Session 6

Coping with missing data

Matt Denwood 2021-06-30

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MNAR: Missing not at random

- There is an unknown (or unrecorded) pattern to the missingness
- It is therefore possible that the prevalence is confounded with missingness

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- There is no possibility of being confounded with prevalence

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- The animal was too aggressive to facilitate a blood sample
- Somebody dropped the samples on the way to the lab

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- Exclude individuals with incomplete data
- Allow template_huiwalter to adjust the model code

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This is a relatively rare kind of missingness, but it does happen

Missing samples occur due to a known pattern

 We can (and must) assess if this is likely be correlated with prevalence

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- Test B was only done if Test A was positive

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- No -> treat as MCAR
- Yes -> we must model the confounding

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- Test A was not done in population 1 because of costs
- Test B was only done if Test A was positive

Solution depends on whether the the missigness is potentially confounded with prevalence

- No -> treat as MCAR
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Very common type of missingness in practice

Missing samples occur due to an unknown/unrecorded pattern

• We must assume that this might be correlated with prevalence

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Examples:

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- Some patients choose to have Test B after knowing the result of Test A

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- Test B was only done if the animal had (unrecorded) diarrhea
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- Exclude segments of the data that may be affected by structural missingness
- Give up and collect a better dataset

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Examples:

- Test B was only done if the animal had (unrecorded) diarrhea
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- Exclude segments of the data that may be affected by structural missingness
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A common type of missingness in secondary data

Missingness and template Hui-Walter

We can simulate MCAR data as follows:

```
set.seed(2021-06-30)
# Parameter values to simulate:
N < -1000
sensitivity <-c(0.8, 0.9, 0.95)
specificity \leftarrow c(0.95, 0.99, 0.95)
Populations <- 2
prevalence \leftarrow c(0.25,0.5)
data <- tibble(Population = sample(seq len(Populations), N,

    replace=TRUE)) %>%

 mutate(Status = rbinom(N, 1, prevalence[Population])) %>%
 mutate(Test1 = rbinom(N, 1, sensitivity[1]*Status +
 mutate(Test2 = rbinom(N, 1, sensitivity[2]*Status +
 mutate(Test3 = rbinom(N, 1, sensitivity[3]*Status +
 select(-Status)
```

Now introduce missingness in all 3 tests:

```
missingness < c(0.1, 0.2, 0.3)
data <- data %>%
 mutate(Test1 = case when(
   rbinom(n(), 1, missingness[1]) == 1L ~ NA_integer_,
   TRUE ~ Test1
 )) %>%
 mutate(Test2 = case when(
   rbinom(n(), 1, missingness[2]) == 1L ~ NA_integer_,
   TRUE ~ Test2
 )) %>%
 mutate(Test3 = case_when(
   rbinom(n(), 1, missingness[3]) == 1L ~ NA_integer_,
   TRUE ~ Test3
 ))
```

```
data %>% count(Missing1 = is.na(Test1), Missing2 = is.na(Test2),
## # A tibble: 8 x 4
## Missing1 Missing2 Missing3
                       n
## <lgl> <lgl> <int>
## 1 FALSE FALSE FALSE 513
## 2 FALSE FALSE TRUE 210
## 3 FALSE TRUE FALSE 126
## 4 FALSE TRUE TRUE
                      56
## 5 TRUE FALSE FALSE 54
## 6 TRUE FALSE TRUE
                        20
## 7 TRUE
         TRUE FALSE
                        14
## 8 TRUE
         TRUE
                TRUE
```

We can simply feed this data to template_huiwalter:

