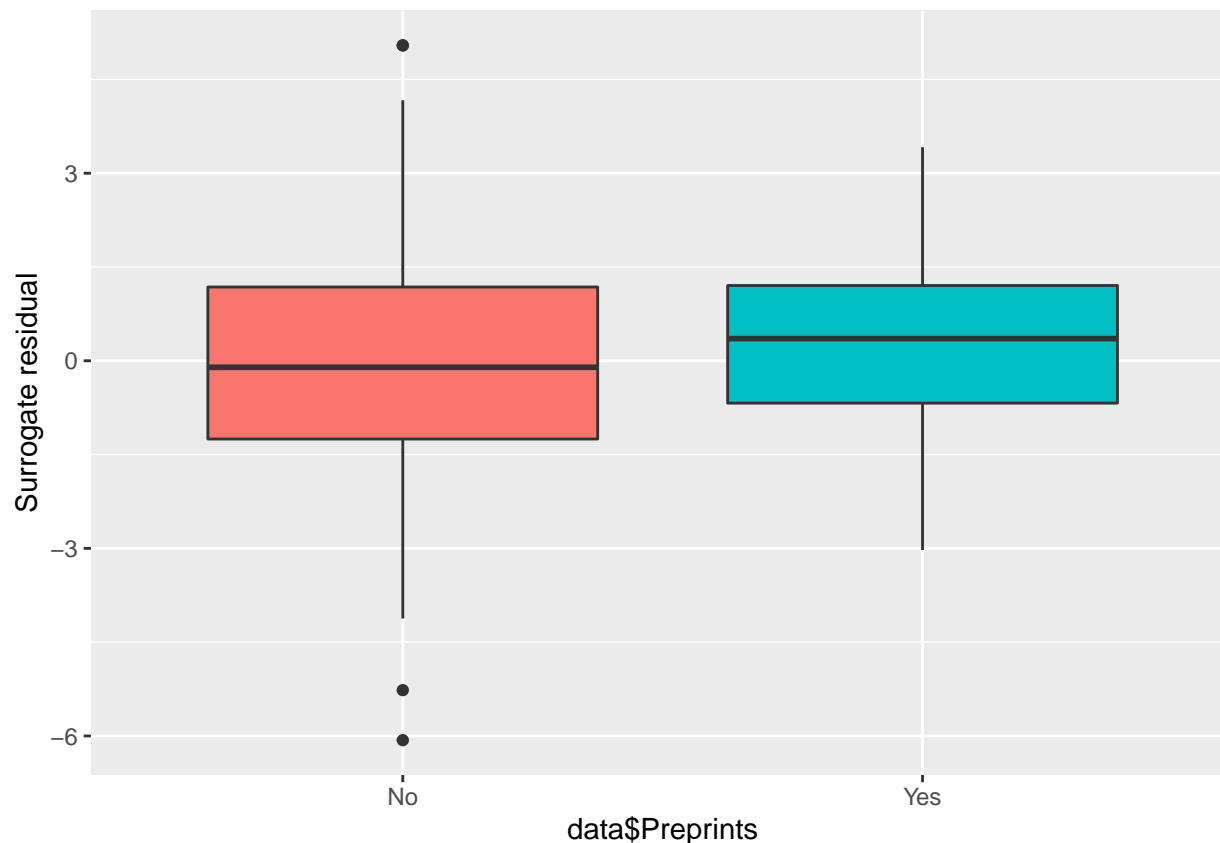
```
autoplot.clm(m1a, what = c("covariate"), x = data$NewDAS) # scale test indicated no variance issues. Th
```

```
## Warning: 'guides(<scale> = FALSE)' is deprecated. Please use 'guides(<scale> =  
## "none")' instead.
```



```
autoplot.clm(m1a, what = c("covariate"), x = data$Preprints)
```

```
## Warning: 'guides(<scale> = FALSE)' is deprecated. Please use 'guides(<scale> =  
## "none")' instead.
```



N.B. - Interpreting residual plots is largely subjective!

#these results were used in the Fifth result paragraph and Table 2 of the publication

```
m1a <- clm(Access ~ NewDAS + Preprints, data = data)
summary_m1a <- summary(m1a)

# Extract coefficients and standard errors
coefs <- summary_m1a$coefficients[, 1] # Coefficients
se_coefs <- summary_m1a$coefficients[, 2] # Standard errors

# Extract coefficients and standard errors
coefs <- summary(m1a)$coefficients
OR <- exp(coefs[,1])
lower_ci <- exp(coefs[,1] - 1.96 * coefs[,2])
upper_ci <- exp(coefs[,1] + 1.96 * coefs[,2])

# Calculate z-values and p-values
z_values <- coefs[,1] / coefs[,2]
p_values <- 2 * (1 - pnorm(abs(z_values)))

# Combine everything into a data frame for easy viewing
results <- data.frame(OR = OR, LowerCI = lower_ci, UpperCI = upper_ci, p_value = p_values)
print(results)
```

##	OR	LowerCI	UpperCI	p_value
----	----	---------	---------	---------


```
## 1|2          0.6034796 0.3984970 0.9139029 1.707011e-02
## 2|3          2.3011558 1.4917473 3.5497420 1.642623e-04
## 3|4          2.8320314 1.8164013 4.4155452 4.349083e-06
## NewDASNot Shared 5.0164438 2.0473496 12.2913588 4.200776e-04
## NewDASShared    5.8615538 3.0241489 11.3611511 1.627745e-07
## PreprintsYes    1.6652166 0.7931505 3.4961165 1.777868e-01
```

#Check the Assumptions

`nominal_test(m1a)` *# This is a test of the proportional odds assumption, odds are proportional in this c*

```
## Tests of nominal effects
##
## formula: Access ~ NewDAS + Preprints
##          Df logLik   AIC   LRT Pr(>Chi)
## <none>      -200.61 413.22
## NewDAS     4 -197.91 415.82 5.4031 0.2484
## Preprints  2 -199.89 415.78 1.4450 0.4855
```

`scale_test(m1a)` *# scale test checks for equal variance/ scale across year, assumptions are not violated*

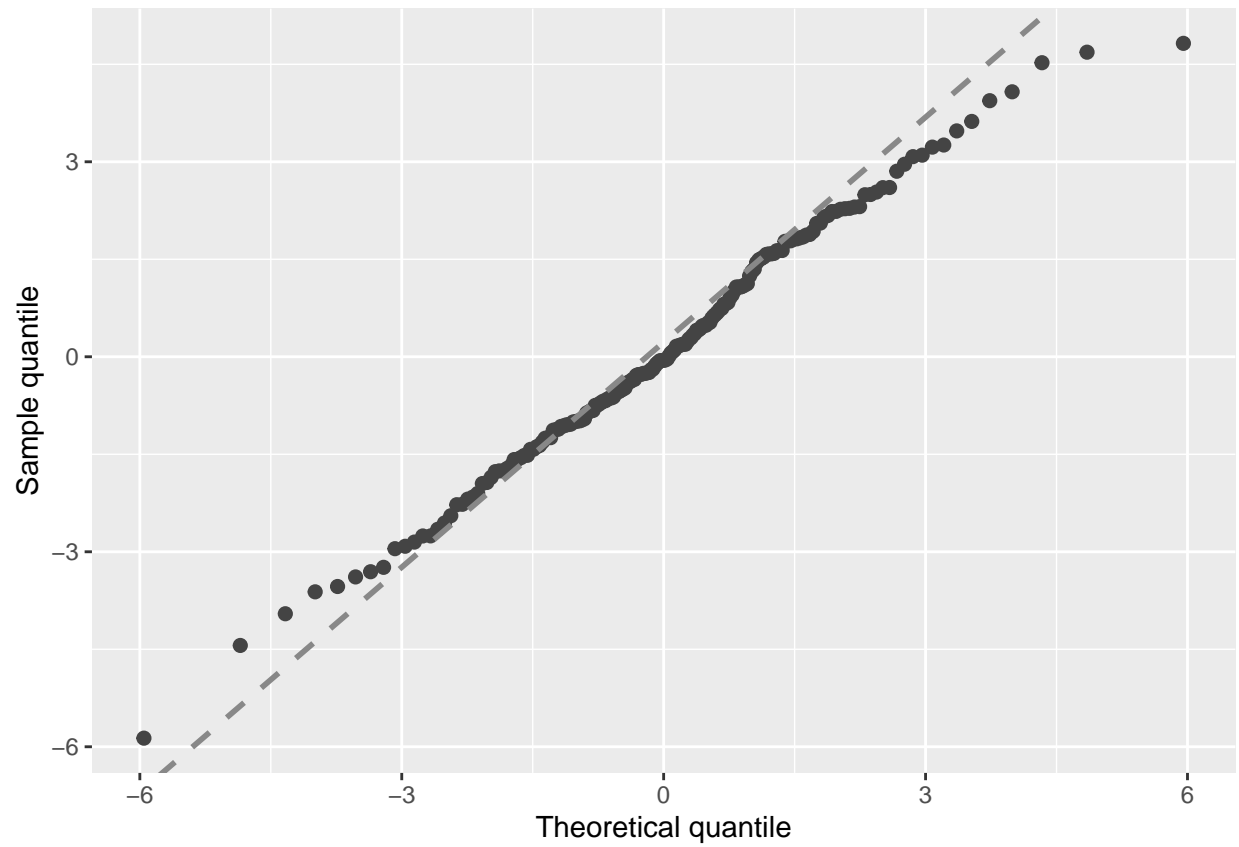
```
## Tests of scale effects
##
## formula: Access ~ NewDAS + Preprints
##          Df logLik   AIC   LRT Pr(>Chi)
## <none>      -200.61 413.22
## NewDAS     2 -200.39 416.78 0.44455 0.8007
## Preprints  1 -200.24 414.47 0.75425 0.3851
```

