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Nominal Hemoptysis Heralds Pseudoaneurysm Induced by a Pulmonary Artery Catheter

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Massive hemoptysis secondary to pulmonary artery rupture is a rare but often fatal complication of pulmonary artery catheters (PACs).1 Several cases in the literature warn of the danger in ignoring nominal hemoptysis as an early sign of pulmonary artery rupture by a PAC.2,3 In all of these cases, the hemoptysis was accompanied by a new infiltrate or nodule on chest x-ray (CXR). We report a case of delayed fatal hemoptysis secondary to rupture of a PAC-induced pseudoaneurysm, which was heralded by a small amount of hemoptysis in association with a normal postinsertion CXR.

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CASE REPORT

An 82-yr-old male was transferred to our hospital for coronary artery bypass graft surgery (CABG). At the time of cardiac catheterization (1 day prior to surgery), a 7-Fr thermoludition PAC was inserted without incident via the right femoral vein. Pulmonary artery pressures were found to be normal. The patient received heparin and nitro-
glycerin infusions and was transferred to our hospital for urgent CABG surgery. The heparin was discontinued 4 h prior to surgery.

Upon the patient’s arrival in the operating room, a radial artery catheter and a right internal jugular introducer were inserted. The baseline activated clotting time (ACT), performed prior to insertion of the central venous introducer, was 115 s. The CXR on admission to our institution revealed the femoral PAC to be in the left main pulmonary artery. In order to have the ability intraoperatively to mani-
upulate the PAC, it was elected to insert a PAC from the right internal jugular site. A 7.5-Fr oximetric PAC (Opticath®, Abbott Critical Care
System) was floated through the right internal jugular introducer to a
wedged position and secured to the skin, at the 47-cm mark, after
deflation of the balloon. Initial pulmonary artery pressures were 21/
10 mmHg. At no time was the balloon on the PAC reinflated after
initial placement. Anesthesia was induced with fentanyl, 50 μg/kg,
and vecuronium 0.15 mg/kg was used for neuromuscular blockade.

After induction, at the surgeons’ request, a rolled sheet was placed
transversely under the patients shoulders. Immediately after the patient was repositioned, the PAC waveform was interpreted by the attending
anesthesiologist to reflect a wedged position, and the PAC promptly
was withdrawn 5 cm until a characteristic pulmonary artery trace was
obtained. Five minutes after surgical positioning of the patient, blood
was noted in the endotracheal tube (ETT), and a volume of less than
10 ml was suctioned from the trachea. Observation of the patient for
15–20 min did not reveal additional blood in the airway and a decision
was made to proceed with surgery due to the critical nature of the coronary disease. The coronary revascularization proceeded uneventfully, and no additional blood was noted from the patient's ETT at any time during the case.

The patient separated from CPB without difficulty and was transported to the surgical intensive care unit (SICU) with his trachea intubated and lungs manually ventilated. The immediate postoperative CXR (fig. 1) obtained in the SICU showed the femoral PAC to be in the left main pulmonary artery; the PAC inserted from the right internal jugular site was in the pulmonary outflow tract. Pulmonary artery pressures were normal in the postoperative period. There was no infiltrate or other parenchymal abnormality noted in either lung field.

The balloon on the PAC was not inflated at any time during the postoperative period, and both PACs were removed on the morning of postoperative day 2. The patient was weaned easily from the ventilator, and his trachea extubated 12 h after admission to the SICU. The routine CXR obtained on the morning of postoperative day 2 also was without abnormality. During the early evening of postoperative

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**Fig. 1.** Postoperative portable chest x-ray.

**Fig. 2.** Arrowhead indicates a ruptured pulmonary artery pseudoaneurysm in communication with a subsegmental bronchus of the lower lobe of the right lung.
day 2, while he was sitting in a chair in the SICU, the patient was observed to have a sudden onset of massive hemoptysis followed by bradycardia and hypotension. Despite immediate resuscitative efforts and intubation with a double-lumen ET T the patient expired.

