IL-21 and probiotic therapy improve Th17 frequencies, microbial translocation, and microbiome in ARV-treated, SIV-infected macaques

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Increased mortality in antiretroviral (ARV)-treated, HIV-infected individuals has been attributed to persistent immune dysfunction, in part due to abnormalities at the gastrointestinal barrier. In particular, the poor reconstitution of gastrointestinal Th17 cells correlates with residual translocation of dysbiotic, immunostimulatory microflora across a compromised intestinal epithelial barrier. We have previously demonstrated that oral probiotics promote increased intestinal CD4+ T-cell reconstitution during ARV treatment in a non-human primate model of HIV infection; however, essential mucosal T-cell subsets, such as Th17 cells, had limited recovery. Here, we sought to promote Th17 cell recovery by administering interleukin (IL)-21 to a limited number of ARV-treated, probiotic-supplemented, Simian Immunodeficiency Virus (SIV)-infected pigtailed macaques. We demonstrate that probiotic and IL-21 supplementation of ARVs are associated with enhanced polyfunctional Th17 expansion and reduced markers of microbial translocation and dysbiosis as compared with infected controls receiving ARVs alone. Importantly, treatment resulted in fewer morbidities compared with controls, and was independent of increased immune activation or loss of viral suppression. We propose that combining ARVs with therapeutics aimed at restoring intestinal stasis may significantly improve disease prognosis of ARV-treated, HIV-infected individuals.

INTRODUCTION

Progressive HIV infection is characterized by persistent innate and adaptive immune activation that is predictive of disease progression and that is not resolved with effective antiretroviral (ARV) therapy.1,2 Although mediators of immune activation are multi-faceted, it is evident that even with effective viral suppression, persistent immune dysfunction at the intestinal epithelial barrier permits the translocation of immunostimulatory luminal contents into intestinal tissues and systemic circulation.3 In non-human primate models of HIV infection, microbial translocation distinguishes progressive from non-progressive models of Simian Immunodeficiency Virus (SIV) infection, with progressive infection models such as rhesus (Macaca mulatta) and pigtailed (Macaca nemestrina) macaques displaying evidence of microbial translocation and immune activation in stark contrast to non-progressive hosts such as sooty mangabeys (Cercocebus atys).4–6 Importantly, although both progressive and non-progressive infected hosts exhibit some loss of intestinal CD4+ T cells, progressive models exhibit a preferential loss of Th17
cells, implicating Th17 cell loss as a primary contributor to microbial translocation and disease progression.6–8

Th17 cells are a subset of differentiated, interleukin (IL)-17-expressing memory CD4+ T cells important for control of the microbiota at mucosal barrier sites.9 Th17 cells develop from naïve CD4+ T cells in response to TGF-β and pro-inflammatory cytokine signaling induced by microbial antigens.10–12 Although Th17 differentiation can be calibrated by the microbiome,13,14 the effects of Th17 cells on gastrointestinal immunity extend beyond interplay with commensal species. Thus, while commensal species are sufficient for the generation of Th17 cells,13,14 the ability of Th17 cells to induce production of antimicrobial peptides and chemotactic cytokines from the intestinal epithelium contributes to resistance against colonization with Salmonella typhimurium, Streptococcus pneumoniae, and Candida albicans.8,9 Implicit in this counter-regulation, microbial dysbiosis—an imbalance in the relative frequencies of commensal and pathogenic species—is associated with a dysregulation of Th17 frequencies and function.15 This, in turn, is implicated in inflammation in a multitude of chronic autoimmune and infectious diseases. In HIV-infected individuals and SIV-infected macaques, microbial dysbiosis has been described as a contributor to Th17 loss and microbial translocation, with some dysbiotic species shown to be enriched in both treated and untreated subjects, and to be associated with markers of disease progression.16,17 Given the contributions of dysbiotic and translocating bacterial species to disease progression in treated, HIV-infected individuals, we previously considered that augmenting ARV therapy in chronically SIV-infected macaques with probiotics might promote a restoration of the commensal microbiota, initiating a sequence of immune restorative processes. Although probiotic supplementation was associated with intestinal CD4+ T-cell restoration and enhanced Th17 functionality, we did not observe a significant improvement in the frequency of Th17 cells or was there any discernible impact on the relative frequencies of fecal microbial communities.18

