Imaging Challenges in Secondary Mitral Regurgitation
Unsolved Issues and Perspectives

Patrizio Lancellotti, MD, PhD; Jose-Luis Zamorano, MD, PhD; Mani A. Vannan, MBBS

Chronic secondary mitral regurgitation (SMR) is a complex entity that is often clinically underappreciated.1 It complicates either ischemic heart disease or dilated cardiomyopathy; its prevalence varies among series but may reach ≤50% in patients with heart failure.2 When present, SMR may exhibit a broad range of severity and confers an adverse prognosis, which is worse with increasing severity of mitral regurgitation (MR).3,4 The management of SMR poses a unique set of challenges, based partly on the complexity of the valve disorder and the still-evolving adoption of the optimal therapeutic approach.5 Noninvasive imaging and, in particular, echocardiography, plays a critical role for the initial and longitudinal assessment, for individual risk stratification and outcome prediction, and for guiding intervention in patients with chronic SMR.6

Cause, Mechanisms, and Structural Remodeling

SMR develops because of a combination of mitral leaflet tethering secondary to left ventricular (LV) dilatation/deformation with papillary displacement/discoordination, annular dilatation/dysfunction, insufficient LV-generated closing forces attributable to reduction of LV contractility, and global LV/papillary muscle dysynchrony.1,5 Tethering of the mitral leaflets is the principal lesion of SMR and results in restriction of systolic leaflet motion, namely type IIb of Carpentier’s classification. SMR does not typically occur in global LV dysfunction without tethering. However, once tethering occurs, leaflet closure is further impaired by LV dysfunction because there is decreased force opposing tethering.7,8

The key event in the pathogenesis of SMR is the distortion of normal LV geometry—regional and global LV remodeling—with subsequent apical and lateral displacement of papillary muscles, which, in turn, draws the chordae tendineae away from the line of coaptation.7,8 The extent of LV systolic dysfunction and dilatation is weakly correlated to the degree of SMR unless accompanied by geometric distortion in the region of the papillary muscles.1,9 With progressive global/regional LV remodeling, the geometric distortion in the region of papillary muscles insertion increases and the SMR is worsened. However, in ischemic cardiomyopathy, SMR may be mitigated by papillary muscle elongation/remodeling via a reduction in leaflet tethering forces.10 Regional mechanical LV dyssynchrony especially that involves the papillary muscles can promote SMR through a reduction in the systolic contraction of the posterior mitral annulus.11 Global LV dysynchrony contributes to SMR through a blunted LV systolic pressure rise resulting in decreased closing forces and leaving the leaflet deforming tethering forces relatively unopposed.12,13

The morphological hallmark in SMR is the deformation of the mitral valve2 the extent of which is the major determinant of the degree of SMR.14,15 The retracted chordae tendineae secondary to LV remodeling and displacement of papillary muscles tether the valve leaflets, preventing normal valve closure and resulting in valvular incompetence. The tethering shape varies according to the site and extent of LV remodeling.16 In asymmetrical tethering, the tenting typically predominates in the region of the posterior-medial scallop of the posterior leaflet (P3) because of apically and posteriorly displaced posterior papillary muscle secondary to localized LV remodeling12 (Figure 1). Tethering of secondary chords contributes to development of a hockey-stick deformity of the anterior leaflet.7 As a result, the coaptation point of the leaflets is displaced posteriorly with respect to the center of the LV cavity. The consequence is anterior leaflet over-ride with a posterior SMR jet. In symmetrical tethering, the coaptation point of the mitral valve is moved apically, with a large tenting and both leaflets are involved to a similar degree, causing a central regurgitant jet. Tethering is thus higher in patients with global LV remodeling16,17 (Figure 2). Alterations in mitral annulus size and shape contribute to the development of SMR as an adjunctive mechanism18 (Figure 3). In fact, patients with isolated mitral annular dilation in the absence of ventricular abnormalities show less MR than those with dilated cardiomyopathy, even after correction for annulus size.19 The normal mitral leaflet area is more than double the area of the annulus, indicating a significant reserve before annular enlargement can lead to noncoaptation. Annular dilatation is not limited to anterior–posterior region but extends to intertrigonal zone and results in loss of the typical saddle shape of the annulus, which reduces leaflet curvature and thereby increases leaflet stress.16 Alterations in annular contraction also interfere with leaflet coaptation and contribute to SMR. Severe asymmetrical LV remodeling in the basal inferoposterior LV and the consequent asymmetrical annular dilatation particularly exacerbate SMR because of less reduction in annular size during systole.5

