visualizemi: Visualization, Effect Size, and Replication of Measurement
Invariance for Registered Reports

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Author Note

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Thank you to K.D. Valentine and Chelsea Parlett-Pelleriti for feedback on some ugly graphs.

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11 Abstract

Latent variable modeling as a lens for psychometric theory is a popular tool for social 12 scientists to examine measurement of constructs (Beaujean, 2014). Journals such as 13 Assessment regularly publish articles supporting measures of latent constructs wherein a 14 measurement model is established. Confirmatory factor analysis can be used to investigate 15 the replicability and generalizability of the measurement model in new samples, while 16 multi-group confirmatory factor analysis is used to examine the measurement model across 17 groups within samples (Brown, 2015). With the rise of the replication crisis and 18 "psychology's renaissance" (Nelson et al., 2018), interest in divergence in measurement has 19 increased, often focused on small parameter differences within the latent model. This 20 manuscript presents visualizemi, an R package that provides functionality to calculate multigroup models, partial invariance, visualizations for (non)-invariance, effect sizes for models and parameters, and potential replication rates compared to random models. Readers will learn how to interpret the impact and size of the proposed non-invariance in 24 models with a focus on potential replication and how to plan for registered reports.

Keywords: multigroup confirmatory factor analysis, measurement invariance,
 visualization, effect size

visualizemi: Visualization, Effect Size, and Replication of Measurement Invariance for Registered Reports

Psychological assessments play a critical role in our ability to measure and analyze 30 constructs to support theories and experimental hypotheses. Defining and creating 31 assessments to validly and reliability measure constructs is often difficult because 32 phenomenon, such as anxiety, are often not directly observable. Instead, we use surveys and 33 questionnaires to indirectly assess the underlying construct (DeVellis & Thorpe, 2022). 34 Latent variable modeling (i.e., structural equation modeling) is a popular tool for the validation of developed survey instruments to verify scale dimensionality, structure, and model fit. A simple search for scale development reveals thousands of articles in psychology that examine new and previously published work, thus, illustrating the interest in both measurement and the use of validation techniques. Unfortunately, except in specialty journals, much of the validity evidence and/or development for measures used in empirical studies is not reported within the journal article (Barry et al., 2014; Weidman et al., 2017). Without this information, it is difficult to interpret individual study conclusions, as validity information allows for judgment of usefulness of the measured values (Flake & Fried, 2020). 43 Further, the current focus on replication (Makel et al., 2012; Makel & Plucker, 2014; Zwaan et al., 2018), reproducibility (Nelson et al., 2018), and the credibility of our results (Vazire et 45 al., 2022) has demonstrated questionable measurement practices - decisions that researchers make like survey selection and scoring that impact the results of the study (Flake & Fried, 2020). Transparent reporting of the use and creation of scales can improve both interpretation and reproducibility when using surveys developed to measure latent constructs (Shadish et al., 2001).

A secondary concern for developed measures is the potential for differential responding and assessment within target populations. For example, Trent et al. (2013) examined for potential variability in the Revised Child Anxiety and Depression Scale in

White and Black youths (Chorpita et al., 2000). They found that the scale mostly

functioned the same for both White and Black individuals but differences in averages on

individual items could potentially affect the scoring and interpretation of the scale results.

This comparison of sub-populations is the test of measurement invariance (Meredith, 1993).

Invariance or equivalence implies that the scale operates in the same fashion for each

sub-group, and thus, differences in the final latent variable scores can interpreted as

differences in populations. Non-invariance suggests that individuals respond or interpret

items differently, and thus, differences in scores may represent different scores on the latent

variable in the population or differences in measurement. Non-invariant measurement may

lead to misleading results when making group comparisons, and assessing invariance has

become a popular technique in scale development (Van De Schoot et al., 2015).

Measurement invariance has been explored and implemented for the last fifty years 65 (Jöreskog, 1971; Sörbom, 1978) and implemented in the most popular structural equation 66 modeling programs (Boker et al., 2011; Jöreskog & Sörbom, 2001; Rosseel, 2012). Byrne et 67 al. (1989) extended the ideas of multi-group testing by suggesting partial invariance 68 (followed by Meredith, 1993). Partial invariance occurs when non-invariance is found but can 69 be attributed to only a few parameter estimate differences between groups (i.e., items 1 and 2 have different factor loadings but all others are the same). This testing provided an 71 advantage to understand where the potential non-invariance may occur for further study and 72 interpretation guidelines. As the field pushes back against favoring cutoff criteria and rules of thumb (Marsh et al., 2004; Putnick & Bornstein, 2016), an effect size measure for translating "how much" non-invariance was developed d_{MACS} (Nye & Drasgow, 2011). This effect size examines the differences in observed variables between the two groups for both the factor loading and the item intercept; thus, any differences in either or both will increase the effect size for non-invariance (Stark et al., 2006).

With d_{MACS} and measurement invariance testing, researchers can begin to quantify

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how and where their construct measurement may vary between groups. Yet, given the large number of studies that show non-invariance, it is clear that equivalence can be hard to meet. 81 It is difficult to know if non-invariance occurs because of random sampling error, true 82 population differences, or differences in replication and reproducibility of the construct in a 83 new sample. The field of psychology is increasingly interested in pre-registration (i.e., registering plans for a study before data collection, Nosek et al., 2018) and the promotion of 85 transparency in study design, implementation, and analysis (Mayo-Wilson et al., 2021), in addition to supporting replication studies (Zwaan et al., 2018). Registered (replication) reports provide an advantageous avenue for the pre-registration of measurement tests, as they allow a researcher the ability to have their study accepted in principle, regardless of the results of a test of construct validity, reliability, or measurement invariance (Hobson, 2019; Nosek & Lakens, 2014). However, there are few tools that can provide effect size measures for models, individual parameters, or visualization for researchers to plan for future studies. d_{MACS} provides the opportunity to begin to think about the smallest effect size of interest or the smallest meaningful effect size for measurement invariance and replication (Anvari & Lakens, 2021; i.e., two studies with overlapping confidence intervals "replicate," even if the test of measurement invariance does not, Lakens, 2017). As mentioned, d_{MACS} has only really been explored for a combined intercept and loadings, and while useful, does not necessarily allow a researcher to pinpoint specific issues within an observed variable. 98

Therefore, purpose of this manuscript is describe an R package, visualizemi, that gg provides functionality to calculate multi-group confirmatory factor analysis, partial 100 invariance tests, visualizations of the size of non-invariance, and potential effect sizes for 101 overall models and individual parameters. No known visualization techniques have been 102 proposed for measurement invariance. By creating panel visualizations, we can supplement a 103 researchers ability to judge the strength of the non-invariance differences and effect size for 104 each item. The proposed effect sizes demonstrate the likelihood of replication with a similar 105 sample as compared to a randomly assigned group model, thus, illustrating what type of 106

measurement one might expect to find, and how different that is from random chance. 107 Within this technique, the individual parameter effect sizes can calculated: both the group 108 differences within a model as compared to random and the likelihood of a parameter 109 replication compared to random groups. Coupled with other indicators (i.e., fit indices 110 differences, d_{MACS}), we can move toward a better understanding of how much measurement 111 non-invariance is meaningful. This tutorial and package will help researchers plan future 112 studies and aid in the ability to estimate a smallest effect of interest for measurement 113 invariance studies, rather than relying on fit indices and rules of thumb alone. 114

By the end of this tutorial manuscript, readers will:

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- 1. Learn how to use *visualizemi* to analyze multi-group confirmatory factor analysis, examine partial invariance, and create visualizations of parameters.
- 2. Learn how to estimate the potential replication of multi-group models and their parameters using bootstrapping compared to a random group model.
- 3. Be able to calculate and interpret effect sizes for model and parameter replication, as well as parameter group differences.
 - 4. Understand the impact of measurement variability on replication and generalizability.

The tutorial will start with simulated data based on known effect sizes using d_{MACS} and demonstrating the package functions for 1) running the multigroup analysis, 2) running a partial invariance analysis, 3) plotting the partial invariance, 4) estimating replication and effect sizes at the model level, and 5) estimating replication and effect sizes at the parameter level. Last, data from Aiena et al. (2014) examining the measurement invariance of the RS-14 (Wagnild, 2009) will be used to demonstrate the application of the package on real data. The *visualizemi* package vignette includes an additional tutorial walk through.

130 Method

Design and Analysis

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Data was simulated using the simulateData function in the R package lavaan132 (Rosseel, 2012) assuming multivariate normality using a μ of 0 and σ of 1 for the data. This 133 function allows you to write lavaan syntax for your model with estimated values to generate data for observed variables (see supplemental for examples). The data included two groups 135 of individuals ("Group 1", "Group 2") for a multi-group confirmatory factor analysis (n_{group}) 136 =250, N=500). The latent variables were assumed to be continuous normal (the package functions do not require this assumption). The model consisted of five observed items 138 predicted by one latent variable (1v = q1 + q2 + q3 + q4 + q5); however, the 139 demonstration in this manuscript extends to multiple latent variables and other 140 combinations of observed variables. Each item was assumed to be related to the latent 141 variable with loadings approximately equal to .40 to .80, except when cases of non-invariance 142 on the loadings was simulated. 143

The Brown (2015) steps of testing measurement invariance are demonstrated in this manuscript for illustration purposes, but in line with Stark et al. (2006) suggestions, the 145 visualizations show the impact of loadings and intercepts together. A convenience function 146 mgcfa is used for these steps or other measurement invariance test orders and combinations. 147 Fit indices for the steps for multi-group models are presented in the appendix for comparison 148 of cutoff rules of thumb (Cheung & Rensvold, 2002) to effect sizes and visualizations 149 presented in this manuscript. Fit indices include Akaike Information Criterion (AIC, Akaike, 150 1998), Bayesian Information Criterion (BIC, Schwarz, 1978), Comparative Fit Index (CFI, 151 Bentler, 1990), Tucker Lewis Index (TLI, Tucker & Lewis, 1973), root mean squared error of 152 approximation RMSEA (Steiger, 1990), and standardized root mean square residual (SRMR, 153 Bentler, 1995). 154

The data was then simulated to represent invariance across all model steps, small,

medium, and large invariance using d_{MACS} estimated sizes from Nye et al. (2019). While 156 d_{MACS} is used primarily for an effect size of the (non)-invariance for intercepts and loadings 157 together, a similar approach was taken for the estimation of small, medium, and large effects 158 on the residuals. The effect size is presented for all models, calculated from the dmacs 159 package (Dueber, 2023; Nye & Drasgow, 2011). Only one item in each model was 160 manipulated from the invariant model to create the non-invariant models. Given the data 161 was simulated with a z-score scaling, the loading values were simulated at .30 points apart 162 (given d_{MACS} suggestions of .2, .4, .7), the intercepts at .25 points apart, and the residuals at 163 .25 points apart. To plan a simulation for your own study, these values can be used to 164 simulate small, medium, and large non-invariance effects by first converting data into z-score. 165

Package Code Examples

The complete code for this manuscript can be found at https://osf.io/wev5f/. This tutorial was registered at https://osf.io/vwf4d, and the example provided at the end of the manuscript was added after that registration. The R package and replication/effect sizes was added after the original manuscript submission.

Multi-group CFA Caculation

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First, we would create our model code in *lavaan* syntax (Rosseel, 2012). The lv latent variable predicts the five measured variables, which are present as columns in our df.invariant data set. The package does generally require raw data for bootstrapping purposes, and an example of how to simulate data from models and covariance/correlations tables sometimes provided in manuscripts (rather than the raw data) is provided in the supplemental documentation.

lavaan automatically sets the mean (i.e., the intercept) for latent variables to zero. If
we wish to visualize the impact of the changes in parameter estimates across groups on the
latent means, we need to allow the latent mean estimation with $lv \sim 1$. However, adding
this estimation into our model will create a non-identified model. To solve this problem, you

can set one of the intercepts of another variable to a value to scale the model. Here we will
set the scale of the model by using q1 ~ 0*1, thus, scaling the expected means to zero.
With simulation, this step is easy to know which variable to pick - we set the intercept on
the variable we know did not show differences. In real data, you may wish to run the model
steps without setting this option, examine the results of a configural or separate models, and
then add the option for the values most similar. Additionally, you could complete partial
invariance steps to determine which value appears most consistent to fix the estimate.

```
# create lavaan model
model.overall <- "
# overall one-factor model
lv =~ q1 + q2 + q3 + q4 + q5
# set the intercept (mean) of q1 to zero
q1 ~ 0*1
# allow the lv intercept to be freely estimated
lv ~ 1"
# look at the data
head(df.invariant)</pre>
```

```
##
                q1
                             q2
                                          q3
                                                     q4
                                                                 q5
                                                                      group
189
   ## 1 -0.8903542 -0.81707530 0.06137292 -1.3236407 -1.7916418 Group 1
190
         1.1054521 -0.03540948 -0.81299606 1.0028340 -0.1909127 Group 1
   ## 2
191
   ## 3
         1.4555852
                     1.54083484
                                 1.59084213 -0.3345967 -0.6865496 Group 1
192
   ## 4 -1.8745187 -1.27880245 -2.53565792 -1.0024193 -1.6253249 Group 1
193
                                1.05507079 -1.2615705 1.7536428 Group 1
   ## 5 -0.4449517 -0.17782974
194
                                1.63251893  0.6449847 -1.0055700 Group 1
         0.2278813 0.71348845
195
```

The mgcfa function is designed to flexibly allow you to leverage lavaan's package functions to calculate multiple measurement steps at once. You would include:

- 1) the model syntax in the model argument
- 199 2) the dataframe in the data argument of our function
- 200 3) the name of the grouping variable in quotes for group
- 4) and the equality constraints you would like to impose in order in group.equal
- 5) ... any other *lavaan* arguments you would like to use such as meanstructure or estimator.

