Severe aortic complications in a patient with variant Shone’s complex and bicuspid aortic valve

Mashrafi Ahmed, Hany Aziz, Leng Jiang

DESCRIPTION

A 47-year-old man presented with new onset exertional dyspnoea. He received a surgical repair of coarctation of the aorta (COA) at the age of 21 years and aortic root grafting with mechanical aortic valve replacement for type A aortic dissection and bicuspid aortic valve (BAV) at the age of 28 years. He had been otherwise healthy and did not follow-up with a cardiologist or surgeon after his last surgery. An echocardiogram (Echo) revealed normal left ventricular size and wall thickness, with an ejection fraction of 45%–50%. There was a parachute mitral valve (MV) (figure 1), with only one severely underdeveloped lateral papillary muscle. The mechanical aortic valve was functioning normally. There was a massively dilated distal aortic arch and proximal descending aorta with maximum dimension of up to 65 mm (figure 2). There was no Doppler evidence of recurrent COA. The CT angiogram confirmed the giant aneurysm, which was absent from the previous site of COA repair (figure 3). With the current Echo finding of parachute MV and history of COA, he was diagnosed with variant Shone’s complex. He then underwent a stent graft repair of the aneurysm.

Shone’s complex is a rare congenital disorder consisting of four obstructive lesions of the left side of the heart which are (1) supra-MV membrane, (2) parachute MV, (3) subaortic stenosis and (4) COA. Since its initial reporting in 1963, a limited number of cases had been reported in English literature. In contrast to the classic Shone’s complex, incomplete forms with two or three lesions were also reported as variant or partial Shone’s complex. Our case is a variant Shone’s complex with parachute MV and COA.

Most of the cases of Shone’s complex were diagnosed during childhood with left-sided outflow obstruction with resultant downstream left heart underdevelopment. Patients were usually symptomatic during early childhood. Typical symptoms include dyspnoea, tachypnoea, poor feeding, failure to thrive, fatigue and signs and symptoms of heart failure and low output syndrome. Recurrent episodes of wheezing and respiratory tract infections with pulmonary congestion and acute pulmonary oedema had been reported as well. Transthoracic Echo was used as diagnostic and follow-up tool. Generally, patients with Shone’s complex undergo a number of left heart interventions, both surgical and transcatheter approaches. COA was associated with an increased likelihood for repeat interventions. The 10-year transplant-free survival was 86%. The presence of...
pulmonary hypertension was associated with increased mortality. Significant left heart structural growth was noticed after surgical intervention. In clinical practice, the definition of Shone’s complex has been broadened beyond the original description. BAV has been increasingly reported in patients with Shone’s complex. Since the first case series published in 1963, subsequent multiple case series reported BAV as an associated feature of Shone’s complex. Our case emphasises that the association of BAV with Shone’s complex should not be neglected since BAV is a disease of both the valve and the aorta. The associated aortopathy may cause severe life-threatening complications as shown in our case. Therefore, for patients with variant Shone’s complex and BAV, cardiac imaging of the thoracic aorta should be performed periodically for surveillance.

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**REFERENCES**

**Learning points**
- Shone’s anomaly is a very rare congenital cardiac malformation characterised by four serial obstructive lesions of the left side of the heart, namely (1) supravalvular mitral membrane, (2) parachute mitral valve, (3) muscular or membranous subaortic stenosis and (4) coarctation of the aorta.
- Although complete Shone’s complex is diagnosed relatively early in childhood, the variant form, characterised by two or three components, may remain undiagnosed owing to the variable severity of each pattern.