Staging classification of aortic stenosis based on the extent of cardiac damage

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Received 25 April 2017; revised 23 May 2017; editorial decision 14 June 2017; accepted 18 June 2017; online publish-ahead-of-print 21 July 2017

Aims
In patients with aortic stenosis (AS), risk stratification for aortic valve replacement (AVR) relies mainly on valve-related factors, symptoms and co-morbidities. We sought to evaluate the prognostic impact of a newly-defined staging classification characterizing the extent of extravalvular (extra-aortic valve) cardiac damage among patients with severe AS undergoing AVR.

Methods and results
Patients with severe AS from the PARTNER 2 trials were pooled and classified according to the presence or absence of cardiac damage as detected by echocardiography prior to AVR: no extravalvular cardiac damage (Stage 0), left ventricular damage (Stage 1), left atrial or mitral valve damage (Stage 2), pulmonary vasculature or tricuspid valve damage (Stage 3), or right ventricular damage (Stage 4). One-year outcomes were compared using Kaplan–Meier techniques and multivariable Cox proportional hazards models were used to identify 1-year predictors of mortality. In 1661 patients with sufficient echocardiographic data to allow staging, 47 (2.8%) patients were classified as Stage 0, 212 (12.8%) as Stage 1, 844 (50.8%) as Stage 2, 413 (24.9%) as Stage 3, or 145 (8.7%) as Stage 4. One-year mortality was 4.4% in Stage 0, 9.2% in Stage 1, 14.4% in Stage 2, 21.3% in Stage 3, and 24.5% in Stage 4 (P trend < 0.0001). The extent of cardiac damage was independently associated with increased mortality after AVR (HR 1.46 per each increment in stage, 95% confidence interval 1.27–1.67, P < 0.0001).

Conclusion
This newly described staging classification objectively characterizes the extent of cardiac damage associated with AS and has important prognostic implications for clinical outcomes after AVR.

Keywords
Aortic stenosis • Aortic valve • Aortic valve replacement • Transcatheter aortic valve replacement • Transcatheter aortic valve implantation • Classification • Staging
Introduction

Current recommendations for aortic valve replacement (AVR) in patients presenting with aortic stenosis (AS) rely solely on the presence of two criteria: (i) The demonstration of a severe stenosis based on a grading system that comprises specific valvular criteria, including peak aortic velocity ($V_{\text{max}}$), mean transvalvular gradient, and aortic valve area or aortic valve area index (AVAI), and (ii) the presence or absence of symptoms (dyspnea, heart failure, angina, or syncope) related to AS. Additionally, risk stratification of patients for AVR is currently based on the presence of comorbidities. There are no recommendations regarding the importance of anatomical or functional cardiac consequences of AS as a component of the AVR decision algorithm, other than reduced left ventricular (LV) systolic function defined as a LV ejection fraction <50%, although the literature clearly documents that the presence of cardiac damage holds prognostic significance. In this report, from a large contemporary population of patients with symptomatic severe AS undergoing AVR (either surgery or transcatheter AVR), we sought to (i) better characterize the prevalence of concomitant cardiac damage at the time of AVR, (ii) categorize the different types of cardiac damage (LV dysfunction, left atrial enlargement, pulmonary hypertension, and right ventricular dysfunction) into distinct stages of cardiac disease involvement, and (iii) assess the impact of these stages on survival and the risk of adverse outcomes after AVR.

Methods

Study population

The design of the Placement of Aortic Transcatheter Valves (PARTNER) 2 trial has previously been described, including a detailed description of eligibility criteria and procedural methods. Data from the PARTNER 2A (N = 2032) and PARTNER 2B (N = 671) trials were pooled for this analysis. The PARTNER 2 Trials were approved by the institutional review board of each participating site and all patients provided written informed consent.

