Isolated aberrant right subclavian artery. A case report
Arteria subclavia derecha aberrante aislada. Reporte de un caso

Eli Pedro Monzón Castillo1,a, Gabriel Tejada Martínez1,b

ABSTRACT
An aberrant right subclavian artery (ARSA) is the most common branch abnormality of the aortic arch. It can be identified by ultrasound scan in 1% of cases. The probability of association with cardiac and/or extracardiac anomalies, as well as chromosomal abnormality, is high. Prevalence of ARSA with Down syndrome is approximately 20%, and this marker may contribute to counseling on Down syndrome during the second trimester and maybe in the first trimester. When ARSA is found, recommendations include a detailed study of the fetal anatomy for other markers of aneuploidy and to obtain a fetal echocardiogram. Invasive studies will be limited to those situations where, in addition to ARSA, other markers or other conditions that increase the risk of Down syndrome are found. However, the finding of an isolated ARSA increases the risk to zero and is considered a normal variant.

Key words: Aberrant right subclavian artery (ARSA), Down syndrome.

CASE REPORT

Introduction
The aberrant right subclavian artery (ARSA) is the most frequent abnormality of the aortic arch, present in approximately 1.2% of postnatal studies and in up to 2% in prenatal series. This anatomic variant has gained special importance because it is associated with congenital heart disease and as a probable marker for either Down syndrome(1), 22q11.2 deletion syndrome (Di George syndrome) and other genetic syndromes associated with abnormalities of the aortic arch(2,3).

In normal conditions, the aortic arch undergoes a complex development during early pregnancy that usually results in the formation of the left aortic arch, where three main branches originate: the first branch is the brachiocephalic trunk (BT) or innominate artery, which branch off into the right subclavian artery (RSA) and the right common carotid artery (RCCA); the second branch is the left common carotid artery; and the third is the left subclavian artery.

The development of the aortic arch is abnormal in approximately 1% to 2% of fetuses(2-4). Sometimes the disruption of the double aortic arch occurs between the RCCA and the RSA, impeding union of these two arteries and originating a left aortic arch with four branches instead of three. The last branch is now the RSA, which is born solitary instead of making part of the BT at the level of the descending aorta and close
to the anterior duct, although it has been described in other locations. This origin of the RSA is on the left and, therefore, goes a longer and straight way behind the trachea and the esophagus to reach its final target, the right arm\textsuperscript{(2,4)}. This abnormality is known as aberrant right subclavian artery or aberrant retroesophageal right subclavian artery or lusoria artery\textsuperscript{(4,5)}. In most cases, the arterial duct persists on the left side, originating an incomplete vascular ring\textsuperscript{(6)}. In 80% of cases, the ARSA goes behind the esophagus, in 15% between the esophagus and the trachea, and in 5% of cases in front of the trachea or the main bronchium\textsuperscript{(5)} (Figure 1).

A normal right subclavian artery is ‘S’-shaped, with anterior concavity in the proximal portion and posterior concavity in the distal portion. It passes in front of the trachea and esophagus and in direction of the right shoulder. If the right subclavian artery is not identified, it is recommended to descend into the axial plane of the three vessels and trachea to evaluate the possibility of an aberrant right subclavian artery. In case of doubt, Doppler color\textsuperscript{(4,7)} or pulse-Doppler would confirm the arterial or venous origin of the vessel in question, as the vein shows the typical vein wave\textsuperscript{(2,8-10)}.

It can also be observed in a longitudinal (sagittal) or coronal plane\textsuperscript{(8)}. The demonstration of the four vessels coming out of the aortic arch in the longitudinal plane is easier to obtain before the 22nd-week of pregnancy than in late pregnancies, as the vessels are closer to each other\textsuperscript{(8)}. Likewise, the coronal plane can be of additional help to find such vascular abnormalities\textsuperscript{(11)}.

