

Combining Multiple Imputation and Cross-Validation for Predicting Survival of ECMO Treatment in ARDS Patients

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1 Introduction

1.1 Discussion of the Context

- Description of Acute Respiratory Syndrome
- Description of ECMO treatment

1.2 Aims of the Proposed Research

Prediction in medical data can often be difficult; imbalanced class distributions and poor predictive covariates. If the sample size is small, then prediction becomes even more difficult. Some of these issues arise from the experimental design of the study but little can be remedied post-hoc. Missing values in the data complicate analysis even further and are often handled either by dropping missing observations or filling in the missing value by the mean. Both methods can be valid if certain assumptions hold, but useful information is either lost to the analysis or the natural distribution of the data is effected.

Multiple imputation is another method for handling missing data that is not yet common in analysis of medical datasets (**Citation**). This method both allows retention of observations in the analysis as well as accounts for the uncertainty of the imputed value. The advantages come at the cost of complexity and increased computation time. Multiple datasets must be imputed and results somehow pooled. This paper investigates the use of multiple imputation in increasing prediction performance in a medical dataset.

1.3 Questions of Interest

The three main questions of interest this paper aims to answer are:

1. Can ECMO treatment survival (**ECMO_Survival**) be accurately predicted by PreECMO biomedical markers?
2. What is the future expected performance of predictions?
3. Which biomedical markers are needed for accurate prediction and which can be dropped?

1.4 Study Population & Data Description

- Description of the study and variables involved

The dataset is composed of 450 observations on patients with Acute Respiratory Distress Syndrome who underwent ECMO treatment. The response variable, **ECMO_Survival**, is a binary categorical variable for survival indication with levels “Y” and “N”. 33 covariates are included in the analysis, two of which are categorical, and 31 continuous. The binary categorical variable **Gender** has two levels for “m”, “f” and **Indication** is a seven level disease

indicator. **Age** is a continuous variable included in the analysis. The remaining variables are biomedical markers from hospital measurements.

2 Methodology

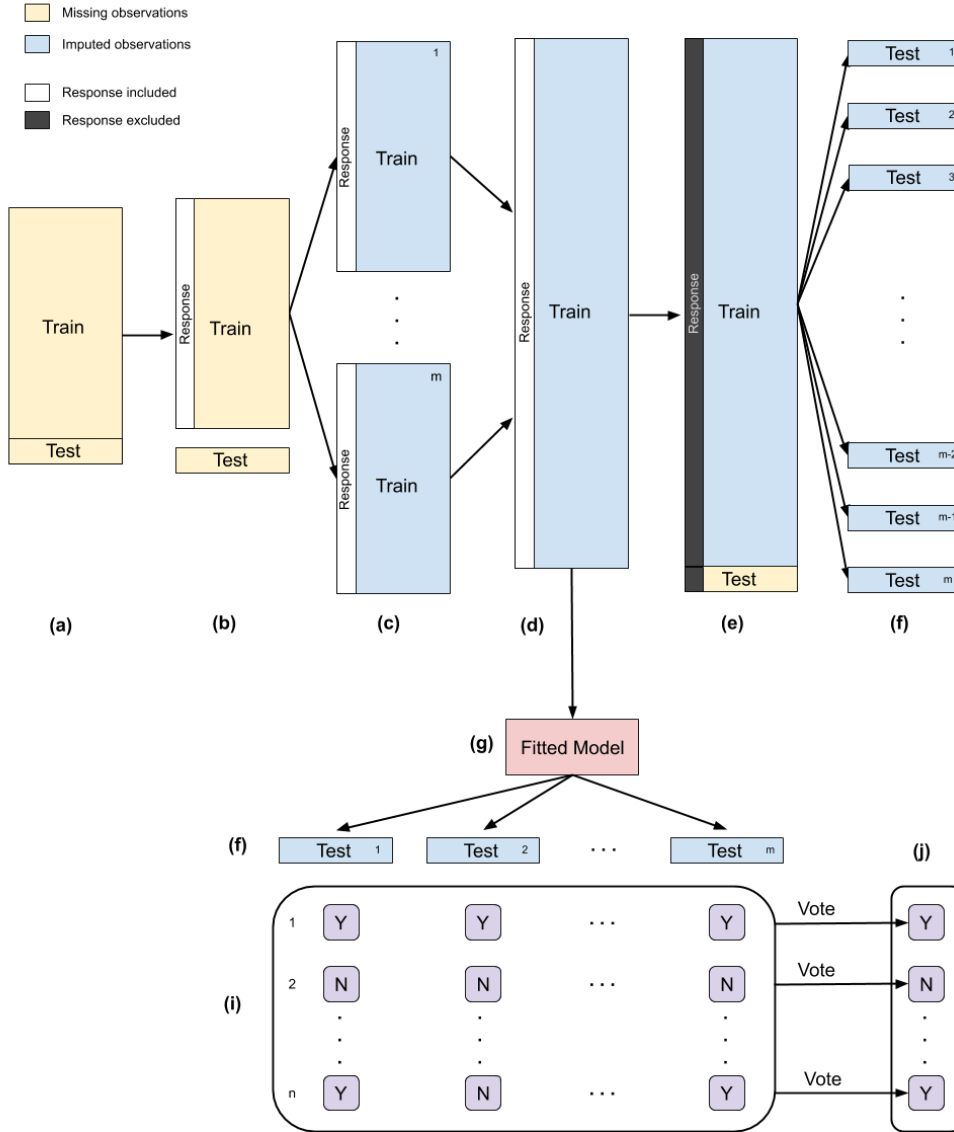


Figure 1: Outline of the algorithm used to pool predictions from multiple imputation. (a) Step 1. (b) Step 2. (c) Step 3. (d) Step 4. (e) Step 5. (f) Step 6.

2.1 Validation & Cross-validation

When building a classification model, it is important to assess its ability to produce valid predictions. If there are ample number of observations, one way to assess model performance is to randomly split the dataset into training, validation, and test sets. The training set is used to fit the model, which is then used to predict the classes for the observations in the validation set; the validation set is used to estimate prediction error and tune hyperparameters.

for model selection; the test set is used to estimate future prediction performance for the model/hyperparameters chosen. To simulate the model predicting on future, unseen data, the test set should be kept isolated. The model can overfit the data if feature manipulation and hyperparameter tuning are done before randomly splitting the data. If standardization and transformation of the covariates is done on the entire dataset, information from the training set can “leak” into the test set and the true test error will be underestimated.

If there is insufficient data to split into three parts then a suitable alternative is K -fold cross-validation. It is one of the simplest and most widely used method for estimating prediction error [Hastie et al., 2009]. The data is randomly split into K folds, where the K^{th} fold is taken as the validation set and the remaining $K - 1$ folds are used for training the model. The procedure is then repeated K times and the prediction error averaged. K -fold cross validation is most useful on sparse datasets as it allows more observations to be used in training the model. The choice of K can effect the variability of the prediction error; if $K = 1$, the model will overfit the data and prediction error will be highly variable and if $K = n$ (the number of observation in the dataset), the model is fit with no validation set for training parameters. Typical values used are $K = 5$ & 10 [Hastie et al., 2009].

a training and a test set, respectively, preserving class proportions using the `createDataPartition()` from the **caret** package.

2.2 Models

There are many classification methods, some perform well on many types of data and others perform better on certain types of data. A variety of classification methods are explored toward the aim of predicting survival of ECMO treatment, including parametric methods with many assumptions and high bias as well as non-parametric methods with higher variability. The five explored on the ARDS dataset in this paper are: Logistic Regression, Linear Discriminant Analysis, Quadratic Discriminant Analysis, K-Nearest Neighbors, and Random Forests.

