Missing Values in Clinical Research (EP16) Multiple Imputation

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Outline

Part I: Multiple Imputation

How does multiple imputation work?

- The ideas behind MI
- Understanding sources of uncertainty
- Implementation of MI and MICE

Part II: Multiple Imputation Workflow

How to perform MI with the **mice** package in R, from getting to know the data to the final results.

Practical: Imputation with mice

Outline (cont.)

Part III: When MICE might fail

Introduction to

- settings where standard use of mice is problematic
- alternative imputation approaches
- alternative R packages

Practical: Imputation in complex settings

Part IV: Multiple Imputation Strategies

Some tips & tricks

Part I Multiple Imputation

Outline of Part I

- 1. What is Multiple Imputation?
 - 1.1 History & Ideas
 - 1.2 Three steps
- 2. Imputation step
 - 2.1 Univariate missing data
 - 2.2 Multivariate missing data
 - 2.3 FCS/MICE
 - 2.4 Checking convergence
- 3. Analysis step
- 4. Pooling
 - 4.1 Why pooling?
 - 4.2 Rubin's Rules

5. A closer look at the imputation step

- 5.1 Bayesian multiple imputation
- 5.2 Bootstrap multiple imputation
- 5.3 Semi-parametric imputation
- 5.4 What is implemented in software?

1. What is Multiple Imputation? 1.1. History & Ideas

- Developed by **Donald B. Rubin** in the 1970s,
- to handle missing values in **public use databases**, e.g., census data provided by the government,
- motivated by the increase in missing values, and
- increased availability of computers.

1.1. History & Ideas

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- increased availability of computers.

Such data should be usable by [11]

- a large number of analysts, who commonly have to rely on
- standard software that can only handle complete data, and usually
- are not experts in handling incomplete data.

1.1. History & Ideas

Rubin's thoughts: [12]

One imputed value can not be correct in general.

→ We need to represent missing values by a **number of imputations**.

To find sensible values to fill in, we need some kind of model.

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Missing data has a distribution.



To find sensible values to fill in, we need some kind of model.



This distribution depends on assumptions that have been made about the model.

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This distribution depends on assumptions that have been made about the model.



What we want to impute is the 'predictive distribution' of the missing values given the observed values.

6 / 225

1.1. History & Ideas

How to obtain that predictive distribution?

Rubin suggests to

- fit a model to the observed data ("respondents"), and to
- obtain for each "nonrespondent" the conditional distribution of the missing data (given the observed data) as if he/she was a respondent.
- → We assume nonrespondents are just like respondents, and obtain the predictive distribution from the model of the respondents data.

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Example: survey about age, gender and height

Boys aged 10 - 12 years old answered (on average) that they are 1.45m tall.

➡ We assume that boys aged 10 to 12 who did not report their height are also around 1.45m tall.

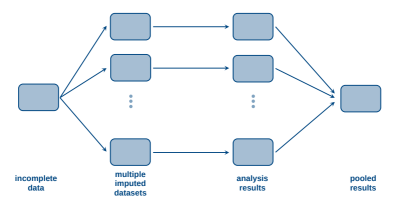
1.1. History & Ideas

How to represent the multiple imputed values?

For each missing value, we now have multiple imputed values.

- For each set of imputed values, create a dataset (those datasets agree in the observed values but imputed values differ).
- Analyse each dataset, and
- take the results from each analysis.
- → We can describe how (much) the **results vary between the imputed datasets**, and calculate summary measures.

1.2. Three steps



In summary:

- 1. **Imputation:** impute multiple times → multiple completed datasets
- 2. Analysis: analyse each of the datasets
- 3. **Pooling:** combine results, taking into account additional uncertainty

How can we actually get imputed values?

For now: assume only one continuous variable has missing values (univariate missing data)

X_1	X_2	<i>X</i> ₃	X_4
√	NA	✓	✓
\checkmark	\checkmark	\checkmark	\checkmark
\checkmark	NA	\checkmark	\checkmark
÷	÷	÷	:

How can we actually get imputed values?

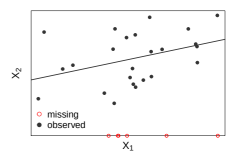
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✓	NA	✓	√
\checkmark	\checkmark	\checkmark	\checkmark
\checkmark	NA	\checkmark	\checkmark
:	÷	÷	:

Idea: Predict values

Model:

$$x_{i2} = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i3} + \beta_3 x_{i4} + \varepsilon_i$$



2. Imputation step

2.1. Univariate missing data

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For now: assume only one continuous variable has missing values (univariate missing data)

X_1	X_2	<i>X</i> ₃	X_4
✓	NA	√	√
\checkmark	\checkmark	\checkmark	\checkmark
\checkmark	NA	\checkmark	\checkmark
÷	÷	÷	:

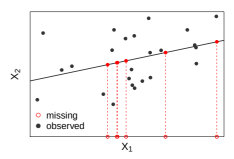
Idea: Predict values

Model:

$$x_{i2} = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i3} + \beta_3 x_{i4} + \varepsilon_i$$

Imputed/predicted value:

$$\hat{x}_{i2} = \hat{\beta}_0 + \hat{\beta}_1 x_{i1} + \hat{\beta}_2 x_{i3} + \hat{\beta}_3 x_{i4}$$



Problem:

- We can obtain only one imputed value per missing value, but we wanted a whole distribution.
- The predicted values do not take into account the added **uncertainty** due to the missing values.

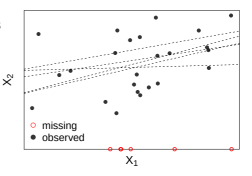
Problem:

- We can obtain **only one imputed value** per missing value, but we wanted a whole distribution.
- The predicted values do not take into account the added **uncertainty** due to the missing values.
- ➡ We need to take into account two sources of uncertainty:
 - The **parameters** are estimated with **uncertainty** (represented by the std. error).
 - There is random variation / prediction error (variation of the residuals).

Taking into account uncertainty about the parameters β :

We assume that β has a distribution, and we can sample realizations of β from that distribution.

When plugging the different realizations of β into the predictive model, we obtain slightly different regression lines.

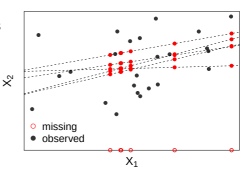


Taking into account uncertainty about the parameters β :

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With each set of coefficients, we also get slightly **different predicted values**.

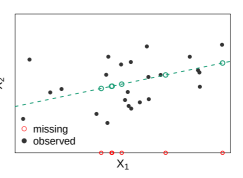


Taking into account the prediction error:

The model does not fit the data perfectly: observations are scattered around the regression lines.

We assume that the data have a distribution, where

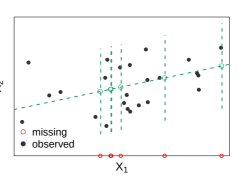
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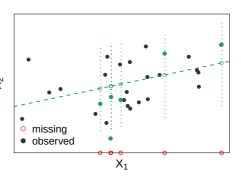
- the **mean** for each value is given by the **predictive model**, and
- the **variance** is determined by the variance of the residuals ε .



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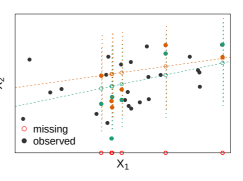
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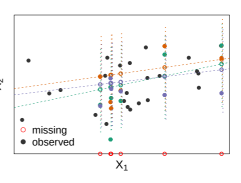
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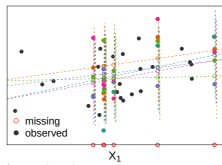


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We assume that the data have a distribution, where

- the mean for each value is given by the predictive model, and
- the **variance** is determined by the variance of the residuals ε .



In the end, we obtain one imputed dataset for each color.

Multivariate missing data:

What if we have missing values in more than one variable?

2. Imputation step

2.2. Multivariate missing data

Multivariate missing data:

What if we have missing values in more than one variable?

In case of **monotone missing values** we can use the technique for univariate missing data in a chain: impute x_4 given x_1 impute x_3 given x_1 and x_4 impute x_2 given x_1 , x_4 and x_3

X_1	X_2	<i>X</i> ₃	X_4
√	NA	√	✓
\checkmark	NA	NA	\checkmark
\checkmark	NA	NA	NA
:	:	:	:

2. Imputation step

2.2. Multivariate missing data

Multivariate missing data:

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In case of **monotone missing values** we can use the technique for univariate missing data in a chain: impute x_4 given x_1 impute x_3 given x_1 and x_4 impute x_2 given x_1 , x_4 and x_3

X_1	X_2	X_3	X_4
√	NA	\checkmark	√
\checkmark	NA	NA	\checkmark
\checkmark	NA	NA	NA
:	÷	÷	÷

When we have **non-monotone missing data** there is no sequence without conditioning on unobserved values.

X_1	X_2	<i>X</i> ₃	X_4
✓	NA	√	✓
NA	\checkmark	NA	NA
\checkmark	NA	\checkmark	NA
:	÷	÷	:

There are **two popular approaches** for the imputation step in **multivariate non-monotone** missing data:

Fully conditional specification

- Multiple Imputation using Chained Equations (MICE)
- sometimes also: sequential regression
- Implemented in SPSS, R, Stata, SAS, ...
- our focus here

There are **two popular approaches** for the imputation step in **multivariate non-monotone** missing data:

Fully conditional specification

- Multiple Imputation using Chained Equations (MICE)
- sometimes also: sequential regression
- Implemented in SPSS, R, Stata, SAS, ...
- our focus here

Joint model imputation

(more details later)

Markov Chain Monte Carlo

is a technique to **draw samples from a complex probability distribution** by creating a chain of random variables (a Markov Chain). The distribution each element in the chain is sampled from depends on the value of the previous element. When certain conditions are met, the chain eventually stabilizes and by continuing to sample elements of the chain a sample from the complex distribution of interest can be obtained.

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Gibbs sampling

is an MCMC method where a **sample from a multivariate distribution** is obtained by repeatedly drawing from each of the univariate full conditional distributions instead.

2. Imputation step 2.3. FCS/MICE

MICE (Multiple Imputation using Chained Equations) or FCS (multiple imputation using Fully Conditional Specification)

extends univariable imputation to the setting with multivariate non-monotone missingness:

MICE/FCS

- imputes multivariate missing data on a variable-by-variable basis,
- using the technique for univariable missing data.

2. Imputation step 2.3. FCS/MICE

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extends univariable imputation to the setting with multivariate non-monotone missingness:

MICE/FCS

- imputes multivariate missing data on a variable-by-variable basis,
- using the technique for univariable missing data.

Moreover, MICE/FCS is

- an iterative procedure, specifically
- a Markov Chain Monte Carlo (MCMC) method,
- uses the idea of the Gibbs sampler, and
- is a Gibbs sampler if the conditional distributions are compatible (we will come back to this)

Notation

- X: $n \times p$ data matrix with n rows and p variables x_1, \ldots, x_p
- R: $n \times p$ missing indicator matrix containing 0 (missing) or 1 (observed)

$$\mathbf{X} = \begin{bmatrix} X_{-2} & X_2 & X_{-2} \\ X_{1,1} & X_{1,2} & \dots & X_{1,p} \\ X_{2,1} & X_{2,2} & \dots & X_{2,p} \\ \vdots & \vdots & \ddots & \vdots \\ X_{n,1} & X_{n,2} & \dots & X_{n,p} \end{bmatrix} \qquad \mathbf{R} = \begin{bmatrix} R_{1,1} & R_{1,2} & \dots & R_{1,p} \\ R_{2,1} & R_{2,2} & \dots & R_{2,p} \\ \vdots & \vdots & \ddots & \vdots \\ R_{n,1} & R_{n,2} & \dots & R_{n,p} \end{bmatrix}$$

$$\mathbf{R} = \begin{vmatrix} R_{1,1} & R_{1,2} & \dots & R_{1,p} \\ R_{2,1} & R_{2,2} & \dots & R_{2,p} \\ \vdots & \vdots & \ddots & \vdots \\ R_{n,1} & R_{n,2} & \dots & R_{n,p} \end{vmatrix}$$

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 \vdots & \vdots & \ddots & \vdots \\
 X_{n,1} & X_{n,2} & \dots & X_{n,p}
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For example:

$$\boldsymbol{X} = \begin{array}{c|cccc} X_1 & X_2 & X_3 & X_4 \\ \hline \checkmark & \mathsf{NA} & \checkmark & \checkmark \\ \checkmark & \checkmark & \mathsf{NA} & \mathsf{NA} \\ \checkmark & \mathsf{NA} & \checkmark & \mathsf{NA} \end{array}$$

$$\Rightarrow R = \begin{vmatrix} 1 & 0 & 1 & 1 \\ 1 & 1 & 0 & 0 \\ 1 & 0 & 1 & 0 \end{vmatrix}$$

$\textbf{Algorithm 1} \ \mathsf{MICE} \ \mathsf{algorithm} \ [17] \ \mathsf{for} \ \textbf{one} \ \mathsf{imputed} \ \mathsf{dataset}$

- 1: **for** j in $1, \ldots, p$: \triangleright Setup
- 2: Specify imputation model for variable X_j $p(X_j^{mis} \mid X_j^{obs}, X_{-j}, R)$
- 3: Fill in starting imputations \dot{X}_{j}^{0} by random draws from X_{j}^{obs} .
- 4: end for

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- 4: end for
- 5: **for** t in 1, ..., T:
- 6: **for** j in 1, ..., p:

- ▷ loop through iterations
- ▷ loop through variables

10: end for

11: end for

Algorithm 1 MICE algorithm [17] for **one** imputed dataset

1: **for** j in 1, ..., p:

⊳ Setup

- 2: Specify imputation model for variable X_j $p(X_j^{mis} \mid X_j^{obs}, X_{-j}, R)$
- 3: Fill in starting imputations \dot{X}_i^0 by random draws from X_i^{obs} .
- 4: end for
- 5: **for** t in 1, ..., T:

▷ loop through iterations

6: **for** j in 1, ..., p:

- ▷ loop through variables
- 7: Define currently complete data except X_j

$$\dot{X}_{-j}^{t} = \left(\dot{X}_{1}^{t}, \dots, \dot{X}_{j-1}^{t}, \dot{X}_{j+1}^{t-1}, \dots, \dot{X}_{p}^{t-1}\right).$$

- 10: end for
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- 8: Draw parameters $\dot{\theta}^t_j \sim p(\theta^t_j \mid X^{obs}_j, \dot{X}^t_{-j}, R)$.
- 10: end for
- 11: end for

