Prevalence of Regional Myocardial Thinning and Relationship With Myocardial Scarring in Patients With Coronary Artery Disease

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Regional myocardial wall thinning is believed to represent chronic myocardial infarction. A recent task force report on the definition of myocardial infarction by the European Society of Cardiology, American College of Cardiology, American Heart Association, and the World Heart Federation concludes that “[i]maging evidence of a region... that is thinned and fails to contract, in the absence of a non-ischaemic cause” is diagnostic for chronic myocardial infarction. Accordingly, many experts state that viability testing is unnecessary for regions with wall thinning, because this is synonymous with scar tissue.2-6

However, recent case reports incorporating the use of delayed-enhancement cardiovascular magnetic resonance (CMR) imaging raise the possibility that this viewpoint is incorrect.7,8 These single-patient reports indicate that myocardial regions with severe wall thinning do not necessarily consist entirely of scar tissue but instead may have minimal or no scar.

Importance Regional left ventricular (LV) wall thinning is believed to represent chronic transmural myocardial infarction and scar tissue. However, recent case reports using delayed-enhancement cardiovascular magnetic resonance (CMR) imaging raise the possibility that thinning may occur with little or no scarring.

Objective To evaluate patients with regional myocardial wall thinning and to determine scar burden and potential for functional improvement.

Design, Setting, and Patients Investigator-initiated, prospective, 3-center study conducted from August 2000 through January 2008 in 3 parts to determine (1) in patients with known coronary artery disease (CAD) undergoing CMR viability assessment, the prevalence of regional wall thinning (end-diastolic wall thickness ≤5.5 mm), (2) in patients with thinning, the presence and extent of scar burden, and (3) in patients with thinning undergoing coronary revascularization, any changes in myocardial morphology and contractility.

Main Outcomes and Measures Scar burden in thinned regions assessed using delayed-enhancement CMR and changes in myocardial morphology and function assessed using cine-CMR after revascularization.

Results Of 1055 consecutive patients with CAD screened, 201 (19% [95% CI, 17% to 21%]) had regional wall thinning. Wall thinning spanned a mean of 34% (95% CI, 32% to 37% [SD, 15%]) of LV surface area. Within these regions, the extent of scarring was 72% (95% CI, 69% to 76% [SD, 25%]); however, 18% (95% CI, 13% to 24%) of thinned regions had limited scar burden (≤50% of total extent). Among patients with thinning undergoing revascularization and follow-up cine-CMR (n=42), scar extent within the thinned region was inversely related to regional (r = −0.72, P < .001) and global (r = −0.53, P < .001) contractile improvement. End-diastolic wall thickness in thinned regions with limited scar burden increased from 4.4 mm (95% CI, 4.1 to 4.7) to 7.5 mm (95% CI, 6.9 to 8.1) after revascularization (P < .001), resulting in resolution of wall thinning. On multivariable analysis, scar extent had the strongest association with contractile improvement (slope coefficient, −0.03 [95% CI, −0.04 to −0.02]; P < .001) and reversal of thinning (slope coefficient, −0.05 [95% CI, −0.06 to −0.04]; P < .001).

Conclusions and Relevance Among patients with CAD referred for CMR and found to have regional wall thinning, limited scar burden was present in 18% and was associated with improved contractility and resolution of wall thinning after revascularization. These findings, which are not consistent with common assumptions, warrant further investigation.

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METHODS
Patients and Protocol
This prospective study, conducted from August 2000 through January 2008, consisted of 3 parts (FIGURE 1). Patients were enrolled at 3 sites: Duke Cardiovascular Magnetic Resonance Center, Durham, North Carolina; The Heart Group PLLC, Nashville, Tennessee; and Northwestern University, Chicago, Illinois. All patients provided written informed consent for this protocol, which was approved by the institutional review board at each site.