Note: you can also use sample.cov, sample.mean, sample.nobs in this step for estimation of multi-group models, but simulated dataframes are needed for bootstrapping replication estimates.

```
# run our mgcfa function to run all models

results.invariant <-
    # name of the saved model syntax

mgcfa(model = model.overall,
    # name of the dataframe
    data = df.invariant,
    # name of the grouping variable
    group = "group",
    # equality constraints to impose in order
    group.equal = c("loadings", "intercepts", "residuals"),
    # other options to send to lavaan cfa function
    meanstructure = T)

# what is saved for you
names(results.invariant)</pre>
```

The following output is saved:

1) model_coef: The parameter estimates for each model with the model step included in
a model column. This set of coefficients can be used for other functions. This
dataframe is created with broom's tidy() function if you wish to recreate this table
without running the mgcfa() function (Robinson et al., 2023).

```
results.invariant$model_coef[1:10 , ]
```

```
## # A tibble: 10 x 13
                                                                        std.lv std.all std.nox
   ##
                           estimate std.error statistic
                                                              p.value
215
          term
                    op
                                                                          <dbl>
   ##
          <chr>>
                    <chr>>
                               <dbl>
                                          <dbl>
                                                      <dbl>
                                                                 <dbl>
                                                                                   <dbl>
                                                                                            <dbl>
216
        1 "lv =~ ~ =~
                                                                         0.803
   ##
                              1
                                         0
                                                    NA
                                                            NA
                                                                                  0.616
                                                                                           0.616
217
        2 "lv =~ ~ =~
                                                     7.44
                                                                                           0.493
                             0.655
                                         0.0880
                                                             9.77e-14
                                                                         0.526
                                                                                  0.493
218
   ##
   ##
        3 "lv =~ ~ =~
                             0.640
                                         0.0895
                                                     7.15
                                                             8.83e-13
                                                                        0.514
                                                                                  0.463
                                                                                           0.463
219
        4 "lv =~ ~ =~
                                                                                           0.209
                             0.277
                                         0.0749
                                                     3.69
                                                             2.24e- 4
                                                                        0.222
                                                                                  0.209
   ##
220
        5 "lv =~ ~ =~
                             0.955
                                                             4.44e-16
                                                                         0.766
   ##
                                         0.117
                                                     8.13
                                                                                  0.656
                                                                                           0.656
221
        6 "q1 ~1 " ~1
                             0
                                         0
                                                    NA
                                                            NA
                                                                         0
                                                                                  0
                                                                                           0
   ##
222
        7 "lv ~1 " ~1
                            -0.0305
                                                    -0.524
                                                             6.00e- 1 -0.0380 -0.0380 -0.0380
   ##
                                         0.0582
223
        8 "q1 ~~ ~ ~~
   ##
                             1.05
                                         0.0995
                                                    10.6
                                                             0
                                                                         1.05
                                                                                  0.620
                                                                                           0.620
224
        9 "q2 ~~ ~ ~~
                             0.860
                                         0.0653
                                                    13.2
                                                             0
                                                                         0.860
                                                                                  0.757
                                                                                           0.757
225
   ## 10 "q3 ~~ ~ ~~
                             0.966
                                         0.0711
                                                    13.6
                                                             0
                                                                         0.966
                                                                                  0.785
                                                                                           0.785
226
   ## # i 4 more variables: model <chr>, block <int>, group <int>, label <chr>
227
```

228 2) model_fit: The model fit indices from fitmeasures() to review for overall model fit
and invariance judgments. The name of the model is included in a model column.

```
head(results.invariant$model_fit)
```

230 ## # A tibble: 6 x 18

```
##
                 AIC
                        BIC
          agfi
                               cfi
                                    chisq
                                           npar
                                                  rmsea rmsea.conf.high
                                                                                      tli
231
         <dbl> <dbl> <dbl> <dbl> <
   ##
                                    <dbl> <dbl>
                                                   <dbl>
                                                                    <dbl>
                                                                             <dbl> <dbl>
232
   ## 1 0.998 7516. 7580. 1
                                    0.650
                                              15 0
                                                                   0
                                                                           0.00616 1.04
233
   ## 2 0.948 3766. 3819. 0.976
                                    7.79
                                              15 0.0473
                                                                           0.0312
                                                                   0.108
                                                                                   0.953
234
   ## 3 0.974 3768. 3820. 1
                                    4.48
                                              15 0
                                                                   0.0831 0.0210
                                                                                    1.01
235
   ## 4 0.961 7533. 7660. 0.991 12.3
                                                                   0.0785 0.0261
                                              30 0.0301
                                                                                    0.982
236
   ## 5 0.965 7528. 7638. 0.994 15.4
                                              26 0.0200
                                                                   0.0660 0.0330
                                                                                    0.992
237
   ## 6 0.969 7522. 7615. 1
                                              22 0
                                                                   0.0542 0.0352
                                                                                    1.00
238
```

- 239 ## # i 8 more variables: converged <lgl>, estimator <chr>, ngroups <int>,
- 240 ## # missing method <chr>, nobs <int>, norig <int>, nexcluded <int>, model <chr>
- 241 3) model_overall: A saved lavaan fitted model of all groups together without any

 equality constraints or grouping variables. These objects can be used with any function

 that normally takes a saved model: parameterEstimates(),
- modificationIndices(), semPlot::semPaths(), and so on (Epskamp, 2022).
- 4) group_models: A list of saved fitted models for each group separately.
- 5) model_configural: A saved fitted model for the configural model that nests together each group into one model with no other constraints.
- of invariance_models: A list of saved fitted models that consecutively adds group.equal constraints.

Visualization of Invariance

251 Package Function

- The results from the model_coef table can then be used directly in plot_mi(). The plot outputs will be described below. First, here are the arguments for the function:
- 1) data_coef: A tidy dataframe of the parameter estimates from the models. This

 function assumes you have used broom::tidy() on the saved model from lavaan and

 added a column called "model" with the name of the model step (Robinson et al.,

2023). This function will only run for models that have used the grouping function (i.e., configural, metric, scalar, and strict or other combinations/steps you wish to examine).

- 259 2) model_step: Which model do you want to plot? You should match this name to the one you want to extract from your model column in the data_coef.
- 3) item_name: Which observed variable from your model syntax do you want to plot?

 Please list this variable name exactly how it appears in the model.
- 4) x_limits: What do you want the x-axis limits to be for your invariance plot? The
 default option is to assume the latent variable is standardized, and therefore, -1 to 1 is
 recommended. Use only two numbers, a lower and upper limit. This value also
 constrains the latent mean diagram to help zoom in on group differences because the
 scale of latent means is usually centered over zero. You can use this parameter to zoom
 out to a more traditional histogram using c(-2, 2).
- 5) y_limits: What do you want the y-axis limits to be for your invariance plot? Given
 that the latent variable is used to predict the observed values in the data, you could
 use the minimum and maximum values found in the data. If that range is large,
 consider reducing this value to be able to visualize the results (i.e., otherwise it may be
 too zoomed out to judge group differences). Use only two numbers, a lower and upper
 limit.
- conf.level: What confidence limit do you want to plot? Use 1 α .
- 7) model_results: In this argument, include the saved *lavaan* output for the model listed in the model_step argument.
- 278 8) lv_name: Include the name of the latent variable, exactly how it is listed in your

 279 lavaan syntax. You should plot the latent variable that the item_name is linked to. If

 280 you have items that load onto multiple latent variables, you will need to make multiple

 281 plots.
- 9) plot_groups: If you include more than two groups in a multi-group model, the
 automatic assumption is that you want the first two groups for this visualization. If

not, include the names of the groups here to plot.

284

```
invariant.plot <-</pre>
 plot_mi(
    # output from model_coef
    data_coef = results.invariant$model_coef,
    # which model do you want to plot
    model_step = "Configural",
    # name of observed item
    item_name = "q4",
    # latent variable limits to graph
    x_{limits} = c(-1,1),
    # Y min and max in data
    y_limits = c(min(df.invariant$q4), max(df.invariant$q4)),
    # what ci do you want
    conf.level = .95,
    # what model results do you want
    model_results = results.invariant$model_configural,
    # which latent variable do you want
    lv name = "lv"
)
names(invariant.plot)
```

```
285 ## [1] "complete" "intercept" "mean" "variance"
```

The outputs from this function are several *ggplot2* objects that can be edited or saved directly using *ggplot2* functionality (Wickham, 2016).

288 1) complete: The output from this model can be found in Figure 1. On the left hand side, the item invariance is plotted, and on the right hand side, the latent mean distributions

for the two groups are plotted. In the item invariance sub-plot, the visualization includes all three components traditionally seen in MGCFA testing steps: loadings, intercepts, and residuals. Each visualization element was designed to match the traditional visualization for that type of output. All parameter estimates are plotted on the unstandardized estimates and their confidence interval based on the standard error of the estimate. All plots are made with *qqplot2* and *cowplot* (Wilke, 2020).

2) intercept: Only the left hand side of the complete plot designed to represent intercepts and factor loadings. Factor loadings represent the slope of the regression equation for the latent variable predicting the scores on the observed variable $(\hat{Y} \sim b_0 + b_1 X + \epsilon)$. The y-axis indicates the observed variable scores, and here, the plot includes the entire range of the scale of the data for item four. The coefficient (b_1) for group 1 was 0.40, while the coefficient for group 2 was 0.21. The ribbon bands around the plotted slopes indicate the confidence interval for that estimate. In this plot, while the coefficients for each group are not literally equal, the overlapping and parallel slope bands indicate they are not different practically.

The item intercepts (b_0) are plotted on the middle line where they would cross the y-axis at a latent variable score of zero. These are represented by a dot with a set of confidence error bars around the point. The intercept for group 1 was 0.07, while the coefficient for group 2 was 0.03. In this invariant depiction, the overlap in the intercepts is clear, indicating they are not different. You can use y_limits to zoom in on the graph if these are too small to be distinguishable.

3) mean: The right hand side of the complete plot graphing the latent variable means and density from the data. The latent variable is shown on the x-axis using standardized values (i.e., z-scores) where -1 indicates one standard deviation below the mean for the latent variable, 0 indicates the mean for the latent variable and so on. The lines indicate the means of the latent variables from the simulated dataset. Group labels are

represented in the figure caption on the bottom. Group 1 is usually the group that is alphabetically first in the data set or whichever group is the first that appears when using the levels() command.

4) variance: A split geom violin plot indicating the variance distribution of the plotted item. Residuals are trickier to plot, as they are the left over error when predicting the observed variables ε. It is tempting to plot this value as the confidence band around the slope, however, that defeats the purpose of understanding that the slopes are estimated separately from the residuals, and both have an associated variability around their parameter estimate. Therefore, residuals are represented in the inset picture at the bottom right of the item invariance plot. The black bars represent the estimated residual for each group (group 1: 0.91, group 2: 1.22). The distributions are plotted to represent the normal spread of values using the standard error of the residuals. The violin plot allows for direct comparison of those residuals and their potential distributions. Note that the placement has nothing to do with the x or y-axis and is designed to always show in the same location, regardless of size/value. The plots are included separately so they can be arranged in a different fashion if desired.

$Simulated \ Results$

The d_{MACS} value for item 4 in the invariant model was 0.16, representing a nil or unimportant difference in this manuscript. It is important to note that while Nye et al. (2019) suggests specific sizes for small, medium, and large, each researcher should determine for themselves what effects represent. Figure 2 displays the results from the small (d_{MACS} = 0.27) difference in loadings, while Figure 3 displays the results from the medium ($d_{MACS} =$ 0.53) difference in loadings, and Figure 4 shows the large ($d_{MACS} = 0.68$) differences. When investigating the slope values, we can clearly see the change in the loading for the second group (the only manipulated variable, although random data set generation may also change intercepts and residuals slightly). At the medium effect size, we see that the confidence

bands do not overlap (at the edges), and at the large effect size, we can see a clear separation of two lines. Note that the intercepts in this model are estimated as equal so the loading 343 representation will not literally separate, but the steepness of the lines is the indicator of the 344 difference between the slopes. You can imagine these lines are interpreted like a simple 345 slopes analysis for interactions in regression (Cohen et al., 2003). When simple slopes for 346 interactions are plotted, if they are parallel, there is no interaction, and if they cross, then 347 there is an interaction. Here, we can use this same logic. If they are parallel, there is likely 348 invariance (they are the same), and the further from parallel they become, the larger the effect size for the differences between group loadings. 350

The latent means in Figure 4 do appear to show differences, albeit visually small. 351 The latent means diagram shows the impact of any group differences that aren't constrained, 352 and this image shows the configural model (as the metric model would force them to be 353 equal). In the simulated model, the *only* manipulated parameter is question 4's loading. In 354 real models, the differences may be larger due to other variation found in the parameter 355 estimates. Therefore, once you discover items you believe would make a model "partially" 356 invariant, you may wish to estimate that model and graph the item again using the partially 357 invariant model to see only the effect of the non-invariant items. Additionally, consider that 358 we set the scaling of the model to 0. The estimate for the ly mean in the large loading model 359 was group 1: 0.00, and group 2: -0.06, which results in 0.06 difference in group means. The 360 practical implications of this difference will depend on the research and interpretations of the 361 researcher. 362

For intercepts, the small (Figure 5), medium (Figure 6), and large (Figure 7)
depictions represent d_{MACS} values of 0.26, 0.47, and 0.70, respectively. Intercept differences
can be clearly seen represented by the spacing out of the intercept locations (and thus, the
overall line as well). While the changes in intercept do not appear to change the latent
means, the caveat to this simulation is that only item four was manipulated. An example is

provided below that demonstrates large changes in latent means.

Last, the effect of the residuals is plotted in small (Figure 8), medium (Figure 9), and 369 large (Figure 10) formats. While d_{MACS} values are not technically available for the residuals, 370 our models showed 0.19, 0.19, and 0.16, respectively. These differences in values are variable 371 due to the random generation of data sets for each measurement invariance manipulation. At 372 first glance, the differences in the small chart may seem large, because the black lines are not 373 touching, but notice that the distributions overlap, indicating a likely small difference. The 374 medium and large differences better illustrate differences in residuals across groups. Further, 375 the impact of the residuals on the shape of the latent mean distribution can also been seen 376 (and unintentionally, in the first figures as well due to random variation). The impact is due 377 to the standard error of the residuals, as smaller standard errors represent lepokurtic 378 distributions (taller), and larger standard errors represent platykurtic distributions (flatter). 379 The effect size difference of the residuals does not appear to change the effects in the latent 380 means.

382 Partial Invariance Calculation

383 Package Function

387

388

The results of the simulated models are presented in the appendix, demonstrating
that each simulated dataset shows partial invariance if item four is allowed to vary between
groups. The function takes the following arguments:

- 1) saved_model: The saved *lavaan* model with the equality constraints at the level of measurement invariance you would like to examine for partial invariance.
- 2) data: The dataframe where the model was estimated.
- 390 3) model: The model syntax for the overall model.
- 391 4) group: The grouping variable column in the dataframe.
- 5) group.equal: The equality constraints including in your original multi-group tests.
- 6) partial step: The level of partial invariance you wish to test.

```
partial.invariant <-
  partial_mi(
    # saved model output with constraints
    saved_model = results.invariant$invariance_models$model.residuals,
    # dataframe from model
  data = df.invariant,
    # model syntax
  model = model.overall,
    # group column name
  group = "group",
    # group equality constraints from your mgcfa
  group.equal = c("loadings", "intercepts", "residuals"),
    # which step you want to examine for partial invariance
  partial_step = "residuals"
  )

names(partial.invariant)</pre>
```

```
394 ## [1] "models" "fit_table"
```

In this function, each parameter with the appropriate *lavaan* syntax is relaxed individually (i.e., ~1 for intercepts, ~~ for residuals, etc.). The fitted models are saved in the models output, and the fit_table output includes all fit indices for each model to investigate potential areas of partial invariance based on the researcher's desired criterion.

