Coexistence of an aberrant right subclavian artery and anomalous origins of bilateral vertebral arteries

A case report

Yihong Wu, MSa, Huayi Zhang, MSb, Chenye Tang, MSc,*

Abstract

Rationale: Anatomical variations in aortic arch (AA) branching are not unusual. Generally, these variations are asymptomatic and are diagnosed incidentally. Here, we report a rare case of a middle-aged female patient with an aberrant right subclavian artery (ARSA) associated with anomalous origins of the bilateral vertebral arteries (VAs).

Patient concerns: The patient treated for urolithiasis complained of repeated dizziness for several years.

Diagnoses: Echocardiography and computed tomography angiography (CTA) confirmed arterial variations. Moreover, mild stenosis was found in the left common carotid artery (LCCA), which was considered to be the cause of dizziness.

Interventions: Congenital anomalous arteries were not necessary to intervene urgently, but aspirin and atorvastatin were administered to prevent potential thrombosis attributed to vascular stenosis after completing the operation for urolithiasis.

Outcomes: Whether the symptoms will be alleviated or not should be continuously followed up, and the patient may accept interventional therapy in the future if necessary.

Lessons: Here, we report the rare variation of AA branches and highlight the importance of preoperative vascular assessment in surgical or interventional procedures for the affected body regions.

Abbreviations: AA = aortic arch, ARSA = aberrant right subclavian artery, CTA = computed tomography angiography, IA = innominate artery, LCCA = left common carotid artery, LSA = left subclavian artery, LVA = left vertebral artery, RCCA = right common carotid artery, RSA = right subclavian artery, RVA = right vertebral artery, SA = subclavian artery, VA = vertebral artery.

Keywords: aberrant right subclavian artery, aortic arch, case report, variation, vertebral artery

1. Introduction

The standard anatomy of the aortic arch (AA) includes 3 branches arising from its superior border from right to left and includes the innominate artery (IA), which then branches into the right subclavian artery (RSA) and the right common carotid artery (RCCA), the left common carotid artery (LCCA), and the left subclavian artery (LSA). As an essential source of blood supply to the posterior circulation, the vertebral artery (VA) usually arises from the posterosuperior aspect of the first part of the ipsilateral subclavian artery (SA). It then travels obliquely to enter the transverse foramen of the C6 vertebra. According to several large-sample studies, 65.9% to 79.2% of the population had a typical AA branching pattern and VA origin, whereas 20.8% to 34.1% had variations. Aberrant right subclavian artery (ARSA) is one of the most abundant AA branching anomalies, with an incidence rate of 0.5% to 1.3%. Anomalous origin of VA is more frequent on the left (6.0%) than on the right side (3.8%), and quite rarer on the bilateral side (approximately 3.0% of all the patients with anomalous origin of VA). Herein, we present the case of a patient with an ARSA associated with anomalous origins of bilateral VAs.

2. Case presentation

A 43-year-old female patient was treated for urolithiasis in the Department of Urology, who complained of repeated dizziness for several years. Preoperative echocardiography showed an ARSA derived from the initial descending aorta right rear to the LSA and an aberrant left vertebral artery (LVA) that directly originated from the AA instead of the LSA (Fig. 1). Then, computed tomography angiography (CTA) was performed, not only revealing the aforementioned variations of the arteries but also confirming an aberrant right vertebral artery (RVA) that
originated from the RCCA rather than the RSA (Fig. 2A, B). The
diameters of the initial segment of ARSA, LVA, and RVA were
about 8.35, 4.14, and 3.22 millimeters, respectively, and no
stenosis was seen in any course of these 3 arteries. The ARSA run
behind the trachea and esophagus to the right side of the body;
the LVA and RVA ascended upward and then entered the
foramen transversarium of the C4 and C5 vertebrae, respectively.
Moreover, mild stenosis was found in the proximal portion of the

---

**Figure 1.** Echocardiography showing the variations of AA branches.

---

**Figure 2.** CTA showing the variations of AA branches and the stenosis in LCCA. (A). The 5 branches of the AA (anterior view) and the stenosis in LCCA. (B). The 5 branches of the AA (left lateral view) and the origin of RVA from the RCCA.
LCCA, adjacent to its aortic origin (Fig. 2A). No other cardiac or vascular anomalies were found on the above-mentioned imageological examinations. To prevent potential thrombosis attributed to vascular stenosis, aspirin and atorvastatin were used after completing the operation for urolithiasis, and the patient was followed up continuously. This study was approved by the Ethics Committee of the Second Affiliated Hospital of Jiaxing University, and written informed consent was obtained from the patient.

3. Discussion and conclusions

Variations in AA branching are usually asymptomatic and detected accidentally by autopsy,[6,7] or imagological examinations performed for other causes.[8,9] In the study by Karacan et al.[13] the AA branching patterns were categorized into 7 types, not including the form of our patient. Only an ARSA was defined as type 5 variation, while coexistence of ARSA and bicarotid trunk was classified as type 6 variation, and their incidences were 0.6% and 0.7%, respectively; meanwhile, the frequency of these 2 types was higher among females than among males.[10] The ARSA results from the embryological degeneration of the right dorsal aorta above the seventh intersegmental artery (generally below the seventh intersegmental artery).[10] The majority of ARSAs (80%) run behind the esophagus and can sometimes cause compression of the esophagus, making it difficult to swallow.[11] Less frequently, the ARSA may travel between the esophagus and trachea (15%) or before the trachea (5%), potentially resulting in cough or airway obstruction.[11] When a bulb-like swelling is present in the proximal portion of the SA near its origin, it is called a Kommerell diverticulum. Although uncommon, it may occur in approximately 60% of patients with ARSA.[12] Once it occurs, the symptoms mentioned above may be more prominent. Additionally, catastrophic consequences may occur, such as distal embolism and diverticular rupture. Therefore, such cases should be actively treated by surgery, if possible. Fortunately, Kommerell diverticulum was not present in our patient.