Subsequent to priming by APCs, Th17 development is augmented by autocrine production of the common γ-chain cytokine, IL-21.19,20 Although not essential for the differentiation of Th17 cells,14,21 blockade of IL-21 during Th17 development limits the expansion and functional capacity of Th17 cells.19,20 In progressive SIV infection, a paucity of IL-21-producing CD4+ T cells correlates with Th17 loss.22 Moreover, although IL-21 treatment during acute SIV infection has been shown to delay Th17 loss independent of ARV therapy, cessation of IL-21 administration results in a rapid decline in Th17 frequencies.23

Herein, we considered that augmenting ARV therapy with combination probiotic and IL-21 therapy might enhance the recovery and maintenance of Th17 cells in chronic SIV infection. As described, we observed that this regimen is associated with increased frequencies of intestinal polynuclear Th17 cells as compared with ARV-only controls, and that supplementation is further associated with improvements in microbial translocation, microbial dysbiosis, and clinical status. Importantly, as our therapy did not promote immune activation or negate viral suppression, we believe that our findings suggest that probiotic and IL-21 supplementation of ARVs may translate into a safe and effective therapeutic for the reduction of co-morbidities suffered by ARV-treated, HIV-infected individuals.

RESULTS
Experimental design
We previously demonstrated that treatment of SIV-infected, virus-suppressed pigtailed macaques (PTMs) with probiotics improved gastrointestinal integrity and immunity; however, probiotic supplementation alone did not promote a restoration of Th17 cells nor was there evidence for a shift toward a significantly less inflammatory microbiome.18 Here, we sought to determine whether treatment with IL-2119,20 enhances probiotic treatment in SIV-infected, ARV- and probiotic-treated PTMs by promoting Th17 recovery and reducing microbial translocation and dysbiosis. As depicted in Figure 1, we infected 11 PTMs with SIVmac239 and, at day 98 post infection (p.i.), all animals began daily treatment with the reverse transcriptase inhibitors, tenofovir and emtricitabine, and the integrase inhibitor, L’812. Of these animals, six were simultaneously treated with the daily probiotic, VLS#3 (consisting of 1011 live Bifidobacterium, Lactobacillus, and Streptococcus species) and with early and late administration courses of IL-21 to assess refractoriness to continued IL-21 therapy. We routinely sampled the blood and stool of all animals, sampled the pulmonary airways through bronchoalveolar lavage (BAL), and obtained jejunal (Jej), lymph node (LN), and rectal (RB) biopsies.

IL-21 and probiotic supplementation does not alter viral suppression in ARV-treated macaques
IL-21 is a pleiotropic cytokine known to promote the development of Th17 cells from naïve CD4+ T-cell precursors.19,20 As mature CD4+ T cells are the preferential targets for HIV and SIV infection, we assessed the impact of the IL-21 and probiotic regimen on viral suppression. With the initiation of therapy, all animals exhibited a rapid decline in plasma viral RNA with all but one probiotic- and IL-21-supplemented animal (PTA2P031) achieving suppression to below 100 copies per ml within 30 days of treatment initiation (Figure 2a). In
Interleukin (IL)-21 and probiotic therapy does not alter viral load or cellular distribution in antiretroviral (ARV)-treated pigtailed macaques (PTMs). (a) Longitudinal, log copies per ml of plasma viral RNA (vRNA) from animals treated with ARVs only (open squares) or with ARVs, probiotic VSL#3, and IL-21 (closed circles). Gray shading indicates treatment initiation and duration. Dotted lines denote IL-21 administration time points. Symbols and shading are consistent throughout the manuscript unless otherwise noted. (b) Copies of viral DNA (vDNA) per 100 colonic CD4^+ TM at the time of necropsy. Triangles indicate samples normalized to the limit of detection.