`convergence(m1a)` *# This is another way to assess the model*

```
## nobs logLik niter max.grad cond.H logLik.Error
## 193 -200.61 6(2) 2.14e-12 1.1e+02 <1e-10
##
##          Estimate Std.Err Gradient Error Cor.Dec Sig.Dig
## 1|2          -0.5050 0.2117 5.49e-13 -1.28e-14 13 13
## 2|3           0.8334 0.2212 1.39e-13 -2.88e-14 13 13
## 3|4           1.0410 0.2266 1.61e-12 -2.95e-14 13 14
## NewDASNot Shared 1.6127 0.4572 -3.03e-13 -2.78e-14 13 14
## NewDASShared    1.7684 0.3376 -1.99e-12 -8.64e-14 12 13
## PreprintsYes    0.5100 0.3784 -2.14e-12 -2.06e-13 12 12
##
## Eigen values of Hessian:
## 398.229 50.296 27.709 8.866 6.325 3.547
##
## Convergence message from clm:
## (0) successful convergence
## In addition: Absolute and relative convergence criteria were met
```

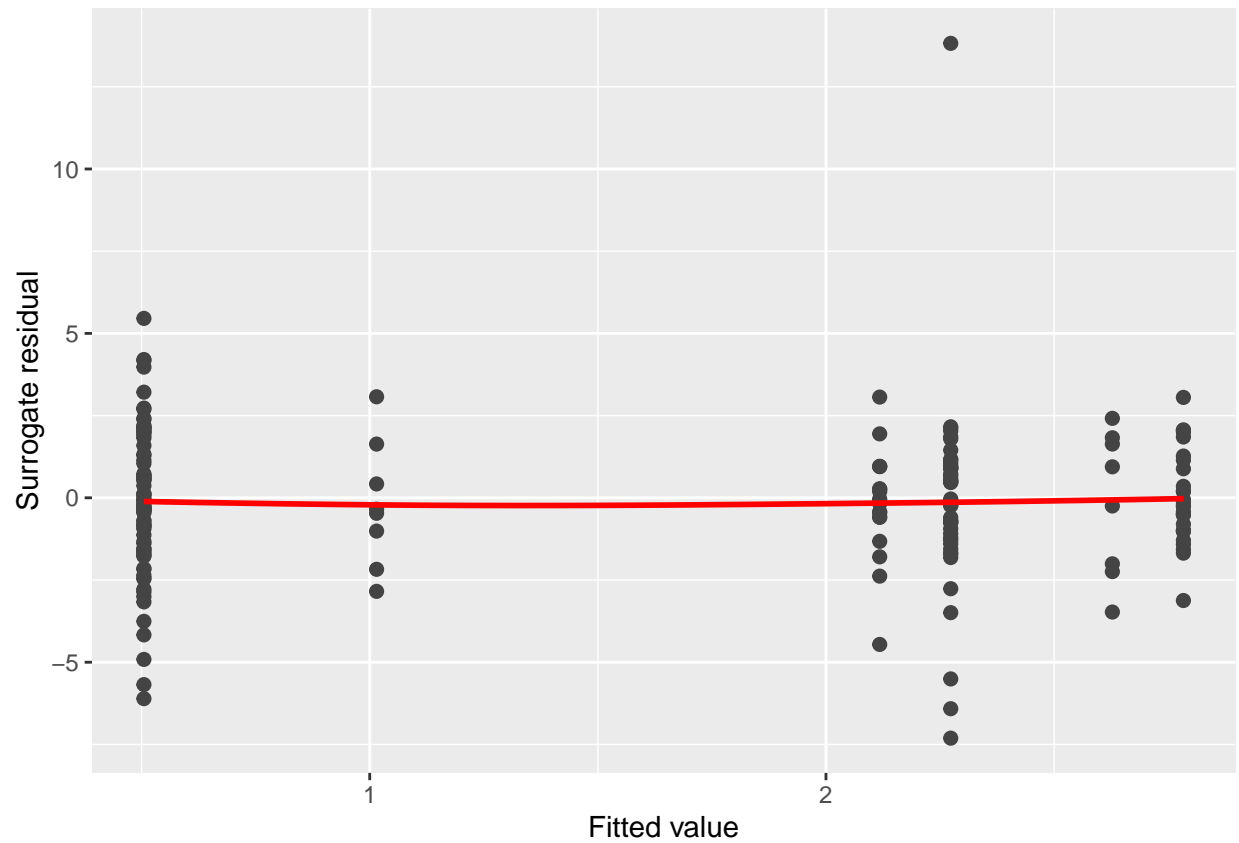
Graphically validate proportional odds using the sure package

`autoplot.clm(m1a, what = c("qq"))` *# most of the points fall along the line, so no violation of linearit*



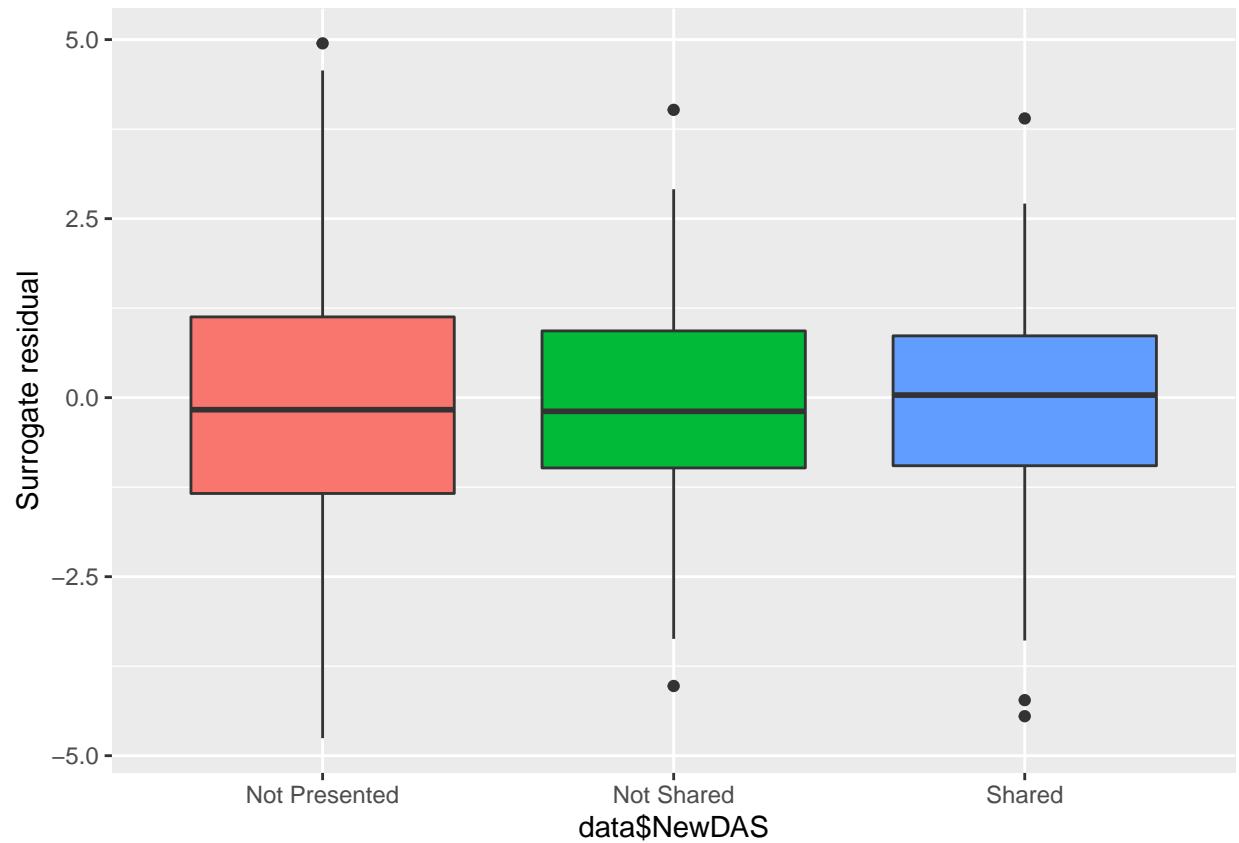
```
autoplot.clm(m1a, what = c("fitted")) # the residuals do not show a clear pattern or trend
```

```
## 'geom_smooth()' using method = 'loess' and formula 'y ~ x'
```



