Interactive effect of HIV infection and smoking on lung immunity and related disorders

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Take-home message of article:
Smoking exacerbates the immunosuppressive/proinflammatory effects of HIV, increasing the risk for pulmonary disease.
ABSTRACT

HIV-infected persons not only have higher rates of smoking than the general population, but are also unusually vulnerable to the associated adverse health effects, both infective and non-infective in origin. Indeed, in the setting of well-organised care and availability of HAART, HIV-infected smokers lose more life years to smoking than to HIV infection per se, presenting a major challenge to healthcare providers. Not surprisingly, the respiratory system is particularly susceptible to the damaging, interactive chronic inflammatory and immunosuppressive effects of HIV and smoking, intensifying the risk for development of opportunistic infections, as well as lung cancer and obstructive lung disorders. The impact of smoking on the immunopathogenesis and frequencies of these respiratory conditions in the setting of HIV infection, as well as on the efficacy of antiretroviral/antimicrobial therapy, represent the primary focus of the current review.

INTRODUCTION

Fifty years after the release of the first report by the US Surgeon General on the health effects of smoking, tobacco use continues to pose a massive public health burden. It remains the single largest preventable cause of death and disease in both men and women, killing nearly 6 million people annually, including 600 000 non-smokers exposed to second-hand smoke [1]. Tobacco use caused approximately 100 million deaths in the 20th century and, if current trends continue, this number may increase to one billion deaths in the 21st century. However, these official rates may be an underestimate since recent research has shown that approximately 17% of the excess mortality among smokers could be attributed to causes not currently associated with smoking [2].

Currently, in excess of 1.1 billion people in the world smoke. Despite the seemingly encouraging estimates of global modeling, which projected that smoking prevalence should have decreased between 1980 and 2012, in reality the opposite was true. The estimated prevalence was offset by the considerable population growth during this period, with the total number of regular smokers and cigarettes consumed actually increasing [3]. In addition, over the last two decades the global burden of disease attributable to tobacco smoking has not changed, since decreases in high-income regions have been tempered by increases in regions such as southeast Asia [4]. While Asia is currently the epicentre of the tobacco epidemic, Africa presents the greatest threat in terms of potential growth in tobacco consumption [4], converging with the human immunodeficiency virus (HIV) pandemic.

Globally, an estimated 35 (33.2–37.2) million people were living with HIV in 2013 and despite enormous achievements in treatment resulting in 12.9 million of these having been initiated on highly-active antiretroviral treatment (HAART), as well as some advances in prevention, new infections continued at a rate of approximately 5,800 daily, totalling 2.1 (1.9–2.4) million annually worldwide. Notably, HIV-infected individuals appear to smoke more than the general population. For example, in the USA, over 85% of HIV-infected individuals have a lifetime history of cigarette smoking, while
studies in European and North American HIV-infected cohorts report a smoking prevalence ranging from 40 to >70%, two to three times that of the general population [5–11]. Notwithstanding the increased prevalence of smoking, HIV-infected smokers have a significant, albeit unexplained, increased mortality and decreased quality of life compared to HIV-uninfected smokers [12].

A major impact of HIV infection is on the respiratory system [13]. This includes the occurrence of both communicable diseases, notably tuberculosis (TB), community-acquired pneumonia (CAP) and *Pneumocystis jirovecii* pneumonia (PCP), as well as non-communicable conditions such as obstructive lung diseases and lung cancer. All of these conditions are associated with concomitant cigarette smoking with the risk seemingly greater in the HIV-infected [14, 15]. Some have even suggested that we are witnessing the convergence of several epidemics, namely smoking, HIV infection, TB, other respiratory infections, obstructive lung disease and lung cancer [16, 17]. While opportunistic respiratory infections associated with advanced immunosuppression dominated the initial clinical picture in the setting of HIV infection, latterly, with the use of HAART, there has been a rise in deaths related to non-AIDS defining illnesses, largely driven by the high rate of smoking among these individuals [15, 18]. In fact, in those areas where HIV care is well organised and HAART is freely available, HIV-infected smokers lose more life-years to smoking than to HIV infection *per se*, the risk being considerable and often attributable to non-AIDS-defining conditions [12, 15, 18, 19].

The impact of smoking on the immunopathogenesis and occurrence of these respiratory conditions in the setting of HIV infection, as well as on the efficacy of antiretroviral/antimicrobial therapy, represent the primary focus of the current review.

**IMMUNOSUPPRESSIVE MECHANISMS OF SMOKING AND HIV INFECTION**

*HIV and pulmonary host defences*

HIV infection has been described as a “disease of secondary lymphatic tissues,” characterised by massive early depletion of the CD4*, CCR5-expressing memory T cell population in the gastrointestinal tract (GIT) [20], with the Th17 subset particularly vulnerable [21]. Although T cell repopulation of the GIT ensues, the recovery is only partial, due to several mechanisms, including defective production of the homing cytokine, CXCL25, by epithelial cells and dendritic cells, impeding the trafficking of gut-homing Th17 cells [22].

