Missing Data in SEM Introduction to SEM with Lavaan



Kyle M. Lang

Department of Methodology & Statistics Utrecht University

Outline

Missing Data Mechanisms

Imputation-Based Solutions

Single Imputation Multiple Imputation MI-Based Analysis

ML-Based Solutions

Maximum Likelihood Estimation
Full Information Maximum Likelihood

Auxiliary Variables



What are Missing Data?

Missing data are empty cells in a dataset where there should be observed values.

 The missing cells correspond to true population values, but we haven't observed those values.



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Missing data are empty cells in a dataset where there should be observed values.

 The missing cells correspond to true population values, but we haven't observed those values.

Not every empty cell is a missing datum.

- Quality-of-life ratings for dead patients in a mortality study
- Firm profitability after the company goes out of business
- Self-reported severity of menstrual cramping for men
- Empty blocks of data following "gateway" items

A Little Notation

 $Y := An N \times P$ Matrix of Arbitrary Data

 $Y_{mis} := \text{The } missing \text{ part of } Y$

 $Y_{obs} := \text{The } observed \text{ part of } Y$

 $R := An N \times P$ response matrix

 $M := An N \times P$ missingness matrix

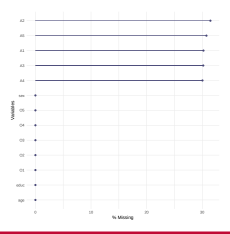
The R and M matrices are complementary.

- $r_{np} = 1$ means y_{np} is observed; $m_{np} = 1$ means y_{np} is missing.
- $r_{np} = 0$ means y_{np} is missing; $m_{np} = 0$ means y_{np} is observed.
- M_p is the *missingness* of Y_p .

```
## Load some useful packages:
library(dplyr)
library(naniar)
library(ggmice)
## Read in some data:
bfi0 <- readRDS("../data/bfi_datasets.rds")</pre>
bfi <- bfi0$incomplete %% select(-matches("N\\d|C\\d|E\\d|male"))
## Compute the variablewise proportions of missing data:
bfi %>% is.na() %>% colMeans() %>% round(2)
 A 1
     A2 A3 A4 A5 O1 O2
                              03
                                   04
                                       05
                                          age sex educ
```

Visualize the percentages missing via **naniar**::gg_miss_var().

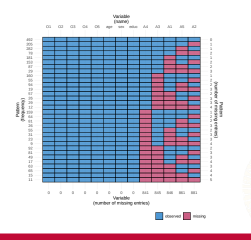
```
gg_miss_var(bfi, show_pct = TRUE)
```





Visualize the missing data patterns via **ggmice**::plot_pattern().

plot_pattern(bfi)





In **lavaan**, we can directly fit a model with incomplete data.

```
library(lavaan)

## Specify the measurement model:
cfaMod <- "
agree = A1 + A2 + A3 + A4 + A5
open = 01 + 02 + 03 + 04 + 05
"

## Estimate the model:
naiveOut <- cfa(cfaMod, data = bfi, std.lv = TRUE)</pre>
```

The model will estimate, just fine...

lavaan	0.6-12.1708	ended nor	mally aft	er 21 ite	rations
Latent	Variables:				
		Estimate	Std.Err	z-value	P(> z)
agree	e =~				
A1		0.561	0.071	7.866	0.000
A2		-0.725	0.056	-13.055	0.000
A3		-0.901	0.063	-14.383	0.000
A4		-0.599	0.074	-8.103	0.000
A5		-0.781	0.059	-13.264	0.000
open	=~				
01		0.490	0.055	8.983	0.000
02		-0.844	0.079	-10.649	0.000
03		0.786	0.058	13.574	0.000
04		0.311	0.055	5.699	0.000
05		-0.874	0.073	-11.987	0.000

Covariances:					
	Estimate	Std.Err	z-value	P(> z)	
agree ~~					
open	-0.254	0.060	-4.204	0.000	
Variances:					
	Estimate	Std.Err	z-value	P(> z)	
.A1	1.662	0.114	14.593	0.000	
.A2	0.781	0.067	11.650	0.000	
.A3	0.865	0.086	10.050	0.000	
.A4	1.776	0.122	14.517	0.000	
.A5	0.863	0.075	11.430	0.000	
.01	0.880	0.064	13.823	0.000	
.02	1.704	0.133	12.776	0.000	
.03	0.658	0.072	9.150	0.000	
.04	0.976	0.065	15.020	0.000	
.05	1.288	0.112	11.460	0.000	

But not everything is as it seems.

	Used	Total
Number of observations	492	2800



Default Approach

Like most software packages, **lavaan** will default to *complete case* analysis when asked to analyze incomplete data.

- In the absence of user input, this is a sensible option.
- That doesn't mean you should actually use deletion to treat the missing data in your analysis.



Default Approach

Like most software packages, **lavaan** will default to *complete case analysis* when asked to analyze incomplete data.

- In the absence of user input, this is a sensible option.
- That doesn't mean you should actually use deletion to treat the missing data in your analysis.

Complete case analysis has two major problems.

- 1. Throws out useful information (potentially a lot of information)
- 2. Probably biases parameter estimates.

To understand the second point, we need to discuss *missing data mechanisms*.

MISSING DATA MECHANISMS



Missing Data Mechanisms

Missing Completely at Random (MCAR)

- $P(R|Y_{mis}, Y_{obs}) = P(R)$
- Missingness is unrelated to any study variables.

Missing at Random (MAR)

- $P(R|Y_{mis}, Y_{obs}) = P(R|Y_{obs})$
- Missingness is related to only the observed parts of study variables.

Missing not at Random (MNAR)

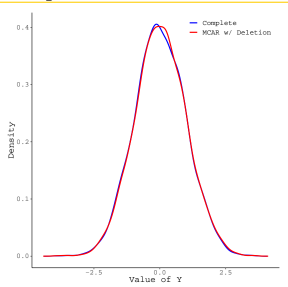
- $P(R|Y_{mis}, Y_{obs}) \neq P(R|Y_{obs})$
- Missingness is related to the unobserved parts of study variables.

Simulate Some Toy Data

```
nObs <- 5000 # Sample Size
pm <- 0.3 # Proportion Missing
sigma \leftarrow matrix(c(1.0, 0.5, 0.3,
                   0.5, 1.0, 0.0,
                   0.3. 0.0. 1.0).
                 ncol = 3
tmp <- rmvnorm(n0bs, c(0, 0, 0), sigma)
x0 \leftarrow tmp[, 1]
y0 <- tmp[ , 2]
z0 \leftarrow tmp[, 3]
cor(y0, x0) # Check correlation between X and Y
[1] 0.4997145
```

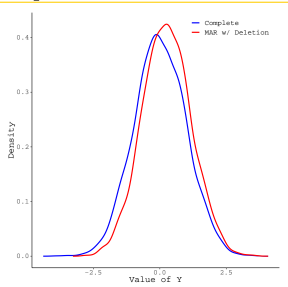
MCAR Example

MCAR Example



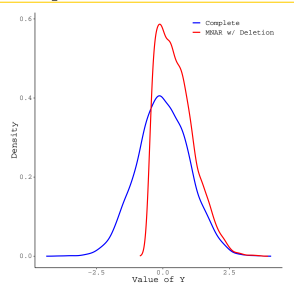
MAR Example

MAR Example



MNAR Example

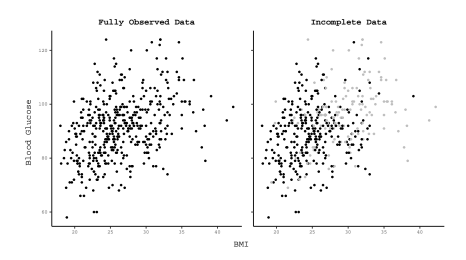
MNAR Example



Effects of Deletion

As we saw in the preceding plots, excluding incomplete cases usually alters the variables' distributions.