Part A: Prevalence of Thinning. The purpose of part A was to evaluate the prevalence of myocardial thinning. Consecutive patients with CAD who were scheduled for CMR viability assessment were screened and asked to participate unless they (1) were younger than 18 years, (2) had experienced an acute myocardial infarction (MI) within last 2 weeks, (3) had nonischemic myocardial disease (eg, hypertrophic cardiomyopathy, myocarditis), or (4) were unable to provide consent. A total of 1134 patients were approached, of whom 56 (5%) declined to participate, 12 (1%) were unable to fit into the scanner, and 11 (1%) did not complete CMR assessment for various reasons (eg, claustrophobia, inability to lie flat). One thousand fifty-five patients completed CMR and were included in the analysis. Similar to prior publications, myocardial thinning was defined as an end-diastolic wall thickness (EDWT) of 5.5 mm or less on cine-CMR.9

Part B: Scar Burden in Thinned Regions. The purpose of part B was to evaluate the prevalence of limited scar burden (defined as ≤50% of total extent) in thinned myocardium. This prespecified cutoff was chosen based on prior delayed-enhancement CMR studies demonstrating that regions with scar burden greater than 50% have low likelihood of functional recovery after revascularization.9,10 However, data were also evaluated using scar extent as a continuous variable.

Part C: Relationship of Scar Burden to Functional Improvement and Tissue Remodeling. The objective of part C was to determine the relationship of scar burden to functional improvement and myocardial tissue remodeling in the thinned region following coronary revascularization. The decision to undergo revascularization was guided by all relevant clinical and imaging data and made at the discretion of the treating physician and the patient.

Cardiovascular Magnetic Resonance
Image Acquisition. Images were acquired on 1.5T scanners (Siemens Sonata/Avanto) as described previously.9 Briefly, cine-CMR images were acquired using a steady-state free-precession sequence. Short-axis views were obtained every 1 cm to cover the entire left ventricle (6 mm thickness, 4 mm gap). Gadolinium contrast (gadopentetate dimeglumine, gadoteridol, or gadoversetamide) was administered intravenously (0.15 mmol/kg), and delayed-enhancement CMR images were acquired 10 to 15 minutes later using a segmented inversion-recovery sequence in the same views used for cine-CMR.

Image Analysis. Cardiovascular magnetic resonance images were evaluated masked to patient identity and clinical information. For patients who underwent revascularization and returned for follow-up cine-CMR, the follow-up scan was analyzed by an observer blinded to all delayed-enhancement data and different from the individual who analyzed the baseline images. Quantitative measurements were made by manual planimetry.

Determination of Wall Thinning. Regional thinning was defined quantitatively as the sector in which EDWT was 5.5 mm or less on the end-diastolic cine frame for each short-axis slice (FIGURE 2A). The global left ventricular (LV) extent of thinning was calculated...
Figure 2. Cardiovascular Magnetic Resonance (CMR) Analyses

A Cine-CMR analysis of regional left ventricular (LV) end-diastolic wall thickness (EDWT)

Figure 2A. Regional thinning was defined as the sector in which LV end-diastolic wall thickness (EDWT) was ≤5.5 mm on the end-diastolic cine frame for each short axis slice. RV indicates right ventricle.

B Cine-CMR analysis of global LV extent of thinning

Figure 2B. A patient was considered to have LV thinning if the global extent of thinning was >5% of total LV circumference (ie, LV surface area).

C Delayed-enhancement CMR analysis of regional extent of scarring

Figure 2C. Patients with ≤50% scarring in the thinned region were classified as having limited scar burden and those with >50% were classified as having extensive scar burden.
by summing the mid-myocardial circumference of the thinned region and dividing by the mid-myocardial circumference of the entire left ventricle (summation of all short-axis cine slices), multiplied by 100 (Figure 2B). On a per-patient basis, thinning was deemed present if the global extent of thinning was more than 5% of total LV circumference (ie, LV surface area).

**Scar Extent in Thinned Regions.** Cine images were used to determine the representative view that went through the central core of the thinned region. Quantitative analysis was then performed on the corresponding delayed-enhancement CMR slice after 4-fold interpolation. The extent of scarring (hyperenhanced region) in the thinned region was expressed as a percentage of the total area of thinned myocardium (Figure 2C). Patients with 50% or less scarring in the thinned region were classified as having limited scar burden and those with more than 50% scarring as having extensive scar burden.

