Successful Treatment of Corticosteroid with Antiviral Therapy for a Neonatal Liver Failure with Disseminated Herpes Simplex Virus Infection

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Abstract

Background  Herpes simplex virus (HSV) infection carries one of the poorest outcomes of neonatal liver failure (NLF). Neonates with disseminated HSV infection can develop hemophagocytic lymphohistiocytosis (HLH), and occasionally need orthotopic liver transplantation. Early interventions may be critical for the cure of NLF.

Case Report  We describe herewith a 6-day-old neonate with fulminant hepatic failure due to disseminated HSV-1 infection, who successfully responded to high-dose corticosteroid therapy 72 hours after the onset of disease. Preceding acyclovir, gamma globulin, and exchange blood transfusion therapies failed to control the disease. Methylprednisolone pulse therapy led to a drastic improvement of liver function and cytokine storms, and prevented the disease progression to HLH. Sustained levels of plasma and cerebrospinal fluid HSV DNA declined after prolonged acyclovir therapy. Bilateral lesions of the periventricular white matter areas, assessed by magnetic resonance imaging, disappeared at 3 months of age. The infant showed normal growth and development at 4 years of age.

Conclusion  Early anti-hypercytokinemia therapy using corticosteroid, and prolonged antiviral therapy might only provide the transplantation-free cure of NLF with HSV dissemination.

Keywords  ► fulminant liver failure  ► newborn  ► herpes simplex virus  ► acyclovir  ► methylprednisolone pulse therapy

Clinical Perspective  Early anti-hypercytokinemia therapy using corticosteroid and prolonged antiviral therapy might only provide the transplantation-free cure of neonatal liver failure with HSV dissemination.

Neonatal liver failure (NLF), occurring as fulminant hepatic failure within the first 28 days of birth, is rare, but has a high mortality.1 It arises from various etiologies including
Neonatal hemochromatosis, inborn errors of metabolism, infections, and hemophagocytic lymphohistiocytosis (HLH). Herpes simplex virus (HSV) infection is one of the major causes of acute liver failure\(^2\) and acquired HLH in early infancy.\(^3\) HSV-1 accounts for more than 60% of herpes meningitis cases in the newborn, which is mostly associated with virus dissemination. The pathophysiology of neonatal HSV infection is explained by systemic inflammatory response syndrome (SIRS),\(^5\) often progress to liver and respiratory failures,\(^6\) and the HLH. The survival rate of NLF patients was reportedly lower than that of the other types of NLF patients even after liver transplantation.\(^1\) Poor neurodevelopmental outcomes occur in approximately 20% of survivors of disseminated HSV disease, and 70% of those with central nervous system (CNS) disease. Prolonged acyclovir therapy is the standard for neonatal HSV disease.\(^7\) However, the effective measures to control HSV-driven NLF and SIRS remains elusive. Corticosteroid therapy is not the rule of NLF management,\(^1\) particularly in cases with the virus dissemination.

We herein present the case of a 6-day-old neonate with NLF and hypercytokinemia because of disseminated HSV infection. This condition was successfully controlled after methylprednisolone pulse therapy started within 72 hours of the disease onset. Early corticosteroid therapy and prolonged antiviral therapy resulted in a significant reduction of serum cytokine levels and the quantity of viral DNA, respectively. Anti-hypercytokinemia therapy in NLF was then discussed.

