Anomalous origin and aneurysm of the right coronary artery associated with congenital bicuspid aortic valve: MDCT findings

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Abstract

Anomalous origin of the coronary artery taking an interarterial course can cause myocardial infarction or sudden death. Association of anomalous origin of the coronary artery with congenital bicuspid aortic valve is rare, and only a few cases have been reported with imaging findings. Coronary artery aneurysms found in young adults are usually non-atherosclerotic. We report MDCT findings of anomalous origin and aneurysm of the right coronary artery associated with congenital bicuspid aortic valve in a 33-year-old man with a history of Kawasaki disease in the childhood, and the key role of MDCT in exact diagnosis and successful management of the complicated disease.

Keywords: Coronary artery anomaly, Coronary artery aneurysm, Bicuspid aortic valve, Kawasaki disease, MDCT

Background

Anomalous origin of the coronary artery from the wrong sinus of Valsalva and its course passing between the two great arteries has been considered potentially malignant. The association of anomalous origin of the coronary artery and bicuspid aortic valve is rare, and only a few cases with imaging findings have been reported in the literature (Aoyagi et al. 1991; Ayusawa et al. 2010). Coronary artery aneurysm that develops as a complication of Kawasaki disease in infants can remain silent until adulthood. We report multidetector computed tomography (MDCT) findings of anomalous origin and aneurysm of the right coronary artery (RCA) associated with congenital bicuspid aortic valve in a young man who had a history of Kawasaki disease in childhood.

Case report

A 33-year-old man underwent CT coronary angiography for the evaluation of coronary artery status. He was referred from another hospital where he had received low dose chest CT for health screening and was informed that coronary artery calcification was detected on CT. He did not complain of acute chest pain at this visit but he recalled having intermittent vague chest discomfort. He did not have any cardiovascular risk factors, but had a history of acute febrile illness at 2 years of age that was diagnosed as Kawasaki disease. Electrocardiography findings were nonspecific. MDCT was performed with a 128-slice dual-source CT system (Somatom Definition Flash; Siemens Healthcare, Forchheim, Germany) using the following parameters: detector collimation, 0.6 mm; slice acquisition, 2 × 128 × 0.6 mm; pitch, 0.25; 300 mAs; 100 kVp; gantry rotation time, 280 ms; temporal resolution, 75 ms. Intravenous nonionic contrast material [iomeprol, 400 mg/mL (Iomeron, Bracco)] was administered at a rate of 4–5 mL/s (total, 70 mL) followed by 30 mL of saline at a rate of 3 mL/s. The patient received 0.6 mg of nitroglycerin sublingually just before CT scanning and a beta-blocker was not administered. The effective radiation dose used in CT scanning was 4.9 mSv.

The volume rendering image of the heart revealed that the RCA arose from the sinotubular junction in between the left and right coronary sinus of Valsalva and coursed anteriorly between the aortic root and the pulmonary artery into the right atrioventricular groove (Fig. 1a). A saccular aneurysm in the proximal portion of the RCA was noted. Curved multiplanar reformation image of the
RCA showed an acute take-off angle of the RCA origin and focal significant narrowing at the proximal portion of the aneurysm (Fig. 1b). Axial images at the level of RCA origin showed further narrowing of the RCA orifice at the systolic phase (Fig. 2), but the anomalous coronary artery did not have an intramural course within the aortic wall. Bicuspid aortic valve had two completely developed cusps without raphe (Fig. 3a). Three-dimensional virtual angioscopic image of the aortic root showed a bicuspid aortic valve and normal ostium of the left main coronary artery. A slit-like orifice of the right coronary artery was located high at the sinotubular junction between the right and left coronary sinus (Fig. 3b) and was nearly effaced in the systolic phase (Fig. 3c). Echocardiography revealed bicuspid aortic valve with mild aortic regurgitation and normal left ventricular function. One month later, the patient underwent bypass graft surgery for the RCA with good recovery.