2.2.1 Logistic Regression

Logistic regression is a widely used approach in machine learning and medicine for binary classification. It is a generalisation of linear regression that models the posterior probabilities of the Y classes. A logit link is used to ensure the posterior probabilities sum to one and are bounded by $[0,1]$. For two classes, the model has the form

$$\text{logit}\left(\Pr(Y|X)\right) = \log\frac{\Pr(Y = 1|X = x)}{\Pr(Y = 2|X = x)} = \mathbf{x}_i^T \boldsymbol{\beta}$$

The posterior probabilities are estimated by maximizing the log-likelihood function to find the parameter estimates, $\hat{\boldsymbol{\beta}}$, to obtain estimates of the probabilities:

$$\Pr(Y = 1|X) = \frac{\exp(\mathbf{x}_1^T \hat{\boldsymbol{\beta}})}{1 + \sum_{i=1}^2 \exp(\mathbf{x}_i^T \hat{\boldsymbol{\beta}})}$$

2.2.2 LDA and QDA

Discriminant Analysis is a widely used set of classification methods. A generalization of Fisher's Linear Discriminant [FISHER, 1936], discriminant functions are created through a combination of the explanatory variables that characterize the classes.

Let $p(X|Y)$ be the densities of distributions of the observations for each class and let π_Y denote the prior probabilities of the classes; that is, the prior probability that a randomly sampled observation belongs to the Y^{th} class based on the class proportions. The posterior probabilities may be written using Bayes Theorem as:

$$p(Y|X) = \frac{p(X|Y) \pi_Y}{p(X)} \propto p(X|Y) \pi_Y \quad (1)$$

Suppose the class distribution for class Y is Multivariate Normal with mean μ_Y and covariance matrix Σ_Y , so that:

$$p(X|Y) = \frac{1}{(2\pi_Y)^{p/2} |\Sigma_Y|^{1/2}} \exp \left[-\frac{1}{2} (X - \mu_Y)^T \Sigma_Y^{-1} (X - \mu_Y) \right] \quad (2)$$

In comparing two classes, it is sufficient to look at the log-ratio:

$$\log \frac{\Pr(Y = 1|X = x)}{\Pr(Y = 2|X = x)} = \log \frac{p(X|Y = 1)}{p(X|Y = 2)} + \log \frac{\pi_1}{\pi_2} \quad (3)$$

and using Bayes Discriminant Rule stating that *an observation should be allocated to the class with the largest posterior probability*. From Equation (1), the posterior probability may be written as

$$p(Y|X) \propto \exp(Q_Y) \quad (4)$$

where

$$Q_Y = (X - \mu_Y) \Sigma_Y^{-1} (X - \mu_Y)^T + \log |\Sigma_Y| - 2 \log \pi_Y \quad (5)$$

defines the Quadratic Discriminant Function for class Y . The Bayes Discriminant Rule is then: *allocated the observation to the class with the largest QDF*. This method of classification is called *Quadratic Discriminant Analysis* (QDA) because the decision boundaries between classes are elliptical and defined by Q_Y , an equation quadratic in X . If the covariance matrix, Σ_Y is assumed to be equal for each class then

$$L_Y = X\Sigma_Y^{-1}\mu_Y^T - \frac{1}{2}\mu_Y\Sigma_Y^{-1}\mu_Y^T - \log \pi_Y \quad (6)$$

defines the *Linear Discriminant Function*. This method has linear decision boundaries between classes defined by L_Y , an equation linear in X , and is known as *Linear Discriminant Analysis* (LDA). The Bayes Discriminant Rule is then: *allocated the observation to the class with the largest LDF*.

There is a bias-variance trade-off; both assume the covariates are normally distributed, there is no multicollinearity, and the observations are independent [Cover, 1965]. LDA additionally assumes equal class covariances. Discriminant Analysis can only utilize continuous covariates with no missing observations. The bias from simple linear or quadratic class boundaries can be acceptable because it is estimated with less variance. Despite the many assumptions and limitations, both LDA and QDA are widely used and perform well on a diverse set of classification tasks [Hastie et al., 2009], even when the classes are not normally distributed.

2.2.3 K-Nearest Neighbors

K-Nearest Neighbors (KNN) is a commonly used non-parametric classification method. To predict the class of a new observation, a distance matrix is constructed between all observations and the K nearest labelled observations to the new observation are considered. The new observation is then assigned the class label that the majority of its neighbors share. In case of only two classes, ties in class assignments are avoided by using odd values of K.

In the event of a tie, a class can be chosen at random. Various distance metrics may be used but it is common to use Euclidean distance to determine the closest training points, though it is advisable to scale variables so that one direction does not dominate the classification.

KNN is sensitive to the local structure of the data. As K increases, the variability of the classification tends to decrease at the expense of increased bias.

2.2.4 Random Forests

Random forests (Breiman, 2001) are one of the most successful general-purpose modern algorithms (Biau and Scornet, 2016). They are an ensemble learning method that can be applied to a wide range of tasks, namely classification and regression. A random forest is created by building multiple decision trees, where randomness is introduced during the construction of each tree. Predictions are made by classifying a new observation to the mode of the multiple decision tree classifications. Random forests often make accurate and robust predictions, even for very high-dimensional problems (Biau, 2012). See (**Appendix X**) for an explanation of the random forests algorithm.

- **State why random forests are good predictors**

2.3 Accuracy Metrics

These are the default metrics used to evaluate algorithms on binary and multi-class classification datasets in caret.

2.3.1 Accuracy, Sensitivity, and Specificity

Accuracy is the percentage of correctly classifies instances out of all instances. It is more useful on a binary classification than multi-class classification problems because it can be less clear exactly how the accuracy breaks down across those classes (e.g. you need to go deeper with a confusion matrix). Learn more about Accuracy [here](#).

Don't use accuracy (or error rate) to evaluate your classifier! There are two significant problems with it. Accuracy applies a naive 0.50 threshold to decide between classes, and this is usually wrong when the classes are imbalanced. Second, classification accuracy is based on a simple count of the errors, and you should know more than this. You should know which classes are being confused and where (top end of scores, bottom end, throughout?)

Table 1: Confusion matrix for two classes.

		Observed	
		N	Y
Predicted	N	a	b
	Y	c	d

For the two class confusion matrix in Table 1 accuracy metrics are defined as:

$$\begin{aligned}\text{sensitivity} &= \frac{a}{a + c} \\ \text{specificity} &= \frac{d}{b + d} \\ \text{accuracy} &= \frac{a + d}{a + b + c + d}\end{aligned}$$

where sensitivity is a measure of how accurately non-survival is predicted, specificity is a measure of how accurately survival is predicted, and accuracy is a measure of how well both survival and non-survival are predicted. While sensitivity and specificity state the accuracy each class prediction, accuracy is a poor measure for model performance in an imbalanced dataset. On the ARDS datasets, for example, if `ECMO_Survival` is predicted to be “Y” for all cases, then the accuracy is 75% but the prediction is no better than the baseline likelihood of the class percentages.

