Title: Granulomatosis with Polyangiitis Mimicking COVID-19 Pneumonia: A Case Report

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Granulomatosis with polyangiitis (GPA), formerly known as Wegener’s granulomatosis, is a necrotizing vasculitis characterized by small to medium-sized vessel involvement and the presence of antineutrophil cytoplasmic antibodies (ANCA). We present a case of a 26-year-old Asian female who was transferred to our center from a nearby hospital, where she presented with shortness of breath, tested positive for COVID-19, and was being managed for COVID-19 pneumonia. She also had hemoptysis, skin lesions, and left foot numbness. Serological markers and VATS-guided lung biopsy confirmed the diagnosis. Treatment with methylprednisolone and rituximab led to stabilization, despite complications of subcutaneous emphysema and lower extremity neuropathic symptoms. Early recognition and appropriate management of GPA are crucial for optimal outcomes.

Keywords: granulomatosis with polyangiitis, ANCA-associated vasculitis, COVID-19

Introduction

Granulomatosis with Polyangiitis (GPA) is a small-medium vessel necrotizing vasculitis and is a component of Anti-Neutrophil Cytoplasmic Antibody (ANCA) Associated Vasculitides. It has a peak incidence at 64 to 75 years of age and is commonly reported in Caucasians without sex predilection. Although almost any organ can be involved in GPA, the upper respiratory tract, lower respiratory tract, and kidneys are commonly affected. The severity of the disease can be heterogeneous. We report a unique case of granulomatosis with polyangiitis (GPA) in a 26-year-old Asian female who was initially transferred from an outside hospital due to shortness of breath presumed to be caused by COVID-19 pneumonia. However, further evaluation led to the diagnosis of GPA highlighting the importance of considering GPA in the differential diagnosis, even in younger individuals and those presenting with respiratory symptoms suggestive of COVID-19.

Case Presentation

A 26-year-old Asian female with a medical history of migraine was transferred to our center from a nearby hospital, where she presented with shortness of breath, tested positive for COVID-19, and was being managed for COVID-19 pneumonia. In the outside center, she was being managed with supplemental oxygen, Remdesivir, Dexamethasone, and Levofloxacin to cover community-acquired pneumonia (CAP). During the
hospitalization, she developed hemoptysis. The tuberculosis workup was negative. She was started on Baricitinib and transferred to our center for specialized care and further workup, given worsening symptoms despite standard treatment.

At the presentation to our center, she complained of chest pain. A review of systems was notable for paresthesia on the dorsal aspect of the left foot. She was tachycardic and required a high-flow nasal cannula to maintain oxygen saturation >92%. Physical examination revealed diffuse bilateral crackles on lung auscultation and nodular non-blanching violaceous skin lesions on bilateral legs, which she attributed to shaving her legs. A patchy area of numbness was appreciated on the dorsal surface of the left leg. Initial laboratory results were significant for leukocytosis, elevated lactate dehydrogenase (LDH), procalcitonin, erythrocyte sedimentation rate (ESR), and C-reactive protein (CRP), as summarized in Table 1. Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) was detected on polymerase chain reaction (PCR). Electrocardiogram (EKG) revealed sinus tachycardia.

Table 1: Initial laboratory values

<table>
<thead>
<tr>
<th>Tests</th>
<th>Result</th>
<th>Reference Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium, serum</td>
<td>137 mmol/L</td>
<td>135-149 mmol/L</td>
</tr>
<tr>
<td>Potassium, serum</td>
<td>4.5 mmol/L</td>
<td>3.4-4.8 mmol/L</td>
</tr>
<tr>
<td>Chloride, serum</td>
<td>99 mmol/L</td>
<td>93-105 mmol/L</td>
</tr>
<tr>
<td>Bicarbonate, serum</td>
<td>27 mmol/L</td>
<td>23-32 mmol/L</td>
</tr>
<tr>
<td>Blood Urea Nitrogen, serum</td>
<td>16.0 mg/dL</td>
<td>7-21 mg/dL</td>
</tr>
<tr>
<td>Creatinine, serum</td>
<td>0.6 mg/dL</td>
<td>0.3-1.1 mg/dL</td>
</tr>
<tr>
<td>White blood count</td>
<td>10.4 k/uL</td>
<td>4.8-10.8 k/uL</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>10.5 g/dL</td>
<td>12-16 g/dL</td>
</tr>
<tr>
<td>Platelets</td>
<td>512 k/uL</td>
<td>150-450 k/uL</td>
</tr>
<tr>
<td>lactate dehydrogenase</td>
<td>482 IU/L</td>
<td>108-199 IU/L</td>
</tr>
<tr>
<td>C-Reactive Protein</td>
<td>13.624 mg/dL</td>
<td>0.0-0.9 mg/dL</td>
</tr>
<tr>
<td>Erythrocyte Sedimentation rate</td>
<td>45 mm/hr</td>
<td>0-20 mm/hr</td>
</tr>
</tbody>
</table>

The patient’s urinalysis showed trace protein, moderate hemoglobin, and urine red blood cell (RBC) count of 25-50 per high power field (HPF). Chest X-ray (CXR) revealed extensive opacity in the right lung and mid to lower lung opacity on the left as shown in Figure 1. Computed Tomography (CT) chest showed dense consolidation in the entire right lung and peripheral infiltrates in the left lung as shown in Figure 2. The patient was admitted to the intensive care unit for COVID-19 treatment. Rheumatology consultation was sought due to hemoptysis, skin lesions, and numbness in the left foot. Laboratory tests revealed an elevated rheumatoid factor, positive cytoplasmic C-ANCA antibody, high C-ANCA titer, and elevated PR-3 antibody, raising suspicion of granulomatosis with polyangiitis (GPA) rather than COVID-19 pneumonia with associated neuropathy and rash.
Figure 1: Chest X-ray showing extensive right lung opacity and left mid to lower lung opacity.
Figure 2: Computed Tomography (CT) chest showing dense consolidation of the entire right lung and peripheral infiltrates in the left lung.

