A ‘crowning’ diagnosis: Fever, neck pain, and diffuse polyarthritis as a fulminant first presentation of pseudogout

Background: Calcium Pyrophosphate Deposition Disease (CPPD) rarely presents as a diffuse polyarthritis with systemic features.

Case presentation: A 75-year-old man with no prior history of rheumatic disease presented with acute-onset diffuse polyarthritis and severe neck pain, accompanied by fevers, leucocytosis, and elevated acute phase reactants. After excluding infectious aetiologies, a synovial fluid aspirate revealed a highly neutrophilic inflammatory infiltrate and rare calcium pyrophosphate dihydrate crystals. Cervical CT confirmed periodontoid calcium deposits consistent with the crowned dens syndrome. The patient was diagnosed with acute CPPD, and his joint and neck pain improved rapidly with colchicine and glucocorticoids.

Conclusion: In addition to arthrocentesis and microscopic crystal analysis, cervical CT can lend support to a diagnosis of pseudogout complicated by CDS. In patients presenting with fever and neck pain, recognition that CDS can be a prominent or isolated feature of acute CPPD facilitates diagnosis, limits unnecessary procedures, and allows for early initiation of appropriate therapy.

Keywords: calcium pyrophosphate deposition disease • pseudo gout • crowned dens syndrome • CT

Abbreviations: CDS: Crowned Dens Syndrome; CPPD: Calcium Pyrophosphate Deposition Disease; CT: Computed Tomography; GCA: Giant Cell Arthritis; PMR: Polymyalgia Rheumatica; RS3PE: Remitting Seronegative Symmetrical Synovitis with Pitting Edema

Introduction

Calcium pyrophosphate dihydrate crystal deposition disease (CPPD), often referred to as pseudogout, is the second most common form of crystalline arthritis. CPPD results from formation of calcium pyrophosphate dihydrate crystals in cartilage, synovium, and soft other tissues, which can trigger local or systemic inflammatory responses [1]. CPPD frequently manifests as episodic acute mono- or oligo arthritis that can mimic gout. While often recognised clinically and radiographically, the gold standard for diagnosis of crystalline arthritis is arthrocentesis with demonstration of positively birefringent calcium pyrophosphate dihydrate crystals under polarized light.

Harrity first reported on the association of CPPD with acute neck pain, later coined the ‘Crowned Dens Syndrome’ (CDS) by Bouvet et al. [2,3] The term CDS has been used to describe both the clinical syndrome arising from inflammation associated with calcium deposits around the odontoid process and the radiographic entity.

Here, we report a case of fulminant polyarthritis complicated by CDS as a rare first presentation of CPPD in an elderly man. Although periodontoid calcification can be observed in asymptomatic patients, identification of CDS by CT was an important aid to the correct diagnosis in this man without prior history of joint disease [4].

Case report

Case presentation

A 75-year-old retired man with hereditary spherocytosis, hypertension, type II diabetes mellitus, and right rotator cuff injury presented with fevers and sudden-onset of diffuse joint pain and swelling.

He was in his usual state of health until three days prior to presentation when he awoke with severe pain in his shoulders, elbows, wrists, hands, knees, and ankles. He noted chills and subjective fever. The following day, the pain spread to include his lower back and neck. He was unable to make a grip or brush his teeth due to pain and stiffness in his hands and shoulders. He presented to a nearby hospital where he was noted to be febrile to 38.2 C. He was empirically started on doxycycline for possible tick-borne illness and discharged home.
Due to persistence of his joint pain, he presented to the emergency department for further evaluation and management. While he endorsed progressive worsening of his diffuse joint pain and swelling to the degree that did no longer allow him to get out of bed, he denied any headache, jaw claudication, visual changes, sore throat, cough, dyspnea, chest pain, abdominal pain, diarrhea, or urinary symptoms. He resided near a wooded area where he regularly walked along the grassy perimeter, but did not recall any tick bites or rashes. He had an extensive travel history. He spent time in Florida from November to May and, the previous summer, he had visited Israel and Italy with his wife. In prior years, he had visited Africa and Asia. He had no history of tuberculosis or malaria. He was up to date in his vaccinations. He had not started any new medications. His family history was significant for coronary artery disease, but not revealing of any autoimmune diseases or malignancies.

On initial examination, he was febrile to 38.3°C (100.9°F), heart rate 103, blood pressure 162/77 mmHg, respiratory rate 16, and SpO2 99%. He was lying in bed without moving his extremities unless prompted. He had conjunctival icterus. He had neck pain/stiffness with flexion or rotational movement; the range of active motion in his neck was significantly limited. His musculoskeletal exam was notable for significant joint tenderness in his shoulders, elbows, wrists, MCPs, PIPs, knees, and ankles. There was synovitis in his elbows, wrists, MCPs, PIPs, and knees bilaterally. There were large bilateral knee effusions and warmth without erythema. There was prominent pitting oedema on the dorsum of both hands.