In contrast to the gastrointestinal tract (GIT), the lung appears relatively resistant to the early HIV-mediated depletion of CD4* T cells [21, 23]. This has been attributed to several mechanisms including: i) resilience of airway Th1 and Th17 cells [21, 23], as well as large, as opposed to small, alveolar macrophages [24] to HIV infection and dysfunction; ii) relatively high frequencies of mucosal, HIV-specific polyfunctional CD4* T cells in the lungs; iii) a lesser mass of HIV-susceptible secondary
lymphoid tissue relative to that of the GIT [20]; and iv) the apparent efficacy of the pulmonary HIV-specific cytotoxic T cell response, manifested clinically as an alveolitis [25].

Nonetheless, these adaptive and innate pulmonary host defences progressively deteriorate, being ultimately subdued, with the following seeming to be the most prominent mechanisms contributing to HIV-mediated immune attrition/exhaustion:

- cytopathic effects of HIV on CD4+ T cells [25, 26]
- elimination of HIV-infected CD4+ T cells by HIV antigen-specific cytotoxic T cells [25, 26]
- permissiveness of follicular helper T cells (Tfh) to HIV infection, compromising production of specific antibodies, while serving as the major reservoir for HIV replication [27]
- induction of various pathways of apoptosis following prolonged exposure of T cells to HIV proteins such as Vpr and Tat, as well as to the family of common γ-chain cytokines, interleukins (IL)-2, -4, -7, -9, -15 and -21 [28–31]. These pathways include Fas/Fas ligand, tumour necrosis factor (TNF)/TNF-R1/R2, TNF-related apoptosis-inducing ligand (TRAIL)/death receptors 4/5 and others, programmed death receptor-1 (PD-1) and its ligand (PD-L1), and apoptotic microparticles released from apoptotic cells [28–31].
- persistent production of interferon-α and indoleamine 2,3 dioxygenase by chronically activated plasmacytoid dendritic cells, promoting apoptosis of CD4+ and CD8+ T cells, and induction of immunosuppressive regulatory T cells (Tregs) [32]. This results predominantly from the interaction of viral nucleic acid with intracellular Toll-like receptor 7 (TLR7), and possibly with TLR9, as well as cytoplasmic pathogen nucleic acid sensors [32].
- abortive infection of quiescent, “bystander” CD4+ T cells, resulting in the cytosolic accumulation of incomplete HIV reverse transcripts (DNA) [33]. These, in turn, are recognised by the interferon-γ-inducible viral cytosolic DNA receptor, IFI16, which forms an inflammasome, resulting in the activation of an intense caspase-1-mediated inflammatory response and a type of inflammation-associated cell death known as pyroptosis [33].
- chronic systemic activation of monocytes [28, 34, 35] and neutrophils [35, 36] by HIV and its proteins, as well as exposure to pro-inflammatory microbial products (DNA and lipopolysaccharide) originating from a damaged GIT by the process of microbial translocation [34] (Figure 1). In the case of activated macrophages, release of pro-inflammatory cytokines and reactive oxygen species (ROS) are likely to drive T cell apoptosis and dysfunction respectively [28, 35]. Neutrophils also acquire an activated phenotype characterised by
increased production of ROS, release of arginase and expression of PD-L1, driving T cell
dysfunction and death [35, 36].

- increased numbers of circulating myeloid suppressor cells as HIV infection progresses [37].

- production of high levels of the counteracting immunosuppressive cytokine, transforming
growth factor-β1 (TGF-β1), by various types of immune, inflammatory and structural cells,
promoting development of Tregs [38] and disruption of T cell colonisation of secondary
lymphoid tissues [20], presumably as a strategy to counter chronic immune activation [39,
40]. TGF-β1 contributes to HIV-related immunosuppression by several mechanisms,
including: i) induction of the transcription factor, Foxp3, a key event in the development of
Tregs [38]; and ii) promotion of collagen deposition and fibrosis in secondary lymphoid
tissues, disrupting T cell colonisation and function [20].

These immunosuppressive mechanisms are summarised in Figure 2.

**Smoking and immunity**

Similarities exist between the effects of HIV infection and those of smoking on immune reactivity. The
wide-ranging suppressive effects of smoking, encompassing both innate and adaptive airway host
defences, have been described elsewhere [41] and are summarised in Figures 1 and 2 and Table 1.
These immunosuppressive and pro-inflammatory effects of smoking result from constant exposure of
the airways to the vast array of low molecular toxicants generated during the combustion of tobacco.
These include various hydrocarbons, gasses, highly-reactive free radicals, and heavy metals [41].
Tobacco also contains various pro-inflammatory microbial products, including bacterial endotoxin,
which remain biologically active in cigarette smoke, albeit at a lower level [42, 43]. Prolonged
exposure to the cytotoxic and pro-inflammatory agents in cigarette smoke results in chronic
inflammation, oxidative stress and immunosuppression, increasing susceptibility to respiratory
infection [41] particularly with the bacterial pathogens, *Streptococcus pneumoniae* and *Mycobacterium
tuberculosis* [41].