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- 4: end for
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 - $\lambda_{-j} = (\lambda_1, \dots, \lambda_{j-1}, \lambda_{j+1}, \dots, \lambda_p)$
- 8: Draw parameters $\dot{\theta}_{j}^{t} \sim p(\theta_{j}^{t} \mid X_{j}^{obs}, X_{-j}^{t}, R)$.
- 9: Draw imputations $\dot{X}_{j}^{t} \sim P(X_{j}^{mis} \mid \dot{X}_{-j}^{t}, R, \dot{\theta}_{j}^{t})$.
- 10: end for
- 11: end for

```
1: for j in 1, ..., p:

    Setup

         Specify imputation model for variable X_i
         p(X_i^{mis} \mid X_i^{obs}, X_{-i}, R)
         Fill in starting imputations \dot{X}_{i}^{0} by random draws from X_{i}^{obs}.
 4: end for
 5: for t = 1:
                                                                        for i=1:
                                                                         6:
              Define currently complete data except X_1
 7:
              \dot{X}_{-1}^{1} = \left(\dot{X}_{2}^{0}, \dot{X}_{3}^{0}, \dot{X}_{4}^{0}\right).
              Draw parameters \dot{\theta}_1^1 \sim p(\theta_1^1 \mid X_1^{obs}, \dot{X}_{-1}^1, R).
 8:
              Draw imputations \dot{X}_1^1 \sim P(X_1^{mis} \mid \dot{X}_1^1, R. \dot{\theta}_1^1).
 9:
         end for
10:
11: end for
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1: for j in 1, ..., p:

    Setup

         Specify imputation model for variable X_i
         p(X_i^{mis} \mid X_i^{obs}, X_{-i}, R)
         Fill in starting imputations \dot{X}_{i}^{0} by random draws from X_{i}^{obs}.
 4: end for
 5: for t = 1:
                                                                        for i = 2:
                                                                         6:
              Define currently complete data except X_2
 7:
              \dot{X}_{-2}^{1} = \left(\dot{X}_{1}^{1}, \dot{X}_{3}^{0}, \dot{X}_{4}^{0}\right).
              Draw parameters \dot{\theta}_2^1 \sim p(\theta_2^1 \mid X_2^{obs}, \dot{X}_{-2}^1, R).
 8:
              Draw imputations \dot{X}_2^1 \sim P(X_2^{mis} \mid \dot{X}_2^1, R, \dot{\theta}_2^1)
 9:
         end for
10:
11: end for
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    Setup

         Specify imputation model for variable X_i
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         Fill in starting imputations \dot{X}_{i}^{0} by random draws from X_{i}^{obs}.
 4: end for
 5: for t = 1:
                                                                       for i = 3:
                                                                        6:
              Define currently complete data except X_3
 7:
             \dot{X}_{-3}^{1} = \left(\dot{X}_{1}^{1}, \dot{X}_{2}^{1}, \dot{X}_{4}^{0}\right).
              Draw parameters \theta_3^1 \sim p(\theta_3^1 \mid X_3^{obs}, X_{-3}^1, R).
 8:
              Draw imputations \dot{X}_3^1 \sim P(X_3^{mis} \mid \dot{X}_3^1, R, \dot{\theta}_3^1).
 9:
         end for
10:
11: end for
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1: for j in 1, ..., p:

    Setup

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         Fill in starting imputations \dot{X}_{i}^{0} by random draws from X_{i}^{obs}.
 4: end for
 5: for t = 1:
                                                                       for i = 4:
                                                                        6:
              Define currently complete data except X_4
 7:
             \dot{X}_{-4}^1 = (\dot{X}_1^1, \dot{X}_2^1, \dot{X}_3^1).
              Draw parameters \dot{\theta}_4^1 \sim p(\theta_4^1 \mid X_4^{obs}, \dot{X}_{-4}^1, R).
 8:
              Draw imputations \dot{X}_{A}^{1} \sim P(X_{A}^{mis} \mid \dot{X}_{A}^{1}, R. \dot{\theta}_{A}^{1})
 9:
         end for
10:
11: end for
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```
1: for j in 1, ..., p:

    Setup

        Specify imputation model for variable X_i
        p(X_i^{mis} \mid X_i^{obs}, X_{-i}, R)
        Fill in starting imputations \dot{X}_{i}^{0} by random draws from X_{i}^{obs}.
 4: end for
 5: for t = 2:
                                                                     for i=1:
                                                                      6:
             Define currently complete data except X_1
 7:
             \dot{X}_{-1}^2 = (\dot{X}_2^1, \dot{X}_3^1, \dot{X}_4^1).
             Draw parameters \dot{\theta}_1^2 \sim p(\theta_1^2 \mid X_1^{obs}, \dot{X}_{-1}^2, R).
 8:
             Draw imputations \dot{X}_1^2 \sim P(X_1^{mis} \mid \dot{X}_1^2, R, \dot{\theta}_1^2)
 9:
         end for
10:
11: end for
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    Setup

         Specify imputation model for variable X_i
         p(X_i^{mis} \mid X_i^{obs}, X_{-i}, R)
         Fill in starting imputations \dot{X}_{i}^{0} by random draws from X_{i}^{obs}.
 4: end for
 5: for t = 2:
                                                                     for i = 2:
                                                                      6:
              Define currently complete data except X_2
 7:
             \dot{X}_{-2}^2 = (\dot{X}_1^2, \dot{X}_3^1, \dot{X}_4^1).
             Draw parameters \dot{\theta}_2^2 \sim p(\theta_2^2 \mid X_2^{obs}, \dot{X}_{-2}^2, R).
 8:
             Draw imputations \dot{X}_2^2 \sim P(X_2^{mis} \mid \dot{X}_2^2, R, \dot{\theta}_2^2)
 9:
         end for
10:
11: end for
```

The imputed values from the last iteration,

$$\left(\dot{X}_1^T,\ldots,\dot{X}_p^T\right),$$

are then used to replace the missing values in the original data.

One run through the algorithm → one imputed dataset.

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→ To obtain *m* imputed datasets: **repeat** *m* **times**

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One run through the algorithm → one imputed dataset.

➡ To obtain *m* imputed datasets: **repeat** *m* **times**

We refer to the **sequence of imputations** for one missing value, from starting value to final iteration, as a **chain**. Each run through the MICE algorithm produces one chain per missing value.

Why iterations?

- Imputed values in one variable depend on the imputed values of the other variables (Gibbs sampling).
- If the starting values (random draws) are far from the actual distribution, imputed values from the first few iterations are not draws from the distribution of interest.

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How many iterations?

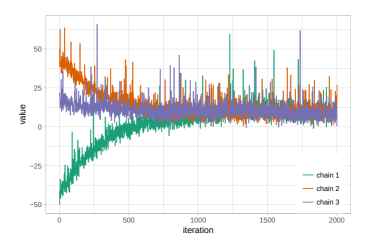
Until convergence

= when the sampling distribution does not change any more (Note: the imputed value will still vary between iterations.)

How to evaluate convergence?

The **traceplot** (x-axis: iteration number, y-axis: imputed value) should show a horizontal band

2. Imputation step 2.4. Checking convergence



Each chain is the sequence of imputed values (from starting value to final imputed value) for the same missing value.

2. Imputation step 2.4. Checking convergence

In imputation we have

- several variables with missing values (e.g., p)
- several missing values in each of these variables
- m chains for each missing value
- → possibly a large number of MCMC chain

To check all chains separately could be very time consuming in large datasets (and storing all iterations from all imputed values is inefficient).

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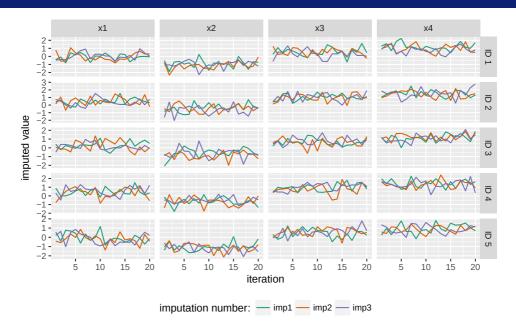
To check all chains separately could be very time consuming in large datasets (and storing all iterations from all imputed values is inefficient).

Alternative: Calculate and plot a summary (e.g., the mean) of the imputed values over all subjects, separately per chain and variable

 \rightarrow only $m \times p$ chains to check

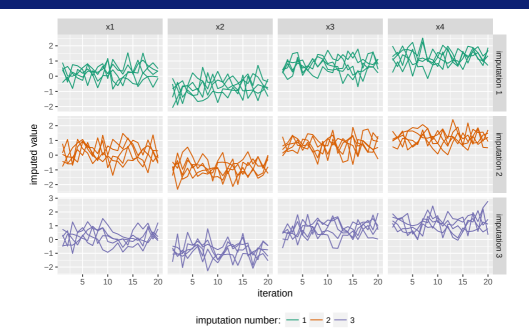
2. Imputation step

2.4. Checking convergence



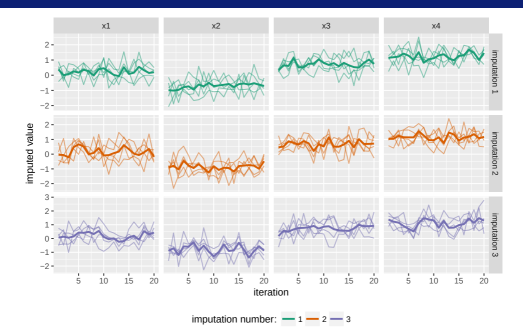
2. Imputation step

2.4. Checking convergence

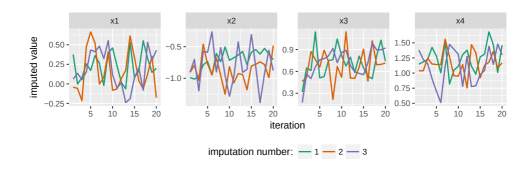


2. Imputation step

2.4. Checking convergence



2. Imputation step 2.4. Checking convergence



3. Analysis step

Multiple imputed datasets:

X_1	X_2	X_3	X_4
1.4	9.2	1.8	2.0
0.5	12.4	2.3	0.1
-0.5	10.7	2.6	-1.6
:	:	:	:

X_1	X_2	X_3	X_4
1.4	13.3	1.8	2.0
0.5	12.4	2.1	0.6
-0.5	10.2	2.6	-1.7
	:	:	:

X_1	X_2	X_3	X_4
1.4	10.0	1.8	2.0
0.5	12.4	2.2	-1.4
-0.5	8.6	2.6	-1.0
:	:	÷	:

3. Analysis step

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$$\begin{array}{c|ccccc} X_1 & X_2 & X_3 & X_4 \\ \hline 1.4 & 13.3 & 1.8 & 2.0 \\ 0.5 & 12.4 & 2.1 & 0.6 \\ -0.5 & 10.2 & 2.6 & -1.7 \\ \vdots & \vdots & \vdots & \vdots \end{array}$$

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Analysis model of interest, e.g.,

$$x_1 = \beta_0 + \beta_1 x_2 + \beta_2 x_3 + \beta_3 x_4$$

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Analysis model of interest, e.g.,

$$x_1 = \beta_0 + \beta_1 x_2 + \beta_2 x_3 + \beta_3 x_4$$

Multiple sets of results:

	est.	se
β_0	0.35	0.21
eta_{1}	0.14	0.02
β_2	-0.64	0.03
eta_3	0.18	0.03

	est.	se
β_0	0.44	0.23
β_1	0.12	0.01
β_2	-0.61	0.03
eta_3	0.22	0.03

	est.	se
β_0	0.19	0.21
β_1	0.14	0.01
β_2	-0.59	0.03
eta_3	0.2	0.03

4. Pooling 4.1. Why pooling?

Recall from slide 6:

We need to represent missing values by a number of imputations.

→ *m* imputed datasets

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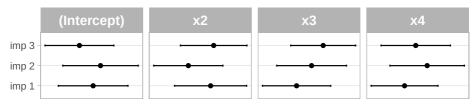
From the different imputed datasets we get **different sets of parameter estimates**, each of them with a standard error, representing the uncertainty about the estimate.

We want to summarize the results and describe how (much) the results vary between the imputed datasets.

4. Pooling 4.1. Why pooling?

In the results from multiply imputed data there are **two types of variation/uncertainty**:

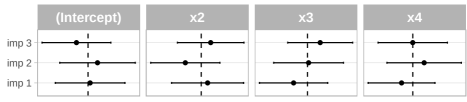
- within imputation (represented by the confidence intervals)
- between imputation (horizontal shift between imputations)



parameter estimate & 95% confidence interval

4. Pooling4.1. Why pooling?

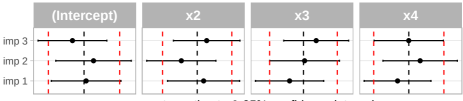
To summarize the results, we can take the mean of the results from the separate analyses. This is the **pooled point estimate**.



parameter estimate & 95% confidence interval

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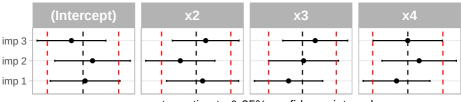


parameter estimate & 95% confidence interval

But does the same work for the std. error (or bounds of the CIs)?

4. Pooling 4.1. Why pooling?

To summarize the results, we can take the mean of the results from the separate analyses. This is the **pooled point estimate**.



parameter estimate & 95% confidence interval

But does the same work for the std. error (or bounds of the CIs)?

The averaged CI's (marked in red) seem to underestimate the total variation (within + between).

4. Pooling 4.2. Rubin's Rules

The most commonly used method to pool results from analyses of multiply imputed data was introduced by Rubin [10], hence **Rubin's Rules**.

Notation:

m: number of imputed datasets

 Q_{ℓ} : quantity of interest (e.g., regr. parameter β) from ℓ -th imputation

 U_{ℓ} : variance of Q_{ℓ} (e.g., $var(\beta) = se(\beta)^2$)

Pooled parameter estimate:

$$ar{Q} = rac{1}{m} \sum_{\ell=1}^m \hat{Q}_\ell$$

4. Pooling 4.2. Rubin's Rules

The variance of the pooled parameter estimate is calculated from the within and between imputation variance.

Average within imputation variance:

$$ar{U} = rac{1}{m} \sum_{\ell=1}^m ar{U}_\ell$$

Between imputation variance:

$$B = rac{1}{m-1} \sum_{\ell=1}^m \left(\hat{Q}_\ell - ar{Q}
ight)^T \left(\hat{Q}_\ell - ar{Q}
ight)$$

Total variance:

$$T = \bar{U} + B + B/m$$

Confidence intervals for pooled estimates can be obtained using the **pooled** standard error \sqrt{T} and a reference t distribution with degrees of freedom

$$\nu = (m-1)(1+r_m^{-1})^2$$
,

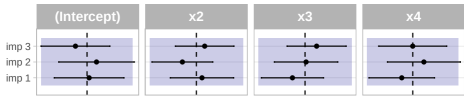
where $r_m = \frac{(B+B/m)}{\bar{U}}$ is the relative increase in variance that is due to the missing values.

The $(1-\alpha)$ 100% confidence interval is then

$$\bar{Q} \pm t_{\nu}(\alpha/2)\sqrt{T}$$

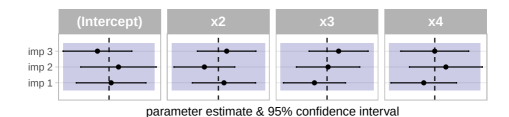
where t_{ν} is the $\alpha/2$ quantile of the t distribution with ν degrees of freedom.

4. Pooling4.2. Rubin's Rules



parameter estimate & 95% confidence interval

4. Pooling 4.2. Rubin's Rules



The corresponding **p-value** is the probability

$$Pr\left\{F_{1,\nu}>\left(Q_{0}-\bar{Q}_{m}\right)^{2}/T\right\},$$

where $F_{1,\nu}$ is a random variable that has an F distribution with 1 and ν degrees of freedom, and Q_0 is the null hypothesis value (typically zero).

Quiz

To reiterate the content of the above sections, you can take the corresponding quiz. An interactive version can be found at

https://emcbiostatistics.shinyapps.io/MICourse_Quiz_PartI

or you can download an html version from Canvas (Files > Principal documents > Multiple Imputation > Quiz_PartI_static.html).

5.1. Bayesian multiple imputation

The imputation step consists itself of two (or three) steps:

- 0. Specification of the imputation model,
- 1. estimation or sampling of the parameters, and
- 2. drawing imputed values from the predictive distribution.

5.1. Bayesian multiple imputation

The imputation step consists itself of two (or three) steps:

- 0. Specification of the imputation model,
- 1. estimation or sampling of the parameters, and
- 2. drawing imputed values from the predictive distribution.

Notation:

Let y be the incomplete covariate to be imputed, and X the design matrix of other (complete or imputed) variables.

5.1. Bayesian multiple imputation

In the Bayesian framework, everything unknown or unobserved is considered as a random variable. Here, this includes for example regression coefficients β , residual variance σ^2 and missing values \mathbf{y}_{mis} and \mathbf{X}_{mis} .

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Random variables have a **probability distribution**. The **expectation** of that distribution quantifies where which **values** of the random variable are **most likely**, the **variance** is a measure of the **uncertainty** about the values.

5.1. Bayesian multiple imputation

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Random variables have a **probability distribution**. The **expectation** of that distribution quantifies where which **values** of the random variable are **most likely**, the **variance** is a measure of the **uncertainty** about the values.

In Bayesian imputation, the information obtained from the observed data is used to estimate the probability distributions for the missing values and unknown parameters, and values are imputed by draws from that posterior (= after having seen the data) distribution.

5.1. Bayesian multiple imputation

To determine the **expectation** of the posterior distribution of the missing values, usually a **regression model** is used, that depends on the unknown coefficients β .

$$\mathbb{E}(\boldsymbol{y}_{mis} \mid \boldsymbol{X}, \boldsymbol{\beta}) = f(\boldsymbol{X}_{mis}\boldsymbol{\beta})$$

The posterior distribution of $\boldsymbol{\beta}$ and σ , $p(\boldsymbol{\beta}, \sigma \mid \boldsymbol{y}_{obs} \mid \boldsymbol{y}_{obs}, \boldsymbol{X}_{obs})$, is estimated from the corresponding regression model on the observed data.

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To impute missing values, while taking into account the uncertainty about β and σ , the estimated posterior distributions of the missing values and parameters are multiplied

$$p(\mathbf{y}_{mis} \mid \mathbf{X}_{mis}, \boldsymbol{\beta}, \boldsymbol{\sigma}) p(\boldsymbol{\beta}, \boldsymbol{\sigma} \mid \mathbf{y}_{obs}, \mathbf{X}_{obs})$$

In practice, this can be implemented by first making a draw from the posterior distributions of β and σ , and plugging the values into the distribution of \mathbf{y}_{obs} .

