Visceral Leishmaniasis Presented as Myelofibrosis and Low grade lymphoma in a Sporadic Region of Iran, Report a Rare Case

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Abstract
We describe a case of leishmaniasis in a 55-year-old male who presented with weakness, fever and anemia. The patient was born and lived all his life in Talaghan, a non-endemic region of kala-azar and there was no history of travel to endemic region for leishmania. In primary diagnosis, the patient suspect has been myelofibrosis and then lymphoma and underwent chemotherapy. His general condition worsened and bone marrow biopsy was performed again and leishmania promastigotes seen in the bone marrow. Specific identification of the parasite was done by RAPD-PCR. After two weeks of treatment, he was transferred to ICU due to heart attack and after 4 days he died due to aortic valve ABE.

Keywords: Visceral leishmaniasis, Myelofibrosis, Low grade lymphoma, Sporadic region

Introduction
Visceral leishmaniasis (VL), known also as Kala-azar, is a chronic infectious disease caused by a protozoan parasite of the genus leishmania; it is transmitted by fly bite.(1) The most highly endemic areas of VL in Iran are parts of Fars and Bushehr Provinces in the south, the districts of Meshkin-shahr and Moghan in northwest, and Qom Province in central Iran. Other parts of Iran are considered as sporadic areas for VL. Visceral leishmaniasis is common (over 98%) among children under 12 years old in different endemic foci in Iran and adult cases frequently present with subclinical and asymptomatic forms in endemic regions.(2, 3) In Iran, the visceral form is mainly due to Leishmania infantum,(3) and the main reservoir of this parasite is infected dogs.(4) Delay and difficulty in diagnosis are common, especially at its early stage of the disease.(5) Here we present a case of visceral leishmaniasis in which the patient was misdiagnosed as myelofibrosis and low grade lymphoma from sporadic region.

Case report
A 55-year-old man who was a resident of Talaghan, an non-endemic region of visceral leishmaniasis in central of Iran, was admitted to the Department of Hematology in Imam Khomeini Hospital affiliated to Tehran University of Medical Sciences in January 2007 with the presentation of weakness, fever, weight loss and anemia. The case was suspected to have myelofibrosis and underwent splenectomy in the other center 7 months before admission to this hospital. Despite of treatment, the
white blood cells (WBCs) $9.9\times10^9$/L. Liver enzymes revealed lactate dehydrogenase (LDH) 542 u/l, SGPT (ALT) 20 u/l, SGOT (AST) 64 u/l, alkaline phosphatase 579 u/l, total bilirubin 2.6 mg/dl, and direct bilirubin 2 mg/dl. Additional blood tests showed triglycerides of 326 mg/dl, Fasting Blood Sugar 133 mg/dl, and Blood Urea Nitrogen 36 mg/dl. Diagnostic tests such as hepatitis B, hepatitis C, tuberculosis (TB), HIV, and serological testing for leishmania antibodies were performed and they were negative. Urine and stool examination revealed no significant pathology and peripheral blood smear was negative for microorganism. Patient was repeatedly suffered from weight loss, so that patient weight was 40 Kg. Bone marrow biopsy was performed and a diagnosis of lymphoma was suspected. After 4 weeks his general condition worsened and bone marrow biopsy was performed again and leishmania promastigotes seen in the bone marrow. (Figure-3) Leishmania promastigotes were analyzed by RAPD-PCR technique and type infantum were reported. However there was no history of travel to endemic region for leishmania or insect bite. All past pathologic preparations were thoroughly reviewed, including bone marrow examinations and liver biopsy. Liver biopsy revealed Leishmania parasites.

After definitive diagnosis of leishmania, treatment was initiated with pentavalent antimony 20 ml/kg daily. It was accompanied with gradual improvement in clinical condition of the patient, and the body temperature returned to the normal value after the seventh day of treatment. The patient was well on discharge. After two weeks he
presented with cough and weakness. Patient was febrile; the temperature was 39°C, and asever hypotension that did not respond to dopamine. Patient was transferred to ICU due to heart attack and alteration in orientation level which caused to coma. Blood culture was positive for staphylococcus and after 4 days he died due to aortic valve ABE. (Figure- 4)

Discussion
There are three important endemic region of VL in Iran: Ardebil, East Azerbaijan, and Fars provinces, and some sporadic region.(6) During 1998–2006, approximately 2,056 cases of VL were reported in Iran, and Ardabil Province contained approximately 30.4% of the cases. More than 90% of VL cases are reported in the pediatric age group and is usually characterized by prolonged fever, hepatosplenomegaly, reversed albumin-to-globulin ratio and proteinuria.(7,8) The first case of visceral leishmaniasis in Iran was reported by Pouya in Mazandaran Province during 1949.(9) Visceral leishmaniasis (VL) or kala-azar caused by L. donovani, L. infantum, L. chagasi, L. amazonensis and rarely L. tropica minor.(10) In humans, L. infantum causes a wide spectrum of clinical manifestations, from asymptomatic or oligosymptomatic infection to acute or chronic disease.(11)

Previous case reports indicate that visceral leishmaniasis can be misdiagnosed as myeloma,(12,13) mixed cryoglobulinaemia(14) malignant lymphoma,(15) lymphoblastic leukemia,(16) and autoimmune hepatitis.(17) Rocha(18) reported that the bone marrow fibrosis is transient (pseudo-myelofibrosis) in human kala-azar and these patients are expected to show regression after treatment. Therefore the presence of associated myelofibrosis it could easily have been mistaken for chronic idiopathic myelofibrosis. Our case was a 55-year-old man living in Talaghan, central Iran, and a non-endemic region of kala-azar. In primary diagnosis, the patient suspect has been myelofibrosis and then lymphoma.

The diagnosis depends on serological studies or direct parasite detection.(19) Spleen biopsy is reported diagnostic of VL in over 95% of cases, liver biopsy and bone marrow sensitivity is reported to be about 70% .In general, serology is not reported to be a very useful test in both visceral and cutaneous leishmaniasis as it may give false positive as well as false negative results.(20) Gagnaire reported that the Leishmania serological test and bone marrow examination included in the work-up may be initially negative, necessitating repeated bone marrow procedures for diagnosis.(22) Serology test was negative in this case. In addition, bone marrow biopsy was performed during admission and leishmania diagnosed on the third bone marrow aspirate (after two previous negative examinations). He was born and lived all his life in Talaghan and there was no history of travel to endemic region for leishmania and he denied having been bitten by insects. Several studies suggest that infection by L. infantum can exist in a subclinical form in healthy individuals without a previous background of cutaneous or visceral leishmaniasis and those with asymptomatic infection can act as reservoirs for parasites.(2,23,24) So it is speculated that he was infected by asymptomatic individuals. Several results have been reported that the undertaken chemo therapy may weaken the immunity system leading to a viscerotropic infection.(25,26) Bacterial infections are often seen in patients with visceral leishmaniasis and Pseudomonas aeruginosa and Staphylococcus aureus are the most common agents.(27) In this case, the patient’s age, an initially negative assay for Leishmania, living in sporadic region and no history of presence in endemic areas caused the disease undiagnosed. After suspicion to lymphoma, treatment with chemotherapy undermines the immune system and disease progression. Blood culture was positive for staphylococcus and patient died due to aortic valve ABE that is one of the major complications of leishmania.

Conclusion
Early detection of the infection is necessary in order to start effective treatment and prevent more serious complications. Therefore the clinical presentation should be considered in the differential diagnosis of malignant neoplasms.

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References