EFA

```
set.seed(42)
library(igraph)
##
## Attaching package: 'igraph'
## The following objects are masked from 'package:stats':
##
##
       decompose, spectrum
## The following object is masked from 'package:base':
##
##
       union
library(QuantPsyc) # for the multivariate normality test
## Loading required package: boot
## Loading required package: dplyr
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:igraph':
##
##
       as_data_frame, groups, union
  The following objects are masked from 'package:stats':
##
##
##
       filter, lag
  The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
##
## Loading required package: purrr
##
## Attaching package: 'purrr'
## The following objects are masked from 'package:igraph':
##
       compose, simplify
##
## Loading required package: MASS
##
## Attaching package: 'MASS'
## The following object is masked from 'package:dplyr':
##
##
       select
```

```
##
## Attaching package: 'QuantPsyc'
## The following object is masked from 'package:base':
##
##
      norm
library(dunn.test)
library(nFactors) # for the scree plot
## Loading required package: lattice
##
## Attaching package: 'lattice'
## The following object is masked from 'package:boot':
##
##
      melanoma
##
## Attaching package: 'nFactors'
## The following object is masked from 'package:lattice':
##
##
      parallel
library(psych) # for PA FA
##
## Attaching package: 'psych'
## The following object is masked from 'package:boot':
##
##
      logit
library(caret) # highly correlated features removal
## Loading required package: ggplot2
##
## Attaching package: 'ggplot2'
## The following objects are masked from 'package:psych':
##
##
      %+%, alpha
##
## Attaching package: 'caret'
## The following object is masked from 'package:purrr':
##
##
      lift
library(tidyverse)
## -- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
## v forcats 1.0.0
                       v stringr
                                   1.5.1
## v lubridate 1.9.3
                        v tibble
                                   3.2.1
                       v tidyr
## v readr
              2.1.5
                                   1.3.1
## -- Conflicts ----- tidyverse_conflicts() --
## x lubridate::%--%()
                           masks igraph::%--%()
```

```
masks psych::%+%()
## x ggplot2::%+%()
## x ggplot2::alpha()
                           masks psych::alpha()
## x tibble::as_data_frame() masks dplyr::as_data_frame(), igraph::as_data_frame()
## x purrr::compose()
                            masks igraph::compose()
## x tidyr::crossing()
                             masks igraph::crossing()
## x dplyr::filter()
                           masks stats::filter()
## x dplyr::lag()
                            masks stats::lag()
## x caret::lift()
                            masks purrr::lift()
                         masks dplyr::select()
## x MASS::select()
                           masks igraph::simplify()
## x purrr::simplify()
## i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become error
library(paletteer) # color palettes
library(conflicted) # to resolve QuantPsyc x dplyr conflicts
conflict_prefer("select", "dplyr")
## [conflicted] Will prefer dplyr::select over any other package.
conflict_prefer("filter", "dplyr")
## [conflicted] Will prefer dplyr::filter over any other package.
```

Load and tidy data

```
pretty_names <- read_csv("../feat_name_mapping.csv")</pre>
## Rows: 85 Columns: 2
## -- Column specification -----
## Delimiter: ","
## chr (2): name_orig, name_pretty
## i Use `spec()` to retrieve the full column specification for this data.
## i Specify the column types or set `show_col_types = FALSE` to quiet this message.
data <- read_csv("../measurements/measurements.csv")</pre>
## Rows: 754 Columns: 108
## -- Column specification -
## Delimiter: ","
## chr (20): fpath, KUK_ID, FileName, FileFormat, FolderPath, subcorpus, Source...
## dbl (85): RuleAbstractNouns, RuleAmbiguousRegards, RuleAnaphoricReferences, ...
## lgl (3): ClarityPursuit, SyllogismBased, Bindingness
## i Use `spec()` to retrieve the full column specification for this data.
## i Specify the column types or set `show_col_types = FALSE` to quiet this message.
.firstnonmetacolumn <- 17
data_no_nas <- data %>%
  select(!c(
   fpath,
    # KUK_ID,
    # FileName,
   FolderPath,
    # subcorpus,
```

```
DocumentTitle,
 ClarityPursuit,
 Readability,
 SyllogismBased,
 SourceDB
)) %>%
# replace -1s in variation coefficients with NAs
mutate(across(c(
  'RuleDoubleAdpos.max allowable distance.v',
  RuleTooManyNegations.max_negation_frac.v,
  `RuleTooManyNegations.max_allowable_negations.v`,
  `RuleTooManyNominalConstructions.max_noun_frac.v`,
  `RuleTooManyNominalConstructions.max_allowable_nouns.v`,
  `RuleCaseRepetition.max_repetition_count.v`,
  `RuleCaseRepetition.max_repetition_frac.v`,
  `RulePredSubjDistance.max distance.v`,
  `RulePredObjDistance.max_distance.v`,
  `RuleInfVerbDistance.max_distance.v`,
  `RuleMultiPartVerbs.max_distance.v`,
  `RuleLongSentences.max_length.v`,
  `RulePredAtClauseBeginning.max order.v`,
  `mattr.v`,
  `maentropy.v`
), ~ na_if(.x, -1))) %>%
# replace NAs with Os
replace na(list(
 RuleGPcoordovs = 0,
 RuleGPdeverbaddr = 0,
 RuleGPpatinstr = 0,
 RuleGPdeverbsubj = 0,
  RuleGPadjective = 0,
 RuleGPpatbenperson = 0,
 RuleGPwordorder = 0.
 RuleDoubleAdpos = 0,
 RuleDoubleAdpos.max allowable distance = 0,
 RuleDoubleAdpos.max_allowable_distance.v = 0,
 RuleAmbiguousRegards = 0,
 RuleReflexivePassWithAnimSubj = 0,
 RuleTooManyNegations = 0,
 RuleTooManyNegations.max_negation_frac = 0,
 RuleTooManyNegations.max_negation_frac.v = 0,
 RuleTooManyNegations.max_allowable_negations = 0,
  RuleTooManyNegations.max_allowable_negations.v = 0,
  RuleTooManyNominalConstructions.max_noun_frac.v = 0,
 RuleTooManyNominalConstructions.max_allowable_nouns.v = 0,
 RuleFunctionWordRepetition = 0,
 RuleCaseRepetition.max_repetition_count.v = 0,
  RuleCaseRepetition.max_repetition_frac.v = 0,
 RuleWeakMeaningWords = 0,
  RuleAbstractNouns = 0.
 RuleRelativisticExpressions = 0,
 RuleConfirmationExpressions = 0,
 RuleRedundantExpressions = 0,
```

```
RuleTooLongExpressions = 0,
   RuleAnaphoricReferences = 0,
   RuleLiteraryStyle = 0,
   RulePassive = 0,
    RulePredSubjDistance = 0,
   RulePredSubjDistance.max_distance = 0,
   RulePredSubjDistance.max_distance.v = 0,
   RulePredObjDistance = 0,
   RulePredObjDistance.max distance = 0,
   RulePredObjDistance.max_distance.v = 0,
   RuleInfVerbDistance = 0,
   RuleInfVerbDistance.max_distance = 0,
   RuleInfVerbDistance.max_distance.v = 0,
   RuleMultiPartVerbs = 0,
   RuleMultiPartVerbs.max_distance = 0,
   RuleMultiPartVerbs.max_distance.v = 0,
   RuleLongSentences.max_length.v = 0,
   RulePredAtClauseBeginning.max_order.v = 0,
   RuleVerbalNouns = 0,
   RuleDoubleComparison = 0,
   RuleWrongValencyCase = 0,
   RuleWrongVerbonominalCase = 0,
   RuleIncompleteConjunction = 0
  ))
data clean <- data no nas %>%
  # norm data expected to correlate with text length
  mutate(across(c(
   RuleGPcoordovs,
   RuleGPdeverbaddr,
   RuleGPpatinstr,
   RuleGPdeverbsubj,
   RuleGPadjective,
   RuleGPpatbenperson,
   RuleGPwordorder,
   RuleDoubleAdpos,
   RuleAmbiguousRegards,
   RuleFunctionWordRepetition,
   RuleWeakMeaningWords,
   RuleAbstractNouns,
   RuleRelativisticExpressions,
   RuleConfirmationExpressions,
   RuleRedundantExpressions,
   RuleTooLongExpressions,
   RuleAnaphoricReferences,
   RuleLiteraryStyle,
   RulePassive,
   RuleVerbalNouns,
   RuleDoubleComparison,
    RuleWrongValencyCase,
   RuleWrongVerbonominalCase,
   RuleIncompleteConjunction,
   num_hapax,
```

```
RuleReflexivePassWithAnimSubj,
 RuleTooManyNominalConstructions,
 RulePredSubjDistance,
 RuleMultiPartVerbs,
 RulePredAtClauseBeginning
), ~ .x / word_count)) %>%
mutate(across(c(
 RuleTooFewVerbs,
 RuleTooManyNegations,
 RuleCaseRepetition,
 RuleLongSentences,
 RulePredObjDistance,
 RuleInfVerbDistance
), ~ .x / sent_count)) %>%
# remove variables identified as "u counts"
select(!c(
 RuleTooFewVerbs,
 RuleTooManyNegations,
 RuleTooManyNominalConstructions,
 RuleCaseRepetition,
 RuleLongSentences,
 RulePredAtClauseBeginning,
 sent_count,
 word_count,
  syllab_count,
  char count
)) %>%
# remove variables identified as unreliable
select(!c(
 RuleAmbiguousRegards,
 RuleFunctionWordRepetition,
 RuleDoubleComparison,
 RuleWrongValencyCase,
 RuleWrongVerbonominalCase
# remove artificially limited variables
select(!c(
 RuleCaseRepetition.max_repetition_frac,
 RuleCaseRepetition.max_repetition_frac.v
)) %>%
# remove further variables belonging to the 'acceptability' category
select(!c(RuleIncompleteConjunction)) %>%
mutate(across(c(
  class,
 FileFormat,
  subcorpus,
 DocumentVersion,
 LegalActType,
 Objectivity,
 AuthorType,
 RecipientType,
 RecipientIndividuation,
 Anonymized
```

```
), ~ as.factor(.x)))
# no NAs should be present now
data_clean[!complete.cases(data_clean), ]
## # A tibble: 754 x 83
##
      KUK_ID
                             FileName FileFormat subcorpus SourceID DocumentVersion
##
      <chr>
                                                  <fct>
                                                            <chr>>
                                                                     <fct>
## 1 673b7a37c6537d54ff062~ 002 Kom~ TXT
                                                  KUKY
                                                            <NA>
                                                                     Original
## 2 673b7a37c6537d54ff062~ 006 Chc~ TXT
                                                  KUKY
                                                            < NA >
                                                                     Redesign
## 3 673b7a37c6537d54ff062~ 004_Nev~ TXT
                                                  KUKY
                                                            <NA>
                                                                     Original
## 4 673b7a37c6537d54ff062~ 008 Pol~ TXT
                                                  KUKY
                                                            <NA>
                                                                     Original
## 5 673b7a37c6537d54ff062~ 005_Och~ TXT
                                                  KUKY
                                                            <NA>
                                                                     Original
## 6 673b7a37c6537d54ff062~ 016_0bc~ TXT
                                                  KUKY
                                                            <NA>
                                                                     Original
## 7 673b7a37c6537d54ff062~ 019_Dět~ TXT
                                                  KUKY
                                                            <NA>
                                                                     Redesign
                                                                     Redesign
## 8 673b7a37c6537d54ff062~ 007_D\u00fc\u00fc~ TXT
                                                  KUKY
                                                            <NA>
## 9 673b7a37c6537d54ff062~ 024_Opa~ TXT
                                                  KUKY
                                                            <NA>
                                                                     Original
## 10 673b7a37c6537d54ff062~ 047_Dav~ TXT
                                                  KUKY
                                                            <NA>
                                                                     Original
## # i 744 more rows
## # i 77 more variables: ParentDocumentID <chr>, LegalActType <fct>,
       Objectivity <fct>, Bindingness <lgl>, AuthorType <fct>,
## #
       RecipientType <fct>, RecipientIndividuation <fct>, Anonymized <fct>,
## #
      `Recipient Type` <chr>, class <fct>, RuleAbstractNouns <dbl>,
       RuleAnaphoricReferences <dbl>,
## #
       RuleCaseRepetition.max_repetition_count <dbl>, ...
data_clean_scaled <- data_clean %>%
  mutate(across(class, ~ .x == "good")) %>%
  mutate(across(.firstnonmetacolumn:length(names(data clean)), ~ scale(.x)))
## Warning: There was 1 warning in `mutate()`.
## i In argument: `across(.firstnonmetacolumn:length(names(data_clean)),
     ~scale(.x))`.
## Caused by warning:
##! Using an external vector in selections was deprecated in tidyselect 1.1.0.
## i Please use `all_of()` or `any_of()` instead.
##
##
     data %>% select(.firstnonmetacolumn)
##
##
     # Now:
##
     data %>% select(all of(.firstnonmetacolumn))
## See <https://tidyselect.r-lib.org/reference/faq-external-vector.html>.
```

