CASE BASED DISCUSSIONS

A 26-year-old woman with respiratory decompensation in the immediate postpartum period at Mount Sinai Medical Center in New York City

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Klaus Meinhof and Robert Hiensch

A 26-year-old previously healthy woman developed sudden-onset hypoxaemic respiratory failure 3 days post partum. During pregnancy, the patient had displayed haematuria (26–50 red blood cells/high-power field on urine dipstick) and proteinuria (30 mg/dL on urine dipstick), which at the time was not further investigated. The pregnancy and vaginal delivery were otherwise uncomplicated. On the morning of planned discharge from the maternity ward, the patient developed sudden dyspnoea, tachypnoea, tachycardia and severe hypoxaemia (PaO2 of 48 mm Hg on room air) accompanied by hospitalised hyperventilation and tachycardia severe hypoxaemia. Adequate oxygenation on the mechanical ventilator was difficult to maintain despite frequent endotracheal suctioning, use of high positive end-expiratory pressure and fraction of inspired oxygen (FiO2), initiation of neumuscular blockade and frequent manual recruitment manoeuvres. Chest X-ray (CXR) showed diffuse, predominantly central opacities (figure 1), and a complete blood count demonstrated a decrease in serum haemoglobin from 12.8 g/dL (prepartum) to 7.1 g/dL.

Christian Becker

Respiratory insufficiency in the peripartum period has many potential causes, including PE, amniotic fluid embolism, pulmonary oedema due to tocolytic therapy and peripartum cardiomyopathy. Venous thromboembolism (VTE) is approximately 80 times more common in the first week post partum when compared with the general population1 and PE is a leading cause of maternal mortality in the developed world (whereas postpartum haemorrhage is the most common cause of death in the developing world2). Haemoptysis in the setting of PE occurs in about 5% of patients, and represents pulmonary infarction.3 Given the acute onset of symptoms and echocardiographic findings, PE is the initial concern for this patient.

Klaus Meinhof and Robert Hiensch

Unfractionated heparin infusion was started. Shortly thereafter, the patient endured worsening dyspnoea and coughed up approximately 100 mL of bright red blood. Her heparin infusion was discontinued, and she was emergently intubated for hypoxaemia. Adequate oxygenation on the mechanical ventilator was difficult to maintain despite frequent endotracheal suctioning, use of high positive end-expiratory pressure and fraction of inspired oxygen (FiO2), initiation of neuromuscular blockade and frequent manual recruitment manoeuvres. Chest X-ray (CXR) showed diffuse, predominantly central opacities (figure 1), and a complete blood count demonstrated a decrease in serum haemoglobin from 12.8 g/dL (prepartum) to 7.1 g/dL.

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Emergent bedside flexible bronchoscopy showed diffuse aerial alveolar haemorrhage (DAH). Bronchoscopy with sequential bronchoalveolar lavage should be performed in the area of radiographic opacities. Alveolar haemorrhage is confirmed when serial lavage aliquots become progressively more haemorrhagic. DAH has many potential causes, including infections, inflammatory conditions and medications. Our patient’s history of unexplored haematuria and proteinuria suggests a previously undiagnosed systemic disease, such as the pulmonary–renal syndromes, which present with DAH and focal segmental necrotising glomerulonephritis. In general, treatment of DAH focuses on respiratory stabilisation and reversal of the underlying cause. Immunosuppressive agents are used in immune-mediated disease settings, and can be used empirically in severe illness if there is high clinical suspicion while awaiting the appropriate confirmatory laboratory studies. In this case, the administration of heparin for presumed PE likely exacerbated her DAH. Other causes of DAH in the peripartum period that should be considered include acquired or congenital heart disease including mitral stenosis, malignancies such as metastatic gestational trophoblastic disease, amniotic fluid embolism and various infectious.4–7

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On further examination, the patient had no palpable purpura or other skin rashes associated with vasculitis and no synovitis. Laboratory results were negative for antinuclear antibodies, rheumatoid factor and antiglomerular basement membrane antibodies. Antineutrophil cytoplasmic antibody (ANCA) immunofluorescence, however, was positive at a dilution of 1:320, and ELISA for antimyeloperoxidase antibody was positive at 186 antibody units/mL. Given her previously noted proteinuria and haematuria, a renal biopsy was performed, showing a pauci-immune crescentic glomerulonephritis (figure 2). The patient was diagnosed with microscopic polyangiitis (MPA).