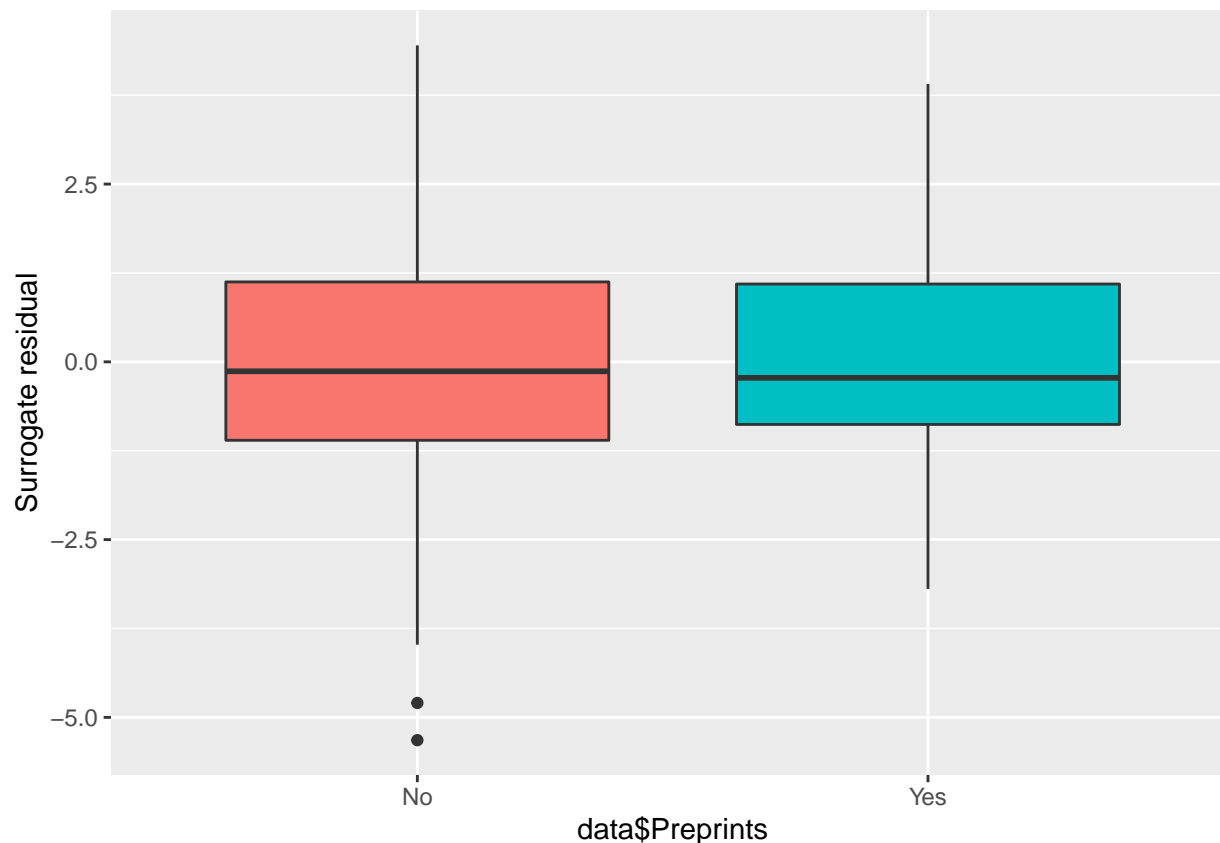
```
autoplot.clm(m1a, what = c("covariate"), x = data$NewDAS) # scale test indicated no variance issues. Th
```

```
## Warning: 'guides(<scale> = FALSE)' is deprecated. Please use 'guides(<scale> =  
## "none")' instead.
```



```
autoplot.clm(m1a, what = c("covariate"), x = data$Preprints)
```

```
## Warning: 'guides(<scale> = FALSE)' is deprecated. Please use 'guides(<scale> =  
## "none")' instead.
```



N.B. - Interpreting residual plots is largely subjective!

#these results were used in the Fifth result paragraph and Table 2 of the publication

```
m1a <- clm(Licence ~ NewDAS + Preprints, data = data)
summary_m1a <- summary(m1a)

# Extract coefficients and standard errors
coefs <- summary_m1a$coefficients[, 1] # Coefficients
se_coefs <- summary_m1a$coefficients[, 2] # Standard errors

# Extract coefficients and standard errors
coefs <- summary(m1a)$coefficients
OR <- exp(coefs[,1])
lower_ci <- exp(coefs[,1] - 1.96 * coefs[,2])
upper_ci <- exp(coefs[,1] + 1.96 * coefs[,2])

# Calculate z-values and p-values
z_values <- coefs[,1] / coefs[,2]
p_values <- 2 * (1 - pnorm(abs(z_values)))

# Combine everything into a data frame for easy viewing
results <- data.frame(OR = OR, LowerCI = lower_ci, UpperCI = upper_ci, p_value = p_values)
print(results)
```

##	OR	LowerCI	UpperCI	p_value
----	----	---------	---------	---------

```
## 1|2          0.6287768 0.4134465 0.9562549 3.007374e-02
## 2|3          0.7442143 0.4924399 1.1247157 1.608644e-01
## 3|4          1.1535025 0.7648622 1.7396180 4.957229e-01
## NewDASNot Shared 5.3128274 1.6908919 16.6930451 4.246679e-03
## NewDASShared    6.3953648 2.8022381 14.5957227 1.045395e-05
## PreprintsYes    2.3785527 0.8260296 6.8490444 1.083171e-01
```

#Check the Assumptions

`nominal_test(m1a)` *# This is a test of the proportional odds assumption, odds are proportional in this c*

```
## Tests of nominal effects
##
## formula: Licence ~ NewDAS + Preprints
##           Df logLik   AIC LRT Pr(>Chi)
## <none>      -149.58 311.16
## NewDAS
## Preprints
```

`scale_test(m1a)` *# scale test checks for equal variance/ scale across year, assumptions are not violated*