**LV Volumes, Ejection Fraction, and Mass.** Endocardial borders on the stack of short-axis cine images in end diastole and end systole, and the epicardial borders in end diastole, were traced using planimetry. Left ventricular ejection fraction (LVEF) was calculated by subtracting end-systolic volume from end-diastolic volume and dividing by end-diastolic volume. Left ventricular mass was calculated by subtracting endocardial from epicardial volumes and multiplying by myocardial density (1.05 g/cm³).11

**Regional Functional Improvement and Myocardial Tissue Remodeling.** For patients who returned for follow-up CMR after revascularization, analyses of end-diastolic and end-systolic wall thickness were performed in the same manner as for baseline measurements. The thinned region identified on the prerevascularization images was registered to the same region on the postrevascularization images using conventional landmarks (right ventricular insertion sites and papillary muscles).

**Intraobserver and Interobserver Agreement.** Images in a random sample of 10% of the population in study part B (21 patients) were reanalyzed 6 months later to determine intraobserver and interobserver agreement for measurements of EDWT, systolic wall thickening, and transmural extent of scarring. Intraobserver and interobserver agreement values were similar, with a bias of less than 0.2 mm for EDWT and systolic wall thickening and less than 1% for extent of scarring within the thinned region, with an SD of the difference of less than 0.3 mm for EDWT and systolic wall thickening and less than 4.7% for extent of scarring.

**Coronary Angiography**

Angiograms were reviewed in conjunction with the cine-CMR images to identify the coronary artery supplying the thinned region. The maximum severity of stenosis was analyzed using standard quantitative coronary angiography methods.12 The extent of collateral flow was determined using the Rentrop classification scheme (0, no collateral flow; 1, collateral flow leading to side branches; 2, collateral flow leading to side branches and a portion of the epicardial segment; 3, collateral flow leading to side branches and the entire epicardial segment).13

**Electrocardiography**

The Minnesota Code, a standardized classification system for electrocardiograms, was used to determine the presence of prior Q-wave myocardial infarction (codes 1-1-1 to 1-2-7) based on Q-wave location, amplitude, and duration.14,15 Infarct size as determined by electrocardiogram was estimated using the 54-criteria, 32-point Selvester QRS scoring system.16

**Statistical Analysis**

Continuous data are presented as mean (SD). Two-sample t tests were used to compare continuous data between 2 groups. Comparisons between discrete data were made using χ² tests; the Fisher exact test was used when cell count was less than 5. We used linear regression analyses to examine the relationships between scar burden and functional and morphological parameters at baseline and changes in these parameters with revascularization. Paired t tests were used to compare regional systolic wall thickening and EDWT before and after revascularization in subgroups with limited scarring and with extensive scarring. Multivariable logistic or linear regression analyses (as appropriate) were performed to identify clinical and imaging characteristics associated with limited scarring, functional improvement, and tissue remodeling. Variables with P < .05 on bivariable analysis were considered candidate variables in the multivariable models unless there were fewer than 3 total candidate variables, in which case the 3 most significant variables were included.

All statistical tests were 2-tailed; P < .05 was considered statistically significant. SAS version 9.2 (SAS Institute Inc) was used to perform the statistical analyses.

**RESULTS**

**Part A: Prevalence of Thinning**

Among the 1055 patients screened and who provided consent, 201 (19% [95% CI, 17% to 21%]) had myocardial wall thinning. The mean circumferential extent of thinning was substantial, encompassing 34% (95% CI, 32% to 37% [SD, 15%]) of total LV surface area. Baseline characteristics are shown in the TABLE. Patients with thinning were predominantly men (79%), had CAD risk factors (mean, 2.3 [SD, 1.0]) and significant LV dysfunction (LVEF, 32.6% [95% CI, 31.0% to 34.2%]; SD, 11.5%), and most had history of MI (71%) and electrocardiographic Q waves (67%).

**Part B: Scar Burden in Thinned Regions**

Of the 201 patients with wall thinning, 198 had scarring within the thinned region. The mean extent of scarring was 72% (95% CI, 69% to 76% [SD, 25%]); however, 18% (95% CI, 13% to 24%) of patients had scarring involving less than 50% of the thinned region (FIGURE 3A). Although there was an association between wall thickness and scar extent, the relationship was poor, because the
extent of scarring was highly variable for each level of wall thickness ($r = -0.22, P = .001$) (Figure 3B).

