Aberrant Right Subclavian Artery: Cadaver Case Report

Jason Kopp,1 Ahmad Irshaid,1 Justin Baker,2 John Fitzsimmons,3 Judith C. Lin.4

Abstract

Different aortic arch branching patterns exist in the general population. These branching patterns can be benign or can cause a variety of symptoms in patients. In the case of a more benign branching pattern, anomalies often go undiagnosed until discovered postmortem. Case: While examining the anatomy of a cadaver in a medical school gross anatomy course, an aortic arch anomaly was discovered. This variant is consistent with an aberrant right subclavian artery (ARSA). In this variant, the right subclavian artery branches from the most distal part of the aortic arch and runs both retrotracheal and retroesophageal as it courses to the right shoulder. This variant is a result of aberrant development of the aortic arch and may present with symptoms such as dysphagia and shortness of breath, if any at all. In addition to the ARSA, there exist a common bicarotid trunk and a direct branching of the left vertebral artery from the aortic arch, both of which are rare anomalies. The cadaver’s medical history includes dysphagia and stretched esophagus, although the severity is unknown. Conclusion: this case report draws attention to these rare anatomical anomalies and includes a discussion of the most common clinical presentation, and surgical implications of an aberrant right subclavian artery anomaly

Key Words: Aberrant Right Subclavian Artery; Case Reports; Aortic Arch; Dysphagia (Source: MeSH-NLM).

Introduction

The aortic arch is a critical structure in the cardiovascular system, as it is the beginning of the systemic arterial circulatory system. Formation of the aortic arch begins during the fourth week of embryonic development and is ultimately derived from multiple structures. At this time, a structure called the aortic sac starts to develop. During the fifth week of development, the aortic sac begins to grow and branch off into the two dorsal aortas and the ventral aorta. Six paired pharyngeal aortic arches develop, which connect the ventral and dorsal aortae. Some of these arches completely regress while portions of the other persist as the mature aorta develops. The primary origins of the aortic arch, from proximal to distal, are the aortic sac, the left fourth aortic arch, and the dorsal aorta.

An aberrant right subclavian artery (ARSA) affects approximately 1% of the population. ARSA develops as a result of the abnormal regression of the right fourth aortic arch and right dorsal aorta distal to the right common carotid artery. In this case, the aberrant right subclavian artery branches from the right seventh segmental artery of the descending aorta and becomes the most distal branch of the aortic arch. The aberrant right subclavian artery passes retrotracheal and retroesophageal in 80% of the cases. Alternatively, it courses between the trachea and the esophagus in 15% of the cases, and anterior to the trachea in 5% of the cases. In addition to an aberrant right subclavian artery, there can also be the occurrence of a bicarotid trunk. The coexistence of both ARSA and the bicarotid trunk has an estimated prevalence of <0.05%.

A majority of individuals with an aberrant right subclavian artery are asymptomatic, and it is estimated that 60-80% of individuals with this anomaly will never develop any symptoms. For those who do develop symptoms, it is more common for them to have dysphagia, caused by mechanical compression of the esophagus by the aberrant artery. This is referred to as dysphagia lusoria. Dysphagia lusoria is seen in 71.2% of symptomatic individuals, but can range from only mild intermittent dysphagia to a potentially severe, and continuous dysphagia of both solids and liquids. These symptoms usually appear in the middle to older age groups. However, the exact reason for this is unclear. Other
symptoms and their prevalence in symptomatic individuals include dyspnea (18.7%), retrosternal pain (17.0%), cough (7.6%), and weight loss (5.9%). This case report highlights a patient with a right aberrant subclavian artery and a bicarotid trunk who experienced dysphagia. Uniquely, our report includes gross anatomical imaging and discusses the clinical implications of such an anomaly.

The Case
While examining cadavers as part of a medical school gross anatomy course, an anomalous origin of the right subclavian was discovered. The variant was consistent with the aberrant right subclavian artery, as previously described. In this case, the right subclavian artery originates from the aortic isthmus rather than the brachiocephalic trunk.

Figure 1 depicts the presentation of the aortic arch and its branches in our cadaver. The left subclavian artery and left vertebral artery branch separately travel to the left shoulder and head, respectively. The right and left common carotid arteries arise from a common bicarotid trunk prior to coursing up the lateral portion of each side of the neck.

Figure 1. Schematic Depiction of the Aortic Arch and its Branches in the Cadaver.

