

Cigarette smoke and HIV synergistically affect lung pathology in cynomolgus macaques

Hitendra S. Chand,^{1,2} Rodrigo Vazquez-Guillamet,³ Christopher Royer,¹ Karin Rudolph,¹ Neerad Mishra,¹ Shashi P. Singh,¹ Shah S. Hussain,² Edward Barrett,¹ Shannon Callen,⁴ Siddappa N. Byrareddy,⁴ Maria Cristina Vazquez Guillamet,³ Jawad Abukhalaf,³ Aryaz Sheybani,³ Vernat Exil,³ Veena Raizada,³ Hemant Agarwal,³ Madhavan Nair,² Francois Villinger,⁵ Shilpa Buch,⁴ and Mohan Sopori¹

¹Lovelace Respiratory Research Institute, Albuquerque, New Mexico, USA. ²Herbert Wertheim College of Medicine, Florida International University, Miami, Florida, USA. ³University of New Mexico Health Sciences Center, Albuquerque, New Mexico, USA. ⁴University of Nebraska Medical Center, Omaha, Nebraska, USA. ⁵New Iberia Research Center, University of Louisiana at Lafayette, New Iberia, Louisiana, USA.

In the era of combined antiretroviral therapy (cART), lung diseases such as chronic bronchitis (CB) and chronic obstructive pulmonary disease (COPD) are common among persons living with HIV (PLWH), particularly smokers. Although smoking is highly prevalent among PLWH, HIV may be an independent risk factor for lung diseases; however, the role of HIV and cigarette smoke (CS) and their potential interaction in the development of chronic lung diseases among PLWH has not been delineated. To investigate this interaction, cynomolgus macaques were exposed to CS and/or simian-adapted human immunodeficiency virus (SHIV) and treated with cART. The development of CB and the lung functions were evaluated following CS±SHIV treatment. The results showed that in the lung, SHIV was a strong independent risk factor for goblet cell metaplasia/hyperplasia and mucus formation, MUC5AC synthesis, loss of tight junction proteins, and increased expression of Th2 cytokines/transcription factors. In addition, SHIV and CS synergistically reduced lung function and increased extrathoracic tracheal ring thickness. Interestingly, SHIV infection generated significant numbers of HIV-gp120⁺ epithelial cells (HGECs) in small airways and alveoli, and their numbers doubled in CS+SHIV-infected lungs. We conclude that even with cART, SHIV independently induces CB and pro-COPD changes in the lung, and the effects are exacerbated by CS.

Introduction

Although cART has dramatically improved the life span of persons living with HIV (PLWH) and transformed the infection to more or less a medically manageable chronic disease (1), the incidence of chronic lung diseases including chronic bronchitis (CB) and chronic obstructive pulmonary disease (COPD) is high in PLWH (2, 3). While the incidence of asthma among HIV-seropositive children is high (4), the frequency of asthma in adult PLWH is debatable (5, 6). Generally, noninfectious chronic lung diseases are associated/affected by CS (7, 8), and smoking is highly prevalent among PLWH (9, 10). Epidemiological data suggest HIV as an independent risk factor for COPD (11, 12) and CS and lung infections synergize to promote COPD (13). Thus, interaction between CS and HIV may affect development of obstructive lung diseases in PLWH who are smokers.

CB or chronic mucus hypersecretion (CMH) (14) results from mucus overproduction by goblet cells, leading to airway obstruction and decline in lung function (15, 16). COPD encompasses a spectrum of diseases including CB, reduced lung function, and emphysema and its severity is linked to CMH (17). We have shown that, even after well-controlled viremia, SIV- and HIV-infected lungs contain large numbers of mucous⁺gp120⁺-immunoreactive

cells (18). Moreover, exposure of normal human bronchial epithelial (NHBE) cells to approximately 8×10^{-11} M HIV-gp120 induces CXCR4- and nicotinic acetylcholine receptor-dependent mucus production and goblet cell hyperplasia (18), suggesting a potential interaction between CS and HIV in CB development. Furthermore, HIV may infect NHBE cells either unproductively (19) or productively (20). Therefore, we investigated the interaction between CS and SHIV infection in the development of lung diseases in cynomolgus macaques.

