Leishmaniasis-Associated Hemophagocytic Syndrome Revisited: Not an Uncommon Clinical Presentation of Leishmaniasis

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Abstract

Visceral leishmaniasis (VL) is an uncommon but important cause of reactive hemophagocytic syndrome (HS) that should be seriously considered in patients coming from endemic areas. We describe two cases of VL-associated HS; a young woman with fever, cytopenias and hyperferritinemia and an immunocompromised patient with fever, splenomegaly, hyperferritinemia and pancytopenia. The diagnosis of leishmaniasis was established with polymerase chain reaction for leishmania and bone marrow examination, whereas leishmaniasis specific treatment with liposomal amphotericin led to full clinical recovery and complete remission of HS. From the clinical point of view, it should be emphasized that the high clinical suspicion along with the use of modern, high yield diagnostic tools, may lead to early diagnosis of VL-associated HS, minimizing unnecessary hospitalization and potentially harmful investigations and treatments.

Keywords: Visceral leishmaniasis; Hemophagocytosis; Hemophagocytic syndrome; Immunosuppression

Introduction

Hemophagocytic syndrome (HS) is an uncommon but life-threatening condition characterized by high fever, lymphadenopathy, hepatosplenomegaly, cytopenias, liver dysfunction and hyperferritinemia. It is caused by a dysregulation in natural killer cell function, leading to an uncontrolled activation of histiocytes/macrophages and lymphocytes, cytokine overproduction and hemophagocytosis [1]. HS can be either primary, due to genetic mutations, or secondary, associated with malignancies, autoimmune diseases and infections. Among the infectious agents, Epstein Barr Virus (EBV) is the most common cause of the syndrome. However, HS can also be associated with other viral infections (other herpes viruses, HIV, influenza, parvovirus, hepatitis viruses) as well as with bacterial, fungal and parasitic infections, including visceral leishmaniasis (VL) in endemic areas [2].

Herein, we describe two cases of VL-associated HS, followed by a short review of the literature.

Case Report

Patient 1

A 22-years-old previously healthy woman was admitted to our department with a 4-day history of fever up to 39 °C, malaise, anorexia and nausea. She also mentioned several episodes of frontal headache without neck stiffness or photophobia, arthralgias, as well as an episode of epistaxis.

The physical examination revealed only mild splenomegaly, while the routine laboratory tests revealed leukopenia (3,500/μL; normal range 4,000 - 10,000/μL), thrombocytopenia (75,000/μL; normal range 150,000 - 400,000/μL), an increase of lactate dehydrogenase (LDH 556 IU/L; upper normal limit (UNL) 230 IU/L), an increase of creatinine phosphokinase (CPK 273 IU/L; UNL 145 IU/L), and normal serum troponin and D-dimer. A chest radiograph was normal, while electrocardiography revealed sinus tachycardia.

During the first week of hospitalization an extensive diagnostic work-up was performed. Blood, urine, bone marrow and cerebrospinal fluid cultures were sterile, whereas serological studies for Brucella, Leishmania, Leptospira, Enteroviruses, Adenoviruses, hepatitis A, B, C viruses and HIV infection tested negative. Testing for EBV and other herpes viruses were consistent with past infection and serum polymerase chain reaction (PCR) for EBV and CMV DNA was negative. Immunological tests (antinuclear antibodies, anti-neutrophil cytoplasmic antibodies, rheumatoid factors) were also negative.
The patient initially received ceftriaxone empirically. However, fever and the other symptoms persisted whereas laboratory tests showed deterioration of the hematological parameters. On clinical suspicion of HS, biochemical and coagulation tests were performed that revealed an extremely high level of ferritin (24,000 ng/mL; normal range 20 - 400 ng/mL) and a mild decrease of fibrinogen. Myelogram confirmed extensive hemophagocytosis. A diagnosis of HS was established, as 6 out of 8 diagnostic criteria for HS were fulfilled: fever, splenomegaly, cytopenias, hyperferritinemia, hypofibrinogenemia and hemophagocytosis in the bone marrow [3].

