Missing Data, Imputation and Variable Selection in Multivariate Modeling

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Motivation and Objective

- Registries – non-interventional studies
  - collect data on procedures and assessments which are considered part of standard practice.

- Leads to a lot of missing data.

- Even if 10-15% is missing per covariate, easily leads to only 40-50% of complete cases.

Objective:

Present a multivariate modeling approach involving variable selection that uses multiple imputation, model presentation and internal and external validation.
• Example – Discrete Case: Analysis of deaths within 180 days. (Note that the actual results are replaced with dummy data in data presentation). Logistic regression Modeling
  – Analysis with missing data, imputation and variable selection
  – Model Representation
  – Internal Validation
  – External Validation

• Example - Continuous Case: Analysis of MM Disease Registry baseline QOL endpoints. Clinical baseline factors as predictive of EQ5D index.
  – Introduction
  – Multivariate model selection
  – Results

• Conclusions
General Approach to Multivariate Logistic Analyses: Endpoint=Death within 180 days (yes, no)

- Univariate Logistic Regression analysis to determine significant variables to be entered into Multivariate analysis.
- 10 datasets created using multiple imputation.
- Datasets stacked and variable selection done using underweighted observations.
- 10 Unstacked multivariate logistic analyses done for the variables selected.
- Inferences combined using Rubin’s method to obtain estimates and p-values.
1a. Imputation Model

Data: \( Y = (Y_{mis}, Y_{obs}) \)

\[ = (Y_{1obs}, Y_{1mis}, Y_{2obs}, Y_{2mis}, Y_{3obs}, Y_{3mis}) \]

Distribution: \( f(y / \theta) \)

1b. Impute the Missing Values

\[ f(Y_{mis} / Y_{obs}, \theta) \]

1c. Produce Multiple Imputations

\[ \hat{Y}_{(1)} = (\hat{Y}_{mis(1)}, Y_{obs}) \quad \hat{Y}_{(m)} = (\hat{Y}_{mis(m)}, Y_{obs}) \quad \hat{Y}_{(M)} = (\hat{Y}_{mis(M)}, Y_{obs}) \]

SAS Code: Multiple Imputation

Default: MCMC Multivariate Normal

• proc mi data =Edeath nimpute = 10 seed = 651467
  out=Edeathm ;
  var age issstage ecog IMWG_risk mhdiabn Beta2_M
  mhhyn mobility VTE plat_ct........ d180; *(partial list of Variables);
  run;

Recoding of discrete variables (Can also use round option of the MI procedure)

• if mhdiabn le 0.5 then mhdiabn = 0;
• if mhdiabn gt 0.5 then mhdiabn = 1;
• if mhhyn le 0.5 then mhhyn = 0;
• if mhhyn gt 0.5 then mhhyn = 1;

Bernaards et. al. (2007) Statistics in Medicine 26:1368–1382
Procedure: Model Selection For Logistic Regression

Selection = score in SAS provides the score statistic for all possible models.

Difference in score statistic - a chi-squared distribution, with degrees of freedom given by the difference in the number of variables in the model.

Starting with best 1 variable model, move in 1 variable increments to the best k variable model, till the incremental score statistic is less than the critical value.

Select that best k variable model.
Theory: Stacked Weighted analysis. Wood et al

They Considered:

• Complete Case (CC)

• Single Stochastic Imputation (Single)

• Separate Imputations
  – S1: select predictors that appear in any model;
  – S2: select predictors that appear in at least half of the models;
  – S3: select predictors that appear in all models.

• Weighted Analysis
  – W1: $w_i = 1/M$.
  – W2: $w_i = (1 - f)/M$
  – W3: $w_i = (1 - f_i)/M$

• Rubin’s Method (RR)

Power defined as the probability of correctly selecting each of the variables in true model.

Type 1 error defined as the probability of wrongly selecting variables not in the true model.

Simulations find that W2 and W3 are close in type I error and power to the ideal but difficult to implement RR method.
• Datasets Stacked and the following CODE. Weight = \(0.9137/10\)

• proc logistic data = Edeathm2 ;
  
  model d180 = age issstage mobility IMWG_risk ecog Beta2_M VTE bm calcium creat plat_ct clcr
  /selection = score
  
  details lackfit ;

  weight wt;

  run;
• Obtain parameter estimates separately for reduced model obtained by the stacked weighted logistic regression (and clinically meaningful).

