Class 3: More complex forms of missing data

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In this class . . .

- Revision of last week
- A bit more on Bayesian models and output
- Some different types of missingness: longitudinal data analysis and time series analysis
- ▶ Not missing at random models: pattern mixture models and selection models

A reminder of terms

- Missing completely at random (MCAR) means the cause of the missingness was completely unrelated to the data.
- Missing at random (MAR) means that the missingness only depends on the observed data. Given the observed information, the data are MCAR.
- Not missing at random (NMAR or MNAR) means the missingness depends on unobserved data that we do not have.

If you are in an MCAR or MAR situation where the parameters ψ and θ are not linked, the missingness is known as **ignorable**

Missing data analysis mathematical notation

Let:

- Y be the variables we are interested in, as a matrix of n observations and p variables
- \triangleright Y_{obs} the observed data
- Y_{mis} the missing data
- M an n×p matrix that defines which observations/variables are missing, with m_{ij} = 1 if observation i and variable j are missing, and 0 otherwise
- $\blacktriangleright \ \psi$ some parameters governing the missing data mechanism

Now:

MCAR means

$$P(m = 1 | Y_{obs}, Y_{mis}, \psi) = P(m = 1 | \psi)$$

MAR means

$$P(m = 1 | Y_{obs}, Y_{mis}, \psi)$$

$$= P(m = 1 | Y_{obs}, \psi)$$

NMAR means

$$P(m = 1 | Y_{obs}, Y_{mis}, \psi)$$

depends on Y_{mis}

Reminder of JAGS code

```
model code ='
model {
  # Likelihood
  for(i in 1:N) {
    y[i] ~ dnorm(intercept + slope*x[i], residual_sd^-2)
  }
  # Priors
  intercept ~ dnorm(0, 10^{-2})
  slope ~ dnorm(0, 10^{-2})
  residual_sd ~ dunif(0,10)
```

Running a JAGS model

```
library(R2jags)
mod_par = c("intercept",
             "slope",
             "residual sd")
model run = jags(data =
                   list(N = length(y),
                        y = y, x = x),
                 parameters.to.save =
                   mod par,
                 model.file =
                   textConnection(model_code))
```

JAGS for binary logistic regression

```
model code = '
model
  # Likelihood
  for (i in 1:N) {
    y[i] ~ dbern(p[i])
    logit(p[i]) = alpha + beta * x[i]
  }
  # Priors
  alpha ~ dnorm(0, 5^{-2})
  beta ~ dnorm(0, 5^{-2})
```

Example for missing data

```
library(mice)
y = as.integer(is.na(nhanes$chl))
x = nhanes$age
plot(jitter(x, 0.1),jitter(y, 0.1))
```



jitter(x, 0.1)

Running the JAGS model

Looking at the output

plot(model_run)



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A quick note on JAGS output

- There are three key parts of the algorithm that affect how good the posterior samples are:
 - 1. The starting values you chose. If you chose bad starting values, you might need to discard the first few thousand iterations. This is known as the *burn-in* period
 - 2. The way you choose your new parameter values. If they are too close to the previous values the MCMC might move too slowly so you might need to *thin* the samples out by taking e.g. every 5th or 10th iteration
 - 3. The total number of iterations you choose. Ideally you would take millions but this will make the run time slower

Plotting the output

You can plot the iterations for all the parameters with traceplot, or for just one with e.g.

```
post = model run$BUGSoutput$sims.list
plot(post$alpha, type = '1')
    2
post$alpha
    0
                     500
                                  1000
                                               1500
                                                           2000
                                                                        2500
                                                                                    3000
                                              Index
```

A good trace plot will show no patterns or runs, and will look like it has a stationary mean and variance

How many chains?

- Beyond increasing the number of iterations, thinning, and removing a burn-in period, JAGS automatically runs *multiple chains*
- This means that they start the algorithm from 3 or 4 different sets of starting values and see if each *chain* converges to the same posterior distribution
- If the MCMC algorithm has converged then each chain should have the same mean and variance.
- ▶ JAGS reports the Rhat value, which is close to 1 when all the chains match
- It's about the simplest and quickest way to check convergence. If you get Rhat values above 1.1, run your MCMC for more iterations