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In addition, mitral valve leaflets undergo progressive remodeling with elongation, stiffening, increase in matrix thickness, and fibrosis in response to the stresses imposed by increased leaflet tethering and mitral annular dilatation.\textsuperscript{20–22} The extent of this remodeling differs in individual patients and for a similar degree of mitral leaflet tethering and LV remodeling, suggesting that leaflet enlargement is not attributable to a straightforward passive adaptation to leaflet stretch. In SMR, mitral leaflet area may increase by \(\leq 35\%\) on average.\textsuperscript{21} However, such an adaptation is

\textbf{Figure 1.} Assessment of mitral valvular deformation and global and regional left ventricular remodeling in patients with secondary mitral regurgitation. A, Symmetrical tenting pattern. B, Asymmetrical tenting pattern. C, Measurements of the tenting area (TA), coaptation distance (CD), and posterolateral angle (PLA). D, Measurements of the sphericity index (D1), interpapillary muscle distance (IPMD; D2), and apical displacement of the posteromedial papillary muscle (D3). PLL indicates posterior leaflet length; and SI, sphericity index.

\textbf{Figure 2.} Patients with secondary mitral regurgitation. A to D, Symmetrical tenting pattern. E to H, Asymmetrical tenting pattern. A and E, Two-dimensional parasternal long-axis view. B and F, Three-dimensional volume rendering of the mitral valve. AML indicates anterior mitral leaflet; and PML, posterior mitral leaflet. C and G, Color flow of the regurgitant jet. D and H, Three-dimensional reconstruction of the tenting region. Arrow indicates zone of regurgitation.
often insufficient to meet the need for increased leaflet area imposed by the change in mitral valve configuration. Therefore, the discrepancy in leaflet area in relation to the required closure area (leaflet area/closure area) is an important determinant of the degree of MR. Interestingly, although valve enlargement can promote mitral coaptation, it can be a counterproductive by inducing maladaptive valve thickening, stiffness, and fibrosis.

**Role of Imaging**

Two-dimensional (2D) transthoracic and transesophageal echocardiography (TEE) play a substantial role in the evaluation of SMR. However, 3D echocardiography (3DE) has contributed most to the understanding of the structural remodeling that accompanies SMR. Three-dimensional echocardiography has demonstrated superiority over 2DE in measuring LV volumes, localizing and assessing the extent of mitral valve deformation, and determining the shape of the regurgitant orifice. In patients with inadequate images or discrepant findings by echocardiography, cardiac magnetic resonance (CMR) can be used to quantify MR severity, evaluate LV anatomy and function, and assess the presence and extent of myocardial scar (late enhancement). Cardiac computed tomography is rarely indicated in the evaluation of SMR, even if the extent of mitral valve deformation and MR severity might be assessed. Cardiac computed tomography could be however useful in some cases to evaluate the coronary anatomy (ie, limited vascular access; Figure 4). Irrespective of imaging technique, the assessment of SMR needs to address findings related to mitral valve configuration, the severity of SMR, and LV remodeling because these features govern the indications for intervention and clinical outcomes.

**Evaluation of Mitral Valve Configuration**

The evaluation of mitral valve configuration should also include consideration of the MR jet direction. A posterior-directed jet (P2–P3) is usually associated with eccentric valve involvement and point of coaptation (asymmetrical tethering), whereas a central jet marks symmetrical involvement and point of coaptation (symmetrical tethering). Despite no evidence of structural mitral valve disease, the displacement of the leaflets into the LV leads to several morphological changes that can quantify the global and regional tethering burden. Annular dimensions, tenting area (region enclosed between the annulus and the mitral valve leaflets body), anterior and posterior leaflet angles, tenting height (vertical distance between the mitral annulus and the leaflet coaptation point), bending distances, and coaptation length/surface (a measure of coaptation reserve) are the commonest parameters measured. With 2D echo, tethering is best appreciated in apical 4-chamber view but can be obtained from the parasternal long-axis view. With CMR, mitral valve anatomy is best imaged by acquisition of standard short-axis, 2-, 3-, and 4-chamber long-axis views in combination with oblique long-axis cines orthogonal to the line of coaptation. Three-dimensional tenting volume correlates better with effective regurgitant orifice area (EROA) in patients with SMR. Three-dimensional echocardiography also represents the method of choice to provide accurate evaluation of leaflets remodeling and mitral annulus dimensions and dynamics. For instance, the leaflet area/closure area and leaflet area/
annular area ratios and the 3D-derived indexes of coaptation are lower in patients with significant SMR. However, it is not clear whether these 3D echo measurements provide incremental information to standard 2D echo measures for predicting outcome after treatment (Table 1).

