# SOC542 Statistical Methods in Sociology II

Missing Data & Model Checking and Robustness

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March 7, 2022

### **Plan**

- Missing data
- ► Model checking
- ► Model robustness

### What is missing data?

- Missing data occurs when we do not have an record of any data for one or more variables for an observation.
- ► This can occur for a variety of reasons including
  - Skip patterns
  - Survey non-response
  - Survey error
  - Data entry errors
  - Data handling errors
- ► The severity of the problem depends on why the data are missing and the amount of missingness

### Missing completely at random (MCAR)

- ► Missing Completely at Random (MCAR)
  - ightharpoonup Probability  $x_i$  is missing is constant across all observations
- Discarding missing cases does not result in any bias

### Missing at random (MAR)

- Missing at Random (MAR)
  - ightharpoonup Probability  $x_i$  is missing depends on observed variables.
- Discarding missing cases does not result in any bias if predictors of missingness are adjusted for.

### Missing not at random

- Missingness that depends on the missing value is considered
   Missing not at Random (MNAR)
  - e.g. Higher income respondents less likely to report income

### Simulating missingness

- We can use simulations to better understand the effects to different kinds of missingness
- Consider the following population model

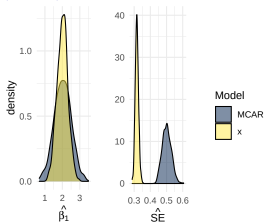
$$z = N(0,1)$$

$$x = 0.5z + N(0,1)$$

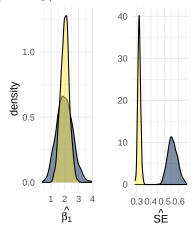
$$y = 2x - 2z + N(0,10)$$

• We can vary the kind of missingess in x and analyze how it effects  $\hat{\beta}_1$  and  $\hat{\sigma}$ .

MCAR, p(Missing) = 0.6

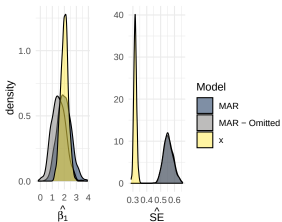


**MAR,** p(Missing) = 0.8 if z > 0

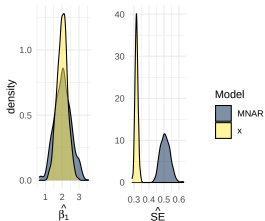




#### MAR, z omitted







### Missingness in practice

- It is often difficult to determine why data are missing
   Knowledge of domain and data generation process helpful
- ► MCAR is a very strong assumption
- ► MAR is more reasonable, but it is difficult to determine whether missingness is due to an unobserved factor
  - Including more predictors in a model helps to reduce concerns

### **Addressing missingness**

- ► Three approaches
  - 1. Delete missing cases
  - 2. Simple imputation
  - 3. Multiple imputation

### **Deleting missing data**

- Complete-case analysis / listwise deletion
  - $\triangleright$  Delete all rows where y or x or z is missing
- ▶ If missingess is not MCAR then results could be biased
- ▶ If many predictors, sample size can reduce substantially

### **Deleting missing data**

- ► Available-case analysis / pairwise deletion
  - Make comparisons where data are available
- In the previous example, one might use the following model to get an estimate of the effect of z on y

$$y = \beta_0 + \beta_1 z + u$$

- ► The variable *x* is ignored, otherwise we drop cases where *x* is missing.
- ▶ One might also estimate the model including x and compare the results.
- ► Like listwise deletion, bias can occur if systematic differences between missing and non-missing cases.

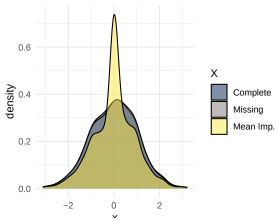
### Simple imputation: Guessing the mean

- ► Instead of removing data, we can try to "guess" the missing values.
- ► A good guess is the mean of the observed data, **mean** imputation.
- ▶ But this can distort the distribution and underestimate the standard deviation.