Like HIV infection [35, 36, 44], smoking also causes a chronic, low-grade, systemic inflammatory
response characterised by increased numbers of activated neutrophils in the circulation and lungs [41,
45]. Smoking-related, systemic inflammation has been attributed to extrapulmonary dispersal of the
neutrophil-mobilising cytokines, interleukin (IL)-1β, IL-6, tumour necrosis factor (TNF) and
granulocyte/macrophage colony-stimulating factor (GM-CSF), as well as the chemokine, IL-8,
produced by resident airway cells [41, 45, 46]. Smokers also have higher levels of circulating bacterial
endotoxin than non-smokers, though the origins (tobacco/tobacco smoke or leakage from the GIT) are
unknown [47]. Ongoing neutrophil-mediated systemic inflammation and endothelial activation [48] are
potential contributors to the unusually high frequency of chronic obstructive pulmonary disease
(COPD) and lung cancer in HIV-infected individuals who smoke [15, 49].
Interactive effects of HIV infection and smoking on immunity

Increased permissiveness of alveolar macrophages from smokers to HIV infection and replication in vitro has been described [50], while exposure of isolated human microglial cells [51] or TZM-bl (CD4/CCR5 co-expressing HeLa cells) and Jurkat T cell lines [52] to nicotine or cigarette smoke extract respectively, also augments HIV infectivity. Smokers also have a more rapid loss of CD4+ T cells during the course of HIV infection [11, 53], consistent with an increased likelihood, or rate, of progression to AIDS [53, 54]. The latter contention was not supported by other studies [55–58], possibly due to differences in the populations studied [59].

To our knowledge, however, no studies have reported possible interactive, immunosuppressive effects of HIV infection and smoking on functional indices of immune reactivity in HAART-naïve subjects. The closest are two recent reports, largely in the setting of virally suppressive HAART, in which smoking was associated with significant elevations in cellular and soluble biomarkers of systemic immune activation/exhaustion/dysfunction [47, 60]. The first of these described increased expression of CD38 and HLA-DR, as well as PD1/Tim3/CTLA4, markers of immune activation and exhaustion, respectively, on both CD4+ and CD8+ T cells of HIV-infected smokers [47]. Because these effects of smoking were associated with raised levels of circulating bacterial endotoxin, the authors proposed that smoking, like HIV infection, causes microbial translocation with consequent augmentation of systemic immune activation [47]. This contention can, however, be challenged on two fronts. Firstly, the levels of circulating bacterial endotoxin were comparable in HIV-infected smokers and nonsmokers [47], and secondly, as mentioned above, endotoxin is present in both tobacco and tobacco smoke [42, 43].

The second study compared the numbers and reactivities of pulmonary mucosal and circulating CD4+ and CD8+ T cells from HIV-infected subjects, many of whom were current smokers and mostly on HAART, with and without COPD, a condition for which both HIV and smoking are major risk factors [60]. Relative to the HIV+/COPD- group, as well as to an HIV-/COPD+ control group, the authors observed significant, selective depletion of pulmonary mucosal CD4+ T cells, including memory and polyfunctional cells, in the HIV+/COPD+ subjects. This was associated with deficient responses of mucosal CD4+ T cells to both HIV- and bacteria-derived antigens in the setting of increased co-expression of the pro-apoptotic markers, Fas death receptor and PD-1 [60]. Although consistent with activation-induced cell death and immune exhaustion, possibly related to an interactive effect of HIV and smoking, this interpretation is complicated by the high number of current smokers in the HIV+/COPD- group and lack of inclusion of numbers of ex-smokers in both HIV-infected groups, as well as data on the use of inhaled corticosteroids and duration of HAART. Interestingly however, the median smoking history (pack years) was shorter, albeit not significantly so, in the HIV+/COPD-group relative to the HIV+/COPD+ group (8.45 versus 13.4 pack years), suggesting that early administration of HAART to HIV infected smokers may slow the development of COPD. Although speculative, this contention is supported by findings that the lung is more responsive than the vascular
compartment to the virally suppressive actions of HAART with a more rapid decline in viral loads in both the acellular compartment and alveolar cells [61]. This is paralleled by partial immunoreconstitution, associated with decreased numbers and activation status of pulmonary CD8+ T cells, and a gradual increase in the numbers of CD4+ T cells in the setting of decreased concentrations of pro-inflammatory cytokines/chemokines [61].