**Evaluation of LV Remodeling/Function**

The location and extent of LV abnormalities/remodeling should be evaluated. The following findings are reported: LV volumes and shape (sphericity index), regional wall motion abnormalities, and markers of transmural necrosis (hyperechogenic segments, diastolic thickness <5.5 mm, severe reduction in segmental strain, no contractile reserve, >50% late gadolinium enhancement [LGE]). LV ejection fraction and sphericity index (actual 3D volume compared with the volume of a sphere whose diameter is derived from the major LV long axis) are obtained preferably using 3D echo. When 3D is not available, the 2D biplane Simpson rule is recommended. With CMR, the LV dimensions are derived from a series of multisection perpendicular to the long axis of the LV (10–12 contiguous slices in short-axis direction). Regardless of the method used, because it is highly dependent on loading conditions, LV ejection fraction often overestimates the systolic function in case of significant SMR. Measurement of LV maximal dP/dt using continuous-wave Doppler of SMR jet can provide noninvasive estimation of global mitral closing forces. The parameters to be used to quantify LV dysynchrony are controversial. Regional remodeling is quantified by the posterior and lateral displacements of 1 or both papillary muscles. Two-dimensional echo and CMR measurements include interpapillary muscle distance and posterior papillary–fibrosa distance. With 3D echo, the extent of papillary muscles displacement can be better evaluated. A specific advantage of CMR lies on the possibility to assess the presence and extent of myocardial fibrosis, which can be either diffuse (quantification of myocardial extracellular expansion based on T1-mapping sequence) or focal (LGE). The more extensive the scar burden/focal fibrosis (LGE-CMR), the lower the likelihood of LV reverse remodeling after revascularization or cardiac resynchronization therapy. Cardiac computed tomography vein assessment in combination with LGE-CMR might be considered as reasonable options among other imaging tests (ie, invasive venous angiography, stress echocardiography, nuclear imaging) in patients thought to have extensive necrosis (Figure 5). The added value of T1-mapping to conventional LGE-CMR evaluation in patients with SMR still needs to be evaluated.

**Assessment of Severity of MR**

Echocardiographic assessment of the severity of SMR relies on an integrated approach using qualitative, semiquantitative, and quantitative parameters. A challenge in the evaluation of the degree of SMR is the temporal variation both within systole of a single cardiac cycle and during interval follow-up imaging related to loading conditions and changes in LV size.

**Current Methods and Limitations**

A combination of 2DE, spectral, and color Doppler parameters form the standard approach to quantify the degree of SMR. Despite the influence of settings, hemodynamic conditions, and mechanism of MR, the presence of a central color jet with a jet area <4 cm² or a jet that measures <10% (MR jet/left atrial area) is highly suggestive of mild SMR. On the contrary, large central MR jets reaching left atrial roof and pulmonary veins (with or without systolic spectral Doppler flow reversal) suggest significant MR. The vena contracta (VC) width in the parasternal long-axis view is another measure of the severity with a cutoff of ≥0.4 cm indicative of severe SMR. It should be noted that although VC is relatively independent...
Table 1. Two- and Three-Dimensional–Derived Echocardiographic Parameters Obtainable in SMR

<table>
<thead>
<tr>
<th>Mitral Parameters</th>
<th>2D TEE</th>
<th>3D TEE</th>
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<td>Annulus</td>
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<td>Intercommisural distance</td>
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<td>Septolateral distance</td>
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<tr>
<td>Annulus dynamics</td>
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<tr>
<td>Leaflets</td>
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<td>Anterior leaflet area</td>
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<td>Leaflet angle</td>
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<td>Anterior leaflet angle</td>
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<tr>
<td>Coaptation</td>
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<td>Coaptation depth</td>
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<td>Coaptation indexes</td>
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<tr>
<td>Leaflet coaptation area</td>
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<td>Tethering</td>
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<td>Tenting area</td>
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<tr>
<td>Tenting volume</td>
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<tr>
<td>Ventricle</td>
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<td>++</td>
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<tr>
<td>LV end-diastolic volume</td>
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<tr>
<td>LV end-systolic volume</td>
<td>+</td>
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<tr>
<td>LV dysynchrony (global/PMs)</td>
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<td>++</td>
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<tr>
<td>Interpapillary distance</td>
<td>+</td>
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<tr>
<td>PM tip</td>
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<td>PM body</td>
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<td>SMR evaluation</td>
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<tr>
<td>PISA shape</td>
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<tr>
<td>Regurgitant orifice geometry</td>
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2D indicates 2-dimensional; LV, left ventricular; PM, papillary muscle; PISA, proximal isovelocity surface area; SMR, secondary mitral regurgitation; and TEE, transesophageal echocardiography.

