Hemolytic uremic syndrome complicating whooping cough

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SUMMARY

Introduction We shall present a case of a two-month old infant who has developed a haemolytic uremic syndrome as an atypical complication of Bordetella pertussis infection. The observation that the development of haemolytic uremic syndrome is a late complication of Bordetella pertussis infection may be a clue for further studies.

Case outline A two-month-old female infant was admitted to the hospital because of fever, intensive cough, shortness of breath and poor feeding. Real-time polymerase chain reaction (PCR) for Bordetella pertussis was positive. A macrolide was introduced in therapy. On the eighth hospital day, the infant’s condition improved, she became afebrile and eupneic. On the 16th hospital day, she developed signs of progressive respiratory distress and oliguric acute kidney injury. Hemolytic uremic syndrome (HUS) was diagnosed, so the therapy with the fresh frozen plasma (FFP) transfusion, therapeutic plasma exchange and peritoneal dialysis was initiated. Levels of ADAMTS13 (a disintegrin and metalloproteinase with a thrombospondin type 1 motif, member 13) were decreased, while the levels of factor H, factor B, and factor I were normal. Despite the full supportive and targeted care, severe multiple organ failure had developed and on the 24th hospital day the infant died.

Conclusion Further studies are necessary to identify the mechanism of potential interaction between pertussis toxins, pathophysiology of the infection and the interaction of complement activation, coagulation and the regulation of these cascades.

Keywords: hemolytic uremic syndrome; pertussis; infant

INTRODUCTION

Typical features of hemolytic uremic syndrome (HUS) are microangiopathic hemolytic anemia and acute thrombocytopenia, accompanied by acute kidney injury. In addition to infections with Shiga-like toxin-producing Escherichia coli (STEC-HUS), complement alternative pathway dysregulation may also predispose to atypical HUS (complement mediated aHUS). HUS may also develop in the context of severe infections, including for example Streptococcus pneumoniae, Shigella dysenteriae and Bordetella pertussis, various viruses, like H1N1 influenza, Cytomegalovirus and Parvovirus B19 [1, 2, 3]. In addition to thrombotic thrombocytopenic purpura (TTP) and STEC-HUS, aHUS is classified in a group of thrombotic microangiopathies (TMA).

The incidence of aHUS and TTP is around 3/1,000.000 among children under 18. Patients with TTP have a good therapeutic response to plasma exchange (PE), whereas this is not the case in patients with aHUS. Mortality in patients with TTP is around 10%, and 25% in patients with aHUS during the acute phase [4, 5].

Despite the fact that vaccine is widely available, Bordetella pertussis infections are still very common. The classic symptoms of pertussis are paroxysmal whooping cough, marked leukocytosis and related pulmonary leukostasis. So far, only four cases of HUS associated with Bordetella pertussis infection have been described in the literature [6–9].

CASE REPORT

A two-month-old infant was admitted to the hospital due to fever, intensive cough, shortness of breath and poor feeding. First symptoms have begun two days before hospital admission.

Five days before admission, she received a combined five-component vaccine (diphtheria, tetanus, pertussis – acellular component, Poliovirus (Inactivated) Vaccine (Adsorbed) and Haemophilus Influenza Type B Conjugate Vaccine). She was the second child of the healthy non-consanguineous parents, born at term, weighting 2,900 g. Prenatal and perinatal period went without any complications.

On admission, she was conscious and alert, febrile, with normal vital parameters and normal body weight of 4,800 g. Lung auscultation revealed wheezing and bilateral, fine, late-inspiratory crackles on the bases of the lungs. All other physical findings were normal.
Initial laboratory findings are shown in Table 1. All other biochemical findings were within the normal ranges. Chest X-rays revealed pulmonary consolidation in perihilar and basal regions of the right lung and, also, regions of hyperinflation on both lungs. Ultrasound of the brain and abdomen showed normal findings.

The initial treatment included inhaled corticosteroids and bronchodilators and parenteral antibiotic therapy (cef-tazidime). Despite the therapy, the child’s condition worsened and she became tachydyspnoic, pale, adynamic and continuously febrile. On the fifth hospital day, marked leukocytosis was registered (Table 1). Peripheral blood smear showed prevalence of mature segmented granulocytes with toxic granulations, with normal red blood cell and platelet count and morphology.

Microbiologic and serologic tests confirmed the diagnosis of *Bordetella pertussis* infection (whooping cough). Real-time polymerase chain reaction (PCR) for *B. pertussis* was positive. Blood cultures were negative.

A macrolide was introduced in therapy. On the eighth hospital day, the infant’s condition improved, she became afebrile and eupneic. This clinical melioration was accompanied with marked improvement of laboratory findings and reduction of WBC count.

