Case 3 - Elderly Female Patient with Acute Myocardial Infarction Presenting Sudden Shock during Thrombolytic Treatment

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A 72-year-old female patient sought medical assistance due to precordial pain that had lasted for two hours.

The patient had received a diagnosis of ischemic heart disease 10 years before. She had been submitted to myocardial revascularization in 1991 and was unable to give any specific details about the treatment or the grafts used at the surgical procedure. Five years after the surgery, she presented an episode of prolonged precordial pain. She sought medical assistance and, at this time, alterations in ventricular repolarization were diagnosed at the electrocardiogram (Figure 1), whereas the serum markers of myocardial injury remained unaltered.

The patient remained asymptomatic for five subsequent years, when she started to present recurrent and daily episodes of precordial pain, which lasted a week, until she was admitted to the hospital. The electrocardiogram (ECG) showed sinus rhythm, HR of 63 bpm, QS complexes in lead aVF and from V6, and ST-segment elevation in leads I, II, III, aVF, and from V4 to V6 (Figure 1). The ECG alterations were not reverted by the administration of sublingual nitrate.

The laboratory assessment disclosed levels of urea: e68 mg/dl, creatinine: 1.2 mg/dl, glycerol: 189 mg/dl and creatine kinase: 42 U/l. A diagnosis of ongoing myocardial infarction was achieved and TNK-tPA (Tecneplase - a genetically modified form of t-PA, with increased PAI-1 resistance, increased fibrin specificity and longer half-life) was administered as bolus of 40 mg with non-fractionated heparin as bolus of 6,060 UI/kg, followed by 1,000 UI/h IV. A residual retrosternal pain persisted.

One hour after the start of the treatment, the patient presented right hypochondrial pain, followed by cardiogenic shock. Volume replacement and dopamine were administered. The control electrocardiogram (April 7, 2001, 2:17 PM) showed sinus tachycardia, HR of 110 bpm, QS complexes with ST-segment elevation in ul, III, aVF, and from V4 to V6 (Figure 3). A rescue angioplasty was indicated. However, the patient presented cardiac arrest with pulseless electrical activity, did not respond to the resuscitation maneuvers and died.

Clinical aspects
In a case such as the one described here, of a 72-year-old female patient, with history of coronary artery disease and myocardial revascularization carried out more than 10 years before, who sought emergency medical assistance due to typical precordial pain, the initial evaluation must always include an electrocardiogram (ECG). A normal ECG or an ECG with non-specific alterations or signs of ischemia would lead to an initial approach that would be different from the one described here. The admission ECG of this patient showed ST-segment elevation.

Considering that the patient had been submitted to a previous myocardial revascularization (MR), the correlation of the diagnostic findings of the ECG deserves special attention. First, the presence of chronic coronary artery disease, as well as the previous MR without the appropriate description of the grafts used in the surgery makes it difficult to establish the electrocardiographic correlation with the culprit artery, due both to the presence of bypasses and the possible existence of an extensive collateral network. It is also important to remember that, in spite of the native artery lesions, the probability of lesions in bypasses is of 38%1. Additionally, the history of chronic coronary disease in elderly diabetic women and the QS in the anterior wall raises the possibility of previous myocardial infarction in the anterior wall, and, as a consequence, the possible correlation of the ST-segment elevation corresponding with the presence of aneurysm in the same wall. Still, the existence of ST-segment elevation in
the inferior wall associated to typical precordial pain justifies the chemical or mechanical reperfusion therapy, as described in the present case.

The description of the initial approach with the use of tenecteplase (TNK) and non-fractionated heparin can be considered the standard approach for a case as the one described here, as the early reperfusion therapy is the treatment of choice for the condition\(^2\). The improvement of the pain can be considered a reperfusion criterion that favors the opening of the artery, although its significance is uncertain\(^2\).

However, the lack of some data in the description hinders a more thorough analysis of the patient’s clinical condition. There is no description of the pulses or chest x-rays that could rule out the presence of an aneurysm in the thoracic aorta leading to the right coronary artery dissection, which is the culprit artery in 80% of the cases of aorta dissection associated with coronary dissection.