Figure 2  Interleukin (IL)-21 and probiotic therapy does not alter viral load or cellular distribution in antiretroviral (ARV)-treated pigtailed macaques (PTMs). Percent complication free status in antiretroviral (ARV)-treated pigtailed macaques (PTMs). Percent of animals remaining free from clinical complications over time. Initial complications per animal are noted alongside changes in the incidence-free curve, with ARV-only animals represented by the dashed line and supplemented animals by the solid line. Significance in the time to the first reported clinical complication between groups was assessed by the Mantel-Cox Log-Rank test.

Figure 3  Interleukin (IL)-21 and probiotic therapy improves clinical status in antiretroviral (ARV)-treated pigtailed macaques (PTMs). Percent complication free status in antiretroviral (ARV)-treated pigtailed macaques (PTMs). Percent of animals remaining free from clinical complications over time. Initial complications per animal are noted alongside changes in the incidence-free curve, with ARV-only animals represented by the dashed line and supplemented animals by the solid line. Significance in the time to the first reported clinical complication between groups was assessed by the Mantel-Cox Log-Rank test.
compared the time to the first diagnosed clinical complication in ARV-only (median day 128 p.i.) and probiotic-, IL-21-supplemented (no complication through study termination) animals and determined that supplementation was associated with a significant difference in the time to first complication ($P = 0.0316$, Mantel-Cox). These results indicate that probiotic and IL-21 supplementation of ARV therapy decreased the incidence of non-AIDS morbidities.

**IL-21 and probiotic supplementation promote polyfunctional Th17 recovery in ARV-treated macaques**

In ARV-treated individuals and macaques, persistent microbial translocation and dysbiosis are associated with continued immune activation, thereby possibly exacerbating lentiviral-mediated Th17 losses. As our previous findings indicated that probiotic supplementation of ARV therapy alone was unable to promote significant restoration of Th17 cells, we considered that additional supplementation with IL-21 might promote sustained restoration of these cells. We assessed the frequency of IL-17-expressing memory CD4$^+$ T cells (Th17) longitudinally in the PBMCs, BAL, Jej, LN, and RB. Among BAL, LN, and PBMC CD4$^+$ T cells, probiotic- and IL-21-supplemented animals exhibited no significant differences in Th17 frequencies as compared with ARV-only controls or as compared with the pre-treatment time point of day 84 p.i. (data not shown). Indeed, preferential loss of Th17 within chronically HIV/SIV-infected individuals is not observed within these specific anatomic sites. However, modest to significant improvements in the frequencies of intestinal Th17 cells were observed in supplemented animals as compared with the pre-treatment baseline time point (Figure 4a and b and Supplementary Figure S3A). Coupled with increased intestinal CD4$^+$ T-cell frequencies (Supplementary Figure S2), the improved relative frequencies of intestinal Th17 cells led to significantly improved absolute frequencies of intestinal Th17 cells (% IL-17-expressing CD4$^+$ TM of CD3$^+$ T cells). We observed increases in the absolute frequency of Jej Th17 cells from day 84 p.i. to days 274 and 365 p.i. ($P = 0.016, 0.031$) and in rectal Th17 cells from day 84 p.i. to days 126, 274, and 365 p.i. ($P = 0.031, 0.031, 0.016$; Figure 4a and b) within the probiotic- and IL-21-supplemented animals. No significant differences were evident in ARV-only controls (Supplementary Table S1). Furthermore, at day 182 p.i. (98 days post-intervention), ARV-only controls exhibited a significantly lower absolute frequency of Jej Th17 cells as compared with probiotic- and IL-21-supplemented animals ($P = 0.0286$). Superior control of disease progression in HIV-infected individuals has been attributed to improved functionality (multiple cytokine expression, or polyfunctionality) among T cells. As such, we further investigated the frequency of intestinal, polyfunctional Th17 cells—as assessed here by the additional expression of IL-2, IL-22, or tumor necrosis factor-α—in our animals. Although no significant differences were observed between groups in the expression of individual cytokines (data not shown), significant differences in the relative frequency of polyfunctional Jej Th17 cells were noted exclusively among probiotic- and IL-21-supplemented animals at necropsy as compared with days 84 and 274 p.i. ($P = 0.031, 0.031$; Supplementary Figure S3B, Supplementary Table S1). As compared with ARV-only controls, the relative frequencies of polyfunctional Th17 cells were lower in supplemented animals at day 274 p.i. within the Jej ($P = 0.033$) and RB ($P = 0.032$) but comparable at all other time points. However, an examination of absolute intestinal polyfunctional Th17 frequencies revealed significant increases specific to probiotic- and IL-21-supplemented animals after the initiation of therapy (Figure 4c). Within Jej tissues, significant differences in the absolute frequencies of polyfunctional Th17 cells were observed from day 84 p.i. to days 274 and 365 p.i. ($P = 0.031, 0.031$) in supplemented animals, in whom a significantly higher absolute frequency of polyfunctional Th17 cells was observed as compared with ARV-only controls at day 182 p.i. ($P = 0.029$). As compared with day 84 p.i., differences were additionally observed among probiotic- and IL-21-supplemented animals in the absolute frequency of RB polyfunctional Th17 cells at days 126, 274, and 365 p.i. ($P = 0.031, 0.031, 0.031$). No significant differences as compared with baseline were observed among the ARV-only controls at any time point (Supplementary Table S1). Thus, probiotic and IL-21 supplementation of SIV-infected, ARV-treated macaques led to a significant improvement in the absolute frequencies of intestinal Th17 cells and in turn, the absolute frequencies of polyfunctional Th17 cells.

