CASE REPORT

Unusual muscle disease in HIV infected patients

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Two patients presented with proximal muscle weakness, a normal or minor elevation of creatine phosphokinase (CPK) and normal findings on electromyography. Muscle biopsy in one patient revealed CD8+ polymyositis, and in the other showed ddi induced myopathy. These cases illustrate the importance of muscle biopsy in identifying the underlying pathology in HIV infected patients with muscle weakness and little or no abnormality in laboratory investigations.

HIV infected patients may develop a variety of muscular disorders including HIV associated polymyositis,1,2 myopathy induced by nucleoside reverse transcriptase inhibitors (NRTI), such as zidovudine,3,4 opportunistic infections including Toxoplasma gondii,5 infiltration by tumour,6 HIV associated vasculitis,7 and rhabdomyolysis caused by HIV itself or by drugs including didanosine.8 Many of these conditions present with overlapping non-specific symptoms of fatigue and myalgia. Biochemical abnormalities of muscle function (creatine phosphokinase (CPK) and aldolase) occur in approximately 15% of asymptomatic HIV infected patients,9 but may not be good markers of muscle inflammation in HIV infected patients undergoing evaluation of possible muscle disease.10-14 In order to define the nature of underlying muscle disease, if any, in a symptomatic HIV infected patient muscle biopsy is necessary. We describe two HIV infected patients with proximal muscle weakness and a normal or minor elevation of CPK and normal electromyography (EMG) findings, in whom muscle biopsy revealed a specific diagnosis.

Case 1
A 50 year old black African woman presented with a 4 week history of leg weakness, with difficulty walking and climbing stairs. She had been diagnosed HIV-1 antibody positive 3 months previously; at which time an eosinophilia of 1.1 (normal range 0.04–0.44) ×10^9/l was noted. On admission to hospital, examination of the legs showed normal tone and proximal weakness (power = MRC grade 4/5). Ankle and knee reflexes were absent and plantars were flexor. The gait was waddling in nature. She had marked difficulty standing from sitting. The thigh muscles were tender to palpation. General examination was normal.

A full blood count (FBC) showed eosinophilia (1.66×10^9/l) and a normochromic, normocytic anaemia: the erythrocyte sedimentation rate (ESR) = 49 mm in the first hour, CPK = 547 (N = 24–173) U/l. Renal, liver, and thyroid function tests were normal, as were calcium and phosphate. The CD4 count was 340 cells ×10^9/l and CD8 count was 3450 (normal = 90–990) cells ×10^9/l; CD4:CD8 ratio = 0.1 (normal = 1.00–2.50). Serological tests for syphilis and an autoimmune screen were negative. Stool microscopy for ova, cysts and parasites, and terminal urine screening for Schistosoma spp ova was negative. Serology for Strongyloides spp, Bartonella spp, Trichinella spp, cysterciosis, Lyme, and Filaria spp was negative.

A plain radiograph of the thighs was normal. Cranial magnetic resonance imaging showed generalised atrophy and widespread leucoencephalopathy. CSF analysis showed protein = 1.72 g/l, glucose = 2.5 (plasma glucose = 5.0) mmol/l, and cell count = 178 lymphocytes/ml (87% were CD8+ lymphocytes); culture for bacteria, mycobacteria, and fungi was negative. Serology for cryptococcal antigen, HTLV-1 and HTLV-2, syphilis, and T gondii was negative. Herpesviruses 1–8, enteroviruses, JC virus and Mycobacterium tuberculosis were not detected by polymerase chain reaction (PCR) amplification.

Nerve conduction studies and EMG were non-diagnostic. Muscle biopsy of left vastus lateralis showed multiple foci of inflammatory cells with occasional infiltration of intact muscle fibres by lymphocytes without granulomata or vasculitis. The inflammatory infiltrate consisted of lymphocytes, macrophages, plasma cells, and eosinophils. Immunohistochemical staining showed the lymphocytes were predominantly CD8+ T cells with smaller numbers of CD4+ T lymphocytes (CD8: C8/144B, Dako, Glostrup, Denmark; CD4: RTU-CD4-1F6, Novacastra, Newcastle, UK) (fig 1) and identified a smaller component of B lymphocytes (CD20: L26, Dako); macrophages expressed CD68 (PG-M1, Immunohistochemical staining demonstrated small numbers of CD4 positive T lymphocytes (C) with larger numbers of lymphocytes expressing CD8 (D). (A) and (B) haematoxylin and eosin; (C) CD4 immunohistochemical staining; (D) CD8 immunohistochemical staining. Bar in (A) represents 50 µm in all panels.

Abbreviations: CPK, creatine phosphokinase; EMG, electromyography; ESR, erythrocyte sedimentation rate; FBC, full blood count; NRTI, nucleoside reverse transcriptase inhibitors.

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Figure 1  Histological examination showed focal endomysial and perivascular inflammation with occasional infiltration of intact fibres by lymphocytes (A). The inflammatory infiltrate was composed predominantly of lymphocytes with additional eosinophils, plasma cells, and macrophages (B, arrows indicate eosinophils). Immunohistochemical staining demonstrated small numbers of CD4 positive T lymphocytes (C) with larger numbers of lymphocytes expressing CD8 (D). (A) and (B) haematoxylin and eosin; (C) CD4 immunohistochemical staining; (D) CD8 immunohistochemical staining. Bar in (A) represents 50 µm in all panels.

