Optic neuropathy in HIV-infected patients results from the HIV infection itself, post-infectious auto-immune disease, opportunistic infections and drugs. Nucleoside reverse transcriptase inhibitors (NRTIs) such as zidovudine and stavudine have known mitochondrial toxicity and can cause mitochondrial myopathies, neuropathies, hyperlactataemia, and can induce mitochondrial genetic disorders. Individuals with the mutation for Leber’s hereditary optic neuropathy (LHON), a mitochondrial disorder, are usually asymptomatic but develop visual loss when exposed to external triggers such as smoking. We report on two HIV-infected patients with LHON mutations (m.14484T>C and m.11778G>A) who developed profound visual loss with antiretroviral therapy. We postulate that the phenotypic expression of LHON in these genetically predisposed individuals was triggered by NRTI drugs lamivudine and tenofovir when used in combination, despite their relatively weak mitochondrial toxic effects.

Table 1. Common causes of HIV-associated optic neuropathy

| Category               | Causes                                      |
|------------------------|---------------------------------------------|
| Immune mediated        | Isolated seroconversion optic neuritis      |
|                        | Acute disseminated encephalomyelitis        |
| Infectious optic neuritis| Bacterial: syphilis, tuberculosis          |
|                        | Viral: cytomegalovirus, varicella zoster virus, herpes simplex virus |
|                        | Parasitic: toxoplasmosis                    |
|                        | Fungal: cryptococcal meningitis             |
| Raised intracranial pressure | Cryptococcal-induced optic neuropathy       |
| Infiltrative optic neuropathy | Lymphoma                                   |
|                        | Nutritional optic neuropathy               |
|                        | Vitamin B₁₂ deficiency                     |
| Drug induced           | Drug induced                                |
|                        | Ethambutol                                  |
|                        | Antiretrovirals                             |

Optic neuropathy in HIV disease is due to the virus itself, para-infectious disease, opportunistic infections and drugs. Nucleoside reverse transcriptase inhibitors (NRTIs) such as zidovudine and stavudine have known mitochondrial toxicity and can cause mitochondrial myopathies, neuropathies, hyperlactataemia, and can induce mitochondrial genetic disorders. Individuals with the mutation for Leber’s hereditary optic neuropathy (LHON), a mitochondrial disorder, are usually asymptomatic but develop visual loss when exposed to external triggers such as smoking. We report on two HIV-infected patients with LHON mutations (m.14484T>C and m.11778G>A) who developed profound visual loss with antiretroviral therapy. We postulate that the phenotypic expression of LHON in these genetically predisposed individuals was triggered by NRTI drugs lamivudine and tenofovir when used in combination, despite their relatively weak mitochondrial toxic effects.

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cause optic nerve dysfunction in over 95% of patients by disrupting complex I-dependent adenosine triphosphate synthesis. The abnormal mitochondrial gene is maternally inherited with incomplete penetrance, as only 50% of males with the LHON mitochondrial mutation present with the disorder. Expression of the disease occurs when triggered by environmental factors such as smoking, alcohol and acute illness. The typical presentation is sequential or bilateral, simultaneous central visual loss in the 2nd to 3rd decade of life in males carrying one of the LHON mitochondrial mis-sense mutations. Ten percent of female carriers are symptomatic. There is initial disc swelling and circumpapillary telangiectasia, which within a few months are replaced by optic atrophy from axonal loss. Visual loss and centro-caecal scotomas progress over months to years. Idebenone, a co-enzyme Q10 derivative, has shown promise in delaying the progression of illness and in some cases substantially reversing the visual and field loss.\textsuperscript{11}

We present two patients from our neuro-ophthalmology unit who presented with central visual loss after starting antiretroviral therapy containing tenofovir, lamivudine and efavirenz.

Patient 1
A 31-year-old HIV-infected man presented with a 4-month history of gradual and progressive loss of bilateral vision. He denied having headaches, pain on eye movements or constitutional symptoms. He was previously unemployed. Two years before, he acquired pulmonary tuberculosis and was found to be HIV-infected; he was then prescribed antiretroviral therapy containing tenofovir, lamivudine and efavirenz. He had no family history of visual loss.

His visual acuity was ‘counting fingers’ on the right and ‘hand movements’ on the left. Central field defects were detected bilaterally on confrontation, but were globally lost on automated Humphrey visual field (HVF) testing. Colour vision was severely impaired bilaterally (0/15 on Ishihara pseudo-isochromatic plates). Both pupils were sluggishly reactive and severe bilateral optic atrophy was present on funduscopes (Fig. 1). The visual evoked potential (VEP) P100 waveforms were obtainable bilaterally using goggles, but were markedly reduced in amplitude.

Magnetic resonance imaging (MRI) of the brain and orbits was normal. Chest radiograph and cerebrospinal fluid (CSF) examination, including neurotropic virus (cytomegalovirus, herpes simplex viruses 1 and 2, varicella zoster virus, Epstein-Barr virus and John Cunningham virus) testing were unre markable and routine haematological investigations were normal. His vitamin B\textsubscript{12}, folate and iron levels were normal. CD4\textsuperscript{+} cell count was 746 cells/µl and viral load was undetectable. The antinuclear factor (ANF), antineutrophil cytoplasmic antibody (ANCA), rapid plasma reagin (RPR) and rheumatoid factor (RF) were negative, and serum angiotensin converting enzyme (SACE) was 19 U/l. Oligoclonal bands (OBs) were not detected in the CSF and aquaporin 4 antibodies were negative. He tested positive for the 14484T>C mitochondrial mutation associated with LHON.