**Case Report**

A 2,746-g male neonate was born to a healthy mother by vaginal delivery at 39 weeks of gestation, without asphyxia. The infant was transferred to the neonatal intensive care unit 6 days after birth, for fever (38.2°C), poor feeding, apnea, and lethargy without vesicular skin lesions. On admission, mild dyspnea needed respiratory support by nasal continuous positive airway pressure. Computed tomography indicated brain edema. Results of hematological examinations were as follows: leukocytes, 5.15 \(\times\) 10\(^9\)/L; hemoglobin, 14.3 g/dL; platelets, 148 \(\times\) 10\(^9\)/L; C-reactive protein, 2.99 mg/dL; alanine aminotransferase (ALT), 455 IU/L (rr, 5–43 IU/L); aspartate aminotransferase (AST), 2,220 IU/L (rr, 12–34 IU/L); lactate dehydrogenase (LDH), 7,510 IU/dL (rr, 115–217 IU/dL); direct bilirubin, 1.3 mg/dL (rr, 0.1–0.3 mg/dL); cholesterol, 78 mg/dL (rr, 130–220 mg/dL); triglyceride, 33 mg/dL (rr, 46–130 mg/dL); ferritin, 141,066 ng/mL (rr, 25.0–280.0 ng/mL); prothrombin time (PT), 15.8 seconds; PT–international normalized ratio (PT–INR), 1.79; fibrinogen, 209 mg/dL (rr, 200–310 mg/dL); and D-dimer, 33.6 mg/L (rr, 0–1 mg/L). Cerebrospinal fluid (CSF) examinations revealed leukocytes, 2 cells/µL (rr, < 5 cells/µL); glucose, 52 mg/dL (rr, 60–80 mg/dL); and protein, 64 mg/dL (rr, 15–45 mg/dL). Under the tentative diagnosis of sepsis and hepatitis, antibiotics and acyclovir were started (60 mg/kg/d). Laboratory examinations performed 8 hours later indicated progressive liver damage with disseminated intravascular coagulation; ALT, 884 IU/L; AST, 4,459 IU/L; LDH, 13,746 IU/dL; platelets, 104 \(\times\) 10\(^9\)/L; PT, 22.4 seconds; PT–INR, 3.11; fibrinogen, 114 mg/dL; and D-dimer, 37.4 mg/L. Infusions of fresh frozen plasma, and whole exchange blood transfusions were instituted. High copy numbers of HSV-1 DNA, but not, the other types of herpes viruses DNA on admission (serum, 1.8 \(\times\) 10\(^7\) copies/mL; CSF, 5.7 \(\times\) 10\(^5\) copies/mL, assessed by real-time polymerase chain reaction\(^8\)) confirmed the diagnosis of NLF because of disseminated HSV-1 infection. Repeated exchange blood transfusions failed to control liver function and coagulopathy (Fig. 1A). Additional high-dose immunoglobulin therapy (2 g/kg/d \(\times\) 2 d) on 8 and 9 days after birth was ineffective. Then, methylprednisolone therapy (30 mg/kg/d \(\times\) 3 d) led to drastic improvement of coagulopathy (PT–INR), declining transaminase levels, and defervescence. Transaminase levels peaked on the 2nd day of treatment (AST: 5,394 IU/L; ALT: 980 IU/L), and no any organ failure developed. The fluid-attenuated inversion recovery (FLAIR) magnetic resonance imaging (MRI) 15 days after birth showed multiple high-signal lesions in the white matter (Fig. 2A). Acyclovir therapy was continued until 43 days after birth. The copy number of CSF HSV DNA decreased to 2.9 \(\times\) 10\(^2\) copy/mL 30 days after birth. The serum copy numbers steadily decreased and became undetectable 49 days after birth (Fig. 1B). No HSV infection recurred after the hospital discharge. Funduscopy, automated auditory brain stem response, and electroencephalogram findings were all normal. FLAIR MRI at 3 months (Fig. 2B) and 2 years and 9 months of age revealed no abnormalities. At the age of 4 years, he is alive and well having 90 score of

![Fig. 1](image) Sequence changes of the liver function and serum cytokine levels during the initial treatment course (A), and those of HSV DNA copy number in serum (•) and cerebrospinal fluid (□) during the overall treatment course (B). ALT, alanine aminotransferase; AST, aspartate aminotransferase; IVIG, intravenous high-dose immunoglobulin therapy; HSV: herpes simplex virus; IFN, interferon; IL, interleukin; PT–INR, prothrombin time–international normalized ratio.
intelligence quotient assessed by the Tanaka–Binet intelligence test.

