INTRODUCTION

Spontaneous isolated dissection of the iliac artery (SID-IA) is a rare pathologic condition. Although the predisposing factors are not fully known, traumatic (trauma [1], extreme physical activity [2], pregnancy [3], and iliac catheterization [4]) and non-traumatic causes (Marfan syndrome [5], Ehlers-Danlos syndrome [6], fibromuscular dysplasia [7,8], and cystic medial degeneration [9]) have been described. The best therapeutic strategy for this condition is still being debated. We present the case of a 59-year-old male with acute right lower limb ischemia characterized by the sudden occurrence of rest pain, hypoesthesia, and paresis. Angiography showed SID-IA extending down to the femoral bifurcation. The patient had no risk factors for SID-IA; however, he survived an electrocution and had arterial hypertension at admission. Endovascular revascularization was successfully performed, with complete restoration of limb blood flow and remission of symptoms. Follow-up ultrasonography at 1 year confirmed stent patency and absence of clinical symptoms. Endovascular stenting is a good therapeutic option for symptomatic SID-IA without rupture.

Key Words: Dissection, Iliac artery, Endovascular technique, Accidental electrocution, Hypertension

CASE

A previously healthy 59-year-old male presented with sudden rest pain, hypoesthesia, and paresis of the right lower limb. At admission an hour after the onset of symptoms, blood pressure was 190/100 mmHg, pulse rate 100 bpm, respiratory rate 12 breaths/min, and SpO2 96%. On physical examination, pulses on the right popliteal, anterior tibial, and posterior tibial arteries were absent, and the right foot was cold. The patient reported a medical history of arterial hypertension for 8 years and an episode of 380 V alternating current electrocution during housework 7 years earlier, with entry point in the right hand and exit point in the right foot, which was not studied with imaging ex-
aminations. He denied high-level sporting exercises or any other trauma aside from the electrocution. No cases of sudden death or connective tissue disorders were reported in his family, and he denied other risk factors for atherosclerosis, such as smoking, diabetes, and dyslipidemia. He did not receive any medical therapy and, on his own admission, he did not check his blood pressure periodically. Blood tests confirmed the absence of hyperglycemia, hypercholesterolemia, and inflammatory or infectious diseases. A total body computed tomography scan (Fig. 1, 2) detected a dissection with a primary entry tear at the origin of the right CIA and compression of the TL at the origin of the superficial femoral artery (SFA) with distal collapse, thereby creating an occlusion of the SFA. No atherosclerotic or non-atherosclerotic wall alterations predisposing to dissection, such as the “string of beads appearance of fibromuscular dysplasia”, were observed.

The patient underwent angiography (Fig. 3) and endovascular revascularization. The procedure was performed through a contralateral femoral access under local anesthesia. Through a 6-Fr sheath, a hydrophilic Terumo guidewire (Terumo Medical Corporation, Somerset, NJ, USA [10]) was used to cross the TL of the dissected CIA into the SFA. The dissected CIA and the external iliac artery were treated with 2 self-expanding overlapped stents, E-Luminexx (Becton Dickinson, Franklin Lakes, NJ, USA [11]); the smaller stent (10 mm×80 mm) was first deployed distally while the larger one (12 mm×40 mm) was overlapped and deployed using the telescoping technique. The left femoral artery access was closed using a MynxGrip (Cordis, Santa Clara, CA, USA [12]) vascular closure device. Completion angiography showed fully restored blood flow into the right iliofemoral axis (Fig. 4).

No reperfusion compartment syndrome occurred after the procedure. The hospital stay was prolonged due to the patient’s high blood pressure, which was successfully managed with olmesartan 20 mg and amlodipine 5 mg once a day. The patient was discharged on the 10th postprocedural day with palpable peripheral pulses. Dual antiplatelet therapy with aspirin and clopidogrel was prescribed for 3 months, followed by lifetime aspirin alone. Follow-up duplex ultrasonography after 1 year confirmed patent stents

Fig. 1. Computed tomographic reconstruction showed dissection from the origin of the right common iliac artery to the femoral bifurcation. Red arrows indicate the entry sites.

Fig. 2. Cross-sectional images of the computed tomography scan showed the dissection of the right external iliac artery.

