INTRODUCTION

Nodular or round pneumonia occurs less frequently than lobar or bronchopneumonia. When present it is rapidly progressive and can be fatal. It is commonly sub-pleural in location but can be central. Sub-pleural consolidation occurs due to haematogenous spread of infection i.e. septic embolism. Rapidly enlarging and necrotizing nodular pneumonia is etiologically consistent with Staphylococcal infection; both aureus and non-aureus species. About 2 to 5% of all community acquired pneumonia is attributed to Staphylococcus aureus(SA). Methicillin resistant staphylococcus aureus(MRSA) producing Panton Valentine Leukocidin(PVL) cytotoxin are associated with rapidly necrotizing nature of pneumonia. But Methicillin sensitive Staphylococcus aureus(MSSA) infection can also produce similar features if it produces PVL cytotoxin. Streptococcus pneumoniae, Klebsiella pneumoniae, Actinomycetes, Nocardia, Aspergillus, Legionella, Q fever, M.tuberculosis, are other pathogens producing similar features.

Our patient developed rapidly progressive nodular pneumonia with sputum culture growing SA that necrotized despite antibiotics. It required six weeks of antimicrobials for the adequate response and healing.

CASE REPORT

Here we have a 58 years old thin built male with history of smoking and treated pulmonary tuberculosis 29 years back. He presented with one spike of fever one week prior to the hospital visit, following which he developed chest pain and dry cough. He had exertional breathlessness with few exacerbations in...
the past. He was febrile (99°F) and breathless on presentation with bilateral crackles in chest and oxygen saturation of 86% in room air. Chest radiograph showed bilateral inhomogeneous nodular lesions. Blood counts revealed leucocyte count of 11,700/mm³ with 80% neutrophils, haemoglobin of 10.8 g/dl. Renal functions and electrolytes were normal except urea of 75 g/dl. Serological screening for HIV, HBsAg and Anti-HCV were all nonreactive. After initial stabilization, the possibility of metastatic malignancy of lung was considered. Hence, ELISA for Carcinoembryonic antigen (CEA), Prostate Specific Antigen (PSA) and Carbohydrate Antigen 19-9 (CA 19-9) were evaluated to identify the primary malignancy. CEA was below reference range but PSA was 7.1 ng/ml. (reference value = <4 ng/ml) and CA 19-9 was 50.2 IU/ml (reference range = <35.0 IU/ml). Colonoscopy was normal and abdominal ultrasound showed minimal pelvic collection but no mass lesions. Ceftriaxone and azithromycin were used as initial empirical antibiotics. Gram’s and acid fast staining of sputum were negative. Blood and urine culture sent from emergency failed to grow any organisms. But sputum grew Staphylococcus aureus sensitive to quinolones, cephalosporins, aminoglycosides, tetracyclines and carbapenems but resistant to penicillins. Hence, the antibiotics were later changed to levofloxacin and ceftriaxone. History and examination reviewed at this time revealed an indurated lesion over the nape of neck, which turned out to be an abscess. The cultures from the drained specimen also showed Staphylococcus aureus. The nodular infiltrates cavitated on the review X-ray done after one week. Contrast tomography of Chest was done suspecting cavitating malignancy. Multiple thick walled cavities of varying sizes were noted in bilateral lung field, predominantly in periphery and upper lobes. Few of these showed air fluid level. Right sided hydropneumothorax with loculated fissural collection was also seen. Incidentally CT also picked up isodense peripheral nodular enhancement in right parasternal region extending from C7 to T6. There were multiple sub-centimetric lymphnodes around the carina. Possibility of septic emboli was considered and patient was continued on two weeks of injectable antibiotics: ceftriaxone and levofloxacin. Orally levofloxacin was continued for next four weeks. Chest radiograph at the end of 1 month of therapy showed reduction of size and number of lesions. Then the patient was followed up after 3 months and the CXR only showed emphysematous changes without any nodules or cavities.