of the driving pressure and flow rate, it is influenced by the dynamic variations in orifice area, which occurs with SMR. Calculation of EROA using the surface area of the proximal flow convergence (PFCR) is another approach to quantify SMR. The PFCR is imaged in the apical 4-chamber or apical long-axis view with appropriate settings, and the radius of the largest flow convergence in either view is used to measure the EROA using the standard formula assuming a hemispheric flow convergence. A cutoff EROA of ≥0.20 cm² is recommended for grading SMR as severe.¹ Dynamic changes in MR, the predominant noncircular regular orifice shape, and constrained PFCR in the setting of regional myocardial deformity and leaflet tethering can limit the use and accuracy of the PFCR-based EROA computation in SMR. Regurgitant volume and fraction (RV/regurgitant fraction) in SMR is a valuable index of severity, but it is not routinely done by 2D echo/Doppler because of the need for cumbersome manual measurements in which small errors result in significant inaccuracies. Furthermore, individual measurements are made in disparate cardiac cycles, which introduce further errors. An RV of ≥30 mL signifies severe SMR (Figure 6).

Newer Methods and Approaches
Recent advances in real-time 3D B-mode and color-flow Doppler (CFD) echocardiography have been valuable in making this technology more practical for routine clinical use.³¹ Furthermore, CMR may prove to be a good alternative and complementary to echocardiography in selective instances. Three-dimensional B-mode TEE and CMR studies show that the anatomic regurgitant orifice tends to track along the leaflet coaptation plane and have an elliptical, irregular, or linear shape in vast majority of patients with SMR.³² Manual planimetry of the orifice area yields anatomic regurgitant orifice area, which is an important adjunct to flow-based measures of EROA. The anatomic regurgitant orifice area closely tracks the color Doppler–based VC area.³³ The latter can be now obtained from 3D color Doppler imaging by both transthoracic echocardiography and TEE. A VC area of ≥0.41 cm² seems to be indicative of severe MR although further validation of this cutoff is necessary given that the comparison standard in most has been conventional 2D estimates of severity of MR. Also, the cutoff value for what constitutes severe SMR may be lower than in degenerative MR.³⁴ Despite these limitations, VC area is probably more accurate than VC width by 2D in SMR.³⁵,³⁶ Imaging the PFCR with 3D CFD has emphasized the considerable limitations of 2D-based computation of EROA. Several studies have confirmed that 3D is superior to 2D in more accurately characterizing the shape and the surface area of the PFCR in SMR. A more recent advancement in which the 3D surface area of the PFCR can be automatically measured and integrated during the duration of systole allows computation of a peak 3D EROA and integrated 3D RV by real-time volume CFD imaging. The latter has been specifically shown to be accurate when compared with CMR RV in SMR, although further confirmation of this is required in larger numbers and in multiple centers.³⁷ The intrinsic limitations of angle dependency of velocity profiles of the PFCR in relation to transducer location still apply to 3D CFD imaging, but it is more accurate than 2D in characterizing the shape, and automation makes it attractive for routine clinical application. Another approach using real-time 3D volume CFD imaging is to measure mitral and LV outflow tract stroke volume and compute RV. The individual stroke volume can be measured automatically by integrating the volume color Doppler data and the area of the mitral annulus and LV outflow tract simultaneously through the cardiac cycle. This automated approach has been shown to be both more accurate and reproducible than using the spectral Doppler velocity profiles and 2D areas.³⁸ Furthermore, when extended specially to SMR, the RV/regurgitant fraction (regurgitant fraction) computed by this automated 3D CFD imaging compares well to CMR.³⁷ This promising single-center experience will need additional validation in a larger population. The presence of high-velocity aliasing in the LV outflow tract can limit the accuracy and the application of this technique just as the presence of multivalve disease. However, it does provide a solution in the presence of multiple MR jets, when VC and PFCR methods are not valid or reliable or to evaluate the residual MR through a dual orifice after MitraClip intervention.
CMR is a valuable adjunct to echocardiography in the evaluation of SMR (Figure 7). When 2DE/3DE is available and optimal, the incremental role of CMR is limited with respect to grading severity of SMR. It is arguably superior to any echocardiographic approaches for the measurement of RV/regurgitant fraction. Similarly, accurate measurement of LV volume/size as an index of severity of MR may be potentially more accurate with CMR although 3DE is proving to be as good a technique and more practical. Although VC width, VC area, and anatomic regurgitant orifice area can be assessed by CMR, it is only necessary when a combination of transthoracic echocardiography and TEE does not provide the information or there is discrepant information. The presence of atrial fibrillation in up to a third of patients with SMR further limits the use of CMR. The potential real strength of routine CMR in SMR may be in the assessment of the LV myocardial scar burden and perhaps mechanical function, which may predict clinical outcomes and success of interventions for SMR.