Note: the partial_step function is used to determine which types of op or operators to freely estimate between groups. If one chooses residuals, you will also freely estimate the residual for the latent variable or any other residuals found in the model. These items may be ignored if they were not meant to be included.

```
head(partial.invariant$fit_table %>%

dplyr::select(free.parameter, cfi, rmsea))
```

```
## # A tibble: 6 x 3
403
         free.parameter cfi
                                      rmsea
404
   ##
         <chr>
                         <lvn.vctr> <lvn.vctr>
405
   ## 1 q1 ~~ q1
                         0.9902679
                                      0.02108648
406
   ## 2 q2 ~~ q2
                         0.9868905
                                      0.02447336
407
   ## 3 q3 ~~ q3
                         0.9958241
                                      0.01381266
408
   ## 4 q4 ~~ q4
                         1.0000000
                                      0.0000000
409
   ## 5 q5 ~~ q5
                         0.9868088
                                      0.02454944
410
   ## 6 lv ~~ lv
                         0.9906154
                                      0.02025143
411
```

Replication and Effect Size: Model

413 Package Function

- The bootstrap_rr function in *visualizemi* was designed to estimate the likely replication of overall model invariance with the assumption that the data used for the estimation represents the larger population. The following arguments are used:
- 1) saved_configural: a saved fitted model at the configural level with no equality
 constraints. This model should include all other lavaan settings you would like to use,
 such as estimator or ordered.
- 2) data: The dataframe where the model was estimated.
- 3) model: The model syntax for the overall model.
- 422 4) group: The grouping variable column in the dataframe.
- 5) nboot: The number of bootstraps to run.
- 6) invariance_index: The fit index you would like to use to determine invariance.
- Please use options and labeling from lavaan see fitmeasures() for options.

7) invariance_rule: The invariance difference score you would like to use as your rule.

426

427

8) group.equal: The equality constraints including in your original multi-group tests.

```
boot.model.invariant <-</pre>
 bootstrap_rr(
    # saved configural model
    saved_configural = results.invariant$model_configural,
    # dataframe
    data = df.invariant,
    # model syntax
    model = model.overall,
    # group variable column in dataframe
    group = "group",
    # number of bootstraps
    nboot = 1000,
    # which fit index you would like to use
    invariance_index = "cfi",
    # what is your criterion for that fit index
    invariance_rule = .01,
    # what equality constraints are you testing
    group.equal = c("loadings", "intercepts", "residuals")
```

The data included in this function will be sampled, with replacement, at the same size as the current dataset, and the included invariance equality constraints are estimated. Each step will be compared to the previous step using the invariance index and comparison rule entered. The output is a dataframe of the proportion of non-invariant bootstraps from the real data and the same bootstrapped dataset with the group labels randomly assigned. The effect size comparison of proportions, h, for non-invariant comparisons:

$$h_{nmi} = 2 \times (asin\sqrt{p_{data}} - asin\sqrt{p_{random}})$$

The alternative, h_{mi} , for effect size of measurement invariance replication would simply be the inverse sign of h_{nmi} and is also included in the table. Two additional columns h_{nmi_p} and h_{nmi_p} represent the h values divided by the upper bound of h (i.e., π), to help with interpretation of the effect size (thus, bounding h to -1 to 1).

438 Simulated Results

Figure 11 portrays the h_{nmi_p} values by simulated non-invariance, strength of non-invariance, and type of equality constraint. This image represents 100 simulations of data by 1000 bootstrapped runs (averaged) to explore the expected pattern of results. The bars are arranged to show what a researcher might inspect when thinking about replication possibilities and their effect sizes (i.e., only three bars for each equality constraint would be calculated).

In the data that was simulated to be invariant between groups, effect sizes are still non-zero (loadings $h_{nmi_p} = 0.28$, intercepts = $h_{nmi_p} = 0.06$, $h_{nmi_p} = 0.00$). This result mirrors the effects found in the literature - that often, many models fail to show invariance, and potentially not because measurement is poor but because of natural random variation in parameter estimates. This result also indicates the need to be able to identify if specific parameters are driving the differences, which is shown in the next section.

Next, Figure 11 demonstrates the patterns one might find for small, medium, and large effects at each type of invariance when data is simulated with *one* difference. For loadings, the pattern shows a larger effect for loadings with zero or negative effect sizes for other effect sizes. The intercept simulations show non-zero effect sizes in the loadings and intercepts, likely for the same reasons d_{MACS} is interpreted as a combined effect size. When intercepts are changed, loadings may naturally shift with those means. Last, the residual results present an unexpected pattern, wherein the effect is primarily seen in the loadings,

rather than the residuals step. However, when distributions of error variance are different,
one may expect that those effects are pushed toward the loadings as well (as values can vary
more, thus potentially weakening the relationship between observed and latent variable).

An example of interpretation on real data is given in a later section. From a research 461 study, only one effect size for each equality constraint would be calculated. The 462 interpretation will often be up to the researcher's smallest effect of interest, and this 463 simulation gives some guidance that the values should not be interpreted with traditional 464 rules of thumb. The pattern of effects is potentially the most useful information: 1) positive 465 effects on the loadings with negative or very close to zero effects on the other parameters may indicate a non-replication in loadings, 2) equal effects on loadings and intercepts with smaller or negative effects may indicate intercepts may be an issue, and 3) residuals may be determined by the same pattern as loadings but with a smaller ratio of loadings to residuals effect (i.e., loadings h_{nmi} / residuals h_{nmi} . The "size" could be determined by the ratio of 470 effect sizes for each constraint. Of course, this represents one simulation study, and results 471 from many studies in a meta-analysis would be fruitful for future work. 472

⁴⁷³ Replication and Effect Size: Parameters

474 Package Function

After examining the overall model potential replication effect size, the individual 475 parameters within a model can be bootstrapped for partial invariance to with that 476 parameter relaxed (overall partial model statistics) and the difference in group parameter 477 estimates (parameter effect size). This function uses arguments seen in other functions, so they will not be repeated here. The general setup consists of using the model you think 479 could be partially invariant in the saved model argument and the fit index for comparison for the model with less constraints in invariance compare. This example examines the 481 loadings in the invariant model, so saved model uses the mgcfa output for equality 482 constraints present on the loadings and compares that model to the configural model with no 483

equality constraints on the loadings. The partial_step argument will be used to determine
which operation syntax (i.e. =~ for loadings) to relax for modeling.

```
boot.partial.invariant <-</pre>
      bootstrap_partial(
        # saved model you want to examine the partial loadings for
        saved_model = results.invariant$invariance_models$model.loadings,
        # the dataset
        data = df.invariant,
        # the model
        model = model.overall,
        # the group variable in the dataset
        group = "group",
        # number of bootstraps
        nboot = 1000,
        # which fit index you would like to use to determine partial invariance
        invariance_index = "cfi",
        # what is the invariance rule
        invariance_rule = .01,
        # what are we comparing the saved model fit index to
        invariance_compare = fitmeasures(results.invariant$model_configural, "cfi"),
        # what step are we using for invariance
        partial_step = "loadings",
        # what equality constraints should be imposed
        group.equal = c("loadings")
```

```
names(boot.partial.invariant)
```

```
## [1] "invariance_plot" "effect_invariance_plot" "density_plot"
```

[4] "boot_DF" "boot_summary" "boot_effects"

The saved output includes several dataframes and plots. The first is the boot_DF
which the summary of each run in a dataframe for plotting or summarization. This
dataframe includes the estimate for each paramter (term) separated by group and type
(boot_1, boot_2 are the bootstrapped estimates for group 1 and group 2, while the same
random columns indicate the randomly assigned groups). The fit index used to determine
invariance is included for bootstrapped and random estimates, and then the differences
between groups and if they were "invariant" or not given the researcher supplied rule.

head(boot.partial.invariant\$boot_DF)

495	##			te	erm	boot_1	boot_2	random_1	random_2	boot_fit	random_fit
496	##	1	lv	=~	q1	0.4548783	0.49928877	0.4627486	0.4651391	0.9296990	1.0000000
497	##	2	lv	=~	q2	0.3599017	0.56241016	0.4100874	0.4980215	0.9441125	1.0000000
498	##	3	lv	=~	q3	0.4254283	0.33640233	0.4274329	0.3422124	0.9377130	1.0000000
499	##	4	lv	=~	q4	0.3930716	0.03320619	0.1380833	0.2628802	0.9750274	1.0000000
500	##	5	lv	=~	q5	0.7306414	0.73512673	0.7093891	0.7532471	0.9266587	1.0000000
501	##	6	lv	=~	q1	0.5537083	0.57086815	0.5732166	0.5475714	0.8958929	0.9814658
502	##		boo	ot_d	difi	ference ran	ndom_differe	ence boot_	index_diff	erence	
503	##	1		-0	. 044	1410454	-0.002390	0463		FALSE	
504	##	2		-0	. 202	2508484	-0.087934	1027		FALSE	
505	##	3		0	. 089	9025927	0.085220)565		FALSE	
506	##	4		0	. 359	9865463	-0.124796	6846		FALSE	
507	##	5		-0	. 004	1485377	-0.04385	7947		FALSE	
508	##	6		-0	.017	7159815	0.02564	5271		FALSE	
509	##		raı	ndor	n_ir	ndex_differ	rence				
510	##	1					TRUE				
511	##	2					TRUE				

```
512 ## 3 TRUE

513 ## 4 TRUE

514 ## 5 TRUE

515 ## 6 TRUE
```

Next, the boot_summary includes a summarized form of the bootstrapped results from separated by bootstrapping versus random and invariant/non-invariant. The d_s for between groups Cohen's d is shown below, and the non-central confidence interval is included. Effect sizes are only calculated when the number of bootstrapped estimates is at least 10% of the data - therefore, you would not receive effect sizes with almost no bootstrapped runs. This dataframe should be used to determine which parameter may be different and at what size between groups in a replication of the study.

```
boot.partial.invariant$boot_summary %>%

dplyr::select(term, d_boot, d_random)
```

```
## # A tibble: 10 x 4
523
      # Groups:
                     term, invariant [10]
524
   ##
          invariant term
                                   d boot d random
525
   ##
          <lgl>
                      <chr>
                                    <dbl>
                                               <dbl>
526
                      lv = ~q1 - 0.0299
   ##
        1 FALSE
                                            0.0583
527
                      lv = ~q1 0.0337
        2 TRUE
                                            0.0116
   ##
528
                      lv = ~q2 - 0.0326
   ##
        3 FALSE
                                            0.0933
529
                      lv = q2 0.146
   ##
        4 TRUE
                                            0.0309
530
                      lv = ~q3 -0.0463
   ##
        5 FALSE
                                            0.113
531
                      lv = ~q3 -0.148
   ##
        6 TRUE
                                            0.0743
532
                      1v = q4 \quad 0.00785 \quad -0.0668
        7 FALSE
   ##
533
                      lv = ~q4 - 0.0157
        8 TRUE
                                            0.0389
   ##
534
        9 FALSE
                      lv = ~q5 -0.00129 -0.169
   ##
535
```

```
lv = q5 0.122
                                 -0.00853
## 10 TRUE
```

The boot_effects table creates a summary similar to the overall model replication 537 table based on the proportion of runs that were considered invariant versus not for each 538 parameter. Note that the effects match the overall results, such that simulated invariant 539 data appears to still show the likelihood that loadings may not replicate in a similar dataset. 540

boot.partial.invariant\$boot effects

A tibble: 5 x 7

546

542	##	term	non_invariant	random_non_invariant	h_nmi	h_mi	h_nmi_p	h_mi_p
543	##	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
544	## 1	lv =~ q1	0.853	0.236	1.34	-1.34	0.427	-0.427

2 lv = q20.858 0.237 1.35 - 1.350.430 - 0.430## 3 lv =~ q3 0.851 0.23 1.35 - 1.350.429 - 0.429

4 lv = ~ q40.84 0.229 1.32 - 1.320.420 - 0.420547

5 lv = q50.819 0.237 1.25 - 1.250.397 - 0.397548

Plots of the results from dataframes can be found within the bootstrap_partial() 549 function. Figure 12 shows the difference between parameters for groups in the bootstrapped 550 and randomly assigned group runs. Figure 13 shows the density plot of the estimates for 551 each group organized by bootstrapped and randomly assigned groups and the invariance 552 decision for each bootstrapped run. Last, Figure 14 indicates the d_s value between groups with an indication of the number of data points in each estimate (i.e., dot size). These visualizations should allow a researcher to understand the likelihood of replication for each 555 parameter, as well as the potential size of the differences. Therefore, one could indicate a 556 specific smallest effect size of interest, rather than a invariance cut-off rule of thumb when 557 planning a replication or registered report. 558

559 Simulated Results

Figure 15 shows the effect size differences within large loadings simulations. The 560 results demonstrate that most of the loadings were considered non-invariant in the 561 bootstrapped models (while holdings all others equal). This result is partially due to 562 simulating very good data, so small changes in loadings results in a drop in fit for our chosen 563 invariance index. However, we can use this graph to show that question four shows a 564 possible effect size ranging from -0.07 to 0.13. The h_{nmi_p} value for question four was 0.27, 565 representing about a quarter of a possible total effect. Last, the density plot in Figure 16 566 shows the separation of the two different groups loadings in item four, thus, illustrating 567 group differences in the findings for their loadings. Each of the other combination of plots 568 can be found in the supplemental materials.

570 An Example Analysis

Aiena et al. (2014) examined the RS-14 (Wagnild, 2009) exploring the factor 571 structure of the Resiliency Scale in a clinical sample receiving treatment services and a 572 college student sample. Measurement invariance was calculated for differences separately for 573 these samples for gender and race finding a partially invariant models with a few item 574 intercepts or residuals that differed between groups. Aiena et al. (2014) did not compare the 575 clinical to the student sample for measurement invariance, and it is reasonable to expect 576 potential differences in these two populations. This example will demonstrate the procedure 577 for researchers who wish to use partial invariance steps and how to interpret real, messy data. 578

```
# load the data, it is called DF
load("manu_data/RS14.Rdata")

# build the one-factor model
model.rs <- "RS =~ RS1+RS2+RS3+RS4+RS5+RS6+RS7+RS8+RS9+RS10+RS11+RS12+RS13+RS14"
# run the multi-group CFA
results.rs <- mgcfa(</pre>
```

```
model = model.rs,
data = DF,
group = "sample",
group.equal = c("loadings", "intercepts", "residuals")
)
```

```
# how to get results in table
results.rs$model_fit %>%

dplyr::select(model, AIC, BIC, cfi, tli, rmsea, srmr)
```