5.1. Bayesian multiple imputation

Example: We assume that y given X is approximately normal.

Then $p(\mathbf{y}_{mis} \mid \mathbf{X}_{mis}, \boldsymbol{\beta}, \boldsymbol{\sigma})$ is a normal distribution and we can

- draw $\tilde{\boldsymbol{\beta}}$ from $p(\boldsymbol{\beta} \mid \boldsymbol{y}_{obs}, \boldsymbol{X}_{obs})$,
- draw $\tilde{\sigma}$ from $p(\sigma \mid \boldsymbol{y}_{obs}, \boldsymbol{X}_{obs})$,
- draw $\tilde{\mathbf{y}}_{mis}$ from a normal distribution with mean (= expectation) $\mathbf{X}\tilde{\boldsymbol{\beta}}$ and variance $\tilde{\sigma}^2$.

This is actually the approach we have seen previously on Slides 12/13 and 19.

5.2. Bootstrap multiple imputation

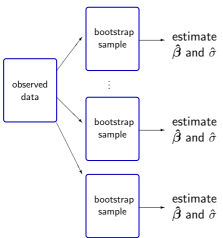
An alternative approach is to capture the uncertainty with **bootstrap** sampling.

In empirical **Bootstrap**, (many) replications of the data are created by repeatedly drawing values from the original data.

5.2. Bootstrap multiple imputation

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In empirical **Bootstrap**, (many) replications of the data are created by repeatedly drawing values from the original data.



Bootstrap samples can contain some observations multiple times and some observations not at all.

The statistic of interest is then calculated on each of the bootstrap samples.

5.2. Bootstrap multiple imputation

In bootstrap multiple imputation,

- one bootstrap sample of the observed data is created per imputation,
- the least-squares estimates of the parameters are calculated from

$$\mathbf{y}_{obs} = \mathbf{X}_{obs} \boldsymbol{\beta} + \varepsilon_{obs} \ \hat{\boldsymbol{\beta}} \quad \hat{\boldsymbol{\phi}}$$
 (step 1).

• Imputed values are sampled from $p(\boldsymbol{y}_{mis} \mid \boldsymbol{X}_{mis}, \boldsymbol{\hat{\beta}}, \hat{\sigma})$ (step 2).

5.2. Bootstrap multiple imputation

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- one bootstrap sample of the observed data is created per imputation,
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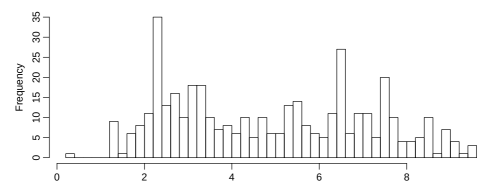
Analogous to Bayesian multiple imputation, for a normal imputation model, p() is the normal distribution and

$$ilde{m{y}}_{mis} = m{X}_{mis}m{\hat{eta}} + ilde{arepsilon}$$

where $\tilde{\varepsilon}$ is drawn independently from $N(0, \hat{\sigma}^2)$.

Both Bayesian and bootstrap multiple imputation sample imputed values from a distribution p() in step 2.

Sometimes, the empirical distribution can not be adequately approximated by a known probability distribution.



Predictive Mean Matching (PMM) was developed to provide a semi-parametric approach to imputation for settings where the normal distribution is not a good choice for the predictive distribution.[8, 9]

The idea is to find cases in the observed data that are similar to the cases with missing values and to fill in the missing value with the observed value from one of those cases.

To find similar cases, the predicted values of complete and incomplete cases are compared.

5.3. Semi-parametric imputation

The steps in PMM:

- 1. Obtain parameter estimates for $\hat{\beta}$ and $\hat{\sigma}$ (see later)
- 2. Calculate the predicted values for the observed data

$$\hat{m{y}}_{obs} = m{X}_{obs}\hat{m{eta}}$$

3. Calculate the predicted value for the incomplete data

$$\hat{m{y}}_{mis} = m{X}_{mis} \hat{m{eta}}$$

- 4. For each missing value, find *d* donor candidates that fulfill a given criterium (details on the next slide).
- 5. Randomly select one of the donors.

5.3. Semi-parametric imputation

Several criteria to select donors have been proposed:

1. The donor is the **(one)** case with the smallest absolute difference $|\hat{y}_{mis,i} - \hat{y}_{obs,j}|$, j = 1, ..., q.

5.3. Semi-parametric imputation

Several criteria to select donors have been proposed:

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- 2. Donor candidates are the d cases with the smallest absolute difference $|\hat{y}_{mis,i} \hat{y}_{obs,j}|$, $j = 1, \ldots, q$. The donor is selected randomly from the candidates.
- 3. Donor candidates are those cases for which the absolute difference is smaller than some limit $\eta\colon |\hat{y}_{mis,i} \hat{y}_{obs,j}| < \eta, \ j=1,\dots,q$. The donor is selected randomly from the candidates.

5.3. Semi-parametric imputation

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- 2. Donor candidates are the d cases with the smallest absolute difference $|\hat{y}_{mis,i} \hat{y}_{obs,j}|$, $j = 1, \ldots, q$. The donor is selected randomly from the candidates.
- 3. Donor candidates are those cases for which the **absolute difference is** smaller than some limit η : $|\hat{y}_{mis,i} \hat{y}_{obs,j}| < \eta, \ j = 1, \dots, q$. The donor is selected randomly from the candidates.
- 4. Select candidates like in 2. or 3., but select the donor from the candidates with probability that depends on $|\hat{y}_{mis,i} \hat{y}_{obs,j}|$.[16]

- Selection criteria 2. 4., require the number of candidates d (or maximal difference η) to be specified. Common choices for d are 3, 5 or 10.
- If the same donor is chosen in many/all imputations (e.g., because only a few similar observed cases are available), the uncertainty about the missing values will be underestimated.

- Selection criteria 2. 4., require the number of candidates d (or maximal difference η) to be specified. Common choices for d are 3, 5 or 10.
- If the same donor is chosen in many/all imputations (e.g., because only a few similar observed cases are available), the uncertainty about the missing values will be underestimated.
 - → PMM may be **problematic** when
 - the dataset is very small,
 - the proportion of missing values is large, or
 - one/some predictor variable(s) are strongly related to the missingness.

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 - one/some predictor variable(s) are strongly related to the missingness.
- Therefore, using d = 1 (selection criterion 1.) is not a good idea. On the other hand, using too many candidates can lead to bad matches.
- Schenker and Taylor [15] proposed an adaptive procedure to select *d*, but it is not used much in practice.

For the **sampling of the parameters** (step 1 on slide 43), different approaches have been introduced in the literature:

- Type-0 point estimates $\hat{\beta}$ are used in both prediction models (least squares or maximum likelihood)
- Type-I $\hat{\beta}$ to predict \hat{y}_{obs} ; $\tilde{\beta}$ to predict \hat{y}_{mis} is sampled from the posterior distribution of β (Bayesian) or bootstrapped
- Type-II $\tilde{\beta}$ to predict \hat{y}_{obs} as well as \hat{y}_{mis}
- Type-III different draws $\tilde{\beta}^{(1)}$ and $\tilde{\beta}^{(2)}$ to predict \hat{y}_{obs} and \hat{y}_{mis} , respectively

The use of point estimates (Type-0 and Type-I matching) underestimates the uncertainty about the regression parameters.

Another point of consideration is the choice of the set of data used to train the prediction models.

In the version presented on slide 43, the same set of data (all cases with observed y) is used to train the model and to produce predicted values of y_{obs} .

The predictive model will likely fit the observed cases better than the missing cases, and, hence, variation will be underestimated.

As an alternative, the **model could be trained on the whole data** (using previously imputed values) or to use a **leave-one-out approach** on the observed data.

5.4. What is implemented in software?

```
mice (in R):
```

- PMM via mice.impute.pmm()
 - specification of number of donors d (same for all variables)
 - Type-0, Type-I, Type-II matching
- PMM via mice.impute.midastouch()
 - allows leave-one-out estimation of the parameters
 - distance based donor selection
 - Type-0, Type-I, Type-II matching
- bootstrap linear regression via mice.impute.norm.boot()
- bootstrap logistic regression via mice.impute.logreg.boot()
- Bayesian linear regression via mice.impute.norm()
- . . .

Summary of Part I

1. What is Multiple Imputation?

- Rubin's two ideas:
 - Missing values need to be represented by multiple imputed values.
 - A model is necessary to obtain good imputations.
- Imputed values are obtained from the **predictive distribution** of the missing data, given the observed data.
- Multiple completed datasets are created from the multiple imputed values.
- Multiple imputation has three steps: Imputation, analysis, pooling

Summary of Part I (cont.)

2. Imputation step

- Two sources of variation need to be taken into account
 - parameter uncertainty
 - random variation
- Two approaches to MI for imputation of non-monotone multivariate missing data
 - MICE/FCS
 - Joint model imputation
- The MICE algorithm re-uses univariate imputation models by iterating through all incomplete variables, multiple times (iterations)
- Multiple runs through the algorithm are necessary to create multiple imputed dataset
- The **convergence of the chains** needs to be checked.

Summary of Part I (cont.)

3. Analysis step

 Analyse each imputed dataset the way you would analyse a complete dataset

4. Pooling

- Results from analyses of multiple imputed datasets can be summarized by taking the average of the regression coefficients
- For the total variance, two sources of variation need to be considered:
 - within imputation variance
 - between imputation variance

Summary of Part I (cont.)

5. A closer look at the imputation step

- Two parametric approaches for imputation:
 - Bayesian (sample from posterior distribution of parameters)
 - Bootstrap (uses bootstrap samples of the data to estimate parameters)
- Predictive mean matching is a semi-parametric alternative (it matches observed and missing cases based on their predicted values).
- In PMM we need to consider
 - donor selection
 - matching type (how parameters are sampled/estimated),
 - the **set of data** used to calculate/estimate the parameters.
- Bayesian and bootstrap imputation take into account the variation, while many choices in PMM lead to underestimation of the variation.

Part II Multiple Imputation Workflow

Outline of Part II

- 6. Know your data
 - 6.1 Missing data patterns
 - 6.2 Data distributions
 - 6.3 Correlations & patterns
 - 6.4 Why are values missing?
 - 6.5 Auxiliary variables
- 7. Imputation with mice()
 - 7.1 Main function arguments
 - 7.2 Imputation methods
 - 7.3 Predictor matrix
 - 7.4 Passive imputation
 - 7.5 Post processing
 - 7.6 Visit sequence
 - 7.7 Good to know

- 8. Convergence & Diagnostics
 - 8.1 Logged events
 - 8.2 Convergence
 - 8.3 Diagnostics
- 9. Analyse & pool the imputed data
 - 9.1 Analysing imputed data
 - 9.2 Pooling results
 - 9.3 Functions for pooled results
- 10. Additional functions in mice()
 - 10.1 Extract & export imputed data
 - 10.2 Combining mids objects
 - 10.3 Adding variables to mids objects
- 11. Multiple Imputation in SPSS
 - 11.1 Where to get help
 - 11.2 Multiple Imputation Features

6. Know your data 6.1. Missing data patterns

There are several packages in R that provide functions to create and **plot the missing data pattern**.

Examples are:

mice, VIM, Amelia, visdat, naniar, ...

To create the plots on the following slides, we need to load the example data

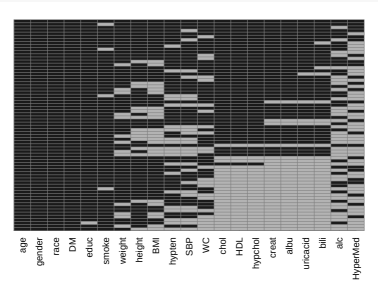
```
datadir <- # fill in path to folder containing the data
load(file.path(datadir, "NHANES.RData"))</pre>
```

6.1. Missing data patterns

```
mdp <- mice::md.pattern(NHANES)</pre>
head(mdp[, -c(7:14)]) # omit some columns to fit it on the slide
##
        age gender race DM educ smoke hypchol creat albu uricacid bili alc HyperMed
    572
                                                                                       1 0
##
    141
##
##
    17
                                                                                       1 1
   1063
                                                                                       0 1
##
     18
                                                                                       1 1
tail(mdp[, -c(7:14)])
##
        age gender race DM educ smoke hypchol creat albu uricacid bili alc HyperMed
##
                                              0
                                                                             0
                                                                                           11
##
                                                                                           12
##
                                                                                           12
##
                                                                                           12
##
                                              0
                                                                                           14
##
                                            175
                                                   184
                                                        184
                                                                  185
                                                                       188 627
                                                                                   1606 3975
```

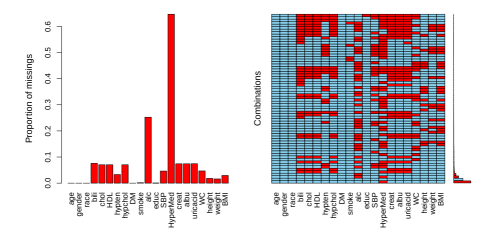
6.1. Missing data patterns

```
par(mar = c(5, 0.5, 1, 3), mgp = c(2, 0.6, 0))
JointAI::md_pattern(NHANES, print = F, printN = F, yaxis_pars = list(yaxt = 'n'))
```



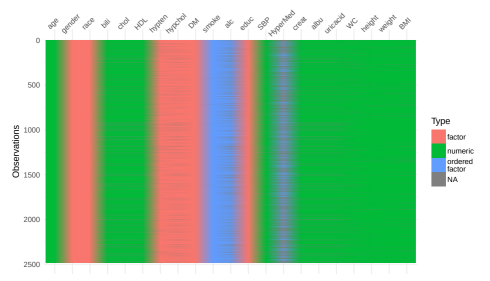
6.1. Missing data patterns

```
par(mar = c(6, 3, 2, 1))
VIM::aggr(NHANES, prop = T, numbers = FALSE)
```



6. Know your data 6.1. Missing data patterns

visdat::vis_dat(NHANES, sort_type = FALSE)



6. Know your data 6.1. Missing data patterns

visdat::vis_miss(NHANES)



6. Know your data 6.1. Missing data patterns

```
# number and proportion of complete cases
Ncc <- cbind(
   "#" = table(complete.cases(NHANES)),
   "%" = round(100 * table(complete.cases(NHANES))/nrow(NHANES), 2)
)
rownames(Ncc) <- c("incompl.", "complete")</pre>
```

6. Know your data 6.1. Missing data patterns

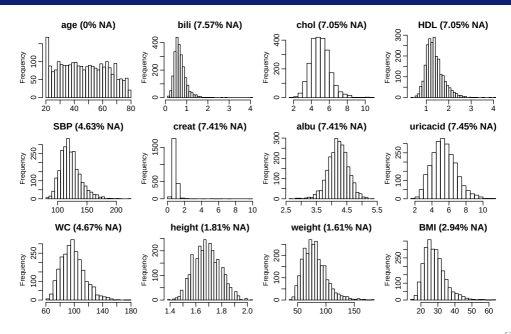
Number and proportion of (in)complete cases

```
## # %
## incompl. 1911 76.96
## complete 572 23.04
```

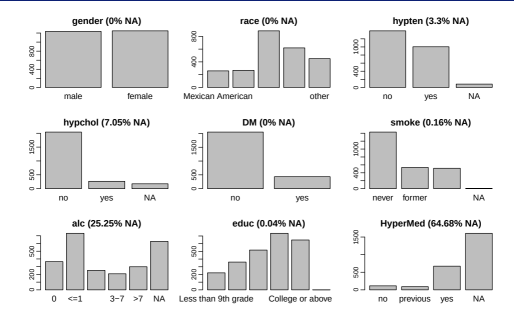
Number and proportion of missing values per variable

```
## # NA % NA
## WC 116 4.67
## chol 175 7.05
## HDL 175 7.05
## hypchol 175 7.05
## creat 184 7.41
## albu 184 7.41
## uricacid 185 7.45
## bili 188 7.57
## alc 627 25.25
## HyperMed 1606 64.68
```

6.2. Data distributions



6.2. Data distributions



6.2. Data distributions

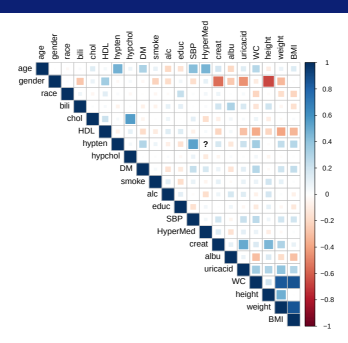
```
# syntax for continuous variables
NHANESnum <- NHANES[, sapply(NHANES, is.numeric)]</pre>
par(mfrow = c(3, 4), mar = c(3, 3.2, 0.5, 0.5), mgp = c(2, 0.6, 0))
for (i in 1:ncol(NHANESnum)) {
 hist(NHANESnum[, i], nclass = 30, xlab = names(NHANESnum)[i], main = "")
 legend("topright", bty = "n",
         legend = paste0(round(mean(is.na(NHANESnum[, i]))*100, 2), "% NA"))
# syntax for factors
NHANESfac <- NHANES[, sapply(NHANES, is.factor)]</pre>
par(mfrow = c(3, 5), mar = c(3, 3.2, 2.5, 0.5), mgp = c(2, 0.6, 0))
for (i in 1:ncol(NHANESfac)) {
 tab <- table(NHANESfac[, i], exclude = NULL)
 names(tab)[is.na(names(tab))] <- "NA"</pre>
  barplot(tab, main = paste0(names(NHANESfac)[i]," (",
                              round(mean(is.na(NHANESfac[, i]))*100,
                                    2), "% NA)"))
```

6. Know your data 6.3. Correlations & patterns

A quick (and dirty) way to check for strong correlations between variables is:

Note: We only use the correlation coefficient for binary variables in this visualization, not as a statistical result!

