Mitral regurgitation (MR) is one of the most prevalent valve disorders and has numerous etiologies, including primary (organic) MR, due to underlying degenerative/structural mitral valve (MV) pathology, and secondary (functional) MR, which is principally caused by global or regional left ventricular remodeling and/or severe left atrial dilation. Diagnosis and optimal management of MR requires integration of valve disease and heart failure specialists, MV cardiac surgeons, interventional cardiologists with expertise in structural heart disease, and imaging experts. The introduction of transcatheter MV therapies has highlighted the need for a consensus approach to pragmatic clinical trial design and uniform endpoint definitions to evaluate outcomes in patients with MR. The Mitral Valve Academic Research Consortium is a collaboration between leading academic research organizations and physician-scientists specializing in MV disease from the United States and Europe. Three in-person meetings were held in Virginia and New York during which 44 heart failure, valve, and imaging experts, MV surgeons and interventional cardiologists, clinical trial specialists and statisticians, and representatives from the U.S. Food and Drug Administration considered all aspects of MV pathophysiology, prognosis, and therapies, culminating in a 2-part document describing consensus recommendations for clinical trial design (Part 1) and endpoint definitions (Part 2) to guide evaluation of transcatheter and surgical therapies for MR. The adoption of these recommendations will afford robustness and consistency in the comparative effectiveness evaluation of new devices and approaches to treat MR. These principles may be useful for regulatory assessment of new transcatheter MV devices, as well as for monitoring local and regional outcomes to guide quality improvement initiatives. (J Am Coll Cardiol 2015;66:278–307) © 2015 by the American College of Cardiology Foundation.
Mitral regurgitation (MR) is the most prevalent valvular disease in the United States and Europe, and along with aortic stenosis, is one of the most frequent valve disorders referred for surgical correction (1-4). In contrast to aortic stenosis, which is typically characterized by severe and homogenous cusp calcification, MR is heterogeneous in etiology, mechanisms, and pathoanatomy. MR may develop either from primary pathology involving any of the components of the mitral valve (MV) apparatus (primary MR, also known as organic MR, usually due to degenerative MV disease) or arise secondarily to left ventricular (LV) dysfunction or occasionally from left atrial (LA) dilation (secondary MR, also known as functional MR) (1,2,5-7). Surgical MV repair is the recommended approach for severe primary MR, with a recently accepted role for transcatheter repair for patients who are at very high or prohibitive surgical risk (1,2,8). Conversely, secondary MR is typically treated with medications and (if indicated) biventricular pacing for heart failure, and coronary revascularization when appropriate, with the utility of MV surgery and transcatheter devices representing active areas of investigation (8). Few randomized trials, however, have been performed to evaluate the safety and efficacy of MV therapies. The introduction of transcatheter MV devices and the performance of a randomized trial comparing 1 such device to MV surgery (8) have exposed the complexities required to properly evaluate MR therapies, specifically regarding the appropriate study population and control group, background medications and procedures, efficacy and safety endpoints, learning curve issues, and analysis cohort and statistical considerations (8,9). Moreover, although the outcomes of patients with MV disorders are sometimes tracked at single centers (10,11) or in national databases (12,13), no standardized endpoints and definitions have been proposed to provide consistency and uniform interpretability of reported results.

The Academic Research Consortium was organized as a collective endeavor between leading academic research organizations and physician-scientists to reach consensus as to what constitutes meaningful clinical endpoints and definitions for evaluation of cardiovascular devices (14). In collaboration with the U.S. Food and Drug Administration (FDA) and supported by device manufacturers, prior Academic Research Consortium initiatives have addressed consensus endpoints for events following percutaneous coronary intervention and transcatheter aortic valve replacement (TAVR) (15-17), as well as bleeding definitions (18), and have been adopted to improve the uniformity and interpretation of clinical studies (19). The Mitral Valve Academic Research Consortium (MVARC) working group was therefore assembled to develop endpoint definitions for clinical studies of MR therapies. In addition, given the complexity of issues that must be considered for MV trials, MVARC has also developed design principles for

SEE PAGE 322
clinical trials and registries investigating transcatheter device therapies to treat MR, which may also be applied to surgical and other approaches. Three in-person meetings were held in 2012 to 2014 in which stakeholders and experts in MV disease and therapeutics from the United States and Europe convened to comprehensively review the principles and elements required to successfully investigate and evaluate the relative risks versus benefits of MV therapies. As listed in the Online Appendix, these multidisciplinary gatherings included specialists in general cardiology and valve disorders, heart failure, cardiac surgery, interventional cardiology, imaging, statistics and epidemiology, and clinical trials. Representatives from the FDA Center for Devices and Radiological Health participated in an advisory role. MVARC was funded by multiple industry sponsors who did not participate in either the sessions or document preparation, but were provided a copy of the report before submission. No fees or honoraria were provided to the writing group or participants.

The present document that resulted from this effort is meant to summarize the current state of knowledge and consensus expert opinion for MR therapies and is organized in 2 parts: recommendations for clinical trial design principles (Part 1), and consensus endpoint definitions (Part 2). We acknowledge that the field of MV therapeutics is highly dynamic and evolving, and we anticipate regular revisions to these recommendations. Finally, we have concentrated our current effort on therapies for primary and secondary MR; however, many of the principles in this document may also be applied to other MV conditions, including treatment of mitral stenosis, degenerated mitral bioprostheses, and failed surgical valvuloplasty.

OVERVIEW: INVESTIGATIVE AND REGULATORY PERSPECTIVES

Clinical trials that are intended to support device regulatory approval or expansion of indications must have clearly documented objectives and be performed in a highly rigorous manner. In Europe, the CE mark process requires demonstration that the device is safe and functions both medically and technically as the manufacturer intends. Effectiveness is usually investigated after CE mark approval, and post-marketing surveillance is an integral part of ongoing clinical evaluation. Either randomized trials or well-performed registries may support CE mark approval. For example, both the MitraClip edge-to-edge device (Abbott Vascular, Santa Clara, California) and the Carillon coronary sinus annuloplasty device (Cardiac Dimensions, Kirkland, Washington) received a CE mark to treat MR on the basis of registry data demonstrating safety.

For U.S. FDA regulatory approval, high-risk class III devices must demonstrate “reasonable assurance” of both safety and effectiveness in a well-defined population for its intended use. Pivotal evaluations of breakthrough technologies such as transcatheter mitral repair systems or percutaneous implantable valves will, in most cases, necessitate randomized controlled trial designs wherein the new device is compared with the currently established standard of care therapy, unless approval for a very limited patient cohort is desired for which randomization is not feasible. For example, the MitraClip was approved in the U.S. to treat symptomatic patients with severe primary MR at prohibitive surgical risk on the basis of high-quality registry data.

For U.S. approval trials, depending on the comparator group, either a superiority or noninferiority design for the primary endpoint may be appropriate. Although superiority in either safety and/or effectiveness is typically preferred for FDA regulatory approval, a new device may demonstrate noninferiority for both and still be approvable as an alternative therapy to the existing standard of care, depending on the benefit-risk balance. In studies addressing an unmet clinical need for a severe disease in which the available therapeutic alternatives are suboptimal, the benefit-risk profile of an investigational device may also be favorable even if effectiveness is somewhat less than that of the comparator if treatment with the investigational device shows evidence of substantial safety benefits (and is more effective than a putative placebo) (20). As knowledge accumulates and technology matures, noninferiority designs (e.g., comparing a new design to a previously approved transcatheter device) and even non-randomized comparisons to performance goals or objective performance criteria may become reasonable to evaluate device iterations and to expand the indications for use (label expansion) of existing approved devices.

Primary effectiveness should be evaluated with a clinically relevant endpoint, either a single event type (e.g., hospitalization for heart failure) or a composite measure (e.g., death or hospitalization for heart failure). Additional support for effectiveness can be obtained through the use of validated instruments demonstrating improved quality-of-life, improvement in symptom status (e.g., New York Heart Association [NYHA] functional classification), and improved exercise performance. Although at the present time these measures are not usually sufficient for principal FDA regulatory device approval,
increasing attention is being paid to patient-centered benefit-risk metrics in device approval decisions. Evidence of meaningful MR reduction by the device that is sustained over time is important to demonstrate, and improvement in ventricular volumes and function during follow-up are additional supportive secondary effectiveness endpoints that should be assessed. Safety assessments may include both short- and long-term procedural and device-related complications, and a primary safety endpoint (separate from the primary effectiveness endpoint) should be pre-specified (see Primary and Secondary Endpoints). Finally, the duration of follow-up must be sufficient to ensure adequate device durability, relevant to the population being studied and comparable to alternative therapies, if available. Late device failures may occur after the primary endpoint of pre-market studies, necessitating robust post-market surveillance to monitor long-term device performance after regulatory approval.

Identifying the intended population for use (e.g., primary vs. secondary MR, high vs. low surgical risk, and so on) may importantly affect decisions on comparator therapies (e.g., medical, surgical, or other transcatheter devices; see Control Group Therapies). As a general principle, because the pathophysiology, prognosis, control groups, and response to therapies for primary and secondary MR vary greatly, these 2 conditions should be studied in separate investigations unless randomization is stratified and each cohort is individually powered for both safety and effectiveness. As a corollary, inclusion and exclusion criteria must be carefully selected to define the population of use (see Inclusion and Exclusion Criteria). Because transcatheter devices for MR are likely to be evaluated over a range of disease severity and comorbidities, detailed anatomic and clinical characterization is required, in addition to key surrogates such as MR quantification and structural cardiac evaluation using imaging techniques (see Assessment of Mitral Regurgitation: Role of Non-invasive Imaging).

Determining operative risk is central to defining the population for intended use of a new device as well as selecting the appropriate comparator arm. Current scoring systems such as the Society of Thoracic Surgeons (STS) and EuroSCORE II indexes (21-23) may not by themselves be sufficient to define risk or operability in all patients. Assessment of patient operability (which may define clinical trial eligibility) should be determined by a local multidisciplinary heart team after comprehensive patient evaluation (including risk score assessment). For MR studies, the heart team should usually include valve and heart failure specialists, MV surgeons, interventional cardiologists experienced in transcatheter MV procedures, imaging experts, and potentially others depending on the specific population and device being studied (see also the subsection Role of the Heart Team).

Several trials may now be cited wherein the use of a sham control helped to demonstrate a lack of device efficacy, contrary to the results of prior unblinded investigations (24-26). Use of sham controls (if possible) are thus desirable and, in most cases, are ethically justifiable (see also discussion on sham controls in Control Group Therapies). When a sham control is not feasible, additional efforts should be considered to blind the patient and participants involved in data collection to the extent possible (e.g., the use of patient headphones to mask device allocation during the procedure; not recording randomization allocation in the chart; and using separate research coordinators and physicians for device implantation and follow-up). Patient-related outcomes, such as quality-of-life, are considered more robust in studies that can be blinded. For pivotal device trials, the use of independent core laboratories and event adjudication and data safety and monitoring committees are mandatory to ensure patient safety, reduce reporting bias, and enhance credibility, accuracy, and interpretability of study findings, especially when patient and physician blinding is not possible.

For both randomized trials and registry studies of MR therapies, written informed consent must be obtained from all patients unless waivers are provided with specific ethical oversight. Within the framework of a randomized trial, study-eligible patients who decline randomization should ideally be followed in a separate registry to provide additional insights into potential study selection bias and the natural history of the control population. If exploratory comparison with randomized trial arms is contemplated, the statistical methodology must be pre-specified and justified (e.g., propensity scoring analysis with appropriate covariates, and so on).