- The statistics upon which we base our analyses generally summarize these distributions.
- Problems with the distributions show up as bias in the results of our anlayses.



```
diabetes1 %>% select(bmi, glu, bp) %>% cor()
         bmi glu bp
bmi 1.0000000 0.38868 0.3954109
glu 0.3886800 1.00000 0.3904300
bp 0.3954109 0.39043 1.0000000
diabetes2 %>% select(bmi, glu, bp) %>% cor(use = "complete")
         bmi glu bp
bmi 1.0000000 0.2566595 0.2052338
glu 0.2566595 1.0000000 0.3011547
bp 0.2052338 0.3011547 1.0000000
```

```
mean(diabetes1$glu)
[1] 91.26018
mean(diabetes2$glu, na.rm = TRUE)
[1] 88.86424
var(diabetes1$glu)
[1] 132.1657
var(diabetes2$glu, na.rm = TRUE)
[1] 115.8254
```

Multiple Imputation (MI)

- ullet Replace the missing values with M plausible estimates
 - Essentially, a repeated application of stochastic regression imputation (with a particular type of regression model)
 - Produces unbiased parameter estimates and predictions
 - Produces "correct" standard errors, CIs, and prediction intervals
 - Very, very flexible
 - Computationally expensive



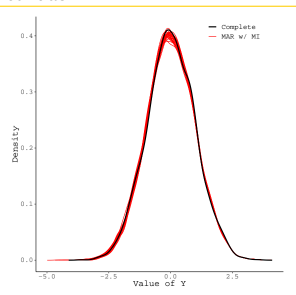
Full Information Maximum Likelihood (FIML)

- Adjust the objective function to only consider the observed parts of the data
 - Models are directly estimated in the presence of missing data
 - The predictors of nonresponse must be included in the model, somehow
 - Unless you write your own optimization program, FIML is only available for certain types of models
 - In linear regression models, FIML cannot treat missing data on predictors (if the predictors are taken as fixed)

What happens when we apply MI to our previous MAR example?

The MI-based parameter estimate looks good.

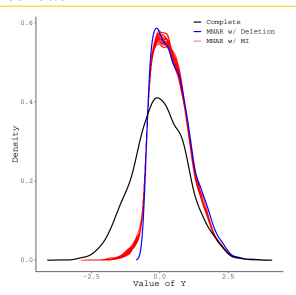
 MI produces unbiased estimates of the parameter when data are MAR.



What about applying MI to our MNAR example?

The MI-based parameter estimate is still biased.

 MI cannot correct bias in parameter estimates when data are MNAR.



MI Example

MI Example

```
## Complete data:
diabetes1 %>% select(bmi, glu, bp) %>% cor()
         bmi
              glu
bmi 1.0000000 0.38868 0.3954109
glu 0.3886800 1.00000 0.3904300
bp 0.3954109 0.39043 1.0000000
## MT:
pooledCorMat(miceOut, c("bmi", "glu", "bp"))
         bmi glu bp
bmi 1.0000000 0.3135162 0.3954109
glu 0.3135162 1.0000000 0.3563903
bp 0.3954109 0.3563903 1.0000000
```

MI Example

```
mean(diabetes1$glu)
[1] 91.26018
with(miceOut, mean(glu)) analyses %>% unlist() %>% mean()
[1] 90.61747
var(diabetes1$glu)
[1] 132, 1657
with(miceOut, var(glu))$analyses %>% unlist() %>% mean()
[1] 123.3748
```

FIML Example

```
fit <- diabetes2 %>%
    select(bmi, glu, bp) %>%
   lavCor(missing = "fiml", output = "sampstat")
## Complete data:
diabetes1 %>% summarize(mean = mean(glu), var = var(glu))
     mean var
1 91.26018 132.1657
## FTMT. .
fit %$% c(mean = mean[["glu"]], var = cov["glu", "glu"])
    mean
          var
 90.82487 125.27146
```

FIML Example

```
## Complete data:
diabetes1 %>% select(bmi, glu, bp) %>% cor() %>% round(3)
     bmi glu bp
bmi 1.000 0.389 0.395
glu 0.389 1.000 0.390
bp 0.395 0.390 1.000
## FTMT.:
fit$cov %>% cov2cor()
   bmi glu bp
bmi 1.000
glu 0.357 1.000
bp 0.395 0.386 1.000
```

IMPUTATION-BASED SOLUTIONS

Prediction Example

To fix ideas, let's consider the *diabetes* data and the following model:

$$Y_{LDL} = \beta_0 + \beta_1 X_{BP} + \beta_2 X_{qluc} + \beta_3 X_{BMI} + \varepsilon$$

Training this model on the first N = 400 patients' data produces the following fitted model:

$$Y_{LDL} = 22.135 + 0.089X_{BP} + 0.498X_{gluc} + 1.48X_{BMI}$$

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Suppose a new patient presents with BP = 121, gluc = 89, and BMI = 30.6. We can predict their LDL score by:

$$\begin{aligned} \hat{Y}_{LDL} &= 22.135 + 0.089(121) + 0.498(89) + 1.48(30.6) \\ &= 122.463 \end{aligned}$$

Imputation is Just Prediction*

In Lecture 3, you heard a bit about missing data imputation.

 Multiple imputation is one of the best ways to treat missing data.

Imputation is nothing more than a type of prediction.

- 1. Train a model on the observed parts of the data, Y_{obs} .
 - Train the imputation model.
- 2. Predict the missing values, Y_{mis} .
 - · Generate imputations.
- 3. Replace the missing values with these predictions.
 - Impute the missing data.

Imputation can be used to support either prediction or inference.

• Our goals will dictate what type of imputation we need to do.

*Levels of Uncertainty Modeling

van Buuren (2018) provides a very useful classification of different imputation methods:

1. Simple Prediction

- The missing data are naively filled with predicted values from some regression equation.
- All uncertainty is ignored.

2. Prediction + Noise

- A random residual error is added to each predicted value to create the imputations.
- o Only uncertainty in the predicted values is modeled.
- The imputation model itself is assumed to be correct and error-free.

3. Prediction + Noise + Model Error

- Uncertainty in the imputation model itself is also modeled.
- Only way to get fully proper imputations in the sense of Rubin (1987).

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The arguments against single imputation can seem archaic and petty. Do we really need to worry about this stuff?

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YES!!! (At least if you care about inference)

The following are results from a simple Monte Carlo simulation:

	Complete Data	Conditional Mean	Stochastic	MI
cor(X, Y)	0.500	0.563	0.498	0.497
Type I Error	0.052	0.138	0.120	0.054

Mean Correlation Coefficients and Type I Error Rates

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Mean Correlation Coefficients and Type I Error Rates

- Conditional mean substitution overestimates the correlation effect.
- Both single imputation methods inflate Type I error rates.
- MI provides unbiased point estimates and accurate Type I error rates.