### Simple imputation

```
X <- rnorm(N)
print(mean(X))
## [1] 0.009998062
print(sd(X))
## [1] 1.020109
X.m <- ifelse(rbinom(N, 1, 1-0.2), X, NA)</pre>
X.g <- ifelse(!is.na(X.m), X, mean(X.m[!is.na(X.m)]))</pre>
print(mean(X.g))
## [1] 0.01318459
print(sd(X.g))
## [1] 0.922013
```

### **Simple imputation**

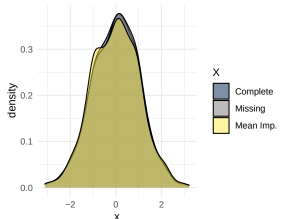


### Simple imputation: Random imputation

- We can address some of the limitations of mean imputation by using the full distribution of the observed variable. This is known as random imputation
- ► However, the imputed values will not necessarily reflect the underlying association between variables and we do not make use of other information.

### Simple imputation: Random imputation

### Simple imputation: Random imputation



- We can improve our imputation model by making use of additional information in other variables.
  - ► For example, we could estimate a model to predict *x* using other covariates and then use these predictions in place of missing vaulues

```
N <- 1000
v <- rnorm(N)
w <- rnorm(N)
x <- 0.5*v + 0.5*w + rnorm(N)
y <- x - 2*w + rnorm(N)

x.m <- ifelse(rbinom(N, 1, 0.4), x, NA) # MCAR

imp.model <- lm(x.m ~ v + w)</pre>
```

	Complete	Listwise	Mean	Random	Predicted
(Intercept)	-0.016	-0.043	-0.016	-0.007	-0.008
	(0.032)	(0.052)	(0.032)	(0.046)	(0.040)
X	0.994	1.034	0.994	0.309	0.998
	(0.030)	(0.048)	(0.030)	(0.040)	(0.052)
w	-2.007	-2.082	-2.007	-1.581	-1.968
	(0.036)	(0.056)	(0.036)	(0.048)	(0.047)
Num.Obs.	1000	390	1000	1000	1000
R2	0.765	0.790	0.765	0.527	0.635
R2 Adj.	0.764	0.788	0.764	0.526	0.634

### **Simple imputation**

- Limitations
  - Each process becomes cumbersome if we have multiple variables with missing data and observations with more than one variable missing.
    - e.g. How do we predict x if we are missing other variables?
  - ► These approaches are *deterministic*, failing to take into account the uncertainity in the imputations.

### Multiple imputation

- ▶ Multiple imputation (MI) methods address both issues:
  - ▶ MI models use observed data to predict missing values.
    - Multiple missing variable can be imputed simultaneously.
    - Prediction uncertainity can be incorporated into the estimates.

### Multiple imputation: Algorithms

- Multiple imputation algorithms work by using existing data to predict missing values across the entire dataset
- Generally, these algorithms use iterative procedures, predicting a subset of the missing values at a time
- These algorithms converge when the distributions of the predicted datasets look like the distributions in the original data
- Often these algorithms use Bayesian techniques such as MCMC sampling<sup>1</sup>

See the mi command in Stata for example.

### Multiple imputation: MI in R

- ► There are a number of different MI packages available in R
- Two commonly used packages are
  - ► MICE (Multivariate Imputation via Chained Equations)
  - Amelia
    - Particularly useful for panel data

### Multiple imputation: Pooling

- ► MI algorithms can produce *M* imputed datasets.
  - In each case, we can compute an estimate,  $\hat{\beta}_{1m}$  and a standard error  $\hat{SE}_m$ .
- ► The overall estimate is an average over the *M* datasets, known as a **pooled** estimate:<sup>2</sup>

$$\hat{\beta} = \frac{1}{M} \sum_{m=1}^{M} \hat{\beta}_{1m}$$

<sup>&</sup>lt;sup>2</sup>See GHV p. 326 for the formula for the standard error of the pooled estimate.