Important features identification

```
feature_importances <- tibble(
  feat_name = character(), p_value = numeric()
)

for (i in .firstnonmetacolumn:ncol(data_clean)) {
  fname <- names(data_clean)[i]</pre>
```

```
formula_single <- reformulate(fname, "class")</pre>
  glm_model <- glm(formula_single, data_clean, family = "binomial")</pre>
  glm_coefficients <- summary(glm_model)$coefficients</pre>
  row_index <- which(rownames(glm_coefficients) == fname)</pre>
  p_value <- glm_coefficients[row_index, 4]</pre>
  feature_importances <- feature_importances %>%
    add_row(feat_name = fname, p_value = p_value)
feature_importances
## # A tibble: 67 x 2
                                                     p_value
##
      feat_name
##
      <chr>>
                                                       <dbl>
## 1 RuleAbstractNouns
                                                  0.00187
## 2 RuleAnaphoricReferences
                                                  0.660
## 3 RuleCaseRepetition.max_repetition_count
                                                  0.0722
## 4 RuleCaseRepetition.max_repetition_count.v 0.00479
## 5 RuleConfirmationExpressions
                                                  0.0985
## 6 RuleDoubleAdpos
                                                  0.312
## 7 RuleDoubleAdpos.max_allowable_distance
                                                  0.000154
## 8 RuleDoubleAdpos.max_allowable_distance.v 0.00000356
## 9 RuleGPadjective
                                                  0.380
## 10 RuleGPcoordovs
                                                  0.828
## # i 57 more rows
selected_features <- feature_importances %>%
 mutate(selected = p value <= 0.05)</pre>
selected_features %>% write_csv("selected_features.csv")
selected_features_names <- selected_features %>%
  filter(selected) %>%
 pull(feat_name)
```

Correlations

See Levshina (2015: 353-54).
analyze_correlation <- function(data) {
 cor_matrix <- cor(data)

cor_tibble_long <- cor_matrix %>%
 as_tibble() %>%
 mutate(feat1 = rownames(cor_matrix)) %>%
 pivot_longer(!feat1, names_to = "feat2", values_to = "cor") %>%
 mutate(abs_cor = abs(cor))

cor_matrix_upper <- cor_matrix
 cor_matrix_upper[lower.tri(cor_matrix_upper)] <- 0

cor_tibble_long_upper <- cor_matrix_upper %>%
 as_tibble() %>%
 mutate(feat1 = rownames(cor_matrix)) %>%
 pivot_longer(!feat1, names_to = "feat2", values_to = "cor") %>%

```
mutate(abs_cor = abs(cor)) %>%
   filter(feat1 != feat2 & abs_cor > 0)
 list(
    cor_matrix = cor_matrix,
    cor_matrix_upper = cor_matrix_upper,
   cor_tibble_long = cor_tibble_long,
    cor tibble long upper = cor tibble long upper
 )
}
data_purish <- data_clean %>% select(any_of(selected_features_names)) %>%
  # remove features expected to have low communalities
  select(!c(
   RuleDoubleAdpos.max_allowable_distance,
   RuleDoubleAdpos.max_allowable_distance.v,
   RuleGPwordorder,
   RuleLiteraryStyle,
   maentropy.v,
   RuleTooManyNegations.max_negation_frac,
   RulePredSubjDistance.max_distance,
   RuleTooManyNegations.max_allowable_negations,
   RuleTooManyNegations.max_allowable_negations.v,
   RuleTooManyNominalConstructions.max_allowable_nouns.v,
   RuleTooFewVerbs.min verb frac.v,
   RulePredObjDistance.max_distance.v,
   RulePredObjDistance.max distance,
   RulePredAtClauseBeginning.max_order.v,
   RuleInfVerbDistance
  )) %>%
  # remove features expected to have low loadings
  select(!c(
   RuleMultiPartVerbs.max_distance.v,
   RulePredSubjDistance.max_distance.v,
    RuleLongSentences.max_length
 ))
```

Extremely non-normal data

```
# # remove where median == 0?
# keep <- character()</pre>
# for (i in seq_along(colnames(data_purish))) {
  cname <- colnames(data purish)[i]</pre>
   q <- quantile(data_purish[, i][[1]], probs = 0.10)[[1]]</pre>
#
#
  if (q > 0) {
#
     keep <- c(keep, cname)
     cat("keep", cname, "\n")
#
   } else {
#
     cat("throw out", cname, "\n")
#
# }
# data_purish <- data_purish %>% select(any_of(keep))
```

High correlations

```
.hcorrcutoff <- 0.9
analyze_correlation(data_purish)$cor_tibble_long %>%
filter(feat1 != feat2 & abs_cor > .hcorrcutoff) %>%
arrange(feat1, -abs_cor) %>%
print(n = 100)
```

```
## # A tibble: 16 x 4
##
      feat1
                 feat2
                              cor abs cor
##
      <chr>
                 <chr>
                            <dbl>
                                     <dbl>
##
    1 ari
                            0.984
                                     0.984
                 fkgl
                            0.978
                                     0.978
##
    2 ari
                 gf
##
    3 ari
                 smog
                            0.951
                                     0.951
##
   4 atl
                 cli
                            0.960
                                     0.960
##
    5 cli
                 atl
                            0.960
                                     0.960
                            0.984
                                     0.984
##
    6 fkgl
                 ari
##
    7 fkgl
                 gf
                            0.967
                                     0.967
##
    8 fkgl
                 smog
                            0.949
                                     0.949
##
    9 gf
                            0.987
                                     0.987
                 smog
## 10 gf
                 ari
                            0.978
                                     0.978
## 11 gf
                            0.967
                                     0.967
                 fkgl
## 12 maentropy mattr
                            0.964
                                     0.964
                                     0.964
## 13 mattr
                 maentropy 0.964
## 14 smog
                            0.987
                                     0.987
                 gf
                            0.951
                                     0.951
## 15 smog
                 ari
## 16 smog
                 fkgl
                            0.949
                                     0.949
```

exclude:

- ari: corr. w/ RuleLongSentences.max_length > 0.94; sentence length seems more universal, let's make it a substitute
- gf: corr. w/ RuleLongSentences.max_length > 0.92; sentence length seems more universal, let's make it a substitute
- maentropy: corr. w/ mattr > 0.96, but mattr is implemented in QuitaUp. besides, the interesting thing about maentropy is its variation
- smog: corr. w/ fkgl almost 0.95, but fkgl coefficients adjusted for Czech are available
- atl: corr. w/ cli around 0.96; unlike cli, atl is not a readability metric

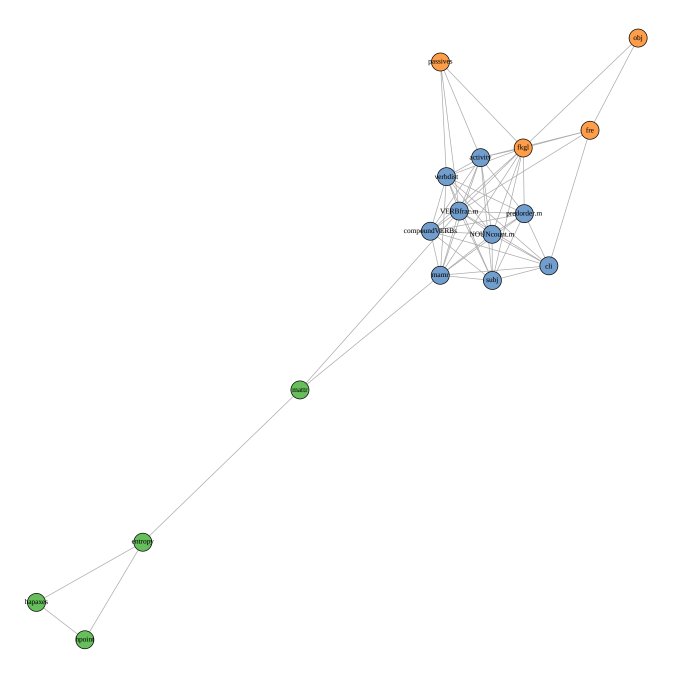
```
high_correlations <- findCorrelation(cor(data_purish), verbose = TRUE)
```

```
## Compare row 20 and column 26 with corr 0.978
##
    Means: 0.399 vs 0.207 so flagging column 20
## Compare row 26 and column 32 with corr 0.987
##
    Means: 0.378 vs 0.195 so flagging column 26
## Compare row 32 and column 24 with corr 0.949
    Means: 0.35 vs 0.184 so flagging column 32
##
## Compare row 21 and column 22 with corr 0.96
    Means: 0.27 vs 0.176 so flagging column 21
##
## Compare row 28 and column 30 with corr 0.964
    Means: 0.194 vs 0.171 so flagging column 28
## All correlations <= 0.9
names(data_purish)[high_correlations]
```

```
data_pureish_striphigh <- data_purish %>% select(!all_of(high_correlations))
analyze_correlation(data_pureish_striphigh)$cor_tibble_long %>%
 filter(feat1 != feat2 & abs_cor > .hcorrcutoff) %>%
  arrange(feat1, -abs_cor) %>%
 print(n = 100)
## # A tibble: 0 x 4
## # i 4 variables: feat1 <chr>, feat2 <chr>, cor <dbl>, abs_cor <dbl>
Low correlations
# 0.35 instead of 0.3 otherwise the FA bootstrapping would freeze
.lcorrcutoff <- 0.35
low_correlating_features <- analyze_correlation(data_pureish_striphigh)$</pre>
  cor_tibble_long %>%
  filter(feat1 != feat2) %>%
  group_by(feat1) %>%
  summarize(max_cor = max(abs_cor)) %>%
  filter(max_cor < .lcorrcutoff) %>%
 pull(feat1)
feature_importances %>% filter(feat_name %in% low_correlating_features)
## # A tibble: 11 x 2
##
     feat name
                                                          p_value
                                                            <dbl>
##
      <chr>>
## 1 RuleAbstractNouns
                                                       0.00187
## 2 RuleCaseRepetition.max_repetition_count.v
                                                       0.00479
## 3 RuleGPdeverbaddr
                                                       0.0112
## 4 RuleGPdeverbsubj
                                                       0.0133
## 5 RuleMultiPartVerbs.max_distance
                                                       0.00320
## 6 RuleRedundantExpressions
                                                       0.0104
## 7 RuleRelativisticExpressions
                                                       0.00205
## 8 RuleTooManyNegations.max_negation_frac.v
                                                       0.0365
## 9 RuleTooManyNominalConstructions.max noun frac.v 0.00000311
## 10 RuleVerbalNouns
                                                       0.0000748
## 11 RuleWeakMeaningWords
                                                       0.0386
data_pure <- data_pureish_striphigh %>%
  select(!any_of(low_correlating_features))
cnames <- map(</pre>
  colnames(data_pure),
  function(x) {
   pull(pretty_names %>%
      filter(name_orig == x), name_pretty)
) %>% unlist()
colnames(data_pure) <- cnames</pre>
```