Also shown in the Table are the characteristics of patients with 50% or less scarring in the thinned region (limited scarring), compared with those with more than 50% (extensive scarring). Characteristics were similar be-

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<td>End-systolic wall thickness</td>
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<td>Systolic wall thickening</td>
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Abbreviations: ACE, angiotensin-converting enzyme; CAD, coronary artery disease; HMG, 3-hydroxy-3-methyl-glutaryl; LV, left ventricular; New York Heart Association.

$^a$Based on 50% or less scarring in the thinned region.

$^b$Available in 195 patients.

$^c$Infarct size as determined by electrocardiogram was estimated using the 32-point Selvester QRS scoring system, in which each point is designed to represent approximately 3% of LV mass.

$^d$Available in 199 patients.
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Figure 3. Correlation Between End-Diastolic Wall Thickness and Scar Burden Within the Thinned Region

A, The mean extent of scarring was 72%. Eighteen percent of patients had scarring involving 50% or less of the thinned region (percentages of patients with scarring were 5% in the region with 0%-25% scarring, 13% in the region with 26%-50% scarring, 27% in the region with 51%-75% scarring, and 54% in the region with 76%-100% scarring). N=201. B, Data were fit using linear regression. N=201.

Bivariate and Multivariable Analysis. eTable 2 reports the results of bivariate and multivariable regression analysis regarding associations with limited scarring in the thinned region who experienced significant improvement in both regional and global function after revascularization.

MYocardial Remodeling. As shown in Figure 5A, there was an inverse relationship between the extent of scarring in the thinned region and increase in EDWT after revascularization (r = -0.84, P < .001). Before revascularization, wall thinning was similar in patients with limited and extensive scarring (4.4 mm [95% CI, 4.1 to 4.7] vs 4.5 mm [95% CI, 4.2 to 4.7], P = .80). After revascularization, the group with limited scarring uniformly demonstrated an increase in EDWT, with a mean change of 3.1 mm (SD, 1.0 mm; range, 1.1-4.9 mm) (P < .001), whereas patients with extensive scarring demonstrated no change in EDWT (Figure 5B).

To ascertain if the increase in EDWT was attributable to active myocardial growth or simply a passive change in LV geometry, we related change in EDWT with change in global LV mass and change in LV end-diastolic volume. Figure 5C demonstrates that after revascularization there was no relationship between change in EDWT and change in LV mass (r = 0.07, P = .67). Conversely, an increase in EDWT after revascularization was associated with a reduction in LV end-diastolic volume (r = -0.42, P = .007) (Figure 5D).

The patient example in Figure 6 demonstrates that limited scarring in the thinned region was associated with disappearance of wall thinning after revascularization. There was also disappearance of Q waves after revascularization. Twenty-two patients had Q waves corresponding to the thinned region before revascularization; among these, the subgroup with reversal of wall thinning after revascularization was more likely to have disappearance of Q waves compared with the subgroup without reversal of wall thinning (4/5 [80%] vs 0/17 [0%]; P < .001).
functional and remodeling changes after revascularization. On multivariable analysis, scar extent had the strongest association with regional contractile improvement (slope coefficient, \(-0.03\) [95% CI, \(-0.04\) to \(-0.02\); \(P<.001\)), global contractile improvement (slope coefficient, \(-0.21\) [95% CI, \(-0.32\) to \(-0.11\); \(P<.001\)), and reversal of thinning (slope coefficient, \(-0.05\) [95% CI, \(-0.06\) to \(-0.04\); \(P<.001\)). The adjusted-model \(R^2\) values representing the proportion of variance explained for the linear regression models of contractile improvement and remodeling are reported in eTable 2.

**COMMENT**

The pathophysiology of regional wall thinning following transmural MI has undergone considerable investigation. Animal models have demonstrated that slippage of myocyte bundles plays a fundamental role in infarct expansion, remodeling, and ultimately wall thinning.\(^{17,18}\) Resorption during infarct healing, with removal of necrotic debris and resolution of tissue edema, may also play a role.\(^{19}\) With either mechanism, only complete rather than partial infarction of the myocardium at risk is postulated to result in substantial wall thinning.\(^{20-23}\) These experimental studies form the physiological basis for the view that regional wall thinning is indicative of transmural MI and absent residual viability.\(^{17,20-24}\) This concept has been perpetuated, despite a paucity of experimental data concerning the effects of chronic ischemia without infarction (ie, hibernating myocardium) on diastolic wall thickness.

