Refractory thrombocytopenia in a severe COVID-19 patient

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Abstract

Severe acute respiratory syndrome-related coronavirus 2 (SARS-CoV-2) infection has spread worldwide. The most of patients presents fever, dyspnea and cough as a typical viral infection, others show peculiar clinical and laboratory signs, such as anosmia, ageusia and thrombocytopenia. We here describe a severe coronavirus disease 2019 patient (76-year old, male) that developed an immune thrombocytopenia and acquired pseudo-thrombocytopenia that were refractory to immunomodulators even after resolution of respiratory failure. The patient developed thrombocytopenia (platelets 88,000/mm$^3$) that got worse the following day (14,000/mm$^3$). We started 1 mg/kg of methylprednisolone i.v. daily. Platelet count increased up to 209,000/mm$^3$, in sodium citrate, but remained about 14,000/mm$^3$ in EDTA 10 days after the beginning of methylprednisolone. The patient showed great improvement in respiratory parameters and radiological finding. About one week after he developed a thrombocytopenia up to 70,000/mm$^3$. We did not modify the steroids dosage. Platelet count slowly began to increase and in about 10 days returned to normal values.

Introduction

Since December 2019, a novel coronavirus [designated severe acute respiratory syndrome-related coronavirus 2 (SARS-CoV-2)] has spread worldwide diffusing a new disease that World Health Organization (WHO) called coronavirus disease 2019 (COVID-19). This disease has peculiar characteristics both respiratory and clinical. As to respiratory findings, patients often present the so-called silent hypoxemia where they refer mild respiratory symptoms despite a severe saturation reduction and a severe hypoxia at arterial blood gases (ABG) analysis. This is due to a diffuse interstitial pneumonia that cause a severe alveolar impairment but preserve the lung compliance, so the patient, in the early stages, can compensate the hypoxia increasing the respiratory effort. This is evident as a respiratory alkalosis at ABG and has been called by some authors as type 1 pneumonia. On the other hand, sometimes patients present a type 2 pneumonia that shows opposite characteristics, so low lung compliance and high lung weight that recalls a typical acute respiratory distress syndrome (ARDS). Moreover, some authors suggest that type 1 pneumonia could evolve to a type 2 in some circumstances causing a greater respiratory effort and perhaps a worse outcome. As to clinical characteristics, there is great variability in presentation and laboratory findings. Even though the most of patients presents fever, dyspnea, fatigue and cough as a typical viral infection, some others show peculiar clinical and laboratory signs, such as anosmia, ageusia and thrombocytopenia. Thrombocytopenia, particularly, can be evident up to 57% of patients among severe cases. Its physiopathology includes: reduced production, increased intravascular destruction and higher consumption. The cytokine storm, described in SARS-CoV-2 infection, could be responsible of all these mechanisms and, moreover, can cause the lung damage that leads to the respiratory failure. Although the greatest part of COVID-19 thrombocytopenia is mild and rarely complicated by hemorrhages, some patients present a severe form. Some authors, indeed, recently reported a case of a severe COVID-19 related thrombocytopenia efficaciously treated with immunomodulators (steroid and immunoglobulins) and eltrombopag. We here describe a severe COVID-19 patient that developed an ITP and acquired pseudo-thrombocytopenia that were refractory to immunomodulators even after resolution of respiratory failure. An informed consent was obtained to write this case report.