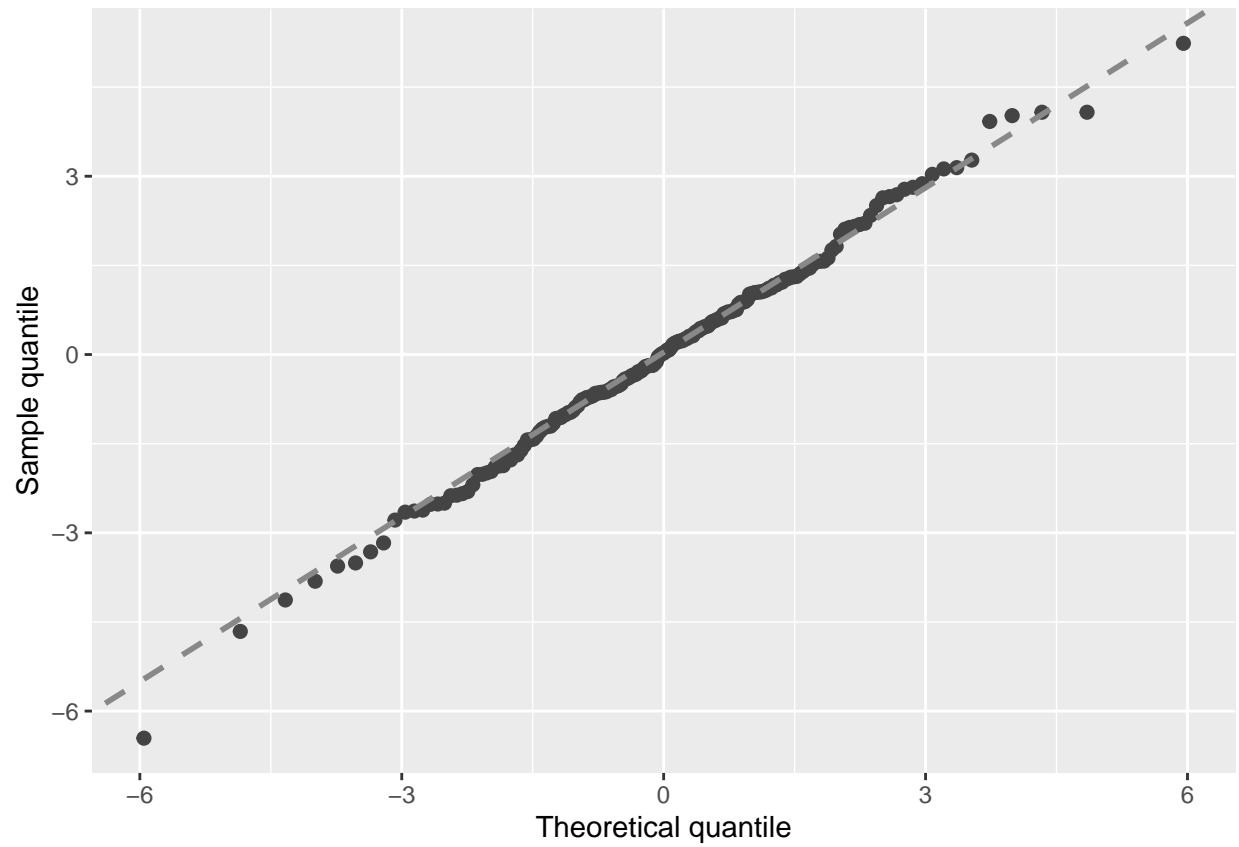
```
## Tests of scale effects
##
## formula: Licence ~ NewDAS + Preprints
##           Df logLik   AIC   LRT Pr(>Chi)
## <none>      -149.58 311.16
## NewDAS      2 -148.95 313.91 1.25533 0.5338
## Preprints    1 -149.10 312.21 0.95565 0.3283
```

`convergence(m1a)` *# This is another way to assess the model*

```
## nobs logLik niter max.grad cond.H logLik.Error
## 193 -149.58 7(2) 5.84e-12 1.6e+02 <1e-10
##
##           Estimate Std.Err Gradient Error Cor.Dec Sig.Dig
## 1|2          -0.4640 0.2139 4.50e-13 -4.50e-14 13 13
## 2|3          -0.2954 0.2107 1.58e-13 -4.58e-14 13 13
## 3|4           0.1428 0.2096 5.71e-12 -5.06e-14 12 12
## NewDASNot Shared 1.6701 0.5841 -7.88e-13 -5.52e-14 12 13
## NewDASShared    1.8556 0.4210 -4.57e-12 -3.21e-13 12 13
## PreprintsYes    0.8665 0.5396 -5.84e-12 -1.33e-12 11 11
##
## Eigen values of Hessian:
## 406.815 113.328 15.099 5.486 3.398 2.567
##
## Convergence message from clm:
## (0) successful convergence
## In addition: Absolute and relative convergence criteria were met
```

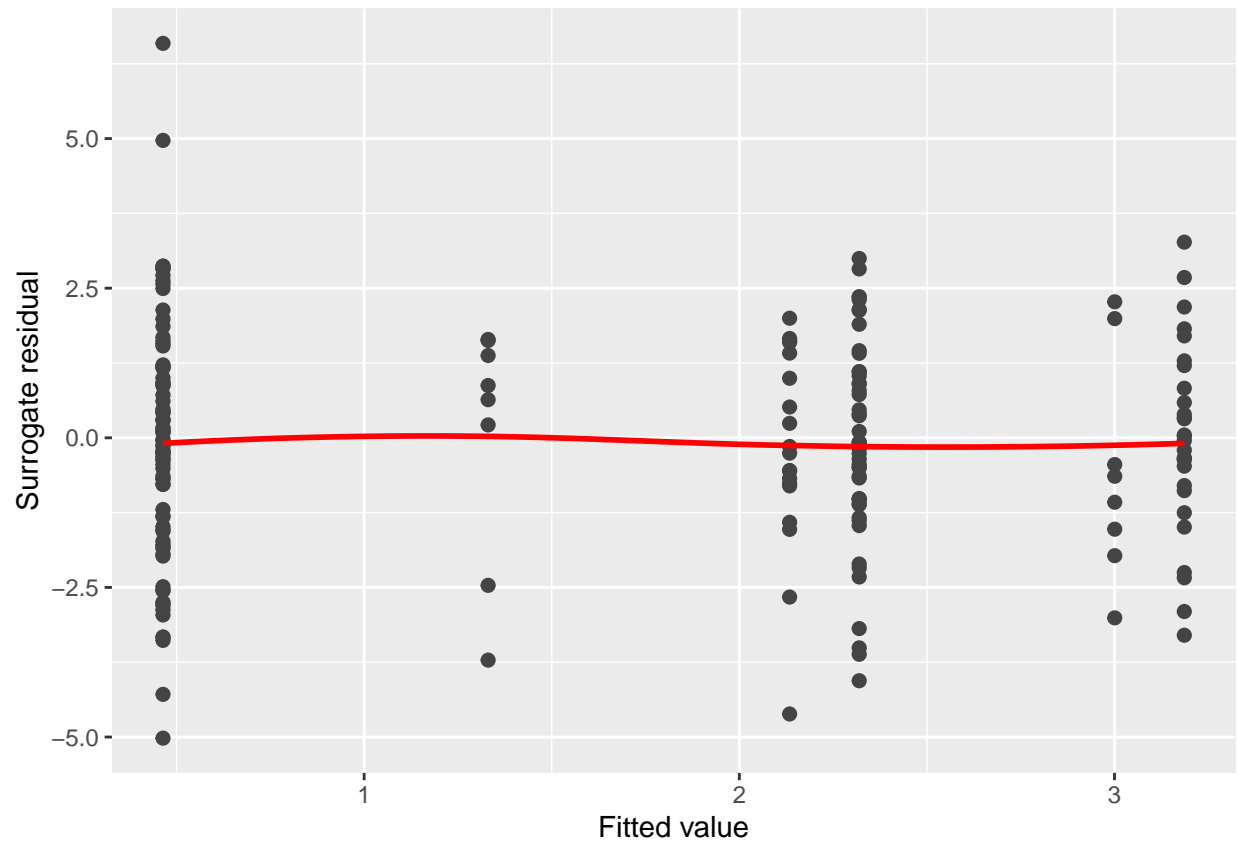
Graphically validate proportional odds using the sure package

`autoplot.clm(m1a, what = c("qq"))` *# most of the points fall along the line, so no violation of linearit*



