Anomalous left coronary artery from the pulmonary artery (ALCAPA) in adult age. A 5-case series successfully repaired with surgery

Origen anómalo de la coronaria izquierda en la arteria pulmonar (ALCAPA) en la edad adulta. Una serie de 5 casos reparados con éxito con cirugía


**Key words:**
Congenital heart disease, pulmonary artery abnormalities, ALCAPA, myocardial ischemia, coronary reimplantation.

**Palabras clave:**
Cardiopatías congénitas, anomalías de la arteria pulmonar, ALCAPA, isquemia miocárdica, reimplantación coronaria.

**ABSTRACT**

**Objective:** To describe conduct, evolution and surgical treatment of ALCAPA in adult patients. **Material and methods:** Series of five cases, > eighteen years old. Clinical data (including surgery and follow-up), and echocardiographic/hemodynamic studies were reviewed. **Results:** The series comprised four women (80%) and a man (20%), with a mean age of 38.2 (range 18-65) years. Myocardial ischemia was found in three patients (60%), detection of a cardiac murmur in one patient (20%) and aborted sudden death in one patient (20%). Physical examination was normal in every case. Two patients (40%) presented grade II cardiomegaly and venocapillary pulmonary hypertension. In three cases, subendocardial ischemia was found on the EKG without necrosis evidence. Echocardiogram was performed in every patient. On the short axis window turbulent (mosaic) retrograde flow on left interventricular septum and left coronary artery absence was documented. Myocardial scintigraphy was performed on four patients finding moderate to severe anterolateral ischemia. CT angiography was performed on 80% of cases. In 40% of patients, left ventricular end diastolic pressure was elevated in cardiac catheterization. Surgical repair was successful in every case. Coronary reimplantation was performed on two patients, Takeuchi procedure was performed on two patients and revascularization with an internal mammary artery-to-anomalous left coronary artery from the pulmonary artery procedure in one patient. After 48 months of follow-up, every patient is alive in functional class I. **Conclusions:** Natural history of ALCAPA includes chronic ischemia, ventricular dysfunction and severe arrhythmias.

**RESUMEN**

**Objetivo:** Describir el comportamiento, evolución y tratamiento quirúrgico de ALCAPA en pacientes adultos. **Material y métodos:** Serie de cinco casos, mayores de 18 años de edad. Se revisaron los datos clínicos, ecocardiográficos, hemodinámicos, quirúrgicos y del seguimiento. **Resultados:** La serie comprendió cuatro mujeres (80%) y un varón (20%), con edad promedio de 38.2 años (rango 18-65 años). Se encontró isquemia miocárdica en 60% de los casos, detección de soplo en 20% y muerte súbita abortada en el 20%. La exploración física fue normal en todos los casos. Dos pacientes con cardiomegalia grado II e hipertensión venocapilar. En tres casos el ECG mostró isquemia subendocárdica pero sin evidencia de necrosis. El ecocardiograma reveló flujo turbulento retrógrado en septum interventricular y ausencia de la arteria coronaria izquierda en el eje corto. El gammagrama cardiaco fue positivo para isquemia, moderada a severa en región anterolateral. Se realizó angiotomografía en 80% de los casos. En 40% de los pacientes, la presión diastólica final del ventrículo izquierdo se encontró elevada en el cateterismo cardíaco. La cirugía fue exitosa en todos los casos. La técnica quirúrgica de elección fue el reimplante de la arteria coronaria en dos pacientes, procedimiento de Takeuchi (túnel intrapulmonar) en dos casos y en un solo caso con revascularización con arteria mamaria interna. Después de 48 meses de seguimiento todos los pacientes están vivos y en clase funcional I. **Conclusiones:** La historia natural de la ALCAPA implica isquemia crónica y disfunción ventricular además de arritmias severas que pueden llevar a la muerte. Son pocos los adultos con esta entidad.
and severe arrhythmias that could lead to death. There are very few cases in adults and the likelihood of reaching this age depends on collateral circulation provided by the right coronary artery. This was found in each case in this series, documented by echocardiography and cardiac catheterization. Surgical procedures for this entity include coronary reimplantation, Takeuchi procedure and internal mammary artery to anomalous left coronary artery from the pulmonary artery procedure (performed in one patient on this series). Prognosis is good, as long as the congenital defect is corrected.

