Clinical Implications of the New York Heart Association Classification

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Background—The New York Heart Association (NYHA) classification has served as a fundamental tool for risk stratification of heart failure (HF) and determines clinical trial eligibility and candidacy for drugs and devices. However, its ability to adequately stratify risk is unclear.

Methods and Results—To compare NYHA class with objective assessments and survival in patients with HF, we performed secondary analyses of 4 multicenter National Institutes of Health–funded HF clinical trials that included patients classified as NYHA class II or III: TOPCAT (Treatment of Preserved Cardiac Function Heart Failure with an Aldosterone Antagonist), DIG (The Effect of Digoxin on Mortality and Morbidity in Patients With Heart Failure), HF-ACTION (Efficacy and Safety of Exercise Training in Patients With Chronic Heart Failure), and GUIDE-IT (Guiding Evidence-Based Therapy Using Biomarker Intensified Treatment in Heart Failure). Twenty-month cumulative survival was compared between classes using Kaplan–Meier curves and the log rank test. NT-proBNP (N-terminal pro–B-type natriuretic peptide), Kansas City Cardiomyopathy Questionnaire (KCCQ) scores, 6-minute walk distances, left ventricular ejection fraction, and cardiopulmonary test parameters were compared using Wilcoxon rank sum tests and percentage overlap using kernel density estimations. Cumulative mortality varied significantly across NYHA classes and HF clinical trials (likelihood ratio, P<0.001). Mortality at 20 months for NYHA class II ranged from 7% for patients in HF-ACTION to 15% in GUIDE-IT, whereas mortality for NYHA class III ranged from 12% in TOPCAT to 26% in GUIDE-IT. There was substantial percentage overlap in values for NT-proBNP levels (79% and 69%), KCCQ scores (63% and 54%), 6-minute walk distances (63% and 54%), and left ventricular ejection fraction (88% and 83%). Similarly, there was substantial overall in values for minute ventilation–carbon dioxide production relationship (71%), maximal oxygen uptake (54%), and exercise duration (53%).

Conclusions—The NYHA system poorly discriminates HF patients across the spectrum of functional impairment. These findings raise important questions about the need for improved phenotyping of these patients to facilitate risk stratification and response to interventions. (J Am Heart Assoc. 2019;8:e014240. DOI: 10.1161/JAHA.119.014240.)

Key Words: clinical trials • heart disease • heart failure • NYHA class

A simple functional classification of heart failure (HF) patients first suggested by the New York Heart Association (NYHA) has been used clinically for almost a century.1 It has long served as a foundational tool for risk stratification of HF and determines clinical trial eligibility and candidacy for drugs and devices. Whereas it is widely acknowledged that NYHA classification is subjective and has low reproducibility, its use is ingrained in both guidelines and contemporary practice, and it serves as a cornerstone of clinical documentation, trial enrollment, and candidacy for therapeutics in HF.2–3 This use has implications for the success of further interventions: currently on ClinicalTrials.gov, 304 ongoing studies have the NYHA classification as an inclusion or exclusion criterion. As a result, guideline recommendations and FDA approval of invasive interventions such as cardiac resynchronization therapy, implantable pulmonary artery pressure monitoring (CardioMEMS HF System (Abbott)), and left ventricular assist devices are firmly anchored in NYHA class.4–6

Despite the ubiquity of the NYHA classification system in HF, its clinical implications are less clear. There is no consistent method for accurate assessment of functional
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Clinical Perspective

What Is New?

• The New York Heart Association (NYHA) functional classification serves as a fundamental descriptor of heart failure and is used clinically and to determine trial eligibility.
• Using data from published trials, we found that NYHA class II versus class III is an unreliable predictor of adverse outcomes in heart failure and poorly discriminates among patients across the spectrum of functional impairment.

What Are the Clinical Implications?

• Continued usage of NYHA class in guidelines and trials, for US Food and Drug Administration approval of therapies, and for clinical decision making may hinder efforts to bring precision medicine to the bedside of heart failure patients.

class, and its relations with objective measures of HF (eg, NT-proBNP [N-terminal pro–B-type natriuretic peptide]) are unknown.7 Consequently, we sought to examine the association of NYHA class with adverse outcomes and objective measures of HF in previously published landmark clinical trials.