Definitions

Patients were categorized into five stages (independent, not additive) depending on the presence or absence of extravalvular (extra aortic valve) cardiac damage or dysfunction as detected by transthoracic echocardiography before AVR. Stage 0: No other cardiac damage detected; Stage 1: LV damage as defined by presence of LV hypertrophy (LV mass index >95 g/m² for women, >115 g/m² for men), severe LV diastolic dysfunction ($E'/e'_0 >1.4$), or LV systolic dysfunction (LV ejection fraction <50%); Stage 2: LA or mitral valve damage or dysfunction as defined by the presence of an enlarged left atrium (>34 mL/m²), the presence of atrial fibrillation, or the presence of moderate or severe mitral regurgitation; Stage 3: Pulmonary artery vasculature or tricuspid valve damage or dysfunction as defined by the presence of systolic pulmonary hypertension (systolic pulmonary arterial pressure >60 mmHg) or the presence of moderate or severe tricuspid regurgitation, and Stage 4: RV damage as defined by the presence of moderate or severe RV dysfunction (Figure 1). Patients were hierarchically classified in a given stage (worst stage) if at least one of the proposed criteria was met within that stage. These criteria were chosen based on their broad acceptance, prior validation as markers of abnormal cardiac function, their simplicity of acquisition, and their potential for future clinical external generalizability. The classification algorithm as well as the statistical models were defined fully a priori. Frailty was defined as the presence of at least two of the following criteria: (i) Katz index of independence in activities of daily living <6; (ii) 15-m walk time ≥24 s, (iii) serum albumin <3.8 g/dL and (iv) grip strength <13 kg (women) or <26 kg (men).

Transthoracic echocardiograms were obtained at baseline and follow-up using a uniform image acquisition protocol. All studies were analysed...
by a central core laboratory with quality and measurement methodology previously reported. All adverse events were adjudicated by an independent committee.

**Statistical analysis**

Continuous data are presented as mean ± standard deviation and were compared between groups using the Student’s t-test or the Wilcoxon rank sum test, as appropriate. Categorical variables were presented as count and per cent, and compared using the χ² or the Fisher exact test. We estimate time-to-event data using Kaplan–Meier techniques.

The unadjusted and adjusted risk of dying within 1 year according to stage of cardiac damage was assessed using Cox proportional hazards regression. Models were stratified by study (PARTNER 2A vs. PARTNER 2B) and treatment assignment (transcatheter vs. surgical), with the following variables included for multivariable adjustment: STS score, frailty, age, sex, O₂-dependent chronic obstructive pulmonary disease (O₂-COPD), coronary artery disease, renal insufficiency, prior myocardial infarction, prior coronary artery bypass grafting, diabetes, Vₘₐₓ, and AVAI. In separate models, we assessed whether the variables STS score, frailty, O₂-COPD, Vₘₐₓ, or AVAI moderated the effect of stage of cardiac damage on mortality by including interaction terms between these variables and the stage of cardiac damage in the Cox models. We also tested for statistical interaction between the staging variable and study or treatment assignment; these latter two models were not stratified by study or treatment assignment. To determine the incremental value of the staging system in regards to 1-year mortality predictability, we calculated the net reclassification index (NRI) and the integrated discriminatory index (IDI) for the stage variable by comparing Cox models including vs. excluding the staging variable in addition to other patient characteristics.

To account for missing variables, we conducted separate sensitivity analyses using multiple imputations and generated 40 imputed datasets. In addition to the individual components of the staging algorithm, the following variables were included in the imputation model: Age, STS score, renal insufficiency, coronary artery disease, previous coronary artery bypass grafting, O₂-COPD, Vₘₐₓ, or AVAI.

