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Multiple imputation of univariate missing data: the mice package School of Mathematics, University of Edinburgh

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Here we will learn how to use the mice package in practice. For now, we will only deal with univariate missingness, we will later expand the scope to the case of several variables with missing values. Before proceeding, I leave the reference to the manual of the package

https://cran.r-project.org/web/packages/mice/index.html

I will start by simulating some data and then imposing MAR missingness.

```
set.seed(1)
n <- 100
x1 <- runif(n, 0, 5)
x2 <- runif(n, 0, 10)
beta0 <- 5
beta1 <- 3
beta2 <- 1
y <- rnorm(n, beta0 + beta1*x1 + beta2*x2, 1)

x2 <- ifelse(x1 > 4.2, NA, x2)
#checking the percentage of missing values
sum(is.na(x2))/n
```

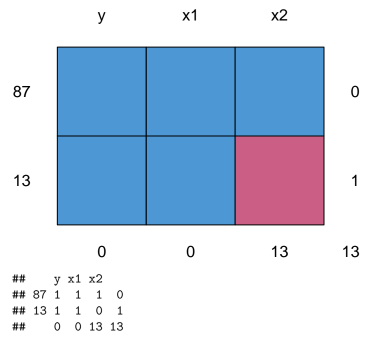
```
## [1] 0.13
```

```
#constructing a dataframe with the 3 variables
simdata <- data.frame("y" = y, "x1" = x1, "x2" = x2)</pre>
```

The package mice has the function cc that returns the complete cases. This function is useful when working with real data as it easily allows some exploratory analyses based on the complete cases.

```
require(mice)
cc(simdata)
nrow(cc(simdata))
```

As we have seen back in week 2, mice also has a function that allows visualising the missing data patterns. md.pattern(simdata)



Another function available in mice is md.pairs, which calculates the number of observations per patterns for all possible pairs of variables. For a pair of variables, there are four possible missing data patterns: both variables are observed (pattern rr), the first variable is observed and the second variable is missing (pattern rm), the first variable is missing and the second variable is observed (pattern mr), and finally the pattern where both variables are missing (pattern mm).

md.pairs(simdata)

```
## $rr
##
         x1 x2
       у
## y 100 100 87
## x1 100 100 87
## x2 87
         87 87
## $rm
     y x1 x2
     0
        0 13
## y
## x1 0
        0 13
## x2 0 0 0
##
## $mr
##
      y x1 x2
## y
      0
        0
           0
## x1 0
         0
            0
## x2 13 13 0
##
## $mm
##
     y x1 x2
## y 0
        0 0
## x1 0
       0 0
## x2 0
       0 13
```

Let us now use the package mice to impute the values in x2. We start with the function mice() to perform step 1, i.e., to impute the missing values. We already know that the default in mice for continuous variables,

as it is the case of x_2 , is predictive mean matching with d=5 donors and $Type\ 1$ matching (between the cases with missing values and those with observed values). Also, by default in mice we have M=5. To know more, type help(mice).

```
imps <- mice(simdata, printFlag = FALSE, seed = 1)</pre>
imps
## Class: mids
## Number of multiple imputations: 5
## Imputation methods:
##
       V
             "" "pmm"
##
## PredictorMatrix:
##
      y x1 x2
## y
      0
         1
## x1 1
         0
            1
## x2 1
```

A few comments apply. We set printFlag = FALSE which results in silent computation of the missing values and we also use seed=1 so that our results are reproducible (any other value would obviously work, but fixing the seed outside the function mice() will not work). A summary of the imputation results can be obtained by calling the imps object. For instance, we see that our saved object imps is of class mids which stands for multiply imputed datasets, which is a special type of object that the mice package has set up for storing multiple imputed datasets. We also obtain information about the imputation method used to impute the variables with missing values. In this case only x2 has missing values and because we have not changed the defaults, unsurprisingly, we have that predictive mean matching was used. Lastly, we have the predictorMatrix which, for instance, tell us that y and x1 were used to impute x2. It also tells us that in case y had missing values, x1 and x2 would be used to impute it and similarly for x1 we would use y and x2. We can also extract this information from imps\$predictorMatrix. The default approach in mice is to impute one variable based on all other variables.

