A Fatal Case of COVID-19 in a Child with ALL: A Cytokine Storm and Hyperferritinemic MODS

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Abstract

Since little is known about dysregulated hyperinflammatory immunological responses causing acute severe infection and multisystem inflammatory syndrome in children associated with coronavirus disease 2019 (COVID-19), the available data on therapies for severe presentations in children are very limited. Describing experiences of severe pediatric COVID-19 presentations in more detail will help improve clinical practice.

In this case report, we describe the complete clinical course of a 9-year-old girl previously diagnosed with Angelman syndrome and high-risk T cell acute lymphoblastic leukemia who had been receiving reinduction chemotherapy, presented with pneumonia and acute respiratory distress syndrome, and progressively developed hyperferritinemic multiple-organ failure, a cytokine storm, and coagulopathy associated with COVID-19. She was treated with therapeutic plasma exchange, tocilizumab, hydrocortisone, and favipiravir, but she died 7 days after her admission into our pediatric intensive care unit.

The utility of therapeutic plasma exchange with other immunomodulatory therapies in severe presentations requires further trials. The spectrum of the inflammatory phenotypes associated with COVID-19 should be investigated and well defined to initiate the optimal treatment strategy on time.

Keywords

► acute lymphoblastic leukemia
► coronavirus disease 2019 (COVID-19)
► critically ill children
► hyperferritinemia
► therapeutic plasma exchange

Introduction

As the worldwide outbreak of a new type of coronavirus spreads, coronavirus disease 2019 (COVID-19), new data has emerged, indicating an unignorable amount of severe presentations among the pediatric population.¹–⁷ Mortality has been reported among children developing progressive multiple-organ failure.²–⁷

Since little is known about dysregulated hyperinflammatory immunological responses causing the acute severe infection and multisystem inflammatory syndrome in children (MISC) associated with COVID-19, the available data on therapies for severe presentations in children are very limited.³,⁴,⁶,⁷ Describing experiences of severe pediatric COVID-19 presentations in more detail will help improve clinical practice.

In this case report, we describe the complete clinical course of a 9-year-old girl diagnosed with high-risk T cell acute lymphoblastic leukemia (ALL) who had been receiving reinduction chemotherapy, presented with pneumonia and acute respiratory distress syndrome (ARDS), and progressively developed hyperferritinemic multiple-organ failure, a cytokine storm, and coagulopathy associated with COVID-19. She was treated with therapeutic plasma exchange (TPE), tocilizumab,
hydrocortisone, and favipiravir, but she died 7 days after her admission into our pediatric intensive care unit (PICU).

Case

Our patient is a 9-year-old girl with mild motor–mental retardation due to Angelman syndrome who had also been diagnosed with high-risk T cell ALL 6 months previously, had been in remission after first-induction chemotherapy, and had been treated with high-risk blocks for the last 3 months. Eight days after the last time she was admitted to the ward to receive her high-risk block, she developed fever. She was administered antibiotics for febrile neutropenia, but she developed cough 4 days later. A chest X-ray was taken, and her initial diagnosis was fungal pneumonia. Empiric antibiotics were widened to vancomycin, meropenem, trimethoprim–sulfamethoxazole, amphotericin B, and ganciclovir. Her cough and fever persisted, and she was admitted to a PICU after she developed mild respiratory distress and oxygen requirement on the second day of her cough and the sixth day of her fever. She was provided noninvasive mechanical ventilation (NIV) with a full-face mask, continuous positive airway pressure 6 cm H₂O, fraction of inspired oxygen (FiO₂) 40%. She had pneumonia with a SpO₂/FiO₂ ratio of 277, normal blood gas analysis, and hemodynamical stability. After 16 hours on NIV, a chest X-ray and thorax computed tomography showed the progression of infiltrates (► Fig. 1A–1I) bilaterally, and she was electively intubated. The patient had mild pediatric ARDS with a Pmax of 250. An endotracheal aspirate was taken for Gram staining, culture, influenzas A and B, and severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2).

