Abstract: Introduction – Takayasu disease, also known as “pulseless disease”, is a very rare autoimmune systemic granulomatous vasculitis characterized by the impairment of the great vessels, predominantly of emerging arteries from the aortic arch. Diagnosis is based on clinical and paraclinical aspects, being necessary having at least 3 out of 6 American College of Rheumatology (ACR) criteria. Case report – We present the case of a 36-year-old patient who comes at the emergency service from Sibiu for hypertension (BP: 230/115 mmHg at right arm) and phenomena of hypertensive encephalopathy. Physical exam: the impossibility of measuring blood pressure in the left upper limb and the absence of pulse. Electrocardiography (ECG): sinusual tachycardia, left ventricular hypertrophy (LVH). Echocardiology: LVH, significant swelling aortic arch with absence of the Doppler flow on the left subclavicular artery. Laboratory tests: a significant inflammatory syndrome. Chest radiography: a left lower elongated heart arc and a much enlarged upper mediastinal opacity. Chest computer tomography (CT) angiography: a giant aneurysm of aortic arch and upper third of descending aorta (55 mm); occlusion of the left subclavicular origin artery; thickened vascular walls of aortic arch and emerging arteries; rich collateral circulation. She was diagnosed with the Takayasu disease. Conclusions – Takayasu is a rare disease that can be easily overlooked as a diagnosis if you do not insist on a thorough clinical examination and search of possible causes in young hypertensive patients. Keywords: Takayasu disease, occlusion of the left subclavicular artery, aneurysm of aortic arch and descending aorta.

INTRODUCTION

Takayasu disease, named also "pulseless disease", is an autoimmune systemic granulomatous vasculitis, with unknown etiopathogenesis, which is characterized by the impairment of the great vessels predominantly of aorta and emerging branches. Young women are more prone to develop this pathology which is often developed in 2nd-3rd decades of life (fewer than 15% of reported cases developed the disease later than 40 years old). Despite the fact that it is a rare disease (incidence 2.6 persons/1 million/year) and is observed more frequently in Asian, Central and South America.

Contact address:
Mădălina Urluescu, MD
Cardiology Department, Sibiu Emergency Clinical Hospital, Corneliu Coposu Boulevard, 2-4, 550245 Sibiu, Romania.
E-mail: madalinaurluescu@yahoo.com
countries, we must not ignore it in formulating a differential diagnosis. This is an inflammatory disease and the lesions produced can be stenotic, occlusive or aneurysmal. We report a case of Takayasu disease and the difficulties in approaching this pathology.

CASE REPORT

We present a case of a 36 years old patient, female, from rural environment, known in history with: grade III hypertension and high cardiovascular risk since the age of 28; repetitive paroxysmal supraventricular tachycardias; obesity; ischemic stroke at 32 years old; with repeated hospital admission in various medical services in our country and in Italy. She comes at the emergency service from Sibiu for high levels of blood pressure (BP: 230/115 mmHg at the right arm), phenomena of hypertensive encephalopathy and mild chest pain, being sent to the Cardiology Department where she is hospitalized. On examination, the relevant elements observed were: obesity, the impossibility of measuring blood pressure and the absence of pulse in the left upper limb, warm skin and rich superficial venous circulation at this level (Figure 1), second grade systolic murmur in the mitral area, no murmurs on the large vessels, right hemiparesis with mild hypotonia, horizontal nystagmus. On the ECG we observed sinus tachycardia (106 bpm), LVH. The echocardiography revealed morphologically normal aortic valves, a minor mitral and aortic insufficiency, LVH, aneurysm of ascending aorta, aortic arch and descending aorta (ascending aorta: 40 mm; aortic arch: 50 mm; descending aorta: 53 mm), with the absence of the Doppler flow on the left subclavian artery, without turbulence in the descending aorta characteristic for aortic coarctation. Laboratory tests found a significant inflammatory syndrome (ESR: 80 mm/1h; C reactive protein: 30; fibrinogen: 512 mg/dl), mild hypochromic microcytic anemia. Chest radiography (Figure 2) revealed a left lower elongated heart archery and a much enlarged upper mediastinal opacity.