Patient 2
A 33-year-old HIV-infected man who worked in construction presented with bilateral visual loss over the previous 3 years. He had been diagnosed with HIV infection after voluntary testing and prescribed tenofovir, lamivudine and efavirenz 3 months before the onset of his visual symptoms. He first noticed loss of central vision in the left eye, followed by similar symptoms in the right a month later. He denied any colour desaturation and pain on eye movements, but did notice intermittent oscillopsia at the onset of his symptoms. He had no other medical history of note and denied any smoking or alcohol use, exposure to any toxins or substance abuse. He had no family history of visual loss.

His visual acuity bilaterally was ‘counting fingers’. Central scotomas were noted on confrontation, but HVF showed bilateral, inferior altitudinal field defects. Colour vision was severely impaired (0/15 on Ishihara pseudo-isochromatic plates) and funduscopes revealed bilateral profound optic atrophy (Fig. 2). His pupils were reactive and a 1+ relative afferent pupillary defect was present on the right. The rest of his neurological examination was normal, and standard haematological, biochemical and CSF examination tests were normal. His CD4\textsuperscript{+} count was 615 cells/µl and viral load was 27 copies/ml. MRI of the brain and orbits was normal, as was his chest radiograph. The VEP P100 wave amplitudes were reduced bilaterally. His vitamin B\textsubscript{12} level was mildly reduced at 117 pmol/l (normal 133 - 675), serum folate was normal and SACE was normal.

Fig. 1. Funduscopy of patient 1, showing bilateral optic atrophy and normal retinas.

Fig. 2. Funduscopy of patient 2, showing bilateral optic atrophy and normal retinas.
27 U/L. ANF, ANCA, RF, RPR, human T lymphotrophic virus 1, CSF OB and aquaporin 4 antibodies were negative. He tested positive for the 11778G>A mitochondrial mutation associated with LHON.

Discussion

Genetic testing of LHON by polymerase chain reaction and restriction enzyme analysis is offered by the Inherited Metabolic Disease Laboratory, National Health Laboratory Service, based at Groote Schuur Hospital in Cape Town. All six LHON mutations in mitochondrial DNA are screened for, viz. m.14484T>C, m.11778G>A, m.3460G>A, m.14459G>A, m.14482C>G, and m.14487T>C. Both patients had a common mitochondrial mutation associated with LHON. Patient 1 was a smoker without a significant pack-year history and patient 2 had mild vitamin B12 deficiency, neither of which seemed to have contributed significantly as triggers for LHON, or to the visual loss on a toxic or nutritional basis. Both patients, however, were HIV-positive, developed their symptoms some time after starting ART and were not on other medication that could have contributed to visual impairment.

LHON following the introduction of the strongly mitochondrial toxic nucleoside reverse transcriptase inhibitors (NRTIs) zidovudine, stavudine and zalcitabine (ddC) has previously been reported in patients with the mitochondrial mutations 11778G>A and 14484T>C. No cases have been described with the weaker mitochondrial toxic NRTI drugs tenofovir and lamivudine until now. The catastrophic sequelae of lactic acidosis that occurs with zidovudine and stavudine, however, is less likely to occur with the tenofovir-lamivudine combination. Tenofovir is a less toxic agent than zidovudine, didanosine, stavudine and zalcitabine, and does not usually cause lactic acidosis, myopathy and peripheral neuropathy. Its mechanism of action, like the other NRTIs, is mitochondrial DNA polymerase inhibition, and thus it is not devoid of toxic effects. Tenofovir-induced nephropathy occurs on the basis of mitochondrial toxicity within the kidneys’ proximal tubular cells, which are rich in mitochondria. Irregularly shaped mitochondria with fragmented cristae ensue. There are currently no data on the effects of tenofovir on the mitochondria of the optic neurons and retina. However, an analogous mechanism of toxicity at the optic nerve head, which also has abundant mitochondria, is conceivable with tenofovir. However, based on existing evidence, the role of lamivudine in mitochondrial toxicity is less clear. Hence, we postulate that in the genetically susceptible individual, tenofovir has the potential to trigger LHON.

Idebenone (co-enzyme Q10 derivative) is not available in SA. However, co-enzyme Q10 is available as an over-the-counter preparation at most pharmacies. Both patients were treated with this formulation and have not shown any deterioration or improvement after 6 months of treatment. ART was discontinued in both patients and when required, both patients will be considered for a combination of antiretrovirals with lower mitochondrial toxicities, such as protease and integrase inhibitors.

Conclusion

LHON can be triggered by NRTIs in HIV-infected patients who harbour the LHON mutations. The expression of LHON can occur regardless of the mitochondrial toxic potential of ART. The routine screening for LHON mutations on all asymptomatic male patients about to commence NRTIs is not cost effective. However, the progression of LHON is reasonably manageable with appropriate therapy; hence testing for LHON mutations in HIV-positive patients with optic neuropathy who are on ART should become standard practice.

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