**Results**

**Study population**

From the PARTNER 2A (n = 1114) and PARTNER 2B (n = 547) trials, the primary study population comprised 1661 patients who had complete echocardiographic assessments at baseline, which allowed for patient staging (see Supplementary material online). A total of 1107 patients underwent transcatheter AVR, and 554 patients underwent surgical AVR. At the time of AVR, 47 (2.8%) patients were in Stage 0 (no cardiac damage), 212 (12.8%) patients were in Stage 1 (LV damage), 844 (50.8%) patients were in Stage 2 (LA or mitral valve damage), 413 (24.9%) patients were in Stage 3 (pulmonary vasculature or tricuspid valve damage), and 145 (8.7%) patients were in Stage 4 (RV damage) (Table 1). Baseline and procedural characteristics according to stage are presented in see Supplementary material online, Table S1. In general, patients in more advanced stages were older, were more often male, had higher body mass index, higher STS score and EuroSCORE, more often had diabetes, more often had previous myocardial infarction or coronary artery bypass grafting, and were more often frail and had O₂-COPD. Rates of each individual cardiac damage component within each stage are presented in see Supplementary material online, Table S2, and rates of each prior stages within each stages are presented in see Supplementary material online, Figure S2.

**Outcomes**

One-year outcomes after AVR stratified by stage of cardiac damage are presented in Table 2 and Figure 2. At 1 year, all-cause death (Figure 2A) and cardiac death (Figure 2B) significantly increased with each stage of worsening cardiac damage. Landmark analysis demonstrated consistent relationship between extent of cardiac damage prior to AVR and the occurrence of death or cardiac death (Figure 2C and D) beyond 30 days. Supplementary material online, Figure S3 presents 1-year adjusted survival curves for death and cardiac death, with consistent results. Supplementary material online, Figure S4 presents 1-year outcomes after excluding patients with prior coronary artery bypass grafting and O₂-COPD. Results of all sensitivity analyses were consistent with the main analysis.

**Predictors of 1-year death**

After multivariable analysis and when tested in multiple models, stage of cardiac damage was shown to be one of the strongest predictors of 1-year death, with an adjusted mortality hazard of ~1.45 with each increase in stage (Table 3). Frailty and O₂-COPD were the only other
identifiable predictors of 1-year death after AVR. Stage of cardiac damage had incremental value over other covariates for prediction of 1-year death after AVR, with NRI and IDI consistently showing improvement after the addition of the staging variable to several different nested models (see Supplementary material online, Table S3). In particular, the addition of stage of cardiac damage as a covariate was shown to reclassify a significant proportion of patients compared with STS (NRI = 0.149) and frailty (NRI = 0.126) variables.
in regard to 1-year mortality prediction. No interaction was detected between AVR strategy (surgical AVR vs. transcatheter AVR) ($P_{\text{interaction}} = 0.28$), STS ($P_{\text{interaction}} = 0.13$), frailty ($P_{\text{interaction}} = 0.98$), $O_2$-COPD ($P_{\text{interaction}} = 1.00$), AVAi ($P_{\text{interaction}} = 0.75$), or $V_{\text{max}}$ ($P_{\text{interaction}} = 0.87$) and stage of cardiac damage in regard to the occurrence of 1-year death.

### Sensitivity analysis
Baseline characteristics and outcomes of patients included and excluded in the present study are presented in Supplementary material online, Tables S4 and S5. Stage of cardiac damage showed a strong unadjusted and adjusted association with mortality after multiple imputation of missing data (see Supplementary material online, Table S6), consistent with the main analysis.

### Discussion
The current report, derived from 1661 patients who underwent comprehensive echocardiographic assessment before AVR, demonstrated a strong relationship between the extent of cardiac damage at baseline and 1-year survival after AVR. The proposed anatomic and functional cardiac damage staging system, was shown to be one of the strongest predictors of mortality. For each stage increment, 1-year mortality risk increased by ~45%.

