Computation of a cardiac severity score with left-censored biomarkers for patients with heart failure

Introduction: Biomarkers reported as being less than the limit of detection (LOD) are challenging to be included in analyses without being dichotomized. Substitutions by the LOD, LOD/2, LOD/4, or zero have been attempted for left-censored values. We calculated a novel Modified Myocardial Injury Summary Score (MMISS) by incorporating four biomarkers (B-type natriuretic peptide, troponin, galectin-3, and suppression of tumorigenicity 2) to efficiently stratify heart failure (HF) patients. The objective of this study is to evaluate the differences in associations between MMISS, calculated by different methods of substitution, and therapeutic intensity index (TII), a composite pharmacologic score.

Methods and results: This is a cross-sectional study including 39 HF patients aged ≥18 years who were treated in the Baylor Health Care System. MMISS was calculated with the left-censored biomarkers substituted by zero, LOD, LOD/2, and LOD/4. Patients with biomarker values more than LOD (complete cases) were also considered separately. The computed TII was regressed separately on MMISS for each substitution method, while controlling for age and gender. All substitution methods yielded negative associations; however, statistical significance for the association was not achieved using substitution by zero or when considering only the complete cases. The association was quite comparable with the substitution of left-censored values by LOD, LOD/2, and LOD/4.

Conclusion: Substantial loss of information is inevitable if only the data with values above the LOD are considered for analysis or when the left-censored values are substituted with zero.

Keywords: cardiac biomarkers, left-censored, limit of detection, therapeutic index, myocardial injury summary score

Introduction
Heart failure (HF) is associated with significant morbidity and mortality. According to the National Vital Statistics, age-adjusted HF mortality in the USA decreased from 2000 through 2012 but increased for the period of 2012–2014.1 HF is a complex condition where either the heart fails to pump blood adequately or has impaired ventricular filling. HF may be chronic when the symptoms appear slowly with progressive worsening of the condition or it may be acute most often following a heart attack or sudden inability to compensate the existing HF. Cardiac biomarkers are routinely used in the treatment of HF for diagnostic and prognostic purposes. Four biomarkers, namely troponin I or T, B-type natriuretic peptide or N-terminal (BNP), galectin-3, and suppression of tumorigenicity 2 (ST2), have been recommended by the American College of Cardiology Foundation (ACCF) and the American Heart Association (AHA) for effective management of HF.2 We created a novel score incorporating these four biomarkers so as to efficiently stratify and therapeutically manage patients with HF. Quantification
of these biomarkers is dependent on the sensitivity of the
diagnostic tool to detect the minimum reportable levels.
In instances where the biomarkers are not detectable, they
are simply reported as being less than the limit of detection
(LOD) by the laboratory. Biomarkers reported as being less
than the LOD are a challenge to be included in analyses
without use of dichotomization, as their quantitative levels
cannot be precisely defined. Substitution is performed in
those instances for the biomarkers reported below the LOD,
as they are essential for calculation of this prognostic score at
the bedside. Traditional methods have advocated substitution
with the LOD, LOD/2, LOD/√2, or zero for the left-censored
values. The objective of this study is to evaluate the differ-
ces in the association between therapeutic intensity index
(TII) and the Modified Myocardial Injury Summary Score
(MMISS), as calculated via different methods of substitution
for the left-censored biomarkers.

Methods
This is a cross-sectional study including HF patients aged
≥18 years who were treated at the outpatient clinics of the
Baylor Health Care System. HF was defined as mentioned in
the 2013 ACCF/AHA guidelines, which further categorized
patients with ejection fraction (EF) ≤40 as HF with reduced
EF (HFrEF). Values of the biomarkers were obtained at base-
line and were used to calculate the MMISS, a novel score
to better stratify HF patients. For biomarkers with values
reported simply as “below the LOD”, we imputed values
using the LOD, LOD/2, LOD/√2, and zero to calculate the
MMISS. Additionally, only those patients with biomarker
values more than LOD were considered separately. The
laboratory equipment was able to detect values as low as
0.02 ng/mL for troponin and 10 pg/mL for BNP.

Calculation of MMISS
The MMISS is calculated as an average of log-transformed
ratios of a biomarker’s present value to its previous value
(obtained during the preceding visit). As this is a cross-
sectional study, subjects did not have biomarker values avail-
able from a preceding visit. Thus, the upper limit normal was
used as a surrogate value for the prior clinical visit. That is

\[ \text{MMISS} = \frac{\sum_{i=1}^{n} \log_{10} \frac{\text{Biomarker}(i)_{\text{baseline}}}{\text{Biomarker}(i)_{\text{ULN}}}}{n} \]

where \( i \) is the biomarker and \( n \) is the total number of avail-
able biomarkers.

Calculation of TII
Based on the medications prescribed at the time of the bio-
marker draw, a TII was calculated. Based on the intensity
of the therapy, scores were assigned for each medication
and was found to range from 0 to 24. The drug classes
included for calculation of the TII were long-acting nitrates,
beta-adrenergic blockers, angiotensin-converting enzyme
inhibitors, angiotensin receptor blockers, mineralocorticoid
receptor antagonists, hydralazine, digoxin, and loop diuretics.

Statistical analysis
The baseline characteristics of the patients included in the
analyses were presented as means or medians for continuous
variables and proportions for categorical variables. Compara-
sions were performed using the Wilcoxon rank-sum test or
Student’s t-test for continuous variables and the chi-square
test or Fisher’s exact test for proportions, where applicable.