Discussion
During a routine examination of cadavers as part of a gross anatomy course, an aberrant right subclavian artery was discovered. This specific aortic arch anomaly is only present in about 1% of people. However, in addition to the aberrant right subclavian artery, there was also the occurrence of a bicarotid trunk, which is an extremely rare variant. The coexistence of both ARSA and the bicarotid has an estimated prevalence of <0.05%. Lastly, the left vertebral artery arises from the aortic arch, which is estimated to occur in around 4% of all individuals. Nevertheless, the combination of all three has not been well documented. Therefore, the incidence cannot be determined in this report.

ARSA variations have been reported in the literature and Adachi-Williams have classified them into three types based on the branches coming off of the aortic arch. Type I (Type-G) has four branches: the right common carotid artery (RCCA), left common carotid artery (LCCA), left subclavian artery (LSA), and aberrant right subclavian artery (ARSA). Type II (Type CG) is the same as type I along with the addition of a branch for left vertebral artery (LVA). Lastly, type III (type H) is seen with a bicarotid trunk, LSA,
and ARSA branches coming from the aorta. The case seen here is a combination of type II and type III variations due to the presence of four branches: the LVA (seen in type II) and the bicaortic trunk (seen in type III), along with LSA, and ARSA.

A recent systematic review, aimed at categorizing left sided aortic arch variants, classified the variants based on the number of emerging branches into type 1b (one branch), 2b, 3b, 4b, 5b and further subclassified based on prevalence. The systemic review concluded that a typical branching pattern had a prevalence of 78% with a 22% prevalence for all other branching patterns. The most common variant was 2b, with bicaortic trunk (LCCA originating from the brachiocephalic arch) being the most common subtype of the 2b variant.

This case report is especially useful for examining an aortic arch anomaly and uniquely presents gross anatomical findings as evidence of the anomaly. This report also includes rare evidence of an ARSA coexisting with a bicaortic trunk. An associated history of dysphagia in the cadaver helps demonstrate possible clinical findings of a patient with an aortic arch anomaly. Due to the nature of the case report, we were unable to examine the patient while they were still living and experiencing symptoms. The findings of the case report were limited to the patient’s brief medical history reported to the anatomy lab. Other cadaver case reports have reported similar findings of an ARSA, and included more information on the clinical severity of the patient’s anomaly. However, future cadaver case reports should include thorough and complete clinical descriptions of patients’ dysphagia as well as record of physician assessments and surgical recommendations made during the clinical course. Diagnosing an ARSA in a living patient is rare and is usually accidently discovered on a coronary angiogram. Surgery may be implicated in extreme cases of dysphagia but is usually foregone in favor of supportive treatment due to the risks of the procedure.

Individuals with ARSA variations can be asymptomatic and those with symptoms often do not present until later in life, which makes a clinical diagnosis challenging. Despite this, it is important that physicians consider an ARSA anomaly when a patient presents with dysphagia, shortness of breath, and/or stridor. Other concerning symptoms for a suspected ARSA include worsening retrosternal pain, cough, and acute limb ischemia. Although a surgical intervention is unlikely except for in rare and severe cases, this case report raises clinical awareness of the possibility of such a condition and can guide more specific intervention and treatment.

Summary – Accelerating Translation

The title of this case report is Aberrant Right Subclavian Artery: Cadaver Case Report. The aorta is the major blood vessel leaving the heart and provides oxygenated blood to the tissues. The aortic arch gives off 3 branches in 78% of the population. Normal and pathological variation in the branching patterns and in the number of branches exists in the general population. This case presentation discusses an aberrant right subclavian artery (ARSA) in a cadaver. In the normal variant, the right subclavian artery branches off the brachiocephalic artery, the most proximal branch of the aortic arch. In our case, the right subclavian artery is the most distal branch of the aortic arch and branches directly off of it as it courses behind the trachea and esophagus to the right shoulder.

Most patients with this anomaly are generally asymptomatic, and the few that develop symptoms tend to develop them later in life. Trouble swallowing, also called dysphagia lusoria, is the most common symptom associated with ARSA. The abnormal position and path of the right subclavian artery mechanically compresses the esophagus causing the dysphagia. Other less common symptoms include shortness of breath, chest pain, cough, and weight loss. Although interventions are generally rare, it’s important for physicians to consider ARSA when working up a patient with dysphagia, and to be aware of the variation in the branching patterns of the aortic arch when planning for cardiothoracic interventions.

References

Acknowledgments
None.

Conflict of Interest Statement & Funding
The Authors have no funding, financial relationships or conflicts of interest to disclose.

Author Contributions

Cite as

This work is licensed under a Creative Commons Attribution 4.0 International License
ISSN 2076-6327
This journal is published by Pitt Open Library Publishing