On postmortem examination a ruptured pseudoaneurysm of the right lower lobe pulmonary artery was found. (fig. 2)

**DISCUSSION**

Rupture of the pulmonary artery is a recognized complication of pulmonary artery catheterization. There are several cases in the literature describing the occurrence of pulmonary artery perforation by a PAC, and the frequency of this rare but often fatal complication is estimated to be 0.06–0.2% of cases. Mortality is reported at 45–64%, and morbidity in the absence of mortality is significant.1–6

Certain factors have been cited as predisposing a patient to pulmonary artery injury from PACs.7 These include age greater than 60 yr and pulmonary hypertension, both of which are associated with degenerative changes in the vessel wall causing its increased fragility. The presence of systemic anticoagulation has been mentioned,7 but it is likely that this predisposes to more severe manifestations of injury rather than to pulmonary artery injury itself. Distal location of the PAC tip and excessive manipulation and migration of the PAC also have been implicated as causes of pulmonary artery rupture.8 The actual mechanism of injury is multifactorial and may include eccentric inflation of the balloon, driving the PAC tip through the arterial wall; inflation of the balloon in the presence of pulmonary hypertension; and over-distention of the balloon, causing pulmonary artery rupture. The actual pressure exerted on pulmonary arterial walls has been investigated by Hardy et al., who found that balloon-generated intraluminal pressures as low as 975 mmHg may cause pulmonary artery rupture.7 Techniques and equipment for limiting generation of dangerous pressures during inflation of PAC balloons have been described.9 During cardiac surgery—the most common clinical situation associated with pulmonary artery perforation—injury may occur during manipulation of the heart, while the heart is empty and the PAC is cold and less pliable, allowing for easy distal migration.8

PAC-induced pseudoaneurysm formation also is uncommon but differs from arterial rupture in pathology and clinical course. A pseudoaneurysm, or false aneurysm, is a dilatation of a vessel or vascular space in which disruption of the normal endothelial lining has occurred.10 The wall of the aneurysm is not composed of elements of the blood vessel wall. Containment of blood within the vascular space is achieved by compression from tissues or hematoma or both, surrounding the perforated vessel.11,12 A layer of fibrin may be present and aid in containment of the hemorrhage. These PAC-induced pseudoaneu-

**REFERENCES**


Successful Anesthetic Management of a Patient with Hypokalemic Familial Periodic Paralysis Undergoing Cardiac Surgery

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Hypokalemic familial periodic paralysis is a rare disease, with obvious anesthetic implications. There are few reports commenting on the anesthetic management for patients with this disease when undergoing general surgery, and we have found only one report of such a patient subjected to cardiac surgery with extracorporeal circulation. This patient's course was complicated early in the postoperative period by an episode of paralysis, which necessitated prolonged mechanical ventilation.

We report a case of a patient with known hypokalemic familial periodic paralysis, who required coronary revascularization. An impending episode of paralysis on the 1st postoperative day was aborted by aggressive management of plasma potassium concentration.

CASE REPORT

A 58-year-old man was admitted with a history of progressive chest pain on exertion that was unresponsive to medical treatment with calcium channel blockers. Beta-adrenergic blockers had been avoided, since they may precipitate symptoms in patients with some types of familial periodic paralysis. Selective coronary angiogram showed critical stenoses of two major vessels with preserved ventricular function. The patient stated that from infancy he had suffered from acute episodes of intense muscular weakness, either associated with stress or infectious diseases or of no apparent cause. Those episodes frequently were of such severity that he was completely unable to move or even to call for help. He knew that several of his relatives, including his father, brother, and cousin, had experienced similar episodes. Diagnosis of hypokalemic familial periodic paralysis had been made several years ago based on a clear association with decreased plasma potassium concentrations: therefore, therapy consisting of potassium supplements and oral acetazolamide was instituted, and for the last 5 yr he had not experienced symptoms of the disease.

His physical examination was normal; he demonstrated adequate muscular tone and strength. Laboratory tests and radiologic examinations were normal. Plasma potassium was 4.2 meq/l.

The patient received 1 mg flunitrazepam the night before surgery and meperidine 1 mg/kg intramuscularly (im) before transfer to the operating room. After percutaneous insertion of peripheral venous, radial arterial, and internal jugular catheters, anesthesia was induced with 20 µg/kg fentanyl and 0.1 mg/kg pancuronium. Halothane 0.5–1% in oxygen was added as required for maintenance of anesthesia. No additional muscle relaxants were used. Monitoring included modified V5 ECG lead, continuous arterial and central venous pressures waveforms, urine output, and temperature. Blood gases and plasma potassium concentration were frequently measured throughout the operation and postoperative period and maintained within normal range. Intravenous glucose was avoided, since it is known to precipitate paralysis in these patients.

Cardiopulmonary bypass was carried out under normothermia, since it has been reported that hypothermia may precipitate paralysis.

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