Mitral Regurgitation Augments Post-Myocardial Infarction Remodeling

Failure of Hypertrophic Compensation

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Objectives

We examined whether mitral regurgitation (MR) augments post-myocardial infarction (MI) remodeling.

Background

MR doubles mortality after MI, but its additive contribution to left ventricular (LV) remodeling is debated and has not been addressed in a controlled fashion.

Methods

Apical MIs were created in 12 sheep, and 6 had an LV-to-left atrial shunt implanted, consistently producing regurgitant fractions of ~30%. The groups were compared at baseline, 1, and 3 months.

Results

Left ventricular end-systolic volume progressively increased by 190% with MR versus 90% without MR (p < 0.02). Pre-load-recruitable stroke work declined by 82 ± 13% versus 25 ± 16% (p < 0.01) with MR, with decreased remote-zone sarcoplasmic reticulum Ca2⁺-ATPase levels (0.56 ± 0.03 vs. 0.76 ± 0.02, p < 0.001), and decreased isolated myocyte contractility. In remote zones, pro-hypertrophic Akt and gp130 were upregulated in both groups at 1 month, but significantly lower and below baseline in the MR group at 3 months. Pro-apoptotic caspase 3 remained high in both groups. Matrix metalloproteinase (MMP)-13 and membrane-type MMP-1 were increased in remote zones of MR versus infarct-only animals at 1 month, then fell below baseline. The MMP tissue inhibitors rose from baseline to 3 months in all animals, rising higher in the MI MR–group border zone.

Conclusions

In this controlled model, moderate MR worsens post-MI remodeling, with reduced contractility. Pro-hypertrophic pathways are initially upregulated but subsequently fall below infarct-only levels and baseline; with sustained caspase 3 elevation, transformation to a failure phenotype occurs. Extracellular matrix turnover increases in MR animals. Therefore, MR can precipitate an earlier onset of dilated heart failure. (J Am Coll Cardiol 2008;51: 476–86) © 2008 by the American College of Cardiology Foundation

Expansion of infarcted tissue begins acutely after myocardial infarction (MI). A more gradual remodeling process, however, also involves the noninfarcted areas (1,2); initially compensatory, this process subsequently becomes mal-adaptive, generating a larger, more spherical, and poorly contracting ventricle (3–7), associated with reduced survival (2,8,9).

Myocardial infarction can also cause mitral regurgitation (MR) by altering ventricular geometry and function (10–12). Such ischemic MR doubles the risk of death after MI (13,14). Severe nonischemic MR has been shown to promote left ventricular (LV) remodeling and reduce survival (15–19). However, whether more typically moderate ischemic MR does so remains controversial. A recent paper maintained that “ischemic MR is a consequence, not a cause, of post-infarction remodeling,” because remodeling persisted despite annuloplasty with apparent MR reduction.

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In patients, ischemic myocardial changes and regurgitation develop concurrently, precluding the ability to separate their effects and analyze their interaction directly (23). This requires a model in which MR can be varied independently in the presence of MI, but without interventions such as infarct patching (24–27) that might themselves influence remodeling. Most experimental studies of post-MI remodeling have used an infero-posterior MI model (27,28); this has the disadvantage of linking the MI-induced remodeling to the development of MR. We therefore tested the incremental contribution of MR in a controlled manner using a model in which the MI by itself does not cause MR, and the MR can be created in a standardized and fixed fashion (29,30).

In such a study, it must be recalled that remodeling is a complex process, involving changes not only in ventricular geometry (3) and in global and regional myocardial function (31), but also in calcium handling, molecular signals promoting hypertrophy, cardiomyocyte viability (32–34), and extracellular matrix turnover (35). Intracellular pathways that promote compensatory hypertrophy also tend, if chronically activated, to drive cells down pathways to cell death by apoptosis and thus to loss of contractile elements, with further deterioration of cardiac function (Fig. 1). According to the importance of protein evidence of remodeling at the molecular, cellular, and whole-heart levels (36). We therefore tested the hypothesis that MR accompanying MI produces greater remodeling than a comparable infarct alone, and may be associated with a different progression of molecular events, with transition from signals promoting hypertrophy to those inducing failure.