Case Report

We describe a case report of a patient (76 years old, male) admitted in the emergency department in April 2020 for severe acute respiratory failure and fever. He presented a past history of chronic obstructive pulmonary disease in first class of GOLD classification, coronary artery disease in conservative treatment, hypertension and mild chronic kidney disease. At admission, the patient was afebrile, presented severe dyspnea, tachypnea and a good hemodynamic compensation (blood pressure 150/80 mmHg, heart rate 80 bpm). He arrived in reservoir mask as oxygen supplementation and was evident, at ABG, hypoxemia with an acute respiratory alkalosis and hyperlactatemia due to the respiratory effort (pH 7.61, pCO$_2$ 16.4 mmHg, pO$_2$ 63 mmHg in FiO$_2$ 80%, lactate 3.71 mmol/L, p/F 78 mmHg). We immediately started a treatment with helmet continuous positive airway pressure with 10 cmH$_2$O of positive end expiratory pressure and a FiO$_2$ of 60%. The physical examination showed only some inspiratory crackles, for the rest was remarkable. As to laboratory findings, he showed mild hyperglycemia and mild worsening in creatinine levels (182 and 1.32 mg/dL respectively), lactic dehydrogenase was 489 IU/L, normal values (n.v.) range from 5 to 248 IU/L, and C-reactive protein was 31.7 mg/L (n.v. 0-7.5). Ferritin, at presentation, was 1500 ng/mL (n.v. 24-336), d-dimer levels were 1590 mg/L (n.v. 50-500), while transaminase, procalcitonin and electrolytes were in normal ranges. The nose swab confirmed a SARS-CoV-2 infection. A complete blood count (CBC) showed a mild anemia (hemoglobin 10.9 gr/dL), but stable compared to previous CBCs, a mild lymphopenia (lymphocytes 670/mm$^3$; n.v. 900-5200) and normal platelets (167,000/mm$^3$), even if they were reduced compared to a...
previous CBC (293,000/mm³). A point of care ultrasound showed a diffuse patchy distribution of interstitial artifactual signs with some pleural consolidations compatible with a COVID-19 interstitial pneumonia, a small and reactive inferior vena cava and normal heart dimension with a mid-range ejection fraction impairment (about 45%). We started a drug therapy, as suggested by our internal protocol, with enoxaparin 4000 UI qd, hydroxychloroquine 200 mg qid the first day and then bid for other 6 days, azithromycin 500 mg qd for 5 days. The day after admission the patient developed thrombocytopenia (platelets 88,000/mm³) that got worse the following day (14,000/mm³). The analyzer flagged the presence of platelet clumps. We did not find any sign of hemolysis and a peripheral blood smear did not show schistocytes but confirmed thrombocytopenia and some aggregates. We performed a platelet count with sodium citrate that confirmed the thrombocytopenia but only mild (69,000/mm³) and did not show platelet clumps in sodium citrate. Coagulation values remained into normal limits. As the patient received heparin also 40 days before, we researched anti PF4 antibodies that resulted negative, while antiplatelet antibodies (specifically anti glycoprotein IIb-IIIa) resulted positive. We started 1 mg/kg of methylprednisolone i.v. daily. We suspended hydroxychloroquine. Five days after admission, the patient still presented a severe respiratory failure, as he needed high flow nasal cannulas (HFNC) at FiO₂ 60% and 60 L/min of flow to achieve a saturation of 92% and the lung ultrasound showed an initial evolution to an ARDS pattern. Platelet count was still 13,000/mm³ in EDTA and 110,000/mm³ in sodium citrate. We decided to administrate tocilizumab as the platelet count in sodium citrate showed a partial improvement and the patient was in the inflammatory phase of the disease. We administered 8 mg/kg of tocilizumab twice at 12 hours of distance from the first dose and maintained the methylprednisolone. In the following days the patient presented progressive improvement of respiratory failure and platelet count. About 3 days after tocilizumab administration, the patient was completely weaned from HFNC and achieved a saturation of 96% with only 2 L per minute of oxygen. Platelet count increased up to 209,000/mm³ in sodium citrate, but remained about 14,000/mm³ in EDTA 10 days after the beginning of methylprednisolone and 6 days after tocilizumab administration. The patient showed great improvement in respiratory parameters and radiological finding as shown in Figure 1. He was transferred to a respiratory rehabilitation ward and continued steroids even if at a lower dosage (methylprednisolone 20 mg/daily). About one week after he developed a thrombocytopenia relapse with a reduction in platelet count of 143,000/mm³ in sodium citrate (EDTA count never exceeded 34,000/mm³).

Discussion

SARS-CoV-2 infection leads to a disease with multiple organ and laboratory involvement. Even if many aspects of COVID-19 are becoming clearer - as research goes on - many others are still unknown. The greatest part of present literature has investigated the acute phase of SARS-CoV-2 infection but poor is known about the post-acute phase. Scarce data are available of post infection period and we are not aware of the possible sequelae. Some authors suggest that the greatest part of hematological and hemostatic alterations (and sometimes the worse outcome) are due to the hyper inflammatory status (sometimes called cytokine storm) related to SARS-CoV-2 infection. On the other hand, it is still unknown how long the effects of

![Figure 1](https://example.com/figure1.jpg)

Figure 1. In this figure we present a high resolution computed tomography (HRCT) of the chest at day 0 (left), showing a severe interstitial COVID-19 pneumonia with multiple and diffuse consolidations. On the right the same HRCT scan after 11 days of therapy.
Even if thrombocytopenia is common among severe COVID-19 patients, it is usually mild and asymptomatic. This case report describes a severe immune thrombocytopenia and a rare case of acquired pseudo thrombocytopenia that, to our knowledge, has not yet been described in COVID-19 patients. Moreover, acquired pseudo thrombocytopenia lasted and the ITP relapsed even after clinical remission and a negative nose swab. This suggests that patients could keep a high risk of COVID-19 complications even after clinical remission and probably after patient dismission, perhaps due to the severe inflammatory state and the cytokine storm typical of these patients. We are aware that clinical research studies are needed to confirm our report; should this be the case, it is reasonable to implement a clinical follow-up after disease resolution, particularly for fragile geriatric patients that are at higher risk of morbidity and mortality related to SARS-CoV-2 infection.

References

6. Panyang Xu, Qi Zhou, Jiancheng Xu. Mechanism of thrombocytopenia in...