Abstract:
Takayasu arteritis, formerly known as “pulseless disease”, is a chronic idiopathic vasculitis which affects the large vessels in the body. First described in the 1800’s, this rare condition is more commonly found in Asian women in their 40’s. Herein, we report the case of a young woman whose exertional angina and claudication were the initial presentation of active Takayasu arteritis. The importance of modern technology of imaging such as CT, MRI and angiography, can often have paramount importance for confirming a diagnosis and the extent of the pathology.

Key Word: Pulse less, Arteritis, Immunosuppressive, Stenosis.

Introduction:
Takayasu arteritis is a rare, systemic, inflammatory large vessel vasculitis of unknown aetiology that most commonly affects women of childbearing age. It is defined as “granulomatous inflammation of the aorta and its major branches” by the Chapel Hill consensus conference on the nomenclature of systemic vasculitis.1 Because of its predilection for the brachiocephalic vessels, this arteritis has been labeled pulseless disease and aortic arch syndrome.2 Takayasu arteritis is commonly occurs in women younger than 50 yrs of age and can manifest as isolated, atypical and/or catastrophic disease. It can involve any or all of the major organ systems. The disease has been reported in all parts of the world, although it appears to be more prevalent in Asians. Takayasu arteritis is observed more in patients of Asian or Indian descent. Japanese patients with Takayasu arteritis have a higher incidence of aortic arch involvement. In contrast, series from India reports higher incidence of abdominal involvement.3,4 In India, the female to male ratio is as low as 1.6:1.1 The underlying pathologic process is inflammatory and several aetiologic factors having proposed, including spirochetes, mycobacterium tuberculosis, streptococcal organisms, and circulating antibodies due to an autoimmune process. Genetic factors may play a role in the pathogenesis.5 Takayasu arteritis has been reported in pediatric patients as young as age 6 months and in adults of every age. In children, Takayasu arteritis is one of the more common aetiologies of renovascular hypertension.6 Takayasu arteritis is also known as pulseless disease, occlusive thromboarteriopathy and Martorell syndrome.7 The aorta can be involved along its entire length, and although any of its branches can be diseased. The most commonly affected are the

1. Professor of Cardiology, Anwer Khan Modern Medical College (AKMMC), Dhaka.
2. Associate Professor, Department of Anatomy, Bangladesh Medical College, Dhaka.
3. Lecturer, Physiology, Marks Medical College, Dhaka.
4. Junior consultant, Cardiology, Anwer Khan Modern Medical College (AKMMC), Dhaka.
5. Professor of Cardiology, Bangabandhu Sheikh Mujib Medical University (BSMMU), Dhaka.
subclavian and carotid arteries. Although the most frequent pattern of disease varies geographically. Stenotic lesions found in >90% of patients, whereas aneurysm are reported in approximately 25%. Clinical features are generalized as well as localized. Non specific systemic symptom like fever, loss of weight, headache, fatigue, general weakness, night sweat, anorexia, arthralgia, Ischemic pain are present. Splenomegaly, cervical and axillary lymph adenopathy are sometimes present.

Case History:
A 37 years old multiparous lady presented with 6 years history of intermittent claudication in right upper and lower limbs. Patient had a history of exertional dysponea initially of NYHA class – I, later progressed to NYHA class – III. She had been suffering from exertional central compressive chest pain radiating to left upper limb for 4 years. During the last 8 months patient felt vertigo without history of unconsciousness, deafness, tinnitus, visual problem or limb weakness.

She had consulted many physicians for her problems and later admitted to NICVD. On examination patient was found anxious with mild pallor and cold periphery. Her pulse was 80 bpm, regular, absent in both upper limbs and low volume in left lower limb. Blood pressure was unrecordable in both upper limbs whilst BP was 180/110 mmHg in right lower limb. Right Carotid, right Subclavian and left renal bruit were present. Precordium revealed normal findings. Investigation revealed – Hb%-10gm/dL, ESR-80mm in 1st hour, CRP-Positive, CXR-mild cardiomegaly and Echocardiography showed anterior wall hypokinesia with mild LV systolic dysfunction (EF-50%). Doppler study of upper and lower limb vessel showed reduced velocity at left common carotid artery with no flow at left subclavian artery with retrograde flow at both vertebral arteries.

CT aortogram and carotid angiogram revealed complete occlusion of left common carotid artery, left subclavian artery and narrow brachiocephalic trunk distally. Mild stenosis at proximal part of the right common carotid artery with collaterals forming left vertebral and axillary artery was noted. Coronary Angiogram (CAG) revealed 70% stenosis at ramus intermedius. Peripheral Angiogram (PAG) revealed non-visualized left common carotid artery and left subclavian artery with 100% stenosis at right subclavian and 50% stenosis at right common carotid artery. Left common iliac artery revealed 90% stenosis (Fig. 1 & 2).

With this findings patient was diagnosed as a case of Takayasu arteritis with coronary artery disease. Patient was managed with high dose of corticosteroid along with anti-ischaemic management for coronary artery disease.

