

The ‘Structural After Measurement’ (SAM) approach to SEM

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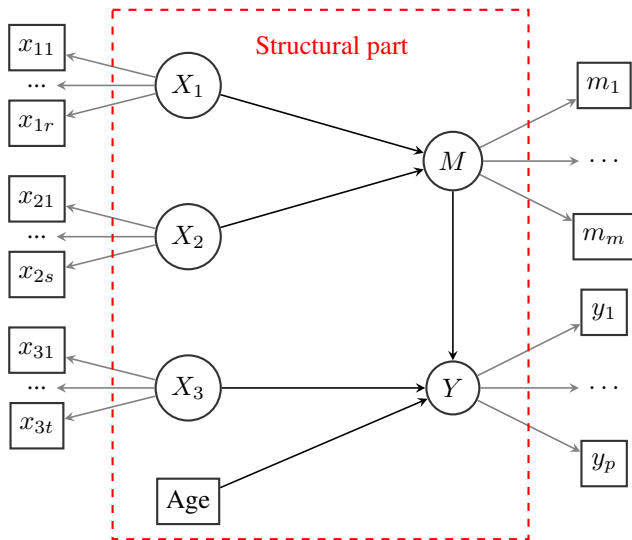
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1 Introduction

1.1 Overview of the workshop

- provide an overview of SAM approaches:
 - replacing the latent variables by ('factor') scores
 - computing summary statistics—the mean vector and (co)variance matrix—for the latent variables
 - the single-indicator (SI) approach
 - global SAM
- why/when should we (not) consider SAM approaches?
- special attention: local SAM

1.2 The setting



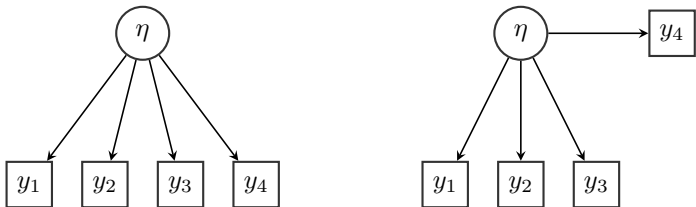
- we focus on ‘large’ models with many (say, > 100) parameters:
 - many constructs (motivation, ability, personality traits, ...)
 - each construct is measured by a set of (observed) indicators
 - many ‘background’ variables (age, gender, ...)
 - multilevel data, missing data, ...
- we are mostly interested in the structural part of the model:
 - if not saturated: how well does the structural part fit?
 - size of direct/indirect effects, hypothesis testing
- **assumption: the measurement instruments for the latent variables are well established, and fit (reasonably) well**
 - SAM approaches should not be used to ‘test’ the quality of your measurement instruments
- in addition: the sample size may not be very large (say, $N = 150$)

1.3 The standard estimation approach in SEM

- all parameters (measurement and structural) are estimated jointly
- we call this ‘joint’ or ‘system-wide’ estimation (Bollen, 1996)
- frequentist: typically using an iterative optimization approach
 - maximum likelihood (normal or Wishart) (ML)
 - generalized least squares (GLS)
 - (diagonally) weighted least squares, (D)WLS
- Bayesian: typically using MCMC
- advantages:
 - one-step, and therefore efficient (in terms of sampling variability)
 - inference is straightforward (standard errors, hypothesis testing)
 - (relatively) easy to handle constraints, missing data, ...
 - well-known, well-studied
 - widely available in software

- works very well if the following conditions are met:
 - correctly specified model
 - large sample size
 - (normally distributed data)
- but under less ideal circumstances, system-wide estimation does not (always) work well:
 - bias
 - instability (small changes in the data lead to very different results)
 - nonconvergence
 - improper solutions
- in addition: combining ‘structural’ and ‘measurement’ is not a good idea, due to a phenomenon called ‘interpretational confounding’

example interpretational confounding (stolen from Roy Levy, 2023)



- left panel: only measurement
- right panel: measurement + structural
- conceptually very different
- mathematically identical (in system-wide SEM)

generate some data

```
> library(lavaan)
> Sigma <- matrix(c(2.0, 1.0, 1.0, 1.0, 1.5,
                  1.0, 2.0, 1.0, 1.0, 0.1,
                  1.0, 1.0, 2.0, 1.0, 0.1,
                  1.0, 1.0, 1.0, 2.0, 0.1,
                  1.5, 0.1, 0.1, 0.1, 2.0), nrow = 5, ncol = 5)
> rownames(Sigma) <- colnames(Sigma) <- c("y1", "y2", "y3", "y4", "z")
> Sigma
```

```
      y1 y2 y3 y4 z
y1 2.0 1.0 1.0 1.0 1.5
y2 1.0 2.0 1.0 1.0 0.1
y3 1.0 1.0 2.0 1.0 0.1
y4 1.0 1.0 1.0 2.0 0.1
z   1.5 0.1 0.1 0.1 2.0
```

```
> set.seed(3)
> Data <- MASS::mvrnorm(n = 200L, mu = rep(0, 5), Sigma = Sigma)
```

R code left panel (model1)

```
> model1 <- '
  f =~ y1 + y2 + y3 + y4
,
> fit1 <- sem(model1, data = Data)
> summary(fit1)
```

lavaan 0.6-22.2560 ended normally after 25 iterations

Estimator	ML
Optimization method	NLMINB
Number of model parameters	8
Number of observations	200

Model Test User Model:

Test statistic	0.140
Degrees of freedom	2
P-value (Chi-square)	0.932

Parameter Estimates:

Standard errors	Standard
Information	Expected
Information saturated (h1) model	Structured

Latent Variables:

	Estimate	Std.Err	z-value	P(> z)
f =~				
y1	1.000			
y2	0.996	0.123	8.122	0.000
y3	0.894	0.116	7.689	0.000
y4	0.911	0.120	7.608	0.000

Variances:

	Estimate	Std.Err	z-value	P(> z)
.y1	1.096	0.148	7.388	0.000
.y2	0.837	0.127	6.601	0.000
.y3	0.987	0.128	7.694	0.000
.y4	1.081	0.138	7.820	0.000
f	1.031	0.208	4.959	0.000

R code right panel (model2)

```
> model2 <- '
  f =~ y1 + y2 + y3
  y4 ~ f
',
> fit2 <- sem(model2, data = Data)
> summary(fit2)
```

lavaan 0.6-22.2560 ended normally after 26 iterations

Estimator	ML
Optimization method	NLMINB
Number of model parameters	8
Number of observations	200

Model Test User Model:

Test statistic	0.140
Degrees of freedom	2
P-value (Chi-square)	0.932

Parameter Estimates:

Standard errors	Standard
Information	Expected
Information saturated (h1) model	Structured

Latent Variables:

	Estimate	Std.Err	z-value	P(> z)
f =~				
y1	1.000			
y2	0.996	0.123	8.122	0.000
y3	0.894	0.116	7.689	0.000

Regressions:

	Estimate	Std.Err	z-value	P(> z)
y4 ~				
f	0.911	0.120	7.608	0.000

Variances:

	Estimate	Std.Err	z-value	P(> z)
.y1	1.096	0.148	7.388	0.000
.y2	0.837	0.127	6.601	0.000
.y3	0.987	0.128	7.694	0.000
.y4	1.081	0.138	7.820	0.000
f	1.031	0.208	4.959	0.000

change outcome variable (y4 becomes z) (model3)

```
> model3 <- '
  f =~ y1 + y2 + y3
  z ~ f
',
> fit3 <- sem(model3, data = Data)
> summary(fit3)
```

lavaan 0.6-22.2560 ended normally after 58 iterations

Estimator	ML
Optimization method	NLMINB
Number of model parameters	8
Number of observations	200

Model Test User Model:

Test statistic	50.659
Degrees of freedom	2
P-value (Chi-square)	0.000

Parameter Estimates:

Standard errors	Standard
Information	Expected
Information saturated (h1) model	Structured

Latent Variables:

	Estimate	Std.Err	z-value	P(> z)
f =~				
y1	1.000			
y2	0.138	0.053	2.601	0.009
y3	0.133	0.051	2.591	0.010

Regressions:

	Estimate	Std.Err	z-value	P(> z)
z ~				
f	0.276	0.087	3.156	0.002

Variances:

	Estimate	Std.Err	z-value	P(> z)
.y1	-4.169	1.801	-2.315	0.021
.y2	1.740	0.173	10.071	0.000
.y3	1.699	0.168	10.086	0.000
.z	1.677	0.211	7.953	0.000
f	6.296	1.715	3.671	0.000

interpretational confounding

- replacing y_4 by z (= changing the structural part) also changes the parameters of the measurement model
- if the resulting parameters of the measurement model imply a different ‘meaning’ of the latent variable than was intended by the researcher, we have a problem
- this problem was coined “interpretational confounding” by Burt (1976)

Burt, R.S. (1976). Interpretational confounding of unobserved variables in structural equation models. *Sociological Methods & Research*, 5(1), 3–52.

- Burt (1976) already suggested the solution: first fit the measurement part of the model, and then fit the structural part of the model

solution: replace sem() by sam()

```
> fit3.sam <- sam(model3, data = Data)
> parameterEstimates(fit3.sam, remove.step1 = FALSE, ci = FALSE,
  output = "text")
```

Latent Variables:

	Step	Estimate	Std.Err	z-value	P(> z)
f =~					
y1	1	1.000			
y2	1	0.969	0.139	6.977	0.000
y3	1	0.884	0.127	6.979	0.000

Regressions:

	Step	Estimate	Std.Err	z-value	P(> z)
z ~					
f	2	0.540	0.121	4.471	0.000

Variances:

	Step	Estimate	Std.Err	z-value	P(> z)
.y1	1	1.067	0.170	6.287	0.000
.y2	1	0.865	0.151	5.723	0.000
.y3	1	0.983	0.142	6.900	0.000
.z	2	1.846	0.195	9.462	0.000
f	2	1.059	0.226	4.678	0.000

