Running a Simple Model & Finding a Routine ${}^{Adam\ Garber}$

Norwegian University of Science and Technology - A Course in ${\tt MplusAutomation}$

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Install the "rhdf5" package to read gh5 files (plot information of the package to read gh5 files (plot information).

```
if (!requireNamespace("BiocManager", quietly = TRUE))
  install.packages("BiocManager")
BiocManager::install("rhdf5")
```

How to install packages?

```
install.packages("tidyverse")
```

Load packages

```
library(tidyverse)
library(haven)
library(here)
library(MplusAutomation)
library(rhdf5)
```

Read in data

```
# object_name <- function_1("dataset_name.sav")
exp_data <- read_spss("https://garberadamc.github.io/project-site/data/explore_lab_data.sav")</pre>
```

View dataframe with labels & response scale meta-data

Note: Use the "print" option to save a PDF as a codebook of metadata.

```
# the {haven} package keeps the meta-data from SPSS files

# package_name::function_within_package()

sjPlot::view_df(exp_data)
```

Types of data for different tasks

- .sav (e.g., spss_data.sav): this data format is for SPSS files & contains variable labels (meta-data)
- .csv (e.g., r_ready_data.csv): this is the preferable data format for reading into R & when estimating models with MplusAutomation (non-labeled data)
- .dat (e.g., mplus_data.dat): this is the data format used to read into Mplus (no column names or strings)

NOTE: Mplus also accepts .txt formatted data (e.g., mplus_data.txt)

Writing, reading, and converting data between 3 formats

Location, location!

NOTE: default directory in an Rproject is the "top-most" project folder

```
here()
```

[1] "/Users/agarber/github/NTNU-workshop"

```
project_location <- here()</pre>
```

Change starting location within the Rproject (.Rproj)

- 1. save stem
- 2. glue() stem into subsequent labs

```
source("rep_functions.R")
change_here(glue("{project_location}/02-run-models"))
here()
```

[1] "/Users/agarber/github/NTNU-workshop/02-run-models"

Prepare datasets, remove SPSS labeling

```
# write a CSV datafile (preferable format for reading into R; !! removes labels !!)
write_csv(exp_data, here("data", "exp_lab1_data.csv"))

# write a SPSS datafile (preferable format for reading into SPSS, labels are preserved)
write_sav(exp_data, here("data", "exp_lab1_data.sav"))

# read the unlabeled data back into R
nolabel_data <- read_csv(here("data", "exp_lab1_data.csv"))

# write a DAT datafile (this function removes header row & converts missing values to non-string)
prepareMplusData(nolabel_data, here("data", "exp_lab1_data.dat"))</pre>
```

THINGS TO NOTE:

- 1. This function produces a minimal template of input syntax for an Mplus input file.
- 2. Behind the scenes mplusObject() will use a similar function to produce an input file & .dat file from the R data.frame that the function is given.
- 3. By default missing values in your R object (NA) are converted to .
- 4. Note that if the following file.string is greater than 90 characters (Mplus limit) your model will not estimate and return an error message.

DATA: FILE = "/Users/agarber/github/NTNU-workshop/02-run-models/data/exp_lab1_data.dat";

Preparing column-names to be MplusAutomation ready

Task: Make all variable names fit within the 8-character name limit (Mplus) while avoiding duplicates.

Renaming columns manually...

What do you do if you have a large dataset with many column names that are > 8 characters?

- first, remove all characters greater than 8 using str_sub()
- second, make sure you don't now have duplicate variable names
- third, locate and change all duplicate names

NOTE: Use regular expressions to remove strings in a more systematic manner to avoid duplicates.

```
# remove characters from the variable names that are greater than 8 characters
names(new_names) <- str_sub(names(new_names), 1, 8)

# check if culumn names are unique
test.unique <- function(df) { ## function to identify unique columns

length1 <- length(colnames(df))
length2 <- length(unique(colnames(df)))
if (length1 - length2 > 0 ) {
   print(paste("There are", length1 - length2, " duplicates", sep=" "))
}

test.unique(new_names)
```

[1] "There are 6 duplicates"

```
# locate duplicates (this will find the first duplicated column)
anyDuplicated(colnames(new_names))
```

[1] 2

A minimal example of writing, running, & reading models

What does the mplusObject() function do:

- 1. It generates an Mplus **input** file (does not need full variable name list, its automated for you!)
- 2. It generates a data file (.dat) specific to each model
- 3. It runs or estimates the model (hopefully) producing the correct output. Always check!

NOTE: Within the mplusObject() function there is a mix of R and Mplus syntax. In general the black colored text will be R code and the green colored text within the quotation marks will be Mplus code (exception; when using {glue}).

PRACTICE: Using MplusObject() method

Model is type = BASIC; (i.e., returns descriptive stats)

Check your model (Always!)

- In the bottom-right pane under the files tab click on the basic_mplus folder
- There should be 3 new files in this location that were produced by mplusObject()
- Click on the .out file to check if the model estimated or if there are any error messages

PRACTICE SUBSETTING: Now explore descriptives for observations that reported as "female"

Add line of syntax: useobs = female == 1;

```
fem_basic <- mplusObject(
   TITLE = "PRACTICE 02 - Explore female observations only",
   VARIABLE =
   "usevar=
   item1 item2 item3 item4 item5
   item6 item7 item8 item9;

   useobs = female == 1; !include observations that report female in analysis",

   ANALYSIS =
   "type = basic;",

   usevariables = colnames(nolabel_data),
   rdata = nolabel_data)

fem_basic_fit <- mplusModeler(fem_basic,</pre>
```

```
dataout=here("basic_mplus", "fem_basic.dat"),
modelout=here("basic_mplus", "fem_basic.inp"),
check=TRUE, run = TRUE, hashfilename = FALSE)
```

PRACTICE: Exploratory Factor Analysis (EFA)

```
## EXPLORATORY FACTOR ANALYSIS LAB DEMONSTRATION
efa_demo <- mplusObject(</pre>
 TITLE = "EXPLORATORY FACTOR ANALYSIS - LAB DEMO",
 VARIABLE =
 "usevar=
 item1 item2 item3 item4 item5
 item6 item7 item8 item9;" ,
 ANALYSIS =
 "type = efa 1 5;
 estimator = MLR;
 parallel=50;",
 MODEL = "",
 PLOT = "type = plot3;",
 OUTPUT = "sampstat standardized residual modindices (3.84);",
 usevariables = colnames(nolabel_data),
 rdata = nolabel_data)
efa_demo_fit <- mplusModeler(efa_demo,</pre>
 dataout=here("basic_mplus", "EFA_Lab_DEMO.dat"),
 modelout=here("basic_mplus", "EFA_Lab_DEMO.inp"),
check=TRUE, run = TRUE, hashfilename = FALSE)
```

END