American College of Rheumatology criteria for the Takayasu's arteritis:

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<th>Criterion</th>
<th>Definition</th>
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<tr>
<td>Age at disease one ≤ 40 years</td>
<td>Development of symptoms or findings related to Takayasu arteritis at age ≤ 40 years</td>
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<tr>
<td>Claudication of extremities</td>
<td>Development and worsening of fatigue and discomfort in muscles of one or more extremities while in use, especially the upper extremities.</td>
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<td>Decreased brachial artery pressure</td>
<td>Decreased pulsation of one or both brachial arteries</td>
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<td>Blood pressure difference &gt; 10 mmHg</td>
<td>Difference of &gt; 10 mmHg in systolic blood pressure between arms</td>
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<tr>
<td>Bruit over subclavian arteries or aorta</td>
<td>Bruit audible on auscultation over one or both subclavian arteries or abdominal aorta.</td>
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<td>Arteriogram abnormality</td>
<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 arteriosclerosis, fibromuscular dysplasia, or similar causes; changes usually focal or segmental.</td>
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For purposes of classification, a patient shall be said to have Takayasu’s arteritis if at least three of these six criteria are present. The presence of any three or more criteria yields a sensitivity of 90.5 percent and a specificity/specificity of 97.8 percent.
Discussion:
Takayasu arteritis is an inflammatory disease of large- and medium-sized arteries, with a predilection for the aorta and its branches. Indian origin aortoarteritis is a chronic granulomatous, necrotizing vasculitis, predominantly affecting the aorta with its branches. The disease is classified based on the site of involvement:

- Type I: Aortic arch involvement
- Type II: Thoracoabdominal involvement
- Type III: Diffuse involvement
- Type IV: Pulmonary involvement
- Type V: Aneurysmal types

The site of arterial disease determines its clinical presentation. There can be dizziness or syncope due to the decreased perfusion to the brain, which can be aggravated by neck movements. This leads to classical drooping position of the head. A classical bruit can be heard over the stenosed carotids. Cardiac involvement in the form of myocardial infarction, valvular pathology, conduction system block or coronary artery involvement can be seen. Pulmonary vasculitis can lead to pulmonary hypertension and ventilation perfusion abnormalities. The other associated findings are renal artery stenosis and musculoskeletal involvement in the form of rheumatoid arthritis and ankylosing spondylitis. Ishikawa graded TA depending on the presence of four major complications, i.e., hypertension, retinopathy, aneurysm formation and aortic regurgitation. Hypertension, the major complication affecting anaesthetic management in patients with TA, is commonly renovascular. It could also result from reduced elasticity and marked narrowing of aorta and major arteries and abnormal function of carotid and aortic sinus baroreceptors. Measurement of blood pressure and proper documentation of all pulses deserves special attention in such patients. NIBP monitoring using the oscillometric method and pulse oximetry can provide simple and reliable blood pressure readings even in patients with pulseless extremities.

Histologic examination during active stages of the disease discloses a granulomatous arteritis similar to giant cell arteritis and to the aortitis associated with the seronegative spondyloarthropathies and Cogan syndrome. In later stages, medial degeneration, fibrous scarring, intimal proliferation, and thrombosis result in narrowing of the vessel, yet there remains a lack of adequate histopathologic criteria for the differential diagnosis of noninfectious arteritides, including Takayasu disease and giant cell aortitis. Aneurysm formation is less common than stenosis, but aneurysm rupture is an important cause of death in patients with Takayasu arteritis. Angiographically, the left subclavian artery is narrowed in approximately 90 percent of patients. The right subclavian artery, left carotid artery, and brachiocephalic trunk follow closely in frequency of stenosis.

Corticosteroid therapy appears effective in suppressing inflammation during the active phase, and favorable results have been reported with immunosuppressive and cytotoxic agents, like Cyclophosphamide & Methotrexate.
Warfarin, Aspirin and Plasmapheresis may be sometimes helpful. Operative treatment may be employed to relieve symptoms caused by arterial obstruction, and percutaneous angioplasty and stenting are associated with favorable results. These procedures are best reserved for patients in whom the acute inflammatory stage of the disease has been controlled.

Medical treatment, Balloon Angioplasty with or without Stenting and surgery is the treatment of Takayasu arteritis. Prednisolone is usually used. Cyclophosphamide, Azathioprine, Methotrexate, Dapsone appears to be useful. Aspirin and warfarin reduces the ischemic symptom.

Follow up of this peripheral vascular disease is very essential. Disease course may be slowed by drugs and some times surgical intervention may be helpful. During follow up ESR and CRP may be done which has got the prognostic value.

Prognosis is some times good, usually downhill course. Immunosuppressive drugs have got good prognostic value.

**Conclusion:**

Takayasu disease is not very uncommon. It affect usually female. It is disabling disease; gradual downhill course of the diseases makes the patient crippled. Early diagnosis, high dose steroid and vascular intervention or surgical treatment can help the patient to some extent.

**Reference:**