6. Know your data 6.3. Correlations & patterns



6. Know your data 6.3. Correlations & patterns

Check out what the problem is with hypertension and HyperMed:

6. Know your data 6.4. Why are values missing?

Knowing your data also means to be able to answer these questions:

- Do missing values in multiple variables always occur together?
 (e.g. blood measurements)
- Are there **structural missing values**? (e.g. pregnancy status in men)
- Are there patterns in the missing values?
 (e.g. only patients with hypertension have observations of HyperMed)
- Are values missing by design?
- Is the assumption of ignorable missingness (MAR or MCAR) justifiable?

6. Know your data 6.5. Auxiliary variables

Auxiliary variables are variables that are not part of the analysis but **can help during imputation**.

Good auxiliary variables

- are related to the probability of missingness in a variable, or
- are related to the incomplete variable itself,
- do not have many missing values themselves and
- are (mostly) **observed** when the incomplete variable of interest is missing.

7. Imputation with mice() 7.1. Main function arguments

The main arguments needed to impute data with mice() are:

- data: the dataset
- m: number of imputed datasets (default is 5)
- maxit: number of iterations (default is 5)
- method: vector of imputation methods
- defaultMethod: vector of default imputation methods for numerical, binary, unordered and ordered factors with > 2 levels (default is c("pmm", "logreg", "polyreg", "polr"))
- predictorMatrix: matrix specifying roles of variables

mice has implemented many **imputation methods**, the most commonly used ones are:

- pmm: predictive mean matching (any)
- norm: Bayesian linear regression (numeric)
- logreg: binary logistic regression (binary)
- polr: proportional odds model (ordered factors)
- polyreg: polytomous logistic regression (unordered factors)

Change the default imputation method:

E.g.: to use norm instead of pmm for all continuous incomplete variables, use:

```
mice(NHANES, defaultMethod = c("norm", "logreg", "polyreg", "polr"))
```

Change the default imputation method:

E.g.: to use norm instead of pmm for all continuous incomplete variables, use:

```
mice(NHANES, defaultMethod = c("norm", "logreg", "polyreg", "polr"))
```

Change imputation method for a single variable:

To change the imputation method for single variables (but also for changes in other arguments) it is convenient to **do a setup run** of mice() without iterations (maxit = 0) and to extract and modify the parameters from there.

Change the default imputation method:

E.g.: to use norm instead of pmm for all continuous incomplete variables, use:

```
mice(NHANES, defaultMethod = c("norm", "logreg", "polyreg", "polr"))
```

Change imputation method for a single variable:

To change the imputation method for single variables (but also for changes in other arguments) it is convenient to **do a setup run** of mice() without iterations (maxit = 0) and to extract and modify the parameters from there.

Exclude variable from imputation:

When a variable that has missing values should not be imputed, the method needs to be set to "".

7.2. Imputation methods

```
library(mice)
imp0 <- mice(NHANES, maxit = 0)</pre>
meth <- imp0$method
meth
                                                                 HDL
##
                 gender
                                          bili
                                                     chol
         age
                              race
           11.11
                      11.11
                                         "pmm"
##
                                 11.11
                                                   "mmq"
                                                               "pmm"
##
      hypten
               hypchol
                                 DM
                                         smoke
                                                      alc
                                                                educ
    "logreg" "logreg"
                                 11.11
                                        "polr"
                                                  "polr" "polyreg"
##
##
         SBP HyperMed
                                         albu
                                                uricacid
                                                                  WC
                             creat
       "pmm" "polr"
##
                             "pmm"
                                        "pmm"
                                                   "pmm"
                                                               "pmm"
##
      height weight
                                BMT
       "pmm"
                "mmm"
                             "pmm"
##
meth["albu"] <- "norm"</pre>
meth["HyperMed"] <- ""</pre>
# imp <- mice(NHANES, method = meth)</pre>
```

7.3. Predictor matrix

The predictorMatrix is a matrix that specifies which variables are used as predictors in which imputation model.

Each row represents the model for the variable given in the rowname.

Variables **not used as predictor** are (or have to be set to) **zero**.

By **default**, **all variables** (except the variable itself) **are used** as predictor. For complete variables all entries are 0.

Important:

A variable that has **missing values needs to be imputed** in order to be used as predictor for other imputation models!!!

Note:

By default, **ALL** variables with missing values are imputed and **ALL** variables are used as predictor variables.

- → Make sure to adjust the predictorMatrix and method to avoid using ID variables or other columns of the data that should not be part of the imputation.
- → Make sure all variables are coded correctly, so that the chosen imputation models are appropriate (e.g., ordered factors).

7. Imputation with mice() 7.3. Predictor matrix

7. Imputation with mice() 7.4. Passive imputation

In some cases, variables are functions of other variables, e.g., $BMI = \frac{weight}{height^2}$.

If we impute BMI directly, its values may be **inconsistent** with the (imputed) values of height and weight.

The imputed values of BMI are impossible given the corresponding values of height and weight.

7. Imputation with mice() 7.4. Passive imputation

Moreover, if some components of a variable are observed we want to use that **information to reduce uncertainty**.

```
table(weight_missing = is.na(NHANES$weight),
    height_missing = is.na(NHANES$height))

## height_missing
## weight_missing FALSE TRUE
## FALSE 2410 33
## TRUE 28 12
```

Here we have 33 + 28 = 61 cases in which either height or weight is observed.

We would like to impute height and weight separately and calculate BMI from the (imputed) values of the two variables.

7. Imputation with mice() 7.4. Passive imputation

If BMI is not a relevant predictor in any of the other imputation models, we could just exclude BMI from the imputation and re-calculate it afterwards.

To use BMI as predictor in the imputation, it has to be **calculated in each iteration** of the algorithm. In **mice** this is possible with **passive imputation**.

7. Imputation with mice() 7.4. Passive imputation

If BMI is not a relevant predictor in any of the other imputation models, we could just exclude BMI from the imputation and re-calculate it afterwards.

To use BMI as predictor in the imputation, it has to be **calculated in each iteration** of the algorithm. In **mice** this is possible with **passive imputation**.

Instead of using a standard imputation method, we can specify a formula to calculate BMI:

```
meth["BMI"] <- "~I(weight/height^2)"  # formula to impute BMI
pred[c("weight", "height"), "BMI"] <- 0  # prevent feedback</pre>
```

To prevent feedback from BMI in the imputation of height and weight the predictorMatrix needs to be modified.

7. Imputation with mice() 7.4. Passive imputation

Since BMI depends on weight, and the two variables are highly correlated (ρ =0.87) it may be beneficial **not to use them simultaneously** as predictors in the other imputation models.

Which one to use may differ between imputation models.

Passive imputation can also be useful in settings where

- imputation models include an **interaction terms** between incomplete variables (see [17, p. 133] for an example), or when
- a number of covariates is used to form a sum score. The sum score, instead of all single elements, can then be used as predictor in other imputation models.

7. Imputation with mice() 7.5. Post processing

mice() has an argument post that can be used to specify functions that modify imputed values.

Helpful functions are

- squeeze() to censor variables at given boundaries
- ifdo() for conditional manipulation (not yet implemented)

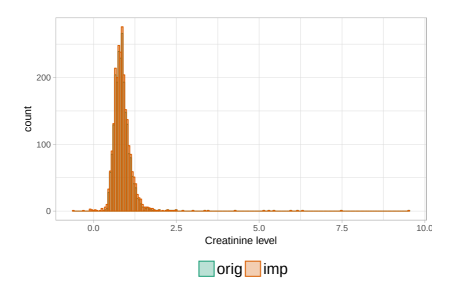
Example:

When inspecting the imputed values from imp, we find that some imputed values in creat are negative.

```
# DF1 is the first imputed dataset we extracted earlier
summary(DF1$creat)

## Min. 1st Qu. Median Mean 3rd Qu. Max.
## -0.6155 0.7000 0.8300 0.8853 0.9900 9.5100
```

7.5. Post processing



7. Imputation with mice() 7.5. Post processing

With the following syntax all imputed values of creat that are outside the interval c(0, 100) will be set to those limiting values.

```
post <- imp$post
post["creat"] <- "imp[[j]][,i] <- squeeze(imp[[j]][,i], c(0, 100))"
imp2 <- update(imp, post = post, maxit = 20, seed = 123)</pre>
```

Note:

When many observations are outside the limits it may be better to **change the imputation model** since the implied **assumption of the imputation model** apparently **does not fit the** (assumption about the) **complete data distribution**.

7. Imputation with mice() 7.5. Post processing

This post-processing of imputed values allows for many more data manipulations and is not restricted to squeeze() (and ifdo()).

Any strings of R commands provided will be evaluated after the corresponding variable is imputed, within each iteration.

For example, if subjects with SBP > 140 should be classified as hypertensive:

```
post["hypten"] <- "imp[[j]][p$data[where[, j], 'SBP'] > 140, i] <- 'yes'"</pre>
```

This also allows for (some) MNAR scenarios, for example, by multiplying or adding a constant to the imputed values or to re-impute values, depending on their current value.

7. Imputation with mice() 7.6. Visit sequence

When the **post-processed or passively imputed values** of a variable depend on other variables, the **sequence in which the variables are imputed** may be important to obtain **consistent values**.

Example:

If BMI is passively imputed (calculated) before the new imputations for height and weight are drawn, the resulting values of BMI, will match height and weight from the **previous iteration**, but not the iteration given in the imputed dataset.

In mice() the argument visitSequence specifies in which order the columns of the data are imputed. By default mice() imputes in the order of the columns in data.

7. Imputation with mice() 7.6. Visit sequence

```
visitSeq <- imp2$visitSequence</pre>
visitSeq
##
       bili
                chol
                          HDL
                                        hypchol smoke
                                                               alc
                                hypten
##
                            6
                                                       10
                                                                11
                 SBP HyperMed creat albu uricacid
##
       educ
                                                                WC
         12
                  13
                           14
                                              16
##
                                    15
                                                       17
                                                                18
##
     height
              weight
                          BMI
##
         19
                  20
                           21
```

Currently, hypten is imputed before SBP, but the imputed values of hypten are post-processed depending on the current value of SBP. To get consistent values of these two variables, we need to change the visitSequence.

7. Imputation with mice() 7.6. Visit sequence

```
visitSeq <- c(visitSeq[-which(names(visitSeq) == "hypten")],</pre>
             visitSeq["hypten"])
visitSeq
      bili
##
               chol
                        HDL
                             hypchol smoke
                                                  alc
                                                         educ
##
                          6
                                          10
                                                   11
                                                           12
       SBP HyperMed creat albu uricacid
                                                  WC
##
                                                       height
##
        13
                 14
                         15
                                  16
                                          17
                                                   18
                                                           19
##
    weight
               BMI
                     hypten
##
        20
                 21
```

The visitSequence may specify that a column is visited multiple times during one iteration. All incomplete variables must be visited at least once.

7. Imputation with mice() 7.6. Visit sequence

```
visitSeq <- c(visitSeq[-which(names(visitSeq) == "hypten")],</pre>
             visitSeq["hypten"])
visitSeq
      bili
##
               chol
                        HDL
                              hypchol smoke
                                                   alc
                                                           educ
##
                         6
                                            10
                                                    11
                                                             12
       SBP HyperMed creat albu uricacid
                                                    WC
##
                                                         height
##
        13
                 14
                          15
                                   16
                                            17
                                                    18
                                                             19
##
    weight BMI
                      hypten
##
        20
                 21
```

The visitSequence may specify that a column is visited multiple times during one iteration. All incomplete variables must be visited at least once.

visitSequence can also be specified using one of the keywords "roman" (left to right), "arabic" (right to left), "monotone" (sorted in increasing amount of missingness), "revmonotone" (reverse of monotone)

7. Imputation with mice() 7.7. Good to know

mice() performs some pre-processing and removes

- incomplete variables that are not imputed but are specified as predictor,
- constant variables, and
- collinear variables.

In each iteration

- linearly dependent variables are removed and
- polr imputation models that do not converge are replaced by polyreg.

Why?

To avoid problems in the imputation models.

7. Imputation with mice() 7.7. Good to know

As a consequence

- imputation models may differ from what the user has specified or assumes is happening, or
- variables that should be imputed are not.
- → Know your data
- Make sure method and predictorMatrix are specified appropriately
- Check the output and log of these automatic actions carefully

7. Imputation with mice() A note

"Please realize that these choices are always needed. Imputation software needs to make default choices. These choices are intended to be useful across a wide range of applications. However, the default choices are not necessarily the best for the data at hand. There is simply no magical setting that always works, so often some tailoring is needed." [17, p. 124]

8. Convergence & Diagnostics 8.1. Logged events

The log of the automatic changes (slide 89) is returned as part of the mids object:

```
head(imp2$loggedEvents)

## it im co dep meth out
## 1 1 1 10 smoke multinom
## 2 1 1 11 alc multinom
## 3 1 1 12 educ multinom
## 4 1 2 10 smoke multinom
## 5 1 2 11 alc multinom
## 6 1 2 12 educ multinom
```

With columns

it	iteration number
im	imputation number
СО	column number in the data
dep	dependent variable
meth	imputation method used
out	names of altered or removed
	predictors

8. Convergence & Diagnostics 8.2. Convergence

Recall from slides 19 and 23:

mice uses an **iterative algorithm** and imputations from the first few iterations may not be samples from the "correct" distributions.

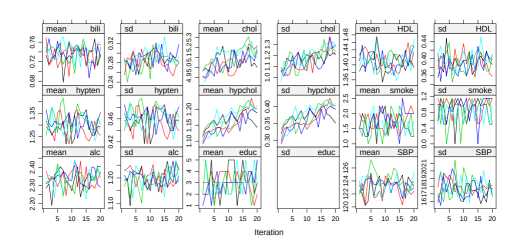
Traceplots can be used to visually assess **convergence**.

In **mice** the function plot() produces traceplots of the mean and standard deviation (across subjects) per incomplete variable (see slide 25).

8. Convergence & Diagnostics

8.2. Convergence

plot(imp2, layout = c(6, 3))



8. Convergence & Diagnostics 8.2. Convergence

The traceplots show that the imputations for chol and hypchol have an upward trend.

Strong trends and traces that show **correlation** between variables indicate **problems of feedback**. This needs to be investigated and resolved in the specification of the predictorMatrix.

Weak trends may be artefacts that often disappear when the imputation is performed with more iterations.

When MCMC chains have converged, the distributions of the imputed and observed values can be compared to investigate differences between observed and imputed data.

Note:

Plots usually show the **marginal** distributions of observed and imputed values, which do not have do be identical under MAR.

Recall:

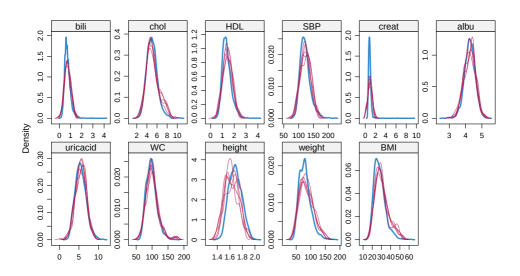
The **conditional** distributions (given all the other variables in the imputation model) of the imputed values are assumed to be the same as the conditional distributions of the observed data.

mice provides several functions for visual diagnosis of imputed values:

- densityplot() (for large datasets and variables with many NAs)
- stripplot() (for smaller datasets and/or variables with few NAs)
- bwplot()
- xyplot()

These functions create lattice graphics, which can be modified analogous to their parent functions from the **lattice** package.