Simulate Some Toy Data

Simulate Some Toy Data

```
## Impose MAR Nonresponse:
dat1 <- dat0
mVec <- with(dat1, x < quantile(x, probs = pm))
dat1[mVec, "y"] <- NA

## Subset the data:
yMis <- dat1[mVec, ]
yObs <- dat1[!mVec, ]</pre>
```

Look at the Data

```
round(head(dat0, n = 5), 3)

y x z

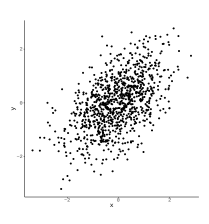
1 0.094 -0.743 0.191

2 1.666 0.542 -0.181

3 0.208 0.617 0.711

4 0.133 -0.827 0.430

5 -0.003 0.441 0.719
```

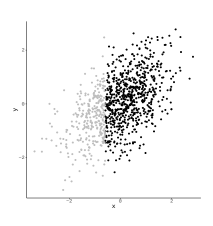


Look at the Data

```
round(head(dat1, n = 5), 3)

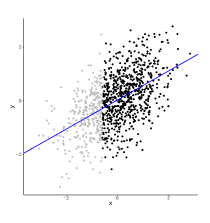
y x z

1 NA -0.743 0.191
2 1.666 0.542 -0.181
3 0.208 0.617 0.711
4 NA -0.827 0.430
5 -0.003 0.441 0.719
```



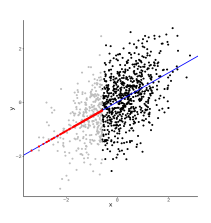
Expected Imputation Model Parameters

```
lsFit \leftarrow lm(y ~x + z, data = y0bs)
beta <- coef(lsFit)</pre>
sigma <- summary(lsFit)$sigma</pre>
as.matrix(beta)
                     Γ,1]
(Intercept) 0.03249194
              0.56993731
х
             -0.12558749
sigma
[1] 0.8452583
```



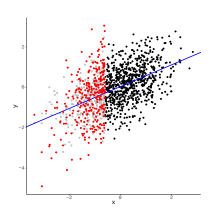
Conditional Mean Substitution

```
## Generate imputations:
imps <- beta[1] +</pre>
    beta[2] * yMis[, "x"] +
   beta[3] * yMis[ , "z"]
## Fill missing cells in Y:
dat1[mVec, "y"] <- imps</pre>
round(head(dat1, n = 5), 3)
1 -0.415 -0.743 0.191
2 1.666 0.542 -0.181
3 0.208 0.617 0.711
4 -0.493 -0.827 0.430
5 -0.003 0.441 0.719
```



Stochastic Regression Imputation

```
## Generate imputations:
imps <- imps +
   rnorm(nrow(yMis), 0, sigma)
## Fill missing cells in Y:
dat1[mVec, "y"] <- imps</pre>
round(head(dat1, n = 5), 3)
1 -0.563 -0.743 0.191
2 1.666 0.542 -0.181
3 0.208 0.617 0.711
4 1.043 -0.827 0.430
5 -0.003 0.441 0.719
```



Flavors of MI

MI simply repeats a single regression imputation M times.

 The specifics of the underlying regression imputation are important.



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MI simply repeats a single regression imputation M times.

 The specifics of the underlying regression imputation are important.

Simply repeating the stochastic regression imputation procedure described above won't suffice.

• Still produces too many Type I errors

	Complete Data	PN-Type	PNE-Type
cor(X, Y)	0.499	0.499	0.498
Type I Error	0.040	0.066	0.046

Mean Correlation Coefficients and Type I Error Rates

 Type I error rates for PN-Type MI are much better than they were for single stochastic regression imputation, but they're still too high.

Proper MI

The problems on the previous slide arise from using the same regression coefficients to create each of the M imputations.

- Implies that you're using the "correct" coefficients.
- This assumption is plainly ridiculous.
 - If we don't know some values of our outcome variable, how can we know the "correct" coefficients to link the incomplete outcome to the observed predictors?



Proper MI

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- Implies that you're using the "correct" coefficients.
- This assumption is plainly ridiculous.
 - If we don't know some values of our outcome variable, how can we know the "correct" coefficients to link the incomplete outcome to the observed predictors?
- Proper MI also models uncertainty in the regression coefficients used to create the imputations.
 - A different set of of coefficients is randomly sampled (using Bayesian simulation) to create each of the *M* imputations.
 - The tricky part about implemented MI is deriving the distributions from which to sample these coefficients.

Setting Up Proper MI

Our imputation model is simply a linear regression model:

$$Y = X\beta + \varepsilon$$

To fully account for model uncertainty, we need to randomly sample both β and $var(\varepsilon) = \sigma^2$.

• Question: Why do we only sample σ^2 and not ε ?



Setting Up Proper MI

Our imputation model is simply a linear regression model:

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To fully account for model uncertainty, we need to randomly sample both β and $var(\varepsilon) = \sigma^2$.

• Question: Why do we only sample σ^2 and not ε ?

For a simple imputation model with a normally distributed outcome and uninformative priors, we need to specify two distributions:

- 1. The marginal posterior distribution of σ^2
- 2. The conditional posterior distribution of β

Marginal Distribution of σ^2

We first specify the marginal posterior distribution for the noise variance, σ^2 .

• This distribution does not depend on any other parameters.

$$\sigma^2 \sim \text{Inv-}\chi^2 \left(N-P, MSE\right)$$
 with $MSE = \frac{1}{N-P} \left(Y - X\hat{\beta}_{ls}\right)^T \left(Y - X\hat{\beta}_{ls}\right)$

• σ^2 follows a scaled inverse χ^2 distribution.



Conditional Distribution of β

We then specify the conditional posterior distribution for β .

• This distribution is conditioned on a specific value of σ^2 .

$$\beta \sim \text{MVN}\left(\hat{\beta}_{ls}, \ \sigma^2(\mathbf{X}^T\mathbf{X})^{-1}\right)$$
 (2)

• β (conditionally) follows a multivariate normal distribution.



PPD of the Missing Data

Once we've sampled our imputation model parameters, we can construct the posterior predictive distribution of the missing data.

- This is the distribution from which we sample our imputed values.
- In practice, we directly compute the imputations based on the simulated imputation model parameters.

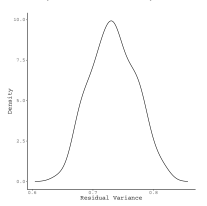
$$Y_{imp} = \mathbf{X}_{mis}\tilde{\boldsymbol{\beta}} + \tilde{\boldsymbol{\varepsilon}}$$
with $\boldsymbol{\varepsilon} \sim N\left(0, \widetilde{\sigma^2}\right)$

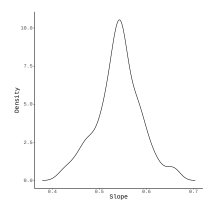
General Steps for Basic MI

With all of the elements in place, we can execute a basic MI by following these steps:

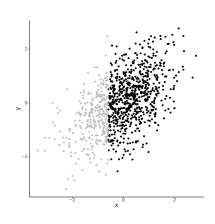
- 1. Find the least squares estimates of β , $\hat{\beta}_{ls}$, by regressing the observed portion of Y onto the the analogous rows of X.
- 2. Use $\hat{\beta}_{ls}$ to parameterize the posterior distribution of σ^2 , given by Equation 1, and draw M samples of σ^2 from this distribution.
- 3. For each of the σ_m^2 , sample a corresponding value of β from Equation 2.
- 4. Plug the M samples of β and σ^2 into Equation 3 to create the M imputations.

Use Bayesian simulation to estimate posterior distributions for the imputation model parameters:





Recall the incomplete data from the single imputation examples.