Missingness in longitudinal data

library(nlme)
head(Orthodont)

| ## | Gı | rouped Dat | ca: d | distance | ~ age | Ι | Subje |
|----|----|------------|-------|----------|-------|---|-------|
| ## | | distance | age | Subject | Sex | | |
| ## | 1 | 26.0 | 8 | MO1 | Male | | |
| ## | 2 | 25.0 | 10 | MO1 | Male | | |
| ## | 3 | 29.0 | 12 | MO1 | Male | | |
| ## | 4 | 31.0 | 14 | MO1 | Male | | |
| ## | 5 | 21.5 | 8 | MO2 | Male | | |
| ## | 6 | 22.5 | 10 | MO2 | Male | | |

These data are distance measures from x-ray images of the skull. We also have age, sex, and repeated individual measurements



A simple model for these data

Fit a mixed effects (hierarchical) model of the form:

$$\begin{split} \mathsf{distance}_{ij} &\sim \textit{N}(\alpha_j + \beta \times (\mathsf{age}_{ij} - \mathsf{a\bar{g}e}), \sigma^2) \\ &\alpha_j &\sim \textit{N}(\mu_\alpha, \sigma_\alpha^2) \end{split}$$

where:

- \blacktriangleright distance_{ij} is the distance measurement for observation i for individual j
- age_{ii} is the age associated observation i for individual j
- α_j is a random intercept term for each individual j with some overall mean μ_{α} and a variability between individuals σ_{α}

```
A JAGS model
   model_code = '
   model
     # Likelihood
     for (i in 1:N) {
       y[i] ~ dnorm(alpha[person[i]] + beta * (age[i] - mean(age)), sigma^-2)
     7
     # Prior for intercept
     for(j in 1:N_people) {
       alpha[j] ~ dnorm(mu_alpha, sigma_alpha^-2)
     }
     # Priors on other parameters
     mu alpha ~ dnorm(0, 100^{-2})
     beta ~ dnorm(0, 10^{-2})
     sigma ~ dgamma(1,1)
     sigma alpha ~ dgamma(1,1)
```

Running this model

Output

plot(model_run)



Bugs model at "5", fit using jags, 3 chains, each with 2000 iterations (first 1000 discarded)

What if some of the data are missing?

- Some distance measurements or ages might be missing. (Subject being missing seems unlikely but possible)
- These might just be individual values (*item non-response* or *analysis dropout*), possibly MCAR or MAR
- We might get treatment discontinuation where some subjects no longer take part. This gets us back to a monotone missingness pattern
- We might get an alternative form of dropout if growth is irregular and subjects are removed form the study - this might be an example of MNAR
- All of these are pretty easy to model in JAGS (especially if MAR)

Adding missing values

Set a large number of the values across the data set to missing

set.seed(123)
Orthodont2 = Orthodont
Orthodont2\$distance[sample(1:
 nrow(Orthodont2), 40)] = NA
Orthodont2\$age[sample(1:
 nrow(Orthodont2), 20)] = NA

These data are distance measures from x-ray images of the skull. We also have age, sex, and repeated individual measurements

plot(Orthodont2)



```
New model
   model code =
   model
   ł
     # Likelihood
     for (i in 1:N) {
       y[i] ~ dnorm(alpha[person[i]] + beta * (age[i] - mean(age)), sigma<sup>-2</sup>)
     }
     # Prior for intercept
     for(j in 1:N_people) {
       alpha[i] ~ dnorm(mu alpha, sigma alpha<sup>-2</sup>)
     }
     # Prior for missing values
     for(k in 1:N_miss_age) {
       age[miss[k]] ~ dunif(min_age, max_age)
     }
     # Priors on other parameters
     mu alpha ~ dnorm(0, 100^{-2})
     beta ~ dnorm(0, 10^{-2})
     sigma ~ dgamma(1,1)
```

Running the model

```
mod par = c("mu alpha", "alpha", "beta", "sigma",
             "sigma alpha", "y", "age")
model run = jags(data =
                   list(N = nrow(Orthodont2)).
                        N_people = length(unique(Orthodont2$Subject)),
                        N miss age = sum(is.na(Orthodont2$age)),
                        miss = which(is.na(Orthodont2$age)),
                        v = Orthodont2 distance.
                        age = Orthodont2 age.
                        min age = min(Orthodont2$age, na.rm = TRUE),
                        max age = max(Orthodont2$age, na.rm = TRUE),
                        person = Orthodont2$Subject),
                 parameters.to.save = mod par,
                 model.file = textConnection(model code))
```

Output

plot(model_run)