Investigations

Laboratory findings were notable for a leucocytosis to 13.7 [NR: 4.5-11], microcytic anaemia with Hb 10.1 [NR: 13.5-17.5] and MCV 72.8 [NR: 80-96], and thrombocytopenia to 92 [NR: 150-450] which was chronically low. ESR was elevated to 76 mm/hr [NR: <22] and CRP 15.3 [NR: <3]; BUN was 28 mg/dL [NR: 7 to 20] and creatinine 1.1 mg/dL [NR: 0.5-1.1] with an elevated BUN/Cr ratio of 25; hyperbilirubinemia to 2.3 mg/dL [NR: 0.1 to 1.2], which was unconjugated; other LFTs were normal. His uric acid level was 6.5 mg/dL [NR: 3.4-7].

A chest X-ray revealed subtle right midlung and left base parenchymal opacities suggestive of atelectasis. An X-ray of the hands showed degenerative changes at the first carpometacarpal joints and distal interphalangeal joints. There was no chondrocalcinosis.

Autoimmune serologies obtained by the primary team were notable for negative ANA, rheumatoid factor, and anti-Cyclic Citrullinated Peptide (CCP) antibodies. His infectious disease workup was negative, and his fevers had not improved on initial empirical antibiotic therapy.

A day after his presentation, the man’s case was reviewed by the in-hospital rheumatology consult service. Left knee arthrocentesis was performed to exclude infectious arthritis and evaluate for crystalline arthritis. A CT with contrast of the neck soft tissue was also ordered in the setting of normal hand and wrist radiographs.

Differential diagnosis

The differential diagnosis for this man’s presentation is broad. We considered crystal-induced arthritis, including gout and CPPD. He had no prior history of rheumatic disease or prior episodes of joint pain. Despite his polyarticular disease, there was sparing of the smaller joints of his feet and notably no podagra, which may have supported a diagnosis of gout. While CPPD can present as arthritis and fever, this fulminant presentation of febrile polyarthritis without prior history of joint pain did not quite fit the typical picture of an acute pseudogout attack. Viral polyarthritis, such as parvovirus B19 and chikungunya (home in Florida, where there was an outbreak in 2014; but no rashes were present), could not yet be excluded from the differential. Other possible aetiology included Polymyalgia Rheumatica (PMR), Giant Cell Arteritis (GCA), endocarditis, and less likely septic arthritis pending arthrocentesis. Lyme disease was also considered unlikely, given lack of a rash, recollection of tick bite, response to doxycycline, and negative Lyme antibodies. Remitting Seronegative Symmetrical Synovitis with Pitting Edema (RS3PE) syndrome, a rare form of at times panarticular tenosynovitis that is more prone to occur in elderly men, was also considered as a possible diagnosis. Adult-onset Still’s disease is exceedingly uncommon in his age group, and he lack additional clinical features.
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that would have prompted consideration of this diagnosis (Table 1).

Diagnosis

Arthrocentesis showed 14,272 leukocytes [non-septic inflammatory range: 2,000-50,000] with a PMN predominance (86%), a lower count than would be expected with septic arthritis. Rare positively birefringent, rhomboid-shaped crystals consistent with calcium pyrophosphate were observed by polarized light microscopy of centrifuged synovial fluid.

CT neck revealed calcific radiodensities at the craniovertebral junction with mild pannus and erosion of the posterior surface of the odontoid process, consistent with CDS. These findings confirmed the diagnosis of acute CPPD (Figures 1 and 2).

Management

The patient was started on colchicine and continued on indomethacin, resulting in some improvement of his pain within 24 hours. Due to the substantial burden of his arthritis and negative blood and synovial fluid cultures, he was given a single dose of methylprednisolone 125 mg IV to gain better control of his symptoms. Within the next 24 hours, he noticed a dramatic improvement in his range of motion and pain which continued over the subsequent days. He was able to get out of bed and started ambulating with assistance.

One week after presentation, he felt able to perform his normal ADLs. He was discharged on a colchicine taper and as-needed prescription for future flares. He received home physical therapy and was advised to use a roller for a few weeks at home. He was discouraged from driving until any residual stiffness in his neck fully resolved.

Discussion

CPPD appears to affect about 1 per 1000 individuals and is primarily a disease of the elderly (mean age at diagnosis of 72). The disease has the potential to mimic several forms of inflammatory arthritis, with at least six distinct presentations: acute CPPD or pseudogout, chronic CPPD (pseudo-osteoarthritis/ pseudo-rheumatoid arthritis), pseudo-polymyalgia rheumatica, and pseudo-neuropathic arthropathy, and most commonly asymptomatic chondrocalcinosis [5].