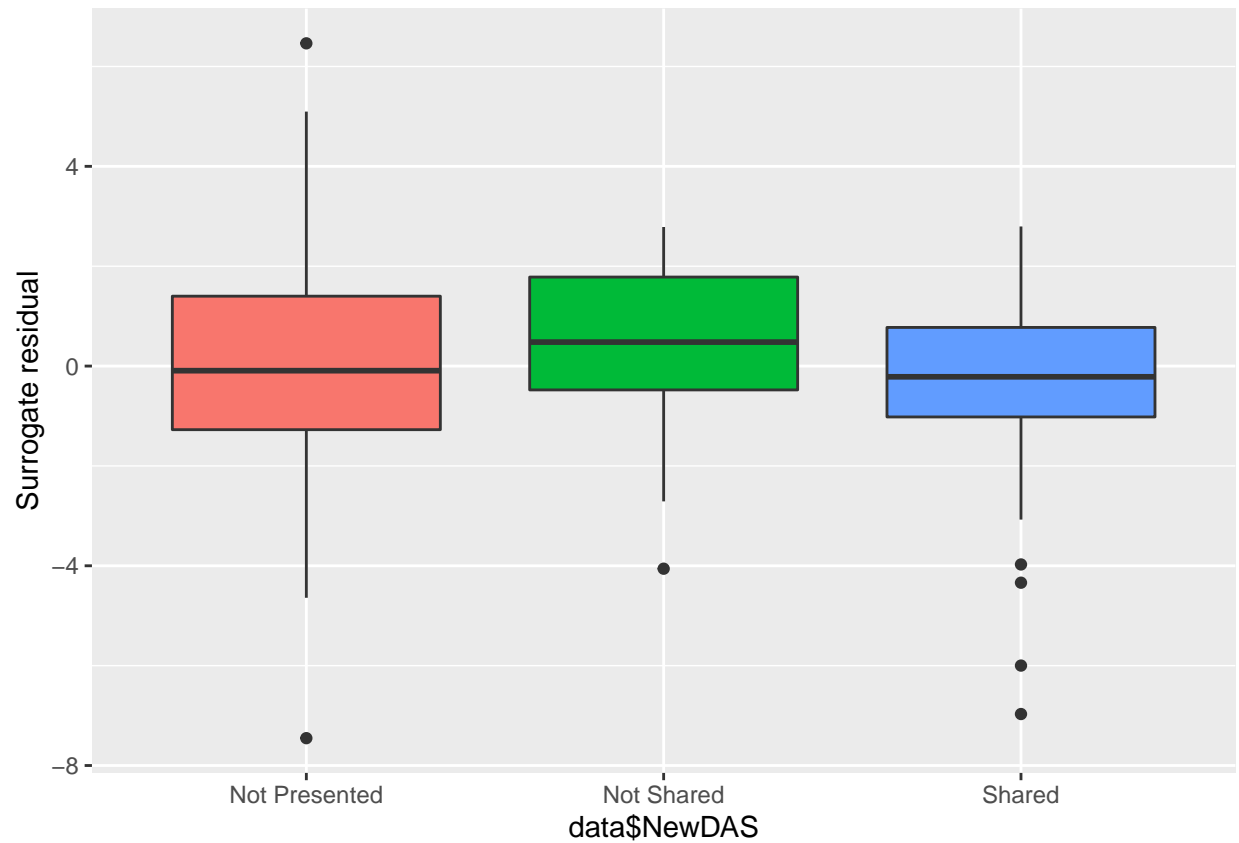
```
autoplot.clm(m1a, what = c("fitted")) # the residuals do not show a clear pattern or trend
```

```
## 'geom_smooth()' using method = 'loess' and formula 'y ~ x'
```



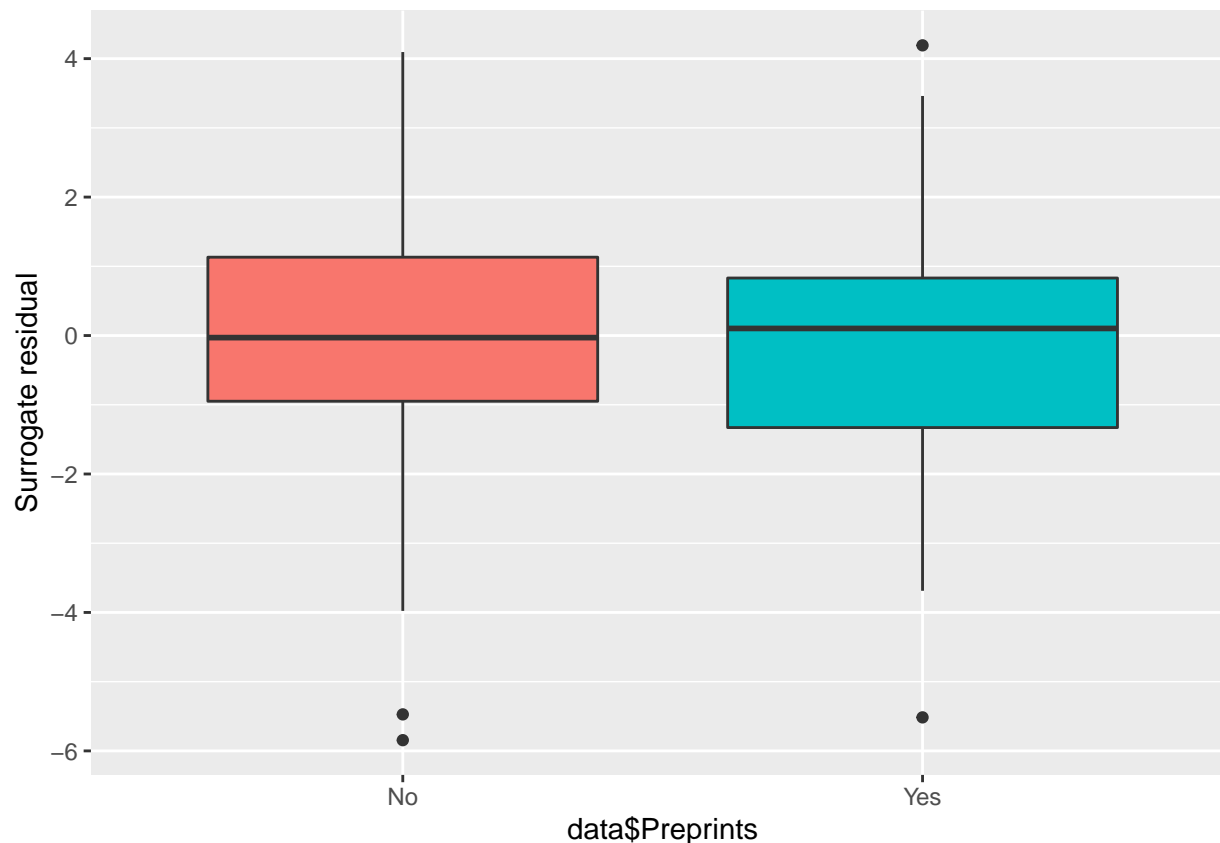
```
autoplot.clm(m1a, what = c("covariate"), x = data$NewDAS) # scale test indicated no variance issues. Th
```

```
## Warning: 'guides(<scale> = FALSE)' is deprecated. Please use 'guides(<scale> =  
## "none")' instead.
```

```
autoplot.clm(m1a, what = c("covariate"), x = data$Preprints)
```

```
## Warning: 'guides(<scale> = FALSE)' is deprecated. Please use 'guides(<scale> =  
## "none")' instead.
```



N.B. - Interpreting residual plots is largely subjective!