INTRODUCTION

Anomalous left coronary artery from the pulmonary artery (ALCAPA) is an infrequent congenital defect, representing 0.25 to 0.5% of congenital heart defects, and affecting one in 300,000 live births. In pediatric population, it can cause cardiac dilatation or myocardial ischemia, whereas in neonatal population, it can be highly lethal, with mortality rates over 90% when it is complicated with myocardial infarction. In this scenario, only 10 to 15% of patients will reach adult age, depending largely on intercoronary collateral circulation. Furthermore, the disorder, besides of causing ischemia, can produce myocardial fibrosis, left ventricular remodeling and chamber dilatation, mitral valve insufficiency and congestive heart failure.

The clinical spectrum of this defect in adults presents from an asymptomatic course to a severe clinical presentation with malignant ventricular arrhythmias and myocardial ischemia. Thus, it is necessary to perform opportune surgical procedure with any of the techniques that establishes a two coronary arteries system: coronary reimplantation, creation of an intrapulmonary aortocoronary tunnel (Takeuchi procedure) or revascularization (either venous or arterial), along with a mitral valve repair if needed.

CASE DESCRIPTION

All cases of patients over the age of 18, diagnosis with ALCAPA in our center, in a 5-year period were included. Patients who did not accept surgical treatment or failed to complete a follow-up were excluded.
An early positive stress test was found on two patients, with subendocardial extensive anterior ischemia, that disappeared at rest (Figure 5).

Echocardiography was the key study to establish ALCAPA diagnosis in every patient. In three cases (60%) we found left ventricular dilatation with anteroseptal and anterior motility alterations. None of the patients suffered from mitral insufficiency. With color Doppler, turbulent (mosaic) retrograde flow was found on apical and four-chamber windows on every patient, ascending from apical portion to basal portion of the left ventricle. When continuous Doppler was used, low gradients and velocities were found (Figure 6). On the short axis window we documented turbulent (mosaic) retrograde flow on left interventricular septum and left coronary artery absence (Figure 7).

Myocardial scintigraphy was performed on four patients, finding moderate to severe anterolateral ischemia (Figure 8). CT angiography was performed on four patients, finding absence of left coronary artery, the origin of the left coronary artery from the pulmonary artery and contralateral coronary circulation (Figure 9).

All patients underwent bilateral selective coronary angiography and left heart catheterization, finding elevated left ventricular end diastolic pressure in two patients. Angiography demonstrated left coronary absence, right coronary circulation and opacification of both the left coronary system and pulmonary artery (Figure 10). In one case, it was possible to selectively cannulate the left coronary artery through the pulmonary artery.

The surgical technique was chosen based on anatomical findings. On three patients, coronary reimplantation was performed. Takeuchi procedure was performed in one patient and in other an internal mammary artery-to-anomalous left coronary artery from the pulmonary artery procedure was performed.

All patients are alive and are asymptomatic after 48-months of follow-up, in functional class I and without evidence of residual ischemia.

The patient who underwent Takeuchi procedure developed mild right ventricle outlet obstruction gradient. At this time, however, surgical procedure is not indicated.

REVIEW

Despite its rarity, this anomaly has been extensively studied, since Brook’s first description in 1886. In 1908, Abbot described the anomaly in detail, but it was until 1933 when Bland, White and Garland coined the current name: ALCAPA, the English acronym of Anomalous Left Coronary Artery from Pulmonary Artery.1

The incidence is low, affecting only 1 in 300,000 live births. Although it represents less than 0.5% of all congenital heart disease, it is the most common congenital defect of the coronary arteries.1,2

Even though ALCAPA usually occurs as an isolated defect, it is sometimes associated to other congenital heart disease (transposition of great vessels, pulmonary atresia with intact ventricular septum, tetralogy of Fallot, double right ventricle outlet and common arterial trunk).4

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Figure 2. Chest roentgenogram, case 1. 18-year old man with no cardiomegaly.

Figure 3. EKG, case 2. Anteroseptal necrosis and systolic overload data.
There are three embryologic theories trying to explain this anomaly: the first one proposes an abnormal disposition of the aortic-pulmonary septum, the second one set forth the persistence of the left coronary primordia (anlage) on the pulmonary artery which permits the anomalous development of the artery in the wrong artery, and the last one that put forward a conotruncus separation defect.¹

Four anatomic/hemodynamic phases are described:
1. Elevated pulmonary artery pressure with blood flow from the pulmonary artery to the left coronary artery without any sign of ischemia.
2. A decrease in both pulmonary artery pressure and pulmonary artery resistance, with flow from the right to the left coronary artery through collateral circulation. Therefore, hypoperfusion, myocardial ischemia and its clinical consequences like angina or infarction, when there is insufficient flow through collateral circulation.