Visualisation

```
my_colors <- paletteer::paletteer_d("ggthemes::Classic_10_Medium")</pre>
network_edges <- analyze_correlation(data_pure)$cor_tibble_long_upper %>%
 filter(abs_cor > 0.3)
network <- graph_from_data_frame(</pre>
  network_edges,
  directed = FALSE
E(network)$weight <- network_edges$abs_cor</pre>
network_communities <- cluster_optimal(network)</pre>
network_membership <- membership(network_communities)</pre>
plot(
 network,
 layout = layout.fruchterman.reingold,
 vertex.color = map(
    network_communities$membership,
    function(x) my_colors[x]
  ) %>% unlist(use.names = FALSE),
  vertex.size = 6,
  vertex.label.color = "black",
  vertex.label.cex = 0.7
```



Scaling

```
data_scaled <- data_pure %>%
  mutate(across(seq_along(data_pure), ~ scale(.x)[, 1]))
final_collist <- data_scaled %>% colnames()
```

Check for normality

```
mult.norm(data_scaled %>% as.data.frame())$mult.test
## Beta-hat kappa p-val
```

```
## Skewness 351.5182 44174.1153 0  
## Kurtosis 858.5678 289.3036 0
```

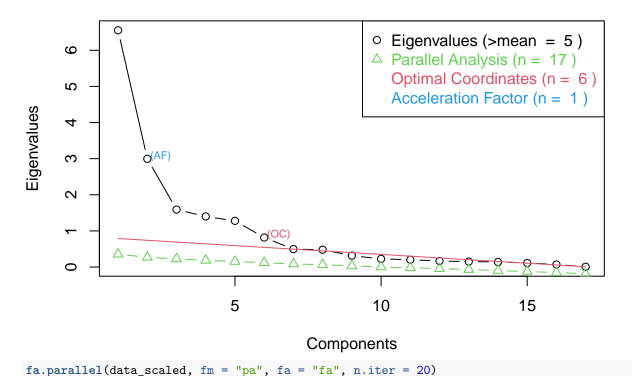
Low (null) p-values show that we can reject the hypothesis that the data would be in a multivariate normal distribution. I.e. the distribution isn't multivariate normal.

$\mathbf{F}\mathbf{A}$

No. of factors

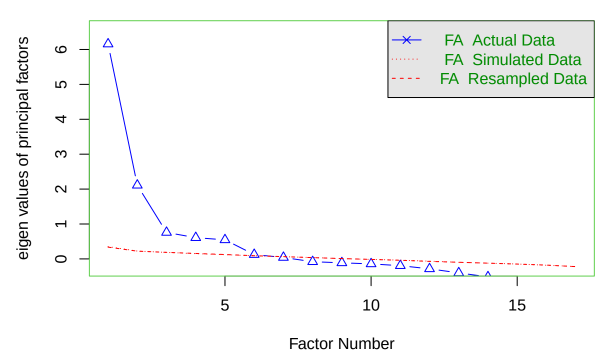
```
eigen <- eigen(cor(data_scaled))
par <- nFactors::parallel(
    subject = nrow(data_scaled),
    var = ncol(data_scaled),
    rep = 100,
    quantile = .95,
    model = "factors"
)
scree <- nScree(x = eigen$values, aparallel = par$eigen$qevpea)
plotnScree(scree)</pre>
```

Non Graphical Solutions to Scree Test



```
## Warning in fa.stats(r = r, f = f, phi = phi, n.obs = n.obs, np.obs = np.obs, :
## The estimated weights for the factor scores are probably incorrect. Try a
## different factor score estimation method.
```

Parallel Analysis Scree Plots



Parallel analysis suggests that the number of factors = 6 and the number of components = N.

Model

https://www.rdocumentation.org/packages/psych/versions/2.5.3/topics/fa

```
# appears to be the happiest when nfactors = 6 or 7
# throws the The estimated weights for the factor scores are probably incorrect.
# Try a different factor score estimation method. warning otherwise
fa_res <- fa(
    data_scaled,
        nfactors = 7,
    fm = "pa",
    rotate = "promax",
    oblique.scores = TRUE,
    scores = "tenBerge",
    n.iter = 100
)</pre>
```

Loading required namespace: GPArotation

```
fa_res
```

```
## Factor Analysis with confidence intervals using method = fa(r = data_scaled, nfactors = 7, n.iter =
      scores = "tenBerge", fm = "pa", oblique.scores = TRUE)
## Factor Analysis using method = pa
## Call: fa(r = data_scaled, nfactors = 7, n.iter = 100, rotate = "promax",
      scores = "tenBerge", fm = "pa", oblique.scores = TRUE)
## Standardized loadings (pattern matrix) based upon correlation matrix
                               PA3
                PA1
                     PA2
                           PA6
                                      PA4
                                           PA5
                                                 PA7
                                                      h2
## compoundVERBs 0.79 0.00 -0.17 -0.01 0.04 0.24 0.51 0.78 0.2219 2.0
                ## passives
```

```
## predorder.m
                -0.72 -0.02 0.13 -0.07 -0.11 -0.05 0.02 0.55 0.4487 1.1
                 0.24 -0.01 0.96 -0.15 -0.02 -0.05 -0.12 0.71 0.2887 1.2
## obj
## subj
                 0.72 0.09 0.02 0.09 -0.15 0.18 -0.01 0.53 0.4684 1.3
                 0.75 -0.01 -0.03 -0.01 -0.03 -0.31 0.24 0.91 0.0932 1.6
## VERBfrac.m
## NOUNcount.m -1.06 0.06 -0.16 0.11 -0.07 -0.13 -0.07 0.86 0.1443 1.1
                 0.72  0.01  0.15 -0.15 -0.02 -0.43  0.18  0.93  0.0695  2.0
## activity
                 0.20 -0.04 -0.14 0.95 0.08 -0.23 -0.03 0.91 0.0852 1.3
## cli
                 0.12 0.74 -0.05 0.04 0.54 0.00 0.03 0.95 0.0461 1.9
## entropy
## fkgl
                -0.40 0.04 0.60 0.07 0.04 0.19 0.08 1.00 0.0013 2.1
                 0.17 -0.03 -0.53 -0.53 -0.04 -0.07 -0.11 0.99 0.0127 2.4
## fre
## hpoint
                 0.01 0.94 0.01 -0.02 -0.01 -0.02 -0.02 0.87 0.1348 1.0
                 0.68 -0.06 -0.04 0.22 -0.32 0.04 0.06 0.75 0.2506 1.7
## mamr
                -0.06 -0.12 -0.01 0.08 0.83 0.04 0.02 0.72 0.2818 1.1
## mattr
                 0.09 -0.94 -0.02 0.03 0.29 0.04 -0.01 0.86 0.1397 1.2
## hapaxes
## verbdist
                -0.94 0.02 -0.27 -0.09 -0.10 0.08 0.07 0.82 0.1803 1.2
##
##
                        PA1 PA2 PA6 PA3 PA4 PA5 PA7
## SS loadings
                        5.52 2.34 1.71 1.31 1.26 1.22 0.35
## Proportion Var
                        0.32 0.14 0.10 0.08 0.07 0.07 0.02
## Cumulative Var
                        0.32 0.46 0.56 0.64 0.71 0.79 0.81
## Proportion Explained 0.40 0.17 0.13 0.10 0.09 0.09 0.03
## Cumulative Proportion 0.40 0.57 0.70 0.79 0.89 0.97 1.00
##
## With factor correlations of
##
        PA1
             PA2
                   PA6 PA3
                             PA4
                                    PA5
                                          PA7
## PA1 1.00 0.02 -0.38 0.04 -0.27 -0.49 -0.09
## PA2 0.02 1.00 0.30 0.17 0.17 0.19 -0.01
## PA6 -0.38 0.30 1.00 0.27 0.13 0.22 0.28
## PA3 0.04 0.17 0.27 1.00 0.10 0.29 0.33
## PA4 -0.27 0.17 0.13 0.10 1.00 0.11 -0.05
## PA5 -0.49 0.19 0.22 0.29 0.11 1.00 0.22
## PA7 -0.09 -0.01 0.28 0.33 -0.05 0.22 1.00
## Mean item complexity = 1.5
## Test of the hypothesis that 7 factors are sufficient.
## df null model = 136 with the objective function = 17.23 with Chi Square = 12859.06
## df of the model are 38 and the objective function was 0.41
## The root mean square of the residuals (RMSR) is 0.01
## The df corrected root mean square of the residuals is 0.02
## The harmonic n.obs is 754 with the empirical chi square 19.64 with prob < 0.99
## The total n.obs was 754 with Likelihood Chi Square = 305.8 with prob < 1.5e-43
## Tucker Lewis Index of factoring reliability = 0.924
## RMSEA index = 0.097 and the 90 % confidence intervals are 0.087 0.107
## BIC = 54.04
## Fit based upon off diagonal values = 1
## Measures of factor score adequacy
                                                   PA1 PA2 PA6 PA3 PA4 PA5
##
## Correlation of (regression) scores with factors 0.99 0.98 0.99 0.99 0.95 0.93
## Multiple R square of scores with factors
                                                  0.97 0.96 0.98 0.98 0.90 0.86
## Minimum correlation of possible factor scores
                                                 0.95 0.91 0.96 0.96 0.80 0.73
```

```
PA7
##
## Correlation of (regression) scores with factors
                                               0.85
## Multiple R square of scores with factors
                                               0.72
## Minimum correlation of possible factor scores
                                               0.44
##
   Coefficients and bootstrapped confidence intervals
                      PA1 upper
                low
                                low
                                     PA2 upper
                                               low
                                                      PA6 upper
                                                                 low
## compoundVERBs 0.69 0.79 0.92 -0.05 0.00 0.03 -0.21 -0.17 -0.08 -0.06 -0.01
## passives
                0.04 0.10 0.14 -0.06 -0.03 0.00 -0.06 0.00 0.04 -0.22 -0.17
               -0.83 -0.72 -0.59 -0.07 -0.02 0.03 0.07 0.13 0.20 -0.22 -0.07
## predorder.m
## obj
                0.57
                     0.72  0.85  0.04  0.09  0.16  -0.07  0.02  0.11  0.03  0.09
## subj
## VERBfrac.m
                0.63 0.75 0.88 -0.05 -0.01 0.01 -0.07 -0.03 0.02 -0.05 -0.01
               -1.19 -1.06 -0.88 0.02 0.06 0.09 -0.20 -0.16 -0.09 0.07 0.11
## NOUNcount.m
                0.60 0.72 0.84 -0.03 0.01 0.03 0.11 0.15 0.21 -0.18 -0.15
## activity
## cli
                0.15
                     0.20 0.24 -0.05 -0.04 -0.01 -0.18 -0.14 -0.11
                                                               0.91 0.95
## entropy
                0.08 0.12 0.15 0.71 0.74 0.78 -0.07 -0.05 -0.02 0.01 0.04
## fkgl
               -0.47 -0.40 -0.32 0.01 0.04 0.06 0.53 0.60 0.69 0.04 0.07
               0.12  0.17  0.20 -0.05 -0.03 -0.01 -0.61 -0.53 -0.46 -0.61 -0.53
## fre
               -0.02 0.01 0.05 0.91 0.94 0.96 -0.02 0.01 0.04 -0.05 -0.02
## hpoint
## mamr
               ## mattr
               -0.11 -0.06 0.00 -0.16 -0.12 -0.09 -0.05 -0.01 0.04 0.04 0.08
               0.04 0.09 0.13 -0.97 -0.94 -0.90 -0.06 -0.02 0.02 0.01 0.03
## hapaxes
               -1.03 -0.94 -0.81 -0.02 0.02 0.06 -0.36 -0.27 -0.16 -0.12 -0.09
## verbdist
##
                           PA4 upper
                                           PA5 upper
               upper
                      low
                                      low
                                                      low
                                                           PA7 upper
## compoundVERBs 0.05 0.01 0.04 0.07 0.14 0.24 0.41 0.32 0.51 0.83
## passives
               -0.12 0.00 0.03 0.06 0.62 0.82 1.09 -0.13 0.12 0.54
               0.03 -0.16 -0.11 -0.03 -0.17 -0.05 0.10 -0.19 0.02 0.22
## predorder.m
               -0.11 -0.05 -0.02 0.01 -0.15 -0.05 0.00 -0.21 -0.12 -0.03
## obj
## subj
               0.15 -0.21 -0.15 -0.09 0.08 0.18 0.29 -0.30 -0.01 0.37
## VERBfrac.m
               0.39
## NOUNcount.m
               0.16 -0.10 -0.07 -0.02 -0.20 -0.13 -0.06 -0.46 -0.07
                                                                0.18
## activity
               -0.12 -0.05 -0.02 0.01 -0.53 -0.43 -0.33 0.13 0.18 0.24
                ## cli
## entropy
                0.07 0.49 0.54 0.57 -0.03 0.00 0.03 -0.01 0.03 0.10
                0.11 0.01 0.04 0.06 0.12 0.19 0.30 -0.02 0.08 0.15
## fkgl
               -0.46 -0.06 -0.04 -0.02 -0.15 -0.07 -0.03 -0.17 -0.11 -0.06
## hpoint
               0.01 -0.04 -0.01 0.02 -0.06 -0.02 0.02 -0.08 -0.02 0.03
                0.27 -0.37 -0.32 -0.25 -0.06 0.04
                                                0.13 -0.21 0.06 0.45
## mamr
                0.11 0.79 0.83 0.87 -0.01 0.04 0.11 -0.09 0.02 0.16
## mattr
                0.07 0.25 0.29 0.33 -0.02 0.04 0.08 -0.08 -0.01 0.10
## hapaxes
               -0.05 -0.13 -0.10 -0.07 -0.01 0.08 0.25 -0.13 0.07 0.24
## verbdist
   Interfactor correlations and bootstrapped confidence intervals
##
           lower estimate upper
## PA1-PA2 -0.0770
                   0.017
                         0.102
## PA1-PA6 -0.5718
                  -0.381 -0.084
                   0.044 0.388
## PA1-PA3 -0.6675
## PA1-PA4 -0.6476
                  -0.270 0.228
## PA1-PA5 -0.6478
                  -0.486
                         0.039
## PA1-PA7 -0.5236
                  -0.093 0.264
## PA2-PA6 0.1799
                   0.297 0.387
## PA2-PA3 -0.0011
                   0.170 0.367
## PA2-PA4 0.0259
                   0.168 0.282
```

```
## PA2-PA5 0.0416
                    0.193 0.282
## PA2-PA7 -0.2526
                   -0.013 0.186
                     0.269 0.454
## PA6-PA3 -0.0012
## PA6-PA4 -0.0020
                     0.134 0.383
## PA6-PA5 0.0516
                     0.219 0.318
## PA6-PA7 -0.0908
                     0.285 0.526
## PA3-PA4 -0.1297
                     0.100 0.456
## PA3-PA5 -0.0808
                     0.288 0.422
## PA3-PA7 -0.2458
                     0.328 0.599
## PA4-PA5 -0.0785
                     0.113 0.283
## PA4-PA7 -0.2606
                   -0.053 0.477
## PA5-PA7 -0.3846
                     0.225 0.555
```