Now let us look at the imputed values. We can extract them from our imps object.

imps\$imp\$x2

```
##
             1
                      2
                                3
                                         4
                                                  5
## 4
     9.240745 7.410786 9.2861520 9.240745 9.286152
     5.476466 5.980924 4.3147369 5.260277 2.126995
     1.103606 3.287773 3.3548749 2.075451 2.075451
## 18 1.891936 1.891936 0.3554058 2.388687 1.482116
## 21 7.410786 7.410786 9.7617069 9.240745 9.286152
## 29 3.354875 3.354875 1.7344233 2.702601 2.832325
## 52 4.781180 7.293096 3.5672691 5.748722 6.547239
## 61 3.179637 3.179637 5.0044097 3.804939 4.531314
## 70 2.126995 4.525708 3.8049389 3.179637 2.126995
## 76 8.864509 8.770575 8.4061455 7.410786 7.410786
## 77 7.828513 6.304141 6.1464497 8.405070 8.405070
## 80 9.286152 9.286152 8.7705754 9.850952 9.850952
## 94 9.240745 8.770575 9.2861520 7.410786 9.240745
```

The row numbers indicate the record number in the original dataset. We can extract, for instance, the imputed values for the first imputed dataset by simply doing the following:

```
imps$imp$x2[,1]
```

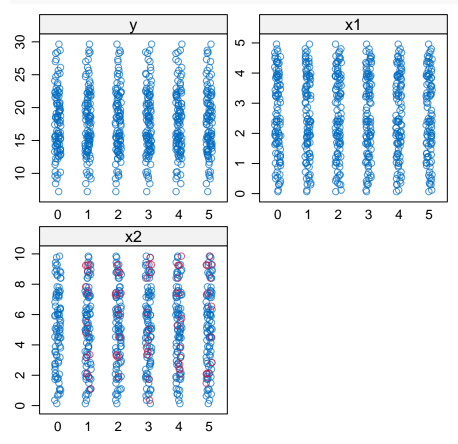
```
## [1] 9.240745 5.476466 1.103606 1.891936 7.410786 3.354875 4.781180 3.179637
## [9] 2.126995 8.864509 7.828513 9.286152 9.240745
```

The (completed) imputed datasets can be extracted by using the complete function. As a way of illustrating the usage of this function, I am extracting the first and second completed datasets.

```
com1 <- complete(imps, 1)
com2 <- complete(imps, 2)
com1
com2</pre>
```

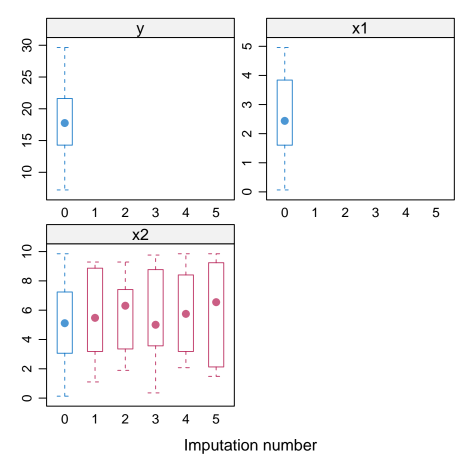
It is also important to visualise the imputation results and the package mice provides several plotting tools. This allows us to check whether imputations are plausible. As van Buuren and Groothuis-Oudshoorn say in their paper describing the mice package (p. 11): "Imputations should be values that could have been obtained had they not been missing. Imputations should be close to the data". One way to do this is through the stripplot function.

stripplot(imps)

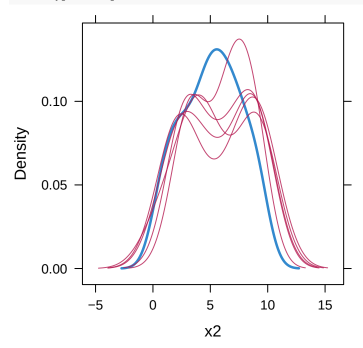


Blue circles denote observed data and red circles imputed data. The panels for y and x1 contain only blue dots because these two variables are fully observed. If there are no large differences between the imputed and observed values then we can conclude that the imputed values are plausible. Here we can see that the red circles follow the blue circles well. If there are discrepancies, interpretation is more difficult, as this may be due to a bad imputation model, due to the missing mechanism not being MAR or due to a combination of both. This plot is most useful when there are not many data points. Alternatively we can use the function bwplot, which produces a boxplot of the observed and imputed data.

bwplot(imps)



There is also the possibility of visualising the kernel density estimates of the observed and imputed data. densityplot(imps)



Aside comment: note that here the densities assign positive mass to negative x2 values and we know that this should not be the case (x2 was simulated from a uniform (0,10) distribution). By looking at the boxplots

in the previous figure, we see that all imputed values are above zero, and so the imputed values are plausible. However, due to the fact that we are using kernel estimates for the densities, which is a nonparametric density estimator, with a reduced sample size, in combination with the fact that the default kernel is the normal one and there values close to zero, leads to mass assigned to negative values.

