Coronary reperfusion therapies have led to a reduction in the incidence of mechanical complications of acute myocardial infarction (AMI), but the associated mortality of these complications has remained high. Ventricular septal rupture is the most common mechanical complication after myocardial infarction and occurs in approximately 0.21% with ST-segment elevation myocardial infarction and in 0.04% with non-ST-segment elevation myocardial infarction. Surgery is the only definitive treatment but it is associated with high operative mortality and morbidity and, in some centers, alternative treatment with mechanical support of circulation and trans-catheter closure of the defect is being used. We continue to believe that immediate surgery offers the best opportunity for long-term survival. Patients should be taken directly to the operating room or via the heart catheterization laboratory if the coronary artery anatomy is unknown. This should be done as soon as the diagnosis is made, as acute rupture of the interventricular septum is a surgical emergency. The operative technique of infarct exclusion has been our preferred method and when performed in hemodynamically stable patients, the operative mortality is around 10%, although much higher in patients in cardiogenic shock. Patch dehiscence and persistent shunts are uncommon after closure of the defect using this technique. Delaying operative intervention in hemodynamically stable patients frequently leads to cardiogenic shock and multi-organ failure. Transcatheter closure of these acute septal lesions is complex, not necessarily feasible in all patients, and there is no evidence that the results are superior to surgery.

Keywords: Acute myocardial infarction (AMI); mechanical complications of myocardial infarction; post-infarction ventricular septal rupture

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Introduction

Surgery for mechanical complications of acute myocardial infarction (AMI) used to be common when I started practicing four decades ago (1), and I was able to accumulate considerable experience in this field and subsequently developed a novel approach to treat a ruptured interventricular septum (2,3). The emergence of coronary artery reperfusion therapies have reduced the frequency of mechanical complications of AMI but the mortality rate has remained high (4). In a recent report by Elbadawi and colleagues (4) based on The National Inpatient Sample (NIS) database from 2003 through 2015, mechanical complications of AMI occurred in 0.27% of patients with ST-segment elevation myocardial infarction (STEMI) and in 0.04% after non-ST-segment elevation myocardial infarction (NSTEMI). However, the mortality was 42.4% in patients with mechanical complications of AMI and 12.7% in patients without, and this did not change during the period of observation (4). These poor outcomes are humbling but challenge us to seek better treatment alternatives for patients with mechanical complications of AMI.

STEMI is more likely to be associated with a transmural infarction than NSTEMI (5). A transmural AMI is a prerequisite for ventricular wall rupture and hence is more common after STEMI than NSTEMI (4,5).
Ventricular septal rupture is the most common type of mechanical complication after AMI (1,4). In Elbadawi's report (4), of 13,767 patients who suffered mechanical complications, 10,344 (75%) developed a ventricular septal defect (VSD) and the remaining patients developed mitral regurgitation due to ruptured papillary muscle (19%) or free wall rupture (6%). The most important determinant of outcome in post-infarction VSD is the development of heart failure (left, right, or both) and cardiogenic shock (2-5). The severity of heart failure is related to the extensiveness of myocardial necrosis and the left-to-right shunt. Cardiogenic shock in patients with anterior VSD is often associated with extensive left ventricular infarction and in patients with posterior VSD is associated with extensive right ventricular infarction (6-9). The magnitude of the shunt is also an important determinant of cardiogenic shock in both cases (7,8). Although the diagnosis of post-infarction VSD can be made by auscultation of the heart, it may be difficult to differentiate from acute mitral regurgitation and a bedside trans-thoracic echocardiogram will usually confirm the diagnosis. Some degree of mitral regurgitation in patients with posterior VSD is common because of posterior wall and papillary muscle infarction. Medical treatment is ineffective and most patients die within 4 weeks (10). Thus, surgery or a catheter-based intervention is needed to save these patients. We believe that immediate surgery offers the best chance of survival after post-infarction VSD (3).