Studies in patients have reinforced the viewpoint that regional wall thinning is indicative of transmural MI and have focused on the presence of focal thinning observable on conventional cardiac imaging as a straightforward, accurate marker of nonviable scar tissue.\(^{3-6}\) Baer et al\(^{7}\) and Schmidt et al\(^{8}\) (using cine-MRI) and Cwaig et al\(^{9}\) and La Canna et al\(^{10}\) (using echocardiography) concluded that a simple measurement of EDWT (<5-6 mm) indicates irreversible myocardial damage and thus obviates the need for viability testing. These investigations were small (43, 40, 45, and 28 patients, respectively), and only a fraction of patients or segments in each cohort had wall thinning. Nonetheless, these reports have become the basis for proposed clinical algorithms in which an early finding of wall thinning (<5-6 mm) precludes further viability testing or coronary revascularization, because thinned regions are assumed to represent permanent scar tissue without residual viability.\(^{3}\)

To our knowledge, the current study is the first to systematically investigate a cohort of patients with regional wall thinning. Of 201 patients identified by CMR as having wall thinning, most had significant LV dysfunction (LVEF, 32.6% [SD, 11.5%]), multivessel CAD, and thinning of a substantial portion of the left ventricle (34.2% [SD, 15.3%] of LV surface area). Among this cohort, 18% (95% CI, 13% to 24%) of thinned regions had limited or no scarring observed using delayed-enhancement CMR. Because the lack of scarring was associated with significant contractile improvement and reverse remodeling with resolution of wall thinning following revascularization, we believe the data indicate that myocardial thinning is potentially reversible and therefore should not be considered a permanent state.

Our results suggest that common clinical characteristics will not be useful in predicting whether thinned regions have limited scar tissue. There were no differences in age, sex, cardiac risk factors, angina or heart failure symptoms, or presence of Q waves between patients with extensive or limited scar burden. Although patients with extensive scarring were less likely to have collateral flow observable on coronary angiography, it is notable that 41% of those with limited scarring had no detectable angiographic collateral flow. The findings suggest that these clinical characteristics should not be
used to assess viability in a region of thinning.

As a group, patients with limited scarring had a very high degree of stenosis in the coronary artery perfusing the thinned region (mean stenosis, 95%). This suggests that lesser grades of stenosis may be sufficient to result in resting dysfunction without infarction but are insufficient to result in the unique condition of resting dysfunction and thinning without infarction. Although speculative, this suggests that thinning of viable regions requires a significant reduction in resting blood flow and represents an extreme form of hibernating myocardium with a delicate balance between resting ischemia and infarction. In this situation, one might expect reduced activity on myocardial perfusion imaging and an exhausted coronary flow reserve with an inability to respond to inotropic stimulation, despite limited or no scar tissue within the thinned region.

Independent of functional changes, it has been suggested that preservation of LV geometry may be an important end point after revascularization, because ongoing LV remodeling is a major determinant of poor prognosis.25-27 Indeed, a few studies have reported that "reverse" remodeling may occur following coronary revascularization, particularly in patients with substantial myocardial viability.26-27 The results of the current study extend previous observations and document a greater range in the progression and regression of LV remodeling as well as a different type of remodeling process. Whereas prior reports focused on global changes in the LV cavity—measures of volumes, sphericity, or both—we have shown that the myocardial wall may thin and revert back to full thickness as long as limited scarring is present. These results indicate that the end stage of remodeling is better determined by tissue composition (ie, scarring) rather than any set level of morphologic changes to the LV cavity or LV wall.

A possible mechanism for the disappearance of wall thinning is suggested by the documentation that reverse remodeling following revascularization was not accompanied by any change in total LV myocardial mass (Figure 5C). We hypothesize that slippage of myocyte bundles,17 the primary mechanism of myocardial infarct expansion, may also occur in the setting of chronic ischemia without infarction and that this process is reversible following coronary revascularization. This hypothesis requires further investigation.

Our study has a number of limitations. First, CMR was performed in patients in whom revascularization was being considered rather than in a randomized fashion. Hence, the results may be subject to selection bias, and the prevalence of wall thinning associated with limited scarring reported in our study population may be different from that found more generally in patients with stable chronic CAD or in those in whom revascularization is not indicated.