Finally, although randomized trials with primary clinical endpoints are strongly recommended, given the logistical, time and cost constraints, MVARC acknowledges that many investigations of MV therapeutics will collect observational or registry data only (preferably compared with either a concurrent or historical control group), or if randomized, will not be powered for clinical endpoints. Potential efficacy endpoints for these studies may include reduction in MR grade, improvement in LV pressures and chamber dimensions, improved quality of life, and enhanced
functional capacity (see Primary and Secondary Endpoints). However, currently none of these endpoints have been sufficiently linked to a major clinical outcome such as death or heart failure hospitalization to be considered a true surrogate, especially as procedural risks must be taken into account when considering the benefit-risk profile of a novel therapy. As such, these studies should be considered hypothesis generating with regard to clinical utility. Nonetheless, such investigations are valuable in their own right, and they provide important supportive data when considering the utility of a new device or approach. Further studies are warranted to strengthen the association between these nonclinical endpoints and clinical outcomes such that, in the future, they might serve as primary endpoints in FDA regulatory trials.

**PRIMARY VERSUS SECONDARY MR: SIMILARITIES, DIFFERENCES, AND IMPLICATIONS FOR TRIAL DESIGN**

**CLASSIFICATION OF MR AND IMPLICATIONS FOR MV THERAPIES.** Accurate diagnosis of the underlying MV anatomy and pathophysiology is essential to understand the etiology, mechanism, lesion localization, and severity of MR; to establish its prognosis; and to design appropriate trials of MR therapies. The MV complex is a dynamic structure including the annulus, the anterior and posterior leaflets and commissures, different level chordae tendineae, the papillary muscles, the underlying LV myocardium, and the LA. Pathological changes in any of the components of the MV can lead to MR, and often lesions are present in more than 1 structural component of the valve. Assessment of MR involves comprehensive evaluation of its etiology and mechanism (the lesion or deformation resulting in valve dysfunction), including the dysfunction type (leaflet motion abnormality) (27–29). Of note, annular dilation is almost universally present in patients with severe MR, regardless of other structural abnormalities, although it typically develops late. One exception is MR arising secondary to LA dilation (often in the setting of atrial fibrillation), in which annular dilation may be the principal mechanism of MR (5,6). Comprehensive characterization of the underlying etiology and MV lesion(s) in each patient is especially critical in the new device era, as many transcatheter devices mechanistically target only a single component of the MV or a single mechanism of MR.

The mechanism of MR may be described by Carpentier’s classification of leaflet motion: type I: normal leaflet motion (e.g., annular dilation, leaflet perforation, or clefts), type II: excessive leaflet motion (e.g., chordal elongation or rupture), and type III: restricted leaflet motion (Figure 1) (30). Type III dysfunction is further subclassified according to restricted leaflet motion predominantly in diastole but also in systole (type IIIa [e.g., rheumatic disease]) versus only in systole (type IIIb [e.g., ischemic or nonischemic LV remodeling with leaflet tethering due to local or diffuse ventricular dilation]). Carpentier’s segmental leaflet anatomy classification is a useful construct when describing MV disease and planning and performing an intervention (30).

**PRIMARY VERSUS SECONDARY MR.** The first and most important distinction that must be drawn is to classify the underlying etiology as either predominantly: 1) primary MR (also commonly known as organic MR), which is due to underlying degenerative/structural MV pathology; or 2) secondary MR (also known as functional MR), which is principally caused by global or regional LV remodeling and/or severe LA dilation, in which case the MV structures are usually normal or exhibit only secondary late fibrosis and/or annular dilation. As discussed in the following text, this distinction currently serves as the central basis for selecting standard of care therapies, which will dictate the choice of control group in randomized trials.

Primary MR usually implies Carpentier type II dysfunction, but may be type I in endocarditis and type IIIa in cases of rheumatic origin. Primary MV disease is the most common form of MR referred for surgical correction and covers a large spectrum of lesions, ranging from abnormalities in an isolated scallop to multisegment (or generalized) prolapse, and from thin/nonredundant leaflets to thickened leaflets with excess tissue (Barlow’s disease) (28). Prolapse location, the presence of valvular/annular calcification, and the severity of annular dilation may affect the feasibility and choice of surgical and transcatheter mitral repair techniques (31).

Secondary MR usually implies a Carpentier type IIIb dysfunction, although type I dysfunction with isolated annular dilation may occur secondary to LA dilation. Secondary MR most commonly develops despite a structurally normal MV due to mitral leaflet tethering secondary to ventricular deformation/remodeling, annular dilation/dysfunction, and insufficient LV-generated closing forces. Assessing global LV function and dilation (diameters, volumes, sphericity, mass) and local remodeling (displacement of papillary muscles) as well as MV deformation (coaptation depth, tenting area, and tenting volume in 3 dimensions) is of paramount importance in
evaluating the potential for reparability and results of treatment (32,33). Tethering may be limited to an isolated leaflet segment on the basis of “localized” ventricular remodeling or be present along the entire MV closure line in end-stage and diffuse ventricular remodeling. The degree of secondary MR may vary greatly depending on loading conditions (more so than in primary MR).

Secondary type IIIb MR can further be subclassified as arising from underlying ischemic heart disease (usually prior myocardial infarction) versus nonischemic dilated cardiomyopathy (whether idiopathic or due to specific causes such as hypertension). The mitral jet is typically eccentric or commissural in the setting of ischemic disease and posterior infarction, resulting in posterior leaflet tethering with medial commissural gap, and is central in most cases when the LV is globally dilated due to anterior infarction or nonischemic cardiomyopathy, resulting in more symmetric displacement of both papillary muscles.

It is particularly important to differentiate and separate populations of patients with primary versus secondary MR in clinical trial design (Table 1), as the comorbidities, prognosis, and therapeutic approaches in these patients vary greatly. Most patients with primary MR due to degenerative MV disease achieve long-term event-free survival similar to an age-matched population after MV surgery, provided MR correction is achieved through valve repair surgery rather than valve replacement, and before significant deterioration in LV geometry or function (1). In contrast, patients with secondary MR have varying degrees of myocardial remodeling and dilation, and usually have significant LV dysfunction. Most patients with secondary MR are treated with heart failure therapies (guideline-directed medical therapy [GDMT] ± cardiac resynchronization therapy [CRT])
TABLE 1 Implications of the Etiology of Mitral Regurgitation

<table>
<thead>
<tr>
<th>Prognosis</th>
<th>Primary Mitral Regurgitation</th>
<th>Secondary Mitral Regurgitation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Principal management strategy (standard of care)</td>
<td>Primarily dependent on the severity of mitral regurgitation and secondarily on left and right ventricular function and pulmonary pressures</td>
<td>Primarily dependent on the degree of underlying left ventricular dysfunction and secondarily on the severity of mitral regurgitation</td>
</tr>
</tbody>
</table>

GDMT = guideline-directed medical therapy.

when appropriate) as well as coronary revascularization if substantial ischemia is present. For patients failing those initial treatments, advanced therapies including LV assist devices and heart transplantation may be considered. In patients with severe LV dysfunction, the long-term prognosis may be dictated more by the extent of ventricular dysfunction and remodeling than the severity of secondary MR. There is currently little evidence that survival or the natural history of the underlying myocardial disease are affected by mitral intervention in patients with secondary MR, although reduction or correction of MR may provide symptomatic relief (34–36).

ASSESSMENT OF MR: ROLE OF NONINVASIVE IMAGING

ECHOCARDIOGRAPHIC EVALUATION OF MR. Echocardiography is fundamental in evaluating the etiology, mechanisms, and severity of MR, and its effect on cardiac structures and function. In addition, serial echocardiography is essential to demonstrate the effects of medical therapy, devices, and surgical MV repair and replacement over time. Routine 2-dimensional (2D) transthoracic echocardiography (TTE) differentiates whether MR is due to primary valve degeneration or is secondary to LV dysfunction or LA dilation. For primary MR, 2D TTE discriminates the specific pathological changes in the MV complex. In the presence of mixed pathologies, classification can be more difficult (e.g., secondary MR with notable annular calcification or leaflet thickening), although usually a predominant etiology can be assigned.

Specific anatomical measurements are also useful in assessment of secondary MR (Figure 2), including leaflet length, leaflet angles (particularly the posterolateral angle, indicating posterior leaflet tethering), coaptation distance (apical displacement of the coaption point), coaptation length, and tenting area. Asymmetric tenting indicates posterior leaflet restriction, whereas symmetric tenting indicates bileaflet restriction. Measurements of global LV remodeling include LV diameters/volumes and the sphericity index. Measurements of local LV remodeling include apical displacement of the postero-medial papillary muscle, second order chords, and the interpapillary muscle distance (Figure 2) (29,37). Finally, echocardiographic measures of annular dimensions (anterior-posterior diameter >35 mm or the ratio of the anterior-posterior diameter to mid-diastolic anterior MV leaflet length >1.3) due to LV dysfunction, dilation, or dyssynchrony have prognostic significance (37,38).

QUANTIFICATION OF MR. Three echocardiographic grades of MR severity are generally recognized: mild, moderate, and severe. Whereas this 3-group classification is preferred, a 4-group quantitative scale is sometimes used as well, wherein 1+ = mild MR, 2+ = moderate MR, 3+ = moderate-to-severe MR, and 4+ = severe MR. Because each echocardiographic measurement has specific limitations and lack of precision, an integrated approach incorporating multiple variables should be used to assess MR severity, with somewhat different criteria for primary and secondary MR (Tables 2 and 3) (29,39). These include qualitative findings (MV morphology, color flow, and continuous wave signals of the MR jet), semiquantitative measures (vena contracta width, pulmonary vein flow, mitral inflow), and quantitative measures (regurgitant volume [RVol] and effective regurgitant orifice area [EROA]), as well as supportive findings (enlarged LV and/or LA, increased pulmonary artery pressure [PAP]) (Figures 3 and 4). MR severity should be evaluated by 2D TTE in the non-sedated, nonanesthetized patient, although 2D and 3-dimensional (3D) transthoracic echocardiography (TTE) may improve assessment, particularly in secondary MR (Figure 5). Moreover, for consideration of patient eligibility for a trial evaluating treatment of chronic MR, the echocardiographic severity of MR must be evaluated during a period of clinical stability. If the patient presents with decompensated LV failure, the degree of MR should not be assessed until at least 30 days after the patient has stabilized on a maximal medical regimen.