• \texttt{proc logistic data=Edeathm2;}
\texttt{model dthbf180 (event = 'Yes') = age ecog VTE mobility ISS plat_ct}
\texttt{/risklimits details lackfit covb;}
\texttt{by _Imputation_;}
\texttt{ods output ParameterEstimates=lgparms CovB=lgcovb;}
\texttt{run;}

\textit{SAS Code: Logistic regression by Imputation}
Rubin’s Inferences behind PROC MI

The MI estimates of the vector of parameters

$$\bar{\theta} = \frac{1}{M} \sum_{m=1}^{M} \hat{\theta}_m$$

Within Imputation Variance

$$\bar{W} = \frac{1}{M} \sum_{m=1}^{M} W_M$$

Between- imputation Variance

$$B = \frac{1}{M-1} \sum_{m=1}^{M} (\hat{\theta}_M - \bar{\theta})^2$$

Total variability of \( \bar{\theta} \)

$$T = \bar{W} + \frac{M + 1}{M} B$$
• Combine the estimates from the separate imputations.

• `proc mianalyze parms=lgparms
covb=lgcovb;
modeleffects Intercept age ecog VTE mobility ISS plat_ct;
ods output ParameterEstimates=est1;
run;`
• Typical Model Presentation. Odds Ratios, Confidence Intervals and P-values.
  - data est1; set est1;
  - OR = exp(Estimate);
  - ORL = exp(LCLMean);
  - ORU = exp(UCLMean);
  - Run;

• Heat Map presentation to aid in physician-patient communication. Matrix presentation of predicted values

\[
\theta_{hi} = \frac{\exp\left\{\alpha + \sum_{k=1}^{l} \beta_k x_{hik}\right\}}{1 + \exp\left\{\alpha + \sum_{k=1}^{l} \beta_k x_{hik}\right\}}
\]

## Typical Model Presentation: Logistic Regression

**Note:** Data redacted with xx.x, estimates changed and variables switched.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Univariate Analysis</th>
<th>Multivariate Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Estimate</td>
<td>95% CI</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Patient Specific</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age (&lt;=75 vs &gt;75)</td>
<td>1.97</td>
<td>(1.26, 3.06)</td>
</tr>
<tr>
<td>Body Mass Index</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>ECOG performance status score</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>History of diabetes</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>History of hypertension</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>History of venous thromboembolism (VTE)</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>Del(17P) from FISH and Cytogenetic forms</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>T(414) from FISH</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td><strong>Disease Specific</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lactic Acid Hydrogena (&lt;= 300 g/dL vs &gt; 300 g/dL)</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>Extramedullary plasmacytoma (Yes/No)</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>Immunoglobulin IgG Class (&lt; 5 g/dL vs &gt;= 5 g/dL)</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>Albumin (&lt;=3.5 g/dL vs &gt;3.5 g/dL)</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>ISS disease stage (calculated)</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>Myeloma bone involvement</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>Hypercalcemia (Serum Calcium &gt;=11.5 mg/dL)</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>Renal Insufficiency (Serum Creatinine &gt; 2 mg/dL)</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>Anemia (Hemoglobin &lt; 10 g/dL or &gt;2 below LLN)</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>Platelet Count (for 100 x 10^9L increments)</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>INWG risk</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>Beta 2 Microglobulin (beta&gt;=5.5 mg/L)</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>HRQoL from Eq5d</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Self care from Eq5D</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>Mobility from Eq5D</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
<tr>
<td>Novel Therapy</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Novel Therapy Use (0,1) vs &gt;=2</td>
<td>x.xx</td>
<td>(x.xx, x.xx)</td>
</tr>
</tbody>
</table>
Note: Estimates changed and variables switched and/or changed

Figure 1: Inputs to Prognostic Chart

<table>
<thead>
<tr>
<th>Attributes</th>
<th>Select Value from Drop-Down List</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Mobility</td>
<td>Some Problem in Walking About</td>
</tr>
<tr>
<td>2. Platelet Count in X10^9/L</td>
<td>&lt;=150</td>
</tr>
<tr>
<td>3. ISS Stage</td>
<td>I, II</td>
</tr>
<tr>
<td>4. Age Group</td>
<td>&lt;=75</td>
</tr>
<tr>
<td>5. ECOG Performance Poor</td>
<td>No</td>
</tr>
<tr>
<td>6. History of VTE</td>
<td>No</td>
</tr>
</tbody>
</table>