The most compelling evidence in support of an interactive, immunosuppressive effect of HIV infection and smoking is, however, derived from clinical-epidemiological studies demonstrating increased frequencies and severity of TB, PCP and pneumococcal pneumonia in HIV-infected smokers described in later sections of this review.

EFFECTS OF SMOKING ON HIV DISEASE PROGRESSION AND TREATMENT

Research has yielded inconsistent results on the effects of smoking on HIV disease progression and outcome on antiretroviral treatment. It is known that the number of leukocytes per microliter in peripheral blood is directly related to cigarette consumption, with HIV–uninfected smokers having higher total white cell counts than non-smokers, so-called ‘smoker’s leukocytosis’ [62]. This effect is non-specific, affecting all leukocytes, including CD4+ T cells. Some authors have suggested that the specific leukocyte population affected is dependent on the amount of smoking, with light smokers having a higher, and heavy smokers a lower, CD4+ to CD8+ ratio [63]. Nevertheless, this difference is attenuated at the time of HIV seroconversion and disappears after 3 years [64]. Some studies have shown that smokers have a more rapid loss of CD4+ T cells during the course of HIV infection [11, 53] and an increased likelihood or rate of progression to AIDS [53, 54]. Such a contention is plausible based on findings that the magnitude of HIV replication is a strong predictor of the rate of progression to AIDS [65] and is supported by several in vitro studies which have shown that smoking enhances HIV-1 replication in alveolar macrophages, microglia and T cells [50, 51, 52]. It has been suggested that iron and oxidative stress are possible mechanisms of enhanced production of HIV-1 by alveolar macrophages in cigarette smokers [66, 67]. In contrast, a number of clinical studies failed to confirm accelerated disease progression in smokers [55–58], related most likely to the specific populations studied [59].

On the other hand, smoking may be associated with decreased efficacy of HAART, resulting in both immunological and virological failure [11, 68]. One study has shown that long-term smoking attenuated the immunological and virological responses to HAART by 40% [68]. These findings are supported by the Women’s Interagency HIV Study (WIHS) which also reported that women who smoked while on HAART had significantly higher morbidity (AIDS-defining conditions (HR=1.36; P=0.01)) and mortality (HR=1.53; P=0.018) than nonsmokers, and that HAART may be less effective. Smokers had a lower chance of achieving a viral (HR=0.79; P=0.006) or immunologic response (HR=0.85; P=0.041), and a greater chance of developing viral (HR=1.39; P=0.013) or immunologic failure (HR=1.52; P=0.001) [11]. In this study, significantly more smokers were African-American,
used illicit drugs, or had a life-time history of illicit injection drug use, were infected with hepatitis C virus, had previously been diagnosed with AIDS, and had lower levels of adherence. Although this analysis controlled for possible confounders, unidentified residual influences cannot be excluded.

Notwithstanding the interactive, counteracting immunosuppressive effect of HIV/smoking on therapy, other possible causes of smoking-related interference with HAART include: i) smoking is a surrogate marker of a non-compliant disposition [69, 70]; ii) higher frequencies of side-effects in smokers including neuropsychiatric symptoms associated with efavirenz-based HAART regimens [71]; iii) negative effects of smoking-related activation of cytochrome P450 enzymes, such as the CPY1A1-m1 variant, which promote oxidative conversion of smoke-derived toxicants to DNA adducts, causing activation of genes which support HIV replication [67, 72]; iv) altered pharmacokinetics of antiretroviral agents, possibly related to smoking-mediated induction of CYP3A4 and aryl-hydrocarbon-hydroxylases [73, 74], which metabolise many protease and non-nucleoside reverse transcriptase inhibitors [75, 76]; and v) smoking-induced mitochondrial oxidative stress, [77] which may exacerbate the adverse effects of antiretroviral drugs that induce mitochondrial toxicity, such as drugs from the nucleoside reverse transcriptase inhibitor class [78].

Smokers may also be at heightened risk for developing HIV-associated drug resistance due to smoking/oxidative stress-associated increased activity of nuclear transcription factor kappa B [79], driving both viral transcription and diversity [80]. However, this remains to be proven in the clinical setting.

**CLINICAL CONSEQUENCES OF THE INTERACTIVE EFFECTS OF SMOKING AND HIV INFECTION ON THE MAJOR HIV-ASSOCIATED PULMONARY CONDITIONS**

**PULMONARY INFECTIONS**

*Community-acquired pneumonia (CAP)*

It is well recognised that smoking is an independent and eminently preventable risk factor for development of CAP in both HIV-infected and -uninfected individuals [13, 81–87]. In HIV-infected patients, the increased risk for CAP, which is evident at all levels of immunosuppression, is estimated to be between 2- and 5-fold, increasing as the CD4 cell count declines [13, 88]. Although rates of CAP in HIV-infected individuals have decreased since the introduction of HAART, they nevertheless remain higher than in HIV-uninfected persons with smoking a major and persistent risk factor [14, 82, 87].