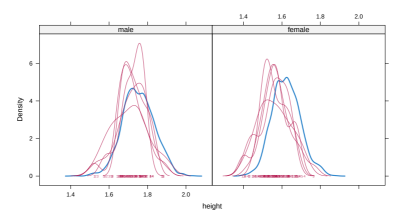
densityplot(imp2)



The densityplot() shows that the distribution of imputed values of creat is wider than the distribution of the observed values and that imputed values of height are smaller than the observed values.

In some cases differences in distributions can be explained by strata in the data, however, here, gender does not explain the difference in observed and imputed values.

densityplot(imp2, ~height|gender, plot.points = T)



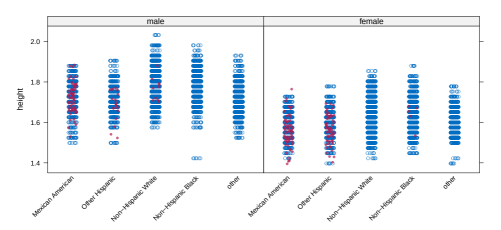
As an alternative, we might consider race to explain the differences

As an alternative, we might consider race to explain the differences

However, there are not enough missing values of height per categories of race to estimate densities.

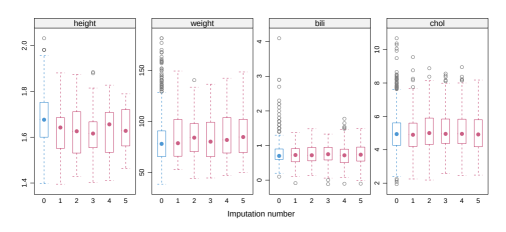
```
with(NHANES, table(race = race, "height missing" = is.na(height)))
##
                     height missing
## race
                     FALSE TRUE
##
    Mexican American
                        233
                             26
##
    Other Hispanic
                   252 16
    Non-Hispanic White 884 2
##
##
    Non-Hispanic Black 618
##
    other
                        451
                              0
```

In that case, a stripplot() may be better suited. Here we can also split the data for gender and race.



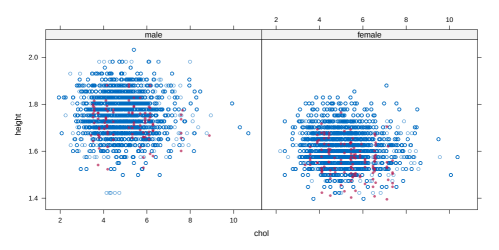
Alternatively, observed and imputed data can be represented by box-and-whisker plots:

bwplot(imp2, height + weight + bili + chol ~.imp)



The function xyplot() allows multivariate investigation of the imputed versus observed values.

```
xyplot(imp2, height ~ chol|gender, pch = c(1,20))
```



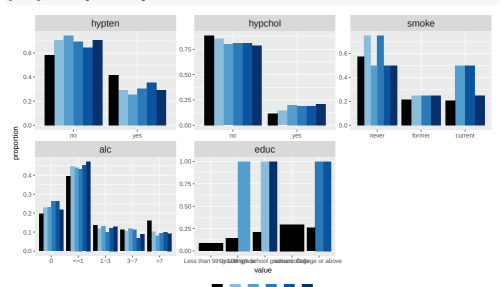
All of the above graphs displayed only continuous imputed variables. For categorical variables we can compare the proportion of values in each category.

mice does not provide a function to do this, but we can write one ourselves, as for instance the function <code>probplot()</code>, for which the syntax can be found here. The function can be loaded into R using:

8. Convergence & Diagnostics

8.3. Diagnostics

probplot(imp2, strip.text = element_text(size = 14))

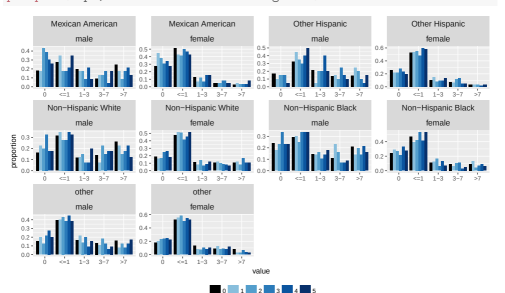


smoke and educ have very few missing values (4 and 1, respectively), so we do not need to worry about differences between observed and imputed data for those variables.

For alc, missing values are imputed by the lower consumption categories more often than we would expect from the observed data, hypten is less frequent and hypchol a bit more frequent, in the imputed data compared to the observed.

If we expect that gender and race might explain the differences for hypten, we can include those factors into the plot.

probplot(imp2, formula = alc ~ race + gender)



Since hypertension is more common in older individuals, we may want to investigate if age can explain the differences in imputed values of hypten.

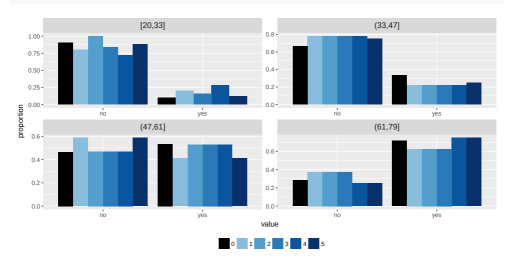
The table shows that the distribution of age in participants with missing hypten is very similar to the distribution of age in participants without hypten.

8. Convergence & Diagnostics

8.3. Diagnostics

Plotting the proportions of observed and imputed <a href="https://hypten.gov/

probplot(imp2, formula = hypten ~ cut(age, quantile(age), include.lowest = T))



9. Analyse & pool the imputed data

9.1. Analysing imputed data

Once we have confirmed that our imputation was successful, we can move on to the **analysis of the imputed data**.

For example, we might be interested in the following logistic regression model:

```
glm(DM ~ age + gender + hypchol + BMI + smoke + alc,
    family = "binomial")
```

To fit the model on each of the imputed datasets, we do not need to extract the data from the mids object, but can use with().

mod1 is an object of class mira.

9. Analyse & pool the imputed data 9.2. Pooling results

Pooled results can be obtained using pool() and its summary.

```
options(width = 90)
res1 <- summary(pool(mod1))</pre>
round(res1, 3)
                                        df Pr(>|t|) lo 95 hi 95 nmis
##
                 est.
                        se
                                 t
                                                                         fmi lambda
  (Intercept) -7.133 0.429 -16.616 2240.573
                                              0.000 -7.975 -6.291
                                                                    NA 0.013
                                                                              0.012
## age
              0.056 0.004 12.952 2468.059
                                             0.000 0.048 0.065
                                                                     0 0.001
                                                                              0.001
## gender2
              -0.422 0.128 -3.304 1749.968 0.001 -0.673 -0.172
                                                                    NA 0.026
                                                                              0.025
## hypchol2
              -0.064 0.188 -0.342 403.591
                                            0.732 -0.434 0.305
                                                                    NA 0.095
                                                                              0.090
## BMI
              0.106 0.009 11.576 2265.730
                                              0.000 0.088
                                                            0.123
                                                                    73 0.012
                                                                              0.011
## smoke2
               0.129 0.144 0.896 2432.312
                                              0.370 -0.153 0.411
                                                                    NA 0.005
                                                                              0.004
## smoke3
              0.080 0.166 0.479 1953.715
                                              0.632 -0.246 0.405
                                                                    NA 0.021
                                                                              0.020
## alc2
              -0.277 0.150 -1.845
                                   475.882
                                              0.066 -0.573 0.018
                                                                    NA 0.085
                                                                              0.082
## alc3
              -0.570 0.220 -2.585 1192.205
                                              0.010 -1.002 -0.137
                                                                    NA 0.042
                                                                              0.041
## alc4
              -0.466 0.246 -1.894 154.639
                                              0.060 -0.952 0.020
                                                                    NA 0.165
                                                                              0.155
## alc5
              -0.741 0.220 -3.375 747.163
                                              0.001 - 1.172 - 0.310
                                                                              0.060
                                                                    NA 0.063
```

9. Analyse & pool the imputed data 9.2. Pooling results

Pooling with mice::pool() is available for most types of models.

Generally, it works for models for which the functions <code>coef()</code> and <code>vcov()</code> can extract the (fixed effects) **coefficients and variance-covariance matrix** of these coefficients.

An alternative is offered by the package **mitools** and the function MIcombine().

9. Analyse & pool the imputed data 9.3. Functions for pooled results

mice currently has two functions available for evaluating model fit / model comparison

For **linear** regression models the pooled R^2 can be calculated using pool.r.squared()

```
mod2 <- with(imp2, lm(SBP ~ DM + age + hypten))
pool.r.squared(mod2, adjusted = TRUE)

## est lo 95 hi 95 fmi
## adj R^2 0.3243655 0.2940335 0.3547556 0.006885114</pre>
```

The argument adjusted specifies whether the adjusted R^2 or the standard R^2 is returned.

9. Analyse & pool the imputed data 9.3. Functions for pooled results

The function pool.compare() allows to compare nested models (i.e., models where one is a special case of the other, with some parameters fixed to zero) using a Wald test.

Example: To test if smoke has a relevant contribution to the model for DM from above we re-fit the model without smoke and compare the two models:

9. Analyse & pool the imputed data

9.3. Functions for pooled results

The package **miceadds** extends **mice**, for example with the following functionality:

Combine χ^2 or F statistics from multiply imputed data:

```
miceadds::micombine.chisquare(dk, df, ...)
miceadds::micombine.F(values, df1, ...)
```

These functions take vectors of statistics computed on each imputed dataset and pool them.

9. Analyse & pool the imputed data

9.3. Functions for pooled results

The package **miceadds** extends **mice**, for example with the following functionality:

Combine χ^2 or F statistics from multiply imputed data:

```
miceadds::micombine.chisquare(dk, df, ...)
miceadds::micombine.F(values, df1, ...)
```

These functions take vectors of statistics computed on each imputed dataset and pool them.

Calculate correlation or covariance of imputed data:

```
miceadds::micombine.cor(mi.res, ...)
miceadds::micombine.cov(mi.res, ...)
```

These functions take mids objects as input.

10. Additional functions in mice() 10.1. Extract & export imputed data

The function complete() allows extraction of the imputed data from a mids object:

```
mice::complete(x, action = 1, include = FALSE)
```

- x: the mids object
- action:
 - 1, ..., m (single imputed dataset)
 - "long": long format (imputed data stacked vertically)
 - "broad": wide format (imputed data combined horizontally; ordered by imputation)
 - "repeated": (like "broad", but ordered by variable)
- include: include the orginal data?(if action is "long", "broad" or "repeated")

10. Additional functions in mice() 10.1. Extract & export imputed data

The function mids2spss() allows the export of imputed data (mids objects) to SPSS.

Data from mids objects can also be exported to MPLUS using mids2mplus().

10. Additional functions in mice() 10.2. Combining mids objects

To increase the number of imputed datasets without re-doing the initial m imputations, a second set of imputations can be done and the two mids objects combined using ibind().

```
# same syntax as before, but different seed
imp2b <- update(imp2, post = post, maxit = 20, seed = 456)
imp2combi <- ibind(imp2, imp2b)</pre>
```

```
# check the new number of impute datasets:
imp2combi$m
## [1] 10
```

10. Additional functions in mice()

10.3. Adding variables to mids objects

The function cbind.mids() allows to add columns to a mids object. The extra columns can either be a data.frame, matrix, vector or factor or another mids object.

For example data columns that should be part of the imputed data for completeness, but are not needed in the imputation.

```
extravar <- rnorm(nrow(NHANES))
impextra <- mice:::cbind.mids(x = imp2, extravar = extravar)</pre>
```

Note: cbind() just adds columns to the data, you need to make sure they are **sorted correctly** so that the rows of the new data are from the same subjects as the corresponding rows in the impute data.

11. Multiple Imputation in SPSS 11.1. Where to get help

A walk-through how to do multiple imputation in SPSS can be found

- for older versions of SPSS
 - > Help
 - > Case Studies
 - > Missing Values Option
 - > Multiple Imputation
 - > Using Multiple Imputation to Complete and Analyze a Dataset
- for newer versions online

```
https://www.ibm.com/support/knowledgecenter/en/SSLVMB_24.0.
0/spss/tutorials/mi_table.html
```

