Streptococcus Intermedius Spondylodiscitis: An Unusual Presentation of Acute Back Pain in an Elderly Gentleman With Parkinson’s Disease

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Abstract

Back pain is a common problem affecting approximately one-third of the UK adult population each year with the etiology in older people commonly attributed to osteoporosis and spondylosis. Uncommon causes, such as spondylodiscitis, can potentially cause significant complications, therefore early diagnosis and management are imperative. We present an unusual case of back pain in an elderly gentleman with Parkinson’s disease (PD) attributed to Streptococcus intermedius spondylodiscitis. A 73-year-old gentleman presented with non-specific chest and abdominal pain. Inflammatory markers were elevated, consolidation demonstrated on chest X-ray, and antibiotics were commenced. Patient suffers from PD managed with selegiline, stalevo, and a self-administered apomorphine subcutaneous pump. He developed a constant severe lower back pain without neurological deficit. Streptococcus intermedius was identified on blood culture and common sources were excluded. Magnetic resonance imaging and minimal-preparation computed tomography (CT) confirmed intervertebral spondylodiscitis. A conservative approach with combination antibiotics and rehabilitation was undertaken and the patient returned to his previous functional level. We report an unusual case of spondylodiscitis in an elderly gentleman with Streptococcus intermedius spondylodiscitis.

Keywords: Streptococcus intermedius; Spondylodiscitis; Elderly; Parkinson’s disease

Introduction

Back pain is a common problem affecting approximately one-third of the UK adult population each year [1]. Currently there is no convincing evidence that age affects the prevalence of back pain [2]. Etiology of back pain in older people has been attributed to osteoporosis and spondylosis [3]. However, uncommon causes such as spondylodiscitis can potentially cause significant complications, thereby early diagnosis is imperative. Despite being rare, the incidence of spondylodiscitis is increasing due to effective diagnostic techniques and increased intravenous drug usage, spinal surgery, and ageing population [4]. We present an unusual presentation of back pain in an elderly gentleman with Streptococcus intermedius spondylodiscitis.

Case Report

A 73-year-old man with known Parkinson’s disease (PD) presented with a sudden onset of central chest pain preceded by a 3-week history of upper abdominal discomfort. Following investigations, biliary colic and acute coronary syndrome were excluded. His PD, diagnosed 17 years ago, was managed with selegiline, stalevo, and a self-administered apomorphine subcutaneous pump. Upon admission, he remained febrile and his inflammatory markers were elevated. A chest radiograph demonstrated left basal consolidation and intravenous co-amoxiclav was initially commenced but subsequently switched to intravenous tazocin when blood parameters failed to improve.

During inpatient stay, he continued having intermittent episodes of pyrexia. A previously described back pain, initially attributed to muscular spasm secondary to dyskinesia, had now become severe and constant with lumbar spine tenderness without neurological deficit. Plain films of the spine demonstrated lower thoracic and lumbar scoliosis. A chest radiograph demonstrated left basal consolidation and intravenous co-amoxiclav was initially commenced but subsequently switched to intravenous tazocin when blood parameters failed to improve.

A contrast CT abdomen and pelvis, initially undertaken to investigate an episode of coffee ground vomit, demonstrated severe spinal degenerative changes with mild spondylolisthesis at L4-L5.
Blood cultures obtained during admission had isolated *Streptococcus intermedius*. Following microbiological advice, infective endocarditis and brain abscess were excluded following transthoracic and trans-oesophageal echocardiograms and a CT brain scan. Furthermore, a minimal-preparation CT abdomen and pelvis was performed to exclude liver abscess, which incidentally identified an acute destructive process at T11-T12, highly suggestive of discitis. Subsequent MRI scan demonstrated vertebral body destruction, an epidural collection at T11-T12 and grade 1 spondylolisthesis, in keeping with intervertebral spondylodiscitis (Fig. 1).