**Methods**

**MR model.** A modification of the chronic MR sheep model of Rankin et al. (29) was implemented in conjunction with Luis Guerrero, a surgeon experienced in physiological modeling, using an 8-cm long, 8-mm diameter reinforced Teflon (PTFE) graft (Edwards Lifesciences, Irvine, California) (cross-sectional area of 0.50 cm²) implanted under sterile conditions into the LV base and left atrial (LA) appendage with intramuscular portions stiffened with epoxy resin (Fig. 2). The regurgitant flow was confirmed during each thoracotomy (see the following text) using a transonic flow probe and color Doppler echocardiography imaging. The standardized shunt diameter and length have produced consistent levels of MR (regurgitant fractions of approximately 30%, corresponding to moderate MR [37]). Animals were treated with heparin (3 days) and then oral aspirin.

**Animal studies.** A total of 12 Dorsett hybrid sheep (20 to 30 kg) were loaded for 3 days with amiodarone (200 mg orally twice a day) then anesthetized with thiopentothal sodium (0.5 ml/kg), intubated, and ventilated at 15 ml/kg with a 2% isoflurane-oxygen mixture. All received glycopyrrolate (0.4 mg intravenously) and prophylastic vancomycin (0.5 g intravenously) and amiodarone (150 mg intravenous drip over the course of the operation). Surface electrocardiogram (ECG) was monitored, and a sterile left thoracotomy performed with pericardial incision and creation of a cradle. A high-fidelity micromanometer-tipped catheter (Millar Instruments Inc., Houston, Texas) was placed into the LV. After baseline 2-dimensional (2D) and 3-dimensional (3D) echocardiography imaging, an anterograde MI was produced by ligating the mid- to distal left anterior descending coronary artery, known to produce a substantial MI without MR (38). An immediate 2D echocardiography apical image was performed to confirm that septal wall motion abnormality extended approximately one-third of the way from LV apex to base for standardization. Mitral regurgitation equivalent volume overload was created for 90 days in 6 sheep with an open LV-to-LA shunt (referred to as the infarct + MR group); in the other 6, the shunt was closed (infarct-only group). In addition to analgesia, propranolol, 1 mg intravenously in 2 divided doses, was given for evident stress and sinus tachycardia (>150) upon extubation. Antibiotics (cephalirin, 0.5 g intravenously) and analgesics (buprenorphine, 0.3 mg twice a day) were administered for the next 5 days, and oral amiodarone (200 mg twice a day) for the next 3.
During repeat sterile thoracotomy at day 30, 3D echocardiography was performed to evaluate LV remodeling and function, with directed biopsies of the noninfarcted myocardium near and remote from the border zone using a TruCut needle (Cook, Winston-Salem, North Carolina). At day 90, 3D echocardiography and blood sampling were repeated at thoracotomy, followed by euthanasia and whole-heart pathological and myocyte size and contractility studies that require considerable tissue. The animal studies conformed to National Institutes of Health Guidelines for Animal Research (Guide for the Care and Use of Laboratory Animals, National Research Council, Washington, DC, 1996) and were approved by the institutional animal care committee.

**3D echocardiography and LV function.** Three-dimensional quantitation is necessary because LV shape changes with chronic MR, as well as MI (39). Rotated apical images were obtained at 10° intervals with an epicardial 5-MHz transesophageal echocardiography probe (Sonos 7500, Philips Medical Systems, Andover, Massachusetts), rotated by software and gated to ECG and respiration. Digital images were analyzed on a workstation with custom programs (40). Endocardial surfaces were traced to calculate LV volumes and ejection fraction (EF), and coded for wall motion abnormality to produce a quantitative 3D wall motion map validated against a 36-crystal sonomicrometer array (41). Remodeling was quantified in terms of increasing LV volumes (42). Regurgitant fraction was calculated as transonic MR-equivalent flow divided by LV ejection volume, with cross-correlation by pulsed Doppler time–velocity integral of shunt flow multiplied by shunt cross-sectional area. At thoracotomy, the initial and final pressure–volume loops were obtained using Millar catheters and 4 sterile subendocardial crystals (Sonometrics Corp., London, Ontario, Canada), with an inferior vena cava occlusion to obtain the end-systolic pressure–volume relation for contractility (43) and pre-load–recruitable stroke work (44). Crystals were located at the LV apex and base (between aorta and mitral annulus) and anteriorly and posteriorly along a midventricular short-axis.

