How sound is the evidence that thrombolysis increases the risk of cardiac rupture?

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Rupture of the free wall of the heart is a catastrophic and usually lethal event. It occurs in up to 10% of patients dying in hospital of acute myocardial infarction. While there is unequivocal evidence for the long-term benefit of thrombolytic therapy in patients with suspected myocardial infarction, in many streptokinase trials there were more deaths during the first 24–36 hours in treated patients than in controls. It has been suggested that thrombolytic therapy increases the risk of cardiac rupture as a consequence of extensive myocardial haemorrhage and that this effect may partly explain this early hazard. The issue may be addressed from the following perspectives:

- Is there evidence from animal studies that thrombolysis promotes myocardial haemorrhage?
- Does myocardial haemorrhage occur and increase the risk of rupture in humans?
- Is there evidence from randomised trials that the use of thrombolytic therapy increases the risk of cardiac rupture?
- What does this mean in the overall context of risk versus benefit when patients are selected for coronary thrombosis?

Myocardial haemorrhage: animal evidence

A review of animal studies indicates that early reperfusion leads to salvage of myocardium but that it may also result in morphological differences, including myocardial haemorrhage, in the necrotic myocardium as compared with non-reperfused infarcts. The potential adverse consequences of myocardial haemorrhage include extension of ischaemic injury, delayed healing, and functional myocardial abnormalities. Haemorrhagic necrosis is directly related to the duration of occlusion and the extent of damage to the microvasculature. It occurs within the ischaemic zone where vascular death has followed myocardial cell death—that is, in regions with irreversible injury. Moreover, myocardial haemorrhage is likely to be related to reperfusion and not to thrombolysis itself. The addition of streptokinase does not lead to a greater degree of haemorrhage than that experienced with reperfusion alone.

Studies suggesting an extension of ischaemic injury caused by myocardial haemorrhage relied on serial serum creatine kinase (CK) concentrations to estimate infarct size. It has been shown, however, that reperfusion enhances the release of CK per gram of infarcted myocardium. Therefore, serial CK determinations are an inaccurate means of predicting extent of myocardial infarction after reperfusion. The effects of thrombolysis on the healing process within infarcted myocardium are controversial. Impaired healing may be suggested because reperfusion leads to an altered early granulocyte response. However, coronary reperfusion after myocardial infarction alters the pattern of injury and the cellular response to the evolving myocardial infarction, so that classic criteria for infarct age may not apply. For infarcts of the same clinical age, those receiving reperfusion therapy were judged to have older histological ages. Reperfusion does not seem to have a detrimental effect on late collagen formation.

There is no evidence that infarct size is increased after reperfusion and myocardial haemorrhage. With short durations of occlusion (<120 minutes) infarct size was smaller than the potential area at risk. This suggests that any deleterious effect of myocardial haemorrhage would be less important than the benefit of reperfusion. With longer periods of occlusion, infarcts were transmural and infarct size was not increased beyond that predicted. There are no data on the effect of myocardial haemorrhage on long-term left ventricular function.

Myocardial haemorrhage: postmortem studies in humans

Myocardial haemorrhage has been observed in humans after cardiac surgery, coronary angioplasty, and thrombolytic therapy. Though cardiac rupture has been found in association with myocardial haemorrhage, it remains unclear whether the risk is increased after thrombolysis or whether any observed association is the result of case selection and reporting.

A postmortem study by Cowan et al suggested that early myocardial haemorrhage may increase the risk of ventricular rupture. Forty three cases of death after reperfusion therapy were collected prospectively from 36 hospitals in Washington state and compared...
with matched controls (no reperfusion therapy) derived from the files of the principal investigator and two local hospitals. Eight of the haemorrhagic infarcts after reperfusion therapy showed ventricular rupture and four of the infarcts were less than 24 hours old. In contrast, 890 files had to be reviewed to match four controls with cardiac rupture within 24 hours of myocardial infarction.

Gertz et al examined the hearts of 61 patients with a first fatal myocardial infarction: 23 received recombinant tissue-type plasminogen activator and 38 did not. Patients died a median of 2-0 and 4-2 days respectively from symptom onset. The frequency of cardiac rupture was lower among those receiving thrombolytic therapy (22% v 47%, p = 0.045). Similarly, the frequency of rupture among the patients who died within 72 hours of symptom onset was also lower among those patients who received recombinant tissue-type plasminogen activator than among those who did not (20% v 64%, p = 0.02).

Gertz et al also examined 52 hearts of patients who had received recombinant tissue-type plasminogen activator for evolving myocardial infarction as part of the Thrombolysis in Myocardial Infarction (TIMI) study and who died at a median of 2-7 days from symptom onset. Myocardial necrosis was confirmed histologically in 43 patients: 23 infarcts were grossly haemorrhagic and 20 were non-haemorrhagic. Haemorrhage was confined to areas of necrosis in all cases. The frequency of cardiac rupture was similar in the haemorrhagic (26%) and non-haemorrhagic (25%) infarctions.