Figure 2: Prognostic Chart with Estimated Probability of Death Before 180 Days

Confined to bed | Some problem in | No problem in |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Confined to bed</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Confined to bed</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Confined to bed</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Selected Attributes are highlighted in Blue and the estimated probability is *Bold, italics and underlined*. For a prognostic chart without the blue highlights or the marked up probability enter blanks for the first two inputs.
• Larger Blocks in the matrix are the larger effects.
• We move to smaller blocks within the larger blocks with factors which have succeeding smaller effects.
• Traffic Color coded. Green for lower probability of dying within 180 days, through Yellow to Red for higher probabilities.
• Designed to show higher risk in the Bottom Left corner and lower risk towards the Top Right corner.
Exit from slide review mode and click excel attachment below
Internal Cross validation of the Model for Deaths Before 180 Days

- Internal validation of the model was done using the concordance index and internal bootstrap re-sampling methods. R rms package.

- Concordance probability (Harrell’s C-Index) is the probability that a randomly selected pair of patients, one with a poorer survival outcome than the other, will be correctly differentially identified based on inputting the two patient’s baseline prognostic characteristics in the fitted model.

- For the logistic model the concordance probability is identical to the area under the receiver operating characteristic (ROC) curve for the model.

Validation Steps – Logistic Analysis

- Uses R package “rms”.
- Import each of the 10 imputed datasets into R and run the following R code for each dataset
  - library("rms")
  - ## Imputation # 1
  - f <- lrm(dthbf180 ~ agen+iss+ecog+platcount+mobility+VTE, data = impt1log, x=TRUE, y=TRUE)
  - validate(f,B=100, dxy = TRUE)
- Use Concordance probability, C-Index =\(0.5*|Dxy|+0.5\) where Dxy is the Somer’s D statistic output by R.
- Find the mean of the Concordance probabilities for test and training over the 10 imputations.
- Obtain the percent reduction in Concordance Probability as
  \[100*\{(C \text{ Avg for training)} – (C \text{ Avg for Test})\}/(C \text{ Avg for training)}\]
- Calculate the bootstrap adjusted C or AUC as C Avg for C-index corrected.
- Calculate the 95% CI for this AUC using expressions from the reference below.

**Confidence Interval for AUC**

Let $AUC$ denote the sample AUC value. For large samples, the distribution of AUC is approximately normal. Hence, a $100(1 - \alpha)\%$ confidence interval for AUC may be computed using the standard normal distribution as follows

$$AUC \pm z_{\alpha/2} SE(AUC)$$

The width of the confidence interval is $2z_{\alpha/2} SE(AUC)$. One-sided limits may be obtained by replacing $\alpha/2$ by $\alpha$. The formula for $SE(AUC)$ was given by Hanley and McNeil (1982) is

$$SE(AUC) = \sqrt{\frac{AUC(1 - AUC) + (N_1 - 1)(Q_1 - AUC^2) + (N_2 - 1)(Q_2 - AUC^2)}{N_1N_2}}$$

where

$$Q_1 = \frac{AUC}{2 - AUC}$$

$$Q_2 = \frac{2AUC^2}{1 + AUC}$$

N1 is the # of deaths in 180 days and N2 is the number alive at 180 days.
• The percent reduction in the concordance probability in the test bootstrap re-sampling estimate compared to the training bootstrap estimate was 2.53%.

• The training optimism adjusted concordance probability of the fitted logistic model was estimated as 73.00% with a 95% confidence interval of (67.29%, 78.69%). A concordance probability significantly greater than 50% is indicative of a good predictive model.
Getting Concordance Probability:

From the combined mianalyze coefficient estimates of the multiple imputation analysis derive predictions in the test dataset as follows:

- `logist <- read.csv("Z:/MedicalAffairs-MM/Connect-MM/Stats/Albert/Death within 180 Days/Validation/CC 4013 015/d_mm_015logit.csv", header = T)`
- `library(rms)`
- `phat <- 1/(1+exp(-(-.68+(0.64*logist$mobf+0.70*logist$ecog+ 0.49*logist$VTE+ 0.50*logist$issstage+0.49*logist$agen+0.81*logist$plat_ct))))`
- `val.prob(phat, logist$dthbf180, xlab="Predicted Probability of Death Before 180 Days ", ylab="Actual Probability of Death Before 180 Days ",lim=c(0, .3), m= 30, cex = 0.5)`
- `text(0.05,0.25, "C-Index = 0.766", cex = 0.8)`

The C-Index (along with a host of other statistics) is provided when you run the analysis without “lim =c(0,0.3)” above.