Figure 4. EKG, case 1. Normal.

Figure 5. Stress test, case 1. Descent in ST segment in anterior and inferior locations.

Figure 6. Echocardiography, short axis at the great vessels, where absence of left coronary is seen.

Figure 7. Echocardiogram, short axis at the left ventricle. Mosaic color corresponds to coronary contralateral circulation.
3. A decrease in both the pulmonary artery pressure and pulmonary artery resistance, but with multiple intercoronary connections and an adequate left coronary artery filling pressure, with no evidence of ischemia.
4. Normal pulmonary artery pressure and pulmonary artery resistance, with left coronary artery flow depending on right coronary artery flow and its collaterals, which steal oxygenated blood from the right coronary artery and the pulmonary trunk causing left-to-right shunt. In this phase, myocardial ischemia may be subclinical.5

Unfortunately, a minority of patients will reach phases 2 or 3 because 85% of them die within the first year of age. The remaining 15% will reach phases 3 or 4 with a life expectancy of 30 to 40 years, dying of heart failure or arrhythmia (sudden death).3,4

The spectrum of the disease depends on collateral circulation and the development of ischemia, conditioning the clinical expression of the disease in two groups of ages; pediatric and adult ALCAPA.

The first group (pediatric ALCAPA) comprises newborns and infants that may have tachypnea, uncontrollable crying spells because of angina, failure to gain weight, irritability, heart failure, mitral insufficiency, cardiac arrhythmias, myocardial infarction and sudden death. Physical examination could reveal a holosystolic murmur due to mitral regurgitation. Approximately 15% of these patients are asymptomatic with a continuous murmur.

The second group (adult ALCAPA) may show no symptoms. This fact largely depends on the magnitude of collateral circulation from the right coronary artery. Some patients will suffer from silent or symptomatic ischemia, mitral insufficiency (dilatation of the mitral valve ring)

Figure 8. Cardiac scintigraphy, case 3. Severe ischemia induced by stress is shown at the anteroapical region.

Figure 9.
CT angiography, case 4.

Figure 10. CT angiography, case 5.
and sudden death which may be the first symptom of the disease (one patient in our series).3,5

EKG will show necrosis with deep Q waves on DI and aVL, left axial deviation and left ventricular hypertrophy.2,4,6

X-ray chest film may show cardiomegaly and left atrial/ventricular enlargement.

Echocardiography is a very important diagnostic tool, disclosing left ventricular function, and furthermore permitting ischemia evaluation (motility disorders and mitral insufficiency secondary to mitral valve ring dilatation). The coronary artery origin is evaluated on the short-axis window allowing to evince the absence of the left coronary artery. With Doppler color, a retrograde flow mosaic, ascending from the apex through interventricular septum (describing anterior descending coronary artery route filled by collateral circulation coming from the right coronary artery) could be shown.7

The previously discussed findings can be confirmed with CT angiography, allowing the location of the artery origin as well as the measure of distances among different structures, helping the surgeon to plan a specific type of repair.

Coronary angiography with aortography is the diagnostic gold standard in ALCAPA because both contrast studies unveil doubtlessly the left coronary artery absence, permitting selective both right coronary artery cannulation to evince collateral circulation and coronary anterior descending artery from the pulmonary artery.3,8

There is no doubt that surgery is the best treatment option for patients with ALCAPA and it can be an emergency intervention in a symptomatic infant. Several techniques have been described establishing a system of two coronary arteries; coronary artery ligation, revascularization, direct anastomosis (using internal mammary artery, saphenous or subclavian vein and prosthetic materials), coronary artery reimplantation or Takeuchi procedure. All of them with mortality rate below 15%. In our center, all our patients are alive after four years of successful surgical correction.9,10

Adverse prognostic factors include: age at onset, ventricular dysfunction and poor collateral circulation. Mitral annuloplasty must be considered in the presence of severe mitral insufficiency. Other treatment options may include left coronary artery occlusion.

As mentioned above, mortality in referral centers is as low as 15%; in our hospital every patient treated is alive after four years of successful surgical treatment.9,10

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