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A SCOTTISH SHOWER: A CASE OF PHLEGMASIACERULEA DUE TO AN AORTOCAVAL FISTULA CAUSED BY AN ABDOMINAL AORTIC ANEURYSM


Key words: abdominal aortic aneurysm, aortocaval fistula, phlegmasiacerulea

ABSTRACT

The abdominal aortic aneurysm is a rare but life-threatening condition. The most-at-risk population is men aged over 50 years, smokers. This pathology is undetectable until the complications become symptomatic but is often diagnosed too late that’s why its mortality rate is very important, especially when symptoms are atypical or complications are unusual. We report a rare case of phlegmasiacerulea due to an aortocaval fistula caused by an abdominal aortic aneurysm.

BACKGROUND

The abdominal aortic aneurysm (AAA) is a rare but life-threatening condition occurring particularly in malesmokers aged over 50 years [1]. The systematic screening of AAA in this most-at-risk population [2;3] and the introduction of endovascular treatment of ruptured AAA in the last decade allowed to decrease the mortality rate [4] of admitted patients to the emergency department (ED) with a ruptured AAA. Nevertheless, the mortality rate of ruptured AAA treated in emergency situations is substantial (ranging from 20 to 60 %) [5;6]. Hence, emergency physicians should know the specific symptomatology and its various complications in order to diagnose this high risk pathology. We present a rare case of a phlegmasia cerulea due to an aortocaval fistula secondary (ACF) to an AAA.

CASE PRESENTATION

We report the case of a 58-year-old man who consulted at the ED for a sudden on set of pain in the right leg. He declared no past medical history except active smoking (20 cigarettes per day during 20 years or 20 packs per year) stopped five years ago. He clearly reported that he felt an intense pain at the inguinal aspect of the right thigh with a sound of crack when he came out of the shower an hour and half earlier. On arrival, his right lower limb was painful (8/10 on the numeric rating scale for pain) and he had difficulty to walk. He confirmed that there was no recent trauma and no previous pain. Vital signs were: afebrile, heart rate of 124 beats per minute and blood pressure of 156/77 mmHg. The physical examination showed a cold, swelling and blue right leg with marbling (Figure 1). All arterial pulses (femoral, popliteal and pedal) were found bilaterally. We noted
a partial sensitive deficit of the foot up to the knee on the right side and marbling on the abdomen (Figures 2 and 3). A vascular bedside ultrasound showed a proximal deep venous thrombosis (to the primitive iliac vein up to the popliteal vein) allowing to evoke the diagnosis of phlegmasia cerulea (Figure 3). We tried to see if the thrombosis concerned the inferior vena cava and discovered fortuitously an AAA. An enhanced thoraco-abdomino-pelvic CT-scan was immediately performed and confirmed a sacciform infra-renal aortic aneurysm of 7 cm diameter, partially thrombosed with ACF (Figures 4 and 5) and a bilateral distal pulmonary embolism. Blood tests showed an increased white blood cell count (13000 mm$^{-3}$), C-reactive protein (115 mg.L$^{-1}$), uremia (9 mmol.L$^{-1}$), creatinine (115 μmol.L$^{-1}$) and arterial lactate (3 mmol.L$^{-1}$). The patient was immediately transferred to a vascular surgery team in another hospital to be treated. Until the surgery, we controlled the blood pressure with a continuous intravenous urapidil infusion and an analgesic therapy which combined two continuous intravenous infusions of sufentanyl and ketamine. He had an aorto-bi-iliac bypass to repair the aorta and a patch on the inferior vena cava to close the fistula. After
a five hours successful surgery, a blood loss of three liters and a
ten days stay at hospital, the patient has finally totally recovered.

DISCUSSION

AAA disease is a common pathology nowadays in the Western population with a prevalence estimated at 2% to 5% among men aged over 50 years [3;4] and its spontaneous rupture is correlated with a high mortality [4]. The prevalence of ACF in patients with AAA is low, about 2% to 6% [5] but it remains a well-known complication. The typical described clinical presentation (abdominal or lower back pain, pulsatile abdominal mass and abdominal bruit) is in fact not so common (< 50%) [5] and it is more likely to show the association of an abdominal pain, a high-output congestive cardiac failure and a large venous inflow with its complications (swelling and cyanotic lower limb, hematuria, acute renal failure, scrotal edema, priapism, etc.) [6]. We describe an authentic case of phlegmasia cerulea which was the only symptom of ACF due to AAA. This association of pathologies is not common as our review of the literature proves it, with only two similar case found [7;8]. The phlegmasia cerulea, first described by Grégoire in 1938 [9], is an acute and massive deep vein thrombosis which needs an urgent medical, and sometimes chirurgical, treatment to avoid necrosis and limb amputation. In our case, the AAA compressing the inferior caval vein caused obstruction of venous outflow which was responsible gradually to a diffuse pre-thrombotic state of the lower limb venous network. The natural evolution of this state allowed to a total and sudden thrombosis of the right common iliac, common femoral and popliteal veins after the migration of a venous embolism. Cinara et al. [10] described a series of 1698 AAA surgically treated among which only 26 cases with ACF. The operative mortality rate of the 26 ACF was 19.2%. Because of the small number of the combination of both phlegmasia cerulea and ACF cases, it doesn’t exist data about mortality but it is easily understandable that it’s management is particularly challenging and mortality rate is very high. Our patient is one of those exceptional cases who had survived.

CONCLUSION

The aortocaval fistula (ACF) is a rare complication of the AAA. Its treatment is associated with a high mortality rate [8] because it is often discovered late. The phlegmasia cerulea belongs to these unusual features which emergency physicians should know.