Introduction
Paracoccidioidomycosis is a systemic fungal infection, endemic in Latin America, formerly known as South American blastomycosis. It is caused by a dimorphic fungus, Paracoccidioides brasiliensis, which is not uniformly distributed in endemic countries. The mycosis is mostly diagnosed among agricultural workers and/or residents of rural areas in humid tropical and subtropical regions of Latin America.1–5

Paracoccidioidomycosis is classified based on its epidemiological, immunopathologic and clinical aspects into two polar forms: (1) the juvenile type or subacute form, observed mainly in patients under 30 years old, or in immunosuppressed individuals, in whom it presents with lymphatic/haematogenous dissemination and characteristic involvement of the phagocytic macrophage system and; (2) the adult type or chronic form, which affects mainly adults, who have a present or past history of working in a rural area, and is frequently associated with pulmonary and skin lesions.6,7

Paracoccidioidomycosis is a rare disease in childhood (about 6 per cent of all cases)8,9 and usually has multisystem manifestations, often presenting with involvement of the superficial and deep lymph nodes, skin, liver, and spleen. Bone lesions have been less frequently described, and are usually part of a multisystem clinical picture.8–12

The present report describes a case of the juvenile form of paracoccidioidomycosis in a 7-year-old child living in the metropolitan area of Río de Janeiro, Brazil, who sought medical care with complaints primarily related to a bone infection of the right foot which was initially diagnosed as a bacterial osteomyelitis.

Case Description
A 7-year-old girl, born and living in the metropolitan area of Río de Janeiro, was admitted to a municipal hospital with complaints of pain in her right foot and difficulty with walking. According to her mother in November 1999 the child began to complain of pain in her foot. She was seen by an orthopaedic specialist, and prescribed anti-inflammatory drugs. In December 1999, with no improvement in the pain, new onset of fever, and increasing difficulty in walking, she was again seen by the orthopaedist who diagnosed an osteomyelitis, based on an X-ray of the right foot showing an osteolytic lesion of the calcaneous bone (Fig. 1). She was admitted to the paediatric ward (HMSF) and i.v. oxacillin and cefazidime administered to treat presumptive bacterial aetiologic agents, including Staphylococcus aureus and Pseudomonas aeruginosa.

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With no improvement after 10 days, incision and drainage was performed. The aspirated material was tested by microscopic examination and culture for bacteria, all of which were negative. She continued with fever, and a haemogram revealed a leukocytosis and an increased sedimentation rate, at which time oxacillin was stopped and vancomycin added. At this time cutaneous lesions similar to molluscum contagiosum, subcutaneous nodules, and hepatosplenomegaly were noted on physical examination. She was transferred to the University Hospital (HUCFF) of the UFRJ in January 2000. On admission the subject was noted to have anaemia, malnutrition, fever, hepatosplenomegaly, subcutaneous nodules, and cervical lymph node enlargement.

Thoracic and abdominal CT scan demonstrated lymph nodes in the mediastinum and in the intra-abdominal region, as well as hepatosplenomegaly,

**Fig. 1.** Roentenogram of right calcaneous with osteolytic lesion.

**Fig. 2.** CT scan of abdomen with enlargement of lymph nodes and hepatosplenomegaly.
but no pulmonary infiltrates were noted (Fig. 2). Bone scintillography detected lesions in the right calcaneous and multiple areas of the skull.

Haemoglobin was 9.2 g; haematocrit was 27.8 per cent, leukocytes were 37 200 cells, and sedimentation rate was 108 mm. Total serum protein was 8.00 g with an elevated gammaglobulin fraction.

Direct microscopy of potassium hydroxide preparations of sputum showed the typical tissue forms of *P. brasiliensis*. Biopsy of a cervical lymph node also demonstrated *P. brasiliensis* (Figs 3 and 4) in granuloma. Blood cultures were negative for bacteria and fungi.

Treatment with amphotericin B was subsequently initiated and following 1 month of therapy the patient began to gain weight, the lymph node enlargement and liver and spleen enlargement subsided, the subcutaneous nodules disappeared, and she was able to walk again without pain. She was discharged in February 2000 to the Pediatric Infectious Disease outpatient clinic (IPPMG) where she continued to receive i.v. amphotericin B every other day.

**Discussion**

This case illustrates an uncommon presentation of paracoccidioidomycosis: the presentation of a young child from the metropolitan area of the city of Rio de Janeiro without rural exposure, with an isolated bone lesion as the first clinical manifestation of the disease and the presence of the fungus in direct examination of sputum, without evidence of a parenchymal lesion in the lungs.