Loadings

fa_res\$loadings

```
##
## Loadings:
                               PA6
                                              PA4
                                                     PA5
                 PA1
                        PA2
                                       PA3
                                                            PA7
## compoundVERBs 0.794
                                -0.165
                                                      0.239 0.505
## passives
                  0.101
                                       -0.169
                                                      0.817 0.123
## predorder.m
                 -0.719
                                0.129
                                              -0.107
## obj
                  0.236
                                0.965 - 0.155
                                                            -0.124
## subj
                  0.721
                                              -0.154 0.183
## VERBfrac.m
                  0.745
                                                     -0.312 0.240
## NOUNcount.m
               -1.060
                               -0.158 0.109
                                                     -0.131
## activity
                  0.716
                                0.151 - 0.147
                                                     -0.432 0.176
## cli
                  0.203
                                -0.138 0.955
                                                     -0.229
## entropy
                  0.121 0.742
                                               0.541
## fkgl
                 -0.404
                                0.605
                                                      0.194
## fre
                  0.171
                                -0.535 -0.527
                                                            -0.113
## hpoint
                         0.936
                  0.685
                                        0.222 -0.318
## mamr
## mattr
                        -0.122
                                               0.826
## hapaxes
                        -0.942
                                               0.294
## verbdist
                 -0.937
                               -0.272
                                              -0.105
##
##
                    PA1
                          PA2
                                PA6
                                       PA3
                                             PA4
## SS loadings
                  5.533 2.351 1.774 1.360 1.227 1.171 0.411
## Proportion Var 0.325 0.138 0.104 0.080 0.072 0.069 0.024
## Cumulative Var 0.325 0.464 0.568 0.648 0.720 0.789 0.813
for (i in 1:fa_res$factors) {
  cat("\n----", colnames(fa_res$loadings)[i], "----\n")
  loadings <- fa res$loadings[, i]</pre>
  load_df <- data.frame(loading = loadings)</pre>
 load_df_filtered <- load_df %>%
    mutate(abs 1 = abs(loading)) %>%
    mutate(str = case_when(
      abs_1 > 0.7 ~ "***",
      abs_1 <= 0.7 & abs_1 > 0.5 ~ "** ",
```

```
abs_1 <= 0.5 & abs_1 > 0.3 ~ "* ",
     abs_1 <= 0.3 & abs_1 > 0.1 ~ ". ",
      .default = ""
   )) %>%
   arrange(-abs_1) %>%
   filter(abs_l > 0.1)
  load_df_filtered %>%
   mutate(across(c(loading, abs_l), ~ round(.x, 3))) %>%
   print()
  cat("\n")
}
##
## ---- PA1 ----
##
                loading abs_l str
                 -1.060 1.060 ***
## NOUNcount.m
## verbdist
                 -0.937 0.937 ***
## compoundVERBs 0.794 0.794 ***
## VERBfrac.m
                 0.745 0.745 ***
## subj
                 0.721 0.721 ***
## predorder.m
               -0.719 0.719 ***
## activity
                0.716 0.716 ***
## mamr
                 0.685 0.685 **
## fkgl
                 -0.404 0.404 *
## obj
                  0.236 0.236 .
## cli
                 0.203 0.203 .
## fre
                 0.171 0.171 .
## entropy
                 0.121 0.121 .
## passives
                 0.101 0.101 .
##
## ----- PA2 -----
          loading abs_l str
## hapaxes -0.942 0.942 ***
## hpoint 0.936 0.936 ***
## entropy 0.742 0.742 ***
## mattr -0.122 0.122 .
##
##
## ---- PA6 ----
##
                loading abs_l str
## obj
                 0.965 0.965 ***
                 0.605 0.605 **
## fkgl
## fre
                 -0.535 0.535 **
## verbdist
                 -0.272 0.272 .
## compoundVERBs -0.165 0.165 .
## NOUNcount.m
                 -0.158 0.158 .
## activity
                 0.151 0.151 .
## cli
                 -0.138 0.138 .
## predorder.m 0.129 0.129 .
##
##
```

```
## ----- PA3 -----
##
             loading abs_l str
## cli
              0.955 0.955 ***
              -0.527 0.527 **
## fre
## mamr
               0.222 0.222 .
              -0.169 0.169 .
## passives
## obj
              -0.155 0.155 .
            -0.147 0.147 .
## activity
## NOUNcount.m 0.109 0.109 .
##
##
## ---- PA4 ----
##
             loading abs_l str
## mattr
              0.826 0.826 ***
## entropy
              0.541 0.541 **
## mamr
               -0.318 0.318 *
## hapaxes
              0.294 0.294 .
## subj
              -0.154 0.154 .
## predorder.m -0.107 0.107 .
## verbdist
             -0.105 0.105 .
##
##
## ----- PA5 -----
                loading abs_l str
##
## passives
               0.817 0.817 ***
                 -0.432 0.432 *
## activity
## VERBfrac.m
                -0.312 0.312 *
## compoundVERBs 0.239 0.239 .
## cli
                -0.229 0.229 .
## fkgl
                0.194 0.194 .
          0.183 0.183 .
## subj
## NOUNcount.m -0.131 0.131 .
##
##
## ----- PA7 -----
##
                loading abs_l str
## compoundVERBs 0.505 0.505 **
## VERBfrac.m
                 0.240 0.240 .
## activity
                 0.176 0.176 .
                 -0.124 0.124 .
## obj
                 0.123 0.123 .
## passives
## fre
                 -0.113 0.113 .
```

hypotheses:

- PA1: register narrativity, richness of expression; shorter clauses (-technical / +narrative)
 - narrativity? (1st and 2nd persons etc.)
- PA2: text length (-short / +long)
 - hapaxes load negatively, because I normed them over word count
- PA6: sentence complexity (more clauses) (-simple / +complex)
 - slightly longer nominal constructions / more objects, more years of education necessary, predicates slightly further in the clause, slightly more verbs
 - fkgl in strong correlation with sentlen.m
- PA3: word length (-short / +long)
 - cli highly correlates with atl, meaning the factor likely expresses mostly token lengths

- slightly more passives, slightly more objects, slightly less verbal overall / slightly longer nom. constructions, slightly morphologically richer, many years of education necessary
- more enumerations? but one would expect higher activity differences to occur if that was the case
- PA4: lexical richness (-poor / +rich)
- **PA5:** passivity (-active / +passive)
 - compound verbs, because that's what passives are in Czech
 - smaller activity, because passive participles count as ADJ in UD.
- PA7: compound verbs (-less / +more)

strong correlations:

- PA1-PA6: (-0.38) narrativity leads to simple clauses
- PA2-PA6: (+0.30) longer texts include more complex sentences
- PA1-PA5: (-0.49, topconf = +0.09) narrative texts more active

NOTE: variables with low communalities are excluded from the analysis, yet still likely play a role in legal writing readability. this includes both those selected for the analysis and the excluded ones.

NOTE: some high-correlating variables were excluded from the FA.