On the 16th hospital day, the infant’s condition suddenly worsened again. She had frequent attacks of heavy cough and developed signs of progressive respiratory distress with consequent oxyhemoglobin desaturation. She became edematous (gained more than 600 g in weight in two days), with a decreased urine output (0.61 ml/kg/h), and hypertensive (BP 128/66 mmHg) so the infant was admitted to the Intensive Care Unit (ICU).

On admission to ICU, laboratory tests showed elevated procalcitonin levels, elevated WBC count, signs of hemolytic anemia and mild thrombocytopenia, as well as the signs of renal failure (Table 1). Peripheral blood smear revealed anisocytosis, polychromasia, presence of schistocytes and erythroblasts. Reticulocytes 23%, hap-toglobin levels were < 0.10 g/l (0.3–2 g/l). Lactate dehydrogenase 64.15 μkta/l (< 7.52 μkta/l). Urea 15.8 mmol/L (1.8–6 mmol/l), creatinine 328.5 μmol/l (14–34 μmol/l), uric acid 1,311 μmol/l (65–319 μmol/l). Blood gas analysis showed decompensated metabolic acidosis. Hemostasis screening tests were normal.

Those clinical and laboratory findings raised suspicion of the presence of HUS, so we started the therapy with the fresh frozen plasma (FFP) transfusion. Hypertension was treated with calcium channel blocker.

Next day (17th hospital day), the infant became tachydyspnoic, hence, she was intubated and mechanical ventilation was initiated. Because the child became anuric, the same day, Tenckhoff catheter for peritoneal dialysis (PD) was inserted and continuous peritoneal dialysis was immediately started. Initially, the dialysis solution with 2.3% glucose was used, and then after the edema was gone it was replaced with the solution with

### Table 1. Laboratory findings and clinical course

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Day 1</th>
<th>Day 5</th>
<th>Day 10</th>
<th>Day 14</th>
<th>Day 16</th>
<th>Day 17</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemoglobin (g/l)</td>
<td>111</td>
<td>105</td>
<td>93</td>
<td>83</td>
<td>61</td>
<td>76</td>
</tr>
<tr>
<td>Platelet count (10^9/l)</td>
<td>552</td>
<td>778</td>
<td>1,137</td>
<td>481</td>
<td>95</td>
<td>46</td>
</tr>
<tr>
<td>White blood cell (10^9/l)</td>
<td>18</td>
<td>90</td>
<td>32</td>
<td>18</td>
<td>32</td>
<td>29</td>
</tr>
<tr>
<td>Creatinine (μmol/l)</td>
<td>/</td>
<td>/</td>
<td>/</td>
<td>/</td>
<td>328.5</td>
<td>398.47</td>
</tr>
<tr>
<td>CRP (mg/l)</td>
<td>45.9</td>
<td>36.7</td>
<td>negative</td>
<td>negative</td>
<td>13.34</td>
<td>/</td>
</tr>
<tr>
<td>Procalcitonine (ng/ml)</td>
<td>/</td>
<td>/</td>
<td>0.38</td>
<td>/</td>
<td>4.39</td>
<td>75.0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Clinical development</th>
<th>Coughing, heavy breathing</th>
<th>Tachydyspnoic, febrile, coughing, adynamic</th>
<th>Better overall condition, coughing</th>
<th>Paroxysmal whooping cough</th>
<th>Oliguria, edema, dyspnea</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Therapy</th>
<th>Antibiotic therapy (ceftazidime)</th>
<th>Antibiotic therapy (macrolide)</th>
<th>Antibiotic therapy (macrolide)</th>
<th>Intensive Care Unit</th>
</tr>
</thead>
</table>
| FFP – frozen fresh plasma; CAPD – continuous ambulatory peritoneal dialysis, TPE – therapeutic plasma exchange

### Table 2. Findings of immunologic tests of parents and the neonate

<table>
<thead>
<tr>
<th>Parameter (reference range)</th>
<th>Patient before FFP infusion (16th hospital day)</th>
<th>Patient after FFP infusion (17th hospital day)</th>
<th>Father</th>
<th>Mother</th>
</tr>
</thead>
<tbody>
<tr>
<td>Classical pathway activity CH50/mL (48–103)</td>
<td>51</td>
<td>42</td>
<td>128</td>
<td>186</td>
</tr>
<tr>
<td>C3 g/L (0.9–1.8)</td>
<td>0.88</td>
<td>0.54</td>
<td>128</td>
<td>122</td>
</tr>
<tr>
<td>C1q mg/L (60–180)</td>
<td>36</td>
<td>40</td>
<td>128</td>
<td>186</td>
</tr>
<tr>
<td>C4 g/L (0.15–0.55)</td>
<td>0.38</td>
<td>0.21</td>
<td>0.27</td>
<td>0.42</td>
</tr>
<tr>
<td>Alternative pathway activity % (70–105)</td>
<td>60</td>
<td>54</td>
<td>128</td>
<td>1209</td>
</tr>
<tr>
<td>Factor B % (70–130)</td>
<td>139</td>
<td>125</td>
<td>146</td>
<td>162</td>
</tr>
<tr>
<td>Factor H mg/L (250–880)</td>
<td>429</td>
<td>358</td>
<td>882</td>
<td>1209</td>
</tr>
<tr>
<td>Factor I % (70–130)</td>
<td>138</td>
<td>99</td>
<td>146</td>
<td>162</td>
</tr>
<tr>
<td>ADAMTS13 activity % (67–147)</td>
<td>43</td>
<td>50</td>
<td>146</td>
<td>162</td>
</tr>
</tbody>
</table>
1.5% glucose. The initial fill volume of the dialysis solution was 10 ml/kg per exchange and was increasing slowly towards 20 ml/kg. Exchanges were done every 20–30 min. at the beginning, and after that hourly. Dwell time was 10 minutes, same as the drain time.

