Actinobacillus actinomycetemcomitans causing a mediastinal abscess

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We report what may be the first documented case of an anterior mediastinal abscess due to infection by Actinobacillus actinomycetemcomitans. This small, fastidious Gram-negative coccobacillus is a normal component of the oral flora of healthy individuals, but rarely a human pathogen. It was first described in 1912 by Klinger and has since been identified as the sole aetiological agent in at least 52 cases of bacterial endocarditis in North America and Europe and in isolated cases of brain abscess, meningitis, urinary tract infection, pneumonia, thyroid abscess, and soft tissue and wound infections.

Case report

A 50-year-old man presented with a 10-week history of pleuritic interscapular pain and a dull pain in the left upper arm radiating to the praecordium. Physical examination and a chest radiograph showed nothing abnormal initially or at review two and five weeks later. After nine weeks a painless, non-pulsatile, erythematous mass 6 × 4 cm developed on the anterior chest wall around the second and third costal cartilages while he was pulling fence posts from the ground. The patient also noted night sweats, palpitations, and tiredness but no weight loss. On admission to hospital four days later he was afebrile and, apart from the chest wall mass, there were no abnormal physical findings. The chest radiographs (figure) showed a large anterosuperior mediastinal mass. Dental and paranasal sinus radiographs were not obtained. Other findings were: Hb 12.9 g/dl, WBC 19.7 × 10⁹/l (80% polymorphs), ESR 82 mm in one hour. Ultrasonography indicated a cystic mass superior and anterior to the aortic root and a gallium scan showed increased uptake limited to this area. Neither fibroptic bronchoscopy nor mediastinoscopy showed any abnormality. Surgical exploration of the chest wall mass showed a subpleural and anterior mediastinal abscess, which was drained. Biopsy specimens of the abscess wall showed non-specific subacute inflammatory infiltration. There was no evidence of fungal, actinomycotic, or tuberculous infection. Small Gram-negative coccobacilli in pure culture were isolated from pus collected from the abscess at surgery. The organism was identified as Actinobacillus actinomycetemcomitans on the criteria of Cowan and Steel, and was sensitive to ampicillin, erythromycin, cotrimoxazole, and tetracycline and resistant to penicillin and lincomycin. The patient's plasma immunoglobulins and cutaneous delayed responses to common antigens injected intradermally were normal. Treatment was commenced with oral amoxycillin, 1 g six hourly, and probenecid, and was continued for four months, during which time the abscess resolved clinically and radiographically. One year later, at the time of writing, the patient remained well and his chest radiograph showed only mild residual mediastinal widening.

Discussion

This patient made an uneventful recovery from subacute supplicative mediastinitis, and we remain uncertain of the portal of entry of this organism or why it became pathogenic. He showed no evidence of dental disease, oesophageal perforation, granulomatous disease, bronchial obstruction or infection, malignancy, parenchymal lung disease, or immune deficiency. The most likely route of infection was through the anterior chest wall, perhaps because of blunt trauma to the sternum or ribs caused by lifting fence posts, though this is not supported by the long history of chest symptoms before the appearance of the chest wall mass and the absence of radiological evidence of sternal erosion.

Since A actinomycetemcomitans is believed to be a mouth commensal, it has been proposed that the mouth or respiratory tract may serve as a portal of entry, either by direct invasion or via paratracheal lymph nodes. In one study a large proportion of mediastinal abscesses were said to have occurred by direct downward extension along the fascial planes of the retropharyngeal or parapharyngeal abscess. Most of these were situated in the superior mediastinum around the oesophagus and spread of infection to the left side was limited by the aortic arch. The anterior mediastinum was affected very rarely, and when it occurred this was usually due to spread from the posterior mediastinum. In our case, however, the absence of obvious oral, cervical, or endobronchial lesions and the location of the abscess make this route of entry unlikely. Despite the failure to grow the organism from bronchial washings and swabs of the teeth and throat, the isolation of A actinomycetemcomitans in pure culture from pus suggests that this was the sole causative organism in this case.

The rarity of isolation of this organism may be related in part to its capnophilic and slow-growing properties. This report highlights the need for awareness of such growth characteristics in the investigation of unusual infections.
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Posteroanterior and lateral chest radiographs showing a large anterosuperior mediastinal mass (arrows).

and documents a successful outcome from severe infection in a site generally associated with a high mortality rate.

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References

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