*Streptococcus pneumoniae* is the most common cause of CAP in both the HIV-infected and -uninfected, accounting for some 20% of cases of all-cause bacterial pneumonia, 40% of cases of CAP in which a specific pathogen is identified, and 70% of cases of bacteraemic pneumonia [13, 85, 88].
The rates of invasive pneumococcal disease (IPD) associated with HIV infection are estimated to be up to 100-fold higher compared to HIV-uninfected persons [85]. Predictably, smoking, as well as alcohol abuse, are also significant risk factors for IPD, with both of these harmful aspects of lifestyle being over-represented in those infected with HIV [88, 89].

Importantly, cessation of cigarette smoking by HIV-infected individuals significantly decreases the risk for development of IPD and other pneumococcal infections irrespective of the degree of immunodeficiency [87, 89–92].

**Tuberculosis (TB)**

Mounting evidence from a number of countries in both the developed and developing worlds attests not only to the predisposing effect of smoking for development of pulmonary TB [93–100], but also for more severe disease progression [101], decreased efficacy of antimicrobial chemotherapy [102], prolonged infectivity [103–105], recurrence of disease [106], poor outcomes of MDR-TB [107], and possibly impaired post-treatment lung function [108].

The link between TB and HIV infection is also well established, with TB recognised as the most important opportunistic infection in HIV-infected patients in both sub-Saharan Africa and other regions of the developing world [16]. Long-term smoking has been reported to double the risk of TB in HIV-seropositive patients on HAART [68]. Although somewhat under-researched, it has been reported that smoking is associated with a poor outcome for opportunistic infections in HIV-infected patients, including TB [14, 16, 109].

With respect to TB therapy-related adverse events, one study reported a significantly increased risk (OR = 2.00) in smokers [110]. In contrast, smoking has been reported to attenuate isoniazid-associated hepatotoxicity by increasing N-acetyltransferase 2 activity with potential implications for efficacy of therapy [111]. Although the impact of smoking on the efficacy of combined HAART/TB therapy, as well as on the frequency of therapy-associated adverse effects, requires additional research, cessation of smoking simply on the basis of its negative effects on the individual types of therapy is clearly warranted.

**Pneumocystis jirovecii**

Cigarette smoking is also associated with an increased risk of PCP. In one study, after controlling for HIV status and HAART on multivariate analysis, cigarette smoking was documented to double the risk of developing PCP [68]. A subsequent study by the same investigators documented not only a high rate of tobacco use among 521 HIV-infected patients admitted to hospital (65% being smokers, 40% of whom smoked more than one pack of cigarettes per day), but also that tobacco use increased the risk of respiratory infections, including CAP, PCP, TB, and non-tuberculous mycobacterial infections [14].
HIV-infected patients were three times more likely to be admitted to hospital with PCP even after controlling for HAART, viral load and CD4 cell count [14]. Importantly, in that study, the increased risk of hospitalisation for CAP and PCP was related in a dose-dependent manner to the number of cigarettes smoked per day. Another study investigating *P. jirovecii* colonisation in HIV-infected men, documented not only a high rate of colonisation, but also identified cigarette smoking as an independent risk factor for colonisation [112]. The authors concluded that the increased rates of colonisation in smokers, as well as being a potential source of *P. jirovecii* infection in the colonised individual, may also increase the risk of spread of the disease to others.

**Other infections**

The relationship between cigarette smoking in HIV-infected persons and other infections is less well established. Influenza is a common respiratory tract infection in the general population and cigarette smoking has been shown to increase both the risk and severity of infection [63, 113–117]. However, the situation with regard to risk in HIV-infected individuals of both seasonal and pandemic influenza is less clear-cut, whether in smokers or in non-smokers [88]. One study from South Africa, documented that influenza-related acute lower respiratory infections were approximately 8-fold more common in HIV-infected persons compared with HIV-uninfected cases [118]. This was associated with a higher frequency of influenza B than influenza A, a greater risk of pneumococcal co-infection and a higher mortality; however, there was no difference in smoking prevalence between the two groups of patients [118]. With respect to fungal infections, strong associations between smoking and infection with *C. neoformans* and *C. gattii* have been described in both HIV-infected and -uninfected individuals [119, 120].

**NON-INFECTIVE PULMONARY CONDITIONS**

**Lung cancer**

Much has been written about the occurrence of lung cancer in HIV-infected patients, with initial studies variously documenting it to be increased, decreased or the same in HIV-infected patients, compared to the general population [121–129]. More recently, several studies have suggested that there is indeed an increased incidence of lung cancer in HIV-infected patients, especially evident in the post-HAART era and associated with a poor outcome [122–126, 128, 129], with the adjusted risk ranging from 2.0 to 7.0 [18]. This increased risk may relate, at least in part, to the fact that HIV-infected persons, especially intravenous drug users, tend to smoke more than the general population [123, 126]. This contention is supported by reports of a high frequency of lung cancer in HIV-infected persons who smoke, as opposed to those who do not [123, 127].