Also, the same day, two-lumen central venous catheter was inserted and the first session of therapeutic plasma exchange (TPE) was performed. During the hospitalization, a total of five TPE sessions have been done. Despite the applied therapy, signs of anuric renal injury, thrombocytopenia, anemia, and elevated lactate dehydrogenase levels continuously persisted.

Additional laboratory tests were carried out for the full work-up and TMAs differentiation. Findings of immunologic tests of the parents and the patient are shown in Table 2.

There was no available material for DNA extraction and genetic analysis of the patient. In both parents, complement profile was normal and the levels of some factors were elevated. Sequencing of the complete CFH gene of both parents was done, and no rare variations were observed, however, both parents turned out to carry the H3 aHUS risk haplotype in heterozygous manner.

Despite the full supportive and targeted care, severe multiple organ failure developed, and on the 24th hospital day the infant died.

DISCUSSION

In the European cohort, 16% of aHUS cases were reported as a secondary HUS [10]. When a secondary HUS occurs, such as in the setting of different diseases, signs and symptoms of the primary disease can confound the diagnosis of aHUS [11].

Non-deficient ADAMTS13 activity (over 5–10%) supports the diagnosis of aHUS (under the condition that STEC test is negative). Of course, in these conditions, a differential diagnosis must be distinguished between aHUS and secondary HUS. Secondary HUS may be associated with different infections and sepsis, use of medications (anticancer molecules, immunotherapeutics, like cyclosporine, tacrolimus, and antiplatelet agents), malignancies and other underlying medical conditions such as autoimmune diseases, scleroderma and antiphospholipid syndrome. Familial aHUS occurs in about 10–20% of cases, the remaining patients having sporadic disease. Hypocomplementemia with a low level of C3 but normal C4 is a sign of alternative pathway dysregulation and activation, however, in 60–80% of the patients with aHUS levels of C3 are normal [12, 13, 2].

Due to the underlying immune-mediated, auto-inflammatory mechanisms and missing complement regulators, transfusions of FFP and plasmapheresis are therapeutic options for this syndrome. The aim of plasma therapy is to replace the mutant elements of the complement with normal elements in order to eliminate the pro-inflammatory and thrombogenic factors responsible for the symptoms. It is assumed that injections of fresh frozen plasma alone are sufficient in the case of quantitative deficits [14]. But the long-term outcomes for this treatment are still not well-known [15].

In the case of the treatment failure (FFP, TPE), application of eculizumab (humanized monoclonal antibody against complement protein C5 terminal) is recommended [16, 17]. Studies that reported the benefits of these therapeutic choices in patients with aHUS are accumulating, but further evaluation is required to guide early and late therapeutic decisions like up-front treatment, treatment duration and discontinuation [18, 19, 20]. Baskin et al. [18] reported 10 paediatric patients with aHUS who did not respond to PE. Eculizumab improved their renal function and quality of life. Greenbaum et al. [19] prospectively evaluated efficacy and safety of eculizumab in paediatric patients with aHUS. Their findings establish the efficacy and safety of eculizumab for the paediatric patients with aHUS and the recommendation is to start the therapy with it as soon as possible after establishing the aHUS diagnosis. Considering the cost of the treatment, the cost/benefit of this cure should be estimated more thoroughly in the future.

This drug was neither registered nor available in Serbia, at the time of our patient's treatment.

B. pertussis produces a number of virulence factors that are involved in the pathogenesis and manifestations of the disease. Toxins of B. pertussis and other virulence factors enable adhesion of bacteria, locally injure epithelium, cause leukocyte dysfunction and macrophage cytotoxicity, increase release of proapoptotic and pro-inflammatory cytokines such as TNF-alpha and IL-6. In addition to the damage to the epithelium, these factors injure the endothelium, which causes a pro-coagulant pathway activation [21].