As IL-21 similarly promotes the differentiation and maintenance of IL-17-expressing memory CD8$^+$ T cells (Tc17), we additionally examined the longitudinal relative frequencies of Tc17 cells. No significant differences between experimental groups were noted at any time point in any tissue examined (Supplementary Figure S3C and data not shown). However, a significant improvement in Jej Tc17 cells was noted in the probiotic- and IL-21-supplemented animals at day 274 p.i. as compared with the pre-treatment time point ($P = 0.0469$)—no differences were noted in relative RB Tc17 frequencies between or among groups. We similarly examined the relative frequencies of polyfunctional Tc17 cells. No differences were noted between experimental groups and, again, significant changes were limited to Jej Tc17 cells in the probiotic- and IL-21-supplemented animals (Supplementary Figure S3D). Significantly higher relative frequencies of polyfunctional, Jej Tc17 cells were noted in these animals at days 274 and 365 p.i. as compared with day 84 p.i. ($P = 0.031, 0.031$). These data indicate that probiotic- and IL-21-supplementation of ARV therapy promotes a recovery of Th17 and Tc17 cells during chronic SIV infection in macaques.

**IL-21 and probiotic supplementation reduce microbial translocation and dysbiosis in ARV-treated macaques**

Persistent immune activation, Th17 depletion, residual microbial translocation, and dysbiosis form a synergistic axis in ARV-treated individuals that likely results in the increased morbidities and mortality observed in ARV-treated subjects.
As our results indicate that probiotic and IL-21 supplementation of ARVs promotes the recovery of intestinal Th17 cells (Figure 4), tempers some markers of immune activation (Supplementary Figure S1), and results in better clinical outcomes (Figure 3), we next considered the possibility that such treatment might further be associated with a reduction in microbial translocation and dysbiosis. We assessed levels of microbial translocation in our animals by immunohistochemical staining for *Escherichia coli* antigens in the colonic lamina propria, mesenteric LN paracortex, and liver tissue from samples obtained at necropsy. *E. coli* antigens were evident in all tissues and only within the liver of probiotic- and IL-21-supplemented animals was there a trend for lower levels of *E. coli* staining ($P = 0.0556$; Supplementary Figure S4 A + B).

