Case report

Pelvic digit as a rare cause of chronic hip pain and functional impairment: a case report and review of the literature
Marc Maegele

Address: Department of Trauma and Orthopedic Surgery, Intensive Care Unit, University of Witten/Herdecke, Cologne-Merheim Medical Center (CMMC), Ostmerheimerstr 200, D-51109 Cologne, Germany
Email: Marc Maegele - Marc.Maegele@t-online.de

Published: 19 November 2009
Received: 15 March 2008
Accepted: 19 November 2009

Abstract

Introduction: Pelvic digit is a rare congenital anomaly where bone develops in the soft tissue adjacent to normal skeletal bone [1]. The condition is benign and is usually discovered accidentally. On a plain radiography, pelvic digit typically appears as a rib- or phalanx-like bone structure with a clear cortex and medulla related to the pelvis, often with a pseudoarticulation at its base.

Case presentation: We present the case of a 40-year-old Caucasian man who presented with chronic pain and tenderness over his right hip together with functional impairment in abduction and external rotation. Radiology identified a bony protuberance at the right anterior inferior iliac spine with fusion of the proximal bony nucleus to the adjacent bone. The pelvic digit was surgically removed and the patient was discharged free of symptoms and with complete range of motion in his right hip joint.

Conclusion: It is important to recognize and distinguish a pelvic digit from post-traumatic ossification and avulsion to avoid unnecessary additional investigations.

Introduction

Pelvic digit is a rare benign congenital anomaly where bone develops in soft tissues adjacent to normal skeletal bone [1]. The condition is usually asymptomatic and is often discovered accidentally [2]. However, pelvic digit may also cause pain and functional impairment and can cause some confusion in trauma cases, especially if the patient is symptomatic in the area [3]. Radiology characteristically shows a rib- or phalanx-like bone structure with a clear cortex and medulla related to the pelvis, often with a typical pseudoarticulation at its base [2]. We present the case of a 40-year-old Caucasian man who presented with chronic pain and tenderness over his right hip together with functional impairment. Radiology identified a bony protuberance at the right anterior inferior iliac spine with fusion of the proximal bony nucleus to the adjacent bone. The pelvic digit was surgically removed and the patient was discharged free of symptoms with complete range of motion in his right hip joint.

Case presentation

A 40-year-old white Caucasian man presented to our outpatient service complaining of right hip pain. He reported having experienced this discomfort since his youth with the condition now worsening, but he denied any history of trauma. Clinical examination revealed tenderness over his right hip together with functional deficits in abduction and external rotation in his right hip joint. Conventional radiography extended by computed tomography (CT) showed a bony protuberance at the right anterior inferior

Published: 19 November 2009

Journal of Medical Case Reports 2009, 3:139 doi:10.1186/1752-1947-3-139
This article is available from: http://www.jmedicalcasereports.com/content/3/1/139
© 2009 Maegele; licensee BioMed Central Ltd.
This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.
iliac spine with fusion of the proximal bony nucleus to the adjacent bone (Figure 1a and 1b). This finding together with the clinical picture was highly suggestive of a pelvic digit. Due to the patient’s complaints including functional impairment in his right hip joint, surgical removal of the pelvic digit was performed (Figure 1c) and the patient was discharged 3 days later free of symptoms with a full range of motion in his right hip joint. Histopathological work-up was consistent with a rib bone (Figure 1d).

**Discussion**

To date, few single cases and case series of pelvic digits have been reported. For example, Sullivan and Cornwell [4] described a 15-year-old girl with a well-defined ‘rib’ in the pelvis. The abnormal bone curved caudad towards the right side of the distal sacral vertebra but was not directly attached to the sacrum. Histological assessment after removal was consistent with a rib, as in our patient. The authors postulated that the abnormal bone originated embryogenically from the first coccygeal vertebra. A similar observation was made by Lame [5] who reported on the case of a 63-year-old man with a finger-shaped bony structure arising from the right iliac crest, terming this anomaly an ‘iliac rib’. This author located the anomaly to the mesenchymal stage of bone growth before the sixth week of fetal development [6], with the posterior segment originating from a displaced rib center and the anterior counterpart from a displaced sternal center [5]. Morphologically, pelvic digits may present as rib-like [7] and phalanx-like structures with one or more (pseudo-) joints within [5-8]. Intermediate appearances with features of both rib and phalanx have also been reported [9]. The pelvic digit is most frequently attached to the ilium [2,3], but also to the sacrum [4], coccyx [10], abdominal wall [8], and, rarely, to the symphysis [6]. Although pelvic digits occur mostly unilaterally, bilateral occurrences have been reported [6].

There are some variations in the numbers of bony segments and (pseudo-) articulations of pelvic digits described in the literature. For example, Lame [5] and Granieri and Bacarini [7] described a total of six cases, all consisting of a bony structure of at least two bony elements and at least one (pseudo-) articulation. Nguyen et al. [8] reported a case series where one patient had one phalanx and one pseudoarticulation, and two other cases with three bony segments and two pseudoarticulations. A similar configuration was reported by Casey et al. [3].