11.1. Where to get help

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```
https://www.ibm.com/support/knowledgecenter/en/SSLVMB_24.0.0/spss/tutorials/mi_table.html
```

The procedure itself is located in the menu

- > Analyze
 - > Multiple Imputation
 - > Impute Missing Data Values

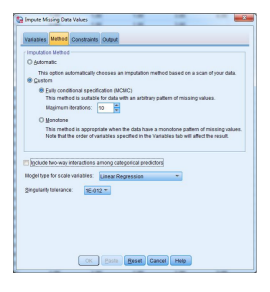
11.2. Multiple Imputation Features



SPSS lets you

specify number of imputations

11.2. Multiple Imputation Features



SPSS lets you

- specify number of imputations
- specify number of iterations
- include interactions
- chose between lin. regression and pmm for continuous variables

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SPSS lets you

- specify number of imputations
- specify number of iterations
- include interactions
- chose between lin. regression and pmm for continuous variables
- restrict variables to certain values
- select which variables to impute
- select which variables are used as predictors

11.2. Multiple Imputation Features



SPSS lets you

- specify number of imputations
- specify number of iterations
- include interactions
- chose between lin. regression and pmm for continuous variables
- restrict variables to certain values
- select which variables to impute
- select which variables are used as predictors
- save the iteration history

11.2. Multiple Imputation Features

SPSS does not let you

- select between linear regression imputation and pmm per variable (only jointly for all variables)
- uses only one donor in predictive mean matching
- uses logistic regression for any categorical variable
- chose per imputation model which variables should be used as predictors
- re-calculate variables during the iterations
- ...

11.2. Multiple Imputation Features

In SPSS the list of models that can be pooled is available in the help under

- > Help
 - > Missing Values Option
 - > Multiple Imputation
 - > Analyzing Multiple Imputation Data

https://www.ibm.com/support/knowledgecenter/en/SSLVMB_24.0.0/spss/mva/mi_analysis.html

Practical

To practice all that we have seen above, go to

https://emcbiostatistics.shinyapps.io/MICourse_MICE

or download the instructions and data for the practical from Canvas (Files > Principal documents > Multiple Imputation > Practical MICE).

Part III When MICE might fail

Outline of Part III

12. Settings where MICE may have problems

12.1 Example: Quadratic effect

12.2 Example: Interaction effect

12.3 Example: Longitudinal outcome

12.4 Example: Survival data

13. Requirements for MICE to work (well)

- 13.1 Joint and conditional distributions
- 13.2 Some conditions and definitions
- 13.3 Why imputation with MICE can go wrong

14. Alternatives to MICE

- 14.1 Joint model imputation
- 14.2 Multivariate Normal Model
- 14.3 Sequential Factorization

Outline of Part III (cont.)

15. Imputation with non-linear functional forms

- 15.1 R package mice
- 15.2 R package JointAl
- 15.3 R package smcfcs
- 15.4 R package jomo
- 15.5 Comparison of results

16. Imputation of longitudinal data

- 16.1 R package mice
- 16.2 R package JointAl
- 16.3 R package jomo
- 16.4 Comparison of results

Outline of Part III (cont.)

17. Imputation of survival data

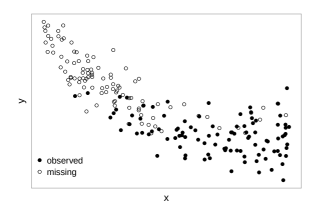
- 17.1 Results from literature
- 17.2 R package mice
- 17.3 R package smcfcs
- 17.4 R package jomo
- 17.5 Comparison of results

12.1. Example: Quadratic effect

Consider the case where the analysis model (which we assume to be true) is

$$y = \beta_0 + \beta_1 x + \beta_2 x^2 + \dots,$$

i.e., y has a quadratic relationship with x, and x is incomplete.



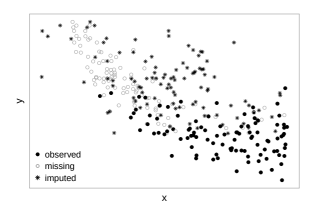
The original data show a curved pattern.

12.1. Example: Quadratic effect

The model used to **impute** x when using MICE (naively) is

$$x = \theta_{10} + \theta_{11}y + \dots,$$

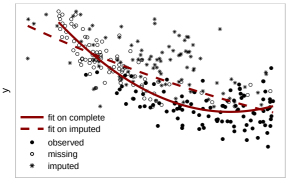
i.e., a **linear relation** between x and y is assumed.



The imputed values distort the curved pattern of the original data.

12.1. Example: Quadratic effect

The model fitted on the imputed data gives **severely biased results**; the non-linear shape of the curve has almost completely disappeared.



	β	95% CI
Original		
Intercept	-0.99	[-1.04, -0.95]
×	-0.61	[-0.66, -0.56]
x^2	0.52	[0.43, 0.62]
Imputed		
Intercept	-0.73	[-0.79, -0.66]
X	-0.53	[-0.62, -0.44]
x ²	0.07	[-0.07, 0.22]

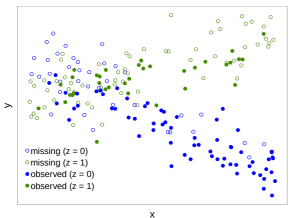
Х

12.2. Example: Interaction effect

Another example occurs when the analysis model (again, assumed to be true) is

$$y = \beta_0 + \beta_x x + \beta_z z + \beta_{xz} xz + \dots,$$

i.e., y has a non-linear relationship with x due to the interaction term.



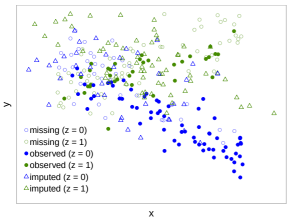
The original data shows a "<" shaped pattern.

12.2. Example: Interaction effect

The model used to impute x when using MICE (naively) is

$$x = \theta_{10} + \theta_{11}y + \theta_{12}z + \dots,$$

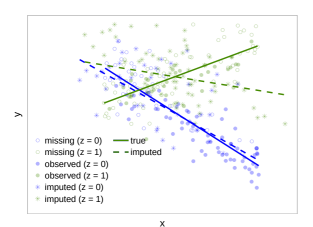
i.e., a linear relation between x and y is assumed.



The "<" shaped pattern of the true data is distorted by the imputed values.

12.2. Example: Interaction effect

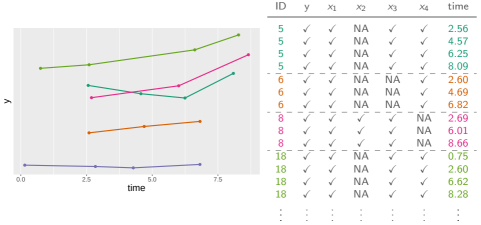
And the analysis on these naively imputed values leads to **severely biased estimates**.



	β	95% CI
Original		
Intercept	-0.96	[-1.00, -0.92]
×	-0.59	[-0.65, -0.53]
Z	0.5	[0.45, 0.56]
x:z	0.94	[0.85, 1.03]
Imputed		
Intercept	-0.96	[-1.01, -0.91]
×	-0.52	[-0.61, -0.44]
Z	0.46	[0.39, 0.54]
X:Z	0.37	[0.24, 0.51]

12.3. Example: Longitudinal outcome

Another setting where imputation with MICE is not straightforward is when the **outcome variable is longitudinal**.



Here, x_1, \ldots, x_4 are baseline covariates, i.e., not measured repeatedly.

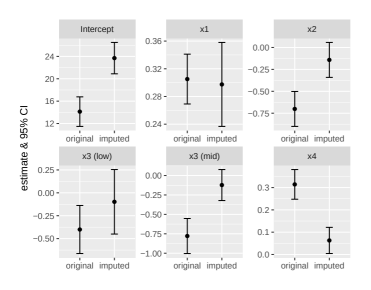
12.3. Example: Longitudinal outcome

If we use MICE in the data in this (long) format, each row would be regarded as independent, which may cause bias and **inconsistent imputations**.

Imputed values of baseline covariates are imputed with different values, creating data that could not have been observed.

ID	У	x_1	<i>x</i> ₂	<i>X</i> 3	<i>X</i> 4	time
5	√	√	boy	√	√	2.56
5	\checkmark	\checkmark	girl	✓	\checkmark	4.57
5	\checkmark	\checkmark	girl	✓	\checkmark	6.25
5	\checkmark	\checkmark	girl	✓	\checkmark	8.09
6			girl	high		2.60
6	\checkmark	\checkmark	boy	mid	\checkmark	4.69
6	\checkmark	\checkmark	girl	high	✓	6.82
8					38.27	2.69
8	\checkmark	\checkmark	\checkmark	\checkmark	38.45	6.01
8	\checkmark	\checkmark	\checkmark	\checkmark	40.71	8.66
18		√	boy			0.75
18	\checkmark	\checkmark	boy	✓	\checkmark	2.60
18	\checkmark	\checkmark	boy	✓	\checkmark	6.62
18	\checkmark	\checkmark	boy	✓	\checkmark	8.28
:	:	:	:	:	:	:

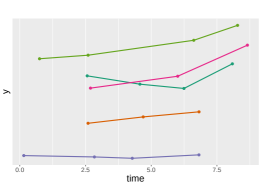
12.3. Example: Longitudinal outcome



Estimates can be severely biased.

12.3. Example: Longitudinal outcome

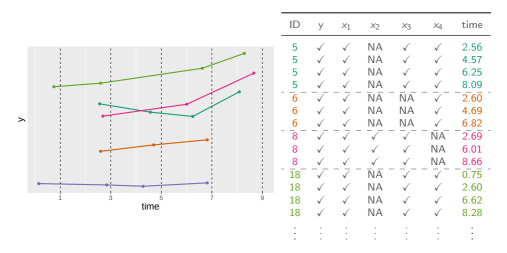
In some settings imputation in wide format may be possible.



ID	У	x_1	<i>x</i> ₂	<i>X</i> 3	<i>X</i> ₄	time
5	√	√	NA	√	√	2.56
5	\checkmark	\checkmark	NA	\checkmark	\checkmark	4.57
5	\checkmark	\checkmark	NA	\checkmark	\checkmark	6.25
5	\checkmark	\checkmark	NA	\checkmark	\checkmark	8.09
6			NA	NA		2.60
6	\checkmark	\checkmark	NA	NA	\checkmark	4.69
6	\checkmark	\checkmark	NA	NA	\checkmark	6.82
8		- 7 -			ΝA	2.69
8	\checkmark	\checkmark	√	\checkmark	NA	6.01
8	\checkmark	\checkmark	√	\checkmark	NA	8.66
18			NĀ			0.75
18	\checkmark	\checkmark	NA	\checkmark	\checkmark	2.60
18	\checkmark	\checkmark	NA	\checkmark	\checkmark	6.62
18	\checkmark	\checkmark	NA	\checkmark	\checkmark	8.28
:	:	:	:	:	:	:
•						•

12.3. Example: Longitudinal outcome

In some settings imputation in wide format may be possible.

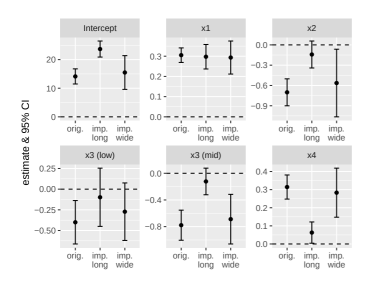


12.3. Example: Longitudinal outcome

id	y.1	y.3	y.5	y.7	y.9	time.1	time.3	time.5	time.7	time.9	
5	NA	35.19	34.94	34.81	35.56	NA	2.56	4.57	6.25	8.09	
6	NA	33.74	33.94	34.09	NA	NA	2.6	4.69	6.82	NA	
8	NA	34.82	NA	35.18	36.13	NA	2.69	NA	6.01	8.66	
18	35.71	35.82	NA	36.28	36.73	0.75	2.6	NA	6.62	8.28	
:	:		:	:	:		:	:	:	:	٠.
	•	•				•	•	•	•	•	

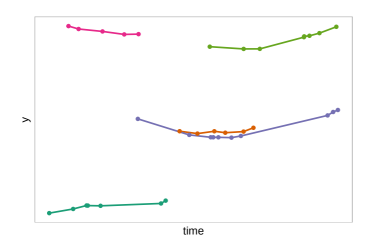
In this **wide format data** frame, missing values in outcome and measurement times need to be imputed (to be able to use them as predictors to impute covariates), even though we would not need to impute them for the analysis (mixed model valid when outcome measurements are M(C)AR).

12.3. Example: Longitudinal outcome



Better, but very large confidence intervals.

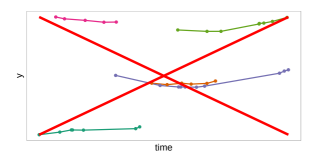
12.3. Example: Longitudinal outcome



When the data is very **unbalanced**, i.e., there are no clear cut-offs in time, transformation to wide format is not possible.

(Or at least transformation to wide format leads to variables with high proportions of missing values.)

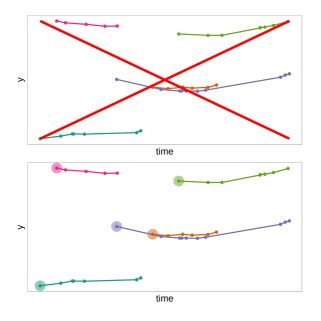
12.3. Example: Longitudinal outcome



Naive approaches that are sometimes used are to

• **ignore the outcome** in the imputation

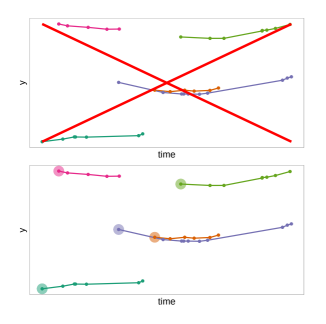
12.3. Example: Longitudinal outcome



Naive approaches that are sometimes used are to

- **ignore the outcome** in the imputation, or to
- use only the first/baseline outcome

12.3. Example: Longitudinal outcome



Naive approaches that are sometimes used are to

- **ignore the outcome** in the imputation, or to
- use only the first/baseline outcome

However, **important information may be lost**, resulting in invalid imputations and biased results.

12.4. Example: Survival data

In survival analysis, the aim is to estimate the effect of covariates on the time until an event of interest happens.

12.4. Example: Survival data

In survival analysis, the aim is to estimate the effect of covariates on the time until an event of interest happens.

In the commonly used method: Cox proportional hazards model

$$h(t) = h_0(t) \exp(x\beta_x + z\beta_z),$$

- h(t): hazard = the instantaneous risk of an event at time t, given that the event has not occurred until time t
- $h_0(t)$: unspecified baseline hazard
- x and z: **incomplete** and **complete** covariates, respectively

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- h(t): hazard = the instantaneous risk of an event at time t, given that the event has not occurred until time t
- $h_0(t)$: unspecified baseline hazard
- x and z: **incomplete** and **complete** covariates, respectively

Survival outcomes are usually represented by the **observed event time** T and the **event indicator** D (D = 1: event, D = 0: censored).

12.4. Example: Survival data

Naive use of MICE treats the columns in the data set containing T and D just like any other variable, and the resulting imputation model for X would have the form

$$p(x \mid T, D, \mathbf{z}) = \theta_0 + \theta_1 T + \theta_2 D + \theta_3 z + \dots$$

12.4. Example: Survival data

Naive use of MICE treats the columns in the data set containing T and D just like any other variable, and the resulting imputation model for X would have the form

$$p(x \mid T, D, \mathbf{z}) = \theta_0 + \theta_1 T + \theta_2 D + \theta_3 z + \dots$$

The correct conditional distribution of x given the other variables is, however,

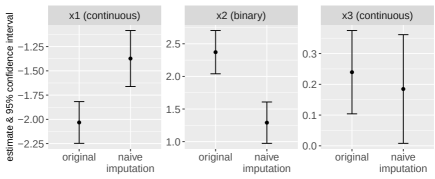
$$\log p(x \mid T, D, z) = \log p(x \mid z) + D(\beta_x x + \beta_z z) - H_0(T) \exp(\beta_x x + \beta_z z) + const.,$$

where $H_0(T)$ is the cumulative baseline hazard.[20]

12. Settings where MICE may have problems

12.4. Example: Survival data

Using the naively assumed imputation model can lead to severe bias:



(Results from MICE imputation with two incomplete normal and one incomplete binary covariate.)

13.1. Joint and conditional distributions

Recall: The MICE algorithm is based on the idea of Gibbs sampling.

Gibbs sampling exploits the fact that a joint distribution is fully determined by its full conditional distributions.



13.1. Joint and conditional distributions

Recall: The MICE algorithm is based on the idea of Gibbs sampling.

Gibbs sampling exploits the fact that a joint distribution is fully determined by its full conditional distributions.



In MICE, the full conditionals are not derived from the joint distribution: we directly specify the full conditionals and hope a joint distribution exists.

13.1. Joint and conditional distributions

The uncertainty about whether a joint distribution exists for the specified set of imputation models is often considered to be mainly a theoretical problem.

In practice, violations only have little impact on results in many applications.

However, as we have seen in the examples on the previous slides, there are settings where the direct specification of the full conditionals/imputation models may lead to problems, causing biased results.

13.2. Some conditions and definitions

Two important definitions:

Compatibility:

A joint distribution exists, that has the full conditionals (imputation models) as its conditional distributions.

Congeniality:

The imputation model is compatible with the analysis model.

13.2. Some conditions and definitions

Important requirements for MICE to work well include:

- Compatibility
- Congeniality
- MAR or MCAR (in the standard implementations)
- all relevant variables need to be included (omission might result in MNAR)
- The outcome needs to be included as predictor variable (but we usually do not impute missing outcome values)
- the imputation models (and analysis model) need to be correctly specified (which is a requirement in any standard analysis)

13.3. Why imputation with MICE can go wrong

What went wrong in our previous examples?

When incomplete variables have **non-linear associations** with the outcome, or with each other, the requirement(s) of **compatibility and/or congeniality are violated**.

Omission, or inadequate inclusion, of the outcome may result in MNAR missing mechanisms. The same is the case when other relevant predictor variables are not used as predictor variables in the imputation.

Furthermore, omission of variables may lead to mis-specified models, however, models may also be mis-specified when all relevant covariates are included, but distributional assumptions or the specified form of associations are incorrect.

14.1. Joint model imputation

To avoid incompatible and uncongenial imputation models, we need to

- specify the joint distribution
- and derive full conditionals / imputation models from this joint distribution instead of specifying them directly.

14.1. Joint model imputation

To avoid incompatible and uncongenial imputation models, we need to

- specify the joint distribution
- and derive full conditionals / imputation models from this joint distribution instead of specifying them directly.

Problem:

Especially in settings with several **variables of mixed type**, the joint distribution is usually not of any known form:

$$\begin{array}{ccc} x_1 \sim \textit{N}(\mu_1, \sigma_1^2) \\ x_2 \sim \textit{N}(\mu_2, \sigma_2^2) \end{array} \Rightarrow \begin{pmatrix} x_1 \\ x_2 \end{pmatrix} \sim \textit{N}\left(\begin{bmatrix} \mu_1 \\ \mu_2 \end{bmatrix}, \begin{bmatrix} \sigma_1^2 & \sigma_{12} \\ \sigma_{12} & \sigma_2^2 \end{bmatrix} \right) \\ \text{but} \quad \begin{array}{c} x_1 \sim \textit{N}(\mu_1, \sigma_1^2) \\ x_2 \sim \textit{Bin}(\mu_2) \end{array} \Rightarrow \begin{pmatrix} x_1 \\ x_2 \end{pmatrix} \sim ??? \end{array}$$

14.1. Joint model imputation

Approach 1: Multivariate Normal Model

Approximate the joint distribution by a known multivariate distribution (usually the normal distribution; this is the approach mentioned in Part I on slide 15)

Approach 2: Sequential Factorization

Factorize the joint distribution into a (sequence of) conditional and a marginal distributions

14.2. Multivariate Normal Model

Assumption:

The outcome and incomplete variables follow a **joint multivariate normal distribution**, conditional on the completely observed covariates \mathbf{X}_c , parameters $\boldsymbol{\theta}$ and, possibly, random effects, \boldsymbol{b} :

$$p(\mathbf{y}, \mathbf{x}_1, \dots, \mathbf{x}_p \mid \mathbf{X}_c, \boldsymbol{\theta}, \boldsymbol{b}) \sim \mathcal{N}(\boldsymbol{\mu}, \boldsymbol{\Sigma})$$

14.2. Multivariate Normal Model

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$$p(\mathbf{y}, \mathbf{x}_1, \dots, \mathbf{x}_p \mid \mathbf{X}_c, \boldsymbol{\theta}, \mathbf{b}) \sim \mathcal{N}(\boldsymbol{\mu}, \boldsymbol{\Sigma})$$

How do we get that multivariate normal distribution?

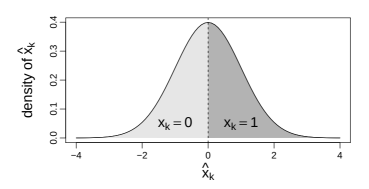
- 1. Assume all incomplete variables and the outcome are (latent) normal.
- 2. Specify linear (mixed) models based on observed covariates.
- 3. Connect using multivariate normal for random effects & error terms.

14.2. Multivariate Normal Model

1. Latent normal assumption:

e.g.: x_k binary \to latent \hat{x}_k is standard normal: $\left\{ \begin{array}{l} x_k = 1 \\ x_k = 0 \end{array} \right.$ if $\left. \begin{array}{l} \hat{x}_k \geq 0 \\ \hat{x}_k < 0 \end{array} \right.$

$$x_k = 1$$
 if $\hat{x}_k \ge 0$
 $x_k = 0$ $\hat{x}_k < 0$



14.2. Multivariate Normal Model

2. Specify models:

$$y = X_c \beta_y + Z_y b_y + \varepsilon_y$$

$$w = X_c \beta_w + Z_w b_w + \varepsilon_w$$

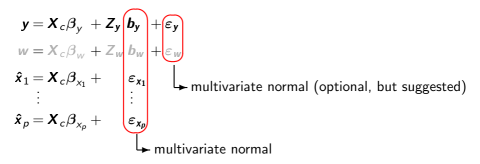
$$\hat{x}_1 = X_c \beta_{x_1} + \varepsilon_{x_1}$$

$$\vdots \qquad \vdots$$

$$\hat{x}_p = X_c \beta_{x_p} + \varepsilon_{x_p}$$

14.2. Multivariate Normal Model

2. Specify models / 3. Connect random effects & error terms:



14.2. Multivariate Normal Model

Advantages:

- Easy to specify
- Relatively easy to implement
- Relatively easy to sample from
- Works for longitudinal outcomes

Imputation with **non-linear associations** or **survival data** is possible with **extensions** of the multivariate normal approach.

Disadvantages:

Assumes linear associations

14.3. Sequential Factorization

The **joint distribution** of two variables y and x can be written as the product of a conditional and a marginal distribution:

$$p(y,x) = p(y \mid x) \ p(x)$$

(or alternatively
$$p(y,x) = p(x \mid y) p(y)$$
)

14.3. Sequential Factorization

The **joint distribution** of two variables y and x can be written as the product of a conditional and a marginal distribution:

$$p(y,x) = p(y \mid x) \ p(x)$$

(or alternatively $p(y,x) = p(x \mid y) p(y)$)

This can easily be extended for more variables:

$$p(y, x_1, \dots, x_p, X_c) = \underbrace{p(y \mid x_1, \dots, x_p, X_c)}_{\text{analysis model}} p(x_1 \mid x_2, \dots, x_p, X_c) \dots p(x_p \mid X_c)$$

where x_1, \ldots, x_p denote incomplete covariates and X_c contains all completely observed covariates.

14.3. Sequential Factorization

That the analysis model is part of the specification of the joint distribution has several advantages:

- The outcome is **automatically included in the imputation** procedure.
- The outcome does not appear in any of the predictors of the imputation models:
 - no need to approximate complex outcomes,
 - no need to summarize complex outcomes.

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- The parameters of interest are obtained directly
 - imputation and analysis in one step

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- The parameters of interest are obtained directly
 - → imputation and analysis in one step
- Non-linear associations or interactions involving incomplete covariates are specified in the analysis model and thereby automatically taken into account

Since the joint distribution usually does not have a known form, Gibbs sampling is used to estimate parameters and sample imputed values.

14.3. Sequential Factorization

Advantages:

- flexible with regards to outcome type
- univariate conditional distributions of incomplete covariates can be chosen according to type of variable
- non-linear associations and interactions can be taken into account
- assures congeniality and compatible imputation models

Disadvantages:

- separate models need to be specified per incomplete variable: takes more time and consideration
- the joint distribution is of unknown form and sampling may be more computationally intensive

15. Imputation with non-linear functional forms

In the following we will not only consider the R package **mice**, but also three additional packages, **JointAI**, **smcfcs** and **jomo**, that provide alternatives to **mice**.

These three packages use **Bayesian methodology** to impute values, but once imputed datasets are obtained, standard complete data methods can be used.

jomo and **smcfcs** perform **multiple imputation** and create imputed datasets that can then be analysed the same way data imputed by **mice** would be analysed.

JointAI works **fully Bayesian** and performs the analysis and imputation simultaneously, so that the results from the analysis model of interest are obtained directly.

There is no strategy for MICE that can guarantee valid imputations when non-linear functional forms and/or interactions are involved, but some settings in **mice** may help to reduce bias in the resulting estimates.

For imputation of variables that have non-linear associations

- pmm often works better than norm,
- Just Another Variable approach can reduce bias in interactions,
- quadratic can help to impute variables with quadratic association.

In the **Just Another Variable (JAV)** approach the non-linear form (or interaction term) is calculated in the incomplete data, added as a column to the dataset and imputed as if it was just another variable.

quadratic provides imputation of covariates that have a quadratic association with the outcome, using the "polynomial combination" method.[17, pp. 139–141], [19].

This is to ensure the imputed values for x and x^2 are consistent, and to reduce bias in the subsequent analysis that uses x and x^2 .

In my experience, using quadratic can lead to numerical problems.

To demonstrate the approaches, we use a simulated example dataset DFnonlin, with

- continuous outcome y
- continuous (normal) covariate x (50% missing values MCAR)
- quadratic effect of x on y
- binary covariate z (complete)
- interaction between x and z

To demonstrate the approaches, we use a simulated example dataset DFnonlin, with

- continuous outcome y
- continuous (normal) covariate x (50% missing values MCAR)
- quadratic effect of x on y
- binary covariate z (complete)
- interaction between x and z

In the naive approach, we leave all settings to the defaults.