Acute arthritis and fever is not an uncommon presentation of CPPD. However, fulminant presentations characterized by acute polyarthritis involving nearly every peripheral joint and the axial skeleton in a patient with no prior history of rheumatic disease is rare.

This patient’s course highlights two instructive clinical issues. Firstly, pseudogout can mimic several diseases, including autoimmune and infectious forms of polyarthritis, meningitis, vasculitis such as giant cell arteritis, and malignancy. In the absence of a prior history of joint disease, the differential diagnosis can be broad [1,6,7]. However, the sudden onset and severity of his joint symptoms, neck pain, and absence of clinical response to empiric antibiotic therapy narrowed the differential diagnosis to suggest a crystalline arthritis or acute viral arthritis.

Table 1. Differential Diagnosis of Fever, Neck Pain & Diffuse Polyarthritis.

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<th>Aetiologies</th>
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<th>Autoinflammatory</th>
<th>Other</th>
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<td>Lyme Disease</td>
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<td>Chikungunya Virus</td>
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Secondly, acute neck pain, particularly in the setting of pseudogout or a suspected pseudogout, should raise clinical suspicion for CDS [4,8]. Recognition of these clinical patterns and a thoughtful approach to workup can help avoid unnecessary investigations (MRI), invasive procedures (e.g., lumbar punctures, temporal artery biopsy), and limit exposure to ineffective treatments (e.g., antibiotics, opioids) [7]. On initial presentation, the patient declined a lumbar puncture that was offered to exclude meningitis. In this patient with few risk factors (e.g., obesity and type 2 diabetes) and otherwise no radiographic evidence of CPPD, CT neck imaging was used to support the clinical suspicion of CDS.

Compared with gout, pseudogout attacks may take longer to reach peak intensity5. Similar to gout, CPPD can manifest with systemic features such as fevers (50%), leucocytosis, and elevated acute phase reactants, but typically has a more insidious onset. Pseudogout attack are typically characterized by mono- or oligoarthritis, with joint pain, tenderness, swelling, and erythema affecting the larger joints such as the knees (>50%), shoulders (35-48%), and ankles (32-35%) [7]. Podagra, which is a common feature of gout, is much less common in CPPD (<20%) [5,7].

While few studies have addressed symptom prevalence in CPPD generally, even fewer have focused on the symptoms at initial diagnosis specifically [1,6,7]. This is a limitation in the scope of literature to date, and there is a need for large-scale investigations into an increasingly prevalent condition among an aging population.

Differentiation from gout or septic arthritis requires arthrocentesis and synovial fluid analysis for cell count with differential, Gram stain, culture, and crystals. Pseudogout is confirmed by the demonstration of CPPD crystals, which often have the appearance of rhomboid-shaped or rod-like structures that exhibit weakly positive or no birefringence by compensated polarized light microscopy [9].

In one study, the prevalence of periodontoid calcification in patients with pseudogout of the peripheral joints (63.4%) or acute neck pain (63.6%) was higher than that in the general population without neck pain (13.5%). The prevalence of calcification increased significantly with age. These results were consistent with those of several previous studies that found pseudogout to be strongly associated with the presence of periodontoid calcification [10].

CDS can often be misdiagnosed as meningitis, epidural abscess, polymyalgia rheumatica, giant cell arteritis, rheumatoid arthritis, cervical spondylitis or even metastatic spinal tumour [9]. To avoid unnecessary invasive procedures and treatment for acute neck pain, cervical CT should be considered in patients presenting with symptoms suggestive of peripheral pseudogout or acute neck pain in the elderly. In addition, the recognition that periodontoid calcification has higher frequency in pseudogout compared to asymptomatic controls suggests that a CT scan can increase diagnostic certainty when the diagnosis is unclear [4].

**Conclusion**

We report a case of fulminant CPPD complicated by CDS as a first presentation of pseudogout in an elderly man. This case highlights the variability in clinical presentation of a relatively common disease. In conjunction with arthrocentesis, CT imaging of the neck can increase diagnostic certainty in cases where there is clinical suspicion for CDS, thus avoiding unnecessary procedures, prolonged hospitalization, and exposure to antibiotic therapy.

**Availability of data and materials**

Data sharing is not applicable to this article since no datasets were generated or analyzed during this study.

**Ethics approval and consent to participate**

This case study did not require ethics approval.

**Consent for publication**

The patient gave written consent for publication of this report.

**Competing interests**

The authors declare that they have no competing interests.

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**References**


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