RUPTURE AND DISSECTION IN PULMONARY ARTERY ANEURYSMS: INCIDENCE, CAUSE, AND TREATMENT—REVIEW AND CASE REPORT

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Since the first report of dissecting aneurysm of the pulmonary artery by Walshe in 1862 (cited by Watson), 48 cases of pulmonary artery dissection have been reported. Dissection mostly develops in a pulmonary artery aneurysm associated with pulmonary hypertension and/or connective tissue disease. Forty-three cases were reported after death, whereas only 5 were diagnosed clinically, indicating that the event is highly lethal.

In this report, we describe the surgical repair of dissection of an aneurysm in the main pulmonary artery resulting from primary pulmonary hypertension.

Clinical summary. A 34-year-old woman was admitted to the emergency room with chest pain and shortness of breath. She had primary pulmonary hypertension for 10 years and had been receiving long-term therapy with coumarin and a continuous intravenous infusion of epoprostenol (prostacyclin) for 4 years, with the dosage increasing to 16 ng · kg⁻¹ · min⁻¹. Seven months previously, she underwent a right heart catheterization that revealed a pulmonary–systemic arterial pressure ratio (Pp/Ps) of 0.53. The electrocardiogram showed right ventricular hypertrophy. A posteroanterior chest x-ray film showed a left hilar mass indicating dilation of the left pulmonary artery. The patient was discharged without any further investigations because the physical examination disclosed no abnormalities. The next day, the patient was readmitted to the hospital because of worsening symptoms. The transthoracic echocardiogram revealed an enlarged main pulmonary artery with a diameter of 8 cm, a competent pulmonary valve, and massive pericardial effusion but failed to demonstrate pulmonary artery dissection. A computed tomogram confirmed dissection of the main pulmonary artery (Fig 1, A). She was transferred to the operating theater with signs of imminent cardiac tamponade and shock. Suprasystemic pulmonary pressures (systolic 120 mm Hg) were noted after induction of anesthesia with a Pp/Ps of 1.9.

Surgical technique. Femoro-femoral bypass was established urgently followed by sternotomy and pericardiotomy, which revealed massive bloody pericardial effusion. At a blood temperature of 32°C with the heart beating, the pulmonary...
The artery was longitudinally incised from the sinotubular junction into the left pulmonary artery up to the left pericardial reflection. The right pulmonary artery incision was made separately between the aorta and superior vena cava. The pulmonary artery dissection was found spreading from the sinotubular junction up to the pericardial reflection on the left and the intermediate trunk on the right. An intimal tear was located between the main pulmonary artery and the intermediate trunk of the right pulmonary artery. The pulmonary valve appeared normal and competent. The torn intima at the intermediate trunk was repaired with 4-0 Prolene running suture (Ethicon, Inc, Somerville, NJ). The beveled end of a 32-mm tube graft was anastomosed end to side to the hilar right pulmonary artery with 5-0 Prolene running suture so that a T-shaped replacement of the central pulmonary artery could be done by the inclusion technique (Fig 1, B). The graft was then pulled through within the central right pulmonary artery to the left side. The free end of the graft was anastomosed in the same way to the hilar left pulmonary artery. Another 32-mm tube graft was interposed between the graft and sinotubular junction with 5-0 Prolene running suture. Finally, the pulmonary artery incision was closed with 4-0 Prolene running suture, burying the T-shaped graft within the pulmonary artery. After cardiopulmonary bypass was discontinued, the patient again had suprasystolic pulmonary pressure. Continuous infusion of alprostadil (prostaglandin E1) at 0.2 µg·kg⁻¹·min⁻¹ and nitric oxide inhalation with 8 ppm was administered before the patient was transferred to the intensive care unit.
On the first postoperative day, alprostadil infusion was changed to epoprostenol (prostaglandin I$_2$) at dosages of 16 to 12 µg · kg$^{-1}$ · min$^{-1}$ and milrinone of 0.4 to 0.2 µg · kg$^{-1}$ · min$^{-1}$. Hemofiltration was performed from postoperative day 1 for 48 hours because of oliguric renal insufficiency. Severe pulmonary hypertension improved gradually with decrement of Pp/Ps from 1.6 to 0.7 (Fig 2).

The fluid requirements decreased dramatically on the third postoperative day with recovery of renal function and decrease in blood lactate concentration. The following course was uneventful. The patient was weaned from nitric oxide inhalation on the sixth postoperative day and was extubated on the eleventh postoperative day. On the thirteenth postoperative day, the patient was transferred to the general ward with still elevated but subsystemic pulmonary pressures (Pp/Ps = 0.6) receiving epoprostenol, 16 ng · kg$^{-1}$ · min$^{-1}$.