Bugs model at "6", fit using jags, 3 chains, each with 2000 iterations (first 1000 discarded)

Missing data in time series models

▶ In a time series analysis we usually have a model of the form:

$$y_t = f(y_t^-, X_t) + \epsilon_t$$

where y_t^- represents previous values of the time series and X represents additional covariates

- The response y_t might be multivariate
- Missingness might occur in y, in X, or even in t
- Some tricks:
 - If you have many missing observations you could move to a continuous time series model
 - JAGS (and Stan) will usually work fine if there are missing values in the time series without any change to the model
 - ▶ With missing X values the approach on the previous slides will work fine
 - With missing t values this starts to get fiddly, but it's still possible provided you can put a prior distribution on t

The imputeTS package

There is a nice R package for single imputation of time series called imputeTS

> It will impute the missing values in y_t via various methods:

na_kalman Missing Value Imputation by Kalman Smoothing
 na_locf Missing Value Imputation by Last Observation Carried Forward
 na_ma Missing Value Imputation by Weighted Moving Average
 na_seadec Seasonally Decomposed Missing Value Imputation
 na_seasplit Seasonally Splitted Missing Value Imputation

Running the stochastic methods multiple times might allow for multiple imputation

More details here: https://steffenmoritz.github.io/imputeTS/

Back to MNAR

- For MNAR data we need to build a model that accounts for the missingness mechanism as well as the prediction model
- There are two popular approaches selection models and pattern-mixture models
- > The two approaches correspond to the way in which we factor the joint distribution
- For regression models, we now need to consider the full data set $y = (Y_{obs}, Y_{mis})$ so we are interested in finding the likelihood:

$$p(y, M|\theta, \psi, x) = \prod_{i=1}^{n} p(y_i, m_i|\theta, \psi, x_i)$$

Selection and pattern mixture models

- We are going to factor this model two different ways:
- ► The *Selection* model way:

 $p(y_i, m_i | \theta, \psi, x_i) = p(y_i | x_i, \theta) \times p(m_i | x_i, y_i, \psi)$

Here the first term is just a standard regression model, and the second term is a binary regression which has y as a covariate (with x) in the model. Both models have to be fitted at the same time

► The *Pattern mixture* model way:

$$p(y_i, m_i | \theta, \psi, x_i) = p(y_i | x_i, m_i, \theta) \times p(m_i | x_i, \psi)$$

Now the first distribution is a regression model that is dependent on whether the data are missing or not, and the second is just a very simple logistic regression model based on covariates x. This usually involves fitting **only** the regression model because m_i is data

Here's a simple example. Suppose we define a pattern mixture model as:

$$y_i|m_i, x_i, \theta \sim \mathcal{N}(\alpha^{(m_i)} + \beta^{(m_i)}x_i, (\sigma^{(m_i)})^2)$$

This means that there are different regression parameters for whether the data is missing or not. Of course, the problem is that you do not have any data y_i when the data is missing $(m_i = 1)!!!$

We need extra assumptions to fit this model

Here's the same example as a selection model

 $y_i | x_i, \theta \sim N(\alpha + \beta x_i, \sigma^2)$ $m_i | x_i, y_i, \psi \sim Binom(1, p_i), \text{ logit}(p_i) = \gamma + \delta x_i + \omega y_i$

This means that the coefficients of the binary missingness indicator will also be hard to estimate if there is a high degree of missingness in y_i

(The probit version of this model is often known as a Heckman model)

Some notes on pattern mixture and selection models

- These models can extend to multivariate y values exactly as in the multiple imputation scenarios we met before.
- You can create hybrid versions of the two (called pattern-set mixture models) but this gets fiddly
- It's called a pattern mixture model because it's essentially a mixture of two regression models (one for the missing and one for the not missing data)
- ▶ I tend to find selection models easier to understand (and fit). Also if $\omega = 0$ in the previous slide then you end up back at MAR!
- It's hard to compare between the two approaches in most real-world scenarios

Some simplifications

A simpler version of the pattern mixture model is:

$$y_i|m_i, x_i, \theta \sim N(\alpha^{(m_i)} + \beta x_i, \sigma^2)$$

... so only a changing intercept between missing and non-missing data

- ► This means that a simple statistic such as α^(m₁) − α^(m₀) will tell you how the missing data are shifted up or down compared to the observed data
- Little and Rubin argue that the pattern mixture model is better because the interpretation of the parameter ω in the selection model is harder, and because imputed data values are easier to simulate from
- However it needs strong assumptions on the missing intercept to fit properly