Imagistic explorations were complemented with the chest CT angiography (Figure 3, 4) which revealed an aneurysm of the ascending aorta (40 mm), of the aortic arch and of the first segment of descending thoracic aorta with a diameter of about 55 mm for a length of 20 cm, without signs of dissection; occlusion of the left subclavian origin artery: thickened vascular walls of the left carotid and along the entire dilated aorta; swelling of the brachio-cephalic trunk and left carotid; king-king of descending thoracic aorta at intervertebral space T8-T9 without damaging the lumen, rich cervico-thoracic, mediastinal and pericardial collateral circulation; arterio-venous shunts between pericardial vessels and the cava vein: small infrarenal aortic aneurysm; significant concentric LVH.

Ocular fundus revealed hypertensive retinopathy. The patient refused coronary angiography and magnetic resonance examination. After the rheumatology consultation carried out in our hospital, we completed the investigations with immunologically tests which revealed high level of complement factor C3 while rheumatoid factor, AntiSm, AntiRo, antibodies and lupus cells were negative. Correlated with clinical examination and other investigations, we have excluded lupus, sarcoidosis and rheumatoid arthritis. Given the clinical and laboratory examinations and considering the fact that we have 4 ACR criteria (Table 1), 1 major (left subclavian artery occlusion) and 6 minor Ishikawa criteria (ESR >20 mm/1h; hypertension; aortic regurgitation; aortic aneurysm of the ascending aorta, of the aortic arch and of the first segment of descending tho-
Unfortunately diagnosed at a late stage in the present case (stage complications). The treatment was completed after 2 weeks in Cluj Napoca Clinical Rheumatology Emergency County Hospital, with methotrexate (20 mg/week) and folic acid, the doses of methylprednisolone being increased to 64 mg/day (following that doses gradually decrease monthly) because the disease is considered severe by biological activity (important nonspecific inflammatory syndrome, ESR still high-90 mm/1h despite already initiated corticotherapy and increased C-reactive protein), by the presence of complications (left subclavian artery obstruction, giant aneurysm of thoracic aorta and stroke in the young woman) and life-threatening (already having important consequences prognosis about quo ad sanationem, quo ad laborem and quality of life). Severe hypertension, impaired fasting glucose, obesity, suggested reducing corticosteroid therapy as soon as possible, leading.

Figures 3 and 4. Computed tomographic angiography images (A, B, C) showing aneurysm of the ascending aorta, of the aortic arch and of the first segment of descending thoracic aorta, king-king of descendent thoracic aorta at T8-T9 intervertebral space, occlusion of the left subclavian origin artery, rich cervico-thoracic, mediastinal and pericardial collateral circulation; arterio-venous shunts; small infrarenal aortic aneurysm.

Figures 3 and 4. Computed tomographic angiography images of the chest, transversal section, showing LVH (A), aneurysm of the ascending aorta, of the aortic arch and of the first segment of descending thoracic aorta, thickened vascular walls and rich collateral circulation (B, C).
to 32 mg/day (one month later) for limiting possible side effects. Thus, the combination of immunosuppressive as early as possible, is one more time justified, given the high activity of the disease and due to preparing for potential vascular surgery which require low doses of corticosteroids and reduced disease activity for a higher rate of success. The most effective, according to the actual data, are anti-TNF alpha and anti-IL6 but they are not available. Cyclophosphamide and azathioprine have greater toxicity and require better adherence to therapy (in the case of our patient collaboration was often difficult), so Methotrexate was preferred for simple dosage, moderate long-term toxicity and easy tracking. It was quickly associated because its effect installs in 8-12 weeks. Unfortunately, despite the efforts, the patient was not compliant because she has discontinued corticotherapy and the treatment with methotrexate for her own initiative after 6 weeks.

