HTLV-I ASSOCIATED MYELOPATHY IN SALVADOR (NORTHEASTERN BRAZIL)

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SUMMARY — Recent studies of tropical spastic paraparesis have confirmed the existence of human T-cell leukemia virus type-1 (HTLVI) in several tropical areas of the world. In order to determine the role of HTLVI as an etiologic agent of myelopathies in Salvador, we conducted a clinical and serological study in 43 patients with non-traumatic and non-tumoral myelopathies. We found 9 patients with HTLVI associated myelopathy (HAM) which points to a new endemic area of HAM.

KEY WORDS: HTLVI, myelopathy, tropical spastic paraparesis.

METHODS

Sera and CSF of 43 patients admitted in a general hospital in Salvador in a period of 5 months with chronic spastic paraparesis were tested for HTLVI. Nine patients with progressive chronic spastic paraparesis had serum and CSF positive to HTLVI.

Antibodies to HTLVI were detected with a commercially available enzyme immunoassay (EIA). EIA repeatedly reactive samples were further confirmed by a new dot blot confirmatory immunoassay using highly purified HTLVI viral and recombinant proteins as an
Antigen source. Samples were considered positive if antibodies against the both gag (p24) and env (p21E) gene products were present. In CSF samples patterns considered were: cytology (cells/mm³ number and cytomorphological profile), protein and glucose contents (mg/dL), and gamma globulins, participation (%) and distribution on the protein profile. Other specific antibodies to syphilis, cysticercosis, schistosomiasis, toxoplasmosis and HIV were also tested.

RESULTS

All the patients were adult, with ages that range from 18 to 56 years. Two of the patients were promiscuous and one of them had blood transfusion 8 months before symptoms. They had similar histories with progressive weakness, first in one leg and after in the other leg, that was associated with paresthesias of several degrees of intensity. None of the patients had sensory level. All of them had vegetative disturbances that where characterized by bladder dysfunction (manifested by increase urinary urgence or incontinence), constipation and impotence in men.

CSF showed slight increase in the cell number with a range of 4.3 to 25 cells/mm³ with predominance of lymphocytes and monocytes. There was no relevant number of macrophages, eosinophils and neutrophils. The glucose was normal and the protein ranged from 28 to 68 mg/dL. There was a slight increase in gamma-globulin rate in two of them. One patient had infection by HIV and HTLV-1. No other associated parasite or fungus infections were found.

COMMENTS

In recent years, many studies have shown the high prevalence of HTLV-I infection and HAM among different races and high risk groups in several populations in the world. In Brazil, Costa & col. reported 10 patients with clinical evidence of TSP and Cortes & col. concluded that HTLV-I was prevalent in groups at risk for AIDS. Our cases confirm former results of HAM in Brazil and points to an endemic zone of HTLV-I associated myelopathy. Among the patients studied two were promiscuous and only one had received blood transfusion suggesting a vertical source of contamination.

As has been pointed out by Spina-França & col., we also found a slight increase in the cell number as well as in the protein content and gamma globulins rate in the CSF.

Since we have in Salvador a population descendent from black Africans and Ibericans this endemia could result from this migration. However, further studies are necessary to assess this question.

REFERENCES