1.4 Running example

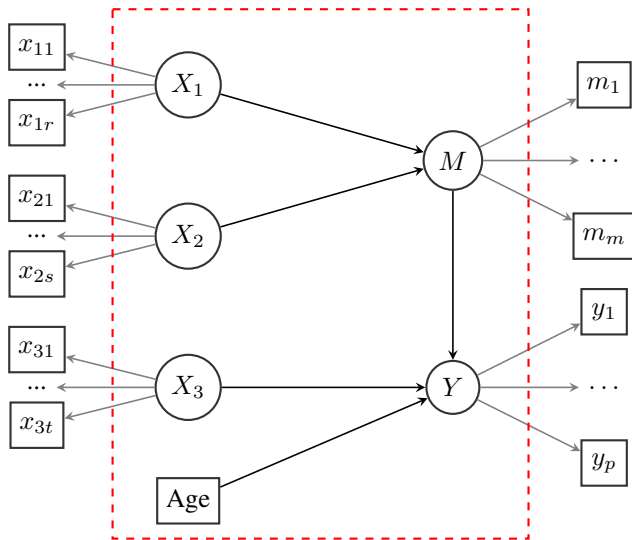
- throughout the workshop, we will make use of a simple ‘latent mediation’ example to illustrate the various estimation approaches
- we generate ‘perfect’ (population-level) data

```
> library(lavaan)
> pop.model <- '
  # factor loadings
  Y =~ 1*y1 + 1.2*y2 + 0.8*y3 + 0.5*y4
  M =~ 1*m1 + 0.5*m2 + 0.5*m3 + 0.7*m4
  X1 =~ 1*x1 + 0.7*x2 + 0.6*x3 + 1.1*x4
  X2 =~ 1*x5 + 0.7*x6 + 0.6*x7 + 0.9*x8
  X3 =~ 1*x9 + 0.7*x10 + 0.6*x11 + 1.1*x12

  # covariances among exogenous X1-X3 and Age
  X1 ~~ 0.4*X2; X1 ~~ -0.2*X3; X2 ~~ 0.4*X3
  Age ~~ 0.6*X1 + 0.7*X2 + 0.2*X3

  # regression part
  Y ~ 0.1*X3 + 0.2*M + (-0.3)*Age
  M ~ -0.40*X1 + 0.5*X2
  ,
> set.seed(1234)
> Data <- simulateData(pop.model, sample.nobs = 200L, empirical = TRUE)
```

diagram



fitting the model using traditional SEM

```

> model <- '
  # measurement part
  Y =~ y1 + y2 + y3 + y4
  M =~ m1 + m2 + m3 + m4
  X1 =~ x1 + x2 + x3 + x4
  X2 =~ x5 + x6 + x7 + x8
  X3 =~ x9 + x10 + x11 + x12

  # structural part
  Y ~ X3 + M + Age
  M ~ X1 + X2
  Age ~~ X1 + X2 + X3
,
> fit.sem <- sem(model, data = Data, estimator = "ML")
> parameterEstimates(fit.sem, ci = FALSE, output = "text")[21:25,]

```

Regressions:

	Estimate	Std.Err	z-value	P(> z)
Y ~				
X3	0.100	0.101	0.994	0.320
M	0.200	0.094	2.137	0.033
Age	-0.300	0.086	-3.472	0.001
M ~				
X1	-0.400	0.125	-3.202	0.001
X2	0.500	0.128	3.895	0.000

upgrading Age to a latent variable (for convenience only)

```

> model2 <- '
  # measurement part
  Y =~ y1 + y2 + y3 + y4
  M =~ m1 + m2 + m3 + m4
  X1 =~ x1 + x2 + x3 + x4
  X2 =~ x5 + x6 + x7 + x8
  X3 =~ x9 + x10 + x11 + x12
  fAge =~ 1*Age; Age ~~ 0*Age

  # structural part
  Y ~ X3 + M + fAge
  M ~ X1 + X2
,
> fit.sem2 <- sem(model2, data = Data, estimator = "ML")
> parameterEstimates(fit.sem2, ci = FALSE, output = "text")[23:27,]

```

Regressions:

	Estimate	Std.Err	z-value	P(> z)
Y ~				
X3	0.100	0.101	0.994	0.320
M	0.200	0.094	2.137	0.033
fAge	-0.300	0.086	-3.472	0.001
M ~				
X1	-0.400	0.125	-3.202	0.001
X2	0.500	0.128	3.895	0.000

model-implied variance-covariance matrix latent variables

- the standard way (in lavaan) to extract the (co)variance matrix of the latent variables:

```
> lavInspect(fit.sem, "cov.lv")  
  
      Y      M      X1      X2      X3  
Y  1.136  
M  0.245  1.250  
X1 -0.240 -0.200  1.000  
X2 -0.102  0.340  0.400  1.000  
X3  0.096  0.280 -0.200  0.400  1.000
```

- unfortunately, this matrix only contains the latent variables, not the observed covariate 'Age'
- that is why (in fit.sem2) we 'upgraded' Age to a latent variable (fAge); lavaan now thinks it is a latent variable, and includes it in the (co)variance matrix of the latent variables

```
> Var.eta <- lavInspect(fit.sem2, "cov.lv")  
> Var.eta
```

	Y	M	X1	X2	X3	fAge
Y	1.136					
M	0.245	1.250				
X1	-0.240	-0.200	1.000			
X2	-0.102	0.340	0.400	1.000		
X3	0.096	0.280	-0.200	0.400	1.000	
fAge	-0.258	0.110	0.600	0.700	0.200	1.000

- fact: if we could somehow compute this matrix, then we can obtain all the regression coefficients using two regressions (one for 'Y', and one for 'M')
- recall the well known formula to estimate the regression coefficients in a linear model:

$$\hat{\beta} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'\mathbf{y}$$

- an equivalent formula based on the sample covariance matrix (\mathbf{S}):

$$\hat{\beta} = (\mathbf{S}_{xx})^{-1}\mathbf{S}_{xy}$$

where \mathbf{S}_{xx} is the part of \mathbf{S} where the rows and columns correspond to the predictors ('x'), while \mathbf{S}_{xy} is the part of \mathbf{S} where the rows correspond to the predictors and the (single) column corresponds to the outcome ('y')

- regression for 'Y':

```
> # compute regression coefficients for Y
> beta.Y <- ( solve(Var.eta[c("X3", "M", "fAge"), c("X3", "M", "fAge")]) %*%
              Var.eta[c("X3", "M", "fAge"), "Y", drop = FALSE] )
> round(beta.Y, 3)
```

```
      Y
X3    0.1
M     0.2
fAge -0.3
```

- regression for 'M':

```
> # compute regression coefficients for M
> beta.M <- ( solve(Var.eta[c("X1", "X2"), c("X1", "X2")]) %*%
              Var.eta[c("X1", "X2"), "M", drop = FALSE] )
> round(beta.M, 3)
```

```
      M
X1 -0.4
X2  0.5
```

2 Structural-after-measurement (SAM) approaches

2.1 Introduction

- SAM is an umbrella term to describe many different (estimation) approaches that have the following in common:
 1. first step: we estimate the parameters related to the measurement part
 2. second step: we estimate the parameters related to the structural part
- SAM is an estimation approach, not a model-building approach; the main goal is to find (good quality) point estimates for the structural (i.e., regression) parameters in the model
- ideally, we also wish to obtain (good quality) standard errors and test statistics for these structural parameters (for statistical inference)
- the term SAM was used by Rosseel & Loh (2024), to avoid the overloaded terms ‘two-step’, ‘two-stage’, ... (and was first used during a presentation I gave at WWU Münster, Jan 2020)

2.2 Early SAM approaches

- some early references:
 - Burt (1976)
 - Hunter & Gerbing (1982)
 - Lance, Cornwell & Mulaik (1988)
- SAM approaches never received much attention in the SEM literature
- but SAM is the default approach in many other fields
 - ‘measurement error models’ in statistics
 - item response theory (IRT) (e.g., plausible values)
 - partial least squares (PLS)
- many applied researchers in psychology, educational sciences, ... replace latent variables by sum scores (or factor scores); this is a SAM approach

2.3 Overview of SAM approaches (for SEM)

- replacing the latent variables by ('factor') scores
 - (uncorrected) factor score regression (UFSR)
 - correlation-preserving factor scores
 - sum scores
- computing ('corrected') summary statistics for the latent variables
 - Wall & Amemiya (2000), Croon (2002), Devlieger et al. (2016, 2017)
 - local SAM (Rosseel & Loh, 2024)
- the single-indicator (SI) approach
- global SAM

2.4 Potential advantages of a SAM approach

- reduced model complexity: the ‘structural’ model is often just a regression model, or a path analysis
- if the model is correctly specified, many SAM approaches (but not all) give identical results as SEM (for the structural part)
- (almost) no convergence issues, increased stability (in particular in small samples)
- less variability (lower MSE) for structural parameters
- in certain cases (but not always): more robust against local (structural) model misspecifications
- no interpretational confounding
- very flexible: the decoupling of ‘measurement’ and ‘structural’ allows for tackling much more complicated models (even with modest sample sizes)
- ...

2.5 Critique of the SAM approach

- the (naive) standard errors in the second step are wrong (because they ignore the uncertainty that stems from the first step)
- in general: inference (e.g., standard errors, confidence intervals, test statistics) is (more) complicated
- multiple step methods are less efficient (more sampling variability)
- Fornell and Yi (1992) gave an example where a misspecified (but well-fitting) measurement model was embedded in a correctly specified structural model; but the (global) model fit of the full model suggested that the model did not fit well, thus incorrectly implying a misspecified structural model
- do (latent) constructs 'exist' on their own? or are they only meaningful when embedded in a context (i.e., a structural model)?
- software packages only allow for joint estimation

2.6 Related techniques

- latent class analysis:

Bakk, Z., & Kuha, J. (2021). Relating latent class membership to external variables: An overview. *British Journal of Mathematical and Statistical Psychology*, 74(2), 340–362.

<https://doi.org/10.1111/bmsp.12227> (Open Access)

- item response theory (IRT):

Kuha, J., & Bakk, Z. (arxiv.org). Two-step estimation of latent trait models.

<https://arxiv.org/pdf/2303.16101.pdf>

- Bayesian SAM:

Levy, R. (2023). Precluding interpretational confounding in factor analysis with a covariate or outcome via measurement and uncertainty preserving parametric modeling. *Structural Equation Modeling: A Multidisciplinary Journal*.

<https://doi.org/10.1080/10705511.2022.2154214>

3 Replacing the latent variables by ('factor') scores

- general idea: we try to compute individual scores for each latent variable that is involved in the structural part of the model
- once we have these scores, we treat them as if they were observed
- we then perform an observed-only procedure (e.g., regression or path analysis) using these scores (and perhaps other observed covariates)
- there are various ways to compute these scores:
 - we compute so-called 'factor scores' (e.g., Bartlett or regression factor scores) for all the latent variables
 - we compute the sum or the mean of the latent variable indicators
 - we just take the observed scores of the first ('marker') indicator of the latent variable

3.1 Factor scores

- the standard factor model:

$$\mathbf{y} = \mathbf{\Lambda}\boldsymbol{\eta} + \boldsymbol{\epsilon}$$

- \mathbf{y} and $\boldsymbol{\epsilon}$ are P -dimensional continuous random vectors
 - $\mathbf{\Lambda}$ is a $P \times M$ matrix of factor loadings
 - $\boldsymbol{\eta}$ is a M -dimensional random vector of latent variables
 - we denote $\text{Var}(\boldsymbol{\epsilon}) = \boldsymbol{\Theta}$
- we can use CFA to estimate all the model parameters
 - here, we only consider linear factor scores of the form

$$\mathbf{f} = \hat{\boldsymbol{\eta}} = \mathbf{A}\mathbf{y}$$

- different types of factor scores can be constructed by choosing a different way to compute the $M \times P$ ‘factor score matrix’ \mathbf{A}
- three important types: 1) ‘regression’ factor scores, 2) Bartlett factor scores, and 3) correlation-preserving factor scores

regression factor scores (Thomson, 1934)

- Thomson regarded the quest for factor scores as a (multivariate) regression problem: we wish to 'predict' $\boldsymbol{\eta}$ as a (linear) function of the observed variables \mathbf{y} :

$$\boldsymbol{\eta} = \mathbf{A}\mathbf{y} + \mathbf{r}$$

- it is well known that the OLS solution is given by

$$\mathbf{A} = \text{Cov}(\boldsymbol{\eta}, \mathbf{y})\text{Var}(\mathbf{y})^{-1}$$