Color flow imaging is not solely used for grading MR severity. Localization, duration, timing, and direction of the regurgitant jet into the LA may be useful to evaluate MR, both at baseline and during follow-up after device or surgical intervention. When feasible, the vena contracta width and the
flow convergence method (proximal isovelocity surface area [PISA], which permits assessment of RVol and EROA) are strongly recommended. Inherent limitations of the PISA method should be appreciated, however, including reduced accuracy with eccentric or multiple jets (especially common in secondary MR or after transcatheter MV repair with certain devices), changes in PISA radius throughout systole, and difficulty in precisely locating the regurgitant orifice. In addition, the assumption that the proximal flow convergence is hemispheric (vs. ellipsoidal or irregularly shaped, as in secondary MR [40], leading to underestimation of MR severity) and that it occurs over a flat surface (requiring angle correction in some cases, including post-MitraClip) are important limitations. By permitting direct planimetry of the vena contracta (as well as multiple jets), 3D-TEE may provide a more accurate assessment of MR severity, especially in secondary MR (41,42). However, both 2D and 3D color flow Doppler may overestimate the orifice area due to aliasing and blooming artifacts. Despite these limitations, PISA is a
practical method that correlates well with the severity of MR and prognosis. Importantly, systolic regurgitant flow lasts only as long as mitral leaflet malcoaptation persists; therefore, EROA and RVol are dynamic. For example, in MV prolapse, the EROA appears or increases in mid-to-late systole, whereas in secondary MR, it decreases in mid systole. EROA is usually holosystolic in severe MR. In the current valve guidelines from both the United States and Europe (1,2), an EROA ≥40 mm² (RVol ≥60 ml) indicates severe primary MR, whereas an EROA ≥20 mm² (RVol ≥30 ml) indicates severe secondary MR. These different thresholds for severe MR due to primary and secondary MV dysfunction have been largely derived from outcome studies demonstrating the prognostic effect of varying degrees of quantitatively measured MR in the 2 conditions (29,43). In both cases, however, the regurgitant fraction is ≥50%. Of note, however, a regurgitant fraction ≥50% can be produced by different values of EROA and RVol, depending on LV volumes and ejection fraction, which can vary widely in secondary MR. Therefore, defining severe MR requires careful integration of all echocardiographic data (Tables 2 and 3) (44).

Exercise echocardiography can demonstrate the dynamic nature of MR (mild-moderate MR increasing to severe MR during exercise) and exercise-induced pulmonary hypertension (45). In asymptomatic patients with primary MR and borderline normal values of LV function and size, worsening of MR (with increasing systolic PAP) and lack of contractile reserve during exercise echocardiography are associated with worse outcomes (46). In patients with secondary MR and chronic LV dysfunction, worsening MR with increase in EROA by ≥13 mm² with exercise is associated with a poor prognosis (47,48). Increasing LV dyssynchrony with increased MR can also occur during exercise and may improve after CRT. Improved regional wall motion during (low-level) exercise indicates re- sidual viability, whereas worsening regional wall motion indicates ischemia. Although exercise echocardiography is increasingly used, the accurate assessment of MR severity during peak exercise remains technically challenging. Pharmacological stress alone is incapable of comprehensively evaluating

### TABLE 2  Grading the Severity of Primary Mitral Regurgitation by Echocardiography

<table>
<thead>
<tr>
<th>MR Severity*</th>
<th>Mild</th>
<th>Moderate</th>
<th>Severe</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Quantitative</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>EROA, mm²</td>
<td>≤20</td>
<td>20–29, 30–39†</td>
<td>≥40</td>
</tr>
<tr>
<td>Regurgitant volume, ml</td>
<td>≤30</td>
<td>30–44, 45–59†</td>
<td>≥60</td>
</tr>
<tr>
<td>LV and LA size</td>
<td>Usually normal</td>
<td>Usually normal or mild dilation</td>
<td>Usually dilated‡</td>
</tr>
<tr>
<td>PA systolic pressure, mm Hg</td>
<td>Usually normal</td>
<td>Usually normal</td>
<td>May be normal; &gt;50 at rest without other cause</td>
</tr>
</tbody>
</table>

**Qualitative**

| MV morphology | Mildly abnormal leaflets (e.g., mild rheumatic thickening, limited prolapse) | Moderately abnormal leaflets (e.g., moderate thickening or prolapse) | Severe valve lesions (e.g., flail leaflet, ruptured papillary muscle, severe retraction, large perforation) |
| Flow convergence zone‡ | Not visible, transient or small | Intermediate in size and duration | Large throughout systole |
| Color flow MR jet | Small LA penetration or not holosystolic | Moderate LA penetration or large penetration and late systolic | Deep LA penetration and holosystolic jet |
| CW signal MR jet | Faint/partial/parabolic | Dense but partial or parabolic and light density | Holosystolic and dense or triangular |

**Semi-quantitative**

| Vena contracta width, mm | <3 | Intermediate | ≥7 (>8 for biplane)† |
| Pulmonary vein flow | Systolic dominance | Systolic blunting§ | May be normal with low LA pressure. Systolic flow reversal |
| Mitral inflow| A-wave dominant | Variable | E-wave dominant (>1.5 cm/s) |
| TVI mitral/TVI aortic ratio | <1.0 | 1.0–1.4 | >1.4 |

*MR severity determined in an individualized manner that accounts for body size, sex, and all other patient characteristics. †With Nyquist limit. ‡For average between apical 2- and 4-chamber views. §Signs are nonspecific and are influenced by many other factors (LV diastolic function, atrial fibrillation, LA pressure). ¶The 2 ranges indicate mild/moderate and moderate/severe MR respectively. EROA 30 to 39 mm² or RVol 45 to 59 ml may be consistent with severe MR in individuals of small body size, particularly women. #LV and LA can be within the “normal” range for patients with acute severe MR or with chronic severe MR who have small body size, particularly women, or with small LV size preceding the occurrence of MR. Modified with permission from Lancellotti et al. (29) and Zoghbi et al. (39).

CW = continuous wave; EROA = effective regurgitant orifice area; LA = left atrium; LV = left ventricular; MR = mitral regurgitation; MV = mitral valve; PA = pulmonary artery; TVI = time velocity integral.
Dynamic changes in MR. Further studies are warranted to evaluate the role of exercise echocardiography in the risk stratification of patients with MR. **Assessing the Consequences of MR.** LV diameters are derived from M-mode echocardiography or 2D imaging. LV end-systolic diameter >40 to 45 mm and left ventricular ejection fraction (LVEF) <60% are indicators of LV systolic dysfunction/dilation in the patient with severe MR. The 2D-based biplane Simpson’s method is recommended for estimation of LV volumes and LVEF; 3D assessment of LV function is generally more accurate than 2D imaging. The LA dilates in chronic volume and pressure overload; the biplane area-length method using apical 2- and 4-chamber views is recommended for assessing LA size. An LA volume index >60 ml/m² predicts a poor prognosis in primary MR. However, LA dilation is more nonspecific than LV dilation, as LA enlargement can also occur in the setting of atrial fibrillation or secondary to an increase in LV enddiastolic pressure, whether due to LV diastolic or systolic dysfunction. A systolic PAP (obtained as the sum of the transtricuspid pressure gradient and the estimated right atrial pressure) >50 mm Hg at rest or >60 mm Hg with exercise is strongly associated with adverse outcomes in primary MR. Elevated PAP results in right ventricular pressure overload, which may induce right ventricular failure, and echocardiographic signs include right ventricular hypertrophy, dilation, impaired function, and increased tricuspid regurgitation (peak jet velocity >3 m/s) (49). Tricuspid annular dilation (≥40 mm or >21 mm/m²) contributes to tricuspid regurgitation after MV surgery, in which case tricuspid annuloplasty may be considered concomitant with MV surgery (29).

**Echocardiographic Eligibility Criteria for Surgical and Transcatheter MV Repair or Replacement.** Surgical and transcatheter MV repair or replacement is generally reserved for severe MR (3+ to 4+) (1,2,50). Echocardiographic eligibility criteria must also carefully consider the likelihood of procedural success for surgery and the experimental transcatheter device. For example, in primary MR,
successful surgical MV repair may be compromised in the setting of multiple complex regurgitant jets, extensive leaflet or valve calcification, and/or when ≥3 scallops (particularly affecting the anterior leaflet) are involved (51,52). In secondary MR, the risk of unsuccessful surgical repair or MR recurrence is increased with the presence of severely altered geometry of the MV apparatus, severe global LV remodeling, and/or extensive basal LV scar or aneurysm (Table 4) (29).

Given their varying mechanisms of action, the echocardiographic determinants of successful transcatheter repair of MR are likely to be device specific. For example, the MitraClip reduces MR by grasping and approximating the anterior and posterior mitral leaflets (8). Echocardiography is indispensable in determining the complexity of the anatomic lesion, and whether the amount of leaflet tissue and coaptation depth and length are sufficient to afford leaflet grasping and approximation by the MitraClip (Table 5) (52). The precise echocardiographic features for procedural success or failure for transcatheter devices that reduce MR by other mechanisms, such as direct or indirect annuloplasty and MV replacement, are notably different and unique to each device.

**IMAGING DURING AND AFTER MV REPAIR AND REPLACEMENT PROCEDURES.** Echocardiography is vital for assessing the acute and late results of both surgical and transcatheter mitral interventions. As regards surgery, 2D TEE (complemented by 3D TEE when available) is performed acutely in the operating theater after surgical MV repair to exclude more than
mild residual valvular MR (e.g., vena contracta width >3 mm). Adequate leaflet coaptation (length ≥8 mm) should be verified. Leakage due to anatomic/technical problems or ring dehiscence and MV stenosis (MV area <1.5 cm², mean transmitral gradient ≥5 mm Hg) should be excluded (53). Systolic anterior motion of the MV and injury to the left circumflex artery (expressed as wall motion abnormalities in the basal and mid inferolateral LV segments) due to the close proximity of sutures needed for annuloplasty ring fixation or compression by the ring itself should also be excluded (53).

LV function may worsen after surgical MV repair and should thus be evaluated in the immediate post-operative period. Historically, this has been attributed to the increase in LV afterload due to reduction in MR. However, after MitraClip repair, cardiac output generally increases, LV filling pressures tend to normalize, and significant LV dysfunction is uncommon, even in patients with severe baseline LV dysfunction (54). This suggests that the LV dysfunction observed in some patients after MV surgery may be attributable to myocardial oxidative stress, systemic inflammation and free radical injury from cardiopulmonary bypass, cardiac arrest, and cardioplegia, rather than to increased afterload due to the reduction in MR (55).

In addition to assessing the acute results of transcatheter device repair or replacement of MR, echocardiography is essential to guide most transcatheter MV procedures. For example, 2D and 3D TEE are used to guide each step of MitraClip implantation (49), complementing fluoroscopy. Immediate post-procedural echocardiographic evaluation includes assessment of residual MR, potential MV stenosis, and exclusion of complications (e.g., pericardial effusion/tamponade, thrombus formation on clips, partial clip detachment, and entrapment of chordae by the clip).

Depending on the device, echocardiographic assessment of MR severity after transcatheter MV procedures may pose unique challenges. For example, MR quantification with color flow Doppler is
TABLE 4 Unfavorable Transthoracic Echocardiographic Characteristics for Surgical Mitral Valve Repair in Secondary Mitral Regurgitation

1. Mitral valve remodeling
   - Coaptation distance ≤10 mm
   - Tenting area >2.5-3.0 cm²
   - Complex regurgitant jets
   - Posterolateral angle >45°

2. Local left ventricular remodeling
   - Interpapillary muscle distance >20 mm
   - Posterior papillary-fibroa distance >40 mm
   - Lateral wall motion abnormality

3. Global left ventricular remodeling
   - End-diastolic diameter >65 mm
   - End-systolic diameter >51 mm (end-systolic volume >140 ml)
   - Systolic sphericity index >0.7

Adapted with permission from Lancellotti et al. (29).

complex in the setting of a double MV orifice after the MitraClip, and artifacts from the clip(s) hamper quantification. Pulse wave Doppler of the pulmonary veins is useful to corroborate a reduction in MR. Specifically, pulmonary vein flow reversal should be eliminated, and there is often a conversion from the pulmonary vein flow from a “D” dominant pattern (consistent with an elevated LA pressure) to an “S” dominant pattern, reflecting a drop in LA pressure secondary to MR reduction. MV stenosis should be excluded, as evidenced by mean transmitral valve gradient <5 mm Hg and MV area ≥1.5 cm². Assessment of paravalvular leak is particularly important in patients undergoing transcatheter and surgical MV replacement (56, 57).

