Bilateral acute acquired toxoplasmic retinochoroiditis after steroid therapy for hantavirus pulmonary syndrome: case report

Retinitis toxoplásica bilateral adquirida após terapia com corticoesteróide para síndrome pulmonar por hantavirus: relato de caso

Rubens Camargo Siqueira¹
Rodrigo Jorge²
Luiz Tadeu de Moraes Figueiredo³

ABSTRACT

Description of a case of acute acquired ocular toxoplasmosis following hantavirus pulmonary syndrome. A 41-year-old man presenting hantavirus pulmonary syndrome, confirmed in the laboratory by detection of IgM antibodies to the virus, was submitted to high doses of intravenous corticosteroids for two months. After clinical improvement of hantavirus pulmonary syndrome the patient presented visual loss in both eyes that was secondary to a toxoplasmosis retinitis. The retinitis resolved with anti-toxoplasma therapy. Acquired toxoplasmic retinochoroiditis can occur following steroid therapy for hantavirus pulmonary syndrome.

Keywords: Hantavirus pulmonary syndrome/drug therapy; Chorioretinitis/etiology; Retinitis; Adrenal cortex hormones

INTRODUCTION

In 1993, a previously unknown group of hantaviruses emerged in the Americas as the cause of acute respiratory disease now termed hantavirus pulmonary syndrome (HPS). Hantavirus causing HPS are associated with wild rodent species of the subfamily *Sigmodontinae* and are transmitted to man by contact or through aerosols of excreta and secretions of infected rodents (1-2). For treatment of HPS, it is recommended to hospitalize the patient in an intensive-care unit, and to perform mechanic ventilation for the pulmonary edema and the capillary leaking syndrome in the lungs leading to respiratory failure. The use of high intravenous doses of corticosteroids has been reported as useful treatment for improvement of respiratory failure.

Toxoplasma gondii is a significant cause of infectious retinitis in humans. In 1973, it was stated that systemic symptoms with acute acquired toxoplasmosis caused ocular involvement in 2-3% of the affected patients (3). Theses authors also concluded that only 2-3% of ocular toxoplasmosis was acquired and the majority of cases arose from reactivation of congenital infection. However, a high incidence of acquired infection followed by ocular toxoplasmosis has been observed in southern Brazil suggesting that it might be a more common route of transmission (3-4).

Toxoplasma gondii is a common cause of retinochoroiditis. We report here a patient having bilateral toxoplasmosis retinochoroiditis acquired after use of high doses of corticosteroids for treatment of a hantavirus pulmonary syndrome (HPS).
CASE REPORT

A 41-year-old man presented with bilateral visual loss after a two-month period of steroid therapy for respiratory failure secondary to HPS.

On examination, best-corrected visual acuity was 20/400 in the right eye and 20/200 in the left. His intraocular pressures were normal. A mild cellular reaction in the anterior chamber and vitreous was observed by slit-lamp biomicroscopy bilaterally. Fundoscopy evaluation showed retinitis in the posterior poles of both eyes (Figure 1A,1B). Fluorescein angiography demonstrated early hypofluorescence in the area of retinitis. Late phases showed staining and leakage around borders of the lesion (active lesion) (Figure 2A, 2B).

Laboratory studies including fluorescent treponemal antibody-antibody screen (FTA-ABS), Lyme disease, complete blood count (CBC), erythrocyte sedimentation rate (ESR), HIV screening, tuberculosis skin testing (purified protein derivative - PPD), chest x-ray and three sets of blood cultures were performed with normal results. However, the patient showed serum positive IgM (>0.600) and elevated IgG (>300 IU/ml) anti-Toxoplasma gondii. The patient was treated with pyrimethamine, folinic acid, and sulfadiazine for 5 weeks. After treatment the lesions were healing (Figure 3A, 3B) and his vision improved to 20/200 (right eye) and 20/100 (left eye) within 2 months.

DISCUSSION

This is the first report of acquired toxoplasmosis following HPS. Immunosuppression by use of high doses of corticosteroids, probably, played a main role in the development of ocular toxoplasmosis in this patient.

The role of cellular immunodeficiency in recurrent toxoplasmic retinochoroiditis: has been demonstrated six Cynomolgus monkeys (Macaca fascicularis) with healed toxoplasmic lesions of the retina were immunosuppressed by total lymphoid irradiation and Toxoplasma specimens were able to
proliferate in ocular tissue following the administration of immunosuppressive therapy⁵⁰.

A possible relationship between systemic corticosteroid use and reactivation of ocular toxoplasmosis has been discussed⁶⁰. These authors showed that the disease occurred at intervals of 20 days to approximately 1 year after the start of corticosteroid therapy. Lesions were typical in appearance, course, and manner in which they responded to antimicrobial therapy.

Other authors described a 58-year-old woman who developed ocular toxoplasmosis while receiving long-term systemic corticosteroid therapy allowing correlation of early histopathologic lesions with their clinical counterpart recorded in fundus photographs⁷⁷.

Acquired ocular toxoplasmosis has been linked to blood transfusion⁰⁷, organ donation⁹⁰, breaks in the dermis or contamination of fingers with oocysts⁹⁴, ingestion of oocytes from cat feces⁸⁰-⁸¹, drinking raw milk⁸²-⁸³, consumption of raw eggs⁸⁴-⁸⁵, and ingestion of uncooked or undercooked beef, pork, and lamb⁸⁶-⁸⁷. In one paper, toxoplasmosis was reported in deer and linked to acute illness in humans in three patients.

The case of a 24-year-old man with acquired ocular toxoplasmosis after immunosuppression of tumoral thymic origin has been reported⁹⁹.

An article described a 48-year-old woman with a previous a liver transplant and a cataract extraction who developed a necrotizing retinochoroiditis in her right eye. As the retinochoroiditis progressed, visual acuity worsened to the point of no light perception, and the eye became blind and painful. An enucleation was performed and histologic examination showed *Toxoplasma gondii* necrotizing retinochoroiditis²⁰.

**RESUMO**

Descrição de um caso de toxoplasmose ocular adquirida pós-síndrome pulmonar por hantavírus. Paciente com 41 anos do sexo masculino apresentando síndrome pulmonar por hantavírus, confirmado no laboratório pela detecção de anticorpos IGM para o vírus, foi submetido a altas doses de corticosteróides intravenosos durante dois meses. Após melhora clínica da síndrome pulmonar por hantavírus, apresentou perda visual em ambos os olhos secundária a retinite por toxoplasmose confirmada com sorologia (IGG e IGM positivo). A retinite resolveu após terapia sistêmica específica. A retinite por toxoplasmose pode ocorrer após terapia sistêmica com esteróide para síndrome pulmonar por hantavírus.

**Descritores:** Síndrome pulmonar por hantavírus/quimioterapia; Coriorretinite/etiologia; Retinite; Corticosteróides

**REFERENCES**

8. Siegel SE, Lunde MN, Gelderman AH, Halterman RH, Brown JA, Levine AS,
Bilateral acute acquired toxoplasmic retinochoroiditis after steroid therapy for hantavirus pulmonary syndrome: case report


II Congresso de Residentes e Pós-Graduandos em Oftalmologia da SBO

14 a 17 de Novembro de 2007
Hotel Novo Mundo
RIO DE JANEIRO - RJ

INFORMAÇÕES

Tel.: (21) 2557-7298
Fax: (21) 2205-2240
e-mail: sbo@sboportal.org.br