#these results were used in the Fifth result paragraph and Table 2 of the publication

#Research Group figures (Internal)

```
# Calculate the frequency and percentage for each 'Complete' score among the research Group
long_data <- data %>%
  count(ResGrp, Complete) %>%
  group_by(ResGrp) %>%
  mutate(Percentage = n / sum(n) * 100) %>%
  mutate(Label = ifelse(Percentage > 5, paste0(round(Percentage, 1), "%"), "")) %>%
  mutate(Cumulative_Percentage = cumsum(Percentage) - (0.5 * Percentage)) %>%
  ungroup()

long_data <- long_data %>%
  mutate(Complete = factor(Complete, levels = c("4", "3", "2", "1")))

# Custom colors
colors <- c("#D0A136", "#36D0A1", "#50689B", "#E5696F")

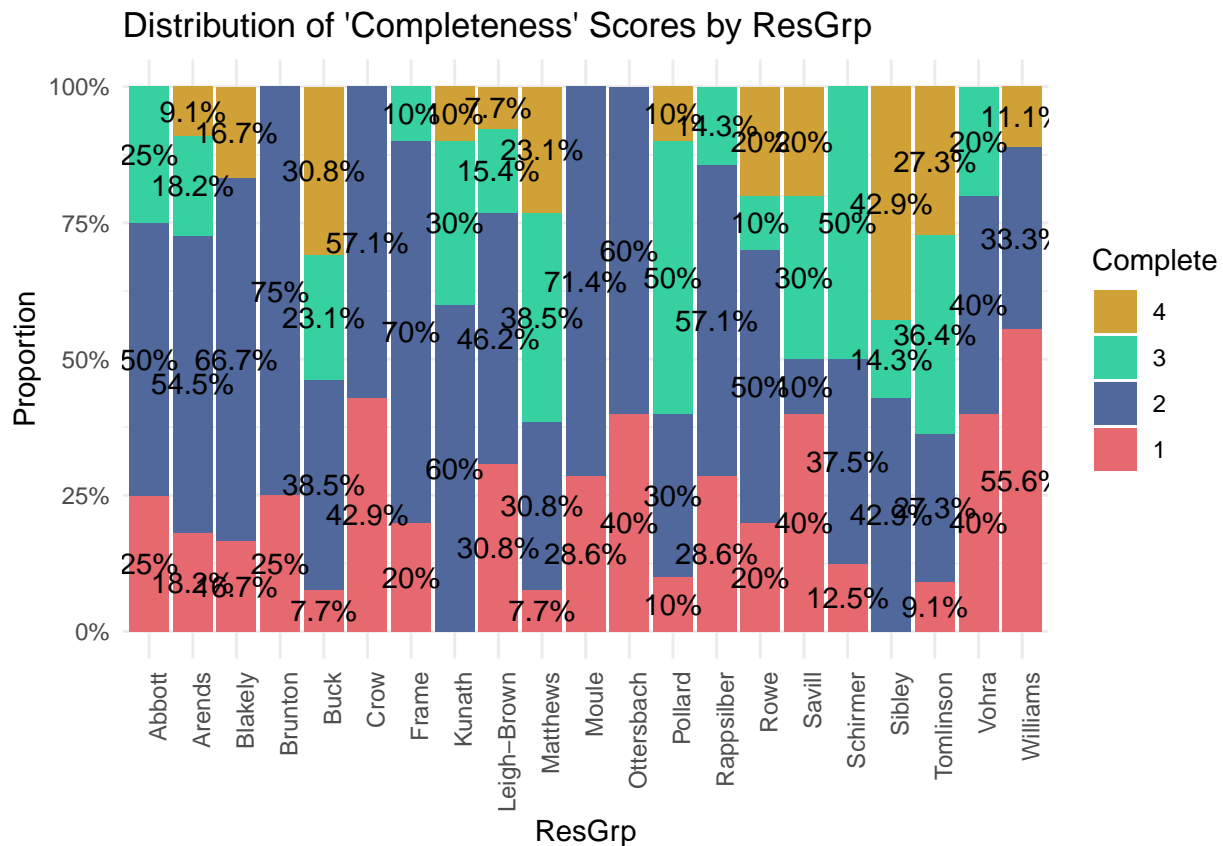
# Creating the stacked bar chart with percentage labels
internal1 <- ggplot(long_data, aes(x = as.factor(ResGrp), y = n, fill = as.factor(Complete))) +
  geom_bar(stat = "identity", position = "fill") + # Scale the bar heights to proportions
  scale_y_continuous(labels = percent_format()) + # Convert the y-axis to percentage
```

```

geom_text(
  aes(label = Label, y = Percentage),
  size = 4,
  color = "black",
  position = position_fill(vjust = 0.5)
) +
scale_fill_manual(values = colors) +
labs(title = "Distribution of 'Completeness' Scores by ResGrp",
     x = "ResGrp",
     y = "Proportion",
     fill = "Complete") +
theme_minimal() +
scale_x_discrete(name = "ResGrp", labels = unique(long_data$ResGrp)) +
theme(axis.text.x = element_text(angle = 90, hjust = 1)) # Adjust text angle for readability

# Print the plot
print(internal1)

```



```

ggsave("internal1.png", internal1, width = 15, height = 10, units = "in", bg= "white")

```

#these results were used for internal university meetings and presentations

```

# Calculate the frequency and percentage for each 'Reuse' score among the research Group
long_data <- data %>%
  count(ResGrp, Reuse) %>%
  group_by(ResGrp) %>%
  mutate(Percentage = n / sum(n) * 100) %>%
  mutate(Label = ifelse(Percentage > 5, paste0(round(Percentage, 1), "%"), "")) %>%
  mutate(Cumulative_Percentage = cumsum(Percentage) - (0.5 * Percentage)) %>%
  ungroup()

long_data <- long_data %>%
  mutate(Reuse = factor(Reuse, levels = c("4", "3", "2", "1")))

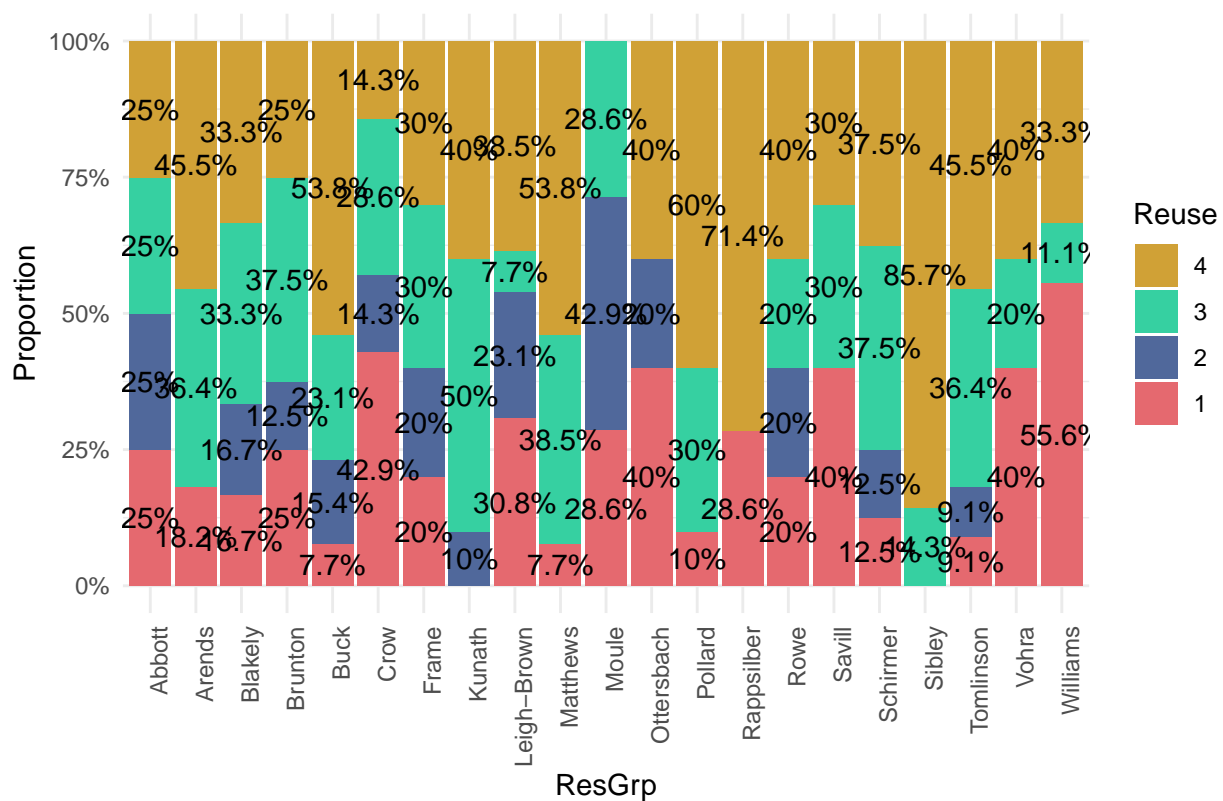
# Custom colors
colors <- c("#D0A136", "#36D0A1", "#50689B", "#E5696F")

# Creating the stacked bar chart with percentage labels
internal2 <- ggplot(long_data, aes(x = as.factor(ResGrp), y = n, fill = as.factor(Reuse))) +
  geom_bar(stat = "identity", position = "fill") + # Scale the bar heights to proportions
  scale_y_continuous(labels = percent_format()) + # Convert the y-axis to percentage
  geom_text(
    aes(label = Label, y = Percentage),
    size = 4,
    color = "black",
    position = position_fill(vjust = 0.5)
  ) +
  scale_fill_manual(values = colors) +
  labs(title = "Distribution of 'Reusability' Scores by ResGrp",
    x = "ResGrp",
    y = "Proportion",
    fill = "Reuse") +
  theme_minimal() +
  scale_x_discrete(name = "ResGrp", labels = unique(long_data$ResGrp)) +
  theme(axis.text.x = element_text(angle = 90, hjust = 1)) # Adjust text angle for readability

# Print the plot
print(internal2)

