COVID-19-associated *Staphylococcus aureus* cavitating pneumonia

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**DESCRIPTION**

A young woman with a background of type 2 diabetes mellitus presented with a 3-day history of shortness of breath and a positive COVID-19 PCR nasopharyngeal swab. She was hypoxic and the chest radiograph revealed bilateral interstitial infiltrates suggestive of COVID-19 pneumonitis (figure 1). A CT pulmonary angiogram ruled out pulmonary embolism and showed ground glass changes without cavitations. On day 1 of hospital admission, she was transferred to the intensive care unit (ICU) due to worsening hypoxia, requiring invasive ventilation 3 days later. The blood cultures taken on days 1 and 5 of admission were negative. After initial improvement, on day 9, she became septic, had increased oxygen requirements and a new inflammatory response (procalcitonin rise from 0.18 μg/L to 63.4 μg/L, C reactive protein 39 mg/L (normal<3.0 mg/L)). She was initially treated for a ventilator-associated pneumonia with piperacillin/tazobactam. The chest radiograph on day 9 showed new bilateral cavitating lung lesions (figure 2). Both blood cultures and endotracheal tube secretions taken on day 10 grew Panton-Valentine leucocidin (PVL)-negative, methicillin-susceptible *Staphylococcus aureus* (MSSA). The antibiotics were initially switched to flucloxacillin 2 g four times a day, and later, due to ongoing clinical deterioration, clindamycin and empirical voriconazole were added to the regimen. Bronchoalveolar lavage samples confirmed MSSA without evidence of tuberculosis or fungal infection (negative mycology cultures, low galactomannan, negative Aspergillus PCR) and serum fungal infection markers (beta-D-glucan, galactomannan) were also low. An extensive respiratory viral PCR panel was performed, which did not detect any concurrent viral infection. A trans-thoracic echocardiogram did not show vegetations. Unfortunately, our patient remained in septic shock despite appropriate antibiotics and maximal supportive therapy and passed away 15 days after admission.

*S. aureus* is a known cause of invasive infections such as bacteraemia and cavitating pneumonia.1 It was a major cause of mortality during the 1918 and the 2009 influenza pandemics.2 3 The host’s immune response is compromised by respiratory viruses, increasing bacterial adhesion to virus-infected cells.4 Increased toxin production is the rationale for including antimicrobials that inhibit toxin synthesis in the treatment of necrotising pneumonitis by *S. aureus*. Bacterial infection seems to be less common in patients with COVID-19 (7%–14%)5 than in those with severe influenza (25%).5 Still, *S. aureus* coinfection causes bacteraemia and pneumonia in patients with COVID-19 (1.2%–1.6%)6 7 and is associated with high 30-day mortality (66.7%).8 Pneumonia was the most common source (19%) in patients with COVID-19 with *S. aureus* bacteraemia with an identified cause.6 Necrotising

**Figure 1** Chest radiograph performed on admission demonstrating bilateral interstitial infiltrates suggestive of COVID-19 pneumonitis. No evidence of cavities were visible on this radiograph.

**Figure 2** Chest radiograph performed 9 days after admission to hospital (6 days after admission to the intensive care unit) demonstrating large bilateral fluid filled cavities in the right and left middle zones.
pneumonia by PVL-producing strains has been described complicating COVID-19. A higher incidence of ICU acquired methicillin-resistant *Staphylococcus aureus* (MRSA) infections were observed in the 2004 SARS outbreak. In our case, the infection was caused by neither a resistant nor a PVL-producing strain but nonetheless caused a very rapidly progressing and destructive pneumonia. We have observed an increased frequency of necrotising/cavitating pneumonias in our patients with COVID-19 in the ICU, as has been described elsewhere. It is therefore important to consider the possibility of infection by *S. aureus* in this cohort of patients.

Consider adding antistaphylococcal antimicrobial agents with good lung penetration and/or antitoxin synthesis effect, for example, linezolid and clindamycin, in the treatment of patients with COVID-19 pneumonia presenting with cavitating pneumonia, particularly if there is rapid progression.

### Patient’s perspective

Unfortunately, we were unable to get an insight into the patient’s perspective in this case as she was critically ill during her admission and subsequently passed away.

### Learning points

- *Staphylococcus aureus* can occur as a secondary bacterial infection in patients infected with COVID-19 and can cause severe cavitating pneumonia as with influenza.
- In the setting of a patient severely ill with COVID-19 pneumonia, *S. aureus* may cause bacteraemia or secondary pneumonia that are associated with high mortality.
- Consider adding antistaphylococcal antimicrobial agents with good lung penetration and/or antitoxin synthesis effect, for example, linezolid and clindamycin, in the treatment of patients with COVID-19 pneumonia presenting with cavitating pneumonia, particularly if there is rapid progression.

### Contributors
SC and CS were involved in writing the manuscript. MA and BC were involved in revising the manuscript. MA helped obtain informed consent from the patient’s next of kin. All authors were involved in the care of the patient.

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### Patient consent for publication
Next of kin consent obtained.

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### REFERENCES


