Primary Hypertrophic Osteoarthropathy
Accompanied by Crohn’s Disease:
A Case Report

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Primary hypertrophic osteoarthropathy is a rare hereditary disease without evidence of underlying diseases. We report a very unusual case of primary HOA accompanied by Crohn’s disease with the primary HOA mimicking secondary HOA, which is a rare manifestation of Crohn’s disease. We also review the literature to find the correlation, if any, between the two.

Key Words: Osteoarthropathy, hypertrophic, Crohn’s disease

Primary hypertrophic osteoarthropathy (HOA, pachydermoperiostosis) is a rare hereditary disease (Martinez-Lavin et al. 1988), characterized by clubbing and hypertrophy of the digits, periosteal new bone formation of tubular bones, and hypertrophic skin changes (pachyderma). Skin manifestations include marked furrowing of the brow, seborrhea and hyperhidrosis (Vogl and Goldfisher, 1962). No underlying diseases have yet been documented. There are just a few reports documenting gastric ulcer, gastric hypertrophy and endocrine abnormalities as accompanying features (Venencie et al. 1988; Matsui et al. 1991).

In contrast, secondary forms of HOA are not uncommon and arise in association with pulmonary diseases such as lung cancer, pulmonary tuberculosis, and chronic obstructive pulmonary diseases. Secondary HOA can be caused by extra-pulmonary diseases such as Crohn’s disease and ulcerative colitis (Neale et al. 1968).

The authors report a case of primary HOA mimicking secondary HOA as a rare manifestation of Crohn’s disease.

CASE REPORT

A 42-year-old man was admitted via our emergency center because of right lower quadrant abdominal pain and diarrhea, which had first been noticed 6 months previously and had become recently aggravated. Twelve years before admission, he had visited our hospital because he noticed a slowly progressive enlargement of his fingers over a 10-year duration without evidence of gastrointestinal trouble at that time. The skin of the hands and face was coarse and clubbing of the digits was noted. Hyperhidrosis, visual impairment, cyanosis, dyspnea, and murmur were absent. No joint tenderness was observed. The following tests were normal at that time: Blood chemistry, serologic test for syphilis (VDRL) and rheumatoid factor, hormonal studies such as T3, T4, fT4, thyroid stimulating hormone, and growth hormone, and he-
matologic studies such as hemoglobin level, red and white cell counts. A postero-anterior radiograph of both hands showed symmetrically thick cortices of phalanges, metacarpals, ulnae and radius including epiphysis (Fig. 1). Soft-tissue thickening of finger tufts was seen. The diaphyses were wider and thicker than the epiphyses. Symmetric, periosteal new bone formations were seen on tibias and fibulas (Fig. 2). No joint abnormalities were found. Two younger brothers and a cousin (Fig. 3) were found to have similar clinical and radiographic features. Chest PA study showed normal appearance. Eight years prior to admission, the patient received an operation (duodenojejunostomy) due to superior mesenteric artery syndrome.

At this admittance, physical examination revealed direct tenderness at the right lower quadrant abdomen. Skin thickening of the face and hands was also noted. Hematologic and blood chemistry data were normal except for 8.8 g/dL of hemoglobin. C-reactive protein, T3, T4, TSH and glycated hemoglobin were not remarkable. Colonoscopy was performed to look for causes of chronic uncontrolled diarrhea and severe abdominal pain. Colonoscopic findings and biopsy results revealed chronic ulcerative colitis with active inflammation and crypt distortion, which were consistent with Crohn’s disease (Fig. 4). A barium study, performed a few days later, showed numerous ulcerations and multiple luminal narrowing of the entire colon (Fig. 5). Eso-
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Fig. 4. A biopsy specimen. The colonic mucosa shows chronic crypt distortion, focal inflammation with chronic inflammatory cells and crypt abscess.

Phagogastroduodenoscopy revealed hypertrophic gastric folds. With sulfasalazine treatment and fluid replacement, gastrointestinal symptoms were relieved. A radiograph of both hands showed no changes of periosteal thickening of metacarpals, phalanges, distal radius and ulnas as compared with his previous admission.

No bowel habit changes suggestive of inflammatory bowel disease were noticed in his relatives.

DISCUSSION

Primary HOA is an inherited disease (Martinez-Lavin et al. 1988), with autosomal dominant transmission with variable expression. One third of previously reported patients have relatives with primary HOA. Any particularly associated diseases have not been described. No gastrointestinal abnormalities have been documented, except for gastric ulcer and hypertrophy, in previous reports (Venencie et al. 1988; Matsui et al. 1991). Our case also showed gastric hypertrophy, but no other diseases which may be related with HOA to define our case as a secondary form.

Histology revealed periosteal proliferation with
round cell infiltration and new bone formation between the periosteum and the original cortex (Vogl and Goldfisher, 1962). Thickened skin showed hyalinization of collagen and elastic fibers with interposed foci of necrosis and inflammation (Vogl and Goldfisher, 1962). Radiography can demonstrate primary HOA, characterized by bilateral, symmetric periostitis of tubular bones. Periosteal new bones showed shaggy, irregular excrescence.

Crohn’s disease is a chronic granulomatous inflammatory bowel disease. The pathogenesis of Crohn’s disease is unknown, although many investigators have studied about immunologic factors, ingestion of exogenous compounds, and genetic linkage (MacDonald, 1993). Crohn’s disease may present osteoarticular manifestations (Pastershank and Tchang, 1972; McAllister et al. 1986). HOA may appear rarely in the course of chronic Crohn’s disease. This secondary form of HOA also presents symmetric periostitis which often has single or laminated appearance (Resnick, 1995). On the basis of radiographic findings, it is well known that differentiation of primary and secondary HOAs is difficult.

The authors believed that the patient of this case report had primary HOA accompanied by Crohn’s disease. Genetic inheritance was evident and no preexisting diseases which may induce secondary HOA. It was also supported by the fact that HOA had been presented 12 years earlier than Crohn’s disease. The association of primary HOA and Crohn’s disease, if it exists at all, has still to be solved.

Although the authors cannot exclude the possibility of the coincidence of primary HOA and Crohn’s disease, both are extremely rare diseases in Asian countries. Here, the authors report a rare case of primary HOA, accompanied by Crohn’s disease metachromatically.

REFERENCES