CASE REPORTS

Prolonged Fever in Adult Still’s Disease – A Case Report and Review of Literature

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Introduction:
In 1897, George Still described a special form of arthritis in children, what is today called Still’s disease. In Still’s original description, he attempted to distinguish a form of chronic joint disease in children from rheumatoid arthritis in adults. The distinct features included pyrexia, splenomegaly and enlargement of lymph nodes. Still’s did not describe the presence of rash, an important feature in the diagnosis of Still’s disease today. The similar illnesses were found in adults and reported by many clinicians, some of which as undiagnosed fever or pyrexia of unknown origin. The first use of the term “adult Still’s disease” was by Eric Bywaters, the eminent English Rheumatologist in 1966.1

Adult Still’s disease (ASD) or adult-onset Still’s disease (AOSD) is a rare, immune-mediated, multisystem inflammatory disorder of unknown etiology characterized by daily high spiking fever, evanescent rash and arthritis, frequently accompanied by sore throat, lymphadenopathy, splenomegaly and neutrophilic leucocytosis. It is frequently underdiagnosed and remains one of the main reasons for hospital admissions due to pyrexia of unknown origin.2 The disease commonly affects young individuals between 16 and 35 years of age and male and female are affected equally.3 The clinical picture is variable with mild to life threatening courses. The disease is self-limiting, intermittently active or chronic.4 Diagnosis is clinical and may be lengthy because it requires exclusion of infectious diseases, neoplasms including lymphoma and leukaemias, and autoimmune diseases. High serum ferritin levels associated with a low fraction of its glycosylated component are useful diagnostic and disease activity markers.5

Case Report:
A 22 - year old married female, short stature, housewife, low socioeconomic status, mother of a one year child, from a rural area of Bangladesh, was admitted into our female medical ward with history of high intermittent fever with chills and rigor, polyarthralgia, sore throat, bodyache and weight loss for 3 months. The fever was high spiking, 103 degree F or above, rose rapidly in the evening and then subsided to or near normal by several hours or the next morning (Fig.-1a). In some days, two fever spikes were noticed. Her sore throat was described as a severe, constant burning pain in the area around pharynx, more prominent during febrile period. Physical examination showed an emaciated (weight-29 kg), toxic girl with moderate anemia, no skin rashes, swollen and tender both wrists, left ankle joint, cervical
lymphadenopathy, mildly enlarged non-tender liver and splenomegaly. Cardiovascular and respiratory system examination revealed normal findings. Deep cervical lymph nodes in the left anterior triangle of the neck were palpable, 3 in number, size 1 – 2 cm in diameter, soft in consistency, non tender, freely mobile and not fixed to underlying structures or overlying skin. During her long febrile period, she was treated with phenoxymethylpenicillin, levofloxacin, doxycycline, cefixime, antimalarial drugs before hospital admission without any benefits. Her laboratory investigations were as follows: Complete blood count & peripheral blood film - Hb 9.3 gm/dl, ESR 106 mm/1st hour, RBC-3.81 million/cumm, MCV-74fl, MCH-24pg, WBC-30,000/cumm (Neutrophil - 87%, L -11%, M - 2% E - 0%), Platelet – 487,000/cumm, Malaria parasite - not found; Widal Test - TO – 1:80, TH – 1:80, AO – 1:40, AH-1:40, BO- 1:40, B H- 1:40, and after 2 weeks, no rising titre; blood culture twice yielded no growth of any bacteria, ASO Titre - 107 IU/ml, S. Creatinine - 0.85 mg/dl, Random blood sugar - 6.2 mmol/L; Urinalysis - normal; Urine Culture repeatedly - no growth. X-ray chest, ECG and transthoracic echocardiography showed normal findings. Abdominal ultrasound revealed normal findings except mild hepatosplenomegaly. Following hospital admission, she was treated empirically with quinine and then ceftriaxon without any improvement. As the patient was not recovering, she was further investigated, some investigations were repeated and the results were as follows: Complete blood count - Hb 8.6 gm/dl, ESR 108 mm/1st hour, WBC-26,000/cumm (N -82%, L-16%, M - 01%, E - 01%); RA Test – negative, ANA - negative, Anti CCP - negative, CRP - 76 mg/L, VDRL – nonreactive, Tuberculin Test - 05 mm, S. Ferritin - 2400 ng/ml (highly elevated), S. Bilirubin-0.4 mg/dl, S. ALT- 26U/L, Urinalysis – normal, S. Fibrinogen- 394 mg/dl, X-ray chest and paranasal sinuses - normal. Serum protein and haemoglobin electrophoresis, bone marrow aspiration was under plan and waiting. Infectious and neoplastic causes of her illnesses were tried to be ruled out by a thorough clinical and laboratory reports evaluation.

Literatures were reviewed and a diagnosis of Adult onset Still’s Disease (AOSD) was made considering Yamaguchi criteria. Tablet prednisolone (1mg/kg/day) and methotrexate (7.5mg weekly) orally were started. Her condition was dramatically improved. She became afebrile (Fig.-1b), appetite and general condition improved.

Subsequent investigations after 6 weeks showed the following: CBC Hb-10.2g/dl, WBC-14000/cumm (N-70%, L-25%, M-4%, E-1%), ESR-15 mm in 1st hour, CRP-6 mg/L.

Follow up visit after 6 month: Healthy smiling girl with mildly tender both wrists with slightly restricted
movements; regularly taking methotrexate 7.5mg weekly orally and laboratory investigations as Hb 11.1g/dl, ESR 20 mm in 1st hour, CRP 6 mg/L, S. Ferritin - 11.5ng/ml.