Results and Discussion

Cynomolgus macaques were exposed to CS for 27 weeks and/or SHIV infection for 16 weeks to obtain 4 experimental groups: fresh air control (FA), CS-exposed, SHIV-infected, and CS+SHIV (see Supplemental Methods and Supplemental Figure 1; supplemental material available online with this article; <https://doi.org/10.1172/JCI121935DS1>). All SHIV-infected macaques received daily cART starting at 2 weeks postinfection until sacrifice. To determine the nature of lung cells that are affected by CS and SHIV, we enumerated HIV-gp120⁺ and pCK⁺ (epithelial cell marker) double-positive cells (i.e., HGECs) in the small airways and alveoli. While SHIV-infected animals had $9.0\% \pm 1.1\%$ and $15.8\% \pm 1.6\%$ HGECs in the airways (Figure 1A) and in alveoli (Figure 1B), respectively, CS nearly doubled the number in both airways and alveoli to $19.0\% \pm 2.2\%$ and $29.6\% \pm 2.6\%$, respectively (Figure 1, A and B). Plasma viral titers were measured at 2 weeks postinfection to analyze the effects of CS on viral replica-

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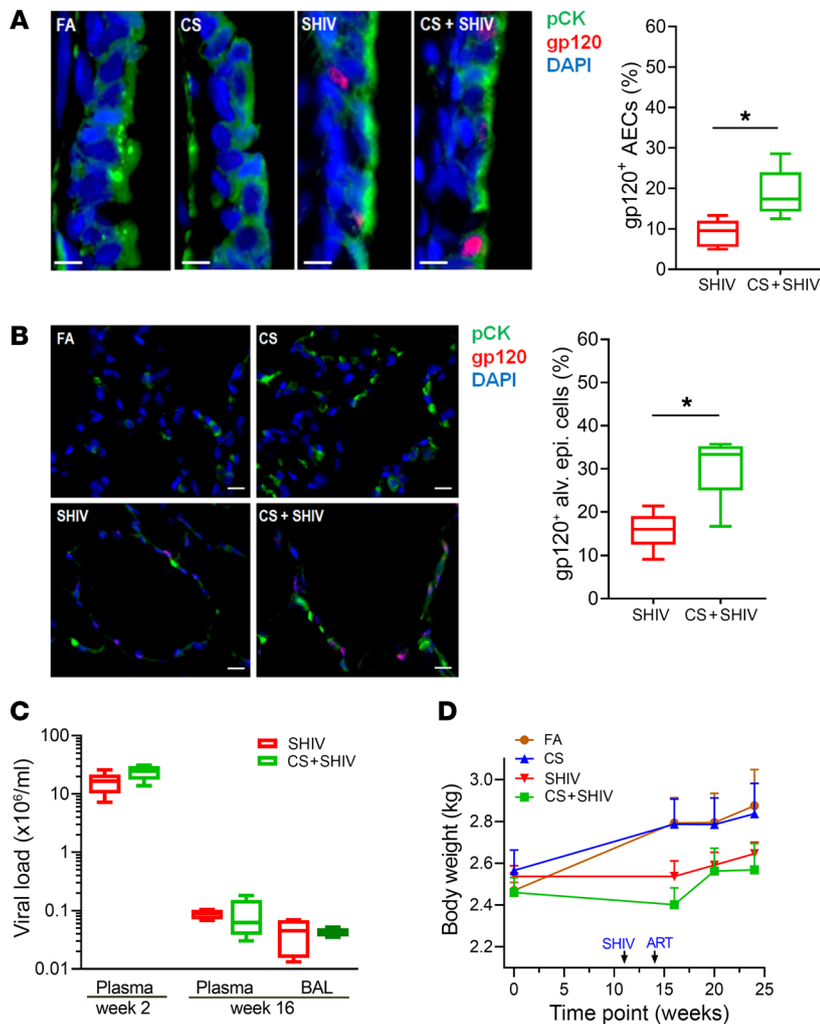