Similarly, variable origins for the digits have been described. According to some authors, the anomaly can originate from a displaced costal process, a displaced sternal center, or the ossification center at the anterior superior iliac spine [4-6,9]. However, these suggestions do not reflect the sites of attachment in the pelvis, for example, at the coccyx, the pelvic walls, and the inferior abdominal wall. Therefore, it appears more likely that pelvic digits originate from an embryonic mesoderm with rib-forming capacity disposed to these regions [2]. At the end of the third week of embryogenesis, embryonic mesoderm cells with rib-forming capacity migrate from the primitive streak and pass around the cloacal membrane, finding their way from the region of the future coccyx via the region of the future pelvic walls, to the region of the lower abdominal wall [2]. In normal rib development, the posterior part of each rib originates as a ‘costal process’ of the mesenchyma, thus forming the vertebral centra [11]. It has been suggested that, in the pelvis, the ‘costal processes’ become incorporated into the lateral parts of the sacrum and coccyx [2]. The ‘costal process' mesenchyma normally degenerates due to apoptosis. Absence of apoptosis may
allow differentiation of 'costal process' mesenchyma into rib tissue. These bony structures may come into contact with the neighboring developing bone [2].

Differential diagnosis of a pelvic digit comprises myositis ossificans, avulsion, heterotopic bone formation, Fong’s disease and osteochondroma. Fong’s disease (onychoosteo-dysplasia) is a hereditary condition with dysplastic or absent nails and absent or hypoplastic kneecaps (nail-patella syndrome). Other characteristic features include iliac horns and abnormality of the elbows interfering with the full range of motion. In some cases, new bone formation after surgery or ossification of the sacrotuberous ligament can resemble a pelvic digit [2]. The pelvic digit is usually identified via radiography and differentiated from post-traumatic myositis ossificans and heterotopic bone formation by its corticated appearance in the absence of trauma [6,7,12]. Additionally, CT confirms the presence of cortical bone [3,8,13,14].

In the absence of clinical symptoms, surgical intervention is not required [12]. Our patient suffered from chronic pain along with functional impairment in his right hip joint. Diagnostic imaging including conventional radiography and CT revealed a right-sided pelvic digit that prompted surgical intervention. Complete removal restored the full range of motion within his right hip joint together with complete relief of symptoms.

Conclusion
Pelvic digit is a benign congenital anomaly where bone develops in the soft tissue adjacent to normal skeletal bone. It is usually identified via radiography and differentiated from post-traumatic myositis ossificans and heterotopic bone by its corticated appearance in the absence of a traumatic event. Surgical removal is indicated in the case of symptoms such as functional impairment. It is important to recognize and distinguish a pelvic digit from post-traumatic ossification and avulsion to avoid unnecessary investigations.

Abbreviations
CT: computed tomography.

Consent
Written informed consent was obtained from the patient for publication of this case report and any accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal.

Competing interests
The author declares that they have no competing interests.

Authors' contributions
MM assembled all relevant data to this case report, performed the literature review and drafted the manuscript.

Acknowledgements
The author acknowledges the patient for his support and for giving his informed consent for this case report to be published.

References
1. Goyen MG, Barkhausen JB, Marksches NA, Debatin JF: The pelvic digit - a rare developmental anomaly. A case report with CT correlation and a review of literature. Acta Radiol 2000, 41:317-319.
2. McGlone BS, Hamilton S, Fitzgerald Mj: Pelvic digit: an uncommon developmental anomaly. Eur Radiol 2000, 10:89-91.
3. Casey MC, Phancao JP, Pressacco J: Answer to case of the month #106: Pelvic Digit. Can Assoc Radiol J 2006, 57:51-53.
4. Sullivan D, Cornwell WS: Pelvic rib. Report of a case. Radiology 1974, 110:265-267.
5. Lame EL: Case report 32. Skeletal Radiol 1997, 2:47-48.
6. Greenspan A, Norman A: The "pelvic digit": an unusual developmental anomaly. Skeletal Radiol 1996, 25:723-726.
7. Granieri GF, Bacarini L: The pelvic digit: five new examples of an unusual anomaly. Skeletal Radiol 1990, 19:110-112.
8. Nguyen VD, Mattches JD, Wunderlich CC: The pelvic digit: CT correlation and review of the literature. Comput Med Imaging Graph 1990, 14:127-131.
9. Van Derslice R, Gembala R, Jesus J: Case report: pelvic rib/ digit. Spine 1992, 17:1264-1266.
10. Pais MJ, Levine A, Pais SO: Coccygeal ribs: development and appearances in two cases. AJR Am J Roentgenol 1978, 131:164-166.
11. Fitzgerald MJT, Fitzgerald M: Human Embryology London (UK): Balliere Tindall; 1994:25-35, 59-65.
12. Hamilton S: Pelvic digit. Br J Radiol 1985, 58:1010-1011.
13. Greenspan A, Norman A: The pelvic digit. Bull Hosp Jt Dis Orthop Inst 1984, 44:72-75.
14. Hoeffel C, Hoeffel JC, Got I: Bilateral pelvic digits. A case report and review of the literature. Rofo 1993, 158:275-276.