The patient began combination antiretroviral therapy with lamivudine, didanosine, and lopinavir and mobilised rapidly. At repeat lumbar puncture, after 2 weeks of antiretroviral therapy, the CSF protein had fallen to 1.1 g/l and the lymphocyte count had fallen to 110 cells ×10⁶/l. After 3 months of antiretroviral therapy the patient was asymptomatic, HIV viral load was undetectable, peripheral blood CD4 count = 310 cells ×10⁶/l, the CD8 count had fallen to 1810 cells ×10⁶/l (CD4:CD8 = 0.17), and CPK and eosinophil count were normal.

**Case 2**

A 51 year old white homosexual man, HIV-1 antibody positive for 10 years, presented with a 2 year history of progressive leg weakness, manifest as difficulty walking and cycling uphill. Six and a half years before the current presentation he had begun combination antiretroviral therapy with zidovudine and lamivudine. This regimen had been modified to didanosine with ritonavir after 3 months, because of marrow toxicity. The patient had taken the latter regimen for over 6 years at the time of presentation with leg weakness. There was no family history of neurological disease. On examination there was weakness of hip flexion (MRC = 4/5) and the thigh muscles were wasted and not tenden: there was no muscle fasciculation. Power, reflexes, sensation, and coordination were otherwise normal in the arms and legs. Gait and cranial nerve examination were normal. General examination was normal.

A FBC, renal, liver, and thyroid function tests, CPK, random glucose, resting venous lactate, autoantibody screen, ESR, and C reactive protein were normal. The CD4 count = 220 cells ×10⁶/l and HIV viral load = 200 copies/ml. An EMG and nerve conduction studies revealed a mild sensorimotor neuropathy (sensory predominance) with no evidence of myopathy. A left vastus lateralis muscle biopsy showed multiple ragged red fibres identified using immunohistochemical staining for mitochondria (ab3298, abcam, Cambridge, UK). There was no evidence of inflammation or fibre necrosis. Ultrastructural examination demonstrated several fibres containing abnormal mitochondria several of which were enlarged with an annular arrangement of the internal membranes while many others contained type I paracrystalline inclusions (fig 2). In the light of the muscle biopsy results didanosine and ritonavir were stopped and abacavir, lamivudine, and efavirenz were started. At follow up 2 months after starting the new antiretroviral drugs the patient was asymptomatic and had an undetectable viral load.

**DISCUSSION**

In both these patients with non-specific presentations, minor abnormalities of CPK and normal results from electromyography, a specific diagnosis was made by muscle biopsy. In the first case the lymphocytic infiltrate of muscle consisted largely of CD8+ lymphocytes, a finding typical of HIV associated polymyositis. The associated CD8+ lymphocytosis in peripheral blood and in CSF, despite the absence of sicca symptoms or parotid enlargement, suggests that this patient had diffuse infiltrative lymphocytosis syndrome (DILS). In one case series of 35 patients with DILS nine (26%) had an inflammatory myositis: in these nine patients the lymphocytic muscle infiltrate was not phenotyped. By extrapolation from other studies of HIV associated polymyositis, it is likely that these patients had a CD8+ lymphocytic infiltrate of muscle. The case described here is also unusual in respect of the finding in CSF of a CD8+ lymphocytic pleocytosis, which responded to antiretroviral therapy. This has not been reported previously as a manifestation of DILS, but Itescu et al, in the original description of DILS, reported that three of 17 patients had “aseptic meningitis.” In these three patients the CSF cell types and their phenotype were not reported and no reponse to antiretroviral therapy was recorded.

Eosinophilic infiltration of muscle has not previously been described in HIV associated polymyositis/DILS. Common infectious causes include the helminths Trichinella, Taenia solis, and the protozoan T gondii. These and other infections were excluded by negative findings on microscopy of the muscle biopsy and by negative serological tests. The negative autoantibody screen and the absence of vasculitis on histology of the muscle excluded connective tissue disease, polymyositis nodosa, and Churg-Strauss as causes of the eosinophilic component of the infiltrate.

In the second case demonstration in muscle of abnormal mitochondria, in the absence of a previous history of muscle disease and a negative family history and in the context of exposure to the NRTI didanosine strongly suggests that this was the aetiological agent. The patient’s recovery when this drug was withdrawn lends further support to this hypothesis. Despite the frequency of reports of symptoms of myalgia and fatigue in clinical trials of NRTI, including didanosine, documented muscle disease caused by didanosine is very rare. In the majority of reports which describe abnormalities of CPK, with or without symptoms of muscle weakness or fatigue, which implicate didanosine as a cause of mitochondrially mediated myotoxicity, patients received zidovudine contemporaneously, or had received it until recently. In vitro, some studies have shown that didanosine exerts a toxic (mitochondrially mediated) effect on muscle, albeit less prominent than that induced by stavudine, zalcitabine, and zidovudine. By contrast, other in vitro studies have failed to demonstrate that didanosine, at therapeutic doses, has a toxic effect on mitochondria in muscle. However, didanosine is clearly myotoxic, as shown by reports of didanosine induced rhabdomyolysis. These two cases add to the differential diagnosis of causes of muscle disease in this patient population. They also serve to underscore the importance of muscle biopsy in identifying...
the underlying pathology in HIV infected patients with non-specific symptoms of muscle weakness, normal or minor elevations of CPK and normal EMG findings.

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