2.3.2 Cohen's Kappa

Kappa or Cohen's Kappa is like classification accuracy, except that it is normalized at the baseline of random chance on your dataset. It is a more useful measure to use on problems that have an imbalance in the classes. Let p_o be the accuracy, the relative observed agreement between observed and predicted classes and let p_e be the probability of chance agreement based on the class probabilities. Cohen's Kappa is defined as:

$$\kappa = \frac{p_o - p_e}{1 - p_e}$$

If all the observations are predicted correctly then $\kappa = 1$. If the observations are predicted no better than expected by the class probabilities, p_e then $\kappa = 0$. If all the observations are predicted incorrectly, then $\kappa = -1$. A positive κ indicates that the model predicts better than would be expected by chance whereas a negative κ indicates that the model predicts worse than would be expected by chance.

$$p_o = \frac{a + d}{a + b + c + d}$$
$$p_e = p_{o,Y} + p_{o,N}$$

$$p_{o,Y} = \frac{a + d}{a + b + c + d} \cdot \frac{a + c}{a + b + c + d}$$
$$p_{o,N} = \frac{c + d}{a + b + c + d} \cdot \frac{b + d}{a + b + c + d}$$

2.4 Missing Data

Missing data is a common problem that must be dealt with in machine learning, statistics, and medicine. Understanding the missing mechanism for the missing observations is important in the analysis. [RUBIN, 1976] defined three types of missing data mechanisms: missing completely at random (MCAR), missing at random (MAR), and missing not at random (MNAR). The data are said to be missing completely at random (MCAR) if the probability of being missing is the same for all cases. This implies the causes of the missing data are unrelated to the data itself. While MCAR is convenient because it allows many complexities that arise because data are missing to be ignored, it is typically an unrealistic assumption [van Buuren, 2012]. The data is said to be MAR if the probability of being missing is the same only within groups defined by the observed data. MAR is a more general and more realistic assumption than MCAR. If neither MCAR nor MAR applies, then the probability of being missing depends on an unknown mechanism and said to be MNAR. Most simple approaches to dealing with missing data are only valid under MCAR assumption. Modern methods to dealing with missing data begin from the MAR assumption.

2.5 Imputation Methods

2.5.1 Complete Case Analysis

Complete case analysis is a convenient method for handling missing data and is the default method in many statistical packages. If there is a missing value in an observation, it is dropped from the analysis. This is often a poor approach as complete cases analysis assumes MCAR. In sparse datasets a complete case analysis can cause an analysis to be underpowered and if MCAR does not hold, can severely bias estimates of means, regression coefficients, and correlations [van Buuren, 2012].

The ARDS dataset considered in this paper has 268/450 observations with missing data.

2.5.2 Mean Imputation

Another common method for handling missing data is mean imputation; the missing value is replaced by the mean of the covariate or the mode for categorical data. Mean imputation is a simple and attractive solution because it retains more of the data. Mean imputation distorts the distribution of the variables toward the mean. If MCAR assumption does not hold it will underestimate the variance and produce biased estimates other than the mean [van Buuren, 2012]. [van Buuren, 2012] suggests mean imputation should only be used only when there are few missing values, and should be generally avoided.

- Can further distort the distribution if the variable is not normally distributed.
- Mean imputation is implemented in this paper because although it is often a poor method of choice for imputing missing values, it is commonly done.
- Variables are transformed via Yeo-Johnson Transformation to reduce distributional distortions.

2.5.3 Multiple Imputation

The aim when imputing data is to recreate the dataset and recreate the missing data as if it were never missing. Multiple imputation is a method that accounts for the uncertainty in the imputed values. The analysis begins with the observed, incomplete dataset. The dataset is imputed multiple times to create $m > 1$ complete datasets. The imputed values are drawn from a distribution specifically modeled for each missing entry. The m datasets are analyzed using the same method that would have been used had the data been complete. The results will differ because of the variation in the input data caused by the uncertainty in the imputed values.

Multiple imputation can handle data that is both MAR and MNAR.

There is uncertainty as to the true value of the unseen data, and that uncertainty should be included in the analysis. Multiple imputation is a method created by Donald Rubin wherein multiple datasets are imputed, the analysis is conducted on each dataset, and the results are pooled using “Rubin’s Rules” [RUBIN, 1976].

- Details of the **MICE** algorithm can be found in Appendix B.

2.5.4 Fully Conditional Specification

2.5.5 Predictive Mean Matching

Predictive Mean Matching (PMM) is a semi-parametric imputation approach. It is similar to the regression method except that for each missing value, it fills in a value randomly from among the a observed donor values from an observation whose regression-predicted values are closest to the regression-predicted value for the missing value from the simulated regression model (Heitjan and Little 1991; Schenker and Taylor 1996). The PMM method ensures that imputed values are plausible; it might be more appropriate than the regression method (which assumes a joint multivariate normal distribution) if the normality assumption is violated (Horton and Lipsitz 2001, p. 246). PMM is fairly robust to transformations of the target variables [van Buuren, 2012], yielding similar results for a Yeo-Johnson transformation or no transformation.

2.6 Ensemble Multiple Imputation

Two approaches have been proposed for pooling results from several SVMs (**Belache et al. 2014**) and Cox regression (**Zavrakidis 2017**) from multiply imputed datasets. The method is to concatenate the m imputed datasets and fit a classifier, and optimize, to the resulting set; this accounts for the variability of the parameter estimates as well as the variability of the training observations in relation to the imputed values (**Belache et al. 2014**). The second procedure fits separate classifiers to each imputed data set and get the pooled (i.e. averaged) performance of the m classifiers. Results from both studies either show similar results between approaches (**Zavrakidis 2017**) or slightly better performance with the first approach (**Belache et al. 2014**). For simplicity and the sake of computational costs, this paper, only considers the first approach.

The steps in the ensemble approach for multiply imputed data in k-fold cross-validation are as follows:

1. Randomly partition the training data into k folds
2. Define the k^{th} as the test set and the remaining $k - 1$ folds as the training set
3. Impute the training set m times, with the response variable `ECMO_Survival` included, to create m imputed training sets
4. Concatenate the m imputed training sets into one extended training set
5. A model is fitted to the extended training set

6. The test set is concatenated with the extended training set
7. Impute the combined test and extended training set, with the response variable `ECMO_Survival` excluded, to create m imputed combined test and extended training sets
8. Extract the m test sets
9. Make m predictions on the m imputed test sets
10. Take the majority vote of the m predictions as the prediction for the fitted model
11. Validate the prediction against the test set by calculating Cohen's Kappa (note there are no missing values for the response variable in the data)
12. Repeat steps 2-11 k times and validate the fitted model on each training set against the test set for each fold
13. Average the k calculated Cohen's Kappas as the estimated in-sample accuracy metric

2.6.1 Number of Imputations

Flexible Imputation Book

"The classic advice is to use a low number of imputation, somewhere between 3 and 5 for moderate amounts of missing information. Several authors investigated the influence of m on various aspects of the results. The picture emerging from this work is that it is often beneficial to set m higher, somewhere in the range of 20-100 imputations.

"Rubin's Rules" provide a simple method for pooling parameters estimates from multiple imputation for linear and generalized linear models but to the author's knowledge, there has been insufficient work on estimating the required number of imputations for estimating posterior probabilities in classification problems. In his book, (**Van Buuren**) states

"Theoretically it is always better to use higher m , but this involves more computation and storage. Setting m very high (say $m=200$) may be useful for low-level estimands that are very uncertain, and for which we want to approximate the full distribution, or for parameters that are notoriously different to estimates, like variance components. On the other hand, setting m high may not be worth the extra wait if the primary interest is on the point estimates (and not on standard errors, p-values, and so on). In that case using $m=5-20$ will be enough under moderate missingness."

2.7 Voting

There has been sufficient exploration into pooling of posterior probabilities resulting from classification problems (**Citation 1**) (**Citation 2**) (**Citation 3**). But not all statistical methods produce posterior probabilities and the comparison of pooled models from multiple imputation is an area ripe for more analysis. Indeed, others have pooled predictions from various machine learning methods by taking the majority vote (**Zavrakidis**) (**Citation 2**), and comparing prediction accuracy.

2.7.1 Majority Vote

The combination can be implemented using a variety of strategies, among which majority vote is by far the simplest, yet it has been found to be just as effective as more complicated schemes [Lam and Suen, 1995].

(Alexandre et al. 2001) There has been some interest on the comparative performance of the sum and product rules (or the arithmetic and geometric means) (Kittler et al., 1996; Tax et al., 1997; Kittler et al., 1998). The arithmetic mean is one of the most frequently used combination rules since it is easy to implement and normally produces good results.

In (Kittler et al., 1998), the authors show that for combination rules based on the sum, such as the arithmetic mean, and for the case of classifiers working in different feature spaces, the arithmetic mean is less sensitive to errors than geometric mean.