Healthiness diagnostics

```
fa_res$loadings[] %>%
  as_tibble() %>%
  mutate(feat = cnames) %>%
  select(feat, everything()) %>%
  pivot_longer(!feat) %>%
  mutate(value = abs(value)) %>%
  group_by(feat) %>%
  summarize(maxload = max(value)) %>%
  arrange(maxload)
```

```
## # A tibble: 17 x 2
##
      feat
                    maxload
##
      <chr>
                       <dbl>
##
   1 fre
                      0.535
##
   2 fkgl
                      0.605
##
   3 mamr
                      0.685
##
   4 activity
                      0.716
##
   5 predorder.m
                      0.719
##
   6 subj
                      0.721
  7 entropy
##
                      0.742
## 8 VERBfrac.m
                      0.745
  9 compoundVERBs
                      0.794
## 10 passives
                      0.817
## 11 mattr
                      0.826
## 12 hpoint
                      0.936
## 13 verbdist
                      0.937
## 14 hapaxes
                      0.942
## 15 cli
                      0.955
                      0.965
## 16 obj
## 17 NOUNcount.m
                      1.06
```

```
fa_res$communality %>% sort()
##
                                     passives
                                                         obj
            subj
                   predorder.m
                                                                      mattr
##
                                    0.5856923
       0.5315955
                     0.5512988
                                                   0.7113441
                                                                  0.7181813
            mamr compoundVERBs
##
                                     verbdist
                                                 NOUNcount.m
                                                                    hapaxes
##
       0.7493974
                     0.7781089
                                    0.8196810
                                                   0.8557424
                                                                  0.8602815
##
          hpoint
                    VERBfrac.m
                                           cli
                                                    activity
                                                                    entropy
##
       0.8652002
                     0.9067914
                                    0.9147991
                                                   0.9305464
                                                                  0.9539003
##
             fre
                           fkgl
       0.9872669
##
                     0.9987222
```

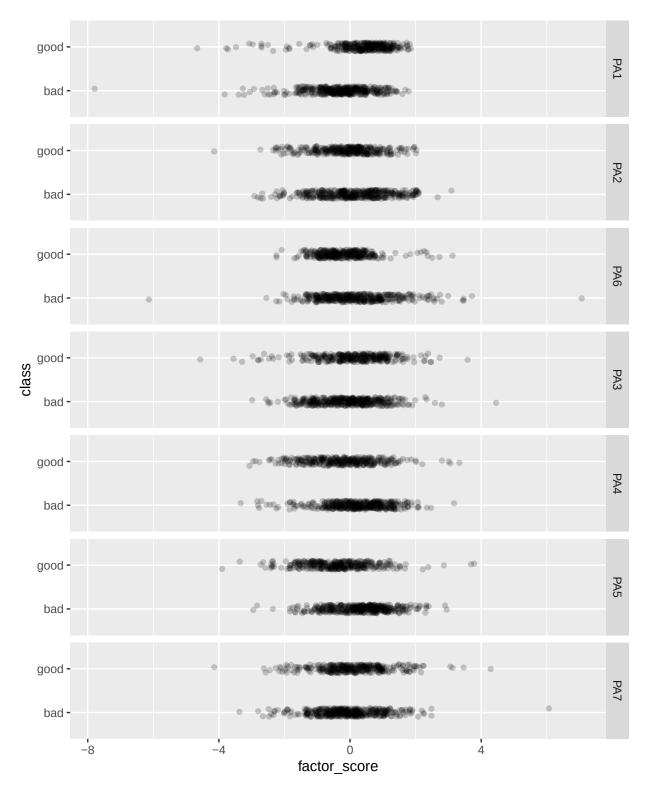
Uniquenesses

```
fa res$uniquenesses %>% round(3)
## compoundVERBs
                       passives
                                  predorder.m
                                                          obj
                                                                       subj
##
           0.222
                          0.414
                                         0.449
                                                        0.289
                                                                      0.468
##
      VERBfrac.m
                   NOUNcount.m
                                     activity
                                                          cli
                                                                    entropy
           0.093
                          0.144
                                                        0.085
                                                                      0.046
##
                                         0.069
##
            fkgl
                            fre
                                       hpoint
                                                        mamr
                                                                      mattr
           0.001
                                         0.135
                                                        0.251
                                                                      0.282
##
                          0.013
##
         hapaxes
                       verbdist
##
           0.140
                          0.180
```

Distributions over factors

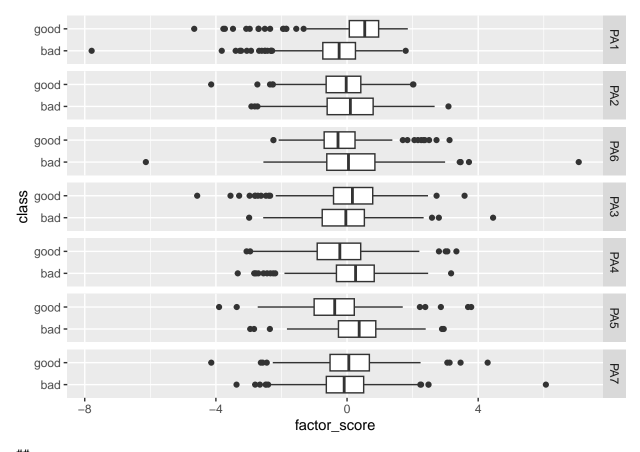
```
analyze_distributions <- function(data_factors_long, variable) {</pre>
  plot <- data_factors_long %>%
    ggplot(aes(x = factor_score, y = !!sym(variable))) +
    geom_boxplot() +
    facet_grid(factor ~ .)
  print(plot)
  formula <- reformulate(variable, "factor_score")</pre>
  factors <- levels(data_factors_long$factor)</pre>
  min p values <- numeric()</pre>
  for (f in factors) {
    data <- data_factors_long %>% filter(factor == f)
    cat(
      "\nTest for the significance of differences in",
      variable, "over", f, ":\n\n"
    )
    dunn <- dunn.test(</pre>
      data$factor_score, data[[variable]],
      altp = TRUE, method = "bonferroni"
    )
    min_p_values <- c(min_p_values, min(dunn$altP.adjusted))</pre>
  }
  cat(
```

```
"\np < 5e-2\tfound in:",
    factors[min_p_values < 0.05],</pre>
    "\np < 1e-2\tfound in:",
    factors[min_p_values < 0.01],</pre>
    "\np < 1e-3\tfound in:",
    factors[min_p_values < 0.001],</pre>
    "\np < 1e-4\tfound in:",
    factors[min_p_values < 0.0001], "\n"</pre>
  )
}
data_factors <- bind_cols(data_clean, fa_res$scores %>% as.data.frame())
cnames <- map(</pre>
  colnames(data_factors),
  function(x) {
    name <- pull(pretty_names %>%
      filter(name_orig == x), name_pretty)
    if (length(name) == 1) {
      return(name)
    } else {
      return(x)
  }
) %>% unlist()
colnames(data_factors) <- cnames</pre>
data_factors_long <- data_factors %>%
  pivot_longer(PA1:PA7, names_to = "factor", values_to = "factor_score") %>%
  mutate(across(
    factor,
    ~ factor(.x, levels = c("PA1", "PA2", "PA6", "PA3", "PA4", "PA5", "PA7"))
  ))
data_factors_long %>%
  ggplot(aes(x = factor_score, y = class)) +
  facet_grid(factor ~ .) +
  theme(legend.position = "bottom") +
  geom_jitter(width = 0, height = 0.1, alpha = 0.2)
```



class

analyze_distributions(data_factors_long, "class")



```
## Test for the significance of differences in class over PA1 :
##
##
     Kruskal-Wallis rank sum test
##
## data: x and group
  Kruskal-Wallis chi-squared = 147.6724, df = 1, p-value = 0
##
##
                               Comparison of x by group
##
##
                                      (Bonferroni)
## Col Mean-|
## Row Mean |
##
##
       good | -12.15205
                 0.0000*
##
##
## alpha = 0.05
## Reject Ho if p <= alpha
\mbox{\tt \#\#} Test for the significance of differences in class over PA2 :
##
##
     Kruskal-Wallis rank sum test
## data: x and group
## Kruskal-Wallis chi-squared = 8.0515, df = 1, p-value = 0
##
```

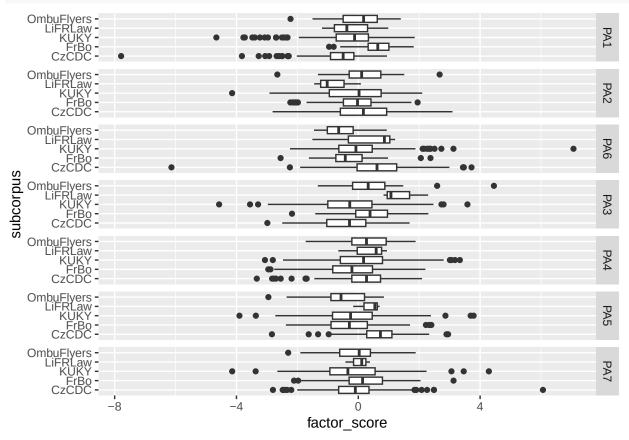
```
##
##
                             Comparison of x by group
                                   (Bonferroni)
##
## Col Mean-|
## Row Mean |
## -----
      good | 2.837517
               0.0045*
##
           ##
## alpha = 0.05
## Reject Ho if p <= alpha
\#\# Test for the significance of differences in class over PA6 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 23.4951, df = 1, p-value = 0
##
##
##
                             Comparison of x by group
##
                                   (Bonferroni)
## Col Mean-|
## Row Mean |
## -----
      good | 4.847174
##
          0.0000*
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in class over PA3 :
##
##
    Kruskal-Wallis rank sum test
## data: x and group
## Kruskal-Wallis chi-squared = 11.9438, df = 1, p-value = 0
##
##
##
                             Comparison of x by group
##
                                   (Bonferroni)
## Col Mean-
## Row Mean |
## -----
      good | -3.455977
               0.0005*
##
           ##
## alpha = 0.05
## Reject Ho if p \le alpha
\#\# Test for the significance of differences in class over PA4 :
##
##
    Kruskal-Wallis rank sum test
##
```

```
## data: x and group
## Kruskal-Wallis chi-squared = 46.5471, df = 1, p-value = 0
##
##
##
                             Comparison of x by group
##
                                   (Bonferroni)
## Col Mean-I
## Row Mean |
## -----
      good | 6.822546
##
##
       - 1
                0.0000*
##
## alpha = 0.05
## Reject Ho if p <= alpha
\#\# Test for the significance of differences in class over PA5 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 101.1707, df = 1, p-value = 0
##
##
                             Comparison of x by group
                                   (Bonferroni)
##
## Col Mean-I
## Row Mean |
                    bad
## -----
      good | 10.05836
##
                0.0000*
##
          - 1
##
## alpha = 0.05
## Reject Ho if p <= alpha
\#\# Test for the significance of differences in class over PA7 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 4.3737, df = 1, p-value = 0.04
##
##
                             Comparison of x by group
##
                                   (Bonferroni)
## Col Mean-|
## Row Mean |
      good | -2.091333
##
       | 0.0365*
##
##
## alpha = 0.05
## Reject Ho if p <= alpha
##
## p < 5e-2 found in: PA1 PA2 PA6 PA3 PA4 PA5 PA7
```

```
## p < 1e-2 found in: PA1 PA2 PA6 PA3 PA4 PA5
## p < 1e-3 found in: PA1 PA6 PA3 PA4 PA5
## p < 1e-4 found in: PA1 PA6 PA4 PA5</pre>
```