Crystals were not placed at day 30 to maximize sterility and survival. dP/dt was obtained by the high-fidelity Millar catheter (Millar Instruments, Houston, Texas).

**Myocyte size and contraction.** Small (5 to 10 g) areas of tissue were removed at euthanasia from the noninfarcted LV and perfused via the corresponding coronary artery for 5 min with low-magnesium Krebs–Henseleit (KH) cardioplegia solution. Myocardial cells were isolated using a well-established method (45,46). The LV sections were perfused with Ca2+-free KH containing collagenase (Worthington Biochem Corp., Lakewood, New Jersey) and hyaluronidase (Sigma, St. Louis, Missouri). Ventricular muscle tissue was minced into small pieces and placed in enzyme buffer I (Sigma) in the presence of trypsin and deoxy-ribonuclease. The tissue was washed in a 3:1 mixture of Ca2+-free KH in Dulbecco’s phosphate-buffered saline (GIBCO, Gaithersburg, Maryland). Cells were loaded with the Ca2+ fluorescent indicator Fura-2 for 15 min in a light-sealed chamber mounted on an inverted microscope. Myocyte length and width were measured offline on 50 consecutive cells using commercial image analysis software to establish proportions of normal short cells versus elongated cells characteristic of eccentric hypertrophy in the failing phenotype. The cells were excited to contract using platinum electrodes in the bath, and a dual excitation spectrofluorometer recorded fluorescence emissions (505 nm from exciting wavelengths 340 nm and 380 nm) to calculate intracellular calcium concentrations. Cell motion was measured by a video-edge motion detector (Crescent Electronics, Sandy, Utah) (45).

**Molecular assays.** We measured levels of several molecular species that determine cell contractility, that modulate cell hypertrophy and death, and that are responsible for extracellular matrix turnover. All protein assays were performed and specimens pooled from the same treatment group and stage.

**Calcium cycling proteins: sarcoplasmic reticulum Ca2+-ATPase (SERCA2a) and phospholamban.** Sarcoplasmic reticulum membranes were obtained from the myocardial biopsy specimens using a sucrose gradient centrifugation method (47). Myocardium was homogenized and centrifuged at 7,700 g for 20 min, and the supernatant filtered and centrifuged at 136,000 g for 60 min. The pellets, resuspended and rehomogenized in 10% sucrose buffer, were centrifuged at 55,000 g in a 10% to 40% sucrose gradient, and the bands between 30% and 40% enriched in sarcoplasmic reticulum were collected, centrifuged at 136,000 g for 60 min, and the pellet resuspended in freezing buffer. Protein was determined by a modified Bradford procedure (using bovine serum albumin for the standard curve). Proteins were separated from the sarcoplasmic reticulum vesicle preparation from tissue frozen in liquid nitrogen, and an immunoblot using monoclonal anti-SERCA2 and phospholamban performed. Proteins levels were normalized against total myocardial protein.

**Pro-hypertrophic and anti-apoptotic cascades.** We measured levels of Akt (protein kinase B) and gp130, which are...
both at their respective levels (cytosol and membrane) important crossroads in pro-hypertrophic signaling; and caspase 3, the final common pathway for intracellular signaling of apoptosis. Western blot analysis was performed on cell lysates from biopsies at baseline and days 30 and 90. Antibodies binding for anti-gp130, anti-phospho Akt, and antiactivated caspase 3 (48) were detected with a peroxidase-conjugated antimouse IgG and chemiluminescence. α-Actin was used as a housekeeping control. Densitometry of the blots was assessed using commercially available software (Photoshop 5, Adobe Systems Inc., San Jose, California).

