Percutaneous Closure of Complex Fistula Between the Internal Mammary Artery and a Lobar Branch of a Pulmonary Artery

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A thorough review of the literature identified only 20 reported cases of fistula involving the internal mammary (internal thoracic) artery and a lobar branch of a pulmonary artery. Surgical closure was frequently done to avoid complications associated with this anomaly. We report the first patient in whom percutaneous treatment was accomplished with a combined technique involving an Amplatzer Duct Occluder device and coils.

Key words: Internal mammary artery. Fistula. Coils.

INTRODUCTION

Fistulas between the internal mammary artery (IMA) and the pulmonary arteries are uncommon vascular abnormalities.1 Most individuals with such abnormalities are usually asymptomatic and the fistula is only detected in the physical examination because of a characteristic continuous murmur.2

For diagnosis and localization of the fistula, aortography and selective angiography of the IMA are used.1 In view of the possible of circulatory overload, degeneration of the arterial wall, infection, traumatic rupture or other complications,3 surgical closure of the defect, as described in the 20 cases reported in the literature, might be indicated.

We think our contribution is of interest because of the novelty of catheter closure of a mammary-pulmonary fistula with double arterial supply using a technique that combines use of coils and an Amplatzer Duct Occluder (ADO) device.

CASE STUDY

A 32-year-old man, who played sports regularly, presented with continuous murmur discovered by chance in his company's medical check-up. He had no cardiovascular risk factors, relevant clinical history, chest trauma or prior surgery. Chest x-ray showed no significant findings and, according to the electrocardiogram (ECG), he had sinus rhythm with a QRS axis of 60°, without increased cavity dimensions, or rhythm or conduction disorders. Transthoracic echocardiography revealed limited dilation of the right chambers without pulmonary hypertension, as the only significant finding. Thoracoabdominal computed tomography (CT) showed a possible vascular fistula between the right IMA and an aberrant artery that connected the abdominal aorta and the inferior lobar branch of the right pulmonary artery. Although the patient was asymptomatic, it was decided to close the defect because of the risk of chest trauma associated with play-
ing sports. The procedure was carried out by retro-
grade puncture of the right femoral artery. A large 5
mm aberrant artery next to the celiac trunk was selec-
tively probed. The aberrant artery flowed towards the
anterior phrenic and right medial region, to give flow
to a vascular tangle neoformation 4 cm in diameter. A
6-mm efferent vessel from this tangle drained the infe-
rior right lobar pulmonary artery. Next, the right mam-
mary artery was selectively probed. It was found to be
well developed, with a diameter of 6 mm. This artery,
before becoming the epigastric artery, gave rise to sev-
eral afferent branches to the vascular tangle found in
the phrenic region with drainage through to the ef-
ferent vessel mentioned above. To close the aortic-
mammary-pulmonary fistula, the aberrant aortic artery
was selectively probed, and 20 MWCE-135HILAL
(COOK) coils, 5 mm long by 4 mm in diameter, were
placed to achieve complete vessel occlusion (Figure
1). Next, a 6-Fr sheaf was placed in the mammary
artery and an ADO device of 6-8 mm and a Flipper
PDA5 coil of 8 mm diameter with 5 turns were im-
planted, and the vessel closed completely (Figure 2).

The difference in arterial oxygen content between the
right ventricle and the right pulmonary artery was
1.3% by volume, and the pressures in the smaller cir-
cuit were normal. Final aortography showed closure of
the double supply by occlusion of both vessels. Pro-
phyllactic antibiotics were administered for 7 days.
Chest contrast CT was performed at 2 months and at
one year post procedure. Both showed that the artery-
artery fistula had disappeared.

**DISCUSSION**

Mammary artery-to pulmonary artery fistula is a
rare malformation—we could only find 20 reports af-
after an exhaustive literature search. The first case of
mammary artery-to pulmonary artery fistula was
published by Burchell and Clagett,2 of the Mayo
Clinic, in 1947. These malformations can be classi-
ified into 3 broad groups according to their anatomical
connections:1 a) those derived from the proximal
part of the IMA that connect with the internal mam-
mary vein, the vena cava or the innominate vein; b)
those that connect the middle third of the IMA with
the pulmonary circulation, and c) those that connect
the distal third with the vitelline veins. Other shunts
between systemic arteries and pulmonary arteries
may originate from intercostal, bronchial, pericardial,
pericardiophrenic, lateral thoracic and esophageal
arteries, or from anomalous branches of the aorta4,5 (as
in our case).

These fistulas may be congenital or acquired from
inflammatory,6-7 neoplastic,8 or traumatic9,10 lesions.
Congenital fistula has been described in association with other anomalies such as the tetralogy of Fallot, with mitral valve prolapse or with sick sinus syndrome.

Several authors have suggested that congenital fistulas arise because pulmonary capillary vessels and the aorta, which connect systemic and pulmonary circulation in the fetus, fail to regress. Congenital fistulas may be apparent at birth but not become functionally active until triggered by external factors such as trauma, infection, inflammation, neoplasia or metabolic disorders.

These fistulas are usually diagnosed in young people in their teens or twenties. The mean age for diagnosis is 22 years. The youngest patient diagnosed was 7 years old and the oldest was 77 years old. In both these patients, the abnormality was diagnosed by abnormal pulmonary circulation in the chest x-ray. The ratio of men to women with such a fistula is approximately 2:1. Clinical presentation depends partly on the functional repercussion of these fistulas, which is proportional to the size of the implicated vessels and the fistula, and where it is located in relation to the heart. All 20 reported cases, and our patient, had a continuous precordial murmur. This poses the problem of differentiating between patients with such fistulas and those with patent ductus arteriosus, coronary artery fistula, rupture of Valsalva sinus, ventricular shunt with aortic valve regurgitation, coarctation of the aorta and arteriovenous fistulas. Chest x-ray, ECG and echocardiography all provide valuable information for differential diagnosis, but in this case, selective angiography is necessary to establish diagnosis of a fistula between the IMA and the pulmonary artery.

With time, artery-artery fistulas may cause vessel dilation and symptoms such as congestive heart failure and bacterial endocarditis. In view of the small number of cases, surgical indication for this abnormality is not clearly established. Some authors recommend that all patients should undergo surgery because of the risk of rupture, endarteritis and congestive heart failure due to progressive dilation of abnormal vessels. In contrast, other authors, who doubt the long-term effectiveness of surgery for an abnormality that may have a benign natural course, opt for conservative treatment of young patients. Most of the 20 cases reported in the literature underwent surgery with vessel ligation, alone or in association with excision of pulmonary tissue. No deaths were reported during surgery.

The Amplatzer Duct Occluder has been successfully used in complete closure of arteriovenous fistulas. This device was initially approved for closure of...
patent ductus arteriosus. The device is self-expandable, and made from nitinol-wire mesh. There is a retainer disk on one face, which is usually 2 mm larger than the diameter of the device. A Dacron mesh facilitates thrombosis and complete closure of the fistula. The main advantages of these devices are that they can be relocated, come in a wide range of sizes, can be placed by a small delivery catheter, have a high rate of complete closure, and are associated with zero mortality. Given the lack of similar reports in the medical literature, we showed the usefulness of this device in association with coils to close a complex artery-artery fistula.

REFERENCES