Follow up visit after 2 years: Reasonably good health but mild tenderness involving both wrists on passive movement, no fever and no other symptoms, CBC- Hb- 12.7g/dl, WBC- 13660/cumm (N-70%, L- 24%, M-06%), ESR- 10mm in 1st hour, CRP – 3mg/L, ANA-Negative, S. Ferritin- 18.2ng/L.

**Discussion:**

Data on diagnosis and treatment of Adult Still’s Disease are limited in medical literature and consists of mainly case reports, small series and some retrospective studies; current knowledge is largely descriptive. Diagnosis is clinical and requires exclusion of infectious, neoplastic and connective tissue diseases. Laboratory tests are nonspecific and show increased immunological response. There is a speculation that the disease has features of nonnecrotising immune complex vasculitis. The disease occurs throughout the world, typically affects 16-35 years old and presents with fever, arthralgia, and sore throat, and skin rash. The fever is typically high and spiking, usually quotidian and occasionally double quotidian. The spike occurs in the late afternoon or evening. The duration of fever is typically brief, and the temperature falls rapidly.

The rash is perhaps the most helpful feature in the diagnosis of Still’s disease, especially when present in association with high fever and arthralgia. It is predominantly a truncal rash but can spread to the arms and legs including the palms and soles, but usually does not involve the face. The typical eruption is a salmon-pink macular or maculopapular rash beginning as small macules that may coalesce; the rash usually is not pruritic, evanescent and may be missed. Patients may not notice it. We did not notice any skin rash in this patient, probably because of black complexion.

**Diagnosis of Adult Still’s Disease**

There is no single diagnostic test for Adult Still’s disease and the diagnosis is clinical and not based on serology. Response to empirical corticosteroid therapy generally helps in diagnosis. Laboratory investigations reveals high WBC count with neutrophilic leukocytosis, raised ESR, high CRP, very high ferritin, high level of AST, ALT and Fibrinogen reflecting systemic inflammation. Rheumatoid factor (RF) and antinuclear antibody (ANA) are classically negative.

Recently, serum ferritin and glycosylated ferritin have received a lot of attention as diagnostic and disease activity marker. The glycosylated fraction of ferritin is a more specific marker of AOSD than ferritin itself. In this patient, glycosylated ferritin was not estimated due to unavailability. In healthy subjects, 50-80% of ferritin is glycosylated; in inflammatory diseases, 20 – 50% of ferritin is glycosylated and in AOSD, less than 20% is glycosylated. Cytokines, such as interleukin(IL)-1, IL-6, interferon(IFN)-gamma, and tumor necrosis factor-alpha, are elevated in patients with AOSD.

At least 7 sets of classification criteria have been devised; of which Yamaguchi, Cush and Fautrel criteria are important. However, the Yamaguchi criteria have
the highest sensitivity. Diagnosis requires at least 5 criteria, with at least 2 of these being major diagnostic criteria.³

Major criteria
• Fever of at least 39 degree centigrade for at least one week
• Arthralgia or arthritis for at least two weeks
• Nonpruritic salmon colored rash (usually over trunk or extremities while febrile)
• Leukocytosis (10000/microL or greater), with granulocyte predominance

Minor criteria
• Sore throat
• Lymphadenopathy
• Hepatomegaly or splenomegaly
• Abnormal liver function tests
• Negative tests for antinuclear antibody and rheumatoid factor

Exclusion criteria
• Infection
• Malignancies
• Other rheumatic diseases

The reported case had almost all the features of Yamaguchi criteria. Skin rash was not obvious, probably because of black complexion. So, a diagnosis of Adult onset Still’s Disease was made.

Treatment of AOSD includes NSAID, corticosteroids, immunosuppressive drugs and biological agents. Immunosuppressants including methotrexate, gold, azathioprine, cyclosporine, leflunomide, terolimus, and cyclophosphamide; along with intravenous immunoglobin are tried. Biological agents or anticytokines (e.g. anti-TNF-alpha, anti-IL-1, anti-IL-6) have been successfully used in refractory cases.²,⁶⁻¹⁸ We treated the patient with NSAID, corticosteroids and methotrexate with satisfactory response.

The clinical course of AOSD can be divided into three main patterns with different prognosis: self-limited or monophasic, intermittent or polycyclic systemic and chronic articular pattern. The reported case seems to be chronic articular pattern. AOSD, though not common, can be from milder form of disease to a life threatening condition.¹⁹⁻²¹

Macrophage-activation syndrome (MAS) is a severe, potentially life threatening complication, associated with Adult-onset Still’s disease. The hallmark clinical and laboratory features include high fever, hepatosplenomegaly, lymphadenopathy, pancytopenia, liver dysfunction, disseminated intravascular coagulation, hypofibrinogenemia, hyperferritinemia and hypertriglyceridaemia. The condition is triggered by a viral infection or a medication. Despite marked systemic inflammation, ESR is depressed because of low fibrinogen level.²²⁻²⁴ Occasionally, AOSD may be complicated by thrombotic thrombocytopenic purpura and haemolytic uraemic syndrome.²⁵

Conclusion:
Adult onset Still’s disease (AOSD) is a rare systemic inflammatory disorder of unknown etiology and pathogenesis, usually presenting with high spiking fever accompanied by systemic manifestations. It is hypothesized that it may be a reactive syndrome where various infectious agents may act as disease triggers in a genetically predisposed host. The suggested etiologies, clinical manifestations and prognoses are diverse. There is no single diagnostic test for AOSD; rather, the diagnosis is based on a set of criteria, the most important of which are indeed clinical. Treatment aims at both minimizing inflammation and halting disease progression. Non-steroidal anti-inflammatory drugs (NSAID), glucocorticoids and disease modifying antirheumatic drugs (DMARD) are used. Novel therapeutic approaches, such as anti-tumor necrosis factor blockade and stem cell transplantation, monoclonal antibodies, may be effective.

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