A surprising diagnosis in a young patient with intermittent claudication: Symptomatic isolated external iliac artery aneurysm associated with cystic media necrosis

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ABSTRACT
Aneurysms of the external iliac artery are extremely rare. We present a case of a middle-aged male patient with calf claudication owing to peripheral arterial embolism on the basis of a thrombosed true aneurysm of the external iliac artery caused by cystic media necrosis. Vascular imaging established the diagnosis and we proceeded to removal of the aneurysm via open repair, with excellent surgical and clinical results. (J Vasc Surg Cases and Innovative Techniques 2020;6:352–6.)

Keywords: Aneurysm; Isolated iliac artery aneurysm; External iliac artery; Intermittent claudication; Cystic medial necrosis

Isolated iliac artery aneurysms (IAA) are reported in up to 7% of all aortoiliac aneurysms and 12% to 48% of all isolated iliac aneurysms (IAA) are bilateral.1-3 True aneurysms of the external iliac artery (EIA) are especially rare.4-6 In the present case, we discuss how we diagnosed and successfully treated a male patient with a symptomatic isolated external IAA caused by cystic medial necrosis (CMN) and review the presentation of this rare entity in the literature.

Written informed consent was obtained from the patient for publication of this case report and accompanying images.

CASE REPORT
A 51-year-old male patient was referred to our department of vascular surgery with a 5-month history of persistent calf and foot claudication of his right lower extremity. No rest pain was reported. The patient was an active smoker (25 pack-years), with no other known cardiovascular risk factors and he was not under any medication. Fourteen years before the acute incident, the patient had suffered a left sided pyelonephritis without sepsis. On physical examination the peripheral pulses of the right lower extremity were absent and the ankle-brachial index was 0. On the standardized treadmill exercise test, the right calf claudication began at 67 m.

We proceeded with magnetic resonance angiography, which showed an obstructed right EIA and a collateralized thromboembolic obstruction of the infrapopliteal arteries (Fig 1, A and B). Owing to presence of slightly dilated external iliac arteries of the contralateral side, we opted to perform computed tomography (CT) angiography, which visualized a thrombosed isolated 2-cm large fusiform right external IAA and a 2-cm saccular aneurysm of the left EIA, without involving the abdominal aorta, the common iliac artery (CIA) or the internal iliac artery (IIA) (Fig 2, A-D).

Histopathologic examination of the excised right iliac artery segment showed a mucoid degeneration of the tunica media, confirming that the true aneurysm was caused by CMN.
A mycotic pathology was ruled out. Fig 4, A-C, illustrates the histopathologic findings.

The operative procedure was uneventful. After a short period of observation in a recovery room, we were able to transfer the patient to our regular vascular surgical ward and there was no need for further intensive care or monitoring. Postoperatively, the pain-free walking distance on the treadmill exercise test increased to more than 500 m and the right ankle-brachial index was 0.7. The patient was discharged on postoperative day 6. On discharge, we recommended strict management of cardiovascular risk factors as well as smoking cessation, antiplatelet therapy, with low-dose aspirin and statin therapy.

Additionally, owing the saccular morphology of the left iliac external artery aneurysm, we planned an elective second procedure of open repair in the same manner 1 month after discharge. Here as well, the postoperative course was uneventful and the patient was discharged on postoperative day 5. At 50 days after repair of the right-sided symptomatic EIA aneurysm, the patient was satisfied and regained much of his everyday mobility with a pain-free walking distance of 4000 m. We were also able to show a preserved flow in both IIA in the postoperative magnetic resonance angiography (Fig 5).