Surgical treatment of post-infarction ventricular septal wall rupture

Cooley and colleagues reported the first case treated with surgery using an approach through the right ventricular outflow tract (11). This approach does not offer adequate exposure of all parts of the septum and a few years later, most surgeons adopted an incision through the infarcted left ventricle (12). The location and length of the incision depends on where the defect is: apical, anterior or posterior septum. Daggett and colleagues from the Massachusetts General Hospital made important contributions to the surgical treatment of this lesion (13,14). They introduced the concept of ‘infarctectomy’ and reconstruction of the interventricular septum and infarcted ventricular wall with Dacron patches (15). Apical VSDs were managed by apical amputation and approximation of the walls over strips of Teflon felt, and more extensive infarcts by infarctectomy and reconstruction of the septum and ventricular wall with separate patches of Dacron fabric (15).

In 1987, we introduced a new operative procedure whereby no infarctectomy was done and the left ventricular cavity was largely excluded from the infarcted myocardium with a pericardial patch sutured to the endocardium of the left ventricle (2,3). We coined the term ‘infarct exclusion’ to describe this approach and we continue to believe it is the best therapy for patients with an extensively infarcted septum, particularly in cases of posterior VSD. Apical VSD can be safely repaired as originally described by Daggett and associates (13-15), and this is illustrated in Figure 1.

Infarct exclusion technique

Myocardial revascularization should be done before septal repair in patients with multi-vessel coronary artery disease. If the VSD is associated with single vessel disease, surgical revascularization will depend on what was done percutaneously prior to the septal rupture. If the single diseased coronary artery was reopened by percutaneous coronary intervention (left anterior descending in anterior VSD and right coronary artery or dominant circumflex in posterior VSD), there is no need for bypass grafting. If the artery remains occluded, revascularization may be beneficial to improve left or right ventricular function. There is no point in bypassing an artery if it supplies only infarcted myocardium.

In patients with anterior VSD, an incision is made in the apex of the left ventricle 10 to 15 mm lateral to the path of the anterior descending artery, through the infarcted muscle, and extended just above where the septal defect is located. The margins of the infarcted septum and anterior wall are identified. A glutaraldehyde-fixed bovine pericardial patch is tailored in the shape of the left ventricular infarct. This patch is usually oval and measures approximately 4 cm × 6 cm in most patients. The pericardial patch is then sutured to the lower part of the non-infarcted endocardium of the interventricular septum with a continuous 3-0 polypropylene suture as illustrated in Figure 2. The patch is also sutured to the non-infarcted endocardium of the anterolateral ventricular wall. If the infarct involves the base of the anterior papillary muscle, the suture is brought outside of the heart and buttressed on a strip of bovine pericardium or Teflon felt, applied to the epicardial surface of the left ventricle. Once the patch is completely secured to the endocardium of the left ventricle, the infarcted septum becomes largely excluded from the left ventricular cavity. The ventriculotomy is closed over two strips of bovine pericardium or Teflon felt, as illustrated in Figure 2. No
infarctectomy is performed unless the necrotic muscle along the ventriculotomy sloughs at the time of its closure. This area will be exposed only to right ventricular pressure once cardiopulmonary bypass is discontinued. The space between the patch and infarcted wall becomes obliterated within a few days.

In patients with posterior VSD, an incision is made in the base of the inferior wall of the left ventricle, 5 mm from the posterior descending artery. It should not be made further into the left side of the ventricle as the posterior papillary is adjacent and can be damaged. This incision starts at the base of the inferior wall, 2 cm from the atrioventricular junction, and extends proximally toward the mitral anulus and distally toward the apex of the ventricle. Stay sutures are passed through the fat pad of the apex of the ventricle to hold the heart upside and to the margins of the ventriculotomy to facilitate exposure of the ventricular cavity. The VSD is carefully located; it is usually in the proximal half of the posterior septum, close to the mitral valve and right ventricle. The posteromedial papillary muscle is often infarcted. A large bovine pericardial patch is used for repair. The patch is sutured to the fibrous anulus of the mitral valve with a continuous 3-0 polypropylene suture starting at the ventricular side of the central portion of P2 of the posterior leaflet of the mitral valve. Some chordae tendineae between the posterior leaflet and ventricular wall may have to be severed if they are obstructing the suture line. This suturing is carried along the fibrous mitral anulus to approximately the level of the medial commissure of the mitral valve, where the suture is interrupted. The large bovine pericardial patch is trimmed to fit the defect. Next, starting at the base of the septum, the patch is sutured to the endocardium of healthy septum towards the apex. The other side of the patch is sutured to the posterior wall of the left ventricle along a line corresponding to the medial margin of the posteromedial