No bacteria was detected in Gram staining, and viral polymerase chain reaction (PCR) was negative for influenza. The patient was hemodynamically stable and received intravenous immunoglobulin, wide-spectrum antibiotics, and supportive care (blood products) without inotropes while waiting for the SARS-CoV-2 test result. Fifteen hours after intubation, she developed sinus tachycardia that was unresponsive to fluids and rapidly developed metabolic acidosis, renal failure, and hyperdynamic shock requiring noradrenalin and adrenaline infusion, and worsening oxygenation simultaneously. Continuous renal replacement therapy (CRRT) started immediately for metabolic acidosis, fluid expansion, inotrope, and fluids were titrated, and mean airway pressure and oxygen needed to be increased with a Pmax/FiO₂ ratio of 56. The result of the PCR test for SARS-CoV-2 was then available, and the patient was diagnosed with COVID-19 simultaneously. She progressively developed shock and multiple-organ failure in 5 hours, and tests showed high inflammatory markers, coagulopathy, and organ failure, consistent with the diagnosis for fulminant COVID-19. Favipiravir, hydrocortisone, tocilizumab, and vitamin C were added to supportive treatment. At the end of the day (on PICU day 3), her pupils were both mildly reactive to light and dilated. On PICU day 4, TPE was begun for severe sepsis with rapidly developing hyperferritinemic multiple-organ dysfunction syndrome (MODS) associated with a cytokine storm of fulminant COVID-19. A second dose of tocilizumab was given after TPE, and CRRT was continued with hydrocortisone, favipiravir, and other supportive care. Fresh frozen plasma was used as replacement fluid, and the patient received 1.5 plasma volume exchange on PICU day 4 and 5 followed by 1 volume exchange on PICU day 6 and 7. Laboratory findings,
|                         | Onco. ward admission | PICU-day 1 | PICU-day 2 | PICU-day 3 | PICU-day 4 | PICU-day 5 | PICU-day 6 | PICU-day 7 |
|-------------------------|----------------------|------------|------------|------------|------------|------------|------------|------------|
| **WBC (mm³)**           | 2,120                | 2,120      | 2,120      | 2,120      | 2,120      | 2,120      | 2,120      | 2,120      |
| **Lymp/Neutro**         | 760/780              | 760/780    | 760/780    | 760/780    | 760/780    | 760/780    | 760/780    | 760/780    |
| **Platelet**            | 115000               | 115000     | 115000     | 115000     | 115000     | 115000     | 115000     | 115000     |
| **LDH (U/L)**           | 140                  | 140        | 140        | 140        | 140        | 140        | 140        | 140        |
| **Ferritin (ng/mL)**    | >100.000             | >100.000   | >100.000   | >100.000   | >100.000   | >100.000   | >100.000   | >100.000   |
| **PT/PTT (sec)**        | 12.4/29.7            | 13.2/39.2  | 13.8/36.8  | 12.1/27.8  | 12.7/43.3  | 13.5/28.4  | 13.5/28.4  | 13.5/28.4  |
| **Fibrinogen (g/L)**    | 4.5                  | 4.5        | 4.5        | 4.5        | 4.5        | 4.5        | 4.5        | 4.5        |
| **AST/ALT (U/L)**       | 42/62                | 110/167    | 115/153    | 862/252    | 1,825/662  | 1,794/760  | 303/232    | 144/91     |
| **BUN (mg/dL)**         | 5.3                  | 7.3        | 14.6       | 22.4       | 14         | 19.1       | 28.7       | 25.4       |
| **Crea. (mg/dL)**       | 0.42                 | 0.5        | 0.9        | 1.6        | 2.4        | 2.1        | 1.8        | 1.6        |
| **T.Bil (mg/dL)**       | 0.43                 | 0.83       | 0.49       | 0.89       | 1.77       | 1.32       | 1.9        | 1.5        |
| **D-dimer (mg/L)**      | 8.74                 | 19.44      | 19.55      | >20        | >88        | >87        | >87        | >87        |
| **IL-6 (pg/mL)**        | 336.7                | 348.8      | 292.5      | 261.4      | 102.9      | 43.7       | 25.9       | 25.9       |
| **CRP (mg/L)**          | 1.1                  | 8.4        | 96.1       | 163.1      | 108        |            |            |            |
| **PCT (µg/L)**          |                      |            |            |            |            |            |            |            |
| **Respiratory support** |                      |            |            |            |            |            |            |            |
| **Vasopressors**        |                      |            |            |            |            |            |            |            |
| **Lactate (mmol/L)**    | 1.6–1.1              | 1–1.6      | 5.8–2.6    | 1.4–2.5    | 1.4–2.1    | 0.9–1.4    | 0.27       |            |
| **Clinical course**     | Stable               | 39°C       | 39°C       | 38.7°C     | 38°C       | 38°C       | 38.2°C     | 37°C       |
|                         | NIV (4p.m)           | Elective intubation (10 µu) | MODS | MODS | MODS | Repeated cultures of tracheal aspiration, blood, urine: negative artérials | Exitus | 30°C (11p.m): Fixed dilated pupil, No gag reflex |
treatment, and notable clinical findings for the patient are shown in Table 1.

