Hemophagocytic Lymphohistiocytosis in Activated PI3K Delta Syndrome: an Illustrative Case Report

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To the Editor,

Hemophagocytic lymphohistiocytosis (HLH) is a life-threatening condition characterized by a dysregulated immune system. HLH is driven by impaired antigen clearance that leads to uncontrolled immune activation [1]. In primary HLH, a genetic mutation impairs the cytotoxic function of natural killer (NK) cells and cytotoxic T-cells, resulting in persistent antigen presentation and dysregulation of histiocyte activation. In secondary HLH (sHLH), unknown or non-genetic triggers result in a state of hyperinflammation. The most common triggers associated with HLH include infections, such as Epstein–Barr virus (EBV), hematological malignancies, and autoimmune diseases.

Activated PI3K delta syndrome (APDS) is a primary immunodeficiency (PID) caused by a gain-of-function mutation in \( \text{PIK3CD} \). As a result of the over-activation of the PI3K/AKT/mTOR pathway, T and B cell senescence is induced [2–4]. APDS is characterized by recurrent sinopulmonary infections, non-malignant lymphoproliferation, (persistent) viral infections, autoimmunity, and increased risk of lymphomas [5].

In APDS patients, HLH might be expected. However, it is only scarcely reported. Here, we describe a young adult with APDS who developed persistent EBV-related HLH. We subsequently discuss the association between HLH and APDS.

Case Report

A 19-year-old man of Indian descent was referred for the evaluation of chronic active EBV disease. The patient’s medical history revealed congenital stenosis of the left bronchus, recurrent sinopulmonary infections, and autism spectrum disorder with an IQ of 80–86. EBV seroconversion was identified at 7 years of age when the patient presented idiopathic abdominal lymphadenopathy. Furthermore, a previous laboratory examination revealed IgA deficiency and low IgG2 and IgG4 levels without an overarching diagnosis. Four months before presentation, the patient developed fatigue, fever, night sweats, hepatosplenomegaly, anemia, hepatitis, weight loss of 10 kg, and lymphadenopathy (Fig. 1). Serum EBV copy numbers were repeatedly > 1500/mL. Histology of the three lymph nodes did not reveal any evidence of malignant lymphoma, but rather a markedly positive EBV-encoded RNA (EBER). Chronically active EBV was diagnosed, and rituximab therapy of 500 mg weekly during 4 weeks was initiated. In the workup of PID, whole exome sequencing (confirmed by Sanger sequencing) revealed a heterozygous c.3074A>C, p.Glu1025Gly mutation in the \( \text{PIK3CD} \) gene (NM_005026.3), which is associated with APDS (Fig. 2) [6]. Neither parent carried this mutation, suggesting a de novo origin.

A stable condition without any additional immunosuppressive drugs was achieved for 3 months after which he developed high spiking fever, pneumonia, progressive lymphadenopathy, severe unexplained cardiomyopathy with progressive anemia, thrombocytopenia, and hyperferritinemia (Table 1). Malignant lymphoma was excluded and hemophagocytosis was detected in the bone marrow (Fig. 3A, B). The patient therefore fulfilled six out of eight diagnostic HLH-04 criteria (Table 2) [7]. His clinical status quickly deteriorated, and he developed acute respiratory distress syndrome and multiple organ failure. Salvage therapy was initiated with high-dose steroids, sirolimus, and intravenous immunoglobulin. PI3K delta inhibitors were withheld.

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due to heart and respiratory failure. Etoposide was withheld due to leucopenia and severe sepsis with systemic infection (hospital acquired pneumonia with H. influenza, not responding to treatment). Within 1 month, the patient died from cardiac failure with refractory pulmonary edema.

