Henoch-Schönlein purpura associated with primary active Epstein-Barr virus infection: a case report

Burcu Karakayali¹, Sila Yılmaz¹, Deniz Çakır¹, Pembe Gül Günes², Sirin Güven¹, Ismail Islek¹,‡

¹University of Health Sciences, Umrahie Research and Training Hospital, Department of Pediatrics, Istanbul, Turkey, ²Haydarpasa Numune Research and Training Hospital, Department of Pathology, Istanbul, Turkey

‡Corresponding author: Ismail Islek, University of Health Sciences, Umrahie Training and Research Hospital, Department of Pediatrics, Istanbul, Turkey

Key words: Henoch-Schönlein purpura, Epstein-Barr virus, children

Received: 05/08/2016 - Accepted: 28/04/2017 - Published: 11/05/2017

Abstract

Henoch-Schönlein purpura (HSP) is the most common form of childhood vasculitis. Various viral and bacterial infections, drugs, vaccines, food allergy and even insect bites have been considered as triggering factors in pathogenesis of HSP. Epstein-Barr virus (EBV) infection, which is associated with HSP, have been rarely reported. Herein we present HSP patient possibly caused by EBV infection. A 8-year old boy was admitted to our department with fever, rashes on legs and arms and intermittent mild abdominal pain. Multiple purpuric rashes were on his extremities, abdomen and buttock. Laboratory investigations revealed that monospot test was positive, EBV serology tests; Anti-EA-D Ig G: 3+, Anti-VCA gp125 Ig G: 3+, Anti-VCA p19 Ig M: 2+, Anti EBNA-1 Ig M: negative, Anti EBNA-1 Ig G: negative. The patient was interpreted as the primary active acute EBV infection. A skin biopsy showed leucocytoclastic vasculitis. The other viral and bacterial investigations were negative. The patient was diagnosed as HSP vasculitis according to EULAR criteria and treated with intravenous hydration and ibuprofen. He was discharged after 15 days with normal laboratory findings and physical examination. We think that EBV infection may be stimulant factor for autoimmune reactions and may cause HSP vasculitis. Hence, it may be useful to investigate the EBV infection in etiology of HSP cases.

Pan African Medical Journal. 2017; 27:29 doi:10.11604/pamj.2017.27.29.10481

This article is available online at: http://www.panafrican-med-journal.com/content/article/27/29/full/

© Burcu Karakayali et al. The Pan African Medical Journal - ISSN 1937-8688. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.
Introduction

Henoch-Schönlein purpura (HSP), which is the most common form of childhood vasculitis, affects mostly skin, joints, gastrointestinal system (GIS), kidneys, and in addition more rarely can have effect on central nervous system (CNS), heart and scrotum. Its pathogenesis and causal factors have not been accurately identified yet [1]. Viral and bacterial infections, drugs, vaccines, food allergy and even insect bites have been considered in the etiology of HSP. Streptococcus, vaccines, viral infections (varicella, measles, rubella, hepatitis A, B), tuberculosis, mycoplasma, Bartonella, helicobacter pylori are stated as the triggering factors in the literature [1-4]. Epstein-Barr virus (EBV) infection associated with HSP cases have been reported rarely [5-7]. Herewith we report the case of an 8-year old boy with primary active Epstein Barr Virus infection triggering HSP.

Patient and observation

A 8-year old boy, who was previously well, presented to our emergency department with fever, fatigue, rashes on his both legs and arms and intermittent mild abdominal pain. On physical examination, the vital signs were as follows: blood pressure 110/60 mmHg, respiratory rate, 24/min; body temperature 38.5°C; heart rate, 110/min. Left submandibular lymph node was 2x1 cm, tender and soft. His lung auscultation and heart sounds were normal without murmur. His liver is palpable 1 cm below right costal margin. Splenomegaly was also found; confirmed by abdominal ultrasonography. Multiple purpuric rashes were on his extremities, abdomen and buttock Figure 1. On admission, laboratory investigations revealed white blood cell (WBC) count of 30,000/mm³, 75% lymphocytes, hemoglobin of 11 g/dl and a platelet count of 357,000/mm³. Biochemical studies revealed serum creatinine of 0.6 mg/dl, blood urea nitrogen of 19 mg/dl, protein 7.9 g/dl, serum albumin of 3.9 g/dl, aspartate amino transferase (AST) 181 IU/L (<52), alanine aminotransferase (ALT) 122 IU/L (<52) and gamma-glutamyltransferase (GGT) 32 g/dL. Erythrocyte sedimentation rate and C-reactive protein level were 48 mm/h and 4.6 mg/dl (0-0.5 mg/dl) respectively. Blood coagulation tests, C3, C4, ANA, other immunoglobulin levels, anti-dsDNA, rheumatoid factor (RF), c-ANCA, p-ANCA and anti streptolysin-O (ASO) were all negative. The investigation of Hepatitis B and C serologies, HIV, VDRL-RPR, VZV, CMV, toxoplasma, rubella Ig M and parvovirus antibody were all negative. Monospot test was positive. The EBV serology tests were revealed as follows Anti-EA-D Ig G: 3+, Anti-VCA gp125 Ig G: 3+, Anti-VCA p19 Ig M: 2+, Anti EBNA-1 Ig G: negative, Anti EBNA-1 Ig M: negative, Anti EBNA-1 Ig G: negative. In light of this findings, the patient was interpreted as the primary active EBV infection. In addition a skin biopsy from left malleol showed leucocytoclastic vasculitis Figure 2. Chest X-ray and an electrocardiogram were normal. The patient was diagnosed as HSP vasculitis according to criteria set by European League against Rheumatism (EULAR) [8]. The patient was treated with intravenous hydration and ibuprofen. The clinical finding improved gradually within 15 days.