What does that look like...?

```
model{
    ## Observation layer:

# Complete observations (N=513):
    for(p in 1:Populations){
        Tally_RRR[1:8,p] ~ dmulti(prob_RRR[1:8,p], N_RRR[p])

        prob_RRR[1:8,p] <- se_prob[1:8,p] + sp_prob[1:8,p]
}</pre>
```

```
# Partial observations (Test1: Recorded, Test2: Missing, Test3:
\hookrightarrow Missing; N=56):
for(p in 1:Populations){
    Tally RMM[1:2,p] ~ dmulti(prob RMM[1:2,p], N RMM[p])
    prob_{RMM}[1:2,p] \leftarrow se_{prob}[c(1,2),p] + sp_{prob}[c(1,2),p] +
                           se prob[c(3,4),p] + sp prob[c(3,4),p] +
                           se prob[c(5,6),p] + sp prob[c(5,6),p] +
                           se_{prob}[c(7,8),p] + sp_{prob}[c(7,8),p]
}
# Partial observations (Test1: Recorded, Test2: Recorded, Test3:
\hookrightarrow Missing; N=210):
for(p in 1:Populations){
    Tally_RRM[1:4,p] ~ dmulti(prob_RRM[1:4,p], N_RRM[p])
    prob RRM[1:4,p] <- se prob[c(1,2,3,4),p] +
    \hookrightarrow sp_prob[c(1,2,3,4),p] +
                           se_prob[c(5,6,7,8),p] +
                           \hookrightarrow sp_prob[c(5,6,7,8),p]
```

```
# Partial observations (Test1: Missing, Test2: Recorded, Test3:
for(p in 1:Populations){
    Tally MRR[1:4,p] ~ dmulti(prob MRR[1:4,p], N MRR[p])
    prob_MRR[1:4,p] \leftarrow se_prob[c(1,3,5,7),p] +
    \hookrightarrow sp prob[c(1,3,5,7),p] +
                        se prob[c(2,4,6,8),p] +
                        \hookrightarrow sp_prob[c(2,4,6,8),p]
}
# Partial observations (Test1: Missing, Test2: Recorded, Test3:
for(p in 1:Populations){
    Tally_MRM[1:2,p] ~ dmulti(prob_MRM[1:2,p], N_MRM[p])
    prob MRM[1:2,p] <- se prob[c(1,3),p] + sp prob[c(1,3),p] +
                        se prob[c(2,4),p] + sp prob[c(2,4),p] +
                        se_prob[c(5,7),p] + sp_prob[c(5,7),p] +
                        se_prob[c(6,8),p] + sp_prob[c(6,8),p]
```

NB: MMM combinations have been removed!

```
## Observation probabilities:
for(p in 1:Populations){
    # Probability of observing Test1- Test2- Test3- from a true
    → positive::
    se_{prob}[1,p] \leftarrow prev[p] * ((1-se[1])*(1-se[2])*(1-se[3])

→ +covse12 +covse13 +covse23)

    # Probability of observing Test1- Test2- Test3- from a true
    → negative::
    sp_prob[1,p] \leftarrow (1-prev[p]) * (sp[1]*sp[2]*sp[3] +covsp12
    \hookrightarrow +covsp13 +covsp23)
    # Probability of observing Test1+ Test2- Test3- from a true
    → positive::
    se_{prob}[2,p] \leftarrow prev[p] * (se[1]*(1-se[2])*(1-se[3]) -covse12
    \hookrightarrow -covse13 +covse23)
    # Probability of observing Test1+ Test2- Test3- from a true
    → negative::
    sp_prob[2,p] \leftarrow (1-prev[p]) * ((1-sp[1])*sp[2]*sp[3] -covsp12
    \hookrightarrow -covsp13 +covsp23)
    # Probability of observing Test1- Test2+ Test3- from a true
    \hookrightarrow positive::
    se_prob[3,p] \leftarrow prev[p] * ((1-se[1])*se[2]*(1-se[3]) -covse12

→ +covse13 -covse23)

    # Probability of observing Test1- Test2+ Test3- from a true
```

```
## Data:
data{
"Populations" <- 2
"N RRR" <- c(233, 280)
"Tally_RRR" <- structure(c(148, 8, 1, 2, 9, 4, 11, 50, 133, 3, 4, 5, 8,
\rightarrow 9, 20, 98), .Dim = c(8, 2))
"N RMR" <- c(65, 61)
"Tally_RMR" <- structure(c(51, 3, 1, 10, 29, 2, 11, 19), .Dim = c(4, 2))
"N RMM" <- c(22, 34)
"Tally_RMM" \leftarrow structure(c(16, 6, 20, 14), .Dim = c(2, 2))
"N RRM" <- c(100, 110)
"Tally_RRM" \leftarrow structure(c(74, 5, 2, 19, 58, 10, 5, 37), .Dim = c(4, 2))
"N MRR" \leftarrow c(27, 27)
"Tally_MRR" <- structure(c(18, 1, 4, 4, 15, 2, 1, 9), .Dim = c(4, 2))
"N MRM" <- c(10, 10)
"Tally MRM" <- structure(c(7, 3, 4, 6), .Dim = c(2, 2))
"N MMR" <- c(6, 8)
"Tally MMR" \leftarrow structure(c(4, 2, 2, 6), .Dim = c(2, 2))
}
```