**DISCUSSION**

Considering the fact that we are dealing with a young hypertensive patient with associated comorbidities, it raises the question of etiology elucidation of these disorders. Measuring blood pressure in both arms is very important considering the fact that, its absence at the left arm wasn’t noticed in our patient, despite repeated presentations to hospital in our country and in Italy. Hypertension is most probably essential, the patient having in her family history a first degree hypertensive mother. However, the king-king of descending thoracic aorta or reducing compliance of aorta through the inflammation processes and secondary fibrosis (which occurs frequently in the chronic phase by affecting all 3 layers and lymphocytes infiltration, occasional giant cells, mucopolysaccharides, smooth muscle cells and fibroblasts or by elastic tissue destruction) could be the causes of secondary hypertension in this case. Rich presence of collateral circulation is an adaptive response to chronic ischemia caused by occlusion of the left subclavian artery\(^{14,17}\). Rich pericardial circulation on may reveal coronary disease that could not be excluded because the patient refused angiography. Such collateral vessels which limits ischemia, plays an important role in maintaining the blood supply in the affected tissue. In Takayasu arteritis specific immunological markers don’t exist therefore, inflammatory syndrome determination is important for assessing disease activity. Recent studies have investigated different biomarkers (matrix metalloproteinases, II-6, II-18, serum amyloid A) but none is sufficient for disease diagnosing and monitoring\(^{13,16}\). In cases where the progression of the disease is fast and extensive, with inappropriate fibrosis, the formation of aneurysms appears. The detection of aneurysms in Takayasu’s disease is described in literature in 6-9% of the cases, often after 40 year\(^{118}\). The most common locations are the ascending aorta, abdominal aorta, the thoracic descending aorta being much less described\(^{15}\). Uncontrolled long-term hypertension, was observed in all patients presenting aortic aneurysms. The annual risk of aneurysmal rupture is relatively low, ranging between 1 and 7% but hypertension must be aggressively approached in these patients\(^{18}\). Given the large aneurysmal dilatation of the aortic arch and descending thoracic aorta (55 mm), the question was of performing surgery or thoracic aortic repair by endovascular approach (TEVAR) using stent-grafts. Occlusion of the left subclavian artery may benefit from subclavicular - carotid bypass if neurological symptoms appear due to verteobasilar suffering. However, any intervention was postponed considering the state of disease activity and because we couldn’t complete the remission treatment, the patient being noncompliant. Considering the presence of numerous complications, we proposed monitoring by a cardiology checkup at every 3 months. Given the fact that our patient has many complications and she refused to follow any treatment, she hasn’t had a good prognosis since, according to Park et al., the 10 years survival rates are approximately 36% in the cases 2, 6. It is very important to assess disease activity, given the

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**Table 1. ACR criteria: Takayasu arteritis if ≥ three of the above six criteria are present.**

<table>
<thead>
<tr>
<th>Sensibility 90.5%, specificity 97.8%</th>
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<td><strong>American College of Rheumatology (ACR) 1990 criteria</strong></td>
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<td>Age on disease onset &lt;40 years</td>
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<td>Claudications of the extremities</td>
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<td>Decreased pulsation of one or both brachial arteries</td>
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<td>Difference of at least 10 mmHg in systolic blood pressure between the arms</td>
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<tr>
<td>Bruit over one or both subclavian arteries or the abdominal aorta</td>
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<tr>
<td>Arteriographic narrowing or occlusion of the entire aorta, its primary branches or large arteries in the proximal upper or lower extremities, not due to atherosclerosis, fibromuscular dysplasia or other causes.</td>
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presence of vascular complications which place the disease in chronic phase, evidenced by relapses and remissions\textsuperscript{10,12}. CT scanning, MRI (where parietal edema can be a sign of disease activity) and angiography evaluate well the extend of the disease but recent studies suggest that, \textit{18F}-fluorodeoxyglucose positron emission tomography allows diagnosis of Takayasu arteritis earlier in the disease course and appreciates better the disease activity\textsuperscript{5,7,9}. Pentraxin 3 is also a biomarker for identifying disease activity, which seems to be more accurate than erythrocyte sedimentation rate and C reactive protein (which can be increased to 50% in the patients in the chronic phase of the disease and 70% in the active phase)\textsuperscript{11,12}.

CONCLUSIONS

Takayasu’s disease is a rare pathology that can be easily overlooked as a diagnosis if you do not insist on a thorough clinical examination and search possible causes in young hypertensive patients. Early and appropriate administration of the treatment accompanied by lifestyle changes and patient adherence may improve the prognosis and obtain disease remission.

Conflict of interest: none declared.

References