### Table 3 One-year predictors of mortality among patients with severe aortic stenosis undergoing aortic valve replacement

| Variables                                           | Unadjusted hazard ratio (95% confidence interval) | $P$-value | Adjusted hazard ratio (95% confidence interval) | $P$-value |
|-----------------------------------------------------|---------------------------------------------------|-----------|------------------------------------------------|-----------|
| **Model 1**                                          |                                                   |           |                                                |           |
| Stage of cardiac damage (by each stage increase)     | 1.46 (1.27–1.67)                                  | <0.0001   | 1.41 (1.20–1.66)                               | <0.0001   |
| Society of Thoracic Surgeons score (by each 1% increase) | 1.05 (1.02–1.08)                                  |           | 1.03 (0.99–1.07)                               | 0.10      |
| Frailty                                             | 1.98 (1.48–2.64)                                  | <0.0001   | 1.83 (1.35–2.49)                               | <0.0001   |
| Age (by each 10 years increase)                      | 1.13 (0.95–1.35)                                  |           | 1.13 (0.89–1.43)                               | 0.32      |
| Male sex                                            | 1.14 (0.89–1.47)                                  |           | 1.25 (0.92–1.70)                               | 0.15      |
| $O_2$-chronic obstructive pulmonary disease          | 1.90 (1.31–2.76)                                  | <0.0001   | 1.99 (1.24–3.17)                               | 0.004     |
| Renal insufficiency                                  | 1.37 (0.90–2.08)                                  |           | 0.79 (0.46–1.35)                               | 0.39      |
| Coronary artery disease                              | 0.89 (0.69–1.15)                                  |           | 0.95 (0.69–1.29)                               | 0.73      |
| Previous coronary artery bypass graft                | 0.91 (0.68–1.22)                                  |           | 0.95 (0.65–1.37)                               | 0.77      |
| $V_{\text{max}}$ (by 0.5 m/s increase)               | 0.91 (0.82–1.01)                                  |           | 0.91 (0.80–1.04)                               | 0.17      |
| Aortic valve area index (by 0.1 cm$^2$ decrease)     | 0.96 (0.84–1.10)                                  |           | 0.99 (0.84–1.16)                               | 0.88      |
| **Model 2**                                          |                                                   |           |                                                |           |
| Stage of cardiac damage (by each stage increase)     | 1.46 (1.27–1.67)                                  | <0.0001   | 1.41 (1.20–1.65)                               | <0.0001   |
| STS (by each 1% increase)                            | 1.05 (1.02–1.08)                                  | <0.0001   | 1.03 (1.00–1.07)                               | 0.09      |
| Frailty                                             | 1.98 (1.48–2.64)                                  | <0.0001   | 1.81 (1.34–2.44)                               | <0.0001   |
| Age (by each 10 years increase)                      | 1.13 (0.95–1.35)                                  |           | 1.07 (0.84–1.36)                               | 0.60      |
| Male sex                                            | 1.14 (0.89–1.47)                                  |           | 1.23 (0.92–1.66)                               | 0.17      |
| $O_2$-chronic obstructive pulmonary disease          | 1.90 (1.31–2.76)                                  | 0.0008    | 2.01 (1.28–3.15)                               | 0.003     |
| Renal insufficiency                                  | 1.37 (0.90–2.08)                                  |           | 0.80 (0.47–1.36)                               | 0.41      |
| Previous myocardial infarction                       | 1.27 (0.94–1.71)                                  |           | 1.25 (0.89–1.77)                               | 0.20      |
| Previous coronary artery bypass graft                | 0.91 (0.68–1.22)                                  | 0.57      | 0.90 (0.63–1.30)                               | 0.58      |
| Diabetes                                            | 0.97 (0.74–1.26)                                  |           | 0.89 (0.65–1.21)                               | 0.46      |
| **Model 3**                                          |                                                   |           |                                                |           |
| Stage of cardiac damage (by each stage increase)     | 1.46 (1.27–1.67)                                  | <0.0001   | 1.44 (1.23–1.70)                               | <0.0001   |
| Frailty                                             | 1.98 (1.48–2.64)                                  | <0.0001   | 1.82 (1.34–2.47)                               | 0.0001    |
| STS (by each 1% increase)                            | 1.05 (1.02–1.08)                                  | <0.0002   | 1.03 (1.00–1.07)                               | 0.07      |
| Age (by each 10 years increase)                      | 1.13 (0.95–1.35)                                  |           | 1.10 (0.86–1.41)                               | 0.43      |
| Male sex                                            | 1.14 (0.89–1.47)                                  |           | 1.26 (0.93–1.71)                               | 0.13      |
| $O_2$-chronic obstructive pulmonary disease          | 1.90 (1.31–2.76)                                  | 0.0008    | 1.99 (1.25–3.17)                               | 0.004     |
| Renal insufficiency                                  | 1.37 (0.90–2.08)                                  |           | 0.77 (0.45–1.32)                               | 0.34      |
| Previous myocardial infarction                       | 1.27 (0.94–1.71)                                  | 0.11      | 1.32 (0.93–1.87)                               | 0.12      |
| Previous coronary artery bypass graft                | 0.91 (0.68–1.22)                                  | 0.55      | 0.89 (0.62–1.29)                               | 0.55      |
| Diabetes                                            | 0.97 (0.74–1.26)                                  |           | 0.91 (0.67–1.25)                               | 0.56      |
| Aortic valve area index (by 0.1 cm$^2$ decrease)     | 0.96 (0.84–1.10)                                  |           | 1.03 (0.89–1.19)                               | 0.68      |
Our report also demonstrated that this new staging system had significant incremental value for prediction of 1-year survival over several well-established predictors of worse outcomes after AVR, including patient frailty and the STS score.