```

Distribution of 'Reusability' Scores by ResGrp



```
ggsave("internal2.png", internal2, width = 15, height = 10, units = "in", bg= "white")
```

#these results were used for internal university meetings and presentations

```
# Calculate the frequency and percentage for each 'Access' score among the research Group
long_data <- data %>%
  count(ResGrp, Access) %>%
  group_by(ResGrp) %>%
  mutate(Percentage = n / sum(n) * 100) %>%
  mutate(Label = ifelse(Percentage > 5, paste0(round(Percentage, 1), "%"), "")) %>%
  mutate(Cumulative_Percentage = cumsum(Percentage) - (0.5 * Percentage)) %>%
  ungroup()
```

```
long_data <- long_data %>%
  mutate(Access = factor(Access, levels = c("4", "3", "2", "1")))
```

Custom colors

```
colors <- c("#D0A136", "#36D0A1", "#50689B", "#E5696F")
```

Creating the stacked bar chart with percentage labels

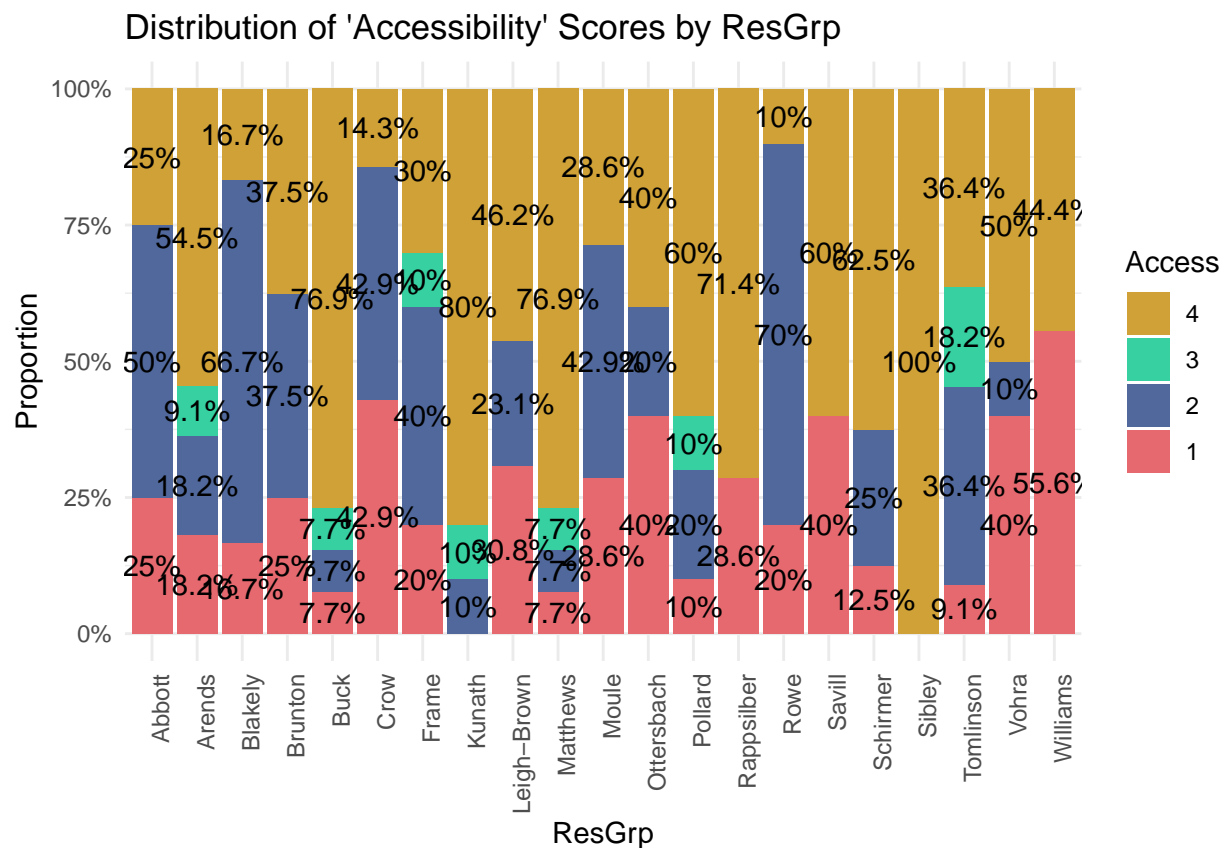
```
internal3 <- ggplot(long_data, aes(x = as.factor(ResGrp), y = n, fill = as.factor(Access))) +
  geom_bar(stat = "identity", position = "fill") + # Scale the bar heights to proportions
  scale_y_continuous(labels = percent_format()) + # Convert the y-axis to percentage
  geom_text(
    aes(label = Label, y = Percentage),
    size = 4,
```

```

    color = "black",
    position = position_fill(vjust = 0.5)
  ) +
  scale_fill_manual(values = colors) +
  labs(title = "Distribution of 'Accessibility' Scores by ResGrp",
       x = "ResGrp",
       y = "Proportion",
       fill = "Access") +
  theme_minimal() +
  scale_x_discrete(name = "ResGrp", labels = unique(long_data$ResGrp)) +
  theme(axis.text.x = element_text(angle = 90, hjust = 1)) # Adjust text angle for readability

# Print the plot
print(internal3)

```



```

ggsave("internal3.png", internal3, width = 15, height = 10, units = "in", bg= "white")

```

#these results were used for internal university meetings and presentations

```

# Calculate the frequency and percentage for each 'Licence' score among the research Group
long_data <- data %>%
  count(ResGrp, Licence) %>%
  group_by(ResGrp) %>%
  mutate(Percentage = n / sum(n) * 100) %>%