Table 1 indicates the results after running the one-factor model. There are several 579 guidelines for assessing assessing a degradation in model fit (Cao & Liang, 2022; Cheung & Rensvold, 2002; Counsell et al., 2020; Jin, 2020; Putnick & Bornstein, 2016) but for the purposes of this illustration $\Delta CFI > .01$ will be used. Table 1 indicates that fit was degraded 582 when the constraint on equal item intercepts was added. The code below provides an 583 example of testing each item individually by relaxing the constraints and recalculating the 584 CFI. If these Items bring the CFI value back up to Δ CFI <= .01 from the metric model, 585 then the model would be considering partially invariant at the scalar level. It seems unlikely 586 that the residuals will show invariance, if partial scalar invariance can be found, as the drop 587 in fit is quite large. 588

```
partial.rs <-
partial_mi(
    saved_model = results.rs\invariance_models\intercepts,
    data = DF,
    model = model.rs,
    group = "sample",
    # be sure to do only up to the step you are interested in
    group.equal = c("loadings", "intercepts"),
    partial_step = "intercepts")</pre>
```

```
partial.rs$fit_table %>%

dplyr::select(free.parameter, cfi)
```

```
## # A tibble: 15 x 2
589
   ##
          free.parameter cfi
590
          <chr>
                            <lu>n.vctr>
   ##
591
        1 "RS1 ~1 "
                            0.9116914
   ##
592
        2 "RS2 ~1 "
                           0.9129976
   ##
593
        3 "RS3 ~1 "
                           0.9117235
   ##
594
        4 "RS4 ~1 "
                           0.9111212
   ##
595
        5 "RS5 ~1 "
                            0.9126742
   ##
596
   ##
        6 "RS6 ~1 "
                           0.9133618
597
          "RS7 ~1 "
                           0.9139287
   ##
598
        8 "RS8 ~1 "
                           0.9111397
   ##
599
   ##
        9 "RS9 ~1 "
                            0.9119702
600
   ## 10 "RS10 ~1 "
                           0.9118309
601
                           0.9110574
   ## 11 "RS11 ~1 "
602
   ## 12 "RS12 ~1 "
                           0.9112309
603
   ## 13 "RS13 ~1 "
                            0.9112367
604
   ## 14 "RS14 ~1 "
                           0.9112015
605
   ## 15 "RS ~1 "
                            0.9108805
606
```

The output indicates that RS6 and RS7 are potential items that could be relaxed to improve model fit and create a partial scalar invariant model (i.e., by picking the largest CFI values). The code below show to check the addition of these items, which are added one at a time. You use the group.partial open to "relax" or freely estimate that parameter for each group separately.

```
# run the partially invariant model with group.partial
partial.rs.1 <- mgcfa(model = model.rs,</pre>
                  data = DF,
                  group = "sample",
                  group.equal = c("loadings", "intercepts"),
                  group.partial = c("RS7~1"),
                 meanstructure = TRUE)
# examine the loadings
partial.rs.1$model_coef %>%
  filter(term == "RS7 ~1 ") %>%
 filter(model == "intercepts") %>%
  dplyr::select(term, group, estimate, std.error)
## # A tibble: 2 x 4
##
                group estimate std.error
     term
     <chr>
                         <dbl>
##
                <int>
                                    <dbl>
## 1 "RS7 ~1 "
                          4.95
                    1
                                   0.0580
## 2 "RS7 ~1 "
                    2
                          4.49
                                   0.0529
# examine the fit indices
partial.rs.1$model_fit %>%
  filter(model == "intercepts") %>%
 dplyr::select(AIC, BIC, cfi, tli, rmsea, srmr)
## # A tibble: 1 x 6
##
         AIC
                  BIC
                        cfi
                               tli rmsea
                                            srmr
                <dbl> <dbl> <dbl> <dbl> <dbl>
       <dbl>
## 1 122454. 122804. 0.914 0.912 0.102 0.0502
```

613

614

615

616

618

619

```
# effect size model
lavaan_dmacs(partial.rs.1$invariance_models$model.intercepts, "Clinical")$DMACS[7]
```

```
621 ## RS7
622 ## 0.282302
```

By examining our estimates, we can see that item seven on the RS-14 is estimated at 623 nearly 5 points for the clinical sample, while the student sample has a lower mean around 4.5 624 points. Generally, students show higher means on the items of the RS-14, but when all 625 loadings and other intercepts are constrained to be equal, and this one item is relaxed, this 626 pattern flips so that clinical groups show higher item intercepts. Given the scale is a 1-7 627 Likert type scale, .5 a point represents a potentially sizable change on the scale. Item seven 628 covers perseverance after hardship, and all items can be found in the user manual for the 629 scale at www.resiliencecenter.com. The effect size from d_{MACS} suggests a small to medium 630 effect, 0.28. In this next code section, we repeat this process for the RS6, as the CFI for our 631 model with only RS7 does not achieve the levels of partial invariance for our Δ CFI criterion 632 (i.e., <= .01 downward change in fit: metric CFI = .925, partial scalar CFI = .914). See Figure 17 for the difference between item intercepts and latent means.

```
filter(term == "RS6 ~1 ") %>%
     filter(model == "intercepts") %>%
     dplyr::select(term, group, estimate, std.error)
   ## # A tibble: 2 x 4
635
   ##
                    group estimate std.error
         term
636
         <chr>>
                    <int>
                              <dbl>
                                         <dbl>
637
   ## 1 "RS6 ~1 "
                               5.00
                                        0.0605
638
   ## 2 "RS6 ~1 "
                        2
                               4.54
                                        0.0533
639
   # examine the fit indices
   partial.rs.2$model_fit %>%
     filter(model == "intercepts") %>%
     dplyr::select(AIC, BIC, cfi, tli, rmsea, srmr)
   ## # A tibble: 1 x 6
                      BIC
   ##
             AIC
                             cfi
                                    tli rmsea
                                                 srmr
641
           <dbl>
                    <dbl> <dbl> <dbl> <dbl> <
642
   ## 1 122363. 122719. 0.917 0.915 0.100 0.0488
   # effect size model
   lavaan_dmacs(partial.rs.2$invariance_models$model.intercepts, "Clinical")$DMACS[6]
             RS6
   ##
   ## 0.2796334
          Again, we see about a half-point difference between our clinical and student samples
```

Again, we see about a nair-point difference between our clinical and student samples for item 6, which is about drive to achieve. The CFI for this model does meet the requirements for partial invariance, .917. The effect size is approximately the same at d_{MACS} = 0.28. See Figure 18 shows the difference between item intercepts and latent means.

```
# plot the image for RS7
plot_mi(
    data_coef = partial.rs.2$model_coef,
    model_step = "intercepts",
    item_name = "RS7",
    x_limits = c(-1,1),
    y_limits = c(min(DF$RS7), max(DF$RS7)),
    conf.level = .95,
    model_results = partial.rs.2$invariance_models$model.intercepts,
    # which latent variable do you want
    lv_name = "RS"
)$complete
```

```
# plot the image for RS6
plot_mi(
  data_coef = partial.rs.2$model_coef,
  model_step = "intercepts",
  item_name = "RS6",
  x_limits = c(-1,1),
  y_limits = c(min(DF$RS6), max(DF$RS6)),
  conf.level = .95,
  model_results = partial.rs.2$invariance_models$model.intercepts,
  # which latent variable do you want
  lv_name = "RS"
)$complete
```

Next, we would examine our replication potential for this model. Given our current results, we may not expect our intercepts to replicate. Given the order of group.equal, the

boot function will select the first non-invariant step in the calculation of the effect size for
potential replication. In our output, we do not see a loadings effect size, and this result
occurs when *none* of the bootstrapped or random results are non-invariant. Therefore, we
would expect the loadings to replicate (and the effect size would be 0 difference between
bootstrapped and random, both showing invariance). The intercepts show a large (i.e., close
to the max possible value) non-invariant effect, and therefore, we should not expect this
model to show invariance in a replication.

```
boot.model.rs <-
bootstrap_rr(
    saved_configural = results.rs$model_configural,
    data = DF,
    model = model.rs,
    group = "sample",
    nboot = 1000,
    invariance_index = "cfi",
    invariance_rule = .01,
    group.equal = c("loadings", "intercepts", "residuals")
)</pre>
```

boot.model.rs

```
## # A tibble: 2 x 7
659
                     non invariant random non invariant h nmi
   ##
         model
                                                                      h_mi h_nmi_p h_mi_p
660
         <chr>
                                                            <dbl>
   ##
                             <dbl>
                                                     <dbl>
                                                                     <dbl>
                                                                              <dbl>
                                                                                       <dbl>
661
   ## 1 intercepts
                             0.998
                                                         0 3.05
                                                                   -3.05
                                                                             0.972
                                                                                    -0.972
662
                                                         0 0.0895 -0.0895 0.0285 -0.0285
   ## 2 residuals
                             0.002
663
```

Next, we would examine the strength of the effects of replication on each parameter

664

at the intercept level. By examining Table 2, it is clear that most of the item means are unlikely to replicate, even though two particular items can be used to create partial invariance. Figures 19 and 20 display the three plots provided in the bootstrap_partial() function. In general, we should expect $M_D = 0.23$ when items are invariant and $M_D = 0.26$ when items are not invariant. The effect size of non-invariant items ranges from 0.43 to 0.62.

The density plot shown at the bottom of Figure 19 illustrates the likely reasons for 670 the differences found in the top plots. It appears that many items show a bimodal 671 distribution within group 1 (Clinical Sample) and when items are invariant, the intercept averages to the same intercept as group 2 (Student Sample). In non-invariant estimates, the 673 same bimodal distributions are found, but they are more extreme than the student samples, 674 and therefore, item show different averages due to the presence of two separate means of 675 data. Further, some items also appear to show two separate student item averages within the 676 data. This result suggests that it would be fruitful to understand a potential predictor of 677 these differences or other confounding variable that separates these samples, creating 678 differences in item averages. 679

In summary, if one were planning a replication, the prediction would be that item 680 intercepts would likely not replicate, with a large effect size (i.e., it is easy to judge h_{nmi_n} 681 close to the max of one as large). While this study found partial invariance by relaxing 682 constraints on two individual items, bootstrapped partial invariance indicates that any item 683 could potentially be problematic with an effect size averaging $d \sim 0.50$ difference in means. 684 While d_{MACS} values represented a "small" effect based on previous publications, this effect may be muted by examining both loadings and intercepts. The results here suggest that the effect is driven by intercepts. The overall average score on items is high: $M_M = 5.04 \ (M_{SD})$ 687 = 1.72). Given the mean standard deviation, a $d \sim 0.50$ represents 0.86 or nearly one whole 688 point on the scale. A researcher could decide that at least d = 0.33 or at least a third of a 689 standard deviation would be an important change and set that as their smallest effect size of 690

interest for invariance. Further, a newly planned study should investigate what variables may predict when and why samples separate into bimodal representations for item means.

```
bootstrap_partial(
    saved_model = results.rs$invariance_models$model.intercepts,
    data = DF,
    model = model.rs,
    group = "sample",
    nboot = 1000,
    invariance_index = "cfi",
    invariance_rule = .01,
    invariance_compare = fitmeasures(results.rs$invariance_models$model.loadings, "cfi")
    partial_step = "intercepts",
    group.equal = c("loadings", "intercepts")
)
```

Discussion

In this tutorial, we examined how to use multiple tools to examine measurement 694 invariance and its potential replication. Model fit comparisons and statistics can be paired 695 with the proposed effect size measures, and a visualization to examine individual items and 696 the overall latent mean scores. The impact of potential replication was estimated on the 697 overall model and the individual parameters. Using real data, the effect of two non-invariant 698 item intercepts was examined and visualized. This tutorial manuscript has provided a concrete way to plan for pre-registration and/or registered reports. Researchers could simulate results based on published or previously collected data to determine the likelihood 701 and size of potential replication. They could plan and pre-register a smallest effect of 702 interest. For example, we may determine that an h_{nmi_p} value above .20 represents an 703 important level of non-invariance for our model overall, while $h_{nmi_p} > .30$ for any individual 704

parameter warrant caution against invariance for groups. Others have begun to discuss the importance of focusing on effects in the scale of the data and their practical importance (Anvari & Lakens, 2021; Cumming, 2012).

From the example, our interpretation may be that the difference between group's latent means is large, as a 0.72 change on a 7 point scale is approximately 10% more resiliency for students when compared to the clinical sample. Practically, 10% in resiliency for an area of the United States (Mississippi) often hit with natural disasters (hurricanes, tornadoes, floods) and high levels of poverty would be very important. Even the smaller difference of .5 point on each individual item could translate into increases in resiliency, and these results may elucidate avenues for further exploration into areas of focus within resiliency, given the items.

What do the results of a study on measurement invariance with these results tell us 716 about replication, generalizability, and validity overall? If a researcher decides their effects 717 are large, they should likely caution against suggesting that these scores are directly 718 comparable without weighting or other adjustment. Let's consider a scenario wherein the 719 change metric between models picked (i.e., Δ CFI, Δ RMSEA) indicates a "significant" 720 change in model fit. However, if both the effect size and a visual inspection of the invariance 721 indicates a small difference, we may decide to lessen the practical importance of those results, 722 much like "just significant" p-values with small effect sizes are treated now. Given that the 723 goal of measurement invariance is to compare estimates, we should expect some differences 724 across samples due to the nature of sampling and estimation. It may be that many of the 725 published models presented represent these effects - small variations between groups due to 726 sampling error or other small crud - but do not represent a fundamental problem with the 727 measurement or generalizability of the results. 728

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Table 1

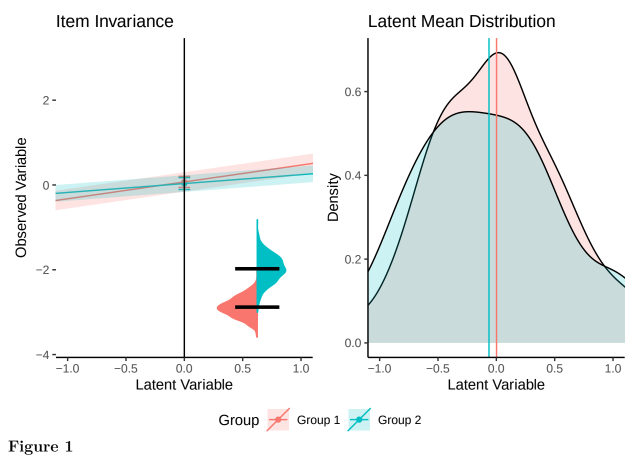
Model Fit for RS-14 Example

Model	AIC	BIC	CFI	TLI	RMSEA	SRMR
Overall	126,722.491	126,888.707	0.934	0.923	0.094	0.036
Group Clinical	52,961.421	53,099.720	0.919	0.904	0.090	0.044
Group Student	69,100.985	69,254.310	0.928	0.915	0.108	0.035
Configural	122,118.406	122,617.055	0.926	0.912	0.102	0.036
Loadings	122,144.532	122,566.010	0.925	0.918	0.098	0.043
Intercepts	122,544.109	122,888.415	0.911	0.910	0.103	0.052
Residuals	126,466.241	126,727.438	0.780	0.793	0.156	0.086