Modulators of extracellular matrix turnover: matrix metalloproteinases (MMPs). Assays were performed for membrane-type matrix metalloproteinase-1 (MT1-MMP), MMP-13, tissue inhibitor of matrix metalloproteinase (TIMP)-1, and TIMP-4 by immunoblotting using methods previously described (35). Left ventricular myocardial samples were homogenized and concentrated to prevent MMP proteolytic activation and degradation. Trypsin (2.5 μM; 5 min) was used to cleave nascent MMPs to fully activate enzyme moieties and/or facilitate the removal of endogenous inhibitors. Relative TIMP-1, TIMP-4, MT1-MMP, and MMP-13 levels were then measured by immunoblotting. A positive control was included in all immunoblots (MMP-13: CC068; MT1-MMP: CC1042) (Chemicon, Temecula, California).

Statistics. Statistical analysis was performed using a Student t test for continuous variables and a chi-square test for categorical variables (Stat View 5, SAS Institute Inc., Cary, North Carolina). A value of p < 0.05 was considered significant. Multiple comparisons were performed using analysis of variance (for repeated measurement where needed) and a Student-Neumann Keuls post-hoc test. Interobserver and intraobserver variability for 3D echo-measured LV volumes and EF in our laboratory is 3.5%.

Results

Remodeling: LV volumes. Left ventricular end-diastolic volume relative to the acute post-MI value increased by an average of 26% in the infarct-only group versus 130% in those with volume overload (p < 0.02). Left ventricular end-systolic volume progressively increased by 190% with MR-type volume overload versus 90% with MI only (p < 0.02) (Figs. 3 and 4).

Remodeling: LV function. Pre-load–recruitable stroke work from pressure-volume loops declined more in the infarct + MR group by 82 ± 13% versus 25 ± 16% of baseline in the infarct-only group (p < 0.01). Left ventricular dP/dt also declined progressively in the MR group to 52 ± 9% of baseline, but only by 22 ± 12% in the shams (p < 0.04) (Fig. 4). These results were corroborated by reduced contractility of isolated cells with reduced calcium transients and increased diastolic intracellular calcium concentration (Tables 1 and 2). The diastolic relaxation constant tau was more prolonged in the infarct + MR group (78 ± 23 ms vs. 44 ± 13 ms, p < 0.01).

Calcium cycling proteins. As shown in Figure 5, SERCA2a abundance was downregulated in the infarct region and border zone compared with the remote region in both infarct + MR and infarct-only sheep LVs. There was a lower level of SERCA2a in the remote zone of infarct + MR than of infarct-only sheep. Phospholamban was unchanged in all regions. These changes are typical for remodeling myocardium.

Intracellular signaling pathways. Akt, a pro-hypertrophic enzyme (49), was upregulated in the remote zones of infarct + MR and infarct-only animals at 1 month, but was lower in animals with than without MR at 3 months (Fig. 6). In the infarct + MR group at 3 months, Akt also fell below baseline of the same group. The same pattern was observed for the interleukin-6-family receptor subunit gp130 (50). Expression of activated caspase 3 (32–34,51), the final common pathway of apoptosis, remained elevated in all regions. These changes are typical for remodeling myocardium.

Cell morphology. Myocytes in the MR group were elongated, characteristic of myocardial failure (45), compared with cells from the MI-only animals (Fig. 7).

Extracellular matrix and MMPs. In the remote myocardium of infarct + MR sheep, at initial follow-up, there was a significant increase in MMP-13 and MT1-MMP, followed by a decrease at the time of sacrifice (Fig. 8, “shunt remote” bars). MMP-13 increased to 238 ± 29% of baseline at follow-up, and then fell to 79 ± 13% of baseline at 3 months sacrifice; MT1-MMP increased to 165 ± 23% of baseline at follow-up, and then fell to 85 ± 16% of baseline at sacrifice. In the remote myocardium of infarct–only sheep, the rise and fall of MMP-13 did not reach statistical
significant (170 ± 85% of baseline at initial follow-up; 75 ± 22% at sacrifice), and MT1-MMP did not significantly increase at 1 month (Fig. 8, “sham remote” bars). In the border zones of the infarct-only sheep, there was a significant decrease in MT1-MMP at sacrifice (to 60 ± 8% of baseline), with levels maintained in the infarct + MR group. In contrast, TIMP-4 showed progressive rises from baseline to initial follow-up and then to sacrifice in the border and remote zones of both groups, with significantly higher levels at sacrifice in the infarct + MR group border zone (207 ± 9% vs. 152 ± 10% of baseline, p < 0.05). Tissue inhibitor of matrix metalloproteinase-1 showed a similar pattern of progressive increases in the infarct + MR group remote zone, with a significant late rise at sacrifice.

