INTRODUCTION

Progressive multifocal leukoencephalopathy (PMLE) is an AIDS defining disease often arising in HIV (human immunodeficiency virus) disease patients with low CD4 T cell count, and rarely among those with more than 500 CD4 T cell/mm³. Definite diagnosis requires JC Virus (JCV) isolation in cerebrospinal fluid (CSF) or in brain tissue. A 31-year-old man was diagnosed with HIV1, never having any AIDS-defining illness since August’18. Combined antiretroviral therapy was started. In February 2019 right hemiparesis along with cognitive and behaviour disturbances. His CD4 T cell count was of 475/mm³. Brain MRI was done showing asymmetrical demyelinating lesions. Normal cerebrospinal fluid (CSF) findings other than mononuclear pleocytosis. PMLE was diagnosed on the basis of clinical and radiological findings. Treatment should not be delayed when a probable diagnosis exists even if CD4 T cell count is above 500/mm³. Repeated lumbar puncture with JCV determinations should be done in the advent of new/worsen neurological symptoms and evidence of demyelination showed in MRI.

Keywords: Progressive multifocal leukoencephalopathy; JC Virus; AIDS; CD4 T cell count.

We describe a clinical case of an HIV infected man under Anti-retroviral therapy (ART) with a high CD4 T cell count that developed neurological symptoms and characteristic PMLE brain lesions.

CASE REPORT

31 year male HIV patient (since Aug’18) presented in Feb’19 with right hemiparesis along with cognitive and behavour dysfunctions since 3 days and one episode of abnormal body movements. Patient was taking ATT for tubercular pleural effusion since 5 months and was on ART since 4 months.

O/E Vitals were stable and GCS was 10/15. Pupils were 2-3mm, equal in size, sluggishly reacting to light. Right side was hypotonice with diminished reflexes. Neck rigidity and kernig’s sign were present. Rest systemic examination was normal.

Blood profile was unremarkable and he had no evidence of infectious disease elsewhere and no signs of systemic infection were found. Thyroid, renal and hepatic profile was normal and there were no vitamin deficiencies. A lympho- or myeloproliferative disorder and an inflammatory or granulomatosis disease were also ruled out.
CD4 count was 475/mm³. CSF examination revealed normal glucose (0.56 g/dl) and protein (0.33 g/L) levels with 60 leucocytes (42 mononuclear cells).

MRI of brain showed asymmetric T2-FLAIR hyperintense lesions in bilateral parieto-occipital white matter and these lesions were predominantly hypointense on T1W images. The lesions also involved perirolandic region, left cerebral peduncle and right cerebellar hemisphere. There was no midline shift.

The patient was diagnosed as progressive multifocal leukoencephalopathy on the basis of clinical and radiological findings.

**DISCUSSION**

PMLE is an AIDS defining disease often seen in HIV patients with low CD4+ count and rarely among those with higher CD4 count. Patients often present with visual deficits, mental impairment, weakness including hemi or monoparesis. Seizures occur in 20% of patients.

This case report shows how a definite PMLE diagnosis is rare and difficult to achieve in a patient with a high CD4 T cell count and under ARV. It is known that in HIV patients, PMLE often arises in severe immunocompromised states, when CD4 T cell count is lower than 200/mm³. Nevertheless, cases have been described with CD4 T cell count higher than that value but rare. It is well known that JCV cannot infect T lymphocytes or bind to T cell membranes. It seems that JCV uses B lymphocytes as a ‘Trojan horse’ to reach the brain exceeding the blood-brain barrier, and it happens when the reactivation process starts, when an immunosuppression state is achieved, because it can induce the loss of specific immune cells that may allow the beginning of active replication and infection as well as changes in JCV regulatory region [6].

Nowadays, it is not known which ‘level’ of immunodepression is needed in an HIV patient for JCV to turn from a latent to a replicative state, or if there are some other means for JCV to reach the brain and infect oligodendrocytes rather than the ‘B lymphocyte way’.

Sometimes in the course of the disease, PMLE diagnosis was questioned, mainly due to the relatively high CD4 T cell count and in spite of the initial findings in MRI. MRI is the best imaging tool to diagnose PMLE, with typical appearance in 90% of the patients, usually with patchy and commonly non-enhancing white matter lesions, usually bilateral and asymmetric, and with no mass effect. Yet, some demyelinating diseases may present almost with the same characteristics, but some of them with mass effect [7]. HIV-leukoencephalitis is the main differential diagnosis, but it seems that lesions are less diffuse, more symmetric, not visible on T1w and less intense on T2w sequence [8, 9].

**CONCLUSION**

Clinical symptoms and signs together with typical MRI findings among patients with high CD4 T cell count and under ARV should be considered and treated as PMLE. The progression of the disease may correlate with a higher replication of the JCV. Several lumbar punctures with JCV PCR in CSF should be done, but a negative result should not delay the start of antiretroviral treatment in those patients with CD4 T cell count even above 500/mm³.
REFERENCES


