Thyrotoxicosis Induced Non-Cardiogenic Pulmonary Edema Requiring Extracorporeal Membrane Oxygenation

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Case

A 44-year-old female with DM2, Graves’ disease and Asthma presented to the emergency department with 2-3 days of flu-like symptoms. She was in respiratory distress and initial Vital signs included BP 175/99, HR 130/min, RR 36/min, with 60-70% O2 sat on room air. Initial labs included leukocytosis of 29k/uL and lactic acid of 6.6 mmol/L. Chest X-ray showed diffuse bilateral infiltrates (Figure 1). She was thought to have acute pulmonary edema and treated with high dose furosemide and BiPAP support. Her initial arterial blood gas (ABG) showed a pH 7.3, PCO2 42 mmHg, PO2 82 mmHg, and SaO2 95% while on BiPAP. Over the next two hours she remained persistently tachypneic with a respiratory rate between 36 and 45. Because of concerns of respiratory fatigue, she was intubated and started on mechanical ventilation. Serial ABGs revealed persistent hypoxia, hypercapnia, and respiratory acidosis despite mechanical ventilation (Table I), so thoracic surgery was consulted and started veno-venous ECMO. Because of her rapid worsening despite diuresis, worsening bilateral infiltrates on CXR, and a PaO2/FiO2 ratio of 201 she was thought to have concomitant ARDS and started on Airway Pressure Release Ventilation (APRV) mode. Return of additional tests raised concern for thyroid storm with Thyroid Stimulating Hormone (TSH) < 0.03 mU/L and elevated free T3 >20 ng/dL and free T4 >4.5 ng/dL. Her Burch-Wartofsky score for thyrotoxicosis was 40. Respiratory viral and vasculitis panels were negative and Bronchoalveolar Lavage (BAL) was negative for viral, bacterial, or fungal organisms. Transesophageal echocardiogram showed mild left ventricular hypertrophy and Pro B-type Natriuretic Peptide (Pro-BNP) was elevated at 1288 pg/mL. She was treated with diuresis, Propylthiouracil (PTU), IV steroids, iodine drops, propranolol, and cholestyramine as well as empiric antibiotics for community acquired pneumonia. Her respiratory failure rapidly improved while on ECMO, with improving lactic acid and she was successfully decannulated and extubated on hospital day 7. However, her thyroiditis persisted despite medical management and she ultimately required two rounds of therapeutic plasmapheresis and a thyroidectomy (Figure 2). After a month long hospitalization, plasmapheresis, and thyroidectomy, the patient was successfully discharged to a rehabilitation facility.

Discussion

Graves’ disease is the most common etiology of thyrotoxicosis and thyroid storm is an uncommon, life-threatening complication. Signs and symptoms include fever, tachycardia, tremor, diarrhea, altered mental status, and heart failure. However, it can also cause multiple organ failure. Diagnosis is based on signs and symptoms in the setting of elevated free T4/T3 with suppressed TSH. The Burch and Wartofsky scoring system has been proposed to assist with the diagnosis. A score of >45 is highly suggestive of thyroid storm and a score of <25 makes the diagnosis unlikely. Treatment typically occurs in the ICU and includes beta blockers, thionamide, iodine, and glucocorticoids which decrease new hormone synthesis, inhibit the release of existing hormone, and block the peripheral effects of thyroid hormone. A bile acid sequestrant may also be added in severe cases to reduce enterohepatic recycling of thyroid hormones. In severe cases, studies have shown clinical utility of ECMO for cardiopulmonary support. This disease is associated with significant mortality, with risk factors including: age >60, CNS dysfunction, mechanical ventilation, and non-use of anti-thyroid drugs or beta blockers.

This case has several aspects for discussion. The etiology of the patient’s pulmonary edema, the concern for concomitant ARDS, and the decision to employ ECMO. The patient was initially thought to have acute cardiogenic pulmonary edema from thyrotoxicosis induced cardiac decompensation. However, the echocardiogram, Pro-BNP, and evolution of the case support a diagnosis of non-cardiogenic pulmonary edema due to thyrotoxicosis. ARDS features include diffuse bilateral opacities on imaging, PaO2/FiO2 ratio of 201, and rapid decline in spite of diuresis. The relatively rapid resolution was not consistent with the typical ARDS course with alveolar destruction, and her improvement may have been due to resolution of her overlying non-cardiogenic pulmonary edema. Regarding the use of ECMO, the patient had persistent hypoxia, hypercapnia, and acidosis despite mechanical ventilation. This prompted consultation with cardiothoracic surgery and initiation of veno-venous ECMO.

While there are other plausible etiologies for the patient’s ARDS, the role of thyrotoxicosis merits further discussion. Thyrotoxicosis may contribute to ARDS directly through the...
pulmonary vasculature. Thyroglobulin aggregation in the pulmonary microcirculation was proposed as cause of ARDS in a patient with metastatic papillary cell carcinoma.\textsuperscript{11} Thyrotoxicosis may also affect surfactant production through hormonal effects. Several studies have demonstrated the role of thyroid transcription factor-1 in surfactant protein production.\textsuperscript{12,13} Another possible mechanism is an antibody mediated immune response in the setting of Graves’ Disease. TSH receptors are located in a variety of cells throughout the body including fibroblasts in the lungs.\textsuperscript{14,15} Furthermore, thyroid hormone levels have been shown to be predictive of mortality rates in patients with ARDS.\textsuperscript{16}

In summary, this patient presented with flu-like symptoms, was found to have thyrotoxicosis with non-cardiogenic pulmonary edema and concomitant ARDS requiring intubation and seven days of veno-venous ECMO support. This was an atypical presentation with a wide differential including CHF exacerbation, pulmonary edema due to thyrotoxicosis, community acquired pneumonia, and ARDS. Thyroid storm in this setting presents a diagnostic challenge. It also highlights how thyroid storm is a true endocrine emergency with life-threatening complications, which may include ARDS.

**Data Table I: Serial ABGs**

<table>
<thead>
<tr>
<th>Clinical Setting</th>
<th>pH</th>
<th>PCO2 (mmHg)</th>
<th>PO2 (mmHg)</th>
<th>HCO3- (mEq/L)</th>
<th>SaO2 (%)</th>
<th>FiO2 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-Intubation on BiPAP</td>
<td>7.3</td>
<td>42</td>
<td>82</td>
<td>21</td>
<td>95</td>
<td>0.21</td>
</tr>
<tr>
<td>Post-Intubation</td>
<td>7.04</td>
<td>93</td>
<td>201</td>
<td>27</td>
<td>99</td>
<td>1.0</td>
</tr>
<tr>
<td>2 Hours Post-Intubation</td>
<td>7.02</td>
<td>96</td>
<td>221</td>
<td>27</td>
<td>98</td>
<td>1.0</td>
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<tr>
<td>Post-Cannulation for ECMO</td>
<td>7.14</td>
<td>74</td>
<td>79</td>
<td>27</td>
<td>94</td>
<td>0.7</td>
</tr>
<tr>
<td>3 Hours After Placement on ECMO</td>
<td>7.38</td>
<td>40</td>
<td>60</td>
<td>24</td>
<td>93</td>
<td>0.4</td>
</tr>
</tbody>
</table>

**Figure 1:** Chest X-ray on day of admission showing extensive diffuse airspace opacification with left retrocardiac opacification consistent with pulmonary edema, pulmonary hemorrhage, pneumonia, or ARDS.

**Figure 2:** Graph showing the trend of thyroid hormone levels throughout the hospitalization. Arrows indicate plasmapheresis on hospital days 19 and 24 and thyroidectomy was on hospital day 25.

**REFERENCES**


