Case report

Acute high-risk pulmonary embolism requiring thrombolytic therapy in a COVID-19 pneumonia patient despite intermediate dosing deep vein thromboprophylaxis

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ABSTRACT

Cytokine storm induced by the coronavirus 19 (COVID-19) profoundly activates the coagulation cascade causing venous thromboembolism (VTE). Initial studies from Wuhan, China showed increased incidence of VTE in patients with no standard deep vein thrombosis (DVT) prophylaxis in COVID-19 pneumonia patients. Few have argued for high intensity or intermediate DVT prophylaxis in COVID-19 patients with the incidence of VTE ranging from 16 to 27% despite standard DVT prophylaxis. However, no guideline recommendations presently exist to prescribe augmented DVT prophylaxis in these patients due to lack of evidence although the risk of VTE was clearly demonstrated. While there are ongoing trials to demonstrate the efficacy of intermediate dosing against standard DVT prophylaxis in the prevention of VTE, we present a 36-year-old male admitted with COVID-19 pneumonia who developed acute high-risk pulmonary embolism (PE) requiring emergent thrombolytic therapy despite intermediate dosing DVT prophylaxis.

1. Introduction

Cytokine storm induced by the coronavirus 19 (COVID-19) profoundly activates the coagulation cascade causing venous thromboembolism (VTE) [1]. Tang et al. retrospectively analyzed that patients affected with COVID-19 showed increased incidence of VTE when compared to no standard deep vein thrombosis (DVT) prophylaxis [2]. Few studies argued for high intensity or intermediate DVT prophylaxis in COVID-19 patients with the incidence of VTE ranging from 16 to 27% despite standard DVT prophylaxis [3,4]. Subsequently many institutions created thromboprophylaxis regimes that were described as high intensity or Intermediate dosing to counteract the overwhelming effects of thrombosis in these patients while guideline recommendations were still awaited. However, here we present a 36-year-old male admitted with COVID-19 pneumonia who developed acute high-risk pulmonary embolism (PE) requiring emergent thrombolytic therapy despite intermediate dosing DVT prophylaxis.

A 36-year male airport baggage handler with no significant past medical history was admitted towards the end of April 2020 to our facility with symptoms of fever, cough, headache, and runny nose of 8 days duration. He has no significant past medical history and does not take any medications. In the emergency department, he was noted to be in mild respiratory distress requiring 4 L/minute nasal cannula, respiratory rate of 23/min. He was febrile 39.1 °C degrees centigrade with a blood pressure (BP) of 125/83 mmHg, and heart rate of 103 beats/min. He weighed 84 kg and calculated BMI was 28. His mental status and other systemic examinations were unremarkable. Electrocardiogram showed sinus tachycardia. Chest x-ray showed opacities in the left mid lung zone. White blood cell count was 7.14 × 10⁹, Hb 150 g/L, and Platelet count 201 × 10⁹. He had normal renal and liver functions. He was admitted to the regular nursing floor for further management.

His COVID-19 PCR nasopharyngeal swab test came back positive. C-reactive protein (CRP) on admission was 59 mg/L, ferritin 3122 μg/L, Interleukin-6 698 ng/L and pro-calcitonin 0.15 μg/L. No anti-viral treatments were prescribed in view of his prolonged QTc (460 seconds). He received empiric piperacillin-tazobactam 4.5 g IV every 6 h for possible secondary bacterial pneumonia. His D-dimer was 0.57 μg/mL and fibrinogen 4.53 g/L on admission. He received enoxaparin 40 mg once daily subcutaneously. On Day 3 his oxygen requirements increased to 10–12 L/min on non-rebreathing mask although patient appeared the same clinically from admission. On day 3, his CRP was 116 mg/L, Ferritin 3104 μg/L, Interleukin-6 6719 ng/L and procalcitonin 0.15 μg/L.

Abbreviations: VTE, Venous thromboembolism; COVID-19, Coronavirus disease 2019; PE, Pulmonary embolism.