Figure 1. CS increases the number of gp120-immunopositive lung epithelial cells in SHIV-infected and cART-treated cynomolgus macaques. Formalin-fixed and paraffin-embedded (FFPE) lung tissue sections (5 μ m) from each group were immunostained for the epithelial cell marker pan-cytokeratin (pCK) and for HIV-gp120. **(A)** Representative micrographs of bronchial airway epithelial cells (AECs) showing pCK (green) and HIV-gp120 (red) immunopositivity along with the DAPI-stained nuclei (blue) from each group. Scale bars: 5 μ m. Percentages of gp120⁺ AECs were quantified. **(B)** Micrographs of alveolar region showing pCK (green) and HIV-gp120 (red) immunopositivity from each group. The percentage of gp120⁺ alveolar epithelial cells were quantified. Scale bars: 10 μ m. Data for **A** and **B** are mean \pm SEM, $n = 7$ /group; * $P \leq 0.05$ by Student's t test. **(C)** Viral titers in the plasma and in the plasma and bronchoalveolar lavage (BAL) of SHIV and CS+SHIV-exposed animals at weeks 2 and 16 post-infection, respectively. **(D)** Body weights of macaques at the baseline and at indicated times during the experiment. Data are mean \pm SEM, $n = 4$ –7/group.

tion. Titers didn't differ significantly ($P = 0.088$) between SHIV ($\sim 1.8 \times 10^7$ copies/ml) and CS+SHIV ($\sim 2.5 \times 10^7$ copies/ml) (Figure 1C). After 14 weeks of cART, the viral titers in SHIV and CS+SHIV dropped by more than 99.9% ($\sim 8 \times 10^3$ copies/ml); viral titers in bronchoalveolar lavage (BAL) were also similar (Figure 1C). At 4 weeks postinfection, while the body weight of SHIV-infected and CS-exposed animals didn't differ ($P = 0.09$), CS+SHIV animals had significantly lower body weight than animals exposed to CS alone ($P = 0.02$). Following cART, CS+SHIV animals started regaining body weight and, at 3 weeks before sacrifice, their weights did not differ significantly from CS-exposed animals (Figure 1D). These results suggest: (a) cynomolgus macaques are extremely sensitive to SHIV_{89.6} infection; (b) the viral replication is sensitive to cART (viral loads decrease by >99.9%); (c) even after successful cART, SHIV infection induces HGECs in small airways and alveoli; and (d) the number of HGECs nearly doubles in animals chronically exposed to CS. However, presently we have no direct evidence suggesting that gp120 is synthesized by the gp120-immunoreactive epithelial cells. Nonetheless, the results support our earlier observation that the lungs of PLWH contain large numbers of HGECs (18) and it is likely that CS increases HGECs by either promoting HIV infection of epithelial cells or inhibiting the clearance of gp120⁺ epithelial cells.

Moreover, CS+SHIV affected the cellularity in BAL and the lung. Thus, compared with FA, BAL from CS+SHIV animals had significantly higher cell numbers (Supplemental Figure 2A) and neutrophil count (Supplemental Figure 2C). However, there was a significant decrease in BAL macrophages in CS+SHIV group (Supplemental Figure 2B) with no changes in BAL lymphocytes (Supplemental Figure 2D). Similarly, CS+SHIV animals exhibited markedly higher peribronchial, parenchymal, and perivascular lymphoid aggregates in the lung (Supplemental Figure 3). Thus, CS and SHIV infection synergistically affect lung pathology.

Smoking is a paramount risk factor for CMH that leads to CB, decline in lung function, COPD, and asthma (8, 15, 21–23). Although smoking is highly prevalent in PLWH, epidemiological evidence suggests HIV as an independent risk factor for CB and COPD (11, 12). MUC5AC and MUC5B are the 2 major secretory mucins in the airways that contribute to CMH and development of obstructive lung diseases (17, 22). When analyzed by quantitative reverse transcription PCR (qRT-PCR), compared with FA, MUC5AC mRNA was strongly upregulated by CS and SHIV and increased synergistically by CS+SHIV (Figure 2A, left panel); however, none of the treatments affected MUC5B levels (Figure 2A, right panel). Similarly, immunostaining of lung sections showed increased MUC5AC (Figure 2B) but not MUC5B (Figure 2C) in response to CS or SHIV.