- because $\text{Var}(\mathbf{y}) = \boldsymbol{\Sigma}$ and $\text{Cov}(\boldsymbol{\eta}, \mathbf{y}) = \text{Var}(\boldsymbol{\eta})\boldsymbol{\Lambda}'$, this results in the following well known expression for \mathbf{A} :

$$\mathbf{A}_{reg} = \text{Var}(\boldsymbol{\eta})\boldsymbol{\Lambda}'\boldsymbol{\Sigma}^{-1}$$

- an equivalent expression is

$$\mathbf{A}_{reg} = (\text{Var}(\boldsymbol{\eta})^{-1} + \boldsymbol{\Lambda}'\boldsymbol{\Theta}^{-1}\boldsymbol{\Lambda})^{-1}\boldsymbol{\Lambda}'\boldsymbol{\Theta}^{-1}$$

- properties:
 - regression factor scores are ‘best’ in the sense that they minimize the MSE
 - but they are not (conditionally) unbiased: $E(\mathbf{f}|\boldsymbol{\eta}) \neq \boldsymbol{\eta}$
 - $\text{Var}(\mathbf{f})$ is *not* equal to $\text{Var}(\boldsymbol{\eta})$ (i.e., not correlation-preserving)
- computing regression factor scores in lavaan:

```
> fs.reg <- lavPredict(fit.sem)
> head(fs.reg)
```

```
      Y           M           X1           X2           X3
[1,] 0.5328842  1.3082196 -2.5327939 -2.098782 -0.7740726
[2,] 0.3395217 -0.6002887  0.1356861 -0.408424  0.2346622
[3,] 0.2342185  0.5654211 -1.0291976 -1.755758 -1.4171193
[4,] 0.6488077  1.2737336  0.2993125  1.157076  0.9422088
[5,] -0.4810281  1.2828249  0.4650392  1.207516 -0.1140724
[6,] -1.2513209 -1.1338941 -0.1388611 -1.528747 -2.1519342
```

- very popular; default in many software packages

Bartlett (1937, 1938) factor scores

- Bartlett used the principle of maximum likelihood to find ‘good’ factor scores
- given \mathbf{y} , $\mathbf{\Lambda}$ and $\mathbf{\Theta}$, we need to find $\boldsymbol{\eta}$ that maximizes the following function:

$$f(\boldsymbol{\eta}) = \text{const} - \frac{1}{2}(\mathbf{y} - \mathbf{\Lambda}\boldsymbol{\eta})^T \mathbf{\Theta}^{-1}(\mathbf{y} - \mathbf{\Lambda}\boldsymbol{\eta})$$

- equivalently, we can minimize the weighted least squares (WLS) criterion

$$f(\boldsymbol{\eta}) = (\mathbf{y} - \mathbf{\Lambda}\boldsymbol{\eta})^T \mathbf{\Theta}^{-1}(\mathbf{y} - \mathbf{\Lambda}\boldsymbol{\eta})$$

- it is well known that the WLS solution for $\boldsymbol{\eta}$ is given by

$$\hat{\boldsymbol{\eta}} = \mathbf{f} = (\mathbf{\Lambda}^T \mathbf{\Theta}^{-1} \mathbf{\Lambda})^{-1} \mathbf{\Lambda}^T \mathbf{\Theta}^{-1} \mathbf{y}$$

- therefore, the factor score matrix used for Bartlett factor scores is given by

$$\mathbf{A}_{bar} = (\mathbf{\Lambda}^T \mathbf{\Theta}^{-1} \mathbf{\Lambda})^{-1} \mathbf{\Lambda}^T \mathbf{\Theta}^{-1}$$

- properties:
 - the MSE is (slightly) larger than the MSE for regression factor scores
 - Bartlett factor scores are conditionally unbiased, that is $E(\mathbf{f}|\boldsymbol{\eta}) = \boldsymbol{\eta}$
 - $\text{Var}(\mathbf{f})$ is *not* equal to $\text{Var}(\boldsymbol{\eta})$ (i.e., not correlation-preserving)
- computing Bartlett factor scores in lavaan:

```
> fs.bar <- lavPredict(fit.sem, method = "Bartlett")
> head(fs.bar)
```

	Y	M	X1	X2	X3
[1,]	-0.1058129	2.2481161	-1.88120459	-1.1520656	-0.6544485
[2,]	0.4105007	-0.7917271	0.43392760	-0.5737966	0.5104814
[3,]	-0.1764400	1.2545300	-0.35873150	-1.3077973	-1.5393525
[4,]	0.9902981	1.5518666	0.01350250	0.8634053	0.9801388
[5,]	-0.4058483	1.6724513	-0.07721199	1.0751197	-0.5591864
[6,]	-1.8000568	-1.1430745	0.19928157	-1.1348063	-2.5876070

correlation-preserving factor scores

- correlation-preserving factor scores have the property that $\text{Var}(\mathbf{f}) = \text{Var}(\boldsymbol{\eta})$
- a better name would have been ‘covariance-preserving’ or ‘structure-preserving’ factor scores
- first introduced by Anderson & Rubin (1956) for orthogonal factors
- general case: Green (1969), McDonald (1981), Krijnen, Wansbeek and ten Berge (1996)
- the ‘regression’ version (Green 1969):

$$\mathbf{A}_{green} = \text{Var}(\boldsymbol{\eta})^{1/2} (\text{Var}(\boldsymbol{\eta})^{3/2} \boldsymbol{\Lambda}^T \boldsymbol{\Sigma}^{-1} \boldsymbol{\Lambda} \text{Var}(\boldsymbol{\eta})^{3/2})^{-1/2} \text{Var}(\boldsymbol{\eta})^{3/2} \boldsymbol{\Lambda}' \boldsymbol{\Sigma}^{-1/2} \boldsymbol{\Sigma}^{-1/2}$$

- the ‘Bartlett’ version (Krijnen’s determinant best version):

$$\mathbf{A}_{det} = \text{Var}(\boldsymbol{\eta})^{1/2} (\text{Var}(\boldsymbol{\eta})^{1/2} \boldsymbol{\Lambda}^T \boldsymbol{\Sigma}^{-1} \boldsymbol{\Lambda} \text{Var}(\boldsymbol{\eta})^{1/2})^{-1/2} \text{Var}(\boldsymbol{\eta})^{1/2} \boldsymbol{\Lambda}' \boldsymbol{\Sigma}^{-1/2} \boldsymbol{\Sigma}^{-1/2}$$

- it turns out that we can also compute them by ‘transforming’ the regression or Bartlett factor scores

- properties:
 - MSE is the largest of the three types
 - not conditionally unbiased; $E(\mathbf{f}|\boldsymbol{\eta}) \neq \boldsymbol{\eta}$
 - but of course correlation-preserving: $\text{Var}(\mathbf{f}) = \text{Var}(\boldsymbol{\eta})$
- computing correlation-preserving (Bartlett) factor scores in lavaan:

```
> fs.cpbar <- lavPredict(fit.sem, method = "Bartlett", transform = TRUE)
> head(fs.cpbar)
```

	Y	M	X1	X2	X3
[1,]	0.2010349	1.7486519	-2.15247206	-1.5655764	-0.6855079
[2,]	0.3693186	-0.6912347	0.26897333	-0.4632965	0.3540204
[3,]	0.0199903	0.8800012	-0.67186207	-1.4787594	-1.4570860
[4,]	0.8232955	1.3953998	0.13470143	0.9961243	0.9504933
[5,]	-0.4293825	1.4569188	0.19173160	1.0971320	-0.3201049
[6,]	-1.5251428	-1.1351608	0.06119329	-1.3210353	-2.3379569

- (unfortunately) almost never used in practice

factor indeterminacy

- there is no single way to compute factor scores; there are many ways, some of them have ‘good’ properties
- this is called ‘factor indeterminacy’
- in particular when the interest is in individual factor scores, this is problematic (different factor scores may result in a different ordering of the individuals)
- but we will only be concerned with the ‘relation’ between factor scores
- therefore, all we care about is the variance-covariance matrix of the factor scores $\text{Var}(\mathbf{f})$
- the fact that $\text{Var}(\mathbf{f}) \neq \text{Var}(\boldsymbol{\eta})$ for regression and Bartlett factor scores has important consequences
- reference: Steiger (1996)

3.2 (Uncorrected) factor score regression (UFSR)

- the idea of factor score regression is simple: we replace all latent variables by factor scores
- then, in a follow-up analysis, we treat these factor scores as if they were observed
- traditionally, only regression (or Bartlett) factor scores have been used in FSR
- but this is problematic, because for these factor scores, we have that $\text{Var}(\mathbf{f}) \neq \text{Var}(\boldsymbol{\eta})$
- as a result, uncorrected FSR is not ‘consistent’: the regression coefficients that we obtain in the follow-up analysis will not match the population values, even if the model is correct and the sample size goes to infinity
- this has been well documented in the (SEM) literature, and has contributed to the widely spread opinion that FSR is not a good practice

example uncorrected FSR with regression factor scores

- first step: compute factor scores (and add observed covariates)

```
> # compute regression factor scores
> FS.reg <- as.data.frame(lavPredict(fit.sem2, method = "regression"))
```

- regression for 'Y':

```
> fit.y <- lm(Y ~ X3 + M + fAge, data = FS.reg)
> round(coef(fit.y), 3)
```

(Intercept)	X3	M	fAge
0.000	0.106	0.257	-0.308

- regression for 'M':

```
> fit.m <- lm(M ~ X1 + X2, data = FS.reg)
> round(coef(fit.m), 3)
```

(Intercept)	X1	X2
0.000	-0.532	0.644

- not the same as the population values

example uncorrected FSR with Bartlett factor scores

- first step: compute factor scores (and add observed covariates)

```
> # compute regression factor scores
> FS.bar <- as.data.frame(lavPredict(fit.sem2, method = "Bartlett"))
```

- regression for 'Y':

```
> fit.y <- lm(Y ~ X3 + M + fAge, data = FS.bar)
> round(coef(fit.y), 3)
```

(Intercept)	X3	M	fAge
0.000	0.086	0.144	-0.291

- regression for 'M':

```
> fit.m <- lm(M ~ X1 + X2, data = FS.bar)
> round(coef(fit.m), 3)
```

(Intercept)	X1	X2
0.000	-0.247	0.319

- not the same as the population values

the fundamental problem with FSR

- for regression and Bartlett factor scores, we have $\text{Var}(\mathbf{f}) \neq \text{Var}(\boldsymbol{\eta})$; as a result, all regressions based on $\text{Var}(\mathbf{f})$ will result in bias
- this is true even if the sample size goes to infinity
- recall that the ‘correct’ $\text{Var}(\boldsymbol{\eta})$ equals

```
> round(Var.eta, 3)
```

	Y	M	X1	X2	X3	fAge
Y	1.136					
M	0.245	1.250				
X1	-0.240	-0.200	1.000			
X2	-0.102	0.340	0.400	1.000		
X3	0.096	0.280	-0.200	0.400	1.000	
fAge	-0.258	0.110	0.600	0.700	0.200	1.000

- $\text{Var}(\mathbf{f})$ for regression factor scores:

```
> N <- nrow(FS.reg)
> FS.reg.cov <- cov(FS.reg) * (N - 1) / N
> round(FS.reg.cov, 3)
```

	Y	M	X1	X2	X3	fAge
Y	0.904	0.229	-0.238	-0.104	0.090	-0.258
M	0.229	0.911	-0.175	0.309	0.272	0.110
X1	-0.238	-0.175	0.798	0.388	-0.170	0.600
X2	-0.104	0.309	0.388	0.800	0.370	0.700
X3	0.090	0.272	-0.170	0.370	0.771	0.200
fAge	-0.258	0.110	0.600	0.700	0.200	1.000