**Imaging Interventions**

Although interventions to treat SMR remain unsatisfactory with respect to survival benefit, there is evidence that reducing or abolishing SMR improves symptoms and attenuates progressive adverse LV remodeling. Interventional approaches have mainly focused on the mitral valve, and although various subvalvular techniques have been explored, there is no consensus on which of these are optimal. Recent advances in device-based therapies have further expanded the options in SMR, all of which have refocused the role of imaging in the selection, guidance, and postprocedural assessment.

**Surgical Approaches**

The predominant surgical technique is restrictive annuloplasty usually in the setting of coronary artery bypass grafting and in the presence of at least moderate SMR. The role of TEE in restrictive annuloplasty is primarily to quantify the degree of MR and to quantify extent of mitral valve tethering. The presence of significant tethering does not preclude annuloplasty, but it seems to be predictive of residual and recurrent MR (Table 2). Whether novel 3D parameters of mitral valve geometry (mitral leaflet size, annulus surface/area, coaptation indexes) by delineating the mechanisms associated with failure of annuloplasty improve the selection of patients and minimize procedural failure remains to be demonstrated. Similarly, whether the extent of preoperative LV remodeling is predictive of recurrent MR is still an ongoing question. Annuloplasty is performed with rings, which are specifically designed to accommodate the mitral annulus distortion (IMR ETlogix and GeoForm rings, Edwards Lifesciences, CA) without replicating the normal mitral annulus. Sizing the ring can be guided by intraoperative TEE. Figure 8 illustrates an example of how this can be done using intraoperative 3D TEE. The role of imaging in subvalvular approaches such as secondary chord cutting and repositioning of the papillary muscles is evolving, but preliminary reports of 3D modeling of papillary muscles and chords to aid interventions are encouraging and may help to reduce postoperative recurrence of MR. When ventricular plication techniques are used, imaging may be a valuable aid to assess ventricular shape and size.

**Device-Based Approaches**

Cardiac resynchronization therapy and percutaneous mitral valve repair techniques can decrease MR in selected groups of patients eligible for these treatments. Furthermore, there is also evidence for combined device approach or surgical–device approach to maximize the benefits of interventions in SMR. Recent studies have applied stringent and varied inclusion criteria for device implantation and therefore have potentially excluded a substantial number of patients who might have benefited from these approaches and have included degenerative
and SMR in 1 cohort applying the same cutoff for severity of MR. Moreover, the disparate approach to defining imaging response to these device based may impede progress in this field. Thus, there is an urgent need to standardize selection of patients, guidance and assessment of procedural success, and definition of response versus nonresponse, especially with the ongoing Clinical Outcomes Assessment of the MitraClip Percutaneous Therapy for Extremely High-Surgical-Risk Patients (COAPT) and Randomized Study of the MitraClip Device In Heart Failure Patients with Clinically Significant Functional Mitral Regurgitation (RESHAPE-HF) studies, which are focused on application of MitraClip to patients with SMR.

**Unresolved Issues in Imaging Response to Treatment**

Imaging can assess absence of MR after annuloplasty or replacement with accuracy, but evaluation of reduction of MR is more challenging especially when the postprocedural...
anatomy is unusual such as the double orifice after MitraClip implantation (Figure 9). It is best to use the same integrative assessment to quantify residual MR as the one that is recommended for quantification of native SMR. The added value of 3D echocardiographic assessment of residual MR severity is unknown, but intuitively this would make sense, and data from ongoing prospective studies will help to ascertain this. At the present time, the definition of success of those procedures, which primarily reduce MR, remains unclear. The current imperfect practice in which an acceptable reduction in SMR is defined as a decrease by 1 grade in severity is perhaps the most practical. In the acute stage, an increase in forward stroke volume (>10%–15%) may also be a marker of good response to treatment. However, whether these responses necessarily translate into chronic improvement of cardiac performance is unknown. Similarly, a positive LV remodeling process after cardiac resynchronization therapy is defined as reduction in LV end-systolic volume (>10%–15%). However, the level to which the reduction in MR plays a role in reverse LV remodeling remains unclear and requires to be evaluated prospectively. It may be also possible to better select patients and predict response in the future based on accumulated postprocedural imaging data on the degree of reduction in MR, positive LV remodeling, and clinical outcomes.