DISCUSSION

An IAA is defined as dilatation of the vessel to more than 1.5 times its normal or expected diameter.\textsuperscript{1,7} Isolated IAA include aneurysms of the CIA, the IIA, the EIA and combinations of those, excluding those of the infrarenal abdominal aorta. They frequently involve the CIA, IIA, or both.\textsuperscript{8,9} Aneurysms of the EIA are extremely rare, possibly owing to the different embryological development between the iliac artery segments.\textsuperscript{10}

The majority of patients with isolated IAA aneurysms are male and diagnosed in the seventh and eighth decades.\textsuperscript{7} Their overall prevalence is reported as less than 2% of the general population and the underlying pathology and type of isolated IAA includes degenerative aneurysm, pseudoaneurysm, penetrating ulcer, postdissection aneurysm, mycotic aneurysm, and traumatic aneurysm.\textsuperscript{11}

CMN was first described by Babes and Mironescu in 1910 and since 1928 was established by Gsell and Erdheim as pathologic finding in the media wall related to aortic aneurysm, dissection, and rupture.\textsuperscript{12} CMN is a disorder of the large arteries, especially the aorta, and commonly seem to be an essential feature of several hereditary conditions, such as Ehlers-Danlos and Marfan syndrome. It is characterized by cyst-like lesions within the media, causing degenerative disruptions of connective tissue proteins and smooth muscles of the tunica media, which may lead to deterioration of the arterial wall.\textsuperscript{13}

Isolated IAs, except for their high risk of rupture of up to 40% with high mortality (31%), rarely can cause thromboembolic peripheral events followed by arterial insufficiency and were recognized only after onset of ipsilateral limb ischemic symptoms, which need further angiographic evaluation.\textsuperscript{14} Elective repair is generally recommended if the aneurysm diameter reaches the
**Fig 2.** A–D, Computed tomography (CT) angiography in axial (C) and coronal (B) view with three-dimensional reconstruction (A and D) shows the isolated 2-cm large thrombosed aneurysm of the right external iliac artery (EIA) (blue arrow) and the 2-cm large sacciform aneurysm of the left EIA (white arrow).

**Fig 3.** A and B, Intraoperative photo after exposure of the external iliac artery (EIA) aneurysm. The common iliac and the distal iliac artery are encircled with vessel loops to obtain vascular control. A, Intraoperative photo after aneurysm exclusion and reconstruction with an 8-mm Dacron interposition graft (B). ICA, Iliac common artery; IIA, internal iliac artery.
threshold of 3.0 to 3.5 cm and the type of operative repair depends on patient anatomy, clinical status, and the presence of concomitant abdominal aortic aneurysm.\textsuperscript{7}

We searched under the keyword "external IAA" in the English-language medical literature and we found 65 EIA aneurysms. We only found three cases of true isolated aneurysms of the EIA caused by CMN until January 2020. Each of them was histologically confirmed and the aneurysms were treated with different types of open repair. The Table presents the cases reviewed.

Crivello et al.\textsuperscript{15} who first described the case of the isolated EIA aneurysm in association with CMN, reported a higher operative risk owing to the friability of the artery wall caused by the media degeneration. In 2009, Kato et al.\textsuperscript{16} reported their experiences in the treatment of an EIA aneurysm, but they could not conform the findings of Crivello et al. Similar to the experience of Kato et al, we also did not find fragile vessel walls in our bilateral open aneurysm repair.

In summary, we present another case of an isolated aneurysm of the EIA with the particularity of bilateral occurrence. We managed the symptomatic right EIA aneurysm via open repair and chose to repair the left sided asymptomatic aneurysm 4 weeks postoperatively, despite the absence of current elective repair threshold of 3.0 to 3.5 cm because of the sacciform morphology. We chose an open approach for two main reasons. First, the patient was young, and we did not want to expose him to followed-up CT radiation. Second, the anatomic features of the arteries were not appropriate for endovascular treatment. Our histopathologic findings conform at large with those of the preceding reports of CMN as underlying pathomechanism of development of isolate iliac aneurysms of the EIA. On reviewing the current literature, we must assert to the fact that there remains
limited information of the rare entity and call on vascular clinicians for more publishing so as to establish the characteristics that may be still unknown.

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