To assess whether probiotic and IL-21 supplementation influence the composition of microbial communities, we measured relative levels of commensal *Bifidobacterium* and *Lactobacillus* 16S DNA in the stool and rectal tissues of our experimental animals. No significant differences were observed in the ARV-only animals as compared with baseline with regards to *Bifidobacterium* levels ($P = 0.0625$, all time points), but these animals exhibited significantly lower frequencies of *Bifidobacterium* as compared with probiotic- and IL-21-supplemented animals at day 154 p.i. ($P = 0.0333$; Figure 5a). Importantly, probiotic- and IL-21-treated animals exhibited significant increases in the relative frequency of *Bifidobacterium* species as compared with the pre-treatment time point at days 154 and 365 p.i. ($P = 0.0156, 0.0313$).

Supplemented animals additionally exhibited significant increases in *Lactobacillus* species at day 154 p.i. as compared with the pre-treatment time point ($P = 0.0156$; Figure 5b)—no significant differences were noted as compared with or among ARV-only animals ($P = 0.0625$, all time points). Thus, probiotic and IL-21 supplementation of ARV therapy during chronic SIV infection promoted a reduction in distal site microbial translocation as well as an increase in the level of probiotic species within the gastrointestinal tract, reflecting a significant improvement in disease progression.

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**Figure 4** Interleukin (IL)-21 and probiotic therapy promotes the absolute expansion of polyfunctional Th17 cells in ARV-treated PTMs. (a) Longitudinal mean (± s.e.m.) percent IL-17$^+$ of CD4$^+$ TM (Th17) from the jejunal (Jej; left) and rectal biopsies (RB; right). (b) Absolute percent Th17 (that is, the percent of IL-17$^+$ CD4$^+$ TM found within CD3$^+$ T cells) from the Jej and RB at the indicated days p.i. (c) Absolute percent polyfunctional Th17 from the Jej and RB at the indicated days p.i. Line through samples represents mean; lines above samples span time points of significance. Significance within groups was assessed by Wilcoxon Signed-Rank test. Significance across groups was assessed by Mann–Whitney $U$-test.
Reduced indoleamine-2,3-dioxygenase (IDO) activity in IL-21-, probiotic-, and ARV-treated macaques

The persistent depletion of Th17 cells in ARV-treated, HIV-infected individuals correlates with elevated expression and activity of IDO, a cytokine-inducible, tryptophan (Trp)-catabolizing enzyme.16,34 Trp metabolites are known to inhibit the generation and maintenance of Th17 cells,34 and persistently elevated IDO activity has been postulated to contribute to poor Th17 recovery in HIV-infected individuals16,34 while also predicting poor CD4⁺ T-cell recovery and increased mortality.

To determine whether probiotic and IL-21 supplementation reduced IDO expression and activity, we first quantified IDO expression by immunohistochemical staining in colon and mesenteric LN tissue samples obtained at necropsy. Tissue staining revealed no significant differences in mesenteric LN IDO between the two groups and only a trend for reduced colonic IDO in probiotic and IL-21 supplemented animals as compared with ARV controls (P = 0.0857; Supplementary Figure S4C + D). We next considered whether Th17 recovery might correspond with a reduction in circulating levels of kynurenine (Kyn), the first immunoactive Trp catabolite generated in the IDO pathway. When we measured the concentration of plasma Kyn, we observed that whereas ARV-only animals exhibited a steady increase in Kyn throughout the study period, probiotic- and IL-21-supplemented animals displayed reduced Kyn from days 154 to 365 p.i. (Figure 5c). Although no significant differences were noted between experimental groups at the time points examined, significantly lower Kyn was apparent between these later time points within the supplemented group (P = 0.0469). To ascertain whether the observed Kyn levels might result from reduced IDO expression and/or activity, we additionally measured the ratio of plasma Kyn/Trp. Similar to the pattern observed for Kyn itself, although ARV-only animals exhibited a steady increase in plasma Kyn/Trp throughout the experimental period, probiotic- and IL-21-supplemented animals exhibited an increase only through the final examined necropsy time point, day 154 p.i. (Figure 5d). There was no significant difference between the final two time points, however, although supplemented animals exhibited a plateau in Kyn/Trp levels. Thus, probiotic and IL-21 supplementation of ARVs during chronic SIV infection is associated with a trend toward reduction in intestinal IDO and an apparent reduction in circulating levels of the Trp catabolite, Kyn.

