Anomalous Right Coronary Artery from the Pulmonary Artery (ARCAPA) in a 63-Year-Old Male Patient with a Stable Angina Pectoris and Positive Stress Test

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ABSTRACT
Anomalous origin of the right coronary artery from the pulmonary artery (ARCAPA) is a very rare congenital cardiac anomaly. Herein, computed tomography and conventional coronary angiography imaging findings of the patient with ARCAPA are presented.

Key Words: Coronary angiography; Computed tomography angiography; Coronary veins

INTRODUCTION
Anomalous origin of the coronary arteries from the pulmonary artery (ACAPA) is a rare congenital anomaly which has two subtypes including right ACAPA (ARCAPA) and left ACAPA (ALCAPA) coronary arteries. The left sided anomalous origin of coronary artery (ALCAPA) is more commonly seen variation compared to ARCAPA with an incidence of one per 300,000 live births and 0.25% - 0.5% of all congenital cardiac diseases. However, ARCAPA is a rare congenital coronary anomaly with reported incidence of 0.002%, which constitutes 0.12% of all coronary artery anomalies [1–3].

Although it is more commonly encountered in the young population, some patients may remain undiagnosed until middle age. The timing of onset of symptoms in ARCAPA patients is variable and is related to the presence of collateral circulation [4]. Early symptoms may be secondary to heart failure, valve insufficiency or myocardial infarction [5]. Surgical treatment is recommended even if patients are asymptomatic due to their high risk profile [6].

The pathophysiology of ARCAPA depends on the direction of blood flow in the coronary artery and its ability to deliver oxygen to the myocardium. Since patients with right dominant blood circulation cannot tolerate ARCAPA well compared to those with left dominant system [2] Coronary steal phenomenon in ARCAPA may cause diastolic pressure differences between systemic and pulmonary artery beds and increased myocardial oxygen demand and risk of myocardial ischemia [7]. Although the symptoms of ARCAPA can range from effort angina to dyspnea or fatigue, it can also manifest itself in the form of ischemic myocardial damage sometimes causing sudden cardiac death [8,9].

In this article, the imaging findings of, ARCAPA with conventional coronary angiography (CCA) and coronary computed tomographic angiography (CCTA) of a 68-year-old male patient are presented.

CASE REPORT
A 68-year-old male patient applied to the hospital with a complaint of stable angina pectoris. There was no referred pain present. No remarkable finding was found on physical examination and electrocardiography. However due to the positive stress test of the patient, CCA examination was done, revealing that no coronary artery originated from the right sinus Valsalva. The left main coronary artery (LMCA) originated from the left coronary sinus, divided into circumflex (Cx) and left anterior descending artery (LAD) branches (Figures 2 & 3). These findings were evaluated as compatible with ARCAPA.

Figure 1) Conventional coronary angiography: The left main coronary artery originating from the left sinus Valsalva was divided into circumflex (Cx) and left anterior descending artery (LAD) branches. From the distal part of LAD and Cx, it was observed that the right coronary artery filling with retrograde grade 3 collaterals and opening to the PA.

showing a course in the right atrioventricular groove. Dilation was observed in LMCA and LAD (Figures 2 & 3). These findings were evaluated as compatible with ARCAPA.

DISCUSSION
Abnormal origin of the coronary arteries, typically from the contralateral side of the sinus Valsalva, is a relatively common finding during routine CCA and is associated with sudden cardiac death in athletes [10]. Coronary arteries originating anomalously from the PA, however, is a rare incidence. Compared to ARCAPA, patients with abnormal origin of the left coronary artery from the pulmonary artery (ALCAPA) are diagnosed early in life as they are usually symptomatic [1]. The diagnosis of ARCAPA, which is even less common than ALCAPA, can be delayed due to presence of collaterals, as was the case with our patient. ARCAPA can be seen alone or can be accompanied with other congenital cardiac defects such as tetralogy of Fallot, bicuspid aortic valve, aortic stenosis, septal defects or aortic coarctation [1,2]. We could not demonstrate any other cardiac defect in this specific patient.

As imaging modalities, transthoracic echocardiography (TTE), magnetic resonance angiography (MRA) and CCTA are used in the diagnosis of ARCAPA; but, MRA and CCTA are more reliable. TTE is a non-invasive method used to for evaluate cardiac defects. Although Doppler US may show the abnormally located ostium and proximal intramural course of coronary artery in children, this cannot be imaged in adults. Intracoronary collaterals within the ventricular septum, thought to be an indicative of ARCAPA, can,
Int J Anat Var 14 No 2 Feb 2021

Sasani H, et al.

Figure 2) Coronary CT angiography examination showed that the left coronary artery originated normally from the left (L) coronary sinus; RCA originated from PA, was at interarterial localization, then showing a course in the right atrioventricular groove. Dilation was observed in LMCA and LAD (RCA: right coronary artery, PA: Pulmonary artery, LCA: coronary artery from left pattern, LCx: circumflex artery, OM: obtuse marginalis branch).

Figure 3) The right coronary artery (RCA; yellow arrow) originates from MPA and shows an interarterial (AS and MPA) course. (AS: ascending aorta, MPA: main pulmonary artery).

however, be shown using color Doppler sonography [11]. In this patient, CCA could demonstrate the anatomy of right coronary circulation due to presence of excellent collaterals from left coronary system. This seems to delay the symptoms of the patient.

By using quantitative blood flow parameters and phase-contrast imaging, cardiac magnetic resonance imaging (CMR) can demonstrate pulmonary to systemic blood flow ratio, can be used in quantification of the flow difference in assessment of the shunt, in evaluation of right ventricular size and function and myocardial viability assessment [12]. Due to a high spatial resolution, CTA can provide 3D volume rendering reconstructions images for better clarifying the variation and also reveal the possible interarterial course of the anomalous coronary artery [13].

CONCLUSION

Although CCTA, CCA and MRA are among the most useful imaging methods in the diagnosis of ARCAPA; as in the current case, the origin and course of the anomalous coronary artery which cannot always be visualized in CCA, was shown on CCTA. Therefore, in order to confirm the diagnosis, the most appropriate imaging method can be selected individually based on the specific patient’s clinical findings.

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REFERENCES