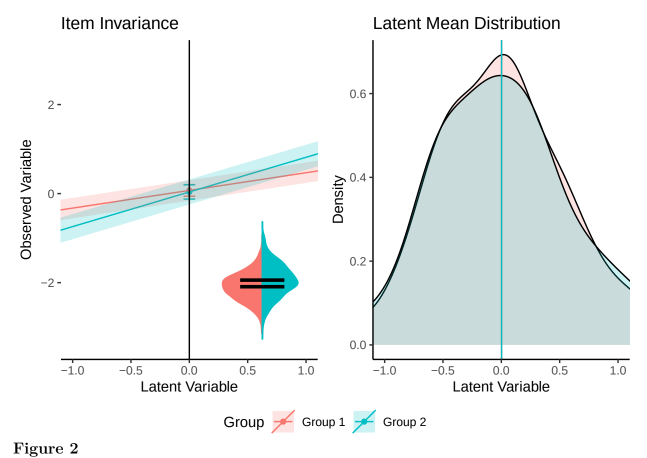
Table 2

Boot Partial Effects Results for RS-14 Intercepts

Term	Non-Invariant	Random Non-Invariant	h_{nmi}	h_{nmi_p}
RS Intercept	0.991	0.007	2.784	0.886
RS1 Intercept	0.989	0.007	2.764	0.880
RS10 Intercept	0.988	0.007	2.755	0.877
RS11 Intercept	0.991	0.007	2.784	0.886
RS12 Intercept	0.991	0.007	2.784	0.886
RS13 Intercept	0.991	0.007	2.784	0.886
RS14 Intercept	0.990	0.007	2.774	0.883
RS2 Intercept	0.985	0.007	2.728	0.869
RS3 Intercept	0.988	0.007	2.755	0.877
RS4 Intercept	0.990	0.007	2.774	0.883
RS5 Intercept	0.984	0.007	2.720	0.866
RS6 Intercept	0.979	0.007	2.683	0.854
RS7 Intercept	0.974	0.007	2.650	0.844
RS8 Intercept	0.991	0.007	2.784	0.886
RS9 Intercept	0.987	0.007	2.746	0.874



Invariant Model Visualization



Small Loadings Model Visualization

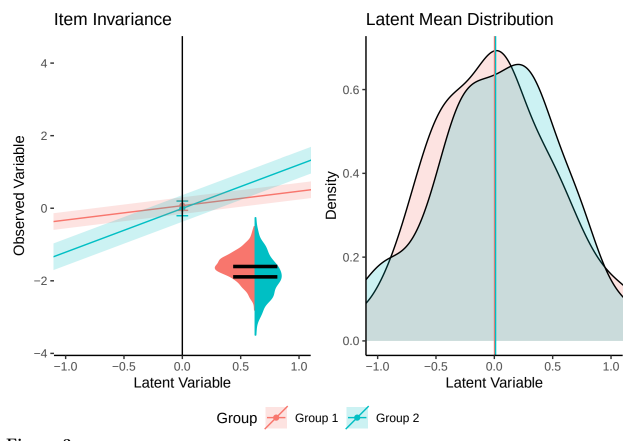
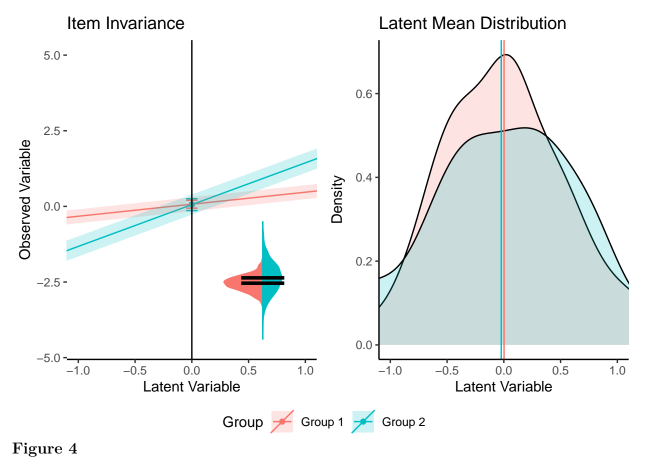
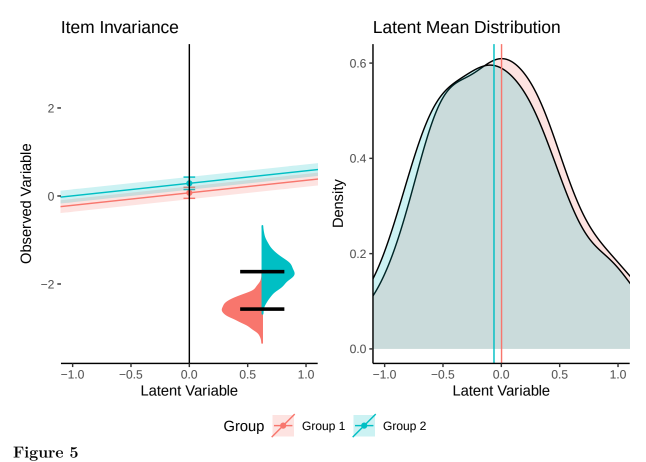


Figure 3

Medium Loadings Model Visualization



Large Loadings Model Visualization



Small Intercepts Model Visualization

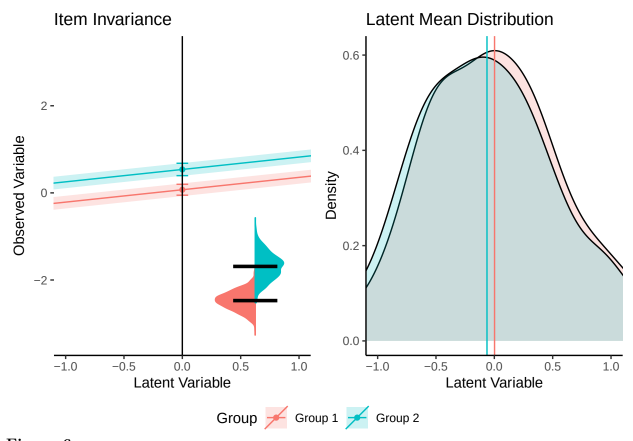
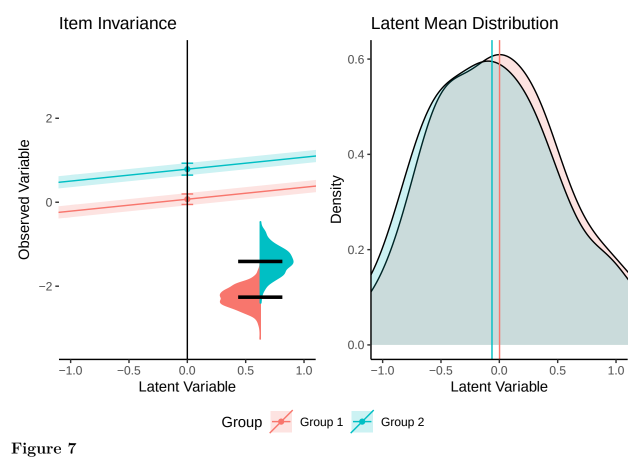
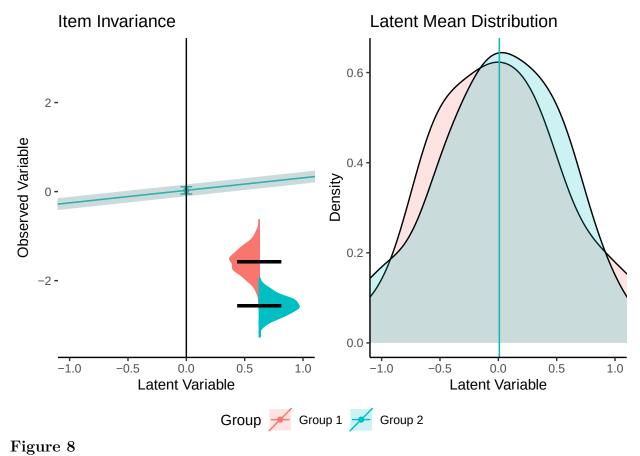


Figure 6

Medium Intercepts Model Visualization



Large Intercepts Model Visualization



Small Residuals Model Visualization

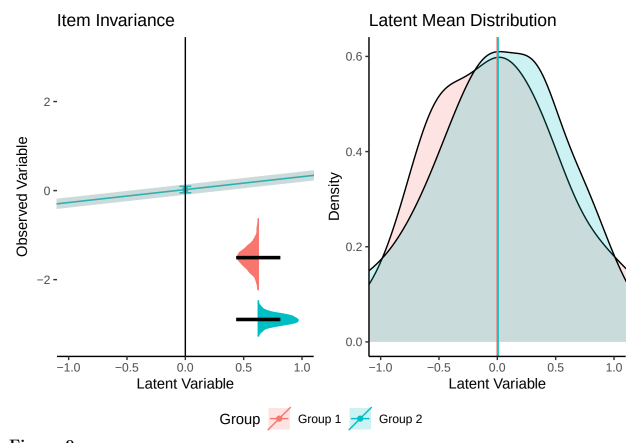
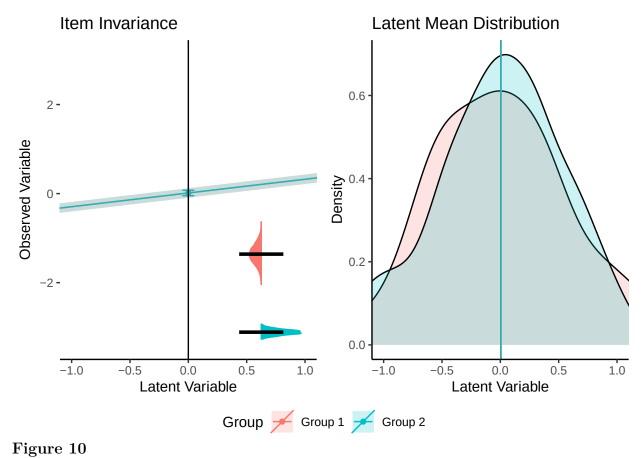


Figure 9

Medium Residuals Model Visualization



Large Residuals Model Visualization

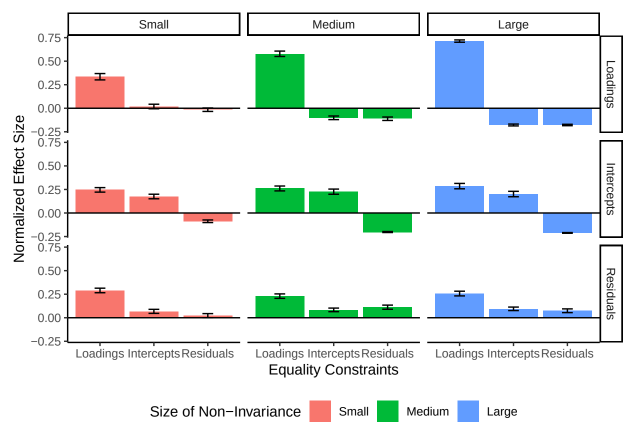


Figure 11

Visualization of the effect size of bootstrapped replication proportions on simulated data. Each panel indicates the simulated data type, colors represent the differences in the strength of the non-invariance, and the bars on the x-axis represent the effect size for the equality constraint.

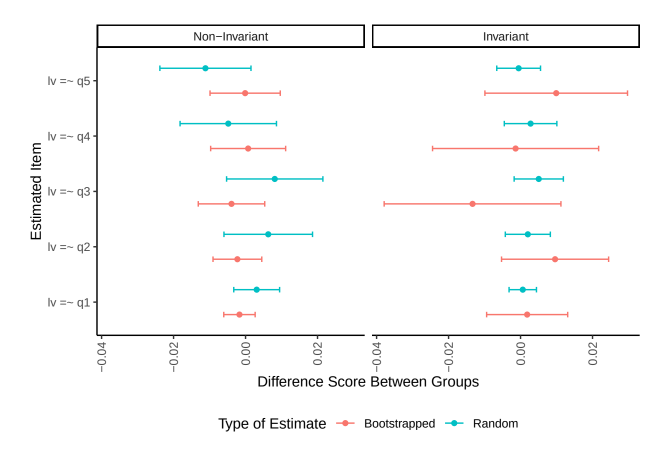


Figure 12
Visualization of the difference score between groups by parameter for invariant and non-invariant bootstrapped and randomly assigned group data.

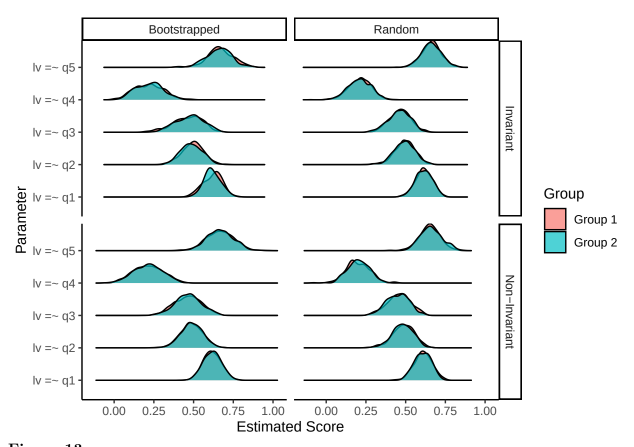


Figure 13

Visualization of the number of estimates for each group by bootstrapped and randomly assigned group runs by their invariance decision.

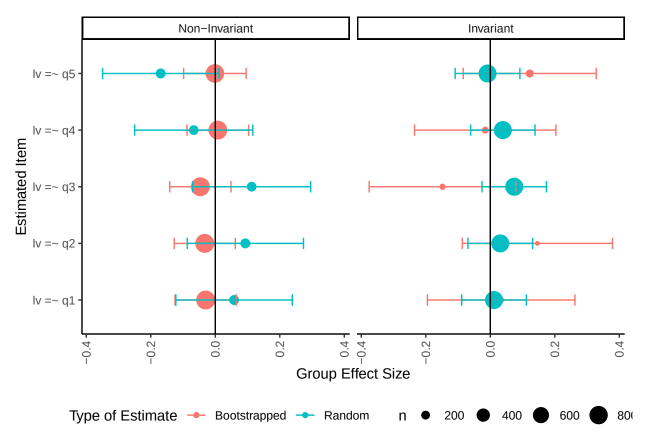
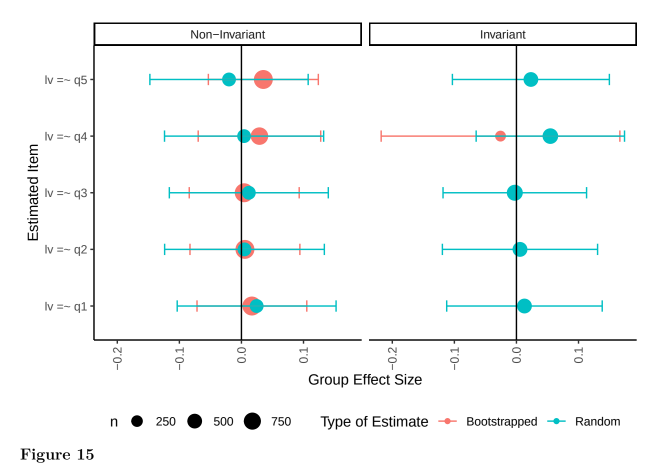


Figure 14

Visualization of effect size between groups by parameter for invariant and non-invariant bootstrapped and randomly assigned group data. The size of the dots indicate the number of data points for that estimate.



Bootstrapped and Random Group effect size differences in loadings for the Large Loading difference simulation. The size of the point represents the number of data points included in that calculation.