To confirm the diagnosis, a Video-assisted thoracoscopic surgery (VATS)-guided lung biopsy was performed, showing characteristic histological features of ANCA-associated vasculitis, including fibrin and blood filling the alveolar space, inflammatory infiltrate with neutrophils, scattered areas of necrosis, endothelial damage, and necrotizing vasculitis as shown in Figure 3 and Figure 4. Despite the absence of well-formed granulomata, the findings supported the diagnosis of GPA. Treatment with methylprednisolone and rituximab led to medical stabilization, although the patient experienced complications of subcutaneous emphysema and progressive lower extremity neuropathic symptoms.

Following successful stabilization, the patient was discharged home with a walker due to foot drop and was scheduled for outpatient neurology and rheumatology follow-up.
GPA, previously known as Wegener’s granulomatosis, is the most common pulmonary vasculitis and is associated with anti-neutrophil cytoplasmic antibodies (ANCA) to proteinase 4 (PR3). Microscopic polyangiitis (MPA) and eosinophilic granulomatosis, and polyangiitis (Churg Strauss syndrome) are also associated with...
ANCAs, often against myeloperoxidase (MPO), and are grouped within ANCA-associated vasculitis4. GPA is a rare disease with an estimated incidence of 0.4-11.9 cases per 1 million person-years predominantly affecting populations of European descent, while MPA predominates in Asian populations5. The typical age of onset is 45-65 years with no sex predilection. Our patient was a 26-year-old Asian woman, demonstrating that any age group and ethnicity might be affected.

GPA is heterogeneous in its spectrum and severity of presentation. Up to 80% of patients present with prodromal clinical features of systemic inflammation, such as fever, weight loss, fatigue, myalgia, and arthralgia4. Rarely, patients may present more acutely over days. It commonly affects the upper respiratory tract (e.g., sinuses, nose, ears, pharynx, trachea), lower respiratory tract, and kidneys. A varying degree of disseminated vasculitis can occur that can affect any organ. Pulmonary involvement is seen in 50-90% of cases6. It can manifest as cough, dyspnea, and a wide variety of imaging abnormalities such as single or multiple nodules, cavitation, ground-glass opacities, consolidation (reflecting inflammation or alveolar hemorrhage), effusions, stenosis of trachea and bronchi, and rarely fibrosis6,7. Glomerulonephritis occurs in 70-85% of patients, with the characteristic lesion being a segmental focal glomerulonephritis, with rapidly progressive glomerulonephritis (RPGN) seen in fulminant cases4. The onset and course are variable, and ESRD develops in 11-32% of the patients8.

Our patient developed sudden onset of respiratory symptoms without any prodrome, which is uncommon. Imaging revealed dense consolidation of the entire right lung and peripheral infiltrates in the left lung, and she tested positive for SARS-CoV-2. This clinical picture resembled COVID-19 pneumonia very closely. No upper-respiratory tract involvement was seen, and microscopic hematuria was seen, which is non-specific. As a result, she was initially managed with therapy directed towards COVID-19 pneumonia. The lack of improvement with COVID-19-directed treatment, the nodular non-blanching violaceous skin lesions on bilateral legs, and left foot numbness (that eventually progressed to distal sciatic neuropathy) raised suspicion for GPA in this case. Skin involvement can present as palpable purpura, nodules, ulcers, and maculopapular rashes and can correlate with active disease. A biopsy can demonstrate leukocytoclastic vasculitis or granulomatous vasculitis, which are not specific to GPA9. Nervous system involvement is less common but presents most commonly as mononeuritis multiplex, as seen in our case. Other reported manifestations to include cerebral infarction or bleeding, seizures, cranial nerve palsies, altered mental status, meningismus, quadripareisis, or parapareisis10,11.

A biopsy was pursued to confirm the diagnosis in our case. A VATS-guided biopsy of the lung was performed rather than a kidney biopsy, as the yield was likely to be low in the presence of only microscopic hematuria. Histopathological features of GPA described in the literature include necrotizing vasculitis involving venules, arterioles, and capillaries, granulomatous inflammation with/without parenchymal necrosis, micro-abscesses, and fibrosis4. Histopathological examination of the lung specimen of our patient revealed alveolar hemorrhage, micro-abscesses, and necrotizing vasculitis consistent with GPA, even in the absence of granulomas.

Treatment of GPA is divided into the induction phase, to achieve remission in 3 months, and the maintenance phase, to maintain remission. Glucocorticoids, along with either cyclophosphamide or rituximab, are the standard of care for inducing remission in severe diseases, such as in our case12. Rituximab is becoming preferred with growing evidence supporting its efficacy and benefits, such as fertility preservation, superiority in PR3-ANCA positive patients, and relapsing disease5. Thus, rituximab was preferred over cyclophosphamide in our patient with subsequent stabilization of her condition over several days.

Conclusion

This case emphasizes the importance of considering GPA in the differential diagnosis, particularly in younger patients with respiratory symptoms. Early recognition and accurate diagnosis are crucial for appropriate management and improved outcomes in GPA. Further studies are needed to explore the potential association between GPA and COVID-19, as well as to better understand the heterogeneous nature of the disease.

References:


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