HS was initially attributed to adult Still’s disease, as the majority of the infectious, immunological and neoplastic diseases related to HS had been ruled out by the aforementioned tests. So, the initial treatment consisted of corticosteroids and gamma globulin infusions as we have described recently [4]. However, from the clinical and laboratory points of view, the patient deteriorated further.

Treatment failure led to a reconsideration of the diagnosis. Other causes of HS were considered, including VL, an endemic parasitosis in Greece that can be manifested as HS. Peripheral blood PCR for Leishmania donovani was positive, while a second bone marrow aspiration revealed Leishmania parasites in macrophage cytoplasm. As the diagnosis of VL-associated HS had been established, the patient received liposomal amphotericin (daily dose 4 mg/kg body weight on days 1 - 5, 14 and 21) leading to a full clinical recovery and restoration of hematological parameters. The patient has been on follow-up by our department for more than 2 years, and no relapse has been noticed.

**Patient 2**

A 40-years-old man was admitted to our department due to 20-days history of high fever and cytopenias. The patient suffered from myasthenia gravis, having a close follow-up by the neurology department. He was under immunosuppressive treatment with prednisone and azathioprine combined with gamma globulin infusions. The patient had been in his usual state of health until 20 days before admission, when he started having everyday fever up to 40 °C along with night sweats, anorexia and weight loss.

Physical examination revealed a 2/6 systolic murmur at the right upper sternal border and splenomegaly. On initial laboratory tests, the total blood count revealed pancytopenia: white cell count 2,300/µL, hemoglobin 8 gr/dL (normal range 12 - 16 gr/dL), platelet count 75,000/µL. Biochemical tests showed mild transaminasemia (aspartate transaminase and alanine transaminase levels of 52 and 43 IU/L respectively; normal range 10 - 35 U/L), a moderate increase of inflammation markers (C-reactive protein 11.3 mg/dL; UNL 0.5 mg/dL) and significant hyperferritinemia (6,329 ng/mL) and hypergamma globulinemia (4.5 g/dL; UNL 3.5 g/dL).

The differential diagnosis included the common infectious and neoplastic causes of febrile splenomegaly. Subsequently, an extensive investigation was performed that ruled out infective endocarditis, brucellosis, leptospirosis, EBV, CMV, HIV or Parvo-virus infection as well as lymphoproliferative disorders. However, due to the typical findings such as fever, splenomegaly, pancytopenia, hypergammaglobulinemia and history of immunosuppression, VL was considered as probable diagnosis. Therefore, peripheral blood and bone marrow PCR for leishmania DNA was performed that were positive, whereas myelogram revealed the presence of leishmania parasites in macrophage cytoplasm, establishing the diagnosis of leishmaniasis. Furthermore, according to the aforementioned criteria of HS [3], a diagnosis of VL-associated HS was established since 5 out of 8 criteria for HS diagnosis were fulfilled [3]; fever, splenomegaly, cytopenias, hyperferritinemia, low natural killer activity as estimated by immunophenotyping.

As the patient was immunocompromised, he received an intensive regimen of liposomal amphotericin (daily dose of 4mg/kg body weight on days 1 - 5, 10, 17, 24, 31, 38) to prevent a possible relapse. Specific treatment resulted in full resolution of patient’s symptoms and signs as well as complete remission of HS.

However, despite the intensive treatment, patient’s symptoms reemerged a few months later and a relapse of VL was confirmed by peripheral blood and bone marrow PCR for leishmania. A second intensive regimen of liposomal amphotericin was administered, followed by prophylactic monthly doses of liposomal amphotericin (4 mg/kg body weight) to prevent new relapses. Since then, the patient is free of symptoms for 8 months while hematological abnormalities have been fully restored.

**Discussion**

We described two cases of VL-associated HS from Greece, a country endemic for this parasitosis. To the best of our knowledge, this is the second report of secondary HS due to VL reported from Greece, the first one having been published more than ten years ago [5].