The most noteworthy fact concerning the patient’s evolution and which deserves further investigation is the rapid clinical worsening with the onset of cardiogenic shock

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**Figure 1** - ECG: ventricular repolarization alterations.

**Figure 2** - ECG: Ongoing infarction in the anterior and inferior walls.
Cardiogenic shock is present in less than 1% of the patients at admission. However, more than 50% of those who will develop cardiogenic shock do so within the first 24 hours, with a mean time of 5 hours.

Cardiogenic shock mortality is 56% to 74% and the most common causes of shock according to the Shock Trial are: LV dysfunction in 79% of the cases; acute mitral regurgitation in 7%; ventricular septal defect in 4%; isolated right ventricular shock in 4% and cardiac tamponade in 7%.

Considering that the patient was adequately monitored and that the second ECG showed tachycardic sinus rhythm without arrhythmias, which can be secondary to the use of dopamine, the most common causes of arrhythmia at the initial post-AMI period become unlikely. Therefore, the differential diagnosis is restricted to other causes of precordial pain, to the associated right ventricular involvement, to the mechanical complications of AMI evolution and the complications inherent to the proposed reperfusion therapy. Each one of these possibilities will be discussed separately.

First, among the clinical pictures of precordial pain associated to ST-segment elevation, which are not basically coronary, only the thoracic aorta dissection with ostium involvement and coronary dissection deserves attention. The acute dissection of the aorta normally affects elderly and hypertensive patients, as the case described here. On the other hand, the involvement of the coronary ostia is rare; only 1% to 2% of the patients present such complication. Of these, the great majority presents right coronary involvement, with ST-segment elevation in the inferior wall, as described in this case. In cases where the dissection goes unnoticed and the case is treated with thrombolysis, mortality reaches 71%, due to the increased risk of cardiac tamponade. As mentioned before, the lack of detailed information on pulses, murmurs and chest x-rays does not allow the diagnosis to be confirmed or ruled out.

The involvement of the right ventricle (RV), associated with the left ventricular AMI condition must always be recalled in cases with AMI of the inferior wall, as the case described here. As mentioned before, the absence of right leads in the admission ECG limits the diagnosis, especially when one considers that the presence of 1-mm ST-segment elevation in V4R has a sensitivity of 88% and a specificity of 78% for the presence of or clinical picture evolution to RV involvement. Even without the right leads, the association of the ST-segment elevation in DII and DIII are contrary to the hypothesis of RV involvement. The presence of ST-segment elevation in DII higher than in DIII has a sensitivity of 97% and a specificity of 57% for right ventricular AMI; however, its absence does not help to elucidate this possibility. Moreover, the presence of crackling rales goes against the condition of right ventricular AMI, as the classic clinical picture courses with infarction and shock in the presence of clear pulmonary auscultation. Other criteria, such as the fact that the patient did not respond to volume replacement or dopamine, which are part of the optimized therapeutics for right ventricular AMI, would make it even less likely for a right ventricular AMI to be responsible for the dramatic evolution. Not only that, but there have been descriptions in the literature of rapid improvement after reperfusion in patients with right ventricular AMI within the first hours. Moreover, there have been reports of patency in the culprit artery right after thrombolysis in more than 75% of the patients that were included in the GUSTO-I study. It is worth remembering that, of the thrombolized cases of right ventricular AMI, only 12% develop shock and 2% die, whereas 82% of the non-thrombolized ones develop shock and these have presented a mortality rate of up to 58% in several studies.
Iatrogenesis must not be ruled out as a diagnostic possibility, secondary to clinical conditions where early aggressive therapy such as chemical thrombolysis is part of the routine therapeutic arsenal. In patients that are thrombolyzed with TNK, as the case described here, it is necessary to emphasize that this is a safe thrombolytic drug, with an incidence of bleeding lower than streptokinase or r-TPA (26 vs 29% in the ASSENT II study, which compared TNK with r-TPA)\(^\text{11}\). However, the presence of clinically significant bleedings is lower than 2% in studies that tested TNK. On the other hand, older age, as in the present case, is one of the most important factors that predispose patients to major bleedings. Additionally, women present a higher risk of bleeding than men. Still, even in cases of major bleeding after thrombolysis, the dramatic clinical evolution and the absence of response to volume replacement are very unlikely, causing such diagnosis to be laid aside.