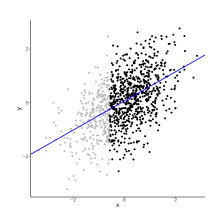
Sample values of β_0 and β_1 :

•
$$\beta_0 = 0.075$$

•
$$\beta_1 = 0.542$$

Define the predicted best-fit line:

$$\hat{Y}_{mis} = 0.075 + 0.542 X_{mis}$$



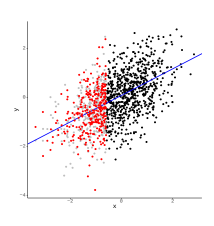
Sample a value of σ^2 :

•
$$\sigma^2 = 0.709$$

Generate imputations using the same procedure described in Single Stochastic Regression Imputation:

$$Y_{imp} = \hat{Y}_{mis} + \varepsilon$$

 $\varepsilon \sim N(0, 0.709)$



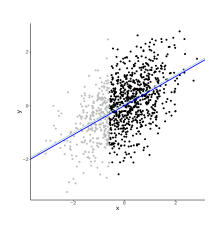
Sample values of β_0 and β_1 :

•
$$\beta_0 = 0.009$$

•
$$\beta_1 = 0.542$$

Define the predicted best-fit line:

$$\hat{Y}_{mis} = 0.009 + 0.542 X_{mis}$$



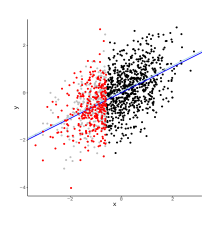
Sample a value of σ^2 :

•
$$\sigma^2 = 0.713$$

Generate imputations using the same procedure described in Single Stochastic Regression Imputation:

$$Y_{imp} = \hat{Y}_{mis} + \varepsilon$$

 $\varepsilon \sim N(0, 0.713)$

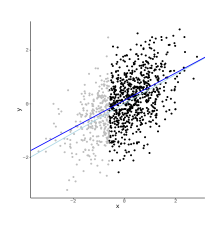


Sample values of β_0 and β_1 :

- $\beta_0 = 0.132$
- $\beta_1 = 0.509$

Define the predicted best-fit line:

$$\hat{Y}_{mis} = 0.132 + 0.509 X_{mis}$$

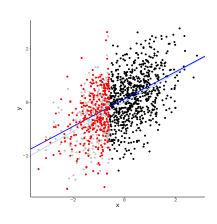


Sample a value of σ^2 :

•
$$\sigma^2 = 0.689$$

Generate imputations using the same procedure described in Single Stochastic Regression Imputation:

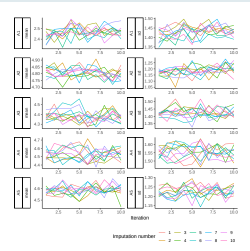
$$Y_{imp} = \hat{Y}_{mis} + \varepsilon$$
$$\varepsilon \sim N(0, 0.689)$$



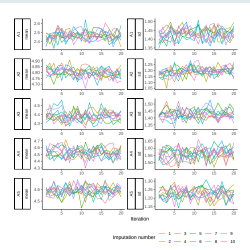
Example

```
## Impute the missing data 10 times:
miceOut <- mice(data = bfi,
              m = 10,
              maxit = 10,
              method = "pmm",
               seed = 235711,
              print = FALSE)
## Check convergence via PSR factor:
rhat(miceOut)
 Variable MissProp RHat_Mean RHat_Variance
1
       A1 30.21429 1.0097875
                           1.118038
2
       A2 31.46429 0.9937958 1.016108
3
       A3 30.17857 1.1010929 1.063382
4
       A4 30.03571 1.0307582 1.039938
       A5 30.75000 1.0201913 1.084875
```

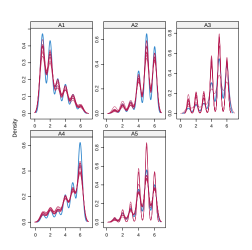
ggmice::plot_trace(miceOut)



ggmice::plot_trace(miceOut)



mice::densityplot(miceOut)

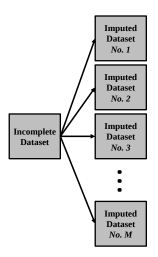


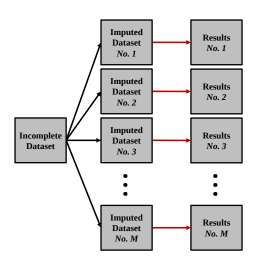
Doing MI-Based Analysis

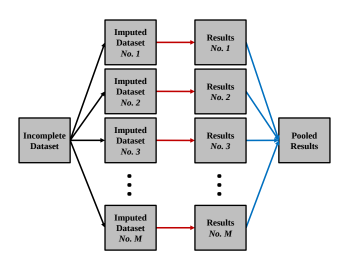
An MI-based data analysis consists of three phases:

- 1. The imputation phase
 - Replace missing values with *M* plausible estimates.
 - Produce *M* completed datasets.
- 2. The analysis phase
 - Estimate M replicates of your analysis model.
 - \circ Fit the same model to each of the M datasets from Step 1.
- 3. The pooling phase
 - Combine the M sets of parameter estimates and standard errors from Step 2 into a single set of MI estimates.
 - Use these pooled parameter estimates and standard errors for inference

Incomplete Dataset







Pooling MI Estimates

Rubin (1987) formulated a simple set of pooling rules for MI estimates.

• The MI point estimate of some interesting quantity, Q^* , is simply the mean of the M estimates, $\{\hat{Q}_m\}$:

$$Q^* = \frac{1}{M} \sum_{m=1}^{M} \hat{Q}_m$$



Pooling MI Estimates

The MI variability estimate, T, is a slightly more complex entity.

• A weighted sum of the *within-imputation* variance, *W*, and the *between-imputation* variance, *B*.

$$W = \frac{1}{M} \sum_{m=1}^{M} \widehat{SE}_{Q,m}^{2}$$

$$B = \frac{1}{M-1} \sum_{m=1}^{M} (\hat{Q}_{m} - Q^{*})^{2}$$

$$T = W + (1 + M^{-1}) B$$

$$= W + B + \frac{B}{M}$$



Inference with MI Estimates

After computing Q^* and T, we combine them in the usual way to get test statistics and confidence intervals.

$$t = \frac{Q^* - Q_0}{\sqrt{T}}$$

$$CI = Q^* \pm t_{crit} \sqrt{T}$$

We must take care with our *df*, though.

$$df = (M-1) \left[1 + \frac{W}{(1+M^{-1})B} \right]^2$$



Fraction of Missing Information

In Lecture 4, we briefly discussed a very desirable measure of nonresponse: *fraction of missing information* (FMI).

$$FMI = \frac{r + \frac{2}{(df+3)}}{r+1} \approx \frac{(1+M^{-1})B}{(1+M^{-1})B+W} \longrightarrow \frac{B}{B+W}$$

where

$$r = \frac{(1+M^{-1})B}{W}$$

The FMI gives us a sense of how much the missing data (and their treatment) have influence our parameter estimates.

• We should report the FMI for an estimated parameter along with other statistics (e.g., t-tests, p-values, effect sizes, etc.).