- $\text{Var}(\mathbf{f})$ for Bartlett factor scores:

```
> FS.bar.cov <- cov(FS.bar) * (N - 1) / N
> round(FS.bar.cov, 3)
```

	Y	M	X1	X2	X3	fAge
Y	1.436	0.245	-0.240	-0.102	0.096	-0.258
M	0.245	1.753	-0.200	0.340	0.280	0.110
X1	-0.240	-0.200	1.327	0.400	-0.200	0.600
X2	-0.102	0.340	0.400	1.376	0.400	0.700
X3	0.096	0.280	-0.200	0.400	1.327	0.200
fAge	-0.258	0.110	0.600	0.700	0.200	1.000

- note: the Bartlett based variance-covariance matrix is very close, except for the variances of the latent variables

3.3 Skrondal & Laake (2001)

- Skrondal & Laake (2001) noticed a remarkable result: if we use (blockwise) ‘Bartlett’ factor scores for the ‘dependent’ variable(s), and (blockwise) regression factor scores for the ‘independent’ variable(s), the resulting regression coefficients are consistent again
- the regression for ‘Y’:

```
> model.y.indep <- '
  X3 =~ x9 + x10 + x11 + x12
  M =~ m1 + m2 + m3 + m4
  fAge =~ Age; Age ~~ 0*Age
'

> fit.y.indep <- cfa(model.y.indep, data = Data)
> SL.y.indep <- lavPredict(fit.y.indep, method = "regression")
> model.y.dep <- ' Y =~ y1 + y2 + y3 + y4 '
> fit.y.dep <- cfa(model.y.dep, data = Data)
> SL.y.dep <- lavPredict(fit.y.dep, method = "Bartlett")
> FS.y <- as.data.frame(cbind(SL.y.dep, SL.y.indep))
> fit.y <- lm(Y ~ X3 + M + fAge, data = FS.y)
> round(coef(fit.y), 3)
```

(Intercept)	X3	M	fAge
0.0	0.1	0.2	-0.3

- the regression for 'M':

```

> model.m.indep <- '
  X1 =~ x1 + x2 + x3 + x4
  X2 =~ x5 + x6 + x7 + x8
  ,
> fit.m.indep <- cfa(model.m.indep, data = Data)
> SL.m.indep <- lavPredict(fit.m.indep, method = "regression")
> model.m.dep <- ' M =~ m1 + m2 + m3 + m4 '
> fit.m.dep <- cfa(model.m.dep, data = Data)
> SL.m.dep <- lavPredict(fit.m.dep, method = "Bartlett")
> FS.m <- as.data.frame(cbind(SL.m.dep, SL.m.indep))
> fit.m <- lm(M ~ X1 + X2, data = FS.m)
> round(coef(fit.m), 3)

```

(Intercept)	X1	X2
0.0	-0.4	0.5

- this works for all recursive models, but a separate analysis is needed for each endogenous latent variable
- but only for the unstandardized solution; the 'standardized' solution is still biased, because we have no (consistent) estimate for the total variance of the outcome variable (recall: $\beta^z = \frac{s_x}{s_y} \beta$)

3.4 Correlation-preserving factor scores

- what if we use correlation-preserving factor scores (instead of regression or Bartlett factor scores)?
- the regression for 'Y':

```
> fs.cpreg <- lavPredict(fit.sem2, method = "regression", transform = TRUE)
> FS.cpreg <- as.data.frame(fs.cpreg)
> fit.y <- lm(Y ~ X3 + M + fAge, data = FS.cpreg)
> round(coef(fit.y), 3)
```

(Intercept)	X3	M	fAge
0.0	0.1	0.2	-0.3

- the regression for 'M':

```
> fit.m <- lm(M ~ X1 + X2, data = FS.cpreg)
> round(coef(fit.m), 3)
```

(Intercept)	X1	X2
0.0	-0.4	0.5

- it works equally well with the 'Bartlett' version

- that we obtain the correct results is of course due to the fact that for correlation-preserving factor scores, we have that $\text{Var}(\mathbf{f}) = \text{Var}(\boldsymbol{\eta})$
- it is remarkable that these correlation-preserving factor scores are almost never used in the applied literature
- in fact, it turns out that correlation-preserving factor scores work really well, as shown in an extensive simulation study reported in

Bogaert, J., Loh, W. W., & Rosseel, Y. (2026). Consistent Factor Score Regression: A Better Alternative for Uncorrected Factor Score Regression? *Educational and Psychological Measurement*
<https://doi.org/10.1177/00131644251399588>

- the authors termed this approach ‘consistent FSR’ (cFSR)
- they are a good alternative for other (consistent) approaches that we will discuss in the next section

3.5 Sum scores

- sum scores are a special type of factor scores
- they (implicitly) assume that:
 1. all factor loadings are the same (say, equal to 1.0)
 2. all residual variances of the indicators are equal to each other
- if this is approximately true, they can easily replace factor scores
- as an additional benefit, no CFA model needs to be fitted in order to obtain the sum scores
- there is a renewed interest (and respect) in the recent SEM literature
- in the example below, we will use mean scores (instead of sum scores), so that the scale of the regression coefficients is more in line with the population values

```

> Ymean <- rowSums(Data[, c("y1", "y2", "y3", "y4")])/4
> Mmean <- rowSums(Data[, c("m1", "m2", "m3", "m4")])/4
> X1mean <- rowSums(Data[, c("x1", "x2", "x3", "x4")])/4
> X2mean <- rowSums(Data[, c("x5", "x6", "x7", "x8")])/4
> X3mean <- rowSums(Data[, c("x9", "x10", "x11", "x12")])/4
> Age <- Data$Age

```

- regression for 'Y':

```

> fit.y <- lm(Ymean ~ X3mean + Mmean + Age)
> round(coef(fit.y), 3)

```

(Intercept)	X3mean	Mmean	Age
0.000	0.088	0.182	-0.254

- regression for 'M':

```

> fit.m <- lm(Mmean ~ X1mean + X2mean)
> round(coef(fit.m), 3)

```

(Intercept)	X1mean	X2mean
0.000	-0.192	0.265

sum scores (for Y) via CFA

```

> model.ysum <- '
  Y =~ 1*y1 + 1*y2 + 1*y3 + 1*y4
  # equal residual variances per factor
  ,
  y1 ~~ vy*y1; y2 ~~ vy*y2; y3 ~~ vy*y3; y4 ~~ vy*y4
'
> fit.ysum <- sem(model.ysum, data = Data, estimator = "ML")
> ysum.reg <- lavPredict(fit.ysum)[,1]
> cor(ysum.reg, Ymean)

[1] 1

> ysum.bar <- lavPredict(fit.ysum, method = "Bartlett")[,1]
> cor(ysum.bar, Ymean)

[1] 1

> head(cbind(ysum.reg, Ymean)) # regression: different scale
      ysum.reg      Ymean
[1,] 0.001068938 0.001417463
[2,] 0.134193914 0.177947526
[3,] 0.086700609 0.114969139
[4,] 0.656095652 0.870014102
[5,] -0.521661208 -0.691747622
[6,] -1.054545684 -1.398377834

```

```
> head(cbind(ysum.bar, Ymean)) # Bartlett: same scale
```

```
      ysum.bar      Ymean
[1,] 0.001417463 0.001417463
[2,] 0.177947526 0.177947526
[3,] 0.114969139 0.114969139
[4,] 0.870014102 0.870014102
[5,] -0.691747622 -0.691747622
[6,] -1.398377834 -1.398377834
```

Reference:

Sijtsma K., Ellis J.L., Borsboom D. (2024). Recognize the Value of the Sum Score, Psychometrics' Greatest Accomplishment. *Psychometrika*, 89(1), 84–117

<https://doi.org/10.1007/s11336-024-09964-7>

3.6 Two-step standard errors

- so far, we only discussed point estimation
- some procedures are consistent, some are not
- but what about standard errors?
- the ‘naive’ standard errors that we obtain in the regression step in FSR are (usually) not correct
- they fail to take the uncertainty of the first step into account
- we can always use the bootstrap to obtain ‘correct’ standard errors for the final regression coefficients (but this can be time consuming)
- (two-step corrected) analytic standard errors for UFSR and cFSR are available in the `sam()` function with `se = "local"`

4 Computing summary statistics for latent variables

4.1 Introduction

- if our goal is (only) to compute the regression coefficients of the structural relations, then we do not need the individual factor scores
- we only need the variance–covariance matrix $\text{Var}(\mathbf{f})$ (and perhaps the mean vector)
- let $\mathbf{S} = \text{Var}(\mathbf{y})$ be the sample covariance matrix of the observed data
- let \mathbf{A} be the factor score matrix
- it is easy to show that:

$$\text{Var}(\mathbf{f}) = \text{Var}(\mathbf{A} \mathbf{y}) = \mathbf{A} \text{Var}(\mathbf{y}) \mathbf{A}^T = \mathbf{A} \mathbf{S} \mathbf{A}^T$$

- with the exception of correlation-preserving factor scores, the main reason why uncorrected FSR is not consistent is because $\text{Var}(\mathbf{f}) \neq \text{Var}(\boldsymbol{\eta})$

from $\text{Var}(\mathbf{f})$ to $\text{Var}(\boldsymbol{\eta})$

- several authors have suggested to ‘correct’ $\text{Var}(\mathbf{f})$ so that it coincides with $\text{Var}(\boldsymbol{\eta})$
 - Dijkstra (1981), Dijkstra & Henseler (2015): consistent PLS
 - Fuller (1987): measurement error models
 - Wall & Amemiya (2000, 2003): 2-stage method of moments (2SMM) for polynomial SEM
 - Croon (2002): ‘(Croon’s) corrections’ for latent variable models
 - Rosseel & Loh (2024): local SAM
- instead of focusing on individual factor scores, these methods try to compute a consistent estimate for $\text{Var}(\boldsymbol{\eta})$ (and perhaps $E(\boldsymbol{\eta})$ if needed)
- once we have these ‘summary statistics’, we can proceed in a second step with a regression (or path analysis) and obtain consistent results for the structural relations (just like SEM)
- in addition, many of them provide ‘two-step corrected’ standard errors

4.2 The method of Croon (2002)

- Croon (2001) considered a very general framework, including discrete latent variables, and discrete observed indicators
- here, we only consider his approach for the continuous case (continuous latent variables, continuous observed indicators)
- Croon's corrections were originally expressed in scalar form; here, we will present them in a more general form (although for simplicity, we will ignore the meanstructure)
- we assume that we fitted all the measurement blocks, either one by one, in subsets, or all together in one joint CFA
- therefore, we have estimates for Λ and Θ
- we also compute the factor score matrix \mathbf{A} (either the regression or the Bartlett version)
- furthermore, we define the matrices $\mathbf{D} = \mathbf{A}\Lambda$ and $\mathbf{E} = \mathbf{A}\Theta\mathbf{A}^T$

- Croon's corrections can then be expressed as follows:

$$\begin{aligned}
 \text{Var}(\boldsymbol{\eta}) &= \mathbf{D}^{-1} [\text{Var}(\mathbf{f}) - \mathbf{E}] \mathbf{D}^{-T} \\
 &= \mathbf{D}^{-1} [\mathbf{A} \mathbf{S} \mathbf{A}^T - \mathbf{E}] \mathbf{D}^{-T} \\
 &= \mathbf{D}^{-1} [\mathbf{A} \mathbf{S} \mathbf{A}^T - \mathbf{A} \boldsymbol{\Theta} \mathbf{A}^T] \mathbf{D}^{-T} \\
 &= \mathbf{D}^{-1} \mathbf{A} [\mathbf{S} - \boldsymbol{\Theta}] \mathbf{A}^T \mathbf{D}^{-T} \\
 &= (\mathbf{A}\boldsymbol{\Lambda})^{-1} \mathbf{A} [\mathbf{S} - \boldsymbol{\Theta}] \mathbf{A}^T (\mathbf{A}\boldsymbol{\Lambda})^{-T}
 \end{aligned}$$

- if we use Bartlett factor scores, we have that $\mathbf{D} = \mathbf{A}\boldsymbol{\Lambda} = \mathbf{I}$ and the last expression simplifies to

$$\text{Var}(\boldsymbol{\eta}) = \mathbf{A}_{bar} [\mathbf{S} - \boldsymbol{\Theta}] \mathbf{A}_{bar}^T$$

- several papers showed that the 'Method of Croon' worked remarkably well, and sometimes even better than SEM
- we will not provide an example here, because the 'Method of Croon' turns out to be a special case of local SAM

some references on the method of Croon

Devlieger, I., Mayer, A., & Rosseel, Y. (2016). Hypothesis testing using factor score regression: A comparison of four methods. *Educational and Psychological Measurement, 76*, 741–770.