For both surgical and transcatheter MV procedures, evaluation of the immediate post-repair/replacement results should be performed when the patient’s blood pressure is at least equal to the basal state (and after the effects of anesthesia have worn off). Comprehensive follow-up TTE is typically recommended at 1 month, at 6 months, and then annually to serially assess MR severity, chamber volumes and pressures, and structural and functional device performance (including the detection of specific device-related technical failure issues and complications as discussed in part 2 of this document).

For clinical trials using serial echocardiographic imaging to assess device performance, study-specific site training and certification in imaging quality before enrollment are recommended, and should be conducted in collaboration with an independent echocardiographic core laboratory.

TABLE 5 Relationship Between the Morphological Characteristics of the Mitral Valve and Suitability for the MitraClip Procedure

<table>
<thead>
<tr>
<th>Ideal Valve Morphology</th>
<th>Unsuitable Valve Morphology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mitral regurgitation originating from the mid portion of the valve (degenerative or functional etiology)</td>
<td>Perforated mitral leaflets or clefts, lack of primary and secondary chordal support</td>
</tr>
<tr>
<td>Lack of calcification in the grasping area</td>
<td>Severe calcification in the grasping area</td>
</tr>
<tr>
<td>Mitral valve area &gt;4 cm²</td>
<td>Hemodynamically relevant mitral stenosis</td>
</tr>
<tr>
<td>Length of posterior leaflet ≥10 mm</td>
<td>Length of posterior leaflet &lt;7 mm</td>
</tr>
<tr>
<td>Nonrheumatic or endocarditic valve disease</td>
<td>Rheumatic valve disease (restriction in systole and diastole) or endocarditic valve disease</td>
</tr>
<tr>
<td>Flail width &lt;15 mm, flail gap &lt;10 mm</td>
<td>3D TEE gap between leaflets &gt;2 mm</td>
</tr>
<tr>
<td>Sufficient leaflet tissue for mechanical coaptation: coaptation depth &lt;11 mm, coaptation length &gt;2 mm</td>
<td></td>
</tr>
</tbody>
</table>

Adapted with permission from Wunderlich et al. (49).

3D = 3-dimensional; TEE = transesophageal echocardiography.

ROLE OF NOVEL IMAGING TECHNOLOGIES: 3D TEE, INTRACARDIAC ECHOCARDIOGRAPHY, CMR, AND MDCT. In MV disease, 2D TTE and 2D TEE are the standard imaging modalities. Three-dimensional TEE has substantially improved visualization of MV anatomy and function, and the spatial relation of the valve with its surrounding structures (Figure 6). Superior diagnostic accuracy for MV prolapse (with anterior leaflet and commissural involvement) (Figure 6), perforations, and clefts has been reported (53). Three-dimensional TEE improves MR quantification (specifically in eccentric or multiple jets), improves vena contracta width assessment, and permits direct measurement of the anatomic EROA (58). Post-processing precisely delineates the mitral annulus, leaflet lengths, leaflet angles, coaptation length, and tenting area (Figure 7) (59). Three-dimensional TEE may also be useful to guide transcatheter MV repair procedures, such as the MitraClip (Figure 8). Conversely, intracardiac echocardiography is rarely used to guide MitraClip procedures, as acquisition of the different views needed during the procedure can be challenging, but may be useful for other transcatheter MV applications (60).

Advanced imaging techniques, including cardiac magnetic resonance (CMR) and multidetector row computed tomography (MDCT), can provide complementary information in patients with MR. Both CMR and MDCT permit assessment of LA and LV volumes, function, sphericity, and scar tissue. Given its high spatial resolution, MDCT can accurately delineate MV anatomy (Figure 9) (59, 61) and is uniquely useful in demonstrating the size and course of the coronary sinus in relation to the mitral annulus and circumflex coronary artery (Figure 10), which is an important consideration for some transcatheter MV devices (62). CMR may have particular value in the precise quantification of MR (Figure 11) (63); however, like all other
FIGURE 6  Assessment of Mitral Valve Morphology With 3-Dimensional Transesophageal Echocardiography in Primary Mitral Regurgitation

(A) LA "en face" view of the normal mitral valve with anterior and posterior mitral leaflets divided in 3 scallops (A1-P1: lateral; A2-P2: central; A3-P3: medial). (B) Prolapse of the anterior mitral leaflet with flail of the A2 scallop (arrow). (C) Isolated prolapse of the P2 scallop. (D and E) Examples of prolapse of the anterior and posterior commissures (arrows), respectively. The aortic valve (Ao) and the left atrial appendage (LAA) are landmarks for orientation of the LA "en face" view of the mitral valve. Abbreviations as in Figure 5.

FIGURE 7  Measurement of Mitral Leaflets and Annulus Dimensions From 3-Dimensional Transesophageal Echocardiography

Accurate measurements of the mitral leaflets and annulus can be obtained by creating 3-dimensional (3D) reconstructions of the mitral valve from 3D transesophageal echocardiography data. The multiplanar reformation planes are aligned across the mitral annulus (A) providing LV outflow tract, bicommissural, and cross-sectional views of the mitral valve. (B) By tracing the leaflets and determining the mitral annulus landmarks, the 3D models are created, and the post-processing software provides semiautomatic measurements of the mitral leaflets and annulus. Reproduced with permission from Shanks et al. (59). LA = left atrium; LV = left ventricle; other abbreviations as in Figure 5.
imaging modalities, the accuracy of CMR in assessing MR severity is reduced in the setting of atrial fibrillation. In the future it is likely that CMR and MDCT will be increasingly used for pre-procedural assessment and planning of both surgical and transcatheter MR repair and replacement procedures, and post-intervention surveillance.

**CONTROL GROUP THERAPIES**

Selection of the appropriate control group is essential to interpreting the benefit-risk profile of a new device. For randomized MR clinical device trials, 3 control groups may be considered: 1) GDMT alone (with or without a sham control) when GDMT is standard of care; 2) GDMT plus surgical therapy when surgical therapy is standard of care; and 3) GDMT plus an active comparator device if an alternative device is available and is considered a standard of care.

Ensuring the use of appropriate GDMT is a requirement for all patients enrolled in randomized controlled trials and registries. It is the basis upon which the safety and incremental efficacy of procedural therapies may be judged. GDMT in symptomatic patients with severe MR includes treatments for heart failure (for all patients with secondary MR due to LV dysfunction, and for those with primary MR with symptoms of heart failure or volume overload (class D), especially those in whom surgery is not performed or will be delayed) (1). GDMT includes not only the use of specific recommended therapies, but also titration of those therapies to recommended target doses, as tolerated. Optimal GDMT use before study enrollment minimizes the likelihood of major changes in medication dosing during the course of a trial, defined for each drug class as an increase in dose by ≥100% or decrease in dose by ≥50% from baseline. Thus, patients should meet pre-defined GDMT dosing stability criteria before randomization, as the initiation, discontinuation, or titration of therapies after randomization (in either the treatment of control groups) may otherwise seriously confound interpretation of the study results. Although it may not be possible to always prevent

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**FIGURE 8 Transesophageal Echocardiogram Evaluation of MitraClip Implantation in a Patient With Severe Secondary Mitral Regurgitation**

From the midesophageal 4-chamber (A) and bicommissural (B) views, the vena contracta width of the central regurgitant jet can be measured. The 3D LA “en face” view shows lack of coaptation between the anterior and posterior mitral leaflets at the central level (C, arrows). With 3D color Doppler data, the convergence flow can be observed along the coaptation line from the LV view (D, arrows). Three MitraClip devices were successfully implanted with significant reduction of MR as observed from the color Doppler biplane views of the MV (E). On 3D transesophageal echocardiogram full volume of the mitral valve, the LA “en face” view shows a double orifice mitral valve after MitraClip implantation (F). The clips were positioned at the central and anterolateral levels (arrow) leading to a large orifice at the posteromedial level and a small anterolateral orifice (F, asterisks). (G) The color Doppler 3D “en face” view of the mitral valve with 2 residual mild regurgitant jets. Abbreviations as in Figures 5 and 7.
major changes in drug dosing (e.g. after improvement in hemodynamics with effective MR therapy), in general such changes should be minimized to isolate the effect of the randomized treatment, unless they are pre-specified and considered as part of the treatment arm strategy (including, for instance, a prospective approach to reduction of heart failure medications).

Achieving and maintaining maximally tolerated guideline recommended doses of beta-blockers, angiotensin-converting enzyme inhibitors or angiotensin receptor blockers, and mineralocorticoid receptor antagonists is especially important before enrollment in secondary MR trials, as reduction in LV dimensions and LV remodeling with effective medical therapy in heart failure may substantially reduce MR in individual patients, obviating the need for advanced or experimental therapies. MR severity and appropriateness for study eligibility should be reassessed at least 30 days (and preferably 90 days) after any major change in GDMT.

Compliance with optimal GDMT in individual patients is often challenging and should be documented at baseline and throughout the course of the study. Before enrollment, the adequacy of GDMT in individual patients (including drug class, dose, and patient compliance) should be verified by a central eligibility committee to reduce bias associated with subjects changing their behavior under observation post-enrollment (Hawthorne type effect) (see also Role of the Central Eligibility Committee). Intolerance to a drug or drug class or limitation in drug dosing should be on the basis of objective clinical criteria, according to the known adverse effects of specific agents, and must be well-documented in the medical chart and study case report form. Examples include symptomatic hypotension with angiotensin-converting enzyme inhibitors, hyperkalemia with

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**FIGURE 9** Multidetector Row Computed Tomography for Assessment of Mitral Valve Geometry in Secondary Mitral Regurgitation

From the reconstructed short-axis view of the mitral valve, orthogonal planes can be placed across the anterolateral, central, and posteromedial levels of the MV leading to the left ventricular outflow tract view at each level. The angles (A as and P as) and tenting (MVTht) of the mitral leaflets can be measured at the anterolateral (A1-P1), central (A2-P2), and posteromedial (A3-P3). Reproduced with permission from Delgado et al. (61). AC = anterior commissure; Ao = aorta; PC = posterior commissure; RA = right atrium; RVOT = right ventricular outflow tract.
mineralocorticoid receptor antagonists, and symptomatic bradycardia with beta-blockers.

