CASE REPORT

Case Report: Severe form of hemolytic-uremic syndrome with multiple organ failure in a child: a case report [version 2; peer review: 2 approved]

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Abstract

Introduction: Hemolytic-uremic syndrome (HUS) is a leading cause of acute renal failure in infants and young children. It is traditionally defined as a triad of acute renal failure, hemolytic anemia and thrombocytopenia that occur within a week after prodromal hemorrhagic enterocolitis. Severe cases can also be presented by acute respiratory distress syndrome (ARDS), toxic megacolon with ileus, pancreatitis, central nervous system (CNS) disorders and multiple organ failure (MOF).

Case presentation: A previously healthy 4-year old Caucasian girl developed acute renal failure, thrombocytopenia and hemolytic anemia following a short episode of abdominal pain and bloody diarrhea. By the end of the first week the diagnosis of the typical HUS was established. During the second week the disease progressed into MOF that included ileus, pancreatitis, hepatitis, coma and ARDS, accompanied by hemodynamic instability and extreme leukocytosis. Nonetheless, the girl made a complete recovery after one month of the disease. She was successfully treated in the intensive care unit and significant improvement was noticed after plasmapheresis and continuous veno-venous hemodialysis.

Conclusions: Early start of plasmapheresis and meticulous supportive treatment in the intensive care unit, including renal placement therapy, may be the therapy of choice in severe cases of HUS presented by MOF. Monitoring of prognostic factors is important for early performance of appropriate diagnostic and therapeutical interventions.

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Introduction

Hemolytic-uremic syndrome (HUS) can be classified as typical, usually when provoked by Shiga-toxin (STx) produced by enterohemorrhagic *Escherichia coli* serotype H157:O7, or atypical, when triggered by other microbes' antigens and toxins. The overall incidence of STx-HUS is estimated to be 2.1 per 100,000 people, with a peak incidence in children younger than 5 years (6.1 per 100,000). Approximately two-thirds of patients have extensive leukocytosis as a hallmark of the disease. In general, HUS patients present with abdominal cramps, vomiting and bloody diarrhea that had appeared within 5 days of disease; the highest WBC count was 94×10^9/L. During the 25th day of disease, her medical condition had not improved and she became anuric, accompanied by combined metabolic and respiratory acidosis, pulmonary edema, bilateral pleural and pericardial effusions, ascites and persistent anemia and thrombocytopenia. Continuous veno-venous hemofiltration (CVVHDF) as well as plasmapheresis were indicated. Access was established via an intravascular catheter placed through the right internal jugular vein guided by ultrasound. CVVHDF was performed for 12 days. Characteristics of dialysis included a dialyzer membrane surface area of 0.8 m², solution containing bicarbonate (multiBic 2 mmol/L potassium solution for haemofiltration; Fresenius Medical Care Deutschland GmbH) and a blood flow rate of 3–6 ml/kg/min. The system was anticoagulated with 100 IU/kg of unfractionated heparin each day. An additional blood flow rate for ultrafiltration was 0.5–2 ml/kg/h.

The renal replacement therapy was stopped for 3 hours a day in order to carry out plasmapheresis which was also initiated on the 8th day of disease. This lasted for 10 days, for 2 hours each day and 40 ml/kg of fresh frozen plasma was used per day. During this period continuous analgosedation with 0.15 mg/kg/h midazolam and 0.075 mg/kg/h morphine, as well as invasive mechanical ventilation were maintained. Continuous adrenergic support was stopped after the 6th day of disease. After 12 days of renal replacement therapy, diuresis started to improve while serum creatinine and urea started to decrease. Despite the polyuria during the initial phase of recovery of renal function, because of the still significant fluid retention, we established negative fluid balance by continuous infusion of furosemide. Renal function had completely recovered by the 25th day of disease (Figure 1). We have reason to believe that the blood levels of creatinine and urea at the time that the CVVHDF was started were falsely lower because of hemodilution caused by the body’s extreme retention of extracellular water.

During the course of plasmapheresis a marked recovery of the platelet count and hemoglobin level was noticed, as well as a significant decline of LDH levels and the degree of *in vivo* hemolysis. After termination of plasmapheresis, an additional time was needed for restoration of platelets and normal erythrocytes to optimal levels because of their biological cell cycle. We were restrictive throughout the transfusion of packed red cells, aiming for a target hemoglobin level of 70–80 g/L.

The most striking laboratory finding during the first two weeks of disease was extreme leukocytosis which persisted during the second week of disease; the highest WBC count was 94×10^9/L. As no other signs of systemic bacterial infection or invasive mycosis were noticed, we believe that leukocytosis was provoked by the systemic inflammatory response to toxemia. We observed the gradual decrease in the WBC count after the initiation of plasmapheresis...
Gradual recovery of renal function. Plasma creatinine and urea levels and urine output during the disease course and applied treatment show significant improvement of renal function by the end of CVVHD and plasmapheresis and normal function is restored in the 4th week of disease without the evidence of permanent damage.