Olivia Ghaw
MPA is an ANCA-associated, nongranulomatous, necrotising vasculitis of the small vessels, with a tropism for the lung and kidney. It typically affects individuals in their fifth and sixth decade of life. Only three cases of MPA have been reported during pregnancy,8–10 and none immediately after parturition. In general, data regarding the effect of pregnancy on the activity of the primary systemic vasculitides are sparse because these syndromes are rare and primarily affect women past their reproductive age. Research on Takayasu’s arteritis, a large-vessel vasculitis that usually affects young women, suggests that pregnancy does not affect disease activity.11

Christian Becker
While the kidney is the most common organ involved in MPA, the lung is another frequent target organ. Pulmonary capillaritis leading to DAH is more commonly seen in MPA than in granulomatosis with polyangiitis (formally Wegener’s granulomatosis),12 13 another ANCA-associated vasculitis. Case series suggest that anywhere from 12%14 to 48%15 of patients with MPA develop DAH during their disease course; this can also predate other extrathoracic manifestations by up to a year.14 Haemoptysis, as seen in our patient, is common in patients with MPA who develop DAH.15 16 For ANCA-associated vasculitis, retrospective data suggest that plasmapheresis improves outcomes in the setting of alveolar haemorrhage.12 The exact role of plasmapheresis in disease induction therapy is the subject of an ongoing trial (http://www.clinicaltrials.gov; PEXIVAS; NCT00987389).

The patient was continued on high-dose prednisone (1 mg/kg) and completed a total of six plasma exchanges. Her gas exchange improved significantly within 48 h of corticosteroid therapy and plasmapheresis, and she was successfully extubated after 6 days of mechanical ventilation. Pharmacological prophylaxis for VTE was not used during her hospitalisation because of her alveolar haemorrhage, though appropriate mechanical VTE prophylaxis was provided.

Olivia Ghaw
For decades, the standard of care for the treatment of ANCA-associated vasculitis was induction with cyclophosphamide. Recent data have shown that rituximab is at least equally effective17 and can also be used as maintenance therapy after induction.18 Seropositivity for antimyeloperoxidase antibodies is associated with lower risk of relapse than antiproteinase-3 antibodies;19 however, lung involvement predicts a higher risk of relapse.20 21

Daniel Bunker
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Daniel Bunker

Once extubated, induction therapy with rituximab was administered (375 mg/m²×4 weekly doses).

Rituximab was chosen because of its favourable side-effect profile, specifically to avoid the risk of cyclophosphamide-induced premature ovarian failure. The excretion of rituximab in breast milk is unknown and thus the patient elected to avoid breast feeding. The patient was discharged in stable condition.

Three weeks later, however, the patient developed acute shortness of breath and orthopnoea. CXR showed marked improvement in her pulmonary infiltrates (figure 3) and transthoracic echocardiogram showed normal biventricular function without a pericardial effusion. CT angiography of the chest revealed filling defects in the lower lobe pulmonary arterial tree, consistent with acute PE. Therapeutic systemic anticoagulation with unfractionated heparin was begun.

Christian Becker

ANCA-associated vasculitis is associated with a markedly increased risk of VTE, especially when the disease is active,22 the patient’s recent ICU stay with neuromuscular blockade and her postpartum state are also risk factors. Given that her current CXR shows resolution of her previous DAH, the benefits of anticoagulation outweigh the risks in this setting.

Klaus Meinhof

The patient’s shortness of breath improved with anticoagulation and she was discharged on warfarin. On follow-up evaluation 3 months later, the patient was doing well with no cardiopulmonary symptoms. Given the high rate of relapse of ANCA-associated vasculitis, maintenance therapy with rituximab is planned. The optimal duration of anticoagulation for VTE that occurs during pregnancy or immediately thereafter is unknown. Generally, the total duration of therapy should be 3–6 months from time of diagnosis for patients with transient risk factors, extending to at least 6 weeks post partum, given the significantly increased risk in the immediate postpartum period.23 24 For patients who have a persistent risk factor, longer duration of therapy is warranted.25 We elected to treat our patient for 6 months with anticoagulation, with close monitoring for further venous thromboembolic events.

Christian Becker

A number of lessons can be drawn from this case:

1. Persistent haematuria and proteinuria during pregnancy should not be dismissed as this may be a sign of a potentially serious underlying disease.
2. One should never be wedded to a diagnosis; although it was treatable empirically for PE in a postpartum patient with acute-onset respiratory failure and haemoptysis, alternate diagnoses, even rare ones such as vasculitis with DAH, should be considered as additional clinical information becomes available or if the patient does not respond to appropriate therapy.
3. The different provider teams should approach each patient encounter with a degree of objectivity; failure to do so when the patient returned to the emergency department after her initial hospitalisation for DAH could have missed the subsequent diagnosis of PE.

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