Methods

Data from the following National Institutes of Health–funded HF clinical trials were used to examine the association of NYHA functional class with survival: TOPCAT (Treatment of Preserved Cardiac Function Heart Failure With an Aldosterone Antagonist), DIG (The Effect of Digoxin on Mortality and Morbidity in Patients With Heart Failure), HF-ACTION (Efficacy and Safety of Exercise Training in Patients With Chronic Heart Failure), and GUIDE-IT (Guiding Evidence-Based Therapy Using Biomarker Intensiﬁed Treatment in Heart Failure).8−11 TOPCAT and DIG trial data were obtained from the publicly available database BioLINCC, whereas HF-ACTION and GUIDE-IT data were obtained from the Duke Clinical Research Institute (DCRI).12 TOPCAT patients from Russia and the Republic of Georgia were excluded because of concerns that these patients were misclassiﬁed as having HF.13 NYHA classes II and III were used, given the low number of patients classiﬁed as class I or IV in all trials. Kaplan–Meier failure curves were created to illustrate time to all-cause mortality up to 20 months from randomization; Kaplan–Meier failure rates at 20 months were reported, and pairwise comparisons were performed comparing all class II and class III rates across trials. Distributions of the following commonly used objective HF variables were overlaid according to NYHA class from HF-ACTION and GUIDE-IT: left ventricular ejection fraction, 6-minute walk distance, NT-proBNP, and Kansas City Cardiomyopathy Questionnaire (KCCQ) overall score. Finally, distributions of variables representing the gold standard for functional status in HF—cardiopulmonary exercise testing—were plotted according to NYHA class in HF-ACTION. Median values with the 25th percentile (first quartile) and 75th percentile (third quartile) are presented in the plots. The Wilcoxon rank sum test was used to evaluate differences in distributions. The percentage overlap between classes II and III was calculated by estimating the overlapping area of the 2 kernel density estimations for each objective measure. Two-tailed P<0.05 was considered statistically significant. All analyses were carried out using SAS v9.4 (SAS Institute) and R v3.4.2 (R Foundation for Statistical Computing). The institutional review boards at Yale University School of Medicine and DCRI approved the study and waived the requirement for informed consent. Because of the sensitive nature of the data collected for this study, requests to access the data set from qualiﬁed researchers trained in human subject conﬁdentiality protocols may be sent to Brooke Alhanti at DCRI (brooke.alhanti@duke.edu).

Results

Cumulative mortality varied signiﬁcantly across NYHA class and clinical trial, ranging from ≈7% to ≈25% (overall likelihood ratio, P<0.001; Figure 1). Those who were NYHA class III in TOPCAT had survival similar to those characterized as NYHA class II in the GUIDE-IT and DIG trials. Mortality at 20 months for NYHA class II was 7.0% for HF-ACTION, 8.1% for TOPCAT, 14.3% for DIG, and 15.0% for GUIDE-IT. Mortality for NYHA class III was 12.1% for TOPCAT, 13.6% for HF-ACTION, 24.3% for DIG, and 26.5% for GUIDE-IT.

Distributions for objective HF variables assessed in GUIDE-IT and HF-ACTION, stratified by NYHA class, are shown in Figure 2. Numbers of patients classiﬁed as NYHA classes II and III, respectively, were as follows: GUIDE-IT, n=447 and n=358; HF-ACTION, n=1477 and n=831. The percentage of overlap among patients who were classiﬁed as NYHA classes II and III, respectively, was as follows: NT-proBNP levels, 79% and 69%; KCCQ, 63% and 54%; 6-minute walk distances, 63% and 54%; and left ventricular ejection fraction, 88% and 83%. At a population level, however, we noted statistically signiﬁcant differences in median levels of NT-proBNP, KCCQ score, and 6-minute walk distance (all P<0.001) but not left ventricular ejection fraction (P=0.76).