On the other hand, several studies have reported that the incidence of lung cancer in HIV-infected individuals remains high even after adjustments for smoking or independently of smoking [121, 124–
In the largest cohort study conducted to date, involving 110,000 veterans, it was reported that HIV-infected cases had a significantly higher risk of lung cancer than HIV-uninfected cases, and that HIV was an independent risk factor for lung cancer after controlling for a number of confounding variables, including smoking [128]. Other studies have documented that lung cancer occurs at a younger age among HIV-infected individuals compared to the general population and at lower exposures to cigarette smoke [121].

Factors other than smoking that may play a role in the increased risk of lung cancer in the setting of HIV infection include HIV-related factors (e.g. oncogenic role of HIV, chronic immunosuppression) and increased susceptibility of HIV-infected persons to mutagens/carcinogens such as chronic inflammation-derived ROS and possibly antiretroviral agents [123, 129]. The role of immunosuppression is unclear, however, with several studies suggesting involvement [124, 127], while others have presented evidence to the contrary [126].

Notably, colonisation with *P. jirovecii* occurs in patients with various chronic lung diseases, and aside from being involved in disease development or transmission, may also be implicated in the development of other lung conditions, including lung cancer [130]. Some studies have documented high rates of colonisation with *P. jirovecii* in patients with lung cancer; one small study noted a colonisation rate of 100% in patients with small-cell carcinoma, as compared to non-small-cell cancer [130]. Other infections that have that been implicated in the pathogenesis of lung cancer include TB and *Chlamydia pneumoniae* infection [131, 132]. The exact mechanisms by which chronic infections may induce cancer development are incompletely understood, but may be related, at least partly, to chronic inflammation, resulting in prolonged exposure of the airways to potentially carcinogenic phagocyte-derived ROS [49].

Clearly, the HAART-associated increase in life expectancy and high smoking rates among HIV-infected patients put them at increased risk of lung cancer, reinforcing the need for a programme of ongoing education and implementation of smoking cessation strategies [126].

**Obstructive lung disease (COPD and asthma)**

In the pre-HAART era, several studies suggested an association of HIV infection with an accelerated form of smoking-induced emphysema in which cytotoxic T lymphocytes were considered to play an important role in pathogenesis [133, 134]. In this context, two studies documenting the impact of smoking in HIV-infected persons in the HAART era have reported an increased frequency of COPD [12, 135]. In the former study, COPD was much more common in current smokers (OR 5.25, 95% CI 1.62-17.01) and ex-smokers (OR 5.25, 95% CI 1.53-17.99) than in never-smokers [12], while in the latter, HIV infection was documented to be an independent risk factor for COPD [135]. However, further studies are needed to determine whether HIV increases the susceptibility to COPD or rather accelerates the course of the disease [135]. Nevertheless, some have suggested that HIV-infected
individuals may be especially susceptible to the effects of cigarette smoke [18]. This contention is supported by the recent study on pulmonary immune dysfunction in HIV-infected subjects with and without COPD mentioned above [60], as well as by the findings from an earlier experimental study, that co-exposure of mice to a viral mimetic (synthetic double-stranded RNA) and cigarette smoke significantly augmented pulmonary inflammation, accelerating development of airway fibrosis and emphysema [136].

Recent research has also focused on the role of *P. jirovecii* colonisation in the pathogenesis and progression of COPD [129, 130, 137]. Although there is clearly an increased rate of colonisation with this pathogen in patients with COPD [130], the link may be purely an epiphenomenon related to structural lung damage and/or immunosuppression due to corticosteroid use. Nevertheless, it is possible that higher rates of airway obstruction in HIV-infected patients, even after adjustments for smoking, could be related aetiologically to both lung and systemic inflammatory responses associated with *P. jirovecii* colonisation [130].

Despite a number of reviews on the topic [129, 134, 138, 139], it remains difficult to determine whether COPD in the setting of HIV infection occurs as a consequence of higher rates of smoking, drug abuse, pulmonary infections, or other risk factors, including occupational or environmental exposures, and low socioeconomic status [138]. These reviews have also focused on potential mechanisms that may be involved in HIV-associated COPD, such as high-risk behaviour (including cigarette smoking), increased susceptibility to pulmonary infection and colonisation, aberrant inflammatory responses, altered oxidant/antioxidant defences, increased apoptosis, and effects of HAART [129, 134, 139]. Irrespective of the mechanisms involved in HIV-associated COPD, cessation of smoking is certainly a priority preventive strategy, while the role of early implementation of HAART remains to be established in this setting.