What about other types of missing?

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MNAR:

- Solution depends entirely on the problem
- And sometimes there is no solution...

But remember: bigger datasets are not always better datasets...

Making your data missing

What happens if we eliminate:

- One population at a time (where we have >2)?
- One test at a time (where we have >2)?
- Do the results change?

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What happens if we eliminate:

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- One test at a time (where we have >2)?
- Do the results change?

If we have >2 populations *and* >2 tests then we can eliminate one combination at a time!

• This is a very useful form of cross-validation

Estimating the full model:

```
template_huiwalter(data, "model_full.txt")
results_full <- run.jags("model_full.txt")
## Loading required namespace: rjags
# Check convergence etc:
# plot(results_full)
# results_full

summary_full <- summary(results_full, vars="^s") %>%
    as.data.frame() %>%
    rownames_to_column("Parameter") %>%
    mutate(Model = "Full") %>%
    select(Model, Parameter, Median, Lower95, Upper95)
```

How can we make a specific population missing?

```
crossval_data <- data %>%
  filter(Population != 1)

template_huiwalter(crossval_data, "model_mp1.txt")
results_crossval <- run.jags("model_mp1.txt")
summary_crossval <- summary(results_crossval, vars="^s") %>%
  as.data.frame() %>%
  rownames_to_column("Parameter") %>%
  mutate(Model = "MP1") %>%
  select(Model, Parameter, Median, Lower95, Upper95) %>%
  bind_rows(summary_full) %>%
  arrange(Parameter, Model)
```

```
summary crossval
##
     Model Parameter Median
                              Lower95 Upper95
## 1 Full se[1] 0.8318452 0.7846137 0.8766986
## 2
    MP1 se[1] 0.8242777 0.7665977 0.8826820
## 3
     Full
               se[2] 0.9026765 0.8564078 0.9441426
## 4 MP1
          se[2] 0.8936380 0.8370178 0.9436194
## 5
    Full se[3] 0.9428753 0.9064733 0.9743135
## 6
     MP1
               se[3] 0.9421457 0.8982132 0.9793507
## 7
     Full
               sp[1] 0.9602720 0.9384173 0.9798948
## 8
     MP1
               sp[1] 0.9713449 0.9383363 0.9976758
## 9
      Full
               sp[2] 0.9888668 0.9741214 0.9999982
## 10
     MP1
               sp[2] 0.9769023 0.9479276 0.9999949
## 11
      Ful1
               sp[3] 0.9488934 0.9221615 0.9737997
## 12
     MP1
               sp[3] 0.9455734 0.8991737 0.9884249
```

How many combinations of test missingness and population do we have?

```
all_combinations <- data %>%
 pivot_longer(-Population, names_to = "Test", values_to = "Result") %>%
 filter(!is.na(Result)) %>%
 count(Population, Test) %>%
 print()
## # A tibble: 6 x 3
    Population Test
##
         <int> <chr> <int>
##
## 1
            1 Test1 420
## 2
            1 Test2 370
## 3
            1 Test.3 331
## 4
          2 Test1 485
## 5
          2 Test2 427
          2 Test3
## 6
                     376
```

