CASE REPORT

Concurrent granulomatous *Pneumocystis carinii* and *Mycobacterium xenopi* pneumonia: an unusual manifestation of HIV immune reconstitution disease

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This report of concurrent granulomatous *Pneumocystis carinii* pneumonia (GPCP) and *Mycobacterium xenopi* pneumonia (MXIP) in a patient with advanced HIV disease 3–5 weeks after commencing antiretroviral therapy (ART) fits the syndrome of HIV immune reconstitution/restoration disease (IRD). It may also be a unique window into the immunity of granulomatous inflammation.

A 35 year old white HIV positive man commenced antiretroviral therapy (ART) with zidovudine, lamivudine, and nelfinavir 7 years after diagnosis (CD4 count 20 cells/mL, viral load 154 000 copies/mL (Chiron III)). Prophylaxis for *Pneumocystis carinii* pneumonia (PCP) was with nebulised pentamidine. Three weeks after commencing ART, with no prior respiratory symptoms, he developed an unproductive cough. Despite 2 weeks of treatment with amoxicillin, his cough persisted and, although he was well, a chest radiograph 5 weeks after initiating ART showed bilateral apical consolidation (fig 1A). Arterial blood gas tensions on air were normal (Pco2 4.53 kPa, Po2 16.3 kPa, pH 7.468). Bronchoscopy examination after 3 days of treatment with high dose intravenous Seprtin showed generalised mucosal inflammation, and the bronchoalveolar lavage (BAL) fluid had scanty *Candida albicans* but was negative for acid fast bacilli and *P carinii*. Ten days after PCP treatment the apical consolidation improved significantly (fig 1B) but the appearance of a new area of right mid-zone consolidation led to diagnostic biopsies to exclude a malignant process. A percutaneous computed tomographic (CT) guided lung biopsy (left apex, 3×20G tissue cores) was undertaken 13 days after admission. The CT scan showed consolidation (fig IC and D) with multiple small and medium sized pulmonary nodules (many cavitated) in both upper and right middle lobes. Histological examination showed granulomatous PCP (fig 1E and F). Blood culture for *M avium* complex and polymerase chain reaction for *M tuberculosis* were both negative. *M xenopi* was isolated from the BAL fluid and three sputum specimens after 6 weeks but tissue culture was negative. Precautionary empirical treatment for PCP with clarithromycin, rifampicin, isoniazid, pyrazinamide and ethambutol was added. Three weeks after commencing MXIP treatment the right middle lobe abnormalities 14 months later. The patient remains well on ART with a CD4 count of 510 cells/mL and an undetectable viral load.

**DISCUSSION**

Race *et al* reported five patients who, within weeks of commencing ART, developed a systemic syndrome with marked inflammatory reactions to latent infection with *M avium* complex. All experienced CD4 increments. Such paradoxical hyper-accentuated inflammatory responses to pre-existing or co-existing infections are recognised complications of immune reconstitution following ART; occurring within days to months and invariably associated with CD4 increments, they are collectively known as immune reconstitution/restoration disease (IRD). The dynamics of T cell increments following ART is a biphasic phenomenon; the first phase (weeks to 6 months) is predominantly of T memory cell redistribution, while the second phase (usually >6 months) is mainly of naïve CD4 cell reconstitution. As IRD invariably coincides with the primary phase of T memory cell redistribution, it has been postulated that this may be a memory T cell versus pathogen/antigen recognition event.

*Pneumocystis carinii* infections have been, uncommonly, reported as IRD. Because co-existence of multiple opportunistic organisms in the respiratory tract of patients with advanced HIV disease is common, diagnosing non-*M tuberculosis* infections can be difficult; in our patient the isolation of *M xenopi* from BAL fluid and sputum specimens and the clinical features all fit the diagnostic criteria for non-*M tuberculosis* infections. The differential time scales for the resolution of the apical radiographic abnormalities (rapid diminution after 10 days of PCP therapy) and the radiographic abnormalities of the right middle lobe (14 months) are compatible with concurrent granulomatous *P carinii* pneumonia (GPCP) and *M xenopi* pneumonia (MXIP).

The granuloma, a localised inflammatory process, can form part of the host’s immune protection response to infections or part of an autoimmune disease process. Patients with AIDS have T cell deficiency and consequently have difficulty in mounting a granulomatous reaction to infections. Despite this, our patient mounted a hyper-accentuated response weeks after commencing HAART.

Bondoc *et al* reported GPCP in three non-HIV patients with malignancy who shared similarities with our patient. GPCP occurred 2 weeks to 2 months after stopping corticosteroids compared with 3–8 weeks after receiving ART in our patient. All had minimal symptoms, a comparable time scale of immune reconstitution, and *P carinii* was not detected in the BAL fluid. These two reports seem to support recent evidence that the main pathway for T cell recruitment during granuloma formation is from the systematically activated T cell pool. Corticosteroid therapy suppresses peripheral circulating lymphocytes and its discontinuation leads to a rebound of peripheral lymphocytes, a process akin to the phase of memory T cell redistribution following ART.

The simultaneous occurrences of GPCP and MXIP are perhaps less surprising when viewed in the context of T memory cell (therefore diverse memory clone) redistribution. Recent evidence has shown that the T cell repertoire of granulomas in response to a primary pathogen is diverse.
Only 30–60% in a single granuloma are specific to the inducing antigen, and systemically activated T cells with no specificity to the granuloma also have access to it.9

Cytokine gene polymorphisms11 and an abnormal cytokine profile12 have been associated with HIV IRD. This, together with a recent report of granulomatous pulmonary disease occurring in a patient following cessation of tumour necrosis factor (TNF)-α antagonist treatment,13 implies that perturbation of the cytokine profile can trigger granulomatous inflammation.

Host and antigenic factors implicated in the pathogenesis of granulomatous lung diseases are also likely to influence the expression of GPCP; in addition, immune dysregulation (systemic T cell pool and cytokine perturbation) may also trigger granulomatous inflammation.

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REFERENCES


LUNG ALERT

Sildenafil and exercise capacity in hypoxic pulmonary hypertension

This study examined the influence of oral sildenafil on pulmonary haemodynamics and exercise tolerance during hypoxia induced pulmonary hypertension in healthy mountaineers and trekkers who were not susceptible to high altitude pulmonary oedema. Fourteen healthy volunteers (12 men, median age 36.5 years) were enrolled and randomised in a double blind, placebo controlled, crossover design. Participants were assigned to receive placebo or one 50 mg dose of sildenafil. Systolic pulmonary artery pressure, cardiac output, and peripheral arterial oxygen saturation at rest and during assessment of maximum exercise capacity on cycle ergometry were measured, both while breathing a hypoxic gas mixture with 10% fraction of inspired oxygen at low altitude and then at the Mount Everest base camp (5400 m).

Sildenafil reduced hypoxic pulmonary hypertension at rest and with exercise, and increased maximum exercise capacity and cardiac output. The study did not examine the effects of sildenafil on normoxic exercise tolerance, and the authors cautioned about the unclear role of sildenafil in the management of acute mountain sickness. In addition, it is not known whether sildenafil increases exercise capacity in hypoxic pulmonary hypertension due to lung diseases such as COPD.

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References

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