In addition to GDMT for heart failure, appropriate patients should also be treated with biventricular pacing (CRT) and coronary revascularization when substantial ischemia is present, according to contemporary clinical practice guidelines, such as those from the American College of Cardiology Foundation/American Heart Association (1,50) and the European Society of Cardiology/European Association for Cardio-Thoracic Surgery (2). CRT is indicated (Class I) in patients with NYHA functional class II to IV symptoms on GDMT with LVEF $\leq 35\%$, sinus rhythm, a left bundle branch block pattern, and QRS duration $\geq 150$ ms (50). In such patients, CRT may substantially decrease LV dimensions and reduce MR in as many as 50% of patients (64-67). CRT may also be considered (Class IIa) for selected patients with a left bundle branch block pattern and QRS duration $<150$ ms, and for those with a non-left bundle branch block and QRS duration $\geq 150$ ms (Class IIa) (50). Surgical or percutaneous coronary revascularization in patients with substantial ischemia may also, on occasion, reduce secondary MR and should be performed in appropriate patients before study enrollment (68,69). After CRT or coronary revascularization, at least 30 days (and preferably 90 days) should pass, after which TTE or other relevant imaging tests are repeated to assess MR severity and appropriateness for study eligibility. Similar to optimal GDMT use, whether CRT and/or coronary revascularization are indicated and utilized should be verified by the central eligibility committee before study enrollment.

**APPROPRIATE SCENARIOS FOR GDMT ALONE (WITH OR WITHOUT A SHAM) AS THE CONTROL GROUP.** GDMT should be used alone as the comparator (control) group when a surgical comparator is either not indicated (i.e., is not standard of care) or is contraindicated due to high surgical risk, and no other active comparator exists. Examples of this scenario are seen in recent studies of TAVR for critical aortic stenosis in extreme surgical risk patients (70) and from a single arm registry of the MitraClip for primary MR in prohibitive surgical risk patients (35). Another example comes from the ongoing COAPT (Cardiovascular Outcomes Assessment of the MitraClip Percutaneous Therapy for Heart Failure Patients with Functional Mitral Regurgitation) trial of the MitraClip for severe secondary MR in chronic heart failure patients (NCT01626079). Although some practice variability exists in this setting, GDMT (rather than MV surgery) is considered the default therapy for most patients with secondary MR, a conclusion supported by both the current United States and European guidelines (1,2). Thus, the control group in COAPT is GDMT alone for patients in whom MV surgery is not considered appropriate after comprehensive individualized evaluation by the local heart team (see also the subsection Role of the Heart Team).

For patients randomized to the control group, a sham control procedure, in which an invasive procedure is performed but the device is not implanted, should be strongly considered when feasible. Although the implanting physician cannot be blinded, use of a sham control minimizes bias by facilitating blinding of study patients as well as the clinicians and investigators responsible for follow-up study assessments. There are now several notable examples in which favorable results from unblinded studies were not supported by sham-controlled randomized trials, including studies of percutaneous myocardial laser revascularization for refractory angina (24), closure of patent foramen ovale for migraines (25), and renal denervation for hypertension (26). The major limitations to the use of sham controls are: 1) ethical
concerns (e.g., risk of harm with no chance of benefit); and 2) difficulties in maintaining the blind. The nature of the sham control will vary according to the control procedure, and should be selected to maximize the goal of maintaining the blind while minimizing patient risk. For example, for procedures in which the experimental procedure requires a trans-septal puncture, the sham control procedure may include femoral venous access and right heart catheterization. Conversely, a sham control may not be possible for an invasive procedure such as apical insertion of a transcatheter mitral valve. Use of a sham control may be less critical if the primary endpoint of the trial is mortality, although even in this case bias in an open-label study may differentially affect medical compliance and crossover to other therapies.

APPROPRIATE SCENARIOS FOR SURGICAL THERAPY AS THE CONTROL GROUP. Surgical therapy (on a background of GDMT) should be considered as the control group when surgical therapy is the standard of care and patients are acceptable surgical candidates. This is the situation for most patients with primary MR who are not considered to be at very high operative risk.

APPROPRIATE SCENARIOS FOR AN ACTIVE COMPARATOR DEVICE AS THE CONTROL GROUP. An active comparator device (on a background of GDMT) may be considered as the control group when another approved device is indicated for use in the population being studied. For example, in the United States, for symptomatic patients with severe primary MR at prohibitive surgical risk (defined by the FDA as an STS score for 30-day mortality of \( \geq 8 \) [replacement calculator] or \( \geq 6 \) [repair calculator] or the presence of 1 or more high-risk features that, in the opinion of an experienced MV surgeon, otherwise precludes surgery), the MitraClip might currently serve as an active comparator for either a randomized trial or single-arm registry, assuming appropriate MV anatomy. Specific

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**FIGURE 11** Quantification of Mitral Regurgitant Volume With 3-Dimensional Velocity Encoded Cardiac Magnetic Resonance

From a 3-dimensional (3D) volume of the heart, the volume of interest is positioned at the atrioventricular level including the systolic excursion of the mitral annulus. The volume is then reformatted in 2-chamber (2CH) and 4-chamber (4CH) views, and the transmitral flow is quantified from the 3D velocity vector field. During diastole, the mitral inflow is acquired, and in systole, the regurgitant flow can be identified. Through-plane motion correction is performed from the longitudinal velocity measured in the lateral wall (green regions of interest). The regurgitant volume is obtained by calculating the Riemann sum of the backward flow during systole in the flow graph. Reproduced with permission from Ajmone Marsan et al. (63).
recommendations for control groups in MR trials appear in Table 6.

**INCLUSION AND EXCLUSION CRITERIA**

Table 7 details numerous considerations for inclusion and exclusion criteria for investigational MR trials that may be used as a framework. Although each trial will need to tailor these criteria to the specific device and patient population being studied, general principles may be applied when selecting patients with primary and secondary MR for enrollment in MV trials.

**RISK SCORES AND SURGICAL CANDIDACY.** A major decision point that must be reached early is whether the patient is an acceptable surgical candidate. Several risk scores are in widespread use to determine short-term morbidity and mortality after cardiac surgery that account for patient comorbidities and ventricular function. General recommendations for the use of risk scores and assessments of comorbidities for patients undergoing TAVR have been recently reviewed in the Valve Academic Research Consortium-2 consensus document (17). MVARC recommends adoption of a similar approach to integration of risk scores and comorbidities for studies of devices treating MR, in particular with regard to the classification of surgical risk related both to prognosis and selection of the appropriate control group.

The STS score and EuroSCORE II are currently most commonly recommended for this purpose (22,23). The STS score provides separate scores for surgical MV repair and MV replacement, and is recommended for use in clinical trials. Conventionally, very high or “prohibitive” surgical risk is defined by an estimated surgical 30-day mortality of ≥8% using the STS replacement calculator or ≥6% using the STS repair calculator; however, such scores, although having good discrimination, have relatively poor calibration and therefore limited accuracy in identifying extreme risk patients (71). Moreover, few patients with prohibitive risk for surgical treatment of MR were included in the cohorts used to develop and validate the STS and EuroSCORE II.

Of note, the STS and EuroSCORE II were developed from outcomes in patients who actually underwent surgery, whereas transcatheter devices for MR may warrant evaluation in patients too sick for surgery, who are not represented by these scoring systems. Thus, similar to the Valve Academic Research Consortium-2 recommendations, other variables that are not captured in these scores should also be considered when deciding whether a patient is at excessive risk for surgery, including frailty, major organ system compromise (e.g., cirrhosis), and procedure-specific impediments (Table 8) (1). Additional factors that may preclude surgery include severe mitral annular calcification, the presence of a hostile chest (e.g., prior mediastinal radiation or chest malformation), patent left internal mammary artery bypass graft crossing the midline, prior tracheotomy, and severe pulmonary hypertension with or without right ventricular dysfunction.

A specific issue unique to MV therapeutics refers to use of the STS repair versus replacement calculator to determine surgical risk. This is an important consideration, as the replacement calculator yields greater predicted perioperative mortality. For studies of primary MR in nonprohibitive risk patients in whom surgical MV repair is generally the standard of care, it is appropriate to use the STS mitral repair calculator to determine surgical risk. Conversely, for studies of secondary MR, MV surgery is not generally considered the standard of care, and GDMT (±CRT as appropriate) is the mainstay therapy. In secondary MR patients who are operated on, MV repair has not been proven superior to MV replacement (34); most patients will be of at least moderate surgical risk given their underlying cardiomyopathy; and most high-risk patients with secondary MR who are operated on currently receive MV replacement rather than repair (at least in the United States) (72). It is, therefore, reasonable to use the STS mitral replacement calculator to determine surgical risk for studies of devices for secondary MR. Additional considerations regarding the choice of the appropriate surgical risk

**Table 6 Recommended Control Groups for Transcatheter Device Trials in Patients With Mitral Regurgitation**

<table>
<thead>
<tr>
<th>Control Group</th>
<th>Acceptable surgical risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary mitral regurgitation</td>
<td>Mitral valve surgery (repair preferable to replacement) + GDMT (if heart failure or left ventricular dysfunction present)</td>
</tr>
<tr>
<td>High surgical risk†</td>
<td>GDMT† or MitraClip</td>
</tr>
<tr>
<td>Secondary mitral regurgitation</td>
<td>Acceptable surgical risk‡</td>
</tr>
<tr>
<td>High surgical risk†‡</td>
<td>GDMT‡</td>
</tr>
</tbody>
</table>

*Maximally tolerated doses of recommended medications for heart failure. Appropriate patients also should have been treated with CRT and/or coronary revascularization before study enrollment. The definition of high surgical risk may vary according to national standards of care. In the United States, per current U.S. Food and Drug Administration guidelines, patients with primary mitral regurgitation should be determined to be at “prohibitive surgical risk” for GDMT or approved transcatheter devices to be considered as an acceptable control group in regulatory trials. Patients for whom the local standard of care for secondary mitral regurgitation is not surgical mitral valve repair or replacement. Mitral valve repair or replacement might also be a suitable control group for selected patients in whom the local standard of care for secondary mitral regurgitation is mitral valve surgery, depending on the experimental device characteristics (e.g., for studies of transcatheter mitral valve replacement). GDMT = guideline-directed medical therapy.
TABLE 7  Recommended Major Inclusion and Exclusion Criteria for Transcatheter Device Trials in Patients with Mitral Regurgitation