**Discussion**

Myocardial infarction, with loss of contracting myocytes, leads to a sequence of events aimed at preserving cardiac output, including increased LV end-diastolic volume to augment pre-load. Cells in noninfarcted areas hypertrophy by addition of contractile elements in series, creating eccentric hypertrophy without increased wall thickness. This allows the LV to dilate, a process that is also promoted by altered MMP activation with extracellular matrix degradation (52), and acceleration of cell death by apoptosis (32,53). This process eventually culminates in systolic enlargement, extensive fibrosis, and LV failure, the extent of which predicts further cardiac events (2,9). Volume and pressure overload increase wall stress, which can aggravate remodeling (54) and promote eccentric hypertrophy through increased fiber stretch (Fig. 1). Conversely, hemodynamically unloading the LV reverses the functional and cellular stigmata of cardiac failure as well as the remodeling process (55,56). Neurohumoral activation and activation of inflammatory mediators and cytokines initially compensate for reduced cardiac output but, in the long term, mediate increased load, cell death, and cardiac fibrosis (50,57–59). Increased wall stress also up-regulates the production and

**Table 1** Isolated Cardiomyocyte Contractility in the Different Groups

<table>
<thead>
<tr>
<th>Contractility Measurements</th>
<th>Normal Control Sheep</th>
<th>Infarct + MR</th>
<th>Infarct Only</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell shortening (%)</td>
<td>5.7 ± 0.4</td>
<td>2.6 ± 0.7†</td>
<td>3.5 ± 0.3††</td>
</tr>
<tr>
<td>t90% (ms)</td>
<td>390 ± 82</td>
<td>676 ± 68†</td>
<td>510 ± 59††</td>
</tr>
</tbody>
</table>

*p < 0.01 compared with normal control sheep; †p < 0.01 compared with infarct + mitral regurgitation (MR) group. t90% = time to 90% relaxation.

**Table 2** Intracellular Calcium Measurements in the Different Groups

<table>
<thead>
<tr>
<th>Calcium Measurements</th>
<th>Normal Control Sheep</th>
<th>Infarct + MR</th>
<th>Infarct Only</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diastolic (Ca²⁺) (nM)</td>
<td>230 ± 41</td>
<td>320 ± 21†</td>
<td>275 ± 22††</td>
</tr>
<tr>
<td>Systolic (Ca²⁺) (nM)</td>
<td>796 ± 24</td>
<td>535 ± 44‡‡</td>
<td>714 ± 43‡‡</td>
</tr>
<tr>
<td>t90% (ms)</td>
<td>361 ± 34</td>
<td>542 ± 38‡</td>
<td>455 ± 43‡†</td>
</tr>
</tbody>
</table>

*p < 0.01 compared with normal control sheep; †p < 0.03 compared with infarct + mitral regurgitation (MR) group; ‡p < 0.005 compared with infarct + MR group. t90% = time to 90% relaxation of the calcium signal.
release of MMPs, which are believed to be pivotal in altering the extracellular matrix in ways that promote ventricular dilatation (60). Their transgenic activation induces congestive heart failure (61), and their inhibition attenuates post-MI remodeling (62,63).

Mitrval regurgitation, caused by alterations in ventricular geometry and function after MI (10,12), can itself initiate the remodeling cascade, and causes progressive deterioration of ventricular function at a cellular and molecular level (15–17). Mitral regurgitation considerably alters the load on the LV. It increases diastolic wall stress, which can induce eccentric LV hypertrophy and subsequent dilatation and failure (18), and it thereby increases early systolic wall stress. Although MR permits LV emptying during systole into the lower-pressure LA, it has been shown actually to increase end-systolic wall stress and, hence, afterload in patients with chronic MR (64,65). Moreover, MR induces further LV dilatation due to activation of neurohumoral and cytokine components of the remodeling cascade (66–68). As MR is both a cause and a result of LV remodeling, it can potentially exacerbate the vicious cycle spiraling down to cardiac failure unless the remodeling or the MR is reversed (69–71).