subcorpus

analyze_distributions(data_factors_long, "subcorpus")



```
##
\#\# Test for the significance of differences in subcorpus over PA1 :
##
##
     Kruskal-Wallis rank sum test
##
## data: x and group
   Kruskal-Wallis chi-squared = 324.6698, df = 4, p-value = 0
##
##
##
                               Comparison of x by group
##
                                      (Bonferroni)
## Col Mean-|
   Row Mean |
                    CzCDC
                                 FrBo
                                            KUKY
                                                     LiFRLaw
##
##
       FrBo |
               -17.35858
##
                  0.0000*
##
##
       KUKY |
               -5.082162
                            11.31002
##
            1
                  0.0000*
                             0.0000*
```

```
##
## LiFRLaw | -0.872398 1.790209 -0.003452
##
    1.0000 0.7342 1.0000
         - 1
##
## OmbuFlye | -4.533034 4.348685 -1.649043 -0.484868
##
        0.0001* 0.0001* 0.9914 1.0000
## alpha = 0.05
## Reject Ho if p <= alpha
##
## Test for the significance of differences in subcorpus over PA2 :
##
   Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 9.8975, df = 4, p-value = 0.04
##
##
##
                          Comparison of x by group
##
                               (Bonferroni)
## Col Mean-|
## Row Mean | CzCDC
                         FrBo KUKY LiFRLaw
## -----+----
     FrBo | 2.438507
##
     0.1475
##
         - 1
            2.006418 -0.192445
##
    KUKY |
             0.4481 1.0000
##
      - 1
##
## LiFRLaw | 1.738831 1.368187 1.394671
              0.8206 1.0000 1.0000
##
          ##
          - 1
## OmbuFlye | -0.091483 -1.356400 -1.214028 -1.712554
                       1.0000
##
         - 1
               1.0000
                                 1.0000
                                          0.8679
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in subcorpus over PA6 :
##
##
   Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 137.7653, df = 4, p-value = 0
##
##
##
                          Comparison of x by group
##
                               (Bonferroni)
## Col Mean-|
               CzCDC
                         FrBo KUKY LiFRLaw
## Row Mean |
     FrBo | 10.82530
##
     1 0.0000*
##
         ##
```

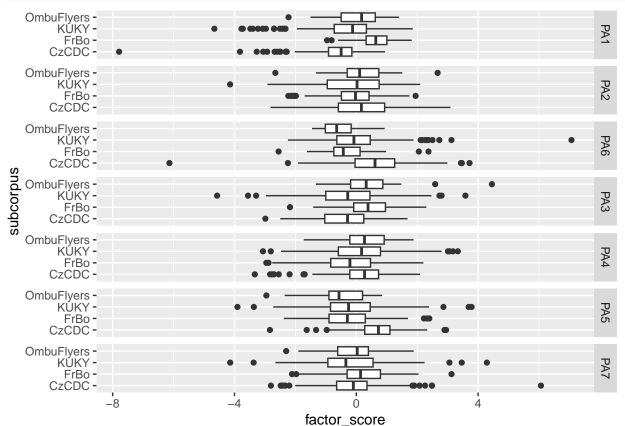
```
KUKY | 6.313404 -3.656522
##
##
       | 0.0000* 0.0026*
##
            0.539206 -1.121281 -0.539826
  LiFRLaw |
##
##
     1
              1.0000 1.0000 1.0000
##
         - 1
## OmbuFlye | 7.526018 2.098278 3.927170 1.686430
              0.0000* 0.3588 0.0009* 0.9171
##
          ##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in subcorpus over PA3 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 112.907, df = 4, p-value = 0
##
##
##
                          Comparison of x by group
##
                               (Bonferroni)
## Col Mean-|
## Row Mean |
            CzCDC
                                    KUKY
                          FrBo
                                           LiFRLaw
  ______
      FrBo | -9.239165
##
      1
             0.0000*
##
          KUKY | -1.300172 7.537538
##
        1.0000 0.0000*
##
##
## LiFRLaw | -3.023849 -1.612043 -2.799330
##
         | 0.0250* 1.0000 0.0512
         - 1
##
## OmbuFlye | -4.182661 0.502797 -3.419066 1.703689
##
              0.0003* 1.0000 0.0063* 0.8844
         ##
## alpha = 0.05
## Reject Ho if p <= alpha
##
## Test for the significance of differences in subcorpus over PA4 :
##
    Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 36.3801, df = 4, p-value = 0
##
##
##
                          Comparison of x by group
##
                               (Bonferroni)
## Col Mean-|
## Row Mean |
                          FrBo
                                    KUKY LiFRLaw
    FrBo | 5.459549
##
```

```
0.0000*
##
         - 1
##
           -
##
      KUKY | 1.732943 -3.411845
##
               0.8311
                        0.0065*
          ##
           LiFRLaw | -0.146970 -0.985287 -0.442916
##
      1
               1.0000
                       1.0000
##
                                    1.0000
##
          ## OmbuFlye | -0.538009 -3.377899 -1.503636 -0.015445
                1.0000 0.0073* 1.0000 1.0000
##
         - 1
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in subcorpus over PA5 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 169.6171, df = 4, p-value = 0
##
##
##
                            Comparison of x by group
                                  (Bonferroni)
## Col Mean-
## Row Mean |
                CzCDC
                           {\tt FrBo}
                                     KUKY LiFRLaw
##
      FrBo | 12.03428
##
       0.0000*
##
##
##
      KUKY |
             9.587804 -1.289058
##
          - 1
               0.0000*
                           1.0000
   LiFRLaw |
             0.509819 -1.336300 -1.128601
##
##
          1.0000 1.0000
                                    1.0000
##
          - 1
## OmbuFlye |
             7.022416 0.956708 1.594639 1.567093
##
           0.0000* 1.0000 1.0000 1.0000
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in subcorpus over PA7 :
##
    Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 30.7162, df = 4, p-value = 0
##
##
##
                            Comparison of x by group
##
                                  (Bonferroni)
## Col Mean-I
## Row Mean |
             CzCDC
                            FrBo
                                       KUKY
                                              LiFRLaw
```

```
##
##
       FrBo |
               -3.966477
                  0.0007*
##
##
##
       KUKY |
                 1.189282
                            5.123866
##
                   1.0000
                             0.0000*
##
    LiFRLaw |
##
                -0.340818
                            0.267296
                                       -0.543730
##
                   1.0000
                               1.0000
                                          1.0000
##
##
   OmbuFlye |
                -0.544032
                            1.497093
                                       -1.205231
                                                    0.170711
                                                      1.0000
##
                   1.0000
                              1.0000
                                          1.0000
##
## alpha = 0.05
## Reject Ho if p <= alpha
##
## p < 5e-2 found in: PA1 PA6 PA3 PA4 PA5 PA7
## p < 1e-2 found in: PA1 PA6 PA3 PA4 PA5 PA7
## p < 1e-3 found in: PA1 PA6 PA3 PA4 PA5 PA7
## p < 1e-4 found in: PA1 PA6 PA3 PA4 PA5 PA7
```

subcorpus wo/ LiFRLaw

```
analyze_distributions(
  data_factors_long %>% filter(subcorpus != "LiFRLaw"), "subcorpus"
)
```



```
##
## Test for the significance of differences in subcorpus over PA1 :
##
    Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 324.9953, df = 3, p-value = 0
##
##
                           Comparison of x by group
##
                                 (Bonferroni)
## Col Mean-|
                                 KUKY
## Row Mean |
                CzCDC
                          FrBo
      FrBo | -17.37448
##
      - 1
             0.0000*
##
          KUKY | -5.087708 11.31941
##
##
       0.0000*
                       0.0000*
##
          ## OmbuFlye | -4.539209 4.350595 -1.652060
    0.0000* 0.0001* 0.5911
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in subcorpus over PA2 :
    Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 7.6553, df = 3, p-value = 0.05
##
##
##
                           Comparison of x by group
                                 (Bonferroni)
##
## Col Mean-
## Row Mean |
             CzCDC
                       FrBo
                                   KUKY
      FrBo | 2.447210
##
      - 1
              0.0864
##
          KUKY I
             2.004739 -0.202682
##
##
              0.2699 1.0000
       ## OmbuFlye | -0.089710 -1.359092 -1.211330
         1.0000 1.0000 1.0000
##
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in subcorpus over PA6 :
##
##
    Kruskal-Wallis rank sum test
```

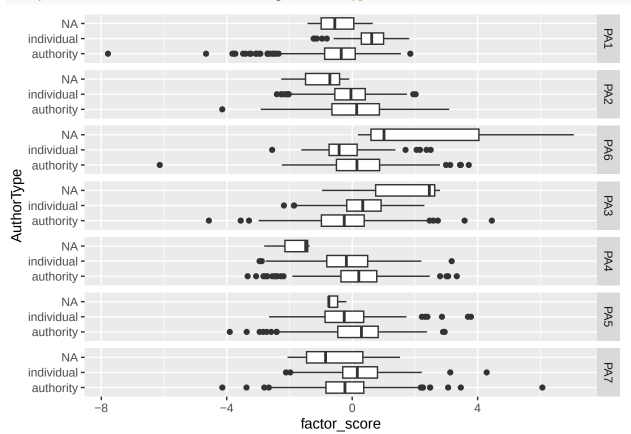
```
##
## data: x and group
## Kruskal-Wallis chi-squared = 137.8128, df = 3, p-value = 0
##
                           Comparison of x by group
##
                                 (Bonferroni)
## Col Mean-
                           FrBo
## Row Mean |
                 CzCDC
                                   KUKY
      FrBo | 10.83598
             0.0000*
##
       ##
          ##
      KUKY | 6.316190 -3.663846
##
               0.0000*
                       0.0015*
          ##
## OmbuFlye |
              7.539865 2.106923 3.939339
               0.0000* 0.2107 0.0005*
##
         ##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in subcorpus over PA3 :
##
##
    Kruskal-Wallis rank sum test
## data: x and group
## Kruskal-Wallis chi-squared = 107.8981, df = 3, p-value = 0
##
##
##
                           Comparison of x by group
##
                                 (Bonferroni)
## Col Mean-|
## Row Mean |
                CzCDC
                           FrBo KUKY
##
      FrBo | -9.256635
##
      0.0000*
##
         ##
      KUKY | -1.298220 7.556555
##
             1.0000 0.0000*
         ## OmbuFlye | -4.189325 0.505022 -3.426765
               0.0002* 1.0000 0.0037*
          ##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in subcorpus over PA4 :
##
    Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 36.0837, df = 3, p-value = 0
##
```

```
##
##
                             Comparison of x by group
                                   (Bonferroni)
##
## Col Mean-|
                                   KUKY
## Row Mean |
                  CzCDC
                            {\tt FrBo}
      FrBo | 5.461216
              0.0000*
##
           -
##
           -
      KUKY |
##
              1.734750 -3.411506
          - 1
                 0.4967
                           0.0039*
##
## OmbuFlye | -0.537587 -3.378330 -1.504230
##
           1.0000
                         0.0044*
                                     0.7951
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in subcorpus over PA5 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 168.5035, df = 3, p-value = 0
##
##
##
                             Comparison of x by group
                                   (Bonferroni)
## Col Mean-|
                  CzCDC
## Row Mean |
                            FrBo
                                       KUKY
## -----
##
      FrBo | 12.01538
             0.0000*
##
        ##
           KUKY |
              9.574393 -1.285257
##
                0.0000*
##
                          1.0000
          ## OmbuFlye |
               7.011861
                          0.955689
                                    1.591681
                0.0000*
##
                          1.0000
                                       0.6687
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in subcorpus over PA7 :
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 30.6296, df = 3, p-value = 0
##
##
                             Comparison of x by group
                                   (Bonferroni)
##
## Col Mean-|
```

```
CzCDC
## Row Mean |
                                FrBo
                                           KUKY
##
               -3.961805
##
       FrBo |
##
                 0.0004*
            ##
##
       KUKY |
                1.188349
                           5.118335
##
                  1.0000
                            0.0000*
            1
##
##
  OmbuFlye |
               -0.542887
                            1.495845 -1.203573
##
                  1.0000
                             0.8082
                                         1.0000
##
## alpha = 0.05
## Reject Ho if p <= alpha
## p < 5e-2 found in: PA1 PA6 PA3 PA4 PA5 PA7
## p < 1e-2 found in: PA1 PA6 PA3 PA4 PA5 PA7
## p < 1e-3 found in: PA1 PA6 PA3 PA4 PA5 PA7
## p < 1e-4 found in: PA1 PA6 PA3 PA4 PA5 PA7
```