```
# naive imputation, using only y, x, z
impnaive <- mice(DF_nonlin, printFlag = F)</pre>
```

We use two different JAV approaches:

JAV: calculating the quadratic and interaction term before imputation

```
# add quadratic term and interaction to data
DF2 <- DF_nonlin
DF2$xx <- DF2$x^2
DF2$xz <- DF2$x * DF2$z

# JAV imputation
impJAV <- mice(DF2, printFlag = F, maxit = 20)</pre>
```

JAV2: additionally using an interaction between z and y

```
# add interaction between y and z to data
DF3 <- DF2
DF3$yz <- DF3$y * DF3$z

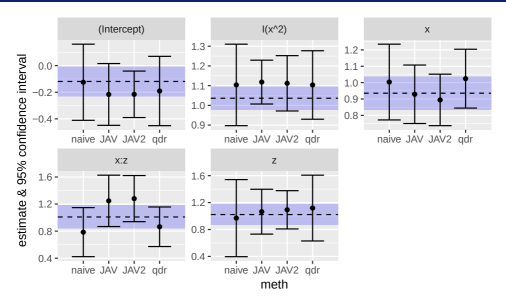
# JAV imputation with additional interaction
impJAV2 <- mice(DF3, printFlag = F, maxit = 20)</pre>
```

We also try using imputation method quadratic.

Note: there were warning messages about numerical issues for this approach.

15. Imputation with non-linear functional forms

15.1. R package mice



For this example, none of the approaches provided satisfying results.

The package **JointAI** uses the **sequential factorization approach** to perform simultaneous analysis and imputation.[4, 3]

JointAI (version 0.1.0) can handle

- linear regression
- generalized linear regression
- linear mixed models

while assuring compatibility between analysis model and imputation models when non-linear functions or interactions are included.

The necessary Gibbs sampling is performed using JAGS (an external program), which is free, but needs to be installed from https://sourceforge.net/projects/mcmc-jags/files/.

Joint AI can be installed from CRAN

```
install.packages("JointAI")
```

The development version (containing bug fixes and other improvements) can be installed from GitHub

```
install.packages("devtools")
devtools::install_github(repo = "JointAI", username = "NErler")
```

A detailed explanation of the functionality is given in the help files of the package, and a vignette with an in-depth example analysis will be available soon.

The syntax we use to analyse and impute the current example using **JointAl** is similar to the specification of a standard linear model using lm().

The syntax we use to analyse and impute the current example using **JointAl** is similar to the specification of a standard linear model using lm().

Convergence of the Gibbs sampler can be checked using a traceplot.

```
traceplot(JointAI_nonlin)
```

Results (no separate analysis & pooling is necessary) can be obtained with the summary() function:

```
res_JointAI_nonlin <- summary(JointAI_nonlin)
```

The package **smcfcs** performs multiple imputation using *substantive model* compatible fully conditional specification, a **hybrid approach between FCS** and sequential factorization.[1]

smcfcs (version 1.3.0) can handle

- linear regression,
- logistic regression,
- poisson regression,
- Cox proportional hazard models, and
- competing risk survival models,

while ensuring compatibility between analysis model and imputation models.

For more information see the help files and the vignette.

The syntax to impute the data in the current example using the package **smcfcs** is:

The convergence of the procedure should be checked, for example with the following syntax:

```
par(mfrow = c(2,3), mar = c(2, 2, 0.5, 0.5), mgp = c(2, 0.6, 0))
for(i in 1:dim(smcfcs_nonlin$smCoefIter)[2]) {
   matplot(t(smcfcs_nonlin$smCoefIter[, i, ]), type = 'l', ylab = '')
}
```

To be able to use the convenient pooling function from the **mice** package we first need to convert the imputed data (which is a list) to a mids object.

This can be done with the function datalist2mids() from the miceadds
package.

```
library(miceadds)
impobj_smcfcs_nonlin <- datalist2mids(smcfcs_nonlin$impDatasets)</pre>
```

The mids object can then be pooled and summarized as we have seen before with mids objects created by mice().

```
models_smcfcs_nonlin <- with(impobj_smcfcs_nonlin, lm(y ~ x*z + I(x^2)))
res_smcfcs_nonlin <- summary(pool(models_smcfcs_nonlin))</pre>
```

The package **jomo** performs **joint model imputation** using the multivariate normal approach, with **extensions to assure compatibility** between analysis and imputation models.[2]

jomo (version 2.6-2) can handle

- linear regression,
- generalized linear regression,
- linear mixed models,
- generalized linear mixed models, and
- Cox proportional hazards models.

Using **jomo** we can impute the data in the current example as follows:

```
library(jomo)
jomo_nonlin <- jomo.lm(y ~ x*z + I(x^2), data = DF_nonlin)</pre>
```

Using jomo we can impute the data in the current example as follows:

```
library(jomo)
jomo_nonlin <- jomo.lm(y ~ x*z + I(x^2), data = DF_nonlin)</pre>
```

To check the convergence of the model, the corresponding function with ending .MCMCchain() as to be used.

```
jomo_nonlinMCMC <- jomo.lm.MCMCchain(y ~ x*z + I(x^2), data = DF_nonlin)</pre>
par(mfcol = c(2, 3), mar = c(3, 2.5, 0.5, 0.5), mgp = c(2, 0.6, 0))
apply(jomo_nonlinMCMC$collectbeta[1, ,], 1, plot, type = "l",
      xlab = 'iteration', ylab = '')
for (k in 1:dim(jomo_nonlinMCMC$collectomega)[1]) {
  apply(jomo_nonlinMCMC$collectomega[k, , ], 1, plot, type = "1",
        xlab = 'iteration', ylab = '')
apply(jomo_nonlinMCMC$collectbetaY[1, ,], 1, plot, type = "l",
      xlab = 'iteration', ylab = '')
plot(jomo_nonlinMCMC$collectvarY, type = '1')
                                                                       175 / 225
```

Again, we need to convert the output to a mids object using datalist2mids(). However, jomo.lm() returns a data frame, in which the original data and all imputed datasets are stacked onto each other.

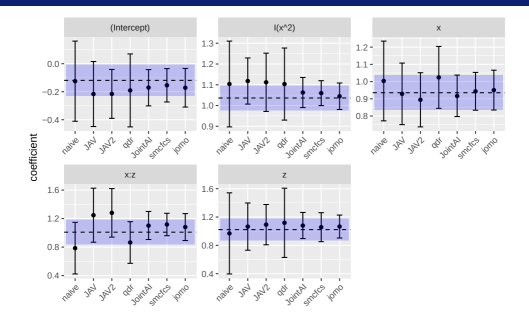
split() splits the dataset by imputation number into a list of datasets, from which we need to exclude the first element (the original/incomplete data).

With the resulting mids object, analysis of the imputed data and pooling of the results works as in the above examples.

```
models_jomo_nonlin <- with(impobj_jomo_nonlin, lm(y ~ x*z + I(x^2)))
res_jomo_nonlin <- summary(pool(models_jomo_nonlin))</pre>
```

15. Imputation with non-linear functional forms

15.5. Comparison of results



Practical

To practice imputation with non-linear forms or interaction terms, go to

https://emcbiostatistics.shinyapps.io/MICourse_MIadvanced

or download the instructions and data for the practical from Canvas (Files > Principal documents > Multiple Imputation > Practical Mladvanced).

mice has functions to allow imputation of longitudinal (2-level) data.

- Level 1: repeated measurements (within subjects) or subjects (within classes)
- Level 2: time-constant/baseline covariates, between subjects effects, variables on the group level

Imputation methods for **level-1** variables:

- 21.pan
- 21.norm
- 21.lmer

Imputation methods for **level-2** variables:

- 2lonly.norm
- 2lonly.pmm
- 2lonly.mean

21. pan uses a linear two-level model with **homogeneous within group** variances using Gibbs sampling [14]. It needs the package **pan** to be installed.

21.pan allows for different roles of predictor variables, that can be specified as different values in the predictorMatrix:

- grouping/ID variable: -2
- random effects (also included as fixed effects): 2
- fixed effects of group means: 3
- fixed effects of group means & random effects: 4

```
# random effects of x in model for y
pred["y","x"] <- 2
# fixed effects of x and group mean of x
pred["y","x"] <- 3
# random effects of x and group mean of x
pred["y","x"] <- 4</pre>
```

21.norm implements a (Bayesian) linear two-level model with **heterogenous** group variances.

In the current implementation all predictors should be specified as random effects (set to 2 in the predictorMatrix, because the algorithm does not handle predictors that are specified as fixed effects).

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In the current implementation all predictors should be specified as random effects (set to 2 in the predictorMatrix, because the algorithm does not handle predictors that are specified as fixed effects).

21.1mer imputes univariate systematically and sporadically missing data using a two-level normal model using lmer() from package lme4 (developed in the context of individual patient meta analysis. [7, 6])

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In the current implementation all predictors should be specified as random effects (set to 2 in the predictorMatrix, because the algorithm does not handle predictors that are specified as fixed effects).

21.1mer imputes univariate systematically and sporadically missing data using a two-level normal model using lmer() from package lme4 (developed in the context of individual patient meta analysis. [7, 6])

2lonly.norm and 2lonly.pmm can be used to impute level-2 variables (in combination with 2l.pan for level-1 variables).

In all case, the group identifier ("id" variable") needs to be set to -2 in the predictorMatrix.

21only.mean imputes values with the mean of the observed values per class. This method should only be used to fill in values that are known to be constant per class and have some values observed in each class.

Example: In a multi-center trial the type of some medical equipment is known to be the same for all patients treated in the same hospital, but not filled in for some patients.

As an example, we will impute the second (unbalanced) longitudinal data example from above. The data contain

- x1 (complete)
- x2 (binary, 30% missing values)
- x3 (3 categories, 30% missing values)
- x4 (continuous/normal, 30% missing values)
- y (longitudinal outcome)
- time (time variable with quadratic effect)
- id (id variable)

Since there is no 2-level method for categorical data, we use 2lonly.pmm to impute x2 and x3.

As usual, we start with the setup run of mice()

```
imp0 <- mice(DFexlong2, maxit = 0)
meth <- imp0$method
pred <- imp0$predictorMatrix</pre>
```

and adjust the imputation method and predictorMatrix

```
meth[c("x2", "x3")] <- "2lonly.pmm"
meth[c("x4")] <- "2lonly.norm"

pred[, "id"] <- -2 # identify id variable
pred[, "ti"] <- 0 # don't use time-point indicator</pre>
```

We can then perform the imputation.

The imputed data can be analysed using either lmer() from the package lme4, or lme() from nlme. Here we use the former.

Currently, there is only limited documentation and examples available that show how to use these functions in **mice**.

Technical details can be obtained from the methodological references given in the help files of the R functions.

A vignette on multi-level imputation with **mice** is available. It gives a more elaborate example of how to analyse such data.

Linear mixed models with incomplete covariates can also be analysed using the package **JointAI**.

The syntax is analogous the syntax used in lme() of the package nlme.

Again, convergence of the Gibbs sampler should be checked using a traceplot,