Adjustments to the defaults used by the predictive mean matching function mice.impute.pmm can be made by simply entering the arguments to be altered into the main mice() call. They will be automatically passed down to mice.impute.pmm. For instance, the number of donors to be sampled from can be set via the donors argument. let us now change this argument to three and we will also create M=10 copies of the dataset (instead of the default M=5).

```
imps_alt <- mice(simdata, m = 10, donors = 3, printFlag = FALSE, seed = 1)</pre>
imps_alt
## Class: mids
## Number of multiple imputations:
## Imputation methods:
##
            x1
      11 11
            "" "pmm"
##
##
  PredictorMatrix:
      y x1 x2
      0 1 1
## y
## x1 1
        0
            1
```

Suppose now that we want to change our method for imputing the missing values. Specifically, suppose that we want to use the method norm.boot. There are two possible ways of doing it. The simplest way and feasible only when the number of variables to be imputed is small is to change the method argument directly in the mice() call.

x2 1

##

1

2.345993

4.452382

9.132930

```
imps_normb <- mice(simdata, method = "norm.boot", printFlag = FALSE, seed = 1)
imps_normb$imp$x2[,1]
## [1] 10.194490  4.595104  2.550149  2.953703  9.353378  2.049476  6.779982</pre>
```

An alternative way is to do a setup run of mice() without iterations (maxit=0) and to extract and modify the method from there.

8.257026 8.847016 10.444725

```
imps0 <- mice(simdata, maxit = 0)</pre>
meth <- imps0$method
meth
##
       У
             x1
             "" "pmm"
##
meth["x2"] <- "norm.boot"</pre>
imps_norm2 <- mice(simdata, method = meth, printFlag = FALSE,</pre>
                     seed = 1)
imps_norm2
## Class: mids
## Number of multiple imputations:
## Imputation methods:
##
                                        x2
              У
             11 11
##
                           "" "norm.boot"
## PredictorMatrix:
##
      y x1 x2
```

```
## y 0 1 1
## x1 1 0 1
## x2 1 1 0
```

imps_norm2\$imp\$x2[,1]

```
## [1] 10.194490 4.595104 2.550149 2.953703 9.353378 2.049476 6.779982 ## [8] 2.345993 4.452382 9.132930 8.257026 8.847016 10.444725
```

x2

"" "norm.boot"

The setup run is also useful to customize our imputation model. Variables in the columns of the predictorMatrix can be switched on or off by using a 1 or a 0 to include or exclude them from the imputation model, respectively. In this way the imputation models for each variable with missing data can be customized (remember that the default is to use all variables in the dataset to impute the variable(s) with missing data). In the hypothetical case that we only want to impute x2 using y, and not both y and x1 (note that this is only to exemplify how to customize the imputation model, I am not saying this is necessarily the way to go in this case).

PredictorMatrix: ## y x1 x2 ## y 0 1 1

x1 1 0 1 ## x2 1 0 0

##

We will now proceed to step 2, and we will use the function with(). Suppose that our substantive model, i.e., our model of interest, is the model we have used to generate the data, that is

```
y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \varepsilon, \varepsilon \sim N(0, 1).
```

Just for the sake of illustration, I will be using the completed datasets stored in the object imps, using mice's defaults.

```
fits <- with(imps, lm(y ~ x1 + x2))
class(fits)</pre>
```

```
## [1] "mira" "matrix"
```

The object fits contains the results of fitting M=5 complete data linear models based on the imputed datasets. The class of fits is mira, which stands for multiply imputed repeated analysis. We can extract the results and corresponding summary of the, say, first and second imputed datasets by doing

