# **Imputation Diagnostics**

Utrecht University Winter School: Missing Data in R



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#### Outline

Imputation Model Convergence

Plausibility of Imputed Values



#### **Example Data**

The example data are synthesized from questionnaire data collected by Lang, Salter, and Adams (2009).

- *N* = 87
- P = 33 Likert-type variables assessing:
  - Perceptions and definitions of racism
  - Political affiliation
  - Support for affirmative action policies
  - Belief in meritocratic ideals

#### The data synthesis involved:

- 1. Resampling the original data to produce a new sample of 250 cases
- 2. Adding Gaussian noise
- 3. Imposing 25% MAR missing
  - MAR Predictors = Political Affiliation, Definition of Racism

# **Imputation Diagnostics**

After we run an MI routine, we need to make sure that the procedure has performed as expected.

Problems can arise to two different places:

- 1. The imputation model may fail to converge.
- 2. The imputed values may be invalid.

We need to examine our results to check for these problems.



# Imputation Model Convergence

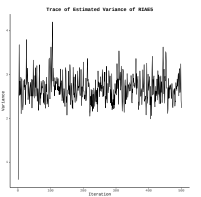
The imputation model is usually estimated through some form of Bayesian simulation.

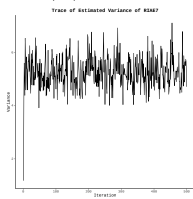
- Gibbs sampled parameters form a Markov Chain.
  - Each draw is dependent on only its immediate predecessor in the chain.
  - $\theta^{(t)}|\theta^{(t-1)}\perp\theta^{(t-j)}$   $\forall j>1$
- Early elements of a Markov chain are similar to the starting values.
  - Samples are poor approximations of the true posterior.
- We must let the sampler iterate for a while to allow the estimates time to separate from their starting values.
  - We call these initial iterations "burn-in" or "warm-up" iterations.

#### Traceplots

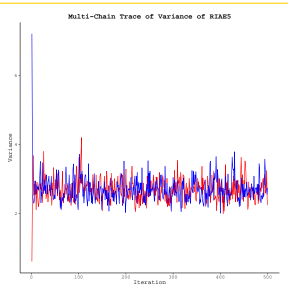
Once converged, each sampled parameter should "bounce" around some equilibrium point.

- The draws will never converge to a single point.
- Deterministic convergence would defeat the purpose of simulation.





# Traceplots



#### Potential Scale Reduction Factor

Suppose we have M length-N Markov chains for the same parameter, heta.

- ullet If these chains have converged, all M chains should be sampling from the same parameter space.
- The pooled *total variance* and the *within-chain* variances should be about the same.

The Gelman and Rubin (1992) Potential Scale Reduction Factor,  $\widehat{R}$ , quantifies this concept:

$$\widehat{R} = \sqrt{\frac{T}{W}}$$

 $\overrightarrow{R}$  will approach 1.0 at convergence.

•  $\widehat{R}$  < 1.1 or 1.2 suggests acceptable convergence.



#### Potential Scale Reduction Factor

$$\widehat{R} = \sqrt{\frac{T}{W}}$$

The total variance, T, is the weighted average of the within-chain variance, W, and the between-chain variance, B.

$$T = \frac{N-1}{N}W + \frac{1}{N}B$$

$$W = \frac{1}{M} \sum_{m=1}^{M} \text{var}(\theta_m)$$

$$B = \frac{N}{M-1} \sum_{m=1}^{M} \left( \bar{\theta}_m - \bar{\theta} \right)^2$$



# Example: Potential Scale Reduction Factor

We can compute  $\hat{R}$  statistics for mice() models using the Rhat.mice() function from the **miceadds** package.

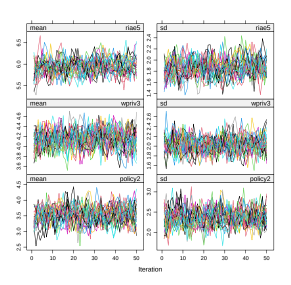
#### More Imputation Model Convergence

A convergent imputation model will produce imputed values that fluctuate around an equilibrium point.