Figure 1 Repair of apical ventricular septal rupture. Simple amputation of the left ventricular apex and approximation of the left ventricular wall, septum and right ventricular wall without excessive compression of the tissues is often an effective treatment.
papillary muscle base. Given the posterior wall of the left ventricle infarct, it is necessary to use full-thickness bites and anchor the sutures on a strip of pericardium or Teflon felt applied on the epicardial surface of the posterior wall of the left ventricle, as shown in Figure 3. Once the patch is completely sutured to the distal part of the posterior wall, the sutures used to secure the patch to the septum and those used to secure the patch to the infarcted posterior wall are brought up toward the epicardium and tied to each other. The posterior wall ventriculotomy is then closed in two layers of sutures buttressed on strips of pericardium or Teflon felt. The infarcted right ventricular wall is left undisturbed. Occasionally, a patient may have severe mitral regurgitation and mitral valve replacement is necessary in such situations. We have done this only twice and in both instances, through the posterior wall ventriculotomy.

Figure 3 shows intra-operative photographs of the various stages of this operation.

Trans-catheter closure of post-infarction ventricular septal wall rupture

The first report of transcatheter closure of a post-infarct VSD was 1988 (16). This is a complex procedure and not all patients are candidates, particularly those with posterior VSD given their proximity to the mitral valve and complexity of the regional anatomy. Schlotter and colleagues (17) conducted a systematic literature review to evaluate the effectiveness of percutaneous closure of post-infarction VSD. They identified 13 series of more than 5 cases per report, for an overall 273 patients (17). Cardiogenic shock was present in 48% of cases, the timing
of the intervention was within 2 weeks from rupture in 42% and successful device implantation rate was 89%. The in-hospital mortality rate in patients successfully treated was 32% (17). The authors concluded that transcatheter closure is an alternative to surgery in selected patients. Larger experience is needed before deciding if this is indeed as effective as surgery to treat post-infarction VSD, particularly in the long-term.

**Discussion**

A report on the surgical outcomes of post-infarction VSD based on the STS Database from 1999 through 2010 indicated that only 2,876 patients underwent surgery in 666 participating institutions with an average of 0.09 to 3.7 patients/year per cardiac unit (18). Thus, very few surgeons operate on more than one patient per year. The overall mortality was 42.9%, higher in patients who had surgery within seven days from the AMI (54.1% vs. 18.4%) (18). I was the invited discussant of that paper and I pointed out that I had reviewed the surgical experience with post-infarction VSD at Toronto General Hospital from 1990 to 2010, with 91 patients identified. The overall operative mortality was 40%, but nine surgeons performed the operations and mortality among surgeons ranged from 20% to 100% — it appears to be related to the number of operations each surgeon performed. Two surgeons operated on 49 patients and nine died whereas seven surgeons operated on 43 patients and 22 died. Repetition makes perfect and it is difficult to gain experience with an operation that is now so rarely performed. In addition, these operations are difficult and likely cannot be mastered by most surgeons largely due to its rarity. Thus, in order to improve the outcomes of post-infarction VSD, patients should be referred to centers where both surgical and catheter-based techniques expertise are available (19,20). I firmly believe that patients with post-infarction VSD should be operated on as soon as the diagnosis is made to prevent
worsening of the clinical presentation. If the expertise to do that is not available, hemodynamically stable patients should be transferred to referral centers as soon as the diagnosis is established, and patients in cardiogenic shock should have a device implanted to mechanically support the circulation (Impella or venous-arterial extracorporeal membrane oxygenation) and then should be transferred to a large center. The probability that the above proposal for the management of these patients will come to fruition may be more utopic and realistic.

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Footnote

Conflicts of Interest: The author has no conflicts of interest to declare.

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Figure 4 Intraoperative photographs of a ruptured posterior septum repair as described in Figure 3.
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