Separate regression models were built using TII as
the dependent variable and MMISS (as calculated via dif-
fferent substitution methods) as the independent variable,
while controlling for age and gender. The objective was
to identify any notable differences in the association of
TII and MMISS, as obtained by the different substitution
methods. The regression coefficient, standard error, and its
statistical significance were compared for all substitution
methods. Analyses were completed using STATA 14.1 and
\( p \) values <0.05 were considered statistically significant.
Posthoc power calculation was conducted using SAS 9.4
(Cary, NC, USA). With a sample size of 39 patients, we had
~88% power to detect a partial correlation of −0.48 between
MMISS and TII while controlling for age and gender. This
study was approved by the Baylor Institutional Review
Board with a waiver of consent as this study involved an
anonymous review of existing medical information posing
minimal risk to the subjects.

Results
There were 39 patients included in this study, and their char-
acteristics are in Table 1. According to the New York Heart
Association (NYHA) functional classification, 13 (33.3%)
patients had class II symptoms, 22 (56.4%) had class III
symptoms, and the remaining 4 (10.3%) patients had class
IV symptoms. The mean left ventricular ejection fraction
was 40.6±17.6; 48.7% with HFrEF. The components of TII
and the frequency of the drugs prescribed in each of the
categories are tabulated in Table 2. There were 4 (10.3%) and 29 (74.4%) instances with left-censored values for BNP and troponin, respectively. Only 9 (23.1%) patients had all biomarker values noted above the LOD and were included in the complete case analysis.

MMISS calculated for the complete cases and by different substitution methods for left-censored biomarkers (LOD, LOD/2, LOD/√2, zero) are represented in Table 3 and graphically by gender (Figure 1) and NYHA functional classification (Figure 2). The regression parameter estimates for MMISS under different substitution methods are in Table 4. Each method revealed a negative association between MMISS and TII in univariate and multivariate analyses. In the multivariate analyses, the regression coefficients of MMISS obtained by including the complete data (patients with available biomarker values more than LOD; \( p = 0.28 \)) and also by substituting zeroes for the left-censored values were not statistically significant \( (p=0.05) \) with an adjusted \( R^2 \) of 0.07. The regression coefficients of MMISS obtained by substituting the left-censored biomarkers with LOD \( (p=0.02) \), LOD/2 \( (p=0.01) \), and LOD/√2 \( (p=0.02) \) were statistically significant with an adjusted \( R^2 \) of 0.11, 0.13, and 0.12, respectively.

**Discussion**

The association between TII and MMISS varied with the different methods of substitution for the left-censored biomarkers. Although all the methods yielded a negative relation, the association did not reach statistical significance when substituted by zero or when only cases with values more than LOD were included. The association was quite comparable with the substitution of left-censored values by LOD, LOD/2, and LOD/√2. We found important relationships between MMISS and NYHA as observed in Figure 2, demonstrating that all four imputation methods had graded increases in MMISS by NYHA classification, suggesting biologic responsiveness to the severity of disease.

Analysis of left-censored biomarker data need special attention as different methods may lead to different conclusions. Often, censored biomarkers are dichotomized so as to explore their relation with the clinical outcomes.\(^3\) In situations where >10% of the values are left-censored, it has been suggested to use multiple imputation techniques for effective estimates. Tobit analysis has also been recommended to statistically model left-censored detections for robust measurements of association.\(^4\) We employed only the traditional substitution methods as our intention was to incorporate the biomarkers in a prognostic clinical score for effective management of patients with HF. It has been suggested that these traditional methods of substitution can be used, provided they do not compromise the validity of the study.\(^5\) Ignoring those values noted as below LOD or substituting them with a zero introduces unnecessary bias resulting in erroneous assumptions. In a clinical setting, values below the LOD may be useful in establishing a diagnosis but would not be able to offer details for further stratification of the severity of the disease.

In 2013, the ACCF and the AHA suggested that four blood biomarkers could be useful in advanced risk strati-
Figure 1 Distribution of modified myocardial injury by different methods of substitution of left-censored biomarkers by gender. Abbreviation: LOD, limit of detection.

Figure 2 Median values of modified myocardial injury by different methods of substitution of left-censored biomarkers by NYHA functional class. Abbreviations: NYHA, New York Heart Association; LOD, limit of detection.

fication and guideline-directed medical therapy in patients with chronic HF. These markers include troponin I or T, BNP or N-terminal BNP, galectin-3, and ST2. Trials have shown BNP or NT-pro BNP guided therapy to be associated with reduced mortality but none of the previous trials have incorporated troponin I or T, galectin-3, or ST2. ST2 has been recommended as prognostic markers in patients suffering from myocardial infarction, HF, and dyspnea. Studies
have also associated ST2 with plaque development and rupture of atheromas. We incorporated the four biomarkers in a prognostic tool that can be used at the bedside for effective clinical management.

The study has all the potential limitations of a cross-sectional study with a limited sample size. No causal inference can be made as the data were collected at a single point in time. As the true regression coefficient is unknown, we are unsure of the acceptable substitution methodology. To further study the performance of the different substitution methods (LOD, LOD/2, and LOD/√2), we would need a larger sample size with biomarker values collected sequentially over time.

Conclusion
Substantial loss of information becomes inevitable if only the data with values above LOD are considered for analysis or when the left-censored values are substituted with a zero. There are several substitution methods for left censoring that yield similar results. The MMISS is not associated with the intensity of medications prescribed for HF, but does appear to be quantitatively related to the physician-assigned severity of illness.

Disclosure
The authors report no conflicts of interest in this work.

References