A Case of Right Sided Aortic Arch with Kommerell Diverticulum

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Abstract

Introduction: Kommerell diverticulum, a rare embryological variant, is an aneurysm of aortic arch at the origin of an aberrant subclavian artery. It is usually asymptomatic and detected incidentally on imaging studies or postmortem. Common symptoms include dysphagia and chest pain. The most common complications include rupture due to an enlarged diverticulum.

Case Report: We present a male patient evaluated for chronic chest pain following traumatic blunt injury, who was incidentally found to have right sided aortic arch with kommerell diverticulum.

Conclusion: Basic investigations like chest X-ray helps in clinching the diagnosis. Vascular anomalies like kommerell diverticulum should be considered as differential diagnosis in the evaluation of chronic cough, dysphagia, dyspnea.

Key words: Right Aortic Arch; Aberrant Subclavian Artery; Kommerell's Diverticulum

Introduction

Right aortic arch with aberrant left subclavian artery is a rare congenital defect of thoracic aorta. It is present in 0.05% to 0.1% of radiology series. This condition is asymptomatic unless aneurysmal disease develops [1]. Kommerell's aneurysm, which was first described by Burchard Kommerell in 1936 in a living patient [2].

Case Report

A 47-year-old male patient presented with a one-year history of continuous, dull aching, non-exertional left sided chest pain. He had no previous comorbid illness. Physical examination was normal except with features suggestive of periarthritis left shoulder showed normal sinus rhythm and no ST-T changes.
Figure 2a: CT Chest showing right sided aortic arch a) At the level of superior mediastinum.

Figure 2b: a CT SCAN chest right sided aortic arch b) At the level of pulmonary artery.

Figure 2c: T Scan chest at the level of ventricles of heart.

Figure 2d: CT Chest right sided aortic arch at the level of the Liver.
Discussion on Embryology of Aorta

Development of aorta occurs in third week of gestation with sequential formation of six pairs of arterial arches connecting the dorsal and ventral aorta. Selective regression and persistence of these arch vessels forms the major arteries of the body. 1st, 2nd, 5th arterial arches disappear [3].

- 3rd arch common carotid artery.
- 4th arch-
  - Right- Proximal part of right subclavian artery.
  - Left - Part of aortic arch between left carotid artery and left subclavian artery.
- 6th arch forms Pulmonary arteries and ductus arteriosus.
Right proximal portion of 6th arch forms proximal segment of right pulmonary artery.

Left proximal portion of 6th arch forms proximal segment of left pulmonary artery while distal part of left 6th arch forms ductus arteriosus.

**Discussion**

Right aortic arch is a developmental anomaly of the thoracic aorta, caused by persistence of the right fourth branchial arch during embryogenesis. It is associated with deletion of 22q11 chromosome in 24% cases. It was first documented by Fioratti and Aglietti in 1763 [4]. This has been classified by Edward in 1948, Felson and Palayew in 1963 and Steward et al in 1964 [5,6].

**Edward's Classification [7]**

**Type 1**: Right sided aortic arch with mirror image branching due to interruption of dorsal segment of left arch between left subclavian artery and descending aorta with regression of right ductus arteriosus in hypothetical double aortic arch.

**Type 2**: Right aortic arch with aberrant left subclavian artery, due to interruption of dorsal segment of left arch between left common carotid artery and left subclavian artery with regression of right ductus arteriosus.

**Type 3**: Right aortic arch with isolated left subclavian artery, due to interruption of arch at two levels, one level between left common carotid and subclavian artery, other between distal to the attachment of left ductus. It is usually associated with verteobasilar insufficiency.

A type 1 and 2 form 98% of the right sided aortic arch. Type 1 right aortic arch is usually occurs with cyanotic congenital heart diseases like Tetralogy of Fallot, Truncus arteriosus [8].

Type 2 right sided aortic arch usually remains asymptomatic and there is no association with cardiac anomalies. However, more symptoms are due to atherosclerotic changes of the anomalous vessels, dissection, aneurysm with compression of adjacent structures causing dysphagia (dysphagia lusoria), dyspnea. As our patient has no symptoms related to the vascular variations, surgery was not indicated [4,9].

The aberrant subclavian artery usually originates from a kommerell's diverticulum, defined as conical dilation of the proximal portion of an aberrant subclavian artery near its origin from the aorta. It is also known as “lusoria diverticulum” [4]. It is the remnant of the left fourth aortic arch in the aberrant left subclavian artery, being located behind the esophagus in 80%, between the trachea and esophagus as in our case in 15% and behind the trachea in 5% [10]. Complications include atherosclerosis, embolism, rupture and dissection [1,10]. The mean size associated with rupture is 5.8cm. Investigations include barium swallow esophagogram, CT scan with contrast, cardiac MRI which demonstrates both diverticulum and its effects on trachea and esophagus [1,9].
Surgical Treatment

It is treated by excising the kommerell's diverticulum and anastomosis of the left subclavian artery to the left common carotid artery [11].

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