• Imputation model convergence can be assessed indirectly by looking at plots of the item-level sufficient statistics for each imputation.

This approach is automated for **mice** via plot.mice().

# More Imputation Model Convergence



# Imputed Value Plausibility

We need to ensure that the imputations are sensible.

- Imputed values shouldn't be too dissimilar from their observed counterparts.
  - What constitutes too much dissimilarity is subjective and problem-specific.

We can assess dissimilarity graphically or through summary statistics.

- Out-of-bounds values for the imputations are perfectly acceptable.
  - MI is NOT designed to maintain the range.
  - We don't want wildly extreme values, though.
- The means of the observed and imputed components of each variable shouldn't differ too much.
  - Again, how much is too much is subjective.

```
## Fill the missing values with imputations:
impList <- complete(miceOut, "all")</pre>
## Computes means:
rawMeans <- colMeans(missData, na.rm = TRUE)
impMeans <- do.call("rbind", impList) %>% colMeans()
## Compute standard deviations:
rawSds <- sapply(missData, sd, na.rm = TRUE)</pre>
impSds <- lapply(impList, function(x) sapply(x, sd)) %>%
    do.call(rbind. .) %>%
    colMeans()
## Compute ranges:
rawRanges <- sapply(missData, range, na.rm = TRUE)
impRanges <- do.call("rbind", impList) %>% sapply(range)
```

Compare observed and imputation-based means:

```
vars <- grep("policy\\d", colnames(missData))
round(rawMeans[vars], 3)
policy1 policy3 policy4 policy5 policy6 policy2
   3.020   3.724   3.564   3.746   4.483   3.558
round(impMeans[vars], 3)
policy1 policy3 policy4 policy5 policy6 policy2
   3.343   4.010   3.223   3.413   4.435   3.666</pre>
```

Compare observed and imputation-based standard deviations:

```
round(rawSds[vars], 3)

policy1 policy3 policy4 policy5 policy6 policy2
   2.045   2.181   2.035   2.015   1.956   2.237

round(impSds[vars], 3)

policy1 policy3 policy4 policy5 policy6 policy2
   2.259   2.309   2.115   2.167   2.093   2.187
```

Compare observed and imputation-based ranges:

```
round(rawRanges[ , vars], 3)

policy1 policy3 policy4 policy5 policy6 policy2

[1,] -1.383 -1.342 -2.260 -1.106 0.126 -2.221

[2,] 8.641 9.551 9.471 8.885 9.098 10.237

round(impRanges[ , vars], 3)

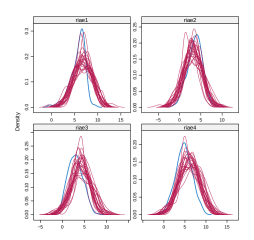
policy1 policy3 policy4 policy5 policy6 policy2

[1,] -3.290 -4.887 -4.717 -2.696 -3.450 -4.241

[2,] 10.751 13.251 9.471 12.595 11.724 12.047
```

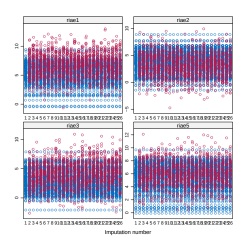
#### **Graphical Imputation Checks**

```
## Overlaid density plots of imputed vs. observed values:
densityplot(miceOut, ~ riae1 + riae2 + riae3 + riae4, layout = c(2, 2))
```



# **Graphical Imputation Checks**

```
## Stripplots of imputed vs. observed values:
stripplot(miceOut, riae1 + riae2 + riae3 + riae5 ~ .imp, layout = c(2, 2))
```



#### References

Gelman, A., & Rubin, D. B. (1992). Inference from iterative simulation using multiple sequences. *Statistical science*, *7*(4), 457–472. Lang, K. M., Salter, P. S., & Adams, G. (2009, April). What drives the relationship between conservatism and racism? A mediation analysis. In *Proceedings of the annual meeting of the Southwestern Psychological Association*.