In fact (Alexandre et al. 2001) show that for classification problems with two classes, that give estimates of the a posteriori probabilities that sum to one the combination rules arithmetic mean (or the sum) and the geometric mean (or the product) are equivalent.

3 Results

3.1 Exploratory Data Analysis

To get an idea of the distribution of the data, the following summary statistics were obtained for the categorical variable `ECMO_Survival` (Table 2) and for the continuous variables (Table 3).

Table 2: Numbers of survivors and nonsurvivors of ECMO treatment.

ECMO_Survival	n	Percent %
N	109	24.22
Y	341	75.78

Table 2 shows that out of the 450 individuals, only 75.78% of the individuals in the study sample survived ECMO treatment (341 survived vs 109 did not survive).

Table 3: Number of males and females.

Gender	n	Percent %
m	305	67.78
w	145	32.22

Table 3 shows that out of the 450 individuals, only 67.78% of the individuals in the study sample are male (305 male vs 145 female).

Table 4: Number of each disease type indication.

Indication	n	Percent %
1	66	14.67
2	181	40.22
3	31	6.89
4	28	6.22
5	71	15.78
6	12	2.67
7	61	13.56

Table 4 shows the distribution of each disease type indication.

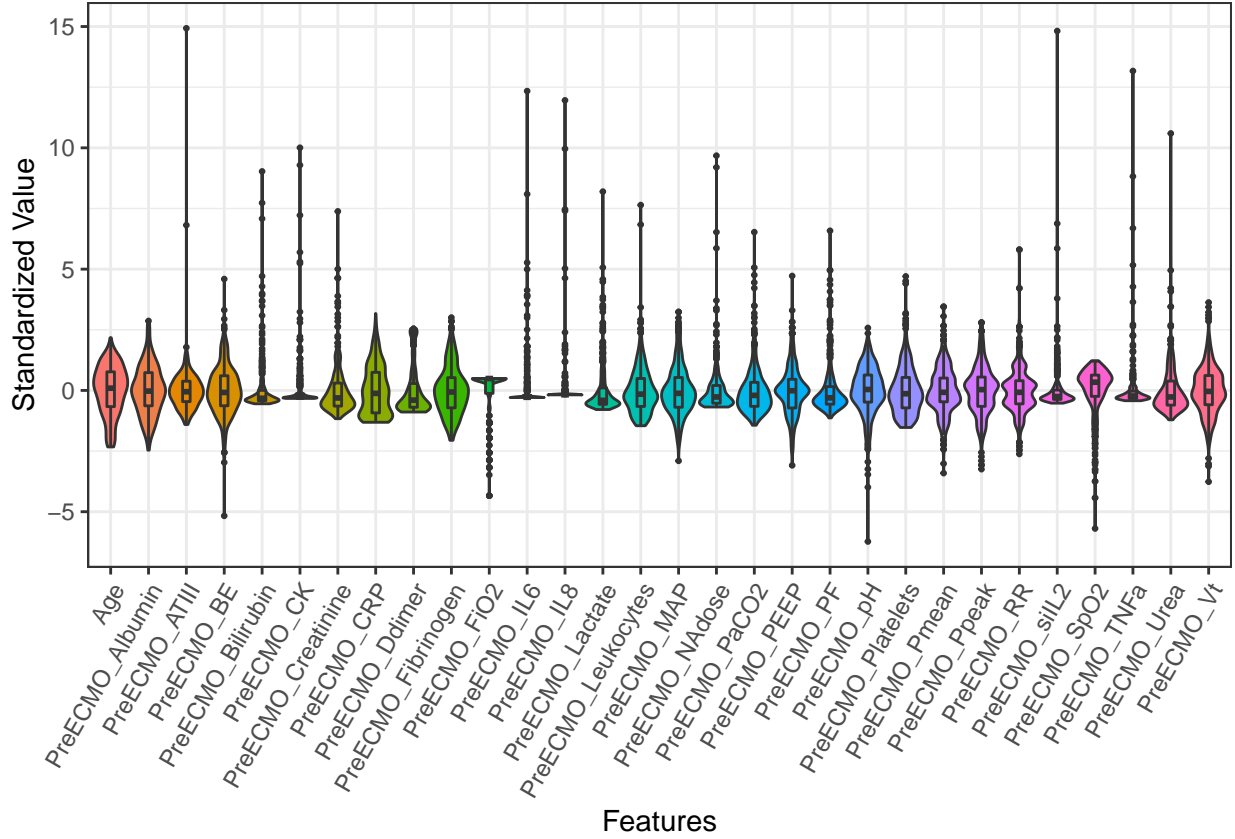


Figure 2: Violin plot of continuous variables.

3.2 Missing Data Patterns

Before imputation, and indeed multiple imputation, it is important to inspect the missingness patterns in the data and check assumptions. Figure 3 shows the missingness patterns in the dataset, where a black bar represents a missing value. Table ?? provides some measures about variable dependence in the dataset. The first row shows the probability of observed values for each variable. The following are coefficients that give insight into how the variables are connected in terms of missingness. **Influx** is the ratio of the number of variables pairs (Y_j, Y_k) with Y_j missing and Y_k observed, divided by the total number of observed data. For a variable that is entirely missing, influx is 1, and 0 for if the variable is complete. **Outflux** is defined in the opposite manner, by dividing the number of pairs (Y_j, Y_k) with Y_j observed and Y_k missing, by the total number of complete cells. For a completely observed variable, outflux will have a value of 1 and 0 if completely missing. Outflux gives an indication of how useful the variable will be for imputing other variables in the dataset, while influx is an indicator for how easily the variable can be imputed. Table 5 shows that all variables will be useful during imputation except `PreECMO_Albumin`. A high outflux variable might turn out to be useless for the imputation procedure if it is unrelated to the incomplete variables, while the usefulness of a highly predictive variable is severely limited by a low outflux value (Van Buuren 2012). **Mention 3)**

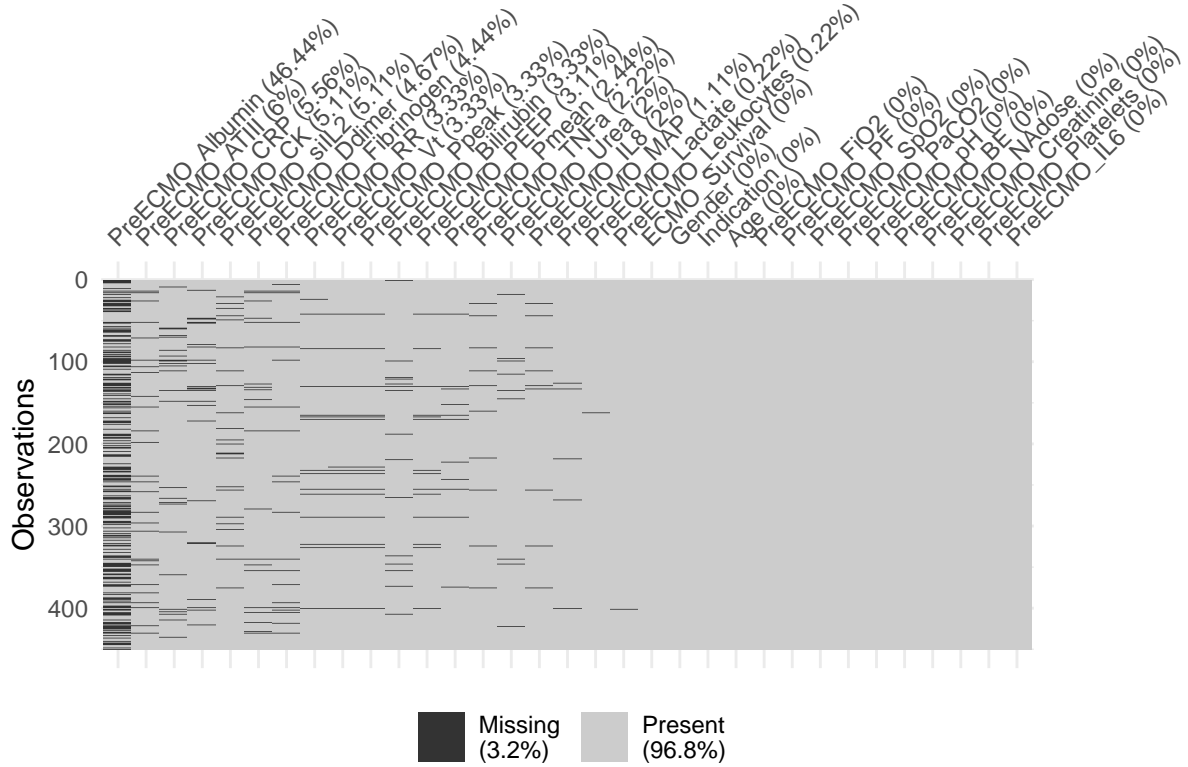


Figure 3: Visual representation of missing observations in the ARDS dataset.