Finally, in the presence of a picture of cardiogenic shock with a dramatic evolution, one must fully evaluate any signs that can be associated with mechanical complications after the AMI. Among them are: ischemic mitral regurgitation, LV free wall rupture leading to tamponade, ventricular septal defect (VSD) due to septal rupture and previous aneurysm or pseudoaneurysm rupture.

Mitral regurgitation classically courses with acute left heart failure with important pulmonary congestion\(^\text{14}\), which is very dissimilar from the patient’s clinical evolution, being highly unlikely.

The ventricular septal defect is a rare complication, with an incidence of 2% in the pre-therapeutic era of reperfusion. With the effective use of reperfusion techniques, the incidence of VSD has become lower than 0.2%, as described in the GUSTO-I study\(^\text{15}\). Traditionally, the onset of VSD takes place between the 3\(^{\text{rd}}\) and 5\(^{\text{th}}\) days, but it can be present in less than 24 hours after the initial event\(^\text{15}\). The clinical evolution courses with shock due to biventricular dysfunction with a predominance of the RV. More than 80% of the patients present murmurs and more than 50% present thrills.

The absence of the description of these signs makes such diagnosis unlikely. Nevertheless, the presence of inferior wall ST-segment elevation associated with anterior wall ST-segment elevation favors this diagnosis, due to the possibility that the picture is associated with AMI in the territory of the anterior interventricular branch (AIB) of the left coronary artery that surpasses the apex and due to the absence of collaterals, leading to important ischemia of the septum and its rupture. On the other hand, the presence of anterior Q5 complexes, as previously described, makes the possibility of anterior wall aneurysm more probable, thus making the hypothesis of AMI due to AIB injury leading to VSD unlikely. Although no echocardiogram or pulmonary artery catheter introduction was carried out for diagnostic clarification, VSD is an unlikely diagnosis.

Left ventricular aneurysms are more commonly found in the anterior wall\(^\text{16}\). Currently, the incidence of aneurysms in thrombolized patients is less than 10%\(^\text{16}\). In spite of the lack of precise data, the presence of anterior Q5 complexes associated to the ST-segment elevation in the same leads raises the possibility of anterior wall aneurysm. The only complication associated with the true aneurysm that can course with such a dramatic evolution, as the one described here, is its rupture. However, a true aneurysm is a fibrotic scar with a stable structure and presents low risk of rupture. Although such diagnosis is uncommon and the clinical evolution is compatible with the described clinical picture, such hypothesis cannot be ruled out, considering the available data.

Pseudoaneurysms are false cavities that establish a communication between the left ventricle and the pericardial cavity. Although they are more common in the inferior wall, their occurrence in the anterior wall is also possible. Although many of them present murmurs, their absence does not rule out this hypothesis. Pseudoaneurysms have a risk of rupture of up to 40% and a mortality rate > 50% when rupture occurs\(^\text{17}\). As the clinical evolution is quite similar to our patient’s evolution, this hypothesis cannot be ruled out either, based on the available data.

The last mechanical complication to be remembered is the LV free wall rupture, which occurs in up to 1% of the cases of AMI\(^\text{18}\). Of the fatal cases of AMI, the LV free wall rupture is the cause of death in 12% of the patients that are thrombolized and in 7% of the patients that are not\(^\text{18}\). Among the risk factors for the rupture, the clinical case patient presented: age older than 70 years; female sex; ST-segment elevation and Q wave at the admission ECG.

