Bronchial artery aneurysm (BAA) is a rare vascular abnormality that may result in life-threatening bleeding if it is left untreated. We present the case of a 35-year-old man with a mediastinal BAA characterized by a short inflow artery segment and tortuous single outflow vessel. The patient's BAA was treated with a novel approach involving placement of a patent ductus arteriosus closure device in the short inflow segment as well as coil embolization of the outflow vessel, successfully excluding the BAA. Two-week follow-up revealed no flow in the embolized artery on computed tomography angiography. This case demonstrates the first successful use of a patent ductus arteriosus occluder device in the treatment of a mediastinal BAA with short inflow segment. (J Vasc Surg Cases and Innovative Techniques 2020;6:93-5.)

**Keywords:** Bronchial artery aneurysm; Patent ductus arteriosus occluder; Endovascular; Embolization

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**CASE REPORT**

A 35-year-old man was referred to Toronto General Hospital for treatment of BAA detected incidentally on CT during follow-up of cystic fibrosis. The aneurysm measured 20 x 25 mm in axial dimensions.

The patient's cystic fibrosis had previously been complicated by *Pseudomonas* infection, *Mycobacterium* abscesses, and pneumothorax. He denied any cardiovascular symptoms, including chest pain and palpitations. His breathing was normal, with limited phlegm production and no hemoptysis. The BAA was technically challenging to treat because of a short inflow segment that spanned a length of approximately 1 mm beyond the origin of the left bronchial artery, arising from the proximal descending aorta (Fig 1). The single outflow vessel was also noted to be significantly tortuous.

Access was gained through the right common femoral artery, and a 100-cm 7F sheath was introduced (Terumo, Shibuya, Japan). A 7F Vista Brite Tip guiding catheter ( Cordis, Miami, Fl) was used to cannulate the origin of the left bronchial artery. Angiography showed brisk flow through the aneurysm sac located just beyond the origin of the left bronchial artery, with a short inflow segment and a dominant, tortuous single outflow vessel noted. There was no evidence of a bronchial artery to pulmonary artery fistula.

Initially, there was difficulty in cannulating the dominant outflow vessel as we were unable to determine the origin of the outflow ostium. The decision was made to use saline injection to propel 3-mm Tornado microcoils (Cook Medical, Bloomington, Ind), some of which lodged in the outflow tract and a few of which remained within the aneurysm sac. This served to provide an estimation of the outflow artery ostia and to approximate the distance the coils would travel if deployment was performed within the aneurysm sac. The outflow vessel was subsequently cannulated with moderate difficulty using a Transcend wire and a 2.4F
Progreat microcatheter (Terumo) and embolized with a combination of 3- to 5-mm Tornado microcoils with good effect.

The wire was exchanged for a 1-cm floppy tip Amplatz wire (Boston Scientific, Marlborough, Mass) and the sheath upsized to 8F. An 8F Mach 1 guiding catheter (Boston Scientific) was inserted with the tip parked in the BAA. An 8-mm Amplatzer PDA occluder (product reference 9-PDA-005; St. Jude Medical, St. Paul, Minn) was placed across the neck and deployed in a satisfactory position (Fig 2, a and b).

Completion angiography demonstrated no filling of the BAA (Fig 2, c). Hemostasis was achieved with an 8F Angio-Seal vascular closure device (Terumo). Because of the patient's thin body habitus, a salinoma was created anterior to the right groin puncture site to accommodate the collagen plug of the Angio-Seal device.

No postprocedural complications were encountered, and the patient was discharged on the same day. Two weeks after BAA embolization, the patient denied any symptoms, and CT angiography evaluation revealed no opacification of the aneurysm, indicating successful treatment (Fig 3). A repeated CT study has been arranged for 6 months after intervention to assess for aneurysmal sac regression.

**DISCUSSION**

BAA is an uncommon vascular phenomenon that often is manifested asymptomatically but can result in massive hemoptysis or mediastinal hemorrhage in the case of rupture. Given the multitude of underlying causes, including bronchiectasis, chronic bronchopulmonary inflammation, trauma, and infection, BAA may be detected incidentally on routine cross-sectional investigation of the lungs. Because of the potential risk of life-threatening hemorrhage, definitive surgical or endovascular treatment is indicated regardless of size.

Traditionally, treatment involved surgical resection or ligation from the midline or lateral thoracotomy. The endovascular approach has become favored, given the
reduced postoperative morbidity and faster recovery time. New technologies are also pushing the boundaries of what can be achieved by minimally invasive methods. Studies have demonstrated that both surgical and endovascular options result in similar outcomes.4

Narrow-necked proximal BAAs, such as that exemplified in this case, are particularly challenging to treat. The use of conventional coils to embolize the inflow segment was deemed too risky, given the high risk of coil migration and subsequent nontarget embolization. Even the controlled use of detachable coils would have been impossible, given the approximately 1-mm length of the inflow segment. Several previous reports have described the use of thoracic stent grafts to exclude proximal mediastinal BAAs that originate close to the thoracic aorta.5,6 In a case described by Hu et al,7 the sac of the proximal BAA was initially embolized with gelatin sponge before the deployment of a thoracic stent graft, resulting in occlusion of both the inflow and outflow of the aneurysm. Although thoracic stent grafts are useful in emergent situations, for example, in cases of acute rupture with active bleeding, their use can lead to complications such as endoleak.

Jagia et al8 and Chadha et al9 have both described cases of short-necked pulmonary artery aneurysms successfully treated with PDA occlusion devices. With regard to the use of closure devices for PDAs, complications have been described in large studies with long-term follow-up; these studies have demonstrated that complications are exceedingly rare.10-12 Hemolytic anemia is the most commonly reported complication, occurring in approximately 1% of cases. There have been no reports of distal emboli in these studies.

CONCLUSIONS

This case report exemplifies the successful use of a PDA closure device for the treatment of ostial BAA, demonstrating it to be a viable treatment option in patients who would otherwise not be candidates for endovascular embolization because of adverse technical factors.

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