Current guidelines recommend risk stratification of patients with AS using the integration of different variables including the severity of AS, the presence or absence of AS-related symptoms, and the presence of other risk factors such as STS score, frailty, or the compromise of other major organ systems (e.g., kidney disease, lung disease)\(^1\)\(^2\), however, no clear recommendation exists on how to incorporate the extent of consequential (or associated) cardiac damage in clinical decision making related to AS. Given the strong association demonstrated in this study between advanced staging of cardiac damage and worse clinical outcomes after AVR, consideration of the stage of AS-related cardiac damage in future recommendations for risk stratification might be useful.

Notably, we demonstrated that the extent of cardiac damage remains one of the strongest independent predictors of 1-year mortality post-AVR after controlling for important prognostic factors such as STS score and the presence of frailty, coronary artery disease, renal disease, or O\(_2\)-COPD. After adjustment, only stage of cardiac damage, frailty, and O\(_2\)-COPD remain predictors of 1-year mortality. This finding is important and may have identified meaningful variables that could be easily incorporated in a simple risk-prediction model beyond the standard STS. In the current report, a high proportion of patients enrolled in the two-pooled studies were at particularly high risk, with multiple comorbidities. Therefore, it is expected that in a lower-risk population (i.e., with minimal comorbidities), the extent of cardiac damage at baseline could play an even greater role in risk prediction. Similarly, extent of cardiac damage was a considerably stronger predictor of adverse outcomes after AVR than both \(V_{\text{max}}\) and AVAI, underscoring the fact that the valvular haemodynamic burden is effectively corrected by AVR whereas the detrimental impact of extravalvular consequences of AS often persists after AVR.

It is important to note that patients classified with more advanced stages also demonstrated higher non-cardiac mortality. This finding is not surprising since patients with more advanced cardiac disease are known to be more vulnerable to any other new insult (such as infection, bleeding, trauma, etc.), and to have a decreased physiological reserve to fight any additional disease process. That being said, identifying the exact cause of death among elderly patients with multiple co-morbidities could be challenging, with often an initial event triggering a cascade of consequences, ultimately leading to death.