Example: Analysis & Pooling

Analyze the multiply imputed datasets and pool results:

```
library(semTools)

## semTools::cfa.mi() will do both the analysis and pooling phases:
miOut <-
     cfa.mi(cfaMod, data = complete(miceOut, "all"), std.lv = TRUE)</pre>
```

Example: Analysis & Pooling

Estimate	Std.Err	t-value	P(> t)	FMI	
0.560	0.034	16.331	0.000	0.303	
-0.770	0.027	-28.983	0.000	0.556	
-1.014	0.030	-33.803	0.000	0.556	
-0.708	0.035	-19.977	0.000	0.399	
-0.817	0.028	-28.815	0.000	0.280	
0.635	0.030	21.272	0.000	0.004	
-0.633	0.041	-15.333	0.000	0.014	
0.838	0.033	25.309	0.000	0.026	
0.344	0.033	10.566	0.000	0.014	
-0.641	0.035	-18.367	0.000	0.010	
	0.560 -0.770 -1.014 -0.708 -0.817 0.635 -0.633 0.838 0.344	0.560 0.034 -0.770 0.027 -1.014 0.030 -0.708 0.035 -0.817 0.028 0.635 0.030 -0.633 0.041 0.838 0.033 0.344 0.033	0.560 0.034 16.331 -0.770 0.027 -28.983 -1.014 0.030 -33.803 -0.708 0.035 -19.977 -0.817 0.028 -28.815 0.635 0.030 21.272 -0.633 0.041 -15.333 0.838 0.033 25.309 0.344 0.033 10.566	0.560 0.034 16.331 0.000 -0.770 0.027 -28.983 0.000 -1.014 0.030 -33.803 0.000 -0.708 0.035 -19.977 0.000 -0.817 0.028 -28.815 0.000 0.635 0.030 21.272 0.000 -0.633 0.041 -15.333 0.000 0.838 0.033 25.309 0.000 0.344 0.033 10.566 0.000	0.560 0.034 16.331 0.000 0.303 -0.770 0.027 -28.983 0.000 0.556 -1.014 0.030 -33.803 0.000 0.556 -0.708 0.035 -19.977 0.000 0.399 -0.817 0.028 -28.815 0.000 0.280 0.635 0.030 21.272 0.000 0.004 -0.633 0.041 -15.333 0.000 0.014 0.838 0.033 25.309 0.000 0.026 0.344 0.033 10.566 0.000 0.014

Example: Analysis & Pooling

Covariances:					
COVALIANCES.	Eatimata	Std.Err	t-value	P(> t)	FMI
~~	Estimate	Sta.Eff	t-value	P(> t)	FMI
agree ""					
open	-0.299	0.029	-10.220	0.000	0.305
Variances:					
	Estimate	Std.Err	t-value	P(> t)	FMI
.A1	1.697	0.056	30.215	0.000	0.389
.A2	0.776	0.032	24.268	0.000	0.325
.A3	0.771	0.041	18.707	0.000	0.423
.A4	1.725	0.059	29.171	0.000	0.373
.A5	0.889	0.036	24.420	0.000	0.469
.01	0.878	0.037	23.974	0.000	0.004
.02	2.048	0.071	28.769	0.000	0.008
.03	0.785	0.046	17.043	0.000	0.037
.04	1.369	0.045	30.594	0.000	0.003
.05	1.357	0.050	26.897	0.000	0.008

We cannot simply average the χ^2 statistics as we do the parameter estimates.

 We need to do some fancy processing to get a correctly distributed fit statistic.

Three different formulations have been proposed, and they are typically designated D_1 , D_2 , and D_3 .

- 1. D₁
- 2. D_2
- $3. D_3$

```
what <- c("chisq", "df", "pvalue", "fmi", "rmsea", "cfi", "tli")
fitMeasures(miOut, test = "D3")[what] %>% round(3)

chisq df pvalue fmi rmsea cfi tli
279.070 34.000 0.000 0.417 0.051 0.909 0.880

fitMeasures(miOut, test = "D2")[what] %>% round(3)

chisq df pvalue fmi rmsea cfi tli
243.995 34.000 0.000 0.472 0.047 0.926 0.902
```

```
## Define a restricted model:
cfaMod2 <- paste(cfaMod, "agree ~~ 0 * open", sep = "\n")
## Estimate the restricted model and pool the results:
mi ()11t.2 <-
   cfa.mi(cfaMod2, data = complete(miceOut, "all"), std.lv = TRUE)
## Test the constraint via nested model comparison:
anova(miOut2, miOut, test = "D3")
         df1 df2 pvalue ariv fmi
93.961 1.000 62.196 0.000 0.323 0.244
anova(miOut2, miOut, test = "D2")
     F df1 df2 pvalue ariv fmi
89.292 1.000 114.627 0.000 0.389 0.280
```

```
## Define a model with some parameter labels:
cfaMod3 <- "
agree = 111 * A1 + 121 * A2 + 131 * A3 + 141 * A4 + 151 * A5
open = ^{\sim} 112 * 01 + 122 * 02 + 132 * 03 + 142 * 04 + 152 * 05
## Estimate the model and pool the results:
miOut3 <-
    cfa.mi(cfaMod3, data = complete(miceOut, "all"), std.lv = TRUE)
## Define some constraints:
cons <- "
111 == 112
121 == 122
131 == 132
141 == 142
151 == 152
```

```
## Define some syntax to allow cross-loadings from the "agree"
## factor to the "open" items:
adds <- "agree =~ 01 + 02 + 03"

## Test the parameter relaxations via a score test:
stD2 <- lavTestScore.mi(miOut, add = adds, test = "D2")
stD1 <- lavTestScore.mi(miOut, add = adds, test = "D1")</pre>
```

stD2

```
$test
total score test:
  test F df1 df2 p.value ariv fmi
1 score 21.809 3 213.475 0 0.211 0.174
$uni
univariate score tests:
      lhs op rhs F df1 df2 p.value riv fmi
1 agree=~01 == 0 3.954 1 681.076 0.047 0.130 0.115
2 agree=~02 == 0 40.767 1 418.527 0.000 0.172 0.147
```

3 agree=~03 == 0 29.511 1 335.876 0.000 0.196 0.164

stD1

```
$test
total score test:
 test F df1 df2 p.value ariv fmi
$uni
univariate score tests:
     lhs op rhs F df1 df2 p.value riv fmi
1 agree=~01 == 0 3.989 1 467.030 0.046 0.090 0.083
2 agree=~02 == 0 42.262 1 277.159 0.000 0.122 0.108
```

3 agree=~03 == 0 31.201 1 474.650 0.000 0.089 0.082

ML-BASED SOLUTIONS



FIML Intuition

FIML is an ML estimation method that is robust to ignorable nonresponse.

 FIML partitions the missing information out of the likelihood function so that the model is only estimated from the observed parts of the data.

After a minor alteration to the likelihood function, FIML reduces to simple ML estimation.

• So, let's review ML estimation before moving forward.



Maximum Likelihood Estimation

ML estimation simply finds the parameter values that are "most likely" to have given rise to the observed data.

- The *likelihood* function is just a probability density (or mass) function with the data treated as fixed and the parameters treated as random variables.
- Having such a framework allows us to ask: "Given that I've observed these data values, what parameter values most probably describe these data?"

Maximum Likelihood Estimation

ML estimation is usually employed when there is no closed form solution for the parameters we seek.

 This is why you don't usually see ML used to fit general linear models.

After choosing a likelihood function, we iteratively optimize the function to produce the ML estimated parameters.

 In practice, we nearly always work with the natural logarithm of the likelihood function (i.e., the loglikelihood).

ML Intuition

Let's say we have the following N = 10 observations.

- We assume these data come from a normal distribution with a known variance of $\sigma^2 = 1$.
- We want to estimate the mean of this distribution, μ .

```
(y <- rnorm(n = 10, mean = 5, sd = 1))

[1] 4.947835 4.183180 3.958366 6.425695 5.759473
[6] 4.159033 2.739487 4.921964 4.409978 6.416356
```

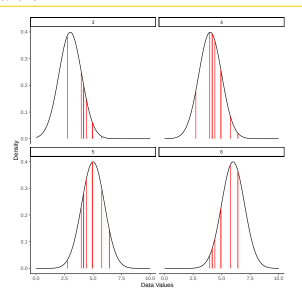
ML Intuition

In ML estimation, we would define different normal distributions.

- Every distribution would have $\sigma^2 = 1$.
- Each distribution would have a different value of μ .

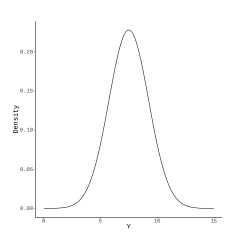
We then compare the observed data to those distributions and see which distribution best fits the data.