By correctly using the assessment technique and knowing the possible diagnostic errors, success in identifying the normal and aberrant right subclavian artery has an effectiveness of 95% to 98% in the second trimester\textsuperscript{(1,2)} and 80% to 85% in the first trimester\textsuperscript{(2,8)}. These percentages are directly related to the operator experience and indirectly to the maternal body mass index (BMI). In the first trimester, visualization is also directly related to gestational age.

CASE REPORT

This 35-year old patient came to our hospital for prenatal control. Her weight before pregnancy was 70 kg, height 156 cm, BMI 28.8. She had regular menstrual cycles, date of her last menstruation on May 28, 2019. Current conception was spontaneous. She was a non-smoker and non-drinker.

First trimester analysis was reported normal, toxoplasmosis non-immune, VDRL negative, HIV and hepatitis C negative, hepatitis B immune, measles immune, blood group A Rh +.
Screening for Down syndrome in the first trimester showed low risk 1/7 732.

There was no relevant personal or family medical history for congenital heart disease or sudden death.

Obstetrical history was G3P2, two normal vaginal deliveries in 2015 and 2017 (newborns 3 500 g and 4 000 g, respectively); probable due date March 3, 2020. She presented transient gestational hyperthyroidism; the endocrinologist prescribed no treatment but control 3-4 months postpartum.

At 20 weeks of pregnancy, fetal anatomy ultrasound showed at the level of the three vessels and trachea, a retrotracheal vessel with suspicion of an aberrant right subclavian artery. No other cardiac or extracardiac structural alteration was identified. Sex was masculine. Figures 2 A and B.

She was referred to another hospital to confirm diagnosis, and fetal echocardiogram reported situs solitus, levocardia, levoapex, normal segmental arrangement, balanced ventricular volumes with normal transvalvular flows, integral interventricular septum, patent foramen oval, left aortic arch with aberrant right subclavian, left ductus arteriosus and incomplete vascular ring. There was a retrotracheal and retroesophageal vascular structure corresponding to aberrant right subclavian artery. No other extracardiac abnormalities were found. Male sex.

The parents were informed on the ultrasound findings. Association of ARSA with Down syndrome was explained, emphasizing on its low incidence when ARSA appears isolated, almost the same as for normal population. Likewise, they were told about symptoms and how to handle them in the newborn, indicating most patients are asymptomatic. In agreement with the parents, decision was not to conduct invasive tests and continue regular prenatal controls.

At 24 weeks, O’Sullivan test showed normal results. At 28 weeks, the mother received a whooping cough vaccine (protocolized in the Spanish National Health System).

At 33 weeks the fetus showed cephalic presentation by ultrasound, with normal growth, normal levels of amniotic fluid, regular fetal movements. Fetal echocardiogram showed the ARSA and no other cardiac or extracardiac structural pathology. Doppler of the umbilical artery revealed a normal pulsatility index. Figure 3.
At 36 weeks, rectovaginal culture was negative for Streptococcus agalactie. The patient was referred for fetal monitoring to start at 39 weeks of pregnancy.

The patient was admitted for finalization of pregnancy at 40 weeks and 6 days. Labor was induced using misofar 25 mcg. On March 10, 2020 at 13:50 hours, and under epidural analgesia, the patient presented normal vaginal delivery.

The newborn Apgar score was 9/10 and weight 3 470 g. He was evaluated by the pediatric cardiology service on March 11, 2020. Physical examination showed good general status, normal color, precordial palpation also normal. Femoral pulses were present and symmetric. Cardiac auscultation was rhythmic, with normal first heart sound, no murmur, and normal second heart sound, free diastole. Abdomen within normal limits. Echocardiograph showed normal anatomical relations. Normal lungs. Regular pulmonary and systemic venous drainage. Integral interatrial septum. Normal atrioventricular and sigma valves. Ventricles of regular size and functioning. Ejection fraction 73.5%. No obstruction of outflow tracts.

Normal pulmonary trunk and branches. Left aortic arch with exit of a wide vessel and bifurcation into two, which impressed as right and left carotids, followed by another vessel of normal diameter, corresponding to the left subclavian artery and, finally, another vessel that was consisting with the aberrant right subclavian, whose course could not be followed. No ductus was observed. Normal abdominal aorta pulsatility. Diagnosis: Aberrant right subclavian artery. Not associated to other cardiac malformations.