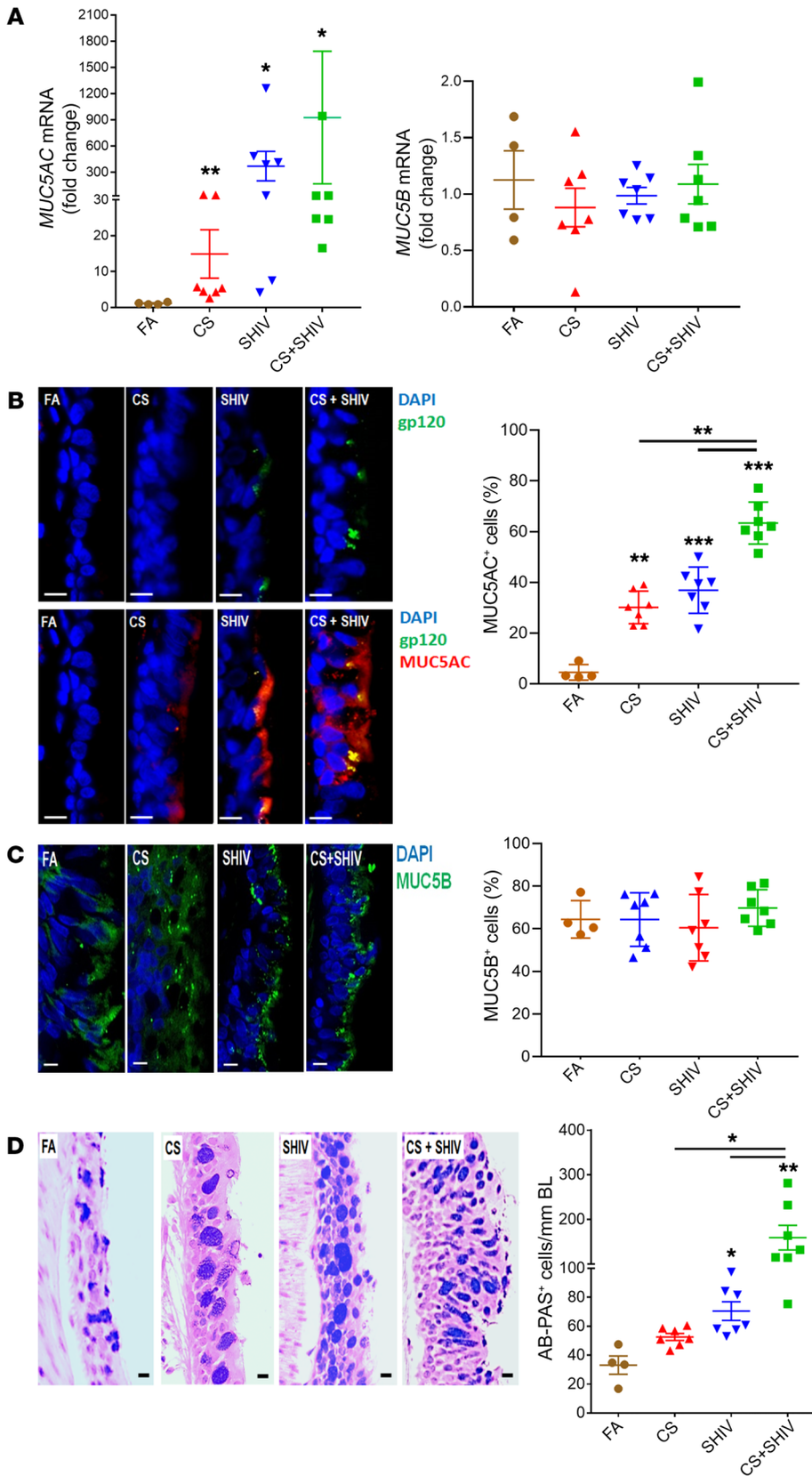


Figure 2. CS exposure and SHIV infection synergistically induce mucous cell hyperplasia and MUC5AC mucin expression.