The patient was discharged on postoperative day 45.

A postoperative computed tomogram showed a sealed leak around the distal tube graft covered with the native pulmonary artery. Pathologic examination of the pulmonary arterial wall showed myxoid degeneration of the endothelial surface and confirmed the presence of dissection within the aneurysmal wall. The histologic examination showed that the specimen did not contain collagen type I or type III.

Because of persistent pulmonary hypertension, the patient was accepted for lung transplantation. She underwent double lung transplantation with the use of an extracorporeal membrane oxygenator 9 months after the emergency operation. Intraoperatively, the sealed leakage was found to be thrombosed. The pulmonary prosthesis was replaced, with a short part of the prosthesis being left in place connecting the main pulmonary artery with the transplants. The patient was discharged 3 weeks after the operation in good condition and remained well 10 months after the transplantation.

**Discussion.** Dissection of the pulmonary artery is a rare but life-threatening event. Although the pathophysiologic cause of dissection in pulmonary artery aneurysm is not clear, pulmonary hypertension and resultant mucoid degeneration of the media and fragmentation of the elastic fibers strongly predispose to this condition. In the literature, dissection of the pulmonary artery always occurs at the site of a pulmonary aneurysm or dilation. It is reasonable to assume that pulmonary artery dissection occurs at the point where pulmonary artery tissue becomes too fragile to support the tension of the pulmonary artery wall. Tissue fragility is caused by infectious and connective tissue diseases, and high wall tension is caused by high internal pressure and a large-radius aneurysm, as predicted by LaPlace’s law. In pulmonary artery dissection, the false lumen tends to rupture rather than to develop a re-entry site, as is usual in aortic dissection. Six cases of pulmonary artery dissection associated with primary pulmonary hypertension have previously been reported.

Yu and colleagues$^5$ showed that epoprostenol exerts inhibitory actions on proliferation and DNA synthesis of rat cardiac fibroblasts and messenger RNA expression for collagen types I and III. In our patient, long-term continuous epoprostenol infusion in the presence of severe pulmonary hypertension might have predisposed to aneurysmal change and subsequent biochemical changes within the vessel wall of the pulmonary artery leading to dissection.

The symptoms of pulmonary artery dissection are nonspecific, with 82% of patients having exertional dyspnea, 67% retrosternal chest pain, and 52% central cyanosis. The diagnosis was made by echocardiography, computed tomography, and/or magnetic resonance imaging. Patients with pulmonary hypertension and those receiving long-term epoprostenol infusion having these symptoms should be examined urgently with echocardiography and computed tomography to exclude pulmonary artery dissection. In our patient, the computed tomogram should have been obtained on the first admission because of her primary pulmonary hypertension.

Because pulmonary artery dissection is highly life-threatening, we believe patients with pulmonary artery dissection should undergo emergency surgery to prevent lethal bleeding. However, the literature contains only one report of a patient with dissection of the pulmonary artery undergoing surgery. That patient had chronic dissection of the pulmonary artery without pulmonary hypertension. To our knowledge, we are reporting on the first successful treatment of a patient who underwent surgical repair of acute pulmonary artery dissection.

Histologic examination was done by Prof Dr M. Susani from the Pathology Department of the University of Vienna.

Received for publication Aug 24, 2000; accepted for publication Oct 24, 2000.

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6.2. General Medical Treatment and Risk Factor Management for Patients With Thoracic Aortic Disease.

9.2. General Medical Treatment and Risk Factor Management for Patients With Thoracic Aortic Disease.

9.2.1. Recommendation for Medical Treatment of Patients With Thoracic Aortic Diseases.

9.2.1.1. Compiled reports were reviewed and additional articles were provided by committee members. We report a case of idiopathic dissection of the MPA in a patient without pulmonary hypertension, cardiac disease or cardiac intervention who presented with acute chest pain. Among the 63 previously reported cases of PA dissection, 34 patients had underlying cardiac disease most commonly congenital heart defects including patent ductus arteriosus and seven patients had rheumatic mitral stenoses. Nine patients with idiopathic PA hypertension with dissection have been reported. Senbaklavaci O, Kaneko Y, Bartunek A: Rupture and dissection in pulmonary artery aneurysms: incidence, cause and treatment - review and case report. J Thorac Cardiovasc Surg. 2001, 121: 1006-1008. Rupture and dissection in pulmonary artery aneurysms: incidence, cause and treatment - review and case report. J Thorac Cardiovasc Surg. 2001; 121: 1006-1008.