DISCUSSION

ARV treatment significantly prolongs the length and quality of life of HIV-infected individuals; however, disease progression in these individuals remains evident even with decades of effective viral suppression.12 In treated HIV-infected individuals, morbidities and mortality have been attributed to a self-reinforcing cycle of microbial translocation and dysbiosis, immune activation, and a persistent loss of antimicrobial Th17
cells.\(^3,35\) Although we have previously demonstrated that probiotic supplementation of ARVs in SIV-infected macaques promotes a restoration of intestinal CD4\(^+\) T cells,\(^18\) in our prior study we did not observe significant Th17 expansion or was there a measurable impact on microbial translocation or on the composition of the gastrointestinal tract microbiome. Herein, we demonstrate that the supplementation of ARVs with probiotics and IL-21 promotes intestinal Th17 recovery, a reduction in measures of microbial translocation, and an enhanced microbiome. Importantly, our treatment regimen did not promote immune activation or a significant expansion of the viral reservoir. Although limited in size, these findings suggest that probiotic and IL-21 supplementation of ARV therapy should be further evaluated as a therapeutic in the treatment of non-AIDS-related co-morbidities.

The development of Th17 cells from naïve CD4\(^+\) T-cell precursors or T\(_{reg}\) is augmented by the autocrine production of IL-21.\(^19–21\) In this capacity, IL-21 enhances ROR\(\gamma\text{t}\) transcription, FOXP3 suppression, and the downstream transcription of IL-17 and additional Th17-associated cytokines such as IL-22.\(^19,20\) Importantly, IL-21 alone is insufficient to induce Th17 development,\(^21\) requiring additional signaling originating from the commensal microbiome. Thus, in murine models, although the presence of the commensal microbiome—specifically, segmented filamentous bacteria—is sufficient to drive the development of Th17 cells \(in\;vivo\),\(^13,14\) an absence or blockade of IL-21 signaling diminishes Th17 differentiation.\(^19,20\) That the inclusion of IL-21 in our regimen promotes Th17 reconstitution (Figure 4) where probiotic supplementation of ARVs alone does not\(^18\) indicates that therapeutically provided probiotic species alone are either insufficient or incorrectly parlayed into the development of Th17 cells during chronic SIV infection. We believe that the prolonged regeneration of Th17 cells in our animals subsequent to IL-21 withdrawal (\(\sim7\) months between early and late administration; Figure 1) suggests that IL-21 functioned directly at the level of CD4\(^+\) T cells to establish a basal threshold of Th17 cells necessary for self-propagation. Although a direct assessment of circulating IL-21 was not performed, attainment of a theoretical threshold is further supported by the observation that late IL-21 administration did not augment already enhanced Th17 frequencies (Figure 4). To this end, it would be of significant interest to determine whether increasing the concentration of administered IL-21 or lengthening early IL-21 therapy might not augment or accelerate Th17 recovery beyond what we describe here. Although we cannot discount alternative roles for IL-21 in improving disease progression in our supplemented animals—such as altered functionality among IL-21-responsive B cells and NK cells\(^36\)—the administration of IL-21 during acute SIV infection in macaques leads to a selective maintenance of Th17 cells that is strictly dependent on the continued administration of IL-21,\(^23\) arguing against an ancillary role for IL-21 in sustaining Th17 frequencies. An assessment of the effects of IL-21 supplementation to ARV therapy independent of probiotic administration will significantly further our understanding of the conditions required for the generation and maintenance of Th17 cells in treated chronic lentiviral infections.