Serum cytokine levels during the acute phase of HSV–NLF (6–12 days after birth) were retrospectively measured by flow cytometry using a cytometric bead array kit (BD Pharmingen, San Diego, CA) according to the manufacturer’s instructions. Seven days after birth, serum levels of interferon (IFN)-γ, interleukin (IL)-6, and IL-10 were remarkably elevated (Fig. 1B). On the contrary, CSF cytokine levels on admission were all undetectable. Serum levels of IFN-γ and IL-6, but not IL-10 showed the nadir (10 days after birth) after methylprednisolone therapy, in concert with the full recovery of PT–INR.

Discussion

Early determination of the causes of NLF is challenging. The survival of affected infants depends on the management according to the progression of metabolic diseases, neonatal hemochromatosis, and primary/secondary HLH. Liver transplantation and hematopoietic stem cell transplantation are the curative treatment at present for neonatal hemochromatosis and familial HLH, respectively. On the contrary, the treatment strategy of HSV–NLF and its-associated HLH has not been established. In this patient, liver dysfunction and CNS disease prompted us to start the acyclovir. Higher copy number of HSV-1 DNA in the serum than seen in CSF indicated disseminated, rather than isolated CNS, infection of the virus. At that time point, the patient fulfilled the diagnostic criteria of HLH but not HLH in the newborn. Corticosteroid therapy is not recommended for the treatment of fulminant hepatic failure. Etoposide therapy does not raise the survival rate of neonatal HSV-associated HLH. Then, methylprednisolone pulse therapy was introduced to the infant for the control of a cytokine storm. Neonatal disseminated infections with HSV-1 or HSV-2 led to SIRS represented by high IL-6 and TNF-α levels, and occasionally to the rapid demise of multiple organ dysfunction syndrome or HLH. Methylprednisolone therapy might be effective for the control of neonatal disseminated HSV infection with HLH. In this reported case, sequential changes in the declining pattern of IL-6, IL-10, and IFN-γ levels were consistent with the results of our observation, although HSV DNA copy number was not monitored. On the contrary, appreciable levels of IL-10 and IFN-γ in the neonates with virus-associated HLH might be associated with a good prognosis. Nagamori et al have recently reported a successfully treated case of neonatal disseminate HSV infection with early anti-inflammatory therapy using prednisolone. On the basis of the kinetics of high-mobility group box protein 1 (HMGB-1) and cytochrome c, they concluded that anti-inflammatory intervention may only be effective if it is undertaken during the early phase of disseminated neonatal HSV infections. Uncontrolled coagulopathy portends an ominous outcome of liver failure. In this line, sequential changes in the PT–INR and cytokine levels after early corticosteroid therapy might represent the improvement of liver damages through the controlled hypercytokinemia.

The neonates with HSV-associated HLH (HSV–HLH) sometimes show a lack of several parameters of diagnostic criteria of HLH. Serum levels of IL-10 and IFN-γ were elevated in the present patient, differentially from non-HSV–HLH in the newborns. The distinct cytokine profiles may explain the clinical expressions and severity of neonatal HLH, according to the causative agents and genetic predispositions of HLH.

The present patient showed the high quantity of HSV DNA but not cytokines in CSF at the diagnosis. Multiple high-signal lesions in the brain MRI images disappeared with the declining viral loads in the CSF and systemic circulation. The declining pattern of the viral loads was much slower than that of circulating cytokines. After the prolonged acyclovir therapy until the attainment of undetectable HSV DNA, the patient showed neither recurrence of disease nor neurological sequelae. Recently, Scoble and Underwood reported a 9-day-old neonate with HSV-2 driven NLF and shock, who presented bilateral periventricular lesions on MRI. The white matter lesions disappeared during the convalescent phase of the disease, similarly in our patient. In this context, the brain lesions during the acute phase of HSV–NLF might represent the focal edema and/or hemorrhage through hypercytokinemia. This observation may emphasize the practical need of early cytokine control under acyclovir therapy for the complete cure of neonatal disseminated HSV infection, to prevent not only the development of HLH/multiple organ dysfunction syndrome but also the impending CNS involvement. Further study should be focused on the targeted therapy of anti-hypercytokinemia in the newborn neonates with HSV dissemination.

Conflicts of Interest

The authors have no conflicts of interest to disclose.

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