Kleiman et al analysed the cause of death in 63 patients who died within the first 18 hours of enrolment in the TIMI Phase II study. Ventricular rupture was deemed to be the cause of death when there was direct evidence from visualisation of the heart or when sudden death occurred in the setting of continuous visualisation and normal blood pressure and rhythm. Pump failure was responsible for 62% of early deaths, cardiac rupture for 16%, arrhythmias for 13%, and complications of therapy for 9%. In the subgroup of patients without initial hemodynamic compromise, rupture accounted for a larger proportion of deaths but was still less common than pump failure. It was more likely to be diagnosed in patients who had necropsy or surgical exploration of the chest, suggesting that clinical suspicion of rupture may have led to more frequent necropsies. The effect of early and deferred beta blockade was assessed within a subgroup of the TIMI II trial. The small number of deaths in each category precluded the demonstration of any effect of beta blocker therapy on rupture. Previously, an effect of beta blockers on early cardiac rupture was proposed after retrospective review of the First International Study of Infarct Survival (ISIS-1), a randomised trial to assess the effect of early beta blockade on survival after myocardial infarction. Almost all of the mortality reduction associated with the use of atenolol occurred on days 0 and 1. Significantly fewer deaths ascribed to cardiac rupture (confirmed by necropsy) or to electromechanical dissociation occurred in the atenolol group than in the controls (20 v 54, 2p < 0.02).

A retrospective review of the 1386 in-hospital deaths in the Gruppo Italiano per lo Studio della Streptocinasi nell'Infarto Miocardico (GISSI) trial showed that the most frequent cause of death was cardiac failure (52%), which was significantly less common in treated patients than in controls. No differences between treatment groups were observed for other causes of death including electromechanical dissociation, sudden cardiac death, and deaths from extracardiac causes. While cardiac rupture accounted for 9-1% of total deaths, it was overrepresented (58%) among the 16% of patients who died and underwent necropsy in this unblinded trial. Diagnostic suspicion may have resulted in a necropsy being performed more often in cases of rupture.

Cardiac rupture: evidence from randomised trials

On the basis of a review of the causes of death in two placebo controlled trials, Vasilomanolakis et al suggested that streptokinase predisposes to cardiac rupture. The differences observed (24% streptokinase, 16% placebo), however, may have been due to chance (p = 0.54). An overview of 33 randomised trials of thrombolysis performed before 1985 did not find an excess rate of cardiac rupture among those treated with a fibrinolytic agent for myocardial infarction. The reliability of these analyses is limited, however, because not all studies reported deaths, and necropsy findings or the rate of necropsy was low. In addition, the thrombolytic dose protocols and concurrent therapies in these trials were not typical of current practice, so any generalisations are tenuous.

In the Intravenous Streptokinase in Acute Myocardial Infarction (ISAM) study 1-2% of those treated with streptokinase had ventricular septal rupture or clinically suspected rupture of the free wall within 21 days of randomisation compared with 1-5% of controls. Of cardiac deaths, rupture (septal and free wall) was the cause among 20-4% (10/49) of streptokinase treated patients and 21-0% (13/62) of controls. Rupture caused only two (1/17 control, 1/2 streptokinase) of the 29 deaths occurring within 14 days in the Western Washington study.

In the ISIS-2 trial cardiac rupture was reported in 0-9% of both the streptokinase and placebo groups. The combination of streptokinase and aspirin was associated with a 0-2% absolute reduction in the rate of cardiac rupture (0-9% v 0-7%). In addition, cardiac arrests from causes other than ventricular fibrillation (possibly electromechanical dissociation) were reduced by streptokinase and the combination of streptokinase and aspirin. Though there were
more deaths in the streptokinase group within the first 24–36 hours the rates of cardiac rupture seemed to be similar. In the GISSI trial there was no difference in the rate of cardiac rupture (1.0% streptokinase, 1.1% controls). Therefore, these studies do not provide evidence that thrombolytic increases the overall risk of cardiac rupture and the timing of thrombolytic therapy for myocardial infarction by performing a meta-analysis of placebo controlled trials where cardiac rupture was reported and the necropsy rate was >50%. The limitations of this meta-analysis have been discussed elsewhere and caution urged in the interpretation of the results. Overall, rates of cardiac rupture among those receiving streptokinase (29/831) and controls (29/807) were similar. However, Honan et al suggested that the risk of rupture was biphasic and directly related to the timing of thrombolytic therapy in relation to the onset of symptoms: early treatment (<7 h) decreases the risk and late treatment (>17 h) increases the risk. The results from the GISSI trial were taken as corroborating evidence despite the low rate of necropsy (16%) and the unblinded nature of the trial.

Clinical context

There is unequivocal evidence supporting the benefit of thrombolytic therapy given up to six hours from the onset of myocardial infarction. There is considerable evidence that there is no strong evidence to suggest that coronary thrombolyis increases the overall risk of cardiac rupture or explains the early risk. Pump failure, not cardiac rupture, continues to be the predominant cause of death in patients receiving thrombolytic therapy. Nevertheless, the results of the study by Honan et al, suggesting an increased risk of rupture with delayed therapy, deserve further investigation. If nothing else, this study would compel researchers to investigate mechanisms of death after thrombolyis and to design adjunctive strategies to reduce myocardial damage and improve the prognosis of patients with myocardial infarction.

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