Devlieger, I., & Rosseel, Y. (2017). Factor Score Path Analysis. *Methodology, 13*, 31–38.

Takane, Y., & Hwang, H. (2017). Comparisons among several consistent estimators of structural equation models. *Behaviormetrika, 45*, 157–188.

Kelcey, B. (2019). A robust alternative estimator for small to moderate sample SEM: Bias-corrected factor score path analysis. *Addictive Behaviors, 94*, 83–98.

Bogaert, J., Loh, W.W., & Rosseel, Y. (2023). A small sample correction for factor score regression. *Educational and Psychological Measurement, 83*(3), 495–519.

4.3 Local SAM

- the measurement model:

$$\mathbf{y} = \boldsymbol{\nu} + \mathbf{\Lambda}\boldsymbol{\eta} + \boldsymbol{\epsilon}$$

- to solve this for $\boldsymbol{\eta}$, we proceed as follows:

$$\boldsymbol{\nu} + \mathbf{\Lambda}\boldsymbol{\eta} + \boldsymbol{\epsilon} = \mathbf{y}$$

$$\mathbf{\Lambda}\boldsymbol{\eta} = \mathbf{y} - \boldsymbol{\nu} - \boldsymbol{\epsilon}$$

$$\mathbf{M}\mathbf{\Lambda}\boldsymbol{\eta} = \mathbf{M}[\mathbf{y} - \boldsymbol{\nu} - \boldsymbol{\epsilon}]$$

$$\boldsymbol{\eta} = \mathbf{M}[\mathbf{y} - \boldsymbol{\nu} - \boldsymbol{\epsilon}]$$

where \mathbf{M} is $M \times P$ mapping matrix such that $\mathbf{M}\mathbf{\Lambda} = \mathbf{I}_M$

- we assume $\mathbf{E}(\boldsymbol{\epsilon}) = \mathbf{0}$ and write $\text{Var}(\boldsymbol{\epsilon}) = \boldsymbol{\Theta}$; it follows that

$$\mathbf{E}(\boldsymbol{\eta}) = \mathbf{M}[\mathbf{E}(\mathbf{y}) - \boldsymbol{\nu}]$$

$$\text{Var}(\boldsymbol{\eta}) = \mathbf{M}[\text{Var}(\mathbf{y}) - \boldsymbol{\Theta}]\mathbf{M}^T$$

local SAM: first stage

- first stage: estimation of the measurement part of the model (only)
- M is the number of latent variables; B is the number of measurement ‘blocks’
- three options:
 1. $B = 1$: single CFA
 2. $B = M$: as many ‘blocks’ as we have latent variables
 3. $B < M$: if some blocks are ‘linked’ together
- we recommend $B = M$ whenever possible (except if the reliabilities are very low, and the number of indicators is small)
- measurement models that are ‘linked’ (due to cross-loadings, correlated residuals, or equality constraints) should be fitted together, leading to $B < M$
- for each block, we can use ML, GLS, ..., or we can use noniterative estimators

local SAM: creating the mapping matrix \mathbf{M}

- recall, the mapping matrix must be chosen such that $\mathbf{M}\mathbf{\Lambda} = \mathbf{I}_M$
- three possible solutions for the mapping matrix \mathbf{M} :

$$\mathbf{M} = (\mathbf{\Lambda}^T \mathbf{\Theta}^{-1} \mathbf{\Lambda})^{-1} \mathbf{\Lambda}^T \mathbf{\Theta}^{-1} \quad (ML)$$

$$\mathbf{M} = (\mathbf{\Lambda}^T \mathbf{S}^{-1} \mathbf{\Lambda})^{-1} \mathbf{\Lambda}^T \mathbf{S}^{-1} \quad (GLS)$$

$$\mathbf{M} = (\mathbf{\Lambda}^T \mathbf{\Lambda})^{-1} \mathbf{\Lambda}^T \quad (ULS)$$

- we then estimate $\mathbb{E}(\boldsymbol{\eta})$ and $\text{Var}(\boldsymbol{\eta})$ as follows:

$$\widehat{\mathbb{E}(\boldsymbol{\eta})} = \hat{\mathbf{M}} [\bar{\mathbf{y}} - \hat{\boldsymbol{\nu}}]$$
$$\widehat{\text{Var}(\boldsymbol{\eta})} = \hat{\mathbf{M}} [\mathbf{S} - \hat{\mathbf{\Theta}}] \hat{\mathbf{M}}^T$$

- \mathbf{M} is called a mapping matrix because it ‘maps’ the (centered and error-corrected) observed data to the latent space

local SAM: second stage

- second stage: $\widehat{E}(\boldsymbol{\eta})$ and $\widehat{\text{Var}}(\boldsymbol{\eta})$ are used to estimate the parameters in the structural part of the model
- this can be done using ‘path analysis’, where we treat everything as observed, and the data are presented via summary statistics
- we can use ML, GLS, ...
- or we can use noniterative estimators: OLS (if the model is recursive) or TSLS (if the model is not recursive)

local SAM: further comments

- two-step corrected standard errors are available (see Appendix C in the SAM paper); recently, we added ‘local’ standard errors
- local fit measures only (for each measurement block, for the structural part)
- the (co)variance matrix of the latent variables is always positive definite
- we can handle missing data (fiml or two-stage), categorical indicators, two-level data (random intercepts only), latent interactions, higher-order measurement models, ...
- in a few special settings, a mini-iterative procedure is needed to obtain $\text{Var}(\boldsymbol{\eta})$:
 - the factor loading matrix ($\boldsymbol{\Lambda}$) is rank deficient (e.g., more factors than observed variables)
 - models (e.g., variance components) where zeroes in the variance-covariance matrix of the latent variables are needed in order to identify the model

example: fit measurement blocks, using $B = M$

```
> fit.Y <- sem('Y =~ y1 + y2 + y3 + y4', data = Data)
> fit.M <- sem('M =~ m1 + m2 + m3 + m4', data = Data)
> fit.X1 <- sem('X1 =~ x1 + x2 + x3 + x4', data = Data)
> fit.X2 <- sem('X2 =~ x5 + x6 + x7 + x8', data = Data)
> fit.X3 <- sem('X3 =~ x9 + x10 + x11 + x12', data = Data)

> # assemble Lambda and Theta
> Lambda <- matrix(0, 20, 5)
> Lambda[ 1:4, 1] <- lavInspect(fit.Y, "est")$lambda
> Lambda[ 5:8, 2] <- lavInspect(fit.M, "est")$lambda
> Lambda[ 9:12, 3] <- lavInspect(fit.X1, "est")$lambda
> Lambda[13:16, 4] <- lavInspect(fit.X2, "est")$lambda
> Lambda[17:20, 5] <- lavInspect(fit.X3, "est")$lambda

> Theta <- lav_matrix_bdiag(lavInspect(fit.Y, "est")$theta,
                           lavInspect(fit.M, "est")$theta,
                           lavInspect(fit.X1, "est")$theta,
                           lavInspect(fit.X2, "est")$theta,
                           lavInspect(fit.X3, "est")$theta)
```

example: compute ML version of the mapping matrix M

```

> Theta.inv <- solve(Theta)
> M <- solve(t(Lambda) %**% Theta.inv %**% Lambda) %**% t(Lambda) %**% Theta.inv

> # add age
> M      <- lav_matrix_bdiag(M,      matrix(1, nrow = 1L, ncol = 1L))
> Theta <- lav_matrix_bdiag(Theta, matrix(0, nrow = 1L, ncol = 1L))
> rownames(M) <- c("Y", "M", "X1", "X2", "X3", "Age")

> # compute (biased) sample covariance matrix 'S'
> N <- nrow(Data)
> S <- cov(Data) * (N - 1L)/N

> # compute Var(Eta)
> Var.eta <- M %**% (S - Theta) %**% t(M)
> round(Var.eta, 3)

```

	Y	M	X1	X2	X3	Age
Y	1.136	0.245	-0.24	-0.102	0.096	-0.258
M	0.245	1.250	-0.20	0.340	0.280	0.110
X1	-0.240	-0.200	1.00	0.400	-0.200	0.600
X2	-0.102	0.340	0.40	1.000	0.400	0.700
X3	0.096	0.280	-0.20	0.400	1.000	0.200
Age	-0.258	0.110	0.60	0.700	0.200	1.000

example: second stage – using OLS

```
> # compute regression coefficients for Y
> beta.Y <- ( solve(Var.eta[c("X3", "M", "Age"), c("X3", "M", "Age")]) %*%
              Var.eta[c("X3", "M", "Age"), "Y", drop = FALSE] )
> round(beta.Y, 3)
```

```
      Y
X3  0.1
M   0.2
Age -0.3
```

```
> # compute regression coefficients for M
> beta.M <- ( solve(Var.eta[c("X1", "X2"), c("X1", "X2")]) %*%
              Var.eta[c("X1", "X2"), "M", drop = FALSE] )
> round(beta.M, 3)
```

```
      M
X1 -0.4
X2  0.5
```

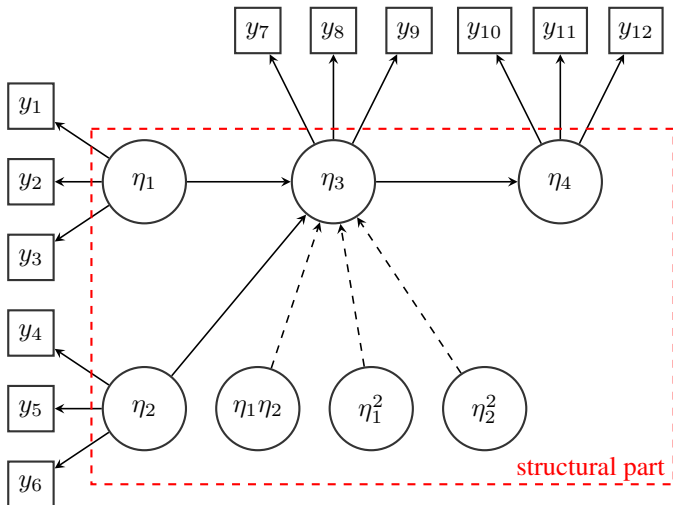
example: using the sam() function

```
> fit.lsam <- sam(model = model, data = Data)
> parameterEstimates(fit.lsam, ci = FALSE, output = "text")[1:5,]
```

