CASE REPORT

FATAL DISSEMINATED PARACOCCIDIOIDOMYCOSIS IN A TWO-YEAR-OLD CHILD

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SUMMARY

A two year-old female child was admitted at the Pediatric Intensive Care Unit in a septic shock associated with a lymphoproliferative syndrome, with history of fever, adynamia and weight loss during the last two months. On admission, the main clinical and laboratory manifestations were: pallor, jaundice, disseminated enlarged lymph nodes, hepatosplenomegaly, crusted warts on face, anemia, eosinophilia, thrombocytopenia, increased direct and indirect bilirubin, alkaline phosphatase, and gammaglutamyl transpeptidase. A parenteral administration of fluids, dobutamine and mechanical ventilation was started, without improvement of the clinical conditions. A direct examination of exudate collected from cervical lymph node revealed numerous oval-to-around cells with multiple budding, like a “pilot wheel” cell, suggesting Paracoccidioides brasiliensis. Even though treatment with intravenous sulfamethoxazole-trimethoprim was soon started, the child died 36 hours after hospital admission. Disseminated paracoccidioidomycosis was confirmed in the autopsy. This is the youngest case of paracoccidioidomycosis in children reported in the literature.

KEYWORDS: Paracoccidioidomycosis; Paracoccidioides brasiliensis; Lymphoproliferative syndrome; Children.

INTRODUCTION

Paracoccidioidomycosis (PCM) is a fungal disease caused by the dimorphic fungus Paracoccidioides brasiliensis, and is considered the predominant systemic mycosis in Latin America. The geographic region where PCM is found stretches from Mexico to Argentina, mainly in Brazil, where 80% of the cases have been described, mainly in the southeastern and mid-western regions of the country. In comparison with adults, PCM in children is uncommon. According to some studies, only 5% of the described cases are under 14 years, most of them between 10 and 14 years old. Since there are no reports of PCM in children under 3 years, the aim of this study is to report a fatal disseminated PCM in a 2-year-old child.

CASE REPORT

A two-year-old female child, coming from an urban area of Mogi Guacu, São Paulo State, was admitted at the Pediatric Intensive Care Unit/Hospital de Clínicas/Universidade Estadual de Campinas, in septic shock associated to a lymphoproliferative syndrome. She presented a two-month history of intermittent fever, adynamia, weight loss (3 kg), progressive appearance of disseminated enlarged lymph nodes and warts on the face. No respiratory complaints were reported. On admission, she was pale, normothermic, with a mild jaundice, coldness of the extremities, a poor peripheral perfusion, a blood pressure difficult to measure (50 mmHg/30 mmHg), tachycardia (180 beats/min), tachypnea (50 breaths/min), crusted warts on the face, disseminated enlarged lymph nodes (bilateral anterior and posterior cervical, right axillary and bilateral inguinal) hardened, coalescent and without fistulization, enlargement of the liver (6 cm below the right costal margin) and of the spleen (6 cm below the left costal margin). The main laboratory features were anemia (7.1 g/dl hemoglobin), leukocytes count (5,400 cells/mm³, with 20% of eosinophils), thrombocytopenia (55,000 platelets/mm³), increased direct (1.2 mg/dL) and indirect bilirubin (1.1 mg/dL), alkaline phosphatase (1,458 IU/L) and gammaglutamyl transpeptidase (628 IU/L) levels.

She was treated with parenteral infusion of crystalloid and colloidal fluids, dobutamine, mechanical ventilation and undergone hemodynamic monitoring (CVP, MAP, HR and ECG). The direct examination of exudate collected from cervical lymph node drainage revealed numerous oval-to-round cells with multiple budding, like a “pilot wheel” cell, suggesting P. brasiliensis. With the hypothesis of disseminated paracoccidioidomycosis intravenous sulfamethoxazole-trimethoprim (10 mg/kg trimethoprim) was started six hours after admission. Despite these therapeutic procedures, no improvement of the clinical conditions was observed. The child died 36 hours after hospital admission, preventing the complete PCM serological investigation and fungus cultures.
The autopsy histological findings showed the presence of granulomatous lesions containing *P. brasiliensis* in the skin, tongue, epiglottis, several lymph nodes, liver (Fig. 1), spleen and bone marrow. Histological examination of the lungs showed hyaline membranes lining the alveoli, suggesting severe acute lung injury, without presence of fungus. Renal congestion and slightly sparse liver cell necrosis were also observed. No abnormalities were observed in adrenal glands.