We ceased the analgosedation on the 12th day of disease in order to evaluate the neurological status of the girl. She appeared comatose, estimated as GCS 4 (E2V1M1) on the Glasgow Coma Scale with flaccid tetraparesis, rotatory nystagmus and symmetrical mydriatic pupils with slow response to light stimulation. An MRI scan of the brain recorded on the 12th day of disease revealed areas of raised signal in the pons, bilaterally in lateral parts of the thalamus (Figure 4), as well as in the right external capsule and the left internal capsule. These findings were analogous with lesions already described in literature believed to be the result of microvascular damage. Analgesation was reinstated and afterwards periodically discontinued every few days for approximately 6 hours in order to examine the girl’s clinical neurological status. After the 18th day her level of consciousness and neurological status started to gradually improve and we decided to definitively cease analgosedation 7 days later. No permanent neurological damage was recorded.

We were concerned by severe gastric retention, absent or silent peristalsis and constipation with episodical soft and rare stools during the second week of disease. An MRI of the abdomen performed on the 12th day of disease showed an edematous intestinal wall and distension of the jejunum, ileum and colon ascendens with several
Figure 2. Extreme leukocytosis during the second week of disease indicated a high intensity of systemic inflammation, as the WBC correlated with LDH plasma levels. Plasmapheresis enabled the clearance of proinflammatory factors as the WBC and LDH level normalized during and after this course of treatment. Red lines denote upper reference limit in our laboratory - for plasma LDH 485 IU/L and WBC $10^9$/L.

Figure 3. CT scan of thorax recorded on the 15th day of disease: massive left-sided hemothorax with atelectatic lung (arrow) that eventually required surgical treatment, as well as a careful strategy of protective invasive mechanical ventilation.

Figure 4. MRI scan of brain recorded on the 12th day of disease: bilateral hyperintensity in lateral thalamic regions (arrows).

air fluid levels (Figure 5). Laboratory findings displayed elevated serum transaminases, total and direct bilirubin and amylase as well as hypoalbuminemia. After prodromal gastroenterocolitis, paralytic ileus obviously evolved and was persistent throughout the second and third week of disease. Therefore, all necessary fluid, macronutrients and micronutrients were completely delivered parenterally via a central venous catheter from the start of disease. In combination with already initiated parenteral nutrition, we carefully started enteral feeding by the end of the second week of disease. As no signs of intraabdominal compartment syndrome, perforation or peritonitis were registered, we wanted to reestablish gastrointestinal function conservatively. Therefore, we applied continuous infusion of 6–10 mg/24h of metoclopramid and intermittent rectal suppositories of 2.5–5 mg of bisacodyl for the next two weeks. We also progressively increased the contribution of the enteral input as soon as peristalsis started to improve. Bowel function slowly recovered by the beginning of the fourth week of disease.

As the patient and her family had been on holiday at the time of admission, an in-person follow up was not possible. However, the
authors made contact with the family in 2013 and the parents reported that their daughter was perfectly healthy and without apparent consequences.

Discussion and conclusion
Here we presented the case of a 4-year old girl who developed a severe form of typical HUS which initially presented classically, but rapidly progressed into MOF which severely affected the central nervous system, renal, gastrointestinal and cardiovascular function, and induced ARDS and hematological disorders.

Diarrhoea-associated HUS is the most severe clinical manifestation of infection with Shiga toxin-producing *Escherichia coli* and is more common in children. Pathogenesis of the syndrome is based on the reaction of the innate immune system to toxemia. Characteristic features of the syndrome are hemorrhagic enterocolitis, hemolytic anemia, thrombocytopenia and acute renal failure, but some patients develop more unusual manifestations that potentially lead to MOF and increase mortality. Even though 70% of children with HUS recover without permanent health consequences, 2–5% of patients die in the acute phase.

Among less usual manifestations, CNS involvement is the most frequent and one that significantly increases morbidity and mortality. Neurological signs range from epileptic seizures to reduced consciousness level and focal motor deficits, but are mostly the result of temporary dysfunction rather than irreversible damage. Brain MRI is the diagnostic method of choice when analyzing possible CNS lesions and usually reveals a pattern of symmetrical hyperintensities in basal ganglia and the thalamus.

The gastrointestinal and hepatobiliary tract may be affected from the esophagus to the perianal area and possible disorders include ileus, intussusception, bowel distension, perforation, necrosis, toxic megacolon, intestinal stricture, rectal prolapse, hepatocellular cholestasis and pancreatitis, which may lead to diabetes mellitus.

Cardiovascular instability can be a consequence of tachyarrhythmia caused by HUS, electrolyte imbalance and toxic myocarditis with subsequent dilatative cardiomyopathy.