(A) Quantification of secretory mucin *MUC5AC* and *MUC5B* mRNA levels in each group. RNA was isolated from frozen lung tissues and assessed using real-time qRT-PCR. (B) Micrographs of small airways showing MUC5AC (red) and HIV-gp120 (green) immunopositivity from each group, and the percentage of MUC5AC⁺ epithelial cells. Scale bars: 5 μm. (C) Micrographs of small airways showing MUC5B (green) immunopositivity from each group, and the percentage of MUC5B⁺ epithelial cells. Scale bars: 5 μm. (D) Representative micrographs of small airways showing mucopolysaccharides stained with AB-PAS. Scale bars: 5 μm. AB-PAS⁺ cells were quantified per unit length (mm) of basal lamina (BL). Data are mean ± SEM, n = 4–7/group, 1-way ANOVA; *P ≤ 0.05; **P < 0.01; ***P < 0.001.

restricted to goblet cells and submucosal glands, respectively. MUC5B is critical for mucociliary clearance and antibacterial responses, whereas MUC5AC is overexpressed in allergic and obstructive lung diseases (24). In humans, MUC5AC and MUC5B are not regulated synchronously (24), thus, the effects of CS and/or SHIV appear localized primarily to goblet cells. Indeed, some MUC5AC⁺ cells are also HIV gp-120 immunoreactive (Figure 2B).

Differentiation of airway goblet cells requires interplay between several transcription factors. SAM-pointed domain-containing Ets-like factor (SPDEF) and forkhead box protein A3 (FOXA3) are selectively expressed by airway epithelial cells and stimulate mucus production, goblet cell hyperplasia/metaplasia, and proasthmatic cytokines like IL-13 and IL-4 (25). Using ELISA, we found that compared with FA, IL-13 levels are significantly upregulated in CS-exposed and SHIV-infected lungs (P < 0.01); CS+SHIV further increased the expression of IL-13 significantly (Supplemental Figure 4A). While FOXA3 and SPDEF are upregulated, the expression of FOXA2 that inhibits goblet cell and Th2 responses is downregulated in asthma and COPD patients (26–29). To assess the effects of CS±SHIV on these transcription factors, we determined the *SPDEF*, *FOXA3*, and *FOXA2* mRNA levels by qRT-PCR (Supplemental Figure 4B). CS and SHIV independently increased *SPDEF*

and *FOXA3* expression, which is increased further in CS+SHIV lungs. Interestingly, *FOXA2* expression was independently reduced by CS and SHIV, and was essentially absent in CS+SHIV lungs. Thus, changes in the airway goblet cell differentiation, prompted by CS and SHIV,

restricted to goblet cells and submucosal glands, respectively. MUC5B is critical for mucociliary clearance and antibacterial responses, whereas MUC5AC is overexpressed in allergic and obstructive lung diseases (24). In humans, MUC5AC and MUC5B are not regulated synchronously (24), thus, the effects of CS and/or SHIV appear localized primarily to goblet cells. Indeed, some MUC5AC⁺ cells are also HIV gp-120 immunoreactive (Figure 2B).