ML Intuition



Suppose we have the following model:

$$Y \sim N(\mu, \sigma^2)$$
.



For a given Y_n , we have:

$$P\left(Y_n|\mu,\sigma^2\right) = \frac{1}{\sqrt{2\pi\sigma^2}} e^{-\frac{(Y_n-\mu)^2}{2\sigma^2}}.$$
 (4)

If we plug estimated parameters into Equation 4, we get the probability of observing Y_n given $\hat{\mu}$ and $\hat{\sigma}^2$:

$$P\left(Y_n|\hat{\mu},\hat{\sigma}^2\right) = \frac{1}{\sqrt{2\pi\hat{\sigma}^2}} e^{-\frac{(Y_n-\hat{\mu})^2}{2\hat{\sigma}^2}}.$$
 (5)

Applying Equation 5 to all N observations and multiplying the results produces a *likelihood*:

$$L\left(\hat{\mu},\hat{\sigma}^{2}\right) = \prod_{n=1}^{N} P\left(Y_{n}|\hat{\mu},\hat{\sigma}^{2}\right).$$

We generally want to work with the natural logarithm of Equation 5. Doing so gives the *loglikelihood*:

$$\mathcal{L}(\hat{\mu}, \hat{\sigma}^2) = \ln \prod_{n=1}^{N} P(Y_n | \hat{\mu}, \hat{\sigma}^2)$$
$$= -\frac{N}{2} \ln 2\pi - N \ln \hat{\sigma} - \frac{1}{2\hat{\sigma}^2} \sum_{n=1}^{N} (Y_n - \hat{\mu})^2$$

ML tries to find the values of $\hat{\mu}$ and $\hat{\sigma}^2$ that maximize $\mathcal{L}(\hat{\mu}, \hat{\sigma}^2)$.

• Find the values of $\hat{\mu}$ and $\hat{\sigma}^2$ that are most likely, given the observed values of Y.

Suppose we have a linear regression model:

$$Y = \beta_0 + \beta_1 X + \varepsilon,$$

$$\varepsilon \sim N(0, \sigma^2).$$

This model can be equivalently written as:

$$Y \sim N \left(\beta_0 + \beta_1 X, \sigma^2\right)$$

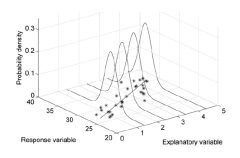


Image retrieved from: http://www.seaturtle.org/mtn/
 archives/mtn122/mtn122p1.shtml

Likelihoods

For a given $\{Y_n, X_n\}$, we have:

$$P(Y_n|X_n, \beta_0, \beta_1, \sigma^2) = \frac{1}{\sqrt{2\pi\sigma^2}} e^{-\frac{(Y_n - \beta_0 - \beta_1 X_n)^2}{2\sigma^2}}.$$
 (6)

If we plug our estimated parameters into Equation 6, we get the probability of observing Y_n given $\hat{Y}_n = \hat{\beta}_0 + \hat{\beta}_1 X_n$ and $\hat{\sigma}^2$.

$$P\left(Y_{n}|X_{n},\hat{\beta}_{0},\hat{\beta}_{1},\hat{\sigma}^{2}\right) = \frac{1}{\sqrt{2\pi\hat{\sigma}^{2}}}e^{-\frac{\left(Y_{n}-\hat{\beta}_{0}-\hat{\beta}_{1}X_{n}\right)^{2}}{2\hat{\sigma}^{2}}}$$
(7)

Likelihoods

So, our final loglikelihood function would be the following:

$$\mathcal{L}\left(\hat{\beta}_{0},\hat{\beta}_{1},\hat{\sigma}^{2}\right) = \ln \prod_{n=1}^{N} P\left(Y_{n}|X_{n},\hat{\beta}_{0},\hat{\beta}_{1},\hat{\sigma}^{2}\right)$$
$$= -\frac{N}{2} \ln 2\pi - N \ln \hat{\sigma} - \frac{1}{2\hat{\sigma}^{2}} \sum_{n=1}^{N} \left(Y_{n} - \hat{\beta}_{0} - \hat{\beta}_{1}X_{n}\right)^{2}.$$



Example

```
## Fit a model:
out1 <- lm(ldl ~ bp + glu + bmi, data = diabetes)
## Extract the predicted values and estimated residual
## standard error.
yHat <- predict(out1)</pre>
s <- summary(out1)$sigma
## Compute the row-wise probabilities:
pY <- dnorm(diabetes$ldl, mean = yHat, sd = s)
## Compute the loglikelihood, and compare to R's version:
sum(log(pY)); logLik(out1)[1]
[1] -2109.939
[1] -2109.93
```

Likelihoods

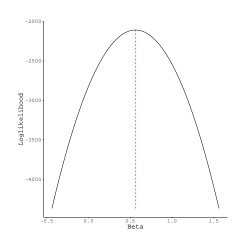
Plugging our estimates into the logliklihood equation gives:

$$\begin{split} \mathcal{L}\left(\hat{\beta},\hat{\sigma}^2\right) &= -221 \times \ln\left(2\pi\right) - 442 \times \ln\left(28.77\right) \\ &- \frac{\sum_{n=1}^{442} \left(Y_n - 26.59 - 0.08X_n^{bp} - 0.56X_n^{glu} - 1.13X_n^{bmi}\right)^2}{1655} \\ &= -2109.93 \end{split}$$

Likelihood Function Visualized

Here, we see the conditional loglikelhood function for β_{qlu} .

• What loglikelihoods do we get by varying the value of $\hat{\beta}_{glu}$ while keep all other parameters fixed at their ML estimates?



Steps of ML

- 1. Choose a probability distribution, $f(Y|\theta)$, to describe the distribution of the data, Y, given the parameters, θ .
- 2. Choose some estimate of θ , $\hat{\theta}^{(i)}$.
- 3. Compute each row's contribution to the loglikelihood function by evaluating: $\ln \left[f\left(Y_n|\hat{\theta}^{(i)}\right) \right]$.
- 4. Sum the individual loglikelihood contributions from Step 3 to find the loglikelihood value, $\mathcal{L}^{(i)}$.
- 5. Choose a "better" estimate of the parameters, $\hat{\theta}^{(i+1)}$, and repeat Steps 3 and 4.
- 6. Repeat Steps 3 5 until the change between $\mathcal{L}^{(i-1)}$ and $\mathcal{L}^{(i)}$ falls below some trivially small threshold.
- 7. Take $\hat{\theta}^{(i)}$ as your estimate.