The LVA directly originating from the AA between the LCCA and LSA seen in our patient is the most common variation of VA origin,[5,8] and was defined as type 3 variation in the series categorized by Karacan et al.[13] If combined with IA and LCCA arising from the AA in a common trunk, the AA branching pattern was categorized as type 4 variation, and the incidence rates of type 3 and 4 variations were 4.1% and 1.2%, respectively.[13] Variations in RVA origin are relatively rare, and the most common is an RSA originating from the extreme proximal RSA; fewer cases of RVA arising from the RCCA are seen in patients with ARSA, with a reported incidence rate of 0.1%.[13] This rare variation in arteries was also observed in this case. Embryologically, the first or second intersegmentary artery failing to regress generates an abnormal origin of the VA from the internal or external carotid, while the permanent artery occurring from the third to sixth intersegmental artery results in an aberrant VA arising from the AA or common carotid artery.[10] Variations in VA origin do not result in clinical symptoms; however, a direct aortic origin of the VA may cause hemodynamic changes that may lead to related complications. Compared with the classic subclavian origin of either the LVA or RVA, an aortic origin of the LVA showed an increased risk of aortic dissection.[14] and it may be an independent risk factor for arterial dissection.[13] Some researchers have postulated that a longer extracranial course may make the vessel wall more vulnerable to shear force, which may lead to intimal tear and dissection.[13] Therefore, if headache and neurological symptoms occur suddenly in a patient with an anomalous VA originating from the AA, a thorough examination should be performed. It is not clear whether VA arising from the common carotid artery would increase the rate of complications when compared with the normal subclavicular origin of VA. Our patient had dizziness, which was thought to have no connection to the anomalous origins of the VAs. It is possible that stenosis of the LCCA was accountable. Conservative medication and long-term follow-up rather than operations were recommended because the atherosclerosis was partial and mild.

In conclusion, the coexistence of an ARSA and anomalous origins of bilateral VAs is an infrequent condition. Misidentification of variant branches may lead to life-threatening complications during surgical or endovascular procedures. Therefore, careful examination and advance knowledge of the anatomic variants are essential for scheduling surgical or interventional procedures involving the thorax, neck, and head.

Author contributions

Case discussion and interpretation: Huayi Zhang.
Conceptualization: Yihong Wu, Chenye Tang.
Validation: Yihong Wu, Huayi Zhang, Chenye Tang.
Writing – original draft: Yihong Wu.
Writing – review & editing: Huayi Zhang, Chenye Tang.

References

[1] Berkos NS, Jain VR, Godelman A, et al. Variants and anomalies of thoracic vasculature on computed tomographic angiography in adults. J Comput Assist Tomogr 2017;41:1105–6.
[2] Rea G, Valente T, Iselli F, et al. Multi-detector computed tomography in the evaluation of variants and anomalies of aortic arch and its branching pattern. Ital J Anat Embryol 2014;119:180–92.
[3] Karacan A, Türkvatan A, Karacan K. Anatomical variations of aortic arch branching: evaluation with computed tomographic angiography. Cardiol Young 2014;24:485–93.
[4] Uchino A, Saito N, Takahashi M, et al. Variations in the origin of the vertebral artery and its level of entry into the transverse foramen diagnosed by CT angiography. Neuroradiology 2013;55:885–94.
[5] Yuan SM. Aberrant Origin of Vertebral Artery and its Clinical Implications. Braz J Cardiovasc Surg 2016;31:52–9.
[6] Singh R. Two rare variants of left vertebral artery. J Craniofac Surg 2017;28:1105–6.
[7] Ma Z, Han J, Li H, et al. A unique variation with five branches of the aortic arch. Interact Cardiovasc Thorac Surg 2018;26:165–6.
[8] Liu YD, Li QZ, Fu JJ, et al. A rare anomalous origin of right vertebral artery with double branch: first case report. Interv Neuroradiol 2018;24:225–8.
[9] Silveira JV, Junqueira FP, Silveira CG, et al. Kommerell diverticulum: right aortic arch with anomalous origin of left subclavian artery and duplicity of right vertebral artery in a 16-Year-old girl. Am J Case Rep 2019;20:228–32.
[10] Kesler WW, Sabat SB. Isolated anomalous origin of the vertebral artery from the common carotid artery. Interact Cardiovasc Thorac Surg 2018;27:615–6.
[11] Guzman ED, Eagleton MJ. Aortic dissection in the presence of an aberrant right subclavian artery. Ann Vasc Surg 2012;26:860.e13–18.
[12] Ota T, Okada K, Takahashi S, et al. Surgical treatment for Kommerell’s diverticulum. J Thorac Cardiovasc Surg 2006;131:574–8.
[13] Moshahedi P, Walker GB, Tavakoli S, et al. Dual origin of the right vertebral artery from the right common carotid and aberrant right subclavian arteries. J Clin Neurosci 2018;53:238–40.
[14] Komiyma M, Morikawa T, Nakajima H, et al. High incidence of arterial dissection associated with left vertebral artery of aortic origin. Neurol Med Chir (Tokyo) 2001;41:8–12.
[15] Dudich K, Bhadelia R, Senivasan J. Anomalous vertebral artery origin may be an independent risk factor for arterial dissection. Eur J Neurol 2005;12:571–2.