Although \(de\;novo\) Th17 development from naïve CD4\(^+\) T-cell precursors is largely reliant on the ability of APCs to integrate microbial sensing into pro-Th17 cytokine signaling,\(^10,11\) Th17 and T\(_{reg}\) cells exist as a transcriptional dichotomy,\(^19,20,37,38\) which may further be reciprocally programmed by aryl hydrocarbon receptor ligand signaling.\(^34,39\) As recent findings have demonstrated a role for the IDO-derived aryl hydrocarbon receptor-agonist Kyn in reducing the Th17/T\(_{reg}\) during lentiviral infection,\(^34\) reduced plasma Kyn (Figure 5c) in our supplemented animals suggests that Th17 recovery may have included mechanisms adjunct to the direct influence of therapeutic IL-21. Given that species of the dysbiotic microbiota encode enzymatic homologues to primate Trp-catabolizing enzymes,\(^16\) we cannot discount the possibility that an IL-21-mediated recuperation of Th17 cells preempted a reduction of dysbiotic species and, hence, Kyn. Indeed, the prolonged recovery of commensal \textit{Bifidobacteria} in our supplemented animals (Figure 5b) in contrast to animals receiving probiotic supplementation alone\(^18\) suggests a role for IL-21 in promoting the colonization of probiotic species. However, as a replete commensal microbiome is itself necessary for Th17 maintenance,\(^14\) prolonged Th17 recovery after IL-21 cessation may thus be dependent upon a reduction in the frequency of dysbiotic species.

Microbial translocation has been extensively described as a significant contributor to immune activation and disease progression in chronic lentiviral infections.\(^3\) In HIV-infected humans and SIV-infected non-human primates, the influence of microbial translocation on immune activation has been causally associated by the correlation of disseminated LPS with systemic interferon-\(\gamma\).\(^4,5\) and formally demonstrated by the prevention of immune activation with acute use of the LPS-sequestering drug sevelamer.\(^40\) Intriguingly, Th17 recovery (Figure 4) and improved disease progression (Figure 3) occurred independent of systemically reduced \textit{E. coli} antigen (albeit a trend for reduced \textit{E. coli} in the liver; Supplementary Figure S4A + B) or persistent immune activation (Supplementary Figure S1). In this regard, immune activation in our supplemented animals may reflect improved immune function rather than a manifestation of innate cell refractoriness or lymphocyte exhaustion.\(^4,41,42\) Indeed, as compared with ARV-only controls, supplemented animals exhibited control over rectal CD4\(^+\) T-cell proliferation (Supplementary Figure S1A) concomitant with improved CD4\(^+\) T-cell reconstitution (Supplementary Figure S2), which is suggestive of homeostatic restoration. A more detailed assessment of immune function, particularly alterations in inflammatory biomarkers, will be necessary to address these hypotheses.

Increased mortality among HIV-infected individuals remains evident even with decades of effective viral suppression,\(^2\) especially in conditions when the IDO-mediated kynurenine pathway is induced and microbial dysbiosis may be present.\(^26\) Consistent with these observations in humans, co-morbidities were evident in our ARV-only animals.
where, despite continued viral suppression (Figure 2), three of five animals displayed repeated complications. That none of our supplemented animals displayed complications despite co-housing with the ARV-only controls is particularly notable given the propensity of PTMs toward microbial translocation and systemic immune activation, even in the absence of SIV infection.49 Although the recovery of Th17 cells in our supplemented animals (Figure 4) likely contributed to improved resistance to co-infection,8,6 a association between Th17 recovery and ARV tolerance is more tenuous. Indeed, ARV complications were reduced in animals supplemented with probiotics alone, where Th17 recovery was absent.18 Although the presence of commensal bacterial species has been demonstrated to improve the efficacy of anticancer therapeutics such as cyclophosphamide,44,45, the influence of probiotics and commensal species in mediating pharmaceutical toxicities has not been formally explored. A thorough characterization of the contributions of our individual therapeutic constituents—including individual probiotic species—to immune function in the presence and absence of ARVs will be necessary to fully uncover the mechanisms, which govern disease progression in treated chronic lentiviral infections.

