Fatal Sulfasalazine-Induced Eosinophilic Myocarditis in a Patient with Periodic Fever Syndrome

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Introduction

Drug-induced hypersensitivity syndrome (DIHS), also known as drug rash with eosinophilia and systemic symptoms – DRESS – syndrome, is a rare (1/1,000–10,000 exposures) but potentially fatal (10%) systemic reaction, which occurs 1–8 weeks after exposure to anticonvulsants, sulfasalazine, allopurinol or antiretrovirals, etc. [1]. DIHS typically manifests with a rash and hepatic dysfunction, but may include nephritis, pneumonitis and rarely hemophagocytosis and eosinophilic myocarditis (EM) [1]. Hyper-IgD syndrome, one of the genetically defined periodic fever syndromes (PFS), also known as autoinflammatory syndromes, is characterized by reduced activity of mevalonate kinase and lifelong recurrent febrile attacks of a noninfectious origin [2]. It is not known how exactly a reduced activity of mevalonate kinase leads to PFS, but it has been reported that proinflammatory cytokine production in patients with hyper-IgD syndrome is strongly enhanced [3]. We hereby report a case of DIHS in a patient with PFS.
Case Report

A 28-year-old man presented to the Emergency Department of the Clinical Center of Serbia, Belgrade, with fever of 39.9 °C, facial/neck edema, erythematous rash, cervical lymphadenopathy, malaise and shortness of breath that had commenced 24 h previously. He was tachycardic with a gallop rhythm, had dull heart sounds, and was tachypneic (20/min) and hypotensive (60/10 mm Hg). No murmurs or rubs were present. An electrocardiogram showed sinus tachycardia (150/min) with occasional premature ventricular extrasystoles. The patient's personal history revealed PFS present since childhood. He had had febrile attacks once a month, accompanied with cervical lymphadenopathy, abdominal cramps and arthritis. His febrile attacks had been treated with nonsteroid anti-inflammatory drugs and short prednisone courses. Previous attacks had been associated with high inflammatory markers, anemia and mild hepatosplenomegaly. His heart function had always been normal. He had been treated with sulfasalazine (500 mg t.i.d.) for ankle joint arthritis 2 weeks before this latest admission.

On admission, blood tests revealed leukocytosis 18.4 × 10⁹ (78% neutrophils without eosinophilia and atypical lymphocytes) and the following elevated inflammatory markers: C-reactive protein (19,455 μg/l), creatine kinase (313 U/l), MB (muscle brain) creatine kinase isoenzyme (81 U/l), troponin T (1.6 μg/l), aspartate aminotransferase (287 U/l) and alanine aminotransferase (238 U/l). Metabolic acidosis (HCO₃⁻ 4.6 mmol/l, pH 7.16, pCO₂ 1.7 kPa) was present. Chest X-rays showed cardiac enlargement. Plasma N-terminal pro-B-type natriuretic peptide was also elevated at 215 ng/l (normal is <100 ng/l).

A diagnosis of cardiogenic shock was made and therapy with oxygen, fluids, inotropic agents, methylprednisolone (120 mg i.v.) and antibiotics was initiated. Despite this treatment, the patient died 8 h after admission.

The autopsy revealed EM with prominent interstitial edema and inflammatory infiltrate composed of eosinophils and mononuclear cells (fig. 1a, b). Vascular congestion of alveolar wall capillaries and alveolar damage were present (fig. 1c). Eosinophilic infiltrates were found in the stomach and skeletal muscles, while massive hemophagocytosis was observed in bone marrow and lymph nodes (fig. 1d). Acute heart failure caused by EM followed by pulmonary edema was established as the cause of death. The diagnosis of DIHS was confirmed by the first 5 of the following 6 diagnostic criteria: (i) a maculopapular rash developing 2–6 weeks after starting new therapy, (ii) lymphadenopathy, (iii) fever (>38 °C), (iv) leukocytosis >10 × 10⁹/l (without atypical lymphocytes and eosinophils but with prominent tissue eosinophilia), (v) hepatitis (alanine aminotransferase, ALT >100 U/ml) and (vi) human herpes virus 6 reactivation (not tested) [1]. Genetic testing results, which were received postmortem, showed two heterozygous mutations in the gene for mevalonate kinase in exone 11 (C.1129G>A leading to P.V377I) and exone 6 (C.564.G>A leading to P.W188X) localized on 12q24.11, thus confirming hyper-IgD syndrome as one of the genetically defined hereditary periodic fever syndromes.

Discussion

Our patient with PFS had a fulminant presentation of EM, with the first symptoms occurring 13 days after the drug introduction, with cardiogenic shock and fatal outcome in the next 24 h. EM is an under-recognized manifestation of DIHS that may occur in an acute form or can lead to progressive heart failure [4, 5]. Ampicillin, aro-
matic anticonvulsants and allopurinol are most frequently responsible for cardiac involvement in DIHS [5]. Myocarditis associated with DIHS often occurs one to several months later and even after the offending drug has been discontinued [5]. Myocardial injury is mediated by toxic cationic proteins, oxygen metabolites and lipid mediators produced by eosinophils [5].

Acute necrotizing EM is a rare and severe complication of DIHS that usually presents with acute chest pain, ST-segment elevation and an increase in cardiac enzymes, frequently with rapid deterioration of systolic function and a mortality rate above 50% [4]. Pericarditis and cardiac arrhythmias are also possible presentations [5]. In most cases the diagnosis is established on autopsy.

Necrotizing EM, presenting as acute myocardial infarction, has been described in a patient with ankylosing spondylitis 6 weeks after starting sulfasalazine therapy [6]. Due to the increase of IL-1 family cytokines in hyper-IgD syndrome [3], our patient had the hyperacute course of DIHS.

The relationship between toxic, viral and immune mechanisms in DIHS remains unresolved. A role of viral reactivation (especially of human herpes virus 6 and cytomegalovirus) is well known [1]. In our patient, contribution of viral reactivation could not be excluded, considering the presence of mixed cellular heart infiltrate and hemophagocytosis. The reactivation of Epstein-Barr virus and hemophagocytosis associated with sulfasalazine treatment have both been described [7].

Variability in clinical and laboratory presentation and concomitant use of anti-inflammatory drugs may delay the diagnosis of DIHS in patients with PFS. No blood eosinophilia together with tissue eosinophilia, as found in our patient, has been previously described [4]. Therefore, the absence of blood eosinophilia does not rule out the diagnosis of drug-induced EM. A positive in vitro lymphocyte transformation test, which was not performed in our patient, may be helpful in the diagnosis of sulfasalazine-induced DIHS. After 6 days of culture, the lymphocyte transformation test measures drug-specific proliferation (³H-thymidine incorporation) in comparison with negative and positive controls [8].

The diagnosis of EM is based on clinical criteria including electrocardiography, echocardiography and cardiac enzymes. The diagnosis is confirmed by endomyocardial biopsy [5]. In subacute and chronic EM, intracardiac thrombi, endomyocardial fibrosis and restrictive cardiomyopathy can be found.

A prompt diagnosis, immediate withdrawal of the offending drug and immunosuppressant therapy may decrease mortality in drug-induced acute EM [1]. High-dose systemic corticosteroids (1 g/day for 3 days with slow tapering), intravenous Ig therapy (1 g/kg), mycophenolate mofetil (1.5 g b.i.d.), azathioprine, rituximab and cyclosporine are possible treatment modalities [5].

In conclusion, this was a case of drug-induced EM in a patient with PFS that had atypical presentation, rapid evolution and a poor outcome.

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Disclosure Statement

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