- It can be difficult or impossible to determine if the data are MCAR. Figure 3 shows that many missing values occur in observations with other missing values. Missing values could be conditionally dependent on other variables, in which case the data would be MAR. The missing values could also be due to some unknown mechanism at the time of recording (*i.e.* a failure of the measurement device) that happens to effect multiple readings (the biomarkers are measured from blood samples and measurements are likely done in batches). In this case the data would be MCAR. Without more information, this analysis assumes the data is MCAR.

3.3 Experiments and Results

The experimentation phase of this study involved three methods for handling missing data: (a) complete case analysis with the variable `PreECMO_Albumin` dropped from the analysis due to 46.44% missingness, (b) mean imputation on variables with missing values, (c) imputation via the MICE algorithm implemented with PMM.

Table 6 shows the averaged Kappa from each analysis in 10-fold cross-validation. In complete case analysis and mean imputaion, LDA is the highest performer. While for predictive mean-matching with $m = 9$ logistic regression has the highest averaged Kappa.

Table 5: Missing pattern statistics for variables in dataset.

	Proportion	Influx	Outflux
ECMO_Survival	1.00	0.00	1.00
Gender	1.00	0.00	1.00
Indication	1.00	0.00	1.00
Age	1.00	0.00	1.00
PreECMO_RR	0.97	0.03	0.85
PreECMO_Vt	0.97	0.03	0.85
PreECMO_FiO2	1.00	0.00	1.00
PreECMO_Ppeak	0.97	0.03	0.85
PreECMO_Pmean	0.98	0.02	0.90
PreECMO_PEEP	0.97	0.03	0.85
PreECMO_PF	1.00	0.00	1.00
PreECMO_SpO2	1.00	0.00	1.00
PreECMO_PaCO2	1.00	0.00	1.00
PreECMO_pH	1.00	0.00	1.00
PreECMO_BE	1.00	0.00	1.00
PreECMO_Lactate	1.00	0.00	0.99
PreECMO_NAdose	1.00	0.00	1.00
PreECMO_MAP	0.99	0.01	0.97
PreECMO_Creatinine	1.00	0.00	1.00
PreECMO_Urea	0.98	0.02	0.94
PreECMO_CK	0.95	0.05	0.87
PreECMO_Bilirubin	0.97	0.03	0.91
PreECMO_Albumin	0.54	0.46	0.26
PreECMO_CRP	0.94	0.05	0.88
PreECMO_Fibrinogen	0.96	0.04	0.85
PreECMO_Ddimer	0.95	0.04	0.86
PreECMO_ATIII	0.94	0.06	0.84
PreECMO_Leukocytes	1.00	0.00	0.99
PreECMO_Platelets	1.00	0.00	1.00
PreECMO_TNFa	0.98	0.02	0.93
PreECMO_IL6	1.00	0.00	1.00
PreECMO_IL8	0.98	0.02	0.93
PreECMO_siIL2	0.95	0.05	0.87

Table 6: Averaged Cohen’s Kappa for each model fitted in cross-validation. The tuned parameters for K-Nearest Neighbors and Random Forests are K=5 and mtry=13, respectively.

	Logit	LDA	QDA	KNN	RF
Complete Case	0.139	0.205	0.038	0.053	0.035
Mean	0.191	0.220	0.040	0.136	0.085
PMM	0.179	0.124	0.106	0.088	0.136

3.3.1 Validation on Test Set

Using the parameters values learned obtained from 10-fold cross-validation in Table 6, models were fit to the full training set and validated against the test set. K-nearest neighbors and random forests were fit with parameters $K = 5$ and $mtry = 13$, respectively.

Table 7: Complete case analysis accuracy metrics. The tuned hyperparameters for K-Nearest Neighbors and Random Forests are $K=5$ and $mtry=11$, respectively.

	Sensitivity	Specificity	Accuracy	Kappa
Logit	0.20	0.814	0.658	0.015
LDA	0.20	0.847	0.684	0.054
QDA	0.00	0.966	0.722	-0.048
KNN	0.30	0.847	0.709	0.161
RF	0.05	0.966	0.734	0.022

Table 8: Mean imputation accuracy metrics. The tuned hyperparameters for K-Nearest Neighbors and Random Forests are $K=5$ and $mtry=11$, respectively.

	Sensitivity	Specificity	Accuracy	Kappa
Logit	0.259	0.871	0.723	0.147
LDA	0.222	0.859	0.705	0.091
QDA	0.111	0.906	0.714	0.021
KNN	0.259	0.835	0.696	0.102
RF	0.074	0.976	0.759	0.071

Table 9: Predictively mean-matching imputation accuracy metrics. The tuned hyperparameters for K-Nearest Neighbors and Random Forests are $K=5$ and $mtry=13$, respectively.

	Sensitivity	Specificity	Accuracy	Kappa
Logit	0.259	0.882	0.732	0.162
LDA	0.259	0.882	0.732	0.162
QDA	0.111	0.906	0.714	0.021
KNN	0.259	0.859	0.714	0.131
RF	0.074	0.953	0.741	0.037

Table 10: 99 Predictively mean-matching imputation accuracy metrics. The tuned hyperparameters for K-Nearest Neighbors and Random Forests are $K=5$ and $mtry=13$, respectively.

	Sensitivity	Specificity	Accuracy	Kappa
Logit	0.259	0.894	0.741	0.178
LDA	0.296	0.882	0.741	0.202
QDA	0.111	0.906	0.714	0.021
KNN	0.222	0.871	0.714	0.106
RF	0.148	0.941	0.750	0.116

4 Discussion

4.1 Model Performance

Logistic Regression

For complete-case analysis, mean imputation, and predictive mean-matching, logistic regression does not meet the “one in ten rule”, a rule of thumb stating that a logistic regression models give stable estimates for the covariates if there are at least 10 observations of the least frequent class per covariate.

LDA

Can perform better than logistic regression when the covariates are normally distributed (CITATION), which they are in this case after Yeo-Johnson transformation.