In addition to COPD, HIV infection has also been found to be associated with an increased prevalence of small airway abnormalities, asthma and bronchial hyperresponsiveness, particularly among smokers [129, 138]. Prior to the introduction of HAART, it was said that asthma was common in patients with HIV infection, possibly related to smoking as well as atopy [129]. In the HAART era, however, findings regarding asthma incidence are somewhat contradictory; some studies have suggested that rates are higher than in the general population with patients reporting the onset of asthma in adulthood, often after the diagnosis of HIV infection [129].

**Other non-infective pulmonary conditions**

Although bronchiectasis appears to be more common in individuals who are HIV-infected, there do not appear to be any studies investigating the role, if any, of cigarette smoking in its pathogenesis [140, 141]. Likewise, the contribution of smoking to the well-recognised association of HIV infection and predisposition for development of pulmonary hypertension [142] is also uncertain with the exception of a single study which assessed the association between HIV, HHV8, socio-demographic information
(including smoking) and pulmonary hypertension [143]. Although no effects of smoking were evident, interpretation is complicated by the high number of smokers in both the HIV-infected and uninfected control group and the lack of objective validation of self-reported smoking history.

POTENTIAL ADJUNCTIVE/ANTI-INFLAMMATORY THERAPIES

While HAART remains the cornerstone of treatment of HIV infection, persistent immune activation/chronic inflammation extending several years beyond the implementation of virally suppressive therapy [144], apparently exacerbated by smoking [47, 60] underscores the necessity for effective adjunctive/anti-inflammatory strategies in this setting. Notwithstanding aggressive implementation of smoking cessation strategies, various therapies targeting chronic immune activation and inflammation which may ameliorate the long-term complications of HIV infection, especially in those who smoke, have been or are currently being investigated (Table 2) [145, 146, 147]. However, research has been restricted to small proof-of-concept trials and should be expanded to larger randomised clinical trials that can evaluate markers of immune activation, as well as clinical events, with special consideration of the outcomes in HIV-infected smokers.

CONCLUSION

Although entirely avoidable, it is abundantly clear that cigarette smoking further compromises a pulmonary immune system already ravaged by HIV, not only exacerbating the predisposition to life-threatening conditions of both infective and non-infective origin, but also countering the benefits of HAART. The magnitude of this problem and the need for prioritisation of effective smoking cessation interventions are underscored by the unusually high frequency of smoking among the HIV-infected, compounded by an apparent lack of awareness of the heightened associated health risks.

SEARCH STRATEGY AND SELECTION CRITERIA

and included systematic reviews and meta-analyses, as well as individual studies when review articles were not available.

CONTRIBUTIONS
All authors contributed equally in conceptualising, writing and editing this work. None of the authors have any conflict of interest or funding sources to declare.

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Table 1: Innate and adaptive pulmonary host defence mechanisms which are impaired by smoking

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<tr>
<th>Innate defences</th>
<th>Adaptive defences</th>
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<td>• mucociliary clearance, epithelial barrier function, production of antiviral and antimicrobial proteins by respiratory epithelium</td>
<td>• production of specific antibodies</td>
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<td>• phagocytic activity of small alveolar macrophages</td>
<td>• cell-mediated immunity (production of interferon-γ)</td>
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<td>• antigen-presenting activities of pulmonary dendritic cells</td>
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<td>• antiviral activities of plasmacytoid dendritic cells</td>
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<tr>
<td>• antitumour and antiviral activities of natural killer cells</td>
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</tbody>
</table>

Described in more detail in reference 41

Table 2: Potential adjunctive/anti-inflammatory therapies

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Potential therapies</th>
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</thead>
<tbody>
<tr>
<td>• enhancing GIT mucosal repair</td>
<td>• bovine serum colostrum, micronutrient supplementation, pro- and pre-biotics</td>
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<tr>
<td>• reducing microbial translocation and endotoxin</td>
<td>• rifaximin, sevelamer carbonate</td>
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<tr>
<td>• intensifying HAART</td>
<td>• maraviroc and raltegravir</td>
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<tr>
<td>• treating co-infections</td>
<td>• valgancyclovir, interferon-α and ribavirin</td>
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<tr>
<td>• reducing activation of plasmacytoid dendritic cells</td>
<td>• chloroquine and hydroxychloroquine</td>
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<tr>
<td>• decreasing TGF-β1 mediated lymph node fibrosis</td>
<td>• pirfenidone, lisinopril</td>
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<tr>
<td>• immune-modulators/anti-inflammatory agents</td>
<td>• HMG CoA reductase inhibitors, minocycline, selective cyclo-oxygenase-2 inhibitors, leflunomide and intravenous immunoglobulin</td>
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<tr>
<td>• inhibiting caspase-1 activity and cleavage of pro-IL-1β</td>
<td>• VX-765</td>
</tr>
</tbody>
</table>

Described in detail in references 145-147
Figure legends (figures attached as separate files)