How can we make a specific combination of test and population missing?

```
all results <- vector('list', length=nrow(all combinations))
all summary <- vector('list', length=nrow(all combinations))</pre>
crossval data <- data %>%
  mutate(Test1 = case when(
    Population == 1 ~ NA integer ,
    TRUE ~ Test1
  ))
template huiwalter(crossval data, "model mc11.txt")
all_results[[1]] <- run.jags("model_mc11.txt")</pre>
# Assess convergence and sample size!
all_summary[[1]] <- summary(all_results[[1]], vars="^s") %>%
  as.data.frame() %>%
  rownames_to_column("Parameter") %>%
  mutate(Model = "MC11") %>%
  select(Model, Parameter, Median, Lower95, Upper95)
```

```
crossval data <- data %>%
  mutate(Test2 = case_when(
    Population == 1 ~ NA_integer_,
    TRUE ~ Test2
  ))
template_huiwalter(crossval_data, "model_mc12.txt")
all_results[[2]] <- run.jags("model_mc12.txt")</pre>
# Assess convergence and sample size!
all_summary[[2]] <- summary(all_results[[2]], vars="^s") %>%
  as.data.frame() %>%
  rownames to column("Parameter") %>%
  mutate(Model = "MC12") %>%
  select(Model, Parameter, Median, Lower95, Upper95)
```

```
crossval data <- data %>%
  mutate(Test2 = case_when(
    Population == 1 ~ NA_integer_,
    TRUE ~ Test2
  ))
template_huiwalter(crossval_data, "model_mc12.txt")
all_results[[2]] <- run.jags("model_mc12.txt")</pre>
# Assess convergence and sample size!
all_summary[[2]] <- summary(all_results[[2]], vars="^s") %>%
  as.data.frame() %>%
  rownames to column("Parameter") %>%
  mutate(Model = "MC12") %>%
  select(Model, Parameter, Median, Lower95, Upper95)
```

Are there any substantial disagreements:

```
bind_rows(list(summary_full, all_summary)) %>% arrange(Parameter, Model)
##
     Model Parameter
                        Median
                               Lower95
                                           Upper95
## 1
     Full
               se[1] 0.8318452 0.7846137 0.8766986
## 2
     MC11
               se[1] 0.8207408 0.7624440 0.8755579
## 3
     MC12
               se[1] 0.8262436 0.7698624 0.8812174
## 4
      Full
               se[2] 0.9026765 0.8564078 0.9441426
## 5
     MC11
               se[2] 0.8941601 0.8405847 0.9445226
## 6
     MC12
               se[2] 0.8952849 0.8393142 0.9443727
## 7
      Full
               se[3] 0.9428753 0.9064733 0.9743135
## 8
     MC11
               se[3] 0.9455431 0.9071161 0.9799713
## 9
     MC12
               se[3] 0.9388186 0.8923919 0.9793301
## 10 Full
               sp[1] 0.9602720 0.9384173 0.9798948
## 11
      MC11
               sp[1] 0.9706544 0.9388596 0.9982201
## 12 MC12
               sp[1] 0.9648173 0.9394740 0.9900894
## 13 Full
               sp[2] 0.9888668 0.9741214 0.9999982
## 14 MC11
               sp[2] 0.9852388 0.9673050 0.9999757
## 15 MC12
               sp[2] 0.9771450 0.9487679 0.9999897
## 16 Full
               sp[3] 0.9488934 0.9221615 0.9737997
## 17 MC11
               sp[3] 0.9502358 0.9168171 0.9819010
## 18 MC12
               sp[3] 0.9526880 0.9190077 0.9875565
```

Practical session 6

Points to consider

- 1. How does MCAR data impact your results?
- 2. What about if you have data using confirmatory tests?
- 3. How can we use cross-validation as a method of checking assumptions?

Summary

- Observations that are MCAR are trivial to deal with using JAGS
- We can also treat MAR observations as if they are MCAR as long as the reason for missingness does not confound with expected prevalence, or we allow prevalence to differ between groups where the structural missingness differs
- MNAR is bad news
- Deliberately making observations missing is a good way to assess model assumptions