Following consultation with our spinal colleagues, a conservative approach was adopted due to the potential risks of surgery in evacuating the epidural collection which was likely to be chronic and not compressing the neural elements. The patient was treated with intravenous meropenem and gentamicin for 6 weeks followed by oral antibiotics for a further 3 months. A thoracolumbar brace was recommended for stabilization of the spine, but was intolerable due to frequent dyskinesia. Within a few weeks, he improved clinically, his inflammatory markers had normalized, and he continued his rehabilitation at a community hospital achieving preadmission functional state.

**Discussion**

Back pain in older people is rarely associated with spondylodiscitis, with the incidence increasing due to spinal surgery, an ageing population, and increased numbers of intravenous drug users and immunocompromised patients [4, 5]. Spondylodiscitis, a spinal infection encompassing spondylitis, discitis, and vertebral osteomyelitis, shares a bimodal age distribution peaking at less than 20 years and those aged 50 - 70 years and is twice more common in men [6]. Our patient was male and between 50 and 70 years of age sharing common demographics described in literature.

Due to the high frequency of back pain, insidious onset, and rarity of the disease, early diagnosis of spondylodiscitis is challenging and often completely missed.

Patients present with non-specific symptoms such as back or neck pain, occasional red flag features, and radicular pain radiating to the chest and abdomen, often leading to misdiagnosis and unnecessary investigations [7]. Initially our patient presented with radicular pain and his lower back pain was initially attributed to dyskinesia. Pyrexia is least commonly elicited by patients with spondylodiscitis, with only half of patients eliciting fever in two studies [6, 8]. However, our patient remained febrile despite being on intravenous tazocin for his chest infection, possibly attributed to his spondylodiscitis.

In pyogenic spondylodiscitis, pathogens commonly enter hematogenously and frequently target the lumbar (58%) and thoracic spine (30%). In our case, spondylodiscitis had targeted the T11-T12 thoracic region. Spondylodiscitis is predominantly caused by *Staphylococcus aureus* [7]. In a study of 1,368 spondylodiscitis patients, the causative agent was identified in 85% of cases, with 48% attributed to Staphylococcus and only 10% to *Streptococcus* [9]. Gouliouris and co-workers reported a 5-20% prevalence of streptococci species (viridans type and β-hemolytic streptococci) in pyogenic spondylodiscitis [7]. These studies demonstrate that streptococcus, the causative organism in our patient, is an uncommon cause of spondylodiscitis.

Common sources of infection for spondylodiscitis in-
clude the genitourinary tract (17%), infective endocarditis (12%), and skin and soft tissue (11%) [8]. Infective endocarditis, brain and liver abscesses were excluded in our patient. Furthermore, the self-administered subcutaneous apomorphine pump was the likely source of infection, augmented by difficult handling during dyskinesia and infrequent replacement of needles.

Elevated inflammatory markers have been reported in the majority of spondylodiscitis cases, with levels normalizing following treatment [10]. Despite intravenous tazocin, inflammatory markers remained persistently elevated in our patient. However, upon alteration to intravenous meropenem and gentamicin, inflammatory markers had normalized within a few weeks.

Due to its high sensitivity (96%), specificity (93%), and accuracy (92%) in delineating anatomical structures, MRI is the modality of choice for identifying spondylodiscitis [8]. The MRI scan was diagnostic of spondylodiscitis in our patient as previous radiological investigations failed to accurately assess the anatomical structures within the spinal canal.

Conclusion

We present an unusual presentation of back pain caused by spondylodiscitis in an elderly gentleman with PD. The patient initially presented with radicular pain followed by low back pain and blood cultures isolated *Streptococcus intermedius*, an unlikely causative agent in pyogenic spondylodiscitis. Furthermore, we propose the likely source of infection to be the subcutaneous apomorphine pump, augmented by difficult handling due to dyskinesia and infrequent replacement of needles during self-administration. These factors highlight the importance of clinical suspicion, early diagnosis and management of patients with spondylodiscitis in order to prevent disease progression and to achieve restoration of functional activity.

Conflict of Interest

The authors declare that there is no conflict of interest.

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References