Regressions:

	Estimate	Std.Err	z-value	P(> z)
Y ~				
X3	0.100	0.101	0.994	0.320
M	0.200	0.094	2.126	0.034
Age	-0.300	0.087	-3.468	0.001
M ~				
X1	-0.400	0.126	-3.164	0.002
X2	0.500	0.132	3.793	0.000

4.4 Application 1: adding latent quadratic and interaction terms



- in the joint setting, adding latent quadratic/interaction terms is not trivial
- two popular methods are the product-indicator (PI) approach, and the so-called ‘Latent Moderated Structural Equations’ (LMS) approach
- none of these scale well: they cannot handle many quadratic and latent interaction terms simultaneously
- but if you can decouple the measurement and structural part, this becomes feasible
- a very general SAM solution (allowing for polynomial relations between latent variables) was already described in Wall & Amemiya (2000)
- the local SAM approach: find an explicit expression for

$$E(\boldsymbol{\eta} \otimes \boldsymbol{\eta}) \quad \text{and} \quad \text{Var}(\boldsymbol{\eta} \otimes \boldsymbol{\eta})$$

where \otimes denotes the tensor (or Kronecker) product

local SAM and interaction/quadratic terms

- creating the 'augmented' vector $\boldsymbol{\eta} \otimes \boldsymbol{\eta}$:
- several elements are duplicated
- based on the model, we select what we need

$$\boldsymbol{\eta} = \begin{pmatrix} 1 \\ \eta_1 \\ \eta_2 \\ \eta_3 \end{pmatrix}$$

$$\boldsymbol{\eta} \otimes \boldsymbol{\eta} =$$

$$\begin{pmatrix} 1 \\ \eta_1 \\ \eta_2 \\ \eta_3 \\ \eta_1 1 \\ \eta_1 \eta_1 \\ \eta_1 \eta_2 \\ \eta_1 \eta_3 \\ \eta_2 1 \\ \eta_2 \eta_1 \\ \eta_2 \eta_2 \\ \eta_2 \eta_3 \\ \eta_3 1 \\ \eta_3 \eta_1 \\ \eta_3 \eta_2 \\ \eta_3 \eta_3 \end{pmatrix} = \begin{pmatrix} 1 \\ \eta_1 \\ \eta_2 \\ \eta_3 \\ \eta_1 \\ \eta_1^2 \\ \eta_1 \eta_2 \\ \eta_1 \eta_3 \\ \eta_2 \eta_1 \\ \eta_2 \\ \eta_2^2 \\ \eta_2 \eta_3 \\ \eta_3 \\ \eta_3 \eta_1 \\ \eta_3 \eta_2 \\ \eta_3^2 \end{pmatrix}$$

the augmented (latent) sample statistics

- augmented mean vector:

$$E(\boldsymbol{\eta} \otimes \boldsymbol{\eta}) = \text{vec}[\text{Var}(\boldsymbol{\eta})] + E(\boldsymbol{\eta}) \otimes E(\boldsymbol{\eta})$$

- augmented variance-covariance matrix (simple version, assuming normality for the measurement error):

$$\text{Var}(\boldsymbol{\eta} \otimes \boldsymbol{\eta}) \approx \text{Var}(\mathbf{f} \otimes \mathbf{f}) - \left[\mathbf{Q} + \mathbf{K}_m \mathbf{Q} + \mathbf{Q} \mathbf{K}_m^T + \mathbf{K}_m \mathbf{Q} \mathbf{K}_m^T + \boldsymbol{\Gamma}_{22}^{*(NT)}(\mathbf{r}) \right]$$

where

$$\mathbf{Q} = \text{Var}(\boldsymbol{\eta}) \otimes \text{Var}(\mathbf{r}) + E(\boldsymbol{\eta})E(\boldsymbol{\eta})^T \otimes \text{Var}(\mathbf{r})$$

and

$$\boldsymbol{\Gamma}_{22}^{*(NT)}(\mathbf{r}) = (\mathbf{I}_{m^2} + \mathbf{K}_m) (\text{Var}(\mathbf{r}) \otimes \text{Var}(\mathbf{r}))$$

- \mathbf{K}_m is the commutation matrix
- $\boldsymbol{\Gamma}_{22}^{*(NT)}(\mathbf{r})$ is the ‘Gamma’ matrix of the measurement error (\mathbf{r})

implementation in lavaan

```
> model <- '  
  # measurement part  
  f1 =~ y1 + y2 + y3  
  f2 =~ y4 + y5 + y6  
  f3 =~ y7 + y8 + y9  
  
  # structural part  
  f3 ~ f1 + f2 + f1:f1 + f2:f2 + f1:f2  
,  
> fit <- sam(model, data = Data)
```

- paper:

Rosseel, Y., Burghgraeve, E., Loh, W.W., Schermelleh-Engel, K. (2025). Structural after Measurement (SAM) approaches for accommodating latent quadratic and interaction effects. *Behavior Research Methods*, 57(101). <https://doi.org/10.3758/s13428-024-02532-y>

4.5 Application 2: noniterative SEM

- for CFA, many noniterative estimators are available; some (i.e., the multiple group method) perform better than ML in terms of mean squared error

Dhaene, S. & Rosseel, Y. (2023). An Evaluation of Non-Iterative Estimators in Confirmatory Factor Analysis. *Structural Equation Modeling: A Multidisciplinary Journal*, 31(1), 1–13.

<https://doi.org/10.1080/10705511.2023.2187285>

- we can use these noniterative estimators for the measurement part in SAM

Dhaene, S., & Rosseel, Y. (2023). An Evaluation of Non-Iterative Estimators in the Structural after Measurement (SAM) Approach to Structural Equation Modeling (SEM). *Structural Equation Modeling: A Multidisciplinary Journal*, 30(6), 926–940

- “[the] local SAM approach outperforms traditional SEM in small to moderate samples (both in terms of convergence and MSE values), especially when reliability drops. ”

4.6 Application 3: comparing structural relations across many groups

- reference (open access):

Perez Alonso, A.F., Rosseel, Y., Vermunt, J.K., & De Roover, K. (2024). Mixture Multigroup Structural Equation Modeling: A Novel Method for Comparing Structural Relations Across Many Groups. *Psychological Methods*. <https://doi.org/10.1037/met0000667>

- relationships between latent variables are often different across groups (e.g., countries); but some groups may be similar in the sense that they have similar values for the regression coefficients; we like to ‘discover’ these hidden clusters of similar groups
- in a first step, we estimated the measurement part across all groups (fixing the factor loadings to be the same across groups); this resulted in (model-implied) latent (co)variance matrices for all the groups
- in a second step, a mixture modeling approach is used to find homogeneous clusters that share similar regression coefficients

5 SAM: other approaches

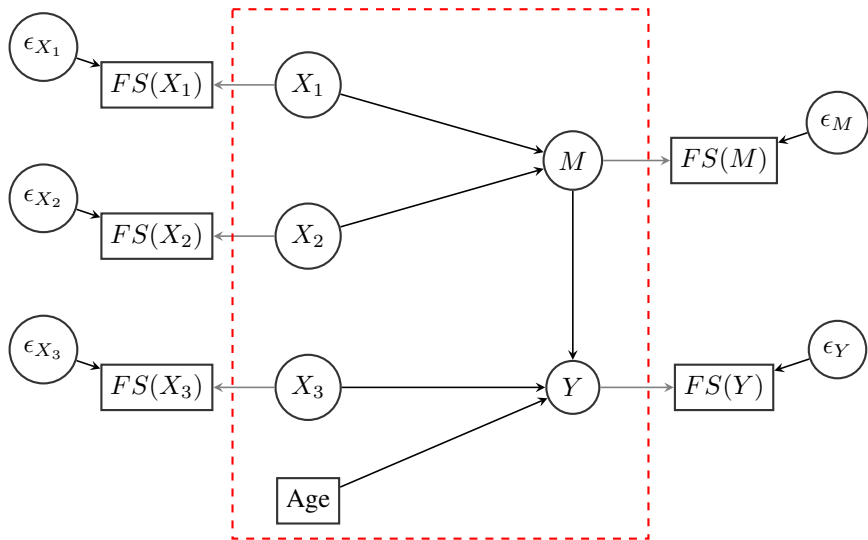
5.1 The single-indicator (SI) approach

- a good reference:

Savalei, V. (2019). A comparison of several approaches for controlling measurement error in small samples. *Psychological methods*, 24(3), 352–370.

- in a first step, we estimate the measurement blocks, typically one by one, and we compute factor scores for each latent variable
- alternatively, we could also directly compute sum (or mean) scores, without fitting a measurement model
- in the second step, we fit the structural part; we keep the latent variables, but we replace their indicators by a single indicator (either the factor scores or the sum scores)

diagram single-indicator approach



setting the reliability of the single indicator

- the (single) factor loading is set to one
- the residual variance of the single indicator is fixed to a constant; let Y_j be the single indicator of the j -th latent variable; we set the residual variance of Y_j as follows:

$$\text{Var}(\epsilon_j) = (1 - \text{REL}_j)\text{Var}(Y_j)$$

- REL_j reflects the ‘reliability’ of Y_j
- there are (at least) three ways to choose the reliability:
 1. we can estimate REL_j based on the measurement model (‘model-based’)
 2. we can provide a fixed guess (say, 0.7 or 0.8)
 3. we can estimate the reliability using, for example, coefficient alpha
- therefore, several variants of this single indicator approach are possible

SI + sum scores + fixed reliability (0.8)

```
> Ymean <- rowSums(Data[, c("y1", "y2", "y3", "y4")])/4
> Mmean <- rowSums(Data[, c("m1", "m2", "m3", "m4")])/4
> X1mean <- rowSums(Data[, c("x1", "x2", "x3", "x4")])/4
> X2mean <- rowSums(Data[, c("x5", "x6", "x7", "x8")])/4
> X3mean <- rowSums(Data[, c("x9", "x10", "x11", "x12")])/4
> CS <- data.frame(Y=Ymean, M=Mmean, X1=X1mean, X2=X2mean,
                  X3=X3mean, Age = Data$Age)

> res.Y <- (1 - 0.8)*var(Ymean)
> res.M <- (1 - 0.8)*var(Mmean)
> res.X1 <- (1 - 0.8)*var(X1mean)
> res.X2 <- (1 - 0.8)*var(X2mean)
> res.X3 <- (1 - 0.8)*var(X3mean)
```

```

> # create model syntax
> model.si <- c("fY =~ 1*Y", paste0("Y ~~ ", res.Y, "*Y"),
               "fM =~ 1*M", paste0("M ~~ ", res.M, "*M"),
               "fX1 =~ 1*X1", paste0("X1 ~~ ", res.X1, "*X1"),
               "fX2 =~ 1*X2", paste0("X2 ~~ ", res.X2, "*X2"),
               "fX3 =~ 1*X3", paste0("X3 ~~ ", res.X3, "*X3"),
               "fAge =~ 1*Age", "Age ~~ 0*Age")
> model.struc <- '
  fY ~ fX3 + fM + fAge
  fM ~ fX1 + fX2
  ,
> fit.si <- sem(c(model.si, model.struc), data = CS)
> parameterEstimates(fit.si, ci = FALSE, output = "text")[13:17,]