Fig. 3. Diagnostic angiography before the endovascular revascularization confirmed the right iliofemoral dissection.
DISCUSSION

SID-IA without involvement of the aorta is extremely rare, and most spontaneous cases are related to connective tissue disorders [5-9] or traumatic causes [1-4]. In the present case, the only predisposing factors identified were hypertension and a previous episode of electrocution. Although arterial hypertension has not been described as a risk factor for SID-IA, it is a known risk factor for aortic dissection [13], predisposing to atherosclerosis and increasing shear stress. Considering that the same mechanisms can potentially act on the iliac arteries as well, we hypothesized that arterial hypertension may have played a role in the occurrence of SID-IA. Moreover, there is evidence that electrocution can lead to vascular injury. Endothelial cell membrane perforation of the aorta and pulmonary artery [14] and a few cases of peripheral arterial rupture (e.g., brachial, radial, ulnar, internal mammary, obturator, and carotid arteries) after electrocution have been reported [15]. In these cases, rupture of the artery immediately followed electrocution. Nonetheless, residual damage caused by electric current may occur in areas of fragility. Therefore, despite the considerable temporal gap with the event in our case, it is not possible to exclude electrocution as a predisposing factor. Our point of view on the risk factors and the suggested treatment concerns a pathology that is still poorly reported in literature because of its rarity. This represents the major limitation of our report, but at the same time, it is what makes it interesting. To our knowledge, this is the first case report of arterial hypertension and electrocution as risk factors for SID-IA and enriches the short list of arterial districts known to have suffered damage from electrocution.

Different strategies have been reported for the management of SID-IA, mainly represented by conservative therapy and endovascular or surgical repair [16]. The clinical presentation usually determines the preferred strategy and timing of intervention. Acute limb ischemia, progressive dissection, and artery rupture are classified as complicated dissection and require urgent intervention [17,18]. No guidelines on the treatment of acute ischemia due to iliac dissection are currently available, and the best therapeutic strategies are still being debated, mainly based on evidence derived from aortic dissection. In this context, open surgery (replacement of the dissected aorta with a graft and removal of the intimal tear) is associated with a high incidence of periprocedural complications, such as bleeding, neurological problems, or multiorgan failure, affecting 40% to 80% of patients undergoing open repair. Furthermore, open surgery is associated with a non-negligible rate of mortality, with in-hospital mortality rates ranging from 25% to 50%. On the other hand, the use of endovascular techniques allows less invasive treatment with lower reported periprocedural complications and in-hospital mortality rates (from 2.6% to 9.8%) [18]. Endovascular repair has become the frontline therapeutic option in complicated acute type B aortic dissection [16,18], and its successful use with low periprocedural mortality and morbidity has also been reported in SID-IA [16,17].

The primary objectives of the treatment of SID-IA are to limit the extent of dissection, preserve blood flow distally through the TL, and prevent artery rupture. Endovascular repair by covering the primary entry tear can reduce blood pressure within the false lumen (FL), possibly inducing FL thrombosis and TL re-expansion [13,18], a process called remodeling. This event has the potential to reduce both early and late complications of dissection, including rupture and aneurysmal enlargement [19].

However, some drawbacks of stent implantation must be acknowledged. Stent restenosis, obliteration of side branches [18], and damage to a friable artery with the creation of new entry tears could complicate the procedure, and the implantation could be limited by vessel tortuosity [17]. Moreover, the risk of continued FL perfusion from distal tears must be considered; in these cases, endovascular embolization of the FL can allow successful stent/stent-graft implantation [17].

Another unresolved technical issue involves the length of the artery that should be covered by the endograft [19,20].
Some studies have shown that a short length of arterial coverage is less likely to result in favorable remodeling. When coverage is too short, the distal end of the stent may damage the friable arterial tissue, creating new entry tears due to movement with the cardiac cycle, while a longer length may physically compress the FL [19]. However, there is no definitive evidence that extended coverage is needed to restore distal perfusion [18] and, traditionally, distal tears remain untreated [19]. We prefer complete re-lining above the inguinal ligament, while we avoid using stents below the ligament if possible because of the change in mechanical characteristics. In summary, our experience confirms that coverage of the primary entry tear may be sufficient to treat the dissection and to induce remodeling observable on follow-up ultrasound as a re-expansion of the TL due to the closure of the entry tear and the consequent pressure drop in the FL. In the iliofemoral axis, we use second-level imaging, such as computed tomography, only in case of complications. Considering its low invasiveness, high technical and clinical success rates, and low periprocedural mortality and morbidity [16,17], endovascular treatment could be the preferred therapeutic option in patients with complicated SID-IA without signs of rupture.

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