# Characterization of missing values in untargeted MS-based metabolomics data and evaluation of missing data handling strategies

Example code for data simulation and imputation

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# Simulation of incomplete data

#### Load required packages and functions

```
require(mvtnorm)
require(parallel)
require(corpcor)
source("gen.data.met.r")
```

### Parameter settings

```
n <- 250
naux <- 5
p <- 2+2*naux
corr <- 0.2
pcor12 <- 0.09977889 # corresponding to Pearson correlation corr = 0.2
pcoraux <- 0.3165772</pre>
```

- n is the number of observations.
- naux is the number of auxiliary metabolites for each metabolite of interest.
- p is the number of variables to be created; in this case, 2 metabolites of interest and 5 auxiliary metabolites for each of them are created.
- corr is the Pearson correlation to achieve between the 2 metabolites of interest.
- pcor12 is the partial correlation between the 2 metabolites of interest such that the set Pearson correlation (corr = 0.2) is achieved in the overall correlation matrix.
- pcoraux is the partial correlation of each metabolite of interest with its 5 auxiliary metabolites. In this case, the value was determined numerically based on our real dataset (KORA F4) such that the 5 auxiliary metabolites achieve an adjusted  $R^2$  of 50% for the metabolite of interest on average to resemble real data example.

#### Generate correlation matrix based on specified parameters

```
matrix(rep(c(0,pcoraux,rep(0,naux),rep(0,naux)),each=naux),nrow=naux))
diag(Pcor) <- 1
Cor <- pcor2cor(Pcor)</pre>
```

- Pcor is the generated partial correlation matrix.
- Cor is the Pearson correlation matrix generated from the partial correlation matrix using function pcor2cor from package corpcor.

#### Generate artificial, complete dataset

Based on the parameters defined above, the function gen.complete.data generates a list datCom, which stores the artificial data in datCom\$x and runday information in datCom\$rundays. datCom\$x is a data.frame with 250 observations and 13 variables. The first and second column correspond to the two main metabolites of interest, columns 3-12 correspond to the auxiliary metabolites, and the last column stores the phenotype (see Supporting Information S3).

- n is the number of observations.
- p is the number of variables.
- Cor is correlation matrix of the variables.
- n.rundays is the number of rundays. In this case, it is specified such that each runday comprises measurements of 34 samples, the everage number observed for the KORA F4 dataset.

#### Generate artificial, incomplete data by introducing missingness

The function gen.miss introduces missing values into the generated artificial data matrix, thereby producing the incomplete data matrix dat.

- datx is the complete data set (output from gen.complete.data).
- incoms is the vector of the proportion (percentage) of incomplete entries to be introduced into the variables; in this example, 10% of the values of the two main metabolites will be set to missing, while all remaining variables will remain complete.
- mech is the missingness mechanism to be introduced into the data (see Supporting Information S3); in this case, runday-specific probabilistic LOD is used.
- betas1 is a vector which specifies the strength of dependency of missingness from the concentration values; in this case, we specify a strong, negative dependency guided by observations from the real dataset (see parameter  $\beta_1$  in Table S8 of Supporting Information S3).

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- strengths.incom.vary is a vector which specifies the variation of missingness across rundays; in this case, we set  $(min(incom, 1-incom)/4)^2$  for moderate variation, which was guided by observations from the real data (see parameter  $\sigma_{miss}^2$  in Table S8 of Supporting Information S3).
- strengths.beta1.vary is a vector which specifies the variation of beta1 across rundays; in this case, we set it to  $(\beta_1/2)^2$  for strong variation, which was guided by observations from the real data (see parameter  $\sigma_{\beta_1}^2$  in Table S8 of Supporting Information S3).
- cors.incom specifies the correlation of missingness across rundays for the two main metabolites (see parameter  $\rho_{miss}$  in Table S8 of Supporting Information S3).
- rundays defines the number of rundays in which each sample was measured.

# Imputation and regression- and correlation-based analysis

## Load required packages and functions

```
require(mice)
require(mitools)
require(pan)
require(ppcor)
require(tmvtnorm)
source("do.imp.met.r")
```

## Perform imputation and evaluations

The function do.imp performs imputation for a given list of methods. The output can either be the imputed data or the already evaluated performance results of the different imputation methods.

- dat is the incomplete data set generated by the function gen.miss; the two columns correspond to the two main metabolites, the third column contains a continuous phenotype without missing values.
- datCom (optional) is the corresponding complete data set generated by the function gen.complete.data for comparison of the results.
- methods is a vector of imputation methods to be applied (see Supporting Information S4).
  - cca: complete case analyses
  - min: minimum imputation
  - mean: mean imputation
  - nie: Richardson & Ciampi (RC)
  - nie.runday: RC on rundays (RC-R)
  - u.tsi: truncated sampling imputation (ITS)
  - u.tsi.runday: ITS on rundays (ITS-R)

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- si.norm: multivariate single imputation by chained eqations and Bayesian regression as primary model (ICE-norm)
- si.pmm: ICE and predictive mean matching as primiary model (ICE-pmm)
- u.tsmi: multiple truncated sampling imputation (MITS)
- u.tsmi.runday: MITS on rundays (MITS-R)
- mi.norm: multiple ICE-norm (MICE-norm)
- mi.pmm: multiple ICE-pmm (MICE-pmm)
- si.pan.incl.runday: ICE with random intercept per runday (ICE-adjR)
- mi.pan.incl.runday: MICE with random intercept per runday (MICE-adjR)
- knn.variable: k-nearest neighbor imputation on variables (KNN-var)
- knn.sample.euc: k-nearest neighbor imputation on samples based on Euclidian distance (KNN-obs)
- knn.sample.euc.sel: KNN-obs with variable pre-selection
- analysis (optional) is a vector of pervormance evaluation methods to be performed.
  - mse1/2/3/4: mean squared error
  - corxy: correlation between the two main metabolites
  - pcorxy: partial correlation between the two main metabolites
  - Imx: linear regression with first metabolite as predictor and second metabolite as response
  - Imy: linear regression with first metabolite as response and second metabolite as predictor
  - logx: logistic regression with phenotype as response and first metabolite as predictor
- K is a vector of neighbors to use for k-nearest neighbor imputation
- min.in.runday specifies the minimum number of observed values in each runday used as a condition to perform ITS imputation; in this example, we set it to half of the average number of observations per runday observed in the real data.
- return.da1 is a logical parameter indicating that the imputed data should be returned (instead of the evaluation results).