Paracoccidioidomycosis is a systemic mycosis of significance in Latin America and, its multisystem manifestations may be missed or misdiagnosed as some other infectious diseases. Due to its rarity in childhood, missing, misdiagnosing and delaying the diagnosis is common in paediatric practice. Gonçalves, *et al.* described 36 cases of paracoccidioidomycosis in children in the state of Rio de Janeiro, all of whom were resident in 15 rural counties scattered in the southwestern part of this state. Paracoccidioidomycosis occurs most commonly in adults with a past history of agricultural work, or those living in rural subtropical areas. In a para-coccidioidin skin test survey performed by Wanke in urban Rio de Janeiro, 2 per cent of children less than 12 years old had a positive reaction and 10 per cent of children in the second decade of life had a positive reaction. In contrast, another study performed in a rural area of the state of Rio de Janeiro showed 32.9 per cent positive skin test in children < 10 years old and 46.2 per cent in older children. Thus, although uncommon, infection by *P. brasiliensis* may occur in children living in metropolitan areas of Latin America, as happened with our young patient, reinforcing the fact that paracoccidioidomycosis should be part of the differential diagnosis even in ‘urban dwellers’.

A study of 46 patients with paracoccidioidomycosis found greater clinical involvement of the reticuloendothelial system in younger individuals as compared to adults. According to Amstalden, *et al.* bone and joint infections are relatively uncommon. They described nine cases (mostly in adults) in whom osteoarticular manifestations were the only or...
primary presentation of the disease; seven of these cases included joint involvement, primarily the large joints and long bones of the extremities.

Paracoccidioidomycosis of the bone is described radiographically as circumscribed, well delineated areas of lysis with or without a rim of sclerosis, and can occur in any bone but is usually seen in clavicles, ribs and scapulae. The differential diagnosis of such bone lesions includes chronic bacterial osteomyelitis, tuberculosis, lymphoma and osteosarcoma, and as the present case illustrates, in endemic areas for P. brasiliensis this systemic mycosis should also be included.

The isolated lesion first seen in the calcaneous bone in our patient led her physicians to treat her initially for a bacterial osteomyelitis, and treatment was targeted towards the principal aetiologic agents of this location (S. aureus and P. aeruginosa). If a fungal aetiology had been suspected initially, the material from the bone drainage could have demonstrated the P. brasiliensis tissue forms if adequately examined with potassium hydroxide preparations, as such lesions are usually teeming with organisms. It was only when other clinical features appeared that other possibilities were considered, and subsequent biopsy of the skin nodules and cervical lymph nodes established the correct aetiology. An early diagnosis and rapid initiation of appropriate treatment can achieve an excellent outcome with minimal complications; a wrong diagnosis can result in a fatal outcome.

Restrepo, et al. described three patients without sign or symptoms of lung involvement, with normal chest X-rays, where P. brasiliensis was found in the sputum, similar to our patient. Such an observation reminds us of the pulmonary genesis of this mycosis, proof of the existence of a pulmonary primary infection (clinical or subclinical), even in patients with the juvenile form of the disease where the lung component may be obscured by predominant lymph node involvement.

In summary, paracoccidioidomycosis in children is not commonly described in the medical literature, and many clinicians have little experience with the juvenile form of this infection, which may lead to the mis-diagnosis of many cases. Paediatricians working in P. brasiliensis endemic areas should be aware of this rare but important presentation in children, and note that treatment with amphotericin B is very effective.

References
5. Mangiaterra ML, Giustano GE, Alonso JM, Gorodner JO. Paracoccidioides brasiliensis infection dans une subtropical
CASE REPORT


Checklist for Authors

Originality
Does the study make an original scientific contribution or new observation on the topic?

Usefulness
Are the findings likely to contribute to improved standards of care?
Would the findings have an impact on preventive/promotive care?

Design Features
Is the objective of the study clearly defined?
Is the study design appropriate for the objective?
Are the subjects for the study, their source, the method of recruitment as well as the inclusion/exclusion criteria defined?
Are the sampling methods likely to give rise to bias?
Is there a statement included about sample size?
Is the method for collection of data clearly described?
Are all laboratory methods used clearly referenced?
Are the study and comparison groups similar in all respects except for the topic of inquiry?
Is the response rate satisfactory?
Is the method of data collection likely to be open to bias?

If intervention has been used was the allocation random and blind?
Have the outcome measures been defined?
Are there any drop outs?
Was the method of outcome measurement open to bias?

Analysis and Presentation
Is the statistical procedure employed (including the software used) clearly stated?
Are the statistical tests used relevant?
Do the results adequately answer the research question?
Is the interpretation of results reasonable?

References
Are the references relevant to the study and up to date?
Are the references cited in the style required?

Ethics
Are the design and conduct of the study ethical?
Has the permission of the local ethical committee been sought and received?

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