```
traceplot(JointAI_long)
```

before obtaining the results:

```
res_JointAI_long <- summary(JointAI_long)</pre>
```

Contrary to the two-level imputation of **mice**, non-linear associations are appropriately handled.

In **jomo**, the functions **jomo.lmer()** and **jomo.glmer()** can be used to impute longitudinal data with normal or non-normal outcomes.

In the multi-level setting, the **level of each variable** needs to be specified (1: repeated measurements, 2: baseline covariates), and **ordered the same way** the variables occur in the data frame.

Like in the example with non-linear effects, convergence of the imputation needs to be checked.

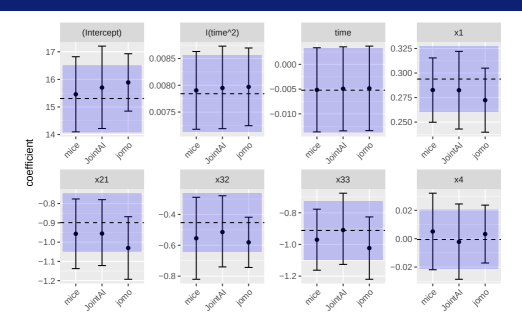
Again, the stacked dataframe returned by <code>jomo.lmer()</code> needs to be split by imputation number and the original data excluded, before fitting the model and pooling the results.

(Note: jomo.lmer() re-names the grouping variable to clus).

As in the examples for non-linear functional forms, congeniality of imputation models is maintained.

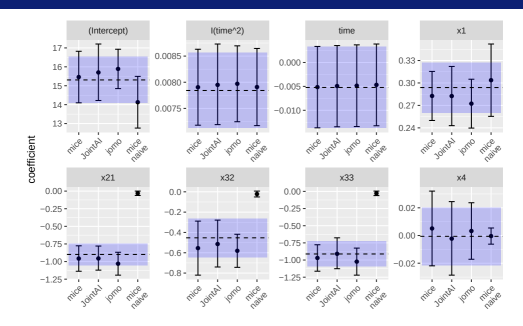
16. Imputation of longitudinal data

16.4. Comparison of results



16. Imputation of longitudinal data

16.4. Comparison of results



Practical

To practice imputation with longitudinal data, continue with the practical at

https://emcbiostatistics.shinyapps.io/MICourse_MIadvanced

or the offline version that can be downloaded from Canvas (Files > Principal documents > Multiple Imputation > Practical Mladvanced).

17.1. Results from literature

On slide 145 we have seen the rather complex formula for imputation of an incomplete covariate in survival data.

White et al. [20] derived versions of this model for different settings (binary or continuous incomplete covariate X, and continuous, categorical or no complete covariate Z) and investigated how to best approximate it.

They found that when **covariate effects and cumulative incidences are rather small**, using Z, D and $H_0(T)$, and possibly an interaction term, as predictor variables in the imputation for X in MICE may work satisfactorily.

However, in practice $H_0(T)$ is unspecified.

17.1. Results from literature

Two main ideas:

- If covariate effects β_x and β_z are small, $H_0(t) \approx H(t)$, which can be approximated by the **Nelson-Aalen estimator**.
- Estimate $H_0(T)$ in an additional step inside the MICE procedure by fitting a Cox model on the imputed data.

Neither of these approaches takes into account uncertainty about $H_0(t)$ (but the impact is likely to be small).

17.1. Results from literature

Two main ideas:

- If covariate effects β_x and β_z are small, $H_0(t) \approx H(t)$, which can be approximated by the **Nelson-Aalen estimator**.
- Estimate $H_0(T)$ in an additional step inside the MICE procedure by fitting a Cox model on the imputed data.

Neither of these approaches takes into account uncertainty about $H_0(t)$ (but the impact is likely to be small).

Based on results from their simulation study, White et al. conclude that **using** Z, D and the Nelson-Aalen estimator $\hat{H}(T)$ as predictors for the imputation of X worked best.

However, some bias towards the null should be expected when covariates have large effects.

17. Imputation of survival data 17.2. R package mice

In mice, nelsonaalen() can be used to calculate the Nelson-Aalen estimator, to use it as covariate in the imputation.

```
survdat$H0 <- nelsonaalen(survdat, timevar = Time, statusvar = event)</pre>
```

Then, we can prepare the imputation using the same steps as in previous examples:

```
# setup run
imp0 <- mice(survdat, maxit = 0)
meth <- imp0$method
pred <- imp0$predictorMatrix

# specify normal imputation for continuous covariates
meth[c("x1", "x3")] <- "norm"

# remove event time from predictor (high correlation with HO)
pred[, "Time"] <- 0</pre>
```

17.2. R package mice

With the modified arguments method and predictorMatrix we run the imputation:

To obtain the pooled results, we first fit the model of interest

```
cox_mice <- with(survimp, coxph(Surv(Time, event) ~ x1 + x2 + x3))</pre>
```

and pool and summarize the results.

```
res_mice_surv <- summary(pool(cox_mice))
## Warning in mice.df(m, lambda, dfcom, method): Large sample
assumed.</pre>
```

The warning message refers to the way the degrees of freedom for the formulas we saw in Part I (slide 32) are calculated and can be ignored.

17. Imputation of survival data 17.3. R package smcfcs

31 11 hadiia83 3iii3133

Using the package **smcfcs**, the same data can be imputed with the following syntax:

Convergence of the procedure should be checked, analogously to the previous example (see slide 172).

After the resulting object is converted to a mids object, fitting the model and pooling the results is identical to what was done with the data imputed by **mice**.

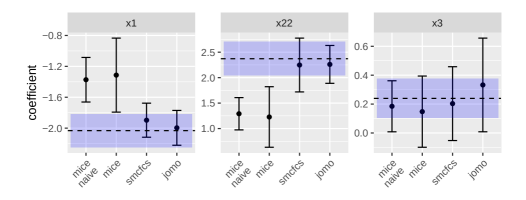
17. Imputation of survival data 17.4. R package jomo

In the package **jomo**, the function **jomo.coxph**() can be used to impute our example survival data:

Note that the convergence of the procedure should be checked using jomo.coxph.MCMCchain() (see the previous examples using jomo).

To analyse & pool the imputed data the steps are identical to the other examples:

17.5. Comparison of results



The naive mice approach, and mice using the Nelson-Aalen estimator give very biased results for the effects of x1 and x2, but performed acceptably well for x3.

Note that the **true effects** (log HR) of x1 and x2 are **very large** (-2 and 2.5, respectively), and represent the setting where the approximation by the Nelson-Aalen estimate is **expected to be biased**.

Practical

To practice imputation with survival data, continue with the practical at

https://emcbiostatistics.shinyapps.io/MICourse_MIadvanced

or the offline version that can be downloaded from Canvas (Files > Principal documents > Multiple Imputation > Practical Mladvanced).

Summary & Conclusion of Part III

- MICE requires congenial & compatible imputation models to work well.
- When this is not the case, (naive) use of MICE can lead to biased results.
- Common settings that require special attention are
 - non-linear functional forms & interaction terms
 - longitudinal data
 - survival data

Summary & Conclusion of Part III (cont.)

- When using the package mice, there are choices that can reduce bias
 - pmm tends to be less biased than norm for interactions or non-linear associations
 - JAV approach reduces bias in settings with interactions or non-linear associations
 - special 2-level imputation methods are available for longitudinal data
 - The **Nelson-Aalen estimator** can be used instead of the time variable for imputing survival data when effects are not too large.
- Generally, problems are more severe when
 - proportions of missing values are large,
 - effect sizes are large,
 - little other **covariate information** is available.

(Note that in the examples we had all of the above.)

Summary & Conclusion of Part III (cont.)

- In settings where MICE may not provide valid imputations, alternative approaches are available and should be considered.
- R packages that provide such alternative approaches are for example:
 - JointAl (non-linear & longitudinal)
 - smcfcs (non-linear & survival)
 - jomo (non-linear, longitudinal & survival)
- These packages are very young.
 - Hence, they may still have some problems.
 - → Use them carefully! (and email the maintainer about problems)
 - They are under active development, so resolutions of bugs and features are frequently added.

Part IV Multiple Imputation Strategies

Outline of Part IV

18. Strategies for using MICE

- 18.1 Imputation methods
- 18.2 Tips & Tricks
- 18.3 Number of imputed datasets
- 18.4 What to do with large datasets?
- 18.5 How much missing is too much?
- 18.6 Imputation of outcomes
- 18.7 Notes of caution & things to keep in mind

19. MICE and MI in the bigger picture

- 19.1 Other R packages that do imputation
- 19.2 Imputation in other software
- 19.3 Other approaches to handle missing values

18.1. Imputation methods

We have focussed on a few imputation methods that cover the most common types of data but there are many more methods implemented.

Imputation methods implemented in the **mice** package:

mice.impute.2l.lmer mice.impute.2l.norm mice.impute.2l.pan mice.impute.2lonly.mean mice.impute.2lonly.norm mice.impute.2lonly.pmm mice.impute.cart mice.impute.lda mice.impute.logreg
mice.impute.logreg.boot
mice.impute.mean
mice.impute.midastouch
mice.impute.norm
mice.impute.norm.boot
mice.impute.norm.nob
mice.impute.norm.predict

mice.impute.passive mice.impute.pmm mice.impute.polr mice.impute.polyreg mice.impute.quadratic mice.impute.rf mice.impute.ri mice.impute.sample

Note: Just because a method is implemented does not mean you need to / should use it.

18. Strategies for using MICE 18.1. Imputation methods

Imputation methods implemented in the **miceadds** package:

mice.impute.2l.binary mice.impute.2l.contextual.norm mice.impute.2l.contextual.pmm mice.impute.2l.continuous mice.impute.2l.eap mice.impute.2l.groupmean mice.impute.2l.groupmean.elim mice.impute.2l.latentgroupmean.mcmc mice.impute.2l.latentgroupmean.ml mice.impute.2l.plausible.values mice.impute.2l.pls mice.impute.2l.pls2 mice.impute.2l.pmm mice.impute.2lonly.function mice.impute.2lonly.norm2 mice.impute.2lonly.pmm2

mice.impute.bygroup mice.impute.eap mice.impute.grouped mice.impute.hotDeck mice.impute.ml.lmer mice.impute.plausible.values mice.impute.pls mice.impute.pmm3 mice.impute.pmm4 mice.impute.pmm5 mice.impute.pmm6 mice.impute.tricube.pmm mice.impute.tricube.pmm2 mice.impute.weighted.norm mice.impute.weighted.pmm

18. Strategies for using MICE 18.2. Tips & Tricks

In complex settings, variables may need to be **re-calculated** or **re-coded** after imputation:

- Use complete() to convert the imputed data from a mids object to a data.frame.
- Perform the necessary calculations.
- Convert the changed data.frame back to a mids object using the functions from the flow-diagram in the second practical (e.g., as.mids(), datalist2mids(), imputationList(), ...)

Not just in imputation: Set a **seed value** to create reproducible results.

- in R: set.seed()
- in mice(): argument seed

18. Strategies for using MICE 18.3. Number of imputed datasets

Early publications on multiple imputation suggested that 3-5 imputations are sufficient and this is still a common assumption in practice.[12]

The reasoning behind using a small number of imputed datasets was that storage of imputed data was "expensive" (which is no longer the case) and a larger number of imputations would only have little advantage.[13]

More **recent work** from various authors [21, 17, 5] considers the efficiency of the pooled estimates, reproducibility of the results, statistical power of tests or the width of the resulting confidence intervals compared to the width of the true confidence intervals.

18. Strategies for using MICE 18.3. Number of imputed datasets

A suggested rule of thumb is that the number of imputed datasets should be similar to the percentage of incomplete cases.[21] Since this percentage depends on the size of the dataset, the average percentage of missing values per variable could be used as an alternative.[17]

Generally, using **more imputed datasets should be preferred**, especially in settings where the computational burden allows for it. Even though results are unlikely to change with a larger number of imputations, it can increase the efficiency and reproducibility of the results.

18. Strategies for using MICE 18.4. What to do with large datasets?

In imputation, generally the advice is to include as much information as possible in the imputation models. Using a large number of predictor variables makes the MAR assumption more plausible (and, hence, reduces bias due to MNAR missingness) and can reduce uncertainty about the missing values.

This can work well in small or medium sized datasets (20 – 30 separate variables, i.e. without interactions, variables derived from others, ...) however, in large datasets (contain hundreds or thousands of variables) this is not feasible.[17]

18.4. What to do with large datasets?

For large datasets a possible strategy is to

- Include all variables used in the analysis model(s) (including the outcome!).
- Include auxiliary variables if they are **strong predictors of missingness**.
- Include auxiliary variables if they have strong associations with the incomplete variables.
- Use auxiliary variables only if they do not have too many missing values themselves (and are observed for most of the incomplete cases of the variable of interest).
- Use **auxiliary variables** only in those imputation models for which they are **relevant** (and exclude them for others using the predictor matrix).
- Calculate summary scores from multiple items referring to the same concept and use the summary score as predictor variable.

18.5. How much missing is too much?

There is **no clear cut-off** for the proportion of missing values that can be handled adequately using MICE (or any other imputation method).

The amount of missingness that can be handeled **depens on the information that is available** to impute it.

- Are there strong predictor variabels available & observed?
- Are there sufficient observed cases to get reliable estimates for the predictive distribution?

18.5. How much missing is too much?

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- Are there strong predictor variabels available & observed?
- Are there sufficient observed cases to get reliable estimates for the predictive distribution?

Example:

- In a set of N = 50 cases, 50% missing values leave 25 cases to estimate the parameters of the predictive distribution.
- In a large set of N = 5000 subjects, 50% missing cases leave 2500 observed cases to estimate parameters.

18. Strategies for using MICE 18.6. Imputation of outcomes

Usually, missing outcome values are not imputed.

Why?

When there are no auxiliary variables, imputation and analysis model are equal.

- Parameters of the imputation model are estimated on observed cases of the outcome.
- Imputed values will fit the assumed model perfectly.
- Including imputed cases in the analysis does not add any information.

Exception:

- When very strong auxiliary variables are available.
- Outcomes may be imputed when one imputation is performed for several analysis models, because not imputing the outcome(s) would mean
 - excluding cases with missing outcome(s) from the imputation, or
 - excluding the outcome variable(s) as predictor(s).

18.7. Notes of caution & things to keep in mind

Multiple imputation is **not** a **quick and easy solution for missing data**. It requires **care and knowledge** about

- the data to be imputed (and the context of the data),
- the statistical method used for imputation, and
- the **software** implementation used.

Moreover

- Never accept default settings of software blindly.
- Question the plausibility of the MAR assumption. If it is doubtful, use sensitivity analysis.

18.7. Notes of caution & things to keep in mind

Remember:

- Use as much information as possible
 - include all covariates and the outcome
 - use auxiliary information
 - use the most detailed version of variables if possible
- Avoid feedback from derived variables to their originals.
- Imputation models must fit the data
 (correct assumption of error distribution and functional forms and possible
 interactions of predictor variables).
- Think carefully how to handle variables that are derived from other variables.
- Consider the impact the **visit sequence** may have.
- Choose an appropriate number of imputations.
- Make sure the imputation algorithm has converged.
- Use common sense when evaluating if the imputed values are plausible.

19. MICE and MI in the bigger picture

19.1. Other R packages that do imputation

In Part III of this course (and the second practical) we have worked with some R packages that perform imputation or provide functionality for missing data other than **mice**.

Currently, there are 217 packages available on CRAN that use the word "missing" in either the titel or description of the package, 132 that use either "impute" or "imputation" and 47 that use the word "incomplete".

Not all of these packages perform imputation or are usefull for our purposes, but even if we excluded those packages, the number of useful packages for dealing with missing data would still be to large to mention them all.

→ The mice package is often a good option, but certainly not the only option to perform imputation!

19. MICE and MI in the bigger picture

19.2. Imputation in other software

In this second half of the course, we have focused on (multiple) imputation using R.

Naturally, R is not the only statistical software that can perform multiple imputation.

- Stata, SAS and MPLUS provide packages/functions to perform multiple imputation and pool the results.
- There are macros and additional packages available, e.g., smcfcs is implemented for Stata as well
- SPSS provides some functionality to perform MI

19. MICE and MI in the bigger picture

19.3. Other approaches to handle missing values

Finally, we should not forget that MICE is not the only method to handle missing values.

Besides MICE, multiple imputation can be performed in a joint model approach (as for instance implemented in the R package jomo).

Furthermore,

- direct likelihood methods,
- fully Bayesian methods(as implemented in JointAI), or
- weighted estimating equations

are valid alternative approaches when data are incomplete.

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