Second, follow-up CMR was performed in a limited number of patients and in none who did not undergo revascularization. Thus, this is another potential source of selection bias, and there remains some uncertainty regarding the conditions and the degree to which thinned regions can recover function. For instance, although we have previously shown that medical therapy can favorably influence LVEF and cavity dimensions in patients with chronic heart failure,28 it remains unknown if optimal medical therapy can result in disappearance of wall thinning.

Third, outcome data on our patient population are not available. Accordingly, we are unable to comment on whether the disappearance of wall thinning in association with recovery of
contractility leads to an improvement in prognosis.
Nevertheless, despite these limitations, we believe our study provides new insights into the pathophysiology of thinned myocardium and more broadly the process of reversible ischemic injury. The data show that thinned myocardium may consist of limited scar tissue and can recover function—concepts that are both inconsistent with current views. Moreover, we have observed that limited scar burden identified by delayed-enhancement CMR prior to revascularization is strongly as-

Figure 6. Cardiovascular Magnetic Resonance (CMR) Imaging and Electrocardiographic Changes in an Example Patient with Wall Thinning and Limited Scar Burden

A, Before revascularization, cine-CMR still frames in systole and diastole demonstrate akinesis and thinning of the anteroseptal, anterior, and apical walls. Delayed-enhancement images demonstrate limited scar burden (≤50%) within the thinned region. B, The electrocardiogram (ECG) demonstrates QS complexes in leads V1 through V3, with poor R-wave progression. C, After revascularization, cine-CMR still frames demonstrate improvement in myocardial contractility along with reversal of thinning in the previously thinned region. End-diastolic wall thickness changed from 4.5 to 9.5 mm after revascularization. D, The ECG, following revascularization, demonstrates presence of r waves in leads V1 through V3, that were not previously present. Full-motion cine sequences can be viewed at http://www.jama.com.
associated with contractile improvement and reversal of wall thinning after revascularization. This is relevant because the viability substudy of the STICH (Surgical Treatment in Ischemic Cardiomyopathy) trial recently reported that conventional viability testing with single-photon emission computed tomography or dobutamine echocardiography cannot help decide who should undergo revascularization or even identify who has improved survival (after adjusting for baseline variables and risk factors). 25 In concordance with the STICH substudy, our findings highlight that the pathophysiology of hibernating myocardium is still incompletely understood and that there is much to improve regarding the assessment of viability. The findings provide rationale for future experimental studies on reversible ischemic injury as well as for clinical studies prospectively testing whether CMR guidance for coronary revascularization decisions can improve patient outcome.

Author Contributions: Drs Shah and R. J. Kim had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

Study concept and design: Shah, H. W. Kim, James, Bonow, R. J. Kim.

Acquisition of data: Shah, James, Wu, Judd, R. J. Kim.

Analysis and interpretation of data: Shah, H. W. Kim, James, Parker, Bonow, Judd, R. J. Kim.

Drafting of the manuscript: Shah, H. W. Kim, Parker, Bonow, R. J. Kim.

Critical revision of the manuscript for important intellectual content: Shah, H. W. Kim, James, Wu, Bonow, Judd, R. J. Kim.

Statistical analysis: Shah, H. W. Kim, Parker, Judd.

Obtained funding: Judd, R. J. Kim.

Administrative, technical, or material support: Shah, James, Bonow, Judd, R. J. Kim.

Study supervision: Shah, Wu, Judd, R. J. Kim.

Conflict of Interest Disclosures: All authors have completed and submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Drs R. J. Kim and Judd reported that they are inventors named on a US patent, owned by Northwestern University, for delayed-enhancement cardiovascular magnetic resonance imaging. Dr Shah reported serving as a consultant to Astellas Pharmaceuticals; receiving grants pending from Siemens Medical Solutions, Astellas Pharmaceuticals, and Methodist Hospital Research Institute; and receiving payment for lectures from AstraZeneca, Lantheus Medical Imaging, and Takeda. No other authors reported disclosures.

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Online-Only Material: Tables 1 and 2 and cine sequences are available at http://www.jama.com.

REFERENCES


