A rare case of spinal epidural abscess in a healthy teenage girl without risk factor: case report and literature review

Dr LEE Yee Ting \(^1\)^, Dr YEUNG Kam Tong \(^1\), Dr MAK Wai Kit \(^1\), Dr CHAN Kit Ying \(^1\), Dr ZHU Xian Lun \(^1\), Dr CHAN Tat Ming \(^1\)

\(^1\) Division of neurosurgery, Department of Neurosurgery, Prince of Wales Hospital Hong Kong
\(^2\) Department of Imaging and Interventional Radiology, Prince of Wales Hospital Hong Kong
Abstract

• We report a case of thoracic spinal epidural abscess in a 15 years old girl who was immunocompetent. Our case is unusual in that a spinal epidural abscess developed in a 15-year-old girl, who was immunocompetent, and lacked any predisposing factors. The main objective of this case report is to raise the awareness for rare spinal epidural abscess in immunocompetent young patient.
Background

- Spinal epidural abscess is an uncommon infection, with annual incidence of 5.1 cases per 10,000 admissions. It is a rare condition in children, with average age of presentation at 57.5 +/- 16.6 years (1). Spinal epidural abscess is rare in individuals who are immunocompetent, rare still is the occurrence of spinal epidural abscess in patient lacking any comorbid condition, such as diabetes mellitus, intravenous drug abuse, chronic renal failure, which can be identified in 84% of spinal epidural abscess cases (2).
Case report

• A 15-year-old girl with good past health and normal development presented to Emergency Unit for 2-day history of severe back pain. On the 2nd Emergency Unit attendance, she developed paraplegia and urinary retention. She was admitted and urgent MRI revealed a T8-10 spinal epidural mass with cord compression. (Figure 1)

• Emergency laminectomy for cord decompression was performed. The intra operative specimen growing Methicillin Sensitive Staphylococcus Aureus (MSSA). Post operative MRI was taken, showing no residual lesion and spinal cord was well decompressed. (Figure 2)
Results

• Workup for primary source was negative without dental abscess and valve vegetation. Precipitating factor for invasive MSSA infection was all negative such as IV drug use, HIV, hepatitis B/C and diabetes mellitus. However, screening CT thorax, abdomen & pelvis revealed features of bilateral sternoclavicular joint septic arthritis. (Figure 3)

• Clinically, patient was asymptomatic of the arthritis. She received more than 6-week course of antibiotics with interval MRI confirmed complete resolution of spinal epidural abscess. (Figure 4) Her lower limb power gradually improved. By two months after the operation, she was able to walk with elbow clutches with assistance, with residual right foot drop and clonus.
Discussion

• Spinal epidural abscess in an immunocompetent paediatric patient without risk factors is an extremely rare condition. In a case report and literature review (3), there were only 31 reported cases. Early diagnosis, prompt surgical decompression and appropriate antibiotics is the key treatment for functional recovery. For the current case, the imaging diagnosis of bilateral sternoclavicular joint septic arthritis is suspected to be part of the infection foci.

Conclusion

• Although spinal epidural abscess in an immunocompetent paediatric patient without risk factors is rare, awareness of it with timely diagnosis is imperative to prevent devastating consequence of permanent neurological deficit or even death.

Reference
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