The association of VL with HS has been known since the late 70s, when a first case of VL mimicking histiocytosis was reported [6]. Since then, only a few cases of HS secondary to VL have been reported, and the majority of them were infants or children living in endemic areas [7]. However, HS as a pathologic finding in VL might be quite more common. In a recent case series examining the hematological features of the bone marrow examination associated with VL, HS was found in 12 out of 16 (75%) patients examined [8].

The recognition of HS can be a diagnostic challenge, as clearly demonstrated in the first case described. According to a recent systematic review of VL-associated HS, the
mean delay in the diagnosis was 9 weeks [7]. This delay is attributed to the overlapping clinical features between VL and primary histiocytosis, as well as to the low sensitivity of diagnostic methods for VL, such as bone marrow examination and serologic testing. The moderate yield of the ‘traditional’ diagnostic tools may hinder the diagnosis. In the same systematic review of Rajagopala et al, the first bone marrow aspiration failed to establish the presence of Leishmania donovani bodies at onset in 36.3% of cases examined, while the diagnostic yield of serology was also low (23.5%) [7]. In another French pediatric series of VL, the parasite was not detected in 22% of the patients, whereas there were cases with Leishmania donovani bodies recognized on the fifth bone marrow aspiration [9, 10]. These diagnostic difficulties are confirmed in the first patient of the present article, where both bone marrow aspiration and serology were initially negative, leading to further investigations and potentially harmful treatment. Therefore, on a strong clinical suspicion for VL-associated HS, a repeated bone marrow evaluation, and especially the use of highly specific and sensitive PCR assays for leishmania might be necessary [11].

The second patient highlights the increasing problem of opportunistic intracellular infections in immunocompromised patients, as a result of the HIV epidemic and the widespread use of enhanced immunosuppressant therapy for various diseases. It has been estimated that HIV related immunosuppression increases the risk of reactivating VL by 100 - 1,000 times [12], while there are many cases of disseminated VL in immunocompromised patients due to corticosteroid therapy, immunosuppression after solid organ transplantation, or hematologic neoplasia [13, 14]. However, HS is probably an atypical manifestation of VL in immunocompromised patients, as till now only two cases of VL-associated HS in immunocompromised patients (one due to immunosuppressant therapy for Harada’s disease and the other due to HIV infection) have been reported [15, 16].

Our cases illustrate the therapeutic challenges of VL-associated HS, both in immunocompetent and immunocompromised patients. Although immunosuppression schedules including etoposide regimen [3] or combination treatment with corticosteroids and intravenous immunoglobulin infusions [4] are used for life-threatening infectious-related HS, the case of VL-associated HS is probably the only type of reactive HS where pathogen-specific treatment is sufficient to control the disease [2]. On the other hand, if the diagnosis of VL eludes, the use of immunosuppressant and chemotherapeutic drugs usually administered for familial and other reactive types of HS, may lead to a dramatic clinical and hematological deterioration, as illustrated in our first case. Therefore, VL should be ruled out in all patients with HS living in endemic areas before immunosuppression is started [17].

Treatment of VL in immunocompromised individuals, including the few cases manifested as HS, consist an unresolved question, due to the poor results of conventional therapy in these cases. Clinical failure and mortality is high, while many patients experience one or more relapses, as in our second case described [14]. Therefore, an intensive regimen of anti-parasitic treatment and especially of liposomal amphotericin [14] should be considered in any immunocompromised patient suffering from VL with or without HS. In the case of relapse despite intensive treatment, a prophylactic regimen (using amphotericin or pentamidine) is an option [18, 19], although there is lack of evidence from large clinical trials to support it.

**Conclusions**

Although traditionally VL is considered as an uncommon cause of reactive HS, this association might be quite more common [8] and due to the high mortality of HS if untreated, it should be seriously considered in patients coming from endemic areas for VL. The correct and early diagnosis will lead to pathogen-specific and highly effective treatment. Therefore, the high clinical suspicion as well as the use of modern, high yield diagnostic tools, such as PCR, may lead to early diagnosis of VL-associated HS, minimizing unnecessary hospitalization and potentially harmful investigations and treatments.

**Conflict of Interest**

There is no conflict of interest in this paper.

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