With vasopressor support for shock, and conventional mechanical ventilation, X-ray findings showed improvement (Fig. 1A–11). Serum ferritin, C-reactive protein (CRP), lactate dehydrogenase (LDH), and interleukin-6 (IL-6) decreased, and the number of platelet suspensions decreased gradually after each TPE session. However, neurological findings remained unchanged, with mildly reactive pupillary reflexes to light, dilatation, and low Glasgow coma scale (E1M3V1) but preserved spontaneous breathing and gag reflex. Retinal hemorrhage and papilledema were detected on fundus examination. Sedation was stopped, levetiracetam was given for potential seizures, and antiedema treatment was given for probable increased intracranial pressure. On PICU day 6, acro-ischemia occurred (Fig. 2), but a lower-extremity ultrasound did not show thrombosis. Repeated cultures of endotracheal aspiration and other sides for bacteria and fungus were negative. She died on PICU day 7 because of severe neurological dysfunction.

Discussion

Turkey’s first COVID-19 cases were confirmed on March 11, 2020. By September 1, 2020, COVID-19 had caused 6,326 deaths in the country, including 12 people under 15 years of age.8

Severe presentations associated with COVID-19 in children are defined as MISC, a postinfectious inflammatory process that has mostly affected children who had previously been healthy. Severe COVID-19 disease is an acute severe infection reported in children and the adult population with existing comorbidities.7,9 Also recently, subgroups of MISC have been defined with different manifestations but a similar process.7 The primary hallmark is a hyperinflammatory state because of a dysregulated immune response triggered by the novel coronavirus, resulting in MODS and coagulopathy, which have been found to predict severity.1,6,9–11 Our patient had profound immunosuppression with very low lymphocyte and neutrophil counts (10/mm³) from the beginning of treatment due to high-risk chemotherapy. She developed severe ARDS and MODS progressively after COVID-19 pneumonia and, finally, acro-ischemia attributed to microthrombosis. She experienced a cytokine storm and COVID-19 coagulopathy that mimicked adult fulminant COVID-19 cases and presented with predictors of a fatal outcome, such as very high ferritin, CRP, D-dimer, and IL-6.8

The patient had several clinical features of secondary hemophagocytic lymphohistiocytosis (HLH), macrophage activation syndrome (MAS), and thrombocytopenia-associated multiple-organ failure (TAMOF)—that is, phenotypes of hyperferritinemia—critically ill diseases with MODS, such as highly elevated ferritin, elevated transaminases, elevated LDH, thrombocytopenia, and lymphopenia. These symptoms have also been reported in acute severe COVID-19, and MISC patients.9,10,14,15 COVID-19–induced cytokine storm with the characteristic coagulopathy of COVID-19 has common pathogenesis and clinical as well as laboratory findings with severe sepsis and MODS, secondary HLH, MAS, and TAMOF.15–17 Hyperferritinemia MODS in severe
In the clinical course of these types of critical diseases, transport issues to the radiology department may have been due to in.