In addition, we assessed the overlap in distributions of variables that reﬂect the gold standard measurement for maximal functional capacity in HF—cardiopulmonary exercise testing—according to NYHA classiﬁcation in HF-ACTION. As shown in Figure 3, although there were statistically signiﬁcant differences in median levels of minute ventilation—carbon dioxide production relationship, maximal oxygen uptake, and exercise duration (P<0.001), we noted substantial overlap in
Discussion
This analysis of 4 landmark HF trials demonstrates that the NYHA system poorly differentiates patients across the spectrum of functional impairment. In this report, we examined both the macro- and microimplications of the NYHA classification system across the spectrum of HF and found that it is an unreliable predictor of survival and a poor discriminator of functional impairment in HF. A heterogeneity of risk is strikingly clear in similar NYHA classifications across studies from lower risk (eg, HF-ACTION) to higher risk (eg, GUIDE IT) and across trials including patients with HF reduced and preserved ejection fraction, implying that the prognostic value of NYHA classification is largely dependent on the baseline risk of the patient in which it is assessed. This suggestion is contrary to the general assumption that the NYHA classification is an accurate measure of mortality risk and is consistent across studies for patients of a similar class.

Whereas the heterogeneity of NYHA class across trials is a recognized consequence of differences in inclusion and exclusion criteria of the studies assessed and the heterogeneity of risk in these studies, use of NYHA symptom severity by regulatory bodies does not necessarily take this limitation into consideration. Once clinical trials are completed, therapies may be approved for specific NYHA classes and suggested based on post hoc analyses of the data. For example, the CardioMEMS HF System is presently approved for NYHA class III, and a clinical trial with expected enrollment of 3600 is ongoing to extend its approval to NYHA class II patients (NCT03387813). Our findings suggest it is time to revisit the use of NYHA class to guide enrollment into trials and approval for therapy on its basis.

Figure 1. Kaplan–Meier curves for all-cause mortality according to clinical trial and New York Heart Association (NYHA) classification. Clinical trials shown are TOPCAT (Treatment of Preserved Cardiac Function Heart Failure With an Aldosterone Antagonist), DIG (The Effect of Digoxin on Mortality and Morbidity in Patients With Heart Failure), HF-ACTION (Efficacy and Safety of Exercise Training in Patients With Chronic Heart Failure), and GUIDE-IT (Guiding Evidence-Based Therapy Using Biomarker Intensified Treatment in Heart Failure).
Other more objective and better calibrated measures of disease severity and patient-reported symptoms such as the KCCQ, the Minnesota Living with Heart Failure Questionnaire, and biomarkers might be better suited to guide enrollment strategies and to appraise the impact of therapeutic interventions on patient symptoms. Although the median levels of almost all objective HF parameters differed significantly between NYHA classes II and III, there was immense overlap in values. These findings, along with the longstanding recognition that the NYHA classification system has poor reproducibility, raise the question of whether our care of HF patients might be enhanced if we elevated the clinical use of disease descriptors that are more objective and precise. Furthermore, these limitations make its centrality to the HF guidelines potentially inconsistent with the goal of improving patient care.

Limitations

Several limitations should be considered. First, we did not include patients classified as NYHA class I or IV because they constituted a small minority in the HF trials we assessed. Second, detailed phenotyping of HF patients that included natriuretic peptide levels and KCCQ assessments was available for only 2 of the trials, and only 1 trial had data on cardiopulmonary exercise testing, the gold standard for measure of functional status in HF. Third, we did not have data on real-world use of NYHA classification, but it would not
be expected to change our conclusions in a meaningful manner. Fourth, we did not limit our analysis to HF patients with reduced ejection fraction trials for the survival analyses. Prior studies have shown that prognosis is similar in HF patients with reduced or preserved ejection fraction. Fifth, the trials covered a long period of time, and therapies for HF have improved; however, the "newest" clinical trial—GUIDE-IT—had patients who did the worst within similar categories of NYHA class, supporting our hypothesis.

Conclusions

The NYHA classification system is an unreliable predictor of adverse outcomes in HF and poorly discriminates among patients across the spectrum of functional impairment. Its continued usage in guidelines, clinical trials, for US Food and Drug Administration approval of therapies, and for clinical decision making may hinder our progress toward bringing precision medicine to the bedside of HF patients.

Disclosures

None.

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