Acute myocardial infarction related to smoke inhalation and myocardial bridging

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Summary: A previously healthy 26 year old woman who was exposed to smoke during a house fire developed acute anterior myocardial infarction complicated by ventricular fibrillation. Subsequent left ventriculography confirmed anterior infarction, but coronary arteriography was normal apart from myocardial bridging over a segment of the left anterior descending artery. The development of acute myocardial infarction in this patient suggests that, in the presence of bridging, carbon monoxide inhalation may cause regional infarction.

Introduction

Acute carbon monoxide toxicity causes tissue hypoxia and evidence of diffuse myocardial damage. In patients with pre-existing coronary artery disease exposure to carbon monoxide may result in acute myocardial infarction. We report a case of acute myocardial infarction following smoke inhalation in which the only coronary artery abnormality was a segment of myocardial bridging. We postulate that such bridging is not always an innocent finding and played a part in the pathogenesis of this patient’s myocardial infarction.

Case report

A previously healthy 26 year old woman, without significant coronary risk factors, was brought to the Accident and Emergency Department in a private car having been rescued from a house fire by her neighbours. On arrival at hospital she was in severe respiratory distress and immediately suffered a cardio-respiratory arrest. She was found to be in ventricular fibrillation and received a 200 joules shock before reverting to a sinus tachycardia. Initially she was deeply unconscious with reduced tendon reflexes and widespread respiratory wheezes.

Following the administration of 60% oxygen by face-mask she recovered consciousness and complained of chest pain radiating to both elbows. An electrocardiograph demonstrated right bundle branch block and ST segment elevation in V2 to V6. She received glyceryl trinitrate and diamorphine, and an infusion of streptokinase (1.5 mU over 60 minutes). During the infusion there was an episode of sustained ventricular tachycardia which resolved after lignocaine was given. Although the ST segment changes resolved rapidly there was evidence of a Q-wave myocardial infarction and the cardiac enzymes were elevated (peak creatine kinase (MB)>155 ng/ml). Her remaining in-hospital course was further complicated by chest infection and she was discharged 15 days after admission.

On assessment 3 months later, a gated isotope study showed an ejection fraction of 54%. She completed stage 4 of the standard Bruce protocol exercise test without evidence of exercise-induced arrhythmias or ischaemia. A left ventriculogram showed a small area of anterior akinesis (Figure 1). The coronary arteries appeared free of atheroma but there was a segment of myocardial bridging over the middle third of the left anterior descending coronary artery (Figure 2). By projecting the angiographic image onto a large screen, and using calipers to allow accurate measurement, there was shown to be a 36% reduction in coronary diameter (systolic frame compared to diastolic frame) at the site of the bridging.

Discussion

Up to 90% of deaths in victims of fire are related to smoke inhalation, rather than burns, and the
Figure 1 Left ventricular angiogram (right anterior oblique projection) demonstrating anteropapical akinesis: (a) diastole; (b) systole.

Figure 2 Left coronary angiogram (left anterior oblique projection) demonstrating myocardial bridging (arrowed) over the middle third of the left anterior descending artery: (a) diastole; (b) systole.

The majority of these are attributable to carbon monoxide toxicity. This toxicity is mediated through tissue anoxia. The reduction in atmospheric oxygen concentration near a fire leads to decreased blood oxygenation and the formation of carboxyhaemoglobin, and carboxymyoglobin causing impaired oxygen delivery to the tissues. Furthermore, cytochrome oxidase activity is reduced by both hydrogen cyanide and carbon monoxide. This latter effect impairs the ability of the heart to recover from transient ischaemia.

The histological findings in the heart after acute carbon monoxide poisoning are diffuse focal areas of necrosis, inflammatory cell infiltrates and punctate haemorrhages. These changes are most marked in the subendocardium of the left ventricle (which is at greatest risk of ischaemia) leading to reversible abnormalities of left ventricular wall motion without evidence of segmental infarction and in the papillary muscles leading to mitral valve prolapse.

Electrocardiographic changes due to this diffuse myocardial damage include sinus tachycardia and ST segment depression; ST elevation is rare. In case reports of acute myocardial infarction after carbon monoxide inhalation the victims have been
presumed to have pre-existing coronary artery disease, though this has not always been proven angiographically.2,11

The young woman reported here suffered an acute Q-wave anterior myocardial infarction, rather than diffuse damage, after smoke inhalation, though there was no preceding history of chest pain or evidence of coronary artery disease. The possible causes of the infarction are coronary vasospasm, primary coronary thrombosis or some contribution from the documented myocardial bridging.

Although the early literature describes a state of 'acute arteriosclerosis' after severe carbon monoxide poisoning, with arterial spasm in extremities and brain,12 it has been shown in canine models that exposure to carbon monoxide results in coronary artery vasodilation and increased myocardial blood flow.13

There is evidence of an increased thrombotic state consequent to acute carbon monoxide inhalation. Total plasma volume is reduced (and vascular permeability increased) after exposure to the gas.14 Carbon monoxide also causes polycythaemia and increased platelet aggregability.15,16

Myocardial bridging is reported in 0.5–5.0% of coronary angiograms,17–19 most commonly affecting the left anterior descending coronary artery and being most obvious in cases of left ventricular hypertrophy and following administration of nitrates or inotropic agents.20 Its role in the production of myocardial ischaemia is controversial. It has been estimated that systolic arterial narrowing may reduce coronary blood flow by only 5–30%.20 A study of 21 patients with typical myocardial bridging demonstrated no evidence of myocardial ischaemia at rest or on exercise.21

However, there are reported cases of myocardial infarction associated with myocardial bridges22,23 and of the relief of chest pain following the resection of periartrial muscle.24 A patient has been described in whom acute coronary occlusion was treated by intracoronary streptokinase. When patency was restored myocardial bridging was demonstrated at the site of the occlusion.23 Moreover, a patient has been reported in whom rapid atrial pacing resulted in coronary spasm at the site of a myocardial bridge, and distal to it.25

We believe that the pathophysiology of the acute myocardial infarction described above involved both the tissue hypoxia secondary to smoke inhalation and the effect of the myocardial bridge on coronary blood flow. What would be in other circumstances a minor reduction in blood flow became a crucial factor at a time of reduced oxygen delivery.

References


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