**DISCUSSION**

Minority of bacterial pneumonia do not produce predictable history and fails to follow the usual natural course. Necrotizing pneumonia is a form of complicated pneumonia which produces multiple areas of consolidation with rapid necrosis. Such phenomenon can also occur in nodular pneumonia but is uncommon.

Haematogenous and local spread of infection producing metastatic focus is a known complication of Staphylococcal infection. Such complications vary in frequency from 16 to 36%. In a prospectively matched case control study published in 2012, 73 % (84 out of 115) patients with gram positive infection had metastatic septic foci and 19% of the metastases were in lungs (16 out of 84) in the form of lung abscess. In a recent study published in 2012, metastatic foci of Staphylococcal infection were considerably higher in outpatients (67.2 %) as compared to inpatients (32.8%). The metastases were most common in the lung (39 out of 67) and the Methicillin sensitive strains were common in the out patients.

The non-detection of a clinically silent focus of infection is a vital factor during clinical management of metastatic gram positive infections. The other important risk factors for metastases of Staphylococcal and other Gram positive infections are delay in treatment, noncompliance with recommended treatment, patient coming directly from the community, positivity of blood culture after 48 hours of admission and persistent fever after 72 hours of admission and treatment. The regular risk factors predisposing to infection also play crucial role in such phenomenon like alcoholism, old age, diabetes mellitus, malignancy and immune-suppression. Our patient came from community after seven days of initial symptoms with a clinically silent abscess over the nape of the neck detected 10 days after admission. Septic pulmonary embolism (SPE) like in our case, almost always has a predisposing infective focus prior to the development of pulmonary features. Racheal J. Cook et al. retrospectively analysed 14 cases of SPE and all patients had extra-pulmonary foci of infection prior to development of SPE.

Initial radiograph of our patient taken at emergency room was suggestive of multiple bilateral lung metastases of neoplastic origin. The necrosis of all the nodular opacities further hinted towards necrotizing malignancy. Only after the detection of the neck abscess the diagnosis metastatic necrotizing nodular pneumonia was considered in this case. Bilateral nodular or round pneumonia are relatively uncommon and undiagnosed in adults as compared to children. More common presentation is solitary pulmonary nodule. Text books define nodular pneumonia to be rarer than lobar or bronchopneumonia. If the mode of spread is haematogenous the lesions are more likely to be bilateral and sub-pleural. The gram positive organisms are to be suspected in such radiological scenario. And if the lesions necrotize then PVL producing Staphylococcus aureus comes into the differential diagnosis. When the
peripherally located infective nodules necrotize then pleural effusion, pneumothorax, hydro-pneumothorax and empyema complicates the picture due to the rupture of the sub-pleural focus. Retrospectively it was seen that all these clinical events did occur in our patient in sequence. However, there are two limitations to our presumption: we were unable to characterize PVL status of the Staphylococcus aureus and blood cultures were negative to confirm bacteraemia.

PVL: the cyto-toxin can be produced by both MSSA and MRSA and is responsible for rapid progression of pneumonia and necrosis with complications. Hence, PVL status of Staphylococcus needs to be verified irrespective of resistance pattern. PVL producers should receive prolonged therapy (4 to 6 weeks) with sensitive antibiotic. Empirical therapy should include beta-lactam (penicillin, ceftriaxone) or respiratory quinolone (levofloxacin) depending on susceptibility and allergic profile. Adding clindamycin, linezolid or rifampicin to beta lactam is known to suppress toxin release in PVL strains. Infectious Disease Society of America (IDSA) and American Thoracic Society (ATS) guidelines on severe community acquired pneumonia states that CA-MRSA requires vancomycin or linezolid. However, first line antibiotic in adequate dose and duration is sufficient for sensitive strains, as in our case.

Figures

Image 1: Multiple nodules in chest x-ray resembling lung metastasis taken on day one of admission

Image 2: CECT chest showing multiple peripherally located cavities with minimal air fluid levels.

Image 3: Nodules converting into cavities on second week of admission with air fluid levels.
REFERENCES