AuthorType

analyze_distributions(data_factors_long, "AuthorType")



```
##
## Test for the significance of differences in AuthorType over PA1 :
##
## Kruskal-Wallis rank sum test
```

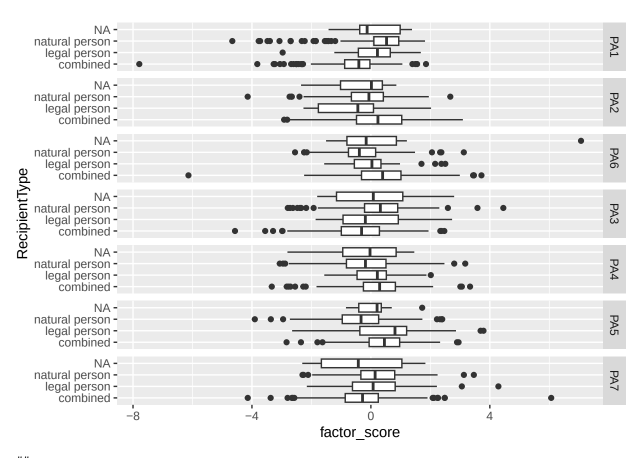
```
##
## data: x and group
## Kruskal-Wallis chi-squared = 300.0919, df = 1, p-value = 0
##
                             Comparison of x by group
                                    (Bonferroni)
##
## Col Mean-|
              authorit
## Row Mean |
## individu | -17.32316
          0.0000*
##
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in AuthorType over PA2 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 9.2401, df = 1, p-value = 0
##
##
##
                             Comparison of x by group
##
                                    (Bonferroni)
## Col Mean-|
## Row Mean |
               authorit
               3.039749
## individu |
##
          0.0024*
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in AuthorType over PA6 :
##
##
    Kruskal-Wallis rank sum test
## data: x and group
## Kruskal-Wallis chi-squared = 58.5159, df = 1, p-value = 0
##
##
##
                             Comparison of x by group
                                    (Bonferroni)
## Col Mean-|
## Row Mean |
              authorit
## -----
## individu | 7.649569
                0.0000*
          ##
## alpha = 0.05
## Reject Ho if p <= alpha
##
```

```
## Test for the significance of differences in AuthorType over PA3 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 81.9629, df = 1, p-value = 0
##
##
                             Comparison of x by group
##
                                   (Bonferroni)
## Col Mean-|
## Row Mean |
              authorit
## -----
## individu | -9.053337
##
          0.0000*
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in AuthorType over PA4:
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 30.2702, df = 1, p-value = 0
##
##
                             Comparison of x by group
                                   (Bonferroni)
##
## Col Mean-|
## Row Mean |
              authorit
## -----
## individu | 5.501835
##
         0.0000*
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in AuthorType over PA5 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 39.5694, df = 1, p-value = 0
##
##
                             Comparison of x by group
##
                                   (Bonferroni)
## Col Mean-|
## Row Mean |
              authorit
## -----
## individu | 6.290419
          1 0.0000*
##
##
```

```
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in AuthorType over PA7 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 40.9073, df = 1, p-value = 0
##
##
##
                             Comparison of x by group
##
                                   (Bonferroni)
## Col Mean-|
## Row Mean |
              authorit
## -----
## individu | -6.395879
##
                0.0000*
       1
##
## alpha = 0.05
## Reject Ho if p <= alpha
## p < 5e-2 found in: PA1 PA2 PA6 PA3 PA4 PA5 PA7
## p < 1e-2 found in: PA1 PA2 PA6 PA3 PA4 PA5 PA7
## p < 1e-3 found in: PA1 PA6 PA3 PA4 PA5 PA7
## p < 1e-4 found in: PA1 PA6 PA3 PA4 PA5 PA7
```

RecipientType

```
analyze_distributions(data_factors_long, "RecipientType")
```



```
## Test for the significance of differences in RecipientType over PA1 :
##
     Kruskal-Wallis rank sum test
##
##
## data: x and group
   Kruskal-Wallis chi-squared = 235.2855, df = 2, p-value = 0
##
##
##
                               Comparison of x by group
##
                                     (Bonferroni)
## Col Mean-|
  Row Mean
                {\tt combined}
                            legal pe
##
   legal pe |
               -3.202351
##
                 0.0041*
##
                           -2.168998
##
   natural
               -15.33836
                  0.0000*
##
                              0.0902
##
  alpha = 0.05
##
   Reject Ho if p <= alpha
##
##
  Test for the significance of differences in RecipientType over PA2:
##
##
     Kruskal-Wallis rank sum test
##
```

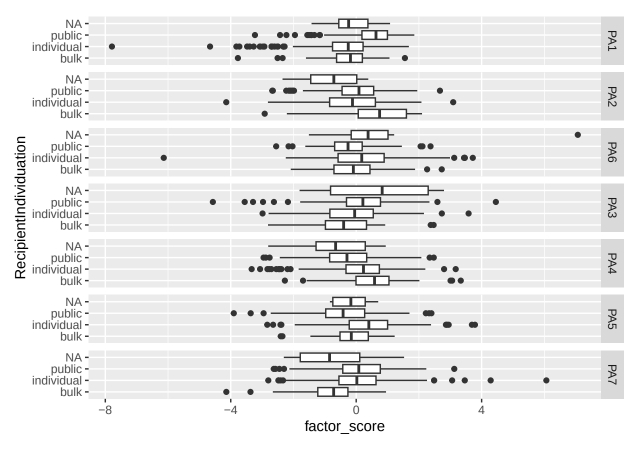
```
## data: x and group
## Kruskal-Wallis chi-squared = 34.4591, df = 2, p-value = 0
##
##
##
                           Comparison of x by group
##
                                 (Bonferroni)
## Col Mean-I
## Row Mean |
             combined
## -----
## legal pe | 3.898285
       0.0003*
             5.079110 -2.144990
## natural |
               0.0000*
##
          0.0959
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in RecipientType over PA6 :
##
    Kruskal-Wallis rank sum test
## data: x and group
## Kruskal-Wallis chi-squared = 97.616, df = 2, p-value = 0
##
##
##
                           Comparison of x by group
                                 (Bonferroni)
## Col Mean-|
## Row Mean | combined legal pe
## -----
## legal pe |
             1.873469
        0.1830
##
          9.879555
## natural |
                        1.587947
               0.0000*
          -
                        0.3369
##
## alpha = 0.05
## Reject Ho if p <= alpha
##
## Test for the significance of differences in RecipientType over PA3 :
##
    Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 88.5372, df = 2, p-value = 0
##
##
##
                           Comparison of x by group
                                 (Bonferroni)
## Col Mean-|
## Row Mean |
            combined legal pe
## -----
## legal pe | -1.401806
```

```
0.4829
##
           1
##
           1
## natural | -9.396916 -1.893909
##
                0.0000*
           0.1747
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in RecipientType over PA4 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 36.6854, df = 2, p-value = 0
##
##
##
                             Comparison of x by group
                                   (Bonferroni)
##
## Col Mean-|
## Row Mean |
              combined
                         legal pe
## -----
## legal pe |
               1.047412
##
           0.8847
##
           6.054595
                         1.074785
## natural |
           0.0000*
                          0.8474
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in RecipientType over PA5 :
##
    Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 127.5444, df = 2, p-value = 0
##
##
##
                             Comparison of x by group
##
                                   (Bonferroni)
## Col Mean-|
## Row Mean |
              combined legal pe
## -----
## legal pe |
               0.058011
                 1.0000
           1
##
               11.07741
                          3.841344
## natural
           - 1
##
                0.0000*
                          0.0004*
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in RecipientType over PA7 :
##
```

```
Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 44.2699, df = 2, p-value = 0
##
##
                            Comparison of x by group
                                  (Bonferroni)
##
## Col Mean-
## Row Mean | combined legal pe
## -----
## legal pe | -1.794855
          0.2180
##
## natural | -6.636304 -0.525457
                0.0000*
##
           1.0000
##
## alpha = 0.05
## Reject Ho if p <= alpha
## p < 5e-2 found in: PA1 PA2 PA6 PA3 PA4 PA5 PA7
## p < 1e-2 found in: PA1 PA2 PA6 PA3 PA4 PA5 PA7
## p < 1e-3 found in: PA1 PA2 PA6 PA3 PA4 PA5 PA7
## p < 1e-4 found in: PA1 PA2 PA6 PA3 PA4 PA5 PA7
court decisions often with RecipientType = combined.
```

${\bf Recipient Individuation}$

```
analyze_distributions(data_factors_long, "RecipientIndividuation")
```