Interestingly, the natural evolution or ‘propagation’ of AS associated cardiac damage doesn’t seem to occur systematically in a sequential fashion. Indeed, among patients with more advanced stages (3 or 4), lower than expected proportion of patients cumulated damage from earlier stages (see Supplementary material online, Figure S2). For instance, among the 145 patients presenting with moderate to severe RV failure, only 25% presented with severe pulmonary hypertension, and ~75% with left atrial enlargement (see Supplementary material online, Table S2). These finding are intriguing and may suggest that the extension of cardiac damages related to a LV pressure overload such as AS may not always be sequential (LV, LA, pulmonary vasculature, and then RV dysfunction), and may vary based on patient susceptibility or genetic predisposition.\(^14\)-\(^16\) In this regard, recent studies demonstrated that the occurrence of LV hypertrophy or LV dysfunction may impact RV function early, either by ventricular interdependence, by contiguous extension of the pathological response to pressure overload, or by systemic hormonal response to left ventricle overload or genetic predisposition to hypertrophy affecting both ventricles.\(^17\)-\(^19\) Further prospective mechanistic investigations related to the natural evolution of cardiac damages among patients with AS are needed to better characterize these hypothesis-generating findings.

### Strengths and limitations

The current report has several strengths. It was derived from a substantial number of patients who underwent comprehensive echocardiographic analysis before AVR evaluated by an independent echocardiographic core laboratory, and it describes for the first time a simple and intuitive physiologic classification, reflecting the natural evolution of AS and bearing prognostic implications. Additionally, outside the current report, the proposed classification may have potential clinical and research utility: (i) It may improve clinical risk stratification of patients prior to AVR; (ii) it provides a tool for clinician to better communicate to patients risks, benefits, and expected prognostic after AVR; (iii) by acknowledging some of the technical challenges, variability, and discordances in echocardiographic acquisition of currently recommended severity grading criteria,\(^20\)-\(^29\) the proposed multi-parametric stratification system may synergistically help to better define the optimal timing of AVR by focusing on the sequences and mechanical repercussion of AS (Figure 3); and (iv) it may represent a standardized tool to better describe, stratify, and quantify cardiac damage during a randomized trial or other research work, either at baseline or following diverse treatment or therapeutic strategies.

The current report has many limitations. First, the described staging classification was retrospectively and not prospectively studied; future ongoing trials will be able to prospectively confirm the prognostic value of this new classification scheme (PARTNER 3 [NCT02675114], TAVR UNLOAD [NCT02661451], PROGRESSA [NCT1679431], EARLY TAVR NCT03042104). Second, a substantial amount of echocardiographic data were missing, leading to the exclusion of a high proportion of patients. That being said, our analysis is by far the largest cohort of patients with core laboratory adjudicated echocardiogram paired with independent events adjudication, making this manuscript unique. Third, the current staging system infers a direct causal role between the presence of AS and the detected cardiac damage. While concomitant comorbidities and diseases (e.g., severe coronary artery disease, severe lung disease) may co-exist and the detected cardiac damage may not be completely due to the AS, given the strong association with mortality, one could argue that this cardiac-oriented classification still has value when applied in a real-world setting. Indeed, patients with severe AS and several concomitant diseases (e.g., chronic obstructive pulmonary disease, ischaemic cardiomyopathy) are much more vulnerable to any new ‘insult’ (e.g., pneumonia), and any potential deterioration or extent of the already existent cardiac damage may not be well tolerated, leading to poor outcomes, even after successful AVR. However, after excluding patients with severe lung disease and prior coronary artery bypass grafting (severe CAD), our results were
unchanged (see Supplementary material online, Figure S3), supporting the value of this classification among all patients with AS. Fourth, more detailed and granular detection of cardiac damage involving different imaging modalities exist. These include detection of reduced LV strain,30–33 LA strain,34 or RV strain35–37 by speckle tracking echocardiography and the presence of myocardial fibrosis by magnetic resonance imaging.38 When available, these findings could be incorporated into the appropriate anatomical staging level (i.e. reduced LV strain in Stage 1, reduced LA strain in Stage 2), resulting in the expansion and improvement of the proposed staging system. Similarly, serum biomarkers, with specific capability to identify LV, LA, or RV overload, could also eventually complement this classification.39–42 Fifth, outcomes were available up to 1 year only; however, longer-term follow-up (up to 5 years) would have most likely amplified the observed difference in mortality between each level of cardiac damage. Finally, the concept behind the proposed staging system could be adapted and applied to other valvular disease.