The mainstay of treatment is supportive and includes control of fluid and electrolyte balance, optimal enteral and parenteral nutrition, use of hemodialysis (required in approximately two-thirds of patients), control of hemodynamic stability and judicious transfusion of blood derivatives. Plasmapheresis with fresh frozen plasma is reserved for the most severe cases, although it was proven ineffective in some recent controlled clinical trials. Platelet transfusions are avoided and limited for control of active bleeding, considering some studies have suggested it could contribute to the microthrombosis and worsen outcome. Use of antibiotics is controversial and should be avoided, since some studies have shown them to be harmful as possible triggers for the development of HUS in patients with enterohemorrhagic *E. coli* infection.

Already known predictors of poor outcome, including death and chronic renal and lung disease, are prolonged oliguria or anuria, need for hemodialysis, neurological impairment, persistent leukocytosis > 20×10⁹/L, hematocrit < 23% on admission and severe dysfunction of the gastrointestinal system. The recorded extreme leukocytosis in this case was related to the particularly high activity of systemic inflammation as a result of toxemia rather than infection. It could be an indicator of HUS complicated with MOF and, therefore, one of the main negative prognostic factors. Despite the fact that all these predictors were recognized in the acute stage of disease, our patient fully recovered without any apparent sequelae during the follow-up period of one year. This case indicates that early diagnosis, thorough supportive treatment, including renal replacement therapy and early plasmapheresis, are crucial interventions for favorable outcomes in severe cases of typical HUS presented by MOF.

Consent
Written informed consent for publication of this case report and corresponding images was obtained from the patient’s parents.

Author contributions
All three authors analyzed the data and critically revised the manuscript. DM is the first author who designed and wrote the majority of the paper and collected the data. AB made images and graphical analysis of the data. ZZ was the primary clinician responsible for the treatment of this patient and had the initial idea about the paper.

Competing interests
No competing interest were disclosed.

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References

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Grant Luxton
Prince of Wales Clinical School, Faculty of Medicine, University of New South Wales, Sydney, NSW, Australia

The changes are minor and I have no issue with them.

Competing Interests: No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 1

Reviewer Report 25 June 2014

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Zeljko Bosnjak
Anesthesiology Research, Medical College of Wisconsin, Milwaukee, WI, USA

I have no further comments.

Competing Interests: No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
Grant Luxton
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This is a case report documenting the response to treatment with best supportive care including dialysis as well as plasmapheresis in a four year old child with "typical" haemolytic uraemic syndrome. As the authors point out, trial evidence is not supportive of plasmapheresis for this condition however there may be situations where it may be of benefit - especially in a patient not responding to best supportive care. I have no issues with the case report and believe it is worth indexing to remind us that plasma exchange should still be considered in selected patients with this disease. With the potential availability of eculizumab however, the role of plasmapheresis may become even more limited.

Competing Interests: No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Dino Mijatovic, University Hospital Rijeka, Rijeka, Croatia

I welcome your comment! We (all the authors) discussed the potential use of eculizumab during the early course of the disease in the second week. However, we didn't have any experience with it and it seemed to be very expensive treatment. Also, we found more data in the recent literature confirming the efficacy of this drug in atypical cases of HUS, rather than typical. Therefore, we decided to initiate the transfusion treatment with full awareness and precautions regarding its potential side-effects, and to leave the eculizumab as the second-line therapy.

Competing Interests: No competing interests are present in the case of commenting this referee's report.
Zeljko Bosnjak
Anesthesiology Research, Medical College of Wisconsin, Milwaukee, WI, USA

The authors report a very interesting but extremely difficult case report that many will find quite helpful. I have few minor suggestions:

- **Abstract (case presentation):** rewrite the second sentence.
- **Figures 1 and 2:** Could easily be combined into one figure, with single labeling on the top, and placing descriptions of the parameters on top of each tracing. Also the units of diuresis should be changed to per day instead. For plasma creatinine, include the micro symbol.

- **Figures 3-5:** Perhaps white arrows would be a better choice.

**Competing Interests:** No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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Author Response 11 Jun 2014

Dino Mijatovic, University Hospital Rijeka, Rijeka, Croatia

Dear Professor Bošnjak,

thank you very much for your comment! Unfortunately, we weren't able to write the response since we have been extremely occupied with organization of the national congress. Anyway, we found ourselves very grateful for your meticulous and thoughtful analysis. We decided to rewrite the second sentence in Case presentation. We also made proper corrections regarding the units of diuresis and plasma creatinine. We have still not combined figures 1 and 2 into one as we want to stress the extreme leukocytosis and its recovery during the PAF treatment, the process that was parallel to the monitored activity of LDH, which is the biomarker of this disease. We believe that this pattern becomes more appreciated when it is stressed in the figure with one or two diagrams. In the uniform picture it would be among 4 different diagrams.

We have left black arrows for figures 3-5 because their tops are a greater contrast to the density of the object on the scan.

**Competing Interests:** No competing interests are present to influence our judgment of this referee's comment.
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