Regarding clinical implications, knowing whether the parallel occurrence of MR and MI causes more pronounced remodeling is critical, because at least MR can be eliminated by repairing or replacing the valve, and thus relieving the volume overload induced by it. Separating these 2 dynamic processes in an experimental model is a challenge, because in most existing models they are linked. One recent study of inferior MIs, in fact, concluded that prevention of ischemic MR does not influence the outcome of remodeling (20), despite benefits in nonischemic models (18) and concerns in nonischemic patients (72). In our model, we therefore implanted an LV-to-LA shunt, which created a moderate and standardized MR-like regurgitant flow, independent of a medium-sized antero-apical MI that by itself does not cause MR.

In this controlled model, moderate MR worsened post-MI remodeling. Specifically, it induced large diastolic and systolic LV volumes reflecting volume overload and reduced global function. Reduced contractility was also apparent in less pre-load–dependent physiological measures such as pre-load–recruitable stroke work. These changes paralleled decreased single-cell contractility in the noninfarcted myocardium with reduced SERCA2a levels, which directly correlate with cell contraction (46).

In contrast to progressive changes in the LV, several mediators of the hypertrophic process underwent biphasic changes in the noninfarcted myocardium of animals with MR-type volume overload. These include gp130, a glycoprotein that forms heterodimers to produce different receptors, notably of the interleukin 6/CT1/LIF family. The activation of these receptors has been linked to cardiomyocyte hypertrophy (50,73), and their reduced abundance, manifested by reduced gp130 levels, has been associated with the transition from hypertrophy to failure (74,75). Akt, or protein kinase B, is a serine-threonine kinase activated by

Figure 5  SERCA2a and Phospholamban Quantification

(A) Quantification of sarcoplasmic reticulum Ca2⁺-ATPase (SERCA2a) activity in remote, border, and infarct regions of experimental and normal control sheep hearts. (B) Immunoblots from different regions of the myocardium in different groups for SERCA2a and phospholamban. Note that in the infarct region, there was residual myocardium that was homogenized and protein extracted. Abbreviations as in Figures 1 and 2.
several pro-survival and -hypertrophic factors, among others cardiotrophin-1 (76), acting through the gp130-containing receptor, or growth factor receptors activating phosphoinositide-3 kinase (77). The constitutive activation of Akt in a model of rat ischemia-reperfusion injury reduced cell death and improved function (78). Like gp130, reduced levels of Akt are related to increased apoptosis and a transition from compensatory hypertrophy to a failure phenotype (49). In our model, while both infarct + MR and infarct-only sheep had elevated expression of Akt and gp130 after 1 month, the MR sheep demonstrated a notable reduction in these protein levels after 3 months, compared with the no-MR animals and with baseline. The pro-hypertrophic process, which begins in both groups at 1 month, continues in the animals without MR but is down-regulated at 3 months in the MR group, in association with continued adverse remodeling and LV dysfunction. This decline may represent earlier termination of the hypertrophic process or transition to a failure phenotype specific to the persistent-MR situation; either way, it represents failure of the hypertrophic process. The ongoing drive to cell death, with the suppressed pro-hypertrophic drive, further promotes the transition to failure. These findings are consistent with the emerging view of LV remodeling in general as a failed attempt at compensation for initial stresses on the myocardium—in this case, exacerbated by the MR volume overload (7,74); in others, it reflects the transition from hypertensive hypertrophy to failure (79,80).

There was also excessive activation of MMPs, notably MMP-13 and MT-MMP1, in the noninfarcted remote zone of the MR sheep. This could induce excess extracellular matrix turnover and potentially exacerbate remodeling. However, these same MMP levels fell at sacrifice, with the fall in remote-zone MT-MMP1 being significantly greater in the MR sheep, and the fall in MMP-13 being significant only for the MR group. As this change parallels the down-regulation of the pro-hypertrophic signaling process, it could be also part of the failure of compensation by limiting the adaptive geometrical changes in myocardial hypertrophy. Matrix metalloproteinase inhibitors showed a different pattern, with a progressive rise from baseline to sacrifice that was significantly greater with MR. This suggests a possible shift from matrix turnover to matrix accumulation and stabilization over time, a response typical of remodeling myocardium (81). However, it is clear that neither the MMP nor the TIMP changes adequately compensate for the remodeling process, as LV volume progressively rose while systolic function fell.