**HLH in APDS**

APDS is caused by a heterozygous gain-of-function mutation in *PIK3CD*. This mutation is known to activate leucocyte development, pushing immune cells towards differentiation and exhaustion by raising CD8+CD57+ and CD8+ effector memory cells [2, 3]. This is indicative of a defective immune response that is incapable of effective elimination of pathogens. Persisting antigen presentation can then evoke HLH. Especially in the case of persistent EBV, a virus that is already able to trigger HLH, APDS patients are theoretically more prone to develop hyperinflammation and thus HLH [8]. Although chronically active EBV and EBV-associated lymphomas are common in APDS, HLH has only been reported in two APDS patients, who are summarized in Table 3 [9, 10]. Both patients were children at the time of APDS diagnosis, which suggests that genetic diagnosis is more often considered in pediatric HLH. Because a high index of suspicion is necessary to identify both HLH and APDS, underdiagnosis is likely.

**Table 1** Laboratory investigations

| Laboratory investigations | At admission | At time of clinical deterioration | Reference |
|---------------------------|--------------|---------------------------------|-----------|
| Hemoglobin (g/dL)         | 6.28         | 5.8                             | 13.9–16.9 |
| Leukocytes (x10^9/L)      | 4.4          | 5.0                             | 3.5–10    |
| Neutrophils (x10^9/L)     | 2.5          | 3.84                            | 1.4–8.0   |
| Platelets (x10^9/L)       | 177          | 24                              | 150–370   |
| CRP (mg/L)                | 190          | 321                             | <10       |
| ASAT (U/L)                | 40           | 312                             | <35       |
| ALAT (U/L)                | 19           | 46                              | <45       |
| Lactate dehydrogenase (U/L) | 263        | 798                             | <248      |
| Gamma-GT (U/L)            | 230          | 391                             | <55       |
| Alkaline phosphatase (U/L) | 918         | 1301                            | <115      |
| Total bilirubin (µmol/L)  | 26           | 66                              | <17       |
| NT-pro-BNP (pmol/L)       | 1028         |                                 | <15       |
| IgA (g/L)                 | 0.76         |                                 | 0.76–3.9  |
| IgG (g/L)                 | 5.70         |                                 | 7.0–16.0  |
| IgM (g/L)                 | 1.24         |                                 | 0.45–2.30 |
| IgG1 (g/L)                | 4.67         |                                 | 4.9–11.4  |
| IgG2 (g/L)                | 0.62         |                                 | 1.5–6.4   |
| IgG3 (g/L)                | 0.32         |                                 | 0.2–1.1   |
| IgG4 (g/L)                | <0.08        |                                 | 0.08–1.4  |
| Ferritin (µg/L)           | 1985         | 10,364                          | 30–240    |
| sIL2R (pg/mL)             | >110,000     | >110,000                        | <2500     |
| Triglycerides (mmol/L)    | 2.41         | 6.58                            | <2        |
| EBV DNA (IU/mL)           | 2.54E4/P     | 1.17E6/P                        |           |

Abbreviations: CRP C-reactive protein, ASAT aspartate aminotransferase, ALAT alanine aminotransferase, Gamma-GT gamma-glutamyltransferase, NT-pro-BNP N-terminal prohormone of brain natriuretic peptide, Ig immunoglobulin, sIL2R soluble interleukin-2 receptor.

**Fig. 1** High glucose metabolism in 18F-fluorodeoxyglucose positron emission tomography positive lymph nodes.

**Fig. 2** Heterozygous c.3074A>C, p.Glu1025Gly mutation in *PIK3CD* identified in the patient. Linear representation of P110δ(PIK3CD) protein domains. The mutation identified in the patient is shown in red. Abbreviations: ABD, adaptor-binding domain; RBD, Ras-binding domain; C2, putative membrane-binding domain.
**Fig. 3** Bone marrow biopsies from the patient with mutated PIK3CD. A Hematoxylin and eosin (H&E) staining shows the hemophagocytosis of erythroid cells by a macrophage. B May–Grünewald–Giemsa staining shows hemophagocytosis of a polynuclear granulocyte by a macrophage.