### Multiple imputation with MICE

### Multiple imputation with MICE

```
# Impute using m=1
simple.imp <- complete(M,1)
mi.s <- lm(y ~ x + w, data = simple.imp)

# Pool over all M
fits <- with(M, lm(y ~ x + w))
mi.M <- pool(fits)</pre>
```

### Multiple imputation with MICE

	Complete	Predicted	MI m	MI M
(Intercept)	-0.016	-0.008	-0.029	-0.029
	(0.032)	(0.040)	(0.033)	(0.044)
X	0.994	0.998	1.007	1.021
	(0.030)	(0.052)	(0.031)	(0.038)
W	-2.007	-1.968	-2.002	-2.030
	(0.036)	(0.047)	(0.037)	(0.050)
Num.Obs.	1000	1000	1000	1000
Num.Imp.				10
R2	0.765	0.635	0.756	0.758
R2 Adj.	0.764	0.634	0.755	0.758

## Model checking

#### **Diagnostics**

- We have already covered several different diagnostics for model checking
  - Residuals and standard error of the residuals
  - Predicted values
  - $ightharpoonup R^2$  and adjusted  $R^2$
  - Standard errors on coefficients and p-values
  - F-statistic

## Model checking

#### **Outliers**

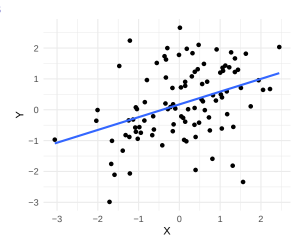
- Outliers are extreme data points that deviate from the distribution of other values
  - Scatterplots of raw data and residual can be helpful for identifying these
- An outlier has **leverage** if the addition of the observation results in a change in the slope of the regression line
- ► Such cases are considered *influential* if they result in substantial changes to the regression results
  - e.g. Differences in statistical sigificance, sign, magnitude

## Model checking

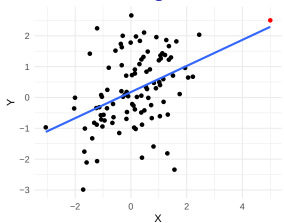
#### **Outliers**

```
N <- 100
X <- rnorm(N)
Y <- 0.5*X + rnorm(N)</pre>
```

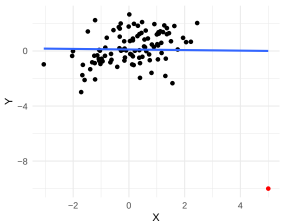
#### **Outliers**



### Outliers: Outlier with low leverage



### Outliers: Outlier with high leverage



#### Underfitting

- ► A model is **underfit** if it does not sufficiently explain the variance in the outcome.
- ► Consider the following population model:

$$y = \beta_0 + \beta_1 x + \beta_2 x^2 + u$$

► The following model underfits because it does not account for the quadratic relationship:

$$\hat{y} = \hat{\beta_0} + \hat{\beta_1} x + u$$

▶ In short, we fail to observe the signal in the data (Molina and Garip 2019)

### **Overfitting**

- A model is **overfit** if it also explains *noise* in addition to the signal in the data (Molina and Garip 2019).
- Using the previous population, consider we estimate the following model:

$$\hat{y} = \hat{\beta}_0 + \hat{\beta}_1 x + \hat{\beta}_2 x^2 + \hat{\beta}_3 z + \hat{\beta}_4 z^2 + u$$

► The additional parameters  $\hat{\beta}_3$  and  $\hat{\beta}_4$  do not correspond to any of the population parameters, but may still explain some of the random variance in the outcome.

### Underfitting, overfitting, and generalization error

- Models suffering from under and overfitting will fail to generalize
  - Underfit models do not sufficiently explain variation in the outcome in the sample or population
  - Overfit models explain patterns in the sample that do not generalize to the population
- ► **Generalization error** refers to the expected prediction error when a model is applied to new data.

#### **Cross-validation**

- Cross-validation is an approach used in machine-learning to assess the extent to which a predictive model can generalize to unseen data.
- ▶ The technique allows us to measure generalization error:
  - 1. Estimate an model using a sample X.
  - Use the fitted model to predict the outcome for a new dataset X'.
  - **3.** Compare the predictive accuracy (e.g. mean squared error) across the two datasets:
    - ▶ Model generalizes well if  $MSE(\hat{y} = f(X)) \approx MSE(\hat{y} = f(X'))$  and both are low
    - ▶ Model overfits if  $MSE(\hat{y} = f(X)) << MSE(\hat{y} = f(X'))$
    - ▶ Model underfit if both MSE scores are high

#### **Cross-validation**

- ▶ Different kinds of cross-validation procedures are often used to evaluate generalization error:
  - **k-fold cross-validation**: data are split into k subsets. Models are estimated using k-1 subsets and predictions made for held-out set. Prediction error is averaged over k held-out sets.
  - ► **Leave-one-out cross-validation**: same procedure where each subset is a single datapoint. Requires estimation of *N* models.
  - **Temporal cross-validation**: Useful with panel data. Estimate a model with data from time t the assess predictions for data recorded at t+1.