Discussion

We herein report the patient with HSP associated with primary active EBV infection. Our patient met clinically HSP criteria and leucocytoclastic vasculitis was demonstrated by skin biopsy. Viral and bacterial investigation showed primary active acute EBV infection and excluded the other infections. A variety of patients involving HSP and infection have been described following bacterial and viral infections [1-4]. EBV is also reported in association with various vasculitis forms such as Kawasaki disease, leucocytoclastic vasculitis, granulomatous vasculitis, systemic lupus erythematosus and ANCA associated vasculitis [9]. Pender stated autoimmune diseases are mainly caused by the infection of autoreactive B cells by EBV [10]. He suggested that autoimmune disease, as a result of EBV infection, is due interaction between B cells, already infected by EBV, and autoreactive T cells. The reason behind the damage of target organ in autoimmune disease is autoreactive T cells proliferation and stimulation to produce cytokines. In our patient, EBV antibodies results indicate primary active EBV infection [10]. Three cases, in which chronic EBV infection lead to a case similar to HSP, have been reported in the literature [4]. In addition, another three cases have been reported for acute EBV infection associated with HSP nephritis [6, 7]. Another case report presents a child patient who has acute EBV infection with immune-complex glomerulonephritis resulting with hemodialysis need and resistant hypertension [5]. Three adult cases shows that EBV related lymph node lesions are similar to autoimmune disease related lymph node reactions [11].

Conclusion

Summary, we think that EBV infection may be stimulant factor for autoimmune reactions and may cause HSP vasculitis. Hence, it may be uselful to investigate the EBV infection in etiology of HSP cases.

Competing interests

The authors declare no competing interests.

Authors’ contributions

Burcu Karakayali: followed the patient, collected the data of patient Sila Yilmaz: followed the patient, collected the data of patient, made english editing Deniz Çakır: made diagnosis in point of infectious disease Pembe Gul Gunes: made diagnosis with pathologic investigation, Sirin Guven: contribute poster preparation (This case was presented at Pediatric Rheumatology National Congress in Antalya, 2016, April) and helped to draft the manuscript Ismail Islek: conceived of the case report, followed the patient and wrote the manuscript.

Figures

Figure 1: Multipl purpura on the lower extremity
Figure 2: Vascular damage characterized by endothelial swelling and necrosis, fibrinoid change in vessel walls (HEX200)

References

1. Petty RE, Laxer RM, Lindsley CB, Wedderburn LR. Textbook of Pediatric Rheumatology - 7th edition. Elsevier, 2016. Google Scholar
2. Islek I, Kalayci AG, Gok F, Muslu A. Henoch-Schönlein purpura associated with hepatitis A infection. Pediatr Int. 2003;45(1):114-116. PubMed | Google Scholar

3. Islek I, Muslu A, Totan M, Gok F, Sanic A. Henoch-Schönlein purpura and pulmonary tuberculosis. Pediatr Int. 2002;44(5):545-546. PubMed | Google Scholar

4. Guissa VR, Aragao PA, Marques HH, Jacob CM, Silva CA. Chronic active Epstein-Barr virus infection mimicking Henoch-Schönlein purpura. Acta Reumatol Port. 2010;35(5):513-517. PubMed | Google Scholar

5. Lande MB, Mowry JA, Houghton DC, White CR Jr, Borzy MS. Immune complex disease associated with Epstein-Barr virus infectious mononucleosis. Pediatr Nephrol. 1998;12(8):651-653. PubMed | Google Scholar

6. Grech V, Vella C. Henoch-Schoenlein purpura with nephritis in two siblings following infectious mononucleosis. Ann Trop Paediatr. 2002;22(3):297-298. PubMed | Google Scholar

7. Kim CJ, Woo YJ, Kook H, et al. Henoch-Schönlein purpura nephritis associated with Epstein-Barr virus infection in twins. Pediatr Nephrol. 2004;19(2):247-248. PubMed | Google Scholar

8. Ozen S, Pistorio A, Lusan SM, et al. EULAR/PRINTO/PRES criteria for Henoch-Schönlein purpura, childhood polyarteritis nodosa, childhood Wegener granulomatosis and childhood Takayasu arteritis: Ankara 2008, Part II: Final classification criteria. Ann Rheum Dis. 2010; 69(5):798-806. PubMed | Google Scholar

9. Yamaguchi M, Yoshioka T, Yamakawa T, Maeda M, Shimizu H, Fujita Y, Maruyama S, Ito Y, Matsuo S. Anti-neutrophil cytoplasmic antibody-associated vasculitis associated with infectious mononucleosis due to primary Epstein-Barr virus infection: report of three cases. Clin Kidney J. 2014;7(1):45-48. PubMed | Google Scholar

10. Pender MP. Infection of autoreactive B lymphocytes with EBV, causing chronic autoimmune diseases. Trends Immunol. 2003;24(11):584-588. PubMed | Google Scholar

11. Kojima M, Yamane Y, Itoh H, et al. Epstein-Barr virus-related lymph node lesion resembling autoimmune disease-like clinicopathological findings in elderly patients, report of three cases. APMIS. 2003;111(12):1083-1088. PubMed | Google Scholar

Figure 1: Multipl purpura on the lower extremity
Figure 2: Vascular damage characterized by endothelial swelling and necrosis, fibrinoid change in vessel walls (HEX200)