**CASE REPORT**

**Serratia liquefaciens** Infection of a Previously Excluded Popliteal Artery Aneurysm

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**Introduction:** Popliteal artery aneurysms (PAAs) are rare in the general population, but they account for nearly 70% of peripheral arterial aneurysms. There are several possible surgical approaches including exclusion of the aneurysm and bypass grafting, or endoaneurysmorrhaphy and interposition of a prosthetic conduit. The outcomes following the first approach are favorable, but persistent blood flow in the aneurysm sac has been documented in up to one third of patients in the early post-operative setting. Complications from incompletely excluded aneurysms include aneurysm enlargement, local compression symptoms, and sac rupture. Notably infection of a previously excluded and bypassed PAA is rare. This is the third reported case of PAA infection after exclusion and bypass grafting and the first due to **Serratia liquefaciens**.

**Methods:** Relevant medical data were collected from the hospital database.

**Results:** This case report describes a 54 year old male patient, diagnosed with acute limb ischaemia due to a thrombosed PAA, submitted to emergency surgery with exclusion and venous bypass. A below the knee amputation was necessary 3 months later. Patient follow-up was lost until 7 years following surgical repair, when he was diagnosed with aneurysm sac infection with skin fistulisation. He had recently been diagnosed with alcoholic hepatic cirrhosis Child—Pugh Class B. The patient was successfully treated by aneurysm resection, soft tissue debridement and systemic antibiotics.

**Conclusion:** PAA infection is a rare complication after exclusion and bypass procedures but should be considered in any patient with evidence of local or systemic infection. When a PAA infection is diagnosed, aneurysmectomy, local debridement, and intravenous antibiotic therapy are recommended. The “gold standard” method of PAA repair remains controversial. PAA excision or endoaneurysmorrhaphy avoids complications from incompletely excluded aneurysms, but is associated with a high risk of neurological damage.

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**INTRODUCTION**

Popliteal artery aneurysms (PAAs) are rare in the general population but can cause significant morbidity and mortality, being the commonest cause of non-traumatic leg amputation. Despite their rarity, PAA are the most common peripheral arterial aneurysms and many of those affected have concomitant abdominal aortic aneurysms (33%) and contralateral PAA disease (50%).

There are several possible surgical approaches including exclusion of the aneurysm and bypass grafting or endoaneurysmorrhaphy and interposition of a prosthetic conduit, usually through a posterior approach. However, the “gold standard” method of repair remains controversial, as both methods have unique merits and risks.

The most common complication after elective PAA surgical repair is late bypass failure, independent of the approach. Outcomes following PAA exclusion are favorable, but persistent blood flow in the aneurysm sac has been documented in up to one third of patients in the early post-operative setting. Complications from incompletely excluded aneurysms include aneurysm enlargement, local compressive symptoms, and sac rupture. Notably infection of a previously excluded and bypassed PAA is rare.

This paper describes one case of **Serratia liquefaciens** infection of a previously excluded popliteal aneurysm 7 years after initial surgical repair. This is the third reported case of PAA infection after exclusion and bypass grafting and the first reported case due to **S. liquefaciens**.

**CASE REPORT**

The patient was a 54 year old male with a previous history of smoking and heavy drinking (mean 70 grams of alcohol...
per day). He was chronically medicated with acetylsalicilic acid and simvastatin.

In 2009 he was referred to the emergency department with acute limb ischaemia. A thrombosed popliteal artery aneurysm was discovered during the diagnostic workup. The runoff was poor, and no distal tibial vessel was identified. The patient was submitted to emergency surgery with aneurysm exclusion and bypass with a venous conduit (great saphenous vein). Owing to an adverse clinical outcome, below the knee amputation was necessary 3 months later. A contralateral PAA was diagnosed, but abdominal aortic aneurysm was excluded.

The patient was lost to follow-up until 7 years later, when he was admitted to the emergency department with a 2 day history of pain and inflammatory signs over the supra-articular incision for the PAA repair. There was no history of local trauma. When specifically asked, he denied pulmonary, urinary or any other symptoms. He had been diagnosed with alcoholic hepatic cirrhosis Child–Pugh B during the lost follow-up period.

Physical examination included hemodynamic stability, apyrexia, and exuberant inflammatory signs over the supra-articular incision with hematoma mixed with purulent drainage. No signs of cirrhosis decompensation were evident on examination.