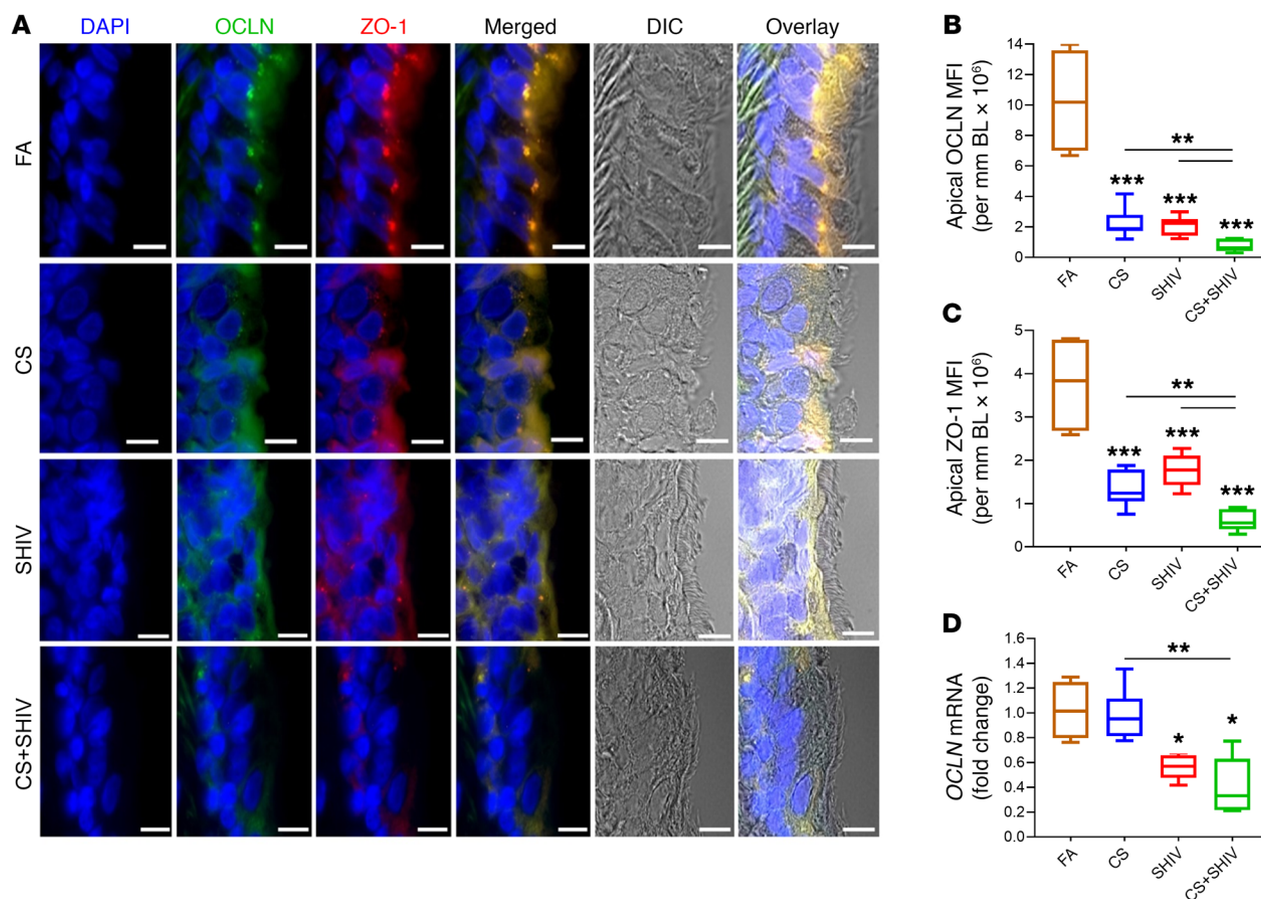


Figure 3. CS and SHIV independently and additively impair the lung epithelial tight junctions. (A) Representative micrographs of small airways showing tight junction proteins occludin (OCLN, green) and ZO-1 (red) along with the DIC (differential interference contrast) imaging. Scale bars: 5 μ m. Quantification of apical OCLN (B) and ZO-1 (C) was assessed by measuring mean fluorescence intensity (MFI) per mm basal lamina (BL) in each group. (D) Quantification of OCLN mRNA levels in each group was determined by qRT-PCR. Data in B–D are mean \pm SEM, $n = 4$ –7/group, 1-way ANOVA; * $P \leq 0.05$; ** $P < 0.01$; *** $P < 0.001$.

parallel the expression of transcription factors and cytokines that regulate mucous responses. Moreover, in human airways, IL-13 is a potent stimulator of MUC5AC but not MUC5B (24). Therefore, it is likely that SHIV infection and CS increases IL-13 expression, which promotes MUC5AC, CMH, and obstructive lung diseases. Although changes in IL-13, goblet cell hyperplasia, *FOXA2*, *FOXA3*, and *SPDEF* are compatible with proasthmatic responses, we have no direct evidence of asthma in these animals.

Given the extent of goblet cell hyperplasia in CS+SHIV airways, it was likely that the combination compromised the airway epithelial barrier that is dependent on the interaction of tight junction proteins such as occludin (OCLN) and cytosolic scaffolding proteins such as zonula occludins 1 (ZO-1) (30). Therefore, we stained airway sections for OCLN and ZO-1 and as shown in Figure 3A, the airway epithelium expressed double positive (OCLN⁺ZO-1⁺) cells in FA. However, CS and SHIV reduced the expression of OCLN and ZO-1 and the changes in the epithelial architecture were clearly evident by DIC (differential interference contrast images) and montage images of the immunohistographs (Figure 3A). Amounts of OCLN (Figure 3B) and ZO-1 (Figure 3C) were decreased independently by CS and SHIV, but the reduction was more prominent in CS+SHIV. Thus, CS and SHIV synergize to compromise the airway tight junctions that are critical in inflammatory lung diseases (31). Analysis of

OCLN mRNA by qRT-PCR (Figure 3D) indicated that, while SHIV downregulated OCLN expression, CS independently did not significantly affect the expression, and CS+SHIV was not significantly lower than SHIV alone. Therefore, CS does not affect the transcription of tight junction proteins but may affect their expression post-translationally such as through phosphorylation (32).

