Primary aortic thrombus in young: a rare site, a novel treatment

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To cite: Raghu TR, Agrawal N, Hegde M, et al. BMJ Case Rep Published online: [please include Day Month Year] doi:10.1136/bcr-2013-010304

DESCRIPTION
A 35-year-old man previously asymptomatic, non-smoker presented with a history of acute onset severe ischaemic pain, paraesthesia and pulselessness involving the left upper limb from the axillary artery distally since 3 days, not preceded by fever or chest pain. ECG was normal and transthoracic echocardiogram showed a normally functioning heart with an abnormal mass in the arch of aorta, which could not be characterised adequately. There was no aortic regurgitation.

CT aortogram revealed a large, immobile thrombus fixed in the arch of aorta causing subtotal occlusion of the ostium of the left subclavian artery, with the rest of the aorta having no evidence of atherosclerosis or aortitis and the coronaries were normal (figures 1–3).

The subsequent workup including analysis for dyslipidaemia, protein C and protein S and antiphospholipid antibody, erythrocyte sedimentation rate, C reactive protein, rheumatoid factor and antinuclear antibodies did not reveal any obvious predisposing factor. There was no historical or investigational evidence suggestive of any malignancy, connective tissue disorder or claudication or any family history of predisposition.

The site of thrombus was not amenable for surgical extraction without significant risks. The patient was subsequently thrombolysed with an infusion of streptokinase because of limb-threatening severe ischaemic pain and pulselessness. Heparin and anticoagulants have been shown to prevent thrombus propagation but its efficacy in dissolution of previously formed thrombus is uncertain. The patient had transient left hemiparesis persisting for 3–4 h during lysis, which was stopped immediately. The reason of selective embolisation to the right carotid from a relatively distant site could not be explained.

There was subsidence of all ischaemic symptoms in the limb and the echocardiogram showed no mass in the aorta and CT of the brain was normal. The patient was discharged on the fourth day on oral anticoagulation which was planned to be continued for at least 6 months. Statins and angiotensin-converting enzyme inhibitors were not

Figure 1 Coronal view of a large thrombus attached to the descending aorta at the origin of the subclavian artery.

Figure 2 Sagittal view of a large thrombus attached to the descending aorta at the origin of the subclavian artery.

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considered as there was no evidence of atherosclerosis and coronary involvement.

Subsequent follow-up since the last 5–6 months has been uneventful. CT aortogram was not been performed during follow-up as the patient was asymptomatic and unwilling.

This unique combination of presence of an aortic thrombus at an unusual site which is subjected to high blood flow and turbulence without any predisposing factor, responsive to thrombolysis has not been reported previously in the literature.1–3

Learning points

▸ Descending aorta can be a rare site of idiopathic thrombus, which can rarely be fixed. CT and blood workup can help in aetiology assessment.
▸ Thrombolysis could provide a successful non-surgical modality of treating thrombus in the aorta, which is not amenable to surgical access in patients having no other contraindications for lysis.
▸ Risk of stroke and peripheral embolism should be taken into account, especially in thrombus in proximity to major end arteries or arteries supplying vital organs.

Competing interests None.
Patient consent Obtained.
Provenance and peer review Not commissioned; externally peer reviewed.

REFERENCES