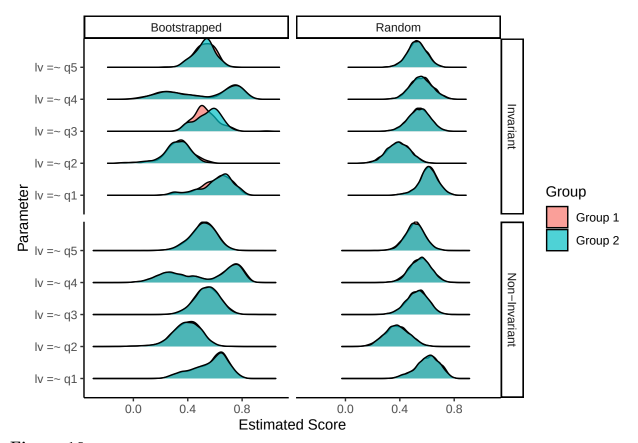


Figure 16

Bootstrapped and Random density plots for invariant and non-invariant bootstrapped partial effects examining only large loadings.

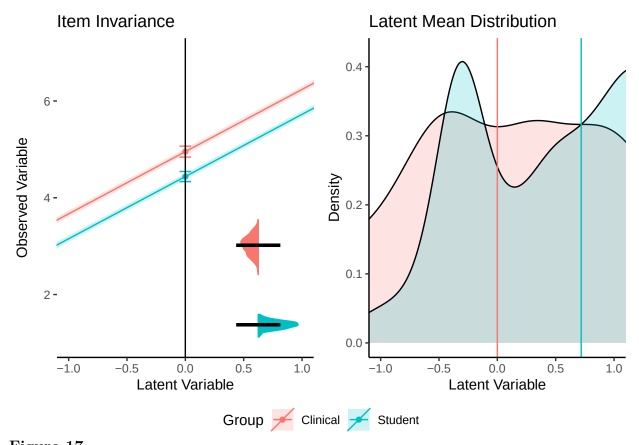


Figure 17

RS7 Item Invariance Visualization

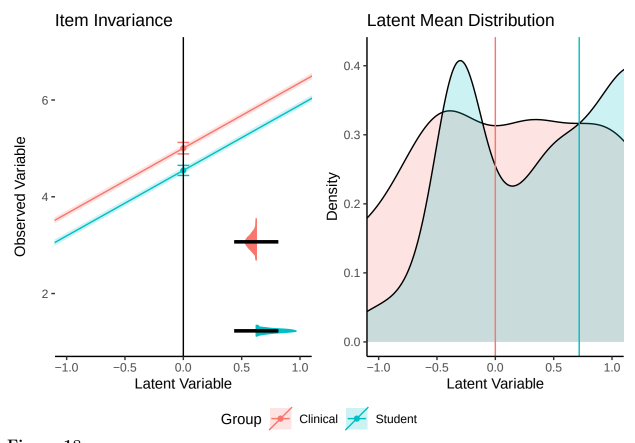


Figure 18

RS6 Item Invariance Visualization

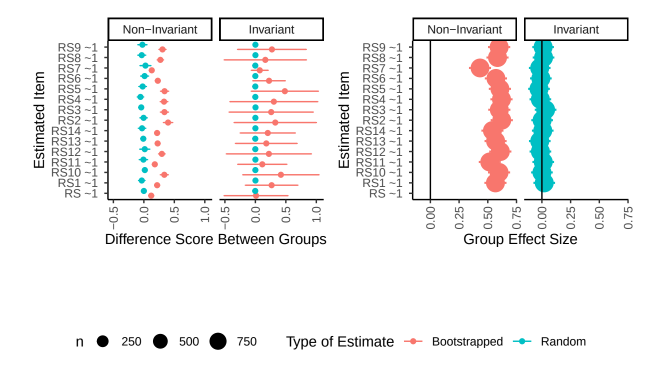
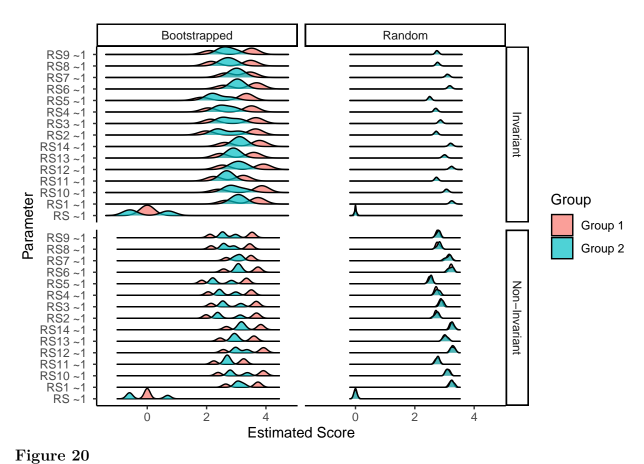


Figure 19

RS-14 scale invariance for item intercepts. The left panel indicates the raw score difference between groups and items, while the right panel indicates the effect size for group differences based on invariance.



RS-14 scale invariance density plots, illustrating invariant versus non-invariant bootstrapped and random runs for each parameter.

Appendix A

Simulating from models

Here's an example of how to simulate directly from a lavaan model:

```
# first build your model
# this example is separate for each group
model.invariant.g1 <- "</pre>
# loadings
1v = .8*q1 + .4*q2 + .6*q3 + .3*q4 + .6*q5
# set the residual for invariance on q4
q4 ~~ 1*q4
# set the intercept for invariance on q4
q4 \sim 0*1
# set the intercept to zero for df purposes
q1 ~ 0*1
# allow the latent mean to be estimated
lv ~ 1"
model.invariant.g2 <- "lv =~ .77*q1 + .43*q2 + .58*q3 + .3*q4 + .61*q5
q4 ~~ 1*q4
q4 \sim 0*1
q1 ~ 0*1
lv ~ 1"
# simulate data invariant separately for each group
df.invariant <- bind_rows(</pre>
  # lavaan function
  simulateData(
    # model with estimates
   model = model.invariant.g1,
```

```
# how many data points
   sample.nobs = 250,
    # mean structure for mgcfa models
   meanstructure = T,
   # model type
   model.type = "cfa",
   # set seed for reproducibility
   seed = 1234) %>%
   # add a group label to the data
   mutate(group = "Group 1"),
  simulateData(
   model = model.invariant.g2,
   sample.nobs = 250,
   meanstructure = T,
   model.type = "cfa",
   seed = 1234) %>%
   mutate(group = "Group 2")
)
```

Appendix B

Simulating from matrices

Here's an example of how to simulate using MASS and covariance or correlation matrices.

```
library(MASS)
# covariance matrix
university.cov <- lav_matrix_lower2full(</pre>
    c(169.00,
      73.710, 182.2500,
      73.229, 88.4250, 171.6100,
      63.375, 72.5625, 127.7250, 156.2500,
      42.120, 67.4325, 122.0265, 123.1875, 182.2500,
      57.226, 63.2610, 117.1926, 154.4250, 138.0240, 201.6400,
      30.875, 32.0625, 60.9805, 62.9375, 76.9500, 79.5910, 90.2500,
      36.075, 38.9610, 61.0722, 58.2750, 65.9340, 70.9290, 81.1965, 123.2100,
      18.096, 21.1410, 26.2131, 39.1500, 44.6310, 46.9452, 48.7635, 56.0106, 75.6900))
# give it names
rownames(university.cov) <-</pre>
    colnames(university.cov) <-</pre>
    c("class", "social", "learn", "chronic", "physical", "sex",
      "depression", "anxiety", "stress")
# means - you need standard deviation if you only have a correlation matrix
university.means \leftarrow c(3.4, 4.3, 3.7, 3.2, 4.5, 1.2, 4.0, 3.5, 4.2)
# use mass function
DF <- mvrnorm(n = 200, mu = university.means, Sigma = university.cov)
```

head(DF)

```
##
               class
                          social
                                     learn
                                             chronic
                                                        physical
                                                                         sex
874
   ## [1,] 12.085294 26.83043663 13.895103 20.634374 16.7903295
                                                                 30.5468832
875
   ## [2,] 3.939545 10.93624431 3.093025 8.316128 -7.2199148
                                                                  9.3579991
876
   ## [3,] 15.659627 0.22811723 5.205657 5.224293 -1.4425219 -1.2710662
877
   ## [4,] 23.086133 12.43649966 1.891769 -5.913170 -19.4937389 -13.5922410
878
   ## [5,] 10.856492 23.30887194 17.124064 11.438840 2.4659294
                                                                   1.7330709
   ## [6,] -4.328380 0.07907149 -1.000636 -1.654947 -0.7365838 -0.3958833
880
           depression
   ##
                         anxiety
                                      stress
881
   ## [1,] 14.174374
                        8.001766
                                 13.6534279
882
   ## [2,]
           3.940819 -6.598153
                                 -7.9229552
883
   ## [3,] -8.927508 -10.335571 -2.8100779
884
   ## [4,] -0.149840
                        6.475669 -15.5300195
885
   ## [5,] -6.492809 -12.524601 0.4924153
886
   ## [6,] 11.917631 3.433587
                                 8.1456346
887
```

Appendix C MGCFA Model Fit Statistics

Model fit statistics are provided for each of the ten model combinations (invariant, three

- sizes for each loadings, intercepts, and residuals). These tables could be used to examine the
- traditional change in fit statistics cutoff rules of thumb (Cheung & Rensvold, 2002), such as
- Δ CFI or Δ RMSEA, to the visualizations presented in the manuscript.

[tbp]

Table C1

Model Fit for Invariant Model

Model	AIC	BIC	CFI	TLI	RMSEA	SRMR
Overall	7,516.454	7,579.673	1.000	1.036	0.000	0.006
Group Group 1	3,765.749	3,818.571	0.976	0.953	0.047	0.031
Group Group 2	3,767.599	3,820.421	1.000	1.008	0.000	0.021
Configural	7,533.348	7,659.786	0.991	0.982	0.030	0.026
Loadings	7,528.476	7,638.056	0.994	0.992	0.020	0.033
Intercepts	7,522.397	7,615.118	1.000	1.003	0.000	0.035
Residuals	7,520.435	7,592.083	0.991	0.992	0.020	0.046

[tbp]

Table C2

Model Fit for Small Differences in Loadings

Model	AIC	BIC	CFI	TLI	RMSEA	SRMR
Overall	7,530.321	7,593.540	0.977	0.955	0.049	0.025
Group Group 1	3,765.749	3,818.571	0.976	0.953	0.047	0.031
Group Group 2	3,785.242	3,838.064	0.979	0.958	0.050	0.029
Configural	7,550.991	7,677.430	0.978	0.956	0.048	0.030
Loadings	7,550.133	7,659.713	0.966	0.952	0.051	0.047
Intercepts	7,542.675	7,635.397	0.979	0.977	0.035	0.047
Residuals	7,534.091	7,605.739	0.993	0.994	0.019	0.054

[tbp]

Table C3

Model Fit for Medium Differences in Loadings

Model	AIC	BIC	CFI	TLI	RMSEA	SRMR
Overall	7,598.681	7,661.900	0.946	0.893	0.078	0.035
Group Group 1	3,765.749	3,818.571	0.976	0.953	0.047	0.031
Group Group 2	3,820.090	3,872.912	0.969	0.938	0.067	0.034
Configural	7,585.839	7,712.277	0.972	0.944	0.058	0.033
Loadings	7,613.434	7,723.014	0.865	0.807	0.107	0.079
Intercepts	7,606.648	7,699.370	0.874	0.860	0.091	0.079
Residuals	7,600.057	7,671.705	0.880	0.895	0.079	0.091

[tbp]

Table C4

Model Fit for Large Differences in Loadings

Model	AIC	BIC	CFI	TLI	RMSEA	SRMR
Overall	7,684.982	7,748.201	0.987	0.973	0.040	0.023
Group Group 1	3,765.749	3,818.571	0.976	0.953	0.047	0.031
Group Group 2	3,857.143	3,909.965	0.992	0.984	0.037	0.029
Configural	7,622.892	7,749.330	0.986	0.972	0.042	0.030
Loadings	7,674.188	7,783.767	0.817	0.738	0.131	0.095
Intercepts	7,667.682	7,760.403	0.824	0.805	0.113	0.096
Residuals	7,683.177	7,754.825	0.762	0.793	0.116	0.138

 $[tbp] \centering % \begin{center} \b$

Table C5

Model Fit for Small Differences in Intercepts

Model	AIC	BIC	CFI	TLI	RMSEA	SRMR
Overall	7,520.471	7,583.690	1.000	1.035	0.000	0.007
Group Group 1	3,765.749	3,818.571	0.976	0.953	0.047	0.031
Group Group 2	3,767.599	3,820.421	1.000	1.008	0.000	0.021
Configural	7,533.348	7,659.786	0.991	0.982	0.030	0.026
Loadings	7,528.476	7,638.056	0.994	0.992	0.020	0.033
Intercepts	7,526.312	7,619.034	0.987	0.986	0.027	0.040
Residuals	7,524.356	7,596.005	0.975	0.978	0.033	0.050

[tbp]

Table C6

Model Fit for Medium Differences in Intercepts

Model	AIC	BIC	CFI	TLI	RMSEA	SRMR
Overall	7,538.375	7,601.594	1.000	1.033	0.000	0.008
Group Group 1	3,765.749	3,818.571	0.976	0.953	0.047	0.031
Group Group 2	3,767.599	3,820.421	1.000	1.008	0.000	0.021
Configural	7,533.348	7,659.786	0.991	0.982	0.030	0.026
Loadings	7,528.476	7,638.056	0.994	0.992	0.020	0.033
Intercepts	7,544.002	7,636.724	0.917	0.907	0.068	0.059
Residuals	7,542.064	7,613.712	0.905	0.917	0.065	0.067

 $[tbp] \centering % \begin{center} \b$

Table C7Model Fit for Large Differences in Intercepts

Model	AIC	BIC	CFI	TLI	RMSEA	SRMR
Overall	7,568.748	7,631.967	1.000	1.032	0.000	0.008
Group Group 1	3,765.749	3,818.571	0.976	0.953	0.047	0.031
Group Group 2	3,767.599	3,820.421	1.000	1.008	0.000	0.021
Configural	7,533.348	7,659.786	0.991	0.982	0.030	0.026
Loadings	7,528.476	7,638.056	0.994	0.992	0.020	0.033
Intercepts	7,574.054	7,666.776	0.797	0.775	0.106	0.084
Residuals	7,572.174	7,643.823	0.785	0.813	0.097	0.090

[tbp]

Table C8

Model Fit for Small Differences in Residuals

Model	AIC	BIC	CFI	TLI	RMSEA	SRMR
Overall	7,462.007	7,525.226	1.000	1.020	0.000	0.013
Group Group 1	3,765.749	3,818.571	0.976	0.953	0.047	0.031
Group Group 2	3,703.797	3,756.619	0.962	0.924	0.061	0.037
Configural	7,469.546	7,595.984	0.969	0.938	0.054	0.034
Loadings	7,471.637	7,581.217	0.944	0.920	0.062	0.049
Intercepts	7,465.722	7,558.443	0.952	0.946	0.051	0.051
Residuals	7,465.986	7,537.635	0.930	0.939	0.054	0.065

Table C9

Model Fit for Medium Differences in Residuals

Model	AIC	BIC	CFI	TLI	RMSEA	SRMR
Overall	7,382.013	7,445.232	0.997	0.995	0.016	0.018
Group Group 1	3,765.749	3,818.571	0.976	0.953	0.047	0.031
Group Group 2	3,602.905	3,655.727	1.000	1.013	0.000	0.023
Configural	7,368.654	7,495.092	0.992	0.983	0.028	0.027
Loadings	7,364.904	7,474.483	0.990	0.986	0.025	0.036
Intercepts	7,358.503	7,451.224	1.000	1.001	0.000	0.037
Residuals	7,385.958	7,457.607	0.864	0.881	0.075	0.098