In summary, we demonstrate that probiotic and IL-21 supplementation of ARV therapy in chronically SIV-infected macaques improves clinical outcomes, as manifested by a reduction in non-AIDS associated morbidities. Importantly, improved clinical outcomes were accompanied with an increase in the absolute frequency of intestinal, polyfunctional Th17 cells, and reduced measures of intestinal microbial dysbiosis. A larger, mechanistic dissection of the immune processes associated with improved disease progression in our supplemented animals will significantly inform the design of future therapeutic treatments for the treatment of HIV-infected individuals.

**METHODS**

**Animal infections and interventions.** Eleven, healthy pigtail macaques (Macaca nemestrina) were assessed for this study and infected with 3,000 TCID₅₀ SIVmac239. At day 98 p.i., all animals began a daily ARV regimen consisting of tenofovir and emtricitabine (30 mg kg⁻¹, twice-daily)—tenofovir was reduced to 20 mg kg⁻¹ after 30 days. Of these animals, 6 additionally received daily probiotic VSL#3 (oral, twice-daily)—tenofovir was reduced to 20 mg kg⁻¹ after 30 days. On a threshold cutoff of 100 cells of the parent population. For Th17 functional analyses, cytokine gates were established on a threshold cutoff. Acquired data were analyzed using FlowJo software (Treestar, Ashland, OR). To quantify cell subset frequencies, we used a threshold cutoff.
Primary antibodies were diluted 1:5,000 in blocking buffer and incubated for 1 h at room temperature. Tissue sections were washed, and either Rabbit Polink-1 or Polink-2 staining systems (Golden Bridge International) were applied for 30 min at room temperature. Sections were developed with Impact 3,3′-diaminobenzidine (Vector Laboratories, Burlingame, CA), counterstained with hematoxylin and mounted in Permount (Fisher Scientific, Hampton, NH). All stained slides were scanned at high magnification (×400) using the Aperio AT2 System (Leica Biosystems, Wetzlar, Germany) yielding high-resolution data for the entire tissue section. Representative high-magnification (400 ×) images were acquired from these whole-tissue scans.

**Quantification of systemic inflammatory markers.** Concentrations of Kyn and Trp were quantified from plasma using commercially available ELISA kits (Antibodies Online, Atlanta, GA) of LPS-binding protein and sCD14 were assessed from plasma using Permount (Fisher Scientific, Hampton, NH). All stained slides were scanned at high magnification (×400) using the Aperio AT2 System (Leica Biosystems, Wetzlar, Germany) yielding high-resolution data for the entire tissue section. Representative high-magnification (400 ×) images were acquired from these whole-tissue scans.

**Assessment of bacterial 16S DNA.** 16S DNA was isolated and quantified from stool or from luminal swabs as previously described.17

**Statistics.** Statistical analyses were performed using Prism (v6.0; GraphPad Software, La Jolla, CA). The Mann–Whitney U-test was used for comparisons between groups; the Wilcoxon Signed-Rank test for paired samples; and the Chi-squared test for categorical variables. The level of significance was set at P < 0.05. All values are presented as mean ± s.e.m.

**SUPPLEMENTARY MATERIAL** is linked to the online version of the paper at http://www.nature.com/mi

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**DISCLOSURE**

The authors declared no conflict of interest.

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