Increased airflow limitation is an early sign of obstructive lung diseases such as COPD (33). Decreased forced expiratory volume at 1 second (FEV₁) reflects airflow obstruction in humans; however, in smaller animals (e.g., cynomolgus macaques), FEV at shorter times is more reproducible (34). We measured FEV_{0.1} and FEV_{0.15} at baseline and at 1 week prior to sacrifice. At baseline, FEV_{0.1} and FEV_{0.15} values were comparable (data not shown). However, while the FEV_{0.1} of FA, CS, and SHIV animals did not change over time — reflecting increased lung volumes in growing animals — the values were significantly lower in CS+SHIV (Figure 4A). Similar results were observed for FEV_{0.15} (data not shown). Thus CS+SHIV cooperate to induce airflow limitations compatible with the development of COPD-like changes.

CB and decreased FEV₁ are directly related to increases in air wall thickness (35), and ultrasound (US) has been successfully used to visualize changes in upper airway structures (36). The epithelia of the cartilaginous airways are composed of mucosal glands, ciliated cells, and goblet cells, and goblet cells are the most common in the

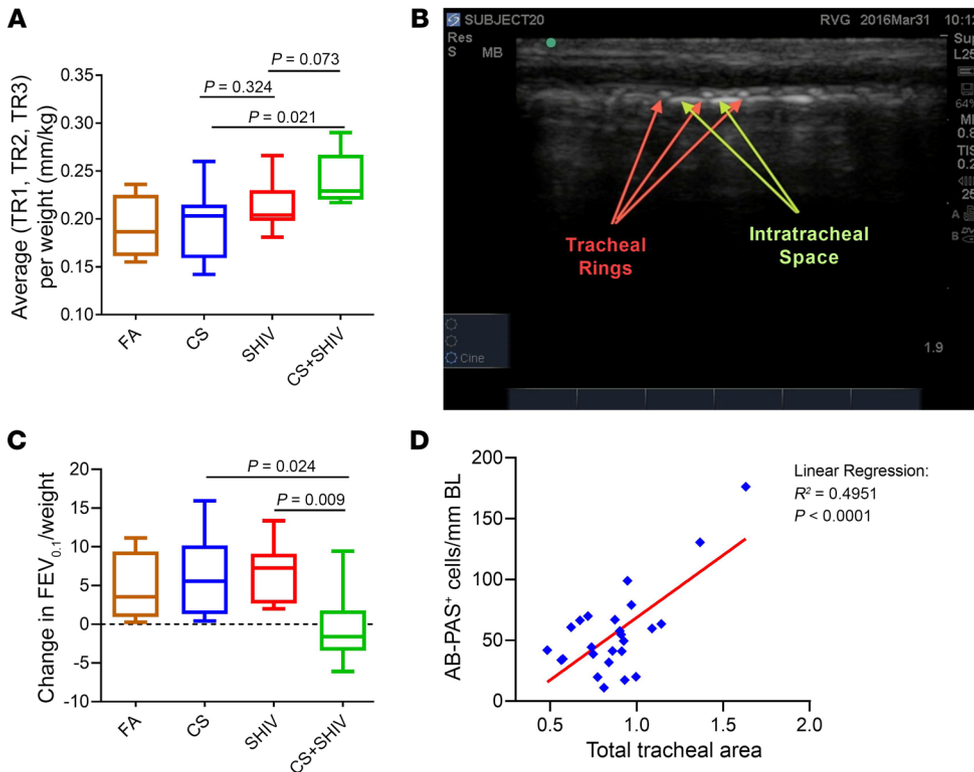


Figure 4. CS exposure and SHIV infection synergistically reduce lung function while augmenting upper airway wall thickness. (A) Changes in lung function parameters as determined by measuring FEV_{0.1} at baseline before the exposures and compared with the FEV_{0.1} values at 1 week prior to sacrifice. **(B)** Representative ultrasonograph of upper airways showing the extrathoracic tracheal wall rings (TR1, TR2, and TR3) whose thickness was measured by ultrasound imaging. **(C)** Quantification of mean tracheal ring thickness among different treatment groups shows synergistic effect of CS and SHIV exposure. Data in **A** and **C** are mean ± SEM, n = 4–7/group, 1-way ANOVA. **(D)** Linear regression analysis between airway mucous cell numbers/mm basal lamina (BL) and mean tracheal wall thickness (TR1 + TR2 + TR3) of all 25 animals.