**Discussion**

Coronary anomalies are detected in about 1% of the general population by coronary angiography and have little clinical significance (Angelini et al. 2002). However, a minority of coronary artery anomalies, particularly in which the coronary artery takes an interarterial course, are known to have a risk of myocardial ischemia or sudden cardiac death (Rigatelli et al. 2005). Several possible causes of myocardial ischemia in cases with anomalous
coronary artery origin from the wrong aortic sinus with
a course between the aorta and the pulmonary artery
have been suggested: acute angle take-off of the coronary
artery producing a slit-like lumen; closure of the abnor-
mal coronary orifice by a valve-like ridge at aortic expan-
sion; compression of the artery when it courses within
the aortic wall (intramural course) or between the aorta
and the pulmonary artery; and spasm of the anomalous
coronary artery (Basso et al. 2000; Virmani et al. 1984).
Virmani et al. (1984), after observing postmortem coro-
nary arteries in victims of sudden death, postulated that
the presence of ostial valve-like ridges with acute angle
take-offs in coronary arteries cause ostial compression
with aortic dilatation. By acting as an occlusive valve,
the ostial valve-like ridges in acute take-offs may cause
coronary spasm and sudden death. In the present case,
MDCT showed narrowing of the RCA origin with an
acute take-off angle from the aorta, and ostial narrowing
was accentuated in the systolic phase although we could
not confirm the presence of an ostial valve-like ridge.

Our patient also had an aneurysm with mural throm-
bus and ring calcification in the proximal portion of the
anomalous RCA, which can be attributed to Kawasaki
disease. In Kawasaki disease, coronary artery aneurysms
may develop as a late complication and remain silent
until adolescence or young adulthood when myocardial
infarction or sudden death occurs (Okura et al. 2013).
Burns et al. (1996) performed a retrospective survey of
74 published cases of coronary artery disease attributed
to antecedent Kawasaki disease in young adults, and
stated that history of Kawasaki disease should be sought
in all young adults who present with acute myocardial
infarction or sudden death. Vasculitis of the coronary
vasa vasorum leads to endothelial damage and coro-
nary artery aneurysm formation, and slow flow within
the aneurysm predisposes to thrombus formation or

**Fig. 3** a 3D CT image in systole shows bicuspid aortic valve with a symmetric and elliptical orifice. Note the right (arrow) and left (long arrow) coro-
nary arteries originating from anterior cusp (A) formed by fusion of the right and left coronary cusp. b Three-dimensional virtual angiographic image
of the aortic root shows the bicuspid aortic valve (arrows), ostium of the left main coronary artery (L), and a slit-like orifice of the right coronary
artery (thin arrows) which is located high at the sinotubular junction. c In the systolic phase, the orifice of the RCA is nearly effaced (thin arrows).
atherosclerotic change (Okura et al. 2013; Burns et al. 1996). Moreover, in our patient, the narrow orifice in the presence of anomalous origin could also have produced hemodynamic alteration and further endothelial injury, contributing to acceleration of aneurysm formation and thrombosis in the RCA (De-Giorgio et al. 2011).

A bicuspid aortic valve is one of the most frequent congenital cardiac malformations in adults and it may be isolated or associated with other congenital heart diseases. There is increased variation in the coronary anatomy in patients with bicuspid aortic valves (Fedak et al. 2002). The association of anomalous origin of coronary arteries and congenital bicuspid aortic valve has only been sporadically reported in the literature (Aoyagi et al. 1991; Ayusawa et al. 2010). According to the observation of embryonic heart development, differentiation of aortic valve is completed during the similar period in which the proximal coronary arteries develop as endothelial outgrowth from the aortic sinus (Hutchins et al. 1988).

Thus, it could be postulated that a common developmental defect produced both anomalies in this patient. Some cardiac surgeons support the routine use of preoperative coronary angiography to assess the status of coronary arteries before aortic valve operation (Barriales-Villa et al. 2003). However, as seen in our case, technically advanced MDCT can replace coronary angiography. Cardiac MDCT can provide comprehensive anatomic information about this complicated anomaly through three-dimensional reconstruction, and also functional information of bicuspid aortic valve. In addition, aortic dilatation, which frequently occurs as a complication of the bicuspid aortic valve, can be evaluated by MDCT.

**Conclusion**

To the best of our knowledge, this is the first reported case of congenital bicuspid aortic valve associated with an anomalous origin of the RCA, complicated by late sequelae of Kawasaki disease and diagnosed with MDCT. MDCT is an effective and useful tool for exact assessment of coronary artery status and associated anomalies, resulting in timely management.

**Consent for publication**

Informed consent for publication of this report and any accompanying images was obtained from the patient.

**Abbreviations**

MDCT: multidetector computed tomography; RCA: right coronary artery; CT: computed tomography.

**Authors’ contributions**

Concept and design: KB. Acquisition of data: SC. Analysis and interpretation of data: KNJ. Drafting the manuscript: SC. Final approval: SC, KNJ, KB. All authors read and approved the final manuscript.

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**Compliance with ethical guidelines**

The authors declare that they have no competing interests.

**Competing interests**

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