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Also, his chest x-ray showed bilateral patchy infiltrates which was worse in comparison to the one on admission. He was moved to the intensive care unit (ICU) for closer monitoring. Infectious disease was consulted, and he was prescribed 400 mg IV Tocilizumab on Day 3. His D-dimer on day 2, 3 and 4 were 1.25 μg/mL, 1.86 μg/mL and greater than 4 μg/mL, respectively. He was switched from enoxaparin 40 mg daily subcutaneously to twice daily based on our institution policy of high intensity DVT prophylaxis (intermediate dosing) although no guideline recommendations from any society existed at the time. A Doppler of lower extremities was performed which was normal. Patient was able to prone position himself multiple times during his stay in the ICU. His oxygen requirements gradually came down to 3–4 L/min nasal cannula and he was appropriately moved out of the ICU on day 8. He was ambulatory on the regular floor showing good signs of recovery and improved symptoms. On day 11, patient suddenly felt lightheaded when he got up to go to the toilet. He was in visible respiratory distress, diaphoretic with a respiratory rate of 40/min, heart rate 120/min, oxygen requirements up to 15 L/min. His BP was 92/57 mmhg. Prior to the event his BP was 124/74 mmHg. His electrocardiogram showed sinus tachycardia with no ST deviation. Emergent computed tomography angiogram of the chest with Pulmonary embolism (CTPE) protocol (Fig. 1) showed a saddle embolus with extension to the right and left pulmonary arteries with significant clot burden. There was evidence of right ventricular (RV) strain with RV diameter/left ventricle diameter ratio ~3 (normal < 0.9). He was quickly moved back to the ICU. Cardiac point of care ultrasound showed lack of RV free wall movement with hyperdynamic apex as well septal flattening and bowing to suggest pressure overload in the RV. NT-BNP was 1187 ng/L (normal < 85 ng/L) and troponin 0.011 (normal < 0.010). His Lupus anti-coagulant was mildly positive. Both his Pulmonary Embolism Severity Index (PESI) was Class V (high risk mortality) and simplified PESI score was high. He was considered to have high-risk PE. With no contraindications to thrombolysis and after obtaining consent, he was thrombolysed with alteplase (100 mg over 2 h). No complications ensued. His hemodynamics stabilized as did his oxygen requirements. Repeat formal Echocardiogram showed no evidence of right heart strain or RV dysfunction. He was moved out of the ICU the next day and discharged on room air the following week with a 3-month prescription of apixaban. He was seen in our Tele-outpatient follow up clinic through our hospital network at about 8 weeks. He remained on room air at the time and was back working full time as an airport baggage handler with excellent level of exercise tolerance. Further tests revealed Factor V Leiden, Pro-thrombin gene mutation, anti-thrombin III, Protein C and S were negative. His Lupus anti-coagulant test was now normal as well as his anti-cardiolipin antibody test was negative.

2. Discussion

The pro-coagulant effects and its manifestations in severe COVID-19 patients have been well established [1,5]. Patho-physiologically there appears to be two distinct processes. Firstly, the virus causes direct infection of the endothelial cells causing diffuse inflammation which leads to widespread apoptosis from recruitment of immune-mediated cells. As a result, there is loss of vascular tone and homeostasis creating microvascular dysfunction primarily by a combination of inflammation, vasoconstriction, tissue edema and clot formation, all of which decrease the vessel lumen size and subsequent organ ischemia [5]. Secondly, due to the significantly increased fibrinogen levels there is a higher incidence of large vessel thrombosis such as DVT and PE [1]. Both these processes result in significantly elevated D-dimer levels.

Profound elevation in D-dimer levels in severe COVID-19 pneumonia patients suggest that the body’s own fibrinolytic system is under enormous pressure to break down clot formation that are forming in many organs including the lungs. Therefore, prescribing anti-coagulation for thromboprophylaxis in these patients is paramount. Tang et al. retrospectively reported that prescribing heparin especially low molecular weight heparin (LMWH) provided mortality benefit against patients with no heparin treatment [2]. More specifically, this was observed in patients with D-dimer greater than 3.0 μg/mL when prescribing heparin showed a 20% reduction in mortality (P = 0.017) [2]. However, in a prospective multi-center French study of 150 ICU patients, 16.7% had Pulmonary embolism despite standard prophylactic anti-coagulation

Fig. 1. CTPE study showing saddle embolism in the main pulmonary trunk extending in to both the right and left main pulmonary artery (a) with RV/LV diameter ratio close to 3 (b). Bilateral ground glass patchy opacities seen in the mid (c) and lower zones (d) of the lung suggestive of COVID-19 pneumonia.
[3]. Similarly, another Dutch retrospective study of 184 ICU patients reported a cumulative incidence of VTE of 27% [4]. Objectively, Ranucci et al. demonstrated that clot firmness and clotting time were on the upper limit of normal on viscoelastic testing in a small study of 16 patients in very ill COVID-19 patients [6]. However, at the time and even presently, no guideline recommendations are available on increasing the intensity of DVT prophylaxis in these patients although the risks were clearly demonstrated. Based on the above, our hospital created a policy to provide high intensity or Intermediate dosing DVT risks were clearly demonstrated. Based on the above, our hospital even presently, no guideline recommendations are available on patients in very ill COVID-19 patients [6]. However, at the time and the upper limit of normal on viscoelastic testing in a small study of 16 Ranucci et al. demonstrated that clot firmness and clotting time were on.

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