<table>
<thead>
<tr>
<th><strong>Inclusion Criteria</strong></th>
<th><strong>Exclusion Criteria</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Age ≥18 yrs</td>
<td>Life expectancy &lt; 1 yr due to noncardiac conditions</td>
</tr>
<tr>
<td>Degree of MR: Severe (or 3+ and 4+)*</td>
<td>NYHA functional class IVb or ACC/AHA stage D heart failure</td>
</tr>
<tr>
<td>LVEF ≥ 20% (primary MR) or ≥ 60% (secondary MR)t</td>
<td>Hypotension (systolic pressure &lt; 90 mm Hg) or requirement for inotropic support or mechanical hemodynamic support</td>
</tr>
<tr>
<td>Symptom status: NYHA functional class II to IVa</td>
<td>Hypertrophic cardiomyopathy, restrictive cardiomyopathy, constrictive pericarditis, or any other structural heart disease causing heart failure other than dilated cardiomyopathy of either ischemic or nonischemic etiology</td>
</tr>
<tr>
<td>Treatment and compliance with optimal guideline-directed medical therapy for heart failure for at least 30 days (preferably 90 days)</td>
<td>Fixed pulmonary artery systolic pressure &gt; 70 mm Hg</td>
</tr>
<tr>
<td>MR mechanism/anatomy: Appropriate to the design specifications of each device</td>
<td>Physical evidence of right-sided congestive heart failure with echocardiographic evidence of moderate or severe right ventricular dysfunction.</td>
</tr>
<tr>
<td>Surgical risk: Specific STS risk score criteria and/or the presence of high-risk features or comorbidities, depending on the specific trial aims</td>
<td>Mitral valve area ≤ 4.0 cm² (if new device therapy may further decrease the mitral orifice area)</td>
</tr>
<tr>
<td>Completion of required functional tests (e.g., 6-min walk) and/or quality-of-life assessments</td>
<td>Any prior mitral valve surgery or transcatheter mitral valve procedure</td>
</tr>
<tr>
<td></td>
<td>Stroke or transient ischemic event within 30 days before randomization</td>
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<tr>
<td></td>
<td>Modified Rankin Scale ≤ 4 disability</td>
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<td></td>
<td>TAVR within 1 month before randomization</td>
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<td></td>
<td>Severe symptomatic carotid stenosis (&lt; 70% by ultrasound)</td>
</tr>
<tr>
<td></td>
<td>Need for emergent or urgent surgery for any reason or any planned cardiac surgery within the next 12 months</td>
</tr>
<tr>
<td></td>
<td>Absence of CRT with Class I indication criteria for biventricular pacing</td>
</tr>
<tr>
<td></td>
<td>Implant or revision of any rhythm management device (CRT or CRT-D) or implantable cardioverter–defibrillator within 1 month before randomization</td>
</tr>
<tr>
<td></td>
<td>Untreated clinically significant coronary artery disease requiring revascularization</td>
</tr>
<tr>
<td></td>
<td>Any percutaneous cardiovascular intervention, cardiovascular surgery, or carotid surgery within 30 days</td>
</tr>
<tr>
<td></td>
<td>Tricuspid valve disease requiring surgery or severe tricuspid regurgitation</td>
</tr>
<tr>
<td></td>
<td>Aortic valve disease requiring surgery</td>
</tr>
<tr>
<td></td>
<td>Need for any cardiovascular surgery (other than for MV disease)</td>
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<tr>
<td></td>
<td>Echocardiographic evidence of intracardiac mass, thrombus, or vegetation</td>
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<tr>
<td></td>
<td>Active endocarditis</td>
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<tr>
<td></td>
<td>Active infections requiring current antibiotic therapy</td>
</tr>
<tr>
<td></td>
<td>Subjects in whom transesophageal echocardiography is contraindicated or high risk</td>
</tr>
<tr>
<td></td>
<td>Any condition making it unlikely the patient will be able to complete all protocol procedures (including compliance with guideline-directed medical therapy) and follow-up visits</td>
</tr>
<tr>
<td></td>
<td>Patient (or legal guardian) unable or unwilling to provide written, informed consent before study enrollment</td>
</tr>
</tbody>
</table>

*Ideally as assessed by an independent echocardiographic core laboratory. Different quantitative criteria may apply for primary and secondary MR. See Assessment of Mitral Regurgitation: Role of Noninvasive Imaging. 1As a starting point for consideration. The upper limit of LVEF should be selected to ensure inclusion of patients with true secondary MR due to LV dysfunction. By unloading the LV, severe MR increases the LVEF, and LVEF > 60% is consistent with LV dysfunction; however, lowering the upper limit of the LVEF range (e.g., to ≤ 50%) may be considered to increase specificity. Similarly, the lower level of LVEF should be selected to ensure exclusion of patients who might not be capable of benefiting from MR reduction. An acute increase in afterload by reducing or eliminating MR may also (rarely) result in hemodynamic compromise in the early post-intervention period. In general, a lower limit LVEF of 20% is recommended. Lower and upper limits for LV dimensions should also be considered on the basis of the specific device being tested. In the case of secondary MR, if patients with both ischemic and nonischemic dilated cardiomyopathy are enrolled, randomization should be stratified by this variable. 4As a starting point for consideration. Patients should be symptomatic, and most patients should be ambulatory (able to complete a 6-min walk test). Dedicated trials, however, may be designed for asymptomatic or end-stage patients. 5Assessed by echocardiography or right heart catheterization, unless active vasodilator therapy in the catheterization laboratory is able to reduce the pulmonary-vascular resistance to < 3 Wood Units or between 3 and 4.5 Wood Units, with v-wave less than twice the mean of the pulmonary capillary wedge pressure. |

ACC = American College of Cardiology; AHA = American Heart Association; CRT = cardiac resynchronization therapy; CRT-D = cardiac resynchronization therapy defibrillator; LVEF = left ventricular ejection fraction; NYHA = New York Heart Association; STS = Society of Thoracic Surgeons; TAVR = transcatheter aortic valve; UNOS = United Network for Organ Sharing.

ROLE OF THE HEART TEAM. The standard of care for any individual patient, including assessment of surgical candidacy, appropriate use of GDMT, and potential clinical trial eligibility, should be determined by a multidisciplinary heart team consisting of local experts experienced in the care of patients with MV disease (1,17). At a minimum, the heart team should include a heart failure/valve cardiologist, an interventional cardiologist skilled in the relevant access and device implantation procedures, an MV cardiac surgeon, and an imaging specialist. Depending on the specific trial, additional members of the heart team might also include an electrophysiologist, a stroke neurologist, an anesthesiologist, and a geriatrician. Other health care professionals, such as pharmacists and behavioral specialists, may also provide needed expertise to the heart team. Each member of the heart team (other than the echocardiographer) should meet and examine the patient, after which appropriate decisions regarding clinical trial eligibility and surgical risk should be reached by
TABLE 8  Risk Assessment in Valvular Heart Disease, Combining Society of Thoracic Surgery Risk Estimates, Frailty, Major Organ System Dysfunction, and Procedure-Specific Impediments for Intervention

<table>
<thead>
<tr>
<th></th>
<th>Low Risk (ALL Criteria in This Column Must Be Present)</th>
<th>Intermediate Risk (At Least 1 Criterion in This Column Must Be Present)</th>
<th>High Risk (At Least 1 Criterion in This Column Must Be Present)</th>
<th>Prohibitive Risk (Any 1 Criterion in This Column Must Be Present)</th>
</tr>
</thead>
<tbody>
<tr>
<td>STS PROM*</td>
<td>&lt;4%</td>
<td>4%-8%</td>
<td>&gt;8%</td>
<td>Predicted risk with surgery of death or major morbidity (all-cause)</td>
</tr>
<tr>
<td>Frailty‡</td>
<td>None</td>
<td>1 index (mild)</td>
<td>≥2 indexes (moderate to severe)</td>
<td>≥50% at 1 yr</td>
</tr>
<tr>
<td>Major organ system compromise not to be improved post-operatively§</td>
<td>None</td>
<td>1 organ system</td>
<td>No more than 2 organ systems</td>
<td>≥3 organ systems</td>
</tr>
<tr>
<td>Procedure-specific impediment</td>
<td>$\text{Possible procedure-specific impediment}$</td>
<td>$\text{Possible procedure-specific impediment}$</td>
<td>$\text{Severe procedure-specific impediment}$</td>
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</table>

*Use of the STS predicted risk of mortality (PROM) to predict risk in a given institution with reasonable reliability is appropriate only if institutional outcomes are within 1 SD of STS average observed/expected ratio for the procedure in question. Seven frailty indexes: Katz Activities of Daily Living (independence in feeding, bathing, dressing, transferring, toileting and urinary continence) and independence in ambulation (no walking aid or assist required for 5-m walk in <6 s). Other scoring systems can be applied to calculate no, mild, or moderate-to-severe frailty. Examples of major organ system compromise: Cardiac: severe LV systolic or diastolic dysfunction or RV dysfunction, or fixed pulmonary hypertension; CKD stage 3 or worse; pulmonary dysfunction with FEV1 <50% or DLCO <50% of predicted; ENS dysfunction: dementia, Alzheimer’s disease, Parkinson’s disease, or CVA with persistent physical limitation; GI dysfuncion: Crohn’s disease, ulcerative colitis, nutritional impairment, or serum albumin <3.0; cancer: active malignancy, and liver: any history of cirrhosis, variceal bleeding, or elevated INR in the absence of VKAs therapy. Examples: transthoracic present, heavily calcified ascending aorta, chest malformation, arterial coronary graft adherence to posterior chest wall, or radiation damage. Adapted with permission from Nishimura et al (1).

†Seven frailty indexes: Katz Activities of Daily Living (independence in feeding, bathing, dressing, transferring, toileting and urinary continence) and independence in ambulation (no walking aid or assist required for 5-m walk in <6 s). Other scoring systems can be applied to calculate no, mild, or moderate-to-severe frailty. Examples of major organ system compromise: Cardiac: severe LV systolic or diastolic dysfunction or RV dysfunction, or fixed pulmonary hypertension; CKD stage 3 or worse; pulmonary dysfunction with FEV1 <50% or DLCO <50% of predicted; ENS dysfunction: dementia, Alzheimer’s disease, Parkinson’s disease, or CVA with persistent physical limitation; GI dysfuncion: Crohn’s disease, ulcerative colitis, nutritional impairment, or serum albumin <3.0; cancer: active malignancy, and liver: any history of cirrhosis, variceal bleeding, or elevated INR in the absence of VKAs therapy. Examples: transthoracic present, heavily calcified ascending aorta, chest malformation, arterial coronary graft adherence to posterior chest wall, or radiation damage. Adapted with permission from Nishimura et al (1).

‡Examples of major organ system compromise: Cardiac: severe LV systolic or diastolic dysfunction or RV dysfunction, or fixed pulmonary hypertension; CKD stage 3 or worse; pulmonary dysfunction with FEV1 <50% or DLCO <50% of predicted; ENS dysfunction: dementia, Alzheimer’s disease, Parkinson’s disease, or CVA with persistent physical limitation; GI dysfunction: Crohn’s disease, ulcerative colitis, nutritional impairment, or serum albumin <3.0; cancer: active malignancy, and liver: any history of cirrhosis, variceal bleeding, or elevated INR in the absence of VKAs therapy. Examples: transthoracic present, heavily calcified ascending aorta, chest malformation, arterial coronary graft adherence to posterior chest wall, or radiation damage. Adapted with permission from Nishimura et al (1).

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consensus during an in-person meeting of the heart team.

The local heart team assessment of operative risk should supersede any single risk score in determining patient eligibility for surgery. Team decision-making should integrate clinical risk scores with other known prognostic variables, including assessment of frailty. During the consideration of surgical eligibility, anticipation of individual expected improvement in symptoms, quality-of-life, and functional status as well as survival must be considered. Importantly, the heart team tailors adjustment of the decision-making process according to local expertise and standards of care (73). Thus, a patient who is considered to be very high risk for MV surgery at 1 institution may appropriately be considered to be at low or intermediate surgical risk at a different center. Clinical trials can accommodate such systematic site-based variability by stratification at the time of randomization on the basis of risk assessment by objective scores or the central eligibility committee. Finally, in clinical practice, patient preferences (shared decision-making) play an important role, and arbitrary age- and risk-score-based cutoffs are no longer the dominant basis for treatment selection. For trials leading to regulatory approval or indication expansion, the local heart team determination of surgical risk and eligibility supersedes other considerations, thus ensuring enrollment of a clinically appropriate control group and minimizing crossovers during trial conduct.