fits\$analyses[[1]]

```
##
## Call:
## lm(formula = y ~ x1 + x2)
##
## Coefficients:
## (Intercept) x1 x2
```

```
##
         4.938
                      2.968
                                   1.018
summary(fits$analyses[[1]])
##
## Call:
## lm(formula = y \sim x1 + x2)
##
## Residuals:
##
      Min
                1Q Median
                                3Q
                                       Max
## -1.8393 -0.6519 -0.1273 0.6897
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.93789
                           0.29489
                                     16.75
                                             <2e-16 ***
## x1
                                     38.90
                2.96771
                           0.07628
                                             <2e-16 ***
## x2
                1.01751
                           0.03838
                                     26.51
                                             <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.015 on 97 degrees of freedom
## Multiple R-squared: 0.9591, Adjusted R-squared: 0.9582
## F-statistic: 1136 on 2 and 97 DF, p-value: < 2.2e-16
fits$analyses[[2]]
##
## Call:
## lm(formula = y \sim x1 + x2)
##
## Coefficients:
## (Intercept)
                         x1
                                      x2
         4.888
                      2.925
                                   1.041
summary(fits$analyses[[2]])
##
## Call:
## lm(formula = y \sim x1 + x2)
##
## Residuals:
##
       Min
                1Q Median
                                3Q
                                       Max
## -1.8787 -0.7282 -0.1435 0.5983 2.8223
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
## (Intercept)
               4.88841
                           0.31110
                                     15.71
                                             <2e-16 ***
## x1
                2.92544
                           0.07997
                                     36.58
                                             <2e-16 ***
## x2
                1.04074
                           0.04140
                                     25.14
                                             <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.063 on 97 degrees of freedom
## Multiple R-squared: 0.9551, Adjusted R-squared: 0.9541
## F-statistic: 1031 on 2 and 97 DF, p-value: < 2.2e-16
```

```
ests <- pool(fits)
#ests
summary(ests, conf.int = TRUE)</pre>
```

The object ests is of class mipo, meaning multiply imputed pooled outcomes. Its printed output resembles the output of an lm object, but note that its content is different: pool gathers the data in mipo in a mira way that makes summarising the statistics using summary easier. One cannot therefore use residuals or predict to obtain residuals or predictions from the final estimated model.

The column estimate correspond to the pooled regression coefficients and their corresponding standard error is available in std.error. By further inspecting the output we have columns corresponding to ubar, which is the within-imputation variance \bar{U} , b corresponds to the between-imputation variance, rvi, which stands for relative increase in variance due to the missing values and as we have learned last week, its expression is given by $\frac{B+\frac{B}{M}}{U}$, the column corresponding to lambda, which is the proportion of variance in the parameter of interest due to the missing values and which is given by $\frac{B+\frac{B}{M}}{VT}$. Finally, fmi contains the fraction of missing information as defined in Rubin (1987), and it depends on rvi but we will not study it further.

We can also only select the columns we are interested in from the summary, as illustrated below. Further note that the argument conf.int = TRUE computes a 95% confidence interval for the (pooled) coefficient estimates.

```
summary(ests, conf.int = TRUE)[, c(2, 3, 7, 8)]

## estimate std.error 2.5 % 97.5 %

## 1 4.948381 0.30048570 4.3514390 5.545323

## 2 2.947302 0.07947479 2.7891265 3.105477

## 3 1.022477 0.04129880 0.9399896 1.104965
```

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

```
pool.r.squared(fits, adjusted = TRUE)
```

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

To conclude, let us check the effect of the choice of M on the results which, of course, in practice, depends on the particular analysis we are doing.