```

```

mutate(Label = ifelse(Percentage > 5, paste0(round(Percentage, 1), "%"), "") %>%
mutate(Cumulative_Percentage = cumsum(Percentage) - (0.5 * Percentage)) %>%
ungroup()

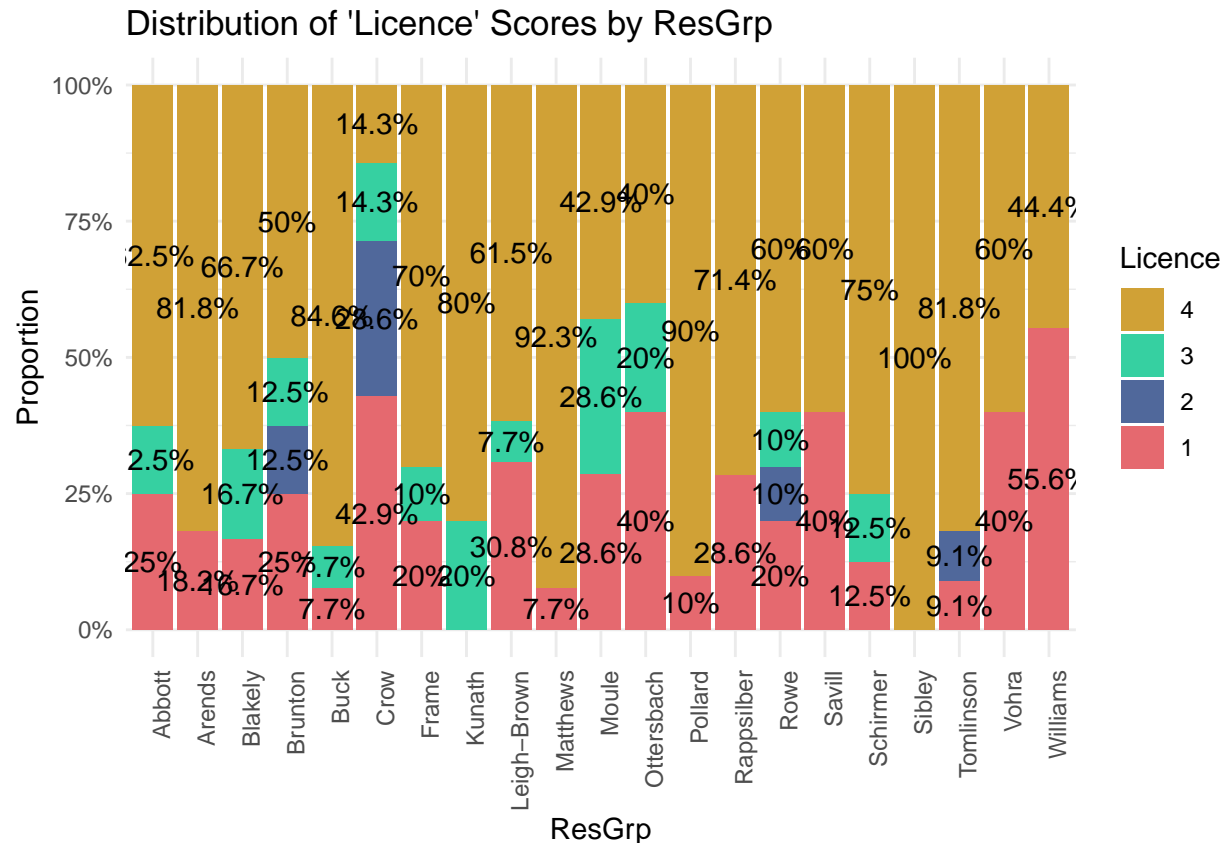
long_data <- long_data %>%
  mutate(Licence = factor(Licence, levels = c("4", "3", "2", "1")))

# Custom colors
colors <- c("#D0A136", "#36D0A1", "#50689B", "#E5696F")

# Creating the stacked bar chart with percentage labels
internal4 <- ggplot(long_data, aes(x = as.factor(ResGrp), y = n, fill = as.factor(Licence))) +
  geom_bar(stat = "identity", position = "fill") + # Scale the bar heights to proportions
  scale_y_continuous(labels = percent_format()) + # Convert the y-axis to percentage
  geom_text(
    aes(label = Label, y = Percentage),
    size = 4,
    color = "black",
    position = position_fill(vjust = 0.5)
  ) +
  scale_fill_manual(values = colors) +
  labs(title = "Distribution of 'Licence' Scores by ResGrp",
    x = "ResGrp",
    y = "Proportion",
    fill = "Licence") +
  theme_minimal() +
  scale_x_discrete(name = "ResGrp", labels = unique(long_data$ResGrp)) +
  theme(axis.text.x = element_text(angle = 90, hjust = 1)) # Adjust text angle for readability

# Print the plot
print(internal4)

```



```
ggsave("internal4.png", internal4, width = 15, height = 10, units = "in", bg= "white")
```

#these results were used for internal university meetings and presentations

Calculate the frequency and percentage for each 'DAS' score among the research Group

```
long_data <- data %>%
  count(ResGrp, NewDAS) %>%
  group_by(ResGrp) %>%
  mutate(Percentage = n / sum(n) * 100) %>%
  mutate(Label = ifelse(Percentage > 5, paste0(round(Percentage, 1), "%"), "")) %>%
  mutate(Cumulative_Percentage = cumsum(Percentage) - (0.5 * Percentage)) %>%
  ungroup()
```

Custom colors

```
colors <- c("Not Presented" = "gray", "Not Shared" = "#E5696F", "Shared" = "#50689B")
```

Creating the stacked bar chart with percentage labels

```
DASinternal <- ggplot(long_data, aes(x = as.factor(ResGrp), y = n, fill = as.factor(NewDAS))) +
  geom_bar(stat = "identity", position = "fill") + # Scale the bar heights to proportions
  scale_y_continuous(labels = percent_format()) + # Convert the y-axis to percentage
  geom_text(
    aes(label = Label, y = Percentage),
    size = 4,
    color = "black",
    position = position_fill(vjust = 0.5)
  ) +
```

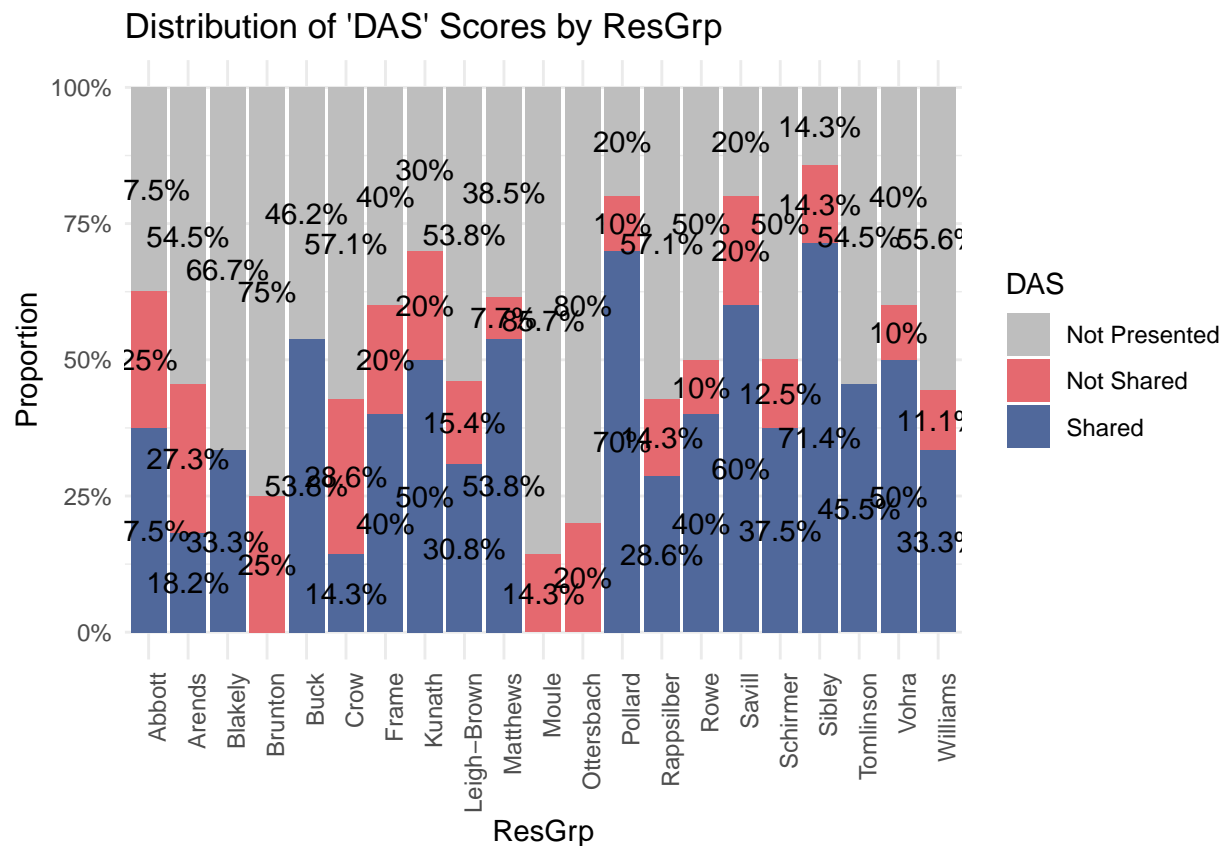


```

scale_fill_manual(values = colors) +
labs(title = "Distribution of 'DAS' Scores by ResGrp",
      x = "ResGrp",
      y = "Proportion",
      fill = "DAS") +
theme_minimal() +
scale_x_discrete(name = "ResGrp", labels = unique(long_data$ResGrp)) +
theme(axis.text.x = element_text(angle = 90, hjust = 1)) # Adjust text angle for readability

# Print the plot
print(DASinternal)

```



```

ggsave("DASinternal.png", DASinternal, width = 15, height = 10, units = "in", bg= "white")

```

#these results were used for internal university meetings and presentations

```

complete4 <- data %>%
  count(Reuse) %>%
  mutate(percentage = n / sum(n)*100)

print(complete4)

```

```
##   Reuse   n percentage
```

## 1	1 42	21.76166
## 2	2 21	10.88083
## 3	3 51	26.42487
## 4	4 79	40.93264