Including extra assumptions

These models often need extra assumptions to fit well. These might include:

- Extra (smaller) data sets on missing values, such as following up non-response
- Imposing restrictions on the model parameters
- Being Bayesian, and putting prior distributions on the parameters

Bayesian approaches to pattern mixture models

- A common approach is to put a prior distribution on the regression parameters for the missing observations
- ▶ In the simpler intercept only example we have:

$$y_i | m_i, x_i, \theta \sim N(\alpha^{(m_i)} + \beta x_i, \sigma^2)$$

$$\alpha^{(m_1)} \sim N(\alpha^{(m_0)}, \sigma_{\alpha}^2)$$

This means it is centered around the non-missing intercept
 The parameter σ_α might also have a prior distribution
 It may also be set to be a function of α^(m₀), e.g. σ_α = kα^(m₀)

```
Example: a selection model in JAGS
   model_code =
   model
     # Likelihood
     for (i in 1:N) {
       y[i] ~ dnorm(alpha + beta * x[i], sigma<sup>-2</sup>)
       m[i] ~ dbern(p[i])
       logit(p[i]) = gamma + omega * (y[i] - mean(y))
     }
     # Priors for regression model
     alpha ~ dnorm(0, 100^{-2})
     beta ~ dnorm(0, 100^{-2})
     sigma ~ dgamma(1, 1)
     # Priors for missingness model
     gamma ~ dnorm(0, 5^{-2})
     omega ~ dnorm(0, 5^{-2})
```

Some selection model NMAR data

```
set.seed(123)
N = 100
x = sort(runif(N))
alpha = 3
beta = 2
sigma = 0.2
y_true = rnorm(N, alpha + beta * x,
               sigma)
gamma = -0.5
omega = 1
m = rbinom(N, 1, plogis(gamma +
      omega * (v true -
                 mean(y_true))))
y = y true
v[m==0] = NA
```

plot(x, y_true) points(x, y, col = 'red', pch = 19)9 4.5 9 3.5 0 0.0 0.2 0.4 0.9

Output



Bugs model at "4", fit using jags, 3 chains, each with 2000 iterations (first 1000 discarded)

How well did it estimate the model parameters



And now a pattern-mixture model

```
model code = '
model
  # Likelihood
  for (i in 1:\mathbb{N}) {
    # Note: m[i] = 1 if not-missing and m[i] = 2 if missing
    y[i] ~ dnorm(alpha[m[i]] + beta * x[i], sigma<sup>-2</sup>)
  }
  # Priors for regression model
  alpha[1] \sim dnorm(0, 100^{-2})
  # This is the key prior
  alpha[2] ~ dnorm(alpha[1] + 0.5, sigma alpha^-2)
  beta ~ dnorm(0, 100^{-2})
  sigma ~ dgamma(1, 1)
  sigma alpha ~ dgamma(1, 1)
```

Some pattern mixture model NMAR data

```
set.seed(123)
N = 100
x = sort(runif(N))
gamma = -0.5
delta = 3
m = rbinom(N, 1, plogis(gamma +
      delta * (x - mean(x))) + 1
alpha = c(3, 3.5)
beta = 2
sigma = 0.2
y_true = rnorm(N, alpha[m] +
          beta * x, sigma)
y = y true
v[m=2] = NA
```

plot(x, y_true)
points(x, y, col = 'red', pch = 19)



Output



Bugs model at "5", fit using jags, 3 chains, each with 2000 iterations (first 1000 discarded)

How well did it estimate the model parameters



Non-ignorable missing data in mice

- ▶ The mice package also allows for some NMAR type multiple imputation
- ▶ There is another package called miceMNAR that specialises in this
- It requires you to specify the regression-type model simultaneously with the imputation
- > As before, it uses the fully conditional specification type model
- There are several different functions including mice.impute.ri and mice.impute.mnar.logreg
- Most of the functions seem to contain examples
- This is very new (only implemented in 2020)



- Missing not at random data is harder to fit because it usually requires some extra assumptions about the missingness mechanism and makes the model more complicated
- We have met the two main types; selection models and pattern mixture models
- They differ in how they decompose the likelihood, and how the results can be interpreted
- ▶ All these models can be fitted in JAGS and Stan; a version can be fitted in mice