Note: Parameter estimates changed to mask data.
Validation Using External Study: Logistic Regression

C-index = 0.766
ANALYSIS OF THE MM DISEASE REGISTRY BASELINE CONTINUOUS QOL ENDPOINTS

1. Univariate Regression Analysis to determine which variables to enter into Multivariate.
2. 10 datasets created using multiple imputation.
3. Datasets stacked and variable selection was done using weighted SSE.
4. 10 Unstacked multivariate regression analyses done for the variables selected.
5. Inferences combined using Rubin’s method to obtain estimates and p-values.
Procedure: Model Selection for Continuous Case

Determine the best 1 variable model to best p variable model.

Compute scaled deviance = SSE/(sigma squared) for the best 1 variable to the best p variable model.

The scaled deviance = -2*LOG(Likelihood) is a chi-squared score statistic.

Difference in scaled deviance is chi-squared with d.f. given by the difference in the number of variables in the Model.

Starting with best 1 variable model, move in 1 variable increments to the best k variable model. Use 1 d.f chi-square critical value.

• SAS PROC REG was run against the stacked dataset with option `selection = maxr`.

• Analysis weighted by mean percent of non-missing observations divided by the number of imputations.

• SSE recorded for each best model

\[ \text{SigmaSqr}(d) = \frac{\text{Saturated SSE}}{(\text{Nobs} - \text{Nparm} - 1)} \]
• Predictor selected in the best 1-variable model was included by default

• For 2-variable model and up,
  – the scaled deviance $= \frac{SSE}{\text{SigmaSqr}}$

• The difference between incremental models
  – $DEV_{dif_i} = DEV_i - DEV_{i-1}$
• Difference in scaled deviance between two models differing by one variable has a chi-squared distribution with 1 d.f.

• Continue adding variables while
  – \{\text{abs}(\text{DEVdif}_i) > 2.71}\}

• For list of variables identified,
  – SAS PROC GLM was run by imputation
  – Combined using SAS PROC MIANALYZE.
### EQ-5D final results after MIANALIZE: Imputed vs. Un-imputed Complete Case Analysis

<table>
<thead>
<tr>
<th>Variable</th>
<th>Estimate</th>
<th>P-val Imputed model</th>
<th>Estimate</th>
<th>P-val Un-imputed model</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECOG</td>
<td>-0.08</td>
<td>&lt;0.001</td>
<td>-0.11</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>BONEINV</td>
<td>-0.06</td>
<td>&lt;0.001</td>
<td>-0.08</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>HISPAN</td>
<td>-0.08</td>
<td>&lt;0.001</td>
<td>-0.11</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>DSSTAGE</td>
<td>-0.01</td>
<td>0.067</td>
<td>-0.02</td>
<td>0.058</td>
</tr>
<tr>
<td>AGEGRP</td>
<td>0.02</td>
<td>0.005</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ISS3</td>
<td>-0.02</td>
<td>0.117</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SEX</td>
<td>0.02</td>
<td>0.045</td>
<td>0.03</td>
<td>0.033</td>
</tr>
<tr>
<td>CALCIUM</td>
<td>-0.04</td>
<td>0.08</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Conclusions

• With Registries missing data analysis can be tied into multivariate analysis efficiently using a few SAS proc calls.

• Features
  – A weighted stacked analysis can be used for variable selection.
  – For the exponential family including the normal continuous case the deviance statistic \((-2*\text{LOG(Likelihood)}\) can be used for variable selection as the difference in deviance is a chi-squared statistic.
  – For logistic regression the SAS score statistic can be used.
  – Ordinary regression one needs the scaled deviance obtained as the (stacked weighted SSE)/(Saturated Sigma Square).
  – Model can be presented in user friendly manner.
  – R package rms can be used for internal and external validation for logistic, regression and survival cases.