Hemophagocytic Syndrome Associated with Cytomegalovirus Infection in a Severely Immunocompromised AIDS Patient: Case Report

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Hemophagocytic syndrome is a clinical condition characterized by the infiltration of the bone marrow and reticuloendothelial system by macrophages and activated histiocytes, leading to uncontrolled phagocytosis of platelets, erythrocytes, lymphocytes and precursor cells. HLH is traditionally considered a rare event, but recent studies have suggested that this condition is underdiagnosed. HLH is a severe inflammatory and aggressive condition, characterized by high fever, hepatosplenomegaly, lymphadenopathy and cytopenia. Central nervous system involvement, cutaneous manifestations, severe coagulation disturbances and multiple organ dysfunctions occur less frequently. Laboratory findings of high levels of ferritin, hypertriglyceridemia and hypofibrinogenemia can also be present, supporting the diagnosis.

This syndrome is classified as familial or acquired, the latter being more frequent. The familial form is related to genetic immune defects, while the acquired form is associated with diverse conditions, such as infections, malignancies and rheumatic diseases. We report a case of HLH associated with cytomegalovirus infection in a patient with acquired-immunodeficiency syndrome and Burkitt’s lymphoma.

Key-Words: Hemophagocytic syndrome, histiocytes, immunosuppression, sepsis, organ dysfunction.

Case Report

A 34 year-old man with AIDS who was on antiretroviral therapy ( stavudine, lamivudine, atazanavir and ritonavir) was admitted to the hospital with a cervical mass. An excisional biopsy yielded a diagnosis of Burkitt’s lymphoma. The CD4 count was 386 cells/mm\(^3\) and the viral load was undetectable. One week later, chemotherapy was initiated, with a significant reduction in the size of the cervical mass, and the patient was discharged. Three days later, he was admitted to the hospital with febrile neutropenia (39 neutrophils/mm\(^3\) ), thrombocytopenia (39,000/mm\(^3\)), and mucositis. Despite the administration of broad-spectrum antibiotics, the patient developed acute respiratory failure with a need for mechanical ventilation; he was transferred to the intensive care unit (ICU). Pneumocystis jiroveci pneumonia was diagnosed and treated with trimethoprim-sulfamethoxazole, prednisone and granulocyte colony-stimulating factor (GCS-F) was started. The patient’s clinical condition improved and his blood cell count normalized. During the ICU stay, the patient developed ventilator-associated pneumonia. Despite appropriate antimicrobial therapy, he evolved with persistent fever and pancytopenia (leukocyte count: 800/mm\(^3\); platelet count: 14,000/mm\(^3\); and hemoglobin level: 6.5 g/dL). GCS-F was administered again, with no response. Several diagnostic procedures to determine the etiology of the pancytopenia were performed. Cytomegalovirus antigenemia was positive, and the bone marrow aspirate revealed hypoplasia, with histiocyte infiltration, as well as phagocytosis of blood cells, consistent with HLH (Figure 1). Based on these findings, therapy was initiated with ganciclovir, resulting in resolution of fever and recovery of the blood cell count. Ten days later, the patient was discharged from the ICU. In the hospital room, cell blood counts returned to normal, CMV antigenemia was negative and the patient was discharged from the hospital.

Discussion

The hemophagocytic syndrome was first described in 1979 in immunosuppressed patients with viral infections [1]. EBV is the most common etiology, while CMV is associated with 30% to 40% of all virus-associated HLH cases [2]. This
syndrome is a result of damage caused by cytotoxic activity of natural killer (NK) cells and T cells; thus the ineffective immune system is constantly stimulated to generate high levels of cytokines, such as tumor necrosis factor α (TNF-α) and interferon γ (IFN-γ), which stimulate the defense cells. High levels of cytokines are responsible for the clinical picture [2,3]. HLH is an uncommon and frequently undiagnosed event, recently described in critically-ill patients [2-4]; it remains a diagnostic challenge, as the clinical presentation of this condition mimics sepsis, a frequent syndrome in ICU patients [5]. Laboratory tests for diagnosis, such as serum triglyceride, ferritin and fibrinogen levels, are frequently not performed in the ICU, and they lack specificity, as they may be significantly altered as a result of critical illness. Moreover, blood transfusions may mask pancytopenia. Our patient had at least four conditions associated with the development of HLH (HIV infection, Burkitt’s lymphoma, bacterial sepsis and CMV infection). The association of HLH and HIV has been described; however, it occurs less frequently than previously expected [6]. HIV-positive patients with lymphoma are at high risk for HLH [1,3]. The association between bacterial infection and HLH is poorly documented, and most case reports indicate that it is due to intracellular bacteria (e.g. *Mycobacteria, Legionella sp.*) [1]. Finally, although the other factors could cause HLH, we believe that CMV was the main etiologic agent involved. The prompt response to ganciclovir allowed clinical and hematological improvement and ICU discharge. This case emphasizes the importance of bone marrow examination and extensive investigation for opportunistic infections in immunocompromised patients presenting with febrile pancytopenia. In most cases, treatment of the underlying condition promotes complete remission of the clinical picture.

Conclusions

We report a case of HLH associated with cytomegalovirus infection in a patient with AIDS and Burkitt’s lymphoma. Although HLH is an uncommon event described in critical patients, it is possibly underdiagnosed. It has been associated with a series of infectious agents, as well as genetic, autoimmune and neoplastic diseases. Awareness concerning its clinical presentation and a high level of clinical suspicion are essential, especially in immunosuppressed patients with fever, hepatosplenomegaly, and pancytopenia that do not respond to GCS-F and antibiotics. Treatment of the underlying condition often leads to clinical improvement.

References