Multivariate Normal Distribution

The PDF for the multivariate normal distribution is:

$$P(\boldsymbol{Y}|\boldsymbol{\mu},\boldsymbol{\Sigma}) = \frac{1}{\sqrt{(2\pi)^P|\boldsymbol{\Sigma}|}} e^{-\frac{1}{2}(\boldsymbol{Y}-\boldsymbol{\mu})^T\boldsymbol{\Sigma}^{-1}(\boldsymbol{Y}-\boldsymbol{\mu})}.$$

So, the multivariate normal loglikelihood is:

$$\mathcal{L}(\boldsymbol{\mu}, \boldsymbol{\Sigma}) = -\left[\frac{P}{2}\ln(2\pi) + \frac{1}{2}\ln|\boldsymbol{\Sigma}| + \frac{1}{2}\right] \sum_{n=1}^{N} (\mathbf{Y}_{n} - \boldsymbol{\mu})^{T} \boldsymbol{\Sigma}^{-1} (\mathbf{Y}_{n} - \boldsymbol{\mu}).$$

Which can be further simplified if we multiply through by -2:

$$-2\mathcal{L}(\mu,\Sigma) = \left[P\ln(2\pi) + \ln|\Sigma|\right] \sum_{n=1}^{N} (\mathbf{Y}_n - \mu)^T \Sigma^{-1} (\mathbf{Y}_n - \mu).$$

Recall the nth observation's contribution to the multivariate normal loglikelihood function:

$$\mathcal{L}\left(\boldsymbol{\mu},\boldsymbol{\Sigma}\right)_{n} = -\frac{P}{2} \ln(2\pi) - \frac{1}{2} \ln|\boldsymbol{\Sigma}| - \frac{1}{2} (\boldsymbol{Y}_{n} - \boldsymbol{\mu})^{T} \boldsymbol{\Sigma}^{-1} (\boldsymbol{Y}_{n} - \boldsymbol{\mu}).$$

It turns out that this function is readily available in R via the **mvtnorm** package:

```
## Vector of row-wise contributions to the overall LL:
110 <- dmvnorm(y, mean = mu, sigma = sigma, log = TRUE)</pre>
```

We can wrap the preceding code in a nice R function:

```
## Complete data loglikelihood function:
11 <- function(par, data) {</pre>
    ## Extract the parameter matrices:
    par <- getParams(par, p = ncol(data))</pre>
    ## Compute the row-wise contributions to the LL:
    110 <- dmvnorm(data.
                    mean = par mu,
                    sigma = par$sigma,
                    log = TRUE)
    sum(110) # Return the overall LL value
```

The **optimx** package can numerically optimize arbitrary functions.

• We can use it to (semi)manually implement ML.

```
## Subset the 'diabetes' data:
dat1 <- diabetes %>% select(bmi, ldl, glu) %>% as.matrix()
## Choose some starting values:
m0 < -rep(0, 3)
s0 <- diag(3) %>% vecChol()
par0 \leftarrow c(m0, s0)
## Use optimx() to numerically optimize the LL function:
mle <- optimx(par = par0,
             fn = 11,
             data = dat1,
             method = "BFGS",
             control = list(maximize = TRUE, maxit = 1000)
             ) %>% quiet()
```

Check convergence and extract the optimized parameters:

Estimate the same parameters using **lavaan**::lavCor().

```
## Estimate the parameters:
fit <- diabetes %>%
    select(bmi, ldl, glu) %>%
    lavCor(meanstructure = TRUE, output = "sampstat")

## Extract the estimated parameters:
muHat2 <- fit$mean
sigmaHat2 <- fit$cov</pre>
```

	bmi	ldl	glu
Manual	26.38	115.44	91.26
Lavaan	26.38	115.44	91.26

Estimated Means

	bmi	ldl	glu		bmi	ldl	glu
bmi	19.48	35.01	19.70	bmi	19.52	35.09	19.74
ldl	35.01	922.86	101.37	ldl	35.09	924.96	101.61
glu	19.70	101.37	131.87	glu	19.74	101.61	132.17

Manual Covariance Matrix

Lavaan Covariance Matrix

From ML to FIML

The nth observation's contribution to the multivariate normal loglikelihood function would be the following:

$$\mathcal{L}(\mu, \Sigma)_{n} = -\frac{P}{2}\ln(2\pi) - \frac{1}{2}\ln|\Sigma| - \frac{1}{2}(\mathbf{Y}_{n} - \mu)^{T}\Sigma^{-1}(\mathbf{Y}_{n} - \mu).$$
 (8)



From ML to FIML

The nth observation's contribution to the multivariate normal loglikelihood function would be the following:

$$\mathcal{L}(\mu, \Sigma)_{n} = -\frac{P}{2}\ln(2\pi) - \frac{1}{2}\ln|\Sigma| - \frac{1}{2}(\mathbf{Y}_{n} - \mu)^{T}\Sigma^{-1}(\mathbf{Y}_{n} - \mu).$$
 (8)

FIML just tweaks Equation 8 a tiny bit:

$$\mathcal{L}\left(\boldsymbol{\mu},\boldsymbol{\Sigma}\right)_{fiml,n} = -\frac{P}{2}\ln(2\pi) - \frac{1}{2}\ln|\boldsymbol{\Sigma}_q| - \frac{1}{2}(\boldsymbol{Y}_n - \boldsymbol{\mu}_q)^T\boldsymbol{\Sigma}_q^{-1}(\boldsymbol{Y}_n - \boldsymbol{\mu}_q).$$

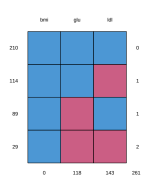
Where q = 1, 2, ..., Q indexes response patterns.

First, punch some holes in our example data.

Visualize the Response Patterns

The data contain 4 unique response patterns.

- We'll define 4 different version of μ and Σ .
- We'll calculate each individual loglikelihood contributions using the appropriate flavor of μ and Σ .



```
## Compute the within-pattern contributions to the LL:
110 <- function(i, mu, sigma, pats, ind, data) {
    ## Define the current response pattern:
   p1 <- pats[i, ]
    if(sum(p1) > 1) # More than one observed variable?
       dmvnorm(x = data[ind == i, p1],
               mean = mu[p1],
               sigma = sigma[p1, p1],
               log = TRUE)
   else
       dnorm(x = data[ind == i, p1],
             mean = mu[p1],
             sd = sqrt(sigma[p1, p1]),
             log = TRUE)
```

```
## FIML loglikelihood function:
llm <- function(par, data, pats, ind) {</pre>
    ## Extract the parameter matrices:
    par <- getParams(par, p = ncol(data))</pre>
    ## Compute the pattern-wise contributions to the LL:
    111 \leftarrow sapply(X = 1:nrow(pats),
                  FUN = 110,
                  mu = par$mu,
                   sigma = par$sigma,
                  pats
                         = pats,
                  ind = ind,
                  data = data)
    sum(unlist(ll1))
```

```
## Summarize response patterns:
pats <- uniquecombs(!is.na(dat2))</pre>
ind <- attr(pats, "index")</pre>
## Choose some starting values:
mO <- colMeans(dat2, na.rm = TRUE)
s0 <- cov(dat2, use = "pairwise") %>% vecChol()
par0 \leftarrow c(m0, s0)
## Use optimx() to numerically optimize the LL function:
mle <- optimx(par = par0,</pre>
              fn = llm,
              data = dat2,
              pats = pats,
              ind = ind,
              method = "BFGS",
              control = list(maximize = TRUE, maxit = 1000)
              ) %>% quiet()
```

Check convergence and extract the optimized parameters:

To use FIML in **lavaan**, we only nee to set the missing = "fiml" argument.