 $[tbp] \label{eq:tbp}$

 $\begin{tabular}{ll} \textbf{Table C10} \\ Model Fit for Large Differences in Residuals \\ \end{tabular}$

Model	AIC	BIC	CFI	TLI	RMSEA	SRMR
Overall	7,300.856	7,364.075	0.998	0.995	0.015	0.018
Group Group 1	3,765.749	3,818.571	0.976	0.953	0.047	0.031
Group Group 2	3,453.099	3,505.921	0.954	0.908	0.069	0.035
Configural	7,218.848	7,345.287	0.965	0.929	0.059	0.033
Loadings	7,217.332	7,326.912	0.955	0.935	0.056	0.045
Intercepts	7,211.566	7,304.287	0.962	0.958	0.046	0.047
Residuals	7,304.566	7,376.215	0.562	0.619	0.137	0.189

Appendix D

Simulated Partial Invariance Results

[tbp]

Table D1

Fit Estimates for Partial Invariance

Residuals on Invariant Data

Estimated Parameter	CFI	RSMEA
q1 ~~ q1	0.990	0.021
q2 ~~ q2	0.987	0.024
q3 ~~ q3	0.996	0.014
q4 ~~ q4	1.000	0.000
q5 ~~ q5	0.987	0.025
lv ~~ lv	0.991	0.020

Table D2

Fit Estimates for Partial Invariance

Loadings for Small Loading Data

Estimated Parameter	CFI	RSMEA
lv =~ q1	0.993	0.019
$lv = \sim q2$	0.989	0.023
lv =~ q3	0.989	0.023
$lv = \sim q4$	1.000	0.000
$lv = \sim q5$	0.994	0.017

[tbp]

Table D3

Fit Estimates for Partial Invariance

Loadings for Medium Loading Data

Estimated Parameter	CFI	RSMEA
$lv = \sim q1$	0.880	0.079
$lv = \sim q2$	0.898	0.074
$lv = \sim q3$	0.878	0.081
$lv = \sim q4$	0.962	0.045
lv =~ q5	0.907	0.071

 $[tbp] \label{eq:tbp}$

Table D4

Fit Estimates for Partial Invariance

Loadings for Large Loading Data

Estimated Parameter	CFI	RSMEA
lv =~ q1	0.762	0.116
$lv = \sim q2$	0.770	0.117
$lv = \sim q3$	0.762	0.119
$lv = \sim q4$	0.971	0.041
lv =~ q5	0.842	0.097

 $[tbp] \centering % \begin{center} \b$

Table D5

Fit Estimates for Partial Invariance

Loadings for Small Intercept Data

Estimated Parameter	CFI	RSMEA
q1 ~1	0.975	0.033
lv ~1	0.975	0.033
q2 ~1	0.972	0.035
q3 ~1	0.972	0.036
q4 ~1	0.988	0.023
q5 ~1	0.971	0.036

Table D6

Fit Estimates for Partial Invariance

Loadings for Medium Intercept Data

Estimated Parameter	CFI	RSMEA
q1 ~1	0.905	0.065
lv ~1	0.905	0.065
q2 ~1	0.901	0.067
q3 ~1	0.901	0.067
q4 ~1	0.988	0.023
q5 ~1	0.902	0.067

[tbp]

Table D7

Fit Estimates for Partial Invariance

Loadings for Large Intercept Data

Estimated Parameter	CFI	RSMEA
q1 ~1	0.785	0.097
lv ~1	0.785	0.097
q2 ~1	0.781	0.100
q3 ~1	0.781	0.100
q4 ~1	0.988	0.023
q5 ~1	0.784	0.099

Table D8

Fit Estimates for Partial Invariance

Loadings for Small Residual Data

Estimated Parameter	CFI	RSMEA
q1 ~~ q1	0.928	0.056
q2 ~~ q2	0.936	0.053
q3 ~~ q3	0.926	0.057
q4 ~~ q4	0.955	0.044
q5 ~~ q5	0.926	0.057
lv ∼∼ lv	0.930	0.054

[tbp]

Table D9

Fit Estimates for Partial Invariance

Loadings for Medium Residual Data

Estimated Parameter	CFI	RSMEA
q1 ~~ q1	0.869	0.075
q2 ~~ q2	0.860	0.078
q3 ~~ q3	0.870	0.075
q4 ~~ q4	0.994	0.016
q5 ~~ q5	0.862	0.077
lv ~~ lv	0.864	0.075

Table D10

Fit Estimates for Partial Invariance

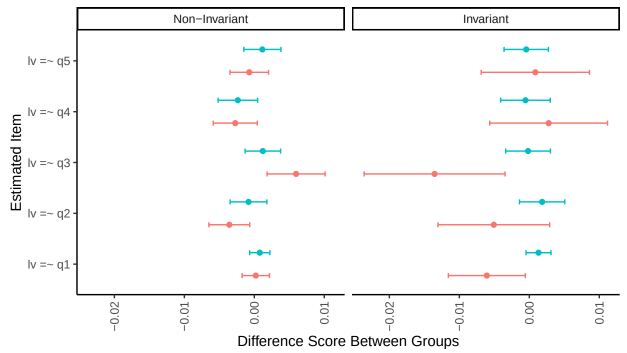
Loadings for Large Residual Data

Estimated Parameter	CFI	RSMEA
q1 ~~ q1	0.558	0.140
q2 ~~ q2	0.559	0.140
q3 ~~ q3	0.560	0.140
q4 ~~ q4	0.972	0.035
q5 ~~ q5	0.559	0.140
lv ∼∼ lv	0.562	0.137

Appendix E

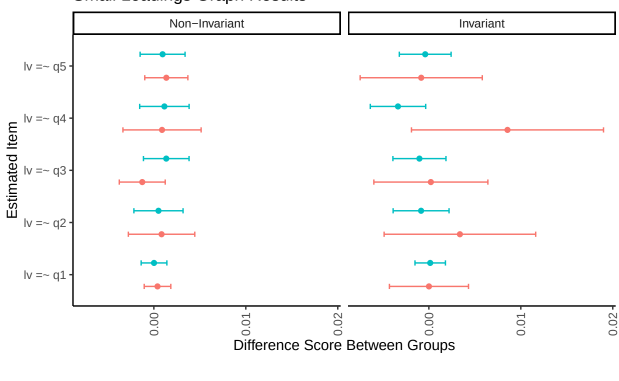
Invariance Plots Difference Scores by Condition

Invariant Graph Results



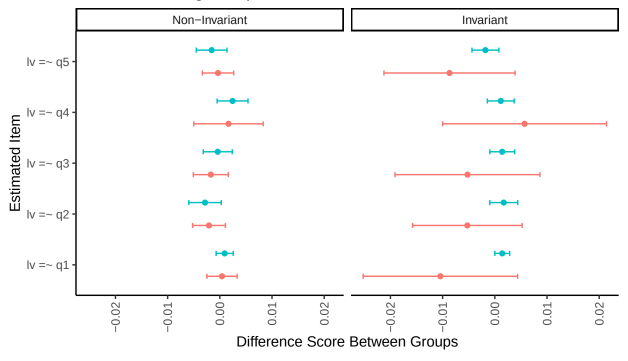
Type of Estimate → Bootstrapped → Random

Small Loadings Graph Results



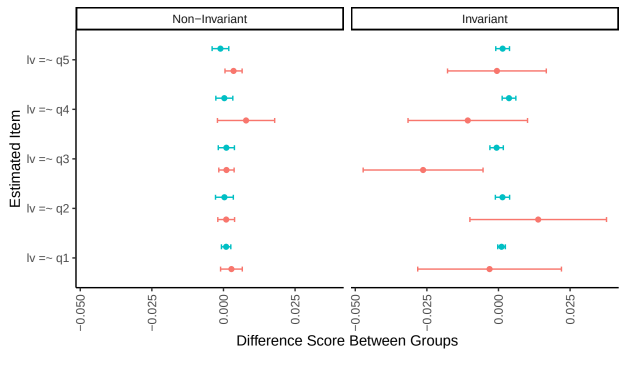
Type of Estimate - Bootstrapped - Random

Medium Loadings Graph Results



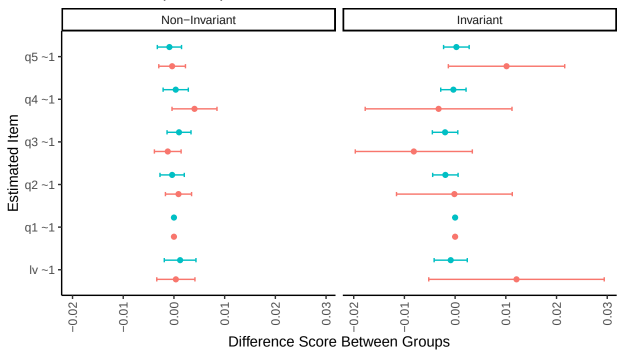
Type of Estimate → Bootstrapped → Random

Large Loadings Graph Results



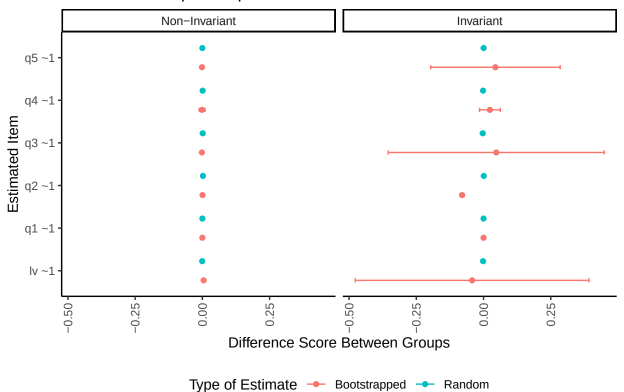
Type of Estimate → Bootstrapped → Random

Small Intercepts Graph Results

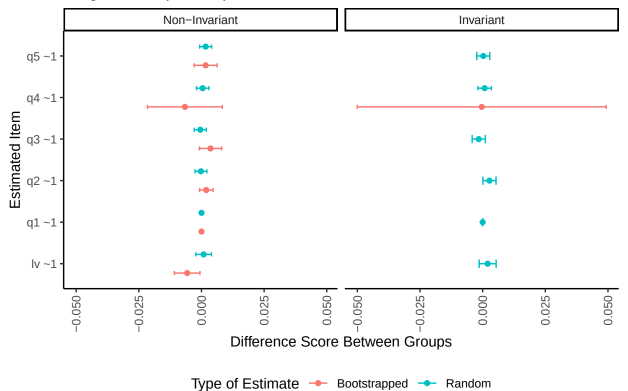


Type of Estimate → Bootstrapped → Random

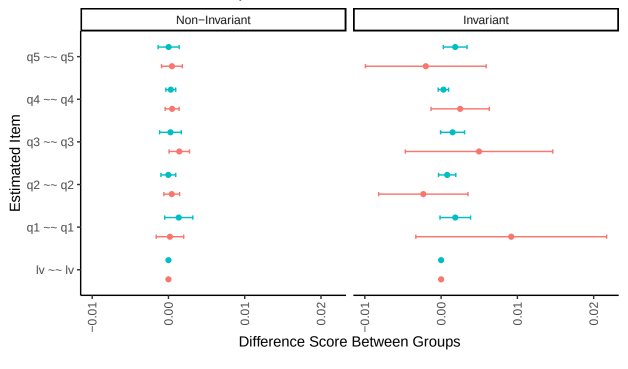
Medium Intercepts Graph Results



Large Intercepts Graph Results

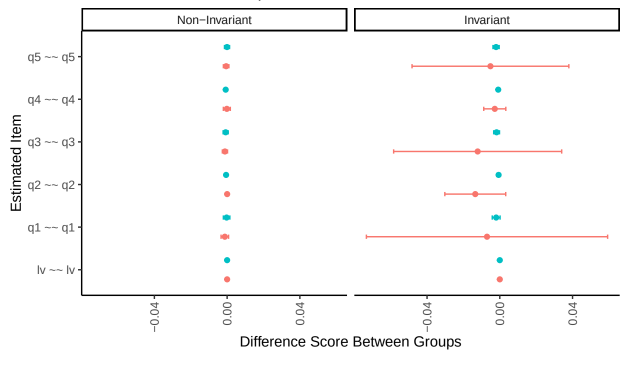


Small Residuals Graph Results



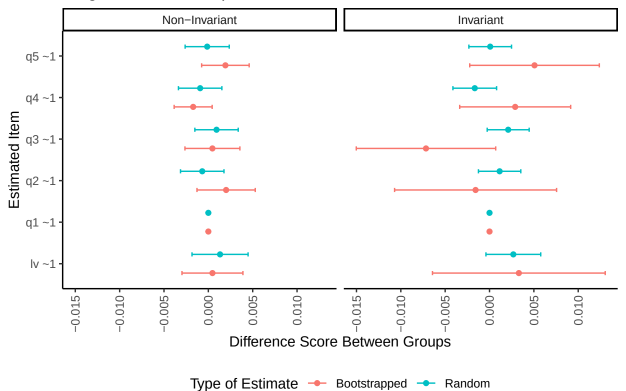
Type of Estimate → Bootstrapped → Random