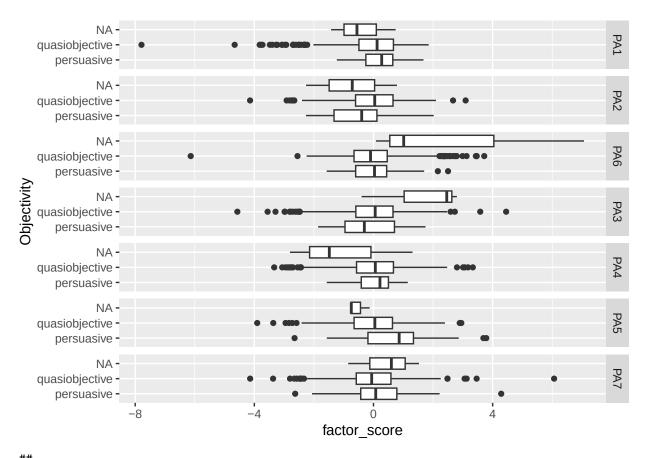
```
## Test for the significance of differences in RecipientIndividuation over PA1 :
##
##
     Kruskal-Wallis rank sum test
##
##
  data: x and group
   Kruskal-Wallis chi-squared = 209.5856, df = 2, p-value = 0
##
##
##
                               Comparison of x by group
##
                                     (Bonferroni)
## Col Mean-|
##
  Row Mean |
                    bulk
                            individu
##
   individu |
                0.552940
                   1.0000
##
##
##
     public |
               -7.610655
                           -14.06096
                 0.0000*
                             0.0000*
##
##
   alpha = 0.05
##
   Reject Ho if p \le alpha
##
  Test for the significance of differences in RecipientIndividuation over PA2 :
##
##
##
     Kruskal-Wallis rank sum test
##
```

```
## data: x and group
## Kruskal-Wallis chi-squared = 33.7947, df = 2, p-value = 0
##
##
##
                            Comparison of x by group
##
                                 (Bonferroni)
## Col Mean-I
## Row Mean |
               bulk
                         individu
## individu | 5.741813
     - 1
             0.0000*
##
          public |
             4.258459 -2.452715
##
##
               0.0001*
                       0.0425*
      - 1
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in RecipientIndividuation over PA6:
##
    Kruskal-Wallis rank sum test
## data: x and group
## Kruskal-Wallis chi-squared = 39.0551, df = 2, p-value = 0
##
##
##
                            Comparison of x by group
                                 (Bonferroni)
## Col Mean-|
## Row Mean |
             bulk individu
## -----
## individu | -2.045441
##
        0.1224
##
          public |
##
             1.596328
                       6.239990
##
                0.3312
                        0.0000*
           -
##
## alpha = 0.05
## Reject Ho if p <= alpha
##
## Test for the significance of differences in RecipientIndividuation over PA3 :
##
    Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 29.2184, df = 2, p-value = 0
##
##
##
                            Comparison of x by group
                                 (Bonferroni)
## Col Mean-|
## Row Mean |
                bulk individu
## -----
## individu | -2.043038
```

```
0.1231
##
          1
##
           -
    public | -4.487184 -4.249809
##
##
               0.0000*
           0.0001*
##
## alpha = 0.05
## Reject Ho if p <= alpha
\#\# Test for the significance of differences in RecipientIndividuation over PA4:
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 58.2474, df = 2, p-value = 0
##
##
##
                            Comparison of x by group
                                  (Bonferroni)
##
## Col Mean-|
## Row Mean |
                 bulk individu
## -----
## individu | 2.146907
##
         0.0954
##
           -
             5.867168
##
   public |
                         6.451235
##
      1
             0.0000*
                        0.0000*
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in RecipientIndividuation over PA5 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 100.7444, df = 2, p-value = 0
##
##
##
                            Comparison of x by group
##
                                  (Bonferroni)
## Col Mean-|
## Row Mean |
                 bulk individu
## -----
## individu | -4.029464
              0.0002*
##
         ##
           public |
              1.783526
                         9.946414
##
##
                0.2235
                        0.0000*
           ##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in RecipientIndividuation over PA7 :
##
```

```
Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 50.2363, df = 2, p-value = 0
##
                            Comparison of x by group
##
                                  (Bonferroni)
##
## Col Mean-|
## Row Mean |
                 bulk individu
## individu | -6.460242
       0.0000*
##
          ##
   public | -6.988232 -1.027106
        0.0000*
##
                           0.9131
##
## alpha = 0.05
## Reject Ho if p <= alpha
## p < 5e-2 found in: PA1 PA2 PA6 PA3 PA4 PA5 PA7
## p < 1e-2 found in: PA1 PA2 PA6 PA3 PA4 PA5 PA7
## p < 1e-3 found in: PA1 PA2 PA6 PA3 PA4 PA5 PA7
## p < 1e-4 found in: PA1 PA2 PA6 PA3 PA4 PA5 PA7
Objectivity
```

analyze_distributions(data_factors_long, "Objectivity")



```
## Test for the significance of differences in Objectivity over PA1 :
##
     Kruskal-Wallis rank sum test
##
##
## data: x and group
  Kruskal-Wallis chi-squared = 0.4589, df = 1, p-value = 0.5
##
##
##
                               Comparison of x by group
##
                                     (Bonferroni)
## Col Mean-|
## Row Mean |
                persuasi
##
  quasiobj |
                0.677452
##
                  0.4981
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in Objectivity over PA2 :
##
     Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 4.2938, df = 1, p-value = 0.04
##
```

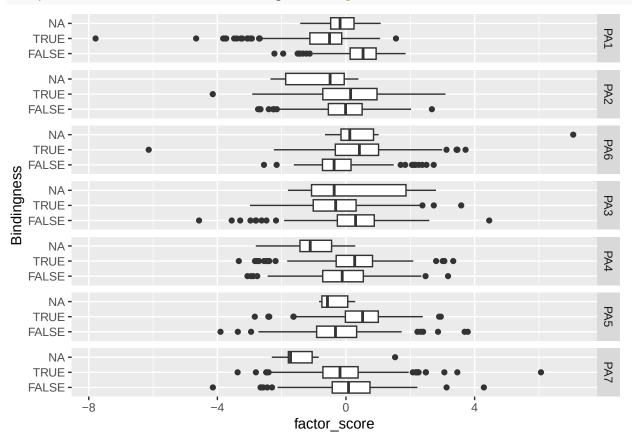
```
##
##
                             Comparison of x by group
                                   (Bonferroni)
##
## Col Mean-|
## Row Mean |
             persuasi
## -----
## quasiobj | -2.072146
                0.0383*
##
           ##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in Objectivity over PA6 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 0.0013, df = 1, p-value = 0.97
##
##
##
                             Comparison of x by group
##
                                   (Bonferroni)
## Col Mean-|
## Row Mean |
             persuasi
## -----
## quasiobj | -0.035709
##
          0.9715
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in Objectivity over PA3 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 1.5825, df = 1, p-value = 0.21
##
##
##
                             Comparison of x by group
##
                                   (Bonferroni)
## Col Mean-
## Row Mean |
             persuasi
## -----
## quasiobj | -1.257979
                 0.2084
##
           ##
## alpha = 0.05
## Reject Ho if p \le alpha
## Test for the significance of differences in Objectivity over PA4 :
##
##
    Kruskal-Wallis rank sum test
##
```

```
## data: x and group
## Kruskal-Wallis chi-squared = 0.0075, df = 1, p-value = 0.93
##
##
##
                             Comparison of x by group
##
                                    (Bonferroni)
## Col Mean-I
## Row Mean |
              persuasi
## -----
## quasiobj | -0.086722
       - 1
                 0.9309
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in Objectivity over PA5 :
##
##
     Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 7.8263, df = 1, p-value = 0.01
##
##
                             Comparison of x by group
                                    (Bonferroni)
##
## Col Mean-|
## Row Mean |
              persuasi
## -----
## quasiobj |
              2.797551
          - 1
                0.0051*
##
##
## alpha = 0.05
## Reject Ho if p \le alpha
## Test for the significance of differences in Objectivity over PA7 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 0.366, df = 1, p-value = 0.55
##
##
                             Comparison of x by group
##
                                    (Bonferroni)
## Col Mean-|
## Row Mean |
               persuasi
## quasiobj |
               0.605013
##
         - 1
                 0.5452
##
## alpha = 0.05
## Reject Ho if p <= alpha
##
## p < 5e-2 found in: PA2 PA5
```

```
## p < 1e-2 found in: PA5
## p < 1e-3 found in:
## p < 1e-4 found in:</pre>
```

Bindingness

```
analyze_distributions(data_factors_long, "Bindingness")
```



```
##
## Test for the significance of differences in Bindingness over PA1 :
##
     Kruskal-Wallis rank sum test
##
##
## data: x and group
  Kruskal-Wallis chi-squared = 322.5625, df = 1, p-value = 0
##
##
##
                               Comparison of x by group
##
                                     (Bonferroni)
## Col Mean-|
   Row Mean |
                    FALSE
##
       TRUE |
                17.96002
##
                 0.0000*
##
            1
## alpha = 0.05
## Reject Ho if p <= alpha
```

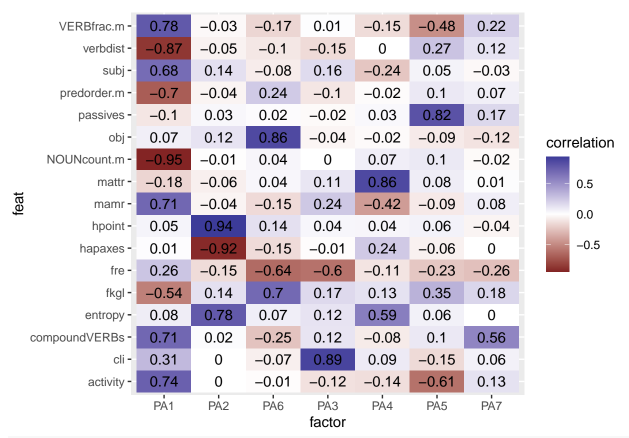
```
##
## Test for the significance of differences in Bindingness over PA2 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 5.0063, df = 1, p-value = 0.03
##
##
                             Comparison of x by group
##
                                   (Bonferroni)
## Col Mean-|
## Row Mean |
                  FALSE
  -----
##
      TRUE | -2.237474
##
           0.0253*
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in Bindingness over PA6 :
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 94.9504, df = 1, p-value = 0
##
##
##
                             Comparison of x by group
                                   (Bonferroni)
##
## Col Mean-|
## Row Mean |
                  FALSE
      TRUE | -9.744250
##
                0.0000*
##
          - 1
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in Bindingness over PA3 :
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 73.3375, df = 1, p-value = 0
##
##
##
                             Comparison of x by group
                                   (Bonferroni)
##
## Col Mean-|
                  FALSE
## Row Mean |
## -----
##
      TRUE | 8.563733
                0.0000*
##
```

```
##
## alpha = 0.05
## Reject Ho if p <= alpha
\#\# Test for the significance of differences in Bindingness over PA4 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 24.7338, df = 1, p-value = 0
##
##
                              Comparison of x by group
                                    (Bonferroni)
##
## Col Mean-|
## Row Mean |
                   FALSE
##
       TRUE | -4.973312
##
            0.0000*
##
## alpha = 0.05
## Reject Ho if p <= alpha
##
## Test for the significance of differences in Bindingness over PA5 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 126.2764, df = 1, p-value = 0
##
##
##
                              Comparison of x by group
                                    (Bonferroni)
##
## Col Mean-|
## Row Mean |
                   FALSE
   -----
##
       TRUE | -11.23727
##
            0.0000*
##
## alpha = 0.05
## Reject Ho if p <= alpha
## Test for the significance of differences in Bindingness over PA7 :
##
##
    Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 17.4925, df = 1, p-value = 0
##
##
##
                              Comparison of x by group
                                    (Bonferroni)
##
## Col Mean-I
## Row Mean |
                   FALSE
```

```
## ------
## TRUE | 4.182400
## | 0.0000*
##
## alpha = 0.05
## Reject Ho if p <= alpha
##
## p < 5e-2 found in: PA1 PA2 PA6 PA3 PA4 PA5 PA7
## p < 1e-2 found in: PA1 PA6 PA3 PA4 PA5 PA7
## p < 1e-3 found in: PA1 PA6 PA3 PA4 PA5 PA7
## p < 1e-4 found in: PA1 PA6 PA3 PA4 PA5 PA7</pre>
```

Feature-factor correlations

```
data_factors_longer <- data_factors_long %>%
  pivot_longer(
    abstractNOUNs:verbdist,
    names_to = "feat", values_to = "feat_value"
  )
data_factors_correlations <- data_factors_longer %>%
  group_by(feat, factor) %>%
  summarize(correlation = cor(feat_value, factor_score))
## `summarise()` has grouped output by 'feat'. You can override using the
## `.groups` argument.
data_factors_correlations %>%
  filter(feat %in% final_collist) %>%
  ggplot(aes(
   x = factor,
    y = feat,
   fill = correlation,
   label = round(correlation, 2)
  )) +
  geom_tile() +
  geom_text() +
  scale_fill_gradient2()
```



```
data_factors_correlations %>%
  filter(!(feat %in% final_collist)) %>%
ggplot(aes(
    x = factor,
    y = feat,
    fill = correlation,
    label = round(correlation, 2)
)) +
geom_tile() +
geom_text() +
scale_fill_gradient2()
```

