Metastatic calcification as a result of extensive bone metastases in a paediatric patient with parameningeal embryonal rhabdomyosarcoma

F Ismail, MB BCh
Z Lockhat, FFRad (D) SA
N Khan, MB BS, FCRad (D) (SA)
I van de Werke, FRCR
Narisha Adroos, MB ChB

Department of Radiology, University of Pretoria

M Sathekge, MB ChB, MMed (Nuclear Medicine)
Department of Nuclear Medicine, University of Pretoria

M Kruger, MMed Paed, MPhil, PhD
D Reynders, MB ChB, MRCPCH FC (Paed)
Department of Paediatrics, University of Pretoria

Abstract
We present a case of parameningeal embryonal rhabdomyosarcoma with the primary lesion arising in the middle and inner ear, with associated diffuse skeletal metastases and metastatic calcifications.

Introduction
Rhabdomyosarcoma is the third most common extracranial solid malignancy in children. Parameningeal rhabdomyosarcoma arises in sites adjacent to the meninges. Severe visceral metastatic calcification in this patient as a result of extensive bone metastases is described.

Case presentation
A 5-year-old boy presented with a chronic non-productive cough and right hip pain. During the clinical work-up, he was found to have a left pre-auricular mass. His biochemistry test results showed corrected cal-

Figs 1a, 1b and 1c. Axial computed tomography in soft tissue (a), in bony windows (b) and sagittal view (c) demonstrates the left temporal mass eroding the temporal bone, extending anterior to the pre-auricular region. There are lytic lesions involving the inner and outer table of the calvarium and a soft-tissue metastatic mass in the occiput.
cium of 4.19 mmol/l and phosphate of 1.73 mmol/l, which were elevated in comparison with normal reference ranges.

Computed tomography (CT) (Figs 1a, b and c) of the brain showed a large, destructive lesion in the left ear cavity, extending anterior to the pre-auricular region and posterior to the mastoid air cells. There was destruction of the left temporal bone and external ear canal. There were lytic lesions in the calvarium and a soft-tissue mass in the occiput. The radiological diagnosis considered at this stage was metastatic neuroblastoma. CT of the chest, abdomen and pelvis was suggested to locate the primary lesion. The scan demonstrated diffuse lytic bony metastases (Fig. 2), involving all visible bones. The adrenal glands and para-aortic regions were normal, as were plain radiographs and CT of the lungs.

A Tc-99m-methylene diphosphonate (MDP) bone scan revealed diffuse metastatic visceral calcifications on the lungs, myocardium and stomach (Figs 3a and b). These calcifications were not found on any of the other modalities.

Wedge biopsy of the pre-auricular lesion revealed a parameningeal embryonal rhabdomyosarcoma. The primary tumour mass was in the left middle and inner ear. The patient presented at a late stage with diffuse bone metastases. He died a few days after the CT scan.

Discussion

Rhabdomyosarcoma is the third most common extracranial solid malignancy in children. It is a tumour of primitive mesenchymal cells committed to skeletal muscle differentiation which can occur in a variety of organs. There are three main histological types, viz. embryonal, alveolar and botryoid.¹

'Parameningeal' sites are those primary sites which are anatomically adjacent to the meninges. These sites include the middle ear, mastoid, ear canal, oral cavity, paranasal sinuses and infratemporal fossa.¹ ²

Tumours originating in the middle ear and mastoid have a particularly bad prognosis. They are generally considered to be aggressive as they are in close proximity to vital structures, with a tendency to spread intracranially and the potential for meningeal involvement.²

Primary lesions can lead to bone remodelling and destruction,² as was the case with our patient. At presentation, 42% of cases have cervical adenopathy and 58% have distant metastases.³

Metastatic calcification usually results from a deposition of calcified products in otherwise normal tissues as a result of hyperphosphataemia with or without hypercalcaemia. It may affect mainly the lungs, blood vessels, stomach, periarticular soft-tissue, kidneys and myocardium, and to a lesser extent skin and skeletal muscles of extremities.⁴

Metastatic calcification affecting the myocardium and lungs is potentially lethal and is rarely detected before death, because of the absence of specific radiographic abnormalities.⁵ This lethal presentation was noted in our patient, as detected by Tc-99m MDP bone scan, and indeed the patient died soon after diagnosis.
Metastatic calcification may occur in patients with chronic renal failure, primary hyperparathyroidism, extensive bone malignancy, hyperparathyroidism, diffuse myelomatosis, and milk-alkali syndrome. Metastatic calcification may occur in the presence or absence of hypercalcaemia and is therefore not a prerequisite. At autopsy, metastatic calcification is reported to be present in up to 60–80% of patients with chronic renal failure. However, calcification is rarely identified on conventional chest radiographs.

McKay et al. recommend evaluation for bone metastases with Tc-99m MDP bone scan or plain skeletal radiographs, and measuring the serum levels of parathyroid hormone (PTH) and PTH-related peptide, as well as the levels of 25-(OH) vitamin D and 1,25-(OH) vitamin D as this will point to the common causes for hypercalcaemia of malignancy and help to rule out non-malignant causes of hypercalcaemia.

The lung is one of the primary sites of metastatic calcification deposition and, although most patients are asymptomatic, respiratory failure may develop. The deposition of calcium may be diffuse, predominantly lobar, or predominantly apical. The choice of imaging techniques is important in making the diagnosis. A chest radiograph may not be effective in the detection of metastatic calcification. CT – in particular, high-resolution CT – may or may not detect metastatic calcification. When pulmonary abnormalities are present, they most commonly consist of parenchymal opacification or poorly defined infiltrates that may simulate pneumonia or pulmonary oedema. On CT scans, multiple pulmonary nodules with or without calcification and calcifications of the vessels of the chest wall may be seen, more frequently on high-resolution chest CT. Other causes of pulmonary calcification include prior infections (tuberculosis, fungal or varicella), silicosis, sarcoidosis, metastatic tumour, rheumatic mitral valve disease, alveolar microthiasis and idiopathic pulmonary ossification. However, these latter conditions do not usually display calcification of vessels of the chest wall. Furthermore, on CT there is relative lack of interlobular septal thickening, since the predominant sites of calcium deposition seen on pathological examination are the alveolar septa and, to a lesser extent, the pulmonary arterioles and bronchioles.

Detecting pulmonary uptake of bone-seeking radiopharmaceuticals is a valuable adjunct to diagnosis, and it has been suggested that bone scanning be used to screen for occult pulmonary calcifications. The scintigraphic pattern of dense uptake in the lung, frequently associated with stomach and kidney uptake in general, correlates with the distribution of calcification found at autopsy.

In conclusion, the case discussed revealed a parameningeal embryonal rhabdomyosarcoma with diffuse skeletal metastases and visceral metastatic calcifications, and highlights the fact that chest radiographs and CT findings may be nonspecific and that the use of Tc-99m MDP scintigraphy is valuable to the diagnosis.

In patients with hypercalcaemia, it is important to exclude other benign causes for hypercalcaemia, which are more easily treatable as hypercalcaemia of malignancy may be resistant to treatment.

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