This study has several limitations. Ischemic MR affecting a native valve often progressively increases (28,69–71), but is inherently linked to the underlying MI and is not standardized. Based on the motivation for this study, it was...
critical to separate the 2 processes of infarction and regurgitation to determine the incremental role of MR most directly, and to do so with a standardized orifice, which provided stable regurgitant fractions of ~30% throughout the study. Although the LV also fills through this valveless orifice, based on prior experience, this should not importantly alter LV filling (29).

We did not assess whether any one of the molecular changes reported in the present study is causative. We intended to test whether differences in remodeling based on MR are associated with molecular changes described in the remodeling process. Determining which factors are causative versus concomitants, and studying all possible factors, is beyond the scope of this study. Nevertheless, we analyzed pathways extensively linked to the pro-hypertrophic and remodeling processes, and causally implicated through transgenic and pharmacologic modification (61,74,78).

While the impact of isolated MR is important to explore, the study specifically addresses the challenge posed by the recent literature (20), which can be stated as an alternative hypothesis that remodeling in ischemic MR is caused only or predominantly by the underlying infarct, not the MR. This required comparing infarct-only versus infarct + MR groups. The data do show important effects of the infarct alone, with end-systolic volume nearly doubling, and increases in gp130, Akt, and caspase. The changes with infarct...
plus MR, however, were greater, with end-systolic volume nearly tripling, and a different pattern of activation followed by exhaustion of hypertrophic- and apoptotic-signaling molecules gp130 and Akt. The findings, therefore, suggest an additive rather than exclusive role of MR. This is supported by our data (unpublished results) from sheep with an identical shunt and no MI, which did not cause significant remodeling in the same 3-month time-frame in terms of either volume or contractility changes.

Recently, contribution of MR to post-MI remodeling has been challenged by Guy et al. (20) in an inferior MI model in which either MR was reduced using an undersized annuloplasty ring repair, or MI was restrained using a mesh. The MR repair group remodeled more, leading to the suggestion that the MR is not causative, but may be a consequence of post-MI remodeling. The main caveat to these conclusions is that undersized ring annuloplasty may itself alter LV mechanics, as recently shown (82), so as to exacerbate remodeling. Thus, the excess remodeling observed may have been caused, at least partially, by the repair itself, which was performed 2 weeks before infarct creation. The difficulty in eliminating MR as a potential contributor to continued remodeling by annuloplasty has also been reported (72). Another recent paper (83) has suggested that repairing MR in a subgroup of patients with LV dysfunction has no survival benefit. While this is intriguing, the patient population in that study had prominent LV dysfunction at baseline (<30% EF), and thus it is possible that repair was performed too late for many to benefit in terms of LV function or outcome. Based on their inclusion criteria, the patients also had at least moderate-to-severe MR—more severe than in our model. Further, unlike this MI model, roughly 40% of the patients did not have coronary artery disease as the basis for their LV dysfunction. Thus, these clinical data cannot readily be compared with our relatively mild MI model with normal baseline LV function and moderate MR. There is definitely a need for a randomized controlled study to clinically address the issue of mitral valve repair in moderate ischemic MR.

Conclusions

This study demonstrates that MR accompanying MI produces greater ventricular remodeling than a comparable infarction alone, with an earlier transition to a failure phenotype. This is the case even for only moderate MR, most typical of the ischemic situation, for which the greatest uncertainty exists regarding impact on the heart and need for intervention. The changes at the level of the whole heart parallel cellular and molecular abnormalities in the noninfarcted myocardium that reflect the complex remodeling process. These molecular events also progress differently with MR than with comparable infarction alone, with an initial rise in pro-hypertrophic and antiapoptotic signals, followed by their exhaustion. Recognizing this pattern can motivate a search for the responsible molecular “switch,” so that decompensation might be limited or reversed, averting the earlier onset of dilated heart failure precipitated by MR that may portend a worse prognosis.

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