**DISCUSSION**

This is the youngest reported case of disseminated form of PCM in children according to the literature reviewed (Medline 1966 - 2003 and LILACS 1982 - 2003).

Clinically, the PCM syndrome in children has been reported as both disseminated and localized forms. Differently from adults, where the lung involvement is common, most children with PCM present reticuloendothelial system involvement, characterized as a febrile lymphoproliferative syndrome. In addition, children older than 10 years tend to present greater involvement of the osteoarticular and tegument systems than children under this age. Younger children are likely to present a more frequent liver and spleen involvement. Gastrointestinal tract, bone marrow, adrenal gland, testis, brain, kidney and heart are occasionally involved. According to the greatest pediatric clinical series of PCM retrospectively reviewed (N = 52, 1944-1974), the lethality rate was 7.7%. Our group have followed, from 1980 to nowadays, 63 children infected with *P. brasiliensis*, including the child herein reported, 93.9% of them with disseminated forms, with a lethality rate of 10% (being submitted, 2003).

While in adults, males are more commonly afflicted than females, at a mean ratio of 10:1, PCM affects children in the same ratio in both sexes. It has been postulated that the female hormones or immunological factors dependent of these hormones could offer a natural protection against this fungal disease.

The non-specific main laboratory changes more frequently reported in children with PCM are anemia, eosinophilia, hypergammaglobulinemia, hypoalbuminemia, increased alkaline phosphatase levels and elevated erythrocyte sedimentation rate. The abnormal liver function tests observed in this case, suggesting cholestasis, could be related to extrinsic obstruction of bile ducts by lymph nodes, liver cell necrosis and the effects of septic inflammatory response. Unfortunately, in this patient, the overwhelming outcome of the septic syndrome prevented the realization of a thorough immunologic investigation, including HIV and PCM serological screens.
The PCM pathogenesis has still not been established. On the other hand, it has been accepted that PCM results from and/or triggers a decrease of the cellular defense mechanisms, increasing the risk of severe systemic infections. However, in this case, it is reasonable to suppose that the septic syndrome was directly related to P. brasiliensis infection, as already described elsewhere. This inference is supported by the negative findings of bacterial pathogens on the blood cultures (two samples) and the autopsy findings of disseminated fungal yeast forms.

The clinical presentation of disseminated histoplasmosis is quite similar to PCM. Although much more rare than PCM in children in Brazil, this etiology should be considered in the differential diagnosis. The morphological aspects of fungus collected from direct examination plus necropsy histological features discharged this diagnosis.

PCM in children is very uncommon under the age of seven. However, this diagnosis should be reminded in children with febrile lymphoproliferative syndrome associated with anemia, eosinophilia and hypergammaglobulinemia, living in endemic areas. In these cases, mainly in the severely ill children, an empirical treatment with intravenous amphotericin B, should be considered while the specific laboratory studies (direct examination, histological studies, cultures and/or serologic tests) are being performed. Since this child was very young in order to consider PCM as the main diagnosis of septic shock upon admission, amphotericin B was not soon started. In addition, we have obtained good results in the treatment of 15 children with severe disseminated forms of PCM, treated exclusively with intravenous sulphamethoxazole-trimetoprin (being submitted, 2003).

RESUMO
Paracoccidioidomycose disseminada fatal em criança de dois anos

Uma menina de dois anos foi internada em Unidade de Terapia Intensiva com o diagnóstico de choque séptico associado a síndrome linfoproliferativa febril, adinâmia e perda de peso nos últimos dois meses. Na admissão, as principais manifestações clínicas e laboratoriais eram: palidez, icterícia, aumento ganglionar em todas as cadeias superficiais, hepatosplenomegalia, lesões cystosomas em face, anemia, eosinofilia, plaquetopenia, elevação de bilirrubina indireta e direta, de fosfatase alcalina e de gama glutamil transferase. A terapêutica instituída foi infusão de fluidos, dobutamina e ventilação mecânica, sem melhora das condições clínicas, seguido da introdução de sulfametoazol-trimetopirina. O exame direto do linfonodo revelou numerosas estruturas ovaladas, com múltiplos brotamentos, como “roda de leme” sugerindo Paracoccidioides brasiliensis. A paciente evoluiu para o óbito 36 horas após a internação. Paracoccidioidomycose disseminada foi confirmada na necropsia. Trata-se do caso mais jovem de paracoccidioidomycose reportado na literatura consultada.

REFERENCES

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