Our patient had profound thrombocytopenia with high fibrinogen and markedly elevated D-dimer levels. Her D-dimer levels remained elevated, with lower levels of fibrinogen and normal ranges for prothrombin time and activated partial thromboplastin time. Laboratory findings and the patient’s clinical course showed overlapping coagulopathies, such as a spectrum of TAMOF (thrombotic thrombocytopenic purpura, thrombotic microangiopathy, and sepsis-induced disseminated intravascular coagulation [DIC]), but she also had unique features of COVID coagulopathy, such as very high D-dimer levels, low-grade DIC, and microthrombosis.

Therapeutic strategies targeting this overactive cytokine response for severe COVID-19 patients are available, but they have not yet been approved. TPE has been shown to remove the proinflammatory cytokines leading to MODS and, furthermore, restore hemostasis and resolve organ dysfunction in patients with hyperferritinemic syndromes, such as severe sepsis and MODS, secondary HLH, MAS, and TAMOF phenotypes. Observational studies have examined TPE’s utility in the clinical course of these types of critical diseases. Moreover, a recent case and case series have reported improved outcomes for severe COVID-19 patients sharing many features with the critical diseases treated using TPE.

Our patient was treated with tocilizumab, favipiravir, and hydrocortisone, combined with TPE. Hemodynamics, organ dysfunction (pulmonary, hepatic, hematologic, and cardiovascular), and inflammatory markers had shown improvements after TPE sessions, beginning from the first TPE session, but severe neurological dysfunction had not improved. Unfortunately, the patient died of central nervous system complications that may have been due to inflammation or infection (brain edema) or coagulopathy (hemorrhage or infarction). Furthermore, we think that the multidrug therapy patient received did not have a negative effect on the brain. Our fateful limitation was that we could not diagnose these complications because of transport issues to the radiology department.

Our patient’s mother and the healthcare workers tested negative for SARS-CoV-2. The management of COVID-19 pneumonia in profoundly immunocompromised patients is far different from other etiologies. Two precious days before intubation might have been the critical point for testing that we missed in this case. Following this case, our institution has implemented a practice of administering SARS-CoV-2 PCR tests for all immunocompromised patients with pneumonia and/or fever, regardless of their history of contact.

The inflammatory phenotypes and courses of organ dysfunction of severe presentations are not homogenous, but some subgroups of MISC and acute fulminant COVID-19 have similar features. These similar features are rapidly progressive and fatal with MODS and hyperinflammation with abnormal coagulopathy. Although there is insufficient evidence to support TPE with other immunomodulatory treatments, such as anakinra or tocilizumab, we think they should begin early in the course, especially before the initiation of invasive mechanical ventilation in the severe group. Close monitoring of laboratory findings such as LDH, ferritin, CRP, D-dimer, and IL-6, if possible, and H Score along with organ dysfunction and oxygen requirement—is important for not missing the critical point of treatment.

The utility of TPE with other immunomodulatory therapies in severe presentations requires further trials. The spectrum of the inflammatory phenotypes associated with COVID-19 should be investigated and well defined to initiate the optimal treatment strategy on time. According to the relevance of overlap, we think TPE might be beneficial individually or in combination with other immunomodulatory treatments for a subgroup of severe (both acute infection and MISC) COVID-19 patients, like experience in secondary HLH and TAMOF.

Conflict of Interest
None declared.

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