```

Regressions:

	Estimate	Std.Err	z-value	P(> z)
fY ~				
fX3	0.104	0.095	1.094	0.274
fM	0.226	0.102	2.216	0.027
fAge	-0.260	0.074	-3.533	0.000
fM ~				
fX1	-0.278	0.086	-3.240	0.001
fX2	0.371	0.089	4.144	0.000

SI + factor scores + model-based reliability

```

> fit.Y <- sem('Y =~ y1 + y2 + y3 + y4', data = Data)
> fit.M <- sem('M =~ m1 + m2 + m3 + m4', data = Data)
> fit.X1 <- sem('X1 =~ x1 + x2 + x3 + x4', data = Data)
> fit.X2 <- sem('X2 =~ x5 + x6 + x7 + x8', data = Data)
> fit.X3 <- sem('X3 =~ x9 + x10 + x11 + x12', data = Data)

> fs.Y <- lavPredict(fit.Y, method = "Bartlett", fsm = TRUE)
> fs.M <- lavPredict(fit.M, method = "Bartlett", fsm = TRUE)
> fs.X1 <- lavPredict(fit.X1, method = "Bartlett", fsm = TRUE)
> fs.X2 <- lavPredict(fit.X2, method = "Bartlett", fsm = TRUE)
> fs.X3 <- lavPredict(fit.X3, method = "Bartlett", fsm = TRUE)
> FS <- data.frame(Y=fs.Y, M=fs.M, X1=fs.X1, X2=fs.X2,
                  X3=fs.X3, Age = Data$Age)

> A <- attr(fs.Y, "fsm")[[1]]; THETA <- lavInspect(fit.Y, "est")$theta
> res.Y <- drop(A %*% THETA %*% t(A))
> A <- attr(fs.M, "fsm")[[1]]; THETA <- lavInspect(fit.M, "est")$theta
> res.M <- drop(A %*% THETA %*% t(A))
> A <- attr(fs.X1, "fsm")[[1]]; THETA <- lavInspect(fit.X1, "est")$theta
> res.X1 <- drop(A %*% THETA %*% t(A))
> A <- attr(fs.X2, "fsm")[[1]]; THETA <- lavInspect(fit.X2, "est")$theta
> res.X2 <- drop(A %*% THETA %*% t(A))
> A <- attr(fs.X3, "fsm")[[1]]; THETA <- lavInspect(fit.X3, "est")$theta
> res.X3 <- drop(A %*% THETA %*% t(A))

```

```

> # create model syntax
> model.si <- c("fY =~ 1*Y", paste0("Y ~~ ", res.Y, "*Y"),
               "fM =~ 1*M", paste0("M ~~ ", res.M, "*M"),
               "fX1 =~ 1*X1", paste0("X1 ~~ ", res.X1, "*X1"),
               "fX2 =~ 1*X2", paste0("X2 ~~ ", res.X2, "*X2"),
               "fX3 =~ 1*X3", paste0("X3 ~~ ", res.X3, "*X3"),
               "fAge =~ 1*Age", "Age ~~ 0*Age")
> model.struc <- '
  fY ~ fX3 + fM + fAge
  fM ~ fX1 + fX2
  ,
> fit.si <- sem(c(model.si, model.struc), data = FS)
> parameterEstimates(fit.si, ci = FALSE, output = "text")[13:17,]

```

Regressions:

	Estimate	Std.Err	z-value	P(> z)
fY ~				
fX3	0.100	0.100	1.000	0.317
fM	0.200	0.091	2.209	0.027
fAge	-0.300	0.084	-3.589	0.000
fM ~				
fX1	-0.400	0.118	-3.388	0.001
fX2	0.500	0.118	4.228	0.000

- note: when using Bartlett factor scores, the diagonal elements of $\mathbf{E} = \mathbf{A} \Theta \mathbf{A}^T$ contain the needed residual variances

5.2 Global SAM

- global SAM is very similar to regular (system-wide) SEM
- in a first step, we estimate the parameters of the measurement blocks (either one by one, or in a single CFA)
- in a second step, we fit the full SEM, but we fix the values of the measurement parameters to the values we obtained in the first step; they are no longer free parameters
- this is the approach that was advocated in the early literature (e.g., Burt 1976), and many people associate ‘SAM’ (or ‘two-step’) with this approach
- because the second step ‘sees’ the full model (measurement and structural), Rosseel & Loh (2024) called this ‘global’ SAM
- two-step corrected standard errors are available
- when the model is correctly specified, global SAM usually provides identical results as local SAM

global SAM using the sam() function

```

> model <- '
  # measurement part
  Y =~ y1 + y2 + y3 + y4
  M =~ m1 + m2 + m3 + m4
  X1 =~ x1 + x2 + x3 + x4
  X2 =~ x5 + x6 + x7 + x8
  X3 =~ x9 + x10 + x11 + x12
  fAge =~ 1*Age; Age ~~ 0*Age

  # structural part
  Y ~ X3 + M + fAge
  M ~ X1 + X2
,
> fit.gsam <- sam(model, data = Data, sam.method = "global")
> parameterEstimates(fit.gsam, ci = FALSE, output = "text")[1:5,]

```

Regressions:

	Estimate	Std.Err	z-value	P(> z)
Y ~				
X3	0.100	0.101	0.994	0.320
M	0.200	0.094	2.126	0.034
fAge	-0.300	0.087	-3.468	0.001
M ~				
X1	-0.400	0.126	-3.164	0.002
X2	0.500	0.132	3.793	0.000

6 Consistency versus MSE

6.1 Consistency

- (informally:) an estimator is said to be consistent if the estimates (produced by the estimator) “converge” to the true value of the parameter being estimated, as the sample size increases (to infinity)
- in statistics, consistency is considered to be a fundamental property of estimators; if an estimator is not consistent, it is often disregarded immediately
- which SAM approaches are consistent:
 - FSR with Skrondal & Laake (2001) factor scores
 - FSR with correlation-preserving factor scores
 - the method of Croon, local SAM
 - single-indicator + Bartlett factor scores + model-based reliability
 - global SAM
- which SAM approaches are NOT consistent:

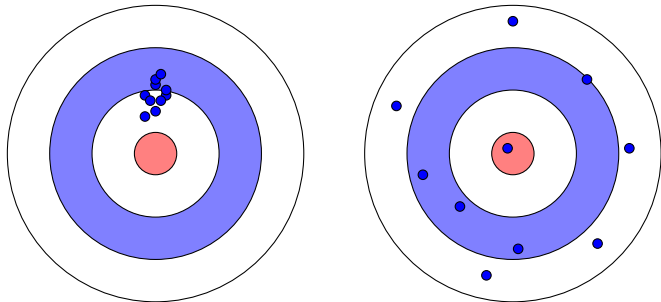
- uncorrected FSR (with Bartlett or regression factor scores)
 - sum (or mean) scores
 - single-indicator + fixed/alpha reliability
- (ML/GLS/WLS as used in SEM are also consistent estimators)
- consistency is not the same as unbiasedness
- (informally:) an estimator is unbiased if the *average* estimate across a very large (infinite) number of samples approaches the true value
- example: the sample variance $\sum_i (X_i - \bar{X})^2 / N$ divided by N (instead of $N - 1$) is biased but consistent
- example: if you always take the first observation (X_1) as the estimate of a mean, then this is unbiased, but not consistent
- in a simulation study, we usually study the bias of an estimator (given a specific sample size)

6.2 Bias, variability and mean squared error (MSE)

- unbiasedness is nice, but there is a price to be paid: unbiased estimators often exhibit increased variability
- on the other hand, a biased estimator may exhibit less variability (in comparison)
- this is known as the bias–variance trade–off
- one way to capture both bias and variance in a single quantity is to use the mean squared error (MSE)
- $MSE = \text{bias}^2 + \text{variance}$
- in statistics, when comparing estimators, we often prefer estimators that have the lowest MSE
- in the SEM literature, MSE has been mostly neglected (in favor of unbiasedness)

low bias versus low MSE

- suppose you are playing darts; which player would you like to be?



- left panel: biased, but low variability
- right panel: unbiased, but high variability

MSE for SAM approaches

- ‘consistent’ approaches typically have the lowest MSE . . . if the sample size is large enough
- however, for small to medium sample sizes, it may very well be that inconsistent approaches (for example: uncorrected FSR) have a lower MSE
- the next two figures are taken from:

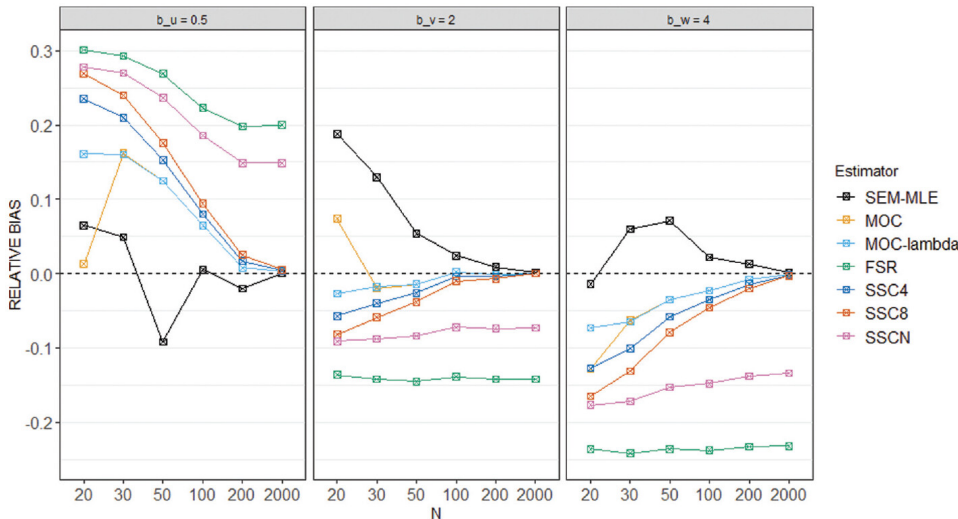
Bogaert, J., Loh, W.W., & Rosseel, Y. (2023). A small sample correction for factor score regression. *Educational and Psychological Measurement*, 83(3), 495–519.

- similar results regarding single-indicator approaches:

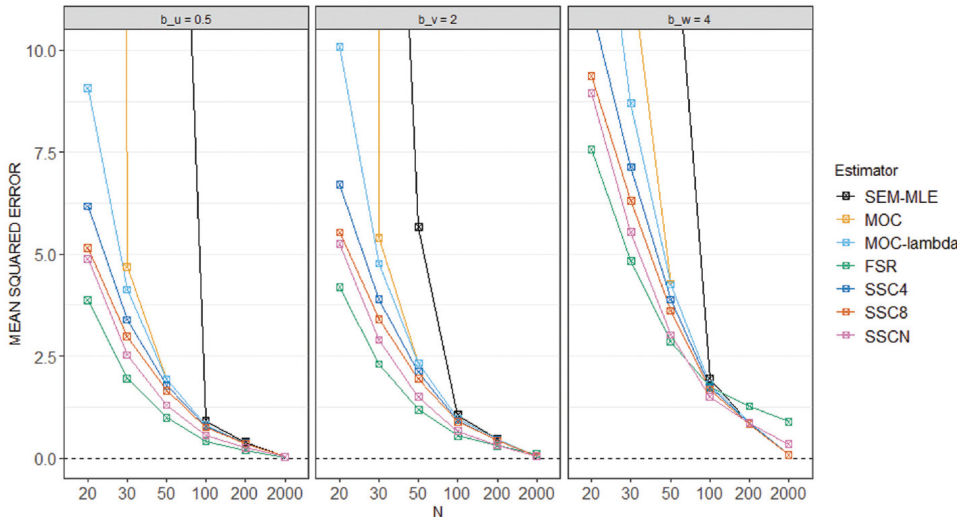
Savalei, V. (2019). A comparison of several approaches for controlling measurement error in small samples. *Psychological methods*, 24(3), 352–370.