Imaging to Predict Progression and Clinical Outcomes

Disease progression in SMR is a complex interplay between LV remodeling which predates MR and the latter which can initiate and promote LV remodeling through increased wall stress and activation of neurohumoral mechanisms. SMR is fundamentally a ventricular disease; hence, the remodeling of the LV, the left atrium, and the mitral annulus are main determinants for predicting the progression of MR. However, SMR begets progressive SMR as shown by Enriquez-Sarano et al who reported an average yearly increase in RV of 1.5±9 mL and 0±5 mm² in EROA in a small group of patients (n=10). In addition, the presence of
The presence and degree of SMR is associated with a 2-fold increase in mortality (38±5% versus 61±6%; P<0.001) after myocardial infarction, with increased mortality even in the presence of mild MR. In patients with ischemic congestive heart failure and minimal or no symptoms and in those with New York Heart Association II and III class, the presence of SMR, and especially in those with EROA ≥20 mm², mortality and morbidity was increased. Rossi et al recently emphasized in a large multicenter study the importance of quantifying SMR to predict all-cause mortality or hospitalization for worsening of heart failure. Interestingly, their cohort included >400 patients with ischemic dilated cardiomyopathy. Even in idiopathic dilated cardiomyopathy, severe SMR, defined as RV >30 mL or EROA ≥20 mm² or VC >0.4 cm, was associated with a 2-fold increased risk of adverse events after adjustment for LV ejection fraction and diastolic function. In a retrospective analysis of 2242 patients, the presence of moderate MR or even mild MR at the time of coronary artery bypass grafting was associated with lower 5-year postoperative survival as compared with patients without SMR (70±1%, 84±1%, and 86±1% for patients with moderate, mild, or no MR, respectively), even after adjustment for the degree of LV dysfunction and other comorbidities. SMR is often dynamic with intermittent changes related to changes in loading conditions. An increase in EROA by ≥13 mm² during exercise is associated with reduced event-free survival, higher risk of heart failure, and major cardiac events compared with patients with no or mild increase or with decrease in MR severity.

Despite all this, debates still exist about whether SMR is only a marker of poor LV function or an independent risk factor for adverse outcome. Indeed, the severity of SMR tends to follow the severity of LV dysfunction causing SMR. Moreover,
correcting SMR does not prolong life, which would be expected to occur if MR affected the outcome.\textsuperscript{11,42} Currently, no outcome studies using contemporary imaging techniques to quantify MR and evaluate LV function/remodeling have been conducted to address this dilemma. In this regard, large-scale studies examining the added value of 3DE and CMR are needed.

Future Directions
Better understanding of the origin and evolution of SMR is necessary to enable building more effective therapeutic strategies of this complex valvular disease. Detailed mapping of the geometric substrates promoting SMR will likely allow developing tailor-made innovative technical approaches targeted toward specific mechanisms. Advances in diagnostic imaging techniques will continue to play a major role in the assessment of SMR. Our role as imagers will be to identify the best imaging modalities in individual patients. The Progressions and Outcome of Secondary Mitral Regurgitation (POMAR) study has been specifically designed to address this question by comparing real-time 3D color transesophageal echocardiography and CMR imaging for the assessment of MR severity and LV remodeling.

Conclusions
A comprehensive evaluation of SMR includes the quantitative description of the structural changes in the LV and the mitral valve apparatus and the severity of MR predominantly using a combination of echocardiography and, if needed, of CMR. Whether newer techniques and technologies enhance the assessment of SMR and improve our ability to predict progression, outcomes, and guidance of therapy remains to be seen. Large multicenter studies that incorporate these newer approaches to image SMR may help to fill the current gaps in knowledge.

Disclosures
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References
Impact of coronary revascularization and transmural extent of scar on clinical and echocardiographic improvement after cardiac revascularization.


**Key Words**: echocardiography • heart failure • mitral valve insufficiency • prognosis
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