estimate std.error p.value 2.5 % 97.5 %

```
## 1 4.975028 0.30434507
                              0 4.3696852 5.580371
## 2 2.926547 0.08158902
                              0 2.7635916 3.089502
## 3 1.023931 0.04061770
                              0 0.9430113 1.104850
summary(ests\_seed3, conf.int = TRUE)[, c(2, 3, 6, 7, 8)]
    estimate std.error
                             p.value
                                         2.5 % 97.5 %
## 1 4.951749 0.32017344 0.000000e+00 4.3076009 5.595897
## 2 2.944345 0.09815355 1.110223e-15 2.7365704 3.152119
## 3 1.023076 0.03941279 0.000000e+00 0.9446045 1.101547
#using the M=20 and changing the seed
ests seed1 20 <- pool(with(mice(simdata, printFlag = FALSE, seed = 1, m = 20), lm(y ~ x1 + x2)))
ests_seed2_20 <- pool(with(mice(simdata, printFlag = FALSE, seed = 11, m = 20), lm(y ~ x1 + x2)))
ests_seed3_20 <- pool(with(mice(simdata, printFlag = FALSE, seed = 111, m = 20), lm(y ~ x1 + x2)))
summary(ests\_seed1\_20, conf.int = TRUE)[, c(2, 3, 6, 7, 8)]
    estimate std.error p.value
                                    2.5 % 97.5 %
## 1 4.926221 0.30482933
                           0 4.3202325 5.532209
## 2 2.959692 0.08561980
                              0 2.7885989 3.130786
## 3 1.022750 0.04034451
                              0 0.9424956 1.103004
summary(ests\_seed2\_20, conf.int = TRUE)[, c(2, 3, 6, 7, 8)]
    estimate std.error p.value
                                    2.5 % 97.5 %
## 1 4.958667 0.30200150
                            0 4.3586337 5.558700
## 2 2.955109 0.08509408
                              0 2.7852253 3.124992
## 3 1.017642 0.03943637
                              0 0.9392827 1.096000
summary(ests_seed3_20, conf.int = TRUE)[, c(2, 3, 6, 7, 8)]
    estimate std.error p.value
                                    2.5 % 97.5 %
## 1 4.938306 0.30942374
                              0 4.3228618 5.553751
## 2 2.947024 0.08790346
                              0 2.7710699 3.122978
## 3 1.024396 0.04017278
                              0 0.9445281 1.104264
#using the M=50 and changing the seed
ests_seed1_50 <- pool(with(mice(simdata, printFlag = FALSE, seed = 1, m = 50), lm(y ~ x1 + x2)))
ests_seed2_50 <- pool(with(mice(simdata, printFlag = FALSE, seed = 11, m = 50), lm(y ~ x1 + x2)))
ests_seed3_50 <- pool(with(mice(simdata, printFlag = FALSE, seed = 111, m = 50), lm(y ~ x1 + x2)))
summary(ests_seed1_50, conf.int = TRUE)[, c(2, 3, 6, 7, 8)]
    estimate std.error p.value
                                    2.5 % 97.5 %
## 1 4.949031 0.30095407
                          0 4.3511043 5.546957
## 2 2.949120 0.08577777
                              0 2.7780191 3.120221
## 3 1.021698 0.04022434
                              0 0.9417208 1.101674
summary(ests\_seed2\_50, conf.int = TRUE)[, c(2, 3, 6, 7, 8)]
    estimate std.error p.value
                                    2.5 % 97.5 %
## 1 4.951577 0.30207745 0 4.3513394 5.551814
## 2 2.951857 0.08441553
                              0 2.7836038 3.120110
## 3 1.020227 0.03971142
                              0 0.9413022 1.099152
summary(ests_seed3_50, conf.int = TRUE)[, c(2, 3, 6, 7, 8)]
    estimate std.error p.value
                                    2.5 % 97.5 %
##
```

```
## 1 4.957735 0.30283071
                               0 4.3559624 5.559507
## 2 2.947614 0.08337180
                               0 2.7815445 3.113683
## 3 1.020432 0.03978913
                               0 0.9413519 1.099513
#using the M=100 and changing the seed
ests_seed1_100 <- pool(with(mice(simdata, printFlag = FALSE, seed = 1, m = 100), lm(y ~ x1 + x2)))
ests_seed2_100 <- pool(with(mice(simdata, printFlag = FALSE, seed = 11, m = 100), lm(y ~ x1 + x2)))
ests_seed3_100 <- pool(with(mice(simdata, printFlag = FALSE, seed = 111, m = 100), lm(y ~ x1 + x2)))
summary(ests_seed1_100, conf.int = TRUE)[, c(2, 3, 6, 7, 8)]
##
     estimate std.error p.value
                                     2.5 %
                                             97.5 %
## 1 4.943259 0.30348831
                               0 4.3401883 5.546330
## 2 2.944795 0.08379677
                               0 2.7779059 3.111683
## 3 1.024108 0.03996190
                               0 0.9446866 1.103530
summary(ests\_seed2\_100, conf.int = TRUE)[, c(2, 3, 6, 7, 8)]
##
     estimate std.error p.value
                                     2.5 %
                                             97.5 %
## 1 4.950286 0.30149773
                               0 4.3512125 5.549359
## 2 2.945878 0.08419970
                               0 2.7781334 3.113623
## 3 1.022392 0.03931929
                               0 0.9442725 1.100511
summary(ests\_seed3\_100, conf.int = TRUE)[, c(2, 3, 6, 7, 8)]
     estimate std.error p.value
                                     2.5 %
                                             97.5 %
## 1 4.957679 0.30407645
                               0 4.3533653 5.561993
## 2 2.941578 0.08564185
                               0 2.7708611 3.112294
## 3 1.022356 0.03970532
                               0 0.9434533 1.101258
```

The (pooled) estimates, standard errors, and the bounds of the intervals get more stable as M increases and we can be more confident in any one specific run. Note that whatever value of M we choose, there will always be some variation in results between repeat runs. The point is that with a sufficiently large M, the results will with high probability only differ by a small amount.