Incomplete primary hypertrophic osteoarthropathy

Avijeet Prasad, Pratyush Shahi ♦, Apoorv Sehgal, Manoj Bhagirathi Mallikarjunaswamy

Department of Orthopaedics, University College of Medical Sciences, New Delhi, Delhi, India

Correspondence to Dr Pratyush Shahi; pratyushshahi@gmail.com

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DESCRIPTION

A 63-year-old man presented with chronic multiple joint pain and swelling. The symptoms had started in adolescence, progressed for about 10–15 years, and then became constant. There was no associated morning stiffness. Symptoms suggesting any chest or abdominal disease were absent. Family history was unremarkable. Physical examination revealed clubbing and bulbous swelling of all fingers and toes, and enlargement of the distal third of both forearms and legs (figure 1). Motion at all joints was mildly restricted. There were no skin changes. Cardiovascular, respiratory, and abdominal examinations were normal. Radiographs showed extensive periosteal changes in the long bones of bilateral forearms and legs, and small bones of hands and feet (figure 2). Chest X-ray, ECG, echocardiogram, and abdominal ultrasonography were normal. All laboratory investigations were normal. A diagnosis of incomplete primary hypertrophic osteoarthropathy (PHOA), which is an extremely rare condition, was made. The patient was prescribed etoricoxib once daily and physical therapy was advised. At 1-year follow-up, the pain had subsided and motion at the joints had improved.

Hypertrophic osteoarthropathy (HOA) is a syndrome of digital clubbing, periosteal new bone formation, and arthritis. It is of two types: (a) secondary HOA associated with a primary underlying disease, and (b) PHOA or idiopathic HOA. PHOA, constituting 3%–5% of all hypertrophic osteoarthropathies, is an autosomal dominant or recessive condition thought to be caused due to impaired metabolism of prostaglandin E2 (PGE2). It is also known as pachydermoperiostosis due to the coarse facial features resulting from thickening and furrowing of skin (pachyderma) and has three clinical variants: (a) complete form, with both pachyderma and periostosis, (b) incomplete form, with periostosis but lacking pachyderma (as seen in our patient) and (c) fruste form, with prominent skin changes but minimal periostosis. It is important to differentiate between the primary and secondary forms as the latter is associated with cardiopulmonary and abdominal diseases and malignancies. In the primary type, there is generally an insidious onset during adolescence and gradual progression until it becomes stationary, there is familial involvement in about 60% of the cases, and the periosteal changes involve the diaphyseal and epiphyseal regions. In contrast, the secondary type has an acute onset and rapid progression along with a waxing-and-waning nature depending on the activity of the aetiologic lesion, is non-hereditary and the epiphyseal region is typically spared. Other similar conditions that can cause confusion include acromegaly, thyroid acropachy, rheumatoid arthritis, multifocal periostitis, osteitis deformans and osteopetrosis. Non-steroidal anti-inflammatory drugs relieve pain and swelling of joints by blocking PGE2 synthesis. The use of bisphosphonates and glucocorticoids has also been proposed. Although patients with PHOA have a good prognosis with a normal lifespan, they need to be under regular and long follow-up as complications like ptosis, deafness, kyphosis, osteonecrosis and carpal tunnel syndrome can arise as a result of increased soft tissue bulk and hyperostosis.
Images in...

Patient’s perspective

I started having these symptoms when I was a teenager. My nails started changing shape and my joints swelled. The symptoms increased until I was about 30 years old, then they came to a standstill. Although I had pain, I never really bothered to get proper treatment as I could do most of my activities. When I visited this hospital, the doctors worked me up and explained to me the nature of the disease. My symptoms have subsided to a large extent on analgesics and physical therapy. I am aware that I have to visit the hospital regularly for follow-up.

Learning points

► Primary hypertrophic osteoarthropathy (PHOA) is a rare condition and has variable expression.
► It is important to differentiate between the primary and secondary types of hypertrophic osteoarthropathies as the latter is associated with a serious underlying disease.
► Patients with PHOA should be given non-steroidal anti-inflammatory drugs for pain and need to be under regular follow-up for monitoring of possible complications.

Contributors

AP: idea of the article, literature search and involved in patient care. PS and AS: literature search and wrote the article. MBM: literature search and involved in patient care.

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ORCID ID

Pratyush Shahi http://orcid.org/0000-0003-4903-9697

REFERENCES