FRAILTY. Assessment of patient frailty deserves special emphasis. The existence of frailty in an elderly population is an important parameter for risk stratification before major cardiovascular interventions and has demonstrated substantial prognostic capability (74-78). Frailty is a geriatric syndrome that reflects a state of decreased physiological reserve and vulnerability to stressors, and is characterized by a progressive decline in muscle mass and strength (74). Multiple frailty criteria and scales have been proposed (74,79), although the single best assessment tool remains uncertain. Most experts agree that the combination of 5-m gait speed, grip strength, unintentional weight loss, inactivity, and exhaustion represent the most validated frailty measurements (74). Disability, defined as the inability or dependency to carry out activities of daily living and/or managing one’s medications or finances (instrumental activities of daily living), is also an essential part of the initial geriatric evaluation. Assessment of baseline and post-procedure cognitive impairment with clinically established scales such as the Mini-Mental State Examination, the modified Telephone Interview of Cognitive Status, and/or the Clinical Dementia Rating Scale should also be considered when evaluating the utility and cost-effectiveness of invasive procedures among an elderly population (80). Involvement of experts in neurocognitive dysfunction (e.g., geriatricians, neurologists) with serial evaluations pre- and post-procedure is essential for meaningful appraisal; however, it should be acknowledged that evaluation and collection of frailty parameters can be time consuming and resource intensive. Further research is required to determine the extent to which frailty assessments should be a routine part of MV clinical trials and registries.
PRINCIPAL AND SECONDARY ENDPOINTS

GENERAL CONSIDERATIONS. Selection of the primary and major secondary clinical endpoints should afford an assessment of new technologies that is meaningful both for regulatory considerations and to guide clinical use. Such endpoints should give a robust determination of the benefit-risk balance afforded to patients receiving the therapy relative to other therapeutic options. Separate pre-specified primary powered safety and effectiveness endpoints are recommended for most trials of new transcatheter MV therapies, and the use of a single composite clinical safety and effectiveness endpoint, especially when the individual components of safety and efficacy may move in opposite directions, is not recommended.

Although a single primary endpoint (or set of endpoints) cannot be pre-determined in this document for all possible clinical trials of MR devices and therapies, general principles may be proposed:

- Major endpoints should address both the safety and effectiveness of the proposed new device. In general, separate safety and effectiveness measures are desirable as coprimary endpoints, and both safety and effectiveness hypotheses should typically be met to declare trial/device success. However, the FDA’s decisions regarding device regulatory approval are ultimately dependent on a benefit-risk determination that also takes into account disease severity, therapeutic options, and unmet clinical needs for life-threatening and life-limiting conditions (20).

- The primary effectiveness endpoint should be a relevant clinical outcome for the population studied, rather than simply a technical or surrogate measure of success. However, continued evidence of device success at the time of primary effectiveness endpoint assessment should be present to support the determination that the observed benefit was due to the device intervention.

- Similarly, meaningful secondary endpoints assessing pathophysiological mechanisms should be measured that are consistent with and linked to meaningful clinical outcomes (e.g., reduction in MR resulting in decreased LA and LV volumes, improved LVEF, and reduced PAP).

- Additional secondary endpoints may include functional measures (e.g., exercise performance as measured by the 6-min walk test or cardiopulmonary exercise testing) (81,82) and patient-reported quality-of-life outcomes such as the Short-Form Health Survey-12 or -36 scales, Minnesota Living with Heart Failure Questionnaire, Rose dyspnea scale, the EuroQol instrument, or the Kansas City Cardiomyopathy Questionnaire (83–87). Efforts to avoid bias in such determinations, such as patient and assessor blinding, should be incorporated into study designs when feasible.

- The primary and secondary safety endpoints should assess procedural and/or device-related complications and incorporate any adverse impact of the intervention on the disease state, future treatments, and prognosis.

- The primary and secondary endpoints should be selected such that meeting these endpoints will demonstrate reasonable assurance of safety and effectiveness and a favorable benefit-risk profile.

- All endpoints must be well defined such that they can be subjected to statistical analysis, and clinical endpoints should rely on the use of independent adjudication processes, blinded when possible. It is acknowledged, however, that blinding the central adjudication committee may be difficult in trials of MR therapies in which the control and experimental treatments vary so greatly, especially if the committee must ascribe the extent to which adverse events are study device-related. This latter issue may be overcome by adjudicating in 2 discrete steps: 1) whether or not an event (stroke, myocardial infarction, and so on) occurred; and then 2) whether or not the event was procedure or device related.

- Quantitative measures, such as imaging parameters and electrocardiographic changes, should be assessed by independent core laboratories that are blinded to treatment assignment when possible.

- Endpoints should be measured at relevant intervals that are appropriate for demonstrating safety and effectiveness, and the analysis should incorporate and pre-specify both early and late endpoints according to previously proposed standards, including acute intraprocedural events as defined in part 2 of this document and in earlier consensus documents (16,17).

- If composite endpoints are necessary to afford reasonable study size, they should be comprised of important clinical outcomes related to effectiveness and/or device safety that may be observed during the relevant period of observation. The individual components of the composite endpoint should share a common pathophysiology or represent specific major complications of device therapy, and should be expected to trend in the same direction. Major and minor events should be clearly distinguished to avoid grouping outcomes of variable clinical significance.
• The statistical analysis of these endpoints should conform to commonly accepted principles, such as accounting for competing risks and multiplicity (see also Statistical Considerations).

Selection of appropriate primary and secondary endpoints to assess device therapies for MR is especially challenging, because of a complicated matrix that includes the underlying risk and comorbidities of the target population, the specific pathogenic mechanisms of MV dysfunction (e.g., primary vs. secondary and ischemic vs. nonischemic etiologies), and whether options for therapy include MV repair or replacement surgery, coronary revascularization, and ventricular resynchronization therapy, in addition to GDMT. For example, whether mortality is the optimal stand-alone primary endpoint may depend on the expected survival rate of the target population with currently available treatment options. For patient cohorts in whom short- or intermediate-term mortality is low, other measures, such as outcomes related to heart failure and functional capacity, may be more clinically meaningful during the course of a clinical trial observation if associated with continued successful device performance.

For studies in which MV surgery is the control group (e.g., for primary MR in acceptable surgical candidates), major endpoints must assess the safety of the new device relative to the complications of surgery, and demonstrating superiority in safety with noninferiority in efficacy (within a reasonable margin reflecting therapeutic interchangeability) is a reasonable goal. Conversely, if the control group is GDMT with or without coronary revascularization and/or CRT as appropriate (e.g., for secondary MR in heart failure), device safety endpoints must be weighed against superiority measures of clinical efficacy that would justify the associated procedural and device-related risks. Whether or not pre-specified endpoints are met, FDA approval is based upon the totality of the data when considered as part of a thorough benefit-risk determination (20).

**PRIMARY ENDPOINTS.** Clinically meaningful effectiveness measures to be considered for MR device therapies are presented in Table 9. All-cause mortality should be incorporated into the primary efficacy endpoint (as either a standalone measure or as part of a composite) if there is a reasonable expectation that MR reduction might improve survival (e.g., for primary MR). For clinical trials in which the mortality rate during the time-course of observation is expected to be high, all-cause mortality as a pre-specified and adequately powered standalone primary efficacy endpoint should strongly be considered, as a significant improvement in survival is likely to support a favorable benefit-risk determination even if other device-related safety issues emerge. It is also the endpoint least affected by lack of blinding. Otherwise, all-cause or cardiac mortality may be part of a carefully constructed primary composite clinical effectiveness endpoint in which the pathophysiology between the components is shared and all events are considered clinically important. In this regard, after all-cause mortality, heart failure-related outcomes may be the best clinical measure of effectiveness to use in trials of MR therapies. As outlined in part 2 of this document, specific criteria defining hospitalization or hospitalization equivalents for heart failure may be crafted to allow independent adjudication of this event. Hospitalization for heart failure may serve as a primary standalone endpoint for conditions in which MR reduction is anticipated to improve quality-of-life, but may not necessarily improve survival (e.g., secondary MR in patients with severe LV dysfunction).

Whether functional measures (e.g., 6-min walk distance) or quality-of-life indexes are sufficiently clinically important and robust to warrant incorporation into a primary effectiveness endpoint is controversial. This consideration is especially relevant in unblinded trials, in which placebo and Hawthorne effects as well as assessment bias may make interpretation of these measures difficult. Proposed
measures of functional performance are presented in Table 9. Such measures might be regarded as having intermediate value as reviewed by Temple (88), because they are meaningful to patients and may eventually result in a favorable effect on survival. If functional measures or quality-of-life outcomes are considered as a primary endpoint (either standalone or as part of a composite endpoint), all possible efforts for blinding should be used (including assessment of the success of the blinding procedures), and the data should demonstrate continued evidence of device performance at the time of assessment and reasonable confidence of lack of harm, including mortality.

The primary safety endpoint is usually a composite endpoint specific to the device and underlying cardiac condition, and should incorporate the need for unplanned MV surgery (or reoperation) due to progressive or recurrent MR or device-related complications.

As an example, these principles have been incorporated into the design of the COAPT trial, an ongoing prospective, multicenter randomized trial performed under an FDA Investigational Device Exemption in which the MitraClip is being compared to GDMT in patients with symptomatic severe secondary MR in whom surgery is not considered appropriate after local heart team evaluation. The primary effectiveness endpoint is hospitalization for heart failure during follow-up (measured by the Andersen-Gill test to take into account the number of heart failure hospitalizations), powered to demonstrate superiority of the MitraClip. The primary safety endpoint is the composite of single leaflet device attachments, device embolization, endocarditis requiring surgery, core laboratory-confirmed mitral stenosis requiring surgery, LV assist device implant, heart transplant, or any device-related complications requiring nonelective cardiovascular surgery at 12 months, to a pre-specified performance goal.

**SECONDARY ENDPOINTS.** Secondary endpoints should include the individual components of the primary endpoint if a composite was used, as well as other measures of effectiveness that were not a part of the primary endpoint, including functional, symptom, and quality-of-life assessments; imaging-related measures of MR reduction and cardiac structure and performance; and major and minor safety outcomes (Table 10). Secondary endpoints of interest that may be affected by the intervention should be pre-specified, but the study may or may not be adequately powered to demonstrate statistical significance for such endpoints even if differences truly exist. Powered secondary endpoints with pre-specified statistical hypotheses are necessary to make labeling claims for approved medical devices in the United States. If not powered, secondary endpoints are considered hypothesis generating, even if pre-specified. Nonetheless, secondary mechanistic endpoints may provide valuable guidance for new device designs or iterations, especially in this early era of MR device development.

Endpoints should be classified according to device- and procedure-relatedness and timing of occurrence as previously proposed: acute, within 24 h; early, after 24 h but within 30 days; intermediate, after 30 days but within 1 year; late, between 1 and 5 years; and very late, after 5 years (17). For secondary outcome measures (as for the primary endpoints), safety should be assessed separately from effectiveness, except possibly for all-cause mortality and stroke, endpoints that reflect both safety and effectiveness of an intervention. Secondary safety endpoints should further evaluate procedural and/or device-related complications and assess any adverse effects of the intervention on the disease state, future treatments, and prognosis (e.g., whether device failure impedes the likelihood to perform successful surgical MV repair). If not already identified as a standalone primary endpoint, all-cause mortality should always be pre-specified as a secondary endpoint (even if not adequately powered) to consider whether a new therapy might result in increased or decreased survival.