Random Forests Fails

- Sparsity - When the data are very sparse, it’s very plausible that for some node, the bootstrapped sample and the random subset of features will collaborate to produce an invariant feature space. There’s no productive split to be had, so it’s unlikely that the children of this node will be at all helpful.
- One surprising consequence is that trees that work well for nearest-neighbor search problems can be bad candidates for forests without sufficient subsampling, due to a lack of diversity. (Tang et al. 2018)
- Data are not axis-aligned - Suppose that there is a diagonal decision boundary in the space of two features, x_1 or x_2 . Even if this is the only relevant dimension to your data, it will take an ordinary random forest model many splits to describe that diagonal boundary. This is because each split is oriented perpendicular to the axis of either x_1 or x_2 .
- XGBoost, Rotation forest (PCA rotation) may do better

5 Conclusion

- Summary of procedure
- Summary of results
- Possible improvements and future work

6 Appendices

6.1 A. Additional Exploratory Data Analysis

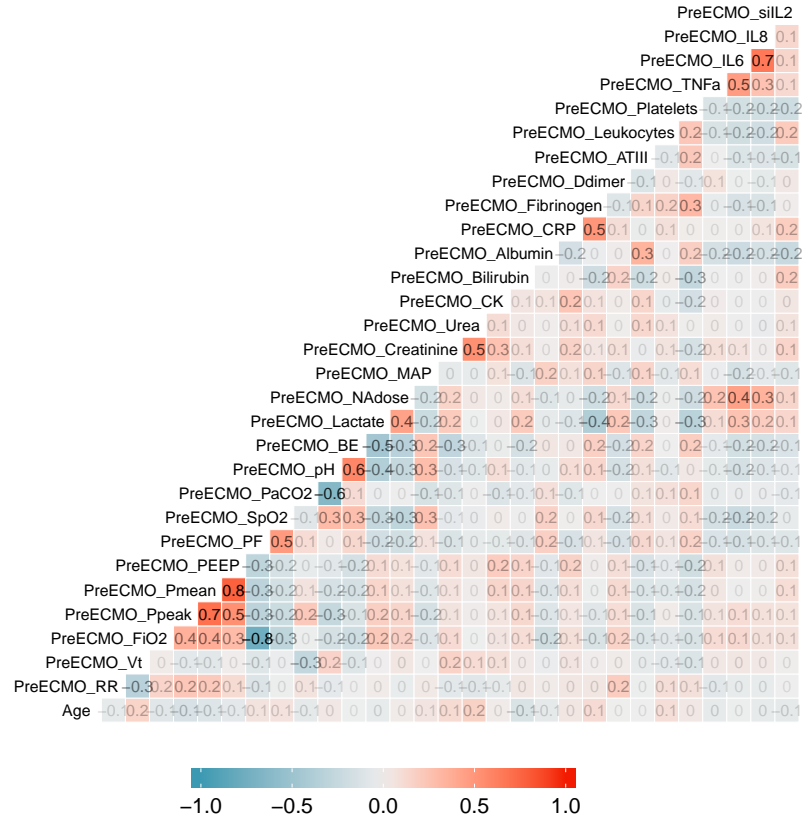


Figure 4: Heatmap of standardized and transformed variables.

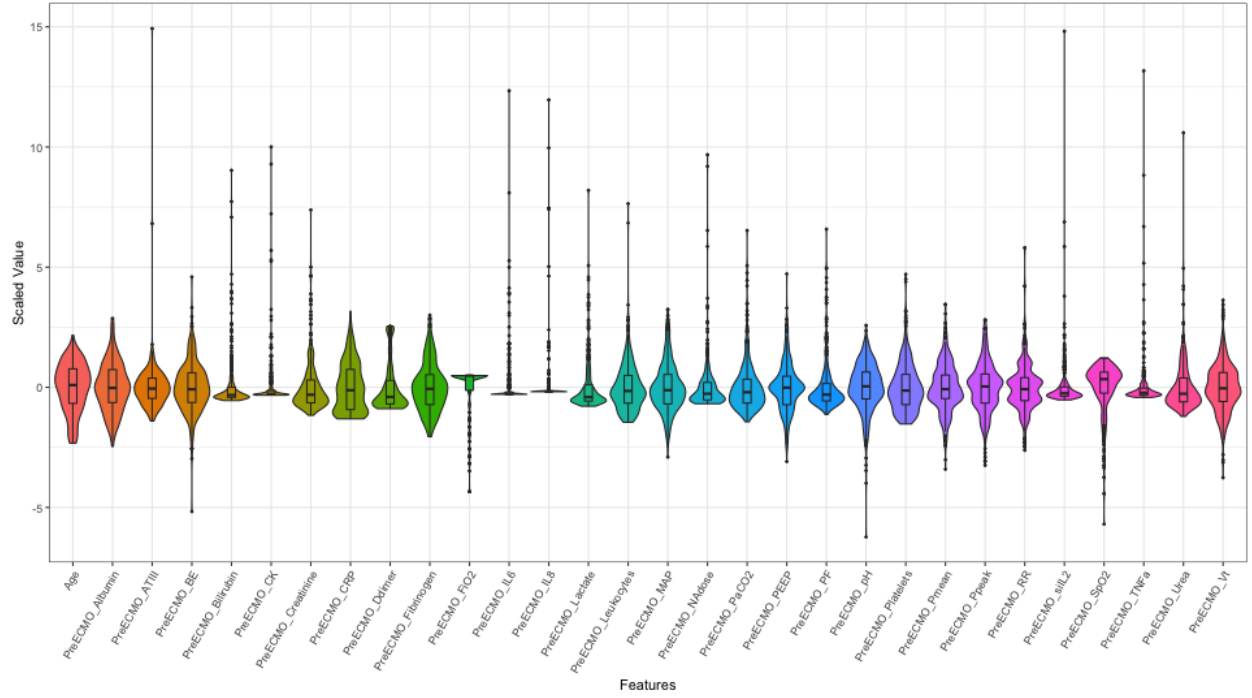


Figure 5: Violin plot of standardized variables.

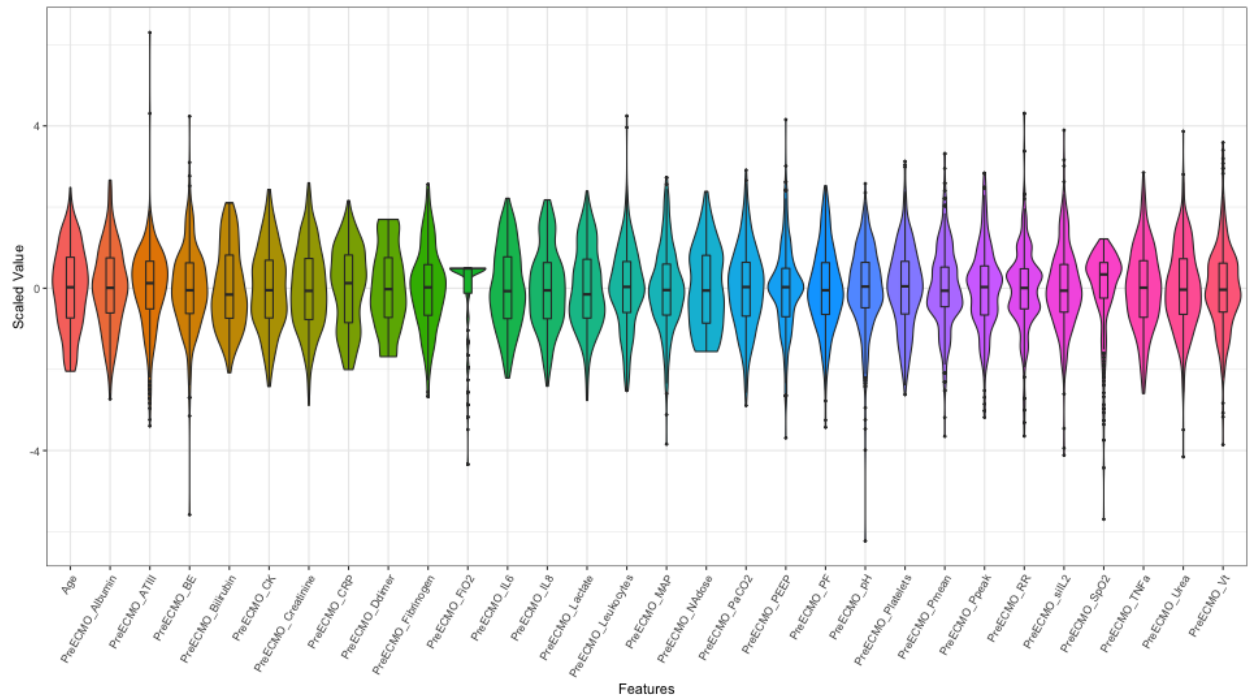


Figure 6: Violin plot of standardized and transformed variables.