Although it normally appears between the third and the fifth days after the AMI, recent data have shown that, with the advent of reperfusion therapy, the occurrence of free wall rupture has become rarer, albeit earlier, and that it can occur within the first hours after the AMI\(^\text{19}\). Two distinctive presentations of rupture have been described\(^\text{19}\) - an early one, in cases of anterior wall AMI with a sudden clinical picture, as the one described here. This form occurs within the first 2 or 3 days after the AMI. The other form usually occurs after 4 days, as an evolution of the infarction area and usually has a slower evolution. Cases presenting the early form usually develop hemopericardium, leading to tamponade with a preferential clinical presentation of right ventricular AMI, followed by respiratory arrest with pulseless electrical activity (PEA). The clinical presentation of PEA without a previous clinical picture of heart failure in patients without a previous history of AMI has an accuracy of 95% for free wall rupture\(^\text{19}\). When the previous presentation included heart failure, the accuracy of the arrest with PEA is only 17%. Although it is not possible to completely rule out the possibility of a previous AMI, these data suggest free wall rupture as a probable diagnosis of the clinical picture, even though it is not possible to state that this was the patient’s first AMI.

**Diagnostic hypothesis**

Acute myocardial infarction, complicated by free wall rupture and tamponade.
Necropsy

The pericardial sac was filled with a large amount of partially coagulated blood, demonstrating the occurrence of cardiac tamponade, which was the immediate cause of death. Transmural myocardial infarction and rupture were observed, with approximately 3 days of evolution, in the anterosuperior and septal walls of the LV (Figure 4), including the apical wall. The rupture occurred exactly at the apex of the heart and there was also hematic infiltration of epicardial fat (Figure 5). The LV myocardium exhibited moderate hypertrophy.

The study of the coronary arteries showed recent occlusive thrombosis of the bypass graft to the anterior interventricular branch of the left coronary artery. Moreover, this saphenous vein also showed the presence of intimal fatty plaques without severe luminal occlusion (Figure 6). Histologically, the coronary arteries showed the presence of atheroma plaques, with various degrees of occlusion, being greater than > 70% in the initial and mid-thirds of the anterior interventricular branch of the left coronary artery and the mid-third of the right coronary artery (Figure 7).

Advanced aortic atherosclerosis was also observed (Figure 8), with dilation of the ascending segment of the aorta. Other necropsy findings were pulmonary emphysema and benign nephrosclerosis associated with systemic arterial hypertension.

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Anatomopathological diagnoses

Ischemic heart disease, with ruptured acute myocardial infarction and cardiac tamponade.

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Comments

The post-infarction ventricular rupture can affect the free wall, the ventricular septum or the papillary muscles.
Anatomopathological Session

No association has been described between the size of the infarction and the risk of rupture; and it is not unusual for the infarction to be small. Data collected at the Laboratory of Pathology of the Heart Institute (InCor) in cases of necropsy have shown that most ruptures occur at the first infarction (72%), after an average of 5.4 days of evolution. The most commonly affected site is the lateral wall, as shown in the aforementioned series and also others in the literature. The rupture of the ventricular apex is rarely described. A Forensic Medicine series reported a prevalence of 6% concerning this rupture site.

There is some controversy regarding the role of systemic arterial hypertension in the outcome as ventricular rupture. Whereas some authors indicate arterial hypertension as a risk indicator for cardiac rupture, others have shown, in series of infarcted patients with hypertension, a lower prevalence of rupture in comparison with normotensive patients. Mann and Roberts observed similar proportions of hypertensive patients between those with and without post-infarction cardiac rupture in an anatomical series.

An immunohistochemical study in humans showed that the patients with myocardial infarction that died due to cardiac rupture had lower levels of catenin alpha-1 than those that died due to other causes. Catenin is a cell adhesion protein present in the intercalated disks of the contractile cardiac cells, the cardiomyocytes. The same finding was also observed outside the infarcted area, which raised the hypothesis that such individuals would carry a genetic defect or at least, a genetic polymorphism. This hypothesis has been tested, but has yet to be proven.

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References