**Table 2** HLH criteria

| HLH-04 criteria [7] | Patient finding |
|---------------------|-----------------|
| Fever               | Present         |
| Splenomegaly        | Present         |
| Cytopenias (affecting ≥2 of 3 lineages in the peripheral blood) | Present |
| Hemoglobin < 90 g/L | 58 g/L          |
| Platelets < 100 × 10⁹/L | 24 × 10⁹/L |
| Neutrophils < 1.0 × 10⁹/L | 3.84 × 10⁹/L |
| Hypertriglyceridemia and/or hypofibrinogenaemia | Not present |
| Fasting triglycerides ≥ 3.0 mmol/L | 6.58 mmol/L |
| Fibrinogen ≤ 1.5 g/L | Not done |
| Hemophagocytosis in bone marrow or spleen or lymph nodes | Present |
| Low or absent NK-cell activity | Not done |
| Ferritin ≥ 500 µg/L | Present (10,364 µg/L) |
| Soluble IL-2 receptor ≥ 2400 U/mL | Present (> 110,000 pg/mL) |

**Table 3** Previously published cases on HLH in APDS patients

| Characteristic                      | Patient 1 [9] | Patient 2 [10] |
|-------------------------------------|---------------|---------------|
| Sex                                 | Male          | Male          |
| Age at the time of APDS diagnosis   | 12 years      | 2 years       |
| Age at the time of HLH onset        | 12 years      | 5 months      |
| Trigger of HLH                      | Idiopathic    | Hodgkin lymphoma |
| Presentation                        | Diffuse lymphadenopathy, hepatosplenomegaly, arthritis, rash | Diffuse lymphadenopathy, hepatosplenomegaly, autoimmune hemolytic anemia |
| PIK3D mutation                      | c.323C > G p.R108L | c.3061G > A p.E1021K |
| Concurrent mutation                 | None noted    | PRF1 p.L236F  |
| Functional NK and T-cell test       | Normal        | Normal function. Lower (but not deficient) perforin levels in NK cells |
| White blood cell count (× 10³/mm³) | 6.14 (4.0–10.8) | 9.08           |
| Neutrophils (× 10³/mm³)             | 4.58 (1.5–8.0)  | 2.39          |
| Lymphocytes (× 10³/mm³)             | 1.21 (0.9–4.0)  | 4.89          |
| IgG (mg/dL)                         | 2119 (700–1600) | 426 (270–792) |
| IgA (mg/dL)                         | 207 (70–400)   | 47 (5.8–58)   |
| IgM (mg/dL)                         | 127 (40–230)   | 258 (18.4–145) |
This case emphasizes the importance of genetic testing for PID (including HLH genes) in patients with the first episode of HLH, regardless of age. An increasing number of genetic mutations in older patients with HLH have been reported [11]. In particular, a medical history of recurrent infections, autoimmunity, developmental delay, and EBV-related HLH in young adults should indicate PID. Correct genetic diagnosis enables personalized treatment and thereby improving survival. Appropriate treatment of APDS, such as hematopoietic stem cell transplantation or mTOR inhibitors, especially the new promising PI3K inhibitor leniolisib. By doing so, the patient is less susceptible to developing HLH. If HLH arises, an additional trigger (e.g., infectious or malignant) should be sought thoroughly and treated rigorously. In severe cases, HLH-directed therapy could be considered to halt the hyperinflammation. This therapy should consist of corticosteroids, with the addition of etoposide in life threatening cases [12]. This case illustrates that earlier diagnosis of APDS could have resulted in more adequate monitoring and targeted treatment before the occurrence of multi-organ failure.

Data Availability  Data transparency.

Code Availability  Not applicable.

Declarations

Ethics Approval  To our opinion, this research does not necessitate ethic approval. It concerns a retrospective study in which no data could be traced back to the presented patient. The patient is not subjected to any intervention and no behavior is imposed.

Consent to Participate  Not applicable.

Consent for Publication  Not applicable.

Conflict of Interest  The authors declare no competing interests.

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