Appointment for control was in one month, but in view of the COVID-19 pandemic in Spain, control was by telephone at 19 days of age. The mother did not report noticeable alterations and was called for control by the pediatric service in July 2020, either personal or by telephone, depending on the state of emergency.

**Discussion**

In most reference centers, evaluation of the right subclavian artery is part of the routine echocardiogram study and is considered a marker of congenital heart disease and Down syndrome\(^1\). Finding of an aberrant right subclavian artery should be followed by comprehensive evaluation of fetal and cardiac morphology to rule out other abnormalities. Considering the prevalence on the general population, ARSA finding does not justify a routine invasive examination for Down syndrome\(^2,8\). This is backed up by the conclusion of a recent meta-analysis which confirmed that an isolated ARSA acts as a variant of normality\(^9\).

The most frequent abnormality of the aortic arch is ARSA, with incidence in the general population and postmortem studies estimated at approximately 0.4% to 2%. Rembouskos et al. prospective study, observed ARSA in 89 of 6 605 fetuses with normal karyotype. Gut et al. investigated the course of the right subclavian artery in 4 125 low-risk pregnancies between 17 and 33 weeks of gestation. ARSA was found in 17 of 4 120 patients (0.4%)\(^3,12\). Average prevalence of ARSA in the normal population is 1%\(^13\).

The presence of an aberrant right subclavian artery represents 20% higher risk of having another abnormality. Among the associated abnormalities, chromosomal alterations are the most frequent, mainly trisomy 21\(^2,3,7,9,10,13\).

Identification of ARSA in fetuses with Down syndrome was first described by Chaoui et al. in 2005\(^12,13\). Prenatal studies report the prevalence of ARSA in 6.8% to 37.5% of fetuses with Down syndrome in the second and third trimesters.
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According to recent publications, the risk for trisomy 21 increases primarily when the finding of ARSA is associated with other markers of trisomy 21. When ARSA presents as the only marker of this chromosomal abnormality, risk is not increased\(^\text{[1,6]}\). Therefore, isolated ARSA may serve as a weak marker for Down syndrome rather than a routine indication for invasive fetal testing\(^\text{[13,14]}\). In our case, we informed and explained the parents on ultrasound findings; agreement included non-invasive testing and routine pregnancy monitoring.

Although association between isolated ARSA and chromosome 22q11 microdeletion has not been proven, its detection in conotruncal heart malformations is the most important marker of this genetic alteration, so examination is recommended\(^\text{[2,8]}\). Offering an invasive procedure to exclude 22q11 microdeletion is under debate. Zidere et al. do not recommend it, unless an extracardiac malformation is present\(^\text{[13]}\).

Cardiac abnormalities are the most frequent structural abnormalities that coexist with ARSA (5.4% to 23.5%)\(^\text{[12]}\). Most frequent associated cardiopathy is tetralogy of Fallot, which was discussed by Blalock in 1948\(^\text{[3]}\). Correlation between ARSA and congenital heart disease was reported by Zapata et al, with a postnatal study that included 11 000 cases. They found ARSA incidence was 2.9% in patients with congenital heart disease and 0.1% in patients with normal heart, suggesting relationship between ARSA and congenital heart disease\(^\text{[12]}\). The ratio of ARSA to extracardiac abnormalities has been found to be approximately 5% to 26.7%\(^\text{[12]}\).

We conclude that it is possible to evaluate the right subclavian artery in the first and second trimesters of pregnancy, with chances of success of 80% and 90%, respectively. Options for achieving good visualization depend on the operator experience, gestational age and maternal body mass index.

Recommendation when an aberrant right subclavian artery is found is to perform a detailed study of the fetal anatomy in search of other aneuploidy markers and a fetal echocardiogram. Invasive studies will be limited to those situations where, in addition to the aberrant right subclavian artery, other markers or conditions that increase the risk of Down syndrome are found.

References