• Do the same analysis via lavaan::cfa().

```
## Define the model in lavaan syntax:
mod <- "
bmi ~~ ldl + glu
ldl ~~ glu
## Fit the model with lavaan::cfa():
fit <- cfa(mod, data = dat2, missing = "fiml")</pre>
## Extract the estimated parameters:
muHat2 <- inspect(fit, "est")$nu</pre>
sigmaHat2 <- inspect(fit, "theta")</pre>
```

	bmi	ldl	glu
Manual	26.38	115.87	90.88
Lavaan	26.38	115.87	90.88

Estimated Means

	bmi	ldl	glu		bmi	ldl	glu
bmi	19.48	24.61	18.48	bmi	19.48	24.64	18.49
ldl	24.61	902.13	76.97	ldl	24.64	902.20	77.01
glu	18.48	76.97	132.55	glu	18.49	77.01	132.56

Manual Covariance Matrix

Lavaan Covariance Matrix

It is very easy to apply FIML to latent variable models in lavaan.

```
## Estimate the CFA from the beginning of the lecture using FIML: fimlOut \leftarrow cfa(cfaMod, data = bfi, std.lv = TRUE, missing = "fiml")
```

```
partSummary(fimlOut, 1:5)
lavaan 0.6-12.1708 ended normally after 41 iterations
  Estimator
                                                      MT.
  Optimization method
                                                  NIMINE
  Number of model parameters
                                                      31
  Number of observations
                                                    2800
                                                      32
  Number of missing patterns
Model Test User Model:
                                                 360.865
  Test statistic
  Degrees of freedom
                                                      34
                                                   0.000
  P-value (Chi-square)
Parameter Estimates:
```

```
partSummary(fimlOut, 7, fmi = TRUE)
Latent Variables:
                   Estimate
                            Std.Err
                                      z-value P(>|z|)
                                                            FMT
  agree =~
   A1
                     0.538
                               0.038
                                      14.109
                                                 0.000
                                                          0.399
   A2
                    -0.764
                               0.030 -25.530
                                                 0.000
                                                          0.417
   A3
                    -0.989
                               0.033 - 29.722
                                                 0.000
                                                          0.434
   A4
                    -0.693
                               0.039 -17.953
                                                 0.000
                                                          0.407
   A5
                    -0.838
                               0.031
                                     -26.622
                                                 0.000
                                                          0.403
  open =~
   Ω1
                      0.635
                               0.025
                                       24.968
                                                 0.000
                                                          0.003
   02
                     -0.640
                               0.036
                                      -17.615
                                                 0.000
                                                          0.004
   0.3
                      0.831
                                                          0.008
                               0.029
                                      28.880
                                                 0.000
   Π4
                      0.345
                               0.028
                                       12.333
                                                 0.000
                                                          0.002
   05
                     -0.647
                               0.031
                                      -20.890
                                                 0.000
                                                          0.005
```

```
partSummary(fimlOut, 9, fmi = TRUE)
Intercepts:
                                       z-value P(>|z|)
                   Estimate
                             Std.Err
                                                              FMI
   .A1
                      2.422
                                0.031
                                        77.128
                                                  0.000
                                                            0.281
   . A2
                                                            0.252
                      4.814
                                0.025
                                       189.726
                                                  0.000
   .A3
                      4.616
                                                  0.000
                                                            0.232
                                0.028
                                       163.443
   . A4
                      4.735
                                0.032
                                       146.063
                                                  0.000
                                                            0.275
   . A5
                      4.573
                                0.027
                                       167.561
                                                  0.000
                                                            0.243
   .01
                      4.816
                                0.021
                                       225,180
                                                  0.000
                                                           -0.000
   .02
                      2.713
                                0.030
                                        91.745
                                                  0.000
                                                           -0.000
   .03
                      4.441
                                0.023
                                       192.731
                                                  0.000
                                                           -0.000
   .04
                      4.894
                                0.023
                                       212.316
                                                  0.000
                                                           -0.000
   .05
                      2.490
                                                           -0.000
                                0.025
                                        99.119
                                                  0.000
                      0.000
   agree
                      0.000
   open
```

```
partSummary(fimlOut, 10, fmi = TRUE)
Variances:
                                        z-value P(>|z|)
                    Estimate
                              Std.Err
                                                               FMI
   .A1
                       1.693
                                0.059
                                         28.649
                                                   0.000
                                                             0.352
   . A2
                                0.037
                                                             0.463
                       0.765
                                         20.491
                                                   0.000
   .A3
                       0.736
                                0.047
                                                   0.000
                                         15.632
                                                             0.489
   . A4
                       1.654
                                0.061
                                         27.307
                                                   0.000
                                                             0.354
   . A5
                       0.876
                                0.043
                                         20.256
                                                   0.000
                                                             0.484
   .01
                       0.878
                                0.031
                                         28,191
                                                   0.000
                                                             0.003
   .02
                       2.039
                                0.062
                                         32,972
                                                   0.000
                                                             0.005
   .03
                       0.797
                                0.040
                                                             0.006
                                         19.848
                                                   0.000
   .04
                       1.369
                                0.038
                                         35.732
                                                   0.000
                                                             0.002
   .05
                       1.349
                                                             0.005
                                0.044
                                         30.369
                                                   0.000
                       1.000
    agree
                       1.000
    open
```

FMI with FIML

As you saw above, we can also estimate the FMI when using FIML.

 The FMI is calculated using the method described by Savalei and Rhemtulla (2012).

Savalei and Rhemtulla (2012) take an information-theoretic approach to defining the FMI.

- Based on the Missing Information Principle of Orchard and Woodbury (1972)
- Their FMI estimates the ratio of missing to complete information for each parameter.

You can use this method to compute the FMI for sufficient statistics via the **semTools**::fmi() function.

AUXILIARY VARIABLES



Satisfying the MAR Assumption

Like MI, FIML also requires MAR data.

• Parameters will be biased with MAR is violated.

Unlike MI, FIML directly treats the missing data while estimating the analysis model.

- The MAR predictors must be included in the analysis model.
- Otherwise, FIML reduces to pairwise deletion.

If the MAR predictors are not substantively interesting variables, naively included them in the analysis model can change the model's meaning.

Saturated Correlates Technique

Graham (2003) developed the *saturated correlates* approach to meet two desiderata:

- 1. Satisfy the MAR assumption by incorporating MAR predictors into the analysis model.
- 2. Do not affect the fit or substantive meaning of the analysis model.

The approach entails incorporating the MAR predictors via a fully-saturated covariance structure:

- Allow all MAR predictors to co-vary with all other MAR predictors.
- 2. Allow all MAR predictors to co-vary with all observed variables in the analysis model (or their residuals).

We can use the lavaan.auxiliary() function (or one of its wrappers) from **semTools** to streamline the analysis.



The cfa.auxiliary() function has automatically added the following paths to our model.

```
age ~~ age
                                       male ~~ male
age ~~ male
                                       male ~~ A1
age ~~ A1
                                       male ~~ A2
age ~~ A2
                                       male ~~ A3
age ~~ A3
                                       male ~~ A4
age ~~ A4
                                       male ~~ A5
age ~~ A5
                                       male ~~ 01
age ~~ 01
                                      male ~~ 02
age ~~ 02
                                       male ~~ 03
age ~~ 03
                                       male ~~ 04
age ~~ 04
                                       male ~~ 05
age ~~ 05
```

The auxiliaries have been correlated with all other variables.



The degrees of freedom have not changed, though.

```
## Naive FIML:
fitMeasures(fimlOut, "df")

df
34

## FIML w/ saturated correlates:
fitMeasures(fimlOut2, "df")

df
34
```

Let's compare the effects of the various missing data treatments on the latent covariance estimates.

	•		Multiple Imputation		
Est	-0.306	-0.254	055	-0.290	-0.295
FMI	—	—		0.147	0.146

Latent Covariances



References

- Graham, J. W. (2003). Adding missing-data-relevant variables to FIML-based structural equation models. *Structural Equation Modeling: A Multidisciplinary Journal*, *10*(1), 80–100. doi: 10.1207/S15328007SEM1001_4
- Orchard, T., & Woodbury, M. A. (1972). A missing information principle: Theory and applications. In *Proceedings of the sixth berkeley symposium on mathematical statistics and probability, volume 1: Theory of statistics.*
- Rubin, D. B. (1987). *Multiple imputation for nonresponse in surveys* (Vol. 519). New York, NY: John Wiley & Sons.
- Savalei, V., & Rhemtulla, M. (2012). On obtaining estimates of the fraction of missing information from full information maximum likelihood. *Structural Equation Modeling: A Multidisciplinary Journal*, 19(3), 477–494.

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