#### Cross-validation: 100

- ► The rstanarm package includes a function loo, which computes an approximation of LOO-CV, avoiding the need to fit N models.
- Models can be compared using the expected log pointwise predictive density (ELPD)}, a quantity that captures the predictive accuracy of the model (McElreath 7.2-4, GHV 11.8).
- ► The function can also be used to compute a k-fold CV score and an information theoretic measure WAIC.<sup>3</sup>

See the documentation and vignette for further details on implementation.

#### Cross-validation: LOO-CV

#### **Cross-validation: LOO-CV**

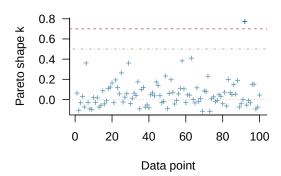
The loo function provides an approximation of the LOO-CV for an estimated model

```
print(loo(m))
##
## Computed from 1000 by 100 log-likelihood matrix
##
##
          Estimate SE
## elpd_loo -145.8 7.0
## p_loo
          3.5 0.7
## looic 291.5 14.0
## Monte Carlo SE of elpd_loo is 0.1.
##
## Pareto k diagnostic values:
##
                         Count Pct.
                                     Min. n_eff
## (-Inf, 0.5] (good) 98 98.0% 612
   (0.5, 0.7]
               (ok)
                          2
##
                               2.0%
                                     460
```

#### Cross-validation: LOO-CV

Individual points are scored using Pareto-Smoothed Importance Sampling (PSIS). High pareto k values (k > .7) indicate observations with high leverage.

#### **PSIS** diagnostic plot



#### Cross-validation: LOO-CV

▶ loo\_compare can be used to compare different models. The results rank the models from best to worst.

```
loo_compare(loo(m), loo(m.u), loo(m.o))

## elpd_diff se_diff

## m 0.0 0.0

## m.o -2.3 0.9

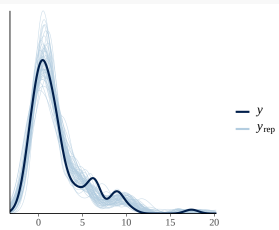
## m.u -118.4 14.9
```

#### Posterior predictive checks

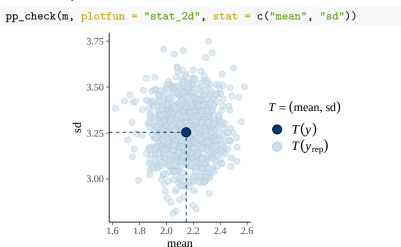
- We can also evaluate Bayesian models by examining the posterior predictive distribution
- Recall that Bayesian models are generative; we can use the posterior distribution to make predictions, creating new, hypothetical datasets
- These predictions can be used to evaluate how well the models fit the data, including key statistics

#### Posterior predictive checks

pp\_check(m)

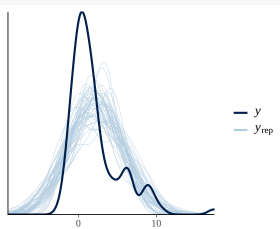


#### Posterior predictive checks

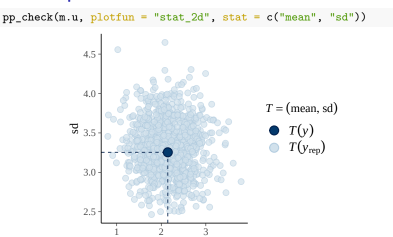


### Posterior predictive checks

pp\_check(m.u)



#### Posterior predictive checks



mean

#### **Prediction and explanation**

- Predictive models are designed to predict  $\hat{y}$ 's with high accuracy.
- In sociology, we typically estimate explanatory models where the primary goal is to estimate one or more  $\hat{\beta}$ 's and prediction is typically used to explore relationships between variables.
- Nonetheless, generalization error can be a useful heuristic for comparing and selecting different models, particularly when we do not have strong theory to guide model specification.<sup>4</sup>