In conclusion, this new staging classification characterizes the extent of anatomical and functional cardiac damage associated with AS prior to AVR and has prognostic implications post-AVR. Further studies are needed to prospectively validate this classification across different AS severities and to better define how it could be integrated to existing grading severity system in guiding AVR timing for patients with AS.

Supplementary material

Supplementary material is available at European Heart Journal online.

### Funding

The PARTNER 2 Trial was funded by Edwards Lifesciences.

### Conflict of interest

P.G. has received consultant fees and speaker fees from Edwards Lifesciences. P.P. holds the Canada research Chair in Valvular Heart Diseases, Canadian Institutes of Health research, Ottawa, ON, Canada, has Core Lab contracts with Edwards Lifesciences for which he receives no direct compensation, and is a consultant for St Jude Medical. M.J.M. is a member of the PARTNER Trial Executive Committee for which he receives no direct compensation. R.R.M. has received grants from Edwards Lifesciences and St Jude Medical, is a consultant for Abbott Vascular, Cordis, and Medtronic, and holds equity in Entourage Medical. W.A.J. has Core Lab contracts with Edwards Lifesciences for which he receives no direct compensation. L.G.S. holds equity in Cardiosolution and Valvexchange, intellectual property with Postthorax, and is a member of the PARTNER Trial Executive Committee for which he receives no direct compensation. E.M.T. is a member of the PARTNER Trial Executive Committee for which he receives no direct compensation. V.H.T. is a consultant for Edwards Lifesciences, Abbott, Sorin Medical, St Jude Medical, and DirectFlow. V.B. is a consultant for Edwards Lifesciences and Abbott Vascular. H.C.H. has received grants from Edwards Lifesciences, St Jude Medical, Medtronic, Boston Scientific, Abbott Vascular, Gore, Siemens, Cardiokinetics, and Mitraspan, is a consultant for Edwards Lifesciences and Siemens, and holds equity in Microinterventional Devices. W.Y.S. is a consultant for Microinterventional Devices. D.J.C. has received research support from Edwards Lifesciences, Medtronic, and Boston Scientific and is a consultant for Edwards Lifesciences and Medtronic. B.R.L.
was supported by K23 HL116660, serves on the advisory board of Roche Diagnostics, and has received grants from Edwards Lifesciences and Roche Diagnostics. M.C.A. is a consultant for Claret Medical. P.D.S. has Core Lab contracts with Edwards Lifesciences for which she receives no direct compensation. R.T.H. has Core Lab contracts with Edwards Lifesciences for which she receives no direct compensation and is a consultant for Philips Healthcare, St Jude Medical and Boston Scientific. S.K.K. has Core Lab contracts with Edwards Lifesciences and holds equity in Thubrikar Aortic Valve, Inc. C.R.S. is a member of the PARTNER Trial Executive Committee for which he receives no direct compensation. D.C.M. is supported by research grant R01 NHLBI #HL67025, has received consulting fees from Abbott Vascular, St Jude Medical, and Medtronic, and is a member of the PARTNER Trial Executive Committee for which he receives no direct compensation. J.G.W. is a consultant for Edwards Lifesciences and a member of the PARTNER Trial Executive Committee, for which he receives no direct compensation. M.B.L. is a member of the PARTNER Trial Executive Committee for which he receives no direct compensation. The other authors report no relevant relationships with industry to disclose.

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