Investigations revealed no leukocytosis \(8.19 \times 10^3/\mu\text{L}\), and marginal elevation in ultra-sensitive C-reactive protein \(5.30 \text{mg/dL}\). Liver workup revealed no cirrhosis decompensation.

Given the patient’s clinical findings, computed tomography angiography (CTA) was performed, which revealed a liquid collection \(79 \times 62 \text{mm}\) with a thin wall in the postero-medial left thigh. There were no clear limits between the vascular structures and the inflammatory collection, so communication with the aneurysmal sac could not be excluded. Persistent aneurysm perfusion and rupture signs were not identified, but could not be excluded (Fig. 1).

The patient was submitted to urgent surgery, with a medial approach over the previous incision. Intra-operative findings included abundant liquefied hematoma mixed with

![Figure 1.](image-url)
purulent drainage, identification of the open aneurysmal sac, and no areas of active bleeding into the sac. The surgical procedure included hematoma and purulent content drainage, debridement of the peri-aneurysmal soft tissue, and aneurysm sac excision. The popliteal artery inflow and outflow were oversewn. The operative field was irrigated and the wound was closed with a negative pressure wound system.

The patient was treated empirically with vancomycin and piperacillin/tazobactam. Blood cultures were negative, but microbiological analysis of the purulent content revealed *S. liquefaciens* infection. The antibiogram revealed resistance to amoxicillin, amoxicillin with clavulanic acid, and cefuroxime, but sensitivity to multiple other antibiotics; hence, he started specific antibiotic treatment with ceftazidime.

Clinical recovery during hospital admission was favorable. He returned to the operating room 3 and 8 days after the first procedure for local debridement and exchange of the negative pressure wound system. Sixteen days after the index procedure, the surgical wound was closed, and the patient was discharged 2 days later with no further complications.

**DISCUSSION**

The “gold standard” method of PAA repair remains controversial, but the most commonly performed surgical procedure for PAA is exclusion and bypass, which involves proximal and distal ligation of the aneurysm sac in combination with a reversed saphenous vein bypass graft. A number of recent reports have highlighted the risk of persistent blood flow in the “excluded” aneurysm sac following exclusion and bypass. More recently, endovascular repair of a PAA has proven to be a viable treatment option. This option exists even for rupture of an excluded PAA.

The largest review of follow-up of PAA repair published to date included 717 legs and revealed a 33% risk of aneurysm sac expansion in the medial approach group and 8.3% in the posterior approach group. About 88% of patients with aneurysm sac expansion were asymptomatic, and 5% underwent surgical re-intervention.

Results from other series in the literature were similar, raising long-term follow-up imaging controversy and routine surveillance of the excluded aneurysm sac recommendations were issued.

Primary mycotic popliteal aneurysms are rare, with an estimated incidence of less than 2% of all popliteal aneurysms. The pathogens most commonly implicated in the infection of popliteal aneurysms are Gram positive cocci such as *Staphylococcus* and *Streptococcus*. Gram-negative bacilli such as *Proteus*, *Escherichia coli*, *Salmonella*, and *Campylobacter* species have occasionally been reported. The source and mechanism of spread is presumably haematogenous.

There are only two case reports in the literature of PAA infection after surgical exclusion. In both procedures the great saphenous vein was used as a conduit, and infectious complications occurred more than 10 years after PAA repair. In both cases clinical presentation included fever, pain, and swelling in the popliteal fossa. The bacteria were *Staphylococcus aureus* and *Campylobacter fetus*.

The *Serratia* genus, as a whole, is responsible for about 2% of nosocomial infections, including urinary tract infections, bloodstream infections, sepsis, pneumonia, meningoccephalitis, and other debilitating infections. However, it is not reported as an agent responsible for mycotic aneurysms. In this specific case, the patient presented with hepatic cirrhosis Child–Pugh class B. Cirrhosis is considered an immunocompromised state; therefore, unusual bacteria are frequently observed and more virulent in patients with cirrhosis relative to those without liver disease.

As a conclusion, PAA infection rarely occurs after an exclusion and bypass procedure but should be considered in any patient with evidence of local or systemic infection. When a PAA infection is diagnosed, aneurysm excision, debridement, and intravenous antibiotic therapy are recommended. Reconstruction with autogenous conduit should be considered if the bypass remained patent.

**CONFLICT OF INTEREST**

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