<sup>&</sup>lt;sup>4</sup>See Watts, Duncan J. 2014. "Common Sense and Sociological Explanations." American Journal of Sociology 120 (2): 313–51. https://doi.org/10.1086/678271 for further elaboration of this idea and Mullainathan, Sendhil, and Jann Spiess. 2017. "Machine Learning: An Applied Econometric Approach." Journal of Economic Perspectives 31 (2): 87–106. https://doi.org/10.1257/jep.31.2.87 for some discussion of the limitations of predictive modeling.

#### Multicollinearity

- Recall that multicollinearity occurs when predictors are highly correlated and results in increased variance
- High pairwise correlations might indicate potential collinearity, but the issue can only be diagnosed after controlling for all relevant predictors

#### Multicollinearity: VIF

- ► The Variance Inflation Factor (VIF) can be used to diagnose highly collinear predictors
- Consider the regression model  $y = \beta_0 + \beta_i x_i + \beta_j x_j + ... + \beta_k x_k + u$
- ► A score is calculated for each *independent variable* using the following approach:
  - For  $x_i$  in  $x_{i=1},...,x_k$ , regress  $x_i = \alpha_0 + \alpha_1 x_j + ... + \alpha_2 x_k + u$
  - ▶ Use  $R^2$  from the model to calculate  $VIF(\hat{\beta}_i) = \frac{1}{1-R_i^2}$
- ▶ VIF scores greater than  $\approx 5-10$  indicate that a predict is highly collinear with one or more of the other predictors

#### Multicollinearity: VIF

#### **Defining robustness**

- ▶ **Model robustness** refers to how *robust* the results of a given model are to alternative specifications.
  - The concern is that a result (such as p < 0.05) is sensitive to a particular specification of a model
- Often we try to mitigate such concerns by estimating several different specifications of a model

#### How robust are our results?

- Recent work calls for greater attention to specification issues as a way to address robustness concerns (Young and Holsteen 2017, Muñoz and Young 2018)
  - Critique: Reporting a handful of ad hoc specifications is insufficient to ensure robustness
  - Solution: Estimate models with every possible combination of independent variables and assess the distribution of coefficients
  - The goal is to explore the entire model space and to construct a distribution of estimates

#### How robust are our results?

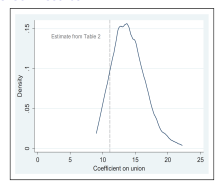
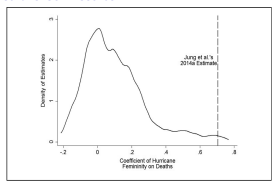


Figure 1. Modeling distribution of union wage premium.

Note: Kernel density graph of estimates from 1,024 models. Vertical line indicates the preferred estimate of an II percent union wage premium as reported in Table 2.

Young and Holsteen 2017.

#### How robust are our results?



**Figure 3.** Model robustness results on Jung et al. (2014a) data. *Note:* Kernel density graph of estimates from 1,152 models. See Table 6 for more information about the modeling distribution.

Muñoz and Young 2018.

### **Bayesian Model Averaging**

- Bayesians have proposed a similar idea known as Bayesian Model Averaging
  - Estimate several models and construct an average across the models, weighted by the model fit, i.e. higher weights to better models
- ► There has been some debate about whether this approach is preferable to the Young-Holsteen-Muñoz technique.
  - ► The latter argue that it is problematic to weight different models if we do not know which is better a priori and that weighting requires more assumptions.<sup>5</sup>

 $<sup>^{5}</sup>$ See Slez' 2017 comment on Young and Holsteen and the rejoinder by the latter and Bruce Western's 2018 comment on Muñoz and Young

### **Conclusions**

- ► Missing data
  - Carefully examine any patterns of missing data
  - Choose an appropriate strategy to address the problem
- Model checking
  - Use diagnostic checks to identify and address potential issues with models
- Model robustness
  - Estimate multiple specifications to ensure results are robust

### Next week

- ► Spring break!
- ► After spring break
  - ► Generalized linear models