6.2 B. Algorithms

6.2.1 Random Forests Algorithm

The random forests algorithm depicted is adapted from [Hastie et al., 2009].

1. For ($b = 1$ to B):
 - (a) Draw a bootstrap sample \mathbf{Z}^* of the size N from the training data.
 - (b) Grow a random-forest tree T_b to the bootstrapped data, by recursively repeating the following steps for each terminal node of the tree, until the minimum node size n_{min} is reached.
 - i. Select $mtry$ variables at random from the p covariates.
 - ii. Pick the best covariate/split-point among the $mtry$.
 - iii. Split the node into two daughter nodes.
2. Output the ensemble of trees $\{T_B\}_1^B$

Let $\hat{Y}_b(x)$ be the class prediction of the b^{th} random-forest tree. Then a new observation, x , is classified as:

$$\hat{Y}_{\text{rf}}^B(x) = \text{majority vote } \left\{ \hat{Y}_b(x) \right\}_1^B$$

Algorithm 1: Random Forest Classifier

6.2.2 MICE Algorithm

The MICE algorithm is adapted from [van Buuren, 2012].

1. Specify an imputation model $P(Y_j^{\text{mis}} | Y_j^{\text{obs}}, Y_{-j}, R)$ for variable Y_j with $j = 1, \dots, p$
2. For each j , fill in starting imputation Y_j^0 by random draws from Y_j^{obs}
3. Repeat for $t = 1, \dots, T$:
4. Repeat for $j = 1, \dots, p$:
5. Define $Y_{-j}^t = (Y_1^t, \dots, Y_{j-1}^t, Y_{j+1}^{t-1}, \dots, Y_p^{t-1})$ as the currently complete data except Y_j
6. Draw $\phi_j^t \sim P(\phi_j^t | Y_j^{\text{obs}}, Y_{-j}^t, R)$.
7. Draw imputations from $Y_j^t \sim P(Y_j^{\text{mis}} | Y_j^{\text{obs}}, Y_{-j}^t, R, \phi_j^t)$.
8. End repeat j .
9. End repeat t .

Algorithm 2: Multiple Imputation via Chained Equations

6.3 C. Additional Missing Data Diagnostics

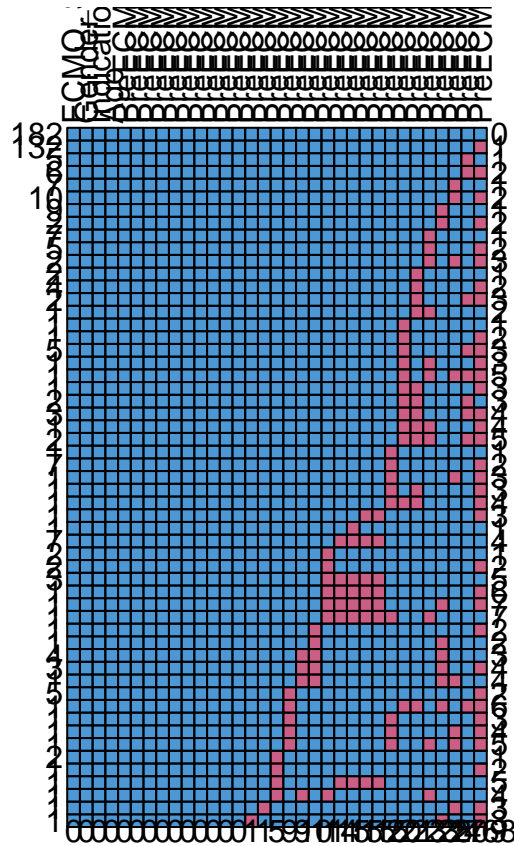


Figure 7: Missing data patterns. Each row corresponds to a missing data pattern (1=observed, 0=missing). Rows and columns are sorted in increasing amounts of missing information. The last column and row contain row and column counts, respectively.

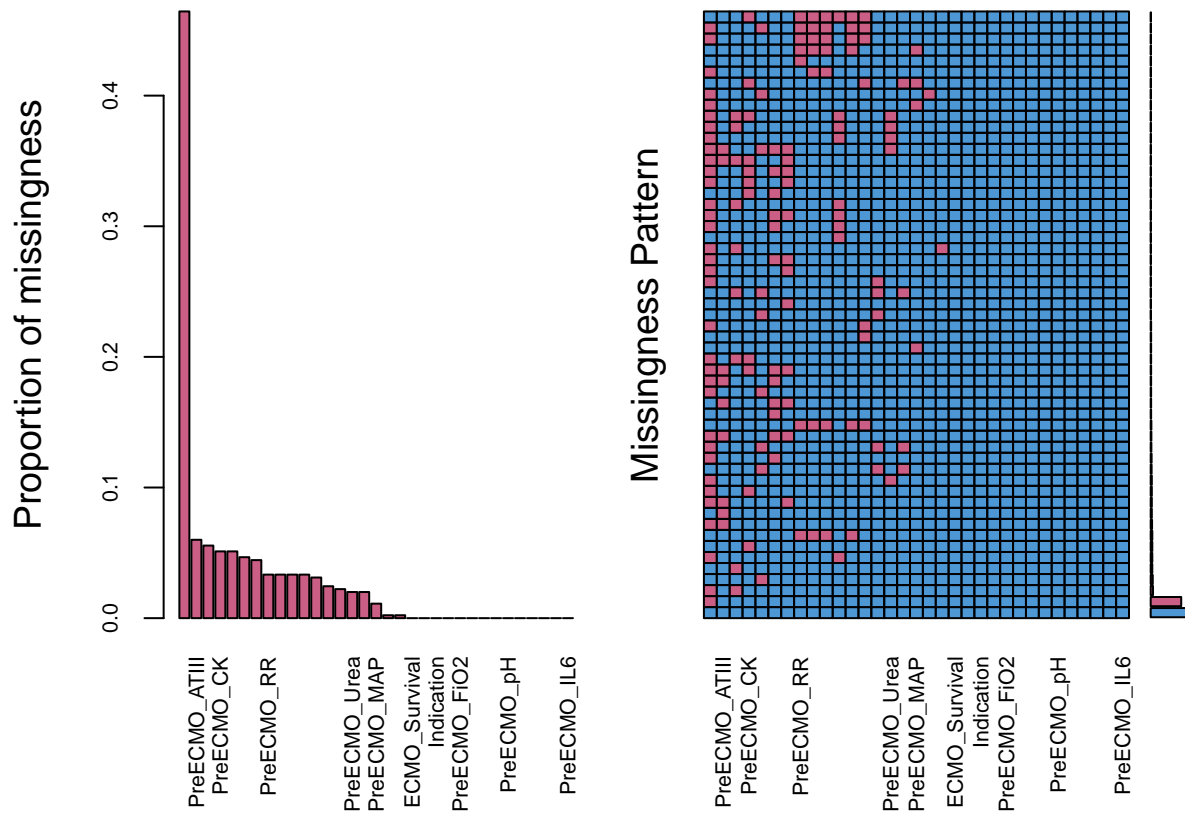


Figure 8: Missing data patterns. Each row corresponds to a missing data pattern (1=observed, 0=missing). Rows and columns are sorted in increasing amounts of missing information. The last column and row contain row and column counts, respectively.