The infant presented in our report has received the first dose of pertussis vaccine two days prior getting the disease, so she has not developed the immunity.

So far, only four cases of HUS associated with Bordetella pertussis infection have been described in the literature [6–9]. The association of Bordetella pertussis infection and HUS was first described by Berners et al. [6] in a newborn with complement factor H (CFH) mutation, which was treated with peritoneal dialysis and ended fatally. In this patient, an abnormal band was identified on Factor H Western-blot, indicative for the presence of an abnormal complement regulator. After that, Pela et al. [9] described the case of a 42-day-old infant treated with hemodialfiltration and FFP infusions. In this case mutations in the genes encoding factor H and MCP were not detected. This patient survived with a completely recovered renal function. Chaturvedi et al. [7] described the case of a 28-day-old baby who was treated with FFP infusions and did not require dialysis. Similarly to previously mentioned case, mutations in the genes encoding factor H, factor I, factor B and ADAMTS13 were not identified. The patient survived, fully recovered and maintained a normal renal function. The most recent case was reported by Cohen-Ganea et al. [8]. In this case, complement (C3, C4, CH50) and factor H levels were within the normal range.
Our patient was a two-month-old infant who had developed HUS three weeks after starting the whooping cough, and who was treated by peritoneal dialysis and TPE without a favorable response. Death of the patient was caused by HUS and severe sepsis which resulted in multiple organ dysfunction.

In the acute phase, very low C1q levels with decreased C3 concentration and combined (classical and alternative pathway) complement consumption were observed. We apprehended that as a result of ongoing severe infection, not as dysregulation of the alternative pathway. In line with this, there was no evidence of Factor H mutation in this patient, since the parents did not carry rare variations in CFH. Hence, we consider the possibility of a cause-effect relationship between B. pertussis infection and HUS in our patient. B. pertussis infection may be capable to stimulate the inflammatory cells to release the cytokines which determine microangiopathy and HUS. No abnormalities of factor H were found in the parents of our patient, however, due to the lack of available samples for DNA extraction we were unable to completely sequence all of the important genes of complement regulators and factors. Therefore, we can only assume that B. pertussis may cause HUS in the children without any predisposing factors. The infant developed HUS in the later phase of the infection after a brief period of clinical improvement (from the eighth to the 14th hospital day). From this aspect our patient was similar to the three previously reported patients who had HUS in the context of an infection caused by B. pertussis. These patients also developed a severe HUS after a period of clinical improvement – in the period between the sixth and 42nd hospital day [6, 7, 9]. The patient described by Cohen-Ganelin et al. [8] was in difficult condition with paroxysmal coughing and extreme excitability, all the time until the moment of HUS diagnosis (the 10th hospital day). So far, all described cases have developed HUS between the 12th and 45th hospital day since the start of the illness (first symptoms of a whooping cough).

Further studies are necessary to identify the mechanism of a potential interaction between pertussis toxins, pathophysiology of the infection and the interaction of complement activation, coagulation and the regulation of these cascades. The observation of the development of HUS as a late complication of B. pertussis infection may be a clue for further studies.

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REFERENCES

САЖЕТАК

Увод
Приказан је случај двомесечног одојчета које је развило хемолитичко-уремички синдром као атипичну компликацију инфекције бактеријом *Bordella pertussis*. Чињеница да је развој хемолитичко-уремичког синдрома касна компликација инфекције бактеријом *Bordetella pertussis* може бити полазна тачка за планирање даљих студија.

Приказ болесника
Двомесечно женско одојче је хоспитализовано на Клиници за педијатрију због повишене температуре, интензивног кашља, диспнеје и одбијања хране. Ланчна реакција полимеразом у реалном времену на бактерију *Bordetella pertussis* може бити полазна тачка за планирање даљих студија. Планрана терапија макролидима и осмог дана хоспитализације клиничко стање детете се побољшало, постало је афебрилно и еутерично. Шеснаестог дана хоспитализације дете је развило клиничке знако прогресивног респираторног дистреса и олигуричног акутног бубрежног оштећења. Дијагностикован је хемолитичко-уремијски синдром, те је започета терапија трансфузима свеже смрзнуте плазме, а затим и терапије заместима плазме и перитонеумска дијализа. Нивои металопротеиназе ADAMTS били су снижени, док су нивои фактора H, В и I били нормални. Упркос спроведеној терапији, у даљем току развија се мултиорганска дисфункција и 24. дана хоспитализације долази до смртног исхода.

Закључак
Потребне су додатне студије да би се утврдили могући механизми интеракције између токсина пертусиса, патофизиологије инфекције и интеракције активације комплемента, коагулације и регулације ових каскада.

Кључне речи: хемолитичко-уремички синдром; велик кашљ; одојче

Хемолитичко-уремички синдром као компликација великог кашља

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