Various measures of mechanistic and technical success are important to include as pre-specified secondary outcomes in MR device trials. Although quantitative reduction in MR is the *sine qua non* for treatment effectiveness, and the associated physiological measures (e.g., reduced LA and LV volumes and PAP, improved LVEF) are consistent with successful device performance over time, these surrogate endpoints are insufficient to serve as primary effectiveness endpoints (either standalone or as a component of a composite measure) because they may not be associated with clinically meaningful improvements. Furthermore, small statistically significant differences in continuous measures may not result in clinically measurable benefits. Nevertheless, a high rate of early technical success, coupled with continued device functional performance and a beneficial physiological response over time, should be present to support the biological plausibility of the primary clinical effectiveness endpoint. These measures should, therefore, be evaluated in all MR trials and reported as secondary efficacy outcomes. MR severity over time is also an important measure of durability of the treatment effect and should be assessed at regular intervals throughout the study.
The analysis plan should incorporate achievement of device- and patient-oriented outcomes to assess overall safety and effectiveness as well as benefit-risk. As detailed in part 2 of this document, device-oriented outcomes include technical success with associated mechanistic outcomes and device- and procedure-related safety endpoints. Patient-oriented outcomes include the components of the primary safety and effectiveness endpoints. A hierarchical analysis plan should be pre-specified beginning with assessment of the primary safety and effectiveness endpoints followed by analysis of powered major secondary endpoints assessing functional and mechanistic outcomes, with attention to preserving type I error (alpha) at the 0.05 level. Thereafter, non-powered secondary endpoints are assessed, as well as subgroup analyses to examine consistency, although these exploratory analyses offer lower levels of evidence and are considered hypothesis-generating in most circumstances.

Finally, it should be noted that over time the level of evidence required for serial iterations of MR devices or even novel devices will likely change as experience grows with therapies for treating MR patients.

### ROLE OF THE CENTRAL ELIGIBILITY COMMITTEE

Even with the use of local heart teams and detailed protocol inclusion/exclusion criteria, site-based variability in patient selection for studies of devices treating MR remains a concern. To enhance interpretability, particularly for regulatory trials of novel MR devices, it is strongly recommended that each patient be presented to a central eligibility committee for evaluation of patient appropriateness for enrollment. The central eligibility committee serves multiple important functions (Table 12). The members of the central eligibility committee for MR trials should include at a minimum a moderator, a heart failure specialist, and an experienced MV surgeon. Each patient should be presented to the committee by

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**TABLE 10 Major Safety, Technical, and Mechanistic Endpoints in Mitral Regurgitation Trials**

<table>
<thead>
<tr>
<th>Endpoint Description</th>
<th>Details</th>
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<tbody>
<tr>
<td>Major safety endpoints</td>
<td>Device or procedure-related adverse events (specific to each device and procedure)</td>
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<tr>
<td></td>
<td>Major bleeding complications (transfusion reported separately)</td>
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<td>Major vascular complications</td>
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<td>Pulmonary complications (device or procedure-related)</td>
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<td>Stroke and other cerebrovascular events (assessed by a stroke neurologist and CT/CMR imaging; disabling and nondisabling; change in modified Rankin score)</td>
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<td>Myocardial infarction</td>
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<td>Acute kidney injury or progression of chronic kidney disease (dialysis reported separately)</td>
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<td>New onset atrial fibrillation</td>
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<td></td>
<td>Unplanned mitral valve surgery due to device/procedure failure or malfunction</td>
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<td></td>
<td>Requirement for valve replacement versus repair</td>
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<td></td>
<td>Unplanned cardiac surgery for any cause</td>
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<td></td>
<td>Requirement/insertion of an implantable cardiac defibrillator</td>
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<td></td>
<td>Requirement/insertion of biventricular pacemaker for cardiac resynchronization therapy</td>
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<td></td>
<td>Device failure resulting in the inability to perform successful surgical mitral valve repair</td>
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</table>

**Technical success**

- Device success (specific definition)
- Implant rate
- Device time and procedure duration
- Ionizing radiation exposure
- Procedural success (specific definition)

**Mechanistic endpoints**

- Imaging measures
  - Mitral regurgitation severity (integrated assessment; see text and Tables 2 and 3)
  - Mitral valve area and mean gradient
  - Left atrial and pulmonary artery pressures
  - End-systolic dimension and volume
  - End-diastolic dimension and volume
  - Left ventricular sphericity
  - Left ventricular ejection fraction
  - Left atrial dimension and volume
  - Right ventricular pressures, dimension, volume, and ejection fraction
  - BNP and/or NT-pro BNP levels

*Absolute levels and incremental change from baseline.

- BNP = B-type natriuretic peptide; CMR = cardiac magnetic resonance; CT = computed tomography; NT-proBNP = N-terminal pro-B-type natriuretic peptide.

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**TABLE 11 Timing of Endpoint Assessment (Follow-Up Intervals)**

- Acute (during procedure or within 24 h)
- Procedural (30 days post-procedure or until discharge from hospital or acute care facility)
- 90 days
- 6 months
- 12 months
- Annual (for a minimum of 5 yrs)
the local principal investigator and, depending on the specific requirements for the trial, a cardiologist with expertise in valvular heart disease, a heart failure specialist, and/or an MV cardiac surgeon. The MV surgeon plays a central role in all MR randomized trials, whether the control arm is surgery (as for most trials of primary MR) or GDMT (as for most trials of secondary MR). Before the committee meeting, a central echocardiographic core laboratory should have reviewed the qualifying TTE (±TEE) to ensure that the MR meets severity criteria and, depending on the trial, to confirm that other eligibility criteria are met (e.g., LV volumes, LVEF, MR etiology and anatomy, absence of mitral stenosis, and so on). Depending on the specific device, the central echocardiographic core laboratory may also be asked to determine whether appropriate anatomy is present for device eligibility. Although implementation of a central eligibility committee and pre-review by a central echocardiographic core laboratory entail extra time and cost, ensuring that only appropriate patients are enrolled in the clinical trial will substantially increase the power of the study and the likelihood of success.

An important distinction must be made between the roles of the local heart team and the central eligibility committee. Both multidisciplinary groups possess expertise to evaluate surgical risk and clinical trial eligibility. However, the thresholds at which MV surgery might be considered reasonable vary from center to center and surgeon to surgeon, depending on local experiences and volume considerations. The subtleties of the patient’s clinical condition (and surgical risk) are also best assessed by those able to speak to and examine the patient. Thus, the local heart team determines the relative surgical risk and operability of a patient, rather than the central eligibility committee. The MV surgeon (and others) on the central eligibility committee may, however, query the local surgeon as to his/her criteria for operability to ensure, for example, that crossover to surgery will not be considered should the patient be randomized to a nonsurgical therapy (unless permitted by the protocol).

**STATISTICAL CONSIDERATIONS**

**GENERAL CLINICAL TRIAL DESIGN ISSUES.** Most clinical studies of new devices addressing MR will be randomized trials. Crossovers should be discouraged, and in all cases not permitted until the primary endpoint has been reached. Contemporary adaptive designs that expose the fewest patients to an inferior therapy may be applicable to randomized trials with short-term primary endpoints, multiple trial arms, and the ability to switch to a long-term endpoint (89). Single-arm trials using historical comparators may be appropriate when the condition to be examined is infrequent or no adequate comparator exists. Single-arm trials may also be appropriate when a body of published data exists that is sufficient for construction and justification of appropriate performance goals or following development of objective performance criteria. As the field matures, single-arm studies may become appropriate for serial device iterations or studies of similar device designs.

**TRIAL ENDPOINTS AND ANALYSIS.** Endpoints fall into 3 categories: 1) early post-procedure events and measures (e.g., death, stroke, valvular regurgitation); 2) time-related events (e.g., death, stroke); and 3) periodically sampled longitudinal data (e.g., valvular regurgitation, chamber dimensions, NYHA functional class). Early events are compared as odds ratios, early measurements are compared as differences, time-related events are compared as actuarial curves and hazard ratios, and longitudinal data are compared as time-related differences in ensemble averages. Although each trial must carefully evaluate and adopt the statistical methodology most appropriate for its goals, the following general principles may be useful to consider.

<table>
<thead>
<tr>
<th>TABLE 12 The Central Eligibility Committee</th>
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<tr>
<td><strong>Purpose</strong></td>
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<td><strong>Format</strong></td>
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<tr>
<td><strong>Outcome</strong></td>
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*C* Ideally determined by echocardiographic core laboratory review of the qualifying imaging studies before the central eligibility committee meeting.

**CR** = cardiac resynchronization therapy; **ICD** = implantable cardioverter-defibrillator; other abbreviations as in Table 2.
**Nonfatal time-related events.** Nonfatal events can repeat (e.g., stroke, rehospitalization). All occurrences should be analyzed, not just time to first occurrence, using the Nelson (90), Andersen-Gill (91), or other estimators (92). These methods make different assumptions with respect to the independence of events, hazard function after each occurrence, and informativeness of death and other competing risks (93).

**Weighted events.** Although generally not done, consideration may be given to weighing nonfatal events (e.g., by applying the National Institutes of Health stroke scale and considering the duration and cost of rehospitalization), which may be further analyzed as cumulative functions, a common industrial method when considering costs (90,94,95).

**Longitudinal data.** Longitudinal data reflect an endpoint’s state at time of assessment; they are not time-to-event data (96). Examples are drug use (binary), functional status (ordinal), and EROA (continuous). Such endpoints should be analyzed by longitudinal repeated measures methods (96). Ensemble averages across time are subject to informative censoring from events with which they are associated.

**Composite endpoints.** The use of composite endpoints to reduce sample size is a practical convention, but if not carefully constructed, may lead to difficulties in interpretation (97). Typically, each component is equally weighted, although the hazard function for each may be different (e.g., the Andersen-Gill method assumes proportional hazards [91]). However, clinical hierarchy or patient preference for each component may differ. If the components can be hierarchically arrayed, tree-structured gatekeeping tests (98) or pair-wise winner-loser strategies (99) may be used. Family-wise tests of individual components emphasize consistent direction of effect (100).

The most controversial composite endpoints combine disparate component categories. Several groups, such as Finkelstein and Schoenfeld (101), have developed methods that combine time-to-event components with periodic longitudinal assessments. Others have extended this to continuous longitudinal data combined with weighted time-to-event data (102). On the horizon are joint models that account simultaneously for different intensity functions of each event and longitudinal components and their interrelations (103).

**CONCLUSIONS**

In contrast to calcific aortic stenosis, a relatively simple disease with limited etiologies and a straightforward pathophysiology, MR is a more complicated entity, due to the greater complexity of the MV structure and the numerous lesions and mechanisms that may lead to its failure. Central to the understanding of MR is recognizing that MR is indeed 2 disorders: 1 of the valve apparatus itself and 1 of the LV (or LA), which secondarily disrupts normal MV function. Continuing the analogy, developing effective therapies (and surgical approaches) for MR and demonstrating their safety and effectiveness in clinical trials is much more challenging than for aortic stenosis, and requires the intimate collaboration between physician-scientists across numerous disciplines, clinical trialists, statisticians, and industry and regulatory authorities. Although each device trial will entail its own nuanced considerations, adopting the principles espoused in this document as a template for clinical investigation of mitral therapeutics should allow sponsors and investigators to avoid the most common errors that can render interpretation of their findings problematic.

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**KEY WORDS** heart failure, mitral regurgitation, mitral valve, valve intervention, valve surgery (or cardiac surgery)

**APPENDIX** For complete information on the MVARC members and participants, please see the online version of this article.