Medium Residuals Graph Results



Type of Estimate → Bootstrapped → Random

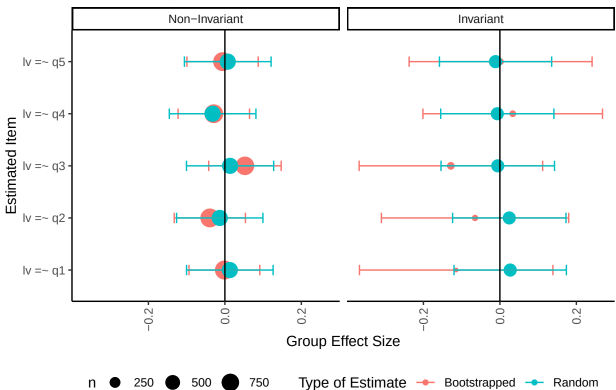
Large Residuals Graph Results



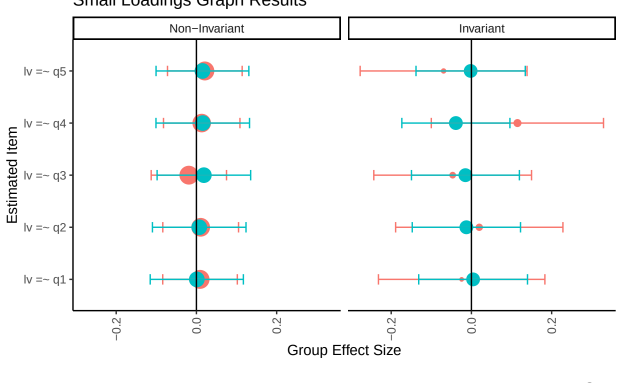
Appendix F

Invariance Plots Effect Sizes by Condition

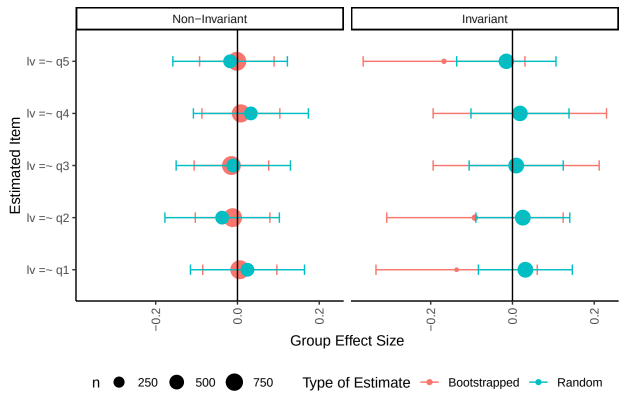
Invariant Graph Results



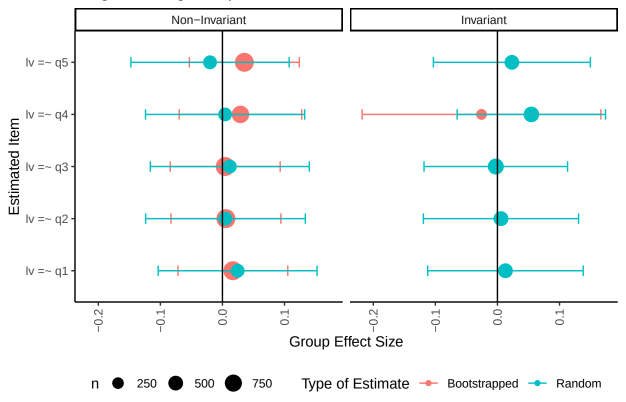
Small Loadings Graph Results



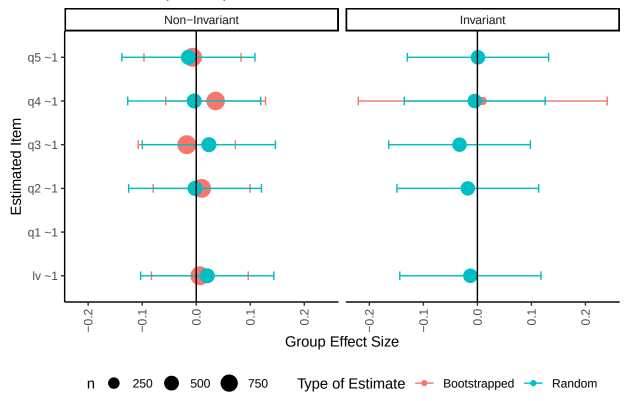
Medium Loadings Graph Results



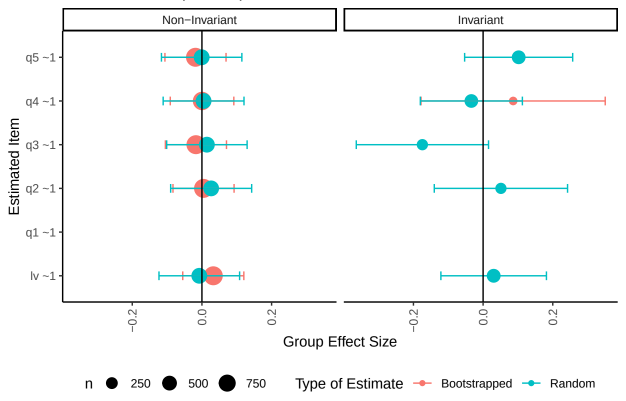
Large Loadings Graph Results



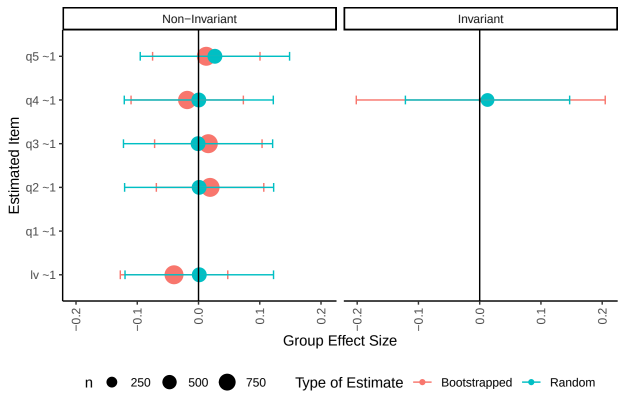
Small Intercepts Graph Results



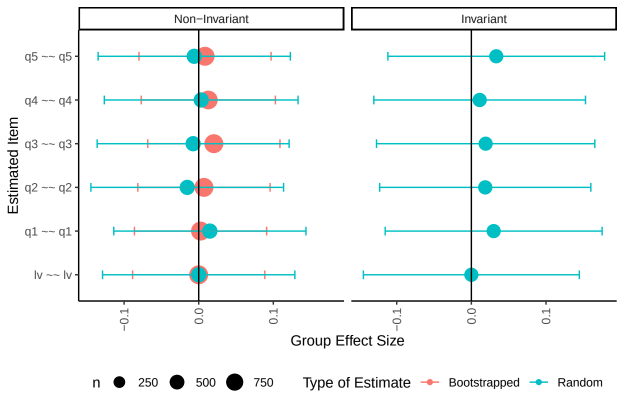
Medium Intercepts Graph Results



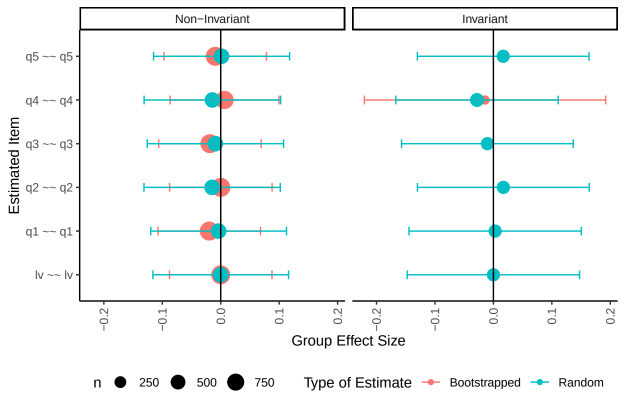
Large Intercepts Graph Results



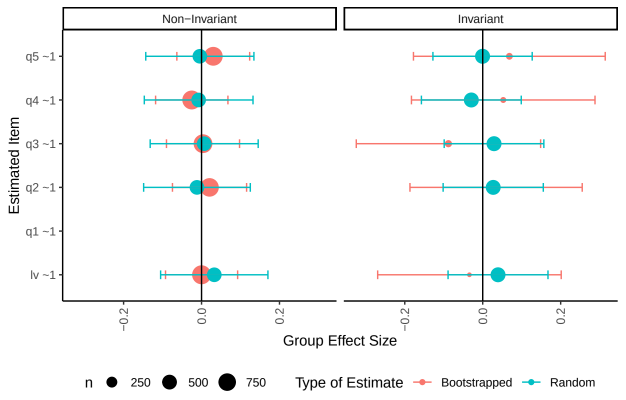
Small Residuals Graph Results



Medium Residuals Graph Results



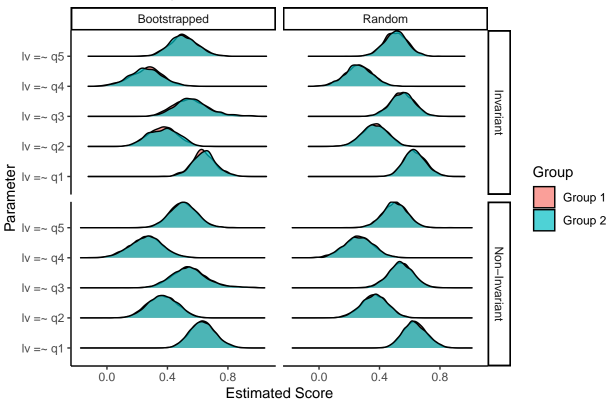
Large Residuals Graph Results



Appendix G

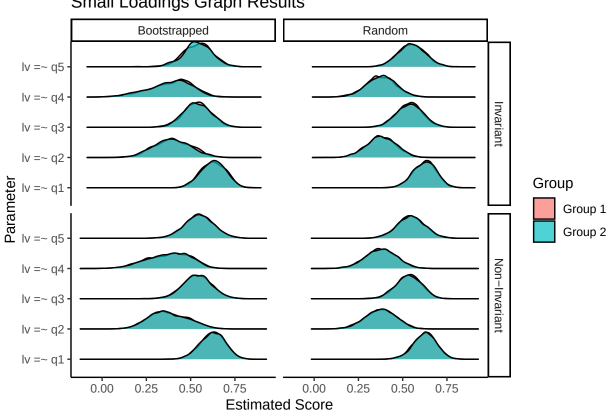
Density Plots by Condition

Invariant Graph Results

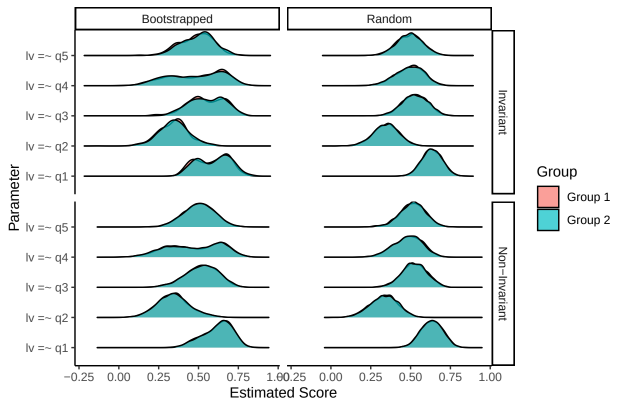


912

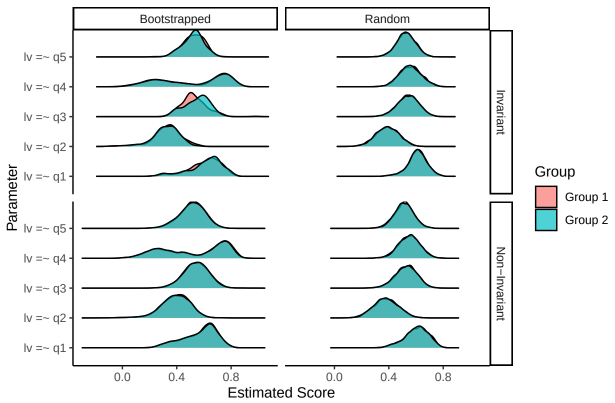
Small Loadings Graph Results



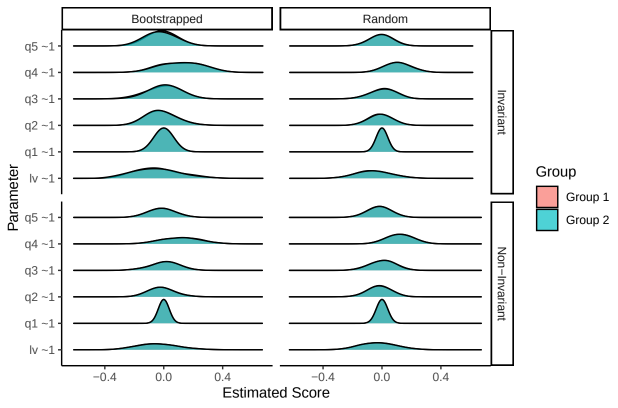
Medium Loadings Graph Results



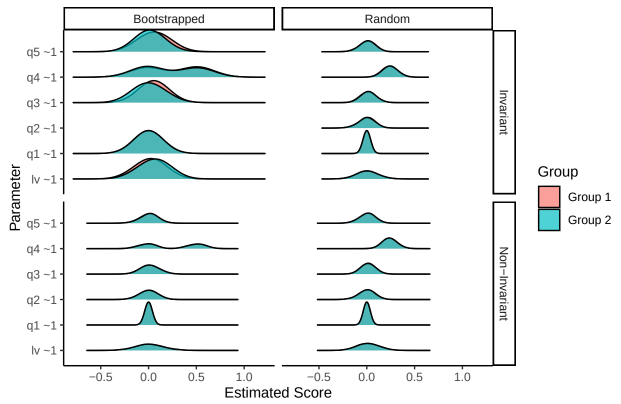
Large Loadings Graph Results



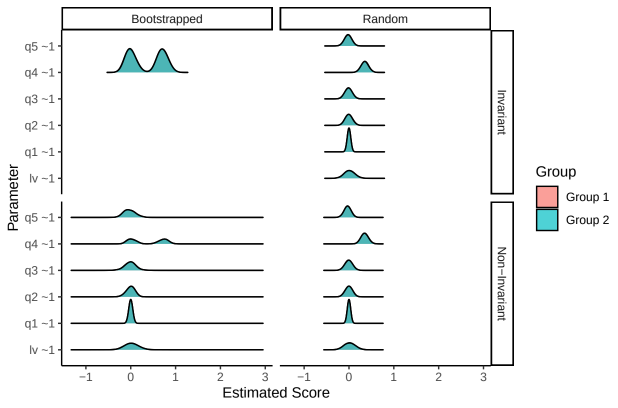
Small Intercepts Graph Results



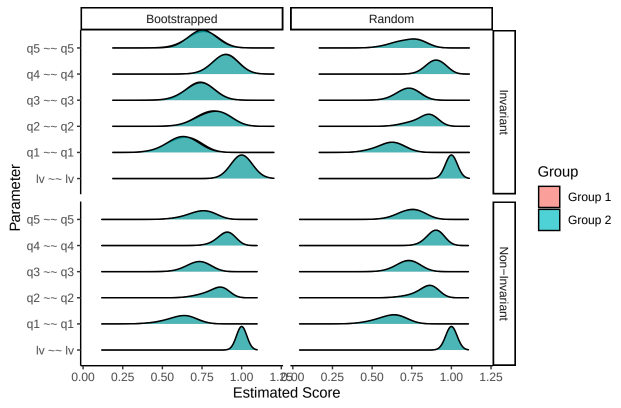
Medium Intercepts Graph Results



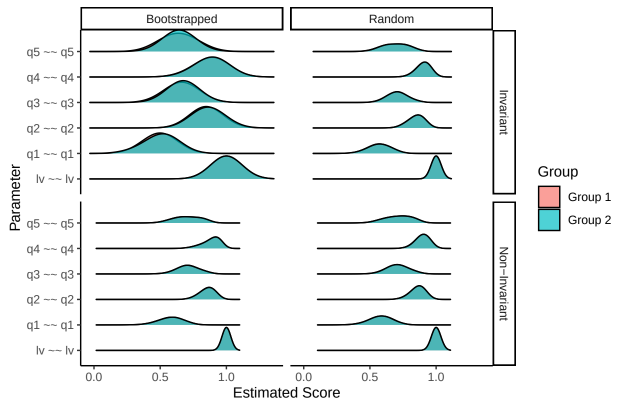
Large Intercepts Graph Results



Small Residuals Graph Results



Medium Residuals Graph Results



Large Residuals Graph Results