cephalad part of the trachea (37). Changes in extrathoracic proximal (subglottic) tracheal cartilage ring size have been reported in animal models and human adults with cystic fibrosis (38, 39). We sought to determine by US imaging whether SHIV and/or CS altered the thickness of the tracheal wall at the level of the tracheal rings (TRs). Longitudinal views of the sagittal plane that spanned from the lower margin of the larynx up to the trachea inferiorly were recorded at 27 weeks of CS exposure and offered the best anatomical details (Figure 4B). Mean thickness of tracheal rings (mTRs) encompassing the first, second, and third TRs and the mean size of intertracheal space between the tracheal rings TR1 and TR2 and TR2 and TR3 were measured and normalized to body weights. Results suggested that none of the treatments caused significant changes in the size of tracheal spaces (data not shown). However, mTR of CS+SHIV was significantly higher than that of CS alone ($P = 0.02$) and was not significantly higher than that of SHIV alone ($P = 0.07$) (Figure 4C). These data suggest that CS+SHIV synergize to stimulate structural changes in the airways that are visualized by US in cynomolgus macaques. Moreover, linear regression analysis suggested a significant correlation between the number of airway mucous cells and the mean tracheal wall thickness (Figure 4D). COPD in HIV-infected patients is underdiagnosed (40) and, although COPD is a progressive disease, early stage interventions may reduce the rate of lung function decline and improve mortality (41). Although we have not done any human testing, it is possible that US of tracheal rings has the potential to monitor CMH/COPD-like changes in PLWH.

Persistent bacterial (42) and viral (43) lung infections have been proposed as COPD inducers. Adenovirus EA1 expression was correlated to accelerated COPD/emphysema in smokers but not in nonsmokers (44). However, to our knowledge, there are no published reports directly linking viral infection alone to the development of CB/COPD.

Differing lifestyles, lung infections, and nearly ubiquitous smoking in PLWH make it difficult to analyze the respiratory effects of HIV; however, epidemiological studies suggest HIV-infection as an independent risk factor for obstructive lung diseases (3, 45, 46). This study provides clear experimental evidence of HIV as an independent risk factor for development of CB- and COPD-like changes in the lung. In addition, changes in the epithelial barrier and upper airway structures underscore the inference that the lung undergoes COPD-like changes in response to CS or SHIV-infection that are accelerated in animals exposed to CS+SHIV. Furthermore, the results suggest that the lung may be an important HIV reservoir and chronic HIV infection, even after cART, promotes obstructive changes in the lung. If so, PLWH who are also smokers are at much higher risk of developing noninfectious chronic lung conditions than PLWH who are nonsmokers.

Methods

For more details, see Supplemental Methods.

Statistics. Statistical methods are described in Supplemental Methods. Briefly, grouped results were expressed as mean ± SEM and P values less than or equal to 0.05 were considered significant.

Study approval. The animals were housed at the Primate Facility of Lovelace Respiratory Research Institute, Albuquerque, NM, in accordance with the Guidelines from the Association for the Assessment and Accreditation for Laboratory Animal Care International. All experimental protocols carried out on these macaques were approved by the Institutional Animal Care and Use Committee.

Author contributions

HSC performed the sample analysis, analyzed the data, and wrote the manuscript. RVG, CR, KR, and NM performed the study and analyzed the data. SPS, SSH, and SNB performed the sample analysis and

analyzed the data. EB, SC, FV, and SB analyzed the data. MS designed the studies, analyzed the data, and wrote the manuscript. MCVG and JA read the ultrasound. AS, VE, VR, and HA read pathology slides. MN designed the HIV experiments. All authors reviewed the manuscript.

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Address correspondence to: Mohan Sopori, Lovelace Respiratory Research Institute, 2425 Ridgecrest Drive SE, Albuquerque, New Mexico 87108, USA. Phone: 505.348.9440; Email: msopori@lrri.org.

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