- in the Savalei paper, SI + sum scores + fixed reliability (0.8) had the lowest MSE when the sample size was small (N=30 to N=200)

relative bias (figure 3)



mean squared error (figure 7)



the alpha correction

- local SAM is consistent, because it uses

$$\widehat{\text{Var}}(\boldsymbol{\eta}) = \text{Var}(\mathbf{f}) - \widehat{\text{Var}}(\mathbf{r})$$

where $\widehat{\text{Var}}(\mathbf{r})$ is the correction (for measurement error) term

- uncorrected FSR (with Bartlett factor scores) is not consistent, because it uses

$$\widehat{\text{Var}}(\boldsymbol{\eta}) = \text{Var}(\mathbf{f})$$

- Bogaert, Loh, & Rosseel (2023) discuss the so-called ‘alpha’ correction

$$\widehat{\text{Var}}(\boldsymbol{\eta}) = \text{Var}(\mathbf{f}) - \alpha^* \widehat{\text{Var}}(\mathbf{r})$$

where α^* is (here) a value between 0 and 1

- if $\alpha^* = 1$, we have local SAM; if $\alpha^* = 0$, we have uncorrected FSR
- future research: for a given model, a given sample size: find the sweet spot: the value for α^* that results (on average) in the lowest MSE

coverage probability

- low MSE is good, but . . .
- for valid inference, we also need good ‘coverage’
- the coverage is the probability that a confidence interval will include the true value of the parameter of interest
- for example, set the nominal coverage probability at 0.95; for a large number of replications, the proportion of replications that produce a confidence interval (for the parameter of interest) that includes the true value should approximate 0.95
- ideally, we have it all: almost no bias, not much variability, and good coverage
- so which SAM method is best? (it will depend on many factors, including sample size)
- as always, more research is needed

7 SAM: software implementation

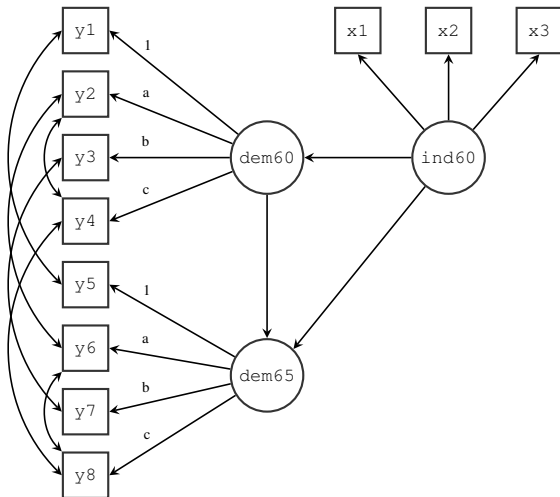
- some of the SAM approaches have been implemented in the `sam()` function in the R package `lavaan`:

- `sam.method = "local"` (default)
- `sam.method = "global"`
- `sam.method = "fsr"` (using Bartlett factor scores)
- `sam.method = "cfsr"` (correlation-preserving factor scores)

- typical call:

```
> fit.sam <- sam(model, data = Data, sam.method = "local",
  local.options = list(M.method = "ML",
    alpha.correction = 0L), # range: 0 -- N-1
  # optionally link measurement blocks
  mm.list = list(block1 = c("Y", "M"),
    block2 = c("X1", "X2", "X3")),
  # measurement options
  mm.args = list(estimator = "ML"),
  # structural options
  struc.args = list(estimator = "GLS"),
  # global options
  meanstructure = FALSE)
```

example: the Political Democracy model



R code (with a misspecification in the structural part)

```
> model <- '  
  # latent variable definitions  
  ind60 =~ x1 + x2 + x3  
  dem60 =~ y1 + a*y2 + b*y3 + c*y4  
  dem65 =~ y5 + a*y6 + b*y7 + c*y8  
  
  # regressions  
  dem60 ~ ind60  
  dem65 ~ ind60 + 0*dem60 # misspecified: fixed-to-zero  
  
  # residual correlations  
  y1 ~~ y5  
  y2 ~~ y4 + y6  
  y3 ~~ y7  
  y4 ~~ y8  
  y6 ~~ y8  
,  
  
> fit.sam <- sam(model, data = PoliticalDemocracy,  
  # link measurement blocks  
  mm.list = list(ind = "ind60", dem = c("dem60", "dem65")),  
  struc.args = list(estimator = "GLS"), # just for fun  
  se = "local") # will be the default soon
```

standard sam() output: only structural part

```
> # standard sam output
> summary(fit.sam)
```

This is lavaan 0.6-22.2560 -- using the SAM approach to SEM

SAM method	LOCAL
Mapping matrix M method	ML
Number of measurement blocks	2
Estimator measurement part	ML
Estimator structural part	GLS
Number of observations	75

Summary Information Measurement + Structural:

Block	Latent	Nind	Chisq	Df
1	ind60	3	0.00	0
2	dem60, dem65	8	15.32	16

Model-based reliability latent variables:

ind60	dem60	dem65
0.966	0.868	0.87

Summary Information Structural part:

```

  chisq df   cfi rmsea  srmr
35.817  1 0.287 0.686 0.505

```

Parameter Estimates:

```

Standard errors
Information
Information saturated (h1) model          Local
                                          Expected
                                          Structured

```

Regressions:

	Estimate	Std.Err	z-value	P(> z)
dem60 ~				
ind60	1.454	0.370	3.932	0.000
dem65 ~				
ind60	1.824	0.363	5.027	0.000
dem60	0.000			

Variances:

	Estimate	Std.Err	z-value	P(> z)
ind60	0.446	0.075	5.975	0.000
.dem60	0.120	0.306	0.394	0.694
.dem65	0.097	0.225	0.432	0.666

sam() output, including measurement part

```
> # just for illustration, we also show the estimated parameters
> # of the measurement blocks
> #
> summary(fit.sam, remove.step1 = FALSE)
```

This is lavaan 0.6-22.2560 -- using the SAM approach to SEM

SAM method	LOCAL
Mapping matrix M method	ML
Number of measurement blocks	2
Estimator measurement part	ML
Estimator structural part	GLS
Number of observations	75

Summary Information Measurement + Structural:

Block	Latent	Nind	Chisq	Df
1	ind60	3	0.00	0
2	dem60, dem65	8	15.32	16

Model-based reliability latent variables:

ind60	dem60	dem65
0.966	0.868	0.87

Summary Information Structural part:

```

chisq df   cfi rmsea  srmr
35.817  1 0.287 0.686 0.505

```

Parameter Estimates:

```

Standard errors
Information
Information saturated (h1) model
Local
Expected
Structured

```

Latent Variables:

		Step	Estimate	Std.Err	z-value	P(> z)
ind60 =~						
	x1	1	1.000			
	x2	1	2.193	0.142	15.403	0.000
	x3	1	1.824	0.153	11.883	0.000
dem60 =~						
	y1	1	1.000			
	y2	(a)	1.213	0.143	8.483	0.000
	y3	(b)	1.210	0.125	9.690	0.000
	y4	(c)	1.273	0.122	10.453	0.000
dem65 =~						
	y5	1	1.000			
	y6	(a)	1.213	0.143	8.483	0.000
	y7	(b)	1.210	0.125	9.690	0.000
	y8	(c)	1.273	0.122	10.453	0.000

Regressions:

	Step	Estimate	Std.Err	z-value	P(> z)
dem60 ~					
ind60	2	1.454	0.370	3.932	0.000
dem65 ~					
ind60	2	1.824	0.363	5.027	0.000
dem60	2	0.000			

Covariances:

	Step	Estimate	Std.Err	z-value	P(> z)
.y1 ~~					
.y5	1	0.577	0.364	1.585	0.113
.y2 ~~					
.y4	1	1.390	0.685	2.030	0.042
.y6	1	2.068	0.733	2.822	0.005
.y3 ~~					
.y7	1	0.727	0.611	1.190	0.234
.y4 ~~					
.y8	1	0.476	0.453	1.049	0.294
.y6 ~~					
.y8	1	1.257	0.583	2.156	0.031

Variances:

	Step	Estimate	Std.Err	z-value	P(> z)
.x1	1	0.084	0.020	4.140	0.000
.x2	1	0.108	0.074	1.455	0.146
.x3	1	0.468	0.091	5.124	0.000
.y1	1	1.879	0.431	4.355	0.000

.y2	1	7.530	1.363	5.523	0.000
.y3	1	4.966	0.966	5.141	0.000
.y4	1	3.214	0.722	4.449	0.000
.y5	1	2.499	0.518	4.824	0.000
.y6	1	4.809	0.924	5.202	0.000
.y7	1	3.302	0.699	4.722	0.000
.y8	1	3.227	0.720	4.482	0.000
ind60	2	0.446	0.075	5.975	0.000
.dem60	2	0.120	0.306	0.394	0.694
.dem65	2	0.097	0.225	0.432	0.666

(optional) extract the various components

```
> out <- sam(model, data = PoliticalDemocracy,
             mm.list = list(ind = "ind60", dem = c("dem60", "dem65")),
             struc.args = list(estimator = "GLS"),
             output = "list")
> names(out)

[1] "MM.FIT"           "Sigma.11"           "step1.free.idx"    "block.mm.idx"
[5] "block.ptm.idx"   "PT.free"            "mm.list"           "PT"
[9] "COV"             "YBAR"               "LAMBDA"            "THETA"
[13] "VETA"            "REL"                "M"                 "lambda"
[17] "alpha"           "MSM"                "MTM"               "FS.mean"
[21] "FS"              "COV.IVETA2"         "LV.NAMES"          "sam.method"
[25] "local.options"  "FIT.PA"             "PT"                "reg.idx"
[29] "step2.free.idx" "extra.id"           "pt.idx"            "pts.idx"
[33] "V2"              "V1"                 "VCOV"              "se"

> out$MM.FIT[[1]]
```

lavaan 0.6-22.2560 ended normally after 21 iterations

Estimator	ML
Optimization method	NLMINB
Number of model parameters	6
Row rank of the constraints matrix	6

Number of observations 75

Model Test User Model:

Test statistic 0.000
Degrees of freedom 0

> out\$FIT.PA

lavaan 0.6-22.2560 ended normally after 28 iterations

Estimator GLS
Optimization method NLMINB
Number of model parameters 5
Number of observations 75

Model Test User Model:

Test statistic 35.817
Degrees of freedom 1
P-value (Chi-square) 0.000

Thank you for attending this workshop!

`https://lavaan.org`

`https://lavaan.ugent.be/about/donate.html`

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