Variables sorted by number of missings:

Variable	Count
PreECMO_Albumin	0.464444444
PreECMO_ATIII	0.060000000
PreECMO_CRP	0.055555556
PreECMO_CK	0.051111111
PreECMO_siIL2	0.051111111
PreECMO_Ddimer	0.046666667
PreECMO_Fibrinogen	0.044444444
PreECMO_RR	0.033333333
PreECMO_Vt	0.033333333
PreECMO_Ppeak	0.033333333
PreECMO_Bilirubin	0.033333333
PreECMO_PEEP	0.031111111
PreECMO_Pmean	0.024444444
PreECMO_TNFa	0.022222222
PreECMO_Urea	0.020000000

PreECMO_IL8 0.020000000
PreECMO_MAP 0.011111111
PreECMO_Lactate 0.002222222
PreECMO_Leukocytes 0.002222222
ECMO_Survival 0.000000000
Gender 0.000000000
Indication 0.000000000
Age 0.000000000
PreECMO_FiO2 0.000000000
PreECMO_PF 0.000000000
PreECMO_SpO2 0.000000000
PreECMO_PaCO2 0.000000000
PreECMO_pH 0.000000000
PreECMO_BE 0.000000000
PreECMO_NAdose 0.000000000
PreECMO_Creatinine 0.000000000
PreECMO_Platelets 0.000000000
PreECMO_IL6 0.000000000

6.3.1 Visual Insepction of Imputations

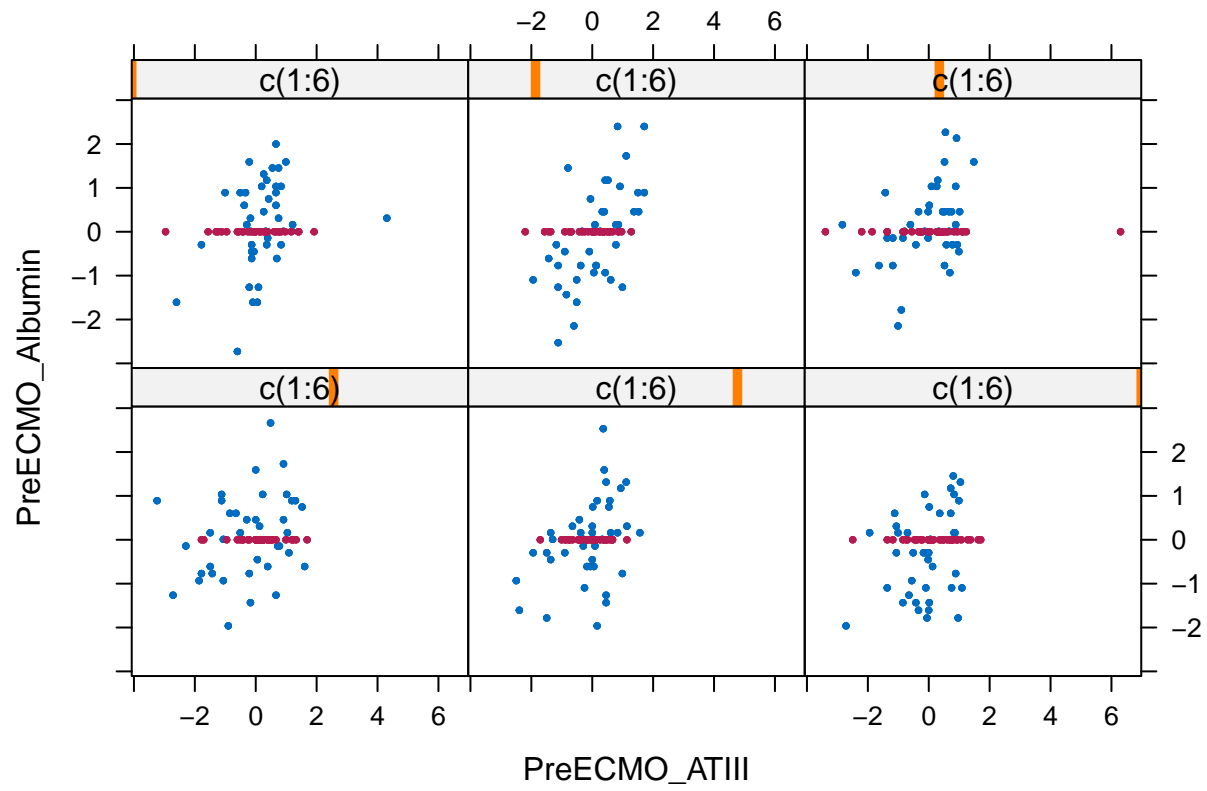


Figure 9: Scatterplot of each imputed dataset

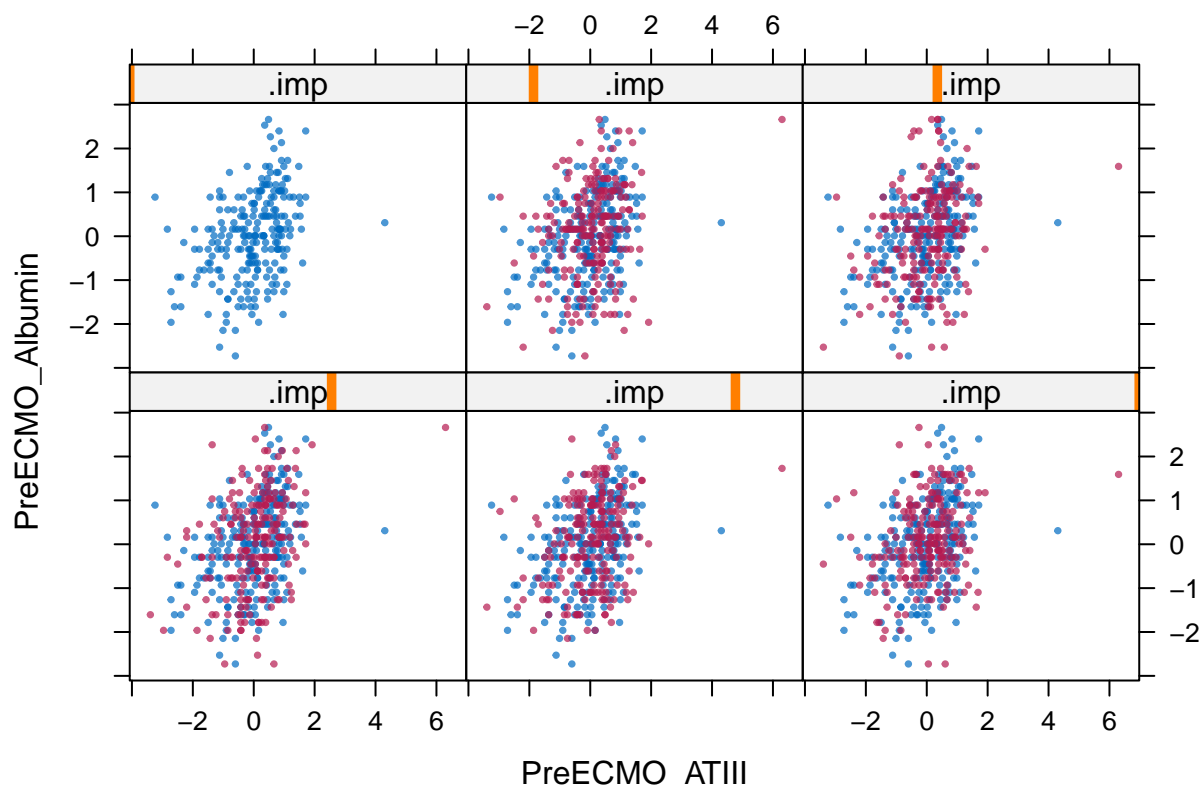


Figure 10: Scatterplot of each imputed dataset

- xyplot checking distributions of original and imputed data for MEAN imputation
- xyplot checking distributions of original and imputed data for PMM imputation
- density plot of original and imputed data for MEAN imputation
- density plot of original and imputed data for PMM imputation

This plot compares the density of observed data with the ones of imputed data. We expect them to be similar (though not identical) under MAR assumption.

6.3.2 Convergence Monitoring

- **

6.4 D. R Code

- Figure showing flow of code

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