Censored Likelihood Multiple Imputation in R

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Installing the clmi package

Installation of the clmi package requires the install_github function in the devtools package. If you have not downloaded the devtools package then run install.packages("devtools").

```
library(devtools)
install_github("https://github.com/bossjona/Single-Pollutant-Multiple-LODs")
```

Load the clmi package and an example dataset

For convenience we have included a example dataset called toy.data, which can be loaded by running data("toy-example"). Let's look at the first 10 entries of the example dataset.

```
library(clmi)
data("toy-example")
head(toy.data, n = 10)
```

##		id	case_cntrl	poll	smoking	gender	batch1
##	1	13707	1	3.588607	0	1	0
##	2	18641	1	NA	0	0	0
##	3	27407	1	2.619124	1	0	0
##	4	45462	1	7.203193	0	1	1
##	5	50357	1	7.336160	1	1	1
##	6	59168	1	NA	0	0	0
##	7	61477	1	5.136974	0	1	0
##	8	76585	1	11.794483	1	1	0
##	9	80681	1	1.280289	0	0	1
##	10	84391	1	5.480510	1	1	0

The id column gives the study ID and is unimportant for the purposes of this example. The case_cntrl takes values 0 or 1, where 1 indicates that the subject has the disease of interest and 0 indicates that the subject is a healthy control. The column poll is the environmental exposure of interest, where NA indicates that the concentration is below the limit of detection (LOD). The variables smoking and gender are covariates that we are going to include in the imputation model. Lastly, batch1 takes two values; 1 if the subject's biosample was assayed in batch 1 and 0 if the subject's biosample was assayed in batch 2. Moreover, the LOD for batch 1 is 0.8 and the LOD for batch 2 is 0.65.

Implementing Censored Likelihood Multiple Imputation

The function that performs censored likelihood multiple imputation is the clmi function. For details see help(clmi).

```
lod.info = data.frame(batch.info = c("1","0"), lod = c(0.8, 0.65)),
n.imps = 20, seed = 12345, t.function = function(x) log(x))
```

Some important things to notice:

- The contaminant, batch, outcome, and contam.covars arguments must correspond to the variable names in toy.data.
- The lod.info argument is where all of the LOD information goes. The first column must be named batch.info and contains all levels of the batch1 variable as character strings. The second column must be named lod and contains the LODs corresponding to each batch. Remember that all of the values below the LOD must be entered as NA so that the function recognizes them as missing.
- The t.function argument specifies whether you want to return an imputed concentrations on the untransformed scale (default) or on a transformed scale. It is often the case that contaminant concentrations are log-transformed to handle right skewed exposure data while still maintaining interpretable effect estimates. To impute on the log-scale specify, t.function = function(x) log(x). As another example, if you want to impute on the square root scale then specify, t.function = function(x) sqrt(x).

The imputed datasets can be extracted as a list using **\$imputed.dfs**:

```
extract.imputed.dfs <- clmi.out$imputed.dfs
```

Fit and pool outcomes models

The pool.clmi function takes the output generated by the clmi function, fits outcome models on each of the imputed datasets, and pools inference across outcome models. For details see help(pool.clmi).

A few things that deserve further comment:

- There are currently two options for the regression.type argument. If you have binary outcome data (as in the current example) use regression.type = "logistic" so that the outcome models fit on the imputed datasets are logistic regression models. If you have continuous outcome data use regression.type = "linear" so that the outcome models fit on the imputed datasets are linear regression models.
- The outcome.covars argument is a vector of strings containing the variable names corresponding to the precision variables that you want to include as adjustment covariates in the outcome models. Note that the strings must correspond to variable names in your dataset. Variables included in the contam.covars when running the clmi function are automatically adjusted for. Therefore, outcome.covars should only contain additional adjustment covariates that were not included in the imputation model. In this case outcome.covars = NULL means that we only want an single-pollutant outcome model that adjusts for the variables in contam.covars, i.e. smoking and gender.

To display the pooled results use **\$output**:

```
results$output
```

```
## (Intercept) 0.36326915 0.07548992 91.98337 5.826613e-06
## smoking -0.08165125 0.11788361 93.48825 4.902506e-01
## gender 0.18971427 0.11566557 91.80907 1.043863e-01
## poll_transform_imputed 0.09303489 0.04861608 89.73839 5.885053e-02
## LCL.95 UCL.95
```

```
## (Intercept) 0.213339286 0.5131990

## smoking -0.315728622 0.1524261

## gender -0.040013900 0.4194424

## poll_transform_imputed -0.003553288 0.1896231
```

If you want to look at the individual regressions fit on each imputed dataset use regression.summaries

```
results$regression.summaries
```

Contact information

If you would like to report a bug in the code, ask questions, or send requests/suggestions e-mail Jonathan Boss at bossjona@umich.edu.