Figure 1: Proposed mechanisms by which cigarette smoking and HIV infection cause translocation of pro-inflammatory microbial products (.decorate) from the lungs and GIT (.decorate) denotes intestinal microorganisms) respectively. Smoking and HIV infection result in increased alveolar epithelial and intestinal epithelial permeability respectively, with consequent translocation of bacterial endotoxin and other microbial products derived from tobacco/cigarette smoke and enteric microorganisms. These microbial products interact with various types of pathogen recognition receptors (PRRs) such as the Toll-like receptors (TLRs) -2, -4, -5 and -9, nucleotide oligomerisation domain-like receptors (NODs) -1, -2 and NOD-like receptors (NLRs) P-1, -3 and -4 expressed on/in various types of immune-inflammatory cells, especially monocytes and neutrophils. PRR-mediated recognition of endotoxin and other microbial products results in systemic activation of these cells, leading, in turn, to the synthesis/release of various pro-inflammatory/apoptosis-inducing/immunosuppressive cytokines (IL-1β, IL-6, IL-8, TNF and others), reactive oxygen and nitrogen species (ROS/RNS) and the granule enzyme arginase, as well as upregulation of expression of the programmed death ligand 1 (PD-L1) on neutrophils, all of which contribute to CD4+ T cell dysfunction and apoptosis.

Figure 2: Proposed mechanism by which HIV infection causes chronic depletion of CD4+ T cells and progressive immune attrition. Notwithstanding HIV-mediated CD4+ T cell cytopathicity and destruction of these cells by HIV-specific CD8+ cytotoxic T cells, chronic exposure to viral antigens also results in activation-induced cell death and pyroptosis of bystander cells due to intracellular accumulation of incomplete viral DNA reverse transcripts. The permissiveness of follicular helper T cells to HIV infection also favours viral persistence and immune dysfunction (decreased production of specific antibodies). These various mechanisms of CD4+ T cell death and dysfunction are exacerbated by the highly pro-inflammatory environment in the lung (and other body compartments) created by HIV infection. This, in turn, results in the unremitting exposure of pulmonary CD4+ T cells to a range of pro-apoptotic mediators derived from both resident and infiltrating immune/inflammatory and structural cells. These include TNF and gamma chain cytokines released by various cell types; interferon-α and indoleamine 2,3-dioxygenase derived from plasmacytoid dendritic cells; reactive oxygen species and arginase from neutrophils and monocytes/macrophages; granzymes and perforins from CD8+ cytotoxic T cells and other types of “killer cells;” and death receptor ligands released by/present on various types of resident and infiltrating immune/inflammatory cells. These interacting mechanisms, exacerbated by microbial translocation and smoking, sustain a self-perpetuating “vicious cycle” of chronic immune activation/inflammation, CD4+ T cell death and immunosuppression. Acquired immunodeficiency is worsened by an ongoing, ineffectual, counteracting anti-inflammatory response involving regulatory T cells (Tregs) and macrophages of the M2 phenotype and their associated cytokines, transforming growth factor-β1 (TGF-β1) and interleukin-10 (IL-10), as well as myeloid suppressor cells.

*Denotes immunosuppressive/ inflammatory mechanisms potentiated by smoking, although the effects of smoking on microbial translocation remain to be conclusively established as denoted by (?).
like receptors (TLRs) -2, -4, -5 and -9, nucleotide oligomerisation domain-like receptors (NODs) -1, -2 and NOD-like receptors (NLRs) P-1, -3 and -4 expressed on/in various types of immune-inflammatory cells, especially monocytes and neutrophils. PRR-mediated recognition of endotoxin and other microbial products results in systemic activation of these cells, leading, in turn, to the synthesis/release of various pro-inflammatory/apoptosis-inducing/immunosuppressive cytokines (IL-1β, IL-6, IL-8, TNF and others), reactive oxygen and nitrogen species (ROS/RNS) and the granule enzyme arginase, as well as upregulation of expression of the programmed death ligand 1 (PD-L1) on neutrophils, all of which contribute to CD4+ T cell dysfunction and apoptosis.

Figure 1
HIV infection/replication

- Chronic CD4 T cell cytopathicity
- CD8+ T cell-mediated cytotoxicity
- Activation-induced cell death
- Pyroptosis
- Permissiveness of T(f)h cells to HIV infection

Chronic immune activation

Pro-inflammatory response *
- Cytokines (TNF, γ-chain cytokines)
- Indoleamine 2,3-dioxygenase
- Reactive oxygen species
- Arginase
- Granzymes/perforins
- Death receptor ligands

Counteracting anti-inflammatory response
- Cellular (T-regs, M2 Mø)
- Cytokines (TGF-β1, IL-10)
- Myeloid suppressor cells

T cell apoptosis

Immune suppression/attrition *

Microbial translocation *
(?)

Figure 2